

Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics

USER GUIDE

Publication Number MAN0018703

Revision B.0

IVD

For In Vitro Diagnostic Use.

ThermoFisher
S C I E N T I F I C



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Products manufactured in Singapore:

Ion PGM™ Dx Instrument System
Ion PGM™ Dx Sequencer
Ion OneTouch™ Dx Instrument
Ion OneTouch™ ES Dx Instrument
Ion PGM™ Dx Chip Minifuge (120V)
Ion PGM™ Wireless Scanner
Ion Torrent™ Server
Torrent Suite™ Dx Software
Veriti™ Dx 96-well Thermal Cycler, 0.2 mL

Products manufactured in Frederick:

Oncomine™ Dx Target Test Kit
Ion Torrent Dx FFPE Sample Preparation Kit
Ion PGM™ Dx Library Kit
Ion OneTouch™ Dx Template Kit
Ion PGM™ Dx Sequencing Kit
Ion 318™ Dx Chip
Ion OneTouch™ Rack Kit
DynaMag™ Dx 96-Well Plate Magnet
DynaMag™ Dx 16 2-mL Magnet

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Revision history: Pub. No. MAN0018703

Revision	Date	Description
B.0	13 September 2021	Updated with addition of claims and supporting material for EGFR exon 20 insertion variants.
A.0	26 August 2021	New Oncomine™ Dx Target Test user guide for FDA submission in support of IDH1 SNV claims.

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About this guide

IMPORTANT! Before using this product, read and understand the information in the “Safety” appendix in this document.

Purpose of this guide

This user guide describes the intended use, theory of operation, and performance characteristics of the Oncomine™ Dx Target Test when used on the Ion PGM™ Dx System. This guide also includes a list of DNA variants and fusion isoforms targeted by the test for companion diagnostic (CDx) claims, as well as variants that show evidence of clinical significance.

Oncomine™ Dx Target Test Kit user guides

This user guide is part of a five-guide set.

- *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*
- *Oncomine™ Dx Target Test Part II: Sample and Library Preparation User Guide*
- *Oncomine™ Dx Target Test Part III: Template Preparation User Guide*
- *Oncomine™ Dx Target Test Part IV: Sequencing and Results Reports User Guide*
- *Oncomine™ Dx Target Test Part V: Torrent Suite™ Dx Software 5.12.5 Reference User Guide*

All five guides are required to complete the entire Oncomine™ Dx Target Test workflow.

Note: The procedures in these guides supersede the instructions in the *Ion PGM™ Dx System User Guide* when using the Ion PGM™ Dx System with the Oncomine™ Dx Target Test.



Product information

Oncomine™ Dx Target Test

The Oncomine™ Dx Target Test is an *in vitro* diagnostic next-generation sequencing test to detect somatic alterations in human DNA and RNA isolated from non-small cell lung cancer (NSCLC) and DNA isolated from cholangiocarcinoma (CC), in formalin-fixed, paraffin-embedded (FFPE) tumor tissue samples. Detection of these variants is performed using the Ion PGM™ Dx System.

The Oncomine™ Dx Target Test (Cat. No. A51695) provides a set of primers in two panels that target key regions of 23 genes related to NSCLC and one related to CC.

Intended use

The Oncomine™ Dx Target Test is a qualitative *in vitro* diagnostic test that uses targeted high-throughput, parallel-sequencing technology to detect single nucleotide variants (SNVs), deletions, and insertions in 23 genes from DNA and fusions in ROS1 and RET from RNA isolated from formalin-fixed, paraffin-embedded (FFPE) tumor tissue samples from patients with non-small cell lung cancer (NSCLC), and IDH1 SNVs from FFPE tumor tissue samples from patients with cholangiocarcinoma (CC) using the Ion PGM™ Dx System.

The test is indicated as a companion diagnostic to aid in selecting NSCLC and CC patients for treatment with the targeted therapies listed in Table 1 in accordance with the approved therapeutic product labeling.

Table 1 List of variants for therapeutic use

Tissue type	Gene	Variant	Targeted therapy
Non-small cell lung cancer (NSCLC)	BRAF	BRAF V600E mutations	TAFINLAR® (dabrafenib) in combination with MEKINIST® (trametinib)
	EGFR	EGFR L858R mutation, EGFR Exon 19 deletions	IRESSA® (gefitinib)
	EGFR	EGFR Exon 20 insertions	EXKIVITY™ (mobocertinib)
	RET	RET fusions	GAVRETO™ (pralsetinib)
	ROS1	ROS1 fusions	XALKORI® (crizotinib)
Cholangiocarcinoma (CC)	IDH1	IDH1 R132C, IDH1 R132G, IDH1 R132H, IDH1 R132L, and IDH1 R132S	TIBSOVO® (ivosidenib)

Safe and effective use has not been established for selecting therapies using this device for the variants other than those listed in Table 1.

Results other than those listed in Table 1 are indicated for use only in patients who have already been considered for all appropriate therapies (including those listed in Table 1). Analytical performance using NSCLC specimens has been established for the variants listed in Table 2.

Table 2 List of variants with established analytical performance only

Gene	Variant ID/type	Amino acid change	Nucleotide change
KRAS	COSM512	p.Gly12Phe	c.34_35delGGinsTT
KRAS	COSM516	p.Gly12Cys	c.34G>T
MET	COSM707	p.Thr1010Ile	c.3029C>T
PIK3CA	COSM754	p.Asn345Lys	c.1035T>A

The test is not indicated to be used for standalone diagnostic purposes, screening, monitoring, risk assessment, or prognosis.

Theory of operation

Overview

DNA and RNA are isolated from tumor tissue samples prepared as FFPE sections on slides. The amounts of DNA and RNA in a sample are quantified, and if they meet the minimum required amounts for the test, cDNA is prepared from the RNA. The DNA and cDNA are made into amplicon libraries using the Ion PGM™ Dx Library Kit and the Oncomine™ Dx Target Test DNA and RNA Panel, which target the variants and gene fusions of interest for the test. No-template libraries and control libraries specific to the test are also prepared.

Each amplicon library is templated onto Ion PGM™ Dx Ion Sphere™ Particles (ISPs), loaded onto an Ion 318™ Dx Chip, and sequenced using the Ion PGM™ Dx Sequencer.

The signal generated by the sequencing reaction is translated into base calls and then reads, which are mapped to a reference sequence. Using parameters in the specific Assay Definition File designed for a particular set of targets and therapies, Torrent Suite™ Dx Software generates reports containing a summary of the samples, test results, and any recommended therapies associated with the detected variants and gene fusions.

Sample and library preparation

The system has been validated with DNA and RNA isolated from FFPE tissue samples using the Ion Torrent Dx FFPE Sample Preparation Kit. Samples are prepared as slide-mounted 5-micron FFPE sections, which are deparaffinized before use. The samples must be macrodissected and enriched for tumor content if the tumor content is less than 20% and the tumor content in the region of interest is greater than or equal to 10%, or if the tissue is highly necrotic. The samples are digested, then the DNA and RNA are isolated and quantified. The minimum concentration and R² values that are required for library preparation are shown in Table 3.

Table 3 Required sample concentrations and R² values from the linear regression of the standards

Sample type	Required concentration	Required R ² value
DNA	≥0.83 ng/μL	≥0.99
RNA	≥1.43 ng/μL	≥0.98

The RNA is transcribed into cDNA using the Ion Torrent Dx cDNA Synthesis Kit, and sample and control amplicon libraries are prepared from the cDNA and DNA using primers and reagents in the Oncomine™ Dx Target Test, Controls, and Diluent Kit and Ion PGM™ Dx Library Kit. Libraries created using these kits have a distinguishing nucleic acid sequence barcode that is incorporated into each amplicon. Information about each sample and its resulting libraries are entered into Torrent Suite™ Dx Software, which tracks the progress of the sample from library preparation through analysis. The specific Assay Definition File for a particular set of targets and therapies defines the sample and library information required and tracked by the software.

Template preparation and sequencing

Using the Ion OneTouch™ Dx Instrument and the process of emulsion PCR, the library molecules are bound to Ion PGM™ Dx ISPs and each nucleic acid sequence is clonally amplified over the ISP surface. The templated ISPs are enriched and collected using the Ion OneTouch™ ES Dx Instrument. Sequencing primer is annealed to the single-stranded template, sequencing enzyme is added, and the ISPs are loaded onto the Ion 318™ Dx Chip. Chip loading occurs through use of the Ion PGM™ Dx Chip Minifuge. The chip is then placed onto the Ion PGM™ Dx Sequencer, where the DNA sequencing reaction occurs.

As the Ion PGM™ Dx Sequencer flows nucleotides over the chip surface, bases are incorporated into the strands on the bead in each well, resulting in the release of protons and a concomitant pH change in the well. The change in pH is detected by sensors at the base of each well on the chip. This initial electrical signal is processed for each well and transmitted to the Ion Torrent™ Server associated with the system.

Throughout this procedure, as the sample is prepared and processed by each instrument, sample and reagent information are recorded and tracked by Torrent Suite™ Dx Software.

Data analysis

On the Ion Torrent™ Server, the initial signals are processed, and bases are called. These calls are assembled into files representing the reads, which are strings of nucleotide bases in the order found in the original library molecules. The reads are then mapped to the reference files provided with the test. Finally, Torrent Suite™ Dx Software assesses the mapped reads at specific nucleotide locations and looks for variation from the sequence information in the human reference sequence.

Results

Using parameters in the specific Assay Definition File designed for a particular set of targets and therapies, Torrent Suite™ Dx Software generates the following electronic results and reports for each sequenced sample and its associated controls.

Table 4 Electronic results and reports generated by the software

Results/report	Description
View Result screen	Contains QC and reference information, detailed sequencing analytics, and all variant and gene fusion calls.
Test Report	A clinical report that lists the variants associated with the cancer type and detected in the sample that are screened by the Oncomine™ Dx Target Test, and any recommended therapies.
Laboratory Report	Contains all the information in the Test Report, as well as sequencing run details and QC evaluation metrics for the sample and controls.

These reports are subject to approval by a lab manager or administrator via electronic signature.

Assay warnings and limitations

- Use of this product must be limited to personnel trained in the techniques of PCR, NGS, and the use of the Oncomine™ Dx Target Test and the Ion PGM™ Dx System.
- The Oncomine™ Dx Target Test has only been validated for use with FFPE tumor slide specimens.
- The Oncomine™ Dx Target Test has been validated to detect the following somatic mutations: single-nucleotide variations (SNVs), multi-nucleotide variations (MNVs), and deletions of 3, 6, 9, 12, 15, and 18 base pairs, and insertions of 3, 6, 9, and 12 base pairs in DNA, and fusions in RNA.
- The Oncomine™ Dx Target Test is only validated for use with the Ion PGM™ Dx System and the Veriti™ Dx 96-well Thermal Cycler, 0.2 mL.
- The Oncomine™ Dx Target Test is only validated for use with 10 ng each of DNA and RNA per sample. Input amounts lower or higher than 10 ng are not recommended.
- Both the DNA and RNA from a single sample extraction must meet the concentration requirements specified in the procedure. Do not use DNA from one extraction with RNA from a different extraction.
- The effects of potential variations in FFPE specimen fixation have not been evaluated.
- Extraction from FFPE sample curls has not been evaluated.
- A potential source of contamination in the procedure is nucleic acid from previous sample processing steps. Follow good laboratory practices and all precautions and guidelines in these user guides to avoid cross-contamination between samples.
- The Oncomine™ Dx Target Test is a qualitative test. The test is not for quantitative measurements of percent mutation.

- For NSCLC, the Oncomine™ Dx Target Test assay definition file includes prevalent but not all rare or newly identified RET isoforms, ROS1 isoforms, and EGFR exon 20 insertions. The Oncomine™ Dx Target Test may miss rare or newly identified:
 - RET isoforms carried by a subset of patients who may derive benefit from pralsetinib
 - ROS1 isoforms carried by a subset of patients who may derive benefit from crizotinib
 - EGFR exon 20 insertions carried by a subset of patients who may derive benefit from mobocertinib
- The safe and effective use of the variants reported in Table 2 has not been established for selecting therapy using this device. The variants for KRAS (COSM512/p.Gly12Phe/c.34_35delGGinsTT and COSM516/p.Gly12Cys/c.34G>T), MET (COSM707/p.Thr1010Ile/c.3029C>T) and PIK3CA (COSM754/p.Asn345Lys/c.1035T>A) have been analytically validated. Performance of all other variants identified by the test, other than the clinically validated therapeutic variants and analytically validated variants, has not been directly demonstrated.

Software compatibility and requirements

The procedures in this guide are designed for use with Torrent Suite™ Dx Software version 5.12.5 or later. To view the current software version, sign in to the software as an Administrator, click the **Settings (⚙)** tab, select **Configuration**, then click the **Software Updates** tab. Version-specific information is provided in the software release notes for the version of the software you are using.

Torrent Suite™ Dx Software is supported on Google™ Chrome™ browser version 64 and later and is best viewed with 1440 × 900 screen resolution. It has not been tested with other browsers.

The Ion Torrent™ Server operating system is Ubuntu™ 18.04 LTS.

Materials provided

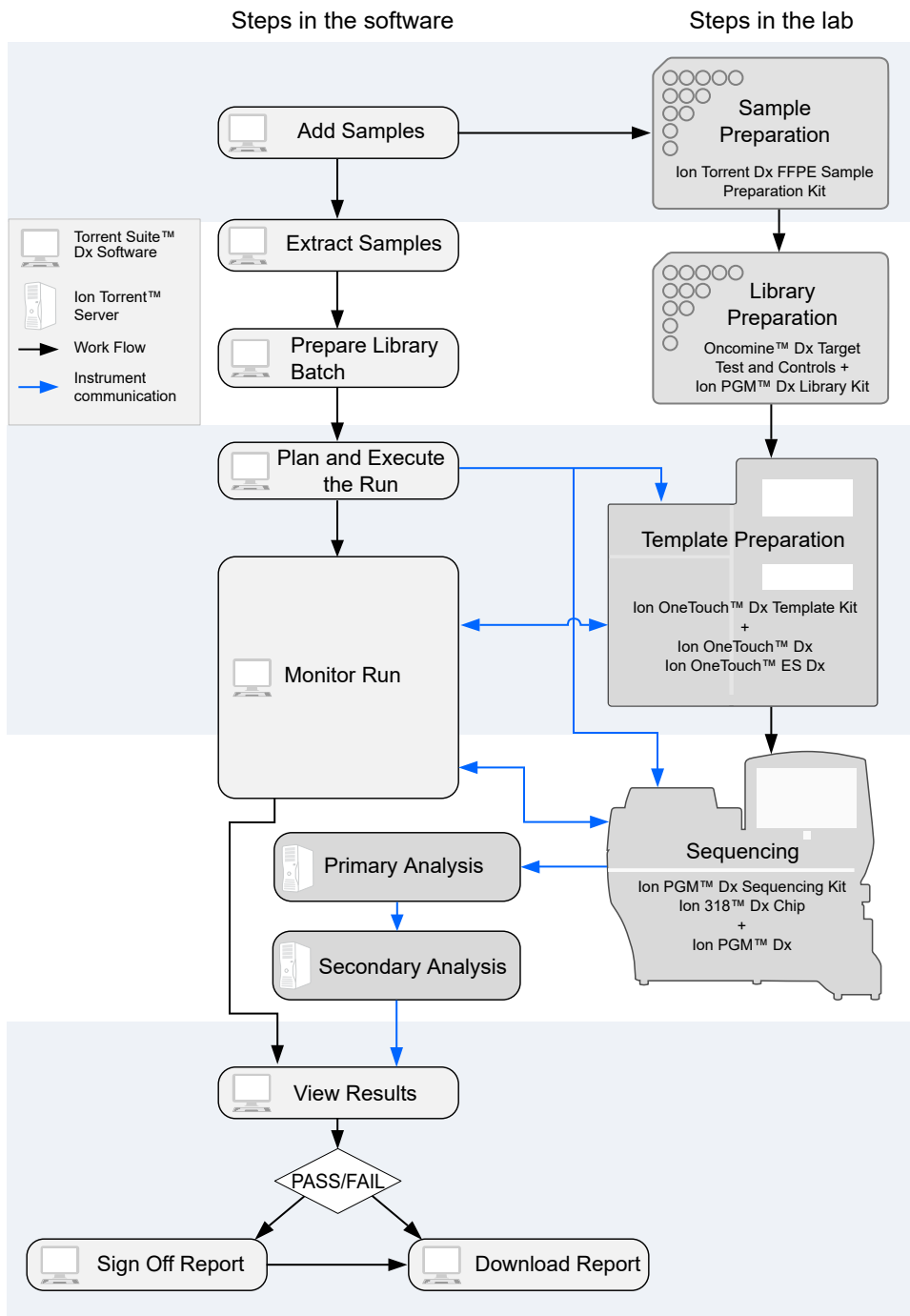
Oncomine™ Dx Target Test Kit

The Oncomine™ Dx Target Test Kit (Cat. No. A51695) includes the following subkits.

IMPORTANT! Refer to the product label for the storage conditions and expiration dates of individual modules and components.

✓	Subkit	Part No.
	Oncomine™ Dx Target Test, Controls, and Diluent Kit	A49756
	Ion Torrent Dx FFPE Sample Preparation Kit	A32445
	Ion PGM™ Dx Library Kit	A49758
	Ion OneTouch™ Dx Template Kit	A49759
	Ion PGM™ Dx Sequencing Kit	A49760
	Ion 318™ Dx Chip Kit	A18937
	Oncomine™ Dx Target Test User Guides and Assay Definition File	A51694

Oncomine™ Dx Target Test system diagram



2

Pass/fail criteria and repeat strategy

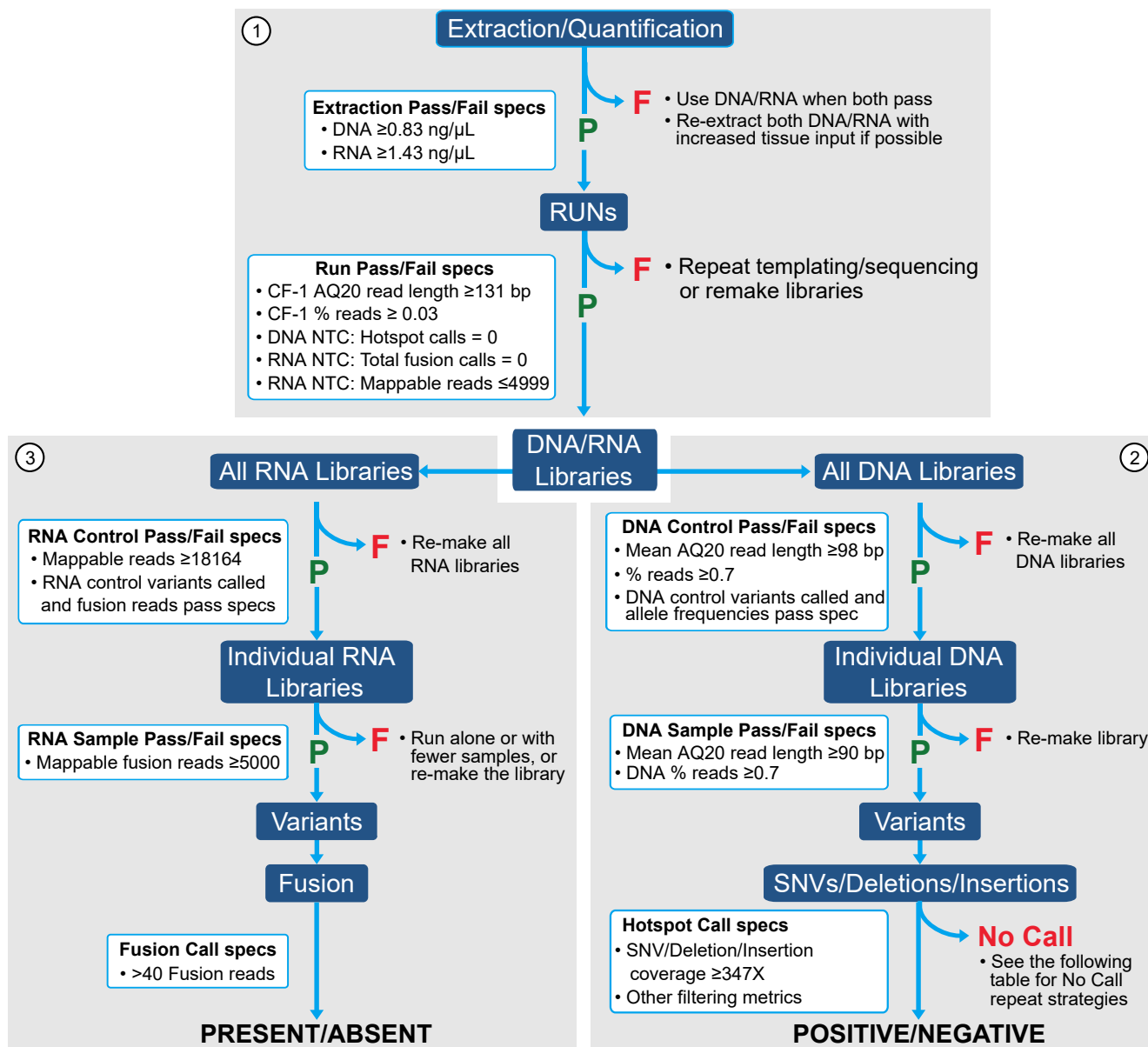
Quality control pass/fail criteria

Metric	Criteria
Run QC	
CF-1 Mean AQ20 Read Length (bp)	≥131
CF-1 Percent Reads (%)	≥0.03
DNA NTC	Hotspot calls = 0
RNA NTC	Total fusion calls = 0
RNA NTC Mappable Reads	≤4999
DNA Library	
Mean AQ20 Read Length (bp)	≥90
Percent Reads (%)	≥0.7
RNA Library	
Mappable Fusion Reads	≥5000
DNA Control	
COSM476_AF	Variant called and AF ≥0.05
COSM521_AF	Variant called and AF ≥0.05
COSM6223_AF	Variant called and AF ≥0.05
COSM6224_AF	Variant called and AF ≥0.05
COSM760_AF	Variant called and AF ≥0.05
COSM28747_AF	Variant called and AF ≥0.05
Mean AQ20 Read Length (bp)	≥98
Percent Reads (%)	≥0.7
RNA Control	
Mappable Reads	≥18164
ROS1 Fusion Reads	Variant called and fusion reads ≥349

Pass/fail specifications and repeat strategy

In general, if you experience a run or sample failure, you can repeat the run or sample preparation at the workflow step in which the failure occurred. The number of libraries that can be made from an extracted sample for repeat testing depends on the quantity of DNA and RNA from the extraction, which varies from sample to sample.

Based on quality control results, you can determine whether a library requires repeat testing. Refer to the following flowchart and table to determine the best course of action.



Point of failure		Tissue type	Recommended repeat strategy
1	RUNs	NSCLC and CC	Repeat templating and sequencing or remake the libraries.
2	DNA Library	NSCLC and CC	Remake all DNA libraries.
3	RNA library	NSCLC	Remake all RNA libraries.
		CC	Unnecessary to remake cholangiocarcinoma sample libraries, because RNA fusion variants are not reported for these samples.

Point of failure	Specification type	Passing criteria	Repeat strategy
1	Extraction/ Quantification	The following minimum concentrations for DNA and RNA are required: <ul style="list-style-type: none"> DNA ≥ 0.83 ng/μL. RNA ≥ 1.43 ng/μL. 	If the minimum concentration requirement is not met for either DNA or RNA, the samples must be re-extracted with increased tissue input. Use the set of extracted samples where both the DNA and RNA meet the minimum concentration requirement for the test.
1	Run	A run must pass the following specifications to have reportable results for any sample within the run: <ul style="list-style-type: none"> CF-1 Mean AQ20 Read Length (bp) must be ≥ 131. CF-1 Percent Reads must be ≥ 0.03. 	If either CF-1 specification fails, the operator may repeat the templating/sequencing run with the same library pool, or re-pool the libraries if a pooling error is suspected. If the issue persists on the repeat run, remake the libraries.
1	No Template Control (NTC)	A run must pass the following NTC specifications to have reportable results for any samples within the run: <ul style="list-style-type: none"> DNA No Template Control (DNA NTC) – Total "Hotspot Calls" must equal zero (0). RNA No Template Control (RNA NTC) – Mappable Reads must be ≤ 4999 and "Total Fusion Calls" must be zero (0). 	<p>If only the DNA NTC fails, remake all of the DNA controls and DNA sample libraries, and re-pool with previously made RNA controls and RNA sample libraries.</p> <p>If only the RNA NTC fails, remake all of the RNA controls and RNA sample libraries, and re-pool with previously made DNA controls and DNA sample libraries.</p> <p>If both the DNA NTC and RNA NTC fail, remake all of the DNA and RNA controls and sample libraries.</p>

(continued)

Point of failure	Specification type	Passing criteria	Repeat strategy
2	DNA Control	<p>The DNA control must pass the following specifications in order for any DNA samples within the run to have any reportable results:</p> <ul style="list-style-type: none"> • AQ20 Mean Read Length (bp) must be ≥ 98. • Percent Reads must be $\geq 0.7\%$. • All variants within the DNA control sample must be called "Present" and pass the allelic frequency range for each variant as specified in the assay definition file. 	If any of these specifications fail, the operator must remake all DNA control and DNA sample libraries.
3	RNA Control	<p>The RNA control must pass the following specifications for any RNA samples within the run to have any reportable results:</p> <ul style="list-style-type: none"> • Mappable Reads must meet the minimum threshold required of ≥ 18164 reads. • All variants within the RNA control sample must be called "Present" and pass the threshold metric set for total fusion reads required for each variant as specified in the assay definition file. 	If either of these specifications fails, the operator must remake all the RNA control and RNA sample libraries.
2	Library DNA Sample	<p>Any individual DNA sample library must meet the following specifications to have reportable results for the DNA sample library:</p> <ul style="list-style-type: none"> • Mean AQ20 Read Length (bp) ≥ 90. • Percent Reads ≥ 0.7. 	<p>Run the DNA sample library alone, or with fewer DNA sample libraries. If the DNA sample library still fails these specifications, remake the library using the same (previously extracted) DNA, if there is sufficient quantity. If not, re-extract using more tissue input, if possible. The repeat libraries must be prepared and run with new DNA controls.</p> <p>Note: Use the original passing RNA control and DNA and RNA sample libraries as placeholders when needed, and re-pool the libraries accordingly in the repeat runs. Any data resulting from the placeholder libraries must be ignored in the repeat run.</p>

(continued)

Point of failure	Specification type	Passing criteria	Repeat strategy
3	Library RNA Sample	Any individual RNA sample library must have ≥ 5000 Mappable Fusion Reads to have reportable results for the RNA sample library.	<p>Run the RNA sample library alone, or with fewer RNA sample libraries. If the RNA sample library still fails this specification, re-make the library using the same (previously extracted) RNA, if there is sufficient quantity. If not, re-extract using more tissue input, if possible. The repeat libraries must be prepared and run with new RNA controls.</p> <p>Note: Use the original passing DNA control and DNA and RNA sample libraries as placeholders when needed, and repool the libraries accordingly in the repeat runs. Any data resulting from the placeholder libraries must be ignored in the repeat run.</p>
2	SNV, Deletion, and Insertion Variant Specifications	All Single Nucleotide Variants (SNVs), Deletions (Dels), and Insertions (Ins) must have coverage ≥ 347 reads and pass all Variant Caller filtering metrics in order to have a reportable result for the variant.	Any SNVs, deletions, and insertions that do not meet the coverage criteria will result in a "No Call" for the variant. The operator may run the sample alone or with fewer samples to obtain reportable results for the variant. If the repeat run fails to meet the minimum coverage requirement, the operator may remake the library to obtain reportable results for the variant.

No Call repeat strategies

Point of failure	Observation (Example FR tag) ^[1]	Reason ^[2]	Repeat strategy
2	<i>MINCOV<347, PosCov<2, NegCov<2*</i>	Coverage	Repeat the run with fewer samples per chip may improve coverage.
2	<i>NODATA</i>	No data	Repeating the run with fewer samples per chip may improve coverage.
2	<i>QualityScore<8</i>	Quality score	Remake the DNA and/or RNA libraries starting from nucleic acid sample to improve the quality score if the quality score is due to low coverage.
2	<i>STDBIAS0.99034>0.96, STDBIASPVAL 0.299<=1</i>	Strand bias	Remake the DNA and/or RNA libraries starting from nucleic acid sample to improve strand coverage.

^[1] The reason for a No Call can be determined by examining the value of the FR tag for a given variant, listed in the output VCF file.

^[2] While presence of ALL of the reasons for No Call is not necessary for a No Call assignment, more than one condition may occur simultaneously. In cases where multiple reasons are observed, such that one recommends repeat while the other does not, repeating sample from library preparation is recommended.

Pass/fail specifications and repeat strategy – mixed runs

If you combine non-small cell lung cancer and cholangiocarcinoma samples in a single Oncomine™ Dx Target Test run and experience a run or sample failure, follow the repeat strategy recommendations according to the tissue type of the samples that require retesting that are listed in “Pass/fail specifications and repeat strategy” on page 14.

For example, if the RNA Control fails in a mixed run, it is not necessary to remake cholangiocarcinoma sample libraries, because RNA fusion variants are not reported for these samples.



Performance characteristics

Cholangiocarcinoma (CC)—Analytical studies

Tissue input study

Fifteen (15) slide-mounted FFPE samples were analyzed to determine if samples extracted using the Ion Torrent Dx Total Nucleic Acid Isolation Kit yield DNA and RNA at the concentrations that are required by the Oncomine™ Dx Target Test when tissue input requirements are met. The test requires DNA at a concentration of ≥ 0.83 ng/ μ L and RNA at a concentration of ≥ 1.43 ng/ μ L.

Five (5) resection samples with $\geq 20\%$ tumor content were prepared without macrodissection, 5 resection samples with $< 20\%$ to $\geq 10\%$ tumor cell content were macrodissected, and 5 samples were collected by core needle biopsy (CNB). For the resection samples with ≥ 100 mm² surface area, 1 x 5 μ m section was used per extraction. For resection samples with < 100 mm², 4 x 5 μ m sections were used per extraction. For CNBs, all of which had a surface area < 30 mm², 9 x 5 μ m sections were used per extraction. DNA and RNA concentrations were determined using the Ion Torrent Dx DNA and RNA Quantification Kits, respectively. No sequencing was performed on the extracted samples.

Of the 15 samples tested, 93.3% (14/15) had a DNA concentration of ≥ 0.83 ng/ μ L and an RNA concentration of ≥ 1.43 ng/ μ L. One CNB sample failed the minimum DNA and RNA concentration specifications, with values of 0.72 ng/ μ L and 0.81 ng/ μ L respectively. The low concentrations were likely caused by insufficient tissue input as the sample only had a single core biopsy per slide.

Guard band study

Guard band testing was performed to evaluate the tolerance levels of the Proteinase K digestion and inactivation steps during FFPE sample preparation when using the Oncomine™ Dx Target Test with CC samples.

The tolerance level for each test condition (volume, temperature, and time for digestion and inactivation) was evaluated by comparing DNA and RNA concentrations across 3 test levels: Low, Standard Operating Protocol (SOP)/Nominal, and High. For each test condition and level, DNA and RNA were extracted from 1 IDH1 variant-positive FFPE CC sample and an FFPE BRAF V600E cell-line (in triplicate) and sequenced using the Oncomine™ Dx Target Test.

No statistically significant difference was observed between the levels for all 3 test conditions for the samples tested.

FFPE block stability study

Stability of CC FFPE tissue blocks at room temperature was established when tested with the Oncomine™ Dx Target Test.

Three (3) IDH1 (2 R132G and 1 R132C) variant-positive clinical sample blocks were tested in duplicate at baseline, 3 months + 2 weeks, 6 months + 2 weeks, and 12 months + 2 weeks' time points.

Linear regression analyses and stability estimates from each sample across the timepoints demonstrated that CC FFPE tissue blocks were stable for up to 12 months.

FFPE slide stability study

Stability of paraffin dipped and un-dipped FFPE CC tissue sections mounted on slides was established when using the Oncomine™ Dx Target Test.

Three (3) IDH1 (2 R132C and 1 R132G) variant-positive clinical samples from dipped and un-dipped FFPE CC tissue sections were tested at baseline, 3 months + 1 week, 6 months + 1 week, 9 months + 1 week, and 12 months + 1 week time points.

Linear regression analyses and stability estimates from each sample across the timepoints demonstrated that CC FFPE tissue sections mounted on slides were stable for up to 12 months.

Sample processing reproducibility study

The reproducibility and repeatability of IDH1 R132 variant detection using the Oncomine™ Dx Target Test were evaluated with 2 IDH1 WT samples and 4 IDH1 R132 (2 R132C and 2 R132G) variant-positive samples at a single test site. The site had 2 Ion PGM™ Dx instrument systems and 2 operators.

Each sample was tested 6 times by each operator, for a total of 12 replicates per sample. After repeat testing, there was a single invalid reaction (1/72 or 1.39%).

The negative call rate, positive call rate, and within-run repeatability were calculated for each IDH1 R132 variant-positive sample at the expected IDH1 R132 variant location. The results are shown in Table 5.

Including no calls the negative call rate for the IDH1 WT sample was 100% at all IDH1 R132 variant locations. Including no calls the positive call rate from the expected IDH1 R132 positive variants was 100%.

Table 5 Reproducibility call rates

Sample	Variant Identification	Variant (amino acid change)	# of valid sample results (N)	# of positive calls (A)	# of negative calls (B)	# of no calls (C)	Positive call rate + 95% C.I.		Negative call rate + 95% C.I.		Within-run repeatability + 95% C.I.	
							Including no calls (A/N)	Excluding no calls (A/(A+B))	Including no calls (B/N)	Excluding no calls (B/(A+B))	Including no calls	Excluding no calls
A	COSM28747	R132C	11	11	0	0	100% (71.5%, 100%)	100% (71.5%, 100%)	0% (0%, 28.5%)	0% (0%, 28.5%)	100% (47.8%, 100%)	100% (47.8%, 100%)
B	COSM28749	R132G	12	12	0	0	100% (73.5%, 100%)	100% (73.5%, 100%)	0% (0%, 26.5%)	0% (0%, 26.5%)	100% (54.1%, 100%)	100% (54.1%, 100%)
D	COSM28747	R132C	12	12	0	0	100% (73.5%, 100%)	100% (73.5%, 100%)	0% (0%, 26.5%)	0% (0%, 26.5%)	100% (54.1%, 100%)	100% (54.1%, 100%)
E	COSM28749	R132G	12	12	0	0	100% (73.5%, 100%)	100% (73.5%, 100%)	0% (0%, 26.5%)	0% (0%, 26.5%)	100% (54.1%, 100%)	100% (54.1%, 100%)

Interfering substances study

Two (2) potentially interfering substances that may be found in cholangiocarcinoma (CC) FFPE tissue samples, hemoglobin and bile acids, were evaluated using the Oncomine™ Dx Target Test on the Ion PGM™ Dx System.

The guidelines for testing are defined in section 7.1 of CLSI EP07A2E, which describes testing substances at a relatively high concentration as an interference screen. One potentially interfering endogenous substance, hemoglobin, was tested at twice the concentration recommended in CLSI EP07A2E, Appendix D.

Table 6 Interfering substances and amounts

Potential interfering substance	Step	Amount of substance
Hemoglobin	After deparaffinization, hemoglobin was added to the Digestion Buffer used to pre-wet the tissue section.	4 mg/mL
Bile acids	After deparaffinization, bile acids were added to the Digestion Buffer used to pre-wet the tissue section.	30 nmol/mL

Three (3) IDH1 R132 (1 R132G and 2 R132C) variant-positive and 1 WT FFPE CC clinical samples (2 replicates each) were extracted in the presence and absence of the excess endogenous substance and processed through the entire assay workflow. The concordance between variant calls in samples with and without interfering substances was calculated for each substance under investigation.

With no calls excluded, the results of testing with hemoglobin and bile acids showed 100% concordance with the control condition for both the IDH1 R132 variant-positive and WT FFPE CC samples. This data supports the claim that hemoglobin, and bile acids do not impact Oncomine™ Dx Target Test assay performance at the level tested.

Limit of Detection (LoD) study

The LoD was evaluated for all 5 IDH1 R132 variants that are detected by the Oncomine™ Dx Target Test in clinical samples. The LoD is the lowest allelic frequency (AF) of the IDH1 R132 variants that can be detected at least 95% of the time. DNA from variant-containing samples or cell lines were blended with DNA from WT FFPE CC samples at multiple levels and used as input DNA for the test. A minimum of 120 data points was generated for each IDH1 R132 variant by testing 6 titration levels, 2 reagent lots, and 10 replicates (per level per lot).

The LoD of the 5 IDH1 R132 variants ranged from 4.5–5.7% AF.

Table 7 LoD of clinical IDH1 variants

Variant	ID	Sample type	Estimated LoD (AF)
R132C	COSM28747	Clinical sample	4.5%
R132G	COSM28749	Clinical sample	5.7%
R132H	COSM28746	Cell line	4.9%
R132L	COSM28750	Cell line	5.1%
R132S	COSM28748	Cell line	5.3%

Assay reproducibility study

The reproducibility and repeatability of IDH1 R132 variant detection using the Oncomine™ Dx Target Test were assessed with 1 IDH1 WT sample and 3 IDH1 R132 variant-positive samples at 2 allelic frequency (AF) levels. Testing was performed at 4 testing sites, each site had 2 Ion PGM™ Dx instrument systems, 2 operators, and completed testing using 4 lots of reagents.

Thirty-six (36) replicates per sample were tested across all sites. Overall, there were 72 sequencing events per variant and samples were run in duplicate for repeatability analysis. After repeat testing, there was a single invalid reaction (1/252 or 0.4%).

The negative call rate, positive call rate, and within-run repeatability were calculated for each IDH1 R132 variant-positive sample at the expected IDH1 R132 variant location. The results are shown in Table 8.

The overall positive call rate for the IDH1 R132 variants was 92.6% when including no calls and 97.1% when excluding no calls. The negative call rate for the IDH1 WT sample was 100% at all IDH1 R132 variant locations.

Table 8 Reproducibility results

Sample	Variant Identification	Variant (amino acid change)	# of valid sample results (N)	# of positive calls (A)	# of negative calls (B)	# of no calls (C)	Positive call rate + 95% CI		Relative LoD
							Including no calls (A/N)	Excluding no calls (A/(A+B))	
D1	COSM28747	R132C	36	36	0	0	100% (90.3%, 100%)	100% (90.3%, 100%)	2.1–2.7X
D2	COSM28747	R132C	36	35	0	1	97.2% (85.5%, 99.9%)	100% (90.0%, 100%)	0.98–1.4X
D3	COSM28749	R132G	36	36	0	0	100% (90.3%, 100%)	100% (90.3%, 100%)	1.9–2.5X
D4	COSM28749	R132G	36	36	0	0	100% (90.3%, 100%)	100% (90.3%, 100%)	0.9–1.3X
D5	COSM28750	R132L	36	36	0	0	100% (90.3%, 100%)	100% (90.3%, 100%)	1.4–1.8X
D6	COSM28750	R132L	35	20	6	9 ^[1]	57.1% (39.4%, 73.7%)	76.9% (56.4%, 91.0%)	0.65–0.94X
D7	Wild-type (WT)	N/A	36	0	0	0	0% (0%, 9.17%)	0% (90.3%, 100%)	N/A

^[1] A number of no calls were seen because the LoD for this variant is close to the assay AF cutoff of 2.5%.

Cholangiocarcinoma (CC)—Clinical studies

IDH1 clinical study

IDH1 study—concordance evaluation

A total of 383 samples were obtained for this study. Both slides cut from FFPE blocks and extracted DNA were used. Of these, 187 were identified by the enrolling clinical trial assay (CTA) as IDH1 variant-positive, 187 were identified as IDH1 variant-negative, and 9 samples were invalid based on the enrolling CTA.

The 187 IDH1 variant-positive samples set were previously enrolled into the AG-120-C-005 clinical study. The IDH1 variant-negative samples were randomly selected from the CTA patient population that had yielded a valid negative result on the CTA assay. The IDH1 invalid samples were randomly selected from the CTA patient population that yielded invalid results.

Of the 187 IDH1 variant-positive samples from the CTA testing, 174 were positive on Oncomine™ Dx Target Test. Of the 174 samples IDH1 positive by Oncomine™ Dx Target Test, 172 were IDH1 positive from the NDA population as 2 samples tested by the CTA were not part of the efficacy population. Of the 187 IDH1 CTA positive samples in the bridging population, 1 sample was called negative, 6 samples were invalid, 6 samples were cancelled due to failure to meet test input requirements due to low or insufficient sample availability.

Of the 187 IDH1 variant-negative samples tested, 0 were identified as IDH1 variant-positive by the Oncomine™ Dx Target Test, 14 were invalid, 5 yielded no calls, and 2 were not tested due to insufficient sample availability, leaving 166 confirmed IDH1 variant-negative samples by the Oncomine™ Dx Target Test.

Of the 9 CTA invalid samples tested, 3 yielded Oncomine™ Dx Target Test positive results, 3 yielded negative results, 2 were confirmed invalid, and 1 was not tested due to insufficient sample availability.

In summary, 9 samples were cancelled (not tested), 22 had invalid Oncomine™ Dx Target Test results, 5 samples were no calls, 177 were IDH1 variant-positive, and 170 were IDH1 variant-negative by the Oncomine™ Dx Target Test.

The PPA was defined as the proportion of IDH1 variant-positive specimens as called by the CTA assay that were also IDH1 variant-positive by the Oncomine™ Dx Target Test. The NPA was defined as the proportion of IDH1 variant-negative specimens as called by the CTA assay that were also IDH1 variant-negative by the Oncomine™ Dx Target Test. The unadjusted concordances by variant and overall concordance (OPA) are shown in Table 9.

Table 9 IDH1 — Bridging concordance results (unadjusted)

Parameter	Agreed	Total	Agreement	Exact 95% CIs
PPA exclude UNK	174	175	99.4%	96.9%, 100.0%
NPA exclude UNK	166	166	100.0%	97.8%, 100.0%
OPA exclude UNK	340	341	99.7%	98.4%, 100.0%
PPA include UNK	174	181	96.1%	92.2%, 98.4%

Table 9 IDH1 — Bridging concordance results (unadjusted) (continued)

Parameter	Agreed	Total	Agreement	Exact 95% CIs
NPA include UNK	166	185	89.7%	84.4%, 93.7%
OPA include UNK	340	366	92.9%	89.8%, 95.3%

Of the enrolled samples, 349 samples were analyzed using the Oncomine™ Dx Target Test to demonstrate positive percent agreement (PPA) and negative percent agreement (NPA) concordance with a validated reference detection method (Sanger assay).

One hundred and sixty-eight (168) specimens from patients that tested positive using the Sanger assay were analyzed using the Oncomine™ Dx Target Test. In addition, 181 specimens that tested negative using the Sanger assay were analyzed using the Oncomine™ Dx Target Test.

Of the IDH1 variant-positive samples, 164 generated valid results from the Oncomine™ Dx Target Test. Three samples had invalid results due to failed QC metrics for the sequencing runs, and one generated a no call due to insufficient coverage.

Of the IDH1 variant-negative samples, 170 generated valid results from the Oncomine™ Dx Target Test. Ten samples had invalid results due to failed QC metrics for the sequencing runs, and one sample generated a no call due to insufficient coverage.

The PPA was defined as the proportion of IDH1 variant-positive specimens as called by the Sanger assay that were also IDH1 variant-positive as called by the Oncomine™ Dx Target Test, and the NPA was defined as the proportion of IDH1 variant-negative specimens as called by the Sanger assay that were also IDH1 variant-negative as called by the Oncomine™ Dx Target Test. The concordances and overall concordance (OPA; overall percent agreement) are shown in Table 10.

Table 10 IDH1 — Accuracy concordance results

Agreement measure	Excluding invalids and no calls		Including invalids and no calls	
	Percent agreement	95% CI	Percent agreement	95% CI
PPA	99.4% (163/164)	(96.7%, 100.0%)	97.0% (163/168)	(93.2%, 99.0%)
NPA	96.5% (164/170)	(92.5%, 98.7%)	90.6% (164/181)	(85.4%, 94.4%)
OPA	97.9% (327/334) ^[1]	(95.7%, 99.2%)	93.7% (327/349)	(90.6%, 96.0%)

^[1] Seven samples were found to be discordant in this analysis, where one was called a false negative and six were called false positives with the Oncomine™ Dx Target Test.

IDH1 study—clinical effectiveness

The clinical effectiveness of the Oncomine™ Dx Target Test was evaluated by measuring progression-free survival (PFS) for patients with CC who tested positive for IDH1 R132 variants (R132C, R132G, R132H, R132L, and R132S) by both the Clinical Trial Assay (CTA) and the Oncomine™ Dx Target Test. Progression-free survival and hazard ratio were calculated for patients who were selected for treatment with ivosidenib.

The primary efficacy outcome measurements between treatment and control arms (including PFS, hazard ratio, and overall survival) were examined in the AG120-C-005 study, based on the evaluable Oncomine™ Dx Target Test results (confirmed positive by the Oncomine™ Dx Target Test; 172 total

patients) and the study population (185 total patients) that included samples not confirmed with the Oncomine™ Dx Target Test.

The clinical efficacy (represented by PFS) determined in the Oncomine™ Dx Target Test positive population (N=115 treatment vs. 57 placebo) showed a HR=0.37 with 95% CI of (0.25, 0.55), and is similar to the Oncomine™ Dx Target Test positive *plus unevaluable* population (N=123 treatment vs. 61 placebo; HR=0.38; 95% CI: 0.26, 0.55) and the overall CTA+ population (primary endpoint of the AG120-C-005 study) (N=124 treatment vs. 61 placebo; HR = 0.37; 95% CI: 0.25, 0.54). These results suggest that no efficacy bias was introduced into the Oncomine™ Dx Target Test positive population.

Refer to the [Drugs@FDA database](#) for the most recent therapeutic product labeling.

Non-small cell lung cancer (NSCLC)—Analytical studies

Limit of blank studies

Limit of Blank (LoB)—Study I

To ensure that a variant-free ("blank") sample does not generate an analytical signal that might be classified as a mutation, wild-type (WT) samples were evaluated at each variant location that can be detected by the Oncomine™ Dx Target Test. Samples that are WT at all locations should produce a "variant not detected" call at each location. By definition (EP17-A2), the 95th percentile of test results on blank samples equals zero.

Operators extracted and sequenced nucleic acid from 5 WT cell lines prepared as FFPE sections on slides. The cell lines have well-characterized genomes and contain no known cancer biomarker sequences.

The study was conducted using two different lots of the Oncomine™ Dx Target Test Kit. For each lot, each cell-line sample was extracted once and made into 6 DNA and 6 RNA libraries. Operators sequenced each library in duplicate, generating 24 different sets of results across the two reagent lots per sample.

For all 5 samples, there were no positive calls at any of the variant locations analyzed by the test. The false positive rate was therefore zero.

Additionally, operators extracted and sequenced nucleic acid from 3 FFPE clinical samples prepared on slides. Each sample was tested using 24 replicates and 2 reagent lots of the Oncomine™ Dx Target Test Kit, resulting in 144 sequencing replicates each for DNA and RNA. For all replicates, there were no positive calls at any of the variant locations. The false positive rate was therefore zero, and the LoB of the test was determined to be zero.

Finally, nucleic acid from three WT FFPE clinical samples was processed from sample preparation through sequencing using the updated Oncomine™ Dx Target Test Kit RNA workflow. The study was conducted using 2 different lot combinations of kits. Each clinical sample generated 48 different replicates across the 2 reagent lot combinations, for a total of 144 replicates for the entire study. The false positive rate of the test was determined to be zero for the ROS1 fusion target, demonstrating that the LoB of the test was zero when tested with clinical samples.

Limit of Blank (LoB)—Study II

A second study was performed to test the frequency of false positive calls for EGFR exon 20 insertion variants detected by the Oncomine™ Dx Target Test in wild-type samples. In this study, nucleic acid was extracted from 4 wild-type (WT) FFPE NSCLC clinical samples and carried through from library preparation to sequencing. Samples that are WT at all EGFR locations should produce a "variant not detected" call at each location. By definition (EP17-A2), the 95th percentile of test results on blank samples equals zero.

Operators extracted and sequenced nucleic acid from the 4 WT samples prepared as FFPE sections on slides. The study was conducted using two different lots of the Oncomine™ Dx Target Test Kit. Operators made 18 library replicates for each sample and kit lot, giving a total of 4 samples × 18 replicates × 2 lots = 144 libraries sequenced.

The result at each variant location for each sample was tabulated. For all 4 samples, there were no positive calls at any of the EGFR exon 20 insertion variant locations analyzed by the test. The false positive rate was therefore zero.

Tissue input study

Sixty slide-mounted FFPE samples were analyzed to determine if samples extracted using the Ion Torrent Dx Total Nucleic Acid Isolation Kit yield DNA and RNA at the concentrations required by the Oncomine™ Dx Target Test when tissue input requirements are met. The test requires DNA at a concentration of ≥ 0.83 ng/ μ L and RNA at a concentration of ≥ 1.43 ng/ μ L.

Thirty resection samples with $\geq 20\%$ tumor content were prepared without macrodissection, 15 resection samples with $< 20\%$ to $\geq 10\%$ tumor cell content were macrodissected, and 15 samples were collected by core needle biopsy (CNB). For the resection samples, 2×5 μ m sections were used per extraction. For CNBs, 9×5 μ m sections were used per extraction. DNA and RNA concentrations were determined using the Ion Torrent Dx DNA and RNA Quantification Kits, respectively. No sequencing was performed on the extracted samples.

Of the 60 samples tested, 98.3% (59/60) had a DNA concentration of ≥ 0.83 ng/ μ L and an RNA concentration of ≥ 1.43 ng/ μ L. One CNB sample failed the minimum DNA and RNA concentration specifications, with values of 0.52 ng/ μ L and 1.23 ng/ μ L respectively. The low concentrations were likely caused by the small tissue size and low tumor content (5%).

Seven slide-mounted FFPE fine needle aspirate (FNA) samples were analyzed to determine if samples extracted using the Ion Torrent Dx Total Nucleic Acid Isolation Kit yield DNA and RNA at the concentrations required by the Oncomine™ Dx Target Test when tissue input requirements are met.

For FNAs, 7×5 μ m sections were used per extraction. DNA and RNA concentrations were determined using the Ion Torrent Dx DNA and RNA Quantification Kits, respectively. 100% of the 7 FNA samples extracted using the Ion Torrent Dx FFPE Sample Preparation Kit yielded DNA at a concentration of ≥ 0.83 ng/ μ L and RNA at a concentration of ≥ 1.43 ng/ μ L.

Guard band study

Guard band testing was performed to define Oncomine™ Dx Target Test tolerances by testing critical assay steps at key points of the NSCLC workflow, from library preparation through sequencing, using a blend of variants with EGFR exon 20 insertions.

Tolerances were tested across 11 test conditions across 3 test levels: Low, Standard Operating Protocol (SOP)/Nominal, and High. A DNA blend of nucleic acids isolated from clinical FFPE samples with the EFGR insertion was used in 10 test conditions related to volumes, temperature, and time, while a DNA control was used in a test condition related to DNA control volume. For a test condition to be considered acceptable, 6 library replicates must pass sample validity metrics (Sample AQ20 mean read length and % reads), and the EGFR variant within the clinical blended sample must be called present using the Oncomine™ Dx Target Test. Following initial testing, narrower acceptable ranges were established for EtOH (1–1.75 μ L) and thermal cycling temperature offset (-1.0°C to 0°C).

The study demonstrated acceptable performance of the test at tolerance ranges that are significant deviations from the SOP-defined condition.

Stability of assay intermediates study

The stability of assay intermediates study was performed to test whether hold times in stopping points specified in the Oncomine™ Dx Target Test user guides affects test performance. The study was conducted in two separate parts that had the same study design and acceptance criteria. DNA corresponding to one of two EGFR exon 20 insertion variants, COSM1238030 (3 bp insertion), and COSM26720 (12 bp insertion), at mean allelic frequency (AF) of 2.5X LoD (1.9X to 3.5X LoD, 10% to 18% AF, respectively) was used in each part of the study to test assay performance at baseline, then compare it to performance after the stopping point holds that are specified in the Oncomine™ Dx Target Test user guides are included in the workflow. The Oncomine™ Dx Target Test workflow allows a total of 9 stopping points, each with a maximum hold time. The no-hold/hold performance was tested in three conditions:

- Nominal (no-hold)
- Library Hold—30-day hold of eluted libraries at –30°C to –10°C
- Combo Hold—8 remaining stopping points in library preparation, template preparation, and sequencing, tested at the hold time specified in the user guide

The study demonstrated that the assay intermediates are stable after pre-defined hold times and Oncomine™ Dx Target Test performance was not affected by the hold times. Both EGFR exon 20 insertion variants (3 and 12 bp) were called correctly 100% of the time in the clinical sample blends across both hold conditions and the control condition. A T-Test was also performed to compare the mean AFs observed across the sample blends tested in each test hold condition compared with the no-hold condition. The p value for each test was >0.05, and together the p values showed no statistically significant differences in the mean variant AF between the hold and the no-hold conditions.

DNA and RNA input studies

DNA and RNA input—Study I

Eight cell-line samples were prepared as FFPE sections, and DNA and RNA were extracted and quantified from multiple sections from each cell line for blending and testing. Sample blends were prepared with known variants at various DNA and RNA input-level combinations within the range of 5–15 ng. The DNA and RNA blends had a target allele frequency of 15% for SNVs and deletions and target fusion reads of 300–600 for the ROS1 variant. A total of 540 individual DNA and RNA libraries were tested, including positive controls and NTC controls, with 6 replicate libraries each for DNA and RNA per test condition.

The study demonstrated a 100% positive variant call rate within the input range tested, supporting the specified input amount of 10 ng each for DNA and RNA for the Oncomine™ Dx Target Test.

The negative variant call rate was >95% for all except 4 sample and DNA/RNA input-level combinations. All cases with a negative variant call rate of <95% were due to no calls, 3 of which occurred with a DNA or RNA input amount of 5 ng and 1 of which occurred in a single sample with DNA and RNA inputs of 10 ng each. There were no false-positive calls.

Additionally, 4 clinical samples prepared as FFPE sections were tested: two samples containing DNA variants and two containing the CD74-ROS1 fusion.

The DNA variant samples were paired with wild-type RNA from the same sample at various input combinations within the range of 5–15 ng, and the RNA variant samples were paired with wild-type DNA at input combinations within the same range.

The study demonstrated positive and negative call rates of >95% for the DNA variants at all input combinations, and 100% for one of the CD74-ROS1 fusions at all input combinations. The second CD74-ROS1 clinical sample showed 100% negative call rates for all test conditions, and 100% positive call rates except for Test Condition 4 (8.5 ng RNA/15 ng DNA), where the call rate was 83%, and Test Condition 6 (15 ng RNA/15 ng DNA), where the call rate was 50%. The false negatives for these test conditions were possibly due to operator error during library preparation, since the remaining replicates in these test conditions had both high total mappable reads and fusion reads, but the cause was not definitively determined.

The results support the DNA and RNA 10-ng input requirement for the Oncomine™ Dx Target Test.

DNA and RNA input—Study II

Two EGFR exon 20 insertion-positive variant DNA blends at allele frequency levels 2–3X the limit of detection were prepared for testing. A total of 96 libraries at various DNA input-level combinations within the range of 5–15 ng were tested, including positive controls and NTC controls, with 6 replicate libraries each test condition.

The study demonstrated a 100% positive variant call rate within the input range tested, supporting the specified input amount of 10 ng each for DNA for the Oncomine™ Dx Target Test.

The results support the DNA and RNA 10-ng input requirement for the Oncomine™ Dx Target Test.

In silico specificity study

An *in silico* cross-reactivity analysis was performed that evaluated the 832 primers in the Oncomine™ Dx Target Test Kit DNA and RNA panels to determine the specificity of the primers to their targeted sequences. The primers were checked for specificity to the human genome, the human transcriptome, and genomes from representative bacteria, fungi, and viruses frequently found in human tissue and lung specimens.

Any unintended amplification products were required to have ≥ 2 base-pair (bp) mismatches to intended amplification product sequences generated by the panels, because mismatches of ≥ 2 bp prevent mapping to the same location on the genome due to a low mapping score.

For the DNA panel primers, *in silico* analysis predicted 20 unintended potential amplicon-generating primer pairings against the human genome. Nineteen of these had unintended amplification products with ≥ 2 bp mismatches, and therefore would have low mapping scores and not cause false results. One unintended primer pairing was predicted to amplify regions identical to an intended product, and therefore would detect the same WT and variant locations and not cause false results.

For the RNA panel primers, analysis predicted 63 unintended primer pairings against the human genome and 7 unintended primer pairings against the human transcriptome. All of these predicted amplicons had mismatches of ≥ 42 bp to intended amplicons, and therefore would not cause false results.

Analysis of representative bacterial, fungi, and viral genomes resulted in one predicted unintended primer pairing with a mismatch of ≥ 61 bp to intended amplicons, which would not cause false results.

Based on these results, the primers in the Oncomine™ Dx Target Test Kit DNA and RNA panels were deemed specific.

Cross-contamination study

A total of 8 FFPE cell line samples were evaluated to determine the percentage of false positive results caused by cross-contamination (contamination from one sample to another within the same sequencing run) and carryover contamination (contamination from a previous run on the same instrument system). Samples that were WT and variant were tested in consecutive runs on the same instruments, and 5 DNA variant locations and 2 RNA variant locations that were expected to be WT for a sample were evaluated for contamination.

Out of 100 DNA and 80 RNA data points analyzed, no false positive results were reported in the DNA variants, and 1 false positive result was reported in a ROS1 fusion. The false positive was likely caused by sample cross-contamination from an adjacent well. Therefore, the false-positive rate at DNA variant locations was 0% (0/100) and the false-positive rate at RNA variant locations was 1.25% (1/80).

Tissue fixation study

A study was performed to evaluate the effect of 10% neutral buffered formalin (NBF) fixation times on cytosine deamination events at the hotspot locations targeted by the Oncomine™ Dx Target Test, and any effect these potential events would have on assay performance. Pellets from the wild-type cell line GM24385 were fixed with 10% NBF for 12, 24, 48, 72, and 84 hours. Sections from each block were cut, mounted on slides, and tested with the Oncomine™ Dx Target Test. These results were compared to results from cell line GM24385 that had not undergone any fixation with 10% NBF.

The average allelic frequency (AF) observed at each of the 103 cytosine deamination-susceptible hotspots was determined for each fixation time tested. The results showed 2 G>A deamination events as a result of the fixation process, one at a 24-hour fixation time for COSM232755 (AF 0.050%) and the other at a 48-hour fixation time for COSM181063 (AF 0.073%). Each resulted in a "no call". DNA and RNA sequencing quality was evaluated by measuring percent reads, no calls, and total mappable reads for each condition tested. Both DNA and RNA demonstrated valid sequencing results with all NBF fixation times tested in this study. Therefore, it was determined that NBF fixation times did not cause deamination events that negatively impacted sequencing results.

Sample processing reproducibility studies

Sample processing reproducibility—Study I

The reproducibility and repeatability of variant detection using the Oncomine™ Dx Target Test were assessed with 2 WT samples and 10 variant-positive samples at 4 testing sites. Each site had 4 Ion PGM™ Dx instrument systems and 4 operators.

Each sample was tested 8 times at each site, for a total of 32 replicates per sample. After repeat testing, the final number of invalid reactions was 15/768 (1.95%), possibly due to low sample quality or lack of sample, though the cause was not definitively determined.

The call rate, no call rate, positive call rate, negative call rate, and within-run repeatability were computed at each variant location of interest. Including no calls and excluding known positive variant locations, the negative call rate at each clinical variant location for all samples was 100%.

The results at positive variant locations are shown in Table 11. Including no calls, all positive call rates from positive variant locations were >84%.

Excluding no calls and combining data across all study samples, the estimate of repeatability was 100% for DNA variants and 87.5% for the RNA variant. The lower limit of the 95% two-sided confidence interval (CI) for repeatability exceeded 96% at all variant locations.

Including no calls from the data, the estimate of repeatability was 100% at 218 out of 605 variant locations, 94–99.9% at 175 out of 605 variant locations, and 71.6–93.9% at 212 out of 605 variant locations. Including no calls, the lower limit of the 95% two-sided confidence interval for repeatability exceeded 64.6% at all variant locations.

Table 11 Call rates at positive variant locations

Sample	Variant identification	Variant location	# of valid sample results (N)	# of positive calls (A)	# of negative calls (B)	# of No Calls (C)	Positive call rate + 95% CI		Negative call rate + 95% CI		Within-run repeatability + 95% CI	
							Including no calls (A/N)	Excluding no calls (A/(A+B))	Including no calls (B/N)	Excluding no calls (B/(A+B))	Including no calls	Excluding no calls
B	COSM6223	EGFR Exon19 del	32	32	0	0	100% (89.1%, 100%)	100% (89.1%, 100%)	0% (0%, 10.9%)	0% (0%, 10.9%)	100% (79.4%, 100%)	100% (79.4%, 100%)
B	COSM763	PIK3CA E545K	32	32	0	0	100% (89.1%, 100%)	100% (89.1%, 100%)	0% (0%, 10.9%)	0% (0%, 10.9%)	100% (79.4%, 100%)	100% (79.4%, 100%)
C	ROS1	N/A	32	30	2	0	93.8% (79.2%, 99.2%)	93.8% (79.2%, 99.2%)	6.3% (0.8%, 20.8%)	6.3% (0.8%, 20.8%)	87.5% (61.7%, 98.4%)	87.5% (61.7%, 98.4%)
D	COSM6225	EGFR Exon19 del	32	32	0	0	100% (89.1%, 100%)	100% (89.1%, 100%)	0% (0%, 10.9%)	0% (0%, 10.9%)	100% (79.4%, 100%)	100% (79.4%, 100%)
E	COSM476	BRAF V600E	32	32	0	0	100% (89.1%, 100%)	100% (89.1%, 100%)	0% (0%, 10.9%)	0% (0%, 10.9%)	100% (79.4%, 100%)	100% (79.4%, 100%)
F	COSM521	KRAS G12D	32	30	0	2	93.8% (79.2%, 99.2%)	100% (88.4%, 100%)	0% (0%, 10.9%)	0% (0%, 11.6%)	87.5% (61.7%, 98.4%)	100% (76.8%, 100%)
F	COSM29313	PIK3CA M1043I	32	30	0	2	93.8% (79.2%, 99.2%)	100% (88.4%, 100%)	0% (0%, 10.9%)	0% (0%, 11.6%)	87.5% (61.7%, 98.4%)	100% (76.8%, 100%)
G	COSM6224	EGFR L858R	32	32	0	0	100% (89.1%, 100%)	100% (89.1%, 100%)	0% (0%, 10.9%)	0% (0%, 10.9%)	100% (79.4%, 100%)	100% (79.4%, 100%)

Table 11 Call rates at positive variant locations (continued)

Sample	Variant identification	Variant location	# of valid sample results (N)	# of positive calls (A)	# of negative calls (B)	# of No Calls (C)	Positive call rate + 95% CI		Negative call rate + 95% CI		Within-run repeatability + 95% CI	
							Including no calls (A/N)	Excluding no calls (A/(A+B))	Including no calls (B/N)	Excluding no calls (B/(A+B))	Including no calls	Excluding no calls
J	COSM87298	KRAS Q61K	32	32	0	0	100% (89.1%, 100%)	100% (89.1%, 100%)	0% (0%, 10.9%)	0% (0%, 10.9%)	100% (79.4%, 100%)	100% (79.4%, 100%)
J	COSM17242 3	ERBB3 V104M	32	32	0	0	100% (89.1%, 100%)	100% (89.1%, 100%)	0% (0%, 10.9%)	0% (0%, 10.9%)	100% (79.4%, 100%)	100% (79.4%, 100%)
K	COSM775	PIK3 H1047R	30 ^[1]	29	0	1	96.7% (82.8%, 99.9%)	100% (88.1%, 100%)	0% (0%, 11.6%)	0% (0%, 11.9%)	93.3% (68.1%, 99.8%)	100% (76.8%, 100%)
M	COSM715	FGR3 S249C	32	32	0	0	100% (89.1%, 100%)	100% (89.1%, 100%)	0% (0%, 10.9%)	0% (0%, 10.9%)	100% (79.4%, 100%)	100% (79.4%, 100%)

^[1] Two replicates for sample K were initially invalid and not retested.

Sample processing reproducibility—Study II

The reproducibility and repeatability of variant detection using the Oncomine™ Dx Target Test were assessed with 2 WT samples and 2 EGFR exon 20 insertion variant-positive samples at 3 testing sites. Each sample was extracted two times and tested using 3 lots of reagents at each site, for a total of 18 replicates per sample. Each site had 2 Ion PGM™ Dx instrument systems and 2 operators.

The call rate, no call rate, positive call rate, negative call rate, and within-run repeatability were computed for each EGFR exon 20 insertion variant. Including no calls and excluding known positive variant locations, the negative call rate at each clinical variant location for all samples was 100%.

The results at positive variant locations are shown in Table 12. Including no calls, all positive call rates from positive variant locations were 100%.

Excluding no calls and combining data across all study samples, the estimate of repeatability was 100% for both EGFR exon 20 insertion variants. The lower limit of the 95% two-sided confidence interval (CI) for repeatability exceeded 100% at all variant locations.

Including no calls from the data, the estimate of repeatability was 100% at all EGFR exon 20 insertion variant locations. Including no calls, the lower limit of the 95% two-sided confidence interval for repeatability exceeded 81.5% at all variant locations.

Table 12 Call rates at positive variant locations

Sample	Variant identification	# of valid sample results (N)	# of positive calls (A)	# of negative calls (B)	# of no calls (C)	Positive call rate + 95% C.I.		Negative call rate + 95% C.I.		Within-run repeatability + 95% C.I.	
						Including no calls (A/N)	Excluding no calls (A/(A+B))	Including no calls (B/N)	Excluding no calls (B/(A+B))	Including no calls	Excluding no calls
D1	COSM12376	18	18	0	0	100% (81.5%, 100%)	100% (81.5%, 100%)	0% (0%, 18.5%)	0% (0%, 18.5%)	100% (66.4%, 100%)	100% (66.4%, 100%)
D2	COSM12380	18	18	0	0	100% (81.5%, 100%)	100% (81.5%, 100%)	0% (0%, 18.5%)	0% (0%, 18.5%)	100% (66.4%, 100%)	100% (66.4%, 100%)

Interfering substances studies

Interfering substances—Study I

Six potentially interfering substances that can be found in FFPE tissue samples or carried over from the deparaffinization and nucleic acid extraction process were evaluated using the Oncomine™ Dx Target Test on the Ion PGM™ Dx System.

The guidelines used at the time of testing are defined in section 7.1 of CLSI EP07A2E, which describes testing substances at a relatively high concentration as an interference screen. One potentially interfering endogenous substance, hemoglobin, was tested at twice the concentration recommended in CLSI EP07A2E, Appendix D.

In addition to the substances tested in this study, the data from the Tumor Necrosis Study demonstrated that the performance of the Oncomine™ Dx Target Test was not impacted by the presence of up to 75% necrotic tissue. However, users should macrodissect highly necrotic areas or select alternate samples if possible.

Table 13 Interfering substances and amounts

Potential interfering substance	Step	Amount of substance
Paraffin	At the deparaffinization step, extra paraffin was added to the xylene bath that contained 250 mL of xylene.	4X of normally expected levels
Xylene	Extra xylene was added into the ethanol bath that contained 250 mL of ethanol.	6X of normally expected residual volume
Ethanol	Extra ethanol was added into the Protease digestion step before digestion.	>4X of normally expected residual volume
Hemoglobin	After deparaffinization, hemoglobin was added to the Digestion Buffer used to pre-wet the tissue section.	4 mg/mL
Protease	Extra Protease was added into the reaction after the digestion step and before column purification.	>10X of expected residual Protease after the heat-kill step
Wash buffer	Wash buffer used to isolate DNA and RNA from deparaffinized and digested samples was added into an aliquot of Dilution Solution, which was later used to dilute the RNA and DNA to the appropriate concentration before library preparation.	1% wash buffer (equivalent to ~10% wash buffer carried over into eluate)
Control	Tissue sections were processed using the standard protocol, without the addition of any potentially interfering substances.	N/A

A total of 6 FFPE samples (1 WT and 5 variants) with 6 replicates each were processed through the entire assay workflow. The variant samples included variants from all variant categories that can be detected by the test. The samples were spiked with additional concentrations or amounts of the listed substances at the relevant processing step, as shown in the table. Replicates of a control sample with no spiked substances were also analyzed. The concordance between variant calls in samples with and without interfering substances was computed for each substance under investigation.

With no calls excluded, for each potential interferent used in sample extraction, the positive and negative concordance with the control condition across all samples was 100%, and the overall concordance with the control condition across all samples was 100%.

With no calls excluded, the results of testing with hemoglobin showed positive concordance with the control condition of 100% (only samples with a positive control condition were analyzed), negative concordance of 99.99%, and overall concordance of 99.99%.

Interfering substances—Study II

The interfering substances study was repeated with the updated RNA workflow to demonstrate that the performance of the Oncomine™ Dx Target Test is not affected by the presence of potentially interfering substances.

The impact on assay performance of the listed interferents (Table 13) was evaluated in this study and the results were compared to the control (no interferents) condition. For the 6 interferents tested, both the positive concordance (no calls excluded) and the overall concordance (no calls excluded) for all samples was 100%. These data support the claim that paraffin, xylene, ethanol, hemoglobin, protease, or wash buffer do not affect assay performance at the level tested.

Interfering substances—Study III

A study was performed to demonstrate that the performance of the Oncomine™ Dx Target Test in detecting RET fusions is not affected by the presence of potentially interfering substances.

A total of 3 FFPE samples (2 RET fusion-positive, and 1 WT) with 3 replicates each were used to evaluate the impact of the listed interferents (Table 13) on assay performance, and the results were compared to the control (no interferents) condition. For the 6 interferents tested, both the positive concordance and the overall concordance for all samples was 100%. These data support the claim that paraffin, xylene, ethanol, hemoglobin, protease, or wash buffer do not affect assay performance at the level tested in detection of the RET fusions.

Interfering substances—Study IV

A study was performed to demonstrate that the performance of the Oncomine™ Dx Target Test in detecting EGFR exon 20 insertion-positive variants is not affected by the presence of potentially interfering substances.

A total of 3 FFPE samples (2 EGFR exon 20 insertion-positive, and 1 WT) with 2 replicates each were used to evaluate the impact of the listed interferents (Table 13) on assay performance, and the results were compared to the control (no interferents) condition. For the 6 interferents tested, both the positive concordance and the overall concordance for all samples was 100%. These data support the claim that paraffin, xylene, ethanol, hemoglobin, protease, or wash buffer do not affect assay performance at the level tested in detection of EGFR exon 20 insertion-positive variants.

Limit of Detection (LoD) studies

Limit of Detection (LoD)—Study I

The LoD was evaluated for 14 representative DNA variants detected by the Oncomine™ Dx Target Test in clinical samples. The LoD is the lowest AF of SNV, MNV, or deletion variants that can be detected at least 95% of the time. Variant-containing samples were blended with WT samples at multiple levels and used as the input DNA for the test.

Due to the large number of variants detected by the Oncomine™ Dx Target Test and the rarity of some of the variants, the LoD was established using a representative variant approach. Variants were selected in the following categories:

- Simple SNVs
- Complex SNVs and MNVs (SNVs in di- or tri-nucleotide repeat regions, SNVs in high-GC (>60%) or low-GC (<40%) content regions, and MNVs)
- Deletions (including deletions of 6, 9, 15, and 18 bp)

Clinical specimens were tested for all variants for which clinical claims are being sought. Seven variants for which analytical claims are being sought were unavailable in clinical specimens, and so plasmid constructs were substituted.

A minimum of 120 data points were generated for each representative variant by testing 6 or more titration levels, 2 reagent lots, and 10 replicates (per level per lot). The claimed LoD for all but 1 variant is the maximum of the LoD obtained from testing each of the 2 lots in this study.

Based on 14 representative DNA variants in 6 genes assessed in clinical samples, the LoDs for DNA variants tested in clinical samples (supported by the results from the assay reproducibility study) were determined to have allelic frequencies (AFs) of 6–8%.

Limit of Detection (LoD)—Study II

The LoD of the Oncomine™ Dx Target Test was calculated by testing 2 clinical ROS1 fusion-positive specimens using the updated Oncomine™ Dx Target Test RNA workflow.

RNA extracted from fusion-positive clinical FFPE specimens was blended with RNA extracted from wild-type (WT) clinical FFPE specimens to achieve 6 dilution levels. For each ROS1 fusion isoform, 10 replicates per 6 dilution levels were tested with 2 reagent lots, giving a total of 120 data points. The LoD of the assay for ROS1 fusion detection was determined to be 516 fusion reads (higher of the LoD observed for the 2 isoforms tested).

Limit of Detection (LoD)—Study III

The LoD of the Oncomine™ Dx Target Test was calculated by testing 2 clinical RET fusion-positive specimens using the updated Oncomine™ Dx Target Test RNA workflow.

RNA extracted from fusion-positive clinical FFPE specimens was blended with RNA extracted from wild-type (WT) clinical FFPE specimens to achieve 6 dilution levels. For each RET fusion isoform, 10 replicates per dilution level were tested with 2 reagent lots, giving a total of 120 data points. The LoD of the assay for RET fusion detection was determined to be 405 fusion reads (higher of the LoD observed for the 2 isoforms tested).

Limit of Detection (LoD)—Study IV

The LoD of the Oncomine™ Dx Target Test was calculated by testing 2 clinical EGFR exon 20 insertion-positive specimens using the Oncomine™ Dx Target Test DNA workflow.

DNA extracted from EGFR exon 20 insertion-positive clinical FFPE specimens was blended with DNA extracted from WT clinical FFPE specimens to achieve 6 dilution levels. For each EGFR exon 20 insertion variant, 10 replicates per dilution level were tested with 2 reagent lots, giving a total of 120 data points. The LoD of the assay for EGFR exon 20 insertion detection was determined to be 4.8–5.2% allelic frequency.

Tumor content studies

Tumor content—Study I

To determine the minimum tumor cell content required in FFPE samples used as input material, 55 pre-characterized clinical samples were analyzed using the Oncomine™ Dx Target Test. They contained SNVs, deletions, and fusions confirmed by validated reference methods. The tumor cell content of each specimen and region of interest was estimated before the study by an external pathology lab.

The samples were analyzed with and without macrodissection. Fifty-four samples contained DNA variants and 1 contained an RNA variant. Three samples contained 2 SNV or deletion variants, for a total of 58 variants analyzed. The observed tumor content had the following distribution:

- 10 samples with tumor content <10%
- 16 samples with tumor content 10–19%
- 13 samples with tumor content 20–29%
- 9 samples with tumor content 30–39%
- 3 samples with tumor content 40–49%
- 4 samples with tumor content 50–60%

In the samples without macrodissection, all 58 variants were detected (called positive) by the Oncomine™ Dx Target Test. In the macrodissected samples, there was one "no call" in a BRAF V600E variant sample with a tumor content of 16%.

Additional studies were performed to validate that the minimum tumor cell content required in FFPE input samples is greater than or equal to 20% when using the updated RNA library preparation workflow. As part of the ROS1 study, 9 ROS1-positive samples were identified by both methods. The observed tumor content in the studies ranged from 20–90%, and had the following distribution:

Table 14 Tumor content range in ROS1-positive samples used in clinical studies

ROS1-positive samples	Tumor-content range
1	20–29%
1	30–39%
0	40–49%
7	50–90%

There was no statistically significant association between tumor content and test results. The results confirm that the minimum tumor cell content in FFPE samples used as input material for the Oncomine™ Dx Target Test is greater than or equal to 20%.

Tumor content—Study II

The minimum tumor cell content that is required in FFPE samples used as input material was calculated for 71 pre-characterized FFPE clinical samples. The samples were analyzed using the updated Oncomine™ Dx Target Test RNA workflow. The tumor cell content of each specimen and region of interest was estimated before the study by an external pathology lab. The samples were analyzed with and without macrodissection.

A total of 9 ROS1 positive and 62 ROS1 negative samples were included in the study analysis. All samples gave valid results for both the Oncomine™ Dx Target Test (passing Run, RNA Control, and RNA Sample QC criteria) and the reference method test. These valid samples were used for the tumor content study analysis. The observed tumor content had the following distribution:

- 0 samples with tumor content <20%
- 18 samples with tumor content 20–29%
- 8 samples with tumor content 30–39%
- 10 samples with tumor content 40–49%
- 14 samples with tumor content 50–69%
- 21 samples with tumor content 70–90%

Table 15 Tumor content range in ROS1-positive samples used in clinical studies

ROS1-positive samples	Tumor-content range
0	<20%
1	20–29%
1	30–39%
0	40–49%
4	50–69%
3	70–90%

The corresponding 95% Clopper Pearson Exact CIs of the PPA, NPA, and OPA overlapped between tumor content levels. This shows that the ROS1 fusion detection performance of the Oncomine™ Dx Target Test was similar at all ranges of tumor content level. The tumor content level of the clinical samples had no impact on the performance of the Oncomine™ Dx Target Test.

Tumor content—Study III

In a further study, the tumor cell content in FFPE samples used as input material was calculated for 149 FFPE clinical samples. The tumor cell content of each specimen and region of interest was estimated before the study by an external pathology lab. The samples were analyzed with and without macrodissection.

A total of 54 EGFR exon 20 insertion-positive and 95 EGFR exon 20 insertion-negative samples were included in the study analysis. All samples gave valid results for both the Oncomine™ Dx Target Test (passing Run, DNA Control, and DNA Sample QC criteria) and the reference method test. These valid samples were used for the tumor content study analysis. The observed tumor content had the following distribution:

- 21 samples with tumor content <20%
- 41 samples with tumor content 20–29%
- 26 samples with tumor content 30–39%
- 27 samples with tumor content 40–59%
- 34 samples with tumor content 60–100%

Table 16 Tumor content range in EGFR exon 20 insertion-positive samples used in clinical studies

EGFR exon 20 insertion-positive samples	Tumor-content range
10	<20%
20	20–29%
6	30–39%
9	40–59%
9	60–100%

The corresponding 95% Clopper Pearson Exact CIs of the PPA, NPA, and OPA overlapped between tumor content levels. This shows that the EGFR exon 20 insertion detection performance of the Oncomine™ Dx Target Test was equivalent at all ranges of tumor content level. The tumor content level of the clinical samples had no impact on the performance of the Oncomine™ Dx Target Test.

Assay reproducibility studies

Assay reproducibility—Study I

The reproducibility and repeatability of the Oncomine™ Dx Target Test was evaluated for 30 representative variants from 18 DNA samples.

The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instrument platforms (reproducibility). Six of the 18 DNA samples were mixtures of plasmid and clinical DNA. Seven of the 12 deletion variants were represented by these plasmid blends. All other variant types were represented by clinical sample DNA.

Due to the large number of variants detected by the test and the rarity of some of the variants, a representative variant approach was used. Variants were selected in the following categories:

Table 17 Representative variant approach—Study I

Variant category	No. of plasmid blends used	No. of clinical specimens used
6-bp deletion	6	0
9-bp deletion	4	2
15-bp deletion	2	4
18-bp deletion	2	4
Simple SNV	0	8
Complex SNVs ^[1] and MNVs	0	6

^[1] Including SNVs in di- or tri-nucleotide repeat regions and SNVs in high-GC (>60%) or low-GC (<40%) content regions

Two of the 18 DNA samples were WT at all locations, and the remaining 16 contained DNA from one or more DNA variants. Each pre-extracted DNA sample was sequenced at 4 sites by 4 operators on 2 systems at each site.

At each site, operators were grouped into 2 pairs, with each pair assigned to 2 instrument systems and responsible for testing 9 DNA samples. Samples were run in duplicate using 2 different reagent lots at 3 of the study sites and on all 3 reagent lots at one study site. The design resulted in a total of 72 test determinations per DNA sample, and all variant locations were assessed for each sample.

The reproducibility results are summarized in the following table.

Table 18 Reproducibility results—Study I

Description	Variants evaluated across the samples	Call rate excluding no calls ^[1]		Call rate including no calls ^[1]	
		Mean	Median	Mean	Median
Variant positive DNA (positive calls)	46	96.6%	97.1%	94.5%	95.8%
WT DNA (negative calls)	872	96.1%	95.0%	96.1%	95.0%

^[1] Analysis includes invalid results.

Excluding no calls, the estimate of repeatability at each DNA variant location across all the samples was $\geq 98.8\%$ (95% CI lower limit of $\geq 97.5\%$). The coefficient of variation (CV) across all DNA clinical variants ranged from 9.8% to 39%. The highest CVs (24.9–39.2%) were observed for the BRAF V600E variant. The higher percent CV for this sample was possibly due to poor sample quality, but the cause was not definitively determined. The CVs for the EGFR L858R variant ranged from 9.8% to 11.3%, and the CVs for the EGFR deletion variants ranged from 11.2% to 25.5%.

Assay reproducibility—Study II

An additional study was performed to evaluate the reproducibility and repeatability of the OncoPrint™ Dx Target Test for 6 representative variants from 11 DNA samples and 4 RNA samples. All 11 DNA samples and 4 RNA samples were clinical sample blends. In addition, 1 WT DNA sample and 4 WT RNA samples were included in the study.

The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instrument platforms (reproducibility). The updated OncoPrint™ Dx Target Test RNA workflow was used.

Due to the large number of variants detected by the test and the rarity of some variants, a representative variant approach was used. Variants were selected in the following categories:

Table 19 Representative variant approach—Study II

Variant category	No. of clinical specimens used	No. of levels tested ^[1]	No. of samples tested
15-bp deletion	1	2	7 ^[2]
Simple SNV	3	2	
Complex SNV ^[3] and MNV	2	2	4
Fusion	2	2	4
WT RNA	2	N/A	4 ^[4]
WT DNA	1	N/A	1

^[1] The number of sample blends with different levels of allele frequency that were tested.

^[2] Includes one sample that contained both a 15-bp deletion and a simple SNV, one sample that contained a 15-bp deletion, and three samples that were simple SNVs.

^[3] Including SNVs in di- or tri-nucleotide repeat regions and SNVs in high-GC (>60%) or low-GC (<40%) content regions

^[4] Each WT RNA sample was tested twice.

One DNA sample was WT at all locations, 1 DNA sample had 2 DNA variants, and the remaining DNA samples had 1 variant each. Four RNA samples were WT for the ROS1 fusion. Each pre-extracted DNA or RNA sample was sequenced at 3 sites by 2 operators on 2 systems at each site.

At each site, 2 operators were assigned to 2 instrument systems and were responsible for testing 12 DNA samples (11 with variants and 1 WT) and 8 RNA samples (4 with variants and 4 WT). Samples were run in duplicate using 3 different reagent lots at all study sites. The study design resulted in a total of 36 test determinations per DNA or RNA sample.

The reproducibility results are summarized in the following table.

Table 20 Reproducibility results (DNA variants)—Study II

Description	Variants evaluated across the samples	Call rate excluding no calls ^[1]		Call rate including no calls ^[1]	
		Mean	Median	Mean	Median
Variant positive DNA (positive calls)	12	99%	100%	98%	99%
WT DNA (negative calls)	367	100%	100%	99%	100%

^[1] Analysis includes invalid results.

Excluding no calls, the estimate of repeatability at each DNA variant location across all the samples was $\geq 94.4\%$ (95% CI lower limit of $\geq 72.7\%$). The CVs across all DNA clinical variants ranged from 9.1% to 22.6%. The CVs for the BRAF V600E variant ranged from 13.1% to 19%, the CVs for the EGFR L858R variant ranged from 11% to 17.6%, and the CVs for the EGFR deletion variants ranged from 10.1% to 15.9%.

Table 21 Reproducibility results (ROS1 fusion)—Study II

Description	Isoforms evaluated across the samples	Call rate excluding or including unknowns ^[1]	
		Mean	Median
ROS1 fusion-positive RNA (positive calls)	4	100%	100%
WT RNA (negative calls)	4	99%	100%

^[1] Unknowns are defined as invalid or no result using the Oncomine™ Dx Target Test.

The estimate of repeatability at each RNA clinical variant location was 100%. One wild-type sample was found to be contaminated with RNA control during the study, resulting in an NPA estimate of 94.4%. No specific sequencer performed differently between three lots of Oncomine™ Dx Target Test reagents for this sample. The CV across both RNA locations ranged from 47.8% to 76.6%.

Assay reproducibility—Study III

A study was performed to evaluate the reproducibility and repeatability of the Oncomine™ Dx Target Test for detection of RET fusions using FFPE RNA from 4 RET fusion-positive samples (blended with WT clinical samples) and 2 RET fusion-negative (WT) samples.

Table 22 Sample description—Study III

RET fusion	No. of clinical specimens used	No. of levels tested	No. of samples tested
KIF5B-RET.K15R12	1	2	2
CCDC6-RET.C1R12	1	2	2
RET-negative/WT RNA	2	N/A	2

The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instrument platforms (reproducibility). The updated RNA library preparation workflow was used.

Six pre-extracted RNA sample blends (4 fusion positive blends and 2 WT specimens) and placeholder DNA libraries were sequenced at 3 sites by 6 operators. At each site, 2 operators were assigned to 2 instrument systems and were responsible for testing the 6 RNA sample blends. Samples were run in duplicate using 3 different reagent lots at all study sites. The study design resulted in a total of 36 test determinations per sample blend tested.

The reproducibility results are summarized in the following table.

Table 23 Reproducibility results (RET fusion)—Study III

Description	Isoforms evaluated across the samples	Call rate excluding or including unknowns ^[1]	
		Mean	Median
RET fusion-positive RNA (positive calls)	4	99%	100%
WT RNA (negative calls)	2	100%	100%

^[1] Unknowns are defined as invalid or no result using the OncoPrint™ Dx Target Test.

Estimates of the repeatability were reported to be 100% for the CCDC6-RET.C1R12 isoform, and 98.1% for the KIF5B-RET.K15R12 isoform. The CV across both RNA locations ranged from 46.8% to 62.7%.

Assay reproducibility—Study IV

A study was performed to evaluate the reproducibility and repeatability of the OncoPrint™ Dx Target Test for detection of EGFR exon 20 insertion variants using FFPE DNA from 2 EGFR variant-positive samples (blended with WT clinical samples) and 2 EGFR variant-negative (WT) samples.

Table 24 Sample description—Study IV

EGFR exon 20 insertion	No. of clinical specimens used	No. of levels tested	No. of samples tested
COSM1238028	1	2	2
COSM12376	1	2	2
EGFR-negative/WT DNA	2	N/A	2

The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instrument platforms (reproducibility).

Six pre-extracted DNA sample blends (4 insertion positive blends and 2 WT specimens) and placeholder RNA libraries were sequenced at 3 sites by 6 operators. At each site, 2 operators were assigned to 2 instrument systems and were responsible for testing the 6 DNA sample blends. Samples were run in duplicate using 2 different reagent lots at all study sites. The study design resulted in a total of 24 test determinations per sample blend tested.

The reproducibility results are summarized in the following table.

Table 25 Reproducibility results (EGFR exon 20 insertions)—Study IV

Description	Variants evaluated across the samples	Call rate excluding no calls ^[1]		Call rate including no calls ^[1]	
		Mean	Median	Mean	Median
EGFR exon 20 insertion-positive DNA (positive calls)	2	100%	100%	100%	100%
WT DNA (negative calls)	2	100%	100%	100%	100%

^[1] Analysis includes invalid results.

Estimates of the repeatability were reported to be 100% for both COSM1238028 and COSM12376. The CV across both DNA locations ranged from 11.3% to 16.7%.

Panel accuracy study

To evaluate the ability of the Oncomine™ Dx Target Test DNA and RNA panels to identify somatic variants in human specimens, 290 FFPE tumor samples were analyzed using the Oncomine™ Dx Target Test to demonstrate positive percent agreement (PPA) and negative percent agreement (NPA) concordance with validated reference detection methods.

The following reference detection methods were used:

- A validated NGS assay, to detect SNV and deletion hotspot variants
- A ROS1 FISH reference test, to detect ROS1 fusions

Variants detected by the Oncomine™ Dx Target Test that were not covered by the reference methods were not included in the PPA/NPA concordance calculation. Variants detected by the Oncomine™ test for which the reference method testing failed and did not yield a valid result were not included in the PPA/NPA calculation.

Accuracy data was analyzed by the following:

- Each variant location
- Bins (or categories) of variants: RNA fusions, simple SNVs, complex SNVs, and deletions
- Each FFPE sample

The results are shown in the following tables.

Table 26 PPA results

PPA measure	Excluding no calls		Including no calls	
	Percent agreement	95% CI	Percent agreement	95% CI
Variant	98.5% (195/198)	(95.6%, 99.7%)	98.5% (195/198)	(95.6%, 99.7%)
Bin	97.2% (176/181)	(93.7%, 99.1%)	97.2% (176/181)	(93.7%, 99.1%)
Sample	96.9% (158/163)	(93.0%, 99.0%)	96.9% (158/163)	(93.0%, 99.0%)

Table 27 NPA results

NPA measure	Excluding no calls		Including no calls	
	Percent agreement	95% CI	Percent agreement	95% CI
Variant	100.0% (118,155/118,159)	(99.99%,100.0%)	96.8% (118,155/122,012)	(96.7%, 96.9%)
Bin	99.8% (942/944)	(99.2%, 100.0%)	70.0% (657/939)	(66.9%, 72.9%)
Sample	98.4% (124/126)	(94.4%, 99.8%)	23.4% (29/124)	(16.3%, 31.8%)

Table 28 OPA results

OPA measure	Excluding no calls		Including no calls	
	Percent agreement	95% CI	Percent agreement	95% CI
Variant	100.0% (118,350/118,357)	(99.99%, 100.0%)	96.8% (118,350/122,210)	(96.74%, 96.94%)
Bin	99.4% (1,118/1,125)	(98.72%,99.75%)	74.4% (833/1,120)	(71.71%, 76.91%)
Sample	97.6% (282/289)	(95.07%,99.02%)	65.2% (187/287)	(59.34%, 70.66%)

Non-small cell lung cancer (NSCLC)—Clinical studies

BRAF clinical study

BRAF study—concordance evaluation

A method comparison evaluated the accuracy of the Oncomine™ Dx Target Test compared to the NSCLC BRAF V600E PCR Assay for the detection of the BRAF V600E mutation in NSCLC samples. Patient samples from the NSCLC BRF113928 clinical trial and an acquired set of negative samples were measured by both assays.

There were a total of 230 samples available for analysis. Of these, 181 samples (67 + 114) provided valid results for both the BRAF V600E PCR assay and the Oncomine™ test. All valid results correlated. Of the remaining samples, 27 samples had invalid results with the Oncomine™ test due to failed control or library QC metrics for the sequencing runs, 9 samples had no calls due to insufficient coverage at the BRAF variant location, and 13 samples were not tested due to insufficient DNA concentration.

The positive percent agreement (PPA) was defined as the proportion of BRAF-positive samples called by the BRAF V600E PCR Assay that were also called by the Oncomine™ Dx Target Test, and the negative percent agreement (NPA) was defined as the proportion of BRAF-negative samples called by the PCR assay that were also identified by the Oncomine™ Dx Target Test. The 95% two-sided exact CIs were determined for PPA, NPA and overall percent agreement (OPA), and the results are shown in the following table:

Table 29 Concordance between the NSCLC BRAF V600E PCR Assay and the Oncomine™ Dx Target Test

Agreement measure	Excluding no calls		Including no calls	
	Percent agreement	95% CI ^[1]	Percent agreement	95% CI ^[1]
PPA	100.0% (67/67)	(94.6%, 100.0%)	91.8%(67/73)	(83.0%, 96.9%)
NPA	100% (114/114)	(96.7%, 100.0%)	97.4%(114/117)	(92.7%, 99.5%)
OPA	100.0% (181/181)	(97.9%, 100.0%)	95.3%(181/190)	(91.2%, 97.8%)

^[1] The 95% CI was calculated using the Pearson-Clopper Exact method.

BRAF study—clinical effectiveness

The clinical effectiveness of the Oncomine™ Dx Target Test was evaluated by measuring the objective response rate (ORR) for patients with stage IV NSCLC who tested positive for the BRAF V600E mutation by both the Local Laboratory Tests (LLTs) and the Oncomine™ test. The ORR was calculated for patients in two cohorts (B and C) who were selected for treatment with TAFINLAR® (dabrafenib) administered in combination with MEKINIST® (trametinib).

The ORR for Cohort B was 68.2% (15/22), which is similar to the 63.2% ORR (36/57) observed in the overall population tested as positive by LLTs. The ORR for Cohort C was 60.9% (14/23), which is similar to the 61.1% ORR (22/36) observed in the overall population tested as positive by LLT.

A secondary objective of the bridging study was to determine the clinical effectiveness of the Oncomine™ Dx Target Test in selecting NSCLC patients for treatment with dabrafenib administered as a single agent and in combination with trametinib by evaluating progression-free survival (PFS), duration of response (DoR), and overall survival (OS) by both investigator assessment and independent review.

For the 15 Cohort B patients with a confirmed tumor response based on independent assessment, the median DoR was not estimable, with an event rate less than 50%. The median DoR for the overall LLT(+) population was 12.6 months. PFS was similar between the Oncomine™ Dx Target Test(+)/LLT(+) population (N = 22) and the total LLT(+) population (N = 57) for both independent and investigator review. Also, the ORR observed by independent assessment was similar to that observed by investigator assessment. The median follow-up time for Cohort B was 16.6 months.

For the 14 Cohort C patients with a confirmed tumor response based on independent assessment, the median DoR was not estimable with an event rate less than 50%. The median DoR for the overall LLT(+) was also not estimable with an event rate less than 50%. PFS was similar between the Oncomine™ Dx Target Test(+)/LLT(+) population (N = 23) and the total LLT(+) population (N = 36) for both independent and investigator review. Also, the ORR observed by independent assessment was similar to that observed by investigator assessment. The median follow-up time for Cohort C was 10.4 months.

Refer to the [Drugs@FDA database](#) for the most recent therapeutic product labeling.

EGFR L858R and exon 19 deletions clinical study

To evaluate the ability of the Oncomine™ Dx Target Test to identify the EGFR biomarker in FFPE NSCLC tumor specimens, 92 specimens from patients that tested positive using the QIAGEN™ *therascreen* EGFR RGQ PCR Kit were analyzed using the Oncomine™ Dx Target Test. In addition, 142 specimens that tested negative using the Qiagen EGFR PCR assay were analyzed using the Oncomine™ Dx Target Test.

Of the EGFR-positive samples, 72 generated valid results from both the Qiagen EGFR PCR assay and the Oncomine™ Dx Target Test. Twenty samples had invalid results due to failed control or library QC metrics for the sequencing runs, or generated no calls due to insufficient coverage.

Of the EGFR-negative samples, 121 generated valid results from both the Qiagen assay and the Oncomine™ test, while 12 had invalid results due to failed QC metrics for the sequencing runs or generated no calls due to insufficient coverage.

In all, 193 samples were used to evaluate concordance between the Oncomine™ test as an investigational method and the Qiagen EGFR PCR assay as the reference method. A total of 70 samples were excluded, and 32 samples were invalid or generated no calls.

The PPA was defined as the proportion of EGFR-positive specimens as called by the EGFR PCR assay that were also EGFR-positive as called by the Oncomine™ Dx Target Test, and the NPA was defined as the proportion of EGFR-negative specimens as called by the EGFR PCR assay that were also

EGFR-negative as called by the Oncomine™ test. The concordances by variant and overall concordance are shown in the following tables:

Table 30 Exon 19 deletion—Concordance

Agreement measure	Excluding no calls		Including no calls	
	Percent agreement	95% CI	Percent agreement	95% CI
PPA	97.6% (41/42)	(87.43%, 99.94%)	74.6% (41/55)	(61.00%, 85.33%)
NPA	99.3% (147/148)	(96.29%, 99.98%)	94.2% (147/156)	(89.33%, 97.33%)
OPA	99.0% (188/190) ^[1]	(96.25%, 99.87%)	89.1% (188/211)	(84.09%, 92.96%)

^[1] Two samples were found to be discordant in this analysis, where one was called a false negative and the other a false positive with the Oncomine™ test.

Table 31 EGFR L858R—Concordance

Agreement measure	Excluding no calls		Including no calls	
	Percent agreement	95% CI	Percent agreement	95% CI
PPA	100% (30/30)	(88.43%, 100%)	93.8% (30/32)	(79.19%, 99.23%)
NPA	100% (167/167)	(97.82%, 100%)	93.3% (167/179)	(88.58%, 96.49%)
OPA	100% (197/197)	(98.14%, 100%)	93.4% (197/211)	(89.12%, 96.33%)

Table 32 Overall concordance

Agreement measure	Excluding no calls		Including no calls	
	Percent agreement	95% CI	Percent agreement	95% CI
PPA	98.6% (71/72)	(92.5%, 100.0%)	81.6% (71/87)	(71.86% , 89.11%)
NPA	99.2% (120/121)	(95.5%, 100.0%)	96.8% (120/124)	(91.95% , 99.11%)
OPA	99.0% (191/193)	(96.31%, 99.87%)	90.5% (191/211)	(85.74% , 94.11%)

EGFR exon 20 insertions clinical study

EGFR exon 20 insertions concordance evaluation—Study II

To evaluate the ability of the Oncomine™ Dx Target Test to identify the EGFR exon 20 insertion biomarker in FFPE NSCLC tumor specimens, 87 specimens from patients that tested positive using local laboratory tests (LLT) were analyzed using the Oncomine™ Dx Target Test and a validated reference NGS assay (henceforth referred to as the reference assay). In addition, 110 samples screened as EGFR exon 20 insertion-negative were analyzed with the Oncomine™ Dx Target Test and the reference assay.

Of the 87 EGFR exon 20 insertion variant-positive samples, 63 were positive by the Oncomine™ Dx Target Test, 12 samples were negative, 6 samples yielded an invalid result, and 6 samples generated no calls. For the reference assay, 55 out of the 87 samples were positive, 13 samples were negative, and 19 samples yielded an invalid result.

Of the 110 CTA-screened expected negative samples, 89 yielded negative calls with the Oncomine™ Dx Target Test, 19 samples yielded an invalid result, and 2 samples generated no calls. For the reference assay, 92 out of the 110 samples were negative, and 18 yielded invalid results.

In all, 160 samples were used to evaluate concordance between the Oncomine™ Dx Target Test as an investigational method and the reference assay. A total of 37 samples were excluded due insufficient material and invalid results between the two tests.

The PPA was defined as the proportion of EGFR-positive specimens as called by the reference assay that were also EGFR-positive as called by the Oncomine™ Dx Target Test, and the NPA was defined as the proportion of EGFR-negative specimens as called by the reference assay that were also EGFR-negative as called by the Oncomine™ Dx Target Test. The concordances by variant and overall concordance are shown in the following table linearly.

Table 33 EGFR exon 20 insertions—concordance

Agreement measure	Excluding unknowns ^[1]		Including unknowns ^[1]	
	Percent agreement	95% CI	Percent agreement	95% CI
PPA	100% (54/54)	(93.4%, 100%)	98.2% (54/55)	(90.3%, 100%)
NPA	100% (95/95)	(96.2%, 100%)	90.5% (95/105)	(83.2.0%, 95.3%)
OPA	100% (149/149)	(97.6%, 100%)	93.1% (149/160)	(88.0%, 96.5%)

^[1] Unknowns are defined as values due to insufficient sample, or sample QC sequencing failure resulting in an invalid result or No Call for the variant.

EGFR exon 20 insertions clinical effectiveness—Study II

The clinical effectiveness of the Oncomine™ Dx Target Test was evaluated by measuring the overall response rate (ORR; complete response or partial response) and duration of response (DOR) for patients with NSCLC that tested positive for EGFR exon 20 insertion variants by the Oncomine™ Dx Target Test. The ORR and DOR were calculated for 102 patients with prior platinum treatment selected for treatment with mobocertinib in the AP32788-15-101 trial, 27 of whom were responders.

The ORR was 26.7% (95% CI: 16.1, 39.7) in the patient population testing positive for EGFR exon 20 insertions with the Oncomine™ Dx Target Test (16/60) in comparison to 18.2% (95% CI: 2.3, 51.8) in the patient population testing negative for EGFR exon 20 insertions (2/11). The 2 samples from responding

patients that tested negative with the Oncomine™ Dx Target Test also tested negative with the reference assay, suggesting that the samples harbored novel variants that are not currently reported by either test.

The 26.7% ORR was similar to the ORR of 26.5% (95% CI: 18.2, 36.1) for the entire CDx or Oncomine™ Dx Target Test population (27/102) due to a large number of samples (n=10) that were unavailable for testing, of which 4 were from patients that showed partial response. The 26.7% ORR was consistent with the response rate of the LLT-positive population.

The median DOR by the Kaplan-Meier (KM) method was 11.07 months (95% CI: 9.13, 16.59) in the patient population testing positive for EGFR exon 20 insertions with the Oncomine™ Dx Target Test (n = 16), in comparison to 13.83 months (95% CI: 11.07, 16.59) for samples testing negative for EGFR exon 20 insertions (n = 2).

In total, 60 Oncomine™ Dx Target Test EGFR exon 20 insertion-positive, prior platinum-treated patients were included in the drug efficacy evaluation, 52 of which were concordant with the reference assay and LLT results, and 8 with only the LLT results available (reference assay unknown).

Refer to the [Drugs@FDA database](#) for the most recent therapeutic product labeling.

RET clinical study

RET study—concordance evaluation

To evaluate the ability of the Oncomine™ Dx Target Test RNA panel to identify RET fusions in human specimens, 238 FFPE tumor samples were analyzed using the Oncomine™ Dx Target Test to demonstrate positive percent agreement (PPA) and negative percent agreement (NPA) with a validated reference detection method (Archer assay).

Of the 238 samples were evaluated in the study, 133 were identified by the enrolling LLTs as RET fusion-positive in the BLU-667-1101 trial, and 105 were stage-matched commercially sourced NSCLC samples, screened with either a local laboratory-validated RET FISH assay, or an NGS-based screening assay. After screening 105 NSCLC samples, 2 were identified as RET fusion-positive by the LLTs and were grouped with the clinical trial samples, resulting in 105 LLT-positive and 103 LLT-negative samples.

Of the 135 RET fusion-positive samples by the LLTs, 54 were cancelled before sequencing by Oncomine™ Dx Target Test due to failure to meet test input requirements—22 samples had insufficient tissue available, 1 sample had an insufficient number of slides, 8 samples did not meet the tumor content requirement, and 23 samples failed the RNA concentration cutoff. Of the remaining 81 samples, 56 were positive by the Oncomine™ Dx Target Test, 24 samples were called negative, and 1 sample yielded an invalid result.

Of the 103 RET fusion-negative samples (by LLT) available for testing, 102 samples were called negative by the Oncomine™ Dx Target Test, and 1 sample yielded an invalid result.

For testing with the Archer assay, of the 135 RET fusion-positive samples by the LLTs, in addition to the 54 cancelled for the Oncomine™ Dx Target Test, 11 more samples were cancelled due to insufficient RNA concentration, and 4 samples were not tested on the Archer Assay. Of the remaining 66 samples, 43 were positive by the Archer assay, 17 samples were called negative, and 6 samples yielded an invalid result. Of the 103 RET fusion-negative samples (by LLT), 1 sample had insufficient RNA volume for testing, 93 were negative, 1 was positive, and 8 were invalid by the Archer assay.

The PPA was defined as the proportion of RET fusion-positive specimens as called by the Archer assay that were also RET fusion-positive as called by the Oncomine™ Dx Target Test, and the NPA was defined as the proportion of RET fusion-negative specimens as called by the Archer assay that were also RET fusion-negative as called by the Oncomine™ Dx Target Test. Concordance between the Oncomine™ Dx Target Test and the Archer assay is shown in Table 34.

Table 34 Concordance between the Archer assay and the Oncomine™ Dx Target Test

Agreement measure	Excluding unknowns		Including unknowns ^[1]	
	Percent agreement	95% CI	Percent agreement	95% CI
PPA	90.9% (40/44)	(78.3%, 97.5%)	90.9% (40/44)	(78.3%, 97.5%)
NPA	91.8% (101/110)	(85.0%, 96.2%)	91.8% (101/110)	(85.0%, 96.2%)
OPA	91.6% (141/154) ^[2]	(86.0%, 95.4%)	91.6% (141/154)	(86.0%, 95.4%)

^[1] The results including and excluding unknowns were identical. Unknowns are defined as invalid or no result using the Oncomine™ Dx Target Test.

^[2] Thirteen samples were found to be discordant in this analysis, where four were false negatives and nine were false positives with the Oncomine™ Dx Target Test.

RET study—clinical effectiveness

The clinical effectiveness of the Oncomine™ Dx Target Test was evaluated by measuring the overall response rate (ORR) for patients with NSCLC that tested positive for a RET fusion by both the LLTs and the Oncomine™ Dx Target Test. The ORR was calculated for patients selected for treatment with the RET inhibitor pralsetinib, with prior platinum treatment, and with no prior systemic treatment. The disease control rate (DCR) and clinical benefit rate (CBR) were also calculated for these patients.

In the efficacy population, the ORR for the NSCLC patients with prior platinum treatment was 70.6% (95% CI: 52.5, 84.9). The ORR for NSCLC patients with no prior systemic treatment was 85.7% (95% CI: 57.2, 98.2).

In the efficacy population, the DCR was 100.0% (95% CI: 91.6, 100.0) for patients with prior platinum treatment, with 34 patients having complete response (CR), partial response (PR), or stable disease (SD). The DCR was 92.9% (95% CI: 66.1, 99.8) for patients with no prior systemic treatment, with 13 patients having complete response (CR), partial response (PR), or stable disease (SD).

In the efficacy population, the CBR was 82.4% for patients with prior platinum treatment, with 28 patients having CR, PR, or SD for at least 16 weeks with 28 days in a cycle. The CBR was 85.7% for patients with no prior systemic treatment, with 12 patients having CR, PR, or SD for at least 16 weeks with 28 days in a cycle.

Refer to the [Drugs@FDA database](#) for the most recent therapeutic product labeling.

ROS1 clinical studies

ROS1 concordance evaluation—Study I

To evaluate the ability of the Oncomine™ Dx Target Test to identify the ROS1 biomarker in NSCLC tumor samples, 19 specimens from patients in the Phase 1 Pfizer Study A8081001 that tested positive using a ROS1 FISH assay were used, together with 13 archival specimens that also tested positive. These 32 positive specimens and 126 archival specimens that tested negative using the ROS1 FISH assay were analyzed using the Oncomine™ Dx Target Test.

Of the ROS1-positive samples, 25 generated valid results from both the FISH assay and the Oncomine™ Dx Target Test. Of the remaining samples, 4 generated invalid sequencing results due to a control or library QC failure, 2 generated insufficient material for FISH assay analysis, and 1 was subsequently determined to be a false positive for ROS1.

Of the ROS1-negative samples, 119 generated valid results from both the FISH assay and the test, while 7 generated invalid sequencing results due to a control or library QC failure. A total of 144 samples were used to evaluate concordance between the assay and the test. Of these, 139 were FFPE specimens and 5 were extracted RNA samples.

