



ISGC NEWSLETTER

ISSUE 26 - DECEMBER 2013

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If you would like to suggest a topic or article for a future newsletter, or to give general feedback, please contact:

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(battey@chgr.mgh.harvard.edu).

Introduction

This is the twenty-sixth issue of the bi-monthly newsletter for the International Stroke Genetics Consortium. The ISGC newsletter will serve to keep investigators updated on ongoing projects, new project proposals, meetings, and publications.

The newsletter will be posted as a PDF document to the ISGC website on the 1st of even months (or nearest workday) and can either be viewed online or downloaded as an attachment.

All Investigators are invited and encouraged to submit content for the newsletter. Please send content suggestions to Thomas Battey (battey@chgr.mgh.harvard.edu).

ISGC Founding Principles

Cerebrovascular disease is a complex disorder influenced by variation in many genetic and non-genetic exposures, each of which contributes only a small influence to disease risk. Therefore large (larger than any single center can assemble on its own) well-characterized samples will be necessary to discover these exposures.

Principles of Collaboration:

- 1) The ISGC is open to all who can contribute
- 2) All contributions will be fairly recognized in publications
- 3) We work together in a spirit of cooperation and open communication in order to promote the best science in the present and the best science in the future.

ISGC Project Updates

New Projects:

There are no new projects at this time.

Ongoing Projets:

Genetics on post stroke functional outcome: Genome wide association study (GWAS) on post ischemic stroke functional outcome.

Contact: Jane Maguire, Arne Lindgren, Christina Jern and Brad Worrall

Date Proposed: February 2013

Aims: (A) To conduct a GWAS to identify genetic variants associated with functional outcome after ischemic stroke;

(B) To replicate the preliminary GWAS results from the discovery cohort.

Status: Study acronym decided: GISCOME Genetics of Ischaemic Stroke Functional Outcome Study. GWAs of Discovery cohort completed. QC ongoing. Replication cohorts contacted, but more are welcome. Primary results expected end of 2013. Grants applications submitted in Australia and Sweden for replication costs. One grant received in Sweden. Contacted SiGN autumn 2013 and asked for permission to use GWAS and CCS data from own sites. Inventory of available phenotype variables at GISCOME sites ongoing.

NIH-funded ICH GWAS

Contact: Jonathan Rosand

Date Proposed: January 2008

Aims: This multi-center genome-wide association study (GWAS) is designed to identify genetic determinants of:

- 1) Risk of intracerebral hemorrhage (ICH) using a case-control design
- 2) Clinical course of ICH using a cohort design of individuals who have suffered an ICH.

Status: Preliminary results of ongoing meta-analyses presented in Charlottesville at the ISGC Workshop in April. Current work is directed at phenotype harmonization between ISGC European and North American samples with those enrolled in Pakistan.

Looking for genetic risk factors of cervical artery dissections

Contact: Stéphanie Debette and Didier Leys, on behalf of the CADISP group

Date Proposed: February 2009

Aims: A replication study to test whether the polymorphisms associated with CAD in the GWAS within the CADISP-consortium are also associated with CAD in other independent populations, in order to exclude spurious associations.

Status: The CADISP manuscript is in the revision process.

ISGC Project Updates

Ongoing Projects (continued):

The International Stroke Genetics Consortium Informatics Platform: A tool for Efficient Collaboration and Rapid Discovery

Contacts: Steve Bevan, Jonathan Rosand

Date Proposed: February 2009

Aims: 1) Compile a publically-available web-based catalog of all clinical characteristics, radiographic studies, genetic data and available biological samples collected for subjects with ischemic stroke and controls.

2) Enrich characterization of stroke patients by classifying all subjects according to the biologically-based Causative Classification of Stroke System and creating a central repository of de-identified neuroimaging data on patients with stroke.

Status: Imaging repository function will be a part of the NINDS-ISGC collaborative U01 grant. Phenotypic and genotypic functionality to be added as a part of a BioInformatics Research Network ARRA grant.

