

Illumina DNA Prep with Exome 2.0 Plus Enrichment

Focused, consistent
exome results from a
single partner

- Cost-effective exome coverage using a focused, comprehensive, up-to-date panel
- Easy-to-use library preparation kit with qualified automation methods
- High-quality end-to-end solution and support

illumina[®]

End-to-end exome sequencing with a single reliable partner

Illumina DNA Prep with Exome 2.0 Plus Enrichment delivers economical human whole-exome sequencing (WES) results with outstanding performance and data quality. The easy-to-use library preparation and enrichment solution are part of an end-to-end workflow that spans from samples to reporting (Figure 1). Illumina Qualified Methods are available on a range of automation platforms through our partners. Illumina DNA Prep with Exome 2.0 Plus Enrichment starts with extracted genomic DNA (gDNA)—or direct blood or saliva input*—and combines rapid on-bead tagmentation library preparation chemistry followed by hybrid-capture exome enrichment (Figure 2).¹ The Illumina DNA Prep with Enrichment chemistry supports integrated normalization of high-quality input DNA (≥ 50 ng), which enables simple volume-based pooling for hybridization and provides even sequencing output from each enriched exome library. Libraries are sequenced on the NovaSeq™ 6000, NextSeq™ 1000, NextSeq 2000, or NextSeq 550 Systems. Award-winning DRAGEN™ secondary analysis performs variant calling.² For genetic diseases, the Emedgene platform applies explainable artificial intelligence (XAI) and automation to streamline interpretation and reporting for exome panels.

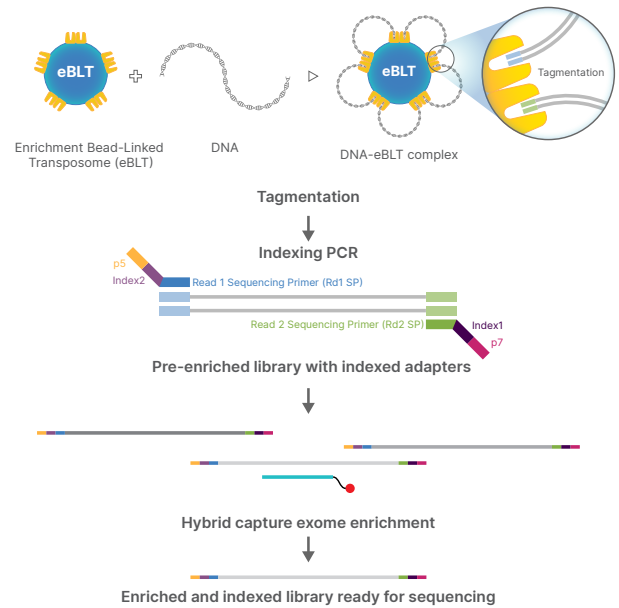


Figure 2: Streamlined tagmentation-based library preparation with exome enrichment—Enrichment bead-linked transposomes (eBLT) mediate a uniform tagmentation reaction with high tolerance to varying DNA sample input amounts.¹ Following hybrid-capture enrichment, exome libraries are ready for sequencing.



Figure 1: From samples to reporting from a single partner—Illumina supports an end-to-end workflow for WES. Extracted gDNA (or DNA from blood or saliva following direct extraction protocols) is input to library prep with Illumina DNA Prep with Exome 2.0 Plus Enrichment. Qualified Methods are available on a range of automation platforms. Sequence according to scale and throughput needs on Illumina instruments. Accurate, rapid secondary analysis and variant calling is performed with the DRAGEN Enrichment App. For genetic diseases, Emedgene enables intuitive interpretation and reporting.

* Direct blood and saliva protocols require the Flex Lysis Reagent Kit. Data shown is generated from extracted gDNA, not blood or saliva. Blood and saliva performance may vary depending on sample quality.

Cost-effective, comprehensive coverage of disease-associated variants in public databases

Illumina DNA Prep with Exome 2.0 Plus Enrichment uses a focused, comprehensive, up-to-date exome enrichment panel. The Twist Bioscience for Illumina Exome 2.0 Plus Panel, included in the kit, is smaller than the Illumina Exome Panel and has improved target region coverage for variants reported in public databases (Table 1 and Table 2). This focused coverage enables a cost-effective WES solution at an optimal number of samples per sequencing run (Table 3).

