```
In [1]: include("/home/nicole/Jupyter/SSBRnoJ/src/SSBR.jl")
using SSBR

In [2]: function getPos(ped,IDs)
    posAi = Array(Int64,size(IDs,1))
    for (i,id) = enumerate(IDs[:,1])
        posAi[i] = ped.idMap[id].seqID
    end
    return posAi
end

Out[2]: getPos (generic function with 1 method)

In [3]: ; cd Data/0.5g2k/G/1
    /home/nicole/Jupyter/JG3/Data/0.5g2k/G/1
```

```
In [4]: ;ls
        Correlation.G5.G.JC.txt
        G0.Genotype.ID
        G0.ID
        G0.noGenotype.ID
        G1.Genotype.ID
        G1.ID
        G1.noGenotype.ID
        G2.Genotype.ID
        G2.ID
        G2.noGenotype.ID
        G3.Genotype.ID
        G3.ID
        G3.noGenotype.ID
        G4.Genotype.ID
        G4.ID
        G4.noGenotype.ID
        G5.Genotype.ID
        G5.ID
        G5.noGenotype.ID
        GenNF.txt
        PedAll.txt
        Phe.txt
        PheAll.txt
        Regression.G5.G.JC.txt
        all.ID
        genotype.ID
        noGenotype.ID
        sim.bv
        sim.phenotype
In [5]:
        ;awk '{print $1}' PedAll.txt | sort -b > all.ID
       ;awk '{print $1}' GenNF.txt | sort -b > genotype.ID
In [6]:
In [7]: |;join -v1 all.ID genotype.ID > noGenotype.ID
In [8]: ;awk '{print $1,$2}' Phe.txt > sim.phenotype
```

```
In [9]: | ;awk '{print $1,$3}' PheAll.txt > sim.bv
In [10]: ; awk 'NR >=1 && NR <=8000 {print $1}' PedAll.txt | sort -b > G0.ID
In [11]: ; awk 'NR >=8001 && NR <=16000 {print $1}' PedAll.txt | sort -b > G1.ID
In [12]: ; awk 'NR >=16001 && NR <=24000 {print $1}' PedAll.txt | sort -b > G2.ID
In [13]: |; awk 'NR >=24001 && NR <=32000 {print $1}' PedAll.txt | sort -b > G3.ID
In [14]: ; awk 'NR >=32001 && NR <=40000 {print $1}' PedAll.txt | sort -b > G4.ID
In [15]: ; awk 'NR >=40001 && NR <=48000 {print $1}' PedAll.txt | sort -b > G5.ID
In [16]: ;join GO.ID genotype.ID > GO.Genotype.ID
In [17]: ;join G1.ID genotype.ID > G1.Genotype.ID
In [18]: ;join G2.ID genotype.ID > G2.Genotype.ID
In [19]: |; join G3.ID genotype.ID > G3.Genotype.ID
In [20]: ;join G4.ID genotype.ID > G4.Genotype.ID
In [21]: |;join G5.ID genotype.ID > G5.Genotype.ID
In [22]: ;join -v1 G0.ID genotype.ID > G0.noGenotype.ID
In [23]: ;join -v1 G1.ID genotype.ID > G1.noGenotype.ID
In [24]: ;join -v1 G2.ID genotype.ID > G2.noGenotype.ID
In [25]: |;join -v1 G3.ID genotype.ID > G3.noGenotype.ID
```

```
In [26]: ;join -v1 G4.ID genotype.ID > G4.noGenotype.ID
In [27]: | ;join -v1 G5.ID genotype.ID > G5.noGenotype.ID
In [28]:
         ;wc G0.Genotype.ID;wc G1.Genotype.ID;wc G2.Genotype.ID;wc G3.Genotype.ID;wc G4.Genotype.ID;wc G5.Genotype
          200 200 1200 GO.Genotype.ID
          200 200 1200 G1.Genotype.ID
          200 200 1200 G2.Genotype.ID
          200 200 1200 G3.Genotype.ID
          200 200 1200 G4.Genotype.ID
          8000 8000 48000 G5.Genotype.ID
         ;wc G0.noGenotype.ID;wc G1.noGenotype.ID;wc G2.noGenotype.ID;wc G3.noGenotype.ID;wc G4.noGenotype.ID;wc G
In [29]:
          7800 7800 46800 GO.noGenotype.ID
          7800 7800 46800 Gl.noGenotype.ID
          7800 7800 46800 G2.noGenotype.ID
          7800 7800 46800 G3.noGenotype.ID
          7800 7800 46800 G4.noGenotype.ID
         0 0 0 G5.noGenotype.ID
In [30]:
         ped,A Mats,numSSBayes = calc Ai("PedAll.txt", "genotype.ID", calculateInbreeding=false)
         nothing
         df
                = read_genotypes("GenNF.txt",numSSBayes)
                                                                                 # without centering
         M_Mats = make_MMats(df,A_Mats,ped);
         y Vecs = make yVecs("sim.phenotype",ped,numSSBayes);
         Z_Mats = make_ZMats(ped,y_Vecs,numSSBayes)
