

AMP-DCC Quality Control Report

FUSION

05/21/2019 (08:56)

Prepared by Ryan Koesterer on behalf of the AMP-DCC Data Analysis Team at Broad Institute

Contact: amp-dcc-dat@broadinstitute.org

This document was generated using Loamstream [10] and the AMP-DCC Data Analysis Pipeline [11]

Contents

1	Introduction	2
2	Data	3
2.1	Samples	3
2.2	Variants	3
3	Sample QC	8
3.1	Ancestry Inference	8
3.2	Duplicates and Excessive Sharing of Identity-by-Descent (IBD)	13
3.3	Sex Chromosome Check	14
3.4	Sample Outlier Detection	14
3.4.1	Principal Component Adjustment and Normalization of Sample Metrics	15
3.4.2	Individual Sample Metric Clustering	15
3.4.3	Principal Components of Variation in PCARM's	16
3.4.4	Combined PCARM Clustering	16
3.4.5	Plots of Sample Outliers	16
3.5	Summary of Sample Outlier Detection	21
4	Variant QC	23
5	Acknowledgements	24

6 References

25

1 Introduction

This document contains details of our in-house quality control procedure and its application to the FUSION dataset. We received genotypes for 4,233 unique samples distributed across 4 different genotype arrays. Quality control was performed on these data to detect samples and variants that did not fit our standards for inclusion in association testing. After harmonizing with modern reference data, the highest quality variants were used in a battery of tests to assess the quality of each sample. Duplicate pairs, samples exhibiting excessive sharing of identity by descent, samples whose genotypic sex did not match their clinical sex, and outliers detected among several sample-by-variant statistics have been flagged for removal from further analysis. Additionally, genotypic ancestry was inferred with respect to a modern reference panel, allowing for variant filtering and association analyses to be performed within population as needed. With the exception of inferring each samples ancestry, QC was performed on each array separately as much as possible, allowing for flexibility in the way the data can be used in downstream analyses.

2 Data

2.1 Samples

Initially, the array was checked for sample genotype missingness. Any samples with extreme genotype missingness (> 0.5) were removed prior to our standard quality control procedures. There were 4 samples removed from this data set.

The following diagram (Figure 1) describes the remaining sample distribution over the 4 genotyping arrays, along with their intersection sizes.

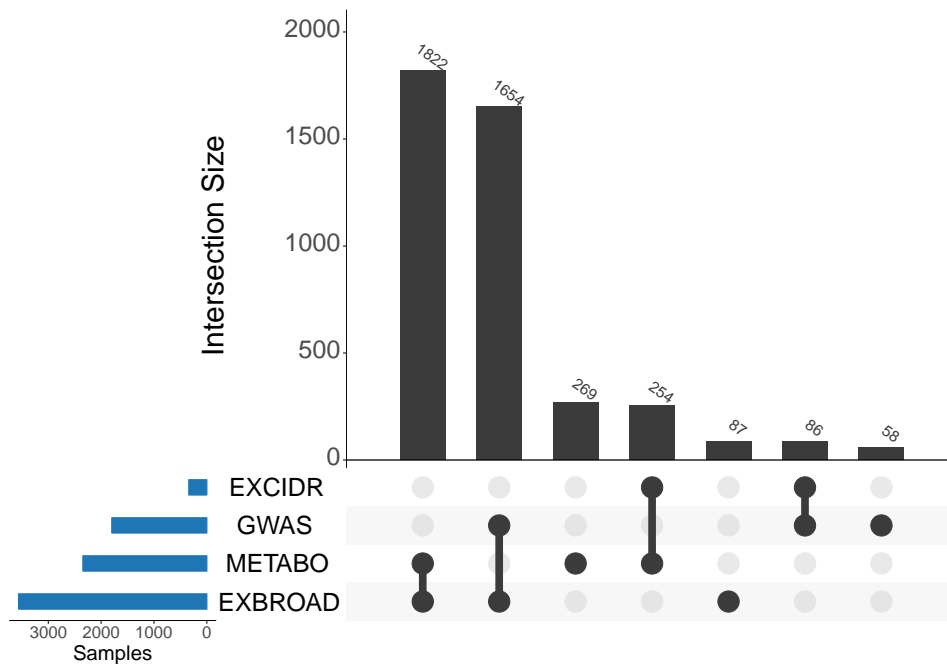


Figure 1: Samples distributed by genotyping array

2.2 Variants

Table 1 gives an overview of the different variant classes and how they distributed across allele frequencies for each dataset. Note that the totals reflect the sum of the chromosomes only. A legend has been provided below the table for further inspection of the class definitions.

Table 1: Summary of raw variants by frequency and classification

Freq = Minor allele frequency (MAF) range

Unpl = Chromosome = 0

Auto = Autosomal variants

X = X chromosome non-pseudoautosomal region (non-PAR) variants

Y = Y chromosome variants

X(PAR) = X chromosome pseudoautosomal (PAR) region variants

Mito = Mitochondrial variants

InDel = Insertion/Deletion variants (I/D or D/I alleles)

Multi = Multiallelic variants (2 or more alternate alleles)

Dup = Duplicated variants with respect to position and alleles

	Freq	Unpl	Auto	X	Y	X(PAR)	Mito	InDel	Multi	Dup	Total
GWAS	[0]	0	0	0	0	0	0	0	0	0	0
	(0,0.01)	0	406	24	0	0	0	0	0	0	430
	[0.01,0.03)	0	3915	160	0	0	0	0	0	0	4075
	[0.03,0.05)	0	8802	282	0	0	0	0	0	0	9084
	[0.05,0.10)	0	32094	864	0	0	0	0	0	0	32958
	[0.10,0.50]	0	261471	7675	0	0	0	0	0	0	269146
	Total	0	306688	9005	0	0	0	0	0	0	315693
EXBROAD	[0]	0	147217	3349	0	0	0	0	0	0	150566
	(0,0.01)	0	48452	972	0	0	0	0	0	0	49424
	[0.01,0.03)	0	7051	160	0	0	0	0	0	0	7211
	[0.03,0.05)	0	2817	57	0	0	0	0	0	0	2874
	[0.05,0.10)	0	4100	79	0	0	0	0	0	0	4179
	[0.10,0.50]	0	21365	438	0	0	0	0	0	0	21803
	Total	0	231002	5055	0	0	0	0	0	0	236057
EXCIDR	[0]	0	197189	4111	78	80	133	0	0	0	201591
	(0,0.01)	0	0	0	0	0	0	0	0	0	0
	[0.01,0.03)	0	10732	143	0	4	17	0	0	0	10896
	[0.03,0.05)	0	2101	91	0	1	0	0	0	0	2193
	[0.05,0.10)	0	4702	69	14	3	3	0	0	0	4791
	[0.10,0.50]	0	22543	411	2	20	11	0	0	0	22987
	Total	0	237267	4825	94	108	164	0	0	0	242458
METABO	[0]	0	51711	0	8	0	44	0	0	3	51763
	(0,0.01)	0	12307	0	6	0	27	0	0	0	12340
	[0.01,0.03)	0	11210	0	0	0	15	0	0	2	11225
	[0.03,0.05)	0	9190	0	4	0	8	0	0	1	9202
	[0.05,0.10)	0	17993	1	3	0	7	0	0	1	18004
	[0.10,0.50]	0	86810	91	0	1	7	0	2	7	86909
	Total	0	189221	92	21	1	108	0	2	14	189443

To facilitate downstream operations on genotype data, such as merging and meta-analysis, each dataset

gets harmonized with modern reference data. The harmonization process is performed in two steps. First, using Genotype Harmonizer [2], the variants are strand-aligned with the 1000 Genomes Phase 3 Version 5 [4] variants. While some variants (A/C or G/T variants) may be removed due to strand ambiguity, if enough information exists, Genotype Harmonizer uses linkage disequilibrium (LD) patterns with nearby variants to accurately determine strand. This step will remove variants that it is unable to reconcile and maintains variants that are unique to the input data. The second step manually reconciles non-1000 Genomes variants with the human reference assembly GRCh37 [7]. This step will flag variants for removal that do not match an allele to the reference and variants that have only a single allele in the data file (0 for the other). Note that some monomorphic variants may be maintained in this process if there are two alleles in the data file and one of them matches a reference allele.

After harmonization, the data is loaded into a Hail [9] matrix table for downstream use. See Figure 2 for final variant counts by genotyping array.

