crimaptools: a tutorial

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This crimaptools package creates input files, runs CRI-MAP and parses output files for analyses of linkage mapping and recombination rate estimation in CRI-MAP v2.504. This packages is still in progress in terms of optimisation (particularly dealing with unusual CRI-MAP syntax), but should run if instructions are followed carefully. To download CRI-MAP, which is currently tirelessly maintained by Jill Maddox, go here. For more information on how to use it from the command line, check out Paris Veltsos's tutorial here.

The package can be installed using devtools:

```
library(devtools)
install_github("susjoh/crimaptools")
```

1. Example dataset.

crimaptools requires two inputs to create, run and parse files:

- 1. A GenABEL gwaa.data object containing genotype information for IDs
- 2. A pedigree object that specifies the individual families used in CRI-MAP, with columns for the ANIMAL, FATHER, MOTHER and FAMILY.

An example dataset from Red deer is included, and can be called using data(deer)

```
data(deer)
ls()

## [1] "deer.abel" "deer.famped" "deer.ped"

deer.famped
```

```
##
      ANIMAL FATHER MOTHER
                                        Family
## 1
        4440
                   0
                          0 Offspring_Mum_108
## 2
        3018
                   0
                          0 Offspring_Mum_108
## 3
        1169
                   0
                          0 Offspring_Mum_108
        1806
                       3018 Offspring_Mum_108
## 4
                1169
## 5
         108
                4440
                       1806 Offspring_Mum_108
## 6
        3049
                   0
                          0 Offspring_Mum_109
## 7
        1289
                   0
                          0 Offspring_Mum_109
## 8
         329
                   0
                          0 Offspring_Mum_109
## 9
                       1289 Offspring_Mum_109
        1911
                 329
## 10
         109
                3049
                       1911 Offspring_Mum_109
## 11
        1303
                   0
                          O Offspring_Mum_110
## 12
        1289
                   0
                          O Offspring_Mum_110
## 13
         329
                   0
                          O Offspring_Mum_110
## 14
        1911
                       1289 Offspring Mum 110
                 329
                       1911 Offspring_Mum_110
## 15
         110
                1303
```

```
## 16
        1920
                   0
                          O Offspring_Mum_111
## 17
        1289
                   0
                          O Offspring_Mum_111
## 18
         329
                   0
                          O Offspring Mum 111
## 19
        1911
                       1289 Offspring_Mum_111
                329
## 20
         111
               1920
                       1911 Offspring_Mum_111
```

2. Creating a CRI-MAP input file.

This is done using the function <code>create_crimap_input</code>. This requires the <code>deer.abel</code> and <code>deer.famped</code> objects, but also requires additional inputs such as <code>analysisID</code>, used as a flag for running CRI-MAP. An ordered list of SNP loci (<code>snplist =</code>) or a chromosome number (<code>chr =</code>) must be specified. It is also possible to specify the directory to which the output should be written, and <code>clear.existing.analysisID</code> will get rid of any previous builds carried out with the same analysisID. Let's run the analysis for chromosome 3, and give it the analysisID 3a.

```
## Recoding alleles to numeric values...
## ...done
## Merging pedigree and genotype information...
## ...done.
## Parsing and writing to crimap/chr3a.gen...
## ...done
```

In the directory crimap, an input file chr3a.gen will have been created.

3. Run prepare and extract and deal with mendelian errors.

The .gen file must now be run through *prepare* to produce the field used for linkage mapping and crossover estimation. This can be done using the function run_crimap_prepare. At present this inefficiently has to define the path to the CRI-MAP exectuable (in future versions it will be bundled with the library). At the moment, the exectuable path is specified relative to the .gen file - for example, below, the CRI-MAP executable is in the same directory as the .gen file:

```
run_crimap_prepare(crimap.path = "crimap2504.exe", genfile = "crimap/chr3a.gen")
dir("crimap")
```

The function has produced the .pre, .loc, .par and .dat files. The .pre file will contain information on mendelian errors between parents and offspring, which can be extracted using the parse_mend_err function. This creates a further file with the extension .mnd, which has a column for each ID and the problematic locus:

