



PRODUCTS & SERVICES 2022

# CELEMICS

Innovative NGS-Based Products  
with Novel Sequencing Technology

# Celemics Takes Customization to the Next Level

NGS industries are rapidly evolving and our customers are in greater need of technical support in various research fields and markets.

Just like a chameleon utilizing its colored pigments and crystals to adapt to its surroundings, we proactively pioneer and adapt to the everchanging NGS markets with our cutting-edge technology to support our customer's needs.

We are keenly focused on bringing the right solution to customers by elaborately customizing to individual customer's needs and goals.



Celemics confidently offers customization services  
and preset panels of the utmost quality

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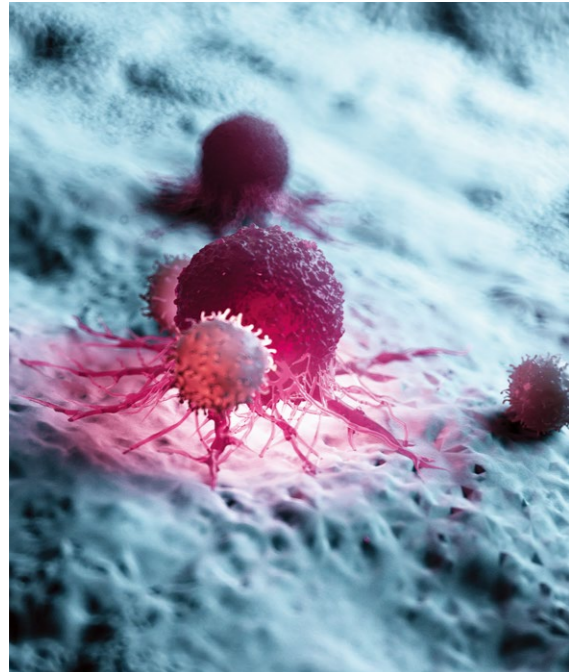
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# CELEMICS PRODUCTS & SERVICES OVERVIEW



## CANCER RESEARCH

- Customized Targeted Sequencing
- BRCA 1/2 (Breast Cancer)
- OncoRisk
- CancerScreen / CancerMaster
- G-Mendeliome Clinical Exome Sequencing
- G-Mendeliome Disease Specific
- Customized RNA Sequencing
- Customized Methylation Sequencing
- Circulating Tumor DNA - Colorectal, Lung, Breast

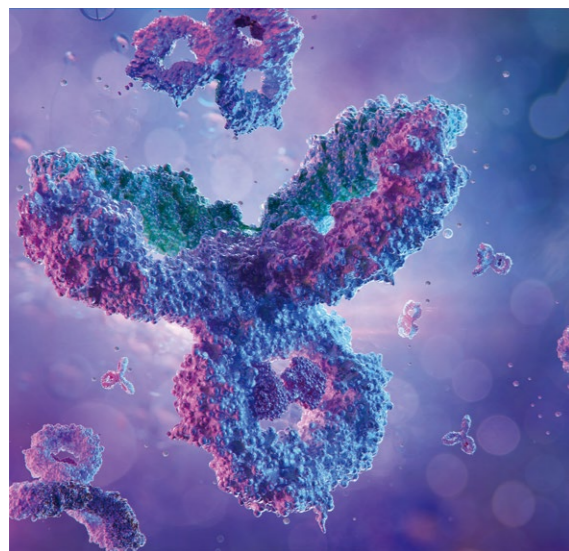
## MOLECULAR BREEDING (ANIMAL / PLANT RESEARCH)

- Customized High-Throughput Genotyping
- Africa Swine Fever Virus



## ANTIBODY DISCOVERY

- TrueRepertoire™
- Immune Repertoire Profiling

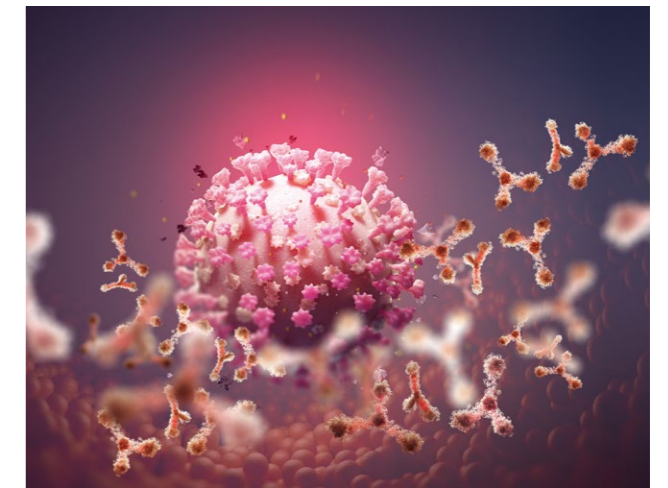


## DIAGNOSTICS & INHERITED DISEASES

- Customized Targeted Sequencing
- G-Mendeliome Clinical Exome Sequencing
- G-Mendeliome Disease Specific
- Cancer Panels (Somatic, Germline, ctDNA)
- Customized RNA Sequencing
- Customized Methylation Sequencing
- PharmaScreen Panel  
: Standard / Epilepsy / Anti-tuberculosis

## MICROBIOLOGY & VIRUS RESEARCHES

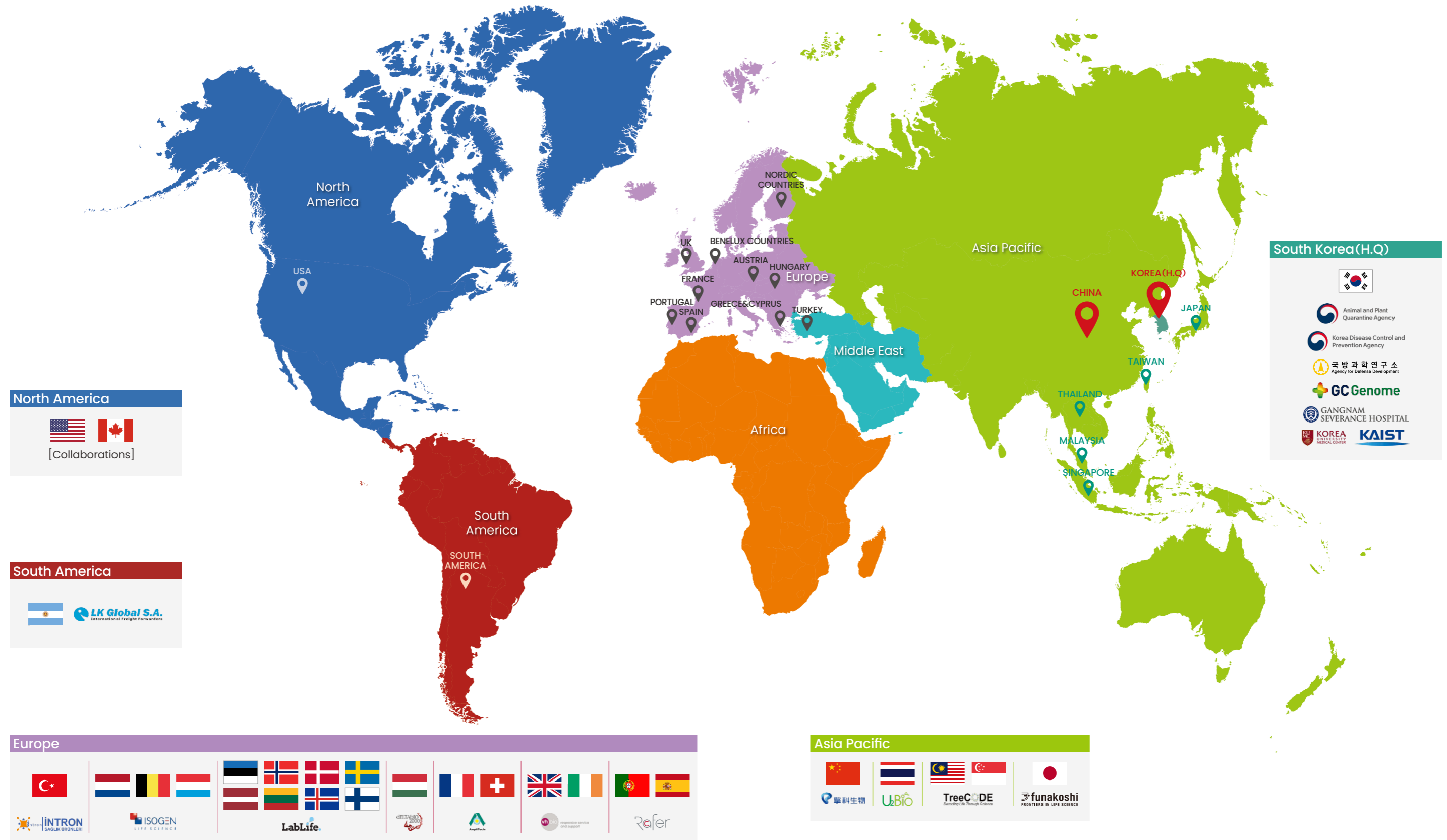
- Customized Targeted Sequencing
- Comprehensive Respiratory Virus
- African Swine Fever Virus
- Customized 16S V4 NGS
- BTSeq™ – Standard
- BTSeq™ – Viral Analysis



## SYNTHETIC BIOLOGY

- BTSeq™ Full Plasmid Sequencing
- BTSeq™ – Standard

# DISTRIBUTORS & COLLABORATIONS



# TARGETED SEQUENCING SOLUTION

CELEMICS PRODUCTS & SERVICES 2022

Targeted Sequencing Overview  
Outstanding Performance of Targeted Sequencing  
Probe Design Technology  
Targeted Sequencing Panel Performance  
Pilot Test & Rebalancing  
Celemics Features & Benefits

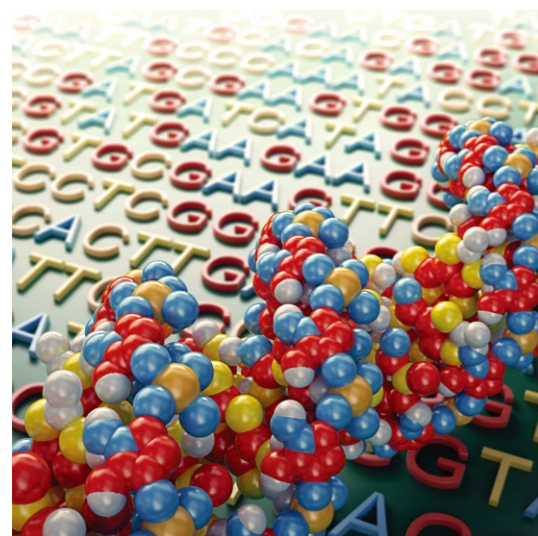


# Targeted Sequencing Overview

Celemics has developed and delivered over 1,000 different customized panels. Our target enrichment method is capable of specifically isolating your genomic loci of interest out of the whole genome and increasing the sensitivity of detecting genetic mutations by producing higher coverage & in-depth sequencing data.



## END-TO-END CUSTOMIZATION



### PANEL DESIGN

- Elaborately designed NGS panels comprised of your genes of interest
- Interactive discussion with customer prior to designing the panel (e.g., GC-rich, Homologous regions)
- Supported by advanced technology for probe design and reagent optimization
- Panel expansion possible through simple gene addition
- Alternative protocols in case required instruments are not available

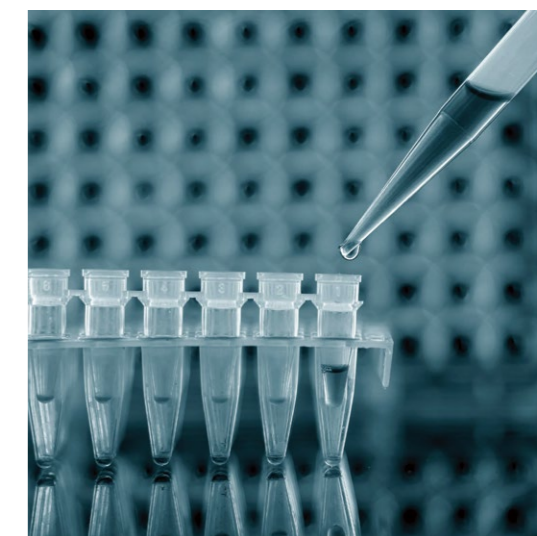
### SUPERIOR PERFORMANCE

- Market leading target enrichment kits
- Maximized cost-effectiveness
- Pre-capture pooling and high panel performance enables additional cost and labor savings



### IN-HOUSE TEST & REBALANCING

- Adjustments to performance and functionality through thorough in-house validation test for every designed panel
- Detailed QC results encompassing wet-lab experiments, NGS run, and bioinformatics analysis provided to customer
- Rebalancing service possible through request
- Able to increase depth and coverage of a specific area if requested
- Finalize your order after reviewing QC results



### DATA ANALYSIS

- Technical support available for customers new to NGS analysis
- Provides bioinformatics analysis services and tools from FASTQ to clinical report by request

## OUTSTANDING PERFORMANCE OF TARGETED SEQUENCING

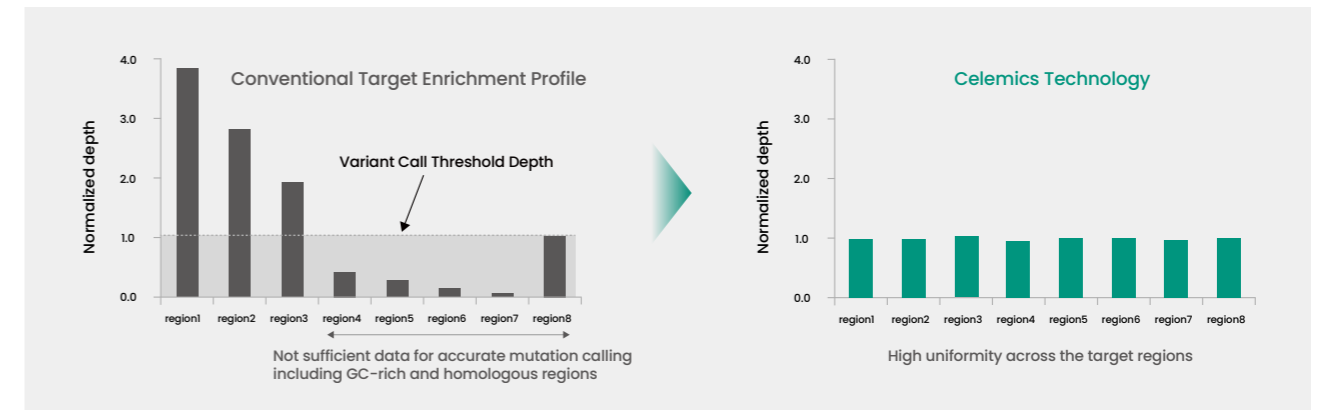
At Celemics, we support our customers through target hybridization-based NGS services and products individually designed and manufactured by experienced researchers and technicians. We have established a robust system for customized design panels and developed a variety of kits according to our customer's needs. All Ready-to-Use kits are completely validated and provide the best performance in the market. Our research team has designed and manufactured over a thousand customized panels and promises to offer the best quality product and service to our customers.

### Key Features

<p><b>1. Exceptional panel performance achieved by hybridization-based target capture method</b></p>	<p>Overcome limitations of amplicon-based NGS analysis with thoroughly validated hybridization-based target capture method</p> <p>High uniformity and coverage achieved by Celemics proprietary probe design technology</p>
<p><b>2. Assess all types of mutations with high sensitivity and specificity</b></p>	<p>Superior analytical performance compared to competitor products in detecting SNV, InDel, CNV, and rearrangement in a single NGS run with maximized sensitivity and specificity and minimized NGS noise enabled by Celemics unique molecular barcode assay and robust bioinformatics pipeline</p>
<p><b>3. Robust performance of assessing DNA and RNA across various specimen quality</b></p>	<p>Compatible with poor-quality and low-amount specimens such as FFPE, solid tumor, liquid biopsy, etc.</p>
<p><b>4. Efficient capture of 'Hard-to-Capture' regions</b></p>	<p>Analyze the clinically significant mutations embedded in GC rich or homologous regions, which are frequently masked by competitors</p>
<p><b>5. Wide compatibility with NGS instruments and automation platforms</b></p>	<p>Compatible with all NGS Instruments from Illumina, Thermo Fisher Scientific, Pacific Bioscience, MGI, and Oxford Nanopore</p> <p>Provides enzymes for DNA fragmentation as a substitute for sonicators</p>
<p><b>6. Flexible panel content: number of reactions of your choice and Gene Add-on Service</b></p>	<p>Save costs by ordering the number of reactions required for your experiment</p> <p>Expand your panel with minimum cost, time, and effort by simply adding or combining panels and genes of your interest</p>

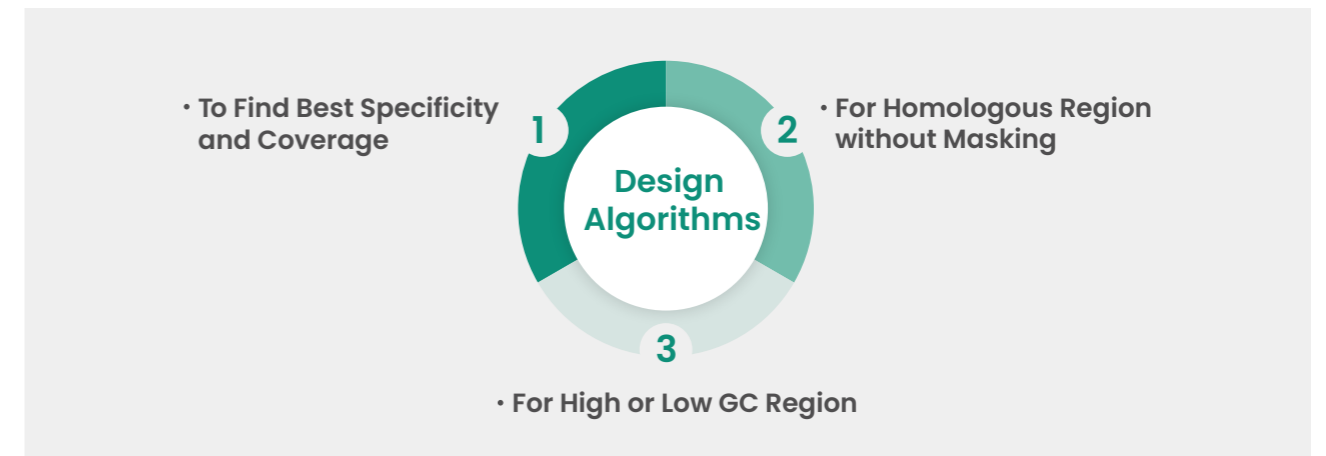
## PROBE DESIGN TECHNOLOGY

### Market Problem and Celemics' Answer



### Proprietary Probe Design Algorithm

Based on extensive wet-lab target capture experimentation for every customized panel



**Customer Testimonial**

“ With Celemics panels, we have obtained successful results with exceptionally high quality in SNV, InDel, and CNV detection.”

-CTO, GC Genome

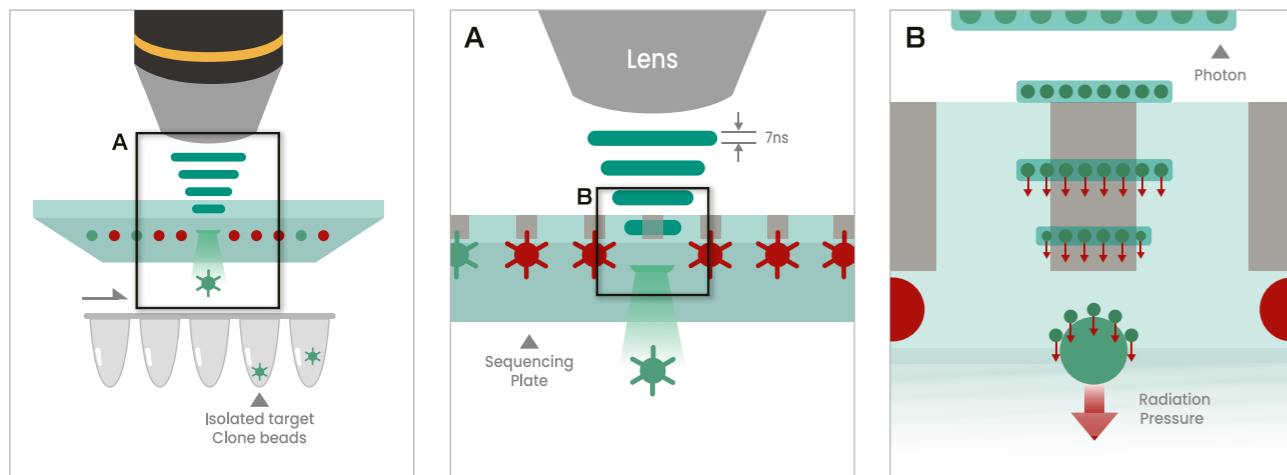
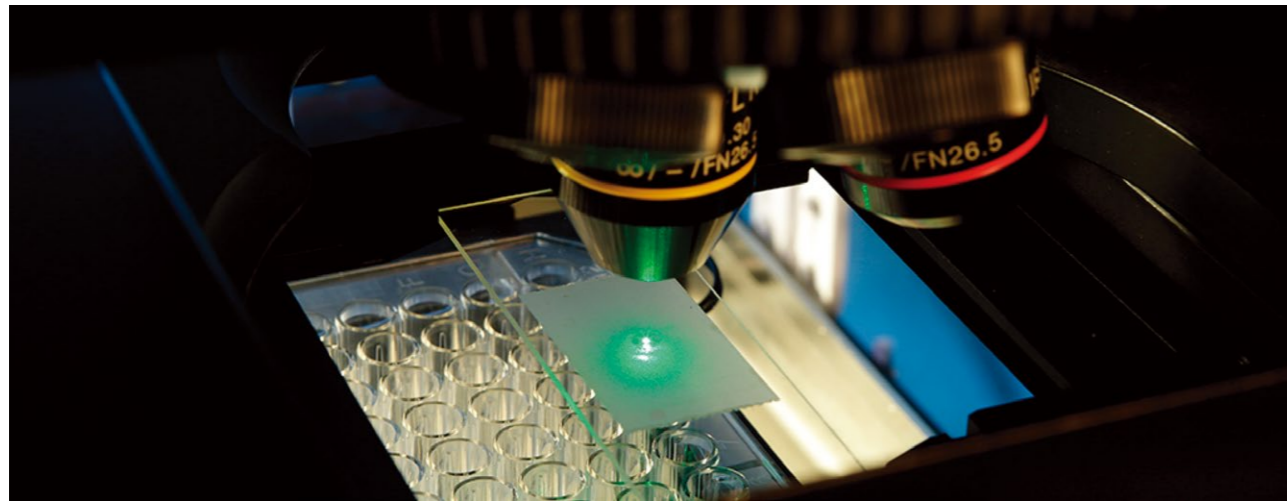


## PROBE DESIGN TECHNOLOGY

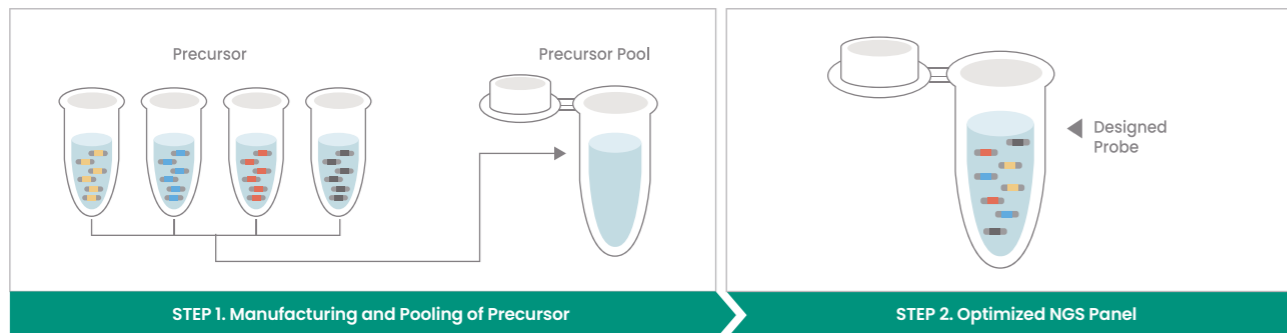
### Proprietary Probe Manufacturing Technology

- Reduces complexity in handling complex oligo pools
- Enables extremely low-biased probe pool with handling individual probe sets
- Allows for cost-effectiveness and high-performance advantage from pool-based probes and individually synthesized probes
- Achieves superior lot-to-lot uniformity for repeated orders due to proprietary 2-step probe synthesis technology

### MSSIC Technology: Massively Separated and Sequence Identified Cloning

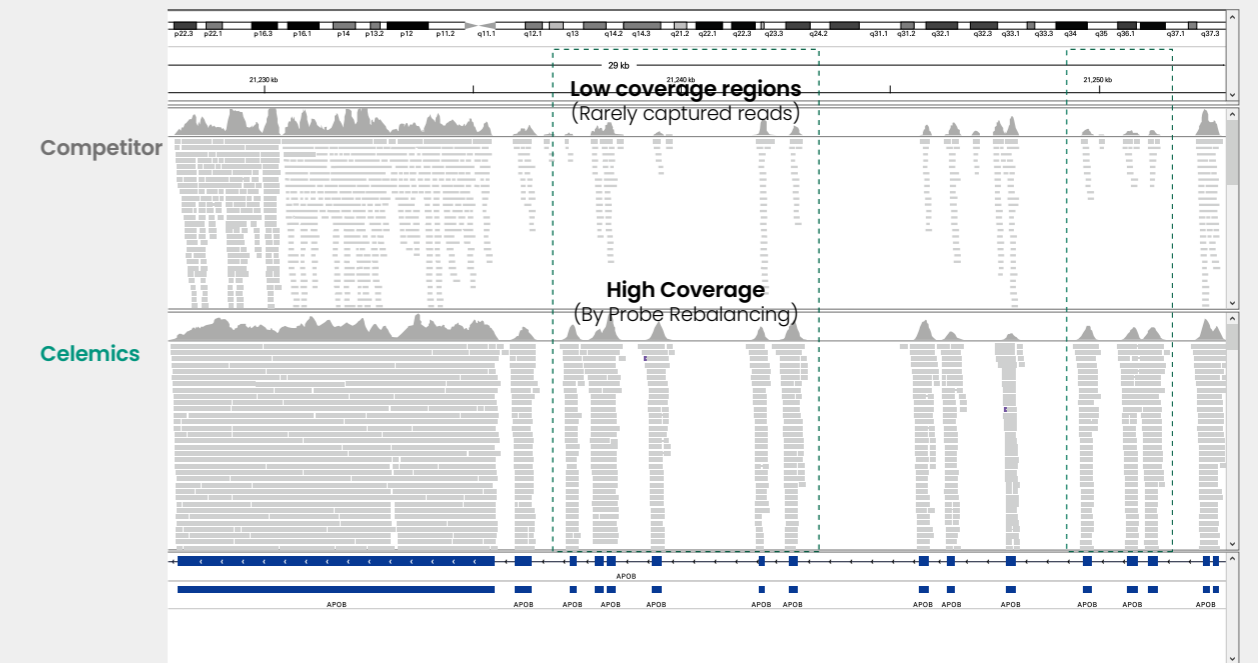


### Two step probe manufacturing

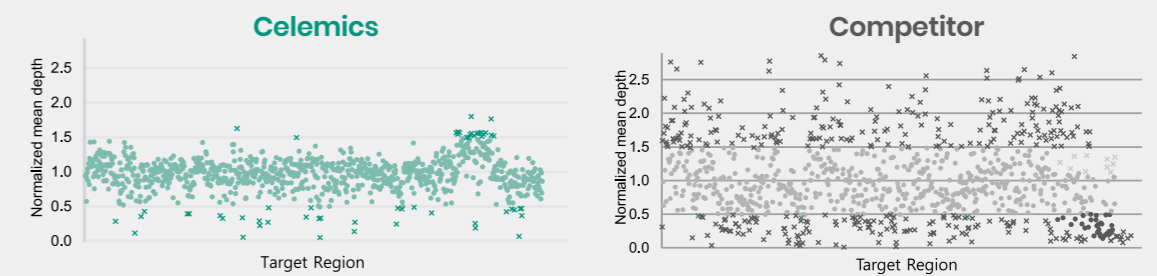


## TARGETED SEQUENCING PANEL PERFORMANCE

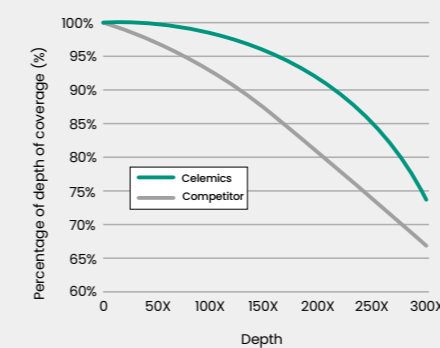
### 1. High Coverage Panel Compared to Competitor Products



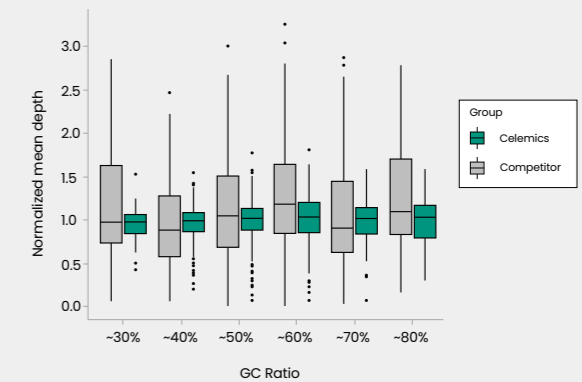
### 2. Higher Uniformity Across Target Regions



### 3. Superior Coverage Depth Over Target Regions



### 4. Superior Capture Performance Across GC Percentage



\*\* Target region of both panels (BED file) are identical.  
 \*\* Number of reads are the same for the results from both panels.

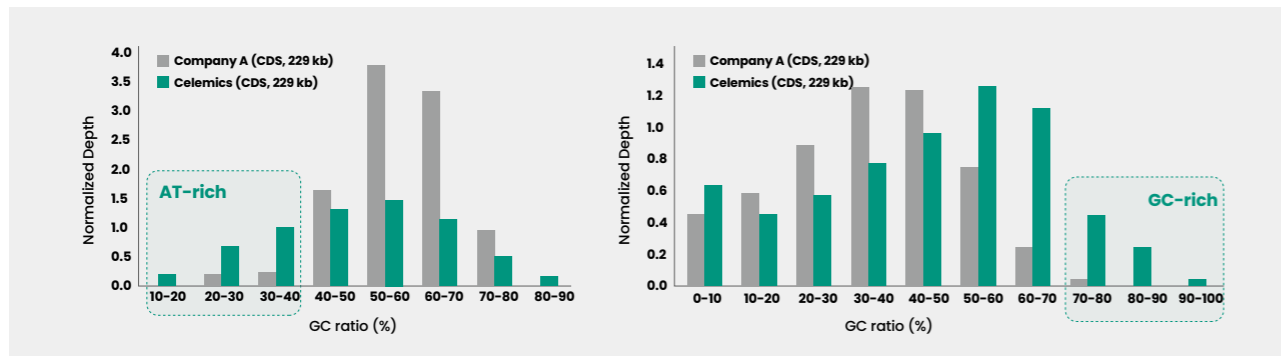
Targeted sequencing allows for sequencing with higher accuracy by specifically targeting the genomic regions of interest. The optimization process of the probes and reagents is essential for each of the different NGS platform types. Celemics has established the design technologies for the probes and reagents for various applications and achieved superior uniformity and depth of coverage compared to competitor products.

# SEQUENCING PERFORMANCE OF CELEMICS PANEL FOR HARD-TO-CAPTURE REGIONS

## 1. Higher Depth compared to Company A Targeting Against the Same Target Area



## 2. Better Uniformity across AT- and GC-rich Regions



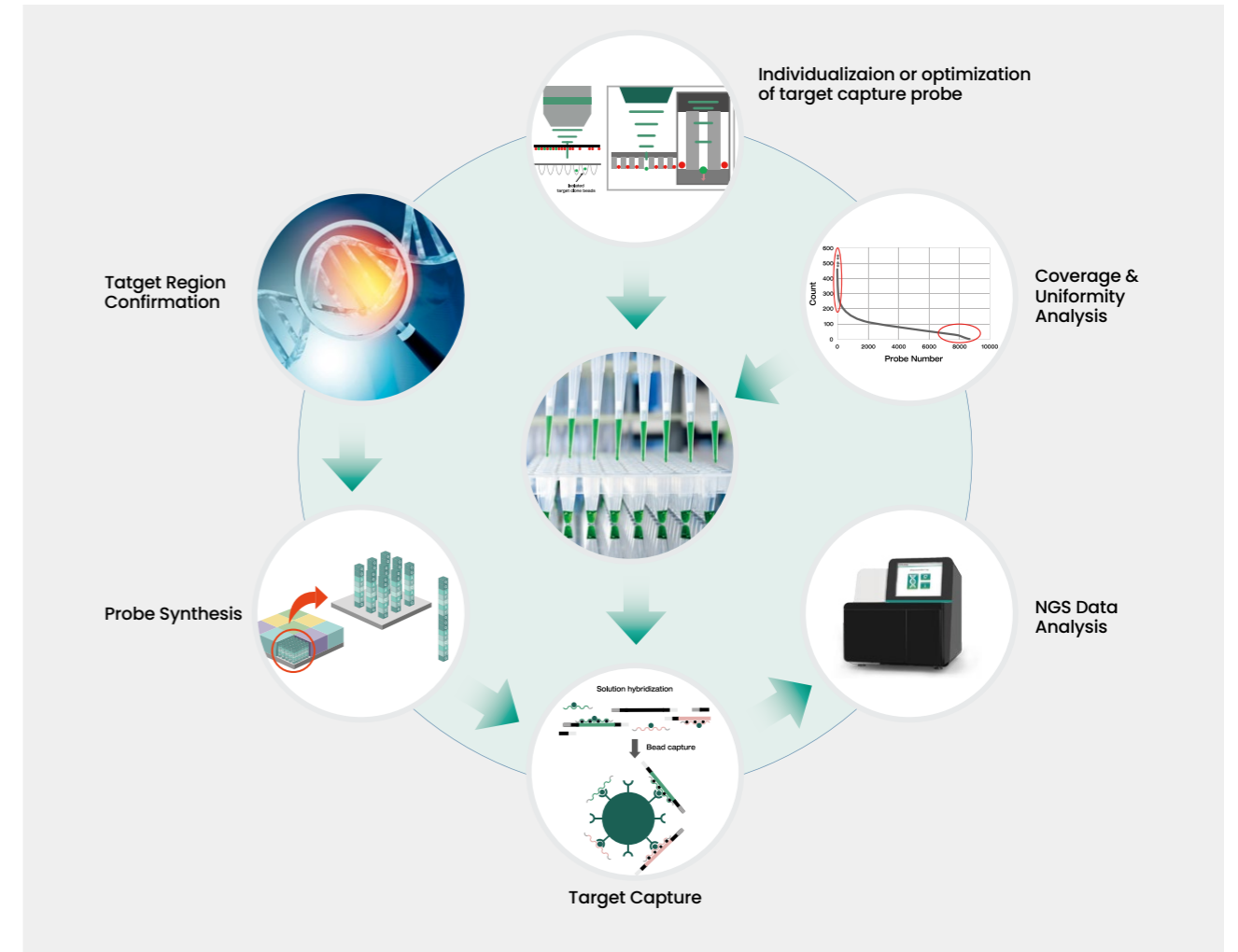
Even the most advanced NGS techniques have been challenged by GC-rich and homologous regions that are often masked or omitted by competitor services. Such a challenge is overcome by Celemics proprietary probe design technology which enables successful sequencing of GC-rich, AT-rich or homologous regions upon request. We also provide Homolog Report when the requested region includes homologous regions. Customers can then decide whether to include the regions in the order.