National Institute of Neurological Disorders and Stroke Ischemic Stroke GWAS

Contacts: Steven Kittner

Date Proposed: March 2009

Aims: Funding of an ischemic stroke genetics consortium in order to perform a genome wide association study in ischemic stroke patients and matched controls.

Status: Grant awarded. Project underway.

White Matter Hyperintensity GWAS

Contacts: Natalia Rost, Jonathan Rosand

Date Proposed: July 2009

Aims: To discover genetic markers of WMH severity using available genome-wide data and WMH volumes measured on MRI in patients with acute ischemic stroke.

Status: (1) interim GWA summary results to be exchanged between MGH/WTCCC to facilitate future meta-analysis; (2) WMH heritability study completed and pending validation; (3) WMH genetic score previously shown to be associated with incident ischemic stroke risk is pending replication.

A multi-center NIH R01 grant application, The MRI-GENetics Interface Exploration (MRI-GENIE) study has been submitted in June 2013 to further explore the genetic determinants of WMH using ischemic stroke cases (n=3,385) from the NINDS Stroke Genetics Network (SiGN).

ISGC Project Updates

Ongoing Projects (continued):

MetaStroke: A meta-analysis of genome-wide studies in ischaemic stroke

Contact: Hugh Markus

Date Proposed: December 2009

Aims: Meta-Analysis of genome-wide data in ischemic stroke

Status: Meta-analysis underway. Additional project proposal for replication cohorts sent out to ISGC in September.

Genome-wide association study of deep cerebral phenotypes

Contact: Dan Woo

Date Proposed: February 2010

Aims: (1) Specific SNPs/gene regions will be associated with deep cerebral phenotypes (deep/periventricular location of ICH, lacunar stroke or white matter disease) and this association will be independent of traditional risk factors including hypertension. (2) Specific SNPs/gene regions will be associated with deep cerebral phenotypes and will be modified by the presence/duration/severity of hypertension.

Status: List of interested studies compiled and in the process of developing a data transfer agreement. In addition, we are awaiting the completion of the Wellcome Trust effort as it would constitute a major contribution of small vessel ischemic strokes.

Genetics of cerebral venous thrombosis (CVT)

Contact: Pankaj Sharma

Date Proposed: May 2010

Aims: To recruit DNA from CVT patients in order to undertake a GWAS.

Status: Exome genotyping of the first 500 cases is now complete

Replication of associations detected in a the Meta-stroke meta-analysis of genome-wide studies in ischaemic stroke

Contact: Hugh Markus

Date Proposed: September 2010

Aims: Replication of findings from the initial MetaStroke collaboration in novel cohorts of Caucasian and other ethnic groups.

Status: Analysis in progress

ISGC Project Updates

Ongoing Projects (continued):

Consortium of Minority Population genomewide-Association Studies of Stroke (COMPASS)

Contact: Brad Worrall

Date Proposed: December 2010

Aims: Meta-analysis of cohorts and case-control studies with GWAS data for individuals of African descent and other minority groups.

Status: No update at this time.

Replication of Ischemic Stroke Genes Discovered from Exome Sequencing

Contact: Steve Rich

Date Proposed: March 2011

Aims: Replication of genes discovered in NHLBI Exome Sequencing Project in ischemic stroke cases with small or large vessel strokes.

Status: The examination of rare variants in ischemic stroke in collaboration with the NHLBI Exome Sequencing Project (ESP) has made significant progress. The ESP Executive Committee has approved the use of 3,000 case samples and 3,000 control samples to be genotyped with the ExomeChip, a custom 300,000 SNP array (as designed by Illumina) with rare variants residing in exomes and additional content. These samples will be split with WHI and the ISGC, and restricted to those samples with existing GWAS data (for imputation), deep phenotyping (for use with other targeted traits), and both cases and controls from the same sites, with IRB approval for extensive sharing of information and deposition of data into dbGaP. The two studies that had samples meeting these criteria and immediately available for shipment to the University of Washington (the ESP genotyping site for ischemic stroke) were GEOS and ISGS. These samples are being sent for evaluation with anticipated completion of genotyping in Q1 2012.

