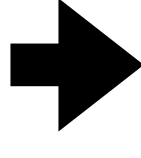
### results/WestNileVirus\_NS3.fna.FITTER.json

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"Standard MG94":{
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         "AIC-c":12923.31190785199,
         "Confidence Intervals":{
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           "non-synonymous/synonymous rate ratio":{
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359
360
             "UB":0.01055858911944712
361
362
          },
         "Log Likelihood":-6413.455134253863,
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         "Rate Distributions":{
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365
           "Substitution rate from nucleotide A to nucleotide G":1,
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           "Substitution rate from nucleotide A to nucleotide T":0.3060223183303938,
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           "Substitution rate from nucleotide C to nucleotide G":0.02087667707983453,
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           "Substitution rate from nucleotide C to nucleotide T":1.979133076079397,
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           "Substitution rate from nucleotide G to nucleotide T":0.2305847598841325,
370
           "non-synonymous/synonymous rate ratio":0.008575064843083209
371
372
         "display order":1,
373
         "estimated parameters":48
374
375
      },
376
     "input":{
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378
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379
       "number of sites":619,
380
       "partition count":1,
381
       "trees":{
382
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383
             (CHIN_01, EG101) Node17) Node15) Node7, (((ITALY_1998_EQUINE, PAAN001) Node22,
             (RO97_50, VLG_4) Node25) Node21, KN3829) Node20) Node6, AST99) Node5, PAH001) Node4, IS_98) Node3, MEX03) "
384
385
     "test results":{
        "non-synonymous/synonymous rate ratio":{
         "LRT": 2512.58476730381,
388
         "p-value":0
389
390
```



# WNV NS3

Model	Log L	# p	dN/dS	LRT	p-value
Null	-7745.48	49	1		
Alternative	-6413.5	50	0.009 [0.007-0.011]	2512.6	~0

# HIV-1 env

	Very strongly conserved					
Model	Log L	# p	dN/dS	LRT	p-value	

# Model Log L # p dN/dS LRT p-value Null -2078.3 40 1 Alternative -2078.2 41 1.122 [0.94-1.33] 0.33 ~0.6

## Not significantly different from neutral

# SARS-CoV-2 spike

Model	Log L	# p	dN/dS	LRT	p-value
Null	-9311.0	176	1		
Alternative	-9292.0	177	0.54 [0.48-0.61]	37.94	~0

Very strongly conserved