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### **Model Calibration in R**

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## Learning Objectives

At the end of this workshop, participants will gain an understanding of the following:

- Concept of model calibration and when it is needed
- Steps and decisions involved in setting up a model calibration
- Different model calibration algorithms and their strengths and weaknesses
- Bayesian model calibration (day 2)
- Implementation of a model calibration in R
  - Can be adapted to calibrate your own models (if using R)
  - Can use as a template for any programming language

### The DARTH Workgroup

- Materials for this workshop were developed in part by the Decision Analysis in R for Technologies in Health (DARTH) Workgroup
- For more information
  - www.darthworkgroup.com
  - Twitter: @DARTHworkgroup

# Why R?

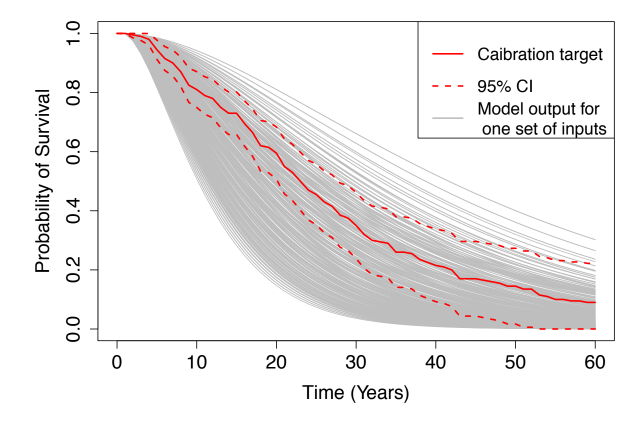
- Open-source, freely available
- Flexible
- Computationally efficient
- Existing packages that implement needed algorithms
- Many other programming languages have these advantages!
  - Can use the calibration framework presented here to write code in your favorite language

#### Motivation

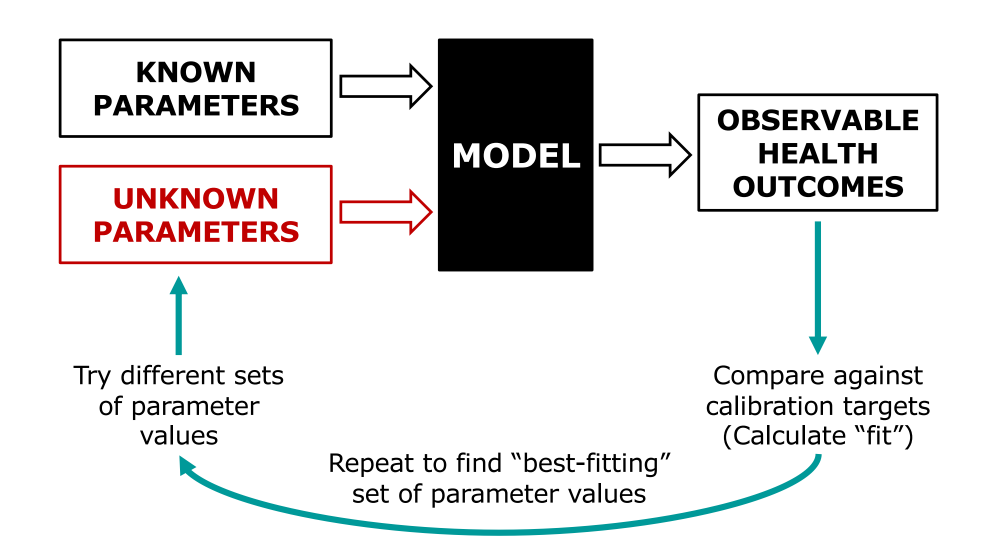
- Mathematical models of disease often involve a subset of parameters whose values are unknown
- Common reasons include physical, feasibility, and ethical limitations
- Estimate values for these parameters by matching model outputs to observed outcomes
  - Model calibration

### Calibration definition

- Process of adjusting model input parameter values to match data on an outcome of interest (e.g., survival, prevalence, or incidence)
- Outcome(s) of interest = "calibration targets"