Genetic studies of recurrent stroke

Contact: Brad Worrall, Michele Sale, Keith Keene

Date Proposed: February 2011

Aims: Meta-analysis and replication of GWAS in recurrent ischemic stroke.

Status: No update at this time.

ISGC Project Updates

Ongoing Projects (continued):

Next Generation Sequencing in Lacunar Stroke and Small Vessel Disease

Contact: Anna Bersano

Date Proposed: April 2011

Aims: Perform Next-Generation Sequencing on subjects with lacunar stroke and small vessel disease. Novel mutations will be replicated via direct genotyping in an additional 1000 cases and 1000 controls .

Status: No update at this time.

GWA meta-analysis of carotid plaque and intima-media thickness (IMT)

Contact: Pankaj Sharma

Date Proposed: June 2011

Aims: Meta-analysis of existing GWA data in those subjects who have had carotid imaging in order to use carotid disease as a surrogate for vascular disease.

Status: Individual Investigators are being approached to determine the extent of phenotyping available.

Genome-wide Heritability of Ischemic Stroke in Caucasians.

Contact: Braxton Mitchell

Date Proposed: August 2011

Aims: Examine the heritability of ischemic stroke using genome wide SNPs for early vs. late onset IS and determining stroke subtype

Status: No update at this time.

GWAS of Stroke/TIA in Patients with Atrial Fibrillation

Contact: Mina Chung, Sudha Seshadri

Date Proposed: October 2011

Aims: Perform a GWAS meta-analysis of stroke/TIA within AF cases

Status: No update at this time.

Genome Wide Association Study of Plasma Fibrinogen

Contact: Christopher O'Donnell

Date Proposed: November 2011

Aims: To conduct a meta-analysis of GWA studies on plasma levels of fibrinogen with the goal of identifying novel loci that underlie variation in plasma fibrinogen concentration.

Status: No update at this time.

ISGC Project Updates

Ongoing Projects (continued):

Pharmacogenomics GWAS of tPA-induced Haemorrhagic Transformation

Contact: Christopher Levi and Jane Maguire

Date Proposed: November 2011

Aims: To identify SNPs associated with haemorrhagic transformation of acute ischemic stroke after intravenous tPA therapy using a pharmacogenomic-focused GWAS.

Status: Although unfunded, we are currently recruiting from two Australian sites, and collaborating with ISGC groups for future pooling of further samples

Genome-wide association study of incident stroke-wave 2

Contact: Stéphanie Debette, Will Longstreth, and Sudha Seshadri on behalf of the CHARGE Consortium

Date Proposed: June 2012

Aims: Our aim is to perform a second wave Incident Stroke GWAS meta-analysis within the CHARGE consortium, including a larger number of cohorts, using 1000G imputation, and including extension to other ethnic groups if large enough samples can be collected.

Status: Meta-analysis for GWAS of all stroke, ischemic stroke, cardioembolic ischemic stroke, non-cardioembolic ischemic stroke was presented at the Lund meeting. 14 prospective longitudinal cohort studies included in analysis, 15th cohort will be added shortly for discovery phase. Replication will be sought immediately after.

GRECAS Project: Genotyping Risk and Efficacy of Clopidogrel or Aspirin following Stroke

Contact: Israel Fernandez-Cadenas and Joan Montaner

Date Proposed: September 2012

Aims: Replication Study. This is a pharmacogenomic study, the main objective is: to find genetic risk factors associated with Aspirin or Clopidogrel clinical resistance, considering clinical resistance as new vascular recurrences during a follow up of one year.

Status: The first plates with the samples for the replication stage have been sent to the genotyping center.

ISGC Project Updates

Ongoing Projects (continued):

International Stroke Genetics Consortium Project Protocols and Standards

Contact: Jennifer Majerisk and Jane Maguire

Date Proposed: January 2013

Aims: The goal of this paper is to describe clinical and research criteria agreed upon by the ISGC to standardize data collection of stroke cases & controls across continents. Criteria will include definitions and requirements for harmonization of phenotypes, neuroimaging, genotyping definitions and requirements, and ethics. (There will be a separate paper by Valerie Valant and others on how to biobank.) It is anticipated that in addition to providing ISGC members with agreed clinical and research parameters, this paper will aid and encourage investigators in under-represented countries & continents to contribute their cohorts to stroke genetic research.

