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# **Celemics Whole Exome Sequencing Panel**

## **Key Features**

- Complete Whole Exome Coverage
- Superior Performance in the Market
- Gene Add-On Service
- FASTQ to Clinical Interpretation Capability
- Rapid Same-Day Workflow
- No Need for Heavy Instruments
- Complete Walkaway Automation
- Flexible Integration with NGS Sequencers

Celemics Whole Exome Sequencing (WES) Panel is a comprehensive solution that covers all target regions of major WES panels available in the market.

With a target size of 37.1 Mb, the panel does not compromise performance in terms of coverage and uniformity, enabling highly efficient and cost-effective sequencing of the human whole exome. The panel coverage spans across exon regions from RefSeq, CCDS, and GENCODE.

Celemics panels also perform well against hard-to-capture regions such as GC-rich regions. Regarding spike-in options, we have the ability to customize our WES panel according to your specific needs by including mitochondrial or intronic regions upon request.

The panel is also fully supported by Celemics Analysis Service (CAS), our end-to-end bioinformatics solution.

## **Complete Whole Exome Coverage**

What differentiates the Celemics Whole Exome Sequencing Panel from other WES panels in the market? Most researchers look for complete coverage especially when it comes to WES. However, most WES panels in the market vary in their target regions and some compromise with coverage, even deleting hard-to-capture regions in order to enhance performance. Celemics has developed a WES panel that covers the regions of all four major WES panels in the market, spanning the coding regions from RefSeq, CCDS and GENCODE. The Celemics WES Panel provides the most comprehensive coverage of protein-coding regions, thereby enabling marker discovery to diagnostics.



### Figure 1. Complete Whole Exome

The Celemics WES Panel covers the target regions of all major whole exome panels in the market, which include WES panels from company A, company I and company T. Among the regions that Company A and T failed to cover from their own target regions, Celemics covers an additional 60 Kb and 306 Kb of Company A and T respectively (data not shown). Most of these regions are challenging to capture due to GC-rich and repeated sequences. Through Celemics' optimized probe design and panel synthesis technology, the Celemics Whole Exome Panel is able to successfully cover the challenging regions that other company products struggle with.



Company A	p- oop
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Celemics	
	p- 60)
Company T	
company	
Sequence 🗕	
Gene	PAFAH181
Celemics Target	PAFAH1B1
ClinVar	Likely_pathogenic Pathogenic Pathogenic Pathogenic Pathogenic Pathogenic Pathogenic Pathogenic Pathogenic Pathogenic
COSMIC	COSM6874584 COSM1286949 COSM436201 COSM6209041
dbSNP	325844564 rs1306889751 rs587784252 rs768437076 rs776953307 rs113994198 rs587784254 rs765448987 rs1226161297 rs58778

#### Figure 2. Superior Capture Performance and Coverage

A mutation in PAFAH1B1 causes Isolated Lissencephaly Sequence (ILS) & Miller-Dieker syndrome. While other competitor panels fail to capture the A-T rich regions in this gene, the Celemics WES Panel successfully covers the region.



C.



#### Figure 3. Exhaustive Coverage for Each Gene

The Celemics WES Panel covers each gene with thorough coverage in comparison to competitor products. The bar graphs indicate the percentage of genes that are covered at (A) 20X depth and (B) 30X depth. The data from the three panels were downsampled to 5.4 Gb. (C) The IGV figure demonstrates the superior coverage performance of the Celemics Whole Exome Panel against the TGID1 gene compared to other competitor products.



## **Exceptional Target Capture Performance**

Celemics provides market-leading target capture performance due to probe design and reagent optimization technology. Despite some companies who resort to masking the hard-to-capture regions (such as GC- or AT-rich regions and homologous regions) or completely omit the regions from their target in order to enhance the result quality, Celemics provides both high coverage and on-target ratio without reducing the number of target regions. With Celemics' proprietary technology, the Celemics WES Panel captures regions that no other companies could capture with quality coverage and uniformity. The all-around performance of Celemics' WES panel allows for highly sensitive, cost-effective and time-saving sequencing of the whole exome.





