gwascat: structuring and querying the NHGRI GWAS catalog

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1 Introduction

NHGRI maintains and routinely updates a database of selected genome-wide association studies. This document describes R/Bioconductor facilities for working with contents of this database.

1.1 Installation

The package can be installed using Bioconductor's *BiocInstaller* package, with the sequence

```
library(BiocInstaller)
biocLite("gwascat")
```

1.2 Attachment and access to documentation

Once the package has been installed, use library(gwascat) to obtain interactive access to all the facilities. After executing this command, use help(package="gwascat") to obtain an overview. The current version of this vignette can always be accessed at www.bioconductor.org, or by suitably navigating the web pages generated with help.start().

1.3 Illustrations: computing

Available functions are:

```
> library(gwascat)
> objects("package:gwascat")
```

```
[1] "DataFrame"
                           "bindcadd_snv"
                                                 "chklocs"
[4] "elementMetadata"
                                                 "getRsids"
                           "g17SM"
 [7] "getTraits"
                                                 "gw6.rs_17"
                           "gg17N"
                                                 "gwdf_2012_03_20"
[10] "gwastagger"
                           "gwcex2gviz"
[13] "gwdf_2012_09_22"
                           "gwrngs"
                                                 "impute.snps"
                           "locs4trait"
                                                 "low17"
[16] "locon6"
[19] "makeCurrentGwascat" "match"
                                                 "mcols"
[22] "mcols<-"
                           "obo2graphNEL"
                                                 "overlapsAny"
[25] "ranges"
                           "riskyAlleleCount"
                                                 "rules_6.0_1kg_17"
[28] "seqnames"
                           "subsetByChromosome" "subsetByTraits"
[31] "topTraits"
                           "traitsManh"
                                                  "values"
```

The extended GRanges instance with all SNP-disease associations is:

> gwrngs

gwasloc instance with 14719 records and 35 attributes per record.

Extracted: 2013-12-03

Excerpt:

GRanges with 5 ranges and 3 metadata columns:

S	eqnames			ranges	stran	d		
	<rle></rle>			Ranges	<rle< td=""><td>> </td><td></td><td></td></rle<>	>		
[1]	chr10	[96405	5502,	96405502]	:	*		
[2]	chr11	[2024	1544,	2024544]	:	*		
[3]	chr4	[137526	5584,	137526584]	:	*		
[4]	chr22	[19691	1502,	19691502]	:	*		
[5]	chr11	[92784	1203,	92784203]	:	*		
				Disease.Tra	ait		${\tt SNPs}$	p.Value
				<characte< td=""><td>er> <c< td=""><td>nara</td><td>cter></td><td><numeric></numeric></td></c<></td></characte<>	er> <c< td=""><td>nara</td><td>cter></td><td><numeric></numeric></td></c<>	nara	cter>	<numeric></numeric>
[1]		Warfar	in ma	intenance d	ose r	s127	77823	5e-12
[2] [NA meth	ylation	(par	ent-of-orig	in)	rs493	30103	5e-16
[3] [NA meth	ylation	(par	ent-of-orig	in) r	s1001	12307	2e-08
[4] I	NA meth	ylation	(par	ent-of-orig	in)	rs481	19833	6e-08
[5] I	NA meth	ylation	(par	ent-of-orig	in)	rs793	31462	2e-09

seqlengths:

chr1 chr2 chr3 chr4 ... chr21 chr22 chrX 249250621 243199373 198022430 191154276 ... 48129895 51304566 155270560

To determine the most frequently occurring traits:

> topTraits(gwrngs)

Obesity-related traits 951	IgG glycosylation 699	Height 522		
Crohn's disease	Type 2 diabetes	Multiple sclerosis		
210	210	177		
Breast cancer	Ulcerative colitis	Body mass index		
156	147	141		
Coronary heart disease				
140				

For a given trait, obtain a GRanges with all recorded associations; here only three associations are shown:

> subsetByTraits(gwrngs, tr="LDL cholesterol")[1:3]

gwasloc instance with 3 records and 35 attributes per record.

