Technical Specifications





Intended Use

FoundationOne Liquid CDx is a next generation sequencing based in vitro diagnostic device that analyzes 324 genes. Substitutions and insertion and deletion alterations (indels) are reported in 311 genes, copy number alterations (CNAs) are reported in 310 genes, and gene rearrangements are reported in 324 genes. The test also detects tumor fraction and the genomic signatures blood tumor mutational burden (bTMB) and microsatellite instability high (MSI-H) status. FoundationOne Liquid CDx utilizes circulating cell-free DNA (cfDNA) isolated from plasma derived from the anti-coagulated peripheral whole blood of cancer patients. The test is intended to be used as a companion diagnostic to identify patients who may benefit from treatment with targeted therapies in accordance with the approved therapeutic product labeling. Additionally, FoundationOne Liquid CDx is intended to provide tumor mutation profiling to be used by qualified health care professionals in accordance with professional guidelines in oncology for patients with malignant neoplasms.

A negative result from a plasma specimen does not mean that the patient's tumor is negative for genomic findings. Patients who are negative for genomic findings should be reflexed to routine biopsy and their tumor mutation status confirmed using an FDA-approved tumor tissue test, if feasible.

FoundationOne Liquid CDx is a single-site assay performed at Foundation Medicine, Inc. in Cambridge, MA.



Summary of Analytical Sensitivity and Specificity

Results from our Limit of Detection (LoD) study are shown below, indicating the median variant allele frequency, tumor fraction or unstable loci at which the test has shown 95% probability of detection. Please refer to our product labeling for a list of the 75 genes baited for enhanced sensitivity and complete product specifications.

ALTERATION TYPE	BAIT SET REGION	MEDIAN LIMIT OF DETECTION (LOD)	
Chauk Varianta	Enhanced Sensitivity	0.40% VAF	
Short Variants	Standard Sensitivity	0.82% VAF	
Rearrangements	Enhanced Sensitivity	0.37% VAF	
	Standard Sensitivity	0.90% VAF	
Copy Number Amplification	NA	21.7% TF	
Copy Number Loss	NA	30.4% TF	
MSI	NA	0.8% Unstable loci	
bTMB (component indels)	NA	1.00% VAF	
bTMB (component subs)	NA	1.00% VAF	

VAF = variant allele frequency; TF = tumor fraction

The accuracy of $\ensuremath{\mathrm{WVAF}}$ / $\ensuremath{\mathrm{WTF}}$ have not been analytically validated

In our Limit of Blank study, which evaluated variant calling in healthy donors, 1,735 unique variants were included in the analysis for a total of 137,065 data points. A total of 18 false positives were observed across 4 unique short variants. The LoB was determined to be the ideal value of zero for short variants, rearrangements and CNAs. The false positive rate was shown to be 0% for rearrangements and CNAs and 0.013% (~1 in 8,000) for short variants (substitutions and indels).



FoundationOne Liquid CDx Gene List[†]

As a professional service, FoundationOne Liquid CDx interrogates 324 genes, including 309 genes with complete exonic (coding) coverage and 15 genes with only select non-coding coverage (indicated with an *); **75 genes (indicated in bold) are captured with increased sensitivity** and have complete exonic (coding) coverage unless otherwise noted. The test also detects tumor fraction and the genomic signatures blood mutational burden (bTMB) and microsatellite instability high (MSI-H) status.

ABL1 [Exons 4-9]	ALOX12B	ASXL1	BAP1	BCR* [Introns 8, 13, 14]	BRIP1	CASP8
	AMER1 (FAM123B)	ATM	BARD1		BTG1	CBFB
ACVR1B	APC	ATR	BCL2	BRAF (Exons 11-18.	BTG2	CBL
AKT1 [Exon 3]	AR	ATRX	BCL2L1	Introns 7-10]	BTK	CCND1
AKT2	ARAF	AURKA	BCL2L2	BRCA1 [Introns 2, 7, 8, 12,	[Exons 2, 15]	CCND2
AKT3	[Exons 4, 5, 7, 11, 13, 15, 16]	AURKB	BCL6	16, 19, 20]	C11orf30 (EMSY)	CCND3
				BRCA2	C17orf39 (GID4)	
ALK [Exons 20-29	ARFRP1	AXIN1	BCOR	[Intron 2]	CALR	CCNE1
Introns 18,19]	ARID1A	AXL	BCORL1	BRD4	CARD11	CD22

(FoundationOne Liquid CDx Gene List continued)