A. Standard Deviation against GC Regions





#### Figure 4. Superior Performance in the Market

Celemics WES Panel shows exceptional performance compared to other competitor products when measured by (A) on-target read ratio, (B) 0.2x mean depth coverage uniformity (higher the better), and (C) Fold-80 base penalty (lower the better). Third-party laboratories (Certified Service Providers) conducted a comparison study between the Celemics WES Panel, Company A and Company T panels. Reference materials NA12878, NA12891, and NA12892 were used with same amount. Illumina instruments were used for the sequencing. The data from the three panels were downsampled to 5.4 Gb.



Figure 5. Exceptional Uniformity across Low and High GC Regions

(A) The Celemics WES Panel demonstrates minimal deviation, yielding 0.166
standard deviations (lower the better) across GC-rich and AT-rich regions in comparison to competitor products yielding 0.199 and 0.356 standard deviations.
(B) The bar graphs shown in different GC ratios also illustrate the consistent uniformity of Celemics WES Panel in comparison to the competitor products.



## **Simple Gene Add-On Customization**



## **Rapid Same-Day Workflow**

Although hybridization capture has great advantages including minimized bias, stable and reliable data results from a variety of sample types, the complexity of the workflow and the long prep time have been obstacles to the users. Celemics has developed a new workflow and have incorporated it into our WES package to significantly simplify the process and reduce the experiment time. The conventional method took 2-3 days to complete one sequencing experiment. Now with Celemics' newly developed method, the whole experiment and the NGS run can be started on the same day.



### Figure 7. Newly Developed Same-Day Workflow

The figure demonstrates that Celemics has significantly reduced the time for performing Whole Exome Sequencing from the conventional 20 hours to 5 hour minimum workflow.



## **Full Bioinformatics Capability: FASTQ to Clinical Interpretation**



CAS (celemics Analysis Service) Workflow for WES Panel

#### **Figure 8. Bioinformatics Support through CAS**

CAS (Celemics Analysis Service) provides easy data transmission by single-click and automated uploads. Due to the complete support from Celemics bioinformatics experts, CAS does not require separate third-party bioinformaticians. CAS also supports real-time troubleshooting throughout primary to tertiary analysis and client-specific customization.

