

Title: Genome-wide Analysis of Insomnia (N=1,331,010) Identifies Novel Loci and Functional Pathways

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Includes **Supplementary Information, Supplementary Figures 1 and 2** in separate pdf file,
and **Supplementary Tables 1-28** in separate excel file

1 **Abstract**

2 Insomnia is the second-most prevalent mental disorder, with no sufficient treatment available.
3 Despite a substantial role of genetic factors, only a handful of genes have been implicated and
4 insight into the associated neurobiological pathways remains limited. Here, we use an
5 unprecedented large genetic association sample (N=1,331,010) to allow detection of a
6 substantial number of genetic variants and gain insight into biological functions, cell types
7 and tissues involved in insomnia. We identify 202 genome-wide significant loci implicating
8 956 genes through positional, eQTL and chromatin interaction mapping. We show
9 involvement of the axonal part of neurons, of specific cortical and subcortical tissues, and of
10 two specific cell-types in insomnia: striatal medium spiny neurons and hypothalamic neurons.
11 These cell-types have been implicated previously in the regulation of reward processing,
12 sleep and arousal in animal studies, but have never been genetically linked to insomnia in
13 humans. We found weak genetic correlations with other sleep-related traits, but strong
14 genetic correlations with psychiatric and metabolic traits. Mendelian randomization identified
15 causal effects of insomnia on specific psychiatric and metabolic traits. Our findings reveal
16 key brain areas and cells implicated in the neurobiology of insomnia and its related disorders,
17 and provide novel targets for treatment.

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21 Insomnia is the second-most prevalent mental disorder¹. One third of the general population
22 reports insomnia complaints. The diagnostic criteria for Insomnia Disorder² (i.e. difficulties
23 with initiating or maintaining sleep with accompanying daytime complaints at least three
24 times a week for at least three months, which cannot be attributed to inadequate
25 circumstances for sleep³) are met by 10%, up to one third in samples of older age⁴. Insomnia
26 contributes significantly to the risk and severity of cardiovascular, metabolic, mood, and
27 neurodegenerative disorders². Despite evidence of a considerable genetic component
28 (heritability 38-59%⁵), only a small number of genetic loci moderating the risk of insomnia
29 have thus far been identified^{6,7}. Recent genome-wide association studies^{6,7} (GWAS) for
30 insomnia complaints (N=113,006) demonstrated its polygenic architecture and implicated
31 three genome-wide significant (GWS) loci and seven genes. A prominent role was reported
32 for *MEIS1*, which showed pleiotropic effects for insomnia complaints and restless legs
33 syndrome (RLS)⁷, yet the role of other genes was not unambiguously shown. We set out to
34 substantially increase the sample size to allow the detection of more genetic risk variants for
35 insomnia complaints, which may aid in understanding its neurobiological mechanisms. By
36 combining data collected in the UK Biobank v2⁸ (UKB; N=386,533) and 23andMe, Inc., a
37 personal genetics company^{9,10} (N=944,477), we obtained an unprecedented sample size of
38 1,331,010 individuals. Insomnia complaints were measured using questionnaire data, and the
39 specific questions were validated to be good proxies of insomnia disorder, using an
40 independent sample (The Netherlands Sleep Register, NSR)¹¹ in which we had access to
41 similar question data, as well as clinical interviews assessing insomnia disorder (see
42 **Supplementary Methods 1.1-1.3**). We find 202 risk loci for insomnia complaints, and
43 extensive functional in silico analyses reveal the involvement of specific tissue and cell types,
44 whereas secondary statistical analyses reveal causal effects of insomnia on metabolic and
45 psychiatric traits.

46 **Meta-analysis yields 202 risk loci**

47 UKB assessed insomnia complaints (hereafter referred to as ‘insomnia’) using a touchscreen
48 device while 23andMe research participants completed online surveys. Assessment of
49 insomnia in both samples shows high accuracy (sensitivity=84-98%; specificity=80-96%) for
50 Insomnia Disorder (see **Supplementary Methods 1.3**). The prevalence of insomnia in the
51 UKBv2 sample was 28.3%, 30.5% in the 23andMe sample, and 29.3% in the combined
52 sample, in keeping with previous estimates for people with advanced age in the UK⁴ and
53 elsewhere^{12,13}. Older people dominate the UKB sample (mean age=56.7, SD=8.0) and the
54 23andMe sample (two-thirds of the sample older than 45, one-third even older than 60 years
55 of age). Prevalence was higher in females (34.6%) than males (24.5%), yielding an odds ratio
56 (OR) of 1.6, close to the OR of 1.4 reported in a meta-analysis¹⁴.

57 Quality control was conducted separately per sample, following standardized, stringent
58 protocols (see **Methods**). GWAS was run separately per sample (UKB; N=386,533,
59 23andMe, Inc.; N=944,477) (**Extended Data Fig. 1**), and then meta-analyzed using
60 METAL¹⁵ by weighing SNP effects by sample size (see **Methods**). We first analyzed males
61 and females separately (**Extended Data Fig. 2, 3**), and observed a high genetic correlation
62 between the sexes ($r_g=0.92$, SE=0.02, **Extended Data Table 1**), indicating strong overlap of
63 genetic effects. Owing to the large sample size, the r_g of 0.92 was significantly different from
64 1 (one-sided Wald test, $P=2.54\times 10^{-6}$) suggesting a small role for sex-specific genetic risk
65 factors, consistent with our previous report⁷. However, since sex-specific effects were
66 relatively small, we here focus on identifying genetic effects important in both sexes and
67 continued with the combined sample (**Supplementary Table 1, 2** and **Supplementary**
68 **Discussion 2.1** provide sex-specific results).

69 We observe significant polygenic signal in the GWAS (lambda inflation factor=1.808) which
70 could not be ascribed to spurious association (LD Score intercept=1.075)¹⁶ (**Extended Data**

71 **Fig. 4a**). Meta-analysis identified 11,990 genome-wide significant (GWS) SNPs ($P < 5 \times 10^{-8}$),
72 represented by 248 independent lead SNPs ($r^2 < 0.1$), located in 202 genomic risk loci (**Fig.**
73 **1a**, **Supplementary Fig. 1** and **Supplementary Table 3, 4**). All lead SNPs showed
74 concordant signs of effect in both samples (**Extended Data Fig. 4b**). We confirm two
75 (chr2:66,785,180 and chr5:135,393,752) out of six previously reported loci for insomnia ^{6,7}
76 (**Supplementary Table 5**). Polygenic score (PGS) prediction in three randomly selected
77 hold-out samples (N=3×3,000) estimated the current results to explain up to 2.6% of the
78 variance in insomnia (**Fig. 1b**, **Extended Data Fig. 5** and **Supplementary Table 6**).

79 The SNP-based heritability (h^2_{SNP}) was estimated at 7.0% (SE=0.002). Partitioning the
80 heritability by functional categories of SNPs (see **Methods**) showed the strongest enrichment
81 of h^2_{SNP} in conserved regions (enrichment=15.8, $P=1.57 \times 10^{-14}$). In addition, h^2_{SNP} was
82 enriched in common SNPs (MAF > 0.3) and depleted in more rare SNPs (MAF < 0.01; **Fig. 1c**
83 and **Supplementary Table 7**).

84 We used FUMA¹⁷ to functionally annotate all 22,068 SNPs in the risk loci that were in LD
85 ($r^2 \geq 0.6$) with one of the independent significant SNPs (see **Methods**). The majority of these
86 SNPs (76.8%) were in open chromatin regions¹⁸ as indicated by a minimum chromatin state
87 of 1-7 (**Fig. 1d** and **Supplementary Table 8**). In line with findings for other traits^{7,19}, about
88 half of these SNPs were in intergenic (35.5%) or non-coding RNA (13.0%) regions (**Fig. 1e**),
89 and of these, 0.72% were highly likely to have a regulatory function as indicated by a
90 RegulomeDB Score < 2 (see **Methods**). However, of these 51.5% were located inside a
91 protein coding gene and 0.81% were exonic. Of the 177 exonic SNPs, 71 were exonic non-
92 synonymous (ExNS, **Supplementary Table 9**). *WDR90* included four ExNS (rs7190775,
93 rs4984906, rs3752493, and rs3803697) all in high LD with the same independent significant
94 SNP (rs3184470). There were two ExNS SNPs with extremely high Combined Annotation
95 Dependent Depletion (CADD) scores²⁰ suggesting a strong deleterious effect on protein

96 function: rs13107325 in *SLC39A8* (locus 56, $P=8.31\times 10^{-16}$) with the derived allele T
97 (MAF=0.03) associated with an increased risk of insomnia, and rs35713889 in *LAMB2* (locus
98 42, $P=1.77\times 10^{-7}$), where the derived allele T of rs35713889 (MAF=0.11) was also associated
99 with an increased risk of insomnia complaints. **Supplementary Table 10** and
100 **Supplementary Discussion 2.2** provide a detailed overview of the functional impact of all
101 variants in the genomic risk loci.

102

103 **Genes implicated in insomnia**

104 To obtain insight into (functional) consequences of individual GWS SNPs we used FUMA¹⁷
105 to apply three strategies to map associated variants to genes (see **Methods**). Positional gene-
106 mapping aligned SNPs to 412 genes by location. Expression Quantitative Trait Loci (eQTL)
107 gene-mapping matched cis-eQTL SNPs to 594 genes whose expression levels they influence.
108 Chromatin interaction mapping annotated SNPs to 159 genes based on three-dimensional
109 DNA-DNA interactions between genomic regions of the GWS SNPs and nearby or distant
110 genes (**Supplementary Fig. 2, Supplementary Table 11** and **Supplementary Discussion**
111 **2.3**). 91 genes were mapped by all three strategies (**Supplementary Table 12**) and 336 genes
112 were physically located outside of the risk loci but were implicated by eQTL associations
113 (306 genes), chromatin interactions (16 genes) or both (14 genes). Several genes were
114 implicated by GWS SNPs originating from two distinct risk loci on the same chromosome
115 (**Fig. 2a** and **2b**): *MEIS1*, located on chromosome 2 in the strongest associated locus (locus
116 20), was positionally mapped by 51 SNPs and mapped by chromatin interactions in 10 tissue
117 types including cross-loci interactions from locus 21, and is a known gene involved in
118 insomnia⁷. *LRGUK*, located on chromosome 7 in locus 106, was positionally mapped by 22
119 SNPs and chromatin interactions in 3 tissue types including cross-loci interactions from locus
120 105. *LRGUK* was also implicated by eQTLs associations of 125 SNPs in 14 general tissue

121 types. *LRGUK* was previously implicated in type 2 diabetes²¹ and autism spectrum disorder²²
122 - disorders with prominent insomnia - but not yet directly implicated in sleep-related
123 phenotypes, and is the most likely candidate to explain the observed association in loci 105
124 and 106.

125 Apart from linking individual associated genetic variants to genes, we conducted a genome-
126 wide gene-based association analysis (GWGAS) using MAGMA²³. GWGAS provides
127 aggregate association *P*-values based on all variants located in a gene, and complements the
128 three FUMA mapping strategies (see **Methods**). GWGAS identified 517 associated genes
129 (**Fig. 2c** and **Supplementary Table 13**). The top gene *BTBD9* ($P=8.51\times 10^{-23}$) on
130 chromosome 6 in locus 81 was also mapped by positional and eQTL mapping (tissue type:
131 left ventricle of the heart), and is part of a pathway regulating circadian rhythms. *BTBD9* has
132 been associated with RLS, periodic limb movement disorder^{24,25} and Tourette Syndrome²⁶.
133 Involvement in sleep regulation was shown in *Drosophila*²⁷, and mouse mutants show
134 fragmented sleep²⁸ and increased levels of dynamin 1²⁹, a protein that mediates the increased
135 sleep onset latency following pre-sleep arousal³⁰.

136 Of the 517 MAGMA-based associated genes, 222 were outside of the GWAS risk loci, and
137 309 were also mapped by FUMA. In total, 956 unique genes were mapped by at least one of
138 the three FUMA gene mapping strategies or by MAGMA (**Extended Data Fig. 6**). Of these,
139 *MEIS1*, *MED27*, *IPO7* and *ACBD4* confirmed previous results^{6,7} (**Supplementary Table 14**).

140 Sixty-two genes were implicated by all four mapping strategies indicating that apart from a
141 GWS gene-based *P*-value, there were (i) GWS SNPs located inside these genes, (ii) GWS
142 SNPs associated with differential expression of these genes and (iii) GWS SNPs that were
143 involved in genomic regions interacting with these genes. We note that genes that were
144 indicated by positional mapping and GWS gene-based *P*-values, but not via eQTL or
145 chromatin interaction mapping (N=54 genes), may be of equal importance, yet are more

146 likely to exert their influence on insomnia via structural changes in the gene products (i.e. at
147 the protein level) and not via quantitative changes in the availability of the gene products.

148

149 **Implicated pathways, tissues and cell-types**

150 To test whether GWS genes converged in functional gene-sets and pathways, we conducted
151 gene-set analyses using MAGMA (see **Methods**). We tested associations of 7,473 gene-sets:
152 7,246 sets derived from the MsigDB³¹, gene expression values from 54 tissues from the
153 GTEx database³², and cell-specific gene expression in 173 types of brain cells (**Fig. 2d**,
154 **Supplementary Table 15**). Competitive testing was used and a Bonferroni corrected
155 threshold of $P < 6.7 \times 10^{-6}$ ($0.05/7,473$) to correct for multiple testing. Of the MsigDB gene-
156 sets, three Gene Ontology (GO) gene-sets survived multiple testing: GO:*locomotory behavior*
157 ($P = 8.95 \times 10^{-7}$), GO:*behavior* ($P = 5.23 \times 10^{-6}$), and GO:*axon part* ($P = 4.25 \times 10^{-6}$). Twelve genes
158 (*LRRK2*, *CRH*, *DLG4*, *DNMI*, *DRD1*, *DRD2*, *DRD4*, *GRIN1*, *NTSRI*, *SNCA*, *CNTN2*, and
159 *CALBI*) were included in all of these gene-sets and two of these (*SNCA*, *DNMI*) had a GWS
160 gene-based *P*-value (**Supplementary Table 16**). *SNCA* encodes alpha-synuclein and has
161 been implicated in REM sleep behavior disorder³³ and Parkinson's disease³⁴. Altered
162 expression in mice changes sleep and wake EEG spectra³⁵ along the same dimensions that
163 have been implicated in insomnia disorder³⁶. *DNMI* encodes the synaptic neuronal protein
164 dynamin 1, which is increased in *BTBD9* mutant mice²⁹ and mediates the sleep-disruptive
165 effect of pre-sleep arousal (see above; *BTBD9* is the top associated gene). Conditional gene-
166 set analyses suggested that the association with the gene-set *behavior* is almost completely
167 explained by the association of *locomotory behavior*, and that the effect of *axon part* is
168 independent of this (**Supplementary Discussion 2.4**). GO:*locomotory behavior* includes 175
169 genes involved in stimulus-evoked movement³⁷. This set included 16 GWS genes: *BTBD9*,
170 *MEIS1*, *DABI*, *SNCA*, *GNAO1*, *ATP2B2*, *NEGR1*, *SLC4A10*, *GIP*, *DNMI*, *GPRC5B*, *GRM5*,

171 *NRG1*, *PARK2*, *TALI*, and *OXR1*). GO:*axon part* reflects a very general cellular component
172 representing 219 genes, of which 14 were GWS (*KIF3B*, *SNCA*, *GRIA1*, *CDH8*, *ROBO2*,
173 *DNMI1*, *RANGAP1*, *GABBR1*, *P2RX3*, *NRG1*, *POLG*, *DAG1*, *NFASC*, and *CALB2*).

174 Tissue specific gene-set analyses showed strong enrichment of genetic signal in genes
175 expressed in the brain. Correcting for overall expression, four specific brain tissues reached
176 the threshold for significance: overall cerebral cortex ($P=3.68\times 10^{-6}$), Brodmann area 9 (BA9)
177 of frontal cortex ($P=5.04\times 10^{-7}$), BA24 of the anterior cingulate cortex ($P=3.25\times 10^{-6}$), and
178 cerebellar hemisphere ($P=5.93\times 10^{-6}$)¹. Several other brain tissues also showed strong
179 enrichment just below threshold, including three striatal basal ganglia (BG) structures
180 (nucleus accumbens, caudate nucleus, putamen). To test whether genes expressed in all three
181 BG structures together would show significant enrichment of low *P*-values, we used the first
182 principal component (BG_{PC}) of these BG structures and found significant enrichment
183 ($P=8.33\times 10^{-8}$). When conditioning the three top cortical structures on the BG_{PC}, they were no
184 longer significantly associated after multiple testing correction (minimum $P=0.03$), which
185 was expected given that the BG_{PC} correlated strongly with gene-expression in cortical (and
186 other) areas ($r>0.96$). Similar results were obtained vice versa, i.e. using the first principal
187 component of all cortical areas and conditioning the three BG structures on this resulted in no
188 evidence of enrichment of low *P*-values for BG structures (minimum $P=0.53$). These results
189 show that (i) genes expressed in brain are important in insomnia, (ii) genes expressed in
190 cortical areas are more strongly associated than genes expressed in BG, (iii) there is a strong
191 correlation between gene expression patterns across brain tissues, which suggests
192 involvement of general cellular signatures more than specific brain tissue structures.

¹ We caution that only a limited set of brain tissues were included and thus we cannot rule out associations with many important areas such as pons, midbrain or thalamus based on this analysis.

193 Brain cell type-specific gene-set analyses was first carried out on 24 broad cell-type
194 categories. Cell type-specific gene expression was quantified using single cell RNA-
195 sequencing of dissociated cells from somatosensory cortex, hippocampus, hypothalamus,
196 striatum and midbrain from mouse (see **Methods**), which closely resembles gene-expression
197 in humans³⁸. Results indicated that genes expressed specifically in the medium spiny neurons
198 (MSN, $P=4.83\times 10^{-7}$) were associated with insomnia, and no other broad cell-types specific
199 gene-set survived our strict threshold of $P<6.7\times 10^{-6}$. MSNs represent 95% of neurons within
200 the human striatum, which is one of the four major nuclei of the subcortical BG. Specifically,
201 the striatum consists of the ventral (nucleus accumbens and olfactory tubercle) and dorsal
202 (caudate nucleus and putamen) subdivisions. The association with MSNs thus likely explains
203 the observed association of the BG striatal structures (nucleus accumbens, caudate nucleus,
204 putamen).

205 Using broad cell classes risks not detecting associations that are specific to distinctive yet rare
206 cell types; to account for this we then tested 149 specific brain cell-type categories, and found
207 significant associations with 7 specific cell types: medio-lateral neuroblasts type 3, 4 and 5
208 ($P=2.36\times 10^{-6}$, $P=1.88\times 10^{-6}$, and $P=1.87\times 10^{-6}$), D2 type medium spiny neurons ($P=2.12\times 10^{-6}$),
209 claustrum pyramidal neurons ($P=3.09\times 10^{-6}$), hypothalamic Vglut2 Morn4 Prrc2a neurons
210 ($P=4.36\times 10^{-6}$), and hypothalamic Vglut2 Hcn16430411 K18 Rik neurons ($P=4.98\times 10^{-6}$),
211 known to have the densest number of melatonin receptors. These results suggest a role of
212 distinct mature and developing cell types in the midbrain and hypothalamus. The
213 hypothalamus contains multiple nuclei that are key to the control of sleep and arousal,
214 including the suprachiasmatic nucleus (SCN) that accommodates the biological clock of the
215 brain³⁹.

216

217

218 **Low genetic overlap with sleep traits**

219 Other sleep-related traits may easily be confounded with specific symptoms of insomnia, like
220 early morning awakening, difficulties maintaining sleep, and daytime sleepiness. The most
221 recent genome-wide studies for other sleep-related traits included 59,128 to 128,266
222 individuals, and assessed genetic effects on morningness^{6,40,41} (i.e. being a morning person),
223 sleep duration^{6,41}, and daytime sleepiness/dozing⁴¹. Using increased sample sizes for each of
224 these sleep-related traits (max N=434,835), we here investigated to what extent insomnia and
225 other sleep-related traits are genetically distinct or overlapping. We performed GWAS
226 analyses for the following six sleep-related traits: morningness, sleep duration, ease of getting
227 up in the morning, naps during the day, daytime dozing, and snoring (**Supplementary**
228 **Methods 1.1-1.2, Extended Data Fig. 7, 9**). Of the 202 risk loci for insomnia, 39 were also
229 GWS in at least one of the other sleep-related traits (**Fig. 3, Supplementary Table 17**). The
230 strongest overlap in loci was found with sleep duration, with 14 out of 49 sleep duration loci
231 overlapping with insomnia. Insomnia showed the highest genetic correlation with sleep
232 duration (-0.47 , $SE=0.02$; **Supplementary Table 18**) compared to other sleep-related traits,
233 which was not surprising given that insomnia also shared the most risk loci with sleep
234 duration (See further discussion sleep phenotypes in **Supplementary Discussion 2.5**).

235 Gene-mapping of SNP associations of sleep-related traits resulted in 973 unique genes
236 (**Extended Data Fig. 9, Supplementary Table 19-23**). Gene-based analysis showed that of
237 the 517 GWS genes for insomnia, 120 were GWS in at least one of the other sleep-related
238 traits, and one gene (*RBFox1*) was GWS in all except napping and dozing (**Supplementary**
239 **Table 24**). The largest proportion of overlap in GWS genes for insomnia was again with
240 sleep duration, with 37 of the 135 (27%) GWS genes for sleep duration also GWS for
241 insomnia. There was overlap in tissue enrichment in cortical structures and basal ganglia
242 between insomnia and both morningness and sleep duration. On the single cell level, the

243 medium spiny neurons were also implicated for morningness and sleep duration, but not for
244 the other sleep-related traits (**Supplementary Table 25**). Taken together, these results
245 suggest that at a genetic level, insomnia shows partial overlap with sleep duration, but
246 minimal overlap with other sleep-related traits. Consistent short sleep across nights occurs
247 only in a minor part of insomnia patients, even in a clinical sample⁴².

248

249 **Strong overlap between insomnia and psychiatric and metabolic traits**

250 We confirm previously reported genetic correlations between insomnia and neuropsychiatric
251 and metabolic traits^{6,7} (**Supplementary Table 26**), and also identify several GWS SNPs for
252 insomnia that have previously been associated with these traits (**Supplementary Table 27**).

253 The strongest correlations were with depressive symptoms ($r_g=0.64$, $SE=0.04$ $P=1.21\times 10^{-71}$),
254 followed by anxiety disorder ($r_g=0.56$, $SE=0.11$ $P=1.40\times 10^{-7}$), subjective well-being
255 ($r_g=-0.51$, $SE=0.03$ $P=4.93\times 10^{-52}$), major depression ($r_g=0.50$, $SE=0.07$ $P=8.08\times 10^{-12}$) and

256 neuroticism ($r_g=0.48$, $SE=0.02$ $P=8.72\times 10^{-80}$). Genetic correlations with metabolic traits
257 ranged between 0.09-0.20. The genetic correlations between insomnia and psychiatric traits
258 were also stronger than the correlations between insomnia and the other sleep-related traits.

259 Since a similar high reliability has been reported for both sleep and psychiatric phenotypes,
260 the findings suggest that genetically insomnia more closely resembles neuropsychiatric traits
261 than it resembles other sleep-related traits (**Fig. 4**). To infer directional associations between

262 insomnia and these correlated traits, we performed bidirectional Multi-SNP Mendelian
263 Randomization (MR) analysis⁴³ (see **Methods**). Results support a direct risk effect of
264 insomnia on metabolic syndrome phenotypes including BMI ($b_{xy}=0.36$, $SE=0.05$,

265 $P=1.25\times 10^{-12}$) type 2 diabetes ($b_{xy}=0.62$, $SE=0.11$, $P=2.29\times 10^{-8}$), and coronary artery disease
266 ($b_{xy}=0.61$, $SE=0.09$, $P=2.88\times 10^{-12}$). In addition, insomnia was bidirectionally associated with

267 educational attainment, with a stronger effect from insomnia on educational attainment

268 ($b_{xy}=-0.32$, $SE=0.02$, $P=1.68\times 10^{-77}$) (i.e. a higher risk for insomnia leads to lower
269 educational attainment) than vice versa ($b_{xy}=-0.10$, $SE=0.01$, $P=2.27\times 10^{-23}$), the same pattern
270 was observed for intelligence. We also found risk effects of insomnia on several psychiatric
271 traits, including schizophrenia ($b_{xy}=0.68$, $SE=0.10$, $P=5.12\times 10^{-11}$), ADHD ($b_{xy}=0.77$,
272 $SE=0.06$, $P=2.50\times 10^{-45}$), neuroticism ($b_{xy}=0.46$, $SE=0.03$, $P=3.92\times 10^{-53}$) and anxiety disorder
273 ($b_{xy}=0.47$, $SE=0.10$, $P=4.11\times 10^{-6}$), with no evidence of large reverse effects, except for a
274 small risk effect of neuroticism on insomnia ($b_{xy}=0.09$, $SE=0.02$, $P=1.24\times 10^{-6}$) and
275 depressive symptoms ($b_{xy}=0.09$, $SE=0.02$, $P=1.24\times 10^{-6}$)². Overall, there was only a small
276 proportion of SNPs showing pleiotropy between insomnia and other traits (**Supplementary**
277 **Table 28** and **Supplementary Discussion 2.6**).

278

279 Discussion

280 In the largest GWAS study to date of 1,331,010 participants we identified 202 genomic risk
281 loci for insomnia. Using extensive functional annotation of associated genetic variants, we
282 demonstrated that the genetic component of insomnia points towards a role of genes involved
283 in locomotory behavior, and genes expressed in specific cell types from the claustrum,
284 hypothalamus and striatum, and specifically in MSNs (**Fig. 5**). MSNs are GABAergic
285 inhibitory cells and represent 95% of neurons in the human striatum, one of the four major
286 nuclei of the BG (for reviews, see ⁴⁴⁻⁴⁶). MSNs receive massive excitatory glutamatergic
287 input from the cerebral cortex and the thalamus, and are targets of dopamine neurons in
288 substantia nigra and the ventral tegmental area. In addition, they receive inhibitory inputs
289 from striatal GABAergic interneurons. MSNs themselves are GABAergic output neurons
290 with exceptionally long projections to globus pallidus (GP), substantia nigra and ventral

² We do note that for major depression the reverse MR could not be carried out due to an insufficient number of SNPs with a low P -value

291 pallidum, and control the activity of thalamocortical neurons. Previous studies during the
292 natural sleep-wake cycle, *in vitro*, and from anesthetized *in vivo* preparations have shown that
293 MSNs show fast, synchronized cyclic firing, i.e. the so-called Up and Down states, during
294 slow-wave sleep and irregular pattern of action potentials during wakefulness. In fact, MSNs
295 were the first neurons in which the Up and Down states characteristic of slow wave sleep
296 were described⁴⁷. Cell body-specific striatal lesions of the rostral striatum induce a profound
297 sleep fragmentation, which is most characteristic of insomnia. A role for BG in sleep
298 regulation is also suggested by the high prevalence of insomnia in neurodegenerative
299 disorders, such as Parkinson's Disease and Huntington's disease in which the BG are
300 affected. Vetrivelan et al.⁴⁴ proposes a cortex-striatum-GP_{external}-cortex network involved in
301 the control of sleep-wake behavior and cortical activation, in which midbrain dopamine
302 disinhibits the GP_{external} and promotes sleep through activation of D2 receptors in this
303 network. Furthermore, brain imaging studies have suggested the caudate nucleus of the
304 striatum as a key node in the neuronal network imbalance of insomnia⁴⁸, and also reported
305 abnormal function in the cortical areas we found to be most enriched (BA9⁴⁹, BA24⁵⁰). Our
306 results support the involvement of the striato-cortical network in insomnia, by showing
307 enrichment of risk genes for insomnia in cortical areas as well as the striatum, and
308 specifically in MSNs. We recently showed that, along with several other cell types, MSNs
309 also mediate the risk for mood disorders⁵¹ and schizophrenia³⁸. MSNs are strongly implicated
310 in reward processing and future work could address whether the genetic overlap between
311 insomnia and mood disorders is mediated by gene function in MSNs.

312 Our results also showed enrichment of insomnia genes in pyramidal neurons of the claustrum.
313 This subcortical brain region is structurally closely associated with the amygdala and has
314 been implicated in salience coding of incoming stimuli and binding of multisensory
315 information into conscious percepts⁵². These functions are highly relevant to insomnia,

316 because the disorder is characterized by increased processing of incoming stimuli⁵³ and by
317 ongoing consciousness even during sleep, a phenomenon known as sleep state
318 misperception⁵⁴. We also found enrichment of insomnia genes in mediolateral neuroblasts
319 from the embryonic midbrain and in two hypothalamic cell types. The role of the
320 mediolateral neuroblasts is less clear; although they were obtained from the embryonic
321 midbrain, it is at present unknown what type of mature neurons they differentiate into. We
322 note that the midbrain is similar on a bulk transcriptomic level to the pons⁵⁵, and lacking cells
323 from that region we cannot conclusively say that midbrain cell-types are enriched.

324 The current findings provide novel insight into the causal mechanism of insomnia,
325 implicating specific cell types, brain areas and biological functions. These findings are
326 starting points for the development of new therapeutic targets for insomnia and may also
327 provide valuable insights for other, genetically related disorders.

328 **Methods:**

329 **Meta-analysis**

330 A meta-analysis on the GWAS results of insomnia and morningness in UKB and 23andMe
331 cohorts was performed using fixed-effects meta-analysis METAL¹⁵, using SNP *P*-values
332 weighted by sample size. To investigate sex-specific genetic effects, we ran the meta-analysis
333 between UKB and 23andMe datasets for males and females separately.

334

335 **Genomic risk loci definition**

336 We used FUMA¹⁷ (<http://fuma.ctglab.nl/>), an online platform for functional mapping and
337 annotation of genetic variants, to define genomic risk loci and obtain functional information
338 of relevant SNPs in these loci. FUMA provides comprehensive annotation information by
339 combining several external data sources. We first identified *independent significant SNPs* that
340 have a genome-wide significant *P*-value ($<5 \times 10^{-8}$) and are independent from each other at
341 $r^2 < 0.6$. These SNPs were further represented by *lead SNPs*, which are a subset of the
342 independent significant SNPs that are in approximate linkage equilibrium with each other at
343 $r^2 < 0.1$. We then defined associated *genomic risk loci* by merging any physically overlapping
344 lead SNPs (linkage disequilibrium [LD] blocks < 250 kb apart). Borders of the genomic risk
345 loci were defined by identifying all SNPs in LD ($r^2 \geq 0.6$) with one of the independent
346 significant SNPs in the locus, and the region containing all these *candidate SNPs* was
347 considered to be a single independent genomic risk locus. LD information was calculated
348 using the UK Biobank genotype data as a reference. Risk loci were defined based on
349 evidence from independent significant SNPs that were available in both 23andMe and UKB.
350 We note that SNPs that were GWS but only available in the 23andMe dataset were not
351 included when defining genomic risk loci and were not included in any follow-up annotations
352 or analyses, because there was no external replication in the UKB sample. If such SNPs were

353 located in a risk locus, they are displayed in Locuszoom plots (grey, as there is no LD
354 information in UKB). When risk loci contained GWS SNPs based solely on 23andMe, we did
355 not count that risk locus, as there were no other SNPs available in both samples that
356 supported these GWS SNPs.

357

358 **Gene-based analysis**

359 SNP-based *P*-values from the meta-analysis were used as input for the gene-based genome-
360 wide association analysis (GWGAS). 18,182 to 18,185 protein-coding genes (each containing
361 at least one SNP in the GWAS, the total number of tested genes can thus be slightly different
362 across phenotypes) from the NCBI 37.3 gene definitions were used as basis for GWGAS in
363 MAGMA²³. Bonferroni correction was applied to correct for multiple testing ($P < 2.73 \times 10^{-6}$).

364

365 **Gene-set analysis**

366 Results from the GWGAS analyses were used to test for association in three types of 7,473
367 predefined gene-sets:

- 368 1. 7,246 curated gene-sets representing known biological and metabolic pathways
369 derived from 9 data resources, catalogued by and obtained from the MsigDB version
370 6.0⁵⁶ (<http://software.broadinstitute.org/gsea/msigdb/collections.jsp>)
- 371 2. Gene expression values from 54 (53 + 1 calculated 1st PC of three tissue subtypes)
372 tissues obtained from GTEx³², log₂ transformed with pseudocount 1 after
373 winsorization at 50 and averaged per tissue
- 374 3. Cell-type specific expression in 173 types of brain cells (24 broad categories of cell
375 types, 'level 1' and 129 specific categories of cell types 'level 2'), which were
376 calculated following the method described in³⁸. Briefly, brain cell-type expression
377 data was drawn from single-cell RNA sequencing data from mouse brains. For each

378 gene, the value for each cell-type was calculated by dividing the mean Unique
379 Molecular Identifier (UMI) counts for the given cell type by the summed mean UMI
380 counts across all cell types. Single-cell gene-sets were derived by grouping genes into
381 40 equal bins based on specificity of expression. Mouse cell gene-expression was
382 shown to closely approximate gene-expression in post-mortem human tissue³⁸.

383 These gene-sets were tested using MAGMA. We computed competitive *P*-values, which
384 represent the test of association for a specific gene-set compared with genes not in the gene-
385 set to correct for baseline level of genetic association in the data⁵⁷. The Bonferroni-corrected
386 significance threshold was $0.05/7,473 \text{ gene-sets}=6.7 \times 10^{-6}$. Conditional analyses were
387 performed as a follow-up using MAGMA to test whether each significant association
388 observed was independent of all others. The association between each gene-set in each of the
389 three categories was tested conditional on the most strongly associated set, and then, if any
390 substantial ($P < 0.05/\text{number of gene-sets}$) associations remained, by conditioning on the first
391 and second most strongly associated set, and so on until no associations remained. Gene-sets
392 that retained their association after correcting for other sets were considered to represent
393 independent signals. We note that this is not a test of association per se, but rather a strategy
394 to identify, among gene-sets with known significant associations and overlap in genes, which
395 set (s) are responsible for driving the observed association.

396

397 **SNP-based heritability and genetic correlation**

398 LD Score regression¹⁶ was used to estimate genomic inflation and SNP-based heritability of
399 the phenotypes, and to estimate the cross-cohort genetic correlations. Pre-calculated LD
400 scores from the 1000 Genomes European reference population were obtained from
401 <https://data.broadinstitute.org/alkesgroup/LDSCORE/>.

402

403 **Genetic correlations**

404 Genetic correlations between sleep-related traits, and between sleep-related traits and
405 previously published GWAS studies of sufficient sample size were calculated using LD Score
406 regression on HapMap3 SNPs only. Genetic correlations were corrected for multiple testing
407 based on the total number of correlations (between 6 sleep-related phenotypes and 27
408 previous GWAS studies) by applying a Bonferroni corrected threshold of
409 ($P < 0.05/33 = 1.51 \times 10^{-3}$).

410

411 **Stratified heritability**

412 To test whether specific categories of SNP annotations were enriched for heritability, we
413 partitioned SNP heritability for binary annotations using stratified LD score regression⁵⁸.
414 Heritability enrichment was calculated as the proportion of heritability explained by a SNP
415 category divided by the proportion of SNPs that are in that category. Partitioned heritability
416 was computed by 28 functional annotation categories, by minor allele frequency (MAF) in
417 six percentile bins and by 22 chromosomes. Annotations for binary categories of functional
418 genomic characteristics (e.g. coding or regulatory regions) were obtained from the LD score
419 website (<https://github.com/bulik/ldsc>). The Bonferroni-corrected significance threshold for
420 56 annotations was set at: $P < 0.05/56 = 8.93 \times 10^{-4}$.

421

422 **Functional annotation of SNPs**

423 Functional annotation of SNPs implicated in the meta-analysis was performed using
424 FUMA¹⁷. We selected all candidate SNPs in genomic risk loci having an $r^2 \geq 0.6$ with one of
425 the independent significant SNPs (see above), a P -value ($P < 1 \times 10^{-5}$), a $MAF > 0.0001$ for
426 annotations, and availability in both UKB and 23andMe datasets. Functional consequences

427 for these SNPs were obtained by matching SNPs' chromosome, base-pair position, and
428 reference and alternate alleles to databases containing known functional annotations,
429 including ANNOVAR⁵⁹ categories, Combined Annotation Dependent Depletion (CADD)
430 scores, RegulomeDB²⁰ (RDB) scores, and chromatin states⁶⁰. ANNOVAR categories identify
431 the SNP's genic position (e.g. intron, exon, intergenic) and associated function. CADD scores
432 predict how deleterious the effect of a SNP is likely to be for a protein structure/function,
433 with higher scores referring to higher deleteriousness. A CADD score above 12.37 is
434 considered to be potentially pathogenic²⁰. The RegulomeDB score is a categorical score
435 based on information from expression quantitative trait loci (eQTLs) and chromatin marks,
436 ranging from 1a to 7 with lower scores indicating an increased likelihood of having a
437 regulatory function. Scores are as follows: 1a=eQTL + Transcription Factor (TF) binding +
438 matched TF motif + matched DNase Footprint + DNase peak; 1b=eQTL + TF binding + any
439 motif + DNase Footprint + DNase peak; 1c=eQTL + TF binding + matched TF motif +
440 DNase peak; 1d=eQTL + TF binding + any motif + DNase peak; 1e=eQTL + TF binding +
441 matched TF motif; 1f=eQTL + TF binding / DNase peak; 2a=TF binding + matched TF motif
442 + matched DNase Footprint + DNase peak; 2b=TF binding + any motif + DNase Footprint +
443 DNase peak; 2c=TF binding + matched TF motif + DNase peak; 3a=TF binding + any motif
444 + DNase peak; 3b=TF binding + matched TF motif; 4=TF binding + DNase peak; 5=TF
445 binding or DNase peak; 6=other;7=Not available. The chromatin state represents the
446 accessibility of genomic regions (every 200bp) with 15 categorical states predicted by a
447 hidden Markov model based on 5 chromatin marks for 127 epigenomes in the Roadmap
448 Epigenomics Project⁶¹. A lower state indicates higher accessibility, with states 1-7 referring
449 to open chromatin states. We annotated the minimum chromatin state across tissues to SNPs.
450 The 15-core chromatin states as suggested by Roadmap are as follows: 1=Active
451 Transcription Start Site (TSS); 2=Flanking Active TSS; 3=Transcription at gene 5' and 3';

452 4=Strong transcription; 5= Weak Transcription; 6=Genic enhancers; 7=Enhancers; 8=Zinc
453 finger genes & repeats; 9=Heterochromatic; 10=Bivalent/Poised TSS; 11=Flanking
454 Bivalent/Poised TSS/Enh; 12=Bivalent Enhancer; 13=Repressed PolyComb; 14=Weak
455 Repressed PolyComb; 15=Quiescent/Low.

456

457 **Gene-mapping**

458 Genome-wide significant loci obtained by GWAS were mapped to genes in FUMA¹⁷ using
459 three strategies:

460 1. Positional mapping maps SNPs to genes based on physical distance (within a 10kb
461 window) from known protein coding genes in the human reference assembly
462 (GRCh37/hg19).

463 2. eQTL mapping maps SNPs to genes with which they show a significant eQTL association
464 (i.e. allelic variation at the SNP is associated with the expression level of that gene). eQTL
465 mapping uses information from 45 tissue types in 3 data repositories (GTEx³², Blood eQTL
466 browser⁶⁰, BIOS QTL browser⁶²), and is based on cis-eQTLs which can map SNPs to genes
467 up to 1Mb apart. We used a false discovery rate (FDR) of 0.05 to define significant eQTL
468 associations.

469 3. Chromatin interaction mapping was performed to map SNPs to genes when there is a
470 three-dimensional DNA-DNA interaction between the SNP region and another gene region.
471 Chromatin interaction mapping can involve long-range interactions as it does not have a
472 distance boundary. FUMA currently contains Hi-C data of 14 tissue types from the study of
473 Schmitt et al⁶³. Since chromatin interactions are often defined in a certain resolution, such as
474 40kb, an interacting region can span multiple genes. If a SNP is located in a region that
475 interacts with a region containing multiple genes, it will be mapped to each of those genes.
476 To further prioritize candidate genes, we selected only interaction-mapped genes in which

477 one region involved in the interaction overlaps with a predicted enhancer region in any of the
478 111 tissue/cell types from the Roadmap Epigenomics Project⁶¹, and the other region is
479 located in a gene promoter region (250bp up and 500bp downstream of the transcription start
480 site and also predicted by Roadmap to be a promoter region). This method reduces the
481 number of genes mapped but increases the likelihood that those identified will indeed have a
482 plausible biological function. We used a $P\text{-FDR} < 1 \times 10^{-5}$ to define significant interactions,
483 based on previous recommendations⁶³, modified to account for the differences in cell lines
484 used here.

485

486 **GWAS catalog lookup**

487 We used FUMA to identify SNPs with previously reported ($P < 5 \times 10^{-5}$) phenotypic
488 associations in published GWAS listed in the NHGRI-EBI catalog⁶⁴, which matched with
489 SNPs in LD with one of the independent significant SNPs identified in the meta-analysis.

490

491 **Polygenic risk scoring**

492 To calculate the explained variance in insomnia by our GWAS results, we calculated
493 polygenic scores (PGS) based on the SNP effect sizes in the meta-analysis. The PGS were
494 calculated using two methods: LDpred⁶⁵ and PRSice⁶⁶, a script for calculating P -value
495 thresholded PGS in PLINK. PGS were calculated using a leave-one-out method, where
496 summary statistics were recalculated each time with one sample of $N=3,000$ from UKB
497 excluded from the analysis. This sample was then used as a target sample for estimating the
498 explained variance in insomnia by the PGS.

499

500 **Mendelian Randomization**

501 To investigate causal associations between insomnia and genetically correlated traits, we
502 analyzed direction of effects using Generalized Summary-data based Mendelian
503 Randomization (GSMR⁴³; <http://cnsgenomics.com/software/gsmr/>). This method uses effect
504 sizes from GWAS summary statistics (standardized betas or log-transformed odds ratios) to
505 infer causality of effects between two traits based on genome-wide significant SNPs. Built-in
506 HEIDI outlier detection was applied to remove SNPs with pleiotropic effects on both traits,
507 as these may bias the results. We tested for causal associations between insomnia and traits
508 that were significantly genetically correlated with insomnia (b_{zx}). In addition, we tested for
509 bi-directional associations by using other traits as the determinant and insomnia as the
510 outcome (b_{zy}). We selected independent ($r^2 < 0.1$) lead SNPs with a GWS P -value ($< 5 \times 10^{-8}$) as
511 instrumental variables in the analyses. For traits with less than 10 lead SNPs (i.e. the
512 minimum number of SNPs on which GSMR can perform a reliable analysis) we selected
513 independent SNPs ($r^2 < 0.1$), with a P -value $< 1 \times 10^{-5}$. If the outcome trait is binary, the
514 estimated b_{zx} and b_{zy} are approximately equal to the natural log of the odds ratio (OR). An OR
515 of 2 can be interpreted as a doubled risk compared to the population prevalence of a binary
516 trait for every SD increase in the exposure trait. For quantitative traits, the b_{zx} and b_{zy} can be
517 interpreted as a one standard deviation increase explained in the outcome trait for every SD
518 increase in the exposure trait.

519

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521

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683 Auton, Robert K. Bell, Katarzyna Bryc, Sarah L. Elson, Pierre Fontanillas, Nicholas A.
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686 Northover, Steven J. Pitts, J. Fah Sathirapongsasuti, Olga V. Sazonova, Janie F. Shelton,
687 Suyash Shringarpure, Chao Tian, and Catherine H. Wilson.

688

689 **Author contributions:** D.P. and E.J.W.V.S. conceived the idea of the study. D.P. supervised
690 the pre- and post gwas analysis pipeline. P.R.J. and K.W. performed the analyses. S.St.
691 performed quality control on the UK Biobank data and wrote the analysis pipeline. K.W.
692 wrote the online platform (FUMA) that was used for follow-up analyses. C.d.L conducted
693 conditional gene-set analyses. J.B., N.S., A.M.M. and J.H.L contributed scRNA information.
694 J.T., D.H., V.V. and the 23andMe Research Team contributed and analyzed the 23andMe

695 cohort data. D.P., E.J.W.V.S. and P.R.J. wrote the paper. All authors discussed the results,
696 and approved the final version of the paper.

697

698 **Materials & Correspondence:** The data analyzed in the current study was partly provided
699 by the UK Biobank Study (www.ukbiobank.ac.uk), received under the UK Biobank
700 application number 16406. Our policy is to make genome-wide summary statistics (sumstats)
701 publicly available. Sumstats from the GWAS's conducted are available for download at
702 <https://ctg.cncr.nl/>. Note that our freely available meta-analytic sumstats (insomnia and
703 morningness) concern results excluding the 23andMe sample. This is a non-negotiable clause
704 in the 23andMe data transfer agreement, intended to protect the privacy of the 23andMe
705 research participants. To fully recreate our meta-analytic results for insomnia and
706 morningness: (a) obtain insomnia and morningness sumstats from 23andMe (see below); (b)
707 conduct a meta-analysis of our sumstats with the 23andMe sumstats. 23andMe participant
708 data are shared according to community standards that have been developed to protect against
709 breaches of privacy. Currently, these standards allow for the sharing of summary statistics for
710 at most 10,000 SNPs. The full set of summary statistics can be made available to qualified
711 investigators who enter into an agreement with 23andMe that protects participant
712 confidentiality. Interested investigators should email dataset-request@23andme.com for more
713 information.

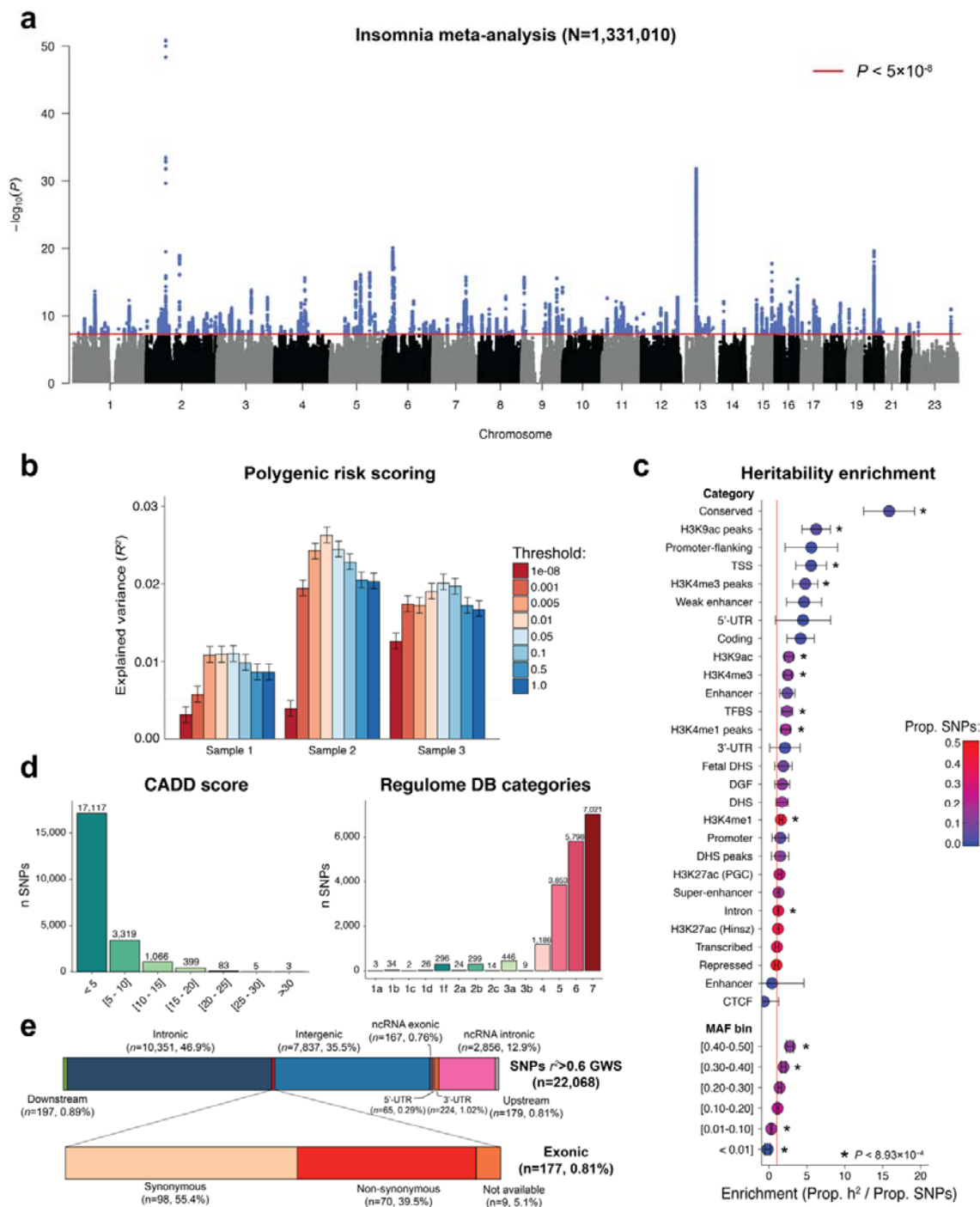
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715 **Author Information:** V.V., D.H., and J.T. are employees of 23andMe. All other authors
716 declare no competing financial interest. Correspondence and requests for materials should be
717 addressed to d.posthuma@vu.nl.

