Basic SIR fitting

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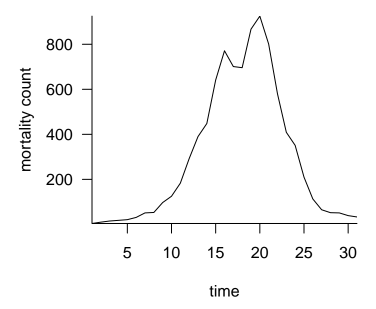
This has been done a million times, but let's try to do it in a reasonably systematic way that could be used in a pedagogical paper.

```
library("fitsir")
library("bbmle") ## need this for now, for coef()
library("plyr")
library("reshape2")
library("ggplot2"); theme_set(theme_bw())
library("RColorBrewer")
```

The current version of fitsir assumes that time and prevalence are stored as columns twee and count within a data frame. Since the bombay data set instead has week (week of epidemic) and mort (mortality), we'll rename it for convenience. (We will for now resolutely ignore issues about fitting weekly mortality counts as prevalences ...)

```
bombay2 <- setNames(bombay,c("tvec","count"))</pre>
```

```
plot(count~tvec,data=bombay2,
          type="l",xaxs="i",yaxs="i",
          xlab="time",ylab="mortality count")
```



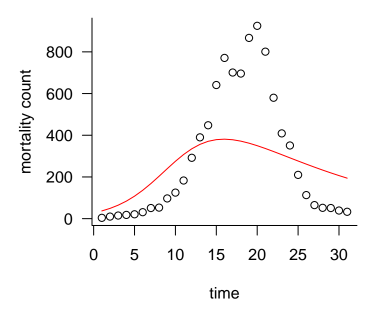
1 Fit the model to the data

Basic fit:

```
m1 <- fitsir(data=bombay2)
```

```
summarize.pars(coef(m1))
## R0 r infper i0
## 5.13774293 0.28629956 14.45249473 0.05235677
```

Seemingly reasonable answers, but \dots



2 Troubleshooting

We're obviously not getting a good answer here. When this happens there are a variety of possibilities.

- optimizer getting stuck
- a small number of local optima
- a large number of local optima, on many different scales (fractal-like or rugged surface)
- a large number of local optima, all similar in scale/height ("fakir's bed" geometry)

Some solutions:

- center/scale parameters and/or reparameterize the model to remove correlation and equalize scales of variation in different parameters
- try to come up with a rule for finding better starting values ("self-starting" fits)

- use a better/more robust local optimizer
- use lots of starting values, randomly or regularly distributed
- use a stochastic global optimizer

```
confint(m1,method="quad")

## Warning: NaNs produced

## 2.5 % 97.5 %

## log.beta -1.453650 -0.6148568

## log.gamma -2.748828 -2.5929057

## log.N NaN NaN

## logit.i -4.049810 -1.7419839
```

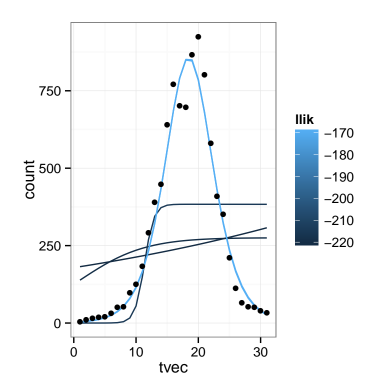
Suggests some sort of unidentifiability ...

What if we try a bunch of starting values?