The PPA was defined as the proportion of ROS1-positive specimens called by the ROS1 FISH assay that were also called by the Oncomine™ Dx Target Test, and the NPA was defined as the proportion of ROS1-negative specimens called by the ROS1 FISH assay that were also called by the test. The OPA was defined as the number of calls where the ROS1 FISH assay and the Oncomine™ Dx Target Test agreed, divided by the total number of calls made. The 95% CIs were determined for PPA, NPA, and OPA, and the results are shown in the following table:

Table 35 Concordance between the ROS1 FISH assay and the Oncomine™ Dx Target Test

Agreement measure	Percent agreement	95% CI
PPA	80.0% (20/25)	59.3%, 93.2%
NPA	100% (119/119)	96.9%, 100%
OPA	96.5% (139/144)	92.08%, 98.86%

Of the 20 concordant ROS1-positive samples, 17 were from FFPE tissue samples and 3 were from RNA extracts. For FFPE specimens alone, excluding invalids, the PPA was 85.0% (17/20) and the NPA was 100% (119/119). For the extracted RNA specimens alone, excluding invalids, the PPA was 60.0% (3/5) and the NPA was not evaluable because all specimens were ROS1 positive. The results from the 5 RNA extraction specimens should be interpreted with caution due to the limited sample size.

All 5 discordant samples were positive for the FISH assay and negative for the test. Three of these also tested negative using a probe hybridization fusion detection method.

ROS1 concordance evaluation—Study II

An additional concordance study was performed using the updated Oncomine™ Dx Target Test RNA library preparation workflow, to verify that the protocol changes did not impact the effectiveness of the test. In this study, results from the updated test workflow were compared to results from the Kreatech™ ROS1 FISH Assay and to results from the original test workflow.

Of the ROS1-positive samples, 9 generated valid results (excluding unknowns) from both the ROS1 FISH assay and the updated Oncomine™ Dx Target Test workflow. Of the ROS1-negative samples, 62 generated valid results from both, for a total of 71 samples with valid results used to evaluate concordance between the test as an investigational method and the ROS1 FISH assay as the reference method.

Of the ROS1-positive samples, 7 generated valid results (excluding unknowns) from both the ROS1 FISH assay and the original Oncomine™ Dx Target Test workflow. Of the ROS1-negative samples, 59 generated valid results from both, for a total of 67 samples with valid results used to evaluate

concordance between the test as an investigational method and the ROS1 FISH assay as the reference method.

Table 36 Overall concordance between the ROS1 FISH assay replicates and the updated OncoPrint™ Dx Target Test RNA library preparation workflow

Agreement measure	Excluding unknowns ^[1]		Including unknowns	
	Percent agreement	95% CI	Percent agreement	95% CI
PPA	100% (9/9)	66.4%, 100.0%	90% (9/10)	55.5%, 99.8%
NPA	100% (62/62)	94.2%, 100.0%	89% (62/70)	78.7%, 94.9%
OPA	100% (71/71)	94.9%, 100.0%	89% (71/80)	79.7%, 94.7%

^[1] Unknowns are defined as invalid or no result using the OncoPrint™ Dx Target Test

Excluding unknowns, the updated OncoPrint™ Dx Target Test workflow showed a 100% PPA, NPA, and OPA with the ROS1 FISH assay.

Table 37 Overall concordance between the updated OncoPrint™ Dx Target Test vs original OncoPrint™ Dx Target Test workflow

Agreement measure	Excluding unknowns ^[1]		Including unknowns	
	Percent agreement	95% CI	Percent agreement	95% CI
PPA	100%	71.5%, 100.0%	100%	71.5%, 100.0%
NPA	98%	92.0%, 99.7%	98%	92.0%, 99.7%
OPA	98%	92.9%, 99.8%	98%	92.9%, 99.8%

^[1] Unknowns are defined as invalid or no result using the OncoPrint™ Dx Target Test

Excluding unknowns, the updated OncoPrint™ Dx Target Test workflow showed a 100% PPA, 98% NPA, and 98% OPA agreement with the original OncoPrint™ Dx Target Test workflow.

ROS1 study—clinical outcomes evaluation

As part of the Study I concordance evaluation described above, clinical outcomes as measured by objective response rate (ORR) and duration of response (DOR) with XALKORI® (crizotinib) were evaluated for 11 patients whose tumors were designated as ROS1-positive by the ROS1 FISH assay and whose tumors were evaluable by the OncoPrint™ Dx Target Test. Of these, 6 samples tested positive by both tests.

The ORR for patients with tumor specimens determined to be ROS1-positive using both tests was 83.3% (5/6) (95% CI: 35.88%, 99.58%).

The mean DOR (N=5) was 17.5 months (95% CI: 10.9, 24.1).

Refer to the [Drugs@FDA database](#) for the most recent therapeutic product labeling.



Variants detected by the Oncomine™ Dx Target Test

DNA variants detected in non-small cell lung cancer

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
AKT1	3	p.Glu17Lys	c.49G>A	COSM33765
ALK	21	p.Gly1128Ala	c.3383G>C	COSM98475
ALK	22	p.Leu1152Arg	c.3455T>G	COSM97185
ALK	22	p.Leu1152Pro	c.3455T>C	COSM1407659
ALK	22	p.Cys1156Tyr	c.3467G>A	COSM99136 Note: Some "no calls" were observed for this analytical variant due to strand bias with plasmid targets. This does not impact clinical test results.
ALK	22	p.Ile1171Asn	c.3512T>A	COSM28498
ALK	22	p.Ile1171Thr	c.3512T>C	COSM4381100 Note: Some "no calls" were observed for this analytical variant due to strand bias with plasmid targets. This does not impact clinical test results.
ALK	23	p.Phe1174Val	c.3520T>G	COSM28054
ALK	23	p.Phe1174Leu	c.3520T>C	COSM28057
ALK	23	p.Phe1174Ile	c.3520T>A	COSM28491
ALK	23	p.Phe1174Cys	c.3521T>G	COSM28059
ALK	23	p.Phe1174Ser	c.3521T>C	COSM53063
ALK	23	p.Phe1174Leu	c.3522C>G	COSM28061
ALK	23	p.Phe1174Leu	c.3522C>A	COSM28055

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
ALK	23	p.Val1180Leu	c.3538G>C	COSM4381101
ALK	23	p.Leu1196Met	c.3586C>A	COSM99137
ALK	23	p.Leu1196Gln	c.3587T>A	COSM1169447
ALK	23	p.Gly1202Arg	c.3604G>A	COSM144250
ALK	23	p.Ser1206Tyr	c.3617C>A	COSM144251
ALK	24	p.Phe1245Val	c.3733T>G	COSM28499
ALK	24	p.Phe1245Ile	c.3733T>A	COSM28492
ALK	24	p.Phe1245Cys	c.3734T>G	COSM28500
ALK	24	p.Phe1245Leu	c.3735C>G	COSM28062
ALK	24	p.Phe1245Leu	c.3735C>A	COSM28493
ALK	25	p.Arg1275Leu	c.3824G>T	COSM28060
ALK	25	p.Arg1275Gln	c.3824G>A	COSM28056
BRAF	11	p.Gly466Val	c.1397G>T	COSM451
BRAF	11	p.Gly466Glu	c.1397G>A	COSM453
BRAF	11	p.Gly469Arg	c.1405G>A	COSM457
BRAF	11	p.Gly469Val	c.1406G>T	COSM459
BRAF	11	p.Gly469Ala	c.1406G>C	COSM460
BRAF	15	p.Asp594Asn	c.1780G>A	COSM27639
BRAF	15	p.Asp594Gly	c.1781A>G	COSM467
BRAF	15	p.Val600Arg	c.1798_1799delGTinsAG	COSM474
BRAF	15	p.Val600Lys	c.1798_1799delGTinsAA	COSM473
BRAF	15	p.Val600Glu	c.1799T>A	COSM476
BRAF	15	p.Val600Glu	c.1799_1800delTGinsAA	COSM475
BRAF	15	p.Val600_Lys601delinsGlu	c.1799_1801delTGA	COSM1133
BRAF	15	p.Lys601Glu	c.1801A>G	COSM478
CDK4	2	p.Lys22Gln	c.64A>C	OM3153
CDK4	2	p.Lys22Arg	c.65A>G	COSM232013
CDK4	2	p.Lys22Met	c.65A>T	COSM3463915

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
CDK4	2	p.Arg24Cys	c.70C>T	COSM1677139
CDK4	2	p.Arg24Ser	c.70C>A	COSM3463914
CDK4	2	p.Arg24Leu	c.71G>T	COSM363684
CDK4	2	p.Arg24His	c.71G>A	COSM1989836
DDR2	5	p.Arg124Trp	c.370C>T	COSM4024594
DDR2	5	p.Arg124Leu	c.371G>T	COSM400880
EGFR	3	p.Arg108Gly	c.322A>G	COSM1451536
EGFR	3	p.Arg108Lys	c.323G>A	COSM21683
EGFR	7	p.Ala289Thr	c.865G>A	COSM21686
EGFR	7	p.Ala289Asp	c.866C>A	COSM21685
EGFR	7	p.Ala289Val	c.866C>T	COSM21687
EGFR	12	p.Ser492Arg	c.1474A>C	COSM236671
EGFR	12	p.Ser492Arg	c.1476C>A	COSM236670
EGFR	15	p.Gly598Ala	c.1793G>C	COSM3412196
EGFR	15	p.Gly598Val	c.1793G>T	COSM21690
EGFR	18	p.Glu709Lys	c.2125G>A	COSM12988
EGFR	18	p.Glu709Ala	c.2126A>C	COSM13427
EGFR	18	p.Glu709Gly	c.2126A>G	COSM13009
EGFR	18	p.Glu709Val	c.2126A>T	COSM12371
EGFR	18	p.Gly719Ser	c.2155G>A	COSM6252
EGFR	18	p.Gly719Cys	c.2155G>T	COSM6253
EGFR	18	p.Gly719Asp	c.2156G>A	COSM18425
EGFR	18	p.Gly719Ala	c.2156G>C	COSM6239
EGFR	19	p.Lys745_Glu749del	c.2233_2247delAAGGAATTAA GAGAA	COSM26038
EGFR	19	p.Lys745_Ala750delinsThr	c.2234_2248delAGGAATTAAG AGAAG	COSM1190791
EGFR	19	p.Glu746_Glu749del	c.2235_2246delGGAATTAAGA GA	COSM28517

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
EGFR	19	p.Glu746_Ala750del	c.2235_2249delGGAATTAAGA GAAGC	COSM6223
EGFR	19	p.Glu746_Thr751delinsIle	c.2235_2252delGGAATTAAGA GAAGCAACinsAAT	COSM13551 Note: The nucleotide change of COSM13551 overlaps that of COSM6223, so a positive COSM13551 sample will also result in a positive call for COSM6223.
EGFR	19	p.Glu746_Ala750del	c.2236_2250delGAATTAAGAG AAGCA	COSM6225
EGFR	19	p.Glu746_Arg748del	c.2239_2247delTTAAGAGAA	COSM6218
EGFR	19	p.Glu746_Thr751del	c.2236_2253delGAATTAAGAG AAGCAACA	COSM12728
EGFR	19	p.Leu747_Ala750delinsPro	c.2239_2248delTTAAGAGAAGi nsC	COSM12382
EGFR	19	p.Leu747_Thr751delinsPro	c.2239_2251delTTAAGAGAAG CAAinsC	COSM12383
EGFR	19	p.Glu746_Thr751delinsAla	c.2237_2251delAATTAAGAGAA GCAA	COSM12678 Note: A false negative call was observed for this variant when tested with plasmid targets for 1 out of 4 of the replicates tested.
EGFR	19	p.Glu746_Ser752delinsAsp	c.2238_2255delATTAAGAGAA GCAACATC	COSM6220
EGFR	19	p.Glu746_Ser752delinsVal	c.2237_2255delAATTAAGAGAA GCAACATCinsT	COSM12384
EGFR	19	p.Glu746_Thr751delinsValAla	c.2237_2253delAATTAAGAGAA GCAACAinsTTGCT	COSM12416
EGFR	19	p.Leu747_Thr751del	c.2240_2254delTAAGAGAAGC AACAT	COSM12369
EGFR	19	p.Leu747_Ser752del	c.2239_2256delTTAAGAGAAG CAACATCT	COSM6255
EGFR	19	p.Leu747_Thr751delinsGln	c.2238_2252delATTAAGAGAA GCAACinsGCA	COSM12419

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
EGFR	19	p.Leu747_Ala750delinsPro	c.2238_2248delATTAAGAGAA GinsGC	COSM12422
EGFR	19	p.Leu747_Pro753delinsGln	c.2239_2258delTTAAGAGAAG CAACATCTCCinsCA	COSM12387 Note: The nucleotide change of COSM12387 overlaps that of COSM6255, so a positive COSM12387 sample will also result in a positive call for COSM6255.
EGFR	19	p.Leu747_Thr751delinsSer	c.2240_2251delTAAGAGAAGC AA	COSM6210
EGFR	19	p.Leu747_Pro753delinsSer	c.2240_2257delTAAGAGAAGC AACATCTC	COSM12370
EGFR	20	NA	NA	COSM26720 ^[1]
EGFR	20	p.Met766_Ala767insAlaThrLeu	c.2302_2303insCGCTGGCCA	COSM12425
EGFR	20	p.Met766_Ala767insAlaSerVal	c.2308_2309insCCAGCGTGG	COSM12376
EGFR	20	p.Met766_Ala767insAlaIle	c.2302_2303insTAGCCA	COSM13559
EGFR	20	p.Ala767_Ser768insSerValAsp	c.2311_2312insGCGTGGACA	COSM13428
EGFR	20	p.Ala767_Ser768insSerValGly	c.2308_2309insGCAGCGTGG	COSM18429
EGFR	20	p.Ala767_Ser768insSerValGly	c.2308_2309insGGAGCGTGG	COSM1235344
EGFR	20	p.Ala767_Ser768insTyrValMet	c.2301_2302insTACGTGATG	COSM1651740
EGFR	20	p.Ser768_Val769insValCys	c.2307_2308insTGCGTG	COSM12379
EGFR	20	p.Ser768_Val769insValAspAsn	c.2313_2314insGTGGACAAC	COSM20885
EGFR	20	p.Ser768_Val769insValGlyVal	c.2308_2309insGGGTCTGG	COSM18430
EGFR	20	p.Ser768_Val769insValAlaAsn	c.2303_2304insTGTGGCCAA	COSM1651741
EGFR	20	p.Ser768Ile	c.2303G>T	COSM6241
EGFR	20	p.Val769dup	c.2308_2309insTGG	COSM6506514
EGFR	20	p.Val769_Asp770insAspGly	c.2310_2311insGGGGAC	COSM85795
EGFR	20	p.Val769_Asp770insAspAsnPro	c.2316_2317insGACAACCCC	COSM1651745
EGFR	20	p.Val769_Asp770insAspGly	c.2310_2311insGGCGAC	COSM22955
EGFR	20	p.Val769_Asp770insGluArgGly	c.2309_2310insGCGTGGAGA	COSM1651742

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
EGFR	20	p.Asp769insAlaSerVal	c.2309_2310delACinsCCAGCG TGGAT	COSM13558
EGFR	20	p.Asp770_Asn771insAsnProHis	c.2319_2320insAACCCCCAC	COSM12381
EGFR	20	p.Asp770_Asn771insTyr	c.2310_2311insTAC	COSM1238030
EGFR	20	p.Asp770delinsGlyTyr	c.2308_2309insGTT	COSM12427
EGFR	20	p.Asp770_Asn771insAsnPro	c.2316_2317insAACCCC	MAN123
EGFR	20	p.Asp770_Asn771insAsnProHis Gly	c.2320_2321insGCAACCCCCA CG	COSM51544
EGFR	20	p.Asp770_Asn771insThr	c.2311_2312insCCA	COSM5023008
EGFR	20	p.Asp770_Asn771insMetAlaThr Pro	c.2311_2312insTGGCCACCCC CA	COSM26719
EGFR	20	p.Asp770_Asn771insGlnArgGly	c.2310_2311insCAGCGTGGC	COSM4970107
EGFR	20	p.Asp770_Asn771insGly	c.2310_2311insGGC	COSM13004
EGFR	20	p.Asp770_Asn771delinsAlaGlyGly	c.2309_2312delACAAinsCTGG TGG	COSM12737
EGFR	20	p.Asp770_Asn771insSerValGlu	c.2311_2312insGCGTCGAAA	COSM1651743
EGFR	20	p.Asp770_Asn771insAlaProTrp	c.2310_2311insGCACCGTGG	COSM20886
EGFR	20	p.Asp770_Asn771insGlyThr	c.2310_2311insGGCACA	COSM1238029
EGFR	20	p.Asp770_Asn771insGlyLeu	c.2310_2311insGGGTTA	COSM48921
EGFR	20	p.Asp770_Asn771insGlyPhe	c.2310_2311insGGGTTT	COSM655155
EGFR	20	p.Asp770_Asn771insGly	c.2310_2311insGGT	COSM12378
EGFR	20	p.Asn771dup	c.2313_2314insAAC	COSM13003
EGFR	20	p.Asn771_Pro772insHis	c.2314_2315insACC	COSM1238031
EGFR	20	p.Asn771_Pro772insProHis	c.2319_2320insCCCCAC	COSM12380
EGFR	20	p.Asn771_Pro772insProThrHis	c.2315_2316insGACACACCC	COSM48923
EGFR	20	p.Asn771_Pro772insArgHis	c.2314_2315insGGCACC	COSM166390
EGFR	20	p.Asn771delinsThrHis	c.2311_2312insCAC	COSM22946
EGFR	20	p.Asn771delinsSerGlyHis	c.2311_2312insGTGGCC	COSM1651744
EGFR	20	p.Asn771delinsValHis	c.2311_2311delAinsGTCC	COSM5023007
EGFR	20	p.Asn771delinsLysLeu	c.2312_2313insACT	COSM6438147



(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
EGFR	20	p.Pro772_His773insHisAla	c.2320_2321insCCCACG	COSM1238028
EGFR	20	p.Pro772_His773insHisAsn	c.2319_2320insAACAC	COSM5023006
EGFR	20	p.Pro772_His773insHisVal	c.2321_2322insCCACGT	COSM18432
EGFR	20	p.Pro772_His773insThrPro	c.2316_2316delCinsAACCCCT	COSM12388
EGFR	20	p.Pro772_His773insVal	c.2316_2317insGTT	COSM255205
EGFR	20	p.Pro773_Val774dup	c.2322_2323insCACGTG	COSM22948
EGFR	20	p.His773dup	c.2319_2320insCAC	COSM12377
EGFR	20	p.His773delinsProAsnProTyr	c.2317_2318insCTAACCCCT	COSM1735761
EGFR	20	p.His773_Val774insThrGlnProPro	c.2319_2320insACACAACCC CC	COSM3727813
EGFR	20	p.His773_Val774insGln	c.2319_2320insCAG	COSM131552
EGFR	20	p.Val774_Cys775insProArg	c.2322_2323insCCACGT	COSM4170223
EGFR	20	p.Cys797Ser	c.2389T>A	COSM6493937
EGFR	20	p.Cys797Ser	c.2390G>C	COSM5945664
EGFR	21	p.Leu858Met	c.2572C>A	COSM12366
EGFR	21	p.Leu858Arg	c.2573T>G	COSM6224
EGFR	21	p.Leu861Gln	c.2582T>A	COSM6213
EGFR	21	p.Leu861Arg	c.2582T>G	COSM12374
ERBB2	8	p.Ser310Tyr	c.929C>A	COSM94225
ERBB2	8	p.Ser310Phe	c.929C>T	COSM48358
ERBB2	17	p.Arg678Gln	c.2033G>A	COSM436498
ERBB2	18	p.Thr733Ile	c.2198C>T	COSM14059
ERBB2	19	p.Leu755Met	c.2263T>A	COSM1205571
ERBB2	19	p.Leu755Pro	c.2263_2264delTTinsCC	COSM683
ERBB2	19	p.Asp769His	c.2305G>C	COSM13170
ERBB2	19	p.Asp769Tyr	c.2305G>T	COSM1251412
ERBB2	20	p.Gly776Val	c.2327G>T	COSM18609
ERBB2	20	p.Val777Leu	c.2329G>T	COSM14062
ERBB2	21	p.Val842Ile	c.2524G>A	COSM14065

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
ERBB2	22	p.Arg896Cys	c.2686C>T	COSM14066
ERBB2	22	p.Arg896His	c.2687G>A	COSM119971
ERBB3	2	p.Met60Leu	c.178A>T	COSM1606366
ERBB3	2	p.Met60Lys	c.179T>A	COSM254678
ERBB3	2	p.Met60Arg	c.179T>G	COSM941484
ERBB3	3	p.Met91Ile	c.273G>A	COSM122890
ERBB3	3	p.Met91Ile	c.273G>C	COSM1299636
ERBB3	3	p.Val104Met	c.310G>A	COSM172423
ERBB3	3	p.Val104Leu	c.310G>C	COSM160824
ERBB3	3	p.Val104Leu	c.310G>T	COSM191840
ERBB3	6	p.Ala232Thr	c.694G>A	COSM4043440
ERBB3	6	p.Ala232Val	c.695C>T	COSM1242239
ERBB3	8	p.Asp297Tyr	c.889G>T	COSM160822
ERBB3	8	p.Asp297Val	c.890A>T	COSM941490
ERBB3	9	p.Glu332Lys	c.994G>A	COSM254677
FGFR2	7	p.Pro253Leu	c.758C>T	COSM537801
FGFR2	7	p.Pro253Arg	c.758C>G	COSM49170
FGFR2	8	p.Ala314Asp	c.941C>A	COSM49171
FGFR2	9	p.Tyr375His	c.1123T>C	COSM1560916
FGFR2	9	p.Tyr375Cys	c.1124A>G	COSM36904
FGFR2	9	p.Cys382Arg	c.1144T>C	COSM36906
FGFR2	9	p.Cys382Tyr	c.1145G>A	COSM915493
FGFR2	12	p.Asn549His	c.1645A>C	COSM250083
FGFR2	12	p.Asn549Ser	c.1646A>G	COSM3665553
FGFR2	12	p.Asn549Lys	c.1647T>G	COSM36902
FGFR2	12	p.Asn549Lys	c.1647T>A	COSM36912
FGFR2	14	p.Lys659Glu	c.1975A>G	COSM36909
FGFR2	14	p.Lys659Met	c.1976A>T	COSM49175

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
FGFR2	14	p.Lys659Asn	c.1977G>T	COSM49173
FGFR2	14	p.Lys659Asn	c.1977G>C	COSM683054
FGFR2	7	p.Ser252Trp	c.755C>G	COSM36903
FGFR3	7	p.Arg248Cys	c.742C>T	COSM714
FGFR3	7	p.Ser249Cys	c.746C>G	COSM715
FGFR3	14	p.Lys650Gln	c.1948A>C	COSM726
FGFR3	14	p.Lys650Glu	c.1948A>G	COSM719
FGFR3	14	p.Lys650Asn	c.1950G>T	COSM1428730
FGFR3	16	p.Gly697Cys	c.2089G>T	COSM24802
HRAS	2	p.Gly12Cys	c.34G>T	COSM481
HRAS	2	p.Gly12Arg	c.34G>C	COSM482
HRAS	2	p.Gly12Val	c.35G>T	COSM483
HRAS	2	p.Gly12Asp	c.35G>A	COSM484
HRAS	2	p.Gly12Ala	c.35G>C	COSM485
HRAS	2	p.Gly13Arg	c.37G>C	COSM486
HRAS	2	p.Gly13Ser	c.37G>A	COSM487
HRAS	2	p.Gly13Cys	c.37G>T	COSM488
HRAS	2	p.Gly13Val	c.38G>T	COSM489
HRAS	2	p.Gly13Asp	c.38G>A	COSM490
HRAS	3	p.Gln61Lys	c.181C>A	COSM496
HRAS	3	p.Gln61Leu	c.182A>T	COSM498
HRAS	3	p.Gln61Arg	c.182A>G	COSM499
HRAS	3	p.Gln61Pro	c.182A>C	COSM500
HRAS	3	p.Gln61His	c.183G>T	COSM502
HRAS	3	p.Gln61His	c.183G>C	COSM503
HRAS	2	p.Gly12Ser	c.34G>A	COSM480
KIT	8	p.Asp419del	c.1255_1257delGAC	COSM29014
KIT	8	p.Asp419_Arg420del	c.1255_1260delGACAGG	COSM1578132

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
KIT	11	p.Trp557_Lys558del	c.1669_1674delTGGAAAG	COSM1217
KIT	11	p.Trp557Arg	c.1669T>A	COSM1216
KIT	11	p.Trp557Arg	c.1669T>C	COSM1219
KIT	11	p.Trp557Gly	c.1669T>G	COSM1221
KIT	11	p.Trp557_Val559delinsPhe	c.1670_1675delGGAAGG	COSM1226
KIT	11	p.Val559del	c.1679_1681delTTG	COSM1247
KIT	11	p.Val559Asp	c.1676T>A	COSM1252
KIT	11	p.Val559Ala	c.1676T>C	COSM1255
KIT	11	p.Val559Gly	c.1676T>G	COSM1253
KIT	11	p.Val560Asp	c.1679T>A	COSM1257
KIT	11	p.Leu576Pro	c.1727T>C	COSM1290
KIT	11	p.Asp579del	c.1735_1737delGAT	COSM1294
KIT	13	p.Lys642Glu	c.1924A>G	COSM1304
KIT	13	p.Val654Ala	c.1961T>C	COSM12706
KIT	17	p.Arg796Lys	c.2387G>A	COSM1600411
KIT	17	p.Asp816His	c.2446G>C	COSM1311
KIT	17	p.Asp816Tyr	c.2446G>T	COSM1310
KIT	17	p.Asp816Val	c.2447A>T	COSM1314
KIT	17	p.Asn822Lys	c.2466T>A	COSM1321
KIT	17	p.Asn822Lys	c.2466T>G	COSM1322
KIT	17	p.Val825Ala	c.2474T>C	COSM1323
KRAS	2	p.Gly12Cys	c.34G>T	COSM516
KRAS	2	p.Gly12Ser	c.34G>A	COSM517
KRAS	2	p.Gly12Arg	c.34G>C	COSM518

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
KRAS	2	p.Gly12Phe	c.34_35delGGinsTT	COSM512 Note: The nucleotide change of COSM512 overlaps that of COSM516, so a positive COSM512 sample will also result in a positive call for COSM516.
KRAS	2	p.Gly12Val	c.35G>T	COSM520
KRAS	2	p.Gly12Ala	c.35G>C	COSM522
KRAS	2	p.Gly12Asp	c.35G>A	COSM521
KRAS	2	p.Gly13Cys	c.37G>T	COSM527
KRAS	2	p.Gly13Ser	c.37G>A	COSM528
KRAS	2	p.Gly13Arg	c.37G>C	COSM529
KRAS	2	p.Gly13Asp	c.38_39delGCinsAT	COSM531 Note: The nucleotide change of COSM531 overlaps that of COSM532, so a positive COSM531 sample will also result in a positive call for COSM532.
KRAS	2	p.Gly13Val	c.38G>T	COSM534
KRAS	2	p.Gly13Ala	c.38G>C	COSM533
KRAS	2	p.Gly13Asp	c.38G>A	COSM532
KRAS	3	p.Ala59Thr	c.175G>A	COSM546
KRAS	3	p.Ala59Gly	c.176C>G	COSM28518
KRAS	3	p.Ala59Glu	c.176C>A	COSM547
KRAS	3	p.Gln61Lys	c.180_181delTCinsAA	COSM87298
KRAS	3	p.Gln61Lys	c.181C>A	COSM549
KRAS	3	p.Gln61Glu	c.181C>G	COSM550
KRAS	3	p.Gln61Pro	c.182A>C	COSM551
KRAS	3	p.Gln61Arg	c.182A>G	COSM552
KRAS	3	p.Gln61Leu	c.182A>T	COSM553

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
KRAS	3	p.Gln61His	c.183A>C	COSM554
KRAS	3	p.Gln61His	c.183A>T	COSM555
KRAS	4	p.Lys117Asn	c.351A>T	COSM28519
KRAS	4	p.Lys117Asn	c.351A>C	COSM19940
KRAS	4	p.Ala146Thr	c.436G>A	COSM19404
KRAS	4	p.Ala146Pro	c.436G>C	COSM19905
KRAS	4	p.Ala146Val	c.437C>T	COSM19900
MAP2K1	2	p.Phe53Ile	c.157T>A	COSM3503329
MAP2K1	2	p.Phe53Leu	c.157T>C	COSM555604
MAP2K1	2	p.Phe53Val	c.157T>G	COSM1562837 Note: The base change c.157T>G in MAP2K1 is associated with Mutation ID COSM5077832 in the COSMIC v.76 database, even though it has been given the Variant HotSpot ID COSM1562837 in the software. This does not impact the test results.
MAP2K1	2	p.Phe53Leu	c.159T>A	COSM1725008
MAP2K1	2	p.Phe53Leu	c.159T>G	COSM2257208
MAP2K1	2	p.Lys57Thr	c.170A>C	COSM4756761
MAP2K1	2	p.Lys57Met	c.170A>T	MAN124
MAP2K1	2	p.Lys57Asn	c.171G>C	COSM5520914
MAP2K1	2	p.Lys57Asn	c.171G>T	COSM1235478
MAP2K1	3	p.Pro124Ser	c.370C>T	COSM235614
MAP2K1	3	p.Pro124Gln	c.371C>A	COSM1167912
MAP2K1	3	p.Pro124Leu	c.371C>T	COSM1315861
MAP2K1	6	p.Glu203Lys	c.607G>A	COSM232755
MAP2K1	6	p.Glu203Val	c.608A>T	COSM3386991
MAP2K2	2	p.Phe57Val	c.169T>G	COSM3534171

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
MAP2K2	2	p.Phe57Leu	c.169T>C	COSM1235618
MAP2K2	2	p.Phe57Leu	c.171T>G	OM3158
MAP2K2	2	p.Phe57Leu	c.171T>A	COSM3389034
MAP2K2	2	p.Gln60Pro	c.179A>C	COSM145610
MET	14	p.Thr1010Ile	c.3029C>T	COSM707
MET	14	p.Tyr1021Asn	c.3061T>A	COSM48564
MET	14	p.Tyr1021Phe	c.3062A>T	COSM339515
MET	—	NA	NA	COSM29633 ^[1]
MET	—	NA	NA	COSM24687 ^[1]
MET	—	NA	NA	COSM35468 ^[1]
MET	16	p.His1112Tyr	c.3334C>T	COSM696
MET	16	p.His1112Arg	c.3335A>G	COSM703
MET	16	p.His1112Leu	c.3335A>T	COSM698
MET	19	p.Tyr1248His	c.3742T>C	COSM690
MET	19	p.Tyr1248Cys	c.3743A>G	COSM699
MET	19	p.Tyr1253Asp	c.3757T>G	COSM700
MET	19	p.Met1268Thr	c.3803T>C	COSM691
MET	19	p.Met1268Ile	c.3804G>A	COSM694
MTOR	30	p.Cys1483Phe	c.4448G>T	COSM462616
MTOR	30	p.Cys1483Tyr	c.4448G>A	COSM462615
MTOR	30	p.Cys1483Trp	c.4449C>G	OM3149
MTOR	39	p.Glu1799Lys	c.5395G>A	COSM180789
MTOR	40	p.Phe1888Val	c.5662T>G	COSM893814
MTOR	40	p.Phe1888Leu	c.5662T>C	COSM3358967
MTOR	40	p.Phe1888Ile	c.5662T>A	COSM3358968
MTOR	40	p.Phe1888Leu	c.5664C>G	COSM462604
MTOR	40	p.Phe1888Leu	c.5664C>A	COSM893813

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
MTOR	43	p.Thr1977Ser	c.5929A>T	COSM1289945 Note: Some "no calls" were observed for this analytical variant due to strand bias with plasmid targets. This does not impact clinical test results.
MTOR	43	p.Thr1977Lys	c.5930C>A	COSM462601
MTOR	43	p.Thr1977Arg	c.5930C>G	COSM462602 Note: Some "no calls" were observed for this analytical variant due to strand bias with plasmid targets. This does not impact clinical test results.
MTOR	43	p.Val2006Leu	c.6016G>C	COSM1134662
MTOR	43	p.Val2006Ile	c.6016G>A	COSM893804
MTOR	43	p.Val2006Phe	c.6016G>T	COSM249481
MTOR	47	p.Ser2215Pro	c.6643T>C	COSM1560108
MTOR	47	p.Ser2215Tyr	c.6644C>A	COSM20417
MTOR	47	p.Ser2215Phe	c.6644C>T	COSM1686998
MTOR	53	p.Leu2427Gln	c.7280T>A	COSM1185313
MTOR	53	p.Leu2427Arg	c.7280T>G	COSM2119114
MTOR	30	p.Cys1483Arg	c.4447T>C	COSM3747775
NRAS	2	p.Gly12Arg	c.34G>C	COSM561
NRAS	2	p.Gly12Ser	c.34G>A	COSM563
NRAS	2	p.Gly12Ala	c.35G>C	COSM565
NRAS	2	p.Gly12Asp	c.35G>A	COSM564
NRAS	2	p.Gly12Val	c.35G>T	COSM566
NRAS	2	p.Gly13Arg	c.37G>C	COSM569
NRAS	2	p.Gly13Ser	c.37G>A	COSM571
NRAS	2	p.Gly13Cys	c.37G>T	COSM570

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
NRAS	2	p.Gly13Ala	c.38G>C	COSM575
NRAS	2	p.Gly13Asp	c.38G>A	COSM573
NRAS	2	p.Gly13Val	c.38G>T	COSM574
NRAS	3	p.Ala59Thr	c.175G>A	COSM578
NRAS	3	p.Gln61Lys	c.181C>A	COSM580
NRAS	3	p.Gln61Glu	c.181C>G	COSM581
NRAS	3	p.Gln61Arg	c.182A>G	COSM584
NRAS	3	p.Gln61Pro	c.182A>C	COSM582
NRAS	3	p.Gln61Leu	c.182A>T	COSM583
NRAS	3	p.Gln61His	c.183A>C	COSM586
NRAS	3	p.Gln61His	c.183A>T	COSM585
NRAS	4	p.Lys117Asn	c.351G>T	MAN13
NRAS	4	p.Ala146Thr	c.436G>A	COSM27174
NRAS	4	p.Ala146Val	c.437C>T	COSM4170228
NRAS	2	p.Gly12Cys	c.34G>T	COSM562
PDGFRA	12	p.Val561Asp	c.1682T>A	COSM739
PDGFRA	14	p.Asn659Tyr	c.1975A>T	COSM22416
PDGFRA	14	p.Asn659Lys	c.1977C>A	COSM22415
PDGFRA	14	p.Asn659Lys	c.1977C>G	COSM22414
PDGFRA	18	p.Asp842_His845del	c.2526_2537delCATCATGCATGA	COSM737
PDGFRA	18	p.Asp842_Met844del	c.2524_2532delGACATCATG	COSM12401
PDGFRA	18	p.Asp842Tyr	c.2524G>T	COSM12396
PDGFRA	18	p.Asp842Val	c.2525A>T	COSM736
PDGFRA	18	p.Ile843_Asp846del	c.2527_2538delATCATGCATGAT	COSM12400
PDGFRA	18	p.Ile843_Ser847delinsThr	c.2528_2539delTCATGCATGAT	COSM12407
PIK3CA	2	p.Arg38Ser	c.112C>A	COSM87310

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
PIK3CA	2	p.Arg38Gly	c.112C>G	COSM40945
PIK3CA	2	p.Arg38Cys	c.112C>T	COSM744
PIK3CA	2	p.Arg38His	c.113G>A	COSM745
PIK3CA	2	p.Glu39Lys	c.115G>A	COSM30625
PIK3CA	2	p.Glu81Lys	c.241G>A	COSM27502
PIK3CA	2	p.Arg88Gln	c.263G>A	COSM746
PIK3CA	2	p.Arg93Trp	c.277C>T	COSM27493
PIK3CA	2	p.Arg93Gln	c.278G>A	COSM86041
PIK3CA	2	p.Gly106Val	c.317G>T	COSM748
PIK3CA	2	p.Arg108His	c.323G>A	COSM27497
PIK3CA	2	p.Lys111Glu	c.331A>G	COSM13570
PIK3CA	5	p.Val344Ala	c.1031T>C	COSM86951
PIK3CA	5	p.Val344Gly	c.1031T>G	COSM22540
PIK3CA	5	p.Asn345Ile	c.1034A>T	COSM94978
PIK3CA	5	p.Asn345Lys	c.1035T>A	COSM754
PIK3CA	6	p.Glu365Lys	c.1093G>A	COSM86044
PIK3CA	6	p.Glu365Gly	c.1094A>G	COSM1420797
PIK3CA	6	p.Glu365Val	c.1094A>T	COSM1484860
PIK3CA	6	p.Cys378Arg	c.1132T>C	COSM756
PIK3CA	6	p.Cys378Tyr	c.1133G>A	COSM1041478
PIK3CA	6	p.Cys378Phe	c.1133G>T	COSM21450
PIK3CA	8	p.Cys420Arg	c.1258T>C	COSM757
PIK3CA	10	p.Pro539Arg	c.1616C>G	COSM759
PIK3CA	10	p.Glu542Lys	c.1624G>A	COSM760
PIK3CA	10	p.Glu542Val	c.1625A>T	COSM762
PIK3CA	10	p.Glu545Lys	c.1633G>A	COSM763
PIK3CA	10	p.Glu545Gln	c.1633G>C	COSM27133
PIK3CA	10	p.Glu545Ala	c.1634A>C	COSM12458



(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
PIK3CA	10	p.Glu545Gly	c.1634A>G	COSM764
PIK3CA	10	p.Glu545Asp	c.1635G>C	COSM27374
PIK3CA	10	p.Glu545Asp	c.1635G>T	COSM765
PIK3CA	10	p.Gln546Lys	c.1636C>A	COSM766
PIK3CA	10	p.Gln546Glu	c.1636C>G	COSM6147
PIK3CA	10	p.Gln546Pro	c.1637A>C	COSM767
PIK3CA	10	p.Gln546Arg	c.1637A>G	COSM12459
PIK3CA	10	p.Glu547Lys	c.1639G>A	COSM29315
PIK3CA	14	p.His701Pro	c.2102A>C	COSM778
PIK3CA	14	p.His701Arg	c.2102A>G	COSM1420881
PIK3CA	14	p.Glu726Lys	c.2176G>A	COSM87306
PIK3CA	14	p.Glu726Gly	c.2177A>G	COSM1420887
PIK3CA	19	p.Cys901Arg	c.2701T>C	COSM1420899
PIK3CA	19	p.Cys901Tyr	c.2702G>A	COSM1420901
PIK3CA	19	p.Cys901Phe	c.2702G>T	COSM769
PIK3CA	21	p.Tyr1021Cys	c.3062A>G	COSM12461
PIK3CA	21	p.Thr1025Ala	c.3073A>G	COSM771
PIK3CA	21	p.Met1043Val	c.3127A>G	COSM12591
PIK3CA	21	p.Met1043Ile	c.3129G>A	COSM29313
PIK3CA	21	p.Met1043Ile	c.3129G>T	COSM773
PIK3CA	21	p.Asn1044Lys	c.3132T>A	COSM12592
PIK3CA	21	p.His1047Tyr	c.3139C>T	COSM774
PIK3CA	21	p.His1047Arg	c.3140A>G	COSM775
PIK3CA	21	p.His1047Leu	c.3140A>T	COSM776
PIK3CA	21	p.Gly1049Ser	c.3145G>A	COSM777
PIK3CA	21	p.Gly1049Arg	c.3145G>C	COSM12597
RAF1	7	p.Ser257Leu	c.770C>T	COSM181063
RAF1	7	p.Ser257Trp	c.770C>G	COSM581519

(continued)

Gene	Exon	Amino acid change	Nucleotide change	Variant ID
RAF1	12	p.Thr421Met	c.1262_1263delCCinsTG	MAN9
RET	10	p.Cys618Arg	c.1852T>C	COSM29803
RET	10	p.Cys618Tyr	c.1853G>A	COSM980
RET	10	p.Cys620Arg	c.1858T>C	COSM29804
RET	11	p.Cys634Arg	c.1900T>C	COSM966
RET	13	p.Glu768Gly	c.2303A>G	COSM1347811
RET	13	p.Glu768Asp	c.2304G>C	COSM21338
RET	15	p.Ala883Phe	c.2646_2648delAGCinsTTT	COSM981 Note: The nucleotide change of COSM981 overlaps that of COSM133167, so a positive COSM981 sample will also result in a positive call for COSM133167.
RET	15	p.Ala883Ser	c.2647G>T	COSM133167
RET	15	p.Asp898_Glu901del	c.2694_2705delTGTTTATGAAG A	COSM962
RET	16	p.Met918Thr	c.2753T>C	COSM965
ROS1	36	p.Leu1951Met	c.5851C>A	COSM1072521
ROS1	38	p.Gly2032Arg	c.6094G>C	MAN11
ROS1	38	p.Gly2032Arg	c.6094G>A	MAN10

[1] Annotations for this variant are not available for reporting.

Fusion isoforms detected in non-small cell lung cancer (NSCLC)

RET fusion isoforms (N=40)	
ACBD5-RET.A11R12	KIF5B-RET.K15R11.COSF1255.1
AFAP1-RET.A3R12	KIF5B-RET.K15R12.COSF1232
AKAP13-RET.A35R12	KIF5B-RET.K16R12.COSF1230
AKAP13-RET.A36R12	KIF5B-RET.K22R12.COSF1253
CCDC6-RET.C1R11	KIF5B-RET.K23R12.COSF1234
CCDC6-RET.C1R11.1	KIF5B-RET.K24R11.COSF1262
CCDC6-RET.C1R12	KIF5B-RET.K24R8.COSF1236
CCDC6-RET.C1R12.COSF1271	KTN1-RET.K29R12.COSF1513
CCDC6-RET.C2R12	NCOA4-RET.N7R12
CCDC6-RET.C8R11	NCOA4_ELE1-RET.E6R12
CCDC6-RET.C8R12full	PCM1-RET.P29R12
CUX1-RET.C10R12	PRKAR1A-RET.P7R12
ERC1-RET.E12R12	RUFY2-RET.R9R12
ERC1-RET.E17R12	SPECC1L-RET.S10R11.NGS.1
ERC1-RET.E7R12	SPECC1L-RET.S10R12
ERC1_ELKS-RET.E11R12.COSF1507	TBL1XR1-RET.T9R11.NGS.1
FKBP15-RET.F25R12	TBL1XR1-RET.T9R12
GOLGA5-RET.G7R12	TRIM24-RET.T9R12
HOOK3-RET.H11R12	TRIM27-RET.T3R12
KIAA1468-RET.K10R12	TRIM33-RET.T16R12

ROS1 fusion isoforms (N=34)	
CCDC6-ROS1.C5R35	LRIG3-ROS1.L16R35.COSF1269
CD74-ROS1.C4R33.NGS	MSN-ROS1.M9R34
CD74-ROS1.C6R32.COSF1202	MYO5A-ROS1.M23R35
CD74-ROS1.C6R34.COSF1200	PPFIBP1-ROS1.P9R35
CD74-ROS1.C6R35	PWWP2A-ROS1.P1R36
CD74-ROS1.C7R34	SDC4-ROS1.S2R32.COSF1265
CEP85L-ROS1.C8R36	SDC4-ROS1.S2R34
CLIP1-ROS1.C19R36	SDC4-ROS1.S4R32.COSF1278
CLTC-ROS1.C31R35	SDC4-ROS1.S4R34.COSF1280
ERC1-ROS1.E11R36	SLC34A2-ROS1.S13R32.COSF1259
EZR-ROS1.E10R34.COSF1267	SLC34A2-ROS1.S13R34.COSF1261
EZR-ROS1.E10R35	SLC34A2-ROS1.S4R32.COSF1197
GOPC-ROS1.G4R36.COSF1188	SLC34A2-ROS1.S4R34.COSF1198
GOPC-ROS1.G8R35.COSF1139	TFG-ROS1.T4R35
HLA_A-ROS1.H7R34	TPM3-ROS1.T3R36
KDEL2-ROS1.K5R35	TPM3-ROS1.T7R35.COSF1273
KIAA1598-ROS1.K11R36	ZCCHC8-ROS1.Z2R36

DNA variants detected in cholangiocarcinoma (CC)

Gene	Amino Acid Change	Nucleotide Change	Variant ID
IDH1	p.Arg132Cys	c.394C>T	COSM28747
IDH1	p.Arg132Ser	c.394C>A	COSM28748
IDH1	p.Arg132Gly	c.394C>G	COSM28749
IDH1	p.Arg132Leu	c.395G>T	COSM28750
IDH1	p.Arg132His	c.395G>A	COSM28746



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Obtaining Certificates of Analysis

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Obtaining Certificates of Conformance

The Certificate of Conformance provides information on conformance testing of each instrument provided with the system. Certificates of Conformance are shipped with the instrument, and are also available by contacting Customer Support at [thermofisher.com/support](https://www.thermofisher.com/support).

Oncomine™ Dx Target Test Part II: Sample and Library Preparation

USER GUIDE

Publication Number MAN0018704

Revision B.0

IVD

For In Vitro Diagnostic Use.

ThermoFisher
S C I E N T I F I C



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Products manufactured in Singapore:

Ion PGM™ Dx Instrument System
Ion PGM™ Dx Sequencer
Ion OneTouch™ Dx Instrument
Ion OneTouch™ ES Dx Instrument
Ion PGM™ Dx Chip Minifuge (120V)
Ion PGM™ Wireless Scanner
Ion Torrent™ Server
Torrent Suite™ Dx Software
Veriti™ Dx 96-well Thermal Cycler, 0.2 mL

Products manufactured in Frederick:

Oncomine™ Dx Target Test Kit
Ion Torrent Dx FFPE Sample Preparation Kit
Ion PGM™ Dx Library Kit
Ion OneTouch™ Dx Template Kit
Ion PGM™ Dx Sequencing Kit
Ion 318™ Dx Chip
Ion OneTouch™ Rack Kit
DynaMag™ Dx 96-Well Plate Magnet
DynaMag™ Dx 16 2-mL Magnet

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Revision history: Pub. No. MAN0018704

Revision	Date	Description
B.0	3 September 2021	Updated with new product catalog numbers.
A.0	11 February 2021	New Oncomine™ Dx Target Test user guide for FDA submission.

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About this guide

IMPORTANT! Before using this product, read and understand the information in the “Safety” appendix in this document.

Purpose of this guide

This user guide provides instructions for sample preparation, sample quantification, and library preparation using the Oncomine™ Dx Target Test. The resulting libraries are ready for template preparation and sequencing on the Ion PGM™ Dx System.

Oncomine™ Dx Target Test Kit user guides

This user guide is part of a five-guide set.

- *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*
- *Oncomine™ Dx Target Test Part II: Sample and Library Preparation User Guide*
- *Oncomine™ Dx Target Test Part III: Template Preparation User Guide*
- *Oncomine™ Dx Target Test Part IV: Sequencing and Results Reports User Guide*
- *Oncomine™ Dx Target Test Part V: Torrent Suite™ Dx Software 5.12.5 Reference User Guide*

All five guides are required to complete the entire Oncomine™ Dx Target Test workflow.

Note: The procedures in these guides supersede the instructions in the *Ion PGM™ Dx System User Guide* when using the Ion PGM™ Dx System with the Oncomine™ Dx Target Test.



Product information

Product description

Oncomine™ Dx Target Test

The Oncomine™ Dx Target Test is an *in vitro* diagnostic next-generation sequencing test to detect somatic alterations in human DNA and RNA isolated from formalin-fixed, paraffin-embedded (FFPE) tissue samples. Detection of these variants is performed using the Ion PGM™ Dx System.

For a complete product description of the Oncomine™ Dx Target Test, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Sample preparation components

The Ion Torrent Dx FFPE Sample Preparation Kit, included as part of the Oncomine™ Dx Target Test Kit, provides the following components for isolating and quantifying DNA and RNA from FFPE tissue samples that are mounted on slides:

- Ion Torrent Dx Total Nucleic Acid Isolation Kit, for extracting and isolating DNA and RNA from FFPE tissue samples
- Ion Torrent Dx DNA Quantification Kit, for quantifying DNA using a fluorometer/fluorescence reader
- Ion Torrent Dx RNA Quantification Kit, for quantifying RNA using a fluorometer/fluorescence reader

The Ion Torrent Dx cDNA Synthesis Kit and the Oncomine™ Dx Target Test, Controls, and Diluent Kit are used to reverse transcribe the quantified RNA into cDNA.

Library preparation components

The Oncomine™ Dx Target Test Kit includes the following components for preparing barcoded libraries from DNA and cDNA for sequencing on the Ion PGM™ Dx System:

- Oncomine™ Dx Target Test, Controls, and Diluent Kit, which includes primer panels for amplifying DNA- and RNA-specific target regions as well as controls
- Ion PGM™ Dx Library Kit, which includes 16 unique barcode adapters (BC 1–BC 16) as well as enzymes and other reagents for library preparation
- Ion PGM™ Dx Library Equalizer™ Reagents, to normalize the concentration of the resulting libraries to ~100 pM without the need for quantification

The library preparation procedure requires 10 ng of DNA and RNA.

Intended use

For the Intended Use statement for the Oncomine™ Dx Target Test, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Theory of operation

For a complete description of the Theory of Operation of the system, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Software compatibility and requirements

The procedures in this guide are designed for use with Torrent Suite™ Dx Software version 5.12.5 or later. For a complete description of software compatibility and requirements, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Materials provided

Oncomine™ Dx Target Test Kit

The Oncomine™ Dx Target Test Kit (Cat. No. A51695) includes the following subkits.

IMPORTANT! Refer to the product label for the storage conditions and expiration dates of individual modules and components.

✓	Subkit	Part No.
	Oncomine™ Dx Target Test, Controls, and Diluent Kit	A49756
	Ion Torrent Dx FFPE Sample Preparation Kit	A32445
	Ion PGM™ Dx Library Kit	A49758
	Ion OneTouch™ Dx Template Kit	A49759
	Ion PGM™ Dx Sequencing Kit	A49760
	Ion 318™ Dx Chip Kit	A18937
	Oncomine™ Dx Target Test User Guides and Assay Definition File	A51694

Subkits used in this guide

The procedures in this user guide use the following subkits from the OncoPrint™ Dx Target Test Kit.

Ion Torrent Dx FFPE Sample Preparation Kit

The Ion Torrent Dx FFPE Sample Preparation Kit (Cat. No. A32445) provides reagents for nucleic acid isolation, quantification, and cDNA synthesis from FFPE samples.

✓	Component	Amount	Storage
Ion Torrent Dx Total Nucleic Acid Isolation Kit box 1 of 2 (36 reactions; Part No. A32434)			
	10X DNase Buffer (white cap)	6 × 46 µL	-30°C to -10°C
	Protease (blue cap)	6 × 34 µL	
	DNase (purple cap)	6 × 34 µL	
Ion Torrent Dx Total Nucleic Acid Isolation Kit box 2 of 2 (36 reactions; Part No. A32435)			
	Elution Solution (red cap)	6 × 900 µL	15°C to 30°C
	Isolation Additive (brown cap)	6 × 1 mL	
	Wash 1 Concentrate (amber cap)	6 × 5.9 mL	
	Wash 2 Concentrate (clear cap)	6 × 3.4 mL	
	Digestion Buffer (green cap)	6 × 200 µL	
	Dilution Solution (black cap)	6 × 1.5 mL	
	Collection Tubes	6 × 6 tubes	
	Filter Cartridges ^[1]	6 × 12 cartridges	
	Low-bind Elution Tubes	6 × 14 tubes	
Ion Torrent Dx DNA Quantification Kit (72 reactions; Part No. A32437)			
	DNA Dye Reagent (blue cap)	6 × 70 µL	2°C to 8°C
	DNA Buffer (white cap)	6 × 14.3 mL	
	DNA Std - 0 ng/µL (white cap)	6 × 150 µL	
	DNA Std - 0.5 ng/µL (green cap)	6 × 150 µL	
	DNA Std - 4 ng/µL (red cap)	6 × 150 µL	
	DNA Std - 10 ng/µL (yellow cap)	6 × 150 µL	

✓	Component	Amount	Storage
Ion Torrent Dx RNA Quantification Kit (72 reactions; Part No. A32438)			
	RNA Dye Reagent (green cap)	6 × 70 µL	2°C to 8°C
	RNA Buffer (blue cap)	6 × 14.3 mL	
	RNA Std - 0 ng/µL (teal cap)	6 × 150 µL	
	RNA Std - 0.5 ng/µL (tan cap)	6 × 150 µL	
	RNA Std - 4 ng/µL (purple cap)	6 × 150 µL	
	RNA Std - 10 ng/µL (orange cap)	6 × 150 µL	
Ion Torrent Dx cDNA Synthesis Kit (48 reactions; Part No. A32436)			
	10X Enzyme Mix (green cap)	6 × 13 µL	-30°C to -10°C
	5X Reaction Mix (red cap)	6 × 22 µL	
Ion Torrent Dx Sample Dilution Kit (Part No. A32439)			
	Dilution Solution (black cap)	8 × 1.5 mL	15°C to 30°C

[1] Includes a filter column pre-inserted in a Collection Tube.

Oncomine™ Dx Target Test, Controls, and Diluent Kit

The Oncomine™ Dx Target Test, Controls, and Diluent Kit (Part No. A49756) provides the following panels and controls.

✓	Component	Amount	Storage
Oncomine™ Dx Target Test DNA and RNA Panel (Part No. A32441)			
	Oncomine™ Dx Target Test—DNA panel (blue cap)	6 × 32 µL	-30°C to -10°C
	Oncomine™ Dx Target Test—RNA panel (yellow cap)	6 × 32 µL	
Oncomine™ Dx Target DNA Control (Part No. A44913)			
	Oncomine™ Dx Target DNA Control (brown cap)	8 × 7 µL (single-use tubes)	-30°C to -10°C
Oncomine™ Dx Target RNA Control (Part No. A32443)			
	Oncomine™ Dx Target RNA Control (white cap)	8 × 7 µL (single-use tubes)	-90°C to -60°C
Oncomine™ Dx Target RNA Control Diluent (Part No. A38872)			
	Oncomine™ Dx Target RNA Control Diluent (blue cap)	8 × 88 µL (single-use tubes)	-90°C to -60°C

✓	Component	Amount	Storage
Ion Torrent Dx No Template Control Kit (Part No. A32444)			
	No Template Control (purple cap)	8 × 30 µL	15°C to 30°C

Ion PGM™ Dx Library Kit

The Ion PGM™ Dx Library Kit (Cat. No. A49758) provides reagents for preparing up to 96 sample libraries.

IMPORTANT! Do not mix components from other library kits.

✓	Component	Amount	Storage
Ion PGM™ Dx Library Reagents (Part No. A18928)			
	LIB HiFi Mix (red cap)	6 × 252 µL	-30°C to -10°C
	LIB FuPa (green cap)	6 × 32 µL	
	LIB Switch Soln (orange cap)	6 × 64 µL	
	LIB DNA Ligase (clear cap)	6 × 32 µL	
	BC 1 through BC 16 (16 unique barcode adapters, numbered 1–16, white cap)	16 × 12 µL	
Ion PGM™ Dx Library Equalizer™ Reagents (Part No. A18929)			
	LIB AMPure™ Reagent (clear cap)	4.4 mL	2°C to 8°C
	LIB Beads (yellow cap)	6 × 48 µL	
	LIB Primers (blue cap)	6 × 36 µL	
	LIB Capture (violet cap)	6 × 160 µL	
	LIB Wash Soln (clear cap)	30 mL	
	LIB Elution Soln (clear cap)	9.6 mL	

Materials and equipment required but not provided

Unless otherwise indicated, all materials are available through thermofisher.com. "MLS" indicates that the material is available from fisherscientific.com or another major laboratory supplier.

Description	Source
Veriti™ Dx 96-well Thermal Cycler, 0.2 mL	4452300
Laminar flow hood	MLS
Dry-bath heaters and aluminum heat blocks (quantity = 3), for use with 1.5-mL tubes	MLS
1.5-mL snap-cap low-retention polypropylene microcentrifuge tubes	MLS
1.5-mL tube rack	MLS
Aluminum cold blocks for use with 96-well plates	MLS
Benchtop cold box for use with 1.5-mL tubes	MLS
Microcentrifuge (must accommodate standard 1.5-mL and 0.2-mL microcentrifuge tubes, and generate 20,000 rcf)	MLS
0.2-mL tube adapters	MLS
Mini centrifuge	MLS
96-well plate centrifuge	MLS
Vortex mixer with a rubber platform	MLS
Fluorometer/fluorescence reader (see additional specifications following)	MLS
Tubes or plates for the fluorometer/fluorescence reader	MLS
DynaMag™ Dx 96-Well Plate Magnet magnet	A31347
DynaMag™ Dx 16 2-mL Magnet	A31346
Slide rack, able to hold standard 3" × 1" (75 × 25 mm) slides	MLS
Staining dish or jar, able to hold sufficient liquid to fully submerge the slide rack	MLS
Disposable scalpel with a sterile #10 blade	MLS
RNase decontamination solution	MLS
Absolute ethanol (ACS grade)	MLS
Xylene (ACS grade, ≥98.5%)	MLS
Nuclease-free water	MLS
Single- and multi-channel pipettes (2-, 20-, 200-, 1000-µL)	MLS

(continued)

Description	Source
Aerosol-barrier pipette tips (2-, 10-, 20-, 200-, 1000- μ L)	MLS
Troughs for multi-channel pipettors	MLS
MicroAmp™ Optical 96-well Reaction Plates	4481191 4481192 (with barcode)
Adhesive PCR Plate Seals	AB0558
15-mL and 50-mL conical tubes and tube holders (for preparing bulk solutions)	MLS
5-mL and 25-mL serological pipettes, and pipette controller (for preparing bulk solutions)	MLS

Fluorometer/ fluorescence reader specifications

For the DNA and RNA quantification procedure, you can use any qualified fluorometer/fluorescence reader that can accommodate the use of a 2–4-point standard curve and is able to operate at the excitation and emission wavelengths listed below:

Dye reagent	Excitation (nm)	Emission (nm)
RNA Dye Reagent	620/15	680/30
DNA Dye Reagent	485/20	528/20

DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet

Note: Do not substitute non-IVD labeled magnets for the DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet.

The DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet, provided with Ion PGM™ Dx System, contain high-energy neodymium magnets and are used as part of the procedure for purifying sample libraries bound to LIB AMPure™ Reagent and LIB Beads. The DynaMag™ Dx 16 2-mL Magnet is also used to prepare TMPL ES Beads as part of template preparation.

The DynaMag™ Dx 96-Well Plate Magnet has 7 bar magnets with a hard plastic top to fit 96-well PCR plates. When you insert a plate, the magnets collect bead-bound biomolecules in suspension at the sides of the plate wells, allowing removal of fluid without disturbing the bead pellets. An extra column in the magnet enables sample mixing by shifting the plate back and forth in the magnet.

The DynaMag™ Dx 16 2-mL Magnet holds 16 standard 1.5-mL or 2-mL microcentrifuge tubes, and collects bead-bound biomolecules in suspension at the sides of the tubes, allowing removal of fluid without disturbing the bead pellets.

Do not use the magnets above 50°C (122°F) and store in a cool, dry environment.



Before you begin

Tissue input requirements for FFPE sample extraction

The starting material for the extraction procedure is an FFPE tissue sample that is unstained and mounted on a slide. Confirm the tumor content of each sample based on the area of a hematoxylin and eosin (H&E) stained section.

The recommended number of slide-mounted 5-micron FFPE sections used in extraction varies depending on the sample collection method:

Sample collection method	Recommended number of sections
Non-small cell lung cancer (NSCLC)	
Resection or surgical biopsies	2 × 5-micron sections
Core needle biopsies	9 × 5-micron sections
Fine needle aspirates	7 × 5-micron sections
Cholangiocarcinoma (CC)	
Resection or surgical biopsies	4 × 5-micron sections
Core needle biopsies	9 × 5-micron sections

Note: Extraction from FFPE sample curls has not been evaluated.

Sample storage and stability

Store FFPE blocks and slides at room temperature (15–30°C).

Blocks and slides (paraffin-dipped or undipped) are stable for up to 5 months at 15–30°C.

Extracted DNA can be stored at –30°C to –10°C for up to 12 months, including 3 freeze-thaw cycles.

Extracted RNA can be stored at –90°C to –60°C for up to 5 months, including 3 freeze-thaw cycles.

Procedural guidelines

Definitions

Throughout this guide:

- Room temperature is defined as the temperature range 15–30°C (59–86°F).
- A pulse centrifugation consists of a 3–5 second centrifugation at maximum speed in a mini centrifuge.

Guidelines to prevent cross-contamination



CAUTION! A primary source of contamination is nucleic acid from previous sample processing steps. Do not introduce amplified DNA into the target amplification preparation area.

- When designing the laboratory layout, dedicate separate areas for pre- and post-amplification activities. Dedicate laboratory supplies and/or equipment to the appropriate area.
- Use a laminar flow hood in the dedicated pre-amplification area for target amplification reaction setup.
- Before and after use, clean all surfaces and equipment in the laminar flow hood with 10% bleach followed by two water rinses.
- Turn on the UV light in the hood for 10 minutes before and after use.
- Use fresh gloves before entering the hood.
- Change tips between pipetting steps.
- Prepare a waste container containing 10% bleach solution for disposing of used tips after pipetting libraries.

Reagent contamination

Before use, verify that any nuclease-free water used in the procedure is not cloudy, a potential indication of contamination. If the water is cloudy, use a different vial.

Guidelines for FFPE samples

- For core needle biopsies, macrodissection is not recommended due to the limiting tissue section surface areas.
- For resection or surgical biopsies, macrodissect and enrich the sample for tumor content if the tumor content is less than 20% and the tumor content in the region of interest is greater than or equal to 10%. Following tumor enrichment, proceed with the extraction protocol.
- Necrotic samples: 10–20% necrotic tissue in the region of interest does not appear to interfere with the assay. However, we recommend that you macrodissect highly necrotic areas or select alternate samples if possible.
- Nucleic acid integrity is important for sample performance. Factors such as age of the block, fixation process used, and sample source can impact the quality of the extracted nucleic acid.
- Nucleic acid yield can be impacted by overall tissue area. Should an initial extraction lead to insufficient concentrations for DNA and RNA, repeat the extractions with more material whenever possible.

Guidelines for RNA

- Wear clean gloves and a clean lab coat.
- Change gloves whenever they may be contaminated.
- Open and close all sample tubes carefully. Avoid splashing or spraying samples.
- Clean lab benches and equipment (including gloves, tube racks, pipettes, centrifuges, and vortexers) with an RNase decontamination solution before and after use.
- Work in a designated RNase-free pre-PCR area.
- Keep RNA on ice or in a –30°C to –10°C chilled benchtop cold box during use.
- Never vortex RNA. Flick 4 times to mix, then pulse centrifuge to collect.

Guidelines for mixing reagents

Immediately before each use:

- Mix enzyme solutions (e.g., Protease and DNase) by flicking the tubes 4 times, followed by a pulse centrifugation.
- Vortex non-enzyme-containing reagents for ~5 seconds, followed by a pulse centrifugation.
- Mix reagent bottles by inverting them 5 times.

Guidelines for pipetting

- Use aerosol-barrier pipette tips. Change pipette tips between samples.
- Avoid introducing air bubbles when pipetting by keeping the pipette tip at the bottom of the solution in the wells.
- Set the pipette to the recommended volume for mixing, and insert tip into the solution with the pipette plunger depressed to avoid introducing air bubbles.
- Visually inspect multi-channel pipette tips to ensure volumes are equivalent during pipetting.

- Touch tip to the side of well and slowly pipet reagent on the side of the well to form a droplet. This enables small volumes to be pipetted accurately and to ensure that the reagent has been added to the well.
- Inspect the pipette tips to verify that the reagent has been adequately dispensed.

Guidelines for freezing and thawing samples

There are stopping points throughout this procedure where you can freeze samples overnight or longer and then thaw the samples before proceeding. If you cannot perform the complete procedure in a day, proceed to a designated stopping point and freeze the samples overnight.

IMPORTANT! Freeze-thaw samples no more than 3 times.

Guidelines for library preparation

- Up to 16 barcode adapters may be used in a single sequencing run.
- Freeze-thaw barcode adapters no more than 6 times.
- Verify that the correct program is selected before starting the Veriti™ Dx program.
- To avoid cross-contamination between samples, skip wells or columns when setting up reactions in a 96-well plate. Circle wells that are used with ethanol-resistant marker to help indicate where the samples are located.
- Because cDNA and DNA amplification reactions require a different number of cycles, they must be set up and run on separate 96-well plates.

Equilibrate materials

Equilibrate the following materials for at least 24 hours before use:

- Equilibrate two 96-well aluminum cold blocks to 2–8°C in a refrigerator.
- Equilibrate a benchtop cold box to –30°C to –10°C in a freezer.
- Equilibrate a separate benchtop cold box to 2–8°C in a refrigerator, or use ice to keep reaction tubes chilled on the bench.

Note: A cold box holds temperature for up to 1 hour on the bench.

Reagent management

Follow the guidelines below for proper reagent storage and use.

Storage

Reagents must be stored under appropriate conditions. Refer to the Product Information section in each user guide for the storage conditions of the kit components used in the procedures in that guide. The Oncomine™ Dx Target Test Kit includes kits with multiple component boxes that require different storage conditions. For example, the Oncomine™ Dx Target Test, Controls, and Diluent Kit includes four boxes, which are stored at different temperatures. To use the Oncomine™ Dx Target Test, Controls, and Diluent Kit, retrieve all boxes from their different storage areas and confirm that they are from the same master lot.

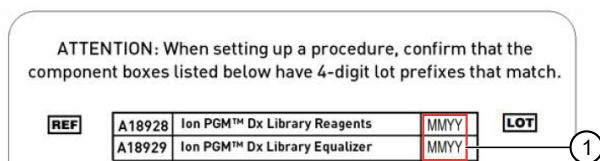
Kit interchangeability and component box lot matching

The top-level kits used for sample preparation, library preparation, template preparation, and sequencing can be mixed and matched. For example, an Ion PGM™ Dx Library Kit can be used with any Ion PGM™ Dx Sequencing Kit. However, the component boxes in a particular kit must be lot-matched with the other boxes in that kit.

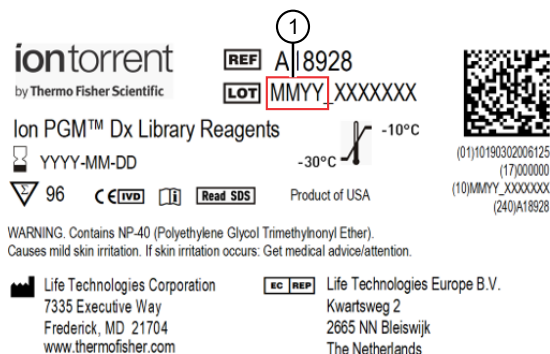
Note: The controls and diluent in the Oncomine™ Dx Target Test, Controls, and Diluent Kit must be lot-matched. However, the controls do not require lot-matching with the Oncomine™ Dx Target Test DNA and RNA Panel.

Each component box lists the 4-digit lot prefixes of the compatible component boxes inside the box lid. Before using a particular kit, check the inside lid of each box to confirm that it is compatible with the other boxes.

An example inside box lid label is shown below:

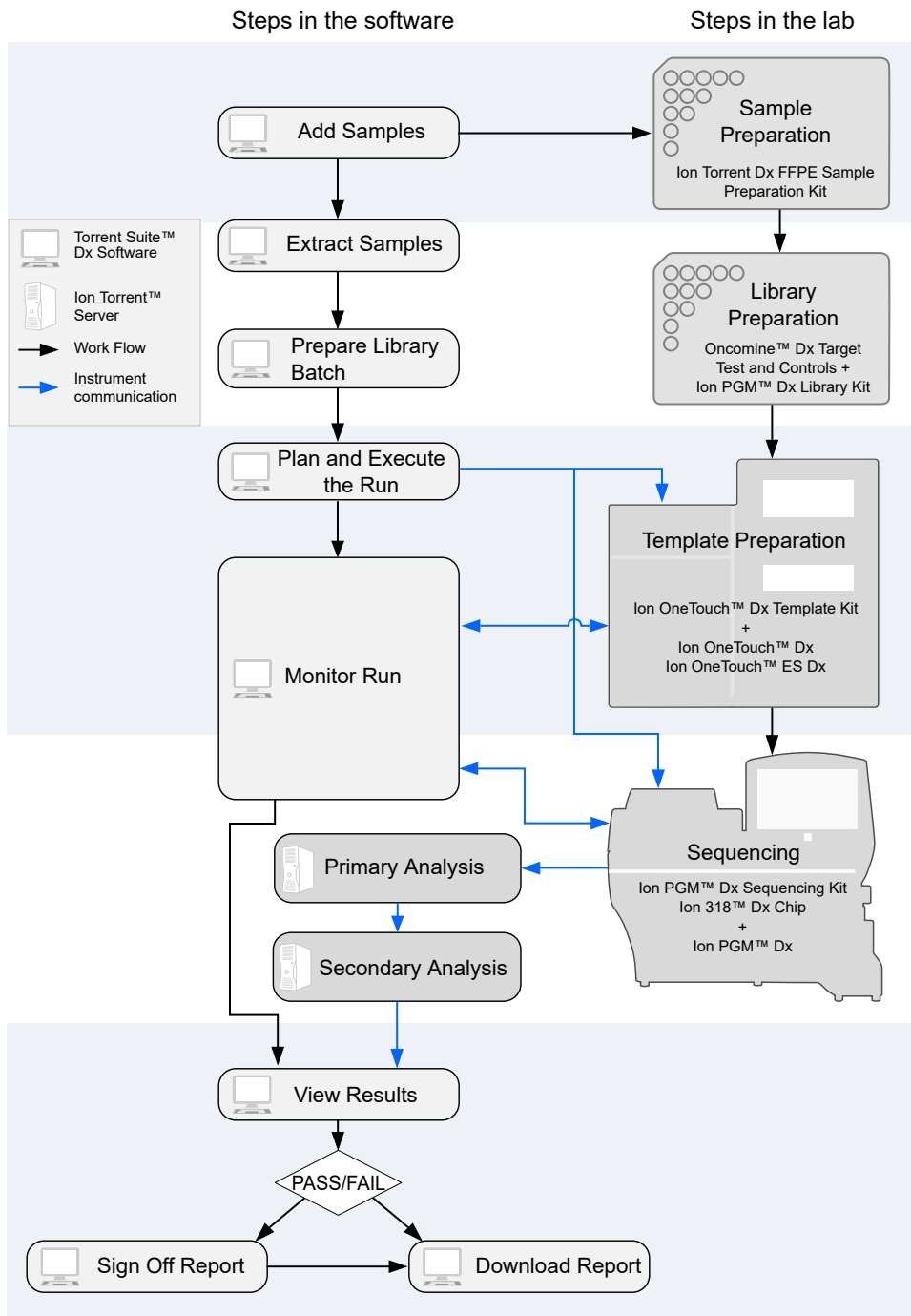


An example box label with lot information is shown below:



① Lot prefix

Oncomine™ Dx Target Test system diagram



Sample preparation workflow

The following workflow summarizes the steps for isolating DNA and RNA from FFPE tumor samples, and preparing cDNA.

Sample setup in Torrent Suite™ Dx Software (page 23)



Deparaffinize and digest samples (page 29)



Separate RNA from DNA (page 33)



Recover the RNA from the flow-through (page 36)



Recover the DNA from the Filter Cartridge (page 40)



DNA and RNA quantification (page 43)



Prepare the cDNA (page 55)



Proceed to library preparation

Library preparation workflow

“Prepare a library batch in Torrent Suite™ Dx Software” on page 60



“Prepare reagents and equipment” on page 63



“Amplify the cDNA” on page 64



“Amplify the DNA” on page 66



“Transfer the cDNA amplicons” on page 70



“Partially digest amplicons” on page 71



“Ligate barcode adapters” on page 73



“Prepare the LIB HiFi Mix plus LIB Primers mix” on page 74



“Purify the barcode-adapted library” on page 75



“Amplify the barcode-adapted library” on page 76



“Prepare the LIB Beads” on page 76



“Add LIB Capture to the amplified sample library” on page 77



“Add the LIB Beads and wash” on page 77



“Elute the library” on page 78



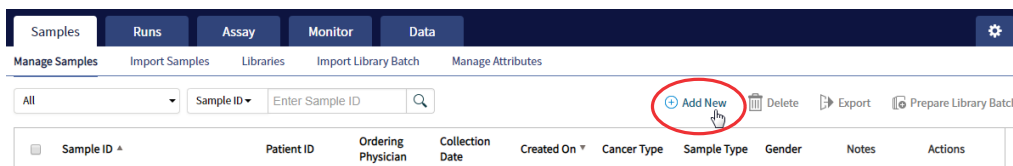
Proceed to *Oncomine™ Dx Target Test Part III: Template Preparation User Guide* (Pub. No. MAN0018705)

3



Sample setup in Torrent Suite™ Dx Software

Add a new sample

1. Under the **Samples** tab, in the **Manage Samples** screen, click **+ Add New**.



2. Complete the **Add New Sample** dialog box. Fields identified with an asterisk (*) are required. If no information is available, substitute dummy data to complete the required fields.

Field	Description
Sample ID*	A unique identifier representing the sample, containing only alphanumeric characters (0–9 and A to Z), full stops/periods (.), underscores (_), or hyphens (-). The Sample ID cannot contain spaces and is limited to a maximum of 20 characters. After a Sample ID is entered into the system, it cannot be edited. It can be deleted unless it has already been used in a library. The software checks all Sample IDs entered or imported to prevent duplication and returns an error message if a non-unique Sample ID is detected.
Patient ID*	An identifier representing the patient. This field accepts all characters including spaces.
Date of Birth*	The patient's date of birth. Click the  button to select the date in the correct format.
Ordering Physician*	The name of the ordering physician. This field accepts all characters including spaces.
Collection Date*	The date the sample was collected from the patient. Click the  button to select the date in the correct format.
Sample Source	Open-entry field that accepts all characters, including spaces. Example entry: Name of the clinic or hospital ordering the test.
Sample Condition*	Open-entry field that accepts all characters, including spaces. Example entry: FFPE surgical resection.
Sample Type*	Open-entry field that accepts all characters, including spaces. Example entry: Lung tumor, invasive adenocarcinoma.

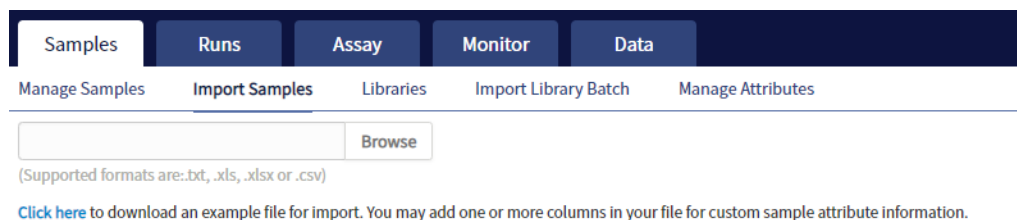
(continued)

Field	Description
Gender*	The biological gender of the sample. This must be Male, Female, or Unknown.
Cancer Type*	Select the cancer type of the sample from the dropdown list.
%Cellularity	The percentage of tumor cellularity in the sample.
%Necrosis	The percentage of cellular necrosis in the sample.
Reference Interval	A normal range of measure for the sample.
Notes	An open-entry field.

3. Click **Save**.
The sample is listed in the **Manage Samples** screen.

Import samples

Under the **Samples** tab in the **Import Samples** screen, you can import sample data in the form of a TXT, XLS, XLSX, or CSV file. The import file includes the same sample attributes that are listed in the **Add New Sample** dialog box.



1. In the **Import Samples** screen, below the **Browse** field, click **Click here** to download a Microsoft™ Excel™ template file.

Note: The template file contains default sample attributes as columns. If additional custom sample attributes have been configured in the software, add these attributes as columns to the template file.

2. In the template file, fill in the information for each sample, one sample per row. See “Predefined sample attributes” on page 25 for more information.
3. Save the file.
4. Click **Browse**, navigate to the saved file, then select it.

5. Click **Import**.

A progress bar followed by an import report displays. If the import process fails, an error message indicates the reason for failure (for example, an invalid character was used). For additional troubleshooting, see “Batch sample import fails” on page 81.

6. Click **Manage Samples** to return to the sample list. Successfully imported samples are listed.

Predefined sample attributes

The software has the following predefined sample attributes, which are listed in the **Add New Sample** dialog box and in the template file for importing samples.

- Sample ID*
- Patient ID*
- Date Of Birth*
- Ordering Physician*
- Collection Date*
- Sample Source
- Sample Condition*
- Sample Type*
- Gender*
- Cancer Type*
- %Cellularity
- %Necrosis
- Reference Interval
- Notes

*Indicates a field required to be filled in during sample creation.

Predefined attributes are locked and cannot be edited. You can create and manage custom sample attributes using the tools in the **Manage Attributes** screen.

Note: LIMS users must create custom attributes before importing sample and Planned Run information from LIMS for the attributes to be propagated through to output files. The software ignores all input file content that is not a recognized attribute.

Enter the Ion Torrent Dx Total Nucleic Acid Isolation Kit barcode

Under the **Samples** tab, in the **Manage Samples** screen, scan the barcode of the Ion Torrent Dx Total Nucleic Acid Isolation Kit used in the extraction process for a particular sample. This barcode is saved with the sample and can be viewed by clicking the Sample ID.

1. Above the samples list, select **To Be Extracted** from the **Filter Samples by...** dropdown list to display only those samples that do not have a kit barcode that is associated with them.
2. Select the checkbox of the sample to be extracted. Select multiple samples if you are using the same kit to process them.

The screenshot shows the 'Manage Samples' interface. At the top, there are tabs for 'Samples', 'Runs', 'Assay', 'Monitor', and 'Data'. Below these are sub-tabs: 'Manage Samples', 'Import Samples', 'Libraries', 'Import Library Batch', and 'Manage Attributes'. A dropdown menu is set to 'To Be Extracted'. To the right, there are buttons for 'Add New', 'Delete', 'Export', 'Extract' (circled in red), and 'Prepare Library Batch'. Below this is a table with the following data:

Sample ID ^	Patient ID	Ordering Physician	Collection Date	Created On ^	Cancer Type	Sample Type	Gender	Notes	Actions
<input checked="" type="checkbox"/> BC1	BC1	Smith	2018-09-05	2018-10-06 02:17	Non-small Cell Lung Cancer	DNA	Male		Edit Audit

At the bottom of the table, there is a pagination control showing '1' of 1 items per page and '1 - 1 of 1 Items'.

3. Click **Extract**. In the dialog box, scan the barcode that is printed on the Ion Torrent Dx Total Nucleic Acid Isolation Kit (box 1 of 2, Part No. A32434).

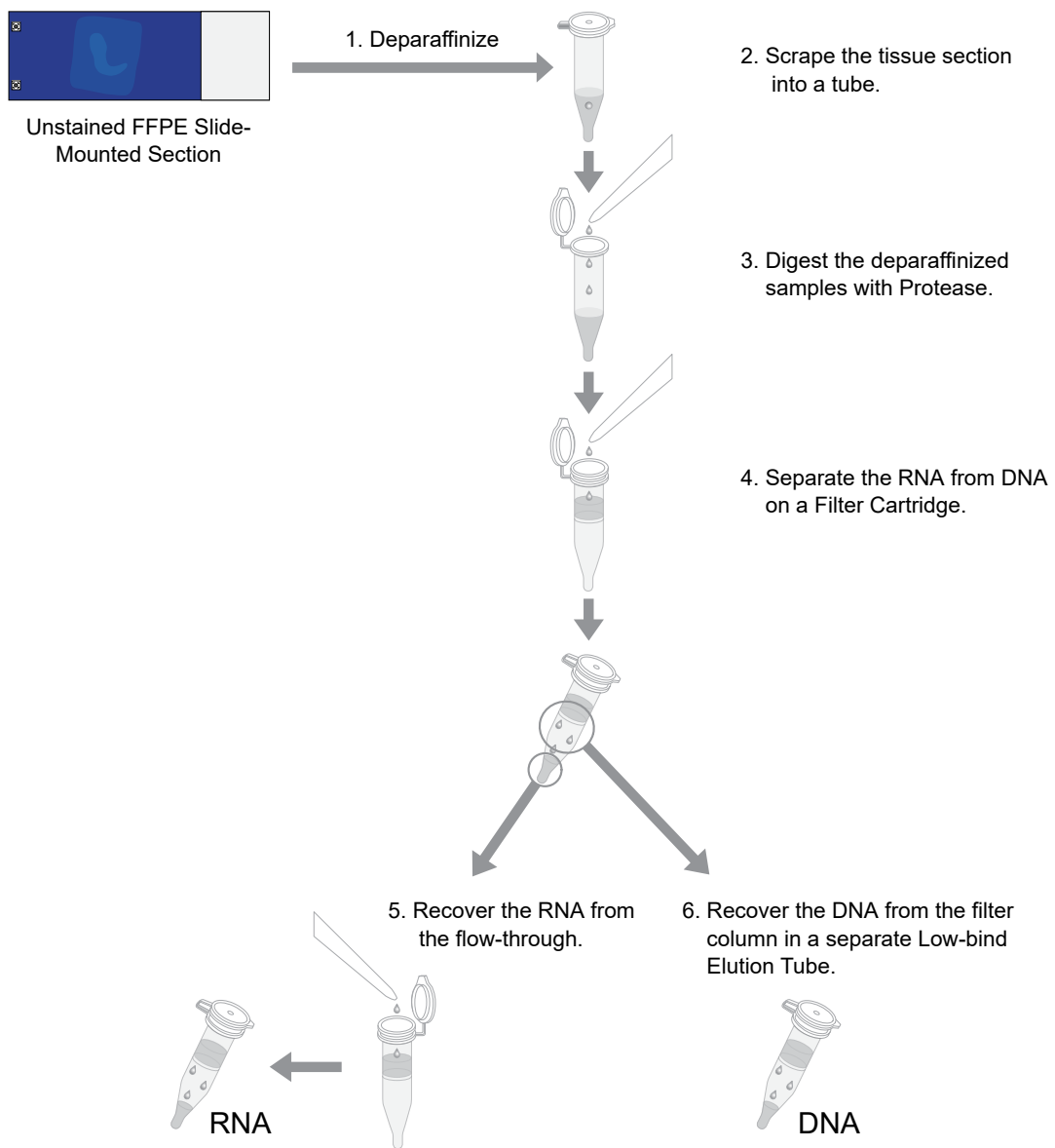
IMPORTANT! Check the expiration date on the box. If the kit is expired, select another kit.

4. Click **Save**.
The sample is no longer listed in the **To Be Extracted** list.

4

Isolate RNA and DNA from FFPE samples

Sample extraction workflow



Review the procedural guidelines

Before you begin the procedures in this section, review the procedural guidelines (see “Procedural guidelines” on page 16).

Ion Torrent Dx FFPE Sample Preparation Kit component lot matching

The six components of the Ion Torrent Dx FFPE Sample Preparation Kit must be lot-matched with each other for use.

Component	Part No.	Storage
Ion Torrent Dx Total Nucleic Acid Isolation Kit (box 1 of 2)	A32434	-30°C to -10°C
Ion Torrent Dx Total Nucleic Acid Isolation Kit (box 2 of 2)	A32435	15°C to 30°C
Ion Torrent Dx DNA Quantification Kit	A32437	2°C to 8°C
Ion Torrent Dx RNA Quantification Kit	A32438	2°C to 8°C
Ion Torrent Dx cDNA Synthesis Kit	A32436	-30°C to -10°C
Ion Torrent Dx Sample Dilution Kit	A32439	15°C to 30°C

Equilibrate the equipment and reagents

Before starting the following procedure:

- Equilibrate a benchtop cold box in a -30°C to -10°C freezer for at least 24 hours before use.

Note: The cold box holds temperature for up to 1 hour on the bench.

- Equilibrate aluminum cold blocks for 96-well plates at 2–8°C before use.
- Power on three dry-bath heaters with aluminum heat blocks 45 minutes before starting the procedure. Set the heaters to 55°C, 90°C, and 95°C.



CAUTION! Use care when working near the heat block to avoid being burned.

Note: Ensure that the heaters are calibrated.

- Remove the Protease and DNase from the freezer, then place them in a cold box equilibrated at -30°C to -10°C.

- Thaw the 10X DNase Buffer at room temperature (15°C to 30°C) and hold at room temperature until use.
- If you plan to quantify the DNA and RNA immediately after the extraction, equilibrate the quantification kit reagents (except the standards) to room temperature for at least 30 minutes before performing the assays.

Prepare wash buffers

Prepare the following buffers before using the Ion Torrent Dx FFPE Sample Preparation Kit. These buffers only need to be prepared once for each kit.

1. To prepare Wash 1 Buffer, add 14 mL of ACS grade 100% ethanol (EtOH) to the bottle labeled "Wash 1 Concentrate". Cap the bottle tightly and mix well by inverting the bottle 5 times.
2. To prepare Wash 2 Buffer, add 14 mL of ACS grade 100% ethanol to the bottle labeled "Wash 2 Concentrate". Cap the bottle tightly and mix well by inverting the bottle 5 times.
3. Mark the bottle labels to indicate that ethanol has been added (" +EtOH," initials, and date). Store the reconstituted Wash 1 and 2 Buffers at room temperature.

Deparaffinize and digest samples

Kit components used in this procedure

Kit component	Box
Digestion Buffer (green cap) Dilution Solution (black cap)	Ion Torrent Dx Total Nucleic Acid Isolation Kit box 2 of 2 (Part No. A32435, stored at 15°C to 30°C)
Protease (blue cap)	Ion Torrent Dx Total Nucleic Acid Isolation Kit box 1 of 2 (Part No. A32434, stored at -30°C to -10°C)

Prepare 1X Digestion Buffer

1. Label a nuclease-free 1.5-mL low-retention microcentrifuge tube for each FFPE tissue sample. Label each tube (cap and side) with its Sample ID using a marker that is resistant to xylene and ethanol.
2. Vortex the Digestion Buffer (green cap) and Dilution Solution (black cap) supplied in the kit for ~5 seconds each, then pulse centrifuge to collect the contents.

3. In a separate 1.5-mL low-retention microcentrifuge tube, prepare a master mix of the 1X Digestion Buffer as follows, where "n" is the number of tissue samples:

	Component	Volume per reaction	
		For ≤6 samples	For ≥7 samples
<input type="checkbox"/>	Digestion Buffer (green cap)	$(n+1) \times 25 \mu\text{L}$	$(n+2) \times 25 \mu\text{L}$
<input type="checkbox"/>	Dilution Solution (black cap)	$(n+1) \times 75 \mu\text{L}$	$(n+2) \times 75 \mu\text{L}$
<input type="checkbox"/>	Total 1X Digestion Buffer	$(n+1) \times 100 \mu\text{L}$	$(n+2) \times 100 \mu\text{L}$

4. Vortex the 1X Digestion Buffer for ~5 seconds to mix, then pulse centrifuge to collect.
5. Add 100 μL of 1X Digestion Buffer to each labeled tube from step 1.

Deparaffinize dipped FFPE slides



WARNING! Xylene is a toxic substance. Read the safety data sheet provided by the manufacturer. Handle it only in a well-ventilated area using personal protection equipment, and discard the waste according to regulations.

IMPORTANT! These instructions are only for paraffin-dipped FFPE slides. For slides that have not been dipped in paraffin, see “Deparaffinize undipped FFPE slides” on page 31.

Note:

- Use fresh xylene and fresh ACS-grade 100% ethanol after two rounds of deparaffinization with dipped slides. Each jar should have ~400 mL of either xylene or ethanol and be clearly marked with the date and initials after replacing the solutions.
- Perform the following steps carefully to avoid tissue loss.

1. Scrape any excess paraffin from each slide.
 - a. Grasp the slide at the slide label, and firmly hold the slide in an upright vertical position with the bottom oriented on the lab-bench paper.
 - b. Using a sterile disposable scalpel, scrape the layer of paraffin from the back of the slide. Use even pressure to scrape the back from top to bottom. Repeat if necessary to remove all the paraffin.

IMPORTANT! Use light pressure to prevent cracking the slide.

- c. If the tissue section cannot be visualized, do not perform this step. Turn the slide so the label and tissue face the operator. Carefully scrape around the tissue section to remove the paraffin.

Note: Scrape away from the tissue section to avoid accidentally removing the section itself.

- d. Repeat steps a–c for each slide, using a new scalpel for each unique sample.

Note: Properly discard used scalpels.

2. Fill a staining dish or jar with ~400 mL of xylene.
3. Place the slides in a slide rack, then completely submerge the rack in the xylene for 5 minutes at room temperature.
4. Incubate the slides for 30 minutes at room temperature. At ~10-minute intervals, lift the rack up and down 3 times to mix.
5. Remove the rack, then drain any excess xylene solution by tilting the rack.
6. Fill a staining dish or jar with ~400 mL of fresh xylene, then completely submerge the slide rack.
7. Incubate the slides for 15 minutes at room temperature. After ~7.5 minutes, lift the rack up and down 3 times to mix.
8. Remove the slides, then drain any excess xylene solution by tilting the slide holder.
9. Inspect the slides. If any paraffin remains, repeat steps 6–8 one more time.
10. Fill a staining dish or jar with ~400 mL of 100% ethanol.
11. Completely submerge the slides in the rack in the 100% ethanol for 5 minutes at room temperature.
12. Remove the rack, then drain any excess ethanol by tilting the rack.
13. Touch the edge of each slide with a clean laboratory wipe to wick any remaining ethanol from the surface, then lay the slide (section-side up) on a clean laboratory wipe.
14. Air dry each slide for at least 15 minutes.

Note: The drying time can vary depending on the section size. Ensure that there are no droplets on the tissue section before scraping.

15. Proceed to “Collect the tissue” on page 32.

Deparaffinize undipped FFPE slides



WARNING! Xylene is a toxic substance. Read the safety data sheet provided by the manufacturer. Handle it only in a well-ventilated area using personal protection equipment, and discard the waste according to regulations.

IMPORTANT! These instructions are only for FFPE slides that have not been dipped in paraffin. For slides that have been dipped in paraffin, see “Deparaffinize dipped FFPE slides” on page 30.

Note:

- Use fresh xylene and fresh ACS-grade 100% ethanol each day. Each jar should have ~400 mL of either xylene or ethanol and be clearly marked with the date and initials after replacing the solutions.
 - Perform the following steps carefully to avoid tissue loss.
-

1. Fill a staining dish or jar with ~400 mL of xylene.
 2. Place the slides with the unstained FFPE tissue sections in a slide rack, then completely submerge the rack in the xylene for 5 minutes at room temperature.
 3. Remove the rack, then drain any excess xylene solution by tilting the rack.
 4. Inspect the slides. If any paraffin remains, repeat steps 1–2 one more time.
 5. Fill a staining dish or jar with ~400 mL of 100% ethanol.
 6. Completely submerge the slides in the rack in the 100% ethanol for 5 minutes at room temperature.
 7. Remove the rack, then drain any excess ethanol by tilting the rack.
 8. Touch the edge of each slide to a clean laboratory wipe to wick any remaining ethanol from the surface, then lay the slide (section-side up) on a clean laboratory wipe.
 9. Air dry each slide for at least 15 minutes.
-

Note: The drying time can vary depending on the section size. Ensure that there are no droplets on the tissue section before scraping.

10. Proceed to “Collect the tissue”.

Collect the tissue

IMPORTANT! Before proceeding, review the tissue input requirements in “Tissue input requirements for FFPE sample extraction” on page 15 and “Guidelines for FFPE samples” on page 17.

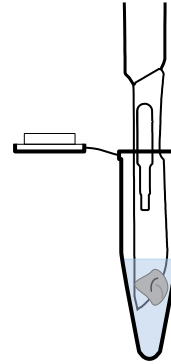
In this procedure, scrape each tissue section into the appropriate labeled 1.5-mL low-retention microcentrifuge tube containing 1X Digestion Buffer (prepared in “Prepare 1X Digestion Buffer” on page 29).

1. Pipet 4 μ L of 1X Digestion Buffer from the labeled tube evenly across the fixed tissue section on the slide to pre-wet the tissue section.
-

Note: Larger sections may need an additional 4 μ L of Digestion Buffer.

2. Using a sterile disposable scalpel, scrape the tissue in a single direction, then collect the tissue into a cohesive mass on the tip of the scalpel blade.

3. Carefully insert the scalpel blade with the tissue mass into the 1X Digestion Buffer in the 1.5-mL low-retention microcentrifuge tube. Rinse the tissue from the blade into the buffer, then ensure that the entire mass is in solution.
4. Remove and inspect the blade to ensure that no tissue remains on it.
5. Inspect the slide to ensure that all the tissue has been removed (the slide should be translucent). Discard the scalpel in a waste container for sharp objects.



Digest the deparaffinized samples

1. Flick-mix the Protease (blue cap) 4 times with your finger, then pulse centrifuge to collect the contents.
2. Add 4 μ L of Protease to each tissue sample tube.
3. Flick-mix each sample tube 4 times, then pulse centrifuge.
4. Incubate the samples at 55°C in a calibrated heat block for 1 hour.

Note: During incubation, proceed to “Label the Filter Cartridges and Collection Tubes” and “Preheat the Elution Solution” to save time.

5. Pulse centrifuge to collect any condensation droplets.
6. Incubate the samples at 90°C in a calibrated heat block for 1 hour.
7. Pulse centrifuge to collect any condensation droplets, then proceed immediately to “Separate RNA from DNA on a Filter Cartridge” on page 35.

Separate RNA from DNA

Kit components used in this procedure

Kit component	Box
Filter Cartridges ^[1]	Ion Torrent Dx Total Nucleic Acid Isolation Kit box 2 of 2 (Part No. A32435, stored at 15°C to 30°C)
Collection Tubes	
Low-bind Elution Tubes	
Elution Solution (red cap)	
Isolation Additive (brown cap)	

^[1] Includes a filter column pre-inserted in a Collection Tube.

Label the Filter Cartridges and Collection Tubes

Note: To save time, label sets of Filter Cartridges (filter column + Collection Tube) and Collection Tubes (tube only) in advance. Use ethanol-resistant markers for labeling. Do not write on the side of the filter column, because the ink may bleed into the sample.

For each FFPE tissue sample, label the following cartridges and tubes as indicated for use in the subsequent DNA and RNA extraction steps:

- Filter Cartridges (2)
- Collection Tubes (1)
- Low-bind Elution Tubes (2)

Component	Label		Material collected	
	Filter column cap	Tube	Filter	Tube
Filter Cartridge (A)	Sample ID and "DNA"	Sample ID and "RNA"	Bound DNA	Flow-through RNA
Collection Tube (B)	—	Sample ID and "DNA Wash"	—	DNA wash
Filter Cartridge (C)	Sample ID and "RNA"	Sample ID and "RNA Wash"	Bound RNA	RNA wash
Low-bind Elution Tube (D)	—	Sample ID, "RNA," date, and operator initials	—	Eluted RNA
Low-bind Elution Tube (E)	—	Sample ID, "DNA," date, and operator initials	—	Eluted DNA

Preheat the Elution Solution

1. For each sample, pipet 125 μ L of Elution Solution (red cap) into a 1.5-mL low-retention microcentrifuge tube.
2. Place the tube(s) of Elution Solution in the 95°C heat block for at least 5 minutes. Keep the Elution Solution in the heat block throughout the following procedure.



CAUTION! Use care when working near the heat block to avoid being burned.

Note: A tube rack may be placed on top of the tubes to prevent the tubes from popping open.

Separate RNA from DNA on a Filter Cartridge

Use the following previously labeled Filter Cartridges and Collection Tubes for the following procedure:

Component	Label		Material collected	
	Filter column cap	Tube	Filter	Tube
Filter Cartridge (A)	Sample ID and "DNA"	Sample ID and "RNA"	Bound DNA	Flow-through RNA
Collection Tube (B)	—	Sample ID and "DNA Wash"	—	DNA wash

1. Place the labeled Filter Cartridge (A) in a tube rack.
2. Add 120 μ L of the Isolation Additive (brown cap) to the digested sample, then mix by pipetting up and down 5 times. The sample appears slightly cloudy.
3. Transfer the digested sample and Isolation Additive mix (~224 μ L) to the Filter Cartridge, then close the lid.
4. Centrifuge the Filter Cartridge at 10,000 rcf for 30 seconds in a microcentrifuge.

IMPORTANT! Do not discard the flow-through in the Collection Tube (labeled with Sample ID and "RNA"). The flow-through contains the RNA.

5. Place the filter column with the bound DNA in a new Collection Tube (B), then store it at 2–8°C for later DNA purification in "Recover the DNA from the Filter Cartridge" on page 40.
6. Proceed to "Recover the RNA from the flow-through" on page 36.

Recover the RNA from the flow-through

Kit components used in this procedure

Kit component	Source/Box
Filter Cartridges Collection Tubes Low-bind Elution Tubes	Previously labeled
Wash 1 Buffer Wash 2 Buffer	Previously prepared from concentrate
Dilution Solution (black cap)	Ion Torrent Dx Total Nucleic Acid Isolation Kit box 2 of 2 (Part No. A32435, stored at 15°C to 30°C)
10X DNase Buffer (white cap) DNase (purple cap)	Ion Torrent Dx Total Nucleic Acid Isolation Kit box 1 of 2 (Part No. A32434, stored at -30°C to -10°C)

Bind the RNA to the Filter Cartridge

Use the following previously labeled Filter Cartridge for the following procedure:

Component	Label		Material collected	
	Filter column cap	Tube	Filter	Tube
Filter Cartridge (C)	Sample ID and "RNA"	Sample ID and "RNA Wash"	Bound RNA	RNA wash

1. Place the new Filter Cartridge (C) in a tube rack.
2. Add 275 μ L of ACS-grade 100% ethanol to the flow-through containing RNA (the tube labeled with the Sample ID and "RNA") from "Separate RNA from DNA on a Filter Cartridge" on page 35.
3. Mix well by pipetting up and down 5 times, then transfer the sample (~450 μ L) to the new Filter Cartridge (C).
4. Centrifuge the Filter Cartridge at 10,000 rcf for 30 seconds in a microcentrifuge.
5. Discard the flow-through in the Collection Tube, then reinsert the filter column into the same Collection Tube.
6. Add 600 μ L of Wash 1 Buffer (prepared in "Prepare wash buffers" on page 29) to the Filter Cartridge.
7. Centrifuge the Filter Cartridge at 10,000 rcf for 30 seconds.

8. Discard the flow-through, then reinsert the filter column in the same Collection Tube.
9. Centrifuge the Filter Cartridge at 10,000 rcf for 30 seconds to remove any remaining fluid.

Treat the RNA bound to the Filter Cartridge with DNase

1. In a 1.5-mL low-retention microcentrifuge tube, prepare a master mix of 1X DNase Solution as follows, where "n" is the number of samples you are preparing.

	Component	Volume per reaction	
		For ≤6 samples	For ≥7 samples
<input type="checkbox"/>	Dilution Solution (black cap)	$(n+1) \times 50 \mu\text{L}$	$(n+2) \times 50 \mu\text{L}$
<input type="checkbox"/>	10X DNase Buffer (white cap)	$(n+1) \times 6 \mu\text{L}$	$(n+2) \times 6 \mu\text{L}$
<input type="checkbox"/>	DNase (purple cap)	$(n+1) \times 4 \mu\text{L}$	$(n+2) \times 4 \mu\text{L}$
<input type="checkbox"/>	Total Volume	$(n+1) \times 60 \mu\text{L}$	$(n+2) \times 60 \mu\text{L}$

2. Flick the 1X DNase Solution tube 4 times to mix, then pulse centrifuge to collect.
3. Pipet 60 μL of the 1X DNase Solution into the center of each filter column (previously labeled with Sample ID and "RNA").

IMPORTANT! To avoid puncturing, do **NOT** touch the pipette tip to the filter.

4. Hold the Filter Cartridge at room temperature for 30 minutes.

Note: If you plan to quantify the DNA and RNA immediately after the extraction, begin equilibrating the following reagents from the quantification kits at room temperature for at least 30 minutes:

- DNA Dye Reagent (blue cap)
- DNA Buffer (white cap)
- RNA Dye Reagent (green cap)
- RNA Buffer (blue cap)

Wash the RNA bound to the Filter Cartridge

Use the following previously labeled Filter Cartridges and Low-bind Elution Tubes for the following procedure.

Component	Label		Material collected	
	Filter column cap	Tube	Filter	Tube
Filter Cartridge (C)	Sample ID and "RNA"	Sample ID and "RNA Wash"	Bound RNA	RNA wash
Low-bind Elution Tube (D)	—	Sample ID, "RNA," date, and operator initials	—	Eluted RNA

1. Add 600 µL of Wash 1 Buffer to the Filter Cartridge (C).
2. Hold the Filter Cartridge for 30 seconds at room temperature, then centrifuge the Filter Cartridge at 10,000 rcf for 30 seconds.
3. Discard the flow-through, then reinsert the filter column in the same Collection Tube.
4. Add 500 µL of Wash 2 Buffer (prepared in “Prepare wash buffers” on page 29) to the Filter Cartridge, then centrifuge the Filter Cartridge at 10,000 rcf for 30 seconds.
5. Discard the flow-through, then reinsert the filter column into the same Collection Tube.
6. Repeat steps 4 and 5 for a second wash.
7. Centrifuge the Filter Cartridge at 20,000–21,000 rcf for 2 minutes to remove any remaining fluid.
8. Remove the filter column from the tube, then touch the bottom of the column with a clean laboratory wipe to wick off any remaining wash buffer.
9. Transfer the filter column to the pre-labeled Low-bind Elution Tube (D).

Elute the RNA

Use the following components for this procedure.

Component	Label		Material collected	
	Filter column cap	Tube	Filter	Tube
Filter column with bound RNA from Filter Cartridge (C)	Sample ID and "RNA"	—	Bound RNA	—
Low-bind Elution Tube (D)	—	Sample ID, "RNA," date, and operator initials	—	Eluted RNA

IMPORTANT!

- Keep the 1.5-mL low-retention microcentrifuge tube containing preheated Elution Solution in the heat block throughout the procedure to maintain a 95°C temperature.
- Change pipette tips between samples when pipetting Elution Solution across multiple samples.

1. Remove the Elution Solution from the heat block, and pulse centrifuge the tube to collect the contents. Return the tube to the heat block.



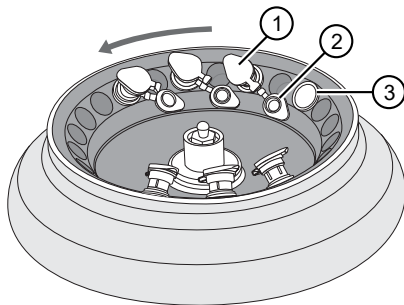
CAUTION! The heat block and Elution Solution are hot. Use care when handling tubes and tube contents to avoid being burned.

2. Wet the pipette tip by slowly pipetting up and down 3 times in the pre-heated Elution Solution.

3. Slowly pipet up 30 μ L of Elution Solution, then confirm that there are no large air gaps in the tip (a small air gap at the bottom of the tip is acceptable). Pipet the solution into the center of the filter column in the Low-bind Elution Tube (D) (from step 9 in “Wash the RNA bound to the Filter Cartridge”).

IMPORTANT! To avoid puncturing the filter, do not touch it with the pipette tip.

4. Close the cap on the filter column, then hold the filter column/Low-bind Elution Tube assembly at room temperature for 1 minute. Close the cap on the Elution Solution tube in the heat block.
5. Insert the filter column/Low-bind Elution Tube assembly in the microcentrifuge in the orientation shown below. To prevent the Low-bind Elution Tube caps from breaking, place a 0.2-mL tube adapter in the position shown.



- ① Filter column cap (closed)
- ② Low-bind Elution Tube cap (open)
- ③ 0.2-mL tube adapter

6. Centrifuge at 20,000–21,000 rcf for 1 minute.

Note: The eluted RNA is in the Low-bind Elution Tube. If the tube cap breaks in the centrifuge, transfer the sample to a new labeled Low-bind Elution Tube.

7. Discard the filter column.
8. Temporarily store the sample at 2–8°C if quantifying on the same day.

STOPPING POINT If you are not quantifying on the same day, store the recovered RNA aliquots at –90°C to –60°C for up to 5 months.

Recover the DNA from the Filter Cartridge

Wash the DNA bound to the Filter Cartridge

Use the following pre-labeled Filter Cartridges and tubes for the following protocol:

Component	Label		Material collected	
	Filter column cap	Tube	Filter	Tube
Filter Cartridge (A)	Sample ID and "DNA"	Sample ID and "RNA"	Bound DNA	Flow-through RNA
Collection Tube (B)	—	Sample ID and "DNA Wash"	—	DNA wash
Low-bind Elution Tube (E)	—	Sample ID, "DNA," date, and operator initials	—	Eluted DNA

1. Retrieve the Filter Cartridge (A) with bound DNA and Collection Tube (B) from 2–8°C storage (previously stored in “Separate RNA from DNA on a Filter Cartridge” on page 35).
2. Add 600 µL of Wash 1 Buffer to the filter column.
3. Hold the Filter Cartridge for 30 seconds at room temperature, then centrifuge at 10,000 rcf for 30 seconds.
4. Discard the flow-through, then reinsert the filter column into the same Collection Tube (B).
5. Add 500 µL of Wash 2 Buffer to the filter column, then centrifuge at 10,000 rcf for 30 seconds.
6. Discard the flow-through, then reinsert the filter column into the same Collection Tube (B).
7. Repeat steps 5 and 6 for a second wash.
8. Centrifuge the Filter Cartridge at 20,000–21,000 rcf for 2 minutes to remove any remaining fluid.
9. Remove the filter column from the tube, then touch the bottom of the column with a clean laboratory wipe to wick off any remaining wash buffer.
10. Transfer the filter column to the pre-labeled Low-bind Elution Tube (E).

Elute the DNA

Use the following components for this procedure.

Component	Label		Material collected	
	Filter column cap	Tube	Filter	Tube
Filter column with bound DNA from Filter Cartridge (A)	Sample ID and "DNA"	—	Bound DNA	—
Low-bind Elution Tube (E)	—	Sample ID, "DNA," date, and operator initials	—	Eluted DNA

IMPORTANT!

- Keep the 1.5-mL low-retention microcentrifuge tube containing preheated Elution Solution in the heat block throughout the procedure to maintain a 95°C temperature.
- Change pipette tips between samples when pipetting Elution Solution across multiple samples.

1. Remove the Elution Solution from the heat block, and pulse centrifuge the tube to collect the contents. Return the tube to the heat block.



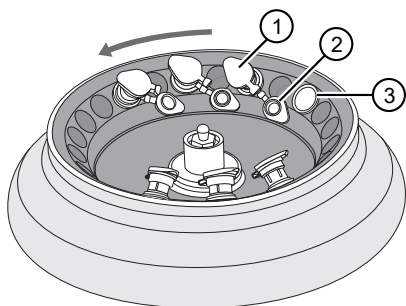
CAUTION! The heat block and Elution Solution are hot. Use care when handling tubes and tube contents to avoid being burned.

2. Wet the pipette tip by slowly pipetting up and down 3 times in the pre-heated Elution Solution.
3. Slowly pipet up 30 µL of Elution Solution, then confirm that there are no large air gaps in the tip (a small air gap at the bottom of the tip is acceptable). Pipet the solution into the center of the filter column in the Low-bind Elution Tube (E) (from step 10 in “Wash the DNA bound to the Filter Cartridge”).

IMPORTANT! To avoid puncturing the filter, do not touch it with the pipette tip.

4. Close the cap on the filter column, then hold the filter column/Low-bind Elution Tube assembly at room temperature for 1 minute. Close the cap on the Elution Solution tube in the heat block.

5. Insert the filter column/Low-bind Elution Tube assembly in the microcentrifuge in the orientation shown below. To prevent the Low-bind Elution Tube caps from breaking, place a 0.2-mL tube adapter in the position shown.



