Speeding up parameter inference with compiled models

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DE models in R are easy to implement, allow simple interactive development using readable code and access to Rs high-level procedures. All of which were part of our motivation to develop deBInfer as an R based inference package. However, the deSolve package also allows the evaluation of DE models that are defined in lower-level languages such as C and FORTRAN. These compiled models have the benefit of increased simulation speed. As the DE model is evaluated many times during the MCMC procedure, even moderate speed-ups from using compiled models may result in large absolute time savings.

We demonstrate the speed-up for an example from the deSolve documentation (Soetaert et al. 2010, Soetaert et al. (2009))

1 A simple ODE example

Assume the following simple ODE (which is from the LSODA source code):

$$\begin{aligned} \frac{dy_1}{dt} &= -k_1 \cdot y_1 + k_2 \cdot y_2 \cdot y_3 \\ \frac{dy_2}{dt} &= k_1 \cdot y_1 - k_2 \cdot y_2 \cdot y_3 - k_3 \cdot y_2 \cdot y_2 \\ \frac{dy_3}{dt} &= k_3 \cdot y_2 \cdot y_2 \end{aligned}$$

where y_1 , y_2 and y_3 are state variables, and k_1 , k_2 and k_3 are parameters.

We first implement and run this model in pure R, then show how to do this in C.

1.1 ODE model implementation in R

An ODE model implemented in pure R should be defined as:

```
yprime = func(t, y, parms, ...)
```

where t is the current time point in the integration, y is the current estimate of the variables in the ODE system, and parms is a vector or list containing the parameter values. The optional dots argument (...) can be used to pass any other arguments to the function. The return value of func should be a list, whose first element is a vector containing the derivatives of y with respect to time, and whose next elements contain output variables that are required at each point in time.

The R implementation of the simple ODE is given below:

```
modelR <- function(t, Y, parameters) {
  with (as.list(parameters), {
    dy1 = -k1*Y[1] + k2*Y[2]*Y[3]</pre>
```

```
dy3 = k3*Y[2]*Y[2]
dy2 = -dy1 - dy3

list(c(dy1, dy2, dy3))
})
}
```

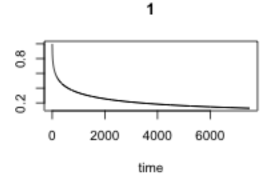
The Jacobian $(\frac{\partial y'}{\partial y})$ associated to the above example is:

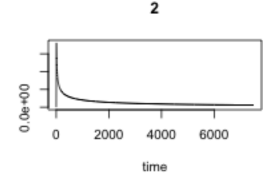
```
jacR <- function (t, Y, parameters) {
    with (as.list(parameters), {

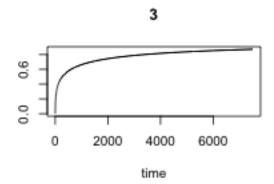
        PD[1,1] <- -k1
        PD[1,2] <- k2*Y[3]
        PD[1,3] <- k2*Y[2]
        PD[2,1] <- k1
        PD[2,3] <- -PD[1,3]
        PD[3,2] <- k3*Y[2]
        PD[2,2] <- -PD[1,2] - PD[3,2]

        return(PD)
    })
}</pre>
```

This model can then be run as follows:







1.2 ODE model implementation in C

In order to create compiled models (.DLL = dynamic link libraries on Windows or .so = shared objects on other systems) you must have a recent version of the GNU compiler suite installed, which is quite standard for Linux. Windows users find all the required tools on [http://www.murdoch-sutherland.com/Rtools/]. Getting DLLs produced by other compilers to communicate with R is much more complicated and therefore not recommended. More details can be found on [http://cran.r-project.org/doc/manuals/R-admin.html].

The call to the derivative and Jacobian function is more complex for compiled code compared to R-code, because it has to comply with the interface needed by the integrator source codes.

