Welcome to stock assessment via TMB

Anders Nielsen an@aqua.dtu.dk

Welcome!

- Please introduce yourselves
 - Name and Organization
 - Experience (TMB, ADMB, R/C coding)
 - What do you hope to learn
 - Special requests

Non-standard models

- Models where you need to write your own likelihood
- Models you cannot write in one line in R
- non-trivial non-linearities
- complex covariance structures
- complicated couplings between fixed and random effects
- different sources of observations needing different likelihood types
- Standard models are very useful, but should not limit us

Formula interfaces are sometimes frustrating ...

• A useful model for longitudinal data:

```
\begin{array}{lll} & \mathbf{lnc} & \sim & N(\mu,\mathbf{V}), \text{ where} \\ & \mu_i & = & \mu + \alpha(\mathtt{treatm}_i) + \beta(\mathtt{month}_i) + \gamma(\mathtt{treatm}_i,\mathtt{month}_i), \text{ and} \\ & & & \left\{ \begin{array}{ll} 0 & , & \text{if } \mathsf{cage}_{i_1} \neq \mathsf{cage}_{i_2} \text{ and } i_1 \neq i_2 \\ \nu^2 + \tau^2 \exp\left\{\frac{-(\mathtt{month}_{i_1} - \mathtt{month}_{i_2})^2}{\rho^2}\right\} & , & \text{if } \mathsf{cage}_{i_1} = \mathsf{cage}_{i_2} \text{ and } i_1 \neq i_2 \\ \nu^2 + \tau^2 + \sigma^2 & , & \text{if } i_1 = i_2 \end{array} \right.
```

• This model is implemented by:

- So many pitfalls and much is hidden. Even difficult to recover model parameters.
- What is τ ? Some hours with manual, but re-implement to be sure...
- Restricted by what someone else has put in there. Giant task to move beyond.

Terbuthylazine

- It is a herbicide
- Free terbuthylazine can be washed into the drinking water
- It can be bound to the soil
- Certain bacterias can mineralize it



$$\frac{dB_t}{dt} = -k_1 B_t + k_2 F_t, B_0 = 0$$

$$\frac{dF_t}{dt} = k_1 B_t - (k_2 + k_3) F_t, F_0 = 100$$

$$\frac{dM_t}{dt} = k_3 F_t, M_0 = 0$$

Simplifying



- The system is closed, so $M_t = 100 B_t F_t$
- Define $X_t = \begin{pmatrix} B_t \\ F_t \end{pmatrix}$
- The simplified system is:

$$\frac{dX_t}{dt} = \underbrace{\begin{pmatrix} -k_1 & k_2 \\ k_1 & -(k_2 + k_3) \end{pmatrix}}_{A} X_t, \qquad X_0 = \begin{pmatrix} 0 \\ 100 \end{pmatrix}$$

• The system is linear, so it can be be solved for instance via the matrix exponential

$$X_t = e^{At} X_0$$

Observations

• The amount of mineralized terbuthylazine was measured 26 times throughout a year

Time	\mathbf{M}	•											
0.77	1.396	•	16 -										
1.69	3.784										0	•	
2.69	5.948		14 - 12 -			· ·							
3.67	7.717				, ° ° ° ° ° °	0	0	0	0				
4.69	9.077				- 00000								
5.71	10.100			O	50								
7.94	11.263			0									
9.67	11.856			0									
11.77	12.251		40	0									
17.77	12.699		10 - 8 -	U									
23.77	12.869	(%		0									
32.77	13.048	(%) W											
40.73	13.222	_		0									
47.75	13.347												
54.90	13.507		6 -	0									
62.81	13.628												
72.88	13.804		4 -										
98.77	14.087			0									
125.92	14.185												
160.19	14.351												
191.15	14.458		2 -	_									
223.78	14.756			0									
287.70	15.262			1	1			1	1			I	
340.01	15.703			0	50	100	1	150	200	250	300	350	
340.95	15.703												
342.01	15.703	Time (days)											