         X_Mats, W_Mats = make_XWMats(Z_Mats,M_Mats,numSSBayes)
                                                                                 # no J
         nothing
```

```
In [31]: | vRes
                = 0.865
                 = 0.865
         vG
         nIter = 50000
         @time aHat1,alphaHat,betaHat,epsiHat =
         ssGibbs(M Mats,y Vecs, Z Mats, X Mats, W Mats, A Mats, numSSBayes, vRes, vG, nIter, outFreq=5000);
         nothing
         This is iteration 5000
         This is iteration 10000
         This is iteration 15000
         This is iteration 20000
         This is iteration 25000
         This is iteration 30000
         This is iteration 35000
         This is iteration 40000
         This is iteration 45000
         This is iteration 50000
         7083.018845 seconds (23.86 G allocations: 737.108 GB, 2.70% gc time)
In [32]:
         betaHat
Out[32]: 1-element Array{Float64,1}:
          31.323
In [33]:
        using DataFrames
In [34]: df = readtable("sim.bv", eltypes =[UTF8String, Float64], separator = ' ',header=false)
         a = Array(Float64, numSSBayes.num ped)
         for (i,ID) in enumerate(df[:,1])
             j = ped.idMap[ID].seqID
             a[j] = df[i,2]
         end
```

```
In [35]: IDs = readtable("all.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor1 = cor(a[posAi],aHat1[posAi])[1,1]
         reg1 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - all.ID : correlation = %6.3f\n", cor1 ) # with epsilon
         @printf("SSBRJC from Gibbs - all.ID : regression of TBV on GEBV = %6.3f\n", req1)
         JCAll = cor1
         SSBRJC from Gibbs - all.ID : correlation = 0.920
         SSBRJC from Gibbs - all.ID : regression of TBV on GEBV = 0.988
Out[35]: 0.9203127838231278
In [36]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[36]: 1.824375949923992
In [37]: IDs = readtable("genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor2 = cor(a[posAi],aHat1[posAi])[1,1]
         reg2 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - genotype.ID : correlation = %6.3f\n", cor2 ) # with epsilon
         @printf("SSBRJC from Gibbs - genotype.ID : regression of TBV on GEBV = %6.3f\n", reg2)
         JCAll = cor2
         SSBRJC from Gibbs - genotype.ID : correlation = 0.920
         SSBRJC from Gibbs - genotype.ID : regression of TBV on GEBV = 1.044
Out[37]: 0.9202621084356952
         GEBV = aHat1[posAi]
In [38]:
         mean (GEBV)
Out[38]: 3.2893753784521444
```

```
In [39]: IDs = readtable("noGenotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor3 = cor(a[posAi],aHat1[posAi])[1,1]
         reg3 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - noGenotype.ID : correlation = %6.3f\n", cor3 ) # with epsilon
         @printf("SSBRJC from Gibbs - noGenotype.ID : regression of TBV on GEBV = %6.3f\n", req3)
         JCAll = cor3
         SSBRJC from Gibbs - noGenotype.ID : correlation = 0.886
         SSBRJC from Gibbs - noGenotype.ID : regression of TBV on GEBV = 0.973
Out[39]: 0.8862083703186387
In [40]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[40]: 1.4862991587251875
In [41]: IDs = readtable("G0.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor4 = cor(a[posAi],aHat1[posAi])[1,1]
         reg4 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G0.ID : correlation = %6.3f\n", cor4 ) # with epsilon
         @printf("SSBRJC from Gibbs - G0.ID : regression of TBV on GEBV = %6.3f\n", req4)
         JCAll = cor4
         SSBRJC from Gibbs - G0.ID : correlation = 0.735
         SSBRJC from Gibbs - G0.ID: regression of TBV on GEBV = 0.928
