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## Readme file for the International Stroke Genetics Consortium (ISGC) distribution files

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### **These are the results files from study:**

“Meta-analysis of Genome-wide Association Studies Identifies 1q22 as a Susceptibility Locus for Intracerebral Hemorrhage”

### **Disclaimer**

GWAS results utilized in the paper were obtained utilizing a p-value based meta-analysis approach. Because this approach does not yield effect estimates or standard errors, the enclosed summary statistics correspond to meta-analyses performed using an inverse variance, fixed effect approach. This meta-analytical approach yields the same conclusions but p-values for each SNP may be slightly different than those described in the paper.

These data are provided "as is", and without warranty, for scientific and educational use only. If you download these data, you acknowledge that the investigator is in compliance with all applicable state, local, and federal laws or regulations and institutional policies regarding human subjects and genetics research; that secondary distribution of the data without approval by secondary parties is prohibited; and that the investigator will cite the appropriate ISGC publication in any communications or publications arising directly or indirectly from these data. Put more bluntly: there are a multitude of ways to erroneously use these data. The onus is on you to use these data correctly. If you intentionally or unintentionally violate the data use instructions, the responsibility falls to you only.

### **Caution**

Some cases and controls were used by multiple studies across the ISGC. This is a non-trivial, non-ignorable issue. Information on the specific cohorts used in each study can be found in the appropriate paper.

### **Identifiability**

There have been extensive discussions in the human genetics community about SNP level data release. Under some circumstances, it is possible to determine if a person was a case or a control in a GWAS. This is not generally the case, and requires near-ideal circumstances and an independent DNA source and genotyping results for a large number of markers. The data included in this public ISGC distribution differ importantly from the idealized scenario. No individual data are being released. No case/control allele frequencies are included (in aggregate across samples or within each sample). Summary data per SNP were generated by imputing individual data onto a common backbone, analysis with PCA and study covariates, and then results from many different studies were combined to yield summary results. The risk of identifiability using this summary level data has been extensively discussed with expert statistical geneticists. The consensus is that the risk of identifiability, even in the strange case in which a DNA sample from a member of an ISGC study was obtained and analyzed, is extremely small. The Principal Investigators of the ISGC approved this plan for release of results.

### **Files**

See the primary papers (above) for full technical details that produced the enclosed results. Again, appropriate use of these files is entirely your responsibility.

### **Citation**

If you use these data, you must (1) acknowledge that “data were accessed through the ISGC Cerebrovascular Disease Knowledge Portal” and (2) cite the appropriate ISGC publication (see above).