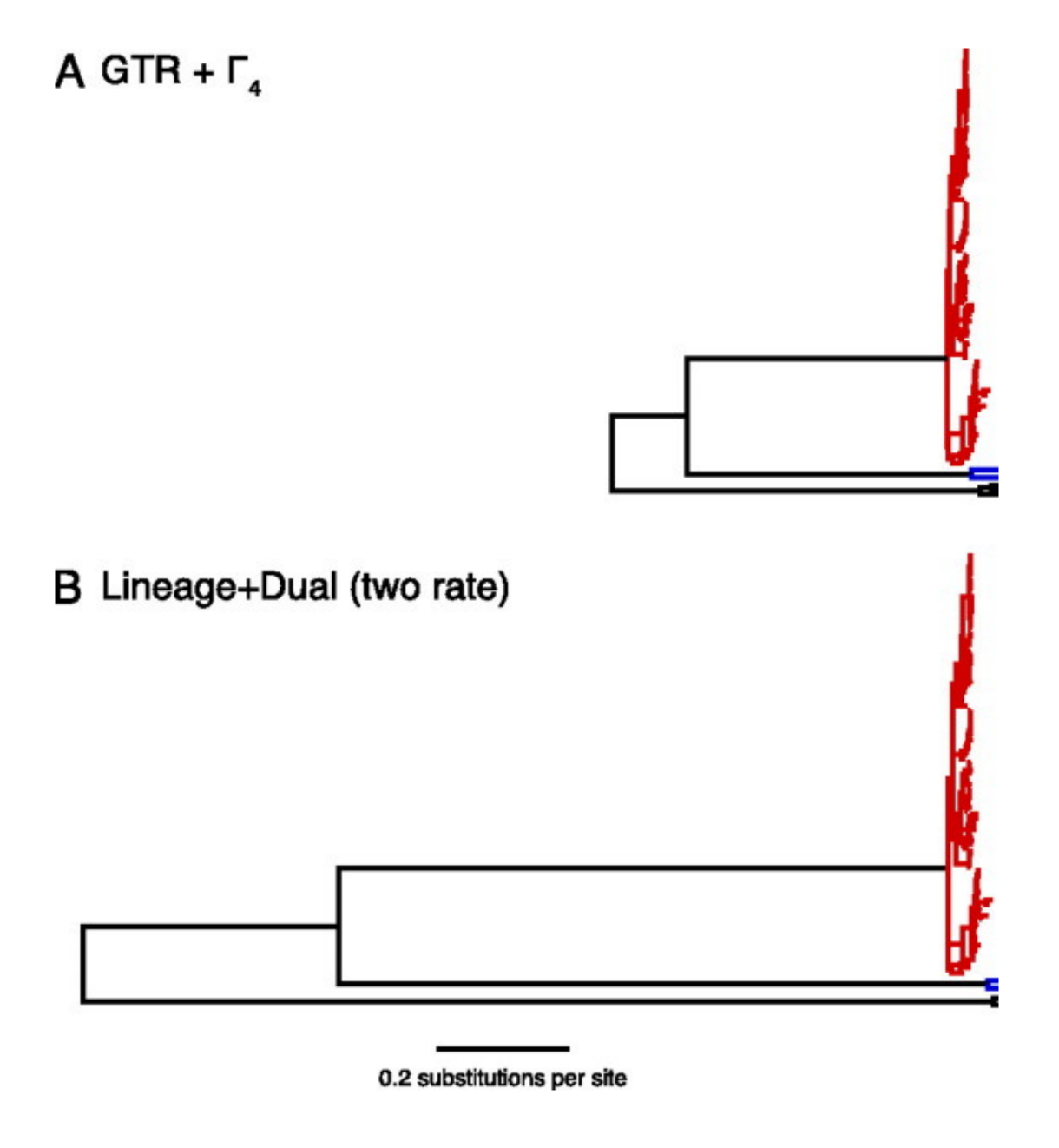
- Using models that do not vary selection pressure across lineages yields a patently false "too young" estimate for the origin of measles (about 600 years ago)
- This estimate is refuted by clear historical records which suggest that measles is at least 1,500-5,000 years old
 - This includes a treatise by a Persian physician Rhazes about differential diagnosis of measles and smallpox published circa 600 AD.
- Same patterns found for coronaviruses, ebola, avian influenza and herpesvirus



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