

TruSight™ Oncology 500 ctDNA

Enabling comprehensive
genomic profiling from liquid
biopsy samples for research

- Leverage minimally invasive blood samples as a complement to tissue biopsy or as an alternative when tissue is not readily available
- Assay DNA biomarkers across > 500 genes plus immuno-oncology signatures such as TMB and MSI
- Realize low limits of detection with UMI-based hybrid-capture library preparation and deep sequencing on the NovaSeq™ 6000 System
- Go from cfDNA to report interpretation in five days with a proprietary DRAGEN™ pipeline and integrated analysis from Pierian



Introduction

Liquid biopsy enables comprehensive analysis of circulating cell-free DNA (cfDNA) in plasma, providing a noninvasive approach for profiling solid tumors. To take advantage of liquid biopsy, it is critical to use a highly sensitive and specific analytical assay capable of detecting somatic mutations at low frequencies. TruSight Oncology 500 ctDNA harnesses the power of proven Illumina next-generation sequencing (NGS) technology to achieve this high analytical sensitivity and enables comprehensive genomic profiling of circulating tumor DNA (ctDNA) found in cfDNA (Figure 1, Table 1). Combining this advanced research solution with the bioinformatics power of the DRAGEN TruSight Oncology 500 ctDNA Analysis Software gives clinical researchers a DNA-to-report solution for evaluating multiple variant types across hundreds of genes in a single assay (Figure 2).


TruSight Oncology 500 ctDNA is compatible with NovaSeq 6000 v1.5 sequencing reagents. In addition to increases in operating efficiencies that result in potential price per sample reductions > 35%, these reagents offer an extended shelf-life of six months and improved Q30 scores.¹

The power of liquid biopsy

Unlike a tissue biopsy that provides information from only a fraction of the tumor, liquid biopsy provides insights about intra- and inter-tumor heterogeneity throughout the body. Studies show that cfDNA analysis detected a significant number of guideline-recommended biomarkers and resistance alterations not found in matched tissue biopsies.² In addition, a non-small cell lung cancer study revealed that cfDNA analyses are highly concordant with tissue-based analyses.³

A foundation of comprehensive content

Content for TruSight Oncology 500 ctDNA was designed with recognized authorities in the oncology community. It includes current and emerging biomarkers with comprehensive coverage of genes involved in key guidelines and clinical trials for multiple tumor types. The panel probe design captures both known and novel gene fusions and includes 523 genes for detecting variants likely to play a role in tumorigenesis. Biomarkers comprise single-nucleotide variants (SNVs), insertions/deletions (indels), copy-number variants (CNVs), gene fusions, and complex immuno-oncology genomic signatures, such as microsatellite instability (MSI) and tumor mutational burden (TMB) (Table 2).

 For a complete list of genes, visit illumina.com/products/by-type/clinical-research-products/trusight-oncology-500-ctdna.html.

