Objects

```
• ped: class Pedigree
     ▶ ped.count: int, #individuals
    ▶ ped.markerCount: int, #markers
     ▶ ped.markerNames: StringArray, marker names
• ped[i]: class Person
     ▶ ped[i].famid/pid/fatid/motid: String, pedigree info
    ▶ ped[i].sex: int, sex
    ▶ ped[i].traits: float*, traits
     ▶ ped[i].markers: class Alleles*
    ▶ ped[i].markers[j][0]/[1]: int, allele
    ▶ ped[i].markers[j].one/two: int, allele, equivalent to item above
    ▶ ped[i].father/mother: class Person*
    ped[i].sibs: class Person**
     ▶ ped[i].sibCount: int, sib count
• ped.GetMarkerInfo(j): class MarkerInfo
     ▶ ped.GetMarkerInfo(j).CountAlleles(): int, #alleles
    ▶ ped.GetMarkerInfo(j).freq[k]: float, allele frequency for allele k
     ▶ ped.GetMarkerInfo(j).GetAlleleLabel(k): String, allele representation
```

Methods

• ped.EstimateFrequencies(1, true): method, calculate allele frequency based on founders

Example

```
CPP
#include <iostream>
#include "Core.hpp"
using namespace SEQLinco;
void showPed(Pedigree & ped)
{
        printf("Loaded %d individuals\n", ped.count);
        for (int i = 0; i < std::min(ped.count, 10); i++) {
                printf("[%s]: %s, %s, %s, %d\t|\t",
                        (const char *)ped[i].pid, (const char *)ped[i].famid,
                        (const char *)ped[i].fatid, (const char *)ped[i].motid,
                        ped[i].sex);
                for (int j = 0; j < ped.markerCount; ++j) {</pre>
                        printf("%d%d\t", ped[i].markers[j].one, ped[i].markers[j].two);
                printf("\n");
        printf("Loaded %d markers\n", ped.markerCount);
        // Estimate allele frequencies for all markers, verbose mode
        ped.EstimateFrequencies(1, false);
        // Get genotype statistics for markers
        for (int i = 0; i < ped.markerNames.Length(); i++) {</pre>
                printf("Statistics for marker [%s]\n", (const char *)ped.markerNames[i]);
```

```
// Allele index starts with 1 not 0
                            for (int j = 1; j <= ped.GetMarkerInfo(i)->CountAlleles(); j++) {
                                          printf("\tFrequency for allele %d: %f\n", j, ped.GetMarkerInfo(i)->freq[j]);
                                          printf("\tName for allele %d: %s\n", j, (const char *)ped.GetMarkerInfo(i)->GetAlleleLabel(j));
                           }
             }
              return;
void readData(Pedigree & ped,
                        const char * datfile, const char * pedfile, const char * mapfile)
{
              // The data file contains a description of the contents of the
              // pedigree file, including for example, a list of marker and
              // trait names
              ped.Prepare(datfile);
              // The pedigree file contains a list of individuals, stored one
              // per row, with specific information about each individual as
              // detailed in the data file.
              ped.Load(pedfile);
              SortFamilies(ped);
              ped.LoadMarkerMap(mapfile);
             return:
}
int main(int argc, char ** argv)
              if (argc != 3 && argc != 4) {
                            printf("usage: \space{0.05cm} \space{0.05cm} source code: 1, 2, 3> \space{0.05cm} \space{0.05cm} \space{0.05cm} source code: 1, 2, 3> \space{0.05cm} \spac
                            return 0;
              }
              PedigreeData ped;
              std::string chrom = "1";
              if (atoi(argv[1]) == 1) readData(ped.data, "haplo.dat", "haplo.ped", "haplo.map");
              else if (atoi(argv[1]) == 2) readData(ped.data, "gene.dat", "gene.ped", "gene.map");
              else if (atoi(argv[1]) == 3) {
                            std::vector<std::string> marker_ids { "V1", "V2", "V3" };
                            {\tt std::vector < std::string > marker\_positions \ \{ \ "1", \ "2", \ "3" \ \};}
                            std::vector< std::vector<std::string> > samples;
                            std::vector<std::string> s0 { "1", "1", "0", "0", "1", "21", "21", "21" };
                            samples.push_back(s0);
                            std::vector<std::string> s1 { "1", "2", "0", "0", "2", "11", "11", "11" };
                            samples.push_back(s1);
                            std::vector<std::string> s2 { "1", "3", "1", "2", "1", "21", "21", "21" };
                            samples.push_back(s2);
                            std::vector<std::string> s3 { "2", "1", "0", "0", "1", "22", "21", "00" };
                            samples.push_back(s3);
                            std::vector<std::string> s4 { "2", "2", "0", "0", "2", "11", "11", "11" };