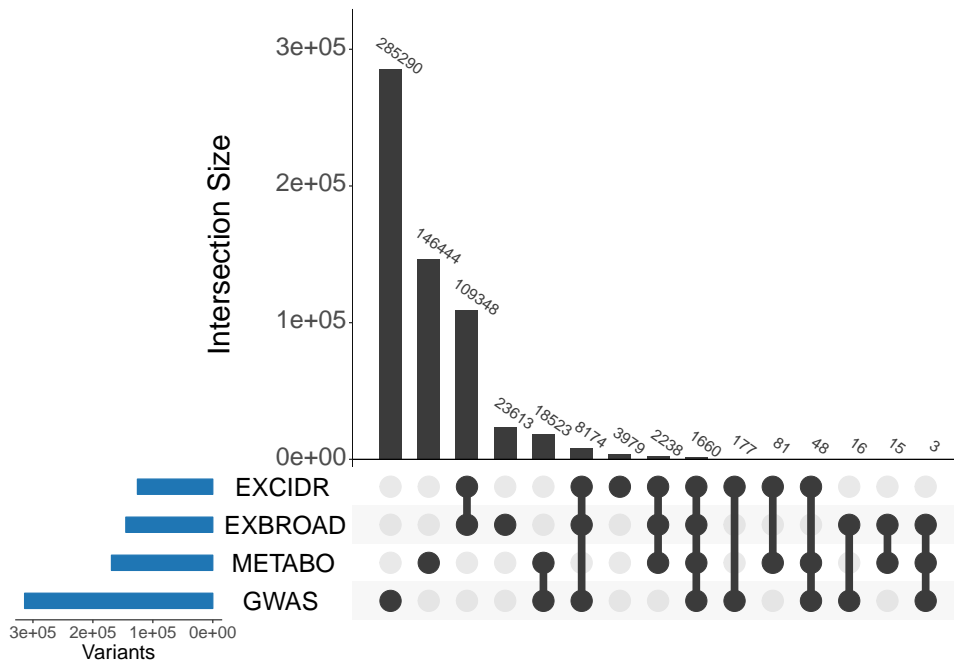


Figure 2: Variants remaining for analysis

3 Sample QC

3.1 Ancestry Inference

Prior to association testing, it is useful to infer ancestry in relation to a modern reference panel representing the major human populations. While our particular sample QC process does not directly depend on this information, it is useful to downstream analysis when stratifying the calculation of certain variant statistics that are sensitive to population substructure (eg. Hardy Weinberg equilibrium). Additionally, ancestry inference may identify samples that do not seem to fit into a well-defined major population group, which would allow them to be flagged for removal from association testing.

Initially, each array was merged with reference data. In this case, the reference used was the entire set of 2,504 1000 Genomes Phase 3 Version 5 [4] samples and our method restricted this merging to a set of 5,166 known ancestry informative SNPs. The merged data consisted of 2,328 GWAS, 2,249 EXBROAD, 2,314 EXCIDR and 414 METABO variants. After merging, principal components (PCs) were computed using the PC-AiR [3] method in the GENESIS R package. This particular algorithm allows for the calculation of PCs that reflect ancestry in the presence of known or cryptic relatedness. The 1000 Genomes samples were forced into the 'unrelated' set and the PC-AiR algorithm was used to find the 'unrelated' samples from the array data. Then PCs were calculated on them and projected onto the remaining samples.

Figures 3, 4, 5, and 6 display plots of the top three principal components along with the 1000 Genomes major population groups.

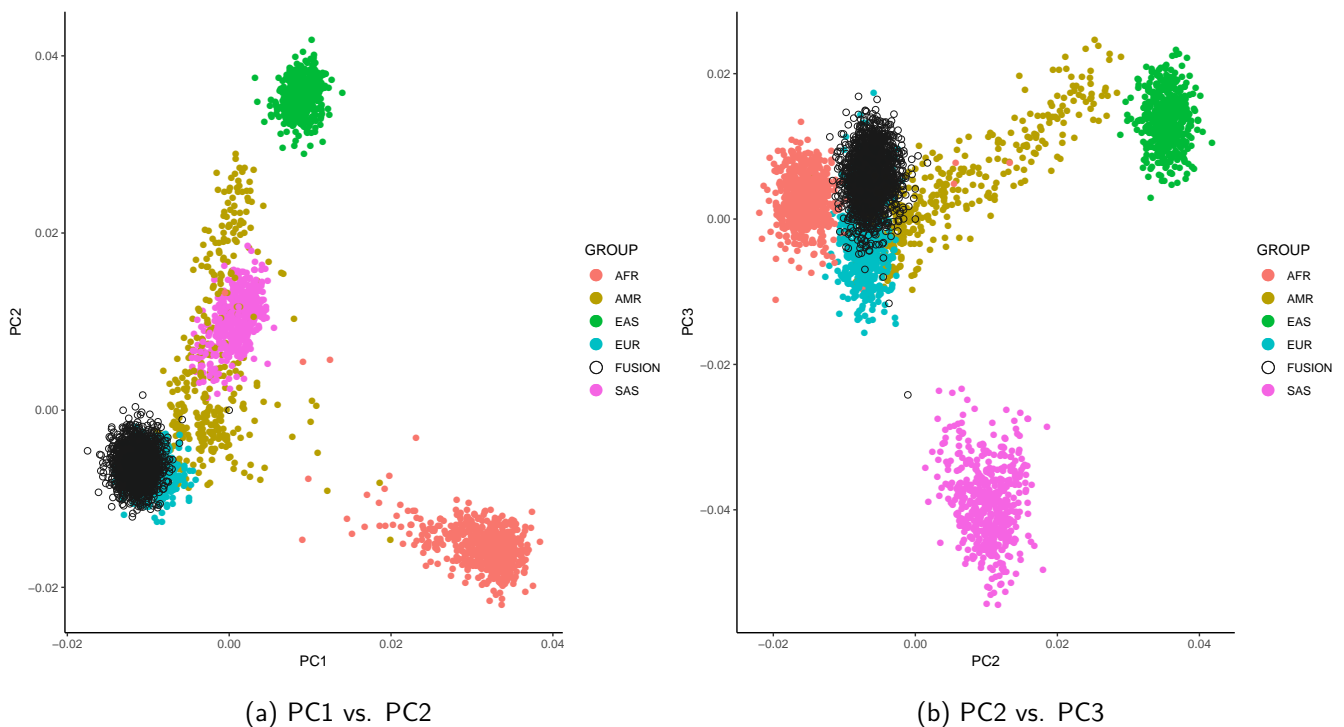
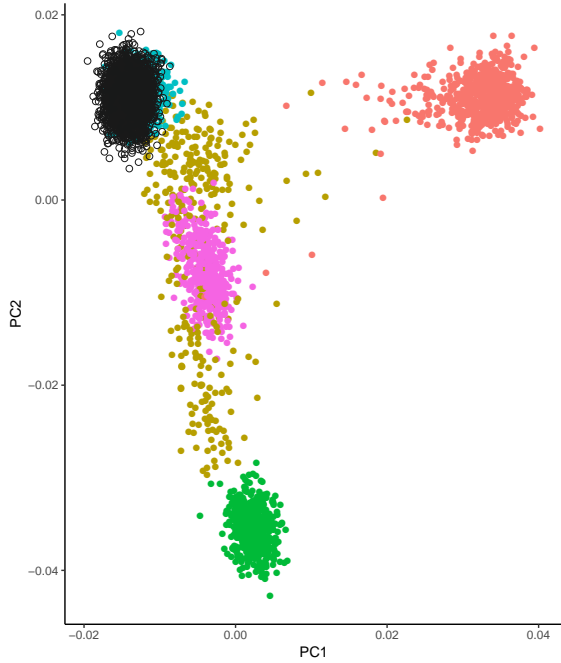
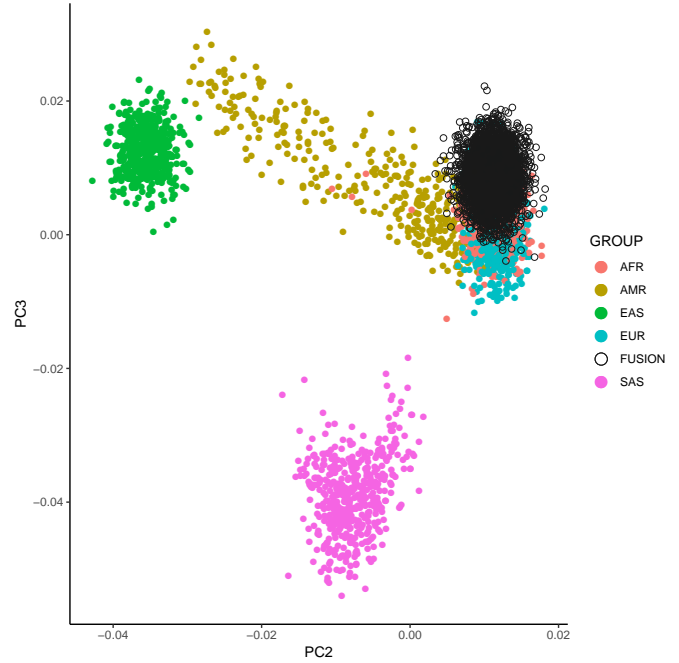


