

SARS-CoV-2 IgG (COV2G)

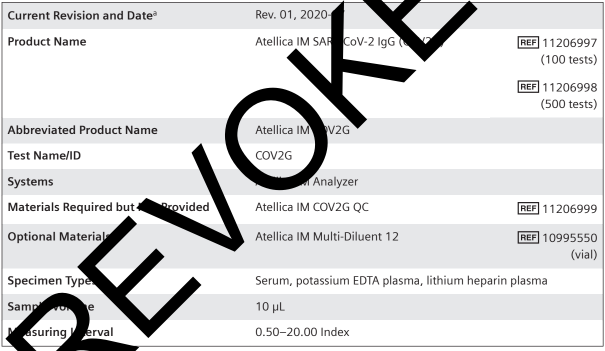
For Use Under Emergency Use Authorization Only

For in vitro diagnostic use.

For prescription use only.

The results of this semi-quantitative test should not be interputed as an inclusion or degree of immunity or protection from reinfection.

Assay for the Detection of IgG Antibodies to SARS-CO-



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Intend Use

The Atellica® IM SARS-CoV-2 IgG (COV2G) assay is a chemiluminescent immunoassay intended for qualitative and semi-quantitative detection of IgG antibodies to SARS-CoV-2 in human serum and plasma (potassium EDTA and lithium heparin) using the Atellica® IM Analyzer. The Atellica IM SARS-CoV-2 IgG (COV2G) assay is intended for use as an aid in identifying individuals with an adaptive immune response to SARS-CoV-2, indicating recent or prior infection. At this time, it is unknown for how long antibodies persist following infection and if the presence of antibodies confers protective immunity. The Atellica IM SARS-CoV-2 IgG (COV2G) assay should not be used to diagnose acute SARS-CoV-2 infection. Testing is limited to laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C 263a, that meet requirements to perform moderate or high complexity tests.

Results are for the detection of SARS-CoV-2 IgG antibodies. IgG antibodies to SARS-CoV-2 are generally detectable in blood several days after initial infection, although the duration of time antibodies are present post-infection is not well characterized. Individuals may have detectable virus present for several weeks following seroconversion.

Laboratories within the United States and its territories are required to report all positive results to the appropriate public health authorities.

The sensitivity of the Atellica IM SARS-CoV-2 IgG (COV2G) assay early after infection is unknown. Negative results do not preclude acute SARS-CoV-2 infection. If acute infection is suspected, direct testing for SARS-CoV-2 is necessary.

False positive results for the Atellica IM SARS-CoV-2 IgG (COV2G) assay may occur due to cross-reactivity from pre-existing antibodies or other possible causes.

The Atellica IM SARS-CoV-2 IgG (COV2G) assay is only for use under the Food Administration's Emergency Use Authorization.

Summary and Explanation

COVID-19 (coronavirus disease 2019) is the illness resulting from unection with RS-60-2 (severe acute respiratory syndrome coronavirus 2) virus.¹⁻⁵ The arus sprands reading from person to person or possibly from environmental exposure.⁶ Expanded apports spread by both asymptomatic and symptomatic individuals.⁷

Antibodies appear approximately 1-3 weeks post-sympto produced in both symptomatic and asymptomatic fecti profiles, near-simultaneous production of both IgM and patients with confirmed SARS-CoV-2. Titer of entibody though additional data is needed to confirm this. onset in high to dients and are ^{8,9} Unlike typical seroconversion has the baserved in symptomatic be higher in symptomatic disease,

nclude spike antibody and Antibodies produced to structural prot hs of the viru nucleocapsid antibody. Data show both gM and IgG a ibodies for these structural proteins appear with seroconversion. IgM (disappea , but IgG remains detectable in most entu patients. Spike is a transmembrane prised of two regions: S1 and S2. S1 lycop mediates recognition and binding o he viral receptor (ACE2) on host cells, and S2 facilitates viral fusion and entry.¹² of S1 is comprised of the receptor binding domain a majori (RBD) that binds dire ighly immunogenic. The S1 RBD in SARS-CoV-2 y to Ac contains both un e and conserved ences compared to other beta-coronaviruses. ment target or include the S1 RBD, as initial data indicate Multiple vaccin in deve can be neutralizing.¹⁴⁻²³ The ability to identify specific antibodies antibodies to t eqi associated with r lization be an important adjunct to the detection of an immune respo. ר∕ור

Princip es of the Procedure

The A area IM COV2G assay is a fully automated 2-step sandwich immunoassay using indirect chemilus nescent technology. The patient specimen is incubated with the Solid Phase Reagent. In Solid Phase contains a preformed complex of streptavidin-coated microparticles and biotinylated SARS-CoV-2 recombinant antigens. The antigen-coated particles subsequently capture SARS-CoV-2 specific antibodies in the specimen. The antibody-antigen complex is washed and Lite Reagent is added. The Lite Reagent consists of an acridinium-ester-labeled anti-human IgG mouse monoclonal antibody. The entire complex is washed and the signal is generated in the presence of Lite Reagent bound to the Solid Phase via the anti-SARS-CoV-2 IgG:SARS-CoV-2 antigen complex.

A direct relationship exists between the amount of SARS-CoV-2 IgG antibody present in the patient sample and the amount of relative light units (RLUs) detected by the system.

A result of reactive or nonreactive is determined according to the Index Value established with the calibrators. Refer to *Interpretation of Results*.

Reagents

Material Description	Storage	Stability
Atellica IM COV2G ReadyPack [®] primary reagent pack ^{a, b}	Unopened at 2–8°C	Until expiration date on product
Lite Reagent 10.0 mL/reagent pack Mouse monoclonal anti-human IgG antibody labeled with acridinium ester (~0.05 µg/mL); buffer; surfactant; bovine serum albumin (BSA); sodium azide (< 0.1%) Solid Phase 10.0 mL/reagent pack Streptavidin-coated paramagnetic microparticles preformed with biotinylated SARS-CoV-2 S1 RBD antigen (~1.0 µg/mL); buffer; bovine serum albumin; horse serum; surfactant; sodium azide (< 0.1%) Ancillary Well Reagent 10.0 mL/reagent pack Buffer; surfactant; bovine serum albumin; horse serum; sodium azide (< 0.1%)	Onboard	28 days
Atellica IM COV2G CAL ^{a, b} COV2G CAL L:	Unon ned at 2-8	Until expiration date on product
1.0 mL/vial Processed* human plasma nonreactive for SAR <u>S-CoV-2</u>	d at 2–8°C	60 days
antibodies; sodium azide (< 0.1%) *Processed plasma is defibrinated and filtend plasma. COV2G CAL H: 1.0 mL/vial Horse serum spiked with human more clone aG antibodies to SARS-CoV-2; sodium azit (< 0.1	1t room temperature	8 hours
Atellica IM Multi-Diluent 2ª, c 20.0 mL/vial	At 2–8°C	Until expiration date on product
Human serum; determents; glycerol, the firm; preservatives	Opened at 2–8°C	21 weeks

^a Store in an up, ht persion.

