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Exome Results & Raw Data Summary

Generated on: 4/26/2012

Congratulations! Your exome has been sequenced and your data is ready for you to download. We have also included this overview of your data to get you started on your exome exploration. Here are a few important points about your exome data:

- Two types of files are available for download: 1) the aligned sequencing reads in BAM format, 2) a file containing variant calls (VCF file).
- The raw data VCF file is a preliminary draft of your exome. Our ability to call variants, especially indels, is greatly improved with each additional exome added to our database. Moreover we will build upon this protocol to include additional steps such as custom treatment of the sex chromosomes. To this end we will update your VCF file at the end of the pilot. We will contact you when this data is available.

Your exome at a glance:
Your exome in numbers
Characterizing your variants
How rare are your variants?
Filtering your variants
See selected variants
Appendix

The Exome Service is a pilot project, and this report contains preliminary data only. 23andMe does not represent that all of this information is accurate. In this report we have used 1000 Genome Project data to report frequencies of variants to determine how common or rare a particular variant is. We have also only provided information about a subset of the many gene-disrupting variants present in the human genome, in a chosen set of genes. Sequencing was performed such that the total number of bases read was at least 80X the size of the exome. As described in the Exome Terms of Use, 23andMe will not be providing the reports and explanations that 23andMe typically provides to customers with respect to their genotyping results for this data. 23andMe Services are for research, informational, and educational use only. We do not provide medical advice. Please keep in mind that genetic information you share with others could be used against your interests.

Your exome in numbers

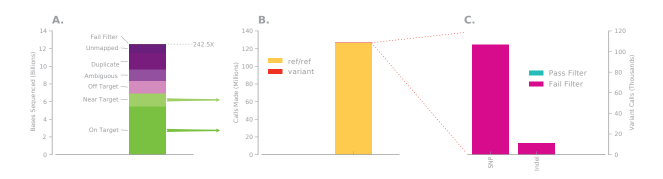
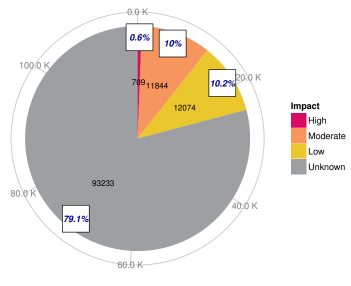


Figure 1: Getting from raw reads to called variants. A) The number of bases obtained by sequencing your exome. The top line indicates total coverage. B) Total number of called bases in your exome. The vast majority are the same as the reference genome. C) An expansion of the small sliver of variants depicted in B. These are the variants present in your VCF file.

Welcome to your exome. Your exome is the 50 million DNA bases of your genome containing the information necessary to encode all your proteins. Your exome data consists of two parts, the raw data (both aligned and unaligned Illumina reads, fig1A) and a draft of the variants present in your exome (fig1C). While this draft is provisional and we will be improving upon it, we wanted to allow you to dig in to your exome as soon as possible so you can tell us what you think is important and should be included.

To create the first draft of your exome we implemented the Broad Institute's "Best Practice" protocol for exome sequencing analysis. You can read a detailed description of it here (for brief summary see Appendix).

Characterizing your variants



Number of variants

Figure 2: Predicting impact of variants on gene function. An overview of your variants and their predicted impact on gene function.

The variants in your VCF file are the positions in your genome that differ from the reference genome. Most of these variants are likely to be functionally neutral and unlikely to cause any severe disorders. Pinpointing genuine disease mutations is still challenging and we used a number of software tools to identify those that may be functionally important. We estimated the impact a variant has on gene function based on the severity of its effect on the gene product:

High impact:

Frame shift Insertion or deletion of bases, not multiple of 3.

Splice site Variant at the 'splicing site' may disrupt the consensus splicing site sequence.

Stop gain Premature termination of peptides, which would disable protein function.

Start loss Loss of the start codon.

Stop loss Loss of the stop codon.

Moderate impact:

Nonsynonymous substitution Non-conservative change altering an amino acid in a protein.

Codon insertion or deletion Insertion or deletion of bases, multiple of 3.