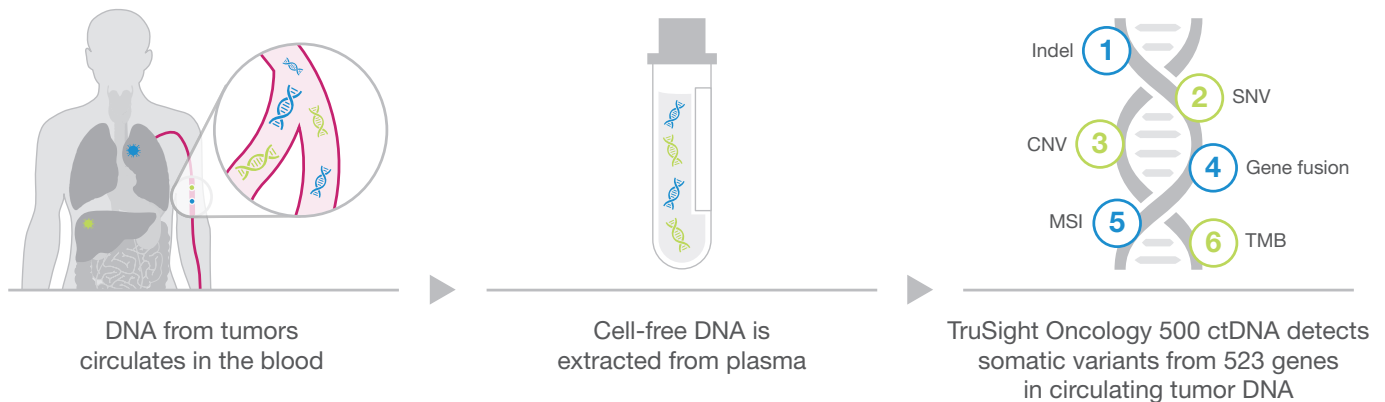


Figure 1: Liquid biopsy enables profiling of biomarkers for multiple variant types and multiple cancer types—Sophisticated variant calling algorithms and high depth of sequencing enable detection of key biomarkers in cfDNA with 0.5% limit of detection (LOD).

Table 1: TruSight Oncology 500 ctDNA at a glance

| Parameter | TruSight Oncology 500 ctDNA |
|---|---|
| System | NovaSeq 6000 System |
| Panel size | 1.94 Mb DNA |
| Panel content | 523 genes 59 genes for CNVs 23 genes for gene fusions MSI (> 2400 loci) TMB |
| DNA input requirement | 30 ng cfDNA ^a |
| Sample type | cfDNA derived from blood |
| Total assay time | 5 days from library prep to variant report |
| Sequence run time | 36 hr run, 10 hr analysis (S2 flow cell) 45 hr run, 22 hr analysis (S4 flow cell) |
| Sequence run | 2 × 151 bp |
| Sample throughput | 8 samples per run (S2 flow cell) 24 samples per run (S4 flow cell) 48 samples per library prep kit |
| Limit of detection | 0.5% VAF for small variants ≥ 1.4-fold change for gene amplifications ≤ 0.6-fold change for gene deletions ≥ 2% tumor fraction for MSI |
| Analytical sensitivity | ≥ 95% (at LOD for all variant types) |
| Analytical specificity | ≥ 95% |
| a. Recommend quantification with Agilent TapeStation or Fragment Analyzer systems | |

Table 2: Examples of variants detected using TruSight Oncology 500 ctDNA

| Variant type | Relevant examples |
|-----------------|----------------------------------|
| SNVs and indels | <i>EGFR, POLE, TMPRSS2, BRAF</i> |
| Gene fusions | <i>ALK, ROS1, NTRK, RET</i> |
| CNVs | <i>HER2</i> |
| MSI | MSI-Score |
| TMB | TMB-Score |

For a complete list of genes, visit illumina.com/products/by-type/clinical-research-products/trusight-oncology-500-ctdna.html

Proven technology for detecting low-level biomarkers

Using proven Illumina sequencing by synthesis (SBS) chemistry, TruSight Oncology 500 ctDNA enables comprehensive genomic profiling from just 30 ng cfDNA, making it an ideal alternative for use when tissue is not readily available or as a complement to tissue analysis. Library preparation takes advantage of target enrichment, using biotinylated probes and streptavidin-coated magnetic beads to enrich for selected targets from DNA-based libraries. Targeted hybridization–capture enrichment uses probes that are large enough to impart high binding specificity, but still allow hybridization to targets containing small mutations. This approach reduces sample dropouts in the presence of both natural allelic variations and sequence artifacts.

