

TruSight™ Oncology 500 ctDNA v2

Enabling sensitive, fast CGP
from liquid biopsy samples

- Detect biomarkers present as low as 0.2% VAF from 20 ng ctDNA (5–30 ng possible)
- Obtain comprehensive results in < 4 days with manual or automated options
- Analyze > 500 genes and immuno-oncology (IO) genomic signatures (MSI, TMB) in one assay
- Leverage transformative economics and batch sizes as low as four samples with the NovaSeq™ X Series



The value ctDNA and liquid biopsy bring to CGP

Understanding the genomic basis of cancer can help pinpoint alterations that fuel the disease and enable advancements in precision medicine. One method for approaching these oncology studies is comprehensive genomic profiling (CGP). CGP is a precision medicine application that takes advantage of next-generation sequencing (NGS) to assess a wide range of biomarkers in a single assay, using less sample and returning results faster than multiple, iterative testing strategies.^{1,2} In addition, CGP tests can identify more clinically relevant variants than conventional testing approaches, such as single-gene tests and hotspot NGS panels.³⁻⁶ This ability to detect more variants grows in importance as an increasing number of biomarkers are discovered, including immuno-oncology (IO) genomic signatures such as tumor mutational burden (TMB) that require large NGS panels (> 1 Mb) for accurate identification.^{7,8}

The standard approach for CGP involves use of solid tumor tissue samples, including formalin-fixed paraffin-embedded (FFPE) samples. Occasionally, sufficient tissue sample may be unavailable (this can occur up to 25% of the time⁹), the tumor may be inaccessible, or results from tissue biopsy may be too delayed. In these cases, performing CGP with circulating tumor DNA (ctDNA) from a blood-based liquid biopsy can provide insights into the genomic landscape of the tumor. ctDNA has been detected through all stages of cancer progression and across multiple solid tumor types,¹⁰ including lung, breast, colorectal, and ovarian cancers. Using ctDNA for CGP has several advantages:

- Access sample easily through a minimally invasive blood draw procedure¹¹
- Capture clones from multiple tumors or even the same tumor,¹² overcoming inherent sampling bias present with solid tumor biopsy and expanding the ability to identify more alterations¹²⁻¹⁵
- Obtain temporal and spatial information about intra- and inter-tumor heterogeneity¹¹
- Repeat analysis to assess clonal selection

Liquid biopsy provides a noninvasive approach to obtaining cell-free DNA (cfDNA), which includes ctDNA, from blood plasma for tumor profiling (Figure 1). For certain diseases, such as non-small cell lung carcinoma (NSCLC), adding CGP analysis from liquid biopsy to tissue analysis can increase the identification of clinically relevant mutations by 15–48%.^{13,14,16} In addition, non-small cell lung cancer studies have revealed that cfDNA analyses are highly concordant with tissue-based analyses.¹⁴ Today, liquid biopsy is increasingly being included in professional guidelines (> 12 diseases) as a method for obtaining sample for molecular profiling.¹⁷⁻¹⁹

To benefit from liquid biopsy, it is critical to use a highly sensitive and specific analytical assay capable of detecting somatic mutations present at low frequencies in ctDNA. The original TruSight Oncology 500 ctDNA assay²⁰ met this challenge, harnessing the power of proven Illumina NGS technology and achieving the high analytical sensitivity needed to enable CGP. Building on this success, TruSight Oncology 500 ctDNA v2 (Table 1, Table 2) offers chemistry and workflow improvements that lead to higher sensitivity and a faster time to answer (Table 3).

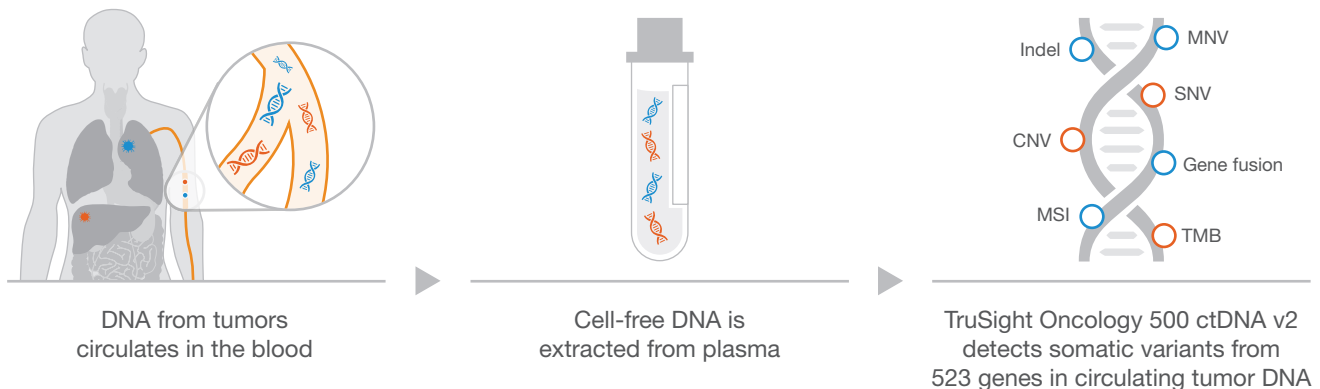


Figure 1: Liquid biopsy enables a noninvasive approach for CGP—ctDNA, found in cell-free DNA in plasma, can be obtained through a simple blood draw analyzed using TruSight Oncology 500 ctDNA v2 to detect the presence of cancer-relevant biomarkers in key guidelines.