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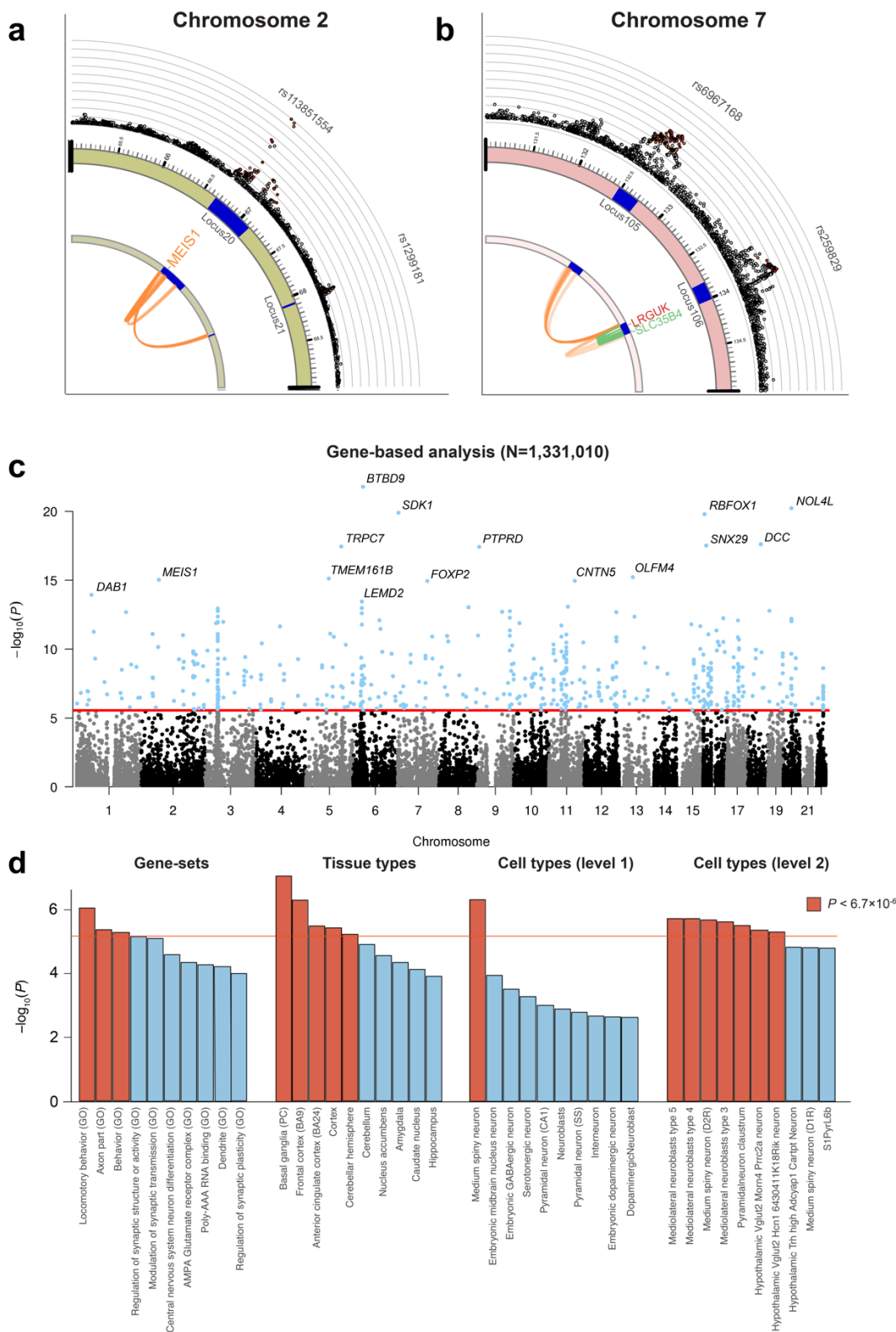
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720 FIGURES
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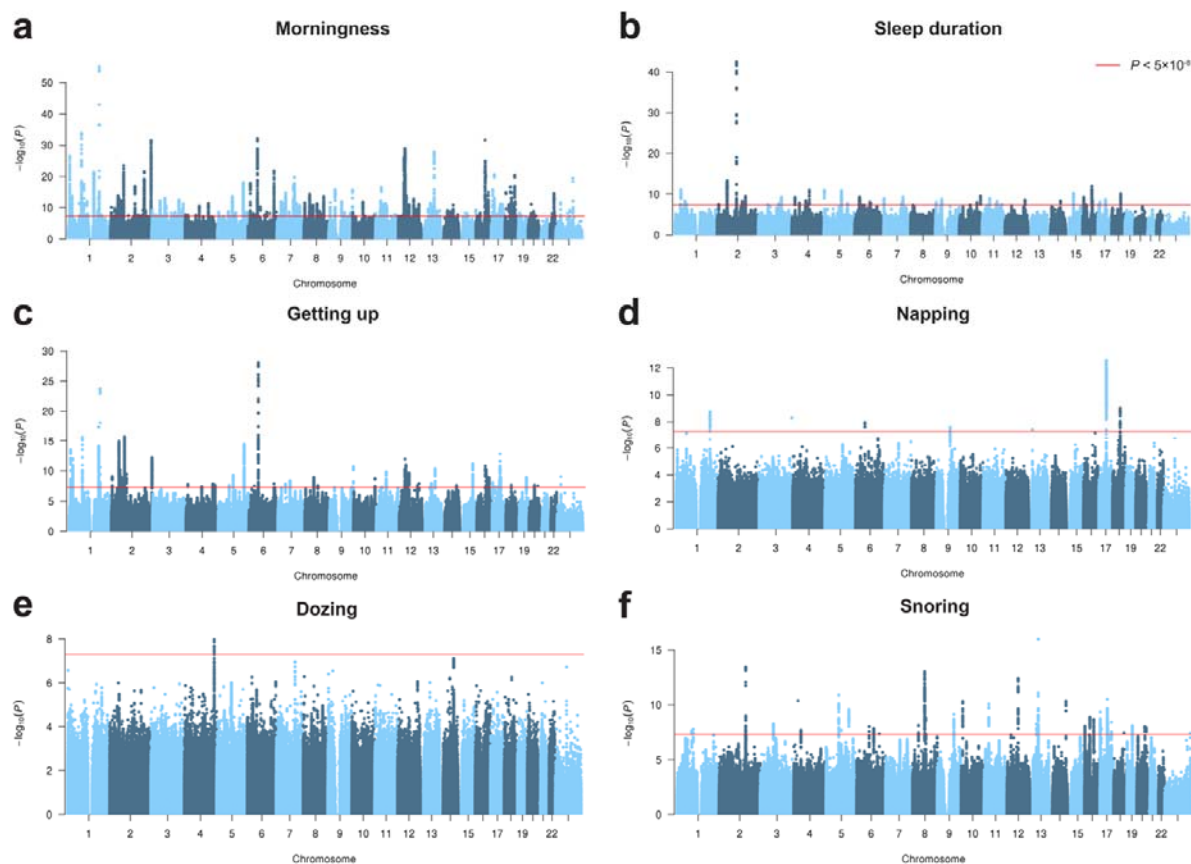
724 **Fig. 1a-e. SNP-based results from the GWAS meta-analysis on insomnia (N=1,331,010).** (a)
725 Manhattan plot of the GWAS of insomnia, showing the $-\log_{10}$ -transformed P -value for each SNP (b)
726 Heritability enrichment for functional SNP categories and minor allele frequency bins (MAF).
727 Enrichment was calculated by dividing the proportion of heritability for each category by the
728 proportion of SNPs in that category, significant enrichments after Bonferroni correction (28 functional
729 categories + 6 MAF bins + 22 chromosomes) are indicated by an asterisk ($P < 0.05/56 = 8.93 \times 10^{-4}$) (c)
730 Polygenic score (PGS) prediction in three hold-out samples (N=3,000), showing the increase in
731 explained variance in insomnia (Nagelkerke's pseudo R^2) and 95% confidence interval for each P -
732 value threshold. All P -value thresholds were statistically significant. (d) Distribution of CADD scores
733 and RegulomeDB category of all annotated SNPs in LD ($r^2 \geq 0.6$) with one of the GWS SNPs
734 ($n=22,068$) and (e) functional consequences of these SNPs.
735



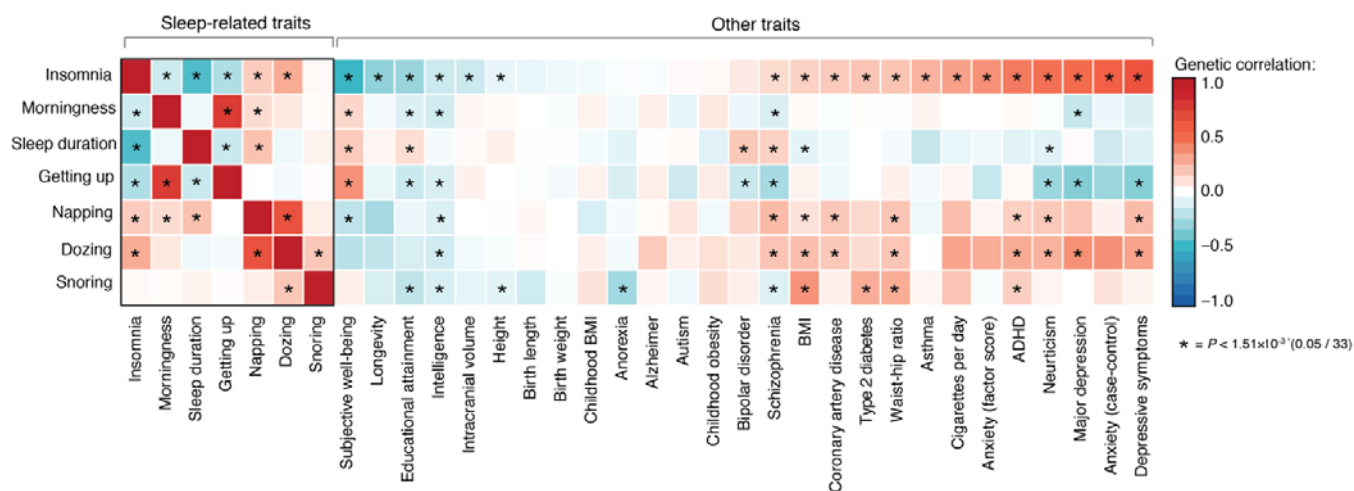
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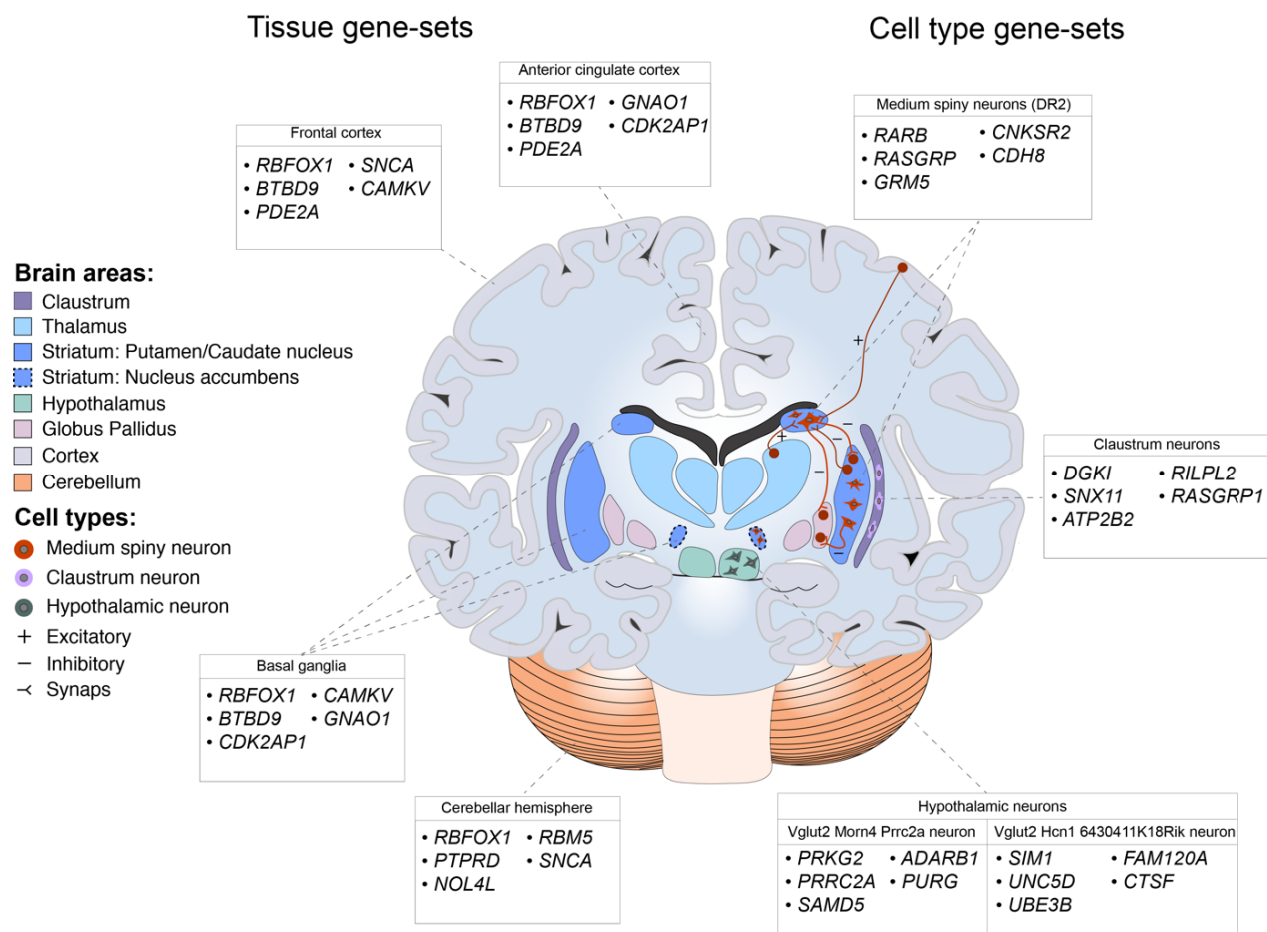
738 **Fig. 2a-d. Gene-based and gene-set analyses of insomnia.** Zoomed-in circos plots showing genes
739 implicated by two genomic risk loci on chromosome 2 **(a)** and chromosome 7 **(b)**, genomic risk loci
740 indicated as blue areas, eQTL associations in green, chromatin interactions in orange. Genes mapped
741 by both eQTL and chromatin interactions are red. The outer layer shows a Manhattan plot
742 containing the negative log₁₀-transformed *P*-value of each SNP in the GWAS meta-analysis of
743 insomnia. Full circos plots of all autosomal chromosomes are provided in **Supplementary Fig. 2.** **(c)**
744 Genome-wide gene-based analysis (GWAS) of 18,185 genes that were tested for association with
745 insomnia in MAGMA. The y-axis shows the negative log₁₀-transformed *P*-value of the gene-based
746 test, the x-axis shows the starting position on the chromosome. The red line indicates the Bonferroni
747 corrected threshold for genome-wide significance ($P=0.05/18,185=2.75\times 10^{-6}$). The top 15 most
748 significant genes are highlighted. **(d)** Gene-set analysis of top 20 for each of the MsigDB pathways,
749 tissue expression of GTEx tissue types, and cell types from single-cell RNA sequencing. Gene-set
750 analyses were performed using MAGMA. The red line shows the Bonferroni significance threshold
751 ($P<0.05/7,473=6.7\times 10^{-6}$), correcting for the total number of tested gene-sets. Red bars indicated
752 significant gene-sets.



753 **Fig. 3a-f. Genome-wide analyses of six sleep-related traits.** Manhattan plots of the genome-wide
754 association analyses of (a) Morningness (N=434,835). (b) Sleep duration (N=384,317) (c) Ease of
755 getting up (N=385,949) (d) Napping (N=386,577) (e) Daytime dozing (N=385,333) and (f) Snoring
756 (N=359,916). The y-axis shows the negative log₁₀-transformed SNP *P*-value, the x-axis the base pair
757 position of the SNPs on each chromosome. The red line indicates the Bonferroni corrected
758 significance threshold ($P < 5 \times 10^{-8}$).



759 **Fig. 4. Genetic overlap of insomnia with other sleep-related traits and psychiatric and metabolic**
 760 **traits.** Heatmap of genetic correlations between insomnia, sleep-related phenotypes and
 761 neuropsychiatric and metabolic traits studies that were calculated using LD Score regression. Red
 762 color indicates a positive r_g while green indicates negative r_g . Correlations that were significant after
 763 Bonferroni correction ($P < 0.05/33 = 1.5110^{-3}$) are indicated with an asterisk (see also **Supplementary**
 764 **Table 18, 26**).



765 **Fig. 5. Overview of brain tissues and cell types associated with insomnia based on GWAS results**
 766 **from 1,331,010 individuals.** For each associated gene-set, the top 5 genes driving the association are
 767 reported. Results for GTEx brain tissue type gene-sets are shown on the left side of the figure, while
 768 results from the level 2 single-cell gene expression are shown on the right.

769 **EXTENDED DATA**

770

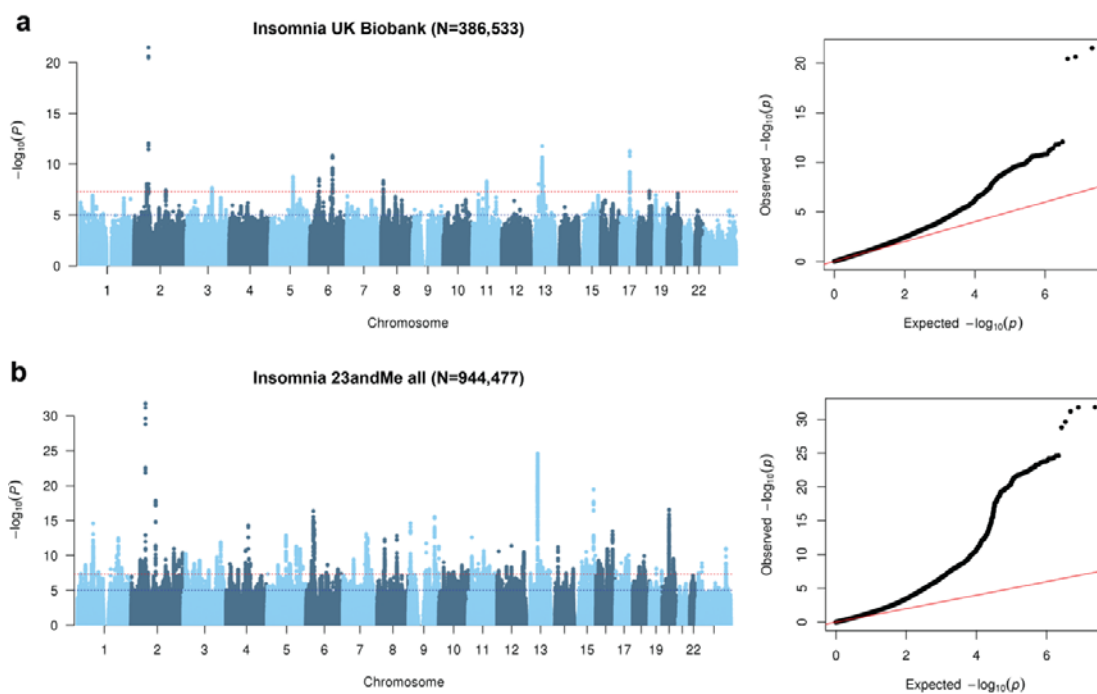
771 **Extended Data Table 1. LD Score regression estimates of the sex-specific GWAS of**
772 **insomnia.** Results are shown for UK Biobank, 23andMe and the sex-specific meta-analyzed
773 sample. H²=estimated SNP-heritability, intercept=LD Score regression intercept, rg=genetic
774 correlation in the same study sample.

775

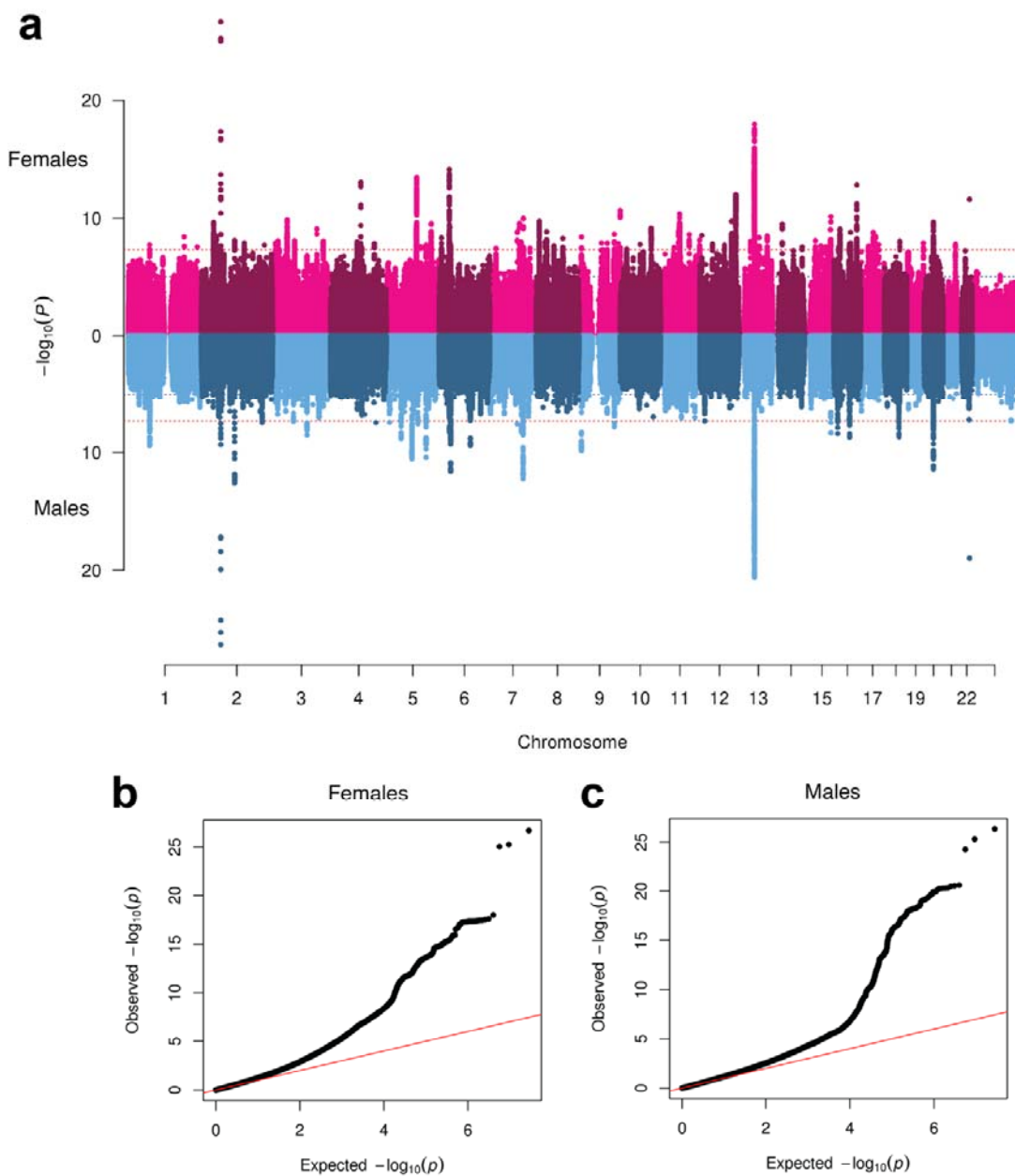
Sample	Sex	N	<i>h</i> ² (SE)	Mean chi ²	Lambda	Intercept (SE)	<i>rg</i> male	<i>rg</i> female
UK Biobank	male	177.817	0.083 (0.007)	1,157	1,143	1.001 (0.008)	1	0.857 (0.051)
	female	208.716	0.092 (0.005)	1,233	1,210	1.011 (0.008)	0.857 (0.051)	1
23andMe	male	443.207	0.080 (0.004)	1,385	1,317	1.016 (0.008)	1	0.925 (0.022)
	female	501.270	0.090 (0.003)	1,580	1,460	1.046 (0.009)	0.925 (0.022)	1
Meta	male	621.024	0.067 (0.003)	1,460	1,382	1.024 (0.009)	1	0.919 (0.018)
	female	709.986	0.078 (0.003)	1,700	1,547	1.042 (0.009)	0.919 (0.018)	1

776

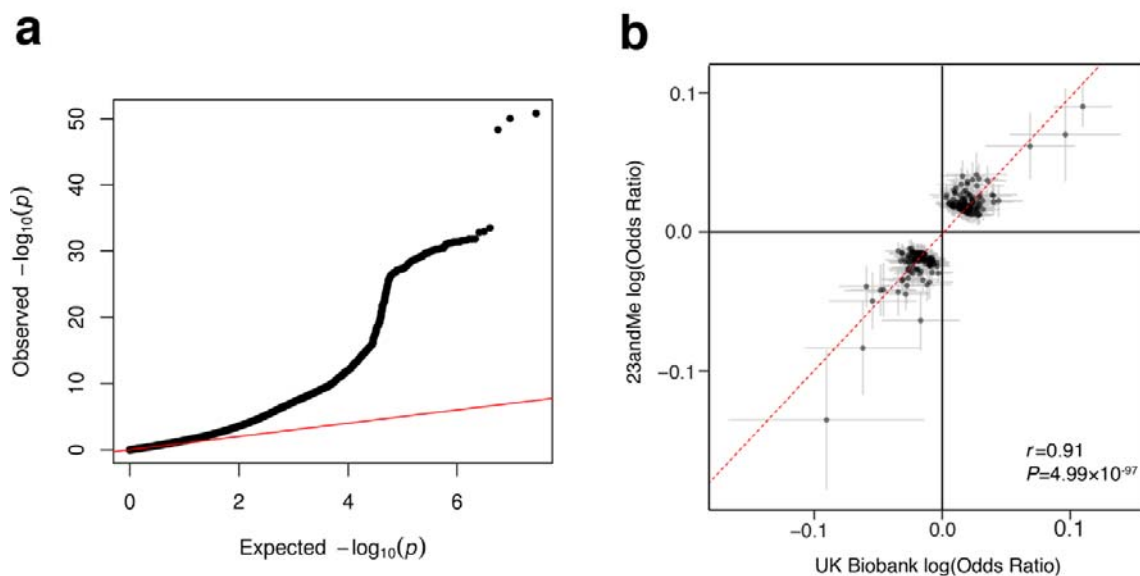
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778 **Extended Data Fig 1a-b. Manhattan and Q-Q plots of the genome-wide analysis of**
779 **insomnia.** Results are shown for the genome-wide analysis in **(a)** UK Biobank and **(b)**
780 23andMe.
781

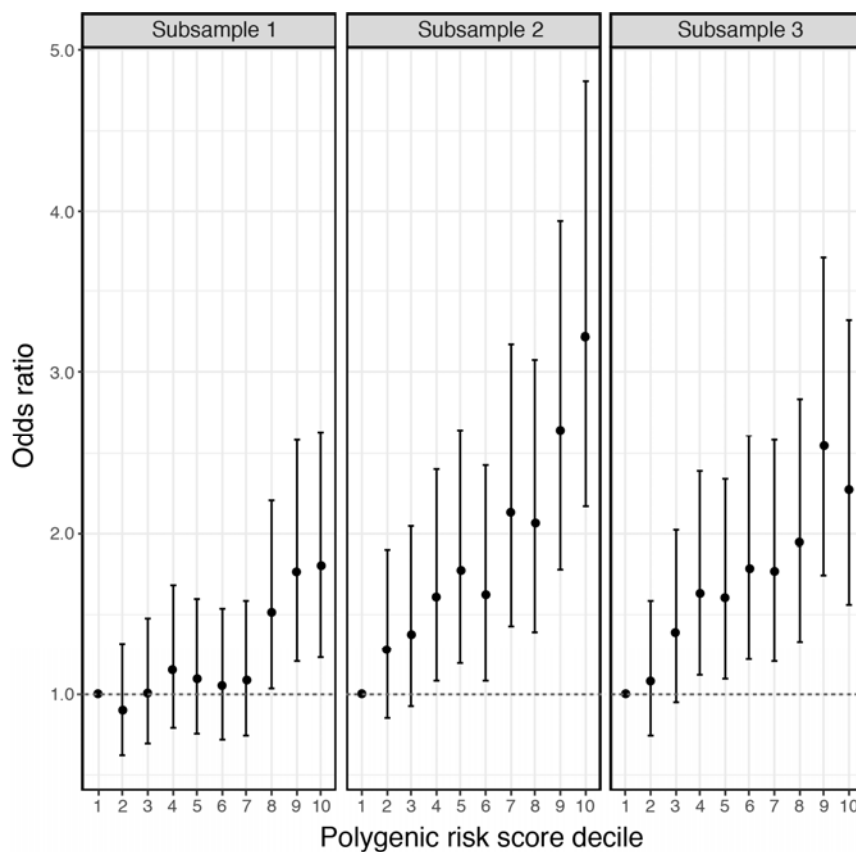


782 **Extended Data Fig. 3a-c. Sex-specific Manhattan plot and Q-Q plot of the insomnia**
783 **meta-analysis in males and females (UK Biobank + 23andMe).** (a) Miami plot showing
784 sex-specific SNP association P-values for females on the upper side and males on the lower
785 side. (b) Q-Q plot in females, and (c) in males.



786 **Extended Data Fig. 4a-b. Q-Q plot and lead SNPs of the GWAS meta-analysis for**
787 **insomnia.** (a) QQ-plot of the insomnia meta-analysis showing the expected negative log10-
788 transformed P -value distribution on the x-axis, and observed negative log10-transformed P -
789 value on the y-axis, (b) effect size plot of the 248 lead SNP of the insomnia meta-analysis
790 (log-transformed odds ratio and 95% confidence interval) in UK Biobank and 23andMe.
791

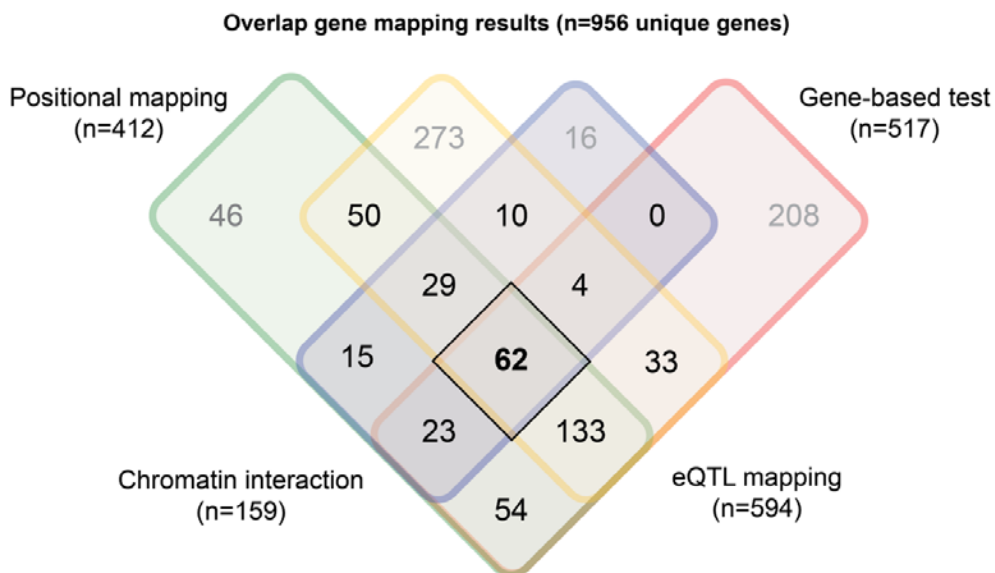
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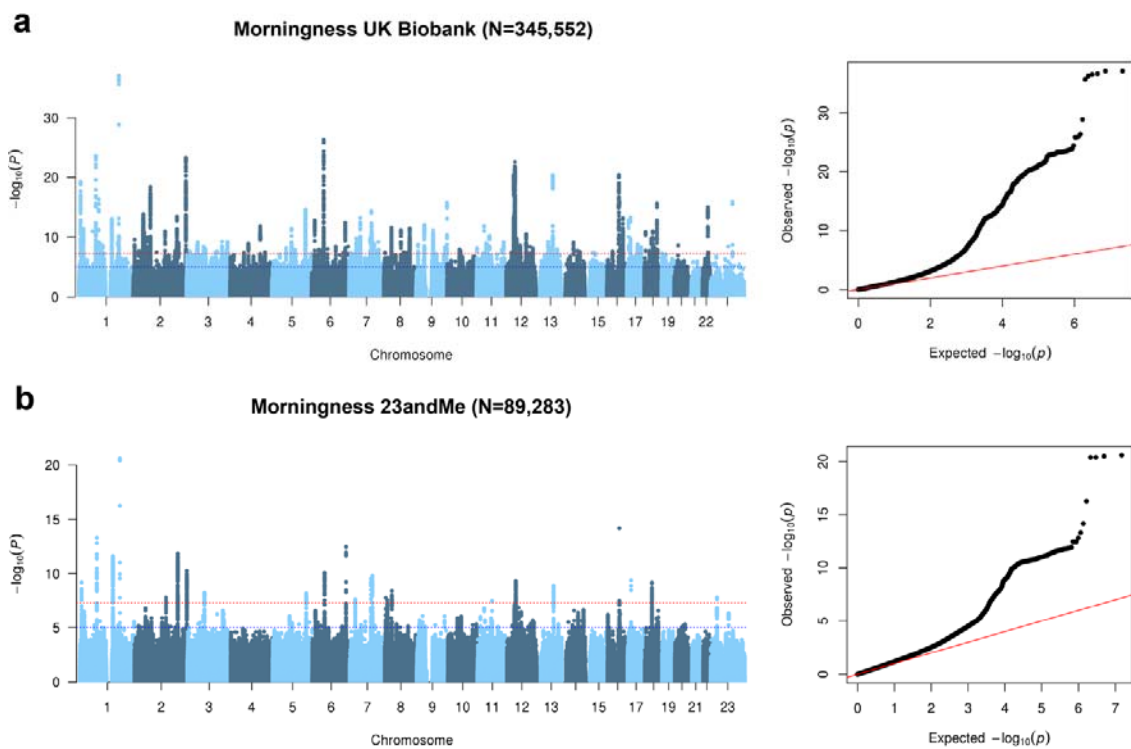
805 **Extended Data Fig. 5. Risk of insomnia per polygenic risk score decile in three**
806 **independent holdout samples (N=3x3000).** Odds ratios and 95% confidence interval for
807 deciles in polygenic risk score were calculated based on a logistic regression model, using the
808 lowest polygenic risk score decile as the reference.

809

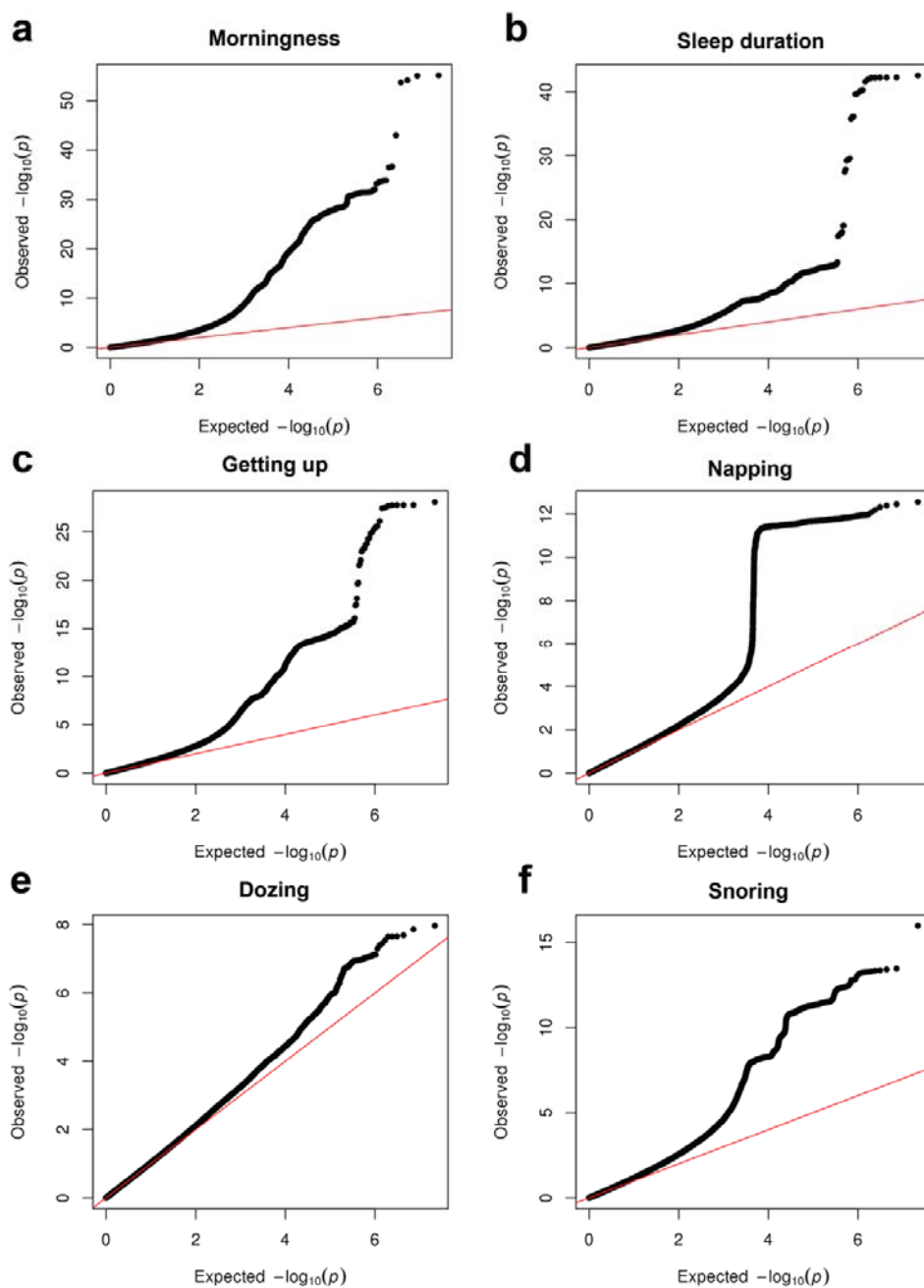
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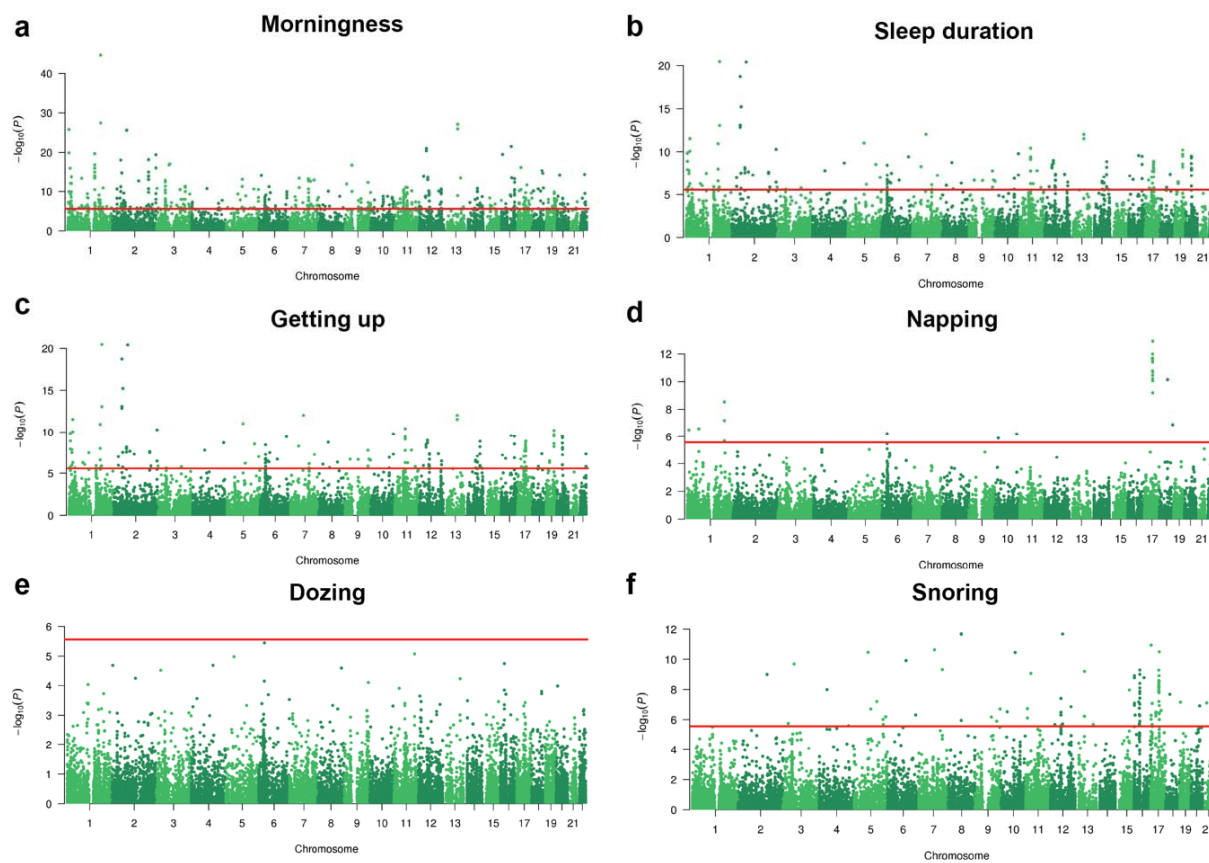
811 **Extended Data Fig. 6. Venn diagram showing the number of genes that were mapped by**
812 **four gene-mapping strategies.** Each square shows the number of overlapping genes between
813 three gene-mapping methods in FUMA (positional mapping, eQTL mapping and chromatin
814 interaction mapping) and significant genes in gene-based tests in MAGMA. The number of
815 genes in bold highlights the number of genes that were implicated by all four methods.
816



817 **Extended Data Fig. 7. Manhattan plot and Q-Q plot of the genome-wide analysis of**
818 **morningness in UK Biobank and 23andMe. Results are shown for (a) UK Biobank and (b)**
819 **23andMe.**
820



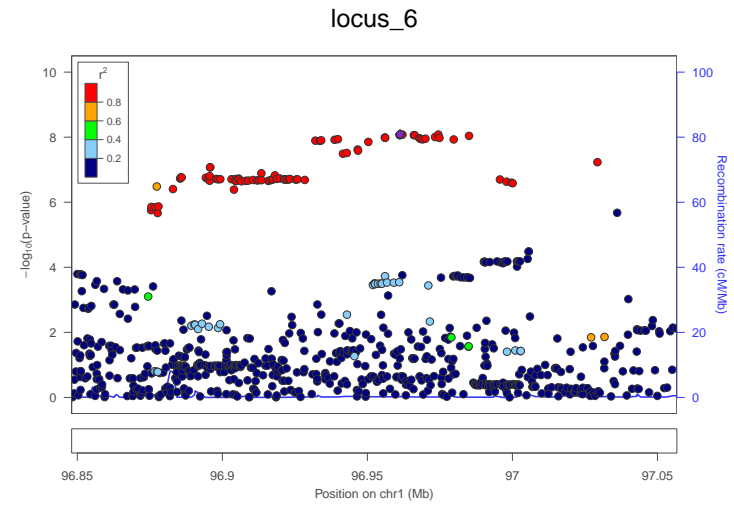
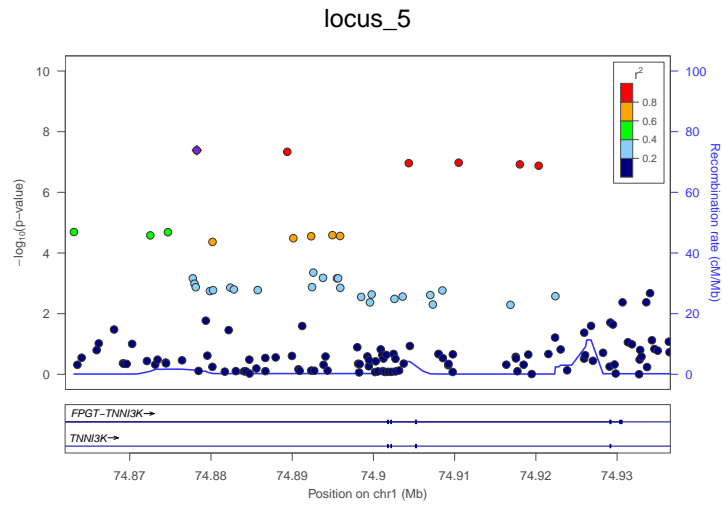
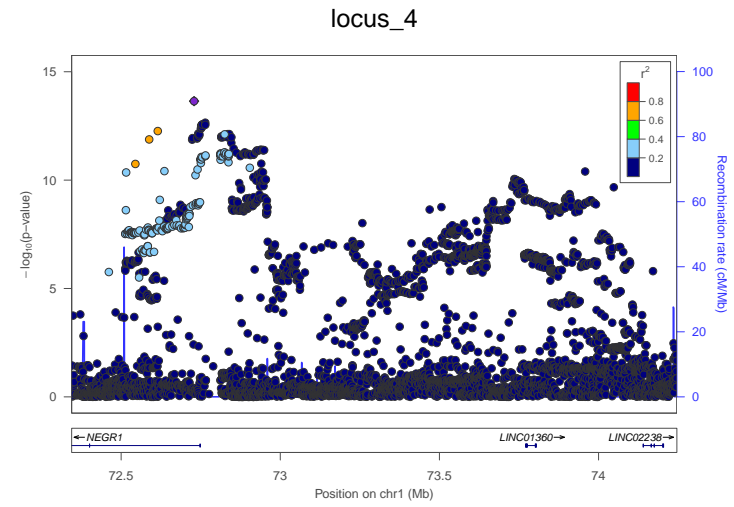
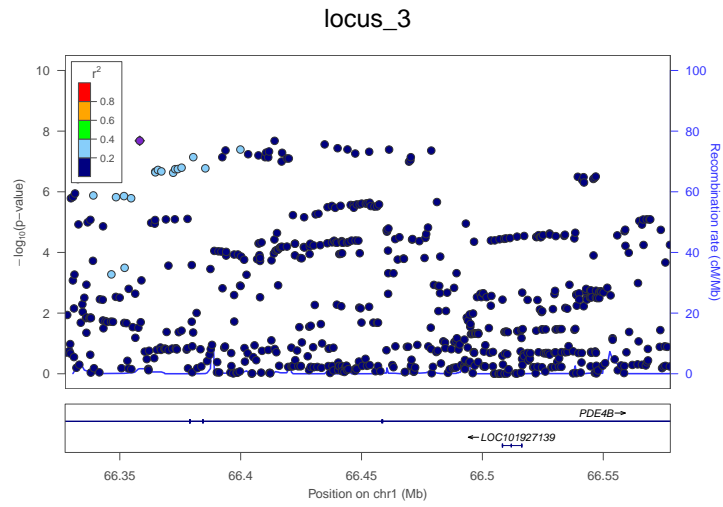
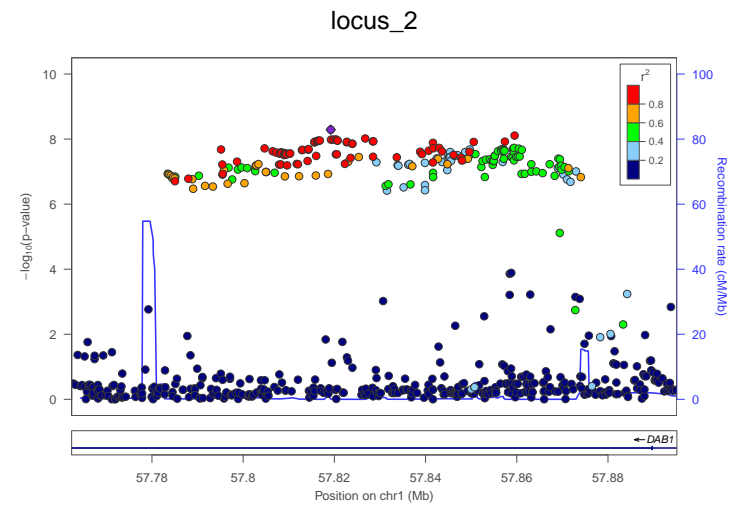
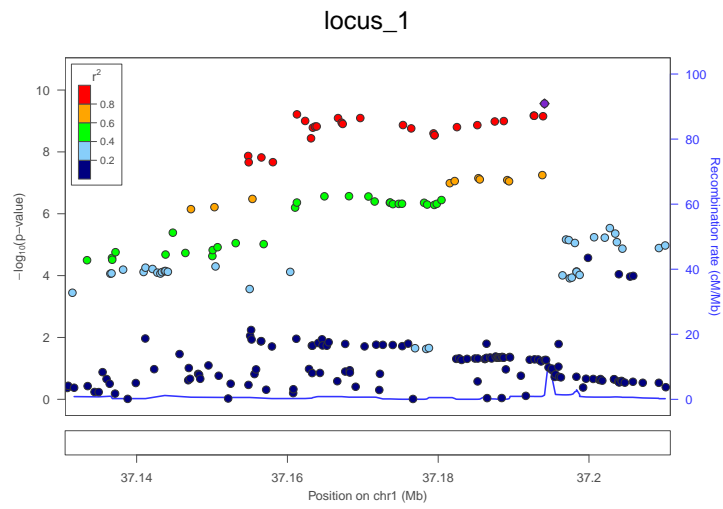
821 **Extended Data Fig. 8 Q-Q plots of the genome-wide analysis of six sleep related traits.**
822 **(a)** morningness (including UKB and 23andMe), **(b)** sleep duration, **(c)** ease of getting up, **(d)**
823 daytime napping, **(e)** daytime dozing, **(f)** snoring. Manhattan plots of the genome-wide
824 analyses are shown in **Fig. 3**.
825



