Blood Collection Instructions

Shipping Instructions

1. Collect specimen by venipuncture according to CLSI GP41¹.

Prevention of Backflow Since Cell-Free DNA BCT contains chemical additives, it is important to avoid possible backflow from the tube. To guard against backflow, observe the following precautions:

- a. Keep patient's arm in the downward position during the collection procedure
- b. Hold the tube with the stopper in the uppermost position so that the tube contents do not touch the stopper or the end of the needle.
- c. Release tourniquet once blood starts to flow in the tube, or within 2 minutes of application.
- 2. Follow recommendations for order of draw outlined in CLSI GP41¹. Cell-Free DNA BCT should be drawn after the EDTA tube and before the fluoride oxalate (glycolytic inhibitor) tube. If a Cell-Free DNA BCT immediately follows a heparin tube in the draw order, Streck recommends collecting a non-additive or EDTA tube as a waste tube prior to collection in

3. Fill tube completely. Remove tube from adapter and immediately mix by gentle inversion 10 times.

Inadequate, delayed, or over mixing may result in incorrect analytical results or poor product performance. One inversion is a complete turn of the wrist, 180 degrees, and back

the Cell-Free DNA BCT.

4. After collection, transport and store tubes within the recommended temperature range.

Note:

 For best results, a 21G or 22G needle is advised. Slower fill times may be observed when using a smaller gauge needle.

- When using a winged (butterfly) collection set for venipuncture and the Streck Cell-Free DNA BCT is the first tube drawn, a non-additive or EDTA discard tube should be partially drawn first in order to eliminate air or "dead space"
- from the tubing. 3. Cell-Free DNA BCT does not dilute blood samples;
- therefore, no dilution factor correction is necessary.

PLASMA ISOLATION

1. Centrifuge blood collection tubes containing whole blood for 10 minutes at 10 °C at 1,600 x g. Carefully unload the tubes

from the centrifuge without disturbing the buffy coat . 2. Pipette maximum 5ml plasma from the specimen tube to a labeled 15ml centrifuge tube. 3. Centrifuge plasma in 15ml centrifuge tube for 10 minutes at 10 °C at 3,220 x g. Note: Centrifuge speed may be adjusted down to 3,200 x g. if centrifuge does not support 3,220 x g.

Rotate

Up

Rotate

Down

 Pour contents of 15ml centrifuge tube for each specimen into an appropriately labeled 5ml conical screw cap tube.
 Isolate cell-free DNA using extraction kit manufacturer's instructions including any adaptations as required.

PLASMA Storage

Plasma may be stored at 2 °C to 8 °C for up to 24 hours or at -80 °C +/- 10 °C for up to 45 days. This has only been validated on the Guardant360 CDx assay. 1. Complete the Test Requisition Form (TRF)

- 2. Fill out barcode labels - Patient/
 - Subject ID - DOB - Collection Date
- **3.** Place a barcode label on each tube with barcode in the vertical position

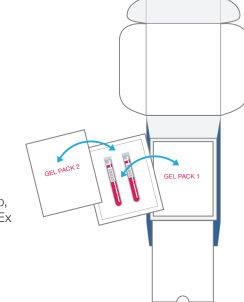
 Place a barcode label on the TRF, then fold and insert the TRF into the outer pocket of the biohazard bag



5. Place filled blood tubes into foam tray



- 7. Place filled biohazard bag into box on top of one of the gel packs.* Place the second gel pack on top of the specimen bag and close the box
- 8. Place the kit into the preprinted FedEx Clinical Pak, seal flap, and schedule a FedEx pick up



*DO NOT freeze Gel Packs. Use as is.

INTENDED USE

The Guardant360[®] CDx Blood Collection Kit is intended for whole blood collection for further processing and testing of cell-free plasma DNA with Guardant360 CDx.

Cell-Free DNA BCT is a direct-draw venous whole blood collection device intended for the collection, stabilization, and transport of venous whole blood samples for use in conjunction with cell-free DNA next- generation sequencing liquid biopsy assays that have been cleared or approved for use with samples collected in the Cell-Free DNA BCT device.

PRINCIPLES OF THE PROCEDURE

Accurate analysis of cell-free DNA (cfDNA) can be compromised by sample handling, shipping and processing, causing lysis of nucleated blood cells and subsequent release of cellular genomic DNA. The preservative reagent contained in Cell-Free DNA BCT reduces the release of cellular genomic DNA. The blood collection tube is intended for the preservation of cell-free DNA. BLOOD COLLECTION TUBE REAGENTS

Cell-Free DNA BCT contains the anticoagulant $K_{\rm 3}\text{EDTA}$ and a cell preservative in a liquid medium. LIMITATIONS

- Performance characteristics for this device have only been established on the Guardant360 CDx assay.
- Do not store outside of established conditions.
- Do not transfer samples drawn into tubes containing other anti-coagulants and/or preservatives into Cell-Free DNA BCT
- Do not use past expiration date printed on label.
- Do not use for clinical chemistry assays or assays not intended for cell-free DNA analysis.
- Do not use for collection of materials to be injected into patients.
- Cell-Free DNA BCT is not intended for the stabilization of RNA nor is it intended for viral or microbial nucleic acids.

PRECAUTIONS

- 1. Wear personal protective equipment (PPE) while handling specimens.
- 2. Do not freeze specimens collected in Cell-Free DNA BCT.
- 3. For single and professional use only.
- 4. Product is intended for use as supplied. Do not dilute or add other components to Cell-Free DNA BCT.
- 5. Specimen transport via pneumatic tube system is not advised.
- 6. Allow the tube to fill completely until blood stops flowing into the tube. Underfilling of tubes with less than 5ml of blood (bottom of the label indicates 5ml fill when tube is held vertically) may lead to incorrect analytical results or poor product performance. This tube has been designed to fill with 10ml of blood.

CAUTION

- Glass has the potential for breakage; precautionary measures should be taken during handling of glass tubes.
- All biological specimens and materials coming in contact with them are considered biohazards and should be treated as if capable of transmitting infection. Dispose of in accordance with federal, state and local regulations. Avoid contact with skin and mucous membranes.
 Product should be disposed with infectious medical waste.
- Remove and reinsert stopper by either gently rocking the stopper from side to side or by grasping with a simultaneous twisting and pulling action. A "thumb roll" procedure for stopper removal is NOT recommended as tube breakage and injury may result.
- 7. SDS(s) can be obtained by emailing sds@guardanthealth.com.

STORAGE AND STABILITY

- 1. Store tubes prior to blood draw at 2°C-30°C until expiration date printed on label
- If stored consistently at 22 °C to 30 °C for longer than 8 months, reagent will acquire a yellow hue. This is normal and does not indicate product degradation.
- 2. Do not freeze empty Cell-Free DNA BCT.
- 3. After draw, whole blood containing Cell-Free DNA BCT should be stored at 18 °C to 25 °C for up to 7 days including shipping.

INDICATIONS OF PRODUCT DETERIORATION

- 1. Cloudiness or precipitate visible in reagent of empty tube.
- If indications of product deterioration occur, contact Guardant Health by emailing sds@guardanthealth.com.

MATERIALS THAT MAY BE REQUIRED BUT ARE NOT PROVIDED

Needle, tube adapter, alcohol swab, bandage, gloves, gauze, tourniquet

REFERENCES

1. Clinical and Laboratory Standards Institute. GP41, Procedures for the collection of diagnostic blood specimens by venipuncture. Approved Standard - Seventh Edition.

GLOSSARY OF HARMONIZED SYMBOLS

STERIL R Sterilized using irradiation	LOT Batch Code	REF Catalog Number	Do Not Re-use	Use By	Biological Risk	Content Sufficient for Number of Tests Specified
EC REP Authorized Representative in the European Community	In Vitro Diagnostic Medical Device	Temperature Limitation	Consult Instructions For Use	Manufacturer	Date of Manufacture	Rx ONLY By Prescription Only

GUARDANT

MANUFACTURED FOR:

Guardant Health 505 Penobscot Drive Redwood City, CA 94063 USA T 1.855.698.8887 USA only | +1.650.290.7575 Worldwide F 1.888.974.4258 clientservices@guardanthealth.com

GH10005, GH10006 assembled in USA GH10000 assembled in Czech Republic





08/2020 D-000348 B1

Blood Collection Kit

FDA APPROVED

Catalog Numbers: GH10005 GH10006 GH10000

GUARDANT 360°CD×

Instructions for use



Physician Insert: Guardant360[®] CDx

For In Vitro Diagnostic Use

Genetic Companion Diagnostic Testing for Targeted Therapy Selection in Non-Small Cell Lung Cancer (NSCLC)

For the most current information on the association of the biomarker and therapeutic outcomes, refer to the therapeutic labels available at Drugs@FDA on the FDA website.

Guardant360 CDx Intended Use

Guardant360[®] CDx is a qualitative next generation sequencing-based *in vitro* diagnostic device that uses targeted high throughput hybridization-based capture technology for detection of single nucleotide variants (SNVs), insertions and deletions (indels) in 55 genes, copy number amplifications (CNAs) in two (2) genes, and fusions in four (4) genes. Guardant360 CDx utilizes circulating cell-free DNA (cfDNA) from plasma of peripheral whole blood collected in Streck Cell-Free DNA Blood Collection Tubes (BCTs). The test is intended to be used as a companion diagnostic to identify non-small cell lung cancer (NSCLC) patients who may benefit from treatment with the targeted therapy listed in **Table 1** in accordance with the approved therapeutic product labeling.

Table 1. Companion Diagnostic Indications

Indication	Biomarker	Therapy
Non-small cell lung cancer (NSCLC)	EGFR exon 19 deletions, L858R and T790M*	TAGRISSO [®] (osimertinib)

A negative result from a plasma specimen does not assure that the patient's tumor is negative for genomic findings. NSCLC patients who are negative for the biomarkers listed in **Table 1** should be reflexed to tissue biopsy testing for **Table 1** biomarkers using an FDA-approved tumor tissue test, if feasible.

*The efficacy of TAGRISSO[®] (osimertinib) has not been established in the *EGFR* T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.

Additionally, the test is intended to provide tumor mutation profiling to be used by qualified health care professionals in accordance with professional guidelines in oncology for cancer patients with any solid malignant neoplasm. The test is for use with patients previously diagnosed with cancer and in conjunction with other laboratory and clinical findings.

Genomic findings other than those listed in **Table 1** are not prescriptive or conclusive for labeled use of any specific therapeutic product.

Guardant360 CDx is a single-site assay performed at Guardant Health, Inc.

Warnings and Precautions

- Alterations reported may include somatic (not inherited) or germline (inherited) alterations. The assay filters germline variants from reporting except for pathogenic *BRCA1*, *BRCA2*, *ATM*, and *CDK12* alterations. However, if a reported alteration is suspected to be germline, confirmatory testing should be considered in the appropriate clinical context.
- The test is not intended to replace germline testing or to provide information about cancer predisposition.
- Somatic alterations in *ATM* and *CDK12* are not reported by the test as they are excluded from the test's reportable range.
- Genomic findings from cfDNA may originate from circulating tumor DNA (ctDNA) fragments, germline alterations, or non-tumor somatic alterations, such as clonal hematopoiesis of indeterminate potential (CHIP).
- Allow the tube to fill completely until blood stops flowing into the tube. Underfilling of tubes with less than 5 mL of blood (bottom of the label indicates 5 mL fill when tube is held vertically) may lead to incorrect analytical results or poor product performance. This tube has been designed to fill with 10 mL of blood.

Test Limitations

- For *in vitro* diagnostic use.
- For prescription use only. This test must be ordered by a qualified medical professional in accordance with clinical laboratory regulations.
- The efficacy of TAGRISSO® (osimertinib) has not been established in the *EGFR* T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.
- TAGRISSO efficacy has not been established in patients with *EGFR* exon 19 deletions < 0.08% MAF, in patients with *EGFR* L858R < 0.09% MAF, and in patients with *EGFR* T790M < 0.03% MAF.
- The test is not intended to be used for standalone diagnostic purposes.
- The test is intended to be performed on specific serial number-controlled instruments by Guardant Health, Inc.

- A negative result for any given variant does not preclude the presence of this variant in tumor tissue.
- Decisions on patient care and treatment must be based on the independent medical judgment of the treating physician, taking into consideration all applicable information concerning the patient's condition, such as patient and family history, physical examinations, information from other diagnostic tests, and patient preferences, in accordance with the standard of care.
- ctDNA shedding rate may be lower in patients with primary central nervous system (CNS) tumors.

Explanation of the Tiered Reporting

Genomic findings other than those listed in **Table 1** are not prescriptive or conclusive for labeled use of any specific therapeutic product. Test results should be interpreted in the context of pathological evaluation of tumors, treatment history, clinical findings, and other laboratory data.

The test report includes genomic finding reported in the following categories (Table 2).

	(Guardant360 CD	X	
Category	Prescriptive use for a Therapeutic Product	Clinical Performance	Analytical Performance	Comments
<u>Category 1:</u> Companion Diagnostic (CDx)	Yes	Yes	Yes	ctDNA biomarkers linked to the safe and effective use of the corresponding therapeutic product, for which Guardant360 CDx has demonstrated clinical performance shown to support therapeutic efficacy and strong analytical performance for the biomarker.
<u>Category 2:</u> ctDNA Biomarkers with Strong Evidence of Clinical Significance in ctDNA	No	No	Yes	ctDNA biomarkers with strong evidence of clinical significance presented by other FDA-approved liquid biopsy companion diagnostics for which Guardant360 CDx has demonstrated analytical reliability but not clinical performance.

Table 2. Category Definitions

Category 3A: Biomarkers with Evidence of Clinical Significance in tissue supported by: strong analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated analytical performance including analytical accuracy, and concordance of blood- based testing to tissue-based testing for the biomarker.
Category 3B: Biomarkers with Evidence of Clinical Significance in tissue supported by: analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated minimum analytical performance including analytical accuracy.
Category 4: Other Biomarkers with Potential Clinical Significance	No	No	Yes	ctDNA biomarkers with emergent evidence based on peer- reviewed publications for genes/variants in tissue, variant information from well-curated public databases, or <i>in-vitro</i> preclinical models, for which Guardant360 CDx has demonstrated minimum analytical performance.

Guardant360[®] CDx

Technical Information

Guardant Health, Inc. 505 Penobscot Dr. Redwood City, CA 94063 USA

1 Intended Use

Guardant360® CDx is a qualitative next generation sequencing-based *in vitro* diagnostic device that uses targeted high throughput hybridization-based capture technology for detection of single nucleotide variants (SNVs), insertions and deletions (indels) in 55 genes, copy number amplifications (CNAs) in two (2) genes, and fusions in four (4) genes. Guardant360 CDx utilizes circulating cell-free DNA (cfDNA) from plasma of peripheral whole blood collected in Streck Cell-Free DNA Blood Collection Tubes (BCTs). The test is intended to be used as a companion diagnostic to identify non-small cell lung cancer (NSCLC) patients who may benefit from treatment with the targeted therapy listed in **Table 1** in accordance with the approved therapeutic product labeling.

Table 1. Companion Diagnostic Indications

Indication	Biomarker	Therapy
Non-small cell lung cancer (NSCLC)	<i>EGFR</i> exon 19 deletions, L858R and T790M*	TAGRISSO [®] (osimertinib)

A negative result from a plasma specimen does not assure that the patient's tumor is negative for genomic findings. NSCLC patients who are negative for the biomarkers listed in **Table 1** should be reflexed to tissue biopsy testing for **Table 1** biomarkers using an FDA-approved tumor tissue test, if feasible.

*The efficacy of TAGRISSO[®] (osimertinib) has not been established in the *EGFR* T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.

Additionally, the test is intended to provide tumor mutation profiling to be used by qualified health care professionals in accordance with professional guidelines in oncology for cancer patients with any solid malignant neoplasm. The test is for use with patients previously diagnosed with cancer and in conjunction with other laboratory and clinical findings.

Genomic findings other than those listed in **Table 1** are not prescriptive or conclusive for labeled use of any specific therapeutic product.

Guardant360 CDx is a single-site assay performed at Guardant Health, Inc.

2 Contraindications

There are no known contraindications.

3 Warnings and Precautions

- Alterations reported may include somatic (not inherited) or germline (inherited) alterations. The assay filters germline variants from reporting except for pathogenic *BRCA1, BRCA2, ATM*, and *CDK12* alterations. However, if a reported alteration is suspected to be germline, confirmatory testing should be considered in the appropriate clinical context.
- The test is not intended to replace germline testing or to provide information about cancer predisposition.
- Somatic alterations in *ATM* and *CDK12* are not reported by the test as they are excluded from the test's reportable range.
- Genomic findings from cfDNA may originate from circulating tumor DNA (ctDNA) fragments, germline alterations, or non-tumor somatic alterations, such as clonal hematopoiesis of indeterminate potential (CHIP).
- Allow the tube to fill completely until blood stops flowing into the tube. Underfilling of tubes with less than 5 mL of blood (bottom of the label indicates 5 mL fill when tube is held vertically) may lead to incorrect analytical results or poor product performance. This tube has been designed to fill with 10 mL of blood.

4 Limitations

- For *in vitro* diagnostic use.
- For prescription use only. This test must be ordered by a qualified medical professional in accordance with clinical laboratory regulations.
- The efficacy of TAGRISSO[®] (osimertinib) has not been established in the *EGFR* T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.
- TAGRISSO efficacy has not been established in patients with *EGFR* exon 19 deletions < 0.08% MAF, in patients with *EGFR* L858R <0.09% MAF, and in patients with *EGFR* T790M < 0.03% MAF.
- The test is not intended to be used for standalone diagnostic purposes.
- The test is intended to be performed on specific serial number-controlled instruments by Guardant Health, Inc.
- A negative result for any given variant does not preclude the presence of this variant in tumor tissue.

- Decisions on patient care and treatment must be based on the independent medical judgment of the treating physician, taking into consideration all applicable information concerning the patient's condition, such as patient and family history, physical examinations, information from other diagnostic tests, and patient preferences, in accordance with the standard of care.
- ctDNA shedding rate may be lower in patients with primary central nervous system (CNS) tumors.

5 Guardant360 CDx Overview

5.1 Test Summary and Explanation

Guardant360 CDx is a next generation sequencing-based test for the detection of genetic alterations in 55 genes frequently mutated in cancer. It is a companion diagnostic to identify non-small cell lung cancer (NSCLC) patients who may benefit from treatment with the targeted therapy listed in **Table 1** of the Intended Use. Additionally, the test is intended to provide tumor mutation profiling to be used by qualified health care professionals in accordance with professional guidelines in oncology for cancer patients with any solid malignant neoplasm.

The test report includes variants reported in the following categories (**Table 2**).

8	Gu	ardant360 CDx		
Category	Prescriptive use for a Therapeutic Product	Clinical Performance	Analytical Performance	Comments
<u>Category 1:</u> Companion Diagnostic (CDx)	Yes	Yes	Yes	ctDNA biomarkers linked to the safe and effective use of the corresponding therapeutic product, for which Guardant360 CDx has demonstrated clinical performance shown to support therapeutic efficacy and strong analytical performance for the biomarker.
<u>Category 2:</u> ctDNA Biomarkers with Strong Evidence of Clinical Significance in ctDNA	No	No	Yes	ctDNA biomarkers with strong evidence of clinical significance presented by other FDA-approved liquid biopsy companion diagnostics for which Guardant360 CDx has demonstrated analytical reliability but not clinical performance.

<u>Category 3A:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: strong analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated analytical performance including analytical accuracy, and concordance of blood-based testing to tissue-based testing for the biomarker.
<u>Category 3B:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated minimum analytical performance including analytical accuracy.
<u>Category 4:</u> Other Biomarkers with Potential Clinical Significance	No	No	Yes	ctDNA biomarkers with emergent evidence based on peer-reviewed publications for genes/variants in tissue, variant information from well- curated public databases, or <i>in-vitro</i> pre-clinical models, for which Guardant360 CDx has demonstrated minimum analytical performance.

5.2 Sample Collection and Test Ordering

To order Guardant360 CDx, the Test Requisition Form (TRF) provided with the Guardant360 CDx Blood Collection Kit must be fully completed and signed by the ordering physician or other authorized medical professional. Refer to the Guardant360 CDx Blood Collection Kit Instructions for Use for further details about collecting blood samples and shipping samples to the Guardant Health Clinical Laboratory.

To order the Guardant360 CDx Blood Collection Kit or obtain an electronic version of the TRF, contact the Guardant Health Client Services department (Tel: 855.698.8887, Fax: 888.974.4258, or Email: <u>clientservices@guardanthealth.com</u>).

5.3 Principles of the Procedure

Guardant360 CDx is performed by a single laboratory, the Guardant Health Clinical Laboratory, located in Redwood City, CA, USA. Guardant360 CDx is composed of the following major processes:

• Whole Blood Collection and Shipping

- Plasma Isolation and cfDNA Extraction
- Library Preparation and Enrichment
- DNA Sequencing
- Data Analysis and Reporting

The Guardant360 CDx Blood Collection Kit is used by the ordering laboratories / physicians to collect whole blood specimens and ship them to the Guardant Health Clinical Laboratory. Whole blood is collected in the provided blood collection tubes, Streck Cell-Free DNA BCTs, which stabilize cfDNA and nucleated blood cells for shipping.

All other reagents, materials and equipment needed to perform the assay are used exclusively in the Guardant Health Clinical Laboratory.

Whole blood specimens are processed in the Guardant Health Clinical Laboratory within 7 days of blood collection. A minimum of 5 mL whole blood must be received in order to achieve optimal performance for the Guardant360 CDx assay. Underfilling of tubes with less than 5 mL of blood may lead to incorrect analytical results or poor product performance. Plasma is isolated via centrifugation and cfDNA is extracted from plasma. cfDNA, 5 to30 ng, is then used to prepare sequencing libraries which are enriched by hybridization capture. The enriched libraries are then sequenced using next generation sequencing on the Illumina NextSeq 550 platform.

Sequencing data are then analyzed using a custom-developed bioinformatics pipeline designed to detect SNVs, indels, CNAs and fusions from cfDNA. Results (detected or not detected) are presented in a results report. A not detected result from a plasma specimen for any given variant does not preclude the presence of this variant in tumor tissue.

The device is designed to detect pre-defined and *de novo* variants in the genes outlined in **Table 3**. Details on all variants reported can be found in the **section 8** *Additional Guardant360 CDx Variant Details*.

Alteration Type	Genes
Single Nucleotide Variants (SNVs)	AKT1, ALK, APC, AR, ARAF, ATM*, BRAF, BRCA1**, BRCA2**, CCND1, CDH1, CDK4, CDK6, CDK12*, CDKN2A, CTNNB1, EGFR, ERBB2, ESR1, FGFR1, FGFR2, FGFR3, GATA3, GNA11, GNAQ, HRAS, IDH1, IDH2, KIT, KRAS, MAP2K1, MAP2K2, MET, MLH1, MTOR, MYC, NF1, NFE2L2, NRAS, NTRK1, NTRK3, PDGFRA, PIK3CA, PTEN, RAF1, RET, RHEB, ROS1, SMAD4, SMO, STK11, TERT, TSC1, VHL
Indels	AKT1, ALK, APC, ATM*, BRAF, BRCA1**, BRCA2**, CDH1, CDK12*, CDKN2A, EGFR, ERBB2, ESR1, FGFR2, GATA3, HNF1A, HRAS, KIT, KRAS, MET, MLH1, NF1, PDGFRA, PIK3CA, PTEN, RET, ROS1, STK11, TSC1, VHL
Copy Number Amplifications (CNAs)	ERBB2, MET
Fusions	ALK, NTRK1, RET, ROS1

Table 3. Genes Containing Alterations Reported by Guardant360 CDx

*Reporting is enabled for pathogenic germline alterations only. Somatic alterations will not be reported.

** Reporting is enabled for both germline and somatic alterations.

5.4 Reagent, Material, and Equipment Usage

Reagents, materials, and equipment needed to perform the test are used exclusively in the Guardant Health Clinical Laboratory. Guardant360 CDx is intended to be performed with the following instruments, as identified by specific serial numbers.

- Agilent Technologies 4200 TapeStation Instrument
- Applied Biosystems Veriti 96-Well Thermal Cycler
- Hamilton Company Microlab STAR
- Hamilton Company Microlab STARlet
- Illumina NextSeq 550 Sequencing System
- Qiagen QIAsymphony SP Instrument

6 Summary of Performance Characteristics

Performance characteristics were established using clinical samples from patients with a wide range of cancer types, including those with NSCLC. The clinical samples consisted of pools of cfDNA from clinical samples from multiple cancer types, pools of cfDNA from clinical samples derived from one cancer type (e.g., samples from patients with NSCLC) or un-pooled clinical samples. Studies include CDx variants as well as a broad range of representative alteration types (SNVs, indels, CNAs, and fusions) in various genomic contexts across a number of genes. Due to limitations in clinical sample availability and due to the rarity of the fusions reported by the Guardant360 CDx, contrived samples were utilized for some non-clinical studies. A contrived sample functional characterization study

was conducted to demonstrate comparable performance of contrived samples made of cell line cfDNA and clinical sample cfDNA so that fusion cell line cfDNA material could be used in some non-clinical studies. Fusion positive clinical samples were used to confirm the estimated limit of detection, analytical accuracy and precision.

6.1 Analytical Accuracy/Concordance

a. Concordance - Comparison to NGS Comparator Method

The detection of alterations by Guardant360 CDx was compared to results of an externally validated NGS assay. Samples from 386 donors with different cancer types were collected for the study. Sixteen (16) samples failed testing with the comparator assay due to instrument failures, while eleven (11) samples failed testing with the Guardant360 CDx assay due to an instrument failure due to a power outage. 359 samples remained comprising three collection sets as follows.

Collection set one consisted of 100 donor samples selected with the comparator assay consecutively without selection for any specific variants. Since the first sample collection was expected to lack many rare variants, in the second collection set, a set of 100 positive samples were selected with the comparator assay. Collection set three consisted of 159 samples selected from the Guardant Health biobank based on Guardant360 LDT results to include additional rare variants including gene fusions which were not available from collection sets 1 and 2.

