```
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/0.1/G/4
    /home/nicole/Jupyter/JG3/Data/0.1/G/4
```

In [4]: ;ls Correlation.G5.G.C.txt Correlation.G5.G.J.txt Correlation.G5.G.JC.txt Correlation.G5.G.PBLUP.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.C.txt

Regression.G5.G.J.txt

Regression.G5.G.JC.txt

Regression.G5.G.PBLUP.txt

all.ID

alphaEstimates

genotype.ID

meanOfSNPGAll

meanOfSNPGG0

meanOfSNPGG1

meanOfSNPGG2

meanOfSNPGG3

meanOfSNPGG4

meanOfSNPGG5

noGenotype.ID

sim.bv
sim.phenotype

```
In [5]: ;awk '{print $1}' PedAll.txt | sort -b > all.ID
 In [6]: ;awk '{print $1}' GenNF.txt | sort -b > genotype.ID
 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
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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
        ; join -v1 G2.ID genotype.ID > G2.noGenotype.ID
In [24]:
In [25]:
         ; join -v1 G3.ID genotype.ID > G3.noGenotype.ID
         ;join -v1 G4.ID genotype.ID > G4.noGenotype.ID
In [26]:
         ; join -v1 G5.ID genotype.ID > G5.noGenotype.ID
In [27]:
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
In [29]:
         ;wc G0.noGenotype.ID;wc G1.noGenotype.ID;wc G2.noGenotype.ID;wc G3.noGenotype.ID;wc G4.noGenotype.ID;wc G
          7800 7800 46800 GO.noGenotype.ID
          7800
                7800 46800 G1.noGenotype.ID
          7800 7800 46800 G2.noGenotype.ID
          7800
                7800 46800 G3.noGenotype.ID
               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)
         M Mats = make MMats(df,A Mats,ped);
                                                                                  # without centering
         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]:
                = 0.871
         vG
         vRes
                = 7.839
         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
         2626.156433 seconds (23.04 G allocations: 723.592 GB, 7.56% gc time)
        betaHat
In [32]:
Out[32]: 1-element Array{Float64,1}:
          9.16032
         using DataFrames
In [33]:
```

```
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", reg1)
         JCAll = cor1
         SSBRJC from Gibbs - all.ID : correlation = 0.762
         SSBRJC from Gibbs - all.ID : regression of TBV on GEBV = 0.988
Out[35]: 0.761675590037221
In [36]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[36]: 1.2961742764659434
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 = 0.938
         SSBRJC from Gibbs - genotype.ID : regression of TBV on GEBV = 1.244
Out[37]: 0.9379937474244471
In [38]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[38]: 2.074963545954978
```

```
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", reg3)
         JCAll = cor3
         SSBRJC from Gibbs - noGenotype.ID : correlation = 0.689
         SSBRJC from Gibbs - noGenotype.ID : regression of TBV on GEBV = 0.957
Out[39]: 0.6889374456865207
In [40]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[40]: 1.1164536758146275
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 = 0.548
         SSBRJC from Gibbs - G0.ID: regression of TBV on GEBV = 1.326
Out[41]: 0.5477065748081605
In [42]: GEBV = aHat1[posAi]
         GOGEBV=mean(GEBV)
Out[42]: 0.12037155730869857
```

```
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 = 0.538
         SSBRJC from Gibbs - G1.ID: regression of TBV on GEBV = 1.107
Out[43]: 0.537586398737016
In [44]: GEBV = aHat1[posAi]
         G1GEBV=mean(GEBV)
Out[44]: 0.9014693989112442
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 = 0.563
         SSBRJC from Gibbs - G2.ID : regression of TBV on GEBV = 1.099
Out[45]: 0.563237097234715
In [46]: GEBV = aHat1[posAi]
         G2GEBV=mean(GEBV)
Out[46]: 1.2699032450624859
```

```
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 = 0.598
         SSBRJC from Gibbs - G3.ID : regression of TBV on GEBV = 1.116
Out[47]: 0.5979168159341982
In [48]: GEBV = aHat1[posAi]
         G3GEBV=mean(GEBV)
Out[48]: 1.5639354551253972
In [49]: IDs = readtable("G4.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor7 = cor(a[posAi],aHat1[posAi])[1,1]
         reg7 = 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 = 0.676
         SSBRJC from Gibbs - G4.ID : regression of TBV on GEBV = 1.198
Out[49]: 0.676433149574001
In [50]: GEBV = aHat1[posAi]
