# Package 'TMB'

# December 25, 2014

Title General random effect model builder tool inspired by ADMB.

Type Package

version 1.1
<b>Date</b> 2014-09-19
Author Kasper Kristensen
Maintainer Kasper Kristensen <kaskr@imm.dtu.dk></kaskr@imm.dtu.dk>
<b>Description</b> With this tool, a user should be able to quickly implement complex random effect models through simple c++ templates. The package combines CppAD (c++ automatic differentiation), Eigen (templated matrix-vector library) and CHOLMOD (sparse matrix routines available from R) to obtain an efficient implementation of the applied Laplace approximation with exact derivatives. Key features are: Automatic sparseness detection, parallelism through BLAS and parallel user templates.
License GPL-2
<b>Depends</b> R (>= 3.0.0),Matrix(>= 1.0-12)
LinkingTo Matrix
R topics documented:
benchmark
compile         2           gdbsource         3
gdbsource
newton
openmp
precompile
Rinterface
runExample
sdreport
template
TMB-internal
Index 13

2 compile

ł	ገԲ	ın	$\boldsymbol{c}$	hr	na	r	k

Benchmark parallel templates

#### **Description**

Benchmark parallel templates

# Usage

```
benchmark(obj, n = 10, expr = NULL, cores = NULL)
```

## **Arguments**

obj	Object from MakeADFun
n	Number of replicates to obtain reliable results.
expr	Optional expression to benchmark instead of default.
cores	Optional vector of cores.

#### **Details**

By default this function will perform timings of the most critical parts of an AD model, specifically

- 1. Objective function of evaluated template.
- 2. Gradient of evaluated template.
- 3. Sparse hessian of evaluated template.
- 4. Cholesky factorization of sparse hessian.

(for pure fixed effect models only the first two). Expressions to time can be overwritten by the user (expr). A plot method is available for Parallel benchmarks.

# **Examples**

```
runExample("linreg_parallel",thisR=TRUE) ## Create obj
ben <- benchmark(obj,n=100,cores=1:4)
plot(ben)
ben <- benchmark(obj,n=10,cores=1:4,expr=expression(do.call("optim",obj)))
plot(ben)</pre>
```

compile

Compile a c++ template to DLL suitable for MakeADFun.

# Description

Compile a c++ template into a shared object file. OpenMP flag is set if the template is detected to be parallel.

# Usage

```
compile(file, flags = "", safebounds = TRUE, safeunload = TRUE,
  openmp = isParallelTemplate(file), libtmb = TRUE, ...)
```

gdbsource 3

#### **Arguments**

file c++ file.

flags Character with compile flags.

safebounds Turn on preprocessor flag for bound checking?
safeunload Turn on preprocessor flag for safe DLL unloading?

openmp Turn on openmp flag? Auto detected for parallel templates.

libtmb Use precompiled TMB library if available (to speed up compilation)?

... Passed as Makeconf variables.

#### **Details**

TMB relies on R's built in functionality to create shared libraries independent on the platform. A template is compiled by compile("template.cpp"), which will call R's makefile with appropriate preprocessor flags. Compiler and compiler flags can be stored in a configuration file. In order of precedence either via the file pointed at by R\_MAKEVARS\_USER or the file ~/.R/Makevars if it exists. Additional configuration variables can be set with . . . argument, which will overwrite any previous selections.

gdbsource

Source R-script through gdb to get backtrace.

#### **Description**

Source R-script through gdb to get backtrace.

#### Usage

```
gdbsource(file, interactive = FALSE)
```

# **Arguments**

file Your R script

interactive Run interactive gdb session?

## **Details**

This function is useful for debugging templates. If a script aborts e.g. due to an out-of-bound index operation it should be fast to locate the line that caused the problem by running gdbsource(file). Alternatively, If more detailed debugging is required, then gdbsource(file,TRUE) will provide the full backtrace followed by an interactive gdb session where the individual frames can be inspected. Note that templates should be compiled without optimization and with debug information i.e. compile(cppfile, "-00 -g") in order to provide correct line numbers.

