## Molecular signatures of selection

 The rate of accumulation of synonymous substitutions (dS) can serve as the neutral background evolutionary rate We can compare the rate of accumulation of non-synonymous substitutions (dN), which alter the

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protein sequence, to **dS** and use their ratio to classify the nature of the evolutionary process

### number of fixed synonymous mutations proportion of random mutations that are synonymous

number of fixed non-synonymous mutations proportion of random mutations that are non-synonymous

 $dN \sim -$ 

#### What can the denominator proportions depend on?

## Molecular signatures of selection

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- The **rate** of accumulation of synonymous substitutions (**dS**) can serve as the neutral background evolutionary rate
- We can compare the **rate** of accumulation of non-synonymous substitutions (**dN**), which alter the protein sequence, to **dS** and use their ratio to classify the nature of the evolutionary process

$$dS \sim \frac{\text{number of fixed synonymous mutations}}{\text{proportion of random mutations that are synonymous}}$$

$$dN \sim \frac{\text{number of fixed non-synonymous mutations}}{\text{proportion of random mutations that are non-synonymous}}$$

# Evolutionary Modes

Positive Selection (Diversifying)

Negative Selection

Neutral Evolution

dS < dN or $\omega := dN/dS > 1$ 

 $dS > dN \text{ or } \omega < 1$ 

 $dS \simeq dN \text{ or } \omega \simeq 1$