

Recommendations: The U.S. Preventative Services Task Force recommends screening for type 2 diabetes in asymptomatic adults with sustained blood pressure (either treated or untreated) greater than 135/80 mmHg.³³

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Markers Tested

Gene/Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
WFS1	rs10010131	G	1.12	Caucasian	67.7%	Validated	18040659
CDKN2B	rs10811661	T	1.27	Asian	57.7%	Validated	18477659
CDKN2B	rs10811661	T	1.20	Caucasian	80.1%	Validated	17463246
MTNR1B	rs10830963	G	1.16	Asian	41.7%	Validated	19241057
MTNR1B	rs10830963	G	1.09	Caucasian	30.0%	Validated	19060907
NOTCH2	rs10923931	T	1.13	Caucasian	9.3%	Validated	18372903
CDKAL1	rs10946398	C	1.16	Caucasian	33.6%	Validated	17463249
HHEX	rs1111875	G	1.27	Asian	31.5%	Validated	18477659
HHEX	rs1111875	G	1.15	Caucasian	58.4%	Validated	17463246
SLC30A8	rs13266634	C	1.17	Asian	53.0%	Validated	18469204
SLC30A8	rs13266634	C	1.12	Caucasian	76.1%	Validated	17463249
IGF2BP2	rs1470579	C	1.18	Asian	26.2%	Validated	18477659
IGF2BP2	rs1470579	C	1.19	Caucasian	29.6%	Validated	17463246
PPARG	rs1801282	C	1.75	Asian	95.2%	Validated	11289058
PPARG	rs1801282	C	1.23	Caucasian	90.3%	Validated	17463249
KCNQ1	rs2237892	C	1.43	Asian	65.5%	Validated	18711367
KCNQ1	rs2237892	C	1.29	Caucasian	92.5%	Validated	18711367
KCNJ11	rs5219	T	1.25	Asian	35.5%	Validated	18162508
KCNJ11	rs5219	T	1.15	Caucasian	36.0%	Validated	17463246
HNF1B	rs7501939	T	1.15	Caucasian	43.4%	Validated	17603484
CDKAL1	rs7754840	C	1.27	Asian	42.3%	Validated	18766326
TCF7L2	rs7903146	T	1.51	African	33.7%	Validated	17601994
TCF7L2	rs7903146	T	1.59	Asian	2.4%	Validated	19012045
TCF7L2	rs7903146	T	1.38	Caucasian	27.9%	Validated	17463246
FTO	rs8050136	A	1.27	Caucasian	46.0%	Validated	17463249
JAZF1	rs864745	T ⁱ	1.10	Caucasian	48.7%	Validated	18372903
ESR1	rs1569788	C	1.25	African	52.9%	Preliminary	18305958
ADIPOQ	rs182052	A	1.24	African	38.7%	Preliminary	19056609
ESR1	rs3020314	C	1.23	Caucasian	25.7%	Preliminary	18854778
Intergenic_6q24	rs4897081	A	1.28	African	51.0%	Preliminary	18560894

^aGene or locus containing the tested marker

^bMarker tested

^cAllele associated with disease risk

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele

^eEthnicity of the population in the corresponding study

^fPercentage of people who have the associated allele in the population studied

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; the PubMed ID (PMID) number identifies the referenced study.

ⁱThis marker can be assayed on either strand of DNA. Therefore, the associated allele could be reported as either an A or a T in the patient report.

Limitations and Warnings: NA

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
250.00 Diabetes mellitus without mention of complication, type II or unspecified type, not stated as uncontrolled	V77.1 Screen-diabetes mellitus

References

1. Ahlqvist E, Ahluwalia TS, Groop L. Genetics of type 2 diabetes. *Clinical chemistry*. 2011;57:241-54.
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Exfoliation glaucoma

Report Type: Health Conditions

About: Exfoliation glaucoma (XFG), also known as pseudoexfoliation glaucoma, accounts for about 12% of glaucoma cases and mainly affects older people.¹ Clinically, XFG is considered a symptom of a systemic condition called exfoliation syndrome (XFS), also known as pseudoexfoliation syndrome. One risk factor for XFS is a family history of the condition,² and research indicates that genetic factors, such as variants in the LOXL1 gene, are also associated with XFG.³

Genetics: The genetic contribution to different forms of glaucoma is believed to include both rare Mendelian mutations and common susceptibility alleles.⁴ For XFG, a common susceptibility allele has been identified.

The genetic marker rs2165241 in the LOXL1 gene, which encodes lysyl oxidase-like protein 1, was found to be associated with XFS in a large study of Caucasians from Iceland and Sweden.³ The LOXL1 protein is believed to assist in the construction and maintenance of the extracellular matrix, which provides structural support to cells. LOXL1 is expressed by many cell types in the eye, and changes in the LOXL1 gene may lead to defective processing of extracellular matrix proteins.³ Approximately 35% of people with European ancestry carry one copy of the risk allele of rs2165241 and about 22% carry two copies. However, the prevalence of XFS in most countries is much lower than the frequency of this allele, indicating that other genetic and environmental factors contribute to risk for XLS. Interestingly, the amount of risk associated with rs2165241 has been observed to be lower in Caucasians in Australia compared to Caucasians in Nordic countries.⁵ The rs2165241 marker is not associated with the risk of developing non-XFS types of glaucoma.³

Recommendations: NA

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Marker Tested

Gene/ Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
LOXL1	rs2165241	T	3.62	Caucasian	43.8%	Preliminary	17690259

^aGene or locus containing the tested marker

^bMarker tested

^cAllele associated with disease risk

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele

^eEthnicity of the population in the corresponding study

^fPercentage of people who have the associated allele in the population studied

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in

genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; the PubMed ID (PMID) number identifies the referenced study.

