MendelGeneticCounseling ASHG Workshop

last update: September 12 2019, 3:20 pm.

The purpose of this tutorial is to demonstrate how to calculate genetic risks for individuals using their family histories and covariate information.

Commands starting with ";" are Mac shell commands that should work with a PowerShell on Windows. If using a Dos cmd window use "type" rather than "cat".

Installation instructions (not needed during workshop)

MendelGeneticCounseling.jl#ASHG2019 currently supports Julia versions 1.0, 1.1 and 1.2, but it is currently an unregistered package. To install, press] to invoke the package manager mode and install these packages by typing:

```
add https://github.com/OpenMendel/SnpArrays.jl
add https://github.com/OpenMendel/MendelSearch.jl
add https://github.com/OpenMendel/MendelBase.jl
add https://github.com/OpenMendel/MendelGeneticCounseling.jl#ASHG2019
build SpecialFunctions
```

Be sure also to download the BRCA and Cholesterol data from https://github.com/OpenMendel/GeneticCounseling_ASHG2019). For convenience, have your input files and this notebook in the same directory.

NOTE: When finished with this notebook, go to the File tab and first select save and checkpoint to save your results. Then under the file tab, select close and halt to prevent copies of Jupyter notebook from running indefinitely in the background.

Check Julia version:

For reproducibility, check the machine information below. To execute a notebook command, hold downShift Enter within the box. This tutorial and corresponding modules have been checked with Julia versions 1.0, 1.1 and 1.2. Please report any issues running the tutorial or the module to Janet Sinsheimer PhD. (jsinshei@g.ucla.edu).

```
In [1]: versioninfo()

Julia Version 1.2.0
Commit c6da87ff4b (2019-08-20 00:03 UTC)
Platform Info:
    OS: macOS (x86_64-apple-darwin18.6.0)
    CPU: Intel(R) Core(TM) i7-6567U CPU @ 3.30GHz
    WORD_SIZE: 64
    LIBM: libopenlibm
    LLVM: libLLVM-6.0.1 (ORCJIT, skylake)
```

When to use MendelGeneticCounseling

MendelGeneticCounseling.jl is capable of calculating the risk of an underlying genotype (e.g. Homozygous_Normal, Heterozygous, Homozygous_Mutant) given a family history and individual risk factors including closely linked genetic markers. It can use parametric models or a penetrance file.

Example 1 uses a parametric model. The parametric models are currently restricted to the following generalized linear model distributions, binomial, exponential, gamma, inverse gaussian, logistic, lognormal, negative binomial, Poisson, and of course, the normal distributions but more distributions can be added to the apply_dist.jl function. The inverse link functions available are for the links: log, logit, cauchit, complementary log log, inverse (1/x), probit, and square root. More can be added to apply_inverse_link_new.jl. We will be adding the capability to run censored survival models soon.

MendelGeneticCounseling.jl is also capable of using a penetrance file that provides the probability that an individual is affected conditional their genotype and risk factors (Example 2).

This module is a prototype and features will be added. Currently we are working on models that can handle censoring, data in snp binary or vcf files, and mutation at the risk locus.

Analysis keywords needed for a glm penetrance function.

| Keyword | Default Value | Allowed value | Description |
|--------------|------------------|--|--|
| glm_mean | 0.0 | symbolic expression | provides the form of x^b |
| glm_response | "Normal" | GLM distribution | One of the following distribution choices: Binomial, exponential, gamma, inverse Gaussian, logistic, lognormal, negative binomial, Poisson, normal |
| glm_link | "IdentityLink" | Inverse Link Function for the appropriate GLM link | CauchitLink,CloglogLink, IdentityLink, InverseLink, LogitLink, LogLink, ProbitLink, Sqrt |
| glm_trait | Affected | String | Defines the column in the pedigree file that contains the trait phenotype when using a glm for the penetrance. Note: Individuals' values must be quantitative at this time |
| glm_scale | 1.0 | Positive real number | measure of the spread of the trait - in the case of the normal distribution, the standard deviation |
| glm_trials | 1 | Positive Integer | For logistic regression glm_trials = 1 |