Extracted: 2013-12-03

Excerpt:

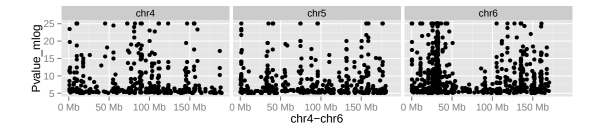
```
GRanges with 3 ranges and 3 metadata columns:
                                                                      SNPs
      seqnames
                              ranges strand |
                                                Disease.Trait
         <Rle>
                           <IRanges>
                                      <Rle> |
                                                   <character> <character>
  [1]
          chr1 [55625548, 55625548]
                                          * | LDL cholesterol rs17111684
          chr2 [21241505, 21241505]
  [2]
                                          * | LDL cholesterol rs12713956
  [3]
          chr2 [44074431, 44074431]
                                          * | LDL cholesterol
                                                                 rs4245791
        p. Value
      <numeric>
  [1]
          2e-17
  [2]
          4e-08
  [3]
          1e-09
  seqlengths:
                             chr3
                                       chr4 ...
                                                     chr21
                                                               chr22
                                                                           chrX
                  chr2
   249250621 243199373 198022430 191154276 ...
                                                 48129895 51304566 155270560
```

2 Some visualizations

2.1 Basic Manhattan plot

A basic Manhattan plot is easily constructed with the ggbio package facilities. Here we confine attention to chromosomes 4:6. First, we create a version of the catalog with $-log_{10}p$ truncated at a maximum value of 25.

```
> gwtrunc = gwrngs
> mlpv = mcols(gwrngs)$Pvalue_mlog
> mlpv = ifelse(mlpv > 25, 25, mlpv)
> mcols(gwtrunc)$Pvalue_mlog = mlpv
> gwlit = gwtrunc[ which(as.character(seqnames(gwtrunc)) %in% c("chr4", "chr5", "chr6
> library(ggbio)
> mlpv = mcols(gwlit)$Pvalue_mlog
> mlpv = ifelse(mlpv > 25, 25, mlpv)
> mcols(gwlit)$Pvalue_mlog = mlpv
> methods:::bind_activation(FALSE)
> autoplot(gwlit, geom="point", aes(y=Pvalue_mlog), xlab="chr4-chr6")
```

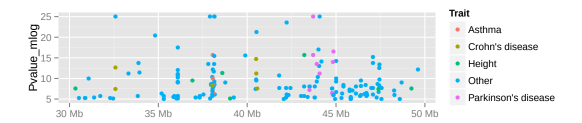


2.2 Annotated Manhattan plot

A simple call permits visualization of GWAS results for a small number of traits. Note the defaults in this call.

> args(traitsManh)

> traitsManh(gwtrunc)



2.3 Integrative view of potential genetic determinants

The following chunk uses GFF3 data on eQTL and related phenomena distributed at the GBrowse instance at eqtl.uchicago.edu. A request for all information at 43-45 Mb was made on 2 June 2012, yielding the GFF3 referenced below. Of interest are locations and scores of genetic associations with DNaseI hypersensitivity (scores identifying dsQTL, see Degner et al 2012).

- $> gffpath = system.file("gff3/chr17_43000000_45000000.gff3", package="gwascat")\\$
- > library(rtracklayer)
- > c17tg = import(gffpath, asRangedData=FALSE)

We make a Gviz DataTrack of the dsQTL scores.