Limitations and Warnings: NA

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
365.9 Unspecified glaucoma	V80.1 Screening for glaucoma

References

1. Kang JH, Loomis S, Wiggs JL, Stein JD, Pasquale LR. Demographic and geographic features of exfoliation glaucoma in 2 United States-based prospective cohorts. *Ophthalmology*. 2012;119:27-35.
2. Damji KF, Bains HS, Stefansson E, et al. Is pseudoexfoliation syndrome inherited? A review of genetic and nongenetic factors and a new observation. *Ophthalmic genetics*. 1998;19:175-85.
3. Thorleifsson G, Magnusson KP, Sulem P, et al. Common sequence variants in the LOXL1 gene confer susceptibility to exfoliation glaucoma. *Science (New York, N.Y.)*. 2007;317:1397-400.
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Hypertension

Report Type: Health Conditions

About: The prevalence of hypertension, also known as high blood pressure, is estimated to be 29% in the U.S.¹ This condition can lead to stroke, heart attack and kidney failure. Risk factors for hypertension include high salt intake, being overweight and high alcohol consumption.² Research indicates that genetic factors are also associated with risk for the condition.^{3,4,5,6}

Genetics: Over 90% of individuals who develop hypertension are considered to have essential or primary hypertension, meaning that no underlying medical cause can be identified. Hypertension can also be secondary to existing medical problems, such as kidney disease. Some rare forms of hypertension are caused by mutations in single genes. Individuals with these mutations usually have a family history of the disease and hypertension occurs at a younger age. This genetic test focuses on essential hypertension, for which several susceptibility alleles have been identified through large population studies.^{3,4,5,6}

Because of the complex interactions of different physiologic pathways regulating blood pressure, it has been difficult to definitively identify risk alleles in candidate genes for hypertension. Contradictory results have been reported for many genes identified in an early study but not replicated in further research. A few large population-based genome-wide scans and candidate gene association studies have identified two genes as risk factors for hypertension: PPARGC1A and BCAT1.^{3,4,5,6}

The PPARGC1A gene encodes the protein PPAR-gamma coactivator 1,⁷ which regulates the expression of genes involved in several metabolic processes related to blood pressure homeostasis. The Gly482Ser (rs7961152) allele causes an amino acid change in the protein, but the functional consequence of this change is unknown. The Ser allele was associated with decreased risk for essential hypertension in a Danish population.⁵ The same allele was also modestly associated with decreased risk for severe essential hypertension in a Chinese population.⁶

The BCAT1 gene encodes branched-chain aminotransferase 1, which catalyzes the first reaction in the catabolism of the essential branched-chain amino acids leucine, isoleucine and valine. The rs7961152 marker is in a non-coding region of the BCAT1 gene, making it difficult to discern the functional impact of a nucleotide change. The variant may be linked to another causative mutation in the same gene or a neighboring gene. The A allele of the rs7961152 marker was associated with an increased risk for essential hypertension in a study of a British Caucasian population.⁴ In a Korean population, the same allele was moderately associated with increased blood pressure.³

Recommendations: The U.S. Preventive Services Task Force recommends screening for high blood pressure in adults aged 18 and older.⁸

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Markers Tested

Gene/ Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
BCAT1	rs7961152	A	1.29	Asian	16.9%	Preliminary	19424278
BCAT1	rs7961152	A	1.16	Caucasian	46.5%	Preliminary	17554300
PPARGC1A	rs8192678	A	0.60	Asian	40.5%	Preliminary	17971240
PPARGC1A	rs8192678	A	0.70	Caucasian	35.0%	Preliminary	15738346

^aGene or locus containing the tested marker

^bMarker tested

^cAllele associated with disease risk

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele

^eEthnicity of the population in the corresponding study

^fPercentage of people who have the associated allele in the population studied

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; thePubMed ID (PMID) number identifies the referenced study.

Limitations and Warnings: NA

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
401.1 Benign essential hypertension	N/A

References

1. Egan BM, Zhao Y, Axon RN. US trends in prevalence, awareness, treatment, and control of hypertension, 1988-2008. *JAMA*. 2010;303:2043-50.
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8. Screening for high blood pressure in adults. U.S. Preventative Services Task Force web site. <http://www.uspreventiveservicestaskforce.org/uspstf/uspshype.htm#summary>. Updated December 2007. Accessed August 3, 2012.

Multiple sclerosis

Report Type: Health Conditions

About: Multiple sclerosis is a chronic inflammatory disease that is presumed to be an autoimmune disorder.^{1,2} This neurodegenerative disease damages the myelin sheaths surrounding neurons, which leads to muscle weakness, difficulties with balance and numbness, among others.³ Though the cause of multiple sclerosis is unknown, genetic factors are associated with the disease.^{2,4}

Genetics: Genetic variation in the major histocompatibility complex region of Chromosome 6 has long been known to play a role in susceptibility to multiple sclerosis. The human leukocyte antigen (HLA) region, a set of genes within the major histocompatibility complex, has been associated with an increased risk for multiple sclerosis in people of European descent.² The HLA region is also associated with other auto-immune conditions, such as type 1 diabetes and rheumatoid arthritis.^{5,6} Two other genes, interleukin 2 receptor alpha (IL2RA) and interleukin 7 receptor (IL7R), are also involved in the immune system and associated with the risk for multiple sclerosis.² Variants in these genes were examined in families of European descent from the U.S. and U.K. While HLA had the strongest association, both IL2RA and IL7R also appeared to play a role in susceptibility to multiple sclerosis.² The test includes variants in the HLA region, as well as the IL2RA and IL7R genes.