^b Prevent exposure counlight are neat.

Reference on a non-al Many rials

rning and Precautions

or Use Under Emergency Use Authorization Only

in vitro diagnostic use only.

For prescription use only.

This test has not been FDA cleared or approved; the test has been authorized by FDA under an Emergency Use Authorization (EUA) for use by laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C 263a, that meet requirements to perform moderate or high complexity tests.

This test has been authorized only for detecting the presence of IgG antibodies against SARS-CoV-2, not for any other viruses or pathogens.

This test is only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of *in vitro* diagnostic tests for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. § 360bbb-3(b)(1), unless the authorization is terminated or revoked sooner.

COV2G

CAUTION

Federal (USA) law restricts this device to sale by or on the order of a licensed healthcare professional.

Safety data sheets (SDS) available on siemens-healthineers.com.



CAUTION POTENTIAL BIOHAZARD

Contains human source material. Each donation of human blood or blood component was tested by FDA-approved methods for the presence of antibodies to human immunodeficiency virus type 1 (HIV-1) and type 2 (HIV-2), as well as for hepatitis B surface antigen (HBsAg) and antibody to hepatitis C virus (HCV). The test results were negative (not repeatedly reactive). No test offers complete assurance that these or other infectious agents are absent; this material should be handled using good laboratory practices and universal precautions

CAUTION

This device contains material of animal origin and should be handled as a potential carrier and transmitter of disease.

Contains sodium azide as a preservative. Sodium azide can reas with paper or leav plumbing to form explosive metal azides. On disposal, flush reagent with the reavel volume of water to prevent buildup of azides. Disposal into drain systems must be in compliance with prevailing regulatory requirements.

Dispose of hazardous or biologically contaminated pater descording to the practices of your institution. Discard all materials in a safe and accepta anner and in compliance with prevailing regulatory requirements.

Storage and Stability

Store all reagents at 2–8°C in an wrigh position, away from light and heat. Do not use products beyond the expiration day print of on the product labeling.

For information about product storage and stating, refer to *Reagents*.

Onboard Stability

Discard products whe end of the onserved stability interval. Do not use products beyond the expiration date winted on the product labeling.

For information product oboard stability, refer to *Reagents*.

Specimer Conection and Handling

Storm and a sense (notassium EDTA and lithium heparin) are the recommended sample types for the activity. Do not use heat-inactivated specimens.

Collecting the Specimen

- Observe universal precautions when collecting specimens. Handle all specimens as if they are capable of transmitting disease.²⁶
- Follow recommended procedures for collection of diagnostic blood specimens by venipuncture.²⁷
- Follow the instructions provided with your specimen collection device for use and processing.²⁸
- Allow blood specimens to clot completely before centrifugation.²⁵

- Keep tubes capped at all times.²⁵
- Test specimens as soon as possible after collecting. Store specimens at 2–8°C if not tested immediately within 8 hours.

Storing the Specimen

- Thawed frozen specimens must be clarified by centrifugation prior to testing. Do not store in a frost-free freezer.
- Freeze samples, devoid of red blood cells, at \leq -20°C for longer storage.

The handling and storage information provided here is based on data or references maintained by the manufacturer. It is the responsibility of the individual laboratory to use all available references and/or its own studies when establishing alternate stability criteria to meet specific needs.

Transporting the Specimen

Package and label specimens for shipment in compliance word applicable federal and international regulations covering the transport of clinical specimens are eticlogical agents.

If shipment is expected to exceed 2 days, ship specine as frozen store samples capped and upright at 2–8°C upon arrival.

Preparing the Samples

This assay requires 10 μ L of sample for a scale of preination. This volume does not include the unusable volume in the sample contained in the additional volume required when performing duplicates or other test of the same sample. For a complete list of appropriate sample containers and information about thermiting the minimum required volume, refer to the system online help.

Do not use samples with opparent contamination.

Before placing samples on the system, ensue that samples are free of:

- Bubbles or foam.
- Fibrin or therp, isulate natter.

Remove particulates by centre pation according to CLSI guidance and the collection device manufacturer's reported ations.²⁵



Procedure

Materials Provided

The following materials are provided:

 Quality control assigned value sheet CONTROL LOT VAL a Additional system fluids are required to operate the system: Atellica IM Wash, Atellica IM Acid, Atellica IM Baa and Atellica IM Cleaner for system fluid instructions for use, refer to the Document Library. 	REF	Contents	Number of Tests
Solid Phase, and Ancillary Well Reagent Atellica IM COV2G master curve and test definition MCTOFF 2 vials Atellica IM COV2G CAL low calibrator CAL II 2 vials Atellica IM COV2G CAL high calibrator CAL II Atellica IM COV2G CAL calibrator assigned value sheet CAL IT VAL Materials Required but Not Provided The following materials are required to perform associate the system control provided:	11206997	Solid Phase, and Ancillary Well Reagent Atellica IM COV2G master curve and test definition MCTDEF 1 vial Atellica IM COV2G CAL low calibrator CAL L 1 vial Atellica IM COV2G CAL high calibrator CAL H	100
Atellica IM Analyzer ^a Atellica IM Analyzer ^a 11206999 Atellica IM COV2G QC (quality control material) 2 x 2.0 mL positive quality control, level 1 control 2 uality control assigned value sheet a Additional system fluids are required to operate the system: Atellica IM Wash, Atellica IM Acid, Atellica IM Ba	Material	Solid Phase, and Ancillary Well Reagent Atellica IM COV2G master curve and test definition MCTDEF 2 vials Atellica IM COV2G CAL low calibrator CAL L 2 vials Atellica IM COV2G CAL high calibrator CAL H Atellica IM COV2G CAL calibrator assigned value sheet CAL S Required but Not Provided	500
11206999 Atellica IM COV2G QC (quality control material) 0 mL negative quality control, level 1 control - 2 x 2.0 mL positive quality control, level 2 control + Quality control assigned value sheet a Additional system fluids are required to operate the system: Atellica IM Wash, Atellica IM Acid, Atellica IM Baa and Atellica IM Cleanant for system fluid instructions for use, refer to the Document Library.	REF	Description	
material) 2 x 2.0 mL positive quality control, level 2 control		Atellica IM Analyzer ^a	
and Atellica IM Cleaner For system fuid instructions for use, refer to the Document Library.	11206999	material) 2 x 2.0 mL positive quality control, level	2 Control + 2
Optional Materials		I system fluids are required to operate the system: Atellica IM Wash, Atellica IM Acid, A ca IM Cleane are system fluid instructions for use, refer to the Document Library.	tellica IM Base,
	Optiona	l M <u>ateri</u> als	
The mowing naterial provided to perform this assay, but are not provided:	The	e nowing naterial of be used to perform this assay, but are not provided:	