## Calibration process



# Calibration targets

- Empirical data to be replicated by the model
- Summary statistics (e.g. mean age of cancer diagnosis) or series of observations (e.g. age-specific incidence)
- Can calibrate to multiple targets (e.g. survival and prevalence) simultaneously
- Model should be able to output the outcomes of interest

# Calculating "fit"

- Goodness-of-Fit (GoF) is the quantitative measure of how the model is replicating the target data
- Different ways to measure GoF
  - Distance
  - Likelihood
- Notation
  - *M* : a mathematical model (e.g., Markov model)
  - $\theta$ : Set of K parameters to be calibrated
  - $\phi = M(\theta)$ : Model output for parameter set  $\theta$
  - *y* : Values of *T* calibration targets

#### Distance GoF measures

Sum of squared errors

$$SSE(\theta) = \sum_{i=1}^{T} (y_i - M_i(\theta))^2$$

- Weighted sum of squared errors
  - Assign different weight to different targets  $(w_i)$
  - Often,  $w_i = \frac{1}{\sigma^2}$

$$WSSE(\theta) = \sum_{i=1}^{T} w_i (y_i - M_i(\theta))^2$$

### Likelihood as GoF

- How likely is the observed data to have come from a model M with set of parameter values  $\theta$ ?
- Assuming targets are independent, overall likelihood is the product of individual likelihoods

$$L(y|M(\theta)) = \prod_{i=1}^{T} L_i(y_i|M(\theta))$$

Generally, we work with log-likelihood

$$\mathcal{L}(y|M(\theta)) = \sum_{i=1}^{T} \log L_i(y_i|M_i(\theta))$$

# Commonly used likelihoods

#### Normal distribution

$$\log L(y_i, \sigma_i | M_i(\theta)) = -\frac{1}{2} \log \left(2\pi\sigma_i^2\right) - \frac{1}{\sigma_i^2} \left(y_i - M_i(\theta)\right)^2$$

• In R: dnorm(x= $y_i$ , mean= $M_i(\theta)$ , sd= $\sigma_i$ , log=T)

#### Binomial distribution

$$\log L(y_i, n_i | M_i(\theta)) = \log \left( \binom{n_i}{y_i} \right) + y_i \log \left( M_i(\theta) \right) + (n_i - y_i) \log \left( 1 - M_i(\theta) \right)$$

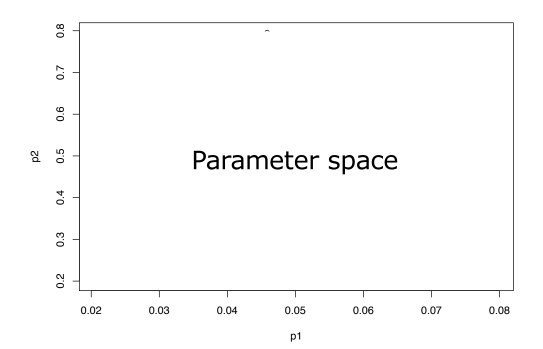
• In R: dbinom(x= $y_i$ , size= $n_i$ , prob= $M_i(\theta)$ , log=T)

#### Multinomial distribution

• In R: dmultinom( $x=y_i$ , prob= $M_i(\theta)$ , log=T)

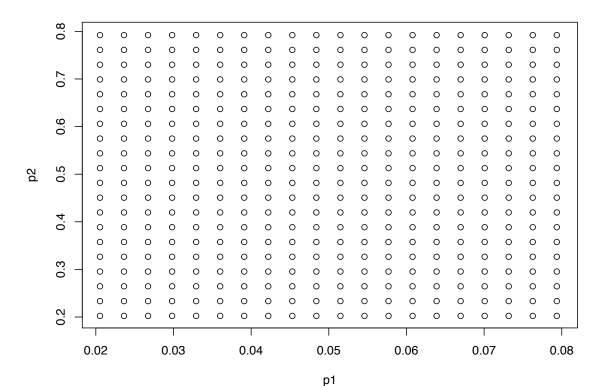
# Search Strategy

- Define plausible ranges for parameter whose values are unknown
- Use a search strategy to "search" through the input parameter space
  - Run the model for sets of parameter values generated by search strategy