Recommendations: NA

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Markers Tested

Gene/ Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
IL2RA	rs12722489	G	1.25	Caucasian	82.7%	Validated	17660530
HLA	rs3135388	T	1.99	Caucasian	19.0%	Validated	17660530
IL7RA	rs6897932	C	1.18	Caucasian	75.7%	Validated	17660530
KIF1B	rs10492972	C	1.34	Caucasian	33.6%	Preliminary	18997785
EVI5	rs10735781	G	1.11	Caucasian	34.5%	Preliminary	17660530
ANKRD15	rs10975200	G	1.14	Caucasian	16.4%	Preliminary	17660530
FAM69A	rs11164838	C	1.11	Caucasian	58.4%	Preliminary	17660530
CD58	rs12044852	C	1.24	Caucasian	87.2%	Preliminary	17660530
CBLB	rs12487066	T	1.09	Caucasian	67.7%	Preliminary	17660530
PDE4B	rs1321172	G	1.08	Caucasian	54.9%	Preliminary	17660530
KLRB1	rs4763655	A	1.10	Caucasian	33.2%	Preliminary	17660530

^aGene or locus containing the tested marker

^bMarker tested

^cAllele associated with disease risk

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele

^eEthnicity of the population in the corresponding study

^fPercentage of people who have the associated allele in the population studied

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; the PubMed ID (PMID) number identifies the referenced study.

Limitations and Warnings: Multiple sclerosis is rare in non-European populations and, thus, has not been widely studied in people of African or Asian descent. Therefore, only data from studies using large populations of European descent are represented. Genetic risk factors found in Caucasians may well apply to people of other ethnicities, but this has not been proven. Only the most common relapsing-remitting form of multiple sclerosis has been well studied, and all the information provided here relates to relapsing-remitting multiple sclerosis.

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
340 Multiple sclerosis Applies to: <ul style="list-style-type: none"> • Disseminated or multiple sclerosis: <ul style="list-style-type: none"> ◦ NOS ◦ brain stem ◦ cord ◦ generalized 	V82.9 Screening for unspecified condition

References

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6. Fugger L, Svejgaard A. Association of MHC and rheumatoid arthritis. HLA-DR4 and rheumatoid arthritis: studies in mice and men. *Arthritis research*. 2000;2:208-11.

Myocardial infarction

Report Type: Health Conditions

About: Myocardial infarction (MI) kills approximately half a million people in the U.S. each year.¹ Symptoms of this condition include chest pain, shortness of breath and other symptoms.^{1,2} Risk factors include a family history of MI, diabetes, hypertension and hypercholesteremia.² Research indicates that genetic factors are also associated with MI.^{2,3}

Genetics: Genetic susceptibility factors contribute to the risk of MI. Their importance is highlighted by the fact that about 15% to 20% of individuals who have an MI lack conventional risk factors.⁴ Studies suggest that many genetic variations associated with the disease are in genes involved in processes involving endothelial cell function, inflammation, lipid metabolism, thrombosis and fibrinolysis.^{2,3}

Many SNPs associated with heart disease are in genes involved in inflammation. A marker in the CXCL12 gene, which encodes a chemokine molecule important for attracting lymphocytes,⁵ was shown to be also associated with the risk for MI in a large study of Caucasians.⁶ Additionally, in a large genome-wide association study identifying markers affecting eosinophil counts, a marker at 12q24 (rs3184504) in the SH2B3 gene, which encodes a protein important for cytokine pathway inhibition, was associated with risk for MI in six different populations.⁷

A number of high-impact studies have identified 9p21.3 as an important genetic region associated with risk for MI or coronary artery disease.⁸ This region contains no annotated genes but lies close to the CDKN2A and CDKN2B genes, which regulate important cell cycle pathways. An allele of the rs10757278 marker in the 9p21.3 region is associated with MI in Icelandic, U.S. Caucasian, and Italian populations.^{9,10,11} When the Italian patient group was subdivided into those with and without a family history of MI, only the individuals with a family history of MI still showed a significant association with the allele.¹¹ This research suggests that in cases with a family history of heart disease and heart attack, the relative risk for an individual carrying a risk allele can be significantly higher.

Another large study carried out by the Myocardial Infarction Genetics Consortium replicated the region of 9p21, as well as identified or replicated other markers in MIA3 (rs17465637), 1p13 (rs646776), 10q11 in CXCL12 (rs1746048), 21q22 (rs9982601), 6p24 in PHACTR1 (rs12526453), and 2q33 in WDR12 (rs6725887).⁶

Recommendations: NA

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Markers Tested

Gene/Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
PSMA6	rs1048990	G	1.21	Asian	37.8%	Validated	16845397
Intergenic_9p21	rs10757278	G	1.28	Caucasian	50.0%	Validated	17478679
PCSK9	rs11206510	T	1.15	Caucasian	83.6%	Validated	19198609
PHACTR1	rs12526453	C	1.12	Caucasian	63.3%	Validated	19198609
CXCL12	rs1746048	C	1.17	Caucasian	85.4%	Validated	19198609
MIA3	rs17465637	C	1.14	Caucasian	26.8%	Validated	19198609
SH2B3	rs3184504	T	1.13	Caucasian	44.5%	Validated	19198610
Intergenic_1p13	rs646776	T ⁱ	1.19	Caucasian	74.6%	Validated	19198609
WDR12	rs6725887	C	1.17	Caucasian	15.9%	Validated	19198609
LGALS2	rs7291467	C ^j	1.23	Asian	73.3%	Validated	15129282
Intergenic_21q22	rs9982601	T	1.20	Caucasian	20.6%	Validated	19198609
LTA	rs1041981	A	1.78	Asian	43.5%	Preliminary	12426569
OR13G1	rs1151640	G	1.31	Caucasian	46.0%	Preliminary	16175505
PRR4	rs1376251	C	1.23	Caucasian	65.3%	Preliminary	16175505
MIAT	rs2331291	T	1.38	Asian	13.1%	Preliminary	17066261

^aGene or locus containing the tested marker.