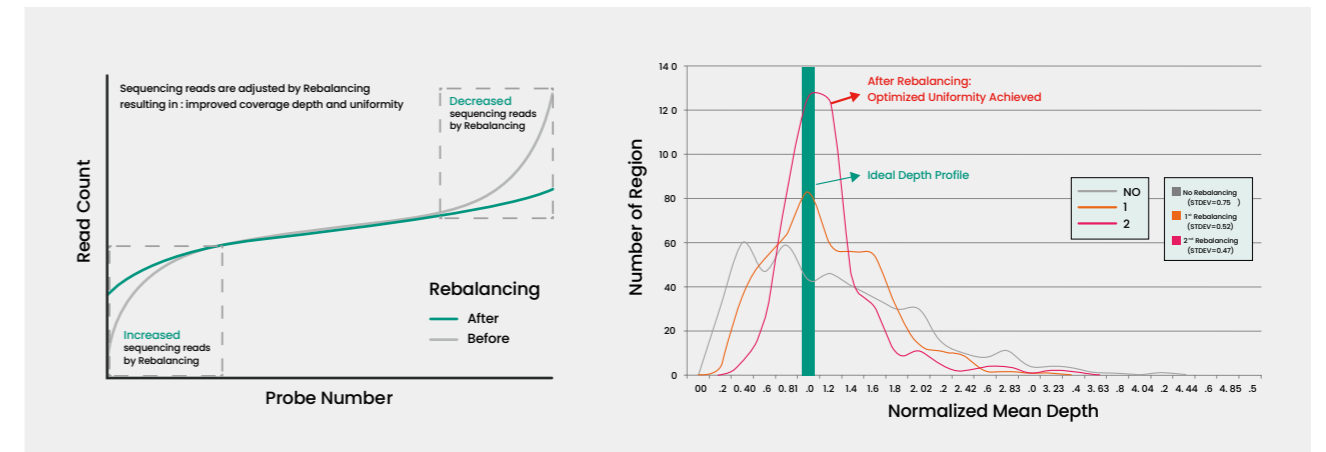
Our customers performed validation tests comparing Celemics' customized panels with our competitors'. For the competitor product, they performed validation tests based on competitor's recommended protocols for the same target regions. They also used the same sequencing amount for the fair experiment. As a result, customers selected our customized panels due to the high capture efficiency even with a lower amount of sequence data.

# PILOT TEST & REBALANCING

## Overview of Celemics Rebalancing Technology



## Capture Uniformity Analysis



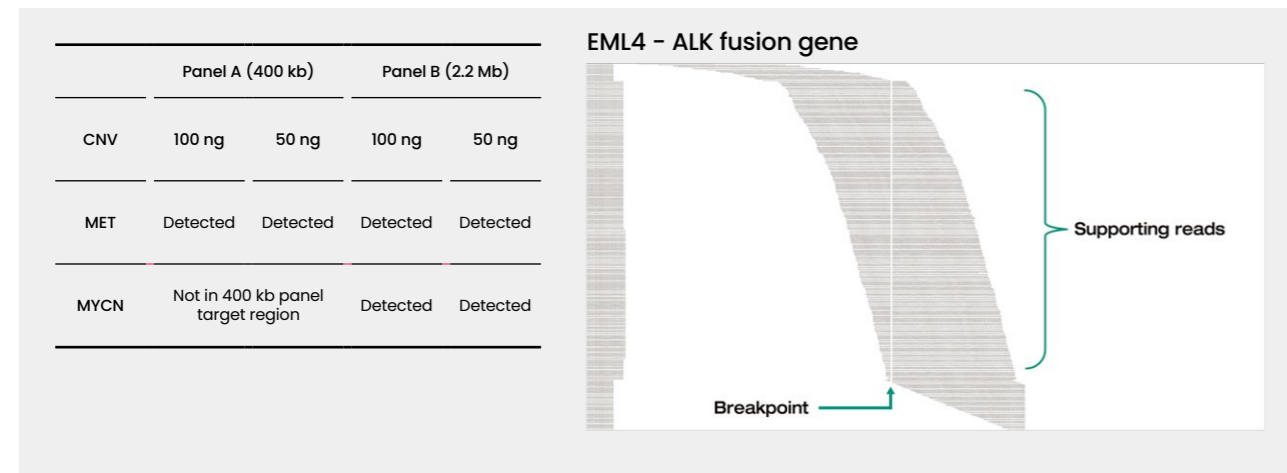
For customized targeted sequencing panels, we conduct in-house performance tests of requested panels and deliver the test results to customers. We also provide rebalancing services in case the customer requests for a specific area or overall performance improvement. The service includes redesigning probes against the requested regions and optimizing reagents to best meet our customers' needs.

## EXAMPLE OF ctDNA ANALYSIS USING PROPRIETARY MOLECULAR BARCODES

Performance Verification using Reference Material:  
100% Sensitivity and Specificity

	Gene	DNA change	AA change	0.5% VAF			1% VAF			WT			
				VAF	VAF	VAF	VAF	VAF	VAF	VAF	VAF	VAF	
Seracare	NRAS	c.182A>G	p.Q61R	0.96%	0.55%	0.78%	1.09%	0.98%	1.44%	0.06%	0.00%	0.00%	
	PIK3CA	c.1633G>A	p.E545K	0.57%	0.69%	0.24%	1.18%	1.13%	0.38%	0.00%	0.00%	0.00%	
	PIK3CA	c.3140A>G	p.H1047R	0.42%	0.33%	0.45%	0.81%	0.93%	0.94%	0.00%	0.00%	0.00%	
	PIK3CA	c.3204_3205insA	p.N1068fs*4	0.51%	0.45%	0.51%	0.86%	0.95%	0.87%	0.00%	0.00%	0.00%	
	EGFR	c.2310_2311insGGT	p.D770_N771insG	0.38%	0.36%	0.42%	0.48%	0.86%	0.78%	0.00%	0.00%	0.00%	
	EGFR	c.2369C>T	p.T790M	0.44%	0.48%	0.48%	0.77%	1.23%	1.05%	0.00%	0.00%	0.00%	
	EGFR	c.2573T>G	p.L858R	0.56%	0.51%	0.74%	1.58%	1.39%	0.85%	0.00%	0.00%	0.00%	
	BRAF	c.1799T>A	p.V600E	0.51%	0.52%	0.47%	0.78%	0.70%	0.45%	0.00%	0.00%	0.00%	
	PTEN	c.741_742insA	p.P248fs*5	0.31%	0.55%	0.51%	1.16%	1.30%	1.52%	0.00%	0.00%	0.00%	
	KRAS	c.35G>A	p.G12D	0.43%	0.34%	0.62%	1.16%	0.89%	0.91%	0.00%	0.00%	0.00%	
	ATK1	c.49G>A	p.E17K	0.69%	0.37%	0.35%	0.65%	0.66%	1.01%	0.00%	0.00%	0.00%	
	TP53	c.818G>A	p.R273H	0.40%	0.47%	0.41%	1.84%	1.14%	0.86%	0.03%	0.05%	0.00%	
	TP53	c.743G>A	p.R248Q	0.47%	0.44%	0.50%	0.90%	0.88%	0.85%	0.02%	0.07%	0.00%	
	TP53	c.723delC	p.C242fs*5	0.43%	0.40%	0.41%	0.87%	0.85%	0.72%	0.00%	0.00%	0.00%	
	TP53	c.524G>A	p.R175H	0.71%	0.66%	0.71%	1.19%	1.13%	1.02%	0.06%	0.05%	0.03%	
	TP53	c.263delC	p.S90fs*33	0.50%	0.81%	0.53%	1.31%	1.55%	1.37%	0.09%	0.01%	0.06%	
	Avg. (%)				0.52%	0.50%	0.51%	1.04%	1.04%	0.94%	0.02%	0.01%	0.01%

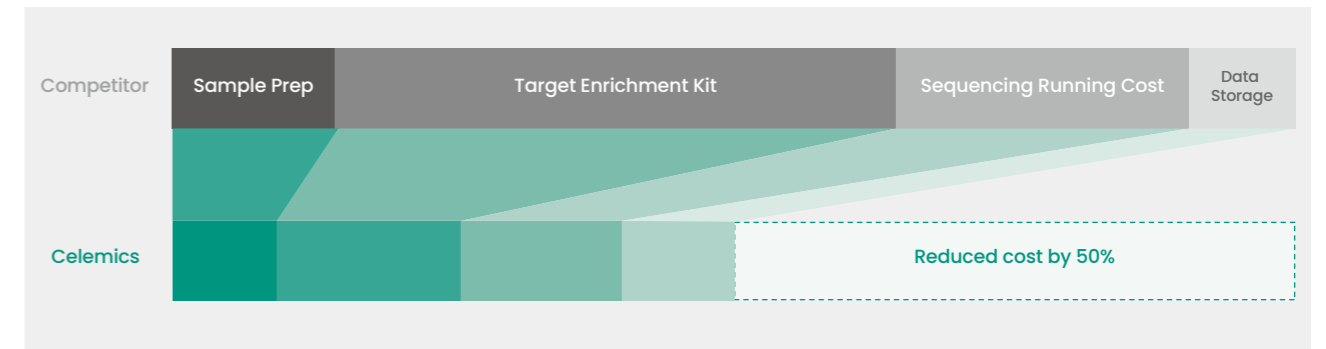
## Accurate CNV and Gene Rearrangement Analysis with FFPE Samples Due to High Coverage Uniformity



We have conducted complete validation test for each Ready-to-Use panel and proved its superior performance compared to competitor products. The products are highly optimized for accurate and efficient assays even with poor quality and low-amount samples such as FFPE, ctDNA, etc. As shown in the table above, we have successfully performed CNV and rearrangement analysis from 50 ng of FFPE samples.

## COST-EFFECTIVE SEQUENCING

Significantly reduced cost in Sample Prep, Target Enrichment Kit, and Sequencing



1. Sample Prep consumables developed and provided by Celeomics for the highest optimization include CeleMag™ Clean-up Bead, CeleMag™ Streptavidin Bead, CLM Polymerase, and EP-kit (one-step workflow from Fragmentation to End-repair and A-tailing).
2. Pre-capture pooling reduces costs per sample.
3. Celeomics has secured technology for proprietary probe design and manufacturing, significantly reducing costs of our Target Enrichment Kit.
4. Celeomics panels have shown superior performance compared to competitor product in terms of uniformity and on-target ratio, enabling high-quality, cost-effective sequencing.

## CELEMICS FEATURES & BENEFITS

1. Hybridization-based capture	2. Maximized Efficiency allows Market Leading Capture Performance	3. Hybridization Enhancer Technology and Enzymatic Library Preparation
4. User-friendly Bioinformatics Software	5. Reduced NGS costs by Pre-capture pooling with no compromise on quality	6. Molecular barcode and bioinformatics for ultra-low VAF mutations
7. CAS for bioinformatics analysis	8. Flexible panel content with Gene Add-on Service	9. Default wet-lab QC for every customized panel
10. Minimal lot variation due to proprietary 2-step probe manufacturing technology	11. Compatible with all NGS instruments and automation platforms	12. Capture the 'Hard-to-Capture' regions
13. Optimization of species-specific blockers for maximum performance for agriculture and animal research	14. Improved Probe Design by Rebalancing Service only available in Celeomics	15. Robust, Rapid, Reliable Customization

# READY-TO-USE PANELS FOR ONCOLOGY

CELEMICS PRODUCTS & SERVICES 2022

BRCA 1/2 Panel  
OncoRisk Panel  
CancerScreen Panel - Core / 50 / 100 / 400  
CancerMaster Panel



# BRCA 1/2 Panel

Germline and Somatic Cancer

## KEY FEATURES

- |  |   |
|--|---|
| 1. Targets the whole CDS (+/- 40) and promoter regions of BRCA 1/2 with high specificity | Target regions not only covering the CDS regions but expanded to +40 and -40 of CDS to detect splicing site variants<br>Probes specifically designed for detecting deletion, duplication, and large rearrangement |
| 2. Compatible with a variety of sample types   | No compromise on panel performance even with using DNA from challenging specimen types such as blood and FFPE   |
| 3. Market-leading panel performance in uniformity and coverage                           | Designed to target whole exon regions of BRCA 1, 2 gene with 100% coverage (RefSeq) and validated to yield 100% coverage  |

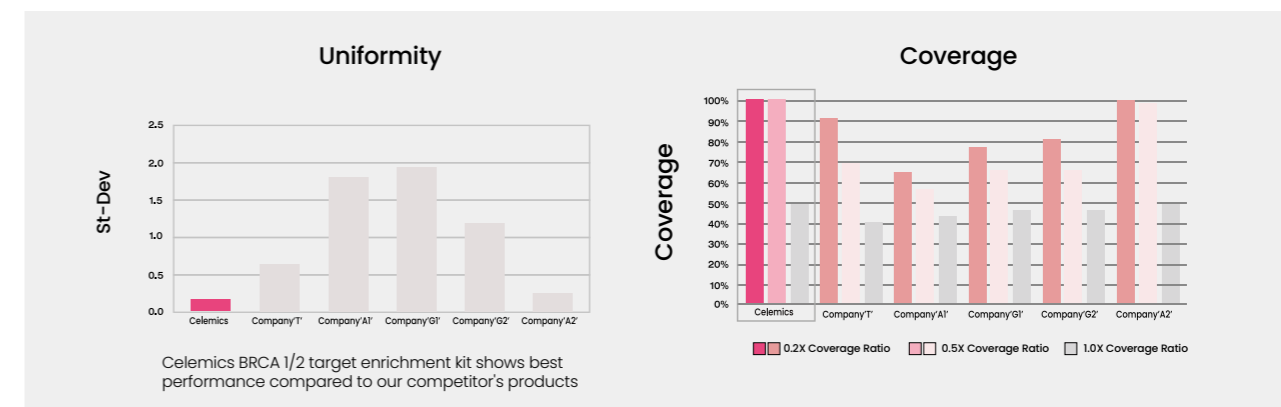
## SPECIFICATION

Gene count*	BRCA 1/2 genes
Covered region	Whole CDS (+/- 40bp), UTR, Promoter
Target size	23 kb
Mutation type	SNV, Indel, CNV
Sample type(amount)	Blood (> 50 ng of fragmented DNA), FFPE
Platform	All sequencers from Illumina, Thermo Fisher, MGI, PacBio, and Oxford Nanopore
Sensitivity	> 95% for all variant types at 5% VAF
Specificity	99.9% (SNV), 99.5% (Indel)
Bioinformatics pipeline	Primary, Secondary and Tertiary analysis result (FASTQ to VCF, VCF to Clinical report)

\* Gene Add-On Service: Genes can be added by customer's request

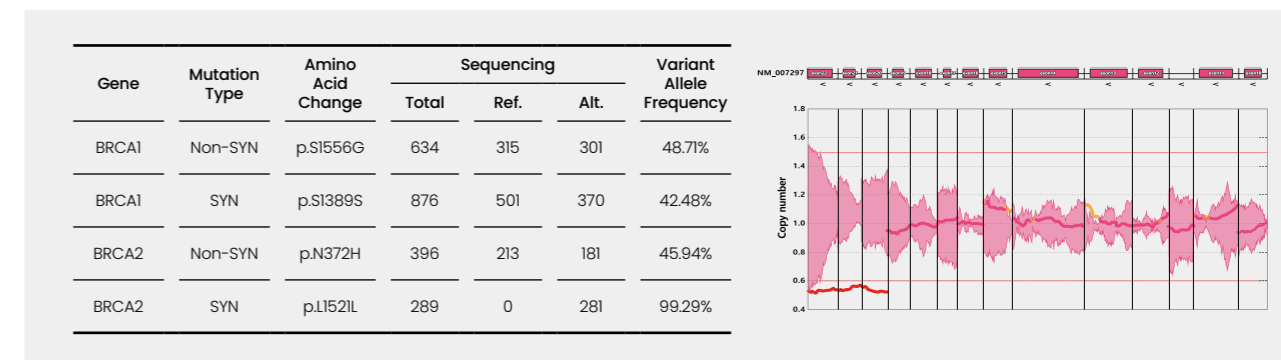
## PANEL PERFORMANCE

### 1. Superior Panel Performance Compared to Competitor Product



### 2. SNV, CNV Analysis

BRCA1, S1556G & S1389S / BRCA2, N372H & L1521L / BRCA1 CNV plot



## PACKAGE COMPOSITION

Package name	Compositions		Package option	Options	
Target Enrichment	Target capture Probe	-	Pooling method	Single Reaction	Pre-capture Pooling
Standard	Target Enrichment reagents	Library prep Kit	Library Preparation kits	Standard Kit	EP-kit
All-In-One	-	Beads / Polymerase	Hybridization Enhancer	Included	Not included



# OncoRisk Panel

Hereditary Cancer  
(Germline Cancer Risk)

## KEY FEATURES

- |   |   |
|---|---|
| 1. Comprehensive analysis of oncogenes                        | Analyze 31 oncogenes associated with inherited cancer and precisely selected from contract research organizations and numerous research studies |
| 2. Robust bioinformatics system for large deletion analysis   | Receive bioinformatics results for large deletion analysis provided by Celeomics proprietary bioinformatics analysis system                     |
| 3. Used for Homologous Recombination Deficiency (HRD) testing | Provides information for HDR grade computation to aid precision medicine for tumor treatment  |

## SPECIFICATION

Gene count*	31 genes
Covered region	Whole CDS
Target size	96 kb
Mutation type	SNV, Indel, CNV, Rearrangement
Sample type (amount)	Blood (> 50 ng of fragmented DNA), FFPE
Platform	All sequencers from Illumina, Thermo Fisher, MGI, PacBio, and Oxford Nanopore
Sensitivity	> 95% for all variant types at 5% VAF
Specificity	99.90% (SNV), 99.50% (Indel)
Bioinformatics pipeline	Primary, Secondary and Tertiary analysis result (FASTQ to VCF, VCF to Clinical report)

\* Gene Add-On Service: Genes can be added by customer's request

## GENE LIST

OncoRisk Panel	APC	ATM	BARD1	BLM	BMPRIA	BRCA1	BRCA2	BRIP1	CDHI	CDK4	CDKN2A	CHEK2	EPCAM
	MLH1	MRE11A	MSH2	MSH6	MUTYH	NBN	PALB2	PMS2	PRSS1	PTEN	RAD50	RAD51C	RAD51D
	SLX4	SMAD4	STK11	TP53	VHL								

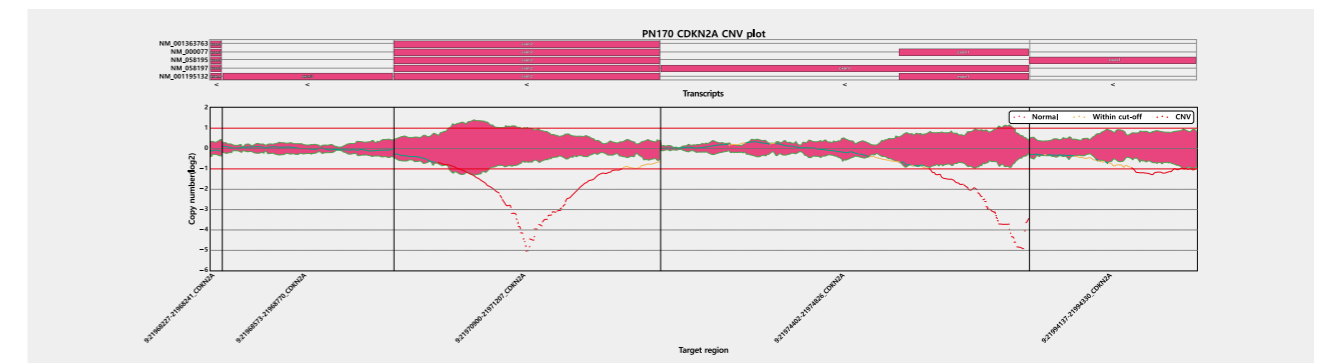
## PANEL PERFORMANCE

### 1. SNV Analysis Example

Gene	Mutation Type	Amino Acid Change	Total Depth	REF Depth	ALT Depth	Variant Allele Frequency
APC	SYN	p.S1738S	1008	590	415	41.17%
ATM	Non-SYN	p.D1853N	417	200	217	52.04%
BARD1	Non-SYN	p.R658C	829	435	394	47.53%
BMPRIA	Non-SYN	p.P2T	621	309	311	50.08%
BRCA1	SYN	p.S1389S	802	460	342	42.64%
BRCA2	SYN	p.V217IV	1026	0	1026	100%
BRIP1	SYN	p.Y1137Y	844	3	840	99.53%
PMS2	Non-SYN	K54I	686	0	646	100%
PRSS1	SYN	p.N246	921	0	921	100%
RAD51D	Non-SYN	p.L152L	971	0	971	100%

### 2. CNV Analysis Example

Higher sequencing depths in the target regions, enabling accurate CNV analysis



## PACKAGE COMPOSITION

Package name	Compositions		Package option	Options	
Target Enrichment	Target capture Probe	-	Pooling method	Single Reaction	Pre-capture Pooling
Standard	Target Enrichment reagents	Library prep Kit	Library Preparation kits	Standard Kit	EP-kit
All-in-One		Beads / Polymerase	Hybridization Enhancer	Included	Not included



# CancerScreen Panel

**Core/50/100/400**

Somatic Cancer

## KEY FEATURES

1. Optimized panel for solid cancer	Assess DNA, RNA, and the whole CDS regions (RefSeq) of up to 407 genes and rearrangement regions associated with solid cancer
2. High sensitivity and specificity	Detect low-frequency and rare variants with high sequencing depths Capture the GC rich and homologous regions with Celeomics proprietary design technology
3. Cost-effective sequencing	Lower sequencing costs for 3 Gb sequencing amount compared to competitor product
4. Assess all variant types	Detect all mutation types including SNV, Indel, Large Indel, CNV, Rearrangement, MSI, and TMB in a single assay

## SPECIFICATION

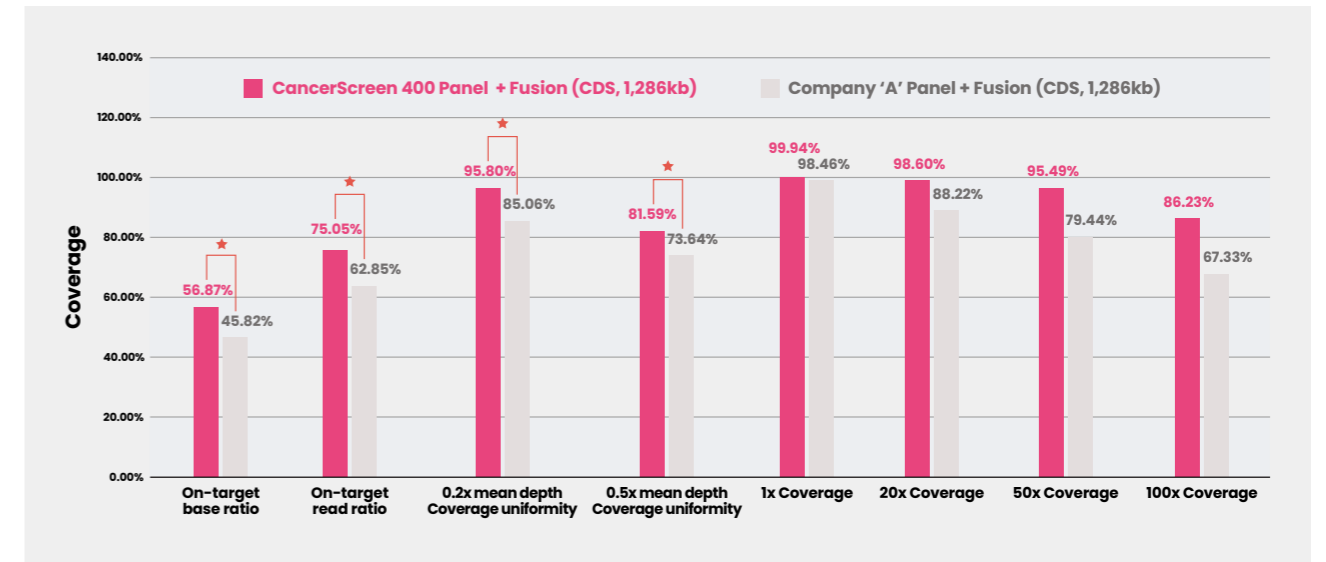
Gene count*	13 / 54 / 99 / 407 genes
Target size	61 / 197 / 299 / 1,123 kb + Rearrangement
Mutation type	SNV, Indel, CNV, Rearrangement, MSI, TMB
Sample type	FFPE, frozen tissue, cfDNA, RNA
Platform	All sequencers from Illumina, Thermo Fisher, MGI, PacBio, and Oxford Nanopore
Bioinformatics pipeline	Primary, Secondary and Tertiary analysis result (FASTQ to VCF, VCF to Clinical report)

\* Gene Add-On Service: Genes can be added by customer's request

## PANEL PERFORMANCE

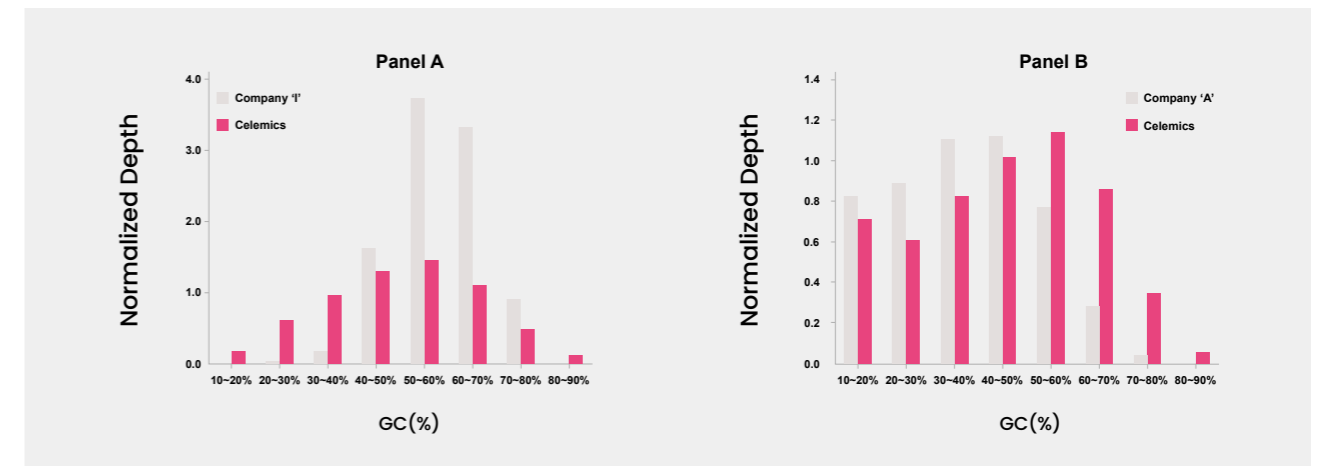
### Performance Comparison with Competitor Product

Higher on-target ratio, uniformity, and coverage at 100X compared to competitor product over the target regions including exons and introns (Compared with the same sequencing depth)



### Performance Comparison over GC-rich Regions

Higher uniform read depths over GC-rich regions compared to competitor product (Compared with the same sequencing depth)



## PACKAGE COMPOSITION

Package name	Compositions		Package option	Options	
Target Enrichment	Target capture Probe	-	Pooling method	Single Reaction	Pre-capture Pooling
Standard	Target Enrichment reagents	Library prep Kit	Library Preparation kits	Standard Kit	EP-kit
All-In-One	-	Beads / Polymerase	Hybridization Enhancer	Included	Not included

# CancerScreen Panel

## Core

### DESCRIPTION

The CancerScreen Core Panel is an NGS assay designed to detect all types of variants in 13 genes associated with somatic cancer. Targeting the selected genes with high sensitivity and specificity enables saving cost and effort. The report consists of the primary, secondary, and tertiary results for the In-depth understanding and interpretation of sequencing data.

### GENE LIST

CancerScreen Core	<b>ALK</b>	APC	BRAF	EGFR	ERBB2	KRAS	MET	NRAS	PIK3CA	<b>RET</b>	<b>ROS1</b>	SMAD4	TP53
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\* Genes in bold indicate fusion analysis

# CancerScreen Panel

## 50

### DESCRIPTION

The CancerScreen 50 Panel is an expanded NGS assay designed to detect all types of variants in over 50 genes associated with somatic cancer. Targeting the selected genes with high sensitivity and specificity enables saving cost and effort. The report consists of the primary, secondary, and tertiary results for the In-depth understanding and interpretation of sequencing data.

### GENE LIST

CancerScreen 50	ABL1	AKT1	<b>ALK</b>	APC	ATM	BRAF	BRCA1	BRCA2	CDH1	CDK4	CDK6	CDKN2A	CSF1R
	CTNNA1	DDR2	EGFR	ERBB2	ERBB4	ESR1	FGFR1	FGFR2	FGFR3	GNAI1	GNAQ	GNAS	HRAS
	IDH1	IDH2	JAK2	KDR	KIT	KRAS	MAP2K1	MET	MLH1	MTOR	MYC	MYCN	NOTCH1
	NRAS	<b>NTRK1</b>	PDGFRA	PIK3CA	PTCH1	PTEN	PTPN11	RBI	<b>RET</b>	<b>ROS1</b>	SMAD4	SMO	SRC
	STK11	TP53											

\* Genes in bold indicate fusion analysis

# CancerScreen Panel

## 100

### DESCRIPTION

The CancerScreen 100 Panel is an NGS assay for the comprehensive analysis of around 100 genes associated with somatic cancer. All types of variants are detected with high sensitivity and specificity. The report consists of the primary, secondary, and tertiary results for the In-depth understanding and interpretation of sequencing data.

### GENE LIST

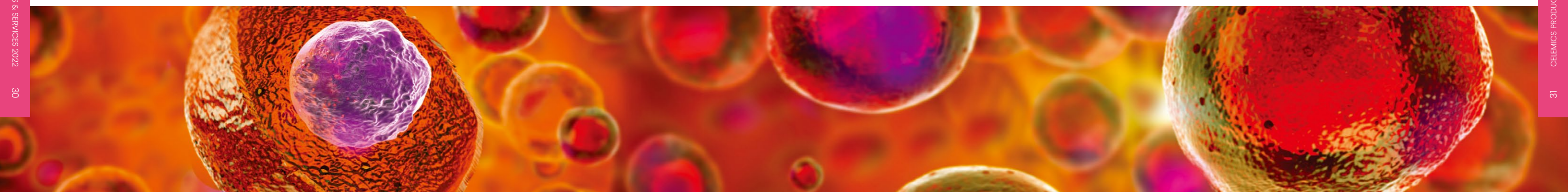
CancerScreen 100	ABL1	AKT1	AKT2	AKT3	ALK	APC	ARID1A	ARID1B	ARID2	ATM	ATRX	AURKA	AURKB
	BARD1	BCL2	BLM	BMP1A	BRAF	BRCA1	BRCA2	BRIPI	CDH1	CDK4	CDK6	CDKN2A	CHEK2
	CSF1R	CTNNA1	DDR2	EGFR	EPCAM	EPHA4	ERBB2	ERBB3	ERBB4	EZH2	FBXW7	FGFR1	FGFR2
	FGFR3	FLT3	GNAI1	GNAQ	GNAS	HNF1A	HRAS	IDH1	IDH2	IGF1R	ITK	JAK1	JAK2
	JAK3	KDR	KIT	KRAS	MDM2	MET	MLH1	MPL	MRE11	MSH2	MSH6	MTOR	MUTYH
	NBN	NF1	NOTCH1	NPM1	NRAS	NTRK1	PALB2	PDGFRA	PDGFRB	PIK3CA	PIK3R1	PMS2	PRSS1
	PTCH1	PTCH2	PTEN	PTPN11	RAD50	RAD51C	RAD51D	RBI	RET	ROS1	SLX4	SMAD4	SMARCB1
	SMO	SRC	STK11	SYK	TERT	TOPI	TP53	VHL					

# CancerScreen Panel

## 400

### DESCRIPTION

The CancerScreen 400 Panel is an NGS assay designed to detect all types of variants in over 400 genes associated with somatic cancer. Targeting the selected genes with high sensitivity and specificity enables saving cost and effort. The report consists of the primary, secondary, and tertiary results for the In-depth understanding and interpretation of sequencing data.





# CancerScreen Panel

## 400

### GENE LIST

ABL1	ABL2	ADGRA2	AKT1	AKT2	AKT3	ALK	AMER1	APC	APCDD1	APEX1	APOB	APOBEC1
AR	ARAF	ARFRP1	ARID1A	ARID1B	ARID2	ASXL1	ATM	ATP11B	ATR	ATRX	AURKA	AURKB
AXINI	AXL	B2M	B3GAT1	BACH1	BAP1	BARD1	BCL2	BCL6	BCL9	BCOR	BCR	BIRC2
BIRC3	BLM	BRAF	BRCA1	BRCA2	BRD2	BRD3	BRD4	BRIPI	BTG1	BTK	BLA	CARD11
CASP5	CASP8	CBFB	CBL	CDK12	CDK4	CDK6	CDK8	CDKN1A	CDKN1B	CDKN2A	CDKN2B	CDKN2C
CDX2	CEBPA	CHD1	CHD2	CHD4	CHEK1	CHEK2	CHUK	CIC	CRBN	CREBBP	CRKL	CRLF2
CSF1R	CSF2	CSF2RA	CSF2RB	CSNK2A1	CTCF	CTLA4	CTNNA1	CTNNB1	CUL3	CUL4A	CUL4B	CXCL10
CXCL11	CXCL9	CXCR3	CYLD	CYP17A1	DAXX	DCUN1D1	DDR2	DICER1	DIS3	DNMT1	DNMT3A	DOCK2
DOT1L	EGFR	ELMO1	EML4	EMSY	EP300	EPHA3	EPHA5	EPHA6	EPHA7	EPHB1	EPHB4	EPHB6
ERBB2	ERBB3	ERBB4	ERCC1	ERCC2	ERG	ERRF1	ESR1	ETV1	ETV4	ETV5	ETV6	EWSR1
EYA2	EZH2	FANCA	FANCC	FANCD2	FANCE	FANCF	FANCG	FANCI	FANCL	FANCM	FAS	FAT1
FAT3	FBXW7	FGF1	FGF10	FGF12	FGF14	FGF19	FGF2	FGF23	FGF3	FGF4	FGF6	FGF7
FGFR1	FGFR2	FGFR3	FGFR4	FH	FLCN	FLT1	FLT3	FLT4	FOXA1	FOXL2	FOXO3	FOXP3
FRS2	FUBP1	GABRA6	GAS6	GATA1	GATA2	GATA3	GATA4	GATA6	GID4	GLI1	GNAI1	GNAI3
GNAQ	GNAS	GRIN2A	GRM3	GSK3B	GUCY1A2	GZMA	GZMB	GZMH	H3F3A	HGF	HIST1H3B	HNFA
HOXA3	HRAS	HSD3B1	HSP90AA1	IDH1	IDH2	IDO1	IDO2	IFITM1	IFITM3	IFNA1	IFNB1	IFNG
IGF1	IGF1R	IGF2	IGF2R	IKBKE	IKZF1	IL12A	IL12B	IL2	IL23A	IL6	IL7R	INHBA
INPP4B	INSR	IRF2	IRF4	IRS2	ITGAE	ITK	JAK1	JAK2	JAK3	JUN	KAT6A	KDM5A
KDM5C	KDM6A	KDR	KEAP1	KEL	KIT	KLF4	KLHL6	KMT2A	KMT2B	KMT2C	KNSTRN	KRAS
LAG3	LMO1	LRP1B	LRP6	LTK	LYN	LZTR1	MAGI2	MAGOH	MAML1	MAP2K1	MAP2K2	MAP2K4
MAP3K1	MAP3K13	MAPK1	MAX	MCL1	MDM2	MDM4	MED12	MEF2B	MEN1	MET	MITF	MLH1
MPL	MRE11	MSH2	MSH6	MTOR	MUTYH	MYB	MYC	MYCL	MYCN	MYD88	MYO18A	NCOA3
NCOR1	NF1	NF2	NFE2L2	NFKB1A	NOTCH1	NOTCH2	NOTCH3	NOTCH4	NPM1	NRAS	NSD1	NSD3
NTRK1	NTRK2	NTRK3	NUP93	NUTM1	PAK3	PAK5	PALB2	PARP1	PARP2	PARP3	PARP4	PAX5
PBRM1	PDCD1	PDCD1LG2	PDGFRA	PDGFRB	PDK1	PGR	PHF6	PHLPP2	PIK3C2B	PIK3C3	PIK3CA	PIK3CB
PIK3CG	PIK3R2	PKHD1	PLCG1	PLCG2	PMS2	PNP	PNRC1	POLD1	POLE	PPARG	PPP2R1A	PRDM1
PREX2	PRF1	PRKARIA	PRKCI	PRKDC	PRPF40B	PRSS8	PTCHI	PTCH2	PTEN	PTK2	PTPN11	PTPRC
PTPRD	QKI	RAB35	RAC1	RAC2	RAD17	RAD50	RAD51	RAD52	RAD54L	RAF1	RANBP2	RARA
RBI	RBM10	REL	RET	RHEB	RHOA	RHOB	RICTOR	ROBO1	ROBO2	ROS1	RPA1	RPS6KB1
RPTOR	RUNX1	RUNX1T1	RUNX3	SDHA	SDHB	SDHC	SDHD	SEMA3A	SEMA3E	SET	SETBP1	SETD2
SF3A1	SF3B1	SH2B3	SKP2	SIT2	SMAD2	SMAD3	SMAD4	SRSF2	SRSF7	STAG2	STAT3	STAT4
TERT	TET2	CD274	TP53									

CancerScreen 400



# CancerMaster Panel

Somatic Cancer

## DESCRIPTION

The CancerMaster Panel is designed to detect all variant types and immuno-oncology markers (MSI and TMB), which are crucial biomarkers for cancer immunotherapy. For CNV analysis, different cut-offs are applied according to the ratio of cancer cells. The panel is also designed to detect Epstein-Barr virus (EBV) and Human Papillomaviruses (HPV), allowing for the comprehensive analysis of cancer-associated genes.

