XPASS

The XPASS package implement the XPASS approach for generating PRS in a target population by integrating multi-ethnic datasets.

Installation

```
#install.packages("devtools")
devtools::install_github("YangLabHKUST/XPASS")
```

Quick start

We illustrate the usage of XPASS using the GWAS summary statistics of BMI from UKB and BBJ. For demonstration, we use the easily accessible 1000 Genomes project genotypes as reference panels. However, because these reference panels only contain 377 EAS amples and 417 EUR samples, optimal prediction accuracy of XPASS is not expected. In practice, it is suggested to use larger datasets as reference panels (n>2000). Therefore, we strongly suggest users to use their own reference panels with sufficiently large sample sizes.

Data preparation

The datasets involved in the following example can be downloaded from here.

Input files of XPASS includ:

- summay statistics file of the target population
- summay statistics file of the auxiliary population
- reference panel of the target population in plink 1 format
- reference panel of the auxiliary population in plink 1 format
- covariates file associated with the target population reference panel (optional)
- covariates file associated with the auxiliary population reference panel (optional)

The XPASS format GWAS summary statistics file has 5 fields:

- SNP: SNP rsid
- N: sample size
- Z: Z-scores
- A1: effect allele
- A2: other allele.

Here, we use the BMI GWAS from BBJ male as the target training set and BMI GWAS from UKB as the auxiliary training set

```
$ head BMI_bbj_male_3M_format_3MImp.txt

SNP N Z A1 A2
rs117086422 159095 -1.20413423957762 T C
rs28612348 159095 -1.25827042089233 T C
rs4475691 159095 -1.19842287777303 T C
rs950122 159095 -1.2014434974188 C G
rs3905286 159095 -1.27046106136441 T C
rs28407778 159095 -1.26746342605063 A G
rs4246505 159095 -1.24706211285128 A G
rs4626817 159095 -1.26366297625074 A G
rs11507767 159095 -1.28611566069053 G A
```

```
$ head height_ukb_3M_format.txt

SNP N Z A1 A2
rs117086422 429312 1.42436004338939 T C
rs28612348 429312 1.48706291417224 T C
rs4475691 429312 1.53977372135067 T C
rs950122 429312 1.37958155329171 C G
rs3905286 429312 1.77045946243262 T C
rs28407778 429312 1.9908370573435 A G
rs4246505 429312 1.90922505355565 A G
rs4626817 429312 1.53216668392479 A G
rs11507767 429312 1.55873328059033 G A
```

We keep the BMI GWAS from BBJ female as the external validation dataset:

```
$ head BMI_bbj_male_3M_format_3MImp.txt

SNP N Z A1 A2
rs117086422 72390 0.897252679561414 T C
rs28612348 72390 0.904738461538462 T C
rs4475691 72390 0.89177374486687 T C
rs950122 72390 0.891523178807947 C G
rs3905286 72390 0.827441738675046 T C
rs28407778 72390 0.816801884323476 A G
rs4246505 72390 0.812148186935463 A G
rs4626817 72390 0.811624558188245 A G
rs11507767 72390 0.811100929441026 G A
```