Low impact:

Synonymous substitution Variant that does not alter the amino acid sequence due to codon degeneracy.

Start gain Variant resulting in the gain of a start codon.

Synonymous stop Variant changing one stop codon into another.

Unknown impact: Variants unlikely to affect gene products.

How rare are your variants?

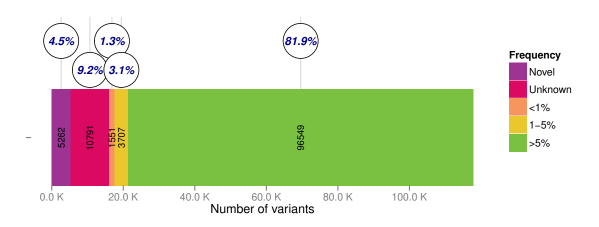


Figure 3: Variant frequencies. The allele frequencies of the variants in your exome. Unknown: allele is present in a public database but no frequency data was available.

One of the advantages of exome sequencing is that we can detect sequence variants that are unique to you! By comparing your variants to all those that have been discovered so far, we can divide your variants into the following categories:

- novel variant hasn't been observed in current public sequence databases
- **unknown** variant has been observed in public databases but allelic frequency has not been calculated and therefore is not available
- rare variant with allelic frequency <1%
- somewhat rare variant with frequency 1-5%
- common frequency of the variant is greater than 5%

One of the most comprehensive human variation public datasets is maintained by the 1000 Genomes Project. We use 1000 Genomes Project data (project release: 08-26-2011) to report frequencies of alleles found in your exome, including reporting if it is absent from the public database (*i.e.* a novel variant).

Filtering your variants

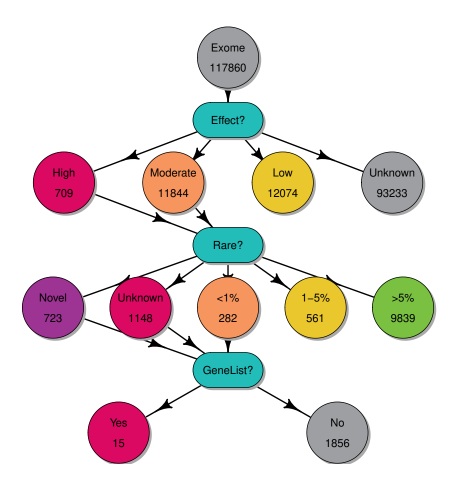


Figure 4: Variant filtering decision tree. A graphical representation of the filtering process that was used to generate your short list of variants of interest.

Most sequence variants in your exome are likely to be neutral and do not cause any severe disorders. A filtering process is often undertaken to prioritize variants discovered through sequencing. To identify potentially interesting and relevant variants with potential functional effects (contributing to disease and other phenotypes of interest) we used three consecutive filters, depicted in the figure above: (1) effect of the variant on the gene product; (2) allele frequency of the variant; (3) location of the variant in one of 592 genes involved in Mendelian disorders (at this point we also exclude indels and variants on the sex chromosomes).

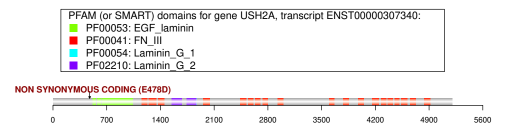
We hope you find this initial list of variants interesting and that it will help you in your journey through your exome. This short list of variants only scratches the surface of what your genome contains and is just the beginning of where your data can take you. Have fun!

List of selected variants

Variant 1:	Gene: CLN5 Your genotype: C/A Location: chr13:77574606					
	Impact: NON SYNONYMOUS CODING	Type: MODERATE				
	1KGenomes: 0.00270	dbSNP: rs138611001				
	Genotype quality: 99	Coverage depth: 63				
	Gene description: ceroid-lipofuscinosis, neuronal 5					
	Transcript: ENST00000535238	AA change: N108K				
	Entrezld: 1203	Ensemblid: ENSG00000102805				
	UniProt: O75503	OMIM: 608102				