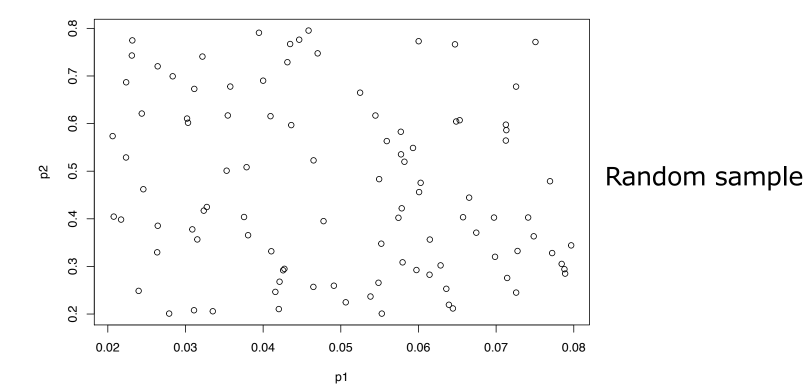
### Grid Search

- Run model for all possible combinations of parameter values
- Often infeasible



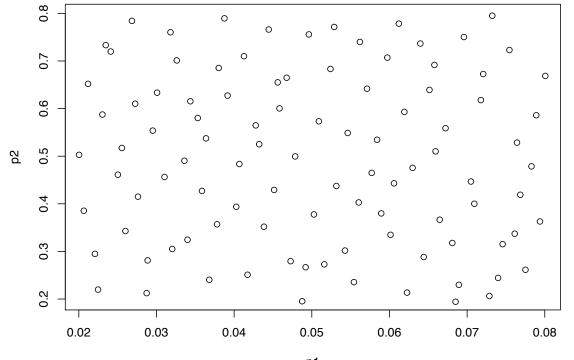
### Random Search

- Randomly sample a large number of parameter value sets from probabilistic distributions
- Use "Latin hypercube sampling" (LHS) to ensure sample captures the full parameter space



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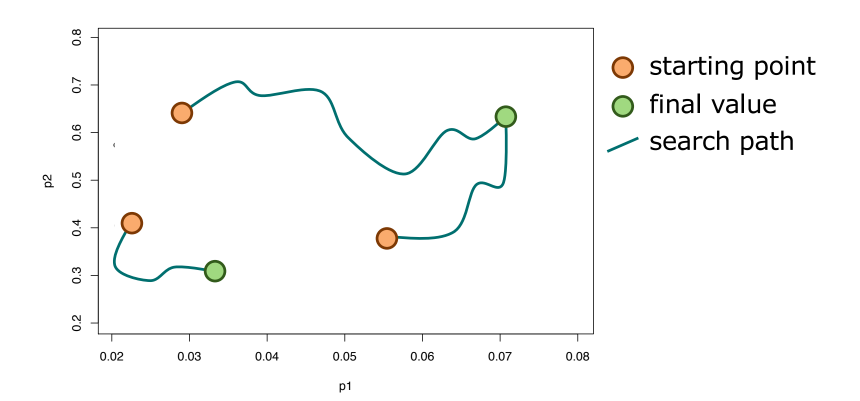
LHS sample

#### **Iterative Search**

- Use fits of past input values to determine which input values to try next
- Directed methods
  - Nelder-Mead (simplex method)
  - Gradient-descent and others
- Meta-heuristic algorithms
  - Genetic algorithms
  - Simulated annealing

# Nelder-Mead Algorithm

- Downhill simplex method
- Must be run multiple times for different starting points to avoid local extrema



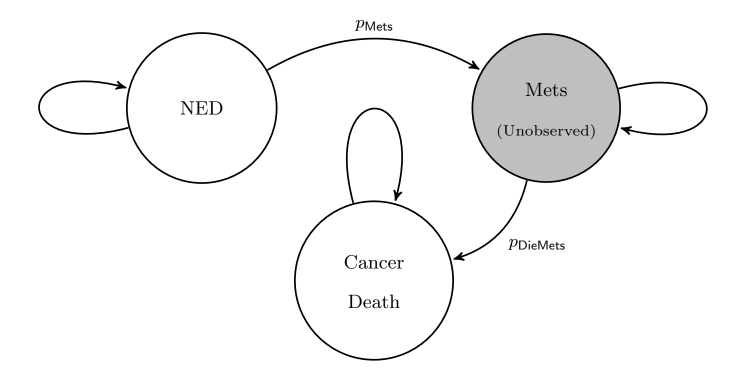
Example: Calibrating a 3-state cancer model

