# Comparison: Michigan Imputation Server / Shapeit+Minimac3 (Chromosome 7)

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### Imputation

```
## Loading required package: BiocGenerics
## Loading required package: parallel
##
## Attaching package: 'BiocGenerics'
## The following objects are masked from 'package:parallel':
##
       clusterApply, clusterApplyLB, clusterCall, clusterEvalQ,
##
       clusterExport, clusterMap, parApply, parCapply, parLapply,
##
       parLapplyLB, parRapply, parSapply, parSapplyLB
## The following objects are masked from 'package:stats':
##
##
       IQR, mad, sd, var, xtabs
## The following objects are masked from 'package:base':
##
       anyDuplicated, append, as.data.frame, basename, cbind,
##
##
       colMeans, colnames, colSums, dirname, do.call, duplicated,
##
       eval, evalq, Filter, Find, get, grep, grepl, intersect,
##
       is.unsorted, lapply, lengths, Map, mapply, match, mget, order,
       paste, pmax, pmax.int, pmin, pmin.int, Position, rank, rbind,
##
       Reduce, rowMeans, rownames, rowSums, sapply, setdiff, sort,
##
       table, tapply, union, unique, unsplit, which, which.max,
##
       which.min
## Loading required package: GenomeInfoDb
## Loading required package: S4Vectors
## Loading required package: stats4
## Attaching package: 'S4Vectors'
## The following object is masked from 'package:base':
##
##
       expand.grid
```

```
## Loading required package: IRanges
## Loading required package: GenomicRanges
## Loading required package: SummarizedExperiment
## Loading required package: Biobase
## Welcome to Bioconductor
##
##
       Vignettes contain introductory material; view with
##
       'browseVignettes()'. To cite Bioconductor, see
       'citation("Biobase")', and for packages 'citation("pkgname")'.
##
## Loading required package: DelayedArray
## Loading required package: matrixStats
## Attaching package: 'matrixStats'
## The following objects are masked from 'package:Biobase':
##
##
       anyMissing, rowMedians
## Loading required package: BiocParallel
##
## Attaching package: 'DelayedArray'
## The following objects are masked from 'package:matrixStats':
##
##
       colMaxs, colMins, colRanges, rowMaxs, rowMins, rowRanges
## The following objects are masked from 'package:base':
##
##
       aperm, apply
## Loading required package: Rsamtools
## Loading required package: Biostrings
## Loading required package: XVector
## Attaching package: 'Biostrings'
## The following object is masked from 'package:DelayedArray':
##
##
       type
```

```
## The following object is masked from 'package:base':
##
##
       strsplit
##
## Attaching package: 'VariantAnnotation'
## The following object is masked from 'package:base':
##
##
       tabulate
## Loading required package: survival
## Loading required package: Matrix
##
## Attaching package: 'Matrix'
## The following object is masked from 'package:VariantAnnotation':
##
##
       expand
## The following object is masked from 'package:S4Vectors':
##
##
       expand
## Loading required package: magrittr
```

#### Imputed data exploration

```
michigan_chr7
```

```
## class: CollapsedVCF
## dim: 192 2275
## rowRanges(vcf):
     GRanges with 5 metadata columns: paramRangeID, REF, ALT, QUAL, FILTER
## info(vcf):
     DataFrame with 4 columns: AF, MAF, R2, ER2
## info(header(vcf)):
##
          Number Type Description
##
      AF 1
                 Float Estimated Alternate Allele Frequency
     MAF 1
##
                 Float Estimated Minor Allele Frequency
##
     R2 1
                 Float Estimated Imputation Accuracy
##
     ER2 1
                 Float Empirical (Leave-One-Out) R-square (available only ...
## geno(vcf):
##
     SimpleList of length 3: GT, DS, GP
## geno(header(vcf)):
##
         Number Type
                       Description
##
     GT 1
                String Genotype
##
     DS 1
                Float Estimated Alternate Allele Dosage : [P(0/1)+2*P(1/1)]
##
      GP 3
                Float Estimated Posterior Probabilities for Genotypes 0/0...
```