- ① Filter column cap (closed)
- ② Low-bind Elution Tube cap (open)
- ③ 0.2-mL tube adapter

6. Centrifuge at 20,000–21,000 rcf for 1 minute.

Note: The eluted DNA is in the Low-bind Elution Tube. If the tube cap breaks in the centrifuge, transfer the sample to a new labeled Low-bind Elution Tube.

7. Discard the filter column.
8. Temporarily store the sample at 2–8°C if quantifying on the same day.

STOPPING POINT If you are not quantifying on the same day, store the recovered DNA aliquots at –30°C to –10°C for up to 12 months.



DNA and RNA quantification

Review the procedural guidelines

Before you begin the procedures in this section, review the procedural guidelines (see “Procedural guidelines” on page 16).

Prepare the reagents and equipment

- If the DNA and RNA samples were frozen for storage, thaw them at room temperature until no ice crystals are present, then transfer them to 2–8°C storage until use.
- Equilibrate a benchtop cold box at –30°C to –10°C for at least 24 hours before use.

Note: The cold box holds temperature for up to 1 hour on the bench.

- Equilibrate the DNA Dye Reagent, RNA Dye Reagent, DNA Buffer, and RNA Buffer to room temperature for at least 30 minutes before use.
- Keep the DNA Std and RNA Std at 2–8°C, until the working plate has been set up and the standards are ready to be added.
- If necessary, set up the fluorometer/fluorescence reader to read the appropriate excitation and emission wavelengths:

Dye reagent	Excitation (nm)	Emission (nm)
RNA Dye Reagent	620/15	680/30
DNA Dye Reagent	485/20	528/20

Set up the DNA quantification assay

Kit components used in this procedure

Kit component	Box
DNA Dye Reagent (blue cap)	Ion Torrent Dx DNA Quantification Kit (Part No. A32437, stored at 2°C to 8°C)
DNA Buffer (white cap)	
DNA Std - 0 ng/μL (white cap)	
DNA Std - 0.5 ng/μL (green cap)	
DNA Std - 4 ng/μL (red cap)	
DNA Std - 10 ng/μL (yellow cap)	

Prepare the DNA Working Solution

1. Determine the number of DNA standards to use with your quantification system.

IMPORTANT! We recommend using 4 standards. If your quantification system does not allow the use of 4 standards, use the maximum allowed by the system. At a minimum, you must use the 0 ng/μL and 10 ng/μL DNA standards. Note that R² values should only be evaluated when 3 or more standards are used.

2. Calculate the number of reactions using the following formula:

$$\mathbf{S \text{ (\# of standards)} + N \text{ (\# of samples)} + 1 = \# \text{ of reactions}}$$

3. Calculate the total volume of DNA Dye Reagent and DNA Buffer required for the number of reactions:

$$\mathbf{\# \text{ reactions} \times 1 \text{ } \mu\text{L (DNA Dye Reagent)} = \text{total volume of DNA Dye Reagent}}$$

$$\mathbf{\# \text{ reactions} \times 199 \text{ } \mu\text{L (DNA Buffer)} = \text{total volume of DNA Buffer}}$$

4. Mix the DNA Buffer and DNA Dye Reagent bottles by inverting 5 times.
5. Prepare the DNA Working Solution: Pipet the calculated volume of DNA Buffer into a pre-labeled tube, then add the calculated volume of DNA Dye Reagent into the same tube.
6. Vortex the tube for ~5 seconds, then proceed to the next steps.

IMPORTANT! If you are not immediately proceeding to the next steps, protect the DNA Working Solution from light. The DNA Working Solution must be used within 3 hours.

Prepare the DNA standards

1. Add 190 μL of DNA Working Solution to each well or tube that will contain a DNA standard.
2. Vortex each DNA standard for ~5 seconds, then pulse centrifuge. Refer to the following table of DNA standards and concentrations.

Note: If you are using fewer than four standards, at a minimum you must use the 0 ng/ μL and 10 ng/ μL DNA standards.

Standard	Concentration
DNA STD 1 (white cap)	0 ng/ μL
DNA STD 2 (green cap)	0.5 ng/ μL
DNA STD 3 (red cap)	4.0 ng/ μL
DNA STD 4 (yellow cap)	10 ng/ μL

3. Pipet 10 μL of each DNA standard into its designated well or tube.

Prepare the DNA samples

1. Add 196 μL of DNA Working Solution to each well or tube that will contain a DNA sample.
2. Vortex each DNA sample for ~5 seconds, then pulse centrifuge.
3. Pipet 4 μL of each DNA sample into its designated well or tube, then proceed to set up the RNA quantification assay.

Set up the RNA quantification assay

IMPORTANT! Wipe down your work surface and pipettes with an RNase decontamination solution. Change gloves before starting and as needed to maintain RNase-free conditions.

Kit components used in this procedure

Kit component	Box
RNA Dye Reagent (green cap)	Ion Torrent Dx RNA Quantification Kit (Part No. A32438, stored at 2°C to 8°C)
RNA Buffer (blue cap)	
RNA Std - 0 ng/ μL (teal cap)	
RNA Std - 0.5 ng/ μL (tan cap)	
RNA Std - 4 ng/ μL (purple cap)	
RNA Std - 10 ng/ μL (orange cap)	

Prepare the RNA working solution

1. Determine the number of RNA standards to use with your quantification system.

IMPORTANT! We recommend using 4 standards. If your quantification system does not allow the use of 4 standards, use the maximum allowed by the system. At a minimum, you must use the 0 ng/μL and 10 ng/μL RNA standards. Note that R² values should only be evaluated when 3 or more standards are used.

2. Calculate the number of reactions using the following formula:

S (# of standards) + N (# of samples) + 1 = # of reactions

3. Calculate the total volume of RNA Dye Reagent and RNA Buffer required for the number of reactions:

reactions × 1 μL (RNA Dye Reagent) = total volume of RNA Dye Reagent

reactions × 199 μL (RNA Buffer) = total volume of RNA Buffer

4. Mix the RNA Buffer and RNA Dye Reagent bottles by inverting 5 times.
5. Prepare the RNA Working Solution: Pipet the total volume of RNA Buffer into a pre-labeled tube, then add the total volume of RNA Dye Reagent into the same tube.
6. Vortex the tube for ~5 seconds, then proceed to the next steps.

IMPORTANT! If you are not immediately proceeding to the next steps, protect the RNA Working Solution from light. The RNA Working Solution must be used within 3 hours.

Prepare the RNA standards

1. Add 190 μL of RNA Working Solution to each well or tube that will contain an RNA standard.
2. Flick mix each RNA Standard 4 times, then pulse centrifuge. Refer to the following table of RNA Standards and concentrations.

Note: If you are using fewer than four standards, at a minimum you must use RNA STD 1 (0 ng/μL) and RNA STD 4 (10 ng/μL).

Standard	Concentration
RNA STD 1 (teal cap)	0 ng/μL
RNA STD 2 (tan cap)	0.5 ng/μL
RNA STD 3 (purple cap)	4.0 ng/μL
RNA STD 4 (orange cap)	10 ng/μL

3. Pipet 10 μL of each RNA Standard into its designated well or tube.

Prepare the RNA samples

1. Add 196 μL of RNA Working Solution to each well or tube that will contain an RNA sample.
2. Flick mix each RNA sample 4 times, then pulse centrifuge.
3. Pipet 4 μL of each RNA sample into its designated well or tube, then proceed to quantification.

Run the quantification assays

1. Incubate the prepared DNA and RNA standards and samples for at least 2 minutes at room temperature before reading.
2. Determine the concentration of the DNA and RNA samples in $\text{ng}/\mu\text{L}$ using a fluorometer/fluorescence reader and linear regression of the standards for DNA and RNA respectively.

The required minimum values for the Oncomine™ Dx Target Test are shown in the following table:

Table 1 Required sample concentrations and R^2 values from the linear regression of the standards

Sample type	Required concentration	Required R^2 value ^[1]
DNA	$\geq 0.83 \text{ ng}/\mu\text{L}$	≥ 0.99
RNA	$\geq 1.43 \text{ ng}/\mu\text{L}$	≥ 0.98

^[1] R^2 values should be evaluated only if the standard curve includes 3 or more points.

IMPORTANT! To proceed with library preparation, both the DNA and RNA concentrations from a single sample extraction must meet the minimum requirements. Do not use DNA from one extraction with RNA from a different extraction.

3. If the samples do not meet the minimum concentration requirements, repeat the extraction with increased tissue input (i.e., more sections) if possible.

STOPPING POINT If you do not dilute the sample on the same day, store the quantified DNA sample at -30°C to -10°C (for up to 12 months) and the quantified RNA sample at -90°C to -60°C (for up to 5 months).

Dilute the samples

Kit components used in this procedure

Kit component	Box
Dilution Solution (black cap)	Ion Torrent Dx Sample Dilution Kit (Part No. A32439, stored at 15°C to 30°C)

Sample Dilution Calculator

The Sample Dilution Calculator is a locked Microsoft™ Excel™ spreadsheet that is provided as an aide to prepare samples for library preparation. The Sample Dilution Calculator provides a uniform method for diluting DNA samples to 0.83 ng/μL and RNA samples to 1.43 ng/μL, including a 10% overage of the final sample dilution volume.

Copy the **SFW0000786_ODxTT_SampleDilution_Calculator_3.02.xlsm** file from the Oncomine™ Dx Target Test media storage device onto your hard drive before use.

IMPORTANT!

- The Sample Dilution Calculator (Part. No. SFW0000786) requires Microsoft™ Excel™ 2010 or later.
 - When a security warning appears after opening the Sample Dilution Calculator file, click **Enable Content** in the message bar to enable the dilution calculator macro.
-

Dilution volume guidelines

- Unless the test or study protocol dictates that multiple libraries must be made from a single sample (for example, replicate libraries), dilute only sufficient sample to prepare a single library.
- The Sample Dilution Calculator includes a 10% overage in its calculations. For manually calculated dilutions, include a 10% overage, except for low-concentration DNA (0.83–0.97 ng/μL) and RNA (1.43–1.93 ng/μL) samples where there is insufficient sample to include an overage.
- The default **Sample Volume (X1)** in the Sample Dilution Calculator is 2 μL. If needed, increase the **Sample Volume (X1)** until the **Volume of Dilution Solution (Y1)** is ≥2 μL. When all volumes are ≥2 μL, red highlighted table cells turn white, indicating sufficient volumes for accurate pipetting.
- In the following procedure, we recommend that you do not pipet volumes <2.0 μL. For samples that are ≥6.01 ng/μL, perform a normalization dilution to an intermediate concentration of 5 ng/μL, then further dilute the sample to a final concentration of 0.83 ng/μL for DNA and 1.43 ng/μL for RNA.
- Low-concentration DNA (0.83–0.97 ng/μL) and RNA (1.43–1.93 ng/μL) samples require pipetting volumes <2 μL to achieve the correct concentration. When necessary, only pipet volumes <2 μL with a pipette designed and calibrated to dispense volumes <2 μL.
- Do not use the Sample Dilution Calculator for low-concentration DNA (0.83–0.97 ng/μL) and RNA (1.43–1.93 ng/μL) samples. Instead, follow the detailed procedures that are provided in the second row of the tables on page 49 and page 52.
- Samples that do not meet the minimum concentration specification (DNA samples <0.83 ng/μL and RNA samples <1.43 ng/μL) must not be used for library preparation.

Thaw frozen samples

If DNA and RNA samples were frozen for storage, thaw them at room temperature until no ice crystals are present before proceeding. Transfer samples to 2–8°C storage until use.

Note: Freeze-thaw samples no more than 3 times.

Dilute DNA samples

IMPORTANT! Do not perform the following dilution procedures until you are ready to proceed directly to library preparation. Library preparation requires accurate input of 10 ng DNA. Pipetting volumes <2 μL is not recommended. When necessary, only pipet volumes <2 μL with a pipette designed and calibrated to dispense volumes <2 μL .

Library preparation requires dilution of DNA samples to a final concentration of 0.83 ng/ μL . See the following table for sample dilution instructions that are based on the starting sample concentration.

DNA concentration	Dilution procedure
<0.83 ng/ μL	Samples do not meet the minimum concentration specification and must not be used for library preparation.
0.83–0.97 ng/ μL	<p>Dilute the DNA sample to 0.83 ng/μL in a total volume of 12 μL.</p> <ol style="list-style-type: none"> Determine the volume of DNA sample required (10 ng \div Sample concentration ng/μL = N μL). Add Dilution Solution if required: <ul style="list-style-type: none"> If N is <12 μL, pipet N μL of DNA sample into a new labeled 1.5-mL low-retention microcentrifuge tube, then add (12 – N μL) of Dilution Solution, for a total volume of 12 μL. If N = 12 μL, skip to step 4. Pipet up and down 5 times to mix, then pulse centrifuge. Proceed directly to “Dilute RNA samples” on page 52. Do not store the diluted DNA samples for longer than necessary. <p>Note: Store the remaining undiluted DNA sample at -30°C to -10°C for up to 12 months.</p>
0.98–6.00 ng/ μL	<p>Perform the Direct Dilution to 0.83 ng/μL.</p> <ol style="list-style-type: none"> Open SFW0000786_ODxTT_SampleDilution_Calculator_3.02.xlsm in Microsoft[™] Excel[™], select the Calculator worksheet tab, then enter the Operator and Date Completed information. In column B, enter the Sample ID. <p>Note: Only alphanumeric characters (0–9 and A to Z), full stops/periods (.), underscores (_), or hyphens (-) are allowed.</p> In column C, enter the DNA sample concentration (C1) in ng/μL determined in the quantification assay. Pipet the indicated volume (X3) of undiluted DNA sample into a new labeled 1.5-mL low-retention microcentrifuge tube, then add the indicated volume of Dilution Solution (Y3). Pipet up and down 5 times to mix, then pulse centrifuge. Proceed directly to “Dilute RNA samples” on page 52. Do not store the diluted DNA samples for longer than necessary. <p>Note: Store the remaining undiluted DNA sample at -30°C to -10°C for up to 12 months.</p>

(continued)

DNA concentration	Dilution procedure
6.01–9.99 ng/ μ L	<p>Perform Normalization Dilution #1 and Normalization Dilution #2 to a final concentration of 0.83 ng/μL.</p> <p>Note: The default Sample Volume (X1) is 2 μL. When all volumes are ≥ 2 μL, red highlighted table cells turn white, indicating sufficient volumes for accurate pipetting have been met.</p> <ol style="list-style-type: none"> 1. Open SFW0000786_ODxTT_SampleDilution_Calculator_3.02.xlsm in Microsoft™ Excel™, select the Calculator worksheet tab, then enter the Operator and Date Completed information. 2. In column B, enter the Sample ID. <p>Note: Only alphanumeric characters (0–9 and A to Z), full stops/periods (.), underscores (_), or hyphens (-) are allowed.</p> 3. In column C, enter the DNA sample concentration (C1) in ng/μL determined in the quantification assay. 4. Increase the value for Sample Volume (X1) until the Volume of Dilution Solution (Y1) is ≥ 2 μL. 5. Pipet the indicated volume (X1) of undiluted DNA sample into a new labeled 1.5-mL low-retention microcentrifuge tube, then add the indicated volume of Dilution Solution (Y1). 6. Pipet up and down 5 times to mix, then pulse centrifuge. 7. Pipet the indicated volume (X2) of diluted DNA sample (5 ng/μL) into a new labeled 1.5-mL low-retention microcentrifuge tube, then add the indicated volume of Dilution Solution (Y2). 8. Pipet up and down 5 times to mix, then pulse centrifuge. 9. Proceed directly to “Dilute RNA samples” on page 52. Do not store the diluted DNA samples for longer than necessary. <p>Note: Store the remaining Normalization Dilution #1 (5 ng/μL) and undiluted DNA sample at -30°C to -10°C for up to 12 months.</p>

(continued)

DNA concentration	Dilution procedure
≥10 ng/μL	<p>Perform Normalization Dilution #1 and Normalization Dilution #2 to a final concentration of 0.83 ng/μL.</p> <p>Note: The default Sample Volume (X1) is 2 μL. When all volumes are ≥2 μL, red highlighted table cells turn white, indicating sufficient volumes for accurate pipetting have been met.</p> <ol style="list-style-type: none"> 1. Open SFW0000786_ODxTT_SampleDilution_Calculator_3.02.xlsm in Microsoft™ Excel™, select the Calculator worksheet tab, then enter the Operator and Date Completed information. 2. In column B, enter the Sample ID. <ul style="list-style-type: none"> Note: Only alphanumeric characters (0–9 and A to Z), full stops/periods (.), underscores (_), or hyphens (-) are allowed. 3. In column C, enter the DNA sample concentration (C1) in ng/μL determined in the quantification assay. 4. Pipet the indicated volume (X1) of undiluted DNA sample into a new labeled 1.5-mL low-retention microcentrifuge tube, then add the indicated volume of Dilution Solution (Y1). 5. Pipet up and down 5 times to mix, then pulse centrifuge. 6. Pipet the indicated volume (X2) of diluted DNA sample (5 ng/μL) into a new labeled 1.5-mL low-retention microcentrifuge tube, then add the indicated volume of Dilution Solution (Y2). 7. Pipet up and down 5 times to mix, then pulse centrifuge. 8. Proceed directly to “Dilute RNA samples” on page 52. Do not store the diluted DNA samples for longer than necessary. <p>Note: Store the remaining Normalization Dilution #1 (5 ng/μL) and undiluted DNA sample at –30°C to –10°C for up to 12 months.</p>

Dilute RNA samples

IMPORTANT! Do not perform the following dilution procedures until you are ready to proceed directly to library preparation. Library preparation requires accurate input of 10 ng RNA. Pipetting volumes <2 μL is not recommended. When necessary, only pipet volumes <2 μL with a pipette designed and calibrated to dispense volumes <2 μL .

Library preparation requires dilution of RNA samples to a final concentration of 1.43 ng/ μL . See the following table for sample dilution instructions that are based on the sample starting concentration.

RNA concentration	Dilution procedure
<1.43 ng/ μL	Samples do not meet the minimum concentration specification and must not be used for library preparation.
1.43–1.93 ng/ μL	<p>Dilute the RNA sample to 1.43 ng/μL in a total volume of 7 μL.</p> <ol style="list-style-type: none"> Determine the volume of RNA sample required (10 ng \div Sample concentration ng/μL = N μL). Add Dilution Solution if required: <ul style="list-style-type: none"> If N is <7 μL, pipet N μL of RNA sample into a new labeled 1.5-mL low-retention microcentrifuge tube, then add (7 – N μL) of Dilution Solution, for a total volume of 7 μL. If N = 7 μL, skip to step 4. Pipet up and down 5 times to mix, then pulse centrifuge. Proceed directly to “Reverse transcribe the RNA” on page 56. Do not store the diluted RNA samples for longer than necessary. <p>Note: Store the remaining undiluted RNA sample at -90°C to -60°C for up to 5 months.</p>
1.94–6.00 ng/ μL	<p>Perform the Direct Dilution to 1.43 ng/μL.</p> <ol style="list-style-type: none"> Open SFW0000786_ODxTT_SampleDilution_Calculator_3.02.xlsm in Microsoft™ Excel™, select the Calculator worksheet tab, then enter the Operator and Date Completed information. In column B, enter the Sample ID. <p>Note: Only alphanumeric characters (0–9 and A to Z), full stops/periods (.), underscores (_), or hyphens (-) are allowed.</p> In column C, enter the RNA sample concentration (C1) in ng/μL determined in the quantification assay. Pipet the indicated volume (X3) of undiluted RNA sample into a new labeled 1.5-mL low-retention microcentrifuge tube, then add the indicated volume of Dilution Solution (Y3). Pipet up and down 5 times to mix, then pulse centrifuge. Proceed directly to “Reverse transcribe the RNA” on page 56. Do not store the diluted RNA samples for longer than necessary. <p>Note: Store the remaining undiluted RNA sample at -90°C to -60°C for up to 5 months.</p>

(continued)

RNA concentration	Dilution procedure
6.01–9.99 ng/μL	<p>Perform Normalization Dilution #1 and Normalization Dilution #2 to a final concentration of 1.43 ng/μL.</p> <p>Note: The default Sample Volume (X1) is 2 μL. When all volumes are ≥2 μL, red highlighted table cells turn white, indicating sufficient volumes for accurate pipetting have been met.</p> <ol style="list-style-type: none"> 1. Open SFW0000786_ODxTT_SampleDilution_Calculator_3.02.xlsm in Microsoft™ Excel™, select the Calculator worksheet tab, then enter the Operator and Date Completed information. 2. In column B, enter the Sample ID. <ul style="list-style-type: none"> Note: Only alphanumeric characters (0–9 and A to Z), full stops/periods (.), underscores (_), or hyphens (-) are allowed. 3. In column C, enter the RNA sample concentration (C1) in ng/μL determined in the quantification assay. 4. Increase the value for Sample Volume (X1) until the Volume of Dilution Solution (Y1) is ≥2 μL. 5. Pipet the indicated volume (X1) of undiluted RNA sample into a new labeled 1.5-mL low-retention microcentrifuge tube, then add the indicated volume of Dilution Solution (Y1). 6. Pipet up and down 5 times to mix, then pulse centrifuge. 7. Pipet the indicated volume (X2) of diluted RNA sample (5 ng/μL) into a new labeled 1.5-mL low-retention microcentrifuge tube, then add the indicated volume of Dilution Solution (Y2). 8. Pipet up and down 5 times to mix, then pulse centrifuge. 9. Proceed directly to “Reverse transcribe the RNA” on page 56. Do not store the diluted RNA samples for longer than necessary. <p>Note: Store the remaining Normalization Dilution #1 (5 ng/μL) and undiluted RNA sample at –90°C to –60°C for up to 5 months.</p>

(continued)

RNA concentration	Dilution procedure
≥10 ng/μL	<p>Perform Normalization Dilution #1 and Normalization Dilution #2 to a final concentration of 1.43 ng/μL.</p> <p>Note: The default Sample Volume (X1) is 2 μL. When all volumes are ≥2 μL, red highlighted table cells turn white, indicating sufficient volumes for accurate pipetting have been met.</p> <ol style="list-style-type: none"> 1. Open SFW0000786_ODxTT_SampleDilution_Calculator_3.02.xlsm in Microsoft™ Excel™, select the Calculator worksheet tab, then enter the Operator and Date Completed information. 2. In column B, enter the Sample ID. <p>Note: Only alphanumeric characters (0–9 and A to Z), full stops/periods (.), underscores (_), or hyphens (-) are allowed.</p> 3. In column C, enter the RNA sample concentration (C1) in ng/μL determined in the quantification assay. 4. Pipet the indicated volume (X1) of undiluted RNA sample into a new labeled 1.5-mL low-retention microcentrifuge tube, then add the indicated volume of Dilution Solution (Y1). 5. Pipet up and down 5 times to mix, then pulse centrifuge. 6. Pipet the indicated volume (X2) of diluted RNA sample (5 ng/μL) into a new labeled 1.5-mL low-retention microcentrifuge tube, then add the indicated volume of Dilution Solution (Y2). 7. Pipet up and down 5 times to mix, then pulse centrifuge. 8. Proceed directly to “Reverse transcribe the RNA” on page 56. Do not store the diluted RNA samples for longer than necessary. <p>Note: Store the remaining Normalization Dilution #1 (5 ng/μL) and undiluted RNA sample at –90°C to –60°C for up to 5 months.</p>

6

Prepare the cDNA

Review the procedural guidelines

Before you begin the procedures in this section, review the procedural guidelines (see “Procedural guidelines” on page 16).

Kit components used in this procedure

Kit component	Box
5X Reaction Mix (red cap) 10X Enzyme Mix (green cap)	Ion Torrent Dx cDNA Synthesis Kit (Part No. A32436, stored at –30°C to –10°C)
Oncomine™ Dx Target RNA Control (white cap; single-use tubes)	Oncomine™ Dx Target RNA Control (Part No. A32443, stored at –90°C to –60°C)
Oncomine™ Dx Target RNA Control Diluent (blue cap; single-use tubes)	Oncomine™ Dx Target RNA Control Diluent (Part No. A38872, stored at –90°C to –60°C)
No Template Control (purple cap)	Ion Torrent Dx No Template Control Kit (Part No. A32444, stored at 15°C to 30°C)

Oncomine™ Dx Target Test Controls lot matching

The following controls and diluent must be lot-matched with each other for use.

Component	Part No.	Storage
Oncomine™ Dx Target DNA Control	A44913	–30°C to –10°C
Oncomine™ Dx Target RNA Control	A32443	–90°C to –60°C
Oncomine™ Dx Target RNA Control Diluent	A38872	–90°C to –60°C
Ion Torrent Dx No Template Control Kit	A32444	15°C to 30°C

Thaw then dilute frozen samples

If DNA and RNA samples were frozen for storage, thaw them at room temperature until no ice crystals are present, then dilute them as described in “Dilute the samples” on page 47 before proceeding. Transfer diluted samples to 2–8°C storage until use.

Note: Freeze-thaw samples no more than 3 times.

Reverse transcribe the RNA

Perform the following steps in a laminar flow hood.

Prepare a master mix for up to 16 cDNA synthesis reactions.

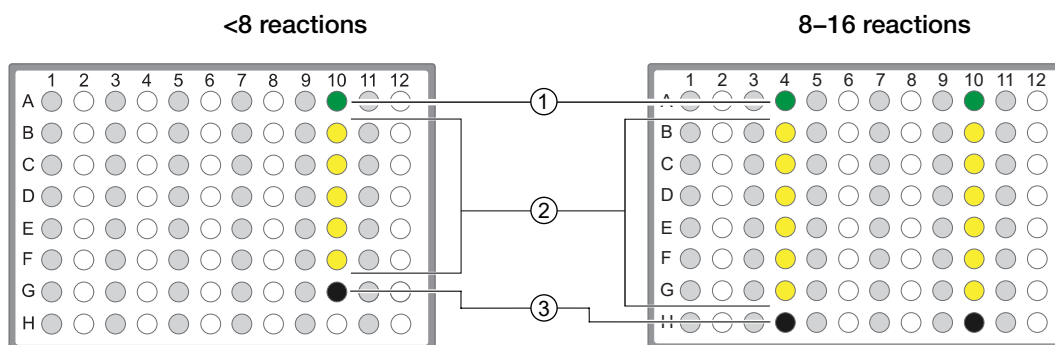
IMPORTANT! For every 6 samples, there must be one No Template Control (NTC) and one Oncomine™ Dx Target RNA Control.

1. Vortex the No Template Control (purple cap) and Dilution Solution (black cap) for ~5 seconds each, then pulse centrifuge.
2. Flick the prediluted RNA sample (1.43 ng/μL), the single-use Oncomine™ Dx Target RNA Control tube (white cap), and the 10X Enzyme Mix (green cap) 4 times each to mix, then pulse centrifuge.

IMPORTANT! Do not vortex the prediluted RNA sample.

3. Label a MicroAmp™ Optical 96-well Reaction Plate with "RNA/cDNA".
4. Place the labeled 96-well plate on a 2–8°C aluminum cold block, then set up the reactions in the designated wells of the plate. Configure the plate for <8 reactions or 8–16 reactions as shown in the figure.

IMPORTANT! Include one No Template Control and one Oncomine™ Dx Target RNA Control well for each column, as illustrated.



- ① Diluted Oncomine™ Dx Target RNA Control
- ② Prediluted RNA samples
- ③ No Template Control

Note: If you are preparing >8 reactions, skip columns to prevent cross-contamination.

5. To each No Template Control well, add 7 μ L of No Template Control.
6. Flick the Oncomine™ Dx Target RNA Control and Oncomine™ Dx Target RNA Control Diluent tubes 4 times to mix, then pulse centrifuge to collect.
7. Dilute the Oncomine™ Dx Target RNA Control. Add the following components to a 1.5-mL low-retention microcentrifuge tube in the order indicated

Note:

- When preparing multiple replicates of the control, create a separate dilution for each replicate.
- The Oncomine™ Dx Target RNA Control and Oncomine™ Dx Target RNA Control Diluent tubes are single-use only. Discard unused volume.

	Order	Component	Volume per reaction
<input type="checkbox"/>	1	Oncomine™ Dx Target RNA Control (white cap)	3 μ L
<input type="checkbox"/>	2	Oncomine™ Dx Target RNA Control Diluent (blue cap)	72 μ L

8. Flick the diluted Oncomine™ Dx Target RNA Control tube 4 times to mix, then pulse centrifuge to collect.
9. To each Oncomine™ Dx Target RNA Control well, add the following components in the order indicated:

	Order	Component	Volume per reaction
<input type="checkbox"/>	1	Diluted Oncomine™ Dx Target RNA Control	3 μ L
<input type="checkbox"/>	2	Dilution Solution (black cap)	4 μ L

10. For each RNA sample reaction, add 7 μ L of prediluted RNA sample into the designated well.

Note: Do not exceed 7 μ L of prediluted RNA, which is equivalent to 10 ng.

11. Prepare a master mix for n+1 reactions. Add the following components to a 1.5-mL low-retention microcentrifuge tube:

	Component	Volume per reaction
<input type="checkbox"/>	5X Reaction Mix (red cap)	(n+1) \times 2 μ L
<input type="checkbox"/>	10X Enzyme Mix (green cap)	(n+1) \times 1 μ L
<input type="checkbox"/>	Total	(n+1) \times 3 μL

12. Flick the master mix tube 4 times to mix, then pulse centrifuge to collect.
13. Pipet 3 μ L of the master mix into each RNA sample, No Template Control, and Oncomine™ Dx Target RNA Control well in the 96-well plate.

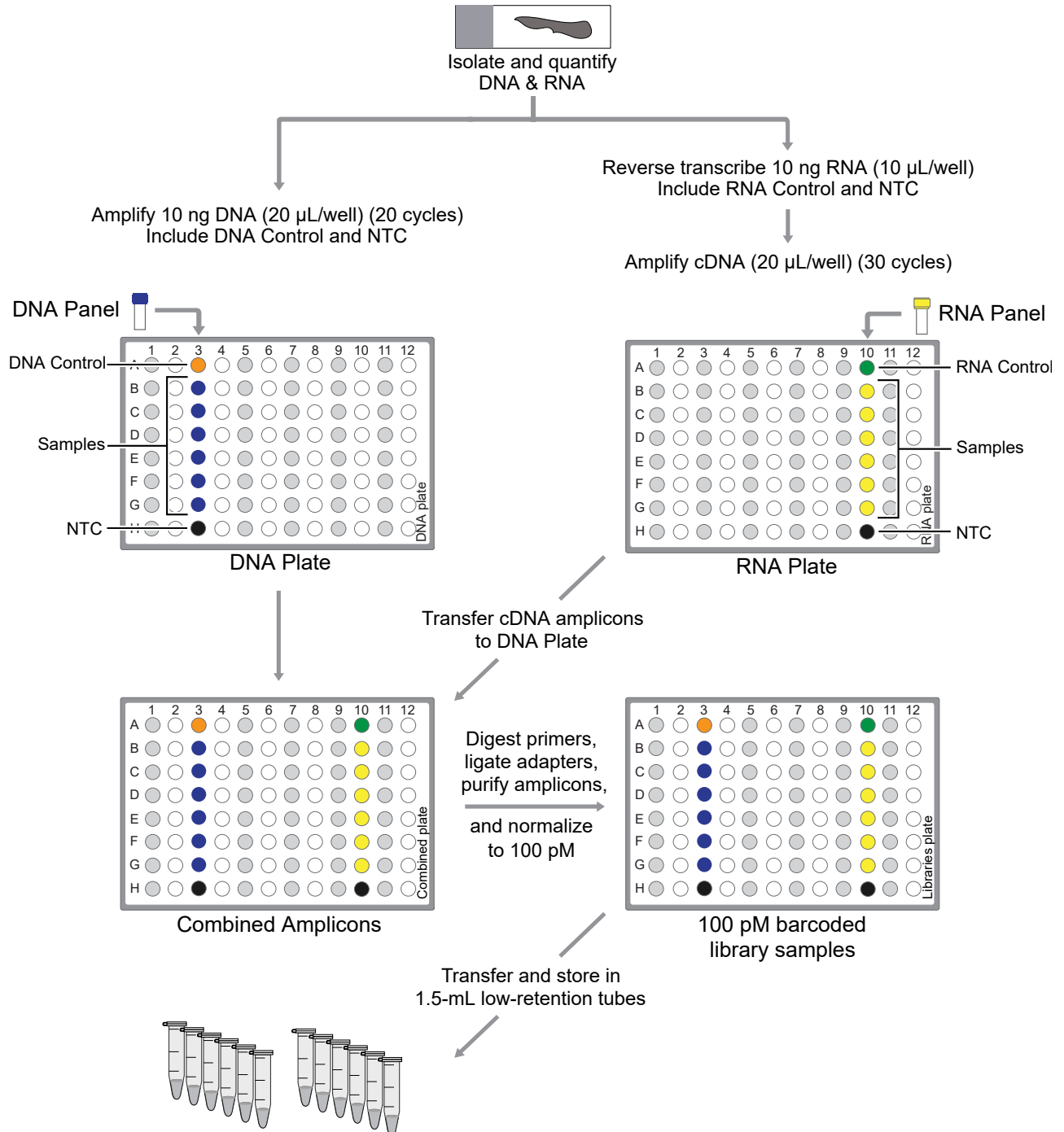
14. Set a pipette to 8 μ L, then pipet the contents of each reaction well up and down 5 times to mix.
15. Seal the plate with an Adhesive PCR Plate Seal, then centrifuge the plate at 100 rcf for 30 seconds.
16. Load the plate in the Veriti™ Dx 96-well Thermal Cycler, then select the **1 ODxTT cDNA Synthesis** program. Select **View**, then confirm that the steps in the program match those in the table below.

	Temperature	Time
<input type="checkbox"/>	42°C	30 minutes
<input type="checkbox"/>	85°C	5 minutes
<input type="checkbox"/>	10°C	Hold (up to 1 hour)

17. When you have confirmed the steps, run the program.

STOPPING POINT The cDNA can be held on the thermal cycler up to 1 hour. Store at -30°C to -10°C for up to 7 days.

Library preparation workflow diagram



Prepare a library batch in Torrent Suite™ Dx Software

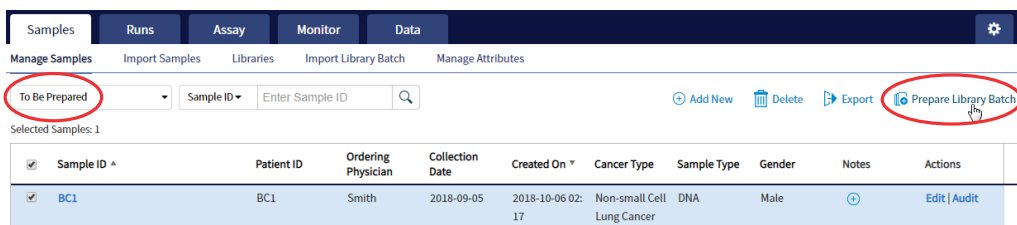
In Torrent Suite™ Dx Software, samples that are entered into the software are placed in library batches for processing and tracking. A library batch consists of a group of libraries that are prepared at the same time.

Note:

- Each library in a library batch must have a unique library name. When combining libraries in the same run, each must also have a unique barcode.
- Control libraries must be included in the same library batch as the sample library they control for.
- Fields identified with a red asterisk (*) are required.

1. Sign in to Torrent Suite™ Dx Software.
2. Under the **Samples** tab, in the **Manage Samples** screen, select **To Be Prepared** from the **Filter Samples by...** dropdown list to display only those samples that have not been placed in a library batch.

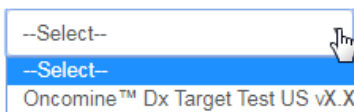
IMPORTANT! Samples that have not been queued for extraction in the software also appear on this tab. Ensure that the samples have been queued for extraction before queuing them for library batch preparation.



3. Select up to 6 samples in the list, then click **Prepare Library Batch**. The **Prepare Library Batch** dialog opens. Required fields are indicated with a red asterisk(*).
4. In the **Select Assay** dropdown list, select **Oncomine™ Dx Target Test**. The assay determines specific parameters of the run, including required controls and post-run data analysis settings.

Prepare Library Batch

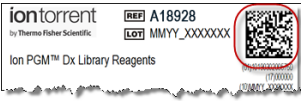

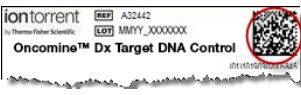
Select Assay:



5. In the following screen, enter a unique identifier for the library batch in the **Library Batch ID** field. Library Batch IDs can only contain alphanumeric characters (0–9 and A to Z), full stop/period (.), underscore (_), and hyphen (-).

- Scan the barcodes from their respective kit boxes into the appropriate fields. Each library batch is associated with a kit lot by scanning the 2D barcode on the appropriate kit box.

IMPORTANT! Check the expiration date on each box. If the kit is expired, select another kit.

Barcode field	Kit	Kit box	Storage	Label scanned
Library Kit Barcode	Ion PGM™ Dx Library Kit	Ion PGM™ Dx Library Reagents	-30°C to -10°C	
Panel Kit Barcode	Oncomine™ Dx Target Test Panel	Oncomine™ Dx Target Test DNA and RNA Panel (box 1 of 3)	-30°C to -10°C	
Control Kit Barcode	Oncomine™ Dx Target Test, Controls, and Diluent Kit	Oncomine™ Dx Target DNA Control (box 2 of 3)	-30°C to -10°C	

- Type a unique library name for each DNA and RNA library in the appropriate field. Library names can only contain alphanumeric characters (0–9 and A to Z), full stop/period (.), underscore (_), and hyphen (-).

Note: The Oncomine™ Dx Target Test Kit requires specific controls, which are automatically listed in the **Prepare Library Batch** dialog.

- Select the Barcode ID of the adapter used to prepare each library. Swap the default barcodes in the dialog between DNA and RNA using the **DNA ↔ RNA** button.

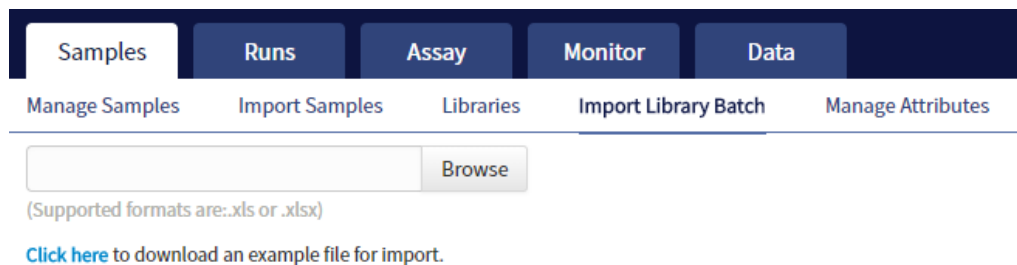
Note: Each library in a library batch must have a different Barcode ID. When preparing the physical libraries, we recommend swapping barcodes between DNA and RNA libraries in consecutive sequencing runs to prevent carryover contamination. See “Alternating barcodes” on page 72.

IMPORTANT! Be careful to ensure that the actual barcodes used to create the libraries match the barcodes that are entered in the **Prepare Library Batch** dialog.

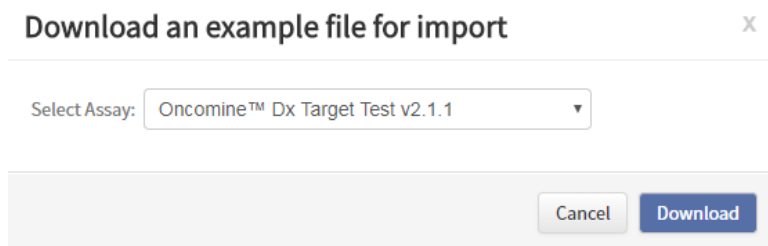
- In the **Input Quantity** field, enter 10 ng for each library.
- Click **Save** to save your selections and close the dialog.
The **Libraries** screen opens, listing the libraries that you created. Libraries that are prepared in the same batch have the same **Library Batch ID**.

Import a library batch

Under the **Samples** tab in the **Import Library Batch** screen, you can import library batch information in the form of an XLS or XLSX file. The import file must include all of the library and kit information that you enter in the **Prepare Library Batch** dialog box.



1. In the **Import Library Batch** screen, below the **Browse** field, click **Click here** to download an example file for import.
2. In the dialog box that opens, select the assay that you are using from the dropdown list, then click **Download**.



The assay name is auto-populated in the Microsoft™ Excel™ template file that downloads to your drive.

3. In the template file, enter or confirm the library batch information.
 - Assay used (auto-populated)
 - Unique Library Batch ID
 - Library kit, control kit, and panel kit barcodes
 - Sample IDs
 - Library names
 - Barcode IDs used for each library and control
 - Nucleic acid type (DNA or RNA)
 - Library input quantity
4. Save the file.
5. Click **Browse**, navigate to the saved file, then select it.

6. Click **Import.**

A progress bar followed by an import report displays. If the import process fails, an error message indicates the reason for failure (for example, an invalid character was used). For additional troubleshooting, see “Library batch import fails” on page 81.

7. Click **Libraries to return to the library batch screen. Your successfully imported library batch is listed.**

Prepare reagents and equipment

- See “Procedural guidelines” on page 16 before setting up the reactions.
- Equilibrate the reagents listed below at room temperature for at least 30 minutes.
 - LIB AMPure™ Reagent
 - LIB Beads
 - LIB Primers
 - LIB Capture
 - LIB Wash Soln
 - LIB Elution Soln
- Place kit components that contain enzymes (LIB HiFi Mix, LIB FuPa, and LIB DNA Ligase) on ice or in a –30°C to –10°C chilled benchtop cold box throughout the procedure until needed. Before use, flick each tube 4 times to mix, then pulse centrifuge.
- Thaw the remaining kit components (except enzymes) at room temperature until no ice is present in the tubes. Vortex for ~5 seconds, then pulse centrifuge before use.
- If there is visible precipitate in the LIB Switch Soln after thawing, vortex for ~5 seconds at room temperature, and pulse centrifuge to collect. Repeat if needed until the solution is clear.

Ion PGM™ Dx Library Kit component lot matching

The two components of the Ion PGM™ Dx Library Kit must be lot-matched with each other for use.

Component	Part No.	Storage
Ion PGM™ Dx Library Reagents	A18928	–30°C to –10°C
Ion PGM™ Dx Library Equalizer™ Reagents	A18929	2°C to 8°C

Oncomine™ Dx Target Test Controls lot matching

The following controls and diluent must be lot-matched with each other for use.

Component	Part No.	Storage
Oncomine™ Dx Target DNA Control	A44913	-30°C to -10°C
Oncomine™ Dx Target RNA Control	A32443	-90°C to -60°C
Oncomine™ Dx Target RNA Control Diluent	A38872	-90°C to -60°C
Ion Torrent Dx No Template Control Kit	A32444	15°C to 30°C

Amplify the cDNA

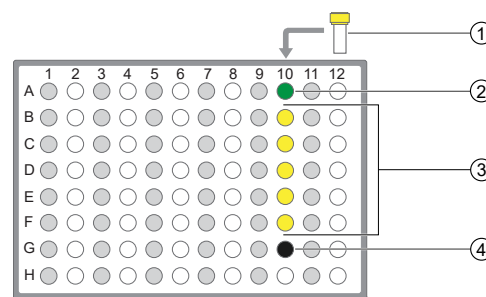
Kit components used in this procedure

Kit component	Box
Oncomine™ Dx Target Test—RNA panel (yellow cap)	Oncomine™ Dx Target Test DNA and RNA Panel (Part No. A32441, stored at -30°C to -10°C)
LIB HiFi Mix (red cap)	Ion PGM™ Dx Library Reagents (Part No. A18928), stored at -30°C to -10°C)

Set up the cDNA amplification reaction (<8 reactions)

If you are preparing <8 reactions, see below. If you are preparing 8–16 reactions, see “Set up the cDNA amplification reaction (8–16 reactions)” on page 65. The number of reactions depends on the configuration of your “RNA/cDNA” plate (prepared as described in “Reverse transcribe the RNA” on page 56).

1. Remove the “RNA/cDNA” plate from the thermal cycler, then centrifuge the plate at 100 rcf for 30 seconds.
2. Transfer the plate to a chilled (2–8°C) 96-well aluminum block.
3. Vortex the Oncomine™ Dx Target Test—RNA panel for ~5 seconds, then pulse centrifuge. Flick the tube of LIB HiFi Mix 4 times to mix, then pulse centrifuge.



- ① Oncomine™ Dx Target Test—RNA panel
- ② Oncomine™ Dx Target RNA Control
- ③ cDNA samples
- ④ No Template Control

- Remove the seal from the plate, then add the following components to each well.

IMPORTANT! The volume of Oncomine™ Dx Target Test—RNA panel is critical and must be accurate.

	Component	Volume
<input type="checkbox"/>	Nuclease-free Water	4 µL
<input type="checkbox"/>	Oncomine™ Dx Target Test—RNA panel (yellow cap)	2 µL
<input type="checkbox"/>	LIB HiFi Mix (red cap)	4 µL
<input type="checkbox"/>	Total volume per well (includes 10 µL from cDNA synthesis)	20 µL

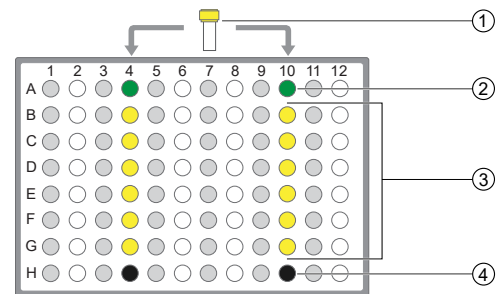
- With the pipettor set to 15 µL, pipet up and down 5 times to mix the contents of each reaction well.
- Proceed to “Amplify the cDNA targets” on page 66.

Set up the cDNA amplification reaction (8–16 reactions)

If you are preparing 8–16 reactions, see below. If you are preparing <8 reactions, see “Set up the cDNA amplification reaction (<8 reactions)” on page 64. The number of reactions depends on the configuration of your “RNA/cDNA” plate (prepared as described in “Reverse transcribe the RNA” on page 56).

For 8–16 amplification reactions (including controls), make a master mix for n+1 reactions, where “n” is the number of reactions you are preparing.

- Remove the “RNA/cDNA” plate from the thermal cycler, then centrifuge the plate at 100 rcf for 30 seconds.
- Transfer the plate to a chilled (2–8°C) 96-well aluminum block.
- Vortex the Oncomine™ Dx Target Test—RNA panel for ~5 seconds, then pulse centrifuge. Flick the tube of LIB HiFi Mix 4 times to mix, then pulse centrifuge.
- Calculate the amounts of the following components needed for n+1 reactions, then add the components to a single 1.5-mL low-retention microcentrifuge tube.



- Oncomine™ Dx Target Test—RNA panel
- Oncomine™ Dx Target RNA Control
- cDNA samples
- No Template Control

IMPORTANT! IMPORTANT: The volume of Oncomine™ Dx Target Test—RNA panel is critical and must be accurate.

	Component	Volume per reaction
<input type="checkbox"/>	Nuclease-Free Water	(n+1) × 4 µL
<input type="checkbox"/>	Oncomine™ Dx Target Test—RNA panel (yellow cap)	(n+1) × 2 µL
<input type="checkbox"/>	LIB HiFi Mix (red cap)	(n+1) × 4 µL
<input type="checkbox"/>	Total	(n+1) × 10 µL

- Vortex the tube for ~5 seconds, then pulse centrifuge to collect.

Note: Keep the master mix on ice or chilled in a 2–8°C benchtop cold box until ready for use.

- Pipet 10 µL of the master mix into each sample or control well in the 96-well plate.
- Set the pipettor to 15 µL, then pipet the contents of each well up and down 5 times to mix.
- Proceed to “Amplify the cDNA targets”.

Amplify the cDNA targets

Note: The Veriti™ Dx 96-well Thermal Cycler, 0.2 mL has been validated with this procedure.

- Seal the 96-well plate with a new adhesive film, then centrifuge the plate at 100 rcf for 30 seconds.
- Load the 96-well plate in the thermal cycler, then select the **2B ODxTT cDNA Target Amp** program. Select **View**, then confirm that the program steps match those listed in the following table:

Stage	Step	Temperature	Time
Hold	Activate the enzyme	98°C	2 minutes
Cycle (30 cycles)	Denature	98°C	15 seconds
	Anneal and extend	60°C	4 minutes
Hold	—	10°C	Hold (up to 24 hours)

- After you have confirmed the steps, run the program.

STOPPING POINT Amplicons can be held in the thermal cycler for up to 24 hours or stored at 2–8°C for up to 1 week. If stored longer than 1 week, prepare new amplicons.

Amplify the DNA

Kit components used in this procedure

Kit component	Box
No Template Control (purple cap)	Ion Torrent Dx No Template Control Kit (Part No. A32444, stored at 15°C to 30°C)
Oncomine™ Dx Target Test—DNA panel (blue cap)	Oncomine™ Dx Target Test DNA and RNA Panel (Part No. A32441, stored at –30°C to –10°C)

(continued)

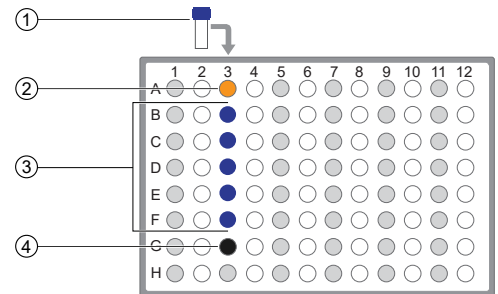
Kit component	Box
LIB HiFi Mix (red cap)	Ion PGM™ Dx Library Reagents (Part No. A18928, stored at –30°C to –10°C)
Dilution Solution (black cap)	Ion Torrent Dx Sample Dilution Kit (Part No. A32439, stored at 15°C to 30°C)
Oncomine™ Dx Target DNA Control (brown cap)	Oncomine™ Dx Target DNA Control (Part No. A44913, stored at –30°C to –10°C)

Set up the DNA amplification reaction (<8 reactions)

If you are preparing <8 reactions, see below. If you are preparing 8–16 reactions, see page 69.

For <8 reactions, set up individual reactions, including a No Template Control (purple cap), an Oncomine™ Dx Target DNA Control (brown cap), and up to 5 clinical samples.

1. Label a 96-well plate "DNA".
2. Place the labeled 96-well plate on a 2–8°C chilled 96-well aluminum block, then set up individual reactions in an odd-numbered column. For every run, include the No Template Control and the Oncomine™ Dx Target DNA Control.
3. Vortex the Oncomine™ Dx Target Test—DNA panel for ~5 seconds, then pulse centrifuge. Flick the tube of LIB HiFi Mix 4 times to mix, then pulse centrifuge.
4. To the No Template Control well, add the following components in the order indicated:



- ① Oncomine™ Dx Target Test—DNA panel
- ② Oncomine™ Dx Target DNA Control
- ③ Prediluted FFPE DNA sample
- ④ No Template Control

	Order	Component	Volume
<input type="checkbox"/>	1	No Template Control (purple cap)	12 µL
<input type="checkbox"/>	2	Oncomine™ Dx Target Test—DNA panel (blue cap)	4 µL
<input type="checkbox"/>	3	LIB HiFi Mix (red cap)	4 µL
<input type="checkbox"/>	—	Total	20 µL

5. To the Oncomine™ Dx Target DNA Control well, add the following components in the order indicated:

	Order	Component	Volume
<input type="checkbox"/>	1	Dilution Solution (black cap)	9 µL
<input type="checkbox"/>	2	Oncomine™ Dx Target DNA Control (brown cap)	3 µL
<input type="checkbox"/>	3	Oncomine™ Dx Target Test—DNA panel (blue cap)	4 µL
<input type="checkbox"/>	4	LIB HiFi Mix (red cap)	4 µL
<input type="checkbox"/>	—	Total	20 µL

Note: The Oncomine™ Dx Target DNA Control tube is single-use only. Discard unused volume.

6. Vortex the prediluted FFPE DNA sample (0.83 ng/µL) for ~5 seconds, then pulse centrifuge to collect.
7. To each sample well, add the following components in the order indicated.

IMPORTANT! If preparing multiple sample libraries, ensure that the appropriate FFPE DNA sample is added to the correct well to avoid patient sample mix-up.

Note: Do not exceed 12 µL of prediluted FFPE DNA, which is equivalent to 10 ng.

	Order	Component	Volume
<input type="checkbox"/>	1	Prediluted sample FFPE DNA (0.83 ng/µL)	12 µL
<input type="checkbox"/>	2	Oncomine™ Dx Target Test—DNA panel (blue cap)	4 µL
<input type="checkbox"/>	3	LIB HiFi Mix (red cap)	4 µL
<input type="checkbox"/>	—	Total	20 µL

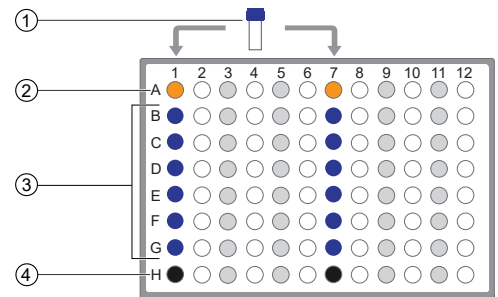
8. Set a 20-µL pipettor to 15 µL, and pipet the contents of each well up and down 5 times to mix.
9. Proceed to “Amplify the DNA targets” on page 70.

Set up the DNA amplification reaction (8–16 reactions)

If you are preparing 8–16 reactions, see below. If you are preparing <8 reactions, see page 67.

For 8–16 reactions (for example, 12 clinical samples plus 4 controls), make a master mix containing every component except prediluted FFPE DNA as follows, where "n" is the number of reactions you are preparing. Include one No Template Control (purple cap) and one Oncomine™ Dx Target DNA Control (brown cap) for each column of reactions as illustrated.

1. Label a 96-well plate "DNA".
2. Place the labeled 96-well plate on a 2–8°C chilled aluminum block, then set up reactions in individual wells in odd-numbered columns of the plate. Skip columns to prevent cross-contamination.
3. To each No Template Control well, add 12 µL of No Template Control (purple cap).
4. To each Oncomine™ Dx Target DNA Control well, add the following components in the order indicated:



- ① Oncomine™ Dx Target Test—DNA panel
- ② Oncomine™ Dx Target DNA Control
- ③ Prediluted FFPE DNA samples
- ④ No Template Control

	Order	Component	Volume per reaction
<input type="checkbox"/>	1	Dilution Solution (black cap)	9 µL
<input type="checkbox"/>	2	Oncomine™ Dx Target DNA Control (brown cap)	3 µL

Note: The Oncomine™ Dx Target DNA Control tube is single-use only. Discard unused volume.

5. Vortex the prediluted FFPE DNA samples (0.83 ng/µL) for ~5 seconds, then pulse centrifuge to collect.
6. To each DNA sample well, add 12 µL prediluted FFPE DNA.

Note: Do not exceed 12 µL of prediluted FFPE DNA, which is equivalent to 10 ng.

7. Vortex the Oncomine™ Dx Target Test—DNA panel for ~5 seconds, then pulse centrifuge. Flick the tube of LIB HiFi Mix 4 times to mix, then pulse centrifuge.
8. Calculate the volume of each component below needed for n+1 reactions, then add that volume to a pre-labeled 1.5-mL low-retention microcentrifuge tube in the order stated:

	Order	Component	Volume
<input type="checkbox"/>	1	Oncomine™ Dx Target Test—DNA panel (blue cap)	(n+1) × 4 µL
<input type="checkbox"/>	2	LIB HiFi Mix (red cap)	(n+1) × 4 µL
<input type="checkbox"/>	—	Total	(n+1) × 8 µL

- Vortex for ~5 seconds, then pulse centrifuge.

Note: Keep the master mix at 2–8°C on ice until ready for use.

- Pipet 8 µL of master mix into each DNA sample, No Template Control, and Oncomine™ Dx Target DNA Control well in the labeled 96-well plate.
- Set a pipettor to 15 µL, then pipet the contents of each well up and down 5 times to mix.
- Proceed to “Amplify the DNA targets”.

Amplify the DNA targets

Note: The Veriti™ Dx 96-well Thermal Cycler, 0.2 mL has been validated with this procedure.

- Seal the 96-well plate with a new adhesive film, then centrifuge the plate at 100 rcf for 30 seconds.
- Load the 96-well plate in the thermal cycler, then select the **3 ODxTT DNA Target Amp** program. Select **View**, and confirm that the program steps match those in the following table:

Stage	Step	Temperature	Time
Hold	Activate the enzyme	99°C	2 minutes
Cycle (20 cycles)	Denature	99°C	15 seconds
	Anneal and extend	60°C	4 minutes
Hold	—	10°C	Hold (up to 24 hours)

- After you have confirmed the steps, run the program.

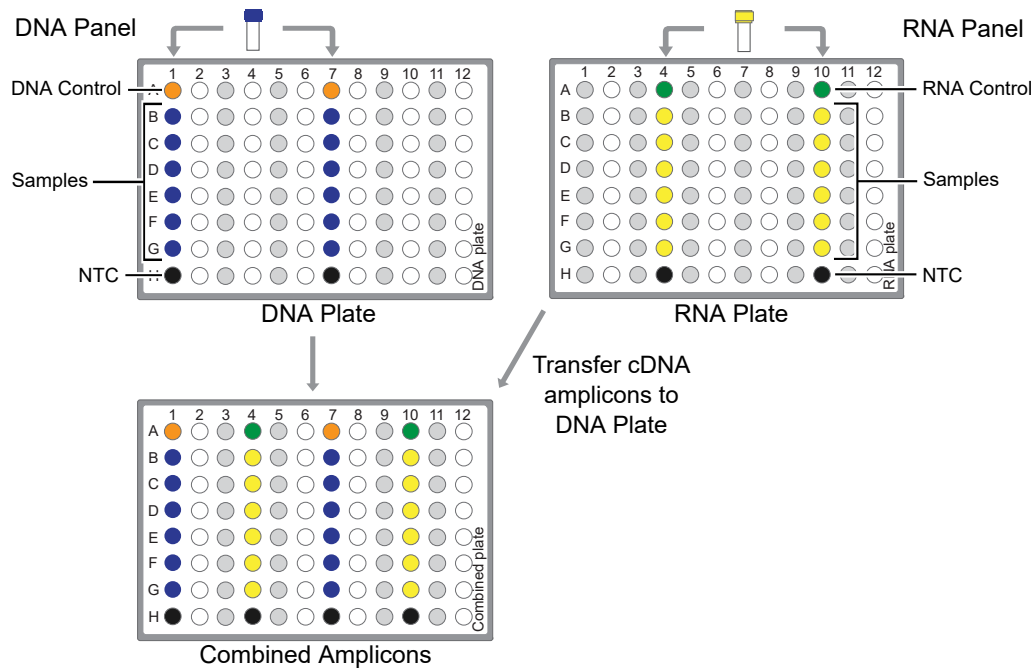
STOPPING POINT Amplicons can be held in the thermal cycler for up to 24 hours or stored at 2–8°C for up to 1 week. If stored longer than 1 week, prepare new amplicons.

Transfer the cDNA amplicons

- After thermal cycling, transfer the 96-well plate to a designated post-PCR preparation area.
- Centrifuge the plates containing amplified cDNA and DNA at 100 rcf for 30 seconds.
- Carefully remove the adhesive film from the plates.

IMPORTANT! Be careful when removing the adhesive film from the plate to minimize cross-contamination.

4. Transfer the cDNA amplicons from the cDNA plate to the corresponding empty wells in even-numbered columns of the DNA plate. Skip columns to prevent cross-contamination, as shown in the example below.



Proceed to “Partially digest amplicons” on page 71.

Partially digest amplicons

1. Place the plate with the amplicons on a 2–8°C cold block.
2. Flick the LIB FuPa tube (green cap) 4 times to mix, then pulse centrifuge to collect.
3. Add 2 μL of LIB FuPa to each reaction well. The total volume is 22 μL per well.

IMPORTANT! LIB FuPa is highly viscous. To avoid carrying over excess enzyme, do not submerge the whole tip in the LIB FuPa solution. Aspirate the solution from just below the surface. The volume is critical and must be accurate. Ensure that no excess solution is added to the sample.

4. Set the pipettor to 15 μL , then slowly pipet the mixture up and down 5 times to mix.
5. Seal the plate with a new adhesive film, then centrifuge at 100 rcf for 30 seconds.

6. Load the plate in the Veriti™ Dx 96-well Thermal Cycler, then select the **4 ODxTT Amplicon Digestion** program. Select **View**, and confirm that the program steps match those listed in the table below:

Temperature	Time
50°C	10 minutes
55°C	10 minutes
60°C	20 minutes
10°C	Hold (for up to 1 hour)

7. After you have confirmed the steps, run the program.

IMPORTANT! Do not leave samples in the thermal cycler for more than 1 hour after cycling.

8. During the run, thaw the LIB Switch Soln (orange cap) and appropriate barcode adapters (white caps) at room temperature for use in subsequent steps.

Alternating barcodes

When preparing libraries, we recommend swapping barcodes between DNA and RNA libraries in consecutive sequencing runs to prevent carryover contamination. The following table provides an example of swapping barcodes between runs.

IMPORTANT! Be careful to confirm that the barcodes used to create the libraries match the barcodes entered in the **Prepare Library Batch** dialog box.

Library type	System Run 1 barcode usage		System Run 2 barcode usage	
	DNA	RNA	DNA	RNA
Positive control	1	9	9	1
Sample	2	10	10	2
Sample	3	11	11	3
Sample	4	12	12	4
Sample	5	13	13	5
Sample	6	14	14	6
Sample	7	15	15	7
No-template control (NTC)	8	16	16	8

Ligate barcode adapters

IMPORTANT! Libraries prepared from DNA and RNA from the same sample must have different barcodes, because the libraries are combined before the amplification reaction on the Ion OneTouch™ Dx Instrument.

1. After thermal cycling, centrifuge the plate at 100 rcf for 30 seconds, then place the plate back on the 2–8°C chilled aluminum block.
2. Vortex the LIB Switch Soln (orange cap) for ~5 seconds, then pulse centrifuge to collect.

IMPORTANT! LIB Switch Soln is highly viscous and must be thoroughly mixed before use. There should be no visible precipitate after vortexing. Inspect the tube and cap carefully for precipitate. If precipitate is visible, secure the cap, invert the tube, then vortex upside down for ~5 seconds or until no visible precipitate is present. Use caution to ensure that the correct volume is delivered while pipetting.

3. Flick the tube of LIB DNA Ligase (clear cap) 4 times to mix, then pulse centrifuge to collect.
4. Ensure that the barcode adapters (BC 1–16) are thawed such that no visible ice is present. Vortex for ~5 seconds, then pulse centrifuge to collect.
5. Carefully remove the adhesive film from the plate, then add the following components to each well containing digested sample in the order shown:

IMPORTANT! When preparing barcoded samples, prevent cross-contamination by opening only one tube of barcode adapter at a time during each addition. We recommend that this step be monitored by a co-technician to prevent sample mix-up and/or cross-contamination.

	Order	Component	Volume
<input type="checkbox"/>	1	LIB Switch Soln (orange cap)	4 µL
<input type="checkbox"/>	2	Barcode adapter (white cap) ^[1]	2 µL
<input type="checkbox"/>	3	LIB DNA Ligase (clear cap)	2 µL
<input type="checkbox"/>	—	Total volume per well (includes 22 µL of sample)	30 µL

^[1] Select from BC 1 through BC 16, based on the sample and your barcode scheme.

6. Set a pipettor to 20 µL, then pipet the volume in each well up and down 5 times.
7. Seal the plate with a new adhesive plate seal, then centrifuge the plate at 100 rcf for 30 seconds.

8. Load the plate in the Veriti™ Dx 96-well Thermal Cycler, then select the **5 ODxTT Adapter Ligation** program. Select **View**, then confirm that the program steps match those listed in the following table.

Temperature	Time
22°C	30 minutes
72°C	10 minutes
10°C	Hold (for up to 1 hour)

9. After you have confirmed the steps, run the program.

IMPORTANT! Do not leave samples in the thermal cycler for more than 1 hour after cycling.

10. During the run, equilibrate the LIB Beads (yellow cap), LIB Capture (violet cap), and LIB Elution Soln to room temperature.

Prepare the LIB HiFi Mix plus LIB Primers mix

- Flick the LIB HiFi Mix (red cap) 4 times to mix, then pulse centrifuge. Keep the LIB HiFi Mix in a -30°C to -10°C chilled benchtop cold box.
- Vortex the LIB Primers (blue cap) for ~5 seconds, then pulse centrifuge.
- Prepare the LIB HiFi Mix plus LIB Primers master mix:
 - <8 libraries—For each library, add components to individual 1.5-mL low-retention microcentrifuge tubes on ice or in a $2-8^{\circ}\text{C}$ chilled benchtop cold box in the following order:

	Order	Component	Volume
<input type="checkbox"/>	1	Nuclease-free Water	40 μL
<input type="checkbox"/>	2	LIB HiFi Mix (red cap)	10 μL
<input type="checkbox"/>	3	LIB Primers (blue cap)	2 μL
<input type="checkbox"/>	—	Total	52 μL

- 8–16 libraries—Calculate the amount of every component needed for $n+1$ libraries, where "n" is the number of libraries being prepared, then add the components to a single 1.5-mL low-retention microcentrifuge tube on ice or in a $2-8^{\circ}\text{C}$ chilled benchtop cold box in the following order:

	Order	Component	Volume
<input type="checkbox"/>	1	Nuclease-free Water	$(n+1) \times 40 \mu\text{L}$
<input type="checkbox"/>	2	LIB HiFi Mix (red cap)	$(n+1) \times 10 \mu\text{L}$
<input type="checkbox"/>	3	LIB Primers (blue cap)	$(n+1) \times 2 \mu\text{L}$
<input type="checkbox"/>	—	Total	$(n+1) \times 52 \mu\text{L}$

4. Flick the master mix 4 times to mix, then pulse centrifuge. Keep at 2–8°C.

Note: You must use the master mix on the same day it was prepared.

Purify the barcode-adapted library

1. Prepare fresh 70% ethanol: combine 230 μL of ethanol with 100 μL of Nuclease-free Water per library, then vortex for 10 seconds to mix.
2. When thermal cycling is complete, centrifuge the 96-well plate at 100 rcf for 30 seconds.
3. Before use, invert the LIB AMPure™ Reagent 10 times, then vortex for 10 seconds until the beads are thoroughly suspended.
4. Carefully remove the adhesive film from the plate, then add 45 μL of LIB AMPure™ Reagent to each well.

IMPORTANT! Ensure that an accurate amount of LIB AMPure™ Reagent is dispensed to each sample, and prevent excess carryover from droplets adhering to the tip.

5. With the pipettor set to 45 μL , pipet up and down 5 times to thoroughly mix the beads in each well. The total volume is 75 μL .
6. Hold the mixture for 5 minutes at room temperature.
7. Place the plate in a DynaMag™ Dx 96-Well Plate Magnet for 3 minutes. The solution in each well must be clear, with beads pelleted to one side.
8. Using a 200- μL pipettor, remove and discard ~75 μL of the supernatant without disturbing the pellet. Use a 20- μL pipettor to remove any remaining supernatant.
9. If you see beads in the pipette tip when removing the supernatant, pipet the supernatant and beads back into their respective wells to re-pellet the beads, then remove and discard the supernatant.
10. Add 150 μL of freshly prepared 70% ethanol to each well.
11. Move the plate from left-to-right on the DynaMag™ Dx 96-Well Plate Magnet, then hold for ~5 seconds to wash and re-pellet the beads.
12. Move the plate from right-to-left on the magnet, then hold for ~5 seconds to wash and re-pellet the beads.
13. Repeat steps 11 and 12 two more times. Keep the plate in the final position on the magnet for 3 minutes or until the solution in each well is clear, with the beads in a pellet to one side.
14. Using a 200- μL pipette, remove and discard ~150 μL of the supernatant without disturbing the pellet. Use a 20- μL pipette to remove any remaining supernatant.

15. Repeat steps 10–14 one more time.
16. Ensure that all the ethanol droplets are removed from the wells. Keeping the plate in the magnet, air-dry the beads at room temperature for 5 minutes.

Amplify the barcode-adapted library

1. Flick the LIB HiFi Mix/LIB Primers master mix (prepared in “Prepare the LIB HiFi Mix plus LIB Primers mix” on page 74) 4 times to mix, then pulse centrifuge.
2. Remove the 96-well plate from the DynaMag™ Dx 96-Well Plate Magnet, then add 52 µL of LIB HiFi Mix/LIB Primers master mix to each well.
3. Set the pipettor to 40 µL, then pipet up and down 10 times to mix until the beads are resuspended.

Note: Visually inspect the sides of the wells to ensure complete resuspension of the beads.

4. Seal the 96-well plate with a new adhesive film. Centrifuge the plate at 100 rcf for 30 seconds.
5. Load the plate in the Veriti™ Dx 96-well Thermal Cycler, then select the **6 ODxTT Library Amplification** program. Select **View**, and confirm that the program steps match those in the table below.

Stage	Temperature	Time
Hold	98°C	2 minutes
7 cycles	98°C	15 seconds
	60°C	1 minute
Hold	10°C	Hold (for up to 30 minutes)

6. After you have confirmed the steps, run the program.

Note: During thermal cycling, you may start to prepare the LIB Beads as described in the next procedure.

Prepare the LIB Beads

Note: LIB Beads must be freshly prepared before every use.

1. Equilibrate the LIB Beads (yellow cap) to room temperature, vortex for 10 seconds or until resuspended, then pulse centrifuge to collect.
2. For each library, combine 3 µL of LIB Beads and 6 µL of LIB Wash Soln (clear cap) in a 1.5-mL low-retention microcentrifuge tube, as follows:
 - For 1–3 libraries, prepare a separate tube of beads and wash solution per library.

- For ≥ 4 libraries, prepare a master mix as shown below.

Number of libraries	Number of reactions to prepare in master mix	Volume of LIB Beads to add	Volume of LIB Wash Soln to add
4–7	$n + 0.5$	$(n + 0.5) \times 3 \mu\text{L}$	$(n + 0.5) \times 6 \mu\text{L}$
8	9	27 μL	54 μL
9–16	$n + 2$	$(n + 2) \times 3 \mu\text{L}$	$(n + 2) \times 6 \mu\text{L}$

- Vortex each tube for ~5 seconds to mix, then pulse centrifuge to collect any beads present on the lid of the tube.
- Place the tube in the DynaMag™ Dx 16 2-mL Magnet for 1 minute.
- Carefully remove and discard the supernatant without disturbing the pellet.
- Remove the tube from the magnet, then add the same volume of LIB Wash Soln as added in step 2.
- Resuspend by pipetting up and down 5 times.
- Keep the prepared beads at room temperature and use them on the same day.

Add LIB Capture to the amplified sample library

- Confirm that the LIB Capture (violet cap) is at room temperature, vortex the tube for ~5 seconds, then pulse centrifuge to collect.
- When thermal cycling is complete, centrifuge the 96-well plate at 100 rcf for 30 seconds.
- Carefully remove the adhesive film from the plate, then add 10 μL of LIB Capture (violet cap) to each well.

IMPORTANT! Accurate volume transfer in this step is critical. Ensure that no excess LIB Capture is carried on the pipette tip by aspirating the solution from just below the surface.

- Set the pipettor to 40 μL , then pipet the mixture up and down 5 times to mix.
- Hold at room temperature for 5 minutes.

Add the LIB Beads and wash

- Mix the prepared LIB Beads by pipetting up and down 5 times, or until the beads are resuspended.
- Add 6 μL of washed LIB Beads to each well.
- Set the pipettor to 40 μL , then pipet the mixture up and down 5 times to mix.
- Hold at room temperature for 5 minutes.

5. Place the 96-well plate in the DynaMag™ Dx 96-Well Plate Magnet for 3 minutes. The solution should be clear.
6. Using a 200- μ L pipette, remove, then discard \sim 68 μ L of the supernatant without disturbing the pellet. Use a 20- μ L pipette to remove any remaining supernatant.
7. Add 150 μ L of LIB Wash Soln to each well.
8. Move the 96-well plate from left-to-right on the DynaMag™ Dx 96-Well Plate Magnet, then hold for 5 seconds to wash and re-pellet the beads.
9. Move the 96-well plate from right-to-left on the magnet, then hold for 5 seconds to wash and re-pellet the beads.
10. Repeat steps 8 and 9 two more times.
11. With the 96-well plate still in the magnet, use a 200- μ L pipette to remove and discard \sim 150 μ L of the supernatant without disturbing the pellet.
12. Repeat the bead wash as described in steps 7–11.
13. Use a 20- μ L pipette to remove any remaining LIB Wash Soln by pipetting without disturbing the pellet.

Elute the library

1. Remove the plate from the plate magnet, then add 100 μ L of LIB Elution Soln to each pellet. Set the pipettor to 80 μ L, then pipet up and down at least 10 times until the beads are resuspended.
2. Seal the plate with a new adhesive film, then centrifuge at 100 rcf for 30 seconds.
3. If beads pellet at the bottom of the wells:
 - a. Carefully remove the adhesive film, and gently resuspend the pellet by pipetting up and down until resuspended.

IMPORTANT! Ensure that the sample remains at the bottom of the well. Avoid introducing bubbles while pipetting.

- b. Seal the plate with a new adhesive film.
4. Load the plate in the Veriti™ Dx 96-well Thermal Cycler, then select the **7 ODxTT Library Elution** program. Select **View** and confirm that the program matches the following table.

Temperature	Time
35°C	5 minutes

5. After you have confirmed the steps, run the program.

IMPORTANT! Remove the plate from the thermal cycler immediately after the 5-minute incubation is complete.

6. During cycling, label a 1.5-mL low-retention microcentrifuge tube for each library. Alternatively, if you are proceeding directly to pooling libraries, label a tube for each library pool.
7. Remove the plate from the thermal cycler, then centrifuge the plate at 100 rcf for 30 seconds.



CAUTION! The sample block and plate are hot. Use care when handling the plate to avoid being burned.

8. Place the plate in the DynaMag™ Dx 96-Well Plate Magnet, then hold at room temperature for 3 minutes. Confirm that the solution is clear.
9. Carefully remove the adhesive film, then transfer the supernatant containing the equalized library (~100- μ L total volume) to a labeled 1.5-mL low-retention microcentrifuge tube. The final concentration of each library is ~100 pM.

STOPPING POINT The eluted libraries can be stored at -30°C to -10°C for up to 30 days. If stored for longer than 30 days, prepare new libraries.



Troubleshooting

Troubleshooting—Sample preparation

Observation	Possible cause	Recommended action
Slide cracked during scraping	Too much pressure was used during scraping.	Repeat the extraction using a fresh slide if possible. Processing a cracked or broken slide can pose a safety hazard to the operator.
Low-bind Elution tube cap breaks off Details:	Low-bind Elution tubes were not properly aligned in the microcentrifuge prior to centrifugation.	Transfer eluted samples to new, pre-labeled Low-bind Elution tubes. Extra Low-bind Elution tubes are provided, however 1.5-mL snap-cap low-retention microcentrifuge tubes may also be used.
DNA/RNA quantification values are not returned	The signal for the sample was oversaturated.	Dilute the DNA and RNA samples with Dilution Solution. Prepare new standards and repeat the quantification assay.
	The signal from the sample was too low.	Prepare new standards and repeat the quantification assay. If the low signal persists, repeat the extraction with increased tissue input (for example, more sections) if available.
DNA/RNA samples do not meet the minimum concentration requirement	Quantification assays performed incorrectly.	Prepare new standards and repeat the quantification assay.
	Insufficient tissue was used in the extraction.	Repeat the extraction with increased tissue input (i.e., more sections) if possible. If the tissue was collected via macrodissection of a resection/surgical biopsy sample, repeat the macrodissection and DNA/RNA extraction with more than two 5-micron sections if available. If only two 5-micron sections remain from the sample, repeat the DNA/RNA extraction with the remaining sections without macrodissection. Note: To proceed with library preparation, both the DNA and RNA from a single sample extraction must meet the minimum concentration requirement. Do not use DNA from one extraction with RNA from a different extraction.
	Elution Solution cooled below 95°C.	Keep the Elution Solution in a 95°C heat block throughout the procedure, including when pipetting.

Observation	Possible cause	Recommended action
R ² values do not meet minimum requirement	Standards were not prepared correctly.	Prepare new standards and repeat the quantification assay.
Batch sample import fails	One or more entries in the sample-import spreadsheet contains special characters, lines breaks, unexpected spaces, incorrect entry length, incorrect date formatting, or other formatting errors.	Check each entry for correct formatting, correct any errors, and repeat the import.
	Blank rows were copied into the sample-import template file from a different source.	Rows that appear empty may contain hidden formatting that conflicts with the import function. Start with a clean sample-import template file, and be careful to copy only those rows that contain actual data.
	The sample import spreadsheet contains a nonunique Sample ID.	Every Sample ID in the software must be unique. Make sure the spreadsheet does not contain any duplicate IDs, and repeat the import. Note that the system check is not case-sensitive, so a Sample ID of ABC1 conflicts with abc1.
	The headings in the sample import spreadsheet do not match the sample attributes in the software.	The headings must match the sample attributes in the software exactly. Check the headings for spelling or other errors.
Library batch import fails	One or more entries in the library batch import spreadsheet contains special characters, lines breaks, unexpected spaces, incorrect entry length, incorrect date formatting, or other formatting errors.	Check each entry for correct formatting, correct any errors, and repeat the import.
	Blank rows were copied into the library batch import template file from a different source.	Rows that appear empty can contain hidden formatting that conflicts with the import function. Start with a clean library batch import template file, and be careful to copy only those rows that contain actual data.
	The library batch import spreadsheet contains a nonunique Library Batch ID.	Every Library Batch ID in the software must be unique. Ensure that the spreadsheet does not contain any duplicate IDs, and repeat the import. Note that the system check is not case-sensitive, so a Library Batch ID of ABC1 conflicts with abc1.



Observation	Possible cause	Recommended action
Library batch import fails <i>(continued)</i>	A Sample ID entered in the library batch import spreadsheet does not match a Sample ID listed in the Manage Samples screen.	Ensure that the Sample IDs entered into the spreadsheet are correct and match an existing sample ID added to the software.
	The Barcode ID name format does not exactly match the format that is used in the Prepare Library Batch dialog box.	Use the following Barcode ID name format: IonDx-1 through IonDx-16.
	An invalid library, control, or panel kit barcode has been entered in the spreadsheet.	Ensure that you have correctly entered a valid kit barcode in the appropriate cell of the spreadsheet.

Warnings and alarms—Veriti™ Dx 96-well Thermal Cycler

Observation	Possible cause	Recommended action
"Fatal Error" message displayed by Veriti™ Dx 96-well Thermal Cycler	Various	For assistance, contact Technical Support (see Appendix E, "Customer and technical support"). Refer to the <i>Veriti™ Dx 96-well Thermal Cycler User Guide</i> (Pub. No. 4453697) for general troubleshooting information for this instrument.



Performance characteristics

For performance characteristics of the Oncomine™ Dx Target Test Kit, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

For performance characteristics of the Ion PGM™ Dx System, see the *Ion PGM™ Dx System Performance Characteristics User Guide* (Pub. No. MAN0018763).



WARNING! GENERAL SAFETY. Using this product in a manner not specified in the user documentation may result in personal injury or damage to the instrument or device. Ensure that anyone using this product has received instructions in general safety practices for laboratories and the safety information provided in this document.

- Before using an instrument or device, read and understand the safety information provided in the user documentation provided by the manufacturer of the instrument or device.
- Before handling chemicals, read and understand all applicable Safety Data Sheets (SDSs) and use appropriate personal protective equipment (gloves, gowns, eye protection, and so on). To obtain SDSs, see the “Documentation and Support” section in this document.

Biological hazard safety



WARNING! BIOHAZARD. Biological samples such as tissues, body fluids, infectious agents, and blood of humans and other animals have the potential to transmit infectious diseases. Conduct all work in properly equipped facilities with the appropriate safety equipment (for example, physical containment devices). Safety equipment can also include items for personal protection, such as gloves, coats, gowns, shoe covers, boots, respirators, face shields, safety glasses, or goggles. Individuals should be trained according to applicable regulatory and company/ institution requirements before working with potentially biohazardous materials. Follow all applicable local, state/provincial, and/or national regulations. The following references provide general guidelines when handling biological samples in laboratory environment.

- U.S. Department of Health and Human Services, *Biosafety in Microbiological and Biomedical Laboratories (BMBL)*, 5th Edition, HHS Publication No. (CDC) 21-1112, Revised December 2009; found at:
<https://www.cdc.gov/labs/pdf/CDC-BiosafetymicrobiologicalBiomedicalLaboratories-2020-P.pdf>
- World Health Organization, *Laboratory Biosafety Manual*, 3rd Edition, WHO/CDS/CSR/LYO/2004.11; found at:
www.who.int/publications/i/item/9789240011311



Chemical safety



WARNING! GENERAL CHEMICAL HANDLING. To minimize hazards, ensure laboratory personnel read and practice the general safety guidelines for chemical usage, storage, and waste provided below. Consult the relevant SDS for specific precautions and instructions:

- Read and understand the Safety Data Sheets (SDSs) provided by the chemical manufacturer before you store, handle, or work with any chemicals or hazardous materials. To obtain SDSs, see the “Documentation and Support” section in this document.
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing).
- Minimize the inhalation of chemicals. Do not leave chemical containers open. Use only with adequate ventilation (for example, fume hood).
- Check regularly for chemical leaks or spills. If a leak or spill occurs, follow the manufacturer's cleanup procedures as recommended in the SDS.
- Handle chemical wastes in a fume hood.
- Ensure use of primary and secondary waste containers. (A primary waste container holds the immediate waste. A secondary container contains spills or leaks from the primary container. Both containers must be compatible with the waste material and meet federal, state, and local requirements for container storage.)
- After emptying a waste container, seal it with the cap provided.
- Characterize (by analysis if necessary) the waste generated by the particular applications, reagents, and substrates used in your laboratory.
- Ensure that the waste is stored, transferred, transported, and disposed of according to all local, state/provincial, and/or national regulations.
- **IMPORTANT!** Radioactive or biohazardous materials may require special handling, and disposal limitations may apply.

Precaution—strong magnet




















Note: Do not substitute non-IVD labeled magnets for the DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet, provided with Ion PGM™ Dx System.

The DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet contain very strong permanent magnets. People wearing a pacemaker or any other medical magnetized implant should not use this product unless advised by a health professional; the implant could be affected or damaged by exposure to a strong magnetic field. Keep tools and objects that could be damaged by the magnetic field out of the working area. This includes, but is not restricted to, credit cards and other products containing magnetic recording devices. Keep away from delicate instruments, watches, electronic equipment, displays and monitors. The magnet may attract steel or other magnetic material with high mechanical forces. Take care during handling. Avoid contact between two magnets. Do not pull the magnets apart if contact has been made; twist off to prevent damage to the unit or fingers. The Health and Safety Officer should take all necessary steps and full responsibility to ensure that the precautions and statements are followed and adhered to.



Medical device symbols

The following table describes symbols that may be displayed on product labels. The symbols that are used on labels conform to standards BS EN ISO 15223-1:2016 and FDA 21 CFR 809.10 "Labeling for in vitro diagnostic products".

Symbol	Description	Symbol	Description
	MANUFACTURER		CONTAINS SUFFICIENT FOR <n> TESTS
	DATE OF MANUFACTURE		USE BY
	BATCH CODE		CATALOG NUMBER
	SERIAL NUMBER		FRAGILE, HANDLE WITH CARE
	LOWER LIMIT OF TEMPERATURE		PROTECT FROM LIGHT
	UPPER AND LOWER LIMITS OF TEMPERATURE		UPPER LIMIT OF TEMPERATURE
	DO NOT REUSE		BIOLOGICAL RISKS
	CAUTION, CONSULT ACCOMPANYING DOCUMENTS		CONSULT INSTRUCTIONS FOR USE
	UPPER AND LOWER LIMITS OF HUMIDITY		OBSERVE PRECAUTIONS FOR HANDLING ELECTROSTATIC SENSITIVE DEVICES
	IN VITRO DIAGNOSTIC MEDICAL DEVICE		



Supplemental sample dilution information

We recommend that you use the Sample Dilution Calculator (**SFW0000786_ODxTT_SampleDilution_Calculator_3.02.xlsm**) when preparing libraries (see “Dilute the samples” on page 47).

Dilute the samples (manual calculation)

If you are not using the Sample Dilution Calculator, dilute DNA samples to a final concentration of 0.83 ng/μL and RNA samples to a final concentration of 1.43 ng/μL, by manually calculating dilutions as follows.

IMPORTANT! Do not perform the following dilution procedure until you are ready to proceed directly to reverse transcription and library preparation.

1. Label two new 1.5-mL low-retention microcentrifuge tubes, one for the DNA sample and the other for the RNA sample. Place the tubes in a pre-chilled benchtop cold box until needed.
2. Use the DNA and RNA sample concentrations (ng/μL) determined in the quantification assays to calculate the volume (X) of each RNA and DNA sample required for 10 ng of sample plus 10% overage. Use the following formula:
$$1.1 \times (10 \text{ ng/DNA or RNA sample concentration in ng/}\mu\text{L}) = X \text{ }\mu\text{L of DNA or RNA}$$

Note: See “Example dilution calculations” on page 88.

3. Calculate the volume (Y) of Dilution Solution required to yield a correctly diluted sample using the following formulas:
DNA samples: $(11 \text{ ng}/0.83 \text{ ng/}\mu\text{L}) - X \text{ }\mu\text{L of DNA} = Y \text{ }\mu\text{L of Dilution Solution}$
RNA samples: $(11 \text{ ng}/1.43 \text{ ng/}\mu\text{L}) - X \text{ }\mu\text{L of RNA} = Y \text{ }\mu\text{L of Dilution Solution}$

IMPORTANT!

- If the sample volume (X) from step 2 is <2.0 μL, use 2.0 μL of the sample and adjust the volume of Dilution Solution accordingly.
 - If volume of Dilution Solution (Y) from step 3 is <2.0 μL, increase the amount of the DNA and/or RNA sample volume until the required volume of Dilution Solution is ≥2 μL. See “Example dilution calculations” on page 88.
-
4. For each DNA or RNA sample, pipet the calculated Y μL of Dilution Solution into the appropriate labeled 1.5-mL low-retention microcentrifuge tube from step 1.
 5. Add the calculated X μL of DNA or RNA sample into the appropriate labeled tube.

6. Pipet up and down 5 times to mix, then pulse centrifuge.
7. Place the diluted DNA and RNA samples back in the chilled benchtop cold box or in a 2–8°C refrigerator, then proceed immediately to “Reverse transcribe the RNA” on page 56.

IMPORTANT! Proceed directly to reverse transcription and then library preparation. Do not store the diluted DNA and RNA samples for longer than necessary.

STOPPING POINT Store the remaining undiluted DNA sample at –30°C to –10°C for up to 12 months, and the remaining undiluted RNA sample at –90°C to –60°C for up to 5 months.

Example dilution calculations

Table 2 Example calculation if the sample volume is $\geq 2 \mu\text{L}$

		DNA concentration = 3 ng/ μL	RNA concentration = 4 ng/ μL
1	Sample volume calculation	$1.1 \times [10 \text{ ng}/(3 \text{ ng}/\mu\text{L})] = 3.67 \mu\text{L DNA sample volume}$	$1.1 \times [10 \text{ ng}/(4 \text{ ng}/\mu\text{L})] = 2.75 \mu\text{L RNA sample volume}$
2	Dilution Solution calculation	$(11 \text{ ng}/0.83 \text{ ng}/\mu\text{L}) - 3.67 \mu\text{L DNA sample} = 9.58 \mu\text{L of Dilution Solution}$	$(11 \text{ ng}/1.43 \text{ ng}/\mu\text{L}) - 2.75 \mu\text{L RNA sample} = 4.90 \mu\text{L of Dilution Solution}$
3	Final concentration check	$(3.67 \mu\text{L} \times 3 \text{ ng}/\mu\text{L}) / (3.67 \mu\text{L} + 9.58 \mu\text{L}) = 0.83 \text{ ng}/\mu\text{L}$	$(2.75 \mu\text{L} \times 4 \text{ ng}/\mu\text{L}) / (2.75 \mu\text{L} + 4.94 \mu\text{L}) = 1.43 \text{ ng}/\mu\text{L}$

Table 3 Example calculation if the sample volume is $< 2 \mu\text{L}$

		DNA concentration = 15 ng/ μL	RNA concentration = 14 ng/ μL
1	Sample volume calculation	$1.1 \times [10 \text{ ng}/(15 \text{ ng}/\mu\text{L})] = 0.73 \mu\text{L DNA sample volume}$	$1.1 \times [10 \text{ ng}/(14 \text{ ng}/\mu\text{L})] = 0.79 \mu\text{L RNA sample volume}$
2	Sample volume adjustment ($\times 3$)	$0.73 \mu\text{L of sample} \times 3 = 2.19 \mu\text{L DNA sample volume}$	$0.79 \mu\text{L of sample} \times 3 = 2.37 \mu\text{L RNA sample volume}$
3	Dilution Solution calculation with adjustment	$[(11 \text{ ng}/0.83 \text{ ng}/\mu\text{L}) \times 3] - 2.19 \mu\text{L DNA sample} = 37.6 \mu\text{L of Dilution Solution}$	$[(11 \text{ ng}/1.43 \text{ ng}/\mu\text{L}) \times 3] - 2.37 \mu\text{L RNA sample} = 20.7 \mu\text{L of Dilution Solution}$
4	Final concentration check	$(2.19 \mu\text{L} \times 15 \text{ ng}/\mu\text{L}) / (2.19 \mu\text{L} + 37.6 \mu\text{L}) = 0.83 \text{ ng}/\mu\text{L}$	$(2.37 \mu\text{L} \times 14 \text{ ng}/\mu\text{L}) / (2.37 \mu\text{L} + 20.7 \mu\text{L}) = 1.43 \text{ ng}/\mu\text{L}$



Table 4 Example calculation if the Dilution Solution volume is <2 µL

		DNA concentration = 0.9 ng/µL	RNA concentration = 1.8 ng/µL
1	Sample volume calculation	$1.1 \times [10 \text{ ng}/(0.9 \text{ ng}/\mu\text{L})] = 12.22 \text{ }\mu\text{L DNA sample volume}$	$1.1 \times [10 \text{ ng}/(1.8 \text{ ng}/\mu\text{L})] = 6.11 \text{ }\mu\text{L RNA sample volume}$
2	Dilution Solution calculation	$(11 \text{ ng}/0.83 \text{ ng}/\mu\text{L}) - 12.22 \text{ }\mu\text{L DNA sample} = 1.03 \text{ }\mu\text{L of Dilution Solution}$	$(11 \text{ ng}/1.43 \text{ ng}/\mu\text{L}) - 6.11 \text{ }\mu\text{L RNA sample} = 1.58 \text{ }\mu\text{L of Dilution Solution}$
3	Dilution Solution adjustment ($\times 2$)	$1.03 \text{ }\mu\text{L of Dilution Solution} \times 2 = 2.06 \text{ }\mu\text{L of Dilution Solution}$	$1.58 \text{ }\mu\text{L of Dilution Solution} \times 2 = 3.16 \text{ }\mu\text{L of Dilution Solution}$
4	Sample volume adjustment ($\times 2$)	$12.22 \text{ }\mu\text{L of sample} \times 2 = 24.44 \text{ }\mu\text{L DNA sample volume}$	$6.11 \text{ }\mu\text{L of sample} \times 2 = 12.22 \text{ }\mu\text{L RNA sample volume}$
5	Dilution Solution calculation with adjustment	$[(11 \text{ ng}/0.83 \text{ ng}/\mu\text{L}) \times 2] - 24.44 \text{ }\mu\text{L DNA sample} = 2.06 \text{ }\mu\text{L of Dilution Solution}$	$[(11 \text{ ng}/1.43 \text{ ng}/\mu\text{L}) \times 2] - 12.22 \text{ }\mu\text{L RNA sample} = 3.16 \text{ }\mu\text{L of Dilution Solution}$
6	Final concentration check	$(24.44 \text{ }\mu\text{L} \times 0.9 \text{ ng}/\mu\text{L}) / (24.44 \text{ }\mu\text{L} + 2.06 \text{ }\mu\text{L}) = 0.83 \text{ ng}/\mu\text{L}$	$(12.22 \text{ }\mu\text{L} \times 1.8 \text{ ng}/\mu\text{L}) / (12.22 \text{ }\mu\text{L} + 3.16 \text{ }\mu\text{L}) = 1.43 \text{ ng}/\mu\text{L}$



Customer and technical support

Visit [thermofisher.com/support](https://www.thermofisher.com/support) for the latest in services and support, including:

- Worldwide contact telephone numbers
- Product support
- Order and web support
- Safety Data Sheets (SDSs; also known as MSDSs)

Additional product documentation, including user guides and Certificates of Analysis, are available by contacting Customer Support.

Obtaining Certificates of Analysis

The Certificate of Analysis provides detailed quality control and product qualification information for each product. Certificates of Analysis are printed and shipped with the product.

Obtaining Certificates of Conformance

The Certificate of Conformance provides information on conformance testing of each instrument provided with the system. Certificates of Conformance are shipped with the instrument, and are also available by contacting Customer Support at [thermofisher.com/support](https://www.thermofisher.com/support).

Oncomine™ Dx Target Test Part III: Template Preparation

USER GUIDE

Publication Number MAN0018705

Revision A.0



For In Vitro Diagnostic Use.



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Block 33 |
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#07-06, Singapore 739256



Life Technologies Corporation |
7335 Executive Way |
Frederick, MD 21704 | USA

Products manufactured in Singapore:

Ion PGM™ Dx Instrument System
Ion PGM™ Dx Sequencer
Ion OneTouch™ Dx Instrument
Ion OneTouch™ ES Dx Instrument
Ion PGM™ Dx Chip Minifuge (120V)
Ion PGM™ Wireless Scanner
Ion Torrent™ Server
Torrent Suite™ Dx Software
Veriti™ Dx 96-well Thermal Cycler, 0.2 mL

Products manufactured in Frederick:

Oncomine™ Dx Target Test Kit
Ion Torrent Dx FFPE Sample Preparation Kit
Ion PGM™ Dx Library Kit
Ion OneTouch™ Dx Template Kit
Ion PGM™ Dx Sequencing Kit
Ion 318™ Dx Chip
Ion OneTouch™ Rack Kit
DynaMag™ Dx 96-Well Plate Magnet
DynaMag™ Dx 16 2-mL Magnet

The information in this guide is subject to change without notice.

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Revision history: Pub. No. MAN0018705

Revision	Date	Description
B.0	3 September 2021	Updated with new product catalog numbers.
A.0	11 February 2021	New Oncomine™ Dx Target Test user guide for FDA submission.

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About this guide



CAUTION! ABBREVIATED SAFETY ALERTS. Hazard symbols and hazard types specified in procedures may be abbreviated in this document. For the complete safety information, see the “Safety” appendix in this document.

IMPORTANT! Before using this product, read and understand the information in the “Safety” appendix in this document.

Purpose of this guide

This user guide provides instructions for using the Ion OneTouch™ Dx System to prepare enriched, template-positive Ion PGM™ Dx Ion Sphere™ Particles (ISPs) from sample and control libraries prepared using the Oncomine™ Dx Target Test Kit. The Ion OneTouch™ Dx System includes the Ion OneTouch™ Dx Instrument and the Ion OneTouch™ ES Dx Instrument.

This user guide is organized as follows:

- Prepare template-positive ISPs from Oncomine™ Dx Target Test libraries using the Ion OneTouch™ Dx Template Kit with the Ion OneTouch™ Dx Instrument.
- Enrich the template-positive ISPs with the Ion OneTouch™ ES Dx Instrument.

Oncomine™ Dx Target Test Kit user guides

This user guide is part of a five-guide set.

- *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*
- *Oncomine™ Dx Target Test Part II: Sample and Library Preparation User Guide*
- *Oncomine™ Dx Target Test Part III: Template Preparation User Guide*
- *Oncomine™ Dx Target Test Part IV: Sequencing and Results Reports User Guide*
- *Oncomine™ Dx Target Test Part V: Torrent Suite™ Dx Software 5.12.5 Reference User Guide*

All five guides are required to complete the entire Oncomine™ Dx Target Test workflow.

Note: The procedures in these guides supersede the instructions in the *Ion PGM™ Dx System User Guide* when using the Ion PGM™ Dx System with the Oncomine™ Dx Target Test.



Product information

Product description

Oncomine™ Dx Target Test

The Oncomine™ Dx Target Test is an *in vitro* diagnostic next-generation sequencing test to detect somatic alterations in human DNA and RNA isolated from non-small cell lung cancer (NSCLC) and cholangiocarcinoma (CC) tumor specimens in formalin-fixed, paraffin-embedded (FFPE) tissue samples. Detection of these variants is performed using the Ion PGM™ Dx System.

For a complete product description of the Oncomine™ Dx Target Test, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Template preparation components

The Ion OneTouch™ Dx Template Kit, included as part of the Oncomine™ Dx Target Test Kit, is used in conjunction with the Ion OneTouch™ Dx System to prepare template-positive Ion PGM™ Dx ISPs for sequencing on the Ion PGM™ Dx Sequencer.

The Ion OneTouch™ Dx System includes the following components:

- Ion OneTouch™ Dx Instrument and accessories
- Ion OneTouch™ ES Dx Instrument and accessories
- Ion OneTouch™ Solutions Rack
- Ion OneTouch™ Assembly Rack
- Ion OneTouch™ Sample Rack

Intended use

For the Intended Use statement for the Oncomine™ Dx Target Test, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Theory of operation

For a complete description of the Theory of Operation of the system, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Software compatibility and requirements

The procedures in this guide are designed for use with Torrent Suite™ Dx Software version 5.12.5 or later. For a complete description of software compatibility and requirements, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Materials provided

Oncomine™ Dx Target Test Kit

The Oncomine™ Dx Target Test Kit (Cat. No. A51695) includes the following subkits.

IMPORTANT! Refer to the product label for the storage conditions and expiration dates of individual modules and components.

✓	Subkit	Part No.
	Oncomine™ Dx Target Test, Controls, and Diluent Kit	A49756
	Ion Torrent Dx FFPE Sample Preparation Kit	A32445
	Ion PGM™ Dx Library Kit	A49758
	Ion OneTouch™ Dx Template Kit	A49759
	Ion PGM™ Dx Sequencing Kit	A49760
	Ion 318™ Dx Chip Kit	A18937
	Oncomine™ Dx Target Test User Guides and Assay Definition File	A51694

Subkits used in this guide

The procedures in this user guide use the following subkits from the Oncomine™ Dx Target Test Kit.

Ion OneTouch™ Dx Template Kit

The Ion OneTouch™ Dx Template Kit (Cat. No. A49759) includes the following modules and components.

IMPORTANT! Refer to the product label for the expiration date of the kit.

✓	Component	Amount	Storage
Ion OneTouch™ Dx Template Supplies (Part No. A18933)			
	TMPL Amplification Plate	8	15°C to 30°C
	TMPL Recovery Router	8	
	TMPL Recovery Tube	16	
	TMPL Sipper	2	
	TMPL Reagent Tube	2	
	TMPL ES Tip	8	
	TMPL ES Strip Tube	1 pack of 12	
	TMPL Cleaning Adapter	8	
	TMPL Emulsion Cartridge	8	
	TMPL Reagent Tube Labels	1 set	
	TMPL Sample Collection Tube ^[1]	1 pack of 50	
Ion OneTouch™ Dx Template Solutions (Part No. A18932)			
	TMPL Oil (white cap)	450 mL	15°C to 30°C
	TMPL Reaction Oil (white cap)	22 mL	
	TMPL Water (yellow cap)	320 µL	
	TMPL Recovery Solution (brown cap)	280 mL	
	TMPL Wash Solution (white cap)	15.2 mL	
	TMPL Rgnt B (blue cap)	2 × 1.2 mL	
	TMPL ES Rsp Soln (orange cap)	1.04 mL	
	TMPL Neutral Soln (red cap)	80 µL	
	TMPL Tween™ Solution (white cap)	2.24 mL	

✓	Component	Amount	Storage
Ion OneTouch™ Dx Template Reagents (Part No. A18930)			
	TMPL Enzyme Mix (brown cap)	400 µL	–30°C to –10°C
	TMPL Rgnt Mix (violet cap)	8 × 500 µL (single-use tubes)	
	TMPL ISP (black cap)	800 µL	
	TMPL CF-1 (clear cap)	40 µL	
Ion OneTouch™ Dx Template ES Beads (Part No. A18931)			
	TMPL ES Beads (green cap)	104 µL	2°C to 8°C

[1] Extra tubes are provided. Use one tube per Ion OneTouch™ ES Dx Instrument run.

Ion PGM™ Dx Instrument System

The Ion PGM™ Dx Instrument System (Cat. No. A25511) includes the following components, which are also sold separately.

✓	Component	Catalog No.
	Ion OneTouch™ Dx Instrument and accessories	A25483
	Ion OneTouch™ ES Dx Instrument and accessories	A25484
	Ion PGM™ Dx Sequencer and accessories	A25485
	Ion PGM™ Wireless Scanner	A25486
	Ion Torrent™ Server (software installed separately)	A28552
	Ion OneTouch™ Rack Kit <ul style="list-style-type: none"> • Ion OneTouch™ Solutions Rack • Ion OneTouch™ Assembly Rack • Ion OneTouch™ Sample Rack 	A24694
	Ion PGM™ Dx Chip Minifuge: <ul style="list-style-type: none"> • 120 VAC • 230 VAC 	A25058 A25482
	DynaMag™ Dx Kit—Tube & Plate <ul style="list-style-type: none"> • DynaMag™ Dx 96-Well Plate Magnet • DynaMag™ Dx 16 2-mL Magnet 	A31755 A31347 A31346

Materials and equipment required but not provided

Unless otherwise indicated, all materials are available through thermofisher.com. "MLS" indicates that the material is available from fisherscientific.com or another major laboratory supplier.

✓	Description	Source
	Laminar flow hood	MLS
	1.5-mL snap-cap low-retention polypropylene microcentrifuge tubes	MLS
	Mini centrifuge	MLS
	Pipettes (2-, 20-, 200-, 1000- μ L)	MLS
	Aerosol-barrier pipette tips (10-, 20-, 200-, 1000- μ L)	MLS
	Vortex mixer with a rubber platform	MLS
	50-mL conical tubes	MLS
	Holder for 50-mL conical tube	MLS
	DynaMag™ Dx 16 2-mL Magnet	A31346
	NaOH, ACS grade (10 M)	MLS
	Nuclease-free Water	MLS
	Benchtop cold box	MLS

DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet

Note: Do not substitute non-IVD labeled magnets for the DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet.

The DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet, provided with Ion PGM™ Dx System, contain high-energy neodymium magnets and are used as part of the procedure for purifying sample libraries bound to LIB AMPure™ Reagent and LIB Beads. The DynaMag™ Dx 16 2-mL Magnet is also used to prepare TMPL ES Beads as part of template preparation.

The DynaMag™ Dx 96-Well Plate Magnet has 7 bar magnets with a hard plastic top to fit 96-well PCR plates. When you insert a plate, the magnets collect bead-bound biomolecules in suspension at the sides of the plate wells, allowing removal of fluid without disturbing the bead pellets. An extra column in the magnet enables sample mixing by shifting the plate back and forth in the magnet.

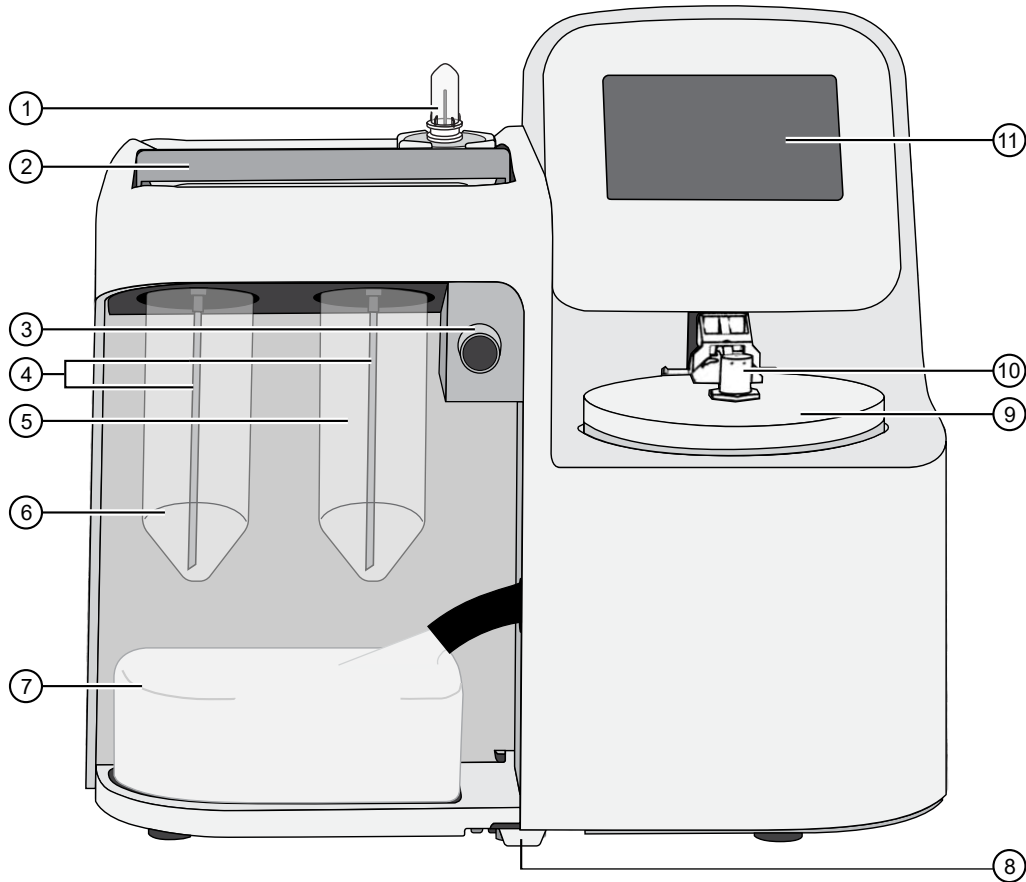
The DynaMag™ Dx 16 2-mL Magnet holds 16 standard 1.5-mL or 2-mL microcentrifuge tubes, and collects bead-bound biomolecules in suspension at the sides of the tubes, allowing removal of fluid without disturbing the bead pellets.

Do not use the magnets above 50°C (122°F) and store in a cool, dry environment.

Recommended materials and equipment

Description	Source
Benchtop absorbent paper or mat	MLS

Ion OneTouch™ Dx Instrument



- ① Reaction Filter
- ② Clamp handle to access the TMPL Amplification Plate in the heat block
- ③ Pinch valve to hold disposable tubing
- ④ TMPL Sippers
- ⑤ TMPL Reagent Tube containing TMPL Recovery Solution 💧
- ⑥ TMPL Reagent Tube containing TMPL Oil 🛢️
- ⑦ Waste Container
- ⑧ Oil waste tray
- ⑨ Centrifuge and TMPL Recovery Router
- ⑩ Injector hub
- ⑪ Touchscreen



Before you begin

Electrical and environmental conditions

IMPORTANT! Protection provided by the equipment may be impaired if the instrument is operated outside the environment and use specifications, the user provides inadequate maintenance, or the equipment is used in a manner not specified by the manufacturer.

IMPORTANT! Observe current Good Clinical Laboratory Practices (GCLP) when using this instrument.

See the *Ion PGM™ Dx System Site Preparation Guide* (Pub. No. MAN0024945) for information about instrument location and setup. Instruments should be shielded from excess exposure to dust, vibration, strong magnetic fields, drafts, excessive moisture, or large temperature fluctuations. Surge protectors or line conditioners should be used if the voltage source is not stable. Sudden voltage spikes can cause damage to the electronics inside the instruments.