Simplest statistical model

• The simplest model we can think of would be:

$$M_{t_i} \sim \mathcal{N}\left(100 - \sum X_{t_i}, \sigma^2\right)$$
, independent, and with $X_{t_i} = e^{At_i}X_0$.

```
> library(Matrix)
> nlogL <- function(theta) {</pre>
      k <- exp(theta[1:3])
      sigma <- exp(theta[4])</pre>
      A \leftarrow rbind(c(-k[1], k[2]), c(k[1], -(k[2] + k[3])))
    x0 < -c(0, 100)
      sol <- function(t) 100 - sum(expm(A * t) %*% x0)
      pred <- sapply(dat[, 1], sol)</pre>
      -sum(dnorm(dat[, 2], mean = pred, sd = sigma, log = TRUE))
+ }
> system.time(fit <- optim(c(-2, -2, -2, -2), nlogL, hessian = TRUE))
         system elapsed
   user
 19.409
         0.004 19.428
> fit$value
[1] 19.26905
> fit$convergence
[1] 0
```

• Try some of the different minimizers in R

```
> library(optimx)
> fit<-optimx(c(-2,-2,-2,-2),nlogL,hessian=TRUE,control=list(all.methods=TRUE))
> fit
  fvalues
              method fns grs conv KKT1 KKT2 xtimes
 153.3056
              bobyqa 144 NA
                                O TRUE FALSE 9.629
              Rcgmin 85 50
 102.7661
                                O TRUE FALSE 22.314
 102.7660
                 nlm
                      NA
                          NA
                                   TRUE FALSE 11.865
 102.7660
                BFGS
                          18
                                   TRUE FALSE 15.005
                      79
 102.7660
              Rvmmin 81
                          15
                                   TRUE FALSE 10.517
 102.7660
                  CG 567 101
                                   TRUE FALSE 91.569
91.17466
              newuoa 696
                          NA
                                0 TRUE FALSE 46.063
 19.26905 Nelder-Mead 223
                                O FALSE FALSE 14.837
                          NA
0.9392184
                                         TRUE 14.953
              ucminf 40
                          40
                                O FALSE
0.9392142
                 spg 198
                               0 FALSE TRUE 55.732
                          NA
0.9392142
            L-BFGS-B 85
                          85
                                         TRUE 50.807
                                O FALSE
0.9392142
              nlminb 33 128
                                O TRUE
                                         TRUE 10.729
```

- Difficult because it is non-linear
- Would possibly be helped by accurate gradient info
- Runs in a fraction of a second in TMB (exercise)
- Notice this is a miniature example with only 4 parameters

What is needed to handle a non-standard problem

- A purely parametric assessment model has more than 100 model parameters and it is non-linear
- Code up the negative log likelihood function
- A good function minimizer

AD aided minimizer

• Want to minimize the negative log likelihood w.r.t. $\theta = (\theta_1, \dots, \theta_n)$

$$\widehat{\theta} = \operatorname*{argmin}_{\theta} \ell(y|\theta)$$

- If the dimension of θ is low (say n less than 5) any method can be used (grid search, random search, finite difference approximations, ...)
- We would like to be able to handle much larger problems
- Important for fixed effects models, and even more for random effects models
- A quasi-Newton minimizer aided by automatic differentiation

Quasi-Newton minimizer



Automatic Differentiation Model Builder

- A Newton minimizer is an iterative algorithm
- Each step assumes that the function $\ell(x,\theta)$ can be approximated locally by a quadratic function
- It uses the first ℓ'_{θ} and second ℓ''_{θ} derivatives to find the minimum
- Instead of calculating ℓ''_{θ} at every step, a quasi-Newton minimizer uses successive first derivatives ℓ'_{θ} to approximate ℓ''_{θ}
- So a fast and accurate way to calculate ℓ'_{θ} is needed