Writing new file crimap/chr3a.mnd

```
read.table("crimap/chr3a.mnd", header = T)
```

```
## ANIMAL SNP.Name
## 1 111 cela1_red_3_74862326
## 2 1920 cela1_red_3_74862326
```

Information from this file can be used to mask mendelian errors in the .gen file, by rerunning the create_crimap_input with use.mnd = TRUE:

```
## dono
```

Recoding alleles to numeric values...

```
## ...done
```

Merging pedigree and genotype information...

```
## ...done.
```

Parsing and writing to crimap/chr3a.gen...

```
## ...done
```

```
run_crimap_prepare(crimap.path = "crimap2504.exe", genfile = "crimap/chr3a.gen")
parse_mend_err(prefile = "crimap/chr3a.pre", genfile = "crimap/chr3a.gen", familyPedigree = deer.famped
```

No Mendelian errors detected. No changes made to .mnd file.

4. Build a linkage map.

Now the dataset is ready for linkage mapping and characterisation of crossovers. This assumes the order of the deer abel dataset when chromosome is specified; if snplist was specified in create_crimap_input then it assumes the order of the snplist itself. The linkage map can be obtained by running run_crimap_map, which may be slow for large numbers of markers/families, and then parsed with parse_map, which provides sex specific maps:

```
run_crimap_map(crimap.path = "crimap2504.exe", genfile = "crimap/chr3a.gen")
dir("crimap")
## [1] "chr3a.dat"
                                          "chr3a.loc"
                                                            "chr3a.map"
                         "chr3a.gen"
## [5] "chr3a.ord"
                         "chr3a.par"
                                          "chr3a.pre"
                                                            "crimap2504.exe"
## [9] "crimapinput1"
deer.map <- parse_map(mapfile = "crimap/chr3a.map")</pre>
head(deer.map)
##
      Order
                        SNP.Name cMPosition.Female cMPosition.Male Female.r
## 1
          0 cela1 sika 3 247473
                                              0.000
                                                                   0
                                                                        0.000
          1 cela1_sika_3_1709255
                                                                        0.211
## 3
                                              0.000
                                                                   0
## 5
          2 cela1_sika_3_1763550
                                             22.546
                                                                   0
                                                                        0.211
                                                                        0.000
## 7
          3 cela1 sika 3 2122831
                                             45.091
                                                                   0
## 9
          4 cela1_red_3_3089490
                                             45.091
                                                                   0
                                                                        0.000
                                                                        0.000
## 11
          5 cela1_red_3_3118111
                                             45.091
                                                                   0
##
      cMdiff.Female Male.r cMdiff.Male analysisID
## 1
              0.000
                         0
                                      0
                                                3a
             22.546
                         0
                                      0
## 3
                                                3a
                                      0
## 5
             22.546
                         0
                                                3a
## 7
              0.000
                         0
                                      0
                                                За
## 9
              0.000
                         0
                                      0
                                                3a
## 11
              0.000
                         0
                                                3a
```

${\bf 5.} \ \ {\bf Characterising} \ \ {\bf recombination} \ \ {\bf events}.$

The recombination events can be obtained by running run_crimap_chrompic, which again may be slow for larger numbers of markers/families. A sex averaged linkage map can be parsed with parse_map_chrompic and crossovers can be extracted with parse_crossovers, which also requires the family pedigree:

```
run_crimap_chrompic(crimap.path = "crimap2504.exe", genfile = "crimap/chr3a.gen")
deer.cmpmap <- parse_map_chrompic(chrompicfile = "crimap/chr3a.cmp")
head(deer.cmpmap)</pre>
```

```
##
      Order
                       SNP.Name cMPosition
                                                r cMdiff analysisID
         1 cela1_sika_3_247473
## 1
                                    0.000 0.000 0.000
                                                                За
## 3
                                     0.000 0.211 22.546
          2 cela1_sika_3_1709255
                                                                 3a
## 5
          3 cela1_sika_3_1763550
                                    22.546 0.211 22.546
                                                                 За
## 7
          4 cela1_sika_3_2122831
                                    45.091 0.000 0.000
                                                                 3a
## 9
          5 cela1_red_3_3089490
                                    45.091 0.000 0.000
                                                                 За
## 11
         6 cela1_red_3_3118111
                                    45.091 0.000 0.000
                                                                 3a
```

```
deer.xovers <- parse_crossovers(chrompicfile = "crimap/chr3a.cmp", familyPedigree = deer.famped)
deer.xovers[1:2,]</pre>
```

```
ANIMAL RecombCount parent
                                            Family No.Inf.Loci First.Inf.Order
##
## 1
        108
                                                            480
                       1 MOTHER Offspring Mum 108
                                                                               2
## 2
        109
                       3 MOTHER Offspring Mum 109
                                                            421
##
     Last.Inf.Order FATHER MOTHER RRID analysisID
## 1
               1845
                       4440
                              1806 1806
## 2
               1786
                       3049
                              1911 1911
                                                 3a
##
                              UniqueID
## 1 3a_Offspring_Mum_108_1806_MOTHER
## 2 3a_Offspring_Mum_109_1911_MOTHER
```

