Constructing G matrix with missing SNPs:

Handling missing SNPs when calculating the **genomic relationship matrix** (**G**; VanRaden (2008)) is relatively straightforward. The standard procedure used for complete data can still be followed, with one key modification: when multiplying the **Z matrix** by its transpose, the missing genotype values should be treated as zero. This ensures that missing markers do not contribute to the calculation, effectively excluding them from the computation as if they were not present.

Example:

Suppose we have the following genotype matrix **M** for 3 animals and 2 SNPs, where SNP #1 for animal #2 is missing:

$$\mathbf{M} = \begin{bmatrix} 1 & 2 \\ NA & 0 \\ 2 & 0 \end{bmatrix}$$

If you are using \mathbf{R} , missing values should be represented by NA. For instance, if missing values were previously coded as 5, you can convert them to NA using:

```
M[M==5]=NA
```

Now, the allele frequencies and the **P matrix** can be calculated in the usual way, without requiring any special treatment for the missing data:

```
n0 \leftarrow apply(M==0, 2, sum, na.rm=T)
n1 <- apply(M==1,2,sum,na.rm=T)</pre>
n2 <- apply(M==2,2,sum,na.rm=T)</pre>
# Compute Z as described in Mrode 2015, page 180.
freq <-((2*n2)+n1)/(2*(n0+n1+n2))
#[1] 0.7500000 0.3333333
freq2 <- 2*(freq)
# create matrix P
P <- matrix(freq2, nrow=1)
P \leftarrow P[rep(1:nrow(P), times = nrow(M)),]
P <- round (P, digits = 2) # This is just to round the outputs!
Р
      [,1] [,2]
#[1,] 1.5 0.67
#[2,] 1.5 0.67
#[3,] 1.5 0.67
```

Next, comput **Z** matrix:

```
Z = M - P
```

```
Z

# [,1] [,2]

#[1,] -0.5 1.33

#[2,] NA -0.67

#[3,] 0.5 -0.67
```

Since we need to multiply **Z** by its transpose, when replacing NAs by 0s, the missing SNPs do not contribute to the result:

The matrix Z above is:

$$\mathbf{Z} = \begin{bmatrix} -0.5 & 1.33 \\ 0 & -0.67 \\ 0.5 & -0.67 \end{bmatrix}$$
 The missing SNP = NA as above

And its transpose is:

$$\mathbf{Z}' = \begin{bmatrix} -0.5 & 0 & 0.5 \\ 1.33 & -0.67 & -0.67 \end{bmatrix}$$

Now we can construct the **ZZ'**:

$$\begin{aligned} \mathbf{ZZ'} \\ &= \begin{bmatrix} (-0.5 \times -0.5) + (1.33 \times 1.33) & (-0.5 \times 0.0) + (1.33 \times -0.67) & (-0.5 \times 0.5) + (1.33 \times -0.67) \\ (0.0 \times -0.5) + (-0.67 \times 1.33) & (0.0 \times 0.0) + (-0.67 \times -0.67) & (0.0 \times 0.5) + (-0.67 \times -0.67) \\ (0.5 \times -0.5) + (-0.67 \times 1.33) & (0.5 \times 0.0) + (-0.67 \times -0.67) & (0.5 \times 0.5) + (-0.67 \times -0.67) \end{bmatrix} \end{aligned}$$

Note that the missing SNP did not contribute to the results (since they were multiplied by 0s).

Accordingly, the **ZZ'** is:

$$\mathbf{ZZ'} = \begin{bmatrix} 2.0189 & -0.8911 & -1.1411 \\ -0.8911 & 0.4489 & 0.4489 \\ -1.1411 & 0.4489 & 0.6989 \end{bmatrix}$$

Another example:

Suppose we have this SNP data (M):

$$\mathbf{M} = \begin{bmatrix} 1 & 0 \\ NA & 2 \\ 0 & 1 \end{bmatrix}$$

And suppose we calculated the allele frequencies and they were:

$$\mathbf{P} = \begin{bmatrix} -0.65 & 0.32 \\ -0.65 & 0.32 \\ -0.65 & 0.32 \end{bmatrix}$$

No,
$$\mathbf{Z} = \mathbf{M} - \mathbf{P}$$

$$\mathbf{Z} = \begin{bmatrix} 1.65 & -0.32 \\ NA & 1.68 \\ 0.65 & 0.68 \end{bmatrix}$$

Replace NAs by 0s:

$$\mathbf{Z} = \begin{bmatrix} 1.65 & -0.32 \\ \mathbf{0} & 1.68 \\ 0.65 & 0.68 \end{bmatrix} \quad \text{and} \quad \mathbf{Z}' = \begin{bmatrix} 1.65 & \mathbf{0} & 0.68 \\ -0.32 & 1.68 & 0.68 \end{bmatrix}$$

Then **ZZ'** is:

```
\mathbf{ZZ'} = \begin{bmatrix} (1.65 \times 1.65) + (-0.32 \times -0.32) & (-0.32 \times 1.68) & (1.65 \times 0.65) + (-0.32 \times 0.68) \\ (1.68 \times -0.32) & (1.68 \times 1.68) & (1.68 \times 0.68) \\ (0.65 \times 1.65) + (0.68 \times -0.32) & (0.68 \times 1.68) & (0.65 \times 0.65) + (0.68 \times 0.68) \end{bmatrix}
```

```
rm(list=ls(all=TRUE))
M <- read.table(textConnection("</pre>
                                  1 0
                                  NA 2
                                  0 1
                                  "), header=F)
M <- as.matrix(M)</pre>
closeAllConnections()
colnames(M) <- NULL
P <- read.table(textConnection("</pre>
                                  -0.65 0.32
                                  -0.65 0.32
                                  -0.65 0.32
                                  "), header=F)
P <- as.matrix(P)</pre>
closeAllConnections()
colnames(P) <- NULL
n0 <- apply(M==0,2,sum,na.rm=T)
n1 <- apply(M==1,2,sum,na.rm=T)</pre>
n2 <- apply(M==2,2,sum,na.rm=T)</pre>
# Compute Z as described in Mrode 2015, page 180.
freq <-((2*n2)+n1)/(2*(n0+n1+n2))
frea
freq2 <- 2*(freq)
Z < - M - P
Z[is.na(Z)] = 0
Zp < -t(Z)
Z%*%Zp
D \leftarrow 2*(sum(freq*(1-freq)))
G <- Z%*%Zp/D
# OR :
G <- tcrossprod(Z)/D
```

References:

VanRaden, Paul M. "Efficient methods to compute genomic predictions." Journal of dairy science 91.11 (2008): 4414-4423.