^bMarker tested.

^cAllele associated with disease risk.

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele.

^eEthnicity of the population in the corresponding study.

^fPercentage of people who have the associated allele in the population studied.

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; the PubMed ID (PMID) number identifies the referenced study.

ⁱThis marker can be assayed on either strand of DNA. Therefore, the associated allele could be reported as either an A or a T in the patient report.

^jThis marker can be assayed on either strand of DNA. Therefore, the associated allele could be reported as either a C or a G in the patient report.

Limitations and Warnings: NA

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
410.9 Acute myocardial infarction of unspecified site Applies to: <ul style="list-style-type: none"> • Acute myocardial infarction NOS • Coronary occlusion NOS • Myocardial infarction NOS 	V81.2 Screening for other and unspecified cardiovascular conditions

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Obesity

Report Type: Health Conditions

About: Obesity (BMI > 30 kg/m²) affects at least 20% of individuals in Western countries, while 50% of people are classified as overweight (BMI > 25 kg/m²) or obese by the World Health Organization's definition. This condition is characterized by an increase in fat mass that can result in adverse health consequences. Obesity is associated with increased risks for cardiovascular disease, type 2 diabetes and various types of cancer. Risk factors for obesity include low physical activity and consumption of high-energy foods. Research indicates that genetic factors are also associated with the disease.¹

Genetics: Some rare inherited forms of obesity are caused by a mutation in a single gene.¹ However, these monogenic forms of obesity only account for a small fraction of obesity cases. The genetic predisposition for obesity is thought to arise from multiple, common variants in several genes.^{2,3} These common variants are referred to as low risk susceptibility alleles because they each contribute a relatively small amount of risk to developing obesity. Low risk susceptibility alleles for obesity have been successfully identified by screening large numbers of individuals in genome-wide association studies.^{4,5}

The FTO gene was the first gene shown to be associated with common obesity in genome-wide association studies.⁵ This association was quickly replicated in many other studies.^{6,7} How FTO affects obesity is not understood,⁸ but studies in rodents suggest that FTO has a role in regulating food intake. Mice genetically engineered to overexpress FTO show increased food intake, which leads to obesity.⁹ The function of FTO in the central nervous system is also supported by the finding that human children carrying an obesity-associated FTO variant have increased energy intake.¹⁰ Besides its effects in the brain, the FTO protein may also have functions in other tissues where it is also expressed.⁸

Genome-wide association studies have also implicated the MC4R gene in obesity.⁴ The MC4R gene encodes the melanocortin 4 receptor, which is expressed in neurons that modulate food intake. Interestingly, rare mutations in this gene can cause monogenic obesity.

Recommendations: The U.S Preventative Services Task Force recommends screening all adults and children aged 6 years and older for obesity.¹¹

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Markers Tested

Gene/Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
MC4R	rs17782313	C	1.12	Caucasian	26.5%	Validated	18454148
FTO	rs9939609	A	1.31	Caucasian	46.0%	Validated	17434869
PCSK1	rs6232	G	1.34	Caucasian	4.4%	Preliminary	18604207
INSIG2	rs7566605	C	1.29	Caucasian	26.5%	Preliminary	17465681
MC4R	rs17782313	C	1.24	Asian	24.4%	Validated	23049848
FTO	rs9939609	A	1.25	Asian	14.9%	Validated	22109280
PCSK1	rs6232	G	N/A	Asian	1.2%	Preliminary	19875984
INSIG2	rs7566605	C	N/A	Asian	33.7%	Preliminary	25028659

^aGene or locus containing the tested marker.

^bMarker tested.

^cAllele associated with disease risk.

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele.

^eEthnicity of the population in the corresponding study.

^fPercentage of people who have the associated allele in the population studied.

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in genetic research and have not met our highest standards for validation. The outcomes for this test are derived using genotype information from validated markers only.

^hPubMed is a service managed by the National Library of Medicine; the PubMed ID (PMID) number identifies the referenced study.

Limitations and Warnings: NA

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
278.00 Obesity, unspecified	V77.8 Screening for obesity
Applies to:	
<ul style="list-style-type: none"> Obesity NOS 	

References

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Osteoarthritis

Report Type: Health Conditions

About: Osteoarthritis is the most common form of arthritis and is a major cause of disability in the U.S.¹ This progressive bone disease is characterized by cartilage loss, which can lead to pain and reduced joint function. Risk factors for osteoarthritis include age, obesity, gender and trauma. Research indicates that genetic factors are also associated with the disease.²

Genetics: Some rare forms of early-onset osteoarthritis (OA) are caused by mutations in single genes, but uncovering the genetic basis of the most common form of OA, which appears after age 45, has been more elusive. Variants in three genes that have been shown to increase the risk of developing OA are included on the test: GDF5, DVWA and PTGS2.

The GDF5 gene encodes a member of the transforming growth factor-beta superfamily and is involved in the development and maintenance of bone and cartilage. Mutations in GDF5 are known to cause disorders of skeletal development including chondrodysplasia, synphalangism and type C brachydactyly. In the largest meta-analysis study of OA to date, researchers found an association of a variant in the GDF5 gene with OA of the knee in Caucasian and Asian women.³

The DVWA gene encodes a protein that binds beta-tubulin, which is the building block of the microtubules that serve a structural and kinetic role in the cell. In Asians, a variant in the DVWA gene is associated with susceptibility to OA of the knee.⁴

The PTGS2 gene encodes prostaglandin G/H synthase 2, which is involved in a key step in the synthesis of prostaglandins. Prostaglandins are regulators of important biological processes such as inflammation, cell division, and formation of new blood vessels. In Caucasians, a variant in the PTGS2 gene is associated with OA in the knee.⁵

Recommendations: NA

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Markers Tested

Gene/Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
GDF5	rs143383	T	1.55	Asian	77.9%	Preliminary	18299287
GDF5	rs143383	T	1.13	Caucasian	66.7%	Preliminary	19479880
PTGS2	rs4140564	C	1.55	Caucasian	8.0%	Preliminary	18471798
DVWA	rs7639618	G ⁱ	1.43	Asian	54.2%	Preliminary	19181678

^aGene or locus containing the tested marker.