         G4GEBV=mean(GEBV)
Out[50]: 1.8264364608882062
```

```
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 = 0.937
         SSBRJC from Gibbs - G5.ID: regression of TBV on GEBV = 1.249
Out[51]: 0.937013383941551
In [52]: GEBV = aHat1[posAi]
         G5GEBV=mean(GEBV)
Out[52]: 2.0949295414996283
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]
         reg9 = 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 = 0.926
         SSBRJC from Gibbs - G0.Genotype.ID : regression of TBV on GEBV = 1.110
Out[53]: 0.9257633059913329
In [54]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[54]: 1.7085693172059597
```

```
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]
         reg9 = linreg(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 = 0.939
         SSBRJC from Gibbs - G1.Genotype.ID : regression of TBV on GEBV = 1.176
Out[55]: 0.939197310730989
         GEBV = aHat1[posAi]
In [56]:
         mean(GEBV)
Out[56]: 1.640197046076589
In [57]: IDs = readtable("G2.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor10 = cor(a[posAi],aHat1[posAi])[1,1]
         reg10 = 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 = 0.930
         SSBRJC from Gibbs - G2.Genotype.ID : regression of TBV on GEBV = 1.198
Out[57]: 0.9301039358389275
In [58]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[58]: 1.8770884470306763
```

```
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 = 0.942
         SSBRJC from Gibbs - G3.Genotype.ID : regression of TBV on GEBV = 1.279
Out[59]: 0.9421895485518019
In [60]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[60]: 2.0657772682849282
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", reg12)
         JCAll = cor12
         SSBRJC from Gibbs - G4.Genotype.ID : correlation = 0.946
         SSBRJC from Gibbs - G4.Genotype.ID : regression of TBV on GEBV = 1.274
Out[61]: 0.9460744464396404
In [62]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[62]: 2.2845458293907233
```

```
In [63]: IDs = readtable("G5.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor13 = cor(a[posAi],aHat1[posAi])[1,1]
         reg13 = 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 = 0.937
         SSBRJC from Gibbs - G5.Genotype.ID : regression of TBV on GEBV = 1.249
Out[63]: 0.937013383941551
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]: 11.194825999999999
In [67]: GEBVG5Gall = aHat1[posAi]
         GEBVG5G=mean(GEBVG5Gall)
Out[67]: 2.0949295414996283
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", reg14)
         JCAll = cor14
         SSBRJC from Gibbs - G0.noGenotype.ID : correlation = 0.541
         SSBRJC from Gibbs - G0.noGenotype.ID : regression of TBV on GEBV = 1.778
Out[68]: 0.5412621757387301
```

```
In [69]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[69]: 0.07964853782415339
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 = 0.523
         SSBRJC from Gibbs - G1.noGenotype.ID : regression of TBV on GEBV = 1.137
Out[70]: 0.5225519379093787
In [71]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[71]: 0.8825276643685429
In [72]: IDs = readtable("G2.noGenotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor15 = cor(a[posAi],aHat1[posAi])[1,1]
         reg15 = 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 = 0.547
         SSBRJC from Gibbs - G2.noGenotype.ID: regression of TBV on GEBV = 1.106
Out[72]: 0.5471768251430771
In [73]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[73]: 1.2543343937299682
```

```
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 = 0.581
         SSBRJC from Gibbs - G3.noGenotype.ID : regression of TBV on GEBV = 1.111
Out[74]: 0.5814831636043375
In [75]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[75]: 1.5510677163264348
In [76]: IDs = readtable("G4.noGenotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor17 = cor(a[posAi],aHat1[posAi])[1,1]
         reg17 = 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", reg17)
         JCAll = cor17
         SSBRJC from Gibbs - G4.noGenotype.ID : correlation = 0.664
         SSBRJC from Gibbs - G4.noGenotype.ID: regression of TBV on GEBV = 1.195
Out[76]: 0.6638238327808826
In [77]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[77]: 1.8146900668240395
In [78]: numSSBayes
Out[78]: SSBR.NumSSBayes(54893,45893,9000,40000,39000,1000,200)
```