# Mitochondrial molecular markers for US lineages of $P.\ infestans$

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## $_{\scriptscriptstyle 1}$ 1 The sample

- 2 The sample includes data which was opportunistically gathered from previous
- 3 publications as well as data which is not yet available to the public (Judelson,
- unpublished).

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Sample	Count	Reference
T30-4	1	[4]
PIC99189 & 90128	2	[9]
13_a2	1	[2]
Yoshida et al.	13	[10]
Martin et al.	3	[7]
$\operatorname{Judelson}$	8	ŇĀ
Total	28	

- The sample 'T30-4' was the first sequenced genome and is considered the reference for nuclear work [4]. This genome was assembled prior to high-throughput sequencing (i.e., Illumina and 454 technologies). The data presented here are not the sequences used for the paper but are part of a project by The Broad[8] to resequence this individual using Illumina and Roche/454 technologies.
- Note that both the Yoshida and Martin papers included ancient DNA in their analyses [10, 7]. Here we have omitted those samples and focused on modern samples.
- For enigmatic reasons, not all of the samples from the Yoshida and Martin papers were actually available online. Therefore our numbers here do not match those presented in the papers.
- The Judelson data include a sample of US1 which was sampled at three different time points (us1\_1, us1\_2 and us1\_3). We suspect that these were different samples and not necessarily the same clone. Therefore differences among these samples may either be due to biological or technical factors.
- The Judelson data includes a sample of US8 which has been characterized as having fungicide resistance [3]. This lineage was also sequenced by Martin et al. [7]. These are most likely different samples so differences among these samples may be interpreted as biological.
  - The Yoshida data includes one sample of *P. mirabilis* (p7722), this should jump out in the analyses.
- For the mitochondrial data we used the type IIa form [1] because it was the longest sequence and we felt this would provide the best alignment.

We've used the term 'SNP' fairly loosely in this document. The term 'variant' may be more appropriate. Until fairly recently the software tools we've been using could only handle SNPs. They now report short indels as well.
We've included both variant types here.

## 35 2 Variant discovery

Reads were mapped to the type IIa mitochondrial reference "AY898627.1".

Reads were mapped using bowtie2 [5]. Variants were called using SAMtools[6].

## 38 2.1 Variant filtering

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As a quality control step, the variant files were filtered by quality, read depth and mapping quality (Figures 1,2). For this we used an in-house R package called vcfR. Here, sequencing depth is cumulative over all samples. Quality here is for each variant over all samples and ranges from 1-999.

The genotype caller in Samtools assumes a diploid, bi-allelic model. Because mitochondria are assumed to be haploid we tried to filter out heterozygous calls. Samples which included high quality heterozygote calls (p1362, p6096, p10650, p12204, p10127) were mostly from the Yoshida et al. [10] paper and were among the low sequencing depth samples they included. Because these samples are not among the US lineages we're interested in, and because they are apparently of low sequencing depth, we omitted them for now. However, the sample nl07434 was among the high sequencing depth samples from this paper and is perhaps noteworthy. T30-4 was called as a heterozygote for one variant and is perplexinng.

In an attept to identify high quality variants we employed a filtering strategy. Filtering of the variant panel was based on quality (QUAL=999), cumulative sequencing depth (1st quartile >= DP >= 3rd quartile) and mapping quality (1st quartile >= MQ >= 3rd quartile). This resulted in 37 variants remaining after filtering (Table 1). We have identified a fraction of these as being diagnostic for a small group of samples (Table 2).

The variants remaining after filtering were visualized as a linear chromosome in Figure 3.

```
## Before filtering:
## [1] 247
## After filtering:
## [1] 37
```

## $_{\scriptscriptstyle 51}$ 3 Variant segregation

In order to visualize how variants segregated among the samples, a phylogeny was inferred. We then used ancestral state reconstruction to map the characters

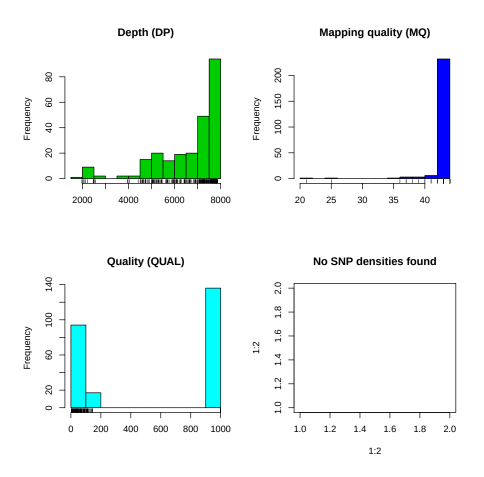


Figure 1: Quality control results for the mtDNA SNP calls before filtering.