#### 3-state cancer model

- Relative survival as reported by the Surveillance, Epidemiology, and End Results (SEER) Program, represents cancer survival in the absence of other causes of death
- Cancer Markov models often have a distant metastasis state, a state not directly observed in SEER, from which cancer deaths are presumed to occur
- These models are used to model interventions that affect transitions to and from this state

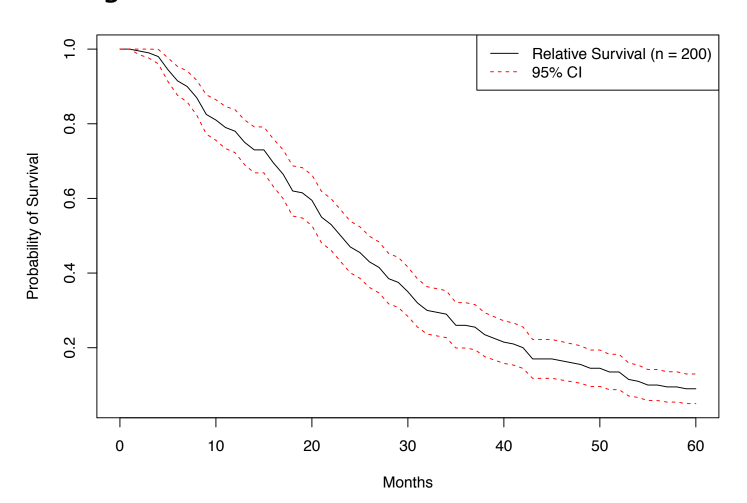
### 3-state cancer model

- Two unknown transition probabilities
  - p.Mets: Monthly risk of developing distant recurrence, range = [0.04, 0.16]
  - p.DieMets: Monthly risk of cancer death, range = [0.04, 0.16]



# Target: Relative survival

 Data frame "CRS\_targets\$Surv" stored in data file "CRS\_CalibTargets.RData"



## Calibration R code template

```
### Load calibration targets ###
### Load model as a function ###
### Specify calibration parameters ###
### Calibrate ###
### Explore best-fitting input sets ###
```

### R Session

# **Bayesian Calibration**

# Bayesian setup

- Instead of a single best-fitting value, we would like an estimate of uncertainty in model parameters,  $\theta$ , given our observed targets, y
  - E.g. a posterior distribution  $p(\theta|y)$
- Recall Bayes theorem

$$p(\theta|y) = \frac{p(y|\theta)p(\theta)}{p(y)}$$

- $p(\theta)$  is the prior distribution
- $p(y|\theta)$  has a "related" likelihood function  $L(y|\theta) \propto p(y|\theta)$
- $p(y) = \int p(y|\theta)p(\theta) d\theta$  is not a function of  $\theta$  and often difficult to calculate

# Bayesian setup

A few steps of math...

$$p(\theta|y) = \frac{p(y|\theta)p(\theta)}{p(y)}$$

$$p(\theta|y) \propto p(y|\theta)p(\theta)$$

$$p(\theta|y) \propto L(y|\theta)p(\theta)$$

 Using this fact, we can sample the posterior distribution using the likelihood function, prior distribution, and some clever algorithms

# Commonly used prior distributions for sampling $n_s$ values

#### Normal distribution

In R: rnorm (n= 
$$n_s$$
, mean= $\bar{\theta}$ , sd= $\sigma_i$ )

#### Uniform distribution

```
In R: runif (n= n_s, min= lb, max=ub)
```

#### Beta distribution

```
In R: rbeta (n= n_s, shape1=\alpha, shape2=\beta)
```

#### Gamma distribution

```
In R: rgamma (n= n_s, shape=\alpha, rate=\beta)
```

# Pros and Cons of Bayesian Calibration

- Pros: We obtain distributions of parameters rather than only point estimates
- Cons: "Harder" to implement than other methods? Mmm not really!

