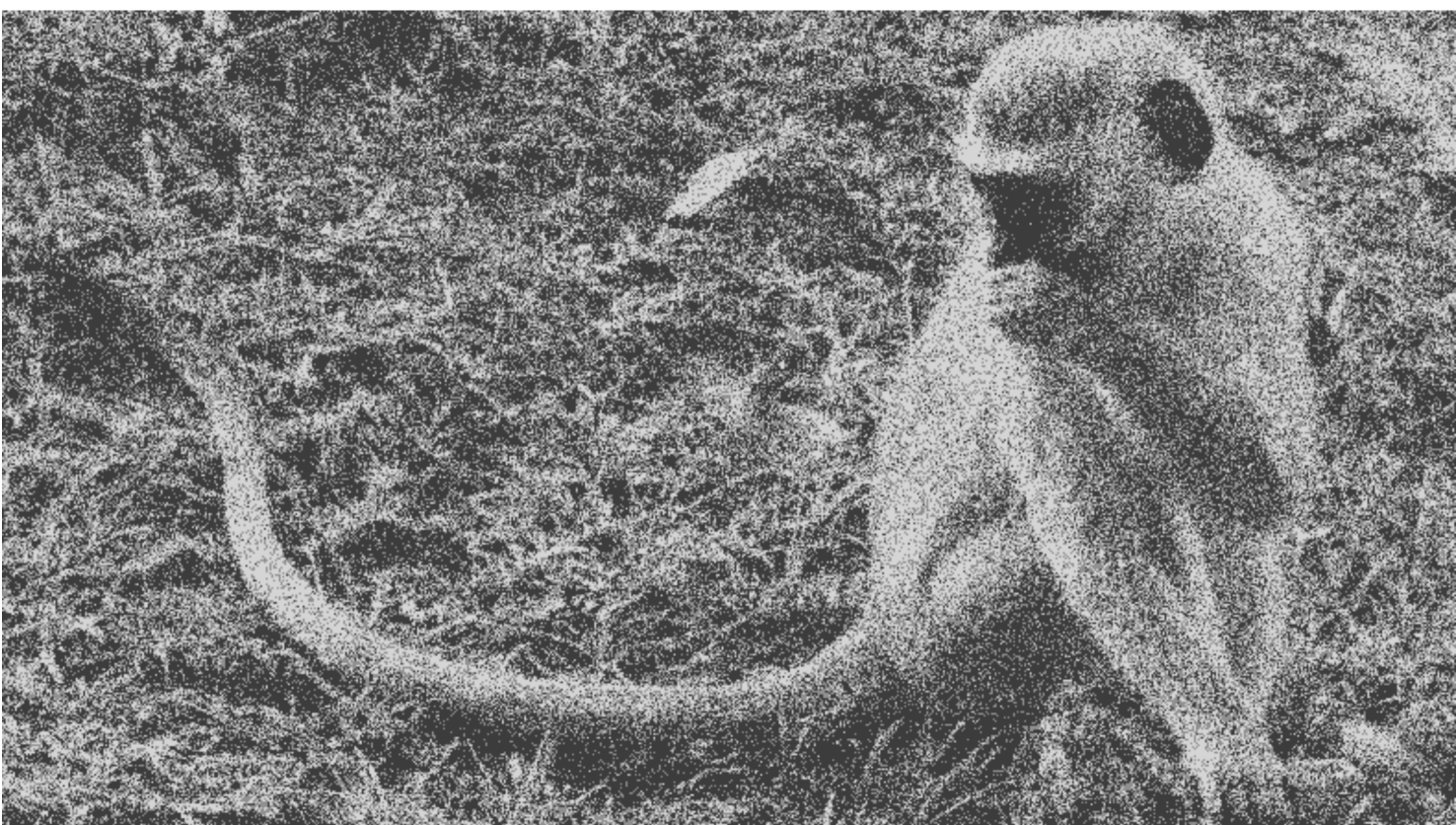
Gene-wide selection random effects over sites and branches [BUSTED]