Figure 3: Principal components of ancestry for GWAS

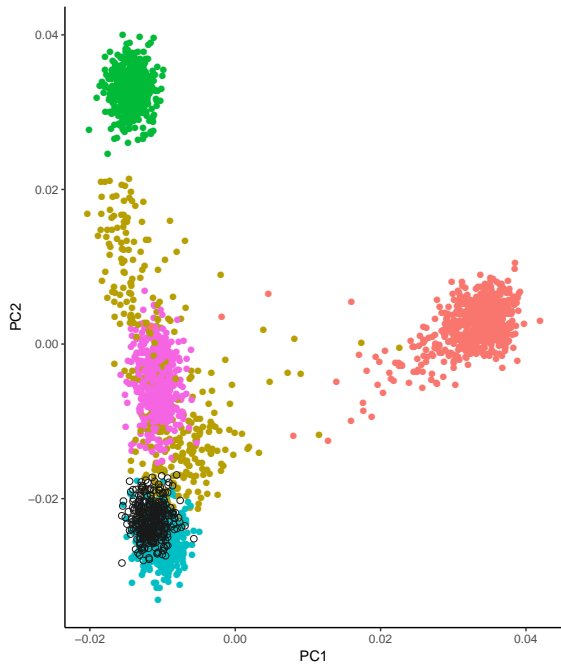


(a) PC1 vs. PC2

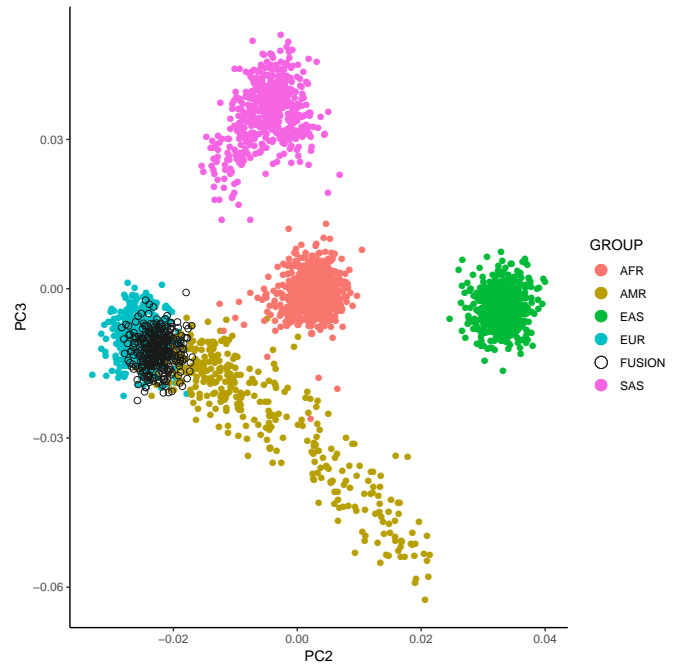


(b) PC2 vs. PC3

Figure 4: Principal components of ancestry for EXBROAD



(a) PC1 vs. PC2



(b) PC2 vs. PC3

Figure 5: Principal components of ancestry for EXCIDR

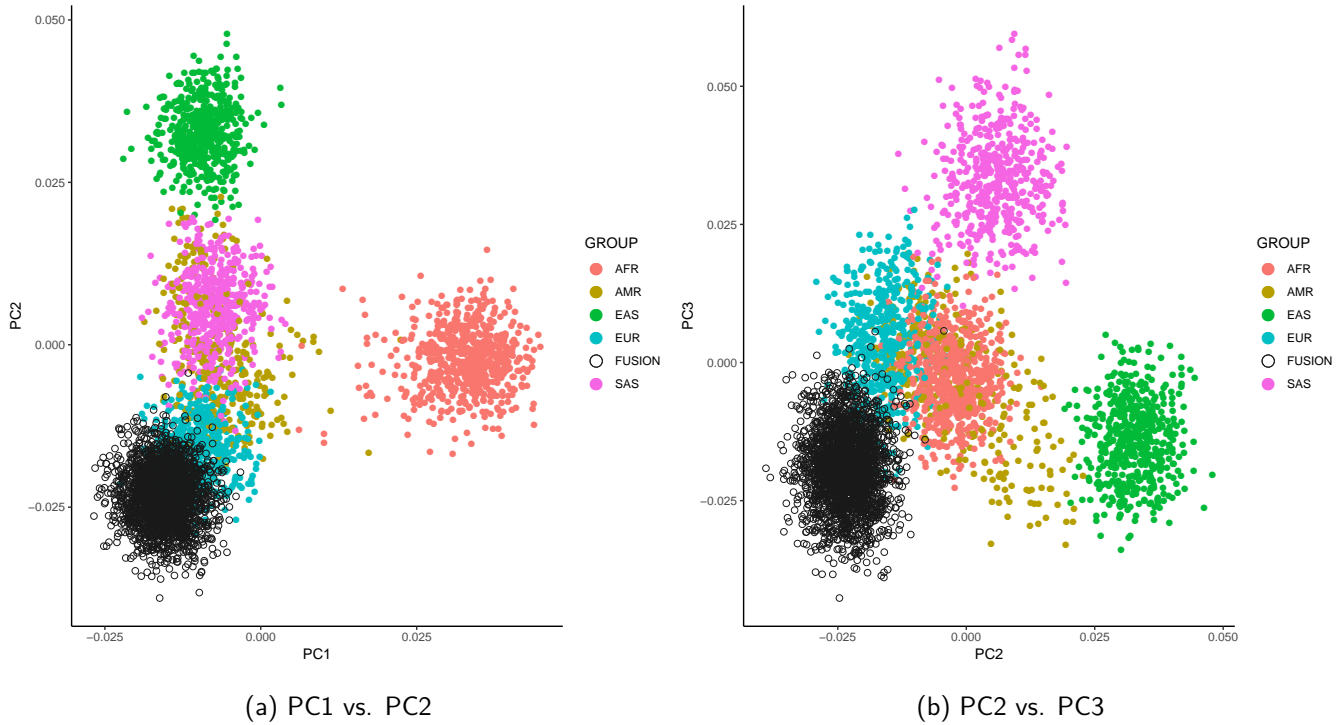


Figure 6: Principal components of ancestry for METABO

Using the principal components of ancestry as features, we employed the signal processing software Klustakwik [5] to model the array as a mixture of Gaussians, identifying clusters, or population groups/subgroups. In order to generate clusters of sufficient size for statistical association tests, we used the first 5 principal components as features in the clustering algorithm. This number of PC's distinctly separates the five major 1000 Genomes population groups: AFR, AMR, EUR, EAS, and SAS. Figures 7, 8, 9, and 10 clearly indicate the population structure in the datasets. In Klustakwik output, cluster 1 is always reserved for outliers, or samples that did not fit into any of the clusters found by the program. Upon further inspection, no samples were manually reinstated during this step. More information is available upon request