Table 1: TruSight Oncology ctDNA v2 at a glance

Parameter	Specification
System	NovaSeq X Series NovaSeq 6000 System NovaSeq 6000Dx Instrument (RUO mode) ^a
Sample throughput	4–48 samples per run
Library preparation kit sizes	24-sample (manual) 48-sample (automated)
Samples per flow cell	4 samples per 1.5B flow cell 8 samples per S2 flow cell 24 samples per 10B flow cell 24 samples per S4 flow cell
Automation capability	Illumina Qualified methods available for the Hamilton Microlab STAR
Panel size	1.94 Mb DNA
Panel content	523 genes for small variants 59 genes for CNVs 23 genes for gene rearrangements MSI (> 2300 loci) TMB (> 1 Mb)
Sample type	cfDNA derived from blood plasma
DNA input requirement	20 ng cfDNA (5–30 ng possible) ^b
Total assay time	From library prep to variant report: • 3 days on the NovaSeq X Series • 4 days on the NovaSeq 6000 System
Hands-on time	8–24 samples (manual): ~2.5 h 8–24 samples (automated): ~1.5 h 48 samples (manual): ~4.5 h 48 samples (automated): ~1.5 h
Library preparation time ^c	8–24 samples (manual): ~8.5 h 8–24 samples (automated): ~9.5 h 48 samples (manual): ~10 h 48 samples (automated): ~11 h
Sequence run time	22–44 h (see Table 5)
Sequence run read length	2 × 151 bp
Sequencing coverage	35,000×
Variant analysis time	8 samples: 9–12 h 24 samples: 20–24 h 48 samples: 40–48 h

a. Onboard server compatibility for the NovaSeq 6000Dx Instrument coming 2H 2024.

b. Recommend quantification with Agilent TapeStation or Fragment Analyzer systems. To learn more about input amounts, read the [Using lower input amounts with TruSight Oncology ctDNA v2 technical note](#).

c. Includes library preparation, enrichment, and bead-based normalization.

Table 2: TruSight Oncology ctDNA v2 performance^a

Parameter	Specification
Limit of detection (LOD)	0.2% VAF for SNVs 0.5% VAF for MNVs and indels 0.5% VAF for gene rearrangements ≥ 1.3-fold change for gene amplifications ≤ 0.6-fold change for gene deletions ≥ 0.3% tumor fraction for MSI
Analytical sensitivity (at LOD)	≥ 90% (at LOD of 0.2% VAF for SNVs) ≥ 95% (at LOD of 0.2% VAF for SNV hotspots) ≥ 95% (at LOD of 0.5% VAF for all other variant types)
Analytical specificity	≥ 99.999%

a. Verified on the NovaSeq 6000 System and NovaSeq X Series.

Table 3: Advances with TruSight Oncology ctDNA v2

Benefit	TruSight Oncology 500 ctDNA v2	TruSight Oncology 500 ctDNA (original)
Improved assay sensitivity	Separate end-repair and A-tailing steps	Combined end-repair and A-tailing
More streamlined workflow and improved user experience	Plate-based indexes/UMI	Tube-based indexes/UMI
Faster, single day workflow	Single hybridization/capture step	Two hybridization and capture steps
More scalability	192 indexes	16 indexes
Broader batch sizes	4–48 samples ^a	8–48 samples
Automation enabled	Yes	No

a. Batches of four samples available on the NovaSeq X Series. UMI, Unique molecular identifier.

Comprehensive content

Content for TruSight Oncology 500 ctDNA v2 was designed with recognized authorities in the oncology community and 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 rearrangements and includes 523 genes for detecting variants likely to play a role in tumorigenesis now and into the future. Biomarkers comprise single-nucleotide variants (SNVs), multi-nucleotide variants (MNVs), insertions/deletions (indels), copy-number variants (CNVs), gene rearrangements, and complex IO genomic signatures, such as blood-based microsatellite instability (bMSI) and blood-based TMB (bTMB) (Table 4).


 For a complete list of genes, visit the [TruSight Oncology 500 ctDNA v2 product page](#).

Table 4: Examples of variant types detected by TruSight Oncology 500 ctDNA v2

Variant type	Example
SNVs and indels	<i>EGFR, POLE, TMPRSS2, BRAF</i>
Gene rearrangements	<i>ALK, ROS1, NTRK1, NTRK2, RET</i>
CNVs	<i>HER2</i>
MSI	MSI-Score
TMB	TMB-Score

Fast, integrated workflow

TruSight Oncology 500 ctDNA v2 is part of an integrated CGP workflow that spans from sample input to final report (Figure 2). Using automated library preparation kits and methods, variant calling tools, and interpretation and reporting software enables a smooth workflow that can be completed in less than four days, less than half the time of other CGP liquid biopsy assays (Figure 3).