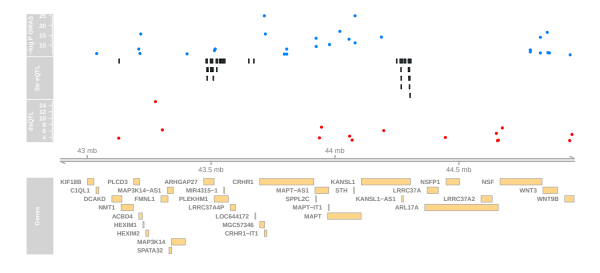
```
> c17td = c17tg[ which(mcols(c17tg)$type == "Degner_dsQTL") ]
> library(Gviz)
> dsqs = DataTrack( c17td, chrom="chr17", genome="hg19", data="score",
+ name="dsQTL")

We start the construction of the graph here.
> g2 = GRanges(seqnames="chr17", IRanges(start=4.3e7, width=2e6))
> basic = gwcex2gviz(contextGR=g2, plot.it=FALSE)

We also collect locations of eQTL in the Stranger 2007 multipopulation eQTL study.
> c17ts = c17tg[ which(mcols(c17tg)$type == "Stranger_eqt1") ]
> eqloc = AnnotationTrack(c17ts, chrom="chr17", genome="hg19", name="Str eQTL")
> displayPars(eqloc)$col = "black"
> displayPars(dsqs)$col = "red"
> integ = list(basic[[1]], eqloc, dsqs, basic[[2]], basic[[3]])

Now use Gviz.
```

> plotTracks(integ)



3 SNP sets and trait sets

3.1 SNPs by name

We can regard the content of a SNP chip as a set of SNP, referenced by name. The pd.genomewidesnp.6 package describes the Affymetrix SNP 6.0 chip. We can determine which traits are associated with loci interrogated by the chip as follows. We work with a subset of the 1 million loci for illustration.

The locon6 data frame has information on 10000 probes, acquired through the following code (not executed here to reduce dependence on the pd.genomewidesnp.6 package, which is very large.

```
> library(pd.genomewidesnp.6)
> con = pd.genomewidesnp.6@getdb()
> locon6 = dbGetQuery(con,
     "select dbsnp_rs_id, chrom, physical_pos from featureSet limit 10000")
Instead use the serialized information:
> data(locon6)
> rson6 = as.character(locon6[[1]])
> rson6[1:5]
[1] "rs2887286" "rs1496555" "rs41477744" "rs3890745" "rs10492936"
   We subset the GWAS ranges structure with rsids that are common to both the chip
and the GWAS catalog. We then tabulate the diseases associated with the common loci.
> intr = gwrngs[ intersect(getRsids(gwrngs), rson6) ]
> sort(table(getTraits(intr)), decreasing=TRUE)[1:10]
                                           Height
                                                3
                         Select biomarker traits
                                   Dental caries
                               IgG glycosylation
```

Immune reponse to smallpox (secreted IFN-alpha)

Metabolic traits

Ulcerative colitis

Age-related macular degeneration

Aging traits

Alcohol dependence

1

3.2 Traits by genomic location

We will assemble genomic coordinates for SNP on the Affymetrix 6.0 chip and show the effects of identifying the trait-associated loci with regions of width 1000bp instead of 1bp.

The following code retrieves coordinates for SNP interrogated on 10000 probes (to save time) on the 6.0 chip, and stores the results in a GRanges instance.

```
> gr6.0 = GRanges(seqnames=ifelse(is.na(locon6$chrom),0,locon6$chrom),
         IRanges(ifelse(is.na(locon6$phys),1,locon6$phys), width=1))
> mcols(gr6.0)$rsid = as.character(locon6$dbsnp_rs_id)
> seqlevels(gr6.0) = paste("chr", seqlevels(gr6.0), sep="")
```

Here we compute overlaps with both the raw disease-associated locus addresses, and with the locus address \pm 500bp.

```
> ag = function(x) as(x, "GRanges")
> ovraw = suppressWarnings(subsetByOverlaps(ag(gwrngs), gr6.0))
> length(ovraw)
[1] 89
> ovaug = suppressWarnings(subsetByOverlaps(ag(gwrngs+500), gr6.0))
> length(ovaug)
[1] 131
```

To acquire the subset of the catalog to which 6.0 probes are within 500bp, use:

```
> rawrs = mcols(ovraw)$SNPs
> augrs = mcols(ovaug)$SNPs
```

> gwrngs[augrs]

gwasloc instance with 131 records and 35 attributes per record.