Of 359 patients, no samples failed QC on Guardant360 CDx, and three samples failed with the comparator NGS assay. In total, 356 donor samples across 18 cancer types, which all passed every QC metric were used for the concordance analysis. The cancer types represented in this study included lung (178), gastrointestinal (82), colon (25), breast (17), head and neck (13), prostate (12), genitourinary (7), bladder (3), stomach (3), pancreas (3), endocrine (2), liver (2), ovarian (2), kidney (2), gynecologic (1), esophagus (1), skin (1), and other (5). A summary of Positive Percent Agreement (PPA) and Negative Percent Agreement (NPA) with 95% confidence intervals (CI) is provided in **Table 4** for CDx alterations in samples from the intended use population, i.e., 176 patients with NSCLC. Agreement rates for each of the CDx variants ranged from 95% to 100% for PPA, and from 98.1% to 99.9% for NPA. The reported PPA and NPA were not adjusted for the distribution of samples from collection set 3 selected using Guardant LDT results. A summary of PPA and NPA for other clinically significant variant categories and for panel wide for SNVs and indels over all sample collections is provided in **Table 4**.

Positive agreement rates were evaluable for nine (9) patients with clinical Category 2 variants, which consisted of clinically relevant *PIK3CA* mutations in breast cancer patients that included E545A, E542K, E545K, H1047R, and H1047L variants. Concordance analysis resulted in 100% PPA and 100% NPA for the Category 2 variants.

Positive agreement rates for clinical Categories 3 and 4 variants resulted in 93.5% PPA and 86.1% PPA, respectively. Variants in clinical category 3 and 4 showed 99.8% and 100.0% NPA.

MET amplifications had a PPA of 56%, which is attributed to differences in reporting of copy number alterations by the Guardant360 CDx and the comparator assay. The Guardant360 CDx reports on only focal amplifications and not chromosome-arm amplifications, while the NGS comparator assay reports all amplifications.

The study demonstrated a PPA of 82.5% for indels, 91.4% for SNVs and >99% NPA for the entire reportable range, i.e., panel-wide, demonstrating the analytical accuracy of the device.

Alteration Type	Guardant360 CDx(+), Comparator (+)	Guardant360 CDx(+), Comparator (-)	Guardant360 CDx(-), Comparator (+)	Guardant360 CDx(-), Comparator (-)	Possible Variants (n)	Patients (n)	PPA (95% CI)	NPA (95% CI)
EGFR T790M	19	3	1	153	1	176	95.0% (75.1%, 99.9%)	98.1% (94.5%, 99.6%)
EGFR L858R	18	1	0	157	1	176	100.0% (81.5%, 100.0%)	99.4% (96.5%, 100.0%)
EGFR exon 19 deletions	30	1	1	1024	6	176	96.8% (83.3%, 99.9%)	99.9% (99.5%, 99.9%)
Category 2 Variants	9	0	0	76	5	17	100.0% (66.4%,100.0%)	100.0% (95.3%, 100.0%)
Category 3 Variants	115	11	8	6191	50	N/A*	93.5% (87.6%, 97.2%)	99.8% (99.7%, 99.9%)
Category 4 Variants	420	58	68	137582	388	356	86.1% (82.7%, 89.0%)	100.0% (99.9%, 100.0%)
MET CNAs	13	3	10	330	1	356	56.5% (34.5%, 76.8%)	99.1% (97.4%, 99.8%)
ERBB2 CNAs	15	0	2	339	1	356	88.2% (63.6%, 98.5%)	100.0% (98.9%, 100.0%)
NTRK1 Fusions	5	0	0	351	1	356	100.0% (47.8%, 100.0%)	100.0% (98.9%, 100.0%)
<i>RET</i> Fusions	11	2	1	342	1	356	91.7% (61.5%, 99.8%)	99.4% (97.9%, 99.9%)
ALK Fusions	10	2	0	344	1	356	100.0% (69.2%, 100.0%)	99.4% (97.9%, 99.9%)
ROS1 Fusions	11	0	0	345	1	356	100.0%	100.0%

Table 4. Summary of Concordance Between Guardant360 CDx and NGSComparator Method

							(71.5%,100.0%)	(98.9%, 100.0%)
Panel-Wide SNVs	428	48	40	1372684 4	38560	356	91.5% (88.5%, 93.8%)	99.9% (99.9%, 99.9%)
Panel-Wide Indels	118	19	25	1571723 8	44150	356	82.5% (75.3%, 88.4%)	99.9% (99.9%, 99.9%)

* For Category 3, no number is given. This is because Category 3 is a merge of many different variants, each with a specific set of cancer types that qualify the variant to belong in Category 3. This means that a different number of patients was associated with each variant within Category 3. For this level, the concordantly negative population was computed as the sum of the concordantly negative populations if each variant in this category was treated independently.

6.2 Contrived Sample Functional Characterization (CSFC) Study

A CSFC study was performed to demonstrate comparable performance between contrived samples that consisted of fusion cell line cfDNA material and fusion positive clinical sample cfDNA material. The CSFC study was performed using 5 ng DNA input (the lowest cfDNA input for the assay) to compare the performance of the Guardant360 CDx with cfDNA derived from cell lines and cfDNA derived from multiple clinical samples from multiple cancer types with ALK, NTRK1, RET, and ROS1 fusions. The cell line and clinical cfDNA sample pools contained known fusion events that were diluted with pools of wild-type (WT) cfDNA from multiple clinical specimens from multiple cancer types to pre-determined MAF levels (targeted levels were above and below LoD; see **Table 5**). Cell line cfDNA sample pools were tested across 13-20 replicates, 13 replicates for level 6, 14 replicates for level 2, and 20 replicates for the other levels at 5 ng cfDNA input. Clinical cfDNA sample pools from multiple cancer types were tested with 14 replicates at 5 ng cfDNA input. Both cell line and clinical cfDNA sample pools were tested with an orthogonal method to confirm MAF level. Detection rates of the 4 fusions, for each titration level, and for each of the two types of pools, are presented in Table 5.

Based on these analyses, the results demonstrate that the performance of the Guardant360 CDx is similar for both fusion positive contrived cfDNA samples and for fusion positive clinical cfDNA samples.

Fusion	Sample	Detection Ra	Detection Rate (95% confidence interval)							
	Туре	Level 1 Target MAF 0.07%	Level 2 Target MAF 0.175%	Level 3 Target MAF 0.35%	Level 4 Target MAF 0.7%	Level 5 Target MAF 1.4%	Level 6 Target MAF 1.8%			
EML4-ALK	Cell line	5.0% (0.1%, 24.9%)	28.6% (8.4%, 58.1%)	50.0% (27.2%, 72.8%)	90.0% (68.3%, 98.8%)	100.0% (83.2%, 100.0%)	100.0% (75.3%, 100%)			

Table 5. Fusion Detection Rate in the CSFC study

EML4-ALK	Clinical	7.1% (0.2%, 33.9%)	28.6% (8.4%, 58.1%)	50.0% (23.0%, 77.0%)	85.7% (57.2%, 98.2%)	100.0% (76.8%, 100.0%)	100.0% (76.8%, 100.0%)
CCDC6- RET	Cell line	15.0% (3.2%, 37.9%)	35.7% (12.8%, 64.9%)	80.0% (56.3%, 94.3%)	95.0% (75.1%, 99.9%)	100.0% (83.2%, 100.0%)	100.0% (75.3%, 100.0%)
TRIM33- RET	Clinical	7.1% (0.2%, 33.9%)	14.3% (1.8%, 42.8%)	64.3% (35.1%, 87.2%)	85.7% (57.2%, 98.2%)	100.0% (76.8%, 100.0%)	100.0% (76.8%, 100.0%)
ROS1- SLC34A2	Cell line	0.0% (0.0%, 16.8%)	21.4% (4.7%, 50.8%)	50.0% (27.2%, 72.8%)	75.0% (50.9%, 91.3%)	100% (83.2%, 100.0%)	100.0% (75.3%, 100%)
ROS1- CD74	Clinical	7.1% (0.2%, 33.9%)	42.9% (17.7%, 71.1%)	85.7% (57.2%, 98.2%)	100.0% (76.8%, 100.0%)	100.0% (83.9%, 100.0%)	ND
TPM3-NTRK1	Cell line	15.0% (3.2%, 37.9%)	50.0% (23.0%, 77.0%)	40.0% (19.1%, 63.9%)	90.0% (68.3%, 98.8%)	100.0% (83.2%, 100.0%)	100.0% (75.3%, 100.0%)
PLEKHA6- NTRK1	Clinical	21.4% (4.7%, 50.8%)	35.7% (12.8%, 64.9%)	85.7% (57.2%, 98.2%)	100.0% (76.8%, 100.0%)	ND	100.0% (76.8%, 100.0%)

ND: Not determined

6.3 Analytical Sensitivity

a. Limit of Blank (LoB)

The LoB was established by evaluating whole blood samples from healthy age-matched donor samples. sixty-two (62) donor samples confirmed to be mutation negative based on sequencing with an externally validated orthogonal method were processed using 30 ng of cfDNA input with the Guardant360 CDx (highest DNA input for the assay) across three lots of reagents, operator groups, and instruments. Of the 62 donor samples, 58 donor samples were tested with 4 replicates, while 4 donors were tested with 2 replicates for a total of 240 replicates analyzed to assess the false positive rate of Guardant360 CDx. This study demonstrated a near zero false positive rate across the entire reportable range, as shown in **Table 6**. The false positive rate was zero for Category 1 (CDx) and Category 2 variants.

Category	Per Position False Positive Rate	Per Sample False Positive Rate
Category 1: <i>EGFR</i> L858R	0%	0 (0/240)
Category 1: EGFR T790M	0%	0 (0/240)

Table 6. LoB Study Summary Results

Category 1: EGFR Exon 19 deletions	0%	0 (0/240)
Category 2	0%	0 (0/240)
Panel-wide SNVs (38,560 bp)	<0.00005% (4/(38,560*240))	1.67% (4/240)
Panel-wide Indels (44,150 bp)	<0.00002% (2/(44,150*240))	0.83% (2/240)
Panel-wide CNAs (2 genes)	0.2% (1/(2*240))	0.42% (1/240)
Panel-wide Fusions (4 genes)	0%	0 (0/240)

b. Limit of Detection (LoD)

The LoD for the Guardant360 CDx variants with CDx claims, representative SNVs and indels, and all reportable CNAs and fusions was established at the lowest and highest claimed cfDNA input amounts (5 and 30ng). LoD established for fusions using cfDNA derived from cell lines was confirmed at 5ng cfDNA input using cfDNA derived from clinical patient samples. LoDs were further confirmed in the clinical pools of relevant cancer types for CDx variants and additional representative variants, including long indels and homopolymers in a combined LoD confirmation and precision study.

For SNVs, indels, including CDx variants and for CNAs, the Guardant360 CDx LoD was established by combining cfDNA from clinical plasma samples from multiple cancers to create pools of material comprising multiple known alterations. The LoD was established with these clinical cfDNA sample pools at 5ng and 30ng input, using a combination of probit and empirical approaches. Samples were titrated at 5 different MAF values that included levels above and below the LoD for SNVs, and indels or copy numbers values for CNAs and tested across 20 replicates for 5 ng input and 14 replicates for 30 ng input across two reagent lots.

The LoDs of three (3) CDx alterations representing *EGFR* T790M, *EGFR* L858R, and *EGFR* exon 19 deletions established using pools of cfDNA from clinical plasma samples from multiple cancer types are summarized in **Table 7**. The LoD was confirmed for CDx variants using cfDNA sample pools from patients with NSCLC only; refer to **Table 9** below.

Table 7. Summary of Estab	lished LoD for Alte	rations Associate	ed with CDx Claims		
using Pools of cfDNA from	using Pools of cfDNA from Clinical Plasma Samples from Multiple Cancer Types				

Alteration	Alteration Type	LoD (5ng input)	LoD (30 ng input)
EGFR T790M	SNV	1.1% MAF	0.2% MAF
EGFR L858R	SNV	1.0% MAF	0.2% MAF
EGFR exon 19 deletion	Indel (15 bp)	1.5% MAF	0.2% MAF

The LoD estimates for SNV, indels, and CNA alterations established using pools of cfDNA from clinical plasma samples from multiple cancer types are summarized in **Table 8**.

For fusions, the Guardant360 CDx LoD was established using cfDNA from cell lines with known fusions titrated into wild-type (WT) cfDNA from clinical plasma samples. Samples were titrated at 5 different MAF values for fusions across 20 replicates for 5 ng cfDNA input and 14 replicates for 30 ng cfDNA input across two reagent lots. The established LoD was then confirmed using fusion positive cfDNA from clinical plasma samples at 5 ng cfDNA input only. Fusion positive cfDNA from clinical samples were titrated across 5 concentrations with 14 replicates across 2 reagent lots.

The higher of the LoD values established using cell lines and confirmed using clinical samples were used to claim the LoD performance levels of the test for fusions at 5 ng (**Table 8**).

Alteration	Alteration Type	LoD, 5 ng (MAF/CN)	LoD, 30 ng (MAF/CN)
BRAF V600E	SNV	1.8%	0.2%
KRAS G12V	SNV	1.5%	0.5%
NRAS Q61R	SNV	3.0%	0.8%
BRCA1 p.E23fs	Indel (2 bp)	2.6%	0.8%
<i>BRCA2</i> p.S1982fs	Indel (1 bp)	1.3%	0.4%
<i>EGFR</i> exon 20 insertion, p.Ala767_Val769dup	Indel (9 bp)	0.8%	0.2%
<i>ERBB2</i> exon 20 insertion, p.A775_G776insYVMA	Indel (12 bp)	1.1%	0.2%
MET	CNA	2.4	2.4
ERBB2	CNA	2.3	2.3
NTRK1	Fusion	0.9% (0.9%)	(0.2%)
RET	Fusion	1.1% (0.7%)	(0.1%)
ROS1	Fusion	1.9% (1.2%)	(0.2%)
ALK	Fusion	1.4% (1.5%)	(0.2%)

Table 8. LoD Establishment Study Summary Results for Representative Variantsusing Pools of cfDNA Clinical Plasma Samples from Multiple Cancer Types

Note: Numbers in parentheses represent LoD established using cell line derived cfDNA. MAF: Mutant Allele Fraction, CN: copy number

The established LoD was confirmed for CDx variants by testing clinical patient pools exclusively from NSCLC patients targeting 1-1.5x LoD of the established LoD (refer to **Table 9**) across at least 20 replicates at 5 ng input using a combined LoD Confirmation and Precision Study. Similarly, the established LoD was confirmed for SNVs and indels

in clinical pools made exclusively from the relevant cancer type source material prepared with 5 ng cfDNA input targeting 1-1.5x LoD and run in at least 20 replicates targeting 5 distinct variants. Established LoD targets were used for 5 variants (*EGFR* L858R, *EGFR* T790M, *EGFR* exon 19 deletion, p.E746_A750del, *KRAS* G12C, and *ROS1* fusions), while in silico LoD targets were used for 10 additional variants to target variants to 1-1.5x LoD.

In this combined LoD and Precision study, (see Section 6.5. below for additional studies demonstrating assay precision starting from cfDNA extraction, and with additional mutation positive and negative samples) samples were tested across three precision combinations that evaluated three operator groups, three instrument combinations, and three SPK reagent lots over at least three different start dates.

The higher of the LoD values established using clinical sample pools from cancer patients and confirmed using clinical samples exclusively from the relevant cancer type source material were used to claim LoD performance of the test at 5 ng input as summarized in **Table 9**.

Alteration	MAF	Alteration Type	Cancer Type	Number Positive / Number Expected	РРА
EGFR L858R	1.5%*	SNV	NSCLC	20/20	100.0%
EGFR T790M	1.4%*	SNV	NSCLC	19/20	95.0%
EGFR exon 19 deletion, p.E746_A750del	1.5%*	Indel (15bp)	NSCLC	20/20	100.0%
<i>EGFR</i> exon 19 deletion, A750_I759delinsPT	2.3%^	Indel (29 bp)	NSCLC	20/20	100.0%
<i>KIT</i> V654A	2.5%^	SNV	Prostate	20/20	100.0%
KRAS G12C	1.8%*	SNV	NSCLC	19/20	95.0%
<i>PIK3CA</i> E545K	2.4%^	SNV	Breast	21/21	100.0%
<i>PIK3CA</i> H1047L	1.7%^	SNV	Breast	21/21	100.0%
EGFR exon 20 insertion, H773_V774insHPH	3.5%^	Indel (9 bp)	NSCLC	22/22	100.0%
<i>MET</i> exon 14 skipping 7.116412041.AAGGTATATT TCAGTT>A	2.7%^	Indel (15 bp)	NSCLC	20/20	100.0%
<i>BRCA2</i> p.T3033fs	4.4%^	Indel (1 bp), homopolymer	NSCLC	21/21	100.0%
<i>BRCA2</i> p.I605fs	5.0%^	Indel (1 bp), homopolymer	Prostate	20/20	100.0%
<i>BRCA2</i> p.V1532fs	4.2%^	Indel (1 bp), homopolymer	Prostate	20/20	100.0%

Table 9. Combined LoD Confirmation and Precision Study Summary Results forCDx Variants and Representative Variants

<i>STK11</i> p.L282fs	4.7%^	Indel (1 bp), homopolymer	NSCLC	21/21	100.0%
ROS1	1.8%*	Fusion	NSCLC	21/21	100.0%

* Observed MAF level in LoD Confirmation Study. LoD confirmed with single cancer type clinical pool and ≥95% detection rate is within 1-1.5x LoD MAF level from the original establishment study range. ^ Observed MAF at the level tested with ≥95% detection rate for variants without direct prior LoD establishment data.

Panel-wide SNV and indels detected by Guardant360 CDx is summarized in **Table 10** as median values.

Table 10. Summary of LoD for Alterations Associated with Panel-Wide Claims

Alteration	Median LoD, 5ng (MAF)	Median LoD, 30ng (MAF)
Panel-wide SNVs	1.8%	0.2%
Panel-wide Indels	2.7%	0.2%

6.4 Analytical Specificity

a. Endogenous and Microbial Interfering Substances

To evaluate the potential impact of endogenous and microbial interfering substances on the performance of Guardant360 CDx, this study evaluated whole blood samples from a total of 50 patients (at least ten patients per interfering substance), representing more than 13 cancer types. The 130 samples that passed QC checks included representative variants.

Substances were considered as non-interfering if, when compared to no interferent controls, the sample level molecule recovery, exon-level molecule recovery, and variant call concordance met pre-defined acceptance thresholds.

Sample level molecule recovery was determined by the depth of non-singleton molecule (NSC) coverage across the panel. Median non-singleton molecule coverage across targeted regions was evaluated to demonstrate that microbial or interfering substances do not impact assay performance to sequence unique molecules. Recovery of unique molecules across interfering substance conditions did not show a negative impact of interfering substances (fold change of median NSC in spike condition over reference condition ranged from 0.88 to 1.08).

Relative exon coverage calculated as the ratio of median exon coverage to sample level coverage for each of the 508 exon regions was compared for each condition-reference sample pair. Aggregating across all samples contributing to the analysis, the total fraction of all exonic regions within expected level of differences defined as $2^* \sigma$, where σ is the pooled standard deviation of the differences observed in historical (σ =0.108) were calculated. Under normal distribution assumption, the fraction of such regions is expected to be 95%. The fraction of exons with relative exon level coverage difference

between condition and reference within 2σ (2 * 0.108) was 94.3-99.7%, which demonstrate that there was no preferential drop-out of relative exon-level coverage exceeding expected levels due to random variation, and the entire panel was covered consistently between reference and interfering substance conditions.

The results were aggregated across all variants across all ten whole blood samples, and concordance was assessed within each treatment category across variants. PPAs were calculated for 62 SNVs, 24 indels, and 3 CNAs. The 6 conditions tested showed variant call concordant PPAs ranging from 83.3%-100.0%. PPA \ge 1x LoD ranged from 90.0%-100.0% for all 6 interferents.

The panel-wide NPAs were also calculated for SNVs and indels within the reportable range. The discordant negative variants were defined as those negative variants that were positive in the non-reference condition. The panel-wide NPA was 99.9%-100.0% for all conditions.

In conclusion, no interference was found in Albumin (60 g/L), Bilirubin (conjugated) (342 μmol/L), Bilirubin (unconjugated) (342 μmol/L), Hemoglobin (2 g/L), Staphylococcus epidermidis (106 cfu) or Triglycerides (15 g/L).

The effect of potential exogenous interfering substances on assay performance was not evaluated. A post-market study will be conducted to evaluate the effect of exogenous interfering substances on assay performance.

b. In silico Analysis

Primer and probe specificity were addressed by mapping panel probes to the human genome. When mapped to the human genome (hg19) with decoy sequences, unplaced contigs, and representative microbial contaminants genomes, 97.6% of probes uniquely map to the genome (MAPQ \geq 60). None of the primers or probes mapped to the representative microbial contaminant genomes.

6.5 Precision

The purpose of the precision studies was to demonstrate the repeatability and withinsite reproducibility of Guardant360 CDx through closeness of agreement between measured qualitative output obtained in replicate testing using different combinations of reagent lots, instruments, operators, and days. Additional runs were conducted (1) on mutation negative samples to demonstrate precision of analytically blank samples, and (2) on plasma samples to understand the influence of extraction on test. All studies were conducted exclusively with patient material and no cell line material was used.

a. Precision across three distinct cfDNA clinical sample pools

Precision was evaluated for alterations associated with CDx claims, as well as representative and specific alterations to support platform-level performance. Repeatability including intra-run performance (run on the same plate under the same conditions) and reproducibility including inter-run performance (run on different plates under different conditions) were assessed and compared across three different precision combinations of instrument sets, reagent lots, and operators over multiple days. This study was carried out on three distinct clinical sample pools from multiple cancer types, containing a total of 16 targeted alterations across the pools, prepared targeting 1-1.5x LoD at 5 ng cfDNA input, included variants associated with CDx claims and additional variants intended to demonstrate panel-wide validation. Ten (10) replicates per three (3) pools were tested for each of three (3) precision combinations (90 replicate samples total) and comprised of three (3) different reagent lots (G360 SPK, Ampure XP beads, and NextSeq 550 sequencing reagent lots), three (3) different instrument sets and three (3) different operator groups. Each combination was tested on two (2) batches, sequenced on four (4) flow cells. The QIAsymphony instrument was not paired within each of the three (3) precision combination sets, since the sample pools were generated from previously extracted and stored cfDNA. Precision starting from cfDNA extraction was evaluated in a separate study described in Section 6.5.b. below. In total, 480 alterations were assessed across 90 samples tested. Qualitative results were used to calculate PPA and NPA.

The final levels for the targeted variants tested ranged from 0.7x to 2.6x LoD. Three variants were below 1x LoD (*ROS1* fusion at 0.9x LoD, *MET* amplification at 0.8X LoD, and *NRAS* Q61R at 0.7x LoD), 8 were within 1-1.5x range, including the CDx variants, and 5 variants were in the 1.7x - 2.6x LoD range.

Across 960 expected negative targeted sites (32 targeted negative variants across 3 sample pools * 30 replicates), the observed NPA was 100.0%.All CDx alterations demonstrated acceptable precision (PPA 96.7%-100.0%), **Table 11**.

The variant level PPA for all targeted variants were above 90.0% across all instrument, reagent, and operator combinations, except for *MET* amplification in pool 1, which may be attributed to the 0.8x LoD range achieved in the titration pool (**Table 11**). *ROS1* fusion detection demonstrated 93.3% PPA, consistent with the achieved 0.9x LoD titration level. *BRCA1* E23fs also resulted in a lower variant level PPA (90.0%) than expected. However, the 90.0% detection rate is consistent with the variant being located in a more challenging area of the panel with respect to coverage. Specifically, the variant is considered to be in a more challenging area because it is in a region with relatively low GC content and has below average DNA molecule recovery.

Across 480 alterations (150 SNVs, 150 indels, 60 CNAs, and 120 fusions), from a set of 90 cfDNA sample replicates containing 16 unique alterations across 3 cfDNA sample pools made from cfDNA from multiple cancer types, all alterations demonstrated PPA of 86.7%-100.0%. Alteration-level repeatability and reproducibility showed high overall positive call rates (**Table 11**).

Alteration Class	Alteration	Number Positive / Number Expected	PPA (95% CI)
SNV	EGFR T790M	30/30	100.0% (88.4%, 100.0%)
SNV	EGFR L858R	30/30	100.0% (88.4%, 100.0%)
Indel	EGFR Exon 19 Del, E746_A750del	29/30	96.7% (82.8%, 99.9%)
SNV	KRAS G12V	30/30	100.0% (88.4%, 100.0%)
SNV	NRAS Q61R	30/30	100.0% (88.4%, 100.0%)
SNV	BRAF V600E	30/30	100.0% (88.4%, 100.0%)
Indel	ERBB2 A775_G776insYVMA	30/30	100.0% (88.4%, 100.0%)
Indel	EGFR A767_V769dup	30/30	100.0% (88.4%-100.0%)
Indel	BRCA1 E23fs	27/30	90.0% (73.5%-97.9%)
Indel	BRCA2 S1982fs	30/30	100.0% (88.4%-100.0%)
CNA	ERBB2	30/30	100.0% (88.4%-100.0%)
CNA	МЕТ	26/30	86.7% (69.3%-96.2%)
Fusion	EML4-ALK	30/30	100.0% (88.4%-100.0%)
Fusion	TPM3-NTRK1	30/30	100.0% (88.4%-100.0%)
Fusion	TRIM33-RET	30/30	100.0% (88.4%-100.0%)
Fusion	ROS1-CCDC6	28/30	93.3% (77.9%-99.2%)
SNV	Panel-wide	150/150	100.0% (97.6%-100.0%)
Indel	Panel-wide	146/150	97.3% (93.3%-99.3%)

Table 11. Summary of Precision PPA Results

The PPA across all targeted alterations for each condition was evaluated. The PPA across all targeted alterations per precision combination (PC) ranged from 96.3%-99.4%.

Precision from clinical pools with samples from a single clinically relevant cancer type was confirmed in the combined LoD confirmation and precision study described in Section 6.3.b above.

b. Precision from plasma evaluation of extraction precision and precision of downstream steps

The purpose of this study was to show the precision of variant calling for the entire sample workflow (from cfDNA extraction through sequencing) with un-pooled clinical samples.

This study utilized clinical plasma samples from 53 unique patients. Each plasma sample with positive variants (as detected by Guardant360 LDT) and high cfDNA yields was split into six aliquots or six replicates per patient.

The LoD was established for inputs of 5 ng and 30 ng, which are the lower and upper limit of cfDNA mass input for library preparation. Since the purpose of this precision study was to test the full spectrum of sample yields that would be observed in normal use, sample inputs ranged from 5 ng to 30 ng of cfDNA input. The corresponding LoD range was between 1x the 30 ng LoD MAFs, and 1.5x the 5 ng LoD MAFs. Variants that were previously observed in this MAF range in the Guardant360 LDT run were selected for this study and evaluated for call agreement.