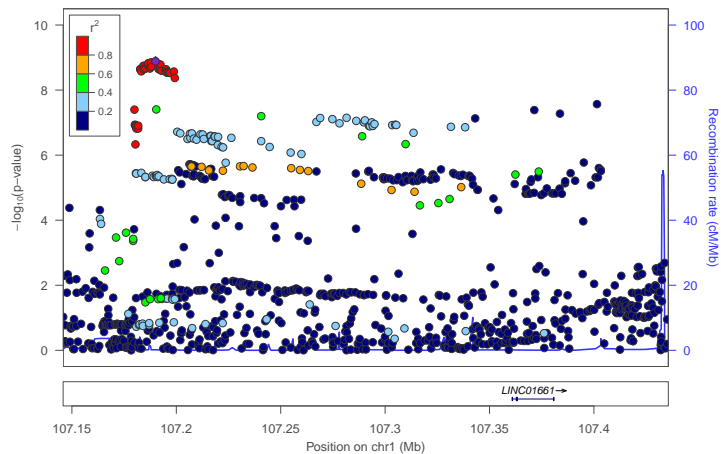
826 **Extended Data Fig. 9a-f. Genome-wide gene-based association analysis of six sleep-**
827 **related phenotypes.** Manhattan plots genome-wide gene-based analysis (GWGAS) results
828 for (a) morningness (b) sleep duration (c) ease of getting up (d) daytime napping (e) daytime
829 dozing (f) snoring. GWGAS was performed in MAGMA. The analysis of morningness was
830 based on GWAS meta-analysis of UKB and 23andMe, while other sleep-related phenotypes
831 were analysed in UKB. The red line indicates Bonferroni corrected significance threshold
832 depending on the number of genes tested.
833

Supplementary Information includes:

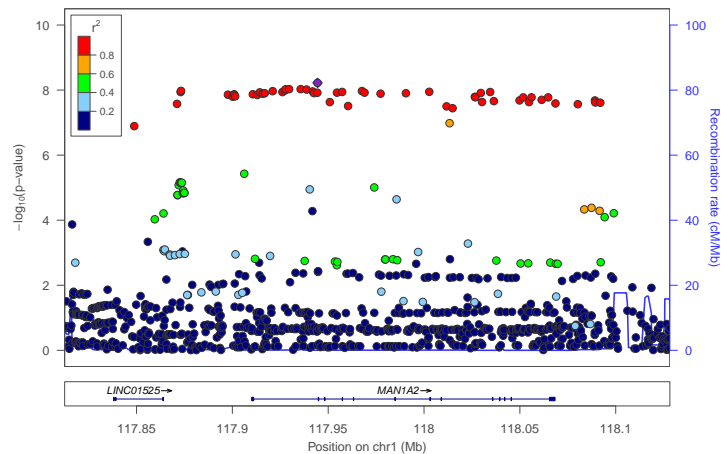
- 1. Supplementary Methods**
 - 1.1 Sample description UK Biobank
 - 1.2 Sample description 23andMe
 - 1.3 Insomnia phenotype validation external sample
- 2. Supplementary Discussion**
 - 2.1. Sex-specific association results for insomnia
 - 2.2. GWAS meta-analysis results for insomnia
 - 2.3. Implicated genes for insomnia
 - 2.4. Gene-set association results for insomnia
 - 2.5. Results sleep-related phenotypes
 - 2.6 Mendelian Randomization
- 3. Supplementary Figures (1 to 2)**
- 4. Supplementary Tables (1 to 28)**



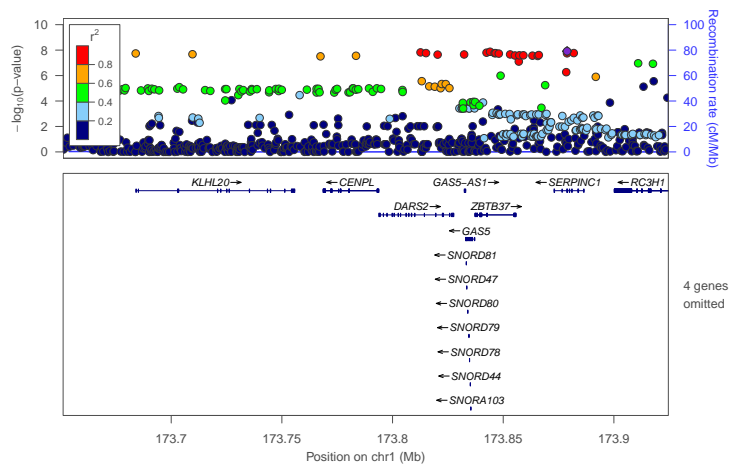
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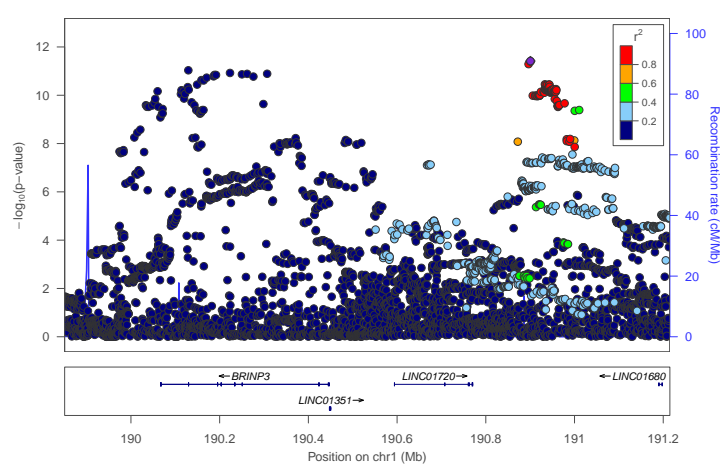
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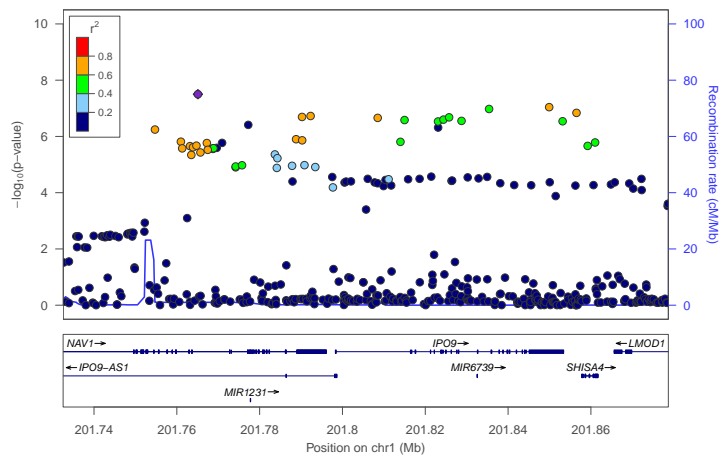
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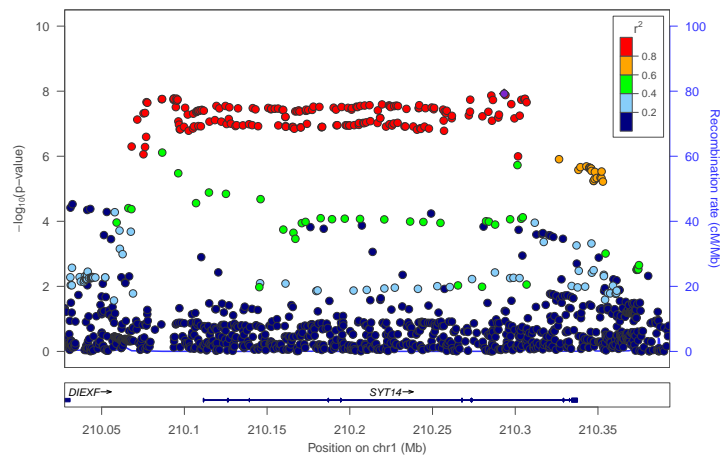
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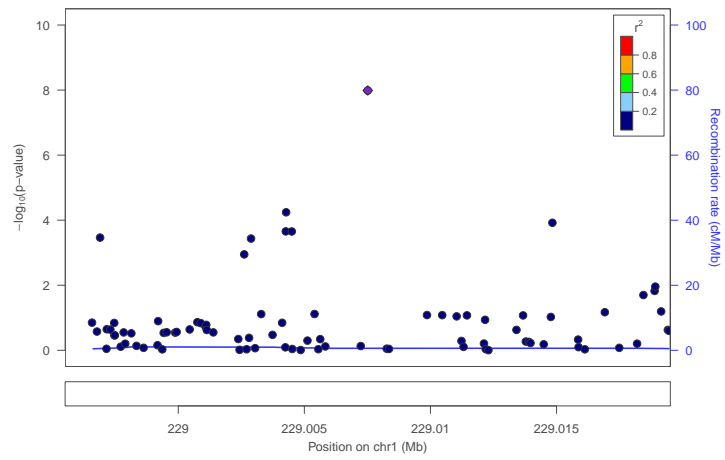
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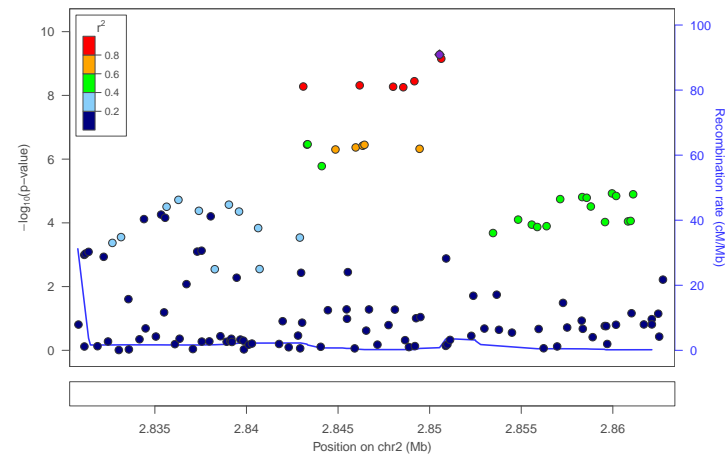
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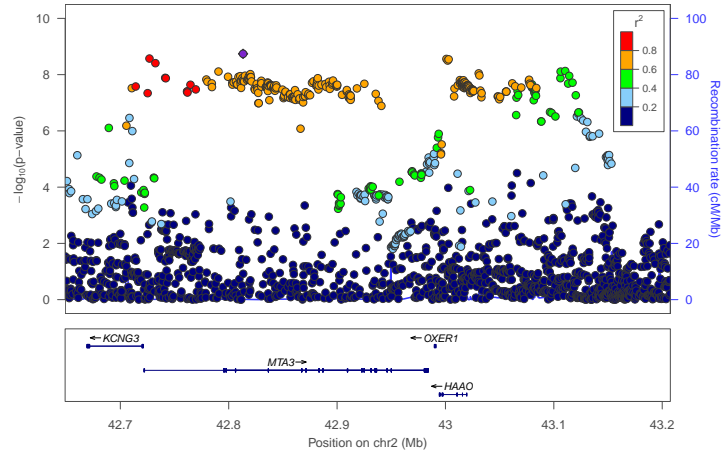
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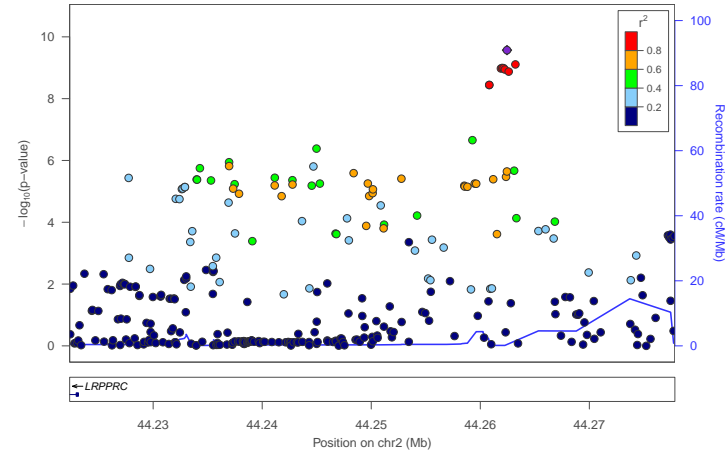
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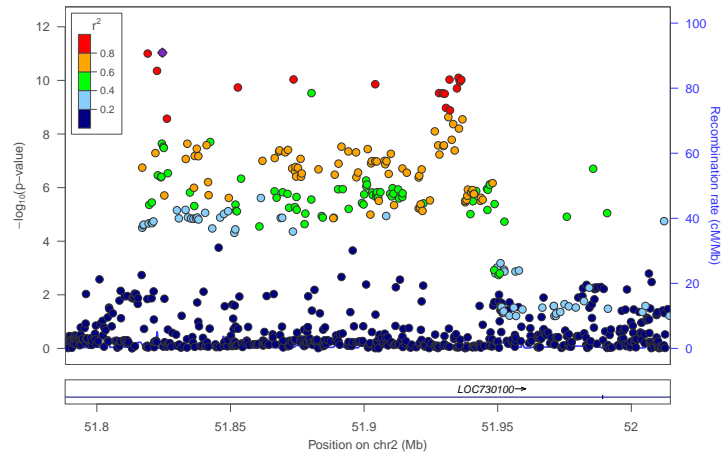
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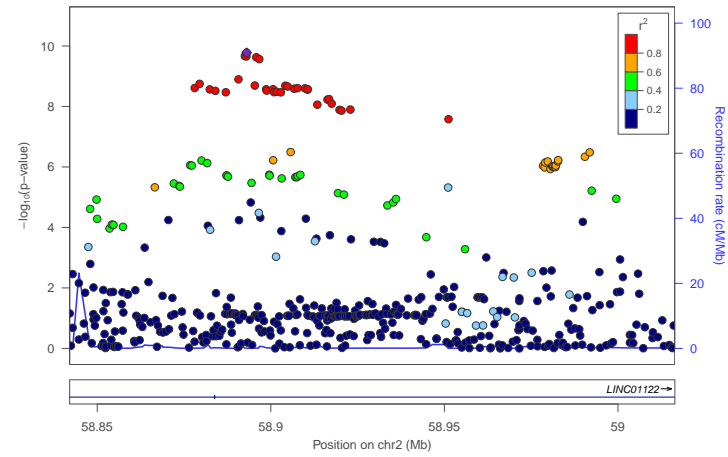
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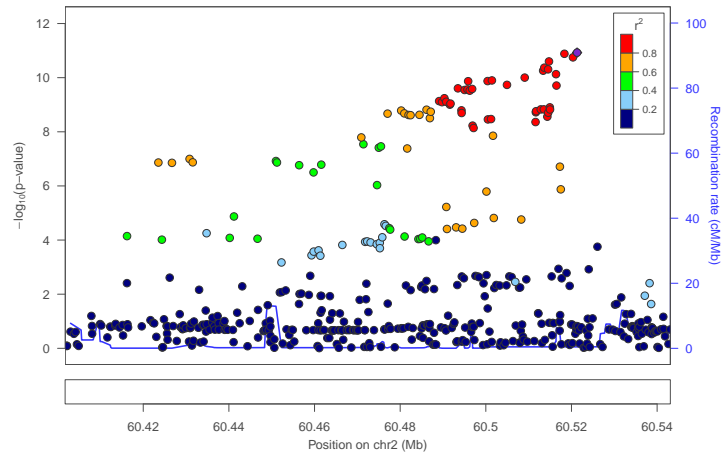
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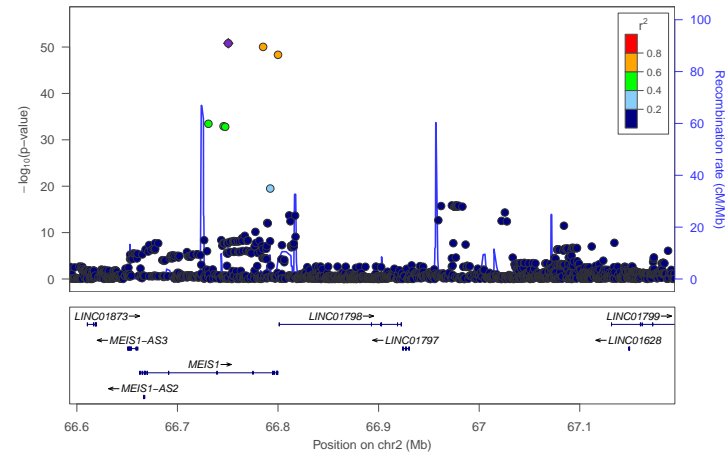
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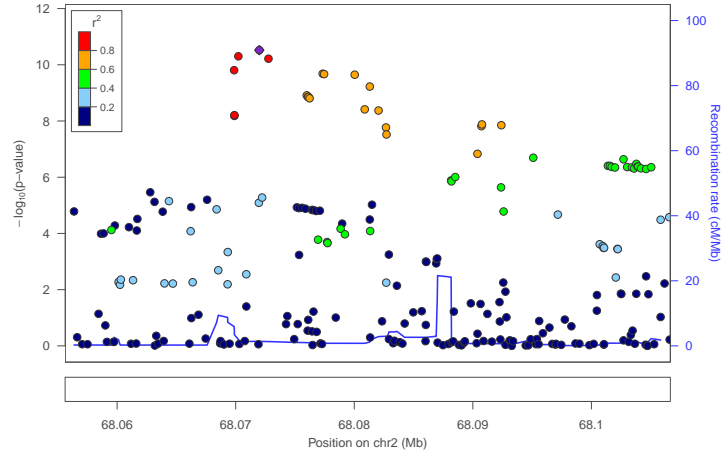
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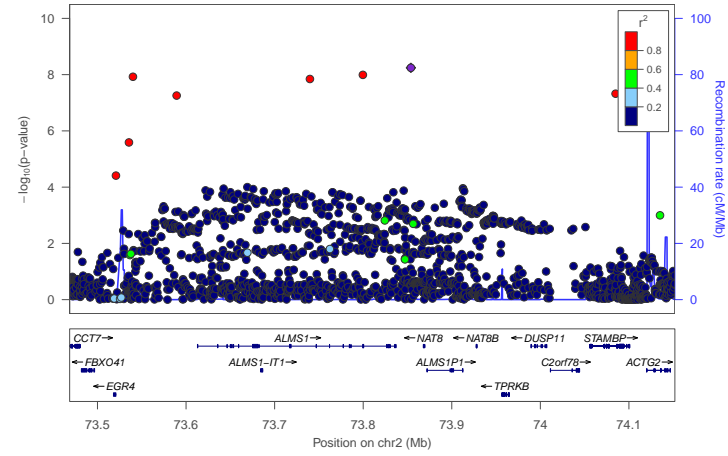
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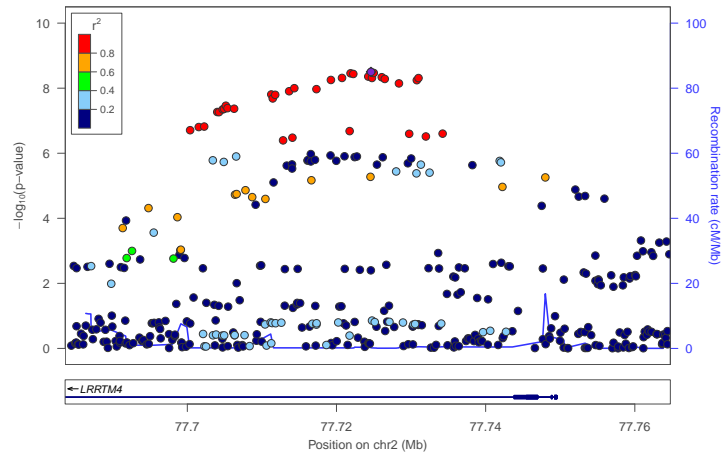
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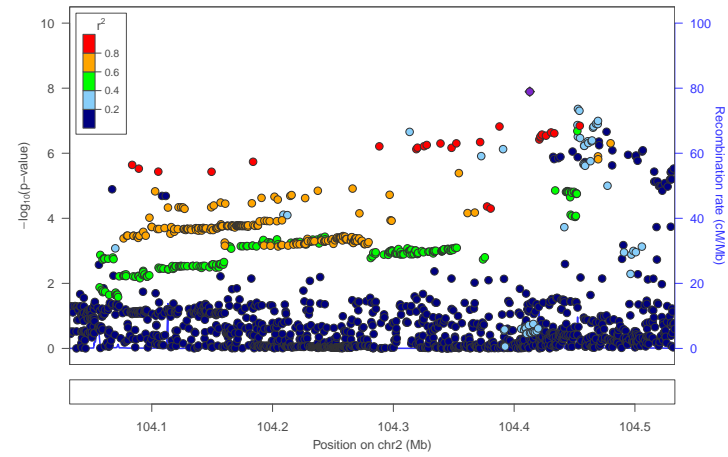
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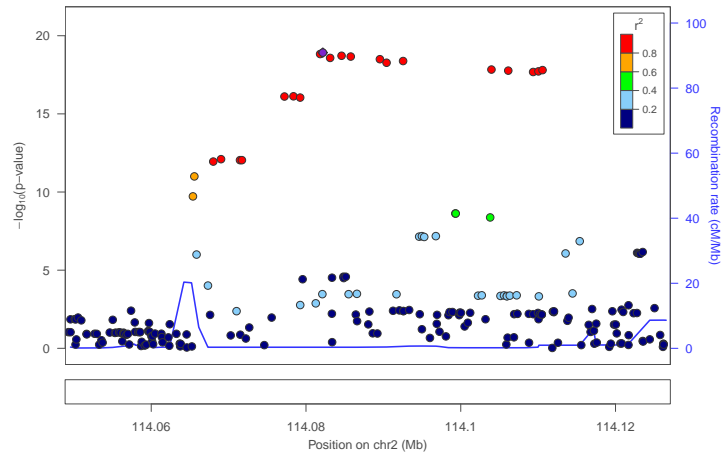
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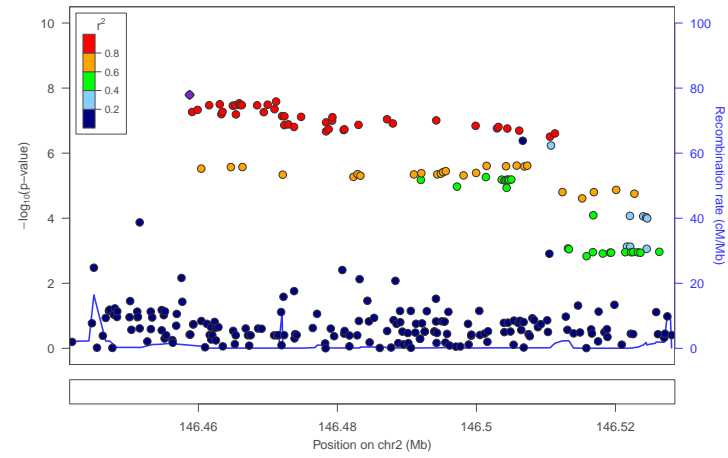
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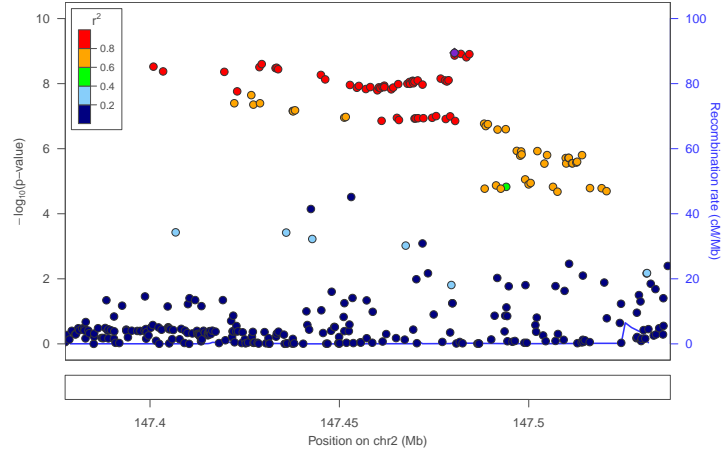
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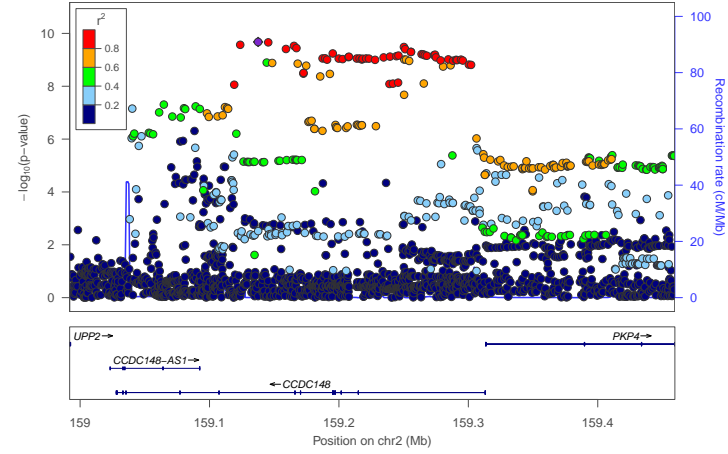
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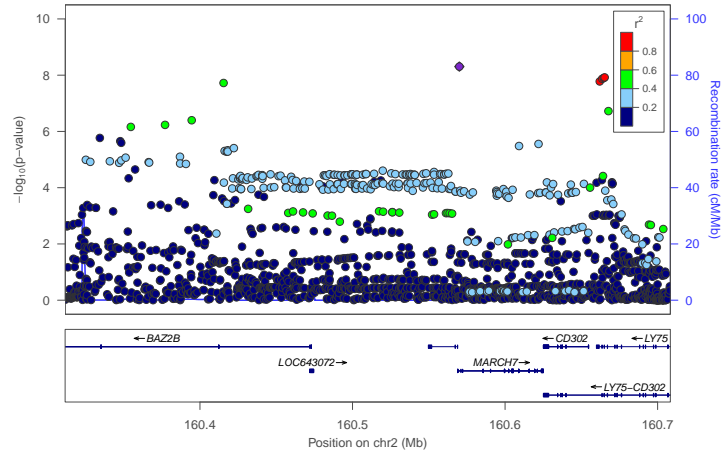
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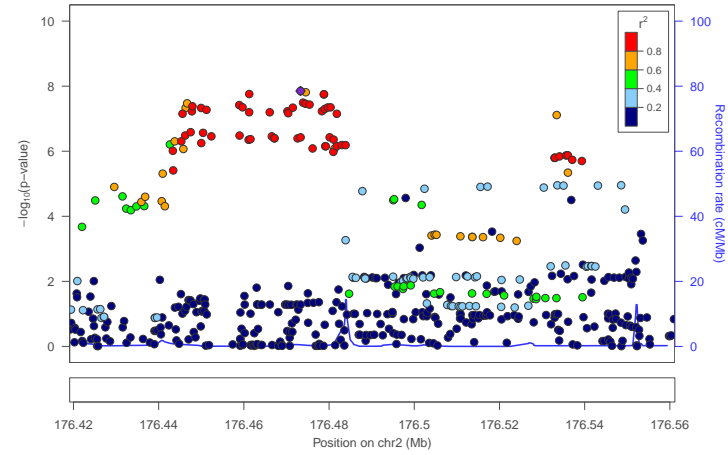
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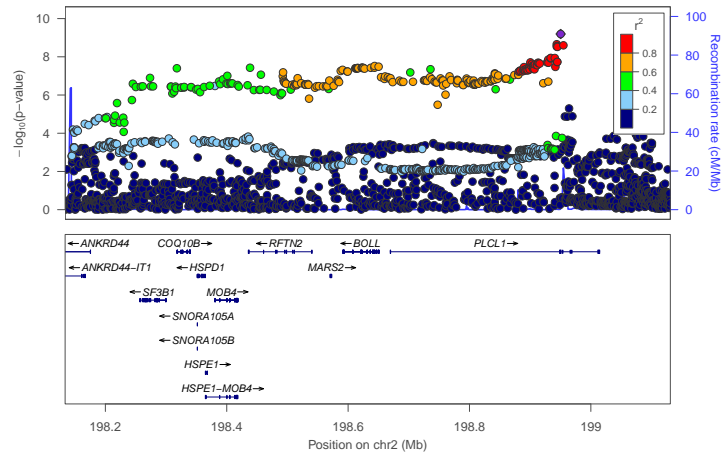
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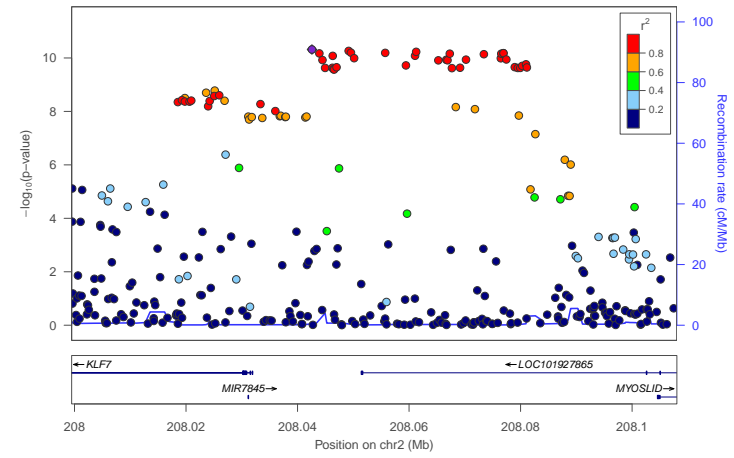
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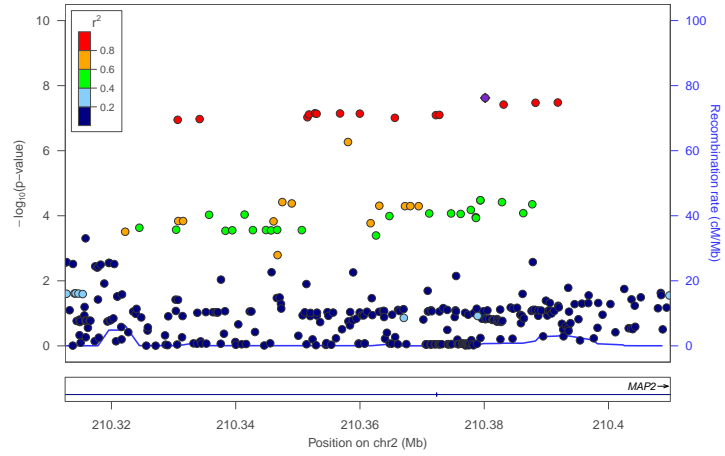
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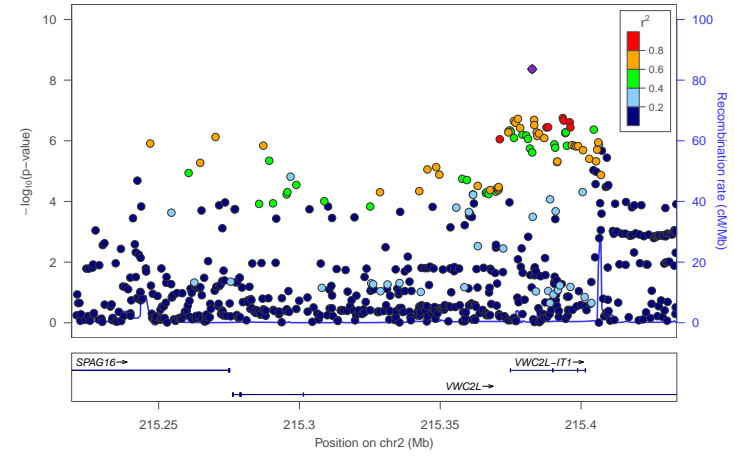
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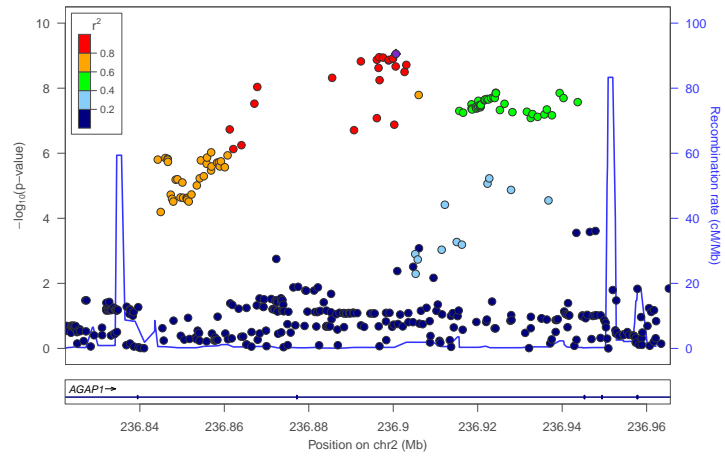
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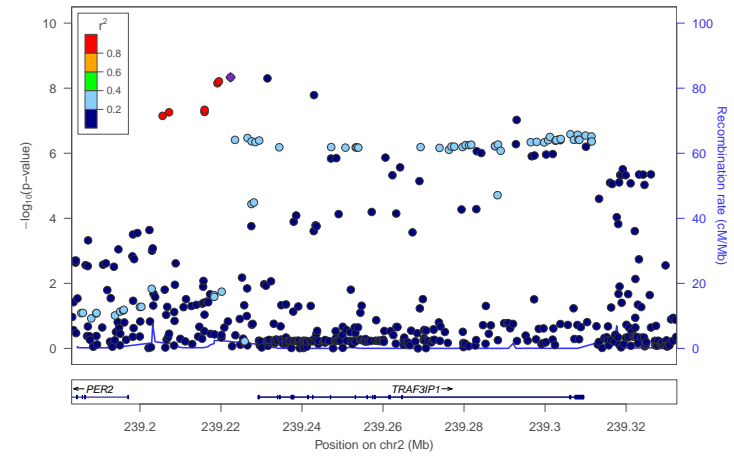
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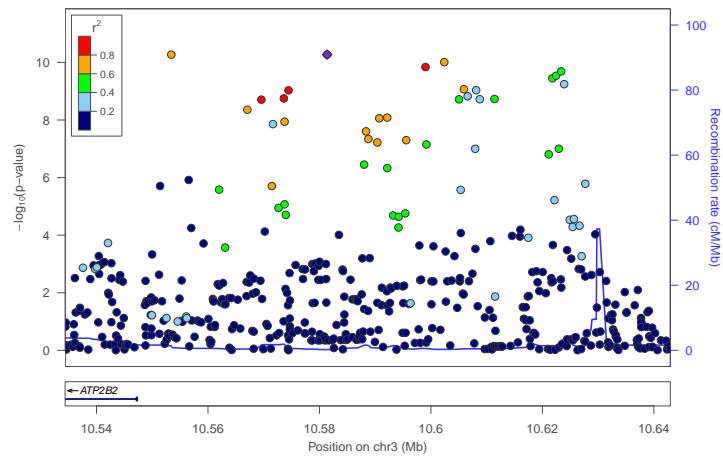
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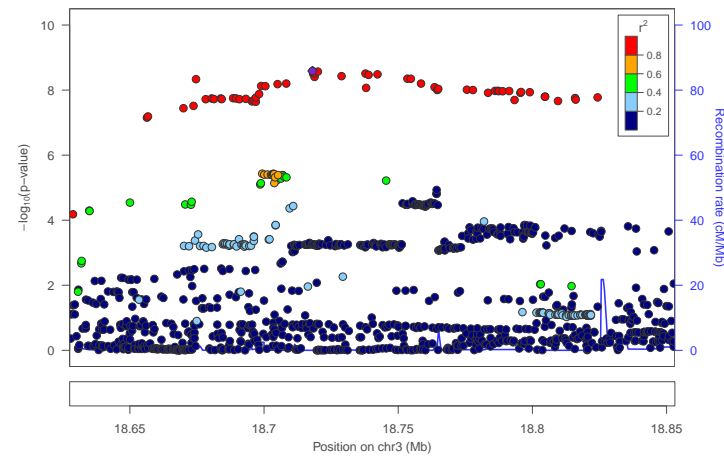
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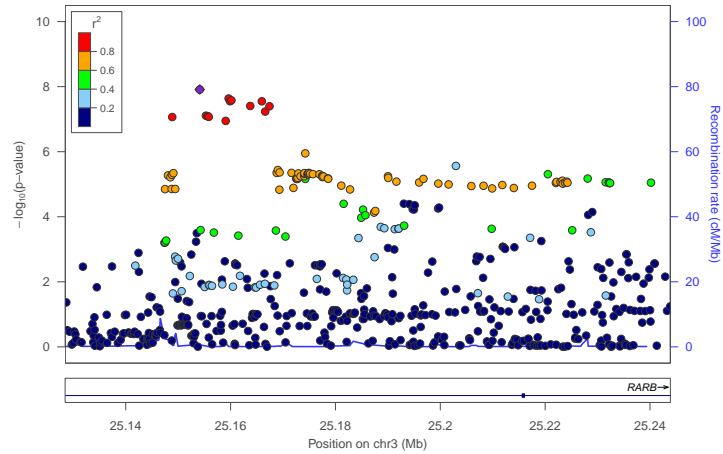
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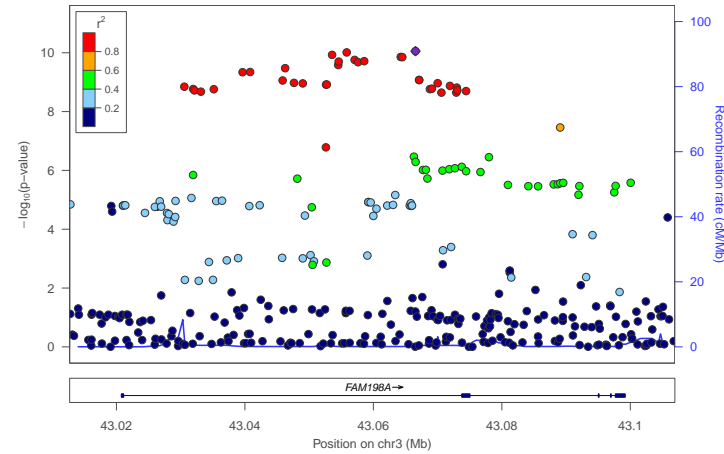
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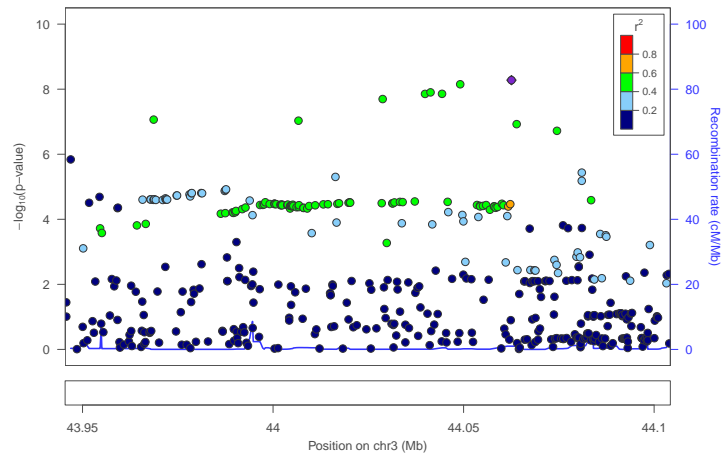
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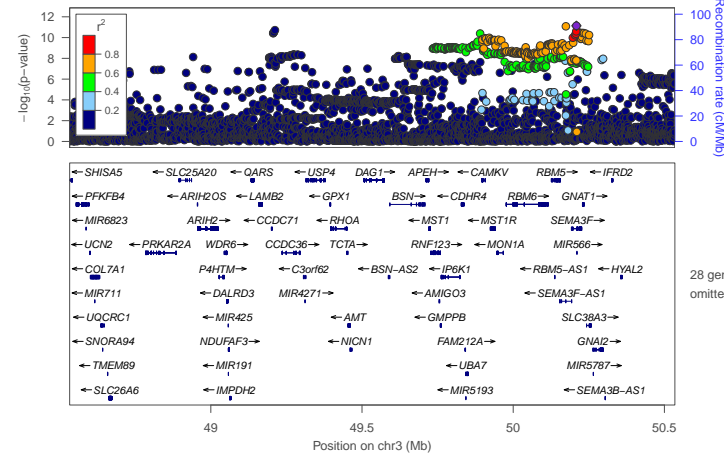
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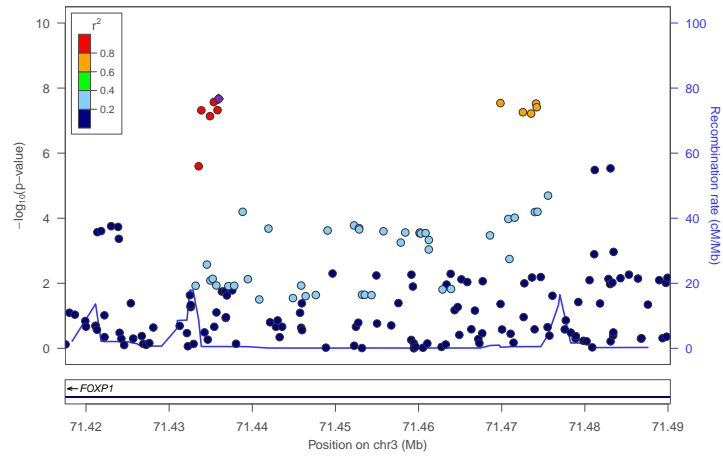
locus_41



