Probability of the alignment given the tree and sequence evolution model

model Tree Prior Transmission model

sequence

Dates

at which each

Transmission model that describes the tree generating process

Average number of secondary infections caused by one case within the same type

Proportion of infections sequenced and included in the analysis

Probability that an infected case is removed from the population after sampling

Model describing the rate of evolution along the branches of the tree

RELAXED CLOCK

Model describing the rates of substitution between nucleotides

HKY

A set of aligned sequences representing the genetic data for each sample

Graphical representation of the evolutionary relationships between samples

Stationary frequencies in evolutionary equilibrium

Rate at which an infected case becomes non-infections/dies/recovers