# Obtaining a Posterior Distribution

- Analytically
  - Not feasible for simulation models
- Markov chain Monte Carlo (MCMC)
- Sampling methods
  - Sampling Importance Resampling (SIR)
  - Incremental Mixture Importance Sampling (IMIS)

#### Markov chain Monte Carlo (MCMC)

- Simulates posterior distribution by jumping in the parameter space in a Markovian fashion
- The transition matrix of the Markov chain is proposed using a proposal distribution
- There are numerous MCMC algorithms, including Metropolis-Hastings (MH)
- R has some packages that implement MCMC algorithms
- Other R packages interface with external MCMC software, including WinBUGS, JAGS, and Stan

#### Markov chain Monte Carlo (MCMC)

#### Pros:

Theoretically will eventually converge to the "true" posterior distribution

#### Cons:

- Computationally intensive
- Autocorrelation is a problem with simulation models and therefore requires a high number of samples
- Use of external software (WinBUGS, JAGS), requires implementing the model within that software's syntax

### Sampling Importance Resampling (SIR)

Used to simulate posterior distributions (Rubin 1988) with three basic steps

- **1. Sampling:** Sample a large number, N, of parameter sets from prior distributions
- 2. Importance:
  - 1. For each parameter set  $\theta_i$  (i = 1, ..., N), run the simulation model and compute the likelihood
  - 2. Compute the (normalized) sampling importance weights  $w_i$  for each parameter set  $\theta_i$  by dividing its likelihood value by the sum of likelihood values of all parameter sets,

$$w_i = \frac{L_i}{\sum_{i=1}^N L_i}$$

**3. Resampling:** Sample from the discrete distribution of  $\{\theta(1), ... \theta(N)\}$  with probabilities  $w_i$ 

### Incremental Mixture Importance Sampling (IMIS)

- SIR can miss areas of high posterior probability as the number of parameter increases
- IMIS addresses limitations of SIR and was proposed by Steele at al. (2006)
- Starts with a modest-size SIR
- But in addition to SIR samples, add in samples from a multivariate normal distribution centered at the point with the highest importance weight
- Recalculate importance weights and sample a new sample of input parameter sets
- At the end, posterior becomes a mixture of multivariate normal distributions and of the prior distribution

### Incremental Mixture Importance Sampling (IMIS)

#### 1. Initialization.

- a) Sample N parameter sets from prior distribution
- b) For each parameter set,  $\theta_i$ , calculate the likelihood  $L_i$  and compute the importance weights  $w_i$
- **2. Importance sampling**. Iterate for k = 1, 2, ...
  - a) Choose parameter set with maximum importance weight as the mean of a multivariate normal (MVN) distribution, and compute covariance matrix from nearby parameter sets. Define  $H_k = \text{MVN}(\theta^k, \Sigma^k)$ .
  - b) Sample B new input sets from  $H_k$
  - c) Re-calculate importance weights,  $w_i^k$ , for all N + Bk samples
  - d) Repeat Step 2 until stopping criteria is met.
- **3. Resampling**. Resample  $B_{re}$  input sets with replacement using importance weights from the last iteration, K. This is a sample from the posterior distribution.

# Incremental Mixture Importance Sampling (IMIS) in R

- Function IMIS() from IMIS library takes the following inputs
  - B: sample size at each iteration of IMIS
  - B.re: number of draws from the posterior
  - number\_k: maximum number of iterations in IMIS
- IMIS() also requires the following functions to be defined by the user
  - prior (x): returns the prior density of x
  - likelihood(x): returns the likelihood of x
  - sample.prior(n): returns n samples from prior distribution of x

# Incremental Mixture Importance Sampling (IMIS) Outputs

- resamples: B.re draws from the posterior distribution
- stat: diagnostic statistics at each IMIS iteration
  - MargLike:
  - UniquePoint: expected number of unique points among resamples
  - MaxWeight: maximum importance weight
  - ESS: effective sample size (the closer to B.re, the better)
- center: center of Gaussian components

### R Session

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