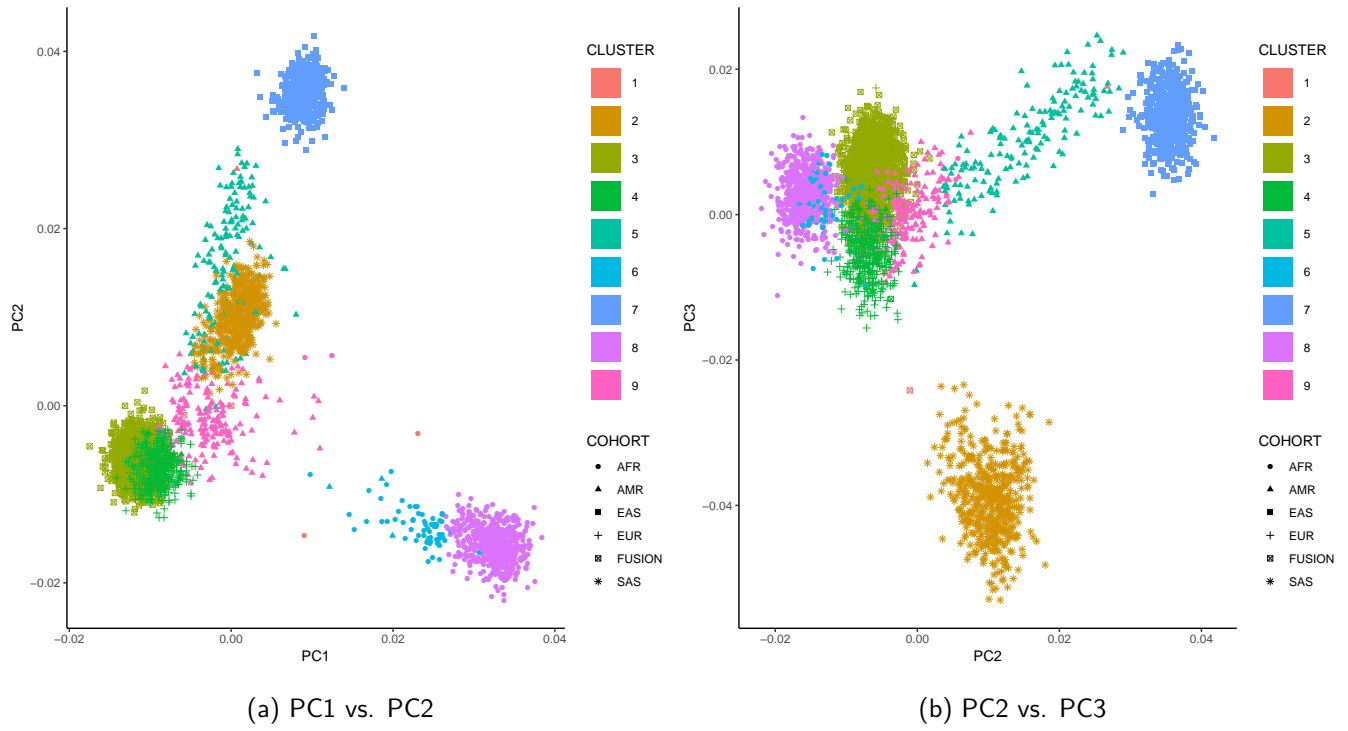


Figure 7: Population clusters for GWAS

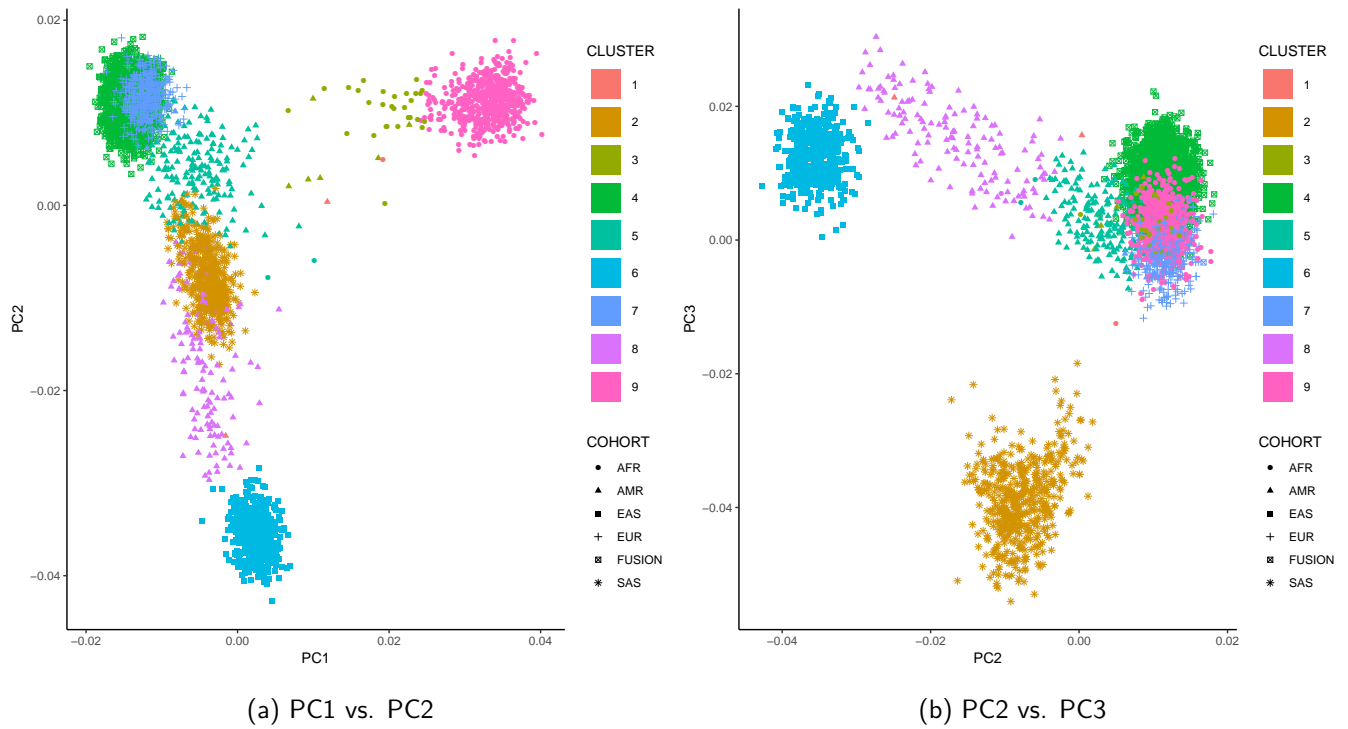


Figure 8: Population clusters for EXBROAD

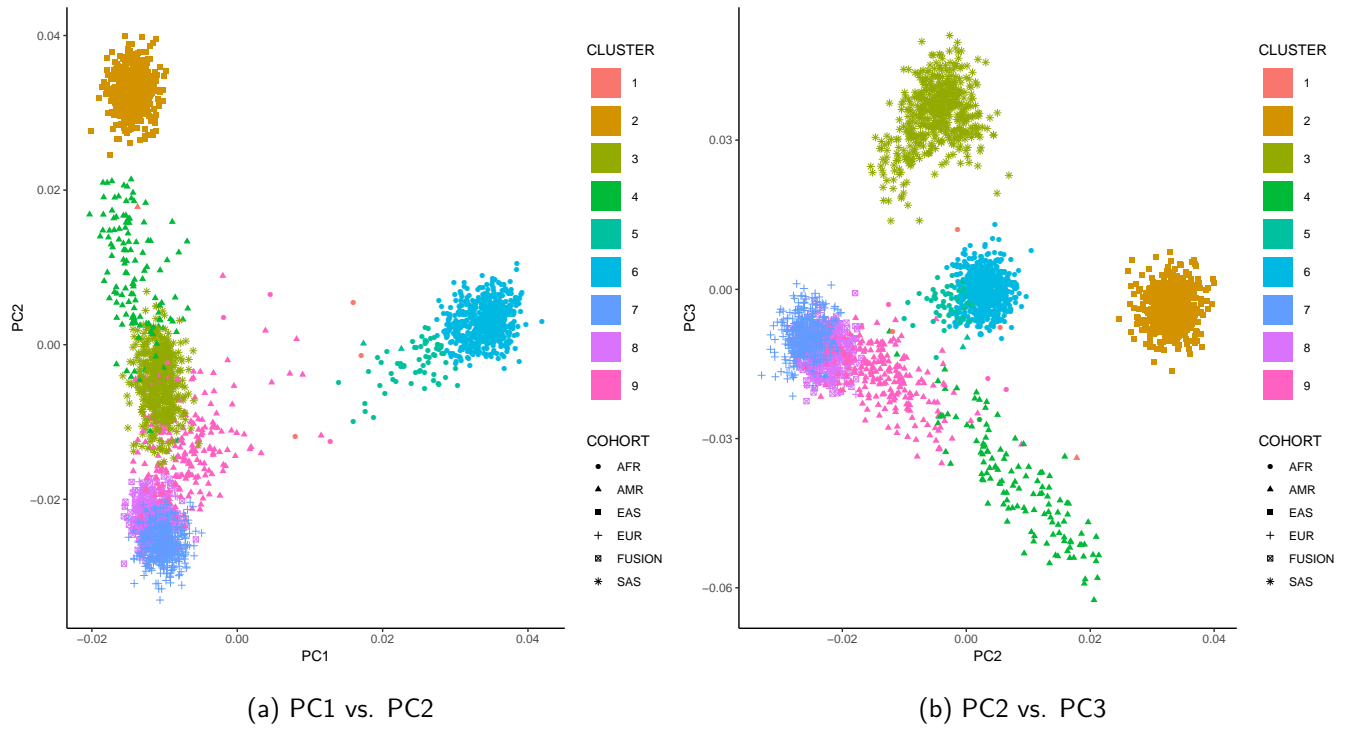


Figure 9: Population clusters for EXCIDR

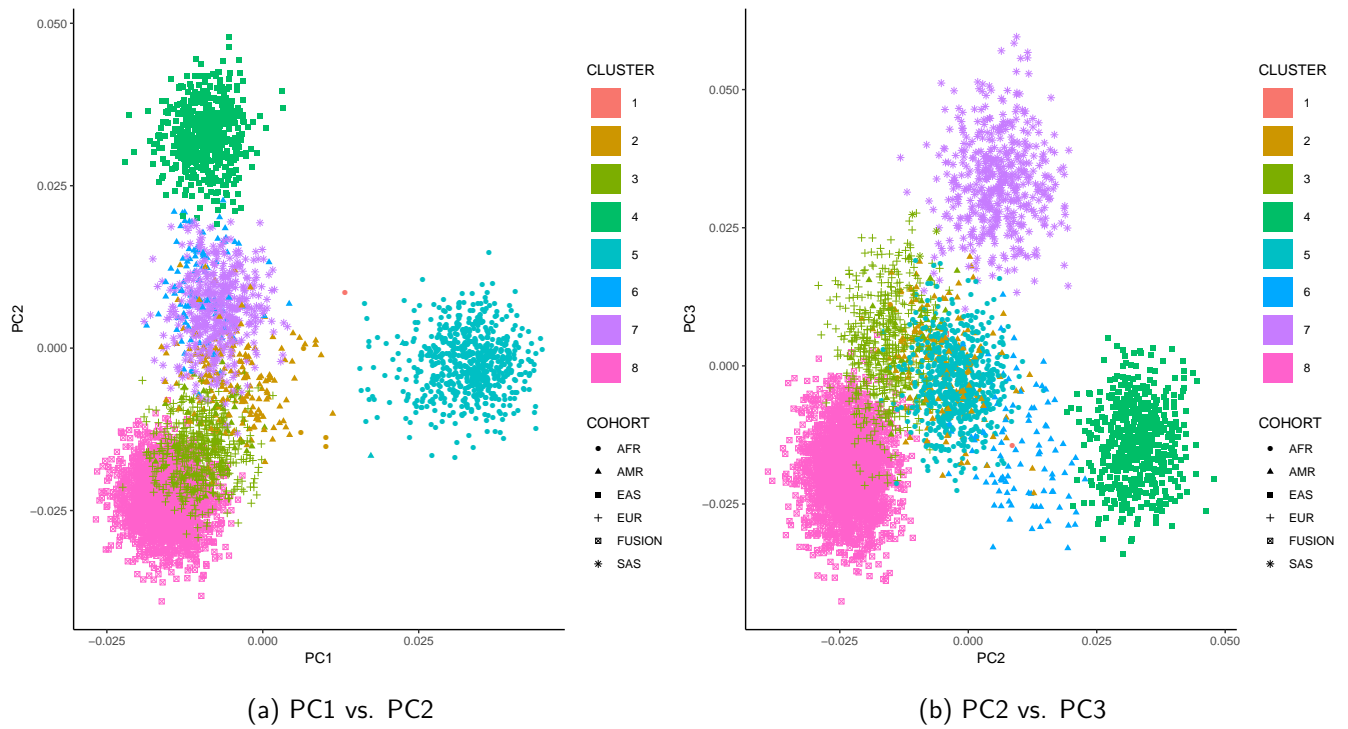


Figure 10: Population clusters for METABO

The resulting clusters are then combined with the nearest 1000 Genomes cohort. Table 2 describes the classification using this method. A final population assignment is determined by setting a hierarchy on the genotyping technologies (GWAS > EXBROAD > EXCIDR > METABO) and assigning each sample to the population determined using the highest technology.

Table 2: Inferred ancestry by dataset and cluster

	Population	Clusters	Samples
GWAS	EUR	3,4	1796
	Outliers	1	2
EXBROAD	EUR	4,7	3563
	Outliers	1	0
EXCIDR	EUR	8	340
	Outliers	1	0
METABO	EUR	8	2344
	Outliers	1	1

Table 3: Final inferred ancestry

Population	Samples
EUR	4227
Outliers	3

3.2 Duplicates and Excessive Sharing of Identity-by-Descent (IBD)

Sample pair kinship coefficients were determined using KING [8] relationship inference software, which offers a robust algorithm for relationship inference under population stratification. Prior to inferring relationships, we used Plink [1] to filter out non-autosomal, non-A/C/G/T, low callrate, and low minor allele frequency variants. Also, variants with positions in known high LD regions [6] and known Type 2 diabetes associated loci were

removed and an LD-pruned dataset was created. The specific filters that were used are listed below.

- --chr 1-22
- --snps-only just-acgt
- --exclude range ...
- --maf 0.01
- --geno 0.02
- --indep-pairwise 1000kb 1 0.2

After filtering there were 79,245 GWAS, 21,790 EXBROAD, 22,376 EXCIDR and 38,585 METABO variants remaining.

In order to identify duplicate pairs of samples, a filter was set to $Kinship > 0.4$. There were no sample pairs identified as duplicate in the array data. Upon manual inspection, if the clinical data for any of the duplicate pairs was nearly identical (same date of birth, etc.), then the sample with the higher call rate was reinstated. If the clinical data did not match or a manual inspection was not performed, both samples were removed. In this case, no samples have been reinstated. More information is available upon request

