PGP-UK Genomics Report for uk6C54D7

1 Summary

This is the genome report was produced using collaborative research tools, including SNPedia and GetEvidence. This section shows an overview of all the small variants which were found in the genome for this individual, when compared with a reference genome. These variants are summarised in Table 1 and the pie-charts in Figures 2, 3 and 4.

This report was generated automatically and is not clinically approved. It is provided for <u>personal and research purposes</u> only.

This document contains hyperlinks, shown in grey, that will take you to external websites where you can find more detailed explanations. Some of the technical terms are also explained in more detail in the Ensembl Glossary. We would welcome your feedback about this report, for example, if you would like more information about anything or if any of the links have become inactive. You can contact us on: pgp-uk@ucl.ac.uk.

This summary shows an overview of all the variants which were found in the genome for this individual. The "variants remaining after filtering" refers to any differences in the DNA identified when compared to the reference genome. Of these, the majority will have already been found in some other sequenced individual and put on a database (existing variants) while others have not yet been annotated (novel variants).

"Overlapped genes" refers to the number of times where a variant was found in a region of the genome containing a gene. The diagram in Figure 1 is a simplification of the usual gene structure. "Exon" refers to the part of the gene which goes on to form a protein, and variants in this part of the gene are more likely to cause changes in the shape of the protein. Upstream, downstream, intronic and intergenic variants are more likely to alter the regulation of that gene but will not change the protein itself.

A transcript for a protein-coding gene can include the exons, introns and other gene features that are transcribed and important for gene function but might not be translated into the final protein. Not all transcripts are for protein-coding genes, with many containing non-coding RNAs that can be overlapping other genes, in introns or in intergenic regions.

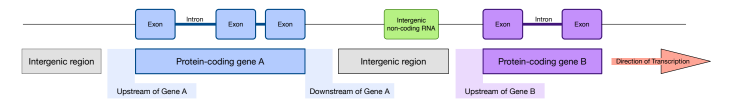


Figure 1: Diagram of gene structure indicating locations of potential variants

Feature	Count
Lines of input read	4898683
Variants filtered out	0
Novel / existing variants	474278 (9.7) / 4413605 (90.3)
Overlapped genes	56781
Overlapped transcripts	67547
Overlapped regulatory features	165499

Table 1: Variant calling summary

There are several different types of genomic variants. The most common change is when one single building block of the DNA (called a nucleotide) is changed, called a single nucleotide variants (SNV). Other variant types include insertions, where the DNA in the individual is longer than the reference sequence due to the insertion of one or more nucleotides; and deletions, where a few nucleotides are missing compared to the reference sequence.

Some of these changes will have no effect on the protein, while some changes may alter the protein function to varying degrees. The PolyPhen analysis software attempts to quantify the effect each mutation will have on the protein function. This ranges from "benign" where no change to the protein function is expected, to "probably damaging" where it is predicted that the mutation will affect protein function. It is nevertheless important to note that what is "damaging" for the protein is not necessarily damaging for the individual.

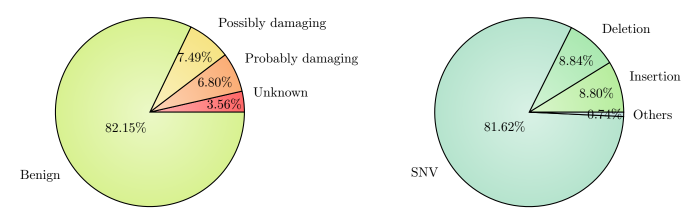


Figure 2: PolyPhen Summary

Figure 3: Variant Class

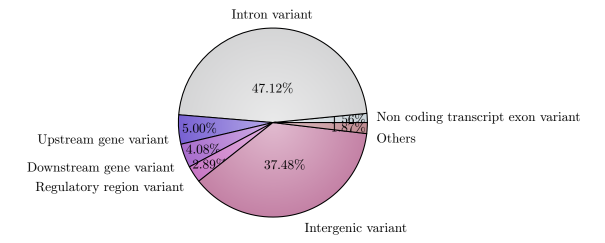


Figure 4: Consequence type

2 Ancestry

This plot shows the distribution of the genomes of different populations. Data from several studies which used whole genome sequencing was used to see the relationships between the genomes of the populations. It shows how closely related certain populations are genetically: Groups which cluster closely are more genetically similar than groups which are further apart. The black star symbol shows where this PGP-UK participant sits in relation to other populations, indicating their ancestry and their most closely related populations according to genetic sequence.

