#### **MKT Menu**











**RESOURCES Databases** with MKT estimates







# **Your MKT Results**

Comparison Number:	Neutral Region:	Testing Region:	Counts correspond <u>to</u> SITES/CHANGES:
Comparison <u>1</u>	1 (synonymous)	1 (non- synonymous)	Changes

Genetic code: Universal Code

## **INPUT SEQUENCES for Region 1:**

Number of Sequences for Species 1: 4 Number of Sequences for Species 2: 2

Length of the Alignment: 399

Percentage of gaps within the alignment (excluding end gaps): -15.0 %

### **View Alignment for Region 1** (Sequences were already aligned)

Input Sequences: 120627.1.txt

SEE RESULTS: ODivergence corrected by Jukes&Cantor 

Without any correction for divergence

**Comparison 1:** 

**NEUTRAL REGION: 1 (synonymous) TESTING REGION: 1 (non-synonymous)** 

**CHANGES** 

#### **Contingency table:**

	Polymorphism	Divergence	TOTAL
Neutral	0		0
Non- Neutral	0		0
TOTAL	0	0	0

### **Estimates:**

Neutrality Index (NI): NULL

Proportion of adaptive substitutions ( $\alpha$ ): NULL

 $\chi^2$  value: NULL p-value: NULL