## KEY FEATURES

### 1. Comprehensive analysis of cancer-associated genes

A broad range of targeting elements including somatic variants, IO-signatures (TMB, MSI), EBV and HPV, for clinical diagnoses of different cancer types and applications to precision medicine

### 2. Extensive validation studies

Robust panel performance supported by extensive validation tests with reference and clinical specimens

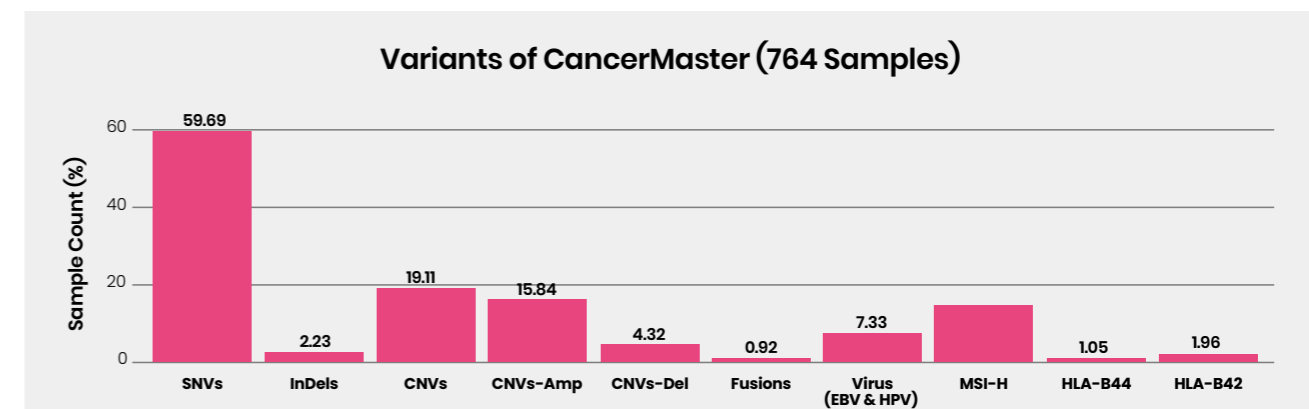
## SPECIFICATION

Gene count*	524 genes
Covered region	Whole CDS, custom regions of oncogenes, immune response genes, and EBV & HPV viruses
Target size	2.5 Mb
Mutation type	SNV, Indel, CNV, Rearrangement, TMB, MSI, EBV, HPV
Sample type	FFPE, Fresh frozen tissue (> 50 ng of fragmented DNA)
Platform	All sequencers from Illumina, Thermo Fisher, MGI, PacBio, and Oxford Nanopore
Bioinformatics pipeline	Primary, Secondary and Tertiary analysis result (FASTQ to VCF, VCF to Clinical report)
Publication	Molecular Characterization of Biliary Tract Cancer Predicts Chemotherapy and PD-1/PD-L1 Blockade Responses, Hepatology, 2021

\* Gene Add-On Service: Genes can be added by customer's request

## PANEL PERFORMANCE

The probes are designed to include the intron regions as well as clinically significant biomarkers. By conducting extensive validation studies with clinical samples, the panel was examined to show its performance with high sensitivity and specificity in detecting the variants in cancer-associated genes.

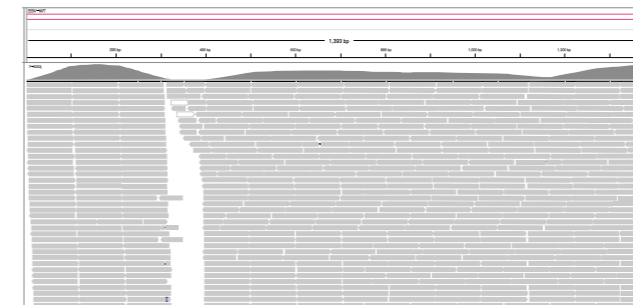


## ANALYSIS OF EBV & HPV

### EBV (Epstein-Barr Virus)

- Related disease – Lymphoma
- Genes – EBV type 1 (EBNA-2)

Validation for detection of EBV type 1 (EBNA-2) in control specimens



### HPV (Human Papillomavirus)

- Related disease – Cervical cancer
- Genes – HPV L1 gene  
(Analysis of a total of 24 types is possible)

Analysis of the following 11 types of HPV types was completed using clinical specimens

Human infection HPV list
Human papillomavirus type 178
Human papillomavirus type 136
Human papillomavirus type 140
Human papillomavirus type 154
Human papillomavirus type 156
Human papillomavirus type 179
Human papillomavirus type 201
Human papillomavirus type 49
Human papillomavirus type 9
Human papillomavirus type 92
Human papillomavirus type 96

## PACKAGE COMPOSITION

Package name	Compositions		Package option	Options	
Target Enrichment	Target capture Probe	-	Pooling method	Single Reaction	Pre-capture Pooling
Standard	Target Enrichment reagents	Library prep Kit	Library Preparation Kits	Standard Kit	EP-kit
All-in-One		Beads / Polymerase	Hybridization Enhancer	Included	Not included

# READY-TO-USE PANELS FOR INHERITED DISEASE

CELEMICS PRODUCTS & SERVICES 2022

G-Mendeliome CES Panel  
: Standard / Expanded  
G-Mendeliome Disease-Specific Panel



# G-Mendeliome CES Panel

## Standard / Expanded

Hereditary Diseases

### DESCRIPTION

The G-Mendeliome CES Panel has overcome the limitations of analyzing clinical diseases with whole exome sequencing. By selectively targeting the clinically significant genes, the panel enables comprehensive analysis with the most effective sequencing throughput.

### KEY FEATURES

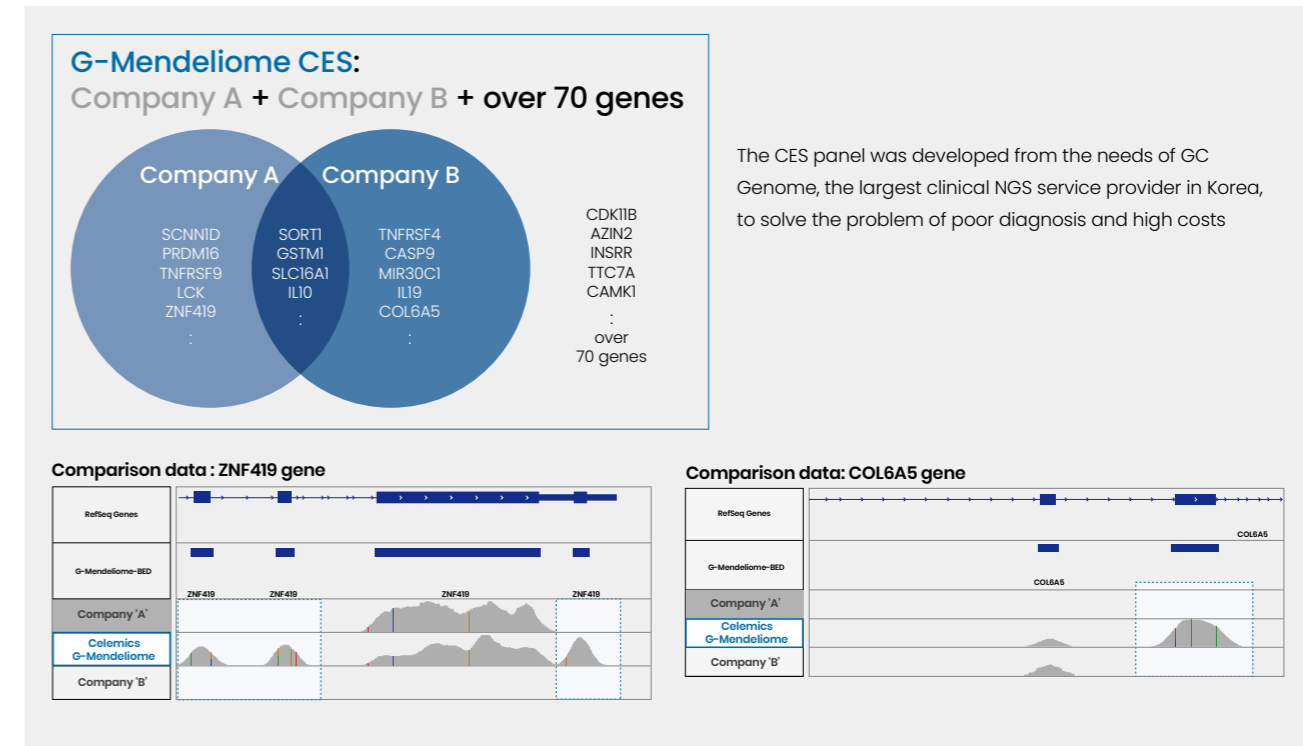
1. Comprehensive genomic profiling of a variety of genetic diseases	Includes 7,000 genes associated with clinically significant genetic diseases
2. A wide range of target regions	Includes all clinically significant regions that are not covered from competitor panels
3. Cost-effective analysis	Able to provide accurate analysis with reduced sequencing costs compared to WES

### SPECIFICATION

Gene count*	5,508 / 7,513 genes
Covered region	CDS, hotspots, Mitochondrial genome
Target size	13.8 / 19.6 Mb
Mutation type	SNV, Indel, CNV
Sample type	Blood (> 50 ng of fragmented DNA)
Platform	All sequencers from Illumina, Thermo Fisher, MGI, PacBio, and Oxford Nanopore
Bioinformatics pipeline	Primary, Secondary and Tertiary analysis result (FASTQ to VCF, VCF to Clinical report)

### PANEL PERFORMANCE

	Celemics	Company A	Company B
On-Target Read Ratio	82.8%	65.9%	80.8%



### PACKAGE COMPOSITION

Package name	Compositions		Package option	Options	
Target Enrichment	Target capture Probe	-	Pooling method	Single Reaction	Pre-capture Pooling
Standard	Target Enrichment reagents	Library prep Kit	Library Preparation kits	Standard Kit	EP-kit
All-in-One	-	Beads / Polymerase	Hybridization Enhancer	Included	Not included



## LIST OF DISEASES ASSESSED BY G-MENDELIOME CES PANEL

Category	Related Diseases
Cardiology	Aortopathy and connective tissue disorders
	Arrhythmia
	Cardiomyopathy
	Congenital heart defect
	Dyslipidemia
	Other cardiovascular diseases
	Pulmonary hypertension
Dermatology	Adams-Oliver syndrome
	Albinism
	Cardiofaciocutaneous syndrome
	Cutis laxa
	Dyskeratosis congenita
	Ectodermal dysplasia
	Ehlers-Danlos syndrome
	Epidermolysis bullosa
	Hereditary acrodermatitis enteropathica
	Hermansky-Pudlak syndrome
	Hypotrichosis
	Ichthyosis
	Neurofibromatosis
	Pachyonychia congenita
	Palmoplantar keratoderma
	Progeria and Progeroid Syndromes
	Skin cancer
	Tuberous sclerosis
	Waardenburg syndrome
Xeroderma pigmentosum	
Endocrinology	Adrenal hyperplasia
	Diabetes
	Hyperinsulinism
	Hyperparathyroidism
	Hypothyroidism
	Kallmann syndrome
	Multiple endocrine neoplasia
	Obesity
	Pancreatitis
	Premature ovarian failure
	ENT
GI/Hepatology	Cholestasis
	Congenital diarrhea
	Congenital hepatic fibrosis
	Gastrointestinal atresia
	Hirschsprung disease
Polycystic liver disease	
Hematology	Anemia
	Bleeding&Thrombotic disorder
	Bone marrow failure
	Congenital neutropenia
	Hemochromatosis
RBC membrane disorder	
Immunology	Antibody deficiencies
	Autoinflammatory disorders
	Combined T/B cell deficiencies
	Complement deficiencies
	Defects in intrinsic and innate immunity
	Immune dysregulation
	Phagocytic defects

Category	Related Diseases
Metabolism	Aminoacidopathies
	Carbohydrate disorders
	Congenital disorders of glycosylation
	Creatine biosynthesis disorders
	Fatty acid oxidation defects
	Lipodystrophy
	Lysosomal storage disorders
	Organic acidemias
	Peroxisomal disorders
	Porphyria
	Purine/Pyrimidine metabolism disorders
	Pyruvate metabolism and tricarboxylic acid cycle defects
	Urea cycle disorders
	Nephrology
Ciliopathies	
Diabetes insipidus	
Hemolytic uremic syndrome	
Hypokalemia	
Hypomagnesemia	
Hypophosphatemic rickets	
Nephrolithiasis	
Nephrotic syndrome/Focal glomerulonephrosis	
Pseudohypoadosteronism	
Renal malformation	
Renal tubular acidosis	
Neurology	
	Movement disorders
	Neurodegenerative disorders
	Neuromuscular disorders
	Neuropathies and related disorders
	Seizures and Brain abnormalities
	Oncology
Colorectal cancer	
Endocrine cancer	
Gastrointestinal cancer	
Hematologic malignancy	
Lung cancer	
Nervous system/brain cancer	
Pancreatic cancer	
Prostate cancer	
Renal cancer	
Sarcoma	
Skin cancer	
Ophthalmology	
	Cataract/Ectopia lentis
	Corneal dystrophy
	Glaucoma
	Microphthalmia/Anophthalmia
	Nystagmus
	Ophthalmoplegia/Oculomotor apraxia
	Optic atrophy
	Retinal dystrophy
	Retinoblastoma
Pulmonology	Bronchiectasis
	Central hypoventilation/Apnea
	Cystic fibrosis
	Cystic lung disease
	Hermansky-Pudlak syndrome
	Interstitial lung disease
	Primary ciliary dyskinesia
Surfactant dysfunction	
Skeletal disorders	Amelogenesis imperfecta
	Arthrogryposes
	Cleft lip palate
	Craniosynostosis
	Exostosis
	Facial dysostosis
	Macrocephaly/Overgrowth syndrome
	Osteopetrosis
	Short stature syndrome
	Skeletal dysplasia

# G-Mendeliome Disease-Specific Panel

## KEY FEATURES

### 1. Comprehensive analysis of a broad range of diseases

Identifying diseases associated with:  
Acute lymphatic leukemia, Acute Myeloid Leukemia, Cardiac disease, Coagulation, Epilepsy, Hearing loss, Inborn errors of metabolism, Lymphoma, Lysosomal storage disease, Common hereditary cancer for a medical checkup, Neuromuscular disease, Parkinson's disease, Alzheimer's disease, Dementia, Dystonia, RASopathies, Retinitis pigmentosa, Short stature, Skin disease, and Somatic cancer

### 2. Collaboration with the leading CRO in the country

Developed 17 different panels for assessing genes of related diseases

## SPECIFICATION

<b>Gene count*</b>	Ranges from 14 to 293 genes
<b>Covered regions</b>	Whole CDS, hotspots
<b>Target size</b>	37-1,159 kb
<b>Mutation type</b>	SNV, Indel, CNV
<b>Sample type</b>	Differs by somatic or germline panel
<b>Platform</b>	All sequencers from Illumina, Thermo Fisher, MGI, PacBio, and Oxford Nanopore
<b>Bioinformatics pipeline</b>	Primary, Secondary and Tertiary analysis result (FASTQ to VCF, VCF to Clinical report)

\* Gene Add-On Service: Genes can be added by customer's request

## PACKAGE COMPOSITION

Package name		Compositions		Package option		Options	
Target Enrichment	Target capture Probe		-	Pooling method	Single Reaction	Pre-capture	Pooling
Standard	Target Enrichment reagents	Library prep Kit	-	Library Preparation kits	Standard Kit	EP-kit	
All-In-One		Beads / Polymerase		Hybridization Enhancer	Included	Not included	

## LIST OF PANELS FOR VARIOUS DISEASES

Panel Name	Related Diseases	Gene List																																																																																																																																																																																							
Alzheimer-Parkinson-Dementia Panel	Alzheimer's disease, Parkinson's disease, Dementia, Dystonia	AARS	ABCA13	ABCA7	ABCB11	ADCY5	ALS2	ANG	ANO3	APP	ATPI3A2	ATPIA3	ATP7B	C19orf12	CACNA1B	CHCHD10	CHMP2B	CHRNA4	CIZ1	COG1	COL4A4	COL6A3	DAO	DCTN1	DNMT1	EVC	FERMT1	FIG4	FREM2	FUS	GBA	GCH1	GNAL	GNAO1	GRM1	GRN	HNRNPA1	HNRNPA2B1	HPCA	HPSE2	IL12RB2	KCTD17	KMT2B	L2HGDH	LAMA3	LRRK2	MAPT	MATR3	MECR	NDUFV3	NEK1	NPHS2	OPTN	PANK2	PARK7	PDP1	PINK1	PLA2G6	PNKD	PRKN	PRKRA	PRNP	PRRT2	PSENI	PSEN2	RELN	SERPIND1	SETX	SGCE	SIGMAR1	SLC12A6	SLC19A3	SLC2A1	SLC30A10	SLC6A3	SNCA	SOD1	SORL1	SOX6	SPG11	SQSTM1	SRY	SUMF1	TAF1	TAF15	TARDBP	TBKI	TDRD7	TH	THAP1	TIMM8A	TORIA	TREM2	TUBA4A	TUBB4A	UBQLN2	VAC14	VAPB	VCAN	VCP	VPSI3A	WNK1																																																																																			
		Bleeding Disorder-Coagulopathy Panel	Bleeding Disorder, Coagulation	AARS	ABCA1	ABCA13	ABCB11	ACTN1	ANKRD26	ANO6	AP3B1	BLOC1S3	BLOC1S6	BRCA1	BRCA2	BRIPI	CD36	CDAN1	COG1	COL4A4	CYCS	DDX41	DKC1	DNMT1	DTNBPI	ELANE	ERCC4	ETV6	EVC	F10	F11	F13A1	F13B	F2	F5	F7	F8	F9	FANCA	FANCB	FANCC	FANCD2	FANCE	FANCF	FANCG	FANCI	FANCL	FANCM	FERMT1	FERMT2	FGA	FBG	FGG	FLII	FREM2	FYB1	GATA1	GATA2	GFI1	GPIIB	GPIBA	GPIBB	GP6	GP9	GRM1	HAX1	HOXA11	HPS1	HPS3	HPS4	HPS5	HPS6	HPSE2	IFNG	IL12RB2	ITGA2B	ITGB3	L2HGDH	LAMA3	LMAN1	LYST	MASTL	MCFD2	MLPH	MPL	MYH9	MYO5A	NBEAL2	NBN	NDUFV3	NHP2	NOPI0	NPHS2	P2RY12	PALB2	PDP1	PLA2G4A	PLAU	PRF1	PRKACG	RAB27A	RAD51C	RASGRP2	RBM8A	RPL11	RPL35A	RPL5	RPS10	RPS19	RPS24	RPS26	RPS7	RUNX1	SBDS	SEC23B	SERPIND1	SERPINE1	SERPINF2	SLC12A6	SLFN14	SLX4	SOX6	SRC	SRP72	SRY	STIM1	SUMF1	TBXA2R	TBXAS1	TDRD7	TERC	TERT	TINF2	UBE2T	VCAN	VIPAS39	VPS33B	VWF	WAS	WIPF1	WNK1	XRCC2																																											
				Cardiovascular Panel	Cardiac diseases	ABCC9	ABCG5	ABCG8	ACTA1	ACTA2	ACTC1	ACTN2	AKAP9	ALMS1	ANK2	ANKRD1	APOA4	APOA5	APOB	APOC2	APOE	BAG3	BRAF	CACNA1C	CACNA2D1	CACNB2	CALM1	CALR3	CASQ2	CAV3	CBL	CBS	CETP	COL3A1	COL5A1	COL5A2	COX15	CREB3L3	CRELD1	CRYAB	CSR3P3	CTFI	DES	DMD	DNAJC19	DOLK	DPP6	DSC2	DSG2	DSP	DTNA	EFEMP2	ELN	EMD	EYA4	FBN1	FBN2	FHL1	FHL2	FKRP	FTN	FXN	GAA	GATADI	GCKR	GJA5	GLA	GPD1L	GPIHBP1	HADHA	HCN4	HFE	HRAS	HSPB8	ILK	JAG1	JPH2	JUP	KCNA5	KCND3	KCNE1	KCNE2	KCNE3	KCNH2	KCNJ2	KCNJ5	KCNJ8	KCNQ1	KLF10	KRAS	LAMA2	LAMA4	LAMP2	LDB3	LDLR	LDLRAP1	LMF1	LMNA	LPL	LTBP2	MAP2K1	MAP2K2	MIB1	MURC	MYBPC3	MYH11	MYH6	MYH7	MYL2	MYL3	MYLK	MYLK2	MYO6	MYOZ2	MYPN	NEXN	NKX2-5	NODAL	NOTCH1	NPPA	NRAS	PCSK9	PDLIM3	PKP2	PLN	PRDM16	PRKAG2	PRKARIA	PTPN11	RAF1	RANGRF	RBM20	RYR1	RYR2	SALL4	SCN1B	SCN2B	SCN3B	SCN4B	SCN5A	SCO2	SDHA	SEPN1	SGCB	SGCD	SGCG	SHOC2	SLC25A4	SLC2A10	SMAD3	SMAD4	SNTA1	SOS1	SREBF2	TAZ	TBX20	TBX3	TBX5	TCAP	TGFB2	TGFB3	TGFBRI	TGFBRI	TMEM43	TMPO	TNNC1	TNNI3	TNNT2	TPM1	TRDN	TRIM63	TRPM4	TTN	TTR	TXNRD2	VCL	ZBTB17	ZHX3	ZIC3						
						Common Hereditary Cancer Panel	Medical checkup	APC	ATM	ATRX	BARD1	BMPRIA	BRAF	BRCA1	BRCA2	BRIPI	CDHI	CDKN2A	CHEK2	EGLN1	EGLN2	EPAS1	EPCAM	FGFR1	FH	H3F3A	HRAS	IDH2	KIF1B	KMT2D	MAX	MDH2	MEN1	MERTK	MET	MLH1	MRE11	MSH2	MSH6	MUTYH	NBN	NFI	NF2	PALB2	PMS2	POLD1	POLE	PRSS1	PTEN	RAD50	RAD51C	RAD51D	RBI	RET	SDHA	SDHAF2	SDHB	SDHC	SDHD	SMAD4	SPINK1	STK11	TMEM127	TP53	TSC1	TSC2	VHL	WT1																																																																																																																					
								Epilepsy Panel	Epilepsy	AARS	ABCA13	ABCB11	ADGRV1	ADSL	ALDH7A1	ALG13	ARHGFE15	ARHGFE9	ARX	ASAH1	ATPIA2	ATP6AP2	CACNA1A	CASK	CDKL5	CHD2	CHRNA2	CHRNA4	CHRNA7	CHRN2	CLCN4	CLN3	CLN5	CLN6	CLN8	CNTNAP2	COG1	COL4A4	CSTB	CTSD	DCX	DEPDC5	DLG3	DNAJC5	DNMI	DNMT1	DOCK7	DYRK1A	EEF1A2	EPM2A	EVC	FERMT1	FOLR1	FOXG1	FREM2	GABRA1	GABRA2	GABRB3	GABRG2	GAMT	GATM	GNAO1	GOSR2	GRIN1	GRIN2A	GRIN2B	GRM1	HCN1	HDAC4	HNRNPU	HPSE2	IL12RB2	IQSEC2	KANSL1	KCNA2	KCNB1	KCNH5	KCNJ10	KCNMA1	KCNQ2	KCNQ3	KCNT1	KCTD7	L2HGDH	LAMA3	LGII	MAGI2	MBD5	MCEP2	MEF2C	MFSD8	NDUFV3	NECAP1	NHLRC1	NPHS2	NR2F1	NRXN1	PCDH19	PDP1	PIGA	PIGO	PIGQ	PIGV	PLCB1	PNKP	PNPO	POLG	PPT1	PRICKLE1	PRICKLE2	PRRT2	QARS	RELN	SCARB2	SCN1A	SCN1B	SCN2A	SCN8A	SCN9A	SERPIND1	SLC12A6	SLC13A5	SLC25A22	SLC2A1	SLC35A2	SLC6A8	SLC9A6	SMS	SOX6	SPTANI	SRPX2	SRY	ST3GAL3	STXBPI	SUMF1	SYN1	SYNGAP1	SYNJ1	SZT2	TBCID24	TCF4	TDRD7	TPP1	TSCI	TSC2	UBE3A	VCAN	WDR45	WNK1	WWOX	ZEB2																																		



# READY-TO-USE PANELS FOR PHARMACO- GENOMICS

CELEMICS PRODUCTS & SERVICES 2022

PharmacoScreen Panel  
· Standard / Epilepsy / Anti-tuberculosis

 **CELEMICS**





# PharmacoScreen Panel

## Standard / Epilepsy / Anti-tuberculosis

### Drug Metabolism

#### DESCRIPTION

The main target of PharmacoScreen Panel is the genes associated with prescribed drugs of the corresponding diseases. The assay allows for precise selection and dosage of prescribed drugs, and detection of genetic variants associated with drug metabolism, epilepsy and anti-tuberculosis.

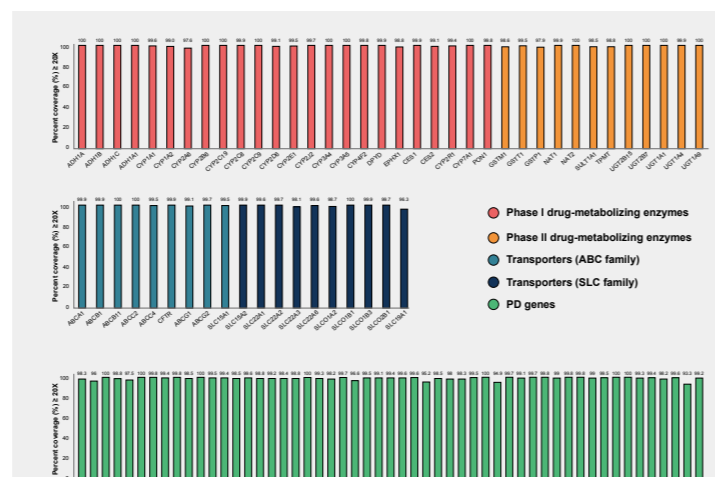
#### KEY FEATURES

<b>1. Assess extensive target regions associated with pharmacogenomics</b>	Target over 120 genes associated with pharmacokinetics and pharmacodynamics
<b>2. Validated panel performance</b>	Collaborated with 4 major university hospitals on a government project Complete validation for clinical application
<b>3. Flexible panel contents</b>	PharmacoScreen Panels for drug metabolism, epilepsy, and anti-tuberculosis.

#### PANEL PERFORMANCE

The panel performance test resulted in 99.9% specificity and 99.7% sensitivity.

- 1.1 Phase I/II drug-metabolizing enzyme (Drug metabolism)
- 1.2 ABC & SLC family transporter genes (Drug effect)
- 1.3 Pharmacodynamics genes (Drug biochemical and physiological)
- 1.4 Modifier genes (Drug ADME enhancement)



#### PACKAGE COMPOSITION

Package name	Compositions		Package option	Options	
Target Enrichment	Target capture Probe	-	Pooling method	Single Reaction	Pre-capture Pooling
Standard	Target Enrichment reagents	Library prep Kit	Library Preparation kits	Standard Kit	EP-kit
All-In-One		Beads / Polymerase	Hybridization Enhancer	Included	Not included

# PharmacoScreen Standard

#### DESCRIPTION

One of the major problems of organ transplantation is tissue damage by rejection and relapse of the disease after transplantation. Although applying immunosuppressive drugs can prevent rejection, determining the proper dosage of immunosuppressive drugs for an individual patient is challenging. The PharmacoScreen Standard Panel is an NGS assay, designed to assess 122 genes associated with pharmacogenomics, including drug metabolism (Phase I, II), Transporters (ABC and SLC families), and Parkinson's disease-related genes (PD genes). The panel is not limited to 122 genes, and more genes of interest can be added through our Gene Add-on service.

#### SPECIFICATION

<b>Gene count*</b>	122 genes
<b>Covered region</b>	Whole CDS, UTR (-50 bp, +10 bp)
<b>Target Size</b>	534 kb
<b>Mutation Type</b>	SNV, Indel, CNV
<b>Sample type (amount)</b>	Blood (> 50 ng of fragmented DNA)
<b>Platform</b>	All sequencers from Illumina, Thermo Fisher, MGI, PacBio, and Oxford Nanopore
<b>Sensitivity / Specificity</b>	100% / 94.5%
<b>Bioinformatics pipeline</b>	Primary, Secondary and Tertiary analysis result (FASTQ to VCF, VCF to Clinical report)
<b>Publications</b>	Targeted Next-Generation Sequencing for Comprehensive Genetic Profiling of Pharmacogenes, Clinical Pharmacology & Therapeutics, 2016

\* Gene Add-On Service: Genes can be added by customer's request

## GENE LIST

- Phase I drug-metabolizing enzymes
- Phase II drug-metabolizing enzymes
- ▲ Transporters (ABC family)
- ▲ Transporters (SLC family)
- PD genes
- ◆ Modifier genes

● ADH1A	● GSTM1	▲ ABCA1	■ ACE	■ ADRB1	■ KCNH2	◆ AHR
● ADH1B	● GSTP1	▲ ABCB1	■ ADRB2	■ ALOX5	■ LDLR	◆ KCNJ11
● ADH1C	● GSTT1	▲ ABCB11	■ BRCA1	■ APOA1	■ MAOA	◆ NR1I3
● ALDH1A1	● NAT1	▲ ABCB2	■ COMT	■ ARID5B	■ NR3C2	◆ NR1I2
● CES1	● NAT2	▲ ABCB3	■ DRD2	■ BDNF	■ NTRK2	◆ POR
● CES2	● SULT1A1	▲ ABCB4	■ F5	■ CACNA1C	■ PEAR1	◆ SOD2
● CYP1A1	● TPMT	▲ ABCB7	■ HMGCR	■ CPS1	■ PTGS1	
● CYP1A2	● UGT1A1	▲ ABCG1	■ MTHFR	■ CRHR1	■ PTGS2	
● CYP2A6	● UGT1A4	▲ ABCG2	■ NQO1	■ DBH	■ RYR1	
● CYP2B6	● UGT1A9	▲ SLC10A1	■ P2RY1	■ DRD1	■ RYR2	
● CYP2C19	● UGT1A10	▲ SLC15A1	■ P2RY12	■ EGFR	■ SCN1A	
● CYP2C8	● UGT2B15	▲ SLC15A2	■ PTGIS	■ ESRI	■ SCN2A	
● CYP2C9	● UGT2B7	▲ SLC19A1	■ SCN5A	■ FKBP5	■ SLC47A1	
● CYP2D6		▲ SLC22A1	■ TYMS	■ GLCC1	■ SLC47A2	
● CYP2E1		▲ SLC22A2	■ VDR	■ GRK4	■ SLC6A3	
● CYP2J2		▲ SLC22A3	■ VKORC1	■ GRK5	■ SLC6A4	
● CYP2R1		▲ SLC22A4		■ G6PD	■ TBXAS1	
● CYP3A4		▲ SLC22A5		■ HTR1A	■ ZNF423	
● CYP3A5		▲ SLC22A6		■ HTR2A		
● CYP4F2		▲ SLC22A8				
● CYP7A1		▲ SLC22A11				
● DPYD		▲ SLC22A12				
● EPHX1		▲ SLC01A2				
● PON1		▲ SLC01B1				
		▲ SLC01B3				
		▲ SLC02B1				

# PharmacoScreen

## Epilepsy

### DESCRIPTION

The PharmacoScreen Epilepsy Panel, designed for research studies on epilepsy, consists of 91 genes associated with anti-epileptic drugs. Epilepsy is one of the most common neurological disorders, with its estimated prevalence is one out of 100 worldwide and constantly increasing. Epilepsy is usually treated by consistent application of anti-epileptic drugs. The aim of the treatment is to prevent seizures with no issues of side effects. Although over 20 different anti-epileptic drugs have been developed, most of the drugs failed to prevent seizures, or faced challenges of determining the proper dosage for an individual patient. The genetic factor is one of clinical factors to be considered.

