Documentation hotspot, v. 1.1: Software to Support Sperm-Typing for Investigating Recombination Hotspots

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1 Introduction

hotspot is a software package for detecting and analyzing recombination hotspots [3]. This is done in three steps: Preparation of allele-specific PCR, the actual PCR, and analysis of the PCR products. hotspot contains the two programs asp and aso for designing allele-specific primers and oligos used in the pre-PCR phase. In addition, it contains xov for computing the rate of recombination during the post-PCR phase of a project and the program six for simulating input to xov. Example input data and a script for downloading the mouse genome sequence including the corresponding SNP data are also provided.

2 Dependencies

- libdivsufsort: calculate suffix arrays, https://github.com/y-256/libdivsufsort/
- tabix: index tab-delimited files [1], part of SAMtools [2], http://www.htslib.org/
- gsl: Gnu Scientific Library, http://www.gnu.org/software/gsl/

All three programs are also available via apt-get and similar package managers.

3 Installation

The four programs that make up hotspot are written in C on a computer running Linux and they should work on any UNIX system. However, please contact me at haubold@evolbio.mpg.de if you have any problems with the programs.

• Clone the github repository

```
git clone https://github.com/evolBioInf/hotspot.git
```

• Change into the newly created directory hot spot

 ${\bf cd}\ {\tt hotspot}$

• Construct the configuration files

```
autoreconf -i
```

- Configure the package
 - ./configure

• Compile

make

• Install

```
sudo make install
```

If instead of using make install you would like to install the binaries by hand, copy them from the src_* directories.

4 Getting Started

The following sections give a tutorial introduction to using the components of hot spot.

4.1 aso

aso is a program for designing allele-specific oligonucleotides that are complementary to SNPs contained in recombination hotspots. The program can also find universal oligos that do not intersect any known SNPs or indels.

• List options

```
aso -h
```

• Take a look at two the example hotspot coordinates supplied with the program

```
$ cat data/mus/exampleHotSpots19.txt
# NB: These mouse hotspot data by Smagulova et al. (2011) are provided
   as part of the hotspot software package for designing
   allele-specific primers and oligos (Odenthal-Hesse et al., 2016).
   The hotspot coordinates by Smagulova et al. refer to mm9 and
#
   will therefore not match the current mouse assembly.
# References:
   1) Smagulova et al. (2011). Genome-wide analysis reveals novel
   molecular features of mouse recombination hotspots. Nature,
   472:375-378.
   2) Odenthal-Hesse et al. (2016). hotspot: software to support
   sperm-typing for investigating recombination hotspots.
   Bioinformatics, 2016, 1-2; doi: 10.1093/bioinformatics/btw195.
# int
       chr
              start
                      end
int1
       chr19 3796569 3799969
               3804782 3808182
int2
       chr19
```

Rows starting with a hash are comments and there can be as many comments as you like. This is followed by the hotspot data in four tab-delimited columns: Hotspot name (interval), chromosome, start, and end.

 The two example hotspots come from chromosome 19. To load the genome and SNP data for chromosome 19, execute

```
make get-chr19-data-mus
```

• Verify that the genome data was downloaded:

```
ls data/mus/genome/
```

• Check that the SNP data was downloaded:

Now run aso

```
aso -g data/mus/genome -s data/mus/vcf data/mus/exampleHotSpots19.txt |
head
int1 rs37745172
                 19 3796745 GGCATAAGCCTGTGGTT GGCATAAGTCTGTGGTT
int1 rs214526021 19 3797057
                             GTGAGTTCGAGGCCAGC GTGAGTTCAAGGCCAGC
int1 rs232006218 19 3797126 AGCTTTTCCTCTTTTCT AGCTTTTCTTTTTCT
int1 rs254266733 19 3797132 TCCTCTTTTCTGCCTTT TCCTCTTTCCTGCCTTT
int1 rs212133110 19 3797168 CAGCCTCCTTGTTACTC CAGCCTCCCTGTTACTC
int1 rs243001379 19 3797209 TAGTAGCAGTTCGGCTC TAGTAGCAATTCGGCTC
int1 rs37665146
                 19 3797213 AGCAGTTCGGCTCATAC AGCAGTTCAGCTCATAC
int1 rs221355178 19 3797237 CATTGTTGCTGTTGCCA CATTGTTGTTGCCA
int1 rs232704341
                 19
                     3797253 ACAGTGGTTTCTTGTGT ACAGTGGTCTCTTGTGT
int1 rs250384293 19
                     3797258 GGTTTCTTGTGTTTGGA GGTTTCTTATGTTTGGA
```