Analysis keywords if using a penetrance file

| Keyword | Default Value | Allowed value | Description | |
|-----------------|------------------|---------------|--|--|
| disease_status | 11 11 | String | Defines the column in the pedigree file that contains the trait phenotype when including a penetrance file. Values affected = 1, unaffected = 0, unknown = -1 | |
| penetrance_file | 11 11 | String | The absence of a penetrance file automatically results in a parametric penetrance file. The presence of a penetrance file results in discrete, user defined values based on risk classes | |

Analysis keywords - input and output common to both options.

| Keyword | Default Value | Allowed value | Description |
|----------------|-------------------|---------------|---|
| locus_file | шш | String | Provides the population specific locus names, allele names and allele frequencies |
| output_file | Mendel_Output.txt | String | OpenMendel generated output file with table of kinship coefficients |
| pedigree_file | | String | Numerator and Denominator pedigree |
| phenotype_file | нн | String | provides genetic model for the underlying genetic locus |

A list of OpenMendel keywords common to most analysis package can be found https://openmendel.github.io/MendelBase.jl/#keywords-table)

Example 1: Probability that an individual is heterozygous given pedigree and covariate information.

Using a gamma distribution to model the penetrance

Data used in Example 1:

The input files for all examples in this tutorial can be obtained from https://github.com/OpenMendel/GeneticCounseling_ASHG2019 (https://github.com/OpenMendel/GeneticCounseling_ASHG2019)

The data are from an example pedigree used in the Mendel version 16.0 release. The pedigree structure and phenotypes are originally from Schrott et al. (1972) Ann Int Med 76:711–720. We have used the pedigree to provide a slightly contrived example in which Mother III13, who has cholesterol value 440 at age 21 is concerned that her young son might also be affected with extreme hypercholesterolemia. This first analysis calculates the probability that person IV11 has a heterozygous genotype given his covariates and their relatives' information.

Step 1: Examine the pedigree file:

Recall what is needed in a valid pedigree structure (https://openmendel.github.io/MendelBase.jl/#pedigree-file).

'MendelGeneticCounseling.jl' calculates the conditional probability of an individual being affected given the pedigree information by calculating the joint probability of the family and the individual's affection status (numerator pedigree) divided by the probability of the family (denominator pedigree). Accordingly the user needs to provide the program with a pedigree file with a numerator pedigree that includes the genotype of the individual of interest and the denominator pedigree doesn't. In our example we have named the pedigrees Top and Bottom but the choice of two names is left to the user as long as they are distinct.

On a Mac you can view the pedigree file by typing "; cat PedChol.csv" This command should work on a PowerShell on a PC. If you are using a cmd window in Dos use "; type PedChol.csv" instead.

