Supplemental Files

- Section 1 eMethods
- Sections 2 eFigures and eTables
- eFigure 1. Flow chart illustrating the patients screening process
- eTable 1. The classification of the stroke etiology for each gene
- eTable 2. List of the 181 genes associated with Mendelian-stroke in custom-designed panel
- eTable 3. Diagnoses of 759 individuals with 1 P/LP variant at risk for one monogenic diseases
- eTable 4. Diagnoses of 29 individuals harbored more than 2 P/LP variants in different genes
- eTable 5. Four individuals with 2 P/LP variants in the ABCC6 gene
- eTable 6. Characteristics of the Patients with one MGD in C3 cohort
- eTable 7. The characteristics of individuals who had been diagnosed monogenic stroke before gene testing

eMethods

NGS data analysis

Updating PVS1, PS1, PP2, and BP1 lists for InterVar

Since the publishing of InterVar software in 2017, a great deal of newly-found pathogenic mutations has been discovered. To incorporate these progresses, we updated the PVS1, PS1, PP2, and BP1 lists of InterVar using recently released ClinVar database (ClinVar 20200622 version). All of the updating process was conducted using the same pipeline of InterVar or under the guidelines of the American College of Medical Genetics and Genomics and the Association for Molecular Pathology.^[1,2]

PVS1 gene list

Genes in PVS1 list harbored loss-of-function (LOF) mutations that were reported to be pathogenic. First, we extracted all of the pathogenic/likely pathogenic (PLP) and LOF mutations from ClinVar. The genetic variants in ClinVar 20200622 version were regarded to be PLP and LOF mutations if: 1) the variant was recorded by MedGen; 2) its minor allele frequency (MAF) < 5%; 3) it was not predicted or reported to be "benign", "likely benign", or "uncertain significance" in ClinVar; 4) the record of the mutation in ClinVar should not contain the string "conflicting"; 5) the mutations resulted in stopgain, stoploss, frameshift, destruction of canonical ± 1 or 2 splice sites (dbscSNV \geq 0.6), loss of initiation codon, or deletion of single or multiple exons. The mutations that fulfill all of the above 5 criteria were regarded to be PLP and LOF mutations. We found 3312 genes that harbored these mutations, and these genes were included in updated PVS1 gene list.

Second, another part of PVS1 gene list were obtained by retrieving the LOF-mutation-intolerant genes from genomAD. A total of 3075 Genes with pLI ≥0.9 were identified and included in PVS1 gene list. After integration and removing duplications for the 3312 and 3075 genes, a total of 5646 genes were resolved. This list was regarded to be the updated PVS1 gene list, and 150 out of the 181 genes were present in this list.

PS1 mutation list

PS1 list contained nonsynonymous mutations that were reported to be pathogenic. Therefore, we first extracted all of the PLP mutations from ClinVar. The genetic variants in ClinVar 20200622 version were regarded to be nonsynonymous PLP mutations if: 1) the variant was recorded by MedGen; 2) its minor allele frequency (MAF) < 5%; 3) it was not predicted or reported to be "benign", "likely benign", or "uncertain significance" in ClinVar; 4) the record of the mutation in ClinVar should not contain the string "conflicting"; 5) the mutations resulted in nonsynonymous mutation. We found 35767 genetic variants that fulfill all of the above 5 criteria, and they were allocated in the updated PS1 list.

PP2 gene list

In a certain gene, if >80% of the pathogenic variants were missense while <10% of missense variants were benign in ClinVar, the gene would be assigned to the PP2 gene list. Accordingly, the PP2 gene list was obtained through 2 steps. First, for each gene that was recorded by ClinVar, we counted the number of PLP variants and calculated the percentage of nonsynonymous mutations among the PLP variants. The genetic variants in ClinVar 20200622 version were regarded to be PLP mutations if: 1) the variant was recorded by MedGen; 2) its minor allele frequency (MAF) < 5%; 3) it was not predicted or reported to be "benign", "likely benign", or "uncertain significance" in ClinVar; 4) the record of the mutation in ClinVar should not contain the string "conflicting". The mutations that fulfill all of the above 4 criteria were regarded to be PLP mutations. Additionally, nonsynonymous PLP mutations should further fulfill a

5th criterion that the PLP variants would result in nonsynonymous mutations. Then the percentage of pathogenic variants that were missense was obtained for each gene.

In the second step, we counted the number of nonsynonymou variants and calculated the percentage of benign/likely benign (BLB) mutations in each gene. Genetic variants were applied in this calculation if:

1) the variant was recorded by MedGen; 2) its minor allele frequency (MAF) < 5%; 3) the mutations were nonsynonymous substitutions. For BLB variants, another 2 criteria should be fulfilled that: 4) the record of the mutation in ClinVar should not contain the string "conflicting"; 5) it was not predicted or reported to be "pathogenic", "likely pathogenic", or "uncertain significance" in ClinVar record. Then the percentage of nonsynonymous variants that were BLB was obtained for each gene.

Afterwards, based on the above 2 percentages, we 490 genes that fulfilled the criteria of PP2 genes, these genes made up the updated PP2 gene list. A total of 22 genes in the 181 candidate genes were present in the updated PP2 gene list.

BP1 gene list

In a certain gene, if >80% of the pathogenic variants were truncating mutation, the gene would be contained in BP1 gene list. Therefore, for each gene that was recorded by ClinVar, we counted the number of PLP variants and calculated the percentage of truncating mutations. The genetic variants in ClinVar 20200622 version were regarded to be PLP mutations if: 1) the variant was recorded by MedGen; 2) its minor allele frequency (MAF) < 5%; 3) it was not predicted or reported to be "benign", "likely benign", or "uncertain significance" in ClinVar; 4) the record of the mutation in ClinVar should not contain the string "conflicting". The mutations that fulfill all of the above 4 criteria were regarded to be PLP mutations.

Afterwards, we calculated the percentage of PLP mutations that resulted in stopgain, stoploss, frameshift, destruction of canonical ± 1 or 2 splice sites (dbscSNV \geq 0.6), loss of initiation codon, or deletion of single or multiple exons, and the genes with this percentage>80% was rated as BP1 genes. In summary, we found 604 BP1 genes and 9 out of the 181 genes were contained in the updated BP1 gene list.

Phenotyping:

All individuals that harbored pathogenic/likely pathogenic variants that had an expected phenotype based off of the inheritance pattern of the disease, and were not given a clinical diagnosis were classified into the following four categories (Unlikely, Possible, Probable, Definite) based off of the criteria listed below after Electronic Health Record (EHR) review.

Dilated Cardiomyopathy (DCM):

Dilated cardiomyopathy is clinically diagnosed based on echocardiography. Its diagnostic criteria are as follows:^[3] 1. The end-diastolic inner diameter of the left ventricle is greater than 5.5 cm in men and 5.0 cm in women. 2. The ejection fraction is less than 45%, or the left ventricular shortening rate is less than 25%. It is more scientific that the end-diastolic inner diameter of the left ventricle per square meter is greater than 2.7cm. Before diagnosing dilated cardiomyopathy, it is necessary to rule out heart enlargement caused by hypertension, coronary heart disease (CAD), valvular heart disease, congenital heart disease, alcoholic cardiomyopathy, etc.

Definite: N/A

Possible: The end-diastolic inner diameter of the left ventricle, greater than 5.5cm in men and 5.0cm in women

Or the ejection fraction is less than 45%;

Or the left ventricular shortening rate is less than 25%;

Or have heart failure in past history;

Or sudden cardiac death in family history;

Unlikely: Normal cardiac imaging (echocardiography or cardiac MRI) with EF>50% OR cardiomyopathy (EF<45%) with significant CAD*

*Significant CAD was defined as ≥75% stenosis in the left main, proximal left anterior descending coronary arteries, or ≥2 epicardial coronary arteries or history of myocardial infarction Hypertrophic cardiomyopathy (HCM)

HCM is typically defined by the presence of unexplained left ventricular hypertrophy (LVH) with a maximum wall thickness ≥15 mm in adults or a z-score >3 in children. [4-6] If there is a family history of HCM, or if genetic testing confirms that a relative has inherited the family's pathogenic sarcomere variant, a maximum LV wall thickness ≥13 mm supports diagnosis. Such LVH occurs in a non-dilated ventricle in the absence of other cardiac or systemic disease capable of producing the observed magnitude of increased LV wall thickness, such as pressure overload or storage/infiltrative disorders.

- Definite: Septal wall thickness ≥ 1.5 cm on echocardiography with no hypertension, coronary heart disease (CAD), valvular heart disease, congenital heart disease, alcoholic cardiomyopathy.
- Possible: Septal wall thickness ≥ 1.3 cm on echocardiography or <1.3 cm and EF<45% in the absence of an ischemic etiology*
- Unlikely: Septal wall thickness < 1.3 cm on echocardiography and EF>50%

An additional requirement for classification was the presence of at least one echocardiography study *Ischemic etiology was defined as \geq 75% stenosis in the left main OR proximal left anterior descending coronary arteries OR \geq 2 epicardial coronary arteries or a history of myocardial infarction Fabry Disease

Suggestive Findings

Fabry disease should be suspected in males and females with the following clinical features:

Periodic crises of severe pain in the extremities (acroparesthesia)

Vascular cutaneous lesions (angiokeratomas)

Sweating abnormalities (anhidrosis, hypohidrosis, and rarely hyperhidrosis)

Characteristic corneal and lenticular opacities

Unexplained stroke

Unexplained left ventricular hypertrophy

Renal insufficiency of unknown etiology including unexplained proteinuria or microalbuminuria

The diagnosis of Fabry disease is established in a male proband by:

Identification of deficient alpha-galactosidase A (α -Gal A) enzyme activity in plasma, isolated leukocytes, and/or cultured cells. The test is a fluorometric assay and uses the substrate 4-methylumbelliferyl- α -D-galactopyranoside.

Males with classic Fabry disease have <1% α -Gal A enzyme activity.

Males with atypical Fabry disease have residual enzyme activity >1% of normal.

Identification of a hemizygous pathogenic variant in GLA by molecular genetic testing

Female proband. The diagnosis of Fabry disease is established in a female proband by identification of

a heterozygous pathogenic variant in GLA by molecular genetic testing.

- Definite: N/A
- Possible: unexplained left ventricular hypertrophy with a hemizygous pathogenic variant in GLA in Males or a heterozygous pathogenic variant in females.
- Unlikely: No above clinical features with a pathogenic variant in GLA.

Unexplained left ventricular hypertrophy

Familial Transthyretin (TTR) Amyloidosis:

Because there is no formal diagnostic criteria for Familial *TTR* amyloidosis, individuals were classified based on the number of neurologic and/or cardiac signs/symptoms that were documented in the EHR of the respective individual. Individuals that harbored pathogenic/likely pathogenic mutations associated with Familial *TTR* amyloidosis were classified into the four categories based off of the following criteria:^[7]

- Definite: N/A
- Probable: One Cardiac and one Neurologic Manifestation
- Possible: One Cardiac or one Neurologic Manifestation
- Unlikely: Normal Cardiac Imaging and no documented Neurologic signs/symptoms

A cardiac manifestation is defined as documentation of at least one of the following:

- Left ventricular hypertrophy in the absence of an alternative cause
- Heart Failure with preserved ejection fraction
- Restrictive cardiomyopathy
- Conduction disease: Atrioventricular block on electrocardiogram (EKG)
- Presence of a pericardial effusion on echocardiography in the absence of an alternative cause

A neurologic manifestation is defined as documentation of at least one of the following:

- Sensory disturbances of unknown etiology
- Neuropathic pain
- Autonomic disturbances

Atrial fibrillation (AF)

Body surface electrocardiogram (ECG) or 24-hour Holter electrocardiogram shows irregular shape and size of f wave instead of P wave, frequency 350-600 beats/min, QRS complex form is standard form or with ventricular differential conduction and wide deformity, ventricular rate is absolute irregular

- Definite: ECG or 24 Holter showed AF signs and had AF history or AD family history.
- Possible: ECG or 24 Holter showed AF signs or other type of arrhythmia
- Unlikely: ECG or 24 Holter showed normal and no arrhythmia history.

Familial Hypercholesterolemia:

Individuals that harbored a pathogenic/likely pathogenic variant associated with familial hypercholesterolemia who were heterozygous for mutations in the LDLR gene were categorized according to the Simon-Broome criteria as follows:

- Definite: Serum LDL-C*≥190 mg/dL or Total Cholesterol (TC)*≥280 mg/dL
- Possible: After the LDL-C and TC values were corrected, Serum LDL-C*≥190 mg/dL or Total Cholesterol (TC)*≥280 mg/dL.
- Unlikely: Serum LDL-C*<190 mg/dL or Total Cholesterol*<280 mg/dL

*If the patient was on a statin, the LDL-C and TC values were corrected using LDL/0.7 and TC/0.8, respectively.

Marfan Syndrome:

Individuals that harbored pathogenic/likely pathogenic mutations associated with Marfan Syndrome were classified into the four categories based off of the Revised Ghent Nosology:^[8]

- Definite: N/A
- Possible: Systemic Score between 4-6 and a causal FBN1 mutation
- Unlikely: Normal Imaging with no mention of Ectopia Lentis or Systemic Score signs/symptoms
- *Aortic Criterion: History of Aortic Dissection or an Aorta Z-score≥2. Z-scores calculated using the patient's aortic diameter, as measured by echocardiography, using the following Z-score calculator: https://www.marfan.org/dx/zscore.

Supravalvular Aortic Stenosis (SVAS):

Individuals that harbored pathogenic/likely pathogenic variants associated with restrictive cardiomyopathy were classified into the four categories as follows:

- Definite: After aortic valve prosthetic valve replacement surgery
- Possible: Moderate Aortic Stenosis gradient with Normal Aortic Valve on echocardiography
- Unlikely: Normal Imaging

Thoracic aortic aneurysms and aortic dissections (TAAD):

A thoracic aortic aneurysm is a permanent, localized dilatation of the thoracic aorta. Thoracic aortic aneurysms may involve different thoracic aortic segments. To evaluate for a thoracic aortic aneurysm, the aortic diameter is measured (perpendicular to the axis of blood flow) by echocardiography, CT, or MRI at reproducible anatomic locations.

- Definite: After aortic valve prosthetic valve replacement surgery,
- Possible: Moderate Aortic Stenosis gradient with Normal Aortic Valve on echocardiography
- Unlikely: Normal Imaging

Neurofibromatosis 1(NF1)

Suggestive Findings

Neurofibromatosis 1 (NF1) should be suspected in individuals who have any of the following findings:

- Six or more café au lait macules >5 mm in greatest diameter in prepubertal individuals and >15 mm in greatest diameter in postpubertal individuals
- Two or more neurofibromas of any type or one plexiform neurofibroma
- Freckling in the axillary or inguinal regions
- Optic glioma
- Two or more Lisch nodules (iris hamartomas)
- A distinctive osseous lesion such as sphenoid dysplasia or tibial pseudarthrosis
- A first-degree relative (parent, sib, or offspring) with NF1 as defined by the above criteria Establishing the Diagnosis

The diagnosis of NF1 is established in a proband who meets the diagnostic criteria for neurofibromatosis 1 (NF1) developed by the National Institutes of Health [NIH 1988]. The NIH diagnostic criteria for NF1 are met in an individual who has two or more of the features listed in Suggestive Findings.^[9]

- Definite: NA;
- Possible: Individuals with a pathogenic variants in the NF1 gene;
- Unlikely: NA

Polycythemia vera

Major criteria

- 1. Hemoglobin >16.5 g/dL in men or > 16 g/dL in women; or hematocrit >49% in men or > 48% in women or increased red blood cell mass
- 2. Presence of JAK2 mutation

Minor criterion

- 1.Bone marrow trilineage proliferation
- 2. Subnormal serum erythropoietin level
- 3. EEC growth

Bone marrow biopsy might not be needed in the presence of hemoglobin >18.5 g/dL (hematocrit 55.5%) in men or >16.5 g/dL (hematocrit 49.5%) in women.

- Definite: both two major criteria + one minor criterion or the first major + two minor criteria;
- Possible: two major criteria;
- Unlikely: Nomal hemoglobin level

Essential thrombocythemia (ET)

Major criteria^[10]

- 1. Platelets \geq 450 x 10⁹/L
- 2. Bone marrow megakaryocyte proliferation and loose clusters
- 3. Not meeting WHO criteria for other myeloid neoplasms
- 4. JAK2/CALR/MPL mutated
 - Definite: all four major criteria
 - Possible: two major criteria;
 - Unlikely: Nomal Platelets level

Polycystic kidney disease (PKD1 and PKD2)[11]

Definite:

The diagnosis of ADPKD is established in a proband with ANY of the following:

Age-specific ultrasound criteria and an affected first-degree relative with ADPKD

Age-specific MRI criteria and an affected first-degree relative with ADPKD

Identification of a heterozygous pathogenic variant in one of the genes listed in Table 3

Ultrasound Criteria for Diagnosis of ADPKD in Individuals at 50% Risk for ADPKD Based on Family History

Age	PKD1	PKD2	Unknown ADPKD Genotype
	≥3 cysts ¹ PPV = 100% SEN = 94.3%	≥3 cysts ¹ PPV = 100% SEN = 69.5%	≥3 cysts ¹ PPV = 100% SEN = 81.7%
30-39 yrs	≥3 cysts ¹ PPV = 100% SEN = 96.6%	≥3 cysts ¹ PPV = 100% SEN = 94.9%	≥3 cysts ¹ PPV = 100% SEN = 95.5%
40-59 yrs	≥2 cysts in each kidney PPV = 100% SEN = 92.6%	≥2 cysts in each kidney PPV = 100% SEN = 88.8%	≥2 cysts in each kidney PPV = 100% SEN = 90%

Possible:

Multiple bilateral renal cysts and the absence of manifestations suggestive of a different renal cystic disease

Cysts in other organs, especially the liver, but also seminal vesicles, pancreas, and arachnoid membrane

Enlargement of the kidneys or liver on physical examination

Hypertension in an individual younger than age 35 years

An intracranial aneurysm

A family history of ADPKD

Unlikely: NA

Cerebral cavernous malformations-1(KRIT1)

- Intracranial thin-walled sinusoidal vessel (cavernous) malformations
- Seizures
- Headache
- Intracranial hemorrhage
- Focal neurologic deficits
- Intracranial calcifications
- Angiographically 'silent'
- MRI is best imaging modality to detect lesions
 - Definite: MRI showed intracranial thin-walled sinusoidal vessel (cavernous)
 - Possible: NA
 - Unlikely: Normal Imaging

Ehlers-Danlos syndrome (EDS)

Individuals that harbored pathogenic/likely pathogenic variants associated with Ehler-Danlos Syndrome (COL5A1,COL5A2, COL1A1, or COL1A2) were classified into the three categories as follows based off of the 2017 international classification of the Ehlers–Danlos syndromes^[12]

- Definite: NA
- Possible: a proband with the minimal clinical diagnostic criteria (intracranial aneurysms and arteriovenous fistulae, may occur in the rare individual) and identification of a heterozygous pathogenic variant in COL1A1 or COL1A2
- Unlikely: None of the above clinical manifestations

Cerebral amyloid angiopathy, GSN-related

- Cranial neuropathy, esp. facial paresis
- Bulbar palsy
- Peripheral polyneuropathy, esp. vibration and touch loss
- Amyloid cardiomyopathy
- Renal failure
- Cutis laxa
- -Corneal lattice dystrophy
- Definite: NA
- Possible: Diagnosed with CAA by imaging
- Unlikely: None of the above clinical manifestations

Thrombophilia due to thrombin defect duo to F2 defect

- Definite: NA
- Possible: Thrombosis, recurrent, cognitive function loss
- Unlikely: None of the above clinical manifestations

Hyperchylomicronemia, late-onset APOA5

- Definite: NA
- Possible: Decreased LDL and HDL, increased TG, stroke
- Unlikely: None of the above clinical manifestations

HTRA1-autosomal dominant disease

HTRA1 heterozygous mutations may lead to a late-onset syndrome characterized by gait disturbances, mood depression, cognitive impairment, stroke, migraine as well as WMHs on MRI.

- Definite: NA
- Possible: HTRA1 heterozygous mutations and a late-onset stroke as well as WMHs
- Unlikely: Just a HTRA1 heterozygous mutations with none of the above clinical manifestations.

Cerebral Autosomal Dominant Arteriopathy with Subcortical Infarcts and Leukoencephalopathy (CADASIL)^[13]

Clinical criteria

- #1 Age at onset (clinical symptoms #2 or white matter lesions) ≤55 years old.
- #2 At least two of the following clinical findings:
- a. Either of subcortical dementia, long tract signs, or pseudobulbar palsy.
- b. Stroke-like episode with a focal neurological deficit.
- c. Mood disorder.
- d. Migraine.
- #3 Autosomal dominant inheritance.
- #4 White matter lesions involving the anterior temporal pole by MRI or CT.
- #5 Exclusion of leukodystrophy (Adrenoleukodystrophy, metachromatic leukodystrophy, etc.).

Genetic criteria

NOTCH3 mutations localize in exons 2-24 and result in the gain or loss of

cysteine residues in the epidermal growth factor-like repeat domain.

Cysteine-sparing variants should be carefully evaluated by skin biopsy and

segregation studies

Pathological criteria

The pathological hallmark of CADASIL is granular osmiophilic material (GOM)

detected by electron microscopy. Immunostaining of NOTCH3 extracellular

domain is also useful.

Definite:

CADASIL is definite when the individual fulfills

- (1) White matter lesions by MRI or CT.
- (2) Clinical criteria #5
- (3) Genetic criteria and/or pathological criteria

Possible:

CADASIL is possible when the individual has abnormal white matter lesions

(Fazekas grade ≥2) and fulfills either of

- (1) ≤55 years old
- (2) At least one of the symptoms in clinical criteria #2

Unlikely:

No phenotype

Retinal vasculopathy with cerebral leukoencephalopathy and systemic manifestations (RVCL-S)

Major features

- #1 Vascular retinopathy typically manifesting as decreased visual acuity and/or visual field defects
- #2 Focal neurologic signs can include but are not limited to hemiparesis, facial weakness, aphasia, and hemianopsia.
- #3 Global brain dysfunction may manifest as progressive cognitive impairment.
- #4 Brain MRI abnormalities are restricted to the white matter
- (1)Focal, non-enhancing T2-hyperintense lesions scattered throughout the periventricular and deep white matter (at an age when nonspecific age-related white matter hyperintensities are infrequent)
- (2)Punctate T2-hyperintense white matter lesions with nodular enhancement
- (3)Hyperintense mass lesions on T2 and hypointense lesions on T1-weighted images, enhanced with gadolinium contrast, and often surrounded by extensive edema. Hemorrhages are rarely reported. Occasionally, restricted diffusion, most often centrally, is observed and is referred to as a "pseudotumor." #5 Family history of middle-age onset of disease manifestations consistent with an autosomal dominant inheritance pattern.