The Twist Bioscience for Illumina Exome 2.0 Plus Panel covers curated coding sequences (CDS) from RefSeq, Consensus Coding Sequence (CCDS), American College of Medical Genetics and Genomics (ACMG), the Cancer Gene Census from the Catalog of Somatic Mutations in Cancer (COSMIC), and the Online Mendelian Inheritance in Man (OMIM) (Table 1) with exceptional coverage of pathogenic or likely pathogenic variants reported in the ClinVar and ACMG databases (Table 2).³⁻⁹

Table 1: Improved target region coverage in a cost-effective size with the Twist Bioscience for Illumina Exome 2.0 Plus Panel

Exome panel	Twist Bioscience for Illumina Exome 2.0 Plus Panel	Illumina Exome Panel
Size	37.5 Mb	42.21 Mb
RefSeq CDS ³	99.1%	98.2%
CCDS CDS ⁴	99.9%	99.5%
ACMG 73 genes CDS ⁵	99.9%	99.3%
COSMIC Cancer Gene Census CDS ^{6,7}	99.9%	99.3%
OMIM ⁸	99.1%	97.7%

Table 2: Clinical research-focused content of the Twist Bioscience for Illumina Exome 2.0 Plus Panel

ClinVar pathogenic/likely pathogenic variants CDS ^{a,9}	98.6%
ACMG 73 pathogenic/likely pathogenic variants CDS ^{b,5}	99.9%

a. The ClinVar public archive reports relationships among human variations and phenotypes with supporting evidence. Pathogenic/likely pathogenic variants are reported based on ClinVar classification guidelines.

b. ACMG pathogenic/likely pathogenic variants is the overlaid variants between the curated coding sequences of ACMG genes and ClinVar pathogenic/likely pathogenic variants.

Table 3: Estimated number of enriched Exome 2.0 Plus libraries per flow cell and sequencing system^a

Desired mean target coverage depth	NextSeq 550 System ^b		NextSeq 2000 System		NovaSeq 6000 System			
	Mid-output	High-output	P2 ^c	P3	SP	S1	S2	S4
50×	6	19	19	57	34	69	176	428 ^d
100×	3	9	9	28	17	34	88	214
200×	1	4	4	14	8	17	45	109

a. Estimates are based on 2 × 101 bp read length, calculated based on typical internal experiments. Number of samples may vary depending on workflow handling, input sample or library quality, and actual sequencing output of each platform and flow cell. Actual data was acquired on the NovaSeq 6000 System using S4 flow cells and extrapolated to other instruments and flow cells.

b. NextSeq 550 reagent kits support 2 × 150 bp read lengths.

c. P2 flow cells with the same sample throughput are also available on the NextSeq 1000 System.

d. Pooling this many enriched libraries requires additional indexes.

Illumina DNA Prep with Exome 2.0 Plus Enrichment sequencing data shows excellent coverage consistency with a high percentage of targets covered at 20× sequence depth or higher using 5 Gb output and two different hybridization times for enrichment (Figure 3).

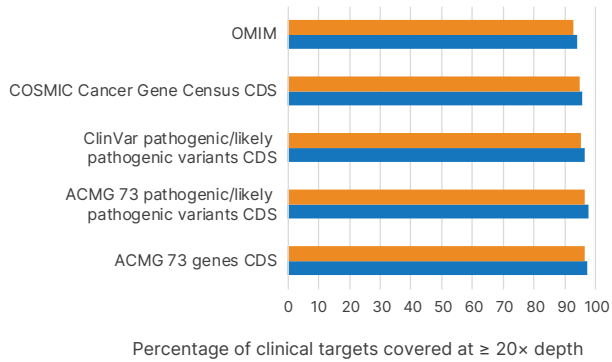


Figure 3: Extensive, comprehensive coverage of disease-associated variants in public databases—Illumina DNA Prep with Exome 2.0 Plus Enrichment using a 1.5-hour (orange) or 16-hour (blue) hybridization time shows high average percent coverage at $\geq 20\times$ of targets from public databases, including ACMG, ClinVar, OMIM, and COSMIC.⁵⁻⁹

High-quality performance

Illumina DNA Prep with Exome 2.0 Plus Enrichment shows outstanding enrichment assay performance. Data was collected using the Illumina DNA Prep with Exome 2.0 Plus Enrichment kit and the included Twist Bioscience for Illumina Exome 2.0 Plus Panel with a 1.5-hour or 16-hour hybridization time. For comparison, the assay was also performed using the original Illumina Exome Panel, following the same protocol except the 1.5-hour hybridization and washing temperatures were reduced to 58°C due to the use of shorter probes (labeled 'IEP 1.5 hr' in Figure 4 and Figure 5).

Performance metrics from the DRAGEN Enrichment App and the Picard pipeline¹⁰ reveal optimal percentage of mappable reads, reads on target (based on percent passing filter for unique reads aligned, percent padded unique read enrichment, and percent padded unique base enrichment), and outstanding mean target coverage for Illumina DNA Prep with Exome 2.0 Plus Enrichment with the Twist Bioscience for Illumina Exome 2.0 Plus probes (Figure 4).

