



Ranking mutations

Team TRS
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Sequence analysis pipeline

Variant calling files (tumor, blood normal, sibling normal) gatk **Extract samples** Filter variants to chromosome 22 **ANNOVAR** Variant annotations Exonic regions excluding synonymous

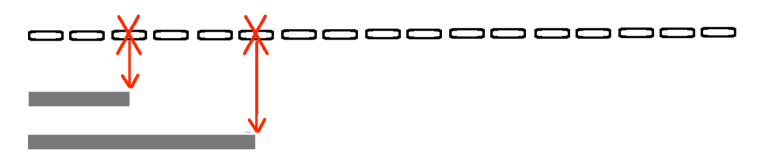
NF2 (neurofibromatosis type 2)

- There are only intronic variants (SNPs and indels) in normal and sibling normal samples
- Tumor sample:
 - Two stop stopgains

Chrom	Start	End	Ref	Alt	Function	Gene	Function	Clinvar
22	30050657	30050657	С	G	exonic	NF2	stopgain	Novel
22	30057302	30057302	С	Т	exonic	NF2	stopgain	Yes

Biological mechanism of NF2 mutations

Both alleles of NF2 were mutated



Double-knockout

Chrom	Start	End	Ref	Alt	Function	Gene	Exonic Function	Clinvar	Genotype
22	30050657	30050657	С	G	exonic	NF2	stopgain	Novel	CG
22	30057302	30057302	С	Т	exonic	NF2	stopgain	Yes	СТ

Summary of variants on chromosome 22

• Intronic, intergenic, non-coding RNA, 5'- and 3'- UTR

	Tumor	Normal blood	Sibling blood	Tumor – Normal_blood	Tumor – Sibling_blood	Normal - Sibling
Total variants	72852	71772	71385	-	-	-
All exonic variants	549	545	538	11	539	536
Non- synonymous exonic variants (genes)	289 (155)	287 (155)	267 (142)	7 (2)	93 (<mark>38</mark>)	92 (38)

7 non-synonymous exonic variants present in tumor compared to blood normal

Chr	Position	Ref	Alt	Function	Gene	Exonic Function
22	16287339	С	Т	exonic	POTEH	nonsynonymous SNV
22	20456765	G	Α	exonic	RIMBP3	nonsynonymous SNV
22	21570373	С	Т	exonic	GGT2	nonsynonymous SNV
22	29075651	G	Α	exonic	TTC28	stopgain
22	29075660	С	Т	exonic	TTC28	nonsynonymous SNV
22	30050657	С	G	exonic	NF2	stopgain
22	30057302	С	Т	exonic	NF2	stopgain

38 genes that have variants in tumor, but not in the sibling

MICAL3, MRPL40, TBX1, COMT, DGCR8, TRMT2A, ZDHHC8, TMEM191B, MED15, P2RX6, CCDC116, MAPK1, DRICH1, DERL3, SLC2A11, GSTT2; GSTT2B, GGT1, MN1, TTC28, NF2, LIF, SEC14L3, SMTN, PLA2G3, PIK3IP1, PRR14L, BPIFC, TMPRSS6, GGA1, APOBEC3B, ATF4, MKL1, EP300, FAM109B, A4GALT, CERK, SELENOO, SHANK3

COMT:

- Controls estrogen, neurotransmitter, and toxin elimination, anxiety and mood swings are prevalent.
- MAPK1: non-synonymous exonic variant
 - $T \rightarrow CSNP$
 - an essential component of the MAP kinase signal transduction pathway
 - the MAPK/ERK cascade mediates diverse biological functions such as cell growth, adhesion, survival and differentiation through the regulation of transcription, translation, cytoskeletal rearrangements.

Insertion in MN1 (Meningioma 1)

- 3-base pair insertion
- In Onno's tumor and blood samples, but not in his sibling
- MN1 Proto-Oncogene, Transcriptional Regulator

Conclusions

- NF2 was sporadic mutations occurred after embryonic development
 - Two stopgains (SNP) in NF2 genes, but no mutations in blood normal or sibling normal
- Discovered MN1 variant in tumor