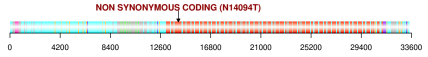


Variant 2:	Gene: USH2A Your genotype: C/G Location: chr1:216496932					
	Impact: NON SYNONYMOUS CODING	Type: MODERATE				
	1KGenomes: 0.00730	dbSNP: rs35730265				
	Genotype quality: 99	Coverage depth: 122				
	Gene description: Usher syndrome 2A (autosomal recessive, mild)					
	Transcript: ENST00000307340	AA change: E478D				
	Entrezld: 7399	Ensemblid: ENSG00000042781				
	UniProt: 075445	OMIM: 608400				

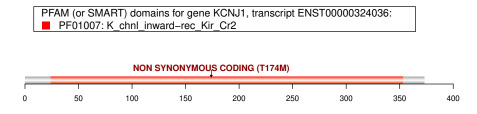


Variant 3:	Gene: TTN Your genotype: T/G Location: chr2:179477267					
	Impact: NON SYNONYMOUS CODING	Type: MODERATE				
	1KGenomes: 0.00830	dbSNP: rs36043230				
	Genotype quality: 99	Coverage depth: 180				
	Gene description: titin Transcript: ENST00000342992 EntrezId: 7273 UniProt: Q8WZ42	AA change: N14094T Ensemblid: ENSG00000155657 OMIM: 188840				

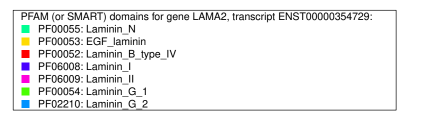
PFAM (or SMART) domains for gene TTN, transcript ENST00000342992:
PF07679: lg_l-set
PF09042: Titin_Z
PF07686: lg_V-set
PF00047: Immunoglobulin
PF02818: PPAK_motif
PF00041: FN_III
PF00069: Se/Thr_kinase-like_dom
PF07714: Ser-Thr/Tyr_kinase

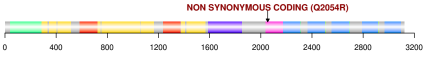


Variant 4:	Gene: KCNJ1 Your genotype: G/A Location: chr11:128709618					
	Impact: NON SYNONYMOUS CODING	Type: MODERATE				
	1KGenomes: 0.00270	dbSNP: rs117535913				
	Genotype quality: 99	Coverage depth: 123				
	Gene description: potassium inwardly-recti Transcript: ENST00000324036 Entrezld: 3758 UniProt: P48048	fying channel, subfamily J, member 1 AA change: T174M Ensemblid: ENSG00000151704 OMIM: 600359				

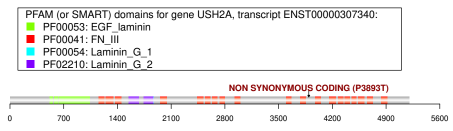


Variant 5:	Gene: LAMA2 Your genotype: A/G Location: chr6:129762036					
	Impact: NON SYNONYMOUS CODING	Type: MODERATE				
	1KGenomes: 0.00230	dbSNP: rs56035053				
	Genotype quality: 99	Coverage depth: 88				
	Gene description: laminin, alpha 2 Transcript: ENST00000354729 EntrezId: 3908 UniProt: P24043	AA change: Q2054R Ensemblid: ENSG00000196569 OMIM: 156225				

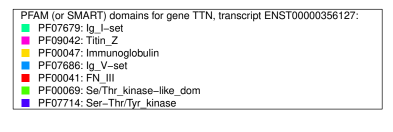


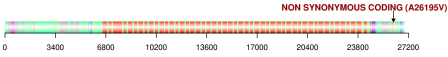


Variant 6:	Gene: USH2A Your genotype: G/T Location: chr1:215914751					
	Impact: NON SYNONYMOUS CODING	Type: MODERATE				
	1KGenomes: 0.00780	dbSNP: rs41303285				
	Genotype quality: 99	Coverage depth: 181				
	Gene description: Usher syndrome 2A (aut Transcript: ENST00000307340 EntrezId: 7399 UniProt: O75445	osomal recessive, mild) AA change: P3893T Ensemblid: ENSG00000042781 OMIM: 608400				

