

Package: BeastJar (via r-universe)

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Type Package

Title JAR Dependency for MCMC Using 'BEAST'

Version 1.10.6

Description Provides JAR to perform Markov chain Monte Carlo (MCMC) inference using the popular Bayesian Evolutionary Analysis by Sampling Trees 'BEAST' software library of Suchard et al (2018) <[doi:10.1093/ve/vey016](https://doi.org/10.1093/ve/vey016)>. 'BEAST' supports auto-tuning Metropolis-Hastings, slice, Hamiltonian Monte Carlo and Sequential Monte Carlo sampling for a large variety of composable standard and phylogenetic statistical models using high performance computing. By placing the 'BEAST' JAR in this package, we offer an efficient distribution system for 'BEAST' use by other R packages using CRAN.

License Apache License 2.0

Encoding UTF-8

LazyData true

Imports rJava

URL <https://github.com/beast-dev/BeastJar>

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RoxygenNote 7.1.1

SystemRequirements Java (>= 1.8)

Suggests testthat

NeedsCompilation no

Author Marc A. Suchard [aut, cre, cph], Andrew Rambaut [cph], Alexei J. Drummond [cph]

Maintainer Marc A. Suchard <msuchard@ucla.edu>

Repository CRAN

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Description

Convenient packaging of the Bayesian Evolutionary Analysis Sampling Trees (BEAST) software package to facilitate Markov chain Monte Carlo sampling techniques including Hamiltonian Monte Carlo, bouncy particle sampling and zig-zag sampling.

Examples

```
# Example MCMC simulation using BEAST
#
# This function generates a Markov chain to sample from a simple normal distribution.
# It uses a random walk Metropolis kernel that is auto-tuning.

if (supportsJava8()) {
  # Set seed
  seed <- 123
  rJava::J("dr.math.MathUtils")$setSeed(rJava::.jlong(seed));

  # Set up simple model - Normal(mean = 1, sd = 2)
  mean <- 1; sd <- 2
  distribution <- rJava::.jnew("dr.math.distributions.NormalDistribution",
                              as.numeric(mean), as.numeric(sd))
  model <- rJava::.jnew("dr.inference.distribution.DistributionLikelihood",
                       rJava::.jcast(distribution, "dr.math.distributions.Distribution"))
  parameter <- rJava::.jnew("dr.inference.model.Parameter$Default", "p", 1.0,
                            as.numeric(-1.0 / 0.0), as.numeric(1.0 / 0.0))
  model$addData(parameter)

  # Construct posterior
  dummy <- rJava::.jnew("dr.inference.model.DefaultModel",
                       rJava::.jcast(parameter, "dr.inference.model.Parameter"))

  joint <- rJava::.jnew("java.util.ArrayList")
  joint$add(rJava::.jcast(model, "dr.inference.model.Likelihood"))
  joint$add(rJava::.jcast(dummy, "dr.inference.model.Likelihood"))

  joint <- rJava::new(rJava::J("dr.inference.model.CompoundLikelihood"), joint)

  # Specify auto-adapting random-walk Metropolis-Hastings transition kernel
  operator <- rJava::.jnew("dr.inference.operators.RandomWalkOperator",
                           rJava::.jcast(parameter, "dr.inference.model.Parameter"),
```

```

        0.75,
        rJava::J(
            "dr.inference.operators.RandomWalkOperator"
        )$BoundaryCondition$reflecting,
        1.0,
        rJava::J("dr.inference.operators.AdaptationMode")$DEFAULT
    )

schedule <- rJava::.jnew("dr.inference.operators.SimpleOperatorSchedule",
    as.integer(1000), as.numeric(0.0))

schedule$addOperator(operator)

# Set up what features of posterior to log
subSampleFrequency <- 100
memoryFormatter <- rJava::.jnew("dr.inference.loggers.ArrayLogFormatter", FALSE)
memoryLogger <-
    rJava::.jnew("dr.inference.loggers.MCLogger",
        rJava::.jcast(memoryFormatter, "dr.inference.loggers.LogFormatter"),
        rJava::.jlong(subSampleFrequency), FALSE)
memoryLogger$add(parameter)

# Execute MCMC
mcmc <- rJava::.jnew("dr.inference.mcmc.MCMC", "mcmc1")
mcmc$setShowOperatorAnalysis(FALSE)

chainLength <- 100000

mcmcOptions <- rJava::.jnew("dr.inference.mcmc.MCMCOptions",
    rJava::.jlong(chainLength),
    rJava::.jlong(10),
    as.integer(1),
    as.numeric(0.1),
    TRUE,
    rJava::.jlong(chainLength/100),
    as.numeric(0.234),
    FALSE,
    as.numeric(1.0))

mcmc$init(mcmcOptions,
    joint,
    schedule,
    rJava::.jarray(memoryLogger, contents.class = "dr.inference.loggers.Logger"))

mcmc$run()

# Summarize logged posterior quantities
traces <- memoryFormatter$getTraces()
trace <- traces$get(as.integer(1))

obj <- trace$getValues(as.integer(0),
    as.integer(trace$valueCount()))

```

```
sample <- rJava::J("dr.inference.trace.Trace")$toArray(obj)

outputStream <- rJava::.jnew("java.io.ByteArrayOutputStream")
printStream <- rJava::.jnew("java.io.PrintStream",
                           rJava::.jcast(outputStream, "java.io.OutputStream"))

rJava::J("dr.inference.operators.OperatorAnalysisPrinter")$showOperatorAnalysis(
  printStream, schedule, TRUE)

operatorAnalysisString <- outputStream$string("UTF8")

# Report auto-optimization information
cat(operatorAnalysisString)

# Report posterior quantities
c(mean(sample), sd(sample))
}
```

supportsJava8

Determine if Java virtual machine supports Java

Description

Tests Java virtual machine (JVM) java.version system property to check if version ≥ 8 .

Usage

```
supportsJava8()
```

Value

Returns TRUE if JVM supports Java ≥ 8 .

Examples

```
supportsJava8()
```

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