Eighteen (18) different tumor types were evaluated in this study to support a pancancer tumor profiling indication for Guardant360 CDx. Each donor specimen was processed in duplicate across three lots for a total of 6 replicates. "Lot" refers to different reagent lots, as well as different combinations of operators, days, and instruments to evaluate precision. The targeted variants evaluated in the study are shown in **Table 12**.

Category	Variant	Number of Eligible Based on MAF/CN
ERBB2	CNA	3
МЕТ	CNA	3
ALK	fusion	2
RET	fusion	2
EGFR exon 19 deletion	indel	6
EGFR exon 20 insertion	indel	2
Long indel (>30 bp)	indel	1
MET exon 14 skipping	indel	1
BRAF V600E	SNV	3
<i>EGFR</i> L858R	SNV	6
EGFR T790M	SNV	4
KRAS G12C	SNV	3
<i>PIK3CA</i> E542K	SNV	3
<i>PIK3CA</i> E545K	SNV	4

 Table 12. Targeted Variants Amongst the 53 Donor Samples Selected for Study

<i>PIK3CA</i> H1047L/R	SNV	2
PIK3CA C420R	SNV	3

A total of 315 replicates passed QC and were analyzed for within-condition and between-condition precision.

For each eligible variant, pairwise comparisons of variant detection were made between the technical replicates in each lot. From the study design with three lots and two replicates within each lot, there were 3 pairs for each variant in calculating withinlot average positive agreement (APA) and 12 pairs for each variant in calculating between-lot APA.

The APA results for eligible SNVs, indels, fusions, CNAs and all three together are shown in **Table 13**. Workflow or sample QC failures mean there were fewer than 3 lots per variant tested in some cases. The within lot APA for all variant types together was 97.3% as shown in **Table 13**.

Variant Type	Variant Lot Comparisons	Concordant (C)	Discordant (D)	APA	
SNV	150	141	9	96.9%	
Indel	35	35 0		100.0%	
CNA	15	13	2	92.9%	
Fusion	12	12	0	100.0%	
ALL	212	201	11	97.3%	

Table 13. Within Reagent Lot APA Summary

The within-lot ANA was 99.9%. This statistic includes all called variant sites panel-wide, not just the eligible variants sites based on LoD in the source samples, so this statistic includes positions with expected stochastic detection due to low mutant molecule count. The number of positions evaluated was 46,217 unique SNV and indel reportable positions, 2 CNAs, and 4 fusions.

The between lot APA for eligible SNVs, indels, fusions, CNAs, and all reportable variants together are shown in **Table 14**. For each of these variants, there were 12 pairwise comparisons.

Variant Type	Variant Lot Comparisons	Concordant	Discordant	АРА	
SNV	47	531	26	97.6%	
indel	11	132	0	100.0%	
CNA	8	53	6	94.6%	
fusion	fusion 4		0	100.0%	
ALL	ALL 70		32	98.0%	

Table 14. Between-Lot APA Summary

The between-lot APA for all variant types together was 98.0% Between lot ANA was 99.9% across all reportable positions and variants. This statistic includes all called variant sites, not just the eligible variants sites based on LoD in the source samples, so includes positions with expected stochastic detection due to low mutant molecule count. The number of positions evaluated was 46,217 unique SNV and indel reportable positions, 2 CNAs, and 4 fusions.

Notably, for *ERBB2* amplifications, within and between lot APA were observed to be 80.0% and 85.0%, respectively, due to variation in focality determination. Specifically, some of the replicates were determined to be focally amplified, and thus reported by the assay, and some were determined to be aneuploid and thus reported negative as the Guardant360 CDx reports CNAs only for focal amplifications and not chromosome-arm amplifications.

In addition to the main study, supplementary samples, starting from plasma, were processed to evaluate precision from extraction. Fusion samples were created by diluting cfDNA extracted from cell lines harboring *ROS1* and *NTRK1* fusions into plasma of clinical lung cancer samples negative for fusions. These contrived plasma samples were evaluated in lieu of clinical samples for this study due to the rarity of these alterations. Plasma was processed from extraction to sequencing on the same batches as the rest of the study samples. The fusion cfDNA was diluted to < 0.2% MAF for *ROS1* and *NTRK1* at ~30 ng input. There was 100% detection (6/6) across reagent lots for both fusions when tested at 0.15% MAF at approximately 30 ng of cfDNA.

c. Precision from mutation negative samples

Samples from healthy donors were pre-screened by an externally validated orthogonal method. Mutation negative samples by the orthogonal method were tested by Guardant360 CDx in three reproducibility conditions (i.e., different reagent lots, operators, instruments, and days). Four replicates from each donor were tested with Guardant360 CDx across the different reproducibility conditions. The study demonstrated a sample-level, within-condition ANA of 97.4% and sample-level between-condition ANA of 97.3%. The within-condition ANA was 99.6% and between-condition. Within-condition ANA values were 100.0% for all CDx variants (*EGFR* L858R, *EGFR* T790M, and *EGFR* exon 19 deletions) and category 2 variants.

6.6 Cross-Contamination/Carry-Over

The carryover/cross-contamination study evaluated the prevalence of crosscontamination when material is transferred between samples in the same batch and carry-over when material is transferred between samples across batches processed sequentially on the same instrument using Guardant360 CDx. A total of 352 plasma samples across 8 batches (44 samples/batch x 8 batches) were run in a consecutive order across instruments within the analytical accuracy study and sequenced on 16 flowcells.

There was no evidence of high positive variants from near-by wells detected in negative samples. In conclusion, no carryover or cross-contamination was observed in 352 samples processed across 8 consecutive batches

6.7 Reagent Lot Interchangeability

Reagents lot interchangeability was assessed by testing two cfDNA sample pools containing 16 alterations, 9 variants in pool 1 and 7 variants in pool 2, in five replicates using two different lots of Guardant360 CDx Sample Preparation Kit in seven different lot combinations. For the sample replicates that proceeded to sequencing, all met the performance metrics. Kit Lot Interchangeability of G360 SPK boxes was evaluated based on the rate of positive agreement for detection of targeted variants.

Out of 70 samples, 68 passed QC metrics (97% pass rate). The rate of qualitative agreement rate (QDR), i.e., the agreement with the majority call for baseline reagent was calculated. QDR was defined as the number of positively detected targeted variants across eligible samples (D) divided by the total number of targeted variants tested across eligible samples (N), expressed as a percentage (100 * D/N). QDR ranged from 91.6% to 98.7%. There was 100.0% negative agreement among expected negative sites within respective pool replicates.

The panel-wide assessment of NPA was 99.9% calculated from negative variant sites across the Guardant360 CDx reportable range that are not detected in the reference condition represents SPK Lot A for all combinations tested.

6.8 Stability

a. Reagent Stability

The stability of the Guardant360 CDx Sample Preparation Kit lots used in sample processing for Guardant360 CDx were evaluated in this study. Three lots of identical reagents were stored under the specified storage conditions for each box and then tested at defined time points using two cfDNA sample pools that contained in total 16 known variants, 9 variants in pool 1 and 7 variants in pool 2. Under the tested conditions, results from each time point, 3, 4, 7, 10, 13 and 19 months were compared against samples tested at day 0 (time point T_0). The G360 SPK boxes were tested at each timepoint with five (5) replicates per each of the two unique sample pools at 5 ng cfDNA input.

Qualitative detection rates (QDR), which is based on the agreement with the majority call at T0 for the number of targeted variants detected, were assessed per lot/per timepoint. QDR was defined as the number of positively detected targeted variants that were positively detected in the baseline condition across eligible samples (D) divided by the total number of positively detected targeted variants

tested across eligible samples (N), expressed as a percentage (100 * D/N). The study showed no significant difference between time points compared to T0 for all three lots (alpha = 0.05), demonstrating that there was no significant decline in detection rates over the course of the study. The qualitative detection rate, calculated from targeted sites, ranged between 95.0% and 100.0% by timepoint. All of the expected negative variants were observed as negative calls across all replicates, indicating 100% negative agreement among all targeted variants expected to be negative across study conditions. The panel-wide assessment of NPA was 99.9% calculated from negative variant sites across the Guardant360 CDx reportable range that are not detected in the reference condition representing time 0 for all time points tested.

Variant detection performance was stable for a claimed shelf life of 18 months.

b. Whole Blood Stability

The objective of this study was to demonstrate the stability of whole blood specimens used for Guardant360 CDx collected in the Guardant360 BCK, that is in Streck Cell-Free DNA BCTs, across the expected range of sample transport and storage conditions for up to 7 days after blood collection prior to plasma isolation. The stability of whole blood used for Guardant360 CDx was evaluated by collecting 4 fresh whole blood samples from 16 cancer patients. From each patient, one tube was processed to plasma 1 day after blood draw (storage at room temperature). Plasma was then shipped on dry ice to Guardant Health. This constituted the reference condition. In addition to the reference tube, three more blood tubes per donor were shipped as whole blood to Guardant Health and subjected to Condition 1 (Summer profile), Condition 2 (Winter profile) or Condition 3 (Room temperature) as follow:

- <u>Reference Condition</u>: Plasma processing 1 day after blood collection)
- <u>Condition 1: Summer Profile Storage</u>: 4h at 22°C, 6h at 37°C, and 56h at 22°C, 6h at 37°C, plus remaining time at room temperature.
- <u>Condition 2: Winter Profile Storage</u>: 4h at 18°C, 6h at 0°C, 56h at 10°C, and 6h at 0°C plus remaining time at room temperature
- <u>Condition 3: Room Temperature Storage</u>: Storage at room temperature 18-25°C

After conditioning, plasma was isolated on the 8th day after blood collection and run on the Guardant360 CDx.

All 64 samples passed all QC and were included in analysis. All storage conditions demonstrated acceptable performance All samples in each group demonstrated acceptable sample-level molecule recovery as assessed by depth of NSC coverage across the panel. Fold change of median NSC in test condition over the reference condition or time zero ranged from 0.90 to 0.97.

Exon-level coverage was also acceptable for all conditions evaluated. The fraction of exons with relative exon level coverage difference between condition and reference (Time zero) within 2σ (2 * 0.108) was 95.3-96.3%, which demonstrate that there

was no preferential drop-out of relative exon-level coverage exceeding expected levels due to random variation, and the entire panel was covered consistently between reference and interfering substance conditions.

PPAs were also calculated for the SNVs and indels in the reportable range: 10 SNVs and 6 indels. All conditions showed variant call concordant PPA of 87.5% - 93.8%. PPA above LoD was 100.0% for all conditions. The data indicate acceptable sensitivity and specificity when using samples across the storage conditions.

The panel-wide NPAs were also calculated for SNVs and indels within the reportable range within 55 genes, CNAs and fusions. The total set of negative variants was set to the reportable range excluding variants found to be positive in the reference condition. The discordant negative variants were defined as those negative variants that were positive in the non-reference condition. The panel wide NPA was 99.9% for condition 1 (739,550 out of 739,552 variants), 99.9% (739,550 out of 739,552 variants) for condition 2, and 99.9% (739,548 out of 739,552 variants) for condition 3.

The whole blood stability study described above was supplemented by an additional study with two objectives: (1) to demonstrate the concordance between samples processed into plasma on the same day as blood collection and the samples processed into plasma the day after collection; (2) robustness to changes in relative humidity (RH) that tubes may be exposed to during shipping.

A total of four BCTs were drawn 19 healthy donors. For each donor, one BCT was processed to plasma within 4 hours after blood collection and shipped to Guardant Health on dry ice on the same day. This served as the reference condition. The other 3 BCTs will be subjected to conditions described below:

- <u>Test condition 1:</u> Intact whole blood in BCTs packed in BCKs was shipped overnight to Guardant Health and plasma isolation was done on the day of receipt (Day 1 after blood collection).
- <u>Test condition 2:</u> Exposure of whole blood in BCT starting on the day of blood collection and for 1 day to low humidity (25% RH, at 23°C) storage profile, followed by storage at Room temperature for 1 day . Plasma isolation occurred on Day 2 after blood collection.
- <u>Test condition 3:</u> Storage of whole blood in BCT starting on the day of blood collection and for 1 day at Room temperature, followed by exposure to high-humidity (90% RH, at 23°C) storage profile for 1 day. Plasma isolation occurred on Day 2 after blood collection.

Out of 76 samples processed, 24 study samples (6 distinct donor samples for all 4 conditions) had cfDNA underloading in some samples and overloading in some other samples due to a Guardant operator error. After QC check, 52 samples from 13 donors passed all sample QC metrics and were included in the analysis. Recovery of unique molecules across the 3 conditions did not show a negative impact of Day 1 processing and exposure of tubes to high (90% RH) and low (25% RH) relative humidity conditions. Fold change of median NSC in storage condition over reference

condition ranged from 0.95 to 0.99. For the reportable range of the device, the fraction of exons with relative coverage within 2σ (2 * 0.108) ranged 98.1 – 99.0%.

Based on the evidence from preservation of overall coverage and relative exon coverage the quantity and quality of cfDNA are not impacted by: (1) whole blood collection at vendor site and overnight shipping to Guardant Health at room temperature, followed by standard plasma isolation on day 1 after collection, (2) exposure of whole blood in BCT starting on the day of blood collection and for 1 day to low relative humidity (25% RH, at 23°C) storage profile, followed by storage at Room temperature for 1 day and plasma isolation on Day 2 after blood collection, and (3) Storage of whole blood in BCT starting on the day of blood collection and for 1 day at Room temperature, followed by exposure to high relative humidity (90% RH, at 23°C) storage profile for 1 day and plasma isolation on Day 2 after blood collection and for 1 day at 23°C) storage profile for 1 day and plasma isolation on Day 2 after blood collection and for 1 day and plasma isolation on Day 2 after blood collection and for 1 day at 23°C) storage profile for 1 day and plasma isolation on Day 2 after blood collection and for 1 day at 23°C) storage profile for 1 day and plasma isolation on Day 2 after blood collection and for 1 day and plasma isolation on Day 2 after blood collection and for 1 day and plasma isolation on Day 2 after blood collection and for 1 day and plasma isolation on Day 2 after blood collection and for 1 day and plasma isolation on Day 2 after blood collection and for 1 day and plasma isolation on Day 2 after blood collection and collection and collection and plasma isolation on Day 2 after blood collection and collection and collection and plasma isolation on Day 2 after blood collection and c

Based on these study results, whole blood may be stored in Cell-Free DNA BCTs tubes for up to 7 days after blood collection and prior to plasma isolation and can withstand winter and summer shipping conditions.

c. Plasma Stability

To define the storage conditions and evaluate the stability of plasma isolated from whole blood, stability at defined temperatures and durations was assessed. Samples were processed and run on Guardant360 CDx immediately after plasma isolation or after storage at -80°C \pm 10°C for 46 days or 2-8°C for 24 hours. Four BCTs from 12 cancer patients, 48 samples in total, were collected and run on Guardant360 CDx, with plasma stored at the specified storage conditions. Plasma from one BCT was processed through cfDNA extraction on the same day as a reference condition, plasma from a second BCT was stored at 2-8°C for 25 hours before cfDNA extraction (for a 24-hour stability claim at 2-8°C; Condition 1), plasma from a third BCT was stored at -80°C \pm 10°C with two freeze/thaw cycles for 46 days before cfDNA extraction to support usage of stored plasma for analytical validation (AV) studies (Condition 3). Extracted cfDNA from each condition was stored at -20°C \pm 5°C until further processing.

Out of 48 samples processed, 40 study samples (11 samples in reference condition, 8 samples in Condition 1, 10 samples in Condition 2 and 11 samples in Condition 3) passed their respective in-process and post-sequencing QC metrics and had at least one reference-condition sample pair, thus were included in the final analysis. In the three tested storage conditions, samples demonstrated acceptable performance. In the three tested storage conditions, samples demonstrated acceptable sample-level molecule recovery, relative exon-level coverage, and variant call concordance.

Sample-level molecule recovery showed fold change of 0.93, 1.10 and 0.9. Exon-level relative coverage demonstrated 92.8%-97.1% fraction of exons within 2σ of expected relative coverage.

PPAs were also calculated for the SNVs and indels in the reportable range within 55 genes that are reportable by test, as well as the reportable CNA and fusion genes: 14 SNVs, 1 indel and 1 CNA. Three conditions showed variant call concordant PPA of 76.9% - 78.6%. PPA above LoD was 90.9% - 91.7% for all conditions (a single variant was discordant). NPA across the reportable range was 99.9%.

Based on these study results, plasma may be stored at 2-8°C for 24 hours or at -80°C \pm 10°C with 2 freeze/thaw cycles for 1 year before cfDNA extraction.

d. cfDNA Stability

To define the storage conditions and evaluate the stability of cfDNA extracted from the plasma of whole blood, stability at defined temperatures and durations was assessed. Eighty-eight (88) samples were collected from 22 patients and run on Guardant360 CDx, with cfDNA stored in the specified storage conditions. Samples were split into two extraction arms (with quantification either before, or after freezing) to establish stability of cfDNA under both measurement workflows.

Sixty-six (66) samples were processed for the reference and 2 conditions below.

- <u>Reference condition A</u>: Post-extraction quantitation: Quantitation, dilution, and library preparation post-extraction on the same day
- <u>Reference condition B</u>: Quantitation, dilution, and library preparation postextraction on the same day
- <u>Condition 1A:</u> Quantitation and dilution post- extraction on the same day, followed by storage of cfDNA at 2-8°C for 25 hours (in FluidX tubes) before library preparation (for a 24-hour stability claim at 2-8°C).
- <u>Condition 1B</u>: Storage of cfDNA at 2-8°C for 25 hours (in Biorad elution plate), followed by quantitation and library dilution, before library preparation (for a 24-hour stability claim at 2- 8°C).
- <u>Condition 2A</u>: Quantitation and dilution post- extraction on the same day, followed by storage of cfDNA at -20°C ± 5°C plus 2 freeze/thaw cycles for 46 days (in FluidX tubes) before library preparation (for a 45-day stability claim at -20°C ± 5°C).
- <u>Condition 2B</u>: Storage of cfDNA at -20°C ± 5°C plus 2 freeze/thaw cycles for 46 days (in Biorad elution plate), followed by quantitation and library dilution, before library preparation (for a 45- day stability claim at -20°C ± 5°C).
- <u>Condition 3A</u>: Quantitation and dilution post-extraction on the same day, followed by storage of cfDNA at -20°C ± 5°C plus 5 freeze/thaw cycles for one year to support usage of stored cfDNA for AV studies in FluidX tubes before library preparation.
- <u>Condition 3B</u>: Storage of cfDNA at -20°C ± 5°C plus 5 freeze/thaw cycles for one year to support usage of stored cfDNA for AV studies (in Biorad elution plate), followed by quantitation and library dilution, before library preparation.

Out of 88 samples processed, 87 study samples passed QC metrics and were included in the final analysis. In the 3 tested storage conditions in both arms, samples demonstrated acceptable performance.

The recovery of unique molecules across storage conditions did not show a negative impact of storage: fold change of median NSC in storage condition over reference condition ranged from 0.93 to 1.06 in arm A (quantitation post-extraction); and from 0.90 to 0.96 in arm B (quantitation post-storage).

Relative exon coverage was also compared for each of the 508 exon regions in 55 genes reported by the test. The fraction of exons with relative exon level coverage difference between condition and reference within 2σ was 92.3-97.3% in Arm A, and 87.4-93.9% in Arm B. The data show that there was no preferential drop out of relative exon-level coverage in excess of what is expected due to random variation, and the panel was covered consistently between reference and storage conditions.

PPAs were also calculated for the SNVs and indels, i.e., 12 SNVs and 3 indels in Arm A, and 11 SNVs and 2 indels in Arm B. Three conditions showed variant call concordant PPA of 93.3%-100% in Arm A and 92.3% -100% in Arm B. PPA above LoD were all 100% for all conditions in Arm A and Arm B.

Together, these results demonstrated that cfDNA was stable at -20°C \pm 5°C for one year and 5 freeze/thaw cycles and 2-8°C for 24 hours. The stability of the stopping point in the workflow for storage of cfDNA at 2-8°C for 24 hours post-extraction pre-quantification was also established.

e. Intermediate Product Stability

To define the storage conditions and evaluate the stability of intermediate products, i.e., library plate, enriched library plate, and sequencing pool, used for repeat testing in the Guardant360 CDx workflow, stability at defined temperatures and durations was assessed. Samples were stored across all conditions (-20°C ± 5°C for 13, 15, or 22 days; or 2-8°C for 31 hours) with an additional thirty (30) samples of fresh intermediate product for reference. Calls from the stored intermediate product were compared to the fresh intermediate product (i.e. the reference condition).

A total of 90 samples containing the sample pools from the precision study from three distinct cfDNA clinical sample pools were used for the study. Sixty samples were processed to test 4 intermediate stability conditions (library plate, enriched library plate, 20 pM sequencing pool, 2.2 pM sequencing pool) and stored as described in **Table 15**.

The intermediate products tested for library plate and enriched library plate were subjected to 2 freeze/thaw cycles. The 20 pM sequencing pool was subjected to 3 freeze/thaw cycles.

Each condition was tested on 3 pools in 5 replicates (3x5) for a total of 15 samples. All 4 sample intermediate product conditions resulted in a total of 60 samples (15x4) passing QC. Additionally, 30 samples from the 2 analytical precision batches (15x2) were used as reference for the analysis of this study.

Intermediate Product	Storage	Target Storage Claim	Stability Testing
Enriched Library Plate	-20°C ± 5°C	14 days (including 2 freeze/thaw cycles)	At least 15 days (including 2 freeze/thaw cycles)
Library Plate	-20°C ± 5°C	21 days (including 2 freeze/thaw cycles)	At least 22 days (including 2 freeze/thaw cycles)
20 pM Pool	-20°C ± 5°C	12 days (including 2 freeze/thaw cycles)	At least 13 days (including 2 freeze/thaw cycles)
2.2 pM Pool	2-8°C	30 hours	At least 31 hours

Table 15. Description of Intermediate Product Storage Conditions

The Qualitative Detection Rate (QDR) for a storage condition was calculated which is equivalent to PPA relative to the reference condition . QDR was defined as the number of positively detected targeted variants that were positively detected in the reference condition across eligible samples (D) divided by the total number of positively detected targeted variants tested across eligible samples (N), expressed as a percentage (100 * D/N). QDR relative to reference conditions ranged from 97.7% to 100% across all stored intermediate product conditions compared to reference conditions. NPA was calculated from all negative variant sites across the Guardant360 CDx reportable range that are not detected in the reference condition. The total number of distinct variants in the final reportable range is 46,223, representing 46,217 SNVs and indels, 2 CNAs and 4 fusions. From this list, all called variants in study samples for each of the 3 pools were removed as expected positive sites for replicates of the same pool in the remaining study conditions. NPA was greater than 99.9%.

Based on these study results, intermediate products may be stored at -20°C ± 5°C for 14 days (enriched library plate), 21 days (library plate), or 12 days (20 pM Pool). Additionally, the 2.2 pM pool intermediate product may be stored at 2-8°C for 30 hours.

6.9 General Lab Equipment and Reagent Evaluation

a. cfDNA Extraction

The performance of the cfDNA extraction from plasma samples was evaluated on the QIAsymphony SP System. A retrospective analysis of clinical whole blood samples processed on the Guardant360 LDT implementation of the Guardant360 CDx device system (N=11,267 processed samples across 79 cancer types), including second tubes re-processed for a quality failure of the first tube or clinical need ,were evaluated to characterize the variability between instruments as well as the variability between runs on the same instrument. The variation in QiaSymphony instrument and/or reagent lot explained <2.1% of variance in cfDNA extraction yield. Each combination of QIAsymphony reagent kits (N=4) / instruments (N=7) resulted in successful extraction of \geq 5ng cfDNA at a rate \geq 94%, with a total success rate of 97.3%.

b. Other Instruments and Reagents

The other general lab instrument/reagent systems (4200 TapeStation, Microlab STAR, Microlab STARlet, NextSeq 550 Sequencing, and Veriti 96-Well Thermal Cycler) were assessed in combination in the precision study. Instruments and reagents varied in 3 precision combinations. Three sample pools were created at 5ng cfDNA inputs. Ten replicates per pool were tested for each of three precision combinations for a total of 6 batches sequenced on 12 flowcells. All 90 study samples passed respective QC metrics and were included in the final analysis.

Acceptable alteration PPA and NPA results were demonstrated across instruments (**Tables 16**). Acceptable sequencing QC parameters were demonstrated across precision combinations (**Table 17**).

Instrument #	PPA	95% CI	NPA	95% CI
1	98.1% (210/214)	[95.3%, 99.5%]	100% (40/40)	[91.2%, 100%]
2	98.1% (52/53)	[89.9%, 100%]	100% (10/10)	[69.2%, 100%]
3	98.1% (156/159)	[94.6%, 99.6%]	100% (30/30)	[88.4%, 100%]
4	96.3% (52/54)	[87.3%, 99.5%]	100% (10/10)	[69.2%, 100%]

Table 16. Sequencer PPA and NPA Across Precision Combinations

Table 17. Sequencing Flowcell Level QC Parameters Across Precision Combinations

QC Parameters (threshold)	Mean	SD	CV%
Cluster Density (≥170000, ≤ 280000)	223,333	9610	4.3
Percentage of Clusters Passing Filter	89.1	1.2	1.3
Quality Score (Q30) in read 1 (≥70.0)	89.1	0.7	0.8
Quality Score (Q30) in read 2 (≥70.0)	87.0	0.8	0.9
Quality Score (Q30) in index (≥70.0)	95.3	0.4	0.5
Prephasing index (≤0.01)	0	0	N/A
Prephasing 1 (≤0.01)	0.0012	0.00008	6.9
Prephasing 2 (≤0.01)	0.0014	0.00005	3.8
Phasing index (≤0.01)	0	0	N/A
Phasing 1 (≤0.01)	0.0014	0.00022	14.9
Phasing 2 (≤0.01)	0.0017	0.00018	10.5

In conclusion, the critical general lab instruments and reagents demonstrated acceptable performance for use with the Guardant360 CDx test.