#6 Exclusion of leukodystrophy AND brain tumor.

Supportive features

Calcifications on brain CT scan, typically not present in healthy controls

Nonspecific MRI white matter lesions that occur more frequently than expected given the age of the individual

Microvascular liver disease, manifested by modest elevations of alkaline phosphatase and gamma-glutamyltransferase

Microvascular kidney disease, typically manifested by a mild-to-moderate increase in serum creatinine or by proteinuria

Pathology

Histologic abnormalities have been demonstrated in all organs involved in RVCL-S, including the following.

- (1)Retina: scattered microinfarcts, thickened hyalinized retinal arterial walls, focal areas of disruption of the ganglion cell layer and inner nuclear layer.
- (2)Brain: Multiple often confluent foci of ischemic necrosis of white matter, Vasculopathy (vessel wall thickening and luminal stenosis; telangiectasias), a modest chronic inflammatory cell infiltrate in some individuals, focal calcifications and reactive astrocytosis, myelin loss.
- (3)Kidney: renal arteriolosclerosis, focal or diffuse glomerulosclerosis
- (4)Liver:Nodular regenerative hyperplasia, micro- and macrovesicular steatosis, periportal inflammation, bridging and portal fibrosis.

Definite:

RVCLS is definite when the individual fulfills

- (1) fulfills either of Major features
- (2) Clinical criteria #6
- (3) Genetic criteria and/or pathological criteria

Possible:

RVSLS is possible when the individual has either of Major features or fulfills more than three of Supportive features

Unlikely:

No phenotype

Brain small vessel disease with or without ocular anomalies (COL4A1)

Major features

- (1)Porencephaly
- (2)Brain small-vessel disease with or without ocular anomalies
- (3)HANAC (hereditary angiopathy with nephropathy, aneurysms, and muscle cramps) syndrome
- (4)Tortuosity of retinal arteries
- (5)Nonsyndromic autosomal dominant congenital cataract

Definite:

- (1) fulfills either of Major features and
- (2) Genetic criteria and/or pathological criteria

Possbile:

Col4a1is possible when the individual has either of Major features

Unlikely:

No phenotype

Brain small vessel disease 2 (COL4A2)

Major features

- (1)Porencephaly
- (2)Brain small-vessel disease with intracranial hemorrhage
- (3)cerebellar and optic atrophy, cataracts, intracranial aneurysms, nephropathy, and myopathy.

Definite:

(1) fulfills either of Major features and

(2) Genetic criteria

Possbile:

Col4a1is possible when the individual has either of Major features

Unlikely:

No phenotype

MOYAMOYA (RNF213)

Diagnostic Criteria

- (1) Cerebral angiography is considered essential for the diagnosis, and must show at least the follow-ing findings:
- (i) Stenosis or occlusion of the terminal portion of the intracranial internal carotid artery or proximal portions of the anterior and/or the middle cerebral artery.
- ii) Abnormal vascular networks in the vicinity of the occlusive or stenotic lesions in the arterial phase.
- (iii) Bilaterality of findings (i) and (ii).
- (2)However, when magnetic resonance imaging(MRI) and magnetic resonance angiographic (MRA) findings meet all of the following criteria, cerebral angiography can be omitted. See the "Guidelines for Diagnostic Imaging by MRI and MRA"
- (i) MRA shows stenosis or occlusion of the terminal portion of the intracranial internal carotid artery or proximal portions of the anterior and/or the middle cerebral artery.
- (ii) MRA shows abnormal vascular networks in the basal ganglia.

Note: When 2 or more visible flow voids are present in the basal ganglia on MRI, at least unilaterally, they can be deemed as representing an abnormal vascular network.

- (iii) Bilaterality of findings (i) and (ii).
- (3) Moyamoya disease is an illness of unknown etiology. The differential diagnosis of this disease includes similar cerebrovascular lesions associated with the following underlying diseases, which should, therefore, be excluded: (i) atherosclerosis, (ii) autoimmune disease, (iii) meningitis, (iv) brain tumors, (v) Down's syndrome, (vi) von Recklinghausen's disease, (vii) head injury, (viii) cerebrovascular lesions after head irradiation, and (ix) others.

Pathological findings that can be used as references for the diagnosis

Thickening of the arterial intima, mainly in the terminal portion of the internal carotid arteries, and narrowing or blockage of the lumen caused by this change, usually bilateral. Occasionally, lipid deposits are also present in the thickened intima.

Arteries such as the anterior, middle, and posterior cerebral arteries forming the circle of Willis occasionally show varying degrees of stenosis or occlusion associated with fibrocellular thickening of the intima, and thinning of the media.

Numerous small vascular channels (perforating and anastomotic barnches)can be seen around the circle of Willis.

Pia mater may also show reticular conglomerates of small vessels.

Diagnostic assessment:

Definite:

All criteria listed in(1)or(2) and in (3)should be met.

Possbile

All criteria are fulfilled except item(1)(iii) and/or item(2)(iii) among the criteria of (1)or(2) and (3). Unlikely:

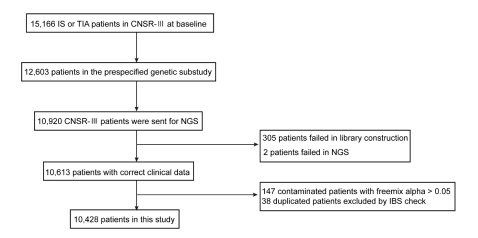
No phenotype

eReference

- 1. Li Q, Wang K. InterVar: Clinical Interpretation of Genetic Variants by the 2015 ACMG-AMP Guidelines. *Am J Hum Genet* 2017;100(2):267-80. doi: 10.1016/j.ajhg.2017.01.004 [published Online First: 2017/01/31]
- 2. Richards S, Aziz N, Bale S, et al. Standards and guidelines for the interpretation of sequence variants: a joint consensus recommendation of the American College of Medical Genetics and Genomics and the Association for Molecular Pathology. *Genet Med* 2015;17(5):405-24. doi: 10.1038/gim.2015.30 [published Online First: 2015/03/06]
- 3. Japp AG, Gulati A, Cook SA, et al. The Diagnosis and Evaluation of Dilated Cardiomyopathy. *J Am Coll Cardiol* 2016;67(25):2996-3010. doi: 10.1016/j.jacc.2016.03.590 [published Online First: 2016/06/25]
- 4. Gersh BJ, Maron BJ, Bonow RO, et al. 2011 ACCF/AHA guideline for the diagnosis and treatment of hypertrophic cardiomyopathy: executive summary: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol* 2011;58(25):2703-38. doi: 10.1016/j.jacc.2011.10.825 [published Online First: 2011/11/15]
- 5. Authors/Task Force m, Elliott PM, Anastasakis A, et al. 2014 ESC Guidelines on diagnosis and management of hypertrophic cardiomyopathy: the Task Force for the Diagnosis and Management of Hypertrophic Cardiomyopathy of the European Society of Cardiology (ESC). *Eur Heart J* 2014;35(39):2733-79. doi: 10.1093/eurheartj/ehu284 [published Online First: 2014/09/01]
- 6. Maron BJ, Towbin JA, Thiene G, et al. Contemporary definitions and classification of the cardiomyopathies: an American Heart Association Scientific Statement from the Council on Clinical Cardiology, Heart Failure and Transplantation Committee; Quality of Care and Outcomes Research and Functional Genomics and Translational Biology Interdisciplinary Working Groups; and Council on Epidemiology and Prevention. *Circulation* 2006;113(14):1807-16. doi: 10.1161/CIRCULATIONAHA.106.174287 [published Online First: 2006/03/29]
- 7. Ando Y, Coelho T, Berk JL, et al. Guideline of transthyretin-related hereditary amyloidosis for clinicians. *Orphanet J Rare Dis* 2013;8:31. doi: 10.1186/1750-1172-8-31 [published Online First: 2013/02/22]
- 8. Loeys BL, Dietz HC, Braverman AC, et al. The revised Ghent nosology for the Marfan syndrome. *J Med Genet* 2010;47(7):476-85. doi: 10.1136/jmg.2009.072785 [published Online First: 2010/07/02]
- 9. Neurofibromatosis. Conference statement. National Institutes of Health Consensus Development Conference. *Arch Neurol* 1988;45(5):575-8. [published Online First: 1988/05/01]
- 10. Tefferi A, Thiele J, Orazi A, et al. Proposals and rationale for revision of the World Health Organization diagnostic criteria for polycythemia vera, essential thrombocythemia, and primary myelofibrosis: recommendations from an ad hoc international expert panel. *Blood* 2007;110(4):1092-7. doi: 10.1182/blood-2007-04-083501 [published Online First: 2007/05/10]
- 11. Bergmann C, Guay-Woodford LM, Harris PC, et al. Polycystic kidney disease. *Nat Rev Dis Primers* 2018;4(1):50. doi: 10.1038/s41572-018-0047-y [published Online First: 2018/12/14]
- 12. Malfait F, Francomano C, Byers P, et al. The 2017 international classification of the Ehlers-Danlos syndromes. *Am J Med Genet C Semin Med Genet* 2017;175(1):8-26. doi: 10.1002/ajmg.c.31552

[published Online First: 2017/03/18]

13. Mizuta I, Watanabe-Hosomi A, Koizumi T, et al. New diagnostic criteria for cerebral autosomal dominant arteriopathy with subcortical infarcts and leukocencephalopathy in Japan. *J Neurol Sci* 2017;381:62-67. doi: 10.1016/j.jns.2017.08.009 [published Online First: 2017/10/11]



eFigure 1. Flow chart illustrating the patients screening process

eTable 1 The classification of the stroke etiology for each gene

Gene	Phenotype	Etiology of stroke
RBM20	Hereditary cardiomyopathies	Embolic stroke
ACTC1	Hereditary cardiomyopathies	Embolic stroke
LMNA	Hereditary cardiomyopathies	Embolic stroke
MYBPC3	Hereditary cardiomyopathies	Embolic stroke
ACTN2	Hereditary cardiomyopathies	Embolic stroke
PRKAG2	Hereditary cardiomyopathies	Embolic stroke
BAG3	Hereditary cardiomyopathies	Embolic stroke
DSG2	Hereditary cardiomyopathies	Embolic stroke
KCNA5	Hereditary cardiac dysrhythm	Embolic stroke
KCNJ5	Hereditary cardiac dysrhythm	Embolic stroke
SCN4B	Hereditary cardiac dysrhythm	Embolic stroke
KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke
GJA5	Hereditary cardiac dysrhythm	Embolic stroke
SCN5A	Hereditary cardiac dysrhythm	Embolic stroke
KCNJ2	Hereditary cardiac dysrhythm	Embolic stroke
KCNH2	Hereditary cardiac dysrhythm	Embolic stroke
KCNE2	Hereditary cardiac dysrhythm	Embolic stroke
SCN3B	Hereditary cardiac dysrhythm	Embolic stroke
NPPA	Hereditary cardiac dysrhythm	Embolic stroke
SCN2B	Hereditary cardiac dysrhythm	Embolic stroke
SCN1B	Hereditary cardiac dysrhythm	Embolic stroke
PRKAR1A	Carney complex, type 1	Embolic stroke
KRAS	Noonan syndrome 3	Embolic stroke
TLL1	Atrial septal defect	Embolic stroke
GJA1	Atrioventricular septal defect 3	Embolic stroke
GATA4	Tetralogy of Fallot	Embolic stroke
NKX2-5	Tetralogy of Fallot	Embolic stroke
ZFPM2	Tetralogy of Fallot	Embolic stroke
GDF1	Congenital heart defects,	Embolic stroke
	multiple types, 6	
TTN	Hereditary cardiomyopathies	Embolic stroke
MFAP5	Aortic aneurysm, familial	Large artery disease
	thoracic 9	
MYLK	Aortic aneurysm, familial	Large artery disease
	thoracic 7	
PRKG1	Aortic aneurysm, familial	Large artery disease
	thoracic 8	
MYH11	Aortic aneurysm, familial	Large artery disease
	thoracic 4	

thoracic 6 ELN Marfan syndrome ELN Supravalvar aortic stenosis COL5A1 Ehlers-Danlos syndrome, classic type COL1A1 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 1 COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 1 COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 1 COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 2 CETP Hyperalphalipoproteinemia RNF213 Moyamoya disease Large artery disease Prothrombotic state Prothrombotic state Prothrombotic state Prothrombotic state Prothrombotic state Thrombophilia due to protein S deficiency Prothrombotic state Arterior and the state state state Prothrombotic state Thrombophilia due to protein S deficiency Prothrombotic state STIMI Stormorken syndrome Prothrombotic state STIMI Stormorken syndrome Prothromboti	Gene	Phenotype	Etiology of stroke
FBNI Marfan syndrome ELN Supravalvar aortic stenosis COL5A1 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 1 COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 1 COL1A3 (Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 2 CETP Hyperalphalipoproteinemia RNF213 Moyamoya disease LDLR Hypercholesterolemia, familial, 1 APOA5 Hyperchylomicronemia, late- onset Thrombophilia due to thrombin defect JAK2 Thrombocythemia Prothrombotic state defect JAK2 Thrombocythemia Prothrombotic state Type ETV6 Thrombocytopenia 5 SERPINC1 Thrombophilia due to heparin cofactor II deficiency SERPINC1 Thrombophilia due to protein S deficiency, autos Thrombophilia due to protein S deficiency, autos STIM1 Stormorken syndrome NOTCH3 CADASIL Brain small vessel disease COL4A2 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral amyloidosis, inerditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP Cerebral amyloid angiopathy, PRNP Cerebral amyloid angiopathy, PRNP Cerebral amyloid angiopathy, PRNP Cetter Torombotics I NET Descriptions Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Prothrombotic state Small vessel disease Thrombophilia due to protein S defectioney Prothrombotic state Small vessel disease Small vessel disease Thrombophilia due to protein S defectioney Small vessel disease Small vessel disease Other disease (small and large artery disease)	ACTA2		Large artery disease
ELN Supravalvar aortic stenosis COL5A1 Ehlers-Danlos syndrome, classic type COL1A1 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome! COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome! COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome ! COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 2 CETP Hyperalphalipoproteinemia RNF213 Moyamoya disease Large artery disease Large artery disease LDLR Hypercholesterolemia, familial, 1 Large artery disease Large artery disease LDLR Hyperchylomicronemia, lateonset Intrombophilia due to thrombin defect Thrombophilia due to thrombin defect JAK2 Thrombophilia due to thrombin defect Vow Willebrand disease, type 1 GP1BA von Willebrand disease, platelettype TVO Thrombophilia due to heparin cofactor II deficiency SERPIND1 Thrombophilia due to heparin cofactor II deficiency SERPINC1 Thrombophilia due to protein S deficiency PROS1 Thrombophilia due to protein S deficiency PROS2 Thrombophilia due to protein S deficiency PROS4 Thrombophilia due to protein S deficiency PROS4 Thrombophilia due to protein S deficiency PROS4 Thrombophilia due to protein S deficiency PROS5 Thrombophilia due to protein S deficiency PROS6 Thrombophilia due to protein S deficiency PROS6 Thrombophilia due to protein S deficiency PROS6 Thrombophilia due to protein S deficiency PROS7 Thrombophilia due Service Small vessel disease COL4A2 Brain small vessel disease Small vessel disease COL4A2 Brain small vessel disease Small vessel disease COL4A2 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Firnish type TTR Amylo	FDN1		Lorge ortery disease
COL5A1 Ehlers-Danlos syndrome, classic type COL1A1 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 1 COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 1 COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 2 CETP Hyperalphalipoproteinemia Large artery disease RNF213 Moyamoya disease LDLR Hypercholesterolemia, familial, Large artery disease LDLR Hypercholesterolemia, familial, Large artery disease LDLR Hypercholesterolemia, familial, Large artery disease LDLR Hyperchylomicronemia, lateonset Thrombophilia due to thrombin defect Thrombophilia due to thrombin defect von Willebrand disease, type 1 Prothrombotic state VWF von Willebrand disease, platelettype ETV6 Thrombocytopenia 5 Prothrombotic state Prothrombotic state Thrombophilia due to heparin cofactor II deficiency SERPIND1 Thrombophilia due to heparin cofactor II deficiency PROS1 Thrombophilia due to protein S deficiency autos FTM Thrombophilia due to protein S deficiency autos STIM1 Stormorken syndrome Prothrombotic state Small vessel disease HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S Small vessel disease TTR Amyloidosis, hiroish type Small vessel disease TTR Amyloidosis, Firnish type Small vessel disease TTR Amyloidosis, bereditary, Small vessel disease TTR Amyloidosis, Firnish type Small vessel disease TTR Amyloidosis, fereditary, Small vessel			
COL1A1 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 1 COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 2 COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 2 CETP Hyperalphalipoproteinemia RNF213 Moyamoya disease LDLR Hypercholesterolemia, familial, 1 APOA5 Hypercholesterolemia, familial, 1 APOA5 Hypercholesterolemia, familial, 1 APOA5 Hyperchylomicronemia, lateonset F2 Thrombophilia due to thrombin defect JAK2 Thromboeythemia 3 Prothrombotic state F3 F2 Thromboeythemia 3 Prothrombotic state F4 Von Willebrand disease, type 1 Prothrombotic state F5 Thrombophilia due to heparin cofactor II deficiency Thrombophilia due to heparin cofactor II deficiency F5 Thrombophilia due to protein C deficiency, autos F6 Thrombophilia due to protein C deficiency, autos F7 Thrombophilia due to protein C deficiency autos F7 Thr			
COL1A1 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 1 COL1A2 Combined osteogenesis imperfecta and Ehlers-Danlos syndrome 2 CETP Hyperalphalipoproteinemia Large artery disease RNF213 Moyamoya disease LDLR Hypercholesterolemia, familial, 1 APOA5 Hypercholesterolemia, familial, 1 APOA5 Hyperchylomicronemia, lateonset onset Thrombophilia due to thrombin defeet JAK2 Thrombocythemia 3 Prothrombotic state GEP1BA von Willebrand disease, type 1 Prothrombotic state GP1BA von Willebrand disease, platelettype ETV6 Thrombophilia due to heparin cofactor II deficiency PROS1 Thrombophilia due to protein S deficiency PROS1 Thrombophilia due to protein S deficiency PROC Thrombophilia due to protein C deficiency autos STIMI Stormorken syndrome NOTCH3 CADASIL Small vessel disease COL4A2 Brain small vessel disease 2 COL4A1 Brain small vessel disease 2 COL4A2 Brain small vessel disease 2 COL4A1 Brain small vessel disease 2 COL4A1 Brain small vessel disease 2 COL4A2 Brain small vessel disease 2 COL4A1 Brain small vessel disease 2 COL4A2 Brain small vessel disease 2 COL4A1 Brain small vessel disease 2 COL4A2 Brain small vessel disease 2 COL4A3 Brain small vessel disease 2 COL4A4 Brain small vessel disease 2 COL4A5 Brain small vessel disease 3 COL4A6 Brain small vessel disease 4 COL4A7 Brain small vessel disease 5 COL4A8 Brain small vessel disease 6 COL4A9 Amyloidosis, Finnish type TTR Amyloidosis, Finnish type TTR Amyloidosis, Finnish type TTR Amyloidosis, Finnish type Cerebral alunyloid angiopathy, PRNP-related PRNP Cerebral amyloid angiopathy, PRNP-related Other disease (small and large artery disease)	COLSINI		Large artery disease
imperfecta and Ehlers-Danlos syndrome 2 CETP Hyperalphalipoproteinemia RNF213 Moyamoya disease LDLR Hypercholesterolemia, familial, 1 APOA5 Hyperchylomicronemia, late- onset F2 Thrombophilia due to thrombin defect JAK2 Thrombocythemia 3 Prothrombotic state VWF von Willebrand disease, type 1 GPIBA von Willebrand disease, platelet- type ETV6 Thrombophilia due to heparin cofactor II deficiency SERPINC1 Thrombophilia due to protein C deficiency, autos Thrombophilia due to protein C deficiency, autos STIM1 Stormorken syndrome NOTCH3 CADASIL HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 COL4A2 Brain small vessel disease vith or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type TTR Amyloidosis, Finnish type PROP Cerebral amyloid angiopathy, PRNP Cerebral amyloid angiopathy	COL1A1	Combined osteogenesis imperfecta and Ehlers-Danlos	Large artery disease
RNF213 Moyamoya disease LDLR Hypercholesterolemia, familial, 1 APOA5 Hyperchylomicronemia, late- onset F2 Thrombophilia due to thrombin defect JAK2 Thrombocythemia 3 Prothrombotic state WWF von Willebrand disease, type 1 GP1BA von Willebrand disease, platelet- type ETV6 Thrombocytopenia 5 Prothrombotic state SERPIND1 Thrombophilia due to heparin cofactor II defficiency SERPIND1 Thrombophilia due to protein C deficiency PROS1 Thrombophilia due to protein S deficiency BTM1 Stormorken syndrome NOTCH3 CADASIL HTRA1 - autosomal dominant disease COL4A2 Brain small vessel disease 2 COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type TSEN I Vescel disease GLA Fabry Disease NF1 Neurofibromatosis 1 Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease Large artery disease)	COL1A2	imperfecta and Ehlers-Danlos	Large artery disease
LDLR Hypercholesterolemia, familial, I Large artery disease APOA5 Hyperchylomicronemia, late- onset F2 Thrombophilia due to thrombin defect JAK2 Thrombocythemia 3 Prothrombotic state VWF von Willebrand disease, type 1 Prothrombotic state GP1BA von Willebrand disease, platelet- type ETV6 Thrombophilia due to heparin cofactor II deficiency SERPIND1 Thrombophilia due to heparin cofactor II deficiency PROS1 Thrombophilia due to protein S deficiency PROC Thrombophilia due to protein S deficiency APOCC Thrombophilia due to protein S APOCC Thromb	CETP		Large artery disease
APOA5 Hyperchylomicronemia, late- onset F2 Thrombophilia due to thrombin defect JAK2 Thrombocythemia 3 VWF von Willebrand disease, type 1 GP1BA Von Willebrand disease, type 1 GP1BA Von Willebrand disease, platelet- type ETV6 Thrombocytopenia 5 SERPIND1 Thrombophilia due to heparin cofactor II deficiency SERPINC1 Thrombophilia due to protein S deficiency PROS1 Thrombophilia due to protein S deficiency, autos STIM1 Stormorken syndrome NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, hereditary, transthyretin-related PSEN1 Alzhemer's disease CH4 Fabry Disease (small and large artery disease) Large artery disease) Prothrombotic state Prothrom	RNF213		Large artery disease
F2 Thrombophilia due to thrombin defect JAK2 Thrombocythemia 3 Prothrombotic state WWF von Willebrand disease, type 1 Prothrombotic state GP1BA von Willebrand disease, platelettype ETV6 Thrombocytopenia 5 Prothrombotic state SERPIND1 Thrombophilia due to heparin cofactor II deficiency SERPINC1 Thrombophilia due to antithrombin III deficiency PROS1 Thrombophilia due to protein S deficiency PROC Thrombophilia due to protein C deficiency, autos STIM1 Stormorken syndrome Prothrombotic state NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease 2 Small vessel disease TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease Other disease Small vessel disease Other disease Other disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	LDLR	1	Large artery disease
JAK2 Thrombocythemia 3 Prothrombotic state VWF von Willebrand disease, type 1 Prothrombotic state GP1BA von Willebrand disease, platelet- type ETV6 Thrombocytopenia 5 Prothrombotic state SERPIND1 Thrombophilia due to heparin cofactor II deficiency SERPINC1 Thrombophilia due to antithrombin III deficiency PROS1 Thrombophilia due to protein S deficiency PROC Thrombophilia due to protein S deficiency, autos STIM1 Stormorken syndrome Prothrombotic state NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease NF1 Neurofibromatosis 1 Other disease (small and large artery disease) NF1 Neurofibromatosis 1 Prothrombotic state Prothrombo	APOA5	onset	Large artery disease
VWF von Willebrand disease, type 1 GP1BA von Willebrand disease, platelet- type ETV6 Thrombocytopenia 5 SERPIND1 Thrombophilia due to heparin cofactor II deficiency SERPINC1 Thrombophilia due to antithrombin III deficiency PROS1 Thrombophilia due to protein S deficiency PROC Thrombophilia due to protein C deficiency, autos STIM1 Stormorken syndrome NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 COL4A1 Brain small vessel disease 2 STIM1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease NF1 Neurofibromatosis 1 Neurofibromatosis 1 Neurofibromatosis 1 Neurofibromatosis 1 Neurofibromatosis 1 Neurofibromatosis 1 Prothrombotic state Prothromboti	F2	•	Prothrombotic state
GP1BA von Willebrand disease, platelet- type ETV6 Thrombocytopenia 5 Prothrombotic state SERPIND1 Thrombophilia due to heparin cofactor II deficiency SERPINC1 Thrombophilia due to antithrombin III deficiency PROS1 Thrombophilia due to protein S deficiency PROC Thrombophilia due to protein C deficiency, autos STIM1 Stormorken syndrome Prothrombotic state NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	JAK2		
type ETV6 Thrombocytopenia 5 Prothrombotic state SERPIND1 Thrombophilia due to heparin cofactor II deficiency SERPINC1 Thrombophilia due to antithrombin III deficiency PROS1 Thrombophilia due to protein S deficiency PROC Thrombophilia due to protein C deficiency, autos STIM1 Stormorken syndrome Prothrombotic state NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type Small vessel disease TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease NF1 Neurofibromatosis 1 Thrombophilia due to protein C Prothrombotic state Pro	VWF	von Willebrand disease, type 1	Prothrombotic state
SERPIND1 Thrombophilia due to heparin cofactor II deficiency SERPINC1 Thrombophilia due to antithrombin III deficiency PROS1 Thrombophilia due to protein S deficiency PROC Thrombophilia due to protein C deficiency, autos STIM1 Stormorken syndrome Prothrombotic state NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	GP1BA	-	Prothrombotic state
cofactor II deficiency SERPINC1 Thrombophilia due to antithrombin III deficiency PROS1 Thrombophilia due to protein S deficiency PROC Thrombophilia due to protein C deficiency, autos STIM1 Stormorken syndrome Prothrombotic state NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type Small vessel disease TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease Small vessel disease	ETV6	Thrombocytopenia 5	Prothrombotic state
SERPINC1 Thrombophilia due to antithrombin III deficiency PROS1 Thrombophilia due to protein S deficiency PROC Thrombophilia due to protein C deficiency, autos STIM1 Stormorken syndrome Prothrombotic state NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type Small vessel disease TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease GLA Fabry Disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	SERPIND1		Prothrombotic state
PROS1 Thrombophilia due to protein S deficiency PROC Thrombophilia due to protein C deficiency, autos STIM1 Stormorken syndrome Prothrombotic state NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type Small vessel disease TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease GLA Fabry Disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	SERPINC1	Thrombophilia due to	Prothrombotic state
PROC Thrombophilia due to protein C deficiency, autos STIM1 Stormorken syndrome Prothrombotic state NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type Small vessel disease TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease GLA Fabry Disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	PROS1	Thrombophilia due to protein S	Prothrombotic state
STIM1 Stormorken syndrome Prothrombotic state NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type Small vessel disease TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease GLA Fabry Disease Small vessel disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	PROC	Thrombophilia due to protein C	Prothrombotic state
NOTCH3 CADASIL Small vessel disease HTRA1 HTRA1-autosomal dominant disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type Small vessel disease TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease GLA Fabry Disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	STIM1	Stormorken syndrome	Prothrombotic state
disease COL4A2 Brain small vessel disease 2 Small vessel disease COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type Small vessel disease TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease GLA Fabry Disease Small vessel disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	NOTCH3		Small vessel disease
COL4A1 Brain small vessel disease with or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type Small vessel disease TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease GLA Fabry Disease Small vessel disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	HTRA1		Small vessel disease
or without ocular anomalies/PADMAL TREX1 Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type Small vessel disease TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease GLA Fabry Disease GLA Fabry Disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	COL4A2	Brain small vessel disease 2	Small vessel disease
cerebral leukoencephalopathy and systemic manifestations/RVCL-S GSN Amyloidosis, Finnish type Small vessel disease TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease GLA Fabry Disease GLA Fabry Disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	COL4A1	or without ocular	Small vessel disease
TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease GLA Fabry Disease Other disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	TREX1	cerebral leukoencephalopathy and systemic manifestations/RVCL-S	Small vessel disease
TTR Amyloidosis, hereditary, transthyretin-related PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease GLA Fabry Disease Other disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	GSN		Small vessel disease
PRNP Cerebral amyloid angiopathy, PRNP-related PSEN1 Alzheimer's disease Small vessel disease GLA Fabry Disease Other disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	TTR		Small vessel disease
PSEN1 Alzheimer's disease Small vessel disease GLA Fabry Disease Other disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	PRNP	Cerebral amyloid angiopathy,	Small vessel disease
GLA Fabry Disease Other disease (small and large artery disease) NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	PSEN1		Small vessel disease
NF1 Neurofibromatosis 1 Other disease (small and large artery disease)	GLA		Other disease
	NF1	Neurofibromatosis 1	Other disease
	ABCC6	Pseudoxanthoma elasticum	Other disease

