

The genomic application that combines a capture-based target enrichment kit with the analytical capabilities and advanced features of the SOPHIA DDM™ platform.

Main Features

SOPHIA Whole Exome Solution v1 covers the coding regions (±5bp of intronic regions) of >19,000 genes and spans 39 Mb of target region (probe footprint: 51 Mb). Probe design is highly optimized to guarantee a high on-target reads percentage and coverage uniformity even in GC-rich regions, including the first exon.

| Gene Panel | Variants Called | Recommendations | Wet Lab |
|--|------------------------------------|---|--|
| <ul style="list-style-type: none"> >19,000 genes | SNVs Indels CNVs (93% genes) | Starting material 200 ng DNA Sample type Blood Samples per run for > 50x coverage depth / Sequencer (Flow Cell) 3 for Illumina NextSeq® 500/550 Mid Output v2 (2x150bp) 12 for Illumina NextSeq® 500/550 High Output v2 (2x150bp) 12 (per lane) for Illumina NovaSeq® 6000 (SP) 24 (per lane) for Illumina NovaSeq® 6000 (S1) 56 (per lane) for Illumina NovaSeq® 6000 (S2) 3 (per lane) Illumina HiSeq® 2500 Rapid Run Mode (2x150bp) 6 (per lane) Illumina HiSeq® 2500 High Output (2x150bp) | Day 1: Library Preparation Day 2: Capture and Sequencing Total library preparation time: 1.5 days |

Analytical Performance

The SOPHIA DDM™ platform analyzes complex NGS data by detecting, annotating and pre-classifying multiple types of genomic variants in all the genes of the panel.

Analysis time¹ from FASTQ: Overnight

| | Observed |
|--|----------|
| Sensitivity ² | >99% |
| Reproducibility | >99% |
| Average on-target rate | >90% |
| Coverage uniformity | >98% |
| Average % of target region with depth >20x | >99% |

One Simple Intuitive Platform: Beyond Analytics

Accelerated assessment and reporting of genomic variants

Dedicated features in SOPHIA DDM™ reduce the complexity of determining the significance of genomic variants and facilitate the interpretation process, thus reducing turnaround time:

- Dual Variant Pre-classification** - Improve assessment of variant pathogenicity with both ACMG scores and SOPHIA GENETICS' machine learning-based predictions
- Virtual Panels** - Restrict the interpretation to sub-panels of genes of interest using the HPO or OMIM® browser
- Cascading Filters** - Apply custom filtering options for quicker screening of relevant variants and save strategies for future analyses
- Familial Variant Analysis (trio-analysis)** - Identify pathogenic variants considering different modes of inheritance, through a family-based approach

After the interpretation, you can generate a customizable variant report including valuable information to support research decision making.

Global support at every step

We offer local support anywhere in the world. Our dedicated bioinformaticians help save time and resources, ensuring fast resolution of workflow disruptions. In addition, our Set Up Program provides assistance with assay set up for a fast and worry-free transition to routine testing.

Secure and unlimited data storage

Access to the SOPHIA DDM™ platform is restricted to registered users only. The platform provides unlimited and unrestricted storage, while keeping data safe by applying the highest industrial standards of encryption in compliance with your local data security policies.

Access to the SOPHIA GENETICS community

In the SOPHIA DDM™ platform, experts from hundreds of healthcare institutions interpret their results and flag the pathogenicity level of variants according to their knowledge and experience. This highly valuable information feeds the variant knowledge base and is anonymously and safely shared among the members of the community.

1. Analysis time may vary depending on the number of samples multiplexed and server load.
 2. Performance metrics are based on high confidence regions in a reference sample, with 80M reads per sample. Sequencing was performed using an Illumina HiSeq® instrument (150bp read length).

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