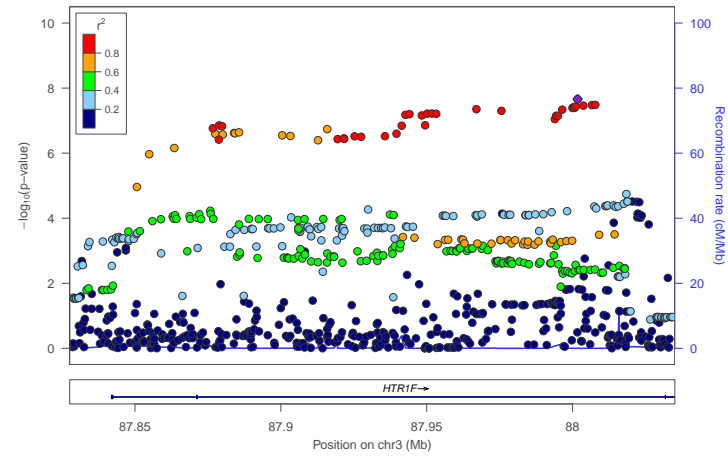
locus_42



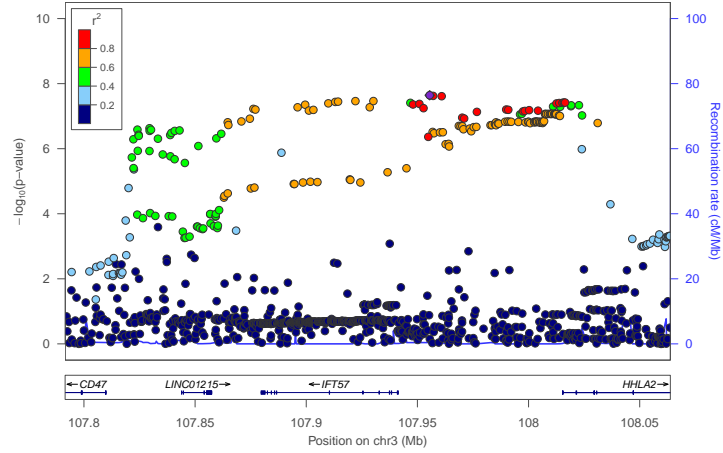
locus_43



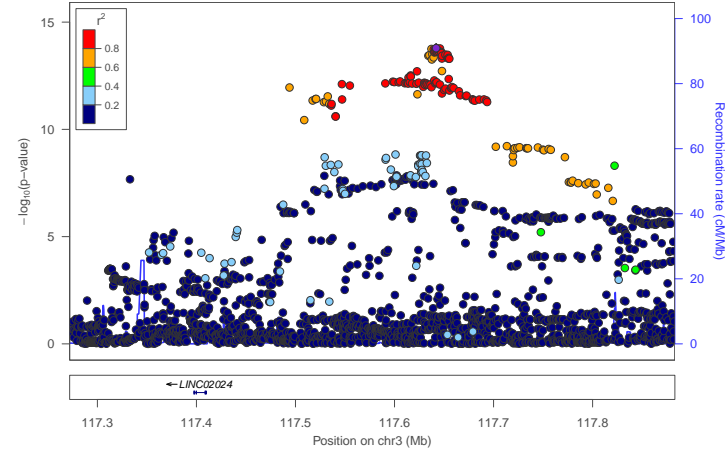
locus_44



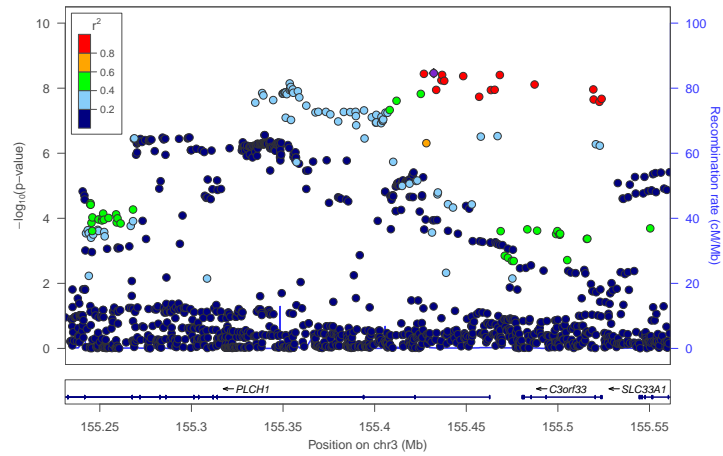
locus_45



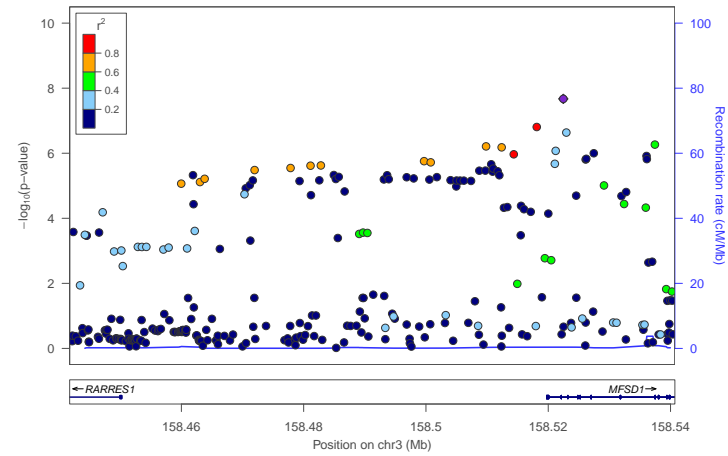
locus_46



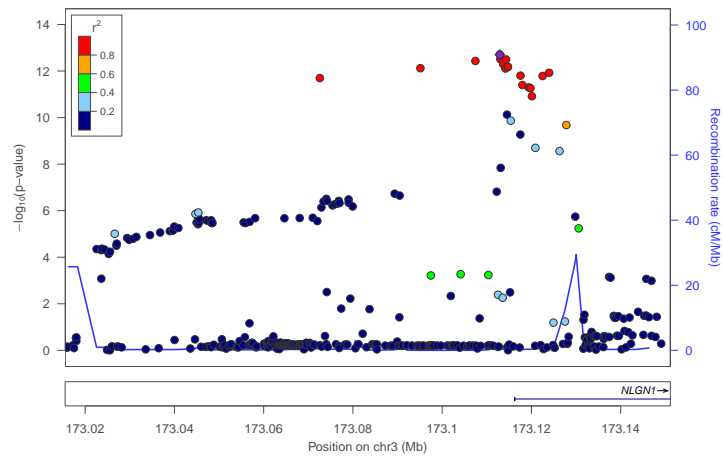
locus_47



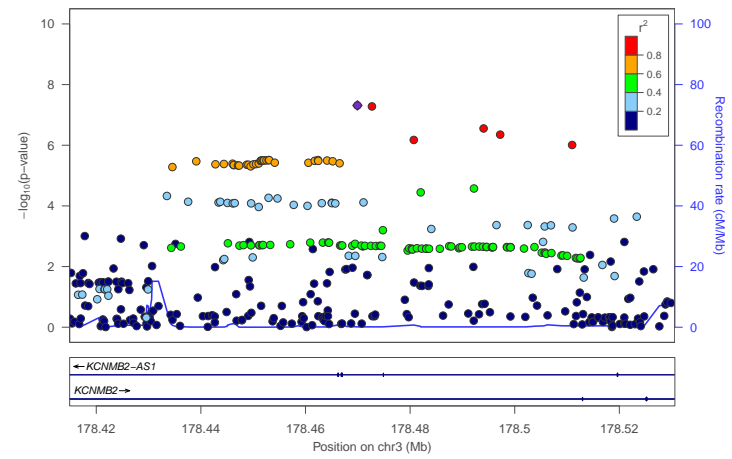
locus_48



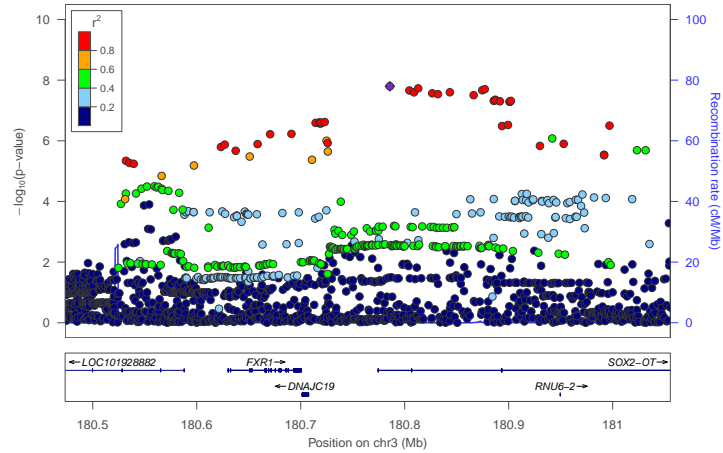
locus_49



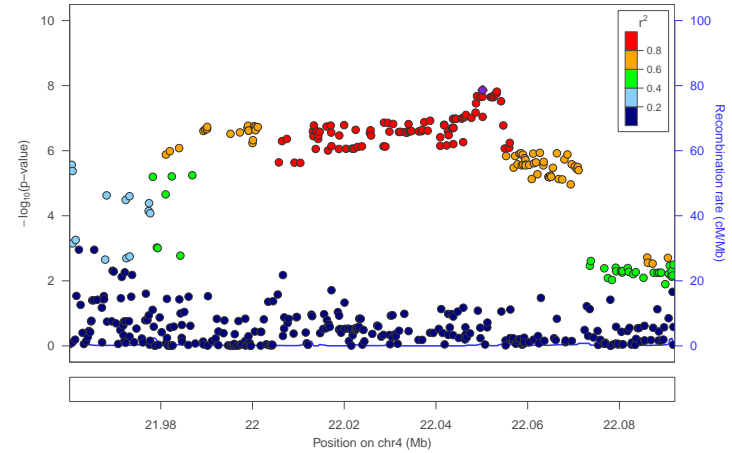
locus_50



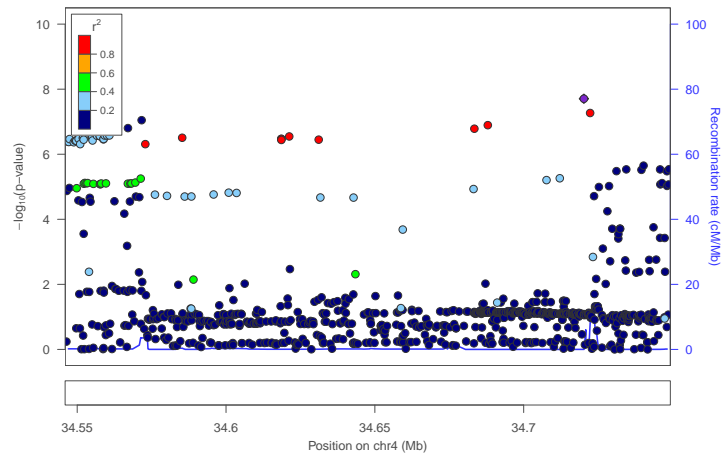
locus_51



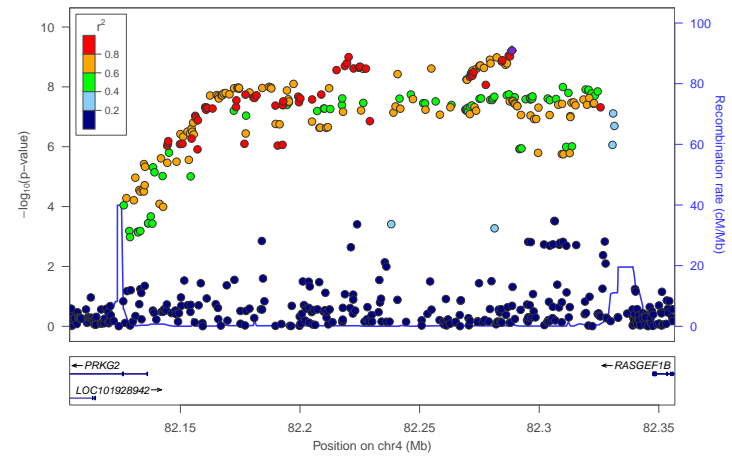
locus_52



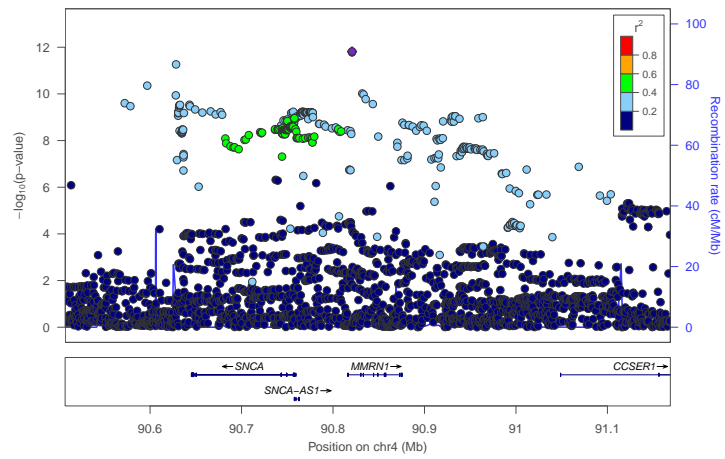
locus_53



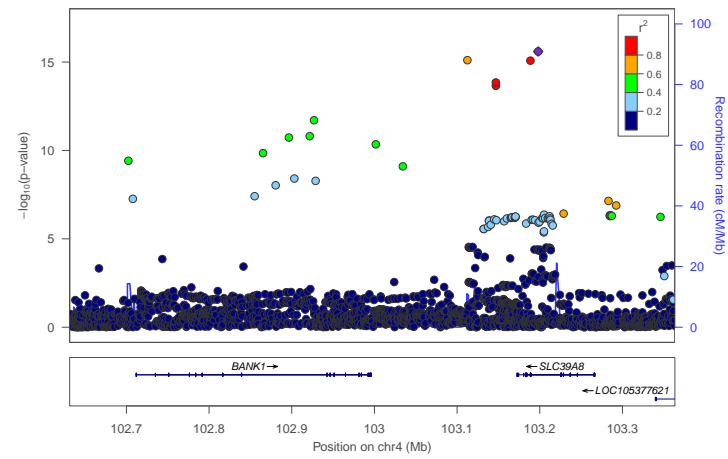
locus_54



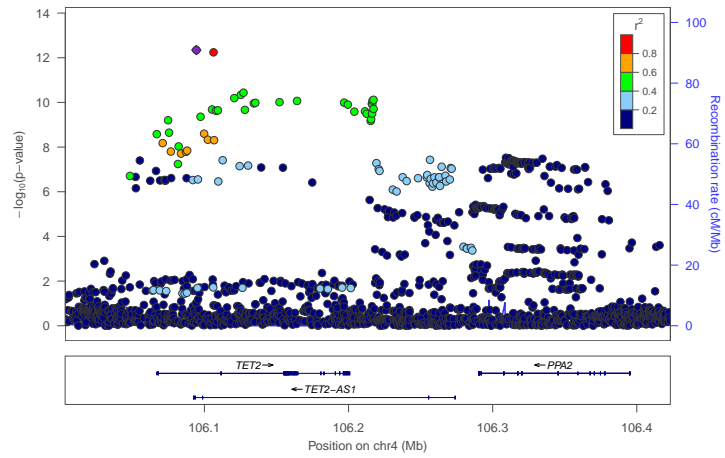
locus_55



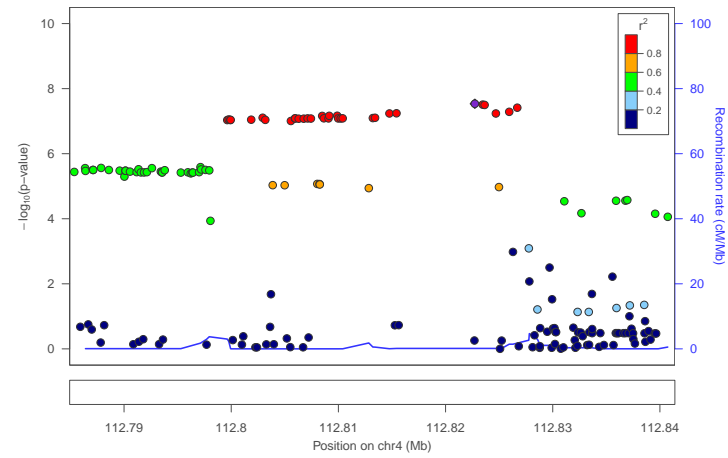
locus_56



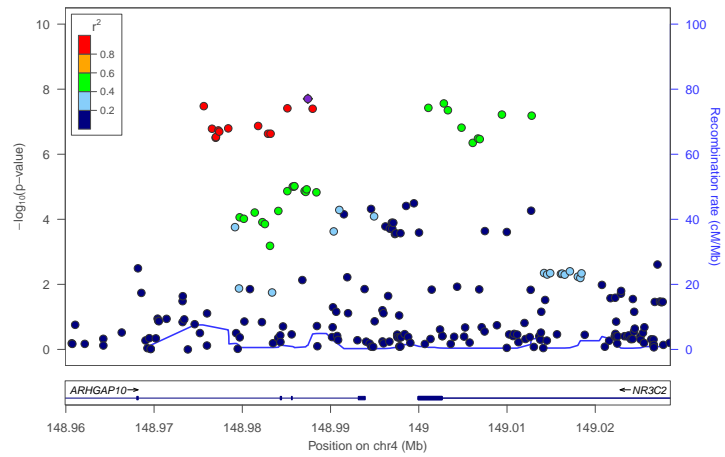
locus_57



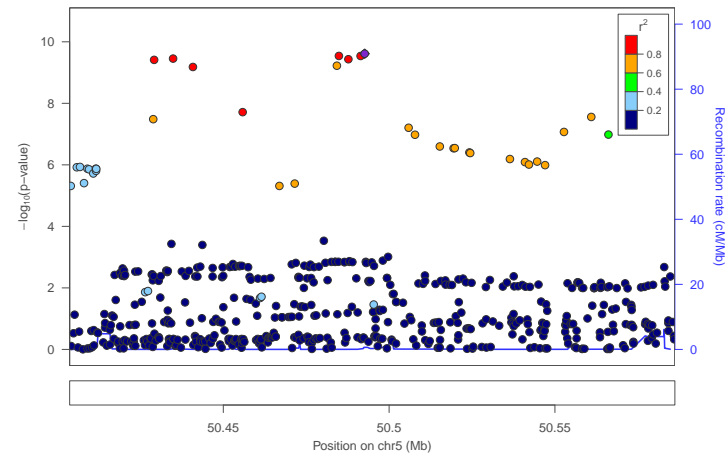
locus_58



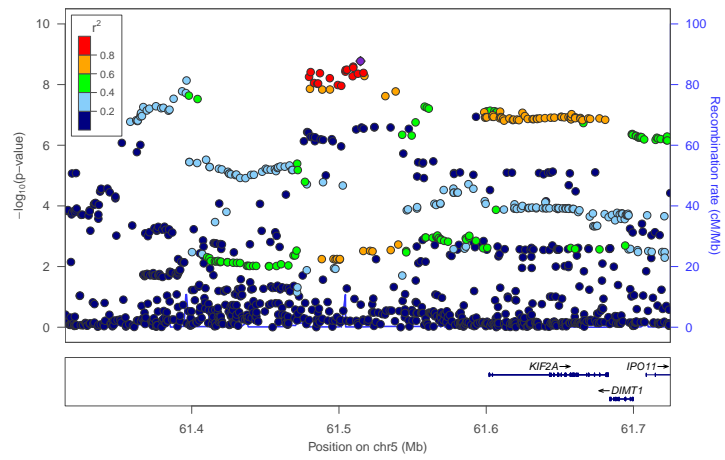
locus_59



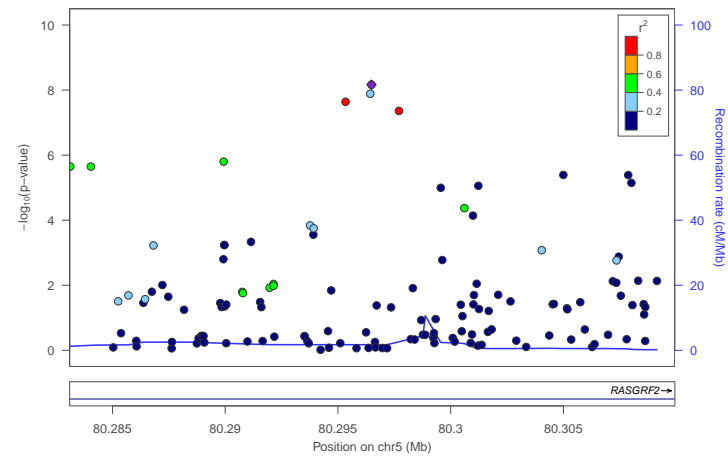
locus_60



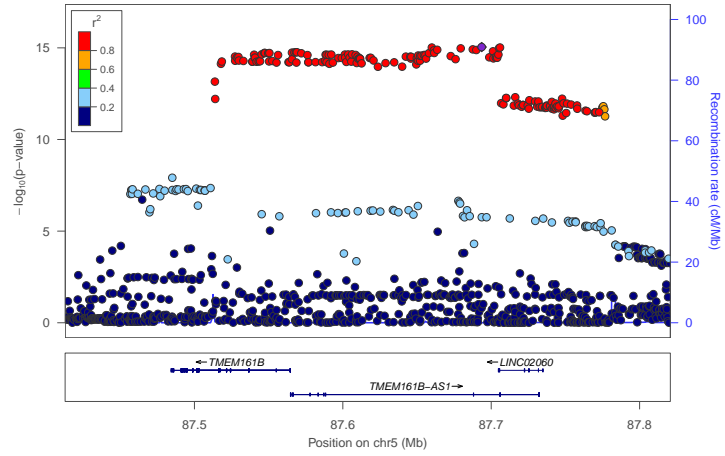
locus_61



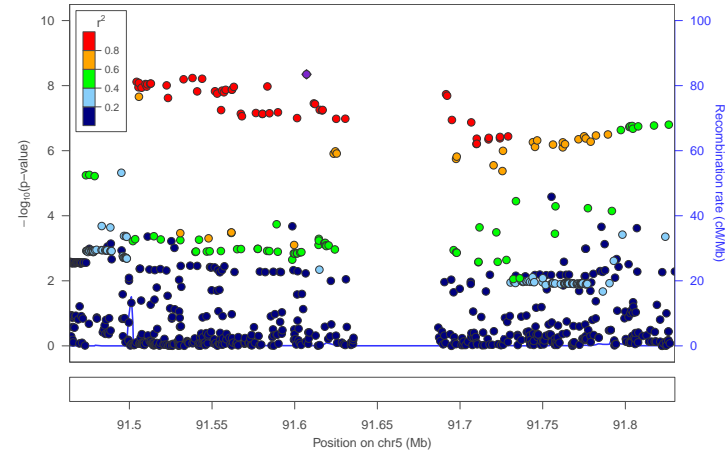
locus_62



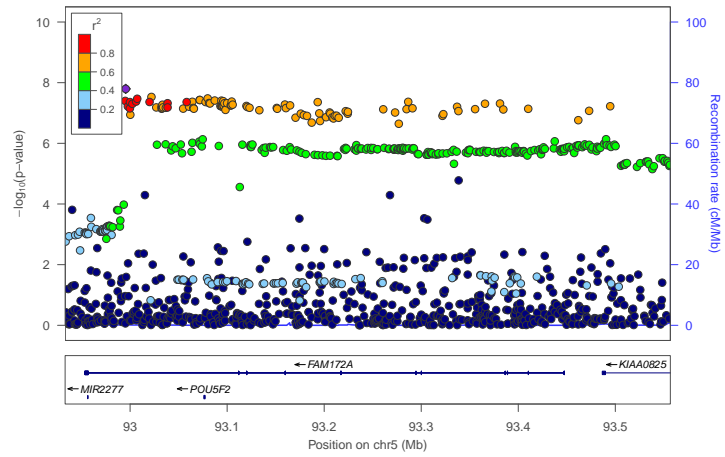
locus_63



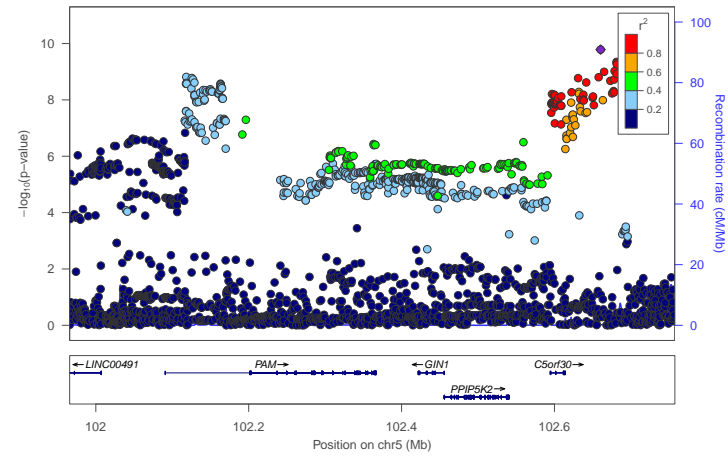
locus_64



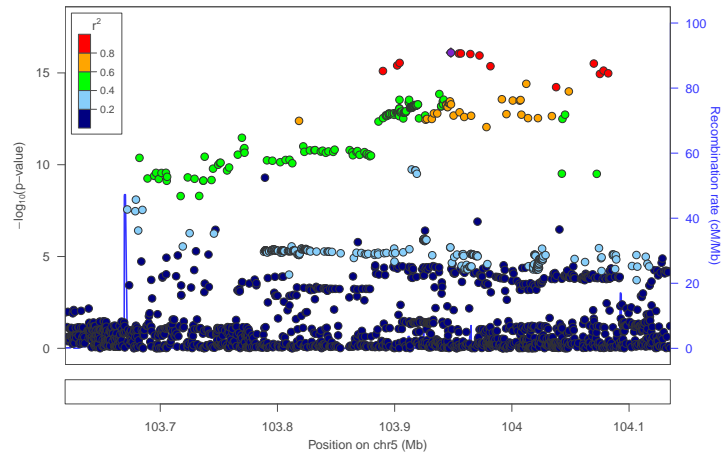
locus_65



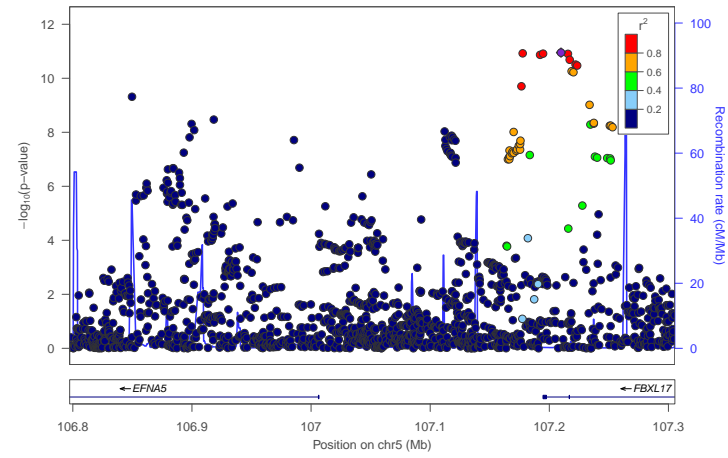
locus_66



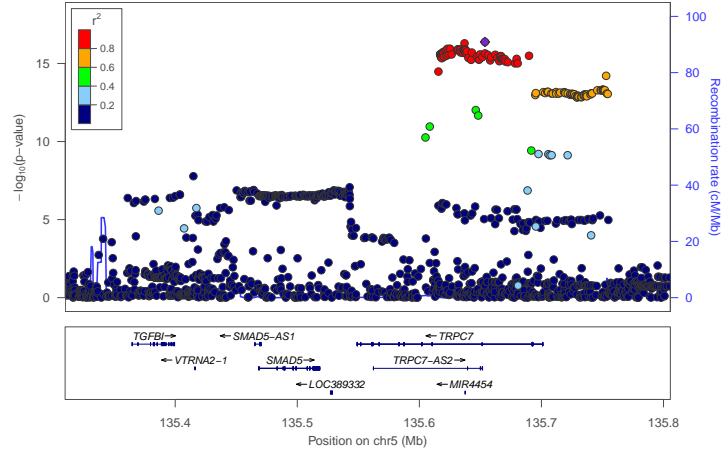
locus_67



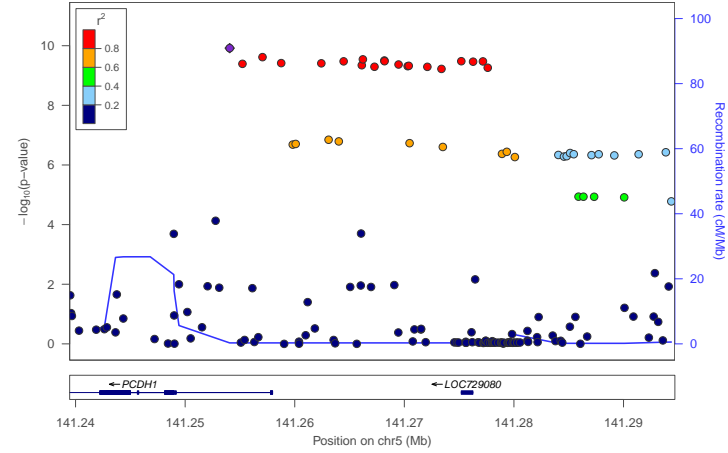
locus_68



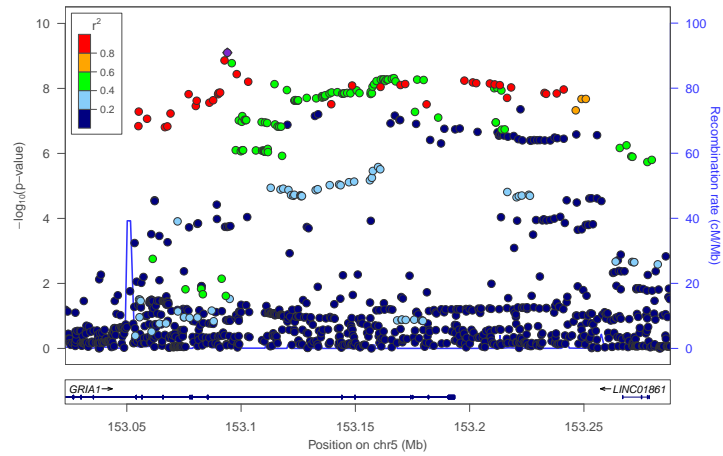
locus_69



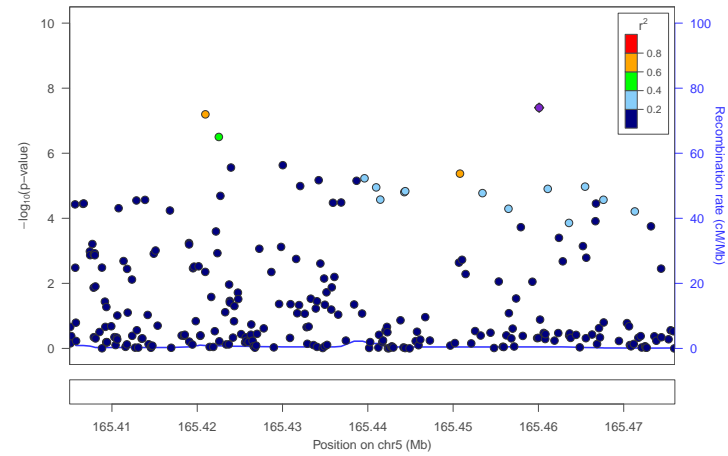
locus_70



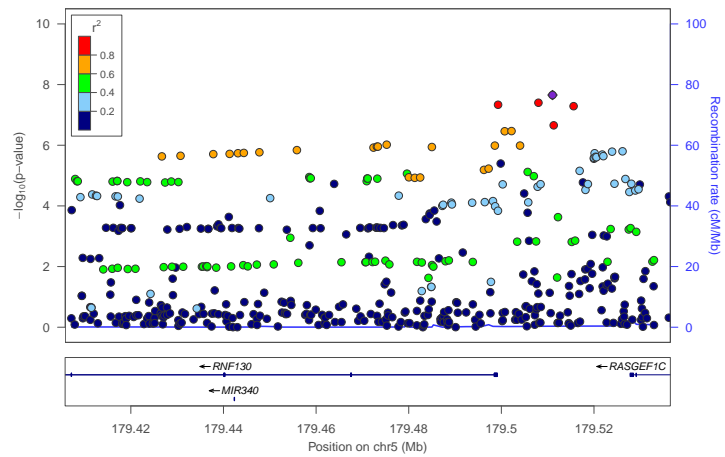
locus_71



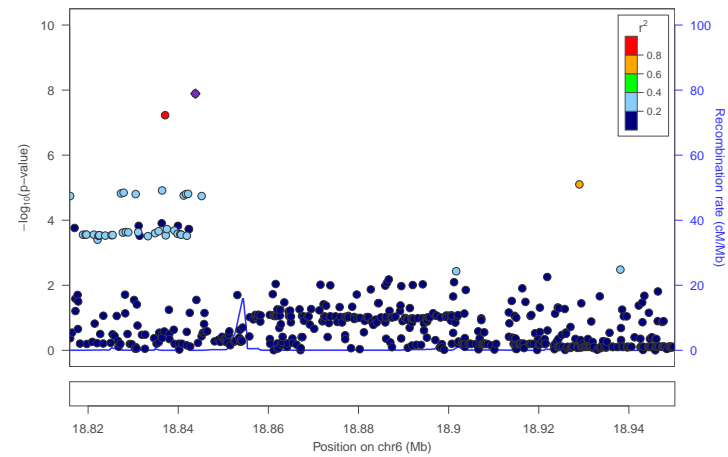
locus_72



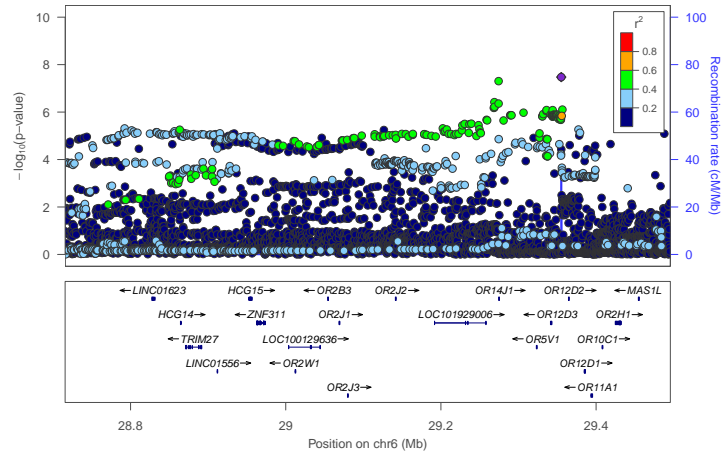
locus_73



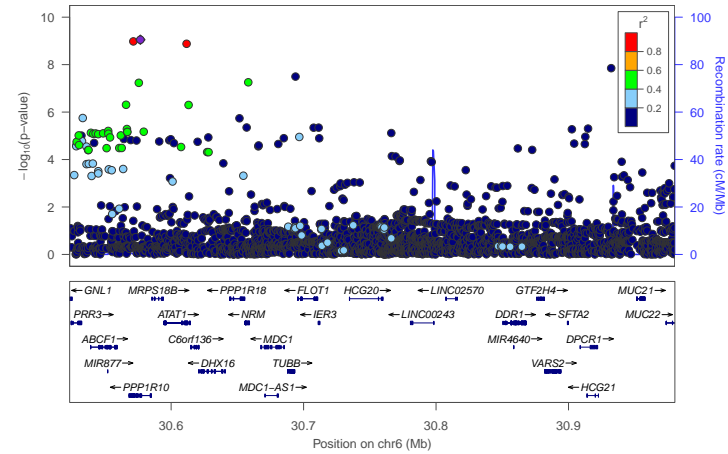
locus_74



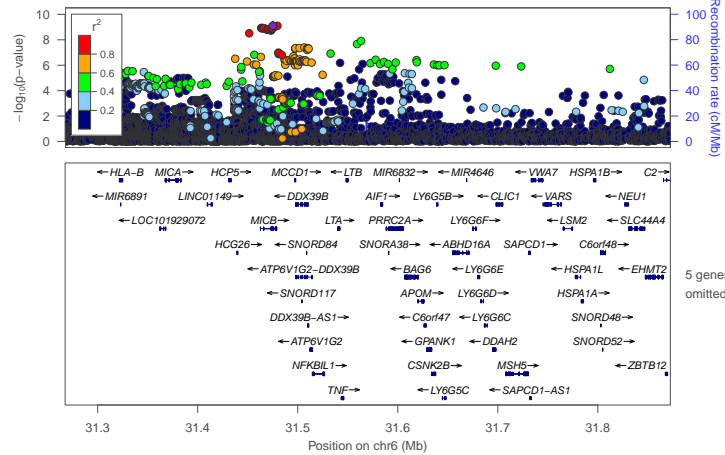
locus_75



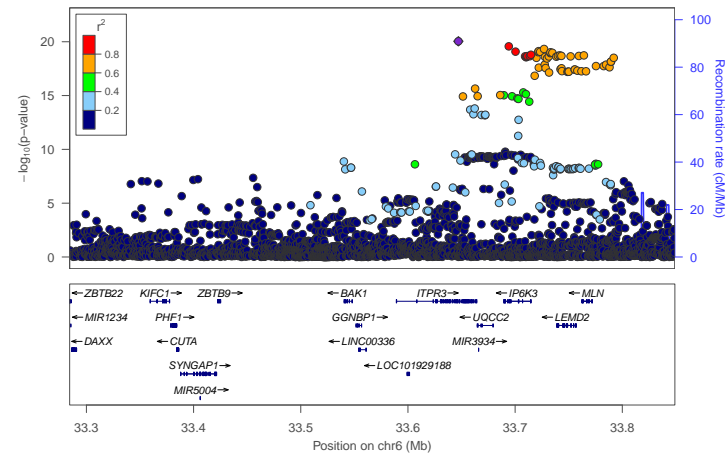
locus_76



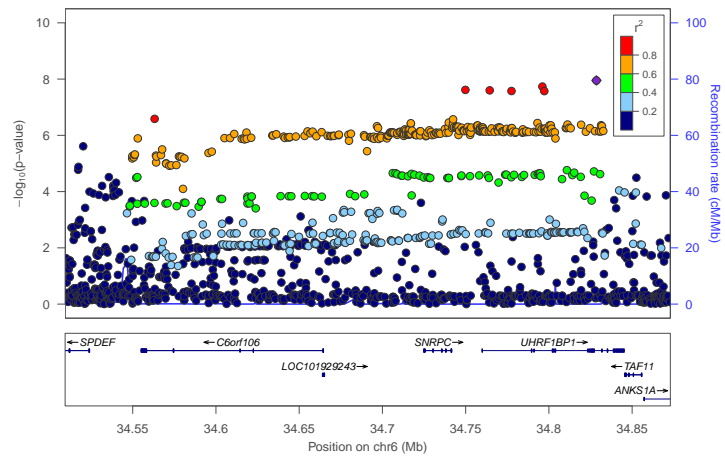
locus_77



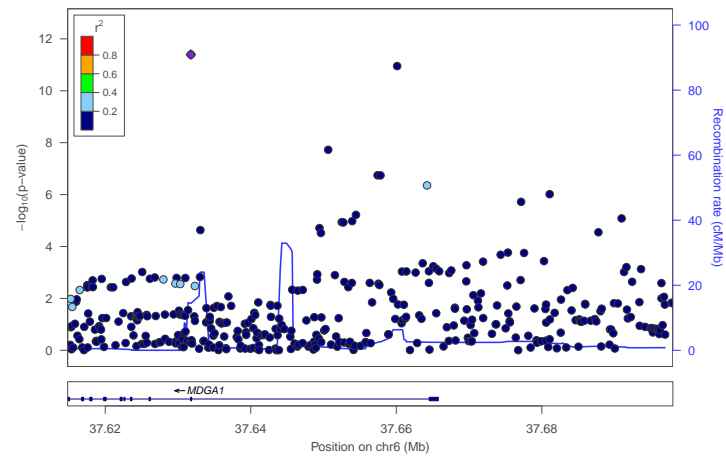
locus_78



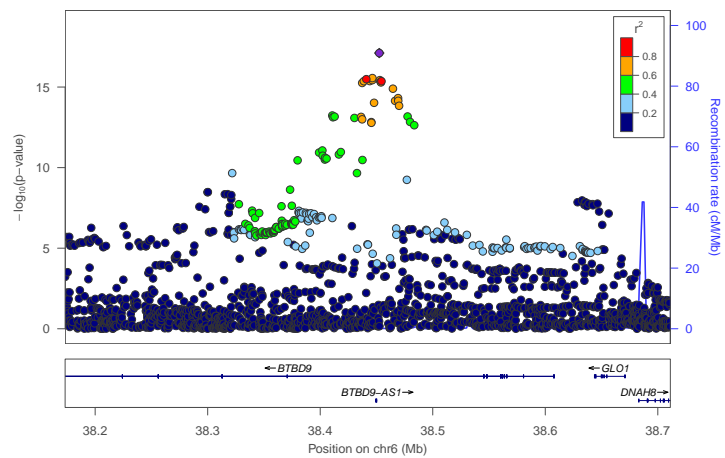
locus_79



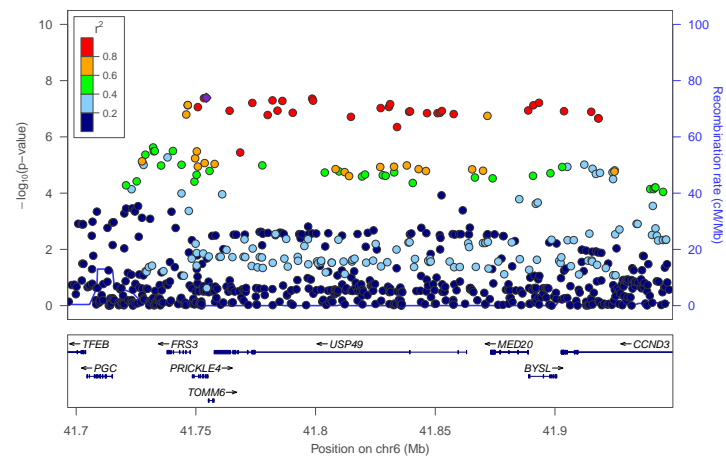
locus_80



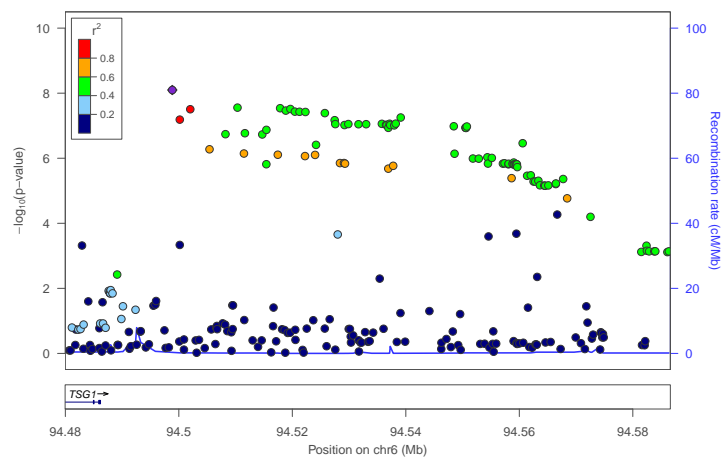
locus_81