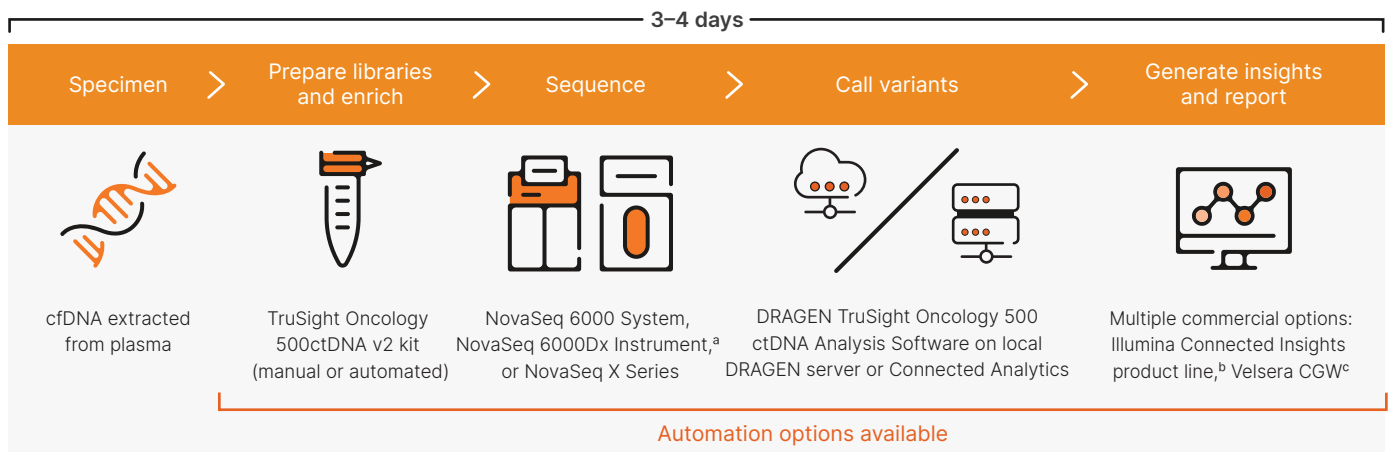


Figure 2: TruSight Oncology 500 ctDNA v2 workflow—TruSight Oncology 500 ctDNA v2 integrates into current lab workflows, going from cfDNA to a variant report in less than four days. DRAGEN TruSight Oncology 500 ctDNA Analysis Software runs locally on a DRAGEN Server or in the cloud version via Illumina Connected Analytics. a. NovaSeq 6000Dx Instrument in RUO mode. b. Available in select countries. Illumina Connected Insights product line supports user-defined tertiary analysis through API calls to third-party knowledge sources. c. Other third-party options are available.

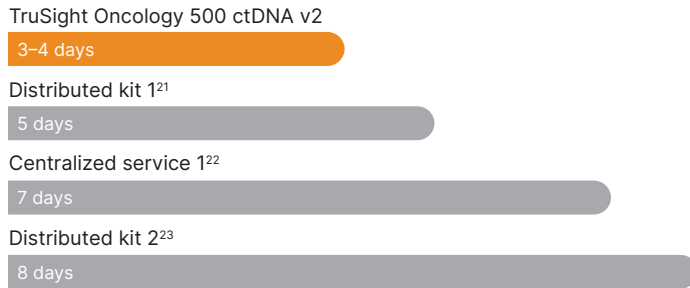


Figure 3: Faster time to report with TruSight Oncology 500 ctDNA v2—A comparison of the time required to go from sample to report for liquid biopsy CGP assays that include the IO biomarkers of bMSI and bTMB.

Optimized library preparation

Using proven Illumina sequencing by synthesis (SBS) chemistry, TruSight Oncology 500 ctDNA v2 enables comprehensive genomic profiling from just 20 ng cfDNA, making it an ideal alternative for use when tissue is not readily available or as a complement to tissue analysis. ctDNA represents a small fraction of cfDNA (often < 5% of total cfDNA), requiring powerful methods to separate signal from noise. To enable ultra-low frequency variant identification, library preparation takes advantage of target enrichment with biotinylated probes and streptavidin-coated magnetic beads to enrich for selected targets from DNA-based libraries and unique molecular identifiers (UMIs)²⁴ to reduce error rates (Figure 4). Advances in product chemistry have decreased the number of hybridizations from two down to one in TruSight Oncology 500 ctDNA v2, allowing for a one-day turnaround for library preparation and a shorter time to results. It has also improved analytical sensitivity down to 0.2% VAF for SNVs. This targeted hybridization–capture approach reduces sample dropouts in the presence of both natural allelic variations and sequence artifacts.



Learn about using input amounts as low as 5 ng with TruSight Oncology ctDNA v2.



Learn about UMIs.

Automation-enabled workflow

TruSight Oncology 500 ctDNA v2 offers manual and automated options to support scalable library prep. Illumina has partnered with Hamilton, a leading liquid-handling manufacturer, to produce a fully automated workflow for TruSight Oncology 500 ctDNA v2 assays on the Hamilton Star. Automation-friendly library preparation kits provide sufficient reagent for preparing 48 libraries and accommodating the dead volume required for the robot with minimal reagent waste. Automated workflows achieve the same high-quality results produced by manual protocols, while reducing hands-on time by 40%, freeing up approximately eight hours for other laboratory activities. Overall, automation enables labs to save on labor costs and improve efficiency.²⁶

Powerful sequencing

TruSight Oncology 500 ctDNA v2 libraries are sequenced on the NovaSeq 6000 Sequencing System, NovaSeq 6000Dx Instrument (RUO mode),* or NovaSeq X Series. The NovaSeq X Series offers a faster workflow, shortening sequencing time by ~40% compared to the NovaSeq 6000 System (Table 5). Regardless of the platform used, sequencing occurs at high depth (400M reads per sample at ~35,000x) to enhance sensitivity. The result is the ability to detect mutations at 0.2% variant allele frequency (VAF) for SNVs, with ≥ 90% analytical sensitivity and ≥ 95% analytical specificity (Table 6). In addition, the NovaSeq X Series offers transformative economics by reducing the cost of sequencing per sample.²⁷

* Sequencing on the NovaSeq 6000Dx Instrument available H2 2024.