REF	2/ESChp.	
10995550	Atellica IM Multi-Diluent 12 (diluent)	20.0 mL/vial DL

Assay Procedure

The system automatically performs the following steps:

- 1. Dispenses 10 μL of sample into a cuvette.
- 2. Dispenses 100 μL of Solid Phase and 100 μL of Ancillary Well Reagent, then incubates for 12 minutes at 37°C.
- 3. Performs a wash sequence using Atellica IM Wash.
- 4. Resuspends the washed particles in 150 μL of Atellica IM Wash.
- 5. Dispenses 100 μL of Lite Reagent, then incubates for 8 minutes at 37°C.

- 6. Performs a wash sequence using Atellica IM Wash.
- 7. Dispenses 300 μ L each of Atellica IM Acid and Atellica IM Base to initiate the chemiluminescent reaction.
- 8. Reports results.

Preparing the Reagents

All reagents are liquid and ready to use. Before loading the packs onto the system, reagents require mixing. For information about mixing the reagents, refer to the system online help.

Preparing the System

A daily cleaning procedure must be completed prior to and after your laboratory's batched testing for the Atellica IM COV2G assay.

Ensure that sufficient materials are loaded on the system. Refer to daterials Provided and Materials Required but Not Provided for guidance about required to gents.

For information about loading products, refer to the system online he

Master Curve Definition

Before initiating calibration on each new lot of reggent, which he assault haster curve and test definition by scanning the MCTOFF 2D barcodes. For information above entering the master curve and test definition, refer to the system online help.

Performing Calibration

For calibration of the Atellica IM JV2G as y, us the calibrators provided with each kit.

Note Calibrators provided in a assay kit mus only be used with the reagent lot provided in the same kit.

Calibration Frequency

Perform a calibration if one more of the following conditions exist:

- When changing he numbers of primary reagent packs.
- At the end of the lot calibration interval, for a specified lot of calibrated reagent on the system.
- At the order of the park calibration interval, for calibrated reagent packs on the system.
- hen in cated y quality control results.
- ter major vaintenance or service, if indicated by quality control results.

te When-bading a new primary reagent pack, a calibration is not required if there is a valid t calibration. For information about lot calibration and pack calibration, refer to the system in *help*.

Stability Interval	Days
Lot Calibration	28
Pack Calibration	14
Reagent Onboard Stability	28

Follow government regulations or accreditation requirements for calibration frequency. Individual laboratory quality control programs and procedures may require more frequent calibration.

Preparing the Calibrators

Calibrators are liquid and ready to use. Allow the calibrators to equilibrate to room temperature. Gently mix and invert the vials to ensure homogeneity of the material.

Use calibrators within the stability limits specified in *Reagents* and discard any remaining material.

Calibration Procedure

The calibrators are provided in dropper vials. Each dispensed drop is approximately 50 µL.

The required sample volume for testing depends on several factors. For information about sample volume requirements, refer to the system online help.

Use the following lot-specific materials to perform calibration:

- For the master curve and assay test definitions, refer to the lot-specific vaster curve and test definition sheet MCTOFF provided with the assay reagents.
- Calibrators provided in an assay kit must only be used with readints from that assay of lot. Do not use calibrators from one assay kit lot with reagents from a different as with lot.
- For the calibrator definitions, refer to the calibrator assignt value neet **CAL LOT VAL** provided with the calibrator materials.
- Generate lot-specific barcode labels to use with the cubrator sa

For instructions about how to perform the calibration providered refer to the system online help.

Performing Quality Control

For quality control of the Atellica IM CC 2G assay, use the Atellica IM COV2G QC at least once during each day that samples are inally d. Use the quality control material in accordance with the quality control instructions for se. For the assigned values, refer to the quality control value sheet CONTROL LOT VAL provided.

A satisfactory level of performance is achieved when the analyte values obtained are within the expected control interval to be system or within your interval, as determined by an appropriate interpendeoratory quant, and the procedure. Follow your laboratory's quality control procedures if the abults obtained do not fall within the acceptable limits. For information as but entrol quality control definitions, refer to the system online help.

Follow generative gulation or accreditation requirements for quality control frequency. Individual law ratory chality control programs and procedures may require more frequent guracy control testing.

ples after a successful calibration.

Taking Corrective Action

If the quality control results do not fall within the expected control interval, do not report results. Perform corrective actions in accordance with established laboratory protocol. For suggested protocol, refer to the system online help.

Results

Calculation of Results

The system determines the result using the calculation procedure described in the system online help. Refer to *Interpretation of Results*.

Dilutions

Sample	Dilution	Minimum Sample Volume (µL)
Serum and plasma	1:2	100
Serum and plasma	1:4	50
Serum and plasma	1:8	25

The system does not perform onboard dilutions for the Atellica IM COV2G assay.

If patient results exceed the upper limit of the analytical measuring interval of the assay, or if laboratory protocol requires manual dilution, manually dilute the patient sample.

For manual dilutions, perform the following actions:

- Use Atellica IM Multi-Diluent 12 (vial) to prepare a manual dution. Refer to optional Materials.
- For information about ordering tests for manually diluced samples, fer to be system online help.
- Ensure that results are mathematically corrected for lilutum. If a dilution factor is entered when scheduling the test, the system automatically concluses the esult.