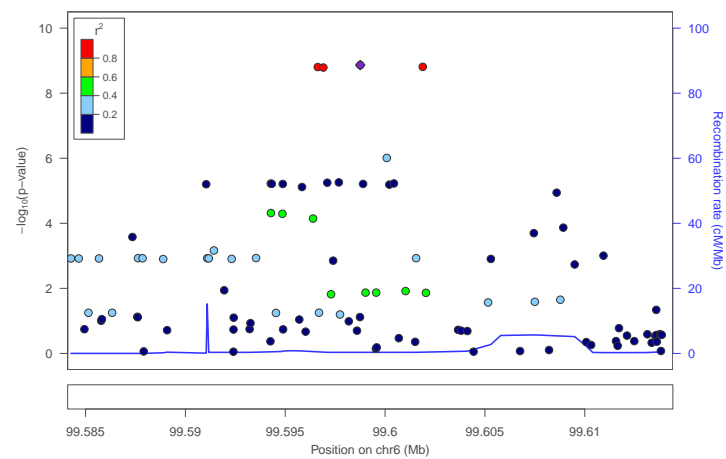
locus_82



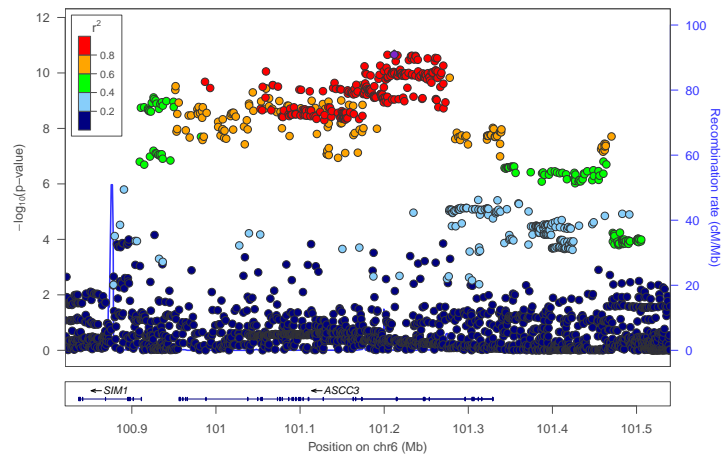
locus_83



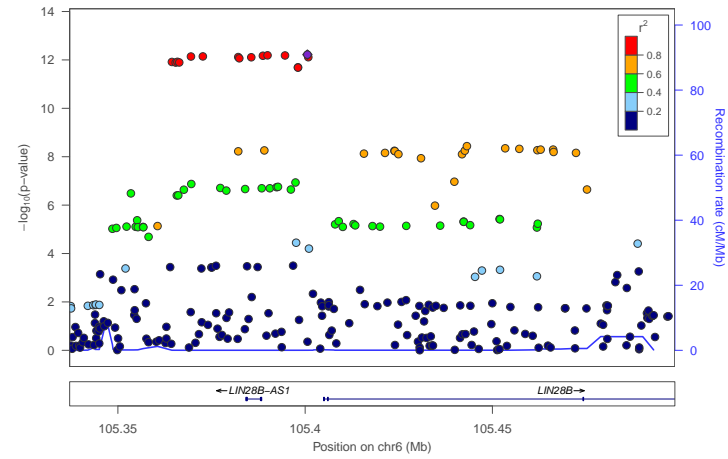
locus_84



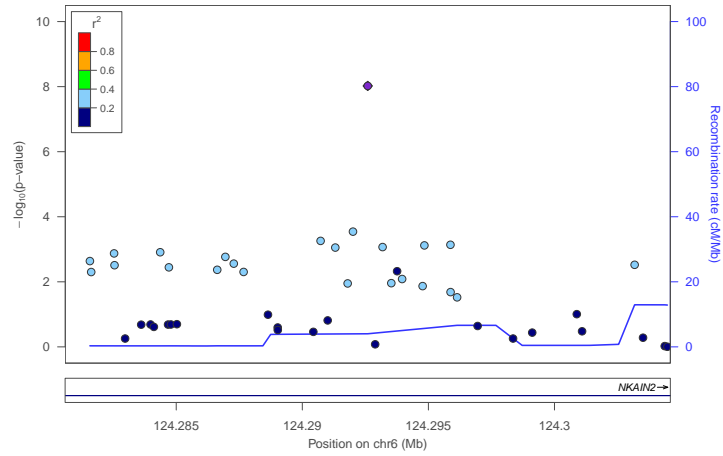
locus_85



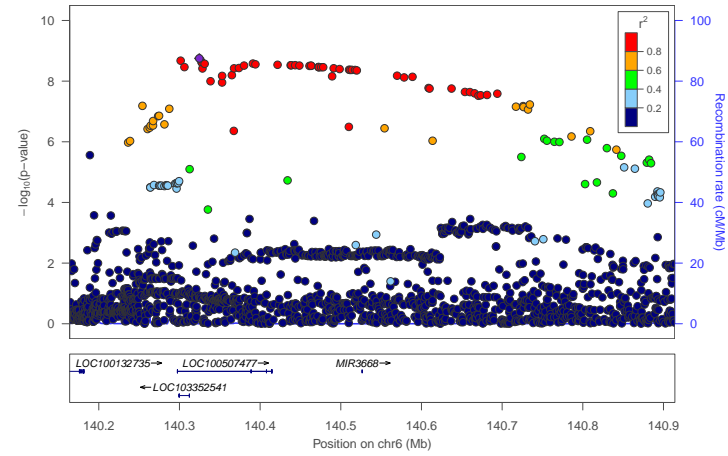
locus_86



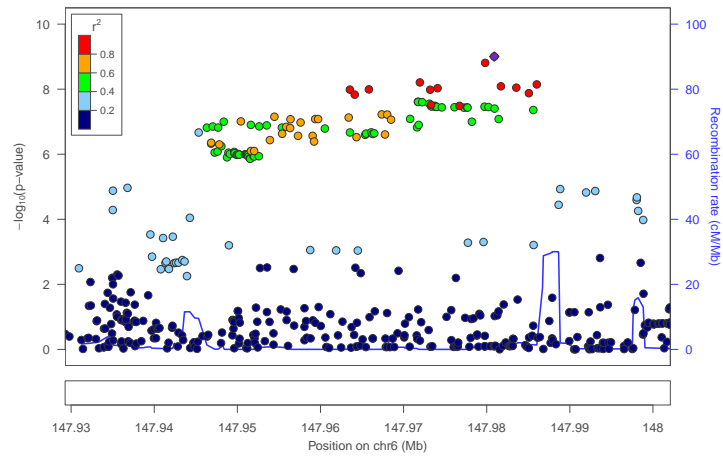
locus_87



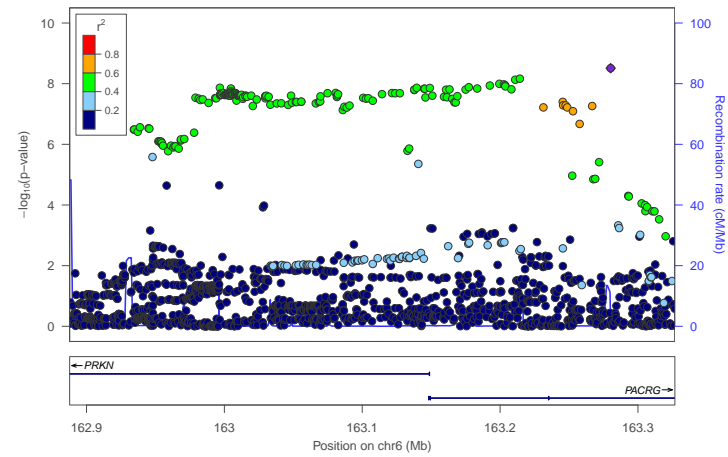
locus_88



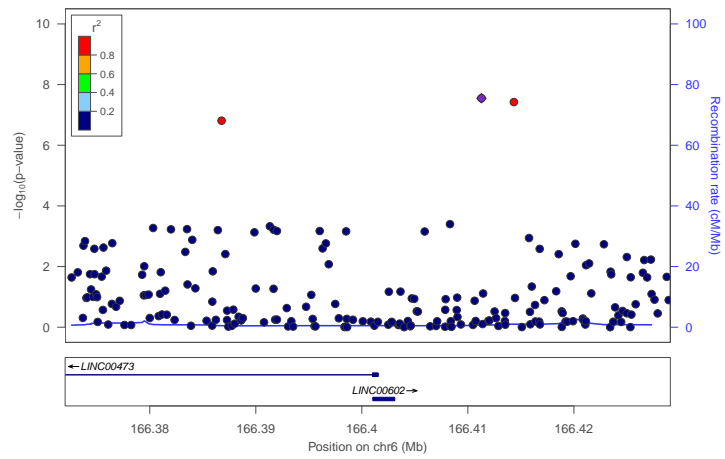
locus_89



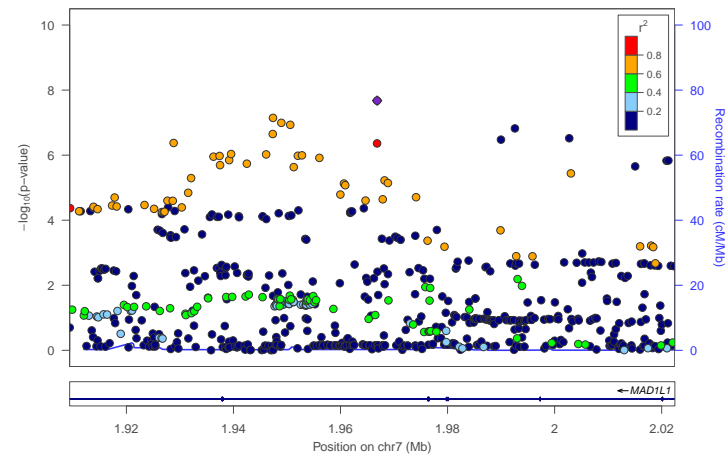
locus_90



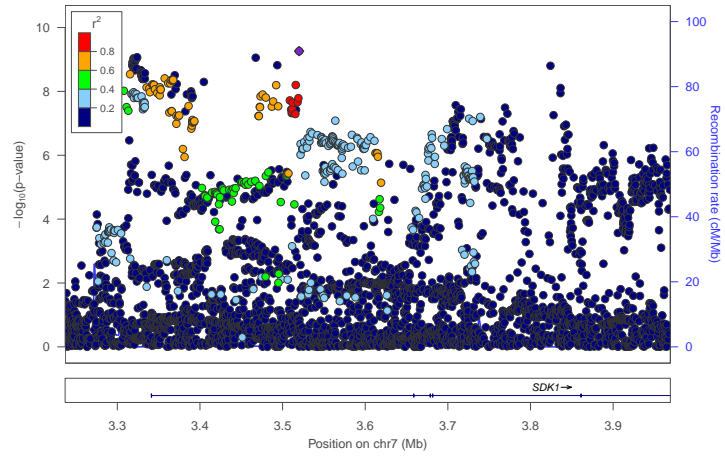
locus_91



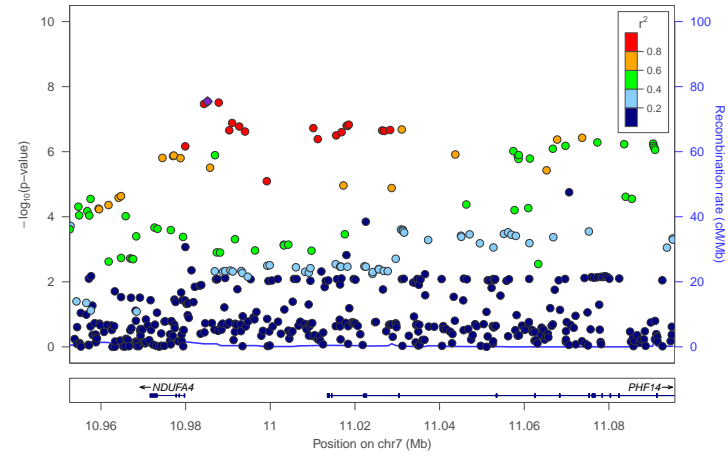
locus_92



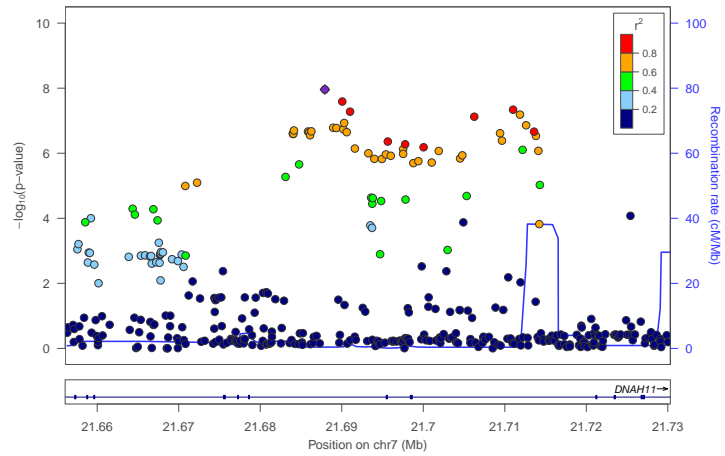
locus_93



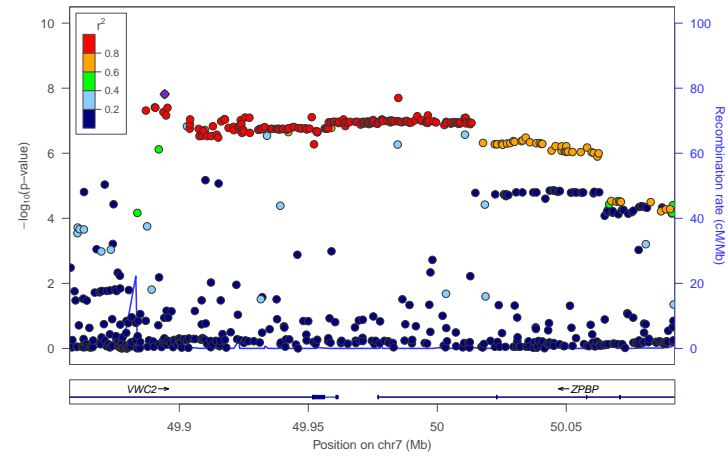
locus_94



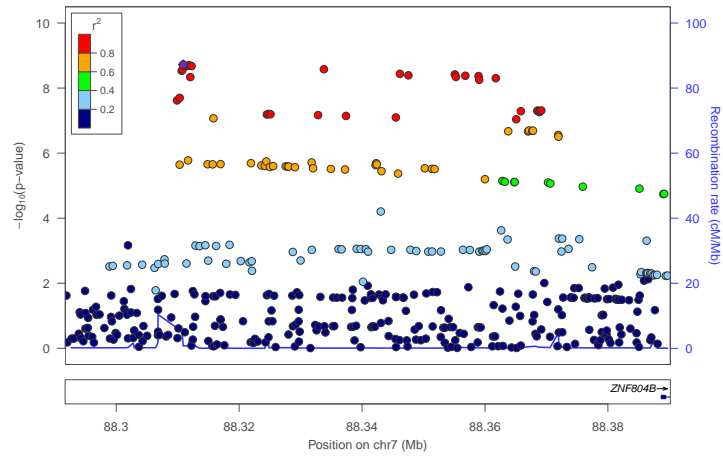
locus_95



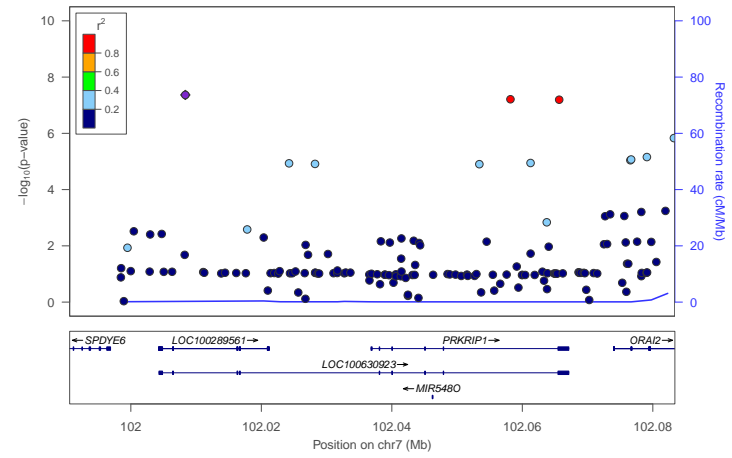
locus_96



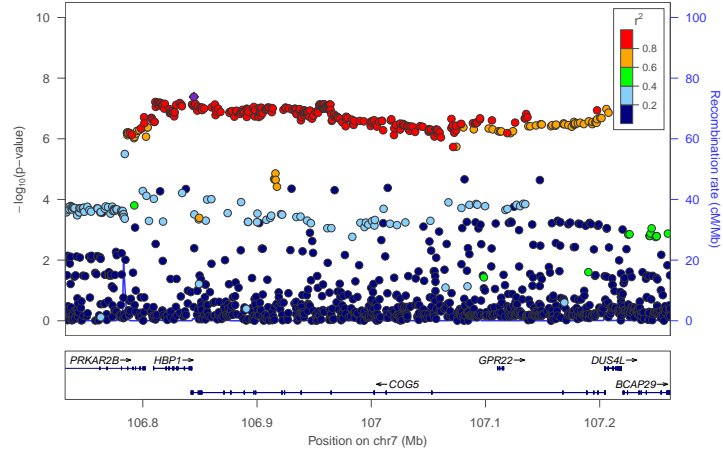
locus_97



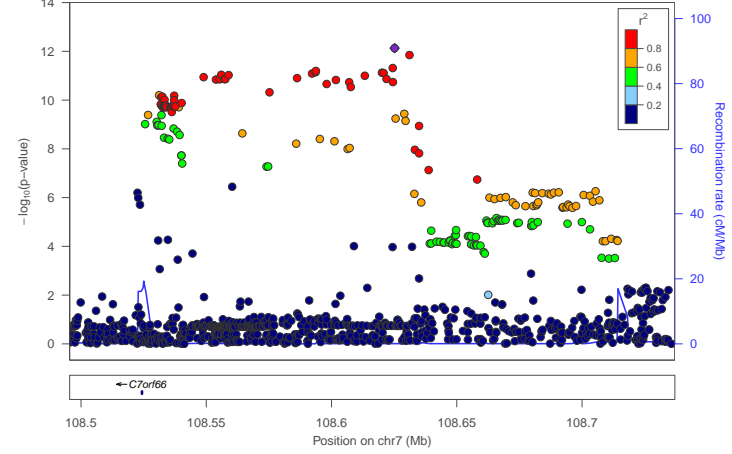
locus_98



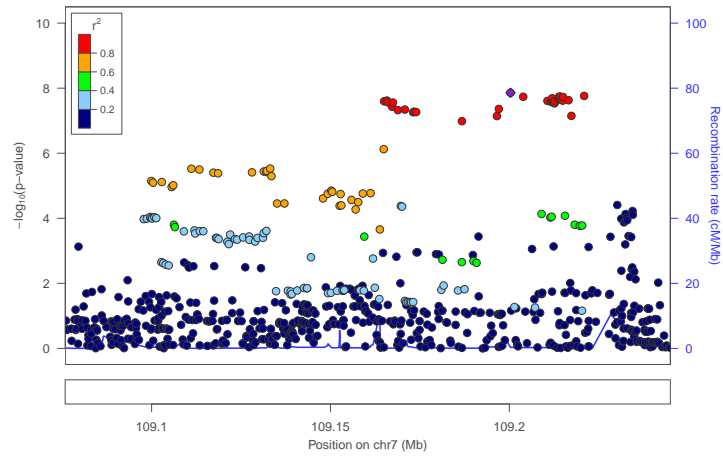
locus_99



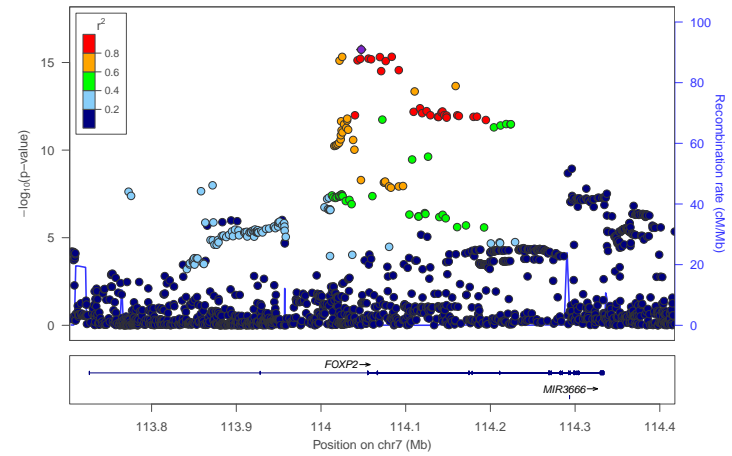
locus_100



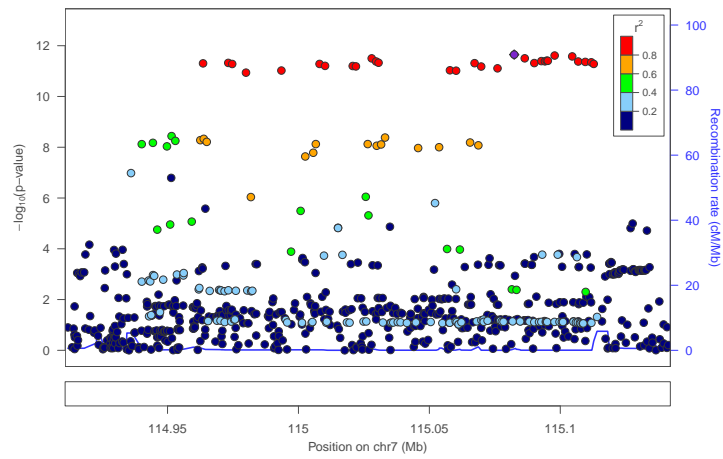
locus_101



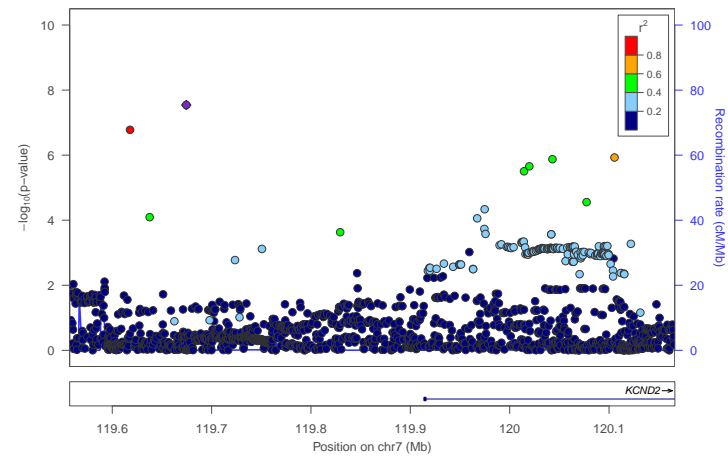
locus_102



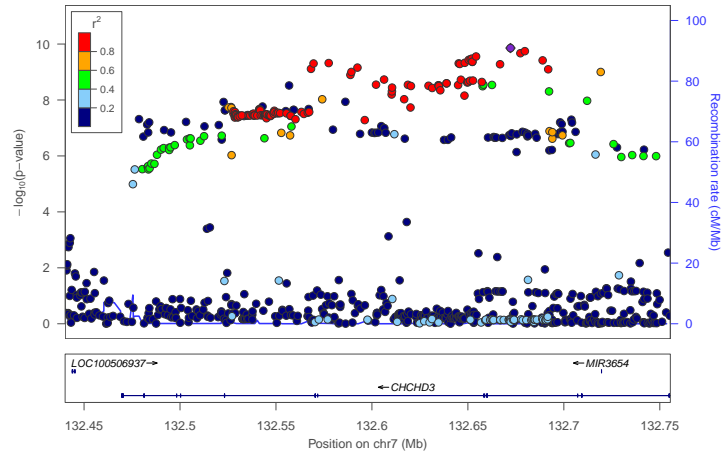
locus_103



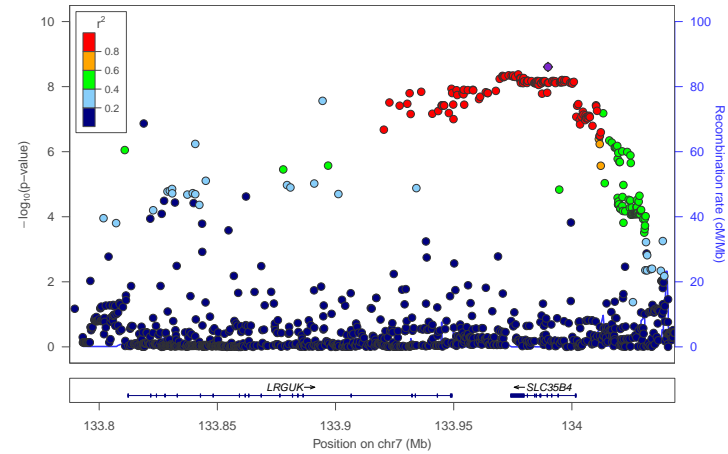
locus_104



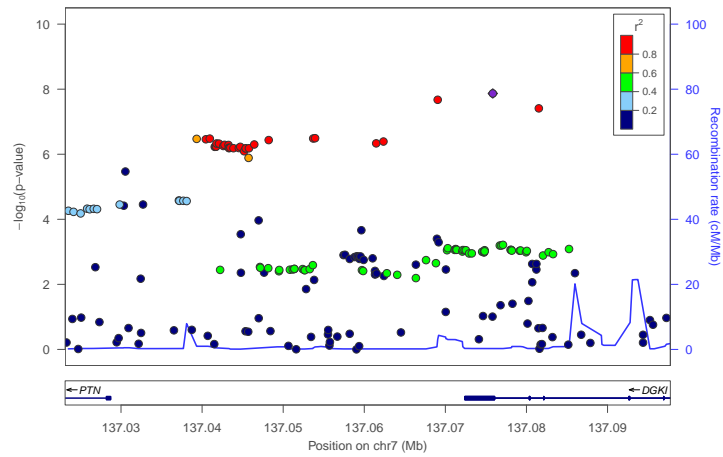
locus_105



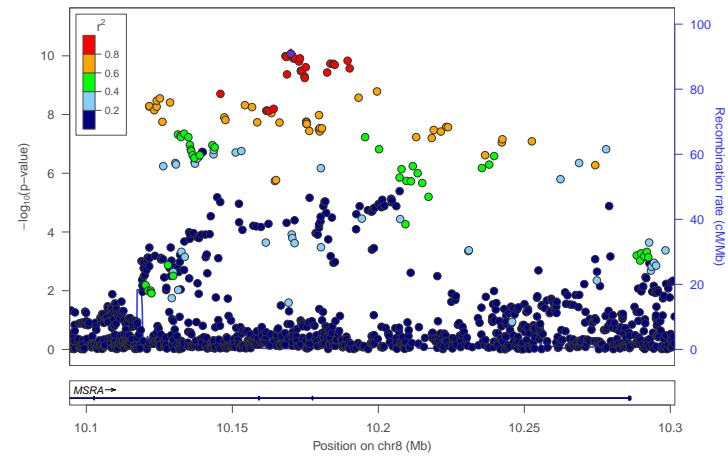
locus_106



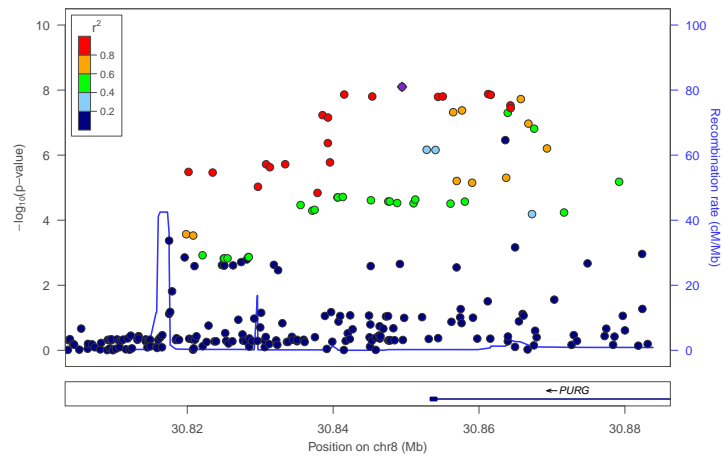
locus_107



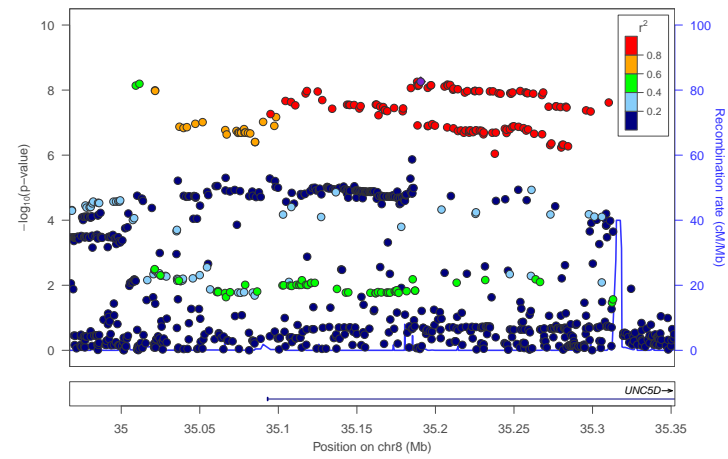
locus_108



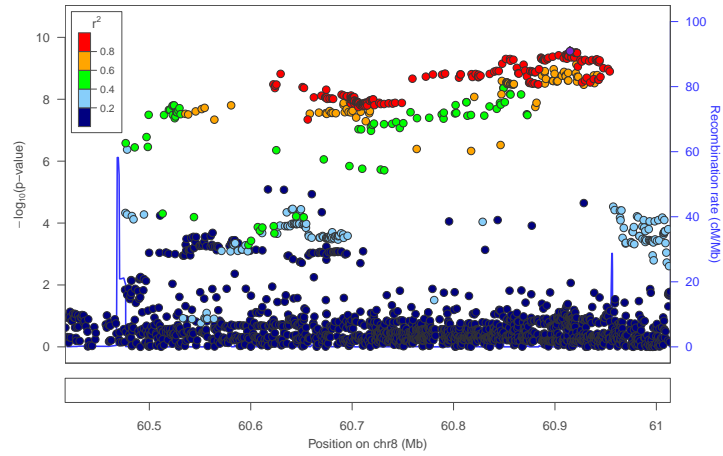
locus_109



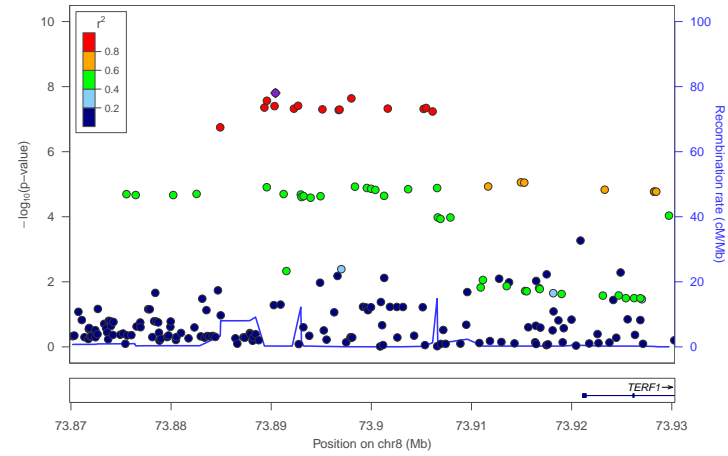
locus_110



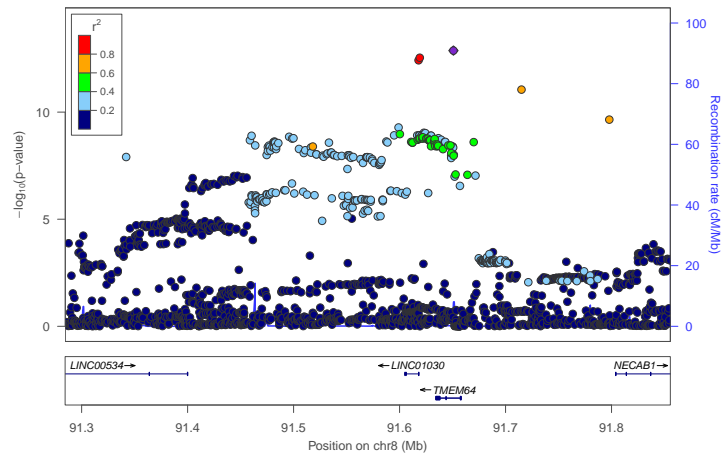
locus_111



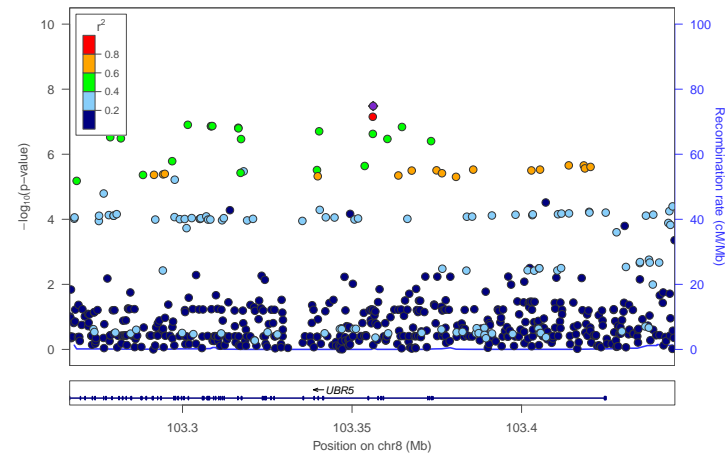
locus_112

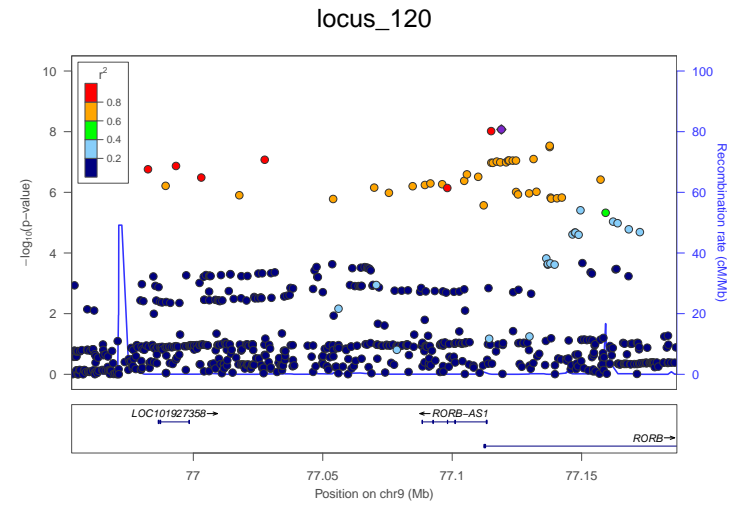
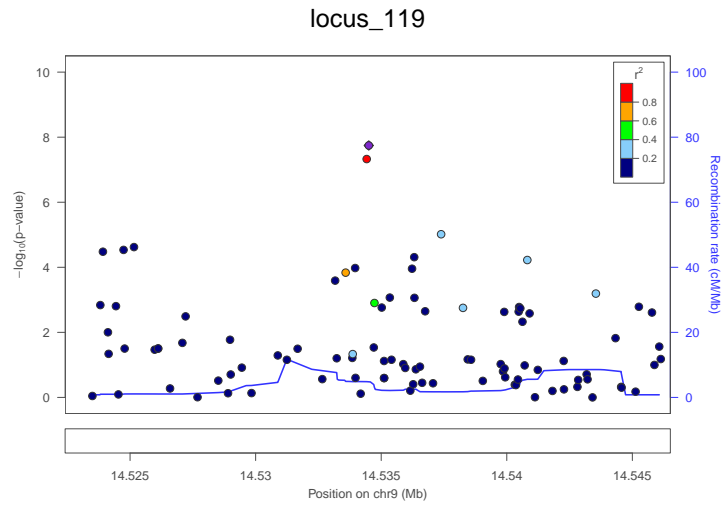
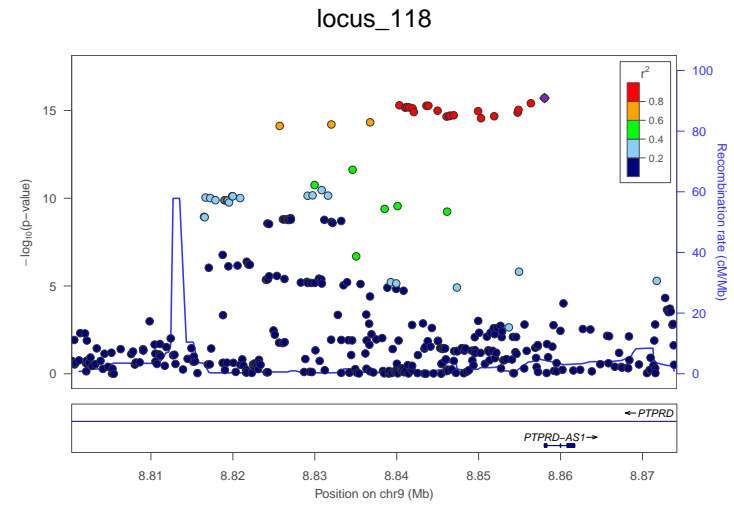
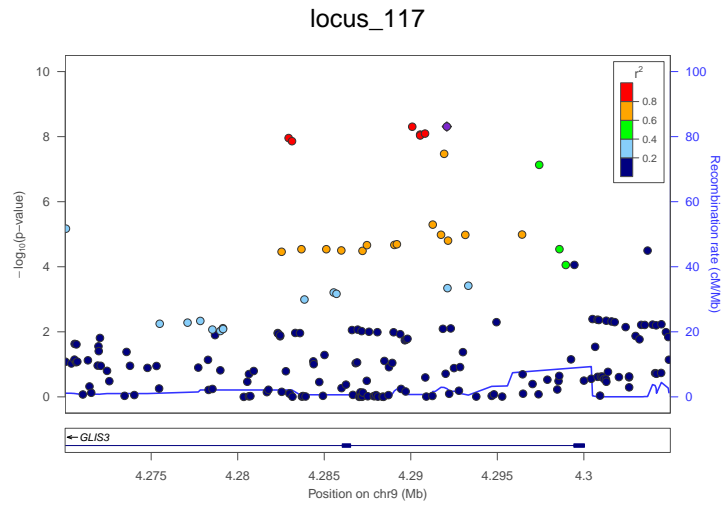
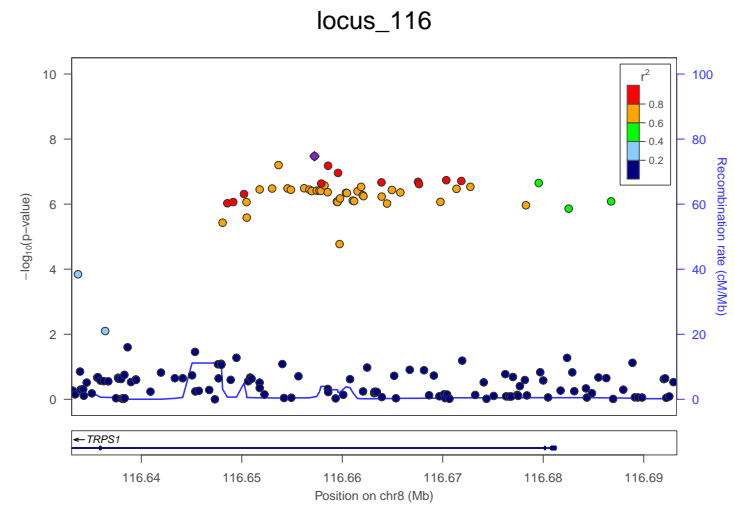
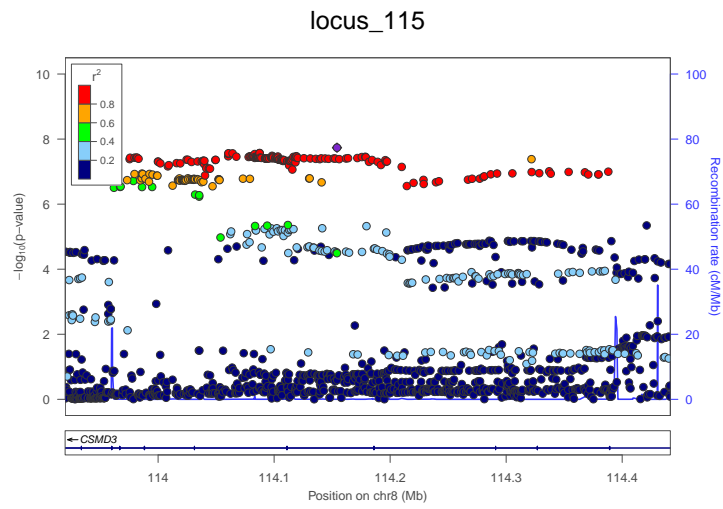


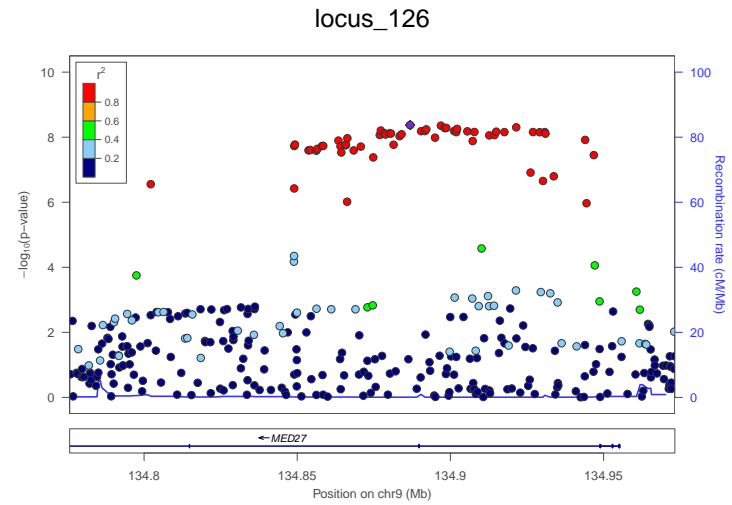
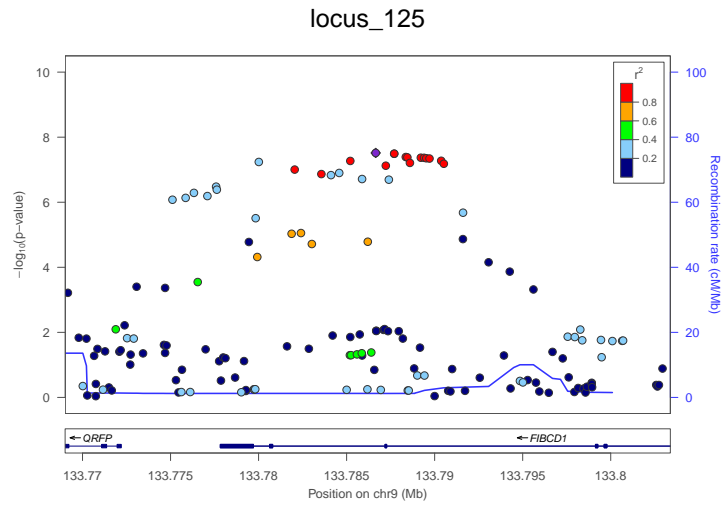
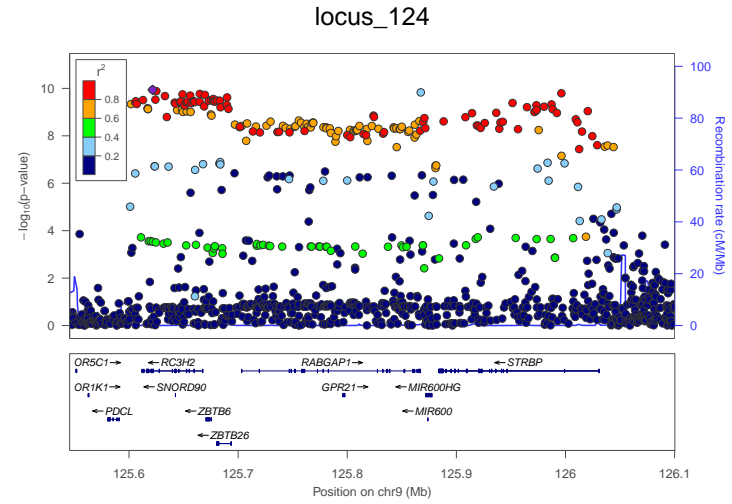
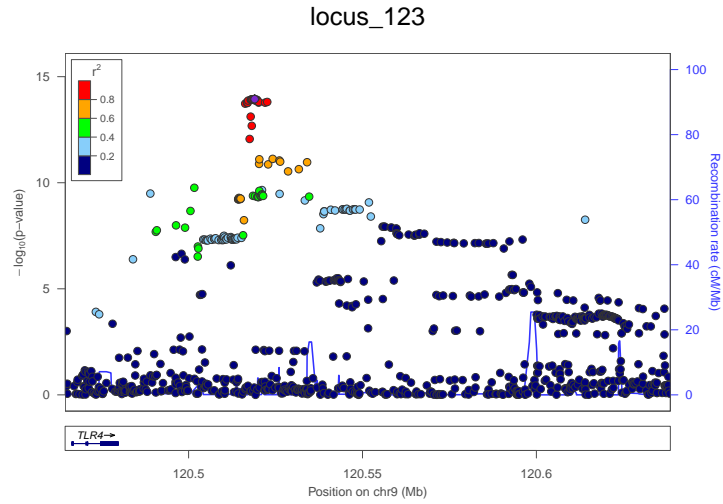
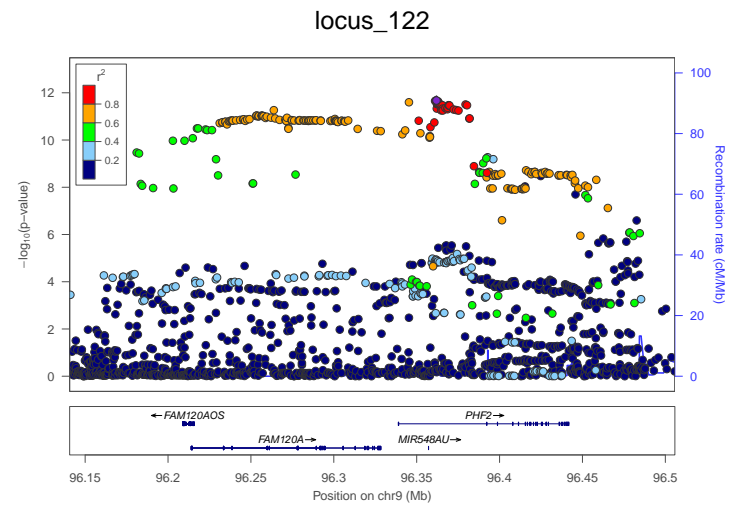
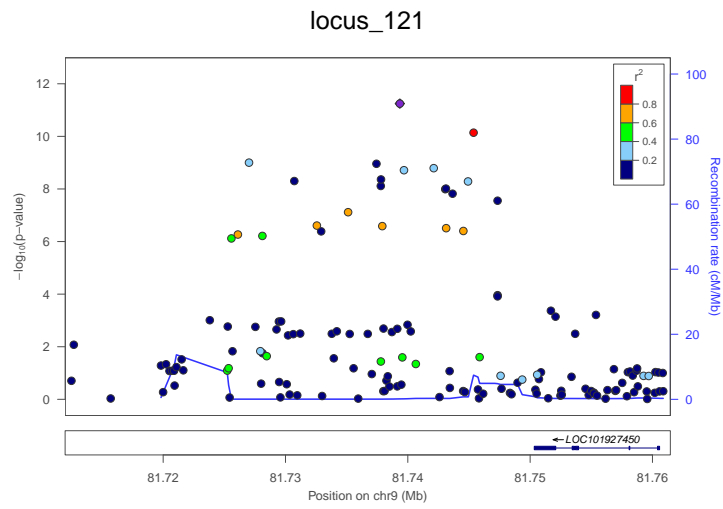
locus_113



