

Dislipidemias-GeneSGKit[®]

Sistemas Genómicos has developed Dislipidemias-GeneSGKit[®], a molecular design of **34 genes whose mutations are associated with various alterations of the metabolism of lipids of monogenic origin that increase the risk of atherosclerotic coronary disease**. Dislipidemias-GeneSGKit[®] allows the diagnosis of different pathologies such as disorders related to hypercholesterolemia and mixed hyperlipidemia, disorders related to hypertriglyceridemia, hyperalphalipoproteinemias, hypocholesterolemia and hypotriglyceridemia, hypoalphalipoproteinemia, hypobetalipoproteinemia and myopathy secondary to statins. Dislipidemias-GeneSGKit[®] also includes the analysis of the **CFTR gene** related with the **cystic fibrosis**.

The workflow is optimised to prepare the libraries in nine hours and the technology used allows the combination of multiple samples and different GeneSGKits[®] in a single run on Illumina sequencing platforms, ensuring an equitable balance of data from the different sample.

HIGHLIGHTS

The most complete solution on the market

Dislipidemias-GeneSGKit[®] is cleared for CE-IVD and contains reagents to process twelve samples with the corresponding bioinformatics analysis and visualization by the GeneSystems[®] platform.

Fast workflow and low input requirements

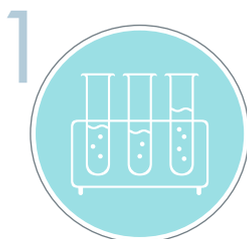
The libraries are ready within nine hours of work starting with 50 ng of DNA.

Complete bioinformatic analysis and data visualization

The bioinformatics pipeline is executed in less than 48 hours, providing the user with a rich variant annotation including public, private and Sistemas Genómicos' reference databases. This comprehensive annotation reports SNPs and indels, which are visualized and easily filtered on GeneSystems[®].

CNVs detection

Sensitive CNVs detection validated in prestigious public and private organisations.



GeneSGKits[®] library preparation



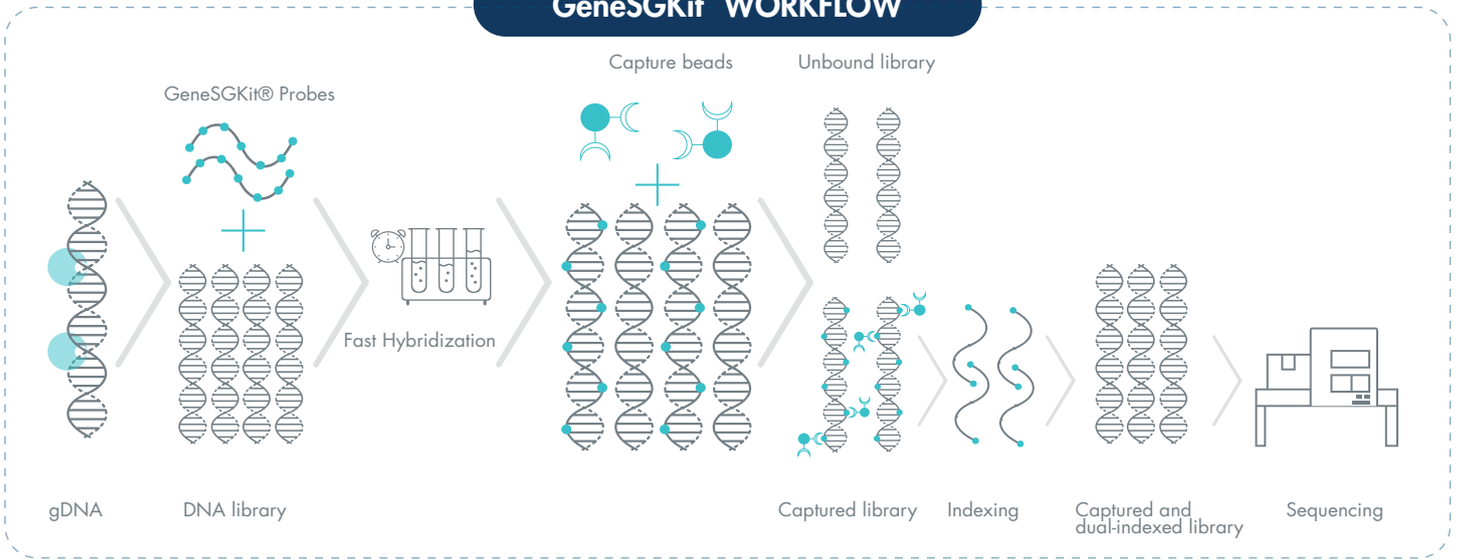
Sample sequencing on Illumina[®]* platforms



Analysis and data interpretation on GeneSystems[®]

*HiSeq and MiSeq. Other sequencing platforms in validation

GeneSGKit® WORKFLOW



ABOUT US

Sistemas Genómicos is a pioneering company in the application of Next Generation Sequencing technology for genetic diagnosis. Accredited and certified by the major entities in quality assurance (ENAC, AENOR and CLIA), we offer solutions to accelerate and standardize clinical genetic testing, enabling clinicians to provide accurate diagnostics to their patients.