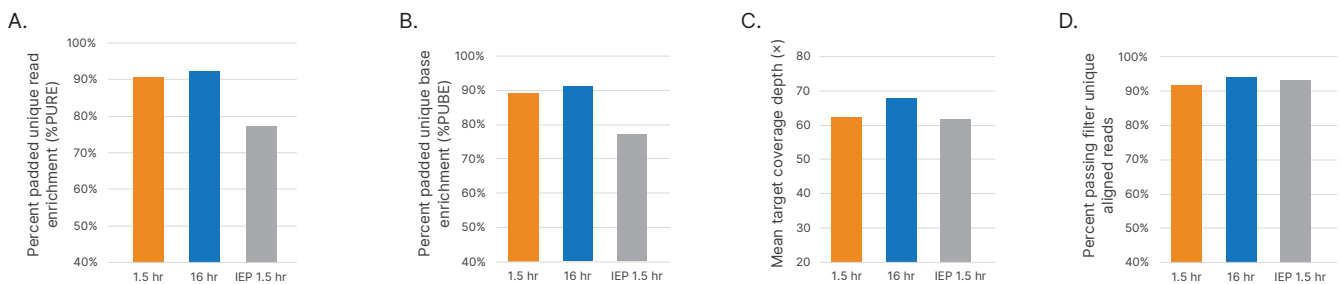


Figure 4: Excellent enrichment assay performance of Illumina DNA Prep with Exome 2.0 Plus Enrichment—Illumina DNA Prep with Exome 2.0 Plus Enrichment using a 1.5-hr (orange) or 16-hr (blue) hybridization time illustrates robust performance compared to the Illumina Exome Panel (IEP 1.5 hr, gray). DRAGEN metrics: (A) percent padded unique read enrichment (% PURE, 150-bp padded size); (B) percent padded unique base enrichment (% PUBE, 150-bp padded size); and (C) mean target coverage depth. Picard metrics: (D) percent passing filter unique reads aligned. Enriched libraries were sequenced on the NovaSeq 6000 System, S4 flow cell with 5 Gb output (50M paired-end reads, 25M clusters) and 2×101 bp read lengths.

Coverage depth and uniformity

Additional analysis demonstrates excellent coverage uniformity of the Illumina DNA Prep with Exome 2.0 Plus Enrichment compared to the Illumina Exome Panel (Figure 5). Illumina DNA Prep with Exome 2.0 Plus Enrichment ensures even coverage with a high percentage of bases at 20× or higher read depth, a low fold-80 base penalty, and a low percentage of zero coverage targets.

The consistent performance between the 1.5-hour and 16-hour hybridization times illustrates how labs can speed up their workflows with 1.5-hour hybridization times or extend their hybridizations overnight if that better suits their workflow.