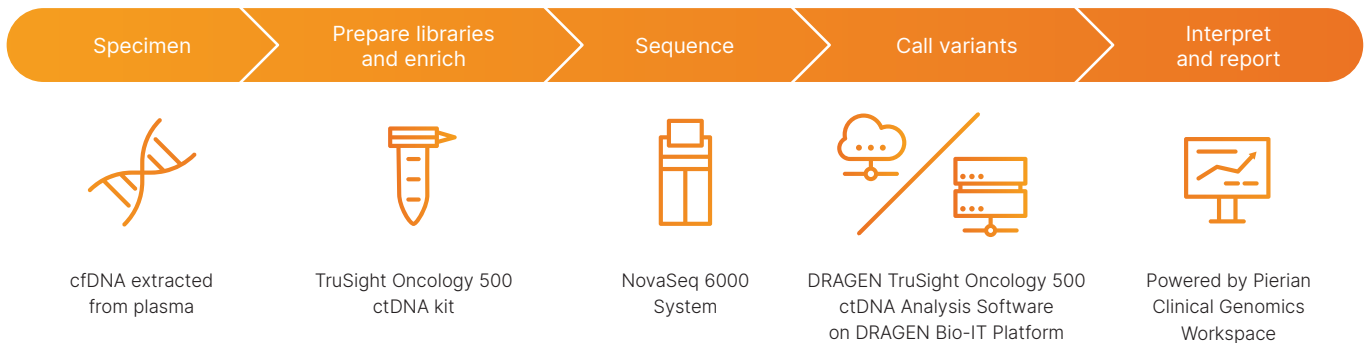


Figure 2: TruSight Oncology 500 ctDNA assay workflow—TruSight Oncology 500 ctDNA assay integrates into current lab workflows, going from cfDNA to a variant report in five days. DRAGEN TruSight Oncology 500 ctDNA Analysis Software can be run locally on a DRAGEN Server or in the cloud via Illumina Connected Analytics.

Because ctDNA represents a small fraction of cfDNA, powerful methods are required to separate signal from noise. Library preparation incorporates unique molecular identifiers (UMIs) that enable ultra-low frequency variant identification.⁴ TruSight Oncology 500 ctDNA libraries are sequenced on the NovaSeq 6000 System at high depth (400M reads per sample at ~35,000x) to enhance sensitivity. The result is the ability to detect mutations at 0.5% variant allele frequency (VAF) for small variants, with 95% analytical sensitivity and > 99.995% analytical specificity (Table 3).

Table 3: Detection of low-level variants with high accuracy

| Variant type | Analytical sensitivity ^a | Analytical specificity ^b |
|---|-------------------------------------|-------------------------------------|
| Small variants (≥ 0.5% VAF) | ≥ 95% | ≥ 99.995% |
| Gene amplifications (≥ 1.4-fold change) | ≥ 95% | ≥ 95% |
| Gene deletions (≤ 0.6-fold change) | ≥ 95% | ≥ 95% |
| Gene fusions (0.5%) | ≥ 95% | ≥ 95% |
| MSI high detection (≥ at 2% tumor fraction) | ≥ 95% | ≥ 95% |

a. Analytical sensitivity is defined as percent detection at the stated variant level
 b. Analytical specificity is defined as the ability to detect a known negative

Accurate, accelerated analysis

DRAGEN TruSight Oncology 500 ctDNA Analysis Software uses accelerated, fully integrated bioinformatics algorithms to ensure optimal assay performance. The software performs sequence alignment, error correction by collapsing the sequence, then variant calling based on the raw data. Duplicated reads and sequencing errors are removed without losing signal for low-frequency variants while yielding high-sensitivity variant calling results. All pipeline components are within the DRAGEN platform, for additional performance improvements and efficiency.

Unlike qualitative results from PCR-based assays, DRAGEN TruSight Oncology 500 ctDNA Analysis Software provides a quantitative MSI score derived from > 2400 homopolymer MSI marker sites. For TMB analysis,

the DRAGEN software optimizes sensitivity by measuring both nonsynonymous and synonymous SNVs and indels. After variant calling and error correction, the accuracy of TMB measurement is further enhanced by filtering germline variants, low-confidence variants, and variants associated with clonal hematopoiesis of indeterminate potential.

DRAGEN TruSight Oncology 500 ctDNA Analysis Software runs locally on an Illumina DRAGEN Server v3 or v4 or in the cloud via Illumina Connected Analytics (ICA). This ultrarapid platform offers enhanced hardware and software that reduce data analysis time by ~85%, or from nine days to ~20 hours (Figure 3). ICA offers labs a secure, cloud-based genomics platform to scale up secondary analysis without the need to acquire and maintain more local infrastructure.⁵ Additionally, DRAGEN TruSight Oncology 500 ctDNA Analysis Software is compatible with DRAGEN v3.10 software, the same version used with the TruSight Oncology 500 assay for analysis of solid tumor tissue. This share platform use offers added flexibility for labs.