Interpretation of Results

The system reports Atellica IM COV2G assay of a model of the system reports Atellica IM COV2G assay of a model of the system reports and as Nonreactive or Reactive:

- Nonreactive: < 1.00 Index these samples are unsidered negative for SARS-CoV-2 IgG antibodies. Report nonreal ive patient realits as < 1.00 Index.
- Reactive: ≥ 1.00 Index. The e samples are considered positive for SARS-CoV-2 IgG antibodies. Report reactive reactive reactive numeric Index Value for semi-quantitative measurements.

Results of this as a should a ways be interpreted in conjunction with the patient's medical history, clinical presentation, and other findings.

Limitatio

The following information pertains to limitations of the assay:

e clin applicability of a quantitative or semi-quantitative result is currently unknown d canno conterpreted as an indication or degree of immunity nor protection from *infection*, nor compared to other SARS-CoV-2 antibody assays.

This device should not be used to diagnose or exclude acute SARS-CoV-2 infection. Direct testing for SARS-CoV-2 with a molecular assay should be performed to evaluate acute affection in symptomatic individuals.

- Performance characteristics have not been established for the assay used in conjunction with other manufacturers' assays for specific SARS-CoV-2 serological markers. Laboratories are responsible for establishing their own performance characteristics.
- The performance of the assay has not been established with cord blood, neonatal specimens, cadaver specimens, or body fluids other than serum or plasma.
- Results obtained with the assay may not be used interchangeably with values obtained with different manufacturers' test methods.
- A positive result may not indicate previous SARS-CoV-2 infection. Consider other information, including clinical history and local disease prevalence, in assessing the need for a second, but different, serology test to confirm an immune response.

- A negative result for an individual subject indicates absence of detectable anti-SARS-CoV-2 antibodies. Negative results do not preclude SARS-CoV-2 infection and should not be used as the sole basis for patient management decisions. A negative result can occur if the quantity of the anti-SARS-CoV-2 antibodies present in the specimen is below the detection limits of the assay, or the antibodies that are detected are not present during the stage of disease in which a sample is collected.
- Performance has only been established with the specimen types listed in the Intended Use. Other specimen types have not been evaluated and should not be used with this assay.
- Results are not intended to be used as the sole basis for patient management decisions. Test results should be interpreted in conjunction with clinical observations, patient history, epidemiological information, and other laboratory findings.
- A reactive test result does not exclude past or present infection by other oronavil as such as SARS-CoV-1, MERS-CoV, HKU1, 229E, NL63, or OC43.
- SARS-CoV-2 antibodies may not be detectable in patients with re-• ions (7–10 days or less) or in samples collected from patients less, n 7 davs a no polymerase chain reaction (PCR) result. Patient specimens ly be no lected during the early (pre-seroconversion) phase of illness or to a c ine in titer over time. In addition, the immune response may be depressed romised, or elde munoco immunosuppressed patients.
- It is not known at this time if the presence of antibod s to SARS-Conc confers immunity to re-infection.
- This test should not be used for donor screening.

Conditions of Authorization for the Laboratory

The Atellica IM SARS-CoV-2 IgG (COV2C assay Letter c Authorization, along with the authorized Fact Sheet for Healthcap Properties, the authorized Fact Sheet for Patients, and authorized labeling are available on the FD.

https://www.fda.gov/medical-devices oronavirus-disease-2019-covid-19-emergency-use-authorizations-medical-devices oronavirus-disease-2019-covid-19-emergency-use-

Authorized laborationes using the Atsociation IM SARS-CoV-2 IgG (COV2G) assay must adhere to the Conditions. Authorization indicated in the Letter of Authorization as listed below:

- Authorized Corportes^a using the Atellica IM SARS-CoV-2 IgG (COV2G) assay will include with the result poorts, all outhorized Fact Sheets. Under exigent circumstances, other authorized methods for disseminating these Fact Sheets may be used, which may include mass mena.
- the zed last cories using the Atellica IM SARS-CoV-2 IgG (COV2G) assay will use the project as outlined in the Instructions for Use. Deviations from the authorized procedures, including the authorized instruments, authorized clinical specimen types, authorized control paterials, authorized other ancillary reagents and authorized materials required to use the Atellica IM SARS-CoV-2 IgG (COV2G) assay are not permitted.
- Authorized laboratories that receive the Atellica IM SARS-CoV-2 IgG (COV2G) assay will notify the relevant public health authorities of their intent to run the assay prior to initiating testing.
- Authorized laboratories using the Atellica IM SARS-CoV-2 IgG (COV2G) assay will have a process in place for reporting test results to healthcare providers and relevant public health authorities, as appropriate.

- Authorized laboratories will collect information on the performance of the Atellica IM SARS-CoV-2 IgG (COV2G) assay and report to DMD/OHT7-OIR/OPEQ/ CDRH (via email: CDRH EUA Reporting@fda.hhs.gov) and Siemens Healthineers Technical Support (https://www.siemens-healthineers.com/en-us/; tel: 1-877-229-3711) any suspected occurrence of false reactive or false nonreactive results and significant deviations from the established performance characteristics of the assay of which they become aware.
- All laboratory personnel using the Atellica IM SARS-CoV-2 IgG (COV2G) assay must be appropriately trained in automated immunoassay techniques and use appropriate laboratory and personal protective equipment when handling this kit, and use the Atellica IM SARS-CoV-2 IgG (COV2G) assay in accordance with the authorized labeling. All laboratory personnel using the assay must also be trained in and be familiar with the interpretation of results of the the Atellica IM SARS-CoV-2 IgG (COV2G) assay.
- Siemens Healthineers, authorized distributors, and authorized la prator ousing the Atellica IM SARS-CoV-2 IgG (COV2G) assay will ensure that apprecords associated with this EUA are maintained until otherwise notified by FDA. Such records will be main available to FDA for inspection upon request.

^a The letter of authorization refers to, "Laboratories certified you're the Choical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, that meet requirement to perform Moderate or high complexity tests" as "authorized laboratories".

Performance Characteristics

Analytical Measuring Interval

0.50–20.00 Index is reported as conreact. (< λ R Index) or Reactive (\geq 1.00 Index).