#### Value

Object of class backtrace

4 MakeADFun

MakeADFun	Construct objective functions with derivatives based on a compiled c++ template.
Makeadi uli	1

# Description

Construct objective functions with derivatives based on the users c++ template.

# Usage

```
MakeADFun(data, parameters, map = list(), type = c("ADFun", "Fun",
   "ADGrad"[!is.null(random)]), random = NULL,
   random.start = expression(last.par.best[random]), hessian = FALSE,
   method = "BFGS", inner.method = "newton", inner.control = list(maxit =
   1000), MCcontrol = list(doMC = FALSE, seed = 123, n = 100),
   ADreport = FALSE, atomic = TRUE, LaplaceNonZeroGradient = FALSE,
   DLL = getUserDLL(), checkParameterOrder = TRUE, regexp = FALSE, ...)
```

## Arguments

•	guments	
	data	List of data objects (vectors,matrices,arrays,factors,sparse matrices) required by the user template (Order does not matter and un-used components are allowed).
	parameters	List of all parameter objects required by the user template (both random and fixed effects).
	map	List defining how to optionally collect and fix parameters - see details.
	type	Character vector defining which operation stacks are generated from the users template - see details.
	random	Character vector defining the random effect parameters. See also regexp.
	random.start	Expression defining the strategy for choosing random effect initial values as function of previous function evaluations - see details.
	hessian	Calculate Hessian at optimum?
	method	Outer optimization method.
	inner.method	Inner optimization method (see function "newton")
	inner.control	List controlling inner optimization
	MCcontrol	List controlling importance sampler (turned off by default)
	ADreport	Calculate derivatives of macro ADREPORT(vector) instead of objective_function return value?
	atomic	Allow tape to contain atomic functions?
	LanlacoNonZoro	Cradiant

 ${\tt LaplaceNonZeroGradient}$ 

Allow Taylor expansion around non-stationary point?

DLL Name of shared object file compiled by user.

 ${\tt checkParameterOrder}$ 

Optional check for correct parameter order.

regexp Match random effects by regular expressions?

... Currently unused.

MakeADFun 5

#### **Details**

A call to MakeADFun will return an object that, based on the users DLL code (specified through DLL), contains functions to calculate the objective function and its gradient. The object contains the following components:

- par A default parameter.
- fn The likelihood function.
- gr The gradient function.
- report A function to report all variables reported with the REPORT() macro in the user template.
- env Environment with access to all parts of the structure.

and is thus ready for a call to R's optim function. Data (data) and parameters (parameters) are directly read by the user template via the macros beginning with DATA\_ and PARAMETER\_. The order of the PARAMETER\_ macros defines the order of parameters in the final objective function. There are no restrictions on the order of random parameters, fixed parameters or data in the template.

Optionally, a simple mechanism for collecting and fixing parameters from R is available through the map argument. A map is a named list of factors with the following properties:

- names(map) is a subset of names(parameters).
- For a parameter "p" length(map\$p) equals length(parameters\$p).
- Parameter entries with NAs in the factor are fixed.
- Parameter entries with equal factor level are collected to a common value.

More advanced parameter mapping, such as collecting parameters between different vectors etc., must be implemented from the template.

Random effects are specified via the argument random: A component of the parameter list is marked as random if its name is matched by any of the characters of the vector random (Regular expression match is performed if regexp=TRUE). If some parameters are specified as random effects, these will be integrated out of the objective function via the Laplace approximation. In this situation the functions fn and gr automatically perform an optimization of random effects for each function evaluation. This is referred to as the 'inner optimization'. Strategies for choosing initial values of the inner optimization can be controlled via the argument random.start. The default is expression(last.par.best[random]) where last.par.best is an internal full parameter vector corresponding to the currently best likelihood. An alternative choice could be expression(last.par[random]) i.e. the random effect optimum of the most recent - not necessarily best - likelihood evaluation. Further control of the inner optimization can be obtained by the argument inner.control which is a list of control parameters for the inner optimizer newton. Depending of the inner optimization problem type the following settings are recommended:

- 1. Quasi-convex: smartsearch=TRUE (the default).
- 2. Strictly-convex: smartsearch=FALSE and maxit=20.
- 3. Quadratic: smartsearch=FALSE and maxit=1.

