

Package ‘parglms’

September 16, 2024

Title support for parallelized estimation of GLMs/GEEs

Version 1.36.0

Author VJ Carey <stvjc@channing.harvard.edu>

Description This package provides support for parallelized estimation of GLMs/GEEs, catering for dispersed data.

Suggests RUnit, sandwich, MASS, knitr, GenomeInfoDb, GenomicRanges, gwascat, BiocStyle, rmarkdown

VignetteBuilder knitr

Depends methods

Imports BiocGenerics, BatchJobs, foreach, doParallel

Maintainer VJ Carey <stvjc@channing.harvard.edu>

License Artistic-2.0

LazyLoad yes

BiocViews statistics, genetics

ByteCompile TRUE

git_url <https://git.bioconductor.org/packages/parglms>

git_branch RELEASE_3_19

git_last_commit 42502e8

git_last_commit_date 2024-04-30

Repository Bioconductor 3.19

Date/Publication 2024-09-15

Contents

parglms-package	2
parGLM-methods	2
Index	4

parGLMs-package	<i>support for parallelized estimation of GLMs/GEEs</i>
-----------------	---

Description

This package provides support for parallelized estimation of GLMs/GEEs, catering for dispersed data.

Details

The DESCRIPTION file: This package was not yet installed at build time.

Index: This package was not yet installed at build time.

In version 0.0.0 we established an approach to fitting GLM from data that have been persistently dispersed and managed by a [Registry](#).

Author(s)

VJ Carey <stvjc@channing.harvard.edu>

Maintainer: VJ Carey <stvjc@channing.harvard.edu>

References

This package shares an objective with the `bigglm` methods of `biglm`. In `bigglm`, a small-RAM-footprint algorithm is employed, with sequential chunking to update statistics in each iteration. In `parGLM` the footprint is likewise controllable, but statistics in each iteration are evaluated in parallel over chunks.

Examples

```
showMethods("parGLM")
```

parGLM-methods	<i>fit GLM-like models with parallelized contributions to sufficient statistics</i>
----------------	---

Description

This package addresses the problem of fitting GLM-like models in a scalable way, recognizing that data may be dispersed, with chunks processed in parallel, to create low-dimensional summaries from which model fits may be constructed.

Methods

signature(formula = "formula", store = "Registry") The model data are assumed to lie in the file.dir/jobs/* folders, with file.dir defined in the store, which is an instance of [Registry](#).

Additional arguments must be supplied:

family a function that serves as a family for stats::glm

binit a vector of initial values for regression parameter estimation, must conform to expectations of formula

maxit an integer giving the maximum number of iterations allowed

tol a numeric giving the tolerance criterion

Failure to specify these triggers a fatal error.

The Registry instance can be modified to include a list element 'extractor'. This must be a function with arguments store, and codei. The standard extraction function is

```
function(store, i) loadResult(store, i)
```

It must return a data frame, conformant with the expectations of formula. Limited checking is performed.

The predict method computes the linear predictor on data identified by jobid in a BatchJobs registry. Results are returned as output of foreach over the jobids specified in the predict call.

Note that setting option parGLM.showiter to TRUE will provide a message tracing progress of the optimization.

Examples

```
if (require(MASS) & require(BatchJobs)) {
  # here is the 'sharding' of a small dataset
  data(anorexia) # N = 72
  # in .BatchJobs.R:
  # best setting for sharding a small dataset on a small machine:
  # cluster.functions = BatchJobs::makeClusterFunctionsInteractive()
  myr = makeRegistry("abc", file.dir=tempfile())
  chs = chunk(1:nrow(anorexia), n.chunks=18) # 4 recs/chunk
  f = function(x) {library(MASS); data(anorexia); anorexia[x,]}
  batchMap(myr, f, chs)
  submitJobs(myr) # now getResult(myr,1) gives back a data.frame
  waitForJobs(myr) # simple dispersal
  # now myr is populated
  oldopt = options()$parGLM.showiter
  options(parGLM.showiter=TRUE)
  pp = parGLM( Postwt ~ Treat + Prewt, myr,
    family=gaussian, binit = c(0,0,0,0), maxit=10, tol=.001 )
  print(summary(theLM <- lm(Postwt~Treat+Prewt, data=anorexia)))
  print(pp$coefficients - coef(theLM))
  if (require(sandwich)) {
    hc0 <- vcovHC(theLM, type="HC0")
    print(pp$robust.variance - hc0)
  }
}
predict(pp, store=myr, jobids=2:3)
options(parGLM.showiter=oldopt)
```

Index

* **methods**

parGLM-methods, [2](#)

* **modeling**

parGLM-methods, [2](#)

* **package**

parGLMs-package, [2](#)

parGLM (parGLM-methods), [2](#)

parGLM, formula, Registry-method
(parGLM-methods), [2](#)

parGLM-methods, [2](#)

parGLMs (parGLMs-package), [2](#)

parGLMs-package, [2](#)

predict (parGLM-methods), [2](#)

print (parGLM-methods), [2](#)

Registry, [2](#), [3](#)

summary (parGLM-methods), [2](#)