```
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/6
         /home/nicole/Jupyter/JG3/Data/0.1/G/6
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
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
         ; awk 'NR >=1 && NR <=8000 {print $1}' PedAll.txt | sort -b > G0.ID
In [10]:
```

```
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
         ; awk 'NR >=24001 && NR <=32000 {print $1}' PedAll.txt | sort -b > G3.ID
In [13]:
         ; awk 'NR >=32001 && NR <=40000 {print $1}' PedAll.txt | sort -b > G4.ID
         ; awk 'NR >=40001 && NR <=48000 {print $1}' PedAll.txt | sort -b > G5.ID
In [16]:
         ; join G0.ID genotype.ID > G0.Genotype.ID
         ; join G1.ID genotype.ID > G1.Genotype.ID
In [17]:
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
         ;join -v1 G0.ID genotype.ID > G0.noGenotype.ID
In [22]:
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 (
               200 1200 GO.Genotype.ID
               200 1200 G1.Genotype.ID
          200
               200 1200 G2.Genotype.ID
               200 1200 G3.Genotype.ID
          200
          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
                7800 46800 G0.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",calculateInbreedia
         nothing
         df
                = read_genotypes("GenNF.txt",numSSBayes)
         M Mats = make MMats(df, A Mats, ped, center=true);
                                                                                  # with
         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
         nothing
                = 0.576
In [31]: vRes
                = 5.191
         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,
         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
         2508.429500 seconds (23.06 G allocations: 724.005 GB, 7.67% gc time)
In [32]: betaHat
Out[32]: 1-element Array{Float64,1}:
          11.2718
In [33]: using DataFrames
In [34]: | df = readtable("sim.bv", eltypes =[UTF8String, Float64], separator = ' ',head
         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 (
         @printf("SSBRJC from Gibbs - all.ID : regression of TBV on GEBV = %6.3f\n",
         JCAll = cor1
         SSBRJC from Gibbs - all.ID : correlation = 0.546
         SSBRJC from Gibbs - all.ID : regression of TBV on GEBV = 0.263
Out[35]: 0.5463080705676527
In [36]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[36]: -0.3874959660026866
In [37]: IDs = readtable("genotype.ID", eltypes =[UTF8String], separator = ' ',header:
         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 ) # |
         @printf("SSBRJC from Gibbs - genotype.ID : regression of TBV on GEBV = %6.3:
         JCAll = cor2
         SSBRJC from Gibbs - genotype.ID : correlation = 0.880
         SSBRJC from Gibbs - genotype.ID : regression of TBV on GEBV = 0.795
Out[37]: 0.8802843791260776
In [38]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[38]: 1.331721103546508
In [39]: IDs = readtable("noGenotype.ID", eltypes =[UTF8String], separator = ' ',heade
         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 );
         @printf("SSBRJC from Gibbs - noGenotype.ID : regression of TBV on GEBV = %6
         JCAll = cor3
         SSBRJC from Gibbs - noGenotype.ID : correlation = 0.438
         SSBRJC from Gibbs - noGenotype.ID : regression of TBV on GEBV = 0.209
Out[39]: 0.43754172095614885
In [40]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[40]: -0.7842383666678855
```

```
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 ei
         @printf("SSBRJC from Gibbs - G0.ID : regression of TBV on GEBV = %6.3f\n", re
         JCAll = cor4
         SSBRJC from Gibbs - G0.ID : correlation = 0.621
         SSBRJC from Gibbs - G0.ID: regression of TBV on GEBV = 0.433
Out[41]: 0.6214620723358467
In [42]: GEBV = aHat1[posAi]
         G1GEBV=mean(GEBV)
Out[42]: -1.4778812687680833
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 e;
         @printf("SSBRJC from Gibbs - G1.ID : regression of TBV on GEBV = %6.3f\n", re
         JCAll = cor4