                            samples.push_back(s4);
                            std::vector<std::string> s5 { "2", "3", "1", "2", "1", "21", "21", "21" };
                            samples.push_back(s5);
                            std::vector<std::string> s6 { "3", "1", "0", "0", "1", "22", "21", "21" };
                            samples.push back(s6):
                            std::vector<std::string> s7 { "3", "2", "0", "0", "2", "11", "11", "21" };
                            samples.push_back(s7);
                            std::vector<std::string> s8 { "3", "3", "1", "2", "1", "21", "21", "21" };
                            samples.push_back(s8);
                           ped.LoadVariants(marker_ids, marker_positions, chrom);
                           ped.LoadSamples(samples);
              } else ;
              if (atoi(argv[2]) == 1) showPed(ped.data);
              else if (atoi(argv[2]) == 2) {
                            MendelianErrorChecker mc;
                           mc.Apply(ped.data);
                            std::cout << "Mendelian Errors " << mc.errorCount << std::endl;
                            GeneticHaplotyper gh(chrom);
                            gh.Apply(ped.data);
                            if (argc == 4) {
                                          for (unsigned f = 0; f < gh.data.size(); f++) {</pre>
                                                       for (unsigned p = 0; p < gh.data[f].size(); p++) {</pre>
                                                                     for (unsigned i = 0; i < gh.data[f][p].size(); i++) {</pre>
                                                                                   std::cout << gh.data[f][p][i] << "\t";
```

```
std::cout << std::endl;
}
std::cout << std::endl;
}

HaplotypeCoder hc(1);
hc.Apply(gh.data);
if (argc == 4) {
    for (unsigned p = 0; p < hc.data.size(); p++) {
        for (unsigned i = 0; i < hc.data[p].size(); i++) {
            std::cout << hc.data[p][i] << "\t";
        }

std::cout << std::endl;
}
}
else;
}</pre>
```

```
CXX = g++
CFLAGS=-03 -I ./ -I./libsrc -I./merlin -I./pdf -I./clusters -D_FILE_OFFSET_BITS=64 -D__ZLIB_AVAILABLE__ -Wall -std=c++11
BINDIR = demo
MERLIN = demo/demo.exe
EXECUTABLES = $(MERLIN)
# MERLIN File Set
MERLINBASE = merlin/AssociationAnalysis merlin/FastAssociation \
 merlin/AnalysisTask merlin/Conquer \
 {\tt merlin/ConquerHaplotyping\ merlin/DiseaseModel\ } \setminus
 merlin/ParametricLikelihood merlin/GenotypeInference \
 merlin/Houdini \
 {\tt merlin/KongAndCox\ merlin/Manners\ merlin/MerlinBitSet}\ \setminus
 merlin/MerlinCluster merlin/MerlinCore \
 merlin/MerlinError merlin/MerlinFamily merlin/MerlinIBD \
 merlin/InformationContent merlin/MerlinCache merlin/MerlinModel \
 merlin/MerlinKinship merlin/MerlinKinship15 \
 merlin/MerlinHaplotype merlin/MerlinMatrix merlin/MerlinParameters \
 {\tt merlin/MerlinPDF\ merlin/MerlinSimulator\ merlin/MerlinSimwalk2\ } \setminus
 merlin/MerlinSort merlin/NPL-ASP merlin/NPL-QTL \
 merlin/Magic merlin/Mantra merlin/Parametric merlin/QtlModel \
 merlin/Tree \
 merlin/TreeBasics merlin/TreeIndex merlin/TreeManager \
 merlin/TreeInfo merlin/TreeFlips merlin/VarianceComponents
MERLINHDR = $(MERLINBASE:=.h) merlin/TreeNode.h Core.hpp
MERLINSRC = $(MERLINBASE:=.cpp) demo/LibmerlinDemo.cpp Core.cpp
MERLINOBJ = $(MERLINSRC:.cpp=.o)
# Files for dealing with clustered markers
CLUSTERS = clusters/HaploFamily clusters/HaploGraph \
 clusters/HaploSet clusters/HaploTree clusters/Likelihood \
 clusters/SparseLikelihood clusters/Unknown
CLUSTERCPP = $(CLUSTERS:=.cpp)
CLUSTERHDR = $(CLUSTERS:=.h)
CLUSTEROBJ = $(CLUSTERS:=.o)
# Utility Library File Set
LIBFILE = libsrc/lib-goncalo.a
LIBMAIN = libsrc/BasicHash libsrc/Error libsrc/FortranFormat \
 libsrc/GenotypeLists libsrc/InputFile libsrc/IntArray libsrc/Hash \
 libsrc/LongArray libsrc/Kinship libsrc/KinshipX libsrc/MapFunction \
 libsrc/MathCholesky libsrc/MathDeriv libsrc/MathFloatVector \
 libsrc/MathGenMin libsrc/MathGold libsrc/MathMatrix libsrc/MathStats \
 libsrc/MathNormal libsrc/MathSVD libsrc/MathVector \
 libsrc/MemoryInfo libsrc/MiniDeflate \
 libsrc/Parameters libsrc/Pedigree libsrc/PedigreeAlleleFreq \
 libsrc/PedigreeDescription libsrc/PedigreeFamily libsrc/PedigreeGlobals \
 libsrc/PedigreePerson libsrc/QuickIndex libsrc/Random libsrc/Sort \
 libsrc/StringArray libsrc/StringBasics libsrc/StringMap \
 libsrc/StringHash libsrc/TraitTransformations
LIBPED = libsrc/PedigreeLoader libsrc/PedigreeTwin libsrc/PedigreeTrim
LIBSRC = $(LIBMAIN:=.cpp) $(LIBPED:=.cpp)
LIBHDR = $(LIBMAIN:=.h) libsrc/Constant.h \
libsrc/MathConstant.h libsrc/PedigreeAlleles.h libsrc/LongInt.h
LIBOBJ = $(LIBSRC:.cpp=.o)
# PDF Library File Sets
```

