

KaKs_Calculator: Calculating Ka and Ks Through Model Selection and Model Averaging

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KaKs_Calculator is a software package that calculates nonsynonymous (Ka) and synonymous (Ks) substitution rates through model selection and model averaging. Since existing methods for this estimation adopt their specific mutation (substitution) models that consider different evolutionary features, leading to diverse estimates, **KaKs_Calculator** implements a set of candidate models in a maximum likelihood framework and adopts the Akaike information criterion to measure fitness between models and data, aiming to include as many features as needed for accurately capturing evolutionary information in protein-coding sequences. In addition, several existing methods for calculating Ka and Ks are also incorporated into this software. **KaKs_Calculator**, including source codes, compiled executables, and documentation, is freely available for academic use at <http://evolution.genomics.org.cn/software.htm>.

Key words: model selection, model averaging, AIC, approximate method, maximum likelihood method

Introduction

Calculating nonsynonymous (Ka) and synonymous (Ks) substitution rates is of great significance in reconstructing phylogeny and understanding evolutionary dynamics of protein-coding sequences across closely related and yet diverged species (1–3). It is known that Ka and Ks, or often their ratio (Ka/Ks), indicate neutral mutation when Ka equals to Ks, negative (purifying) selection when Ka is less than Ks, and positive (diversifying) selection when Ka exceeds Ks. Therefore, statistics of the two variables in genes from different evolutionary lineages provides a powerful tool for quantifying molecular evolution.

Over the past two decades, several methods have been developed for this purpose, which can generally be classified into two classes: approximate method and maximum likelihood method. The approximate method involves three basic steps: (1) counting the numbers of synonymous and nonsynonymous sites, (2) calculating the numbers of synonymous and nonsynonymous substitutions, and (3) correcting for multiple

substitutions. On the other hand, the maximum likelihood method integrates evolutionary features (reflected in nucleotide models) into codon-based models and uses the probability theory to finish all the three steps in one go (4). However, these methods adopt different substitution or mutation models based on different assumptions that take account of various sequence features, giving rise to varied estimates of evolutionary distance (5). In other words, Ka and Ks estimation is sensitive to underlying assumptions or mutation models (3). In addition, since the amount and the degree of sequence substitutions vary among datasets from diverse taxa, a single model or method may not be adequate for accurate Ka and Ks calculations. Therefore, a model selection step, that is, to choose a best-fit model when estimating Ka and Ks, becomes critical for capturing appropriate evolutionary information (6, 7).

Toward this end, we have applied model selection and model averaging techniques for Ka and Ks estimations. We use a maximum likelihood method based on a set of candidate substitution models and adopt the Akaike information criterion (AIC) to measure fitness between models and data. After choosing the

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best-fit model for calculating Ka and Ks, we average the parameters across the candidate models to include as many features as needed since the true model is seldom one of the candidate models in practice (8). Finally, these considerations are incorporated into a software package, namely KaKs_Calculator.

Algorithm

Candidate models

Substitution models play a significant role in phylogenetic and evolutionary analyses of protein-coding sequences by integrating diverse processes of sequence evolution through various assumptions and providing approximations to datasets. We focused on a set of time-reversible substitution models (9–16) as shown in Table 1 (17, 18), ranging from the Jukes-Cantor (JC) model, which assumes that all substitutions have equal rates and equal nucleotide frequencies, to the general time-reversible (GTR) model that considers six different substitution rates and unequal nucleotide frequencies. Subsequently, we incorporated the parameters in each nucleotide model into a codon-based model (19, 20). As a result, a general formula of the substitution rate q_{ij} from any sense codon i to j ($i \neq j$) is given for all candidate models (19):

$$q_{ij} = \begin{cases} 0 & \text{if } i \text{ and } j \text{ differ by more than one} \\ & \text{difference} \\ \kappa_{xy} \pi_j & \text{if } i \text{ and } j \text{ differ by a synonymous} \\ & \text{substitution of } x \text{ for } y \\ \omega \kappa_{xy} \pi_j & \text{if } i \text{ and } j \text{ differ by a nonsynony-} \\ & \text{mous substitution of } x \text{ for } y \end{cases}$$