Technically, the user template is processed several times by inserting different types as template parameter, selected by argument type:

- "ADFun" Run through the template with AD-types and produce a stack of operations representing the objective function.
- "Fun" Run through the template with ordinary double-types.

6 newton

• "ADGrad" Run through the template with nested AD-types and produce a stack of operations representing the objective function gradient.

Each of these are represented by external pointers to c++ structures available in the environment env.

Further objects in the environment env:

- validpar Function defining the valid parameter region (by default no restrictions). If an invalid parameter is inserted fn immediately return NaN.
- parList Function to get the full parameter vector of random and fixed effects in a convenient list format.
- random An index vector of random effect positions in the full parameter vector.
- last.par Full parameter of the latest likelihood evaluation.
- last.par.best Full parameter of the best likelihood evaluation.
- tracepar Trace every likelihood evaluation?
- tracemgc Trace mgc of every gradient evaluation?
- silent Pass 'silent=TRUE' to all try-calls ?

A high level of tracing information will be output by default when evaluating the objective function and gradient. This is useful while developing a model, but may eventually become annoying. The following will disable all tracing from an object 'obj' returned by 'MakeADFun':

```
• obj$env$tracemgc <- FALSE
```

- obj\$env\$inner.control\$trace <- FALSE
- obj\$env\$silent <- TRUE

#### Value

List with components (fn,gr, etc) suitable for an optim call.

newton

Generalized newton optimizer.

## **Description**

Generalized newton optimizer used for the inner optimization problem.

#### Usage

```
newton(par, fn, gr, he, trace = newtonOption("trace"),
  maxit = newtonOption("maxit"), tol = newtonOption("tol"), alpha = 1,
  smartsearch = newtonOption("smartsearch"),
  mgcmax = newtonOption("mgcmax"), super = TRUE, silent = TRUE,
  ustep = 1, power = 0.5, u0 = 1e-04, grad.tol = tol, step.tol = tol,
  tol10 = 0.001, env = environment(), ...)
```

newton 7

#### **Arguments**

par Initial parameter.
fn Objective function.
gr Gradient function.

he Sparse hessian function. trace Print tracing information?

maxit Maximum number of iterations.

tol Convergence tolerance.

alpha Newton stepsize in the fixed stepsize case.

smartsearch Turn on adaptive stepsize algorithm for non-convex problems?

mgcmax Refuse to optimize if the gradient is too steep.

super Supernodal Cholesky?

silent Be silent?

ustep Adaptive stepsize initial guess between 0 and 1.

power Parameter controlling adaptive stepsize.

u0 Parameter controlling adaptive stepsize.

grad.tol Gradient convergence tolerance.
step.tol Stepsize convergence tolerance.

Try to exit if last 10 iterations not improved more than this.

env Environment for cached Cholesky factor.

... Currently unused.

#### **Details**

If smartsearch=FALSE this function performs an ordinary newton optimization on the function fn using an exact sparse hessian function. A fixed stepsize may be controlled by alpha so that the iterations are given by:

$$u_{n+1} = u_n - \alpha f''(u_n)^{-1} f'(u_n)$$

If smartsearch=TRUE the hessian is allowed to become negative definite preventing ordinary newton iterations. In this situation the newton iterations are performed on a modified objective function defined by adding a quadratic penalty around the expansion point  $u_0$ :

$$f_t(u) = f(u) + \frac{t}{2} ||u - u_0||^2$$

This functions hessian (f''(u) + tI) is positive definite for t sufficiently large. The value t is updated at every iteration: If the hessian is positive definite t is decreased, otherwise increased. Detailed control of the update process can be obtained with the arguments ustep, power and u0.

#### Value

List with solution similar to optim output.

8 precompile

openmp

Control number of openmp threads.

## **Description**

Control number of openmp threads.

#### Usage

```
openmp(n = NULL)
```

#### **Arguments**

n

Requested number of threads, or NULL to just read the current value.

#### Value

Number of threads.

precompile

Precompile the TMB library in order to speed up compilation of templates.