Gene	Phenotype	Etiology of stroke
		(small and large artery disease)
PKD1	Polycystic kidney disease 1	Other disease
		(Cerebrovascular
		malformations)
PKD2	Polycystic kidney disease 2	Other disease
		(Cerebrovascular
		malformations)
KRIT1	Cerebral cavernous	Other disease
	malformations-1	(Cerebrovascular
		malformations)
ACVRL1	Telangiectasia, hereditary	Other disease
	hemorrhagic, type 2	(Cerebrovascular
		malformations)
ENG	Telangiectasia, hereditary	Other disease
	hemorrhagic, type 1	(Cerebrovascular
		malformations)
FLCN	Birt-Hogg-Dube syndrome	Other disease
		(Unknown mechanism)
DYRK1B	Abdominal obesity-metabolic	Other disease
	syndrome 3	(Unknown mechanism)
PDE4D	Acrodysostosis 2, with or	Other disease
	without hormone resistance	(Unknown mechanism)
APOA1	Amyloidosis, 3 or more types	Other disease
		(Unknown mechanism)
VHL	Pheochromocytoma	Other disease
		(Unknown mechanism)
RET	Medullary thyroid carcinoma or	Other disease
	Pheochromocytoma	(Unknown mechanism)
BMPR2	Pulmonary hypertension,	Other disease
	primary	(Unknown mechanism)
CBL	Noonan syndrome-like disorder	Other disease
	with or without juvenile	(Unknown mechanism)
	myelomonocytic leukemia	· · · · · · · · · · · · · · · · · · ·
KIF1B	Pheochromocytoma	Other disease
		(Unknown mechanism)
TGIF1	Holoprosencephaly 4	Other disease
	_ , ,	(Unknown mechanism)

eTable 2 List of the 181 genes associated with Mendelian-stroke in custom-designed panel

Gene	Location	OMIM ID	Inheri- tance	Associated Mendelian Disorder
ABCA1	9q31.1	600046	AR	Tangier disease
ABCC6	16p13.11	603234	AR	Arterial calcification, generalized, of infancy, 2; Pseudoxanthoma elasticum
ACTA2	10q23.31	102620	AD	Moyamoya disease 5; Aortic aneurysm, familial thoracic 6; Multisystemic smooth muscle dysfunction syndrome
ACTC1	15q14	102540	AD	Atrial septal defect 5; Cardiomyopathy, dilated, 1R; Cardiomyopathy, hypertrophic, 11; Left ventricular noncompaction 4
ACTN2	1q43	102573	AD	Cardiomyopathy, dilated, 1AA, with or without LVNC; Cardiomyopathy, hypertrophic, 23, with or without LVNC; Myopathy, congenital with structured cores and Z-line abnormalities; Myopathy, distal, 6, adult onset
ACVRL1	12q13.13	601284	AD	Telangiectasia, hereditary hemorrhagic, type 2
ANK2	4q25-q26	106410	AD	Cardiac arrhythmia, ankyrin-B-related; Long QT syndrome 4
APOA1	11q23.3	107680	AD	Amyloidosis, 3 or more types
APOA5	11q23.3	606368	AD	Hyperchylomicronemia, late-onset
APP	21q21.3	104760	AD	Alzheimer disease 1, familial; Cerebral amyloid angiopathy, Dutch, Italian, Iowa, Flemish, Arctic variants
ATP7A	Xq21.1	300011	XLR	Menkes disease; Occipital horn syndrome; Spinal muscular atrophy, distal, X-linked 3
B4GALT1	9p21.1	137060	AR	Congenital disorder of glycosylation, type IId
BAG3	10q26.11	603883	AD	Cardiomyopathy, dilated, 1HH
BMPR2	2q33.1- q33.2	600799	AD	Pulmonary hypertension, familial primary, 1, with or without HHT; Pulmonary hypertension, primary, fenfluramine or dexfenfluramineassociated; Pulmonary venoocclusive disease 1
BRCC3	Xq28	300617	XLR	Moyamoya 4 X-link
CBL	11q23.3	165360	AD	juvenile myelomonocytic leukemia and Noonan syndrome-like disorder
CBS	21q22.3	613381	AR	Homocystinuria, B6-responsive and nonresponsive types; Thrombosis, hyperhomocysteinemic
CCER2	19q13.2	617634	AD	Moyamoya disease, susceptibility to
CCM2	7p13	607929	AD	Cerebral cavernous malformations 2
CDKN1C	11p15.4	600856	AD	Beckwith-Wiedemann syndrome; IMAGE syndrome
CECR1/ ADA2	22q11.1	615688	AR	Vasculitis, autoinflammation, immunodeficiency, and hematologic defects syndrome
CEP19	3q29	615586	AR	Morbid Obesity And Spermatogenic Failure
CETP	16q13	118470	AD	Hyperalphalipoproteinemia; High density lipoprotein cholesterol level QTL 10
COL1A1	17q21.33	120150	AD	Combined osteogenesis imperfecta and Ehlers- Danlos syndrome 1

Gene	Location	OMIM ID	Inheri- tance	Associated Mendelian Disorder
COL1A2	7q21.3	120160	AD	Combined osteogenesis imperfecta and Ehlers- Danlos syndrome 2
COL3A1	2q32.2	120180	AD, AR	Ehlers-Danlos syndrome, vascular type; Polymicrogyria with or without vascular-type EDS
COL4A1	13q34	120130	AD	Angiopathy, hereditary, with nephropathy, aneurysms, and muscle cramps; Brain small vessel disease with or without ocular anomalies; Anterior segment dysgenesis with cerebral involvement; Porencephaly 1; Retinal artery tortuosity
COL4A2	13q34	120090	AD	Brain small vessel disease 2
COL5A1	9q34.3	120215	AD	Ehlers-Danlos syndrome, classic type, 1; Fibromuscular dysplasia, multifocal
CPT2	1p32.3	600650	AR	CPT II deficiency
CST3	20p11.21	604312	AD	Cerebral amyloid angiopathy
CTC1	17p13.1	613129	AR	Cerebroretinal Microangiopathy With Calcifications And Cysts
CTSA	20q13.12	613111	AR	Galactosialidosis
CYP27A1	2q35	606530	AR	Cerebrotendinous xanthomatosis
DES	2q35	125660	AD	Cardiomyopathy, dilated, 1I;Scapuloperoneal syndrome, neurogenic, Kaeser type
DOCK8	9p24.3	611432	AR	Hyper-IgE recurrent infection syndrome, autosomal recessive
DSG2	18q12.1	125671	AD	Arrhythmogenic right ventricular dysplasia, familial, 10
DYRK1B	19q13.2	604556	AD	Abdominal obesity-metabolic syndrome 3
ELN	7q11.23	130160	AD	Cutis laxa, autosomal dominant; Supravalvar aortic stenosis
ENG	9q34.11	131195	AD	Hereditary hemorrhagic telangiectasia, type 1
ENPP1	6q23.2	173335	AD, AR	Arterial calcification, generalized, of infancy, 1;Cole disease
EPHX2	8p21.2- p21.1	132811	AD, AR	Hypercholesterolemia, familial, due to LDLR defect, modifier of
ETV6	12p13.2	600618	AD	Thrombocytopenia 5
F10	13q34	613872	AR	Factor X Deficiency
F13A1	6p25.1	134570	AR	Factor XIII Subunit A Deficiency
F13B	1q31.3	134580	AR	Factor XIII Subunit B Deficiency
F2	11p11.2	176930	AD, AR	Dysprothrombinemia; Hypoprothrombinemia; Thrombophilia due to thrombin defect;
F3	1p21.3	134390	NA	NA
F5	1q24.2	612309	AD, AR	Factor V deficiency; Thrombophilia due to activated protein C resistance; Thrombophilia, susceptibility to, due to factor V Leiden
F7	13q34	613878	AR	Factor VII Deficiency
F8	Xq28	300841	XLR	Hemophilia A
FBN1	15q21.1	134797	AD	Marfan syndrome
FGA	4q31.3	134820	AD, AR	Congenital Afibrinogenemia; Amyloidosis, familial visceral
FGB	4q31.3	134830	AR	Congenital Afibrinogenemia
FGG	4q32.1	134850	AR	Congenital Afibrinogenemia
FLCN	17p11.2	607273	AD	Birt-Hogg-Dube syndrome; Pneumothorax, primary spontaneous

Gene	Location	OMIM ID	Inheri- tance	Associated Mendelian Disorder
GATA4	8p23.1	600576	AD	Atrial Septal Defect 2; Tetralogy of Fallot
GATA6	18q11.2	601656	AD	Atrial Septal Defect 9; Tetralogy of Fallot
GATAD1	7q21.2	614518	AR	Cardiomyopathy, dilated, 2B
GDF1	19p13.11	602880	AD	Congenital heart defects, multiple types, 6
GDF2	10q11.22	605120	AD	Hereditary hemorrhagic telangiectasia, type 5
GGCX	2p11.2	137167	AR	Vitamin K-Dependent Clotting Factors, Combined Deficiency of, 1
GJA1	6q22.31	121014	AD	Atrioventricular septal defect 3
GJA5	1q21.2	121013	AD	Atrial fibrillation, familial, 11
GLA	Xq22.1	300644	XL	Fabry disease; Fabry disease, cardiac variant
GP1BA	17p13.2	606672	AD, AR	Bernard-Soulier syndrome, type A2; von Willebrand disease, platelet-type
GP6	19q13.42	605546	AR	Bleeding Disorder Platelet Type 11
GSN	9q33.2	137350	AD	Amyloidosis, Finnish type
GUCY1A3	4q32.1	139396	AR	Moyamoya 6 with achalasia
HBB	11p15.4	141900	AR	Sickle cell anemia
HTRA1	10q26.13	602194	AD, AR	Cerebral autosomal recessive arteriopathy with subcortical infarcts and leukoencephalopathy (CARASIL); Cerebral arteriopathy, autosomal dominant, with subcortical infarcts and leukoencephalopathy, type 2
ITGA2	5q11.2	192974	NA	platelet disorder
ITGA2B	17q21.31	607759	AD, AR	Bleeding Disorder Platelet Type 16;Glanzmann thrombasthenia 1
ITGB3	17q21.32	173470	AD, AR	Bleeding disorder, platelet-type, 24, autosomal dominant; Glanzmann thrombasthenia 2
ITM2B	13q14.2	603904	AD	Retinal dystrophy with inner retinal dysfunction and ganglion cell abnormalities; Dementia
ITPKC	19q13.2	606476		Kawasaki disease, associated
JAG1	20p12.2	601920	AD	Tetralogy of Fallot; Alagille syndrome 1
JAK2	9p24.1	147796	AD	Thrombocythemia 3
KCNA5	12p13.32	176267	AD	Familial Atrial Fibrillation 7
KCNE1	21q22.12	176261	AD, AR	Long QT syndrome 5; Jervell and Lange-Nielsen syndrome 2
KCNE2	21q22.11	603796	AD	Familial Atrial Fibrillation 4; Long QT syndrome 6
KCNH2	7q36.1	152427	AD	Long QT syndrome 2; Short QT syndrome 1
KCNJ2	17q24.3	600681	AD	Familial Atrial Fibrillation 9; Short QT syndrome 3; Andersen syndrome
KCNJ5	11q24.3	600734	AD	Long QT syndrome 13; Hyperaldosteronism, familial, type III
KCNQ1	11p15.5- p15.4	607542	AD	Atrial fibrillation, familial 3; Long QT syndrome 1; Short QT syndrome 2
KIF1B	1p36.22	605995	AD	Pheochromocytoma; Charcot-Marie-Tooth disease, type 2A1
KRAS	12p12.1	190070	AD	Arteriovenous malformation of the brain, somatic; Cardiofaciocutaneous syndrome 2; Noonan syndrome 3; RAS-associated autoimmune leukoproliferative disorder
KRIT1	7q21.2	604214	AD	Cerebral cavernous malformations 1
LAMP2	Xq24	309060	XLD	Danon disease
LDLR	19p13.2	606945	AD, AR	Hypercholesterolemia, familial

Gene	Location	OMIM ID	Inheri- tance	Associated Mendelian Disorder
LIPC	15q21.3	151670	AR	High density lipoprotein cholesterol level QTL 12; Hepatic lipase deficiency
LMBRD1	6q13	612625	AR	Methylmalonic aciduria and homocystinuria, cblF type
LMNA	1q22	150330	AD, AR	Charcot-Marie-Tooth disease, type 2B1; Cardiomyopathy, dilated, 1A
LPL	8p21.3	609708	AD, AR	Hepatic Lipase Deficiency; Combined hyperlipidemia, familial; High density lipoprotein cholesterol level QTL 11
MFAP5	12p13.31	601103	AD	Aortic aneurysm, familial thoracic 9
MMACHC	1p34.1	609831	AR	Methylmalonic aciduria and homocystinuria, cblC type
MPL	1p34.2	159530	AD, AR	Thrombocythemia 2; Thrombocytopenia, congenital amegakaryocytic
MTCP1	Xq28	300116	XLR	Moyamoya disease
MTHFR	1p36.22	607093	AR	Homocystinuria due to MTHFR deficiency
MTR	1q43	156570	AR	Homocystinuria-megaloblastic anemia, cblG complementation type
MTRR	5p15.31	602568	AR	Homocystinuria-megaloblastic anemia, cbl E type
MTTP	4q23	157147	AD, AR	Abdominal obesity-metabolic syndrome 1; Abetalipoproteinemia
MMUT	6p12.3	609058	AR	Methylmalonic aciduria, mut (0) type
MYBPC3	11p11.2	600958	AD	Cardiomyopathy, hypertrophic, 4; Cardiomyopathy, dilated, 1MM; Left ventricular noncompaction 10
MYH11	16p13.11	160745	AD	Aortic aneurysm, familial thoracic 4
МҮН9	22q12.3	160775	AD	Macrothrombocytopenia and granulocyte inclusions with or without nephritis or sensorineural hearing loss; Deafness, autosomal dominant 17
MYLK	3q21.1	600922	AD	Aortic aneurysm, familial thoracic 7
NF1	17q11.2	613113	AD	Neurofibromatosis, type 1
NKX2-5	5q35.1	600584	AD	Atrial Septal Defect 7, with or without AV Conduction Defects; Tetralogy of Fallot
NOS3	7q36.1	163729	AD	Alzheimer disease, type 1
NOTCH3	19p13.12	600276	AD	Cerebral arteriopathy with subcortical infarcts and leukoencephalopathy (CADASIL1)
NPPA	1p36.22	108780	AD	Familial Atrial Fibrillation 6
NUP155	5p13.2	606694	AR	Familial Atrial Fibrillation 15
P2RY12	3q25.1	600515	AR	Bleeding Disorder Platelet Type 8
PCNT	21q22.3	605925	AR	Microcephalic osteodysplastic primordial dwarfism, type II
PDCD10	3q26.1	609118	AD	Cerebral cavernous malformations 3
PDE4D	5q11.2- q12.1	600129	AD	Acrodysostosis 2, with or without hormone resistance
PGM1	1p31.3	171900	AR	Congenital disorder of glycosylation, type It
PHACTR1	6p24.1	608723	AD	Spontaneous Bilateral Cervical Internal Carotid and Vertebral Artery Dissection
PIGA	Xp22.2	311770	XLR	Multiple congenital anomalies-hypotonia- seizures syndrome 2
PITX2	4q25	601542	AD	Axenfeld-Rieger syndrome, type 1
PKD1	16p13.3	601313	AD	Polycystic kidney disease 1

