Based on the likelihood ratio test, there **is** evidence of *episodic diversifying* selection in this dataset (p=0.000).

BUSTED analysis (v4.0) was performed on the alignment from /Users/sergei/Dropbox/Talks/VEME-current/data/HIV-sets.fas using HyPhy v2.5.40. This analysis **did not include** site-to-site synonymous rate variation.

Suggested citation: Gene-wide identification of episodic selection, Mol Biol Evol.

32(5):1365-71, Synonymous Site-to-Site Substitution Rate Variation Dramatically Inflates False Positive Rates of Selection Analyses: Ignore at Your Own Peril, Mol Biol Evol. 37(8):2430-2439

sequences in the

alignment

Evidence ratio 10 Update threshold 16 288 codon sites in the

26 used for testing

variation

alignment

None synonymous rate variation

0.0 p-value for episodic diversityfing selection

13 Sites with ER≥10 for positive selection



Alignment-wide results

Log (L) AIC-c Params. Rate distribution Model Rate plot -2039.96 4170.83 Unconstrained model 45 **Tested ω** 0.5596 (86.941%) 0.9885 (10.960%) 96.09 (2.0981%) Mean = 2.611, CoV = 5.24220 40 60 80100 Constrained model -2078.31 4245.48 44 Tested ω 1.000 (14.819%) 1.000 (20.229%) 1.000 (64.952%) Mean = 1.000, CoV = NaN 73

Gene-wide selection analysis using a branch-site method (BUSTED), HIV-1 env

hyphy busted --srv No --alignment data/HIV-sets.nex --starting-points 5

Produces HIV-sets.nex.BUSTED.json file View in http://vision.hyphy.org/BUSTED

Figure 1. Empirical Bayes Factors for $\omega > 1$ at a particular branch and site (only tested branches are included).