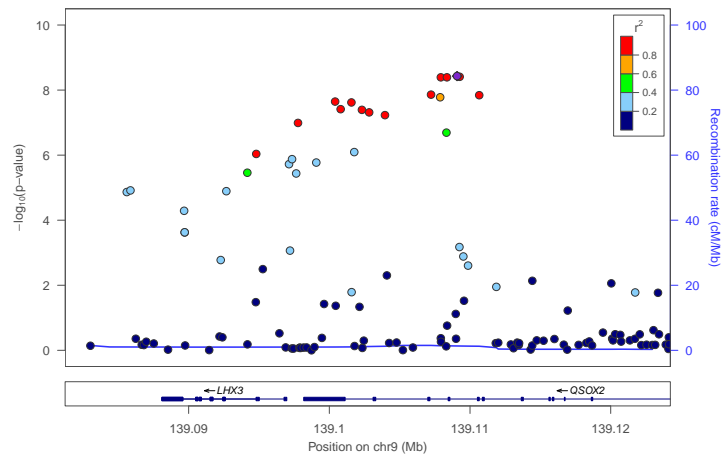
locus_114



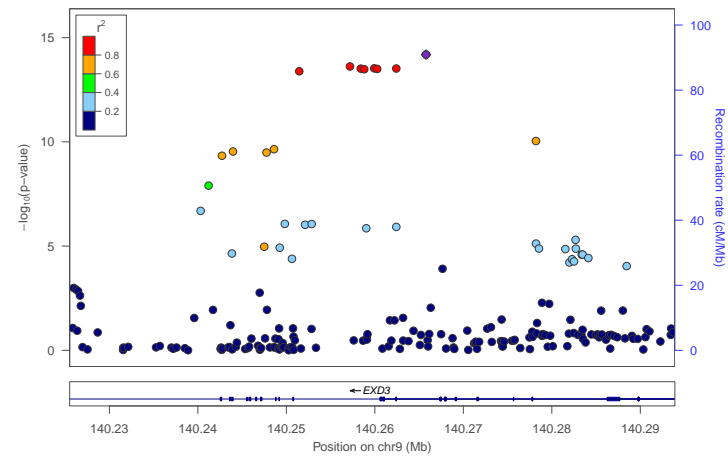




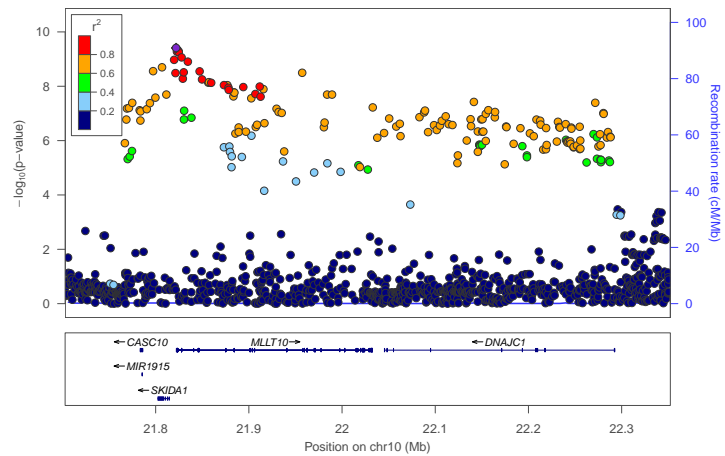
locus_127



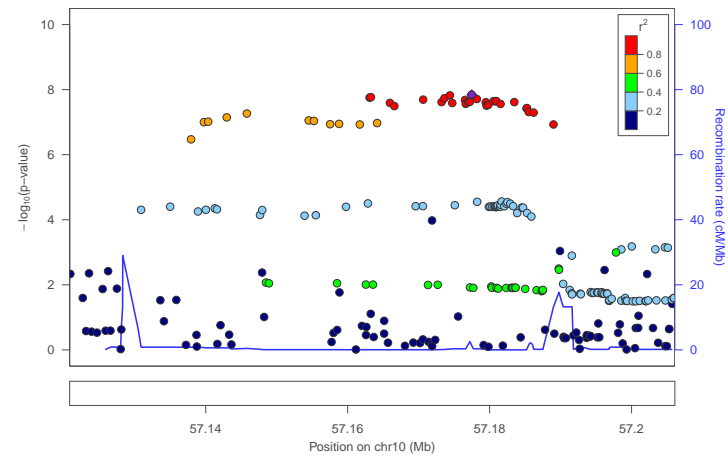
locus_128



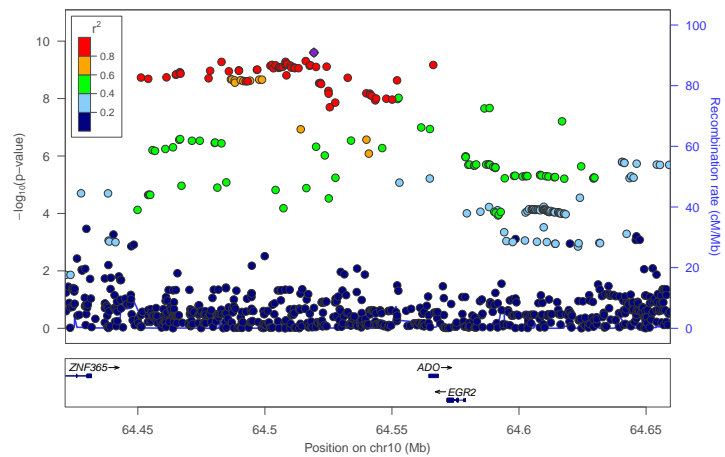
locus_129



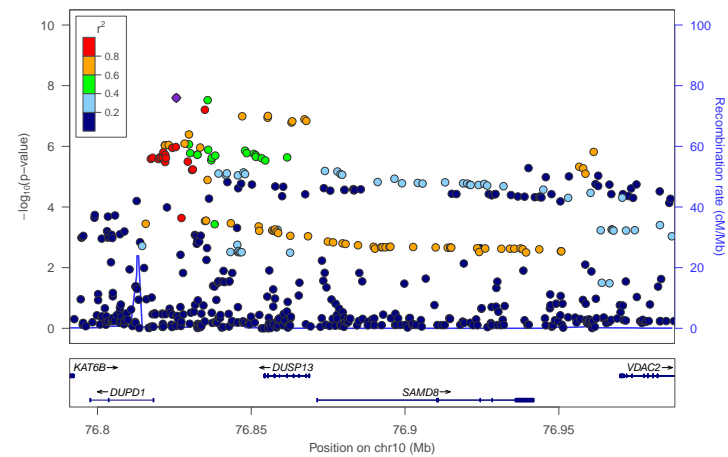
locus_130



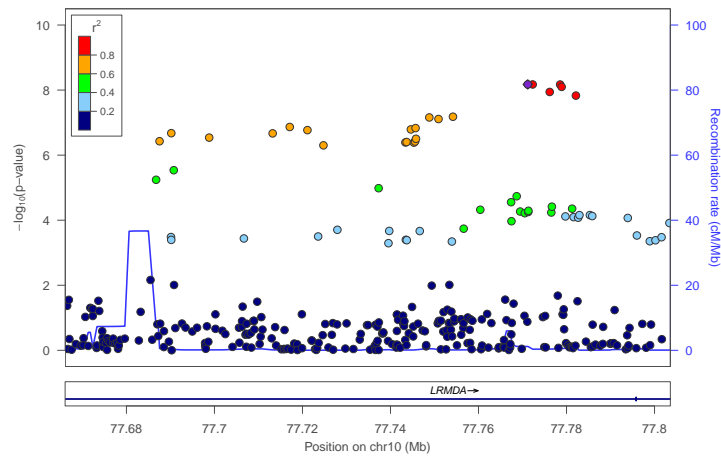
locus_131



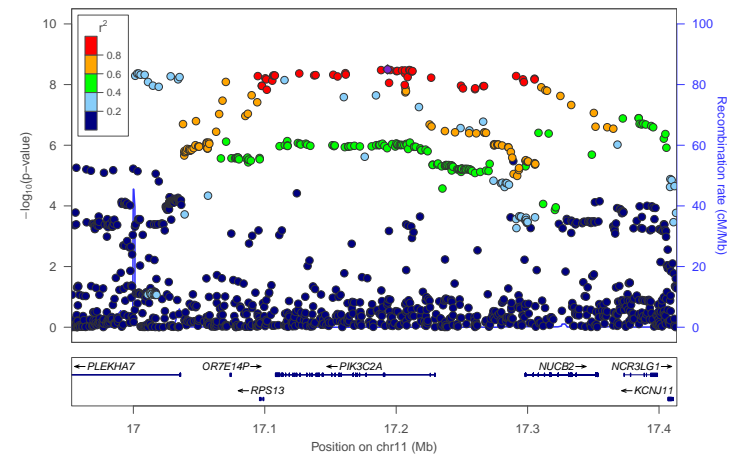
locus_132



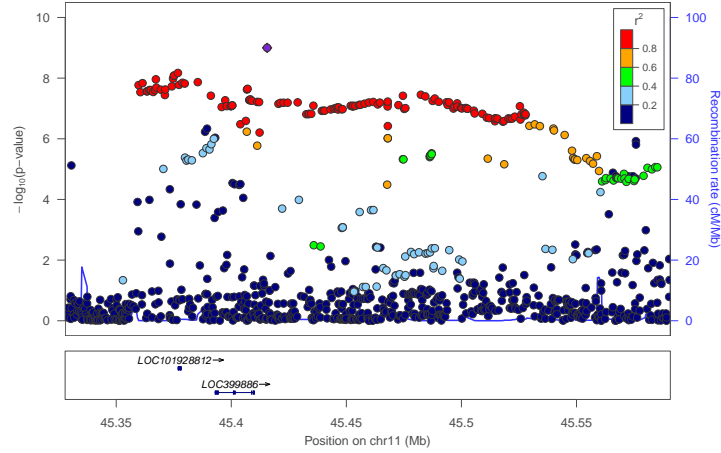
locus_133



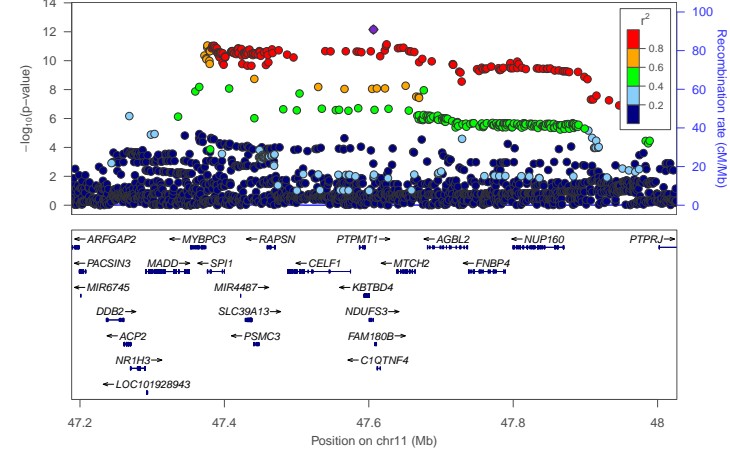
locus_134



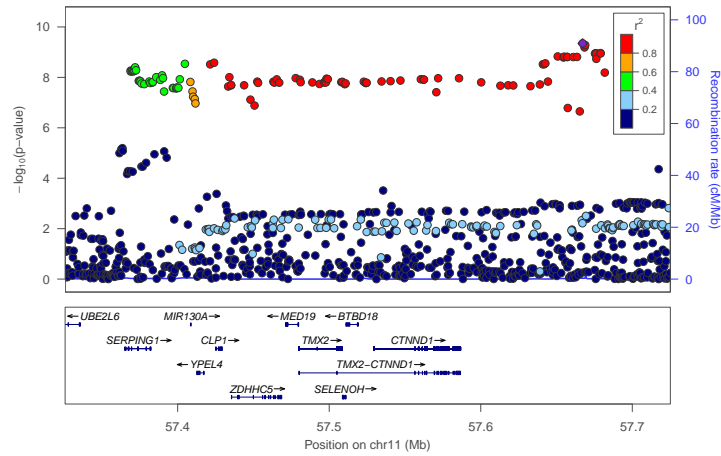
locus_135



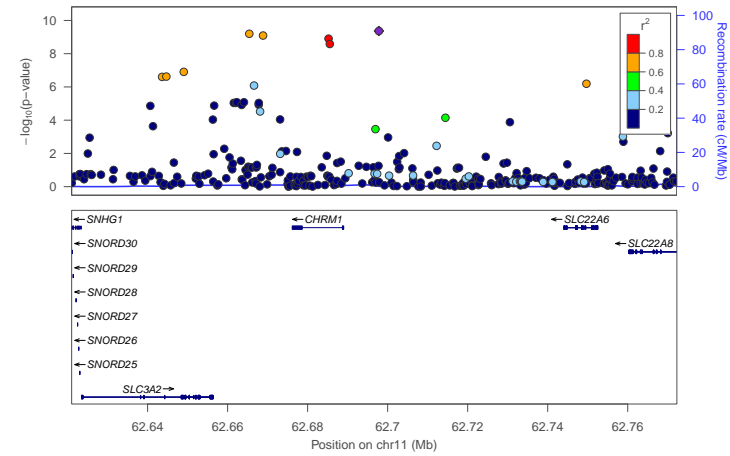
locus_136



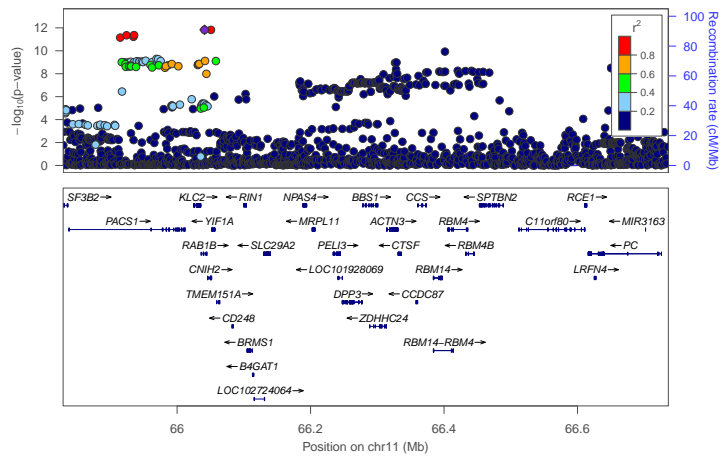
locus_137



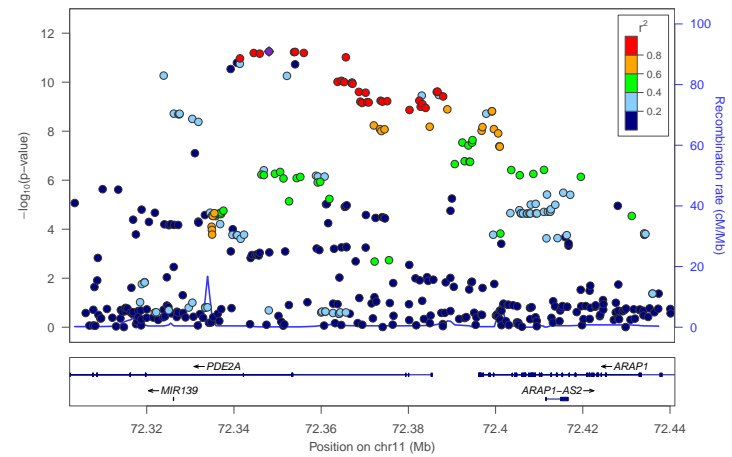
locus_138



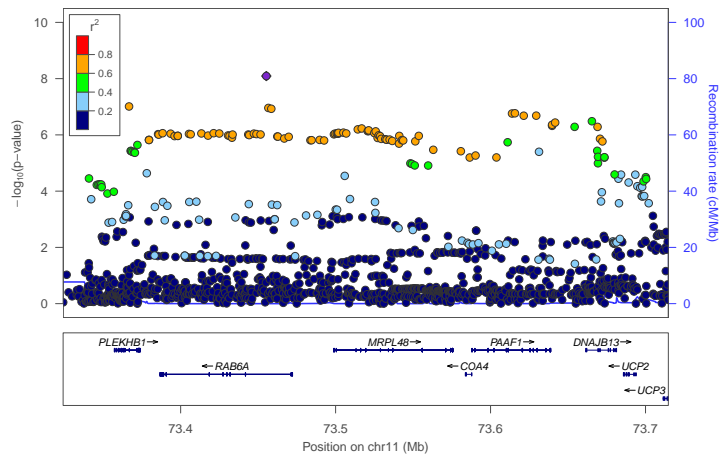
locus_139



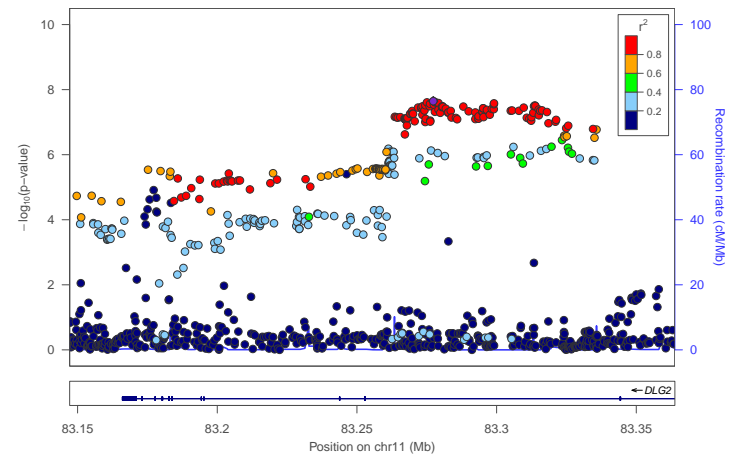
locus_140



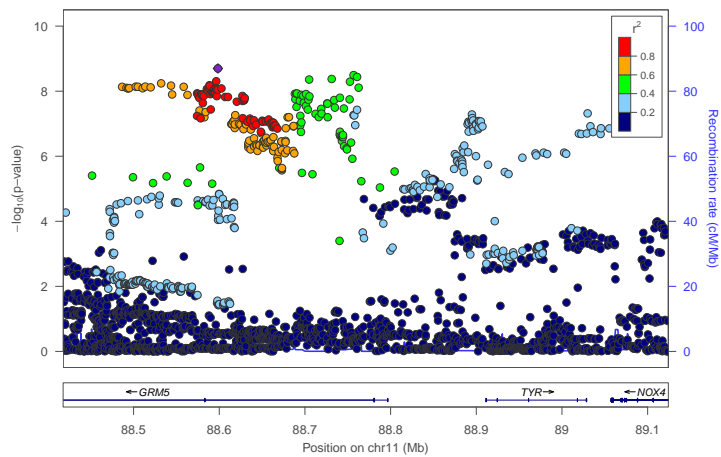
locus_141



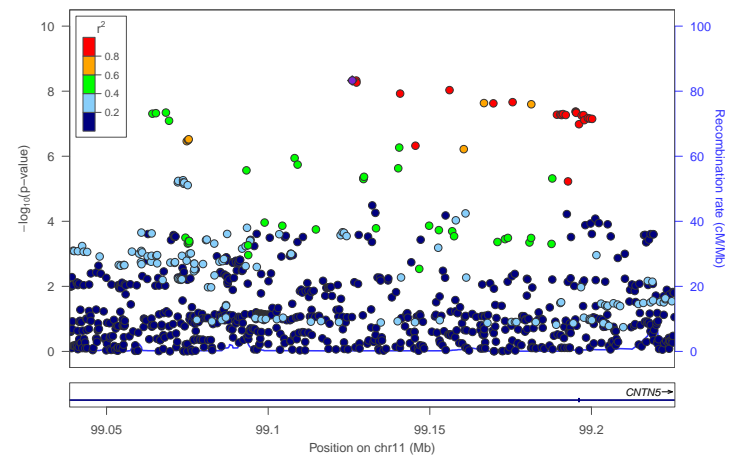
locus_142



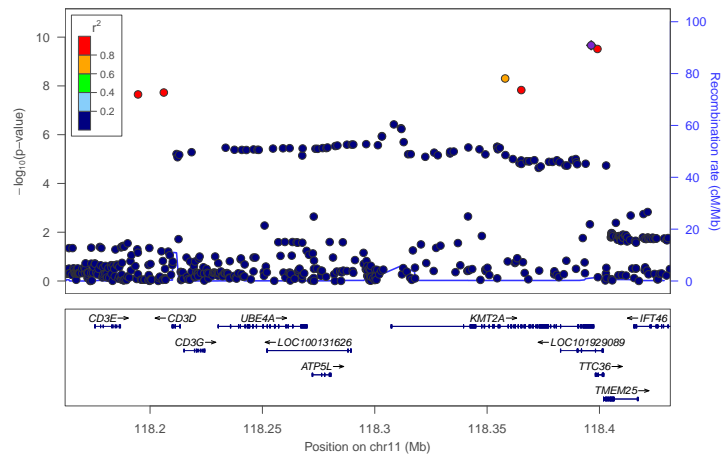
locus_143



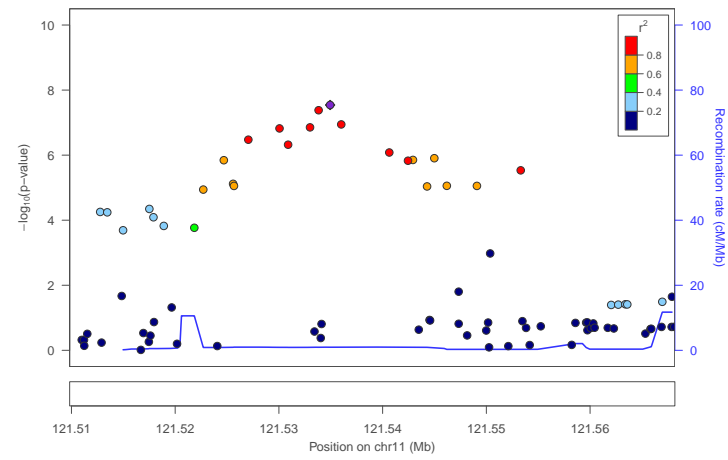
locus_144



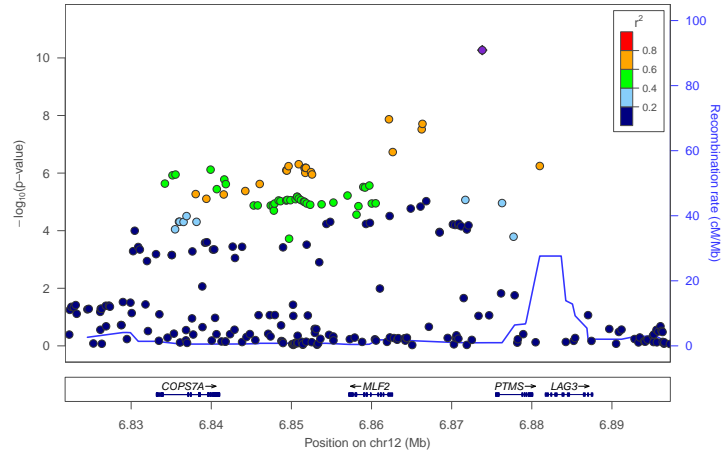
locus_145



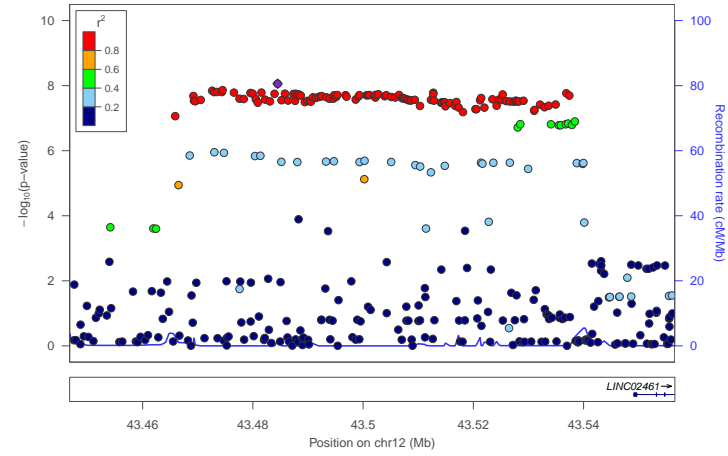
locus_146



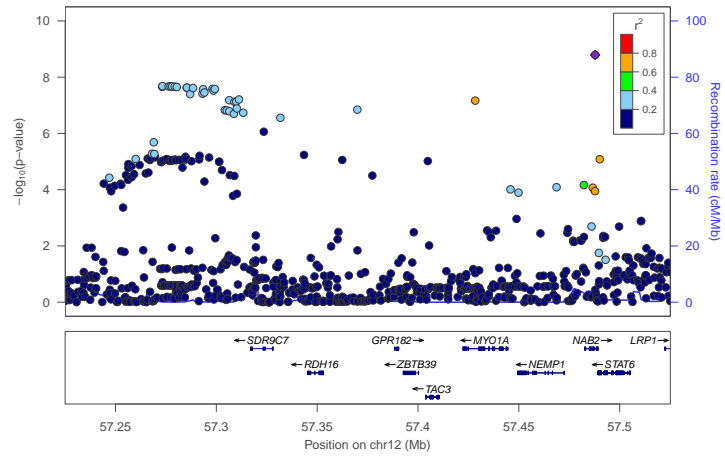
locus_147



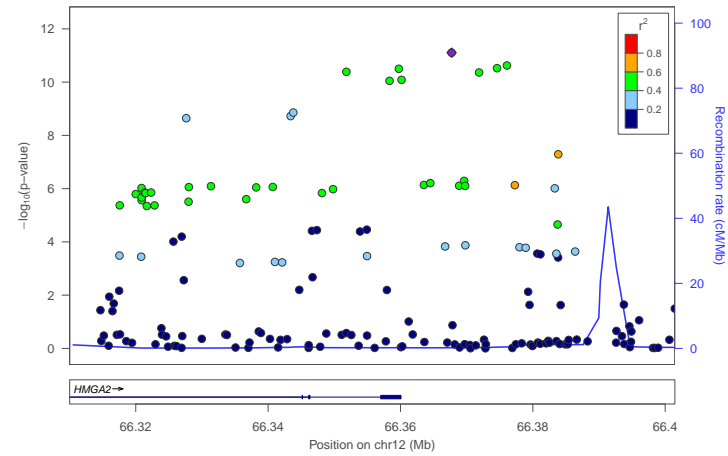
locus_148



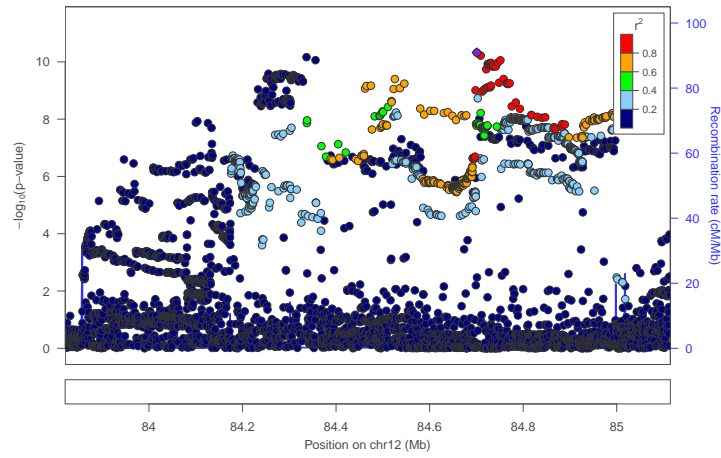
locus_149



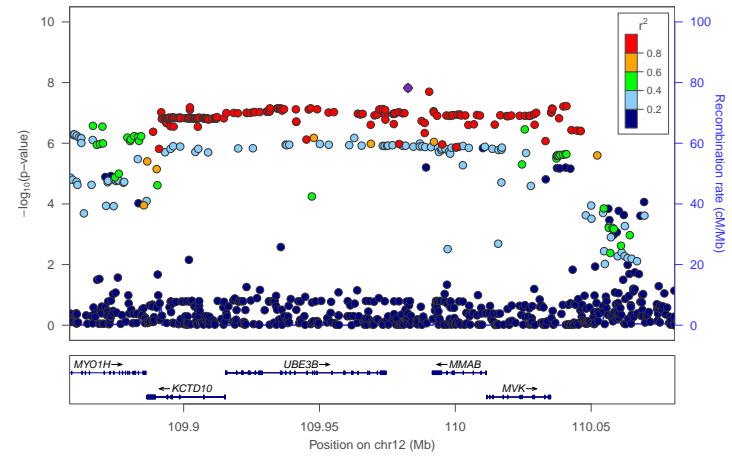
locus_150



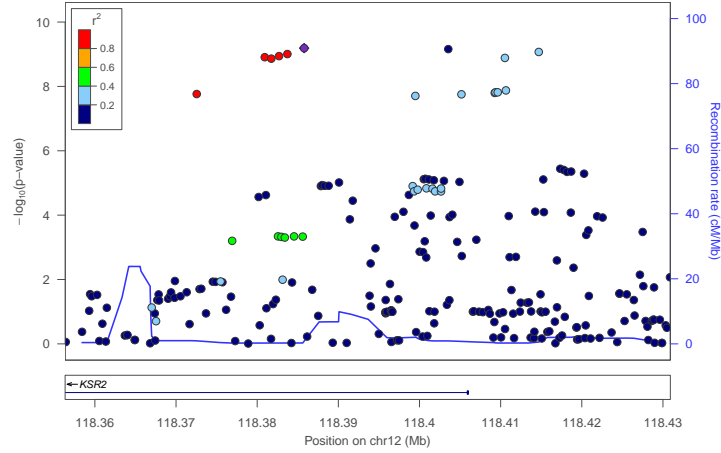
locus_151



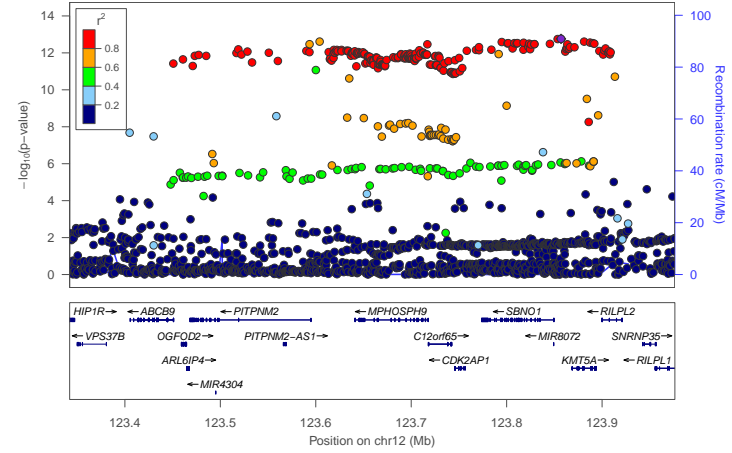
locus_152



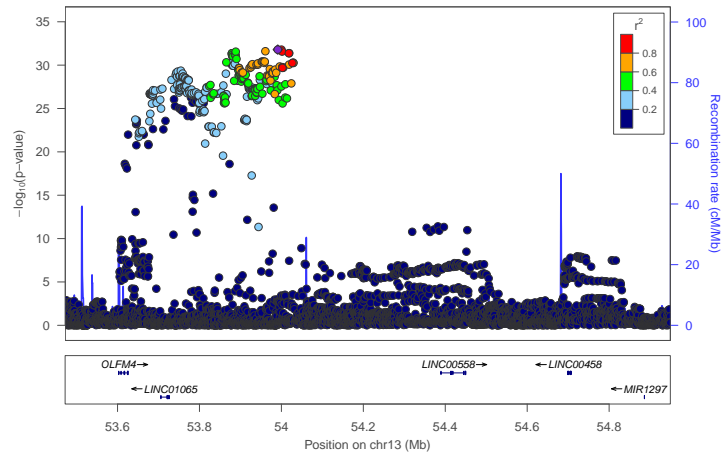
locus_153



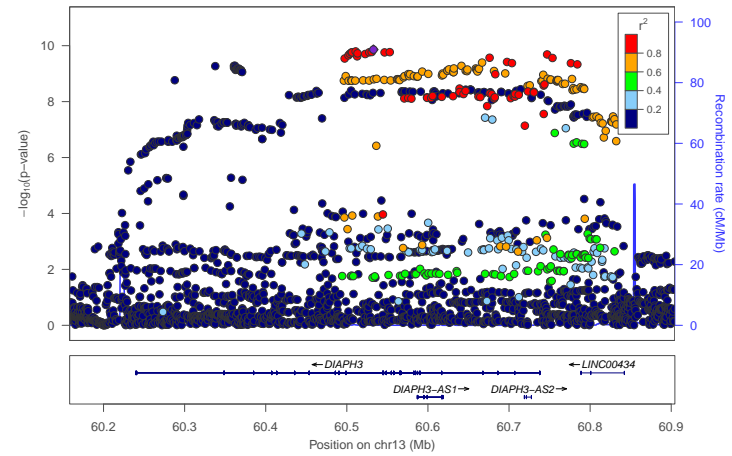
locus_154



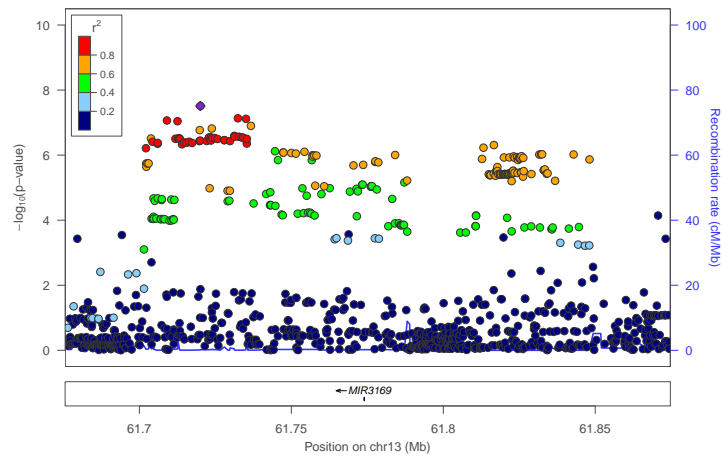
locus_155



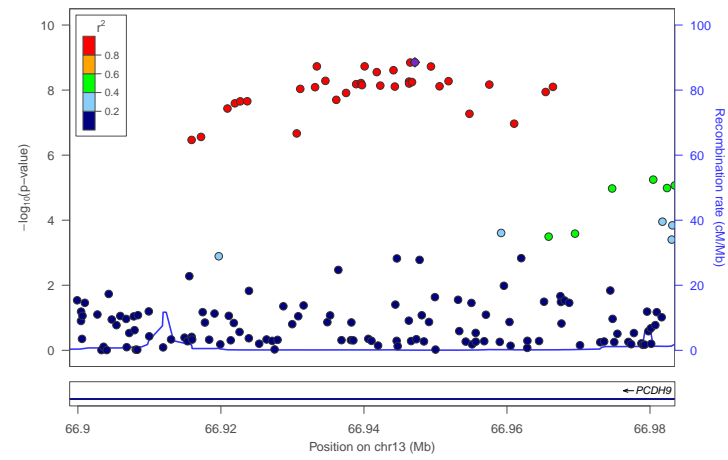
locus_156



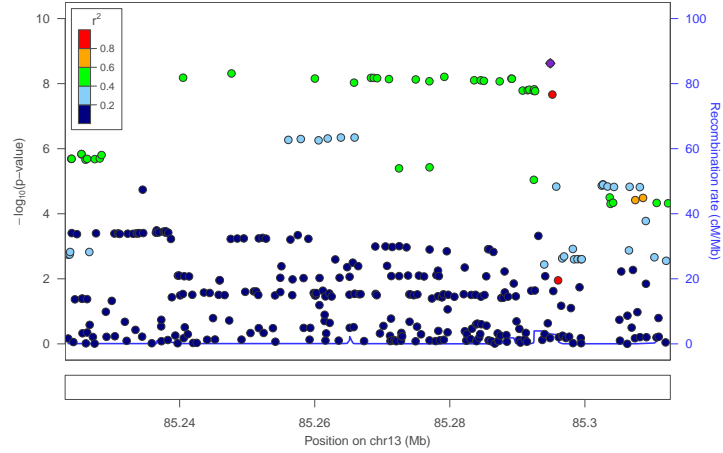
locus_157



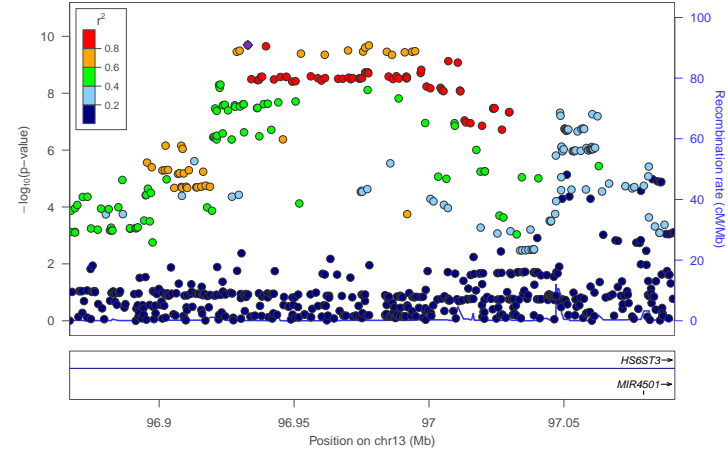
locus_158



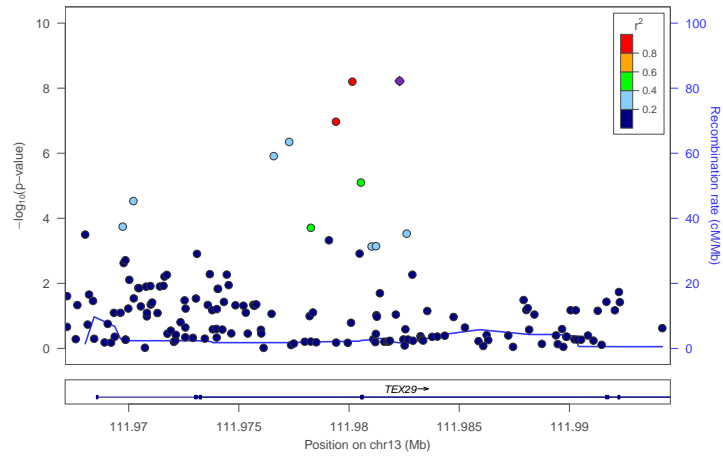
locus_159



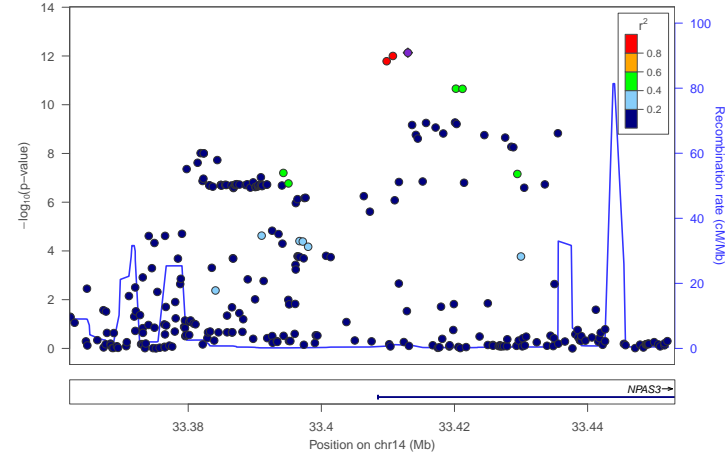
locus_160

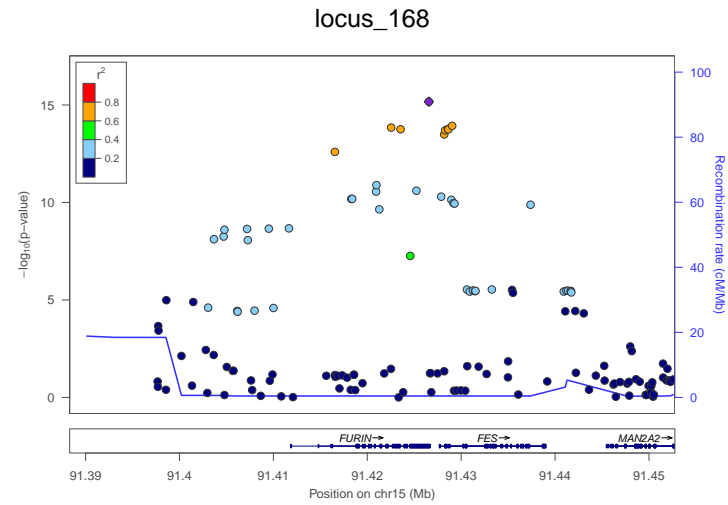
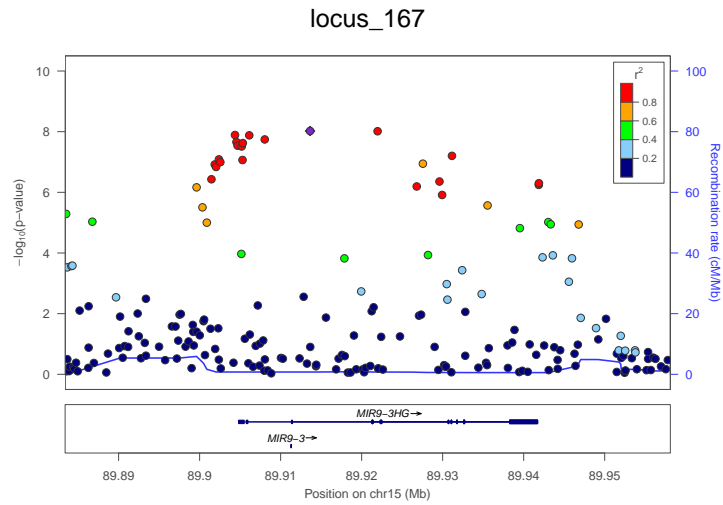
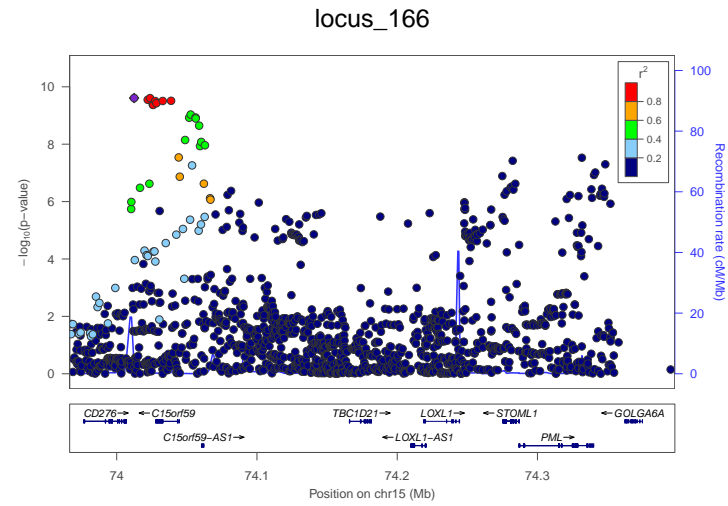
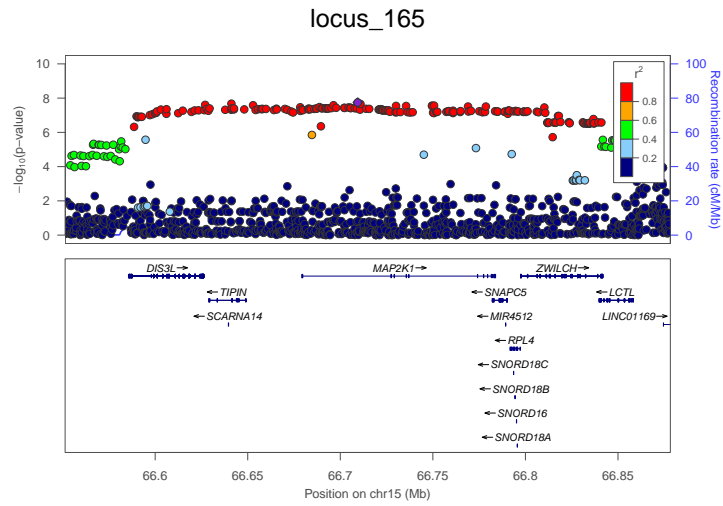
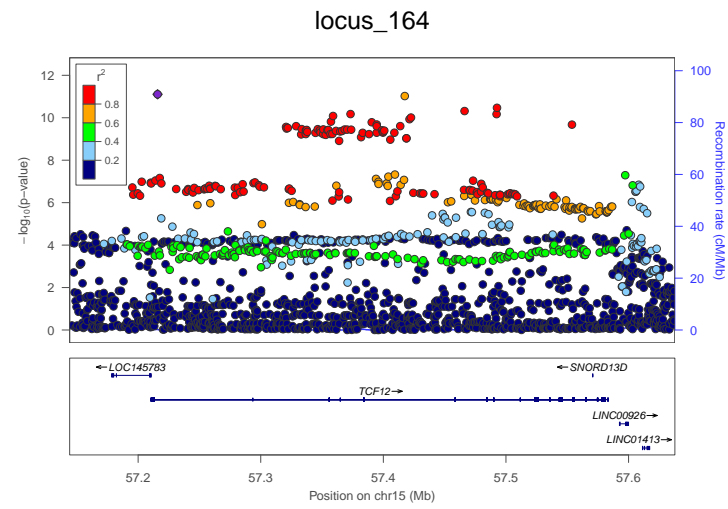
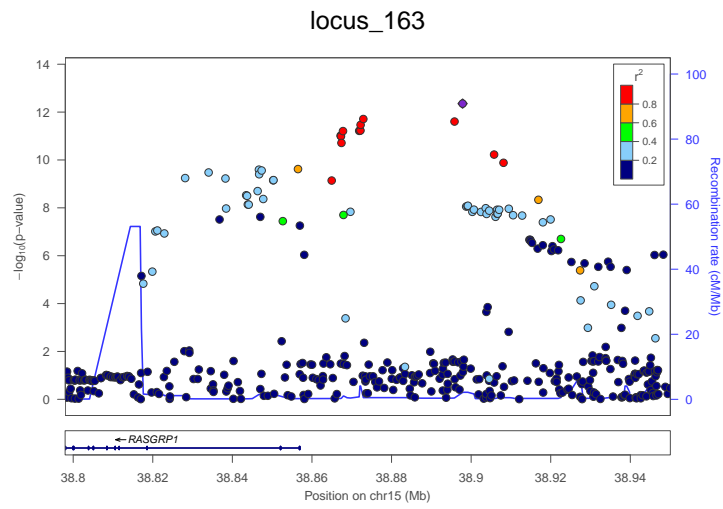


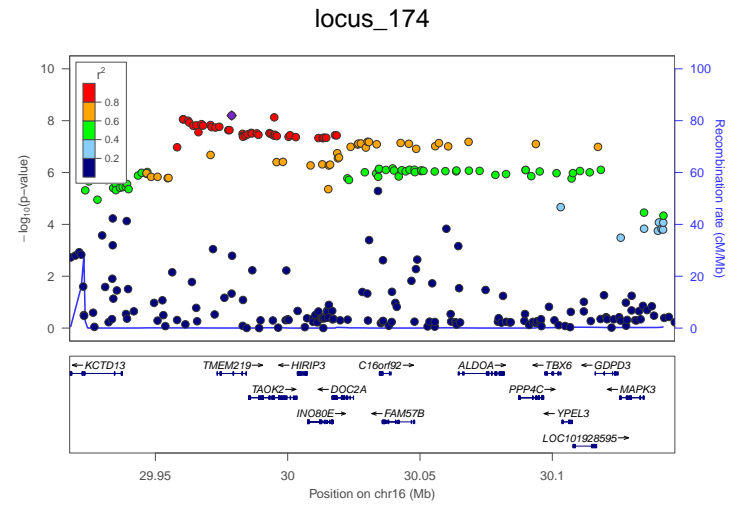
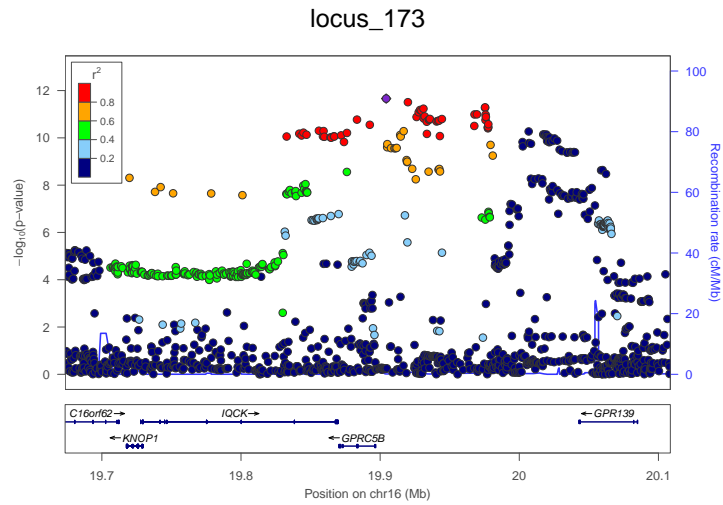
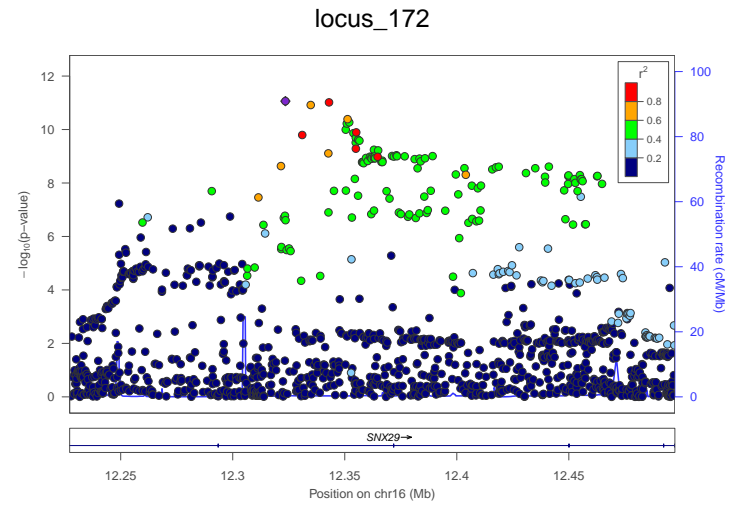
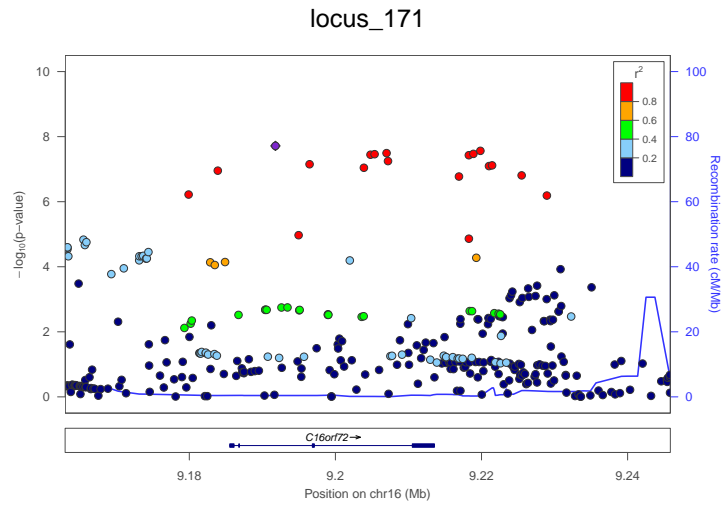
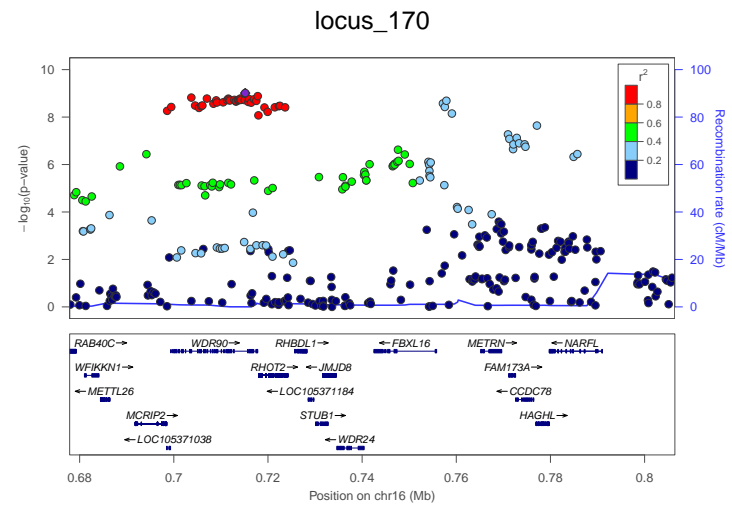
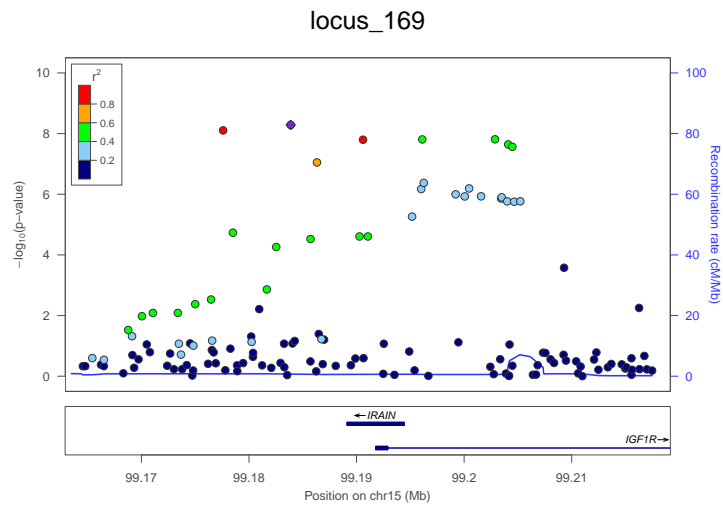
locus_161

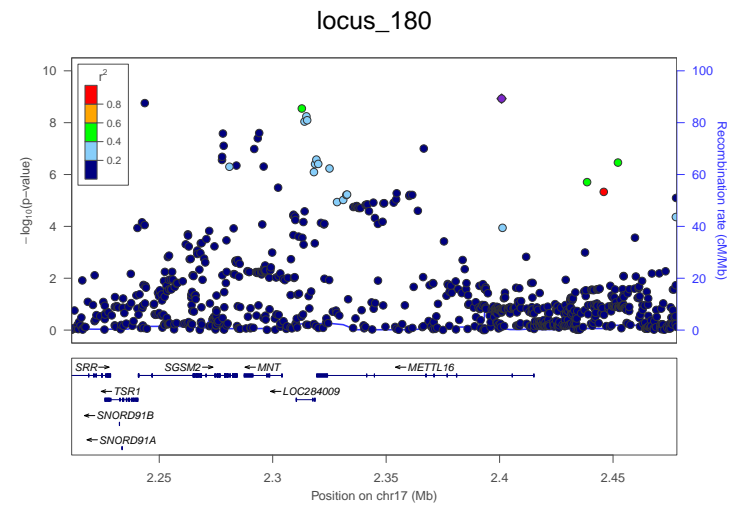
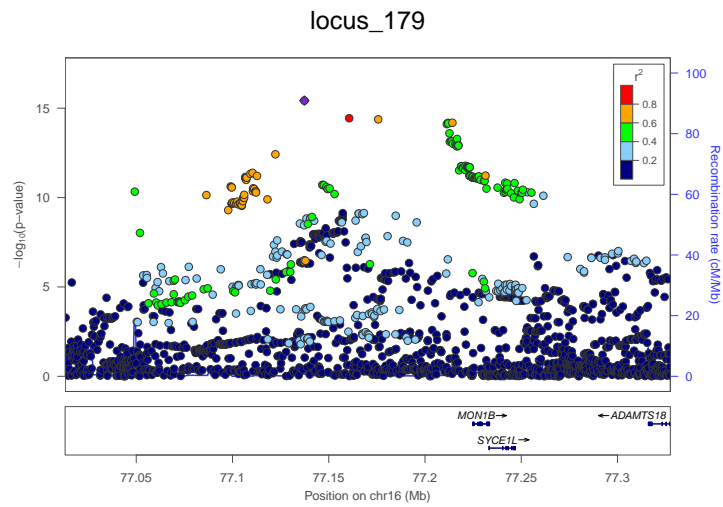
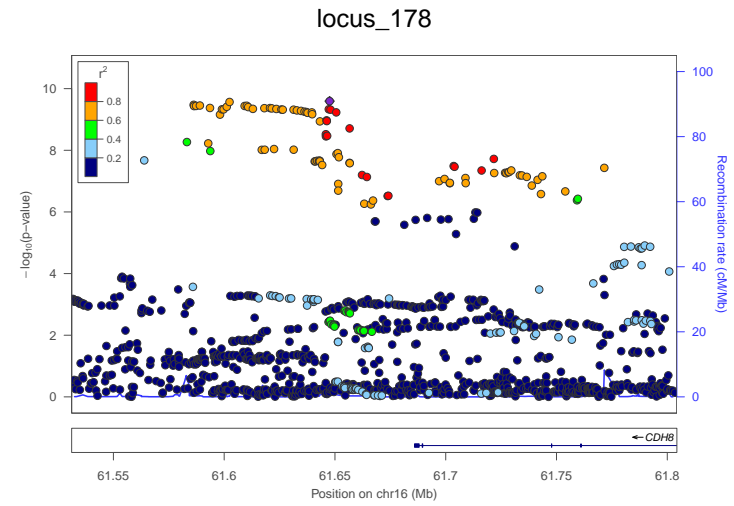
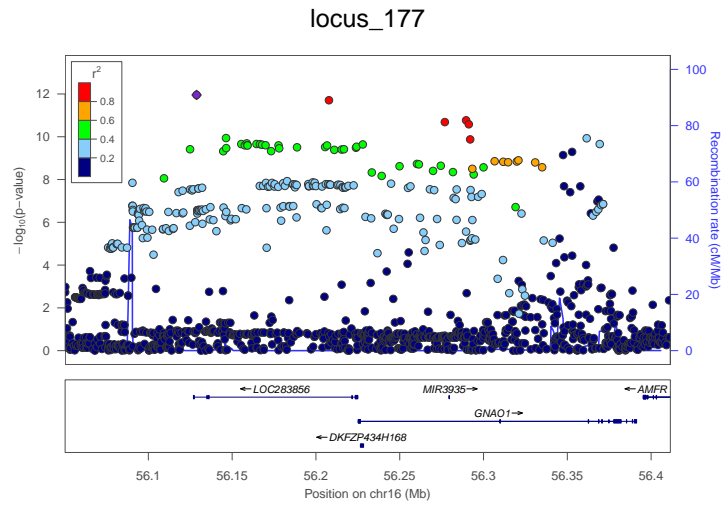
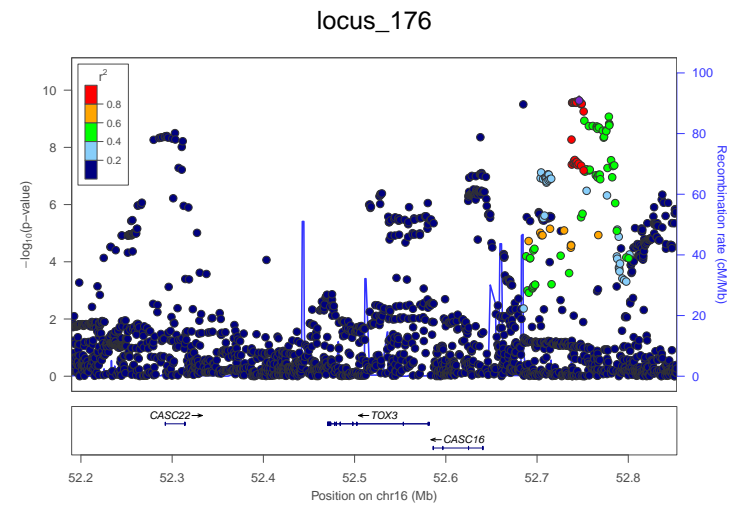
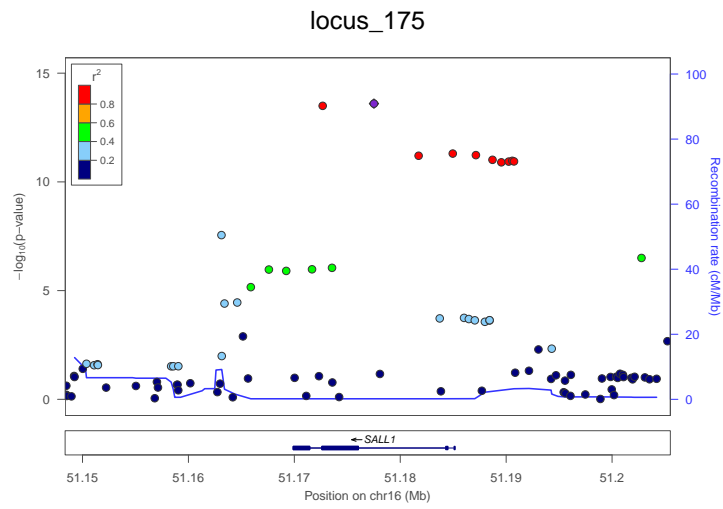