Status: The group is working on common phenotyping definitions at this time.

The ISGC: Establishment of a biorepository for stroke genetic research

Contact: Valerie Valant, Thomas Battey and Jonathan Rosand

Date Proposed: February 2013

Aims: To describe how one academic health center and an international consortium of researchers collaborated to organize and implement a stroke genetic research biorepository. This paper will not only provide details on how to set-up a research biorepository, but will also help under-represented countries & continents to contribute their cohorts to stroke genetic research. (There will be a separate paper by Jenny Majersik and Jane Maguire on ISGC phenotyping protocols and standards).

Status: Manuscript has been drafted and will be circulated to co-authors shortly.

ISGC Project Updates

Ongoing Projects (continued):

A Joint meta-analysis of gene x aspirin interactions in ischemic stroke to identify variants related to aspirin resistance

Contact: Christopher John Oldmeadow and John Attia

Date Proposed: February 2013

Aims: Discovery study to confirm loci involved in gene x aspirin interaction in stroke patients . We performed a genome-wide gene x environment-interaction study on a predominantly elderly European cohort of stroke cases with self-reported medication history and found evidence of a locus on chromosome 7p32 (3 SNPs). We propose to meta-analyse this region (consisting of 4 SNPs) using summary statistics from participating groups using the Joint Meta Analysis (JMA) method (described in Manning et al, Genetic Epidemiology 2011).

Status: Seeking ISGC Collaborators

Genetics on post stroke functional outcome: Genome wide association study (GWAS) on post ischemic stroke functional outcome

Contact: Jane Maguire, Arne Lindgren, Christina Jern and Brad Worrall

Date Proposed: February 2013

Aims: (A) To conduct a GWAS to identify genetic variants associated with functional outcome after ischemic stroke; (B) To replicate the preliminary GWAS results from the discovery cohort

Status: GWAs of Discovery cohort in progress. Replication cohorts contacted, but more are welcome. Primary results expected end of 2013. Grants applications submitted in Australia and Sweden for replication costs.

ISGC Project Updates

Completed Projects:

A genome-wide association study of early onset ischemic stroke

Contact: Braxton Mitchell, Steven Kittner

Date Proposed: January 2008

Aims: To carry out a GWAS of early onset stroke

- 1) Conduct a staged GWAS in the U of Maryland sample
- 2) Replicate associations detected in Aim 1 in an independent set of young-onset stroke cases controls from collaborators in the ISGC.
- 3) Determine if SNPs robustly associated with early onset stroke in both the Maryland and IGSC cohorts are also associated with older onset stroke.

Status: Manuscript in press at *G3: Genes, Genomes, Genetics*

*Cheng Y-C, O'Connell JR, Cole, JW, Stine OC, Dueker N, McArdle PF, Sparks MJ, Shen J, Laurie CC, Nelson S, Doheny KF, Ling H, Pugh EW, Bott TG, Brown Jr. RD, Meschia JF, Nalls M, Rich SS, Worrall B, Andreson CD, Biffi A, Cortellini L, Furie KL, Rost NS, Rosand J, Manolio TA, Kittner SJ, Mitchell BD. Genome-wide association analysis of ischemic stroke in young adults. *G3: Genes, Genomes, Genetics*. 2011 Nov 1; 1(6):505-514.

International Validation of a Computerized Algorithm for Etiologic Classification of Ischemic Stroke: The Causative Classification of Stroke System (CCS)

Contact: Hakan Ay, Jonathan Rosand

Date Proposed: March 2008

Aims: This is an ISGC-wide study to validate a computerized system for etiologic classification of ischemic stroke.

Status: Manuscript published.

*Arsava EM, Ballabio E, Benner T et. al.; on behalf of the International Stroke Genetics Consortium. The Causative Classification of Stroke system: An international reliability and optimization study. *Neurology*. 2010 Oct 5;75(14):1277-1284.

