**Introduction**

PONDEROSA is an algorithm designed to assist in pedigree construction. PONDEROSA works well in datasets with high-quality long-range phasing. We have found that this can be better achieved in endogamous populations. Even in datasets with poor phasing, PONDEROSA can still distinguish avuncular from grandparent-grandchildren and maternal half-siblings from paternal half-siblings and will work well in datasets with few half-siblings or datasets with a narrow age range (which may be unlikely to have avuncular or grandparent-grandchildren pairs). PONDEROSA works best in datasets with existing pedigree structure, which is necessary for training the machine-learning classifiers. PONDEROSA will work out this existing pedigree structure from tracing parent-offspring lineages; therefore, every parent-offspring pair as inferred by KING must be present in the .fam file.

Please note that PONDEROSA is designed to *assist* pedigree construction and further steps by the user are required to construct the pedigree. We hope to change this in future versions of PONDEROSA. For now, PONDEROSA largely infers relationships in a vacuum (i.e. without considering the context of the pedigree). Relationship inference should be double-checked against the existing pedigree structure.

**Requirements**

**python3.6**

**scikit-learn** and its dependencies (install here: https://scikit-learn.org/stable/install.html). Alternatively, we recommend running python directly from anaconda3, which has all the packages needed to run PONDEROSA.

**Input**

*Required arguments*

**--king** KING .seg file (or any .seg-formatted IBD file).

**--map** PLINK-formatted .map files. The chromosome number should be replaced with “%s”. Note that PONDEROSA expects a .map file for each chromosome but only one **--map** flag (see example script). The .map file must be the same .map file used to generate IBD segments.

**--fam** PLINK-formatted .fam file. All PO present in the KING file must be present in the .fam file. If age data is unavailable/unreliable and the parent/offspring cannot be distinguished in the pair, PONDEROSA can be run and the haplotype scores of the individuals can be used to make the distinction (and then run again).

**--match** GERMLINE-formatted match file where the chromosome number is replaced with “%s”. Again, PONDEROSA expects a .match file for each chromosome but only one **--match** flag (see example script). If GERMLINE file, must be generated with GERMLINE’s **--haploid** flag (we suggest GERMLINE v1.5.3). iLASH .match files can also be used, but PONDEROSA’s **--ilash** flag must be used.

*Optional arguments*

**--out** Output file prefix. *Default: “PONDEROSA”*

**--chr** Number of autosomes. Change only for non-human samples. *Default: 22*

**--ilash** For use if .match file is in iLASH format.

**--haps** If PONDEROSA has already been run, supplying the haplotype score file

will skip the haplotype score calculation step.

**--age** Age file where the first column corresponds to the individual ID and the second column corresponds to the age. Note that not all individuals need an age.

**--gp\_gap** Minimum age-gap for a grandparent-grandchild pair. Note that if you do not want PONDEROSA to consider age, use 0 for this flag. *Default: 30*

**­--mhs\_gap** Maximum age-gap for maternal half-siblings. If you do not want PONDEROSA to consider age here, use an arbitrarily large age gap (e.g. 100). *Default: 30*

**--po\_gap** Minimum age-gap for parent-offspring. If you do not want PONDEROSA to consider age here, use 0 for this flag. *Default: 15*

**--ped** PLINK-formatted .ped file used by PONDEROSA to stitch IBD segments together. If no .ped file is supplied, PONDEROSA stitches together two segments that are within 1 cM of each other. If .ped file is supplied, PONDEROSA only stitches two segments that are within 1 cM (can be changed with **--cm\_gap** flag) of each other *and* have, at most, one discordant homozygote (can be changed with **--disc\_homoz** flag).

**--cm\_gap**  Maximum gap in cM between IBD segments for them to be considered a single segment (see **--ped** flag for more detail). *Default: 1*

**--disc\_homoz** Maximum number of discordant homozygotes between two IBD segments in order for them to be considered the same IBD segment. Only use if **--ped** file is used. *Default: 1*

**--likelihood** Minimum likelihood (0.5 - 1) required for a pair to be inferred as a 2nd degree pair. We recommend being more conservative here. *Default: 0.80*

**Output**

PONDEROSA outputs five files.

**.pedigree file**

Outputs every pairwise relationship found and the confidence. For inferred 2nd degree pairs, the confidence reported is P(2nd degree relationship | 2nd degree). Note that for each of these pairs, the P(2nd degree) > 0.80 under default parameters, but can be changed using **–likelihood** flag. The confidence reported for these 2nd degree relatives is a posterior probability, calculated after considering existing pedigree structure and age discrepancies (note: see the age parameters for instructions on how to ignore age); for the probabilities of each relationship class before this step (i.e. the probabilities calculated on the haplotype scores and number of IBD segments alone), refer to the .score file. Each line corresponds to a pair with the following fields:

1. Individual ID 1 (child in PO pair, grandchild in GP pair, niece/nephew in AV pair)
2. Individual ID 2 (parent in PO pair, grandparent in GP pair, aunt/uncle in AV pair)
3. Pedigree relationship
4. Confidence

**.log file**

Provides information about runtime, number of training relationships, and any warnings about inferred relationships. PONDEROSA assumes that all reported parent-offspring are true, but will output warnings when purported PO have IBD1 < 0.80. PONDEROSA will also report errors when FS or 2nd degree pairs have IBD1 < 0.30. Typically, these pairs have low IBD1 values because of an error with the reported PO. These should be double-checked, the .fam file should be updated, and PONDEROSA rerun.

**.training file**

Gives the haplotype scores, haplotype score ratios, and the number of IBD segments for each known second degree pair. This file is useful for assessing whether or not the phase quality of the dataset is high (e.g. by plotting). Each line corresponds to a pair with the following fields:

1. Individual ID 1 (child in PO pair, grandchild in GP pair, niece/nephew in AV pair)
2. IID1’s haplotype score
3. Individual ID 2 (parent in PO pair, grandparent in GP pair, aunt/uncle in AV pair)
4. IID2’s haplotype score
5. Haplotype score ratio
6. Number of IBD segments
7. Relationship

**.score file**

Provides information about each putative second degree relative. Each line corresponds to a pair with the following fields:

1. Individual ID 1 (putative grandchild/niece/nephew)
2. IID1’s haplotype score
3. Individual ID 2 (putative grandparent/aunt/uncle)
4. IID2’s haplotype score
5. Haplotype score ratio
6. Number of IBD segments
7. P(PHS)
8. P(MHS)
9. P(GP)
10. P(AV)

**.fam file**

As of v1.2, the only difference between this newly generated fam file and the user-supplied fam file is that full-siblings are assigned dummy parents if they are missing one or both parents. In future versions, we hope to assign dummy parents to all non-founder samples missing one or both parents.

**Pipeline**

Scripts for the pipeline and sample data are available on Github. Note that the data provided is simulated data of west African descent.