In [2]: ;cat PedChol.csv

```
Pedigree,Person,Father,Mother,Sex,HC,Age ,lnChol,Chol,High_Chol
               , II2
                           , II1
                                       ,2,,22,6.39,595,1
  TOP , III11
  TOP , II16
                    12
                              I1
                                       ,1,,38,6.17,479,1
  TOP , III56
                   II19
                             II18
                                       ,2,,4,6.12,454,1
                            , I1
  TOP , II18
                   12
                                       ,2,,36,6.1,448,1
                   II2
                            , II1
  TOP , III13
                                       ,2,,21,6.09,440,1
                            , II17
 TOP , III50
                   II16
                                       ,2,,10,6.07,433,1
                           , II1
 TOP , III7
                   II2
                                       ,1,,26,6.05,425,1
                ,
                            , II1
  TOP , III1
                   II2
                                       ,1,,31,6.05,422,1
                ,
                            , I1
                                       ,1,,27,6.03,416,1
  TOP , II29
                ,
                           , II21
                                       ,1,,6,6.02,413,1
  TOP , III58
                   II20
                            , II18
  TOP , III55
                   II19
                                       ,1,,10,6.02,412,1
                                       ,1,,20,5.97,391,1
  TOP , III15
                             II1
                   II2
  TOP ,
        III41
                              II14
                                       ,2,,15,5.94,381,1
                    II15
 TOP ,
        II20
                    Ι2
                               I1
                                       ,1,,35,5.94,379,1
                ,
  TOP ,
        III54
                    II19
                               II18
                                       ,2,,13,5.93,377,1
  TOP ,
        II22
                    Ι2
                               Ι1
                                       ,1,,32,5.93,376,1
  TOP ,
        III26
                    II7
                              II8
                                       ,1,,20,5.93,376,1
 TOP ,
        III61
                    II22
                              II23
                                       ,2,,6,5.92,374,1
                            ,
 TOP ,
        III43
                    II15
                               II14
                                       ,2,,10,5.91,369,1
 TOP ,
        IV10
                    III12
                               III11
                                       ,2,,7,5.46,235,1
                                       ,1,,41,5.9,365,1
 TOP ,
        II12
                   Ι2
                              Ι1
 TOP ,
        III17
                   II2
                              II1
                                       ,2,,15,5.9,365,1
 TOP ,
        III46
                   II16
                              II17
                                       ,2,,16,5.9,365,1
                                       ,1,,12,5.83,341,1
 TOP ,
                   III1
                              III2
        IV1
 TOP ,
        III38
                   II12
                              II13
                                       ,2,,4,5.82,337,1
  TOP ,
        III9
                   II2
                              II1
                                       ,2,,24,5.8,331,1
                            , II14
 TOP ,
        III39
                   II15
                                       ,1,,20,5.78,324,1
 TOP ,
                            , II18
                                       ,1,,16,5.77,320,1
        III53
                   II19
 TOP ,
        III27
                   II7
                             II8
                                       ,1,,17,5.76,317,1
                 ,,,1,,35,5.75,313,1
 TOP ,
        II19
                            , II17
 TOP ,
                 , II16
                                       ,1,,11,5.73,309,1
        III49
  TOP ,
        II14
                 , 12
                               Ι1
                                       ,2,,39,5.7,299,1
 TOP ,
                 ,,,1,,51,5.69,296,1
        II2
                 , II12
  TOP ,
        III36
                          , II13
                                       ,1,,16,5.69,295,1
                 ,,,2,,41,5.66,286,1
  TOP ,
        II6
                , II2
  TOP ,
                           , II1
        III16
                                       ,1,,16,5.64,281,1
 TOP ,
                , II15
                                       ,2,,19,5.63,280,1
        III40
                              II14
 TOP ,
                 ,,,2,,29,5.6,271,1
        III4
 TOP ,
                , II20
                          , II21
        III59
                                       ,2,,5,5.58,265,1
                , II12
 TOP , III35
                            , II13
                                       ,2,,17,5.55,257,1
                 ,,,2,,36,5.55,256,1
  TOP , II4
                , II3
                          , II4
                                       ,2,,12,5.53,252,1
  TOP , III18
                , II3
  TOP , III19
                             II4
                                       ,2,,11,5.52,250,1
                   III14
                              III13
  TOP , IV11
                                       ,1,Heterozygous,1,5.51,248,1
                , 12
  TOP ,
        II24
                              Ι1
                                       ,1,,30,5.51,246,1
                , II32
  TOP, III68
                               II33
                                       ,2,,1,5.51,246,1
                ,,,2,,30,5.5,245,1
  TOP , III2
                 ,,,1,,48,5.5,244,1
  TOP , II28
                , II2
                            , II1
                                       ,1,,25,5.49,242,1
  TOP , III8
                 , II10
  TOP , III32
                               II9
                                       ,1,,8,5.48,241,1
                 ,,,1,,47,5.46,236,1
  TOP , II15
 TOP ,
                            , III4
                                       ,2,,7,5.46,235,1
        IV4
                   III3
 TOP ,
        III20
                    II3
                               II4
                                       ,1,,10,5.46,234,1
  TOP ,
                 ,,,2,,36,5.45,232,1
        II17
  TOP ,
                            , II14
        III42
                   II15
                                       ,1,,12,5.44,231,1
 TOP ,
        III3
                    II2
                               II1
                                       ,1,,30,5.4,221,0
 TOP ,
        III31
                    II10
                               II9
                                       ,2,,12,5.4,221,0
 TOP ,
                                       ,1,,29,5.39,220,0
        III5
                   II2
                               II1
 TOP ,
                   II24
                              II25
                                       ,2,,4,5.39,220,0
        III63
 TOP ,
                 , II24
        III62
                            , II25
                                       ,2,,6,5.39,219,0
                 ,,,2,,26,5.38,216,0
  TOP , II25
 TOP ,
                 , II12
        III37
                           , II13
                                       ,1,,15,5.37,214,0
 TOP ,
        III65
                    II26
                               II27
                                       ,2,,5,5.37,214,0
  ΨΩР .
                    TTT1
                               TTT2
                                       _2__9_5_37_214_0
        TV2
```