### SPECIFICATION

Gene count*	91 genes
Covered region	Whole CDS + UTR (-50 bp, +10 bp)
Target size	575 kb
Mutation type	SNV, Indel, CNV
Sample type (amount)	Blood (> 50 ng of fragmented DNA)
Platform	All sequencers from Illumina, Thermo Fisher, MGI, PacBio, and Oxford Nanopore
Bioinformatics pipeline	Primary, Secondary and Tertiary analysis result (FASTQ to VCF, VCF to Clinical report)

\* Gene Add-On Service: Genes can be added by customer's request

### GENE LIST

PharmacoScreen Panel Epilepsy	ANKK1	CACNA1A	CACNA1B	CACNA1D	CACNA1E	CACNA1F	CACNA1G	CACNA1H	CACNA1I	CACNA1S	CACNA2D1	CACNA2D2	CACNA2D3	
	CACNA2D4	CACNB1	CACNB2	CACNB3	CACNB4	CACNG1	CACNG2	CACNG3	CACNG4	CACNG5	CACNG6	CACNG7	CACNG8	
	CDHI3	CLCN2	EFHC1	GABRA1	GABRA2	GABRA3	GABRA4	GABRA5	GABRA6	GABRB1	GABRB2	GABRB3	GABRD	
	GABRE	GABRG1	GABRG2	GABRG3	GABRP	GABRQ	GABRR1	GABRR2	GABRR3	GRIAI	GRIA2	GRIA3	GRIA4	
	GRIK1	GRIK2	GRIK3	GRIK4	GRIK5	GRIN1	GRIN2A	GRIN2B	GRIN2C	GRIN2D	GRIN3A	GRIN3B	HNF4A	
	HTR1B	KCNA2	KCNB1	KCNC1	KCND3	KCNH1	KCNJ10	KCNQ2	KCNQ3	KCNT1	KCNTD7	LEPR	MAOA	
	MAOB	RBFOX1	SCN1A	SCN2A	SCN3A	SCN8A	STS	TPH1	TPH2	UGT1A10	UGT1A6	UGT1A7	UGT1A9	

# PharmacoScreen

## Anti-tuberculosis

### DESCRIPTION

The PharmacoScreen Anti-tuberculosis Panel assesses genes associated with liver injury. Drug-induced liver injury (DILI), which is an important cause of acute liver failure, can be a threat to a patient and a common reason why some drug development projects are discontinued. According to a spontaneous reporting database from a research network of pharmacovigilance institutions in Korea, anti-tuberculosis drugs are reported to be the most common factor that leads to DILI demanding precise and personalized medicine.

### SPECIFICATION

Gene count*	132 genes
Covered regions	Whole CDS + UTR (-50 bp, +10 bp)
Target size	186 kb
Mutation type	SNV, Indel, CNV
Sample type (amount)	Blood (> 50 ng of fragmented DNA)
Platform	All sequencers from Illumina, Thermo Fisher, MGI, PacBio, and Oxford Nanopore
Bioinformatics pipeline	Primary, Secondary and Tertiary analysis result (FASTQ to VCF, VCF to Clinical report)

\* Gene Add-On Service: Genes can be added by customer's request

### GENE LIST

PharmacoScreen Panel Anti-tuberculosis	ABHD5	ADA	ADORA2A	ALAS1	ALPK2	ANO10	ASAH1	BACH1	BAX	BCL2	BTLA	CARD8	CASP1
	CASP3	CASP8	CASP9	CAT	CCL2	CD274	CD276	CD28	CD40	CD40LG	CD80	CD86	CPA6
	CTLA4	CYBA	DDX10	DPP4	ENTPD1	FAHD2A	FAS	FASLG	FBXW8	FOXP3	GCLC	GCLM	GGT1
	GPX1	GPX3	GPX4	GSR	GSS	GSTA1	GSTA2	GSTA3	GSTA4	GSTA5	GSTK1	GSTM2	GSTM3
	GSTM4	GSTM5	GSTO1	GSTO2	GSTT2	GSTZ1	HAVCR2	HIF1A	HMOX1	HMOX2	HSPAIL	ICOS	ICOSLG
	IDO1	IDO2	IFNG	IFNGR1	IFNGR2	IL10	IL10RA	IL12A	IL12B	IL12RB1	IL12RB2	IL17A	IL17RA
	IL18	IL18R1	IL18RAP	IL1A	IL1B	IL1R1	IL4	IL4R	IL6	IL6R	KCNE3	KCNIP3	KEAP1
	KSR2	LAG3	LGALS9	MAFK	MIR4272	MPO	NFE2L2	NLRP3	NOS1	NOS2	NOS3	NT5E	PDCD1
	PDCD1LG2	PLXNA4	POLD3	PROM2	PSD3	SOD1	SOD3	SRXN1	STAT3	TGFB1	TGFBRI	THSD7B	TNFRSF4
	TNF	TNFAIP3	TNFRSF14	TNFRSF1A	TNFRSF1B	TNFRSF9	TNFSF10	TNFSF14	TNFSF4	TNFSF9	TRIM43	TXNRD1	USP44
	VTCN1	ZNF804B											



# READY-TO-USE PANELS FOR LIQUID BIOPSY

CELEMICS PRODUCTS & SERVICES 2022

Circulating Tumor DNA Panel  
: Colorectal / Breast / Lung



# Circulating Tumor DNA Panel

## Colorectal / Breast / Lung

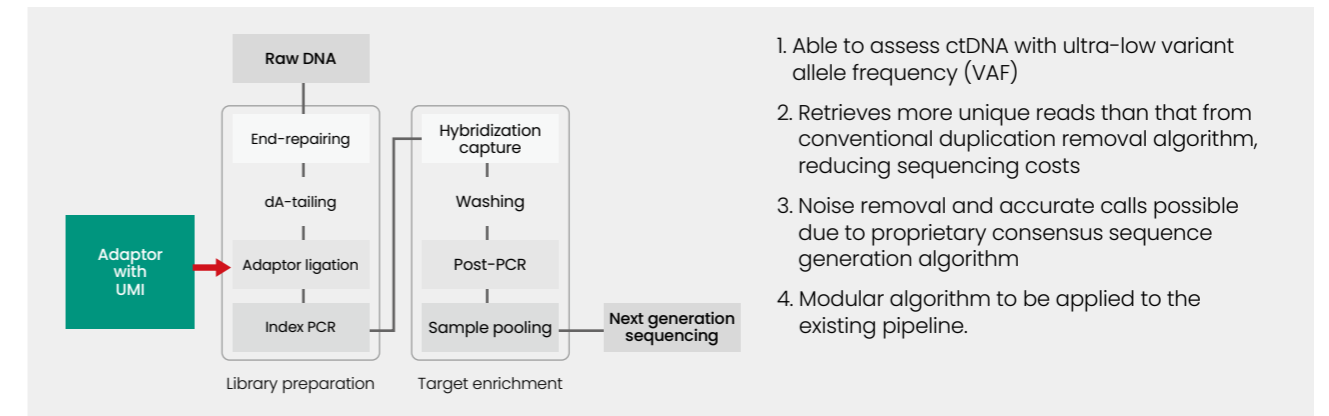
### OVERVIEW

The detection sensitivity for low-frequency variants from a limited amount of sample is of great importance to ctDNA analysis kits. Celemics has developed ctDNA kits for colon, breast, and lung cancer assay through collaborative research with Seoul National University Hospital (SNUH) since 2017. We have integrated our market-leading proprietary technologies including probe design algorithms, noise removal techniques, and reagents optimization. The panels are thoroughly validated and ready to use for clinical diagnosis.

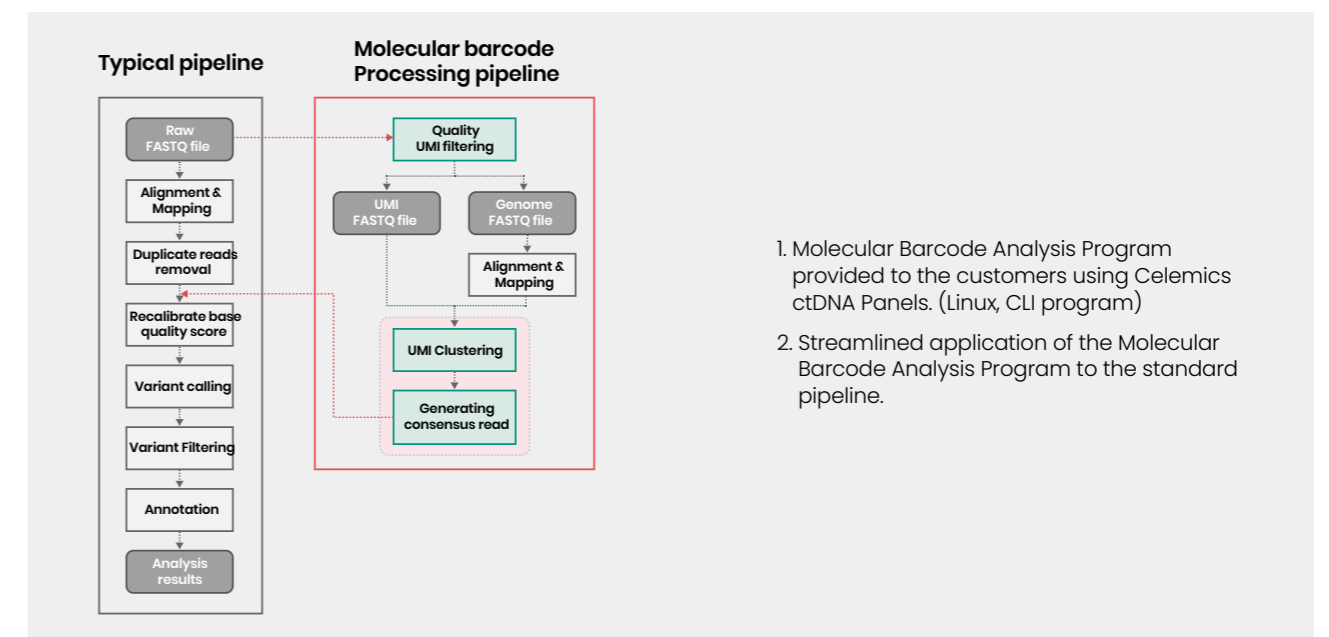
### KEY FEATURES

- |  |   |
|--|---|
| <p><b>1. Detects ctDNA for colorectal cancer, breast cancer, and lung cancer</b></p>     | <p>Assess 16 key genes for colorectal cancer, 27 for breast cancer, 28 for lung cancer</p>  |
| <p><b>2. Highly optimized panel for clinical testing with exceptional accuracy</b></p>   | <p>Complete validated panel performance conducted with patient samples through collaborative research with Seoul National University Hospital</p>                               |
| <p><b>3. Provides Unique Molecular Identifiers (UMI) and Bioinformatics Software</b></p> | <p>Receive high-quality data supported by Celemics proprietary UMI algorithms and analysis software, enabling efficient duplication removal and minimizing sequencing noise</p> |

### MODULAR UNIQUE MOLECULAR IDENTIFIER



### MODULAR BIOINFORMATICS PIPELINE



### PACKAGE COMPOSITION

Package name	Compositions		Package option	Options	
Target Enrichment	Target capture Probe	-	Pooling method	Single Reaction	Pre-capture Pooling
Standard	Target Enrichment reagents	Library prep Kit	Library Preparation kits	Standard Kit	EP-kit
All-In-One		Beads / Polymerase	Hybridization Enhancer	Included	Not included



# Circulating-tumor DNA Colorectal Cancer Panel

## SPECIFICATION

Gene count*	16 genes
Covered region	Whole CDS
Target size	18 kb
Mutation type	SNV, Indel
Sample type (amount)	Plasma (> 20 ng of cfDNA)
Platform	All sequencers from Illumina and MGI

**Bioinformatics pipeline**

1. Primary and Secondary analysis result (FASTQ to VCF)
2. Tertiary analysis result (VCF to Clinical report)
3. Linux-based consensus read generation software provided

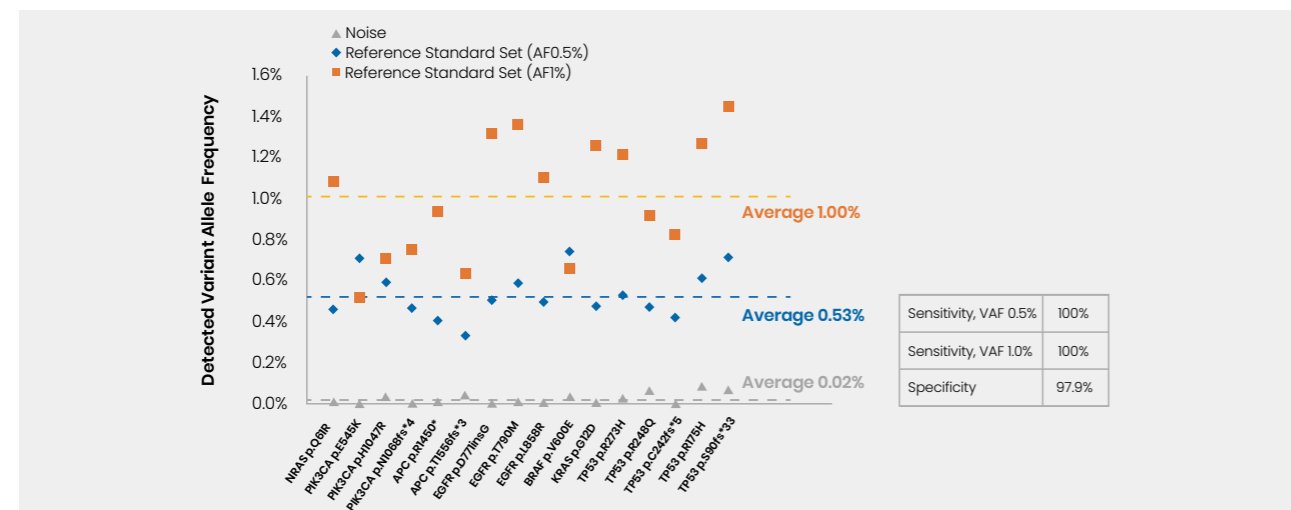
\* Gene Add-On Service: Genes can be added by customer's request

## GENE LIST

ctDNA Panel Colorectal Cancer	APC	BRAF	EGFR	ERBB2	ERBB3	FGFR1	HRAS	IRS1	KRAS	KRAS	MET	NRAS	PDGFRB
	PIK3CA	PTEN	TP53										

## PANEL PERFORMANCE

Detection of 16 variants with 100% sensitivity and 97.9% specificity at 0.5% VAF and 1% VAF



# Circulating-tumor DNA Breast Cancer Panel

## SPECIFICATION

Gene count*	27 genes
Covered region	Whole CDS
Target size	99 kb
Mutation type	SNV, Indel
Sample type (amount)	Plasma (> 20 ng of cfDNA)
Platform	All sequencers from Illumina and MGI

**Bioinformatics pipeline**

1. Primary and Secondary analysis result (FASTQ to VCF)
2. Tertiary analysis result (VCF to Clinical report)
3. Linux-based consensus read generation software provided

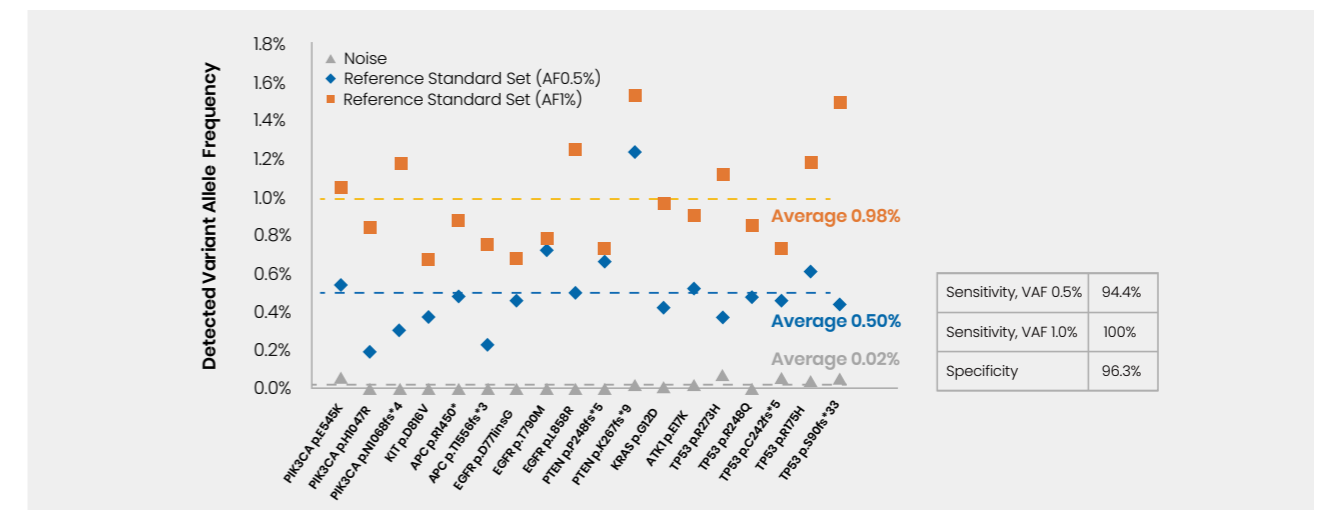
\* Gene Add-On Service: Genes can be added by customer's request

## GENE LIST

ctDNA Panel Breast Cancer	AKT1	APC	AR	BRCA1	BRCA2	CCND1	CDHI	EGFR	ERBB2	ESR1	FGFR1	FGFR2	GATA3
	IGF1R	KIT	KRAS	MAP2K4	MAP3K1	MDM2	MYC	NFI	PIK3CA	PIK3R1	PTEN	RBI	TOP2A
	TP53												

## PANEL PERFORMANCE

Detection of 27 variants with 96.3% specificity and 94.4% sensitivity at 0.5% VAF and 100% at 1% VAF



# Circulating-tumor DNA Lung Cancer Panel

## SPECIFICATION

Gene count*	28 genes
Covered region	Whole CDS
Target size	47 kb
Mutation type	SNV, Indel
Sample type (amount)	Plasma (> 20 ng of cfDNA)
Platform	All sequencers from Illumina and MGI

**Bioinformatics pipeline**

1. Primary and Secondary analysis result (FASTQ to VCF)
2. Tertiary analysis result (VCF to Clinical report)
3. Linux-based consensus read generation software provided

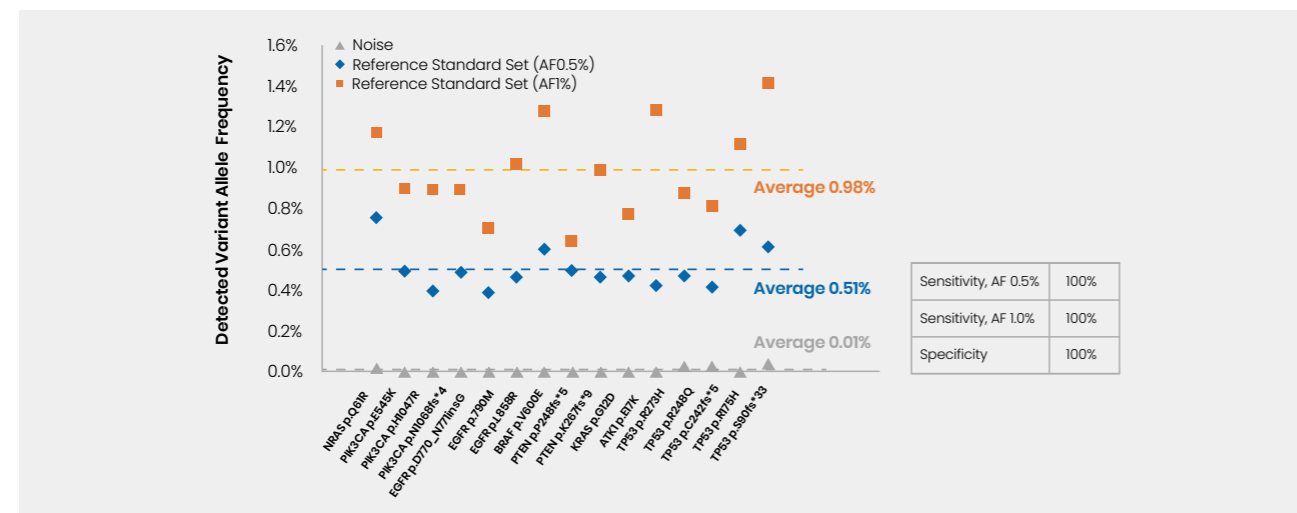
\* Gene Add-On Service: Genes can be added by customer's request

## GENE LIST

ctDNA Panel Lung Cancer	AKT1	ALK	ARAF	ARID1A	BRAF	CBL	CDKN2A	EGFR	ERBB2	HRAS	KEAP1	KRAS	MAP2K1
	MET	MTOR	NFI	NRAS	NTRK1	NTRK2	PIK3CA	PTEN	RBI	RIT1	ROS1	SETD2	STK11
	TP53	U2AF1											

## PANEL PERFORMANCE

Detection of 28 variants with 100% sensitivity and 100% specificity at 0.5% VAF and 1% VAF detection



# READY-TO-USE PANELS FOR MITOCHONDRIAL DNA

CELEMICS PRODUCTS & SERVICES 2022

Mitochondrial DNA Sequencing Panel





# Mitochondrial DNA Sequencing Panel

Mitochondrial Diseases  
: Metabolic and neurological disorders and cancers

## DESCRIPTION

Celemics has specifically designed capture probes and adjusted the concentration of the panel for each respective use with our own proprietary rebalancing technologies to provide complete, consistent coverage of the whole mitochondrial genome while taking into consideration small target regions. This enables the same high level of target capture efficiency regardless of small target sizes even with a stand-alone panel.

## KEY FEATURES

1. High-fidelity sequencing	Guarantees maximum capture efficiency in custom panels without affecting target specificity
2. Highly uniform coverage and mean depth	High coverage and uniformity across the entire human mitochondrial genome
3. Flexible customization	Convenient addition to other Celeomics target enrichment panels such as G-Mendeliome panels for further mtDNA-derived rare disease analysis

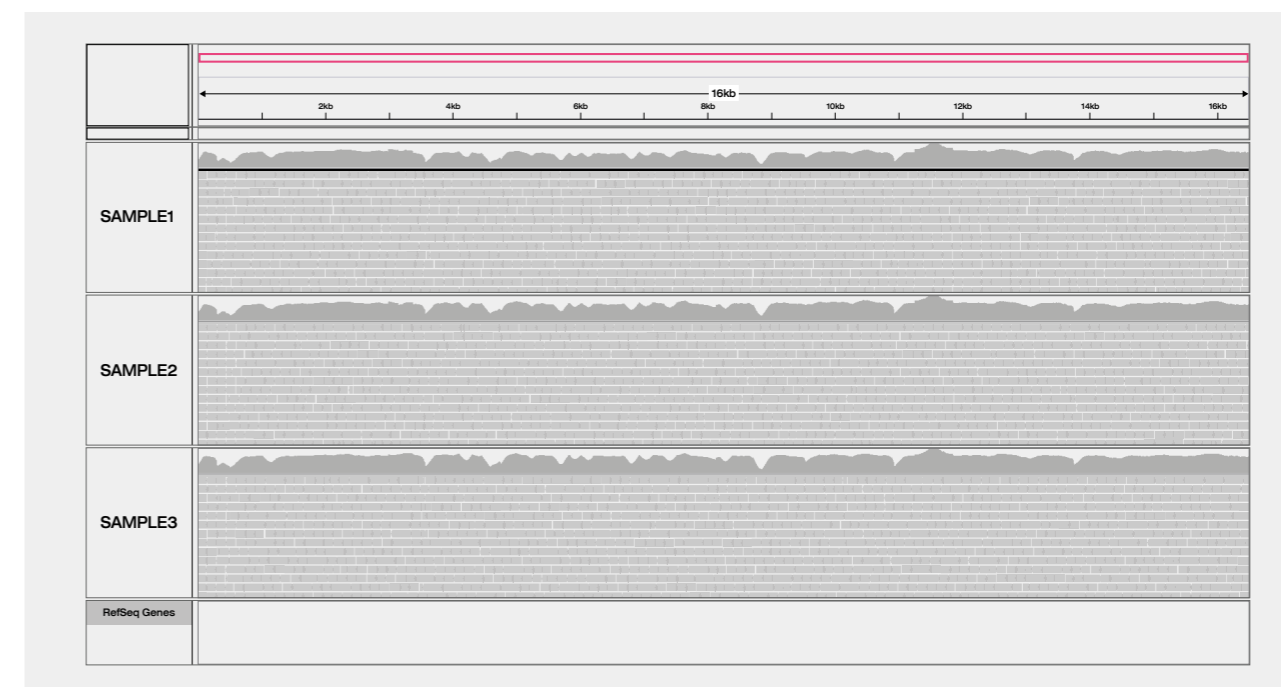
## SPECIFICATION

Covered region*	Whole mitochondrial genome
Target size	16.6 kb
Mutation type	SNV, Indel
Sample type (amount)	Blood (> 50 ng of fragmented DNA)
Platform	All sequencers from Illumina, Thermo Fisher, MGI, PacBio, and Oxford Nanopore
Bioinformatics pipeline	Primary, Secondary and Tertiary analysis result (FASTQ to VCF, VCF to Clinical report)

## PERFORMANCE

NGS Sequencing Amount	On-Target Base Ratio	Mean Depth	Coverage		
			10x	50x	100x
10Mb	97.93%	493x	99.98%	99.91%	99.87%

## IGV EXAMPLE OF CELEMICS mtDNA SEQUENCING PANEL



Celemics mtDNA Sequencing Panel shows 99% coverage with uniformity

## PACKAGE COMPOSITION

Package name	Compositions		Package option	Options	
Target Enrichment	Target capture Probe	-	Pooling method	Single Reaction	Pre-capture Pooling
Standard	Target Enrichment reagents	Library prep Kit	Library Preparation Kits	Standard Kit	EP-kit
All-In-One		Beads / Polymerase	Hybridization Enhancer	Included	Not included



# TARGET ENRICHMENT KITS FOR RNA SEQUENCING

CELEMICS PRODUCTS & SERVICES 2022

Targeted RNA Sequencing Panel



# Targeted RNA Sequencing Panel

Transcriptome Analysis

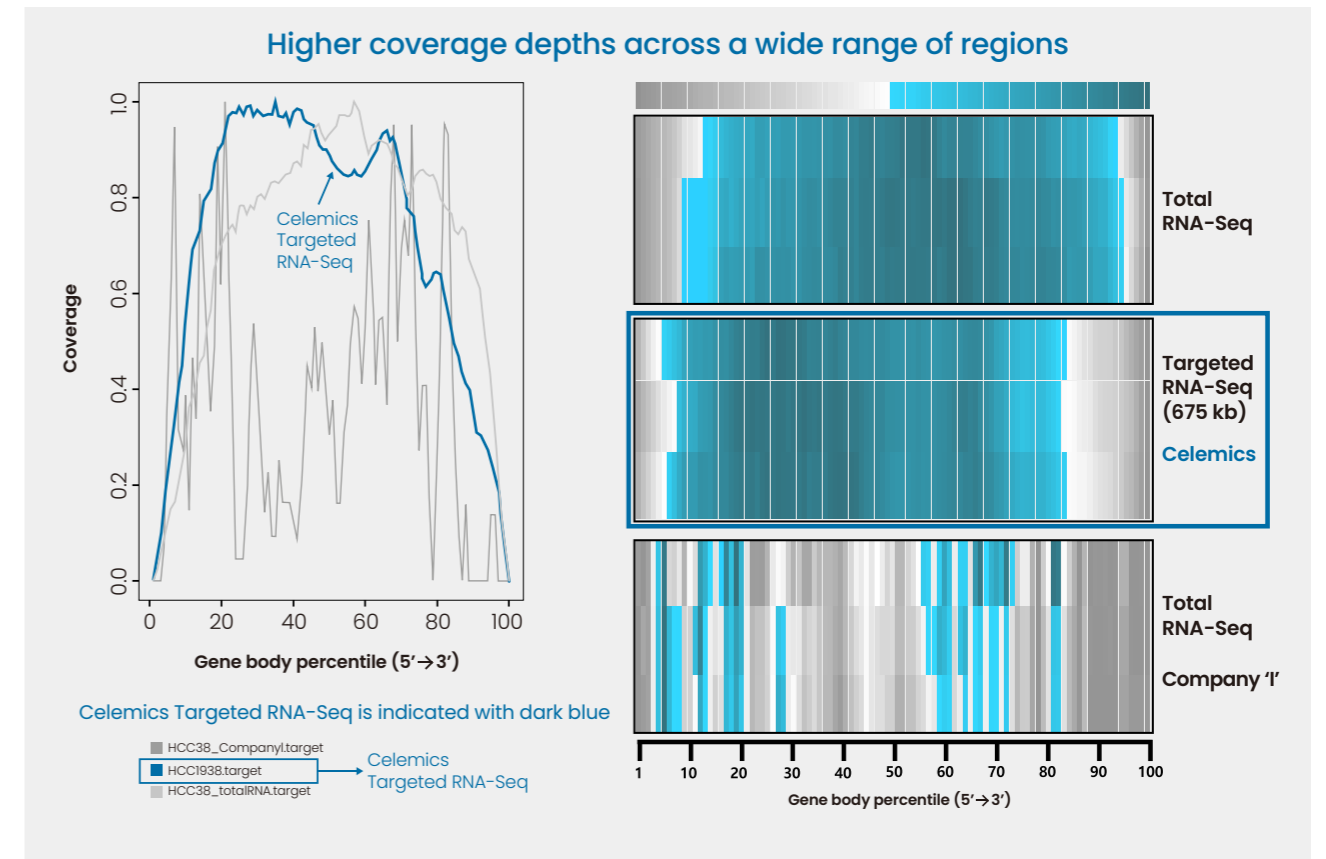
## KEY FEATURES

1. Cost-effective high-quality analysis	Accurate analysis of expression levels enabled by higher depth of coverage due to specific targeting of genes of interest, compared to total RNA sequencing
2. Compatible with a variety of sample types	Receive reliable results from poor-quality samples such as FFPE and low-amount samples such as cfRNA
3. Expression level in all regions of genes of interest	Covers all gene regions, allowing for the assessment of expression levels across all exons
4. Gene rearrangement analysis	Detects rearrangement and all other types of variants
5. Isoform analysis	Identify isoform expression levels by assessing the entire regions of targeted genes.

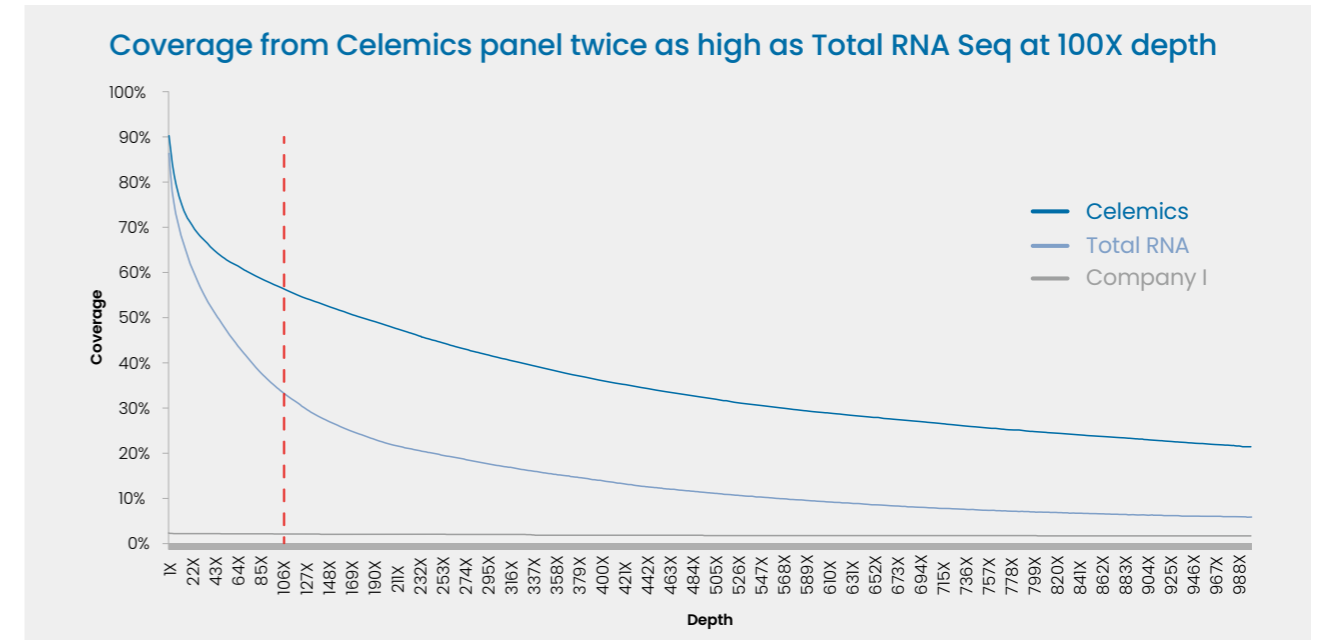
## PACKAGE COMPOSITION

Package name	Compositions	Package option	Options
Target Enrichment	Target capture Probe	Pooling method	Single Reaction    Pre-capture Pooling
Standard	Target Enrichment reagents	Library Preparation kits	Standard Kit
All-in-One	Library prep Kit Beads / Polymerase	Hybridization Enhancer	Included    Not included

## PANEL PERFORMANCE



Celeomics Targeted RNA Sequencing assesses the expression level of selective genes with sufficient level of coverage depth that is higher than that of total mRNA sequencing. Compared to competitor products that targets only parts of an exon, the Targeted RNA Sequencing developed by Celeomics showed relatively higher coverage across a wide range of regions.



The comparison test between Celeomics Targeted RNA Sequencing and total RNA sequencing shows that the coverage from the Celeomics product is 15% higher at 50X and twice as high at 100X.