6.10 Pan-Cancer Analysis

Guardant360 CDx performance characteristics were established using cfDNA derived from a wide range of cancer types. In total, 929 patient samples representing 20 cancer categories were included across the analytical validation studies performed for Guardant360 CDx.

cfDNA fragment size distributions were compared across samples from multiple cancer types. For this analysis, clinical samples were selected from analytical validation studies representing 8 different cancer types: NSCLC, breast, colorectal cancer (CRC), prostate, and uterine. The electropherograms of cfDNA post-extraction from plasma on the TapeStation show a mono-nucleosomal peak that is consistent across cancer types and with published literature. Based on these observations, cfDNA fragment size distributions are similar across cancer types and would generate qualitatively similar inputs into the assay workflow.

To further understand the performance of the Guardant360 CDx across cancer types, presequencing quality metrics (cfDNA extraction and library enrichment), post-sequencing quality metrics (non-singleton coverage, in-process contamination, coverage exceptions, GC bias, and on target rate), as well as the clinically relevant metrics of overall QC success rate and detectable levels of tumor shedding (as measured by the maximum allelic fraction of detected somatic variants) across samples tested with Guardant360 CDx candidate assay implemented in Guardant's CLIA laboratory as an LDT test were analyzed. The Guardant360 LDT assay in this analysis refers to an LDT implementation of the CDx utilizing the exact configuration. This test has been operated in the Guardant Health Clinical Laboratory to process over 10,000 clinical samples. The quality thresholds are equivalent between both versions with the exception of an additional 5 ng minimum input amount requirement for Guardant360 CDx and an upper limit to the cluster density per flowcell. These additional requirements were applied retrospectively to the Guardant360 LDT results to infer success rates for Guardant360 CDx (note that a single flowcell, out of 640, fails the upper limit of cluster density for the Guardant360 CDx).

The pan-cancer analysis evaluated 11,097 samples processed across 23 cancer categories. For each cancer category, quality pass rates were measured, and the overall patient success rate was >98% for all cancer categories. The frequency of failures for each of the individual metrics was similar across cancer types (**Table 18**).

Category Data		Sample Pr	eparation Q Pass		Patient Sar		nple Sequencing QC ss (median value)		Patient Outcome Metrics		
Cancer Category	Total Patients	First Tube Success	cfDNA Ex. Sample QC Pass %	Library Enrich. Sample QC Pass %	In process Contam- ination %	Coverage Exception	GC Bias	Non- singleton Coverage	On Target Rate	Overall Sample Pass Rate	Maximum MAF: median (standard deviation)
Breast	1516	95.2	96.6	99.1	100 (0.01)	99.2 (0.0)	99.7 (1.36)	99.8 (2766)	99.3 (88.04)	99.9	2.9 (17.5)
CUP	258	95.0	98.8	99.2	100 (0.01)	96.9 (0.0)	99.2 (1.38)	99.2 (2981)	98.4 (88.63)	100	4.9 (19.7)
Cholangio- carcinoma	302	96.0	98.6	99.3	99.7 (0.01)	99.0 (0.0)	99.3 (1.45)	100 (2911)	99.3 (88.95)	100	1.2 (13.5)
Colorectal	1041	96.5	98.8	99.5	100 (0.01)	97.8 (0.0)	98.7 (1.36)	99.8 (2832)	99.3 (88.33)	100	5.3 (21.1)
Gastroeso- phageal	443	96.2	99.0	100	100 (0.01)	98.2 (0.0)	98.4 (1.37)	100 (2790)	99.7 (88.34)	100	3.1 (17.7)
Gyneco- logical	322	95.4	98.0	99.7	100 (0.01)	97.5 (0.0)	98.7 (1.30)	100 (2771)	99.7 (88.15)	99.1	3.1 (18.5)
Head and Neck	98	94.9	96.7	100	99.0 (0.01)	99.0 (0.0)	100 (1.23)	99.0 (2399)	100 (87.85)	100	2.8 (17.0)
Liver	67	91.0	100	100	100 (0.01)	97.0 (0.0)	100 (1.50)	98.5 (2880)	97.0 (88.68)	100	1.2 (16.5)
Lung Squamous Cell Carcinoma	584	97.6	98.2	99.6	100 (0.01)	99.8 (0.0)	100 (1.27)	100 (2812)	99.7 (88.31)	100	2.2 (14.7)
Lung cancer, NOS	152	93.4	95.6	100	100 (0.01)	98.7 (0.0)	98.7 (1.39)	100 (2837)	99.3 (88.01)	99.3	4.1 (19.1)
Melanoma	174	90.8	90.4	99.4	100 (0.01)	99.4 (0.0)	100 (1.25)	100 (2439)	100 (87.90)	98.8	1.3 (15.3)
Mesoth- elioma	12	100	100	100	100 (0.01)	100 (0.0)	100 (1.20)	100 (2968)	100 (87.72)	100	0.3 (2.5)
NSCLC	4111	96.1	97.6	99.4	100 (0.01)	99.0 (0.0)	99.5 (1.29)	99.9 (2671)	99.4 (88.04)	99.9	1.7 (14.3)
Neuro- endocrine	100	90	93.6	98.9	100 (0.01)	98 (0.0)	100 (1.41)	100 (2758)	98 (87.91)	98	2.5 (21.7)
Other	419	95.7	97.95	99.5	100 (0.01)	97.8 (0.0)	99.3 (1.30)	99.3 (2730)	98.8 (88.11)	99.0	2.0 (17.3)

Table 18. Sample Success Rate Across 23 Cancers

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Pancreatic	581	95.9	97.6	98.5	100 (0.01)	99.0 (0.0)	100 (1.35)	100 (2843)	99.3 (88.12)	100	0.9 (13.9)
Primary CNS	47	93.6	93.3	100	100 (0.01)	100 (0.0)	100 (1.35)	100 (2431)	100 (88.28)	100	0.2 (0.3)
Prostate	770	94.9	98.0	99.3	100 (0.01)	97.53 (0.0)	99.09 (1.34)	99.9 (2706)	98.6 (88.14)	99.5	3.0 (19.6)
Renal	89	95.5	97.6	98.8	100 (0.01)	100 (0.0)	100 (1.28)	100 (2739)	98.9 (87.63)	100	0.8 (6.8)
SCLC	136	95.6	98.5	99.3	100 (0.01)	99.26 (0.0)	100 (1.34)	100 (2701)	98.5 (88.34)	100	3.0 (24.5)
Soft Tissue	91	98.9	98.9	100	100 (0.01)	100 (0.0)	100 (1.36)	100 (2844)	100 (88.26)	100	1.2 (12.8)
Thyroid	47	97.9	97.6	100	100 (0.01)	100 (0.0)	100 (1.33)	100 (2809)	100 (87.76)	100	0.5 (3.2)
Urothelial	147	99.3	99.3	100	100 (0.01)	98.64 (0.0)	98.64 (1.26)	100 (2660)	100 (87.82)	100	2.6 (15.2)

To assess the impact of cancer type on the variation of continuous QC metrics and ctDNA shedding level, the percent of variation explained by cancer type with variance component analysis was estimated. Variant component analysis was performed for cfDNA yield, enrichment molarity, GC bias, non-singleton coverage, on target rate, and maximum MAF. Cancer types explained no more than 2.9% of the variance across all metrics tested, including factors linked to assay sensitivity such as cfDNA yields, depth of coverage after library preparation and sequencing, and the levels of ctDNA shedding.

ctDNA shedding levels are shown below (**Figure 1**) by cancer type. Maximum MAF served as a proxy for ctDNA shedding, and maximum MAF ranges were similar for all cancer types, except primary CNS tumors. The difference in ctDNA shedding rated may be explained by CNS tumors being located behind the blood-brain barrier, which impairs the transfer of ctDNA from the CNS to the periphery, with a concomitant decrease in typical ctDNA level and detection rate. ctDNA detection is high in NSCLC and CRC, in which the most common genomic alterations are represented on the Guardant360 CDx panel; however, ctDNA detection rates are lower in mesothelioma and renal cell carcinoma, as mutations in the Guardant360 CDx reportable range are less common in these tumor types, resulting in lower ctDNA detection rate.

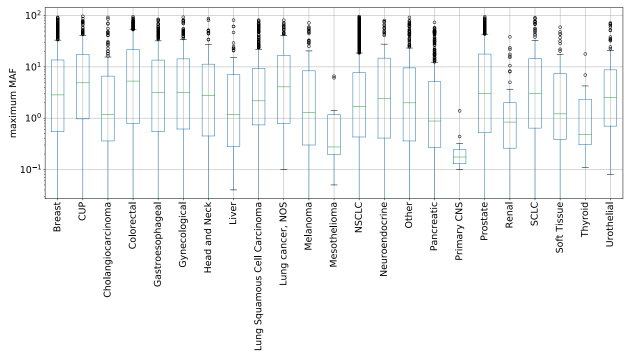


Figure 1. Maximum MAF Distribution by Cancer Type

In addition to these QC metrics, cfDNA fragment distributions in a large cohort of clinical patient samples was examined to demonstrate similarity of profiles across cancer types. Similar to other QC metrics, cancer type explained less than 1% of the variance in the locations of the cfDNA fragment size profile peak.

6.11 Concordance - Guardant360 CDx Comparison to Guardant360 LDT

A study was performed to establish the concordance between Guardant360 CDx and Guardant360 LDT. The purpose of this study was to compare the Guardant360 CDx against a Guardant360 LDT configuration used to generate historical data and is intended to support the use of those results as representative of Guardant360 CDx results.

The design and composition of these two devices is similar, as they share the same principles of operation. The primary differences in design are the panel with which the device is operated. The Guardant360 LDT version used for data generation in support of concordance to the for Guardant360 CDx test in this study was operated with version 2.10 of the panel, which covers 73 genes. The Guardant CDx is operated with version 2.11 of the panel, which covers 74 genes. While the Guardant360 CDx can detect alterations in 74 genes, it only reports select SNVs and indels in 55 genes, CNAs in two (2) genes, and fusions in four (4) genes. The concordance analysis between the Guardant360 CDx and the Guardant360 LDT is limited to 55 gene restricted reportable range. This concordance analysis utilized the bioinformatics pipeline software corresponding to each assay version.

This study evaluated a set of 258 samples with alterations in genes interrogated by both assays, after removing 2 samples that failed QC metrics. The study included cfDNA derived

from 22 cancer types, comprising two distinct sample sets. The first set was selected consecutively from among samples from patients with NSCLC positive for Guardant360 CDx variants according to Guardant360 LDT variant calling rules, targeting to obtain a minimum of 50 valid sample results for *EGFR* L858R, 50 for *EGFR* exon 19 deletions, and 75 for *EGFR* T790M mutation. The second set was selected consecutively without consideration for tumor type or previous testing results. Per the study protocol samples with specific set of rare variants were excluded from the study. "Rare" here was defined by Guardant Health as <1% prevalence or to rare fusion events (e.g. *NTRK1, ROS1*), and *MET* exon 14 skipping variants. In addition, when known to Guardant Health based on prior LDT testing or pathology reports, samples from patients for whom tumors are considered tumor mutational burden (TMB) high, microsatellite instability high (MSI-H), or PD-L1 positive were also excluded. In total, only 1 sample was excluded, as it contained an *ALK* fusion.

The cancer types represented in this concordance study were obtained from patients with NSCLC (195), gastrointestinal tumors (22), genitourinary tumors (20), breast cancer (14), gynecological tumors (4), and other solid tumors (4).

PPA and NPA between Guardant360 CDx and Guardant360 LDT, using the Guardant360 LDT assay as the reference method, was calculated for all alterations. A total of 279 SNVs, 117 indels, and 23 CNAs met the alteration inclusion criteria. A summary of PPA and NPA is provided in **Table 19**. PPA for the CDx variants as well as panel-wide SNVs, indels, and clinically significant variants showed was above 94% in all cases, whereas positive agreement levels were low for *ERBB2* and *MET* amplifications. Agreement levels were low for *ERBB2* and *MET* amplification levels for 70% of samples tested were near the decision boundary (< 1.5x LoD). High NPA was observed in all classes.

Concordance between the Guardant360 CDx and the Guardant360 LDT for the four fusions reported by the Guardant360 CDx (*ROS1, ALK, NTRK1, and RET*) is unknown as it was not evaluated.

Alteration Type	CDx+ LDT+	CDx- LDT+	CDx+ LDT–	CDx- LDT-	PPA (95% CI)	NPA (95% CI)
EGFR T790M	87	4	5	99	95.6% (89.1%, 98.8%)	95.2% (89.1%, 98.4%)
EGFR L858R	52	1	4	138	98.1% (89.9%, 100%)	97.2% (92.9%, 99.2%)
EGFR exon 19 deletions	89	3	2	101	96.7% (90.8%, 99.3%)	98.1% (93.2%, 99.8%)
Clinically Significant	282	16	14	97498	94.6% (91.4%,96.9%)	99.98% (99.97%,99.99%)

Table 19. Summary of Concordance Between Guardant360 CDx and Guardant360LDT

Panel-Wide SNV	242	15	21	105647	94.2% (90.6%,96.7%)	99.98% (99.97%,99.99%)
Panel-Wide Indel	102	5	7	50768	95.3% (89.4%,98.5%)	99.99% (99.97%,99.99%)
MET CNA	12	4	0	242	75.0% (47.6%,92.7%)	100% (98.49%,100%)
ERBB2 CNA	5	2	0	251	71.4% (29.04%,96.33%)	100% (98.54%,100%)

The concordance study also compared the Guardant360 CDx to the Guardant360 LDT which was also used in the FLAURA and AURA3 clinical studies to support the EGFR CDx indication.

The concordance analysis presented below in **Table 20** is for the EGFR CDx variants in NSCLC patient samples only (195 out of 258). Concordance analyses between the Guardant360 CDx and Guardant360 LDT utilized the bioinformatics pipeline software corresponding to the Guardant360 CDx applied to the Guardant360 LDT results.

Table 20. Summary of Concordance Between Guardant360 CDx and Guardant360LDT

Alteration Type	CDx+ LDT+	CDx- LDT+	CDx+ LDT-	CDx- LDT-	PPA (95% CI)	NPA (95% CI)
EGFR T790M	87	4	5	99	95.6% (89.1%, 98.8%)	95.2% (89.1%, 98.4%)
EGFR L858R	52	1	4	138	98.1% (89.9%, 100%)	97.2% (92.9%, 99.2%)
<i>EGFR</i> exon 19 deletions	89	3	2	101	96.7% (90.8%, 99.3%)	98.1% (93.2%, 99.8%)

In addition to the concordance study described above, the analytical performance with regards to LoD and precision was found to be comparable between the Guardant 360 CDx and the Guardant360 LDT with regards to the *EGFR* CDx variants.

6.12 Additional Studies

a. Blood Collection Tube Concordance

The purpose of this study was to establish concordance between the Streck Cell-Free DNA BCTs and BCTs used in the clinical trials (hereafter referred to as BCT-CTA) to

enable use of Guardant360 CDx data generated from the FLAURA and AURA3 clinical trials (refer to Section 7 below).

Blood from NSCLC Stage III or IV patients, prescreened externally for CDx positive and negative markers *EGFR* L858R, *EGFR* T790M, *EGFR* exon 19 deletions), were collected by utilizing two BCT-CTAs and two Streck Cell-Free DNA BCTs. The second BCT-CTA was not processed for this study. A total of 59 patients were enrolled, some with and others without CDx variants, and whole blood samples were tested from three tubes, two Streck Cell-Free DNA BCTs and one BCT-CTA.

The performance of BCT-CTAs relative to Streck Cell-Free DNA BCTs was evaluated through a call agreement analysis which tests the difference of the PPA of Streck Plasma Aliquot 2 (S2) to Streck Plasma Aliquot 1 (S1) and the PPA of BCT-CTA Plasma Aliquot 1 (C1) to S1 (difference denoted as Δ PPA1). Δ PPA2 is calculated similarly except that S2 is considered the reference instead of S1. For negative agreement, Δ NPA1 and Δ NPA2 are also calculated in a similar fashion.

Of the one-hundred and seventy-seven (177) aliquots (59 samples across 3 tube designations), 176 (99.4%) passed in-process and post-sequencing QC metrics. Of the 176 passing post-sequencing metrics, 2 failed sample QC, leaving 174 of 177 (98.3%) samples passing QC metrics. Three of the 59 patients with S1, S2, and C1 runs were excluded from call concordance analyses because of QC failures of at least one of 3 replicates.

In total 56 patients met study criteria for inclusion, including 26 distinct CDx variants observed in at least one tube. The PPA and NPA values across the entire set of CDx variants (aggregated), and for each CDx variants were calculated. BCT-CTAs and Streck Cell-Free DNA BCTs demonstrated expected levels of positive agreement, PPA 92 % – 95.5 % for CDx variants. Discordant detection was observed below LoD, with agreement above LoD being 100%. BCT-CTAs and Streck tubes demonstrated expected levels of negative agreement, NPA 97.3%– 100 % for CDx variants. The delta PPA and delta NPA values were within acceptable limits.

7 Summary of Primary Clinical Studies

Guardant Health, Inc. performed two clinical bridging studies to establish the safety and effectiveness of Guardant360 CDx to select patients with NSCLC whose tumors have *EGFR* exon 19 deletions, L858R mutations, and/or T790M mutations for osimertinib (TAGRISSO®, AstraZeneca) therapy. In the first study, pre-treatment plasma samples and clinical outcome data from patients randomized in the AstraZeneca FLAURA clinical study (NCT02296125) were used to support the safety and effectiveness of Guardant360 CDx to aid in the selection of previously untreated metastatic NSCLC patients with *EGFR* exon 19 deletions or L858R mutations for TAGRISSO therapy. In the second study, pretreatment plasma samples and clinical outcome data from the AstraZeneca AURA3 clinical study (NCT02151981) were used to assess the safety and effectiveness of the Guardant360 CDx

to aid in identifying NSCLC patients whose disease has progressed on or after *EGFR* tyrosine kinase inhibitor (TKI) therapy and who may be eligible for TAGRISSO therapy based on a *EGFR* T790M mutation-detected result.

7.1 Guardant360 CDx Clinical Bridging Study for *EGFR* Exon 19 Deletions or L858R Mutations

FLAURA Clinical Study Design

The FLAURA clinical study was a phase III, double-blind, randomized study assessing the efficacy and safety of osimertinib versus standard of care (SoC) EGFR tyrosine kinase inhibitor (TKI) therapy (gefitinib or erlotinib) in the first-line treatment of patients with locally advanced and metastatic NSCLC whose tumors have *EGFR* exon 19 deletions or exon 21 L858R mutations. Patients were enrolled based on the presence of *EGFR* exon 19 deletions or exon 21 L858R mutations in their tumor as determined by the cobas[®] EGFR Mutation Test at a central laboratory or testing at a CLIA-certified or accredited laboratory. This clinical study was used to support the approval of TAGRISSO under NDA 208065 Supplement 8.

Guardant360 CDx EGFR Exon 19 Deletions or L858R Mutations Bridging Study Design

Pre-treatment blood samples and clinical outcome data from patients positive for *EGFR* mutations by tissue testing randomized in the FLAURA clinical study were used to assess the safety and effectiveness of Guardant360 CDx for the selection of previously untreated metastatic NSCLC patients with *EGFR* exon 19 deletions or L858R mutations for TAGRISSO therapy.

Pretreatment plasma samples from 189 FLAURA patients (34% of the randomized population) were tested with Guardant360 LDT as part of an exploratory analysis. This Guardant360 LDT testing took place before the diagnostic clinical bridging study was initiated.

All patient samples would ideally have been tested using Guardant360 CDx for this diagnostic study's efficacy analysis. However, pre-treatment plasma samples were only available for the 252 patients (45% of the randomized population) not previously tested with Guardant360 LDT.

The use of this population alone in the diagnostic study was not feasible due to the bias introduced by selection of patients for exploratory testing. Specifically, patients selected for exploratory testing using Guardant360 LDT were those who had progressed and/or discontinued treatment at the time of sample selection for testing, which created a selection bias that is expected to result in longer PFS in patients tested with Guardant360 CDx relative to those tested with Guardant360 LDT and, therefore, relative to the FLAURA randomized population as a whole.

In order to minimize this selection bias, the diagnostic study primary objective analysis includes all FLAURA patients with pretreatment plasma available for testing using Guardant360 CDx, supplemented by patients for whom data was previously generated on Guardant360 LDT. This combined patient group is expected to represent the full

randomized patient population in a more robust manner. The analytical concordance study described above, supplemented by demonstration of the comparability of key performance characteristics, i.e., LoD and precision between the Guardant360 CDx and LDT, was performed to support the validity of combining data generated on Guardant360 CDx and LDT test versions for the detection of *EGFR* Exon 19 deletions or L858R mutations (Refer to Section 6.10.a. Guardant360 CDx-LDT Concordance Study results). The potential impact of the discordance observed from these studies on the effectiveness of the device was further evaluated through sensitivity analyses (see below). Further a blood collection concordance study establishing the concordance between samples collected in Streck Cell-Free DNA BCTs and the BCT-CTAs was conducted to support the validity of the data generated by testing samples collected in BCT-CTAs (Refer to Section 6.12.a).

No plasma from FLAURA patients negative for *EGFR* mutations by tissue testing was available to represent the Guardant360-positive, tissue-negative portion of the Guardant360-positive intended use population. As such, supplemental matched tissue and plasma samples from the <u>N</u>oninvasive vs. <u>Invasive Lung Evaluation clinical study</u> (the NILE study, NCT03615443) were used to estimate the prevalence of patients positive for *EGFR* exon 19 deletions or L858R mutations by Guardant360 but negative by tissue testing to evaluate the potential impact of this population on clinical efficacy.

- a. Bridging Study Inclusion and Exclusion Criteria
 - Inclusion Criteria for plasma samples from the FLAURA clinical study
 - Patient screened for the FLAURA clinical study with documented informed consent for blood sample use for diagnostic development
 - Pre-treatment time point plasma sample available for testing using Guardant360
 - Exclusion Criteria for plasma samples from the FLAURA clinical study
 - Absence of plasma for testing on Guardant360
 - Informed consent withdrawn
 - China mainland patients
 - Inclusion Criteria for samples from the NILE clinical study
 - Patient enrolled in the NILE clinical study with documented informed consent
 - Pre-treatment plasma sample available for testing with Guardant360 CDx
 - Availability of unstained slides and/or a tissue block of formalin-fixed paraffin-embedded tissue with sufficient tumor content and quantity for testing as defined by the central testing laboratory requirements for cobas[®] EGFR Mutation Test testing. Tumor tissue must be from the same disease process as the NILE study plasma sample
 - Exclusion Criteria for samples from the NILE clinical study
 - Absence of available plasma or tissue for Guardant360 CDx and cobas[®] EGFR Mutation Test testing, respectively

• Informed consent withdrawn

b. Follow-up Schedule

The Guardant360 CDx *EGFR* exon 19 deletions or L858R mutations bridging study involved only retrospective testing of plasma samples; as such, no additional patient follow-up was conducted.

c. Clinical Endpoints

The clinical endpoint used to assess osimertinib efficacy in the FLAURA clinical study primary objective was investigator-assessed progression-free survival (PFS), which was defined as the time interval between randomization and the first RECIST progression or mortality event. The Guardant360 CDx *EGFR* exon 19 deletions or L858R mutations bridging study uses the same clinical endpoint for its primary objective.

• Diagnostic Objective and Endpoint

The primary objective of the diagnostic study was to demonstrate the safety and effectiveness of the Guardant360 CDx for the selection of metastatic NSCLC patients with *EGFR* exon 19 deletions or L858R mutations for treatment with TAGRISSO. This objective was assessed by comparing the efficacy, PFS to RECIST v1.1 by investigator assessment, of single-agent TAGRISSO compared with SoC EGFR TKI therapy in the tissue-positive, Guardant360 CDx-positive patients enrolled in FLAURA.

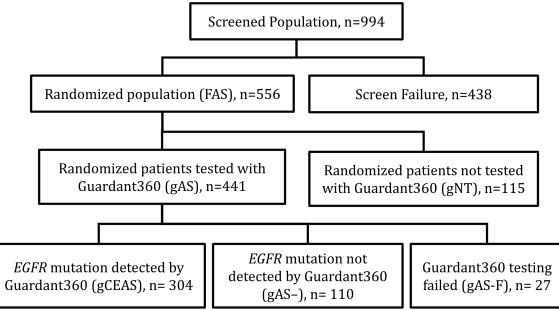
The possible influence of tissue-negative Guardant360 CDx-positive patients in the effectiveness of the Guardant360 CDx was assessed through a sensitivity analysis. As no plasma samples from FLAURA patients negative for *EGFR* mutations by tissue testing were available to represent the Guardant360 CDx-positive, tissue-negative portion of the Guardant360 CDx-positive intended use population, samples from the NILE clinical study were tested with Guardant360 CDx and the cobas® EGFR Mutation Test using tissue to calculate the NPA for the sensitivity analysis to evaluate the potential impact of this hypothetical population on clinical efficacy. The sensitivity analysis was performed using data generated by analyzing supplemental tissue samples from the NILE clinical study using the cobas® EGFR Mutation Test and by analyzing residual plasma samples from those same patients using Guardant360 CDx.

Accountability of PMA Cohort

The FLAURA diagnostic study included 441 of the total 556 (79.3%) patients randomized in the FLAURA clinical study (**Figure 2**). The analysis sets comprise diagnostic data generated using Guardant360 CDx (252/441, 57.1%) supplemented by data previously generated on Guardant360 LDT (189/441, 42.9%) as described above. Hereafter, Guardant360 CDx and LDT test versions results combined are referred to as Guardant360 results.

Of these, 304 patients (54.7% of the total population) tested positive by the Guardant360 were included in the primary objective analysis set, while 110 (24.9%) tested negative, and 27 (6.1%) failed testing.

Figure 2. Guardant360 CDx *EGFR* Exon 19 Deletions or L858R Mutations Bridging Study Patient Accountability and Analysis Set Definitions



Study Population Demographics and Baseline Parameters

Demographic and baseline clinical characteristics of patients enrolled in the FLAURA clinical study (FAS) were categorized relative to the Guardant360 CDx *EGFR* exon 19 deletions or L858R mutations bridging study populations as defined by Guardant360 results (gCEAS) and assessed for treatment arm balance. As shown in **Table 21**, demographics and baseline clinical characteristics in the clinical efficacy analysis subgroups were well-balanced between treatment arms, maintaining approximately a 1:1 randomization within each group.