The top line provides the column names. The first 5 columns indicate the necessary pedigree information including sex. The 6th column label, HC, is the risk locus genotype, which is unknown for most individuals. The 7th column label, Age, is age in years. The 8th column label, InChol, is log base e of the cholesterol value, the 9th column label, Chol, is total cholesterol in mg/dl, and 10th column label is a dichotomize trait where total cholesterol greater than or equal to 225 is denoted as 1 for affected and less than 225 is denoted as 0. Missing values are denoted as NA.

If you examine the file carefully you will see that all the information in the numerator pedigree is repeated in the denominator pedigree except that we specify individual IV11's genotype in the numerator pedigree but not the denominator pedigree. The likelihood of the numerator pedigree divided by the likelihood of denominator pedigree gives us $P(G_{IV11} = 1/2 | \mathbf{G}, \mathbf{Chol}, \mathbf{Age}, \mathbf{Sex})$, the "risk."

Step 2: Examine the control file

A control file gives specific instructions to MendelGeneticCounseling. We specify the dependent variable with the keyword glm trait=Chol. The trait name must correspond exactly to the column name in the pedigree file.

In this example we treat the cholesterol values, which are highly right tailed, as gamma distributed. This information is specified by the keyword glm_response = GammaDist and glm_link= LogLink.

The glm_mean is the linear equation, x^tb .

Specifically, $f(y) = (\frac{\sigma}{\mu})^{\sigma} \frac{y^{\sigma-1}e^{\frac{-\sigma y}{\mu}}}{\Gamma(\mu)}$ where the scale $\sigma = 44.68$ and the mean, $\mu = g(x^tp) = e^{(4.691+0.562(\max(allele1,allele2))+0.00194Age+0.036Sex)}$ and g is the exponential function which is the inverse link function for \ln . Note that allele1/allele2 represents the current genotype presented to the penetrance function.

We specify the "normal allele" as 1 and the "mutant allele" as 2 according to their order of appearance in the locus file. By using the expression $\max(allele1, allele2)$, we are specifying that genotype 1/1 adds 0.562, and genotype 1/2 or 2/2 adds 1.124 to $\ln \mu$. In other words, we are modeling the locus as dominant. Sex is coded here as 1 for males and 2 for females. Male adds 0.036 to and female adds 0.072 to $\ln \mu$. Age is measured in years; each year adds 0.00194 to $\ln \mu$. Thus a 50 year old female with genotype 1/1 has an expected cholesterol value of $e^{(4.691+0.562+0.00194(50)+0.036(2))} = 226.3$.