# TARGET ENRICHMENT KITS FOR EPIGENETICS

CELEMICS PRODUCTS & SERVICES 2022

Targeted Methylation Sequencing Panel



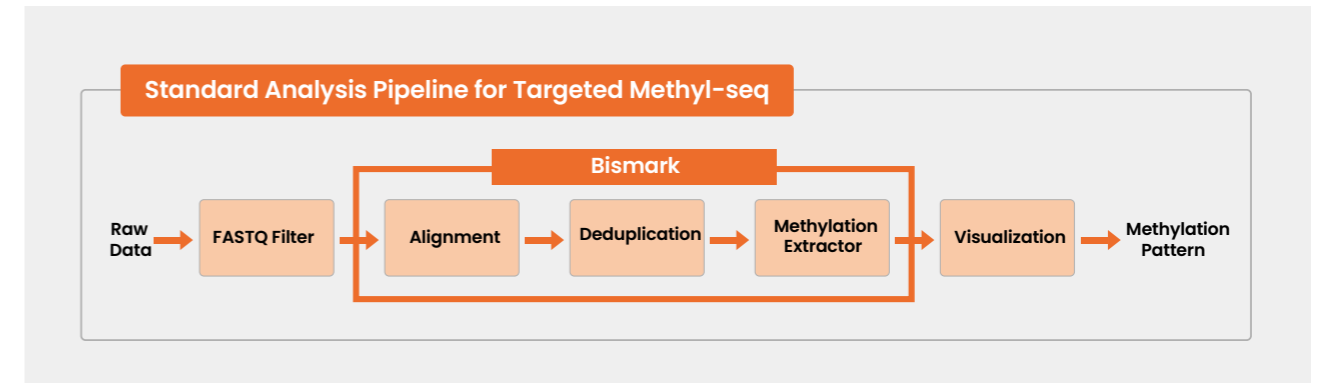
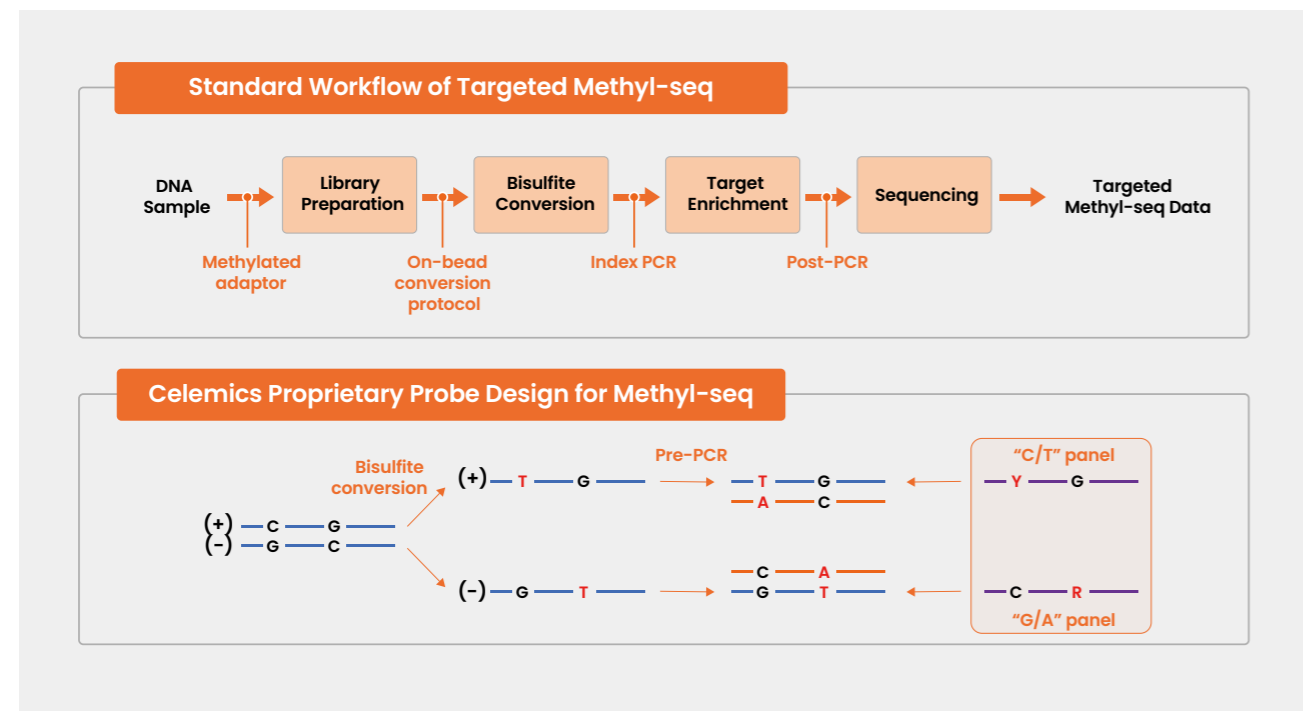
# Targeted Methylation Sequencing Panel

Epigenetics

## KEY FEATURES

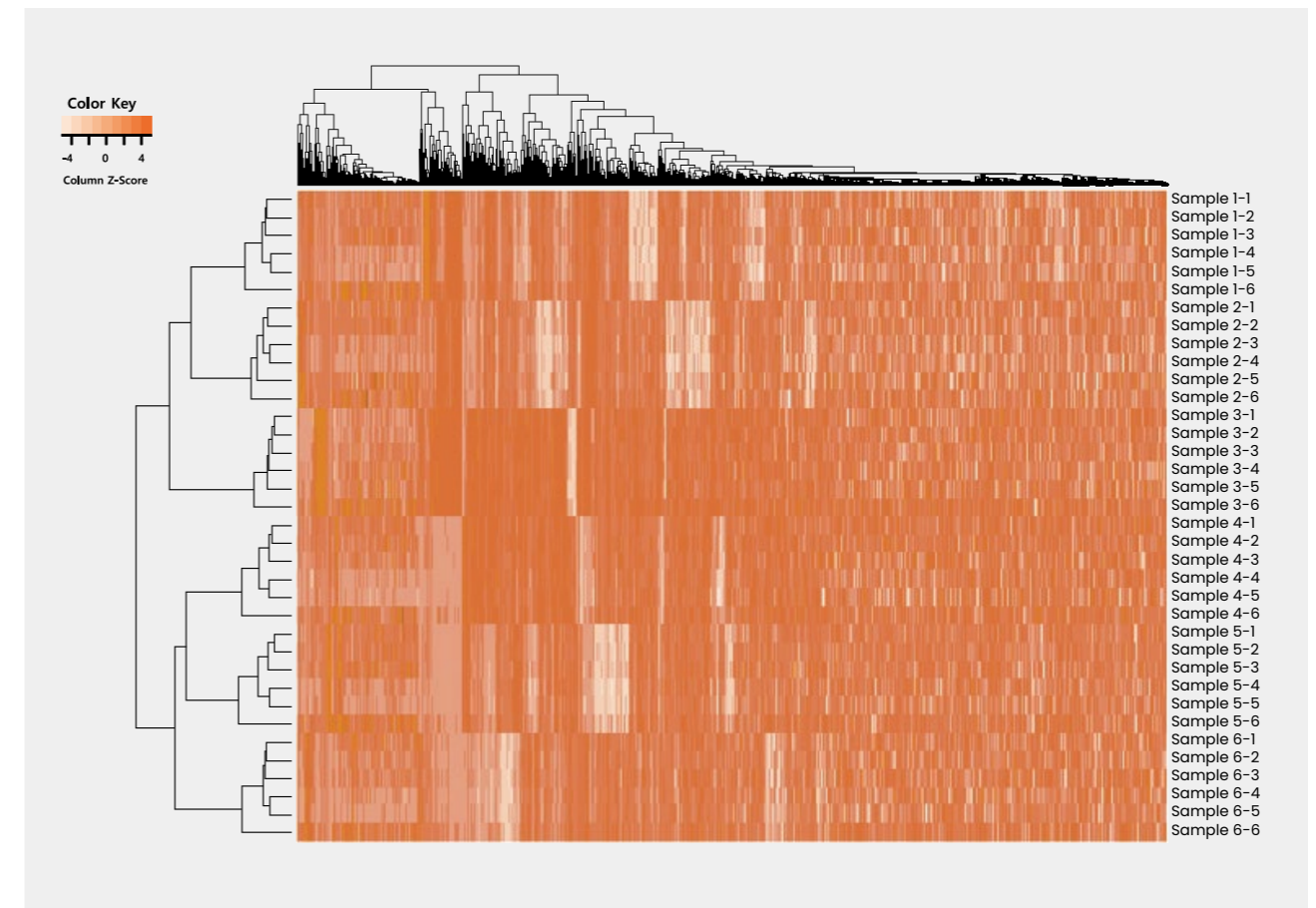
- |   |   |
|---|---|
| 1. Probe specifically designed for Methyl-seq | Elaborate design considering the sequence alteration by bisulfite conversion<br>Thorough comparison analysis of the sequences before and after bisulfite conversion, enabling accurate detection of methylation sites |
| 2. Compatible with all sample types           | Perform methylation analysis with gDNA and cfDNA  |

## PANEL PERFORMANCE



The Targeted Methylation Sequencing is proceeded with including a bisulfite conversion process in the NGS workflow. The hybridization probe and methylated adaptors are designed considering the sequence alteration by bisulfite conversion, enabling an accurate comparison analysis of the sequences before and after the conversion. Selective genes are targeted for the analysis, allowing for cost-effective sequencing.

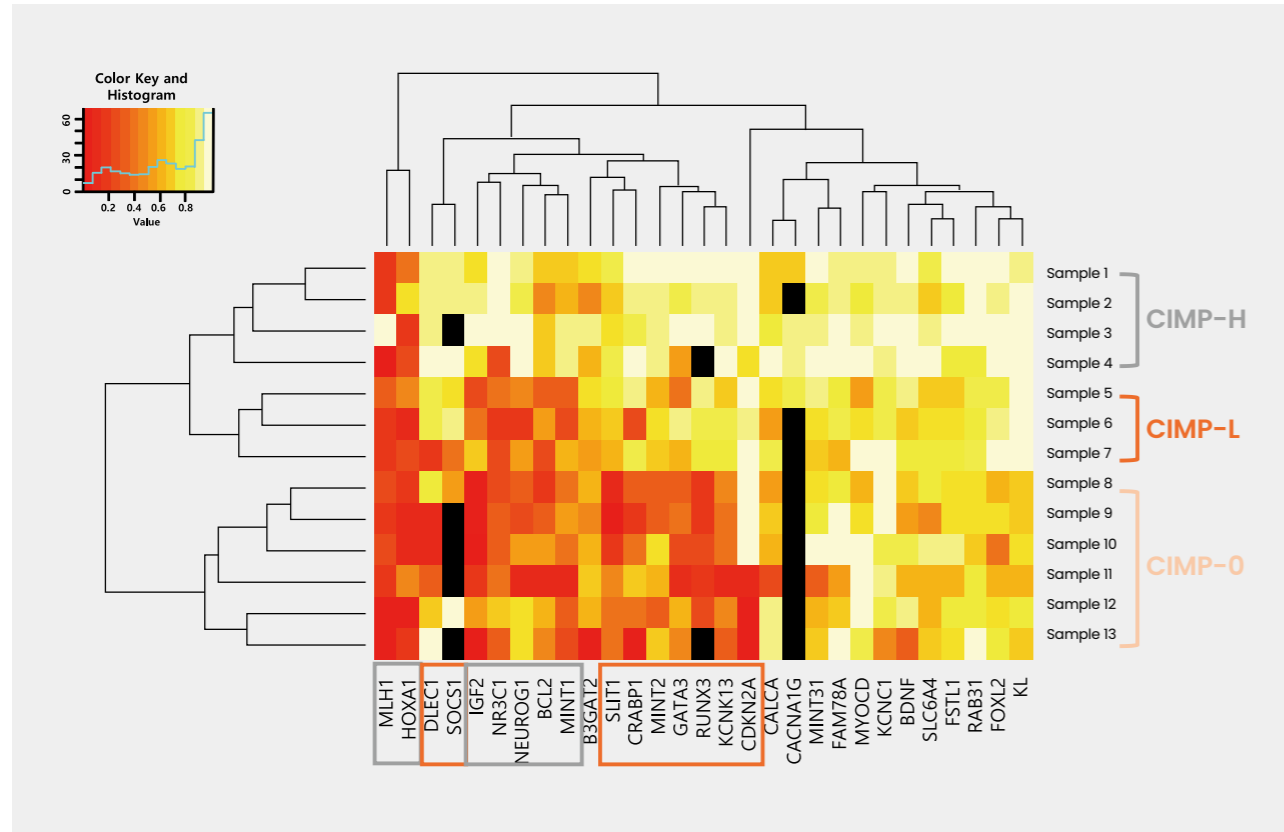
## HIGH REPRODUCIBILITY OF METHYLATION PATTERN ANALYSIS



The results demonstrate high reproducibility of the analysis, yielding the same methylation patterns when repeatedly tested with the identical specimens.

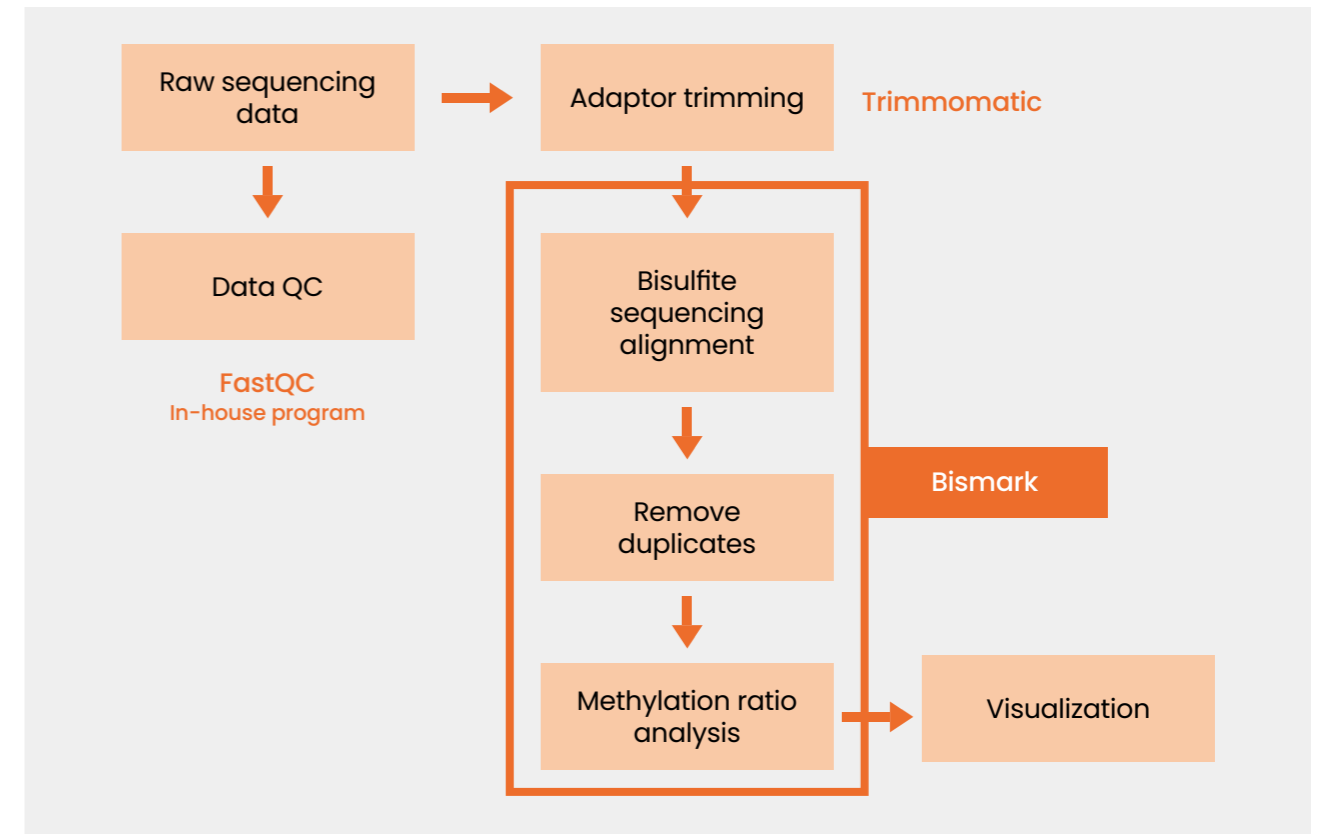


## HIGH CONCORDANCE OF METHYLATION PATTERN ANALYSIS WITH CLINICAL INFORMATION



The clustering result from pattern analysis showed high concordance with the clinical data information.

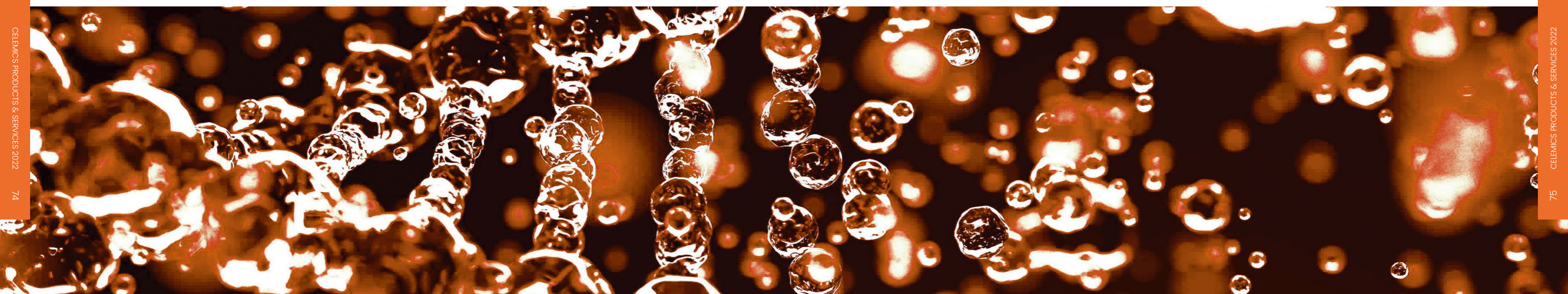
## WORKFLOW OF TARGETED METHYLATION SEQUENCING ANALYSIS



Customers who are new to methylation analysis are supported by Celemics bioinformatics software service for fast and accurate analysis.

## PACKAGE COMPOSITION

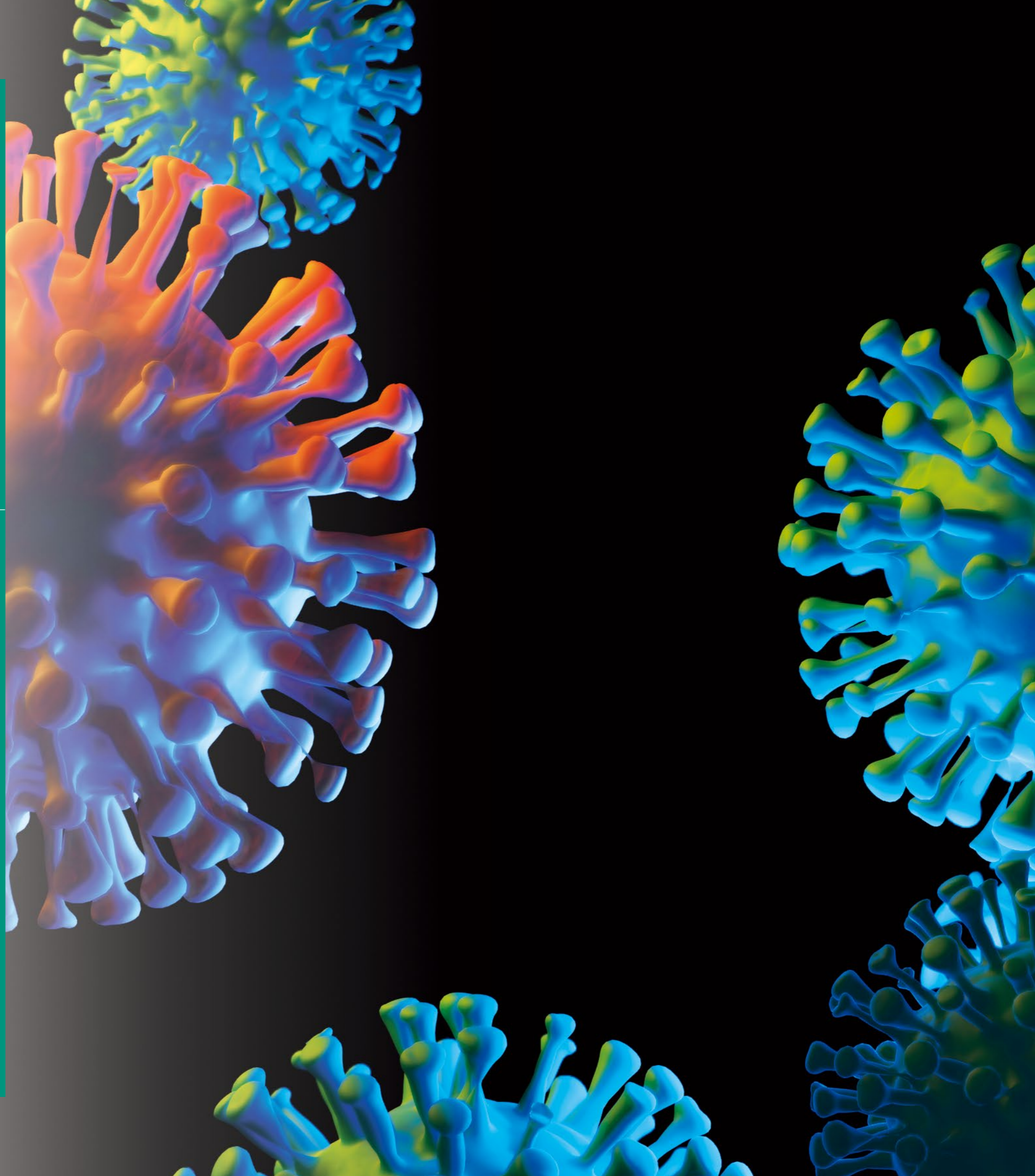
Package name	Compositions		Package option	Options	
Target Enrichment	Target capture Probe	-	Pooling method	Single Reaction	Pre-capture Pooling
Standard	Target Enrichment reagents	Library prep Kit	Library Preparation kits	Standard Kit	EP-kit
All-in-One		Beads / Polymerase	Hybridization Enhancer	Included	Not included



# TARGET ENRICHMENT KITS FOR VIRUS RESEARCH

CELEMICS PRODUCTS & SERVICES 2022

Comprehensive Respiratory Virus Panel  
African Swine Fever Virus Panel



# Comprehensive Respiratory Virus Panel

Research and Diagnosis of Respiratory viruses

## DESCRIPTION

The CRV Panel is designed for the comprehensive analysis of clinically significant respiratory viruses that are widely assessed by medical institutions around the globe. The panel validation test with clinical samples showed superior whole genome sequencing (WGS) success rates compared to other competitor kits on the market. The panel tests for multiple infections by assessing all types of respiratory viruses including SARS-CoV-2. The panel includes all required kits including the RNA-to-cDNA Kit and cDNA-to-Captured Library Kit. The hybridization enhancer technology is used for rapid one-day workflow. Our customers can receive stand-alone bioinformatics software, 'Celemics Virus Verifier', which provides in-depth analysis information while ensuring the security of client sequence information.

## KEY FEATURES

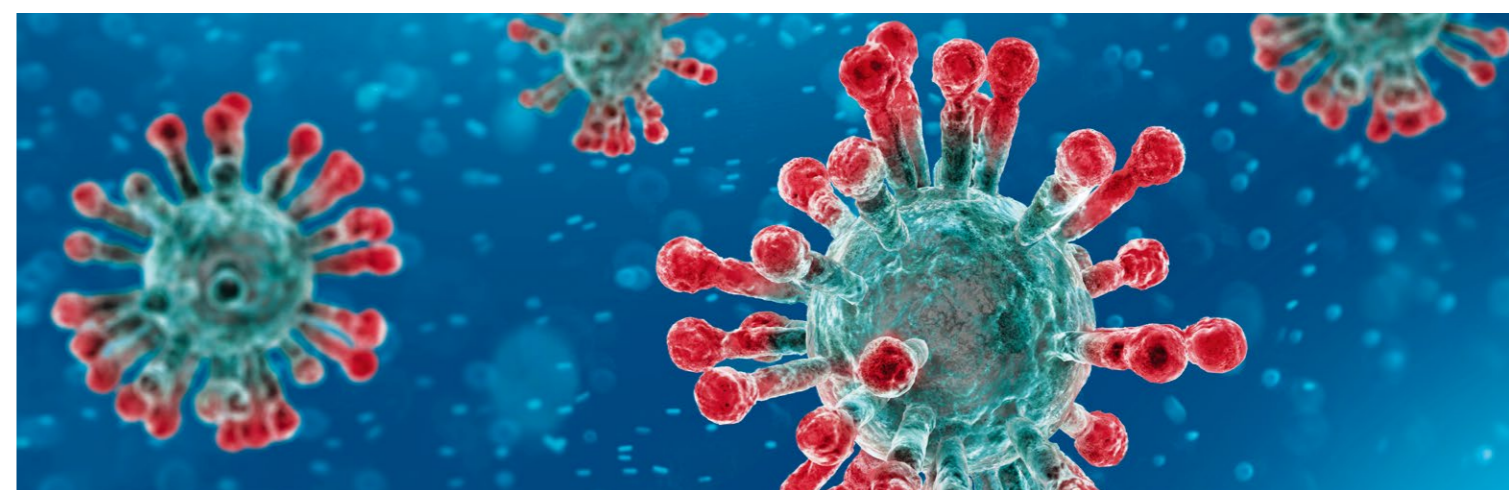
<b>1. Coverage of wide range of respiratory pathogens</b>	Assess WGS of 39 variants for 9 different virus types (SARS-CoV-2 solo analysis also available) Includes all types of respiratory viruses that are assessed by medical institutions around the globe
<b>2. Superior WGS success rate even with poor quality specimen</b>	Able to detect pathogens from patient specimens as well as poor quality environmental specimens Exceptional success rate of variant detection and WGS Significantly reduced gap formation
<b>3. Double pandemic / coinfection detection</b>	Detect all relevant viral strains in a single assay and test for multiple infections
<b>4. Inclusion of Celemics Virus Verifier or bioinformatics analysis</b>	Receive stand-alone bioinformatics SW Protect your easily-compromised data with our EU-GDPR compliant cloud system

## SPECIFICATION

<b>Target viruses*</b>	9 types / 39 virus strains, including SARS-CoV-2
<b>Target size</b>	706 kb
<b>Mutation type</b>	Viral variants detection, Viral mutation (SNV, Indel) from generated Whole Genome Sequence
<b>Sample type</b>	Upper respiratory tract, Nasopharyngeal, Oropharyngeal specimens, and others
<b>Platform</b>	All sequencers from Illumina and Thermo Fisher
<b>Kit composition</b>	Provides all required reagents, including RNA to cDNA kit, cDNA to captured library kit, and bioinformatics software
<b>Bioinformatics pipeline</b>	Provides stand-alone bioinformatics software 'Celemics Virus Verifier' (FASTQ to Report)
<b>Related publication</b>	Evidence of long-distance droplet transmission of SARS-CoV-2 by direct air flow in a restaurant in Korea, J Korean Med Sci. (2020)

## PATHOGEN LIST

Human Adenovirus	Coronavirus	Parainfluenza Virus	Respiratory Syncytial Virus
Human Adenovirus Type 1 (HAdV-C1)	Coronavirus HKU1	Parainfluenza 1 (PIV 1)	Respiratory Syncytial Virus A (RSV A)
Human Adenovirus Type 2 (HAdV-C2)	Coronavirus NL63	Parainfluenza 2 (PIV 2)	Respiratory Syncytial Virus B (RSV B)
Human Adenovirus Type 3 (HAdV-B3)	Coronavirus 229E	Parainfluenza 3 (PIV 3)	Human Metapneumovirus
Human Adenovirus Type 4 (HAdV-E4)	Coronavirus OC43	Parainfluenza 4 (PIV 4) A	
Human Adenovirus Type 5 (HAdV-C5)	SARS-CoV-2	Parainfluenza 4 (PIV 4) B	
Human Adenovirus 7 (HAdV-B7)			
Human Adenovirus 14 (HAdV-B14)			
Human Adenovirus 21 (HAdV-B21)			
	<b>Influenza A</b>		
	Influenza A Virus (Flu A)	EV-C104	Human Rhinovirus A
		EV-C105	Human Rhinovirus B
<b>Bocavirus 1/2/3/4 (HBoV)</b>	Influenza A-H1 Virus (Flu A-H1)	EV-C109	Human Rhinovirus C
Human Bocavirus 1	Influenza A-H3 Virus (Flu A-H3)	EV-C117	
Human Bocavirus 2		EV-C118	
Human Bocavirus 3	<b>Influenza B</b>	CV-A21	
Human Bocavirus 4	Influenza B Virus (Flu B)	EV-D68	
			<b>Human Enterovirus</b>
			<b>Human Rhinovirus (A/B/C)</b>



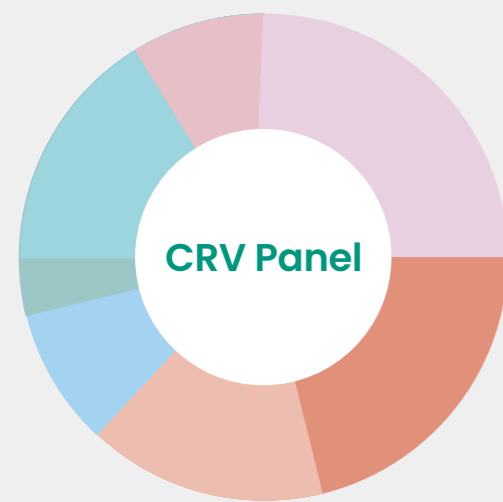


## PERFORMANCE

### High coverage of whole genome from reference samples using CRV Panel

Sample Type	Coverage [1X]	Coverage [10X]	Coverage [100X]
Reference sample (Illumina 2x75 bp)	99.95%	99.87%	98.95%

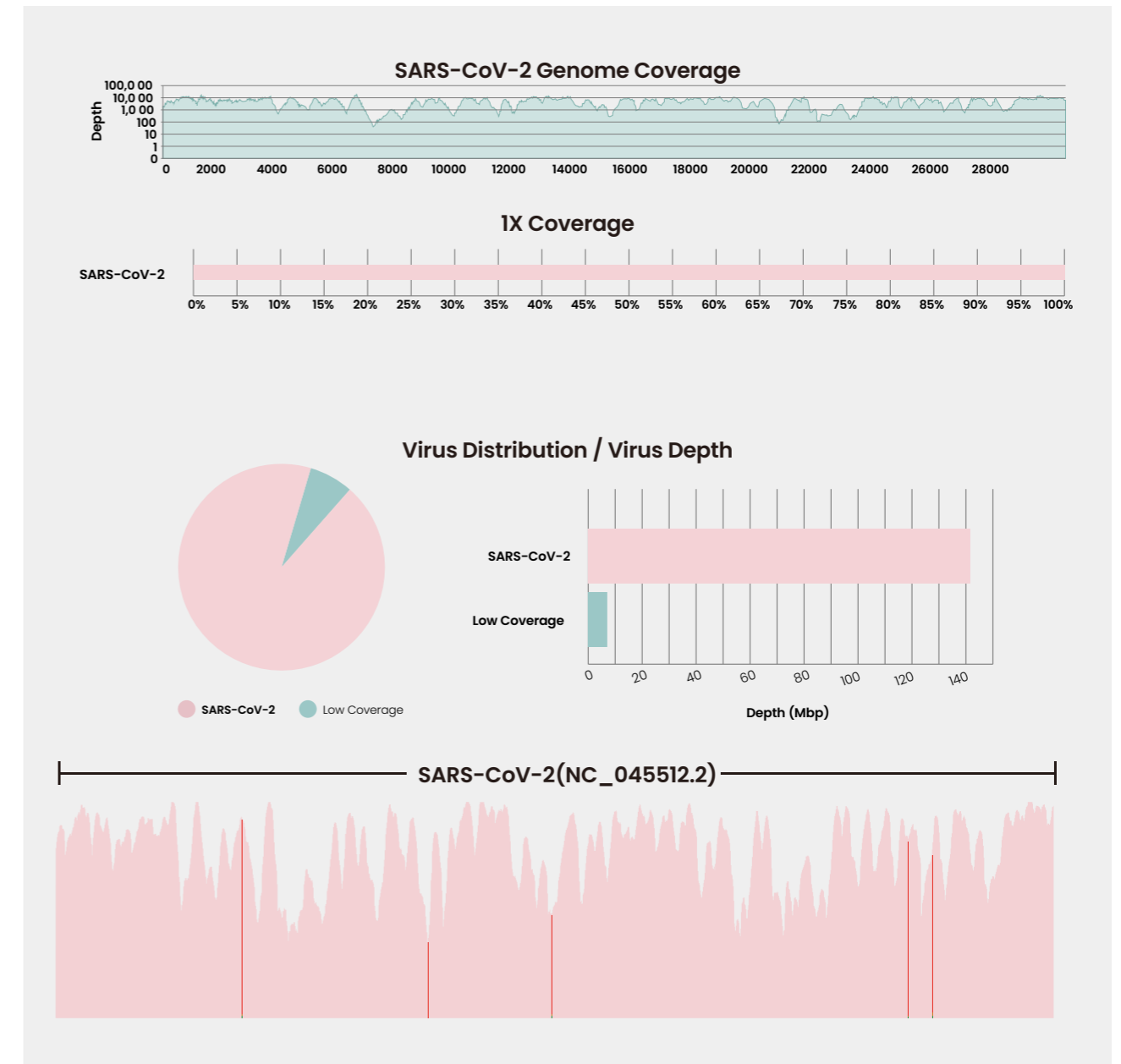
### Example of viral strain identification in reference sample



- Human\_Enterovirus\_EV-D68
- Influenza\_virus\_A-H1\_H1N1
- Influenza\_virus\_A-H3\_H3N2
- Low Coverage
- Influenza\_virus\_B
- SARS-CoV-2
- Parainfluenza\_virus\_4A

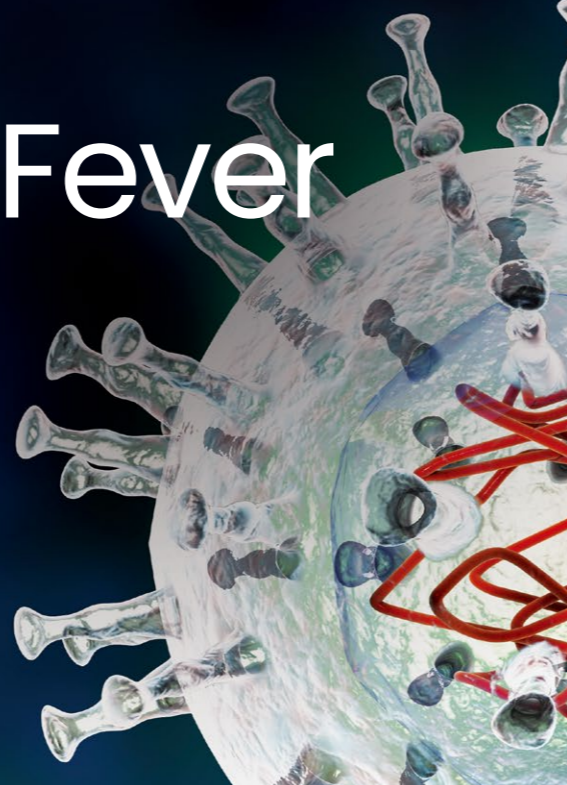
## CRV PANEL RESULTS GENERATED THROUGH CELEMICS VIRUS VERIFIER (STAND-ALONE SOFTWARE)

Celemics provides stand-alone software for bioinformatics analysis, allowing customers to access the detailed data analysis information and ensuring the security of client sequence information.



# African Swine Fever Virus Panel

Virus Research, Virus WGS Analysis



## DESCRIPTION

The high morbidity and mortality of African swine fever (ASF) has a severe impact on the global swine industry. However, currently there is no effective treatments or vaccines commercially available. The ASFV panel is designed to identify 26 strains of genotype II virus in a single NGS run. The panel can be utilized for identifying the cause and infection route.