Out[41]: 0.7351203779264184
In [42]: GEBV = aHat1[posAi]
         G0GEBV=mean(GEBV)
Out[42]: 0.14596087957972595
```

```
In [43]: IDs = readtable("G1.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor4 = cor(a[posAi],aHat1[posAi])[1,1]
         reg4 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G1.ID : correlation = %6.3f\n", cor4 ) # with epsilon
         @printf("SSBRJC from Gibbs - G1.ID : regression of TBV on GEBV = %6.3f\n", req4)
         JCAll = cor4
         SSBRJC from Gibbs - G1.ID : correlation = 0.776
         SSBRJC from Gibbs - G1.ID: regression of TBV on GEBV = 1.003
Out[43]: 0.7764219959095869
In [44]: GEBV = aHat1[posAi]
         G1GEBV=mean(GEBV)
Out[44]: 0.9305624511889384
In [45]: IDs = readtable("G2.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor5 = cor(a[posAi],aHat1[posAi])[1,1]
         reg5 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G2.ID : correlation = %6.3f\n", cor5 ) # with epsilon
         @printf("SSBRJC from Gibbs - G2.ID : regression of TBV on GEBV = %6.3f\n", reg5)
         JCAll = cor5
         SSBRJC from Gibbs - G2.ID : correlation = 0.768
         SSBRJC from Gibbs - G2.ID : regression of TBV on GEBV = 0.997
Out[45]: 0.7678455943838043
In [46]: GEBV = aHat1[posAi]
         G2GEBV=mean(GEBV)
Out[46]: 1.5809909200812964
```

```
In [47]: IDs = readtable("G3.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor6 = cor(a[posAi],aHat1[posAi])[1,1]
         reg6 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G3.ID : correlation = %6.3f\n", cor6 ) # with epsilon
         @printf("SSBRJC from Gibbs - G3.ID : regression of TBV on GEBV = %6.3f\n", reg6)
         JCAll = cor6
         SSBRJC from Gibbs - G3.ID : correlation = 0.768
         SSBRJC from Gibbs - G3.ID: regression of TBV on GEBV = 1.007
Out[47]: 0.7678587278183745
In [48]: GEBV = aHat1[posAi]
         G3GEBV=mean(GEBV)
Out[48]: 2.1699369771983084
In [49]: IDs = readtable("G4.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor7 = cor(a[posAi],aHat1[posAi])[1,1]
         req7 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G4.ID : correlation = %6.3f\n", cor7 ) # with epsilon
         @printf("SSBRJC from Gibbs - G4.ID : regression of TBV on GEBV = %6.3f\n", req7)
         JCAll = cor7
         SSBRJC from Gibbs - G4.ID : correlation = 0.769
         SSBRJC from Gibbs - G4.ID : regression of TBV on GEBV = 1.027
Out[49]: 0.7690288587158385
In [50]: GEBV = aHat1[posAi]
         G4GEBV=mean(GEBV)
Out[50]: 2.7707065941373465
```

```
In [51]: IDs = readtable("G5.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor8 = cor(a[posAi],aHat1[posAi])[1,1]
         reg8 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G5.ID : correlation = %6.3f\n", cor8 ) # with epsilon
         @printf("SSBRJC from Gibbs - G5.ID : regression of TBV on GEBV = %6.3f\n", reg8)
         JCAll = cor8
         SSBRJC from Gibbs - G5.ID : correlation = 0.906
         SSBRJC from Gibbs - G5.ID: regression of TBV on GEBV = 1.032
Out[51]: 0.9059704718182444
In [52]: GEBV = aHat1[posAi]
         G5GEBV=mean(GEBV)
Out[52]: 3.3480978773583363
In [53]: IDs = readtable("G0.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor9 = cor(a[posAi],aHat1[posAi])[1,1]
         \#TBV = a[posAi]
         #GEBV = aHat1[posAi]
         \#reg = linreg(X,Y)
         \#reg = linreg(GEBV, TBV)[2,1]
         req9 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G0.Genotype.ID : correlation = %6.3f\n", cor9 ) # with epsilon
         @printf("SSBRJC from Gibbs - G0.Genotype.ID : regression of TBV on GEBV = %6.3f\n", reg9)
