### Manual for *HierAdmix*

#### 1. Introduction

*HierAdmix* is a software that can be used to reconstruct the 4-way admixture history considering the complex admixture structure. It could automatically select the optimal admixture model between the hierarchical admixture model and the sequential admixture model. *HierAdmix* is an extension of the *MultiWaver* series software.

## 2. Input Files

The format of input ancestral tracts file is similar to *MultiWaver* series software but with two additional columns. Each line represents an ancestral tract with the genetic distance of start point, the genetic distance of end point, from which ancestry the tract originates, the index of admixed haplotype, the haplotype label, and the chromosome label. The unit of first two columns are both in Morgan. Here takes the first few rows of "Hier.seg" as an example:

0	0.01113699	Anc3	hap0	1
0.01113699	0.02255031	Anc2	hap0	1
0.02255031	0.02763909	Anc3	hap0	1
0.02763909	0.03211435	Anc4	hap0	1
0.03211435	0.08753307	Anc3	hap0	1
0.08753307	0.09612572	Anc4	hap0	1
•••••	•••••		•••••	

### 3. Output Files

The format of output model file is same as *MultiWaver* series software. Here takes the output file "Hier.txt" as an example, which is inferred based on the input file "Hier.seg":

```
Best Model:
               Hierarchical Admixture Model
     Anc1
               143(G)
                         0.208525
     Anc2
               143(G)
                         0.791475
     Anc3
               147(G)
                         0.700884
     Anc4
               147(G)
                         0.299116
     Anc1 Anc2 50(G)
                         0.398402
     Anc3_Anc4 50(G)
                         0.601598
Bootstrapping details
Bootstrapping support ratio:100.0% (100/100)
               [137, 147](G) [0.20302, 0.214607]
     Anc1
     Anc2
               [137, 147](G)
                             [0.785393, 0.79698]
     Anc3
               [144, 149](G) [0.694946, 0.706197]
               [144, 149](G) [0.293803, 0.305054]
     Anc4
     Anc1 Anc2 [48, 51](G)
                               [0.389099, 0.40447]
     Anc3_Anc4 [48, 51](G)
                               [0.59553, 0.610901]
```

# 4. Arguments and Options

Arguments	Type	Description	Note
input	string	Input file name	required
lower	float	Lower bound to discard short tracts	optional, default = 0
bootstrap	int	Number of bootstrapping	optional, default = 0
ci	float	Confidence level of bootstrapping confidence interval	optional, default = 0.95
output	string	Prefix of output file	optional, default="out"
-h	/	Print help message	/

### 5. Notes

A special hierarchical admixture model needs special attention, within which the admixture proportions of two populations of left-side are same, and the admixture proportions of two populations of right-side are same too. In this case, the admixture time derived from the length distribution of ancestral tracts doesn't have the unique solution and the estimated time of *HierAdmix* can be unreliable.

## 6. Questions and Trouble-shooting

Questions and suggestions are welcome, feel free to contact Shi Zhang 19121640@bjtu.edu.cn or Rui Zhang zhangrui2018@picb.ac.cn.