Fitting: simple approaches

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Fitting: simple approaches

- Overview
- Stochastic simulation
  - Discrete time
  - Continuous time
- 3 Fitting: simple approaches
  - Trajectory matching
  - Gradient matching
  - Comparison

Fitting: simple approaches

- Overview

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Typical stats	Typical math
stochastic	deterministic
static	dynamic
phenomenological	mechanistic

- Time-series models: mostly phenomenological and linear (e.g. ARIMA, spectral/wavelet analyses)
- Biomath models: mostly mechanistic and nonlinear (e.g. Lotka-Volterra, SIR, Fitzhugh-Nagumo)

### Modeling

- time: continuous or discrete
- state: continuous (e.g. quantitative genetics) or discrete (e.g. Mendelian)
- evolution: deterministic or stochastic

#### e.g.

ODEs: continuous-time, continuous-state, deterministic

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 branching processes: continuous-time, discrete-state, stochastic

- For stochastic models need to define both a process model and an observation model (= measurement model)

  Process model  $Y(t+1) \sim F(Y(t))$ Measurement model  $Y_{\text{obs}}(t) \sim Y(t)$
- Only process error affects the future dynamics of the process (usually)
- Might decompose process model into a deterministic model for the expectation and (additive?) noise around the expectation: e.g.  $Y(t) = \mu + \epsilon$ ,  $Y(t) \sim \text{Poisson}(\exp(\eta))$

Measurement model  $Y_{obs}(t) \sim Y(t)$ 

Overview

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#### Consequences

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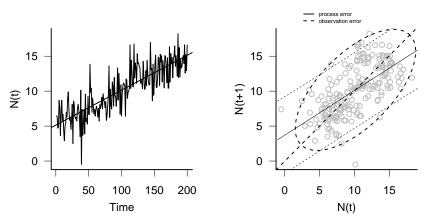
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 Simple statistical methods (i.e. uncorrelated, equal variance) are incorrect

#### Linear example



How should we interpret this single realization?

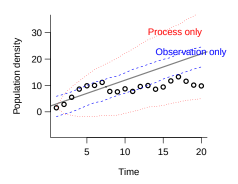
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- Stochastic simulation
  - Discrete time
  - Continuous time

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- understand dynamics
- test methods in best-case scenario
- explore precision/power
- quantify properties of statistical estimators
- evaluate robustness

#### Linear model



$$egin{aligned} & extstyle N(1) = a \ & extstyle N(t+1) \sim extstyle extstyle Normal(N(t)+b, \sigma_{ extstyle proc}^2) \ & extstyle N_{ extstyle Obs}(t) \sim extstyle Normal(N(t), \sigma_{ extstyle obs}^2) \end{aligned}$$

# R code (version 1)

```
## set up parameters etc.
nt <- 20: a <- 6: b <- 1
sd_proc <- sqrt(2)
sd_obs <- sqrt(2)
N <- Nobs <- numeric(nt)
set.seed(101) ## for reproducibility
## actual model
N[1] \leftarrow a
Nobs[1] \leftarrow rnorm(1,N[1],sd_obs)
for (i in 1:nt) {
  N[i+1] \leftarrow rnorm(1,N[i]+b,sd_proc)
  Nobs[i+1] \leftarrow rnorm(1,N[i+1],sd_proc)
```

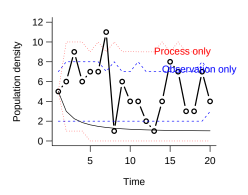
# R code (version 2)

```
library(deSolve)
linfun <- function(t,y,parms) {</pre>
    ## with() is magic to use param names directly
    g <- with(as.list(c(y,parms)), {
        N_new <- rnorm(1,mean=N+b,sd=sd_proc)</pre>
        c(N=N_new, Nobs=rnorm(1, mean=N_new, sd=sd_obs))
  })
  return(list(g)) ## deSolve needs this format
set.seed(101)
NO <- c(N=a, Nobs=rnorm(1,a,sd_obs))
linparms <- c(a=6,b=1,sd_proc=sd_proc,sd_obs=sd_obs)</pre>
ode(NO,1:nt,linfun,linparms,method="iteration")
```