Below is an implementation of this model in C:

```
/* file mymod.c */
#include <R.h>
static double parms[3];
#define k1 parms[0]
#define k2 parms[1]
#define k3 parms[2]

/* initializer */
void initmod(void (* odeparms)(int *, double *))
{
   int N=3;
   odeparms(&N, parms);
}

/* Derivatives and 1 output variable */
void derivs (int *neq, double *t, double *y, double *ydot,
```

```
double *yout, int *ip)
{
    if (ip[0] <1) error("nout should be at least 1");</pre>
    ydot[0] = -k1*y[0] + k2*y[1]*y[2];
    ydot[2] = k3 * y[1]*y[1];
    ydot[1] = -ydot[0]-ydot[2];
    yout[0] = y[0]+y[1]+y[2];
}
/* The Jacobian matrix */
void jac(int *neq, double *t, double *y, int *ml, int *mu,
           double *pd, int *nrowpd, double *yout, int *ip)
{
  pd[0]
                      = -k1;
  pd[1]
                      = k1;
                      = 0.0;
 pd[2]
  pd[(*nrowpd)]
                      = k2*v[2];
 pd[(*nrowpd) + 1]
                     = -k2*y[2] - 2*k3*y[1];
  pd[(*nrowpd) + 2]
                     = 2*k3*y[1];
  pd[(*nrowpd)*2]
                      = k2*y[1];
 pd[2*(*nrowpd) + 1] = -k2 * y[1];
 pd[2*(*nrowpd) + 2] = 0.0;
/* END file mymod.c */
```

The structure of the implementation in C is described in detail in Soetaert et al. (2009).

2 Running ODE models implemented in compiled code

To run the model described above, the C code must first be compiled¹. This can be done using the R system statement

```
system("R CMD SHLIB mymod.c")
```

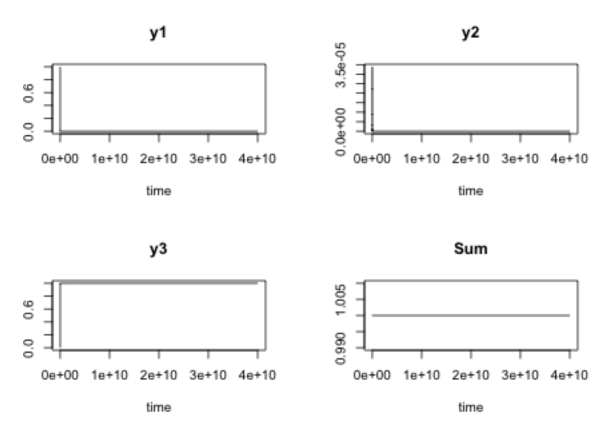
This will create file mymod.dll on Windows, or mymod.so on other platforms.

We load the compiled model with

```
dyn.load(paste("mymod", .Platform$dynlib.ext, sep = ""))
```

The model can now be run as follows:

¹This requires a correctly installed GNU compiler.



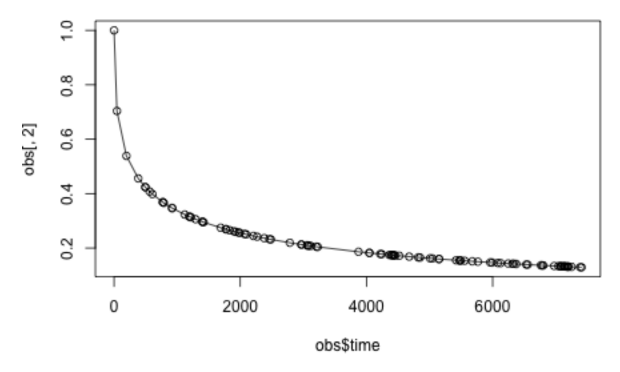
We can determine the speed improvement using the microbenchmark package, which shows us that the C version of the models is solved about 40 times faster than the R version.

```
comp <- microbenchmark::microbenchmark(</pre>
  C = ode(Y, times, func = "derivs", parms = parms,
           jacfunc = "jac", dllname = "mymod",
           initfunc = "initmod", nout = 1, outnames = "Sum"),
  R = ode(Y, times, modelR, parms = parms, jacfunc = jacR)
print(comp, unit = "ms")
## Unit: milliseconds
##
   expr
              min
                                            median
                         lq
                                  mean
                                                           uq
                                                                    max neval
       C 0.43757 0.527498 0.5988335 0.5901185 0.6464945
##
                                                              1.111103
                                                                           100
       R 21.12073 21.963163 24.7362956 24.2408290 24.9036375 81.611968
                                                                           100
##
```