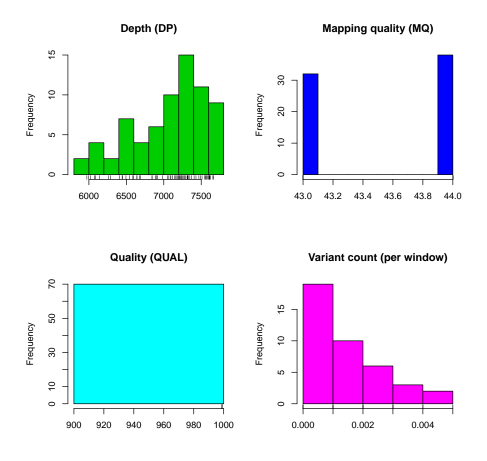


Figure 2: Quality control results for the mtDNA SNP calls after filtering and windowizing variants.

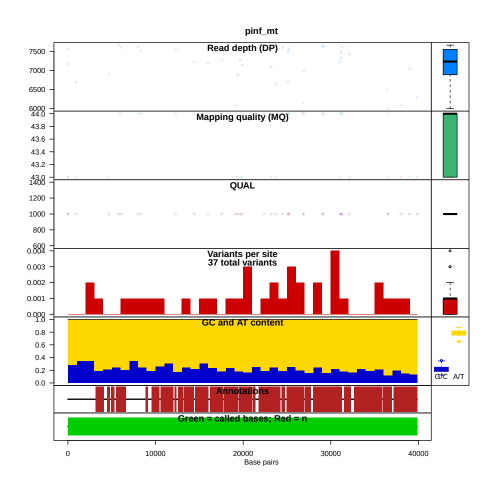


Figure 3: Whole mtDNA genome scan for the P. infestans samples.

- to the tree. At this time we're not trying to say anything bold about phylogeny
- or character evolution. We're simply using these tools to visualize how the
- 66 variants segregate.

#### <sub>67</sub> 3.1 Phylogenetic reconstruction

- Using the whole genome alignment (28 sequences, 39,870 nucleotides) we per-
- formed a whole-genome phylogeny using maximum likelihood (RAxML) and
- Bayesian inference (BEAST). We used RAxML using no partitions, 1000 boot-
- strap replicates, a GTR+I+G model of nucleotide evolution to obtain a biparti-
- tioned tree with the boostrap values mapped to the branches. For BEAST,
- we specified p7722 ( $P.\ mirabilis$ ) as the outgroup. We used a HKY+G+I model
- of nucleotide substitutions, a strict molecular clock, a constant population size
- prior, UPGMA starting tree and 10 million Markov chains. The best tree is
- shown in Figure 4.

## 3.2 Mapping the SNP's in the BEAST tree

- 78 To map the variants found in the mtDNA genome to the coalescent tree, we
- used Mesquite. We did a removal of invariable regions and ancestral state re-
- construction for all 37 SNPs using a parsimony reconstruction state. Select
- variants which we felt were diagnostic for a small number of lineages are pre-
- sented as trees with mapped characters (see tree figures).