- Characteristic		gC	EAS	FAS		
		TAGRISSO (n=146)	EGFR TKI (gefitinib or	TAGRISSO (n=279)	EGFR TKI (gefitinib or	
			erlotinib) (n=158)		erlotinib) (n=277)	
Age (years)	Median (range)	63 (32-83)	63 (35-87)	64 (26-85)	64 (35-93)	
Age group (years),	<65	81 (55.5)	92 (58.2)	153 (54.8)	142 (52.3)	
n (%)	≥65	65 (44.5)	66 (41.8)	126 (45.2)	132 (47.7)	
Sex, n (%)	Female	95 (65.1)	103 (65.2)	178 (63.8)	172 (62.1)	
Race, n (%)	Asian	83 (56.8)	94 (59.5)	174 (62.4)	173 (62.5)	
Smoking status, n	Never	99 (67.8)	100 (63.3)	182 (65.2)	175 (63.2)	
(%)	Current	1 (0.7)	4 (2.5)	8 (2.9)	9 (3.2)	
	Former	46 (31.5)	54 (34.2)	89 (31.9)	93 (33.6)	
AJCC staging at	I-III	15 (10.3)	15 (9.5)	52 (18.6)	47 (17.0)	
diagnosis	IV	131 (89.7)	143 (90.5)	226 (81.0)	230 (83.0)	
	Unknown	0 (0)	0 (0)	1 (0.4)	0 (0)	
Overall disease	Metastatic	141 (96.6)	155 (98.1)	264 (94.6)	262 (94.6)	
classification	Locally advanced	4 (2.7)	3 (1.9)	14 (5.0)	15 (5.4)	
	Missing	1 (0.7)	0 (0)	1 (0.4)	0 (0)	
Histology type	Adenocarcinoma	137 (93.8)	145 (91.8)	246 (88.2)	251 (90.6)	
	Other	9 (6.2)	13 (8.2)	33 (11.8)	26 (9.4)	

Table 21. Clinical Effectiveness Analysis Subgroup Demographics and Baseline	
Clinical Characteristics	

Demographic and baseline clinical characteristics of patients enrolled in the FLAURA clinical study, full analysis set (FAS), were also categorized relative FLAURA patients with plasma available for testing in this diagnostic study (gAS) and those without (gNT) to evaluate comparability (**Table 22**).

Baseline clinical characteristics were well-balanced within each population by treatment arm for all demographics and baseline clinical characteristics.

Demographics and baseline clinical characteristics between gAS and gNT were well-balanced with the exception of age ≥ 65 (48.3% gAS vs. 39.1% gNT, p = 0.0791), never smoking status (62.8% gAS vs. 69.6% gNT, p = 0.1785), AJCC stage at diagnosis I-III (16.1% gAS vs. 24.3% gNT, p = 0.0354), and metastatic overall disease classification (95.5% gAS vs. 91.3% gNT, p = 0.0603).

Table 22. Comparison of Demographics and Baseline Clinical CharacteristicsBetween FLAURA Patients with Plasma Available for Testing (gAS) and ThoseWithout (gNT)

Characteristics			gAS			gNT		
		TAGRISSO (n=219)	EGFR TKI (n=222)	Total (n=441)	TAGRISSO (n=60)	EGFR TKI (n=55)	Total (n=115)	2-sided p value [a]
Age group	<65	112 (51.1)	116 (52.3)	228 (51.7)	41 (68.3)	29 (52.7)	70 (60.9)	0.0791
(years), n (%)	≥65	107 (48.9)	106 (47.7)	213 (48.3)	19 (31.7)	26 (47.3)	45 (39.1)	
Sex, n (%)	Female	137 (62.6)	142 (63.5)	279 (63.3)	41 (68.3)	30 (54.5)	71 (61.7)	0.7628
Race, n (%)	Asian	137 (62.6)	141 (63.5)	278 (63.0)	37 (61.7)	32 (58.2)	69 (60.0)	0.5117
Smoking status	Never	137 (62.6)	140 (63.1)	277 (62.8)	45 (75.0)	35 (63.6)	80 (69.6)	0.1785
	Current/ Former	82 (37.4)	82 (36.9)	164 (37.2)	15 (25.0)	20 (36.4)	35 (30.4)	
AJCC stage at	I-III	38 (17.4)	33 (14.9)	71 (16.1)	14 (23.3)	14 (25.5)	28 (24.3)	0.0354
diagnosis	IV	181 (82.6)	189 (85.1)	370 (83.9)	45 (75.0)	41 (74.5)	86 (74.8)	
	Missing	0	0	0	1 (1.7)	0	1 (0.9)	
Overall disease	Metastatic	208 (95.0)	213 (95.9)	421 (95.5)	56 (93.3)	49 (89.1)	105 (91.3)	0.0603
classification	Locally advanced	10 (4.6)	9 (4.1)	19 (4.3)	4 (6.7)	6 (10.9)	10 (8.7)	
	Missing	1 (0.5)	0	1 (0.2)	0	0	0	
Histology type Other	Adenocarci- noma	209 (95.4)	204 (91.9)	413 (93.7)	56 (93.3)	54 (98.2)	110 (95.7)	0.4185
	Other	10 (4.6)	18 (8.1)	28 (6.3)	4 (6.7)	1 (1.8)	5 (4.3)	

[a] 2-sided p-value is based on Chi-square test for the comparisons. Statistical comparison is based on nonmissing values.

Table 23 shows that demographic and baseline clinical characteristics of patients screened for the FLAURA and enrolled in the NILE clinical studies were well-balanced between the subgroups used in the supplementary Guardant360-positive, tissuenegative prevalence analysis. with the exception of race and smoking status.

Characteristic			FLAURA Patients				
		FAS	Screen Failure	Total	Patients		
		(n=556)	(n=438)	(n=994)	(n=92)		
Age Group	<65	298 (53.6)	249 (56.8)	547 (55.0)	40 (43.5)		
(years), n (%)	≥65	258 (46.4)	189 (43.2)	447 (45.0)	52 (56.5)		
Sex, n (%)	Female	350 (62.9)	228 (52.1)	578 (58.1)	57 (62.0)		
Race, n (%)	Asian	347 (62.4)	221 (50.5)	568 (57.1)	5 (5.4)		
Smoking Status	Never	357 (64.2)	251 (57.3)	608 (61.2)	21 (22.8)		
	Current	17 (3.1)	57 (13.0)	74 (7.4)	22 (23.9)		
	Former	182 (32.7)	130 (29.7)	312 (31.4)	46 (50.0)		
	Missing	0	0	0	3 (3.3)		
	I-III	99 (17.8)	0	99 (10.0)	17 (18.5)		

Table 23. Supplementary Guardant360-Positive, Tissue-Negative PrevalenceAnalysis Subgroup Demographics and Baseline Clinical Characteristics

AJCC staging at	IV	456 (82.0)	0	456 (45.9)	75 (81.5)
diagnosis	Missing	1 (0.2)	438 (100)	439 (44.2)	0
Overall disease	Metastatic	526 (94.6)	0	526 (52.9)	89 (96.7)
classification	Locally advanced	29 (5.2)	0	29 (2.9)	3 (3.3)
	Missing	1 (0.2)	438 (100)	439 (44.2)	0
Histology type	Adenocarcinoma	523 (94.1)	0	523 (52.6)	88 (95.7)
	Other	33 (5.9)	0	33 (3.3)	4 (4.3)
	Missing	0	438 (100)	438 (44.1)	0

Safety and Effectiveness Results

a. Safety Results

Data regarding the safety and efficacy of TAGRISSO therapy were presented in the original drug approval and are summarized in the drug label. Refer to the TAGRISSO label for more information. No adverse events were reported in the conduct of the diagnostic studies as these involved retrospective testing of banked specimens only.

b. Effectiveness Results

PFS in Patients Positive by Guardant360 for *EGFR* Exon 19 Deletions or L858R Mutations

The efficacy of single-agent TAGRISSO relative to EGFR TKI therapy in patients randomized in FLAURA positive for *EGFR* exon 19 deletions or L858R mutations by tissue and by Guardant360 (gCEAS) is shown in **Table 24**. The observed PFS hazard ratio (HR) of 0.41 (95% CI 0.31, 0.54) is similar to that for the full FLAURA randomized population (FAS, PFS HR 0.46, 95% CI 0.37, 0.57). The clinical efficacy observed in the tissue and plasma positive portion of the Guardant360 intended use population, gCEAS, is consistent with that in the FAS.

Kaplan-Meier analysis of PFS in the gCEAS is presented in Figure 3.

Population	Treatment	N	Number (%) of patients with events [a]	Comparison bet Hazard Ratio (95% CI)	ween treatments 2-sided p-value
	TAGRISSO	146	83 (56.8)		.0.0001
gCEAS [b]	EGFR TKI	158	132 (83.5)	0.41 (0.31, 0.54)	<0.0001
	TAGRISSO	279	136 (48.7)	0.46 (0.37, 057)	< 0.0001
FAS [b]	EGFR TKI	277	206 (74.4)		

Table 24. Investigator-Assessed PFS in the gCEAS and FAS

[a] Progression events that do not occur within 2 scheduled visits (plus visit window) of the last evaluable assessment (or randomization) are censored and therefore excluded in the number of events. Progression includes deaths in the absence of RECIST (v1.1) progression.

[b] The analysis was performed using a log rank test stratified by mutation status and race. A hazard ratio < 1 favors TAGRISSO

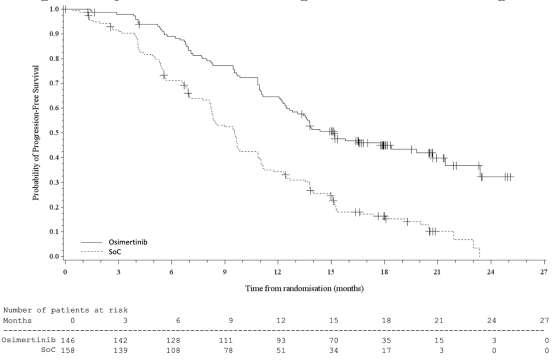


Figure 3. Kaplan-Meier Plot of Investigator-Assessed PFS for the gCEAS

Sensitivity Analysis

Imputation of Missing Guardant360 Test Results Primary Analysis for the investigatorassessed PFS

The robustness of the study conclusions was assessed by evaluating the impact of missing Guardant360 results on the effectiveness of the device. The missing Guardant360 results were imputed in the randomized (tissue positive) population using an imputation model under missing at random assumption.

There were 115 out of 556 (21%) randomized patients in FLAURA without Guardant360 test results. One of the 115 patients had missing baseline covariates and is therefore removed from the analysis as this patient's probability Guardant360 positive (G360+) could not be predicted from the selected model. Baseline covariates included in the Logit model were:

- PFS (in months, post-baseline data)
- Age group (<65 years, ≥65 years)
- Smoking status (never, current/former)
- AJCC stage at diagnosis (I-III, IV)
- Overall disease classification (Metastatic, locally advanced)
- Cobas[®] EGFR Mutation Test using plasma test result (positive, negative, failure, missing)

Results based on 1,000 imputations are presented in **Table 25** which shows robust and consistent TAGRISSO benefit in both the gCEAS defined by existing Guardant360 test results and the gCEAS (observed and imputed), in which missing Guardant360 test results were imputed via the specified Logit model. These results demonstrate that the missing data has no meaningful impact on the robustness of the efficacy result observed in the FLAURA study.

Table 25. Primary Analysis for the Investigator-Assessed PFS for the gCEAS
(observed) and gCEAS (observed and imputed)

		Comparison between treatments			
Population	Treatment	N	Number (%) of patients with events [a]	Hazard Ratio	95% Confidence Interval
gCEAS	TAGRISSO	146	83 (56.8)	0.41	0.31, 0.54
(observed)	EGFR TKI	158	132 (83.5)	0.41	0.31, 0.34
gCEAS	TAGRISSO	173	93 (53.8)	0.42	0.27.057
(observed and imputed) [b]	EGFR TKI	192	154 (80.2)	0.42	0.37, 057

[a]Log rank method with adjustment of the study stratification factors is used for the comparison between treatments.

[b] For each imputation, the analysis was performed using a log rank test stratified by mutation status and race. The average HR with 95% CI from 1,000 imputations is presented.

<u>PFS Imputation Analysis to Evaluate the Effect of Observed Guardant360 CDx-LDT</u> <u>Discordance</u>

An imputation analysis modeling the potential effect of Guardant360 CDx- Guardant360 LDT discordance on the PFS HR observed in the primary objective analysis was conducted. The sensitivity analysis by imputation analysis modelling was performed based on the NPA and PPA accounting for MAF between the Guardant360 CDx and Guardant360 LDT. The potential effect of Guardant360 CDx-Guardant360 LDT discordance on the PFS HR was calculated by the Log rank model. The identity between the observed investigator- assessed PFS HR of 0.41 (95% CI 0.31, 0.54) and the imputation results (0.40, 95% confidence 0.31, 0.54) demonstrates that the level of observed Guardant360 CDx-LDT discordance does not impact the observed results. These results support the combination of data derived from Guardant360 LDT and Guardant360 CDx for the primary objective analysis.

Sensitivity analysis for the investigator-assessed PFS in the Guardant360 positive population

A sensitivity analysis was performed by assuming a range of clinical efficacies in the Guardant360-positive, tissue-negative population (i.e. assumed HR (tissue-, G360+)), and the analysis results are presented in **Table 26**. The sensitivity analysis results

support the primary analysis results, with consistent clinical benefit, due to the high PPV of Guardant360 relative to tissue tests. The PPV calculation shown in **Table 26** for patients screened in FLAURA used a prevalence of 67%.

	Estimated P(Tissue+ Guardant360+) with 95% CI		Estimated HR (Guardant360+) with 95% CI			
	PPV Point Estimate	95% CI	Assumed HR (Tissue- and Guardant360+)	Estimated HR	95% CI	
gCEAS (observed)						
	0.99	0.97, 1.00	0.41	0.41	0.31, 0.54	
			0.50	0.41	0.31, 0.54	
			0.75	0.41	0.31, 0.54	
			1.00	0.41	0.31, 0.54	
gCEAS (observed and						
imputed)	0.99	0.97, 1.00	0.42	0.42	0.32, 0.54	
			0.50	0.42	0.32, 0.54	
			0.75	0.42	0.32, 0.54	
			1.00	0.42	0.32, 0.55	

Table 26. Sensitivity Analysis for Investigator-Assessed PFS (Guardant360 positiveirrespective of tissue result)

Log rank method with adjustment of the study stratification factors is used to estimate HR with 95% CI for the patients in the gCEAS (observed) and gCEAS (observed and imputed).

Further, because the demographic and baseline clinical characteristics of patients screened for the FLAURA and enrolled in the NILE clinical studies were not well-balanced for race and smoking status, an additional analysis was conducted to determine the minimum PPV that will lead to a unity (1.0) hazard ratio at the two-sided 95% upper confidence bound for Guardant360 positive population. Assuming fixed prevalence of the *EGFR* marker and PPA observed from the FLAURA samples, the NPA corresponding to this tipping point PPV was determined to help to address the robustness of the study results. This analysis demonstrated that NPA value corresponding to the PPV tipping point associated with an HR upper limit of the 95% CI = 1.0 was significantly less than the observed NPA of 98.7% (in **Table 28** below) supporting the robustness of the study results.

Concordance Between Guardant360 and the cobas® EGFR Mutation Test Using Tissue

Concordance between Guardant360, i.e., Guardant360 CDx and LDT test versions results combined, and the cobas[®] EGFR Mutation Test using tissue for all matched plasma-tissue from the FLAURA study is shown in **Table 27**.

EGFR Exon 19 Delet	EGFR Exon 19 Deletions cobas [®] EGFR Mutation Test Using Tissue						
	Positive	Negative	Failed	Total			
Guardant360							
Positive	185	1	2	188			
Negative	53	141	3	197			
Failed	14	12	1	27			
Total	252	154	6	412			
PPA (95% CI) [a]	77.7% [71.9%, 82.9%]						
NPA (95% CI) [a]	99.3% [96.1%, 100.0%]						
EGFR L858R Mutation		EGFR Mutation	Test Using T	<u>'issue</u>			
	Positive	Negative	Failed	Total			
Guardant360							
Positive	96	2	2	100			
Negative	40	242	3	285			
Failed	12	14	1	27			
Total	148	258	6	412			
PPA (95% CI) [a]	70.6% [62.2%, 78.1%]						
NPA (95% CI) [a]	99.2% [97.1%, 99.9%]						
EGFR Exon 19 Delet	ions or <u>cobas®</u>	EGFR Mutation	Test Using T	'issue			
L858R Mutations	Positive	Negative	Failed	Total			
Country 100	1031176	Negative	Falleu	Iotai			
Guardant360	201	2	4	207			
Positive	281	2	4	287			
Negative	93	4	1	98			
Failed	26	0	1	27			
Total	400	6	6	412			
PPA (95% CI) [a]	75.1% [70.4%, 79.4%]						
NPA (95% CI) [a]	NC						

Table 27. Concordance Between Guardant360 and the cobas® EGFR Mutation Test Using Tissue in Samples from the FLAURA Clinical Study

[a] PPA and NPA with 95% CIs are calculated based on valid test results (positive or negative). The 95% exact (Clopper-Pearson) CI is calculated. NC = not calculated

Concordance relative to Guardant360 CDx alone is similar to the concordance obtained with the Guardant360 combined data i.e., Guardant360 CDx and LDT test versions results combined. The point estimates of PPA and NPA and corresponding 95% CIs for *EGFR* Exon 19 Deletions are 73.8% (65.7%, 80.8%) and 100% (95%, 100%) respectively. The point estimates of PPA and NPA and corresponding 95% CIs for *EGFR* L858R mutations are 68.6% (56.4%,79.1%) and 98.6% (95.0%, 99.8%) respectively. The PPA for *EGFR* Exon 19 Deletions or L858R was 72.0% with a corresponding 95% CI of 65.5%, 78.0%.

As no plasma samples from FLAURA patients negative for *EGFR* mutations (Exon 19 Deletions or L858R) by tissue testing were available, NPA could not be calculated using samples from FLAURA. The NPA for *EGFR* Exon 19 Deletions or L858R relative to the

cobas[®] EGFR Mutation Test using tissue was calculated using samples from the NILE clinical study shown in **Table 28**. Of note, the single sample that tested positive for by Guardant360 CDx but negative by the cobas[®] EGFR Mutation Test using tissue comprised an uncommon *EGFR* exon 19 deletion, p.T751_I759delinsN, which is not targeted by the cobas[®] EGFR Mutation Test.

EGFR Exon 19 Deletion L858R Mutations	<u>s or</u> <u>cobas</u> ®_	cobas [®] EGFR Mutation Test Using Tissue				
	Positive	Negative	Failed	Total		
Guardant360						
Positive	14	1	0	15		
Negative	0	73	2	75		
Failed	0	2	0	2		
Total	14	76	2	92		
PPA (95% CI) [a]	100% [76.8%, 100.0%]					
NPA (95% CI) [a]	98.7% [92.7%, 100.0%]					

Table 28. Concordance Between Guardant360 and the cobas® EGFR Mutation Test
Using Tissue in Samples from the NILE Clinical Study

[a] PPA and NPA with 95% CIs are calculated based on valid test results (positive or negative). The 95% exact (Clopper-Pearson) CI is calculated.

7.2 Guardant360 CDx Clinical Bridging Study for *EGFR* T790M Mutations

AURA3 Clinical Study Design

AURA3 was a Phase III, multicenter international, open-label, randomized study to assess the efficacy and safety of TAGRISSO versus platinum-based doublet chemotherapy as second-line therapy in patients with locally advanced or metastatic *EGFR* T790M mutationpositive NSCLC, who had progressed following treatment with 1 line treatment with an approved EGFR-TKI agent. Patients were randomized in a 2:1 ratio to TAGRISSO or pemetrexed plus cisplatin / carboplatin.

Patients were enrolled based on the presence of *EGFR* T790M in their tumor as determined by the cobas[®] EGFR Mutation Test in a central laboratory. This clinical study was used to support the approval of TAGRISSO under NDA 208065 Supplement 6.

Guardant360 CDx AURA3 Bridging Study Design

Pretreatment blood samples were collected and clinical outcome data from the AURA3 clinical study were used to assess the safety and effectiveness of Guardant360 CDx for the selection of patients for TAGRISSO therapy with *EGFR* T790M mutation-positive metastatic NSCLC whose disease has progressed on or after EGFR TKI therapy.

Pretreatment samples from 287 AURA3 patients (68% of the randomized population) were tested with Guardant360 LDT in the research setting as part of an exploratory

analysis. This Guardant360 LDT testing took place before this diagnostic study was initiated.

All patient samples would ideally have been tested using Guardant360 CDx for this diagnostic study's efficacy analysis. However, pre-treatment plasma samples were available for only 265 patients (63% of the randomized population). As such, this sample set was supplemented by 35 patients for whom data was previously generated on Guardant360 LDT but for whom no plasma remains available for testing with Guardant360 CDx. The analytical concordance study described above, supplemented by demonstration of the comparability of key performance characteristics, i.e., LoD and precisions between the Guardant360 CDx and LDT, was performed to support the validity of combining data generated on Guardant360 CDx and LDT test versions for the detection of *EGFR* T790M mutation (Refer to Section 6.10.a, Guardant360CDx-LDT Concordance Study results). Further a blood collection concordance study establishing the concordance between samples collected in Streck Cell-Free DNA BCTs and the BCT-CTA was conducted to support the validity of the data generated by testing samples collected in BCT-CTA (Refer to Section 6.12.b.).

- a. Bridging Study Inclusion and Exclusion Criteria
 - Inclusion Criteria for plasma samples from the AURA3 clinical study
 - Patient screened for the AURA3 clinical study with documented informed consent for blood sample use for diagnostic development
 - Pre-treatment time point plasma sample available for testing using Guardant360
 - Exclusion Criteria for plasma samples from the AURA3 clinical study
 - Absence of plasma for testing on Guardant360
 - Informed consent withdrawn
 - China mainland patients

b. Follow-up Schedule

The Guardant360 CDx *EGFR* T790M bridging study involved only retrospective testing of plasma samples; as such, additional patient follow-up was conducted.

c. Clinical Endpoints

The clinical endpoint used to assess TAGRISSO efficacy in the AURA3 clinical study primary objective was investigator-assessed PFS, which was defined as the time interval between randomization and the first RECIST progression or mortality event. The Guardant360 CDx *EGFR* T790M bridging study uses the same clinical endpoint for its primary objective.

• Diagnostic Objective and Endpoint

The primary objective of the study was to demonstrate the safety and effectiveness of Guardant360 CDx for the selection of NSCLC patients who have progressed on or after EGFR TKI therapy with *EGFR* T790M mutations for treatment with TAGRISSO. This objective was assessed by comparing the efficacy as determined by PFS to RECIST v1.1

by investigator assessment of single-agent TAGRISSO compared with chemotherapy in the tissue-positive, Guardant360 CDx-positive patients enrolled in AURA3.

The possible influence of tissue-negative Guardant360 CDx-positive patients in the effectiveness of the Guardant360 CDx was assessed through sensitivity analysis based on randomly selected tissue-negative AURA3 screen-failure samples.

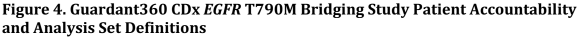
Accountability of PMA Cohort

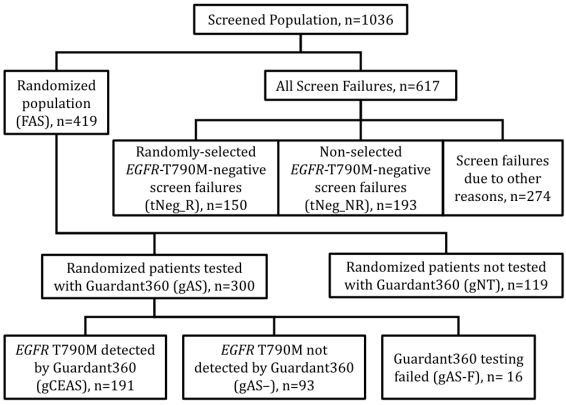
The AURA3 diagnostic study included 300 of the total 419 (71.6%) patients randomized in the AURA3 clinical study (**Figure 4**). Of these, 191 patients (45.6% of the total population) tested positive by Guardant360 and were included in the primary objective analysis set, 93 (31.0%) tested negative, and 16 (5.3%) failed testing. The analysis sets comprise diagnostic data generated using Guardant360 CDx (265/300, 88.3%) supplemented by data previously generated on Guardant360 LDT (35/300, 11.7%) as described above. Hereafter, Guardant360 CDx and LDT test versions results combined are referred to as Guardant360 results.

As AURA3 randomized patients comprised only those positive by tissue testing for *EGFR* T790M mutations, a sensitivity analysis to assess the possible influence of tissue-negative, Guardant360 plasma-positive patients was also performed using 150 randomly selected samples derived from the screened population of AURA3 that failed screening due to a negative *EGFR* T790M tissue test result (150/343, 43.7%).

Study Population Demographics and Baseline Parameters

Demographic and baseline clinical characteristics of patients enrolled in the AURA3 clinical study (FAS) were categorized relative to the Guardant360 CDx *EGFR* T790M bridging study populations as defined by Guardant360 results (gCEAS) and assessed for treatment arm balance. As shown in **Table 29**, demographics and baseline clinical characteristics in the clinical efficacy analysis subgroups were well-balanced between treatment arms, maintaining approximately a 2:1 randomization within each group.





Characteristic		gCE	AS	FAS		
		TAGRISSO (n=138)	Chemo- therapy	TAGRISSO (n=279)	Chemo- therapy	
			(n=53)		(n=140)	
Age (years)	Median (range)	61.0 (34,82)	63.0 (20,80)	62.0 (25, 85)	63.0 (20, 90)	
Age group	<65	86 (62.3)	28 (52.8)	165 (59.1)	77 (55.0)	
(years), n (%)	≥65	52 (37.7)	25 (47.2)	114 (40.9)	63 (45.0)	
Sex, n (%)	Male	50 (36.2)	13 (24.5)	107 (38.4)	43 (30.7)	
	Female	88 (63.8)	40 (75.5)	172 (61.6)	97 (69.3)	
Race, n (%)	Asian	74 (53.6)	35 (66.0)	182 (65.2)	92 (65.7)	
Smoking status, n	Never	95 (68.8)	39 (73.6)	189 (67.7)	94 (67.1)	
(%)	Current	5 (3.6)	1 (1.9)	14 (5.0)	8 (5.7)	
	Former	38 (27.5)	13 (24.5)	76 (27.22)	38 (27.1)	
AJCC staging at	I-III	20 (14.5)	10 (18.9)	52 (18.6)	31 (22.1)	
diagnosis	IV	117 (84.8)	43 (81.1)	225 (80.6)	109 (77.9)	
	Missing	1 (0.7)	0	2 (0.7)	0	
Overall disease	Metastatic	134 (97.1)	53 (100.0)	266 (95.3)	138 (98.6)	
classification	Locally advanced	4 (2.9)	0	13 (4.7)	2 (1.4)	
Histology type	Adenocarcinoma	137 (99.3)	53 (100.0)	277 (99.3)	140 (100)	
	Other	1 (0.7)	0	2 (0.7)	0	

Table 29. Baseline Demographics and Clinical Characteristics

Also, of interest in this analysis is the comparison between AURA3 patients with plasma available for testing in this diagnostic study (gAS) and those without (gNT) to evaluate comparability (**Table 30**).

