VARIANTPlex-HT Pan Solid Tumor

Description

The VARIANT*Plex*-HT Pan Solid Tumor panel is a balanced pool of gene-specific primer (GSP) oligonucleotides that is optimized for use with VARIANT*Plex*-HT reagents and molecular barcode (MBC) adapters to produce targeted NGS libraries. This product insert should be used in conjunction with VARIANT*Plex*-HT protocol for Illumina® (RA-DOC-058).

VARIANT*Plex*-HT Pan Solid Tumor contains **9878** GSPs targeting **185** genes commonly mutated in solid tumors as well as microsatellite instability (**MSI**) and tumor mutational burden (**TMB**) status.

Description	Part number	Storage
VARIANT <i>Plex</i> -HT Pan Solid Tumor GSP1, 24 reactions or VARIANT <i>Plex</i> -HT Pan Solid Tumor GSP1, 96 reactions	SA20196241 or SA20196961	−20°C ± 10°C
VARIANT <i>Plex</i> -HT Pan Solid Tumor GSP2, 24 reactions or VARIANT <i>Plex</i> -HT Pan Solid Tumor GSP2, 96 reactions	SA20196242 or SA20196962	— -20 C ± 10 C

Required reagent volumes

Protocol reference	Protocol step	Reagent	Volume per reaction (µL)
Α	Ligation Step 2 Elution	5mM NaOH	20
В	First PCR	VARIANT <i>Plex</i> -HT Pan Solid Tumor GSP1	8
С	First PCR	10mM Tris-HCl pH 8.0	18
D	First PCR	Purified PCR1 eluate	16
Е	Second PCR	VARIANT <i>Plex</i> -HT Pan Solid Tumor GSP2	8

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Recommended PCR cycling

	Step	Temperature (°C)	Time	Cycles
	1	95	3 min	1
	2	95	30 sec	
	3	58	10 sec	 15
First PCR reaction	4	60	10 min (100% ramp rate)	
	5	72	3 min	1
	6	4	Hold	1
	1	95	3 min	1
Second PCR reaction	2	95	30 sec	
	3	58	10 sec	
	4	65	10 min (100% ramp rate)	
	5	72	3 min	1
	6	4	Hold	1

[†]The number of PCR2 cycles may be decreased if you regularly experience library yields greater than 200 nM.

Recommended reads and multiplexing

VARIANT*Plex*-HT Pan Solid Tumor libraries should be sequenced to a minimum of **25M** reads. Starting read depth recommendations for standard profiling may be adjusted based on user needs.

Archer™ Analysis settings

Sequencing data should be processed using Archer Analysis (v7.2, or greater). The VARIANT*Plex*-HT Pan Solid Tumor panel works with selection of the *SNV/Indel, Copy Number Variation, Structural Variation, MSI, and TMB* pipelines, found under the *DNA* Input Type. The recommended TMB settings in Archer Analysis are *High Threshold for TMB* = 20 and *Low Threshold* for TMB = 5 for VARIANT*Plex* Pan Solid Tumor. As with most Analysis settings, these may be adjusted based on user needs. Selection of the DNA Target Coverage pipeline is optional (see the Archer Analysis User Guide for more details on setting up your analysis).

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Archer Analysis processing of VARIANT*Plex*-HT Pan Solid Tumor libraries requires a one-time upload of the Panel GTF. When performing DNA Target Coverage analysis, users must also select a Region of Interest BED file. Users may optionally add a Targeted Mutations VCF file for targeted SNV/Indel detection. Files can be obtained by contacting archer-tech@idtdna.com

Assay targets

Integrative genomic biomarkers MSI, TMB

Genes

The following genes in the panel have full coding sequence coverage for SNV/Indel analysis. Coding sequence was determined using NCBI RefSeq transcripts†.