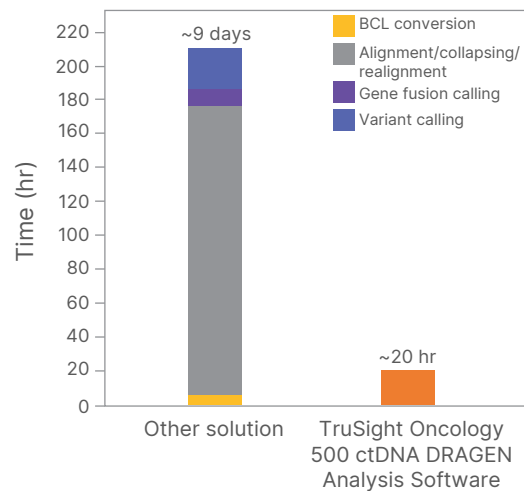


Figure 3: Onsite DRAGEN Server v4 reduces data analysis time—TruSight Oncology 500 ctDNA DRAGEN Analysis Software consolidates various data analysis steps into a single process that requires ~20 hrs, an ~85% reduction compared to another solution. Analysis times compare DRAGEN TruSight Oncology 500 ctDNA v2.1 pipeline on a DRAGEN v4 server for 24 samples using an S4 flow cell to other solution using single node (128G memory, 24 cores CPU), nonparallelized pipeline for 24 samples using an S4 flow cell.

Pierian Clinical Genomics Workspace completes the workflow with tertiary analysis. Simply upload variant report files directly into the Clinical Genomics Workspace cloud from a local or ICA-based secondary analysis environment. Clinical Genomics Workspace performs variant annotation and filtering for smooth interpretation and reporting. From thousands of variants in the genome, Clinical Genomics Workspace filters and prioritizes biologically relevant variants for the final automated, customizable genomic report. The entire workflow, from cfDNA to consolidated variant reporting, takes only five days (Figure 2).

Extensive validation delivers accurate and highly reproducible results

To demonstrate the high-quality results achieved with TruSight Oncology 500 ctDNA, Illumina performed various studies evaluating the ability to call SNVs, CNVs, gene fusions, TMB, and MSI (Figures 4 and 5, Tables 5 and 6).

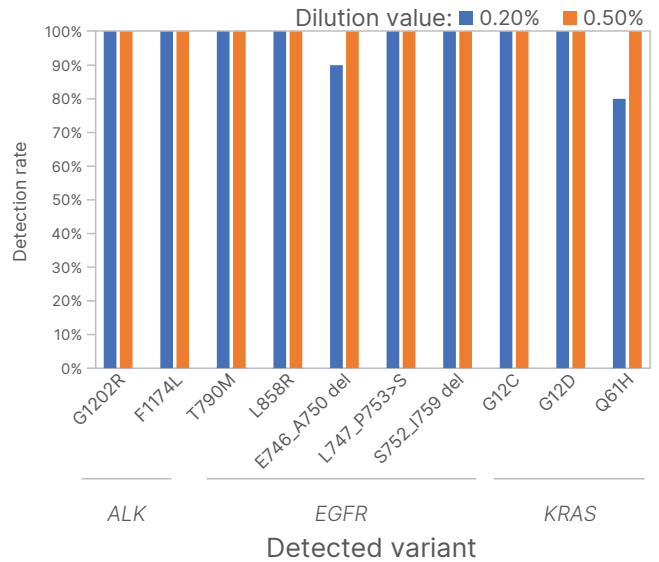


Figure 4: Small variant detection at low VAF—Samples with known VAF for each variant were diluted to values ranging from 0.10–1.00% VAF. Five replicates of each sample were analyzed with TruSight Oncology 500 ctDNA using 30 ng of commercial reference control DNA.