Makefile

PDFLIB = pdf/libpdf.a

```
PDFFILES = pdf/PDF pdf/PDFfont pdf/PDFinfo pdf/PDFpage \
pdf/PDFchartbasics pdf/PDFchartbar pdf/PDFlinechart \
pdf/PDFhistogram \
pdf/PDFchartaxis pdf/PDFchartlegend pdf/PDFchartmarker \
pdf/PDFchartline pdf/PDFchartobject
PDFSRC = $(PDFFILES:=.cpp)
PDFHDR = $(PDFFILES:=.h)
PDFOBJ = $(PDFFILES:=.o)
# make everything
all : $(EXECUTABLES)
$(EXECUTABLES) : $(BINDIR)
$(BINDIR) :
       mkdir -p $(BINDIR)
# dependencies for executables
$(MERLIN) : $(LIBFILE) $(PDFLIB) $(MERLINOBJ) $(CLUSTEROBJ)
       $(LIBFILE) : $(LIBOBJ) $(LIBHDR)
       ar -cr $@ $(LIBOBJ)
       ranlib $@
$(PDFLIB) : $(PDFOBJ)
      ar -cr $@ $(PDFOBJ)
       ranlib $@
$(MERLINOBJ) : $(MERLINHDR) $(CLUSTERHDR) $(LIBHDR)
$(CLUSTEROBJ) : $(CLUSTERHDR) $(MERLINHDR) $(LIBHDR)
$(LIBOBJ) : $(LIBHDR)
$(PDFOBJ) : $(PDFHDR)
clean :
       rm -f */*.a */*.o $(EXECUTABLES)
.c.o :
       $(CXX) $(CFLAGS) -o $@ -c $*.c
#.cpp.X.o :
       $(CXX) $(CFLAGS) -o $0 -c $*.cpp -D__CHROMOSOME_X_
.cpp.o :
       $(CXX) $(CFLAGS) -o $@ -c $*.cpp
.SUFFIXES : .cpp .c .o .X.o $(SUFFIXES)
```