For this particular example, we can cheat because the process error doesn't affect the future dynamics — it just accumulates:

```
N \det <- a+b*(0:(nt-1))
set.seed(101) ## for reproducibility
proc_noise <- rnorm(nt-1,mean=0,sd=sd_proc)</pre>
N <- N_det+cumsum(c(0,proc_noise))</pre>
N_obs <- rnorm(nt,mean=N,sd=sd_obs)</pre>
```

#### Hyperbolic nonlinear model



$$egin{aligned} \mathcal{N}(1) &= \mathcal{N}_0 \ \mathcal{N}(t+1) \sim \mathsf{Poisson}\left(rac{a\mathcal{N}(t)}{b+\mathcal{N}(t)}
ight) \ \mathcal{N}_{\mathsf{obs}}(t) \sim \mathsf{Binomial}(\mathcal{N}(t),p) \end{aligned}$$

#### R code

```
hypfun <- function(t,y,parms) {</pre>
    g <- with(as.list(c(y,parms)), {</pre>
         N_{det} \leftarrow a*N/(b+N)
         N <- rpois(1,lambda=N_det)</pre>
         N_obs <- rbinom(1,size=N,prob=prob_obs)</pre>
         c(N=N.Nobs=N obs)
  })
  return(list(as.numeric(g))) ## deSolve needs numeric() (
set.seed(101)
NO \leftarrow c(N=4, N_{obs}=4)
hypparms \leftarrow c(a=6,b=1,prob_obs=0.9)
ode(NO, times=1:nt, func=hypfun,
    parms=hypparms, method="iteration")
```

#### Stochastic ODEs

- continuous-time, continuous-state
- ordinary differential equations plus dW (= derivative of a Brownian motion/Wiener process)
- delicate analysis (For biologists: Turelli (1977); Roughgarden (1995). For mathematicians: Øksendal (2003))

- More common for cellular/physiological than population models
- Solve via Euler-Maruyama (= Euler + appropriately scaled Gaussian noise)

### continuous-time, discrete-state

 specify (limits of) probabilities of transitions per unit time, e.g.  $P(N \rightarrow N+1)$  in the interval (t, t+dt) is rN(t) dt

- Even harder than SDEs to analyze rigorously . . .
- But computationally straightforward: Gillespie algorithm and variations (Gillespie, 2007): exponentially distributed time between transitions

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• Easiest: simulate the deterministic version of the model (i.e., with neither observation nor process error) and compare

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- Because measurement/observation error is (typically) independent at each observation, overall log-likelihood is sum of log-likelihood
- for Normally distributed, equal-variance error, maximum likelihood estimation equivalent to least-squares
- Very common for ODE models, e.g. Gani and Leach (2001); van Veen et al. (2005)
- Brute force can be slow/unstable: use sensitivity equations (Raue et al., 2013)

#### Pseudo-code

```
## deterministic dynamics:
## function of parameters, possibly including ICs
determ_fun <- function(determ_params) {</pre>
    ## code...
## objective function (neg. log-likelihood, SSQ, ...)
## 'params' includes process and observation parameters
obj_fun <- function(params,data) {</pre>
  estimate <- determ_fun(params[determ_params]))</pre>
  obj <- likfun(estimate,data,params[obs_params])
  return(obj)
find_minimum(obj_fun,starting_params,...)
```

# Real code #1 (for loops)

```
determ_fun <- function(p,nt) {</pre>
  with (as.list(p), a+b*(1:nt))
}
obj_fun <- function(p,nt,Nobs) {
  estimate <- determ_fun(p[c("a", "b")],nt)
  ## negative log-lik. of Normal
  obj <- -sum(dnorm(Nobs, estimate, p["sd"], log=TRUE))
  return(obj)
optim(fn=obj_fun,par=c(a=5,b=2,sd=1),nt=20,Nobs=linN)
```