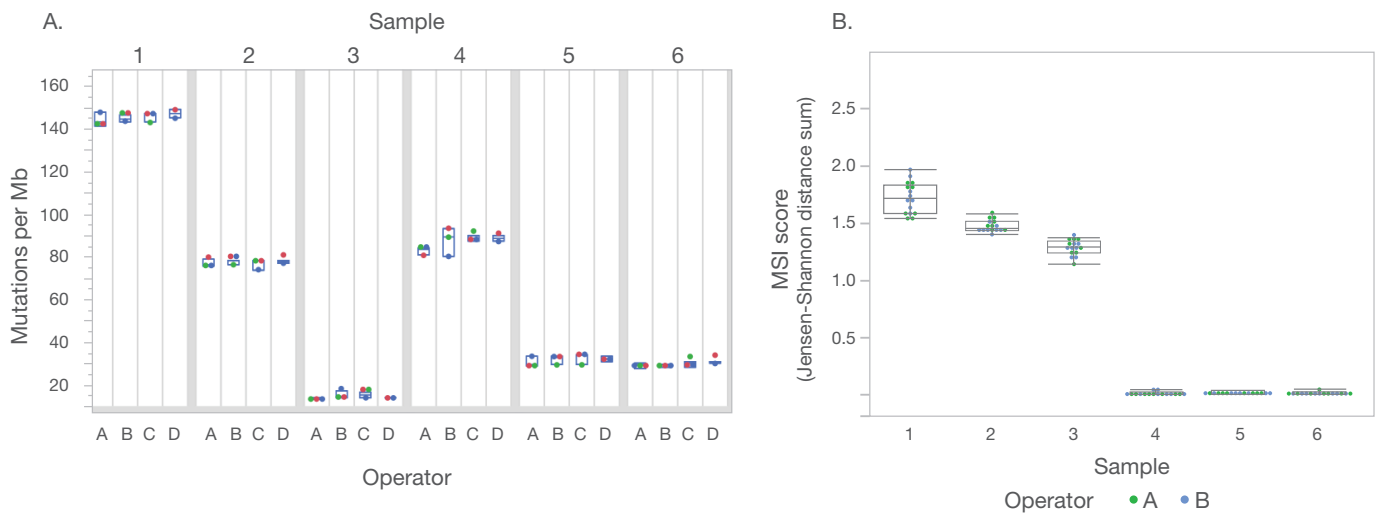


Figure 5: Reproducible TMB and MSI measurement—(A) TMB was evaluated in six different plasma samples (1–6) across four operators (A, B, C, D) in triplicate (green, blue, red dots). (B) MSI was evaluated in three nucleosomal prepped cell lines with known MSI-high status (samples 1–3) and three cfDNA samples from low prevalence MSI-high tumors (samples 4–6) across two different operators (A-green, B-blue).

Table 5: Sensitive detection of CNVs

| Gene | Expected fold-change | Observed mean | Standard deviation | Detection rate |
|-----------------------|----------------------|---------------|--------------------|----------------|
| Amplifications | | | | |
| <i>BRCA2</i> | 1.8 | 1.4 | 0.01 | 100% |
| <i>CCND3</i> | 1.5 | 1.3 | 0.01 | 100% |
| <i>FGF14</i> | 1.3 | 1.5 | 0.01 | 100% |
| <i>FGF3</i> | 1.6 | 1.4 | 0.01 | 100% |
| <i>FGF4</i> | 1.7 | 1.4 | 0.01 | 100% |
| <i>FGFR2</i> | 1.6 | 1.4 | 0.01 | 100% |
| <i>MET</i> | 1.5 | 1.3 | 0.01 | 100% |
| <i>MYC</i> | 1.9 | 1.7 | 0.02 | 100% |
| Deletions | | | | |
| <i>BRCA1</i> | 0.7 | 0.7 | 0 | 100% |
| <i>BRCA2</i> | 0.6 | 0.6 | 0.01 | 100% |

Samples with known fold-changes for gene amplifications and deletions were evaluated using TruSight Oncology 500 ctDNA with 30 ng of cfDNA input. Five replicates of each sample were analyzed.

Table 6: Gene fusion detection at low VAF

| Gene fusion | Expected VAF | Observed VAF | Standard deviation | Detection rate |
|-----------------------|--------------|--------------|--------------------|----------------|
| <i>FGFR2-COL14A1</i> | 4.1% | 4.1% | 0.5% | 100% |
| <i>NPM1-ALK</i> | 3.4% | 0.6% | 0.1% | 100% |
| <i>FGFR3-BAIAP2L1</i> | 3.4% | 0.8% | 0.2% | 100% |
| <i>NPM1-ALK</i> | 2.4% | 0.6% | 0.1% | 100% |
| <i>EML4-ALK</i> | 1.7% | 0.5% | 0.1% | 100% |
| <i>CCDC6-RET</i> | 1.0% | 0.7% | 0.1% | 100% |
| <i>FGFR2-COL14A1</i> | 0.9% | 0.4% | 0.2% | 100% |
| <i>EML4-ALK</i> | 0.7% | 0.2% | 0.1% | 100% |
| <i>EML4-ALK</i> | 0.5% | 0.5% | 0.3% | 100% |
| <i>NPM1-ALK</i> | 0.5% | 0.2% | 0.0% | 100% |
| <i>NCOA4-RET</i> | 0.5% | 0.2% | 0.0% | 100% |
| <i>CCDC6-RET</i> | 0.2% | 0.1% | 0.1% | 100% |

Samples with known gene fusion allele frequencies ranging from ~0.5–4% were evaluated. Five replicates of each sample were analyzed using TruSight Oncology 500 ctDNA with 30 ng cfDNA input. Gene fusion directionality reported based on known expression. Consult the [TruSight Oncology 500 ctDNA Local App User Guide](#) for more information on DNA-based fusion directionality.