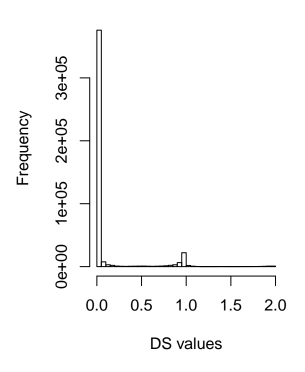
#### minimac\_chr7

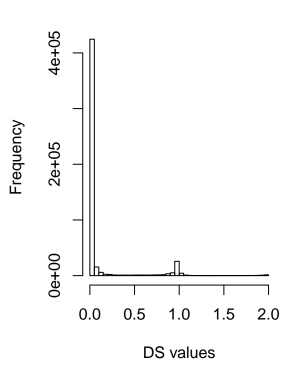
```
## class: CollapsedVCF
## dim: 223 2275
## rowRanges(vcf):
    GRanges with 5 metadata columns: paramRangeID, REF, ALT, QUAL, FILTER
## info(vcf):
    DataFrame with 4 columns: AF, MAF, R2, ER2
## info(header(vcf)):
##
         Number Type Description
##
     AF 1
                Float Estimated Alternate Allele Frequency
##
     MAF 1
                Float Estimated Minor Allele Frequency
##
     R2 1
                Float Estimated Imputation Accuracy
##
     ER2 1
                Float Empirical (Leave-One-Out) R-square (available only ...
## geno(vcf):
##
    SimpleList of length 3: GT, DS, GP
## geno(header(vcf)):
##
        Number Type Description
##
     GT 1
              String Genotype
               Float Estimated Alternate Allele Dosage : [P(0/1)+2*P(1/1)]
##
     DS 1
     GP 3
##
               Float Estimated Posterior Probabilities for Genotypes 0/0...
```

#### DS values

# **Michigan Imputation Server**

### Minimac3





# DS correlation by individuals
min(cor\_by\_ind)

## [1] 0.4930321

max(cor\_by\_ind)

## [1] 0.9999955

mean(cor\_by\_ind)

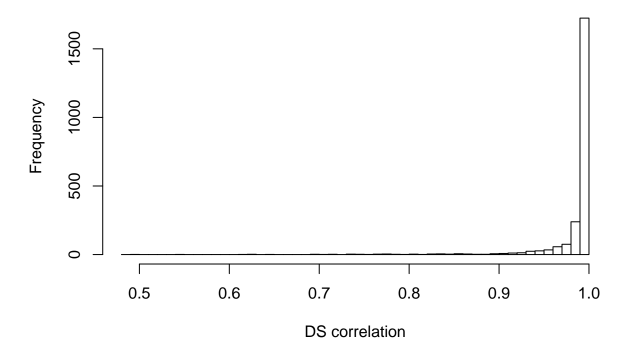
## [1] 0.9857534

mean(cor\_by\_ind > 0.95)

## [1] 0.9358242

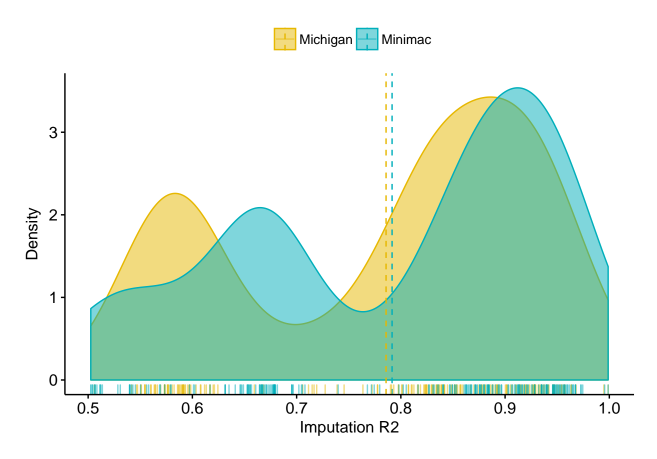
mean(cor\_by\_ind > 0.99)