Column data is the inheritance pattern from grandparents, with each character representing an ordered SNP. 0 is grandmaternal, 1 is grandpaternal. RRID is the parent in which the meiosis took place.

6. Investigating doubles crossovers.

Genotyping and/or phasing errors can lead to erroneous calls of double crossovers. These can be investigated by running check_double_crossovers on the parsed crossovers:

```
deer.doubles <- check_double_crossovers(parsed.xovers = deer.xovers)</pre>
```

Splitting chromosome into segments of shared grandparental origin

```
head(deer.doubles)
```

```
##
     Phase StartPos StopPos StartSpan StopSpan InfCount Segment Segment.Count
## 1
         1
                   4
                          62
                                                        19
                                      1
                                              117
                                                                  1
         0
                                                                                 2
## 2
                 117
                        1845
                                     62
                                             1845
                                                        461
                                                                  2
## 3
         0
                   2
                            2
                                      1
                                                8
                                                          1
                                                                  1
                                                                                 4
                                                                  2
                                      2
                                              402
                                                        96
                                                                                 4
## 4
         1
                   8
                         395
## 5
         0
                 402
                         402
                                    395
                                              403
                                                          1
                                                                  3
                                                                                 4
## 6
                 403
                                    402
                                             1786
                                                       323
                                                                  4
         1
                        1786
                                                          Family RRID parent
##
      Type
                                     UniqueID
## 1 First 3a_Offspring_Mum_108_1806_MOTHER Offspring_Mum_108 1806 MOTHER
      Last 3a_Offspring_Mum_108_1806_MOTHER Offspring_Mum_108 1806 MOTHER
## 3 First 3a_Offspring_Mum_109_1911_MOTHER Offspring_Mum_109 1911 MOTHER
## 4
       Mid 3a_Offspring_Mum_109_1911_MOTHER Offspring_Mum_109 1911 MOTHER
## 5
       Mid 3a Offspring Mum 109 1911 MOTHER Offspring Mum 109 1911 MOTHER
     Last 3a_Offspring_Mum_109_1911_MOTHER Offspring_Mum_109 1911 MOTHER
##
     analysisID Singleton
## 1
             3a
## 2
             3a
                        no
## 3
             3a
                       yes
## 4
             3a
                        no
## 5
             3a
                       yes
## 6
             3a
                        no
```

This takes the phasing information and returns information on the Phase fragments per chromosome (i.e. runs from a single grandparent. Phase: 0 = grandmaternal, 1 = grandpaternal; StartPos and StopPos are the first and last informative positions of the fragment, StartSpan and StopSpan are the closest informative SNPs on either side of the fragment (if first or last, then is first or last SNP); InfCount is the number of informative SNPs in the fragment; Segment is the order of the fragment, numbered 1:N; Type is whether the fragment

was the first, last or occured in the middle of the chromosome; RRID is the individual in which the meiosis occurred.