The covariates files should not include row names and column names. The rows should be exactly corresponding to the individuals in the .fam file of reference genotypes. Each column corresponds to one covariate. A column of one should not be included in the file.

```
$ head 1000G.EAS.QC.hm3.ind.pc5.txt

-0.0242863 0.0206888 -0.0028171 0.0343263 0.0211044
-0.025051 0.0275325 -0.0332156 -0.0233166 0.0588989
-0.0198603 0.0286747 0.008464 -0.0215478 0.0189802
-0.0117008 0.0251493 -0.0228276 -0.0670019 0.0186454
-0.0246233 0.0281415 -0.0418795 0.0059947 0.0196607
-0.0270411 0.0127586 -0.0376545 0.0182809 0.0344744
-0.0193744 0.0367263 -0.0176168 -0.0307868 0.0254125
-0.0212299 0.022658 0.0225698 -0.0249273 0.0144324
-0.0159054 0.00558952 -0.00609582 -0.033497 0.0518336
-0.0258843 0.0476758 0.0073353 0.0164056 0.0072118
```

```
$ head 1000G.EUR.QC.hm3.ind.pc20.txt
0.0290072\ 0.0717627\ -0.0314029\ 0.0317316\ 0.0618357\ 0.0385132\ 0.122857
-0.0289581 \ -0.0114267 \ -0.0205926 \ -0.0466983 \ 0.0836711 \ 0.00690379 \ 0.0345008
-0.0179313 0.0109661 -0.0214763 0.0014544 0.0182944 0.0399625
0.0378568 0.0499758 -0.00811504 0.0363021 -0.0579984 0.0422509 0.141279
-0.0167868 -0.0181999 0.0165593 -0.0304088 0.0423324 0.0226001 0.00843853
0.0212477 - 0.0666462 - 0.0787379 \ 0.0136196 \ 0.108933 \ 0.0801246
0.0417318 0.0732938 -0.0409508 0.0176872 0.0801957 0.0124742 0.0252689
-0.0444921 -0.00305238 0.0035535 -0.0070929 0.0240194 -0.00543616 0.0272464
-0.0048309 -0.0207223 0.0415044 0.0494025 0.0213837 -0.0021093
0.0348071 0.0715395 -0.0266058 0.00280025 -0.0166164 0.0440144 0.135709
0.017364 0.0276564 -0.00286321 0.0314583 0.00299185 0.0792055 0.012042
-0.0337269 -0.00999033 0.0186435 -0.0699027 0.00791191 -0.0131168
0.0444763 \ 0.0713348 \ 0.00162451 \ -0.0107805 \ 0.0868909 \ 0.0218014 \ 0.0314216
-0.0429928 -0.0137937 0.00913544 -0.062828 0.0555199 0.0378234 -0.0162297
0.000344947 - 0.0164497 \ 0.0523967 - 0.0861731 - 0.038893 \ 0.028166
0.0300713 0.0522082 -0.0116997 -0.029994 0.0539977 0.0162067 -0.0522507
0.0022036\ 0.0255471\ 0.0129012\ 0.0371803\ 0.0701241\ 0.0314957\ 0.00870374
0.00566795 0.116672 0.0267188 0.0451948 0.00288655 -0.0427304
0.0339424 \ 0.0718272 \ 0.000614329 \ -0.0183555 \ 0.0162768 \ -0.0600599 \ 0.00343218
0.0149793 - 0.0236301 - 0.0267658 \ 0.0387814 - 0.00624387 - 0.0364751 \ 0.00486515
-0.0341221 0.0415286 -0.0274807 -0.013188 0.0695243 0.0495376
0.0423378 \ 0.0656126 \ -0.0331807 \ -0.0361484 \ -0.0155739 \ 0.0557459 \ 0.00428138
0.0840953 - 0.034451 \ 0.0753096 - 0.0180153 \ 0.0595412 - 0.0367107 - 0.0285888
-0.0986386 0.00845412 -0.00388558 -0.0641134 -0.05815 0.0242433
0.0429698 0.0431213 -0.0357636 -0.00270477 -0.0567958 0.0892002 0.0980711
0.0468321 -0.0359592 0.011195 -0.000235849 -0.0192522 -0.00271491 0.0381155
-0.00845755 -0.0171629 -0.026532 -0.0415778 0.0274635 0.122063
0.0290721 \ 0.0541744 \ -0.0238317 \ 0.0254426 \ 0.0986334 \ 0.0706142 \ 0.0977585
0.00427919 - 0.0381976 \ 0.0020029 - 0.0161052 - 0.016666 - 0.00627125 - 0.00490556
-0.0410802 0.0125096 -0.0175252 0.0320359 0.00866061 0.0736791
```

Run XPASS

Once the imput files are formatted, XPASS will automatically process the datasets, including SNPs overlapping and allele matching.