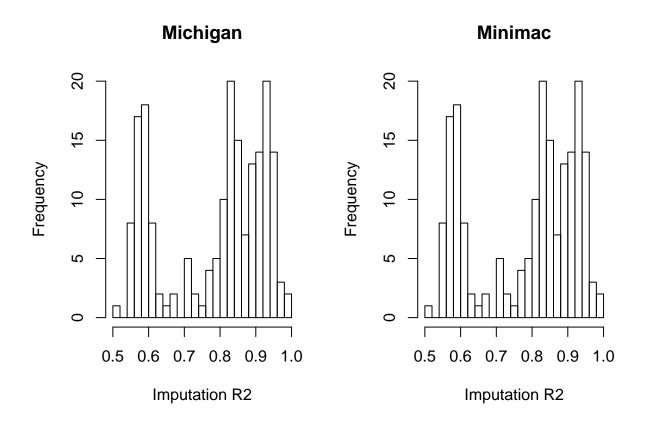
### DS correlation values by individuals



#### $\mathbf{R^2}$

Density and histogram plots comparing the RS2 values in both methods ("ismich = TRUE" indicates the values for the Michigan imputation, whereas "ismich = FALSE" shows the values for the Minimac imputation)





#### Genotype predictions

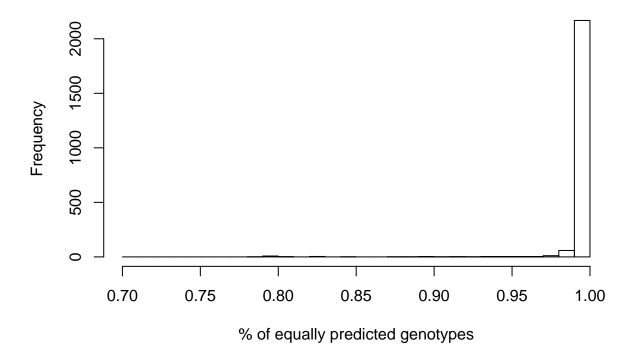
```
\mbox{\tt \#\#} non-single nucleotide variations are set to NA \mbox{\tt \#\#} non-single nucleotide variations are set to NA
```

Compare the genotype predictions (BestGuess) with each method by individuals. "perc\_by\_ind" is the % of SNPs by individual predicted equally in both methods

```
min(perc_by_ind)
## [1] 0.7864583
max(perc_by_ind)
## [1] 1
```

```
mean(perc_by_ind)
```