         SSBRJC from Gibbs - G1.ID : correlation = 0.359
         SSBRJC from Gibbs - G1.ID: regression of TBV on GEBV = 0.143
Out[43]: 0.35929207332153495
In [44]: GEBV = aHat1[posAi]
         G1GEBV=mean(GEBV)
Out[44]: -0.9680681139121986
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 ep
         @printf("SSBRJC from Gibbs - G2.ID : regression of TBV on GEBV = %6.3f\n", re
         JCAll = cor5
         SSBRJC from Gibbs - G2.ID : correlation = 0.355
         SSBRJC from Gibbs - G2.ID : regression of TBV on GEBV = 0.140
Out[45]: 0.35477279963294844
In [46]: GEBV = aHat1[posAi]
         G2GEBV=mean(GEBV)
Out[46]: -0.6931039166368335
```

```
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 ei
         @printf("SSBRJC from Gibbs - G3.ID : regression of TBV on GEBV = %6.3f\n", re
         JCAll = cor6
         SSBRJC from Gibbs - G3.ID : correlation = 0.356
         SSBRJC from Gibbs - G3.ID: regression of TBV on GEBV = 0.142
Out[47]: 0.3563946326652258
In [48]: GEBV = aHat1[posAi]
         G3GEBV=mean(GEBV)
Out[48]: -0.4420616361573864
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 e;
         @printf("SSBRJC from Gibbs - G4.ID : regression of TBV on GEBV = %6.3f\n", re
         JCAll = cor7
         SSBRJC from Gibbs - G4.ID : correlation = 0.355
         SSBRJC from Gibbs - G4.ID: regression of TBV on GEBV = 0.147
Out[49]: 0.35459655937259243
In [50]: GEBV = aHat1[posAi]
         G4GEBV=mean(GEBV)
Out[50]: -0.10506716980453498
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 ep
         @printf("SSBRJC from Gibbs - G5.ID : regression of TBV on GEBV = %6.3f\n", re
         JCAll = cor8
         SSBRJC from Gibbs - G5.ID : correlation = 0.879
         SSBRJC from Gibbs - G5.ID : regression of TBV on GEBV = 0.795
Out[51]: 0.8788523433894616
In [52]: GEBV = aHat1[posAi]
         G5GEBV=mean(GEBV)
Out[52]: 1.361206309262917
```

```
In [53]: IDs = readtable("G0.Genotype.ID", eltypes =[UTF8String], separator = ' ',heac
         posAi = getPos(ped,IDs)
         cor9 = cor(a[posAi],aHat1[posAi])[1,1]
         reg9 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G0.Genotype.ID : correlation = %6.3f\n", cor9 )
         @printf("SSBRJC from Gibbs - G0.Genotype.ID : regression of TBV on GEBV = %6
         JCAll = cor9
         SSBRJC from Gibbs - G0.Genotype.ID : correlation = 0.854
         SSBRJC from Gibbs - G0.Genotype.ID : regression of TBV on GEBV = 0.783
Out[53]: 0.8537986706667252
In [54]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[54]: 0.6630898950720062
In [55]: IDs = readtable("G1.Genotype.ID", eltypes =[UTF8String], separator = ' ', head
         posAi = getPos(ped,IDs)
         cor9 = cor(a[posAi],aHat1[posAi])[1,1]
         reg9 = linreg(aHat1[posAi], a[posAi])[2,1]
         @printf("SSBRJC from Gibbs - G1.Genotype.ID : correlation = %6.3f\n", cor9 )
         @printf("SSBRJC from Gibbs - G1.Genotype.ID : regression of TBV on GEBV = %6
         JCAll = cor9
         SSBRJC from Gibbs - G1.Genotype.ID : correlation = 0.833
         SSBRJC from Gibbs - G1.Genotype.ID: regression of TBV on GEBV = 0.805
Out[55]: 0.8331376166642711
In [56]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[56]: 0.7743266463680412
In [57]: IDs = readtable("G2.Genotype.ID", eltypes =[UTF8String], separator = ' ', head
         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
         @printf("SSBRJC from Gibbs - G2.Genotype.ID : regression of TBV on GEBV = %6
         JCAll = cor10
         SSBRJC from Gibbs - G2.Genotype.ID : correlation =
         SSBRJC from Gibbs - G2.Genotype.ID : regression of TBV on GEBV = 0.826
Out[57]: 0.8828775562286081
In [58]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[58]: 1.0433095794324148
```

```
In [59]: IDs = readtable("G3.Genotype.ID", eltypes =[UTF8String], separator = ' ',heac
         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
         @printf("SSBRJC from Gibbs - G3.Genotype.ID : regression of TBV on GEBV = %6
         JCAll = cor11
         SSBRJC from Gibbs - G3.Genotype.ID : correlation = 0.886