The lower limit of the analytic measuring in rval is defined by the LoQ (0.50 Index). However, report nonreactive plaient results as a 1.00 Index. When sample results exceed the upper limit of the analytic lime, wring interval, refer to Dilutions.

Detection Capability

Detection canability are determined in accordance with CLSI Document EP17-A2.²⁹ The following readilts were obtained

Method	Result (Index)
Limit control (LoB)	0.40
Light of Detection (LoD)	0.50
Line of antitation (20Q)	0.50

ts obtained at individual laboratories may vary from the data presented.

The LoB corresponds to the highest measurement result that is likely to be observed for a blank sample with a probability of 95%. The estimate of the LoB based on 2 reagent lots is 0.40 Index.

The LoD corresponds to the lowest concentration of IgG antibodies to SARS-CoV-2 that can be detected with a probability of 95%. The estimate of the LoD based on 2 reagent lots is 0.50 Index.

The LoQ corresponds to the lowest concentration of IgG antibodies to SARS-CoV-2 in a sample at which the within laboratory CV is \leq 20%. The LoQ of the assay based on 2 reagent lots is 0.50 Index.

The lower limit of the analytical measuring interval is defined by the LoQ (0.50 Index). However, report nonreactive patient results as < 1.00 Index.

Seroconversion Sensitivity

A total of 131 specimens were collected serially from 28 subjects with a clinical diagnosis of COVID-19 based on a positive SARS-CoV-2 polymerase chain reaction (PCR) method. Of these, seroconversion was observed in 8 panels with 3 or more blood draws. The results are shown in the table below:

Numbe		Number	First Draw			ive	First Reactive Draw		Last Draw	
Panel	Number of Draws	of Reactive Draws	Days Post PCR Positive	Index	Days Post PCR Positive	Index	Days Post PCR Positive	Ind	Days Post PCR Positive	dex
А	8	5	5	0.10	7	0.99	8	3.18	2	05
В	7	6	6	0.08	6	0.08	9	1.50	17	3.81
С	4	3	0	0.00	0	0.00	6	.85		6.13
D	4	2	5	0.02	6	0.08	9	52	10	3.56
E	5	2	0	0.01	4	24	5	1.1.	12	6.93
F	3	2	0	0.88	0	0.8	2	6.60	3	4.44
G	5	3	5	0.02		4		1.25	10	5.52
Н	8	4	2	0.10	1	0.4	5	1.44	7	5.64

Clinical Agreement

Positive percent agreement and negative percent agreement were determined in accordance with CLSI Document EP12-3. The performance of the Atellica IM COV2G assay was determined by testing a total of 2. The supples using the Atellica IM Analyzer.

Positive Percent Agreement

Positive percent uses and was betermined by testing 197 retrospective samples collected over the other of the from a unique donor subjects with a clinical diagnosis of COVID-19 based on a public SA 2-COV-2 polymerase chain reaction (PCR) method. The following table densibles polytive percent agreement by time of sampling following a positive PCR result:

Days After PCR Method	mber Tested	Reactive	Nonreactive	Positive Percent Agreement (95% CI)
0–6	91	51	40	56.04% (45.25%–66.44%)
7–13	64	59	5	92.19% (82.70%–97.41%)
≥14	42	42	0	100.00% (91.59%–100.00%)

Negative Percent Agreement

Negative percent agreement was determined by testing 1841 samples collected prospectively prior to the COVID-19 outbreak (before November 2019) from apparently healthy individuals and apparently healthy pregnant women in the United States. The results are shown in the table below:

Group	Number Tested	Nonreactive	Reactive	Negative Percent Agreement (95% CI)
Apparently Healthy	1744	1743	1	99.94% (99.68%–100.00%)
Apparently Healthy Pregnant Women	97	97	0	(96.2) 100.00%)
Total	1841	1840	1	99.95% (99.70%– 0.00%)

Precision

Single-Site Precision

A single-site precision study for the Atellica IM CO CLSI Document EP05-A3.³¹ Samples and a by co Control) were assayed in duplicate, in 2 runs or a Results for the precision of the Atellicate COV2 as

/2G assay was unducted in accordance with trols (a Negative Control and a Positive ay for 2000 Ays using the Atellica IM Analyzer. assay are presented in the following table:

			A	A leatabuilty		oratory Precision
Specimen Type	Nª	lean dex)	SD ^b (Index)	CV ^c (%)	SD (Index)	CV (%)
Serum A	80	0.8	0.02	2.7	0.03	3.8
Serum B		1.72	0.04	2.0	0.05	2.8
Serum C	80	7.25	0.33	4.5	0.58	8.0
Plasma, Lithiu Heparin	80	1.82	0.04	2.0	0.07	4.0
Plasma, CDTA	80	1.76	0.04	2.3	0.08	4.5
Cor of 1		0.01	0.01	N/A ^d	0.01	N/A
Cerrol 2	80	1.83	0.07	3.9	0.11	6.2

^a Num of measurements.

^b Standal eviation.

^c Coefficient f variation.

^d Not applicable.

Results obtained at individual laboratories may vary from the data presented.

Instrument and Lot Reproducibility

Reproducibility of the Atellica IM COV2G assay was evaluated on 2 Atellica IM instruments using 2 reagent lots. Samples and assay controls (a Negative Control and a Positive Control) were assayed in duplicate in 2 runs per day for 3 days. The data were analyzed to calculate the following components of precision: repeatability, between-run, between-day, between-lot, between-instrument, and reproducibility (total). Results for the reproducibility of the Atellica IM COV2G assay are presented in the following table:

			Repeata	Between- atability Run		1-			Between-Lot		Between- Instrument		Reproducibility	
Sample	Nª	Mean (Index)	SD ^b (Index)	CV ^c (%)	SD (Index)	CV (%)	SD (Index)	CV (%)	SD (Index)	CV (%)	SD (Index)	CV (%)	SD (Index)	CV (%)
Serum A	48	0.89	0.02	2.6	0.02	2.2	0.02	2.0	0.05	6.1	0.00	J.O		7.3
Serum B	48	2.00	0.04	2.0	0.02	1.2	0.07	3.3	0.17	8.5	0.	0.0	0.19	9.4
Serum C	48	8.99	0.37	4.1	0.28	3.1	0.00	0.0	0.26	2.9	0.17	9	0.56	6.2
Serum D	48	19.51	1.31	6.7	0.91	4.6	0.74	3.8	0.00		0.89	4.5		10.1
Plasma, Lithium Heparin	48	1.97	0.04	2.2	0.02	0.9	0.04	2.2	0.09	6	JB	1.3	0.11	5.7
Plasma, EDTA	48	2.00	0.04	2.1	0.02	1.1	0.04	2.2	(4	6.9	V	0.0	0.15	7.6
Control 1	48	0.01	0.00	N/A ^d	0.00	N/A	0.00	N/A	0	N/A	0.01	N/A	0.01	N/A
Control 2	48	2.03	0.07	3.4	0.00	0.0	0.01	0.	0.0	4.5	0.05	2.3	0.12	6.1

^a Number of measurements.