^bMarker tested.

^cAllele associated with disease risk.

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele.

^eEthnicity of the population in the corresponding study.

^fPercentage of people who have the associated allele in the population studied.

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; the PubMed ID (PMID) number identifies the referenced study.

ⁱThis marker can be assayed on either strand of DNA. Therefore, the associated allele could be reported as either a C or a G in the patient report.

Limitations and Warnings: NA

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
715.89 Osteoarthritis involving, or with mention of more than one site, but not specified as generalized, multiple sites	N/A

References

1. Murphy L, Schwartz TA, Helmick CG, et al. Lifetime risk of symptomatic knee osteoarthritis. *Arthritis and rheumatism*. 2008;59:1207-13.
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Parkinson's disease

Report Type: Health Conditions

About: By the year 2030, the prevalence of Parkinson's disease (PD) is projected to double from numbers in 2005 (4.1 to 4.6 million).¹ This neurodegenerative disease is characterized by tremors, akinesia, bradykinesia and balance difficulties.² Risk factors include age and head trauma,³ and research indicates that genetic factors, such as a variant in the LRRK2 gene, are also associated with the disease.²

Genetics: The genetic contribution to PD includes high-risk genetic mutations as well as low-risk susceptibility factors.² Familial PD, which accounts for 15% to 25% of individuals with PD, is caused by rare, high-penetrance mutations (LRRK2, SNCA, parkin, UCHL1, PINK1 and DJ-1). Most PD patients, however, do not have known high-risk mutations, have no family history of the disease and are classified as having sporadic or idiopathic PD. Compared to familial PD, individuals with sporadic PD tend to have a later age of onset but still exhibits similar clinical and pathological features.⁴

The test includes a common variant (rs34778348) in the LRRK2 gene that is associated with sporadic PD in Asians.^{5,6} The LRRK2 (leucine-rich repeat kinase 2) gene, also known as dardarin, encodes a kinase that is found in the brain. High-risk genetic mutations in LRRK2 are pathogenic for familial PD in families of European descent.⁷ In Asians, high risk PD-causing mutations of the LRRK2 gene appear to be extremely rare.

The A allele of rs34778348 in the LRRK2 gene has been found in PD patients in three Asian populations (Han Chinese, Japanese and Malay) but has not yet been found in other ethnic populations. This allele is associated with only a small increase in PD risk. Thus, rs34778348 is considered a low-risk susceptibility marker. Many people who carry the A allele may not develop PD. The A allele leads to an LRRK2 protein that causes cell death only when combined with oxidative stress.⁸ This finding is consistent with the idea that the A allele may result in PD only when additional environmental, lifestyle or other personal risk factors are present.

Recommendations: NA

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Marker Tested

Gene/ Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
LRRK2	rs34778348	A	2.55	Asian	3.0%	Validated	19343804

^aGene or locus containing the tested marker.

^bMarker tested.

^cAllele associated with disease risk.

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele.

^eEthnicity of the population in the corresponding study.

^fPercentage of people who have the associated allele in the population studied.

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; the PubMed ID (PMID) number identifies the referenced study.

Limitations and Warnings: The test does not include high-penetrance pathogenic mutations that cause PD and are inherited in a dominant manner. Most notably, the test does not include pathogenic mutations of the LRRK2 gene, which have been identified in PD families of European descent.⁷

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
332.0 Paralysis agitans	N/A
Applies to:	
<ul style="list-style-type: none"> • Parkinsonism or Parkinson's disease: <ul style="list-style-type: none"> ◦ NOS ◦ idiopathic ◦ primary 	

References

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Peripheral arterial disease

Report Type: Health Conditions

About: Peripheral arterial disease (PAD) affects 20% of individuals in the U.S. over the age of 55 and is strongly associated with a risk for myocardial infarction and stroke.¹ This progressive disease is characterized by the accumulation of plaque in the arterial system, especially the abdominal aorta and arteries in the legs.^{1,2} Risk factors for PAD include age, sex, smoking, diabetes, hypertension and hyperlipidemia.¹ Research indicates that genetic factors are also associated with PAD.³

Genetics: Peripheral arterial disease (PAD) is caused by atherosclerosis, a complex disorder involving both genetic risk factors as well as traditional risk factors, such as cigarette smoking, cholesterol levels, hypertension and stress. Within various populations, genetic differences are thought to account for greater than 50% of the risk for atherosclerosis.³ It is estimated that the genetic risk of atherosclerosis involves variants in hundreds of genes with a variety of functions in regulating blood pressure, lipid and cholesterol metabolism, pro-inflammatory processes, cell adhesion and migration.³

The CHRNA3 gene encodes a subunit of the nicotinic acetylcholine receptor, which binds to nicotine.⁴ A large study of PAD in Caucasians from Iceland, New Zealand, Austria, Sweden and Italy⁵ showed that a variant in CHRNA3 increased the risk for PAD by approximately 20%. No significant differences in risk were shown between males and females. The results from this study suggest that this variant may also partly confer a risk for cardiovascular disease indirectly through its relationship with exposure to nicotine as a function of smoking history.

Recommendations: NA

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Marker Tested

Gene/ Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
CHRNA3	rs1051730	T	1.19	Caucasian	38.5%	Validated	18385739

^aGene or locus containing the tested marker

^bMarker tested

^cAllele associated with disease risk

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele

^eEthnicity of the population in the corresponding study

^fPercentage of people who have the associated allele in the population studied

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in

genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; the PubMed ID (PMID) number identifies the referenced study.