Based on the populations defined in the 1000 genomes project (1kGP), the ancestry composition for this individual is inferred to be 100.0 percent European [British in England and Scotland].

Please note that this analysis is limited by the populations available in the 1kGP data. If there are European subpopulations reported, and the ancestry of the participant does not correspond to any of the 1kGP populations, the closest 1kGP sampled subpopulation will be shown (even though it might be different from the participant's actual ancestry).

Ancestry uk6C54D7

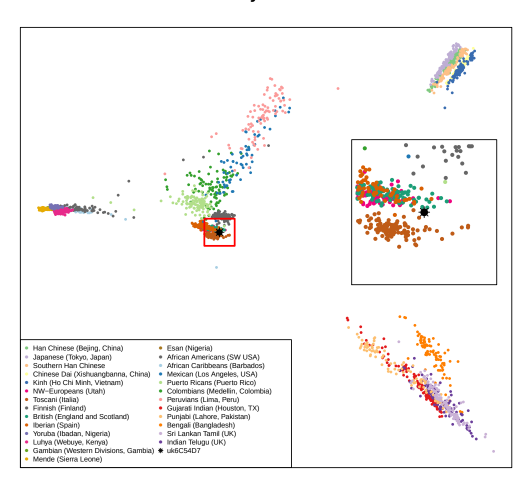


Figure 5: Ancestry Principal Component Analysis

3 Traits (based on SNPedia information)

Existing research has associated many variants with phenotypic traits, some of which can be perceived as beneficial while others appear to have a harmful effect. Some traits are complex and can be affected by several variants. It is likely that some of these would confer a higher risk while others a lower risk of trait manifestation. These can not be combined linearly to produce an actual risk of disease.

It is important to note that in most cases genomic data is probabilistic, not deterministic- i.e. having a genetic predisposition for a disease is not a diagnosis; rather, it shows an increased likelihood of developing that disease. Also, one person can have both potentially beneficial and harmful variants in the same gene, or associated with the same disease.

Some variants can also affect certain populations more, or will only affect a particular gender. For example, a variant for higher risk of endometriosis in the sequence of a male will not directly affect that person, but can be passed on to descendants.

While many traits are the result of a unique variant, many are the combination of several variants throughout the genome. In SNPedia, these are called genosets. These can integrate some of the information already present in the single variant tables, or be the combination of variants that have no phenotypic effect on their own, but contribute to a trait when together.

The variants in the following tables are sorted by magnitude. This is an subjective measure defined in SNPedia to highlight the perceived importance of the genotype described. At the moment this scale goes from 0 to 10. You can read more about it by visiting their explanatory webpage.

As our knowledge grows, the interpretation of the effect of certain variants might change. Clicking on the links in the genome report tables will take you to websites containing more information about each variant.

