

TwinStrand Duplex Sequencing[™] Mutagenesis Panel (Human-50), v1.0

Tube Label:



Mutagenesis Panel (Human-50) v1.0

Part Number: 06-1005-XX

Pipeline ID: human-muta-v1.0

Total Panel Size: 48 kb **Number of Target Intervals:** 20

Target Type: representative genomic

regions

Genome Reference: hs38DH (Human)

This human mutagenesis panel is designed to quantify and characterize background and induced mutations in the nuclear genome of human DNA samples. Probes target 20 representative genomic regions not believed to be under significant positive or negative somatic selection. Outputted mutation data are used to estimate genome-wide per-nucleotide mutation frequency, spectrum and trinucleotide signatures for different genotoxic exposures.

Target Regions in Panel						
Chromosome	Start	End				
chr1	84,597,127	84,599,527				
chr2	40,162,767	40,165,167				
chr4	22,386,244	22,388,644				
chr6	155,239,014	155,241,414				
chr7	11,732,774	11,735,174				
chr8	51,513,056	51,515,456				
chr9	23,709,463	23,711,863				
chr10	128,969,037	128,971,437				
chr11	108,510,787	108,513,187				
chr12	114,115,043	114,117,443				
chr13	75,803,912	75,806,312				
chr14	74,661,755	74,664,155				
chr15	46,089,737	46,092,137				
chr16	51,754,103	46,092,137				
chr17	70,672,726	70,675,126				
chr18	5,749,264	5,751,664				
chr19	31,831,021	31,833,421				
chr20	24,153,684	24,156,084				
chr21	23,665,976	23,668,376				
chr22	48,262,370	48,264,770				

Coordinates in the BED format (0 start, half-open)



Panel-Specific Protocol Recommendations

Cot-1 DNA Recommended:Human: 03-2020-XXNumber of Final PCR Cycles:14 (PCR-2), 6 (PCR-3)Number of Captures Required:1Conditioning Reagent:Strongly recommended

Typical On-Target Percentage: 90%-95% (1 capture), 95%-99% (2 captures)

Panel-Specific Protocol Notes

DNA Conditioning: For mutagenesis panels, which require extreme sensitivity in the 10⁻⁷ mutations per nucleotide range, the DNA conditioning step should always be performed.

1 vs. 2 Captures: With probe panels of this size, which have been specifically designed to avoid pseudogenes and other difficult regions, a single round of hybrid capture is generally sufficient to obtain 90% or greater on-target bases. However, we recommend starting with 2 captures the first time through the protocol to generate data that can be compared against. An increase in off-target reads means that raw sequencing must be increased proportionally to generate the same amount of informative data, so a trade-off exists between time saved with only 1 capture vs. additional sequencing cost. The user should consider the pros and cons in the context of their specific situation.

Final PCR Cycles: The number of final PCR cycles listed in the table above assumes 500 ng of high quality genomic DNA input at the beginning of library preparation. However, the number of final PCR cycles necessary may vary with genomic DNA input mass, DNA quality, probe panel size and nature, number of captures, thermocycler calibration and other factors. The values in the table should be considered a starting point. If 2 rounds of hybrid capture are performed then PCR-3 is the final PCR, and with 1 round of capture PCR-2 is the final PCR.

Statistical Power: Mutagenesis experiments should typically target a specific number of Informative Duplex Bases per sample such that a sufficient number of mutagenic events are observed. In general, a minimum of 10-15 mutations are ideal for simple frequency measurements, and at least 100 mutations are needed to assess trinucleotide spectra. When calculating overall mutation frequency, unique mutation events are the numerator, and Informative Duplex Bases are the denominator. Mutation frequency in genomic DNA from control tissue samples can be as low as 1x10⁻⁷, but that must be determined empirically for each sample type. In the absence of prior data, we recommend targeting between 500 million and one billion Duplex Bases to start.

DNA Input vs. Projected Duplex Sequencing Data							
DNA Input (ng)*	Number of Clusters (Million)**	Number of Paired-End Reads (Million)**	Illumina Sequencing Cost (NovaSeq, S4, List Price)***	Mean Duplex Depth****	Max Duplex Depth****	Informative Duplex Bases (Million)****	
250	50	100	\$144	5,500x	7,000x	310	
500	100	200	\$288	11,000x	14,000x	620	
1,000	200	400	\$576	22,000x	28,000x	1,240	

Assumes high molecular weight genomic DNA (quantified pre-fragmentation). DNA requirements to achieve a given Duplex Depth when using cell-free DNA or heavily damaged DNA such as that from FFPE will be lower and higher, respectively.

TwinStrand Duplex Sequencing data yields are dependent on accurate quantification, and the quality of input DNA. We recommend Qiagen silica column-based kits or "salting out" methods for DNA extraction. Phenol-chloroform extraction can damage DNA and reduce data yields. DNA should be quantified using a double-stranded DNA-binding fluorescent dye (e.g. Qubit fluorometer, Thermo Fisher Scientific) rather than UV spectrometry which can overestimate concentration if contaminants are present, which will reduce data yields. If possible, check genomic DNA quality using a 0.5% agarose gel, or a fragment analyzer system (e.g. Agilent TapeStation system), DNA integrity number (DIN) >7 on a fragment analyzer or a single high molecular weight band on a gel generally indicates good quality DNA, although oxidation or other forms of non-fragmentation chemical damage may still be present and not appreciable.

©2020 TwinStrand Biosciences, Inc. All rights reserved. All trademarks are the property of TwinStrand Biosciences, Inc. or their respective owners. This Information Sheet may contain trademarks, service marks and trade names of other parties, which are the property of their respective owners. Our use or display of third parties' trademarks, service marks, trade names or products herein is not intended to, and does not imply a relationship with, or endorsement or sponsorship by them or us.



PDS 000003 Rev. 1.0

Approximate sequencing required to achieve a peak tag family size ~10.

S4 list price = \$7,200/lane (2.5 billion clusters/lane) as of December 2019.

^{****} Data estimates assume 150 bp paired-end reads