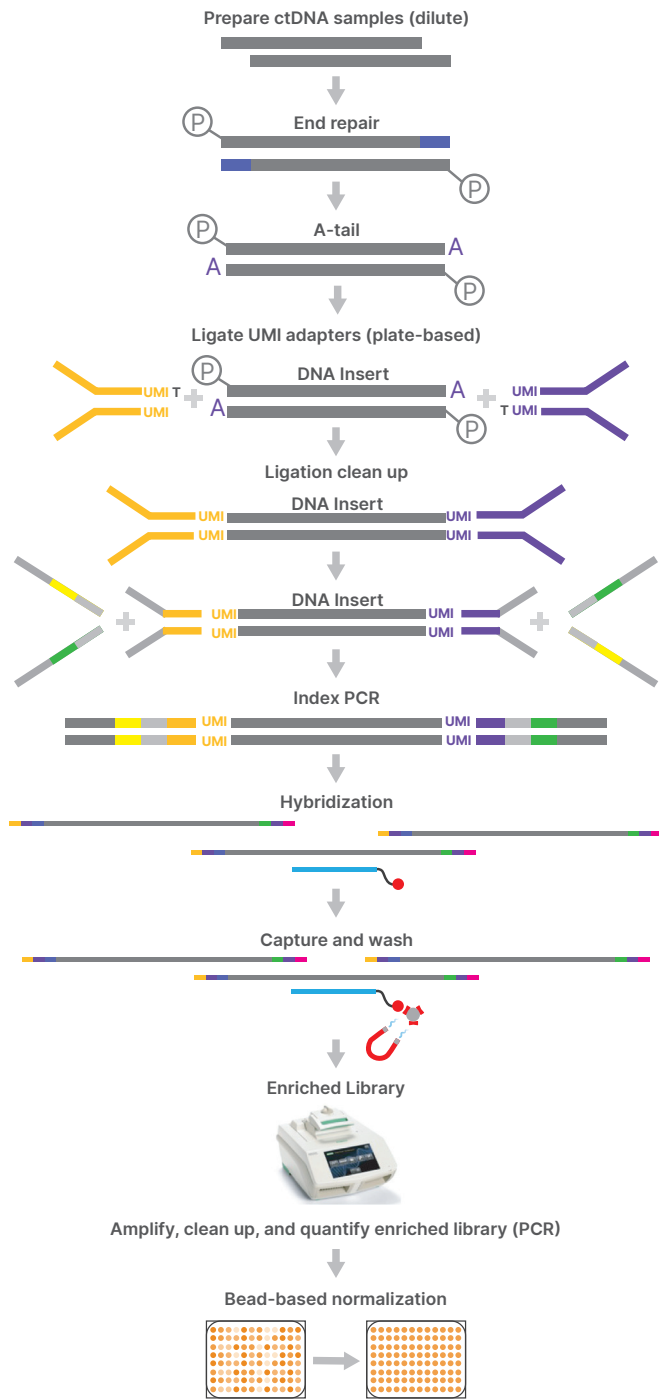


Figure 4: UMI-based hybrid-capture library preparation— Targeted enrichment uses probes large enough to impart high binding specificity, but still allow hybridization to targets containing mutations. UMI reagents reduce error rates, increasing analytical specificity and yielding higher confidence variant calls.²⁵

Table 5: Estimated sequencing run times

System	NovaSeq 6000 System ^a		NovaSeq X Series	
	S2	S4	1.5B	10B
Flow cell				
No. of samples	Sequencing run time (No. of flow cells)			
4	—	—	22 h (1)	—
8	36 h (1)	—	22 h (2)	—
24	—	44 h (1)	—	25 h (1)
48	—	44 h (2)	—	25 h (2)

a. Sequencing run times apply to the NovaSeq 6000Dx Instrument in RUO mode.

Table 6: Accurate detection of low-level biomarkers^a

Variant type	Analytical sensitivity ^b	Analytical specificity ^c
Small nucleotide variants (≥ 0.2% VAF)	≥ 90%	≥ 99.9994%
Multi-nucleotide variants (≥ 0.5% VAF)	≥ 90%	≥ 95%
Insertions/deletions (≥ 0.5% VAF)	≥ 90%	≥ 95%
Gene amplifications (≥ 1.3-fold change)	≥ 95%	≥ 95%
Gene deletions (≤ 0.6-fold change)	≥ 95%	≥ 95%
Gene rearrangements (≥ 0.5%)	≥ 95%	≥ 95%
MSI high detection (≥ at 0.3% tumor fraction)	≥ 95%	≥ 95%

a. Performance verified on the NovaSeq 6000 System and NovaSeq X Series.

b. Analytical sensitivity is defined as percent detection at the stated variant level.

c. Analytical specificity is defined as the ability to detect a known negative.

Accurate, accelerated analysis

Comprehensive, efficient variant calling

The DRAGEN™ TruSight Oncology 500 ctDNA Analysis pipeline uses accelerated, fully integrated bioinformatics algorithms to perform 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.