## **Description**

Precompile the TMB library

#### Usage

```
precompile(...)
```

# **Arguments**

... Passed to compile.

## Details

The precompilation should only be run once, typically right after installation of TMB. Note that the precompilation requires write access to the TMB package folder. Two versions of the library - with/without the openmp flag - will be generated. After this, compilation times of templates should be reduced.

- To precompile on Linux run precompile().
- To precompile on OS X run precompile(PKG\_LIBS = "-install\_name `pwd`/\$@").

Note that precompilation has side effects: It is not possible to work with more than one model at a time for a single R instance.

Rinterface 9

Create minimal R-code corresponding to a cpp template.

# Description

Create a skeleton of required R-code once the cpp template is ready.

# Usage

```
Rinterface(file)
```

# **Arguments**

file

cpp template file.

## **Examples**

```
file <- system.file("examples/simple.cpp", package = "TMB")
Rinterface(file)</pre>
```

runExample

Run one of the test examples.

# Description

Compile and run a test example (runExample() shows all available examples).

# Usage

```
runExample(name = NULL, all = FALSE, thisR = TRUE, clean = FALSE,
  exfolder = NULL, ...)
```

# Arguments

name	Character name of example.
all	Run all the test examples?
thisR	Run inside this R?
clean	Cleanup before compile?
exfolder	Alternative folder with examples.
	Passed to compile.

10 sdreport

sdreport

General sdreport function.

## Description

After optimization of an AD model, sdreport is used to calculate standard deviations of all model parameters, including non linear functions of random and fixed effects parameters specified through the ADREPORT() macro from the user template.

#### Usage

```
sdreport(obj, par.fixed = NULL, hessian.fixed = NULL,
  getJointPrecision = FALSE)
```

#### **Arguments**

obj Object returned by MakeADFun

par.fixed Optional. Fixed effect parameter estimate (will be known to obj when an opti-

mization has been carried out).

hessian.fixed Optional. Hessian wrt. fixed effects (will be calculated from obj if missing).

getJointPrecision

Optional. Return full joint precision matrix of random and fixed effects?

#### **Details**

First, the Hessian wrt. the fixed effect parameter vector  $(\theta)$  is calculated. The fixed effects covariance matrix is approximated by

 $V(\hat{\theta}) = -\nabla^2 l(\hat{\theta})^{-1}$ 

where l denotes the log likelihood function (i.e. -objfn).

For non-random effect models the standard delta-method is used to calculate the covariance matrix. Let  $\phi(\theta)$  denote some non-linear function of  $\theta$ . Then

$$V(\phi(\hat{\theta})) \approx \nabla \phi V(\hat{\theta}) \nabla \phi'$$

For random effect models a generalized delta-method is used. First the joint covariance of random and fixed effects is estimated by

$$V \left( \begin{matrix} \hat{u} \\ \hat{\theta} \end{matrix} \right) \approx \left( \begin{matrix} H_{uu}^{-1} & 0 \\ 0 & 0 \end{matrix} \right) + JV(\hat{\theta})J'$$

where  $H_{uu}$  denotes random effect block of the full joint Hessian of obj\$env\$f and J denotes the Jacobian of  $\begin{pmatrix} \hat{u}(\theta) \\ \theta \end{pmatrix}$  wrt.  $\theta$ . Here, the first term represents the expected conditional variance given the fixed effects and the second term represents the variance of the conditional mean wrt. the fixed effects. Now the delta method can be applied on a general non-linear function  $\phi(u,\theta)$  of random effects u and fixed effects  $\theta$ :

$$V(\phi(\hat{u}, \hat{\theta})) \approx \nabla \phi V \begin{pmatrix} \hat{u} \\ \hat{\theta} \end{pmatrix} \nabla \phi'$$

The full joint covariance is not returned by default, because it may require large amounts of memory. It may be obtained by specifying getJointPrecision=TRUE, in which case  $V\begin{pmatrix} \hat{u} \\ \hat{\theta} \end{pmatrix}^{-1}$  will be part

template 11

of the output. This matrix must be manually inverted using solve(jointPrecision) in order to get the joint covariance matrix. Note, that the parameter order will follow the original order (i.e. obj\$env\$par).