Summary

TruSight Oncology 500 ctDNA is an NGS-based, multiplex research assay that analyzes hundreds of cancer-related biomarkers from plasma simultaneously. Assay content is aligned with current guidelines and research from clinical trials. The single, comprehensive assay can detect multiple variant types from 523 genes implicated in various tumor types, without requiring multiple samples for iterative testing. TruSight Oncology 500 ctDNA also provides assessment of immuno-oncology and emerging biomarkers (TMB, MSI, *NTRK*, and *ROS1*). Taking advantage of extensive genomic content, industry-leading sequencing technology, and enhanced software, TruSight Oncology 500 ctDNA provides an integrated solution for accelerating clinical research projects with minimal operational and analysis complexity.

Learn more

TruSight Oncology 500 ctDNA, illumina.com/tso500-ctDNA

NovaSeq 6000 System, illumina.com/systems/sequencing-platforms/novaseq.html

DRAGEN Bio-IT Platform, illumina.com/products/by-type/informatics-products/dragen-bio-it-platform.html

Illumina Connected Analytics, illumina.com/products/by-type/informatics-products/connected-analytics.html

Ordering information

| Product | Catalog no. |
|---|-------------|
| TruSight Oncology 500 ctDNA Kit (48 samples, 16 indexes) | 20039252 |
| TruSight Oncology 500 ctDNA Kit plus Pierian Interpretation Report (48 samples, 16 indexes) | 20043410 |
| Sequencing reagent kits | |
| NovaSeq 6000 S2 Reagent Kit v1.5 (300 cycles) | 20028314 |
| NovaSeq 6000 S4 Reagent Kit v1.5 (300 cycles) | 20028312 |
| NovaSeq Xp 4-Lane Kit v1.5 | 20043131 |
| On-premise variant reporting | |
| Illumina DRAGEN Server v4 | 20051343 |
| Illumina DRAGEN Server Advance Exchange Plan | 20032797 |
| Illumina DRAGEN Server Installation | 20031995 |
| Cloud-based variant reporting | |
| ICA Basic Annual Subscription | 20044874 |
| ICA Professional Annual Subscription | 20044876 |
| ICA Enterprise Annual Subscription | 20038994 |
| ICA Enterprise Compliance Add-on (applies to Basic only) | 20066830 |
| ICA Training and Onboarding | 20049422 |
| ICA Data Storage: Illumina Analytics, 1 credit | 20042038 |
| ICA Data Storage: Illumina Analytics Starter Pack, 1000 credits | 20042039 |
| ICA Data Storage: Illumina Analytics, 5000 credits | 20042040 |
| ICA Data Storage: Illumina Analytics, 50,000 credits | 20042041 |
| ICA Data Storage: Illumina Analytics, 100,000 credits | 20042042 |

References

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3. Leighl NB, Page RD, Raymond VM, et al. [Clinical Utility of Comprehensive Cell-free DNA Analysis to Identify Genomic Biomarkers in Patients with Newly Diagnosed Metastatic Non-small Cell Lung Cancer](#). *Clin Cancer Res*. 2019;25(15):4691-4700. doi:10.1158/1078-0432.CCR-19-0624
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5. Illumina. Illumina Connected Analytics Security Brief. [illumina.com/content/dam/illumina/gcs/assembled-assets/marketing-literature/ica-security-brief-m-gl-00683/ica-security-brief-m-gl-00683.pdf](https://www.illumina.com/content/dam/illumina/gcs/assembled-assets/marketing-literature/ica-security-brief-m-gl-00683/ica-security-brief-m-gl-00683.pdf). Published 2022. Accessed March 16, 2022.



1.800.809.4566 toll-free (US) | +1.858.202.4566 tel
techsupport@illumina.com | www.illumina.com

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M-GL-00843 v4.0