In addition to identifying duplicate samples, any single individual that exhibited kinship values indicating a 2nd degree relative or higher relationship with 10 or more others was flagged for removal. The relationship count indicated no samples that exhibited high levels of sharing identity by descent. Upon further inspection, no samples were manually reinstated during this step. More information is available upon request

3.3 Sex Chromosome Check

Each array was checked for genotype / clinical data agreement for sex. There was 1 sample that was flagged as a 'PROBLEM' by Hail because it was unable to impute sex and there were no samples that were flagged for removal because the genotype based sex did not match their clinical sex. Upon further inspection, no samples were manually reinstated during this step. More information is available upon request

3.4 Sample Outlier Detection

Each sample was evaluated for inclusion in association tests based on 10 sample-by-variant metrics (Table 4), calculated using Hail [9]. Note that for the metrics `n_called` and `call_rate`, only samples below the mean are filtered.

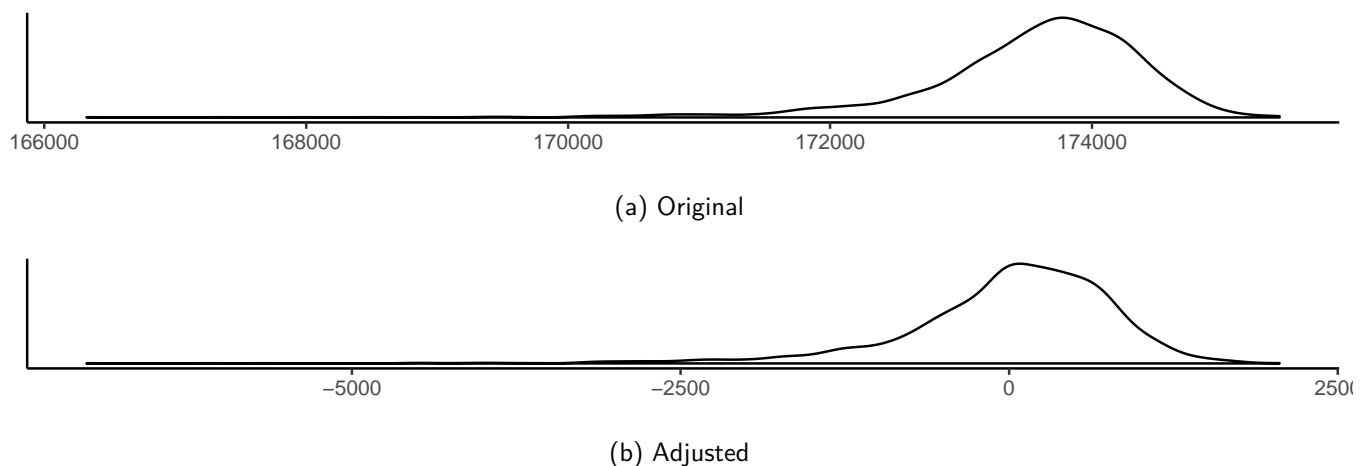
Table 4: Sample Metrics

n_non_ref	n_het + n_hom_var
n_het	Number of heterozygous variants
n_called	n_hom_ref + n_het + n_hom_var
call_rate	Fraction of variants with called genotypes
r_ti_tv	Transition/transversion ratio
het	Inbreeding coefficient
het_high	Inbreeding coefficient for variants with $MAF \geq 0.03$
het_low	Inbreeding coefficient for variants with $MAF < 0.03$
n_hom_var	Number of homozygous alternate variants
r_het_hom_var	het/hom_var ratio across all variants

3.4.1 Principal Component Adjustment and Normalization of Sample Metrics

Due to possible population substructure, the sample metrics exhibit some multi-modality in their distributions. To evaluate more normally distributed data, we calculated principal component adjusted residuals of the metrics using the top 10 principal components (PCARM's). Figure 11 shows the n_non_ref metric for GWAS samples before and after adjustment.