Unlike qualitative results from PCR-based assays, DRAGEN TruSight Oncology 500 ctDNA Analysis pipeline provides a quantitative bMSI score derived from > 2300 homopolymer MSI marker sites. For bTMB analysis, the DRAGEN pipeline optimizes sensitivity by measuring both nonsynonymous and synonymous SNVs and indels. After variant calling and error correction, the accuracy of bTMB 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 pipeline runs locally on an Illumina DRAGEN Server v4 or in the cloud via Illumina Connected Analytics (ICA). ICA offers options for automated data transfer and analysis launch, as well as a secure, cloud-based genomics platform to scale secondary analysis without the need to acquire and maintain local infrastructure.²⁸ Enhanced DRAGEN hardware and software that reduce data analysis time by ~85% (Table 7).

Table 7: Reduced time for data analysis for 24 samples using an S4 flow cell

Data analysis step	Solution A ^a	DRAGEN TruSight Oncology 500 ctDNA Analysis pipeline
BCL conversion	6 hr	1 hr
Alignment + collapsing + realignment	170 hr	11 hr
Gene rearrangement calling	10 hr	2 hr
Variant calling	24 hr	8 hr
Total time	~9 days	~20 hr (~85% reduction)

a. Single node (128G memory, 24 cores CPU), nonparallelized pipeline.

Streamlined data interpretation

After variant class and biomarker type identification via secondary analysis, the next step is to interpret the data to extract biologically relevant meaning. Illumina Connected Insights,[†] Velsera Clinical Genomics Workspace, and third-party apps can be used.

Variant calling files produced locally or via the cloud with Illumina Connected Analytics can be automatically ingested into Illumina Connected Insights. When combined with sequencing system integration and the autolaunch capabilities of Connected Analytics, the analysis workflow can be fully automated with Connected Insights, removing the need for manual data transfers, resulting in a final customizable report.

Reliable, reproducible results

TruSight Oncology 500 ctDNA v2 provides sensitive detection of genomic variants and biomarkers in a cfDNA sample, even when present at low levels. To demonstrate the high-quality results achieved with TruSight Oncology 500 ctDNA v2, Illumina performed various studies evaluating the ability to call small DNA variants, CNVs, gene rearrangements, TMB, and MSI. Performance results were verified on the NovaSeq 6000 System and NovaSeq X Series.

SNVs and indels

One benefit of target enrichment chemistry is the use of probes designed large enough to impart high binding specificity, but also allow hybridization to targets containing small mutations. As SNVs have been associated with cancer susceptibility in various cancer types, it is critical that any CGP method can detect these variants at low levels. TruSight Oncology 500 ctDNA v2 reproducibly detects SNVs and indels present at levels as low as 0.2% or 0.5% VAF, respectively (Figure 5 and Figure 6).

[†] Not available in all countries. Illumina Connected Insights supports user-defined tertiary analysis through API calls to third-party knowledge sources.

CNVs

Copy-number changes in genes and tumor types can be associated with tumorigenesis.²⁸ TruSight Oncology 500 ctDNA v2 includes analysis of 59 CNV-associated genes and can call amplifications with a limit of detection at ≥ 1.3 fold for amplifications and ≤ 0.6 for deletions (Table 8).

Table 8: TruSight Oncology 500 ctDNA v2 analytical performance for CNVs

Gene	Expected fold-change	Observed fold-change	Detection rate
Amplifications			
<i>ERBB2</i>	1.5	1.50	100%
<i>MET</i>	1.5	1.55	100%
<i>MYC</i>	1.5	1.27	100%
<i>ERBB2</i>	1.4	1.73	100%
<i>MET</i>	1.4	1.46	100%
<i>MYC</i>	1.4	1.22	100%
<i>ERBB2</i>	1.3	1.35	100%
<i>MET</i>	1.3	1.38	100%
<i>MYC</i>	1.3	1.19	8%
<i>ERBB2</i>	1.2	1.19	100%
<i>MET</i>	1.2	1.22	100%
<i>MYC</i>	1.2	N/A	0
Deletions			
<i>BRCA1</i>	0.85	0.86	16%
<i>BRCA2</i>	0.85	N/A	0
<i>BRCA1</i>	0.80	0.79	100%
<i>BRCA2</i>	0.80	0.80	100%
<i>BRCA1</i>	0.70	0.69	100%
<i>BRCA2</i>	0.70	0.69	100%

Samples with known fold changes for gene amplifications using synthetic controls and cell lines for deletions were evaluated using TruSight Oncology 500 ctDNA v2. CNVs were diluted to three VAF levels. LOD ≥ 1.3 fold change for gene amplifications ≤ 0.6 for deletions. Note the strong correlation between expected and observed fold changes. Data shown are for sequencing on the NovaSeq 6000 System; similar performance was observed on the NovaSeq X Series.

Gene rearrangements

Gene rearrangements can act as genomic drivers for cancer, making the ability to detect them essential to studies focusing on understanding the foundation of the disease. TruSight Oncology 500 ctDNA v2 detects and characterizes gene rearrangements agnostic from the partner, even when present at low concentrations (Table 9).