3.1 Possibly Beneficial Traits

Mag.	Identifier	Genotype	Summary	ExAC	GetEvidence	ClinVar
3	rs7294919	(C;T)	Moderately enhanced hippocampal volume			
2.4	rs9272346	(G;G)	0.08x risk type-1 diabetes		Link	
2.1	rs2511989	(A;G)	0.63x decreased age-related macular degeneratio		Link	
2	rs1012053	(A;C)	0.625x reduced risk of Bipolar Disorder.		Link	
2	rs10468017	(C;T)	Associated with higher HDL cholesterol		Link	
2	rs11045585	(A;A)	24% chance (lower than average) of docetaxel-in		Link	
2	rs12979860	(C;C)	~80% of such hepatitis C patients respond to tr		Link	Link
2	rs1800972	(G;G)	Reduced risk for Crohn's disease; reduced risk	Link		
2	rs1864163	(G;G)	Associated with higher HDL cholesterol		Link	
2	rs2056202	(T;T)	Rare decreased risk of autism	Link		
2	rs2243250	(C;T)	0.6x decreased risk for myocardial infarction i			
2	rs2292813	(T;T)	Decreased risk of autism			
2	rs2542052	(C;C)	Better odds of living to 100			
2	rs3738579	(C;T)	0.5x decreased risk for cervical cancer: HNSCC:			
2	rs3750817	(C;T)	0.78x reduced risk for breast cancer			
2	rs3782179	(C;T)	3x lower odds of testicular cancer risk for men			
2	rs4149268	(A;G)	Associated with higher HDL cholesterol		Link	
2	rs6495446	(T;T)	0.64x reduced risk for chronic kidney disease			
2	rs6807362	(G;G)	Decreased autism risk	Link	Link	
2	rs6855911	(A;G)	0.62x decreased risk for gout		Link	
1.8	rs1128535	(A;G)	0.77x risk for Crohn's disease			
1.8	rs266729	(C;G)	0.73x decreased risk for colorectal cancer		Link	
1.8	rs3814113	(C;T)	0.8x decreased risk for ovarian cancer		Link	
1.8	rs4714156	(C;C)	<0.61x risk for restless legs			
1.6	rs1061170	(T;T)	Lower risk for AMD: generally longer live than	Link	Link	Link
1.5	rs11212617	(A;C)	Somewhat increased likelihood of treatment succ			Link
1.5	rs4149274	(C;T)	Associated with higher HDL (good) cholesterol			

Mag.	Identifier	Genotype	Summary	ExAC	GetEvidence	ClinVar
1.5	rs4939883	(C;T)	Associated with higher HDL cholesterol		Link	
1.4	rs1165205	(A;T)	0.85x decreased gout risk		Link	
1.4	rs2294008	(C;C)	Lower risk of gastric and bladder cancer	Link	Link	
1.4	rs6700125	(C;C)	0.7x decreased risk for ALS			
1.2	rs11246226	(A;C)	Decreased risk of schizophrenia in limited stud		Link	
1.2	rs4320932	(A;G)	0.87x decreased risk for ovarian cancer			
1.2	rs6048	(G;G)	Slightly lower risk (10-20%) of deep vein throm	Link	Link	Link
1.2	rs9306160	(C;T)	0.75x (reduced) risk for metastasis in LN-/ER+	Link	Link	
1.1	rs2293347	(G;G)	Among NSCLC patients: better Gefitinib response	Link		Link
1.1	rs4988235	(T;T)	Can digest milk			Link
1.1	rs7568369	(G;T)	0.90x reduced risk of obesity			
1	rs182549	(T;T)	Can digest milk.			Link
1	rs2351299	(G;T)	Possible reduced risk of Autism			
1	rs2546890	(G;G)	Lower risk of multiple sclerosis			
1	rs33927012	(C;T)	Currently evaluated as benign in ClinVar	Link	Link	Link
1	rs7850258	(A;G)	Typical odds of developing primary hypothyroidi			
1	rs800292	(C;T)	1% decreased risk of macular degeneration	Link	Link	Link
0.1	rs891512	(G;G)	Lower blood pressure than those with an A allel	Link		
0	rs1047781	(A;A)	ABH blood group "Secretor" status if Japanese	Link	Link	Link
0	rs12252	(T;T)	More resistant to influenza	Link		Link
0	rs16990018	(A;A)	PrP Codon 171 Asn - Non-pathogenic variant	Link		Link
0	rs17244841	(A;A)	More responsive to statin treatment		Link	Link
0	rs1799782	(C;C)	Lower risk for skin cancer	Link	Link	
0	rs1800562	(G;G)	Not a C282Y hemochromatosis carrier.	Link	Link	Link
0	rs28933385	(G;G)	Prion protein Codon 200 (E) - Non pathogenic va			Link
0	rs312481	(C;C)	Better response to certain calcium channel bloc			
0	rs5065	(A;A)	1.12x risk on diuretic; if hypertensive: better	Link	Link	Link
0	rs6259	(G;G)	Best inverse correlation between tea-drinking:	Link	Link	
0	rs7305115	(A;A)	Individuals showed a significantly lower risk o	Link	Link	
0	rs74315403	(G;G)	PrP codon 178 (D) - non pathogenic variant			Link
0	rs7495174	(A;A)	Blue/gray eyes more likely		Link	