Limitations and Warnings: NA

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
443.9 Peripheral vascular disease, unspecified Applies to: <ul style="list-style-type: none"> • Intermittent claudication NOS • Peripheral: <ul style="list-style-type: none"> ◦ angiopathy NOS ◦ vascular disease NOS • Spasm of artery 	V81.2 Screening for other and unspecified cardiovascular conditions

References

1. Muir RL. Peripheral arterial disease: Pathophysiology, risk factors, diagnosis, treatment, and prevention. *Journal of vascular nursing : official publication of the Society for Peripheral Vascular Nursing*. 2009;27:26-30.
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Psoriasis

Report Type: Health Conditions

About: Psoriasis is a common skin disease that is characterized by itchy, scaly skin and is caused by a defect in the immune system.¹ Risk factors for psoriasis include smoking, alcohol consumption, family history, diet, BMI, infections and stress.² Research indicates that genetic factors are also associated with the disease.³

Genetics: Psoriasis has long been known to have a heritable component, with the siblings of an individual with psoriasis having a significantly greater likelihood of also developing the condition compared to unrelated individuals. For example, a study in Australia looked at psoriasis in almost 4,000 twins.⁴ They found that if one twin had psoriasis, the probability that the second twin also had the condition was 35% if they were identical, but only 12% if they were fraternal twins.

As with other autoimmune conditions, the genes in the HLA (human leukocyte antigen) region of chromosome 6 strongly influence susceptibility to psoriasis. Variants in other genes related to the immune system have also been implicated in the risk of developing the disorder.

One of the largest studies looking at HLA and psoriasis identified an association between variants in the HLA region of Chromosome 6 and susceptibility to psoriasis. HLA was also found to play a role in susceptibility to psoriatic arthritis, a subtype of psoriasis.³ The study also revealed associations between psoriasis and genes involved in the body's immune response, namely interleukin genes IL12B and IL23R. A study by a consortium of academic groups in the US and Germany that specifically looked at interleukin genes confirmed the association with psoriasis.⁵ This report also examined whether the interleukin genes were interacting with genes in the HLA region but found no evidence of such an interaction, suggesting that the HLA and the interleukin genes act as independent risk factors for psoriasis.

Recommendations: NA

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Markers Tested

Gene/Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
HLA	rs10484554	T	2.80	Caucasian	13.4%	Validated	18369459
IL23R	rs11209026	G	1.40	Caucasian	95.9%	Validated	18219280
TNIP1	rs17728338	A	1.59	Caucasian	8.0%	Validated	19169254
STAT2	rs2066808	A ⁱ	1.34	Caucasian	92.9%	Validated	19169254
IL12B	rs3212227	A	1.62	Caucasian	81.0%	Validated	18219280
TNFAIP3	rs610604	G ^j	1.19	Caucasian	42.5%	Validated	19169254
Intergenic_1q21	rs4112788	C	1.41	Caucasian	59.7%	Preliminary	19169253
SPATA2	rs495337	C	1.25	Caucasian	57.0%	Preliminary	18364390

^aGene or locus containing the tested marker.

^bMarker tested.

^cAllele associated with disease risk.

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele.

^eEthnicity of the population in the corresponding study.

^fPercentage of people who have the associated allele in the population studied.

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; the PubMed ID (PMID) number identifies the referenced study.

ⁱThis marker can be assayed on either strand of DNA. Therefore, the associated allele could be reported as either an A or a T in the patient report.

^jThis marker can be assayed on either strand of DNA. Therefore, the associated allele could be reported as either a C or a G in the patient report.

Limitations and Warnings: NA

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
696.1 Other psoriasis	N/A
Applies to: <ul style="list-style-type: none"> • Acrodermatitis continua • Dermatitis repens • Psoriasis: <ul style="list-style-type: none"> ◦ NOS ◦ any type, except arthropathic 	

References

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Rheumatoid arthritis

Report Type: Health Conditions

About: Rheumatoid arthritis (RA) affects approximately 1% of the general population in the U.S.¹ Symptoms of this autoimmune disease include joint pain, stiffness and swelling.² Risk factors for RA include diet, smoking and age, among others.³ Research indicates that genetic factors are also associated with the disease.^{4,5}

Genetics: RA is known to have a heritable component. As with other auto-immune conditions, the genes in the HLA (human leukocyte antigen) region of chromosome 6, strongly influence susceptibility to rheumatoid arthritis. The HLA Class II region contains genes that are part of the major histocompatibility complex. HLA Class II molecules are strongly associated with development of RA in all ethnic groups and account for 30% to 50% of the overall genetic risk in RA.⁶

Other genes shown to be associated with RA are also involved in the immune response. Among these, the CTLA4 (Cytotoxic T-Lymphocyte Antigen 4) gene encodes a protein that plays a key regulatory role in the immune system, particularly in T-cells. A variant in the rs3087243 marker is associated with decreased production of soluble CTLA-4.^{4,5} Abatacept is a drug used to treat RA that mimics the function of CTLA-4 by binding and inhibiting receptors on T-cells.⁷ The CTLA-4 variant and RA is rare in African populations.⁸

The CD40 and TRAF1 genes encode proteins that act together to inhibit T-cells through intracellular signaling pathways. CD40 is also involved in the development of “memory” in B-cells and is found in the synovial fluid of RA patients. A variant of the rs4810485 marker is associated with a change in the efficiency of CD40 protein production.⁴ A variant of the rs3761847 marker may impact TRAF1 protein levels.⁹

The FCRL3 gene encodes an immune receptor that has activating and inhibitory functions. Mutations in this gene are associated with RA, systemic lupus erythematosus and autoimmune thyroid disease. A variant in the rs7528684 marker may regulate FCRL3 protein levels in Asians. Evidence of a similar association has been found for some Caucasian subgroups.¹⁰

The CD244 gene encodes a receptor found on T-cells that regulates their cell-killing function. A variant in the rs3766379 marker is associated with RA in Asians.¹¹ The marker is located in a non-coding region of the gene, and the effect of the variant on protein function is unknown.