Table 9: TruSight Oncology 500 ctDNA v2 analytical performance for gene rearrangements

Fusion	Expected VAF	Observed VAF	Detection rate
<i>ALK:EML4</i>	0.60%	0.48%	100%
<i>GOPC;ROS1:CD74</i>	0.60%	0.39%	100%
<i>RET:NCOA4</i>	0.60%	0.31%	100%
<i>ALK:EML4</i>	0.50%	0.43%	100%
<i>GOPC;ROS1:CD74</i>	0.50%	0.33%	100%
<i>RET:NCOA4</i>	0.50%	0.27%	100%
<i>ALK:EML4</i>	0.40%	0.36%	100%
<i>GOPC;ROS1:CD74</i>	0.40%	0.24%	100%
<i>RET:NCOA4</i>	0.40%	0.19%	100%
<i>ALK:EML4</i>	0.20%	0.18%	88%
<i>GOPC;ROS1:CD74</i>	0.20%	0.11%	100%
<i>RET:NCOA4</i>	0.20%	0.12%	83%

Samples with three known DNA fusions diluted to VAF levels ranging from 0.2% to 0.6% were evaluated with TruSight Oncology 500 ctDNA v2. The LOD for gene rearrangements = 0.5%. NGS-based assessment using TruSight Oncology 500 ctDNA v2 interrogates > 2300 homopolymer sites sized 6–7 bp, which helps reduce error rates and decrease potential false positives commonly found in homopolymer sequencing. Data shown are for sequencing on the NovaSeq 6000 System; similar performance was observed on the NovaSeq X Series. VAF, variant allele frequency.

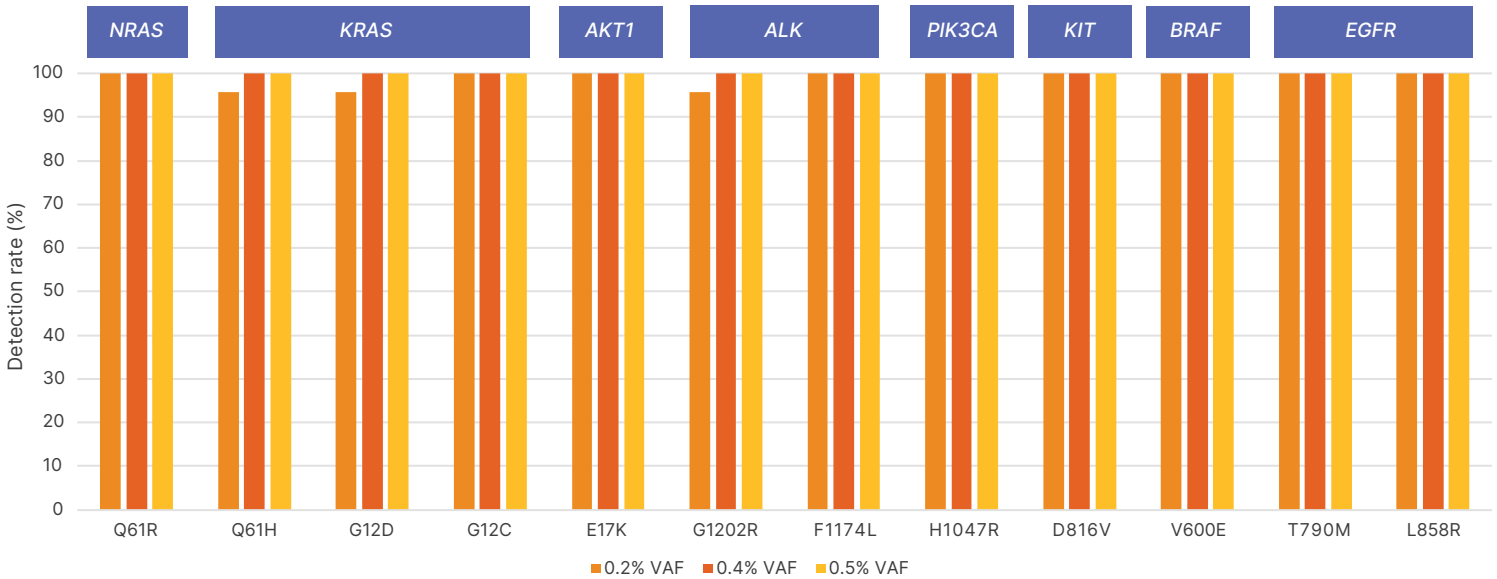


Figure 5: High analytical performance for key SNVs at LOD (0.2% VAF)—Synthetic control samples with known VAF for each single nucleotide variant were diluted to values ranging from 0.20%–0.50% VAF and analyzed using TruSight Oncology 500 ctDNA v2. SNVs present at levels as low as 0.2% were detectable. Data shown are for sequencing on the NovaSeq 6000 System; similar performance was observed on the NovaSeq X Series.



Figure 6: High analytical performance for indels at LOD (0.5% VAF)—Synthetic control samples with known VAF for each insertion or deletion were diluted to values ranging from 0.20%–0.50% VAF and analyzed by TruSight Oncology 500 ctDNA v2. *BRCA1* detection was lower due to the variant being in a highly homopolymer region, resulting in a high level of background noise. Data shown are for sequencing on the NovaSeq 6000 System; similar performance was observed on the NovaSeq X Series.