3.2 Possibly Harmful Traits

Mag.	Identifier	Genotype	Summary	ExAC	GetEvidence	ClinVar
3	rs10897346	(C;C)	If depressed: 2.6x more likely to not respond t			
3	rs2237717	(T;T)	Reduced abilities related to neurocognition and			
3	rs2306402	(C;C)	1.18x increased risk for late-onset Alzheimer's			
3	rs3892097	(A;A)	CYP2D6 poor metabolizer; many associations rela	Link	Link	Link
3	rs6920220	(A;G)	1.2x risk Rheumatoid Arthritis		Link	
3	rs7754840	(C;C)	1.3x increased risk for type-2 diabetes		Link	
2.5	rs1121980	(C;T)	1.67x risk for obesity		Link	
2.5	rs13266634	(C;T)	Increased risk for type-2 diabetes	Link	Link	Link
2.5	rs1421085	(C;T)	~1.3x increased obesity risk		Link	Link
2.5	rs16969968	(A;G)	Slightly higher risk for nicotine dependence: l	Link	Link	Link
2.5	rs17595731	(C;G)	~5 fold higher risk for Fuchs' dystrophy: a cor			
2.5	rs187238	(G;G)	Hypertension increases risk 3.75x for sudden ca			
2.5	rs2004640	(T;T)	1.4x increased risk for SLE		Link	Link
2.5	rs2241880	(C;C)	2x-3x increased risk for Crohn's disease in Cau	Link	Link	Link
2.5	rs339331	(T;T)	Prostate cancer risk			
2.5	rs3738919	(C;C)	1.94x risk of developing rheumatoid arthritis			
2.5	rs5888	(C;T)	3x higher risk for age-related macular degenera	Link		
2.5	rs613872	(G;T)	~5 fold higher risk for Fuchs' dystrophy: a cor			
2.5	rs664143	(C;T)	Higher risk for number of cancers			
2.5	rs8034191	(C;T)	1.27x lung cancer risk		Link	
2.3	rs7966230	(C;G)	Slightly lower levels of plasma VWF			
2.2	rs2231137	(G;G)	~1.5-3x increased risk for ischemic stroke	Link	Link	Link
2.2	rs2305089	(T;T)	Higher risk for chordoma	Link	Link	
2.1	rs10427255	(C;C)	Highest odds of photic sneeze reflex			
2.1	rs10811661	(T;T)	1.2x increased risk for type-2 diabetes		Link	
2.1	rs17070145	(C;C)	Reduced memory abilities			Link
2.1	rs2231142	(A;C)	1.74x increased gout risk; gefinitib takers 4x	Link	Link	Link
2.1	rs2254958	(C;T)	1.24x increased risk for Alzheimer's			
2.1	rs4149056	(C;T)	Reduced breakdown of some drugs; 5x increased m	Link	Link	Link
2.1	rs4363657	(C;T)	4.5x increased myopathy risk for statin users		Link	
2.1	rs4693596	(C;C)	2x odds of myopathy if taking statins			
2.1	rs646776	(A;A)	1.2x risk of coronary artery disease		Link	
2.1	rs7837688	(G;T)	1.7x increased risk for prostate cancer			
2.1	rs944289	(C;T)	1.3x increased thyroid cancer risk		Link	
2	rs10086908	(C;T)	1.7x increased risk for prostate cancer			
2	rs10090154	(C;T)	1.4x increased risk for prostate cancer			
2	rs1024611	(C;T)	Increased risk of exercise induced ischemia			Link
2	rs10248420	(A;A)	7x less likely to respond to certain antidepres		Link	
2	rs1045642	(C;T)	Slower metaboliser for some drugs	Link	Link	Link
2	rs1051730	(C;T)	1.3x increased risk of lung cancer	Link	Link	Link
2	rs10984447	(A;G)	1.17x increased risk for multiple sclerosis		Link	
2	rs1160312	(A;G)	1.6x increased risk of Male Pattern Baldness.		Link	
2	rs11983225	(T;T)	7x less likely to respond to certain antidepres		Link	
2	rs12037606	(A;A)	1.52x risk of developing Crohn's disease			
2	rs1333048	(A;C)	1.3x increased coronary artery disease risk			
2	rs1360780	(C;T)	1.3x increased risk for depression		Link	
2	rs137853242	(A;G)	Associated with MODY3; maturity onset of diabet	Link		Link
2	rs1585215	(A;G)	2x increased risk for Hodgkin lymphoma			
2	rs1691053	(A;G)	Increased risk of developing prostate cancer			
2	rs16944	(G;G)	Increased risk of mental disorders		Link	
2	rs1734791	(A;A)	1.4x increased risk for lupus			
2	rs17576	(A;G)	Higher risk for MI and lung cancer: and COPD in	Link	Link	
2	rs1800896	(A;A)	1.8x increased prostate cancer risk			