# Real code #2 (using mle2())

```
library(bbmle)
obj_fun <- function(a,b,sd,nt,Nobs) {
  estimate <- determ_fun(a,b,nt)
  ## negative log-lik. of Normal
  obj <- -sum(dnorm(Nobs, estimate, sd, log=TRUE))</pre>
 return(obj)
determ_fun <- function(a,b,nt) a+b*(1:nt)
mle2(obj_fun,
     data=list(Nobs=linN.nt=nt).
     start=list(a=5,b=2,sd=1.01).
     method="Nelder-Mead")
```

mle2() simplifies computation of confidence intervals, likelihood profiles, etc..

Formula interface further simplifies getting predicted values, etc. (but may make debugging harder!)

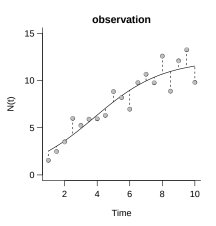
- wrapper for optim
- assumes objective function is negative log-likelihood

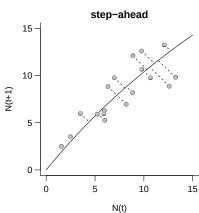
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- uses method="BFGS" by default (maybe switch to Nelder-Mead)
- unlike optim, obj. function takes parameters separately: objfun(alpha, beta) instead of objfun(params)
- use trace=TRUE to track parameters and obj fun value
- nicer accessors (coef(), logLik(), etc.: see methods(class="mle2"))

# Logistic model fit





- use sensible starting values
  - for GA/MCMC, use values that are different (allow exploration) but not crazy (crash/get stuck)
- Nelder-Mead is slower and more robust than BFGS
- test objective/mean function externally
- use cat() to print parameter values, see where you're running into trouble
- use constraints (method="L-BFGS-B", or BOBYQA from nloptr package) or transform parameters (e.g. fit  $\log(\beta)$  rather than  $\beta$ )

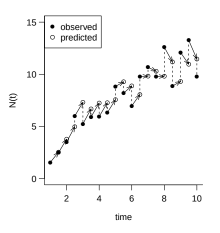
## Gradient matching

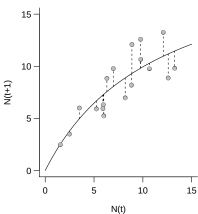
- Next-easiest approach: assume only process error (no measurement error)
- N(t+1) depends only on N(t) (which we know exactly): conditional independence

- One-step-ahead prediction
- Simple for discrete-time models (we need to specify  $N(t+1) \sim N(t)$  anyway)
- More complicated for continuous-time models (Ellner et al., 2002): fit a smooth curve to data, then fit to derivatives of the curve

```
## deterministic dynamics:
## function of parameters and previous values
onestep_fun <- function(determ_params,Nt) { ... }</pre>
## objective function (neg. log-likelihood, SSQ, ...)
obj_fun <- function(params,data) {</pre>
  obj <- ... ## numeric vector of length (nt-1)
 for (i in 1:(nt-1)) {
     estimate <- onestep_fun(N[i],params[determ_params])</pre>
     obj[i] <- fun(estimate, N[i+1], params[obs_params])
  }
  return(sum(obj))
find_minimum(obj_fun,starting_params,...)
```

# Logistic growth fit





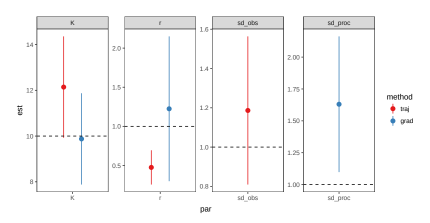
#### How can we use these?

 Try both and hope the answers are not importantly different . . .

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 Use biological knowledge of whether process ≫ observation error or vice versa



#### References

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