Ensure that the room where the instruments have been installed is maintained under correct environmental conditions. Avoid placing the instruments adjacent to heaters, cooling ducts, or in direct sunlight. Place the instruments at least a meter away from major sources of electronic noise, such as refrigerators or microwaves.

Electrical requirements



CAUTION! Do not unpack or plug in any components until a field service representative has configured them for the proper operating voltage.



WARNING! For safety, the power outlet used for powering the instrument must be accessible at all times. In case of emergency, you must be able to immediately disconnect the main power supply to all the equipment. Allow adequate space between the wall and the equipment so that the power cords can be disconnected in case of emergency.

- Electric receptacle required: 2-prong with ground pin
- Main AC line voltage tolerances must be at most $\pm 10\%$ percent of nominal voltage.

- Power cords are provided with the instruments. If not suitable for installation in your region, ensure any power cord you do use is:
 - Maximum 10 feet (3 meters) in length
 - Grounding type
 - Compatible with the power supply receptacles used to connect to main power
 - Suitable for the rating of the instrument and main power supply
 - Compliant with local safety requirements (for example, UL Listed for North America, JIS approved for Japan, HAR or agency certified for Europe)
- (Ion OneTouch™ Dx Instrument only) Fuse Rating: 6 A, 250 VAC, Type M. Replace only with the same fuse type and rating.



WARNING! FIRE HAZARD. For continued protection against the risk of fire, replace fuses only with fuses of the type and rating specified for the instrument.

Device	Rated voltage ^[1,2]	Rated frequency	Rated current ^[3]
Ion PGM™ Dx Sequencer	110/120VAC 220/240VAC	50/60 Hz	9 A
Ion Torrent™ Server ^[4]	110/120VAC 220/240VAC	50/60 Hz	11 A
Ion OneTouch™ Dx Instrument with power supply	110/120VAC 220/240VAC	50/60 Hz	5.5 A
Ion OneTouch™ ES Dx Instrument	110/120VAC 220/240VAC	50/60 Hz	375 mA 160 mA
Ion PGM™ Dx Chip Minifuge	120 VAC 220–240 VAC	50/60 Hz	130 mA 65 mA

^[1] In Japan, rated voltages of 100 VAC and 200 VAC are acceptable.

^[2] If the supplied power fluctuates beyond the rated voltage, a power line regulator may be required. High or low voltages can adversely affect the electronic components of the instrument.

^[3] Based on rated current at minimum input voltage.

^[4] Minimum Efficiency: 65% (Energy Star Qualified); 85% Efficient Power Supply.

Environmental requirements

Ensure that the room where the instruments have been installed is maintained under the correct environmental conditions. Avoid placing the instruments next to heaters, cooling ducts, or in direct sunlight. Place the sequencer at least a meter away from major sources of electronic noise, such as refrigerators or microwaves.



CAUTION! Use of the instruments in an unspecified manner may result in the protection provided by the instruments to be impaired.

Component	Acceptable range
Altitude	Between sea level and 2,000 meters (6,500 feet) above sea level
Humidity: operating	10–90% relative humidity
Humidity: transportation and storage	20–80% relative humidity
Temperature: operating	15–30°C (59–86°F) At or above 1,800 meters (5,906 feet), the system must not be used if the temperature is above 29.5°C.
Temperature: transportation and storage	–30°C to 60°C (–22°F to 140°F)
Vibration	Ensure that benches where instruments are to be installed are free of vibration and have no contact with equipment that causes vibration (freezers, pumps, and similar equipment). Vibration can reduce the quality of sequencing measurements.
Pollution	The system has a Pollution Degree rating of II (2). The system may only be installed in an environment that has nonconductive pollutants, such as dust particles or wood chips. Typical environments with a Pollution Degree II (2) rating are laboratories, sales, and commercial areas.
Overvoltage category	The instruments have an installation (overvoltage) category of II (2).
Other conditions	For indoor use only. Keep away from any vents that could expel particulate material on the system components.

Precautions before using the Ion OneTouch™ Dx instruments

For additional safety information, see Appendix C, “Safety”.

Instrument installation by trained personnel only

IMPORTANT! The Ion PGM™ Dx System is installed by trained service personnel and must not be relocated without assistance from trained service personnel. See Appendix F, “Customer and technical support”.

Nucleic acid contamination

IMPORTANT! A primary source of contamination is DNA fragments from previously processed samples. See the notes about avoiding contamination in the procedural guidelines.

Reagent contamination

Before use, verify that the TMPL Water and TMPL Tween™ Solution are not cloudy, a potential indication of contamination. If these reagents are cloudy, use a different vial.

Procedural guidelines

Definitions

Throughout this guide:

- Room temperature is defined as the temperature range 15–30°C (59–86°F).
- A pulse centrifugation consists of a 3–5 second centrifugation at maximum speed in a mini centrifuge.

Guidelines for pipetting

Pipet viscous solutions slowly and ensure complete mixing.

Guidelines to prevent cross-contamination

- Use good laboratory practices to minimize cross-contamination of products and reagents.
- When designing the laboratory layout, dedicate separate areas for pre- and post-amplification activities. Dedicate laboratory supplies and/or equipment to the appropriate area.
- Before and after use, clean all working surfaces with fresh 10% bleach followed by two water rinses.
- Use a laminar flow hood in the dedicated pre-amplification area when combining libraries and preparing the amplification solution.

- Before and after use, clean all surfaces and equipment in the laminar flow hood with fresh 10% bleach followed by two water rinses.
- Turn on the UV light in the hood for 10 minutes before and after use.
- Use fresh gloves before entering the hood.
- Change tips between pipetting steps.
- Prepare a waste container containing fresh 10% bleach solution for disposing of used tips after pipetting libraries.
- To collect the contents before opening, pulse centrifuge tubes containing libraries and library pools.
- When simultaneously preparing more than one amplification solution, only open one library-pool tube at a time.
- Use fresh reagents if a contamination event occurs or is suspected.
- Use fresh gloves when installing new consumables.
- To avoid contamination of TMPL Reagent Tubes (which may be reused), discard waste from the tubes in a separate container from other Ion OneTouch™ Dx Instrument waste and 50-mL conical-tube waste.
- Always change gloves after handling Ion OneTouch™ Dx Instrument waste oil, used amplification plates, and used cleaning adapters.
- To prevent cross-contamination, we do not recommend running the same barcode for the same type of library sample (DNA or RNA) in a consecutive system run.

Guidelines for Ion OneTouch™ Dx Instrument operation

- To prevent debris from entering the system, keep the TMPL Reagent Tubes installed on the Ion OneTouch™ Dx Instrument when not in use.
- After a Planned Run is executed on the Ion Torrent™ Server, the run must be started immediately on the Ion OneTouch™ Dx Instrument.
- A run on the Ion OneTouch™ Dx Instrument can be performed overnight. Enrichment on the Ion OneTouch™ ES Dx Instrument must start within 24 hours after completion of the Ion OneTouch™ Dx Instrument run.
- If a run is aborted for any reason, you must restart the Ion OneTouch™ Dx Instrument by power cycling.
- Ensure that you clean up any spilled oil immediately. Place a nonslip floor mat in front of the instrument to prevent slips.



CAUTION! Spilled oil from the Ion OneTouch™ Dx Instrument can present a slip hazard.

Reagent management

Follow the guidelines below for proper reagent storage and use.

Storage

Reagents must be stored under appropriate conditions. Refer to the Product Information section in each user guide for the storage conditions of the kit components used in the procedures in that guide. The Oncomine™ Dx Target Test Kit includes kits with multiple component boxes that require different storage conditions. For example, the Oncomine™ Dx Target Test, Controls, and Diluent Kit includes four boxes, which are stored at different temperatures. To use the Oncomine™ Dx Target Test, Controls, and Diluent Kit, retrieve all boxes from their different storage areas and confirm that they are from the same master lot.

Kit interchangeability and component box lot matching

The top-level kits used for sample preparation, library preparation, template preparation, and sequencing can be mixed and matched. For example, an Ion PGM™ Dx Library Kit can be used with any Ion PGM™ Dx Sequencing Kit. However, the component boxes in a particular kit must be lot-matched with the other boxes in that kit.

Note: The controls and diluent in the Oncomine™ Dx Target Test, Controls, and Diluent Kit must be lot-matched. However, the controls do not require lot-matching with the Oncomine™ Dx Target Test DNA and RNA Panel.

Each component box lists the 4-digit lot prefixes of the compatible component boxes inside the box lid. Before using a particular kit, check the inside lid of each box to confirm that it is compatible with the other boxes.

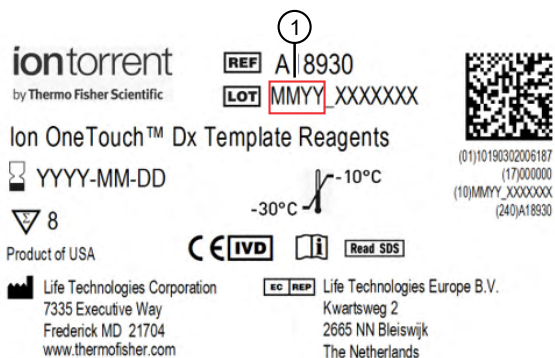
An example inside box lid label is shown below:

ATTENTION: When setting up a procedure, confirm that the component boxes listed below have 4-digit lot prefixes that match.

REF		LOT
A18930	Ion OneTouch™ Dx Template Reagents	MMYY
A18931	Ion OneTouch™ Dx Template ES Beads	MMYY
A18932	Ion OneTouch™ Dx Template Solutions	MMYY
A18933	Ion OneTouch™ Dx Template Supplies	MMYY

①

An example box label with lot information is shown below:

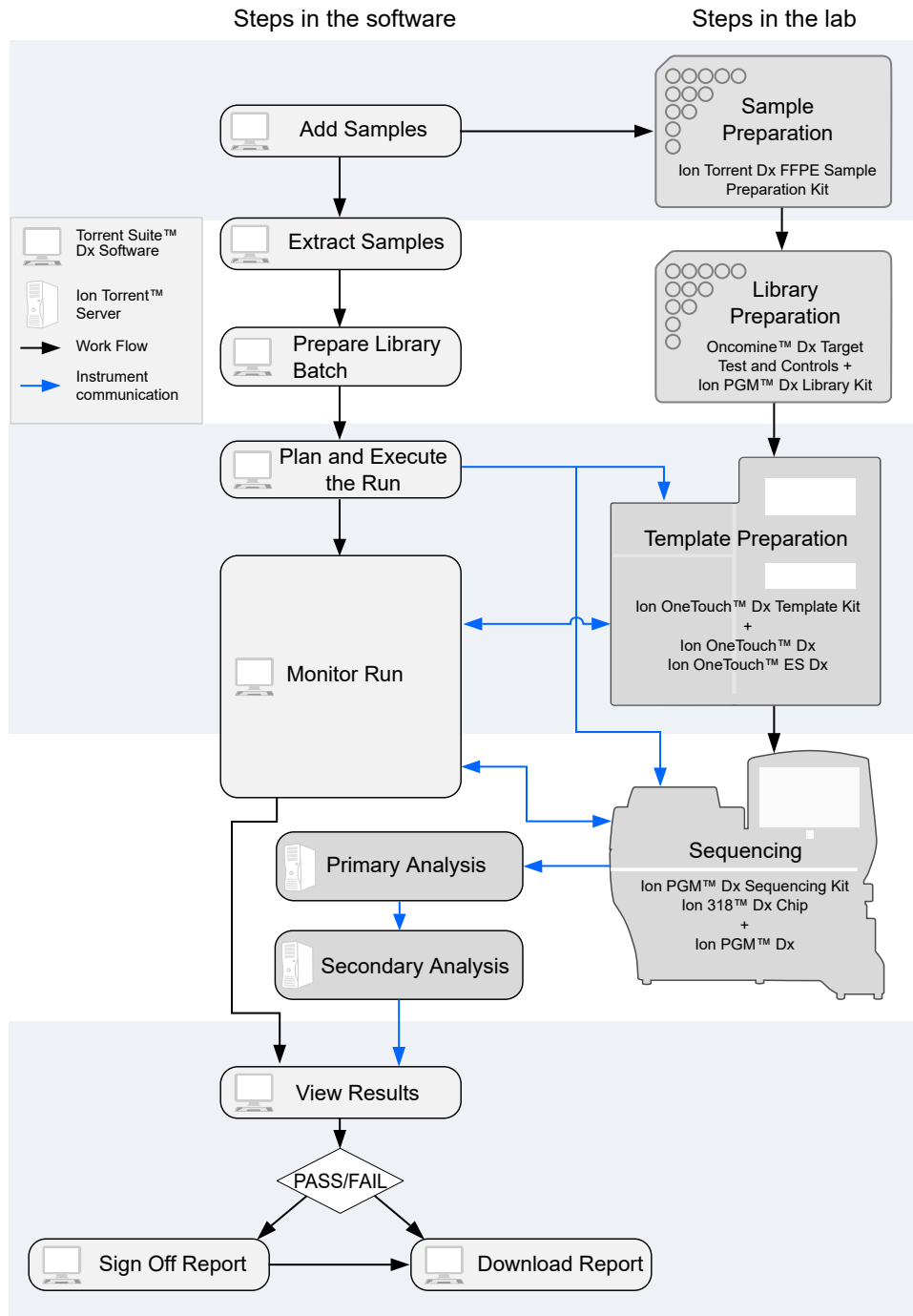


① Lot prefix

Library preparation

Libraries must be prepared as described in the *Oncomine™ Dx Target Test Part II: Sample and Library Preparation User Guide*.

Oncomine™ Dx Target Test system diagram



Template preparation workflow

The total workflow time is 7 hours, including 1 hour of hands-on time.

Previous guide: *Oncomine™ Dx Target Test Part II: Sample and Library Preparation User Guide*



Plan and execute a run on the Torrent Suite™ Dx Software (page 23)



Pool sample and control libraries (page 27)



Clean the Ion OneTouch™ Dx Instrument before a run (page 30)



Run the Ion OneTouch™ Dx Instrument (page 43)



Prepare the Ion OneTouch™ ES Dx Instrument and perform the run (page 49)



Clean the Ion OneTouch™ Dx Instrument after the run (page 51)



Proceed to: *Oncomine™ Dx Target Test Part IV: Sequencing and Results Reports User Guide*



Plan and execute a run on the Torrent Suite™ Dx Software


Components such as libraries, kits, and chips used in a diagnostic assay must be uniquely identified, and the identification must be stored so that the record can be audited. Torrent Suite™ Dx Software records these components in the Planned Run, which is prepared in the software and then transferred to the system instruments for tracking and verification at each stage of the run.

The software also uses the Planned Run to verify that only unexpired kits and chips are used and the correct assay is performed on the correct sample.

For more information, see the *Oncomine™ Dx Target Test Part V: Torrent Suite™ Dx Software 5.12.5 Reference User Guide*.

Create a Planned Run

Libraries that are ready to be entered into a Planned Run are listed under the **Samples** tab in the **Libraries** screen.

Note: You can also plan a run from the **Runs** tab in the **Planned Runs** screen (click  **Add New**).

1. Sign in to Torrent Suite™ Dx Software.
2. In the **Libraries** screen, select the library or libraries to be run by selecting checkboxes in the list. To view only those libraries that have not yet been added to a Planned Run, select **To Be Planned** in the **Filter Libraries by** dropdown list.

Note:

- Libraries prepared with the same assay in the same library batch can be combined and run together, as long as they have unique library names and Barcode IDs.
 - To plan a run with the Oncomine™ Dx Target Test assay, a report template must be created and associated with the assay by an Administrator or Manager. Under the **Assay** tab, go to **Presets ▶ Report Templates** to add a new report template.
 - Up to 16 libraries (including controls) can be sequenced in a single run.
 - If libraries need to be rerun due to a run failure, they can be added to a new Planned Run.
-

3. Click **+** **Plan a Run**.

Manage Samples Import Samples **Libraries** Import Library Batch Manage Attributes

To Be Planned Library Name Enter Library Name **Plan a Run** Delete

Selected Libraries: 1

<input checked="" type="checkbox"/>	Library Batch ID	Assay Name	Sample ID	Library Name	Library Type	Barcode ID	Notes	Actions
<input checked="" type="checkbox"/>	21081007_BC1	OncoPrint™ Dx Target Test v2.1.1	BC1	BC1_DNA BC1_RNA	DNA RNA	IonDx-2 IonDx-10		Edit Audit

1 - 1 of 1 Items

4. In the **Add New Plan** dialog box, enter a name for the run, then select the appropriate report template.

The selected library or libraries are listed in the dialog box, and the control libraries are automatically listed.

Add New Plan X

Name: * Warr_1

Assay Name: OncoPrint™ Dx Target Test v2.1.1

Select Report Template: Report_AutoInserted_Oncomi

Notes:

Number of Sample Libraries: 2

<input type="checkbox"/>	Library Batch ID	Sample ID	Library Name	Barcode ID	Library Type
<input type="checkbox"/>	21081007_ BC1	BC1	BC1_DNA BC1_RNA	IonDx-2 IonDx-10	DNA RNA
<input type="checkbox"/>		NA	internalControl_14	IonDx-9	RNA Control

Remove **+ Add more Libraries**

Cancel Save

5. To remove libraries from the run, select the appropriate checkboxes, then click **Remove**. To add libraries, click **+ Add more Libraries**, then select them from the **Add Libraries** dialog box.

Any added libraries must be from the same library batch and have unique library names and Barcode IDs.

6. Click **Save**.

The new Planned Run is automatically assigned a Run Short Code and is displayed at the top of the list under the **Runs** tab in the **Manage Runs** screen.

Execute a Planned Run

Planned Runs are listed under the **Runs** tab in the **Manage Runs** screen. Runs that are ready to be performed have the **Execute** command available in the **Actions** column.

Executing a Planned Run in the software queues the run for initiation on the Ion OneTouch™ Dx Instrument. After a Planned Run has been queued for execution, the operator should immediately start template preparation.

In the **Manage Runs** screen:

1. Click **To Be Started** in the **Filter Run by...** dropdown list to limit the list of Planned Runs to only those runs yet to be started.
2. Find the Planned Run in the list, then under the **Actions** header, click **Execute**.



The **Execute Planned Run** dialog box opens.

3. In the **Tube Label** field, enter the text that is used to label the tubes that contain the final combined libraries.

The tube label text can be any combination of letters and numbers. The system uses this text to track the sample throughout the run, so be careful to label each tube legibly at the points noted in the procedure. The software does not allow use of the same Tube Label text within 7 days.

4. Click the **Template Prep Kit barcode** field, then scan the barcode from the Ion OneTouch™ Dx Template Reagents box.

IMPORTANT! Ensure that you scan the barcode from the actual reagents box that is used in the run.



5. Click **Save** to save your changes.
The **Review Planned Run** dialog box opens.
6. Write down the **Run Short Code** and/or click **Print** to print the scannable barcode. The code must be entered into the Ion OneTouch™ Dx Instrument and Ion PGM™ Dx Sequencer for tracking and verification before the start of the instrument run.
7. Click **Close** to close the dialog box and send the run to the instrument.

Note: The last 5 executed Planned Runs are listed under the **Monitor** tab in the software.

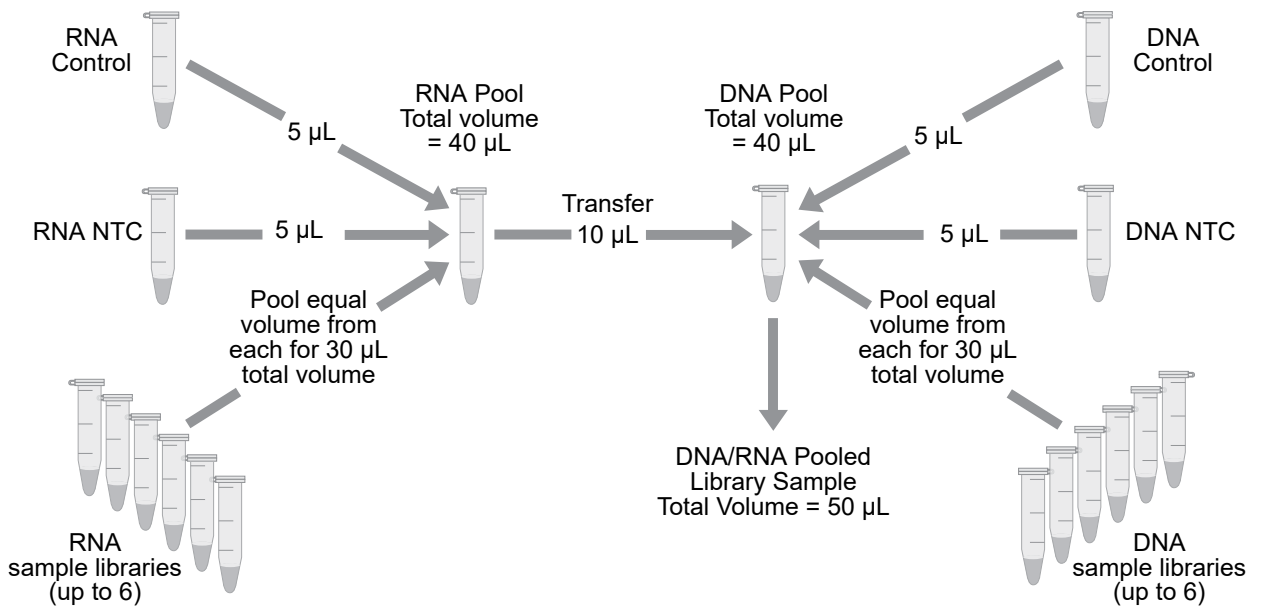
4

Pool sample and control libraries

Strategy for combining libraries

You can multiplex up to six RNA sample libraries and six DNA sample libraries into a single Ion PGM™ Dx System run. You must also include separate DNA Control, RNA Control, and No Template Control libraries with each run.

A strategy for combining multiple libraries and controls is diagrammed below.



Create an RNA combined library

Perform the following steps in a laminar flow hood in a designated post-PCR area. Change pipette tips between libraries. See “Guidelines to prevent cross-contamination” on page 17 for additional guidelines to avoid cross-contamination.

1. Before pipetting, vortex each library tube for ~5 seconds, then pulse centrifuge for 3–5 seconds to collect the contents.
2. Label a new, nuclease-free 1.5-mL low-retention microcentrifuge tube with the text "RNA Combined Library" and the date.
3. Add 5 µL of the RNA Control library to the tube.
4. Add 5 µL of the RNA NTC library to the same tube.

- Based on the number of RNA sample libraries to combine, transfer the volume shown in the following table to the tube.

Note: The total volume of the combined libraries and controls equals 40 μL .

# of RNA sample libraries	Volume per RNA sample library
1	30 μL
2	15 μL
3	10 μL
4	7.5 μL
5	6 μL
6	5 μL

- Vortex the combined library for ~5 seconds, then pulse centrifuge for 3–5 seconds to collect the contents.

Create a DNA combined library

Perform the following steps in a laminar flow hood. Change pipette tips between libraries. See “Guidelines to prevent cross-contamination” on page 17 for additional guidelines to avoid cross-contamination.

- Before pipetting, vortex each library tube for ~5 seconds, then pulse centrifuge for 3–5 seconds to collect the contents.
- Label a new, nuclease-free 1.5-mL low-retention microcentrifuge tube with the text "DNA/RNA Combined Library" and the date.
- Add 5 μL of the DNA Control library to the tube.
- Add 5 μL of the DNA NTC library to the tube.
- Based on the number of DNA sample libraries to combine, add the volume shown in the following table to the tube.

Note: The total volume of the combined libraries and controls equals 40 μL .

# of DNA sample libraries	Volume per DNA sample library
1	30 μL
2	15 μL
3	10 μL
4	7.5 μL

(continued)

# of DNA sample libraries	Volume per DNA sample library
5	6 µL
6	5 µL

- Vortex the combined library for ~5 seconds, then pulse centrifuge for 3–5 seconds to collect the contents.

Create a DNA/RNA combined library

- Transfer 10 µL of the RNA combined library (from step 6, “Create an RNA combined library” on page 27) to the tube containing 40 µL of DNA combined library (labeled “DNA/RNA Combined Library”).

Note:

- The total volume of the DNA/RNA combined library equals 50 µL.
 - The remaining RNA combined library can be stored at –30°C to –10°C for up to 30 days.
-

- Vortex the DNA/RNA combined library for ~5 seconds, then pulse centrifuge for 3–5 seconds to collect the contents.

STOPPING POINT Proceed to “Clean the Ion OneTouch™ Dx Instrument before a run” on page 30, or store the DNA/RNA combined library at –30°C to –10°C for up to 30 days.



Clean the Ion OneTouch™ Dx Instrument before a run

Track use of TMPL Reagent Tubes and TMPL Sippers

TMPL Reagent Tubes and TMPL Sippers can be used up to 8 times.

1. To track use of the reagent tubes and sippers, label the tubes with the labels that are provided in the kit, then mark the labels after each use.
2. After 8 uses, discard the used reagent tubes and sippers in an appropriate waste container, then label new reagent tubes.

Power cycle the Ion OneTouch™ Dx Instrument

The Ion OneTouch™ Dx Instrument can be left on overnight and on weekends, but should be power cycled under the following conditions:

To power cycle the instrument, turn the instrument off, wait 3 seconds, then turn the instrument back on.

- Power cycle the instrument before installing TMPL Reagent Tubes from a new kit.
- Power cycle the instrument after daylight-saving time changes.

IMPORTANT! Allow up to 20 minutes for the Ion OneTouch™ Dx Instrument to resynchronize with the Ion Torrent™ Server after power cycling. Failure to resynchronize generates an alarm until synchronization is complete.

Clean the Ion OneTouch™ Dx Instrument before a run

Before you perform a new run on the Ion OneTouch™ Dx Instrument, you must clean the instrument. Until you clean the instrument, the **Run** button on the instrument remains disabled.

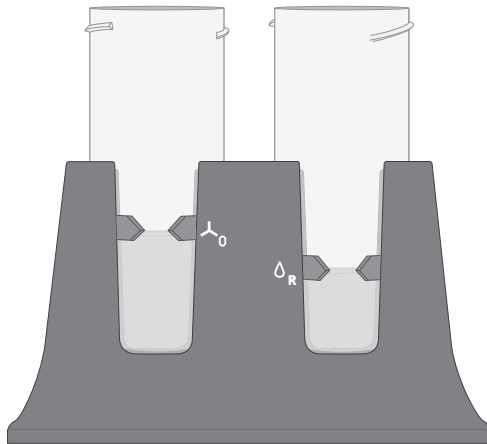
IMPORTANT! Change gloves after handling instrument waste during disposal and before handling samples. See the product SDS for guidance regarding proper disposal and handling of Ion OneTouch™ Dx Instrument waste products.

Note: Do not press the instrument touchscreen buttons too rapidly (>1 per second), or the touchscreen may freeze.

1. Log in to the instrument using the touchscreen.
2. Press the **Clean** button, then follow the touchscreen prompts.
3. Remove the TMPL Reagent Tube containing TMPL Oil from the position marked with an "O" on the instrument (the tube on the left when facing the instrument), then discard the contents in an appropriate hazardous waste container.

IMPORTANT! When removing reagent tubes, do not touch the reagent sippers or allow them to come into contact with any surfaces. Ensure that the reagent tubes do not come into contact with the waste container when emptying contents.

4. Place the TMPL Reagent Tube in the position labeled "O" on the Ion OneTouch™ Solutions Rack.



5. Change gloves, then invert the TMPL Oil bottle (white cap) 5 times to mix. Fill the reagent tube with oil to the level marked on the rack, so the meniscus is within the fill indicator arrows (volume = 50 mL).
6. Confirm that the sipper is securely attached to the instrument, then screw the reagent tube containing the oil back into the position marked with an "O" on the instrument until the tube can no longer rotate. Press **Next**.



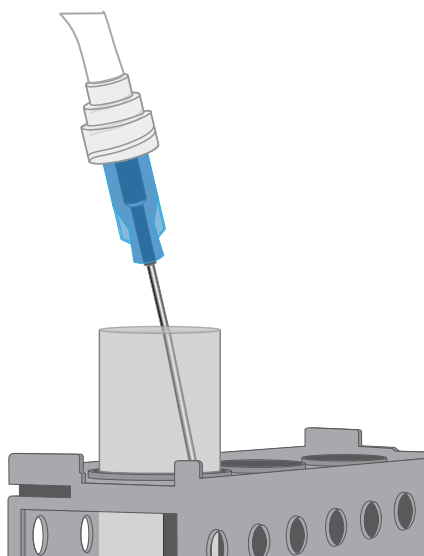
CAUTION! Be careful to attach the reagent tube to the correct position on the instrument. Attaching the tube to the wrong position may damage the instrument.

7. Remove the TMPL Reagent Tube containing TMPL Recovery Solution from the position marked with an "R" on the instrument. Discard the contents in an appropriate waste container.

IMPORTANT! When removing reagent tubes, do not touch the reagent sippers or allow the reagent sippers or reagent tubes to come into contact with any surfaces.

8. Place the TMPL Reagent Tube in the position marked with an "R" on the Ion OneTouch™ Solutions Rack.

9. Change gloves, then invert the TMPL Recovery Solution bottle (brown cap) 5 times to mix. Fill the reagent tube with recovery solution to the level marked with arrows on the "R" position of the rack (volume = 35 mL).
10. Confirm that the sipper is securely attached to the instrument, then screw the reagent tube containing the recovery solution back into the position marked with an "R" on the instrument until the tube can no longer rotate. Press **Next**.
11. Confirm that both the TMPL Cleaning Adapter and TMPL Amplification Plate from the post-run cleaning are in place on the instrument, then check that the tubing from the amplification plate is removed from the pinch valve.
12. Confirm that the injector tip is placed in an empty 50-mL conical tube to collect waste.



13. Press **Next** to start cleaning.

Note: The cleaning procedure takes 13 minutes to complete, and consumes 10 mL of oil.

14. When cleaning is complete, the screen displays "Cleaning Complete". Wearing clean gloves, press **Next**.
15. Remove and appropriately discard the used TMPL Cleaning Adapter, TMPL Amplification Plate, disposable injector, and tubing from the instrument.



CAUTION! Hot Surface. Use care when working near this area to avoid being burned by hot components.

- a. Remove the used TMPL Cleaning Adapter, then discard in an appropriate waste container.
- b. Lift, then push back the instrument handle to open the heat block.

- c. Remove the disposable tubing, remove the needle, then discard each in appropriate waste containers.



CAUTION! Piercing Hazard: Remove the needle from the tubing by unscrewing it or cutting it off with scissors, and discard in an appropriate hazardous waste container for sharp objects.

- d. Gently pull back the TMPL Amplification Plate from the inlet and outlet holes of the instrument.
 - e. Remove the plate from the heat block, then discard in an appropriate waste container.
 - f. Leave the heat block open.
16. Appropriately discard the 50-mL conical tube waste, then press **Next** to return to the main screen.

IMPORTANT! Always change gloves after handling the waste oil, used amplification plates, and cleaning adapters.

6

Perform an Ion OneTouch™ Dx System run

This chapter describes how to prepare templated ISPs on the Ion OneTouch™ Dx Instrument from a DNA/RNA combined library, and then enrich the templated ISPs on the Ion OneTouch™ ES Dx Instrument.

Prepare reagents and library tube for template preparation

1. Label the library tube with the Tube Label text that you entered into the software.
2. Equilibrate the library, TMPL Rgnt Mix, TMPL ISP, and TMPL CF-1 to room temperature for 30 minutes.

Ion OneTouch™ Dx Template Kit component lot matching

The four components of the Ion OneTouch™ Dx Template Kit must be lot-matched with each other for use.

Component	Part No.	Storage
Ion OneTouch™ Dx Template Supplies	A18933	15°C to 30°C
Ion OneTouch™ Dx Template Solutions	A18932	
Ion OneTouch™ Dx Template Reagents	A18930	-30°C to -10°C
Ion OneTouch™ Dx Template ES Beads	A18931	2°C to 8°C

Select the Planned Run

1. Change gloves, then press the **Run** button on the Ion OneTouch™ Dx Instrument touchscreen.
2. Enter the Run Short Code (generated by the Torrent Suite™ Dx Software when you executed the run). Make sure the Planned Run information on the screen is correct, then press **Next**.

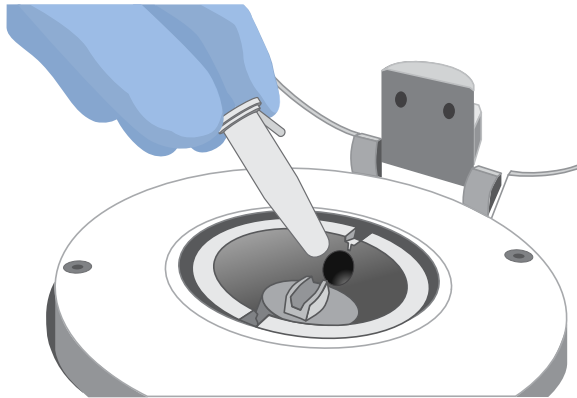
3. Enter the Tube Label text that you entered into the software when you executed the run, then press **Next**.

Note: For tracking purposes, the Tube Label text must exactly match the text you entered in the **Tube Label** field in the Torrent Suite™ Dx Software.

4. Confirm the run type displayed on the instrument screen, then press **Next**.

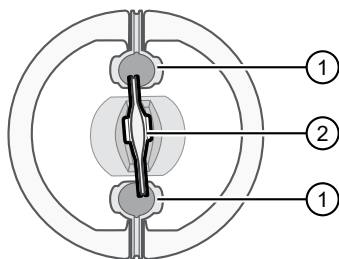
Install the TMPL Recovery Tubes, TMPL Recovery Router, and TMPL Amplification Plate

1. Label two TMPL Recovery Tubes with the Tube Label text entered into the software.
2. Insert the tubes into the holes in the Ion OneTouch™ Dx Instrument centrifuge, making the sure the tube arm is inserted into the slot next to each hole.



3. Pinch the sides of the TMPL Recovery Router, then push the router down into the center slot of the centrifuge until it is seated flat and secure in the center of the rotor.

Note: The router is not directly aligned with the tubes, but is intentionally positioned at an angle.

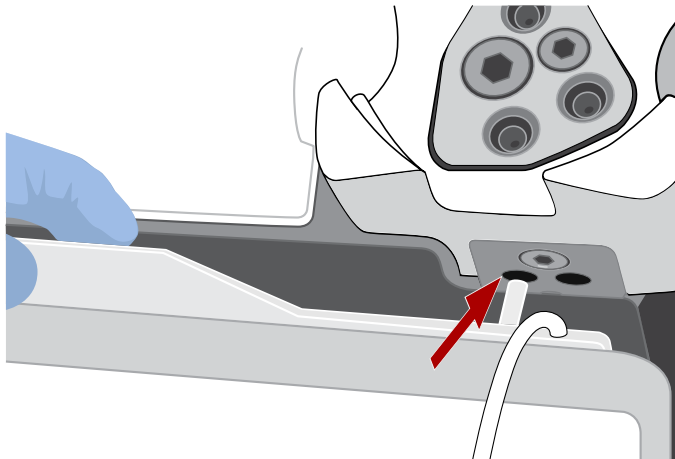


- ① TMPL Recovery Tube
- ② TMPL Recovery Router

4. Close the lid of the centrifuge, then press **Next**.

5. Insert a new TMPL Amplification Plate into the open instrument heat block.

Note: Ensure that the plate port is intact, then carefully align the plate port with the left outlet hole on the instrument. The plate includes disposable tubing and a disposable injector.



CAUTION! Hot Surface. Use care when working near the heat block to avoid burns from the hot components.

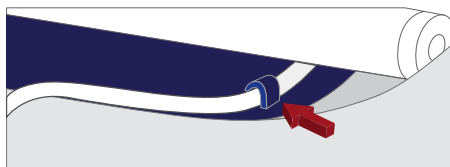


CAUTION! PHYSICAL INJURY HAZARD. The pointed end of the disposable injector can puncture your skin. Keep your hand away from the point of the injector.

6. Pull the handle forward to secure the plate. The tubing should be under the handle. Press **Next**.

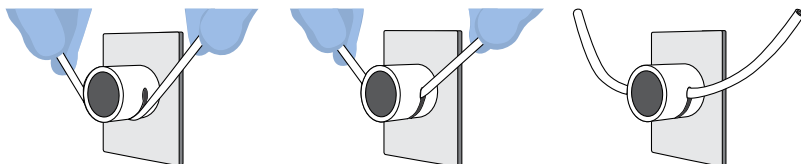
Note: In the following steps, ensure that the tubing is not kinked or twisted at any point along its length.

7. Thread the tubing through the tubing holder.



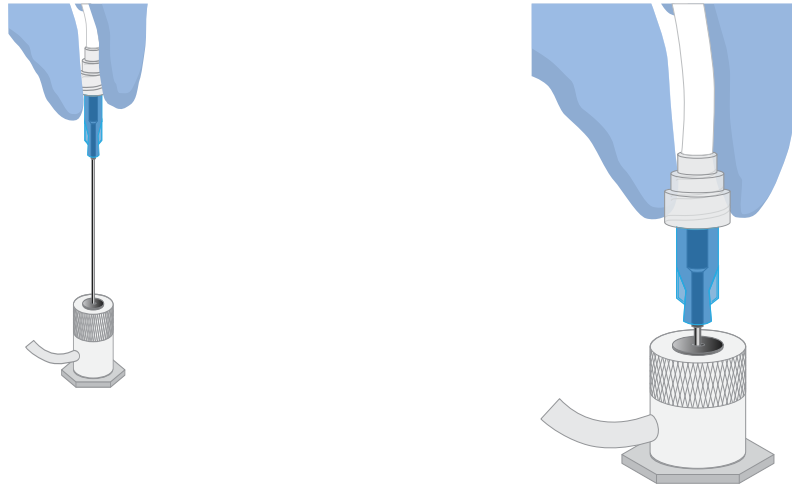
Note: Examine the tubing to ensure there are no kinks in the tubing before proceeding.

8. Align the tubing with the slot that runs along the bottom of the pinch valve. Gently pull the tubing up into the slot until it is secure in the notch.



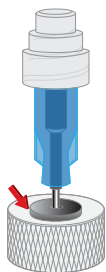
- Adjust the tubing so that it is straight but not too taut. The injector needle should reach the injector hub without stretching the tubing. Press **Next**.
- Ensure that the injector needle is screwed tightly onto the tubing. Hold the centrifuge lid down with one hand, and with your other hand insert the injector needle straight down into the injector hub. Push down until the injector touches the hub.

Note: The spring-loaded top of the injector hub clicks upon release, which indicates that the tip of the needle is the correct distance from the hub surface.

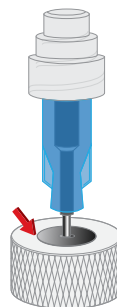


- Gently push the injector down again and release. You should hear a click from the hub. Then press **Next**.

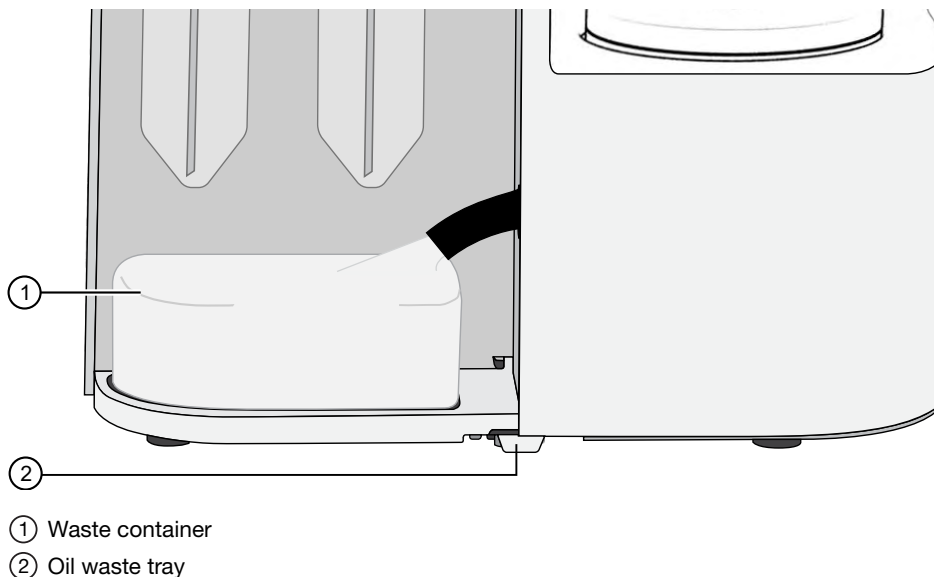
Up position



Down position



12. Pull the waste container from the external waste tubing, and empty the container into an appropriate hazardous waste receptacle.



13. Inspect the oil waste tray on the instrument.



CAUTION! If the oil waste tray is full, do not run the instrument. The instrument must be inspected and repaired by a field application specialist.

14. Put the empty waste container back on the tray, then push the waste tube back into the container port. Press **Next**, then proceed to prepare the amplification solution.

Prepare the amplification solution

Kit components used in this procedure

Kit component	Box
TMPL Water (yellow cap) TMPL Rgnt B (blue cap)	Ion OneTouch™ Dx Template Solutions (Part No. A18932, stored at 15°C to 30°C)
TMPL Rgnt Mix (violet cap) TMPL Enzyme Mix (brown cap) TMPL ISP (black cap) TMPL CF-1 (clear cap)	Ion OneTouch™ Dx Template Reagents (Part No. A18930, stored at -30°C to -10°C)

Prepare the amplification solution

Perform the following steps in a laminar flow hood in a designated preamplification area. See “Guidelines to prevent cross-contamination” on page 17 for additional guidelines to avoid cross-contamination.

IMPORTANT! The volume of TMPL ISP reagent used in the amplification solution is critical and must be accurate.

1. Put on new gloves after emptying the Ion OneTouch™ Dx Instrument waste.
2. Ensure that the library, TMPL Rgnt Mix, TMPL ISP, and TMPL CF-1 are completely thawed.
3. Vortex the TMPL Rgnt B and TMPL Rgnt Mix tubes for 5 seconds each, then pulse centrifuge for 3–5 seconds.
4. Flick the TMPL Enzyme Mix tube with the tip of your finger 4 times, then pulse centrifuge for 3–5 seconds. Place in a benchtop cold box at 2–8°C until needed.
5. In a new nuclease-free 1.5-mL low-retention microcentrifuge tube at room temperature, add the following components in the designated order:

	Order	Reagent	Cap color	Volume
<input type="checkbox"/>	1	TMPL Water	Yellow	40 µL
<input type="checkbox"/>	2	TMPL Rgnt Mix	Purple	500 µL
<input type="checkbox"/>	3	TMPL Rgnt B	Blue	300 µL
<input type="checkbox"/>	4	TMPL Enzyme Mix	Brown	50 µL

6. Cap the 1.5-mL tube and vortex for ~5 seconds, then pulse centrifuge for 3–5 seconds.
7. Vortex the TMPL ISP tube for 30 seconds, then pulse centrifuge for 3–5 seconds.
8. Vortex the library and TMPL CF-1 tubes for 5 seconds each, then pulse centrifuge for 3–5 seconds.
9. Add the following to the tube in the designated order. After each addition, cap the tube, vortex for ~5 seconds, then pulse centrifuge for 3–5 seconds.

	Order	Reagent	Cap color	Volume
<input type="checkbox"/>	1	TMPL ISP	Black	100 µL
<input type="checkbox"/>	2	TMPL CF-1	Clear	5 µL
<input type="checkbox"/>	3	DNA/RNA combined library	—	5 µL

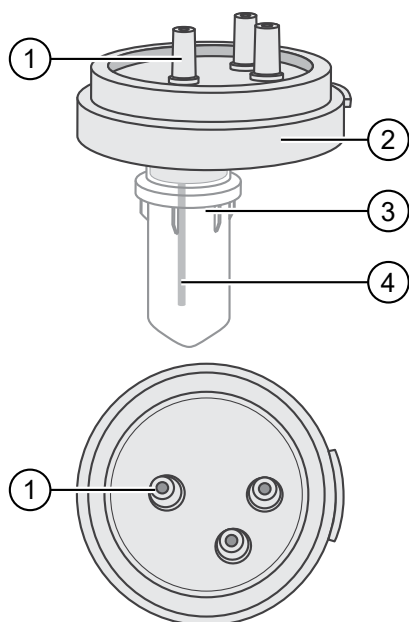
Proceed immediately to “Fill the TMPL Emulsion Cartridge” on page 40.

Fill the TMPL Emulsion Cartridge

1. Clean the Ion OneTouch™ Assembly Rack with fresh 10% bleach followed by two water rinses.
2. Label a pre-assembled TMPL Emulsion Cartridge with the Tube Label text entered into the software.

Note: Make sure that the short tube is attached to the bottom of the sample port and extends into the Reaction Tube, then push the Reaction Tube firmly onto the cartridge to ensure it is securely fastened to the cartridge.

3. Use a marker to circle the sample port on the top of the TMPL Emulsion Cartridge to distinguish it from the other two ports.



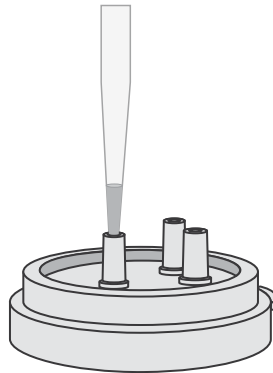
- ① Sample port (above the Reaction Tube)
- ② TMPL Emulsion Cartridge
- ③ Reaction Tube
- ④ Short tube from the sample port into the Reaction Tube

4. Place the TMPL Emulsion Cartridge into the Ion OneTouch™ Assembly Rack with the ports facing up.



① Sample port (marked as in step 3)

5. Collect the full volume of the amplification solution (~1000 μ l) using a P1000 pipettor, then insert the pipette tip vertically into the circled sample port on the cartridge with a snug fit.



6. Slowly pipet the amplification solution into the sample port.

Note: Keep the pipette and pipette tip vertical and the pipette plunger depressed when removing the tip from the port to avoid withdrawing any reaction mix or introducing air bubbles.

7. Invert the TMPL Reaction Oil 5 times to mix.
8. Using a new pipette tip, collect 750 μ L of TMPL Reaction Oil using a P1000 pipettor, then insert the pipette tip vertically into the sample port on the cartridge.
9. Slowly pipet the oil into the sample port to layer the oil over the aqueous reaction mix.

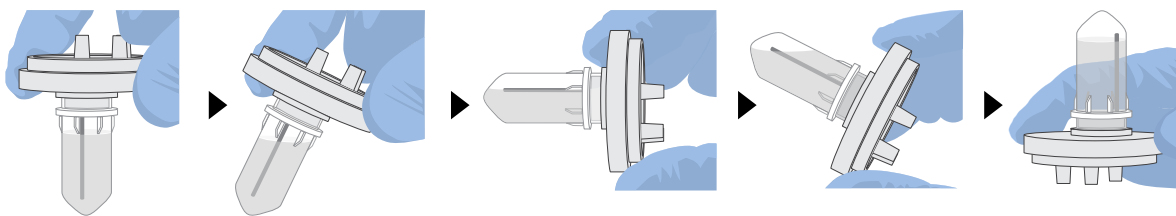
Note: Keep the pipette plunger depressed when removing the tip from the port.

10. Replace the pipette tip to avoid contamination, and repeat steps 8–9 one more time, adding another 750 µL of TMPL Reaction Oil (1.5 mL total).
11. Immediately install the TMPL Emulsion Cartridge on the instrument. Do not mix or shake the cartridge.

Install the filled TMPL Emulsion Cartridge

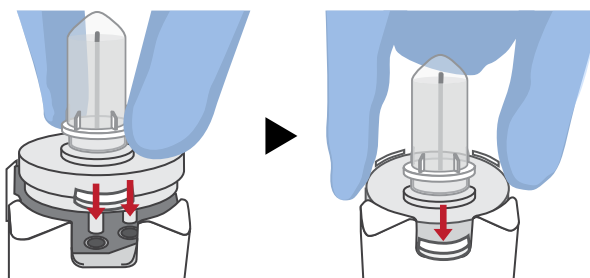
IMPORTANT! Do not mix or shake the TMPL Emulsion Cartridge during the following steps.

1. Keep the TMPL Emulsion Cartridge in the Ion OneTouch™ Assembly Rack, then orient the assembly so that the sample port is on your *left*.
2. Lift the TMPL Emulsion Cartridge straight out of the tube rack.
3. With the short tube in the Reaction Tube to the left, slowly rotate the assembly to your *right* until the Reaction Tube is inverted.



Note: This ensures minimal exposure of the short tubing in the Reaction Tube with the aqueous phase.

4. Insert the three ports of the TMPL Emulsion Cartridge into the three holes on the top of the manifold. Press firmly on all sides to ensure a secure fit on the manifold.



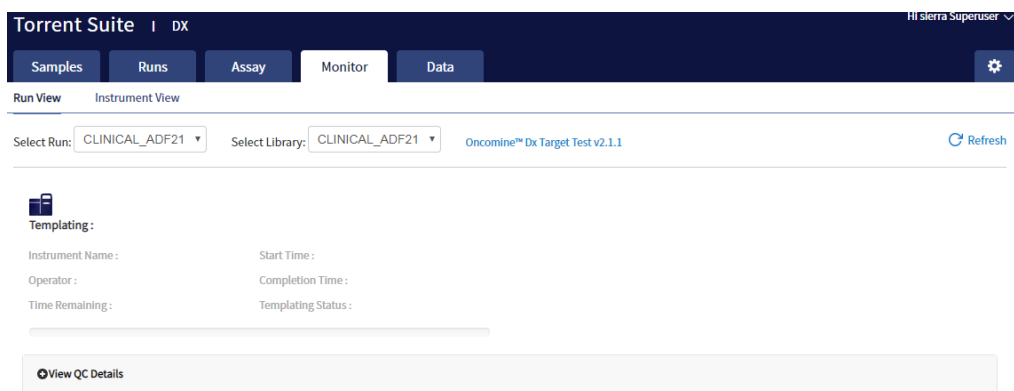
Note: After inserting the TMPL Emulsion Cartridge, bubbles may shoot up into the Reaction Tube. This is normal.

Run the Ion OneTouch™ Dx Instrument

IMPORTANT! ISPs can remain on the Ion OneTouch™ Dx Instrument for up to 24 hours after a run. You must begin enriching the ISPs on the Ion OneTouch™ ES Dx Instrument within 24 hours.

1. Ensure that the centrifuge lid of the Ion OneTouch™ Dx Instrument is closed.
2. Press **Next** on the instrument touchscreen to start the run.

Note: The run takes approximately 5 hours and 30 minutes to complete. The time remaining and a progress bar is displayed on the instrument screen during the run and under the **Monitor** tab on the Torrent Suite™ Dx Software.

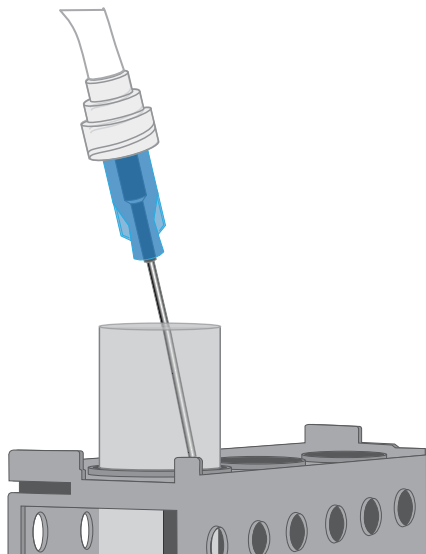


- To cancel a run, press **Abort**, then press **Yes** to confirm. If there is a high-pressure event on the instrument, the instrument aborts the run automatically.
 - If a run is aborted, you must power-cycle the Ion OneTouch™ Dx Instrument before repeating the run preparation and run. Not doing so can cause the next run to fail.
3. After the run has completed, press **Re-Spin** on the touchscreen. The time since the end of the run is displayed (*hh:mm:ss*).
 4. Based on the time displayed, do one of the following:
 - If it has been ≤ 30 minutes since the end of the run, no re-spin is required. Press **No**, then press **Next** to bypass this step. Proceed to recover the ISPs.
 - If it has been > 30 minutes since the end of the run, press **Yes** to repellet the ISPs for 10 minutes. After re-spin is complete, press **Next** and proceed to recover the ISPs.
 - If it has been > 30 minutes but the re-spin step was accidentally skipped, repellet the ISPs.
 - a. Leave the ISPs in the recovery tube on the instrument, and proceed to “Clean the Ion OneTouch™ Dx Instrument after the run” on page 51. (You can perform the cleaning procedure while the ISPs remain on the instrument.)
 - b. After cleaning is complete (~12 minutes), press **Options** on the touchscreen (Managers and Administrators only). Then select **Re-spin** to pellet the ISPs.
 - c. When the spin is complete, proceed to recover the ISPs.

Recover the ISPs from the Ion OneTouch™ Dx Instrument

The instrument touchscreen will guide the user through each step in the process. These steps are described below in more detail.

1. Remove the disposable injector from the injector hub, and carefully release the flexible tubing from the pinch valve. Place the injector into an empty 50-mL conical tube to collect waste. Press **Next**.



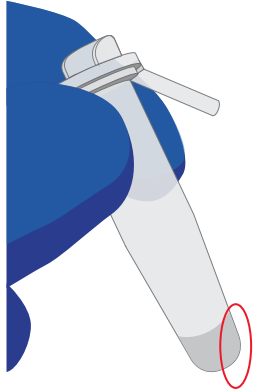
2. Press **Open Lid** on the Ion OneTouch™ Dx Instrument touchscreen, and wipe up any residue from the inside of the lid with a new disposable wipe.

Note: When using multiple instruments, use a new disposable wipe for every instrument.

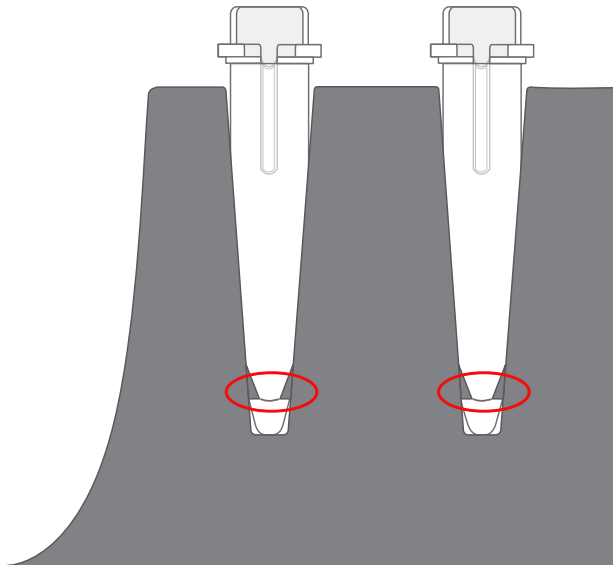
3. Remove and discard the TMPL Recovery Router.
4. Carefully remove the two TMPL Recovery Tubes from the instrument and insert them in the Ion OneTouch™ Sample Rack, with each tube arm inserted into the slot on the back of the rack. Close the instrument lid.

IMPORTANT! Do not agitate the tubes when handling. Make sure the tubes are clearly labeled with the Tube Label text.

Note: The pelleted ISPs are located at the bottom of each tube on the same side as the tube arm, as shown below. The liquid in each tube may appear cloudy and the pellet will not be visible. This is normal.



5. Depress the plunger on a P-1000 pipette and carefully insert the tip into a TMPL Recovery Tube, avoiding the pelleted ISPs. Slowly remove ~1 mL of supernatant from the top down. Switch to a P-200 pipette and use the same procedure to remove the supernatant until the meniscus at the top of the liquid aligns with the bottom of the triangular arrow guides.



IMPORTANT! Top-down removal is essential to avoid loss of ISPs. Avoid the pellet and do not eject liquid back into the tube when pipetting.

Note: The volume remaining in each tube is 50 μ L.

Prepare the TMPL ES Strip Tube

Kit components used in this procedure

Kit component	Box
TMPL ES Rsp Soln (orange cap) TMPL Wash Solution (white cap) TMPL Neutral Soln (red cap) TMPL Tween™ Solution (white cap)	Ion OneTouch™ Dx Template Solutions (Part No. A18932, stored at 15°C to 30°C)
TMPL ES Strip Tube TMPL Sample Collection Tube	Ion OneTouch™ Dx Template Supplies (Part No. A18933, stored at 15°C to 30°C)
TMPL ES Beads (green cap)	Ion OneTouch™ Dx Template ES Beads (Part No. A18931, stored at 2°C to 8°C)

Prepare the TMPL ES Beads with TMPL ES Rsp Soln

If you are processing multiple template preparations at the same time, prepare a master mix by increasing the volumes of TMPL ES Beads and TMPL ES Rsp Soln according to the table in step 5.

1. Vortex the TMPL ES Beads (green cap) for 30 seconds to resuspend the beads.
2. In a new 1.5-mL low-retention microcentrifuge tube, add 14.3 µL of TMPL ES Beads per template preparation (13 µL of beads plus 10% extra to mitigate pipetting errors).
3. Fill the tube with 1 mL of TMPL Wash Solution.
4. Cap the tube, vortex for 10 seconds, then place the tube on the DynaMag™ Dx 16 2-mL Magnet for 1 minute to capture the beads. Without disturbing the pellet, carefully remove and discard the supernatant.
5. Add 143 µL of TMPL ES Rsp Soln (orange cap) per template preparation to the tube (130 µL of solution plus 10% extra to mitigate pipetting errors).

Number of template preparations	Volume [1]	
	TMPL ES Beads	TMPL ES Rsp Soln
1	14.3 µL	143 µL
2	28.6 µL	286 µL
3	42.9 µL	429 µL
4	57.2 µL	572 µL
5	71.5 µL	715 µL
6	85.8 µL	858 µL

(continued)

Number of template preparations	Volume [1]	
	TMPL ES Beads	TMPL ES Rsp Soln
7	100.1 µL	1001 µL
8	114.4 µL	1144 µL

[1] Includes 10% extra

6. Cap the tube, then vortex for 30 seconds to resuspend the pellet.
7. If some beads are stuck to the lid of the tube, pulse centrifuge the tube for 3 seconds. Leave the tube at room temperature until ready to use.

Prepare fresh Melt-Off Solution

1. Prepare 1 M NaOH by adding 1 mL of 10 M NaOH to 9 mL Nuclease-Free water. 1 M NaOH must be prepared fresh weekly.
2. In a new 1.5-mL low-retention microcentrifuge tube, combine the following components in order.

	Order	Component	Volume
<input type="checkbox"/>	1	TMPL Tween™ Solution	280 µL
<input type="checkbox"/>	2	1 M NaOH	40 µL
		Total	320 µL

3. Cap the tube, vortex for 10 seconds, then pulse centrifuge for 3 seconds.

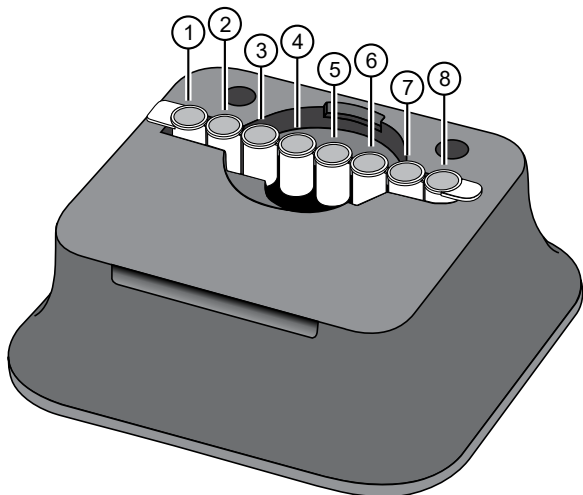
Prepare the strip tube

1. Using a marker, label a TMPL ES Strip Tube on the square tab with the Tube Label text, then place it in the Ion OneTouch™ Assembly Rack.
When facing the rack, make sure that the square tab of the TMPL ES Strip Tube is on the left and the round tab is on the right.
2. Resuspend the contents of each TMPL Recovery Tube containing the templated ISPs in 50 µL (from “Recover the ISPs from the Ion OneTouch™ Dx Instrument” on page 44, step 5) by vigorously pipetting up and down 30 times.

Note: Set a pipette to 40 µL and keep the pipette tip at the bottom of the tube when mixing to minimize air bubbles, which can lead to the loss of ISPs.

3. Pool the entire volume of templated ISPs from both recovery tubes into well 1 of the TMPL ES Strip Tube. The total pooled volume will be 100 µL.

4. Fill the remaining wells in the strip as follows:



	Well number	Reagent
<input type="checkbox"/>	Well 1	Template-positive ISPs (~100 µL), added in step 3
<input type="checkbox"/>	Well 2	130 µL of TMPL ES Beads in TMPL ES Rsp Soln
<input type="checkbox"/>	Well 3	300 µL of TMPL Wash Solution
<input type="checkbox"/>	Well 4	300 µL of TMPL Wash Solution
<input type="checkbox"/>	Well 5	300 µL of TMPL Wash Solution
<input type="checkbox"/>	Well 6	Empty
<input type="checkbox"/>	Well 7	300 µL of freshly-prepared Melt-Off Solution
<input type="checkbox"/>	Well 8	Empty

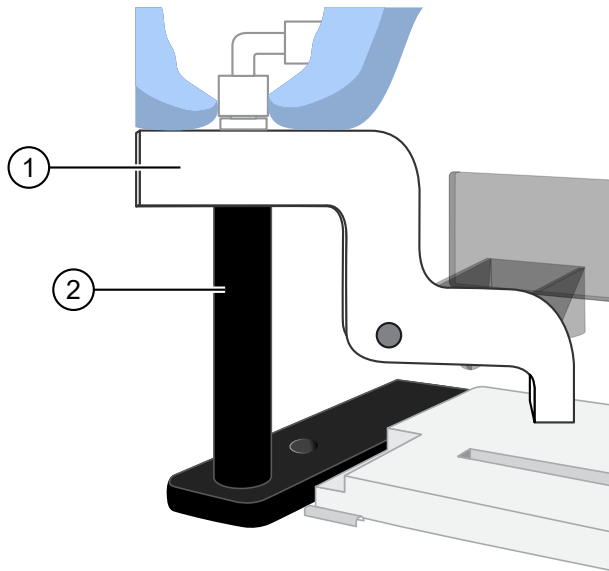
5. With a marker, label a 0.2-mL TMPL Sample Collection Tube with the Tube Label text.
6. Add 10 µL of TMPL Neutral Soln (red cap) to the TMPL Sample Collection Tube, then place it in the tube holder in the Ion OneTouch™ Sample Rack.

IMPORTANT! The volume of TMPL Neutral Soln added to the TMPL Sample Collection Tube is critical and must be accurate.

Prepare the Ion OneTouch™ ES Dx Instrument and perform the run

1. Place a new TMPL ES Tip in the Tip Loader. Remove the Tip Arm from its cradle on the Ion OneTouch™ ES Dx Instrument.
2. Grip the Tip Arm with two fingers, then align the metal fitting of the Tip Arm with the top of the tip. Firmly press the Tip Arm straight down onto the tip for 3 seconds with even pressure to ensure proper installation.

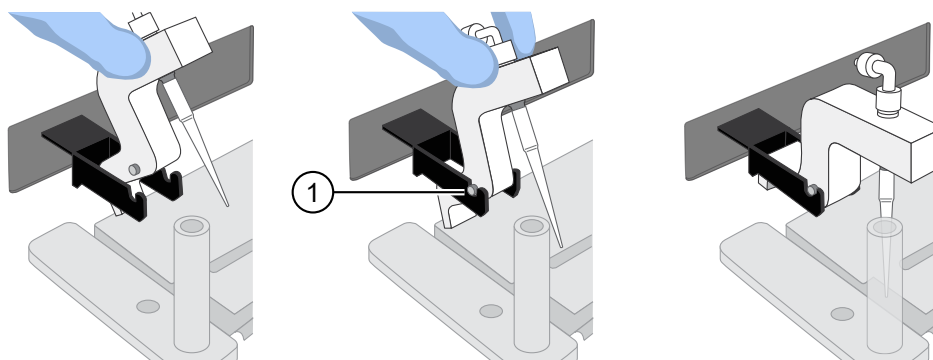
IMPORTANT! Do not repeatedly jam the Tip Arm up and down onto the tip.



- ① Tip Arm
- ② Tip Loader

3. Lift the Tip Arm straight up to pull the installed TMPL ES Tip from the Tip Loader tube.
4. Return the Tip Arm to its cradle on the instrument. Tilt the Tip Arm back and align the pins with the round notches in the cradle, then lower the Tip Arm into the home position.

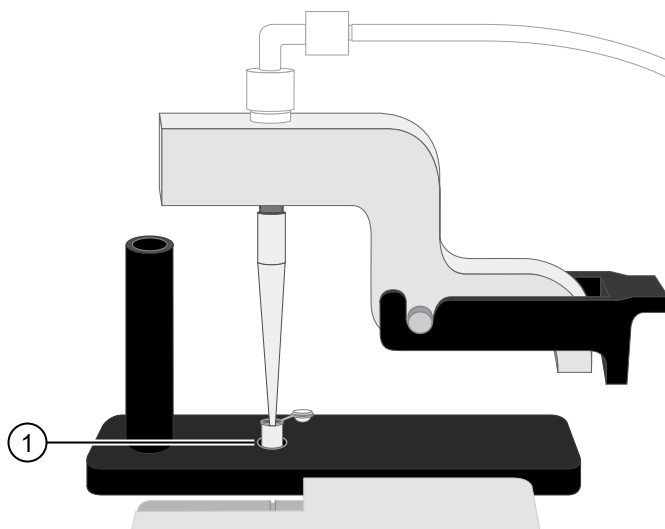
Note: Ensure that the back end of the Tip Arm is not resting on top of the thumb screw, causing the Tip Arm to tilt forward.



① Tip Arm pins resting in the notches in the cradle

5. Remove the 0.2-mL TMPL Sample Collection Tube containing TMPL Neutral Soln from the Ion OneTouch™ Sample Rack. Place it in the hole in the base of the Tip Loader. When the Tip Arm is lowered, the tip will fit inside the TMPL Sample Collection Tube, as shown below.

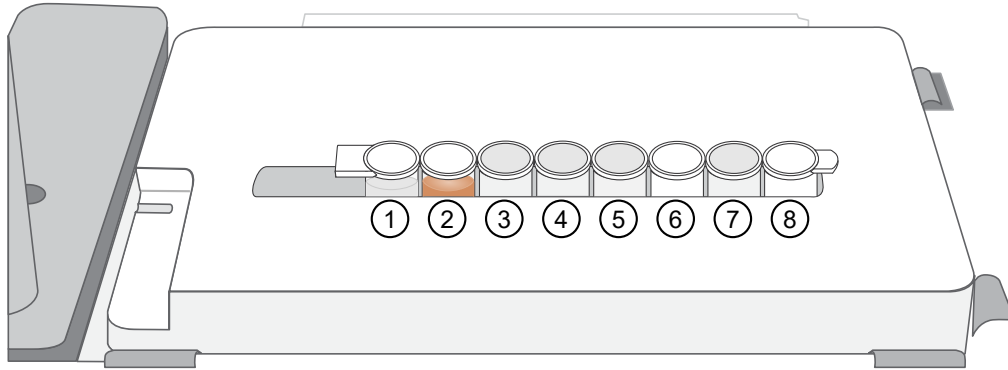
IMPORTANT! Be sure that the Tip Loader is aligned properly in its trough. If the Tip Loader is too far forward or back, the tip will miss the tube and eject sample onto the tip loader.



① TMPL Sample Collection Tube

6. Confirm that the square-shaped tab of the 8-well strip is on the left, then insert the filled strip into the **right end of the slot** on the Ion OneTouch™ ES Dx Instrument tray.

IMPORTANT! Make sure that the strip is pushed all the way to the right in the slot, away from the Tip Arm.



7. Press the **Start/Stop** button on the instrument to start the run.
The screen displays "Run" during the run, which takes ~35 minutes.
8. If you need to stop the instrument during a run, press **Start/Stop** again. The instrument completes the current step, then stops the run and displays "End". Press **Start/Stop** again to return the Tip Arm to the home position. It is not possible to restart where you left off after stopping a run.
9. During the run, proceed to "Clean the Ion OneTouch™ Dx Instrument after the run" on page 51.
10. At the end of the run, the Ion OneTouch™ ES Dx Instrument displays "End" and beeps every 60 seconds. Press the **Start/Stop** button to silence this alarm, then reset the instrument for the next run. Proceed to "Collect the sample from the Ion OneTouch™ ES Dx Instrument" on page 52.

The enriched ISPs can be left on the instrument for up to 2 hours. The instrument can be left on between runs.

Clean the Ion OneTouch™ Dx Instrument after the run

IMPORTANT! Refer to the product SDS for guidance regarding proper disposal and handling of Ion OneTouch™ Dx Instrument waste products.

1. Return to the Ion OneTouch™ Dx Instrument, then press **Next** on the touchscreen to continue with the post-run cleaning procedure.
2. Wipe any residue from the inside of the lid with a disposable wipe.
3. Remove the used TMPL Emulsion Cartridge and invert it to visually inspect that the aqueous phase was completely injected. Only one phase should be present. Discard the cartridge in the appropriate waste container.

Note: If more than one phase remains, sample injection from the assembly did not occur. See Appendix A, "Troubleshooting".

- Temporarily remove the used TMPL Amplification Plate and visually inspect it to check that there is no remaining emulsion or excessive air bubbles in the plate.

Note:

- Only clear oil should remain in the plate. Any white or cloudy areas indicate the presence of emulsion.
 - If the TMPL Amplification Plate is filled with emulsion, sample injection from the plate did not occur. See Appendix A, “Troubleshooting”. Trace amounts of emulsion are okay.
-

- Reinstall the TMPL Amplification Plate.
- Check the reagent tubes to ensure that the appropriate volume of oil and recovery solution was used. Approximately 20 mL of oil and 11 mL of TMPL Recovery Solution will be left in their respective reagent tubes. Press **Next**.

Note: Inappropriate volumes may indicate an instrument failure during the run. See Appendix A, “Troubleshooting”.

- Install a new TMPL Cleaning Adapter.
- Confirm that the disposable injector has been placed into a 50-mL conical tube.
- Press **Next** on the Ion OneTouch™ Dx Instrument touchscreen to start the cleaning run.

Note: The cleaning run takes 13 minutes. During cleaning, ensure that oil is flowing from the disposable injector. No flow of oil could indicate a clog in the manifold or in the cleaning adapter. See Appendix A, “Troubleshooting”.

- When cleaning is complete, press **Next**.

Note: If the touchscreen indicates that cleaning failed, see Appendix A, “Troubleshooting”.

- Remove the disposable injector from the 50-mL conical tube, then discard the tube and its waste into appropriate containers. Place the injector in a new 50-mL conical tube.

IMPORTANT! Do not remove the TMPL Cleaning Adapter or TMPL Amplification Plate. Leave them on the instrument.

- Press **Next** to return to the main screen, then log out of the instrument touchscreen.

Collect the sample from the Ion OneTouch™ ES Dx Instrument

- After the Ion OneTouch™ ES Dx Instrument run ends, remove the TMPL Sample Collection Tube containing the enriched ISPs and securely close the tube.

Note: The enriched ISPs can be left on the instrument for up to 2 hours.

2. Ensure that the TMPL Sample Collection Tube has >200 µL of solution.

Note: If the tube has <200 µL of solution, see Appendix A, “Troubleshooting”.

3. Remove, then discard the used tip and 8-well strip tube from the Ion OneTouch™ ES Dx Instrument.

STOPPING POINT Proceed to sequencing within 2 hours after the end of the Ion OneTouch™ ES Dx Instrument run, or transfer the template-enriched ISPs to 2–8°C storage. The ISPs may be stored for up to 1 week. If stored longer than 1 week, prepare new template-enriched ISPs from the library.



Troubleshooting

Ion OneTouch™ Dx Instrument

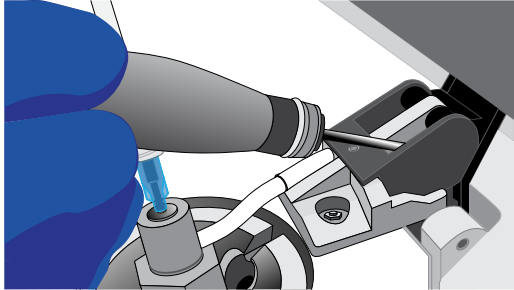
Observation	Possible cause	Recommended action
Display message: Sensor unable to measure pressure.	Hardware issue	Contact Technical Support (see Appendix F, “Customer and technical support”). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm.
Display message: Coolant pump does not flow.	Hardware issue	Contact Technical Support (see Appendix F, “Customer and technical support”). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm.
Display message: Connection failure with Ion Torrent™ Server	Ion OneTouch™ Dx Instrument and Ion Torrent™ Server connection is not established	Check that a network connection to the Ion Torrent™ Server is established, then reboot the Ion OneTouch™ Dx Instrument. Note: A sample created during a run with this alarm raised can still be used.
Display message: Failed to connect to the Ion Torrent™ Server. Check your connection.	Ion OneTouch™ Dx Instrument and Ion Torrent™ Server connection is not established during startup	Check your network connection to the Ion Torrent™ Server to make sure the connection is established, then reboot the Ion OneTouch™ Dx Instrument.
Display message: Failed to set up system time at startup. Check your connection to the Ion Torrent™ Server.	Ion OneTouch™ Dx Instrument and Ion Torrent™ Server connection is not established	Check your network connection to the Ion Torrent™ Server to make sure the connection is established, then reboot the Ion OneTouch™ Dx Instrument.
	Instrument is still in the process of establishing a connection	Allow 10 minutes to see if the display message clears.
Display message: Motor current too high. Reboot the instrument to clear the alarm.	Hardware issue	Contact Technical Support (see Appendix F, “Customer and technical support”). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm. Note: A sample created during a run with this alarm raised can still be used.
Display message: Pressure too high. Reboot the instrument to clear the alarm.	<ul style="list-style-type: none"> • Hardware issue • Clogged TMPL Emulsion Cartridge due to contaminated reagents or defective emulsion cartridge 	Reboot the Ion OneTouch™ Dx Instrument to clear the alarm. Use a new TMPL Emulsion Cartridge and fresh reagents to repeat the run. Contact Technical Support (see Appendix F, “Customer and technical support”) if the issue persists.
Display message: Sensor unable to measure instrument temperature	Hardware issue	Contact Technical Support (see Appendix F, “Customer and technical support”). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm.

Observation	Possible cause	Recommended action
Display message: Set temperature out of range. Reboot the instrument to clear the alarm.	Hardware issue	Contact Technical Support (see Appendix F, “Customer and technical support”). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm. IMPORTANT! If this alarm is raised, you cannot use the prepared template.
Display message: Software versions incompatible. Go to the Options menu and update the software.	The system software was updated, but the instrument software was not.	After the system software has been updated, update the instrument software as follows: <ol style="list-style-type: none"> 1. On the main menu of the instrument, press Options and follow the instructions to check for and install updates. 2. When installation is complete, follow the onscreen prompts to restart the instrument. IMPORTANT! You must restart the instrument before proceeding.
Display message: TEC current too high. Reboot the instrument to clear the alarm.	Hardware issue	Contact Technical Support (see Appendix F, “Customer and technical support”). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm. IMPORTANT! A sample created during a run with this alarm raised must NOT be used.
Ion OneTouch™ Dx Instrument displays a blue or grey screen with folders while the instrument is idle	Instrument issue	Power cycle the Ion OneTouch™ Dx Instrument using the On/Off switch. If alarms appear or the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).
Ion OneTouch™ Dx Instrument displays a blue or grey screen with folders during cleaning or a run	Instrument issue	Power cycle the Ion OneTouch™ Dx Instrument using the On/Off switch, then restart the cleaning or run. If alarms appear or the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).
Ion OneTouch™ Dx Instrument touchscreen freezes	Touchscreen button is pressed more than once per second.	Wait 5 minutes. If still unresponsive: <ol style="list-style-type: none"> 1. Click on the Abort button if displayed, and restart the process. 2. If Abort button is not displayed, power-cycle the instrument using the On/Off switch, then restart the process. Note: If the touchscreen freezes during Ion OneTouch™ Dx Instrument run setup, the software will remember that the instrument has already been cleaned and will not require the cleaning to be performed a second time.
During cleaning, no liquid comes out of the disposable injector	Loose or damaged reagent tube sipper in the "O" position	Reinstall the sipper in the "O" position, then restart cleaning. If the problem persists, install a new sipper.



Observation	Possible cause	Recommended action
During cleaning, no liquid comes out of the disposable injector <i>(continued)</i>	Improper installation of the TMPL Cleaning Adapter and TMPL Amplification Plate	Reinstall the TMPL Cleaning Adapter and TMPL Amplification Plate, then restart cleaning. If the problem persists, use a new TMPL Cleaning Adapter and a new TMPL Amplification Plate.
	Instrument issue	Contact Technical Support (see Appendix F, “Customer and technical support”).
Run did not complete	<ul style="list-style-type: none"> Operator did not power cycle the Ion OneTouch™ Dx Instrument after an "Abort" operation Instrument issue 	Power cycle the Ion OneTouch™ Dx Instrument using the On/Off switch, then start a new run. If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).
Cleaning was aborted due to high pressure	Improper installation of the TMPL Cleaning Adapter or TMPL Amplification Plate	Reinstall the TMPL Cleaning Adapter and TMPL Amplification Plate, then restart the cleaning protocol. If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).
Run was aborted due to high pressure	Clogged TMPL Emulsion Cartridge	Install a new TMPL Emulsion Cartridge, then restart the run. If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).
	Reagent contamination	Replace the TMPL Reaction Oil, TMPL Reagent Tubes, and TMPL Sippers, then restart the run.
	Instrument issue	Contact Technical Support (see Appendix F, “Customer and technical support”).
Oil is leaking from the instrument	Improper installation of consumables	Wipe up any leakage and reinstall TMPL Sippers and TMPL Reagent Tubes.
	Waste was not emptied before the run	Wipe up any leakage and empty the waste container before the next run.
	Instrument issue	Contact Technical Support (see Appendix F, “Customer and technical support”).
Excessive oil in waste tray	Instrument issue	Power cycle the Ion OneTouch™ Dx Instrument using the On/Off switch. If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).
Waste backup observed after completion of Ion OneTouch™ Dx Instrument run, waste leaks from the waste line after removal of the waste container	Filter in waste container clogged causing back-pressure	Remove or clean filter in waste container.
	Improper installation of consumables	Wipe up any leakage and reinstall TMPL Sippers and TMPL Reagent Tubes.
	Waste was not emptied before the run	Wipe up any leakage and empty the waste container before the next run.
	Instrument issue	Contact Technical Support (see Appendix F, “Customer and technical support”).

Observation	Possible cause	Recommended action
The centrifuge keeps running and the run never completes	Instrument hardware issue	Press Abort and power cycle the Ion OneTouch™ Dx Instrument using the On/Off switch. Press Open Lid to remove and discard the recovery tubes, then restart template preparation. If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).
Two phases are present in the sample cup at the end of the Ion OneTouch™ Dx Instrument run, sample injection from the TMPL Amplification Plate did not occur	<ul style="list-style-type: none"> The TMPL Emulsion Cartridge was not inserted properly into the instrument Problem with the instrument 	<ol style="list-style-type: none"> Power cycle the Ion OneTouch™ Dx Instrument using the On/Off switch. Repeat run preparation, then the run, being careful to seat the TMPL Emulsion Cartridge as described. <p>If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).</p>
Large air gap (1 mL or greater) is present in the reaction cup	<ul style="list-style-type: none"> Reagent tube not filled with TMPL Oil to start the run Problem with the instrument 	Repeat run preparation, then the run, being careful to follow all steps as described. If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).
Emulsion is detected in the TMPL Amplification Plate after completion of the Ion OneTouch™ Dx Instrument run	<ul style="list-style-type: none"> Reagent tube not filled with oil to start the run Improper installation of the consumables Problem with the instrument 	<ol style="list-style-type: none"> Ensure the consumables are installed correctly and not defective. Power cycle the Ion OneTouch™ Dx Instrument using the On/Off switch. Repeat run preparation, then the run, being careful to follow all steps as described. <p>If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).</p>
Inappropriate volumes of TMPL Oil and TMPL Recovery Solution are left after completion of the Ion OneTouch™ Dx Instrument run	Incorrect volumes of TMPL Oil and TMPL Recovery Solution used to start	Repeat the run, carefully check all volumes during run setup.
	<ul style="list-style-type: none"> Improper installation of the consumables Instrument issue 	<ol style="list-style-type: none"> Ensure the consumables are installed correctly and not defective. Power cycle the Ion OneTouch™ Dx Instrument using the On/Off switch. Repeat run preparation, then the run, being careful to follow all steps as described. <p>If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).</p>
Centrifuge makes a loud sound during centrifugation	There is an obstruction in the centrifuge chassis	Turn off the instrument, then contact Technical Support (see Appendix F, “Customer and technical support”).

Observation	Possible cause	Recommended action
Centrifuge lid does not open Details:	<ul style="list-style-type: none"> • Power failure • Software crash 	<ol style="list-style-type: none"> 1. Slide a 1/8-inch L-wrench (hex wrench) or equivalent tool into the right-hand hole at the top edge of the centrifuge hinge: <div data-bbox="922 394 1432 682" style="text-align: center;">  </div> 2. Press tool into the hole until there is a slight compression of the tool against the instrument and the centrifuge lid unlocks and opens. 3. Remove the tool from the hole, then open the lid. Do <i>not</i> force the lid open. <p>If the problem persists, Contact Technical Support (see Appendix F, “Customer and technical support”).</p>
Centrifuge does not spin	Ion OneTouch™ Dx Instrument centrifuge lid was not closed properly Note: Centrifuge will not operate unless the lid is fully closed.	<ol style="list-style-type: none"> 1. Open and properly close the centrifuge lid, then press re-spin. 2. After re-spin completes, power cycle the Ion OneTouch™ Dx Instrument using the On/Off switch. 3. Repeat run preparation, then the run, beginning with your pooled library sample. Be careful to properly close the centrifuge lid. <p>If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).</p>

Observation	Possible cause	Recommended action
Centrifuge does not spin <i>(continued)</i>	Instrument failure (fuse, motor driver board, and/or interlock switch failures)	<p>Confirm that the centrifuge is not operating. On the touchscreen press Options, then press respin.</p> <ul style="list-style-type: none"> • If the centrifuge does not begin to spin, contact Technical Support (see Appendix F, “Customer and technical support”). • If the centrifuge begins to spin. After re-spin completes: <ol style="list-style-type: none"> a. Power cycle the Ion OneTouch™ Dx Instrument using the On/Off switch. b. Repeat run preparation, then the run, beginning with your pooled library sample. Be careful to properly close the centrifuge lid.

Ion OneTouch™ ES Dx Instrument

Observation	Possible cause	Recommended action
Final sample volume is <200 uL	Loose or cracked tip or loose tip fitting on the Ion OneTouch™ ES Dx Instrument Tip Arm	Tighten the tip and tip fitting. If the tip is cracked, replace it.
	Improper calibration	Perform a residual volume check; if the residual volume check fails, perform calibration (see “Calibrate the Ion OneTouch™ ES Dx Instrument” on page 65). If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).
Excessive foaming	Improperly calibrated or inadequate volume in one or more wells of the TMPL ES Strip Tube	Use recommended volumes for all wells. Perform a residual volume check; if the residual volume check fails, perform calibration (see “Calibrate the Ion OneTouch™ ES Dx Instrument” on page 65). If the problem persists, contact Technical Support (see Appendix F, “Customer and technical support”).
	Loose or cracked tip, or loose tip fitting on the Ion OneTouch™ ES Dx Instrument Tip Arm	Tighten the tip and tip fitting. If the tip is cracked, replace it.



Observation	Possible cause	Recommended action
E4, E12, or E22 error displays when the Ion OneTouch™ ES Dx Instrument is initializing	<ul style="list-style-type: none"> Fuse is installed incorrectly Instrument is below operating temp Bad program or calibration setting Tip Arm is not moving 	<ol style="list-style-type: none"> Ensure that the fuse module is installed correctly and that the unit is within its recommended operating temperature range of 68°F to 86°F (20°C to 30°C). Reboot the instrument: Power OFF the instrument, wait 3 seconds, then power ON the instrument. If the error persists, restore the factory defaults, then recalibrate the instrument (see “Calibrate the Ion OneTouch™ ES Dx Instrument” on page 65).
Solution overflow during a run	Overloaded reagent volumes in TMPL ES Strip Tube	Repeat enrichment with correct reagent volumes.
Tip is causing the 8-well strip to lift out of its slot during run	Tip is not aligned vertically	Try tightening the tip. If the problem persists, perform vertical axis calibration (see “Vertical axis calibration” on page 65).
Strip lifts up during strip push	Instrument is not calibrated properly	Perform horizontal position calibration.
Strip lifts up when tip is raised from well	Instrument is not calibrated properly	Perform vertical calibration.
Immediately after strip push, the strip is not in contact with the magnet	Instrument is not calibrated properly	Perform horizontal position calibration.
Tip grinds into the base of the instrument and Code "1999" displays	<ul style="list-style-type: none"> Vertical calibration setting too low or out of range Instrument is not calibrated properly 	<ol style="list-style-type: none"> Erase the memory on the instrument: Hold down the vertical-adjust button while powering ON the instrument. The instrument beeps several times. Perform a residual volume test. Recalibrate the instrument if residual volume check failed.
Tip is hitting the top of tray at start of run	<ul style="list-style-type: none"> Instrument tray or tip is not properly seated in the instrument Tip adapter is loose 	<ol style="list-style-type: none"> Check for debris between the tray and the instrument, then reinstall the tray and tip. Check the tip adapter to make sure it is tight.
Error displays	Various	<ol style="list-style-type: none"> Power the instrument OFF then ON. If the error continues to display, erase the memory on the instrument. Hold down the vertical adjust button while powering ON the instrument. The instrument beeps several times. Perform residual volume check. Recalibrate the instrument if the residual volume check failed.

Observation	Possible cause	Recommended action
Instrument does not aspirate or dispense liquids	Loose fittings	<ul style="list-style-type: none"> • Ensure that the connections at the elbow on the Tip Arm and at the tubing on the rear syringe pump are finger-tight. • Ensure that the metal tip adapter fitting on the Tip Arm is finger-tight. <p>IMPORTANT! After any changes to the metal tip adapter, perform a remaining volume test, and recalibrate the instrument.</p>
Ion OneTouch™ ES Dx Instrument has a blown fuse	Various	Contact Technical Support (see Appendix F, “Customer and technical support”).



Supplemental procedures and instruments

Fuse replacement on the Ion OneTouch™ ES Dx Instrument

IMPORTANT! The Ion OneTouch™ ES Dx Instrument is supplied with a Fuse Module and two different types of spare fuses, which should only be replaced by trained field service engineers. The Fuse Module is installed by the field service engineer into the Power Entry Module located on the back of the instrument in the proper orientation for the voltage in your area. If you are not sure of the setting that is right for your area, contact your local power company.

Line voltage	Replacement fuse type required
110/120 VAC	375 mA TT (Slow Blow) 1/4 inches x 1--1/4 inches
220/240 VAC	160 mA TT (Slow Blow) 5 x 20 mm



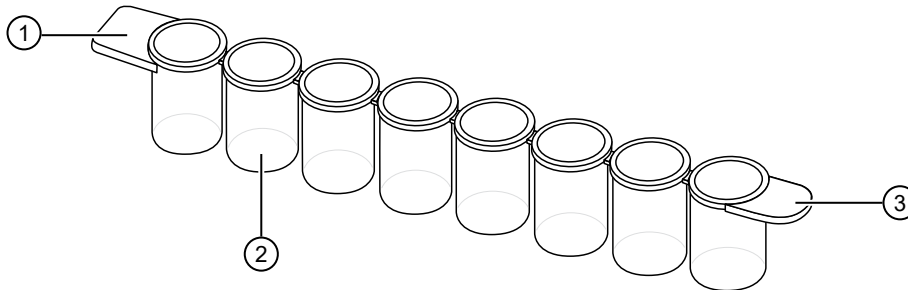
WARNING! ELECTRICAL SHOCK HAZARD. Severe electrical shock, which could cause physical injury or death, can result from working on an instrument when the high voltage power supply is operating. To avoid electrical shock, disconnect the power supply to the instrument, unplug the power cord, and wait at least 1 minute before working on the instrument.

Ion OneTouch™ ES Dx Instrument residual volume test

1. Install a new tip on the Ion OneTouch™ ES Dx Instrument Tip Arm.

Note: For the residual volume test, you do not need to put a TMPL Sample Collection Tube in the Tip Loader.

2. Load 80 µL water or TMPL Wash Solution into the second well (Well 2) from the square-tab end of the 8-well strip:



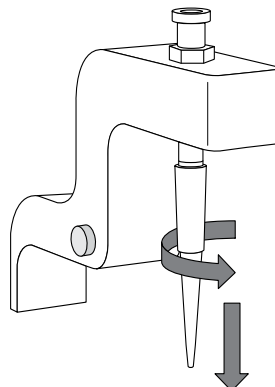
- ① Square tab
- ② Second well
- ③ Round tab

3. Load the 8-well strip into the right end of the slot on the Ion OneTouch™ ES Dx Instrument tray so that the square tab is on the left and the rounded tab is on the right.

IMPORTANT! Before proceeding, carefully read and familiarize yourself with the following steps, which require you to manually start and stop the test run and manipulate the strip tube during the run. During the test, confirm that the tip is centered in the wells when moving in or out of a well.

4. Turn the instrument ON.
5. Wait for the instrument to initialize. The screen displays "rdy". The Tip Arm performs a series of movements and returns to the home position (~5 seconds).
6. Press **Start/Stop**.
7. Wait for the instrument to aspirate the solution from Well 2 and completely remove the tip from Well 2, then *manually* push the 8-well strip to the left so that Well 4 is positioned directly under the Tip Arm.
8. Wait for the instrument to dispense the tip contents into Well 4.
9. Press **Start/Stop** to stop the test run, then press **Start/Stop** again to return the Tip Arm to the home position.
10. Using a P10 pipette, aspirate the entire residual volume from Well 2, then estimate the residual volume.

11. Remove the used tip: with the Tip Arm in its cradle and while standing above the Tip Arm, twist the tip *counterclockwise* and pull it downward to remove and discard the tip.



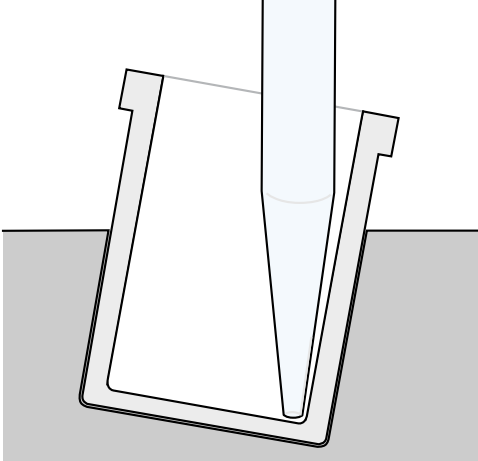
IMPORTANT! Improper removal of tips can loosen the metal tip adapter fitting on the Tip Arm and affect instrument operation.

12. Remove and discard the used 8-well strip.
13. After performing the residual volume test, take one or more of the following actions:

Observation	Pass/Fail	Possible cause	Recommended actions
Residual volume in Well 2 is $\leq 5 \mu\text{L}$	Pass	—	—
Residual volume in Well 2 is $> 5 \mu\text{L}$ IMPORTANT! The volume is measured from the <i>bottom</i> of the well, not from the sides.	Fail	The tip height is too high during aspiration.	Calibrate the instrument (see “Calibrate the Ion OneTouch™ ES Dx Instrument” on page 65).
The 8-well strip lifts as the tip rises to the top of the well	Fail	The tip is angled too far forward or the tip height is set too low.	Verify that the tip is vertical and positioned directly over the notch in the calibration shelf. If the tip is positioned correctly, restore defaults, then calibrate the instrument (see “Calibrate the Ion OneTouch™ ES Dx Instrument” on page 65).

Calibrate the Ion OneTouch™ ES Dx Instrument

Perform horizontal and vertical calibrations so that during operation the tip is optimally positioned in the well of the 8-well strip:



Note that the 8-well strip is always tilted at a fixed 10-degree angle in the slot. The pipette tip is vertical. When the tip is aligned properly during calibration so that it is in line with the notch in the calibration shelf, the tip touches the front-bottom edge of the well during the run.

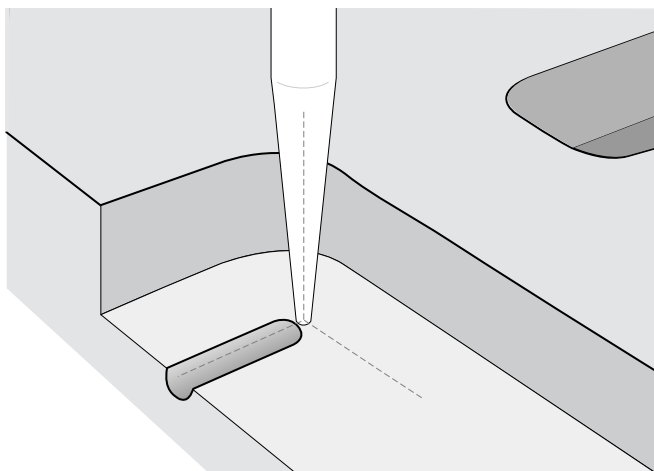
IMPORTANT! If you use more than one Ion OneTouch™ ES Dx Instrument, do not switch Trays or Tip Arms between instruments. Each Tray and Tip Arm is calibrated with a particular instrument. To track the Tray and Tip Arm, each component has a printed label with the matching serial number of the instrument.

Vertical axis calibration

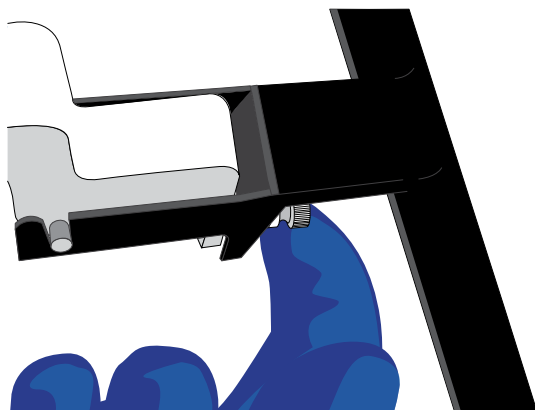
1. Install a new tip.
2. Restore the factory default settings:
 - a. Power OFF the instrument, then wait 3 seconds.
 - b. While holding down **Vert. Adjust**, power ON the instrument.
3. Put the instrument into calibration mode:
 - a. Power OFF the instrument.
 - b. While holding down **Select/Calibrate**, power the instrument ON. Keep holding down **Select/Calibrate** until "P1" is displayed.
 - c. Press **Select/Calibrate** for ~3 seconds until the instrument beeps 2 times and "CAL" is displayed.

Note: The instrument will cycle through several values before "CAL" is displayed.

4. Press **Vert. Adjust**. The instrument displays "ASP" (Aspirate or z-bottom position).
5. Press **Start/Stop**. The Tip Arm lowers to bring the tip near the notch in the calibration shelf on the left side of the Tray.
6. The tip should be positioned in line with the slot in the calibration shelf, and the tip should be touching the shelf. If necessary, adjust the tip as follows:



- a. To adjust the alignment of the tip with the slot, turn the thumbscrew at the back of the Tip Arm.



- b. To adjust the height of the tip, press the ▼ (minus) button repeatedly until the tip touches the shelf. Press the button eight more times to lower the tip further. This will account for variations in tip lengths and installation.

Note: It is better to have the ASP (aspiration) height be too low than too high.

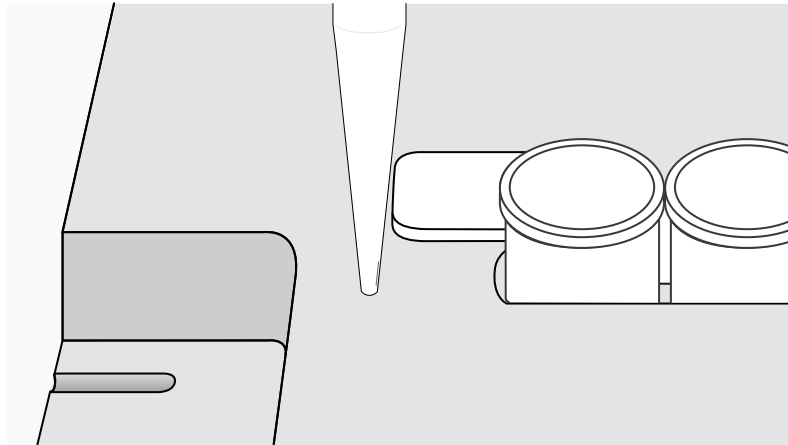
7. Press **Start/Stop**, then wait for the Tip Arm to stop moving and for "P1" to display.



Horizontal axis calibration

1. Press **Select/Calibrate** for ~3 seconds until the instrument beeps 2 times and “CAL” is displayed.

Note: While you press the button, the instrument cycles through several values before “CAL” is displayed.

2. Press **Horiz. Adjust**. The instrument displays “FLA”. Press **Start/Stop**.
3. Place an empty 8-well strip in the slot in the Tray, with the square tab on the left.
4. Push the 8-well strip as far to the left in the slot as possible.
5. Observe the position of the 8-well strip relative to the position of the tip. When properly calibrated, the 8-well strip is within 1 mm of touching the tip, but not pushing on it. To clearly see the relationship between the pipette tip, calibration shelf, and notch during calibration, mark each of them with a felt-tip pen:



6. Adjust the horizontal position of the Tip Arm so that the tip just touches the square tab on the left of the 8-well strip when the 8-well strip is pushed to the far left of the slot in the Tray:
 - a. Apply slight pressure to keep the 8-well strip to the far left.
 - b. Press the  (plus) button repeatedly until the tip touches the 8-well strip. Each press of the  (plus) key moves the Tip Arm to the *right* by ~0.002 inches (~50 µm), which may be difficult to detect.
7. Press **Start/Stop** to save the setting, then wait for “P1” to display.
8. Power the instrument OFF, wait >3 seconds, then power the instrument ON to return to normal operating mode.
9. Perform a residual volume test (see “Ion OneTouch™ ES Dx Instrument residual volume test” on page 62).

Removal of instruments from use for repair or disposal

To remove the Ion OneTouch™ Dx Instrument or Ion OneTouch™ ES Dx Instrument from use for repair or disposal, perform the following steps:

IMPORTANT! This procedure does not guarantee total decontamination of the Ion OneTouch™ Dx Instrument or Ion OneTouch™ ES Dx Instrument.

1. Wear disposable gloves, safety glasses, and a lab coat.
2. Use a cleaning pad wetted with a solution of 1 part chlorine bleach in 9 parts water (10% bleach solution) to clean all outside surfaces of the Ion OneTouch™ Dx Instrument or Ion OneTouch™ ES Dx Instrument. Use care to avoid getting bleach solution inside the chassis.
3. Dry the surfaces of the instrument with paper towels or other disposable wipes.
4. Use cotton swabs to clean and dry areas that are difficult to access.
5. Properly dispose of all used consumables (including tubes, sippers, amplification plates) and cleaning materials to ensure that no one becomes exposed to contaminants.

If returning the instrument, contact your local Thermo Fisher Scientific representative to schedule a pickup of the instrument.



WARNING! GENERAL SAFETY. Using this product in a manner not specified in the user documentation may result in personal injury or damage to the instrument or device. Ensure that anyone using this product has received instructions in general safety practices for laboratories and the safety information provided in this document.

- Before using an instrument or device, read and understand the safety information provided in the user documentation provided by the manufacturer of the instrument or device.
- Before handling chemicals, read and understand all applicable Safety Data Sheets (SDSs) and use appropriate personal protective equipment (gloves, gowns, eye protection, and so on). To obtain SDSs, see the “Documentation and Support” section in this document.

Chemical safety



WARNING! GENERAL CHEMICAL HANDLING. To minimize hazards, ensure laboratory personnel read and practice the general safety guidelines for chemical usage, storage, and waste provided below. Consult the relevant SDS for specific precautions and instructions:










- Read and understand the Safety Data Sheets (SDSs) provided by the chemical manufacturer before you store, handle, or work with any chemicals or hazardous materials. To obtain SDSs, see the “Documentation and Support” section in this document.
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing).
- Minimize the inhalation of chemicals. Do not leave chemical containers open. Use only with adequate ventilation (for example, fume hood).
- Check regularly for chemical leaks or spills. If a leak or spill occurs, follow the manufacturer's cleanup procedures as recommended in the SDS.
- Handle chemical wastes in a fume hood.
- Ensure use of primary and secondary waste containers. (A primary waste container holds the immediate waste. A secondary container contains spills or leaks from the primary container. Both containers must be compatible with the waste material and meet federal, state, and local requirements for container storage.)
- After emptying a waste container, seal it with the cap provided.
- Characterize (by analysis if necessary) the waste generated by the particular applications, reagents, and substrates used in your laboratory.
- Ensure that the waste is stored, transferred, transported, and disposed of according to all local, state/provincial, and/or national regulations.
- **IMPORTANT!** Radioactive or biohazardous materials may require special handling, and disposal limitations may apply.



Symbols on this instrument

Symbols may be found on the instrument to warn against potential hazards or convey important safety information. In this document, the hazard symbol is used along with one of the following user attention words:

- **CAUTION!**—Indicates a potentially hazardous situation that, if not avoided, may result in minor or moderate injury. It may also be used to alert against unsafe practices.
- **WARNING!**—Indicates a potentially hazardous situation that, if not avoided, could result in death or serious injury.
- **DANGER!**—Indicates an imminently hazardous situation that, if not avoided, will result in death or serious injury.

Symbol	English	Français
	Caution, risk of danger Consult the manual for further safety information.	Attention, risque de danger Consulter le manuel pour d'autres renseignements de sécurité.
	Caution, risk of electrical shock	Attention, risque de choc électrique
	Caution, piercing hazard	Attention, danger de perforation
	Caution, hot surface	Attention, surface chaude
	Potential biohazard	Danger biologique potentiel
	On	On (marche)
	Off	Off (arrêt)
	Earth (ground) terminal	Borne de (mise à la) terre
	Protective conductor terminal (main ground)	Borne de conducteur de protection (mise à la terre principale)



(continued)

Symbol	English	Français
	<p>Do not dispose of this product in unsorted municipal waste</p> <p>⚠ CAUTION! To minimize negative environmental impact from disposal of electronic waste, do not dispose of electronic waste in unsorted municipal waste. Follow local municipal waste ordinances for proper disposal provision and contact customer service for information about responsible disposal options.</p>	<p>Ne pas éliminer ce produit avec les déchets usuels non soumis au tri sélectif.</p> <p>⚠ MISE EN GARDE ! Pour minimiser les conséquences négatives sur l'environnement à la suite de l'élimination de déchets électroniques, ne pas éliminer ce déchet électronique avec les déchets usuels non soumis au tri sélectif. Se conformer aux ordonnances locales sur les déchets municipaux pour les dispositions d'élimination et communiquer avec le service à la clientèle pour des renseignements sur les options d'élimination responsable.</p>

Conformity symbols on the instrument

Conformity mark	Description
	Indicates conformity with safety requirements for Canada and U.S.A.
	Indicates conformity with European Union requirements for safety and electromagnetic compatibility.
	Indicates conformity with Australian standards for electromagnetic compatibility.






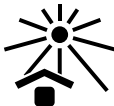









Medical device symbols

The following table describes symbols that may be displayed on product labels. The symbols that are used on labels conform to standards BS EN ISO 15223-1:2016 and FDA 21 CFR 809.10 "Labeling for in vitro diagnostic products".

Symbol	Description	Symbol	Description
	MANUFACTURER		CONTAINS SUFFICIENT FOR <n> TESTS
	DATE OF MANUFACTURE		USE BY







(continued)

Symbol	Description	Symbol	Description
	BATCH CODE		CATALOG NUMBER
	SERIAL NUMBER		FRAGILE, HANDLE WITH CARE
	LOWER LIMIT OF TEMPERATURE		PROTECT FROM LIGHT
	UPPER AND LOWER LIMITS OF TEMPERATURE		UPPER LIMIT OF TEMPERATURE
	DO NOT REUSE		BIOLOGICAL RISKS
	CAUTION, CONSULT ACCOMPANYING DOCUMENTS		CONSULT INSTRUCTIONS FOR USE
	UPPER AND LOWER LIMITS OF HUMIDITY		OBSERVE PRECAUTIONS FOR HANDLING ELECTROSTATIC SENSITIVE DEVICES
	IN VITRO DIAGNOSTIC MEDICAL DEVICE		



Safety alerts on this instrument

Additional text may be used with one of the symbols described above when more specific information is needed to avoid exposure to a hazard. See the following table for safety alerts found on the instrument.

English	Français
 CAUTION! Hazardous chemicals. Read the Safety Data Sheets (SDSs) before handling.	 MISE EN GARDE ! Produits chimiques dangereux. Lire les fiches signalétiques (FS) avant de manipuler les produits.
 CAUTION! Hazardous waste. Refer to SDS(s) and local regulations for handling and disposal.	 MISE EN GARDE ! Déchets dangereux. Lire les fiches signalétiques (FS) et la réglementation locale associées à la manipulation et à l'élimination des déchets.

Safety information for third-party instruments

Refer to the manufacturer's documentation for information on the safe use of third-party products provided with the instrument system.

Instrument safety

General



CAUTION! Do not remove instrument protective covers. If you remove the protective instrument panels or disable interlock devices, you may be exposed to serious hazards including, but not limited to, severe electrical shock, laser exposure, crushing, or chemical exposure.



CAUTION! Solvents and Pressurized fluids. Wear eye protection when working with any pressurized fluids. Use caution when working with any polymeric tubing that is under pressure:

- Extinguish any nearby flames if you use flammable solvents.
- Do not use polymeric tubing that has been severely stressed or kinked.
- Do not use polymeric tubing with tetrahydrofuran or nitric and sulfuric acids.
- Be aware that methylene chloride and dimethyl sulfoxide cause polymeric tubing to swell and greatly reduce the rupture pressure of the tubing.
- Be aware that high solvent flow rates (~40mL/min) may cause a static charge to build up on the surface of the tubing and electrical sparks may result.



Physical injury



CAUTION! Moving Parts. Moving parts can crush, pinch and cut. Keep hands clear of moving parts while operating the instrument. Disconnect power before servicing.



CAUTION! ROTATION HAZARD. Wait until rotation stops before opening. Rotating parts can cause injury

Electrical safety



WARNING! Fuse Installation. Before installing the instrument, verify that the fuses are properly installed and the fuse voltage matches the supply voltage. Replace fuses only with the type and rating specified for the unit. Improper fuses can damage the instrument wiring system and cause a fire.



WARNING! Ensure appropriate electrical supply. For safe operation of the instrument:

- Plug the system into a properly grounded receptacle with adequate current capacity.
- Ensure the electrical supply is of suitable voltage.
- Never operate the instrument with the ground disconnected. Grounding continuity is required for safe operation of the instrument.



WARNING! Power Supply Line Cords. Use properly configured and approved line cords for the power supply in your facility.



WARNING! Disconnecting Power. To fully disconnect power either detach or unplug the power cord, positioning the instrument such that the power cord is accessible.