Mag.	Identifier	Genotype	Summary	ExAC	GetEvidence	ClinVar
2	rs1867277	(A;A)	2x increased risk for thyroid cancer			
2	rs2066843	(T;T)	4.09x higher risk for Crohn's disease	Link	Link	
2	rs2073963	(G;T)	Increased risk of baldness			
2	rs2143340	(C;T)	Increased risk of dyslexia and poor reading per			
2	rs2156921	(A;G)	1.29x increased risk for depression			
2	rs2201841	(T;T)	2.4x increased risk for Graves' disease		Link	
2	rs2230201	(G;G)	>1.4x risk of lupus	Link		
2	rs2235015	(G;G)	Somewhat less likely to respond to certain anti	Link	Link	
2	rs2235040	(G;G)	7x less likely to respond to certain antidepres	Link	Link	
2	rs2235067	(G;G)	7x less likely to respond to certain antidepres			
2	rs2305480	(C;T)	3.5x increase in risk of asthma for Han Chinese	Link	Link	
2	rs2305795	(A;G)	1.28x higher risk of narcolepsy compared to (G;			Link
2	rs2352028	(T;T)	Increased risk of lung cancer in non-smokers an		Link	
2	rs2383206	(A;G)	1.4x increased risk for heart disease			
2	rs2383207	(A;G)	Increased risk for heart disease			
2	rs241448	(C;T)	1.51x increased risk for Alzheimer's	Link		Link
2	rs25487	(A;G)	2x higher risk for skin cancer; possibly other	Link	Link	Link
2	rs3129934	(C;T)	Increased risk of Multiple Sclerosis.		Link	
2	rs3212227	(C;C)	Significantly increased risk of developing cerv			
2	rs326	(A;A)	Lower HDL cholesterol		Link	Link
2	rs351855	(C;T)	1.2x increased risk for prostate cancer	Link	Link	Link
2	rs358806	(C;C)	1.78x increased risk of developing Type-2 diabe		Link	
2	rs3745516	(A;A)	Increased risk of developing primary biliary ci			
2	rs4148739	(A;A)	7x less likely to respond to certain antidepres		Link	
2	rs4242382	(A;G)	1.7x increased risk for prostate cancer		Link	
2	rs4444903	(A;G)	3.5x risk of hep-cancer in cirrhosis patients;			Link
2	rs5174	(A;G)	1.3x increased risk for heart disease	Link	Link	Link
2	rs520354	(A;G)	Increased risk in men for biliary conditions		T. 1	
2	rs5759167	(T;T)	Higher prostate cancer risk		Link	
2	rs6498169	(A;A)	>1.14x risk of multiple sclerosis	T . 1	Link	T . 1
2	rs669	(G;G)	3.8x or higher increased risk for Alzheimers	Link	Link	Link
2	rs6896702	(T;T)	Increased risk of developing Parkinson's Diseas	T · 1	T · 1	T · 1
2	rs6897932	(C;C)	1.08x increased risk for multiple sclerosis	Link	Link	Link
2	rs6908425	(C;C)	1.95x increased risk of developing Crohn's dise		Link	
2	rs6997709	(G;G)	1.5x higher risk for hypertension	T :1-	T :1.	
2 2	rs7530511 rs7536563	(T;T) (A;A)	9x increased risk for Graves disease >1.12x risk of multiple sclerosis	Link	Link Link	
$\frac{2}{2}$	rs7794745	(A;A) (A;T)	Slightly increased risk for autism		Link	Link
$\frac{2}{2}$	rs7923837	(G;G)	3.2x risk for T2D		TILLK	LIIIK
$\frac{2}{2}$	rs828907	(G;G) (G;T)	Slightly increased risk of bladder cancer and 2			
$\frac{2}{2}$	rs854560	(A;T)	Higher risk for heart disease: diabetic retinop	Link	Link	Link
2	rs9525638	(T;T)	Weaker bones	TITIK	THIK	THIK
$\frac{2}{2}$	rs965513	(A;G)	1.77x increased thyroid cancer risk		Link	
$\frac{2}{2}$	rs9954153	(G;T)	~2.5x higher risk for Fuchs' dystrophy: a corne		221111	
$\frac{2}{2.0}$	rs17221417	(G;G)	1.9x higher risk for Crohn's disease		Link	
2.0	rs4911414	(G;T)	2-4x higher risk of sun sensitivity if part of		Link	
2.0	rs9642880	(T;T)	1.5x increased bladder cancer risk		Link	
1.9	rs721048	(A;A)	Slightly increased prostate cancer risk		Link	Link
1.8	rs37973	(A;G)	Among asthmatics: 1.5x more likely to show less			Link
1.8	rs4474514	(A;G)	3x increased testicular cancer risk for men		Link	
1.6	rs11523871	(C;C)	>1.6x increased breast cancer risk for women ov	Link	Link	
1.6	rs1260326	(T;T)	Slightly higher risk for gout	Link	Link	Link
1.6	rs2981745	(T;T)	>1.6x increased risk for breast cancer in femal			
1.6	rs3764880	(A;A)	1.2 - 1.8x increased tuberculosis risk	Link	Link	
1.6	rs3775948	(C;G)	Slightly higher risk for gout			