where π_j is the frequency of codon j , ω is the Ka/Ks ratio, and κ_{xy} is the ratio of r_{xy} to r_{CA} , $x, y \in \{A, C, G, T\}$ (Table 1). For example, in the JC model, κ_{xy} and π_j are equal to 1 owing to equal substitution rates and equal nucleotide frequencies assumed. In the Hasegawa-Kishino-Yano (HKY) model, κ_{TC} and κ_{AG} become equivalent to the transition/transversion rate ratio and π_j can be estimated from sequences, similar to the method reported by Goldman and Yang (19). Other models can be accommodated by making obvious modifications. Therefore, we could acquire maximum likelihood scores in various values generated from individual candidate model by implementing the codon-based models in a maximum likelihood framework (19, 20).

Model selection

AIC (21) has been widely used in model selection aside from other methods such as the likelihood ratio test (LRT) and the Bayesian information criterion (BIC) (8). AIC characterizes the Kullback-Leibler distance between a true model and an examined model, and this distance can be regarded as quantifying the information lost by approximating the true model. KaKs_Calculator uses a modification of AIC (AIC_C), which takes account of sampling size (n), maximum likelihood score ($\ln L_i$), and the number of parameters (k_i) in model i as follows:

$$AIC_{Ci} = AIC_i + \frac{2k_i(k_i + 1)}{n - k_i - 1} = -2 \ln L_i + 2k_i + \frac{2k_i(k_i + 1)}{n - k_i - 1}$$

AIC_C is proposed to correct for small sampling size, and it approaches to AIC when sampling size comes to infinity. Consequently, we could use this equation to compute AIC_C for each candidate model and then identify a model that possesses the smallest AIC_C , which is a sign for appropriateness between models and data.

Model averaging

Model selection is merely an approximate fit to a dataset, whereas a true evolutionary model is seldom one of the candidate models (8). Therefore, an alternative way is model averaging, which assigns each candidate model a weight value and engages more than one model to estimate average parameters across models. Accordingly, we first need to compute the Akaike weight (w_i , where $i = 1, 2, \dots, m$) for each model in a set of candidate models:

$$w_i = \frac{\exp[-\frac{1}{2}(AIC_{Ci} - \min AIC_C)]}{\sum_{j=1}^m \exp[-\frac{1}{2}(AIC_{Cj} - \min AIC_C)]}$$

where $\min AIC_C$ is the smallest AIC_C value among candidate models. We can then estimate model-averaged parameters. Taking κ_{TC} as an example, a model-averaged estimate can be calculated by:

$$\kappa_{TC} = \frac{\sum_{i=1}^m [w_i \times I(\kappa_{TC,i}) \times \kappa_{TC,i}]}{\sum_{i=1}^m [w_i \times I(\kappa_{TC,i})]}$$

where $\kappa_{TC,i}$ is κ_{TC} in model i and

$$I(\kappa_{TC,i}) = \begin{cases} 1 & \text{if } r_{TC} \neq r_{CA} \text{ in model } i \\ 0 & \text{otherwise} \end{cases}$$

Table 1 Candidate Models for Model Selection and Model Averaging in KaKs_Calculator

Model	Description (Reference)	Nucleotide frequency	Substitution rate*
JC	Jukes-Cantor model (9)	Equal	$r_{TC} = r_{AG} = r_{TA} = r_{CG} = r_{TG} = r_{CA}$
F81	Felsenstein's model (10)	Unequal	
K2P	Kimura's two-parameter model (11)	Equal	$r_{TC} = r_{AG} \neq r_{TA} = r_{CG} = r_{TG} = r_{CA}$
HKY	Hasegawa-Kishino-Yano model (12)	Unequal	
TNEF	TN model with equal nucleotide frequencies	Equal	$r_{TC} \neq r_{AG} \neq r_{TA} = r_{CG} = r_{TG} = r_{CA}$
TN	Tamura-Nei model (13)	Unequal	
K3P	Kimura's three-parameter model (14)	Equal	$r_{TC} = r_{AG} \neq r_{TA} = r_{CG} \neq r_{TG} = r_{CA}$
K3PUF	K3P model with unequal nucleotide frequencies	Unequal	
TIMEF	Transition model with equal nucleotide frequencies	Equal	$r_{TC} \neq r_{AG} \neq r_{TA} = r_{CG} \neq r_{TG} = r_{CA}$
TIM	Transition model	Unequal	
TVMEF	Transversion model with equal nucleotide frequencies	Equal	$r_{TC} = r_{AG} \neq r_{TA} \neq r_{CG} \neq r_{TG} \neq r_{CA}$
TVM	Transversion model	Unequal	
SYM	Symmetrical model (15)	Equal	$r_{TC} \neq r_{AG} \neq r_{TA} \neq r_{CG} \neq r_{TG} \neq r_{CA}$
GTR	General time-reversible model (16)	Unequal	