Extracted: 2013-12-03

Excerpt:

GRanges with 5 ranges and 3 metadata columns:

```
seqnames
                              ranges strand |
       <Rle>
                           <IRanges>
                                       <Rle> |
[1]
       chr10 [ 57549296, 57549296]
                                            * |
[2]
        chr1 [109392837, 109392837]
                                            * |
[3]
       chr10 [124214448, 124214448]
                                            * |
[4]
        chr1 [209712889, 209712889]
                                            * |
[5]
        chr1 [150940625, 150940625]
                                            * |
```

Disease.Trait

```
<character>
  [1]
                    Post-traumatic stress disorder (asjusted for relatedness)
  [2] Adverse response to chemotherapy (neutropenia/leucopenia) (carboplatin)
  [3]
                                              Age-related macular degeneration
  Γ41
                                                 Amyotrophic lateral sclerosis
  [5]
                                             Rhegmatogenous retinal detachment
             SNPs
                    p.Value
      <character> <numeric>
  [1] rs16907840
                      8e-06
  [2] rs1277203
                      9e-06
  [3] rs10490924
                      8e-27
  [4] rs6703183
                      3e-08
  [5]
         rs267738
                      1e-07
  seqlengths:
        chr1
                  chr2
                            chr3
                                       chr4 ...
                                                    chr21
                                                               chr22
                                                                          chrX
   249250621 243199373 198022430 191154276 ...
                                                 48129895 51304566 155270560
  Relaxing the intersection criterion in this limited case leads to a larger set of traits.
> setdiff( getTraits(gwrngs[augrs]), getTraits(gwrngs[rawrs]) )
 [1] "Adverse response to chemotherapy (neutropenia/leucopenia) (carboplatin)"
 [2] "Body mass index"
 [3] "Response to taxane treatment (docetaxel)"
 [4] "Neuroblastoma"
 [5] "Response to amphetamines"
 [6] "Lentiform nucleus volume "
 [7] "Venous thromboembolism"
 [8] "Fasting glucose-related traits (interaction with BMI)"
 [9] "Response to angiotensin II receptor blocker therapy"
[10] "Capecitabine sensitivity"
[11] "Response to hepatitis C treatment"
[12] "Response to antidepressant treatment"
[13] "Phospholipid levels (plasma)"
[14] "Endometrial cancer"
[15] "MRI atrophy measures"
[16] "Self-rated health"
[17] "Neonatal lupus"
[18] "Crohn's disease"
[19] "Optic disc size (cup)"
[20] "Response to statin therapy"
[21] "Tanning"
```

- [22] "Obesity"
- [23] "Osteonecrosis of the jaw"
- [24] "Hip geometry"
- [25] "Parkinson's disease"

4 Counting alleles associated with traits

We can use riskyAlleleCount to count risky alleles enumerated in the GWAS catalog. This particular function assumes that we have genotyped at the catalogued loci. Below we will discuss how to impute from non-catalogued loci to those enumerated in the catalog.

> data(gg17N) # translated from GGdata chr 17 calls using ABmat2nuc > gg17N[1:5,1:5]

	rs6565733	rs1106175	rs17054921	rs8064924	rs8070440
NA06985	"G/G"	"A/G"	"C/C"	"G/G"	"G/G"
NA06991	"G/G"	"A/A"	"C/C"	"G/G"	"G/G"
NA06993	"G/G"	"A/A"	"C/C"	"G/G"	"G/G"
NA06994	"A/G"	"A/G"	"C/C"	"A/G"	"G/G"
NA07000	"G/G"	"A/A"	"C/C"	"G/G"	"G/G"