The keyword glm_trials is ignored in this analysis. It is used when running logistic regression mean models, where glm_response = BinomialDist and glm_link = LogitLink, binomial and negative binomial models.

```
In [3]:    ;cat ControlParametricPenetranceExample.txt

#
    # Input and Output files.
#
    locus_file = LocusChol.txt
    pedigree_file = PedChol.csv
    phenotype_file = PhenoChol.txt
    output_file = CholHeterozygousRisk.txt
#
    # Analysis parameters for Genetic Counseling option.
#
    glm_mean = 4.691+0.562(max(allele1,allele2))+0.00194Age+0.036Sex
    glm_response = GammaDist
    glm_link = LogLink
    glm_trait = Chol
    glm_scale = 44.68
    glm trials = 1
```

Step 3: Examine the Locus and Phenotype Files.

The locus file provides the name of the putative disease locus. The name must match exactly the column name in the pedigree file. In this simple example, this locus is unobserved and so no one has a genotype in the pedigree file except person IV11. The locus HC has two alleles "+" and "-". The locus is on an autosomal chromosome. The "-" allele has frequency 0.98 and the "+" allele has frequency 0.02. The phenotype file provides the genetic model for this locus. The "-/-" genotype has phenotype "Homozygous_Normal", the "+/-" genotype has phenotype "Heterozygous", and the "+/+" genotype has phenotype "Homozygous_Mutant."

Step 4: Run the analysis in Julia REPL or directly in notebook

The first command using MendelGeneticCounseling loads the MendelGeneticCounseling module. This command needs to be issued just once during this tutorial. The next command GeneticCounseling reads the control file, reads in the data and runs the analysis.

```
In [6]: using MendelGeneticCounseling
        GeneticCounseling("ControlParametricPenetranceExample.txt")
             Welcome to OpenMendel's
         Genetic Counseling Analysis Option
        Reading the data.
        The current working directory is "/Users/janets/Documents/lap top janet/janet/sh
        ort courses 2019/ASHGproposal/GeneticCounseling ASHG2019-master".
        Keywords modified by the user:
          control file = ControlParametricPenetranceExample.txt
          glm link = LogLink
          glm mean = 4.691+0.562(max(allele1,allele2))+0.00194Age+0.036Sex
          glm response = GammaDist
          glm scale = 44.68
          glm trait = Chol
          glm trials = 1
          locus file = LocusChol.txt
          output_file = CholHeterozygousRisk.txt
          pedigree file = PedChol.csv
          phenotype_file = PhenoChol.txt
         no penetrance file
        Analyzing the data.
         The risk = 0.27557.
        Mendel's analysis is finished.
```

Step 4: Interpreting the result

MendelGeneticCounseling should have generated a file CholHeterozygousRisk.txt in your local directory. The value is the conditional probability that person IV11 is heterozygous given the information provided regarding the family and the individuals' age and genotype. The probability is 0.27557.

Step 5: Test yourself.

- (1) Modify the pedigree file and the control file to calculate the probability that individual IV11 has the homozygous normal genotype. Then rerun MendelGeneticCounseling with these new data. You should find the probability is 0.71335.
- (2) Modify the pedigree file to determine the probability individual III13 has the heterozygous genotype. You should find the probability is 0.96698.

Help on modifying the jupyter notebook so you can rerun MendelGeneticCounseling without destroying your original results:

First go to the file tab and press "Save and checkpoint" then go to the insert and click "insert cell below". Then insert command to run GeneticCounseling with your new control file.

Example 2: Using a penetrance file.

In this example we illustrate how to use a penetrance file. We thank Brian Shirts for pointing us to the "Analyze My Variant" website (http://analyzemyvariant.com (http://analyzemyvariant.com (http://analyzemyvariant.com/brca1-info (<a

BrianspedigreeMay162019.jpg

Step 1: Examine the Penetrance file

We have set up the penetrance file to have 5 columns. The first column is for the risks for homozygous wild type genotype (1/1) for each sex and risk decade (penetrance class). The second column is for the risks for heterozyous genotype (1/2) by penetrance class. The third column is for the risks for the homozygous high risk genotype (2/2) by penetrance class. The next column corresponds to the sex of the individual. The final column corresponds to the risk decade of the individual (1: $0 \le age < 20$, 2: $20 \le age < 30$, 3: $30 \le age < 40$, 4: $40 \le age < 50$, 5: $50 \le age < 60$, 6: $60 \le age < 70$, and 7: $70 \le age$).