IO gene signatures: MSI and TMB

MSI and TMB detection relies upon analysis of multiple genomic loci. NGS-based assessment with the TruSight Oncology 500 ctDNA v2 interrogates > 2300 homopolymer sites sized 6–7 bp, which helps reduce error rates and decrease potential false positives commonly found in homopolymer sequencing.²⁸ Featuring sensitive library preparation chemistry combined with advanced bioinformatics, TruSight Oncology 500 ctDNA v2 provides MSI detection achieved down to 0.3% tumor fraction (Figure 7).

Obtaining a precise and reproducible bTMB value at low mutation levels can be challenging with smaller panels.⁷ TruSight Oncology 500 ctDNA v2 combines comprehensive genomic content with a 1.94 Mb panel and sophisticated informatics algorithms to provide accurate bTMB estimations. The proprietary DRAGEN TruSight Oncology 500 ctDNA bioinformatics pipeline applies advanced filtering for both germline and clonal hematopoiesis variants, resulting in a highly concordant tumor-only and tumor-normal workflows ($R^2 = 0.992$) (Figure 8).²⁹

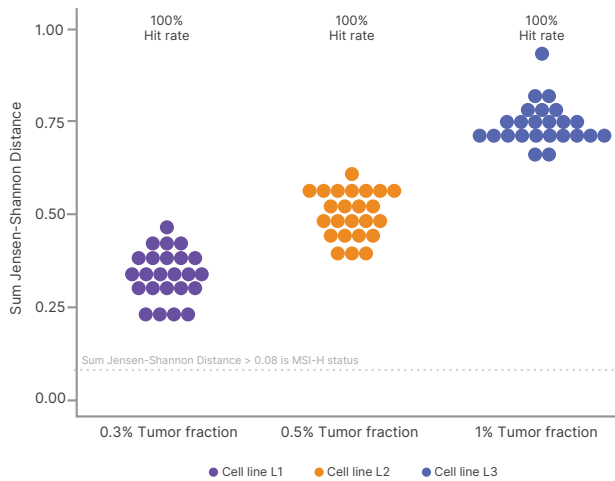


Figure 7: Sensitive MSI performance for IO research—Tumor fractions produced by titrating nucleosomal prepared cell lines with previously known MSI-H scores titrated into wildtype cell background. High MSI analytical sensitivity is achieved with proprietary DRAGEN TruSight Oncology 500 ctDNA v2.1 Analysis Software. More than 2300 homopolymer sites were assessed. Data shown are for sequencing on the NovaSeq 6000 System. Similar performance was observed on the NovaSeq X Series.

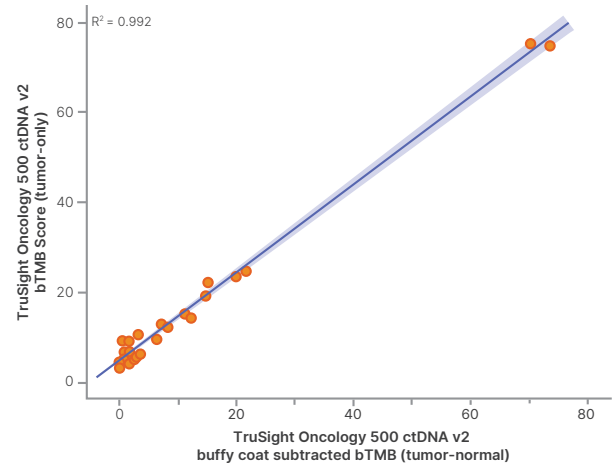


Figure 8: High bTMB data correlation between tumor-only and tumor-normal analysis workflows—Tumor-only bTMB scores produced with TruSight Oncology 500 ctDNA v2 with advanced bioinformatics and a panel large enough to detect TMB (> 1 Mb) show high concordance to bTMB scores produced from a paired tumor-normal workflow using plasma and buffy coat cfDNA. Data shown are for sequencing on the NovaSeq 6000 System; similar performance was observed on the NovaSeq X Series.

Enhanced product attributes

Illumina offers high levels of service and support to ensure operational success for laboratories. To enable greater efficiency, TruSight Oncology 500 ctDNA v2 features:

- Advanced change notification—Illumina notifies laboratories six months before any significant changes are made to TruSight Oncology 500 ctDNA
- Certificate of Analysis[‡]—Every TruSight Oncology 500 ctDNA v2 is issued with a certificate of analysis (CoA) by the Illumina Quality Assurance Department that ascertains the product has met its predetermined product release specifications and quality
- Extended shelf life—The minimum guaranteed shelf life for TruSight Oncology 500 ctDNA v2 reagents is extended to six months, reducing the risk of product expiration and enabling labs to use reagents according to current testing needs
- Single-lot shipments—Available for manual kits now and coming in H2 2024 for automation kits, single-lot shipments reduce the burden of lot qualification and incoming QC testing

Integrated solution enabling CGP from liquid biopsy

TruSight Oncology 500 ctDNA v2 is an NGS-based, multiplex research assay that analyzes hundreds of cancer-related biomarkers aligned with current guidelines and research from clinical trials from plasma samples simultaneously. The comprehensive assay detects multiple variant types in blood from 523 genes implicated in various tumor types and assesses IO and emerging biomarkers (bTMB, bMSI, *NTRK*, and *ROS1*), without requiring multiple samples for iterative testing.