Physical map infomation for the markers can also be added by specifying map positions in a data frame with headers SNP.Name, Position, Order and analysisID:

```
library(GenABEL)
```

```
## Loading required package: MASS
```

Loading required package: GenABEL.data

Splitting chromosome into segments of shared grandparental origin

```
head(deer.doubles)
```

```
Phase StartPos StopPos StartSpan StopSpan InfCount Segment Segment.Count
## 1
                          62
                                                                                2
         1
                   4
                                      1
                                             117
                                                        19
                                                                  1
                                                                                2
## 2
         0
                 117
                        1845
                                     62
                                            1845
                                                       461
                                                                  2
                                                                                4
## 3
         0
                           2
                                                                  1
                  2
                                      1
                                               8
                                                         1
## 4
         1
                   8
                         395
                                      2
                                             402
                                                        96
                                                                  2
                                                                                4
                                                                  3
## 5
         0
                 402
                         402
                                    395
                                             403
                                                         1
                                                                                4
                        1786
                                            1786
##
  6
                                                                  4
                                                                                4
         1
                 403
                                    402
                                                       323
##
      Type
                                     UniqueID
                                                          Family RRID parent
## 1 First 3a_Offspring_Mum_108_1806_MOTHER Offspring_Mum_108 1806 MOTHER
      Last 3a_Offspring_Mum_108_1806_MOTHER Offspring_Mum_108 1806 MOTHER
## 3 First 3a_Offspring_Mum_109_1911_MOTHER Offspring_Mum_109_1911_MOTHER
       Mid 3a_Offspring_Mum_109_1911_MOTHER Offspring_Mum_109 1911 MOTHER
## 5
       Mid 3a_Offspring_Mum_109_1911_MOTHER Offspring_Mum_109 1911 MOTHER
      Last 3a_Offspring_Mum_109_1911_MOTHER Offspring_Mum_109 1911 MOTHER
## 6
     analysisID StartPos.GenomePos StopPos.GenomePos StartSpan.GenomePos
##
## 1
             3a
                            2122831
                                               7067364
                                                                      247473
## 2
             3a
                           10792068
                                             121375733
                                                                     7067364
## 3
             3a
                            1709255
                                               1709255
                                                                      247473
             3a
## 4
                            3155501
                                              29013193
                                                                     1709255
## 5
             3a
                           29656063
                                              29656063
                                                                    29013193
## 6
                           29689850
                                             117062248
                                                                    29656063
##
     StopSpan.GenomePos PosLength SpanLength Singleton
## 1
                10792068
                           4944533
                                      10544595
              121375733 110583665
## 2
                                     114308369
                                                       no
## 3
                 3155501
                                 0
                                       2908028
                                                      yes
## 4
               29656063
                          25857692
                                      27946808
                                                      no
## 5
               29689850
                                  0
                                        676657
                                                      yes
## 6
              117062248 87372398
                                      87406185
                                                       no
```

This outputs additional columns with the genome positions for Pos and Span values, and also PosLength and SpanLength, which is the difference between the start and stop positions.

The user then has some choice of which lines may be erroneous. For example, singletons may be removed.

deer.xovers.clean

```
deer.remove <- subset(deer.doubles, Singleton == "yes")
deer.xovers.clean <- revise_double_crossovers(parsed.xovers = deer.xovers, removesections = deer.remove
## [1] "Fixing Problem 1 of 3"</pre>
```

```
##
ANIMAL RecombCount parent
                        Family No.Inf.Loci First.Inf.Order
             1 MOTHER Offspring_Mum_108
## 1
    108
                                  480
                                            2
## 2
    109
             O MOTHER Offspring_Mum_109
                                  421
## 3
    110
             2 MOTHER Offspring_Mum_110
                                  421
                                            2
                                            2
             1 MOTHER Offspring Mum 111
## 4
    111
                                  417
  Last.Inf.Order FATHER MOTHER RRID analysisID
## 1
        1845
             4440
                 1806 1806
                            3a
## 2
        1786
             3049
                 1911 1911
                            3a
## 3
        1786
             1303
                 1911 1911
                            3a
## 4
        1778
             1920
                 1911 1911
                            3a
##
                 UniqueID
## 1 3a_Offspring_Mum_108_1806_MOTHER
## 2 3a_Offspring_Mum_109_1911_MOTHER
## 3 3a_Offspring_Mum_110_1911_MOTHER
## 4 3a_Offspring_Mum_111_1911_MOTHER
```