HEVA pro

Hereditary and somatic variants profiling in Breast and Ovary Cancer

HEVA pro

4bases HEVA pro is kit for the identification of mutations in genes related to Breast and Ovary cancer, Familial adenomatous polyposis (FAP) and hereditary nonpolyposis colorectal cancer (HNPCC), two syndromes of colorectal cancer predisposition, inherited in an autosomal dominant fashion.

HEVA pro allows the search for variants linked to a defect in homologous recombination repair (HRD), similarly to pathogenic variants in *BRCA1* or *BRCA2*. The defect in HR causes error-prone DNA repair pathways in the cell, resulting in increased genomic instability, which may be responsible for susceptibility to poly- (ADP) -ribose polymerase (PARPis) inhibitors.

HEVA pro is a kit for the analysis of 50 genes through a molecular protocol based on Next Generation Sequencing (NGS) technologies. The kit is validated for germline and somatic analysis (SNPs, indels, CNVs) of DNA extracted from blood or body tissues (fresh, frozen, FFPE, FNA) samples. HEVA pro kit contains all reagents required for the preparation of the capture of specifically designed probes and for the NGS analysis using Illumina sequencers.

TECHNOLOGY

The HEVA pro kit is part of a DNA-to-variant solution that offers streamlined content, easy-to-perform library preparation, push-button sequencing systems, and simplified data analysis.

WORKFLOW

Library preparation follows a straightforward, capture-based protocol that can be completed in as little as 36 hours, with < 3 hours hands-on time. Resulting libraries can be normalized, pooled, and then loaded on to a flow cell for sequencing. Prepared libraries are sequenced on any compatible Illumina sequencers.

VALIDATION

To demonstrate assay capabilities, clinical samples were run in a clinical setting. DNA quality and quantity of the libraries prepared were verified using Qubit and Agilent Bioanalyzer.

Profile of the prepared libraries in Figure .

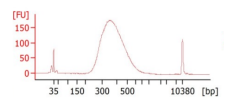


Figure 1. HEVA PRO Bioanalyzer library profile

Table 1: List of genes in HEVA pro

MSH6	FAM175A	LZTR1
MUTYH	MRE11	CTNNB1
NBN	PIK3CA	KRAS
PALB2	PMS2CL	ERBB2
PMS2	XRCC2	ALK
PTEN	NF1	EGFR
RAD50	SPRED1	RB1
RAD51C	MEN1	CDKN2A
RAD51D	RET	CDK4
STK11	SMARCB1	FLCN
TP53	NF2	RUNX1
WRN	PTCH1	CDKN1C
	NBN PALB2 PMS2 PTEN RAD50 RAD51C RAD51D STK11 TP53	MUTYH MRE11 NBN PIK3CA PALB2 PMS2CL PMS2 XRCC2 PTEN NF1 RAD50 SPRED1 RAD51C MEN1 RAD51D RET STK11 SMARCB1 TP53 NF2 WRN PTCH1

SAMPLE PER RUN

Instrument	Samples per run*	
	Germline	Somatic
iSeq i1 100 Kit (300-cycles)	16	0
MiSeq Micro Kit v2 (300-cycles)	16	0
MiSeq Kit v2 (300-cycles)/(500-cycles)	56	0
MiSeq Kit v3 (600-cycles)	56	0
MiniSeq Mid Output Kit (300-cycles)	32	0
MiniSeq High Output Kit (300-cycles)	96	8
NextSeq 550 Mid-Output Kit	536	32
NextSeq 550High-Output Kit	1656	96

*the estimated maximum number of samples per cartridge assumes a reading depth of 300x for the germline and 5000x for the somatic. The optimal number of samples can be empirically estimated on the local setup.

The volume present in the kit is calculated to allow the subdivision into multiples of 8 analysis sessions. Dividing the kit in different ways decrease the total number of tests that can be performed.

ORDERING INFORMATION

Product**	REF	
HEVA pro	C3040-16X (16 test)	
HEVA pro	C3040-96X (96 test)	
UDI***		
UDI Primers Set A	7001 (96 test)	
UDI Primers Set B	7002 (96 test)	
UDI Primers Set C	7003 (96 test)	
UDI Primers Set D	7004 (96 test)	
UDI Primers Set 16	7005 (16 test)	

^{**}the kit is also available in its version only for research use (RUO).

^{***}UDI are always included in CEIVD kit.