Replication of Chr. 9q21 region in stroke cases and matched controls in Chinese population

Contacts: Xingyu Wang, Lisheng Liu

Date Proposed: March 2008

Aims: To replicate the findings of the Chromosome 9p21 projects of the ISGC within a Chinese cohort.

Status: The project has been stalled due to a lack of sample collection.

ISGC Project Updates

Completed Projects (continued):

Australian GWAs in ischaemic stroke

Contact: Christopher Levi, John Attia, Jane Maguire and Liz Holliday

Date Proposed: May 2008

Aims: To identify snps associated with acute ischaemic stroke

Status: We demonstrated that the most common mechanistic form of ischaemic stroke, large artery atherosclerosis (LAA) is influenced by a genetic component. We identified a new LAA susceptibility locus on chromosome 6p21.1. We then replicated this susceptibility locus in 1,715 LAA cases and 52,695 population controls from 10 independent population cohorts. We are currently participating in discovery and replication cohorts with various ISGC members. Manuscript published.

Holliday EG, Maguire JM, Evans TJ, et al, on behalf of the International Stroke Genetics Consortium. Common variants at 6p21.1 are associated with large artery atherosclerotic stroke. *Nat Genet.* 2012;44(10):1147-51.

Relationship of genetic markers for common risk factors for stroke with ischemic cerebrovascular disease

Contact: Vincent Thijs

Date Proposed: June 2008

Aims: Determine whether SNPs associated with well known risk factors for ischemic stroke like diabetes, elevated LDL, myocardial infarction and atrial fibrillation are associated with ischemic cerebrovascular disease using a case control design.

Status: Manuscript published.

*Lemmens R, Buysschaert I, Geelen V, et.al. International Stroke Genetics Consortium. The Association of the 4q25 Susceptibility Variant for Atrial Fibrillation With Stroke Is Limited to Stroke of Cardioembolic Etiology. *Stroke.* 2010;41(9):1850-7.

ISGC Project Updates

Completed Projects (continued):

Chromosome 12 and risk of ischemic stroke: A replication study

Contacts: James Meschia, Andrew Singleton, Jonathan Rosand

Date Proposed: April 2009

Aims: Replication effort through the ISGC of the CHARGE discovery of two SNPs on chromosome 12 that were over-represented among cases with ischemic stroke, compared to controls.

Status: Manuscript published.

*International Stroke Genetics Consortium; Wellcome Trust Case-Control Consortium 2. Failure to validate association between 12p13 variants and ischemic stroke. *New England Journal of Medicine*. 2010;362(16):1547-1550.

Are established candidate gene polymorphisms for blood pressure, coronary heart disease, atrial fibrillation, lipid metabolism and hemostatic and inflammatory pathways also related to ischemic stroke risk in populations from the Southwest of Sweden?

Contacts: Arne Lindgren, Christina Jern, Olle Melander

Date Proposed: July 2009

Aims: To examine if SNPs related to phenotypes are related to ischemic stroke risk in a homogenous population sample from the Southwest of Sweden.

Status: Manuscript published.

*Olsson S, Melander O, Jood K, Smith JG, Lökvist H, Sjögren M, Engström G, Norrving B, Lindgren A, Jern C, the International Stroke Genetics Consortium (ISGC). A genetic variant on chromosome 12p13 does not show association to ischemic stroke in three Swedish case-control studies. *Stroke*. 2010; 42(1):214-6.

ISGC Project Updates

Completed Projects (continued):

Genes and Response to Aspirin in Secondary Stroke Prevention, GRASSP

Contact: Agnieszka Slowik, Joanna Pera

Date Proposed: June 2010

Aims: To establish genetic markers of aspirin efficiency, aspirin resistance, aspirin intolerance, and increased risk aspirin-related adverse effects in ischemic stroke patients with different stroke etiologies. To develop clinically useful and cost-effective test(s) allowing predict responses to aspirin treatment, and to avoid/reduce adverse effects .