## KEY FEATURES

1. Swine-specific blocking reagent	Provides swine-specific blocking reagent that effectively blocks repetitive sequences and allows for selectively retrieving the ASFV sequence
2. Comprehensive analysis of ASFV subtypes	Detect genotype II virus subtypes with specifically designed probes
3. Convenient testing	Highly accurate results from blood samples, often considered more challenging due to the lower viral load compared to concentrated culture supernatant or spleen tissue sample

## SPECIFICATION

Target viruses*	ASFV 26 strains
Target size	192 kb
Mutation type	Virus detection, Virus genome assembly
Sample type (amount)	Swine blood (50 ng of fragmented DNA)
Platform	All sequencers from Illumina, Thermo Fisher, MGI, PacBio, and Oxford Nanopore
Bioinformatics pipeline	Celemics ASFV Pipeline (FASTQ to Result)

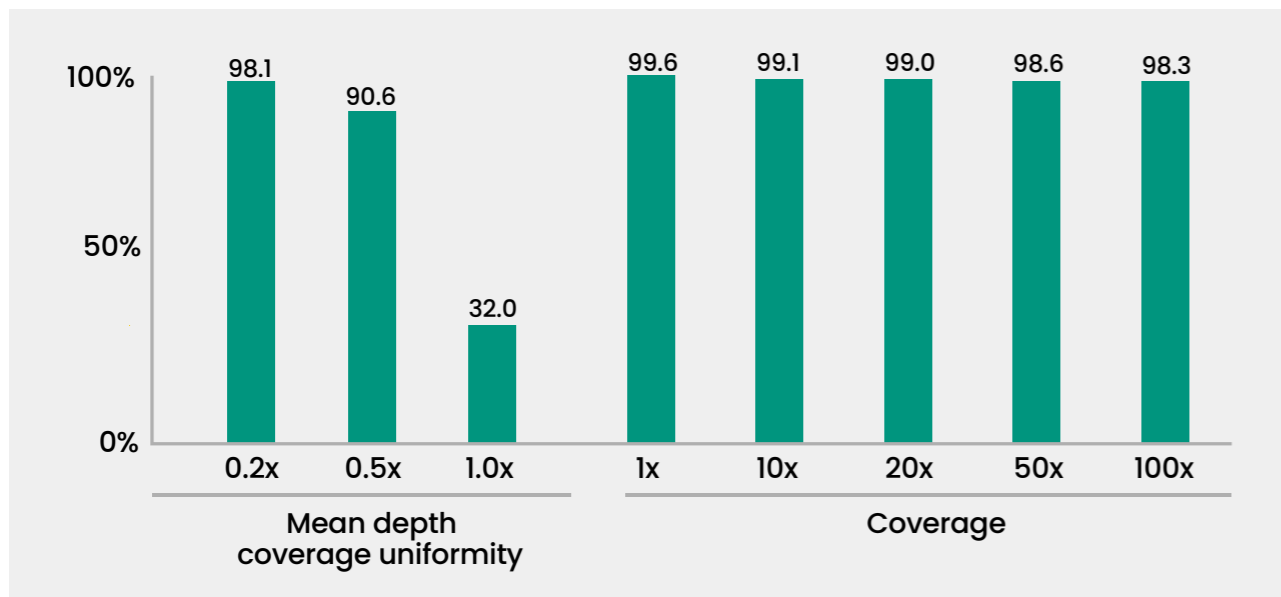
\* Gene Add-On Service: Genes can be added by customer's request

## ASFV DETECTION WORKFLOW

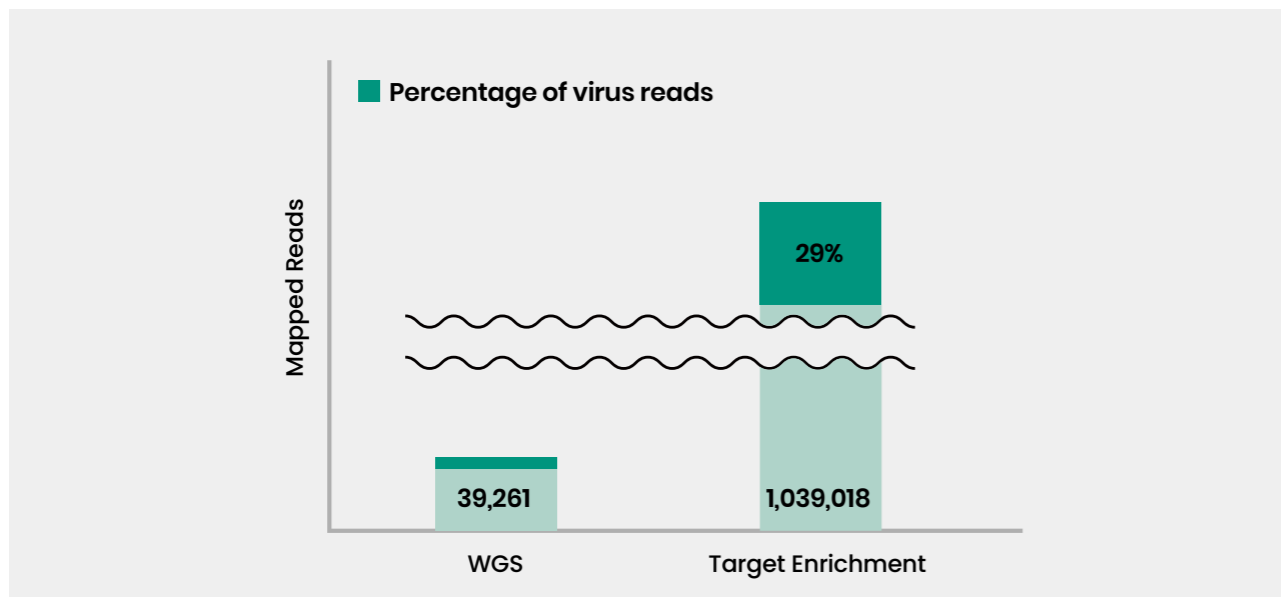


## PERFORMANCE

Advanced target enrichment technology enabling exceptional capture performance with high coverage and uniformity



The panel validation result shows high uniformity and high coverage at all levels.



With the same sequencing amount, target enrichment NGS yielded 29% virus reads out of a total of 1,039,018 reads, while whole genome sequencing (WGS) yielded 0.5% virus reads (green) out of a total of 39,261 reads.



# TARGET ENRICHMENT KITS FOR AGRICULTURE & ANIMAL RESEARCH

CELEMICS PRODUCTS & SERVICES 2022

Customized High-Throughput Genotyping Panel



# Customized High-Throughput Genotyping Panel

Plant and animal research

## DESCRIPTION

For molecular breeding, the availability and easy accessibility of genomic resources is a prerequisite. Although technological advances have provided a range of resources like molecular markers, genetic linkage maps, whole genome sequences and transcriptomes, agricultural genomics has faced many challenges. Celeemics provides a solution with the High-Throughput Genotyping Panel. We have utilized NGS methods, whereby a high number of regions of interest are simultaneously enriched using specifically designed probes to provide new insights into different agricultural genomics research.

## KEY FEATURES

<b>1. NGS-based target enrichment sequencing assay</b>	Utilize NGS-based target enrichment methods for higher accuracy and cost-effectiveness compared to conventional methods such as conventional GBS, PCR, and microarray
<b>2. Comprehensive analysis with high accuracy</b>	Perform comprehensive assay of 100 to 10,000 markers with minimized false-negatives and false-positives Discover novel SNPs
<b>3. Cost-effective analysis</b>	Benefit from Celeemics' library preparation kits, target capture technology, and multiplexing indices specifically designed for high-throughput genotyping
<b>4. Outstanding performance regardless of various origins</b>	Receive high-quality results enabled by species-specifically designed blocking oligos across all types of origins

## PACKAGE COMPOSITION

Package name	Compositions			Package option	Options	
Target Enrichment	Target capture Probe	-		Pooling method	Single Reaction	Pre-capture Pooling
Standard	Target Enrichment reagents	Library prep Kit	-	Library Preparation kits	Standard Kit	EP-kit
All-In-One		Beads / Polymerase		Hybridization Enhancer	Included	Not included

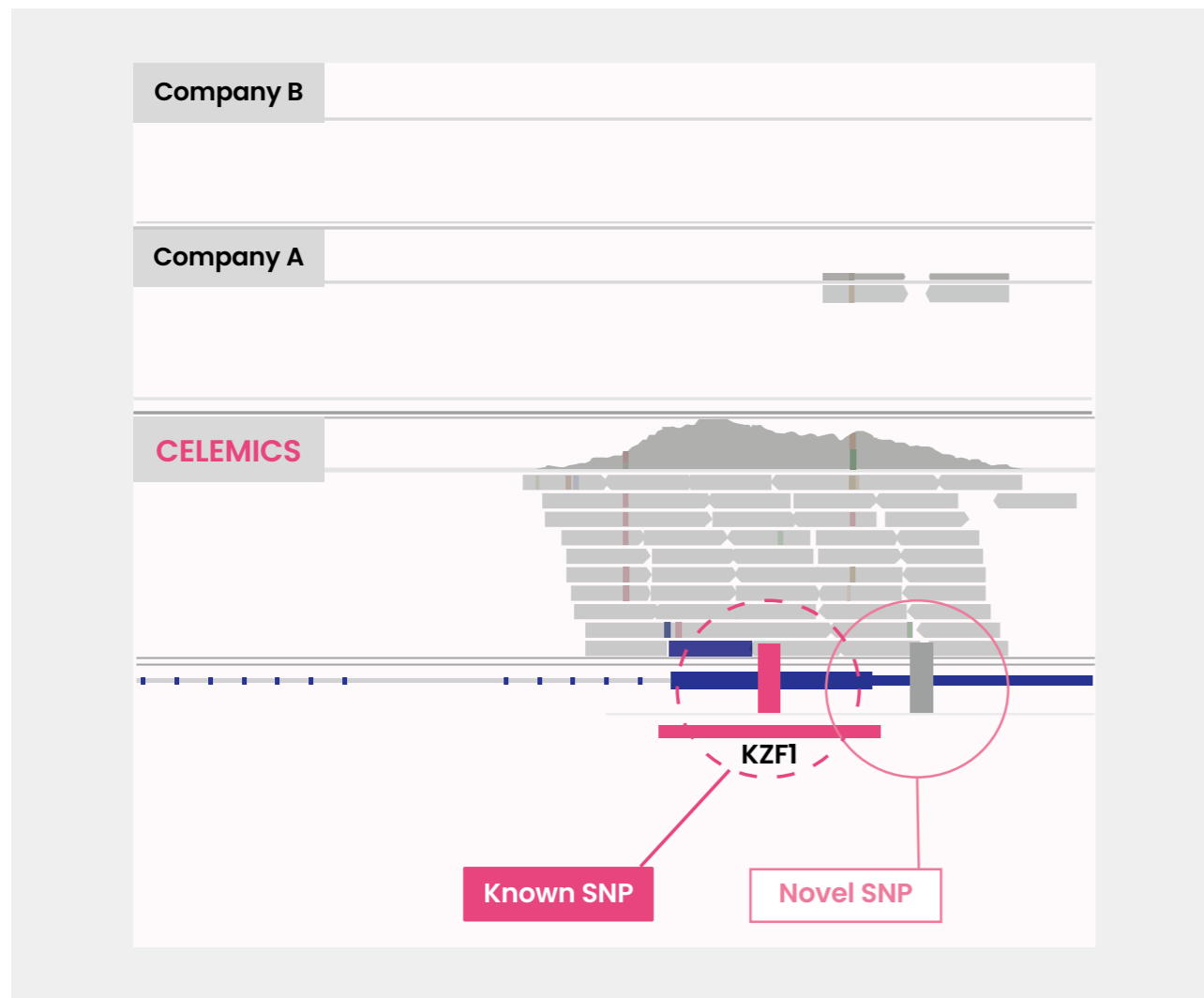
## COMPARISON WITH CONVENTIONAL TECHNOLOGIES

	Advantages	Disadvantages
<b>Conventional GBS</b>	1. Sequencing of multiple samples due to lower amount of data required compared to WGS	1. Limited biomarkers available due to limited conserved regions, reducing overall resolution 2. Unable to detect SNPs in the restriction sites
<b>Microarray</b>	1. Higher reproducibility than conventional GBS	1. Hard to customize new targets (novel biomarkers) 2. Low flexibility to meet various kinds of genotyping
<b>PCR</b>	1. Cost-effective for low number of samples 2. Easy and fast analysis	1. Limited number of biomarkers to analyze at once 2. Inappropriate for mass-analysis of biomarkers
<b>Celeemics Target Enrichment</b>	<p><b>1. Cost saving</b> : Highly cost-effective when assessing multiple samples</p> <p><b>2. Flexible customization</b> : Novel biomarkers can be added or removed</p> <p><b>3. Comprehensive analysis</b> : Including novel SNP discovery</p> <p><b>4. Exceptional performance</b> : Celeemics proprietary blocking oligo design technology</p> <p><b>5. Wide compatibility</b> : Compatible with a wide range of sample types</p>	



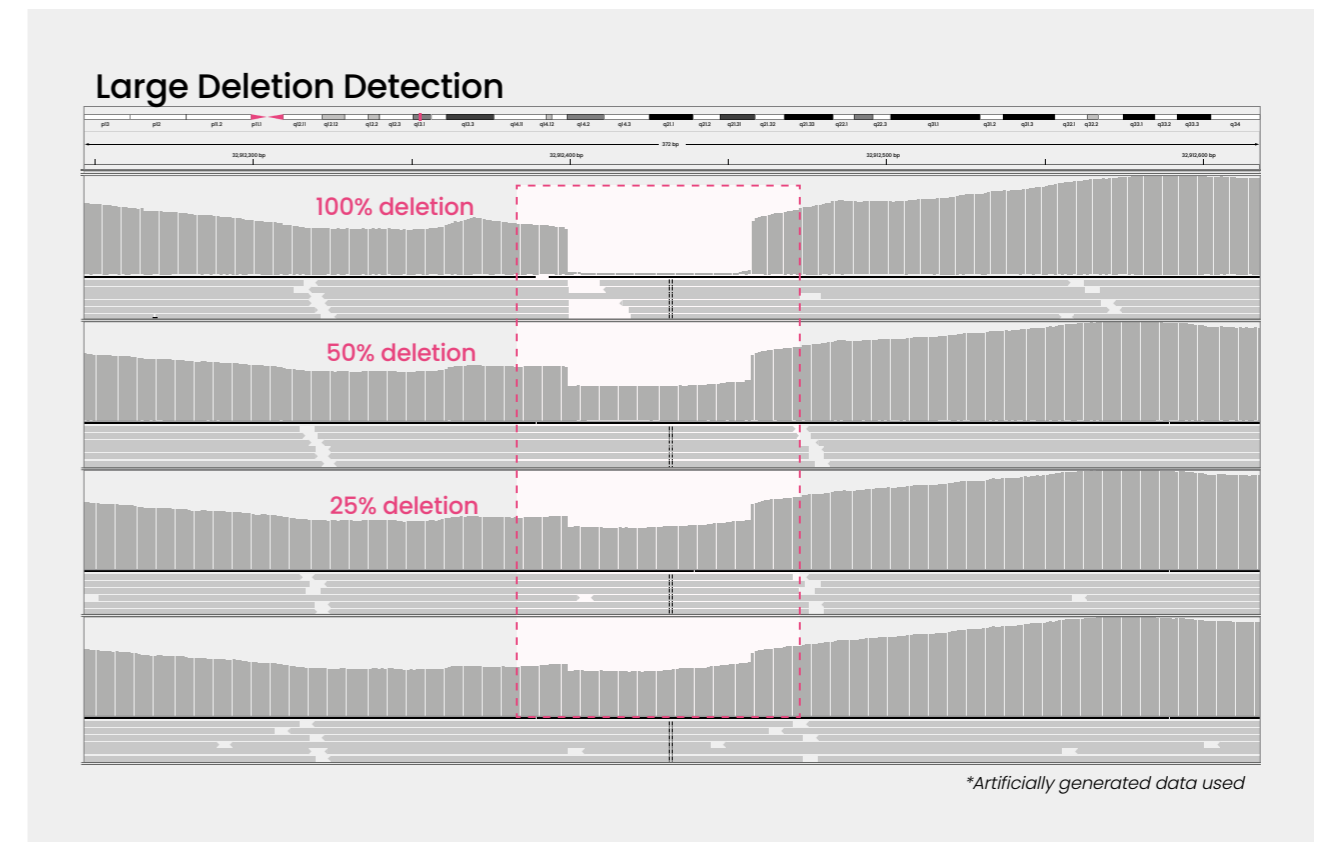
## PERFORMANCE

Hybridization-based NGS target enrichment enables discovery of novel SNPs near target regions



## PERFORMANCE

Hybridization-based NGS target enrichment enables accurate analysis of all mutation types including large deletion and rearrangement.



# CELEMICS SOLUTIONS FOR METAGENOMIC SEQUENCING

CELEMICS PRODUCTS & SERVICES 2022

Metagenomic Sequencing Service and Kit



# Metagenomic Sequencing Service and Kit

16S V4 / 16S V3-V4 / 18S ITS1 / 18S ITS1-ITS2

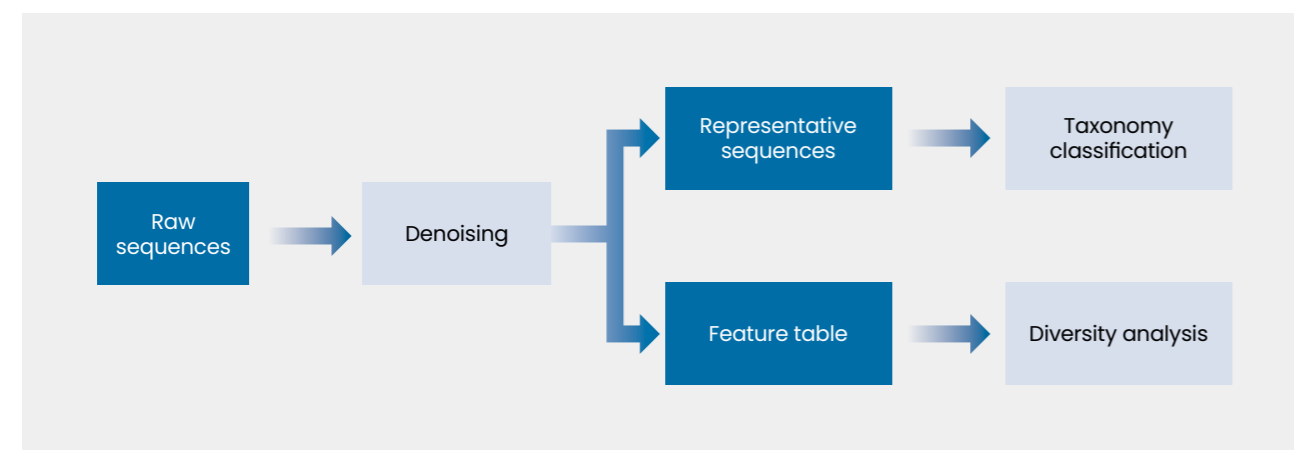
## DESCRIPTION

Metagenomic Sequencing Service and Kit is used for microbiome and mycobiome studies. The service allows for characterizing and differentiating a myriad of microbial species. The 16S V4 (or V3-V4) region of bacteria and archaea and 18S ITS1 (or ITS1-ITS2) region of fungi is amplified by PCR. After cleaning up using CeleMag beads, the indices and adapters are attached for NGS and bioinformatics analysis. According to the purpose of customer's studies, various analysis reports are provided by the Celeemics robust analysis pipeline. Please contact us for further information.

## EXPERIMENT WORKFLOW

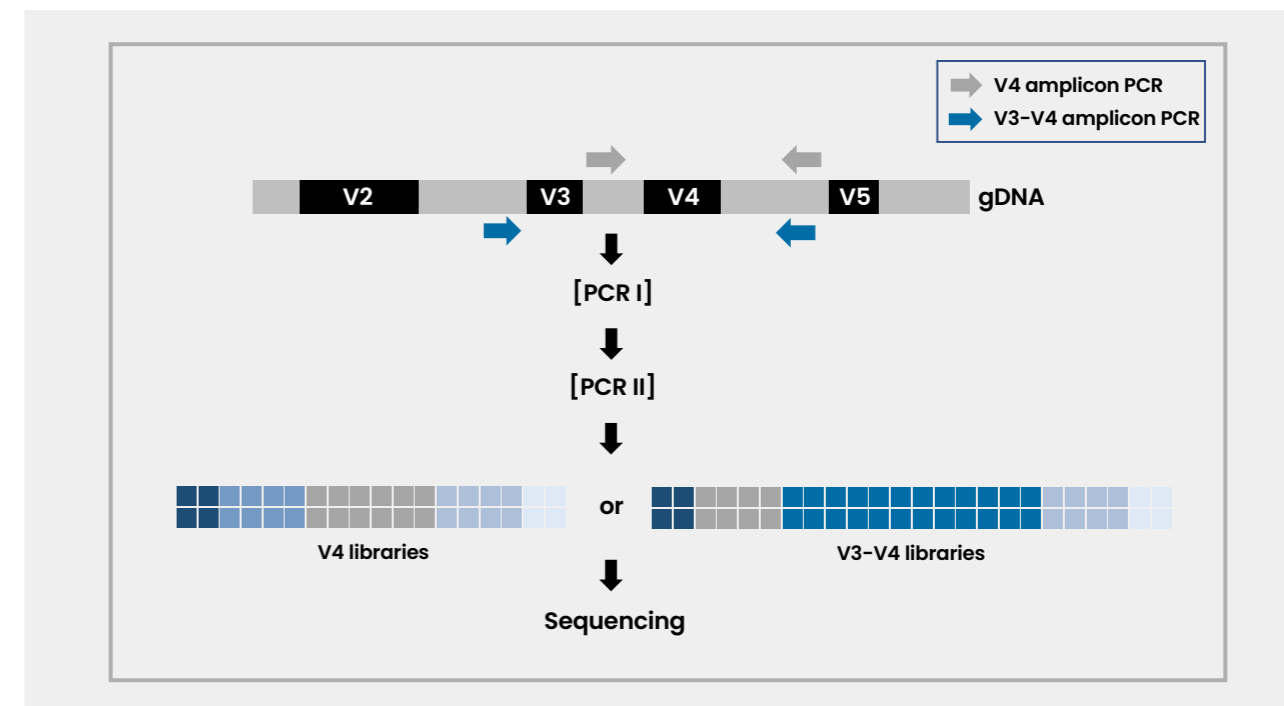
1. PCR amplification against gDNA using 16S region or ITS region specific primers
2. Bead cleanup
3. Index and adapter ligation with Nextera Index sets
4. Bead cleanup
5. Library pooling
6. NGS Sequencing

## NGS-BASED METAGENOME ANALYSIS WORKFLOW

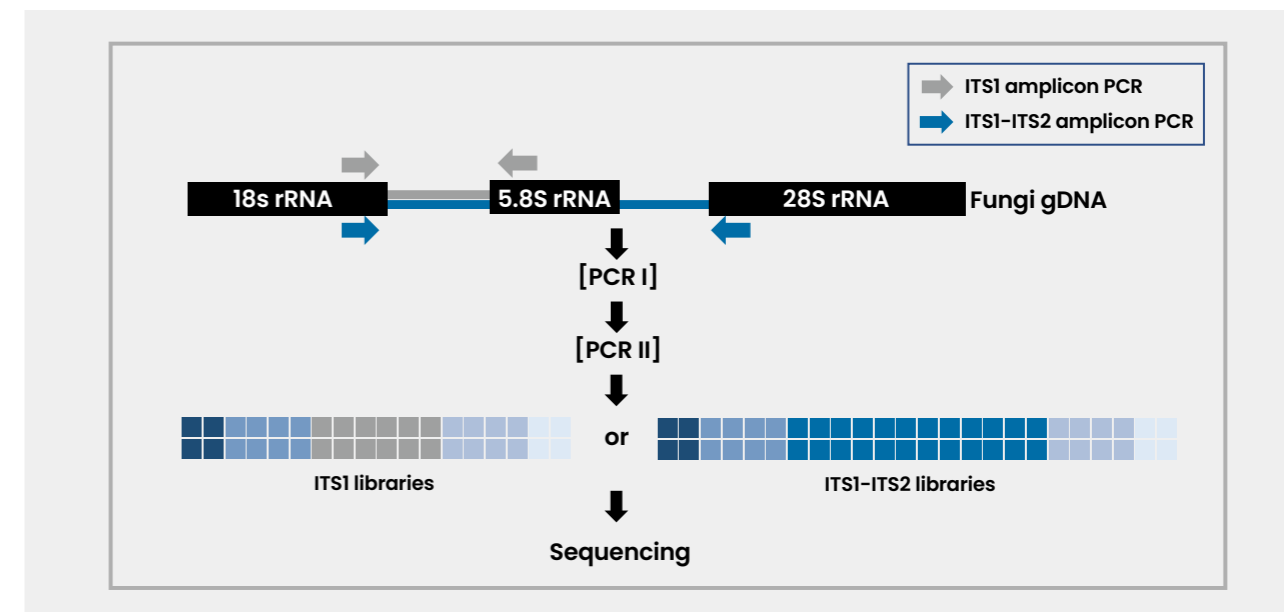


## SEQUENCING WORKFLOW

16S rRNA V4 and V3-V4

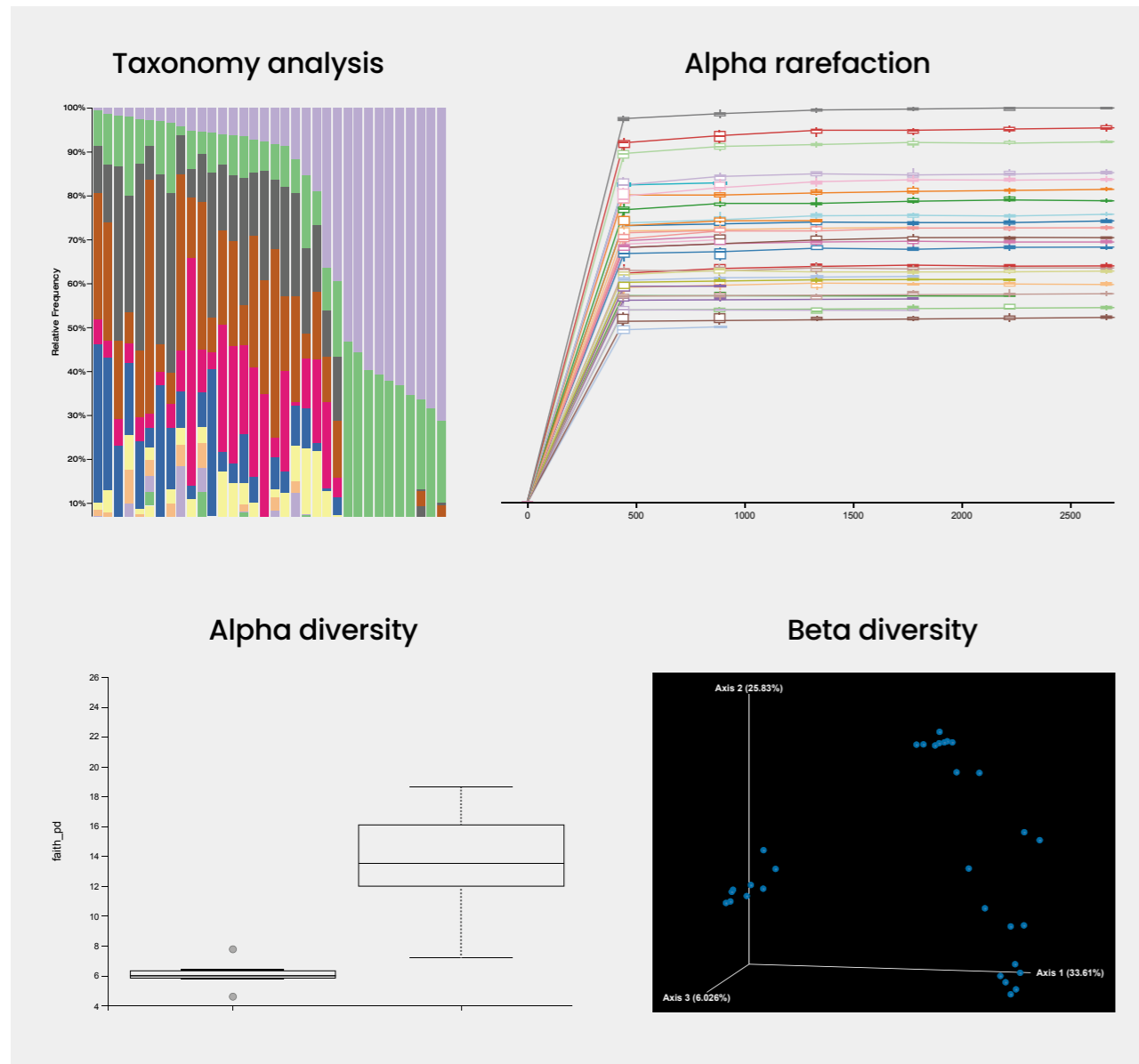


18S ITS1 and 18S ITS1-ITS2





## EXAMPLE OF METAGENOMIC SEQUENCING ANALYSIS REPORT



Results presented above are a few selected examples of the metagenomics sequencing results that Celemics provides. Contact us for more information.



# Barcode Tagged Sequencing™ (BTSeq™)

CELEMICS PRODUCTS & SERVICES 2022

BTSeq™ – Standard Service and Kit  
BTSeq™ – Viral Analysis Service  
BTSeq™ Mitochondrial DNA Sequencing Service  
BTSeq™ Full Plasmid Sequencing Service



# Barcode-Tagged Sequencing™ (BTSeq™)

# BTSeq™ – Standard Service and Kit



## Wide Range of DNA Sizes

- No limitation of DNA size: 200 bp - 20 kb or longer
- Plasmid sequencing with large insert DNA



## NGS-based, High Sequencing Accuracy

- NGS-based high sequencing quality
- Digitized sequencing results



## Fast TAT, No Need for Primer Walking

- NGS-based result, within 24 hours after sample arrival
- No need for primer synthesis
- No need of repetitive Sanger sequencing cycles



## Cost-effective

- Unparalleled cost-effectiveness compared to Sanger
- Only sequencing primer information required, eliminating the need for synthesizing the primers



## No Limitation of Origin

- Sequencing samples of various species
- Virus, Bacteriophage, Mycobiome, etc.



## No Need of High Concentration Sample

- Compatible with unpurified PCR products
- Low-amount sample requirements as little as 10 ng/μl

## DESCRIPTION

For the last few decades, Sanger Sequencing has been the standard for analyzing DNA sequences. Due to its need for repetitive primer design, primer synthesis, and sequencing steps during Primer Walking when analyzing long sequences, however, it requires lengthy experimental time and large costs to perform. Additionally, issues such as high re-experimentation rates, intermittent errors, and a less than 1 kb read length limitation have made sequence analysis difficult for clients. To overcome these limitations, Celemics created an NGS-based molecular barcoding technology and NGS error elimination algorithm solution, allowing for the analysis of sequences with lengths greater than 1kb without the need of sequencing primers.

## KEY FEATURES

### Long DNA sequencing, No need of sequencing primer

Analyze from 200 bp to 20 kb and longer length in a single reaction  
No need of sequencing primer\*  
No need of repetitive primer walking for long DNA de novo sequencing

### Cost-effective, highly accurate, rapid turnaround time

Novel NGS-based proprietary enzyme and bioinformatics technology  
Cost-effective sequencing compared to Sanger sequencing  
Secure sequencing accuracy with NGS-based sequencing that yields more reliable results than Sanger sequencing  
Receive digitized results within 1-2 business days

### Wide compatibility

Various applications with no limitation on DNA size or sample types across a broad range of origins  
Compatible with unpurified PCR products\*\*

## BTSeq™ SERVICE

- BTSeq™ – Viral Analysis Service
- Mitochondrial DNA Sequencing Service
- Full Plasmid Sequencing Service
- Microbial Identification Service

\* Only primer sequence information is required

\*\* Only samples that have single bands from gel electrophoresis are accepted

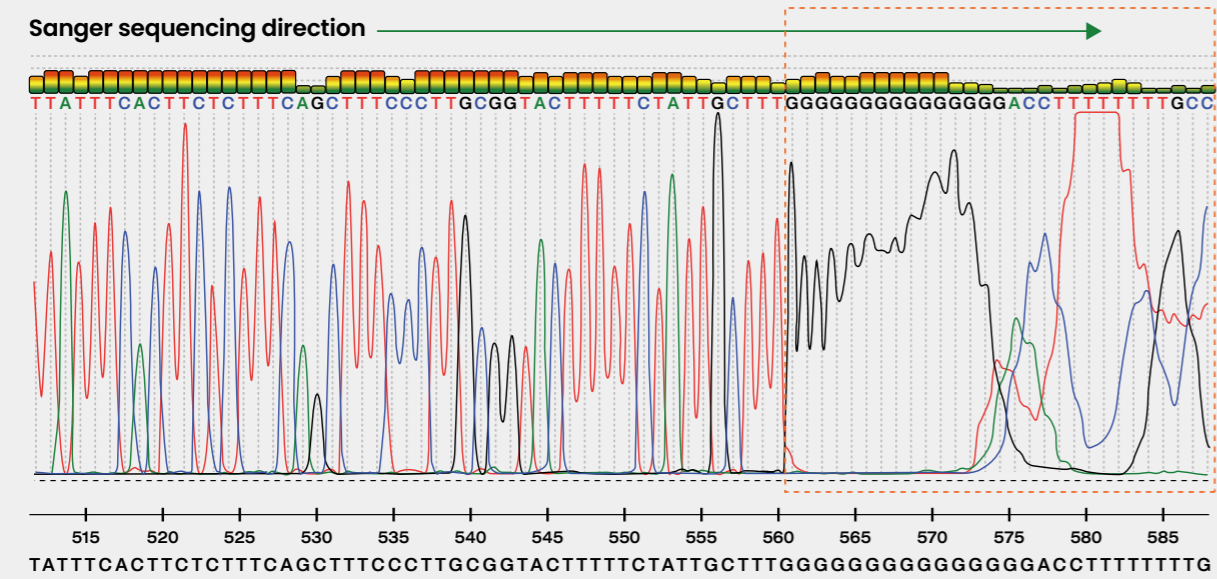
## HIGH ACCURACY ACHIEVED BY NGS-BASED BTSeq™ SEQUENCING SERVICE

Sanger sequencing has been the gold standard sequencing method. Although Sanger sequencing service providers have supported researchers for several decades, the high competition among providers led to cost reduction in Sanger sequencing reagents. Most Sanger sequencing service providers started diluting the reagents and applying methods that are not recommended for the best quality result. This has resulted in inaccurate sequencing results and repetitive sequencing cycles.

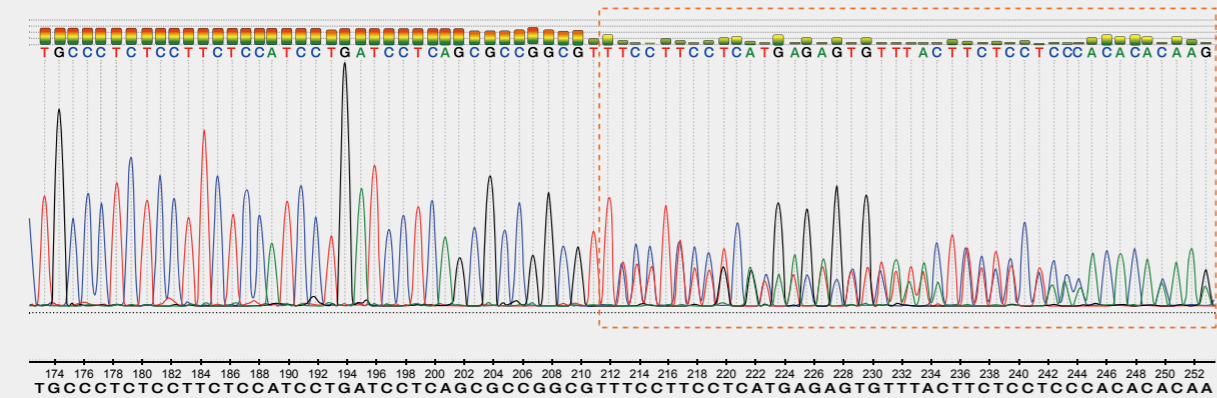
While Sanger sequencing may have many limitations such as homopolymer sequencing, multi-peak problems, and detecting Indel or frameshift mutations, BTSeq™ overcomes such limitations and provides accurate sequencing data even from poor quality or low-amount samples.