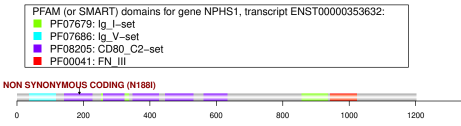


Variant 7:	Gene: TTN Your genotype: G/A Location: chr2:179395554					
	Impact: NON SYNONYMOUS CODING	Type: MODERATE				
	1KGenomes: 0.00530	dbSNP: rs66961115				
	Genotype quality: 99	Coverage depth: 201				
	Gene description: titin Transcript: ENST00000356127 Entrezld: 7273 UniProt: Q8WZ42	AA change: A26195V Ensemblid: ENSG00000155657 OMIM: 188840				





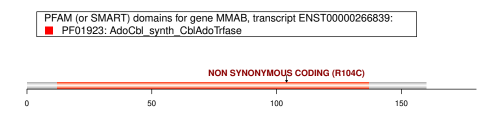
Variant 8:	Gene: NPHS1 Your genotype: T/A Location: chr19:36341311					
	Impact: NON SYNONYMOUS CODING	Type: MODERATE				
	1KGenomes: 0.00290	dbSNP: rs145125791				
	Genotype quality: 99	Coverage depth: 68				
	Gene description: nephrosis 1, congenital, Transcript: ENST00000353632 EntrezId: 4868 UniProt: O60500	Finnish type (nephrin) AA change: N188I EnsemblId: ENSG00000161270 OMIM: 602716				



Variant 9:	Gene: CEP290 Your genotype: C/T Loc	cation: chr12:88472996		
	Impact: NON SYNONYMOUS CODING	Type: MODERATE		
	1KGenomes: 0.00330	dbSNP: rs61941020		
	Genotype quality: 99	Coverage depth: 126		
	Gene description: centrosomal protein 290 Transcript: ENST00000552810 Entrezld: 80184 UniProt: O15078	DkDa AA change: R1746Q Ensemblid: ENSG00000198707 OMIM: 610142		

NON SYNONYMOUŞ CODING (R1746Q)								
	1	1	1	1	1	1	Г	
0	350	700	1050	1400	1750	2100	2450	28

Variant 10:	Gene: MMAB Your genotype: G/A Location: chr12:109998846			
	Impact: NON SYNONYMOUS CODING	Type: MODERATE		
	1KGenomes: 9e-04	dbSNP: NA		
	Genotype quality: 99	Coverage depth: 52		
	Gene description: methylmalonic aciduria Transcript: ENST00000266839 Entrezld: 326625 UniProt: Q96EY8	(cobalamin deficiency) cblB type AA change: R104C EnsemblId: ENSG00000139428 OMIM: 607568		



Variant 11:	Gene: SLC34A2 Your genotype: G/C Lo		
	Impact: NON SYNONYMOUS CODING	Type: MODERATE	
	1KGenomes: 0.00320	dbSNP: rs76404281	
	Genotype quality: 99	Coverage depth: 85	
	Gene description: solute carrier family 34 Transcript: ENST00000503434 EntrezId: 10568 UniProt: O95436	(sodium phosphate), member 2 AA change: L413F Ensemblid: ENSG00000157765 OMIM: 604217	

PFAM (or SMART) domains for gene SLC34A2, transcript ENST00000503434: PF02690: Na/Pi_cotranspt

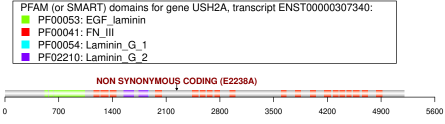


Variant 12:	Gene: KCNE1 Your genotype: C/T Location: chr21:35821680			
	Impact: NON SYNONYMOUS CODING	Type: MODERATE		
	1KGenomes: 0.00490	dbSNP: rs1805128		
	Genotype quality: 99	Coverage depth: 90		
	Gene description: potassium voltage-gated Transcript: ENST00000337385 Entrezld: 3753 UniProt: P15382	channel, lsk-related family, member 1 AA change: D85N Ensemblid: ENSG00000180509 OMIM: 176261		