Mag.	Identifier	Genotype	Summary	ExAC	GetEvidence	ClinVar
1.5	rs10757272	(C;T)	1.30x increased risk for Coronary artery diseas			
1.5	rs10980705	(C;T)	2.3x increased risk for knee osteoarthritis			
1.5	rs1169300	(A;G)	~1.5x increased lung cancer risk			
1.5	rs12431733	(C;T)	Slightly increased risk of developing Parkinson		Link	
1.5	rs12469063	(A;G)	Slightly increased risk of developing restless			
1.5	rs13149290	(C;C)	Slightly increased risk of developing prostate			
1.5	rs13181	(G;T)	1.12x increased risk for cutaneous melanoma	Link	Link	Link
1.5	rs13376333	(C;T)	1.5x higher risk of atrial fibrillation		Link	
1.5	rs140701	(A;G)	Increased risk for anxiety disorders			
1.5	rs144848	(G;T)	Very slightly increased breast cancer risk	Link	Link	Link
1.5	rs165599	(G;G)	May indicate increased susceptibility to schizo		Link	
1.5	rs1801274	(C;T)	Complex; generally greater risk for cancer prog	Link	Link	Link
1.5	rs2007153	(G;G)	Increased risk of schizophrenia in limited stud			
1.5	rs2240340	(A;G)	Slightly increased (1.5x) risk for RA	Link		
1.5	rs2272127	(C;C)	Associated with herpes and schizophrenia			
1.5	rs2280714	(A;A)	1.4x increased risk of SLE			
1.5	rs2464196	(C;T)	~1.5x increased lung cancer risk	Link	Link	Link
1.5	rs27388	(A;G)	Slightly increased risk of developing schizophr			
1.5	rs2881766	(T;T)	Slightly increased risk for pregnancy-induced h			
1.5	rs3087243	(A;G)	Increased risk for auto-immune diseases		Link	
1.5	rs309375	(T;T)	Larger mosquito bites			
1.5	rs3814570	(T;T)	1.3x increased risk for Crohn's disease with il			
1.5	rs3825776	(A;G)	1.3x increased risk for ALS		Link	
1.5	rs393152	(A;A)	Increased risk of both PD and AD	Link	Link	
1.5	rs401681	(C;C)	~1.2x increased risk for several types of cance		Link	
1.5	rs4027132	(A;G)	1.39x increased risk of developing bipolar diso			
1.5	rs464049	(C;T)	Increased risk of schizophrenia in limited stud			
1.5	rs4845618	(G;T)	1.7x increased melanoma risk			
1.5	rs486907	(A;G)	1.5x increased prostate cancer risk	Link	Link	Link
1.5	rs5746059	(A;G)	Slightly higher fat mass			
1.5	rs6601764	(C;T)	1.16x increased risk of developing Crohn's dise		Link	
1.5	rs6710341	(A;G)	Slightly increased risk of developing restless			
1.5	rs7341475	(G;G)	1.58x increased schizophrenia risk for women		Link	
1.5	rs807701	(C;T)	Slightly increased dyslexia risk			
1.5	rs872071	(G;G)	~1.5x increased risk for chronic lymphocytic le		Link	
1.5	rs9303277	(C;T)	1.46x Slightly increased risk of developing pri			
1.5	rs9652490	(A;G)	Slightly increased risk of developing Parkinson		Link	
1.4	rs12770228	(A;G)	1.4x increased risk for meningioma			
1.4	rs1447295	(A;C)	1.4x increased risk of prostate cancer		Link	
1.4	rs1801157	(A;G)	1.4x higher risk for breast cancer			
1.4	rs3131296	(G;G)	1.4x increased risk for schizophrenia		Link	
1.4	rs3184504	(C;T)	Slightly increased risk for celiac disease	Link	Link	
1.4	rs6010620	(G;G)	1.4x higher risk for glioma development; but th		Link	
1.3	rs1042713	(A;G)	1.3x increased risk that pediatric inhaler use	Link	Link	Link
1.3	rs10947262	(C;C)	1.3x increased risk for osteoarthritis			
1.3	rs110419	(A;G)	1.3x increased risk for neuroblastoma			
1.3	rs1375144	(C;T)	1.32x increased risk of developing bipolar diso			
1.3	rs1434536	(A;G)	1.29x increased breast cancer risk			
1.3	rs16847548	(C;T)	1.3x increased risk for sudden cardiac death in			
1.3	rs1746048	(C;C)	1.03 increased risk for coronary heart disease		Link	
1.3	rs2024513	(A;G)	1.3x higher risk for schizophrenia (among Han C			
1.3	rs2059693	(C;T)	1.3x increased risk for testicular cancer			
1.3	rs2736100	(G;T)	1.3x higher risk for glioma development: 2.1x r		Link	
1.2	rs1344706	(T;T)	1.2x increased risk for schizophrenia		Link	
1.2	rs143383	(C;T)	1.1x increased risk for osteoarthritis		Link	Link