Automatic Differentiation

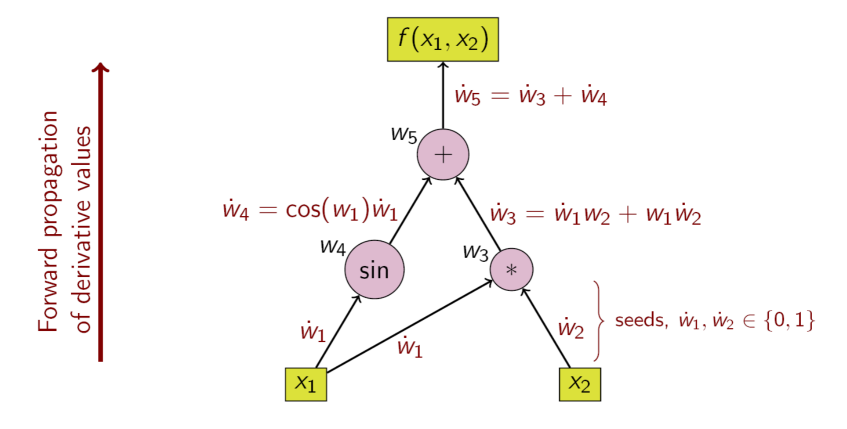
- We need to write a program to compute $\ell(\theta, x)$ anyway
- A computer program is a long list of simple operations: '+', '-', '*', '/', 'exp', 'log', 'sin', 'cos', 'tan', 'sqrt', and so on
- We know how to derive each of these operations
- The chain rule tells us how to combine: (f(g(x)))' = f'(g(x))g'(x)
- So if the computer is instructed to:
 - keep track of all the simple operations used when calculating $\ell(\theta, x)$
 - use the simple derivative formulas and the chain rule
- Then once $\ell(\theta, x)$ is computed, we also have ℓ'_{θ} with a minimum of extra calculations
- This is fast and accurate, and the difficult part is built into TMB(!)
- Alternatives:
 - Finite difference: $(\ell'_{\theta})_i \approx \frac{\ell(\theta_i + \Delta\theta_i, x) \ell(\theta, x)}{\Delta\theta_i}$ Simple, but slow and inaccurate
 - Analytical: Excellent option, but difficult in larger models

```
#include <math.h>
#include <iostream.h>
class result {
  private: double v,d;
 public: result(){v = 0;d= 0;};
          result(double val){v = val; d = 0;};
          result(double val, double der) {v = val; d = der;};
          double Value(){return v;};
          double Der(){return d;};
};
class parameter: public result {
  public: parameter(double pval) : result(pval,1.0) {};
          parameter() : result(0.0,1.0) {};
};
result sin(result n){
  return result(sin(n.Value()), cos(n.Value())*n.Der());
};
result operator*(result n1, result n2){
  return(result(n1.Value()*n2.Value(), n1.Der()*n2.Value() + n2.Der()*n1.Value()));
};
ostream& operator << (ostream& o, result n) {
  o << n.Value() << " (Derative: " << n.Der() << ") ";
  return o;
int main(int argc, char* argv[]){
  parameter theta(2);
  result y;
 v = sin(theta*theta);
  cout << "The result is " << y << endl;</pre>
}
```