Cleaning and decontamination



CAUTION! Cleaning and Decontamination. Use only the cleaning and decontamination methods specified in the manufacturer's user documentation. It is the responsibility of the operator (or other responsible person) to ensure the following requirements are met:

- No decontamination or cleaning agents are used that could cause a HAZARD as a result of a reaction with parts of the equipment or with material contained in the equipment.
- The instrument is properly decontaminated a) if hazardous material is spilled onto or into the equipment, and/or b) prior to having the instrument serviced at your facility or sending the instrument for repair, maintenance, trade-in, disposal, or termination of a loan (decontamination forms may be requested from customer service).
- Before using any cleaning or decontamination methods (except those recommended by the manufacturer), users should confirm with the manufacturer that the proposed method will not damage the equipment.



Laser



CAUTION! LASER HAZARD, Bar Code Scanner. The bar code scanner included with the instrument system is a Class 2 laser. To avoid damage to eyes, do not stare directly into the beam or point into another person's eyes.

Safety and electromagnetic compatibility (EMC) standards

The instrument design and manufacture complies with the following standards and requirements for safety and electromagnetic compatibility.

Safety

Reference	Description
Directive 2006/95/EC	European Union "Low Voltage Directive"
IEC 61010-1 EN 61010-1 CSA C22.2 61010-1	Safety requirements for electrical equipment for measurement, control, and laboratory use
IEC 61010-2-010 EN 61010-2-010 CSA C22.2 61010-2-010	Safety requirements for electrical equipment for measurement, control and laboratory use — Part 2-010: Particular requirements for laboratory equipment for the heating of materials
IEC 61010-2-020 EN 61010-2-020 CSA C22.2 61010-2-020	Safety requirements for electrical equipment for measurement, control and laboratory use — Part 2-020: Particular requirements for laboratory centrifuges
IEC 61010-2-081 EN 61010-2-081 CSA C22.2 61010-2-081	Safety requirements for electrical equipment for measurement, control and laboratory use — Part 2-081: Particular requirements for automatic and semi-automatic laboratory equipment for analysis and other purposes
IEC 61010-2-101	Safety Requirements for Electrical Equipment for Measurement, Control and Laboratory Use — Part 2-101: Particular Requirements for In Vitro Diagnostic (IVD) Medical Equipment



EMC

Reference	Description
Directive 2004/108/EC	European Union “EMC Directive”
EN 61326-1	<i>Electrical Equipment for Measurement, Control and Laboratory Use – EMC Requirements – Part 1: General Requirements</i>
EN 61326-2-6	<i>Electrical Equipment for Measurement, Control and Laboratory Use – EMC Requirements – Part 26: Particular requirements – In vitro diagnostic (IVD) medical equipment)requirements</i>
FCC Part 15	U.S. Standard “Industrial, Scientific, and Medical Equipment”
AS/NZS CISPR 22:2009	<i>Limits and Methods of Measurement of Electromagnetic Disturbance Characteristics of Industrial, Scientific, and Medical (ISM) Radiofrequency Equipment</i>
ICES-003, Issue 5	<i>Industrial, Scientific and Medical (ISM) Radio Frequency Generators</i>

Environmental design

Reference	Description
Directive 2012/19/EU	European Union “WEEE Directive” – Waste electrical and electronic equipment
Directive 2011/65/EU	European Union “RoHS Directive” – Restriction of hazardous substances in electrical and electronic equipment

Precaution—strong magnet

Note: Do not substitute non-IVD labeled magnets for the DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet, provided with Ion PGM™ Dx System.

The DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet contain very strong permanent magnets. People wearing a pacemaker or any other medical magnetized implant should not use this product unless advised by a health professional; the implant could be affected or damaged by exposure to a strong magnetic field. Keep tools and objects that could be damaged by the magnetic field out of the working area. This includes, but is not restricted to, credit cards and other products containing magnetic recording devices. Keep away from delicate instruments, watches, electronic equipment, displays and monitors. The magnet may attract steel or other magnetic material with high mechanical forces. Take care during handling. Avoid contact between two magnets. Do not pull the magnets apart if contact has been made; twist off to prevent damage to the unit or fingers. The Health and Safety Officer should take all necessary steps and full responsibility to ensure that the precautions and statements are followed and adhered to.



Performance characteristics

For performance characteristics of the Oncomine™ Dx Target Test Kit, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

For performance characteristics of the Ion PGM™ Dx System, see the *Ion PGM™ Dx System Performance Characteristics User Guide* (Pub. No. MAN0018763).



Instrument warranty

For new Ion Torrent™ instruments, Life Technologies warrants to and only to buyer for twelve (12) months from the date of shipping, that the Ion Torrent™ software and Ion Torrent™ instruments are free from defects in material and workmanship and conform to Life Technologies' published specifications in all material respects. Where a valid and timely claim in respect of breach of Ion Torrent™ warranty is submitted to Life Technologies, Life Technologies may, at its discretion, replace, repair or modify the Ion Torrent™ instrument. Any agreed replacement shall be at 1:1, like-kind basis, at no cost to the buyer. For Ion Torrent™ chips or reagents reasonably determined by Life Technologies to be defective, independent of user error, shall be replaced by Life Technologies on a 1:1, like-kind basis at no cost to buyer, provided that such defective Ion Torrent™ chips or reagents were used by buyer prior to their expiration date, or if there is no expiration date, the Ion Torrent™ chips or reagents were used within six (6) months of receipt, and the defect was promptly reported with appropriate detail to Life Technologies' technical support.

NO OTHER WARRANTIES SHALL BE APPLICABLE TO ION TORRENT PRODUCTS (WHETHER OR NOT ANY FURTHER WARRANTY DOCUMENTATION MAY BE INCLUDED IN THE SHIPMENT), WITH THE EXCEPTION OF THIRD PARTY WARRANTIES WITH RESPECT TO THIRD PARTY PRODUCT. ANY THIRD PARTY PRODUCTS ARE NOT COVERED BY THIS SECTION AND ANY WARRANTIES FOR THIRD PARTY PRODUCTS ARE PROVIDED BY THE ORIGINAL MANUFACTURER OF THE THIRD PARTY PRODUCT. Warranties are made only to buyer purchasing the Ion Torrent™ Product directly from Life Technologies, are not transferable and do not extend to the benefit of any other person or entity, unless otherwise expressly stated in writing by Life Technologies. ANY PRODUCT NOT COVERED BY AN EXPRESS WRITTEN WARRANTY IS SOLD AND PROVIDED "AS IS," WITHOUT WARRANTY OF ANY KIND, STATUTORY, EXPRESS OR IMPLIED. Any description of Ion Torrent™ Product recited in Life Technologies' quotation is for the sole purpose of identifying Ion Torrent™ Product, and any such description is not part of any contract between Life Technologies and buyer and does not constitute a warranty that Ion Torrent™ Product shall conform to that description. Any sample or model used in connection with Life Technologies' quotation is for illustrative purposes only, and is not part of any contract between Life Technologies and buyer and does not constitute a warranty that Ion Torrent™ Product will conform to the sample or model. No affirmation of fact or promise made by Life Technologies, whether or not in Life Technologies' quotation, shall constitute a warranty that Ion Torrent™ Product will conform to the affirmation or promise. Unless otherwise specified in writing in documentation shipped with Ion Torrent™ Product or otherwise agreed by Life Technologies in writing. Life Technologies does not provide service or support for custom products or other products made to buyer's specifications. THE WARRANTIES IDENTIFIED IN THIS CLAUSE ARE LIFE TECHNOLOGIES' SOLE AND EXCLUSIVE WARRANTIES WITH RESPECT TO Ion Torrent™ PRODUCT AND ARE IN LIEU OF ALL OTHER WARRANTIES, STATUTORY, EXPRESS OR IMPLIED, ALL OF WHICH OTHER WARRANTIES ARE EXPRESSLY DISCLAIMED, INCLUDING WITHOUT LIMITATION ANY IMPLIED WARRANTY OF MERCHANTABILITY, FITNESS FOR A PARTICULAR PURPOSE, NON-INFRINGEMENT, OR REGARDING RESULTS OBTAINED THROUGH THE USE OF ANY PRODUCT (INCLUDING, WITHOUT LIMITATION, ANY CLAIM OF INACCURATE, INVALID OR INCOMPLETE RESULTS), WHETHER ARISING FROM A STATUTE OR OTHERWISE IN LAW OR FROM A COURSE OF PERFORMANCE, DEALING OR USAGE OF TRADE.



Customer and technical support

Visit [thermofisher.com/support](https://www.thermofisher.com/support) for the latest in services and support, including:

- Worldwide contact telephone numbers
- Product support
- Order and web support
- Safety Data Sheets (SDSs; also known as MSDSs)

Additional product documentation, including user guides and Certificates of Analysis, are available by contacting Customer Support.

Obtaining Certificates of Analysis

The Certificate of Analysis provides detailed quality control and product qualification information for each product. Certificates of Analysis are printed and shipped with the product.

Obtaining Certificates of Conformance

The Certificate of Conformance provides information on conformance testing of each instrument provided with the system. Certificates of Conformance are shipped with the instrument, and are also available by contacting Customer Support at [thermofisher.com/support](https://www.thermofisher.com/support).

Oncomine™ Dx Target Test Part IV: Sequencing and Results Reports

USER GUIDE

Publication Number MAN0018706

Revision B.0



For In Vitro Diagnostic Use.

ThermoFisher
S C I E N T I F I C



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Life Technologies Corporation |
7335 Executive Way |
Frederick, MD 21704 | USA

Products manufactured in Singapore:

Ion PGM™ Dx Instrument System
Ion PGM™ Dx Sequencer
Ion OneTouch™ Dx Instrument
Ion OneTouch™ ES Dx Instrument
Ion PGM™ Dx Chip Minifuge (120V)
Ion PGM™ Wireless Scanner
Ion Torrent™ Server
Torrent Suite™ Dx Software
Veriti™ Dx 96-well Thermal Cycler, 0.2 mL

Products manufactured in Frederick:

Oncomine™ Dx Target Test Kit
Ion Torrent Dx FFPE Sample Preparation Kit
Ion PGM™ Dx Library Kit
Ion OneTouch™ Dx Template Kit
Ion PGM™ Dx Sequencing Kit
Ion 318™ Dx Chip
Ion OneTouch™ Rack Kit
DynaMag™ Dx 96-Well Plate Magnet
DynaMag™ Dx 16 2-mL Magnet

The information in this guide is subject to change without notice.

DISCLAIMER

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Revision history: Pub. No. MAN0018706

Revision	Date	Description
B.0	3 September 2021	<ul style="list-style-type: none"> Updated with new product catalog numbers. Deleted Chapter 7, "Pass/fail criteria and repeat strategy". Users referred to the part I user guide for this content.
A.0	11 February 2021	New Oncomine™ Dx Target Test user guide for FDA submission.

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About this guide

IMPORTANT! Before using this product, read and understand the information in the “Safety” appendix in this document.

Purpose of this guide

This user guide provides instructions for sequencing enriched, template-positive Ion PGM™ Dx Ion Sphere™ Particles (ISPs) that have been prepared from Oncomine™ Dx Target Test libraries. Sequencing is performed using the Ion PGM™ Dx System with the Ion PGM™ Dx Sequencing Kit and the Ion 318™ Dx Chip.

Oncomine™ Dx Target Test Kit user guides

This user guide is part of a five-guide set.

- *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*
- *Oncomine™ Dx Target Test Part II: Sample and Library Preparation User Guide*
- *Oncomine™ Dx Target Test Part III: Template Preparation User Guide*
- *Oncomine™ Dx Target Test Part IV: Sequencing and Results Reports User Guide*
- *Oncomine™ Dx Target Test Part V: Torrent Suite™ Dx Software 5.12.5 Reference User Guide*

All five guides are required to complete the entire Oncomine™ Dx Target Test workflow.

Note: The procedures in these guides supersede the instructions in the *Ion PGM™ Dx System User Guide* when using the Ion PGM™ Dx System with the Oncomine™ Dx Target Test.



Product information

Product description

Oncomine™ Dx Target Test

The Oncomine™ Dx Target Test is an *in vitro* diagnostic next-generation sequencing test to detect somatic alterations in human DNA and RNA isolated from formalin-fixed, paraffin-embedded (FFPE) tissue samples. Detection of these variants is performed using the Ion PGM™ Dx System.

For a complete product description of the Oncomine™ Dx Target Test, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Sequencing components

The Ion PGM™ Dx Sequencing Kit, included as part of the Oncomine™ Dx Target Test Kit, is used to prepare template-positive Ion PGM™ Dx ISPs for 200 base-read sequencing on the Ion 318™ Dx Chip and Ion PGM™ Dx System.

Intended use

For the Intended Use statement for the Oncomine™ Dx Target Test, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Theory of operation

For a complete description of the Theory of Operation of the system, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Software compatibility and requirements

The procedures in this guide are designed for use with Torrent Suite™ Dx Software version 5.12.5 or later. For a complete description of software compatibility and requirements, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Materials provided

Oncomine™ Dx Target Test Kit

The Oncomine™ Dx Target Test Kit (Cat. No. A51695) includes the following subkits.

IMPORTANT! Refer to the product label for the storage conditions and expiration dates of individual modules and components.

✓	Subkit	Part No.
	Oncomine™ Dx Target Test, Controls, and Diluent Kit	A49756
	Ion Torrent Dx FFPE Sample Preparation Kit	A32445
	Ion PGM™ Dx Library Kit	A49758
	Ion OneTouch™ Dx Template Kit	A49759
	Ion PGM™ Dx Sequencing Kit	A49760
	Ion 318™ Dx Chip Kit	A18937
	Oncomine™ Dx Target Test User Guides and Assay Definition File	A51694

Subkits used in this guide

The procedures in this user guide use the following subkits from the Oncomine™ Dx Target Test Kit.

Ion PGM™ Dx Sequencing Kit

The Ion PGM™ Dx Sequencing Kit (Cat. No. A49760) includes the following modules and components.

✓	Component	Amount	Storage
Ion PGM™ Dx Sequencing Supplies (Part No. A18936)			
	SEQ Wash Bottle Sipper	8 long, 16 short	15°C to 30°C
	SEQ Reagent Tube Sipper	32	
	SEQ Reagent Tube plus label	32	
	SEQ Wash 1 Bottle (250-mL bottle)	1	
	SEQ Wash 2 Bottle (2-L bottle)	1	
	SEQ Wash 3 Bottle (250-mL bottle)	1	

✓	Component	Amount	Storage
Ion PGM™ Dx Sequencing Reagents (Part No. A18934)			
	SEQ dGTP (black cap)	2 × 40 µL	–30°C to –10°C
	SEQ dCTP (blue cap)	2 × 40 µL	
	SEQ dATP (green cap)	2 × 40 µL	
	SEQ dTTP (red cap)	2 × 40 µL	
	SEQ Enzyme (yellow cap)	24 µL	
	SEQ Primer (white cap)	96 µL	
Ion PGM™ Dx Sequencing Solutions (Part No. A18935)			
	SEQ W2 Solution (white cap)	8 × 126.25 mL	2°C to 8°C (store SEQ W2 Solution bottles in the sealed plastic bag provided)
	SEQ Cleaning Tablet	8 tablets	
	SEQ Sample Buffer (brown cap)	160 µL	
	SEQ W3 Solution (white cap)	4 × 100 mL	

Ion 318™ Dx Chip Kit

The Ion 318™ Dx Chip Kit (Cat. No. A18937) includes the following components.

✓	Component	Amount	Storage
	Ion 318™ Dx Chip	8	15°C to 30°C

Ion PGM™ Dx Instrument System

The Ion PGM™ Dx Instrument System (Cat. No. A25511) includes the following components, which are also sold separately.

✓	Component	Catalog No.
	Ion OneTouch™ Dx Instrument and accessories	A25483
	Ion OneTouch™ ES Dx Instrument and accessories	A25484
	Ion PGM™ Dx Sequencer and accessories	A25485
	Ion PGM™ Wireless Scanner	A25486
	Ion Torrent™ Server (software installed separately)	A28552

(continued)

✓	Component	Catalog No.
	Ion OneTouch™ Rack Kit <ul style="list-style-type: none"> • Ion OneTouch™ Solutions Rack • Ion OneTouch™ Assembly Rack • Ion OneTouch™ Sample Rack 	A24694
	Ion PGM™ Dx Chip Minifuge: <ul style="list-style-type: none"> • 120 VAC • 230 VAC 	A25058 A25482
	DynaMag™ Dx Kit—Tube & Plate <ul style="list-style-type: none"> • DynaMag™ Dx 96-Well Plate Magnet • DynaMag™ Dx 16 2-mL Magnet 	A31755 A31347 A31346

Materials and equipment required but not provided

Unless otherwise indicated, all materials are available through thermofisher.com. "MLS" indicates that the material is available from fisherscientific.com or another major laboratory supplier.

✓	Description	Source
	Tank of compressed nitrogen (grade 4.8, 99.998% or better)	MLS
	Multistage (dual-stage) gas regulator (0-50 PSI, 2-3 Bar output)	MLS
	18-MΩ water purification system (see the following description)	MLS
	0.45-μm vacuum filtration system and filters (nylon or PVDF filters, 1 L vol.)	MLS
	Pipettes (2-, 20-, 200-, and 1000-μL)	MLS
	Aerosol-barrier pipette tips (10-, 20-, 200-, and 1000- μL)	MLS
	Vortexer with a rubber platform	MLS
	Mini centrifuge	MLS
	Microcentrifuge (must accommodate standard 1.5- and 0.2-mL tubes and generate 15,000 rcf)	MLS
	Veriti™ Dx 96-well Thermal Cycler, 0.2 mL	4452300
	Graduated cylinders (1 L or 2 L volume)	MLS
	Glass bottle (1 L)	MLS
	15-mL conical tubes	MLS
	NaOH, ACS grade (10 M)	MLS

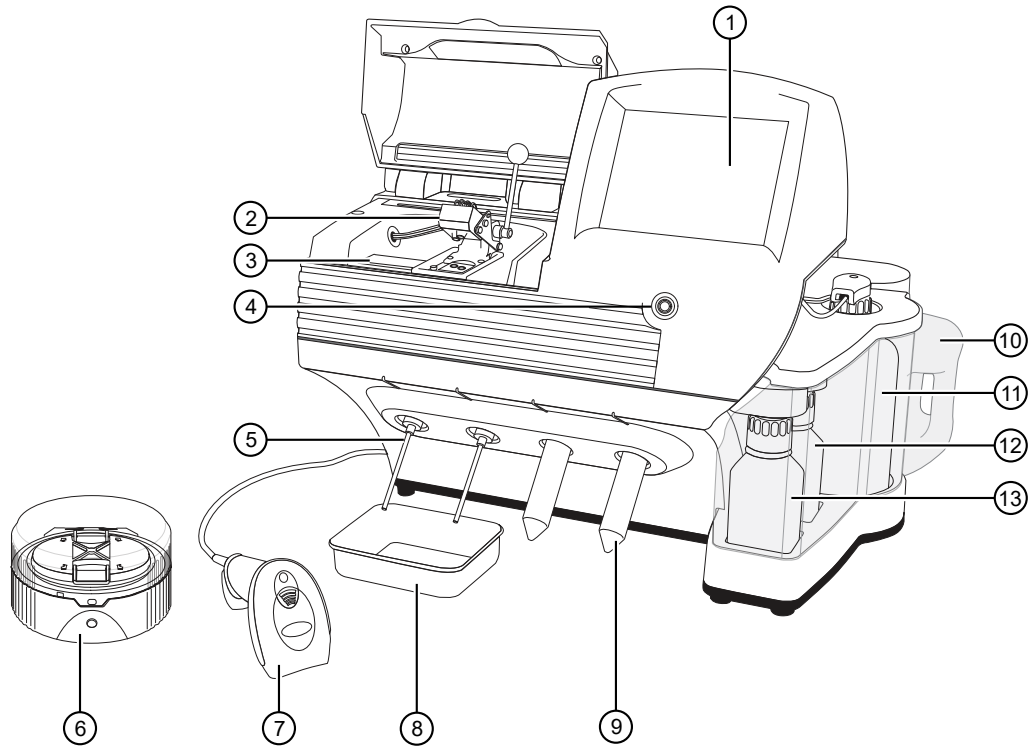
(continued)

✓	Description	Source
	Nuclease-free water	MLS
	Benchtop cold box	MLS
	5-mL and 10-mL serological pipettes, and pipette controller (for preparing the NaOH solution)	MLS

18-M Ω water purification system

The Ion PGM™ Dx Sequencer requires an 18-M Ω water purification system to prepare water for solutions used on the instrument. Such a system is essential to remove ions and organic carbons from the water that might interfere with the chip surface or sequencing enzyme. Water purchased from vendors or stored for any length of time is not acceptable.

Ion PGM™ Dx System with Reagent and Wash Bottles attached



- | | |
|-----------------------------|-------------------------------|
| ① Touchscreen | ⑧ Collection tray |
| ② Chip clamp | ⑨ Reagent Tube |
| ③ Grounding plate | ⑩ Waste Bottle |
| ④ Power button | ⑪ Wash 2 Bottle (W2 position) |
| ⑤ Reagent Tube Sipper | ⑫ Wash 3 Bottle (W3 position) |
| ⑥ Ion PGM™ Dx Chip Minifuge | ⑬ Wash 1 Bottle (W1 position) |
| ⑦ Barcode scanner | |

2

Before you begin

Installation and special requirements

Special requirements

- The Torrent Suite™ Dx Software requires the use of Google™ Chrome™ browser.
- The Ion PGM™ Dx Sequencer requires the use of an 18-MΩ water purification system installed in the laboratory. Such a system is essential for ensuring that any ions or organic carbons that might interfere with the chip surface or sequencing enzyme are removed from the water immediately before use on the instrument. Water purchased from vendors or stored for any length of time is not acceptable.

Gas cylinders

You must supply the required nitrogen gas cylinder and accessories for the installation. This instrument requires a pressurized house line or one size 1-A nitrogen gas cylinder that holds approximately 7.2 m³ (257 ft³) of gas when full. Use only prepurified nitrogen of 99.998% (grade 4.8) or greater purity.



CAUTION! Damage to the instrument and its products can result from using impure gas, gases other than nitrogen, or an inadequate amount of gas.



WARNING! EXPLOSION HAZARD. Pressurized gas cylinders are potentially explosive. Always cap the gas cylinder when it is not in use, and attach it firmly to the wall or gas cylinder cart with approved brackets or chains.



WARNING! Gas cylinders are heavy and may topple over, potentially causing personal injury and tank damage. Cylinders should be firmly secured to a wall or work surface. Please contact your Environmental Health and Safety Coordinator for guidance on the proper installation of a gas cylinder.

Perform a leak test

To perform a leak test on the gas cylinder:

1. Open the main tank shutoff valve. The high-pressure gauge of the gas tank regulator reads approximately 2,000–2,500 psi for a full tank.
2. Adjust the pressure to the instrument by slowly turning the pressure adjustment valve clockwise until the low-pressure gauge reads 30 psi.
3. Close the needle valve, then close the main tank valve.
4. Monitor the high-pressure gauge of the gas tank regulator for 5 minutes. There should be no noticeable drop in pressure.

If the pressure	Action
Drops in 5 minutes	There can be a leak at either the needle valve or the gas tank regulator itself. Check the fittings and resolve any problems, then continue with step 5.
Does not drop in 5 minutes	The instrument passes the leak test. Reopen the main tank valve and skip the following steps.

5. Open the main tank valve and the needle valve for at least 15 seconds to pressurize the instrument.
6. Close the main tank valve.
7. Monitor the high-pressure gas tank regulator gauge. There should be no more than a 100-psi drop in pressure after 5 minutes. Locate, then resolve any leaks. Turn the main tank valve back on.

Electrical and environmental conditions

IMPORTANT! Protection provided by the equipment may be impaired if the instrument is operated outside the environment and use specifications, the user provides inadequate maintenance, or the equipment is used in a manner not specified by the manufacturer.

IMPORTANT! Observe current Good Clinical Laboratory Practices (GCLP) when using this instrument.

See the *Ion PGM™ Dx System Site Preparation Guide* (Pub. No. MAN0024945) for information about instrument location and setup. Instruments should be shielded from excess exposure to dust, vibration, strong magnetic fields, drafts, excessive moisture, or large temperature fluctuations. Surge protectors or line conditioners should be used if the voltage source is not stable. Sudden voltage spikes can cause damage to the electronics inside the instruments.

Ensure that the room where the instruments have been installed is maintained under correct environmental conditions. Avoid placing the instruments adjacent to heaters, cooling ducts, or in direct sunlight. Place the instruments at least a meter away from major sources of electronic noise, such as refrigerators or microwaves.

Electrical requirements



CAUTION! Do not unpack or plug in any components until a field service representative has configured them for the proper operating voltage.



WARNING! For safety, the power outlet used for powering the instrument must be accessible at all times. In case of emergency, you must be able to immediately disconnect the main power supply to all the equipment. Allow adequate space between the wall and the equipment so that the power cords can be disconnected in case of emergency.

- Electric receptacle required: 2-prong with ground pin
- Main AC line voltage tolerances must be at most $\pm 10\%$ percent of nominal voltage.
- Power cords are provided with the instruments. If not suitable for installation in your region, ensure any power cord you do use is:
 - Maximum 10 feet (3 meters) in length
 - Grounding type
 - Compatible with the power supply receptacles used to connect to main power
 - Suitable for the rating of the instrument and main power supply
 - Compliant with local safety requirements (for example, UL Listed for North America, JIS approved for Japan, HAR or agency certified for Europe)
- (Ion OneTouch™ Dx Instrument only) Fuse Rating: 6 A, 250 VAC, Type M. Replace only with the same fuse type and rating.



WARNING! FIRE HAZARD. For continued protection against the risk of fire, replace fuses only with fuses of the type and rating specified for the instrument.

Device	Rated voltage ^[1,2]	Rated frequency	Rated current ^[3]
Ion PGM™ Dx Sequencer	110/120VAC 220/240VAC	50/60 Hz	9 A
Ion Torrent™ Server ^[4]	110/120VAC 220/240VAC	50/60 Hz	11 A
Ion OneTouch™ Dx Instrument with power supply	110/120VAC 220/240VAC	50/60 Hz	5.5 A
Ion OneTouch™ ES Dx Instrument	110/120VAC 220/240VAC	50/60 Hz	375 mA 160 mA
Ion PGM™ Dx Chip Minifuge	120 VAC 220–240 VAC	50/60 Hz	130 mA 65 mA

^[1] In Japan, rated voltages of 100 VAC and 200 VAC are acceptable.

^[2] If the supplied power fluctuates beyond the rated voltage, a power line regulator may be required. High or low voltages can adversely affect the electronic components of the instrument.

^[3] Based on rated current at minimum input voltage.

^[4] Minimum Efficiency: 65% (Energy Star Qualified); 85% Efficient Power Supply.

Environmental requirements

Ensure that the room where the instruments have been installed is maintained under the correct environmental conditions. Avoid placing the instruments next to heaters, cooling ducts, or in direct sunlight. Place the sequencer at least a meter away from major sources of electronic noise, such as refrigerators or microwaves.



CAUTION! Use of the instruments in an unspecified manner may result in the protection provided by the instruments to be impaired.

Component	Acceptable range
Altitude	Between sea level and 2,000 meters (6,500 feet) above sea level
Humidity: operating	10–90% relative humidity
Humidity: transportation and storage	20–80% relative humidity
Temperature: operating	15–30°C (59–86°F) At or above 1,800 meters (5,906 feet), the system must not be used if the temperature is above 29.5°C.
Temperature: transportation and storage	–30°C to 60°C (–22°F to 140°F)

(continued)

Component	Acceptable range
Vibration	Ensure that benches where instruments are to be installed are free of vibration and have no contact with equipment that causes vibration (freezers, pumps, and similar equipment). Vibration can reduce the quality of sequencing measurements.
Pollution	The system has a Pollution Degree rating of II (2). The system may only be installed in an environment that has nonconductive pollutants, such as dust particles or wood chips. Typical environments with a Pollution Degree II (2) rating are laboratories, sales, and commercial areas.
Overvoltage category	The instruments have an installation (overvoltage) category of II (2).
Other conditions	For indoor use only. Keep away from any vents that could expel particulate material on the system components.

Precautions before using the Ion PGM™ Dx System

For additional safety information, see Appendix C, “Safety”.

Instrument installation by trained personnel only

IMPORTANT! The Ion PGM™ Dx System is installed by trained service personnel and must not be relocated without assistance from trained service personnel. See “Customer and technical support” on page 106.

Nucleic acid contamination

IMPORTANT! A primary source of contamination is DNA fragments from previously processed samples. Do not introduce amplified DNA into the library preparation laboratory or work area.

IMPORTANT! Possible contamination can occur during the transfer of dNTPs into Reagent Tubes. Be careful to avoid cross contamination of dNTP stocks. Barrier tips are required for all pipetting steps. Change gloves after handling concentrated dNTP stocks.

Reagent contamination

Before use, verify that any nuclease-free water used in the procedure is not cloudy, a potential indication of contamination. If the water is cloudy, use a different vial.

CO₂ contamination

IMPORTANT! Dry ice (solid CO₂) must be kept away from areas where buffers, wash solutions, or sources of molecular biology grade water for the Ion PGM™ Dx System are used. High air concentrations of subliming CO₂ may change the pH of such buffers during or after their preparation. The stability of the pH of these buffers is a critical factor in the performance of the Ion PGM™ Dx System.

Instrument vibration and clearances

IMPORTANT! Significant vibration during sequencing may add noise and reduce the quality of the measurements. The Ion PGM™ Dx Sequencer must be installed on a bench that is free from vibrations or in contact with equipment that can cause vibrations to the bench, such as freezers, pumps, large benchtop centrifuges, and other similar equipment. Mini and microcentrifuges may be used near the sequencer. An air table is not required, nor is securing the sequencer to the bench.

IMPORTANT! Position the Ion PGM™ Dx Sequencer so that the front bezel is a minimum of 12 in. (30.5 cm) and the Reagent Tubes containing dNTPs are a minimum of 8 in. (20.3 cm) from the front of the laboratory bench. Place the instrument at least 40 in. (1 meter) away from major sources of electronic noise such as refrigerators or microwaves.

Static electricity

IMPORTANT! To avoid possible damage to chips from static electricity, see “Guidelines for chip handling and use” on page 20.

Ventilation requirements



WARNING! Instrumentation must be installed and operated in a well-ventilated environment, defined as having a minimum airflow of 6–10 air changes per hour. Assess the need for ventilation or atmospheric monitoring to avoid asphyxiation accidents from inert gases and/or oxygen depletion, and take measures to clearly identify potentially hazardous areas through training or signage. Please contact your Environmental Health and Safety Coordinator to confirm that the instruments will be installed and operated in an environment with sufficient ventilation.

Procedural guidelines

Definitions

Throughout this guide:

- Room temperature is defined as the temperature range 15–30°C (59–86°F).
- A pulse centrifugation consists of a 3–5 second centrifugation at maximum speed in a mini centrifuge.

Guidelines to prevent cross-contamination

- Use good laboratory practices to minimize cross-contamination of products and reagents.
- When designing the laboratory layout, consider separating pre- and post-amplification activities. To significantly reduce the potential for contamination, dedicate laboratory supplies and/or equipment to the appropriate space.

Guidelines for pipetting

- Use aerosol-barrier pipette tips. Change pipette tips between samples.
- Avoid introducing air bubbles when pipetting by keeping the pipette tip at the bottom of the solution in the wells.
- Set the pipette to the recommended volume for mixing, and insert tip into the solution with the pipette plunger depressed to avoid introducing air bubbles.
- Visually inspect multi-channel pipette tips to ensure volumes are equivalent during pipeting.
- Touch tip to the side of well and slowly pipet reagent on the side of the well to form a droplet. This enables small volumes to be pipetted accurately and to ensure that the reagent has been added to the well.
- Inspect the pipette tips to verify that the reagent has been adequately dispensed.

Guidelines for chip handling and use

IMPORTANT! To avoid damage to Ion 318™ Dx Chips or the Ion PGM™ Dx Sequencer due to electrostatic discharge:

- **Remove your gloves when handling chips**, especially before transferring chips on or off the instrument. Follow the steps in the sequencing procedure for taking off and putting on gloves.
 - **Hold chips by their edges when handling.**
 - **Do not place chips directly on the bench or any other surface.** Always place chips either on the grounding plate on the Ion PGM™ Dx Sequencer or in the Ion PGM™ Dx Chip Minifuge bucket.
-

Note: Ion 318™ Dx Chips can be handled without gloves during all stages of chip preparation, loading, and sequencing without risk of contamination.

Used chips cannot be reused for sequencing. Used chips must be discarded or clearly marked for cleaning and initialization.

Guidelines for initializing the sequencer

- Handle the SEQ dGTP, SEQ dCTP, SEQ dATP, and SEQ dTTP tubes carefully to avoid cross-contamination. Always change gloves after removing used sippers from the Ion PGM™ Dx System to avoid cross-contamination of the nucleotides. Also change gloves after handling concentrated dNTP stocks.
- Replace the SEQ Reagent Tubes and sippers every time you initialize.
- After 8 sequencing runs, do not use the SEQ Wash 1 Bottle, SEQ Wash 2 Bottle, or SEQ Wash 3 Bottle for initialization or sequencing to avoid possible breakage or leaking. You can continue to use the SEQ Wash 1 Bottle and SEQ Wash 3 Bottle as extra cleaning bottles.

Guidelines for sequencing runs

- One or two sequencing runs may be performed from a single initialization based on the option selected when starting the initialization, but both runs must be started within 27 hours after the start of initialization.
- If you press the **Abort** button on the sequencer touchscreen, the touchscreen may freeze. You may need to restart the sequencer.

IMPORTANT! After aborting a run, do not open the chip clamp, reagent tubes, or wash bottles until a new run or cleaning is initiated. Doing so can cause a fluid or gas leak if the sequencer was in a pressurized state when the run was aborted. From the main menu, select either **Clean** or **Run**, then follow the touchscreen prompts to depressurize the system.

Reagent management

Follow the guidelines below for proper reagent storage and use.

Storage

Reagents must be stored under appropriate conditions. Refer to the Product Information section in each user guide for the storage conditions of the kit components used in the procedures in that guide. The Oncomine™ Dx Target Test Kit includes kits with multiple component boxes that require different storage conditions. For example, the Oncomine™ Dx Target Test, Controls, and Diluent Kit includes four boxes, which are stored at different temperatures. To use the Oncomine™ Dx Target Test, Controls, and Diluent Kit, retrieve all boxes from their different storage areas and confirm that they are from the same master lot.

Kit interchangeability and component box lot matching

The top-level kits used for sample preparation, library preparation, template preparation, and sequencing can be mixed and matched. For example, an Ion PGM™ Dx Library Kit can be used with any Ion PGM™ Dx Sequencing Kit. However, the component boxes in a particular kit must be lot-matched with the other boxes in that kit.

Note: The controls and diluent in the Oncomine™ Dx Target Test, Controls, and Diluent Kit must be lot-matched. However, the controls do not require lot-matching with the Oncomine™ Dx Target Test DNA and RNA Panel.

Each component box lists the 4-digit lot prefixes of the compatible component boxes inside the box lid. Before using a particular kit, check the inside lid of each box to confirm that it is compatible with the other boxes.

An example inside box lid label is shown below:

ATTENTION: When setting up a procedure, confirm that the component boxes listed below have 4-digit lot prefixes that match.

REF	Product Name	LOT
A18934	Ion PGM™ Dx Sequencing Reagents	MMYY
A18935	Ion PGM™ Dx Sequencing Solutions	MMYY
A18936	Ion PGM™ Dx Sequencing Supplies	MMYY

An example box label with lot information is shown below:

iontorrent
by Thermo Fisher Scientific

REF A18934
LOT MMYX XXXXXX

Ion PGM™ Dx Sequencing Reagents

YYYY-MM-DD

8

Product of USA

CE IVD

Life Technologies Corporation
7335 Executive Way
Frederick MD 21704
www.thermofisher.com

Life Technologies Europe B.V.
Kwartsweg 2
2665 NN Bleiswijk
The Netherlands

(01)10190302006224
(17)000000
(10)MMYX_XXXXXXX
(240)A18934

1

1 Lot prefix

Instrument operation and maintenance

Service and maintenance

You will be alerted by the Ion PGM™ Dx System when annual maintenance service is required. A notification will appear on the instrument touchscreen and in the Torrent Suite™ Dx Software.

Shut down/restart the sequencer

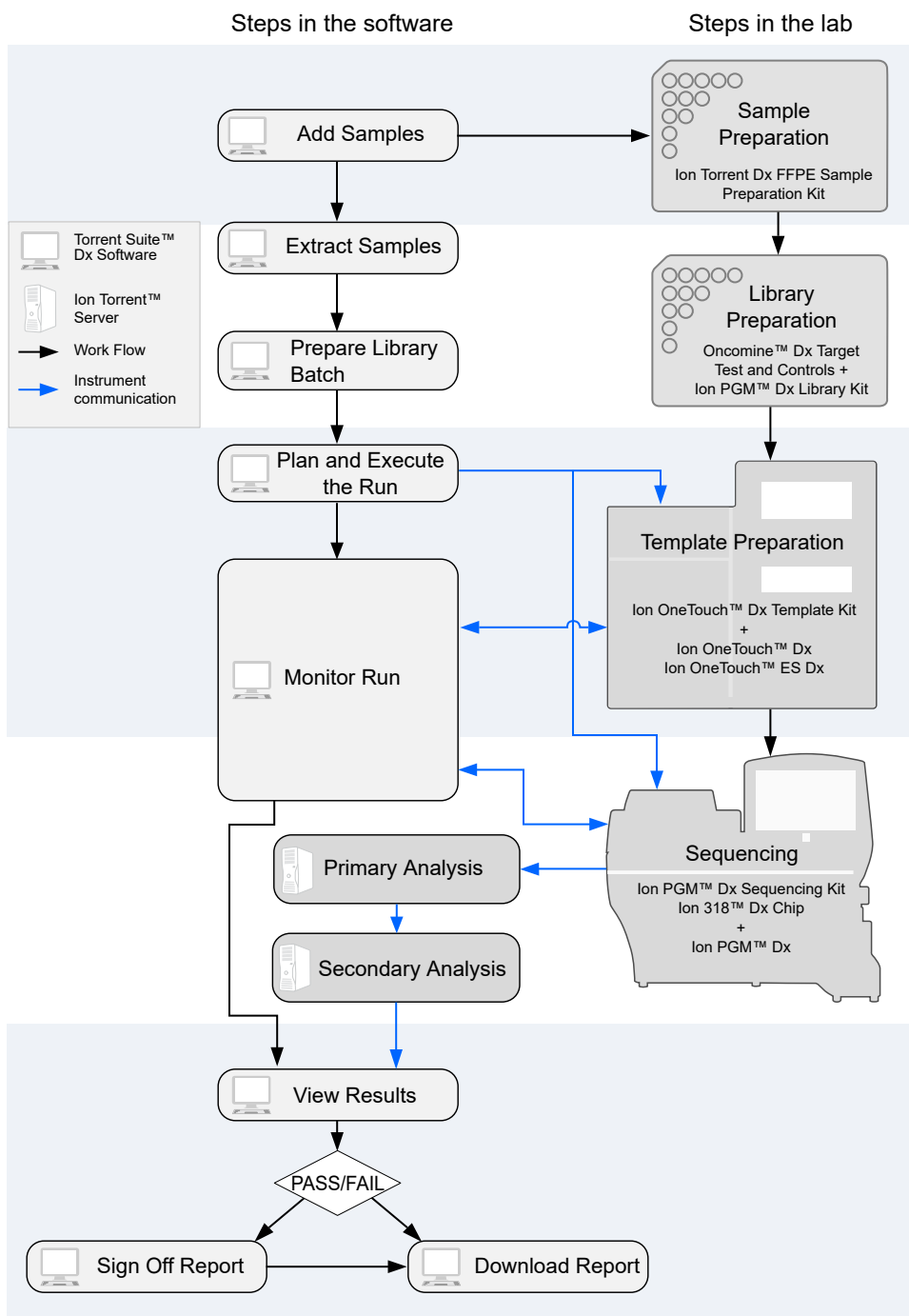
In general, the Ion PGM™ Dx Sequencer can remain on all the time, including overnight and over weekends. If shutdown is necessary, there are two methods, depending on the state of the instrument:

Type of shutdown	Typically used when	Method
Routine shutdown (Managers and Administrators only)	<ul style="list-style-type: none">The instrument will not be used for an extended period of time (>1-2 weeks).The instrument needs to be moved or serviced.	<ol style="list-style-type: none">Press the Options button on the main touchscreen and press Shut Down.If the instrument will not be used for an extended period of time, select the Cleaning checkbox to perform a water clean first, then press Shut Down. Otherwise, just press Shut Down.In the next screen, press the Halt button and then press OK to power down the instrument.
Forced shutdown	The instrument isn't responding normally (e.g., the touchscreen is frozen).	Hold down the power button below the touchscreen for ~5 seconds to power down the instrument.

To restart the sequencer:

- Hold down the power button below the touchscreen for ~5 seconds.

Oncomine™ Dx Target Test system diagram



Sequencing workflow

Note: Up to two sequencing runs may be performed from one initialization, if both runs are started within 27 hours after start of initialization.

Previous guide: *Oncomine™ Dx Target Test Part III: Template Preparation User Guide*



“Clean the Ion PGM™ System” on page 26



“Initialize the sequencer” on page 34



“Prepare enriched ISPs for sequencing” on page 41



“Set up the sequencing run” on page 43



“Load the sample on the chip” on page 46



“Perform the run” on page 49

Proceed to: *Oncomine™ Dx Target Test Part V: Torrent Suite™ Dx Software 5.12.5 Reference User Guide*



Clean and initialize

Clean the Ion PGM™ System

Cleaning schedule

The Ion PGM™ Dx Sequencer requires cleaning with either 18 MΩ water or chlorite solution according to the following schedule.

Clean with:	Schedule:
18 MΩ water	<p>The Ion PGM™ Dx Sequencer requires cleaning with 18 MΩ water when one of the following conditions is met:</p> <ul style="list-style-type: none">• The last water cleaning was completed more than 27 hours ago, and initialization was performed• If during the last initialization, the One run per initialization checkbox was selected, and one run was performed since the last water cleaning• If during the last initialization, the Two runs per initialization checkbox was selected, and two runs were performed since the last water cleaning
Chlorite solution	<p>The Ion PGM™ Dx Sequencer requires cleaning with chlorite solution when one of the following conditions is met:</p> <ul style="list-style-type: none">• The last chlorite cleaning was completed more than 7 days ago, and one or more runs have been performed since that cleaning• The instrument has been left with reagent for more than 48 hours <p>Note: If the Ion PGM™ Dx Sequencer will not be used for more than 3 days, a chlorite cleaning is strongly recommended within 48 hours after run completion.</p>

Before you begin

Mark the cleaning bottles

Three 250-mL bottles are provided as part of instrument installation. Mark these bottles as described below and use them in the cleaning procedures.

Note: After you have used the 250-mL bottles provided in each sequencing kit for 8 runs, you can mark them as extra cleaning bottles.

- Write "Waste" on one 250-mL bottle, to be used for both water and chlorite washes.
- Write "Water" on one 250-mL bottle, to be used for water washes only.
- Write "Chlorite" on one 250-mL bottle, to be used for chlorite solution only.

Mark the cleaning chips

After a sequencing run, used chips can be marked for use in cleaning and initialization. (Chips cannot be reused for sequencing.) Mark used chips as described below.

- To designate a used chip for chlorite washes only, write "CL" on the corner. Do not use this chip for initialization.
- To designate a used chip for water washes and initialization, write "W" on the corner. Do not use this chip for chlorite washes.

Chip marked for chlorite wash only:



Chip marked for water wash/initialization:



Start the cleaning

The sequencer touchscreen will guide you through each step in the cleaning process. The touchscreen provides a brief summary of the steps, which are described below in more detail.

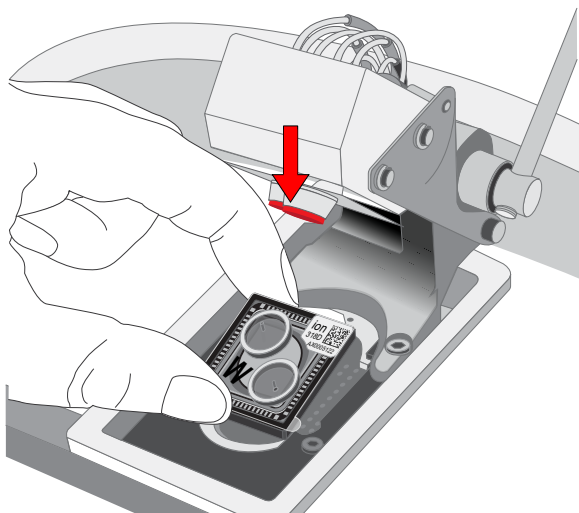
1. Log into the Ion PGM™ Dx Sequencer touchscreen.
2. On the main menu, press the **Clean** button.
The instrument touchscreen will display the following:
 - **Last water clean: [YYYY-MM-DD] [hh]:[mm]:[ss]**
 - **Last chlorite clean: [YYYY-MM-DD] [hh]:[mm]:[ss]**
3. The type of cleaning required is selected by default (**18 MΩ water cleaning** or **Chlorite cleaning**). Press **Next**.

18-MΩ water cleaning

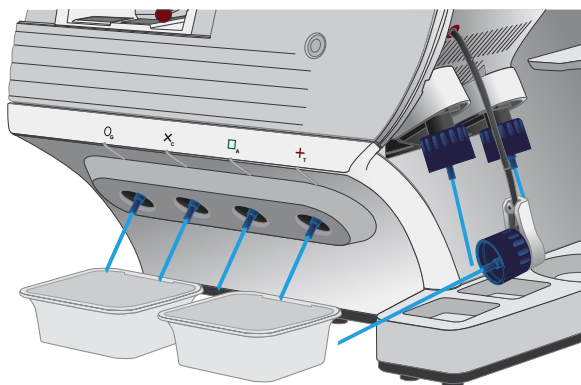
IMPORTANT! For all the following steps, use 18-MΩ water directly from the purification system. Do not use water that has been collected or stored in any other containers.

1. Touch the grounding plate on the Ion PGM™ Dx Sequencer with a bare finger, then use ungloved hands to insert the used chip marked "W" into the chip clamp.

IMPORTANT! Confirm that both red rubber port gaskets are securely in place in the clamp. Loose or missing gaskets can result in a spill hazard and instrument damage.



2. Close the clamp, then press **Next** on the touchscreen.
3. Put on new gloves, then remove all wash bottles and reagent tubes attached to the instrument. Keep the sippers in place at all positions. Press **Next**.



4. Rinse the bottle marked "Water" twice with ~100 mL of 18-M Ω water. Press **Next**.
5. Fill the bottle marked "Water" with 250 mL of 18-M Ω water. Press **Next**.
6. Remove and rinse the W1 sipper with 18-M Ω water, then reattach the sipper. Press **Next**.
7. Attach the bottle marked "Water" to the W1 position, ensuring that the W1 cap is screwed on tightly. Press **Next**.

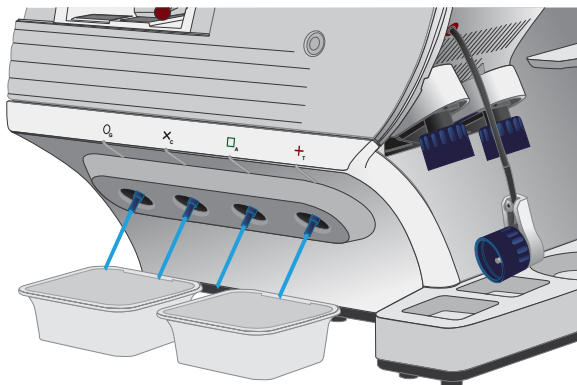
8. With the W3 sipper still in its cap, insert the sipper into the empty bottle marked "Waste". Do not screw on the cap. Insert the W2 sipper into the same bottle. Press **Next**.



9. Place collection trays below the reagent tube sippers in the dNTP positions.

Note: To avoid spills, ensure that the collection trays are properly aligned to catch fluid flowing out from the sippers.

10. Press **Next** to begin cleaning.
The cleaning will take ~15 minutes to complete.
11. When cleaning is complete, leave the reagent sippers and collection trays in place. Remove the bottles and sippers from the W1, W2, and W3 positions. Press **Next** to return to the main menu, then proceed to initialization.



Chlorite cleaning

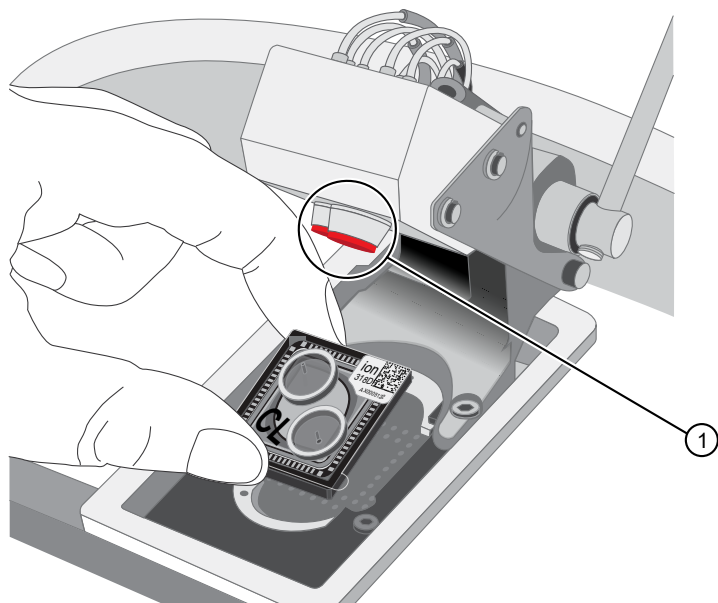
Note: This procedure uses 1 M NaOH, diluted from 10 M NaOH each week using nuclease-free water. You can use the 1 M NaOH that was prepared during template preparation.

1. Fill a glass bottle with 1 L of 18-M Ω water, then add a SEQ Cleaning Tablet (chlorite tablet). Allow the tablet to completely dissolve (~10 minutes). Press **Next** on the touchscreen.
2. Add 1 mL of 1 M NaOH and filter the solution using a 0.22- μ m or 0.45- μ m filter. Press **Next**.

Note: Use the chlorite solution within 2–3 hours. Discard the unused solution in an appropriate waste container after this time.

3. Touch the grounding plate on the Ion PGM™ Dx Sequencer with a bare finger, then use ungloved hands to insert the used chip marked "CL" into the chip clamp.

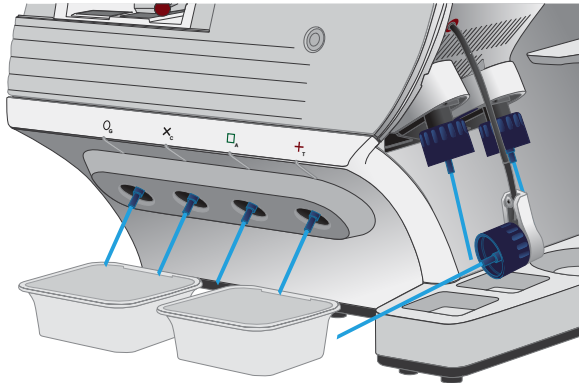
IMPORTANT! Confirm that both red rubber port gaskets are securely in place in the clamp. Loose or missing gaskets can result in a spill hazard and instrument damage.



① Rubber port gaskets

4. Close the clamp, then press **Next**.

5. Put on new gloves, then remove all wash bottles and reagent tubes that are attached to the instrument. Keep the sippers in place at all positions. Press **Next**.



6. Rinse the bottle marked "Chlorite" and the bottle marked "Water" twice each with ~150 mL of 18-M Ω water.
7. Add 250 mL of filtered chlorite solution to the bottle marked "Chlorite". Press **Next**.
8. Remove and rinse the sipper in the W1 position with 18-M Ω water, then reattach it to the instrument. Press **Next**.
9. Attach the bottle marked "Chlorite", containing the filtered chlorite solution, to the W1 position. Make sure that the W1 cap is tight. Press **Next**.
10. With the W3 sipper still in its cap, insert the tube into the empty bottle marked "Waste". Do not screw on the cap. Insert the W2 sipper into the same bottle. Press **Next**.



11. Place collection trays below the reagent tube sippers in the dNTP positions.

Note: To avoid spills, ensure that the collection trays are properly aligned to catch fluid flowing out from the sippers.

12. Press **Next** to begin cleaning. The cleaning will take ~13 minutes to complete.
13. When prompted, remove the bottle marked "Chlorite" from the W1 position. Press **Next**.
14. Remove and rinse the W1 sipper with 18-MΩ water, then reattach the sipper. Press **Next**.
15. Fill the bottle marked "Water" with 250 mL of 18-MΩ water and attach the bottle in the W1 position. Make sure the cap is tight.



16. Press **Next**. The water rinse will take ~15 minutes to complete.
17. When cleaning is complete, leave the reagent sippers and collection trays in place. Remove the bottles and sippers from the W1, W2, and W3 positions. Press **Next** to return to the main menu.
18. Rinse the "Chlorite" and "Water" bottles twice each with ~150 mL of 18-MΩ water, then proceed to initialization.

Initialization

Initialization guidelines

- When performing one or two sequencing runs from the same initialization, the run(s) must be started within 27 hours after the start of initialization.
- Handle the SEQ dGTP, SEQ dCTP, SEQ dATP, and SEQ dTTP tubes carefully to avoid cross-contamination. Always change gloves after removing used sippers from the Ion PGM™ Dx System to avoid cross-contamination of the nucleotides. Also change gloves after handling concentrated dNTP stocks.
- Replace the SEQ Reagent Tubes and sippers every time you initialize.
- After eight sequencing runs, do not use the SEQ Wash 1 Bottle, SEQ Wash 2 Bottle, or SEQ Wash 3 Bottle for initialization or sequencing to avoid possible breakage or leaking. You can continue to use the SEQ Wash 1 Bottle and SEQ Wash 3 Bottle as extra cleaning bottles, as described in “Mark the cleaning bottles” on page 26.

Ion PGM™ Dx Sequencing Kit component lot matching

The three components of the Ion PGM™ Dx Sequencing Kit must be lot-matched with each other for use.

Component	Part No.	Storage
Ion PGM™ Dx Sequencing Supplies	A18936	15°C to 30°C
Ion PGM™ Dx Sequencing Reagents	A18934	-30°C to -10°C
Ion PGM™ Dx Sequencing Solutions	A18935	2°C to 8°C

Before initialization

Check the tank pressure for the nitrogen gas. When the tank pressure drops below 500 psi, change the tank (see also “Gas cylinders” on page 14).

Initialize the sequencer

The instrument touchscreen provides a summary of the steps in the initialization process, which are described below in more detail.

IMPORTANT! Do not remove the old sippers from the dNTP ports until instructed to do so. **Do not let the new sippers touch any surfaces.**

IMPORTANT! Load the bottles as quickly as possible to prevent atmospheric CO₂ from reducing the pH of the Wash 2 solution.

1. After cleaning is complete, the **Initialize** button on the touchscreen main menu will turn blue. Press the button to start the initialization process.
2. Depending on the number of sequencing runs per initialization, select either the **One 200 bp run per initialization** or the **Two 200 bp runs or One 400 bp run per initialization** checkbox. Press **Next**.

Note: If you are performing two runs, both must start within 27 hours after start of the initialization.

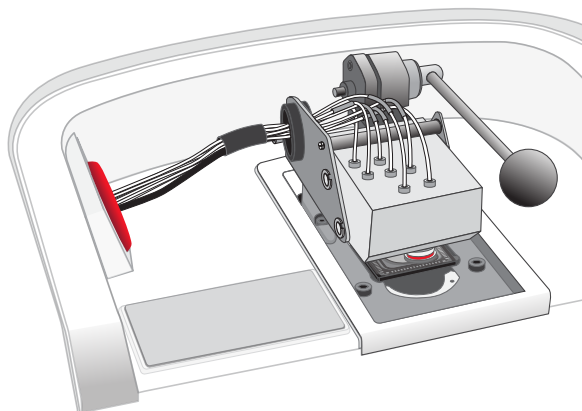
Note: 400-bp runs are performed for Assay Development purposes only and are not used with the Oncomine™ Dx Target Test.

3. Press the **Keyboard** button below the **Sequencing kit barcode** field. Using the barcode scanner attached to the sequencer, scan the barcode on the Ion PGM™ Dx Sequencing Reagents box. Press **OK**, then press **Next**.



4. Ensure that the used chip marked "W" (from the cleaning procedure) is secured in the chip clamp on the sequencer.

IMPORTANT! Do *not* use the chip marked "CL" for initialization. (Use it for chlorite cleaning only.)



5. Ensure that the old reagent sippers and collection trays are in place. Press **Next**.
6. Rinse the SEQ Wash 1 Bottle, SEQ Wash 2 Bottle, and SEQ Wash 3 Bottle 3 times each with 18-M Ω water. Use 150 mL for the Wash 1 and Wash 3 bottles, and 500 mL for the Wash 2 bottle.

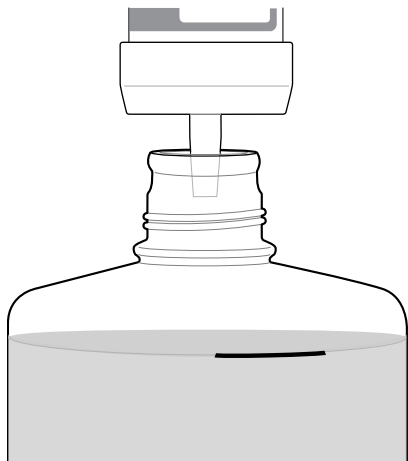
Note: The bottles can be rinsed and reused for up to 8 sequencing runs. To track the number of uses, mark each bottle each time a run has finished as shown below. After 8 uses, the SEQ Wash 1 Bottle and SEQ Wash 3 Bottle bottles can be marked and used for cleaning only.



7. Prepare 500 μ L of fresh 100 mM NaOH daily by diluting 50 μ L of 1 M NaOH in 450 μ L of nuclease-free water.

Note: Prepare a stock of fresh 1 M NaOH weekly.

8. If your 18-M Ω water system has a spigot, extend it into **but not below** the neck of the SEQ Wash 2 Bottle. Otherwise, position the nozzle as close to the mouth of the bottle as possible.



IMPORTANT! If your water system has a digital display, verify that it reads "18 M Ω " when filling the bottle. If it does not, see Appendix A, "Troubleshooting".

9. Fill the bottle to the mold line with 18-M Ω water.

Note: The total volume of water is ~2 liters. You can mark the mold line on the bottle for clarity. If you are preparing bottles for multiple sequencers, cap each bottle immediately after filling. Leave them capped until you are ready to add SEQ W2 Solution.

10. Add an entire bottle of SEQ W2 Solution to the SEQ Wash 2 Bottle.

Note: Store the remaining SEQ W2 Solution bottles in the sealed mylar bag they were provided in.

11. Using a P200 pipette, add 70 μ L of 100 mM NaOH to the SEQ Wash 2 Bottle.

Note: If the initialization consistently undershoots the pH target (see step 21), add more than the recommended 70 μ L of 100 mM NaOH when preparing the Wash 2 Bottle. After adding the NaOH, the Wash 2 Bottle must be between pH 6.0–6.5 at first pH iteration before you start initialization.

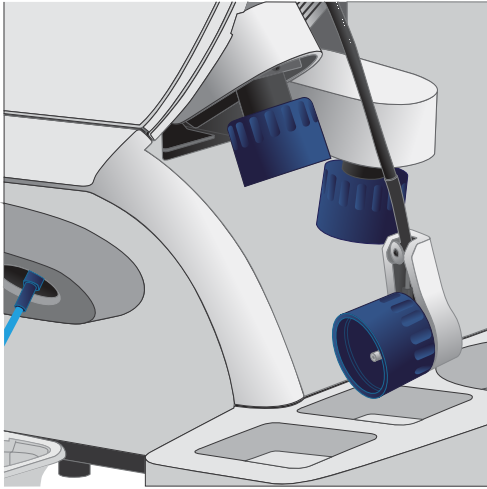
12. Cap the bottle and invert 5 times to mix, then immediately proceed through the remainder of the initialization procedure.

IMPORTANT! Do not store the mixed SEQ Wash 2 Bottle.

13. SEQ Wash 1 Bottle: Add 350 μ L of freshly prepared 100 mM NaOH to a clean SEQ Wash 1 Bottle. Ensure that the pipette tip touches the bottom of the bottle to dispense.

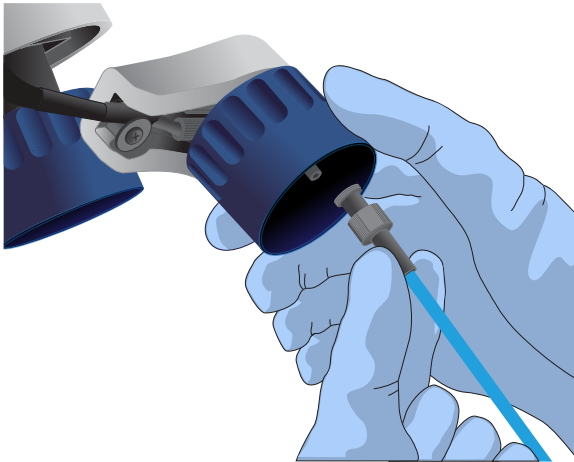
14. SEQ Wash 3 Bottle: Add SEQ W3 Solution to the 50-mL line marked on a clean SEQ Wash 3 Bottle. Press **Next**.

15. Remove the old sippers from the W1, W2, and W3 positions on the instrument, then discard in a waste container.

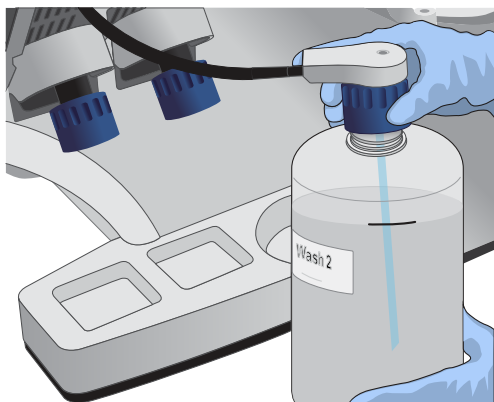


16. Wearing clean gloves, firmly attach a new, long sipper to the cap in the W2 position. **Do not let the sipper touch any surfaces.**

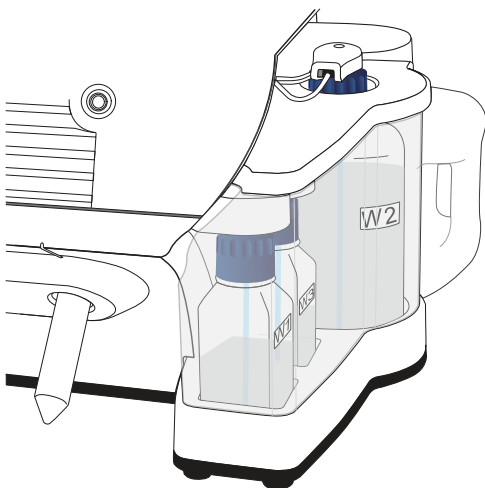
IMPORTANT! Loosely attached sippers can adversely affect results.



17. Immediately attach the prepared SEQ Wash 2 Bottle to the cap in the W2 position, then tighten the cap. Press **Next**.



18. Change gloves and firmly install new, short sippers to the caps in the W1 and W3 positions.
19. Immediately attach the prepared SEQ Wash 3 Bottle to the cap in the W3 position, then attach the SEQ Wash 1 Bottle to the cap in the W1 position. Tighten the caps.



20. Ensure that the collection trays are properly aligned to catch fluid flowing out of the sippers. Press **Next** to start the auto-pH process.
21. When auto-pH is complete:
 - If auto-pH fails (for example, the instrument overshoots or undershoots the target pH), the touchscreen displays an error message. Do not press **Next**. See Appendix A, “Troubleshooting” for information about specific errors.
 - If auto-PH passes, the touchscreen will display to the next step.

22. Thaw the following at room temperature until no ice crystals are visible in the tubes (~15 minutes). Keep the tubes on ice or in a chilled benchtop cold box until ready to use.
- SEQ Primer (white cap)
 - SEQ dGTP (black cap)
 - SEQ dCTP (blue cap)
 - SEQ dATP (green cap)
 - SEQ dTTP (red cap)

23. The kit includes labels for each type of dNTP. Attach one of each label type to four SEQ Reagent Tubes.

24. Verify that no ice crystals are visible in each thawed dNTP stock solution tube (SEQ dGTP, SEQ dCTP, etc.) from step 22. Vortex each dNTP for ~5 seconds, then pulse centrifuge to collect.

Note: To avoid cross-contamination in the next step, open only one dNTP tube at a time, and use a fresh pipette tip for each aliquot.

25. Using a P20 pipette and separate tips, aliquot each dNTP into its appropriately labeled reagent tube as follows:

- Aliquot 10 μ L of dNTP per reagent tube if performing one 200 bp run per initialization.
- Aliquot 20 μ L of dNTP per reagent tube if performing two 200 bp runs per initialization.

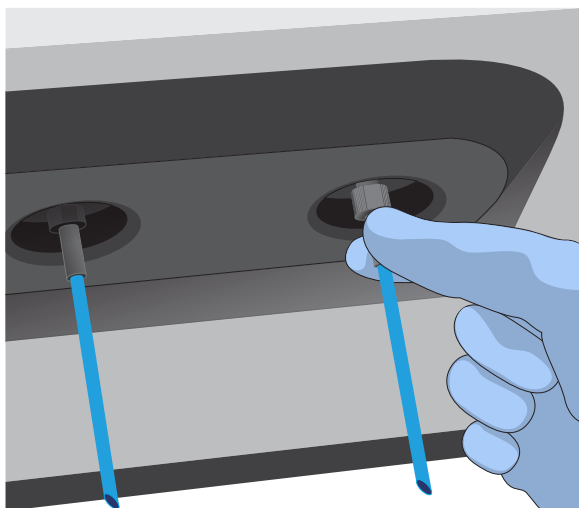
Note: Reagent tubes can be capped and left at 2–8°C or on ice until ready for use. Place the remaining dNTP stocks back into –30°C to –10°C for storage. The dNTPs can be freeze-thawed up to 4 times. We recommend marking the box with each freeze-thaw to indicate how many have occurred.

26. Press **Next**. Remove the old reagent tube sippers, then discard them in a waste container.

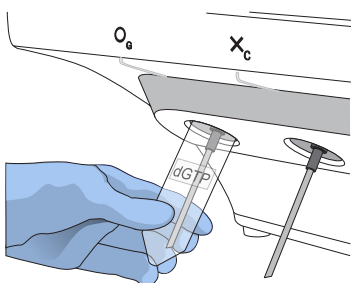
27. Remove the collection trays, discard any waste in a sink, then rinse the trays.

28. Using new gloves, firmly insert a new SEQ Reagent Tube Sipper (blue) into each dNTP port. Do not let the sipper touch any surfaces.

IMPORTANT! Be careful to push each sipper onto the port firmly. Loosely attached sippers can adversely affect results.



29. Attach each SEQ Reagent Tube to the correct dNTP port (for example, the dGTP tube on the port marked "G"), then tighten firmly by hand until it can no longer rotate.



30. After each reagent tube is securely installed, press **Next**.

The instrument automatically completes the rest of the initialization, which takes ~20 minutes. You can monitor the progress on the touchscreen.

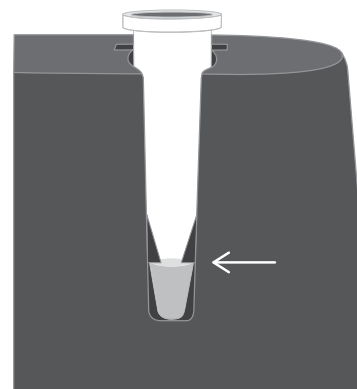
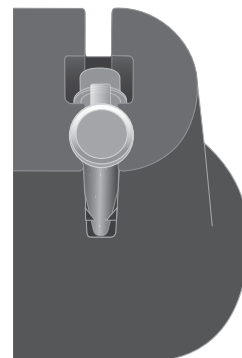
- If initialization is successful, the touchscreen indicates that initialization has "Passed" (highlighted in green). Press **Next** to return to the main menu.
- If initialization fails, see Appendix A, "Troubleshooting".

4

ISP preparation

Prepare enriched ISPs for sequencing

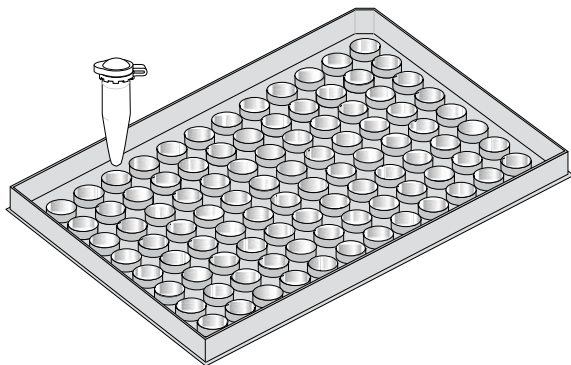
1. Confirm that the SEQ Primer is completely thawed before use. No ice crystals should be visible.
2. Vortex the SEQ Primer for ~5 seconds, then pulse centrifuge to collect the contents. Leave the tube on ice or in a chilled benchtop cold box until use.
3. Remove the enriched ISP sample in the TMPL Sample Collection Tube from the Ion OneTouch™ ES Dx Instrument or 2–8°C storage.
4. Place the tube in a microcentrifuge with a 0.2-mL tube adapter. Orient the tab of the tube lid so that it is pointing away from the center of the centrifuge (this will indicate where the sample ISPs are being pelleted).
5. Centrifuge at 15,000 rcf for 2 minutes.
6. Insert the tube into the appropriate slot of the Ion OneTouch™ Sample Rack. Open the tube lid carefully to avoid spillage, then lift the tube out of the slot, fold the lid flat against the back of the tube, and insert the tube and folded lid into the slots in the rack.
7. Use a P200 pipette and keep the pipette plunger depressed as you insert the tip into the tube containing the enriched ISPs. Carefully remove ~200 μ L of supernatant from the top down, avoiding the side of the tube where the ISPs are pelleted (i.e., the side with the tab on the tube lid). Discard the supernatant.
8. Change to a P20 pipette. Keep the pipette plunger depressed as you insert the tip into the tube, and carefully remove the remaining supernatant to the volume marked by the arrow guides on the rack. Again, avoid the side of the tube where the ISPs are pelleted. The final volume in the tube will be ~15 μ L.



9. Remove the tube from the sample rack, then add 12 μ L of thawed SEQ Primer (white cap) to the ISPs in the tube and vigorously pipet the mixture up and down 30 times.

IMPORTANT! Make sure that the pipette tip is at the bottom of the tube during mixing to avoid introducing air bubbles into the sample.

10. Insert the tube into the 96-well tray supplied with the Veriti™ Dx 96-well Thermal Cycler.



11. Load the tube and tray assembly into the thermal cycler. Select the **8 ODxTT Seq Primer Hyb** program on the thermal cycler touchscreen. Select **View** and confirm that the program steps match those listed in the table below.

Stage	Temperature	Time
Hold	95°C	2 minutes
Hold	37°C	2 minutes
Hold	25°C	Hold (up to 30 minutes)

12. When you have confirmed the program steps, run the cycling program.

After cycling, keep the tube containing the primer-annealed ISPs at room temperature. Proceed immediately to set up the sequencing run.



Sequencing protocol

Set up the sequencing run

The instrument touchscreen provides a summary of the steps in the process, which are described below in more detail.

IMPORTANT!

- To avoid damage to the chips or instrument due to electrostatic discharge:
 - **Remove your gloves when handling chips**, especially before transferring chips on or off the instrument. Hold chips by their edges when handling.
 - **Do not place chips directly on the bench or any other surface.** Always place chips either on the grounding plate on the Ion PGM™ Dx Sequencer or in the Ion PGM™ Dx Chip Minifuge bucket.

Note: Steps without gloves can be performed without risk of contamination.

- When performing one or two sequencing runs from the same initialization, the runs must be started within 27 hours after the start of initialization.

-
1. After initialization and sample preparation, go to the main menu of the Ion PGM™ Dx Sequencer, then press the **Run** button to begin the run setup.
 2. Remove the waste bottle from the instrument and completely empty the bottle. Return the waste bottle to its position on the instrument. Press **Next**.

IMPORTANT! Removing any waste before each run is critical. Waste overflow can result in a spill hazard and instrument damage.

3. Make sure the chip used for initialization is still in the chip clamp, then press **Next**. The instrument cleans the fluid lines, then proceeds automatically to the next screen.
4. Press the **Keyboard** button next to the **Planned Run** field. Using the barcode scanner attached to the sequencer, scan the **Run Short Code/Barcode** on the **View Planned Run** printout. Alternatively, type the code (displayed below the barcode) using the touchscreen keyboard. Press **OK**.

Run Short Code/ Barcode:



Note: If the Ion Torrent™ Server has ≤1 terabyte (TB) of free disk space, an alert notifies the user that there is insufficient disk space to perform the run. The run cannot proceed until data on the server is archived and deleted. Contact your IT system administrator to manually archive and delete data. An administrator-level user in the software can also change the **Archive Settings** to

reduce the **Auto archive after** interval. See the *Oncomine™ Dx Target Test Part V: Torrent Suite™ Dx Software 5.12.5 Reference User Guide*.

5. Confirm the Planned Run information displayed on the touchscreen, then press **Next**.

- Planned Run name
- Sample name if a single sample, or number of samples if multiple samples
- Assay name
- Created by [user name]
- Created date [YYYY-MM-DD]
- Ion PGM™ Dx Sequencing Kit barcode [scanned during initialization]

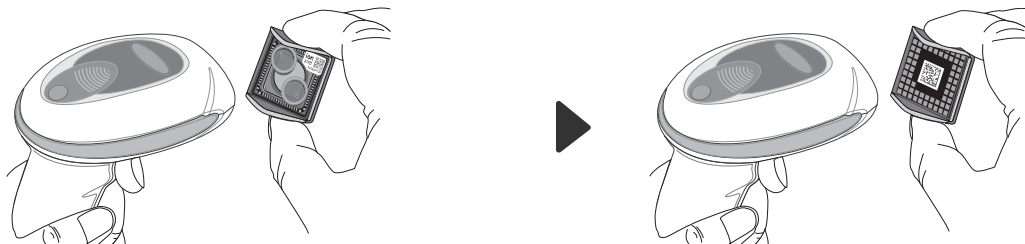
6. Remove your gloves.

Note: The following steps can be performed without gloves without risk of contamination.

7. Open the chip clamp, then remove the chip used for initialization. Press **Next**.

8. Touch the grounding plate with a bare finger, then remove a new chip from its packaging.

9. Press the **Keyboard** button next to the **Top barcode** field. Using the barcode scanner attached to the sequencer, scan the barcode on the top of the chip, then press **OK**.

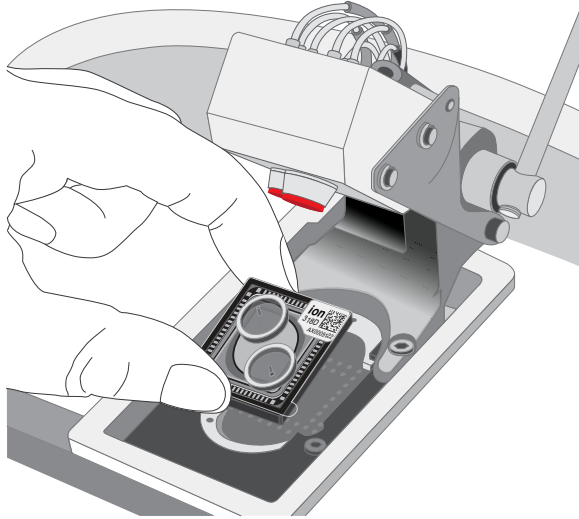


10. Press the **Keyboard** button next to the **Bottom barcode** field. Scan the barcode on the bottom of the chip, then press **OK**.

Note: If the barcode does not scan, use a new chip and contact Technical Support for a replacement.

11. Using a marker, label a corner of the chip with the **Tube Label** text.

12. Secure the new chip in the chip clamp.



13. Press **Next** to calibrate the chip. The screen prompts you to perform certain steps during calibration.
14. When prompted, visually inspect the chip in the clamp for liquid leaking from the chip case into and around the clamp area.



CAUTION! If a leak occurs, press the **Abort** button immediately, then see “Leak of unknown origin during chip calibration (before sample has been loaded on the chip)” on page 78.

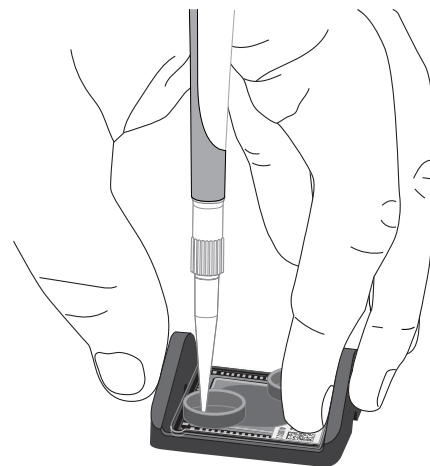
15. Close the instrument lid when prompted.
16. When chip calibration is complete, a "Calibration Passed" or "Calibration Failed" message appears.
 - If calibration passes, proceed to chip loading.
 - If calibration fails or you get an error message, see “Chip calibration failure (before sample has been loaded on the chip)” on page 77.

Load the sample on the chip

For additional instructions on using the Ion PGM™ Dx Chip Minifuge, see “Ion PGM™ Dx Chip Minifuge” on page 90.

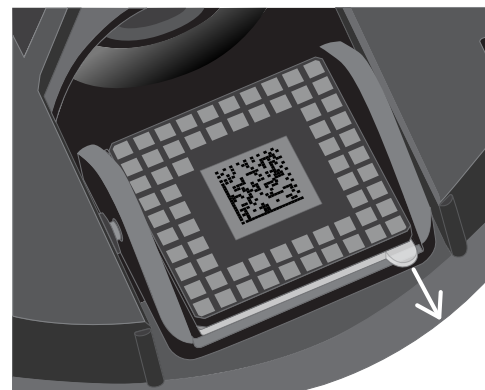
IMPORTANT! The following steps (including chip loading) should be performed without gloves, except when adding SEQ Enzyme as noted. The steps can be performed without gloves without risk of contamination.

1. Enter the Tube Label text into the Ion PGM™ Dx Sequencer touchscreen. Press **Next**.
2. Touch the grounding plate on the instrument with a bare finger, then remove the new Ion 318™ Dx Chip from the clamp.
3. Insert the used chip from initialization (marked with a "W") back into the chip clamp.
4. Place the new chip in the removable bucket from the Ion PGM™ Dx Chip Minifuge, then place the bucket on a firm, flat surface.
5. Set a P200 pipette to 30 µL. Insert the tip firmly into the chip loading port while holding the bucket and chip steady with your other hand. Remove as much liquid as possible from the port, then discard the liquid.



6. Place the chip **upside-down** in the bucket, then transfer the bucket to the Ion PGM™ Dx Chip Minifuge **with the chip tab pointing out** (away from the center of the minifuge).
7. Balance the minifuge with another upside-down chip in the opposing bucket.

Note: If you have prepared only one loaded chip, balance the minifuge with an empty used chip. Mark the used chip with a marker to differentiate it from the loaded chip.



8. Centrifuge for 3–5 seconds to completely dry the chip surface.



CAUTION! Allow the minifuge to come to a complete stop before opening the lid.

9. Remove the bucket containing the chip from the minifuge. Remove the chip from the bucket, then wipe down the bucket with a disposable wipe to remove residual liquid. Place the chip right-side up in the bucket.
10. Put on new gloves.
11. Remove the SEQ Enzyme (yellow cap) from storage and flick 4 times to mix. Pulse centrifuge the tube to collect the contents, then place the tube on ice or in a chilled benchtop cold box until use.
12. Remove the primer-annealed ISPs from the Veriti™ Dx 96-well Thermal Cycler (from “Prepare enriched ISPs for sequencing” on page 41).
13. Add 3 μL of SEQ Enzyme to the primer-annealed ISPs. Set a P200 pipette with a low-bind tip to 20 μL , then gently pipet the mix up and down 10 times.

Note: If there is condensation on the walls of the tube, push down with the end of the pipette tip to mix the condensation with the sample.

14. Incubate at room temperature for 5 minutes.
15. Remove your gloves.

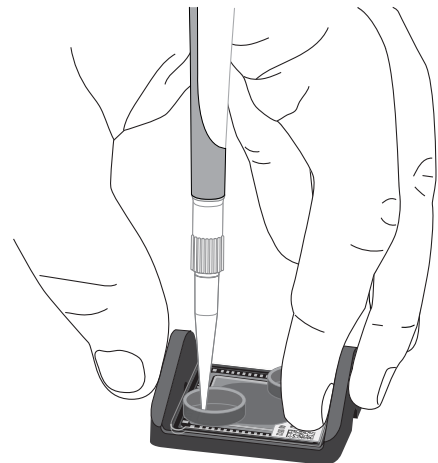
Note: The following steps can be performed without gloves without risk of contamination.

16. Place the chip in the minifuge bucket on a firm, flat surface. Following the 5-minute polymerase incubation, collect 30 μL of sample into a P200 pipette tip. Discard any remaining sample.

Note: If the sample is less than $\sim 30 \mu\text{L}$, there will be a visible air gap at the end of the tip. To correct the volume, see “Sample volume is $<30 \mu\text{L}$ ” on page 83.

17. Insert the pipette tip firmly into the loading port of the chip. Apply gentle pressure between the tip and chip throughout the loading process.

IMPORTANT! Do not remove the pipette tip from the port or dial up the pipette during loading.



18. With the pipette unlocked, dial down the pipette to deposit the ISPs. Ensure that the entire sample is loaded onto the chip. To avoid introducing air bubbles, leave a small amount of sample in the pipette tip (~0.5 μL).

Note: A small amount of sample may leak from the outlet port, which is acceptable. However, if a large amount of sample leaks from the outlet port (~15 μL or greater), do not continue with the run, and contact Technical Support.



19. Transfer the chip in the bucket to the minifuge **right-side up with the chip tab pointing in** (toward the center of the minifuge).
20. Balance the minifuge with another right-side up chip in the opposing bucket, then centrifuge for 30 seconds.

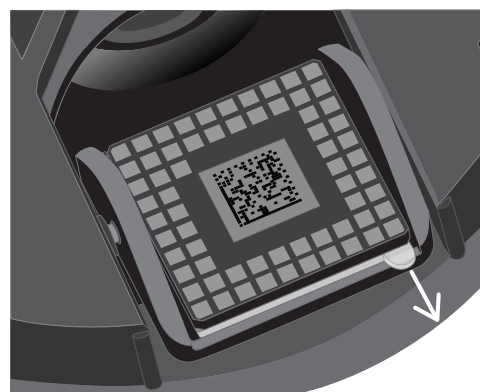
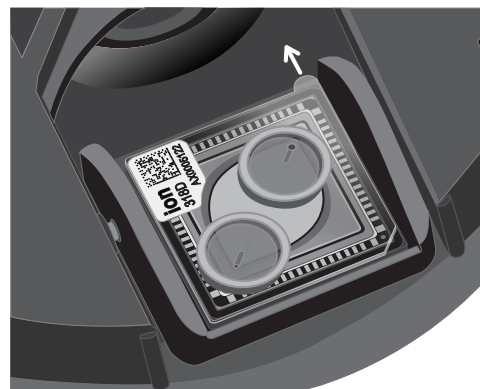


CAUTION! Allow the minifuge to come to a complete stop before opening the lid.

21. Remove the bucket containing the chip from the minifuge and place it on a flat surface. Set a P200 pipette to 40 μL , then pipet the liquid out through the loading port. Remove as much liquid from the chip as possible.

Note: It is normal to have some liquid remain in the chip after this step.

22. Turn the chip upside-down in the bucket, then place the bucket back in the minifuge with the chip tab pointing out.
23. Balance the minifuge with another upside-down chip in the opposing bucket, then pulse centrifuge for 3–5 seconds at maximum speed to dry the chip surface completely.
24. Remove the chip from the bucket, then wipe down any residual liquid left on the bucket with a disposable wipe.



Proceed immediately to “Perform the run”.

Perform the run

1. On the Ion PGM™ Dx Sequencer touchscreen, press **Next**, then press the **Keyboard** button next to the **Top barcode** field. Using the barcode scanner attached to the sequencer, scan the barcode on the top of the new loaded chip, then press **OK**.
2. Remove the used chip from the instrument, then use ungloved hands to insert the loaded chip into the chip clamp.

IMPORTANT! Confirm that both red rubber port gaskets are securely in place in the clamp.

3. Press **Next** to calibrate the loaded chip. At the start of calibration, visually inspect the chip for leaks before closing the instrument cover. Close the lid when prompted to do so.



CAUTION! If a leak occurs, press the **Abort** button immediately, then see “Leak of unknown origin during chip calibration (after sample has been loaded on the chip)” on page 80.

4. After ~1 minute, the touchscreen indicates if calibration was successful.
 - If the chip fails calibration, reseal the chip and press **Re-try** to recalibrate. If the chip continues to fail calibration, see “Error message: Calibration failed (after sample has been loaded on the chip)” on page 81.
 - If a message says that the chip is not seated correctly, see “Error message: Calibration failed (after sample has been loaded on the chip)” on page 81.
5. When the chip passes calibration, press **Next** to start the sequencing run. The sequencing run takes ~4.5 hrs to complete.

IMPORTANT! Confirm the run has started before leaving the sequencer. During a run, do not touch the instrument or the attached bottles or tubes, because this can reduce the quality of the measurements.

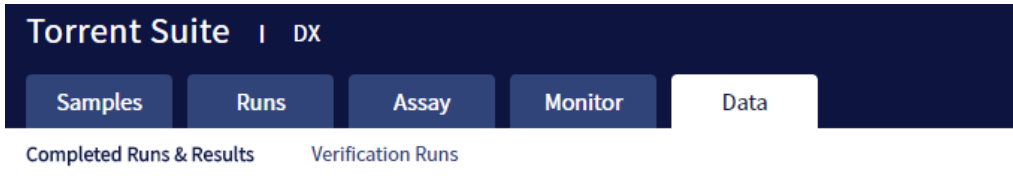
6. After the run, the touchscreen returns to the main menu.

Review the results and generate reports as described in the next chapter.

Note: Used chips cannot be reused for sequencing. Used chips can be marked for cleaning and initialization as described previously.

6

Review data and results



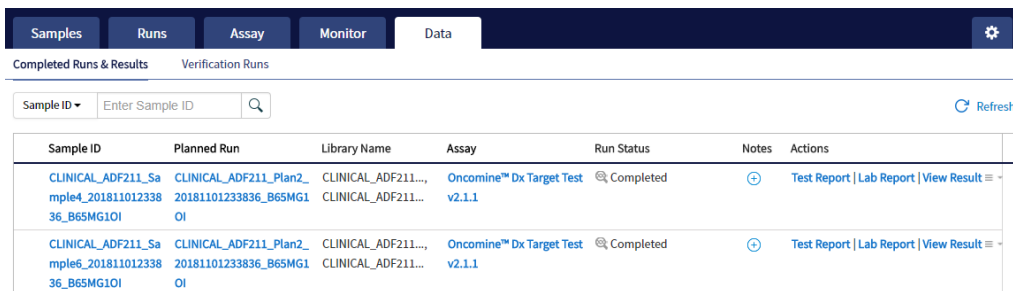
In the **Data** tab, you can review run results and perform data analysis and data management tasks:

- Select **Completed Runs & Results** (the default window) to review completed sample run results and reports. Run results are listed by **Sample ID**.
- Select **Verification Runs** to review data from completed verification runs performed during installation or PQ validation.

Completed Runs & Results screen

Under the **Data** tab, in the **Completed Runs & Results** screen, samples that have been sequenced are listed by Sample ID.

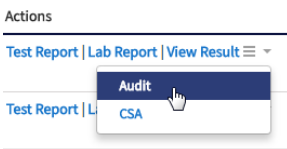
You can search the list of results by Sample ID or Planned Run name. You can also filter and sort the list.



The following information is displayed in the screen:

Column	Description
Sample ID	The unique identifier created when the sample was entered into the software. Click on the link to display the details of the sample.
Planned Run	The name of the Planned Run given when it was created in the software, after preparation of the library batch. Click on the name to display details of the Planned Run.

(continued)

Column	Description
Library Name	The names of the DNA and RNA sample libraries given during preparation of the library batch.
Assay	The assay selected when the sample was placed in a library batch prior to creating the Planned Run. Click on the assay name to display the details of the assay.
Run Status	The status of the run (for example: completed, failed, terminated).
Notes	User-entered notes about the sample. To view previously written notes, click . To add new notes, click .
Actions	<p>Click the appropriate link to:</p> <ul style="list-style-type: none"> Download the Test Report. Download the Lab Report. Open the View Result screen. View the audit trail for the Planned Run. Click , then click Audit.  <ul style="list-style-type: none"> Download Customer Support Archive (CSA) log files for the run to aid in troubleshooting. Click , then click CSA.

Download the Lab Report or Test Report

1. To download the Lab Report or Test Report for a sample of interest, in the **Completed Runs & Results** screen, click the appropriate link in the **Actions** column for the sample. A ZIP file containing all languages of the report downloads automatically.

Completed Runs & Results Verification Runs

Sample ID ▾ Refresh

Sample ID	Planned Run	Library Name	Assay	Run Status	Notes	Actions
CLINICAL_ADF211_Sample 4_20181101233836_B65MG 10I	CLINICAL_ADF211_Plan2_201811 01233836_B65MG10I	CLINICAL_ADF211..., CLINIC AL_ADF211...	OncoPrint™ Dx Target Test v2.1.1	Completed		Test Report Lab Report View Result

2. Extract the downloaded files, then open the PDF file of the desired language in an appropriate viewer.

Test Report

The Test Report (available in the **Completed Runs & Results** screen) is a clinical report generated by the software that can be downloaded in PDF format. It is identical to the Lab Report, except that it does not include the Sequencing Run Details and Control and Sample QC Evaluation Metrics sections. The Test Report contains the following sections and information.

Note: The display of variants in the Test Report depends on cancer type.

Section	Description
Sample Details	The sample and patient information entered into the software.
Results for Sequence Variations for Therapeutic Use	Displays the results for the clinical variants and gene fusions in the sample, and any recommended therapies that are clinically indicated for the selected cancer type.
Results for Analytical Sequence Variations Detected	Displays the results for analytical variants and gene fusions detected in the sample, depending on the variants associated with the cancer type.
Test Description	A description of the test and the gene variants in associated tissue types that it screens for.
Results for Analytical Sequence Variations Not Detected	Displays the results for analytical variants and gene fusions not detected in the sample, depending on the variants associated with the cancer type.

Lab Report

The Lab Report (available in the **Completed Runs & Results** screen) is a report generated by the software that can be downloaded in PDF format. The Lab Report contains the following sections and information.

Note: The display of variants in the Lab Report depends on cancer type.


Section	Description
Sample Details	The sample information entered into the software.
Results for Sequence Variations for Therapeutic Use	The results for the clinical variants and gene fusions in the sample, and any recommended therapies that are clinically indicated for the selected cancer type. Allele frequencies are also reported.
Results for Analytical Sequence Variations Detected	A list of the analytical variants and gene fusions detected by the assay, and associated information for each, depending on the variants associated with the cancer type.
Test Description	A description of the assay.
Analytical Sequence Variations Not Detected	A list of all the analytical variants and gene fusions not detected by the test, and associated information for each, depending on the variants associated with the cancer type.

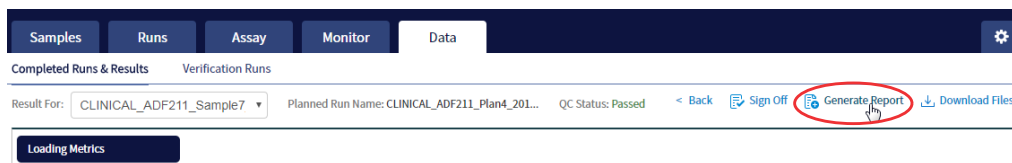
(continued)

Section	Description
Sequencing Run Details	A list of all the kits and instruments used to perform the test.
Control and Sample QC Evaluation Metrics	A summary of the quality control metrics.

Generate customized reports

By default, the Test Report and Lab Report are generated in the language that is selected in the report template. To generate these reports in another language, or change the types of variant calls that are reported, do the following:

1. In the **Completed Runs and Results** screen, click **View Result**.
2. In the **View Result** screen, click  **Generate Report**.

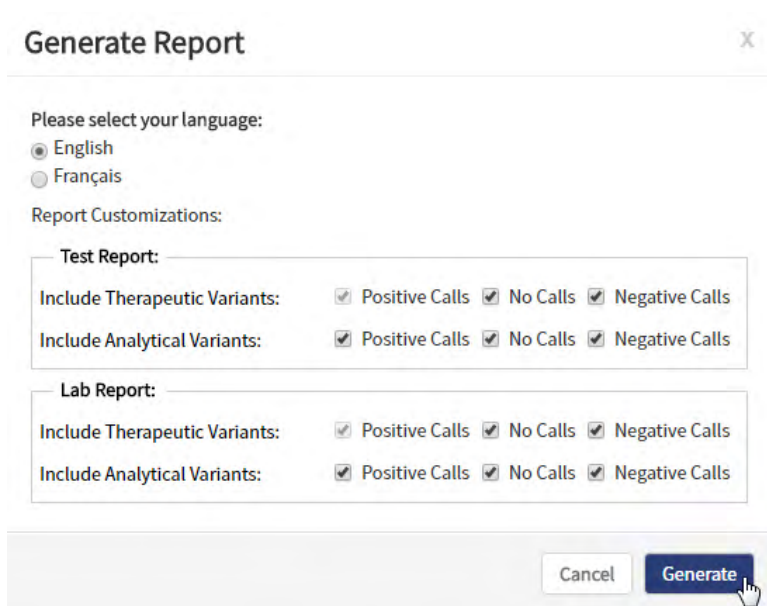


The **Generate Report** dialog box opens.

3. Do one or both of the following:

- Select the desired language.
- Select the checkboxes for the positive calls, no calls, or negative calls that you want to report in the Test Report and/or Lab Report.

Note: These checkboxes are only available in IVD mode if they are enabled for the selected assay.



Generate Report X

Please select your language:

English
 Français

Report Customizations:

Test Report:

Include Therapeutic Variants: Positive Calls No Calls Negative Calls

Include Analytical Variants: Positive Calls No Calls Negative Calls

Lab Report:

Include Therapeutic Variants: Positive Calls No Calls Negative Calls

Include Analytical Variants: Positive Calls No Calls Negative Calls

Cancel Generate

Note: Positive calls for therapeutic variants cannot be deselected.

4. Click **Generate**.

A message displays when the report has been generated. The new report overwrites the existing report.

- 5.**
- Click
- [↓](#)
- Download Files**
- to open a dialog box to select the new report and other run-related files for download.

View results

The run results shown in the screens described in this section can also be downloaded in the form of results files (see “Results files” on page 66).

- In the **Completed Runs & Results** screen, click **View Result** to view the sequencing run results for a sample of interest.

The **Results** screen for the selected sample opens, with the **Loading Metrics** screen displayed. Result categories are arranged vertically on the left side of the screen. The QC status for the run is listed at the top of the screen.

QC Status: Passed

	Name	Count	Percentage
Loading Metrics	Total Addressable Wells	11287275	-
	Wells With ISPs	8533877	75.6 %
	Live ISPs	8527652	99.9 %
	Control ISPs	49654	0.6 %
Filtering Metrics	Library ISPs	8477998	99.4 %
	Filtered: Polyclonal	2567923	30.3 %
	Filtered: Primer-Dimer	16404	0.2 %
	Filtered: Low Quality	1602223	18.9 %
	Usable Library Reads	4291448	50.6 %

- ① QC status
- ② Selected sample
- ③ Result categories

IMPORTANT! A **QC Status** of "Passed" does not guarantee that the genotypes of all analytically relevant variants are determined. See the **Test Result** column in the Lab Report or Test Report for any "No Call" results when interpreting the results.

- Click the links on the left side of the screen to display results of interest, including QC results and results for different categories of variants.

- To view results for another sample in the same run, select the sample from the **Result For:** dropdown list. All samples in a run share the same **Loading Metrics** data. The other result categories are sample-specific.

Completed Runs & Results Verification Runs

Result For: CLINICAL_ADF211_Sample3 Planned Run Name: CLINICAL_ADF

Run Results

CLINICAL_ADF211_Sample3_20181006014359_FEYI1LM1

CLINICAL_ADF211_Sample7_20181006014359_FEYI1LM1



CLINICAL_ADF211_Sample6_20181006014359_FEYI1LM1

QC Report CLINICAL_ADF211_Sample5_20181006014359_FEYI1LM1

Summary CLINICAL_ADF211_Sample2_20181006014359_FEYI1LM1

CLINICAL_ADF211_Sample4_20181006014359_FEYI1LM1

Therapeutic Loading Metrics Live ISPs

- To generate the Lab Report and Test Report in other languages, click  **Generate Report** (see “Generate customized reports” on page 53).
- To download individual results files for further analysis, click  **Download Files** (see “Results files” on page 66).
- To return to the **Completed Runs & Results** screen, click **< Back**.

Loading metrics

The **Loading Metrics** link in the **View Result** screen displays the following loading and filtering metrics for the run:

Metric	Description
Loading Metrics	
Total Addressable Wells	The total number of wells on the chip – excluded wells.
Wells with ISPs	The number (count) and percentage of chip wells that contain ISPs. The percentage is expressed as a percent of total addressable wells.
Live ISPs	The number (count) and percentage of chip wells containing live ISPs (ISP's templated with library or control fragment), with the percentage expressed as a percent of wells with ISPs.
Control ISPs	The number (count) and percentage of ISPs that have a key signal identifying them as internal controls, with the percentage expressed as a percent of live ISPs.
Library ISPs	The number (count) and percentage of ISPs that have a key signal identical to the library key signal, with the percentage expressed as a percent of live ISPs.

(continued)

Metric	Description
Filtering Metrics^[1]	
Filtered: Polyclonal	ISPs carrying clones from two or more templates, with the percentage expressed as a percent of library ISPs.
Filtered: Primer-Dimer	ISPs with an insert length of less than 8 bp, with the percentage expressed as a percent of library ISPs.
Filtered: Low Quality	ISPs with low or unrecognizable signal, with the percentage expressed as a percent of library ISPs.
Usable Library Reads ^[2]	Number (count) and percentage of library ISPs passing all filters.

^[1] Filtering Metrics only apply to ISPs templated with library fragments, not control fragment.

^[2] Values in the "Filtered:" rows are subtracted from the Library ISPs value (Loading Metrics) to give the Usable Library Reads value.

QC Report

The **QC Report** link displays metrics for the sample libraries and internal controls. This information is also accessible through the **Monitor** tab for the last five runs.

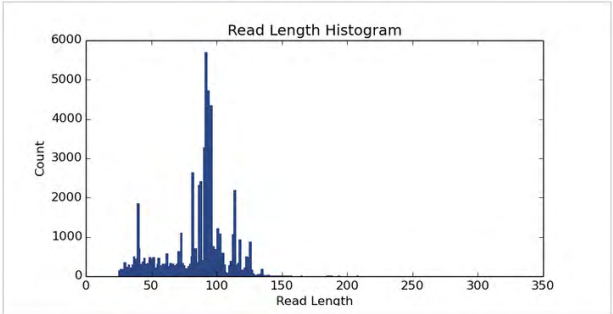
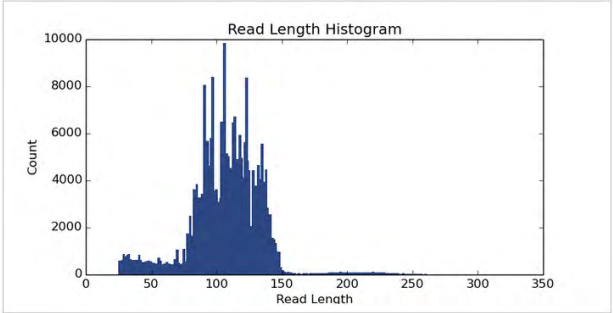
IMPORTANT! The following table describes all the quality control metrics that can be displayed. Whether particular metrics are displayed depends on the assay configuration.

Metric	Description
Library QC Evaluation Metrics	
Library QC: Library RNA	<ul style="list-style-type: none"> • Mean AQ20 Read Length (bp): The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of a library. • Mean Read Length (bp): The average length, in base pairs, of all reads reported for a given library. • Total Mappable Reads: The number of reads that are mapped to the fusion reference file.^[1]
Library QC: Library DNA	<ul style="list-style-type: none"> • Mean AQ20 Read Length (bp): The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of a library. • Mean Read Length (bp): The average length, in base pairs, of all reads reported for a given library. • Percent Reads: The number of library reads normalized by the total addressable wells in a run.

(continued)

Metric	Description
Control QC Evaluation Metrics	
Control QC: CF-1	<ul style="list-style-type: none"> • Base Call Accuracy: 1 – (total number of errors for all positions in the control / total number of aligned bases) for Control Fragment-1 (CF-1) reads. • Key Signal: The average peak signal from the incorporation trace measured across the 1-mer incorporations in the CF-1 key sequences. [2] • Mean AQ20 Read Length (bp): The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of CF-1. • Percent Reads: The number of all usable library reads that aligned with the CF-1 sequence divided by the total number of addressable wells.
Control QC: DNA Control	<ul style="list-style-type: none"> • COSMIC ID variant calls and allelic frequencies: The individual variant positions and wild-type positions that are assessed in the DNA control reagent for presence or absence of the variant. • Mean AQ20 Read Length (bp): The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of a control. • Percent Reads: The number of all usable library reads that aligned with the control fragment sequence divided by the total number of addressable wells.
Control QC: RNA Control	<ul style="list-style-type: none"> • Fusion calls and number of reads: The individual fusion calls and number of reads that are assessed in the RNA control reagent for the presence or absence of the fusion. • Total Mappable Reads: the number of reads that are mapped to the fusion reference file.
Control QC: DNA NTC	<ul style="list-style-type: none"> • Hotspot Calls: The total number of hotspots where a call was made in the no-template control.
Control QC: RNA NTC	<ul style="list-style-type: none"> • Total Fusion Calls: The total number of fusion locations where a call was made in the no-template control. • Total Mappable Reads: The number of reads in the no-template control that are mapped to the fusion reference file.

(continued)

Metric	Description
Histogram of Read Length ^[3]	<ul style="list-style-type: none"> A library-specific read-length histogram: <div style="display: flex; flex-direction: column; align-items: center;"> <div style="margin-bottom: 20px;"> <p>RNA</p>  <p>The RNA histogram shows a distribution of read lengths from 0 to 350. The y-axis is labeled 'Count' and ranges from 0 to 6000. The x-axis is labeled 'Read Length' and ranges from 0 to 350. The distribution is centered around 100, with a peak count of approximately 5500.</p> </div> <div> <p>DNA</p>  <p>The DNA histogram shows a distribution of read lengths from 0 to 350. The y-axis is labeled 'Count' and ranges from 0 to 10000. The x-axis is labeled 'Read Length' and ranges from 0 to 350. The distribution is centered around 100, with a peak count of approximately 9500.</p> </div> </div>

^[1] May not equal the sum of the individual fusions, since some detected fusions are not included in the fusion BED file.

^[2] The minimum Key Signal value for the CF-1 control (i.e., the Templating Control) is set to 0 in all preinstalled assays and new custom assays in Torrent Suite™ Dx Software version 5.8 and later. Custom assays created in previous versions of the software may have a higher Key Signal value for the control. If a run using an older custom assay fails QC due to this metric, reanalyze the run using a minimum Key Signal of 0 for the control. For future runs, create a new custom assay based on the old assay, changing the minimum Key Signal to 0.

^[3] Not displayed in the Monitor tab.

Summary

The **Summary** link displays the following information for the run:

Field	Description
Run and Configuration Summary	
Run Name	The name of the run.
Assay Name	The name of the assay.
Reference Genome	The reference genome used for analysis.
Target Region	The name of the targeted regions BED file used.
Hotspot Regions	The name of the hotspot regions BED file used.
Fusion Reference	The name of the fusion reference used for analysis, when applicable.
Fusion Panel	The name of the fusion panel used, when applicable.
Library Name	The names of the DNA and DNA libraries prepared from the sample, entered during library batch preparation.
Amplicon Summary	
Number of Targets	The number of amplicons in the panel.
On Target Reads	The number of reads mapped to the target amplicons.
Percent On Target Reads	The percentage of reads mapped on target to total reads.
Percent Full Length On Target Reads	The percentage of full length reads mapped on target to total reads.
Average Coverage	The ratio of the depth of coverage at each base in the target region to the length of the target region.
Target Coverage at 20X	The percentage of base positions with depth of coverage $\geq 20X$ in the target region.
No Strand Bias	The percentage of bases with a strand bias between 30% and 70%.
Coverage Uniformity	The ratio of passed number of targets to total number of targets, where passed number of targets is the number of amplicons that have at least 0.2X mean coverage.
Variant Summary^[1]	
Number of SNVs/MNVs	The number of single- and multi-nucleotide variations (SNVs/MNVs).
Number of INDELS	The number of insertions or deletions.
Number of Fusions	The total number of fusion calls.

^[1] Includes Therapeutic, Analytical, and Level2 variants in the **View Result** screen.

Therapeutic variant results

Therapeutic variant results for the selected sample are provided under the **Therapeutic** section of the left navigation bar in the **Results** screen. Depending on the assay, results for these variants may be divided into multiple subscreens, each with a separate link (**SNV/INDEL**, **Fusion**, etc.).

The results are determined by the reference sequences installed on the server, the QC controls used in the run, the assay used for the run, and any Reporting Gene List associated with that assay.

Columns in the screen or subscreens are described in the following table.

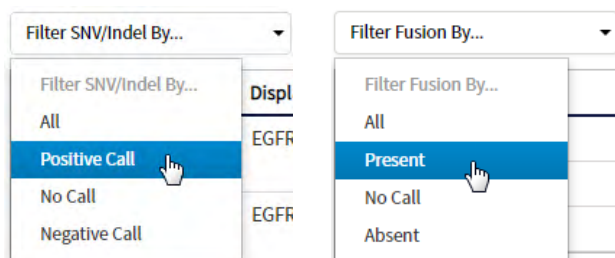
Column	Description
Summary	
Gene Fusion Present Variants Present	The therapeutic gene fusion or SNV/INDEL variant detected in the sample
Associated Therapy	The therapy indicated for each variant listed
SNV / INDEL	
Gene	The gene name, which provides a link to the View Annotation Sources popup with additional information about the HotSpot ID (see “View annotation sources” on page 64)
Display Name	The gene name with the exonic location of the deletion or insertion
Amino Acid Change	The amino acid change using HGVS-nomenclature
Nucleotide Change	Identification of the DNA-level nucleotide change using HGVS-nomenclature
Test Result	The presence or absence of the variant
Allele Frequency	The most likely frequency of the variant allele ^[1]
HotSpot ID	The name of the hotspot as defined in the BED file
Locus	The one-base position of the variant in the reference genome
Type	The type of variation detected: <ul style="list-style-type: none"> • snv/mnv (single nucleotide variation/multi-nucleotide variation) • ins (insertion) • del (deletion) • complex (Variant that includes both a deletion and insertion of DNA sequences relative to the reference sequence)
Genotype	The genotype at the locus
Ref	The reference base or bases at the locus

(continued)

Column	Description
Quality Score	The relative probability of either the "reference" hypothesis interval [0,cutoff) or the "variant" hypothesis interval [cutoff,1], on a Phred scale (-10*log10). This provides a measure of the strength of the evidence for the variant call. A higher score means more evidence for the call. Quality scores are capped at 100.
Coverage	The number of reads covering the variant position after down-sampling
Fusion	
Gene	The gene that regulates expression of the gene fusion
Display Name	The gene name of the fusion
Read Count	The number of valid reads aligned to the specific fusion sequence
Test Result	The presence or absence of that fusion variant

[1] The allele frequency is the most likely variant frequency in the reads after corrections are made for probable errors. The software uses this corrected frequency and the uncertainty in the observations to calculate the probability that the variant frequency in the sample falls within defined intervals that can be set by the user. The user sets a threshold frequency for deciding between genotypes: this defines intervals $[0,c)[c,1-c)[1-c,1]$ corresponding to the three diploid genotypes (reference, heterozygous, homozygous) respectively, where "c" is the minimum variant frequency set during the creation of the assay. The software reports the genotype corresponding to the interval with the highest probability of containing the variant frequency in the sample. In most cases, the reads are highly accurate and do not have a high probability of error. In such cases, the observed frequency and the most likely frequency are similar, and are contained within a single interval leading to a high quality genotype. In some cases, if very few reads are observed or the error rate is high, the observed counts and the most likely counts can be different, and the uncertainty in the real sample frequency can be high. When this happens, it is often impossible to exclude a heterozygous population in the sample, and the genotype is assigned to be heterozygous, since a significant portion of the probability falls in that interval.