PFAM (or SMART) domains for gene KCNE1, transcript ENST00000337385: ■ PF02060: K_chnl_volt-dep_bsu_KCNE



Variant 13:	Gene: USH2A Your genotype: T/G Location: chr1:216166454				
	Impact: NON SYNONYMOUS CODING	Type: MODERATE			
	1KGenomes: 0.00970	dbSNP: rs41277212			
	Genotype quality: 99	Coverage depth: 198			
	Gene description: Usher syndrome 2A (au Transcript: ENST00000307340 Entrezld: 7399 UniProt: O75445	utosomal recessive, mild) AA change: E2238A Ensemblid: ENSG00000042781 OMIM: 608400			
PEAM (or SMART) domains for gene USH2A, transcript ENST00000307340;					



Variant 14:	Gene: TTN Your genotype: C/A Location: chr2:179395555			
	Impact: NON SYNONYMOUS CODING	Type: MODERATE		
	1KGenomes: 0.00530	dbSNP: rs67254537		
	Genotype quality: 99	Coverage depth: 201		
	Gene description: titin Transcript: ENST00000356127 EntrezId: 7273 UniProt: Q8WZ42	AA change: A26195S Ensemblid: ENSG00000155657 OMIM: 188840		

M (or SMART) domains for gene TTN, transcript ENST00000356127: PF07679: lg_l-set
PF09042: Titin_Z
PF00047: Immunoglobulin
PF07686: Ig_V-set
PF00041: FN_III
PF00069: Se/Thr_kinase-like_dom
PF07714: Ser-Thr/Tyr_kinase

						NON SYNON	IYMOUS CO	DING (A26195S)
0	3400	6800	10200	13600	17000	20400	23800	27200

Variant 15:	Gene: SMPD1 Your genotype: G/A Location: chr11:6413167			
	Impact: NON SYNONYMOUS CODING	Type: MODERATE		
	1KGenomes: 5e-04	dbSNP: rs1803161		
	Genotype quality: 99	Coverage depth: 138		
	Gene description: sphingomyelin phosphod Transcript: ENST00000530395 Entrezld: 6609 UniProt: P17405	liesterase 1, acid lysosomal AA change: R18H EnsemblId: ENSG00000166311 OMIM: 607608		

PFAM (or SMART) domains for gene SMPD1, transcript ENST00000530395: PF00149: Metallo_PEstase_dom

NON SYNONYMOUS CODING (R18H)

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Appendix

To create the first draft of your exome we implemented the Broad Institute's "Best Practice" protocol for exome sequencing analysis. You can read a detailed description of it here, however a brief summary of it follows:

- 1. We took your raw reads and aligned them against the reference genome (these are the alignments available in the BAM file of the encrypted download).
- 2. We used these alignments to identify probable contamination (unaligned reads) and artifacts of sample preparation (PCR duplicates) which are then removed from subsequent steps.
- 3. From this point on we focus on the reads that align either to one of the exons or within the regions 250 bases up and downstream of it.
- 4. To improve the quality of the alignments we carry out a more accurate alignment of the reads that overlap known indels or are likely to contain indels themselves.
- 5. We also recalibrate the base quality scores of the reads to bring them in line with the empiricallydetermined values.
- 6. Using these realigned+recalibrated reads we generate allele calls at every position with enough high-quality data and filter out those that are homozygous for the allele present in the reference genome (the vast majority of these are at such a high frequency in the population they're unlikely to be interesting). The remaining SNP and indel calls (variants) are the ones available in the VCF file that you downloaded.
- 7. As yet no sequencing technology is 100% accurate and the highly duplicated nature of the human genome makes variant calling a challenging task. Consequently, a small proportion of the variant calls in your VCF are likely to be incorrect. To reduce this proportion we applied the filters recommended by the Broad Institute to remove technical artifacts. Variants that pass all filters are marked in your VCF file with a PASS. As the exome pilot progresses and we gather more data we will be able to use more advanced techniques identify potential errors and improve the quality of your exome.