Recommendations: NA

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Markers Tested

Gene/Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
PADI4	rs2240340	A	1.31	Asian	41.1%	Validated	18087673
PTPN22	rs2476601	A	1.53	Caucasian	11.7%	Validated	17982455
CTLA4	rs3087243	G	1.22	Asian	80.4%	Validated	19404967
CTLA4	rs3087243	G	1.11	Caucasian	54.0%	Validated	18794853
TRAF1	rs3761847	G	1.32	Caucasian	47.8%	Validated	17804836
CD244	rs3766379	T	1.31	Asian	48.2%	Validated	18794858
MMEL1	rs3890745	T ⁱ	1.12	Caucasian	66.8%	Validated	18794853
CD40	rs4810485	G	1.15	Caucasian	75.2%	Validated	18794853
HLA	rs6457617	T	2.36	Caucasian	52.2%	Validated	17554300
Intergenic_4q27	rs6822844	G	1.28	Caucasian	85.4%	Validated	19404967
Intergenic_6q23	rs6920220	A	1.24	Caucasian	16.5%	Validated	18794853
STAT4	rs7574865	T	1.27	Asian	35.2%	Validated	19404967
STAT4	rs7574865	T	1.24	Caucasian	23.4%	Validated	19404967
IL1B	rs16944	G	1.10	Caucasian	64.2%	Preliminary	18838388
FCRL3	rs7528684	G	1.16	Asian	40.5%	Preliminary	18087673

^aGene or locus containing the tested marker.

^bMarker tested.

^cAllele associated with disease risk.

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele.

^eEthnicity of the population in the corresponding study.

^fPercentage of people who have the associated allele in the population studied.

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; the PubMed ID (PMID) number identifies the referenced study.

ⁱThis marker can be assayed on either strand of DNA. Therefore, the associated allele could be reported as either an A or a T in the patient report.

Limitations and Warnings: NA

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing

laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
714.0 Rheumatoid arthritis	V82.1 Screening for rheumatoid arthritis
Applies to: <ul style="list-style-type: none"> • Arthritis or polyarthritis: <ul style="list-style-type: none"> ◦ atrophic ◦ rheumatic (chronic) 	

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Systemic lupus erythematosus

Report Type: Health Conditions

About: Systemic lupus erythematosus (SLE) is the most common form of lupus.¹ This chronic autoimmune disease is characterized by the production of autoantibodies that can cause renal failure, arthritis, thrombosis, vasculitis and seizures.² Risk factors for SLE include exposure to silica dust, hormonal and reproductive factors and cigarette smoke.^{2,3} Research indicates that genetic factors are also associated with the disease^{4,5,6}

Genetics: Systemic lupus erythematosus (SLE or lupus) can occur either sporadically or be recurrent in families. Genes connected to the immune system are associated with the development of SLE, including the HLA Class II molecules of the major histocompatibility complex (MHC) and genes involved in the complement system.

The rs2187668 marker is located in the HLA Class II region and is linked to a high-risk set of alleles in three tightly linked genes in Caucasians.⁴ The high risk alleles are HLA-DRB1*0301-HLA-DQA1*0501-DQB1*0201. This combination of alleles is very common, but it is the pairing of these alleles with others that can initiate SLE as well as other autoimmune diseases. The minor allele, "A", of rs2187668 has been associated with increased SLE risk in Caucasians.

Other genes included in the test are also involved in the immune system. Among these, the FCGR2A gene encodes a receptor that is expressed on immune cells and interacts with antibody complexes. The T allele of the rs1801274 marker causes an altered interaction with IgG and is thought to perpetuate autoimmune responses in SLE in Caucasians and Asians.^{5,6}

In another example, the IRF5 gene encodes a transcription factor that regulates other genes that respond to interferon-alpha (IFN alpha). IFN alpha is normally produced in response to viral infections, but its levels are abnormally high in SLE patients. Increased IFN alpha levels result in increased IRF5 levels. The T allele of the rs2004640 marker encodes an alternative form of IRF5 and is associated with increased risk for SLE in individuals of Caucasian, Asian, African and Hispanic descent.⁷ IRF5 has also been shown to have additive effects with STAT4 for increased risk for SLE.

The STAT4 gene encodes a transcription factor that regulates inflammatory gene expression. The T allele of the rs7574865 marker is associated with increased STAT4 expression and is thought to contribute to earlier onset and more severe disease in Caucasians, Asians and people of Hispanic ancestry.^{8,9,10}