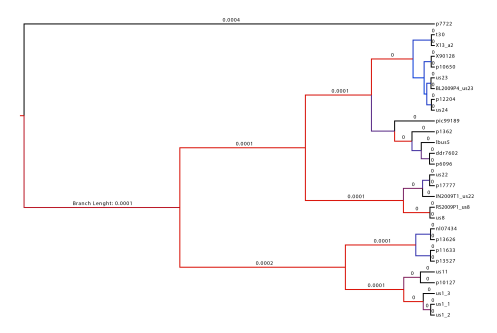
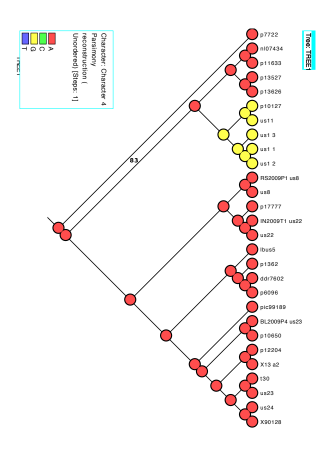
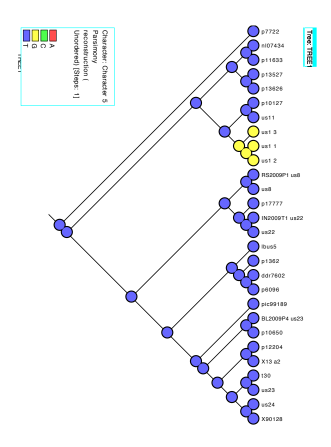
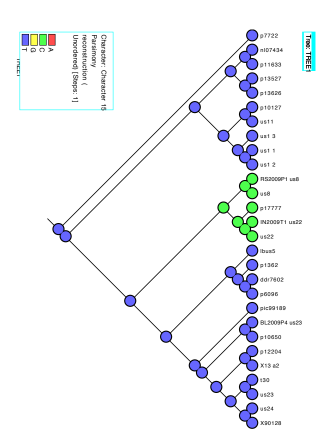
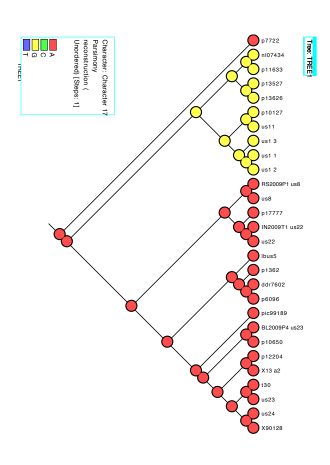


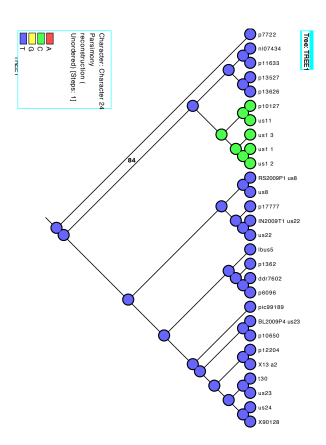
Figure 4: Bayesian coalescent tree of the whole mtDNA genome of *P. infestans* using BEAST 1.8.0. Values above branches represent branch lengths (topology is concordant to the bifurcating model of a coalescent reconstruction). Branches are colored based on their posterior probability values (legend indicates color scheme).

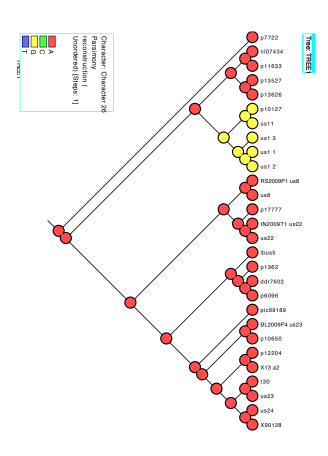


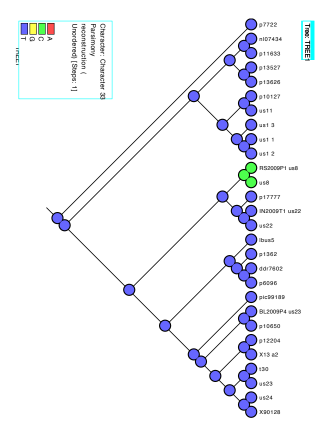












## 55 4 Session information

```
sessionInfo()
## R version 3.0.2 (2013-09-25)
## Platform: x86_64-pc-linux-gnu (64-bit)
##
## locale:
    [1] LC_CTYPE=en_US.UTF-8
                                    LC_NUMERIC=C
##
##
    [3] LC_TIME=en_US.UTF-8
                                    LC_COLLATE=en_US.UTF-8
##
    [5] LC_MONETARY=en_US.UTF-8
                                    LC_MESSAGES=en_US.UTF-8
##
    [7] LC_PAPER=en_US.UTF-8
                                    LC NAME=C
##
    [9] LC_ADDRESS=C
                                    LC_TELEPHONE=C
  [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
##
##
## attached base packages:
  [1] stats
                 graphics grDevices utils
                                                datasets
                                                          methods
                                                                     base
##
## other attached packages:
  [1] xtable_1.7-1 vcfR_0.1
                                  knitr_1.5
##
## loaded via a namespace (and not attached):
  [1] ape_3.0-11
                       evaluate_0.5.1 formatR_0.10
                                                        grid_3.0.2
  [5] lattice_0.20-24 nlme_3.1-111 stringr_0.6.2
                                                        tools_3.0.2
```