9 1		J				
ABL1	BMPR1A	CTNNB1*	FGF19	HNF1A	MDM4	NOTCH2
ACVR1	BRAF*	DAXX	FGFR1	HRAS	MED12	NOTCH3
AKT1	BRCA1*	DDR2	FGFR2	IDH1	MEN1	NOTCH4
AKT2	BRCA2*	DDX3X	FGFR3	IDH2	MET*	NPM1
AKT3	BRIP1	DICER1	FGFR4	JAK1	MLH1	NRAS
ALK	CCND1	EGFR	FH	JAK2	MPL	NTRK1
APC	CCND2	EIF1AX	FLCN	JAK3	MRE11A	NTRK2
AR	CCND3	EP300	FLT1	KDM6A	MSH2	NTRK3
ARID1A	CCNE1	EPCAM	FLT3	KDR	MSH3	PALB2
ARID1B	CDH1	ERBB2	FLT4	KEAP1	MSH6	PBRM1
ARID2	CDK12	ERBB3	FOXA1	KIT*	MTOR	PDGFRA*
ATM*	CDK4	ERBB4	FOXL2	KLF4	MUC16	PIK3CA
ATR	CDK6	ERCC1	FUBP1	KMT2C	MUTYH	PIK3CB
ATRX	CDKN2A	ERCC2	GNA11	KMT2D (MLL2)	MYC	PIK3R1*
AURKA	CDKN2B	ESR1	GNAQ	KRAS	MYCN	PLCB4
B2M	CHD1	EZH2	GNAS	LZTR1	NBN	PMS2
BAP1	CHEK1	FANCA	H3F3A	MAP2K1 (MEK1)	NF1	POLD1
BARD1	CHEK2*	FANCI	H3F3B	MAP2K2 (MEK2)	NF2	POLE
BCOR	CIC	FANCL	HIST1H3B	MAP3K1	NKX2-1	PPP2R1A

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BLM	CSF1R	FBXW7	HIST1H3C	MDM2	NOTCH1*	PPP2R2A
PRKD1	RAD51C	RICTOR	SETD2	SRC	TP53	VHL
PTCH1	RAD51D	RNF43	SF3B1	SRSF2	TP63	XRCC2
PTEN	RAD54L	ROS1	SMAD2	STAG2	TRAF7	XRCC3
PTPN11	RAF1	SDHA	SMAD4	STK11	TSC1	
RAD50	RB1	SDHB	SMARCA4	SUFU	TSC2	
RAD51	RET*	SDHC	SMARCB1	TERT	TSHR	
RAD51B	RHOA	SDHD	SMO	TGFBR2	U2AF1	

[†]Contact archer-tech@idtdna.com for the panel target file to view complete list of targeted regions.

Genes targeted for CNV

AKT1	BRIP1	ERBB2	JAK2	PDGFRA
AKT2	CCND1	ERBB3	KDR	PIK3CA
AKT3	CCND2	ERCC1	KEAP1	PIK3CB
ALK	CCND3	ERCC2	KIT	PTEN
AR	CCNE1	ESR1	KRAS	RAD51B
ATM	CDK12	FGF19	MDM2	RAF1
AURKA	CDK4	FGFR1	MDM4	RET
BARD1	CDK6	FGFR2	MET	RICTOR
BRAF	CHEK1	FGFR3	MYC	STK11
BRCA1	CHEK2	FGFR4	MYCN	TERT
BRCA2	EGFR	FLT3	NRAS	

Please contact archer-tech@idtdna.com to inquire about enabling additional genes for CNV detection.

SNPs and sites targeted for sample tracking

rs560681	rs430046	rs987640	rs10776839	rs12393891
18000001	18430046	18907040	1810776639	1812393691
rs740598	rs8078417	rs6444724	rs6530357	chrX:4429309
rs1498553	rs9951171	rs6811238	rs5971553	chrX:11314433
rs10773760	rs576261	rs13182883	rs5953060	chrY:6738552
rs1058083	rs1109037	rs214955	rs6524626	chrY:19490214
rs4530059	rs1523537	rs321198	rs5940270	
rs1821380	rs221956	rs4606077	rs722847	

SNPs may be used in combination to uniquely tag and track samples over time. Contact archer-tech@idtdna.com for further details.

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^{*}Indicates that select regions of this gene are enabled for Structural Variant analysis in the GTF.

Limitations of use

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Revision History

Document Number	Date	Description of change
RA-DOC-065/REV01	October 2023	Initial release.
RA-DOC-065/REV02	November 2023	Updated First and Second PCR cycling conditions to include separate anneal and extended steps.
		Updated branding.

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