To view the file type:

```
In [7]:
        ;cat PenBRCAExample.csv
        Homozygous Normal, Heterozygous, Homozygous Mutant, Sex, Risk decade
        0.000000885,0.001025896,0.001025896,female,1
        0.000040997,0.047524,0.047524,female,2
        0.00189916,0.18042,0.18042,female,3
        0.00878848,0.3736,0.3736,female,4
        0.0275136,0.5752,0.5752,female,5
        0.05646,0.6889,0.6889,female,6
        0.0793,0.785,0.785,female,7
        7.58E-08,1.07E-05,1.07E-05,male,1
        0.0000012,0.00017,0.00017,male,2
        0.000019,0.0012,0.0012,male,3
        0.000085,0.003,0.003,male,4
        0.00027,0.0062,0.0062,male,5
        0.00067,0.012,0.012,male,6
        0.0012,0.018,0.018,male,7
```

Step 2: Examine the pedigree file

In this example we are interested in determining the probability that individual 18, a 38 year old female who is currently unaffected with breast cancer is heterozygous for a BRCA1 mutation. In this case, some members of her family have genotypes at the locus.

Like the previous example, the pedigree is present in two copies. The first copy (called Top) has the genotype of individual 18 as heterozygous. The second copy (called Bottom) has her genotype missing. The likelihood of the first pedigree divided by the likelihood of second pedigree gives us $P(G_{IV11} = 1/2 | \mathbf{G}, \mathbf{CancerStatus}, \mathbf{Age}, \mathbf{Sex})$, the "risk."

```
In [8]: ;cat PedBRCAExample.csv
          Pedigree, Person, Father, Mother, Sex, BRCA, Age, Risk decade, Proband, Cancer
          TOP, 1, 0, 0, male, , 79, 7, 0, -1
          TOP, 2, 0, 0, female, 78, 7, 0, -1
          TOP, 3, 1, 2, female, Heterozygous, 40, 4, 0, 1
          TOP, 4, 1, 2, female, ,79, 7, 0, 0
          TOP, 5, 1, 2, female, ,85, 7, 0, 1
          TOP, 6, 1, 2, female, , 43, 4, 0, 1
          TOP,7,0,0,male,,80,7,0,0
          TOP,8,7,3,male,,73,7,0,0
          TOP, 9, 7, 3, male, ,41, 4, 0, 0
          TOP, 10, 0, 0, female, 89, 7, 0, -1
          TOP, 11, 7, 3, male, Heterozygous, 30, 3, 0, 0
          TOP, 12, 0, 0, female, ,80, 7, 0, 0
          TOP, 13, 9, 10, female, Heterozygous, 41, 4, 0, 1
          TOP, 14, 9, 10, male, ,60,6,0,0
          TOP, 15, 9, 10, female, Heterozygous, 50, 5, 0, 1
          TOP, 16, 9, 10, female, Heterozygous, 60, 6, 0, 0
          TOP, 17, 11, 12, female, Heterozygous, 49, 4, 1, 1
          TOP, 18, 11, 12, female, Heterozygous, 38, 3, 0, 0
          TOP, 19, 11, 12, male, Heterozygous, 36, 3, 0, 0
          TOP, 20, 11, 12, female, Heterozygous, 48, 4, 0, 1
          BOTTOM, 1, 0, 0, male, , 79, 7, 0, -1
          BOTTOM, 2, 0, 0, female, ,78, 7, 0, -1
          BOTTOM, 3, 1, 2, female, Heterozygous, 40, 4, 0, 1
          BOTTOM, 4, 1, 2, female, , 79, 7, 0, 0
          BOTTOM, 5, 1, 2, female, , 85, 7, 0, 1
          BOTTOM, 6, 1, 2, female, , 43, 4, 0, 1
          BOTTOM, 7, 0, 0, male, ,80, 7, 0, 0
          BOTTOM, 8, 7, 3, male, , 73, 7, 0, 0
          BOTTOM, 9, 7, 3, male, , 41, 4, 0, 0
          BOTTOM, 10, 0, 0, female, ,89, 7, 0, -1
          BOTTOM, 11, 7, 3, male, Heterozygous, 30, 3, 0, 0
          BOTTOM, 12, 0, 0, female, ,80, 7, 0, 0
          BOTTOM, 13, 9, 10, female, Heterozygous, 41, 4, 0, 1
          BOTTOM, 14, 9, 10, male, ,60,6,0,0
          BOTTOM, 15, 9, 10, female, Heterozygous, 50, 5, 0, 1
          BOTTOM, 16, 9, 10, female, Heterozygous, 60, 6, 0, 0
          BOTTOM, 17, 11, 12, female, Heterozygous, 49, 4, 1, 1
          BOTTOM, 18, 11, 12, female, , 38, 3, 0, 0
          BOTTOM, 19, 11, 12, male, Heterozygous, 36, 3, 0, 0
          BOTTOM, 20, 11, 12, female, Heterozygous, 48, 4, 0, 1
```