^b Standard deviation.

- ^c Coefficient of variation.
- ^d Not applicable.

Specimen Equivalency

Matched sample s a, and lithium heparin plasma) from the same serum, EDTA atrix comparison studies. Samples contained SARS-CoV-2 IgG levels donors were us for the suring interval. Specimen equivalency was determined by testing the r distributed ac OV2G assay using the Atellica IM Analyzer. Using a linear the samples wit tellica IM lts fro plasma samples were compared to serum results in accordance regres .³² The following results were obtained:

Tube (y) vs. 5 (x)	N ^a	Sample Interval	Slope (95% Cl)	Intercept (95% CI)	r ^b
EDTA (plasma)	36	0.61–17.06	1.05 (1.02–1.07)	-0.09 (-0.27–0.10)	0.997
lithium heparin (plasma)	36	0.55–16.15	1.02 (0.99–1.06)	-0.06 (-0.30–0.18)	0.995

^a Number of samples tested.

^b Correlation coefficient.

Agreement of the specimen types may vary depending on the study design and sample population used. Assay results obtained at individual laboratories may vary from the data presented.

Interferences

Interference testing was performed in accordance with CLSI Document EP07-ed3.³³ The impact of potentially interfering substances on the detection of SARS-CoV-2 IgG antibodies with the Atellica IM COV2G assay was evaluated with endogenous substances commonly found in serum and plasma specimens, including hemoglobin, conjugated bilirubin, unconjugated bilirubin, triglycerides, biotin, cholesterol, and protein. Serum samples were spiked with SARS-CoV-2 IgG at the following levels: unspiked, near cut-off (~1.0 Index), and low positive (~2.5 Index). Testing demonstrated a \leq 10% change for each substance at the indicated concentration.

Substance Test Concentration
1000 mg/dL
40 mg/dL
40 mg/dL
2000 mg/dL
3500 ng/mL
500 mg/dL
12 g/dL

Cross-Reactivity

Cross-reactivity was determined a accordance with CLSI Document EP07-ed3.³³ The assay was evaluated for potential cross-reactivity using the cellica in COV2G assay with the Atellica IM Analyzer. No false positive result were observed with the potential cross-reactants listed in the following table:

Number Tested	Number Reactive with Atellica IM COV2G Assay
24	0
5	0
5	0
4	0
10	0
4	0
10	0
14	0
25	0
12	0
14	0
8	0
15	0
	24 5 5 4 10 4 10 14 25 12 14 8

Clinical Category	Number Tested	Number Reactive with Atellica IM COV2G Assay
Human chorionic gonadotropin (hCG)	10	0
Human immunodeficiency virus (HIV) antibody	10	0
Influenza antibody	29	0
Influenza A antibody	6	0
Influenza B antibody	29	0
Measles antibody	5	0
Mycoplasma pneumoniae IgG	9	0
Parvovirus B19 antibody	7	0
Respiratory pathogen antibodies ^b	19	0
Respiratory syncytial virus (RSV) antibody	20	
Treponema pallidum (Syphilis) IgG	7	9
Varicella zoster virus (VZV) IgG	16	
Varicella zoster virus (VZV) IgM	5	
Total	322	

^a This group consists of samples from 24 subjects with actoining the discrete states, including anti-nuclear antibody (ANA; N = 6), Graves' disease (N = 6), rho matoid factor RF; N \rightarrow), Sjogren's syndrome (N = 3), and systemic lupus erythematosus (SLE; N = 2).

^b This panel consists of samples from 19 subjects we antibodies to fultiple respiratory pathogens, including Adenovirus antibodies (N = 6), *Bordetella perussis* (N = 17), flamydia pneumoniae IgG (N = 18), *Chlamydia psittaci* IgM (N = 1), *Haemophilus afluenza* (111 agG (N = 10), Influenza A IgG (N = 17), Influenza A IgM (N = 1), Influenza B IgG (N = 1), and *Mycoplasma pneumoniae* IgG (N = 4).

Results obtained at individual borato es may vary from the data presented.

Linearity

Linearity testin, was purformed in accordance with CLSI Document EP06-A.34

Patient productions for the product of SARS-CoV-2 IgG (1 serum, 1 EDTA plasma, and 1 lithium heparic plasm) were filuted with negative basepool to prepare a dilution series comprised of nine (9) level. Each level was tested in 3 replicates using an Atellica IM Analyzer. Linearity was do constrained of the analytical measuring interval of 0.50–20.00 Index with deviations from linear worthin 15%.

Taking in consideration the estimates of LoB, LoD, LoQ, precision, and linearity, the analytical hosuring interval of the Atellica IM COV2G assay is 0.50–20.00 Index.

Extended Measuring Interval (Dilutions)

Two serum samples, three lithium heparin plasma samples, and one EDTA plasma sample in the range of 12.65–32.97 Index were manually diluted 1:2, 1:4, and 1:8 with Atellica IM Multi-Diluent 12 and assayed for recovery. The recoveries ranged from 82.7%–111.6%.

The extended measuring interval of the Atellica IM COV2G assay by manual dilution of 1:2, 1;4, and 1:8 with Atellica IM Multi-Diluent 12 is 20.00–160.00 Index.