### SNPs (genotypes) equally predicted with Michigan and Minimac3

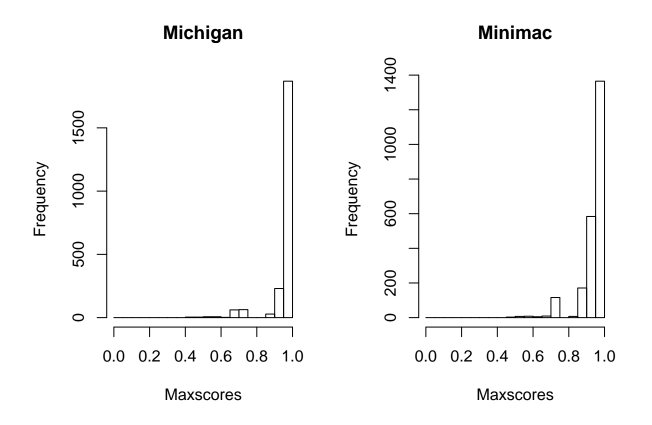


### Inversion prediction

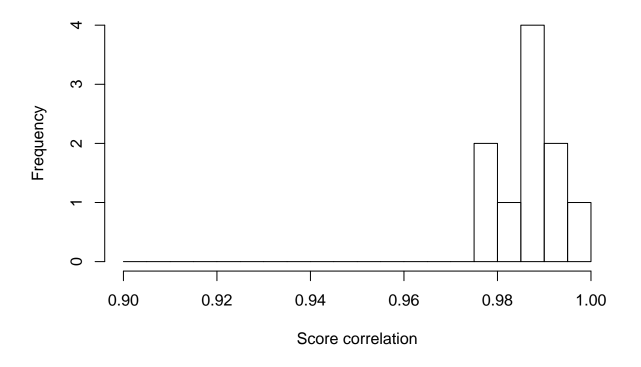
Predicted inversions with scoreInvHap

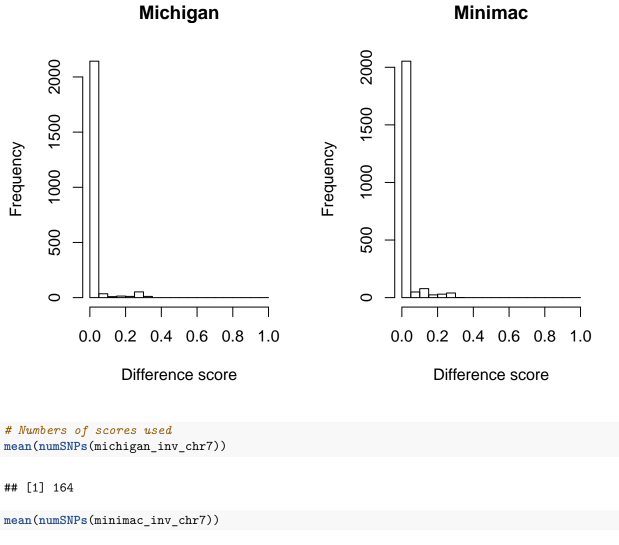
```
michigan_inv_chr7
```

```
## scoreInvHapRes
## Samples: 2275
## Genotypes' table:
## IaIa
            IaIb
                            NaIa
                                    NaIb
                                            NaNa
                                                    NaNb
                                                            NbIa
                                                                   NbIb
                                                                            NbNb
                    IbIb
        5
            367
                    631
                            0
                               1036
                                            21 123
## - Inversion genotypes' table:
## NN
        NI II
## 1039
            775
                    461
## - Inversion frequency: 37.30%
minimac_inv_chr7
## scoreInvHapRes
## Samples: 2275
## Genotypes' table:
                                                            NbIa
                                                                    NbIb
                                                                            NbNb
## IaIa
            IaIb
                    IbIb
                            NaIa
                                    NaIb
                                            NaNa
                                                    NaNb
        70 433
                    565
                            0
                               970
                                      2
                                            21 118
## - Inversion genotypes' table:
## NN
        NI II
## 979 704
                592
## - Inversion frequency: 41.49%
# Comparison table
scoreinvhap_table
          Minimac
##
## Michigan IaIa IaIb IbIb NaIa NaIb NaNa NaNb NbIa NbIb NbNb
##
      IaIa
             89
                   0
                        0
                             0
                                  0
                                       0
                                            0
##
      IaIb
              0
                   4
                        0
                             1
                                  0
                                       0
                                            0
                                                 0
                                                      0
                                                           0
      IbIb
                                      23
##
              0
                   0 344
                             0
                                  0
                                            0
                                                 0
##
      NaIa
              0
                  66
                        1 564
                                  0
                                       0
                                            0
                                                      0
      NaIb
                  0
                                       0
##
              0
                        0
                             0
                                  0
                                            0
                                                 0
                                                      0
                                                           0
                 0
##
      NaNa
            0
                       88
                             0
                                  0
                                     947
                                            1
                                                 0
                                                      0
                                                           0
##
      NaNb
            0 0
                      0
                                     0
                                            0
                                                 0
##
      NbIa
              0
                 0
                        0
                             0
                                  0
                                       0
                                            0
                                                21
                                                      0
                                                           0
##
      NbIb
              0
                   0
                        0
                             0
                                  0
                                       0
                                            1
                                                 0 115
                                                           7
##
      NbNb
              0
                   0
                             0
                                       0
                                            0
                                                      3
sum(diag(scoreinvhap_table))/sum(scoreinvhap_table)
## [1] 0.916044
# Comparison of the results for both imputation methods
par(mfrow=c(1,2))
hist(maxscores(michigan_inv_chr7), breaks=seq(0, 1, by=0.05), main="Michigan", xlab="Maxscores")
hist(maxscores(minimac_inv_chr7), breaks=seq(0, 1, by=0.05), main="Minimac", xlab="Maxscores")
```



