MEME analysis (v3.0) was performed on the alignment from

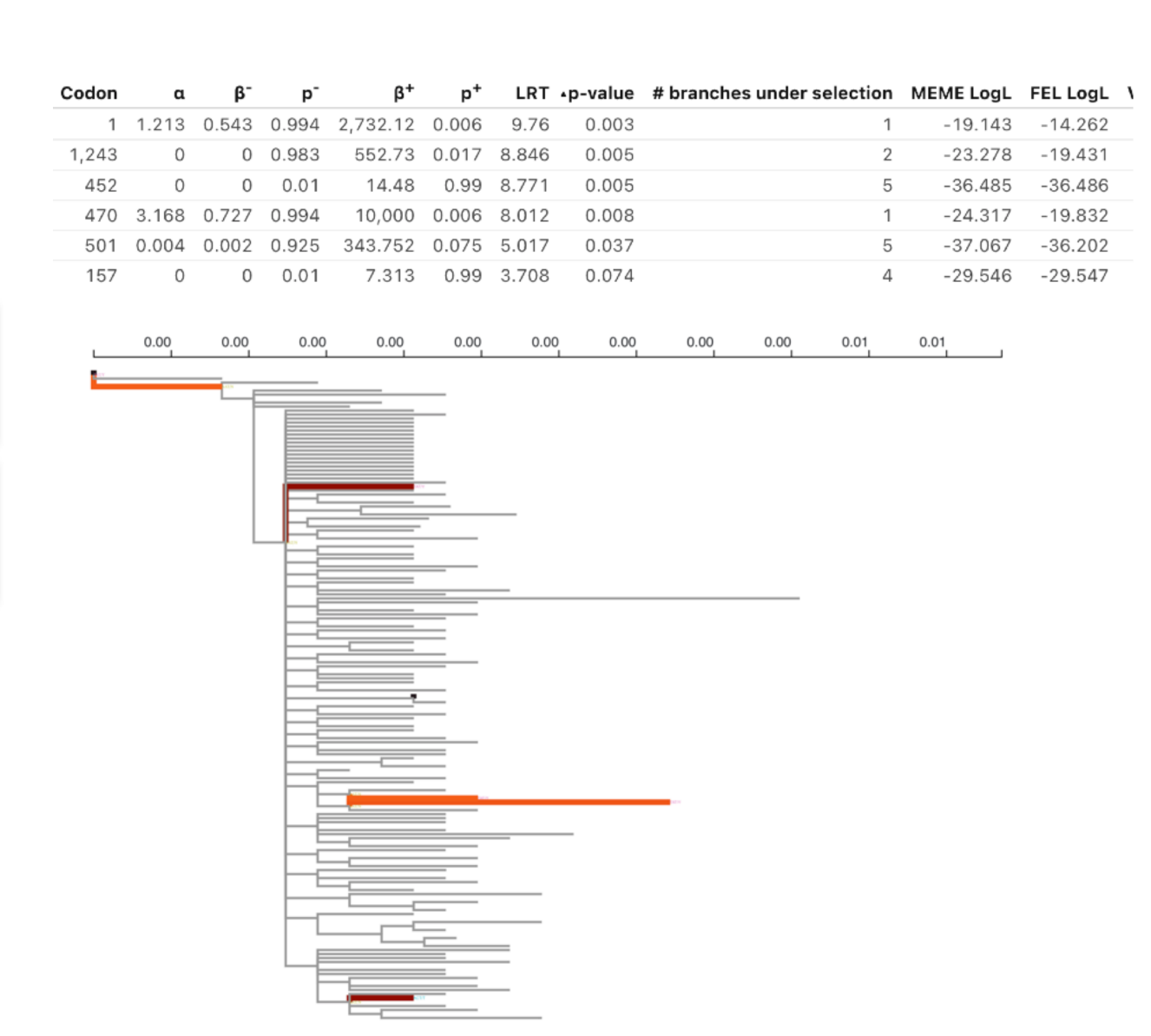
/Users/sergei/Dropbox/Talks/VEME-current/data/spike.fas using HyPhy v2.5.40.

p-value threshold 0.1 Update

Suggested citation: Detecting Individual Sites Subject to Episodic Diversifying Selection.

PLoS Genet 8(7): e1002764.

118 sequences in the alignment	1273 codon sites in the alignment	1 partitions
162 median branches/partition used for testing	N/A bootstrap replicates	6 sites subject to episodic diversifying selection
3.00 median branches with support for selection/selected site	$\frac{3}{\text{sites with variable }\omega}$ across $\frac{4}{9}$	



hyphy meme --alignment data/spike.fas --tree data/spike.tree

Interpreting dN/dS for intra-host and intra-species pathogen

- dN/dS can be estimated for all sorts of sequence data (e.g., it has been done for cancer SNP data)
- Traditional interpretation of dN/dS is based on the assumption that substitution ~ fixation
- Not the same for intra-species / intra-host pathogens

- Much of variation is due to polymorphism, or even dead-end mutations
- This is because selection has not had a chance to "filter" mutations (except for patently deleterious ones)
- This often manifests as differences in selective "regimes" between tips and internal branches