Chemistry improvements and expanded sequencing system compatibility have decreased the overall turnaround time to 3–4 days, reduced the input requirement to 20 ng cfDNA, and lowered the limit of detection to 0.2% VAF (for SNVs). In addition, the automation-enabled workflow reduces hands-on time and minimizes the burden on lab personnel, creating a streamlined lab for greater efficiency. Taking advantage of extensive genomic content, industry-leading sequencing technology, and enhanced software, TruSight Oncology 500 ctDNA v2 provides an integrated solution that enables CGP-based clinical research projects with minimal operational and analysis complexity.

Learn more

[TruSight Oncology 500 ctDNA v2](#)

[NovaSeq X Series](#)

[NovaSeq 6000 System](#)

[NovaSeq 6000Dx Instrument](#)

[DRAGEN secondary analysis](#)

[Illumina Connected Analytics](#)

[Illumina Connected Insights](#)

[‡] CoA available in 2024.

Ordering information—Library preparation kits (manual)

Product	Catalog no.
TruSight Oncology 500 ctDNA v2 (24 samples)	20105899
TruSight Oncology 500 ctDNA v2 for use with NovaSeq 6000 S2 (24 samples)	20105901
TruSight Oncology 500 ctDNA v2 for use with NovaSeq 6000 S4 (24 samples)	20105902
TruSight Oncology 500 ctDNA v2 (24 samples) plus Velsera Interpretation Report	20105905
TruSight Oncology 500 ctDNA v2 plus Velsera Interpretation Report, for use with NovaSeq 6000 S2 (24 samples)	20105907
TruSight Oncology 500 ctDNA v2 plus Velsera Interpretation Report, for use with NovaSeq 6000 S4 (24 samples)	20105908
TruSight Oncology 500 ctDNA v2 (24 samples) plus Connected Insights Interpretation Report	20105911
TruSight Oncology 500 ctDNA v2 plus Connected Insights Interpretation Report, for use with NovaSeq 6000 S2 (24 samples)	20105913
TruSight Oncology 500 ctDNA v2 plus Connected Insights Interpretation Report, for use with NovaSeq 6000 S4 (24 samples)	20105914

Ordering information—Library preparation kits (automated)

Product	Catalog no.
TruSight Oncology 500 ctDNA v2 for Automation (48 samples)	20105900
TruSight Oncology 500 ctDNA v2 Automation Kit, for use with NovaSeq 6000 S2 (48 samples)	20105903
TruSight Oncology 500 ctDNA v2 Automation Kit, for use with NovaSeq 6000 S4 (48 samples)	20105904
TruSight Oncology 500 ctDNA v2 for Automation (48 samples) plus Velsera Interpretation Report	20105906
TruSight Oncology 500 ctDNA v2 Automation Kit plus Velsera Interpretation Report, for use with NovaSeq 6000 S2 (48 samples)	20105909
TruSight Oncology 500 ctDNA v2 Automation Kit plus Velsera Interpretation Report, for use with NovaSeq 6000 S4 (48 samples)	20105910
TruSight Oncology 500 ctDNA v2 for Automation (48 samples) plus Connected Insights Interpretation Report	20105912
TruSight Oncology 500 ctDNA v2 Automation Kit plus Connected Insights Interpretation Report, for use with NovaSeq 6000 S2 (48 samples)	20105915
TruSight Oncology 500 ctDNA v2 Automation Kit plus Connected Insights Interpretation Report, for use with NovaSeq 6000 S4 (48 samples)	20105916

Ordering information—Index adapters

Product	Catalog no.
IDT for Illumina UMI DNA/RNA UD Indexes Set A, Ligation (96 Indexes, 96 Samples)	20034701
IDT for Illumina UMI DNA/RNA UD Indexes Set B, Ligation (96 Indexes, 96 Samples)	20034702
IDT for Illumina UMI DNA/DNA Index Anchors Set A for Automation	20066404
IDT for Illumina UMI DNA/DNA Index Anchors Set B for Automation	20063213

Ordering information—Sequencing reagents

Product	Catalog no.
NovaSeq 6000 S2 Reagent Kit v1.5 (300 cycles)	20028314
NovaSeq 6000 S4 Reagent Kit v1.5 (300 cycles)	20028312
NovaSeq™ X Series 1.5B Reagent Kit (300 cycles)	20104705
NovaSeq X Series 10B Reagent Kit (300 cycles)	20085594

Ordering information—Analysis

Product	Catalog no.
Local variant reporting	
Illumina DRAGEN Server v4	20051343
Illumina DRAGEN Server Installation	20031995
Illumina DRAGEN Server v4 Support Plan	20085832
Field Delivered Applications Training	15032919
Cloud-based variant reporting	
ICA Basic Annual Subscription	20044874
ICA Professional Annual Subscription	20044876
ICA Enterprise Annual Subscription	20038994
ICA Enterprise Compliance Add-on	20066830
Subscription ICA Training and Onboarding	20049422
Variant interpretation	
Illumina Connected Insights – Annual Subscription	20090137
Illumina Connected Insights—Research – Annual Subscription	20112516
Illumina Connected Insights – Oncology Genome Equivalent Samples (VCF)	20090138
Illumina Connected Insights Training – Remote	20092376
Informatics Professional Services	20071787
Cloud storage	
Illumina Analytics – 1 iCredit	20042038
Illumina Analytics Starter Package – 1,000 iCredits	20042039
Illumina Analytics – 5,000 iCredits	20042040
Illumina Analytics – 50,000 iCredits	20042041
Illumina Analytics – 100,000 iCredits	20042042

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

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