Note: You can filter the variant lists using options in the **Filter SNV/Indel By...** and **Filter Fusion By...** dropdown lists in the upper left corner of the results report.



Analytical variant results

Analytical variant results for the selected sample are provided under the **Analytical** section of the left navigation bar in the **Results** screen. Depending on the assay, results for these variants may be divided into multiple subscreens, each with a separate link (**SNV/INDEL**, **Fusion**, etc.).

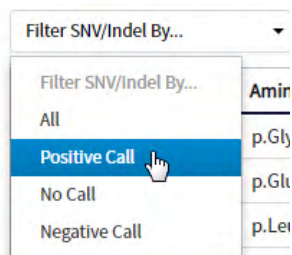
The results are determined by the reference sequences installed on the server, the QC controls used in the run, the assay used for the run, and any Reporting Gene List associated with that assay.

Columns in the screen or subscreens are described in the following table.

Column	Description
Summary	
Variants Present	Lists the analytical SNV or INDEL variants detected in the sample
SNV / INDEL	
Gene	The gene name, which provides a link to the View Annotation Sources popup with additional information about the HotSpot ID (see “View annotation sources” on page 64)
Amino Acid Change	The amino acid change using HGVS-nomenclature
Nucleotide Change	The DNA-level nucleotide change using HGVS-nomenclature
Test Result	The presence or absence of the variant
Allele Frequency	The most likely frequency of the variant allele
HotSpot ID	The name of the hotspot as defined in the BED file
Locus	The one-base position of the variant in the reference genome
Type	The type of variation detected: <ul style="list-style-type: none"> • snv/mnv (single nucleotide variation/multi-nucleotide variation) • ins (insertion) • del (deletion) • complex (Variant that includes both a deletion and insertion of DNA sequences relative to the reference sequence)
Genotype	The genotype at that locus
Ref	The reference base or bases at the locus
Quality Score	The relative probability of either the "reference" hypothesis interval [0,cutoff) or the "variant" hypothesis interval [cutoff,1], on a Phred scale (-10*log10). This provides a measure of the strength of the evidence for the variant call. A higher score means more evidence for the call. Quality scores are capped at 100.
Coverage	The number of reads covering the variant position after down-sampling
Fusion^[1]	
Gene	The gene that regulates expression of the gene fusion
Display Name	The gene name of the fusion
Read Count	The number of valid reads aligned to the specific fusion sequence
Test Result	The presence or absence of that fusion variant

^[1] Note that for the OncoPrint™ Dx Target Test, no analytical fusions are reported except for expression controls.

Note: You can filter the variant list using options in the **Filter SNV/Indel By...** dropdown list in the upper left corner of the results report.



View annotation sources

You can view additional information for each hotspot ID listed under the **SNV/Indel** links of the **Therapeutic** and **Analytical** sections in the **View Results** screen. The gene name in the **SNV/Indel** report is a link that opens the **View Annotation Sources** dialog box, which provides information for the particular hotspot.

View Annotation Sources

Transcript: NM_005228.3
 PolyPhen: NA
 Sift: NA
 Alternate Allele Count: 0,254,0
 p-value: 1.1694993910198652E-85
 Variant Effect: ["nonframeshiftBlockSubstitution"], ["nonframeshiftDeletion"]
 Filtered Coverage: 1936
 UCSC common SNPs: NA
 MAF: NA
 Zygosity: NA
 rs121913423:rs121913422:rs121913436:rs121913425:rs121913435:rs121913424:rs121913229:rs121913438:rs121913437:rs121913426:rs121913439:rs121913463:rs121913441:rs121913231:rs121913440:rs121913421:rs121913464:rs121913442
 Allele Ratio: 0,0,0.13120000064373016,0,0
 Grantham: NA
 Allele Read-Count: 254

Close

Example of transcript information in the VCF

The COSM1074639 hotspot ID has multiple transcripts associated with it, which include a mutation from 'A' to 'C'. These are listed in the VCF file as shown below:

chr6 152419923 COSM1074639;COSM1074637 A C,G 122.19 PASS
AF=0,0;AO=0,0;DP=1116;FAO=0,0;FDP=1116;FR=.;FRO=1116;FSAF=0,0;FSAR=0,0;FSRF=661;FSRR=455;F
WDB=0.00426152,-
0.0087378;FXX=0;HRUN=1,1;HS;LEN=1,1;MLLD=103.308,103.438;QD=0.437949;RBI=0.0394298,0.03043
16;REFB=-2.22051E-5,-2.32713E-5;REVB=-0.0391988,-
0.0291502;RO=1115;SAF=0,0;SAR=0,0;SRF=661;SRR=454;SSEN=0,0;SSEP=0,0;SSSB=-6.44874E-8,-
6.44874E-
8;STB=0.5,0.5;STBP=1,1;TYPE=snp,snp;VARB=0,0;OID=COSM1074639,COSM1074637;OPOS=152419923,
152419923;OREF=A,A;OALT=C,G;OMAPALT=C,G;FUNC=[

```
{'normalizedRef':'A','transcript':'NM_001122742.1','grantham':'144.0','gene':'ESR1','location':'exonic','ori  
gAlt':'C','origPos':'152419923','origRef':'A','normalizedPos':'152419923','exon':'10','function':'missense',  
protein':'p.Tyr537Ser','normalizedAlt':'C','gt':'neg','codon':'TCT','coding':'c.1610A>C'},
```

```
{'normalizedRef':'A','transcript':'NM_001122742.1','grantham':'194.0','gene':'ESR1','location':'exonic','ori  
gAlt':'G','origPos':'152419923','origRef':'A','normalizedPos':'152419923','exon':'10','function':'missense',  
protein':'p.Tyr537Cys','normalizedAlt':'G','gt':'neg','codon':'TGT','coding':'c.1610A>G'},
```

```
{'normalizedRef':'A','transcript':'NM_001122741.1','grantham':'144.0','gene':'ESR1','location':'exonic','ori  
gAlt':'C','origPos':'152419923','origRef':'A','normalizedPos':'152419923','exon':'9','function':'missense',  
protein':'p.Tyr537Ser','normalizedAlt':'C','gt':'neg','codon':'TCT','coding':'c.1610A>C'},
```

```
{'normalizedRef':'A','transcript':'NM_001122741.1','grantham':'194.0','gene':'ESR1','location':'exonic','ori  
gAlt':'G','origPos':'152419923','origRef':'A','normalizedPos':'152419923','exon':'9','function':'missense',  
protein':'p.Tyr537Cys','normalizedAlt':'G','gt':'neg','codon':'TGT','coding':'c.1610A>G'},
```

```
{'normalizedRef':'A','transcript':'NM_001122740.1','grantham':'144.0','gene':'ESR1','location':'exonic','ori  
gAlt':'C','origPos':'152419923','origRef':'A','sift':'0.0','normalizedPos':'152419923','exon':'9','function':'mi  
ssense',protein':'p.Tyr537Ser','gt':'neg','normalizedAlt':'C','codon':'TCT','polyphen':'0.979','coding':'c.161  
0A>C'},
```

```
{'normalizedRef':'A','transcript':'NM_001122740.1','grantham':'194.0','gene':'ESR1','location':'exonic','ori  
gAlt':'G','origPos':'152419923','origRef':'A','sift':'0.0','normalizedPos':'152419923','exon':'9','function':'mi  
ssense',protein':'p.Tyr537Cys','gt':'neg','normalizedAlt':'G','codon':'TGT','polyphen':'0.998','coding':'c.16  
10A>G'},
```

Results files

The following files can be downloaded from the **Results Report** window. To download the files, click **Download Files**, select the files to download, then click **Download**.

File name	Description
Test Report	A report of the completed analysis in PDF format
Lab Report	A clinical lab report of the completed analysis in PDF format; includes both clinical and analytical results.
PlannedRun-AuditTrail.pdf	Contains all audit records pertaining to the Planned Run.
Info.csv	Contains information about the run and analysis, such as software, sequencing information, instrument information, analysis information, QC details etc.
<RNABarcode>_rawlib.basecaller.bam	Unmapped RNA BAM File; output of base calling, contains unmapped reads.
Snvindel.tab	A tab-delimited file that contains information about non-targeted SNVs and indels
<RNABarcode>_rawlib.basecaller_alignments.bam	Mapped RNABarcode BAM file; output after reads have been mapped to the fusion reference.
Target_Summary.tab	A tab-delimited file that contains a targeted test results summary
<RNABarcode>_rawlib.basecaller_alignments.bam.bai	Mapped RNABarcode BAM index file
<RNABarcode>_rawlib.basecaller.fastq	FASTQ file generated from unmapped BAM file of the RNA barcode used.
<DNABarcode>_rawlib.basecaller.bam	Unmapped DNA barcode BAM file; output of base calling, contains unmapped reads.
raw_peak_signal	Key signal gives the percentage of LiveSPs with a key signal that is identical to the library key signal.
<LibPrepID>_<analysisID>.final.vcf	A VCF file containing all the variants detected as a result of the analysis, along with information such as test result, read count, gene name, quality scores, etc.
Summary.tab	A tab-delimited file that contains the on-targeted test results summary
<LibPrepID>_rawlib.stats.cov.txt	Amplicon statistics file


(continued)

File name	Description
Fusion.tab	A tab-delimited file that contains non-targeted (analytical) fusion details in a table format. Note: The information displayed in the file for each isoform of a particular fusion is identical, because specific isoform and locus information is not included in this table. Detailed isoform and locus information is available in the <LibraryPrepID>.<AnalysisID>.final.vcf file, available under Download Files .
readLenHisto.png	Gives the read-length distribution of FASTQ files in the form of a histogram. A thumbnail histogram of the read lengths for a particular barcode.
<LibPrepID>_rawlib.bam.bai	Mapped DNA barcode BAM index file (index file of DNA barcode-mapped BAM file)
Basecaller.log	Base Caller log file
analysis.log	Analysis log file
sigproc.log	Signal processing log file
Bead_density_contour.png	Loading density image; a pseudo-color density image of the Ion Chip plate showing percent loading across the physical surface
<DNABarcode>_rawlib.basecaller.fastq	FASTQ file of the DNA barcode used
Target_cnv.tab	Targeted CNV detail table (CNV results for targeted variants from a sequencing run) Note: For use with IVD tests that include CNV reporting.
cnv.tab	Non-targeted CNV detail table (analytical CNV results from a sequencing run) Note: For use with IVD tests that include CNV reporting.
Target_fusion.tab	A tab-delimited file that contains targeted (clinical) fusion details in a table format
<LibPrepID>_rawlib.bam	Mapped DNA barcode BAM File; output after mapping reads to reference.
Iontrace_Library.png	Key incorporation trace image showing the average signal readings for flows of the bases T, C, and A in the library key.
rawtf.basecaller.fastq	FASTQ file for the test fragment

Sign the run results

In the **View Result** screen, Managers/Administrators can provide their electronic signature on the run results. The signature information appears in the **QC Report** in the **View Result** screen, and in the downloaded Test Report and Lab Report PDFs.

Multi-language support for PDF report generation is provided. By default reports are generated in the language that is selected in the **Report Template** used. When reports are generated in multiple languages, **Sign Off** occurs only in the report of the default language.

1. At the top of the **View Result** screen, click  **Sign Off**, then enter your user name, password, and comments in the dialog box. Fields identified with a red asterisk (*) are required fields.
2. In the **Footer Field**, enter any text.
3. Click **Sign Off** to confirm your electronic signature.

Files in the Reports folder

When a manager- or administrator-level user signs a report, a folder named with the Sample ID is created in the Reports folder on the server (`/results/analysis/output/reports`), and the following files are copied into it:

Info file (.csv)	Non-targeted Test Results Summary (.tab)
Signal processing log file (.log)	Targeted SNV/INDEL Detail Table (.tab)
Targeted Fusion Detail Table (.tab)	Amplicon Stats (<code>_rawlib.stats.cov.txt</code>)
Targeted Test Results Summary (.tab)	RNA FASTQ File (.fastq)
Analysis log file (.log)	RNA Mapped BAM file (.bam)
VCF file (.vcf)	RNA Unmapped BAM file (.bam)
DNA Mapped BAM file (.bam)	Test Fragment FASTQ File (.fastq)
DNA Unmapped BAM file (.bam)	Read Length Histogram (.png)
Key Signal	Test PDF Report (<i>optional</i>) (.pdf)
Key Incorporation Trace (.png)	Lab PDF Report (.pdf)
Fusion Detail Table (.tab)	Planned Run Audit (.pdf)
DNA Mapped BAM Index file (.bam.bai)	Basecaller command files (.json)
RNA Mapped BAM Index file (.bam.bai)	checksum file
DNA FASTQ File (.fastq)	Pipeline commands (<code>_pipeline.json</code>)
Base Caller Log File (.log)	Experimental log file (<code>_final.txt</code>)
Targeted CNV Detail Table (.tab) ^[1]	Wells with beads (<code>_beadogram.png</code>)
Non-targeted CNV Detail Table (.tab) ^[1]	Bead find stats file (.stats)
SNV/INDEL Detail Table (.tab)	Loading Density Figure (.png)

^[1] For use with IVD tests that include CNV reporting.

Pass/fail criteria and repeat strategy

For pass/fail criteria and repeat strategy, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.



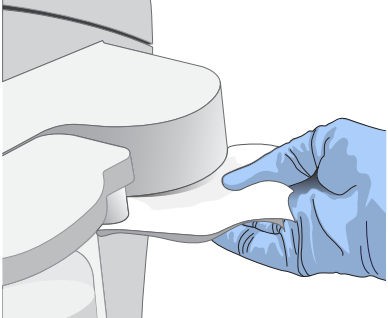
Troubleshooting

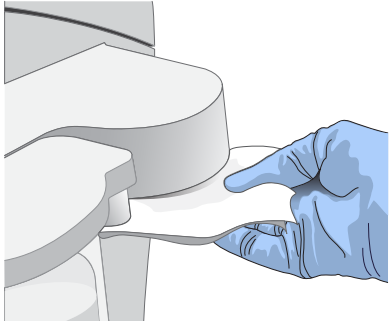
Ion PGM™ Dx Sequencer initialization

Observation	Possible cause	Recommended action
Pressure alarm	Gas cylinder may be turned off or empty	<ul style="list-style-type: none"> Verify that the cylinder has at least 500 PSI and 30 PSI at the outlet of the regulator. Confirm that all valves between the cylinder and the sequencer are open. Once you confirm gas pressure leading into the instrument, press Yes to retry verification of gas pressure. If the test continues to fail, contact Technical Support (see “Customer and technical support” on page 106).
Error message: Leak check failed, make sure the reagent tubes are not attached to the sequencer and check Wash 1, 2, or 3 for leak	<ul style="list-style-type: none"> Caps are not tightened on the Wash 1, 2, or 3 bottles. Bottles may be damaged or defective. 	<ol style="list-style-type: none"> Inspect all the bottles for damage or visible leaks. If a bottle appears damaged, replace it. Finger-tighten all the bottle caps, make sure that the reagent tubes are not attached to the instrument, and then press Retry. If leak check continues to fail, contact Technical Support (see “Customer and technical support” on page 106).
Error message: UNDERSHOT TARGET (instrument touchscreen may also be frozen)	Water used did not meet specifications, or insufficient amount of NaOH added	<p>If the touchscreen is frozen, reboot the sequencer and restart initialization. If you receive the same error and the touchscreen continues to freeze, contact Technical Support (see “Customer and technical support” on page 106). Otherwise, proceed to the following steps.</p> <ol style="list-style-type: none"> Press Retry to reattempt pH adjustment. If you receive the same error message after multiple attempts, press Abort. Check your water purification system and troubleshoot per the manufacturer's directions, or identify a new source of 18-MΩ water. Restart the initialization procedure and prepare new solutions, including a new aliquot of 100 mM NaOH. If the problem persists, contact Technical Support (see “Customer and technical support” on page 106).

Observation	Possible cause	Recommended action
Error message: UNDERSHOT TARGET (instrument touchscreen may also be frozen) <i>(continued)</i>	Auto-pH did not add enough Wash 1 Solution to the Wash 2 Solution before the maximum iterations occurred.	If the touchscreen is frozen, reboot the sequencer and restart initialization. If you receive the same error and the touchscreen continues to freeze, contact Technical Support (see “Customer and technical support” on page 106). Otherwise, proceed to the following steps. <ol style="list-style-type: none"> 1. A blockage may have occurred. See “Error message: There may be a blockage or no NaOH in W1. Check W1 and run line clear then try again” on page 73. 2. Press Retry to restart the pH check. If you still get the "Undershot target pH" error, replace the chip with a new (unused) chip and restart the pH check. <p>Note: The new chip can be used for sequencing after initialization completes.</p>
	Loose W1 sipper	Tighten the sipper and retry.
Error message: Please insert a chip and press Start	Instrument cannot detect the chip in chip socket.	<ol style="list-style-type: none"> 1. Open the chip clamp and remove the chip. 2. Check for debris under the chip and in the chip socket. <p>IMPORTANT! Never rub or wipe the socket. Rubbing the socket can damage it and cause it to fail.</p> <ol style="list-style-type: none"> 3. Look for liquid outside the flow cell of the chip: <div data-bbox="971 1213 1198 1352" data-label="Image"> </div> <ol style="list-style-type: none"> 4. If you see liquid, replace the chip with a new (unused) one. <p>Note: The new chip can be used for sequencing after initialization completes.</p> <ol style="list-style-type: none"> 5. Close the clamp, then press Start to restart the process. 6. If the new chip also fails, there could be a problem with the chip socket. Contact Technical Support (see “Customer and technical support” on page 106).

Observation	Possible cause	Recommended action
Error message: OVERSHOT TARGET	<ul style="list-style-type: none"> Wrong amount or concentration of NaOH was added to the Wash 1 Bottle Auto-pH added more NaOH from the Wash 1 Bottle to the Wash 2 Bottle than was needed 	Do not press Next . Prepare fresh reagents, then press Retry to retry the initialization. If the problem persists, contact Technical Support (see “Customer and technical support” on page 106).
	Clog in waste line(s)	Do not press Next . See “Error message: There may be a blockage or no NaOH in W1. Check W1 and run line clear then try again” on page 73.
	Clog in chip	Do not press Next . See “Error message: There may be a blockage or no NaOH in W1. Check W1 and run line clear then try again” on page 73. If the waste lines are not clogged, replace the chip, then click Retry .
W2 pH consistently undershoots target	pH of water is too low before any NaOH is added	<ul style="list-style-type: none"> Confirm that the 18 MΩ water supply meets specifications. If necessary, when preparing the Wash 2 Bottle, add more than the recommended 70 μL of 100 mM NaOH. After adding the NaOH, the Wash 2 Bottle must be in the range of pH 6.0–6.5 at first pH iteration before you begin initialization.
Error Message: Leak detected – Check dNTP Conical tubes	<ul style="list-style-type: none"> dNTP Reagent Tube seal is not tight. dNTP Reagent Tube may be damaged or defective. 	<ol style="list-style-type: none"> Remove and reinstall each dNTP Reagent Tube. Ensure that each tube is securely tightened (finger-tighten), then press OK to re-check pressure. If the error message persists, set up dNTPs in new tubes, secure new tubes on the instrument, then press OK. If leak check continues to fail, contact Technical Support (see “Customer and technical support” on page 106).

Observation	Possible cause	Recommended action
<p>Error message: There may be a blockage or no NaOH in W1. Check W1 and run line clear then try again</p>	<p>The waste lines may be blocked.</p>	<ol style="list-style-type: none"> 1. Remove the waste bottle and place lab wipes under the waste arm. 2. Gently wipe the waste arm with a lab wipe to clear liquid from around the waste line.  <ol style="list-style-type: none"> 3. Press Flow check one or more times to observe the flow rates from both lines. One line should drip slightly faster than the other. If one or both lines are blocked (no flow), or the drip rates are significantly different, go to the next step. If the flow rates are normal, see “Error message: There may be a blockage or no NaOH in W1 (continued from previous page)” on page 74 below. 4. Press Line Clear. Follow the prompts and use the syringe supplied with the Ion PGM™ Dx System. 5. After Line Clear, press Flow check, then check for normal flow rates from the waste lines. 6. If the flow rates are still not normal, perform Line Clear one more time. 7. If the line(s) remain blocked, contact Technical Support (see “Customer and technical support” on page 106). Otherwise, press Start to restart auto-pH.
	<p>Wash 1 or Wash 2 sipper may be loose</p>	<ol style="list-style-type: none"> 1. Loosen the Wash 1 cap and retighten the sipper. Since the gas flows when the cap is loose, tighten the sipper as quickly as possible. (The gas is not harmful to the NaOH solution and is not a hazard.) 2. Loosen the Wash 2 cap and retighten the sipper. Since the gas flows when the cap is loose, tighten the sipper as quickly as possible. (The gas is not harmful to the W2 Solution and is not a hazard.) 3. Press Start to restart the auto-pH process.

Observation	Possible cause	Recommended action
Error message: There may be a blockage or no NaOH in W1 (continued from previous page)	Forgot to add NaOH to the Wash 1 Bottle: Chip does not detect a large enough pH difference between the NaOH (W1) and W2 Solutions.	<ol style="list-style-type: none"> 1. If you forgot to add NaOH to the Wash 1 Bottle, loosen the cap and add 350 µL of 100 mM NaOH to the Wash 1 Bottle. (The flowing gas is not harmful to the NaOH solution and is not a hazard.) 2. Recap the bottle and shake gently to mix. 3. Press Start to restart auto-pH.
	Damaged chip	<ol style="list-style-type: none"> 1. Replace the chip with a new (unused) one. Insert the chip in the socket, then press Start. Note: The new chip can be used for sequencing after initialization completes. 2. If the error persists, there could be a problem with the chip clamp. Contact Technical Support (see “Customer and technical support” on page 106).
Error message: W2 average not stable. Try reseating/replacing chip	Reading for W2 solution is not stabilizing quickly enough	<ol style="list-style-type: none"> 1. Remove the waste bottle and gently wipe excess fluid from the waste lines with a lab wipe.  2. Check for leaks and reseat the chip. Replace the chip with a new (unused) one if needed. Note: The new chip can be used for sequencing after initialization completes. 3. Loosen the Wash 2 cap and retighten the sipper. Since the gas flows when the cap is loose, tighten the sipper as quickly as possible. (The gas is not harmful to the W2 Solution and is not a hazard.) 4. After performing one or more above steps, press Start to retry for auto-pH. If auto-pH fails even after replacing the chip, contact Technical Support (see “Customer and technical support” on page 106).
	The waste line may be blocked.	See “Error message: There may be a blockage or no NaOH in W1. Check W1 and run line clear then try again” on page 73.


Observation	Possible cause	Recommended action
Error message: W2 out of range	<ul style="list-style-type: none"> Chip measurements very unstable Chip is damaged 	See troubleshooting tips for "W2 average not stable" above.
Error message: Chip reading inconsistent. Please replace chip and try again.	<ul style="list-style-type: none"> pH response of the chip is not uniform or reliable Ran out of SEQ W3 Solution or volume too low 	<ol style="list-style-type: none"> Verify that there is enough SEQ W3 Solution (>25 mL) in the Wash 3 Bottle and that the sipper is secure. If necessary, loosen the Wash 3 Bottle cap, tighten the sipper, and add more SEQ W3 Solution to fill to 50 mL. Since the gas flows when the cap is loose, perform these operations as quickly as possible. (The gas is not harmful to the SEQ W3 Solution and is not a hazard.) If there is enough SEQ W3 Solution, replace the chip with a new (unused) one. Insert the chip in the socket, then press "re-try". <p>Note: The new chip can be used for sequencing after initialization completes.</p>
Error message: Added too much W1 to W2	<ul style="list-style-type: none"> Poor water quality 18-MΩ water exposed to air for too long Incorrect solution added to the SEQ W2 Solution Too little NaOH added to SEQ Wash 1 Bottle Damaged chip 	<ol style="list-style-type: none"> Check whether the water meets the 18-MΩ specification and that the 100 mM NaOH and SEQ W2 Solutions were added correctly. If solutions are incorrect or water does not meet specifications, correctly prepare the solution(s) and/or use 18-MΩ water. Abort the initialization and restart using correct solutions/water. If the solutions and water are correct, abort the initialization and try reinitializing with a different chip.
WARNING: AutoPH is within expected pH range, 0 milliliters of W1 was added to W2. Press "next" if expected, otherwise press "retry" to restart AutoPH	After auto-pH undershot the target, the user pressed the Retry button to restart auto-pH, but no additional NaOH was added to the Wash 2 bottle.	The pH reading was close enough to the target that no additional NaOH was added. Press Next to proceed with initialization.

Initialization: Reagent pH verification

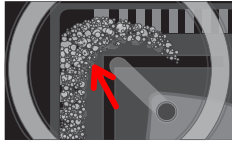
Observation	Possible cause	Recommended action
Failure screen	<ul style="list-style-type: none"> • One or more reagents are not within the target pH range • Chip is damaged 	<ol style="list-style-type: none"> 1. Press Start to repeat the pH measurement. 2. If the test still fails, replace the chip with a new (unused) chip and press Start to repeat. <p>Note: The new chip can be used for sequencing after initialization completes.</p> 3. If the test still fails with the new chip, clean and reinitialize the instrument with fresh reagents and a new chip. 4. If the test still fails, contact Technical Support (see “Customer and technical support” on page 106).

Chip calibration

Observation	Possible cause	Recommended action
Chip calibration failure (before sample has been loaded on the chip)	<ul style="list-style-type: none"> • Debris on the chip socket • Chip not seated correctly in the chip clamp • Chip is damaged • Problem with chip clamp or socket 	<ol style="list-style-type: none"> 1. Open the chip clamp, remove the chip, and check for damage, leaks, and/or debris under the chip and on the chip socket. <ul style="list-style-type: none"> • If debris or leaks are visible, gently dab the socket with a lab wipe tissue dampened with 18-MΩ water, then reseal the chip in the socket. • If no leaks or debris is visible, reseal the chip in the socket. <p>IMPORTANT! Never rub or wipe the socket. Rubbing the socket can damage it and cause it to fail.</p> 2. After reseating the chip, press Retry. 3. If the chip passes, press Next to start the experiment. If the chip still fails, reseal the chip again, then press Retry. 4. If chip calibration continues to fail with the same chip, retry with a new chip. 5. If the new chip fails as well: <ol style="list-style-type: none"> a. Press Abort to abort the run. <p>IMPORTANT! Be sure to abort the run before rebooting the sequencer in the subsequent steps.</p> b. Reboot the sequencer by holding down the Power button on the front to shut it down, then press again to restart. c. Restart the run with the new chip. 6. If the new chip continues to fail, there may be an issue with the chip socket. Contact Technical Support (see “Customer and technical support” on page 106).
Error message: Bad pixels error	Chip not seated correctly in the chip clamp.	<ol style="list-style-type: none"> 1. Make sure the chip is seated correctly and the clamp is closed. 2. Press down on the chip clamp with even pressure until the message instructing to close the lid appears. 3. If the new chip continues to fail, there may be an issue with the chip socket. Contact Technical Support (see “Customer and technical support” on page 106).

Observation	Possible cause	Recommended action
Leak of unknown origin during chip calibration (before sample has been loaded on the chip)	<ul style="list-style-type: none"> • Chip clamp not closed properly. • Chip is damaged. • Problem with the chip clamp or socket 	<ol style="list-style-type: none"> 1. Press Abort to immediately stop the run. Note: If the system is pressurized during abort, initiate the clean step or a run to depressurize the system and stop the flow of buffer, then abort the clean step. 2. Open the chip clamp, remove the chip, and gently dab the chip socket with a lab wipe tissue to absorb any fluid. IMPORTANT! Never rub or wipe the socket. Rubbing the socket can damage it and cause it to fail. 3. Make sure that the rubber gaskets on the chip clamp are properly installed and not loose or out of position. 4. Rinse the socket with 18-MΩ water and gently absorb most of the water with the lab wipe. 5. Repeat the rinse, then gently dab the chip socket with a lab wipe until dry. 6. Place a lab wipe on the grounding plate and dampen it with 18-MΩ water. Wipe the bottom of the chip on this wipe to remove salts from the chip contacts, and then gently dab the bottom of the chip with a dry lab wipe. 7. Remove the lab wipe, dry the grounding plate, and place the chip on it. Confirm that there is no condensation outside the chip flow cell:  8. If there is condensation or fluid, the chip is damaged and cannot be run. If no chip damage is apparent, reseal the chip and press Run to restart the experiment.

Observation	Possible cause	Recommended action
Leak of unknown origin during chip calibration (before sample has been loaded on the chip) <i>(continued)</i>		9. When prompted to install the chip, make certain that the chip clamp is fully closed. 10. If the chip leaks again, press Abort , clean the chip socket, then restart the run with a different chip. If the new chip leaks, it may indicate a problem with the chip clamp or socket. Contact Technical Support (see “Customer and technical support” on page 106).

Observation	Possible cause	Recommended action
Leak of unknown origin during chip calibration (after sample has been loaded on the chip)	<ul style="list-style-type: none"> • Chip clamp not closed properly • Chip is damaged. • Problem with the chip clamp or socket 	<ol style="list-style-type: none"> 1. Press Abort to immediately stop the run. Note: If the system is pressurized during abort, initiate the clean step to de-pressurize the system to stop the flow of buffer, then abort the clean step. 2. Open the chip clamp, remove the chip, and gently dab the chip socket with a lab wipe tissue to absorb any fluid. IMPORTANT! Never rub or wipe the socket. Rubbing the socket can damage it and cause it to fail. 3. Make sure that the rubber gaskets on the chip clamp are tight and in position. 4. Place a lab wipe on the grounding plate and dampen it with 18-MΩ water. Wipe the bottom of the chip on this wipe, and then gently dab with a dry lab wipe. 5. Dry the grounding plate with a lab wipe and place the chip on it. Confirm that there is no condensation outside the chip flow cell:  6. If there is condensation or fluid, the chip is damaged and cannot be run. Prepare a new run starting with template preparation (skip the following troubleshooting steps). 7. If no chip damage is apparent, you can attempt to rerun the loaded chip. Press Run on the Main Menu to start a new run. 8. When prompted, place the used chip from initialization in the chip clamp and press Next to clean the fluid lines. 9. When prompted, scan or enter the Planned Run Short Code. 10. When prompted, scan the top and bottom barcodes on the loaded chip. 11. Secure the loaded chip in the chip clamp and proceed with initial chip calibration.

Observation	Possible cause	Recommended action
<p>Leak of unknown origin during chip calibration (after sample has been loaded on the chip) <i>(continued)</i></p>		<p>12. If the chip leaks again, press Abort, then clean the chip socket. There may be a problem with the chip clamp or socket. Contact Technical Support (see “Customer and technical support” on page 106).</p> <p>13. If the chip does not leak and initial calibration passes, skip the chip loading steps and continue with the run.</p>
<p>Error message: Calibration failed (after sample has been loaded on the chip)</p>	<ul style="list-style-type: none"> • Debris on the chip socket • Chip clamp not closed properly • Chip is damaged • Problem with chip clamp or socket 	<ol style="list-style-type: none"> 1. Open the chip clamp, remove the chip, and check for damage, leaks, or debris under the chip and on the chip socket. <ul style="list-style-type: none"> • If debris or leaks are visible, gently dab the socket with a lab wipe tissue dampened with 18-MΩ water, then re-clamp the chip. • If no leaks or debris are visible, re-clamp the chip. <p>IMPORTANT! Never rub or wipe the socket. Rubbing the socket can damage it and cause it to fail.</p> 2. After re-clamping the chip, press Retry. 3. If the chip passes, press Next to start the experiment. If the chip still fails, re-clamp the chip and press Retry. <p>Note: You have a total of 3 attempts to retry the chip before aborting the run.</p> 4. If calibration continues to fail, there may be an issue with the loaded chip, chip clamp, or chip socket. To test the clamp and socket: <ol style="list-style-type: none"> a. Abort the run, then restart the run. b. Do not use the loaded chip. Instead, place a <i>used</i> chip in the chip clamp, then follow the steps for the initial (unloaded) chip calibration. <ul style="list-style-type: none"> • If the used chip fails initial calibration, there may be an issue with the clamp or socket. Abort the run and contact Technical Support (see “Customer and technical support” on page 106). • If the used chip passes calibration, the previously loaded chip may be bad. Abort the run, prepare fresh template, and run with a new chip.

Observation	Possible cause	Recommended action
Error message: Chip not seated correctly. Please reset the chip and click OK to continue	<ul style="list-style-type: none"> • Debris on the chip socket • Chip not seated correctly in the chip clamp • Chip is damaged • Problem with chip clamp or socket 	<ol style="list-style-type: none"> 1. Open the chip clamp, remove the chip, and check for damage, leaks, and/or debris under the chip and on the chip socket. <ul style="list-style-type: none"> • If debris or leaks are visible, gently dab the socket with a lab wipe tissue dampened with 18-MΩ water, then reseal the chip in the socket. • If no leaks or debris are visible, reseal the chip in the socket. <p>IMPORTANT! Never rub or wipe the socket. Rubbing the socket can damage it and cause it to fail.</p> 2. Press Retry. 3. If the chip passes, press Next to start the experiment. If the chip still fails, try reseating the chip again and pressing Retry. <p>Note: You have a total of 3 attempts to reseal the chip before aborting the run.</p> 4. If you continue to receive the "Chip not seated correctly" error message, contact Technical Support (see "Customer and technical support" on page 106). There may be an issue with the chip clamp or socket.
Chip calibration status bar does not progress	During chip calibration, the touchscreen status progress bar does not progress.	<ol style="list-style-type: none"> 1. Press the Abort button to abort the run. 2. Reboot the sequencer by holding down the Power button on the front to shut it down, then press the button again to restart. 3. Restart the run with the same chip. <p>Note: Reinitialization is <i>not</i> required after rebooting the sequencer if the initialization successfully completed before the run was aborted.</p>
Barcode on the bottom of the chip does not scan	The barcode is printed incorrectly.	Use a new chip and contact Technical Support for a replacement (see "Customer and technical support" on page 106).

Sample loading

Observation	Possible cause	Recommended action
Sample volume is <30 μ L If the sample volume is less than <30 μ L, there will be a visible air gap at the end of the pipette tip when you collect the sample for chip loading.	Volume was lost at some point during sample preparation.	<p>Note: Equilibrate SEQ Sample Buffer (brown cap) to room temperature for 15 minutes before performing the following steps.</p> <ol style="list-style-type: none"> 1. With the sample loaded in the pipette tip, dial down the pipette until the liquid reaches the end of the tip. Record the volume shown on the pipette (X). 2. Dispense the sample back into the sample tube. 3. Subtract the volume shown on the pipette from 30 μL ($30 - X = Y \mu$L) to determine the missing volume (Y). 4. Change the pipette tip, and add Y μL of SEQ Sample Buffer (brown cap) to the sample. 5. Pipet up and down 4 times to mix the contents, then continue loading the sample into the chip.
Large amount of the sample volume ($\geq 15 \mu$ L) leaks from the outlet port during chip loading.	The chip loading port has a clog or obstruction.	Do not continue with the sequencing run. Contact Technical Support.

Warnings and alarms—Ion PGM™ Dx System

The following warnings and alarms appear on the Ion PGM™ Dx Sequencer touchscreen and in the Torrent Suite™ Dx Software under the **Monitor** tab.

Observation	Possible cause	Recommended action
Display message: Pressure too high	Internal pressure regulator was not set correctly	Contact Technical Support (see “Customer and technical support” on page 106).
Display message: Pressure too low.	<ul style="list-style-type: none"> • Gas line is not connected to the instrument • Gas cylinder may be turned off or empty 	<ol style="list-style-type: none"> 1. Verify that the gas line is connected to the instrument. 2. Verify that the gas cylinder has at least 500 psi on the pressure gauge inside the tank and 30 psi on the gauge that regulates pressure to the outlet. (Both pressure gauges are mounted on the gas cylinder.) 3. Confirm that the outlet valve on the regulator is turned on. 4. If the problem persists, contact Technical Support (see “Customer and technical support” on page 106).



Observation	Possible cause	Recommended action
Display message: Instrument idle temperature too high	<ul style="list-style-type: none"> Room temperature is too high. Clogged filter or blocked airway on the instrument Hardware issue (fan is not running or running too slowly) 	<p>Note: The data created during a run with this alarm raised may still be used if all the QC metrics are met.</p> <p>See the recommended action for “Display message: Instrument temperature too high” on page 87.</p>
Display message: Instrument idle temperature too low	<ul style="list-style-type: none"> Ambient room temperature is below 20°C. Hardware issue 	Bring the ambient temperature up to 20°C. If the problem persists, contact Technical Support (see “Customer and technical support” on page 106).
Display message: Bad results data drive	<ul style="list-style-type: none"> On some machines, the warning appears before the reboot completes. There is a hardware issue. 	Wait for a few minutes to see if the error message disappears. If the error message disappears, data obtained during a run with this alarm raised can still be used. If the problem persists, contact Technical Support (see “Customer and technical support” on page 106).
Display message: Kernels do not match	Hardware and/or software issue	Contact Technical Support (see “Customer and technical support” on page 106).
Display message: Failed to set up system time at startup. Check your connection to the Ion Torrent™ Server.	The connection between the Ion PGM™ Dx Sequencer and the Ion Torrent™ Server has been lost.	<ol style="list-style-type: none"> Check the network connection to the Ion Torrent™ Server to make sure the connection is established, then reboot the instrument. If the problem persists, replace the network cable(s) to the instrument and server. If the problem persists, contact Technical Support (see “Customer and technical support” on page 106).
	Instrument is still in the process of establishing a connection	Allow 10 minutes to see if display message clears.
Display message: Log on failed. Could not authenticate the user.	Incorrect username or password entered by the user.	Enter the correct username and password.
	The connection between the instrument and the server has been lost.	Check the connection to the Ion Torrent™ Server, then reboot the Ion PGM™ Dx Sequencer. Ensure that no alarms involving the connection to the Ion Torrent™ Server appear.
Display message: Lost connection to the Ion Torrent™ Server	The connection between the instrument and the server has been lost.	Check the network connection to the Ion Torrent™ Server, and then reboot the Ion PGM™ Dx Sequencer. If this alarm appears during a run, the data created during that run can still be used.
Display message: UBoots do not match	Hardware issue	Contact Technical Support (see “Customer and technical support” on page 106).

Observation	Possible cause	Recommended action
Display message: Bad boot drive detected	Hardware issue	Contact Technical Support (see “Customer and technical support” on page 106). If this alarm appears during a run and data for the run is generated, that data may still be used.
Display message: Results drive not accessible. Reboot and try again.	<ul style="list-style-type: none"> On some machines, the warning appears before the reboot completes Hardware issue 	Wait for a few minutes to see if the error message disappears. If the error message appears and disappears during a run, data obtained during that run can still be used. If the alarm persists, contact Technical Support (see “Customer and technical support” on page 106).
Display message: Lost chip connection	The instrument cannot detect a chip in the chip clamp	See the instructions under “Chip calibration failure (before sample has been loaded on the chip)” on page 77.
Display message: Sensor unable to measure instrument temperature	Hardware issue	Contact Technical Support (see “Customer and technical support” on page 106).
Display message: Results drive check failed	Hardware issue	If the error message disappears when you return to the main instrument screen, this alarm can be ignored. Otherwise, contact Technical Support (see “Customer and technical support” on page 106).
Display message: Software versions incompatible. Go to the Options menu and update the software.	The system software was updated, but the instrument software was not.	<p>After the system software has been updated, update the instrument software as follows:</p> <ol style="list-style-type: none"> On the main menu of the instrument, press Options and follow the instructions to check for and install updates. When installation is complete, follow the onscreen prompts to restart the instrument. <p>IMPORTANT! You must restart the instrument before proceeding.</p>
Display message: Sensor unable to measure gas pressure. Check supply gas pressure.	<ul style="list-style-type: none"> Gas line is not connected to the instrument Gas cylinder may be turned off or empty Hardware issue 	<ol style="list-style-type: none"> Verify that the gas line is connected to the instrument. Verify that the gas cylinder has at least 500 psi on the pressure gauge inside the tank and 30 psi on the gauge that regulates pressure to the outlet. (Both pressure gauges are mounted on the gas cylinder.) Confirm that the outlet valve on the regulator is turned on. If the problem persists, contact Technical Support (see “Customer and technical support” on page 106).

Observation	Possible cause	Recommended action
Display message: Failed to locate the barcode scanner. Check if the scanner is attached.	The connection between the barcode scanner and the Ion PGM™ Dx Sequencer has been lost.	Make sure the scanner is plugged into a USB port on the instrument. If it is connected and the alarm still appears, try plugging the scanner into a second USB port. If the alarm persists, contact Technical Support (see “Customer and technical support” on page 106).
Display message: Failed to set the pressure to target range. Check the gas connection and try again.	<ul style="list-style-type: none"> Gas line is not connected to the instrument Gas cylinder may be turned off or empty Hardware issue (regulator malfunction) 	<ol style="list-style-type: none"> Verify that the gas line is connected to the instrument. Verify that the gas cylinder has at least 500 psi on the pressure gauge inside the tank and 30 psi on the gauge that regulates pressure to the outlet. (Both pressure gauges are mounted on the gas cylinder.) Confirm that the outlet valve on the regulator is turned on. If the problem persists, contact Technical Support (see “Customer and technical support” on page 106).
Display message: Failed to set up FTP connection. Check your connection to the Ion Torrent™ Server .	The network connection is not established or an incorrect IP address was used.	Confirm that the server information is correct for the Ion Torrent™ Server. Contact your local network administrator for support if the issue persists.
Display message: Unable to mount the file system	Hardware and/or software issue	<ol style="list-style-type: none"> Reboot the instrument to clear the alarm. If the alarm is not cleared after reboot, contact Technical Support (see “Customer and technical support” on page 106).
Display message: Instrument temperature too low	<ul style="list-style-type: none"> Room temperature is below 20°C. Hardware issue 	<p>Note: The data created during a run with this alarm raised may still be used if all the QC metrics are met.</p> <p>If the ambient room temperature is below 20°C, raise it. If the problem persists, contact Technical Support (see “Customer and technical support” on page 106).</p>

Observation	Possible cause	Recommended action
<p>Display message: Instrument temperature too high</p>	<ul style="list-style-type: none"> • Room temperature is too high. • Clogged filter or blocked airway on the instrument • Hardware issue (fan is not running or running too slowly) 	<p>Note: The data created during a run with this alarm raised may still be used if all the QC metrics are met.</p> <ol style="list-style-type: none"> 1. If the ambient room temperature is above 30°C, lower it. 2. Make sure that the round filter on the back panel of the instrument has unrestricted airflow. If the filter is clogged with dust, clean it as follows: <ol style="list-style-type: none"> a. Pinch the dirty filter with your fingers, then remove it from the instrument. <div data-bbox="1024 699 1414 1035" data-label="Image"> <p>The illustration shows a hand in a blue glove pinching a dark, circular filter. The filter is being held between the thumb and index finger, demonstrating the correct technique for removal from the instrument's back panel.</p> </div> <ol style="list-style-type: none"> b. Shake the filter over a waste container to remove most of the dust. c. Rinse the filter with running water to remove any remaining dust. The water flow should be from the inside-facing surface to the outside-facing surface through the filter. d. Air dry the filter. e. Blot any remaining dust from the filter using tape. f. Reinsert the filter. <p>If the problem persists, contact Technical Support (see “Customer and technical support” on page 106).</p>
<p>Display message: Chip temperature too low</p>	<p>Hardware issue</p>	<p>IMPORTANT! The data created during a run with this alarm raised should <i>not</i> be used.</p> <p>Contact Technical Support (see “Customer and technical support” on page 106).</p>



Observation	Possible cause	Recommended action
Display message: Chip temperature too high	<ul style="list-style-type: none"> Room temperature is too high. Clogged filter or blocked airway on the instrument Hardware issue (instrument fan is not running or running too slowly) 	<p>IMPORTANT! The data created during a run with this alarm raised should <i>not</i> be used.</p> <p>See the recommended action for “Display message: Instrument temperature too high” on page 87.</p>
Display message: Lost communication with valve board	Hardware issue	Contact Technical Support (see “Customer and technical support” on page 106).
Display message: Fan current too low	Hardware issue	Contact Technical Support (see “Customer and technical support” on page 106).
Display message: Heater current too low	Hardware issue	<p>IMPORTANT! If the chip temperature is also out of range, data created during a run should <i>not</i> be used.</p> <p>Contact Technical Support (see “Customer and technical support” on page 106). If no chip temperature alarms are raised, data created during a run may still be used if all the QC metrics are met.</p>
Error message: A non-recoverable error has occurred	Hardware issue	Prepare new template from the same library and plan a new run. If the issue persists, contact Technical Support (see “Customer and technical support” on page 106).

Sequencer software issues

(For additional software anomalies, see the release notes provided with your version of the software.)

Observation	Possible cause	Recommended action
Sequencer touchscreen is frozen	The user pressed multiple buttons on the screen in rapid succession.	Wait 5 minutes for the touchscreen to unfreeze. If the touchscreen remains frozen, reboot the sequencer by holding down the Power button on the front to shut it down, then press again to restart.
	The touchscreen is in a locked state.	Reboot the sequencer by holding down the Power button on the front to shut it down, then press again to restart.

Warnings and alarms—Veriti™ Dx 96-well Thermal Cycler

Observation	Possible cause	Recommended action
"Fatal Error" message displayed by Veriti™ Dx 96-well Thermal Cycler	Various	For assistance, contact Technical Support (see "Customer and technical support" on page 106). Refer to the <i>Veriti™ Dx 96-well Thermal Cycler User Guide</i> (Pub. No. 4453697) for general troubleshooting information for this instrument.



Ion PGM™ Dx Chip Minifuge and barcode scanner

Ion PGM™ Dx Chip Minifuge

The Ion PGM™ Dx Chip Minifuge is supplied with one custom rotor and two buckets. The buckets are designed to hold two chips: one in each bucket. The rotor and bucket design enables effective and efficient reagent loading of chips.



Safety precautions



CAUTION!

- Make sure your supply voltage matches the voltage label on the minifuge, i.e., never plug a 120V minifuge into an 220–240 VAC outlet. Operating the minifuge with a supply voltage outside the specified range may cause a fire or electric shock.
- Do not run the minifuge for more than 30 seconds.
- Never operate the minifuge without a rotor properly attached to the shaft.
- Never operate with only one chip in place. A chip must be present in each bucket to balance the rotor. If necessary, you can balance a loaded chip with a used chip of any type.
- Never put hands in the rotor area unless the rotor is completely stopped.
- Never move the minifuge while the rotor is spinning.
- Do not leave the minifuge running when not in use.

Note: The rotor can be balanced with a used chip from a previous reaction without risk of contamination.

Voltage selection

Two different minifuges are available, depending on your supply voltage: 120 VAC and 220–240 VAC. Make sure that the voltage specification on the label of your minifuge matches the supply voltage. If they do not match, change your supply voltage or contact Customer Support to request the appropriate minifuge.



CAUTION! Never plug a 120V minifuge into an 220–240 VAC outlet, or vice versa. Operating the minifuge with a supply voltage outside of the range specified on the label may cause a fire or electric shock.

Voltage, RPM, and RCF

The following tables list the revolutions per minute (RPM) and relative centrifugal force (RCF) at different voltages.

120/50 VAC, 60 Hz	RPM	RCF
90	4100	836
100	4550	1030
110	4960	1224
120	5330	1424
130	5710	1628

230/50 VAC, 60 Hz	RPM	RCF
210	5070	1279
220	5310	1403
230	5515	1513
240	5705	1619
250	5900	1732

Operation

1. Place the Ion PGM™ Dx Chip Minifuge on a level, clean surface near an accessible power outlet so that the cord and outlet are within easy reach of the operator.
2. Make sure the power switch on the minifuge is in the "off" position.
3. Load a chip into each bucket.

IMPORTANT! A chip must be present in each bucket to balance the rotor. If necessary, you can balance a loaded chip with a used chip of any type.

4. Turn the power switch on.
5. To begin centrifugation, close the lid of the minifuge. (The centrifugation time will vary depending on the step in the chip-loading protocol.)
6. To stop centrifugation, press down on the lid release tab on the front of the minifuge.



CAUTION! Do not attempt to open the lid or remove the chips until the unit has come to a complete stop.

7. After the rotor has stopped, open the lid by grabbing it with the thumb on the front and fingers on the back, then lifting the lid back on the hinge.

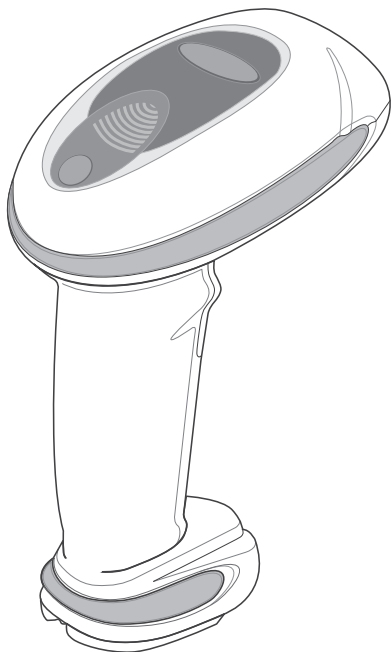
Cleaning

To clean the minifuge, use a damp cloth and a mild, noncorrosive detergent (pH <8). After cleaning, ensure that all parts are dried thoroughly before attempting to operate the unit. Do not immerse the centrifuge in liquid or pour liquids over it.

Note: Use only the cleaning protocol described above.

Barcode scanner

The barcode scanner provided with the Ion PGM™ Dx System uses a low-power, visible-light diode.



CAUTION! As with any bright light source, you should avoid staring directly into the light beam or shining the beam into other people's eyes. Momentary exposure to a Class 2 laser is not known to be harmful.



CAUTION! Use of controls, adjustments, or performance of procedures other than those specified in this guide can result in hazardous laser light exposure.

The barcode scanner specifications are listed below.

Wavelength	Rated Power
630–680 nm	1 mW



WARNING! GENERAL SAFETY. Using this product in a manner not specified in the user documentation may result in personal injury or damage to the instrument or device. Ensure that anyone using this product has received instructions in general safety practices for laboratories and the safety information provided in this document.

- Before using an instrument or device, read and understand the safety information provided in the user documentation provided by the manufacturer of the instrument or device.
- Before handling chemicals, read and understand all applicable Safety Data Sheets (SDSs) and use appropriate personal protective equipment (gloves, gowns, eye protection, and so on). To obtain SDSs, see the “Documentation and Support” section in this document.

Chemical safety



WARNING! GENERAL CHEMICAL HANDLING. To minimize hazards, ensure laboratory personnel read and practice the general safety guidelines for chemical usage, storage, and waste provided below. Consult the relevant SDS for specific precautions and instructions:

- Read and understand the Safety Data Sheets (SDSs) provided by the chemical manufacturer before you store, handle, or work with any chemicals or hazardous materials. To obtain SDSs, see the “Documentation and Support” section in this document.
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing).
- Minimize the inhalation of chemicals. Do not leave chemical containers open. Use only with adequate ventilation (for example, fume hood).
- Check regularly for chemical leaks or spills. If a leak or spill occurs, follow the manufacturer's cleanup procedures as recommended in the SDS.
- Handle chemical wastes in a fume hood.
- Ensure use of primary and secondary waste containers. (A primary waste container holds the immediate waste. A secondary container contains spills or leaks from the primary container. Both containers must be compatible with the waste material and meet federal, state, and local requirements for container storage.)
- After emptying a waste container, seal it with the cap provided.
- Characterize (by analysis if necessary) the waste generated by the particular applications, reagents, and substrates used in your laboratory.
- Ensure that the waste is stored, transferred, transported, and disposed of according to all local, state/provincial, and/or national regulations.
- **IMPORTANT!** Radioactive or biohazardous materials may require special handling, and disposal limitations may apply.

Biological hazard safety





WARNING! BIOHAZARD. Biological samples such as tissues, body fluids, infectious agents, and blood of humans and other animals have the potential to transmit infectious diseases. Conduct all work in properly equipped facilities with the appropriate safety equipment (for example, physical containment devices). Safety equipment can also include items for personal protection, such as gloves, coats, gowns, shoe covers, boots, respirators, face shields, safety glasses, or goggles. Individuals should be trained according to applicable regulatory and company/ institution requirements before working with potentially biohazardous materials. Follow all applicable local, state/provincial, and/or national regulations. The following references provide general guidelines when handling biological samples in laboratory environment.

- U.S. Department of Health and Human Services, *Biosafety in Microbiological and Biomedical Laboratories (BMBL)*, 5th Edition, HHS Publication No. (CDC) 21-1112, Revised December 2009; found at:
<https://www.cdc.gov/labs/pdf/CDC-BiosafetymicrobiologicalBiomedicalLaboratories-2020-P.pdf>
- World Health Organization, *Laboratory Biosafety Manual*, 3rd Edition, WHO/CDS/CSR/LYO/2004.11; found at:
www.who.int/publications/i/item/9789240011311

Symbols on this instrument

Symbols may be found on the instrument to warn against potential hazards or convey important safety information. In this document, the hazard symbol is used along with one of the following user attention words:

- **CAUTION!**—Indicates a potentially hazardous situation that, if not avoided, may result in minor or moderate injury. It may also be used to alert against unsafe practices.
- **WARNING!**—Indicates a potentially hazardous situation that, if not avoided, could result in death or serious injury.
- **DANGER!**—Indicates an imminently hazardous situation that, if not avoided, will result in death or serious injury.

Symbol	English	Français
	Caution, risk of danger Consult the manual for further safety information.	Attention, risque de danger Consulter le manuel pour d'autres renseignements de sécurité.
	Caution, risk of electrical shock	Attention, risque de choc électrique






(continued)

Symbol	English	Français
	Caution, piercing hazard	Attention, danger de perforation
	Caution, hot surface	Attention, surface chaude
	Potential biohazard	Danger biologique potentiel
	On	On (marche)
	Off	Off (arrêt)
	Earth (ground) terminal	Borne de (mise à la) terre
	Protective conductor terminal (main ground)	Borne de conducteur de protection (mise à la terre principale)
	Terminal that can receive or supply alternating current or voltage	Borne pouvant recevoir ou envoyer une tension ou un courant de type alternatif
	Do not dispose of this product in unsorted municipal waste CAUTION! To minimize negative environmental impact from disposal of electronic waste, do not dispose of electronic waste in unsorted municipal waste. Follow local municipal waste ordinances for proper disposal provision and contact customer service for information about responsible disposal options.	Ne pas éliminer ce produit avec les déchets usuels non soumis au tri sélectif. MISE EN GARDE ! Pour minimiser les conséquences négatives sur l'environnement à la suite de l'élimination de déchets électroniques, ne pas éliminer ce déchet électronique avec les déchets usuels non soumis au tri sélectif. Se conformer aux ordonnances locales sur les déchets municipaux pour les dispositions d'élimination et communiquer avec le service à la clientèle pour des renseignements sur les options d'élimination responsable.






















Conformity symbols on the instrument

Conformity mark	Description
	Indicates conformity with safety requirements for Canada and U.S.A.
	Indicates conformity with European Union requirements for safety and electromagnetic compatibility.
	Indicates conformity with Australian standards for electromagnetic compatibility.



Medical device symbols





The following table describes symbols that may be displayed on product labels. The symbols that are used on labels conform to standards BS EN ISO 15223-1:2016 and FDA 21 CFR 809.10 "Labeling for in vitro diagnostic products".

Symbol	Description	Symbol	Description
	MANUFACTURER		CONTAINS SUFFICIENT FOR <n> TESTS
	DATE OF MANUFACTURE		USE BY
	BATCH CODE		CATALOG NUMBER
	SERIAL NUMBER		FRAGILE, HANDLE WITH CARE
	LOWER LIMIT OF TEMPERATURE		PROTECT FROM LIGHT
	UPPER AND LOWER LIMITS OF TEMPERATURE		UPPER LIMIT OF TEMPERATURE
	DO NOT REUSE		BIOLOGICAL RISKS
	CAUTION, CONSULT ACCOMPANYING DOCUMENTS		CONSULT INSTRUCTIONS FOR USE
	UPPER AND LOWER LIMITS OF HUMIDITY		OBSERVE PRECAUTIONS FOR HANDLING ELECTROSTATIC SENSITIVE DEVICES
	IN VITRO DIAGNOSTIC MEDICAL DEVICE		



Safety alerts on this instrument

Additional text may be used with one of the symbols described above when more specific information is needed to avoid exposure to a hazard. See the following table for safety alerts found on the instrument.

English	Français
 CAUTION! Hazardous chemicals. Read the Safety Data Sheets (SDSs) before handling.	 MISE EN GARDE ! Produits chimiques dangereux. Lire les fiches signalétiques (FS) avant de manipuler les produits.
 CAUTION! Hazardous waste. Refer to SDS(s) and local regulations for handling and disposal.	 MISE EN GARDE ! Déchets dangereux. Lire les fiches signalétiques (FS) et la réglementation locale associées à la manipulation et à l'élimination des déchets.

Safety information for third-party instruments

Refer to the manufacturer's documentation for information on the safe use of third-party products provided with the instrument system.

Instrument safety

General



CAUTION! Do not remove instrument protective covers. If you remove the protective instrument panels or disable interlock devices, you may be exposed to serious hazards including, but not limited to, severe electrical shock, laser exposure, crushing, or chemical exposure.



CAUTION! Solvents and Pressurized fluids. Wear eye protection when working with any pressurized fluids. Use caution when working with any polymeric tubing that is under pressure:

- Extinguish any nearby flames if you use flammable solvents.
- Do not use polymeric tubing that has been severely stressed or kinked.
- Do not use polymeric tubing with tetrahydrofuran or nitric and sulfuric acids.
- Be aware that methylene chloride and dimethyl sulfoxide cause polymeric tubing to swell and greatly reduce the rupture pressure of the tubing.
- Be aware that high solvent flow rates (~40mL/min) may cause a static charge to build up on the surface of the tubing and electrical sparks may result.



Electrical



WARNING! Ensure appropriate electrical supply. For safe operation of the instrument:

- Plug the system into a properly grounded receptacle with adequate current capacity.
- Ensure the electrical supply is of suitable voltage.
- Never operate the instrument with the ground disconnected. Grounding continuity is required for safe operation of the instrument.



WARNING! Power Supply Line Cords. Use properly configured and approved line cords for the power supply in your facility.



WARNING! Disconnecting Power. To fully disconnect power, either detach or unplug the power cord. Position the instrument such that the power cord is accessible.



WARNING! Radio interference. This equipment has been designed and tested to CISPR 11 Class A. In a domestic environment it may cause radio interference, in which case you may need to take measures to mitigate the interference.



WARNING! Electromagnetic radiation. Do not use this device in close proximity to sources of strong electromagnetic radiation (e.g., unshielded intentional RF sources), as these may interfere with proper operation.

Cleaning and decontamination



CAUTION! Cleaning and Decontamination. Use only the cleaning and decontamination methods specified in the manufacturer's user documentation. It is the responsibility of the operator (or other responsible person) to ensure the following requirements are met:

- No decontamination or cleaning agents are used that could cause a HAZARD as a result of a reaction with parts of the equipment or with material contained in the equipment.
- The instrument is properly decontaminated a) if hazardous material is spilled onto or into the equipment, and/or b) prior to having the instrument serviced at your facility or sending the instrument for repair, maintenance, trade-in, disposal, or termination of a loan (decontamination forms may be requested from customer service).
- Before using any cleaning or decontamination methods (except those recommended by the manufacturer), users should confirm with the manufacturer that the proposed method will not damage the equipment.



Laser



CAUTION! LASER HAZARD, Bar Code Scanner. The bar code scanner included with the instrument system is a Class 2 laser. To avoid damage to eyes, do not stare directly into the beam or point into another person's eyes.

Gas safety

Verify that your installation room can accommodate gas cylinders.



WARNING! Instrumentation must be installed and operated in a well-ventilated environment as defined as having a minimum airflow of 6-10 air changes per hour. Assess the need for ventilation or atmospheric monitoring to avoid asphyxiation accidents from inert gases and/or oxygen depletion, and take measures to clearly identify potentially hazardous areas through training or signage. Please contact your Environmental Health and Safety Coordinator to confirm that the instruments will be installed and operated in an environment with sufficient ventilation.



WARNING! Pressurized gas cylinders are potentially explosive. Always cap the gas cylinder when it is not in use, and attach it firmly to the wall or gas cylinder cart with approved brackets or chains.



WARNING! Gas cylinders are heavy and may topple over, potentially causing personal injury and tank damage. Cylinders should be firmly secured to a wall or work surface. Please contact your Environmental Health and Safety Coordinator for guidance on the proper installation of a gas cylinder.

Safety and electromagnetic compatibility (EMC) standards

The instrument design and manufacture complies with the following standards and requirements for safety and electromagnetic compatibility.

Safety

Reference	Description
EU Directive 2006/95/EC	European Union "Low Voltage Directive"
IEC 61010-1 EN 61010-1 UL 61010-1 CSA C22.2 No. 61010-1	<i>Safety requirements for electrical equipment for measurement, control, and laboratory use – Part 1: General requirements</i>



(continued)

Reference	Description
IEC 61010-2-010 EN 61010-2-010	<i>Safety requirements for electrical equipment for measurement, control and laboratory use – Part 2-010: Particular requirements for laboratory equipment for the heating of materials</i>
IEC/EN 61010-2-101	<i>Safety requirements for electrical equipment for measurement, control and laboratory use - Part 2-101: Particular requirements for in vitro diagnostic (IVD) medical equipment</i>

EMC

Reference	Description
Directive 2004/108/EC	European Union “EMC Directive”
EN 61326-1	<i>Electrical Equipment for Measurement, Control and Laboratory Use – EMC Requirements – Part 1: General Requirements</i>
EN 61326-2-6	<i>Electrical Equipment for Measurement, Control and Laboratory Use – EMC Requirements – Part 26: Particular requirements – In vitro diagnostic (IVD) medical equipment)requirements</i>
FCC Part 15	U.S. Standard “Industrial, Scientific, and Medical Equipment”
AS/NZS CISPR 22:2009	<i>Limits and Methods of Measurement of Electromagnetic Disturbance Characteristics of Industrial, Scientific, and Medical (ISM) Radiofrequency Equipment</i>
ICES-003, Issue 5	<i>Industrial, Scientific and Medical (ISM) Radio Frequency Generators</i>

Environmental design

Reference	Description
Directive 2012/19/EU	European Union “WEEE Directive” – Waste electrical and electronic equipment
Directive 2011/65/EU	European Union “RoHS Directive” – Restriction of hazardous substances in electrical and electronic equipment

Precaution—strong magnet

Note: Do not substitute non-IVD labeled magnets for the DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet, provided with Ion PGM™ Dx System.

The DynaMag™ Dx 96-Well Plate Magnet and DynaMag™ Dx 16 2-mL Magnet contain very strong permanent magnets. People wearing a pacemaker or any other medical magnetized implant should not use this product unless advised by a health professional; the implant could be affected or damaged by exposure to a strong magnetic field. Keep tools and objects that could be damaged by the magnetic



field out of the working area. This includes, but is not restricted to, credit cards and other products containing magnetic recording devices. Keep away from delicate instruments, watches, electronic equipment, displays and monitors. The magnet may attract steel or other magnetic material with high mechanical forces. Take care during handling. Avoid contact between two magnets. Do not pull the magnets apart if contact has been made; twist off to prevent damage to the unit or fingers. The Health and Safety Officer should take all necessary steps and full responsibility to ensure that the precautions and statements are followed and adhered to.



Performance characteristics

For performance characteristics of the Oncomine™ Dx Target Test Kit, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

For performance characteristics of the Ion PGM™ Dx System, see the *Ion PGM™ Dx System Performance Characteristics User Guide* (Pub. No. MAN0018763).



Instrument warranty

For new Ion Torrent™ instruments, Life Technologies warrants to and only to buyer for twelve (12) months from the date of shipping, that the Ion Torrent™ software and Ion Torrent™ instruments are free from defects in material and workmanship and conform to Life Technologies' published specifications in all material respects. Where a valid and timely claim in respect of breach of Ion Torrent™ warranty is submitted to Life Technologies, Life Technologies may, at its discretion, replace, repair or modify the Ion Torrent™ instrument. Any agreed replacement shall be at 1:1, like-kind basis, at no cost to the buyer. For Ion Torrent™ chips or reagents reasonably determined by Life Technologies to be defective, independent of user error, shall be replaced by Life Technologies on a 1:1, like-kind basis at no cost to buyer, provided that such defective Ion Torrent™ chips or reagents were used by buyer prior to their expiration date, or if there is no expiration date, the Ion Torrent™ chips or reagents were used within six (6) months of receipt, and the defect was promptly reported with appropriate detail to Life Technologies' technical support.

NO OTHER WARRANTIES SHALL BE APPLICABLE TO ION TORRENT PRODUCTS (WHETHER OR NOT ANY FURTHER WARRANTY DOCUMENTATION MAY BE INCLUDED IN THE SHIPMENT), WITH THE EXCEPTION OF THIRD PARTY WARRANTIES WITH RESPECT TO THIRD PARTY PRODUCT. ANY THIRD PARTY PRODUCTS ARE NOT COVERED BY THIS SECTION AND ANY WARRANTIES FOR THIRD PARTY PRODUCTS ARE PROVIDED BY THE ORIGINAL MANUFACTURER OF THE THIRD PARTY PRODUCT. Warranties are made only to buyer purchasing the Ion Torrent™ Product directly from Life Technologies, are not transferable and do not extend to the benefit of any other person or entity, unless otherwise expressly stated in writing by Life Technologies. ANY PRODUCT NOT COVERED BY AN EXPRESS WRITTEN WARRANTY IS SOLD AND PROVIDED "AS IS," WITHOUT WARRANTY OF ANY KIND, STATUTORY, EXPRESS OR IMPLIED. Any description of Ion Torrent™ Product recited in Life Technologies' quotation is for the sole purpose of identifying Ion Torrent™ Product, and any such description is not part of any contract between Life Technologies and buyer and does not constitute a warranty that Ion Torrent™ Product shall conform to that description. Any sample or model used in connection with Life Technologies' quotation is for illustrative purposes only, and is not part of any contract between Life Technologies and buyer and does not constitute a warranty that Ion Torrent™ Product will conform to the sample or model. No affirmation of fact or promise made by Life Technologies, whether or not in Life Technologies' quotation, shall constitute a warranty that Ion Torrent™ Product will conform to the affirmation or promise. Unless otherwise specified in writing in documentation shipped with Ion Torrent™ Product or otherwise agreed by Life Technologies in writing. Life Technologies does not provide service or support for custom products or other products made to buyer's specifications. THE WARRANTIES IDENTIFIED IN THIS CLAUSE ARE LIFE TECHNOLOGIES' SOLE AND EXCLUSIVE WARRANTIES WITH RESPECT TO Ion Torrent™ PRODUCT AND ARE IN LIEU OF ALL OTHER WARRANTIES, STATUTORY, EXPRESS OR IMPLIED, ALL OF WHICH OTHER WARRANTIES ARE EXPRESSLY DISCLAIMED, INCLUDING WITHOUT LIMITATION ANY IMPLIED WARRANTY OF MERCHANTABILITY, FITNESS FOR A PARTICULAR PURPOSE, NON-INFRINGEMENT, OR REGARDING RESULTS OBTAINED THROUGH THE USE OF ANY PRODUCT (INCLUDING, WITHOUT LIMITATION, ANY CLAIM OF INACCURATE, INVALID OR INCOMPLETE RESULTS), WHETHER ARISING FROM A STATUTE OR OTHERWISE IN LAW OR FROM A COURSE OF PERFORMANCE, DEALING OR USAGE OF TRADE.

Customer and technical support

Visit [thermofisher.com/support](https://www.thermofisher.com/support) for the latest in services and support, including:

- Worldwide contact telephone numbers
- Product support
- Order and web support
- Safety Data Sheets (SDSs; also known as MSDSs)

Additional product documentation, including user guides and Certificates of Analysis, are available by contacting Customer Support.

Obtaining Certificates of Analysis

The Certificate of Analysis provides detailed quality control and product qualification information for each product. Certificates of Analysis are printed and shipped with the product.

Obtaining Certificates of Conformance

The Certificate of Conformance provides information on conformance testing of each instrument provided with the system. Certificates of Conformance are shipped with the instrument, and are also available by contacting Customer Support at [thermofisher.com/support](https://www.thermofisher.com/support).

Oncomine™ Dx Target Test Part V: Torrent Suite™ Dx Software 5.12.5 Reference

USER GUIDE

Publication Number MAN0018707

Revision B.0

IVD

For In Vitro Diagnostic Use.

ThermoFisher
S C I E N T I F I C



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Block 33 |
Marsiling Industrial Estate Road 3 |
#07-06, Singapore 739256



Life Technologies Corporation |
7335 Executive Way |
Frederick, MD 21704 | USA

Products manufactured in Singapore:

Ion PGM™ Dx Instrument System
Ion PGM™ Dx Sequencer
Ion OneTouch™ Dx Instrument
Ion OneTouch™ ES Dx Instrument
Ion PGM™ Dx Chip Minifuge (120V)
Ion PGM™ Wireless Scanner
Ion Torrent™ Server
Torrent Suite™ Dx Software
Veriti™ Dx 96-well Thermal Cycler, 0.2 mL

Products manufactured in Frederick:

Oncomine™ Dx Target Test Kit
Ion Torrent Dx FFPE Sample Preparation Kit
Ion PGM™ Dx Library Kit
Ion OneTouch™ Dx Template Kit
Ion PGM™ Dx Sequencing Kit
Ion 318™ Dx Chip
Ion OneTouch™ Rack Kit
DynaMag™ Dx 96-Well Plate Magnet
DynaMag™ Dx 16 2-mL Magnet

The information in this guide is subject to change without notice.

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Revision history: Pub. No. MAN0018707

Revision	Date	Description
B.0	3 September 2021	<ul style="list-style-type: none"> Updated antivirus software recommendations Minor corrections and formatting updates.
A.0	11 February 2021	New Oncomine™ Dx Target Test user guide for FDA submission.


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About this guide

Purpose of this guide

This user guide provides instructions for using Torrent Suite™ Dx Software to analyze sequencing data generated from sample libraries prepared using the Oncomine™ Dx Target Test Kit.

Oncomine™ Dx Target Test Kit user guides

This user guide is part of a five-guide set.

- *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*
- *Oncomine™ Dx Target Test Part II: Sample and Library Preparation User Guide*
- *Oncomine™ Dx Target Test Part III: Template Preparation User Guide*
- *Oncomine™ Dx Target Test Part IV: Sequencing and Results Reports User Guide*
- *Oncomine™ Dx Target Test Part V: Torrent Suite™ Dx Software 5.12.5 Reference User Guide*

All five guides are required to complete the entire Oncomine™ Dx Target Test workflow.

Note: The procedures in these guides supersede the instructions in the *Ion PGM™ Dx System User Guide* when using the Ion PGM™ Dx System with the Oncomine™ Dx Target Test.



Product information

Product description

Oncomine™ Dx Target Test

The Oncomine™ Dx Target Test is an *in vitro* diagnostic next-generation sequencing test to detect somatic alterations in human DNA and RNA isolated from non-small cell lung cancer (NSCLC) and cholangiocarcinoma (CC) tumor specimens in formalin-fixed, paraffin-embedded (FFPE) tissue samples. Detection of these variants is performed using the Ion PGM™ Dx System.

For a complete product description of the Oncomine™ Dx Target Test, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Software system

Torrent Suite™ Dx Software is a software system for managing samples and libraries, as well as creating, executing, and analyzing templating and sequencing runs on the Ion PGM™ Dx System. The Oncomine™ Dx Target Test assay is a locked assay in the software that provides configuration and analysis settings for using the Oncomine™ Dx Target Test Kit.

Intended use

For the Intended Use statement for the Oncomine™ Dx Target Test, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Theory of operation

For a complete description of the Theory of Operation of the system, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

Software description

Torrent Suite™ Dx Software is hosted on the Ion Torrent™ Server, part of the Ion PGM™ Dx System.

The software is used with the OncoPrint™ Dx Target Test Assay Definition File and the Ion PGM™ Dx Instrument System control software to support the system workflow from sample preparation through library preparation, template preparation, sequencing, primary and secondary analysis, and generation of OncoPrint™ Dx Target Test reports.

Software component	Main functions
Torrent Suite™ Dx Software	<ul style="list-style-type: none"> Generates base calls and quality values (primary analysis) Filters and trims reads to ensure high sequence accuracy Performs alignments and variant calling and generates reports using parameters in the OncoPrint™ Dx Target Test Assay Definition File (secondary analysis) Hosts the Torrent Suite™ Dx Software web application
OncoPrint™ Dx Target Test Assay Definition File	Installed with the software, this file contains gene lists and parameters for sequencing and analysis of specific DNA and RNA variants targeted by the OncoPrint™ Dx Target Test Kit.
Ion OneTouch™ Dx Instrument control software	<ul style="list-style-type: none"> Provides control of the Ion OneTouch™ Dx Instrument via a touchscreen interface Enables users to prepare template-positive Ion PGM™ Dx Ion Sphere™ Particles using the instrument Enables users to perform instrument maintenance
Ion PGM™ Dx Sequencer control software	<ul style="list-style-type: none"> Provides control of the Ion PGM™ Dx Sequencer via a touchscreen interface Enables users to perform sequencing workflows using the instrument Collects and compresses sequencing data, then transfers data to the Ion Torrent™ Server for primary and secondary analysis Enables users to perform instrument maintenance

Features of Torrent Suite™ Dx Software

Torrent Suite™ Dx Software enables users to:

- Generate reports on detected sequence variations in DNA and RNA in genes targeted by the Oncomine™ Dx Target Test
- Enter and manage information about samples and create libraries and library batches
- Plan runs to be executed on the Ion PGM™ Dx System instruments
- Monitor the progress of active instrument runs
- View and download run results and analysis reports
- View QC settings
- View workflow settings
- Manage user information and privileges
- View audit trails
- Configure archiving, reporting, and other administrative functions

Note: This guide provides instructions for using the Torrent Suite™ Dx Software in IVD Mode only.

Limitations and Precautions


Torrent Suite™ Dx Software has the following limitations:

- User names can contain alphanumeric characters and underscores, periods, and hyphens. Passwords can only contain alphanumeric characters.
- Torrent Suite™ Dx Software is intended to be used only with the Ion PGM™ Dx System and does not accept any software plugins.

Software compatibility and requirements

The procedures in this guide are designed for use with Torrent Suite™ Dx Software version 5.12.5 or later. For a complete description of software compatibility and requirements, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

About screen

Under the  tab, the **About** screen contains following links:

- **Support Contact Information** is a link to thermofisher.com/technical-resources, where you can access technical support and product information.
- **Instrument Diagnostics** opens a page that lists the active sequencers connected to the Ion Torrent™ Server. For each sequencer, you can click on additional links to download or open the following files:
 - Diagnostic archive file (ZIP file)
 - Init.log file
 - Installation Acceptance Report (PDF file)
- **End User License Agreement** opens the End User License Agreement (EULA) in PDF format.



Get started with Torrent Suite™ Dx Software

User-access levels

Users at this level...	Can...
Operator	<ul style="list-style-type: none">• Add and import samples• Prepare library batches• Create and save Planned Runs• Execute Planned Runs• Monitor runs• View results and reports
Manager	Operator functions plus: <ul style="list-style-type: none">• Create and edit sample attributes• Delete Planned Runs• Approve reports• Amend reports• Manage reference sequences and panel, hotspot, and other sequence files• Access services information
Administrator	Operator and Manager functions plus: <ul style="list-style-type: none">• View, export, and print audit records• Configure network settings• View and manage software updates• Configure data archive and storage settings• Manage instrument and software log files• Add and manage users

Ion PGM™ Dx System synchronization requirement

Allow up to 20 minutes for synchronization with the Ion Torrent™ Server after you power on any of the following instruments or the server itself:

- Ion PGM™ Dx Sequencer
- Ion OneTouch™ Dx Instrument

If you attempt to use an instrument before synchronization is complete, it will generate an alarm.

Network and password security requirements

Network configuration and security

The network configuration and security settings of your laboratory or facility (such as firewalls, anti-virus software, network passwords) are the sole responsibility of your facility administrator, IT, and security personnel. This product does not provide any network or security configuration files, utilities, or instructions.

If external or network drives are connected to the software, it is the responsibility of your IT personnel to ensure that such drives are configured and secured correctly to prevent data corruption or loss. It is the responsibility of your facility administrator, IT, and security personnel to prevent the use of any unsecured ports (such as USB, Ethernet) and ensure that the system security is maintained.

Password security

Thermo Fisher Scientific strongly recommends that you maintain unique passwords for all accounts in use on this product. All passwords should be reset upon first sign in to the product. Change passwords according to your organization's password policy.

It is the sole responsibility of your IT personnel to develop and enforce secure use of passwords.

Antivirus software

Thermo Fisher Scientific has tested Torrent Suite™ Dx Software with the following antivirus software products and found them compatible as antivirus solutions:

- Bitdefender GravityZone™ Business Security
- Kaspersky™ Endpoint Security for Linux
- Symantec™ Endpoint Protection

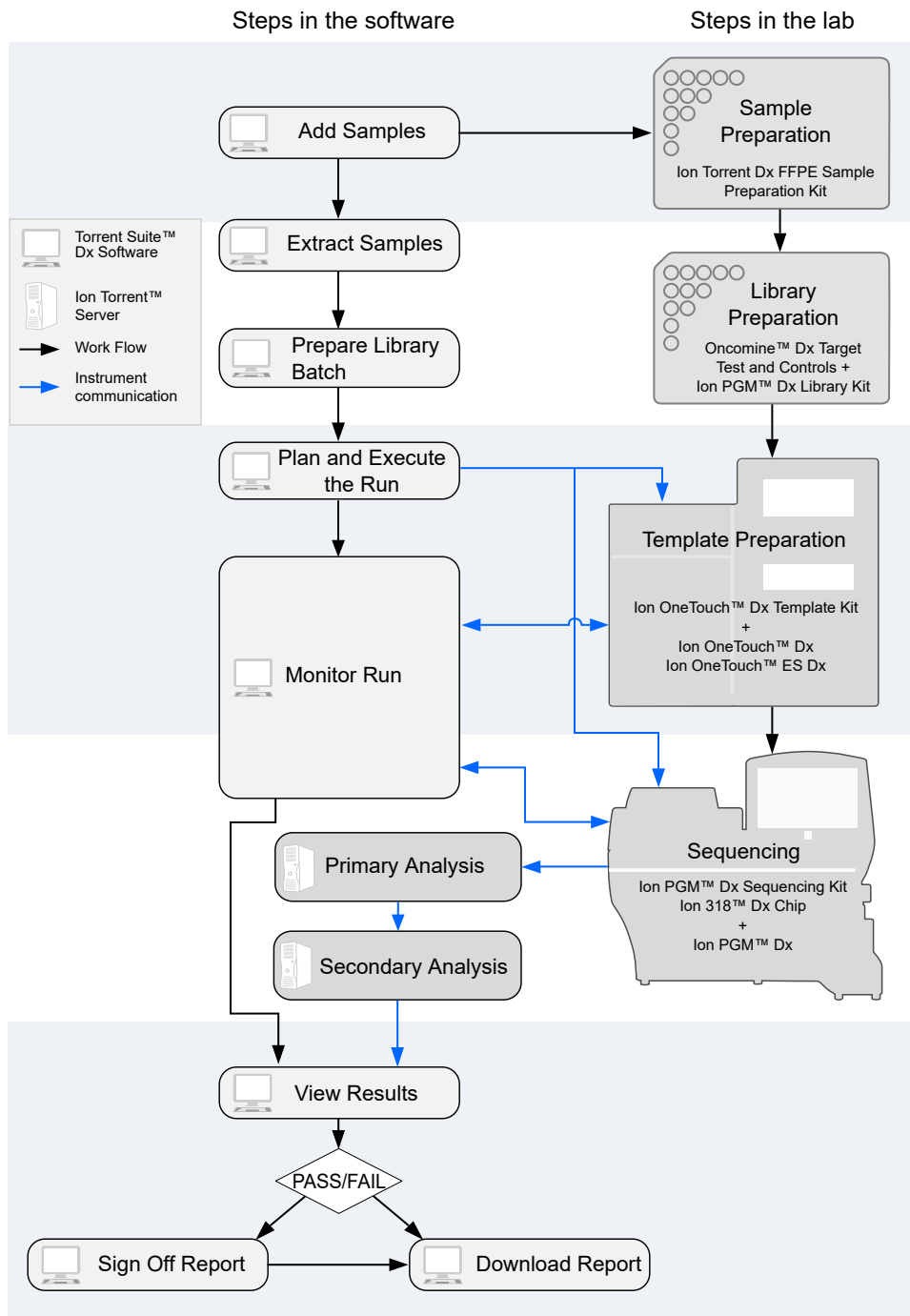
System tracking

The system tracks and checks user, sample, workflow, reagents and QC metrics for auditable records. If the software detects an error at any step—for example, a scanned barcode is inconsistent with the information given for the run—the software alerts the user and does not proceed with the run.

Reagent management

Users should put in place a reagent management system to inventory and track reagent use. Proper use of a reagent management system promotes regulatory compliance and ensures optimal use of kits, chips, and reagents.

Oncomine™ Dx Target Test system diagram



Installation and verification run

Do not attempt to power on any Ion PGM™ Dx System instruments or sign in to Torrent Suite™ Dx Software until a field service engineer (FSE) has installed and performed a performance qualification (PQ) verification run and provided a basic system overview.

The software and all connected instruments are locked until a verification run has been performed and the report is signed by the FSE.

Request and sign in to a new account

Only administrator-level users can create user accounts.

After account creation, the Ion PGM™ Dx System automatically sends an email to the new user with the user name and password information.

- To request a new account, contact your local administrator.
- To sign in to a new account for the first time:
 - a. Go to the Torrent Suite™ Dx Software home page, then enter your user name in the **Username** field.
 - b. Enter the system-generated password in the **Password** field.
 - c. Click **Enter**, or click **Sign In**.
 - d. Click **Accept** to accept the End User Software License Agreement.
 - e. In the **Change Password** screen, enter your temporary password in the **Current Password** field. Type a new password in the **New Password** field, then confirm the password.
 - Passwords must be between 6 and 10 characters.
 - Passwords must contain at least one alphabetic character (aA–zZ).
 - Passwords must contain at least one numeric character (0–9).
 - Passwords must contain only alphanumeric characters (0–9, aA–zZ), no spaces or special characters.
 - Passwords are case-sensitive.
 - f. Click **Change**.

Sign in

To sign in to the software:

1. Open the software home page.
2. Under **Mode Switch** at the bottom of the page, verify that **IVD Mode** is selected.

Note: If you are not in **IVD Mode**, contact a system administrator to switch modes.

3. Select your preferred language from the dropdown list in the upper right corner of the page.
4. Enter your user name and password, then press **Enter** or click **Sign In**.

IMPORTANT! Your user name and password must be unique and not shared with other users.

The software opens to the **Samples** tab and the **Manage Samples** screen.

User interface

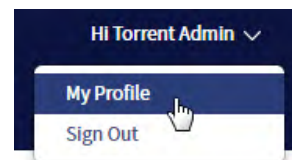
Torrent Suite™ Dx Software has a browser-based interface with five tabs containing the main functions of the software:



- **Samples** tab—Create and manage samples and libraries; plan new runs from libraries.
- **Runs** tab—Create and manage Planned Runs (executable instructions for individual runs). The system uses the Planned Run to verify at each step of the run that the correct kit is used to perform the correct assay on the correct sample.
- **Assay** tab—Manage assays (protocol specifications for templating, sequencing, and data analysis).
- **Monitor** tab—View the status of your system instrument(s) and current runs. Near real-time information is provided on your runs, so that you know early on about any instrument problems.
- **Data** tab—View summaries of completed runs, and detailed run reports; review the run plan settings; download or print output files and run reports.

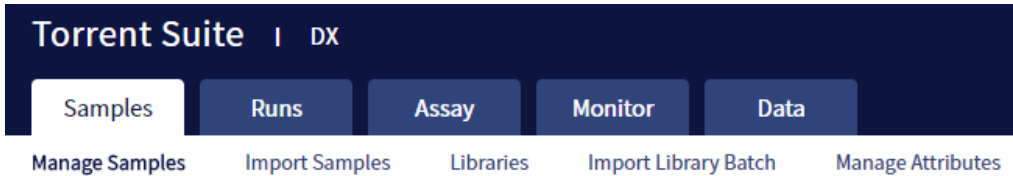
In addition to the main tabs, the **Settings** tab enables access to audit records and log files, server configuration and data management settings, reference file information, and user accounts.

Click **My Profile** from the user name dropdown list in the upper right corner to view your account information and change your password and email address. Click **Sign Out** to sign out of the software.



3

Enter sample and library information



Sample and library information is entered into Torrent Suite™ Dx Software under the **Samples** tab. This information is tracked by the software throughout the entire system workflow.

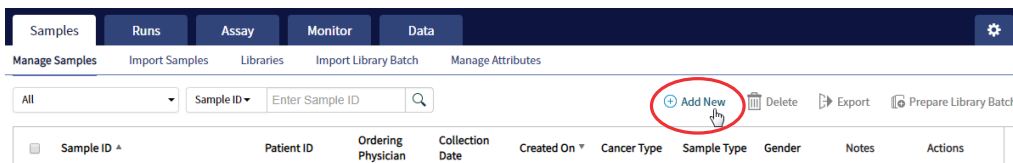
In the **Samples** tab, you can:

- Create, import, and manage samples and their attributes
- Enter the barcode on the kit used to extract each sample
- Prepare libraries and library batches

Add or import samples

Add a new sample



1. Under the **Samples** tab, in the **Manage Samples** screen, click **+ Add New**.



2. Complete the **Add New Sample** dialog box. Fields identified with an asterisk (*) are required. If no information is available, substitute dummy data to complete the required fields.

Field	Description
Sample ID*	<p>A unique identifier representing the sample, containing only alphanumeric characters (0–9 and A to Z), full stops/periods (.), underscores (_), or hyphens (-). The Sample ID cannot contain spaces and is limited to a maximum of 20 characters.</p> <p>After a Sample ID is entered into the system, it cannot be edited. It can be deleted unless it has already been used in a library. The software checks all Sample IDs entered or imported to prevent duplication and returns an error message if a non-unique Sample ID is detected.</p>

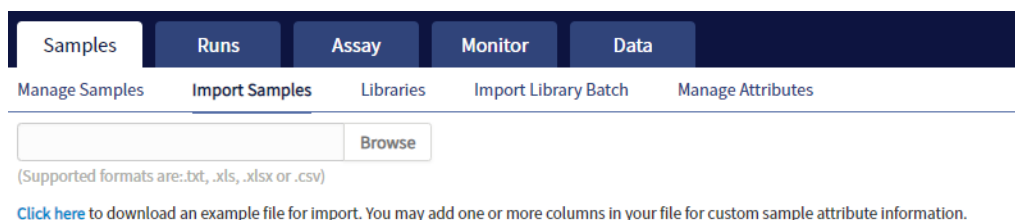
(continued)

Field	Description
Patient ID*	An identifier representing the patient. This field accepts all characters including spaces.
Date of Birth*	The patient's date of birth. Click the  button to select the date in the correct format.
Ordering Physician*	The name of the ordering physician. This field accepts all characters including spaces.
Collection Date*	The date the sample was collected from the patient. Click the  button to select the date in the correct format.
Sample Source	Open-entry field that accepts all characters, including spaces. Example entry: Name of the clinic or hospital ordering the test.
Sample Condition*	Open-entry field that accepts all characters, including spaces. Example entry: FFPE surgical resection.
Sample Type*	Open-entry field that accepts all characters, including spaces. Example entry: Lung tumor, invasive adenocarcinoma.
Gender*	The biological gender of the sample. This must be Male, Female, or Unknown.
Cancer Type*	Select the cancer type of the sample from the dropdown list.
%Cellularity	The percentage of tumor cellularity in the sample.
%Necrosis	The percentage of cellular necrosis in the sample.
Reference Interval	A normal range of measure for the sample.
Notes	An open-entry field.

3. Click **Save**.The sample is listed in the **Manage Samples** screen.

Import samples

Under the **Samples** tab in the **Import Samples** screen, you can import sample data in the form of a TXT, XLS, XLSX, or CSV file. The import file includes the same sample attributes that are listed in the **Add New Sample** dialog box.



The screenshot shows the 'Import Samples' screen. At the top, there is a navigation bar with tabs: 'Samples' (selected), 'Runs', 'Assay', 'Monitor', and 'Data'. Below this, there is a sub-navigation bar with links: 'Manage Samples', 'Import Samples' (active), 'Libraries', 'Import Library Batch', and 'Manage Attributes'. The main content area features a file upload field with a 'Browse' button. Below the field, it states '(Supported formats are: .txt, .xls, .xlsx or .csv)'. At the bottom, there is a link: 'Click here to download an example file for import. You may add one or more columns in your file for custom sample attribute information.'

1. In the **Import Samples** screen, below the **Browse** field, click **Click here** to download a Microsoft™ Excel™ template file.

Note: The template file contains default sample attributes as columns. If additional custom sample attributes have been configured in the software, add these attributes as columns to the template file.

2. In the template file, fill in the information for each sample, one sample per row. See “Predefined sample attributes” on page 20 for more information.
3. Save the file.
4. Click **Browse**, navigate to the saved file, then select it.
5. Click **Import**.
A progress bar followed by an import report displays. If the import process fails, an error message indicates the reason for failure (for example, an invalid character was used). For additional troubleshooting, see “Batch sample import fails” on page 92.
6. Click **Manage Samples** to return to the sample list. Successfully imported samples are listed.

Predefined sample attributes

The software has the following predefined sample attributes, which are listed in the **Add New Sample** dialog box and in the template file for importing samples.

- Sample ID*
- Patient ID*
- Date Of Birth*
- Ordering Physician*
- Collection Date*
- Sample Source
- Sample Condition*
- Sample Type*
- Gender*
- Cancer Type*
- %Cellularity
- %Necrosis
- Reference Interval
- Notes

*Indicates a field required to be filled in during sample creation.

Predefined attributes are locked and cannot be edited. You can create and manage custom sample attributes using the tools in the **Manage Attributes** screen.

Note: LIMS users must create custom attributes before importing sample and Planned Run information from LIMS for the attributes to be propagated through to output files. The software ignores all input file content that is not a recognized attribute.

Create a new sample attribute

1. Under the **Samples** tab in the **Manage Attributes** screen, click **+ Add New Attribute**.
2. Enter an attribute name in the dialog.

Note: Attribute names are limited to ≤20 alphanumeric characters (0–9 and A–Z), full stops/periods (.), underscores (_), or hyphens (-).

3. Specify whether the attribute is a text or number in the **Data Type** dropdown list.
4. Select the **Required** checkbox to designate the new attribute as required in the **Add New Sample** dialog.

Note: Users will not be able to create a new sample if this information is not entered in the **Add New Sample** dialog.

5. Click **Save**.

The new sample attribute is listed in the **Attribute Name** column and is available when you add a new sample. After an attribute has been created, it cannot be deleted. To remove an attribute from use, see “Obsolete an attribute” on page 21.

Note:

- When editing samples created before a new attribute was created, the new attribute appears in the **Edit Sample** dialog. If the new attribute was designated as required, a valid entry must be entered into the new attribute field to save the sample information.
 - Samples can be edited after they have been added to a library batch, but samples cannot be edited after execution of a Planned Run that uses the samples in a library batch.
-

Obsolete an attribute

Under the **Samples** tab in the **Manage Attributes** screen, you can obsolete user-created sample attributes to remove them from use in the software. Obsoleted attributes can be reactivated, and a record of their use is maintained in the audit trail of samples that are created using that attribute.

Note: You can only obsolete user-created attributes, not system-installed attributes.

- To obsolete an attribute, click **Obsolete** in the **Actions** column, then confirm the action. **Reactivate** replaces **Obsolete** in the **Actions** column.
- To reactivate an attribute, click **Reactivate** in the **Actions** column.

Note: All active sample attributes are listed in the **Add New Sample** dialog box.

Enter the Ion Torrent Dx Total Nucleic Acid Isolation Kit barcode

Under the **Samples** tab, in the **Manage Samples** screen, scan the barcode of the Ion Torrent Dx Total Nucleic Acid Isolation Kit used in the extraction process for a particular sample. This barcode is saved with the sample and can be viewed by clicking the Sample ID.

1. Above the samples list, select **To Be Extracted** from the **Filter Samples by...** dropdown list to display only those samples that do not have a kit barcode that is associated with them.
2. Select the checkbox of the sample to be extracted. Select multiple samples if you are using the same kit to process them.

The screenshot shows the 'Manage Samples' interface. The 'Filter Samples by...' dropdown is set to 'To Be Extracted'. The 'Extract' button is circled in red. Below the filter, a table lists one sample with the following details:

Sample ID	Patient ID	Ordering Physician	Collection Date	Created On	Cancer Type	Sample Type	Gender	Notes	Actions
BC1	BC1	Smith	2018-09-05	2018-10-06 02:17	Non-small Cell Lung Cancer	DNA	Male		Edit Audit

3. Click **Extract**. In the dialog box, scan the barcode that is printed on the Ion Torrent Dx Total Nucleic Acid Isolation Kit (box 1 of 2, Part No. A32434).

IMPORTANT! Check the expiration date on the box. If the kit is expired, select another kit.

4. Click **Save**.
The sample is no longer listed in the **To Be Extracted** list.

Manage samples



You can find tools for creating, searching, sorting, editing, deleting, exporting samples, and viewing the sample history under the **Samples** tab in the **Manage Samples** screen.

The screenshot shows the 'Manage Samples' interface with a list of three samples. The 'Filter Samples by...' dropdown is set to 'Sample ID'. The table lists the following samples:

Sample ID	Patient ID	Ordering Physician	Collection Date	Created On	Cancer Type	Sample Type	Gender	Notes	Actions
TestSample_01	Patient_01	Dr. Smith	2017-03-06	2018-11-02 10:07	Non-small Cell Lung Cancer	FFPE	Male		Edit Audit
TestSample_02	Patient_02	Dr. Smith	2017-03-07	2018-11-02 10:07	Non-small Cell Lung Cancer	FFPE	Male		Edit Audit
TestSample_03	Patient_03	Dr. Smith	2017-03-08	2018-11-02 10:07	Non-small Cell Lung Cancer	FFPE	Male		Edit Audit

Search samples

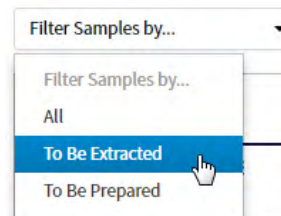
Under the **Samples** tab, in the **Manage Samples** screen:

1. Enter the full or partial sample ID into the **Enter Sample ID** field.
2. Click .
The sample or samples matching the search parameters are listed.
3. Click  to return to the complete list of samples.

Sort samples


Under the **Samples** tab in the **Manage Samples** screen, the list of samples is displayed with the most recently created sample on top by default. To return to the default display, click **Manage Samples**.

- To list only those samples that have not been extracted, click **To Be Extracted** from the **Filter Samples by** dropdown list.
- To list only those samples that have not been prepared as a library, click **To Be Prepared**.
- To list all samples, click **All**.
- To sort the list:
 - a. Click a column header to sort the list by the entries in that column.
 - b. Click the column header again to reverse the order.
 - c. Click **Manage Samples** to return to the default order (most recently created on top).





Export and print samples

The **Export** function generates a XLS file of the sample details.

1. Under the **Samples** tab in the **Manage Samples** screen, in the **Sample ID** column, select the checkbox next to each sample to be exported. Select the checkbox above the column to select all of the samples.
2. Click  **Export**.
An XLS file is generated. Depending on your browser settings, the software automatically downloads the file or prompts you to open or save the file.
3. Open the XLS file in an appropriate viewer to print.

View notes or add a note to a sample

1. Click  in the **Notes** column for a sample to view notes that have been added for the sample.
2. To add a new note, click  **Add Note**. Enter the note in the text field, then click **Save**.
3. Click **Close** to return to the **Manage Samples** screen.

Edit sample

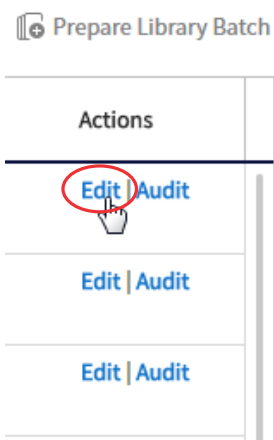
Under the **Samples** tab, in the **Manage Samples** screen, samples that can be edited are identified by the **Edit** link in the **Actions** column.

A sample can be edited at any point before execution of a Planned Run containing that sample, after which the **Edit** link is no longer available. You can edit sample information after a run using the **Edit and Amend Report** command (for more information, see “Edit a sample and amend the report after a run”).

1. Click **Edit** in the row of the sample to be edited.
2. In the **Edit Sample** dialog box, click a field to edit the information.

Note: The Sample ID cannot be edited. For a description of the fields, see “Add a new sample” on page 18.

3. Click **Save**.



Edit a sample and amend the report after a run

After a sequencing run and its analysis have completed, you can edit a sample and amend the Test and Lab Reports for up to 30 days. After 30 days, only the audit link displays. Editing a sample after a run triggers an automatic update of the reports and other files associated with the sample. Under the **Samples** tab, in the **Manage Samples** screen, editable samples are identified by the **Edit and Amend Report** link for the sample in the **Actions** column.

1. To edit a sample after a run, click **Edit and Amend Report** in the row of the sample to be edited.
2. In the **Edit Sample and Amend Report** dialog box, click a field to edit the information.

Note: The Sample ID cannot be edited. For a description of the fields, see “Add a new sample” on page 18.

3. Click **Save**.
4. In the dialog box that appears, click **Yes** to confirm the edit and continue.

Prepare Library Batch

Actions
Edit and Amend Report Audit
Audit
Audit

Edit Sample and Amend Report

Sample edit will trigger an update of Test_Report(s), Lab_Report(s), tab files and Info.csv file. The current version of these files will be replaced and no longer available. Do you want to continue?

The sample information is edited and associated reports and files are updated.

Review sample history


The entire history of a sample is available for review, export, or printing.

1. Under the **Samples** tab in the **Manage Samples** screen, click **Audit** for the sample of interest in the **Actions** column.
The **Audit Trail** dialog box opens, listing each event that modified the selected sample.
2. Click [Details](#) under the **Record** header to view the details of the change made.
3. In the **Audit Record Details** dialog box, click **Export** to export a PDF of the record.

Delete samples

Under the **Samples** tab, in the **Manage Samples** screen, you can delete samples that have not been assigned to a library.

Note: Samples assigned to libraries are locked and cannot be edited or deleted. Locked samples only display **Audit** under the **Sample ID**.

1. Select the sample or samples to be deleted by selecting the checkbox adjacent to the **Sample ID**. Select all the samples on the page by selecting the checkbox above the column.
2. Click  (**Delete**).
The **Delete Sample** dialog box opens with the message "Are you sure you want to delete selected sample(s)?"
3. Click **Yes** to delete the selected samples. Click **No** to return without deleting.

Prepare a library batch in Torrent Suite™ Dx Software

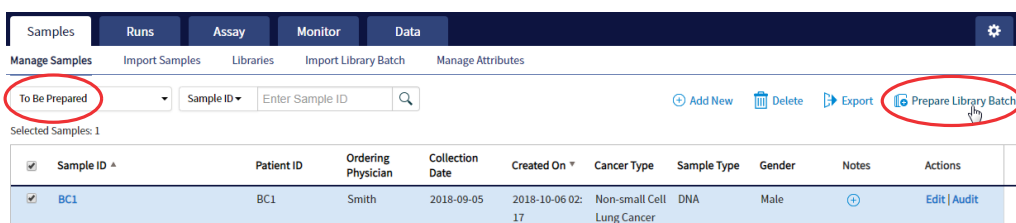
In Torrent Suite™ Dx Software, samples that are entered into the software are placed in library batches for processing and tracking. A library batch consists of a group of libraries that are prepared at the same time.

Note:

- Each library in a library batch must have a unique library name. When combining libraries in the same run, each must also have a unique barcode.
- Control libraries must be included in the same library batch as the sample library they control for.
- Fields identified with a red asterisk (*) are required.

1. Sign in to Torrent Suite™ Dx Software.
2. Under the **Samples** tab, in the **Manage Samples** screen, select **To Be Prepared** from the **Filter Samples by...** dropdown list to display only those samples that have not been placed in a library batch.

IMPORTANT! Samples that have not been queued for extraction in the software also appear on this tab. Ensure that the samples have been queued for extraction before queueing them for library batch preparation.

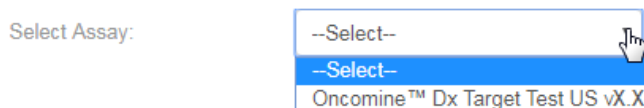


Sample ID *	Patient ID	Ordering Physician	Collection Date	Created On *	Cancer Type	Sample Type	Gender	Notes	Actions
<input checked="" type="checkbox"/> BC1	BC1	Smith	2018-09-05	2018-10-06 02:17	Non-small Cell Lung Cancer	DNA	Male		Edit Audit

3. Select up to 6 samples in the list, then click  **Prepare Library Batch**.
The **Prepare Library Batch** dialog opens. Required fields are indicated with a red asterisk(*).

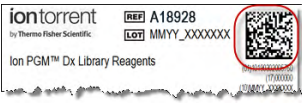

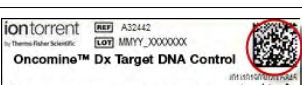
- In the **Select Assay** dropdown list, select **Oncomine™ Dx Target Test**. The assay determines specific parameters of the run, including required controls and post-run data analysis settings.

Prepare Library Batch



- In the following screen, enter a unique identifier for the library batch in the **Library Batch ID** field. Library Batch IDs can only contain alphanumeric characters (0–9 and A to Z), full stop/period (.), underscore (_), and hyphen (-).
- Scan the barcodes from their respective kit boxes into the appropriate fields. Each library batch is associated with a kit lot by scanning the 2D barcode on the appropriate kit box.

IMPORTANT! Check the expiration date on each box. If the kit is expired, select another kit.

Barcode field	Kit	Kit box	Storage	Label scanned
Library Kit Barcode	Ion PGM™ Dx Library Kit	Ion PGM™ Dx Library Reagents	–30°C to –10°C	
Panel Kit Barcode	Oncomine™ Dx Target Test Panel	Oncomine™ Dx Target Test DNA and RNA Panel (box 1 of 3)	–30°C to –10°C	
Control Kit Barcode	Oncomine™ Dx Target Test, Controls, and Diluent Kit	Oncomine™ Dx Target DNA Control v2 (box 2 of 3)	–30°C to –10°C	

- Type a unique library name for each DNA and RNA library in the appropriate field. Library names can only contain alphanumeric characters (0–9 and A to Z), full stop/period (.), underscore (_), and hyphen (-).

Note: The Oncomine™ Dx Target Test Kit requires specific controls, which are automatically listed in the **Prepare Library Batch** dialog.

8. Select the Barcode ID of the adapter used to prepare each library. Swap the default barcodes in the dialog between DNA and RNA using the [DNA ↔ RNA](#) button.

Note: Each library in a library batch must have a different Barcode ID. When preparing the physical libraries, we recommend swapping barcodes between DNA and RNA libraries in consecutive sequencing runs to prevent carryover contamination. See “Alternating barcodes” on page 28.

IMPORTANT! Be careful to ensure that the actual barcodes used to create the libraries match the barcodes that are entered in the **Prepare Library Batch** dialog.

9. In the **Input Quantity** field, enter 10 ng for each library.
10. Click **Save** to save your selections and close the dialog.
The **Libraries** screen opens, listing the libraries that you created. Libraries that are prepared in the same batch have the same **Library Batch ID**.

Alternating barcodes

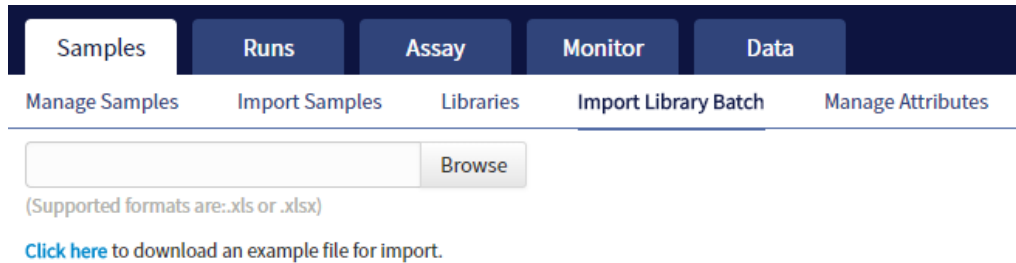
When preparing libraries, we recommend swapping barcodes between DNA and RNA libraries in consecutive sequencing runs to prevent carryover contamination. The following table provides an example of swapping barcodes between runs.

IMPORTANT! Be careful to confirm that the barcodes used to create the libraries match the barcodes entered in the **Prepare Library Batch** dialog box.

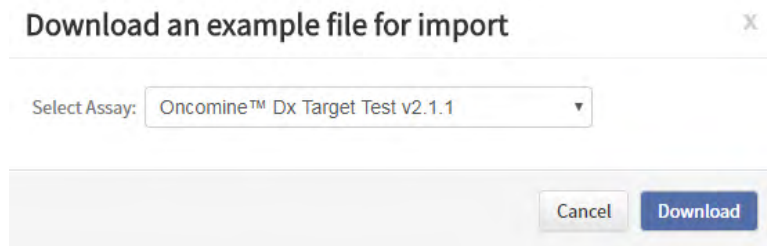
Library type	System Run 1 barcode usage		System Run 2 barcode usage	
	DNA	RNA	DNA	RNA
Positive control	1	9	9	1
Sample	2	10	10	2
Sample	3	11	11	3
Sample	4	12	12	4
Sample	5	13	13	5
Sample	6	14	14	6
Sample	7	15	15	7
No-template control (NTC)	8	16	16	8

Import a library batch

Under the **Samples** tab in the **Import Library Batch** screen, you can import library batch information in the form of an XLS or XLSX file. The import file must include all of the library and kit information that you enter in the **Prepare Library Batch** dialog box.



1. In the **Import Library Batch** screen, below the **Browse** field, click **Click here** to download an example file for import.
2. In the dialog box that opens, select the assay that you are using from the dropdown list, then click **Download**.



The assay name is auto-populated in the Microsoft™ Excel™ template file that downloads to your drive.