#### Value

Object of class sdreport

# Examples

```
runExample("linreg_parallel",thisR=TRUE) ## Fixed effect example
sdreport(obj)
runExample("rw",thisR=TRUE) ## Random effect example
rep <- sdreport(obj)
summary(rep,"random") ## Only random effects
summary(rep,"fixed",p.value=TRUE) ## Only fixed effects
summary(rep,"report") ## Only report</pre>
```

template

Create cpp template to get started.

## **Description**

Create a cpp template to get started.

## Usage

```
template(file = NULL)
```

# Arguments

file

Optional name of cpp file.

## **Details**

This function generates a c++ template with a header and include statement. Here is a brief overview of the c++ syntax used to code the objective function.

Macros to read data and declare parameters:

C++ type	R type
vector <type></type>	vector
matrix <type></type>	matrix
Type	numeric(1)
int	integer(1)
vector <int></int>	factor
vector <int></int>	integer
Eigen::SparseMatrix <type></type>	dgTMatrix
array <type></type>	array
matrix <type></type>	matrix
vector <type></type>	vector
array <type></type>	array
Type	numeric(1)
	vector <type> matrix<type> Type int vector<int> vector<int> Eigen::SparseMatrix<type> array<type> matrix<type> vector<type> array<type> array<type></type></type></type></type></type></type></int></int></type></type>

TMB-internal

#### Basic calculations:

Template Syntax	Explanation
REPORT(x)	Report x back to R
ADREPORT(x)	Report x back to R with derivatives
vector $<$ Type $>$ v(n1);	R equivalent of v=numeric(n1)
matrix <type> m(n1,n2);</type>	R equivalent of m=matrix(0,n1,n2)
array < Type > a(n1,n2,n3);	R equivalent of $a=array(0,c(n1,n2,n3))$
v+v,v-v,v*v,v/v	Pointwise binary operations
m*v	Matrix-vector multiply
a.col(i)	R equivalent of a[,,i]
a.col(i).col(j)	R equivalent of a[,j,i]
a(i,j,k)	R equivalent of a[i,j,k]
exp(v)	Pointwise math
m(i,j)	R equivalent of m[i,j]
v.sum()	R equivalent of sum(v)
m.transpose()	R equivalent of t(m)

Some distributions are avaliable as c++ templates with syntax close to R's distributions:

#### **Function header**

# dnbinom2(x,mu,var,int give\_log=0) dpois(x,lambda,int give\_log=0) dlgamma(y,shape,scale,int give\_log=0) dnorm(x,mean,sd,int give\_log=0)

#### Distribution

Negative binomial with mean and variance Poisson distribution as in R log-gamma distribution Normal distribution as in R

# **Examples**

template()

TMB-internal	Internal TMB Functions

# Description

Internal TMB functions

# **Details**

These are not to be called by the user (or in some cases are just waiting for proper documentation to be written:).

# **Index**

```
benchmark, 2
checkSparseHessian (TMB-internal), 12
compile, 2
config (TMB-internal), 12
dynlib (TMB-internal), 12
flagsDefaults (TMB-internal), 12
gdbsource, 3
getUserDLL (TMB-internal), 12
grepRandomParameters (TMB-internal), 12
info (TMB-internal), 12
isParallelTemplate(TMB-internal), 12
MakeADFun, 4
newton, 6
newtonDefaults (TMB-internal), 12
newtonOption (TMB-internal), 12
openmp, 8
parallelBenchmark (TMB-internal), 12
plot.parallelBenchmark(TMB-internal),
        12
precompile, 8
print.backtrace(TMB-internal), 12
print.sdreport (TMB-internal), 12
Rinterface, 9
runExample, 9
runSymbolicAnalysis(TMB-internal), 12
sdreport, 10
setDefaults (TMB-internal), 12
sparseHessianFun (TMB-internal), 12
summary.sdreport (TMB-internal), 12
template, 11
TMB-internal, 12
tmbOption (TMB-internal), 12
updateCholesky (TMB-internal), 12
```