### Limitations of Sanger sequencing

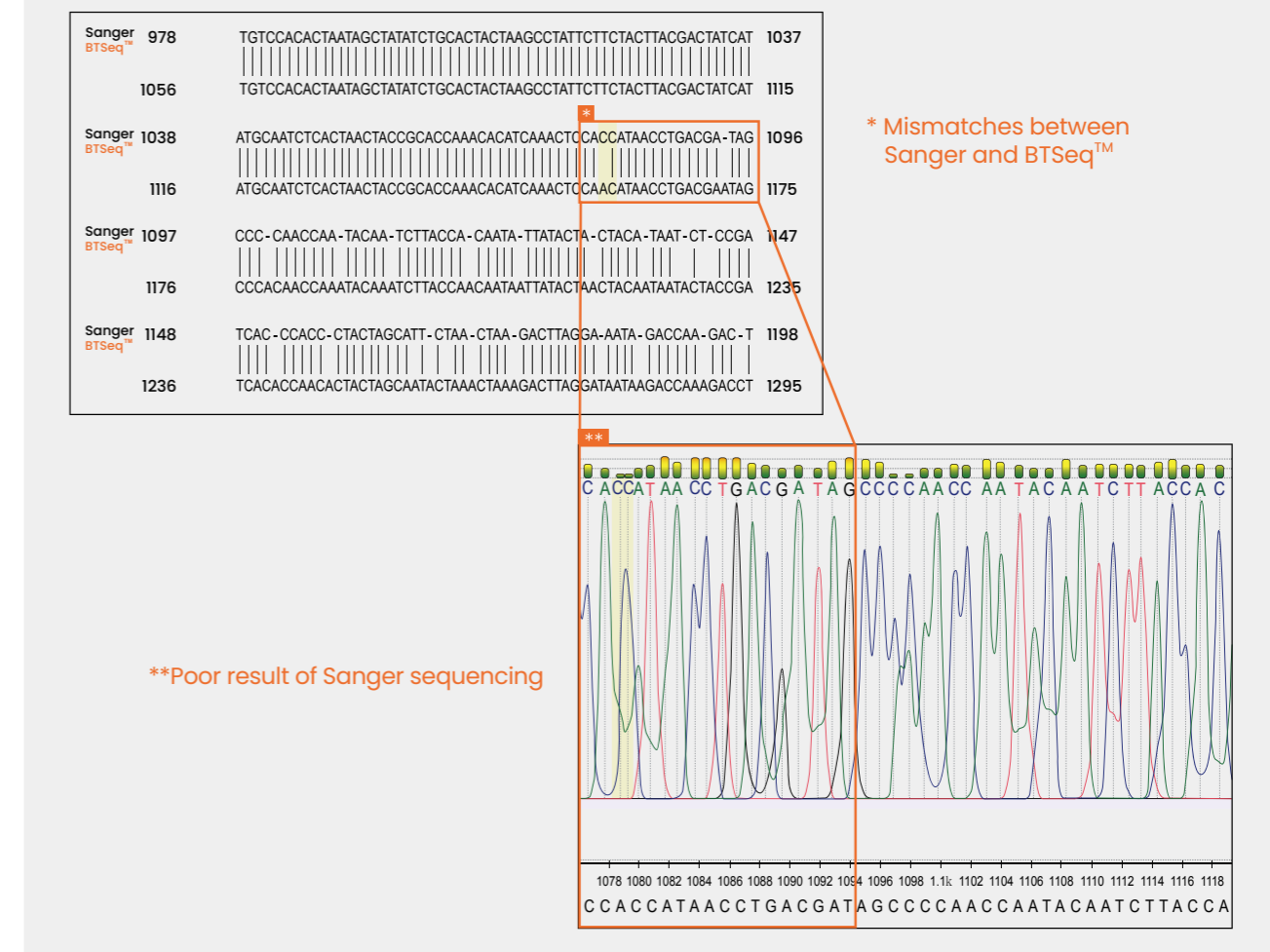
Example of homopolymer problem in Sanger sequencing



Example of multi-peak problem in Sanger sequencing



### Accurate sequencing of BTSeq™



Most mismatches between BTSeq™ and Sanger sequencing results were due to the minor peaks or poor-quality results from Sanger sequencing.

### Comparison between Sanger and BTSeq™

	Sanger	BTSeq™
Data type	Analog	Digital
Data quality	Ambiguous	Clear
Analysis size	Up to 1 kb	Up to 20 kb or longer
Sample concentration	> 100 ng/μl	> 10 ng/μl
Sample amount	> 20 μl	> 10 μl

## COMPARISON TEST

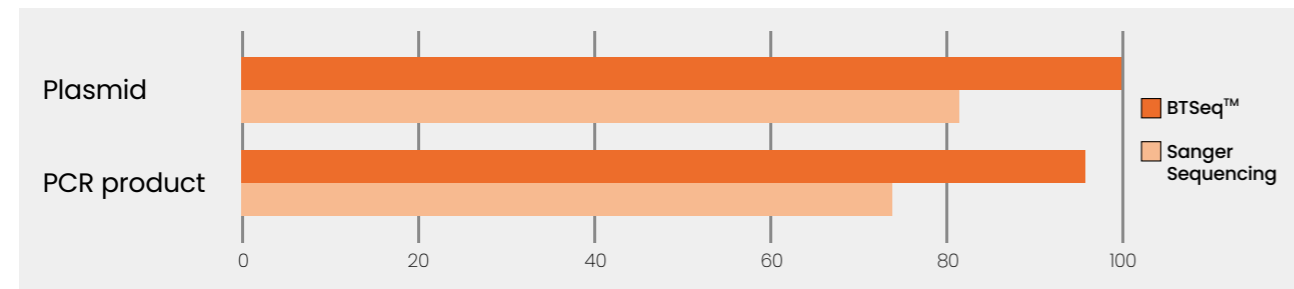
To assess the performance of BTSeq™, we have conducted multiple comparison tests with Sanger sequencing method. The samples that were sequenced by Sanger were provided by a Sanger Sequencing partner and randomly selected for BTSeq™ validation test. More than 80% of PCR product samples were not purified. The sample concentration ranged from 0.1 ng/μl to 200 ng/μl and 1 μl (0.1 ng – 200 ng) of each sample were used for BTSeq™. The results show high concordance of BTSeq™ with Sanger sequencing with even higher accuracy.

### BTSeq™ shows errorless sequencing results

Number of Samples	Plasmid (n=454)		PCR product (n=801)	
	Method	BTSeq™	Sanger Sequencing	BTSeq™
Unidentified	0	85**	36*	211**
Identified	454	369	765	590
Analysis success rate (%)	100.0%	81.3%	95.5%	73.7%

\* Long repeated sequences  
\*\* Poor sequencing results

### Analysis Success Rate (%)



## DIGITIZED RESULTS\*

The BTSeq™ service, an NGS-based sequencing service, provides digitized results by standalone bioinformatics analysis software enabling various options for result data

**Output type1 : ABI file**

sample .abi

**Output type3 : FASTQ file**

sample .fastq

```
@Sample_1
ACCCCTGAATTGACTCTCTCCGGGCGCTATCATGCCA
+
HHHHHHHHHHHHHHHHHHHHHHHHHHHHHHHHHHHHHH
```

**Output type2 : FASTA file**

sample .fasta

```
>Sample_1
ACCCCTGAATTGACTCTCTCCGGGCGCTATCATGCCA
```

**Output type4 : Base frequency table file**

sample .xlsx

Pos.	Ref.	A	C	T	G
1	A	121	1	1	0
2	C	0	126	0	0
3	C	0	126	0	0
4	C	0	126	0	0
5	T	0	0	126	0

\* Different options provided for different applications. Contact us for more information.

## BTSeq™ SERVICE OPTIONS

Product Group	Service Option	Sample Type	Description
BTSeq™	BTSeq™ – Standard	PCR product / Plasmid	Primer sequence information is required*
	Plasmid Extraction	E. coli	-
	BTSeq™ – Raw Data	PCR product / Plasmid	Provides FASTQ file only

## BTSeq™ SERVICE PROCESS

Celemics has developed sample preparation techniques and bioinformatics software enabling cost-effective workflow. The BTSeq™ sequencing provides highly accurate results with short turnaround time (TAT) by effectively

correcting sequencing errors and generating consensus sequence with Celemics proprietary techniques.



# BTSeq™ – Viral Analysis Service

## DESCRIPTION

In most cases, RNA of the host cell is separated and purified along with viral RNA during extraction. This leads to an excessive amount of data being required to perform typical Total RNA-seq compared to the entire viral genome, leading to low-quality data and high costs. Celeemics solves this issue by developing extremely uniform amplification technology and bioinformatics software, which in turn provides quality data by efficiently eliminating any gaps generated from bias in the RT-PCR step.

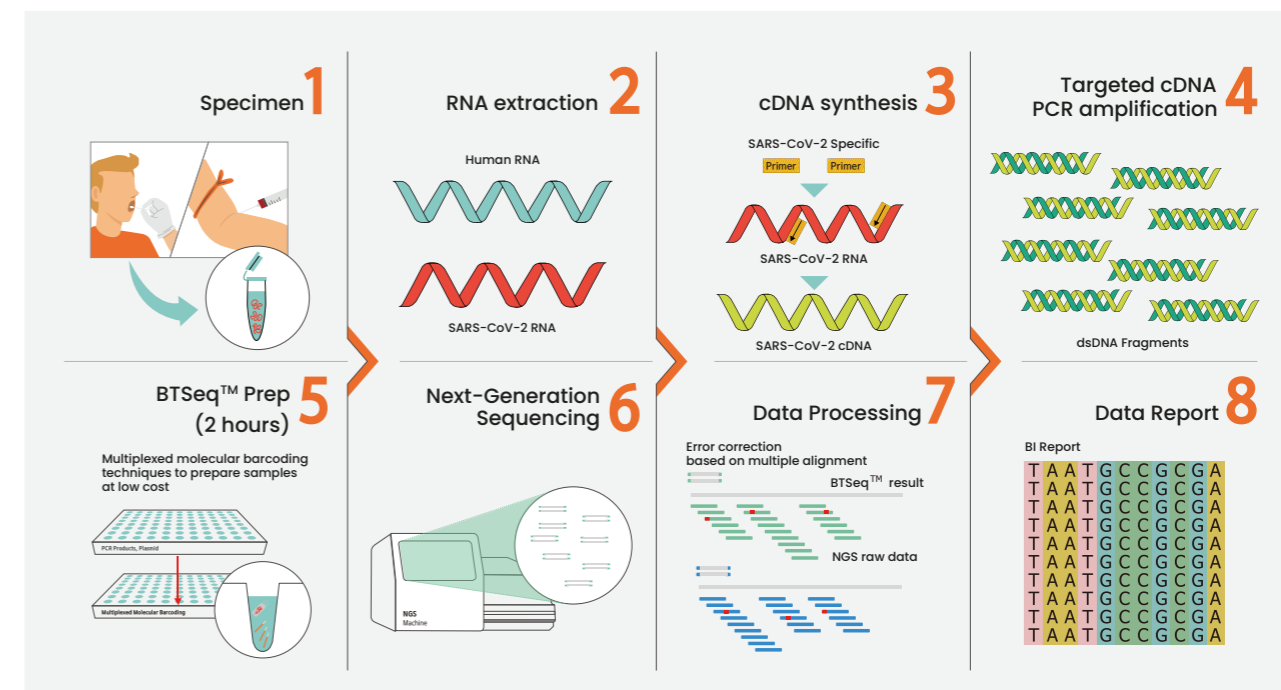
## KEY FEATURES

<b>1. High quality result generation from minuscule low-quality RNA samples</b>	Enabling of high-quality whole genome analysis, even in minuscule low-quality RNA samples extracted from upper respiratory tract, nasopharyngeal, oropharyngeal swab clinical specimens
<b>2. Results provided within 24 hours</b>	Provision of whole novel coronavirus genome within 24 hours using Celeemics' proprietary reagent and bioinformatics technology
<b>3. High-quality data generation at cost-effective price</b>	High-quality result generation, even from minuscule amounts of clinical samples

## REQUIREMENTS

<b>Sample type</b>	RNA
<b>Concentration</b>	Ct value < 25
<b>Volume</b>	40 µl
<b>Turnaround time</b>	Within 3-5 business days from sample collection
<b>Shipment</b>	Shipping on dry ice (essential)

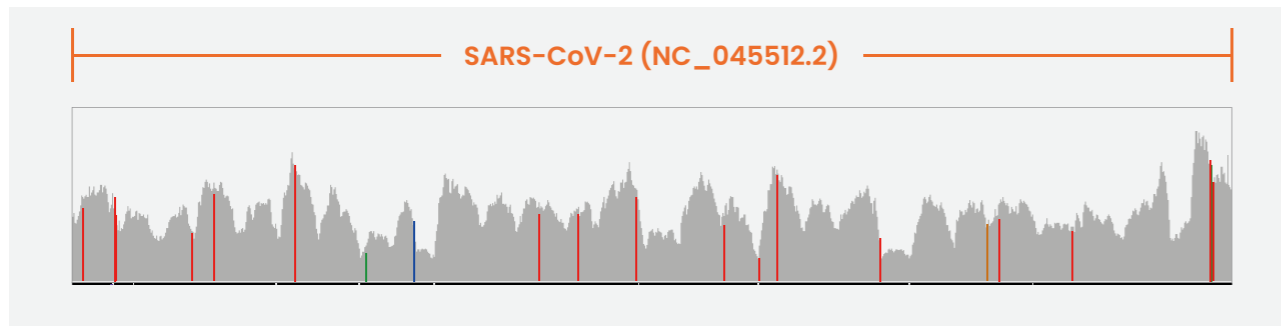
## SERVICE PROCESS



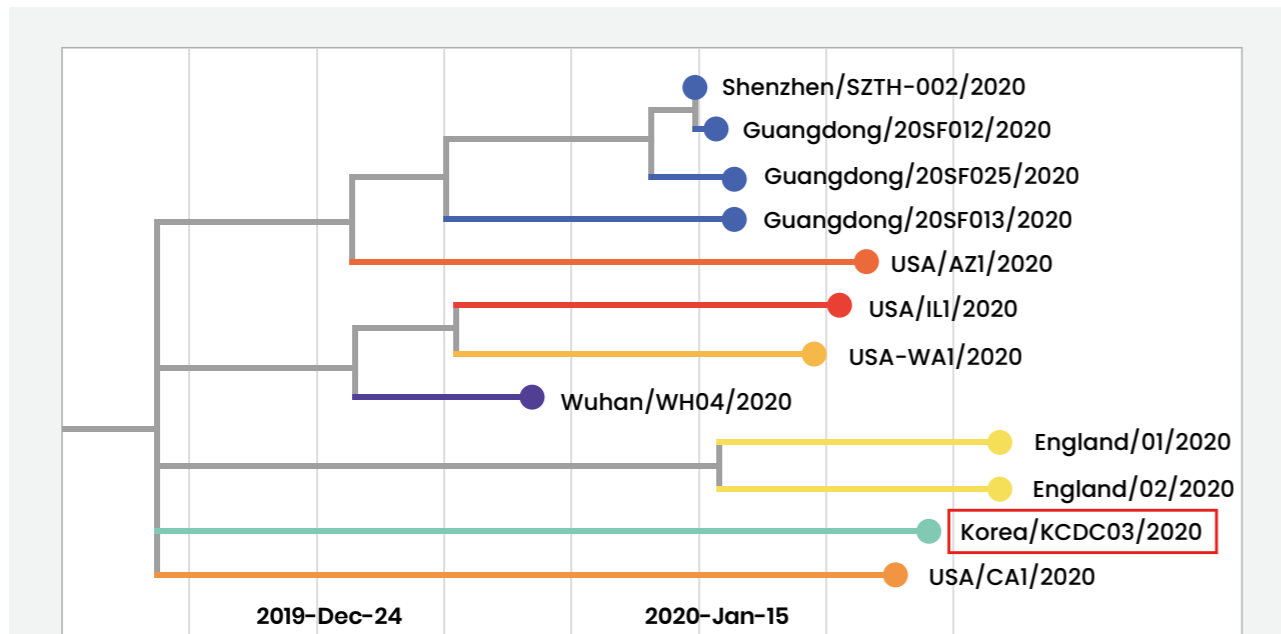
## COMPARISON BETWEEN TOTAL RNA SEQUENCING AND BTSeq™

	Total RNA-Sequencing	BTSeq™
<b>cDNA Synthesis</b>	<b>Total RNA (host and viral) into cDNA</b> : Leads to unnecessary data and high sequencing cost	<b>Only viral RNA to cDNA</b> : Utilizes Virus specific multiplex primers to selectively amplify viral RNAs from total RNAs
<b>Target Enrichment</b>	<b>No target enrichment required</b> : Viral RNA coexists with host RNA when cDNA synthesis is performed	<b>Viral genome is specifically amplified</b> : Only a small amount of viral RNA is required
<b>Library Preparation</b>	RNA library prep using RNA library kit: <b>4 hours</b>	Simple library prep using BTSeq™ reagent: <b>2 hours</b>
<b>Data Analysis</b>	: Mapped to the viral genome : Read/assembly based classification	: Mapped to the viral genome : Read/assembly based classification
<b>Turnaround Time</b>	<b>2-3 weeks</b>	<b>1-2 days</b>

## FULL COVERAGE OF SARS-COV-2 WGS ANALYZED BY BTSeq™ FROM PATIENT SPECIMENS



## IDENTIFICATION OF KOREA/KCDC03/2020 USING BTSeq™ – VIRAL ANALYSIS SERVICE



### Korea/KCDC03/2020

<b>Collection date</b>	2020-01-26
<b>Authors</b>	Kim et al
<b>Title</b>	Newly discovered betacoronavirus, 2019-2020
<b>Country</b>	South Korea
<b>Admin division</b>	Gyeonggi
<b>Host</b>	Human
<b>Location</b>	Gyeonggi

## REFERENCE OF BTSeq™ – VIRAL ANALYSIS SERVICE

GENOME SEQUENCES

### Genome Sequences of Two GH Clade SARS-CoV-2 Strains Isolated from Patients with COVID-19 in South Korea

Minwoo Kim,<sup>a</sup> Youn-Jung Lee,<sup>b</sup> Jae Sun Yoon,<sup>b</sup> Jin Young Ahn,<sup>b</sup> Jung Ho Kim,<sup>b</sup> Jun Yong Choi,<sup>b</sup> Jong-Won Oh<sup>b</sup>

<sup>a</sup>Department of Biotechnology, Yonsei University, Seoul, South Korea  
<sup>b</sup>Division of Infectious Diseases, Department of Internal Medicine, Yonsei University College of Medicine, Seoul, South Korea

Minwoo Kim and Youn-Jung Lee contributed equally to this work. Author order was determined by drawing straws.

**ABSTRACT** We report the genome sequences of two GH clade severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) strains isolated from nasopharyngeal swabs from patients with coronavirus disease 2019 (COVID-19) in South Korea. These strains had two mutations in the untranslated regions and seven nonsynonymous substitutions in open reading frames, compared with Wuhan/Hu-1/2019, showing 99.96% sequence identity.

Using the QIAamp viral RNA minikit (Qiagen, Hilden, Germany), RNA was extracted from the virus, which had been purified by passaging the swab samples three times on Vero cells (ATCC CCL-81) by the limiting dilution method (4). Viral cDNA synthesized using ProtoScript II reverse transcriptase (New England Biolabs, Ipswich, MA, USA) was amplified as described previously (5, 6), using in-house-designed primer sets and the Illumina platform-based BTSeq SARS-CoV-2 whole-genome sequencing (WGS) kit (Celemics, Seoul, South Korea) for multiplex amplicon sequencing on a MiSeq sequencer (150-bp paired-end mode; Illumina, San Diego, CA, USA). After dual-index

patient 8. All of the studies were approved by the institutional review board (IRB) of Severance Hospital, Yonsei University Healthcare System, with written informed consent from the patients (IRB protocol number 4-2020-0076).

Using the QIAamp viral RNA minikit (Qiagen, Hilden, Germany), RNA was extracted from the virus, which had been purified by passaging the swab samples three times on Vero cells (ATCC CCL-81) by the limiting dilution method (4). Viral cDNA synthesized using ProtoScript II reverse transcriptase (New England Biolabs, Ipswich, MA, USA) was amplified as described previously (5, 6), using in-house-designed primer sets and the Illumina platform-based BTSeq SARS-CoV-2 whole-genome sequencing (WGS) kit (Celemics, Seoul, South Korea) for multiplex amplicon sequencing on a MiSeq sequencer (150-bp paired-end mode; Illumina, San Diego, CA, USA). After dual-index filtering and adapter trimming using in-house scripts, reads (69,447 and 66,754 reads for isolates YS006 and YS008, respectively) were mapped to the reference sequence of Wuhan/Hu-1/2019 (GenBank accession number MN988668) (nucleotides 1 to 29870) (7) with BWA v0.7.17-r1188 (8), generating consensus genome sequences of strains SARS-CoV-2/human/KOR/YS006/2020 (29,825 nucleotides) and SARS-CoV-2/human/KOR/YS008/2020 (29,826 nucleotides) isolated from patients 6 and 8, respectively, with average coverage depths of 98.65× and 95.5×, respectively. The consensus sequences for YS006 (nucleotides 16 to 29840) and YS008 (nucleotides 16 to 29841) had no indels. The nearly complete genomes of these isolates, which lack 15 nucleotides and 29 or 30

**Citation** Kim M, Lee Y-J, Yoon JS, Ahn JY, Kim JH, Choi JY, Oh J-W. 2021. Genome sequences of two GH clade SARS-CoV-2 strains isolated from patients with COVID-19 in South Korea. *Microbiol Resour Announc* 10:e01384-20. <https://doi.org/10.1128/MRA.01384-20>.

**Editor** Simon Raux, DOE Joint Genome Institute

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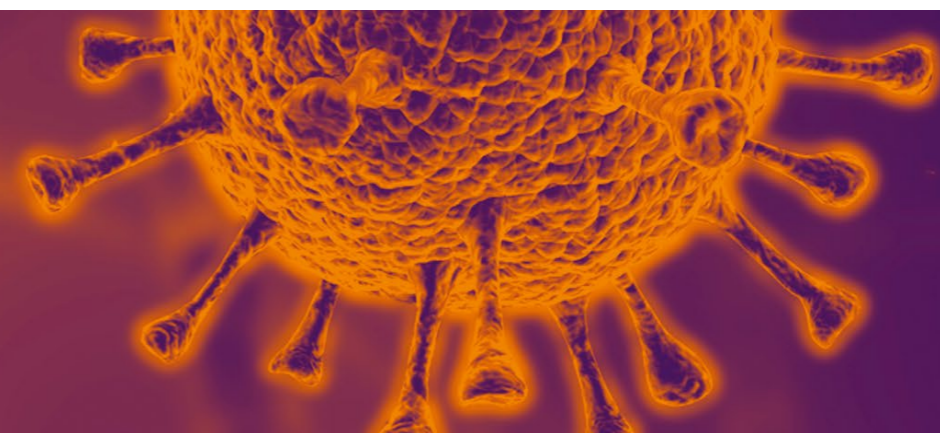
**Address correspondence** to Jun Yong Choi, seran@yuh.ac, or Jong-Won Oh, jwch@yonsei.ac.kr.

**Received** 4 December 2020  
**Accepted** 14 December 2020  
**Published** 7 January 2021

January 2021 Volume 10 Issue 1 e01384-20

mra.asm.org 1

\*Reference: Genome Sequences of Two GH Clade SARS-CoV-2 Strains Isolated from Patients with COVID-19 in South Korea, American Society Microbiology. (2020)



# BTSeq™ Mitochondrial DNA Sequencing Service

## DESCRIPTION

The BTSeq™ Mitochondrial DNA Sequencing enables accurate analysis of clinical variability and genetic heterogeneity. By sequencing 17 kb-long mtDNA with newly developed NGS-based technology, customers can decipher the instability and variations of mtDNA associated with many metabolic and neurologic disorders and cancers. The service provides highly accurate results with fast TAT and cost-effectiveness.

## REQUIREMENTS

Sample Type	gDNA
Concentration	50 ng/μl
Volume	10 μl
Turnaround time	Within 4 business weeks from sample arrival
Shipment	Ship on ice

## RESULT EXAMPLE OF BTSeq™ MITOCHONDRIAL DNA SEQUENCING

### Example of mitochondrial variant analysis report

Sample	Gene	Amino Acid Change	Type	Allele		Sequencing Depth			VAF	Associated Disease
				Ref	Alt	Total	Ref	Alt		
Sample 1	ND2	p.Leu237Met	Missense	C	A	16370	0	16361	-	Blood iron metabolism
	ATP8	p.Leu17Phe	Missense	C	T	16155	8	16139	-	Longevity
	ND5	p.Asn30fs	Frameshift	-	A	17593	17197	197	1.12%	
Sample 2	ND2	p.Thr122Ala	Missense	A	G	16759	14	8005	-	AD, PD
	ATP6	p.Met58Thr	Missense	T	C	16909	15721	12	4.53%	-
	ND3	p.Thr114Ala	Missense	A	G	20141	8	219	1.24%	Breast cancer risk
Sample 3	ND2	p.Leu237Met	Missense	C	A	5100	0	5100	-	Blood iron metabolism
	ATP8	p.Leu17Phe	Missense	C	T	16353	6	16340	-	Longevity
	ND5	p.Asn30fs	Frameshift	-	A	16960	16625	193	1.14%	405

### Example of summary report of NGS operation (target size: 16.6 kb)

Sample name	Raw read	Raw base	Total read	Filtered ratio	On target read ratio	On target base ratio	Uncovered	20x coverage	50x coverage	100x coverage
Sample 1	3,521,438	531,737,138	3,486,316	99.00%	90.78%	95.36%	0.00%	100.00%	100.00%	100.00%
Sample 2	3,514,296	530,658,696	3,479,540	99.01%	91.39%	95.82%	0.00%	100.00%	100.00%	99.99%
Sample 3	3,526,146	532,448,046	3,489,580	98.96%	90.12%	95.24%	0.00%	100.00%	100.00%	100.00%
Sample 4	3,500,420	528,563,420	3,463,806	98.95%	90.85%	95.67%	0.00%	100.00%	100.00%	100.00%





# BTSeq™ Full Plasmid Sequencing Service

## DESCRIPTION

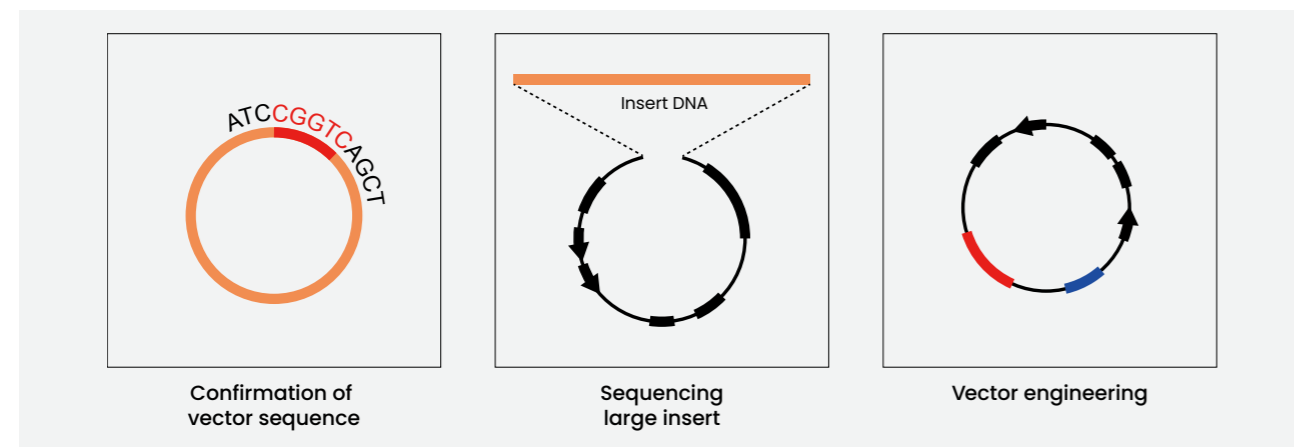
The BTSeq™ Full Plasmid Sequencing Service allows for the most effective analysis of the full-length sequencing of plasmids with shorter TAT and lower cost than Sanger sequencing. The service is ideal for protein engineering, vector engineering, antibody optimization, synthetic biology, and various other applications.

## REQUIREMENTS

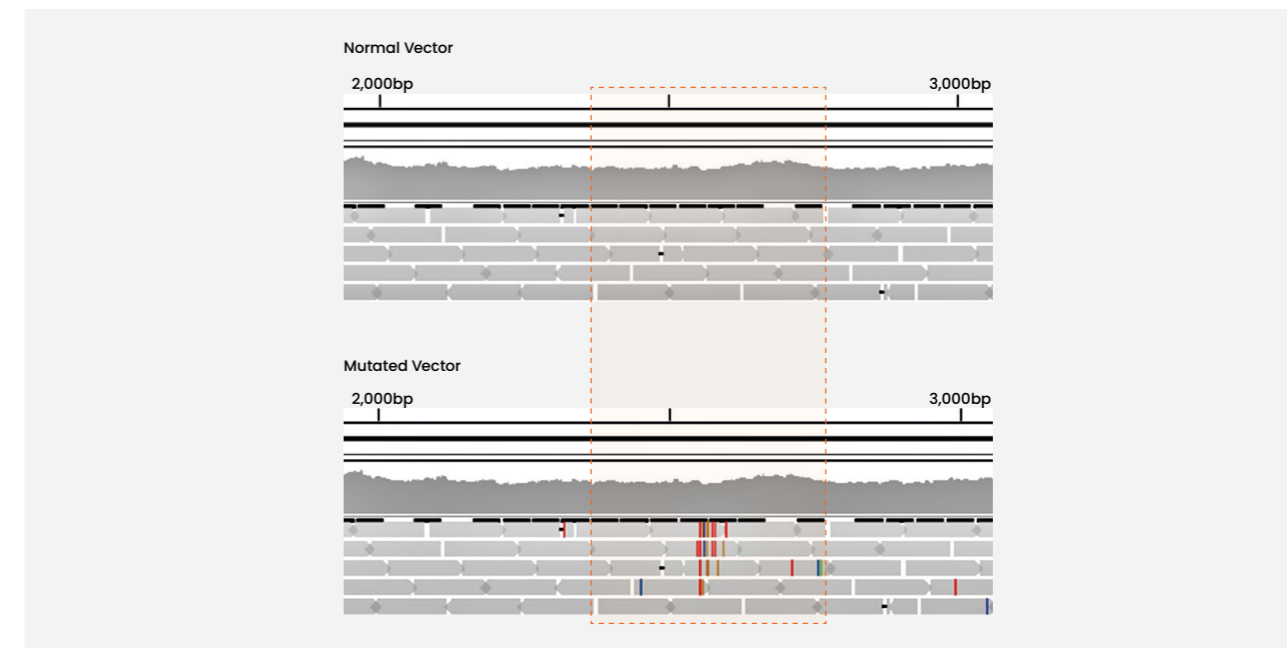
Sample type	PCR product, Plasmid*
Concentration	10 ng/μl
Volume	10 μl
Turnaround time	Within 1 business day from sample arrival
Packaging	1) RT 2) Ship on ice (Recommended)

\* Contact us for plasmids longer than 20kb

## APPLICATIONS OF BTSeq™ FULL PLASMID SEQUENCING



## COMPARISON IGV DATA BETWEEN NORMAL AND MUTATED VECTORS



# CELEMICS SOLUTIONS FOR IMMUNE REPERTOIRE SEQUENCING

CELEMICS PRODUCTS & SERVICES 2022

Immune Repertoire Profiling Service  
TrueRepertoire™ Service



# Immune Repertoire Profiling Service

## DESCRIPTION

Immune repertoire often represents an individual's current immunological status; whether the person is healthy, vaccinated, diseased, or infected. Only high-throughput NGS analysis can comprehensively profile an individual's immune repertoire. The Immune Repertoire Profiling Service provides effective data acquisition, integration, and interpretation for the customers.

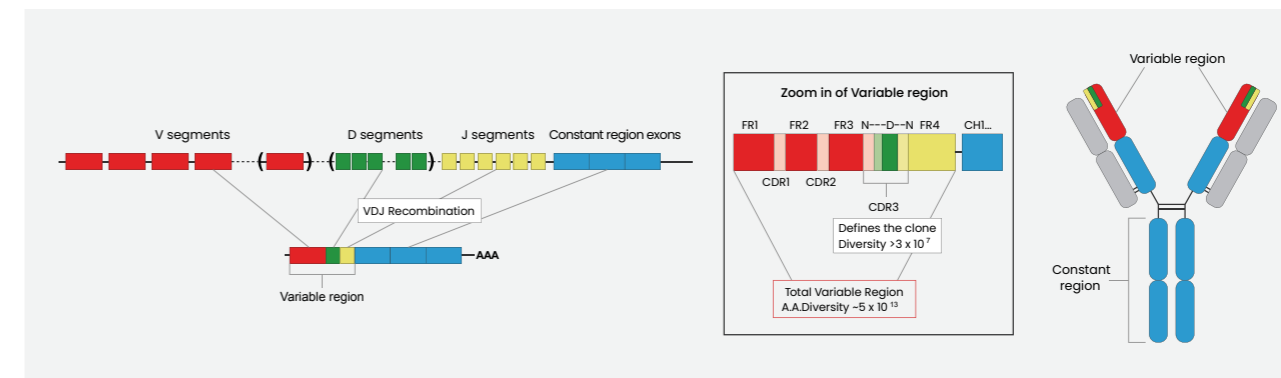
## KEY FEATURES

<b>1. Quantitative analysis of library diversity</b>	<ul style="list-style-type: none"> <li>- NGS-based analysis of complex antibody library consisting of millions (<math>10^6</math>-<math>10^{12}</math>) of sequences in a single experiment</li> <li>- Analysis of immunoglobulin and T-cell receptor repertoire; analysis of BCR/TCR for each clone</li> <li>- Frequency analysis of individual antibody clones within the library, identifying major and minor clones</li> </ul>
<b>2. Tracking of clonal frequencies for each sample</b>	<ul style="list-style-type: none"> <li>- For antibody discovery, analysis of library diversity according to its panning degree enabling monitoring changes in clonal frequency</li> <li>- Minimized omission of potentially significant antibody clones</li> <li>- Analysis of immune repertoire characteristics from blood sample and monitoring of each clone</li> </ul>
<b>3. Various analysis options for immune system studies</b>	<p>Perform the experiment with drastically reduced time and cost enabled by the advanced technology of MSSIC developed by Celemics</p>

## REQUIREMENTS

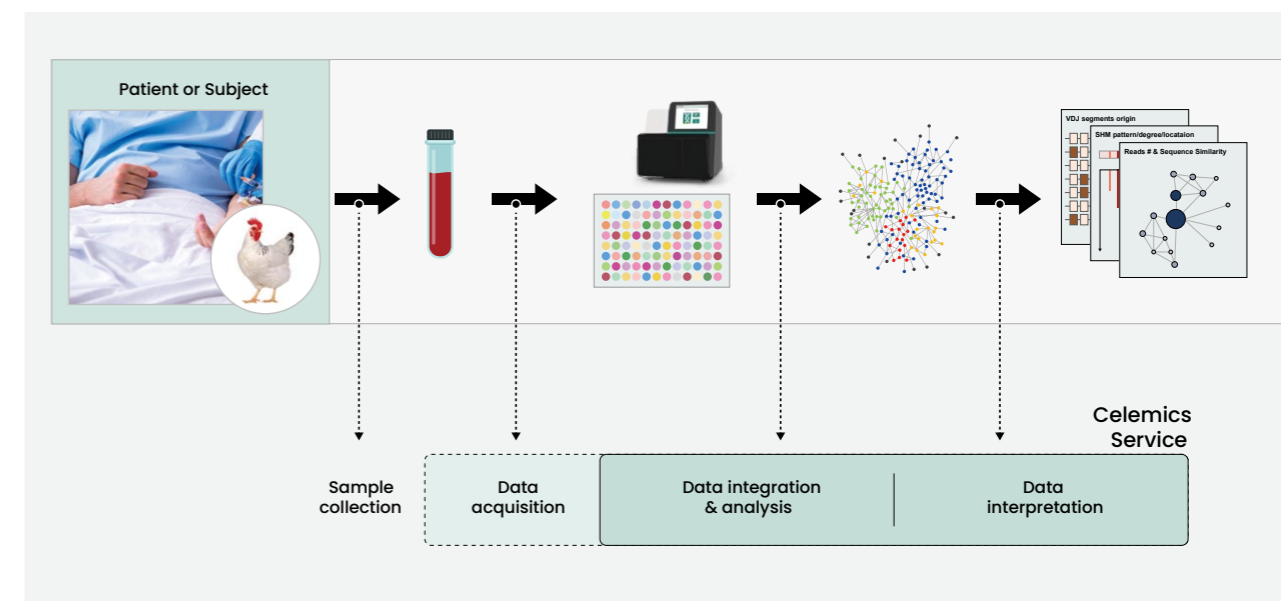
<b>Sample type</b>	Total RNA from B-Cell or/and T-Cell, DNA from B-Cell or/and T-Cell, DNA/RNA Amplicons
<b>Concentration</b>	100 ng/ $\mu$ l
<b>Amount</b>	1 $\mu$ g
<b>Turnaround time</b>	Within 4-6 business weeks from sample collection
<b>Temperature</b>	RT for storage and shipment

## DIVERSITY OF ANTIBODY



The antibody genes are composed of many different segments. The antibodies are presented in B cells with great diversity of  $10^{13}$  repertoires.

