

NHGRI GWAS Catalog: Description of Column Headings for HTML Catalog

DATE ADDED TO CATALOG: Date added to catalog
PUBMEDID: PubMed identification number
FIRST AUTHOR: Last name of first author
DATE: Publication date (online (epub) date if available)
JOURNAL: Abbreviated journal name
LINK: PubMed URL
STUDY: Title of paper (linked to PubMed abstract)
DISEASE/TRAIT: Disease or trait examined in study
INITIAL SAMPLE SIZE: Sample size for Stage 1 of GWAS
REPLICATION SAMPLE SIZE: Sample size for subsequent replication(s)
REGION: Cytogenetic region associated with rs number (NCBI)
REPORTED GENE (S): Gene(s) reported by author
MAPPED GENE(S): Gene(s) mapped to the strongest SNP (NCBI). If the SNP is located within a gene, that gene is listed. If the SNP is intergenic, the upstream and downstream genes are listed, separated by a hyphen. Gene info is linked to Entrez Gene.
STRONGEST SNP-RISK ALLELE: SNP(s) most strongly associated with trait + risk allele (? for unknown risk allele). May also refer to a haplotype. SNP info is linked to dbSNP.
SNPS: Strongest SNP; if a haplotype is reported above, may include more than one rs number (multiple SNPs comprising the haplotype). Asterisk indicates that the published rs number has since been merged with a different rs number (linked to current record in dbSNP).
CONTEXT: SNP functional class (NCBI)
RISK ALLELE FREQUENCY: Reported risk allele frequency associated with strongest SNP
P-VALUE: Reported p-value for strongest SNP risk allele (linked to dbGaP Association Browser)
P-VALUE (TEXT): Information describing context of p-value (e.g. females, smokers). Note that p-values are rounded to 1 significant digit (for example, a published p-value of 4.8×10^{-7} is rounded to 5×10^{-7}).
OR or BETA: Reported odds ratio or beta-coefficient associated with strongest SNP risk allele
95% CI (TEXT): Reported 95% confidence interval associated with strongest SNP risk allele
PLATFORM (SNPS PASSING QC): Genotyping platform manufacturer used in Stage 1; also includes notation of pooled DNA study design or imputation of SNPs, where applicable
CNV: Study of copy number variation (yes/no)

Updated: May 19, 2011