Demographics and baseline clinical characteristics were well-balanced between treatment arms for both the gAS and gNT with the exception of Asian race (89.1% osimertinib vs. 65.5% chemotherapy) and sex (56.3% osimertinib vs. 70.9% chemotherapy) in the gNT. Demographics and baseline clinical characteristics between gAS and gNT were comparable, with the exception of age \geq 65 (45.0% gAS vs. 35.3% gNT, p = 0.0697), Asian race (60.3% gAS vs. 78.2% gNT, p = 0.0005), and never smoking status (65.7% gAS vs. 72.3% gNT, p = 0.1931).

Table 30. Comparison between AURA3 Patients with Plasma Available for Testing in this Diagnostic Study (gAS) and Those Without (gNT)

Characteristic		gAS			gNT			
		TAGRISSO (n=215)	Chemo- therapy (n=85)	Total (n=300)	TAGRISSO (n=64)	Chemo- therapy (n=55)	Total (n=119)	2-sided p value [a]
Age group	<65	121 (56.3)	44 (51.8)	165 (55.0)	44 (68.8)	33 (60)	77 (64.7)	0.0697
(years), n (%)	≥65	94 (43.7)	41 (48.2)	135 (45.0)	20 (31.2)	22 (40)	42 (35.3)	
Sex, n (%)	Female	136 (63.3)	58 (68.2)	194 (64.7)	36 (56.3)	39 (70.9)	75 (63.0)	0.7520

Race, n (%)	Asian	125 (58.1)	56 (65.9)	181 (60.3)	57 (89.1)	36 (65.5)	93 (78.2)	0.0005
Smoking statu	s Never	141 (65.6)	56 (65.9)	197 (65.7)	48 (75.0)	38 (69.1)	86 (72.3)	0.1931
	Current/ Former	74 (34.4)	29 (34.1)	103 (34.3)	16 (25.0)	17 (30.9)	33 (27.7)	
AJCC stage at	I-III	39 (18.1)	23 (27.1)	62 (20.7)	13 (20.3)	8 (14.5)	21 (17.6)	
diagnosis	IV	174 (80.9)	62 (72.9)	236 (78.7)	51 (79.7)	47 (85.5)	98 (82.4)	0.4657
	Missing	2 (0.9)	0 (0)	2 (0.7)	0 (0)	0 (0)	0 (0)	
Overall disease	e Metasta- tic	204 (94.9)	84 (98.8)	288 (96.0)	62 (96.9)	54 (98.2)	116 (97.5)	
	Locally advan- ced	11 (5.1)	1 (1.2)	12 (4.0)	2 (3.1)	1 (1.8)	3 (2.5)	0.5712
Histology type	Adeno- carcino-	214 (99.5)	85 (100)	299 (9.7)	64 (100)	55 (100)	119 (100)	1.000
	ma Other	1 (0.5)	0 (0)	1 (0.3)	0 (0)	0 (0)	0 (0)	

[a] 2-sided p-value is based on Chi-square test for the comparisons. Statistical comparison is based on nonmissing values.

Safety and Effectiveness Results

a. Safety

Data regarding the safety of TAGRISSO therapy were presented in the original drug approval and are summarized in the drug label. Refer to the TAGRISSO label for more information. No adverse events were reported in the conduct of the diagnostic studies as these involved retrospective testing of banked specimens only.

b. Effectiveness Results

PFS in Patients Positive by Guardant360 for EGFR T790M Mutations

The efficacy of single-agent TAGRISSO relative to chemotherapy in patients positive for *EGFR* T790M mutations by Guardant360 (gCEAS) is shown in **Table 31**. The observed PFS HR of 0.34 (95% CI 0.22, 0.53) was similar to the full AURA3 randomized population (FAS, PFS HR 0.30, 95% CI 0.23, 0.41). This demonstrates clinically relevant osimertinib efficacy in the Guardant360 intended use population.

Kaplan-Meier analysis of PFS in the gCEAS is presented in **Figure 5**.

				Comparison between treatments			
Population	Treatment	N	Number (%) of patients with events [a]	Hazard Ratio (95% CI)	2-sided p-value		
gCEAS [b]	TAGRISSO	138	85 (61.6)		0.0004		
	Chemotherapy	53	48 (90.6)	0.34 (0.22, 0.53)	<0.0001		
FAS [b]	TAGRISSO	279	140 (50.2)	0.20 (0.22, 0.41)	-0.0001		
	Chemotherapy	140	110 (78.6)	0.30 (0.23, 0.41)	<0.0001		

Table 31. Investigator-Assessed PFS in the gCEAS and FAS

[a] Progression events that do not occur within 2 scheduled visits (plus visit window) of the last evaluable assessment (or randomization) are censored and therefore excluded in the number of events. Progression includes deaths in the absence of RECIST (v1.1) progression.

[b] The analysis was performed using a log rank test stratified by race. A hazard ratio < 1 favors TAGRISSO

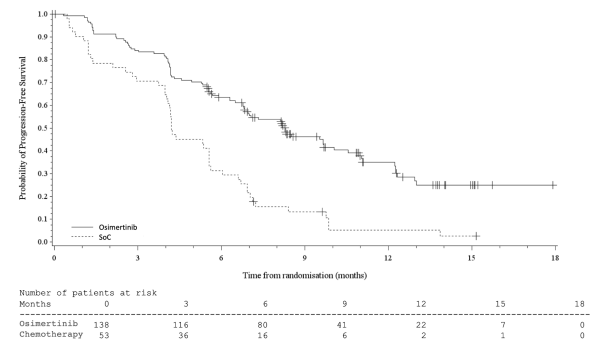


Figure 5. Kaplan-Meier Plot of Investigator-Assessed PFS for gCEAS

Sensitivity Analysis

Imputation of missing Guardant360 test results Primary analysis for the investigator-assessed PFS

The robustness of the study conclusions was assessed by evaluating the impact of missing Guardant360 results on the effectiveness of the device. The missing Guardant360 results were imputed in the randomized (tissue positive) population using an imputation model under missing at random assumption. There are 119 (300/419, 28%) randomized patients in AURA3 with missing Guardant360 test results, each of the 119 patients with missing Guardant360 test results is to be imputed via a specified Logit model. Baseline covariates included in the Logit model are:

- PFS (in months, post-baseline data)
- Age group (<65 years, ≥65 years)
- Race (Asian, Non-Asian)
- Smoking status (never, current/former)
- cobas[®] EGFR Mutation Test using plasma test result (positive, negative, failed, not tested, missing)

Results based on 1,000 imputations are presented in **Table 32** and show robust and consistent TAGRISSO benefit in the gCEAS defined by the observed Guardant360 test results and the gCEAS (observed and imputed), in which missing Guardant360 test results were imputed via the specified Logit model. The consistency of these results demonstrates that the missing G360 data have no meaningful impact on the robustness of the efficacy result observed in the AURA3 study.

Table 32. Primary analysis for the investigator-assessed PFS for the gCEAS (observed) and gCEAS (observed and imputed)

				Comparison between		
				treatr	nents	
Population	Treatment	Ν	Number (%) of	Hazard Ratio	95%	
			patients with		Confidence	
			events [a]		Interval	
gCEAS	TAGRISSO	138	85 (61.6)	0.34	0.22, 0.53	
(observed)	Chemotherapy	53	48 (90.6)	0.34	0.22, 0.33	
gCEAS	TAGRISSO	182	102 (56.0)	0.35	0.24, 0.51	
(observed	Chemotherapy	92	74 (80.4)			
1						

and

imputed) [b]

[a]Log rank method with adjustment of the study stratification factors is used for the comparison between treatments.

[b] For each imputation, the analysis was performed using a log rank test stratified by mutation status and race. The average HR with 95% CI from 1,000 imputations is presented.

PFS Imputation Analysis to Evaluate the Effect of Observed Guardant360 CDx-LDT Discordance

An imputation analysis modeling the potential effect of Guardant360 CDx-Guardant360 LDT discordance on the PFS HR observed in the primary objective analysis was conducted. The sensitivity analysis by imputation analysis modelling was performed accounting for MAF. The potential effect of Guardant360 CDx-Guardant360 LDT discordance on the PFS HR was calculated by the Log rank model. The identity between the observed investigator- assessed PFS HR of 0.34 (95% CI 0.22, 0.53) and the imputation results (0.34, 95% confidence 0.22, 0.53) demonstrates that the level of observed Guardant360 CDx-LDT discordance does not impact the observed results. These results support the combination of data derived from Guardant360 LDT and Guardant360 CDx for the primary objective analysis.

<u>Sensitivity analysis for the investigator-assessed PFS in the Guardant360</u> <u>positive population</u>

The analysis above demonstrated TAGRISSO efficacy in the Guardant360positive, tissue-positive subset of the Guardant360 CDx intended use population. As shown in **Table 33**, sensitivity analysis modeling efficacy in the entire Guardant360 CDx intended use population demonstrates robustness to the contribution of the Guardant360-positive, tissue-negative patients not represented in the AURA3 clinical study, with statistically-significant efficacy maintained across the entire Guardant360 CDx intended use population, including the modeled Guardant360-positive, tissue-negative subgroup. The PPV calculation shown in **Table 33** for the patients screened in AURA3 used a prevalence of 55%.

	Estimated P(Tissue+ Guardant360+) with 95% CI		Estimated HR (Guardant360+) with 95% CI			
	PPV Point Estimate	95% CI	Assumed HR (Tissue- and Guardant360+)	Estimated HR	95% CI	
gCEAS (observed)						
	072	0.66, 0.77	0.34	0.34	0.22, 0.53	
			0.50	0.38	0.27, 0.53	
			0.75	0.43	0.30, 0.60	
			1.00	0.46	0.33, 0.65	
gCEAS (observed +						
imputed)	0.72	0.66, 0.77	0.35	0.36	0.24, 0.51	
			0.50	0.39	0.29, 0.52	
			0.75	0.43	0.32, 0.59	
			1.00	0.47	0.35, 0.64	

Table 33. Sensitivity Analysis for Investigator-Assessed PFS (Guardant360 positive irrespective of tissue result)

Log rank method with adjustment of the study stratification factors is used to estimate HR with 95%CI for the patients in the gCEAS (observed) and gCEAS (observed + imputed).

Concordance Between Guardant360 and the cobas® EGFR Mutation Test Using Tissue

Concordance between Guardant360, i.e., Guardant360 CDx and LDT test versions results combined and the cobas[®] EGFR Mutation Test using tissue for all matched plasma-tissue samples from the AURA3 study is shown in **Table 34**.

<u>EGFR T790M</u>	cobas [®] EGFR Mutation Test Using Tissue						
	Positive	Negative	Failed	Total			
Guardant360							
Positive	190	48	0	238			
Negative	92	98	0	190			
Failed	15	4	0	19			
Total	297	150 [b]	0	447			
PPA (95% CI) [a]	67.4% [61.6 - 72.8%]						
NPA (95% CI) [a]	67.1% [58.9 – 74.7%]						

Table 34. Concordance Between Guardant360 and the cobas® EGFR Mutation Test Using Tissue

[a] PPA and NPA with 95% CIs are calculated based on valid test results (positive or negative). The 95% exact (Clopper-Pearson) CI is calculated. [b] Includes 2 patients negative for *EGFR* T790M randomized into the FAS in error.

Concordance relative to Guardant360 CDx alone is similar. The point estimates of PPA and NPA and corresponding 95% CIs for *EGFR* T790M are 66.9% (60.7%, 72.8%) and 67.1% (58.9%, 74.7%) respectively.

8 Additional Guardant360 CDx Variant Details

Table 35. Guardant360 CDx Reportable Alterations Based on cDNA and Amino Acid	l
Changes	

Gene (Transcript ID)	Reportable cDNA and Amino Acid Changes				
AKT1 (NM_001014432)	E17K, R69_C77dup				
<i>ALK</i> (NM_004304)	V1123S; T1151M; L1152P; L1152R; L1152V; C1156T; C1156Y; L1156Y; I1171N; I1171S; I1171T; F1174C; F1174L; F1174V; F1174I; F1174X; F1175C; F1175L; V1180L; L1196M; L1196Q; L1198F; G1202R; G1202del; D1203N; S1206C; S1206F; S1206Y; E1210K; D1225N; E1242K; F1245C; G1269A; R1275Q; P43A; R557C				
<i>APC</i> (NM_001127511)	c.1312+1G>A; c.1312+1G>T; c.1409-1G>A; c.1548+1G>C; c.1744-1G>A; c.532- 1G>A; c.730-1G>A; c.834+1G>A; c.834+2T>C; c.835-1G>A Y1000*; N1026S; K1030*; Y1031*; Q1045*; W1049*; I1055fs; K1061*; Q1062fs; R1066fs; S1068*; E1080*; S1104*; E1111*; R1114*; G1120E; Q1123*; N1142fs; E1149*; E1156*; E1156fs; K1165*; E1168*; Q1175*; K1182*; Y1183*; K1192*; S1196*; Q1204*; E1209*; S1213fs; Q1244*; Q1260fs; S1281*; S1282*; E1286*; I1287fs; E1288*; G1288*; G1288fs; Q1291*; Q1294*; Q1294fs; E1295*; E1295fs; A1296fs; S1298fs; T1301fs; L1302fs; Q1303*; I1304fs; E1306*; E1306fs; I1307fs; E1309*; E1309fs; K1310*; K1310fs; I1311fs; G1312*; G1312fs; R1314fs; S1315*; E1317*; P1319fs; E1322*; E1322fs; S1327*; Q1328*; R1331*;				

Gene (Transcript ID)	Reportable cDNA and Amino Acid Changes
Gene (Transcript ID)	Reportable cDNA and Amino Acid Changes R1331fs; Q1338*; Q1338fs; L1342fs; E1345*; S1346*; S1346fs; Q1349*; V1352fs; E1353*; E1353fs; S1355fs; S1356*; G1357*; Q1360*; S1364fs; G1365fs; Q1367*; K1370*; K1370fs; E1374*; Y1376*; Y1376fs; Q1378*; E1379*; M1383fs; R1386*; C1387*; S1392*; D1394fs; S1395C; F1396fs; E1397*; R1399fs; S1400L; S1400fs; A1402V; Q1406*; S1407fs; E1408*; Q1411*; S1411fs; V1414*; V1414fs; S1415fs; I1417fs; I1418fs; S1421fs; D1422fs; L1423fs; P1424fs; P1427fs; Q1429*; T1430fs; M1431fs; S1434fs; R1435fs; Q1447*; K1449*; K1449fs; R1450*; R14450fs; E1451*; V1452fs; N1455fs; A1457fs; E1461*; E1464fs; S1465fs; G1466R; Q1469fs; V1472fs; Q1477*; V1479fs; Q1480*; A1485fs; D1486fs; T1487fs; L1488fs; L1489fs; H1490fs; F1491fs; A1492fs; T1493fs; E1521*; Q1529*; E1530*; N1531fs; E1536*; E1538*; E1538fs; S1539*; E1544*; S1545*; N1546fs; E1547*; N1548fs; Q1549*; E1550*; E1576*; E1576fs; C1578fs; I1579fs; K1593fs; P1594fs; Q1621*; D1636fs; R1687*; D170fs; L1713fs; P173fs; N1792fs; R1858*, A1879fs; R1920*; A199V; H2063fs; S21*; E211*; R213*; S2140*; R216*; R250W; I2615fs; E2619*; R222*; R2237; E225*; R230C; S2307L; S2310*; R232*; G2332fs; Q236*; R2387; S2441*; Q247*; W2504*; S2555*; W2564*; R259W; I2615fs; E2619*; R2144*; R405*; L548*; L548fs; W553*; R554*; R564*; E574*; K299*; Q532*; K534*; L540*; L548fs; W553*; R55
	G721*; S747*; Q757*; Q767*; S770*; E771*; F773fs; L779*; D78fs; K782*; R786C; Q789*; Y796*; Y799fs; R805*; F814fs; L822fs; Y825fs; L826fs; P832fs; S837*; S843fs; D849fs; R854fs; E855*; E855fs; N869fs; R876*; V915fs; E918*; Y935*; Y935fs; N936fs; S940*; E941*; N942fs; S943*; C947fs; K953*; R976fs; G977fs; Q978*; E984*; E991*; K993*; Y997fs; Q999*
AR (NM_000044)	A270T; R630Q; Q641*; L702H; V716M; W742C; M750L; G796R; F814V; E873Q; H875Q; H875Y; T878A; T878S; M887I; S889G; D891H; M896V
ARAF (NM_001654)	S214A; S214C; S214F; S214Y; S214P

Gene (Transcript ID)	Reportable cDNA and Amino Acid Changes
BRAF (NM_004333)	S365L; R444W; R462E; R462I; I463S; G464V; G466V; G466A; G466E; G466R; S467L; F468C; G469A; G469E; G469L; G469V; G469R; G469S; V471F; L485F; K499E; E501K; L505H; L525R; N581H; N581S; N581T; N581Y; N581K; D587A; D587E; I592M; I592V; D594E; D594N; D594A; D594G; D594H; D594V; D594Y; F595S; G596C; G596D; G596R; G596S; G596V; L597Q; L597R; L597S; L597V; T599R; V600D; V600E; V600G; V600K; V600M; V600R; V600A; V600L; K601E; K601N; K601Q; K601R; S605N
<i>BRCA1</i> (NM_007294)	M?; M1R; S1164I; Q1395Q; L1407P; K1487R; R1495K; R1495M; R1495T; E1559K; E1559Q; M1652K; V1653M; S1655F; G1656D; L1657P; E1660G; T1685A; T1685I; H1686Q; H1686R; M1689R; M1689T; T1691I; T1691K; D1692H; D1692Y; D1692N; V1696L; C1697R; R1699L; R1699Q; R1699W; T1700A; K1702E; Y1703H; Y1703S; F1704S; L1705P; G1706E; G1706R; A1708E; A1708V; V1713A; V1714G; S1715C; S1715N; S1715R; W1718C; W1718L; W1718S; S1722F; F1734L; F1734S; V1736A; V1736D; V1736G; G1738R; G1738E; D1739E; D1739G; D1739V; D1739Y; V1741G; G1743R; H1746N; P1749R; R1751P; A1752P; A1752V; R1753T; Q1756C; F1761I; F1761S; G1763V; L1764P; I1766S; G1770V; T1773I; M1775K; M1775R; M1775E; L1780P; C1787S; G1788V; G1788D; A1789T; M18T; G1803A; I1807S; V1809F; V1810G; Q1811R; P1812A; W1815*; E1817*; A1823T; V1833E; V1833M; R1835P; E1836K; W1837C; W1837G; W1837R; V1838E; S1841A; S1841N; S1841R; A1843P; A1843T; Y1853C; L1854P; L22S; C24R; C27A; E33A; T37R; T37K; C39Y; C39R; H41R; C44Y; C44F; C44S; C47G; C61G; A622V; C64G; C64W; C64Y; R71G; R71K; R71T; C1787_G1788delinsSD
<i>BRCA2</i> (NM_000059)	M1?; A1393V; S142I; V159M; G173C; R174C; D191G; S196N; S206C; V211I; V211L; E2258K; R2336C; R2336H; R2336P; R2336L; P2532L; R2602T; W2626C; I2627F; L2647P; L2653P; R2659K; R2659T; E2663V; S2670L; I2675V; S2695L; T2722R; D2723A; D2723G; D2723H; G2748D; R2784W; N2829R; R2842C; E2918E; E3002K; P3039P; R3052W; D3095E; E3167E; E3342K
<i>CCND1</i> (NM_053056)	P287H; T286A; T286I; P287L; P287A; P287S; P287T
<i>CDK4</i> (NM_000075)	K22M; K22A; R24H; R24L; R24S; R24C
<i>CDK6</i> (NM_001259)	R87Q
<i>CDKN2A</i> (NM_058195, NM_000077)	E10*; G101W; D108G; D108H; D108N; D108V; D108Y; W110*; P114H; P114L; P114T; S12*; E120*; G125R; A128D; Y129*; W15*; G23D; R24P; E27del; V28_E33del; R29_A34del; L32_L37del; G35_A36del; G35del; A36_N39delinsD; L37_Y44delinsVR; N39_N42del; Y44*; P48L; Q50*; Q50H; M53I; R58*; V59G; A60T; E61*; G67S; E69*; E69A; N71S; D74N; D74Y; D74A; G75V; R80*; R80Q; P81L; G83V; H83Q; H83R; H83Y; H83N; D84H; D84N; D84A; D84Y; R87W; E88*; E88K; A97G; A97V; R98L; H98P
<i>CTNNB1</i> (NM_001904)	D32A; D32G; D32H; D32N; D32V; D32Y; S33A; S33C; S33F; S33P; S33T; S33Y; G34E; G34R; G34V; G34A; S37A; S37C; S37F; S37P; S37Y; T41A; T41I; T41N; S45C; S45F; S45P; S45Y; S45A

Gene (Transcript ID)	Reportable cDNA and Amino Acid Changes
<i>EGFR</i> (NM_005228)	Y1069C; R108G; R108K; E114K; R222C; S229C; R252P; T263P; A289D; A289T; A289V; R324L; R324C; E330K; V441D; V441G; R451C; S464L; G465E; G465R; K467T; I491M; I491R; S492G; S492R; P546S; D587H; P596L; G598A; G598V; C624Y; T638M; S645C; R671C; Q684H; P691S; L692F; L703P; L703V; E709A; E709G; E709K; E709Q; E709V; T710A; L718Q; L718V; G719A; G719C; G719D; G719R; G719S; S720P; A722V; F723L; G724S; T725M; V726M; Y727H; W731*; W731L; P733L; E734K; E734Q; G735S; V742A; K745R; E746G; E746K; E746Q; E746V; L747P; L747F; L747S; L747V; E749Q; A750P; A750E; T751I; S752Y; P753S; E758G; D761N; D761Y; V765A; S768I; V769M; V769L; N771D; H773L; H773Y; V774A; V774M; R776H; R776C; R776G; T783A; S784F; T785A; T790M; L792F; L792H; L792R; L792V; L792X; G796D; G796R; G796S; G796A; C797S; C797Y; C797G; C797D; C797W; Y801H; V802F; E804G; K806A; G810S; S811F; N826S; N826Y; R831H; L833V; V834L; H835L; R836C; D837N; L838P; L838V; L844V; V851I; T854S; T854A; T854I; G857E; L858R; L858M; L858Q; A859T; L861Q; L861R; L861F; L861P; A864V; A864T; E868G; H870R; A871G; E884K; Y891D
<i>ERBB2</i> (NM_004448)	E265K; G279A; G279E; S280F; S280Y; G292R; G309A; G309E; S310F; S310Y; E321G; S653C; V659E; G660D; R678W; R678Q; L726F; L726I; T733I; D739Y; G746S; L755A; L755P; L755R; L755S; L755F; L755M; L755W; L755V; V762L; V762M; I767F; I767M; D769H; D769V; D769Y; D769N; L770P; V773A; G776D; G776S; G776V; V777A; V777L; V777M; P780L; V794M; T798I; T798M; D808N; D821N; N827S; V842I; N857S; T862A; L866M; L869R; H878Y; E884K
<i>ESR1</i> (NM_001122742)	K303R; E380Q; V392I; S436P; S463P; L469V; R503W; V534E; P535H; L536H; L536P; L536R; L536Q; L536G; L536K; Y537S; Y537C; Y537D; Y537H; Y537N; D538G; D538E; T594R
FGFR1 (NM_023110)	S125L; P252T; M515V; N544K; N546D; N546K; N577K; K656N; K656E; K687E
FGFR2 (NM_000141)	D101Y; R203C; S252L; S252W; P253R; T268dup; F276C; K310R; S320C; C342Y; S354C; D374G; Y375C; C382R; C382Y; Y382H; C383Y; T524A; M536I; M537I; M538I; I547V; I548L; N549H; N549K; N550K; V564F; E565A; N638T; N639K; K658E; K658N; K659E; K659M; K659N; K660E; E731K
FGFR3 (NM_000142)	R248C; S249C; E322K; G370C; Y373C; Y375C; G380R; Y648S; K650E; K650M; K650N; K650Q; K650R; K650T; Y650F; G699C
GNA11 (NM_002067)	R183C; Q209L; Q209P
GNAQ (NM_002072)	R183Q; Q209L; Q209P; Q209R; T96S
HNF1A (NM_000545)	P291fs; G292fs
HRAS (NM_005343)	K117N; K117R; G12C; G12R; G12V; G12D; G12S; G12A; G13dup; G13R; G13V; G13C; G13D; A146T; A146V; A59G; A59T; Q61K; Q61L; Q61R; Q61H
<i>IDH1</i> (NM_005896)	R132C
<i>IDH2</i> (NM_002168)	R172G; R172K; R172M; R172S