Run XPASS with the following comand:

```
# library(devtools)
# install github("https://github.com/YangLabHKUST/XPASS")
library(XPASS)
library(data.table)
library(RhpcBLASctl)
blas set num threads(30)
# reference genotypes for EAS (prefix of plink file bim/bed/fam)
ref EAS <- "1000G.EAS.QC.hm3.ind"</pre>
# covariates of EAS reference genotypes
cov_EAS <- "1000G.EAS.QC.hm3.ind.pc5.txt"</pre>
# reference genotypes for EUR (prefix of plink file bim/bed/fam)
ref_EUR <- "1000G.EUR.QC.hm3.ind"</pre>
# covariates of EUR reference genotypes
cov EUR <- "1000G.EUR.QC.hm3.ind.pc20.txt"
# genotype file of test data (plink prefix).
# Note: for demonstration, we assume that the genotypes of prediction target
are used as the reference panel of target population.
# In practice, one can also use genotypes from other sources as reference
panel.
BMI_test <- "1000G.EAS.QC.hm3.ind"
# sumstats of height
BMI_bbj_male <- "BMI_bbj_male_3M_format_3MImp.txt" # target
BMI ukb <- "BMI ukb sumstat format all.txt" # auxiliary
BMI_bbj_female <- "BMI_bbj_female_3M_format_3MImp.txt" # external validation
fit bbj <-XPASS(file z1 = BMI bbj male,file z2 = BMI ukb,file ref1 = ref EAS,
                file_ref2 = ref_EUR,
                file cov1 = cov EAS, file cov2 = cov EUR,
                file predGeno = BMI test,
                compPRS=T,
                pop = "EAS",sd_method="LD_block",compPosMean = T,
                file_out = "BMI_bbj_ukb_ref_TGP")
Summary statistics file 1: BMI_bbj_male_3M_format_3MImp.txt
```

```
Summary statistics file 2: BMI ukb sumstat format all.txt
Reference file 1: 1000G.EAS.QC.hm3.ind
Reference file 2: 1000G.EUR.QC.hm3.ind
Covariates file 1: 1000G.EAS.QC.hm3.ind.pc5.txt
Covariates file 2: 1000G.EUR.QC.hm3.ind.pc20.txt
Reading data from summary statisitcs...
3506148 and 3777871 SNPs found in summary statistics files 1 and 2.
Reading SNP info from reference panels...
1209411 and 1313833 SNPs found in reference panel 1 and 2.
746454 SNPs are matched in all files.
0 SNPs are removed because of ambiguity; 746454 SNPs remained.
Calculating kinship matrix from the both reference panels...
127749 SNPs in the second reference panel are alligned for alleles according to
the first.
14337 SNPs have different minor alleles in population 1, z-scores are corrected
according to reference panel.
14332 SNPs have different minor alleles in population 2, z-scores are corrected
according to reference panel.
Assigning SNPs to LD Blocks...
Calculate PVE...
                                 h12
[1,] 0.165790395 0.247603545 0.129399058 0.63866483
[2,] 0.007631346 0.008542654 0.006856057 0.02002658
Predicting PRS from test genotypes...
Done.
```

XPASS output

XPASS returns a list of results, the some key are:

• H: a table of estimated heritabilities, co-heritability and genetic correlation (first row) and their corresponding standard erros (second row).

```
> fit_bbj$H

h1 h2 h12 rho

[1,] 0.165790395 0.247603545 0.129399058 0.63866483

[2,] 0.007631346 0.008542654 0.006856057 0.02002658
```

 mu: a data frame storing the posterior means computed by LDpred-inf using only the target dataset (mu1) and only the auxiliary dataset (mu2), and the posterior mean computed by XPASS (mu_XPASS). SNPs information is also returned: A1 is the effect allele, A2 is the other allele.

```
> head(fit_bbj$mu)
CHR SNP POS A1 A2 mu1 mu2 mu_XPASS

1 1 rs4475691 846808 T C -1.271443e-04 -0.0001837393 -0.0003248579

2 1 rs7537756 854250 G A -4.778206e-05 -0.0002169705 -0.0002979870

3 1 rs3748592 880238 A G -3.201406e-04 -0.0008911591 -0.0007477507

4 1 rs2340582 882803 A G -3.396992e-04 -0.0009056487 -0.0008076512

5 1 rs4246503 884815 A G -3.318260e-04 -0.0009265197 -0.0008148705

6 1 rs3748597 888659 T C -3.327131e-04 -0.0009142760 -0.0008113772
```

• PRS (if file_predGeno provided and compPRS=T): a data frame storing the PRS generated using mu1, mu2 and mu_XPASS, respectively.

```
> head(fit bbj$PRS)
     FID IID
                      PRS1
                                 PRS2 PRS_XPASS
1 HG00403 HG00403 0.14310343 0.92040953 0.55960909
2 HG00404 HG00404 -0.19478307 0.63289820 0.03773899
3 HG00406 HG00406 -0.09440555 0.69099450 0.14173516
4 HG00407 HG00407 0.05533466 0.07036618 0.05091082
5 HG00409 HG00409 0.25063980 1.54556843 0.83527097
6 HG00410 HG00410 0.13454734 0.23143305 0.10214319
# One can also compute PRS after fitting the model:
> PRS <- predict XPASS(fit bbj$mu,ref EAS)
> head(PRS)
     FID IID PRS1 PRS2 PRS XPASS
1 HG00403 HG00403 0.14310343 0.92040953 0.55960909
2 HG00404 HG00404 -0.19478307 0.63289820 0.03773899
3 HG00406 HG00406 -0.09440555 0.69099450 0.14173516
4 HG00407 HG00407 0.05533466 0.07036618 0.05091082
5 HG00409 HG00409 0.25063980 1.54556843 0.83527097
6 HG00410 HG00410 0.13454734 0.23143305 0.10214319
```

XPASS will also write above outputs into the files with file_out prefix, if provided.