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  123 Kamoun, Johannes Krause, et al. Correction: The rise and fall of the
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  125 2, 2013.

	CHROM	POS	REF	ALT
6	Supercontig 1.1	2701	С	T
7	Supercontig 1.1	2728	$\mathbf{C}$	$\mathbf{A}$
12	Supercontig 1.1	3519	G	${f T}$
36	Supercontig 1.1	6872	accc	acc
42	Supercontig 1.1	7857	$\mathbf{A}$	G
44	Supercontig_1.1	8118	${ m T}$	G
59	Supercontig_1.1	9857	${ m T}$	$\mathbf{C}$
62	$Supercontig\_1.1$	10146	$\mathbf{A}$	G
85	$Supercontig\_1.1$	13684	$\mathbf{A}$	G
106	$Supercontig_1.1$	15563	G	$\mathbf{A}$
114	$Supercontig_1.1$	16986	G	$\mathbf{A}$
119	$Supercontig_1.1$	18440	$\mathbf{A}$	${f T}$
123	$Supercontig_1.1$	19793	G	$\mathbf{A}$
124	$Supercontig_1.1$	20005	$\mathbf{C}$	$\mathbf{A}$
125	$Supercontig\_1.1$	20290	G	${f T}$
128	$Supercontig\_1.1$	20464	${f T}$	$^{\mathrm{C}}$
141	$Supercontig\_1.1$	22711	G	$\mathbf{A}$
147	$Supercontig\_1.1$	23431	G	$\mathbf{A}$
150	$Supercontig\_1.1$	23872	G	${f T}$
154	$Supercontig\_1.1$	24611	$\mathbf{C}$	${f T}$
159	$Supercontig\_1.1$	25142	$\mathbf{A}$	$^{\mathrm{C}}$
160	$Supercontig\_1.1$	25204	${ m T}$	$^{\mathrm{C}}$
161	$Supercontig\_1.1$	25237	G	${f T}$
169	$Supercontig\_1.1$	26684	$\mathbf{A}$	G
171	$Supercontig\_1.1$	26767	${ m T}$	$^{\mathrm{C}}$
186	$Supercontig\_1.1$	28680	$\mathbf{C}$	$\mathbf{A}$
187	$Supercontig\_1.1$	28783	$\mathbf{A}$	G
192	$Supercontig\_1.1$	30427	${ m T}$	G
193	$Supercontig\_1.1$	30552	$\mathbf{A}$	G
194	$Supercontig\_1.1$	30591	$\mathbf{A}$	G
195	$Supercontig\_1.1$	30660	$^{\mathrm{C}}$	${ m T}$
201	$Supercontig\_1.1$	31403	${f T}$	$^{\mathrm{C}}$
221	$Supercontig\_1.1$	35296	G	${f T}$
222	${ m Supercontig}\_1.1$	35367	taaaaaaaaaaa	taaaaaaaaaa
233	$Supercontig\_1.1$	36663	${f T}$	$^{\mathrm{C}}$
239	$Supercontig\_1.1$	37562	$\mathbf{C}$	${f T}$
242	$Supercontig\_1.1$	38345	$\mathbf{C}$	A

Table 1: Variants remaining after filtering.

Table 2: Diagnostic SNP positions for the mtDNA genome after filtering. The tree number corresponds to the character legend on the tree figures

Position	SNP	Diagnostic for	Character number
7857	A/G	p10127, us11 and us1	4
8118	T/G	us1	5
20464	T/C	p17777, us22 and us8	15
22711	G/A	p17777, us22 and us8	17
26767	T/C	p10127, us11 and us1	24
28783	A/G	p10127, us11 and us1	26
36663	T/C	us8	33