J-SSBRJ-JC-G

```
In [4]: ;ls
        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.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
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]:
         ;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]:
         vG
                = 0.872
                = 7.848
         vRes
         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
         2702.570166 seconds (23.03 G allocations: 723.343 GB, 7.34% gc time)
In [32]: betaHat
Out[32]: 2-element Array{Float64,1}:
          13.637
           2.78264
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", reg1)
         JCAll = cor1
         SSBRJC from Gibbs - all.ID : correlation = 0.766
         SSBRJC from Gibbs - all.ID : regression of TBV on GEBV = 0.954
Out[35]: 0.7662203112592455
In [36]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[36]: -1.6017850322378557
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 = 0.943
         SSBRJC from Gibbs - genotype.ID : regression of TBV on GEBV = 1.072
Out[37]: 0.9431897455260283
In [38]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[38]: -0.8279472169972151
```

```
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 = 0.699
         SSBRJC from Gibbs - noGenotype.ID : regression of TBV on GEBV = 0.948
Out[39]: 0.6987568423064265
In [40]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[40]: -1.7803629896010806
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 = 0.545
         SSBRJC from Gibbs - G0.ID : regression of TBV on GEBV = 1.317
Out[41]: 0.5450265796824896
In [42]: GEBV = aHat1[posAi]
         GOGEBV=mean(GEBV)
Out[42]: -2.7397032119896774
```

```
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 = 0.549
         SSBRJC from Gibbs - G1.ID: regression of TBV on GEBV = 1.038
Out[43]: 0.5490985007056757
In [44]: GEBV = aHat1[posAi]
         G1GEBV=mean(GEBV)
Out[44]: -1.9856854971639846
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 = 0.589
         SSBRJC from Gibbs - G2.ID : regression of TBV on GEBV = 1.031
Out[45]: 0.5891088422274579
In [46]: GEBV = aHat1[posAi]
         G2GEBV=mean(GEBV)
Out[46]: -1.652200787062204
```

```
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 = 0.646
         SSBRJC from Gibbs - G3.ID : regression of TBV on GEBV = 1.080
Out[47]: 0.6459939812754207
In [48]: GEBV = aHat1[posAi]
         G3GEBV=mean(GEBV)
Out[48]: -1.3613670837561076
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 = 0.689
         SSBRJC from Gibbs - G4.ID : regression of TBV on GEBV = 1.031
Out[49]: 0.6886357355874572
In [50]: GEBV = aHat1[posAi]
         G4GEBV=mean(GEBV)
Out[50]: -1.0641748231874661
```

```
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 = 0.942
         SSBRJC from Gibbs - G5.ID: regression of TBV on GEBV = 1.076
Out[51]: 0.9419197934946048
In [52]: GEBV = aHat1[posAi]
         G5GEBV=mean(GEBV)
Out[52]: -0.8075787902676952
In [53]: IDs = readtable("G0.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         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 ) # 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 = 1.020
Out[53]: 0.9261203289559194
In [54]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[54]: -1.217095563733515
```

```
In [55]: IDs = readtable("G1.Genotype.ID", eltypes =[UTF8String], separator = ' ',header=false)
         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 ) # 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 = 0.937
         SSBRJC from Gibbs - G1.Genotype.ID : regression of TBV on GEBV = 1.057
Out[55]: 0.9373556279164482
In [56]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[56]: -1.3134724479058608
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 = 0.955
         SSBRJC from Gibbs - G2.Genotype.ID : regression of TBV on GEBV = 1.103
Out[57]: 0.9545686302416471
In [58]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[58]: -1.067998616084926
```

```
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 = 0.957
         SSBRJC from Gibbs - G3.Genotype.ID : regression of TBV on GEBV = 1.026
Out[59]: 0.9573440202182532
In [60]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[60]: -0.7722227004360134
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 = 1.035
Out[61]: 0.9436173597393124
In [62]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[62]: -0.5836838260065516
```

```
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 = 0.942
         SSBRJC from Gibbs - G5.Genotype.ID : regression of TBV on GEBV = 1.076
Out[63]: 0.9419197934946048
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]: 12.725047124999998
In [67]: GEBVG5Gall = aHat1[posAi]
         GEBVG5G=mean(GEBVG5Gall)
Out[67]: -0.8075787902676952
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 = 0.537
         SSBRJC from Gibbs - G0.noGenotype.ID : regression of TBV on GEBV = 1.732
Out[68]: 0.5367571514843124
```

```
In [69]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[69]: -2.7787444337398353
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 = 0.533
         SSBRJC from Gibbs - G1.noGenotype.ID : regression of TBV on GEBV = 1.055
Out[70]: 0.5326955427910182
In [71]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[71]: -2.002921729196244
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 = 0.570
         SSBRJC from Gibbs - G2.noGenotype.ID: regression of TBV on GEBV = 1.030
Out[72]: 0.5700659097811646
In [73]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[73]: -1.6671803299077754
```

```
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 = 0.633
         SSBRJC from Gibbs - G3.noGenotype.ID: regression of TBV on GEBV = 1.087
Out[74]: 0.6330760955301457
In [75]: GEBV = aHat1[posAi]
         mean (GEBV)
Out[75]: -1.3764733499950843
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 = 0.679
         SSBRJC from Gibbs - G4.noGenotype.ID : regression of TBV on GEBV = 1.031
Out[76]: 0.6793714213232942
In [77]: | GEBV = aHat1[posAi]
         mean (GEBV)
Out[77]: -1.0764951051664642
In [78]: numSSBayes
Out[78]: SSBR.NumSSBayes(54874,45874,9000,40000,39000,1000,200)
```

```
In [79]: J1 = sortrows(J_Vecs.J1)
Out[79]: 45874x1 Array{Float64,2}:
          -0.999675
          -0.987323
          -0.986174
          -0.985395
          -0.985332
          -0.984131
          -0.98413
          -0.982808
          -0.982653
          -0.982593
          -0.982364
          -0.982323
          -0.981637
           6.77188e-17
           7.393e-17
           7.4291e-17
           7.44698e-17
           8.3077e-17
           8.77235e-17
           8.87952e-17
           8.88495e-17
           1.04174e-16
           1.05731e-16
           1.07411e-16
           1.11144e-16
```

```
In [80]: J1[J1 .< 0.0,:]
Out[80]: 43825x1 Array{Float64,2}:
          -0.999675
          -0.987323
          -0.986174
          -0.985395
          -0.985332
          -0.984131
          -0.98413
          -0.982808
          -0.982653
          -0.982593
          -0.982364
          -0.982323
          -0.981637
          -5.41167e-36
          -5.4077e-36
          -7.63874e-65
          -7.60646e-65
          -7.60646e-65
          -3.81937e-65
          -3.80323e-65
          -2.90722e-65
          -2.59338e-65
          -6.99614e-66
          -9.01683e-67
          -1.78007e-67
```