External validation using independent GWAS data

Because the WeGene data is currently not available online, we use the GWAS of female BMI from BBJ as the external validation dataset to approximate the prediction \mathbb{R}^2 . Specifically we use the following equation:

$$R^2 = corr(y,\hat{y})^2 = \left(rac{cov(y,\hat{y})}{\sqrt{var(y)var(\hat{y})}}
ight)^2 = \left(rac{z^T ilde{\mu}/\sqrt{n}}{\sqrt{ ilde{\mu}^T\Sigma ilde{\mu}}}
ight)^2,$$

where z is the z-score of external summsry statistics, n is its sample size, $\tilde{\mu}$ is the posterior mean of effect size at the standardized genotype scale, Σ is the LD reference panel.

While the reference panels have only limmited samples, XPASS still achieves 30% relative improvement compared to LDpred-inf in terms of \mathbb{R}^2 .

Prediction of Type 2 Diabetes

We provide an additional example with common disease Type 2 Diabetes (T2D). We use the GWAS of males from BBJ as traing set and that of females as testing data, UKB GWAS as auxiliary dataset, and 1000 Genomes as reference panels. Datasets are available here.

```
# reference genotypes for EAS (prefix of plink file bim/bed/fam)
ref_EAS <- "1000G.EAS.QC.hm3.ind"</pre>
# covariates of EAS reference genotypes
cov_EAS <- "1000G.EAS.QC.hm3.ind.pc5.txt"</pre>
# reference genotypes for EUR (prefix of plink file bim/bed/fam)
ref_EUR <- "1000G.EUR.QC.hm3.ind"</pre>
# covariates of EUR reference genotypes
cov_EUR <- "1000G.EUR.QC.hm3.ind.pc20.txt"</pre>
# genotype file of test data (plink prefix).
T2D test <- "1000G.EAS.QC.hm3.ind"
# sumstats of height
T2D bbj_male <- "T2D_bbj_male_3M_format.txt" # target</pre>
T2D_ukb <- "T2D_UKB_summary_format.txt" # auxiliary</pre>
T2D_bbj_female <- "T2D_bbj_female_3M_format.txt" # external validation
fit bbj <-XPASS(file z1 = T2D bbj male,file z2 = T2D ukb,file ref1 = ref EAS,
                file_ref2 = ref_EUR,
                file cov1 = cov EAS, file cov2 = cov EUR,
                file_predGeno = T2D_test,
                compPRS=T,
                pop = "EAS",sd_method="LD_block",compPosMean = T,
                file_out = "T2D_bbj_ukb_ref_TGP")
Summary statistics file 1: T2D_bbj_male_3M_format.txt
Summary statistics file 2: T2D_UKB_summary_format.txt
```

```
Reference file 1: 1000G.EAS.QC.hm3.ind
Reference file 2: 1000G.EUR.QC.hm3.ind
Covariates file 1: 1000G.EAS.QC.hm3.ind.pc5.txt
Covariates file 2: 1000G.EUR.QC.hm3.ind.pc20.txt
Reading data from summary statisitcs...
3701030 and 11971734 SNPs found in summary statistics files 1 and 2.
Reading SNP info from reference panels...
1209411 and 1313833 SNPs found in reference panel 1 and 2.
754466 SNPs are matched in all files.
0 SNPs are removed because of ambiguity; 754466 SNPs remained.
Calculating kinship matrix from the both reference panels...
128611 SNPs in the second reference panel are alligned for alleles according to
the first.
233891 SNPs have different minor alleles in population 1, z-scores are
corrected according to reference panel.
460962 SNPs have different minor alleles in population 2, z-scores are
corrected according to reference panel.
Assigning SNPs to LD Blocks...
Calculate PVE...
                        h2
                                    h12
              h 1
                                               rho
[1,] 0.096949968 0.04137762 0.041067706 0.64840131
[2,] 0.006767725 0.00166678 0.002257857 0.02400119
Predicting PRS from test genotypes...
```

We compute the prediction \mathbb{R}^2 as

For T2D, XPASS achieves 23.5% relative improvement interms of prediction \mathbb{R}^2 .

Development

The XPASS package is developed by Mingxuan Cai (mcaiad@ust.hk).

Contact information

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