         SSBRJC from Gibbs - G3.Genotype.ID: regression of TBV on GEBV = 0.820
Out[59]: 0.8862213810337987
In [60]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[60]: 1.290938202141833
In [61]: IDs = readtable("G4.Genotype.ID", eltypes =[UTF8String], separator = ' ', head
         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
         @printf("SSBRJC from Gibbs - G4.Genotype.ID : regression of TBV on GEBV = %6
         JCAll = cor12
         SSBRJC from Gibbs - G4.Genotype.ID : correlation = 0.876
         SSBRJC from Gibbs - G4.Genotype.ID : regression of TBV on GEBV = 0.845
Out[61]: 0.8759296639935704
In [62]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[62]: 1.707532966061882
In [63]: IDs = readtable("G5.Genotype.ID", eltypes =[UTF8String], separator = ' ', head
         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
         @printf("SSBRJC from Gibbs - G5.Genotype.ID : regression of TBV on GEBV = %6
         JCAll = cor13
         SSBRJC from Gibbs - G5.Genotype.ID : correlation =
         SSBRJC from Gibbs - G5.Genotype.ID: regression of TBV on GEBV = 0.795
Out[63]: 0.8788523433894616
In [64]: | writedlm("Correlation.G5.G.C.txt",cor13)
In [65]: writedlm("Regression.G5.G.C.txt",reg13)
```

```
In [66]: GEBVG5Gall = aHat1[posAi]
         GEBVG5G=mean(GEBVG5Gall)
Out[66]: 1.361206309262917
In [67]: IDs = readtable("G0.noGenotype.ID", eltypes =[UTF8String], separator = ' ',he
         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", corl
         @printf("SSBRJC from Gibbs - G0.noGenotype.ID : regression of TBV on GEBV = '
         JCAll = cor14
         SSBRJC from Gibbs - G0.noGenotype.ID : correlation = 0.597
         SSBRJC from Gibbs - G0.noGenotype.ID : regression of TBV on GEBV = 0.427
Out[67]: 0.5968095876904267
In [68]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[68]: -1.532777965276804
In [69]: IDs = readtable("G1.noGenotype.ID", eltypes =[UTF8String], separator = ' ',he
         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", corl
         @printf("SSBRJC from Gibbs - G1.noGenotype.ID : regression of TBV on GEBV = '
         JCAll = cor14
         SSBRJC from Gibbs - G1.noGenotype.ID : correlation = 0.346
         SSBRJC from Gibbs - G1.noGenotype.ID : regression of TBV on GEBV = 0.138
Out[69]: 0.34603313216081005
In [70]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[70]: -1.0127449026373327
In [71]: IDs = readtable("G2.noGenotype.ID", eltypes =[UTF8String], separator = ' ',he
         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", cor1!
         @printf("SSBRJC from Gibbs - G2.noGenotype.ID : regression of TBV on GEBV = '
         JCAll = cor15
         SSBRJC from Gibbs - G2.noGenotype.ID : correlation = 0.341
         SSBRJC from Gibbs - G2.noGenotype.ID: regression of TBV on GEBV = 0.134
Out[71]: 0.34106147559708677
```

```
In [72]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[72]: -0.737627339612968
In [73]: IDs = readtable("G3.noGenotype.ID", eltypes =[UTF8String], separator = ' ',he
         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", cor1
         @printf("SSBRJC from Gibbs - G3.noGenotype.ID : regression of TBV on GEBV = '
         JCAll = cor16
         SSBRJC from Gibbs - G3.noGenotype.ID : correlation = 0.343
         SSBRJC from Gibbs - G3.noGenotype.ID: regression of TBV on GEBV = 0.136
Out[73]: 0.3430739016846085
In [74]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[74]: -0.48649752944711006
In [75]: IDs = readtable("G4.noGenotype.ID", eltypes =[UTF8String], separator = ' ',he
         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", cor1
         @printf("SSBRJC from Gibbs - G4.noGenotype.ID : regression of TBV on GEBV = '
         JCAll = cor17
         SSBRJC from Gibbs - G4.noGenotype.ID : correlation = 0.340
         SSBRJC from Gibbs - G4.noGenotype.ID : regression of TBV on GEBV = 0.141
Out[75]: 0.339797634521568
In [76]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[76]: -0.15154409636521235
In [77]: numSSBayes
Out[77]: SSBR.NumSSBayes(54923,45923,9000,40000,39000,1000,200)
```