The tab-delimited output consists of six columns:

- 1. Interval name
- 2. SNP name
- 3. Chromosome
- 4. SNP position
- 5. First oligo
- 6. Second oligo

The two oligos only differ in the middle, the SNP position.

• Instead of printing allele-specific oligos, aso can also find oligos that do not span any polymorphism. These "universals" are usually longer, say 100 bp (-1) and are constructed when using -u:

```
aso -u -l 100 -g data/mus/genome
-s data/mus/vcf data/mus/exampleHotSpots19.txt |
head
                  3796669 0.894 GAA...
int1 19 3796570
int1 19 3796869
                 3796968
                         1.000
                                 GAG...
int1 19 3797168
                 3797267
                          0.981 TTG...
int1
    19 3797467
                  3797566
                          0.941
                                 GGG...
int1 19 3797766
                 3797865
                          1.000
                                 CCG...
int1 19 3798065
                 3798164
                         1.000
                                 GTG...
int1 19 3798364
                 3798463 0.914
                                 TTT...
int1
    19
         3798663
                  3798762 0.894
                                 TCT...
int1 19 3798962 3799061 0.382 GCA...
int1 19 3799261 3799360 1.000 TAA...
```

The columns indicate

- 1. Interval name
- 2. Start
- 3. End
- 4. Complexity. This measure lies between 0 and 1; a sequence consisting of a single nucleotide would have zero complexity; random sequences with equi-probable nucleotides have an expected complexity of 1. Complexities greater than 1 are truncated to 1. We can sort the output according to complexity:

```
aso -n -l 100 -g data/mus/genome
-s data/mus/vcf data/mus/exampleHotSpots19.txt
sort -n -k 5
head
int1
     19
          3798962
                   3799061
                             0.382
                                    GCA...
          3796570
                   3796669
                             0.894
int1
     19
                                    GAA...
                             0.894
int1
     19
          3798663
                   3798762
                                    TCT...
                                    TTT...
int1
     19
          3798364
                   3798463
                             0.914
int1
     19
          3797467
                   3797566
                             0.941
                                    GGG...
int1 19
          3797168
                   3797267
                             0.981
                                    TTG...
int1 19
          3796869
                   3796968
                             1.000
                                    GAG...
     19
                             1.000
                                    CCG...
int1
          3797766
                   3797865
int1
      19
          3798065
                   3798164
                             1.000
                                    GTG...
          3799261 3799360
                            1.000
int1
      19
                                    TAA...
```

Notice the long microsatellelite in the top sequence.

4.2 asp

asp is a program for designing PCR primers that have a 3'-end complementary to SNPs in the regions flanking a recombination hotspot. The user can set a maximal and a minimal primer length. Within these bounds the program searches for the primer length that comes closest to an optimal GC-content, which the user can also set.