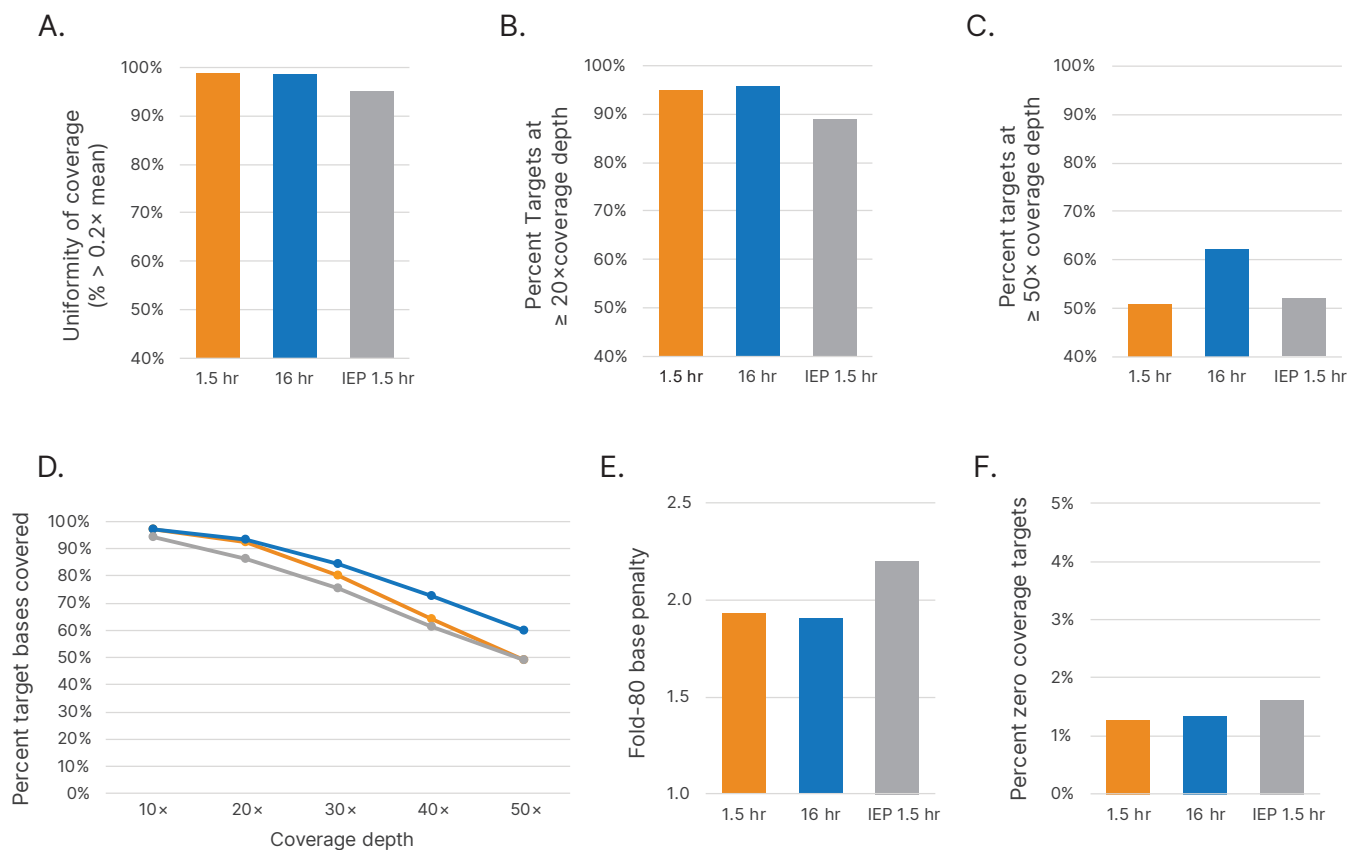


Figure 5: Excellent coverage and uniformity independent of the hybridization time—Illumina DNA Prep with Exome 2.0 Plus Enrichment using a 1.5-hr (orange) or 16-hr (blue) hybridization time shows excellent coverage uniformity compared to the Illumina Exome Panel (IEP 1.5 hr, gray) run in parallel. DRAGEN metrics: (A) coverage uniformity (percent > 0.2x mean coverage); (B) percent targets covered at $\geq 20\times$; and (C) percent targets covered at $\geq 50\times$. Picard metrics: (D) percent target bases with 10x, 20x, 30x, 40x, and 50x coverage depth; (E) fold-80 base penalty (fold over-coverage necessary to raise 80% of bases in nonzero coverage targets to the mean coverage level in those targets); and (F) percent zero coverage targets. Enriched libraries sequenced on the NovaSeq 6000 System, S4 flow cell with 5 Gb output (50M paired-end reads, 25M clusters) and 2×101 bp read lengths.

Summary

Illumina DNA Prep with Exome 2.0 Plus Enrichment offers a well-designed, reliable human WES solution that is effective and efficient. The included Twist Bioscience for Illumina Exome 2.0 Plus Panel provides comprehensive, up-to-date content covering disease-associated variants within the public databases. The optimized enrichment panel enables high sample throughput for economical exome sequencing. Additional efficiency gains can be achieved by adopting Illumina Qualified Methods on a range of automation platforms, available through our partners. Excellent, uniform coverage facilitates downstream analysis and interpretation. Labs can now benefit from a high-quality end-to-end exome sequencing workflow—from samples to reporting—from a single partner.

Learn more

[Illumina DNA Prep with Exome 2.0 Plus Enrichment](#)

[Illumina Qualified Methods for automation](#)

[DRAGEN secondary analysis](#)

[Emedgene tertiary analysis](#)

Ordering information

Product	Catalog no.
Illumina DNA Prep with Exome 2.0 Plus Enrichment, (S) Tagmentation Set B (96 samples, 12-plex) ^a	20077595
Illumina DNA Prep with Exome 2.0 Plus Enrichment, (S) Tagmentation Set D (96 samples, 12-plex) ^a	20077596
Flex Lysis Reagent Kit (96 reactions) ^b	20018706
IDT for Illumina DNA/RNA Unique Dual Indexes Set A, Tagmentation (96 indexes, 96 samples) ^c	20027213
IDT for Illumina Nextera™ DNA/RNA UD Indexes Set C, Tagmentation (96 indexes, 96 samples) ^c	20042666

- Kits include Illumina DNA Prep with Enrichment library preparation and hybridization reagents, Illumina Purification Beads for cleanup/size selection, the Twist Bioscience for Illumina Exome 2.0 Plus Panel enrichment probes, and an index adapter plate.
- Kit required for direct blood input.
- Choose a different index set if preferred.

References

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