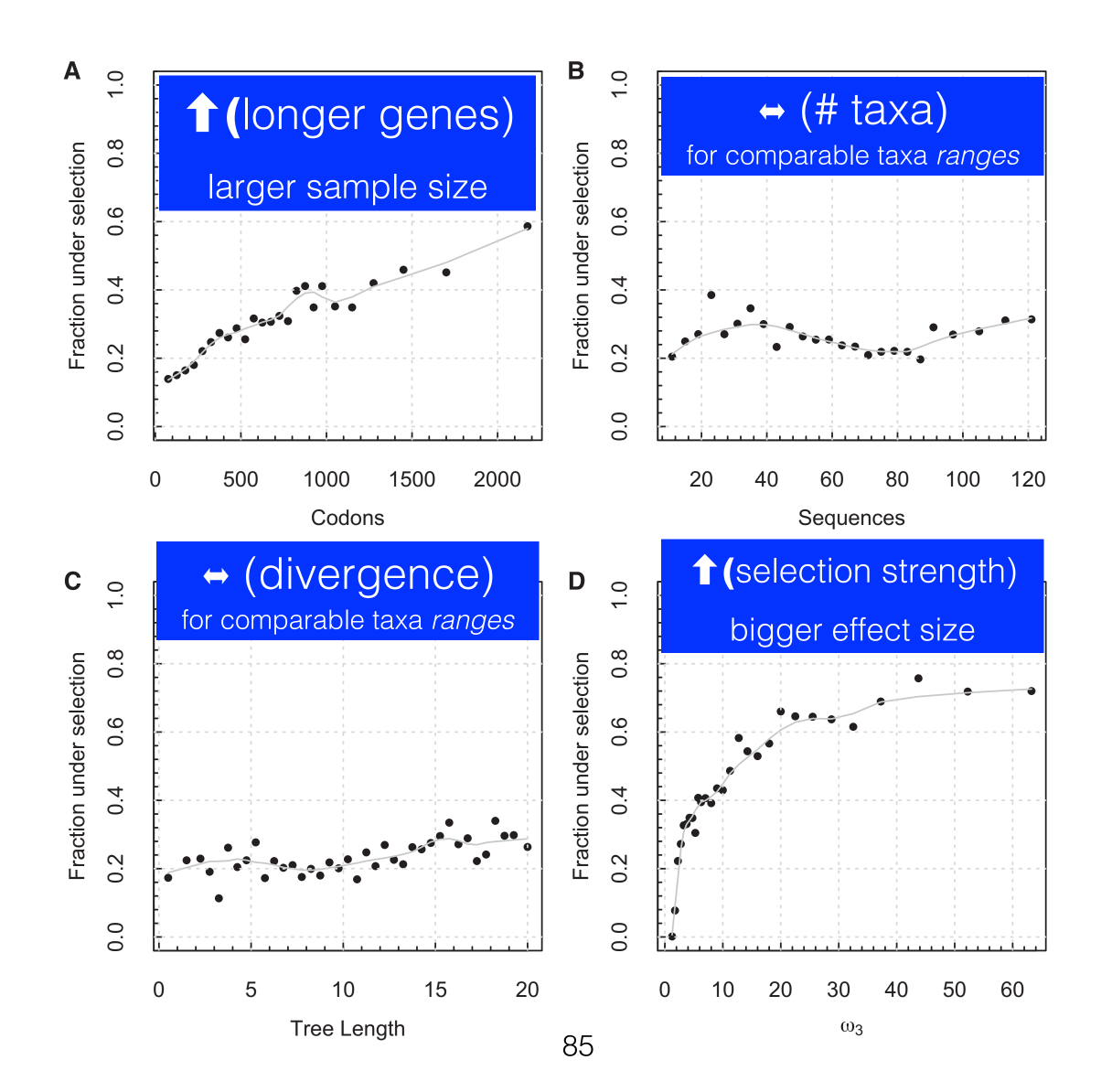
Where does the power come from for BUSTED?

An analysis of ~9,000 curated gene alignments from selectome.unil.ch



Any questions on the previous material?

We covered the following methods in HyPhy: FitMG94, BUSTED

Which estimates a mean gene-wide dN/dS (FitMG94)
Or estimates dN/dS through a branch-site method (BUSTED)