* r_{ij} indicates the rate of substitution of i for j , where $i, j \in \{A, C, G, T\}$.

Application

KaKs_Calculator is written in standard C++ language. It is readily compiled and run on Unix/Linux or workstation (tested on AIX/IRIX/Solaris). In addition, we use Visual C++ 6.0 for graphic user interface and provide its Windows version that can run on any IBM compatible computer under Windows operating system (tested on Windows 2000/XP). Compiled executables on AIX/IRIX/Solaris and setup application on Windows, as well as source codes, example data, instructions for installation and documentation for KaKs_Calculator is available at <http://evolution.genomics.org.cn/software.htm>.

Different from other existing tools (22, 23), KaKs_Calculator employs model-selected and model-averaged methods based on a set of candidate models to estimate Ka and Ks. It integrates as many features as needed from sequence data and in most cases gives rise to more reliable evolutionary information (see the comparative results on simulated sequences at <http://evolution.genomics.org.cn/doc/SimulatedResults.xls>) (24). KaKs_Calculator also provides comprehensive information estimated from compared sequences, including the numbers of synonymous and nonsynonymous sites and substitutions, GC contents, maximum likelihood scores, and AIC_C . Moreover, KaKs_Calculator incorporates several other methods (19, 25-31) and allows users to choose one or more methods at one running time (Table 2).

Table 2 Methods Incorporated in KaKs_Calculator

Method	Approximate method			Reference
	Mutation model ^{#1}			
	Step 1	Step 2	Step 3	
NG	JC	JC	JC	26
LWL	JC	K2P	K2P	28
MLWL	K2P	K2P	K2P	30
LPB	—*	—*	K2P	25, 29
MLPB	—*	—*	K2P	30
YN	HKY	HKY	HKY	27
MYN	TN	TN	TN	31
Method	Maximum likelihood method			Reference
	Mutation model ^{#2}			
	GY	HKY		
MS	a model that has the smallest AIC_C among 14 candidate models		Model-selected method proposed in this study	
MA	a model that averages parameters across 14 candidate models		Model-averaged method proposed in this study	

^{#1}The approximate method involves three basic steps: Step 1: counting the numbers of synonymous and nonsynonymous sites; Step 2: calculating the numbers of synonymous and nonsynonymous substitutions; Step 3: correcting for multiple substitutions. ^{#2}The maximum likelihood method uses the probability theory to finish the three steps in one go (4). *No specific definition of synonymous and nonsynonymous sites or substitutions.

Although there exist 203 time-reversible models of nucleotide substitution (8), model selection in practice is often limited to a subset of them (32), and thus model averaging can reduce biases arising from model selection. Therefore, model-averaged methods should be preferred for general calculations of Ka and Ks. Some planned improvements include application of model selection and model averaging to detect positive selection at single amino acid sites, which requires high-speed computing for maximum likelihood estimation, especially when an adopted model becomes complex.

In conclusion, KaKs_Calculator incorporates as many features as needed for accurately extracting evolutionary information through model selection and model averaging, therefore it may be useful for in-depth studies on phylogeny and molecular evolution.

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Authors' contributions

ZZ designed and programmed this software, and drafted the manuscript. JL carried out computer simulations to generate sequences. XQZ performed test for earlier versions of the software. JW and GKSW contributed in conceiving this software and participated in software design. JY supervised the study and revised the manuscript. All authors read and approved the final manuscript.

Competing interests

The authors have declared that no competing interests exist.

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