Mag.	Identifier	Genotype	Summary	ExAC	GetEvidence	ClinVar
1.2	rs1800693	(A;G)	Slight (1.2x) increase in risk for multiple scl	Link	Link	Link
1.2	rs2072590	(G;T)	1.2x increased risk for ovarian cancer			
1.2	rs2252586	(A;G)	1.2x higher risk for glioma development			
1.2	rs419788	(A;G)	2.0x risk for lupus	Link		
1.2	rs4496877	(T;T)	For type-1 diabetics: 1.6x increased nephropath			
1.2	rs4686484	(A;A)	Slightly increased risk for celiac disease			
1.2	rs4795067	(A;G)	Slight increase in risk for psoriatic arthritis			
1.2	rs498872	(C;T)	1.2x higher risk for glioma development		Link	
1.2	rs8050136	(A;C)	1.2x increased risk for T2D in some populations		Link	
1.17	rs3802842	(A;C)	1.17x increased risk of colorectal cancer		Link	
1.15	rs748404	(C;T)	Very slightly increased risk (1.15) for lung ca		Link	
1.1	rs11037909	(C;T)	1.27x type II diabetes risk	Link		
1.1	rs11110912	(C;C)	1.3x high blood pressure risk			
1.1	rs1800450	(A;G)	Carrier of mannose binding deficiency but of lo	Link	Link	Link
1.1	rs2295190	(G;T)	Slightly increased risk for ovarian cancer in w	Link	Link	Link
1.1	rs2651899	(A;G)	1.1x higher risk for migraines			
1.1	rs2653349	(G;G)	2-6x increased risk for cluster headaches	Link	Link	
1.1	rs34516635	(G;G)	Less longevity for Ashkenazi Jewish women.	Link		Link
1.1	rs3740878	(A;G)	1.26x type II diabetes risk	Link		Link
1.1	rs3818361	(C;T)	1.15x increased risk for late-onset Alzheimer's			
1.1	rs4324715	(C;T)	1.5x increased testicular cancer risk for men			
1.1	rs6897876	(C;T)	Slight increase in testicular cancer risk for m			
1.1	rs7412	(C;C)	More likely to gain weight if taking olanzapine	Link	Link	Link
1.1	rs925391	(C;C)	More likely to go bald; common			
1	rs10504861	(G;G)	Major allele: normal risk of migraine			
1	rs1804197	(A;C)	Increased risk of familial colorectal cancer an			Link
1	rs2282679	(A;C)	Somewhat lower vitamin D levels			
1	rs3194051	(A;G)	1.12x risk of type-1 diabetes	Link	Link	Link
1	rs6932590	(T;T)	1.1x increased risk for schizophrenia		Link	
1	rs798766	(T;T)	Increased susceptibility urinary bladder cancer			
1	rs987525	(A;C)	2.5x increased risk for cleft lip		Link	
0.1	rs601338	(G;G)	Susceptible to Norovirus infections	Link	Link	Link
0	rs1004819	(C;C)	1.5x risk of Crohn's disease		Link	
0	rs10239794	(T;T)	>1.3x risk for ALS			
0	rs10761659	(A;A)	1.5x risk of Crohn's disease		Link	
0	rs3761418	(A;A)	1.3x increased risk for depression			
0	rs3813929	(C;C)	Possible weight gain if taking olanzapine		Link	Link
0	rs4293393	(T;T)	1.25x Increased Risk of CKD for T allele in			
0	rs440446	(G;G)	Increased risk in men for biliary conditions	Link		
0	rs4712653	(T;T)	2x increased risk for neuroblastoma			
0	rs6314	(C;C)	Higher risk for RA	Link	Link	
0	rs7787082	(G;G)	7x less likely to respond to certain antidepres		Link	