• Run asp to find forward primers in a window of 5kb upstream of the start of the candidate interval:

```
asp -g data/mus/genome -s data/mus/vcf data/mus/exampleHotSpots19.txt | head
int1 rs36588234 19 3791591 GATTCAGCCAACAA 0.43 34.4 GATTCAGCCAACAG 0.50
                                                                           37.4
                             TTTAGATAGTTGGT 0.29 28.6 TTTAGATAGTTGGG
int1
     rs266164282 19
                    3791605
                                                                     0.36
                                                                           31.5
int1 rs233070165 19 3791753 AGAAGAGCAGTCGG 0.57 40.3 AGAAGAGCAGTCGA
                                                                           37.4
                                                                     0.50
int1 rs586881213 19 3791766 GGTGCTCTTACCCA 0.57 40.3 GGTGCTCTTACCCT 0.57
int1 rs258256279 19 3791984 AGTTCCAGGACAGT 0.50 37.4 AGTTCCAGGACAGC
                                                                     0.57
                                                                           40.3
int1 rs215047461 19
                     3792005 GCACATAAAAACTT 0.29 28.6 GCACATAAAAACTC
                                                                     0.36
                                                                           31.5
                 19 3792235 TGATTGCAGAAATT 0.29 28.6 TGATTGCAGAAATC
int1
     rs38604711
                                                                     0.36
                                                                           31.5
int1 rs212603417 19 3792310 TTAAACCTCCCATT 0.36 31.5 TTAAACCTCCCATC 0.43
                                                                           34.4
int1 rs36316803 19 3792353 TATTGTCATGAGCA 0.36 31.5 TATTGTCATGAGCG 0.43
int1 rs579758610 19 3792405 CATTACCCATGAGC 0.50 37.4 CATTACCCATGAGG 0.50
```

The output columns are

- 1. Interval name
- 2. SNP name
- 3. Chromosome
- 4. SNP position
- 5. Primer for first allele
- 6. GC-content for first primer
- 7. Melting temperature for first primer
- 8. Primer for second allele
- 9. CG-content for second primer
- 10. Melting temperature for second primer
- To get the reverse primers in the upstream region, run

```
int1 rs253442444 19 3800434 CTCTCAAGTGCTGG
                                                  0.57 40.3 CTCTCAAGTGCTGA
                                                                                0.50 37.4
int1 rs214858261 19
                     3800484 TGCTCTATGAACCA
                                                  0.43 34.4 TGCTCTATGAACCT
                                                                                0.43
                                                                                      34.4
     rs232338632 19
                     3800485
                             TTGCTCTATGAACC
                                                  0.43 34.4 TTGCTCTATGAACT
                                                                                0.36 31.5
int1
                             TGGCTGTGGCTGTC
                                                             TGGCTGTGGCTGTT
                                                                                     40.3
int1 rs261835571 19
                     3800506
                                                  0.64
                                                      43.2
                                                                                0.57
     rs222895727
                 19
                     3800511
                             TTATGTGGCTGTGG
                                                  0.50
                                                             TTATGTGGCTGTGT
                                                                                0.43
                                                                                      34.4
                                                       37.4
int1
     rs579285301 19
                     3800535 TATTTATTTATTTTTGAGG 0.16
                                                       36.0 ATTTATTTATTTTTGAGA 0.11
                                                                                      32.1
```

Notice that the two primers need not have the same length. By default, asp searches for a primer of length 14–19 with a GC-content as close to 0.5 as possible.

4.3 xov

xov implements a maximum likelihood procedure for estimating the number of crossover events from the observed number of positive and negative allele-specific PCR reactions. In addition, it uses the likelihood ratio method for estimating a confidence interval. By default this interval is set to 95%, but the user can set it to any level desired.

• Take a look at example input for xov

```
cat data/mus/exampleResults.txt
# This mock data set is taken from
    Kauppi et al. (2009). Analysis of human
    recombination products from human sperm.
    In: Keeney, S. (ed.), Meiosis, Volume 1,
    Molecular and Genetic Methods, Volume 557.
# Int
       Chr Start End
                         d|n-k|k
                                     d \mid n-k \mid k
                                                d \mid n-k \mid k
Int1
            1
                    500
                         2000|2|6
                                     600|1|7
                                                200|0|8
                    250
Int2
            1
                         2000|1|5
                                     600|1|7
                                                200 | 1 | 7
Int3
            1
                    250
                         2000|3|3
                                     600|1|7
                                                200|0|8
            1
                    500
                         2000|5|3
                                     6001018
Int4
       1
                                                2001018
Int5
            1
                   1500
                         2000|0|8
                                     600|0|8
                                                200|0|8
                    500 60|5|7
Int6
            1
                                     120 | 11 | 1
                                               240|12|0
```