		OMIM ID	Inheri- tance	Associated Mendelian Disorder					
PKD2	4q22.1	173910	AD	Polycystic kidney disease 2					
PLA2G7	6p12.3	601690	AR	Platelet-Activating Factor Acetylhydrolase Deficiency					
PLAU	10q22.2	191840	AD	Quebec Platelet Disorder					
PLOD3	7q22.1	603066	AR	Lysyl hydroxylase 3 deficiency					
PNP	14q11.2	164050	AR	Immunodeficiency due to purine nucleoside phosphorylase deficiency					
POLD1	19q13.33	174761	AD	Mandibular Hypoplasia, Deafness, Progeroid Features, And Lipodystrophy syndrome					
PRKAG2	7q36.1	602743	AD	Cardiomyopathy, hypertrophic 6;Glycogen storage disease of heart, lethal congenital; Wolff-Parkinson-White syndrome					
PRKAR1A	17q24.2	188830	AD	Carney complex, type 1					
PRKG1	10q11.2- q21.1	176894	AD	Aortic aneurysm, familial thoracic 8					
PRNP	20p13	176640	AD	Cerebral amyloid angiopathy, PRNP-related					
PROC	2q14.3	612283	AD, AR	Thrombophilia due to protein C deficiency					
PROS1	3q11.1	176880	AD, AR	Thrombophilia due to protein S deficiency					
PROZ	13q34	176895	NA	deep venous thrombolic disease, associated					
PSEN1	14q24.2	104311	AD	Cardiomyopathy, dilated, 1U;Alzheimer disease, type 3					
PSEN2	1q42.13	600759	AD	Cardiomyopathy, dilated, 1V;Alzheimer disease-4					
RASA1	5q14.3	139150	AD	Capillary malformation-arteriovenous malformation					
RBM20	10q25.2	613171	AD	Cardiomyopathy, dilated, 1DD					
RET	10q11.21	164761	AD	Pheochromocytoma					
RNF213	17q25.3	613768	AD, AR	Moyamoya disease 2, susceptibility to					
RYR1	19q13.2	180901	AD, AR	Central core disease					
RYR2	1q43	180902	AD	Arrhythmogenic Right Ventricular Dysplasia/Cardiomyopathy 2; Ventricular arrythmias due to cardiac ryanodine receptor calcium release deficiency syndrome; Ventricular tachycardia, catecholaminergic polymorphic, 1					
SCN1B	19q13.11	600235	AD	Atrial fibrillation, familial, 13					
SCN2B	11q23.3	601327	AD	Atrial fibrillation, familial, 14					
SCN3B	11q24.1	608214	AD	Atrial fibrillation, familial, 16;Brugada syndrome 7					
SCN4B	11q23.3	608256	AD	Atrial fibrillation, familial,17; Long QT syndrome 10					
SCN5A	3p22.2	600163	AD	Atrial fibrillation, familial, 10; Brugada syndrome 1; Cardiomyopathy, dilated, 1E					
SERPINC1	1q25.1	107300	AD, AR	Thrombophilia due to antithrombin III deficiency					
SERPIND1	22q11.21	142360	AD	Thrombophilia due to heparin cofactor II deficiency					
SHOC2	10q25.2	602775	AD	Noonan-like syndrome and moyamoya disease					
SLC19A2	1q24.2	603941	AR	Thiamine-Responsive Megaloblastic Anemia Syndrome					

Gene	Location	OMIM ID	Inheri- tance	Associated Mendelian Disorder
SLC2A10	20q13.12	606145	AR	Arterial tortuosity syndrome
SMAD3	15q22.33	603109	AD	Loeys-Dietz syndrome, type 3
SMAD4	18q21.2	600993	AD	Juvenile polyposis/hereditary hemorrhagic
				telangiectasia syndrome
SMARCAL1	2q35	606622	AR	Schimke immunoosseous dysplasia
STIM1	11p15.4	605921	AD, AR	;Immunodeficiency 10
TALDO1	11p15.5	602063	AR	Transaldolase deficiency
TAZ	Xq28	300394	XLR	3-Methylglutaconic aciduria, type II (Barth syndrome)
TBX1	22q11.21	602054	AD	Tetralogy of Fallot
TBX20	7p14.2	606061	AD	Atrial septal defect 4
TBXA2R	19p13.3	188070	AD	Bleeding disorder, platelet-type, 13, susceptibility to
TCAP	17q12	604488	AD	Cardiomyopathy, hypertrophic, 25
TCN1	11q12.1	189905	NA	NA
TGFB2	1q41	190220	AD	Loeys-Dietz syndrome, type 4
TGFB3	14q24.3	190230	AD	Arrhythmogenic right ventricular dysplasia 1;Loeys-Dietz syndrome 5
TGFBR1	9q22.33	190181	AD	Loeys-Dietz syndrome, type 1A; Loeys-Dietz syndrome, type 2A
TGFBR2	3p24.1	190182	AD	Loeys-Dietz syndrome 2
TGIF1	18p11.31	602630	AD	Holoprosencephaly 4
THBD	20p11.21	188040	AD	Thrombophilia Due To Thrombomodulin Defect
TLL1	4q32.3	606742	AD	Atrial septal defect 6
TMEM173	5q31.2	612374	AD	STING-associated vasculopathy, infantile-onset
TREX1	3p21.31	606609	AD	Vasculopathy, retinal, with cerebral leukodystrophy
TTN	2q31.2	188840	AD	Cardiomyopathy, dilated, 1G
TTR	18q12.1	176300	AD	Amyloidosis, hereditary, transthyretin-related
VHL	3p25.3	608537	AD	von Hippel-Lindau syndrome
VKORC1	16p11.2	608547	AD	Vitamin K-Dependent Clotting Factors, Combined Deficiency of, 2; Warfarin resistance
VWF	12p13.31	613160	AD, AR	Von Willebrand Disease
YY1AP1	1q22	607860	AR	Grange syndrome
ZFPM2	8q23.1	603693	AD	Tetralogy of Fallot

eTable 3 Diagnoses of 759 individuals with 1 P/LP variant at risk for one monogenic diseases

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinva r	ACM G	Defin ite	Possi ble	Undete rmined	Insufficien t Informati on
CNSR30 0069	F	58	ACTC1	Hereditary cardiomyopathies	Embolic stroke	NM_005159:exon3:c.T 213A:p.Y71X	NA	P			Y	
CNSR30 2746	F	71	ACTC1	Hereditary cardiomyopathies	Embolic stroke	NM_005159:exon7:c.G 1093A:p.D365N	NA	LP			Y	
CNSR30 1704	M	55	ACTN2	Hereditary cardiomyopathies	Embolic stroke	NM_001103:exon16:c. G1919A:p.R640H	LP	VUS			Y	
CNSR30 5213	M	63	ACTN2	Hereditary cardiomyopathies	Embolic stroke	NM_001103:exon16:c. G1919A:p.R640H	LP	VUS				Y
CNSR30 8476	F	64	ACTN2	Hereditary cardiomyopathies	Embolic stroke	NM_001103:exon14:c. C1618T:p.Q540X	NA	P			Y	
CNSR30 3363	M	63	BAG3	Hereditary cardiomyopathies	Embolic stroke	NM_004281:exon2:c.1 81-1G>A	NA	P			Y	
CNSR30 9438	M	52	DSG2	Hereditary cardiomyopathies	Embolic stroke	NM_001943:exon10:c. G1311A:p.W437X	NA	P			Y	
CNSR30 0880	M	51	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052.5(GATA4):c.997+103G>T	P	VUS			Y	
CNSR30 2453	M	84	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:exon6:c.G 1075A:p.E359K	P	VUS			Y	
CNSR30 2668	F	80	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:exon6:c.G 1075A:p.E359K	P	VUS			Y	
CNSR30 2724	F	53	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:c.997+10 3G>T	P	VUS			Y	
CNSR30 2956	M	53	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:exon7:c.C 1325T:p.A442V	P	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 3353	M	53	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:exon5: c.C946G:p.Q316E	P	VUS			Y	
CNSR30 3946	M	41	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:exon6: c.G1075A:p.E359K	P	VUS			Y	
CNSR30 5783	M	60	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:c.997 +103G>T	P	VUS			Y	
CNSR30 6009	M	77	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:c.997 +103G>T	P	VUS			Y	
CNSR30 9014	M	70	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:exon7: c.C1325T:p.A442V	P	VUS			Y	
CNSR30 9505	M	74	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:exon7: c.C1325T:p.A442V	P	VUS			Y	
CNSR30 9956	M	51	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:exon3: c.777delC:p.R260fs	NA	LP			Y	
CNSR30 9997	M	46	GATA4	Tetralogy of Fallot	Embolic stroke	NM_002052:c.997 +103G>T	P	VUS			Y	
CNSR30 4597	M	47	GDF1	Congenital heart defects, multiple types, 6	Embolic stroke	NM_001492:exon7: c.159delC:p.P53fs	NA	LP			Y	
CNSR30 6192	M	71	GDF1	Congenital heart defects, multiple types, 6	Embolic stroke	NM_001492:exon7: c.289dupG:p.V97fs	NA	LP			Y	
CNSR30 7760	F	72	GDF1	Congenital heart defects, multiple types, 6	Embolic stroke	NM_001492:exon7: c.C262T:p.Q88X	NA	P			Y	
CNSR30 0610	F	72	GJA1	Atrioventricular septal defect 3	Embolic stroke	NM_000165:exon2: c.A305T:p.K102M	NA	LP			Y	
CNSR30 4700	F	45	GJA1	Atrioventricular septal defect 3	Embolic stroke	NM_000165:exon2: c.A305T:p.K102M	NA	LP			Y	
CNSR30 5166	F	77	GJA1	Atrioventricular septal defect 3	Embolic stroke	NM_000165:exon2: c.G913C:p.A305P	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30	M	66	GJA1	Atrioventricular	Embolic	NM_000165:exon2:	NA	LP			Y	
5429				septal defect 3	stroke	c.A305T:p.K102M					<u> </u>	
CNSR30	M	83	GJA1	Atrioventricular	Embolic	NM_000165:exon2:	NA	LP			Y	
6575				septal defect 3	stroke	c.A305T:p.K102M					<u> </u>	
CNSR31	M	76	GJA1	Atrioventricular	Embolic	NM_000165:exon2:	NA	LP			Y	
0100				septal defect 3	stroke	c.A524G:p.Y175C						
CNSR30	M	60	GJA5	Hereditary cardiac	Embolic	NM_005266:exon2:	NA	LP			Y	
0869				dysrhythm	stroke	c.T269C:p.L90P						
CNSR30	F	66	GJA5	Hereditary cardiac	Embolic	NM_005266:exon2:	NA	LP			Y	
3120				dysrhythm	stroke	c.T269C:p.L90P						
CNSR30	M	74	GJA5	Hereditary cardiac	Embolic	NM 005266:exon2:	NA	LP		Y		
3449				dysrhythm	stroke	c.G1055T:p.R352M						
CNSR30	M	66	GJA5	Hereditary cardiac	Embolic	NM 005266:exon2:	NA	LP		Y		
5175				dysrhythm	stroke	c.T686C:p.L229P						
CNSR30	F	62	GJA5	Hereditary cardiac	Embolic	NM 005266:exon2:	NA	LP		Y		
6594				dysrhythm	stroke	c.G317C:p.R106P						
CNSR30	F	61	GJA5	Hereditary cardiac	Embolic	NM 005266:exon2:	NA	LP			Y	
9131				dysrhythm	stroke	c.G1028A:p.R343H						
CNSR30	M	64	GJA5	Hereditary cardiac	Embolic	NM 005266:exon2:	NA	LP		Y		
9756				dysrhythm	stroke	c.C316T:p.R106C						
CNSR31	F	89	GJA5	Hereditary cardiac	Embolic	NM 005266:exon2:	NA	LP		Y		
0261				dysrhythm	stroke	c.T581C:p.V194A						
CNSR30	M	64	KCNA5	Hereditary cardiac	Embolic	NM 002234:exon1:	P	VUS			Y	
0158				dysrhythm	stroke	c.C1727T:p.A576V						
CNSR30	F	43	KCNA5	Hereditary cardiac	Embolic	NM 002234:exon1:	P	VUS			Y	
0624				dysrhythm	stroke	c.C1727T:p.A576V						
CNSR30	M	47	KCNA5	Hereditary cardiac	Embolic	NM 002234:exon1:	P	VUS			Y	
1595				dysrhythm	stroke	c.C1727T:p.A576V						
CNSR30	M	45	KCNA5	Hereditary cardiac	Embolic	NM 002234:exon1:	P	VUS		1	Y	
1642	1			dysrhythm	stroke	c.C1727T:p.A576V					_	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30	M	70	KCNA5	Hereditary cardiac	Embolic	NM 002234:exon1:	P	VUS		Y		
2182				dysrhythm	stroke	c.G1828A:p.E610K						
CNSR30	M	56	KCNA5	Hereditary cardiac	Embolic	NM_002234:exon1:	P	VUS			Y	
2320				dysrhythm	stroke	c.C1727T:p.A576V						
CNSR30	F	79	KCNA5	Hereditary cardiac	Embolic	NM_002234:exon1:	P	VUS	Y			
2364				dysrhythm	stroke	c.C1727T:p.A576V						
CNSR30	M	45	KCNA5	Hereditary cardiac	Embolic	NM_002234:exon1:	P	VUS			Y	
4681				dysrhythm	stroke	c.G1828A:p.E610K						
CNSR30	M	54	KCNA5	Hereditary cardiac	Embolic	NM_002234:exon1:	P	VUS			Y	
5072				dysrhythm	stroke	c.C1727T:p.A576V						
CNSR30	M	60	KCNA5	Hereditary cardiac	Embolic	NM_002234:exon1:	P	VUS	Y			
5124				dysrhythm	stroke	c.C1727T:p.A576V						
CNSR30	M	59	KCNA5	Hereditary cardiac	Embolic	NM_002234:exon1:	P	VUS			Y	
7143				dysrhythm	stroke	c.C1727T:p.A576V						
CNSR30	M	75	KCNA5	Hereditary cardiac	Embolic	NM_002234:exon1:	P	VUS			Y	
7545				dysrhythm	stroke	c.C1727T:p.A576V						
CNSR30	M	66	KCNA5	Hereditary cardiac	Embolic	NM_002234:exon1:	P	VUS			Y	
8123				dysrhythm	stroke	c.C1727T:p.A576V						
CNSR30	M	72	KCNA5	Hereditary cardiac	Embolic	NM_002234:exon1:	P	VUS			Y	
8261				dysrhythm	stroke	c.G1828A:p.E610K						
CNSR30	M	47	KCNE2	Hereditary cardiac	Embolic	NM_172201:exon2:	LP	VUS			Y	
3689				dysrhythm	stroke	c.G205A:p.V69M						
CNSR30	M	66	KCNE2	Hereditary cardiac	Embolic	NM_172201:exon2:	LP	VUS		Y		
4562				dysrhythm	stroke	c.G205A:p.V69M						
CNSR30	F	53	KCNE2	Hereditary cardiac	Embolic	NM_172201:exon2:	LP	VUS		Y		
5932				dysrhythm	stroke	c.G205A:p.V69M						
CNSR30	M	61	KCNH2	Hereditary cardiac	Embolic	NM_000238:exon7:	LP	VUS			Y	
3137				dysrhythm	stroke	c.G1888A:p.V630I		<u> </u>				
CNSR30	M	53	KCNH2	Hereditary cardiac	Embolic	NM_000238:exon2:	NA	LP			Y	
3306				dysrhythm	stroke	c.G121A:p.V41I						

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30	F	68	KCNH2	Hereditary cardiac	Embolic	NM_000238:exon6:	LP	VUS			Y	
5582				dysrhythm	stroke	c.C1352T:p.P451L						
CNSR30 6897	M	57	KCNH2	Hereditary cardiac dysrhythm	Embolic stroke	NM_000238:exon2: c.G271T:p.E91X	NA	P			Y	
CNSR30 2460	F	70	KCNJ2	Hereditary cardiac dysrhythm	Embolic stroke	NM_000891:exon2: c.A971G:p.H324R	NA	LP			Y	
CNSR30 5866	M	52	KCNJ2	Hereditary cardiac dysrhythm	Embolic stroke	NM_000891:exon2: c.A593G:p.N198S	NA	LP			Y	
CNSR30 0301	M	44	KCNJ5	Hereditary cardiac dysrhythm	Embolic stroke	NM_000890:exon2: c.G862A:p.E288K	NA	LP			Y	
CNSR30 6658	M	58	KCNJ5	Hereditary cardiac dysrhythm	Embolic stroke	NM_000890:exon2: c.G532A:p.V178I	NA	LP			Y	
CNSR30 7982	M	33	KCNJ5	Hereditary cardiac dysrhythm	Embolic stroke	NM_000890:exon3: c.G1039T:p.D347Y	NA	LP			Y	
CNSR30 8486	F	81	KCNJ5	Hereditary cardiac dysrhythm	Embolic stroke	NM_000890:exon2: c.G632A:p.R211Q	NA	LP			Y	
CNSR30 0462	M	71	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon5: c.C758G:p.S253C	P	VUS	Y			
CNSR30 0983	F	71	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon3: c.C589T:p.P197S	LP	VUS			Y	
CNSR30 1428	M	42	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon6: c.G911A:p.W304X	NA	P		Y		
CNSR30 3201	F	48	KCNQ1	Hereditary cardiac	Embolic stroke	NM_000218:exon7: c.C965T:p.T322M	P	VUS			Y	
CNSR30 3313	M	58	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon1 1:c.1489dupC:p.L4 96fs	NA	LP			Y	
CNSR30 3540	F	54	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon1 5:c.C1780T:p.R594 X	Р	LP		Y		

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 3754	M	68	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon1 3:c.G1664A:p.R555 H	P/LP	VUS		Y		
CNSR30 4030	F	53	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon7: c.C961T:p.Q321X	P	P			Y	
CNSR30 4199	M	71	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon1 3:c.C1630T:p.Q544 X	NA	P			Y	
CNSR30 4883	M	83	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon3: c.C520T:p.R174C	P/LP	VUS			Y	
CNSR30 5135	F	59	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon1 6:c.1887dupC:p.G6 29fs	P/LP	VUS			Y	
CNSR30 5142	M	58	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon9: c.1148dupC:p.A383 fs	NA	LP			Y	
CNSR30 5149	F	78	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon1 3:c.C1639T:p.Q547 X	NA	P			Y	
CNSR30 6632	M	58	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon2: c.G436T:p.E146X	NA	P			Y	
CNSR30 6754	F	60	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon4: c.C674T:p.S225L	P/LP	VUS	Y			
CNSR30 8491	M	71	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon1 1:c.1445delC:p.T48 2fs	P	LP	Y			
CNSR30 8806	F	61	KCNQ1	Hereditary cardiac dysrhythm	Embolic stroke	NM_181798:exon1: c.5dupA:p.D2fs	NA	LP			Y	
CNSR30 0909	M	51	KRAS	Noonan syndrome 3	Embolic stroke	NM_033360:exon5: c.G463C:p.A155P	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 1758	M	69	KRAS	Noonan syndrome 3	Embolic stroke	NM_033360:exon5: c.T476C:p.L159S	NA	LP			Y	
CNSR30 3935	F	62	KRAS	Noonan syndrome 3	Embolic stroke	NM_033360:exon5: c.T467C:p.F156S	NA	LP			Y	
CNSR30 7860	M	47	KRAS	Noonan syndrome 3	Embolic stroke	NM_033360:exon5: c.T467C:p.F156S	NA	LP			Y	
CNSR30 0439	M	66	LMNA	Hereditary cardiomyopathies	Embolic stroke	NM_170707:exon6: c.G949A:p.E317K	P/LP	VUS		Y		
CNSR30 3884	F	30	LMNA	Hereditary cardiomyopathies	Embolic stroke	NM_170707:exon6: c.G1157A:p.R386K	P	VUS			Y	
CNSR30 6920	F	69	LMNA	Hereditary cardiomyopathies	Embolic stroke	NM_170707:exon1 1:c.G1745A:p.R582 H	P	VUS			Y	
CNSR30 0696	M	57	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon3 2:c.3628-2A>G	NA	P			Y	
CNSR30 1225	M	59	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon1 3:c.1153_1168del:p .V385fs	P/LP	LP		Y		
CNSR30 1773	M	51	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon2: c.G109T:p.G37X	NA	P		Y		
CNSR30 1808	M	68	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256.3(MY BPC3):c.821+1G>	P	P			Y	
CNSR30 3691	M	61	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon3 1:c.3624delC:p.K12 09fs	P	LP		Y		
CNSR30 4072	M	19	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon2 2:c.2308+1G>C	NA	P		Y		
CNSR30 6521	M	81	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon1 5:c.1377delC:p.P45 9fs	P	LP	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 7147	M	84	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon1 5:c.1377delC:p.P45 9fs	P	LP		Y		
CNSR30 8693	M	72	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon1 5:c.C1387T:p.Q463 X	P	P		Y		
CNSR30 9002	F	80	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon1 3:c.1153_1168del:p .V385fs	P/LP	LP	Y			
CNSR31 0109	M	89	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon1 4:c.G1256A:p.R419 H	LP	VUS			Y	
CNSR31 0251	M	58	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon1 3:c.G1187A:p.W39 6X	NA	P		Y		
CNSR31 0405	M	46	MYBP C3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon1 4:c.G1256A:p.R419 H	LP	VUS			Y	
CNSR30 0926	F	48	NKX2- 5	Tetralogy of Fallot	Embolic stroke	NM_001166176:ex on2:c.386delT:p.L1 29X	NA	LP			Y	
CNSR30 1214	M	67	NKX2- 5	Tetralogy of Fallot	Embolic stroke	NM_001166176:ex on2:c.386delT:p.L1 29X	NA	LP			Y	
CNSR30 2885	F	62	NKX2- 5	Tetralogy of Fallot	Embolic stroke	NM_001166176:ex on2:c.386delT:p.L1 29X	NA	LP			Y	
CNSR30 3343	M	68	NKX2- 5	Tetralogy of Fallot	Embolic stroke	NM_001166176:ex on2:c.C349T:p.R11 7X	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 3636	F	68	NKX2- 5	Tetralogy of Fallot	Embolic stroke	NM_001166176:ex on2:c.386delT:p.L1 29X	NA	LP			Y	
CNSR30 4524	M	54	NKX2- 5	Tetralogy of Fallot	Embolic stroke	NM_001166176:ex on2:c.386delT:p.L1 29X	NA	LP			Y	
CNSR30 9531	F	48	NKX2- 5	Tetralogy of Fallot	Embolic stroke	NM_001166176:ex on2:c.386delT:p.L1 29X	NA	LP			Y	
CNSR31 0093	F	44	NKX2- 5	Tetralogy of Fallot	Embolic stroke	NM_001166176:ex on2:c.386delT:p.L1 29X	NA	LP			Y	
CNSR31 0271	F	75	NKX2- 5	Tetralogy of Fallot	Embolic stroke	NM_001166176:ex on2:c.386delT:p.L1 29X	NA	LP			Y	
CNSR30 4051	F	34	NPPA	Hereditary cardiac dysrhythm	Embolic stroke	NM_006172:exon2: c.C319T:p.R107X	NA	LP			Y	
CNSR30 4789	F	81	NPPA	Hereditary cardiac dysrhythm	Embolic stroke	NM_006172:exon2: c.C181T:p.Q61X	NA	P			Y	
CNSR30 3354	M	45	PRKA G2	Hereditary cardiomyopathies	Embolic stroke	NM_016203:exon7: c.914delC:p.P305fs	NA	LP			Y	
CNSR30 4520	F	60	PRKA G2	Hereditary cardiomyopathies	Embolic stroke	NM_016203:exon4: c.G547A:p.E183K	P	VUS		Y		
CNSR30 8259	M	57	PRKA G2	Hereditary cardiomyopathies	Embolic stroke	NM_024429:exon7: c.445_446delinsAA :p.P149N	NA	LP			Y	
CNSR30 9141	M	60	PRKA G2	Hereditary cardiomyopathies	Embolic stroke	NM_016203:exon7: c.G922C:p.E308Q	NA	LP			Y	
CNSR30 9562	F	84	PRKA G2	Hereditary cardiomyopathies	Embolic stroke	NM_016203:exon1 3:c.A1402C:p.K468 Q	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 9699	F	52	PRKA G2	Hereditary cardiomyopathies	Embolic stroke	NM_016203:exon4: c.G547A:p.E183K	P	VUS			Y	
CNSR30	M	49	PRKAR	Carney complex, type	Embolic	NM 212472.2(PR	P	VUS			Y	
4696	IVI	49	1A	1	stroke	KAR1A):c.709- 7 709-2del		V03			1	
CNSR30 0155	M	38	RBM20	Hereditary cardiomyopathies	Embolic stroke	NM_001134363:ex on2:c.471delA:p.A 157fs	NA	LP			Y	
CNSR30 1927	M	51	RBM20	Hereditary cardiomyopathies	Embolic stroke	NM_001134363:ex on2:c.699_702del:p .K233fs	NA	LP		Y		
CNSR30 6544	M	51	RBM20	Hereditary cardiomyopathies	Embolic stroke	NM_001134363:ex on6:c.1668+1G>T	NA	P			Y	
CNSR31 0082	M	50	RBM20	Hereditary cardiomyopathies	Embolic stroke	NM_001134363:ex on10:c.2615_2616i nsAG:p.E872fs	NA	LP			Y	
CNSR30 6204	F	77	SCN1B	Hereditary cardiac dysrhythm	Embolic stroke	NM_001037.5(SC N1B):c.590+1G>A	LP	VUS			Y	
CNSR30 4738	F	68	SCN2B	Hereditary cardiac dysrhythm	Embolic stroke	NM_004588:exon2: c.G172A:p.V58M	NA	LP	Y			
CNSR30 5339	M	84	SCN2B	Hereditary cardiac dysrhythm	Embolic stroke	NM_004588:exon2: c.C142G:p.L48V	NA	LP	Y			
CNSR30 6524	M	61	SCN2B	Hereditary cardiac	Embolic stroke	NM_004588:exon2: c.G172A:p.V58M	NA	LP		Y		
CNSR30 3696	M	61	SCN3B	Hereditary cardiac dysrhythm	Embolic stroke	NM_018400:exon3: c.A326G:p.N109S	NA	LP			Y	
CNSR30 0328	M	62	SCN4B	Hereditary cardiac dysrhythm	Embolic stroke	NM_174934:exon2: c.C140A:p.T47K	NA	LP			Y	
CNSR30 1522	M	72	SCN4B	Hereditary cardiac dysrhythm	Embolic stroke	NM_174934:exon5: c.G602A:p.C201Y	NA	LP		Y		