Sample	Dilution	Observed (Index)	Expected (Index)	Recovery (%)
Serum 1	_	31.47	_	_
	1:2	16.18	15.74	102.8
	1:4	8.45	7.87	107.3
	1:8	3.88	3.93	98.6
	Mean			102.9
Serum 2	_	32.97	-	
	1:2	13.63	16.	8
	1:4	7.80	8.24	7
	1:8	3.95	2	95.9
	Mean			91.1
Lithium heparin plasma 1	—	21 0		_
	1:2	10	10.70	99.5
	1:4		5.35	94.1
	8	2.25	2.68	84.2
	lean			92.6
Lithium heparin plasma 2		20.46	_	
	1:2	9.27	10.23	90.6
	1:4	4.71	5.11	92.1
	.8	2.29	2.56	89.4
	Mean			90.7
Lithium in place 3	_	12.65	_	_
	1:2	7.02	6.33	110.9
	1:4	3.19	3.16	100.9
	1:8	1.48	1.58	93.5
	Mean			101.8
EDTA plasma 1	_	21.19	_	_
	1:2	9.67	10.60	91.2
	1:4	5.47	5.30	103.3
	1:8	2.96	2.65	111.6
	Mean			102.0
Mean				97.1

current

Assay results obtained at individual laboratories may vary from the data presented.

Standardization

The Atellica IM COV2G assay standardization is traceable to an internal standard based on agreement with known positive and negative SARS-CoV-2 samples.

Currently no reference standard is available for this assay.

Technical Assistance

For customer support, contact your local technical support provider or distributor.

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Definition of Symbols

The following symbols may appear on the product labeling: Symbol Title and Description Symbol Consult instructions for use **i** Version of instructions for use Rev. 01 nstruction Internet URL address to access the electronic siemens.com/healthcare i siemens.com/document-library Revision REVISION Rev. Caution Consult instructions for e or accompar ng documents for cautionary information such as warnings autions that c not, for a variety of reasons, be presented nd pi on the medical de **Biological risks** Potenti logical ris are associated with the medical device. ronment erous to Oral, dermal, or inhalation hazard Inhalation hazard Respiratory or internal health Flammable Flammable to extremely flammable Oxidizing

Symbol	Symbol Title and Description
	Explosive
	Τοχίς
\Diamond	Compressed gas
紊	Keep away from sunlight Prevent exposure to sunlight and heat.
<u>††</u>	Up Store in an upright position.
	Do not freeze
2°C / ^{8°C}	Temperature limit Upper and lower limits of temperature indicators a pedjacent to the upper and lower horizontal lines.
	Handheld barcode
IVD	In vitro diagnotic medical devi
∑∑(n)	Contains & ficient Comparents Total number of IVD tests the system can perform with the IVD kit reagents appears Discent to the symbol.
RxOnly	Prescription (e. tre (US only) Applies only to United States-registered IVD assays. CAUTION: Federal (USA) law restricts this device to sale by or on the order of a license healthcare professional.
	Minung of substances A product before use.
g Com	Reconstitute and mix lyophilized product before use.
	Target
← →	Interval
	Legal Manufacturer
EC REP	Authorized Representative in the European Community
R	Use-by date Use by the designated date.

Symbol	Symbol Title and Description
LOT	Batch code
REF	Catalog number
E.	Recycle
PRINTED WITH SOY INK	Printed with soy ink
CE	CE Mark
CE xxxx	CE Mark with notified body ID number Notified body ID number can vary.
YYYY-MM-DD	Date format (year-month-day)
CHECKSUM	Variable hexadecimal number that ensures the Mass. Cruce and Calibrator definition values entered are valid.
MC DEF	Master Curve Definition
LOT DTL	Lot Details
UNITS C	Common Units
UNITS SI	International System of Leits
MATERIAL	Material
MATERIAL ID	Unique matens a lentification number
CONTROL NAME	Name of control
CONTROL TYPE	is the contraction of contraction of the contractio

Legal Morph

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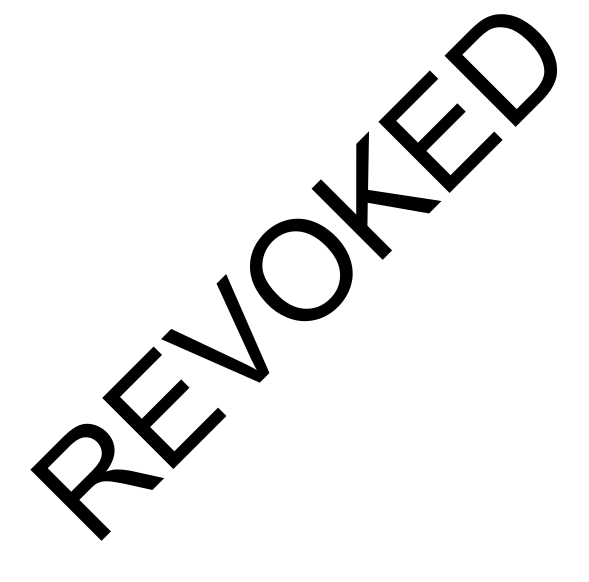
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SARS-CoV-2 IgG Quality Control (COV2G QC)

Current Revision and Date ^a	Rev. 01, 2020-07
Product Name	Atellica IM SARS-CoV-2 IgG Quality Control (COV2G
Abbreviated Product Name	Atellica IM COV2G QC
	2 x 2.0 mL negative quality control, level 1 CONTRACT 1 F 11206999 2 x 2.0 mL positive quality control, level 2 CONTROL + Quality control assigned value shere CONTROL LOVE VAL
Systems	Atellica IM Analyzer
^a A vertical bar in the page mar	gin indicates technical content hat differs the theorevious version.

^a A vertical bar in the page margin indicates technical content

FOR USA:

For Use Under Emergency Use Authorization

For in vitro diagnostic use.

For Professional Use.

Intended Use

gG Quality Control (COV2G QC) is for in vitro diagnostic use in The Atellica® IM SARS-CoVmonitoring the ision and ccuracy of the Atellica[®] IM SARS-CoV-2 IgG (COV2G) assay using the Atellica® Analy

hly

Material Pescription

Material Descript	Storage	Stability
Atela a IM C /2G QC C 2G Cont / 1:	At 2–8°C	Until expiration date on product
Processory human plasma nonreactive for SARS-CoV-2	Opened at 2–8°C	60 days
antiboo · sodium azide (< 0.1%) *Processer Jasma is defibrinated and filtered plasma.	At room temperature	8 hours
COV2G Convol 2: 2.0 mL/vial	Atellica [®] Sample Handler ^a	
Horse serum spiked with human monoclonal IgG antibodies to SARS-CoV-2; sodium azide (< 0.1%)		

Refer to the supplementary document "Atellica Sample Handler Calibrator and QC Storage and Stability" for а information about storage and stability of materials in the Cal-QC tube storage area.