This function can use genotype information in the A/B format, assuming that B denotes the alphabetically later nucleotide. Because we have direct nucleotide coding in our matrix, we set the matIsAB parameter to false in this call.

```
> h17 = riskyAlleleCount(gg17N, matIsAB=FALSE, chr="ch17")
> h17[1:5,1:5]
```

	rs7217319	rs684232	rs623323	rs2360111	rs11078597
NA06985	0	0	2	0	0
NA06991	0	0	2	1	0
NA06993	0	0	2	2	0
NA06994	0	0	2	2	1
NA07000	0	0	2	0	0

> table(as.numeric(h17))

It is of interest to bind the counts back to the catalog data.

```
> gwr = gwrngs
> gwr = gwr[colnames(h17),]
> mcols(gwr) = cbind(mcols(gwr), DataFrame(t(h17)))
> sn = rownames(h17)
> gwr[,c("Disease.Trait", sn[1:4])]
gwasloc instance with 323 records and 5 attributes per record.
            2013-12-03
Extracted:
Excerpt:
GRanges with 5 ranges and 5 metadata columns:
      segnames
                            ranges strand |
                                                                   Disease.Trait
         <Rle>
                         <IRanges>
                                     <Rle> |
                                                                      <character>
  [1]
         chr17 [ 38924,
                            38924]
                                                                AIDS progression
  [2]
         chr17 [ 618965,
                           618965]
                                                                 Prostate cancer
  [3]
         chr17 [ 700020,
                           700020]
                                                                 Type 2 diabetes
  [4]
         chr17 [ 831667,
                                         * | Economic and political preferences
                           831667]
  [5]
         chr17 [1618363, 1618363]
                                                             Serum albumin level
                                         * |
        NA06985
                   NA06991
                             NA06993
                                        NA06994
      <integer> <integer> <integer> <integer>
  [1]
              0
                         0
                                    0
  [2]
              0
                         0
                                    0
                                              0
                         2
                                              2
  [3]
              2
                                    2
                                    2
                                              2
  [4]
              0
                         1
  [5]
              0
                         0
                                    0
                                              1
  seqlengths:
                                                                chr22
                   chr2
                             chr3
                                        chr4 ...
                                                      chr21
        chr1
                                                                            chrX
   249250621 243199373 198022430 191154276 ...
                                                  48129895
                                                             51304566 155270560
```

Now by programming on the metadata columns, we can identify individuals with particular risk profiles.

5 Imputation to unobserved loci

If we lack information on a specific locus s, but have reasonably dense genotyping on a subject, population genetics may allow a reasonable guess at the genotype at s for this subject. Many algorithms for genotype imputation have been proposed. Here we use a very simple approach due to David Clayton in the snpStats package.

We use the "low coverage" 1000 genomes genotypes for the CEU (central European) HapMap cohort as a base for constructing imputation rules. We focus on chromosome 17 for illustration.

The base data are

```
> data(low17)
> low17

A SnpMatrix with 60 rows and 196327 columns
Row names: NA06985 ... NA12874
Col names: chr17:1869 ... chr17:78654554
```

A somewhat sparser set of genotypes (HapMap phase II, genomewide 4 million loci) on chromosome 17 is archived as g17SM. This has a compact SnpMatrix encoding of genotypes.

```
> data(g17SM)
> g17SM

A SnpMatrix with 90 rows and 89701 columns
Row names: NA06985 ... NA12892
Col names: rs6565733 ... rs4986109
```

For a realistic demonstration, we use the subset of these loci that are present on the Affy 6.0 SNP array.

```
> data(gw6.rs_17)
> g17SM = g17SM[, intersect(colnames(g17SM), gw6.rs_17)]
> dim(g17SM)
```

> if (!exists("rules_6.0_1kg_17")) data(rules_6.0_1kg_17)