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 2551	F	57	SCN4B	Hereditary cardiac dysrhythm	Embolic stroke	NM_174934:exon3: c.G373A:p.D125N	NA	LP			Y	
CNSR30 3785	M	71	SCN4B	Hereditary cardiac dysrhythm	Embolic stroke	NM_174934:exon4: c.G514A:p.G172R	NA	LP	Y			
CNSR30 5818	F	70	SCN4B	Hereditary cardiac dysrhythm	Embolic stroke	NM_174934:exon5: c.G607C:p.V203L	NA	LP			Y	
CNSR30 6117	F	82	SCN4B	Hereditary cardiac dysrhythm	Embolic stroke	NM_174934:exon2: c.C76T:p.P26S	NA	LP		Y		
CNSR30 9161	M	54	SCN4B	Hereditary cardiac dysrhythm	Embolic stroke	NM_174934:exon4: c.T565G:p.F189V	NA	LP			Y	
CNSR30 9232	M	65	SCN4B	Hereditary cardiac dysrhythm	Embolic stroke	NM_174934:exon4: c.T509C:p.V170A	NA	LP			Y	
CNSR30 2210	M	61	SCN5A	Hereditary cardiac dysrhythm	Embolic stroke	NM_198056:exon7: c.G845A:p.R282H	P	VUS			Y	
CNSR30 4301	M	66	SCN5A	Hereditary cardiac dysrhythm	Embolic stroke	NM_198056:exon2 4:c.4246delG:p.A1 416fs	NA	LP			Y	
CNSR30 5555	F	53	SCN5A	Hereditary cardiac dysrhythm	Embolic stroke	NM_198056:exon1 6:c.C2440T:p.R814 W	P/LP	VUS		Y		
CNSR30 5563	M	77	SCN5A	Hereditary cardiac dysrhythm	Embolic stroke	NM_198056:exon2 8:c.G4931A:p.R164 4H	P	VUS			Y	
CNSR30 9188	M	40	SCN5A	Hereditary cardiac dysrhythm	Embolic stroke	NM_198056:exon2 8:c.G4931A:p.R164 4H	P	VUS			Y	
CNSR30 0443	F	49	TLL1	Atrial septal defect	Embolic stroke	NM_012464:exon1 9:c.2566delC:p.P85 6fs	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 0397	F	68	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on326:c.C76717T:p .R25573X	LP	P			Y	
CNSR30 0902	M	51	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.T13254G:p.Y44 18X	NA	LP			Y	
CNSR30 0950	F	66	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on326:c.69869_698 70insAAGA:p.D23 290fs	NA	LP			Y	
CNSR30 1273	F	77	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on50:c.C14864G:p. S4955X	NA	P			Y	
CNSR30 1296	M	56	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on304:c.C62299T:p .Q20767X	NA	P			Y	
CNSR30 1340	M	66	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.G10819T:p. E3607X	NA	LP			Y	
CNSR30 1524	M	65	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.C11806T:p.R39 36X	NA	LP			Y	
CNSR30 1692	F	81	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.T13254G:p.Y44 18X	NA	LP			Y	
CNSR30 1708	M	51	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on322:c.C68449T:p .R22817X	P/LP	LP			Y	
CNSR30 1710	M	60	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on283:c.54989delC :p.T18330fs	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 1726	M	71	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on358:c.C104947T: p.Q34983X	LP	P			Y	
CNSR30 1743	F	65	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on61:c.G17856A:p. W5952X	NA	P			Y	
CNSR30 1816	F	61	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on240:c.44335delG :p.E14779fs	NA	LP			Y	
CNSR30 1883	M	38	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on266:c.49991delC :p.T16664fs	NA	LP			Y	
CNSR30 1922	M	80	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on62:c.C18055T:p. Q6019X	NA	P			Y	
CNSR30 2004	F	73	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on92:c.26483- 1G>A	NA	P			Y	
CNSR30 2024	F	59	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on361:c.107377+1 G>C	NA	P			Y	
CNSR30 2155	M	55	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on103:c.G29590T:p .E9864X	NA	P			Y	
CNSR30 2306	M	47	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.13482delC:p.A4 494fs	NA	LP			Y	
CNSR30 2383	M	54	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.G10819T:p. E3607X	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 2462	F	59	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.C14113T:p.R47 05X	NA	LP			Y	
CNSR30 2468	M	53	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on143:c.33826+2T >A	NA	P		Y		
CNSR30 2498	F	63	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on314:c.T66159G:p .Y22053X	NA	LP			Y	
CNSR30 2527	F	28	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on338:c.G92001A: p.W30667X	NA	P			Y	
CNSR30 2655	F	43	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on92:c.26483- 1G>A	NA	P			Y	
CNSR30 2687	F	62	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.T13254G:p.Y44 18X	NA	LP			Y	
CNSR30 2703	F	63	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on332:c.88636_886 37del:p.V29546fs	NA	LP			Y	
CNSR30 2733	M	76	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.14580delT:p.Y4 860X	NA	LP			Y	
CNSR30 2739	M	42	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.13962delG:p.T4 654fs	NA	LP			Y	
CNSR30 2803	M	65	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on346:c.96144delG :p.W32048X	NA	LP		Y		

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 2812	M	84	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on360:c.G106945T: p.E35649X	NA	P			Y	
CNSR30 2836	F	56	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133378.4:c.59 644+1G>A	LP	VUS			Y	
CNSR30 2853	M	55	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.G10819T:p. E3607X	NA	LP			Y	
CNSR30 2922	M	46	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on319:c.C67495T:p .R22499X	P/LP	P			Y	
CNSR30 3144	M	69	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on358:c.C104947T: p.Q34983X	LP	P		Y		
CNSR30 3232	M	65	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on265:c.G49659A: p.W16553X	NA	P			Y	
CNSR30 3373	M	45	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.G10819T:p. E3607X	NA	LP			Y	
CNSR30 3397	M	59	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on48:c.11884delG: p.V3962fs	NA	LP		Y		
CNSR30 3414	M	82	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on289:c.56275delA :p.T18759fs	NA	LP			Y	
CNSR30 3473	F	68	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on304:c.C63025T:p .R21009X	P/LP	P			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 3486	M	70	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on52:c.G15304T:p. G5102X	NA	P			Y	
CNSR30 3552	F	60	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on45:c.C10405T:p. Q3469X	NA	P			Y	
CNSR30 3676	M	48	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on326:c.76397_763 98del:p.I25466fs	P	LP			Y	
CNSR30 3961	M	82	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.13897dupC:p.Q 4633fs	NA	LP			Y	
CNSR30 3985	F	75	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on159:c.G35500T:p .E11834X	NA	P			Y	
CNSR30 4024	M	72	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on138:c.33340+2T >C	NA	P			Y	
CNSR30 4056	F	51	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550.2:c .49345+2T>C	LP	VUS			Y	
CNSR30 4056	F	51	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.12571delA:p.T4 191fs	NA	LP			Y	
CNSR30 4076	F	70	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on271:c.C51436T:p .Q17146X	P	P			Y	
CNSR30 4115	F	72	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on248:c.46201_462 04del:p.T15401fs	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 4236	M	72	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.G10819T:p. E3607X	NA	LP			Y	
CNSR30 4295	M	46	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on100:c.28907delG :p.C9636fs	NA	LP			Y	
CNSR30 4443	F	66	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on251:c.46821_469 06del:p.P15607fs	NA	LP			Y	
CNSR30 4630	F	77	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on87:c.25083_2508 6del:p.F8361fs	NA	LP		Y		
CNSR30 4797	M	57	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.13550delT:p.M 4517fs	NA	LP			Y	
CNSR30 5023	M	72	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on63:c.C18541T:p. R6181X	NA	P			Y	
CNSR30 5065	F	68	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.14463_14464de l:p.T4821fs	NA	LP			Y	
CNSR30 5258	M	63	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550.2:c .49345+2T>C	LP	VUS			Y	
CNSR30 5258	M	63	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.12571delA:p.T4 191fs	NA	LP			Y	
CNSR30 5322	F	53	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_003319:exon4 5:c.11505_11506ins AATTAATTCATT AACA:p.V3836_E 3837delinsNX	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 5448	M	61	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on45:c.C10405T:p. Q3469X	NA	P			Y	
CNSR30 5694	M	60	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on55:c.16112delA: p.N5371fs	NA	LP			Y	
CNSR30 5735	M	62	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on352:c.98381delG :p.G32794fs	NA	LP			Y	
CNSR30 5812	M	71	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on163:c.35813delA :p.K11938fs	NA	LP			Y	
CNSR30 6351	M	57	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550.2:c .49345+2T>C	LP	VUS			Y	
CNSR30 6351	M	57	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.12571delA:p.T4 191fs	NA	LP			Y	
CNSR30 6385	F	67	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on56:c.C16404A:p. C5468X	NA	P			Y	
CNSR30 6470	M	70	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on358:c.C104947T: p.Q34983X	LP	P			Y	
CNSR30 6608	M	48	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on63:c.C18541T:p. R6181X	NA	P			Y	
CNSR30 7049	M	79	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on280:c.C54067T:p .R18023X	LP	P			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 7096	M	78	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on92:c.26483- 2A>G	NA	P		Y		
CNSR30 7169	M	61	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.11782_11783del :p.M3928fs	NA	LP			Y	
CNSR30 7344	F	70	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.11416_11419del :p.I3806fs	NA	LP			Y	
CNSR30 7493	F	68	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on347:c.96504delA :p.G32168fs	NA	LP			Y	
CNSR30 7561	M	55	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on322:c.68527+1G >T	NA	P			Y	
CNSR30 7677	M	65	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.T13254G:p.Y44 18X	NA	LP			Y	
CNSR30 7734	M	63	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550.2:c .49345+2T>C	LP	VUS			Y	
CNSR30 7734	M	63	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.12571delA:p.T4 191fs	NA	LP			Y	
CNSR30 7750	M	52	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on339:c.A92854T:p .R30952X	NA	P			Y	
CNSR30 7875	M	73	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.G10819T:p. E3607X	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 8005	M	48	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.C10735T:p. Q3579X	NA	P			Y	
CNSR30 8032	M	72	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.G10819T:p. E3607X	NA	LP		Y		
CNSR30 8096	M	58	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.G10819T:p. E3607X	NA	LP			Y	
CNSR30 8211	M	87	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on91:c.G26254T:p. E8752X	NA	P			Y	
CNSR30 8221	M	75	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on137:c.33247+1G >A	NA	P			Y	
CNSR30 8406	F	80	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on324:c.69179_691 80insTTAC:p.T230 60fs	NA	LP			Y	
CNSR30 8440	F	58	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on360:c.G106914A :p.W35638X	NA	P			Y	
CNSR30 8503	F	65	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on343:c.95246delA :p.E31749fs	NA	LP			Y	
CNSR30 8517	M	47	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on328:c.87432delT: p.P29144fs	NA	LP			Y	
CNSR30 8532	M	59	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on93:c.26979dupT: p.G8994fs	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 8562	M	78	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.14236delA:p.T4 746fs	NA	LP			Y	
CNSR30 8731	F	72	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on82:c.23896_2389 7del:p.S7966fs	NA	LP			Y	
CNSR30 8774	M	73	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on87:c.25113delG: p.E8371fs	NA	LP			Y	
CNSR30 8791	M	64	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on7:c.1205delC:p.A 402fs	NA	LP			Y	
CNSR30 9122	M	50	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on11:c.1732dupG:p .E578fs	NA	LP			Y	
CNSR30 9205	M	72	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.15075delT:p.V5 025fs	NA	LP			Y	
CNSR30 9225	F	53	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.C11806T:p.R39 36X	NA	LP			Y	
CNSR30 9332	M	52	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on354:c.C99052T:p .Q33018X	NA	P			Y	
CNSR30 9344	M	65	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550.2:c .49345+2T>C	LP	VUS			Y	
CNSR30 9344	M	65	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.12571delA:p.T4 191fs	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 9386	F	71	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.G10819T:p. E3607X	NA	LP			Y	
CNSR30 9744	M	48	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on335:c.90597delA :p.G30199fs	NA	LP			Y	
CNSR30 9871	M	46	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.G10819T:p. E3607X	NA	LP			Y	
CNSR31 0007	M	61	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.C10522T:p.Q35 08X	NA	LP			Y	
CNSR31 0370	M	58	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:ex on46:c.G10819T:p. E3607X	NA	LP			Y	
CNSR31 0422	M	62	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon4 6:c.T13254G:p.Y44 18X	NA	LP			Y	
CNSR30 2718	M	66	ZFPM2	Tetralogy of Fallot	Embolic stroke	NM_012082:exon6: c.739+1G>A	NA	P			Y	
CNSR31 0062	M	60	ZFPM2	Tetralogy of Fallot	Embolic stroke	NM_012082:exon7: c.G779A:p.R260Q	P	VUS			Y	
CNSR30 3911	M	82	ACTA2	Aortic aneurysm, familial thoracic 6	Large artery disease	NM_001613:exon3: c.170delG:p.G57fs	NA	LP	Y			
CNSR30 0368	M	64	APOA5	Hyperchylomicronem ia, late-onset	Large artery disease	NM_052968:exon2: c.G30A:p.W10X	NA	P			Y	
CNSR30 7179	F	50	APOA5	Hyperchylomicronem ia, late-onset	Large artery disease	NM_052968:exon2: c.G30A:p.W10X	NA	P		Y		
CNSR30 0355	F	79	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon2: c.T222G:p.Y74X	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 1124	M	75	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	
CNSR30 1461	F	53	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon6: c.G537A:p.W179X	NA	P			Y	
CNSR30 1827	M	35	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon2: c.T222G:p.Y74X	NA	LP			Y	
CNSR30 2864	F	64	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon2: c.T222G:p.Y74X	NA	LP			Y	
CNSR30 3299	M	51	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	
CNSR30 3732	F	48	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon1 1:c.1102delC:p.P36 8fs	NA	LP			Y	
CNSR30 4614	M	62	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.786dupC:p.L262f s	NA	LP			Y	
CNSR30 4790	F	73	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	
CNSR30 5067	M	55	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.786delC:p.L262fs	NA	LP			Y	
CNSR30 5247	M	76	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	
CNSR30 5437	M	77	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.C853T:p.R285X	NA	LP			Y	
CNSR30 5830	M	45	СЕТР	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon2: c.T222G:p.Y74X	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 5982	M	75	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon1 1:c.982-1G>C	NA	LP			Y	
CNSR30 6316	F	46	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon1 4:c.1321+1G>A	P	P			Y	
CNSR30 6620	F	54	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon1 2:c.1208delA:p.D4 03fs	NA	LP			Y	
CNSR30 6695	M	59	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon2: c.T222G:p.Y74X	NA	LP			Y	
CNSR30 6834	M	85	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon2: c.T222G:p.Y74X	NA	LP			Y	
CNSR30 7652	M	70	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	
CNSR30 7689	M	50	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	
CNSR30 7818	M	88	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	
CNSR30 7964	M	47	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon1 4:c.1321+1G>A	P	P			Y	
CNSR30 8010	M	82	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon2: c.C160T:p.R54X	NA	LP			Y	
CNSR30 8189	M	64	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon2: c.T222G:p.Y74X	NA	LP			Y	
CNSR30 8467	F	81	СЕТР	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 9068	M	56	СЕТР	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	
CNSR30 9319	F	75	СЕТР	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	
CNSR30 9412	F	76	СЕТР	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	
CNSR30 9459	M	46	СЕТР	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon1 1:c.1102delC:p.P36 8fs	NA	LP			Y	
CNSR30 9700	M	27	CETP	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon2: c.T222G:p.Y74X	NA	LP			Y	
CNSR31 0111	F	79	СЕТР	Hyperalphalipoprotei nemia	Large artery disease	NM_000078:exon9: c.783_786del:p.D2 61fs	NA	LP			Y	
CNSR30 4557	F	56	COL1A 1	CeAD,FMD,TAA,A AA	Large artery disease	NM_000088:exon5: c.441delC:p.P147fs	P	LP		Y		
CNSR30 6617	M	49	COL1A 1	CeAD,FMD,TAA,A AA	Large artery disease	NM_000088:exon3 7:c.G2594A:p.R865 H	LP	VUS			Y	
CNSR30 6717	F	63	COL1A 1	CeAD,FMD,TAA,A AA	Large artery disease	NM_000088:exon5: c.386_388delinsT:p .G129Lfs*39	NA	P			Y	
CNSR30 9324	M	52	COL1A 1	CeAD,FMD,TAA,A AA	Large artery disease	NM_000088:exon2 1:c.1354-1G>C	NA	P		Y		
CNSR30 5489	F	77	COL1A 2	CeAD,FMD,TAA,A AA	Large artery disease	NM_000089:exon1 9:c.G946A:p.G316 S	P	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 6054	M	49	COL1A 2	CeAD,FMD,TAA,A AA	Large artery disease	NM_000089:exon1 7:c.G874A:p.G292 S	P	VUS		Y		
CNSR30 6112	M	69	COL1A 2	CeAD,FMD,TAA,A AA	Large artery disease	NM_000089:exon5: c.207delC:p.G69fs	NA	LP		Y		
CNSR30 4228	M	72	COL5A	Ehlers-Danlos syndrome, classic type,	Large artery disease	NM_000093:exon6 6:c.T5486G:p.F182 9C	LP	VUS			Y	
CNSR30 6147	F	67	COL5A 1	Ehlers-Danlos syndrome, classic type,	Large artery disease	NM_001278074:ex on64:c.C5095T:p.Q 1699X	NA	P			Y	
CNSR30 8149	M	71	COL5A	Ehlers-Danlos syndrome, classic type,	Large artery disease	NM_000093:exon4 8:c.G3781A:p.G12 61R	LP	VUS			Y	
CNSR30 0432	M	48	ELN	SVAS	Large artery disease	NM_000501:exon1 4:c.686-2A>G	NA	P			Y	
CNSR30 0818	M	83	ELN	SVAS	Large artery disease	NM_000501:exon2 4:c.1621+1G>A	LP	P			Y	
CNSR30 1992	M	72	ELN	SVAS	Large artery disease	NM_000501:exon1 4:c.686- 2A>G2A>G	NA	P			Y	
CNSR30 2656	M	50	ELN	SVAS	Large artery disease	NM_000501:exon1 4:c.686-2A>G	NA	P			Y	
CNSR30 5968	F	78	ELN	SVAS	Large artery disease	NM_001278939:ex on26:c.1933+2T>A	NA	P		Y		
CNSR30 6387	M	55	ELN	SVAS	Large artery disease	NM_001278939:ex on26:c.1933+2T>A	NA	P		Y		
CNSR30 7595	M	60	ELN	SVAS	Large artery disease	NM_000501:exon1 4:c.686-2A>G	NA	P			Y	
CNSR30 8754	M	39	ELN	SVAS	Large artery disease	NM_000501:exon1 1:c.571+1G>A	NA	P			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR31 0118	M	45	ELN	SVAS	Large artery disease	NM_000501:exon2 4:c.1621+1G>A	LP	P	Y			
CNSR30 0886	M	75	FBN1	Marfan syndrome	Large artery disease	NM_000138:exon2 2:c.A2613C:p.L871 F	P	VUS			Y	
CNSR30 1821	M	64	FBN1	Marfan syndrome	Large artery disease	NM_000138:exon6 3:c.T7754C:p.I2585 T	P/LP	VUS		Y		
CNSR30 2393	M	51	FBN1	Marfan syndrome	Large artery disease	NM_000138:exon2 2:c.A2613C:p.L871 F	P	VUS			Y	
CNSR30 3067	F	58	FBN1	Marfan syndrome	Large artery disease	NM_000138:exon2 2:c.A2613C:p.L871 F	P	VUS			Y	
CNSR30 4747	M	73	FBN1	Marfan syndrome	Large artery disease	NM_000138:exon2 2:c.A2613C:p.L871 F	P	VUS			Y	
CNSR30 5922	M	50	FBN1	Marfan syndrome	Large artery disease	NM_000138:exon2 2:c.A2613C:p.L871 F	P	VUS			Y	
CNSR30 9171	M	78	FBN1	Marfan syndrome	Large artery disease	NM_000138:exon1 0:c.G1091A:p.R364 Q	NA	LP			Y	
CNSR30 0049	M	53	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon9: c.C1216A:p.R406R	P	VUS			Y	
CNSR30 0051	F	42	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 1:c.A1691G:p.N56 4S	LP	VUS	Y			
CNSR30 0104	F	65	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 1:c.T1592A:p.M53 1K	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 0212	F	48	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 1:c.T1592A:p.M53 1K	NA	LP			Y	
CNSR30 0289	M	62	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 0:c.A1454T:p.H485 L	NA	LP		Y		
CNSR30 0310	M	41	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon4: c.510delC:p.D170fs	P	LP	Y			
CNSR30 0371	F	76	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon4: c.C516A:p.D172E	NA	LP			Y	
CNSR30 0418	M	58	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 6:c.A2344T:p.K782 X	P	LP		Y		
CNSR30 0451	F	81	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 0:c.A1525G:p.K50 9E	LP	VUS	Y			
CNSR30 0867	F	73	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 3:c.G1879A:p.A62 7T	P/LP	VUS			Y	
CNSR30 1304	M	57	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 1:c.T1592A:p.M53 1K	NA	LP		Y		
CNSR30 1669	F	70	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 1:c.A1691G:p.N56 4S	LP	VUS		Y		
CNSR30 1850	M	67	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon7: c.T1016C:p.L339P	LP	VUS	Y			
CNSR30 2713	M	42	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon4: c.C487T:p.Q163X	Р	LP	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 3264	M	60	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 3:c.C1880T:p.A627 V	LP	VUS	Y			
CNSR30 3708	F	69	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 3:c.G1864T:p.D622 Y	NA	LP			Y	
CNSR30 4012	M	82	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 1:c.T1592A:p.M53 1K	NA	LP			Y	
CNSR30 4637	F	70	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 3:c.G1879A:p.A62 7T	P/LP	VUS		Y		
CNSR30 4929	M	79	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon6: c.A924T:p.E308D	LP	VUS			Y	
CNSR30 5027	F	64	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon9: c.G1247A:p.R416Q	P/LP	VUS	Y			
CNSR30 5733	M	59	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon4: c.T400C:p.C134R	P/LP	VUS			Y	
CNSR30 6253	M	83	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 4:c.G2026C:p.G676 R	LP	VUS	Y			
CNSR30 6440	M	62	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon7: c.C1048T:p.R350X	P	LP	Y			
CNSR30 6967	M	56	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon7: c.G1049A:p.R350Q		VUS			Y	
CNSR30 7020	F	52	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_001195798.2:c .313+1dup	P/LP	VUS		Y		
CNSR30 7426	M	54	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon9: c.G1247A:p.R416Q	P/LP	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 7613	F	71	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 4:c.G2026C:p.G676 R	LP	VUS	Y			
CNSR30 7649	M	49	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 2:c.1745_1746del:p .L582fs	P	LP	Y			
CNSR30 7861	M	66	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 0:c.1570_1582del:p .V524fs	NA	LP		Y		
CNSR30 8053	M	69	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon1 4:c.G2026C:p.G676 R	LP	VUS			Y	
CNSR30 8235	M	52	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon9: c.C1216A:p.R406R	P	VUS	Y			
CNSR30 8766	M	61	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon4: c.G502A:p.D168N	P/LP	VUS			Y	
CNSR30 9242	M	53	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon2: c.T100G:p.C34G	P/LP	VUS			Y	
CNSR30 9264	M	42	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon5: c.G796C:p.D266H	NA	LP		Y		
CNSR30 9633	M	69	LDLR	Hypercholesterolemia , familial, 1	Large artery disease	NM_000527:exon6: c.908delG:p.R303fs	NA	LP			Y	
CNSR30 0114	M	77	MFAP5	Aortic aneurysm, familial thoracic 9	Large artery disease	NM_003480:exon3: c.C88T:p.R30X	LP	VUS			Y	
CNSR30 1352	M	45	MFAP5	Aortic aneurysm, familial thoracic 9	Large artery disease	NM_003480:exon1 0:c.C472T:p.R158X	P	VUS			Y	
CNSR30 1444	M	41	MFAP5	Aortic aneurysm, familial thoracic 9	Large artery disease	NM_003480:exon1 0:c.C472T:p.R158X	P	VUS			Y	
CNSR30 2503	F	68	MFAP5	Aortic aneurysm, familial thoracic 9	Large artery disease	NM_003480:exon1 0:c.C472T:p.R158X	P	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 2574	M	77	MFAP5	Aortic aneurysm, familial thoracic 9	Large artery disease	NM_003480:exon1 0:c.C472T:p.R158X	P	VUS			Y	
CNSR30 3517	F	65	MFAP5	Aortic aneurysm, familial thoracic 9	Large artery disease	NM_003480:exon9: c.338dupT:p.L113fs	NA	LP			Y	
CNSR30 5631	M	65	MFAP5	Aortic aneurysm, familial thoracic 9	Large artery disease	NM_003480:exon1 0:c.C472T:p.R158X	P	VUS			Y	
CNSR30 6162	M	57	MFAP5	Aortic aneurysm, familial thoracic 9	Large artery disease	NM_003480:exon1 0:c.C472T:p.R158X	P	VUS			Y	
CNSR30 6421	F	53	MFAP5	Aortic aneurysm, familial thoracic 9	Large artery disease	NM_003480:exon1 0:c.C472T:p.R158X	P	VUS			Y	
CNSR30 0828	M	73	MYH11	Aortic aneurysm, familial thoracic 4	Large artery disease	NM_002474:exon2 8:c.T3791A:p.L126 4O	NA	LP			Y	
CNSR30 6393	M	44	MYH11	Aortic aneurysm, familial thoracic 4	Large artery disease	NM_002474:exon1 9:c.A2254T:p.K752 X	NA	P			Y	
CNSR30 7728	M	48	MYH11	Aortic aneurysm, familial thoracic 4	Large artery disease	NM_002474:exon2 6:c.G3466T:p.E115 6X	NA	P			Y	
CNSR30 0206	M	54	MYLK	Aortic aneurysm, familial thoracic 7	Large artery disease	NM_053025:exon1 6:c.C2371T:p.Q791 X	NA	P			Y	
CNSR30 2047	F	73	MYLK	Aortic aneurysm, familial thoracic 7	Large artery disease	NM_053025:exon1 8:c.2463-2A>G	NA	P			Y	
CNSR30 4181	F	64	MYLK	Aortic aneurysm, familial thoracic 7	Large artery disease	NM_053025:exon1 2:c.1517-2A>G	NA	P			Y	
CNSR30 5870	F	61	MYLK	Aortic aneurysm, familial thoracic 7	Large artery disease	NM_053025:exon1 8:c.C2692T:p.R898 X	NA	P			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 7523	F	67	MYLK	Aortic aneurysm, familial thoracic 7	Large artery disease	NM_053025:exon1 4:c.C1915T:p.Q639 X	NA	P			Y	
CNSR30 7730	M	56	MYLK	Aortic aneurysm, familial thoracic 7	Large artery disease	NM_053025:exon1 8:c.C2665T:p.Q889 X	NA	P			Y	
CNSR30 9070	M	61	MYLK	Aortic aneurysm, familial thoracic 7	Large artery disease	NM_053025:exon6: c.422+1G>A	NA	P			Y	
CNSR30 9296	M	58	MYLK	Aortic aneurysm, familial thoracic 7	Large artery disease	NM_053025:exon1 0:c.1228dupG:p.D4 10fs	NA	LP			Y	
CNSR30 0785	M	76	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon1 3:c.T1473A:p.H491 Q	NA	LP			Y	
CNSR30 2491	F	48	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon2: c.A320G:p.D107G	NA	LP			Y	
CNSR30 3483	M	72	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon8: c.T962C:p.I321T	NA	LP			Y	
CNSR30 3769	M	65	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon2: c.A320G:p.D107G	NA	LP			Y	
CNSR30 4124	F	58	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon3: c.T541C:p.F181L	NA	LP			Y	
CNSR30 5284	F	68	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon8: c.T962C:p.I321T	NA	LP			Y	
CNSR30 5514	F	80	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon2: c.A350G:p.D117G	NA	LP			Y	
CNSR30 5685	F	57	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon3: c.T539C:p.V180A	NA	LP			Y	
CNSR30 6077	F	65	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon8: c.T962C:p.I321T	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 6372	М	60	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon2: c.G391A:p.V131M	NA	LP			Y	
CNSR30 7814	M	60	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon1 0:c.1128delC:p.N37 6fs	NA	LP			Y	
CNSR30 9202	F	44	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_001098512:ex on1:c.A257C:p.K86 T	NA	LP			Y	
CNSR30 9352	F	49	PRKG1	Aortic aneurysm, familial thoracic 8	Large artery disease	NM_006258:exon2: c.A320G:p.D107G	NA	LP		Y		
CNSR30 0151	M	61	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 0449	F	65	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 0542	M	65	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 0570	F	60	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 0650	M	71	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 1176	M	60	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS		Y		
CNSR30 1302	M	66	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS		Y		