A crude Latin-hypercube-like strategy: pick evenly spaced values on sensible log scales, then permute to get random (but even) coverage of the space.

```
qlhcfun <- function(n=5,seed=NULL) {</pre>
    require("plyr")
    if (!is.null(seed)) set.seed(seed)
    R0vec \leftarrow 1+10^seq(-1,1.5,length=n)
    infpervec <- sample(10^seq(-1,2,length=n))</pre>
    Nvec <- sample(10^seq(2,5,length=n))</pre>
    i0vec <- sample(10^seq(-3,-1,length=n))</pre>
    startlist <- alply(cbind(R0=R0vec,infper=infpervec,N=Nvec,i0=i0vec),1,
                         function(x) {
                             with(as.list(x), {
                                 beta <- RO/infper
                                  gamma <- 1/infper
                                  c(log.beta=log(beta),log.gamma=log(gamma),
                                    log.N=log(N),logit.i=qlogis(i0))
                             })
                         })
    return(startlist)
startlist <- qlhcfun(n=5, seed=101)
```

```
fitlist <- llply(startlist,fitsir,data=bombay2,
    method="Nelder-Mead",control=list(maxit=1e5))</pre>
```



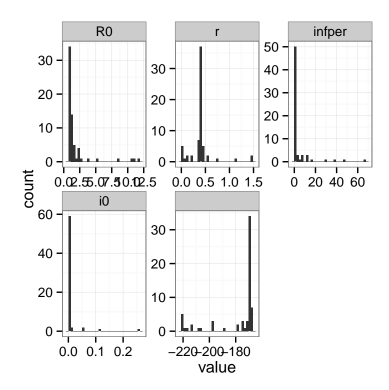
Now try a much larger sample:

```
testOK <- function(x,max.R0=100,max.r=1000,max.infper=400) {
   if (is.null(x)) return(FALSE)
    ss <- summarize.pars(coef(x))
    return(ss["R0"]<max.R0 & ss["r"]<max.r & ss["infper"] < max.infper)
}
fitlist100.0K <- fitlist100[sapply(fitlist100,testOK)]
length(fitlist100.0K)

## [1] 65

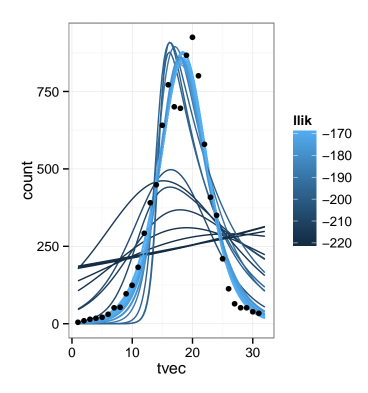
fittab <- laply(fitlist100.0K,function(x) c(summarize.pars(coef(x)),logLik(x)))
ggplot(melt(fittab),aes(x=value))+geom_histogram()+facet_wrap(~Var2,scale="free")

## Warning: position_stack requires constant width: output may be incorrect</pre>
```

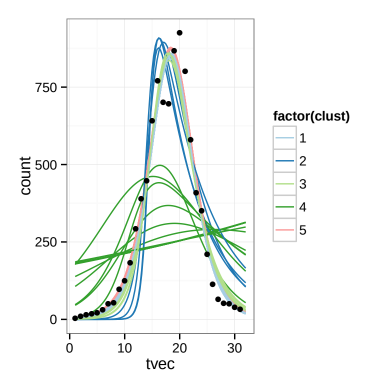


```
likframe100 <- setNames(ldply(fitlist100.0K,logLik),c(".id","llik"))
fittraj100 <- ldply(fitlist100.0K,gettraj,tvec=seq(1,32,length=101))
fitmat100 <- acast(fittraj100,tvec~.id,value.var="count")
fittraj100 <- merge(fittraj100,likframe100)
## plot together</pre>
```

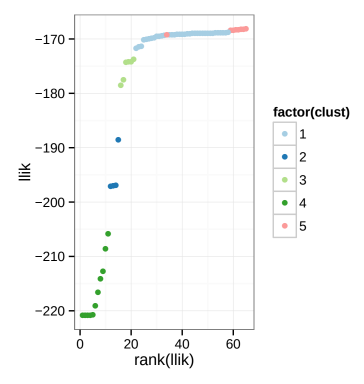
```
ggplot(fittraj100,aes(tvec,count,colour=llik,group=.id))+geom_line()+
    geom_point(data=bombay2,colour="black",aes(group=NA))
```



We can identify clusters \dots



Check out clustering on log-likelihood cumulative distribution curve:



I'm not 100% sure (yet) what this tells us. The clusters aren't so well separated that I necessarily believe that they are distinct modes.