Status: Grant submitted, project not funded at this time.

Association of myocardial infarction-associated SNPs with ischemic stroke: a meta-analysis of European Caucasian populations

Contact: Braxton Mitchell, Yu-Ching Cheng

Date Proposed: July 2010

Aims: The goal of this project is to extend previous work (e.g., the ISGC analysis of the chr 9 SNP on stroke) to determine if: (1) other MI-associated SNPs are associated with ischemic stroke; and (2) if associations of these additional SNPs are dependent on stroke subtype and/or age of stroke onset.

Status: Manuscript published.

Cheng YC, Anderson CD, Bione S, et al, on behalf of the International Stroke Genetics Consortium. Are myocardial infarction—associated single-nucleotide polymorphisms associated with ischemic stroke? *Stroke*. 2012;43(4):980-986.

Other Projects Involving ISGC Members

Genetic and Environmental Risk Factors for Hemorrhagic Stroke

Contact: Daniel Woo

Australian Stroke Genetics Collaborative Group

Contact: Chris Levi, John Attia

NHLBI Initiative on White Matter Disease

Contact: Paul Nyquist

INTERSTROKE

Contact: Guillaume Pare

Mitochondrial Genetics and Risk of Stroke

Contact: Jonathan Rosand

Status: Manuscript published

METASTROKE

Contact: Martin Dichgans

ISGC Grants Awarded

- Wellcome Trust Genome-Wide Association Study for Ischemic Stroke (WTCCC2)
- Australian Stroke Genetics Collaborative Group
- The Baltimore-Washington Young Stroke Study (GEI)
- Gene Discovery for Warfarin-Related Intracerebral Hemorrhage (ICH GWAS)
- Ethnic/Racial Variations of Intracerebral Hemorrhage (ERICH)
- NINDS-Stroke Genetics Network (SiGN) Study

ISGC Publications (2009-2011)

2009:

Gschwendtner A, Bevan S, Cole JW, et. al.; on behalf of the International Stroke Genetics Consortium. Sequence variants on chromosome 9p21.3 confer risk for atherosclerotic stroke. *Ann Neurol.* 2009;65(5):531-9.

2010:

International Stroke Genetics Consortium; Wellcome Trust Case-Control Consortium 2. Failure to validate association between 12p13 variants and ischemic stroke. *N Engl J Med.* 2010;362(16):1547-50.

Lemmens R, Buysschaert I, Geelen V, et. al.; on behalf of the International Stroke Genetics Consortium. The association of the 4q25 susceptibility variant for atrial fibrillation with stroke is limited to stroke of cardioembolic etiology. *Stroke.* 2010;41(9):1850-7.

Arsava EM, Ballabio E, Benner T, et. al.; on behalf of the International Stroke Genetics Consortium. The Causative Classification of Stroke system: An international reliability and optimization study. *Neurology.* 2010; 75(14):1277-1284.

Biffi A, Sonni A, Anderson CD, et al.; on behalf of the International Stroke Genetics Consortium. Variants at APOE Influence Risk of Deep and Lobar Intracerebral Hemorrhage. *Ann Neurol.* 2010; 68(6):934-43.

2011:

Anderson CD, Biffi A, Rahman R, et. al, on behalf of the International Stroke Genetics Consortium. Common mitochondrial sequence variants in ischemic stroke. *Ann Neurol.* 2011; 69(3):471-80.

Olsson S, Melander O, Jood K, Smith JG, Lövkvist H, Sjögren M, Engström G, Norrving B, Lindgren A, Jern C, the International Stroke Genetics Consortium (ISGC). A genetic variant on chromosome 12p13 does not show association to ischemic stroke in three Swedish case-control studies. *Stroke.* 2011; 42(1):214-6.

Biffi A, Anderson CD, Jagiella JM, et.al., on behalf of the International Stroke Genetics Consortium. APOE genotype and extent of bleeding and outcome in lobar intracerebral haemorrhage: a genetic association study. *Lancet Neurology.* 2011;10(8):702-709.