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 1408	M	70	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 1446	M	57	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 1456	F	46	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 1695	M	45	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS		Y		
CNSR30 1768	M	62	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 2170	M	37	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS				Y
CNSR30 2455	F	27	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS	Y			
CNSR30 2619	M	65	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 2741	F	67	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS		Y		
CNSR30 2775	M	35	RNF21 3	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 3338	M	65	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 3394	M	54	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS				Y
CNSR30 3404	M	57	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 3451	F	53	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS	Y			
CNSR30 3695	F	70	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS		Y		
CNSR30 4133	F	54	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 4137	F	60	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 4305	M	61	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 4368	M	56	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 4370	M	67	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 4477	M	69	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS				Y
CNSR30 4840	M	30	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS	Y			
CNSR30 5285	M	52	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 5511	M	45	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 5688	F	51	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 5701	F	58	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 5969	F	67	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS	Y			
CNSR30 6018	M	79	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 6251	F	48	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS	Y			
CNSR30 6474	M	51	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS				Y

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 6719	F	56	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 7009	F	65	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS				Y
CNSR30 7043	M	62	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 7110	M	62	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 7307	M	72	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 7323	M	66	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 7418	F	68	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS		Y		
CNSR30 7541	F	61	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 8107	M	51	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 8482	M	53	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 8599	M	59	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 8643	M	60	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 8924	F	44	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS	Y			
CNSR30 9034	M	58	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 9437	F	51	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS		Y		
CNSR30 9554	F	53	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 9631	M	62	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS	Y			
CNSR31 0183	M	41	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS		Y		
CNSR31 0309	M	51	RNF21	Moyamoya disease	Large artery disease	NM_001256071:ex on60:c.G14429A:p. R4810K	P	VUS			Y	
CNSR30 3114	M	71	ETV6	Thrombocytopenia 5	Prothrombot ic state	NM_001987:exon2: c.C115T:p.R39X	NA	P			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 0244	F	60	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon1 0:c.C1195T:p.R399 C	NA	LP			Y	
CNSR30 0779	M	60	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon1 3:c.G1679A:p.R560 Q	NA	LP			Y	
CNSR30 1431	M	49	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon4: c.G290A:p.R97Q	NA	LP			Y	
CNSR30 2043	M	75	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon7: c.A715G:p.S239G	NA	LP			Y	
CNSR30 2610	F	62	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon7: c.A715G:p.S239G	NA	LP			Y	
CNSR30 2706	M	65	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon4: c.G290A:p.R97Q	NA	LP			Y	
CNSR30 2815	F	64	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon7: c.C683T:p.T228I	NA	LP			Y	
CNSR30 2834	M	54	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon1 1:c.G1307A:p.R436 Q	NA	LP			Y	
CNSR30 3002	M	53	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon4: c.G290A:p.R97Q	NA	LP			Y	
CNSR30 3616	M	70	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon7: c.A715G:p.S239G	NA	LP			Y	
CNSR30 3837	M	68	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon2: c.A136G:p.T46A	NA	LP			Y	
CNSR30 4831	M	45	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon4: c.G290A:p.R97Q	NA	LP			Y	
CNSR30 4932	M	64	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon1 0:c.G1196T:p.R399 L	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 5605	M	66	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon4: c.G290A:p.R97Q	NA	LP			Y	
CNSR30 6073	M	51	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon7: c.A715G:p.S239G	NA	LP		Y		
CNSR30 6469	M	69	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon7: c.C677A:p.A226E	NA	LP			Y	
CNSR30 6519	F	71	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon1 1:c.G1299T:p.R433 S	NA	LP			Y	
CNSR30 6646	F	81	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon5: c.419_420insTGA G:p.P140fs	NA	LP			Y	
CNSR30 6759	M	44	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon1 1:c.G1299T:p.R433 S	NA	LP			Y	
CNSR30 7305	M	45	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon1 1:c.C1306T:p.R436 X	NA	P			Y	
CNSR30 7437	M	82	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon2: c.G119A:p.R40Q	NA	LP			Y	
CNSR30 7824	M	76	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon9: c.G1054A:p.E352K	P	LP			Y	
CNSR30 7908	M	58	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon1 0:c.C1147T:p.R383 W	NA	LP			Y	
CNSR30 8145	F	74	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon4: c.G290A:p.R97Q	NA	LP			Y	
CNSR30 8905	M	56	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon7: c.A715G:p.S239G	NA	LP			Y	
CNSR30 8999	M	62	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon8: c.G1003A:p.D335N	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 9128	M	52	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon1 1:c.G1299T:p.R433 S	NA	LP			Y	
CNSR31 0389	M	53	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon7: c.A715G:p.S239G	NA	LP			Y	
CNSR31 0428	M	60	F2	Thrombophilia due to thrombin defect	Prothrombot ic state	NM_000506:exon2: c.C239T:p.T80M	NA	LP			Y	
CNSR30 1130	F	87	GP1BA	von Willebrand disease, platelet-type	Prothrombot ic state	NM_000173:exon2: c.A449G:p.N150S	LP	VUS			Y	
CNSR30 0210	M	54	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 0974	F	61	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 1140	F	67	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 1453	M	74	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 1596	M	81	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 1724	F	53	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 1871	M	70	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	Р	VUS	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 2013	F	62	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 2241	M	76	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 2246	F	43	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 2755	F	59	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 2832	M	63	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 2950	F	60	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 3845	M	61	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 3914	F	51	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 4686	M	64	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 6131	F	76	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	Р	VUS	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 6885	M	65	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 6975	M	53	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 6992	F	62	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 7039	M	66	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 7258	M	73	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 7566	F	62	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 8276	M	67	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 8823	M	67	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS			Y	
CNSR30 8929	M	79	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS			Y	
CNSR30 9106	M	77	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	Р	VUS	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 9615	F	79	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS			Y	
CNSR30 9872	M	75	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR31 0420	M	69	JAK2	Thrombocythemia 3	Prothrombot ic state	NM_004972:exon1 4:c.G1849T:p.V617 F	P	VUS	Y			
CNSR30 5210	M	58	PROC	Thrombophilia due to protein C deficiency, autos	Prothrombot ic state	NM_000312:exon9: c.G889C:p.D297H	P	VUS			Y	
CNSR30 6872	F	64	PROC	Thrombophilia due to protein C deficiency, autos	Prothrombot ic state	NM_000312:exon3: c.G199C:p.E67Q	NA	LP			Y	
CNSR30 7697	M	58	PROC	Thrombophilia due to protein C deficiency, autos	Prothrombot ic state	NM_000312:c.678 +9C>T	P	VUS			Y	
CNSR30 7931	M	29	PROC	Thrombophilia due to protein C deficiency, autos	Prothrombot ic state	NM_000312:exon7: c.C658T:p.R220W	P	VUS			Y	
CNSR30 9298	F	64	PROC	Thrombophilia due to protein C deficiency, autos	Prothrombot ic state	NM_000312:exon3: c.G76C:p.V26L	NA	LP			Y	
CNSR31 0362	M	74	PROC	Thrombophilia due to protein C deficiency, autos	Prothrombot ic state	NM_000312:exon9: c.G889C:p.D297H	P	VUS			Y	
CNSR30 3734	M	40	PROS1	Thrombophilia due to protein S deficiency	Prothrombot ic state	NM_000313:exon1 0:c.C1063T:p.R355 C	Р	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 3874	M	48	PROS1	Thrombophilia due to protein S deficiency	Prothrombot ic state	NM_000313:exon6: c.A586G:p.K196E	P	VUS			Y	
CNSR30 4216	M	65	PROS1	Thrombophilia due to protein S deficiency	Prothrombot ic state	NM_000313:exon1 4:c.T1680A:p.Y560 X	LP	LP			Y	
CNSR30 4306	F	61	PROS1	Thrombophilia due to protein S deficiency	Prothrombot ic state	NM_000313:exon1 0:c.C1063T:p.R355 C	P	VUS			Y	
CNSR30 5014	F	67	PROS1	Thrombophilia due to protein S deficiency	Prothrombot ic state	NM_000313:exon1 0:c.C1063T:p.R355 C	P	VUS			Y	
CNSR30 7585	M	56	PROS1	Thrombophilia due to protein S deficiency	Prothrombot ic state	NM_000313:exon1 4:c.1753delG:p.E58 5fs	NA	LP			Y	
CNSR30 7679	M	63	PROS1	Thrombophilia due to protein S deficiency	Prothrombot ic state	NM_000313:exon1 0:c.C1063T:p.R355 C	P	VUS			Y	
CNSR30 8628	F	59	PROS1	Thrombophilia due to protein S deficiency	Prothrombot ic state	NM_000313:exon1 2:c.1427_1428insA :p.L476fs	NA	LP			Y	
CNSR30 0805	M	53	SERPI NC1	Thrombophilia due to antithrombin III deficiency	Prothrombot ic state	NM_000488:exon2: c.C218T:p.P73L	P/LP	VUS			Y	
CNSR30 1590	M	70	SERPI NC1	Thrombophilia due to antithrombin III deficiency	Prothrombot ic state	NM_000488:exon3: c.T442C:p.S148P	P	VUS			Y	
CNSR30 5745	M	71	SERPI NC1	Thrombophilia due to antithrombin III deficiency	Prothrombot ic state	NM_000488:exon3: c.T442C:p.S148P	P	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 5890	F	41	SERPI NC1	Thrombophilia due to antithrombin III deficiency	Prothrombot ic state	NM_000488:exon3: c.A572G:p.Q191R	LP	VUS			Y	
CNSR30 6312	M	50	SERPI NC1	Thrombophilia due to antithrombin III deficiency	Prothrombot ic state	NM_000488:exon2: c.C235T:p.R79C	LP	VUS			Y	
CNSR30 9720	M	60	SERPI NC1	Thrombophilia due to antithrombin III deficiency	Prothrombot ic state	NM_000488:exon3: c.A572G:p.Q191R	LP	VUS			Y	
CNSR30 0699	F	69	SERPI ND1	Thrombophilia due to heparin cofactor II deficie	Prothrombot ic state	NM_000185:exon2: c.657_660del:p.R21 9fs	NA	LP			Y	
CNSR30 2505	M	54	SERPI ND1	Thrombophilia due to heparin cofactor II deficie	Prothrombot ic state	NM_000185:exon2: c.C415T:p.R139X	NA	P			Y	
CNSR30 4069	M	62	SERPI ND1	Thrombophilia due to heparin cofactor II deficie	Prothrombot ic state	NM_000185:exon2: c.556_557del:p.F18 6fs	NA	LP			Y	
CNSR30 5242	F	70	SERPI ND1	Thrombophilia due to heparin cofactor II deficie	Prothrombot ic state	NM_000185:exon3: c.949_950del:p.E31 7fs	NA	LP			Y	
CNSR30 7508	M	55	SERPI ND1	Thrombophilia due to heparin cofactor II deficie	Prothrombot ic state	NM_000185:exon3: c.917delA:p.E306fs	NA	LP			Y	
CNSR30 9254	M	66	SERPI ND1	Thrombophilia due to heparin cofactor II deficie	Prothrombot ic state	NM_000185:exon2: c.668_677del:p.D2 23fs	NA	LP			Y	
CNSR30 5747	M	62	STIM1	Stormorken syndrome	Prothrombot ic state	NM_001277961:ex on11:c.1621_1624d el:p.S541fs	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 0050	M	60	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 0111	M	59	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.2289dupG:p.S7 64	NA	LP			Y	
CNSR30 0534	M	68	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 4:c.C1677A:p.C559 X	NA	LP			Y	
CNSR30 0609	F	74	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 0627	F	67	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon2 8:c.C4135T:p.R137 9C	P	VUS			Y	
CNSR30 1287	M	48	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 1361	M	62	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon3 2:c.5462delC:p.T18 2	NA	LP			Y	
CNSR30 1580	M	74	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon2 8:c.C4696T:p.R156 6X	P	P			Y	
CNSR30 1972	M	66	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 2390	M	54	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon4 3:c.C7390T:p.R246 4C	P/LP	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 2557	F	67	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 3068	M	73	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon2 8:c.C3797A:p.P126 6Q	LP	VUS			Y	
CNSR30 3106	M	60	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 3246	F	64	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.C2372T:p.T791 M	LP	VUS			Y	
CNSR30 3642	M	64	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 3701	M	77	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 3871	M	63	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon4 4:c.C7464T:p.G248 8G	LP	VUS			Y	
CNSR30 4193	M	68	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 4338	F	57	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon2 8:c.C3797A:p.P126 6Q	LP	VUS			Y	
CNSR30 4493	M	64	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 4667	M	52	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 4733	M	75	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon2 0:c.2658delC:p.P88 6fs	NA	LP			Y	
CNSR30 4865	M	50	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 5150	M	63	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon4 3:c.G7332A:p.W24 44X	NA	P			Y	
CNSR30 6284	M	68	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 6560	M	76	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 6641	F	70	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon4 4:c.C7464T:p.G248 8G	LP	VUS			Y	
CNSR30 6919	M	68	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon2 0:c.G2561A:p.R854 Q	P/LP	VUS			Y	
CNSR30 7271	M	63	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon3 9:c.C6835T:p.Q227 9X	NA	P			Y	
CNSR30 8117	F	79	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 8135	M	53	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon2 8:c.T3774A:p.Y125 8X	NA	LP			Y	
CNSR30 8576	M	25	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon1 8:c.G2303A:p.R768 Q	LP	VUS			Y	
CNSR30 8978	F	75	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon7: c.C813G:p.Y271X	NA	LP			Y	
CNSR30 9553	M	62	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon2 8:c.G3970A:p.G13 24S	Р	VUS			Y	
CNSR30 9563	M	50	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon2 8:c.4094delT:p.L13 65fs	NA	LP			Y	
CNSR31 0361	M	62	VWF	von Willebrand disease, type 1	Prothrombot ic state	NM_000552:exon3 0:c.G5235A:p.W17 45X	NA	P			Y	
CNSR30 1005	M	51	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001303110:ex on6:c.G343A:p.G11 5S	NA	LP	Y			
CNSR30 2264	M	73	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001303110:ex on9:c.G502A:p.G1 68R	NA	LP	Y			
CNSR30 2968	F	81	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001845:exon5 0:c.G4718A:p.G15 73E	NA	LP	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 2973	M	58	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001303110:ex on15:c.G823A:p.G 275	NA	LP			Y	
CNSR30 3061	F	55	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001303110:ex on9:c.G502A:p.G1 68R	NA	LP			Y	
CNSR30 3631	F	79	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001303110:ex on21:c.G1277T:p.G 42	NA	LP	Y			
CNSR30 3650	F	88	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001845:exon2 7:c.G1937C:p.G646 A	NA	LP	Y			
CNSR30 4740	M	52	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001845:exon4 0:c.3407-1G>A	NA	P			Y	
CNSR30 4947	M	43	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001845:exon3 7:c.C3187T:p.R106 3X	NA	LP	Y			
CNSR30 5943	M	70	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001303110:ex on6:c.G343A:p.G11 5S	NA	LP	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 6510	M	78	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001845:exon4 9:c.G4471A:p.G14 91S	NA	LP			Y	
CNSR30 7106	M	58	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001303110:ex on23:c.G1420A:p. G474R	NA	LP	Y			
CNSR30 8031	M	70	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001845:exon2 7:c.G1937C:p.G646 A	NA	LP		Y		
CNSR30 8223	F	69	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001845:exon2 5:c.G1640C:p.G547 A	NA	LP			Y	
CNSR30 9282	F	76	COL4A	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001303110:ex on15:c.G823C:p.G2 75R	NA	LP			Y	
CNSR31 0132	F	81	COL4A 1	Brain small vessel disease with or without ocular anomalies/PADMAL	Small vessel disease	NM_001845:exon3 9:c.G3379A:p.G112 7S	NA	LP	Y			
CNSR30 0117	M	75	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon3 3:c.G2954T:p.G985 V	NA	LP	Y			
CNSR30 0633	M	64	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon2 8:c.G2105C:p.G702 A	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 0646	M	65	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon3 7:c.3420delA:p.G11 40fs	NA	LP			Y	
CNSR30 0683	F	79	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon1 9:c.G1180A:p.G394 R	NA	LP		Y		
CNSR30 1611	M	53	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon3 6:c.G3338A:p.G111 3E	NA	LP			Y	
CNSR30 1656	M	57	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon4 8:c.G4909A:p.G16 37S	NA	LP	Y			
CNSR30 1698	M	60	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon1 9:c.G1180A:p.G394 R	NA	LP			Y	
CNSR30 1913	F	68	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon3 3:c.G2954T:p.G985 V	NA	LP			Y	
CNSR30 2077	M	44	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon4 8:c.G4906A:p.G16 36S	NA	LP			Y	
CNSR30 2272	F	61	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon2 0:c.1306delC:p.P43 6fs	NA	LP	Y			
CNSR30 2290	F	72	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon4 6:c.G4465A:p.G14 89S	NA	LP	Y			
CNSR30 2556	M	57	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon5: c.220delC:p.P74fs	NA	LP		Y		