# Score correlation by individuals





```
## [1] 164

mean(numSNPs(minimac_inv_chr7))

## [1] 184

# Number of samples in both imputation methods before and after QC filtering
length(classification(michigan_inv_chr7))

## [1] 2275

length(classification(michigan_inv_chr7, minDiff = 0.1, callRate = 0.9))

## [1] 97

length(classification(michigan_inv_chr7, minDiff = 0.1, callRate = 0.9))/
length(classification(michigan_inv_chr7))
```

```
length(classification(minimac_inv_chr7))

## [1] 2275

length(classification(minimac_inv_chr7, minDiff = 0.1, callRate = 0.9))

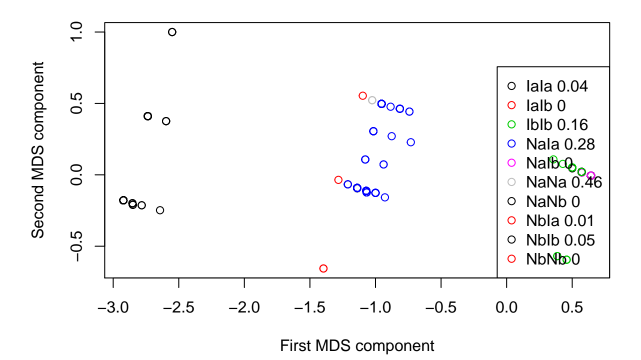
## [1] 173

length(classification(minimac_inv_chr7, minDiff = 0.1, callRate = 0.9))/
    length(classification(minimac_inv_chr7))
```

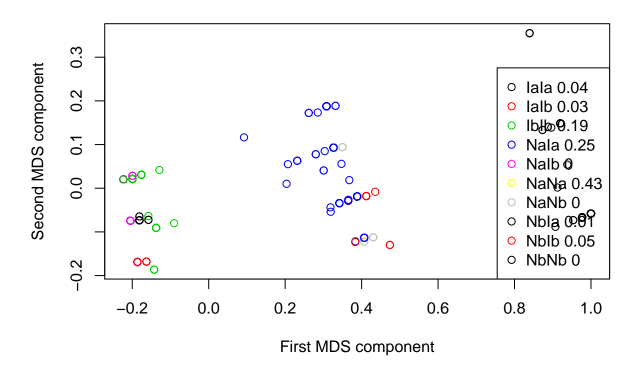
## [1] 0.07604396

#### Plots with invClust

```
# Michigan
par(mfrow=c(1,1))
plotInv(michigan_invclust_chr7, classification = classification(michigan_inv_chr7))
```



```
# Minimac
plotInv(minimac_invclust_chr7, classification = classification(minimac_inv_chr7))
```