Step 3: Examine the Locus and Phenotype File

These files are very similar to the ones we used in the first example.

```
In [9]: ; cat LocusBRCAExample.txt

Locus,Allele,Chromosome,European
BRCA,"1",Autosome,0.998
BRCA,"2",Autosome,0.002

In [10]: ; cat PhenoBRCAExample.txt

Locus,Phenotype,Genotypes
BRCA,Homz_rare,"2/2"
BRCA,Heterozygous,"1/2"
BRCA,Homz common,"1/1"
```

Step 4: Examine the control file

This control file has some of the same features as parametric ones but it doesn't need to specify the GLM values because there is a penetrance file. Besides the input and output file names, the only needed information is the name of the Column in the pedigree that contains the information regarding breast cancer status.

```
In [11]:    ;cat ControlBRCAexample.txt

#
    # Input and Output files.
#
    locus_file = LocusBRCAExample.txt
    pedigree_file = PedBRCAExample.csv
    phenotype_file = PhenoBRCAExample.txt
    penetrance_file = PenBRCAExample.csv
    output_file = BRCAExampleOut.txt
    #
    disease_status = Cancer
```

Step 5: Running the analysis

```
In [12]: using MendelGeneticCounseling
         GeneticCounseling("ControlBRCAexample.txt")
              Welcome to OpenMendel's
          Genetic Counseling Analysis Option
         Reading the data.
         The current working directory is "/Users/janets/Documents/lap_top_janet/janet/sh
         ort_courses_2019/ASHGproposal/GeneticCounseling_ASHG2019-master".
         Keywords modified by the user:
           control_file = ControlBRCAexample.txt
           disease status = Cancer
           locus file = LocusBRCAExample.txt
           output file = BRCAExampleOut.txt
           pedigree_file = PedBRCAExample.csv
           penetrance file = PenBRCAExample.csv
           phenotype_file = PhenoBRCAExample.txt
         this problem has 2 factors called Symbol[:Sex, :Risk decade]
         Analyzing the data.
          The risk = 0.45091.
         Mendel's analysis is finished.
```

Step 6: Interpreting the Result

Again you should get a file with the results. The probability of that individual 18 is heterozygous is 0.45091.

Step 7: Test yourself.

- (1) Modify the pedigree file and the control file to calculate the probability that individual 18 has the homozygous normal genotype. Then rerun MendelGeneticCounseling with these new files.
- (2) How would the risk change if individual 18 were 20 years old instead 38 years old? What if she were 68 years old? (To determine, change her risk decade in the pedigrees and rerun the analysis).

Final Comments

The Julia version of Mendel, provides an opportunity for the user to easily modify the code to suit their own needs. All the source code is provided and Julia is both accessible and very fast.

NOTE: When Finished with this notebook. Go to the File tab and first select save and checkpoint to save your results. Then under the file tab, select close and halt to prevent copies of Jupyter notebook from running indefinitely in the background