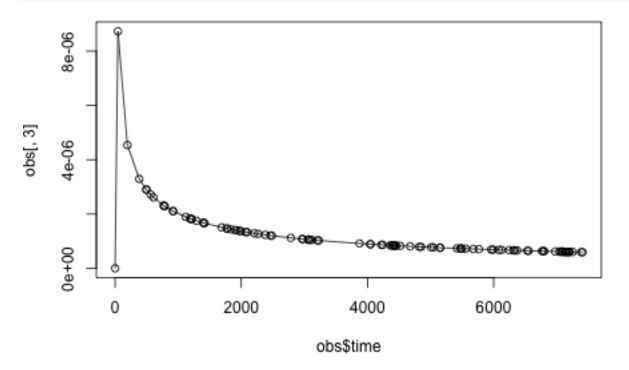
The Again, please refer to the deSolve documentation (Soetaert et al. 2009) for a detailed description of the syntax.

We can now simulate a dataset to demonstrate the inference procedure

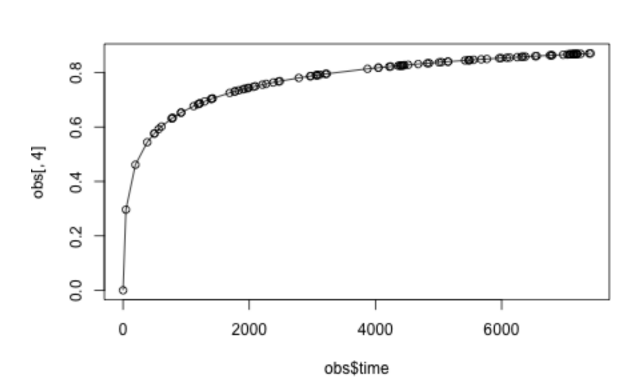
```
set.seed(143)
#force include the first time-point (t=0)
obs <- as.data.frame(outR[c(1,runif(100, 0, nrow(outR))),])
obs <- obs[order(obs$time),]
plot(obs$time, obs[,2], type='o')</pre>
```



plot(obs\$time, obs[,3], type='o')



plot(obs\$time, obs[,4], type='o')



3 Defining an observation model and parameters for inference

For simplicity we assume a normal log-likelihood for these data

$$\ell(\mathcal{Y}|\boldsymbol{\theta}) = \sum_{i} \sum_{t} \ln \left(\frac{1}{\sigma_{obs} \sqrt{2\pi}} \exp\left(-\frac{(\tilde{y}_{t,i} - (y_{t,i})^2}{2\sigma_{obs}^2}\right) \right) \tag{1}$$

where $\tilde{y}_{t,i}$ are the observations, and $y_{t,i}$ are the predictions of the DE model given the current MCMC sample of the parameters $\boldsymbol{\theta}$.

```
# the observation model
obs_model <- function(data, sim.data, samp){

llik.y1 <- sum(dnorm(obs[,2], mean = sim.data[,2], sd = samp[['sd.y1']], log = TRUE))

llik.y2 <- sum(dnorm(obs[,3], mean = sim.data[,3], sd = samp[['sd.y2']], log = TRUE))

llik.y3 <- sum(dnorm(obs[,4], mean = sim.data[,4], sd = samp[['sd.y3']], log = TRUE))

return(llik.y1 + llik.y2 + llik.y3)
}</pre>
```

We declare the DE model parameter r, assign a prior $r \sim \mathcal{N}(0,1)$ and a random walk sampler with a Normal kernel (samp.type="rw") and proposal variance of 0.005. Similarly, we declare $K \sim \ln \mathcal{N}(1,1)$ and $\ln(\sigma_{obs}^2) \sim \mathcal{N}(0,1)$. Note that we are using the asymmetric uniform proposal distribution $\mathcal{U}(\frac{a}{b}\theta^{(k)}, \frac{b}{a}\theta^{(k)})$ for the variance parameter (samp.type="rw-unif"), as this ensures strictly positive proposals.