Warnings and Precautions

FOR USA:

For Use Under Emergency Use Authorization Only

For in vitro diagnostic use.

For Professional Use.

CAUTION

Federal (USA) law restricts this device to sale by or on the order of a licensed healthcare professional.

Safety data sheets (SDS) available on siemens-healthineers.com.

CAUTION POTENTIAL BIOHAZARD

Contains sodium azide as a preservative. Sodium azide can react with copper of read publicing to form explosive metal azides. On disposal, flush reagents with a large volume of the reader to prevent buildup of azides. Disposal into drain systems must be a compliance with prevailing regulatory requirements.

Dispose of hazardous or biologically contaminated mater is according to the practices of your institution. Discard all materials in a safe and accertable anner and in compliance with prevailing regulatory requirements.

Storage and Stability

Store quality control materials in an up the position. Uality control materials are stable until the expiration date on the product when stored at 2-8 $\stackrel{\circ}{_{\sim}}$. Discard products at the end of the onboard stability interval. Do not use pullucts beyong the expiration date printed on the product labeling.

For information about product storage and stability, refer to Material Description.

Note Refer to the supplementary document "Atellica Sample Handler Calibrator and QC Storage and Stability for informational, but storage and stability of materials in the Cal-QC tube storage are

Performing Qual Cristrol

Performance vality control recedure at least once during each day that samples are analyzed. Test gality control same to after a successful calibration.

For w governments for quality control frequency. Indiversible values aboratory quality control programs and procedures may require more frequent quality entrol testing.

Treat all que y control samples the same as patient samples.

Preparing the Quality Control Materials

Quality control materials are liquid and ready to use. Gently mix and invert the vials to ensure homogeneity of the material.

Note Use quality control material within the stability limits specified in *Material Description* and discard any remaining material.

Quality Control Procedure

The product is provided in dropper vials. Each dispensed drop is approximately 50 μ L.

The required sample volume for testing depends on several factors. For information about sample volume requirements, refer to the system online help.

Use the following lot-specific materials to perform quality control:

- For the quality control (QC) definitions, refer to the lot-specific value sheet CONTROL LOT VAL provided with the quality control materials.
- Generate lot-specific barcode labels to use with the quality control samples.

For instructions about how to perform the quality control procedure, refer to the system online help.

Taking Corrective Action

If the quality control results do not fall within the assigned values, do protocorresults. Perform corrective actions in accordance with established laborator protocorrest suggested protocol, refer to the system online help.

Expected Values

For the assigned values, refer to the quality control va ovided. A sheet satisfactory level of performance is achieved when the alues obtained are within the nalyt II, as det expected control interval for the system or within nined by an our l appropriate internal laboratory quality control sc eme. Fo vour poratory's quality control procedures if the results obtained do not fall wit h the accep limits. For information about entering QC definitions, refer to the nline help. ster

The assigned values are traceable to the stand value zation of the assay. For additional information, refer to the assay instruction for us

Limitations

The Atellica IM COV2G Que is founce only with the Atellica IM COV2G assay. Assay values have not been established for a lays of the than the Atellica IM COV2G assay.

The results obtained using q ality control material depend on several factors. Erroneous results can occur a causes such as improper storage, inadequate mixing, reconstitution errors, or sample hand, a error associated with system or assay procedures.

The assigned control values should be used as a guide in evaluating performance. The control target and intervals should be adapted to each laboratory's individual requirements. Values obtained here d fall within the established interval. Each laboratory should establish exercise transition of the established interval. Each laboratory should establish exercise transition of the established interval. Follow the applicable government is used and local guidelines for quality control.

thni di sistance

r customer support, contact your local technical support provider or distributor.

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Definition of Symbols The following symbols may appear on the product labeling: Symbol Symbol Title and Description Consult instructions for use i Version of instructions for use Rev. 01 Internet URL address to access the elect nstruction siemens.com/healthcare or use siemens.com/document-library Revision REVISION Rev. Caution Consult instruction mying documents for cautionary information or use recautions that cannot, for a variety of reasons, be presented such as warnings and on the l device gical risks ological risks are associated with the medical device. Dangerous to environment Irritant Oral, dermal, or inhalation hazard Inhalation hazard Respiratory or internal health Flammable Flammable to extremely flammable

Symbol	Symbol Title and Description
	Oxidizing
\diamond	Explosive
	Тохіс
\bigotimes	Compressed gas
挙	Keep away from sunlight Prevent exposure to sunlight and heat.
<u>††</u>	Up Store in an upright position.
	Do not freeze
2°C-4 ^{8°C}	Temperature limit Upper and lower units of the peraturn indicators are adjacent to the upper and lower horizont plines.
	Handheld bardele scanner
IVD	In vitro diagnostic medical device
$\overline{\Sigma}(n)$	Contains 1 ⁴⁵ Cont for <n> tests Total number 6.1VD tests the system can perform with the IVD kit reagents appears djacent to the symbol.</n>
RxOnly	Prescription device (US only) Apples only to United States-registered IVD assays. UTION: Federal (USA) law restricts this device to sale by or on the order of a licensed healthcare professional.
	Mixing of substances Mix product before use.
	Reconstitute and mix lyophilized product before use.
→■ ←	Target
← →	Interval
	Legal Manufacturer

Symbol	Symbol Title and Description
EC REP	Authorized Representative in the European Community
8	Use-by date Use by the designated date.
LOT	Batch code
REF	Catalog number
Ê	Recycle
PRINTED WITH SOY INK	Printed with soy ink
CE	CE Mark CE Mark with notified body ID number
	CE Mark with notified body ID number Notified body ID number can vary.
YYYY-MM-DD	Date format (year-month-day)
CHECKSUM	Variable hexadecimal number that ensuring the Master Curve and Calibrator definition values entered an value.
UNITS C	Common Units
UNITS SI	International System of Unio
MATERIAL	Material
MATERIAL ID	Useque material is. Sife ion number
CONTROL NAME	Name a control
CONTROL TYPE	Type of conten

Legal In prelation

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