Gene (Transcript ID)	Reportable cDNA and Amino Acid Changes
<i>KIT</i> (NM_000222)	C443Y; N463S; E490K; F504L; N505I; D52N; D52G; F522C; V530I; K550N; Y553N; Y553C; W557G; W557R; W557C; W557S; K558N; K558E; K558Q; K558P; V559C; V559D; V559G; V560D; V560G; V560A; V560E; N566D; V569G; Y570H; D572A; L576P; Y578C; Y578S; R634W; E635K; L641P; K642E; K642N; K642Q; V643A; L647P; I653T; V654A; V654E; N655K; N655S; N655T; T670E; T670I; N680K; H697Y; S709F; D716N; S746A; L783V; R804W; C809G; D816; D814V; D816F; D816H; D816V; D816Y; D816A; D816E; D816G; D816N; D820A; D820E; D820G; D820Y; D820H; D820V; D820N; S821F; N822H; N822I; N822K; N822Y; N822T; Y823D; V825A; A829P; P838L; I841V; S864F
<i>KRAS</i> (NM_004985)	G10dup; A11_G12dup; N116H; K117N; K117F; K117R; D119N; D119H; G12A; G12C; G12D; G12F; G12R; G12S; G12V; G12E; G12I; G12L; G12W; G12_G13dup; G13A; G13C; G13D; G13E; G13G; G13R; G13S; G13V; G13H; G13dup; G12_G13insAG; V14I; V14L; A146P; A146T; A146V; A146S; A18D; L19F; Q22E; Q22K; Q22R; Q22L; I24N; D33E; P34L; P34R; I36M; K5N; K5E; T50I; T58I; A59E; A59G; A59T; G60R; G60D; Q61H; Q61K; Q61L; Q61R; Q61E; Q61P; E62K; S65N; S65I; Y71H; Y71C; T74P; R97K
<i>MAP2K1</i> (NM_002755)	I111N; I111S; I111A; I111P; I111R; H119P; E120D; C121R; C121S; P124L; P124S; P124Q; G128D; G128V; E203K; V211D; L215P; P264S; N382H; F53C; F53I; F53L; F53V; F53Y; F53S; Q56P; K57N; K57E; K57T; D67N; I99T
MAP2K2 (NM_030662)	C125S; P128Q; P128R; Y134H; Y134C; V215E; F57C; F57L; F57V; Q60P
<i>MET</i> (NM_000245)	Y1003C; Y1003F; Y1003N; P1009S; D1010H; D1010N; D1010Y; Y1021C; Y1021F; Y1021N; V1070A; V1070E; V1070R; V1088A; V1088E; V1088R; V1092I; V1092L; H1094L; H1094R; H1094Y; H1106D; V1110I; V1110L; H1112Y; H1112L; H1112R; N1118Y; H1124D; M1131T; M1149T; G1163R; T1173I; G1181R; V1188L; T1191I; L1195V; L1195F; V1206L; L1213V; F1218I; V1220I; D1228H; D1228N; Y1230C; Y1230H; Y1230S; Y1230F; Y1230N; Y1235D; Y1235H; V1238I; D1246H; D1246N; D1246V; Y1248C; Y1248H; Y1248S; Y1248D; M1250T; Y1253D; Y1253H; K1262R; M1268I; M1268T
<i>MTOR</i> (NM_004958)	L1433S; K1452N; W1456G; W1456R; A1459P; L1460P; C1483F; C1483W; C1483Y; E1799K; F1888L; F1888I; F1888V; T1977K; T1977I; T1977R; E2014K; S2215F; S2215T; S2215Y; L2230V; L2427P; L2427Q; I2500F; I2500M
<i>NFE2L2</i> (NM_006164)	W24C; W24R; W24S; I28T; D29H; D29N; D29Y; L30F; L30P; G31A; G31R; G31V; V32G; R34G; R34Q; E63Q; E63V; D77G; D77H; E79D; E79K; E79Q; T80K; T80A; T80R; G81S; G81V; G81D; G81R; E82D; E82A; E82G; E82V
NRAS (NM_002524)	K117R; G12A; G12C; G12D; G12S; G12V; G12R; G12L; G13D; G13A; G13C; G13R; G13S; G13V; A146T; K170N; A18T; Q22K; D33E; K5N; T50I; T58I; A59G; A59T; G60E; Q61H; Q61K; Q61P; Q61R; Q61*; Q61E; Q61L; S65R
<i>NTRK1</i> (NM_002529)	R342Q; T434M; L564H; V573M; R583P; F589L; G595R; G595L; A608D; F646I; G667S; G667C; D679G; R692C; R692H
<i>NTRK3</i> (NM_001012338)	G623R; G696A
<i>PDGFRA</i> (NM_006206)	E229K; L275F; Y288C; V469A; V536E; V536M; Y555C; E556K; V561A; V561D; E563K; D568N; P577S; Q579R; A633T; H650Q; V658A; N659K; N659R; N659S; R748G; R841K; D842I; D842V; H845Y; D846Y; N848K; Y849C; Y849S; G853D; V859M

Gene (Transcript ID)	Reportable cDNA and Amino Acid Changes
<i>PIK3CA</i> (NM_006218)	Y1021C; Y1021H; T1025A; T1025S; D1029Y; P104L; M1043I; M1043L; M1043T; M1043V; N1044K; N1044Y; H1047L; H1047Q; H1047R; H1047Y; G1049R; G1049S; G106D; G106R; G106V; N1068Kfs; *1069fs; R108H; E110K; K111E; K111N; K111R; G118D; V344G; V344M; V344A; N345H; N345K; N345S; N345T; N345I; D350G; E365K; C378R; C378Y; R38C; R38G; R38H; R38L; R38S; E39K; E418K; C420G; C420R; P449T; E453A; E453D; E453K; E453Q; P539R; E542A; E542G; E542K; E542Q; E542V; E545A; E545D; E545G; E545K; E545Q; E545V; Q546H; Q546K; Q546L; Q546P; Q546R; Q546E; D549N; D578G; E579K; C604R; H701P; E726A; E726K; E81K; R88Q; C901F; G914R; R93Q; R93W
<i>RAF1</i> (NM_002880)	R143Q; R143W; S257L; S257W; S259A; S259F; S259P; T260R; P261L; P261R; N262K; V263A; W368S; L397M; S427G; I448V; L613V; R73Q
<i>RET</i> (NM_020975)	A373V; Y606C; C618Y; P628_L633del; P628_L633delinsH; L629_D631delinsH; C630_D631del; D631_L633delinsE; D631_L633delinsA; D631_L633delinsV; E632_L633del; E632_T636delinsSS; L730I; L730V; E732K; V738A; V778I; V804E; V804L; V804M; Y806C; Y806N; A807V; G810A; G810S; G810R; R833C; I852M; V871I; R873W; A883F; S904F; M918T; S922F; G949R; F998V;
<i>RHEB</i> (NM_005614)	Y35N; Y35C; Y35H
<i>ROS1</i> (NM_002944)	A1921G; L1951R; E1974K; V1979A; V1979M; 1981Tins; L1982F; L1982V; S1986F; S1986Y; E1990G; F1994L; M2001T; K2003I; F2004C; F2004I; F2004V; I2009L; L2028; E2020K; F2024C; F2024V; L2026M; L2026R; D2033; G2032R; D2033N; F2075C; F2075I; F2075V; V2089M; G2101A; N2112K; D2113G; R2116K; W2127*; M2128T; M2134I; L2155S; L2223*; N2224K
<i>SMAD4</i> (NM_005359)	Q245*; E330A; E330G; E330K; D351G; D351H; D351N; D351Y; P356L; P356R; P356S; G358*; R361C; R361H; R361P; R361S; R361G; G386A; G386C; G386V; Y412*; R445*; D493N; D493A; D493H; R515*; W524C; W524L; W524R; D537E; D537H; D537V
<i>SMO</i> (NM_005631)	T241M; W281L; V321A; V321M; A324T; I408V; L412F; D473H; D473N; D473Y; G497W; S533N; W535R; W535L; R562Q
<i>TERT</i> (NM_198253)	c124C>T; c146C>T; c57A>C; c45G>T; c236G>A; c124C>A; c138C>T; c 139C>T; c1G>A; c54C>A

Gene (Transcript ID)	Alteration Type	Exon	Codon		
BRAF (NM_004333)	Indel	12; 15	-		
EGFR (NM_005228)	SNV	-	436; 441; 442; 451; 464; 465; 466; 489; 491; 492; 497; 498		
EGFR (NM_005228)	Indel	18; 19; 20	-		
ERBB2 (NM_004448)	Indel	19; 20	-		
ESR1 (NM_001122742)	Indel	8; 10	-		
<i>KIT</i> (NM_000222)	Indel	All in-frame, excluding			
<i>MET</i> (NM_000245)	SNV, Indel	14	-		
<i>MET</i> (NM_000245)	SNV	19	-		
MYC (NM_002467)	SNV	-	74, 161, 251		
NFE2L2 (NM_006164)	SNV	-	24, 26, 27, 28, 29, 30, 31,32, 34, 77, 79, 80, 81, 82		
<i>PDGFRA</i> (NM_006206)	Indel	All in-frame, excluding splice site	-		
<i>PIK3CA</i> (NM_006218)	Indel	2; 8	-		
<i>ROS1</i> (NM_002944)	Indel	37	-		

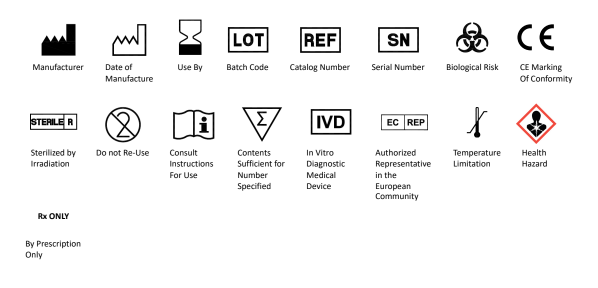
Table 36. Guardant360 CDx Reportable Alterations Based on Exons and Codons

Table 37. Guardant360 CDx Re	portable Alterations Based on Loss of Function
Tuble 57. dual dant 500 CDX Re	por table miter ations based on boss of runction

Gene (Transcript ID)	Reportable cDNA and Amino Acid Changes
BRCA1 (NM_007294)	Loss of function alterations found in all exons.
BRCA2 (NM_000059)	Loss of function alterations found in all exons.
<i>CDH1</i> (NM_004360)	Loss of function alterations found in exons 3, 8, and 9.
GATA3 (NM_001002295)	Loss of function alterations found in exons 5 and 6.
MLH1 (NM_000249)	Loss of function alterations found in exon 12.
NF1 (NM_001042492)	Loss of function alterations found in exons 11 and 29.
PTEN (NM_000314)	Loss of function alterations found in all exons.
<i>STK11</i> (NM_000455)	Loss of function alterations found in all exons.
<i>TSC1</i> (NM_000368)	Loss of function alterations found in exons 15 and 23.
VHL (NM_000551)	Loss of function alterations found in all exons.

9 Additional Information

9.1 Symbols



10 References

Meijuan Li. Statistical consideration and challenges in bridging study of personalized medicine. *J. Biopharma Stat.* (2015); 25: 397-407.

GUARDANT360 CDx INTENDED USE

Guardant360[®] CDx is a gualitative next generation sequencing-based in vitro diagnostic device that uses targeted high throughput hybridization-based capture technology for detection of single nucleotide variants (SNVs), insertions and deletions (indels) in 55 genes, copy number amplifications (CNAs) in two (2) genes, and fusions in four (4) genes. Guardant360 CDx utilizes circulating cell-free DNA (cfDNA) from plasma of peripheral whole blood collected in Streck Cell-Free DNA Blood Collection Tubes (BCTs). The test is intended to be used as a companion diagnostic to identify non-small cell lung cancer (NSCLC) patients who may benefit from treatment with the targeted therapy listed in **Table 1** in accordance with the approved therapeutic product labeling.

Table 1. Companion Diagnostic Indications

Indication	Biomarker	Therapy
Non-small cell lung cancer (NSCLC)	EGFR exon 19 deletions, L858R and T790M*	TAGRISSO [®] (osimertinib)

A negative result from a plasma specimen does not assure that the patient's tumor is negative for genomic findings. NSCLC Patients who are negative for the biomarkers listed in Table 1 should be reflexed to tissue biopsy testing for Table 1 biomarkers using an FDA-approved tumor tissue test, if feasible.

*The efficacy of TAGRISSO® (osimertinib) has not been established in the EGFR T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.

Additionally, the test is intended to provide tumor mutation profiling to be used by qualified health care professionals in accordance with professional guidelines in oncology for cancer patients with any solid malignant neoplasm. The test is for use with patients previously diagnosed with cancer and in conjunction with other laboratory and clinical findings.

Genomic findings other than those listed in **Table 1** are not prescriptive or conclusive for labeled use of any specific therapeutic product.

Guardant360 CDx is a single-site assay performed at Guardant Health, Inc.

CONTRAINDICATIONS

There are no known contraindications.

WARNINGS AND PRECAUTIONS

- Alterations reported may include somatic (not inherited) or germline (inherited) alterations. The assay filters germline variants from reporting except for pathogenic BRCA1, BRCA2, ATM, and CDK12 alterations. However, if a reported alteration is suspected to be germline, confirmatory testing should be considered in the appropriate clinical context.
- The test is not intended to replace germline testing or to provide information about cancer predisposition.
- Somatic alterations in ATM and CDK12 are not reported by the test as they are excluded from the test's reportable range.
- Genomic findings from cfDNA may originate from circulating tumor DNA (ctDNA) fragments, germline alterations, or non-tumor somatic alterations, such as clonal hematopoiesis of indeterminate potential (CHIP).
- Allow the tube to fill completely until blood stops flowing into the tube. Underfilling of tubes with less than 5 mL of blood (bottom of the label indicates 5 mL fill when tube is held vertically) may lead to incorrect analytical results or poor product performance. This tube has been designed to fill with 10 mL of blood.

LIMITATIONS

- For in vitro diagnostic use.
- For prescription use only. This test must be ordered by a gualified medical professional in accordance with clinical laboratory regulations.
- The efficacy of TAGRISSO[®] (osimertinib) has not been established in the EGFR T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.
- TAGRISSO efficacy has not been established in patients with EGFR exon 19 deletions < 0.08% MAF, in patients with EGFR L858R <0.09% MAF, and in patients with EGFR T790M < 0.03% MAF.
- The test is not intended to be used for standalone diagnostic purposes.
- The test is intended to be performed on specific serial number-controlled instruments by Guardant Health, Inc.
- A negative result for any given variant does not preclude the presence of this variant in tumor tissue.
- on patient care and treatment must be based on the independent medical judgment of the treating physician, taking into consideration all applicable information concerning the patient's condition, such as patient and family history, physical examinations, information from other diagnostic tests, and patient preferences, in accordance with the standard of care.
- ctDNA shedding rate may be lower in patients with primary central nervous system (CNS) tumors.



1.855.698.8887 client services / clientservices@guardanthealth.com / www.guardanthealth.com

GUARDANT 360°CDx Assay Specifications

Guardant360 CDx provides guideline-recommended genomic results in 7 days from a routine blood draw. Guardant360 CDx provides genomic profiling for advanced cancer patients. With demonstrated concordance to tissue in multiple prospective studies, Guardant360 CDx ensures fast and reliable results.

TEST SPECIFICATIONS*

Two 10 mL tubes of whole blood. Do not freeze or refrigerate. Ship same	Test turnaround time 7 calendar days from sample receipt to results.	P
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*Blood collection kit instructions for use

GUARDANT360 CDx PANEL

All four major classes of alterations detected

Point Mutations (SNVs) (54 Genes)					Deletion Variants (Indels) (30 Genes)			Amplifications (2 Genes)	Fusions (4 Genes)		
AKT1	ALK	APC	AR	ARAF	ATM*	BRAF	ALK	ATK1	APC	ERBB2	ALK
BRCA1**	BRCA2**	CCND1	CDH1	CDK4	CDK6	CDK12*	ATM*	BRAF	BRCA1**	MET	NTRK1
CDKN2A	CTNNB1	EGFR	ERBB2	ESR1	FGFR1	FGFR2	BRCA2**	CDH1	CDK12*		RET
FGFR3	GATA3	GNA11	GNAQ	HRAS	IDH1	IDH2	CDKN2A	EGFR	ERBB2		ROS1
KIT	KRAS	MAP2K1	MAP2K2	MET	MLH1	MTOR	ESR1	FGFR2	GATA3		1037
MYC	NF1	NFE2L2	NRAS	NTRK1	NTRK3	PDGFRA	HNF1A	HRAS	KIT		
IVIYC	INFI	INFE2L2	NRAS	NIKKI	NIRKJ	PDGFRA	KRAS	MET	MLH1		
PIK3CA	PTEN	RAF1	RET	RHEB	ROS1	SMAD4	NF1	PDGFRA	PIK3CA		
SMO	STK11	$TERT^{\dagger}$	TSC1	VHL			PTEN	RET	ROS1		
							STK11	TSC1	VHL		

NSCLC guideline-recommended genes shown in bold / † Includes TERT promoter region

*Reporting is enabled for pathogenic germline alterations only. Somatic alterations will not be reported. ** Reporting is enabled for both germline and somatic alterations.

ANALYTICAL PERFORMANCE

Alteration Type	Analytical Sensitivity*	Allelic Fr Copy N		Analytical Specificity***	Threshold for Positivity	
		5ng	30ng			
SNVs	≥ 95%	≥ 1.8%	≥ 0.2%	98.33%	≥ 0.001% MAF	*Analytical Sensitivity defined as
						the Detection Rate, that is, limit of detection (LoD)
Indels	≥ 95%	≥ 2.65%	≥ 0.2%	99.17%	≥ 0.01% MAF	**Demonstrated Allelic Fraction/Copy Number at 95% Analytical Sensitivity with 5ng and
CNAs	≥ 95%	≥ 2.3-2.4 copies	≥ 2.3-2.4 copies	99.58%	≥ 2.16-2.18 copies	30ng cfDNA input ***Analytical Specificity defined as 1 minus the per-sample false positive rate
Fusions	≥ 95%	≥ 0.7-1.5%	≥ 0.1-0.2% [†]	100%	≥ 2 unique molecules	[†] Data based on cell-line samples. See Technical Information document for further information

Actual CNA and Fusion 95% Limit of Detection for (5ng/30ng): CNAs - ERBB2 (2:3/2:3 copies), MET (2:4/2:4 copies) Fusions - NTRK1 (0:9/0.2% MAF), RET (0:72/0.1% MAF), ROS1 (1:2/0.2% MAF), ALK (1.4/0.2% MAF)







REPORTING		PHYSICIAN
Report Date:	MMM-DD-YYYY	First and Last Name
Receipt Date:	MMM-DD-YYYY	Site Name
Collection Date:	MMM-DD-YYYY	Site Address
Specimen:	Blood	Ph: (xxx) xxx-xxxx Fax: (xxx) xxx-xxxx
Status:	[Status]	Additional Recipient: First and Last Name

Companion Diagnostic

Biomarker	Status	Additional Information	
[Insert biomarker as appropiate]			
[Dynamic] *placed after EGFR exon 19 deletion, L858R, and/or T790M			

[Dynamic] *The MAF for EGFR exon 19 detection for this patient is <0.08%. Please refer below to Limitations section. [Dynamic] *The MAF for EGFR L858R for this patient is <0.09%. Please refer below to Limitations section. [Dynamic] *The MAF for EGFR T790M for this patient is <0.03%. Please refer below to Limitations section.

Other Biomarkers Identified

Results reported in this section are not prescriptive or conclusive for labeled use of any specific therapeutic product. See professional services section for additional information.

ctDNA Biomarkers with Strong Evidence of Clinical Significance in ctDNA †

Biomarker	Status	Additional Information
[Insert alteration as appropiate]		

† Please refer below to Performance Characteristics and Definitions sections for descriptions of categories.

Biomarkers with Evidence of Clinical Significance in Tissue and ctDNA †

Biomarker	Status	Additional Information	
[Insert alteration as appropiate]			

† Please refer below to Performance Characteristics and Definitions sections for descriptions of categories.

Other Biomarkers with Potential Clinical Significance

Clinical significance has not yet been established for biomarkers in this section. See the professional services section for additional information. - [Insert biomarker as appropriate]





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Table 1. Companion Diagnostic Indications

Indication	Biomarker	Therapy
Non-small cell lung cancer (NSCLC)	EGFR exon 19 deletions, L858R, and T790M*	TAGRISSO® (osimertinib)

A negative result from a plasma specimen does not assure that the patient's tumor is negative for genomic findings. NSCLC patients who are negative for the biomarkers listed in Table 1 should be reflexed to tissue biopsy testing for Table 1 biomarkers using an FDA-approved tumor tissue test, if feasible.

*The efficacy of TAGRISSO® (osimertinib) has not been established in the *EGFR* T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.

Additionally, the test is intended to provide tumor mutation profiling to be used by qualified health care professionals in accordance with professional guidelines in oncology for cancer patients with any solid malignant neoplasm. The test is for use with patients previously diagnosed with cancer and in conjunction with other laboratory and clinical findings.

Genomic findings other than those listed in Table 1 are not prescriptive or conclusive for labeled use of any specific therapeutic product.

Guardant360 CDx is a single-site assay performed at Guardant Health, Inc.

Warnings and Precautions

- Alterations reported may include somatic (not inherited) or germline (inherited) alterations. The assay filters germline variants from reporting except for pathogenic *BRCA1*, *BRCA2*, *ATM*, and *CDK12* alterations. However, if a reported alteration is suspected to be germline, confirmatory testing should be considered in the appropriate clinical context.
- The test is not intended to replace germline testing or to provide information about cancer predisposition.
- Somatic alterations in ATM and CDK12 are not reported by the test as they are excluded from the test's reportable range.
- Genomic findings from cfDNA may originate from circulating tumor DNA (ctDNA) fragments, germline alterations, or non-tumor somatic alterations, such as clonal hematopoiesis of indeterminate potential (CHIP).
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Limitations

- For in vitro diagnostic use.

- For prescription use only. This test must be ordered by a qualified medical professional in accordance with clinical laboratory regulations.
- The efficacy of TAGRISSO® (osimertinib) has not been established in the *EGFR* T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.
- TAGRISSO efficacy has not been established in patients with EGFR exon 19 deletions < 0.08% MAF, in patients with EGFR L858R < 0.09% MAF, and in patients with
- EGFR T790M < 0.03% MAF.
- The test is not intended to be used for standalone diagnostic purposes.
- The test is intended to be performed on specific serial number-controlled instruments by Guardant Health, Inc.
- A negative result for any given variant does not preclude the presence of this variant in tumor tissue.
- Decisions on patient care and treatment must be based on the independent medical judgment of the treating physician, taking into consideration all applicable information concerning the
 patient's condition, such as patient and family history, physical examinations, information from other diagnostic tests, and patient preferences, in accordance with the standard of care.
 ctDNA shedding rate may be lower in patients with primary central nervous system (CNS) tumors.

Definition of Categories

The test report includes genomic finding reported in the following categories:

Category	Prescriptive use for Therapeutic Product	Clinical Performance	Analytical Performance	Comments
<u>Category 1</u> : Companion Diagnostic (CDx)	Yes	Yes	Yes	ctDNA biomarkers linked to the safe and effective use of the corresponding therapeutic product, for which Guardant360 CDx has demonstrated clinical performance shown to support therapeutic efficacy and strong analytical performance for the biomarker.
Category 2: ctDNA Biomarkers with Strong Evidence of Clinical Significance in ctDNA	No	No	Yes	ctDNA biomarkers with strong evidence of clinical significance presented by other FDA-approved liquid biopsy companion diagnostics for which Guardant360 CDx has demonstrated analytical reliability but not clinical performance.
<u>Category 3A:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: strong analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated analytical performance including analytical accuracy, and concordance of blood-based testing to tissue-based testing for the biomarker.
<u>Category 3B:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated minimum analytical performance including analytical accuracy.
Category 4: Other Biomarkers with Potential Clinical Significance	No	No	Yes	ctDNA biomarkers with emergent evidence based on peer-reviewed publications for genes/variants in tissue, variant information from well-curated public databases, or <i>in-vitro</i> preclinical models, for which Guardant360 CDx has demonstrated minimum analytical performance.

Performance Characteristics

Please refer to product label, www.guardant360cdx.com/technicalinfo. Clinical Performance has not been established for biomarkers in categories 2, 3A, 3B, and 4. Guardant360 CDx is indicated to report the following SNVs (*AKT1, ALK, APC, AR, ARAF, ATM*, BRAF, BRCA1**,BRCA2**, CCND1, CDH1, CDK4, CDK6,CDK12*, CDKN2A, CTNNB1, EGFR, ERBB2, ESR1, FGFR1, FGFR2, FGFR3, GATA3, GNA11, GNAQ, HRAS, IDH1, IDH2, KIT, KRAS, MAP2K1, MAP2K2, MET, MLH1, MTOR, MYC, NF1, NFE2L2, NRAS, NTRK1, NTRK3, PDGFRA, PIK3CA, PTEN, RAF1, RET, RHEB, ROS1, SMAD4, SMO, STK11, TERT, TSC, VHL). Indels (<i>ALK, ATK1, APC, ATM*, BRAF, BRCA1**, BRCA2**, CDH1, CDK12*, CDKN2A, EGFR,ERBB2, ESR1, FGFR2, GATA3, HNF1A, HRAS , KIT, KRAS, MET, MLH1, NF1, PDGFRA, PIK3CA, PTEN, RET, ROS1, STK11, TSC1, VHL)*, Fusion (*ALK , NTRK1, RET, ROS1*), and Amplifications (*ERBB2, MET*).

*Reporting is enabled for pathogenic germline alterations only. Somatic alterations will not be reported. | ** Reporting is enabled for both germline and somatic alterations.

Testing performed at: Guardant Health

Laboratory Director: [First and Last Name] | CLIA ID: [XXXXXX] | CAP #: [XXXXXX] | 505 Penobscot Drive Redwood City, CA, 94063, USA



NSCLC patient with *EGFR* exon 19 deletion (alteration below lowest MAF from clinical study)



REPORTING		PHYSICIAN
Report Date:	MAR-20-2017	Dougie Houser
Receipt Date:	MAR-04-2017	Center for People Who are Sick and Want to Get Better
Collection Date:	MAR-03-2017	123 Four St., Metropolis, NY, 12345, USA
Specimen:	Blood	Ph: (808) 555-1234 Fax: N/A
Status:	FINAL	Additional Recipient: N/A

Biomarker	Status	Additional Information
EGFR exon 19 Deletions*	DETECTED	EGFR E746_A750del TAGRISSO [®] (osimertinib) is FDA approved for this indication
EGFR T790M	NOT DETECTED	
EGFR L858R	NOT DETECTED	

[Dynamic] *The MAF for EGFR exon 19 detection for this patient is <0.08%. Please refer below to Limitations section.





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Table 1. Companion Diagnostic Indications

Indication	Biomarker	Therapy
Non-small cell lung cancer (NSCLC)	EGFR exon 19 deletions, L858R, and T790M*	TAGRISSO® (osimertinib)

A negative result from a plasma specimen does not assure that the patient's tumor is negative for genomic findings. NSCLC patients who are negative for the biomarkers listed in Table 1 should be reflexed to tissue biopsy testing for Table 1 biomarkers using an FDA-approved tumor tissue test, if feasible.