## GENERAL WORKFLOW



Celemics provides service for data acquisition, integration, and analysis, and interpretation.





# TrueRepertoire™ Service

## DESCRIPTION

The TrueRepertoire™ is a NGS-based antibody library sequencing platform developed to overcome the key issues of existing methods such as sequencing error, short-read length, and high-cost gene synthesis for further characterization. Celemics has developed a cloning microchip, barcode assay technology, and laser-based non-contact clone retrieval system and integrated into the newly developed platform, TrueRepertoire™ assay. This service allows for full sequence analysis of over 10,000 clones in a single experiment and thereby discovering rare clones. The TrueRepertoire™ service contains the client's antibody clone of interest within the library itself, eliminating the need to perform new gene synthesis and significantly reducing time and cost.

## KEY FEATURES

<b>1. Provision of antibody DNA sequence library containing over 10,000 errorless strains</b>	NGS-based sequence analysis and high-capacity clone separation and molecular barcode assays using Celemics proprietary MSSIC technology
<b>2. V<sub>H</sub>-V<sub>L</sub> linkage analysis of each antibody</b>	Receive V <sub>H</sub> -V <sub>L</sub> linkage information, an area difficult to analyze through NGS due to its short read length
<b>3. Provision of physical property analysis of each antibody through bioinformatics analysis</b>	Clone frequency distribution within the library V <sub>H</sub> -V <sub>L</sub> sequence length distribution, post-translation modification information, CDR and frame amino acid information, etc.
<b>4. Retrieval of selected physical antibody allowing for convenient workflow</b>	Eliminates the need to perform new gene synthesis and reduces time and cost due to the antibody clones within the library itself, enabling isolation of physical DNA for further characterization

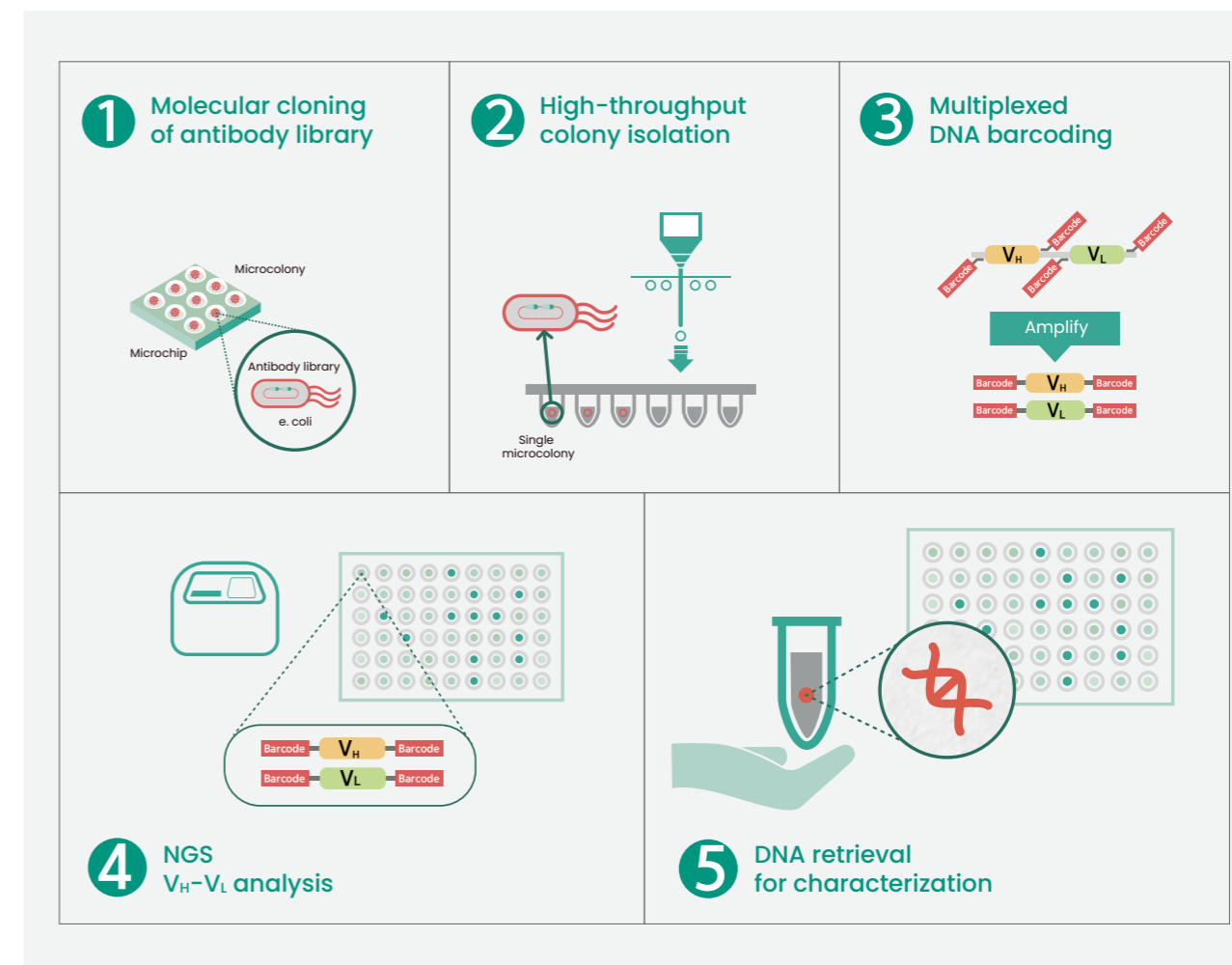
## REQUIREMENTS

<b>Sample type*</b>	Total RNA from B-Cell or/and T-Cell, DNA from B-Cell or/and T-Cell, DNA/RNA Amplicons
<b>Concentration</b>	100 ng/μl
<b>Amount</b>	1 μg
<b>Turnaround time</b>	Within 4-6 business weeks from sample collection**
<b>Temperature</b>	RT for storage and shipment

\* ~30 bp of Consensus upstream & downstream sequence over V<sub>H</sub> and V<sub>L</sub> region required

\*\* TAT depends on colony size

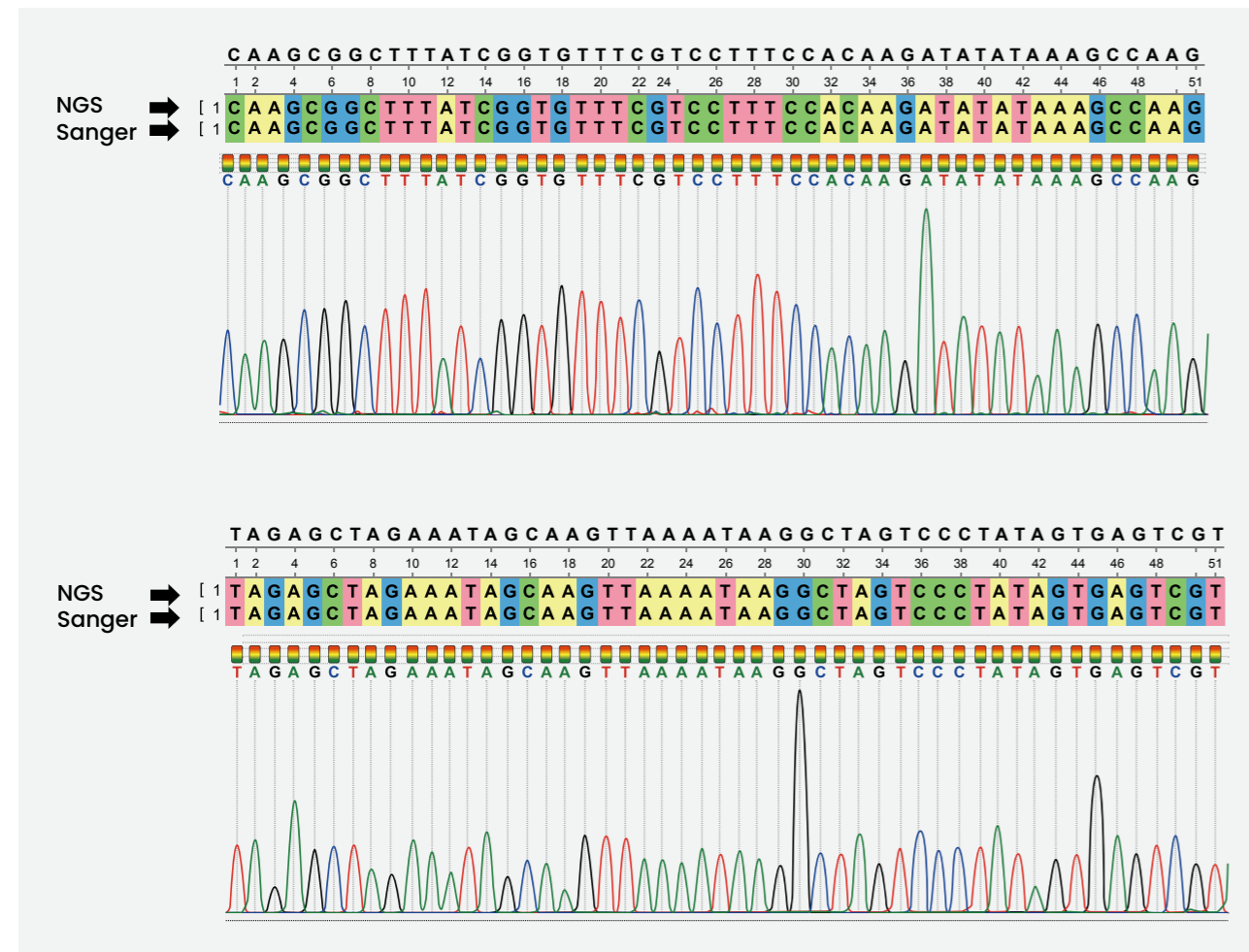
## HOW TrueRepertoire™ WORKS



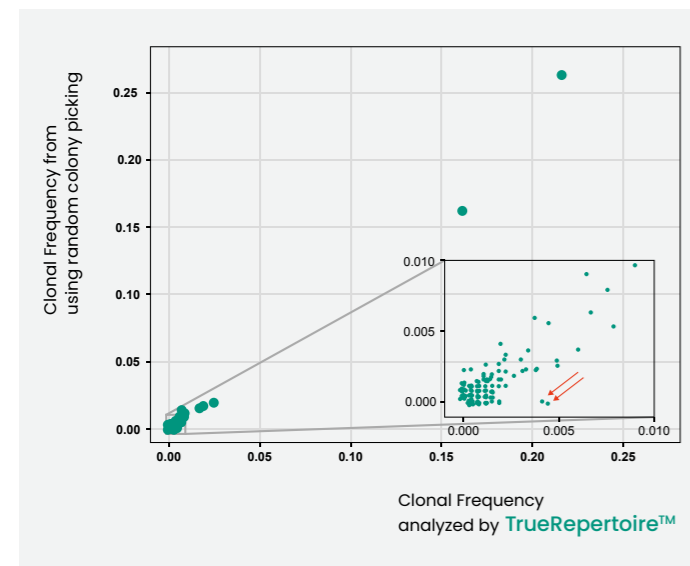
1. Celemics proprietary microcolony chip formation with high density, each colony starts from a single E. coli.
2. Extraction of the colonies from the microchip into microwell by Celemics' proprietary laser system
3. Multiplex PCR with barcoded primers from the isolated colonies
4. NGS and computation of the consensus sequences with cognate pairing of V<sub>H</sub> and V<sub>L</sub>
5. Clonal DNA retrieval based on the consensus sequence for further characterization

## VALIDATION TESTS

Validation I. Result of 480 randomly selected antibody clones from TrueRepertoire™ perfectly matched (480/480) Sanger sequencing results of their physical DNA

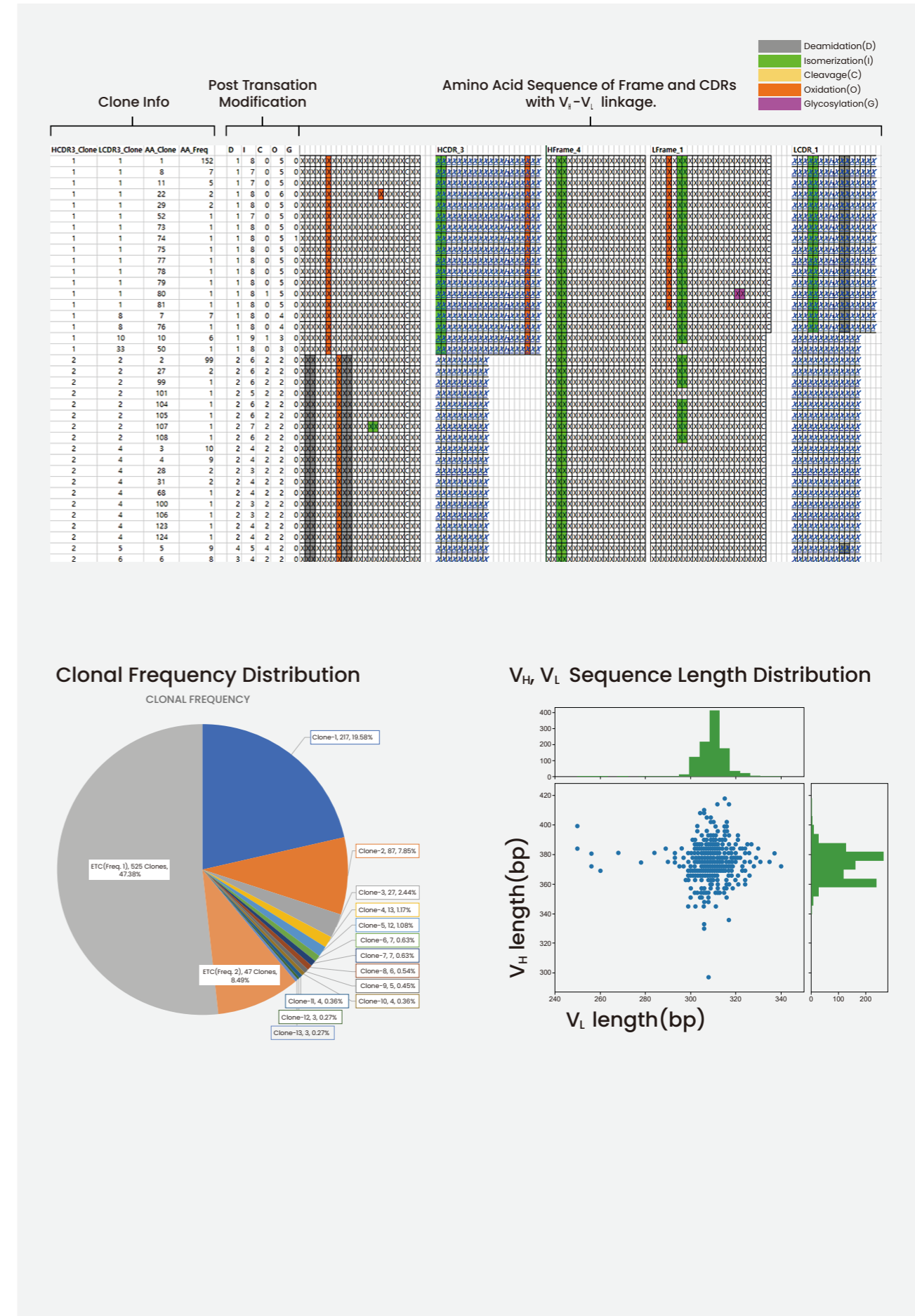


Validation II. Similar clonotype frequencies of major clones between TrueRepertoire™ and random colony picking



1. Major clones showed similar clonotype frequency in both platforms – random colony picking followed by Sanger sequencing and TrueRepertoire™
2. The result showed that there were newly identified clones found only in the TrueRepertoire™ results (red arrows)

## USER FRIENDLY TrueRepertoire™ REPORT



# MODULAR ACCESSORIES

CELEMICS PRODUCTS & SERVICES 2021

Library Preparation Kit - Standard / EP  
Double-Stranded cDNA Synthesis Kit  
Hybridization Enhancer  
CeleMag™ Clean-up Bead  
CeleMag™ Streptavidin Bead  
CLM Polymerase  
Bioinformatics Software



# Celemics Library Preparation Kit Standard / EP

## DESCRIPTION

Celemics Library Preparation Kit is optimized for high-efficiency Celemics panels. The Library Preparation Kits include End-repair, A-tailing enzyme mix, index primers (single or dual), adapters and buffers.

## LIBRARY PREPARATION WORKFLOW FOR TARGET ENRICHMENT NGS

DNA Fragmentation		
Standard Fragmentation	EP Fragmentation	Fragmentase
Option 1. Sonication	Option 2. Fragmentase	
Bead Purification & Quantification		
NGS Library Preparation		
ER/A	ER/A	ER/A
Adapter Ligation (Single/Dual Index)		
Bead Purification		
Index PCR		
Target Enrichment		

Celemics provides two methods for the library preparation step, Standard Library Preparation Kit and Enzymatic Preparation Kit (EP Kit). The Standard Library Preparation Kit includes all reagents for End repair (ER), A-tailing (A), and Adapter Ligation steps. For DNA fragmentation from Standard Library Preparation Kit, customers can use ultra-sonication devices or fragmentase. Fragmentase is provided by Celemics and included in the kit upon request. While the Standard Kit is composed of 4 different steps, the EP Kit includes all steps from enzymatic fragmentation to ER/A in a single reaction enabling convenient workflow. Since the purification step is not needed for EP Kit, the kit allows for minimal DNA loss which is a crucial factor for damaged DNA samples such as FFPE. EP Kit, provided by Celemics, includes all reagents required for library preparation.

Note.

For Option 1, ultra sonicator is not provided with the kit.  
For Option 2, the inclusion of the fragmentase in the kit is optional.

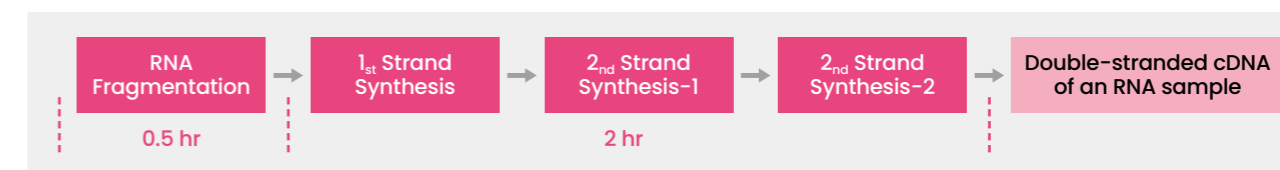


# Celemics Double-Stranded cDNA Synthesis Kit

## DESCRIPTION

Celemics Double-Stranded cDNA Synthesis Kit is optimized for NGS-based RNA sequencing. The kit includes all components from RNA fragmentation to double-stranded cDNA synthesis for NGS library preparation. The robust performance of the kit allows for the cDNA synthesis even from low amounts of RNA samples with high accuracy and reduced reaction time.

## cDNA SYNTHESIS WORKFLOW



Sample amount : 10 ng to 1 µg \*

Assay time : 30 minutes for RNA fragmentation and 2 hours for double-stranded cDNA synthesis

\* Carrier RNA is required for sample amount < 25 ng

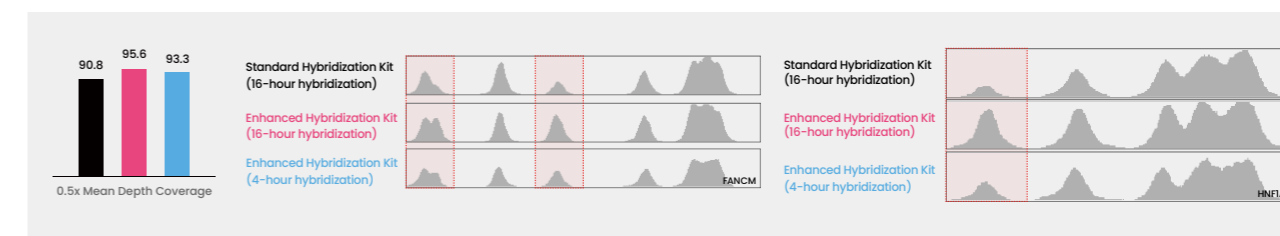


## DESCRIPTION

Celemics Hybridization Enhancer is developed for the hybridization step in the library preparation using Celemics Target Enrichment Kits (Enhanced Hybridization Kit). It enables 4 hours of hybridization with no compromise on the performance quality.

## PERFORMANCE

Improved uniformity and coverage with Hybridization Enhancer





# CeleMag™ Clean-up Bead



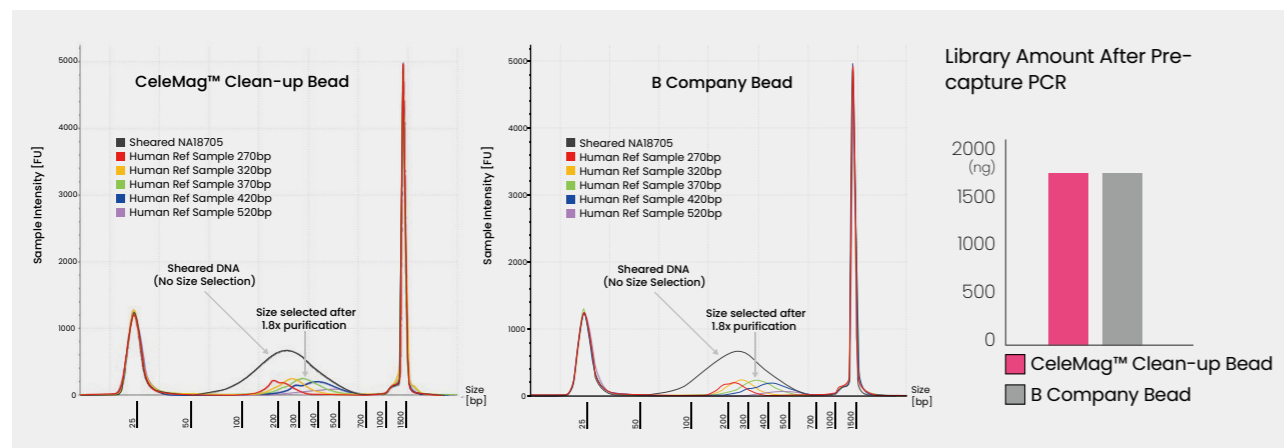
## DESCRIPTION

The CeleMag™ Clean-up Bead utilizes unique magnetic bead-based chemistry enabling a simple, flexible and reproducible workflow for purification and size selection of nucleic acids.

## KEY FEATURES

1. Market leading purification and size selection efficiency
2. Highly optimized with Celeemics Target Enrichment Kits
3. Consistent size selection with flexibility

## PERFORMANCE



CeleMag™ Clean-up Bead provides highly comparable performance to competitor product in size selection workflows, achieving consistent DNA size distributions and yielding desired library sizes. CeleMag™ Clean-up Bead also provides equivalent NGS Library preparation recovery efficiency compared to competitor product.

# CLM Polymerase



## DESCRIPTION

The role of polymerase is critical in NGS process. Due to the complexity of the library, high performance polymerase is required for high uniformity and yield. As a service provider, Celeemics has been providing CLM polymerase with market-leading performance, exhibiting high yield and accuracy with minimized PCR bias. The product includes all reaction components for PCR. Contact us for more information.



# CeleMag™ Streptavidin Bead



## DESCRIPTION

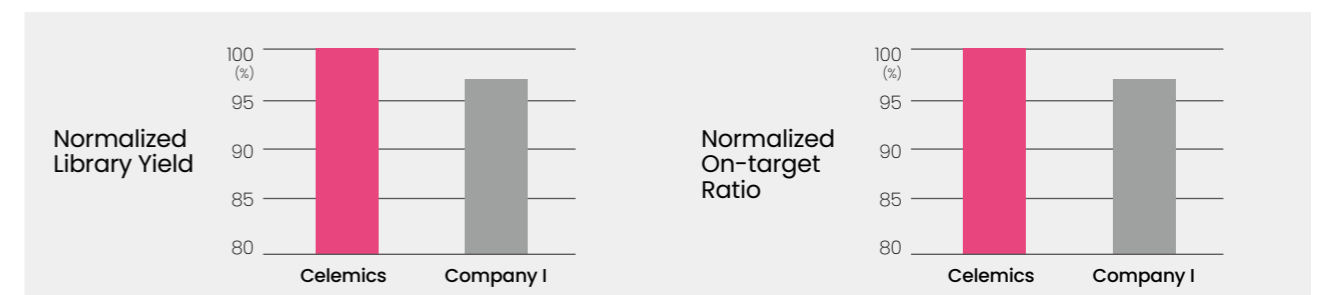
The CeleMag™ Streptavidin Bead selectively isolates biotinylated ligand, using binding properties of biotin. Its high performance enables isolating targeted genes that are bound to probes and minimizes DNA loss during the target enrichment process.

## KEY FEATURES

1. High biotin-streptavidin binding capacity
2. Superior target enrichment efficiency

## PERFORMANCE

Superior performance of CeleMag™ Streptavidin Bead compared to competitor product



# Bioinformatics Software

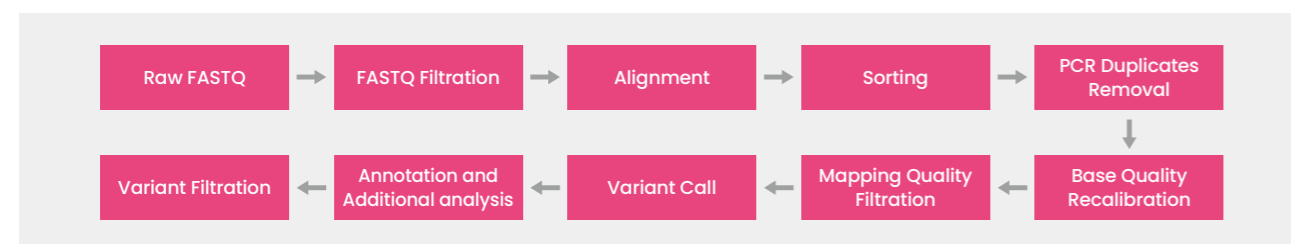
## DESCRIPTION

As a part of Celemics' intellectual property, a unique NGS bioinformatics pipeline is developed to process and analyze massive amounts of genomic data into a readable format with clinically significant biomarkers obtained through Next Generation Sequencing.

## KEY FEATURES

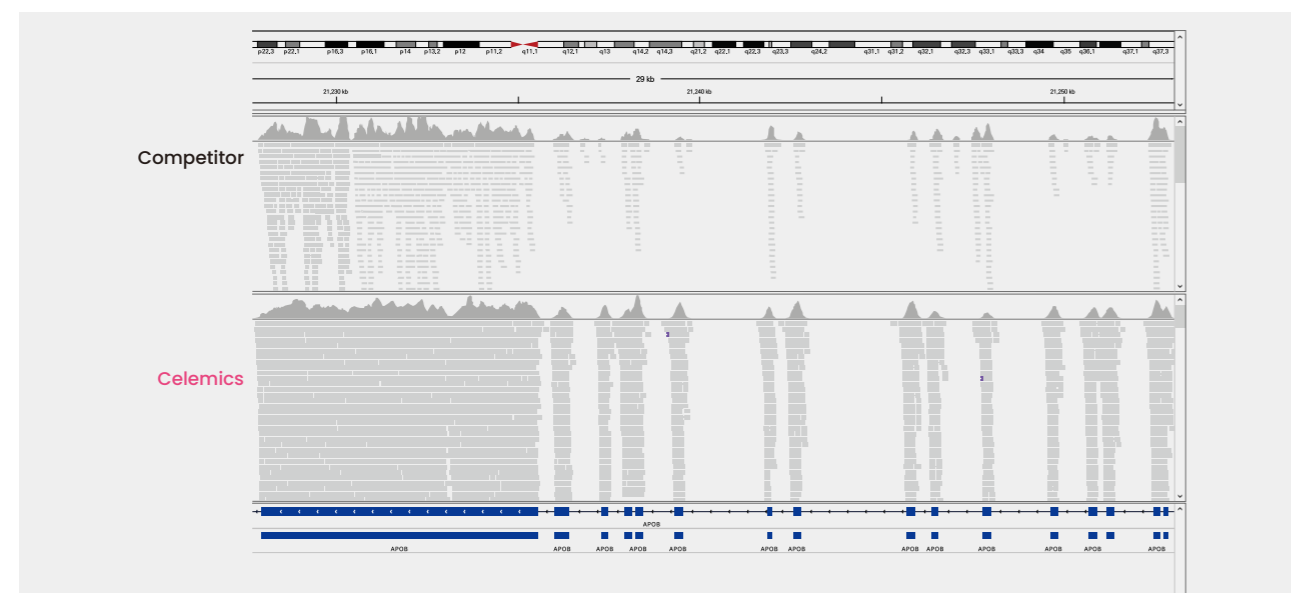
1. Built-in service for all panel kits and services
2. Provides FASTQ to VCF and interpretation
3. Robust pipelines for detecting and analyzing all types of variants including SNV, Indel, CNV, Rearrangements, MSI, TMB, and ultra-low variants

## NGS DATA ANALYSIS PIPELINE

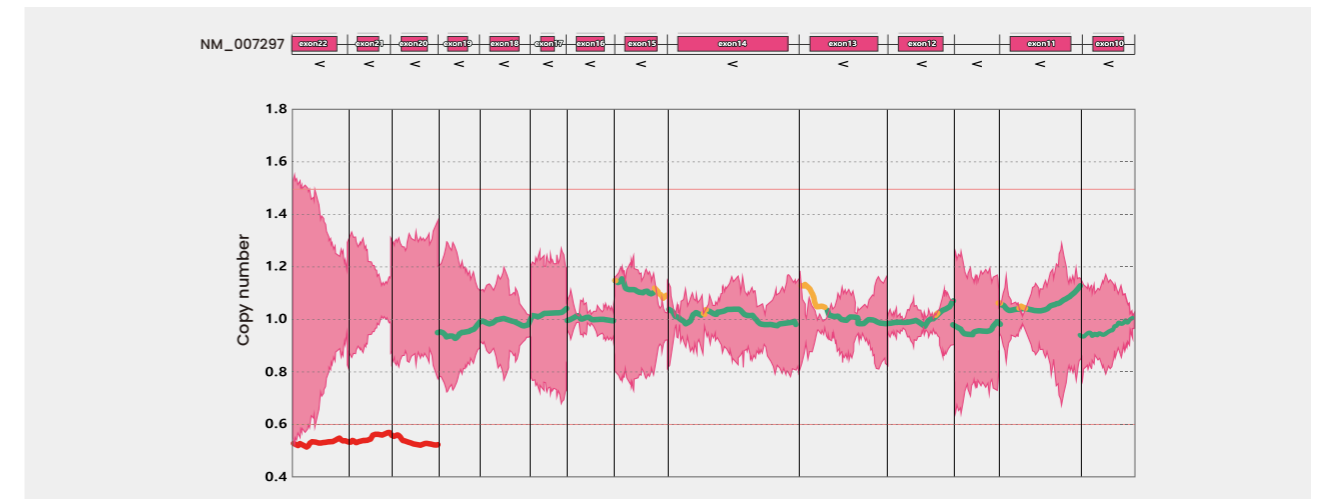


## EXAMPLES OF BIOINFORMATICS ANALYSIS REPORT

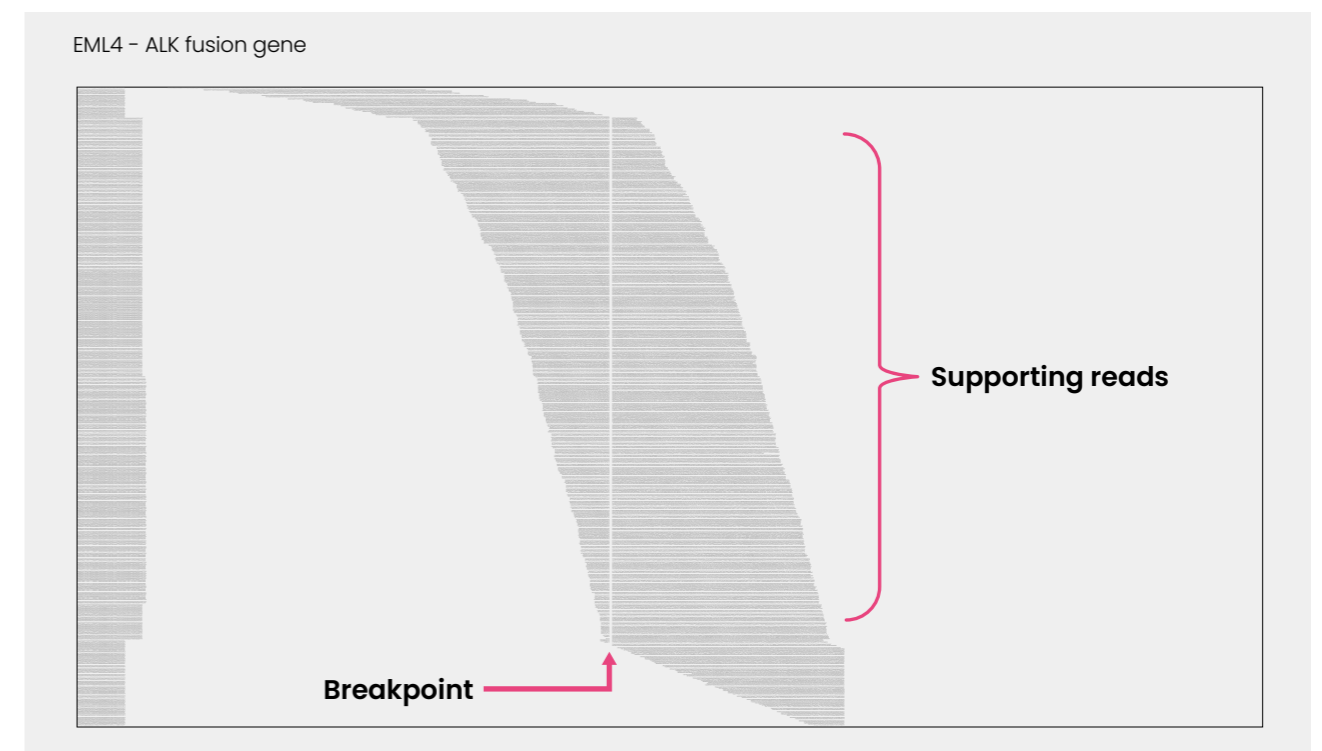
### Comparison of IGV results from Celemics and competitor product



## CNV Analysis Example – Deletion



## Gene rearrangement analysis with FFPE samples





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