Recommendations: NA

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Markers Tested

Gene/ Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
ITGAM	rs1143679	A	1.55	African	11.0%	Validated	18204448
ITGAM	rs1143679	A	1.78	Caucasian	10.0%	Validated	18204448
ITGAM	rs1143679	A	2.26	Hispanic	9.0%	Validated	19129174
TNFSF4	rs1234314	G	1.26	Caucasian	42.5%	Validated	19092840
BLK	rs13277113	A	1.30	Asian	75.0%	Validated	19225526
BLK	rs13277113	A	1.39	Caucasian	23.5%	Validated	18204098
BANK1	rs17266594	T	1.64	Asian	84.5%	Validated	19357697
BANK1	rs17266594	T	1.42	Caucasian	73.9%	Validated	18204447
MECP2	rs1734787	C	1.55	Asian	88.5%	Validated	18320046
MECP2	rs1734787	C	1.35	Caucasian	18.5%	Validated	19333917
FCGR2A	rs1801274	C	1.87	African	45.3%	Validated	12115187
FCGR2A	rs1801274	C	1.61	Asian	31.5%	Validated	12867584
FCGR2A	rs1801274	C	1.24	Caucasian	50.9%	Validated	12115187
HLA	rs2187668	A	1.76	Caucasian	8.0%	Validated	19493061
CTLA4	rs231775	G	1.25	Asian	69.6%	Validated	15688186
PTPN22	rs2476601	A	1.35	Caucasian	11.7%	Validated	19493061
PTPN22	rs2476601	A	2.06	Hispanic	5.2%	Validated	16868974
CTLA4	rs3087243	G	1.32	Caucasian	54.0%	Validated	15248219
TNFAIP3	rs5029939	G	2.09	Caucasian	2.7%	Validated	19387456
STAT4	rs7574865	T	1.71	Asian	35.2%	Validated	19225526
STAT4	rs7574865	T	1.56	Caucasian	23.4%	Validated	18516230
STAT4	rs7574865	T	1.62	Hispanic	40.0%	Validated	18432273
CRP	rs3093062	G	1.45	African	79.9%	Preliminary	18182444

^aGene or locus containing the tested marker.

^bMarker tested.

^cAllele associated with disease risk.

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele.

^eEthnicity of the population in the corresponding study.

^fPercentage of people who have the associated allele in the population studied.

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; the PubMed ID (PMID) number identifies the referenced study.

Limitations and Warnings: NA

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
710.0 Systemic lupus erythematosus	V82.9 Screening for unspecified condition
Applies to:	
<ul style="list-style-type: none"> • Disseminated lupus erythematosus • Libman-Sacks disease 	

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Ulcerative colitis

Report Type: Health Conditions

About: Ulcerative colitis (UC) has been on the rise for the past 50 years, but incidence rates seem to be plateauing.¹ This chronic, idiopathic condition is characterized by ulcers in the rectum and colon that can cause pain and bloody diarrhea.² Risk factors include a family history of the disease and smoking.^{3,4} Research indicates that genetic factors are also associated with UC.^{5,6,7}

Genetics: Both genetics and environmental factors are known to contribute to the risk of developing UC. A role for genetics is shown by the observation that people with a family history of UC have an increased risk of developing the disease. Up to 20% of UC cases occur in families, with a higher incidence in those of northern European and Jewish ancestry.⁸ The genetic contribution to ulcerative colitis (UC) can be divided into two groups: HLA (human leukocyte antigen) and non-HLA susceptibility loci.

Like other autoimmune conditions, there is a strong association between susceptibility to UC and variants of the HLA region on Chromosome 6.⁵ The involvement of HLA genes in UC has long been known, but the exact mechanisms are not well understood. The HLA-associated UC susceptibility loci in the test include the rs2395185 marker in the HLA Class II region⁹ and the rs1800629 marker in the TNFA gene, which encodes the immune stimulatory TNF-alpha protein.¹⁰ The genetic association to TNFA alleles has only been reliably demonstrated in Asians. TNF-alpha is a target of antibody drugs used to treat ulcerative colitis.¹¹

Non-HLA UC susceptibility genes in the test include IL10 (interleukin 10)⁶ and IL23R (interleukin 23 receptor),⁷ among others. Several pro-inflammatory interleukin proteins, including IL10 and IL23, are targets for ulcerative colitis drug therapies that are either available or in clinical development.¹¹

Recommendations: NA

Possible Outcomes: Increased Risk, Above Average Risk, Average Risk

Markers Tested

Gene/Locus ^a	Marker ^b	Associated Allele ^c	Odds Ratio ^d	Ethnicity ^e	Population Frequency ^f	Scientific Strength ^g	PMID ^h
NKX2-3	rs10883365	G	1.20	Caucasian	45.6%	Validated	18438406
IL23R	rs11209026	G	1.81	Caucasian	95.9%	Validated	19122664
IFNG	rs1558744	A	1.35	Caucasian	40.3%	Validated	19122664
HLA	rs2395185	G	1.77	Caucasian	56.7%	Validated	19122664
IL10	rs3024505	T	1.46	Caucasian	18.1%	Validated	18836448
MST1	rs3197999	T	1.20	Caucasian	26.1%	Validated	18438406
RNF186	rs3806308	G	1.37	Caucasian	60.2%	Validated	19122664
Intergenic_1p36	rs6426833	A	1.45	Caucasian	50.9%	Validated	19122664
BSN	rs9858542	A	1.31	Caucasian	25.7%	Validated	18438406
TNFA	rs1800629	A	2.27	Asian	8.9%	Preliminary	18827481

^aGene or locus containing the tested marker

^bMarker tested

^cAllele associated with disease risk

^dMeasure of the likelihood that an individual will get the disease if carrying a specific allele

^eEthnicity of the population in the corresponding study

^fPercentage of people who have the associated allele in the population studied

^gValidated markers represent the highest quality genetic markers available; preliminary markers represent the latest in genetic research and have not met our highest standards for validation.

^hPubMed is a service managed by the National Library of Medicine; thePubMed ID (PMID) number identifies the referenced study.

Limitations and Warnings: NA

Compatible ICD-9 Codes: Insurance coverage is not required for genetic testing. The diagnostic code (ICD-9) information provided herein is for insurance information purposes only and does not guarantee insurance coverage for any genetic test, nor is it intended to be a definitive list of diagnosis codes that may be applicable for any individual patient. The testing laboratory will pursue reimbursement directly from the patient should the patient's insurance carrier refuse to provide coverage.

Primary ICD-9 Code(s)	Screening ICD-9 Code(s)
556.6 Universal ulcerative (chronic) colitis	N/A
Applies to:	
<ul style="list-style-type: none"> Pancolitis 	

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