mitoGenome Assembly

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Install the software

- 1. MITObim¹: https://github.com/chrishah/MITObim
 - make sure to follow install instructions, including dependecies: MIRA, PERL, etc.
- 2. MUSCLE²: https://www.drive5.com/muscle/
- 3. MAFFT³: https://mafft.cbrc.jp/alignment/software/

If you're on a mac, downloading this repository will include MUSCLE and MAFFT suited for a mac, so you can plug 'n' play. If you're on a Windows or Linux machine, you'll have to do this separately, and adjust the call to the programs in 'mitoAlign.R'. Sorry, I'm just too dumb.

¹Hahn, C., Bachmann, L., Chevreux, B. Reconstructing mitochondrial genomes directly from genomic next-generation sequencing - a baiting and iterative mapping approach. Nucleic Acids Research 41:13. doi:10.1093/nar/gkt371 (2013).

²Edgar, R.C. MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Research 32(5). doi:10.1093/nar/gkh340 (2004).

³Katoh, Standley. MAFFT multiple sequence alignment software version 7: improvements in performance and usability. Molecular Biology and Evolution 30:772-780 (2013).

Multiple mtGenome assembly/alignment using MITObim from R

Why Bother?

As a result of the exon capture process for Anchored Hybrid Enrichment projects, we're getting a considerable amount of mitochondrial bycatch in our raw reads. This mtDNA can provide a lot of information including a separate phylogenetic history for the group of interest, or even past introgression events. Also, :: free data::.

File Preparation

We'll use the MITObim pipeline, run from R to assemble our mtGenomes against a reference (find one on GenBank, distantly related is OK). We'll then pull out the final assembly for each sample and align them against each other using MUSCLE.

Start by creating a directory to hold:

- the MITObim pipeline ('MITObim.pl')
- the MIRA sequence assembler and mapper directory ('mira.4.0.2...')
- the MUSCLE executable ('muscle3.8.31...')
- the MAFFT folder ('mafft-mac')
- a project-specific directory holding all your mtDNA reads for each sample in subdirectories ('Frogs'), and your reference mtGenome as a fasta file, labelled '[taxon]_mtGenome.fasta'

Here's a quick schematic of what the structure should look like:

```
/PATH_TO_PARENT_DIRECTORY/mtGenomes
|-- MITObim.pl
|-- mira_4.0.2_darwin13.1.0_x86_64_static
| |-- bin (et al.)
|-- muscle3.8.31_i86darwin64
|-- Frogs
|-- [taxon]_mtGenome.fasta (reference genome)
|-- Taxon1
| |-- Taxon1_R1.fastq.gz
| |-- Taxon2_R1.fastq.gz
|-- Taxon2_R1.fastq.gz
|-- Taxon2_R2.fastq.gz
```

Executing the Code

Open a new R script, and source the functions we'll need

```
source("/PATH_TO_CODE/mtGenome_Assembly.R")
```

Then set your working directory to the project-specific directory (I've called this directory 'Frogs'). This holds a subdirectory for each sample, and your reference mitogenome.

```
setwd("/PATH_TO_DIR/mtGenomes/Frogs")
```

The first function (*mitoAssemble*) assumes that in the directory 'Frogs', there are one or more subdirectories (1 per sample), with one or more gzipped fastq files of the raw reads. For each sample (subdirectory), it will:

- merge (concatenate) any *fastq.gz files together.
- copy your reference genome (fasta file)
- call MITObim and attempt to assemble the mtGenome
- $\bullet\,$ copy the assembled mtGenome to a directory of all assemblies

The function requires as input, three things:

```
mitoAssemble(num.iter, reference.name, project.name)
```

- num.iter is the number of assembly iterations MITObim should try before it times out and moves on to the next sample. I usually leave this set to 100, sometimes it takes 4 iterations, sometimes 60, hard to know.
- reference.name is the unique name of your reference genome fasta file which precedes '_mtGenome.fasta'
- project.name is what you'd like the generated directories to be called (where the assemblies are stored)

The function *mitoAssemble* will spit out all the assemblies to a new directory, and tell you where it is:

```
Assembly(s) completed and saved to: /Users/Ian/MITObim/Assa/Assa_mtGenomes
```

The second function mitoAlign will:

- combine all assembled mtGenomes into a single fasta alignment ([project.name]_Assembly_Alignment.fasta)
- $\bullet \ \ a lign \ the \ assemblies \ using \ MUSCLE \ or \ MAFFT, into \ a single \ final \ a lignment \ ([project.name_Aligned_Assemblies.fasta])$

The function mitoAlign will spit out the final alignment into your new directory, and tell you where it is:

```
your alignment of mtGenome assemblies is called:
/Users/Ian/MITObim/Assa/Assa_mtGenomes/Assa_Aligned_Assemblies.fasta
```

The function requires little information to do its job well. Bare minimum it should be given the project name, and your choice of aligner ("MUSCLE" or "MAFFT"). You can also designate the reference genome for aligning purposes:

```
mitoAlign(project.name, "MUSCLE", reference.name)
```

• project.name what you've been calling the project, so it can find the folder with all your mitoGenome assemblies.

• reference.name if you'd like to improve your multi-mitoGenome alignment, you can align them back to your reference genome. By default, this is NULL, to specify it, just give it the name you provided in the mitoAssemble step.

At the moment, I'd probably suggest using the MAFFT aligner with the reference option if possible. Aligning the assembled mitogenomes without a reference can result in some poor behavior from poorly assembled genomes.

The third function *mitoCheck* will:

- read in your alignment, and provide information about missing/gaps in your data
- remove taxa with low amounts of sequence if you'd like

You just have to give it the alignment and a little info:

```
mitoCheck(project.name, alignment, count.gaps=T, missing.threshold=NULL)
```

- project.name is again, just what you've been calling the project so far (name of the directory)
- alignment is the name of the output .fasta file from the mitoAlign step
- count.gaps is just a TRUE/FALSE statement about whether to count gaps as missing data
- missing.threshold if you want to remove taxa that exceed a certain percentage of missing data, indicate that here (0 < x < 1)

If you do decide to trim taxa based on missing data, the function will produce a new alignment and tell you the name:

```
your reduced alignment is now called:
/Users/Ian/MITObim/Assa/Assa_mtGenomes/Assa_Aligned_Assemblies_Reduced.fasta
```

I thought about using a different metric such as base composition/content, but I haven't thought about it enough to implement it well. Instead, it always identifies the outgroups as being funky, so I've just stuck with missing data.

The last function *mitoChop* will:

- take the whole mitoGenome alignment and split it up by locus
- this function requires the R packages ape and seqinr to read and write the files.

The function will kick out the alignment for each mitochondrial locus (CDS/gene, rRNA, tRNA, et al.) separately into a new folder:

```
your separated mitochondrial loci alignments are in a folder called::
/Users/Ian/MITObim/Assa/Assa_mtGenomes/Assa_mitoLoci
```

You just have to provide a character set file as a .CSV so it knows where to cut it up

```
mitoChop(project.name, alignment, character.sets=NULL)
```

- project.name is again, just what you've been calling the project so far (name of the directory)
- alignment the name of the output .fasta file from the mitoAlign step

-	is name, the starting position (<i>Minimum</i>), and final position (<i>Maximum</i>) for each included in the respository ("mitoGenome Annotations Example.csv").
ocus. The example is	meraded in the respository (introductions_transference).
- v	of other slick things I could do after this, like have it plot the individual assembly it's worth it at the moment. Let me know if there's something specific you're
Good luck!	
-	

 \bullet character.sets is the .CSV file with three columns "Name", "Minimum", and "Maximum". These