OUR MISSION

We strive to provide the best diagnostic solutions with the best technology and knowledge available to maximise patient benefit. Our integrated systems have helped laboratories for the last two decades to improve their diagnostics and research work, by developing a unique range of CE-IVD solutions.



DESIGN

The GeneSGKits® offer excellent performance from only 50 ng of DNA. The capture probes are based on ultra-long RNA baits (120 nucleotides) with high sensitivity and specificity for the detection of SNPs, indels and CNVs. Our probe design captures all the coding regions of each targeted gene, including splicing sequences, with a recommended average coverage of 200x.



PERFORMANCE

With GeneSGKits® clinicians and researchers can be confident in the accuracy and reproducibility of the sequencing data. Each sample is sequenced with a minimum coverage of 20x in over 98% of the regions of interest for relevant pathologies, guaranteed by strict validation protocols.



SUPPORT

For Sistemas Genómicos, customer is at the center of our work. This includes a personalised service to enable a seamless implementation of diagnostic next generation sequencing. We offer personalised training programs to ensure a quick acquisition of the skill set required to successfully and reliably harness the latest advances in sequencing technology to maximize patient benefit.

 GeneSGKit®



GeneSystems®

SISTEMAS GENÓMICOS

GeneSystems® is a leading platform for the analysis of NGS data for diagnosis and genetic research which is registered as a health product with a CE Marking for diagnostic use in vitro for processing information from massive DNA sequencing to detect genetic variants and help to diagnose at the molecular level heterogeneous genetic diseases by means of directed resequencing. Developed in a private, secure and reliable Cloud Computing environment, it includes:



**Uploading and
downloading data**



**Complete access to all the data, from
raw data to annotated variants**



**Visualization and
prioritization of results**

It is developed by a multidisciplinary team of geneticists, bioinformatic and engineers, and validated by more than 200 companies, including hospitals, laboratories, research centers, universities, pharmaceutical companies. The platform has been audited independently and allows the detection of SNP and INDELS.

DATA ANALYSIS

GeneSystems® is designed to solve the different stages of the analytical data cycle of massive sequencing and provides solutions from the FASTQ files to the clinically-relevant data.

GeneSystems® captures annotations at gene, variant and transcript level and incorporates the most clinically relevant databases for the interpretation of variants, such as Ensemble, 1000Genomes, HGMD, dbSNP, RefSeq, Interpro, Prosite, Cosmic, HapMap, OMIM, ClinVar, ExAC, etc. Furthermore, GeneSystems® users have access to the proprietary variant database of Sistemas Genómicos. Within the platform there is also information on conservation scores, pathogenicity in silico predictors, flanking regions, among others. A one stop solution to find clinical meaning in your sequencing data.

GeneSystems® provides a highly sensitive CNVs detection validated in prestigious public and private organisations, which optimizes the MLPA experiments thus reducing costs and diagnostic turnaround times.

High data quality.

With the GeneSGKits® technology, clinicians and researchers will have the best tools for genetic diagnosis, thanks to the excellent quality of the sequencing data of the samples prepared with GeneSGKit®. Each sample will be sequenced with low variance in the coverage values with 98% of the regions sequenced at a minimum depth of 20x.

Dislipidemias-GeneSGKit[®] CATALOG

Technical data

GeneSGKit [®]	Genes	Size (MB)	% COVERAGE	
			12x	20x
Dislipidemias-GeneSGKit [®]	35	0.12	99.45	98.39

References

Dislipidemias-GeneSGKit[®] Ref.: LV3367 (Kit CE-IVD) / LV3366 (Kit RUO)

Genes description

Dislipidemias-GeneSGKit[®]

ABCA1	APOA5	CETP	GPD1	LDLRAP1	MTTP	PCSK9
ABCG5	APOB	CH25H	GPIHBP1	LIPA	NPC1	SAR1B
ABCG8	APOC2	CYP7A1	INSIG2	LIPC	NPC1L1	SLCO1B1
ANGPTL3	APOC3	GBA	LCAT	LMF1	NPC2	STAP1
APOA1	APOE	GCKR	LDLR	LPL	OSBPL5	CFTR