*The efficacy of TAGRISSO® (osimertinib) has not been established in the *EGFR* T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.

Additionally, the test is intended to provide tumor mutation profiling to be used by qualified health care professionals in accordance with professional guidelines in oncology for cancer patients with any solid malignant neoplasm. The test is for use with patients previously diagnosed with cancer and in conjunction with other laboratory and clinical findings.

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- For in vitro diagnostic use.

- For prescription use only. This test must be ordered by a qualified medical professional in accordance with clinical laboratory regulations.
- The efficacy of TAGRISSO® (osimertinib) has not been established in the EGFR T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.
- TAGRISSO efficacy has not been established in patients with EGFR exon 19 deletions < 0.08% MAF, in patients with EGFR L858R < 0.09% MAF, and in patients with
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 ctDNA shedding rate may be lower in patients with primary central nervous system (CNS) tumors.

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<u>Category 3B:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated minimum analytical performance including analytical accuracy.
Category 4: Other Biomarkers with Potential Clinical Significance	No	No	Yes	ctDNA biomarkers with emergent evidence based on peer-reviewed publications for genes/variants in tissue, variant information from well-curated public databases, or <i>in-vitro</i> preclinical models, for which Guardant360 CDx has demonstrated minimum analytical performance.

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Please refer to product label, www.guardant360cdx.com/technicalinfo. Clinical Performance has not been established for biomarkers in categories 2, 3A, 3B, and 4. Guardant360 CDx is indicated to report the following SNVs (*AKT1, ALK, APC, AR, ARAF, ATM*, BRAF, BRCA1**,BRCA2**, CCND1, CDH1, CDK4, CDK6,CDK12*, CDKN2A, CTNNB1, EGFR, ERBB2, ESR1, FGFR1, FGFR2, FGFR3, GATA3, GNA11, GNAQ, HRAS, IDH1, IDH2, KIT, KRAS, MAP2K1, MAP2K2, MET, MLH1, MTOR, MYC, NF1, NFE2L2, NRAS, NTRK1, NTRK3, PDGFRA, PIK3CA, PTEN, RAF1, RET, RHEB, ROS1, SMAD4, SMO, STK11, TERT, TSC, VHL). Indels (<i>ALK, ATK1, APC, ATM*, BRAF, BRCA1**, BRCA2**, CDH1, CDK12*, CDKN2A, EGFR, ERBB2, ESR1, FGFR2, GATA3, KIT, KRAS, MET, MLH1, NT1, PDGFRA, PIK3CA, PTEN, RET, ROS1, STK11, TSC1, VHL)*, Fusion (*ALK , NTRK1, RET, ROS1*), and Amplifications (*ERBB2, MET*).

*Reporting is enabled for pathogenic germline alterations only. Somatic alterations will not be reported. | ** Reporting is enabled for both germline and somatic alterations.

Testing performed at: Guardant Health

Laboratory Director: [First and Last Name] | CLIA ID: [XXXXXX] | CAP #: [XXXXXX] | 505 Penobscot Drive Redwood City, CA, 94063, USA



NSCLC patient with *EGFR* L858R (alteration below lowest MAF from clinical study)



REPORTING		PHYSICIAN
Report Date:	MAR-20-2017	Dougie Houser
Receipt Date:	MAR-04-2017	Center for People Who are Sick and Want to Get Better
Collection Date:	MAR-03-2017	123 Four St., Metropolis, NY, 12345, USA
Specimen:	Blood	Ph: (808) 555-1234 Fax: N/A
Status:	FINAL	Additional Recipient: N/A

Biomarker	Status	Additional Information
EGFR L858R*	DETECTED	EGFR L858R* TAGRISSO [®] (osimertinib) is FDA approved for this indication
EGFR T790M	NOT DETECTED	
EGFR exon 19 Deletions	NOT DETECTED	

[Dynamic] *The MAF for EGFR L858R for this patient is <0.09%. Please refer below to Limitations section.





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*The efficacy of TAGRISSO® (osimertinib) has not been established in the *EGFR* T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.

Additionally, the test is intended to provide tumor mutation profiling to be used by qualified health care professionals in accordance with professional guidelines in oncology for cancer patients with any solid malignant neoplasm. The test is for use with patients previously diagnosed with cancer and in conjunction with other laboratory and clinical findings.

Genomic findings other than those listed in Table 1 are not prescriptive or conclusive for labeled use of any specific therapeutic product.

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Warnings and Precautions

- Alterations reported may include somatic (not inherited) or germline (inherited) alterations. The assay filters germline variants from reporting except for pathogenic *BRCA1*, *BRCA2*, *ATM*, and *CDK12* alterations. However, if a reported alteration is suspected to be germline, confirmatory testing should be considered in the appropriate clinical context.
- The test is not intended to replace germline testing or to provide information about cancer predisposition.
- Somatic alterations in ATM and CDK12 are not reported by the test as they are excluded from the test's reportable range.
- Genomic findings from cfDNA may originate from circulating tumor DNA (ctDNA) fragments, germline alterations, or non-tumor somatic alterations, such as clonal hematopoiesis of indeterminate potential (CHIP).
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Limitations

- For in vitro diagnostic use.

- For prescription use only. This test must be ordered by a qualified medical professional in accordance with clinical laboratory regulations.
- The efficacy of TAGRISSO® (osimertinib) has not been established in the EGFR T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.
- TAGRISSO efficacy has not been established in patients with EGFR exon 19 deletions < 0.08% MAF, in patients with EGFR L858R < 0.09% MAF, and in patients with
- EGFR T790M < 0.03% MAF.
- The test is not intended to be used for standalone diagnostic purposes.
- The test is intended to be performed on specific serial number-controlled instruments by Guardant Health, Inc.
- A negative result for any given variant does not preclude the presence of this variant in tumor tissue.
- Decisions on patient care and treatment must be based on the independent medical judgment of the treating physician, taking into consideration all applicable information concerning the
 patient's condition, such as patient and family history, physical examinations, information from other diagnostic tests, and patient preferences, in accordance with the standard of care.
 ctDNA shedding rate may be lower in patients with primary central nervous system (CNS) tumors.

Definition of Categories

The test report includes genomic finding reported in the following categories:

Category	Prescriptive use for Therapeutic Product	Clinical Performance	Analytical Performance	Comments
<u>Category 1</u> : Companion Diagnostic (CDx)	Yes	Yes	Yes	ctDNA biomarkers linked to the safe and effective use of the corresponding therapeutic product, for which Guardant360 CDx has demonstrated clinical performance shown to support therapeutic efficacy and strong analytical performance for the biomarker.
Category 2: ctDNA Biomarkers with Strong Evidence of Clinical Significance in ctDNA	No	No	Yes	ctDNA biomarkers with strong evidence of clinical significance presented by other FDA-approved liquid biopsy companion diagnostics for which Guardant360 CDx has demonstrated analytical reliability but not clinical performance.
<u>Category 3A:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: strong analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated analytical performance including analytical accuracy, and concordance of blood-based testing to tissue-based testing for the biomarker.
<u>Category 3B:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated minimum analytical performance including analytical accuracy.
Category 4: Other Biomarkers with Potential Clinical Significance	No	No	Yes	ctDNA biomarkers with emergent evidence based on peer-reviewed publications for genes/variants in tissue, variant information from well-curated public databases, or <i>in-vitro</i> preclinical models, for which Guardant360 CDx has demonstrated minimum analytical performance.

Performance Characteristics

Please refer to product label, www.guardant360cdx.com/technicalinfo. Clinical Performance has not been established for biomarkers in categories 2, 3A, 3B, and 4. Guardant360 CDx is indicated to report the following SNVs (*AKT1, ALK, APC, AR, ARAF, ATM*, BRAF, BRCA1**,BRCA2**, CCND1, CDH1, CDK4, CDK6,CDK12*, CDKN2A, CTNNB1, EGFR, ERBB2, ESR1, FGFR1, FGFR2, FGFR3, GATA3, GNA11, GNAQ, HRAS, IDH1, IDH2, KIT, KRAS, MAP2K1, MAP2K2, MET, MLH1, MTOR, MYC, NF1, NFE2L2, NRAS, NTRK1, NTRK3, PDGFRA, PIK3CA, PTEN, RAF1, RET, RHEB, ROS1, SMAD4, SMO, STK11, TERT, TSC, VHL). Indels (<i>ALK, ATK1, APC, ATM*, BRAF, BRCA1**, BRCA2**, CDH1, CDK12*, CDKN2A, EGFR, ERBB2, ESR1, FGFR2, GATA3, KIT, KRAS, MET, MLH1, NT1, PDGFRA, PIK3CA, PTEN, RET, ROS1, STK11, TSC1, VHL)*, Fusion (*ALK , NTRK1, RET, ROS1*), and Amplifications (*ERBB2, MET*).

*Reporting is enabled for pathogenic germline alterations only. Somatic alterations will not be reported. | ** Reporting is enabled for both germline and somatic alterations.

Testing performed at: Guardant Health

Laboratory Director: [First and Last Name] | CLIA ID: [XXXXXX] | CAP #: [XXXXXX] | 505 Penobscot Drive Redwood City, CA, 94063, USA



NSCLC patient with *EGFR* T790M (alteration below lowest MAF from clinical study)



REPORTING		PHYSICIAN
Report Date:	MAR-20-2017	Dougie Houser
Receipt Date:	MAR-04-2017	Center for People Who are Sick and Want to Get Better
Collection Date:	MAR-03-2017	123 Four St., Metropolis, NY, 12345, USA
Specimen:	Blood	Ph: (808) 555-1234 Fax: N/A
Status:	FINAL	Additional Recipient: N/A

Biomarker	Status	Additional Information
EGFR T790M*	DETECTED	EGFR T790M TAGRISSO [®] (osimertinib) is FDA approved for this indication
EGFR L858R	NOT DETECTED	
EGFR exon 19 Deletions	NOT DETECTED	

[Dynamic] *The MAF for EGFR T790M for this patient is <0.03%. Please refer below to Limitations section.





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Table 1. Companion Diagnostic Indications

Indication	Biomarker	Therapy
Non-small cell lung cancer (NSCLC)	EGFR exon 19 deletions, L858R, and T790M*	TAGRISSO® (osimertinib)

A negative result from a plasma specimen does not assure that the patient's tumor is negative for genomic findings. NSCLC patients who are negative for the biomarkers listed in Table 1 should be reflexed to tissue biopsy testing for Table 1 biomarkers using an FDA-approved tumor tissue test, if feasible.

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Genomic findings other than those listed in Table 1 are not prescriptive or conclusive for labeled use of any specific therapeutic product.

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- The efficacy of TAGRISSO® (osimertinib) has not been established in the EGFR T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.
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- The test is intended to be performed on specific serial number-controlled instruments by Guardant Health, Inc.
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 patient's condition, such as patient and family history, physical examinations, information from other diagnostic tests, and patient preferences, in accordance with the standard of care.
 ctDNA shedding rate may be lower in patients with primary central nervous system (CNS) tumors.

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Category 2: ctDNA Biomarkers with Strong Evidence of Clinical Significance in ctDNA	No	No	Yes	ctDNA biomarkers with strong evidence of clinical significance presented by other FDA-approved liquid biopsy companion diagnostics for which Guardant360 CDx has demonstrated analytical reliability but not clinical performance.
<u>Category 3A:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: strong analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated analytical performance including analytical accuracy, and concordance of blood-based testing to tissue-based testing for the biomarker.
<u>Category 3B:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated minimum analytical performance including analytical accuracy.
Category 4: Other Biomarkers with Potential Clinical Significance	No	No	Yes	ctDNA biomarkers with emergent evidence based on peer-reviewed publications for genes/variants in tissue, variant information from well-curated public databases, or <i>in-vitro</i> preclinical models, for which Guardant360 CDx has demonstrated minimum analytical performance.

Performance Characteristics

Please refer to product label, www.guardant360cdx.com/technicalinfo. Clinical Performance has not been established for biomarkers in categories 2, 3A, 3B, and 4. Guardant360 CDx is indicated to report the following SNVs (*AKT1, ALK, APC, AR, ARAF, ATM*, BRAF, BRCA1**,BRCA2**, CCND1, CDH1, CDK4, CDK6,CDK12*, CDKN2A, CTNNB1, EGFR, ERBB2, ESR1, FGFR1, FGFR2, FGFR3, GATA3, GNA11, GNAQ, HRAS, IDH1, IDH2, KIT, KRAS, MAP2K1, MAP2K2, MET, MLH1, MTOR, MYC, NF1, NFE2L2, NRAS, NTRK1, NTRK3, PDGFRA, PIK3CA, PTEN, RAF1, RET, RHEB, ROS1, SMAD4, SMO, STK11, TERT, TSC, VHL). Indels (<i>ALK, ATK1, APC, ATM*, BRAF, BRCA1**, BRCA2**, CDH1, CDK12*, CDKN2A, EGFR, ERBB2, ESR1, FGFR2, GATA3, KIT, KRAS, MET, MLH1, NT1, PDGFRA, PIK3CA, PTEN, RET, ROS1, STK11, TSC1, VHL)*, Fusion (*ALK , NTRK1, RET, ROS1*), and Amplifications (*ERBB2, MET*).

*Reporting is enabled for pathogenic germline alterations only. Somatic alterations will not be reported. | ** Reporting is enabled for both germline and somatic alterations.

Testing performed at: Guardant Health

Laboratory Director: [First and Last Name] | CLIA ID: [XXXXXX] | CAP #: [XXXXXX] | 505 Penobscot Drive Redwood City, CA, 94063, USA



Breast cancer patient with PIK3CA C420R



REPORTING		PHYSICIAN
Report Date:	MAR-20-2017	Dougie Houser
Receipt Date:	MAR-04-2017	Center for People Who are Sick and Want to Get Better
Collection Date:	MAR-03-2017	123 Four St., Metropolis, NY, 12345, USA
Specimen:	Blood	Ph: (808) 555-1234 Fax: N/A
Status:	FINAL	Additional Recipient: N/A

No reportable alterations with companion diagnostic (CDx) claims

Other Biomarkers Identified

Results reported in this section are not prescriptive or conclusive for labeled use of any specific therapeutic product. See professional services section for additional information.

ctDNA Biomarkers with Strong Evidence of Clinical Significance in ctDNA †

Biomarker	Status	Additional Information
PIK3CA Activating SNVs	DETECTED	<i>PIK3CA</i> C420R See professional services section for additional information

† Please refer below to Performance Characteristics and Definitions sections for descriptions of categories.





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Table 1. Companion Diagnostic Indications

Indication	Biomarker	Тherapy
Non-small cell lung cancer (NSCLC)	EGFR exon 19 deletions, L858R, and T790M*	TAGRISSO® (osimertinib)

A negative result from a plasma specimen does not assure that the patient's tumor is negative for genomic findings. NSCLC patients who are negative for the biomarkers listed in Table 1 should be reflexed to tissue biopsy testing for Table 1 biomarkers using an FDA-approved tumor tissue test, if feasible.

*The efficacy of TAGRISSO® (osimertinib) has not been established in the *EGFR* T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.

Additionally, the test is intended to provide tumor mutation profiling to be used by qualified health care professionals in accordance with professional guidelines in oncology for cancer patients with any solid malignant neoplasm. The test is for use with patients previously diagnosed with cancer and in conjunction with other laboratory and clinical findings.

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Limitations

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 ctDNA shedding rate may be lower in patients with primary central nervous system (CNS) tumors.

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<u>Category 3B:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated minimum analytical performance including analytical accuracy.
Category 4: Other Biomarkers with Potential Clinical Significance	No	No	Yes	ctDNA biomarkers with emergent evidence based on peer-reviewed publications for genes/variants in tissue, variant information from well-curated public databases, or <i>in-vitro</i> preclinical models, for which Guardant360 CDx has demonstrated minimum analytical performance.

Performance Characteristics

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Testing performed at: Guardant Health

Laboratory Director: [First and Last Name] | CLIA ID: [XXXXXX] | CAP #: [XXXXXX] | 505 Penobscot Drive Redwood City, CA, 94063, USA



CRC patient with KRAS Q61R



REPORTING		PHYSICIAN
Report Date:	MAR-20-2017	Dougie Houser
Receipt Date:	MAR-04-2017	Center for People Who are Sick and Want to Get Better
Collection Date:	MAR-03-2017	123 Four St., Metropolis, NY, 12345, USA
Specimen:	Blood	Ph: (808) 555-1234 Fax: N/A
Status:	FINAL	Additional Recipient: N/A

No reportable alterations with companion diagnostic (CDx) claims

Other Biomarkers Identified

Results reported in this section are not prescriptive or conclusive for labeled use of any specific therapeutic product. See professional services section for additional information.

Biomarkers with Evidence of Clinical Significance in Tissue and ctDNA †

Biomarker	Status	Additional Information
KRAS Activating SNVs	DETECTED	KRAS Q61R See professional services section for additional information

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Table 1. Companion Diagnostic Indications

Indication	Biomarker	Тherapy
Non-small cell lung cancer (NSCLC)	EGFR exon 19 deletions, L858R, and T790M*	TAGRISSO® (osimertinib)

A negative result from a plasma specimen does not assure that the patient's tumor is negative for genomic findings. NSCLC patients who are negative for the biomarkers listed in Table 1 should be reflexed to tissue biopsy testing for Table 1 biomarkers using an FDA-approved tumor tissue test, if feasible.

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Testing performed at: Guardant Health

Laboratory Director: [First and Last Name] | CLIA ID: [XXXXXX] | CAP #: [XXXXXX] | 505 Penobscot Drive Redwood City, CA, 94063, USA



Melamona patient with BRAF V600E



REPORTING		PHYSICIAN
Report Date:	MAR-20-2017	Dougie Houser
Receipt Date:	MAR-04-2017	Center for People Who are Sick and Want to Get Better
Collection Date:	MAR-03-2017	123 Four St., Metropolis, NY, 12345, USA
Specimen:	Blood	Ph: (808) 555-1234 Fax: N/A
Status:	FINAL	Additional Recipient: N/A

No reportable alterations with companion diagnostic (CDx) claims

Other Biomarkers Identified

Results reported in this section are not prescriptive or conclusive for labeled use of any specific therapeutic product. See professional services section for additional information.

Biomarkers with Evidence of Clinical Significance in Tissue and ctDNA †

Biomarker	Status	Additional Information
BRAF V600E	DETECTED	BRAF V600E See professional services section for additional information

† Please refer below to Performance Characteristics and Definitions sections for descriptions of categories.





Guardant360[®] CDx is a qualitative next generation sequencing-based in vitro diagostic device that uses targeted high throughput hybridization-based capture technology for detection of single nucleotide variants (SNVs), insertions and deletions (indels) in 55 genes, copy number amplifications (CNAs) in two (2) genes, and fusions in four (4) genes. Guardant360 CDx utilizes circulating cell-free DNA (cfDNA) from plasma of peripheral whole blood collected in Streck Cell-Free DNA Blood Collection Tubes (BCTs). The test is intended to be used as a companion diagnostic to identify non-small cell lung cancer (NSCLC) patients who may benefit from treatment with the targeted therapy listed in Table 1 in accordance with the approved therapeutic product labeling.

Table 1. Companion Diagnostic Indications

Indication	Biomarker	Тherapy
Non-small cell lung cancer (NSCLC)	EGFR exon 19 deletions, L858R, and T790M*	TAGRISSO® (osimertinib)

A negative result from a plasma specimen does not assure that the patient's tumor is negative for genomic findings. NSCLC patients who are negative for the biomarkers listed in Table 1 should be reflexed to tissue biopsy testing for Table 1 biomarkers using an FDA-approved tumor tissue test, if feasible.

*The efficacy of TAGRISSO® (osimertinib) has not been established in the *EGFR* T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.

Additionally, the test is intended to provide tumor mutation profiling to be used by qualified health care professionals in accordance with professional guidelines in oncology for cancer patients with any solid malignant neoplasm. The test is for use with patients previously diagnosed with cancer and in conjunction with other laboratory and clinical findings.

Genomic findings other than those listed in Table 1 are not prescriptive or conclusive for labeled use of any specific therapeutic product.

Guardant360 CDx is a single-site assay performed at Guardant Health, Inc.

Warnings and Precautions

- Alterations reported may include somatic (not inherited) or germline (inherited) alterations. The assay filters germline variants from reporting except for pathogenic *BRCA1*, *BRCA2*, *ATM*, and *CDK12* alterations. However, if a reported alteration is suspected to be germline, confirmatory testing should be considered in the appropriate clinical context.
- The test is not intended to replace germline testing or to provide information about cancer predisposition.
- Somatic alterations in ATM and CDK12 are not reported by the test as they are excluded from the test's reportable range.
- Genomic findings from cfDNA may originate from circulating tumor DNA (ctDNA) fragments, germline alterations, or non-tumor somatic alterations, such as clonal hematopoiesis of indeterminate potential (CHIP).
- Allow the tube to fill completely until blood stops flowing into the tube. Underfilling of tubes with less than 5 mL of blood (bottom of the label indicates 5 mL fill when tube is held vertically) may lead to incorrect analytical results or poor product performance. This tube has been designed to fill with 10 mL of blood.

Limitations

- For in vitro diagnostic use.

- For prescription use only. This test must be ordered by a qualified medical professional in accordance with clinical laboratory regulations.
- The efficacy of TAGRISSO® (osimertinib) has not been established in the EGFR T790M plasma-positive, tissue-negative or unknown population and clinical data for T790M plasma-positive patients are limited; therefore, testing using plasma specimens is most appropriate for consideration in patients from whom a tumor biopsy cannot be obtained.
- TAGRISSO efficacy has not been established in patients with EGFR exon 19 deletions < 0.08% MAF, in patients with EGFR L858R < 0.09% MAF, and in patients with
- EGFR T790M < 0.03% MAF.
- The test is not intended to be used for standalone diagnostic purposes.
- The test is intended to be performed on specific serial number-controlled instruments by Guardant Health, Inc.
- A negative result for any given variant does not preclude the presence of this variant in tumor tissue.
- Decisions on patient care and treatment must be based on the independent medical judgment of the treating physician, taking into consideration all applicable information concerning the
 patient's condition, such as patient and family history, physical examinations, information from other diagnostic tests, and patient preferences, in accordance with the standard of care.
 ctDNA shedding rate may be lower in patients with primary central nervous system (CNS) tumors.

Definition of Categories

The test report includes genomic finding reported in the following categories:

Category	Prescriptive use for Therapeutic Product	Clinical Performance	Analytical Performance	Comments
<u>Category 1</u> : Companion Diagnostic (CDx)	Yes	Yes	Yes	ctDNA biomarkers linked to the safe and effective use of the corresponding therapeutic product, for which Guardant360 CDx has demonstrated clinical performance shown to support therapeutic efficacy and strong analytical performance for the biomarker.
Category 2: ctDNA Biomarkers with Strong Evidence of Clinical Significance in ctDNA	No	No	Yes	ctDNA biomarkers with strong evidence of clinical significance presented by other FDA-approved liquid biopsy companion diagnostics for which Guardant360 CDx has demonstrated analytical reliability but not clinical performance.
<u>Category 3A:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: strong analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated analytical performance including analytical accuracy, and concordance of blood-based testing to tissue-based testing for the biomarker.
<u>Category 3B:</u> Biomarkers with Evidence of Clinical Significance in tissue supported by: analytical validation using ctDNA	No	No	Yes	ctDNA biomarkers with evidence of clinical significance presented by tissue-based FDA-approved companion diagnostics or professional guidelines for which Guardant360 CDx has demonstrated minimum analytical performance including analytical accuracy.
Category 4: Other Biomarkers with Potential Clinical Significance	No	No	Yes	ctDNA biomarkers with emergent evidence based on peer-reviewed publications for genes/variants in tissue, variant information from well-curated public databases, or <i>in-vitro</i> preclinical models, for which Guardant360 CDx has demonstrated minimum analytical performance.

Performance Characteristics

Please refer to product label, www.guardant360cdx.com/technicalinfo. Clinical Performance has not been established for biomarkers in categories 2, 3A, 3B, and 4. Guardant360 CDx is indicated to report the following SNVs (*AKT1, ALK, APC, AR, ARAF, ATM*, BRAF, BRCA1**,BRCA2**, CCND1, CDH1, CDK4, CDK6,CDK12*, CDKN2A, CTNNB1, EGFR, ERBB2, ESR1, FGFR1, FGFR2, FGFR3, GATA3, GNA11, GNAQ, HRAS, IDH1, IDH2, KIT, KRAS, MAP2K1, MAP2K2, MET, MLH1, MTOR, MYC, NF1, NFE2L2, NRAS, NTRK1, NTRK3, PDGFRA, PIK3CA, PTEN, RAF1, RET, RHEB, ROS1, SMAD4, SMO, STK11, TERT, TSC, VHL). Indels (<i>ALK, ATK1, APC, ATM*, BRAF, BRCA1**, BRCA2**, CDH1, CDK12*, CDKN2A, EGFR,ERBB2, ESR1, FGFR2, GATA3, HNF1A, HRAS , KIT, KRAS, MET, MLH1, NF1, PDGFRA, PIK3CA, PTEN, RET, ROS1, STK11, TSC1, VHL)*, Fusion (*ALK , NTRK1, RET, ROS1*), and Amplifications (*ERBB2, MET*).

*Reporting is enabled for pathogenic germline alterations only. Somatic alterations will not be reported. | ** Reporting is enabled for both germline and somatic alterations.

Testing performed at: Guardant Health

Laboratory Director: [First and Last Name] | CLIA ID: [XXXXXX] | CAP #: [XXXXXX] | 505 Penobscot Drive Redwood City, CA, 94063, USA



Breast patient with only Category 4 variant



REPORTING		PHYSICIAN
Report Date:	MAR-20-2017	Dougie Houser
Receipt Date:	MAR-04-2017	Center for People Who are Sick and Want to Get Better
Collection Date:	MAR-03-2017	123 Four St., Metropolis, NY, 12345, USA
Specimen:	Blood	Ph: (808) 555-1234 Fax: N/A
Status:	FINAL	Additional Recipient: N/A

No reportable alterations with companion diagnostic (CDx) claims

Other Biomarkers Identified

Results reported in this section are not prescriptive or conclusive for labeled use of any specific therapeutic product. See professional services section for additional information.

Other Biomarkers with Potential Clinical Significace

Clinical significance has not yet been established for biomarkers in this section. See the professional services section for additional information

-BRAF V600K





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