# Haplotype phasing using EM & divide and conquer

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# The application

We developed a lightweight software for haplotype phasing, consisting of two scripts: main.py. the main interface and phase.py, containing the implementations. Usage of the scripts are described in readme.txt.

# Methodology

EM is widely used for haplotype phasing; however it has poor scalability because its computational time increases exponentially with the length of input genotype sequence. To phase arbitrarily long genotypes, we divided the input genotypes (sequences of  $\{0,1,2\}$ ) into atomic l-length segments, apply EM to each of the l-length segments, and finally merge the individually phased segments together in a reasonable way.

#### $\mathbf{EM}$

We extend the EM algorithm such that it deals with individuals coming from an admixture of populations. Let  $p_{hk}$  be the frequency of haplotype h in population k,  $q_k$  be the probability that a randomly chosen haplotype is from population k. Let  $C(x_i)$  be the set of all compatible haplotypes with genotype  $x_i$ . Starting with an initial guess of p and q as  $p^0$ ,  $q^0$ , repeatedly update with:

$$\begin{split} a_{i,k_1,k_2,h_1,h_2}^t &= \frac{q_{k_1}^t q_{k_2}^t p_{h_1 k_1}^t p_{h_2 k_2}^t}{\sum_{k,k'} \sum_{h,h' \in C(x_i)} q_{k}^t q_{k'}^t p_{hk}^t p_{h'k'}^t} \\ q_k^{t+1} &= \text{Sum of all } a \text{ with at least } 1 \ k \\ &\text{in subscript} \end{split}$$

 $p_{hk}^{t+1} = \text{Sum of all } a \text{ with at least } 1 \text{ } k$ and at least 1 h in subscript p and q are normalized such that all q sum to 1 and all p for a particular k sum to 1. After convergence, the most probable phase (and population for each haplotype) is derived.

### Merging

After the phase for each of the two neighboring segments A and B are found, an additional round of EM is applied to a w-length window C centering between A and B. This will determine whether the two segments are relatively trans- or cis-phased. For example if w=6, A=212210 and B=112200, we attempt to find the haplotype frequencies within the window 210112. If we have already phased A to be 101110, 101100 and B to be 111100, 001100, and by doing EM on C we discovered that the the pair 100111, 110001 has higher likelihood than 110111, 100001, we can assume that the more probable phase for AB is 101110001100, 101100111100.

### Additional notes

To account for the fact that this merging method is greedy and that the optimal phase for a segment may not be in the optimal phase for the full genotype, a command line option allows the program to save the top k phases for a segment, and during merging these phases are re-ranked based how how likely they appear in the window C. Experiments suggest that doing so consistently improves switch accuracy. In addition, the initial guess for q is randomized but the user can input a custom guess. As a result with multiple populations the performance is highly dependent on this initial guess. Therefore, using only 1 population is recommended.