Figure 11: Comparison of n_non_ref distributions before and after adjustment / normalization



3.4.2 Individual Sample Metric Clustering

For outlier detection, we clustered the samples into Gaussian distributed subsets with respect to each PCARM using the software Klustakwik [5]. During this process, samples that did not fit into any Gaussian distributed set

of samples were identified and flagged for removal.

3.4.3 Principal Components of Variation in PCARM's

In addition to outliers along individual sample metrics, there may be samples that exhibit deviation from the norm across multiple metrics. In order to identify these samples, we calculated principal components explaining 95% of the variation in 8 of the 10 PCARMs combined. The adjusted residuals for metrics 'call_rate' and 'n_called' are characterized by long tails that lead to the maximum value, which is not consistent with the other metrics. In order to avoid excessive flagging of samples with lower, yet still completely acceptable, call rates, these metrics were left out of principal component calculation.

3.4.4 Combined PCARM Clustering

All samples were clustered into Gaussian distributed subsets along the principal components of the PCARM's, again using Klustakwik [5]. This effectively removed any samples that were far enough outside the distribution on more than one PCARM, but not necessarily flagged as an outlier on any of the individual metrics alone.

3.4.5 Plots of Sample Outliers

The distributions for each PCARM and any outliers (cluster = 1) found are shown in Figures 12, 13, 14, and 15. Samples are labeled according to Table 5.

Table 5: Sample Legend for Outlier Plots

Grey	Clustered into Gaussian distributed subsets (not Flagged)
Orange	Flagged as outlier based on individual PCARM's
Blue	Flagged as outlier based on PC's of PCARM's
Green	Flagged as outlier for both methods