cpp/ad.cpp

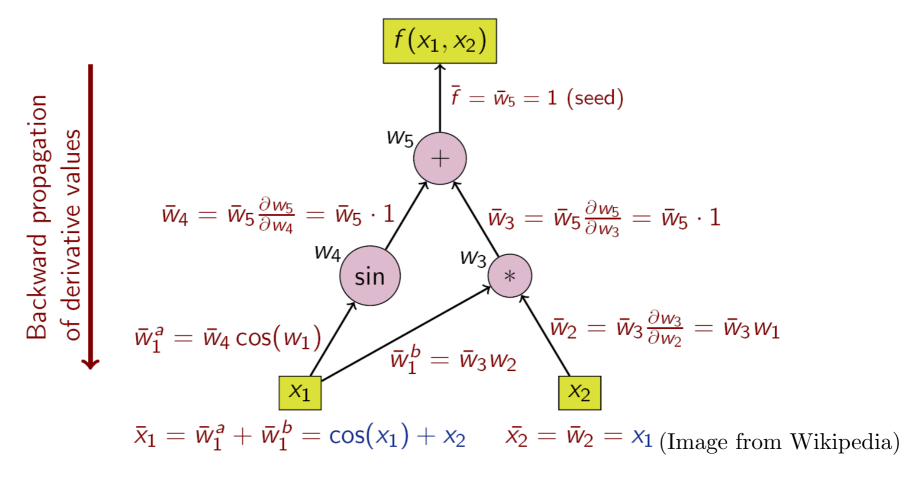
The result is -0.756802 (Derivative: -2.61457)

Forward and reverse mode



(Image from Wikipedia)

- Forward mode is easy to understand and implement
- Not efficient when θ is high dimensional



- Requires recording a stack of all operations
- Efficient in number of operations $(C(\ell'_{\theta}) < 4C(\ell)^{\mathbf{a}})$
- TMB uses reverse mode
- Except for random effects models where a combo of forward and reverse mode is used

^aGriewank, A., 2000. Evaluating Derivatives: Principles and Techniques of Algorithmic Differentiation. SIAM, Philadelphia, PA.

Template Model Builder (TMB):

- Developered by Kasper Kristensen (DTU-Aqua)
- ADMB inspired R-package
- Combines external libraries: CppAD, Eigen, CHOLMOD
- Continuously developed since 2009
- Implements Laplace approximation for random effects
- C++ Template based
- Automatic sparseness detection
- Parallelism through BLAS
- Parallel user templates
- Parallelism through parallel package

Example

• Assume that these 15 numbers follow a negative binomial distribution:

13 5 28 28 15 4 13 4 10 17 11 13 12 17 3

• The TMB code becomes

nbin.cpp

```
library(TMB)
compile("nbin.cpp")
dyn.load(dynlib("nbin"))
dat <- list()</pre>
dat\$Y \leftarrow c(13, 5, 28, 28, 15, 4, 13, 4,
            10, 17, 11, 13, 12, 17, 3)
par <- list()</pre>
par$logsize <- 0
par$p <- 0.5
obj <- MakeADFun(dat, par, DLL="nbin")</pre>
opt <- nlminb(obj$par, obj$fn, obj$gr)</pre>
summary(sdreport(obj))
```

nbin.R.

Exercise 1:

- Installing TMB on your computer via the instructions on (if you have not already): https://github.com/kaskr/adcomp/wiki/Download
- Try replicating the negative binomial example

Final comments:

- Fisheries research has inspired some tools which are useful for statisticians
- These tools have made the jump from standard to non-standard models smaller
- Writing own models give greater insights
- Read more in:
 - Fournier DA, HJ Skaug, J Ancheta, J Ianelli, A Magnusson, MN Maunder, A Nielsen, J Sibert 2012. AD
 Model Builder: using automatic differentiation for statistical inference of highly parameterized complex nonlinear models. Optimization Methods and Software 27 (2), 233-249
 - Kristensen, K, A. Nielsen, C.W. Berg, H.J. Skaug, B. Bell 2016. TMB: Automatic differentiation and laplace approximation. Journal of Statistical Software 70 (5), 1-21
 - Nielsen, A. and C.W. Berg 2014. Estimation of time-varying selectivity in stock assessments using state-space models. Fisheries Research 158, 96-101
 - Thygesen, U.H., C.M. Albertsen, C.W. Berg, K. Kristensen, and A. Nielsen 2017. Validation of state space models fitted as mixed effects models. (Subm. EES).
 - http://tmb-project.org
 - http://admb-project.org
 - https://github.com/fishfollower/SAM