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 2914	F	70	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon4 4:c.C4228T:p.R141 0	NA	P		Y		
CNSR30 3301	M	61	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon7: c.G451A:p.G151S	NA	LP			Y	
CNSR30 3811	M	68	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon3 9:c.G3589A:p.G119 7S	NA	LP	Y			
CNSR30 3908	M	47	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon2 2:c.G1525A:p.G50 9R	NA	LP			Y	
CNSR30 5035	F	75	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon2 0:c.1306delC:p.P43 6fs	NA	LP	Y			
CNSR30 5298	M	67	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon2 8:c.G2105C:p.G702 A	NA	LP			Y	
CNSR30 5789	M	59	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon3 4:c.G3088A:p.G10 30S	NA	LP			Y	
CNSR30 6020	M	52	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon1 8:c.G1078A:p.G36 0S	NA	LP			Y	
CNSR30 6307	F	76	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon1 7:c.G995A:p.G332 E	NA	LP			Y	
CNSR30 6619	M	58	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon2 5:c.G1778A:p.G59 3D	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 7166	M	72	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon1 2:c.G719A:p.G240 D	NA	LP	Y			
CNSR30 7249	M	62	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon4 2:c.G3985A:p.G13 29R	NA	LP				Y
CNSR30 7401	М	64	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon4 4:c.4269delC:p.G14 23fs	NA	LP	Y			
CNSR30 7591	F	53	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon2 0:c.1306dupC:p.G4 3	NA	LP		Y		
CNSR30 7722	M	71	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon2 0:c.C1291T:p.R431 X	NA	P	Y			
CNSR30 8082	F	67	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon3 4:c.G3071C:p.G102 4A	NA	LP		Y		
CNSR30 8217	M	70	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon2 5:c.G1792A:p.G59 8S	NA	LP	Y			
CNSR30 8251	M	65	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon3 3:c.G2954T:p.G985 V	NA	LP	Y			
CNSR30 8728	F	72	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon3 9:c.G3589A:p.G119 7S	NA	LP	Y			
CNSR30 9273	M	67	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon2 3:c.G1597T:p.G533 X	NA	P	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 9581	M	68	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon3 4:c.G3088A:p.G10 30S	NA	LP				Y
CNSR30 9645	M	59	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon2 2:c.G1525A:p.G50 9R	NA	LP		Y		
CNSR30 9799	F	52	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon4 0:c.G3649A:p.G12 17R	NA	LP			Y	
CNSR30 9962	M	62	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon7: c.G451A:p.G151S	NA	LP			Y	
CNSR31 0003	M	53	COL4A 2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon3 9:c.G3607A:p.G12 03S	NA	LP	Y			
CNSR30 1748	F	52	GSN	Amyloidosis, Finnish type	Small vessel disease	NM_000177:exon7: c.G1034A:p.W345 X	NA	P			Y	
CNSR30 3348	M	63	GSN	Amyloidosis, Finnish type	Small vessel disease	NM_000177:exon1 2:c.1736dupT:p.V5 79fs	NA	LP		Y		
CNSR30 3408	M	52	GSN	Amyloidosis, Finnish type	Small vessel disease	NM_000177:exon1 1:c.C1534T:p.Q512 X	NA	P			Y	
CNSR30 3542	M	72	GSN	Amyloidosis, Finnish type	Small vessel disease	NM_000177:exon1 3:c.1899_1909del:p .A6	NA	LP			Y	
CNSR30 3781	F	71	GSN	Amyloidosis, Finnish type	Small vessel disease	NM_000177:exon1 0:c.1408delG:p.G4 70fs	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 3822	M	50	GSN	Amyloidosis, Finnish type	Small vessel disease	NM_000177:exon1 0:c.1408delG:p.G4 70fs	NA	LP			Y	
CNSR30 4726	F	55	GSN	Amyloidosis, Finnish type	Small vessel disease	NM_000177:exon2: c.349+1G>T	NA	P			Y	
CNSR30 6001	M	71	GSN	Amyloidosis, Finnish type	Small vessel disease	NM_000177:exon1 0:c.G1349A:p.W45 0X	NA	P				Y
CNSR30 9374	M	54	GSN	Amyloidosis, Finnish type	Small vessel disease	NM_001127663:ex on3:c.85dupT:p.L2 8fs	NA	LP		Y		
CNSR30 5312	F	58	HTRA1	CARASIL	Small vessel disease	NM_002775:exon2: c.G517A:p.A173T	NA	LP	Y			
CNSR30 7080	M	59	HTRA1	CARASIL	Small vessel disease	NM_002775:exon4: c.778-2A>G	NA	P	Y			
CNSR30 9218	F	45	HTRA1	CARASIL	Small vessel disease	NM_002775:exon7: c.C1156T:p.R386X	NA	P			Y	
CNSR30 0221	F	69	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 7:c.G2687T:p.C896 F	NA	LP	Y			
CNSR30 0468	M	75	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			
CNSR30 0673	M	74	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 5:c.C2299T:p.R767 C	NA	LP		Y		
CNSR30 0768	М	43	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 0946	M	42	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon2 1:c.C3427T:p.R114 3C	NA	LP	Y			
CNSR30 1247	M	55	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 3:c.A2129G:p.Y71 0C	NA	LP	Y			
CNSR30 1511	M	37	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon2 0:c.G3313T:p.G110 5C	NA	LP	Y			
CNSR30 1757	M	46	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 8:c.C2898A:p.C966 X	NA	P	Y			
CNSR30 1888	F	61	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 6:c.G2459T:p.C820 F	NA	LP	Y			
CNSR30 1929	M	65	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1759T:p.R587 C	NA	LP	Y			
CNSR30 1933	M	61	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 3:c.C2038T:p.R680 C	NA	LP	Y			
CNSR30 2163	M	46	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 5:c.C2299T:p.R767 C	NA	LP	Y			
CNSR30 2623	M	46	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon4: c.T547A:p.C183S	NA	LP	Y			
CNSR30 2884	M	55	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 2919	M	53	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 2:c.C1918T:p.R640 C	NA	LP			Y	
CNSR30 3044	M	58	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 7:c.G2689T:p.G897 C	NA	LP	Y			
CNSR30 3149	M	58	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1759T:p.R587 C	NA	LP	Y			
CNSR30 3251	F	64	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			
CNSR30 3417	F	81	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon2 2:c.C3601T:p.R120 1C	NA	LP	Y			
CNSR30 3714	F	67	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1759T:p.R587 C	NA	LP	Y			
CNSR30 3980	F	56	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 2:c.C1918T:p.R640 C	NA	LP	Y			
CNSR30 4138	F	46	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon5: c.G798C:p.W266C	NA	LP	Y			
CNSR30 4414	F	66	NOTC H3	disease	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			
CNSR30 4488	M	54	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon3: c.G260T:p.C87F	NA	LP	Y			
CNSR30 4558	F	61	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 0:c.G1547T:p.C516 F	NA	LP	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 4990	M	45	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 4:c.C2149T:p.R717 C	NA	LP	Y			
CNSR30 5140	M	59	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon2 2:c.C3568T:p.R119 0C	NA	LP	Y			
CNSR30 5147	M	50	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 4:c.C2182T:p.R728 C	LP	VUS	Y			
CNSR30 5228	M	56	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon2 0:c.C3298T:p.R110 0C	NA	LP	Y			
CNSR30 5245	F	63	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P			Y	
CNSR30 5301	M	57	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			
CNSR30 5508	M	57	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			
CNSR30 5632	M	79	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon2 1:c.C3427T:p.R114 3C	NA	LP	Y			
CNSR30 6139	F	61	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 6:c.T2498G:p.F833 C	NA	LP	Y			
CNSR30 6600	M	52	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1759T:p.R587 C	NA	LP	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 6663	F	68	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 5:c.C2353T:p.R785 C	NA	LP	Y			
CNSR30 6806	F	56	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon6: c.889 894delinsTG	NA	P			Y	
CNSR30 6879	M	61	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			
CNSR30 6929	M	39	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			
CNSR30 6956	M	82	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 5:c.C2353T:p.R785 C	NA	LP	Y			
CNSR30 7019	F	63	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 2:c.T1931A:p.V644 D	NA	LP	Y			
CNSR30 7044	F	52	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon4: c.G671A:p.C224Y	P	VUS	Y			
CNSR30 7182	M	44	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1759T:p.R587 C	NA	LP	Y			
CNSR30 7435	M	74	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 8:c.2984delC:p.P99	NA	LP				Y
CNSR30 7912	M	65	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			
CNSR30 8185	M	61	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon4: c.C619T:p.R207C	P/LP	VUS		Y		

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 8416	F	75	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1759T:p.R587 C	NA	LP	Y			
CNSR30 8638	F	52	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon9: c.1488delC:p.P496	NA	LP			Y	
CNSR30 8656	M	65	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P		Y		
CNSR30 8967	M	63	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1819T:p.R607 C	P/LP	VUS	Y			
CNSR30 9724	M	49	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			
CNSR31 0191	M	61	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon1 1:c.C1630T:p.R544 C	P/LP	P	Y			
CNSR31 0413	M	75	NOTC H3	CADASIL	Small vessel disease	NM_000435:exon2 1:c.C3427T:p.R114 3C	NA	LP			Y	
CNSR30 0708	F	67	PRNP	Cerebral amyloid angiopathy, PRNP- related	Small vessel disease	NM_000311:exon2: c.G538A:p.V180I	P/LP	VUS			Y	
CNSR30 1128	M	76	PRNP	Cerebral amyloid angiopathy, PRNP- related	Small vessel disease	NM_000311:exon2: c.G538A:p.V180I	P/LP	VUS			Y	
CNSR30 4209	M	71	PRNP	Cerebral amyloid angiopathy, PRNP- related	Small vessel disease	NM_000311:exon2: c.G628A:p.V210I	P	VUS			Y	
CNSR30 4380	M	69	PSEN1	Alzheimer's disease	Small vessel disease	NM_000021:exon7: c.C658T:p.R220X	NA	P			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 7124	F	71	TREX1	Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL -S	Small vessel disease	NM_016381:exon1: c.1024_1041del:p.3 42_347del	P	VUS			Y	
CNSR30 8620	M	70	TREX1	Vasculopathy, retinal, with cerebral leukoencephalopathy and systemic manifestations/RVCL -S	Small vessel disease	NM_016381:exon1: c.G832A:p.A278T, TR	LP	VUS			Y	
CNSR30 0109	F	67	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon3: c.A326G:p.E109G	NA	LP			Y	
CNSR30 0873	M	69	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon4: c.C347G:p.T116R	NA	LP		Y		
CNSR30 0907	M	45	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon2: c.C170A:p.A57D	NA	LP			Y	
CNSR30 1040	M	74	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon2: c.T119C:p.V40A	NA	LP			Y	
CNSR30 1165	F	54	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon2: c.C170A:p.A57D	NA	LP			Y	
CNSR30 2353	M	78	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon2: c.C170A:p.A57D	NA	LP		Y		

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 3371	M	37	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon2: c.C170A:p.A57D	NA	LP			Y	
CNSR30 4525	M	46	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	c.G424A:p.V142I	P/LP	VUS			Y	
CNSR30 4550	F	57	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon3: c.A287G:p.K96R	NA	LP			Y	
CNSR30 5919	M	61	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon3: c.G307A:p.G103S	NA	LP				Y
CNSR30 6558	F	45	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon3: c.G241A:p.E81K	P	VUS			Y	
CNSR30 6867	M	70	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon2: c.C170A:p.A57D	NA	LP			Y	
CNSR30 7119	M	64	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon4: c.C347G:p.T116R	NA	LP			Y	
CNSR30 7312	F	64	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon2: c.T119C:p.V40A	NA	LP			Y	
CNSR30 7403	F	60	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon3: c.A287G:p.K96R	NA	LP			Y	
CNSR30 7497	M	60	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon2: c.C170A:p.A57D	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 7816	M	71	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	c.C170A:p.A57D	NA	LP			Y	
CNSR30 7956	M	58	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon2: c.C170A:p.A57D	NA	LP			Y	
CNSR30 8636	F	64	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon4: c.C419T:p.A140V	NA	LP			Y	
CNSR30 9004	M	62	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon4: c.G361A:p.G121S	NA	LP			Y	
CNSR31 0000	M	45	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon4: c.G349T:p.A117S	P/LP	VUS			Y	
CNSR31 0227	F	62	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon4: c.G361A:p.G121S	NA	LP			Y	
CNSR31 0334	M	47	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon2: c.G148A:p.V50M	P	LP			Y	
CNSR31 0356	M	69	TTR	Amyloidosis, hereditary, transthyretin-related	Small vessel disease	NM_000371:exon4: c.C413T:p.T138I	NA	LP			Y	
CNSR30 2278	F	75	ACVR L1	Telangiectasia, hereditary hemorrhagic, type 2	Other disease	NM_000020:exon7: c.C936A:p.H312Q	NA	LP				Y
CNSR30 2466	M	55	ACVR L1	Telangiectasia, hereditary hemorrhagic, type 2	Other disease	NM_000020:exon6: c.G682A:p.V228I	LP	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 4271	M	68	ACVR L1	Telangiectasia, hereditary hemorrhagic, type 2	Other disease	NM_001077401:ex on6:c.817_818delin sTG:p.L273W	NA	LP			Y	
CNSR30 0263	M	79	APOA1	Amyloidosis, 3 or more types	Other disease	NM_000039:exon3: c.127dupG:p.V43fs	NA	LP			Y	
CNSR30 4280	F	47	APOA1	Amyloidosis, 3 or more types	Other disease	NM_000039:exon3: c.116_117insTGGC :p.A39fs	NA	LP			Y	
CNSR30 2473	M	61	BMPR2	Pulmonary hypertension, primary	Other disease	NM_001204:exon2: c.77-1G>C	NA	P			Y	
CNSR30 3094	M	62	BMPR2	Pulmonary hypertension, primary	Other disease	NM_001204:exon1 2:c.G1687A:p.V56 3M	P	VUS			Y	
CNSR30 6598	M	76	BMPR2	Pulmonary hypertension, primary	Other disease	NM_001204:exon1 2:c.G1687A:p.V56 3M	P	VUS			Y	
CNSR31 0185	M	44	BMPR2	Pulmonary hypertension, primary	Other disease	NM_001204:exon1 2:c.G1687A:p.V56 3M	P	VUS			Y	
CNSR30 1648	M	64	CBL	Noonan syndrome- like disorder with or without juvenile myelomonocytic leukemia	Other disease	NM_005188:exon1 3:c.2153+1G>T	NA	P				Y
CNSR30 8533	F	65	CBL	Noonan syndrome- like disorder with or without juvenile myelomonocytic leukemia	Other disease	NM_005188:exon8: c.T1111C:p.Y371H	P	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 9022	F	51	DYRK1 B	Abdominal obesity- metabolic syndrome 3	Other disease	NM_004714:exon1 0:c.C1462T:p.R488 X	NA	P			Y	
CNSR30 9338	F	69	DYRK1 B	Abdominal obesity- metabolic syndrome 3	Other disease	NM_004714:exon3: c.149_150del:p.V5 0fs	NA	LP			Y	
CNSR30 3561	M	60	ENG	Telangiectasia, hereditary hemorrhagic, type 1	Other disease	NM_000118:exon2: c.G219A:p.T73T	P	VUS			Y	
CNSR30 4704	M	45	FLCN	Birt-Hogg-Dube syndrome	Other disease	NM_144997:exon1 2:c.1381dupA:p.S4 61fs	NA	LP			Y	
CNSR30 7128	M	72	FLCN	Birt-Hogg-Dube syndrome	Other disease	NM_144997:exon4: c.7delG:p.A3fs	NA	LP			Y	
CNSR30 9913	F	58	FLCN	Birt-Hogg-Dube syndrome	Other disease	NM_144997:exon1 1:c.1285dupC:p.H4 29fs	P	LP			Y	
CNSR30 9961	M	47	FLCN	Birt-Hogg-Dube syndrome	Other disease	NM_144997:exon1 1:c.1285dupC:p.H4 29fs	P	LP			Y	
CNSR31 0041	M	61	FLCN	Birt-Hogg-Dube syndrome	Other disease	NM_144997:exon9: c.C1015T:p.Q339X	NA	P			Y	
CNSR30 4882	F	44	GLA	Fabry Disease	Other disease	NM_000169:exon3: c.514delT:p.C172fs	NA	LP		Y		
CNSR30 6744	M	76	GLA	Fabry Disease	Other disease	c.639+919G>A	P	VUS		Y		
CNSR30 1291	F	84	KIF1B	Pheochromocytoma	Other disease	NM_015074:exon6: c.C463T:p.R155X	NA	P			Y	
CNSR30 8367	M	84	KIF1B	Pheochromocytoma	Other disease	NM_183416:exon2 1:c.1996delG:p.G6 66fs	NA	LP			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 0249	M	56	KRIT1	Cerebral cavernous malformations-1	Other disease	NM_194454:exon1 3:c.G1391A:p.W46 4X	P	P	Y			
CNSR30 2326	F	51	KRIT1	Cerebral cavernous malformations-1	Other disease	NM_194456:exon8: c.457dupA:p.T153f s	NA	LP			Y	
CNSR30 4825	F	60	KRIT1	Cerebral cavernous malformations-1	Other disease	NM_194456:exon6: c.262+1G>A	NA	P			Y	
CNSR30 5230	M	63	KRIT1	Cerebral cavernous malformations-1	Other disease	NM_194456:exon5: c.T33G:p.Y11X	NA	LP			Y	
CNSR30 9657	M	51	KRIT1	Cerebral cavernous malformations-1	Other disease	NM_194456:exon9: c.T585A:p.Y195X	NA	LP			Y	
CNSR30 5024	M	52	NF1	Neurofibromatosis 1	Other disease	NM_000267:exon4 1:c.C6349T:p.Q211 7X	NA	P		Y		
CNSR30 5340	M	53	NF1	Neurofibromatosis 1	Other disease	NM_000267:exon1 8:c.2027dupC:p.T6 76fs	P	LP		Y		
CNSR30 7410	M	73	NF1	Neurofibromatosis 1	Other disease	NM_000267:exon4: c.G479A:p.R160K	LP	VUS		Y		
CNSR30 4809	M	53	PDE4D	Acrodysostosis 2, with or without hormone resistance	Other disease	NM_001165899:ex on3:c.G94T:p.G32 X	NA	P			Y	
CNSR30 4838	F	62	PDE4D	Acrodysostosis 2, with or without hormone resistance	Other disease	NM_001197223:ex on1:c.18_21del:p.Y 6fs	NA	LP			Y	
CNSR30 0154	M	60	PKD1	ADPKD	Other disease	NM_001009944.3(PKD1):c.7065+9C> T	P	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 0265	M	53	PKD1	ADPKD	Other disease	NM_001009944:ex on10:c.1987delC:p. Q663fs	NA	LP		Y		
CNSR30 1236	M	69	PKD1	ADPKD	Other disease	NM_001009944:ex on44:c.G12036A:p. W4012X	P	P		Y		
CNSR30 1587	F	64	PKD1	ADPKD	Other disease	NM_001009944:ex on29:c.C9829T:p.R 3277C	P/LP	VUS			Y	
CNSR30 4664	F	49	PKD1	ADPKD	Other disease	NM_001009944:ex on11:c.T2534G:p.L 845W	NA	LP			Y	
CNSR30 6045	F	56	PKD1	ADPKD	Other disease	NM_001009944:ex on19:c.G7494A:p. W2498X	NA	P			Y	
CNSR30 6128	M	65	PKD1	ADPKD	Other disease	NM_001009944:ex on34:c.10499+1G> A	NA	P			Y	
CNSR30 9549	M	47	PKD1	ADPKD	Other disease	NM_001009944:ex on11:c.T2534C:p.L 845S	P/LP	VUS			Y	
CNSR30 9966	F	50	PKD1	ADPKD	Other disease	NM_001009944:ex on45:c.G12391T:p. E4131X	P	P			Y	
CNSR30 1946	M	67	PKD2	ADPKD	Other disease	NM_000297:exon3: c.779delC:p.T260fs	NA	LP			Y	
CNSR30 1999	F	54	PKD2	ADPKD	Other disease	NM_000297:exon4: c.C958T:p.R320X	P	P	Y			
CNSR30 3365	M	53	PKD2	ADPKD	Other disease	NM_000297:exon4: c.1094+1G>C	NA	P	Y			