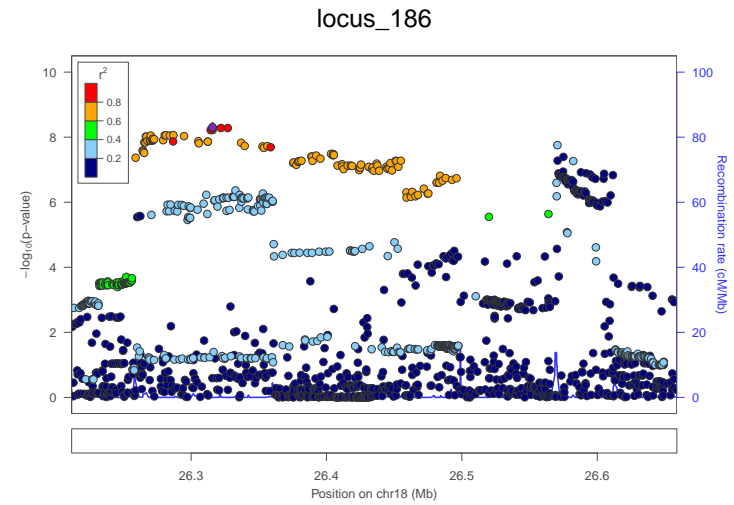
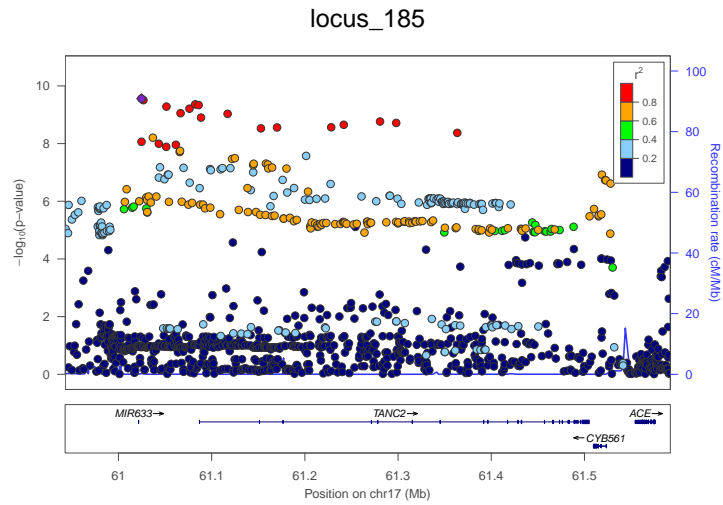
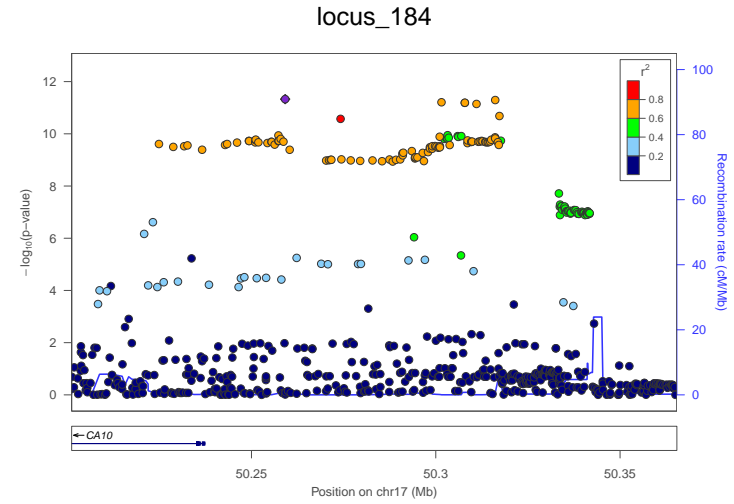
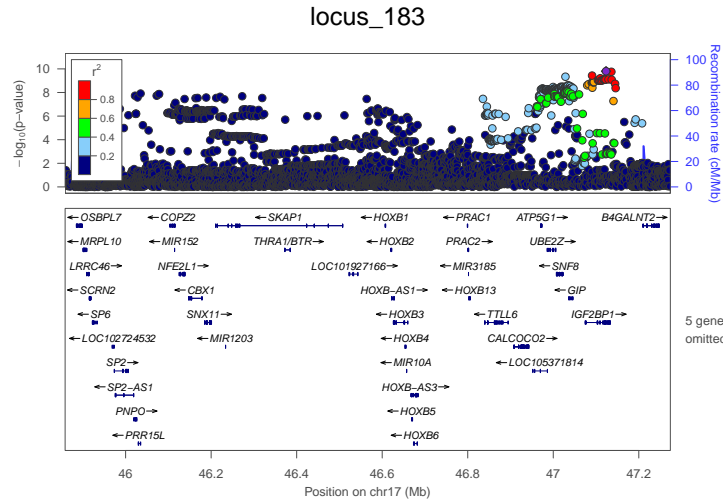
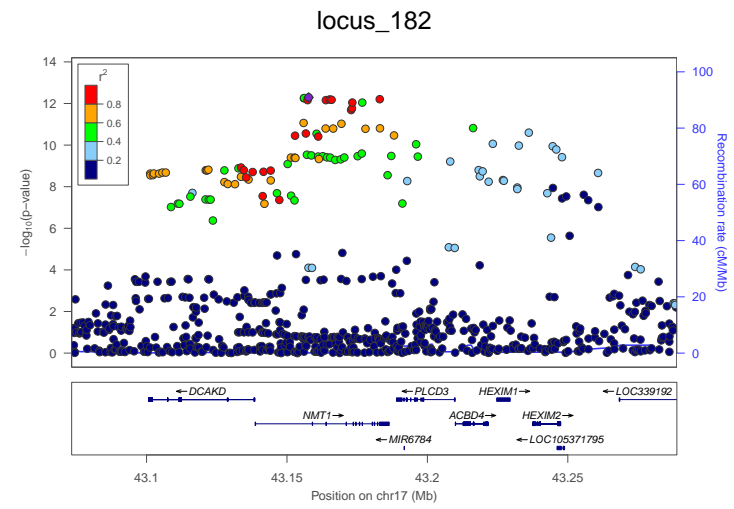
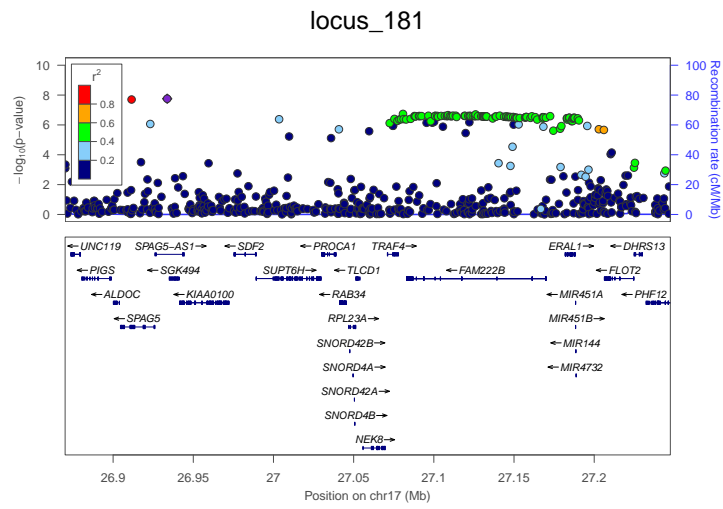
locus_162

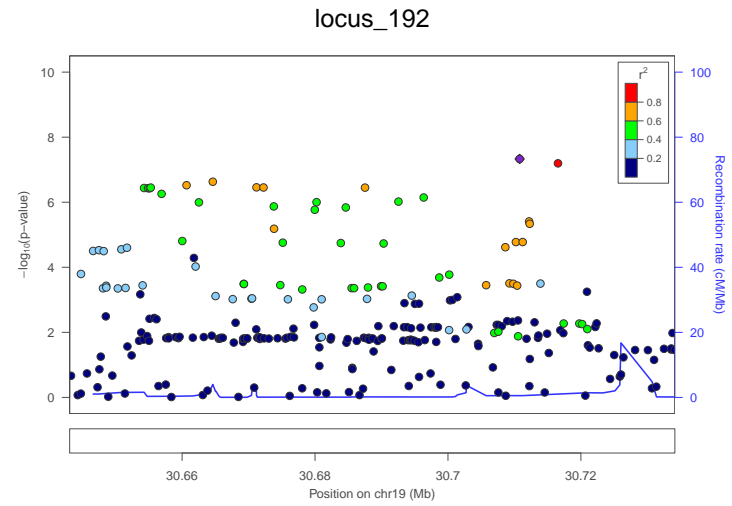
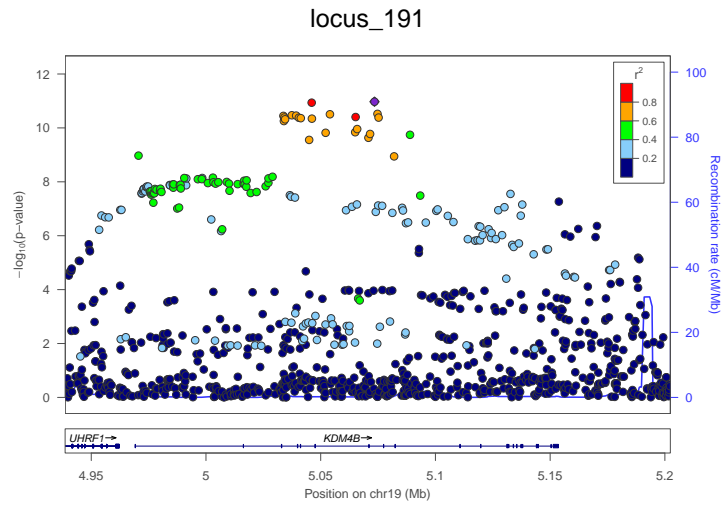
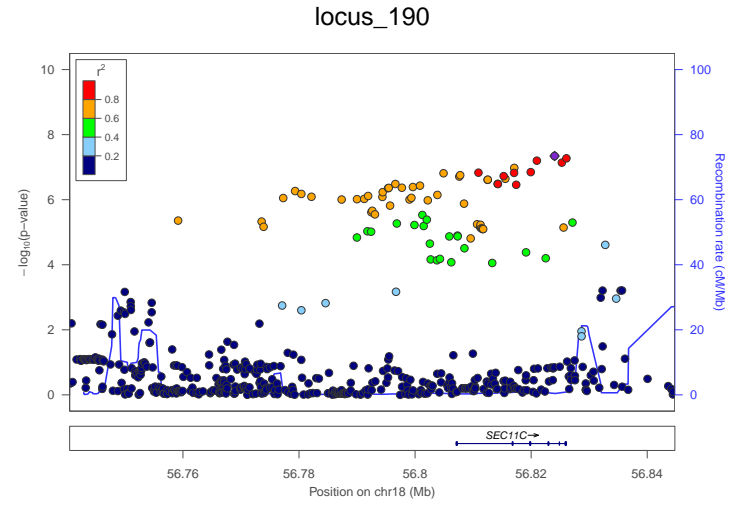
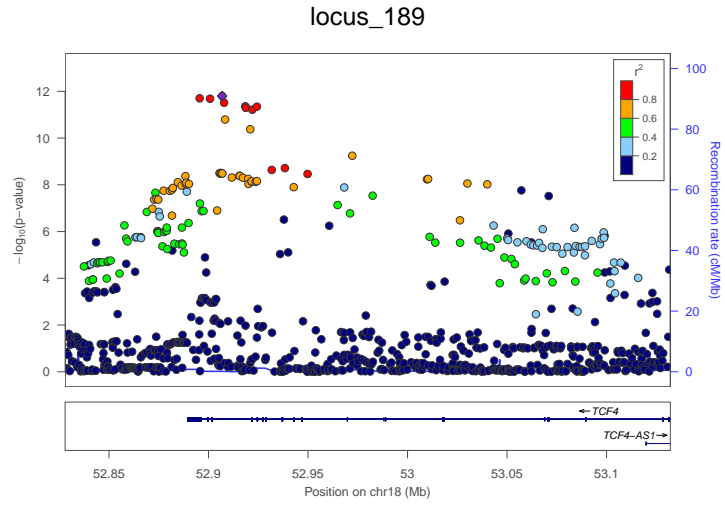
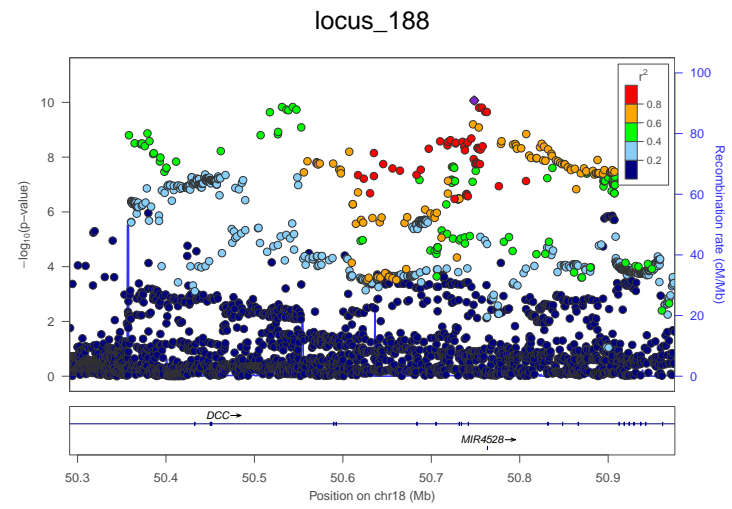
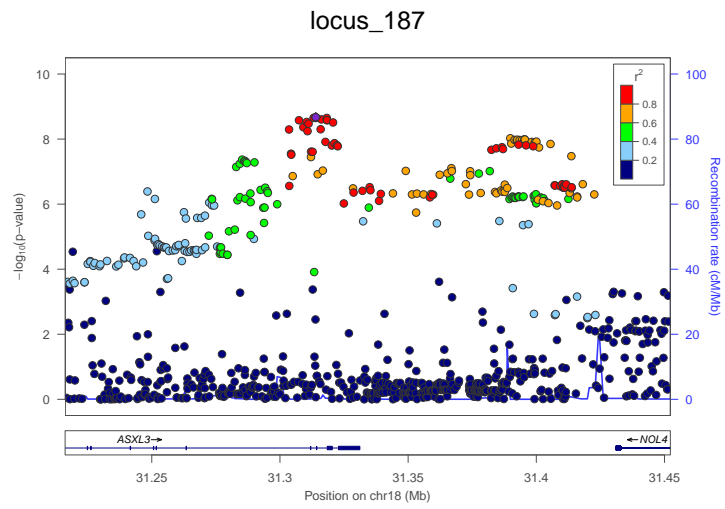