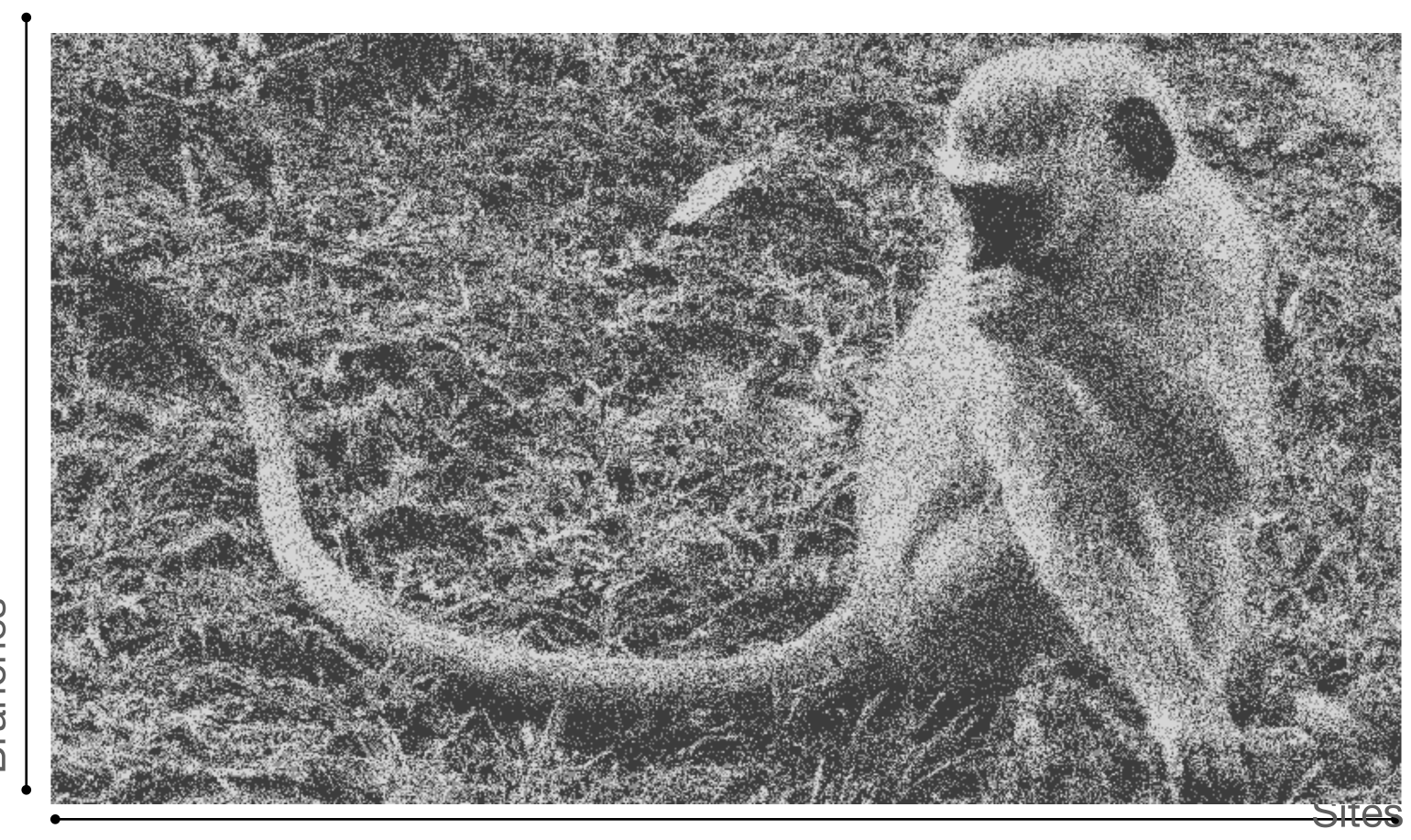


0	Is there enough image area that is sufficiently bright; allow each pixel to be one of $K (=3)$ colors, chosen adaptively, e.g. to minimize perceptual differences
*	[BUSTED]: each branch-site combination is a drawn from a K-bin (dS,dN) distribution. The distribution is estimated from the entire alignment. Tests if dN/dS>1 for some branch/site pairs in the alignment



Gene-wide selection

random effects over sites and branches [BUSTED]



Is there enough **image area** that is sufficiently bright; allow each pixel to be one of K (=3) colors, chosen adaptively, e.g. to minimize perceptual differences



[BUSTED]: each branch-site combination is a drawn from a K-bin (dS,dN) distribution. The distribution is estimated from the entire alignment. Tests if dN/dS>1 for some branch/site pairs in the alignment

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

Evidence ratio 10 Update threshold 16 288 sequences in the codon sites in the alignment alignment 26 None synonymous rate variation used for testing variation 13 0.0 Sites with ER≥10 for p-value for episodic diversityfing selection positive selection

Alignment-wide results

Model Log (L) AIC-c Params. Rate distribution Rate plot Unconstrained model -2039.96 4170.83 45 Tested ω 0.5596 (86.941%) 0.9885 (10.960%) 96.09 (2.0981%) Mean = 2.611, CoV = 5.242 0 20 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

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

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

Produces *HIV-sets.fas.BUSTED.json* file View in http://vision.hyphy.org/BUSTED or https://observablehq.com/@spond/busted