

Methods

Study Selection

We sought papers published by investigators referencing datasets from The Genetic Association Information Network (GAIN). We searched through papers included in The GAIN Data Access Committee (GAIN DAC) annual reports of investigators requesting access to GAIN genotyping data.

Only publications written by study investigators acknowledging use of GAIN datasets were included in the Publications Using GAIN Genotyping Data bibliography. These publications were scanned for search terms relating to GAIN or dbGaP (ie: *GAIN*, *dbGaP*, *data accession number*, etc).

Publications Using GAIN Genotyping Data

1. Jian XQ, Wang KS, Wu TJ, Hillhouse JJ, Mullersman JE. [Association of ADAM10 and CAMK2A Polymorphisms with Conduct Disorder: Evidence from Family-Based Studies](#). J Abnorm Child Psychol 39: 773-82. 2011.
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8. Sun J, Wan C, Jia P, Fanous AH, Kendler KS, Riley BP, Zhao Z. [Application of systems biology approach identifies and validates GRB2 as a risk gene for schizophrenia in the Irish Case Control Study of Schizophrenia \(ICCS\) sample](#). Schizophr Res 125: 201-8. 2011.
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