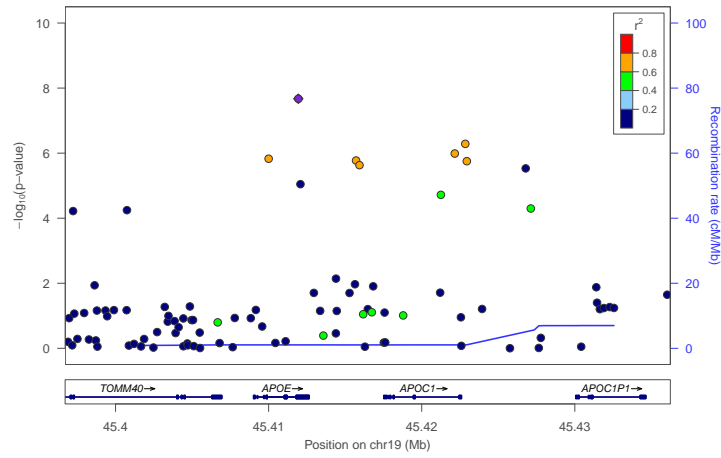




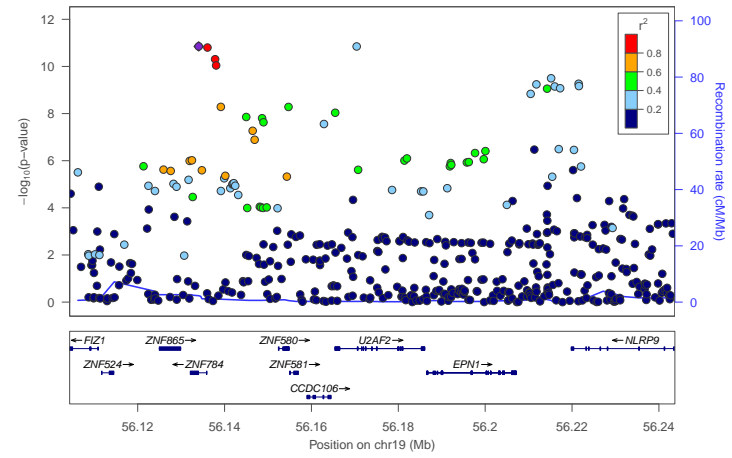




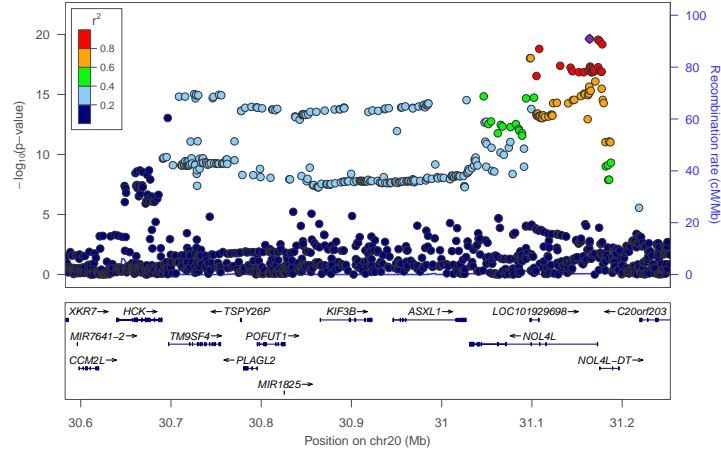
locus_193



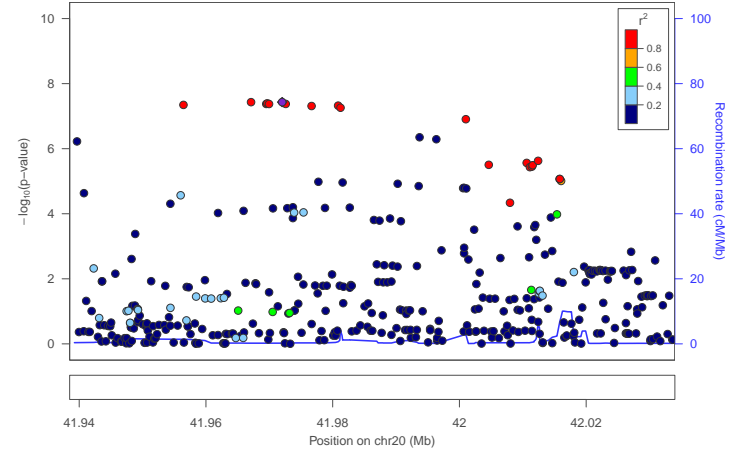
locus_194



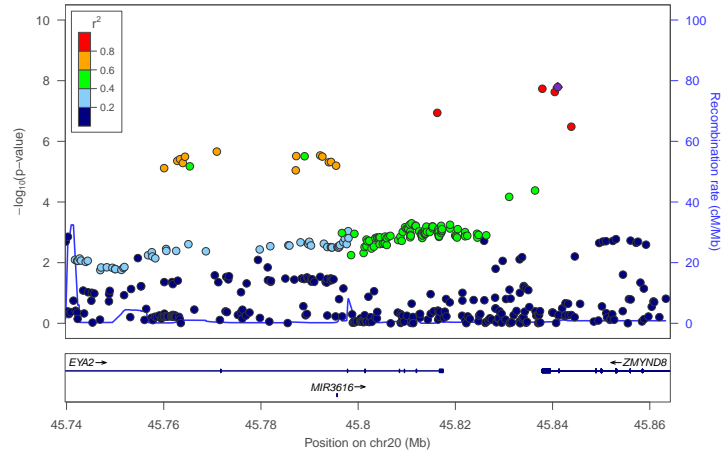
locus_195



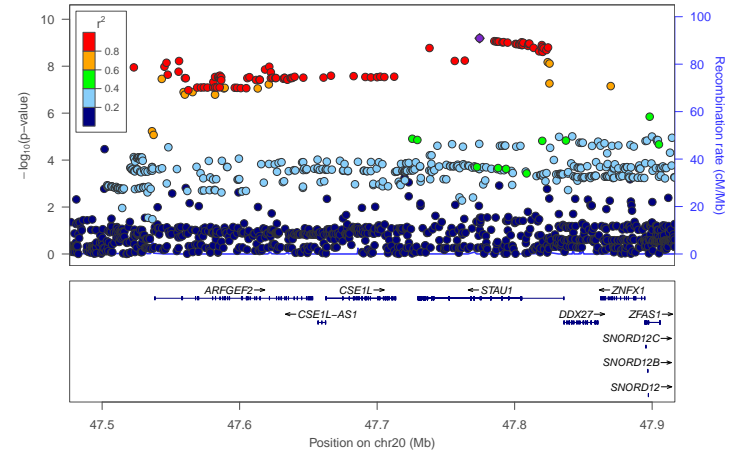
locus_196



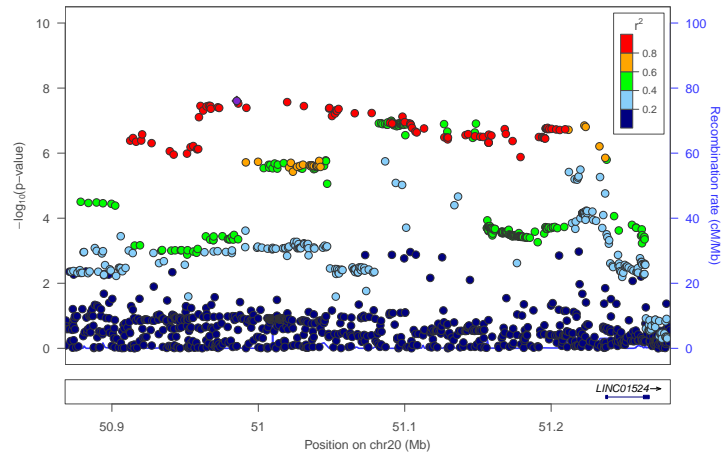
locus_197



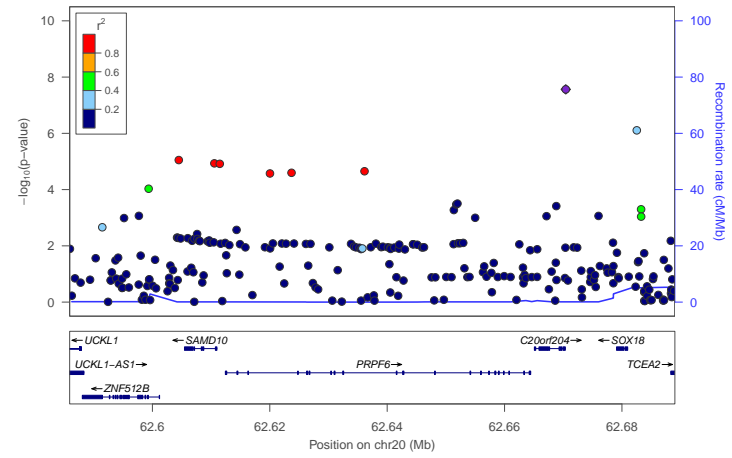
locus_198



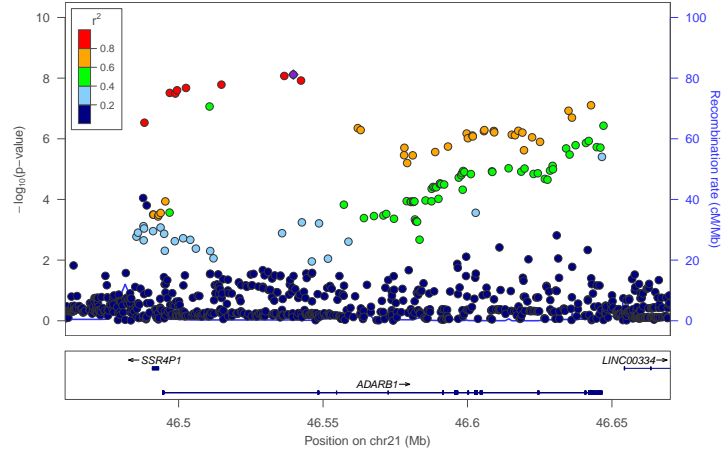
locus_199



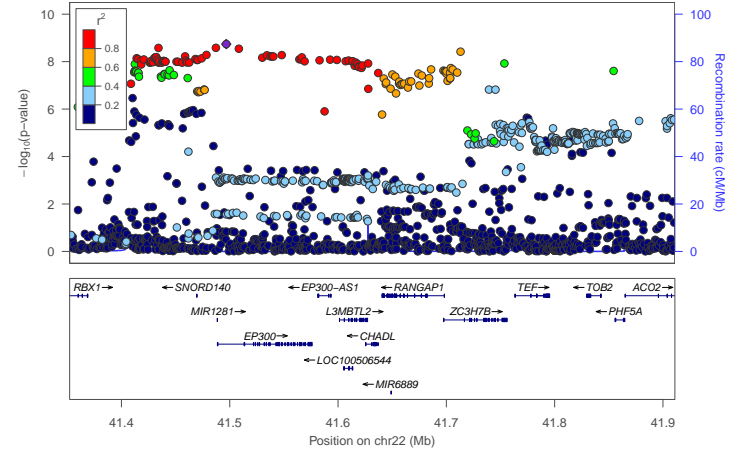
locus_200



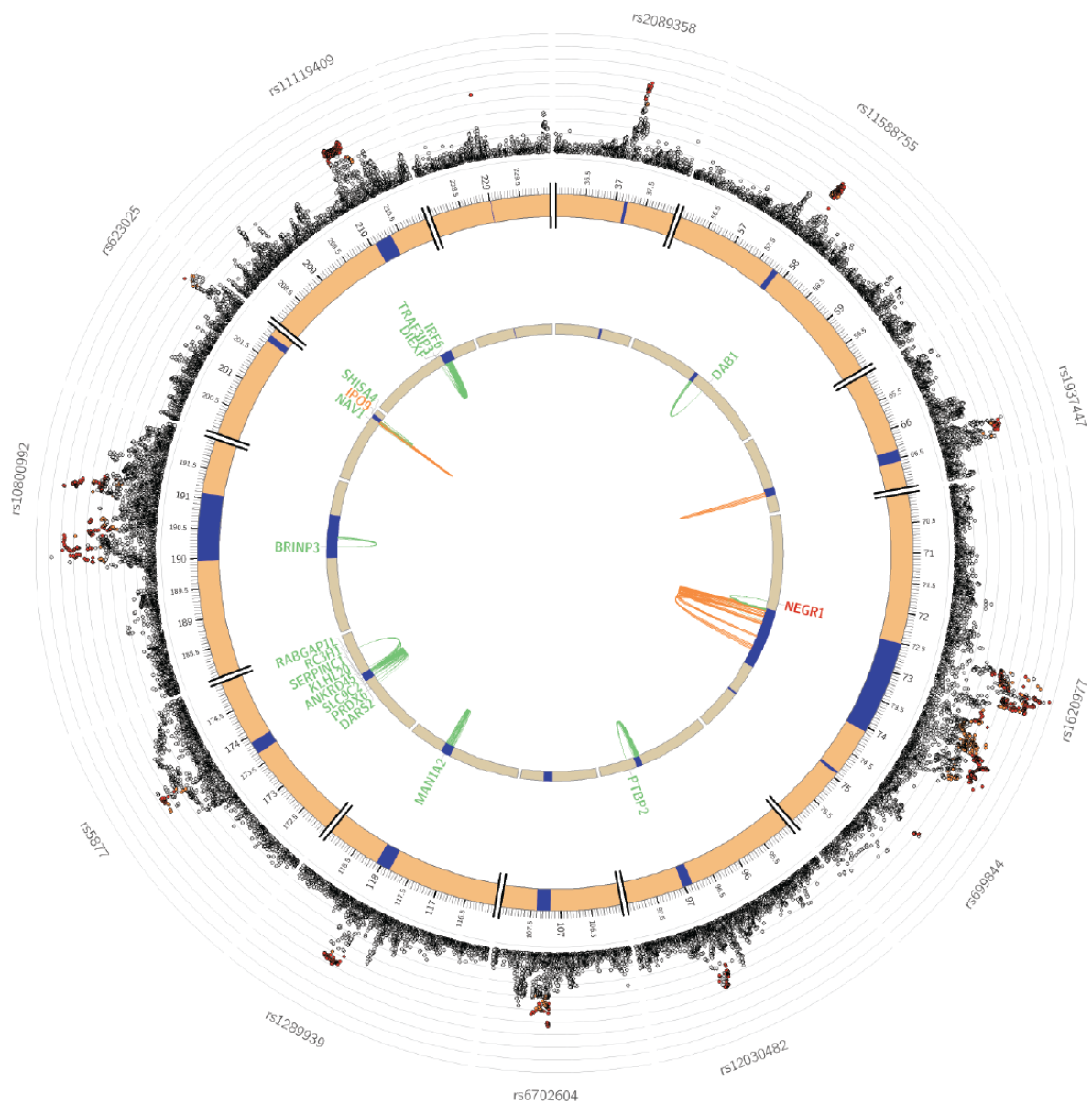
locus_201



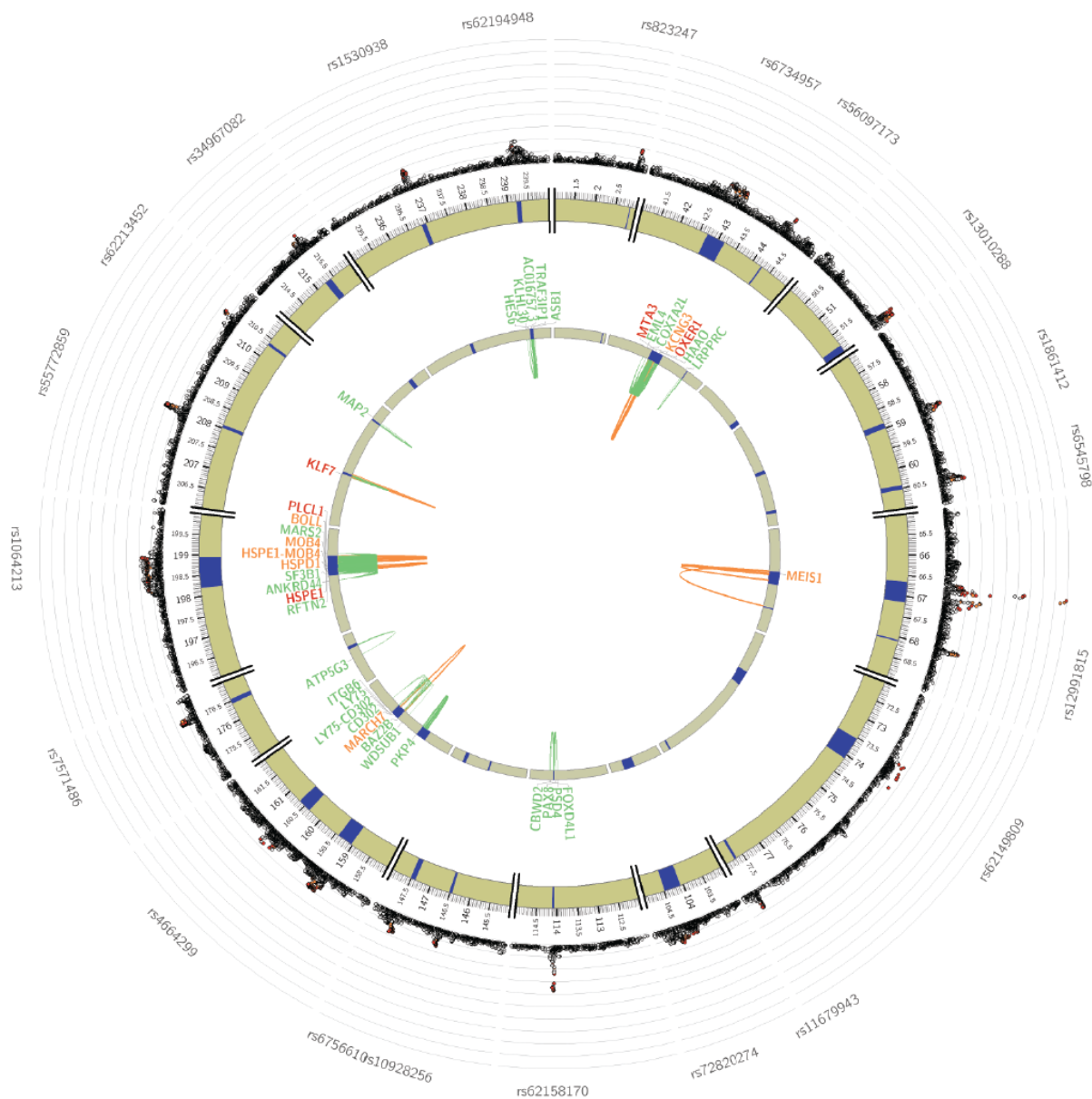
locus_202



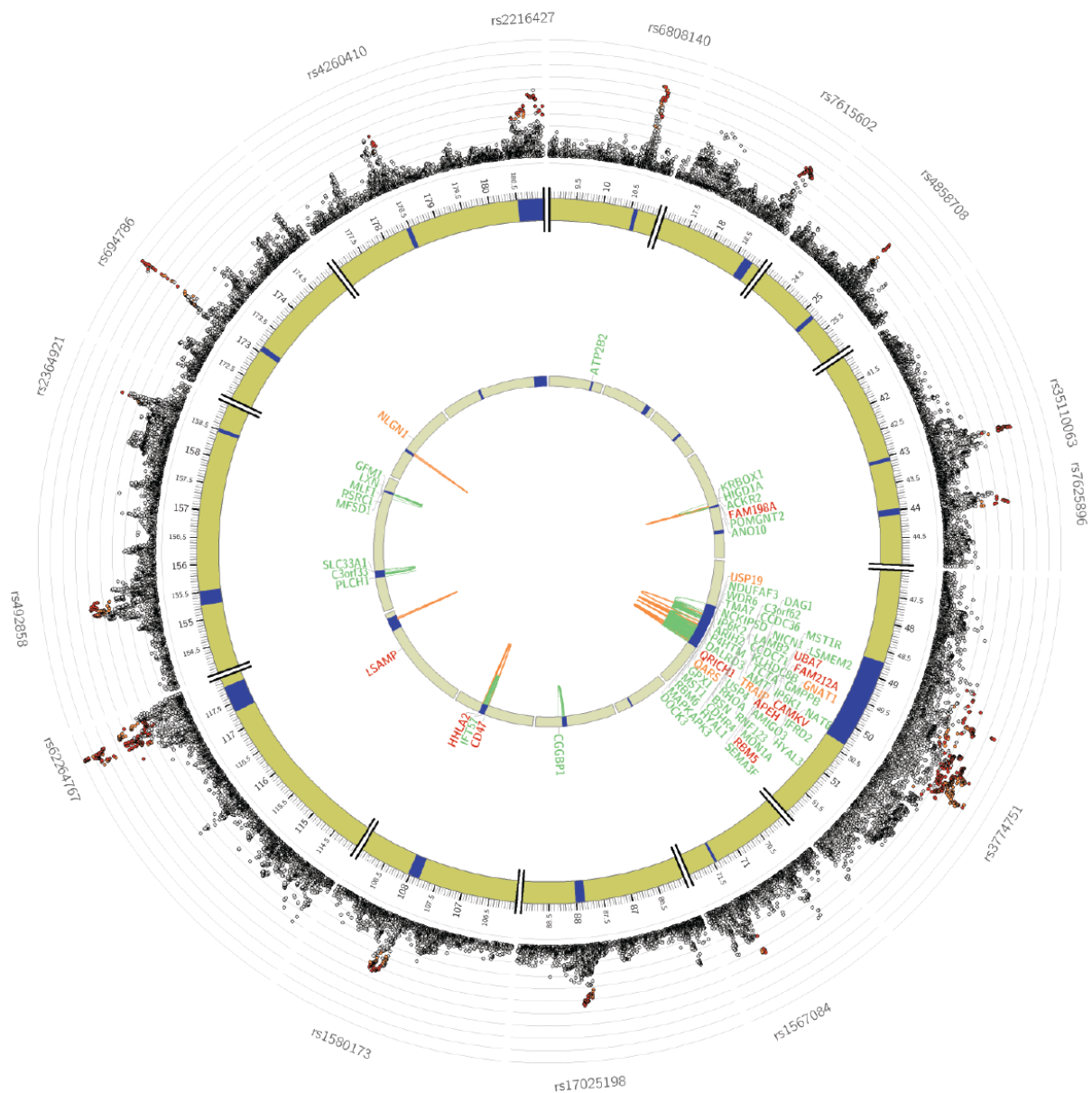
Chromosome 1



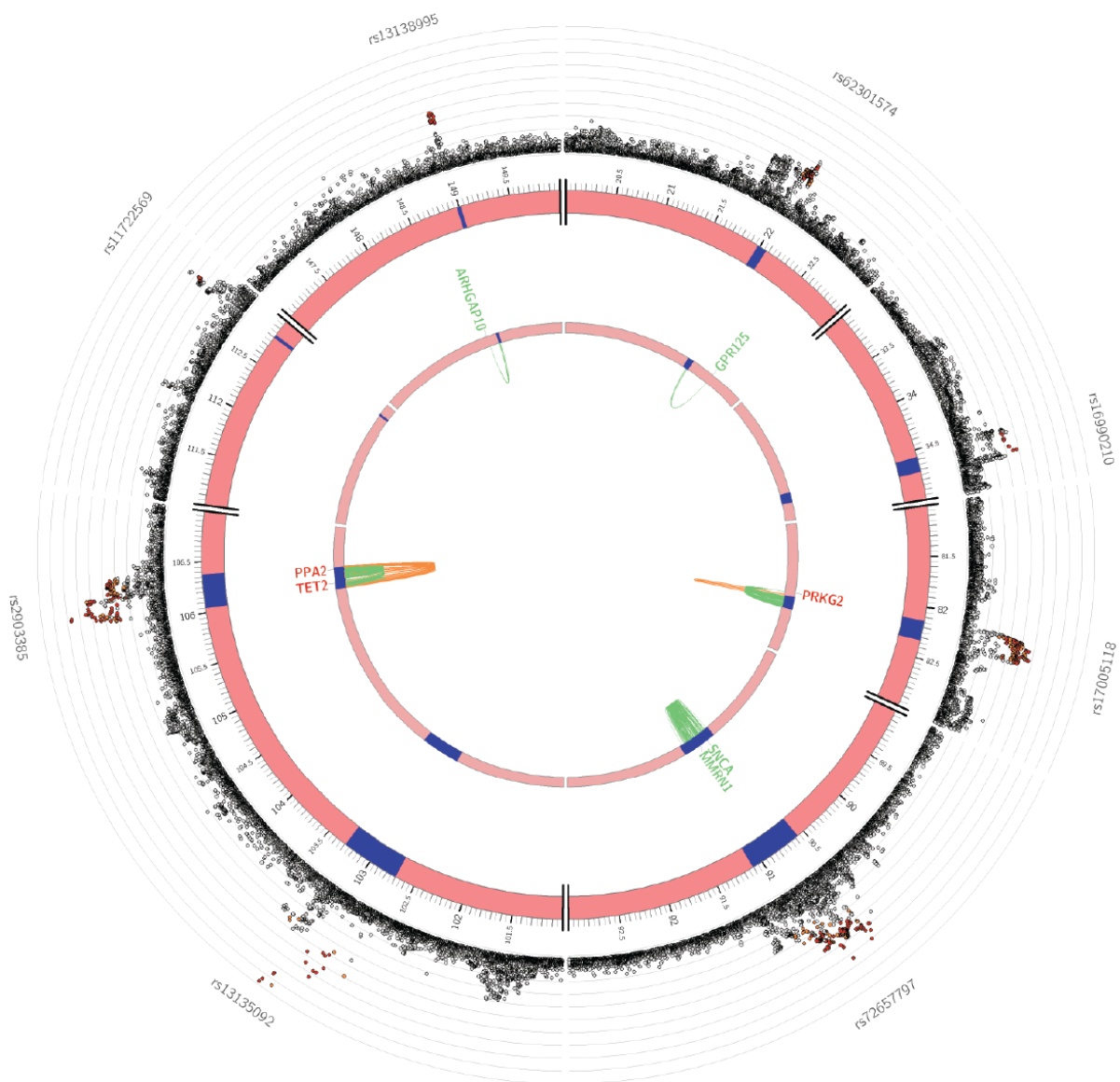
Chromosome 2



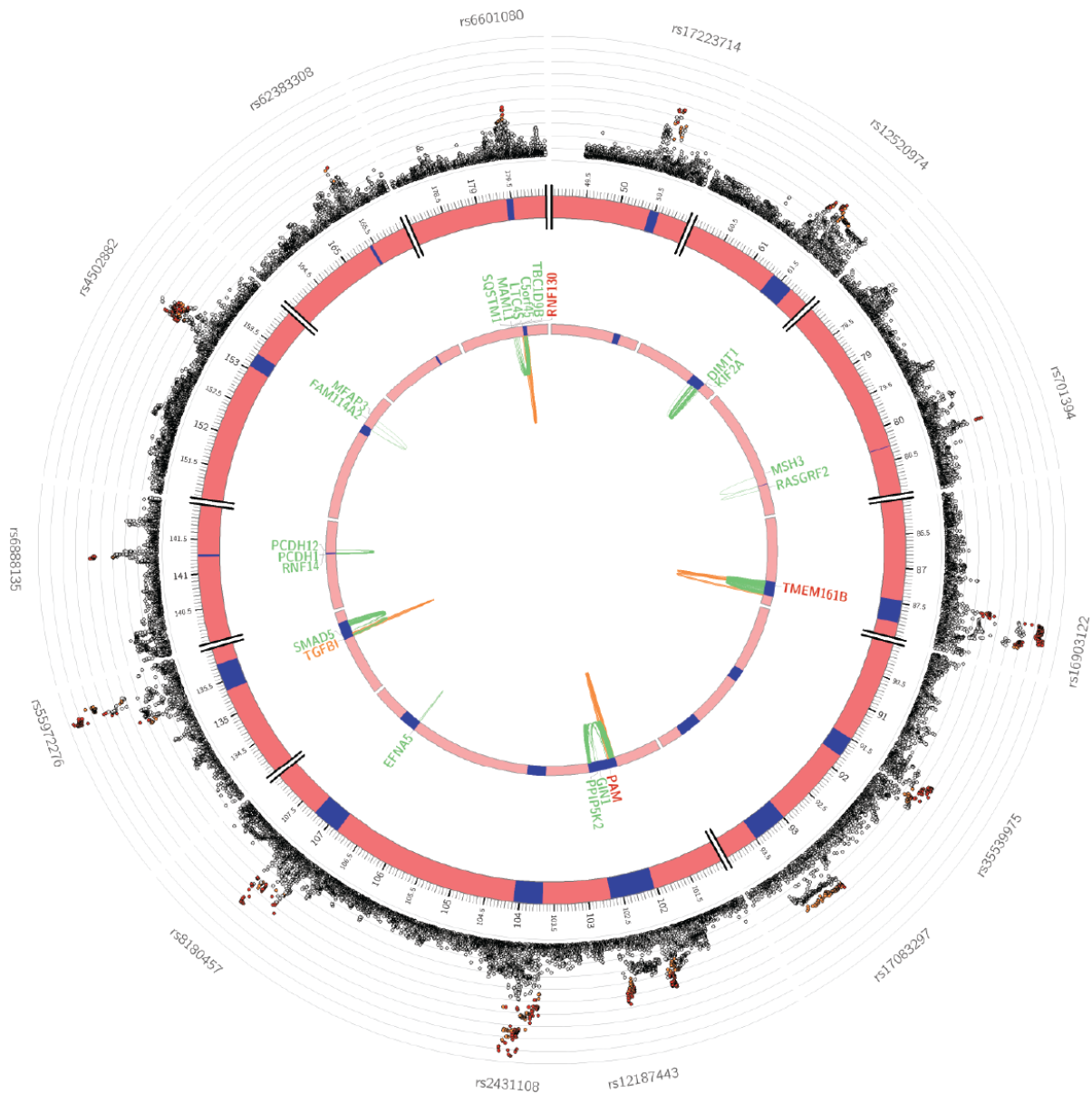
Chromosome 3



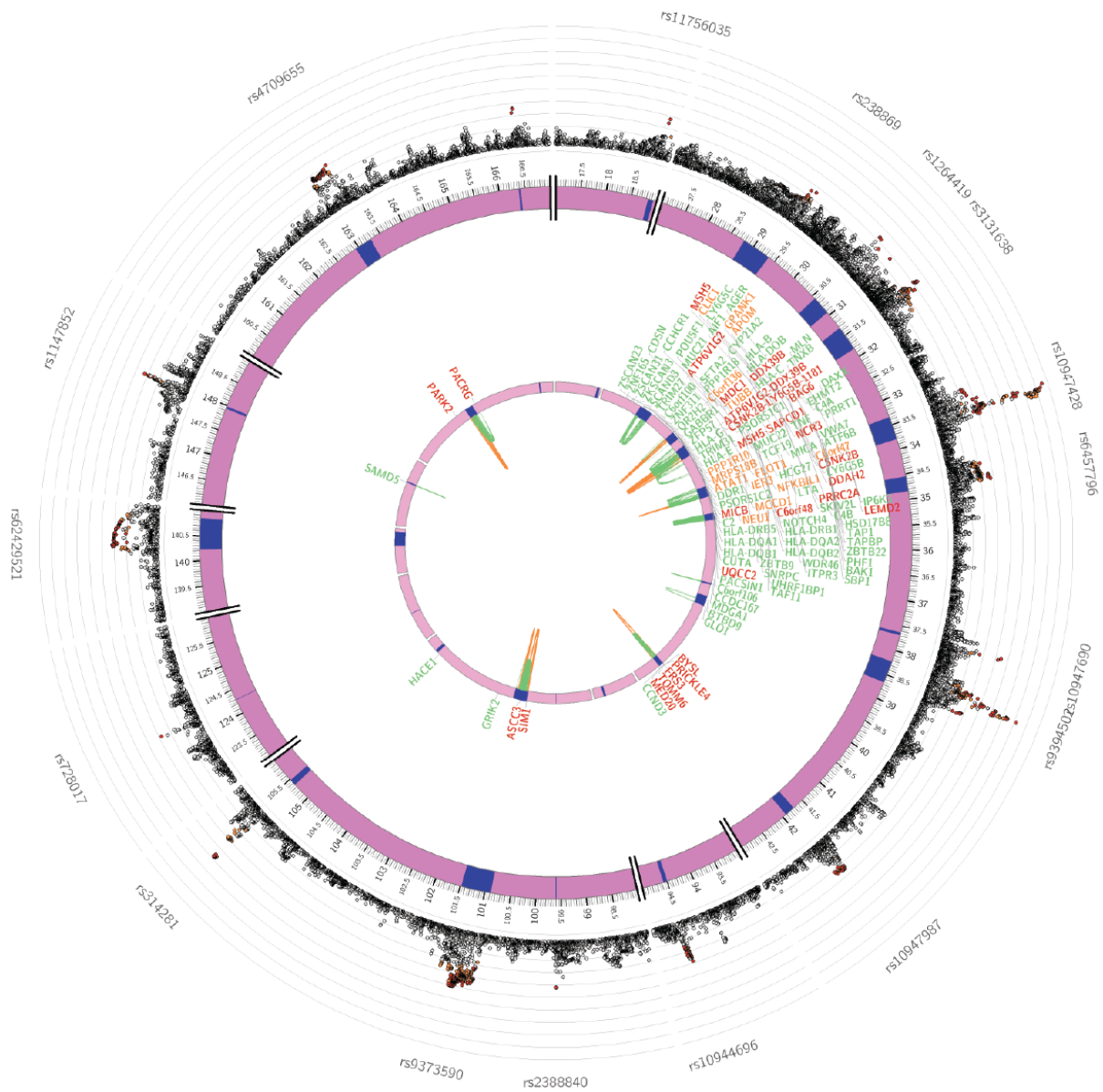
Chromosome 4



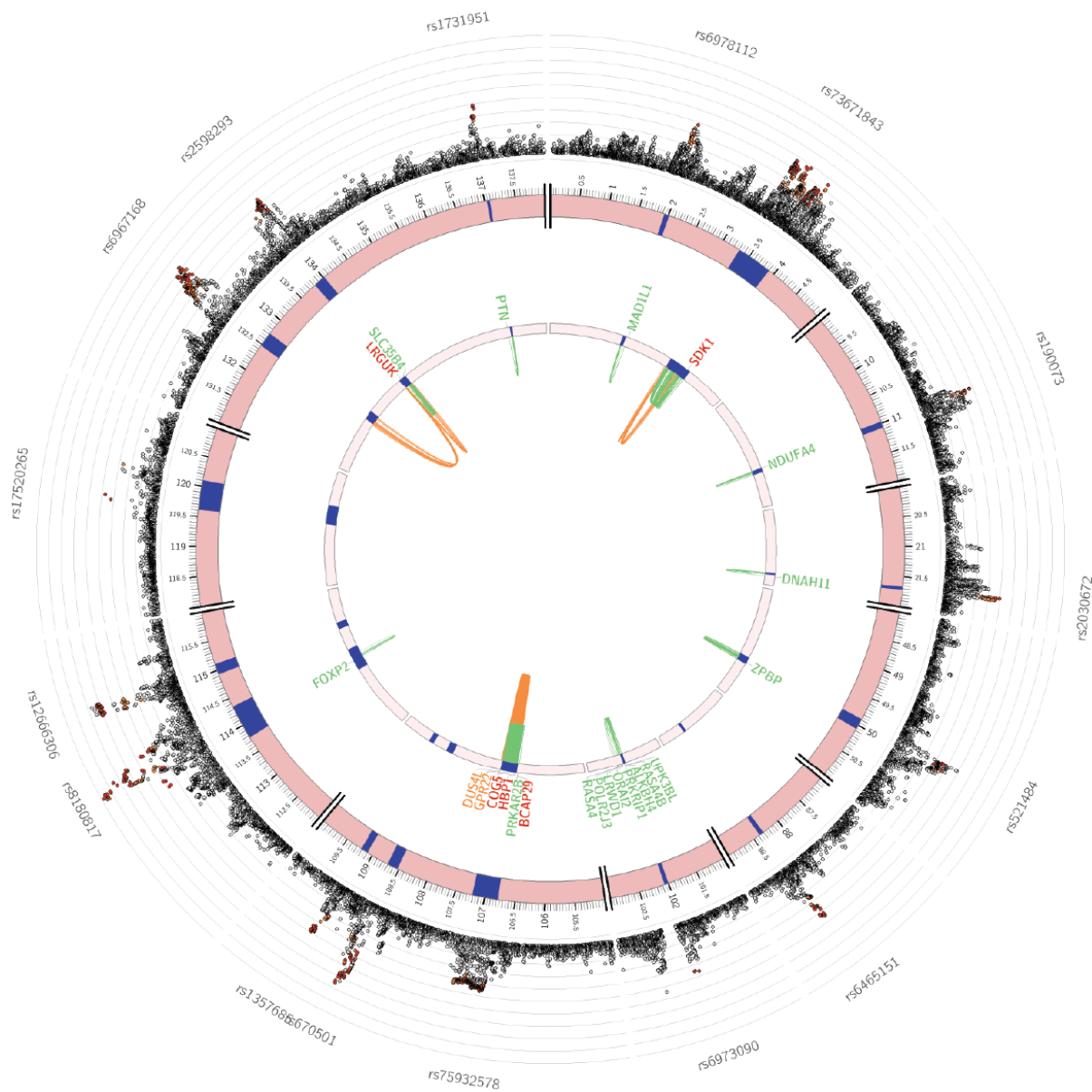
Chromosome 5



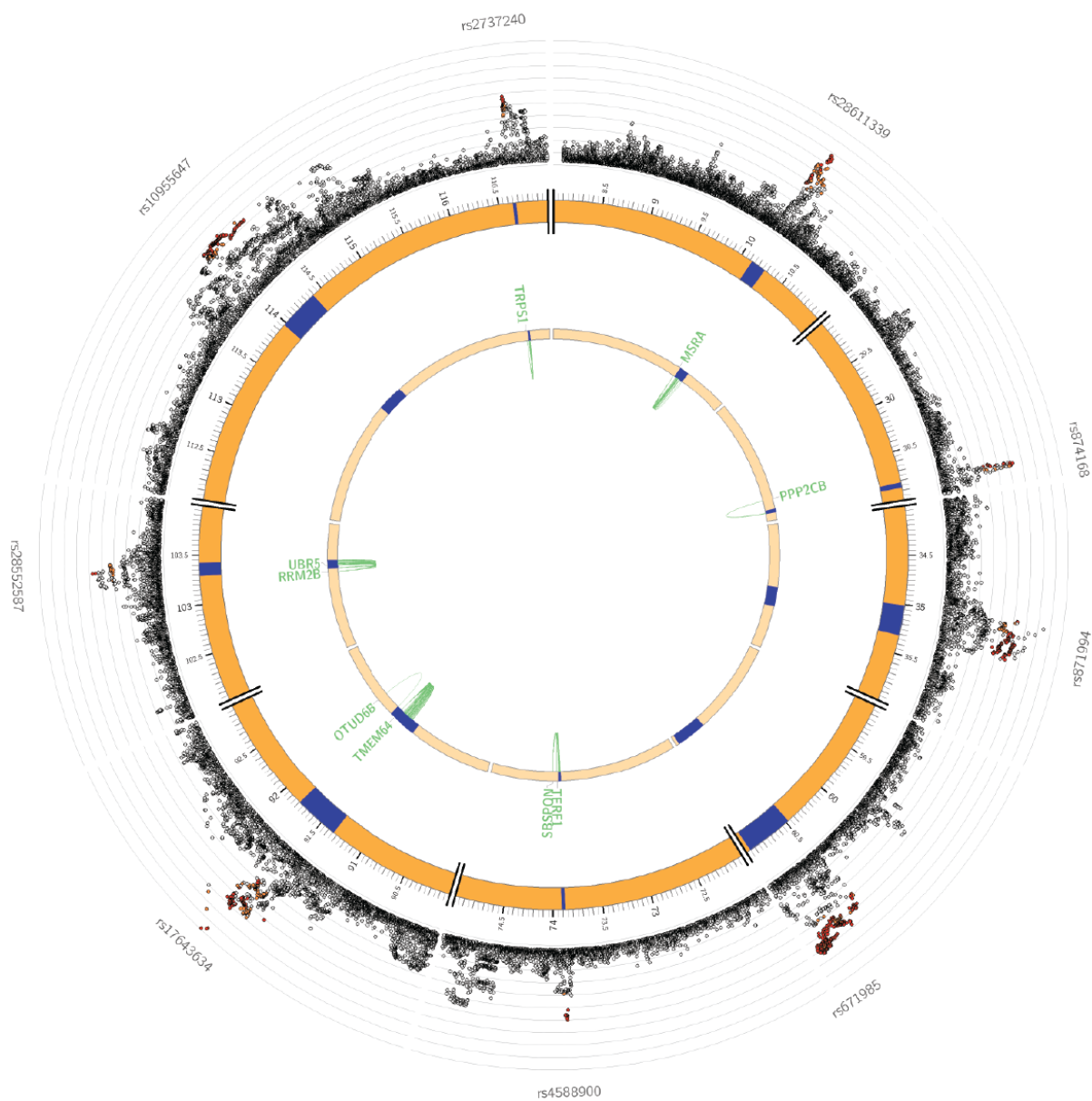
Chromosome 6



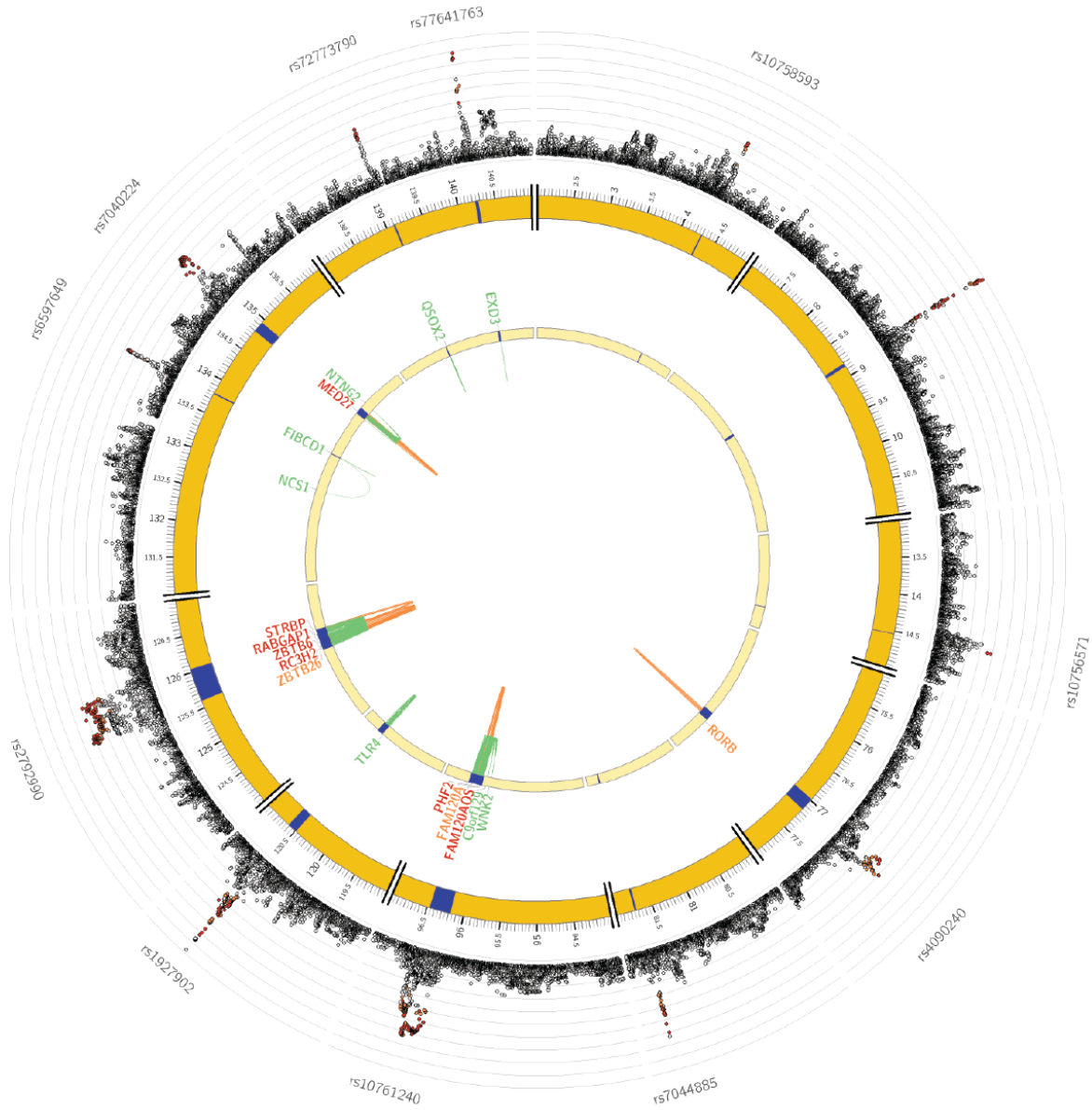
Chromosome 7



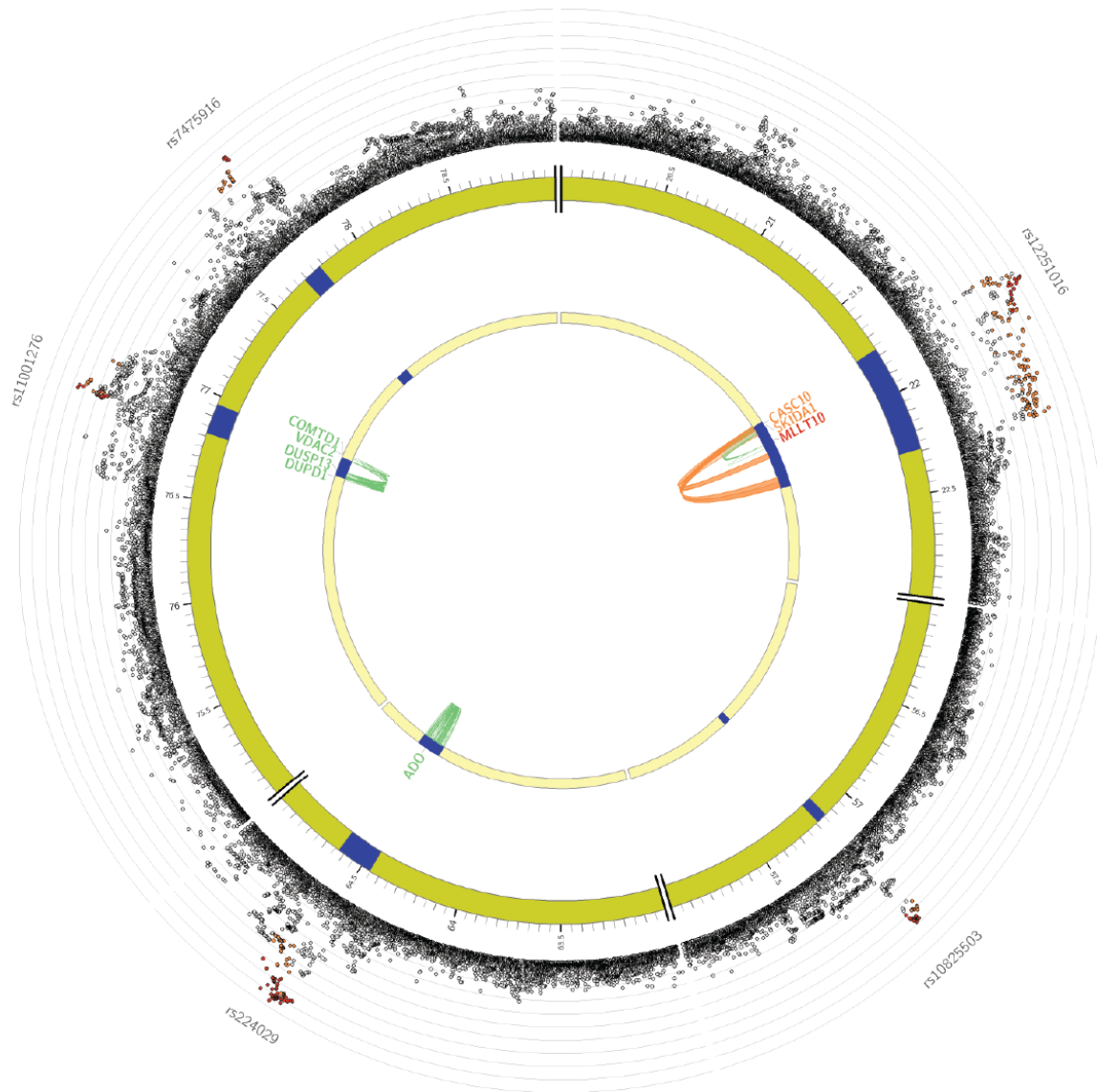
Chromosome 8



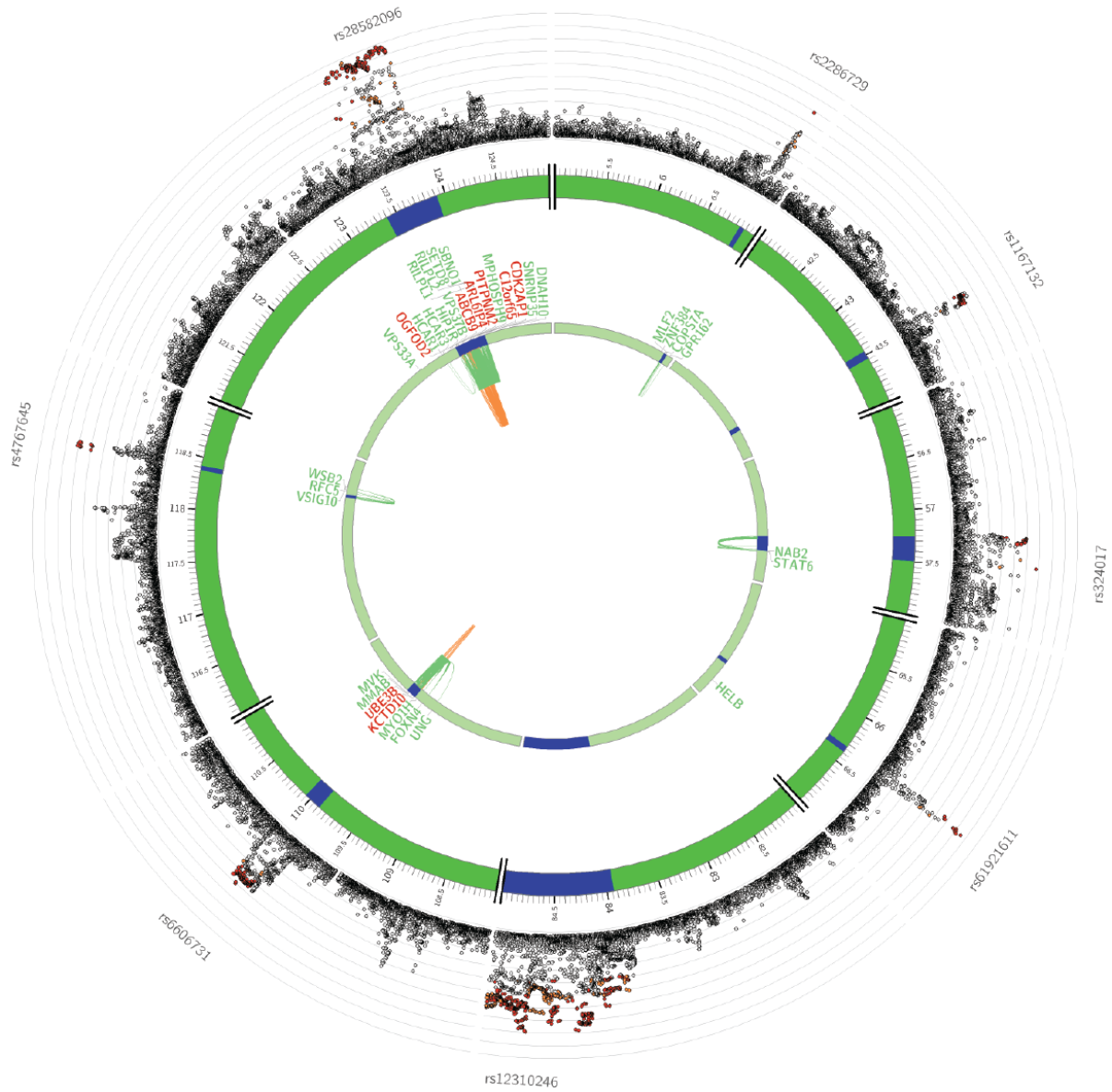
Chromosome 9



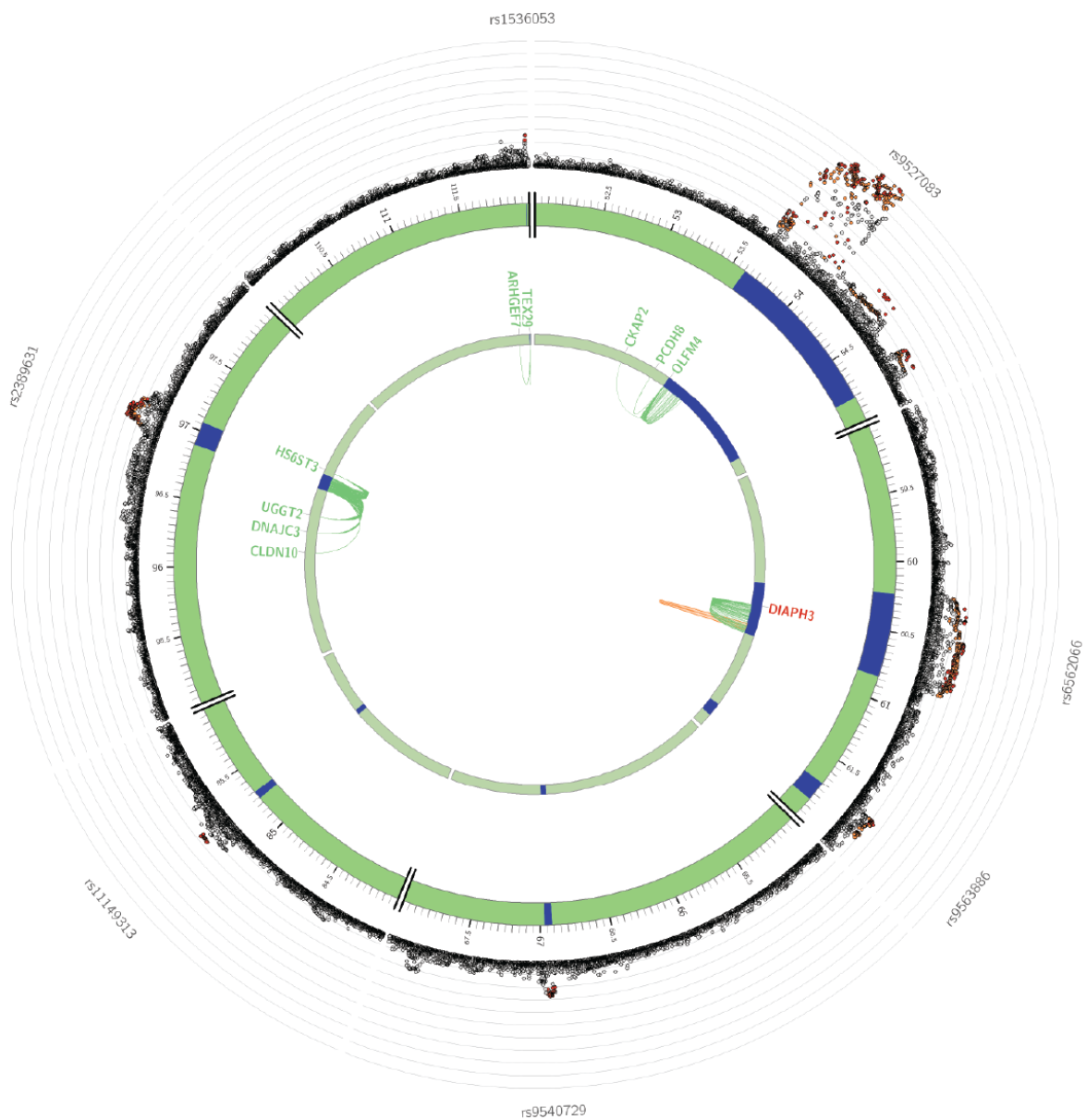
Chromosome 10



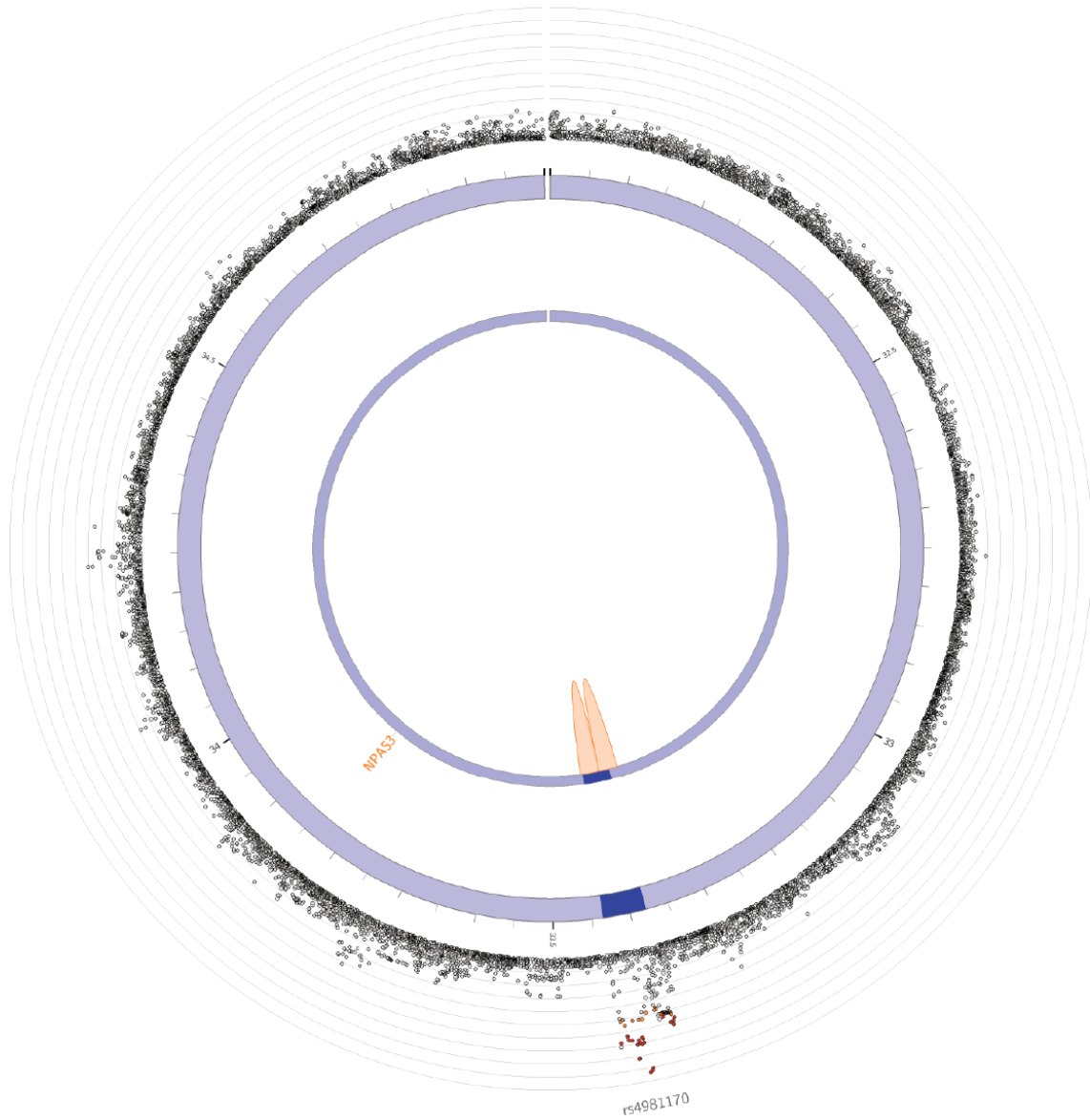
Chromosome 12



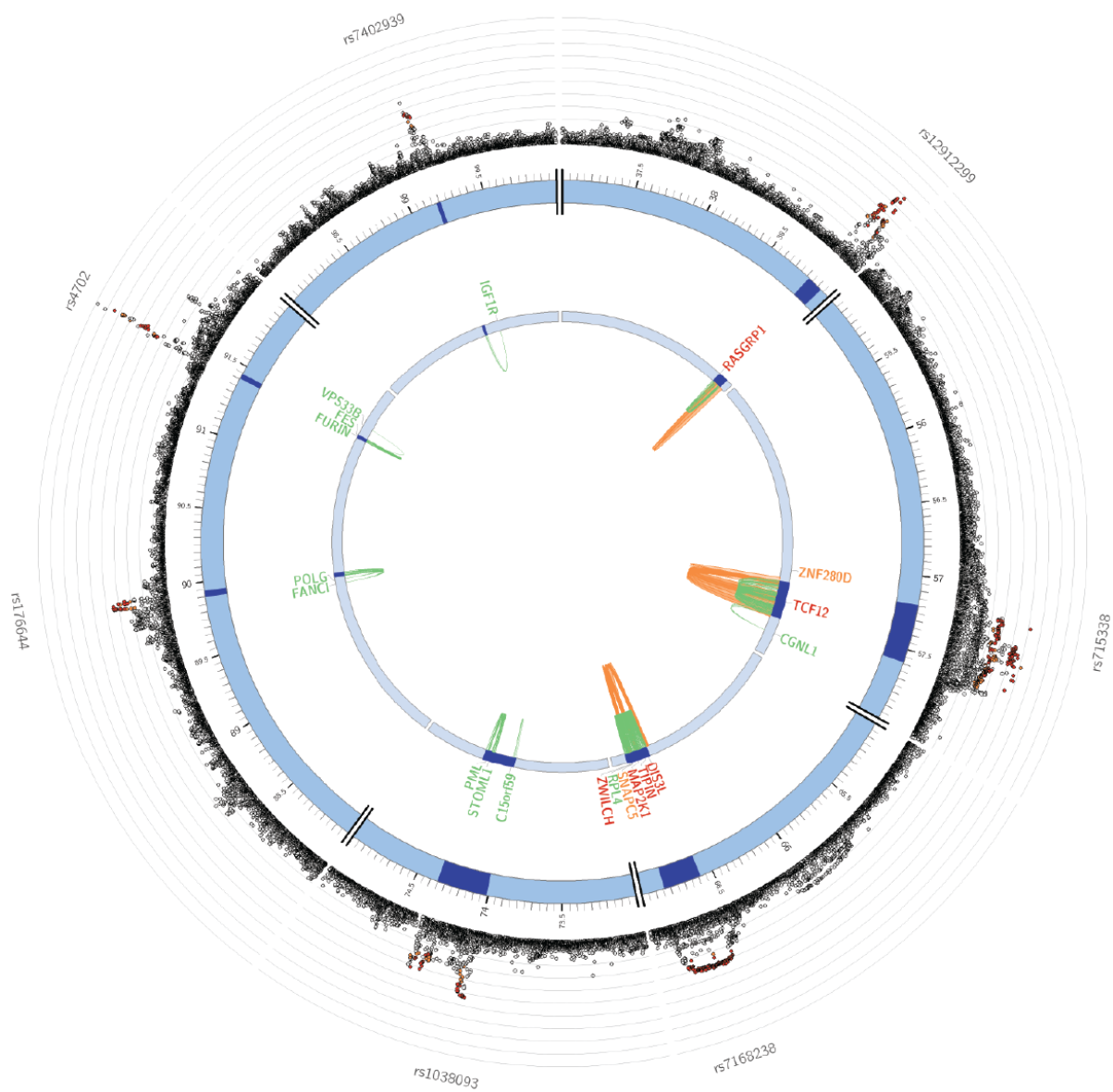
Chromosome 13



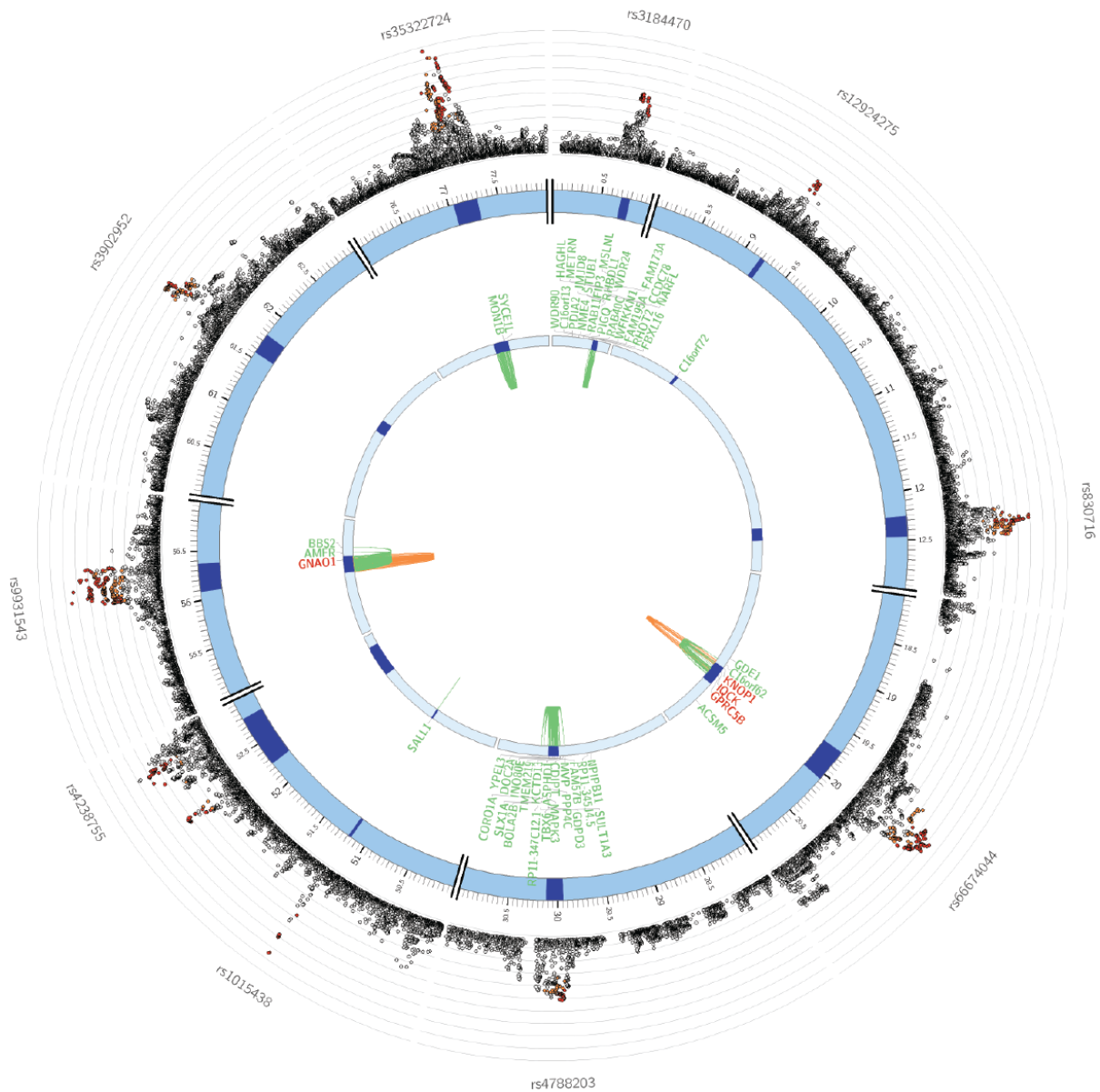
Chromosome 14



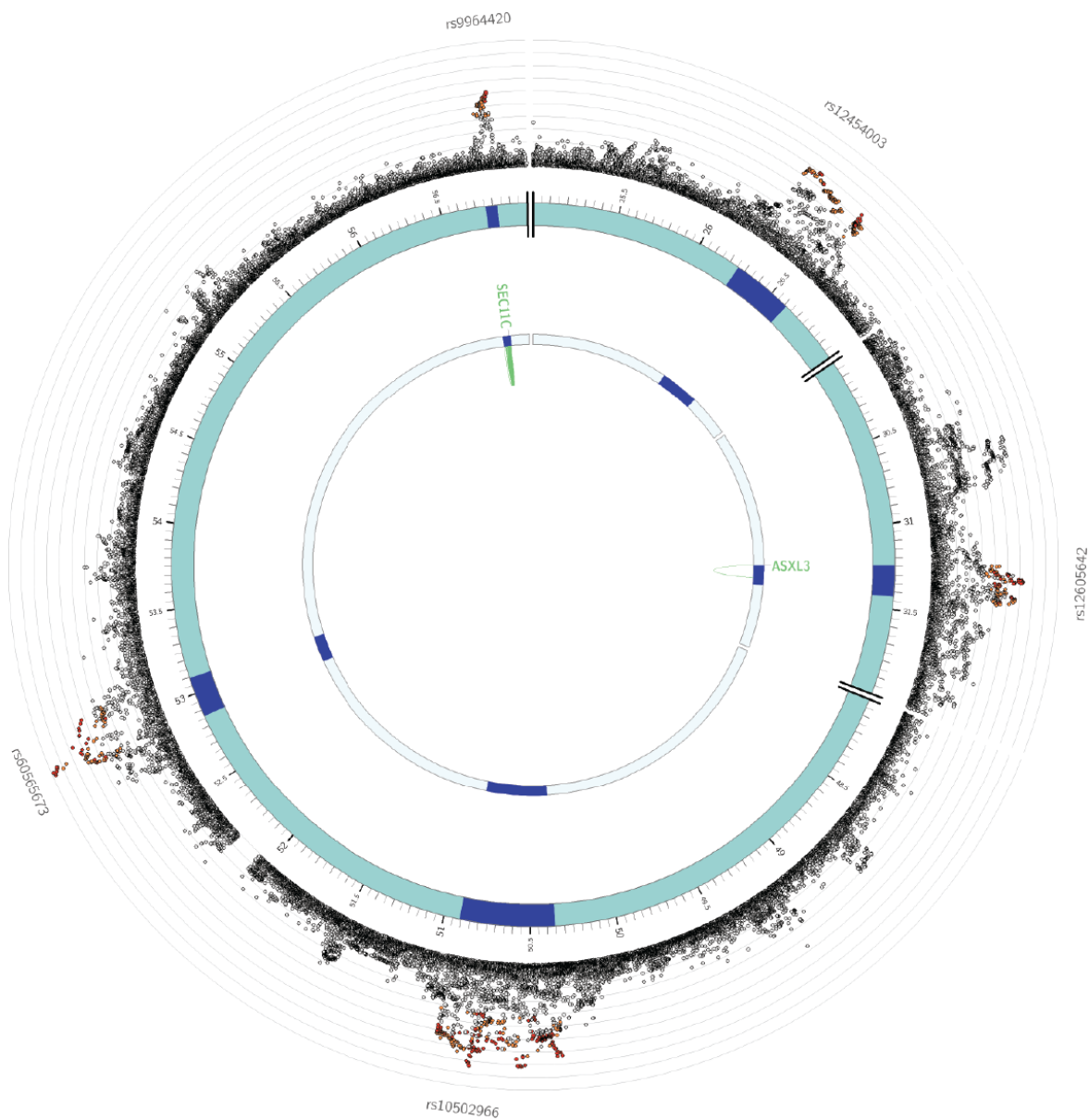
Chromosome 15



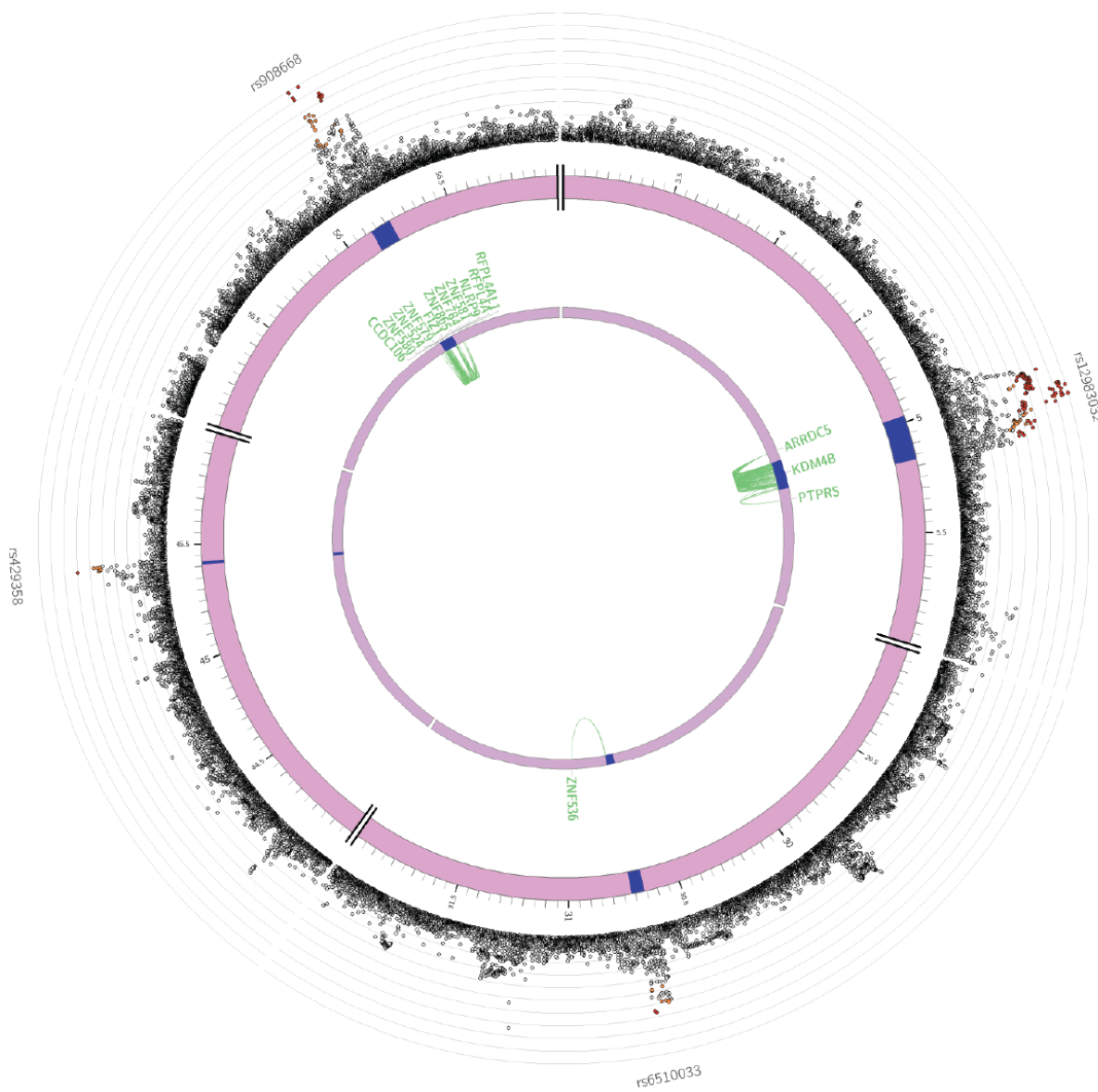
Chromosome 16



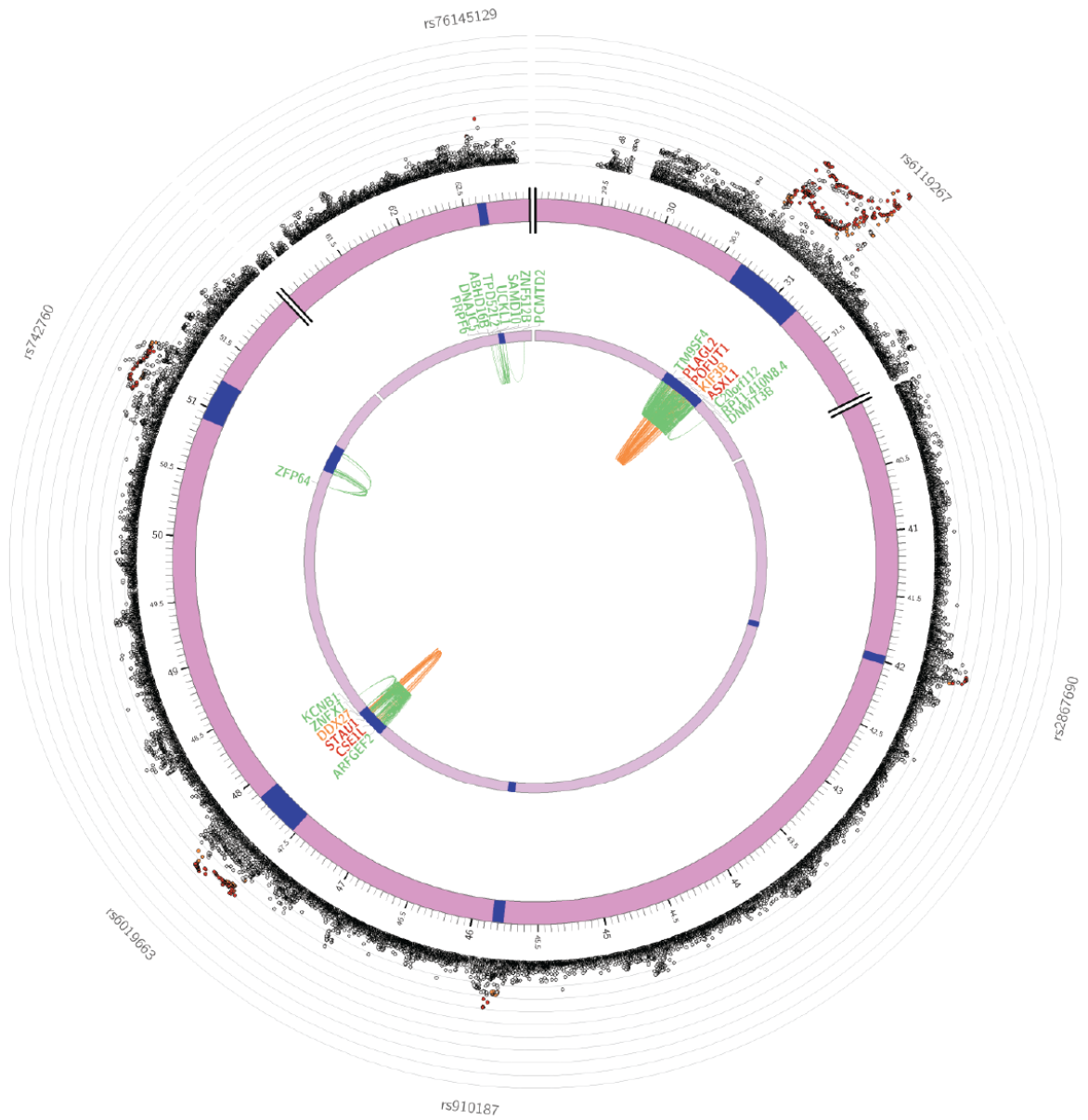
Chromosome 18



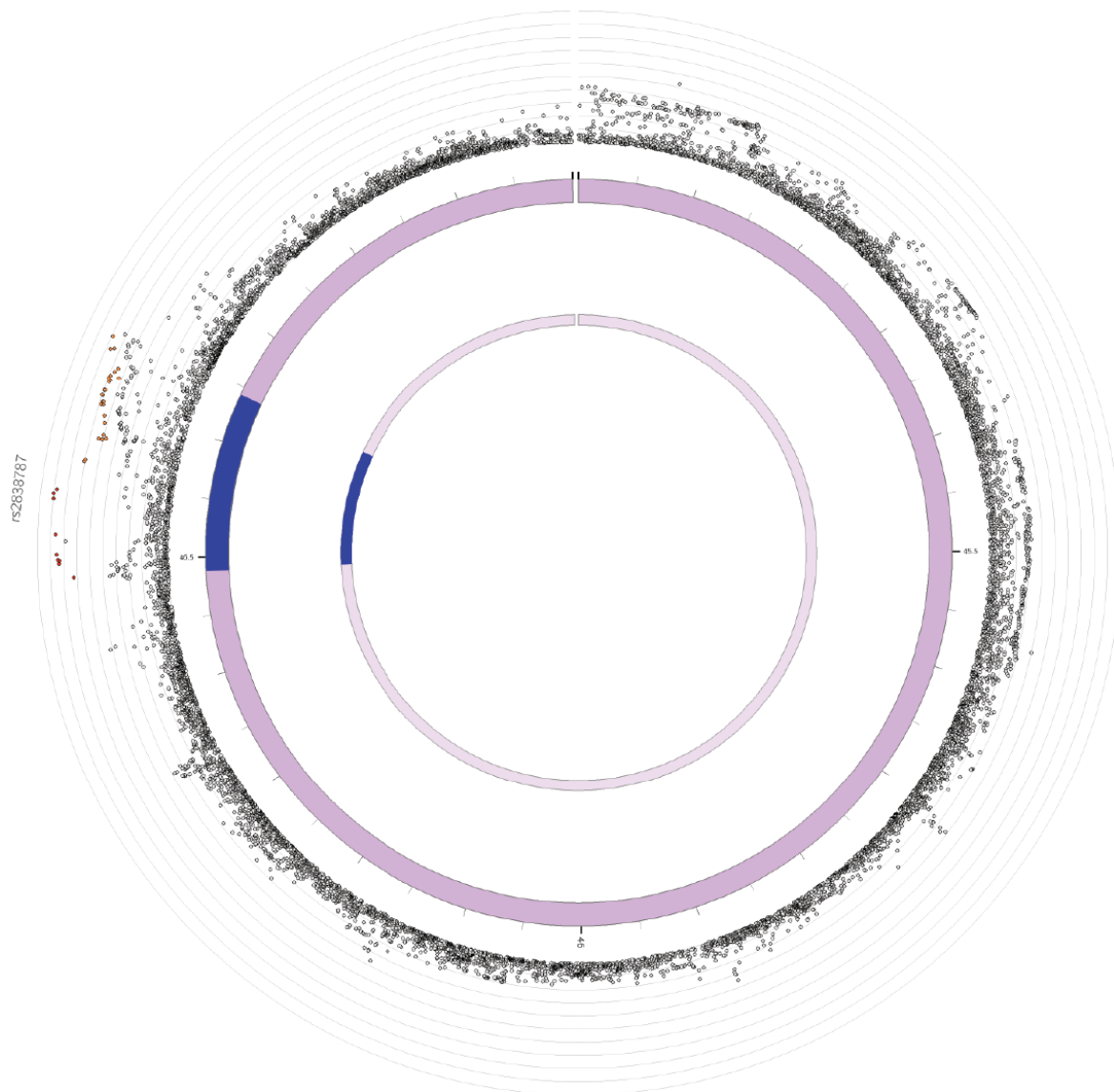
Chromosome 19



Chromosome 20



Chromosome 21



Chromosome 22