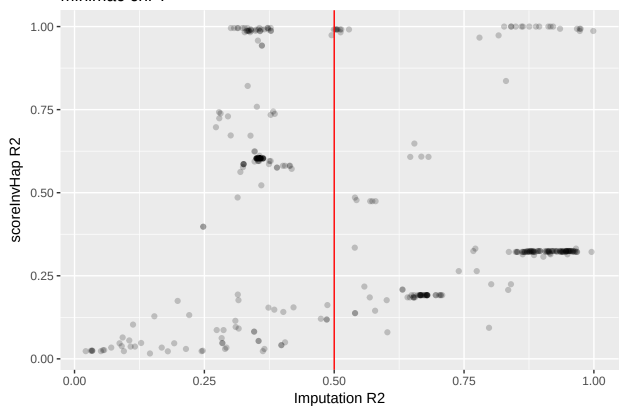
### No filtered imputed data

```
nofilter_minimac_7
```

```
## class: CollapsedVCF
## dim: 1543 2275
## rowRanges(vcf):
     GRanges with 5 metadata columns: paramRangeID, REF, ALT, QUAL, FILTER
## info(vcf):
    DataFrame with 4 columns: AF, MAF, R2, ER2
##
##
  info(header(vcf)):
##
          Number Type Description
                 Float Estimated Alternate Allele Frequency
##
      AF
##
      MAF 1
                 Float Estimated Minor Allele Frequency
##
      R2 1
                 Float Estimated Imputation Accuracy
      ER2 1
                 Float Empirical (Leave-One-Out) R-square (available only ...
##
## geno(vcf):
     SimpleList of length 3: GT, DS, GP
## geno(header(vcf)):
```

```
##
         Number Type Description
##
     GT 1
                String Genotype
                Float Estimated Alternate Allele Dosage : [P(0/1)+2*P(1/1)]
##
      GP 3
                Float Estimated Posterior Probabilities for Genotypes 0/0...
##
nofilter_michigan_7
## class: CollapsedVCF
## dim: 1543 2275
## rowRanges(vcf):
     GRanges with 5 metadata columns: paramRangeID, REF, ALT, QUAL, FILTER
## info(vcf):
    DataFrame with 4 columns: AF, MAF, R2, ER2
## info(header(vcf)):
##
          Number Type Description
##
                 Float Estimated Alternate Allele Frequency
##
     MAF 1
                 Float Estimated Minor Allele Frequency
##
                 Float Estimated Imputation Accuracy
     R2 1
##
     ER2 1
                 Float Empirical (Leave-One-Out) R-square (available only ...
## geno(vcf):
     SimpleList of length 3: GT, DS, GP
## geno(header(vcf)):
##
         Number Type
                       Description
##
                String Genotype
##
                Float Estimated Alternate Allele Dosage : [P(0/1)+2*P(1/1)]
     DS 1
##
      GP 3
                Float Estimated Posterior Probabilities for Genotypes 0/0...
nofilter_minimac_inv_7
## scoreInvHapRes
## Samples: 2275
## Genotypes' table:
## IaIa
             IaIb
                                                             NbIa
                                                                      NbTb
                                                                              NbNb
                     IbIb
                             NaIa
                                     NaIb
                                             NaNa
                                                     NaNb
## 89
        261
                 185
                         368
                                 710
                                         508
                                                 2
                                                     27 117
                                                                 8
## - Inversion genotypes' table:
## NN
        NI II
## 518 1222
                 535
## - Inversion frequency: 50.37%
nofilter_michigan_inv_7
## scoreInvHapRes
## Samples: 2275
## Genotypes' table:
                     IbIb
## IaIa
             IaIb
                                                     NaNb
                                                             NbIa
                                                                      NbIb
                                                                              NbNb
                             NaIa
                                     NaIb
                                             NaNa
## 89
            536
                     595
                                             22 121
         40
                             0
                                 866
                                         1
## - Inversion genotypes' table:
## NN
         NI II
## 872 738
                 665
## - Inversion frequency: 45.45%
```

#### Minimac chr 7



#### Filtered chr7 No filtered chr7 Frequency Frequency 0.8 0.0 0.2 0.4 0.6 1.0 0.0 0.2 0.4 0.6 8.0 1.0 scoreInvHapR2 scoreInvHapR2

#Correlation between Imputation R2 and scoreInvHap R2 (NO filtered data)
cor(info(nofilter\_minimac\_7)[snps\_minimac\_7,]\$R2, SNPsR2\$inv7p11.2[snps\_minimac\_7])

## [1] -0.08281116

# Comparison table scoreInvHap with filtered and no filtered data scoreinvhap\_table\_filt

```
##
                Filtered
## No_filtered IaIa IaIb IbIb NaIa NaIb NaNa NaNb NbIa NbIb NbNb
##
           IaIa
                    89
                           0
                                 0
                                                   0
                                                               0
                                                                     0
                                                                           0
           IaIb
                                    223
##
                     0
                          38
                                 0
                                             0
                                                   0
                                                               0
                                                                     0
                                                                           0
##
           IbIb
                     0
                           0
                                71
                                       0
                                                114
                                                         0
                                                                     0
                                                                           0
           NaIa
                          32
                                    336
##
                     0
                                 0
                                                   0
##
           NaIb
                     0
                           0
                              272
                                       0
                                             0
                                                438
                                                         0
                                                                     0
           NaNa
                     0
                           0
                                90
                                       0
                                             0
                                                418
                                                               0
                                                                     0
##
##
           NaNb
                     0
                           0
                                       0
                                                   0
                                                               0
                                                                     1
                                                   0
                                                              21
##
           NbIa
                     0
##
           NbIb
                     0
                           0
                                 0
                                       0
                                             0
                                                   0
                                                         1
                                                               0
                                                                  116
                                                                           0
##
           NbNb
                     0
                                       0
                                                   0
                                                         0
```

sum(diag(scoreinvhap\_table\_filt))/sum(scoreinvhap\_table\_filt)