[1] 90 20359

The base data were used to create a set of rules allowing imputation from genotypes in the sparse set to the richer set. Some rules involve only a single locus, some as many as 4. The construction of rules involves tuning of modeling parameters. See snp.imputation in snpStats for details.

```
> rules_6.0_1kg_17[1:5,]

chr17:1869 ~ rs9915268+rs11247571+rs9895105+rs6598837 (MAF = 0.06666667, R-squared = chr17:2220 ~ rs4790867+rs10454094+rs2586238+rs7207284 (MAF = 0.125, R-squared = 0.706 chr17:6689 ~ rs4424950+rs4790867+rs7225087+rs11658347 (MAF = 0.125, R-squared = 0.592 rs34663111 ~ rs11658079+rs1609550+rs4985594+rs9788983 (MAF = 0.1166667, R-squared = 0.266 rs62054999 ~ rs17609440+rs2740351+rs2589492+rs16956017 (MAF = 0.125, R-squared = 0.266 rs17609440+rs2740351+rs2589492+rs16956017 (MAF = 0.125609440+rs2586440+rs17609440+rs2586440+rs17609440+rs2586440+rs17609440+rs2586440+rs17609440+rs2586440+rs17609440+rs2586440+rs17
```

The summary of rules shows the degree of association between the predictors and predictands in terms of \mathbb{R}^2 . Many potential targets are not imputed.

```
> summary(rules_6.0_1kg_17)
```

	SNPs us	ed			
R-squared	1 tags	2 tags	3 tags	4 tags	<na></na>
[0,0.1)	655	785	276	56	0
[0.1,0.2)	7	664	926	868	0
[0.2,0.3)	0	158	916	3054	0
[0.3,0.4)	0	28	411	5104	0
[0.4,0.5)	0	20	203	6365	0
[0.5,0.6)	0	21	121	6052	0
[0.6,0.7)	0	29	104	5623	0
[0.7,0.8)	0	54	108	6330	0
[0.8,0.9)	0	141	225	9506	0
[0.9,0.95)	652	700	572	8056	0
[0.95,0.99]	7274	1689	1388	6158	0
[0.99,1]	33660	1353	2326	10152	0
<na></na>	0	0	0	0	53601

The overlap between the 6.0-resident g17SM loci and the catalog is

> length(intersect(colnames(g17SM), mcols(gwrngs)\$SNPs))

[1] 132

The new expected B allele counts are

> exg17 = impute.snps(rules_6.0_1kg_17, g17SM)

The number of new loci that coincide with risk loci in the catalog is:

> length(intersect(colnames(exg17), mcols(gwrngs)\$SNPs))

[1] 197

6 Formal management of trait vocabularies

6.1 Diseases: Disease Ontology

The Disease Ontology project Osborne et al. (2009) formalizes a vocabulary for human diseases. Bioconductor's DO.db package is a curated representation.

- > library(DO.db)
- > DO()

Quality control information for DO:

```
This package has the following mappings:
```

```
DOANCESTOR has 6310 mapped keys (of 6311 keys)
DOCHILDREN has 1796 mapped keys (of 6311 keys)
DOOBSOLETE has 2370 mapped keys (of 2370 keys)
DOOFFSPRING has 1796 mapped keys (of 6311 keys)
DOPARENTS has 6310 mapped keys (of 6311 keys)
DOTERM has 6311 mapped keys (of 6311 keys)
```

Additional Information about this package:

```
DB schema: DO_DB
```

DB schema version: 1.0

All tokens of the ontology are acquired via:

```
> alltob = unlist(mget(mappedkeys(DOTERM), DOTERM))
```

> allt = sapply(alltob, Term)

> allt[1:5]

DOID:0000000 DOID:0000405 DOID:0001816
"gallbladder disease" "vascular tissue disease" "angiosarcoma"
DOID:0002116 DOID:0014667
"pterygium" "disease of metabolism"