ISGC Publications (2012-2013)

2012:

International Stroke Genetics Consortium (ISGC); Wellcome Trust Case Control Consortium 2 (WTCCC2), Bellenguez C, et al. Genome-wide association study identifies a variant in HDAC9 associated with large vessel ischemic stroke. *Nature Genetics*. 2012;44(3):328-333.

Cheng YC, Anderson CD, Bione S, et al, on behalf of the International Stroke Genetics Consortium. Are myocardial infarction—associated single-nucleotide polymorphisms associated with ischemic stroke? *Stroke*. 2012;43(4):980-986.

Falcone GJ, Biffi A, Devan WJ, et al, on behalf of the International Stroke Genetics Consortium. Burden of Risk Alleles for Hypertension Increases Risk of Intracerebral Hemorrhage. *Stroke*. 2012;43(11):2877-2883.

Holliday EG, Maguire JM, Evans TJ, et al, on behalf of the International Stroke Genetics Consortium. Common variants at 6p21.1 are associated with large artery atherosclerotic stroke. *Nat Genet*. 2012;44(10):1147-51.

Traylor M, Farrall M, Holliday EG, et al, on behalf of the International Stroke Genetics Consortium. Genetic risk factors for ischaemic stroke and its subtypes (the METASTROKE Collaboration): a meta-analysis of genome-wide association studies. *Lancet Neurol*. 2012;11(11):951-962.

2013:

Williams FM, Carter AM, Hysi PG, et al, on behalf of the International Stroke Genetics Consortium. Ischemic stroke is associated with the ABO locus: The EuroCLOT study. *Ann Neurol*. 2013;73(1):16-31.

Anderson CD, Biffi A, Nalls MA, et al, on behalf of the International Stroke Genetics Consortium. Common variants within oxidative phosphorylation genes influence risk of ischemic stroke and intracerebral hemorrhage. *Stroke*. 2013;44(3):612-9.

Falcone GJ, Biffi A, Devan WJ, et al, on behalf on the GOCHA investigators. Burden of Blood Pressure-Related Alleles is Associated with Larger Hematoma Volume and Worse Outcome in Intracerebral Hemorrhage. *Stroke*. 2013. In press.

Devan WJ, Falcone GJ, Anderson CD, on behalf of the International Stroke Genetics Consortium. Heritability Estimates Identify a Substantial Genetic Contribution to Risk and Outcome of Intracerebral Hemorrhage. *Stroke*. 2013. In press.

ISGC Publications (2013-Present)

2013 (Cont.):

Adib-Samii P, Rost N, Traylor M, et al. on behalf of the International Stroke Genetics Consortium. 17q25 Locus is associated with white matter hyperintensity volume in ischemic stroke, but not with lacunar stroke status. *Stroke*. 2013; 44(6):1609-1615.

Yadav S, Cotlarciuc I, Munroe PB, et al. on behalf of the International Stroke Genetics Consortium. Genome-wide analysis of blood pressure variability and ischemic stroke. *Stroke*. 2013; 44(10):2703-2709.

Biffi A, Anderson, CD, Falcone GJ, et al. on behalf of the International Stroke Genetics Consortium. Novel insights into the genetics of intracerebral hemorrhage. *Stroke*. 2013; 44(6 Suppl 1):S137.