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Patient	Haplotype Patient	Sample ID	HaplotypeCaller	Test	CEL-PAN405	Standard Drugs					
inder	Female	Specimen Type	FFPE Block	Type	Full Report	Alectinib	Enhanced Response				
	60 Years	Block ID		Sample Collected	28-Feb-2021	Alectilib		ker(s)			
N		Tumor Content		Sample Received	01-Mar-2021	Markers					
ferred by	Dr. Lea-Bennett	Specimen Site		Report Generated	03-Mar-2021			_			ed non-sma
ferring Institution	The University of Oklahoma Health Sciences Center					ALK		_			10 M - 0
linical In	diantiana					Evidence Details		_	or BRAF p.Val		28475456
Non-email cell lung carcinoma (NSCLC) Note A variant was detected in ALK and its therapeutic implications are summarized below.				below.		Alectinib is a second generi small cell lung cancer (NSC positive NSCL c patients in to alectinib and critozinib ir in terms of cinical benefit v vs. 48.7% with crizotinib), e treated group and a respon	Addetable is a second protection QAX (inhibitor approved for the treatment of patients with ALK-rearrangement politikin non- sensitivity and paraneter (BSCC). Addetable has whose multitable settial in bio facultable pre-severate data batterine train VAX, positive MSCLC patients' in multiple clinical studies (12,10,31,41,51,81). The phase III ALX study comparing response to adjective and orizotable in analysis of the ALX study Comparing response to a studies in the study of the adjective and the study of the adjective and the study of the adjective ad			ith ALK-positive cal studies have n treatment with a upon crizotinib positive NSCLC YFS). In patients (the patients (4), eported an ORR ORR of 100% in cancer indicates	ancer and of <u>36718</u> ] sistance to Fi VID: <u>277808</u> sr. <i>N. Engl.</i> ell lung can sancer. <i>Eur.</i>
Reported Variants						Marker Details					
						EML4 (NM_019063, E	kon 1-6) : ALK (NM_004304, Exon 20-29)	Enhanced			
/IL4 <sup>5</sup> : ALK <sup>3</sup>	translocation	ocation Tier-I				Oncogenicity Established b	Oncogenicity Established by in-vitro Studies				
EML4 <sup>S</sup> : ALK <sup>S</sup> translocation Tier-I				EML4-ALK translocation is represents 2-7% of the lur chromosome 2 and has con	EML4-ALK translocation is the most common aberration of the ALK gene, which is seen exclusively in lung cancer and represents 2-7% of the lung cancers cases (7). The fusion gene results from an inversion event on the short arm of chromosome 2 and has constitutive ALK kinase activity [8]. Till date, over 11 EML4-ALK fusion variants have been identified						
*kerted categorization tens as per the AMP guidatines Summary for Standard Drugs						EML4 (NM_001145076 Oncogenicity Established by EML4-ALK translocation is represente 3.7% of the lute	In lang cancer, composed of varying EMA transcript length tasks with exon 20 of ALK [9]. EMIA (1MC,01148075, Exon 1-5); ALK (LMC, MO,04304, Exon 20-239) Oncogenicity Established by In-vitro Studies EMIA-ALK translocation is the most common alternation of the ALK gene, which is seen exclusively in lung cancer and			lung cancer and he short arm of e been identified	
nuas NOT INF	CATED Based on	FDA Mandated/ Gu	ideline Recommended	Markers (see n	age 3 for details)	chromosome 2 and has con	stitutive ALK kinase activity [8]. Till date, over 11 EML4-ALK fusion variants have been identified	Pemetrexed,			
herapy		Tested Marker(s)		Relevant N	arker(s)	in lung cancer, composed o	f varying EML4 transcript length fused with exon 20 of ALK [9].			lung cancer and	
abrafenib		BRAF		None		Drug Description		Irinotecan,		e been identified	
metinib		BRAF		None		An orally available inhibitor	of the receptor tyrosine kinase anaplastic lymphoma kinase (ALK) with antineoplastic activity, in binds to and inhibits ALK kinase. ALK fusion proteins as well as the patekeeper mutation		Non-detection		
atinib		EGFR		None		ALKL1196M known as one o	f the mechanisms of acquired resistance to small-molecule kinase inhibitors. The inhibition leads				
comitinib		EGFR		None		to disruption of ALK-mediat belongs to the insulin recep and gene rearrangements a	to disruption of ALX-mediated signaling and versitually inhibits tumor cell growth in ALX-overexpressing tumor cells, ALX betrage to the insulin receptor superfamily and pages in important (see in nervous system development, ALX dysregulation development, and the insulin receptor superfamily and page in the insuling and the insuling and the insuling and Source: The National Cancer Institute's Cancer Drug Information			ibits ALK kinase d EGFR kinase,	
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### **Figure 9. Clinical Interpretation Report**

Celemics provides robust clinical interpretation services through a CAP-accredited partner that combines bioinformatics algorithms, public data from external sources/knowledge databases, visualization interfaces and reporting capabilities. The report includes pathogenicity and drug associated information.



## **No Need for Heavy Instruments**

In order to perform Library Preparation prior to Target Enrichment and Sequencing, it is often required to have heavy instruments (such as a vacuum concentrator or sonicator, etc.), which are barriers against complete automation. Even with an automation protocol, using these heavy instruments is inevitable and is often burdensome to the users. Celemics has successfully eliminated the need for heavy instruments by substituting them with a more convenient solution of enzymes and beads. After rigorous validation that consistently showed reliable performance, we have optimized this workflow to enable the benefit of a complete walkaway solution.



# Flexible Integration with NGS Sequencers & Complete Walkaway Automation



### Figure 11. Compatibility with NGS Sequencers and Automation

The Celemics WES Panel is seamlessly integrated with all NGS instruments from Illumina, MGI, and Ion Torrent. Since there are no heavy instruments required, the experiment can be carried out with complete automation.

# **Ordering Information**



Produc	Applied Platform			Package Option	Product Unit	
Category	Sub-category					
Ready-to-use panels	Whole Exome Sequencing	Illumina	lon Torrent	MGI	1) All-in-one Package 2) Standard Package 3) Target Enrichment Package	1) Reaction basis: 16, 48, 96 rxn 2) Sample basis: 16, 96, and more options *Pre-capture pooling options: 4, 8, 12

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