         JCAll = cor9
         SSBRJC from Gibbs - G0.Genotype.ID : correlation = 0.927
         SSBRJC from Gibbs - G0.Genotype.ID: regression of TBV on GEBV = 1.009
Out[53]: 0.9268286090553739
         GEBV = aHat1[posAi]
In [54]:
         mean (GEBV)
Out[54]: 1.7796062650279225
```

```
In [55]: IDs = readtable("G1.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor9 = cor(a[posAi],aHat1[posAi])[1,1]
         \#TBV = a[posAi]
         #GEBV = aHat1[posAi]
         \#reg = linreg(X,Y)
         #reg = linreg(GEBV, TBV)[2,1]
         req9 = linreq(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G1.Genotype.ID : correlation = %6.3f\n", cor9 ) # with epsilon
         @printf("SSBRJC from Gibbs - G1.Genotype.ID : regression of TBV on GEBV = %6.3f\n", reg9)
         JCAll = cor9
         SSBRJC from Gibbs - G1.Genotype.ID : correlation = 0.930
         SSBRJC from Gibbs - G1.Genotype.ID : regression of TBV on GEBV = 0.994
Out[55]: 0.9304342338612851
         GEBV = aHat1[posAi]
In [56]:
         mean(GEBV)
Out[56]: 2.277972970049029
In [57]: IDs = readtable("G2.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor10 = cor(a[posAi],aHat1[posAi])[1,1]
         req10 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G2.Genotype.ID : correlation = %6.3f\n", cor10 ) # with epsilon
         @printf("SSBRJC from Gibbs - G2.Genotype.ID : regression of TBV on GEBV = %6.3f\n", req10)
         JCAll = cor10
         SSBRJC from Gibbs - G2.Genotype.ID : correlation = 0.933
         SSBRJC from Gibbs - G2.Genotype.ID : regression of TBV on GEBV = 1.012
Out[57]: 0.933312436286517
         GEBV = aHat1[posAi]
In [58]:
         mean (GEBV)
Out[58]: 2.7913136856163776
```

```
In [59]: IDs = readtable("G3.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor11 = cor(a[posAi],aHat1[posAi])[1,1]
         reg11 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G3.Genotype.ID : correlation = %6.3f\n", cor11 ) # with epsilon
         @printf("SSBRJC from Gibbs - G3.Genotype.ID : regression of TBV on GEBV = %6.3f\n", req11)
         JCAll = cor11
         SSBRJC from Gibbs - G3.Genotype.ID : correlation = 0.918
         SSBRJC from Gibbs - G3.Genotype.ID : regression of TBV on GEBV = 1.027
Out[59]: 0.9177517404220936
In [60]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[60]: 3.3625312087596226
In [61]: IDs = readtable("G4.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor12 = cor(a[posAi],aHat1[posAi])[1,1]
         reg12 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G4.Genotype.ID : correlation = %6.3f\n", cor12 ) # with epsilon
         @printf("SSBRJC from Gibbs - G4.Genotype.ID : regression of TBV on GEBV = %6.3f\n", req12)
         JCAll = cor12
         SSBRJC from Gibbs - G4.Genotype.ID : correlation = 0.878
         SSBRJC from Gibbs - G4.Genotype.ID : regression of TBV on GEBV = 0.942
Out[61]: 0.8779792467935449
         GEBV = aHat1[posAi]
In [62]:
         mean (GEBV)
Out[62]: 3.886552806560079
```

```
In [63]: IDs = readtable("G5.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor13 = cor(a[posAi],aHat1[posAi])[1,1]
         req13 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G5.Genotype.ID : correlation = %6.3f\n", cor13 ) # with epsilon
         @printf("SSBRJC from Gibbs - G5.Genotype.ID : regression of TBV on GEBV = %6.3f\n", req13)
         JCAll = cor13
         SSBRJC from Gibbs - G5.Genotype.ID : correlation = 0.906
         SSBRJC from Gibbs - G5.Genotype.ID : regression of TBV on GEBV = 1.032
Out[63]: 0.9059704718182444
In [64]: writedlm("Correlation.G5.G.N.txt",cor13)