The columns list

- 1. Interval
- 2. Chromosome
- 3. Start
- 4. End
- 5. Number of molecules, number of positive PCR reactions, number of negative PCR reactions; this triplet of values is repeated for each experiment, three in this example
- Run xov

xov data/mus/exampleResults.txt # Int Chr Start End Len [% %] [cM/Mb cM/Mb cM/Mb] 500 500 0.004 0.015 0.039 30.004 Tnt.1 7.448 78.119 Tnt2 1 250 250 0.004 0.018 0.046 17.596 70.852 184.259 Int3 1 1 250 250 0.008 0.027 0.063 33.291 107.923 253.801 21.009 500 500 0.011 0.030 0.065 Tnt4 1 1 59.128 129.232 1500 1500 0.000 0.000 0.009 0.000 0.000 Int5 1 1 5.716 Tnt.6 500 500 0.931 1.499 2.362 1862.604 2997.521 4723.249

Two sets of confidence intervals are computed: one for the %-recombination frequency, the other for the standard measure of recombination, centi-Morgans per megabase (cM/Mb). These are obtained from the %-frequencies as

$$\mbox{cM/Mb} = \frac{\mbox{\%-Freq}}{\mbox{Len}} \cdot 10^6. \label{eq:cm/mb}$$

4.4 six

six simulates typing data given some crossover rate. This can be used for checking the accuracy of xov, and for testing assay designs.

• Run six

```
six
# Int Chr Start
                       End
                                  m|n-k|k m|n-k|k m|n-k|k
           15,000,001 15,002,000 60|8|4
                                          120 | 10 | 2 240 | 12 | 0
Intl 1
Int2 1
           15,000,001 15,002,000 60|7|5
                                          120|9|3
                                                   240|11|1
Int3 1
           15,000,001 15,002,000 60|8|4
                                          120|7|5
                                                  240|12|0
Int4 1
           15,000,001 15,002,000 60|7|5
                                          120|10|2 240|11|1
Int5 1
           15,000,001 15,002,000
                                  60|5|7
                                          120|8|4
                                                   2401913
```

where the columns have the same meaning as explained above for exampleResults.txt.

• Pipe the results of six through xov

```
six | xov
# Int Chr Start
                                      용
                                               %] [cM/Mb cM/Mb
                   End
                            Len [%
                                                                   cM/Mbl
     1
           15000001 15002000 2000 0.542 0.869 1.332 270.847 434.393 665.759
Int1
           15000001 15002000 2000 0.895 1.404 2.142 447.448 701.838 1070.429
Int2
      1
          15000001 15002000 2000 0.688 1.083 1.645 343.656 541.430 822.117
Int3 1
Int4 1 15000001 15002000 2000 0.534 0.855 1.307 266.739 427.079 652.990
           15000001 15002000 2000 0.905 1.447 2.259 452.265 723.175 1129.031
Int5 1
```

• Check accuracy of xov

The simulated %-crossover frequency (-x) is 1.5, but the average estimate is 1.59. In other words, our estimator has an upward bias [3].

References

- [1] H. Li. Tabix: fast retrieval of sequence features from generic TAB-delimited files. *Bioinformatics*, 27:718–719, 2010.
- [2] H. Li, B. Handsaker, A. Wysoker, T. Fennell, J. Ruan, N. Homer, G. Marth, G. Abecasis, R. Durbin, and 1000 Genome Project Data Processing Subgroup. The sequence alignment/map format and SAMtools. *Bioinformatics*, 25:2078–2079, 2009.
- [3] L. Odenthal-Hesse, J. Y. Dutheil, F. Klötzl, and B. Haubold. hotspot: Software to support sperm-typing for investigating recombination hotspots. *Bioinformatics*, 32:2554–2555, 2016.