3. In the template file, enter or confirm the library batch information.
 - Assay used (auto-populated)
 - Unique Library Batch ID
 - Library kit, control kit, and panel kit barcodes
 - Sample IDs
 - Library names
 - Barcode IDs used for each library and control
 - Nucleic acid type (DNA or RNA)
 - Library input quantity
4. Save the file.
5. Click **Browse**, navigate to the saved file, then select it.




6. Click **Import**.

A progress bar followed by an import report displays. If the import process fails, an error message indicates the reason for failure (for example, an invalid character was used). For additional troubleshooting, see “Library batch import fails” on page 93.

7. Click **Libraries** to return to the library batch screen. Your successfully imported library batch is listed.

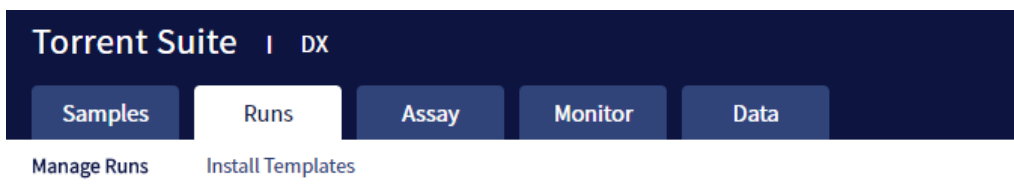
Manage libraries and library batches

Use the following tools in the **Libraries** screen under the **Samples** tab to perform the specified actions.

To...	Do the following...
View more library information	Hover over a name in the library list.
View all libraries	Click All in the Filter Libraries by dropdown list.
View libraries that are not in a Planned Run	Click To Be Planned in the Filter Libraries by dropdown list.
Search the list	Select search by Library Name or Library Batch ID , type a full or partial Library Name or Library Batch ID in the search field above the list, then click Q . Click x to clear the search criteria and display the complete list.
Sort the list	Click a column header to sort the list by the information in that column.
View multiple pages of libraries	Click the scroll buttons below the list.
Delete libraries	Select the checkbox next to the library or related libraries (DNA and RNA), then click  Delete . Note: You can only delete a library if none of the libraries in the same library batch have been assigned to a Planned Run.
Edit a library batch	Click Edit for the library batch in the Actions column, and modify the information in the dialog box.
View or export the audit trail of a library batch	<ol style="list-style-type: none"> 1. Click Audit for the library batch in the Actions column. The Audit Trail dialog box opens, displaying the user, action, and time of each action performed on the batch. 2. To view the action details, click  in the Record column of the Audit Trail dialog box. The Audit Record Details dialog box opens. 3. To export the record as a PDF, click Export in the Audit Record Details dialog box.
Add or modify notes for a library	Click  in the Notes column.

4

Create and manage Planned Runs



Planned Runs contain all the sample, library, and assay information required to perform a run on template preparation and sequencing instruments. The Planned Run information is tracked by the system from template preparation through sequencing, data analysis, and final results.

Prerequisites to create a Planned Run

Before creating a Planned Run, check that:

- Sample information is correctly entered
- Library batches have been prepared
- Each library has been assigned a unique Barcode ID

The software returns an error message when any of these conditions are not met when creating a Planned Run.

The system identifies and tracks kit components and chips by barcodes. The system includes a cordless barcode reader and a barcode reader that is attached to the Ion PGM™ Dx Sequencer.

Note: All kits and chips that are used in a diagnostic assay must be uniquely identified, and the identification must be stored so that the record can be audited.

Create a Planned Run

Libraries that are ready to be entered into a Planned Run are listed under the **Samples** tab in the **Libraries** screen.

Note: You can also plan a run from the **Runs** tab in the **Planned Runs** screen (click **+ Add New**).

1. Sign in to Torrent Suite™ Dx Software.
2. In the **Libraries** screen, select the library or libraries to be run by selecting checkboxes in the list. To view only those libraries that have not yet been added to a Planned Run, select **To Be Planned** in the **Filter Libraries by** dropdown list.

Note:

- Libraries prepared with the same assay in the same library batch can be combined and run together, as long as they have unique library names and Barcode IDs.
- To plan a run with the OncoPrint™ Dx Target Test assay, a report template must be created and associated with the assay by an Administrator or Manager. Under the **Assay** tab, go to **Presets ▶ Report Templates** to add a new report template.
- Up to 16 libraries (including controls) can be sequenced in a single run.
- If libraries need to be rerun due to a run failure, they can be added to a new Planned Run.

3. Click **+ Plan a Run**.

The screenshot shows the 'Libraries' screen in the software. The 'Libraries' tab is active. Below the navigation bar, there are options for 'To Be Planned' (selected), 'Library Name', and a search icon. A table of libraries is displayed with the following data:

Library Batch ID	Assay Name	Sample ID	Library Name	Library Type	Barcode ID	Notes	Actions
21081007_BC1	OncoPrint™ Dx Target Test v2.1.1	BC1	BCL_DNA BCL_RNA	DNA RNA	IonDx-2 IonDx-10		Edit Audit

The 'Plan a Run' button in the Actions column of the first row is circled in red. The page also shows 'Selected Libraries: 1' and a pagination control for '20 Items per page'.

4. In the **Add New Plan** dialog box, enter a name for the run, then select the appropriate report template.

The selected library or libraries are listed in the dialog box, and the control libraries are automatically listed.

Add New Plan X

Name: *

Assay Name: Oncomine™ Dx Target Test v2.1.1

Select Report Template:

Notes:

Number of Sample Libraries: 2

<input type="checkbox"/>	Library Batch ID	Sample ID	Library Name	Barcode ID	Library Type
<input type="checkbox"/>	21081007_ BC1	BC1	BC1_DNA	IonDx-2	DNA
			BC1_RNA	IonDx-10	RNA
<input type="checkbox"/>		NA	internalCo ntrol_14	IonDx-9	RNA Contr ol
<input type="checkbox"/>		NA	internalCo ntrol_14	IonDx-10	DNA Me T

Remove

- To remove libraries from the run, select the appropriate checkboxes, then click **Remove**. To add libraries, click **+ Add more Libraries**, then select them from the **Add Libraries** dialog box. Any added libraries must be from the same library batch and have unique library names and Barcode IDs.
- Click **Save**.
The new Planned Run is automatically assigned a Run Short Code and is displayed at the top of the list under the **Runs** tab in the **Manage Runs** screen.

Execute a Planned Run

Planned Runs are listed under the **Runs** tab in the **Manage Runs** screen. Runs that are ready to be performed have the **Execute** command available in the **Actions** column.

Executing a Planned Run in the software queues the run for initiation on the Ion OneTouch™ Dx Instrument. After a Planned Run has been queued for execution, the operator should immediately start template preparation.

In the **Manage Runs** screen:

1. Click **To Be Started** in the **Filter Run by...** dropdown list to limit the list of Planned Runs to only those runs yet to be started.
2. Find the Planned Run in the list, then under the **Actions** header, click **Execute**.



The **Execute Planned Run** dialog box opens.

3. In the **Tube Label** field, enter the text that is used to label the tubes that contain the final combined libraries.

The tube label text can be any combination of letters and numbers. The system uses this text to track the sample throughout the run, so be careful to label each tube legibly at the points noted in the procedure. The software does not allow use of the same Tube Label text within 7 days.

4. Click the **Template Prep Kit barcode** field, then scan the barcode from the Ion OneTouch™ Dx Template Reagents box.

IMPORTANT! Ensure that you scan the barcode from the actual reagents box that is used in the run.



5. Click **Save** to save your changes.
The **Review Planned Run** dialog box opens.
6. Write down the **Run Short Code** and/or click **Print** to print the scannable barcode. The code must be entered into the Ion OneTouch™ Dx Instrument and Ion PGM™ Dx Sequencer for tracking and verification before the start of the instrument run.
7. Click **Close** to close the dialog box and send the run to the instrument.

Note: The last 5 executed Planned Runs are listed under the **Monitor** tab in the software.

Manage Planned Runs

Under the **Runs** tab, in the **Planned Runs** screen, a list of all Planned Runs is displayed, with the most recently created run on top. In this screen, you can:

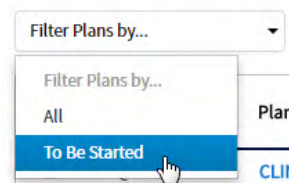
- Sort and search the list of Planned Runs
- Review and print a Planned Run
- View the audit trail of a Planned Run
- Edit a Planned Run
- Delete a Planned Run (Manager/Administrator level users only)

Search and sort Planned Runs

Under the **Run** tab, in the **Manage Runs** screen, the list of Planned Runs is displayed, with the most recently created on top by default.

- To search the Planned Run list:
 - a. Enter the full or partial Planned Run name into the **Planned Run** field.

- b. Click **Q** .
The Planned Run(s) matching the search parameters are listed.
- c. Click **x** to return to the complete list of Planned Runs.
- To sort the Planned Runs list:
 - a. Click the header name of interest.
The list of Assays reorders alphabetically or numerically based on the header name selected: Run Short Code, Planned Run Name, Assay, and Tube Label.
 - b. Click the header name a second time to reverse the order of samples displayed.
- Click **To Be Started** in the **Filter Plans by...** dropdown list to limit the list of Planned Runs to only those runs that are ready to be executed.




Review/Print a Planned Run

You can review and print the information in a **Planned Run** when you execute it. Alternatively:

1. Under the **Runs** tab, in the **Manage Runs** screen, click the Planned Run name in the list.
2. In the **View Planned Run** dialog box, review the information that was entered or generated when you created the run, then click **Print** to print it.

Review a Planned Run audit trail

The entire history of a Planned Run is available for review, export, or printing.


1. Under the **Runs** tab, in the **Manage Runs** screen, click **Audit** in the **Actions** column for the desired run.
The **Audit Trail** dialog box opens. Each modifying event for the selected Planned Run is listed.
2. Click the details  icon under the **Record** header to view the details of the change made.
The **Audit Record Details** dialog box opens detailing the edits that were made.
3. In the **Audit Record Details** dialog box, click
 - a. **Export** to export a Print-Ready PDF of the record.
 - b. **Cancel** or the **X** icon to return to the **Audit Trail** dialog box.
4. Open the downloaded PDF file in an appropriate viewer, then print the record from within the open document.
5. Click **Cancel**, or the **X** icon, to return to the **Manage Runs** screen.

Edit a Planned Run

You can edit Planned Runs that have not been executed yet.

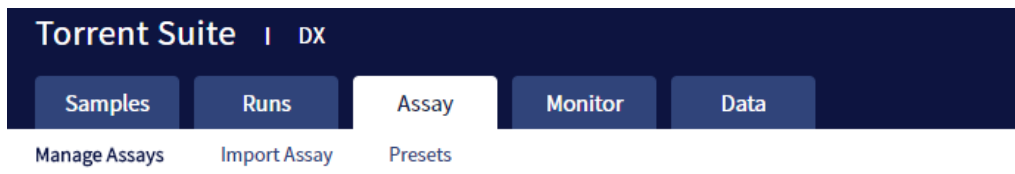
1. Under the **Runs** tab, in the **Manage Runs** screen, click **Edit** for the Planned Run in the **Actions** column.
2. In the **Edit Planned Run** dialog box, change the settings, then click **Save**.

Delete a Planned Run (Manager/Administrator only)

1. Under the **Runs** tab, in the **Manage Runs** screen, select the Planned Run to be deleted by selecting the checkbox next to the **Run Short Code**. Select all of the Planned Runs on the screen by selecting the checkbox above the column.
2. Click  **Delete**.
The **Delete Plan** dialog box opens with the message "Are you sure you want to delete the selected planned run(s)?"
3. Click **Yes** to delete the Planned Run, or click **No** to return to the **Manage Runs** screen without deleting.

System installation templates (service engineers only)

IMPORTANT! The templates used by service engineers during Ion PGM™ Dx System installation are listed under the **Runs** tab in the **Install Templates** screen. These should not be used or modified in any way.



The **Assay** tab has tools for viewing the Oncomine™ Dx Target Test assay and managing and creating report templates.

Oncomine™ Dx Target Test assay

The Oncomine™ Dx Target Test assay is a locked assay that contains the settings and parameters for analyzing sequencing results and determining valid variant calls for the Oncomine™ Dx Target Test. It also defines the kits and chips that can be used with the test, and specifies the threshold values for quality control and variant detection.

IMPORTANT! The Oncomine™ Dx Target Test assay cannot be edited or used as the basis for a Reanalysis assay.

Assay	Application	DNA Panel	RNA Panel	Status	Created By	Created On *	Actions
★ Oncomine™ Dx Target Test vX.X	DNA+RNA	Oncomine™ Dx Target Panel vX.X Regions	Oncomine™ Dx Target Panel vX.X Fusions	Locked	System Installed	2018-11-01 22:53	Audit

Search and sort assays

Under the **Assay** tab in the **Manage Assays** screen, the most recently created assay is listed first by default. To return to the default display, click **Manage Assays** to refresh the screen.

- To search the assay list:
 - a. Enter the full, or partial, assay name into the **Assay Name** field.
 - b. Click .
The assay or assays matching the search parameters are listed.
 - c. Click to return to the complete list of assays.

- To sort the assay list:
 - a. Click the desired column header.
The list of assays reorders based on the header name selected: Assay, Application, Status, Created By, Created On.
 - b. Click the header name a second time to reverse the order.


View assay details

The **View Assay Details** dialog displays all the settings for an assay. To open this dialog:

1. In the **Manage Assays** screen, click an assay name in the list.
2. Scroll down to view the all the settings.

Review the assay audit trail

Managers and Administrators can review and export the details of all the actions that are performed on an assay.

1. In the **Manage Assays** screen, click **Audit** for the assay in the **Actions** column.
2. Each action that is performed on the assay is listed in the **Audit Trail** dialog. Click  in the **Record** column to view the details of each action.
3. In the **Audit Record Details** dialog, click **Export** to export a PDF of the selected record.

Assay presets



Report templates

You can create and manage custom report templates used to generate PDF reports after a run. You select a report template when you set up a Planned Run.

The tools for setting up report templates are located in the **Report Templates** subtab under the **Assay** tab in the **Presets** screen.

Manage report templates

The tools for managing report templates are found under the **Assay** tab in the **Presets** screen under the **Report Templates** subtab.

- To view the details of a report template, click the template name.
- To preview the template layout, click **Preview** in the **Actions** column.
- To make a template unavailable in the software, click **Obsolete** in the **Actions** column.
- To add notes to a template, click  in the **Notes** column.
- To view notes for a template, click  in the **Notes** column.

Create a report template

To create a report template:

1. Under the **Assay** tab in the **Presets** screen, click the **Report Templates** subtab.
Existing report templates are listed on this page, including any system-installed templates.
2. Click **+ Add New**.

Note: To plan a run with the OncoPrint™ Dx Target Test assay, a report template must be created and associated with the assay by an Administrator or Manager.

3. Enter the **Report Name**, then select the **Assay** from the dropdown list.
4. In the **Select Language** dropdown list, select the language for generating the report.
5. Fill out the information in the remaining sections. Fields identified with a red asterisk (*) are required.

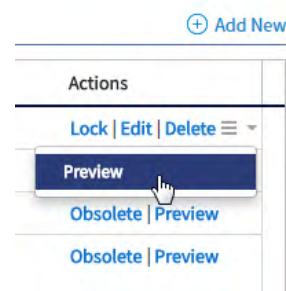
Note: For locked assays, some fields and attributes cannot be edited and are grayed out.

6. Under **Sample Details**, drag and drop the attributes to display in the report into one of the three column placeholders. You can also drag and drop the additional attributes below the columns into the columns. Select the checkbox next to an attribute to include the attribute in each page header.
7. When you are done, click **Save**.
The new report template appears in **Report Templates** list.

8. To preview the report layout, click **☰ ▼**, then click **Preview** in the **Actions** column in the row of the new template. Click **Edit** to make changes.

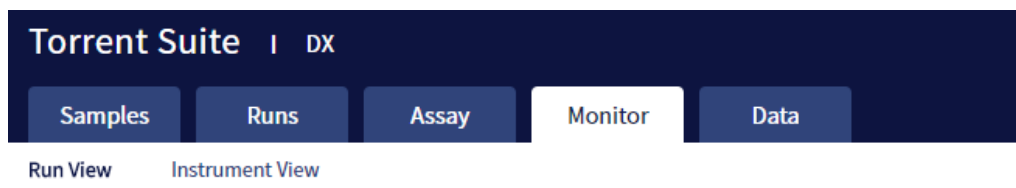
9. When you are finished, click **Lock**.

Note: The template must have a locked status to be used in a Planned Run.





Monitor runs



In the **Monitor** tab, you can view the status of any jobs running on instruments that are connected to the Ion Torrent™ Server. To monitor the status of runs versus instruments, click **Run View** or **Instrument View** under the tab.



Run information is displayed for the last 5 runs that are in progress, have failed, or have completed data analysis.

Information displayed

The **Run View** and **Instrument View** screens under the **Monitor** tab display the following information:

View	Information displayed
Run View	Ion OneTouch™ Dx Instrument and Ion PGM™ Dx Sequencer <ul style="list-style-type: none"> • Instrument name • Operator • Start and Completion times • Time remaining • Run Status
	Ion PGM™ Dx Sequencer only <ul style="list-style-type: none"> • Flow Transfer number • Analysis Status • QC Details (expand to view after analysis is complete) <ul style="list-style-type: none"> – Run QC – Sample QC – Internal Control QC
Instrument View	Ion OneTouch™ Dx Instrument and Ion PGM™ Dx Sequencer <ul style="list-style-type: none"> • Current state • Instrument status • Last cleaning • PQ status • PQ due date • Instrument serial number • Any instrument alarms

View a run

Under the **Monitor** tab, in the **Run View** screen, the last 5 sequencing runs (active, completed, or failed) are listed. The instrument type is indicated by either an Ion OneTouch™ Dx Instrument icon , or an Ion PGM™ Dx Sequencer icon  adjacent to the instrument serial number in the respective panes. To view a run:

1. Click **Refresh** to update the displayed information.
2. In the **Select Run** dropdown list, select the Planned Run to be viewed.
The instruments used for template preparation and sequencing will be displayed.
3. In the **Select Library** dropdown list, select the library in the run that you want to view.

To view details of the assay used in the designated run, click on the assay name adjacent to the dropdown lists.

View the status and quality control metrics of a run

Under the **Monitor** tab, in the **Run View** screen, you can review the status and quality control metrics of the last 5 template preparation and sequencing runs (active, completed, or failed).

1. In the **Select Run** dropdown list, select the Planned Run.
The status of the instruments used in the run are displayed. The **Analysis Status** field indicates whether post-sequencing analysis ran to completion or failed.
2. To update the status of the run, click **Refresh**.
3. To display additional information about the run, select from the following options:
 - To view details of the assay used in the run, click the assay name next to the dropdown lists.
 - To view the quality control information for a library in the run, select the library from the **Select Library** dropdown list, then click **+View QC Details**.

Note: The QC metrics are described in “Quality control metrics” on page 44.
See the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide* for detailed pass/fail criteria and a repeat testing strategy.

Restart analysis (Administrator)

Under the **Monitor** tab, in the **Run View** screen, the **Analysis Status** field indicates whether data analysis has started, is running, or has completed. An analysis listed as "RUNNING" for more than 12 hours may be stuck in the pipeline. After 12 hours, a **Restart Analysis** button appears under the tab for Administrator-level users.

Click the button to restart the analysis from the beginning.

The screenshot shows the 'Run View' interface with the 'Monitor' tab selected. At the top, there are navigation tabs: Samples, Runs, Assay, Monitor, and Data. Below these, there are dropdown menus for 'Select Run' (OCP_AMR_SGE_S) and 'Select Library' (OCP_AMR_IonDx-7). A red circle highlights the 'Restart Analysis' button. To the right is a 'Refresh' button. The main content area is divided into two sections: 'Templating' and 'Sequencing'. The 'Templating' section shows instrument name 'GoldenData', operator 'Vidya', and status 'COMPLETED'. The 'Sequencing' section shows instrument name 'GoldenData', operator 'ion.reporter@lifetech.com', and status 'SigProcActor : RUNNING', which is also circled in red. Below the sequencing section, it says 'Flow Transfer : 2 of 500 transferred'.

Repopulate QC (Administrator)

Under the **Monitor** tab, in the **Run View** screen, the **Repopulate QC** button appears for Administrator-level users after a run is complete. If the **QC Status** of a run is **Failed** and one or more QC metrics are listed as **N/A**, the QC metrics may not have been updated in the database correctly after the run.

The screenshot shows the 'Monitor' tab in the 'Run View' screen. The 'Repopulate QC' button is circled in red. Below the button, the 'View QC Details' section is expanded, showing a table of QC metrics:

	Metric Name	Value	Reference Range	QC Status
Run QC	Library RNA	Total Mappable Reads	N/A	Failed
	Library DNA	Percent Reads	N/A	Failed
Library QC	Library RNA	Mean AQ20 Read Length (bp)	N/A	Failed
	Library DNA	Percent Reads	N/A	Failed

Administrator-level users can click **Repopulate QC** to repopulate the QC metrics in the database.

Note: This function does not recalculate or change the QC data from the run. It repopulates the database with the previously calculated metrics and regenerates the Lab and Test Reports. The report status (draft or signed) is retained.

1. Under the **Monitor** tab in the **Run View** screen, under **View QC Details**, check the **QC Status** of the run. If the status is **Failed** and any QC metric is listed as **N/A**, proceed to the next step.
2. Click **Repopulate QC** to repopulate all QC metrics in the database.

Note: Repopulation takes 15–30 minutes.

3. If any metric is still listed as **N/A** and the **QC Status** is still **Failed**, repeat the run starting at template preparation from the DNA/RNA combined library, and ensure that you add all controls. If the problem continues, contact Technical Support.

Quality control metrics

In the **Run View** screen, once analysis of a sequencing run is complete, the following quality control details can be viewed.

QC detail	QC metric and description
Library QC: DNA	<ul style="list-style-type: none"> • Mean AQ20 Read Length (bp): The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of a library. • Percent Reads: The number of library reads normalized by the total addressable wells in a run.


(continued)

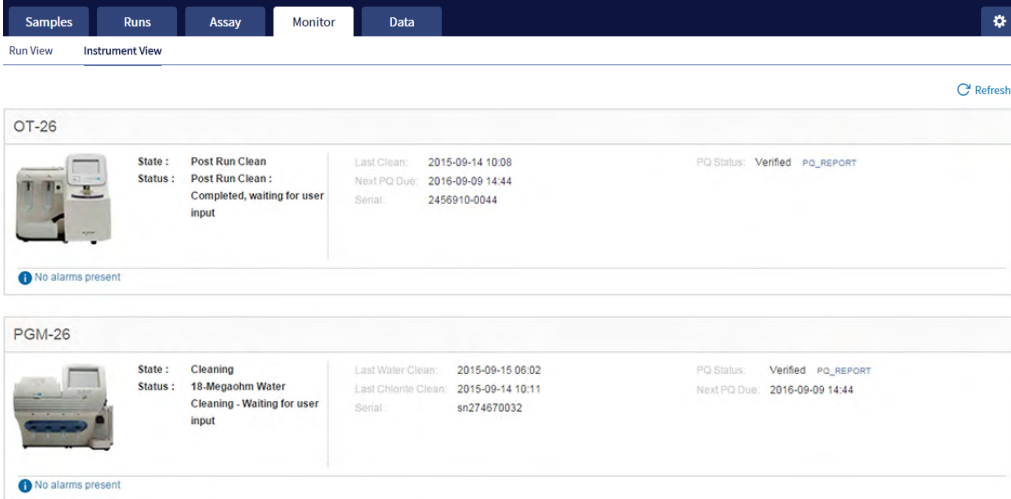
QC detail	QC metric and description
Library QC: RNA	<ul style="list-style-type: none"> • Mappable Fusion Reads: The number of reads that are mapped to the fusion reference file.
Control QC: CF-1	<ul style="list-style-type: none"> • Mean AQ20 Read Length (bp): The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of CF-1. • Percent Reads: The number of all usable library reads that aligned with the CF-1 sequence divided by the total number of addressable wells.
Control QC: DNA	<ul style="list-style-type: none"> • Individual COSMIC variant calls and allelic frequencies: Individual variant positions are assessed in the DNA control for presence or absence of the variant. • Mean AQ20 Read Length (bp): The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of a control fragment. • Percent Reads: The number of all usable library reads that aligned with the control fragment sequence divided by the total number of addressable wells.
Control QC: RNA	<ul style="list-style-type: none"> • Mappable Reads: The number of reads that are mapped to the fusion reference file. • ROS1 Fusion: Detection of ROS1 fusion gene. • ROS1 Fusion Reads: The number of target reads mapping to the ROS1 gene fusion.
Control QC: DNA NTC	<ul style="list-style-type: none"> • Hotspot Calls: The total number of hotspots where a call was made.
Control QC: RNA NTC	<ul style="list-style-type: none"> • Mappable Reads: The number of reads that are mapped to the fusion reference file. • Total Fusion Calls: The total number of fusion calls made.

Quality control pass/fail criteria

Metric	Criteria
Run QC	
CF-1 Mean AQ20 Read Length (bp)	≥131
CF-1 Percent Reads (%)	≥0.03
DNA NTC	Hotspot calls = 0
RNA NTC	Total fusion calls = 0
RNA NTC Mappable Reads	≤4999
DNA Library	
Mean AQ20 Read Length (bp)	≥90
Percent Reads (%)	≥0.7
RNA Library	
Mappable Fusion Reads	≥5000
DNA Control	
COSM476_AF	Variant called and AF ≥0.05
COSM521_AF	Variant called and AF ≥0.05
COSM6223_AF	Variant called and AF ≥0.05
COSM6224_AF	Variant called and AF ≥0.05
COSM760_AF	Variant called and AF ≥0.05
COSM28747_AF	Variant called and AF ≥0.05
Mean AQ20 Read Length (bp)	≥98
Percent Reads (%)	≥0.7
RNA Control	
Mappable Reads	≥18164
ROS1 Fusion Reads	Variant called and fusion reads ≥349

Instrument View screen

The instruments connected to the Ion Torrent™ Server are listed under the **Monitor** tab in the **Instrument View** screen. To update the information in this screen during a run or following completion of a run, click  **Refresh**.



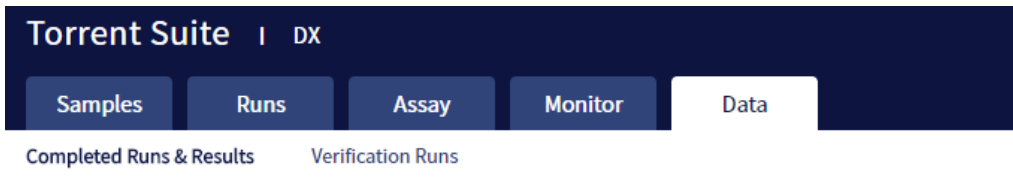
The screenshot shows the 'Monitor' tab selected in the top navigation bar. Below the navigation, there are two instrument cards. The first card is for 'OT-26' and shows a state of 'Post Run Clean' with a status of 'Completed, waiting for user input'. The second card is for 'PGM-26' and shows a state of 'Cleaning' with a status of '18-Megohm Water Cleaning - Waiting for user input'. Both cards include details for 'Last Clean', 'Next PQ Due', 'Serial', and 'PQ Status' (Verified) with a link to 'PQ_REPORT'. A 'Refresh' button is located in the top right corner of the main content area.

The following information is displayed in this screen:

Label	Description
State	The function that the instrument is currently performing (for example, Clean , Initialize , or Run).
Status	The current status of the active function, including data analysis.
Last Clean	The date and time that the instrument was last cleaned. For the Ion OneTouch™ Dx Instrument, this refers to the pre-run cleaning and is not updated following the post-run cleaning.
Serial	The serial number of the instrument.
PQ Status	The performance qualification (PQ) status of the instrument (Verified , Not Verified , or Expired). Click the link next to the status to download and view the PQ report. Note: If the PQ status is Expired , users cannot perform a non-PQ run on the instrument until a PQ run is performed and verified.
Next PQ Due	The expiration date of the current PQ run.

7

Review data and results



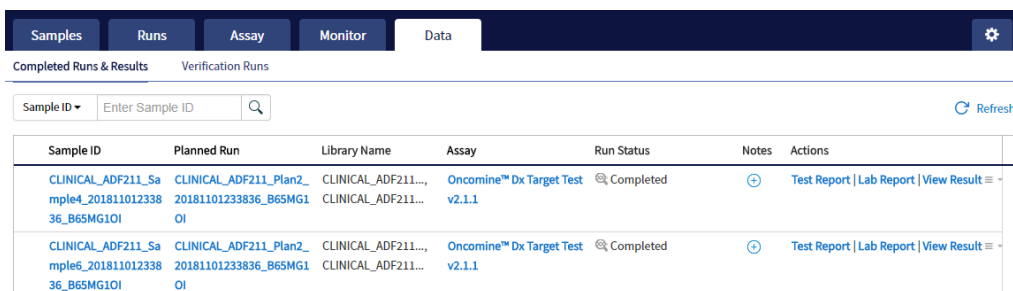
In the **Data** tab, you can review run results and perform data analysis and data management tasks:

- Select **Completed Runs & Results** (the default window) to review completed sample run results and reports. Run results are listed by **Sample ID**.
- Select **Verification Runs** to review data from completed verification runs performed during installation or PQ validation.

Completed Runs & Results screen

Under the **Data** tab, in the **Completed Runs & Results** screen, samples that have been sequenced are listed by Sample ID.

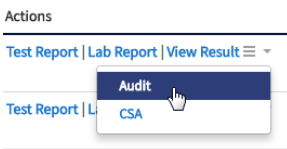
You can search the list of results by Sample ID or Planned Run name. You can also filter and sort the list.



The following information is displayed in the screen:

Column	Description
Sample ID	The unique identifier created when the sample was entered into the software. Click on the link to display the details of the sample.
Planned Run	The name of the Planned Run given when it was created in the software, after preparation of the library batch. Click on the name to display details of the Planned Run.

(continued)

Column	Description
Library Name	The names of the DNA and RNA sample libraries given during preparation of the library batch.
Assay	The assay selected when the sample was placed in a library batch prior to creating the Planned Run. Click on the assay name to display the details of the assay.
Run Status	The status of the run (for example: completed, failed, terminated).
Notes	User-entered notes about the sample. To view previously written notes, click . To add new notes, click .
Actions	<p>Click the appropriate link to:</p> <ul style="list-style-type: none"> Download the Test Report. Download the Lab Report. Open the View Result screen. View the audit trail for the Planned Run. Click , then click Audit.  <ul style="list-style-type: none"> Download Customer Support Archive (CSA) log files for the run to aid in troubleshooting. Click , then click CSA.

Download the Lab Report or Test Report

1. To download the Lab Report or Test Report for a sample of interest, in the **Completed Runs & Results** screen, click the appropriate link in the **Actions** column for the sample. A ZIP file containing all languages of the report downloads automatically.

Completed Runs & Results Verification Runs

Sample ID ▾ Refresh

Sample ID	Planned Run	Library Name	Assay	Run Status	Notes	Actions
CLINICAL_ADF211_Sample 4_20181101233836_B65MG 10I	CLINICAL_ADF211_Plan2_201811 01233836_B65MG10I	CLINICAL_ADF211..., CLINIC AL_ADF211...	OncoPrint™ Dx Target Test v2.1.1	Completed		Test Report Lab Report View Result

2. Extract the downloaded files, then open the PDF file of the desired language in an appropriate viewer.

Test Report

The Test Report (available in the **Completed Runs & Results** screen) is a clinical report generated by the software that can be downloaded in PDF format. It is identical to the Lab Report, except that it does not include the Sequencing Run Details and Control and Sample QC Evaluation Metrics sections. The Test Report contains the following sections and information.

Note: The display of variants in the Test Report depends on cancer type.

Section	Description
Sample Details	The sample and patient information entered into the software.
Results for Sequence Variations for Therapeutic Use	Displays the results for the clinical variants and gene fusions in the sample, and any recommended therapies that are clinically indicated for the selected cancer type.
Results for Analytical Sequence Variations Detected	Displays the results for analytical variants and gene fusions detected in the sample, depending on the variants associated with the cancer type.
Test Description	A description of the test and the gene variants in associated tissue types that it screens for.
Results for Analytical Sequence Variations Not Detected	Displays the results for analytical variants and gene fusions not detected in the sample, depending on the variants associated with the cancer type.

Lab Report

The Lab Report (available in the **Completed Runs & Results** screen) is a report generated by the software that can be downloaded in PDF format. The Lab Report contains the following sections and information.

Note: The display of variants in the Lab Report depends on cancer type.


Section	Description
Sample Details	The sample information entered into the software.
Results for Sequence Variations for Therapeutic Use	The results for the clinical variants and gene fusions in the sample, and any recommended therapies that are clinically indicated for the selected cancer type. Allele frequencies are also reported.
Results for Analytical Sequence Variations Detected	A list of the analytical variants and gene fusions detected by the assay, and associated information for each, depending on the variants associated with the cancer type.
Test Description	A description of the assay.
Analytical Sequence Variations Not Detected	A list of all the analytical variants and gene fusions not detected by the test, and associated information for each, depending on the variants associated with the cancer type.

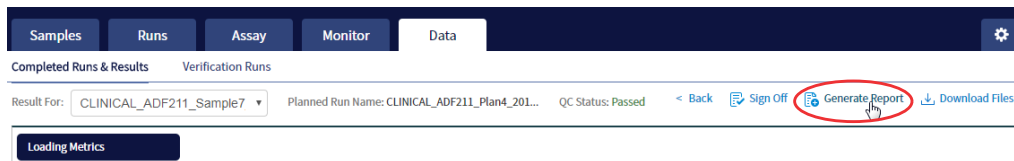
(continued)

Section	Description
Sequencing Run Details	A list of all the kits and instruments used to perform the test.
Control and Sample QC Evaluation Metrics	A summary of the quality control metrics.

Generate customized reports

By default, the Test Report and Lab Report are generated in the language that is selected in the report template. To generate these reports in another language, or change the types of variant calls that are reported, do the following:

1. In the **Completed Runs and Results** screen, click **View Result**.
2. In the **View Result** screen, click  **Generate Report**.

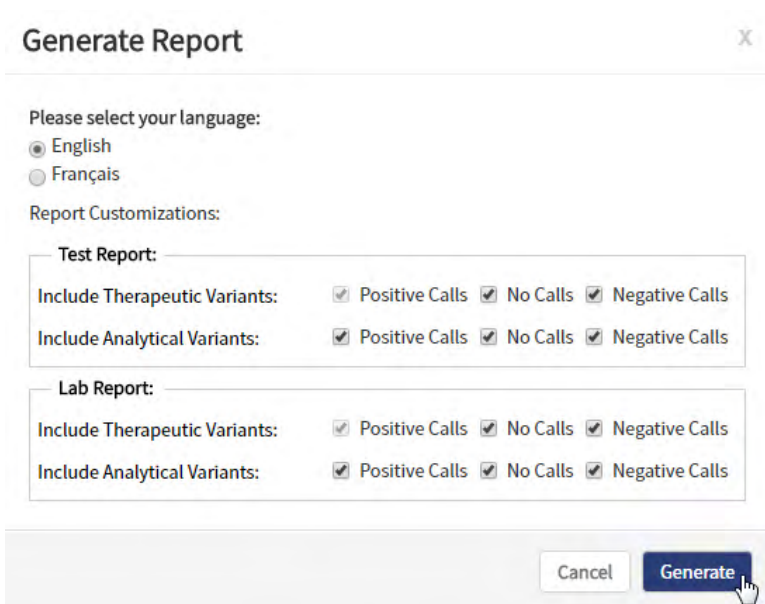


The **Generate Report** dialog box opens.

3. Do one or both of the following:

- Select the desired language.
- Select the checkboxes for the positive calls, no calls, or negative calls that you want to report in the Test Report and/or Lab Report.

Note: These checkboxes are only available in IVD mode if they are enabled for the selected assay.



Generate Report X

Please select your language:

English
 Français

Report Customizations:

Test Report:

Include Therapeutic Variants: Positive Calls No Calls Negative Calls

Include Analytical Variants: Positive Calls No Calls Negative Calls

Lab Report:

Include Therapeutic Variants: Positive Calls No Calls Negative Calls

Include Analytical Variants: Positive Calls No Calls Negative Calls

Cancel Generate

Note: Positive calls for therapeutic variants cannot be deselected.

4. Click **Generate**.

A message displays when the report has been generated. The new report overwrites the existing report.

- 5.** Click [↓](#) **Download Files** to open a dialog box to select the new report and other run-related files for download.

Restore archived results

Torrent Suite™ Dx Software can be configured to automatically transfer older run data, results files, and signed reports from an Ion Torrent™ Server to an external server, based on when the results were generated (see “Disk space usage and data archiving” on page 86). Archived results and reports can then be restored to the Ion Torrent™ Server and downloaded from the **Completed Runs & Results** screen.

IMPORTANT!

- If results are restored from runs that were archived in an older version of Torrent Suite™ Dx Software (version 5.6.4 and earlier), the software may not display all the variants from the restored results in the user interface, and reports generated from those restored results may not contain all variants. Carefully review the restored data in the user interface to determine whether all variants are present. Do not generate new reports from results restored from these older archives. Note that all variants are preserved in the restored source files, and can be downloaded using the **Download Files** command. This issue was corrected in version 5.8 of the software.
- In older versions of Torrent Suite™ Dx Software (version 5.6.4 and earlier), results reports should be generated and signed before results are archived. See “Sign the run results” on page 67.

Archived results are listed in the **Completed Runs & Results** screen with the **Restore** link active in the **Actions** column. To restore and download archived results:

1. In the **Completed Runs & Results** screen, locate the archived result, then click **Restore** in the **Actions** column.

Sample ID	Planned Run	Library Name	Assay	Run Status	Notes	Actions
CLINICAL_ADF211_Sample 2_20181101233836_B65MG 1OI	CLINICAL_ADF211_Plan2_201811 01233836_B65MG1OI	CLINICAL_ADF211..., CLINIC AL_ADF211...	Oncomine™ Dx Target Test v2.1.1	Completed		Restore
CLINICAL_ADF211_Sample 3_20181101233836_B65MG 1OI	CLINICAL_ADF211_Plan2_201811 01233836_B65MG1OI	CLINICAL_ADF211..., CLINIC AL_ADF211...	Oncomine™ Dx Target Test v2.1.1	Completed		Test Report Lab Report View Result ▾

2. Click **OK** in the confirmation dialog box.
In the **Actions** column, the **View Result** button is now active.
3. Click **View Result**, then click [↓](#) **Download Files** to download and view the restored results files.

View results

The run results shown in the screens described in this section can also be downloaded in the form of results files (see “Results files” on page 65).

- In the **Completed Runs & Results** screen, click **View Result** to view the sequencing run results for a sample of interest.

The **Results** screen for the selected sample opens, with the **Loading Metrics** screen displayed. Result categories are arranged vertically on the left side of the screen. The QC status for the run is listed at the top of the screen.

The screenshot shows the 'View Results' interface. At the top, there is a navigation bar with tabs for 'Samples', 'Runs', 'Assay', 'Monitor', and 'Data'. Below this, the 'Completed Runs & Results' section is active, showing 'Verification Runs'. A dropdown menu is set to 'Sample15_MT', and the 'Planned Run Name' is 'PR2_OCP-AMR_RD_MT'. The 'QC Status' is 'Passed'. There are links for '< Back', 'Sign Off', 'Generate Report', and 'Download Files'. On the left, a sidebar lists result categories: 'Loading Metrics', 'QC Report', 'Summary', 'Therapeutic', 'SNV / INDEL', 'Fusion', 'Analytical', and 'Summary'. The main content area displays a table with columns 'Name', 'Count', and 'Percentage'.

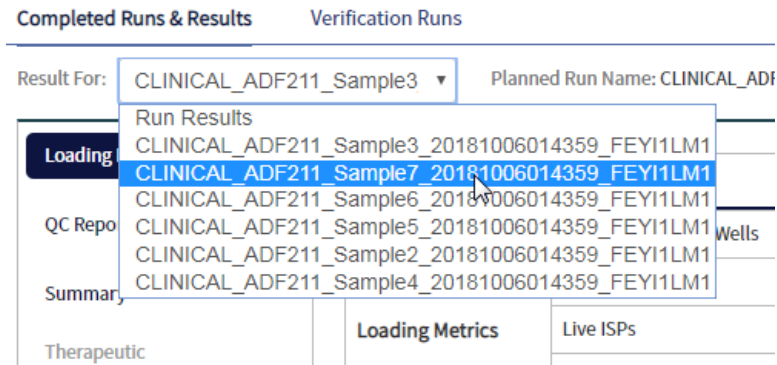
	Name	Count	Percentage
Loading Metrics	Total Addressable Wells	11287275	-
	Wells With ISPs	8533877	75.6 %
	Live ISPs	8527652	99.9 %
	Control ISPs	49654	0.6 %
	Library ISPs	8477998	99.4 %
Filtering Metrics	Filtered: Polyclonal	2567923	30.3 %
	Filtered: Primer-Dimer	16404	0.2 %
	Filtered: Low Quality	1602223	18.9 %
	Usable Library Reads	4291448	50.6 %

- ① QC status
- ② Selected sample
- ③ Result categories

IMPORTANT! A **QC Status** of "Passed" does not guarantee that the genotypes of all analytically relevant variants are determined. See the **Test Result** column in the Lab Report or Test Report for any "No Call" results when interpreting the results.

- Click the links on the left side of the screen to display results of interest, including QC results and results for different categories of variants.

- To view results for another sample in the same run, select the sample from the **Result For:** dropdown list. All samples in a run share the same **Loading Metrics** data. The other result categories are sample-specific.



- To generate the Lab Report and Test Report in other languages, click [Generate Report](#) (see “Generate customized reports” on page 51).
- To download individual results files for further analysis, click [Download Files](#) (see “Results files” on page 65).
- To return to the **Completed Runs & Results** screen, click **< Back**.

Loading metrics

The **Loading Metrics** link in the **View Result** screen displays the following loading and filtering metrics for the run:

Metric	Description
Loading Metrics	
Total Addressable Wells	The total number of wells on the chip – excluded wells.
Wells with ISPs	The number (count) and percentage of chip wells that contain ISPs. The percentage is expressed as a percent of total addressable wells.
Live ISPs	The number (count) and percentage of chip wells containing live ISPs (ISP's templated with library or control fragment), with the percentage expressed as a percent of wells with ISPs.
Control ISPs	The number (count) and percentage of ISPs that have a key signal identifying them as internal controls, with the percentage expressed as a percent of live ISPs.
Library ISPs	The number (count) and percentage of ISPs that have a key signal identical to the library key signal, with the percentage expressed as a percent of live ISPs.

(continued)

Metric	Description
Filtering Metrics^[1]	
Filtered: Polyclonal	ISPs carrying clones from two or more templates, with the percentage expressed as a percent of library ISPs.
Filtered: Primer-Dimer	ISPs with an insert length of less than 8 bp, with the percentage expressed as a percent of library ISPs.
Filtered: Low Quality	ISPs with low or unrecognizable signal, with the percentage expressed as a percent of library ISPs.
Usable Library Reads ^[2]	Number (count) and percentage of library ISPs passing all filters.

^[1] Filtering Metrics only apply to ISPs templated with library fragments, not control fragment.

^[2] Values in the "Filtered:" rows are subtracted from the Library ISPs value (Loading Metrics) to give the Usable Library Reads value.

QC Report

The **QC Report** link displays metrics for the sample libraries and internal controls. This information is also accessible through the **Monitor** tab for the last five runs.

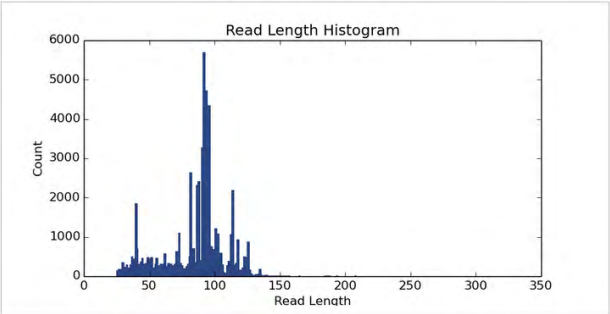
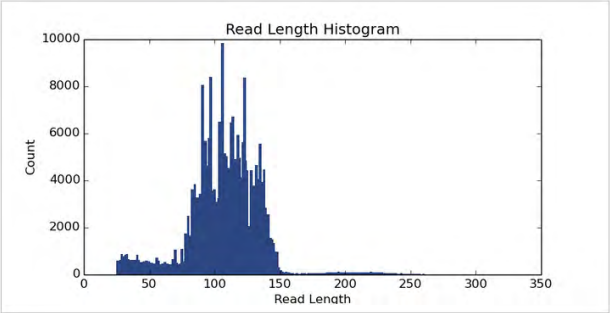
IMPORTANT! The following table describes all the quality control metrics that can be displayed. Whether particular metrics are displayed depends on the assay configuration.

Metric	Description
Library QC Evaluation Metrics	
Library QC: Library RNA	<ul style="list-style-type: none"> • Mean AQ20 Read Length (bp): The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of a library. • Mean Read Length (bp): The average length, in base pairs, of all reads reported for a given library. • Total Mappable Reads: The number of reads that are mapped to the fusion reference file.^[1]
Library QC: Library DNA	<ul style="list-style-type: none"> • Mean AQ20 Read Length (bp): The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of a library. • Mean Read Length (bp): The average length, in base pairs, of all reads reported for a given library. • Percent Reads: The number of library reads normalized by the total addressable wells in a run.

(continued)

Metric	Description
Control QC Evaluation Metrics	
Control QC: CF-1	<ul style="list-style-type: none"> • Base Call Accuracy: $1 - (\text{total number of errors for all positions in the control} / \text{total number of aligned bases})$ for Control Fragment-1 (CF-1) reads. • Key Signal: The average peak signal from the incorporation trace measured across the 1-mer incorporations in the CF-1 key sequences. [2] • Mean AQ20 Read Length (bp): The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of CF-1. • Percent Reads: The number of all usable library reads that aligned with the CF-1 sequence divided by the total number of addressable wells.
Control QC: DNA Control	<ul style="list-style-type: none"> • COSMIC ID variant calls and allelic frequencies: The individual variant positions and wild-type positions that are assessed in the DNA control reagent for presence or absence of the variant. • Mean AQ20 Read Length (bp): The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of a control. • Percent Reads: The number of all usable library reads that aligned with the control fragment sequence divided by the total number of addressable wells.
Control QC: RNA Control	<ul style="list-style-type: none"> • Fusion calls and number of reads: The individual fusion calls and number of reads that are assessed in the RNA control reagent for the presence or absence of the fusion. • Total Mappable Reads: the number of reads that are mapped to the fusion reference file.
Control QC: DNA NTC	<ul style="list-style-type: none"> • Hotspot Calls: The total number of hotspots where a call was made in the no-template control.
Control QC: RNA NTC	<ul style="list-style-type: none"> • Total Fusion Calls: The total number of fusion locations where a call was made in the no-template control. • Total Mappable Reads: The number of reads in the no-template control that are mapped to the fusion reference file.

(continued)

Metric	Description
Histogram of Read Length ^[3]	<ul style="list-style-type: none"> A library-specific read-length histogram: <div style="display: flex; flex-direction: column; align-items: center;"> <div style="display: flex; align-items: center; margin-bottom: 10px;"> RNA  </div> <div style="display: flex; align-items: center;"> DNA  </div> </div>

^[1] May not equal the sum of the individual fusions, since some detected fusions are not included in the fusion BED file.

^[2] The minimum Key Signal value for the CF-1 control (i.e., the Templating Control) is set to 0 in all preinstalled assays and new custom assays in Torrent Suite™ Dx Software version 5.8 and later. Custom assays created in previous versions of the software may have a higher Key Signal value for the control. If a run using an older custom assay fails QC due to this metric, reanalyze the run using a minimum Key Signal of 0 for the control. For future runs, create a new custom assay based on the old assay, changing the minimum Key Signal to 0.

^[3] Not displayed in the Monitor tab.

Summary

The **Summary** link displays the following information for the run:

Field	Description
Run and Configuration Summary	
Run Name	The name of the run.
Assay Name	The name of the assay.
Reference Genome	The reference genome used for analysis.
Target Region	The name of the targeted regions BED file used.
Hotspot Regions	The name of the hotspot regions BED file used.
Fusion Reference	The name of the fusion reference used for analysis, when applicable.
Fusion Panel	The name of the fusion panel used, when applicable.

(continued)

Field	Description
Library Name	The names of the DNA and DNA libraries prepared from the sample, entered during library batch preparation.
Amplicon Summary	
Number of Targets	The number of amplicons in the panel.
On Target Reads	The number of reads mapped to the target amplicons.
Percent On Target Reads	The percentage of reads mapped on target to total reads.
Percent Full Length On Target Reads	The percentage of full length reads mapped on target to total reads.
Average Coverage	The ratio of the depth of coverage at each base in the target region to the length of the target region.
Target Coverage at 20X	The percentage of base positions with depth of coverage $\geq 20X$ in the target region.
No Strand Bias	The percentage of bases with a strand bias between 30% and 70%.
Coverage Uniformity	The ratio of passed number of targets to total number of targets, where passed number of targets is the number of amplicons that have at least 0.2X mean coverage.
Variant Summary^[1]	
Number of SNVs/MNVs	The number of single- and multi-nucleotide variations (SNVs/MNVs).
Number of INDELS	The number of insertions or deletions.
Number of Fusions	The total number of fusion calls.

^[1] Includes Therapeutic, Analytical, and Level2 variants in the **View Result** screen.

Therapeutic variant results

Therapeutic variant results for the selected sample are provided under the **Therapeutic** section of the left navigation bar in the **Results** screen. Depending on the assay, results for these variants may be divided into multiple subscreens, each with a separate link (**SNV/INDEL**, **Fusion**, etc.).

The results are determined by the reference sequences installed on the server, the QC controls used in the run, the assay used for the run, and any Reporting Gene List associated with that assay.

Columns in the screen or subscreens are described in the following table.

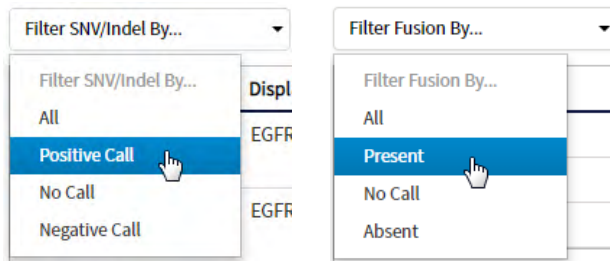
Column	Description
Summary	
Gene Fusion Present Variants Present	The therapeutic gene fusion or SNV/INDEL variant detected in the sample
Associated Therapy	The therapy indicated for each variant listed
SNV / INDEL	
Gene	The gene name, which provides a link to the View Annotation Sources popup with additional information about the HotSpot ID (see "View annotation sources" on page 63)
Display Name	The gene name with the exonic location of the deletion or insertion
Amino Acid Change	The amino acid change using HGVS-nomenclature
Nucleotide Change	Identification of the DNA-level nucleotide change using HGVS-nomenclature
Test Result	The presence or absence of the variant
Allele Frequency	The most likely frequency of the variant allele ^[1]
HotSpot ID	The name of the hotspot as defined in the BED file
Locus	The one-base position of the variant in the reference genome
Type	The type of variation detected: <ul style="list-style-type: none"> • snv/mnv (single nucleotide variation/multi-nucleotide variation) • ins (insertion) • del (deletion) • complex (Variant that includes both a deletion and insertion of DNA sequences relative to the reference sequence)
Genotype	The genotype at the locus
Ref	The reference base or bases at the locus
Quality Score	The relative probability of either the "reference" hypothesis interval [0,cutoff) or the "variant" hypothesis interval [cutoff,1], on a Phred scale (-10*log10). This provides a measure of the strength of the evidence for the variant call. A higher score means more evidence for the call. Quality scores are capped at 100.
Coverage	The number of reads covering the variant position after down-sampling
Fusion	
Gene	The gene that regulates expression of the gene fusion

(continued)

Column	Description
Display Name	The gene name of the fusion
Read Count	The number of valid reads aligned to the specific fusion sequence
Test Result	The presence or absence of that fusion variant

^[1] The allele frequency is the most likely variant frequency in the reads after corrections are made for probable errors. The software uses this corrected frequency and the uncertainty in the observations to calculate the probability that the variant frequency in the sample falls within defined intervals that can be set by the user. The user sets a threshold frequency for deciding between genotypes: this defines intervals $[0, c][c, 1-c][1-c, 1]$ corresponding to the three diploid genotypes (reference, heterozygous, homozygous) respectively, where “c” is the minimum variant frequency set during the creation of the assay. The software reports the genotype corresponding to the interval with the highest probability of containing the variant frequency in the sample. In most cases, the reads are highly accurate and do not have a high probability of error. In such cases, the observed frequency and the most likely frequency are similar, and are contained within a single interval leading to a high quality genotype. In some cases, if very few reads are observed or the error rate is high, the observed counts and the most likely counts can be different, and the uncertainty in the real sample frequency can be high. When this happens, it is often impossible to exclude a heterozygous population in the sample, and the genotype is assigned to be heterozygous, since a significant portion of the probability falls in that interval.

Note: You can filter the variant lists using options in the **Filter SNV/Indel By...** and **Filter Fusion By...** dropdown lists in the upper left corner of the results report.



Analytical variant results

Analytical variant results for the selected sample are provided under the **Analytical** section of the left navigation bar in the **Results** screen. Depending on the assay, results for these variants may be divided into multiple subscreens, each with a separate link (**SNV/INDEL**, **Fusion**, etc.).

The results are determined by the reference sequences installed on the server, the QC controls used in the run, the assay used for the run, and any Reporting Gene List associated with that assay.

Columns in the screen or subscreens are described in the following table.

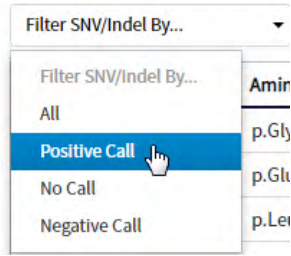
Column	Description
Summary	
Variants Present	Lists the analytical SNV or INDEL variants detected in the sample
SNV / INDEL	
Gene	The gene name, which provides a link to the View Annotation Sources popup with additional information about the HotSpot ID (see “View annotation sources” on page 63)

(continued)

Column	Description
Amino Acid Change	The amino acid change using HGVS-nomenclature
Nucleotide Change	The DNA-level nucleotide change using HGVS-nomenclature
Test Result	The presence or absence of the variant
Allele Frequency	The most likely frequency of the variant allele
HotSpot ID	The name of the hotspot as defined in the BED file
Locus	The one-base position of the variant in the reference genome
Type	The type of variation detected: <ul style="list-style-type: none"> • snv/mnv (single nucleotide variation/multi-nucleotide variation) • ins (insertion) • del (deletion) • complex (Variant that includes both a deletion and insertion of DNA sequences relative to the reference sequence)
Genotype	The genotype at that locus
Ref	The reference base or bases at the locus
Quality Score	The relative probability of either the "reference" hypothesis interval [0,cutoff) or the "variant" hypothesis interval [cutoff,1], on a Phred scale (-10*log10). This provides a measure of the strength of the evidence for the variant call. A higher score means more evidence for the call. Quality scores are capped at 100.
Coverage	The number of reads covering the variant position after down-sampling
Fusion^[1]	
Gene	The gene that regulates expression of the gene fusion
Display Name	The gene name of the fusion
Read Count	The number of valid reads aligned to the specific fusion sequence
Test Result	The presence or absence of that fusion variant

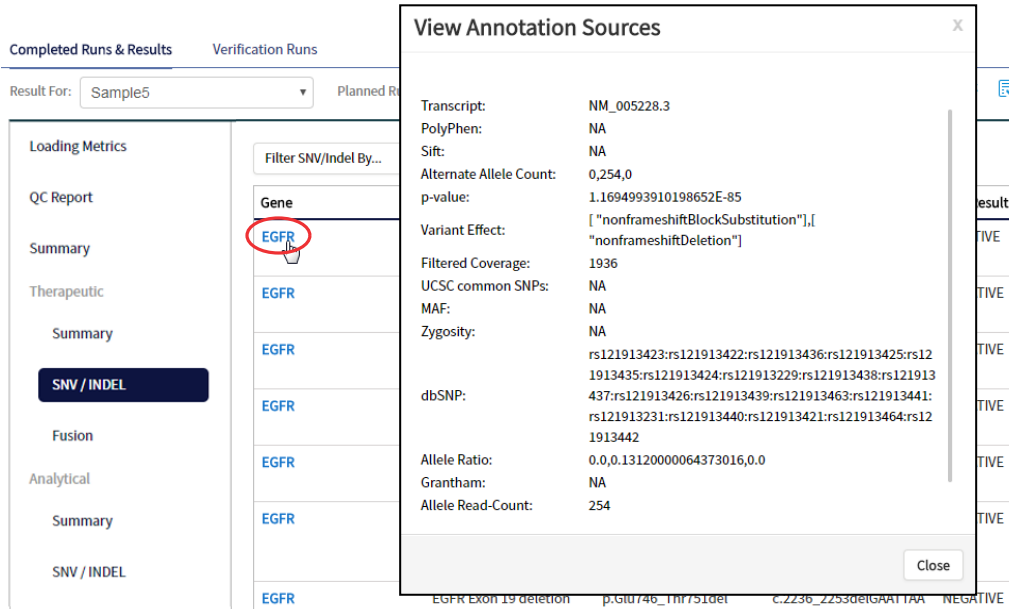
[1] Note that for the OncoPrint™ Dx Target Test, no analytical fusions are reported except for expression controls.

Note: You can filter the variant list using options in the **Filter SNV/Indel By...** dropdown list in the upper left corner of the results report.



View annotation sources

You can view additional information for each hotspot ID listed under the **SNV/Indel** links of the **Therapeutic** and **Analytical** sections in the **View Results** screen. The gene name in the **SNV/Indel** report is a link that opens the **View Annotation Sources** dialog box, which provides information for the particular hotspot.



Example of transcript information in the VCF

The COSM1074639 hotspot ID has multiple transcripts associated with it, which include a mutation from 'A' to 'C'. These are listed in the VCF file as shown below:

chr6 152419923 COSM1074639;COSM1074637 A C,G 122.19 PASS
 AF=0,0;AO=0,0;DP=1116;FAO=0,0;FDP=1116;FR=.;FRO=1116;FSAF=0,0;FSAR=0,0;FSRF=661;FSRR=455;F
 WDB=0.00426152,-
 0.0087378;FXX=0;HRUN=1,1;HS;LEN=1,1;MLLD=103.308,103.438;QD=0.437949;RBI=0.0394298,0.03043
 16;REFB=-2.22051E-5,-2.32713E-5;REVB=-0.0391988,-
 0.0291502;RFO=1115;SAF=0,0;SAR=0,0;SRF=661;SRR=454;SSEN=0,0;SSEP=0,0;SSSB=-6.44874E-8,-
 6.44874E-
 8;STB=0.5,0.5;STBP=1,1;TYPE=snp,snp;VARB=0,0;OID=COSM1074639,COSM1074637;OPOS=152419923,
 152419923;OREF=A,A;OALT=C,G;OMAPALT=C,G;FUNC=[

{'normalizedRef':'A','transcript':'NM_001122742.1','grantham':'144.0','gene':'ESR1','location':'exonic','ori
 gAlt':'C','origPos':'152419923','origRef':'A','normalizedPos':'152419923','exon':'10','function':'missense',
 protein':'p.Tyr537Ser','normalizedAlt':'C','gt':'neg','codon':'TCT','coding':'c.1610A>C'},

{'normalizedRef':'A','transcript':'NM_001122742.1','grantham':'194.0','gene':'ESR1','location':'exonic','ori
 gAlt':'G','origPos':'152419923','origRef':'A','normalizedPos':'152419923','exon':'10','function':'missense',
 protein':'p.Tyr537Cys','normalizedAlt':'G','gt':'neg','codon':'TGT','coding':'c.1610A>G'},

{'normalizedRef':'A','transcript':'NM_001122741.1','grantham':'144.0','gene':'ESR1','location':'exonic','ori
 gAlt':'C','origPos':'152419923','origRef':'A','normalizedPos':'152419923','exon':'9','function':'missense',
 protein':'p.Tyr537Ser','normalizedAlt':'C','gt':'neg','codon':'TCT','coding':'c.1610A>C'},

{'normalizedRef':'A','transcript':'NM_001122741.1','grantham':'194.0','gene':'ESR1','location':'exonic','ori
 gAlt':'G','origPos':'152419923','origRef':'A','normalizedPos':'152419923','exon':'9','function':'missense',
 protein':'p.Tyr537Cys','normalizedAlt':'G','gt':'neg','codon':'TGT','coding':'c.1610A>G'},

{'normalizedRef':'A','transcript':'NM_001122740.1','grantham':'144.0','gene':'ESR1','location':'exonic','ori
 gAlt':'C','origPos':'152419923','origRef':'A','sift':'0.0','normalizedPos':'152419923','exon':'9','function':'mi
 ssense',protein':'p.Tyr537Ser','gt':'neg','normalizedAlt':'C','codon':'TCT','polyphen':'0.979','coding':'c.161
 0A>C'},

{'normalizedRef':'A','transcript':'NM_001122740.1','grantham':'194.0','gene':'ESR1','location':'exonic','ori
 gAlt':'G','origPos':'152419923','origRef':'A','sift':'0.0','normalizedPos':'152419923','exon':'9','function':'mi
 ssense',protein':'p.Tyr537Cys','gt':'neg','normalizedAlt':'G','codon':'TGT','polyphen':'0.998','coding':'c.16
 10A>G'},

Results files

The following files can be downloaded from the **Results Report** window. To download the files, click **Download Files**, select the files to download, then click **Download**.

File name	Description
Test Report	A report of the completed analysis in PDF format
Lab Report	A clinical lab report of the completed analysis in PDF format; includes both clinical and analytical results.
PlannedRun-AuditTrail.pdf	Contains all audit records pertaining to the Planned Run.
Info.csv	Contains information about the run and analysis, such as software, sequencing information, instrument information, analysis information, QC details etc.
<RNABarcode>_rawlib.basecaller.bam	Unmapped RNA BAM File; output of base calling, contains unmapped reads.
Snvindel.tab	A tab-delimited file that contains information about non-targeted SNVs and indels
<RNABarcode>_rawlib.basecaller_alignments.bam	Mapped RNABarcode BAM file; output after reads have been mapped to the fusion reference.
Target_Summary.tab	A tab-delimited file that contains a targeted test results summary
<RNABarcode>_rawlib.basecaller_alignments.bam.bai	Mapped RNABarcode BAM index file
<RNABarcode>_rawlib.basecaller.fastq	FASTQ file generated from unmapped BAM file of the RNA barcode used.
<DNABarcode>_rawlib.basecaller.bam	Unmapped DNA barcode BAM file; output of base calling, contains unmapped reads.
raw_peak_signal	Key signal gives the percentage of LiveSPs with a key signal that is identical to the library key signal.
<LibPrepID>_<analysisID>.final.vcf	A VCF file containing all the variants detected as a result of the analysis, along with information such as test result, read count, gene name, quality scores, etc.
Summary.tab	A tab-delimited file that contains the on-targeted test results summary
<LibPrepID>_rawlib.stats.cov.txt	Amplicon statistics file


(continued)

File name	Description
Fusion.tab	A tab-delimited file that contains non-targeted (analytical) fusion details in a table format. Note: The information displayed in the file for each isoform of a particular fusion is identical, because specific isoform and locus information is not included in this table. Detailed isoform and locus information is available in the <LibraryPrepID>.<AnalysisID>.final.vcf file, available under Download Files .
readLenHisto.png	Gives the read-length distribution of FASTQ files in the form of a histogram. A thumbnail histogram of the read lengths for a particular barcode.
<LibPrepID>_rawlib.bam.bai	Mapped DNA barcode BAM index file (index file of DNA barcode-mapped BAM file)
Basecaller.log	Base Caller log file
analysis.log	Analysis log file
sigproc.log	Signal processing log file
Bead_density_contour.png	Loading density image; a pseudo-color density image of the Ion Chip plate showing percent loading across the physical surface
<DNABarcode>_rawlib.basecaller.fastq	FASTQ file of the DNA barcode used
Target_cnv.tab	Targeted CNV detail table (CNV results for targeted variants from a sequencing run) Note: For use with IVD tests that include CNV reporting.
cnv.tab	Non-targeted CNV detail table (analytical CNV results from a sequencing run) Note: For use with IVD tests that include CNV reporting.
Target_fusion.tab	A tab-delimited file that contains targeted (clinical) fusion details in a table format
<LibPrepID>_rawlib.bam	Mapped DNA barcode BAM File; output after mapping reads to reference.
Iontrace_Library.png	Key incorporation trace image showing the average signal readings for flows of the bases T, C, and A in the library key.
rawtf.basecaller.fastq	FASTQ file for the test fragment

Sign the run results

In the **View Result** screen, Managers/Administrators can provide their electronic signature on the run results. The signature information appears in the **QC Report** in the **View Result** screen, and in the downloaded Test Report and Lab Report PDFs.

Multi-language support for PDF report generation is provided. By default reports are generated in the language that is selected in the **Report Template** used. When reports are generated in multiple languages, **Sign Off** occurs only in the report of the default language.

1. At the top of the **View Result** screen, click  **Sign Off**, then enter your user name, password, and comments in the dialog box. Fields identified with a red asterisk (*) are required fields.
2. In the **Footer Field**, enter any text.
3. Click **Sign Off** to confirm your electronic signature.


Files in the Reports folder

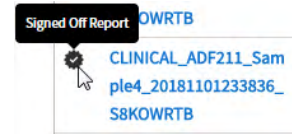
When a manager- or administrator-level user signs a report, a folder named with the Sample ID is created in the Reports folder on the server (`/results/analysis/output/reports`), and the following files are copied into it:


Info file (.csv)	Non-targeted Test Results Summary (.tab)
Signal processing log file (.log)	Targeted SNV/INDEL Detail Table (.tab)
Targeted Fusion Detail Table (.tab)	Amplicon Stats (<code>_rawlib.stats.cov.txt</code>)
Targeted Test Results Summary (.tab)	RNA FASTQ File (.fastq)
Analysis log file (.log)	RNA Mapped BAM file (.bam)
VCF file (.vcf)	RNA Unmapped BAM file (.bam)
DNA Mapped BAM file (.bam)	Test Fragment FASTQ File (.fastq)
DNA Unmapped BAM file (.bam)	Read Length Histogram (.png)
Key Signal	Test PDF Report (<i>optional</i>) (.pdf)
Key Incorporation Trace (.png)	Lab PDF Report (.pdf)
Fusion Detail Table (.tab)	Planned Run Audit (.pdf)
DNA Mapped BAM Index file (.bam.bai)	Basecaller command files (.json)
RNA Mapped BAM Index file (.bam.bai)	checksum file
DNA FASTQ File (.fastq)	Pipeline commands (<code>_pipeline.json</code>)
Base Caller Log File (.log)	Experimental log file (<code>_final.txt</code>)
Targeted CNV Detail Table (.tab) ^[1]	Wells with beads (<code>_beadogram.png</code>)
Non-targeted CNV Detail Table (.tab) ^[1]	Bead find stats file (.stats)
SNV/INDEL Detail Table (.tab)	Loading Density Figure (.png)

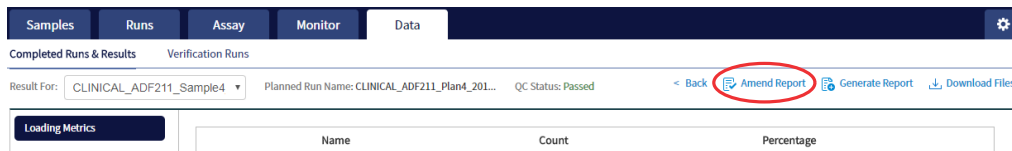
^[1] For use with IVD tests that include CNV reporting.

Amend a signed-off report

Managers and administrators can amend a report within 30 days of signing off. A sample that has a signed-off report has a  symbol to the left of the Sample ID in the **Completed Runs & Reports** screen. You can change sign-off or laboratory comments, or the text that appears in the report footer. Follow these steps to amend a signed-off report:



1. In the **Completed Runs & Reports** screen, click **View Result** for the sample of interest.
2. Click  **Amend Report**.



3. In the **Electronic Signature** dialog box, make your changes in the appropriate fields, then click **Amend Report**.
4. Click **Test Report** or **Lab Report** to download the amended report.

Repeat testing

Based upon quality control results, users can determine whether a library requires repeat testing. Refer to the following table to determine the best course of action. See the *OncoPrint™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide* for the details of each procedure.

Quality Control			Pooling Error ^[1]	Retest Sample	Procedures to follow
Library	Run	Control			
Pass	Pass	Pass	No	N/A	Sign off on Library.
Fail	Pass	Pass	No	gDNA	Starting from library preparation. ^[2]
Fail	Pass	Pass	Yes	Library	Starting from preparing the library pool. If the retest of the sample library fails, retest the gDNA sample starting from library preparation.
Fail	Pass	Fail	No	gDNA	Starting from library preparation.
Fail	Pass	Fail	Yes	Library	Starting from preparing the library pool. If the retest of the library fails, retest the gDNA sample starting from library preparation.
Fail	Fail	Pass	No	gDNA	Starting from library preparation.

(continued)

Quality Control			Pooling Error ^[1]	Retest Sample	Procedures to follow
Library	Run	Control			
Fail	Fail	Pass	Yes	Library	Starting from preparing the library pool. If the retest of the library fails, retest the gDNA sample starting from library preparation.
Pass (all samples)	Pass	Fail	No	Pooled Library	Starting from the pooled sample libraries, repeat template preparation. If the CF-1 fails QC a second time, reprepare the library pool.
Pass (all samples)	Fail	Fail	No	Pooled Library	Starting from the pooled sample libraries, repeat template preparation. If the CF-1 fails QC a second time, reprepare the library pool.
Pass (all samples)	Fail	Pass	No	Pooled Library	Starting from the pooled sample libraries, repeat template preparation. If the CF-1 fails QC a second time, reprepare the library pool.

^[1] Pooling error is defined as no reads detected for a Barcode ID.

^[2] **IMPORTANT!** If a library fails Library QC in a second run that passes both Run and Control (CF-1) QC metrics, the sample is considered invalid.

Data files and flow

During an Ion PGM™ Dx Sequencer run, sequence raw data (DAT) files are transferred to the Ion Torrent™ Server via a network cable in a process controlled by the ionCrawler service. After the data from the initial flows on the sequencer are available on the server, the Torrent Suite™ Dx Software begins processing the data, producing the <1> wells file. Basecalling is performed on the <1> wells file data, producing an unmapped BAM (uBAM) file. Subsequent analysis produces mapped reads (BAM) and variant calls (VCF) files.

The following table shows the flow of data and typical file size generated as the Torrent Suite™ Dx Software processes data from an Ion 318™ Dx Chip.

Process	File Type	File Size
		Ion 318™ Dx Chip
Sequencing ^[1]	DAT	~350 GB
▼		
Signal Processing	Wells	12 GB
▼		
Base Calls (reads)	uBAM	1.5 GB
▼		
Mapped Reads	BAM	4.3 GB
▼		
Variant Calls	VCF	120 Kb

^[1] The sequencing raw data (DAT) files are deleted from the server 72 hours after data analysis to conserve Ion Torrent™ Server disk space.

Note: Data from approximately 200 sequencing runs using an Ion 318™ Dx Chip can be accommodated on an Ion Torrent™ Server before disk space becomes a limiting factor and data archiving is required. See “Data Management (administrator)” on page 84 for more information.

Verification runs on the Ion PGM™ Dx System

Verification runs are sequencing runs performed during Ion PGM™ Dx System installation by Thermo Fisher Scientific support specialists to validate the performance of the instruments.

Verification runs are performed using Torrent Suite™ Dx Software. Verification runs can also be performed using Torrent Suite™ Assay Development Software, but are not required.

You can view verification run reports, but only qualified support specialists can perform the runs. Under the **Data** tab, in the **Verifications Runs** screen, you can search, filter, sort, and view completed verification runs and reports.

The following information is displayed in the **Verification Runs** screen:

Column	Description
Planned Run	<p>The name of the run, created when the run was created. Click the name to open the Review Planned Run dialog box.</p> <ul style="list-style-type: none"> Click Audit to display the list of users who created/edited the Planned Run. You can export and print information from the list from the Audit Trail dialog box. Click CSA (Customer Support Archive) to download all the instrument log files, which can be useful for troubleshooting.
Field Engineer Name	The name of the support specialist who performed the run.
Instrument Name	The name of the validated instrument.
Templating Completion	The completion date and time of the Ion OneTouch™ Dx System run. Click the link to open a dialog box that includes additional information about the instrument, operator, and template kit used.
PGM Completion	The completion date and time of the Ion PGM™ Dx System run. Click the link to open a dialog box that includes additional information about the sequencer, operator, sequencing kit, and chip used.
Analysis Completion	The completion date and time of the post-sequencing run analysis.
Run status	The current status of the full run, including analysis.
QC Status	Indicates whether a run passed or failed, based on the QC metrics selected for the assay.
PQ Report	If the results of the run have been approved and signed off by a Manager or Administrator, the approved PQ Report are listed in this column.
Actions	<ul style="list-style-type: none"> Click View Results to view the results of completed runs that have not yet been approved. Managers and Administrators can also sign off on the results after viewing them. Click View Report to view a report of the run.

Verification run results

Under the **Data** tab, in the **Verifications Runs** screen, click **View Results** in the **Actions** column to view the performance qualification (PQ) report for a verification run.

Note: Manager- and administrator-level users can sign off on this report, at which point it becomes a locked PQ Report. However, we recommend that only qualified support specialists sign off on PQ reports.

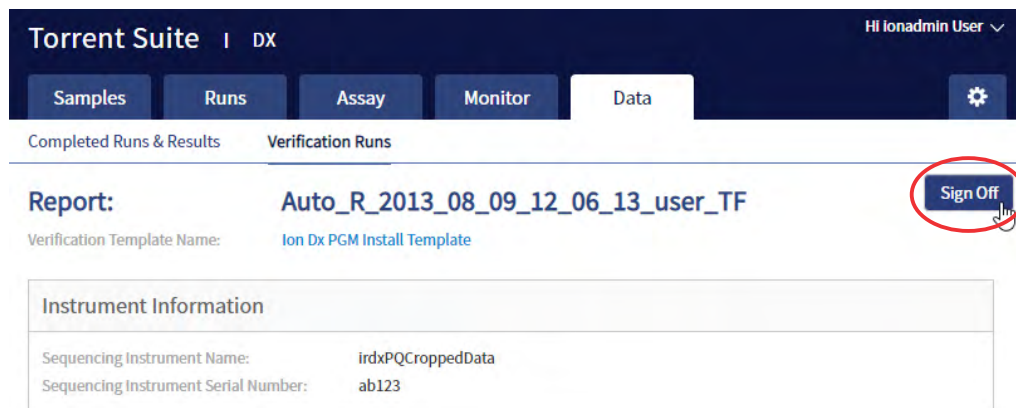
In addition to basic information about the verification run, the following data are reported for the controls that are used in the run:

Metric	Description
Base Call Accuracy	1 – (total number of errors for all positions in the control / total number of aligned bases)
Mean AQ20 read length (bp)	The average length, in base pairs, at which the error rate is $\leq 1\%$ for all aligned reads of a control fragment
Percent Reads	The number of all usable library reads that align with the control fragment sequence, divided by the total number of addressable wells

Sign verification run reports (manager/administrator)

Manager- and administrator-level users can sign results reports for verification runs. However, we recommend that only qualified support specialists sign PQ reports.

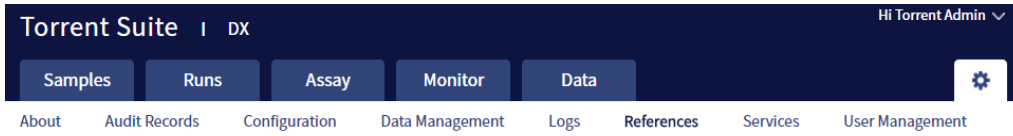
1. Click **Data** ▶ **Verification Runs**.
2. In the **Report** window, click **Sign Off** in the upper right corner above the report.



3. In the **Sign off Electronic Signature** dialog box, enter your user name, password, and any comments, then click **Sign Off**.
4. The report is locked and listed in the **PQ Report** field for the run in the **Verification Runs** screen.
5. Click the PQ report name to view, print, or download the report in a PDF file.



System administration and management



In the **tab**, Managers and Administrators can:

- View, create, and manage reference files, including genome references, panel and hotspot BED files, primers, and others.

Note: The Oncomine™ Dx Target Test assay is locked. None of the files for this assay can be edited; they can only be viewed.

- View system information and settings.

In addition, Administrators can:

- View and export audit records.
- View and change configuration settings.
- Create and manage user accounts.
- Manage log files.

Access to tab functions by user level


In the **tab**:

Users at this level...	Can access...
Operator	<p>About: More Information and Assistance</p> <ul style="list-style-type: none">• Support Contact Information: a link directing users to the thermofisher.com/technical-resources website.• Instrument Diagnostics: performs a diagnostic check of the Ion PGM™ Dx Sequencer.• End User License Agreement: opens the End User License Agreement (EULA).

(continued)

Users at this level...	Can access...
<p>Manager</p>	<p>Operator functions plus:</p> <p>References: displays and allows management of:</p> <ul style="list-style-type: none"> • Reference Sequences • Panels • Hotspots • DNA Barcodes • Fusion Reference • Fusion Panel • Reporting Gene List • Control Fragments • Primers <p>Services: displays system status report for:</p> <ul style="list-style-type: none"> • Jobs Server • Active Jobs • ionCrawler Service Details • Smart Monitoring Service Details
<p>Administrator</p>	<p>Manager functions plus:</p> <p>Audit Records: perform an audit: search, sort, view, export, and print any record.</p> <p>Configuration: Configure settings for the:</p> <ul style="list-style-type: none"> • Network • Email • Instruments • Lab Information • Software Updates <p>Data Management: Manage archiving, view available disk space, and manually delete incomplete runs.</p> <p>Logs: Manage retention of instrument logs and Torrent Suite™ Dx Software logs.</p> <p>User Management: Add, assign user level, and edit user information.</p>

References (manager/administrator)

Under the  tab in the **References** screen, manager- and administrator-level users can access the following reference files under the subtabs.

Note: These files are defined by the OncoPrint™ Dx Target Test Assay Definition File. They are locked and can only be viewed or exported.


- Reference sequence files
- Panel files (BED format)
- Hotspots files (BED format)
- DNA barcode sequences
- Fusion reference sequences (FASTA format)
- Fusion panel files (BED format)
- Reporting Gene Lists (Microsoft Excel™ format)
- Control fragments
- Primer sequences (FASTA format)

Manage reference sequences

Manager- and administrator-level users manage the DNA reference sequence files that are available in the software


Manage panel files

Manager- and administrator-level users can view and manage the panel BED files that are available in the software.

1. Click  ► **References**
2. Select the **Panels** tab.
3. To export a panel file, in the **Actions** column, click **Export**.


Manage hotspot files

Manager- and administrator-level users can view and manage the hotspot BED files that are available in the software

1. Click  ► **References**
2. Select the **Hotspots** tab.
3. To export a hotspot file, in the **Actions** column click **Export**.

View DNA barcodes


Manager- and administrator-level users can view the barcode IDs and sequences for each barcode set that is available in the software.

1. Click  ► **References**.
2. Select the **DNA Barcodes** tab.

3. In the **DNA Barcodes** screen, click the name of a barcode set in the list.
A window opens showing the name and sequence of each barcode in the set.
4. Click **Back** to return to list on the **DNA Barcodes** screen.

Manage the Reporting Gene List

Manager- and administrator-level users can view and manage reporting gene lists for the Oncomine™ Dx Target Test. A gene list is the list of the genes and mutations that are used to generate reports for an assay.

1. Click  **References**
2. Select the **Reporting Gene List** tab.
3. In the **Reporting Gene List** screen, to view the contents of the gene list:
 - a. In the **Name** column, click the name of the gene list.
A screen opens showing the gene names, mutation IDs, and other information about the genes in the list.
 - b. Click **Back** to return to the **Reporting Gene List** screen.
4. To export the gene list, in the **Actions** column, click **Export**.


Control fragments

Manager- and administrator-level users can review information about control fragments. These internal controls are predefined and cannot be modified by users.

View the control fragment information by clicking  **References** **Control Fragments**. The **Control Fragments** screen provides the following information.

Column	Description
Name	The name of the control fragment.
Sequence	The single letter nucleotide sequence of the control fragment.
Control	The process in which the control fragment is used.
Status	Indicates that the control fragment information is locked and cannot be modified.

Manage primers

Under the  tab, in the **References** screen on the **Primers** tab, manager- and administrator-level users can view the primer sequences in the software.

Services (manager/administrator)

On the **Services** screen, manager- and administrator-level users can view the status and details of active jobs, ionCrawler service, Smart Monitoring service, and the jobs server. Viewing this information can be useful when troubleshooting error messages during an active job or why an analysis failed.

Services

Service	Function
Jobs Server	Lists server information and active (running) software.
Active Jobs	<ul style="list-style-type: none"> Lists active and queued analysis jobs on the Ion Torrent™ Server. When no job is currently active, displays a “No active jobs” message.
ionCrawler Service Details	Displays information about the data transfer process from the Ion PGM™ Dx Sequencer to the Ion Torrent™ Server.
Smart Monitoring Service Details	Allows remote monitoring of Ion PGM™ Dx System.

View or terminate an active job

Manager- and administrator-level users can view jobs that are active on the server and terminate an active job if required. A job is active when the status indicates Job is running.

1. Click  ▶ **Services**
2. In the **Active Jobs** section, click **Terminate** to stop the active job.

Active Jobs


Job ID	Job Name	Job Type	Status Message	
6592	job-0	Analysis	Job is running	Terminate
6595	job-1	Analysis	Job is running	Terminate

A confirmation message opens, “Are you sure you want to terminate Job Name ?”

3. Click **Yes** to terminate the active job.

Disable or enable the ionCrawler service

Manager- or administrator-level users can manage the IonCrawler service. The ionCrawler service typically remains enabled at all times, but can be disabled for remote service troubleshooting.

- To disable the service:
 - a. Click  ▶ **Services**.

- b. In the **ionCrawler Service Details** section, click **Stop**.


ionCrawler Service Details

Status: Online **Stop**

- c. In the confirmation message, click **Yes** to stop the service.
The **Stop** option changes to **Start**.

ionCrawler Service Details

Status: Offline **Start**



- To enable the service:
 - a. Click  **Services**.
 - b. In the **ionCrawler Service Details** section, click **Start**.
 - c. In the confirmation message, click **Yes** to start the service.
The **Start** option changes to **Stop**.

Enable or disable the Smart Monitoring service

Manager- and administrator-level users can manage the Smart Monitoring service.

The Smart Monitoring service allows Thermo Fisher Scientific personnel to remotely monitor the status of the Ion PGM™ Dx System through an internet connection. Smart Monitoring employs multiple layers of security, including a Secure Sockets Layer (SSL) and Lightweight Directory Access Protocol (LDAP) authentication, to provide real-time troubleshooting and problem resolution for the Ion PGM™ Dx System.

The Smart Monitoring service is inactive by default and can be enabled by manager- or administrator-level users. The service can also be disabled by manager- or administrator-level users, if needed.

- To enable the service:
 - a. Click  **Services**.
 - b. In the **Smart Monitoring Service Details** section, click **Start**.
 - c. In the confirmation message, click **Yes** to enable the Smart Monitoring service.
The **Start** option changes to **Stop**.
- To disable the service:
 - a. Click  **Services**.
 - b. In the **Smart Monitoring Service Details** section, click **Stop**.
 - c. In the confirmation message, click **Yes** to disable the Smart Monitoring service.
The **Stop** option changes to **Start**.

Audit records (administrator)

In the **Audit Records** screen, administrator-level users can use the tools that are provided to review, sort, export, and print audit records.

All components in a diagnostic assay must be uniquely identified, and the identification must be stored so that the record can be audited.

Note: Library Batches and Planned Runs created in a batch with other objects in a single LIMS transaction do not have a "Create" action listed in the **Audit Records** screen. However, you can view their complete audit record from the **Libraries** and **Planned Runs** screens, and any subsequent actions that are performed on them are listed in the **Audit Records** screen. A Library Batch or Planned Run created individually in a LIMS transaction is listed.

Search audit records

Administrator-level users can use the toolbar to search for an existing audit record.

The image shows a search toolbar with the following elements from left to right: a 'Start Date' field with a calendar icon, an 'End Date' field with a calendar icon, a 'First Name Last N.' dropdown menu, a 'Select Action' dropdown menu, a 'Sample' dropdown menu, a 'Search' button, and a 'Clear' button.

1. Click ► **Audit Records**.
2. In the **Audit Records** screen, select the search parameters:
 - a. To select a date range, click next to **Start Date** and **End Date**.
If an end date is not selected, the search results include all records from the start date through the current date.
 - b. To limit the results to actions that are performed by a specific user, select the user from the **First Name Last Name** list.
 - c. To limit the results to specific actions, select from the **Select Action** list.
 - d. To limit the results to a particular sample, select from the **Sample** list.
3. Click **Search**.
4. Click **Clear** to return to the complete list of records.

Sort audit records

In the **Audit Records** screen, the list of audit records is displayed with the oldest record on top by default. Administrator-level users can sort the **Audit Records** screen to find audit records. To return to the default display, click **Audit Records** or **Clear**.


1. Click ► **Audit Records**.
2. In the **Audit Records**, click the column heading of interest.
The list of records is reordered based on the heading name selected. **User**, **Action Performed**, and **Data Object Name** sort alphabetically (**A** ► **Z**).

Click the **Timestamp** heading to reverse the default setting with the most recent record on top.

3. Click the column heading a second time to reverse the order of records displayed.
4. Click **Audit Records** or **Clear** to return to the default display.


Export and print Audit Records

The **Export** function generates a print-ready PDF file of the selected Audit Record.


1. Select the records to be exported by clicking the checkbox next to the record of interest. Select all the records on the screen by selecting the checkbox above the column.
2. Click  **Export**.
A PDF is generated. Depending on your browser settings, the software automatically downloads the file or prompts you to open or save the file.
3. Open the PDF in an appropriate viewer to print.

Update the Audit Configuration

Audit Configuration allows the Administrator to require that a reason for the change is included as part of changing designated objects. To update the Audit Configuration:

1. Click  **Audit Configuration**.
2. In the **Audit Configuration** dialog box, click the **Require Reason** checkbox next to the Data Object Name.
3. Click **Save**.
To remove a reason for change requirement, open the **Audit Configuration** dialog box, deselect the checkbox, then click **Save**.

Configuration (Administrator)

Under the  tab, in the **Configuration** screen, the settings parameters are grouped by function into 5 subtabs:

- Network Settings
- Email Settings
- Instruments
- Lab Information
- Software Updates

Torrent Server Network Settings (Audit Trail)

Mac Address: b0:83:fe:e8:9e:ee

Public IP: 12.27.71.34
 DHCP
 Static

IP Address:

Subnet:

Gateway:

Name Servers:

Proxy Servers:

Proxy Login:

Ethernet 0	Detected ✓
IP Address	Detected ✓
Default route	Detected ✓
security.ubuntu.com:80	Detected ✓
drm.appliedbiosystems.com:443	Detected ✓
updates.iontorrent.com:80	Detected ✓
us.archive.ubuntu.com:80	Detected ✓
rssh.iontorrent.com:22	Detected ✓

Network Settings

If a problem occurs with the Ion Torrent™ Server, administrator-level users can use network settings information to troubleshoot the cause. Information about the Ion Torrent™ Server is displayed in the **Configuration** screen on the **Network Settings** tab.

An audit trail of the Ion Torrent™ Server network settings is available to view, export, and print by clicking **Audit Trail** on the **Network Settings** tab.

Configure Email Settings

Administrator-level users configure the Ion Torrent™ Server to allow sending email notifications.


Before you begin, obtain SMTP server URL and port information from your IT department.

1. Click ► **Configuration**.
2. Select the **Email Settings** tab.
3. In the **Torrent Server Email Settings** screen, enter the appropriate settings.
Required fields are indicated with a red asterisk (*).
4. Click **Update**.
5. Click **Send Test Email** to send a test email to the email address provided in the user account.

Instrument configuration

Administrator-level users can view the instrument configuration for an instrument that is connected to the Ion Torrent™ Server, such as:

- **Instrument Type**
- **Instrument Identifier** (name)
- **Instrument State**
- **Last PQ Date**

Click  **Configuration** **► Instruments**, then from the **Select Instrument** list, select an instrument and view the instrument details.



Lab Information

IMPORTANT! Your network system administrator should review support personnel contact information periodically to ensure that it is current and accurate.

Administrator-level users can review information about support personnel, listing points of contact if a problem with the Ion Torrent™ Server or a connected Ion PGM™ Dx System instrument occurs.

Click  **Configuration** **► Lab Information** to view the lab information.


- **Lab Contact**—This is the person in your organization who should be notified during a support request of problems related to the instrument.
- **IT Contact**—This is the person in your organization who should be notified during a support request of problems related to the Ion Torrent™ Server hardware or the network environment.

Update the software


IMPORTANT! Before updating the software, ensure that all instruments that are connected to the server are idle and no analysis jobs are running on the server or are queued to run.

Administrator-level users can update the software. Use the **Configuration** screen to:

- See the currently installed software version for each software module and component application.
- Check for software updates.
- Install software updates from a USB drive using the following steps.

1. Click  ► **Configuration**.
2. Select the **Software Updates** tab.
3. Click the desired instrument software module link to expand the information.
4. After viewing the software version information, click the software module heading a second time to collapse the information.
5. To check for updates, click **Check for updates**.
If an available update is found, **Download & Update** appears next to **Check for updates**. If no updates are found, “No updates available” displays.
6. To load an update from a USB drive, connect the USB drive, then click **Check for updates**.
The software searches the USB drive for updates and lists them.
7. Click **Update Server**.
8. In the confirmation dialog box, click **OK** to start the update.
After successful completion of the software update, the Ion Torrent™ Server automatically starts rebooting within 2 minutes.
9. After the server has rebooted (~10 minutes), click the browser **Refresh** button to return to the software home page.
After the update is complete, confirm the software version number in the **Software Updates** tab in the **Configuration** screen.


Data Management (administrator)

Administrator-level users can perform the following functions in the **Data Management** screen under the  tab.

- Monitor Ion Torrent™ Server disk space usage
- Manage archive settings, and view an audit trail of any changes made

Note: 72 hours after completion of data analysis, raw data (DAT) files are deleted from the server to conserve Ion Torrent™ Server disk space.

About the data output directory

The data output directory is the primary location on the Ion Torrent™ Server where data generated by the Ion PGM™ Dx System is stored. The path to the directory is listed in the **Data Management** screen under the  tab.

Note: The directory location is locked and cannot be changed. Users should configure their LIMS to access this folder location to receive sequencing output files.

After a sequencing run and data analysis are complete, the software creates a separate results folder for each sample in the data output directory using the following naming convention:


Dx_ <Library Name>_<Assay Name>_<PlannedRunShortCode>_<PlannedRunShortCode>

The following files are added to the results folder:

- Run log files: basecaller.log, sigproc.log, analysis.log
- PlannedRun-AuditTrail.pdf: audit trail of the planned run in PDF format
- Assay-specific variant files in VCF and XLS formats
- Dx_allele_counts.xls for each barcode: allele coverage for bases in hot spot regions
- <barcode>_rawlib.basecaller.bam: unmapped BAM file
- <LibPrepID>_rawlib.bam: mapped BAM file
- <LibPrepID>_rawlib.bam.bai: mapped BAM index file
- A checksum file containing checksums for each output file
- analysis.bfmask.stats: contains analysis statistics of wells in the bead find stage
- BaseCaller.json: a JSON format file of the statistics of basecaller, bead summary, filtering, phasing, and training subset
- datasets_pipeline.json: a JSON format file of the settings needed by the pipeline to run the basecaller module
- explog_final.txt: final run settings needed for analysis
- ion_params_00.json: a JSON format file of the detailed settings of the run and analysis arguments
- TFStats.json: a JSON format file of control statistics
- wells_beadogram.png: a figure of statistics to characterize wells
- Test_Report.pdf: an assay-specific PDF report of targeted clinical and analytical variants; can be generated in multiple languages if selected (filenames are appended with language suffix)
- Lab_Report.pdf: similar to the Test Report, but includes all analytical variants detected; can be generated in multiple languages if selected (filenames are appended with language suffix)
- Summary.tab: non-targeted test results summary
- Snvindel.tab: non-targeted SNV/INDEL detail table
- Fusion.tab: non-targeted fusion detail table
- Cnv.tab: non-targeted CNV detail table
- Target_summary.tab: targeted test results summary
- Target_hotspot.tab: targeted SNV/INDEL detail table
- Target_fusion.tab: targeted fusion detail table
- Target_cnv.tab: targeted CNV detail table
- <LibPrepID>_<AnalysisID>.final.vcf: VCF file
- Info.csv: contains sample attributes, reagent information, QC values, and instrument information
- <barcode>_rawlib.basecaller.bam: unmapped BAM file – DNA
- <barcode>_rawlib.basecaller.fastq: FASTQ file – DNA
- <LibPrepID>_rawlib.bam: mapped BAM file – DNA
- <LibPrepID>_rawlib.bam.bai: mapped BAM index file – DNA
- <barcode>_rawlib.basecaller.bam: unmapped BAM file – RNA
- <barcode>_rawlib.basecaller.fastq: FASTQ file – RNA

- <barcode>_rawlib.basecaller.alignments.bam: mapped BAM file – RNA
- <barcode>_rawlib.basecaller.alignments.bai: mapped BAM index file – RNA
- Bead_density_contour.png: loading density figure
- iontrace_Library.png: key incorporation traces figure
- raw_peak_signal: key signal for controls and library
- readLenHisto.png: histogram of the read length
- <LibPrepID>_rawlib.stats.cov.txt: amplicon statistics
- basecaller.log: base caller log file
- sigproc.log: signal processing log file
- analysis.log: analysis log file
- PlannedRun-AuditTrail.pdf: Planned Run audit file

Disk space usage and data archiving

Ion Torrent™ Server disk space usage is shown in the **Data Management** screen under the  tab. To maintain sufficient disk space, we recommend implementing a systematic plan to archive older run data, results files, variants information, and signed reports to an external file storage system. Consult your Field Service Engineer to discuss archive and database backup options. Your local system administrator is responsible for establishing your data archive system.

After an archive system has been established, administrator-level users can configure the archive schedule in the **Data Management** screen. Archived results and reports can be restored to the Ion Torrent™ Server and downloaded from the **Completed Runs & Results** screen. See “Restore archived results” on page 53.

Note: If the Ion Torrent™ Server has ≤ 1 terabyte (TB) of free disk space, an alert notifies the user that there is insufficient disk space when setting up the run on the sequencer. The run cannot proceed until data on the server is archived and deleted. Contact your local system administrator to manually archive and delete data. You should also change the archive schedule.

IMPORTANT!

- If results are restored from runs that were archived in an older version of Torrent Suite™ Dx Software (version 5.6.4 and earlier), the software may not display all the variants from the restored results in the user interface, and reports generated from those restored results may not contain all variants. Carefully review the restored data in the user interface to determine whether all variants are present. Do not generate new reports from results restored from these older archives. Note that all variants are preserved in the restored source files, and can be downloaded using the **Download Files** command. This issue was corrected in version 5.8 of the software.
 - In older versions of Torrent Suite™ Dx Software (version 5.6.4 and earlier), results reports should be generated and signed before results are archived. See “Sign the run results” on page 67.
-

Update archive settings

Administrator-level users can view and edit archive settings in the **Data Management** screen. Consult your Field Service Engineer to set up a database archive and backup system.

1. Click  **Data Management**.
2. Edit the archive settings.

Setting	Procedure
Auto archive after	Select the number of months for data to remain on the Ion Torrent™ Server before it is archived on an external server. To free up additional space on the Ion Torrent™ Server, reduce this interval.
Archive Directory	Enter the file path to the external archive directory. The default path must be changed before data can be archived.

Note: The **Data Output Directory** is listed in this screen but cannot be changed.

3. Click **Save**.

Archive notifications

Administrator-level users receive email notifications about archives when any of the following conditions occur:


- The archive location has ≤120 gigabytes (GB) of free disk space remaining.
- The archive location has not been changed from the default location (`/results/analysis`). The archive path must be changed to that of the external server before data can be archived.
- The archive location is not accessible when the system tries to archive a run.

To receive email notifications about archives, a valid administrator email address must be entered into Torrent Suite™ Dx Software. For more information, see “User Management (administrator)” on page 88.

Additionally, the Ion Torrent™ Server must be configured to allow sending email notifications. For more information, see “Configure Email Settings” on page 82.


Logs (administrator)

Select a log category

Within the  **Logs** screen, administrator-level users choose from the **Select Category** list to view and set auto-deletion rules for **Torrent Suite** and **Instrument** logs.

Manage logs

Administrator-level users can manage logs to set retention times and set up auto-deletion of logs.

1. Click  **Logs**.
2. Use the **Select Category** list to view and set auto-deletion rules for **Torrent Suite** or **Instrument** logs.
3. Click **Manage Log**.
4. In the **Manage Log File** dialog box, from the **Retention Period** list, select the number of months that logs are to be retained on the server.
5. Select **Enable Auto Deletion** to automatically delete log files after the designated retention period.
6. Click **Save**.



User Management (administrator)

Administrator-level users can use the  tab, in the **User Management** screen, to perform the following actions:

- Add users
- Assign user privileges (roles)
- Edit user information
- View user account audit trails
- Manage user account policies

Add a new user

Note: The email settings on the Ion Torrent™ Server must be configured before new users can receive emails. See “Configure Email Settings” on page 82 for more information.

1. To add a new user account, under the  tab, in the **User Management** screen, click  **Add New**.
2. In the **Create New User** dialog box, enter the account information.


Note: Required fields are indicated by a red asterisk (*).

3. Select the user-access level (Administrator, Manager, or Operator) from the **Role** list.
4. To enable the ability to sign reports (Manager- or Administrator-level users only), select the **Electronic Signature** checkbox.
5. Click **Save**.

The Ion Torrent™ Server sends a temporary password to the email address of the new user.


Set security policies

Administrator-level users set user account security policies. User account security policies include the permissible number of failed sign-in attempts, password lifetime, and the length of inactivity before a user is automatically signed out of the software.

1. Click  **▶ User Management**.
2. In the **User Management** screen, click **Policies**.
3. Update the **Policies** dialog box. Accept the default values or select the appropriate values to set the user account suspension, password, and session policies, then click **Save**.
4. Select **Enabled** to enable the session policy settings.
5. Click **Save**.


Sort users

Administrator-level users can sort the list of users to help find a user account to work with.

1. Click  **▶ User Management**.
2. Click the header name of interest to sort alphabetically.
3. Click the header name a second time to reverse the order of users displayed.
4. Click **User Management** to return to the default display.

Edit user accounts

Administrator-level users can edit user account information.

1. Click  **▶ User Management**.
2. In the **User Name** column, click **Edit** under a user name.
3. In the **Edit User Details** dialog box, enter the desired changes to the account information.
 - From the **Role** list, select the user-access level (**Administrator**, **Manager**, or **Operator**).
 - To allow manager- or administrator-level users to sign reports, select the **Electronic Signature** checkbox.
 - In the **Status** list, select the appropriate status (**Active**, **Suspend**, or **Disable**).
4. Click **Save** to make the changes.

Reset password



Administrator-level users can reset user account passwords.

Note: Operator level users can reset their passwords by following the directions included in the system-generated email notification of a pending password expiration.

1. In the Edit User Details window, click **Reset Password**.
2. In the **Reset User Password** dialog box, click **Send password in email**, to email a new password to the email address entered in the user account.
3. Alternatively, enter a new password into the **Password** field.
4. Reenter the new password into the **Re-type Password** field, then click **Save**.
The confirmation message "Password reset successfully" appears. Click **Back**, then **Save** to return to the **User Management** window.

View user audit trail

The user audit trail is a record of when the user account was created and modified.

1. Under the  tab, in the **User Management** screen, in the **Actions** column, click **Audit**.
2. In the **Audit Trail** dialog box, click  (**Details**) under **Record**.
3. In the **Audit Record Details** dialog box, click **Export** to export a PDF of the record.



Anomalies

Software anomalies

For Torrent Suite™ Dx Software anomalies, refer to the release notes for your version of the software, included on the software USB drive.



Troubleshooting

HotSpot ID corrections

The following HotSpot IDs reported in the software are inaccurate, and correspond to the actual COSMIC IDs listed below. These anomalies do not impact test results.

HotSpot ID in the software	Actual COSMIC ID	Gene	Amino acid change	Nucleotide change
OM3157	COSM1235478	MAP2K1	p.Lys57Asn (p.K57N)	c.171G>T
COSM1235478	N/A ^[1]	MAP2K1	p.Lys57Met (p.K57M)	c.170A>T
COSM1562837	COSM5077832	MAP2K1	p.Phe53Val (p.F53V)	c.157T>G

^[1] This variant is not in the COSMIC database.

Torrent Suite™ Dx Software

Observation	Possible cause	Recommended action
Cannot sign in to the Ion Torrent™ Server	You have either forgotten your password or are signed out due to several failed login attempts.	Contact a Torrent Suite™ Dx Software administrator-level user.
Cannot sign in to the Ion PGM™ Dx Sequencer	The Ion PGM™ Dx Sequencer lost its connection to the Ion Torrent™ Server.	Contact a Torrent Suite™ Dx Software administrator-level user.
Data and user profiles are missing from the software	You are signed into the incorrect mode of the software. Data and user profiles created in Torrent Suite™ Dx Software will not be visible in Torrent Suite™ Assay Development Software, and vice versa.	Confirm that you are signed into the correct mode of the software.
Batch sample import fails	One or more entries in the sample-import spreadsheet contains special characters, lines breaks, unexpected spaces, incorrect entry length, incorrect date formatting, or other formatting errors.	Check each entry for correct formatting, correct any errors, and repeat the import.

Observation	Possible cause	Recommended action
Batch sample import fails (continued)	Blank rows were copied into the sample-import template file from a different source.	Rows that appear empty may contain hidden formatting that conflicts with the import function. Start with a clean sample-import template file, and be careful to copy only those rows that contain actual data.
	The sample import spreadsheet contains a nonunique Sample ID.	Every Sample ID in the software must be unique. Make sure the spreadsheet does not contain any duplicate IDs, and repeat the import. Note that the system check is not case-sensitive, so a Sample ID of ABC1 conflicts with abc1.
	The headings in the sample import spreadsheet do not match the sample attributes in the software.	The headings must match the sample attributes in the software exactly. Check the headings for spelling or other errors.
Library batch import fails	One or more entries in the library batch import spreadsheet contains special characters, lines breaks, unexpected spaces, incorrect entry length, incorrect date formatting, or other formatting errors.	Check each entry for correct formatting, correct any errors, and repeat the import.
	Blank rows were copied into the library batch import template file from a different source.	Rows that appear empty can contain hidden formatting that conflicts with the import function. Start with a clean library batch import template file, and be careful to copy only those rows that contain actual data.
	The library batch import spreadsheet contains a nonunique Library Batch ID.	Every Library Batch ID in the software must be unique. Ensure that the spreadsheet does not contain any duplicate IDs, and repeat the import. Note that the system check is not case-sensitive, so a Library Batch ID of ABC1 conflicts with abc1.
	A Sample ID entered in the library batch import spreadsheet does not match a Sample ID listed in the Manage Samples screen.	Ensure that the Sample IDs entered into the spreadsheet are correct and match an existing sample ID added to the software.
	The Barcode ID name format does not exactly match the format that is used in the Prepare Library Batch dialog box.	Use the following Barcode ID name format: lonDx-1 through lonDx-16.
	An invalid library, control, or panel kit barcode has been entered in the spreadsheet.	Ensure that you have correctly entered a valid kit barcode in the appropriate cell of the spreadsheet.

Observation	Possible cause	Recommended action
<p>Cannot execute my Planned Run</p> <p>Details: "System error" message is displayed</p>	<p>The Tube Label you assigned the Planned Run is not unique.</p> <p>Note: The Ion PGM™ Dx Instrument Control software does not allow use of the same Tube Label text within 7 days.</p>	<p>Assign the Planned Run a new, unique Tube Label.</p>
	<p>Incorrect template prep kit barcode was entered.</p>	<p>Rescan the correct template prep kit barcode.</p>
<p>The results of my run are not showing up in the Data tab</p>	<p>Ion Torrent™ Server disk space is full.</p>	<p>Clear disk space on the Ion Torrent™ Server. After sufficient disk space is cleared, data transfer from the Ion PGM™ Dx Sequencer to the Ion Torrent™ Server recommences and data analysis proceeds.</p>
	<p>Ion PGM™ Dx Sequencer lost connection to the Ion Torrent™ Server.</p>	<p>Reestablish the Ion PGM™ Dx Sequencer Ethernet connection to the Ion Torrent™ Server. Once the connection has been restored, data transfer from the Ion PGM™ Dx Sequencer to the Ion Torrent™ Server recommences and data analysis proceed.</p>
<p>Cannot find file links to the BAM and VCF file</p>	<p>The run failed. Links to BAM and VCF files are not available for runs that fail QC.</p>	<p>Repeat the run.</p>
<p>Loading metrics are reported as 0</p>	<ul style="list-style-type: none"> • Chip failure • Defective pariposer • Chip leak 	<p>Refer to sequencing troubleshooting for possible causes of chip or pariposer failure.</p>
	<p>Sample preparation failure resulting in loss of sample</p>	<p>Refer to troubleshooting for sample, library, and template preparation.</p>
<p>QC Status is Failed due to one or more QC metrics listed as N/A</p> <p>Details: Under the Monitor tab in the Run View screen, under View QC Details, one or more of the QC metrics (Library RNA, Library DNA, CF-1, DNA Control, RNA Control, DNA NTC, and RNA NTC) have a value of N/A and the QC Status is Failed.</p>	<ul style="list-style-type: none"> • The control associated with the QC metric was not added to the reaction or otherwise failed to generate a measurable value. • The QC status was not updated properly in the database due to a software issue. This is uncommon. 	<p>For Administrator-level users, the Repopulate QC button is displayed under the Monitor tab in the Run View screen after a run.</p> <ol style="list-style-type: none"> 1. Click Repopulate QC to update all QC metrics in the database. <p>Note: Repopulation takes 15–30 minutes.</p> <ol style="list-style-type: none"> 2. If any metric is still listed as N/A and the QC Status is still Failed, repeat the run starting at template preparation from the DNA/RNA combined library, and ensure that you add all controls. If the problem continues, contact Technical Support.