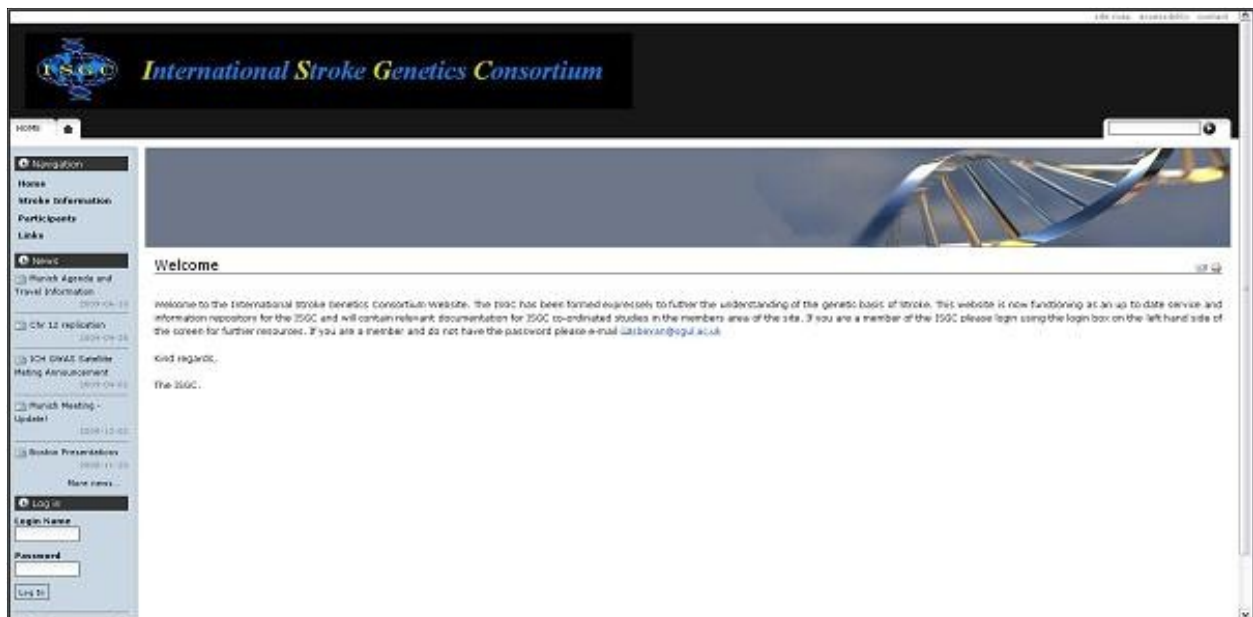
Dichgans M, Malik R, König IR, et al. on behalf of the METASTROKE Consortium; the CARDIoGRAM consortium; the C4D consortium; the International Stroke Genetics Consortium. Shared Genetic Susceptibility to Ischemic Stroke and Coronary Artery Disease: A Genome-Wide Analysis of Common Variants. *Stroke*. In press.

Website Update

The ISGC website can be found at www.strokegenetics.org. Please consider submitting a short bio and a picture for the website! If you are interested, please email them to Steve.

If you have additional website content or layout suggestions, please email Steve Bevan (snb31@medschl.cam.ac.uk) with your ideas.

Thank you!



Upcoming Conference Deadlines

2014

International Stroke Conference

Abstract Deadline: Passed

Meeting Date: February 12-14, 2014

Meeting Location: San Diego, CA

International Stroke Genetics Consortium Meeting

Meeting Date: April 3-4, 2014

Meeting Location: Salt Lake City, UT (USA)

American Academy of Neurology

Abstract Deadline: Passed

Meeting Date: April 26 - May 3, 2014

Meeting Location: Philadelphia, PA

European Stroke Conference

Abstract Deadline: January 12, 2014

Meeting Date: May 4-9, 2014

Meeting Location: Nice, France

American Neurological Association

Call for Abstracts: January 2014

Meeting Date: October 12-14, 2014

Meeting Location: Baltimore, MD

American Society of Human Genetics

Abstract Deadline: June 4, 2014

Meeting Date: October 18-22, 2014

Meeting Location: San Diego, CA

14th International Workshop: Lund, Sweden

Thank you to Arne Lindgren, Jennifer Majersik, and Christina Jern for putting on a successful ISGC Workshop in Lund!

The 14th International Workshop of the ISGC was held November 14-15, 2013 in Lund, Sweden.



International Stroke Genetics Consortium



15th International Workshop: Salt Lake City, Utah, USA

The 15th International Workshop of the ISGC will take place on April 3-4, 2014 in Salt Lake City, Utah, USA and will be hosted by Jennifer Majersik (Jennifer.Majersik@hsc.utah.edu) and co-chairs to be determined.

More detailed information regarding event programs and travel arrangements, as well as other items, will become available as they are developed.

Stay tuned for a conference website and other information!

