```
In [1]: include("/home/nicole/Jupyter/SSBRJ/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/1
         /home/nicole/Jupyter/JG3/Data/0.1/G/1
 In [4]: ;ls
         GenNF.txt
         PedAll.txt
         Phe.txt
         PheAll.txt
 In [5]: |;awk '{print $1}' PedAll.txt | sort -b > all.ID
 In [6]: ;awk '{print $1}' GenNF.txt | sort -b > genotype.ID
         ; 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
```

```
;wc G0.Genotype.ID;wc G1.Genotype.ID;wc G2.Genotype.ID;wc G3.Genotype.ID;wc G4.Genotype.ID;wc G5.Genotype
In [28]:
          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 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
         ped,A Mats,numSSBayes = calc Ai("PedAll.txt", "genotype.ID", calculateInbreeding=false)
In [30]:
         nothing
         df
                = read genotypes("GenNF.txt", numSSBayes)
                                                                                 # with centering
         M_Mats = make_MMats(df,A_Mats,ped,center=true);
         y_Vecs = make_yVecs("sim.phenotype",ped,numSSBayes)
         J_Vecs = make_JVecs(numSSBayes,A_Mats)
         Z_Mats = make_ZMats(ped,y_Vecs,numSSBayes)
         X Mats, W Mats = make XWMats(J Vecs, Z Mats, M Mats, numSSBayes)
                                                                                 # with J
         nothing
```

```
In [31]: | vRes
                = 7.831
         vG
                 = 0.870
         nIter = 50000
         @time aHat1,alphaHat,betaHat,epsiHat =
         ssGibbs(M Mats,y Vecs,J 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
         4574.941825 seconds (23.05 G allocations: 723.771 GB, 6.82% gc time)
In [32]:
        betaHat
Out[32]: 2-element Array{Float64,1}:
          13.4164
           3.63842
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 = 0.761
         SSBRJC from Gibbs - all.ID : regression of TBV on GEBV = 0.998
Out[35]: 0.7609941054822827
In [36]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[36]: -2.1335119480735765
In [37]: IDs = readtable("genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor2 = cor(a[posAi],aHat1[posAi])[1,1]
         req2 = 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 = 0.943
         SSBRJC from Gibbs - genotype.ID : regression of TBV on GEBV = 1.202
Out[37]: 0.9432619918402971
         GEBV = aHat1[posAi]
In [38]:
         mean (GEBV)
Out[38]: -1.3864362527326517
```

```
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 = 0.687
         SSBRJC from Gibbs - noGenotype.ID : regression of TBV on GEBV = 0.968
Out[39]: 0.6870078494670717
In [40]: | GEBV = aHat1[posAi]
         mean(GEBV)
Out[40]: -2.30591403161379
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 = 0.538
         SSBRJC from Gibbs - G0.ID: regression of TBV on GEBV = 1.326
Out[41]: 0.5379504106992943
In [42]: GEBV = aHat1[posAi]
         G0GEBV=mean(GEBV)
Out[42]: -3.2708631316265793
```

```
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 = 0.535
         SSBRJC from Gibbs - G1.ID: regression of TBV on GEBV = 1.120
Out[43]: 0.5353632588318308
In [44]: GEBV = aHat1[posAi]
         G1GEBV=mean(GEBV)
Out[44]: -2.51353496718105
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 = 0.585
         SSBRJC from Gibbs - G2.ID : regression of TBV on GEBV = 1.133
Out[45]: 0.5852496004050483
In [46]: GEBV = aHat1[posAi]
         G2GEBV=mean(GEBV)
Out[46]: -2.1540293775447323
```

```
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 = 0.599
         SSBRJC from Gibbs - G3.ID: regression of TBV on GEBV = 1.089
Out[47]: 0.5993090318567967
In [48]: GEBV = aHat1[posAi]
         G3GEBV=mean(GEBV)
Out[48]: -1.874161234249001
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 = 0.676
         SSBRJC from Gibbs - G4.ID : regression of TBV on GEBV = 1.150
Out[49]: 0.6758401793054039
In [50]: GEBV = aHat1[posAi]
         G4GEBV=mean(GEBV)
Out[50]: -1.6191981544923164
```

```
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 = 0.943
         SSBRJC from Gibbs - G5.ID: regression of TBV on GEBV = 1.205
Out[51]: 0.9425909574693728
In [52]: GEBV = aHat1[posAi]
         G5GEBV=mean(GEBV)
Out[52]: -1.3692848233477803
In [53]: IDs = readtable("G0.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor9 = cor(a[posAi],aHat1[posAi])[1,1]
         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 = 0.906
         SSBRJC from Gibbs - G0.Genotype.ID : regression of TBV on GEBV = 1.139
Out[53]: 0.9059338483811672
         GEBV = aHat1[posAi]