Observation	Possible cause	Recommended action
<p>Run analysis does not complete</p> <p>Details: If the Analysis Status for a run is listed in the Monitor tab as "RUNNING" for more than 12 hours, a Restart Analysis button appears in the tab (Administrator-level users only).</p>	<p>Run analysis has stalled in the analysis pipeline.</p>	<p>Click Restart Analysis to restart the analysis from the beginning.</p>
<p>Run analysis does not complete, and Restart Analysis is not successful</p>	<p>A known software issue exists where the database status of a run is not updated properly. This behavior is uncommon.</p>	<p>If clicking Restart Analysis in the Monitor tab does not result in a completed analysis, repeat the run starting at template preparation from the DNA/RNA combined library. If the problem continues, contact Technical Support.</p>

Library preparation

Observation	Possible cause	Recommended action
Display message: Invalid kit barcode or Expired kit	Kit is beyond expiration date.	Use a nonexpired kit.
	Incorrect kit scanned.	Rescan the barcode of the correct kit.
Run QC passed but low coverage uniformity (<95%)	Poor purification causes loss of short amplicons	Vortex AMPure™ XP reagent thoroughly before use, and be sure to dispense accurate volume.
Run QC passed, low percent full length On Target Reads, but Control passed	Library prep failure	Reprepare the sample library.
Samples passed but CF-1 Mean AQ20 Read Length failed	Reads filtered out due to high polyclonal ISPs caused by too much library added to the Ion OneTouch™ Dx amplification reaction	Repeat Ion OneTouch™ Dx run using less library in the amplification reaction.
Run QC passed, but one or more sample libraries failed Percent Reads QC metric	Library prep failed due to unwashed beads.	Be sure to wash the library beads prior to use.
	Library preparation failed due to wrong library amplification primers.	Use the LIB Primers provided in the Ion PGM™ Dx Library Kit.
	Library prep failed due to residual salt after wash.	Carefully remove all wash solution prior to elution.
	Library prep failed due to mis-quantification of input DNA.	Requantify input DNA.
	Library prep failed due to inefficient PCR, digestion, or ligation.	Ensure that you properly dispense and mix the viscous components at each step.
		Ensure that you use the correct thermal cycling conditions.
	Library prep failed due to the library being discarded during purification of the amplified library.	Be sure to save the supernatant during first-round purification and save the library pellet during the second round purification of the amplified library.
	Library prep failed due to over-drying of the AMPure™ XP beads.	Do not dry the AMPure™ XP beads more than 5 minutes.
		Ensure that you dispense exactly 10 µL of capture reagent to the amplified library.
Library failed due to ineffective capture of the amplified library.	Ensure that the library capture reagent is at room temperature before use.	
	Ensure that you dispense exactly 10 µL of capture reagent to the amplified library.	
	Make sure to mix completely and incubate for 5 minutes at room temperature.	

Observation	Possible cause	Recommended action
Run QC passed, but one or more library samples failed Percent Reads QC metric - continued	Reads filtered out due to primer-dimers.	In unamplified library purification, ensure that you use the correct amount of AMPure™ XP reagent. Do not combine Barcode Adapters, LIB DNA Ligase, and LIB Switch Soln prior to addition. Ensure that Barcode Adapters are diluted properly.
	Reads filtered out due to low quality ISPs.	Repeat the Ion OneTouch™ Dx run. If it still happens, then reprepare the library.
	Reads filtered out due to high polyclonal ISPs caused by failed consumables.	Repeat the Ion OneTouch™ Dx run. If it still happens, then reprepare the library.
	Reads filtered out due to high polyclonal ISPs caused by too much library added to the Ion OneTouch™ Dx amplification reaction.	Requantify input DNA or RNA, and remake the library.
	Forgot to add the library to amplification reaction.	Repeat the Ion OneTouch™ Dx run. If observation recurs, then reprepare the library.

Template preparation

Observation	Possible cause	Recommended action
Display message: Failed to set up system time at startup. Check your connection to the Ion Torrent™ Server.	Ion OneTouch™ Dx Instrument and Ion Torrent™ Server connection is not established	Check your network connection to the Ion Torrent™ Server to make sure the connection is established, then reboot the Ion OneTouch™ Dx Instrument.
	Instrument is still in the process of establishing a connection	Allow 10 minutes to see if the display message clears.
Display message: Failed to connect to the Ion Torrent™ Server. Check your connection.	Ion OneTouch™ Dx Instrument and Ion Torrent™ Server connection is not established during startup	Check your network connection to the Ion Torrent™ Server to make sure the connection is established, then reboot the Ion OneTouch™ Dx Instrument.
Display message: Pressure too high. Reboot the instrument to clear the alarm.	<ul style="list-style-type: none"> • Hardware issue • Clogged TMPL Emulsion Cartridge due to contaminated reagents or defective emulsion cartridge 	Reboot the Ion OneTouch™ Dx Instrument to clear the alarm. Use a new TMPL Emulsion Cartridge and fresh reagents to repeat the run. Contact Technical Support (see Appendix E, “Customer and technical support”) if the issue persists.

Observation	Possible cause	Recommended action
Display message: TEC current too high. Reboot the instrument to clear the alarm.	Hardware issue	Contact Technical Support (see Appendix E, "Customer and technical support"). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm. IMPORTANT! A sample created during a run with this alarm raised must NOT be used.
Display message: Coolant pump does not flow.	Hardware issue	Contact Technical Support (see Appendix E, "Customer and technical support"). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm.
Display message: Sensor unable to measure instrument temperature	Hardware issue	Contact Technical Support (see Appendix E, "Customer and technical support"). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm.
Display message: Software versions incompatible. Go to the Options menu and update the software.	The system software was updated, but the instrument software was not.	After the system software has been updated, update the instrument software as follows: <ol style="list-style-type: none"> 1. On the main menu of the instrument, press Options and follow the instructions to check for and install updates. 2. When installation is complete, follow the onscreen prompts to restart the instrument. IMPORTANT! You must restart the instrument before proceeding.
Display message: Sensor unable to measure pressure.	Hardware issue	Contact Technical Support (see Appendix E, "Customer and technical support"). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm.
Display message: Connection failure with Ion Torrent™ Server	Ion OneTouch™ Dx Instrument and Ion Torrent™ Server connection is not established	Check that a network connection to the Ion Torrent™ Server is established, then reboot the Ion OneTouch™ Dx Instrument. Note: A sample created during a run with this alarm raised can still be used.
Display message: Motor current too high. Reboot the instrument to clear the alarm.	Hardware issue	Contact Technical Support (see Appendix E, "Customer and technical support"). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm. Note: A sample created during a run with this alarm raised can still be used.

Observation	Possible cause	Recommended action
Display message: Set temperature out of range. Reboot the instrument to clear the alarm.	Hardware issue	Contact Technical Support (see Appendix E, “Customer and technical support”). Reboot the Ion OneTouch™ Dx Instrument to clear the alarm. IMPORTANT! If this alarm is raised, you cannot use the prepared template.
TMPL CF-1 fails QC metric(s); all sample libraries pass QC metrics	TMPL CF-1 reagent not added	Repeat Ion OneTouch™ Dx Instrument run and Ion PGM™ Dx Sequencer run, and ensure that the TMPL CF-1 reagent is added per procedure description.
	The minimum Key Signal value of the CF-1 control required for the assay is >0.	The minimum Key Signal value for the CF-1 control (i.e., the Templating Control) is set to 0 in all preinstalled assays and new custom assays in Torrent Suite™ Dx Software 5.12.5. Custom assays created in previous versions of the software may have a higher Key Signal value for the control.

Sequencing

Observation	Possible cause	Recommended action
Cannot import my sample	Sample ID is greater than 20 characters.	Limit Sample ID to 20 characters or fewer.
	Expired library kit.	Use a nonexpired library kit.
	Incorrect library kit barcode.	Rescan the correct library kit barcode.
Run report says run failed	Run failed because one or more QC metrics were not met	
I created a Planned Run but the Ion PGM™ Dx Sequencer does not recognize the Short Code	Planned Run was not executed.	In Torrent Suite™ Dx Software: 1. Select the Planned Run, then click Execute . 2. Complete the dialog box, then click Save .
	The Ion PGM™ Dx Sequencer and Ion Torrent™ Server have lost connection.	Check the network connection to the Ion Torrent™ Server to ensure that the connection is established, then reboot the instrument.
TMPL CF-1 fails QC metric(s); all sample libraries fail QC metrics	<ul style="list-style-type: none"> • Forgot Enzyme • Forgot Seq primer • Used wrong annealing program 	Repeat Ion OneTouch™ Dx Instrument run and Ion PGM™ Dx Sequencer run.
Display message: Lost connection to the Ion Torrent™ Server	The connection between the instrument and the server has been lost.	Check the network connection to the Ion Torrent™ Server, and then reboot the Ion PGM™ Dx Sequencer. If this alarm appears during a run, the data created during that run can still be used.

Observation	Possible cause	Recommended action
Display message: Failed to set up system time at startup. Check your connection to the Ion Torrent™ Server.	The connection between the Ion PGM™ Dx Sequencer and the Ion Torrent™ Server has been lost.	<ol style="list-style-type: none"> 1. Check the network connection to the Ion Torrent™ Server to make sure the connection is established, then reboot the instrument. 2. If the problem persists, replace the network cable(s) to the instrument and server. 3. If the problem persists, contact Technical Support (see Appendix E, “Customer and technical support”).
	Instrument is still in the process of establishing a connection	Allow 10 minutes to see if display message clears.
Display message: Bad boot drive detected	Hardware issue	Contact Technical Support (see Appendix E, “Customer and technical support”). If this alarm appears during a run and data for the run is generated, that data may still be used.
Display message: UBoots do not match	Hardware issue	Contact Technical Support (see Appendix E, “Customer and technical support”).
Display message: Kernels do not match	Hardware and/or software issue	Contact Technical Support (see Appendix E, “Customer and technical support”).
Display message: Bad results data drive	<ul style="list-style-type: none"> • On some machines, the warning appears before the reboot completes. • There is a hardware issue. 	Wait for a few minutes to see if the error message disappears. If the error message disappears, data obtained during a run with this alarm raised can still be used. If the problem persists, contact Technical Support (see Appendix E, “Customer and technical support”).
Display message: Lost chip connection	The instrument cannot detect a chip in the chip clamp.	Refer to the <i>Oncomine™ Dx Target Test Part IV: Sequencing and Results Reports User Guide</i> troubleshooting section.
Display message: Lost communication with valve board	Hardware issue	Contact Technical Support (see Appendix E, “Customer and technical support”).
Run aborted during presequencing step	Critical alarm present on the Ion PGM™ Dx Sequencer	Reboot the Ion PGM™ Dx Sequencer to clear the alarm, then restart the sequencing run.
Display message: Sensor unable to measure instrument temperature	Hardware issue	Contact Technical Support (see Appendix E, “Customer and technical support”).

Observation	Possible cause	Recommended action
<p>Display message: Instrument temperature too high</p>	<ul style="list-style-type: none"> • Room temperature is too high. • Clogged filter or blocked airway on the instrument • Hardware issue (fan is not running or running too slowly) 	<p>Note: The data created during a run with this alarm raised may still be used if all the QC metrics are met.</p> <ol style="list-style-type: none"> 1. If the ambient room temperature is above 30°C, lower it. 2. Make sure that the round filter on the back panel of the instrument has unrestricted airflow. If the filter is clogged with dust, clean it as follows: <ol style="list-style-type: none"> a. Pinch the dirty filter with your fingers, then remove it from the instrument. <div data-bbox="1029 701 1414 1031" data-label="Image"> <p>The illustration shows a hand with blue gloves pinching a dark, circular filter. The filter is being held between the thumb and index finger, demonstrating the correct way to remove it from the instrument.</p> </div> <ol style="list-style-type: none"> b. Shake the filter over a waste container to remove most of the dust. c. Rinse the filter with running water to remove any remaining dust. The water flow should be from the inside-facing surface to the outside-facing surface through the filter. d. Air dry the filter. e. Blot any remaining dust from the filter using tape. f. Reinsert the filter. <p>If the problem persists, contact Technical Support (see Appendix E, “Customer and technical support”).</p>
<p>Display message: Instrument temperature too low</p>	<ul style="list-style-type: none"> • Room temperature is below 20°C. • Hardware issue 	<p>Note: The data created during a run with this alarm raised may still be used if all the QC metrics are met.</p> <p>If the ambient room temperature is below 20°C, raise it. If the problem persists, contact Technical Support (see Appendix E, “Customer and technical support”).</p>

Observation	Possible cause	Recommended action
Display message: Chip temperature too high	<ul style="list-style-type: none"> Room temperature is too high. Clogged filter or blocked airway on the instrument Hardware issue (instrument fan is not running or running too slowly) 	<p>IMPORTANT! The data created during a run with this alarm raised should <i>not</i> be used.</p> <p>See the recommended action for “Display message: Instrument temperature too high” on page 101.</p>
Display message: Chip temperature too low	Hardware issue	<p>IMPORTANT! The data created during a run with this alarm raised should <i>not</i> be used.</p> <p>Contact Technical Support (see Appendix E, “Customer and technical support”).</p>
Display message: Instrument idle temperature too high	<ul style="list-style-type: none"> Room temperature is too high. Clogged filter or blocked airway on the instrument Hardware issue (fan is not running or running too slowly) 	<p>Note: The data created during a run with this alarm raised may still be used if all the QC metrics are met.</p> <p>See the recommended action for “Display message: Instrument temperature too high” on page 101.</p>
Display message: Instrument idle temperature too low	<ul style="list-style-type: none"> Ambient room temperature is below 20°C. Hardware issue 	Bring the ambient temperature up to 20°C. If the problem persists, contact Technical Support (see Appendix E, “Customer and technical support”).
Display message: Fan current too low	Hardware issue	Contact Technical Support (see Appendix E, “Customer and technical support”).
Display message: Heater current too low	Hardware issue	<p>IMPORTANT! If the chip temperature is also out of range, data created during a run should <i>not</i> be used.</p> <p>Contact Technical Support (see Appendix E, “Customer and technical support”). If no chip temperature alarms are raised, data created during a run may still be used if all the QC metrics are met.</p>
Display message: Pressure too high	Internal pressure regulator was not set correctly	Contact Technical Support (see Appendix E, “Customer and technical support”).

Observation	Possible cause	Recommended action
Display message: Pressure too low.	<ul style="list-style-type: none"> • Gas line is not connected to the instrument • Gas cylinder may be turned off or empty 	<ol style="list-style-type: none"> 1. Verify that the gas line is connected to the instrument. 2. Verify that the gas cylinder has at least 500 psi on the pressure gauge inside the tank and 30 psi on the gauge that regulates pressure to the outlet. (Both pressure gauges are mounted on the gas cylinder.) 3. Confirm that the outlet valve on the regulator is turned on. 4. If the problem persists, contact Technical Support (see Appendix E, “Customer and technical support”).
Display message: Sensor unable to measure gas pressure. Check supply gas pressure.	<ul style="list-style-type: none"> • Gas line is not connected to the instrument • Gas cylinder may be turned off or empty • Hardware issue 	<ol style="list-style-type: none"> 1. Verify that the gas line is connected to the instrument. 2. Verify that the gas cylinder has at least 500 psi on the pressure gauge inside the tank and 30 psi on the gauge that regulates pressure to the outlet. (Both pressure gauges are mounted on the gas cylinder.) 3. Confirm that the outlet valve on the regulator is turned on. 4. If the problem persists, contact Technical Support (see Appendix E, “Customer and technical support”).
Display message: Failed to set the pressure to target range. Check the gas connection and try again.	<ul style="list-style-type: none"> • Gas line is not connected to the instrument • Gas cylinder may be turned off or empty • Hardware issue (regulator malfunction) 	<ol style="list-style-type: none"> 1. Verify that the gas line is connected to the instrument. 2. Verify that the gas cylinder has at least 500 psi on the pressure gauge inside the tank and 30 psi on the gauge that regulates pressure to the outlet. (Both pressure gauges are mounted on the gas cylinder.) 3. Confirm that the outlet valve on the regulator is turned on. 4. If the problem persists, contact Technical Support (see Appendix E, “Customer and technical support”).
Display message: Failed to locate the barcode scanner. Check if the scanner is attached.	The connection between the barcode scanner and the Ion PGM™ Dx Sequencer has been lost.	Make sure the scanner is plugged into a USB port on the instrument. If it is connected and the alarm still appears, try plugging the scanner into a second USB port. If the alarm persists, contact Technical Support (see Appendix E, “Customer and technical support”).

Observation	Possible cause	Recommended action
Display message: Results drive not accessible. Reboot and try again.	<ul style="list-style-type: none"> On some machines, the warning appears before the reboot completes Hardware issue 	Wait for a few minutes to see if the error message disappears. If the error message appears and disappears during a run, data obtained during that run can still be used. If the alarm persists, contact Technical Support (see Appendix E, "Customer and technical support").
Display message: Results drive check failed	Hardware issue	If the error message disappears when you return to the main instrument screen, this alarm can be ignored. Otherwise, contact Technical Support (see Appendix E, "Customer and technical support").
Display message: Unable to mount the file system	Hardware and/or software issue	<ol style="list-style-type: none"> Reboot the instrument to clear the alarm. If the alarm is not cleared after reboot, contact Technical Support (see Appendix E, "Customer and technical support").
Display message: Failed to set up FTP connection. Check your connection to the Ion Torrent™ Server .	The network connection is not established or an incorrect IP address was used.	Confirm that the server information is correct for the Ion Torrent™ Server. Contact your local network administrator for support if the issue persists.

No Calls

Observation	Possible cause	Recommended action
MINCOV<347, PosCov<2, NegCov<2*	No Call = Coverage ^[1]	Repeat the run with fewer samples per chip may improve coverage.
NODATA	No Call = No data ^[1]	Repeating the run with fewer samples per chip may improve coverage.
QualityScore<8	No Call = Quality score ^[1]	Remake the DNA and/or RNA libraries starting from nucleic acid sample to improve the quality score if the quality score is due to low coverage.
STDBIAS0.99034>0.96, STDBIASPVAL0.299<=1	No Call = Strand bias ^[1]	Remake the DNA and/or RNA libraries starting from nucleic acid sample to improve strand coverage.

^[1] The reason (Observation) for a No Call can be determined by examining the value of the FR tag for a given variant, listed in the output VCF file.

Observation	Possible cause	Recommended action
<p>PREDICTIONSHIFTx0.435739</p> <p>Note: In rare cases, one specific variant may interfere with EGFR S768I (COSM6241) causing a No Call to be assigned. Internal validation studies indicate that no positive calls are missed due to such a No Call.</p>	<p>No Call = Signal</p>	<p>Repeat not needed.</p>
<p>REJECTION</p>	<p>Noise from nearby positive variants</p>	<p>Repeat not needed.</p>



Performance characteristics

For performance characteristics of the Oncomine™ Dx Target Test Kit, see the *Oncomine™ Dx Target Test Part I: Test Description and Performance Characteristics User Guide*.

For performance characteristics of the Ion PGM™ Dx System, see the *Ion PGM™ Dx System Performance Characteristics User Guide* (Pub. No. MAN0018763).



Instrument warranty

For new Ion Torrent™ instruments, Life Technologies warrants to and only to buyer for twelve (12) months from the date of shipping, that the Ion Torrent™ software and Ion Torrent™ instruments are free from defects in material and workmanship and conform to Life Technologies' published specifications in all material respects. Where a valid and timely claim in respect of breach of Ion Torrent™ warranty is submitted to Life Technologies, Life Technologies may, at its discretion, replace, repair or modify the Ion Torrent™ instrument. Any agreed replacement shall be at 1:1, like-kind basis, at no cost to the buyer. For Ion Torrent™ chips or reagents reasonably determined by Life Technologies to be defective, independent of user error, shall be replaced by Life Technologies on a 1:1, like-kind basis at no cost to buyer, provided that such defective Ion Torrent™ chips or reagents were used by buyer prior to their expiration date, or if there is no expiration date, the Ion Torrent™ chips or reagents were used within six (6) months of receipt, and the defect was promptly reported with appropriate detail to Life Technologies' technical support.

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Customer and technical support

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- Product support
- Order and web support
- Safety Data Sheets (SDSs; also known as MSDSs)

Additional product documentation, including user guides and Certificates of Analysis, are available by contacting Customer Support.

Obtaining Certificates of Analysis

The Certificate of Analysis provides detailed quality control and product qualification information for each product. Certificates of Analysis are printed and shipped with the product.

Obtaining Certificates of Conformance

The Certificate of Conformance provides information on conformance testing of each instrument provided with the system. Certificates of Conformance are shipped with the instrument, and are also available by contacting Customer Support at [thermofisher.com/support](https://www.thermofisher.com/support).



**Oncomine
Dx Target
Test**

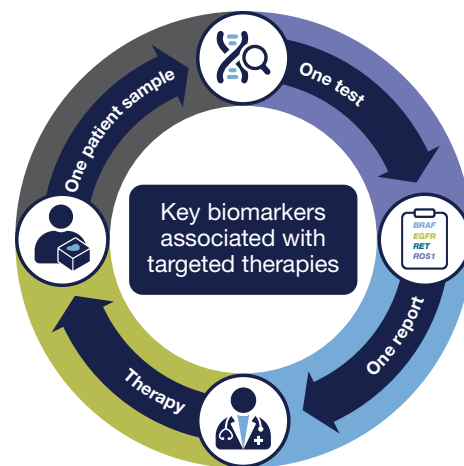
A new paradigm in testing for targeted therapies in NSCLC and CC with an FDA-approved NGS CDx test

The Ion Torrent™ Oncomine™ Dx Target Test is the first targeted next-generation sequencing (NGS) *in vitro* diagnostic (IVD) test for non-small cell lung cancer (NSCLC) and cholangiocarcinoma (CC), simultaneously delivering multiple biomarker results for multiple targeted therapies from one sample within 4 days.

This test is now reimbursed by Medicare and the top 40 commercial payers, covering over 200 million US lives.

The Oncomine Dx Target Test enables:

- **Fast results**—The single streamlined sequencing workflow enables concurrent analysis of both DNA and RNA targets. From sample extraction to clinical test report, the total workflow turnaround time is 4 days.
- **Clinical performance**—Based on Ion AmpliSeq™ technology, the test is designed to deliver robust and reproducible results in 23 genes clinically associated with NSCLC and one gene in CC, all from 10 ng of DNA and RNA from formalin-fixed, paraffin-embedded (FFPE) tissue.
- **Automated clinical report**—The Oncomine Dx Target Test results are presented in a single two-part Clinical Test Report that incorporates companion diagnostic (CDx) biomarker results, with associated therapy indications, and other detected cancer-associated gene variant results.



Cancer type	Gene	Targeted therapies
NSCLC	<i>BRAF</i>	TAFINLAR® (dabrafenib) in combination with MEKINIST® (trametinib)
	<i>EGFR</i> L858R, exon 19 deletions	IRESSA® (gefitinib)
	<i>EGFR</i> exon 20 insertions	EXKIVITY™ (mobocertinib)
	<i>RET</i>	GAVRETO™ (pralsetinib)
	<i>ROS1</i>	XALKORI® (crizotinib)
CC	<i>IDH1</i>	TIBSOVO® (ivosidenib)

Figure 1. List of genes for therapeutic use.

Complete system: from sample to actionable result, powered by proven Ion Torrent and Ion AmpliSeq NGS technology

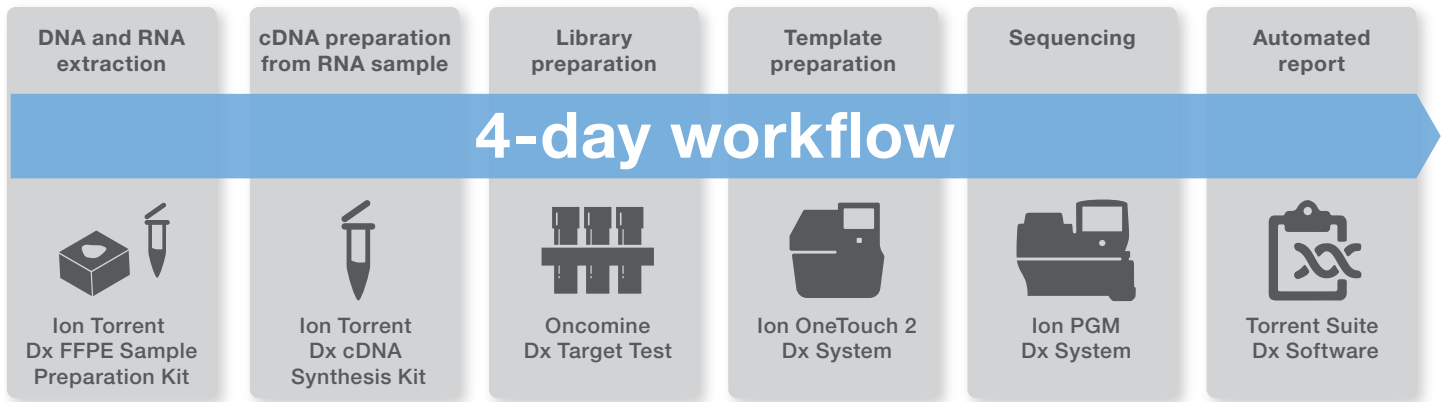


Figure 2. The Oncomine Dx Target Test utilizes a single streamlined NGS workflow for detecting cancer-associated biomarkers, incorporating reagents, instrument systems, and bioinformatics. The turnaround time, from FFPE sample to report, is 4 days.

Optimized for challenging FFPE samples

Based on Ion AmpliSeq technology, the Oncomine Dx Target Test requires as little as 10 ng of input DNA and RNA. This enables analysis of small and challenging samples. Alternative NGS methods require more FFPE slides and hundreds of nanograms of DNA and RNA, making them less practical for routine analysis of FFPE tumor samples.

Quality controls included

The Oncomine Dx Target Test incorporates DNA, RNA, and no-template controls for automatic assessment of run success.

Ion PGM Dx Sequencer

Targeted sequencing is performed on an Ion PGM™ Dx Sequencer using an Ion 318™ Dx Chip, which can accommodate up to 5.5 million reads and 6 patient samples per run (barcode adapters for multiplexing included). Run setup is fast with an easy user interface, and sequencing run time is approximately 4.5 hours. Data analysis and reporting are fully automated and streamlined using Torrent Suite™ Dx Software v5.12.5 or later.

A complete and flexible system

The Oncomine Dx Target Test is used in conjunction with the Ion PGM Dx sequencing system, which includes a complete NGS system of instruments, reagents, and software, initially validated using challenging germline variants and now validated with the Oncomine Dx Target Test for somatic mutation reporting for FFPE samples. The Ion PGM Dx sequencing system is a Class II 510K Medical Device and incorporates combined functionality, with both “IVD Mode” for molecular diagnostic tests and “Assay Development Mode” for clinical research. The system also facilitates 21 CFR Part 11 compliance, role-based workflows, sample and reagent tracking, QC metrics, and audit trails.



Oncomine Dx Target Test for NSCLC

Oncomine Dx Target Test—content

The Oncomine Dx Target Test includes targets for cancer-associated genes that all play an important role in NSCLC pathogenesis. Five of them are companion diagnostics to aid in selecting patients for approved targeted therapies, while others are currently being investigated in clinical trials and are potentially actionable in the future as referenced in Figure 3. The Oncomine Dx Target Test is indicated as a companion diagnostic to aid in selecting NSCLC patients for treatment with the five targeted therapies listed in Table 1, in accordance with the approved therapeutic product labeling. See Drugs@FDA Database.

Gene targets included for NSCLC				
Gene targets for therapeutic use				
<i>BRAF</i> : V600E	<i>EGFR</i> : L858R, exon 19 deletions, and exon 20 insertions	<i>ROS1</i> : fusions	<i>RET</i> : fusions	
Analytically validated targets				
	<i>KRAS</i>	<i>MET</i> *	<i>PIK3CA</i>	
Additional targets**				
<i>AKT1</i>	<i>ERBB2</i>	<i>HRAS</i>	<i>MTOR</i>	<i>RET</i>
<i>ALK</i> *	<i>ERBB3</i>	<i>KIT</i>	<i>NRAS</i>	<i>ROS1</i>
<i>CDK4</i>	<i>FGFR2</i>	<i>MAP2K1</i>	<i>PDGFRA</i>	
<i>DDR2</i>	<i>FGFR3</i>	<i>MAP2K2</i>	<i>RAF1</i>	

Figure 3. Complete gene list. * The test reports fusion/translocation variants for *ROS1* and *RET* only. The test only reports *ALK* and *MET* mutations.

** Performance for the additional gene target variants has been validated based on a representative method.

Table 1. Companion diagnostic biomarkers and therapies.

Gene	Variant status	Targeted therapies
<i>BRAF</i>	<i>BRAF</i> V600E	TAFINLAR® (dabrafenib) in combination with MEKINIST® (trametinib)
<i>EGFR</i>	<i>EGFR</i> L858R, exon 19 deletions	IRESSA® (gefitinib)
<i>EGFR</i>	<i>EGFR</i> exon 20 insertions	EXKIVITY™ (mobocertinib)
<i>RET</i>	<i>RET</i> fusions	GAVRETO™ (pralsetinib)
<i>ROS1</i>	<i>ROS1</i> fusions	XALKORI® (crizotinib)

Method comparison studies evaluated the accuracy of the Oncomine Dx Target Test for the detection of *BRAF* V600E, *EGFR* exon 19 deletions, L858R, and exon 20 insertions, *ROS1* fusions, and *RET* fusions, using a *BRAF* V600E PCR assay, *therascreen*[™] *EGFR* PCR kit, *ROS1* FISH assay, and validated NGS assays respectively. A summary of the concordance studies' results is included in Table 2. For details, see the User Guide.

Validation of performance for additional gene targets

The Oncomine Dx Target Test also detects DNA sequence variations in an additional 19 genes (approximately 343 targets) that are clinically associated with NSCLC. The variants for *KRAS*, *MET*, and *PIK3CA* have been analytically validated. Performance of all other variants identified by the test, other than clinically validated therapeutic variants and analytically validated variants, has not been directly demonstrated and has been validated based on a representative method.

Table 2. Concordance between the Oncomine Dx Target Test and reference methods for five companion diagnostic biomarkers.

Variants for therapy selection	Validated comparator methods	Excluding no-calls or unknowns*			Including no-calls or unknowns*		
		Positive percent agreement	Negative percent agreement	Overall percent agreement	Positive percent agreement	Negative percent agreement	Overall percent agreement
<i>BRAF</i> V600E	Validated <i>BRAF</i> V600E qPCR test	100% (67/67)	100% (114/114)	100% (181/181)	91.8% (67/73)	97.4% (114/117)	95.3% (181/190)
<i>EGFR</i>	<i>therascreen</i> [™] <i>EGFR</i> PCR kit	98.6% (71/72)	99.2% (120/121)	99.0% (191/193)	81.6% (71/87)	96.8% (120/124)	90.5% (191/211)
<i>EGFR</i> exon 19 deletions		97.6% (41/42)	99.3% (147/148)	99.0% (188/190)	74.6% (41/55)	94.2% (147/156)	89.1% (188/211)
<i>EGFR</i> exon 21 L858R		100% (30/30)	100% (167/167)	100% (197/197)	93.8% (30/32)	93.3% (167/179)	93.4% (197/211)
<i>EGFR</i> exon 20 insertions	Validated NGS Assay	100% (54/54)	100% (95/95)	100% (149/149)	98.2% (54/55)	90.5% (95/105)	93.1% (149/160)
<i>ROS1</i> fusions	Validated <i>ROS1</i> FISH test	100% (9/9)	100% (62/62)	100% (71/71)	90.0% (9/10)	88.6% (62/70)	88.8% (71/80)
<i>RET</i> fusions	Validated NGS Assay	90.9% (40/44)	91.8% (101/110)	91.6% (141/154)	90.9% (40/44)	91.8% (101/110)	91.6% (141/154)

* No-calls are for DNA variants and unknowns are for RNA fusions.

Oncomine Dx Target Test performance for NSCLC

Accuracy study

To evaluate the ability of the Oncomine Dx Target Test DNA and RNA panels to identify somatic variants in human specimens, 290 FFPE tumor samples were analyzed using the Oncomine Dx Target Test to demonstrate positive percent agreement (PPA) and negative percent agreement (NPA) concordance with validated reference detection methods. The following reference detection methods were used:

- Validated NGS method to detect single-nucleotide variant (SNV) and deletion hotspot variants
- Validated *ROS1* FISH test, to detect *ROS1* fusions

The study demonstrated a variant level PPA of 98.5%, NPA of 100%, and OPA of 100%, excluding invalids and no-calls; and a PPA level of 98.5%, NPA of 96.8%, and OPA of 96.8% including no-calls. A summary of the data is included in Table 3. For details, see the User Manual.

Establishment of limit of detection

Four limit of detection (LoD) studies were performed to evaluate DNA variants, *ROS1* fusions, *RET* fusions, and *EGFR* exon 20 insertions.

Study I: The LoD was evaluated for 14 representative DNA variants representing 3 variant categories detected by the Oncomine Dx Target Test. The LoD is the lowest allele frequency of SNV, multi-nucleotide polymorphism (MNP), or deletion variants that can be detected at least 95% of the time. The study demonstrated that the Oncomine Dx Target Test can detect DNA variants with allele frequencies between 6 and 8%.

Study II: The LoD was calculated for 2 clinical *ROS1* RNA fusion isoforms using the updated RNA library preparation workflow, and determined at 516 fusion reads.

Study III: The LoD was calculated for 2 clinical *RET* fusion isoforms using the updated RNA library preparation workflow, and determined at 405 fusion reads.

Study IV: The LoD was calculated for 2 clinical *EGFR* exon 20 insertion positive samples, and determined to be 4.8-5.2% allele frequencies.

Assay reproducibility study

Four reproducibility studies were performed to evaluate DNA variants, *ROS1* fusions, *RET* fusions, and *EGFR* exon 20 insertions.

Study I: The reproducibility and repeatability of the Oncomine Dx Target Test was evaluated for 30 representative variants from 18 DNA samples. The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instruments (reproducibility). Due to the large number of variants detected by the test and the rarity of some of the variants, a representative variant approach was used. Variants were selected in the following categories:

- Simple SNVs
- Complex SNVs and MNPs, including SNVs in di- or tri-nucleotide repeat regions and SNVs in high-GC (>60%) or low-GC (<40%) content regions
- Deletions (including deletions of 6, 9, 15, and 18 bp)

Table 3. Variant level accuracy study results.

Variant level measure of agreement	Percent agreement excluding no-calls (N)	Percent agreement including no-calls (N)
Positive percent agreement	98.5% (195/198)	98.5% (195/198)
Negative percent agreement	100.0% (118, 155/118, 159)	96.8% (118, 155/122, 012)
Overall percent agreement	100.0% (118, 350/118, 357)	96.8% (118, 350/122, 210)

Excluding no-calls, the percent of correct calls is >96%. The estimate of repeatability at each DNA variant location across all the samples was $\geq 98.8\%$ (95% CI lower limit of $\geq 97.5\%$). A summary of the results of Study I is included in Table 4. For details, see the User Manual.

Study II: An additional study was performed to evaluate the reproducibility and repeatability of the OncoPrint Dx Target Test for 6 representative variants from 11 DNA samples and 4 RNA samples. One wild-type (WT) DNA sample and 4 WT RNA samples were included in the study.

The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instrument platforms (reproducibility). The updated RNA library preparation workflow was used. Due to the large number of variants detected by the test and the rarity of some variants, a representative variant approach was used. Variants were selected in the following categories:

- 15 bp deletion
- Simple SNVs
- Complex SNVs and MNPs
- Fusions

Excluding no-calls, the estimate of repeatability at each DNA variant location across all the samples was $\geq 94.4\%$ (95% CI lower limit of $\geq 72.7\%$). The estimate of repeatability at each RNA clinical variant location was 100%. A summary of the results of Study II is included in Tables 5 and 6.

Study III: An additional study was performed to evaluate the reproducibility and repeatability of the OncoPrint Dx Target Test for 4 *RET* fusion positive samples and 2 *RET* fusion-negative samples.

The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instrument platforms (reproducibility). The updated RNA library preparation workflow was used.

Excluding unknowns, estimates of the repeatability ranged from 98.1% to 100% for two *RET* variants. A summary of the results of Study III is included in Table 7.

Study IV: A study was performed to evaluate the reproducibility and repeatability of the OncoPrint Dx Target Test for detection of *EGFR* exon 20 insertion variants using FFPE DNA from 2 *EGFR* variant-positive samples (blended with WT clinical samples) and 2 *EGFR* variant-negative (WT) samples.

The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instrument platforms (reproducibility).

Excluding no-calls, estimates of the repeatability is 100% for both *EGFR* exon 20 insertion variants. A summary of the results of Study IV is included in Table 8.

Table 4. Study I—assay reproducibility study results.

Description	No. of variants	Call rate excluding no-calls		Call rate including no-calls	
		Mean	Median	Mean	Median
DNA positive variants (positive calls)	46	96.60%	97.10%	94.50%	95.80%
WT DNA variant locations (negative calls)	872	96.10%	95.00%	96.10%	95.00%

Table 5. Study II—assay reproducibility study results (DNA variants).

Description	No. of variants	Call rate excluding no-calls		Call rate including no-calls	
		Mean	Median	Mean	Median
DNA positive variants (positive calls)	11	99%	100%	98%	99%
WT DNA variant locations (negative calls)	367	100%	100%	99%	100%

Table 6. Study II—assay reproducibility study results (*ROS1* fusions)

Description	No. of variants	Call rate including or excluding unknowns	
		Mean	Median
<i>ROS1</i> positive variants (positive calls)	4	100%	100%
WT RNA variant locations (negative calls)	4	99%	100%

Table 7. Study III—assay reproducibility study results (*RET* fusions)

Description	No. of variants	Call rate including or excluding unknowns	
		Mean	Median
<i>RET</i> positive variants (positive calls)	4	99%	100%
WT RNA variant locations (negative calls)	2	100%	100%

Table 8. Study IV—assay reproducibility study results (*EGFR* exon 20 insertions).

Description	No. of variants	Call rate excluding no-calls		Call rate including no-calls	
		Mean	Median	Mean	Median
<i>EGFR</i> insertion positive variants (positive calls)	2	100%	100%	100%	100%
WT DNA variants (negative calls)	2	100%	100%	100%	100%

OncoPrint Dx Target Test report for Non-small Cell Lung Cancer

The Clinical Test Report for the OncoPrint Dx Target Test is automatically generated as a PDF and incorporates relevant patient, sample, and test information required to help ensure high performance standards, and to assist with regulatory compliance and quality control. The test results are presented in two parts: companion diagnostic biomarker results with associated therapy indications, and analytically detected NSCLC-associated biomarker results in a separate section. The report is laboratory information management system (LIMS)-compatible and customizable for sample details.

Test Labs
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 Tel: +1 650 867 7979 | Fax: +1 650 867 7980
 contactus@testlabs.com
 www.testlabs.com

Clinical Test Report: OncoPrint™ Dx Target Test US

Patient ID: _____ Date Of Birth: _____

Sample Details

Cancer Type:	Non-small Cell Lung Cancer	Ordering Physician:	Sample Type:
Accession Number:	Physician Org:	Sample ID:	Collection Date:
Patient ID:	Physician Phone:	Receive Time:	%Cellularity:
Gender:	Physician Fax:	Sample Source:	Reference Interval:
Date Of Birth:	Pathologist:	% Necrosis:	
Sample Condition:	Pathology Lab Org:		
MRN:	Pathology Lab Phone:		
	Pathology Lab Fax:		

Results for Sequence Variations for Therapeutic Use (For illustrative purposes only. EGFR, BRAF, ROS1, and RET are mutually exclusive.)

DNA Sequence Variants

Gene	Display Name	Amino Acid Change	Nucleotide Change	Test Result	Hotspot ID	Associated Therapy
EGFR	EGFR L858R	p.Leu858Arg	c.2573T>G	POSITIVE	COSM6224	IRESSA® (gefitinib)
EGFR	EGFR exon 20 insertions	p.Ala767_Ser768 insSerValAsp	c.2311_2312ins GCGTGGACA	POSITIVE	COSM13428	EXKIVITY™ (mobocertinib)
BRAF	BRAF V600E	p.Val600Glu	c.1799T>A	POSITIVE	COSM476	TAFINLAR®+ MEKINIST® (dabrafenib in combination with trametinib)

Gene Fusions

Gene	Display Name	Test Result	Associated Therapy
ROS1	ROS1 Fusions	POSITIVE	XALKORI® (crizotinib)
RET	RET Fusions	POSITIVE	GAVRETO™ (pralsetinib)

Results for Analytical Sequence Variations Detected


DNA Sequence Variants Detected

No DNA sequence variations detected

Gene	Display Name	Nucleotide Change	Test Result	Hotspot ID
MET	p.His112Arg	c.3335A>G	NEGATIVE	COSM703
KRAS	p.Ala146Pro	c.436G>C	NEGATIVE	COSM19905
FGFR2	p.Lys659Asn	c.1977G>T	NO CALL	COSM49173
AKT1	p.Glu17Lys	c.49G>A	NEGATIVE	COSM33765
.....				

Lab Director: _____ CLIA number: _____

Report generated by Life Technologies PGM Dx Torrent Suite Software v5.12.5
For In Vitro Diagnostic Use.



1

Section 1. Includes the patient ID, date of birth, date of the report, and specifics such as the cancer type, sample type, and quality, source, and pathologic characteristics customizable by the lab.

2

Section 2. Includes results of the companion diagnostic markers, with associated therapy indications. For illustrative purposes only. EGFR, BRAF, ROS1, and RET are mutually exclusive.

3

Section 3. Contains results of the additional analytically detected DNA biomarkers—here, for illustrative purposes, only a few rows are shown. The real report will, however, contain results of all remaining over 300 variants detectable by the test, and will therefore be several pages long.

Figure 4. Example of OncoPrint Dx Target Test report format.

Oncomine Dx Target Test for Cholangiocarcinoma (CC)

Oncomine Dx Target Test—content

The Oncomine Dx Target Test includes *IDH1* R132 mutations as a companion diagnostics to aid in selecting cholangiocarcinoma (CC) patients for TIBSOVO® (ivosidenib), in accordance with the approved therapeutic product labeling, referenced in Table 9. See Drugs@FDA Database.

Table 9. Gene targets for CC.

	Gene Targets	Targeted Therapies
List of Genes for Therapeutic Use	<i>IDH1</i> R132C <i>IDH1</i> R132G <i>IDH1</i> R132H <i>IDH1</i> R132L <i>IDH1</i> R132S	TIBSOVO® (ivosidenib)

Establishment of limit of detection

The limit of detection (LoD) was evaluated for 5 *IDH1* R132 variants detected by the Oncomine Dx Target Test. The LoD is the lowest allele frequency of SNV that can be detected at least 95% of the time. The study demonstrated LoD of the 5 *IDH1* R132 variants ranged from 4.5-5.7% allele frequencies, including 4.5% for R132C, 5.7% for R132G, 4.9% for R132H, 5.1% for R132L, and 5.3% for R132S.

Assay reproducibility study

The reproducibility and repeatability of *IDH1* R132 variant detection using Oncomine Dx Target Test were assessed with 1 *IDH1* WT sample and 3 *IDH1* R132 variant positive samples at two allelic frequency (AF) levels. Testing was performed at 4 testing sites, each site had 2 PGM Dx instrument systems, 2 operators, and using 4 lots of reagents. The overall positive call rate for *IDH1* R132 variants was 92.6% when including no calls and 97.1% when excluding no calls. The negative call rate for *IDH1* WT sample were 100% at all *IDH1* R132 variant locations. (Table 10).

Clinical Study

To evaluate the ability of the Oncomine Dx Target Test to identify five *IDH1* biomarkers in FFPE cholangiocarcinoma tumor specimens, 168 specimens from patients tested positive and 181 specimens tested negative using Sanger assay were tested using Oncomine Dx Target Test to demonstrate positive percent agreement (PPA) and negative percent agreement (NPA) concordance with Sanger assay as validated reference detection method.

The study demonstrated PPA of 99.4%, NPA of 96.5%, and OPA of 97.9%, excluding invalids and no calls; and PPA level of 97.0%, NPA of 90.6% and OPA of 93.7% including no calls. A summary of the data is included in Table 11. For details, see the User Manual.

Oncomine Dx Target Test for Cholangiocarcinoma (CC)

Table 10. Reproducibility results

Sample COSMIC ID, Variant	No. of valid sample results	Call rate (95% CI)	
		Including no calls	Excluding no calls
D1 COSM28747, R132C	36	100% (90.3%, 100%)	100% (90.3%, 100%)
D2 COSM28747, R132C	36	97.2% (85.5%, 99.9%)	100% (90.0%, 100%)
D3 COSM28749, R132G	36	100% (90.3%, 100%)	100% (90.3%, 100%)
D4 COSM28749, R132G	36	100% (90.3%, 100%)	100% (90.3%, 100%)
D5 COSM28750, R132L	36	100% (90.3%, 100%)	100% (90.3%, 100%)
D6 COSM28750, R132L	35	57.1% (39.4%, 73.7%)	76.9% (56.4%, 91.0%)
D1-D6 All Variants, R132	215	92.6% (88.2%, 95.7%)	97.1% (93.7%, 98.9%)
D7 Wild-Type	36	100% (90.3%, 100%)	100% (90.3%, 100%)

Table 11. Concordance between the Oncomine Dx Target Test and reference method for *IDH1* R132 mutations

Variants for therapy selection	Validated comparator methods	Excluding invalid results and no-calls			Including invalid results and no-calls		
		Positive percent agreement	Negative percent agreement	Overall percent agreement	Positive percent agreement	Negative percent agreement	Overall percent agreement
<i>IDH1</i> R132	Validated Sanger assay	99.4% (163/164)	96.5% (164/170)	97.9% (327/334)	97.0% (163/168)	90.6% (164/181)	93.7% (327/349)

Oncomine Dx Target Test Report for Cholangiocarcinoma (CC)

The Clinical Test Report for the Oncomine Dx Target Test is automatically generated as a PDF and incorporates relevant patient, sample, and test information required to help ensure high performance standards, and to assist with regulatory compliance and quality control. The test results are presented in two parts: companion diagnostic biomarker results with associated therapy indications, and analytically detected biomarker results in a separate section. The report is laboratory information management system (LIMS) system-compatible and customizable for sample details.

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 contactus@testlabs.com
 www.testlabs.com

Clinical Test Report: Oncomine™ Dx Target Test US

Patient ID: _____ Date Of Birth: _____ Date: _____

Sample Details

Cancer Type: Cholangiocarcinoma	Ordering Physician: _____	Sample Type: FFPE, Block
Accession Number: 0826_100	Physician Org: _____	Sample ID: _____
Patient ID: _____	Physician Phone: _____	Collection Date: _____
Gender: _____	Physician Fax: _____	Receive Time: _____
Date Of Birth: _____	Pathologist: _____	%Cellularity: _____
Sample Condition: _____	Pathology Lab Org: _____	Sample Source: _____
MRN: _____	Pathology Lab Phone: _____	Reference Interval: _____
	Pathology Lab Fax: _____	% Necrosis: _____


Results for Sequence Variations for Therapeutic Use (for illustrative purposes only)

DNA Sequence Variants

Gene	Display Name	Amino Acid Change	Nucleotide Change	Test Result	Hotspot ID	Associated Therapy
IDH1	IDH1 R132	p.Arg132Gly	c.394C>G	POSITIVE	CO5M28749	TIBSOVO® (ivosidenib)

Lab Director: _____ **CLIA number:** _____

Report generated by Life Technologies PGM Dx Torrent Suite Software v5.12.5
For In Vitro Diagnostic Use.



1

Section 1. Includes the patient ID, date of birth, date of the report, and specifics such as the cancer type, sample type and quality, source, and pathologic characteristics customizable by the lab.

2

Section 2. Includes results of the companion diagnostic markers, with associated therapy indications.

Figure 5. Example of Oncomine Dx Target Test report format.

Ordering information

Product	Cat. No.
Oncomine Dx Target Test includes:	A51695
Ion Torrent Dx FFPE Sample Preparation Kit	A32445
Oncomine Dx Target Test, Controls, and Diluent Kit	A49756
Ion PGM Dx Library Kit	A49758
Ion PGM Dx OneTouch Template Kit	A49759
Ion PGM Dx Sequencing Kit	A49760
Ion PGM Dx 318 Chip Kit	A18937
Oncomine Dx Target Test User Guides and Assay Definition File	A51694
Ion PGM Dx Instrument System includes:	A25511
• Ion PGM Dx Sequencer	
• Ion OneTouch Dx Instrument	
• Ion PGM Dx System Installation and Training Kit	
• Ion PGM Dx Chip Minifuge	
• Ion PGM Wireless Scanner	
• Ion Torrent Server with Ion PGM Dx Software Pack v5.12.5 (Torrent Suite Dx Software v5.12.5 and Torrent Suite Assay Development Software v5.12.5)	

Find out more at thermofisher.com/oncomine-dxtarget

ThermoFisher
SCIENTIFIC

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Abbreviated Intended Use: The Oncomine Dx Target Test is a qualitative *in vitro* diagnostic test that uses targeted high-throughput, parallel-sequencing technology to detect single-nucleotide variants (SNVs), deletions, and insertions in 23 genes from DNA and fusions in *ROS1* and *RET* from RNA isolated from formalin-fixed, paraffin-embedded (FFPE) tumor tissue samples from patients with non-small cell lung cancer (NSCLC) and *IDH1* R132 mutations from FFPE tumor tissue samples from patients with cholangiocarcinoma (CC) using the Ion PGM Dx System (MAN0018810).

Oncomine Dx Target Test

Epidemiology of lung cancer

Lung cancer is the leading cause of cancer deaths in the United States [1]. In 2020, an estimated 228,820 new cases (116,300 in men and 112,520 in women) of lung and bronchial cancer will be diagnosed, and 135,720 deaths (72,500 in men and 63,220 in women) are estimated to occur because of the disease [2]. Only 19.4% of all patients with lung cancer live 5 years or more after diagnosis [3].

Genetic companion diagnostic testing for targeted therapy selection in NSCLC

Lung cancer comprises two main histologic subtypes: non-small cell lung cancer (NSCLC) and small cell lung cancer (SCLC). Over the past decade, several biomarkers associated with therapeutic benefit have emerged for NSCLC. To obtain comprehensive molecular biomarker profiling, multiplexing technology such as next-generation sequencing (NGS) is recommended by IASLC/AMP* NSCLC testing guidelines, given the limited tissue.

For the most current information concerning the essential biomarkers for lung cancer and their association with therapeutic outcomes, refer to the therapeutic labels available at “Drugs@FDA” on the FDA website.

EGFR: *EGFR* exon 19 deletions and the L858R mutation are found in approximately 10% of Caucasian patients with NSCLC and up to 50% of Asian NSCLC patients [4]. These mutations result in activation of the tyrosine kinase domain, and are associated with sensitivity to small molecule tyrosine kinase inhibitors (TKIs), such as erlotinib, gefitinib, and afatinib [5]. Data show that erlotinib, gefitinib, or afatinib (instead of standard first-line chemotherapy) should be used in patients with *EGFR* exon 19 deletions and the L858R mutation [6-11]. *EGFR* companion diagnostic tests have been approved by the FDA for specific drug indications, including the *therascreen*[™] *EGFR* RGQ PCR Kit by Qiagen for gefitinib and afatinib, the cobas[™] *EGFR* Mutation Test v2 by Roche for erlotinib, and the Ion Torrent[™] Oncomine[™] Dx Target Test by Thermo Fisher Scientific for gefitinib.

EGFR: *EGFR* exon 20 insertions are much less common and seen in approximately 2% of non-squamous NSCLC (in the Caucasian population), or 12% of all *EGFR* mutations [32]. *EGFR* exon 20 insertions are typically represented by in-frame insertion of 3 to 21 base pairs, or 1 to 7 amino acids, involving codons 761 to 775 [33]. NSCLC with *EGFR* exon 20 insertions, with the exception of the A763_Y764insFQEA variant, do not typically respond to first- and second-generation tyrosine kinase inhibitors (TKIs) or anti-PD-L1 treatments [32]. Over 60 unique variants of *EGFR* exon 20 insertions have been identified through comprehensive genomic profiling, the majority of which are rare variants [32]. Mobocertinib is approved by the FDA to treat metastatic NSCLC patients with *EGFR* exon 20 insertions who have received prior platinum-based chemotherapy. The Oncomine Dx Target Test is approved by the FDA as a companion diagnostic test for detection of *EGFR* exon 20 insertions.

BRAF: It is estimated that *BRAF* mutations occur in about 3–5% of patients with NSCLC [17]. Dabrafenib in combination with trametinib is approved by the FDA to treat NSCLC patients with a *BRAF* V600E mutation. The Oncomine Dx Target Test is approved by the FDA as a companion diagnostic test for detection of *BRAF* V600E mutation.

ALK: It is estimated that 2–7% of patients with NSCLC have an *ALK* gene rearrangement [12]. Crizotinib is approved by the FDA to treat people with NSCLC that has spread to other parts of the body and is caused by either an *ALK* fusion or a *ROS1* fusion. Molecular diagnostic testing using FISH and immunohistochemistry (IHC), which are the standard methods for *ALK* in NSCLC, have been approved by the FDA for detecting *ALK* fusions and *ALK* expression, respectively [13,14].

While NGS can also be used to assess the presence of an *ALK* fusion, the Oncomine Dx Target Test does not detect *ALK* fusions. Two *ALK* companion diagnostic tests have been approved by the FDA for use with crizotinib, including the Vysis™ *ALK* Break Apart FISH Probe Kit by Abbott Molecular, and the VENTANA™ *ALK* (D5F3) CDx Assay by Roche.

***ROS1*:** It is estimated that *ROS1* fusions occur in about 1–2% of patients with NSCLC [15]. *ROS1* is very similar to *ALK* (77% amino acid sequence homology in the ATP binding sites of the tyrosine kinase domain) and both are members of the insulin receptor family. Crizotinib is very effective for NSCLC patients with *ROS1* rearrangements [16]. The Oncomine Dx Target Test is approved by the FDA as a companion diagnostic test for detection of *ROS1* fusions.

***RET*:** *RET* fusions occur in ~1–2% of lung carcinomas [18]. Pralsetinib is approved by the FDA to treat NSCLC patients with *RET* fusions. The Oncomine Dx Target Test is approved by FDA as a companion diagnostic test for detection of *RET* fusions.

Epidemiology of Cholangiocarcinoma

Cholangiocarcinoma, i.e., bile duct cancer, is classified according to its anatomical location in relation to the liver, being either intrahepatic or extrahepatic. Extrahepatic cholangiocarcinomas, aka perihilar (or referred to as Klatskin tumor) or distal bile duct cancers, are more common than intrahepatic cholangiocarcinomas.[19] Each year, about 8,000 people in the United States are diagnosed with cholangiocarcinoma.[20] Average 5-year survival rate (2008-2014) is estimated at about 8% for intrahepatic bile duct cancers, with 24% for localized and as low as 1% for metastatic, and 10% for extrahepatic bile duct cancers, at 13% for localized and only 1% for metastatic tumors.[21]

***IDH1/2*:** *IDH1/2* mutations are the most commonly observed genomic alteration found in intrahepatic cholangiocarcinomas (10-23% of cases). [22-28] Mutations in *IDH1* have been found in approximately 70% of Grade 2 to 3 gliomas,[29] 50% of chondrosarcomas,[30] and up to 20% of cholangiocarcinomas.[31] Ivosidenib has been approved by FDA for the treatment of adult patients with locally advanced or metastatic cholangiocarcinoma who have been previously treated with gemcitabine- or fluorouracil (5-FU)-based regimens with an *IDH1* mutation as detected by an FDA-approved test. For more information of the biomarker and therapeutic outcomes, refer to the therapeutic labels available at Drugs@FDA on the FDA website.

Test intended use/indications for use

The Oncomine Dx Target Test is a qualitative *in vitro* diagnostic (IVD) test that uses high-throughput targeted, parallel sequencing technology to detect single-nucleotide variants (SNVs), deletions, and insertions in 23 genes from DNA and fusions in *ROS1* and *RET* from RNA isolated from formalin-fixed, paraffin-embedded (FFPE) tumor samples from patients with NSCLC, and *IDH1* R132 mutations from FFPE tumor tissue samples from patients with CC using the Ion PGM™ Dx System.

The test is indicated to aid in selecting NSCLC and CC patients for treatment with the targeted therapies listed in Table 1 in accordance with the approved therapeutic product labeling.

Table 1. Variants for therapeutic use.

Non-small Cell Lung Cancer (NSCLC)

Gene	Variant status	Targeted therapy
<i>BRAF</i>	<i>BRAF</i> V600E	TAFINLAR® (dabrafenib) in combination with MEKINIST® (trametinib)
<i>EGFR</i>	<i>EGFR</i> L858R, exon 19 deletions	IRESSA® (gefitinib)
<i>EGFR</i>	<i>EGFR</i> exon 20 insertions	EXKIVITY™ (mobocertinib)
<i>RET</i>	<i>RET</i> fusions	GAVRETO™ (pralsetinib)
<i>ROS1</i>	<i>ROS1</i> fusion	XALKORI® (crizotinib)

Cholangiocarcinoma (CC)

Gene	Variant status	Targeted therapy
<i>IDH1</i>	<i>IDH1</i> R132C <i>IDH1</i> R132G <i>IDH1</i> R132H <i>IDH1</i> R132L <i>IDH1</i> R132S	TIBSOVO® (ivosidenib)

Safe and effective use has not been established for selecting therapies using this device for the variants other than those listed in Table 1. Results other than those listed in Table 1 are indicated for use only in patients who have already been considered for all appropriate therapies (including those listed in Table 1).

Analytical performance using NSCLC specimens has been established for the variants listed in Table 2. Safety and effectiveness of these three genes and four variants have not been established and they are not intended to be used to direct therapy.

Table 2. Variants with established analytical performance only.

Gene	Variant ID	Amino acid change	Nucleotide change
<i>KRAS</i>	COSM512	p.Gly12Phe	c.34_35delGGinsTT
<i>KRAS</i>	COSM516	p.Gly12Cys	c.34G>T
<i>MET</i>	COSM707	p.Thr1010Ile	c.3029C>T
<i>PIK3CA</i>	COSM754	p.Asn345Lys	c.1035T>A

The test is not indicated to be used for stand-alone diagnostic purposes, screening, monitoring, risk assessment, or prognosis.

Test performance and characteristics - NSCLC

Oncomine Dx Target Test detects over 300 variants in 23 genes, with active clinical trials and/or having demonstrated association with NSCLC in the literature. Summary of reported variants in NSCLC:

DNA: *AKT1*, *ALK*, *BRAF*, *CDK4*, *DDR2*, *EGFR*, *ERBB2*, *ERBB3*, *FGFR2*, *FGFR3*, *HRAS*, *KIT*, *KRAS*, *MAP2K1*, *MAP2K2*, *MET*, *MTOR*, *NRAS*, *PDGFRA*, *PIK3CA*, *RAF1*, *RET*, and *ROS1*

RNA: *RET* and *ROS1*

Analytical validation of the Oncomine Dx Target Test was established through a series of studies to assess the accuracy, sensitivity, specificity, and reproducibility of the assay for the detection of SNVs, deletions, insertions, and fusions [34].

Based on the data observed, the test demonstrated a limit of detection (LoD) of between 6–8% allele frequencies for DNA variants, 4.8–5.2% allele frequencies for *EGFR* insertions, 516 fusion reads for *ROS1* fusions, and 405 fusion reads for *RET* fusions, with 95% confidence.

Additionally, based on the representative variants that were tested in the accuracy study, the test detected variants with 98.5% positive percent agreement (PPA) and 100% negative percent agreement (NPA) against validated reference methods (excluding no-calls).

Four studies were conducted to evaluate the repeatability and reproducibility of the test for DNA variants, *EGFR* exon 20 insertions, *ROS1* fusions, and *RET* fusions. Repeatability is >94% for DNA variants, 100% for *EGFR* exon 20 insertions, 100% for *ROS1* fusions, and >98% for *RET* fusions. Reproducibility is >99% for DNA variants, *EGFR* exon 20 insertions, *ROS1* fusions, and *RET* fusions (excluding unknowns).

Method comparison studies evaluated the concordance of the test for the detection of *BRAF* V600E, *EGFR* exon 19 deletions, L858R, and exon 20 insertions, *ROS1* fusions, and *RET* fusions, using a *BRAF* V600E PCR assay, *therascreen*™ *EGFR* PCR kit, *ROS1* FISH assay, and validated NGS assays, respectively. The studies show:

- 100% overall percent agreement (OPA), positive percent agreement (PPA) and negative percent agreement (NPA) for *BRAF*, *EGFR* exon 20 insertions, and *ROS1* fusions
- 99% OPA, PPA, and NPA for *EGFR* exon 19 deletions and L858R
- 92% OPA, 91% PPA, and 92% NPA for *RET* fusions

Test performance and characteristics - CC

Oncomine Dx Target Test only reports *IDH1* R132 in CC. Analytical validation of the Oncomine Dx Target Test in CC was established through LoD and precision studies [34].

Based on the data observed, the test confirmed a limit of detection (LoD) of 4.5-5.7% allele frequencies for 5 *IDH1* R132 mutations, including 4.5% for R132C, 5.7% for R132G, 4.9% for R132H, 5.1% for R132L, and 5.3% for R132S. Repeatability and reproducibility study demonstrated overall positive call rate of 92.6% for the *IDH1* R132 variants when including no calls and 97.1% when excluding no calls. The negative call rates for the *IDH1* WT sample were 100%.

A clinical concordance study was conducted to evaluate the ability of the Oncomine Dx Target Test to identify five *IDH1* biomarkers in FFPE cholangiocarcinoma tumor specimen compared to a validated Sanger assay. The study shows 99.4% positive percent agreement (PPA), 96.5% negative percent agreement (NPA), and 97.9% overall percent agreement (OPA) excluding invalid results and no-calls.

Guide to interpreting results

Test results should be interpreted in the context of pathological evaluation of tumors, treatment history, clinical findings, and other laboratory data.

All clinical interpretations of the variants detected should be made by a board-certified pathologist or equivalent. It is recommended that the physician ordering the test consult with a board-certified pathologist. Patients are advised to seek information from their oncologist or certified health care provider.

Additional information may be obtained from NCCN Guidelines™ and IASLC/AMP NSCLC testing guidelines.

The molecular profile of a tumor can vary between primary and metastatic sites, as well as change over time in response to treatment, leading to the development of mutations that could confer resistance to therapeutic agents.

Test limitations and warnings

The test is designed to interrogate over 300 variants in 23 genes associated with NSCLC and one gene in CC. However, when certain quality metrics and controls established for the specimen testing are not met, accuracy of the test cannot be assured and therefore mutation status in the exons is reported. Variants detected by the panel that are not clinically or analytically validated should not be used for selecting treatment.

- This test does not detect genomic copy number variants.
- The Oncomine Dx Target Test does not detect *ALK* fusions.
- This test does not detect structural variants in genes other than *ROS1* and *RET*.
- Rare polymorphisms exist that could lead to false-negative or false-positive results.
- A negative (wild-type) result does not rule out the presence of a mutation that is below the limits of detection of this assay (6–8%).
- The product is designed to detect a targeted set of known variants in the genes. New variants that are not included in the test may be discovered in the future.
- For NSCLC, the Oncomine Dx Target Test assay definition file includes prevalent but not all rare or newly identified *RET* isoforms, *ROS1* isoforms, and *EGFR* exon 20 insertions. The Oncomine Dx Target Test may miss rare or newly identified:
 - *RET* isoforms carried by a subset of patients who may derive benefit from GAVRETO™ (pralsetinib)
 - *ROS1* isoforms carried by a subset of patients who may derive benefit from XALKORI® (crizotinib)
 - *EGFR* exon 20 insertions carried by a subset of patients who may derive benefit from EXKIVITY™ (mobocertinib)

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Find out more at thermofisher.com/oncomine-dxtarget

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Oncomine Dx Target Test— FDA approved

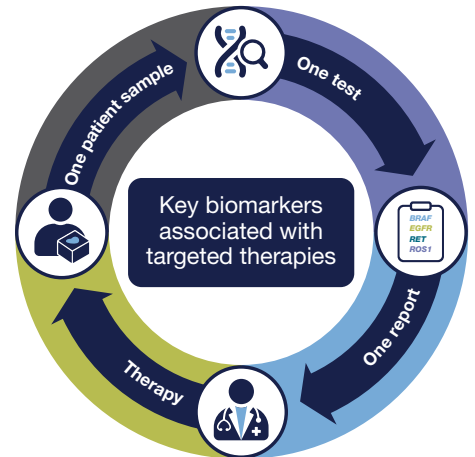
Tell your oncologist about Oncomine Dx Target Test

A new paradigm in testing for targeted therapies in NSCLC and CC

This test is now reimbursed by Medicare and the top 40 commercial payers, covering over 200 million US lives.

The Ion Torrent™ Oncomine™ Dx Target Test is the first targeted next-generation sequencing (NGS) *in vitro* diagnostic test for non-small cell lung cancer (NSCLC) and cholangiocarcinoma (CC), simultaneously delivering multiple biomarker results for multiple targeted therapies from one sample within 4 days.

- **Identify patients for multiple therapies**—one test indicated as a companion diagnostic (CDx) device to aid in selecting NSCLC and CC patients for treatment with targeted therapies.
- **Multiple biomarkers from one limited sample**—one test for detection of 23 genes, minimizing the risk of depleting tissues and requiring additional biopsies. Based on Ion AmpliSeq™ technology, the required input is as low as 10 ng of DNA and RNA.
- **One workflow, helps save time**—laboratory results can be generated within four days.
- **Established performance**—concordance with FDA approved or validated reference methods based on FISH, PCR, Sanger, or NGS was established for all CDx biomarkers:
 - 100% overall percent agreement (OPA), positive percent agreement (PPA) and negative percent agreement (NPA) for *BRAF*, *EGFR* exon 20 insertions, and *ROS1* fusions
 - 99% OPA, PPA, and NPA for *EGFR* exon 19 deletions and L858R
 - 92% OPA, 91% PPA, and 92% NPA for *RET* fusions
 - 98% OPA, 99% PPA, and 97% NPA for *IDH1*



Cancer type	Gene	Targeted therapies
NSCLC	<i>BRAF</i>	TAFINLAR® (dabrafenib) in combination with MEKINIST® (trametinib)
	<i>EGFR</i> L858R, exon 19 deletions	IRESSA® (gefitinib)
	<i>EGFR</i> exon 20 insertions	EXKIVITY™ (mobicertinib)
	<i>RET</i>	GAVRETO™ (pralsetinib)
CC	<i>ROS1</i>	XALKORI® (crizotinib)
	<i>IDH1</i>	TIBSOVO® (ivosidenib)

Figure 1. List of genes for therapeutic use.

A complete and flexible system

The Oncomine Dx Target Test is used in conjunction with the Ion PGM™ Dx System, which includes a complete NGS system of instruments, reagents, and software, now validated with the Oncomine Dx Target Test for somatic mutation reporting for FFPE samples (see Figure 2 for workflow). The Ion PGM Dx sequencing system is a Class II Medical Device and incorporates combined functionality, with both “IVD Mode” for molecular diagnostic tests and “Assay Development Mode” for clinical research. The system also facilitates 21 CFR Part 11 compliance, with role-based workflows, sample and reagent tracking, QC metrics, and audit trails.

Oncomine Dx Target Test—gene content

The Oncomine Dx Target Test includes targets for cancer-associated genes. Six of them are companion diagnostics to aid in selecting patients for approved targeted therapies,



Figure 2. The Oncomine Dx Target Test utilizes a single streamlined NGS workflow for detecting cancer-associated biomarkers, incorporating reagents, instrument systems, and bioinformatics. The turnaround time from FFPE sample to report is 4 days.

including five in NSCLC and one in CC, while remaining genes are currently being investigated in clinical trials and may be potentially actionable in the future as referenced in Figure 3.

Oncomine Dx Target Test—report

The Oncomine Dx Target Test report is automatically generated as a PDF and incorporates relevant patient, sample, and test information required to help ensure high performance standards, regulatory compliance, and quality control. The test results are presented in two parts: companion diagnostic biomarker results with associated therapy indication (Figure 4), and other analytically detected biomarker results in a separate section (not shown). The report is laboratory information management system (LIMS) compatible.

Genes targets for NSCLC			
Gene targets for therapeutic use			
<i>BRAF</i> : V600E	<i>EGFR</i> : L858R, exon 19 deletions, and exon 20 insertions	<i>ROS1</i> : fusions	<i>RET</i> : fusions
Analytically validated targets			
<i>KRAS</i>	<i>MET</i> *	<i>PIK3CA</i>	
Additional targets**			
<i>AKT1</i>	<i>ERBB3</i>	<i>MAP2K1</i>	<i>RAF1</i>
<i>ALK</i> *	<i>FGFR2</i>	<i>MAP2K2</i>	<i>RET</i>
<i>CDK4</i>	<i>FGFR3</i>	<i>MTOR</i>	<i>ROS1</i>
<i>DDR2</i>	<i>HRAS</i>	<i>NRAS</i>	
<i>ERBB2</i>	<i>KIT</i>	<i>PDGFRA</i>	

Figure 3. Complete gene list. * The test reports fusion/translocation variants for *ROS1* and *RET* only. The test only reports *ALK* and *MET* mutations. ** Performance for the additional gene target variants has been validated based on a representative method. Only *IDH1* is reported for CC.

NSCLC results for sequence variations for therapeutic use (for illustrative purposes only; *EGFR*, *BRAF*, *ROS1*, and *RET* are mutually exclusive)

DNA sequence variants						
Gene	Display name	Amino acid change	Nucleotide change	Test result	Hotspot ID	Associated therapy
<i>EGFR</i>	<i>EGFR</i> L858R	p.Leu858Arg	c.2573T>G	POSITIVE	COSM6224	IRESSA® (gefitinib)
<i>EGFR</i>	<i>EGFR</i> exon 20 insertions	p.Ala767_Ser768 insSerValAsp	c.2311_2312ins GCGTGGACA	POSITIVE	COSM13428	EXKIVITY™ (mabocertinib)
<i>BRAF</i>	<i>BRAF</i> V600E	p.Val600Glu	c.1799T>A	POSITIVE	COSM476	TAFINLAR® + MEKINIST® (dabrafenib in combination with trametinib)
Gene fusions (RNA)						
Gene	Display name			Test result	Associated therapy	
<i>ROS1</i>	<i>ROS1</i> fusions			POSITIVE	XALKORI® (crizotinib)	
<i>RET</i>	<i>RET</i> fusions			POSITIVE	GAVRETO™ (pralsetinib)	

CC results for sequence variations for therapeutic use (for illustrative purposes only)

DNA sequence variants						
Gene	Display name	Amino acid change	Nucleotide change	Test result	Hotspot ID	Associated therapy
<i>IDH1</i>	<i>IDH1</i> R132	p.Arg132Gly	c.394C>G	POSITIVE	COSM28749	TIBSOVO® (ivosidenib)

Figure 4. Example of Oncomine Dx Target Test report format. The report includes a section with results of the validated biomarkers and information about relevant treatment indication, as well as a separate section with the other biomarkers not validated for treatment selection (not shown).

Find out more at thermofisher.com/oncomine-dxtarget



**Oncomine Dx
Target Test—
FDA approved**

Ask your pathologist about Oncomine Dx Target Test

A new paradigm in testing for targeted therapies in
NSCLC and CC

This test is now reimbursed
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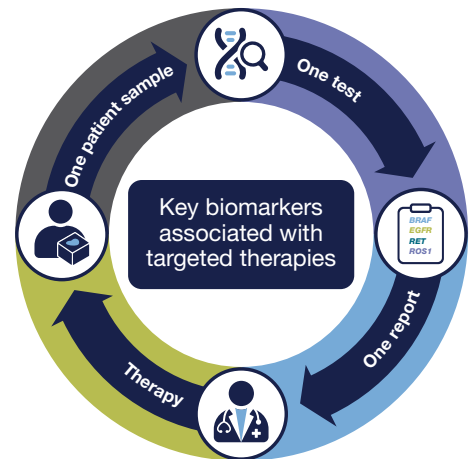
Did you know:

- Many biopsy samples are so small that they cannot be analyzed by some NGS tests, especially panels containing hundreds of genes, leading to tissue exhaustion
- It can take several weeks to get results with alternative NGS tests, potentially delaying treatment decision

Choosing the right NGS test can make a difference for your patient

The Oncomine Dx Target Test is an FDA-approved NGS CDx test that can:

- Identify patients for multiple therapies—one test indicated as a companion diagnostic (CDx) device to aid in selecting NSCLC and CC patients for treatment with targeted therapies
- Accept small samples (10 ng DNA and RNA), for more patients to potentially access targeted therapies
- Generate results in a laboratory within four days, enabling faster treatment decisions



Cancer type	Gene	Targeted therapies
NSCLC	<i>BRAF</i>	TAFINLAR® (dabrafenib) in combination with MEKINIST® (trametinib)
	<i>EGFR</i> L858R, exon 19 deletions	IRESSA® (gefitinib)
	<i>EGFR</i> exon 20 insertions	EXKIVITY™ (mobocertinib)
	<i>RET</i>	GAVRETO™ (pralsetinib)
	<i>ROS1</i>	XALKORI® (crizotinib)
CC	<i>IDH1</i>	TIBSOVO® (ivosidenib)

Figure 1. List of genes for therapeutic use.

Genes targets for NSCLC			
Gene targets for therapeutic use			
<i>BRAF</i> : V600E	<i>EGFR</i> : L858R, exon 19 deletions, and exon 20 insertions	<i>ROS1</i> : fusions	<i>RET</i> : fusions
Analytically validated targets			
<i>KRAS</i>	<i>MET</i> *	<i>PIK3CA</i>	
Additional targets**			
<i>AKT1</i>	<i>ERBB3</i>	<i>MAP2K1</i>	<i>RAF1</i>
<i>ALK</i> *	<i>FGFR2</i>	<i>MAP2K2</i>	<i>RET</i>
<i>CDK4</i>	<i>FGFR3</i>	<i>MTOR</i>	<i>ROS1</i>
<i>DDR2</i>	<i>HRAS</i>	<i>NRAS</i>	
<i>ERBB2</i>	<i>KIT</i>	<i>PDGFRA</i>	

Figure 2. Complete gene list. * The test reports fusion/translocation variants for *ROS1* and *RET* only. The test only reports *ALK* and *MET* mutations. ** Performance for the additional gene target variants has been validated based on a representative method. Only *IDH1* is reported for CC.

Oncomine Dx Target Test—performance

Concordance with FDA approved or validated reference methods based on FISH, PCR, Sanger, or NGS was established for all CDx biomarkers:

- 100% overall percent agreement (OPA), positive percent agreement (PPA) and negative percent agreement (NPA) for *BRAF*, *EGFR* exon 20 insertions, and *ROS1* fusions
- 99% OPA, PPA, and NPA for *EGFR* exon 19 deletions and L858R
- 92% OPA, 91% PPA, and 92% NPA for *RET* fusions
- 98% OPA, 99% PPA, and 97% NPA for *IDH1*

Oncomine Dx Target Test—report

NSCLC results for sequence variations for therapeutic use (for illustrative purposes only; *EGFR*, *BRAF*, *ROS1*, and *RET* are mutually exclusive)

DNA sequence variants						
Gene	Display name	Amino acid change	Nucleotide change	Test result	Hotspot ID	Associated therapy
<i>EGFR</i>	<i>EGFR</i> L858R	p.Leu858Arg	c.2573T>G	POSITIVE	COSM6224	IRESSA® (gefitinib)
<i>EGFR</i>	<i>EGFR</i> exon 20 insertions	p.Ala767_Ser768 insSerValAsp	c.2311_2312ins GCGTGGACA	POSITIVE	COSM13428	EXKIVITY™(mobocertinib)
<i>BRAF</i>	<i>BRAF</i> V600E	p.Val600Glu	c.1799T>A	POSITIVE	COSM476	TAFINLAR® + MEKINIST® (dabrafenib in combination with trametinib)
Gene fusions (RNA)						
Gene	Display name			Test result	Associated therapy	
<i>ROS1</i>	<i>ROS1</i> fusions			POSITIVE	XALKORI® (crizotinib)	
<i>RET</i>	<i>RET</i> fusions			POSITIVE	GAVRETO™ (pralsetinib)	

CC results for sequence variations for therapeutic use (for illustrative purposes only)

DNA sequence variants						
Gene	Display name	Amino acid change	Nucleotide change	Test result	Hotspot ID	Associated therapy
<i>IDH1</i>	<i>IDH1</i> R132	p.Arg132Gly	c.394C>G	POSITIVE	COSM28749	TIBSOVO® (ivosidenib)

Figure 3. Example of Oncomine Dx Target Test report format. The report includes a section with results of the validated biomarkers and information about relevant treatment indication, as well as a separate section with the other biomarkers not validated for treatment selection (not shown).

If your pathology laboratory does not perform the Oncomine Dx Target Test, you can send samples to one of these reference laboratories.

Reference lab	Telephone number	Website
Integrated Oncology (a division of LabCorp)	800-447-5816	integratedoncology.com
Quest Diagnostics, Inc.	866-697-8378	questdiagnostics.com
NeoGenomics Laboratories, Inc.	866-776-5907	neogenomics.com
Phenopath, a Quest Diagnostics Company	888-927-4366	phenopath.com

Find out more at thermofisher.com/oncomine-dxtarget



Oncomine IVD solutions— it's about time

Get faster answers for more NSCLC and CC patients

When your patients' samples are analyzed with the NGS-based Ion Torrent™ Oncomine™ Dx Target Test in your own hospital laboratory, you can get answers within a week, even from small biopsies, allowing you to make faster therapeutic decisions for more patients. No need to rely on external laboratories. Now you can bring precision oncology closer to your patients and fully own their care journey.

Find out more at thermofisher.com/oncomine-dxtarget

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Oncomine IVD solutions— it's about time

Test faster to enable faster treatment

When your patients' samples are analyzed with the NGS-based Ion Torrent™ Oncomine™ Dx Target Test in your own hospital laboratory, you can get answers within a week, even from small biopsies, allowing you to make faster therapeutic decisions for more patients. Ask your pathologist about Ion Torrent™ Oncomine™ IVD solutions.

Find out more at thermofisher.com/oncomine-dxtarget

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Oncomine IVD solutions— faster answers for more patients

When your patients' samples are analyzed with the NGS-based Ion Torrent™ Oncomine™ Dx Target Test in your own hospital laboratory, you can get answers within a week, even from small biopsies, allowing you to make faster therapeutic decisions for more patients. Ask your pathologist about Ion Torrent™ Oncomine™ IVD solutions.

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Ask for Oncomine IVD solutions

Get faster answers for more NSCLC and CC patients

When your patients' samples are analyzed with the NGS-based Ion Torrent™ Oncomine™ Dx Target Test in your own hospital laboratory, you can get answers within a week, even from small biopsies, allowing you to make faster therapeutic decisions for more patients. Ask your pathologist about Ion Torrent™ Oncomine™ IVD solutions.

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Ask for Oncomine IVD solutions

Get faster answers for more NSCLC patients

When your patients' samples are analyzed with the NGS-based Ion Torrent™ Oncomine™ Dx Target Test in your own hospital laboratory, you can get answers within a week, even from small biopsies, allowing you to make faster therapeutic decisions for more patients. No need to rely on external laboratories. Now you can bring precision oncology closer to your patients and fully own their care journey.

Find out more at thermofisher.com/oncomine-dxtarget

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Oncomine Dx Target Test—US



A new paradigm in testing for targeted therapies in NSCLC and CC

In the era of personalized medicine, molecular profiling has become essential for the treatment of cancer patients. With an increasing number of genomic alterations becoming clinically relevant, sequential testing of individual mutations becomes a significant challenge for clinical laboratories. Next-generation sequencing (NGS), which can detect multiple alterations at once from a small amount of tissue, offers a solution.

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[Download flyer for clinicians ›](#)

[Download flyer for laboratory professionals ›](#)

[Download brochure for laboratory professionals ›](#)

The Oncomine Dx Target Test is currently available in the following commercial labs:

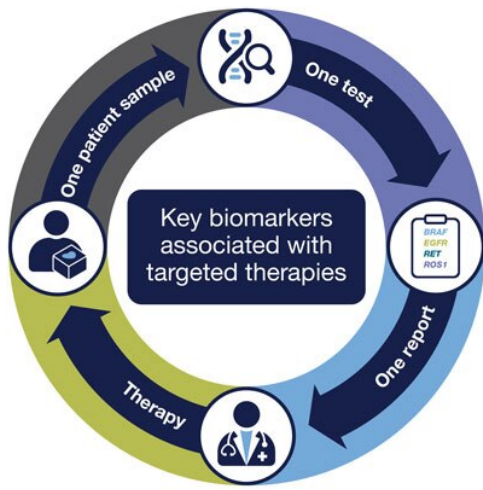
Reference lab	Telephone number	Website
Integrated Oncology (a division of LabCorp)	800-447-5816	integratedoncology.com
Quest Laboratories, Inc.	866-697-8378	questdiagnostics.com
NeoGenomics Laboratories, Inc.	866-776-5907	neogenomics.com
Phenopath, a Quest Diagnostics Company	888-927-4366	phenopath.com

One test that can expedite treatment selection decisions

The Ion Torrent Oncomine Dx Target Test is the first targeted NGS-based *in vitro* diagnostic test for non-small cell lung cancer (NSCLC) and cholangiocarcinoma (CC), simultaneously delivering multiple biomarker results for multiple targeted therapies from one sample within four days.

Cancer type	Gene	Targeted therapies
NSCLC	<i>BRAF</i>	TAFINLAR® (dabrafenib) in combination with MEKINIST® (trametinib)
	<i>EGFR</i> L858R and exon 19 deletions	IRESSA® (gefitinib)
	<i>EGFR</i> exon 20 insertions	EXKIVITY™ (mobocertinib)
	<i>RET</i>	GAVRETO™ (pralsetinib)
	<i>ROS1</i>	XALKORI® (crizotinib)
CC	<i>IDH1</i>	TIBSOVO® (ivosidenib)

Figure 1. List of genes for therapeutic use.



- **Identify patients for multiple therapies**—one test indicated as a companion diagnostic (CDx) device to aid in selecting NSCLC and CC patients for treatment with targeted therapies
- **Multiple biomarkers from one limited sample**—one test for detection of 23 genes, minimizing the risk of depleting tissues and requiring additional biopsies. Based on Ion AmpliSeq™ technology, the required input is as low as 10 ng DNA and RNA.
- **One workflow, helps save time**—laboratory results can be generated within 4 days
- **Established performance**—Concordance with FDA approved or validated reference methods based on FISH, PCR, Sanger, or NGS was established for all CDx biomarkers:
 - 100% overall percent agreement (OPA), positive percent agreement (PPA) and negative percent agreement (NPA) for *BRAF*, *EGFR* exon 20 insertions, and *ROS1* fusions
 - 99% OPA, PPA, and NPA for *EGFR* exon 19 deletions and L858R
 - 92% OPA, 91% PPA, and 92% NPA for *RET* fusions
 - 98% OPA, 99% PPA, and 97% NPA for *IDH1*

Technical and validation data for the Oncomine Dx Target Test

Oncomine Dx Target Test content

Gene targets included for NSCLC				
Gene targets for therapeutic use				
<i>BRAF</i> : V600E	<i>EGFR</i> : L858R, exon 19 deletions, and exon 20 insertions	<i>ROS1</i> : fusions	<i>RET</i> : fusions	
Analytically validated targets				
<i>KRAS</i>	<i>MET</i> *	<i>PK3CA</i>		
Additional targets**				
<i>AK1</i>	<i>ERBB2</i>	<i>HRAS</i>	<i>MTOR</i>	<i>RET</i>
<i>ALK</i> *	<i>ERBB3</i>	<i>KIT</i>	<i>NRAS</i>	<i>ROS1</i>
<i>CDK4</i>	<i>FGFR2</i>	<i>MAP2K1</i>	<i>PDGFRA</i>	
<i>DDR2</i>	<i>FGFR3</i>	<i>MAP2K2</i>	<i>RAF1</i>	

Figure 2. Complete gene list. *The test reports fusion/translocation variants for *ROS1* and *RET* only. The test only reports *ALK* and *MET* mutations. **Performance for the additional gene target variants has been validated based on a representative method. Only *IDH1* is reported for CC.

Give Feedback

The power of next generation sequencing

Next-generation sequencing (NGS) can sequence hundreds to thousands of genes and detect multiple biomarkers at the same time. The sequencing takes place in a chip that contains millions of wells (flow cells) with separate sequencing reactions taking place in each well, allowing many genes to be sequenced at once and multiple variations to be detected simultaneously, unlike traditional companion diagnostic technologies such as FISH, IHC, or PCR, which only analyze one target gene at the time.

Learn how Ion Torrent NGS technology works in short video

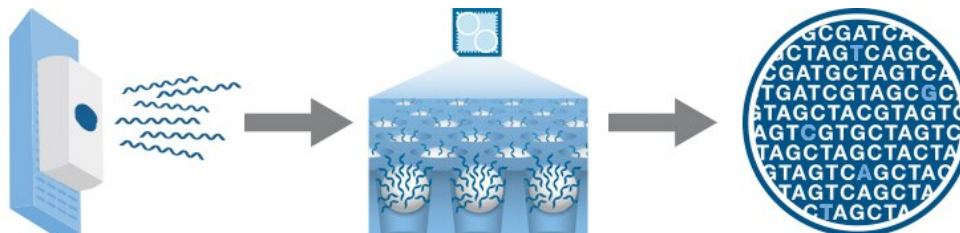


Figure 3. The NGS process starts with extraction of the DNA and/or RNA, which is processed in the chip in the Ion PGM Dx instrument, and results are analyzed and reported by a dedicated bioinformatics solution.

Oncomine Dx Target Test report

The Oncomine Dx Target Test Clinical Test Report is automatically generated as a PDF and incorporates relevant patient, sample, and test information required to help ensure high-performance standards, regulatory compliance, and quality control. The test results are presented in two-parts: companion diagnostic marker results with associated therapy indications and cancer driver analytical-only biomarker results in a separate section. The report is customizable and LIMS system-compatible.

NSCLC results for sequence variations for therapeutic use (for illustrative purposes only: *EGFR*, *BRAF*, *ROS1*, and *RET* are mutually exclusive)

DNA Sequence Variants

Gene	Display name	Amino acid change	Nucleotide change	Test result	Hotspot ID	Associated therapy
<i>EGFR</i>	<i>EGFR</i> L858R	p.Leu858Arg	c.2573T>G	POSITIVE	COSM6224	IRESSA® (gefitinib)
<i>EGFR</i>	<i>EGFR</i> exon 20 insertions	p.Ala767_Ser768 insSerValAsp	c.2311_2312ins GCGTGGACA	POSITIVE	COSM13428	EXKIVITY™ (mobocertinib)
<i>BRAF</i>	<i>BRAF</i> V600E	p.Val600Glu	c.1799T>A	POSITIVE	COSM476	TAFINLAR® + MEKINIST® (dabrafenib in combination with trametinib)

Gene Fusions (RNA)

Gene	Display name	Test result	Associated therapy
<i>ROS1</i>	<i>ROS1</i> Fusion	POSITIVE	XALKORI® (crizotinib)
<i>RET</i>	<i>RET</i> Fusions	POSITIVE	GAVRETO™ (pralsetinib)

CC results for sequence variations for therapeutic use (for illustrative purposes only)

DNA Sequence Variants

Gene	Display name	Amino acid change	Nucleotide change	Test result	Hotspot ID	Associated therapy
<i>IDH1</i>	<i>IDH1</i> R132	p.Arg132Gly	c.394C>G	POSITIVE	COSM28749	TIBSOVO® (ivosidenib)

Figure 4. An example of the OncoPrint Dx Target Test report format. The report includes a section with results of the validated biomarkers and information about relevant treatment indication, as well as a section with the other biomarkers not validated for treatment selection (not shown).

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[Request a quote](#)

Abbreviated Intended Use: The OncoPrint Dx Target Test is a qualitative *in vitro* diagnostic test that uses targeted high-throughput, parallel-sequencing technology to detect single-nucleotide variants (SNVs), deletions, and insertions in 23 genes from DNA and fusions in *ROS1* and *RET* from RNA isolated from formalin-fixed, paraffin-embedded (FFPE) tumor samples from patients with non-small cell lung cancer (NSCLC) and *IDH1* R132 mutations from FFPE tumor tissue samples from patients with cholangiocarcinoma (CC) using the Ion PGM Dx System.

Test limitations and warnings

- Use of this product must be limited to personnel trained in the techniques of PCR, NGS, and the use of the OncoPrint Dx Target Test and the Ion PGM Dx System.
- The OncoPrint Dx Target Test has only been validated for use with NSCLC and CC FFPE tumor slide specimens.
- The OncoPrint Dx Target Test has been validated to detect the following somatic mutations: RNA fusions, single-nucleotide variations (SNVs), multi-nucleotide variations (MNVs), and deletions of 3, 6, 9, 12, 15, and 18 base pairs (bps), and insertions of 3, 6, 9, and 12 base pairs (bps) from DNA.
- The OncoPrint Dx Target Test is only validated for use with the Ion PGM Dx System and the Veriti Dx 96-Well Thermal Cycler.
- The OncoPrint Dx Target Test is only validated for use with 10 ng each of DNA and RNA per sample. Input amounts lower or higher than 10 ng are not recommended.
- Both the DNA and RNA from a single sample extraction must meet the concentration requirements specified in the procedure. Do not use DNA from one extraction with RNA from a different extraction.
- The effects of potential variations in FFPE specimen fixation have not been evaluated.
- Extraction from FFPE sample curls has not been evaluated.
- A potential source of contamination in the procedure is nucleic acid from previous sample processing steps. Follow good laboratory practices and all precautions and guidelines in these user guides to avoid cross-contamination between samples.
- The OncoPrint Dx Target Test is a qualitative test. The test is not for quantitative measurements of percent mutation.
- The Ion OneTouch Rack Kit has only been designed to work with GeneMate SnapStrip 8-Strip 0.2 mL PCR Tubes. Tubes from other manufacturers may not fit properly in the rack, resulting in a higher risk of user error.
- For NSCLC, the OncoPrint Dx Target Test assay definition file includes prevalent but not all rare or newly identified *RET* isoforms, *ROS1* isoforms, and *EGFR* exon 20 insertions. The OncoPrint Dx Target Test may miss rare or newly identified:
 - *RET* isoforms carried by a subset of patients who may derive benefit from GAVRETO™ (pralsetinib)
 - *ROS1* isoforms carried by a subset of patients who may derive benefit from XALKORI® (crizotinib)
 - *EGFR* exon 20 insertions carried by a subset of patients who may derive benefit from EXKIVITY™ (mobocertinib)

For *In Vitro* Diagnostic Use.



Oncomine Dx Target Test Technical and Validation Information



Established performance

The Ion Torrent Oncomine Dx Target Test is the first targeted next-generation sequencing (NGS) *in vitro* diagnostic test simultaneously delivering multiple biomarker results to aid selection of targeted therapies for NSCLC and CC patients. Concordance with FDA approved or validated reference methods based on FISH, PCR, Sanger, or NGS was established for *EGFR*, *BRAF*, *ROS1*, and *RET* in NSCLC, and for *IDH1* in CC. The variants for *KRAS*, *MET*, and *PIK3CA* were analytically validated in NSCLC. The safety and effectiveness of these three genes have not been established and they are not intended to be used to direct therapy. The performance of all other variants identified by the test, other than clinically validated therapeutic variants and analytically validated variants, was validated based on a representative method.

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Oncomine Dx Target Test content

Gene targets included for NSCLC

Gene targets for therapeutic use				
<i>BRAF</i> : V600E	<i>EGFR</i> : L858R, exon 19 deletions, and exon 20 insertions	<i>ROS1</i> : fusions	<i>RET</i> : fusions	
Analytically validated targets				
<i>KRAS</i>	<i>MET</i> *	<i>PIK3CA</i>		
Additional targets**				
<i>AKT1</i>	<i>ERBB2</i>	<i>HRAS</i>	<i>MTOR</i>	<i>RET</i>
<i>ALK</i> *	<i>ERBB3</i>	<i>KIT</i>	<i>NRAS</i>	<i>ROS1</i>
<i>CDK4</i>	<i>FGFR2</i>	<i>MAP2K1</i>	<i>PDGFRA</i>	
<i>DDR2</i>	<i>FGFR3</i>	<i>MAP2K2</i>	<i>RAF1</i>	

Figure 1. Complete gene list. *The test reports fusion/translocation variants for *ROS1* and *RET* only. The test only reports *ALK* and *MET* mutations. ** Performance for the additional gene target variants has been validated based on a representative method. Only *IDH1* is reported for CC

Clinical concordance for companion diagnostics markers for targeted therapies selection - NSCLC

Method comparison studies evaluated the accuracy of the Oncomine Dx Target Test for the detection of *BRAF* V600E, *EGFR* exon 19 deletions, L858R, and exon 20 insertions, *ROS1* fusions, and *RET* fusions, using a *BRAF* V600E PCR assay, the Therascreen™ *EGFR* PCR Kit, *ROS1* FISH assay, and NGS assays respectively. A summary of the concordance studies results are included in Table 1. For details see the User Manual.

Variants for therapy selection	Validated comparator methods	Excluding no calls or unknowns*			Including no calls or unknowns*		
		Positive percent agreement	Negative percent agreement	Overall percent agreement	Positive percent agreement	Negative percent agreement	Overall percent agreement
<i>BRAF</i> V600E	Validated <i>BRAF</i> V600E qPCR test	100% (67/67)	100% (114/114)	100% (181/181)	91.8% (67/73)	97.4% (114/117)	95.3% (181/190)
<i>EGFR</i>	Therascreen™ <i>EGFR</i> PCR Kit	98.6% (71/72)	99.2% (120/121)	99.0% (191/193)	81.6% (71/87)	96.8% (120/124)	90.5% (191/211)
<i>EGFR</i> exon 19 deletions		97.6% (41/42)	99.3% (147/148)	99.0% (188/190)	74.6% (41/55)	94.2% (147/156)	89.1% (188/211)
<i>EGFR</i> exon 21 L858R		100% (30/30)	100% (167/167)	100% (197/197)	93.8% (30/32)	93.3% (167/179)	93.4% (197/211)
<i>EGFR</i> exon 20 insertions	Validated NGS Assay	100% (54/54)	100% (95/95)	100% (149/149)	98.2% (54/55)	90.5% (95/105)	93.1% (149/160)
<i>ROS1</i> fusions	Validated <i>ROS1</i> FISH test	100% (9/9)	100% (62/62)	100% (71/71)	90.0% (9/10)	88.6% (62/70)	88.8% (71/80)
<i>RET</i> fusions	Validated NGS Assay	90.9% (40/44)	91.8% (101/110)	91.6% (141/154)	90.9% (40/44)	91.8% (101/110)	91.6% (141/154)

* No-calls are for DNA variants and unknowns are for RNA fusions

Table 1. Method comparison between OncoPrint Dx Target Test and reference methods for five companion diagnostic biomarkers in NSCLC.

Analytical validation performance - NSCLC

The OncoPrint Dx Target Test also detects DNA sequence variations in an additional 19 genes in NSCLC. The variants for *KRAS*, *MET*, and *PIK3CA* have been analytically validated. Safety and effectiveness of these three genes have not been established and they are not intended to be used to direct therapy. The performance of all other variants identified by the test, other than clinically validated therapeutic variants and analytically validated variants, has not been directly demonstrated and was validated based on a representative method.

Limit of detection

Four LoD studies were performed to evaluate DNA variants, *ROS1* fusions, *RET* fusions, and *EGFR* exon 20 insertions.

Study I: The limit of detection (LoD) was evaluated for 14 representative DNA variants representing 3 variant categories detected by the OncoPrint Dx Target Test. The LoD is the lowest allele frequency of SNV, multi-nucleotide polymorphism (MNP), or deletion variants, that can be detected at least 95% of the time. The study demonstrated that the OncoPrint Dx Target Test can detect DNA variants with allele frequencies between 6 and 8%.

Study II: The LoD was calculated for 2 clinical *ROS1* RNA fusion variants using the updated RNA library preparation workflow, and determined at 516 fusion reads.

Study III: The LoD was calculated for 2 clinical *RET* fusion variants using the updated RNA library preparation workflow, and determined at 405 fusion reads.

Study IV: The LoD was calculated for 2 clinical *EGFR* exon 20 insertion positive samples, and determined to be 4.8–5.2% allele frequencies.

Accuracy

To evaluate the ability of the OncoPrint Dx Target Test DNA and RNA panels to identify somatic variants in human specimens, 290 FFPE tumor samples were analyzed using the OncoPrint Dx Target Test to demonstrate positive percent agreement (PPA) and negative percent agreement (NPA) concordance with validated reference detection methods.

The following reference detection methods were used:

- Validated NGS method, to detect SNV and deletion hotspot variants
- Validated *ROS1* FISH test, to detect *ROS1* fusions

The study demonstrated variant level PPA of 98.5%, NPA of 100%, and OPA of 100%, excluding invalids no-calls; and PPA level of 98.5%, NPA of 96.8%, and OPA of 96.8% including no-calls. A summary of the data are included in Table 2. For details see the User Manual.

Variant level measure of agreement	Percent agreement (N) excluding no-calls	Percent agreement (N) including no-calls
Positive percent agreement	98.5% (195/198)	98.5% (195/198)
Negative percent agreement	100.0% (118,155/118,159)	96.8% (118,155/122,012)
Overall percent agreement	100.0% (118,350/118,357)	96.8% (118,350/122,210)

Table 2. Variant level accuracy study results

Reproducibility

Four reproducibility studies were performed to evaluate DNA variants, *ROS1* fusions, *RET* fusions, and *EGFR* exon 20 insertions.

Study I:

The reproducibility and repeatability of the OncoPrint Dx Target Test was evaluated for 30 representative variants from 18 DNA samples. The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instruments (reproducibility). Due to the large number of variants detected by the test and the rarity of some of the variants, a representative variant approach was used. Variants were selected in the following categories:

- Simple SNVs
- Complex SNVs and MNPs, including SNVs in di- or tri-nucleotide repeat regions and SNVs in high-GC (>60%) or low-GC (<40%) content regions
- Deletions (including deletions of 6, 9, 15, and 18 bp)

Excluding no calls, the percent of correct calls is >96%. The estimate of repeatability at each DNA variant location across all the samples was ≥98.8% (95% CI lower limit of ≥97.5%). A summary of results of the assay reproducibility study are included in Table 3. For details see the User Manual.

Description	No. of variants	Call rate excluding no-calls		Call rate including no-calls	
		Mean	Median	Mean	Median
DNA positive variants (positive calls)	46	96.60%	97.10%	94.50%	95.80%
WT DNA variant locations (negative calls)	872	96.10%	95.00%	96.10%	95.00%

Table 3. Assay reproducibility study I.

Study II:

An additional study was performed to evaluate the reproducibility and repeatability of the OncoPrint Dx Target Test for 6 representative variants from 11 DNA samples and 4 RNA samples. 1 WT DNA sample and 4 WT RNA samples were included in the study.

The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instrument platforms (reproducibility). The updated RNA library preparation workflow was used. Due to the large number of variants detected by the test and the rarity of some variants, a representative variant approach was used.

Variants were selected in the following categories:

- 15 bp deletion
- Simple SNVs
- Complex SNVs and MNPs
- Fusions

Excluding no calls, the estimate of repeatability at each DNA variant location across all the samples was $\geq 94.4\%$ (95% CI lower limit of $\geq 72.7\%$). The estimate of repeatability at each RNA clinical variant location was 100%. A summary of the results of Study II is included in Table 4 and 5.

Description	No. of variants	Call rate excluding no-calls		Call rate including no-calls	
		Mean	Median	Mean	Median
DNA positive variants (positive calls)	11	99%	100%	98%	99%
WT DNA variant locations (negative calls)	367	100%	100%	99%	100%

Table 4. Study II—assay reproducibility study results (DNA variants)

Description	No. of variants	Call rate including or excluding unknowns	
		Mean	Median
<i>ROS1</i> positive variants (positive calls)	4	100%	100%
WT RNA variant locations (negative calls)	4	99%	100%

Table 5. Study II—assay reproducibility study results (*ROS1* fusions)

Study III:

An additional study was performed to evaluate the reproducibility and repeatability of the OncoPrint Dx Target Test for 4 *RET* fusion positive samples and 2 *RET* fusion-negative samples. The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instrument platforms (reproducibility). The updated RNA library preparation workflow was used. Excluding unknowns, estimates of the repeatability ranged from 98.1% to 100% for two *RET* variants. A summary of the results of Study III is included in Table 6.

Description	No. of variants	Call rate including or excluding unknowns	
		Mean	Median
<i>RET</i> positive variants (positive calls)	4	99%	100%
WT RNA variant locations (negative calls)	2	100%	100%

Table 6. Study III—assay reproducibility study results (*RET* fusions)

Study IV:

A study was performed to evaluate the reproducibility and repeatability of the OncoPrint™ Dx Target Test for detection of *EGFR* exon 20 insertion variants using FFPE DNA from 2 *EGFR* variant-positive samples (blended with WT clinical samples) and 2 *EGFR* variant-negative (WT) samples.

The study was designed to evaluate within-run precision performance (repeatability) and variability across sites, operators, and instrument platforms (reproducibility). Excluding no-calls, estimates of the repeatability is 100% for both *EGFR* exon 20 insertion variants. A summary of the results of Study IV is included in Table 7.

Description	No. of variants	Call rate excluding no-calls		Call rate including no-calls	
		Mean	Median	Mean	Median
<i>EGFR</i> insertion positive variants (positive calls)	2	100%	100%	100%	100%
WT DNA variants (negative calls)	2	100%	100%	100%	100%

Table 7. Study IV—assay reproducibility study results (*EGFR* exon 20 insertions)

Clinical concordance for companion diagnostics marker for targeted therapies selection - CC

A clinical concordance study was conducted to evaluate the ability of the Oncomine Dx Target Test to identify five *IDH1* biomarkers in FFPE cholangiocarcinoma tumor specimen compared to a validated Sanger assay. The study demonstrated OPA of 97.9%, excluding invalids and no calls. A summary of the data is included in Table 8.

Table 8. Concordance between the Oncomine Dx Target Test and reference method for *IDH1* R132

Variants for therapy selection	Validated comparator methods	Excluding invalid results and no-calls			Including invalid results and no-calls		
		Positive percent agreement	Negative percent agreement	Overall percent agreement	Positive percent agreement	Negative percent agreement	Overall percent agreement
<i>IDH1</i> R132	Validated Sanger assay	99.4% (163/164)	96.5% (164/170)	97.9% (327/334)	97.0% (163/168)	90.6% (164/181)	93.7% (327/349)

Analytical validation performance - CC

Limit of detection

The limit of detection (LoD) was evaluated for 5 *IDH1* R132 variants detected by the Oncomine Dx Target Test. The LoD is the lowest allele frequency of SNV that can be detected at least 95% of the time. The study demonstrated LoD of the 5 *IDH1* R132 variants ranged from 4.5-5.7% allele frequencies, including 4.5% for R132C, 5.7% for R132G, 4.9% for R132H, 5.1% for R132L, and 5.3% for R132S.

Assay reproducibility

The reproducibility and repeatability of *IDH1* R132 variant detection using Oncomine Dx Target Test were assessed with 1 *IDH1* WT sample and 3 *IDH1* R132 variant positive samples at two allele frequency (AF) levels. Testing was performed at 4 testing sites, each site had 2 PGM Dx instrument systems, 2 operators, and using 4 lots of reagents. The overall positive call rate for *IDH1* R132 variants was 92.6% when including no calls and 97.1% when excluding no calls. The negative call rate for *IDH1* WT sample were 100% at all *IDH1* R132 variant locations. (Table 9).

Table 9. Reproducibility results

Sample COSMIC ID, Variant	No. of valid sample results	Call rate (95% CI)	
		Including no calls	Excluding no calls
D1 COSM28747, R132C	36	100% (90.3%, 100%)	100% (90.3%, 100%)
D2 COSM28747, R132C	36	97.2% (85.5%, 99.9%)	100% (90.0%, 100%)
D3 COSM28749, R132G	36	100% (90.3%, 100%)	100% (90.3%, 100%)
D4 COSM28749, R132G	36	100% (90.3%, 100%)	100% (90.3%, 100%)
D5 COSM28750, R132L	36	100% (90.3%, 100%)	100% (90.3%, 100%)
D6 COSM28750, R132L	35	57.1% (39.4%, 73.7%)	76.9% (56.4%, 91.0%)
D1-D6 All Variants, R132	215	92.6% (88.2%, 95.7%)	97.1% (93.7%, 98.9%)
D7 Wild-Type	36	100% (90.3%, 100%)	100% (90.3%, 100%)

A complete and flexible system

The Oncomine Dx Target Test is used in conjunction with the Ion PGM Dx System, which includes a complete NGS system of instruments, reagents, and software. The Ion PGM Dx System was initially validated using challenging germline variants and is now additionally validated with the Oncomine Dx Target Test for somatic mutation reporting for FFPE tissue samples. The Ion PGM Dx sequencing system is a Class II 510 K Medical Device and incorporates combined functionality, with both "IVD Mode" for molecular diagnostic tests and "Assay Development Mode" for clinical research. The system also facilitates 21CFR Part 11 compliance, role-based workflows, sample and reagent tracking, QC metrics, and audit trails.

The Oncomine Dx Target Test workflow—all results in 4 days

The Oncomine Dx Target Test workflow is a fully validated IVD workflow from beginning to end and includes all the reagents, consumables, instruments, and software to perform the test. It is possible to run 1–6 samples per run, plus 2 controls within 4 days (Figure 2).



Figure 2. The Oncomine Dx Target Test utilizes a single streamlined NGS workflow for detecting cancer-associated biomarkers, incorporating reagents, instrument systems, and bioinformatics. The turnaround time, from FFPE sample to report, is 4 days.

Request more information

Abbreviated Intended Use: The Oncomine Dx Target Test is a qualitative *in vitro* diagnostic test that uses targeted high-throughput, parallel-sequencing technology to detect single-nucleotide variants (SNVs), deletions, and insertions in 23 genes from DNA and fusions in *ROS1* and *RET* from RNA isolated from formalin-fixed, paraffin-embedded (FFPE) tumor samples from patients with non-small cell lung cancer (NSCLC) and *IDH1* R132 mutations from FFPE tumor tissue samples from patients with cholangiocarcinoma (CC) using the Ion PGM Dx System.

Test limitations and warnings

- Use of this product must be limited to personnel trained in the techniques of PCR, NGS, and the use of the Oncomine Dx Target Test and the Ion PGM Dx System. The
- Oncomine Dx Target Test has only been validated for use with NSCLC and CC FFPE tumor slide specimens.
- The Oncomine Dx Target Test has been validated to detect the following somatic mutations: RNA fusions, single-nucleotide variations (SNVs), multi-nucleotide variations (MNVs), deletions of 3, 6, 9, 12, 15, and 18 base pairs (bps), and insertions of 3, 6, 9, and 12 base pairs (bps) from DNA.
- The Oncomine Dx Target Test is only validated for use with the Ion PGM Dx System and the Veriti Dx 96-Well Thermal Cycler.
- The Oncomine Dx Target Test is only validated for use with 10 ng each of DNA and RNA per sample. Input amounts lower or higher than 10 ng are not recommended.
- Both the DNA and RNA from a single sample extraction must meet the concentration requirements specified in the procedure. Do not use DNA from one extraction with RNA from a different extraction.
- The effects of potential variations in FFPE specimen fixation have not been evaluated.
- Extraction from FFPE sample curls has not been evaluated.
- A potential source of contamination in the procedure is nucleic acid from previous sample processing steps. Follow good laboratory practices and all precautions and guidelines in these user guides to avoid cross-contamination between samples.
- The Oncomine Dx Target Test is a qualitative test. The test is not for quantitative measurements of percent mutation.
- The Ion OneTouch Rack Kit has only been designed to work with GeneMate SnapStrip 8-Strip 0.2 mL PCR Tubes. Tubes from other manufacturers may not fit properly in the rack, resulting in a higher risk of user error.
- For NSCLC, the Oncomine Dx Target Test assay definition file includes prevalent but not all rare or newly identified *RET* isoforms, *ROS1* isoforms, and *EGFR* exon 20 insertions. The Oncomine Dx Target Test may miss rare or newly identified:
 - *RET* isoforms carried by a subset of patients who may derive benefit from GAVRETO™ (pralsetinib)
 - *ROS1* isoforms carried by a subset of patients who may derive benefit from XALKORI® (crizotinib)
 - *EGFR* exon 20 insertions carried by a subset of patients who may derive benefit from EXKIVITY™ (mabocertinib)

For In Vitro Diagnostic Use.



United States

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