3.3 Genosets (Multi-variant Phenotypes)

Magnitude	Identifier	Summary
3.1	gs191	Problem metabolizing NSAIDs
3	gs127	Intermediate warfarin metabolizer
2.5	gs155	CYP3A5 non-expressor
2.5	gs161	CYP2C9 Intermediate Metabolizers
2.5	gs242	Increase risk of prostate cancer patients dying
2.5	gs281	Part of the 88% of the population claimed not t
2.5	gs285	You will lose 2.5x as much weight on a low fat
2.4	gs297	Lower heart attack risk than average
2	gs101	Probably able to digest milk
2	gs104	Restless legs syndrome risk
2	gs154	NAT2 Slow metabolizer
2	gs239	Reduced conversion of beta-carotene to retinol
2	gs246	APOE3/APOE3
2	gs279	Mild trimethylaminuria
1.5	gs247	Parkinson's Disease Risk
0	gs158	CYP1A2 normal metabolizer

4 Raw Data

The raw data used to create this report has been assigned the identifier ERS1176618 in the European Nucleotide Archive (ENA) hosted at the European Bioinformatics Institute (EBI).

These data will not be accessible unless the report is approved. This will happen by default one month after the report is issued, or if the report is approved for immediate release within the one month period. Participants can also withdraw from the study at any time in which case the report and the data will not be released and will be deleted.

If the data has already been released, it can be accessed at: http://www.ebi.ac.uk/ena/data/view/ERS1176618

5 Report Metadata

Resource	Version	Website
Genome	GRCh38	Link
BWA	0.7.12	Link
SAMtools	1.3	Link
GATK	3.4-46	Link
PLINK	v1.90b3.35	Link
VEP	88	Link
SNPedia	30-Jul-2017	Link
ExAC	v0.3.1	Link
GetEvidence	16-Dec-2016	Link
ClinVar	16-Dec-2016	Link

Table 5: Analysis Pipeline Versions

Report generated on August 2, 2017.