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30 4191	F	57	PKD2	ADPKD	Other disease	NM_000297:exon1 4:c.C2533T:p.R845 X	P	P	Y			
CNSR30 8318	M	59	PKD2	ADPKD	Other disease	NM_000297:exon1 2:c.G2305T:p.E769 X	LP	P	Y			
CNSR30 8733	F	63	PKD2	ADPKD	Other disease	NM_000297:exon6: c.T1506G:p.Y502X	NA	P	Y			
CNSR30 1009	F	76	RET	Medullary thyroid carcinoma or Pheochromocytoma	Other disease	NM_020975:exon1 4:c.G2410A:p.V80 4M	P/LP	VUS			Y	
CNSR30 1675	M	62	RET	Medullary thyroid carcinoma or Pheochromocytoma	Other disease	NM_020975:exon1 4:c.G2410A:p.V80 4M	P/LP	VUS			Y	
CNSR30 3165	F	61	RET	Medullary thyroid carcinoma or Pheochromocytoma	Other disease	NM_020975:exon1 4:c.G2410A:p.V80 4M	P/LP	VUS			Y	
CNSR30 5395	M	60	RET	Medullary thyroid carcinoma or Pheochromocytoma	Other disease	NM_020975:exon1 3:c.G2370T:p.L790 F	P	VUS			Y	
CNSR30 9526	F	64	RET	Medullary thyroid carcinoma or Pheochromocytoma	Other disease	NM_020975:exon1 3:c.G2370T:p.L790 F	P	VUS			Y	
CNSR31 0343	M	68	RET	Medullary thyroid carcinoma or Pheochromocytoma	Other disease	NM_020975:exon1 4:c.G2410A:p.V80 4M	P/LP	VUS			Y	
CNSR30 3341	M	44	TGIF1	Holoprosencephaly 4	Other disease	NM_173208:exon3: c.C83T:p.S28F	NA	LP			Y	
CNSR30 8090	F	71	TGIF1	Holoprosencephaly 4	Other disease	NM_173208:exon4: c.A451G:p.T151A	P	VUS			Y	

Code_n	Ge nde r	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinvar	ACMG	Defin ite	Possi ble	Undete rmined	Insufficient Information
CNSR30	M	66	TGIF1	Holoprosencephaly 4	Other	NM_173208:exon4:	P	VUS			Y	
8126					disease	c.A451G:p.T151A						
CNSR30	M	62	VHL	Pheochromocytoma	Other	NM_000551:exon3:	NA	LP				Y
4572					disease	c.A479G:p.E160G						

eTable 4 Diagnoses of 29 individuals harbored more than 2 P/LP variants in different genes

CTADIC T	Diagno	303 01	2) IIIui	viduais nai boi cu i	more than 2 1711	variants in unierent	genes					
Code_n	Gend er	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinv ar	ACM G	Defin ite	Possi ble	Undet ermin ed	Insufficient Information
CNSR30 0094	M	70	СЕТР	Hyperalphalipoprot einemia	Large artery disease	NM_000078:exon2:c.C1 60T:p.R54X	NA	LP			Y	
CNSR30 0094	M	70	VWF	von Willebrand disease, type 1	Prothrombotic state	NM_000552:exon28:c.C 4909T:p.Q1637X	NA	P		Y		
CNSR30 0101	М	68	LDL R	Hypercholesterolem ia, familial, 1	Large artery disease	NM_000527:exon3:c.49 7delinsGGATCCCCA GCTGCATCCCCAG: p.A166GfsX48	LP	VUS		Y		
CNSR30 0101	M	68	RNF2 13	Moyamoya disease	Large artery disease	NM_001256071:exon60: c.G14429A:p.R4810K	P	VUS			Y	
CNSR30 0197	F	67	JAK2	Thrombocythemia 3	Prothrombotic state	NM_004972:exon14:c.G 1849T:p.V617F	P	VUS	Y			
CNSR30 0197	F	67	SERP INC1	Thrombophilia due to antithrombin III deficiency	Prothrombotic state	NM_000488:exon2:c.C2 35T:p.R79C	LP	VUS			Y	
CNSR30 0499	F	82	ELN	SVAS	Large artery disease	c.639+919G>A	NA	P			Y	
CNSR30 0499	F	82	GLA	Fabry Disease	Other disease	NM_000501:exon14:c. 686-2A>G	P	VUS		Y		
CNSR30 0830	M	64	KCN Q1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon3:c.T 560C:p.L187P	P/LP	VUS		Y		
CNSR30 0830	M	64	RNF 213	Moyamoya disease	Large artery disease	NM_001256071:exon6 0:c.G14429A:p.R4810 K	P	VUS			Y	
CNSR30 1010	M	55	LDL R	Hypercholesterole mia, familial, 1	Large artery disease	NM_000527:exon5:c.G 796A:p.D266N	P/LP	VUS			Y	

Code_n	Gend er	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinva r	AC MG	Defin ite	Possi ble	Undeter mined	Insufficient Information
CNSR30 1010	M	55	PDE 4D	Acrodysostosis 2, with or without hormone resistance	Other disease	NM_001197220:exon 1:c.65+1G>A	NA	P			Y	
CNSR30 1223	M	53	SCN2 B	Hereditary cardiac dysrhythm	Embolic stroke	NM_004588:exon2:c.C 142G:p.L48V	NA	LP			Y	
CNSR30 1223	M	53	LDL R	Hypercholesterole mia, familial, 1	Large artery disease	NM_000527:exon5:c.G 805A:p.G269S	P/LP	VU S	Y			
CNSR30 1496	M	47	JAK2	Thrombocythemia 3	Prothrombotic state	NM_004972:exon14:c. G1849T:p.V617F	P	VU S	Y			
CNSR30 1496	M	47	NF1	Neurofibromatosi s 1	Other disease	NM_000267:exon25:c. 3240delA:p.L1080fs	NA	LP		Y		
CNSR30 1738	F	64	FBN1	Marfan syndrome	Large artery disease	NM_000138:exon27:c. C3268G:p.P1090A	NA	LP			Y	
CNSR30 1738	F	64	RNF2 13	Moyamoya disease	Large artery disease	NM_001256071:exon6 0:c.G14429A:p.R4810 K	P	VU S		Y		
CNSR30 1797	M	77	F2	Thrombophilia due to thrombin defect	Prothrombotic state	NM_000506:exon11:c. G1303A:p.E435K	NA	LP			Y	
CNSR30 1797	M	77	VWF	von Willebrand disease, type 1	Prothrombotic state	NM_000552:exon44:c. G7450A:p.V2484I	LP	VU S			Y	
CNSR30 2006	M	46	KCN A5	Hereditary cardiac dysrhythm	Embolic stroke	NM_002234:exon1:c.C 1727T:p.A576V	P	VU S			Y	
CNSR30 2006	M	46	RNF2 13	Moyamoya disease	Large artery disease	NM_001256071:exon6 0:c.G14429A:p.R4810 K	P	VU S			Y	
CNSR30 2050	F	76	KCN Q1	Hereditary cardiac dysrhythm	Embolic stroke	NM_000218:exon6:c.G 815A:p.G272D	P	VU S	Y			
CNSR30 2050	F	76	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:exon3 26:c.C85223G:p.S2840 8X	NA	P			Y	

Code_n	Gend er	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinva r	AC MG	Defin ite	Possi ble	Undeter mined	Insufficient Information
CNSR30 2050	F	76	LDL R	Hypercholesterole mia, familial, 1	Large artery disease	NM_000527:exon13:c. G1898T:p.R633L	NA	P	Y			
CNSR30 2176	M	63	СЕТР	Hyperalphalipopr oteinemia	Large artery disease	NM_000078:exon2:c.T 222G:p.Y74X	NA	LP			Y	
CNSR30 2176	М	63	F2	Thrombophilia due to thrombin defect	Prothrombotic state	NM_000506:exon7:c.G 691A:p.G231R	NA	LP			Y	
CNSR30 3412	F	70	GJA1	Atrioventricular septal defect 3	Embolic stroke	NM_000165:exon2:c.G 158A:p.R53H	NA	LP			Y	
CNSR30 3412	F	70	KCN A5	Hereditary cardiac dysrhythm	Embolic stroke	NM_002234:exon1:c.C 1727T:p.A576V	P	VU S			Y	
CNSR30 3485	M	50	JAK2	Thrombocythemia 3	Prothrombotic state	NM_004972:exon14:c. G1849T:p.V617F	P	VU S			Y	
CNSR30 3485	М	50	DYR K1B	Abdominal obesity-metabolic syndrome 3	Other disease	NM_004714:exon8:c.C 1072T:p.R358X	NA	P			Y	
CNSR30 3619	М	65	СЕТР	Hyperalphalipopr oteinemia	Large artery disease	NM_000078:exon11:c. 1115_1127del:p.Q372f s	NA	LP			Y	
CNSR30 3619	M	65	COL 4A2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon39:c. G3589A:p.G1197S	NA	LP			Y	
CNSR30 3839	М	55	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_001267550:exon3 24:c.69145delA:p.I230 49fs	NA	LP		Y		
CNSR30 3839	M	55	SERP INC1	Thrombophilia due to antithrombin III deficiency	Prothrombotic state	NM_000488:exon3:c.T 442C:p.S148P	P	VU S			Y	
CNSR30 3839	M	55	VWF	von Willebrand disease, type 1	Prothrombotic state	NM_000552:exon14:c. 1614delC:p.P538	NA	LP			Y	
CNSR30 4342	M	53	KCN A5	Hereditary cardiac dysrhythm	Embolic stroke	NM_002234:exon1:c.C 1727T:p.A576V	P	VU S			Y	

Code_n	Gend er	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinva r	AC MG	Defin ite	Possi ble	Undeter mined	Insufficient Information
CNSR30 4342	M	53	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon46:c. C13231T:p.Q4411X	NA	LP			Y	
CNSR30 6238	F	74	СЕТР	Hyperalphalipopr oteinemia	Large artery disease	NM_000078:exon13:c. 1225_1226insAGACT: p.K409fs	NA	LP			Y	
CNSR30 6238	F	74	PRN P	Cerebral amyloid angiopathy, PRNP-related	Small vessel disease	NM_000311:exon2:c.G 538A:p.V180I	P/LP	VU S		Y		
CNSR30 6857	M	74	TTN	Hereditary cardiomyopathies	Embolic stroke	NM_133379:exon46:c. 13109delC:p.S4370X	NA	LP			Y	
CNSR30 6857	M	74	RNF2 13	Moyamoya disease	Large artery disease	NM_001256071:exon6 0:c.G14429A:p.R4810 K	P	VU S			Y	
CNSR30 6857	M	74	F2	Thrombophilia due to thrombin defect	Prothrombotic state	NM_000506:exon2:c.T 80C:p.V27A	NA	LP			Y	
CNSR30 6857	M	74	VWF	von Willebrand disease, type 1	Prothrombotic state	NM_000552:exon20:c. G2561A:p.R854Q	P/LP	VU S			Y	
CNSR30 7276	M	53	PRO C	Thrombophilia due to protein C deficiency, autos	Prothrombotic state	NM_000312:exon9:c.G 1000A:p.G334S	P	VU S			Y	
CNSR30 7276	M	53	COL 4A2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon30:c. 2577_2578insGG	NA	LP				Y
CNSR30 7448	M	64	MFA P5	Aortic aneurysm, familial thoracic 9	Large artery disease	NM_003480:exon10:c. C472T:p.R158X	P	VU S			Y	
CNSR30 7448	M	64	TGIF 1	Holoprosencephal y 4	Other disease	NM_173208:exon4:c.A 451G:p.T151A	P	VU S			Y	
CNSR30 7803	M	64	VWF	von Willebrand disease, type 1	Prothrombotic state	NM_000552:exon22:c. C2965T:p.Q989X	NA	P			Y	
CNSR30 7803	M	64	NOT CH3	CADASIL	Small vessel disease	NM_000435:exon11:c. C1759T:p.R587C	NA	LP	Y			

Code_n	Gend er	Age	Gene	Phenotype	Etiology of stroke	Mutation	Clinva r	AC MG	Defin ite	Possi ble	Undeter mined	Insufficient Information
CNSR30 8769	M	58	ELN	SVAS	Large artery disease	splicing	P	P			Y	
CNSR30 8769	M	58	LDL R	Hypercholesterole mia, familial, 1	Large artery disease	NM_000527:exon10:c. G1567A:p.V523M	P	VU S		Y		
CNSR30 8831	M	42	COL 4A2	Brain small vessel disease 2	Small vessel disease	NM_001846:exon12:c. G701T:p.G234V	NA	LP			Y	
CNSR30 8831	M	42	PKD 1	ADPKD	Other disease	NM_001009944:exon4 5:c.G12391T:p.E4131 X	P	P			Y	
CNSR30 9790	M	57	RBM 20	Hereditary cardiomyopathies	Embolic stroke	NM_001134363:exon2 :c.870delA:p.S290fs	NA	LP			Y	
CNSR30 9790	M	57	TTR	Amyloidosis, hereditary, transthyretin- related	Small vessel disease	NM_000371:exon2:c.C 170A:p.A57D	NA	LP				Y
CNSR31 0131	М	60	MYB PC3	Hereditary cardiomyopathies	Embolic stroke	NM_000256:exon12:c. 1042_1043insCGGCA: p.M348fs	P	LP			Y	
CNSR31 0131	M	60	NOT CH3	CADASIL	Small vessel disease	NM_000435:exon11:c. C1630T:p.R544C	P/LP	VU S	Y			
CNSR31 0244	M	53	GAT A4	Tetralogy of Fallot	Embolic stroke	NM_001308093:c.100 0+103G>T	P	VU S			Y	
CNSR31 0244	M	53	NF1	Neurofibromatosi s 1	Other disease	NM_000267:exon14:c. 1541_1542del:p.Q514f s	P	LP		Y		
CNSR31 0283	M	59	GJA5	Hereditary cardiac dysrhythm	Embolic stroke	NM_005266:exon2:c.C 292T:p.H98Y	NA	LP		Y		
CNSR31 0283	M	59	NOT CH3	CADASIL	Small vessel disease	NM_000435:exon3:c.C 328T:p.R110C	P	VU S	Y			

eTable 5 Four individuals with 2 P/LP variants in the ABCC6 gene

Code	Gender	Age	Gene	Mutation	Clinvar	ACMG	Phenotype
CNSR309580	F	66	ABCC6	NM_001171: exon26: c.C3703T: p.R1235W	P	VUS	Pseudoxanthoma elasticum
CNSR301384	M	53	ABCC6	NM_001171: exon31: c.4404-1G>A	NA	P	Pseudoxanthoma elasticum
CNSR301384	M	53	ABCC6	NM_001171: exon24: c.C3490T: p.R1164X	P	P	Pseudoxanthoma elasticum
CNSR309564	M	37	ABCC6	NM_001171: exon26: c.C3703T: p.R1235W	P	VUS	Pseudoxanthoma elasticum
CNSR309564	M	37	ABCC6	NM_001171: exon28: c.G3892A: p.V1298I	NA	LP	Pseudoxanthoma elasticum
CNSR310336	M	48	ABCC6	NM_001171: exon26: c.3735+1G>T	NA	P	Pseudoxanthoma elasticum
CNSR310336	M	48	ABCC6	NM_001171: exon23: c.C3304T: p.Q1102X	NA	P	Pseudoxanthoma elasticum

Supplemental material

Characteristic	ToTal (N=759)	Embolic stroke (N=245, 32.28)	Large artery disease (N=184, 24.24)	Prothrombotic state (N=124, 16.34)	Small vessel disease (N=148, 19.50)	Other disease (N=58, 7.64)
Age at time of study entry, mean±SD	61.53±11.50	61.91±12.23	60.43±12.22	62.78±10.54	61.52±10.59	60.71±10.05
≤45 yr	66 (8.70)	21 (8.57)	21 (11.41)	8 (6.45)	13 (8.78)	4 (6.90)
> 45 yr	693 (91.30)	226 (92.24)	163 (88.59)	116 (93.55)	135 (91.22)	54 (93.10)
Male, n (%)	504 (66)	162 (66.12)	119 (64.67)	89 (71.77)	98 (66.22)	36 (62.07)
Medical history, n (%)						
Ischaemic stroke	193 (25.43)	56 (22.86)	42 (22.83)	30 (24.19)	47 (31.76)	18 (31.03)
Coronary heart diseases	98 (12.91)	36 (14.69)	15 (8.15)	17 (13.71)	22 (14.86)	8 (13.79)
Atrial fibrillation	64 (8.43)	25 (10.20)	14 (7.61)	10 (8.06)	9 (6.08)	6 (10.35)
Hypertension	452 (59.55)	139 (56.73)	119 (64.67)	71 (57.26)	81 (54.73)	42 (72.41)
Diabetes mellitus	177 (23.32)	57 (23.27)	45 (24.46)	29 (23.39)	28 (18.92)	18 (31.03)
Dyslipidemia	68 (8.96)	26 (10.61)	10 (5.43)	8 (6.45)	18 (12.12)	6 (10.35)
Stroke type						
IS	718 (94.60)	233 (95.10)	171 (92.93)	119 (95.97)	142 (95.95)	53 (91.38)
TIA	41 (5.40)	12 (4.90)	13 (7.07)	5 (4.03)	6 (4.05)	5 (8.62)
Family history of Stroke	97 (12.78)	38 (15.51)	24 (13.04)	13 (10.48)	15 (10.14)	7 (12.07)
CCS						
Large artery atherosclerosis	220 (28.99)	66 (26.94)	70 (38.04)	33 (26.61)	40 (27.03)	11 (18.97)
Cardioaortic embolism	57 (7.51)	27 (11.02)	10 (5.43)	6 (4.84)	8 (5.41)	6 (10.35)
Small arterial occlusion	191 (25.16)	66 (26.94)	36 (19.57)	27 (21.77)	43 (29.05)	19 (32.76)
Other etiologies	11 (1.45)	1 (0.41)	6 (3.26)	1 (0.81)	3 (2.03)	0 (0.00)
Undetermined etiology	280 (36.89)	85 (34.69)	62 (33.70)	57 (45.97)	54 (36.49)	22 (37.93)

Supplemental material

Code_n	Mendelian caused of stroke through EHR reviewed	Gene	Age	Gender
CNSR301140	Thrombocythemia	JAK2;exon11:c.G1402T:p.V468F,	67	Female
CNSR301496	Thrombocythemia	JAK2;exon11:c.G1402T:p.V468F,	47	Male
CNSR302832	Thrombocythemia	JAK2;exon11:c.G1402T:p.V468F,	63	Male
CNSR307039	Thrombocythemia	JAK2;exon11:c.G1402T:p.V468F,	66	Male
CNSR307566	Thrombocythemia	JAK2;exon11:c.G1402T:p.V468F,	62	Female
CNSR309872	Thrombocythemia	JAK2;exon11:c.G1402T:p.V468F,	75	Male
CNSR303251	Small vessel disease, CADASIL	NOTCH3;exon11:c.C1630T:p.R544C	64	Female
CNSR305508	Small vessel disease, CADASIL	NOTCH3;exon11:c.C1630T:p.R544C	57	Male
CNSR306929	Small vessel disease, CADASIL	NOTCH3;exon11:c.C1630T:p.R544C	39	Male
CNSR307912	Small vessel disease, CADASIL	NOTCH3;exon11:c.C1630T:p.R544C	65	Male
CNSR308967	Small vessel disease, CADASIL	NOTCH3;exon11:c.C1819T:p.R607C	63	Male
CNSR301757	Small vessel disease, CADASIL	NOTCH3;exon18:c.C2898A:p.C966X	46	Male
CNSR310283	Small vessel disease, CADASIL	NOTCH3;exon3:c.C328T:p.R110C	59	Male
CNSR307044	Small vessel disease, CADASIL	NOTCH3;exon4:c.G671A:p.C224Y	52	Female
CNSR302455	Moyamoya	RNF213;exon60:c.G14429A:p.R4810K	27	Female
CNSR304840	Moyamoya,PFO?	RNF213;exon60:c.G14429A:p.R4810K	30	Male
CNSR306251	Moyamoya	RNF213;exon60:c.G14429A:p.R4810K	48	Female