



Pillar® ONCO/Reveal™ Custom Panels

ONCO/Reveal Custom Panels are robust NGS assays that interrogate a customer-defined set of gene regions of interest. Custom assays are designed using the ampPD™ intelligent primer design platform. All Pillar assays feature proprietary Stem-Loop Inhibition-Mediated amplification (SLIMamp®) technology, a tiled amplicon-based library prep chemistry for efficient single-tube target enrichment. Variant calls are made using the Pillar Variant Analysis Toolkit (PiVAT®) bioinformatics pipeline software.

Disease states	Oncology, genetic diseases, ophthalmology, familial hypercholesterolemia, thalassemia, metabolism, carrier screening, NIPT, PGx, IBD, and more
Sample types	Solid tumors, liquid biopsy, myeloid, germline, FFPE, blood, plasma, serum, other biofluids
Variant types	Fusions, CNV, SNV, indels
Coverage	Hotspots or full gene CDS (overlapping amplicons) from single-tube amplification
PiVAT	Cloud or local installation
Support	Field application scientist team

Simple NGS library prep workflow

Maintain control of samples and results with single-tube, tiled amplification that can be performed in-house by any NGS lab

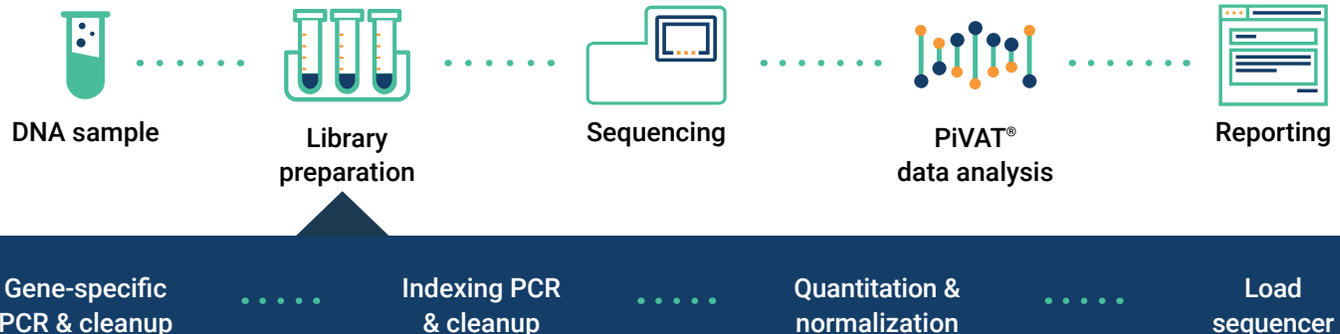
Sensitive and robust chemistry

Achieve variant detection as low as 1% VAF⁺ without UIDs[‡] and <1% with UIDs even with limited DNA input or poor sample quality

Reduced fully-loaded lab costs

Improve lab efficiency and reduce “no calls”, repeat testing, and difficult interpretation decisions

Simple, one-day workflow



⁺VAF, variant allele frequency; [‡]UID, unique ID; also known as unique molecular ID (UMI)

Panel specifications

Enrichment chemistry	Multiplex PCR using tiled amplicons
Number of pools	1 pool
Number of amplicons	Up to 5,000
Variant types	SNVs, indels, fusions, CNVs
Limit of detection (LoD)	1%-2% VAF without UIDs (10ng input DNA) or <1% VAF with UIDs (20ng input DNA)
Input DNA range	1ng to 80ng (10ng recommended)
Sample types	Genomic DNA, Cell-free DNA, FFPE DNA, fresh frozen tissue, RNA
Design coverage	>90%
Mapping rate*	>90%
% on-target aligned reads*	>90%
Coverage uniformity (% targets with >0.2X mean coverage)*	>90%
Total assay time (from sample to sequencer)	<8 hours
Interactive design and functional testing	Yes
Designer; Analysis software	ampPD; PiVAT
Sequencing platforms	Illumina®, MGISEQ™, and Ion Torrent™
Configurable kit contents	Primer pool, indexing primers, UDG, beads

*Only guaranteed with testing and optimization

Ordering information

Select one of the index kit options listed below to go with your custom panel.

Pillar Index Kit options	Part number
Pillar Kit A, indices PI501-8, PI701-4 (32 combinations)	IDX-PI-1001-96
Pillar Kit B, indices PI501-8, PI705-8 (32 combinations)	IDX-PI-1002-96
Pillar Kit C, indices PI501-8, PI709-12 (32 combinations)	IDX-PI-1003-96
Pillar Kit D, indices PI501-8, PI701-12 (96 combinations)	IDX-PI-1004-192
Pillar Kit E, indices PI501-8, PI701-12 (96 combinations)	IDX-PI-1004-384

Ion Torrent Index Kit options	Part number
Pillar Biosciences IonXpress Indexing Kit A (24 combinations) <i>Includes ION Common Index</i>	IDX-TI-1001-96

MGI Index Kit options	Part number
Pillar Biosciences MGI Indexing Kit A (24 combinations) <i>Includes BGI Common Index</i>	IDX-BI-1002-96

Design and delivery timelines



without testing
and optimization



with testing
and optimization

Learn more at
pillar-biosciences.com

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PILLAR
BIOSCIENCES

Pillar Biosciences, Inc.
9 Strathmore Rd
Natick, MA 01760
(800) 514-9307
info@pillar-biosciences.com