In [54]:
         mean (GEBV)
Out[54]: -1.7354424282254073
```

```
In [55]: IDs = readtable("G1.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         posAi = getPos(ped,IDs)
         cor9 = cor(a[posAi],aHat1[posAi])[1,1]
         req9 = 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 = 0.944
         SSBRJC from Gibbs - G1.Genotype.ID : regression of TBV on GEBV = 1.198
Out[55]: 0.9439184640041216
In [56]:
         GEBV = aHat1[posAi]
         mean(GEBV)
Out[56]: -1.7902230544598199
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 = 0.950
         SSBRJC from Gibbs - G2.Genotype.ID : regression of TBV on GEBV = 1.157
Out[57]: 0.949812934619145
         GEBV = aHat1[posAi]
In [58]:
         mean (GEBV)
Out[58]: -1.5827372471469212
```

```
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 = 0.946
         SSBRJC from Gibbs - G3.Genotype.ID: regression of TBV on GEBV = 1.139
Out[59]: 0.9456690264031616
In [60]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[60]: -1.3656648349020275
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 = 0.944
         SSBRJC from Gibbs - G4.Genotype.ID : regression of TBV on GEBV = 1.210
Out[61]: 0.9435793496177692
         GEBV = aHat1[posAi]
In [62]:
         mean (GEBV)
Out[62]: -1.1441708743239491
```

```
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 = 0.943
         SSBRJC from Gibbs - G5.Genotype.ID: regression of TBV on GEBV = 1.205
Out[63]: 0.9425909574693728
In [64]: writedlm("Correlation.G5.G.JC.txt",cor13)
In [65]: writedlm("Regression.G5.G.JC.txt",reg13)
In [66]: TBVG5Gall = a[posAi]
         TBVG5G=mean(TBVG5Gall)
Out[66]: 12.014813624999999
In [67]: GEBVG5Gall = aHat1[posAi]
         GEBVG5G=mean(GEBVG5Gall)
Out[67]: -1.3692848233477803
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 = 0.528
         SSBRJC from Gibbs - G0.noGenotype.ID : regression of TBV on GEBV = 1.736
Out[68]: 0.5276352056389705
```

```
In [69]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[69]: -3.3102328932522505
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 = 0.519
         SSBRJC from Gibbs - G1.noGenotype.ID : regression of TBV on GEBV = 1.148
Out[70]: 0.518908042145604
In [71]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[71]: -2.5320814264815943
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 = 0.569
         SSBRJC from Gibbs - G2.noGenotype.ID : regression of TBV on GEBV = 1.140
Out[72]: 0.5691611846945711
In [73]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[73]: -2.1686778937087783
```

```
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", req16)
         JCAll = cor16
         SSBRJC from Gibbs - G3.noGenotype.ID : correlation = 0.584
         SSBRJC from Gibbs - G3.noGenotype.ID: regression of TBV on GEBV = 1.088
Out[74]: 0.5844515794422067
In [75]: GEBV = aHat1[posAi]
         mean(GEBV)
Out[75]: -1.8871996034630258
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 = 0.664
         SSBRJC from Gibbs - G4.noGenotype.ID: regression of TBV on GEBV = 1.147
Out[76]: 0.6641845419542929
In [77]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[77]: -1.6313783411633
In [78]: numSSBayes
Out[78]: SSBR.NumSSBayes(54905,45905,9000,40000,39000,1000,200)
```

```
In [79]: J1 = sortrows(J_Vecs.J1)
Out[79]: 45905x1 Array{Float64,2}:
          -0.987503
          -0.986037
          -0.986007
          -0.985991
          -0.985903
          -0.985634
          -0.985614
          -0.985579
          -0.985507
          -0.985288
          -0.985284
          -0.985085
          -0.984751
           6.02079e-17
           6.07214e-17
           6.0938e-17
           6.3236e-17
           6.6105e-17
           6.61774e-17
           6.63649e-17
           6.65614e-17
           6.66877e-17
           8.88178e-17
           1.06976e-16
           1.14306e-16
```

```
In [80]: J1[J1 .< 0.0,:]
Out[80]: 43957x1 Array{Float64,2}:
          -0.987503
          -0.986037
          -0.986007
          -0.985991
          -0.985903
          -0.985634
          -0.985614
          -0.985579
          -0.985507
          -0.985288
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          -0.985085
          -0.984751
          -1.08217e-35
          -1.0817e-35
          -7.22085e-36
          -7.22081e-36
          -7.21871e-36
          -7.2166e-36
          -7.21238e-36
          -7.21237e-36
          -7.21237e-36
          -5.41087e-36
          -2.55363e-65
          -5.29562e-67
```