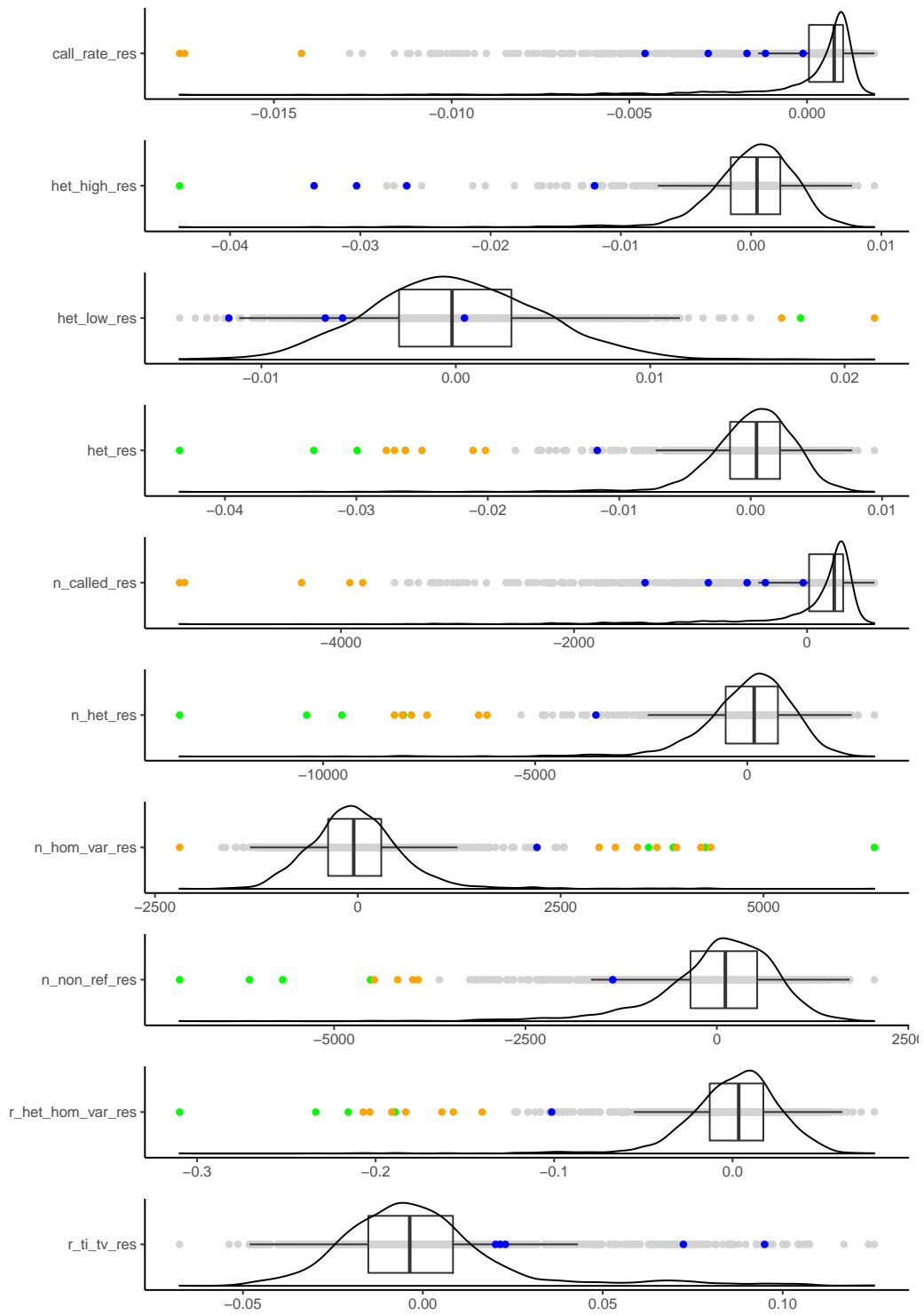


Figure 12: Adjusted sample metric distributions for GWAS

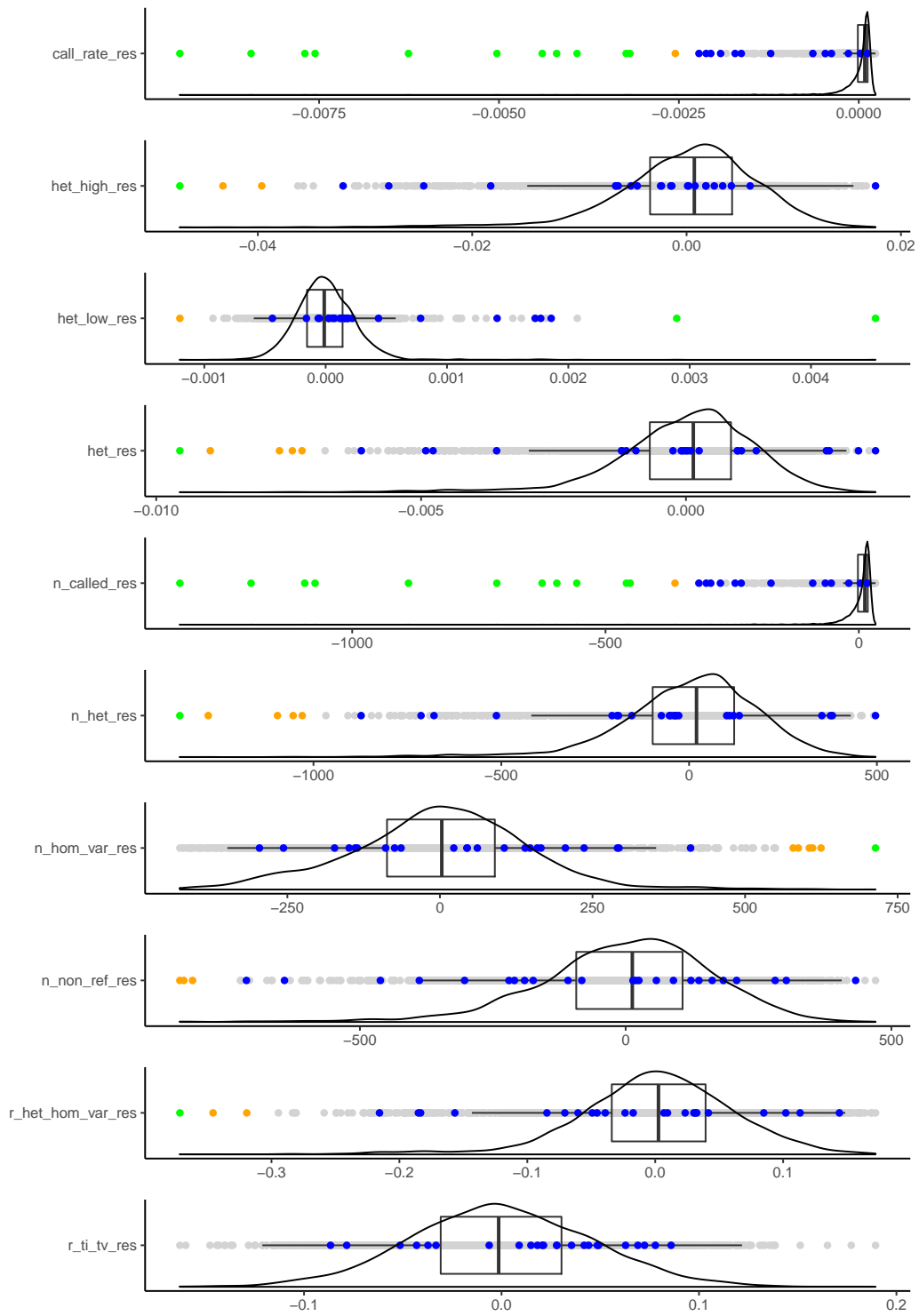


Figure 13: Adjusted sample metric distributions for EXBROAD

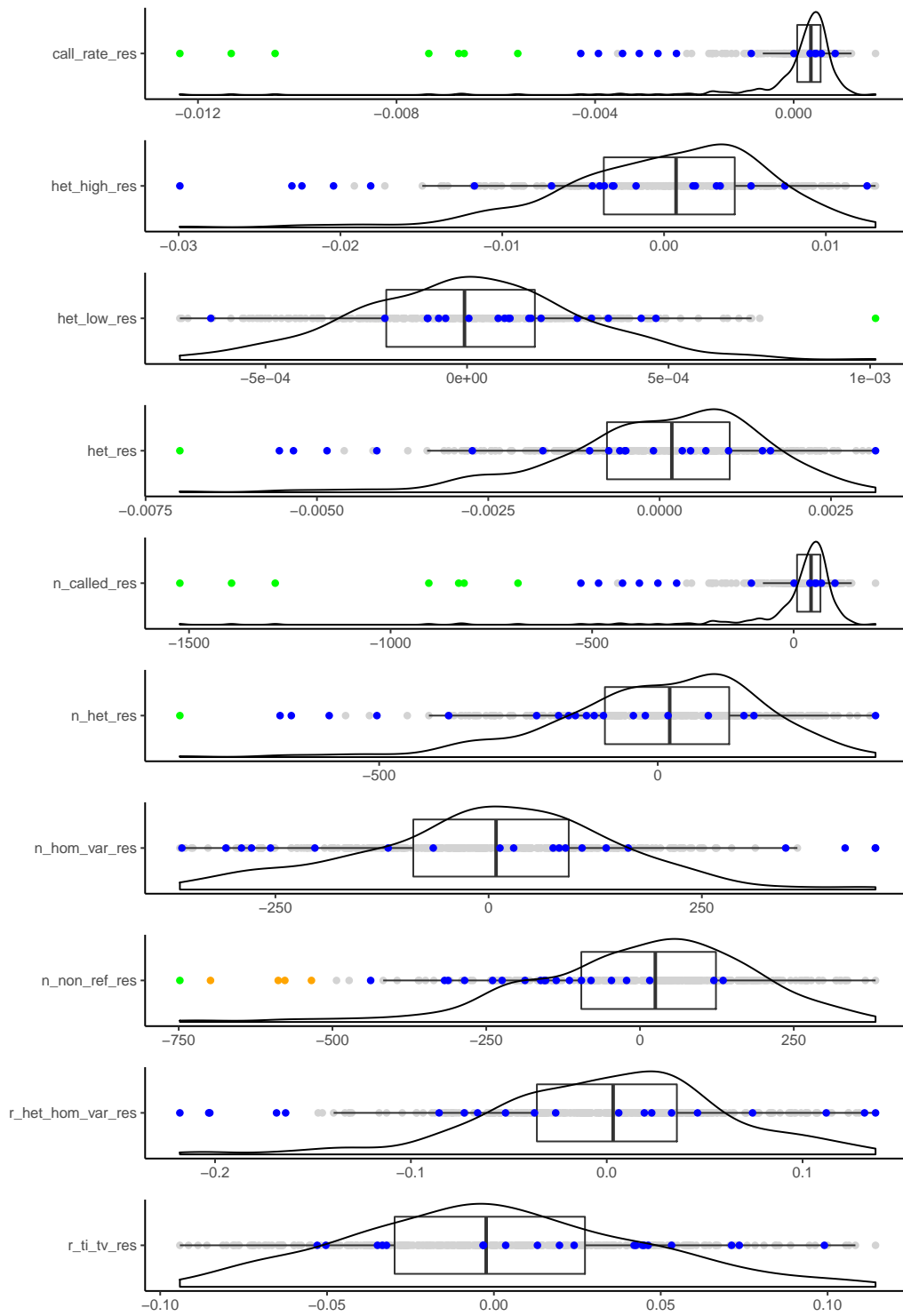


Figure 14: Adjusted sample metric distributions for EXCIDR

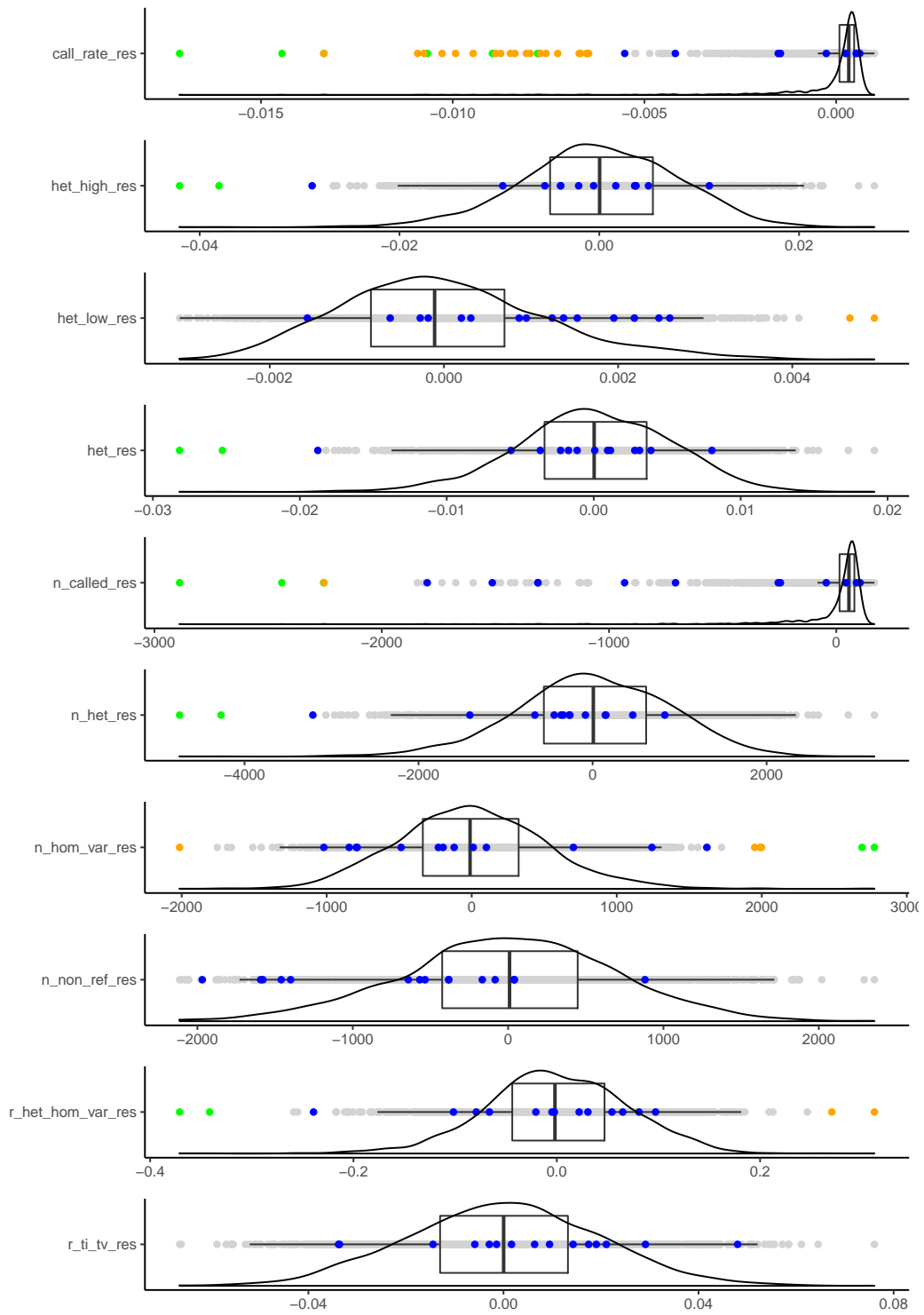


Figure 15: Adjusted sample metric distributions for METABO

3.5 Summary of Sample Outlier Detection

Table 6 contains a summary of outliers detected by each method and across all genotyping technologies. Note that 'PCA(Metrics)' results from the clustering of the PCs of the 8 PCARM's combined, so 'Metrics + PCA(Metrics)' is the union of samples flagged by that method with samples flagged by each of the 10 individual metric clusterings. Figure 16 summarizes the samples remaining for analysis. Upon further inspection, no samples were manually reinstated during this step. More information is available upon request