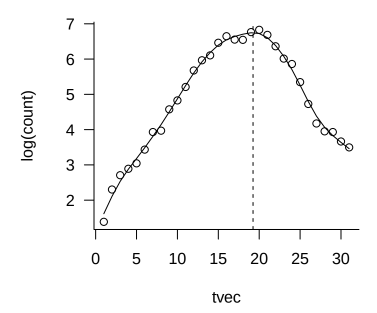
```
startmat100 <- do.call(rbind,startlist100)</pre>
```

to do: characterize starting value sets by cluster (or "bad"), plot, look for regularities. Suspect that large i_0 is a problem?

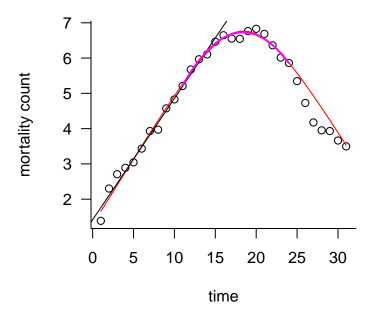
3 Self-starting strategies

Try smooth.spline with spar=0.5 to identify max.; linear regression through times up to 1/2 tmax to identify i(0)N and r; then try a range of other parameters?

```
tvec <- bombay2$tvec
ss <- with(bombay2,smooth.spline(tvec,log(count),spar=0.5))
ss.tmax <- uniroot(function(x) predict(ss,x,deriv=1)$y,c(0,40))$root
plot(log(count)~tvec,data=bombay2)
lines(predict(ss,tvec))
abline(v=ss.tmax,lty=2)</pre>
```



```
ss.thalf <- min(tvec)+(ss.tmax-min(tvec))/2
```



Looks like this works. Is there a way to get a crude starting guess for R_0 and N?

Quadratic fit to peak of log trajectory is very good: what do these parameters tell us about the epidemic?

We want $d^2(\log I)/dt^2$ at the peak ... $d\log I/dt = \beta S/N - \gamma$ and $\hat{S} = \gamma N/\beta$ so we have (at the peak where $d\log I/dt = 0$)

$$d^2 \log I/dt^2 = \beta S'/N = \beta (-\beta SI/N)/N = -\beta^2/N^2 (\gamma N/\beta)I = -\beta \gamma I/N \quad (1)$$

Second derivative of $a + bt + ct^2 = 2c$

OK, I guess, although I expected a little better? Second derivative at the smoothing spline peak is a little bit easier (we don't have to decide on a range

over which to fit the quadratic), and in this case is actually closer to the theoretical value (proportional error 0.11 vs. -0.16) although it's possibly more sensitive to weird shapes at the peak.

This should get us one more parameter.

With Q, a_0 , and b_0 , this gives me so far:

$$a_0 = \log i_0 + \log N \quad \text{(initial number infected, log scale)}$$

$$b_0 = \log \beta - \log \gamma \quad (r) \tag{2}$$

$$\log(-Q) - \log(I_{\max}) = \log \beta + \log \gamma + \log N$$

 $S = \gamma N/\beta$ at the peak time is approximately $N - I_0 - \sum_{t=0}^{\hat{t}} S(t)$ (we have to be careful to decide whether we're counting incidence or prevalence, and correct for γ accordingly: prevalence = incidence/ γ).

Number of counts up to peak:

```
sumcount.tmax <- with(subset(bombay2,tvec<ss.tmax),sum(count))</pre>
```

Should be able to do something with this??

Should use $N(1-i_0)$, not N, as starting condition for S.