In [65]: writedlm("Regression.G5.G.N.txt",reg13)
In [66]: TBVG5Gall = a[posAi]
         TBVG5G=mean(TBVG5Gall)
Out[66]: 34.67961825
In [67]: GEBVG5Gall = aHat1[posAi]
         GEBVG5G=mean(GEBVG5Gall)
Out[67]: 3.3480978773583363
In [68]: IDs = readtable("G0.noGenotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor14 = cor(a[posAi],aHat1[posAi])[1,1]
         reg14 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G0.noGenotype.ID : correlation = %6.3f\n", cor14 ) # with epsilon
         @printf("SSBRJC from Gibbs - G0.noGenotype.ID : regression of TBV on GEBV = %6.3f\n", req14)
         JCAll = cor14
         SSBRJC from Gibbs - G0.noGenotype.ID : correlation = 0.714
         SSBRJC from Gibbs - G0.noGenotype.ID: regression of TBV on GEBV = 0.941
Out[68]: 0.7144778389615832
```

```
In [69]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[69]: 0.10407253636310551
In [70]: IDs = readtable("G1.noGenotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor14 = cor(a[posAi],aHat1[posAi])[1,1]
         reg14 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G1.noGenotype.ID : correlation = %6.3f\n", cor14 ) # with epsilon
         @printf("SSBRJC from Gibbs - G1.noGenotype.ID : regression of TBV on GEBV = %6.3f\n", reg14)
         JCAll = cor14
         SSBRJC from Gibbs - G1.noGenotype.ID : correlation = 0.759
         SSBRJC from Gibbs - G1.noGenotype.ID: regression of TBV on GEBV = 1.011
Out[70]: 0.7594601246340887
In [71]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[71]: 0.8960134635258591
In [72]: IDs = readtable("G2.noGenotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor15 = cor(a[posAi],aHat1[posAi])[1,1]
         req15 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G2.noGenotype.ID : correlation = %6.3f\n", cor15 ) # with epsilon
         @printf("SSBRJC from Gibbs - G2.noGenotype.ID : regression of TBV on GEBV = %6.3f\n", reg15)
         JCAll = cor15
         SSBRJC from Gibbs - G2.noGenotype.ID : correlation = 0.750
         SSBRJC from Gibbs - G2.noGenotype.ID: regression of TBV on GEBV = 0.996
Out[72]: 0.7498234181988137
In [73]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[73]: 1.5499570030162941
```

```
In [74]: IDs = readtable("G3.noGenotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor16 = cor(a[posAi],aHat1[posAi])[1,1]
         reg16 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G3.noGenotype.ID : correlation = %6.3f\n", cor16 ) # with epsilon
         @printf("SSBRJC from Gibbs - G3.noGenotype.ID : regression of TBV on GEBV = %6.3f\n", reg16)
         JCAll = cor16
         SSBRJC from Gibbs - G3.noGenotype.ID : correlation = 0.750
         SSBRJC from Gibbs - G3.noGenotype.ID : regression of TBV on GEBV = 1.004
Out[74]: 0.7499372920058434
In [75]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[75]: 2.139357637927506
In [76]: IDs = readtable("G4.noGenotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor17 = cor(a[posAi],aHat1[posAi])[1,1]
         req17 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G4.noGenotype.ID : correlation = %6.3f\n", cor17 ) # with epsilon
         @printf("SSBRJC from Gibbs - G4.noGenotype.ID : regression of TBV on GEBV = %6.3f\n", req17)
         JCAll = cor17
         SSBRJC from Gibbs - G4.noGenotype.ID : correlation = 0.752
         SSBRJC from Gibbs - G4.noGenotype.ID : regression of TBV on GEBV = 1.024
Out[76]: 0.7522391730259095
In [77]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[77]: 2.7420951527931736
In [78]: numSSBayes
Out[78]: SSBR.NumSSBayes(54920,45920,9000,40000,39000,1000,2000)
```