Table 6: Samples flagged for removal

	EXBROAD	EXCIDR	GWAS	METABO	Total
call_rate	25	20	8	35	87
het_high	26	20	5	15	65
het_low	25	20	7	17	69
het	28	20	11	15	70
n_called	25	20	10	16	70
n_het	28	20	11	15	70
n_hom_var	29	20	13	19	77
n_non_ref	27	24	9	15	74
r_het_hom_var	26	20	12	17	73
r_ti_tv	24	20	5	15	64
PCA(Metrics)	24	20	5	15	64
Metrics+PCA(Metrics)	36	24	19	42	114
Extreme Missingness	0	0	4	0	4
Duplicates	0	0	0	0	0
Cryptic Relatedness	0	0	0	0	0
Sexcheck	0	0	0	1	1
Ancestry Outlier	0	0	0	0	0
Total	39	27	25	46	121

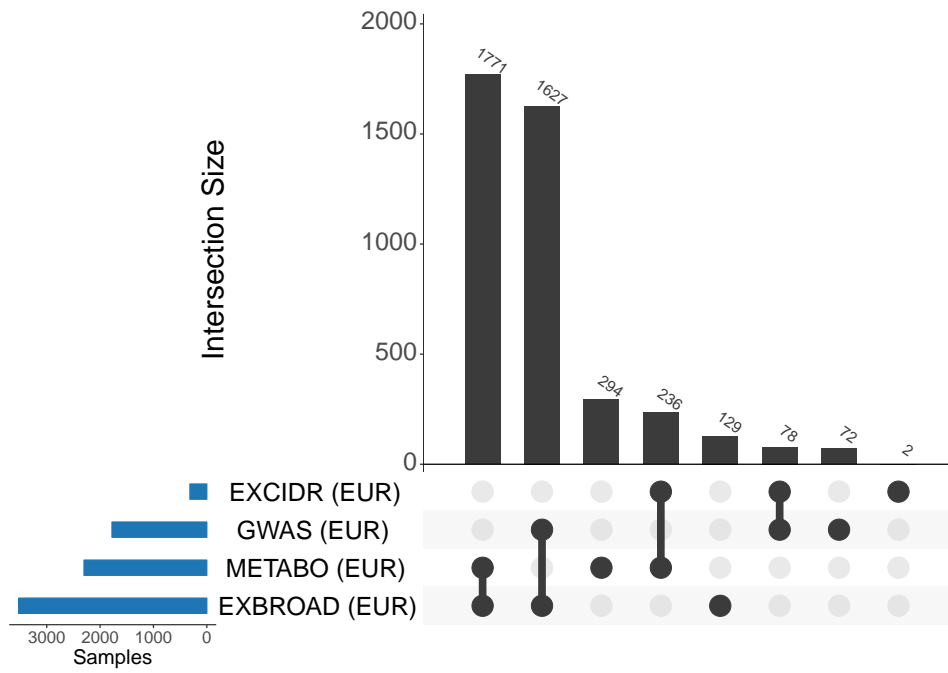


Figure 16: Samples remaining for analysis

4 Variant QC

Variant quality was assessed using call rate and Hardy Weinberg equilibrium (HWE). We calculate HWE using controls only within any of 4 major ancestral populations; EUR, AFR, SAS and EAS. There must have been at least 100 samples in a population to trigger a filter. This conservative approach minimizes the influence from admixture in other population groups. This procedure resulted in flagging 5,374 GWAS, 324 EXBROAD, 685 EXCIDR and 1,935 METABO variants for removal. Figure 17 shows the number of variants remaining for analysis after applying filters.

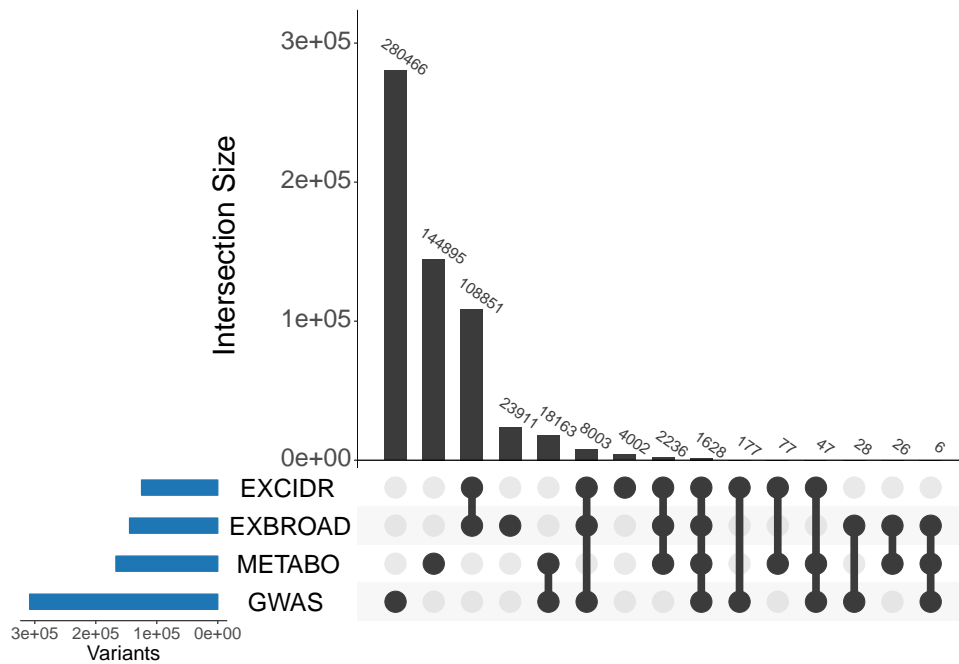


Figure 17: Variants remaining for analysis

5 Acknowledgements

We would like to acknowledge the following people for their significant contributions to this work.

Ryan Koesterer

Jason Flannick

Marcin von Grotthuss

6 References

- [1] Plink1.9, <https://www.cog-genomics.org/plink2>.
- [2] Deelan P, Bonder MJ, Joeri van der Velde K, Westra HJ, Winder E, Hendriksen D, Franke L, Swertz MA. Genotype harmonizer: automatic strand alignment and format conversion for genotype data integration. BMC Research Notes; 2014. 7:901. doi:10.1186/1756-0500-7-901. <https://github.com/molgenis/systemsgenetics/wiki/Genotype-Harmonizer>.
- [3] Conomos MP. GENetic ESTimation and Inference in Structured samples (GENESIS): Statistical methods for analyzing genetic data from samples with population structure and/or relatedness, <https://www.rdocumentation.org/packages/GENESIS/versions/2.2.2>.
- [4] 1000 Genomes Phase 3 v5, https://mathgen.stats.ox.ac.uk/impute/1000GP_Phase3.html.
- [5] Klustakwik, <http://klustakwik.sourceforge.net/>.
- [6] [http://genome.sph.umich.edu/wiki/Regions_of_high_linkage_disequilibrium_\(LD\)](http://genome.sph.umich.edu/wiki/Regions_of_high_linkage_disequilibrium_(LD)).
- [7] <https://www.ncbi.nlm.nih.gov/grc/human/data?asm=GRCh37>.
- [8] <http://people.virginia.edu/~wc9c/KING/>.
- [9] Seed C, Bloemendal A, Bloom JM, Goldstein JI, King D, Poterba T, Neale BM. Hail: An Open-Source Framework for Scalable Genetic Data Analysis. In preparation. <https://github.com/hail-is/hail>.
- [10] Gilbert C, Ruebenacker O, Koesterer R, Massung J, Flannick J. Loamstream. loamstream 1.4-SNAPSHOT (1.3-211-g92f442f) branch: master commit: 92f442fb6952fbc26e474fbf49b6db15a7370677 built on: 2019-05-03T16:01:23.778Z. <https://github.com/broadinstitute/dig-loam-stream>.
- [11] Koesterer R, Gilbert C, Ruebenacker O, Massung J, Flannick J. AMP-DCC Data Analysis Pipeline. dig-loam-2.5.22. <https://github.com/broadinstitute/dig-loam>.