CD70	ERBB2	FOXL2	KLHL6	NF1	PPARG	SMAD2
CD74*	ERBB3	FUBP1	KMT2A (MLL)	NF2	PPP2R1A	SMAD4
[Introns 6-8]	[Exons 3, 6, 7, 8, 10, 12, 20, 21, 23, 24, 25]	GABRA6	[Introns 6, 8-11, Intron 7]	NFE2L2	PPP2R2A	SMARCA4
CD79A	ERBB4	GATA3	KMT2D	NFKBIA	PRDM1	SMARCB1
CD79B	ERCC4	GATA4	(MLL2)	NKX2-1	PRKAR1A	SMO
CD274 (PD-L1)	ERG	GATA6	KRAS	NOTCH1	PRKCI	SNCAIP
CDC73	ERRFI1	GNA11	LTK	NOTCH2	PTCH1	SOCS1
CDH1	ESR1	[Exons 4, 5]	LYN	[Intron 26]	PTEN	SOX2
CDK12	[Exons 4-8]	GNA13	MAF	NOTCH3	PTEN PTPN11	
CDK4	ETV4*	GNAQ	MAP2K1 (MEK1)	NPM1		SOX9
CDK6	[Intron 8]	[Exons 4, 5]	[Exons 2, 3]	[Exons 4-6, 8, 10]	PTPRO	SPEN
CDK8	ETV5* [Introns 6,7]	GNAS [Exons 1, 8]	MAP2K2 (MEK2) [Exons 2-4, 6, 7]	NRAS [Exons 2, 3]	QKI	SPOP
CDKN1A	ETV6*	GRM3	MAP2K4	NSD3 (WHSC1L1)	RAC1	SRC
CDKN1B	[Introns 5,6]	GSK3B	MAP3K1	NT5C2	RAD21	STAG2
CDKN2A	EWSR1* [Introns 7-13]	H3F3A	MAP3K13	NTRK1	RAD51	STAT3
CDKN2B	EZH2	HDAC1	MAPK1	[Exons 14, 15, Introns 8-11]	RAD51B	STK11
CDKN2C	[Exons 4, 16, 17, 18]	HGF	MCL1	NTRK2	RAD51C	SUFU
CEBPA	EZR* [Introns 9-11]	HNF1A	MDM2	[Intron 12]	RAD51D	SYK
CHEK1	FAM46C	HRAS	MDM4	NTRK3 [Exons 16, 17]	RAD52	TBX3
CHEK2	FANCA	[Exons 2, 3]	MED12	NUTM1*	RAD54L	TEK
CIC	FANCC	HSD3B1	MEF2B	[Intron 1]	RAF1 [Exons 3, 4, 6, 7,	TERC* {ncRNA}
CREBBP	FANCG	ID3	MEN1	P2RY8	10, 14, 15, 17, Introns 4-8]	TERT* {Promoter}
CRKL	FANCL	IDH1 [Exon 4]	MERTK	PALB2	RARA	TET2
CSF1R	FAS	IDH2	MET	PARK2	[Intron 2]	TGFBR2
CSF3R	FBXW7	[Exon 4]	MITF	PARP1	RB1	TIPARP
CTCF	FGF10	IGF1R	MKNK1	PARP2	RBM10	TMPRSS2* [Introns 1-3]
CTNNA1	FGF12	IKBKE	MLH1	PARP3	REL	TNFAIP3
CTNNB1	FGF14	IKZF1	MPL	PAX5	RET [Introns 7, 8, Exons 11,	TNFRSF14
[Exon 3]	FGF19	INPP4B	[Exon 10]	PBRM1	13-16, Introns 9-11]	TP53
CUL3	FGF23	IRF2	MRE11A	PDCD1 (PD-1)	RICTOR	TSC1
CUL4A	FGF3	IRF4	MSH2	PDCD1LG2 (PD-L2)	RNF43	
CXCR4		IRS2	[Intron 5]	PDGFRA	ROS1	TSC2
CYP17A1	FGF4	JAK1	MSH3	[Exons 12, 18, Introns 7, 9, 11]	[Exons 31, 36-38, 40, Introns 31-35]	TYRO3
DAXX	FGF6	JAK2	MSH6	PDGFRB	RPTOR	U2AF1
DDR1	FGFR1 [Introns 1, 5, Intron 17]	[Exon 14]	MSTIR	[Exons 12-21, 23]	RSPO2*	VEGFA
DDR2		JAK3 [Exons 5, 11, 12,	MTAP	PDK1	[Intron 1]	VHL
[Exons 5, 17, 18]	FGFR2 [Intron 1, Intron 17]	13, 15, 16]	MTOR [Exons 19, 30, 39, 40,	PIK3C2B	SDC4* [Intron 2]	WHSC1
DIS3	FGFR3	JUN	43-45, 47, 48, 53, 56]	PIK3C2G	SDHA	WT1
DNMT3A	[Exons 7, 9 (alternative designation exon 10),	KDM5A	MUTYH	PIK3CA Exons 2, 3, 5-8, 10, 14,	SDHB	XPO1
DOT1L	14, 18, Intron 17]	KDM5C	MYB* [Intron 14]	19, 21 (Coding Exons 1, 2, 4-7, 9, 13, 18, 20)	SDHC	XRCC2
EED	FGFR4	KDM6A	MYC	PIK3CB	SDHD	ZNF217
EGFR [Introns 7, 15, 24-27]	FH	KDR	[Intron 1]	PIK3R1	SETD2	ZNF703
EP300	FLCN	KEAP1	MYCL (MYCL1)	PIM1	SF3B1	
EPHA3	FLT1	KEL	MYCN	PMS2	SGK1	
EPHB1	FLT3 [Exons 14, 15, 20]	KIT [Exons 8, 9, 11, 12, 13, 17,	MYD88 [Exon 4]	POLD1	SLC34A2*	
EPHB4		Intron 16]	NBN	POLE	[Intron 4]	
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^{*}Current as of August 2020. Please visit foundationmedicine.com for the most up-to-date gene list.