

Bayesian Evolutionary Analysis Sampling Trees Version v2.4.1, 2002-2016

BEAST XML File:	tutorial_run1.xml Choose File
	default: only write new log files \$
Random number seed:	1985
Thread pool size:	Automatic ‡
Use BEAGLE library if available:	
Prefer use of: CPU ‡	
Prefer precision: Double \$	
Show list of available BEAGLE resources and Quit	
BEAGLE is a high-performance phylogenetic library that can make use of additional computational resources such as graphics boards. It must be downloaded and installed independently of BEAST: http://beagle-lib.googlecode.com/	

Quit

Run