sfs: Compute Site Frequency Spectra

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1 Introduction

For a sample of size n let $f_i(n)$ be the number of sites where i individuals carry a mutation. The vector

$$f_1(n), f_2(n), ..., f_{n-1}(n)$$

is called the *site frequency spectrum*. The program sfs takes multiple haplotype samples simulated with ms as input and prints the average site frequency spectrum.

2 Getting Started

sfs was written in C on a computer running Linux and should work on any standard UNIX system. However, please contact me at haubold@evolbio.mpg.de if you have any problems with the program.

• Change into the program's directory

cd sfs

and list its contents

ls

• Generate sfs

make

• List its options

```
./sfs -h
```

• Test the program on a data set consisting of 2 simulated haplotype samples:

```
./sfs data/msOdd.dat
```

• Compare to folded SFS

```
./sfs -f data/msOdd.dat
```

• Repeat for an even-numbered sample size

```
./sfs data/msEven.dat
./sfs -f data/msEven.dat
```

• Apply sfs to 10⁴ newly simulated haplotypes

```
ms 10 10000 -t 10 \mid ./sfs
```

where ms is the coalescent simulator by Hudson (2002).

3 Listing

The following listing documents the driver program for sfs.

```
/**** sfs.c ***********************
   * Description:
   * Author: Bernhard Haubold, haubold@evolbio.mpg.de
   * Date: Wed Sep 20 07:50:43 2017
   ****************
6 #include <stdio.h>
  #include <stdlib.h>
  #include "interface.h"
  #include "eprintf.h"
  #include "sample.h"
#include "spectrum.h"
  void scanFile(FILE *fp, Args *args) {
    Sample *sa;
    Spectrum *sp;
    int i, n;
    sa = initializeSample(fp, args);
    sp = newSpectrum(sa->nsam);
    n = 0;
    while((sa = getSample(0)) != NULL) {
      sp = computeSpectrum(sa, sp);
      n++;
    if(!args->r){
      for (i=0; i<sp->n; i++)
26
        sp->spectrum[i] /= (double)n;
    if(args->f)
      foldSpectrum(sp);
    printSpectrum(sp);
    freeSample();
  int main(int argc, char *argv[]){
    int i;
    char *version;
    Args *args;
    FILE *fp;
    Spectrum *sp;
41
    version = "0.7";
    setprogname2("sfs");
    args = getArgs(argc, argv);
    if (args->v)
      printSplash(version);
    if(args->h || args->e)
      printUsage(version);
    if (args->t) {
      sp = getArtificialSpectrum(args);
      if (args->f)
```

```
foldSpectrum(sp);
   printSpectrum(sp);
   return 0;
}

if (args->numInputFiles == 0) {
    fp = stdin;
       scanFile(fp, args);
}else{
    for(i=0;i<args->numInputFiles;i++) {
       fp = efopen(args->inputFiles[i],"r");
       scanFile(fp, args);
       fclose(fp);
    }
}

free(args);
free(progname());
return 0;
```

4 Change Log

- Version 0.1 (September 25, 2017)
 - First working version.
- Version 0.2 (October 23, 2017)
 - Polished interface.
- Version 0.3 (November 17, 2017)
 - Implemented folding of SFS (−F).
- Version 0.4 (November 29, 2017)
 - Enable analytic computation of SFS (-T to specify θ and -n to specify sample size.
- Version 0.5 (December 1, 2017)
 - Fixed error in folding.
- Version 0.6 (December 18, 2017)
 - Cleaned up interface.
- Version 0.7
 - Implemented -r for printing raw counts.
- June 13, 2018
 - Posted on github.

References

R. R. Hudson. Generating samples under a Wright-Fisher neutral model of genetic variation. *Bioinformatics*, 18: 337–338, 2002.