Lastly, we provide an initial value $N_0 = 0.1$ for the DE:

```
y1 <- debinfer_par(name = "y1", var.type = "init", fixed = TRUE, value = 1)
y2 <- debinfer_par(name = "y2", var.type = "init", fixed = TRUE, value = 0)
y3 <- debinfer_par(name = "y3", var.type = "init", fixed = TRUE, value = 0)</pre>
```

The declared parameters are then collated using the setup_debinfer function. Note that for models with more than one state variable, the initial values must be entered in the same order, as they are specified in the DE model function, as the solver matches these values by position, rather than by name. More details can be found in ?deSolve::ode. The remaining parameters can be entered in any order.

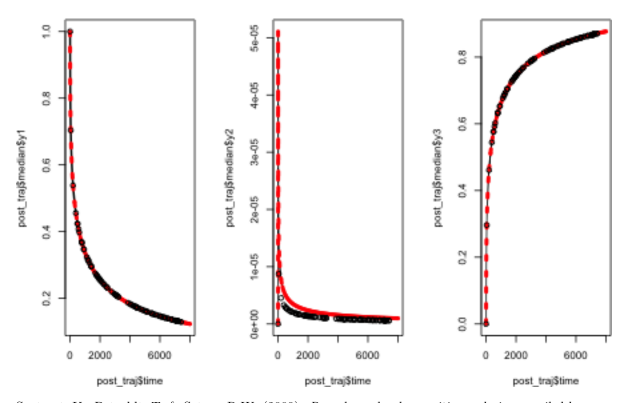
```
mcmc.pars <- setup_debinfer(k1, k2, k3, sd.y1, sd.y2, sd.y3, y1, y2, y3)
```

4 Conduct inference

Finally we use deBInfer to estimate the parameters of the original model. de_mcmc is the workhorse of the package and runs the MCMC estimation. The progress of the MCMC procedure can be monitored using the cnt, plot and verbose options: Every cnt iterations the function will print out information about the current state, and, if plot=TRUE, traceplots of the chains will be plotted. Setting verbose=TRUE will print additional information. Note that frequent plotting will substantially slow down the MCMC sampler, and should be used only on short runs when tuning the sampler.

```
## Order of initial conditions is y1, y2, y3
## initial posterior probability = -372.440801942676
```

```
Ct <- system.time(mcmc_samplesC <- de_mcmc(N = iter, data = obs, de.model = "derivs",
           jacfunc = "jac", dllname = "mymod",
           initfunc = "initmod", nout = 1, outnames = "Sum",
                          obs.model = obs_model, all.params = mcmc.pars,
                          Tmax = max(obs$time), data.times = obs$time, cnt = 50,
                          plot = FALSE, solver = "ode", verbose.mcmc = FALSE))
## Order of initial conditions is y1, y2, y3
## initial posterior probability = -372.440801942676
print(Rt)
     user system elapsed
## 51.352
           0.494 54.747
print(Ct)
##
     user system elapsed
##
     6.351
           0.044 6.552
system.time(post_traj <- post_sim(mcmc_samples, n=100, times=0:8000, burnin=100, output = 'all', prob =
##
      user system elapsed
           0.503 36.137
## 32.406
system.time(post_trajC <- post_sim(mcmc_samplesC, n=100, times=0:8000, burnin=100, output = 'all', prob
##
     user system elapsed
##
     3.161
           0.245
                    3.435
par(mfrow=c(1,3))
plot(post_traj$time, post_traj$median$y1, type='l',lwd=2)
lines(post_trajC$time, post_trajC$median$y1, col="red", lty=3, lwd=4)
points(obs$time, obs[,2])
plot(post_traj$time, post_traj$median$y2, type='1')
lines(post_trajC$time, post_trajC$median$y2, col="red", lty=3, lwd=4)
points(obs$time, obs[,3])
plot(post_traj$time, post_traj$median$y3, type='1')
lines(post_trajC$time, post_trajC$median$y3, col="red", lty=3, lwd=4)
points(obs$time, obs[,4])
```



Soetaert, K., Petzoldt, T. & Setzer, R.W. (2009). R package desolve, writing code in compiled languages. Soetaert, K., Petzoldt, T. & Setzer, R.W. (2010). Solving differential equations in R: Package deSolve. Journal of Statistical Software, 33, 1–25.