Direct mapping from disease trait tokens in the catalog to this vocabulary succeeds for a modest proportion of records.

```
> cattra = mcols(gwrngs)$Disease.Trait
> mat = match(tolower(cattra), tolower(allt))
> catDO = names(allt)[mat]
> na.omit(catDO)[1:50]
```

```
[1] "DOID:3347"
                 "DOID:3347"
                               "DOID:3347"
                                            "DOID:3347"
                                                         "DOID:3347"
[6] "DOID:3347"
                 "DOID:1094"
                               "DOID:1094"
                                            "DOID:1094"
                                                         "DOID:1094"
[11] "DOID:2055"
                               "DOID:12365" "DOID:12365" "DOID:12365"
                 "DOID:2055"
[16] "DOID:12365" "DOID:12365" "DOID:12365" "DOID:12361" "DOID:12361"
[21] "DOID:12361" "DOID:12361" "DOID:12361" "DOID:12361" "DOID:332"
[26] "DOID:332"
                 "DOID:8689"
                               "DOID:8689"
                                            "DOID:8689"
                                                         "DOID:8689"
[31] "DOID:8689"
                 "DOID:8689"
                              "DOID:8689"
                                           "DOID:8689"
                                                        "DOID:8689"
```

```
[36] "DOID:8689" "DOID:8689" "DOID:8689" "DOID:8689" "DOID:8689" [41] "DOID:8689" "DOID:12129" "
```

Approximate matching of unmatched tokens can proceed by various routes. Some traits are not diseases, and will not be mappable using Disease Ontology. However, consider

> unique(cattra[is.na(catDO)])[1:20]

```
[1] "Warfarin maintenance dose"
 [2] "DNA methylation (parent-of-origin)"
 [3] "DNA methylation (variation)"
 [4] "Insomnia"
 [5] "Sleep depth"
 [6] "Sleep duration"
 [7] "Sleep quality"
 [8] "Sleep time"
 [9] "Response to antidepressant treatment (citalogram)"
[10] "Educational attainment"
[11] "LDL cholesterol"
[12] "Triglycerides"
[13] "Blood trace element (Cu levels)"
[14] "Blood trace element (Se levels)"
[15] "Blood trace element (Zn levels)"
[16] "Post-traumatic stress disorder (asjusted for relatedness)"
[17] "Congenital heart disease"
[18] "Congenital heart malformation"
[19] "Adverse response to radiation therapy"
[20] "Primary tooth development (number of teeth)"
> nomatch = cattra[is.na(catD0)]
> unique(nomatch)[1:5]
[1] "Warfarin maintenance dose"
                                          "DNA methylation (parent-of-origin)"
                                          "Insomnia"
[3] "DNA methylation (variation)"
[5] "Sleep depth"
```

Manual searching shows that a number of these have very close matches.

6.2 Other phenotypic traits: Human Phenotype Ontology

Bioconductor does not possess an annotation package for phenotype ontology, but the standardized OBO format can be parsed and modeled into a graph.

```
> hpobo = gzfile(dir(system.file("obo", package="gwascat"), pattern="hpo", full=TRUE)
> HPOgraph = obo2graphNEL(hpobo)
> close(hpobo)
  The phenotypic terms are obtained via:
> hpoterms = unlist(nodeData(HPOgraph, nodes(HPOgraph), "name"))
> hpoterms[1:10]
                                HP:000001
                                      " A ] ] "
                                HP:000002
              "Abnormality of body height"
                                HP:000003
            "Multicystic kidney dysplasia"
                                HP:000004
               "Onset and clinical course"
                                HP:000005
                     "Mode of inheritance"
                                HP:000006
          "Autosomal dominant inheritance"
                                HP:000007
         "Autosomal recessive inheritance"
                                HP:000008
"Abnormality of female internal genitalia"
                                HP:000009
   "Functional abnormality of the bladder"
                                HP:000010
      "Recurrent urinary tract infections"
  Exact hits to unmatched GWAS catalog traits exist:
> intersect(tolower(nomatch), tolower(hpoterms))
 [1] "insomnia"
                                      "scoliosis"
 [3] "eating disorders"
                                      "hypertriglyceridemia"
 [5] "coronary artery calcification" "sagittal craniosynostosis"
 [7] "glioma"
                                      "autism"
 [9] "atrial fibrillation"
                                      "stroke"
[11] "iga nephropathy"
                                      "nephropathy"
[13] "glomerulosclerosis"
                                      "freckling"
[15] "knee osteoarthritis"
                                      "hearing impairment"
```

More work on formalization of trait terms is underway.

7 CADD scores

[18] -0.695270

Kircher et al. (Kircher et al., 2014) define combined annotation-dependent depletion scores measuring variant pathogenicity in an integrative way. Small requests to bind scores for SNV to GRanges can be resolved through HTTP; large requests can be carried out on a local tabix-indexed selection from their archive.

```
> g3 = as(gwrngs, "GRanges")
> bg3 = bindcadd_snv( g3[which(seqnames(g3)=="chr3")][1:20] )
> inds = ncol(mcols(bg3))
> bg3[, (inds-3):inds]
```

This requires cooperation of network interface and server, so we don't evaluate in vignette build but on 1 Apr 2014 the response was:

GRanges with 20 ranges and 4 metadata columns:

0.987

O		O					
	seqnames		ranges	strand		Ref	Alt
	<rle></rle>		Ranges	<rle></rle>		<character></character>	<character></character>
[1]	3	[109789570,	109789570]	*		A	G
[2]	3	[25922285,	25922285]	*		G	Α
[3]	3	[109529550,	109529550]	*	- 1	Т	C
[4]	3	[175055759,	175055759]	*	- 1	Т	G
[5]	3	[191912870,	191912870]	*	- 1	C	T
[16]	3	[187716886,	187716886]	*		A	G
[17]	3	[160820524,	160820524]	*		G	C
[18]	3	[169518455,	169518455]	*		T	C
[19]	3	[179172979,	179172979]	*		G	T
[20]	3	[171785168,	171785168]	*		G	C
	CScore	PHRED					
	<numeric></numeric>	<pre> <numeric></numeric></pre>					
[1]	-0.182763	3.110					
[2]	-0.289708	2.616					
[3]	0.225373	5.216					
[4]	-0.205689	3.003					
[5]	-0.172189	3.161					
[16]	-0.019710	3.913					
[17]	-0.375183	2.235					

8 Appendix: Adequacy of location annotation

A basic question concerning the use of archived SNP identifiers is durability of the association between asserted location and SNP identifier. The chklocs function uses a current Bioconductor SNPlocs package to check this.

For example, to verify that locations asserted on chromosome 20 agree between the Bioconductor dbSNP image and the gwas catalog,

```
> if ("SNPlocs.Hsapiens.dbSNP.20120608" %in% installed.packages()[,1]) {
+ library(SNPlocs.Hsapiens.dbSNP.20120608)
+ suppressWarnings(chklocs("20"))
+ }
[1] TRUE
```

This is not a fast procedure but has succeeded for all chromosomes 1-22 when checked off line.

References

Martin Kircher, Daniela M Witten, Preti Jain, Brian J O'Roak, Gregory M Cooper, and Jay Shendure. A general framework for estimating the relative pathogenicity of human genetic variants. *Nat Genet*, Feb 2014. doi: 10.1038/ng.2892.

John D Osborne, Jared Flatow, Michelle Holko, Simon M Lin, Warren A Kibbe, Lihua Julie Zhu, Maria I Danila, Gang Feng, and Rex L Chisholm. Annotating the human genome with disease ontology. *BMC Genomics*, 10 Suppl 1:S6, Jan 2009. doi: 10.1186/1471-2164-10-S1-S6. URL http://www.biomedcentral.com/1471-2164/10/S1/S6.