

Session 3

Multi-population Hui-Walter models

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Recap

- Fitting models using MCMC is easy with JAGS / runjags
- But we must **never forget** to check convergence and effective sample size!
- More complex models become easy to implement
 - For example imperfect diagnostic tests, and Hui-Walter models
 - But remember to be realistic about what is possible with your data
 - Also carefully consider the influence of your priors

Multi-population Hui-Walter models

Hui-Walter models with multiple populations

- Basically an extension of the single-population model
- Works best with multiple populations each with differing prevalences
 - Including an unexposed population works well
 - BUT be wary of assumptions regarding constant sensitivity/specificity across populations with very different types of infections

Independent intercepts for populations

```
model{
  for(p in 1:Populations){
    Tally[1:4, p] ~ dmulti(prob[1:4, p], TotalTests[p])
    # Test1- Test2- Pop1
    prob[1, p] <- (prev[p] * ((1-se[1])*(1-se[2]))) + ((1-prev[p]) *
↪ ((sp[1])*(sp[2])))
    ## etc ##
    prev[p] ~ dbeta(1, 1)
  }

  se[1] ~ dbeta(HPSe[1,1], HPSe[1,2])T(1-sp[1], )
  sp[1] ~ dbeta(HPSp[1,1], HPSp[1,2])
  se[2] ~ dbeta(HPSe[2,1], HPSe[2,2])T(1-sp[2], )
  sp[2] ~ dbeta(HPSp[2,1], HPSp[2,2])

  #data# Tally, TotalTests, Populations, HPSe, HPSp
  #monitor# prev, prob, se, sp
  #inits# prev, se, sp
}
```

Random intercepts for a larger number of populations

```
model{
  for(p in 1:Populations){
    Tally[1:4, p] ~ dmulti(prob[1:4, p], TotalTests[p])
    # Test1- Test2- Pop1
    prob[1, p] <- (prev[p] * ((1-se[1])*(1-se[2]))) + ((1-prev[p]) *
↪ ((sp[1])*(sp[2])))
    ## etc ##

    logit(prev[p]) <- intercept + population_effect[p]
    population_effect[p] ~ dnorm(0, tau)
  }

  tau ~ dgamma(0.01, 0.01)
  intercept ~ dnorm(0, 0.33)

  se[1] ~ dbeta(HPSse[1,1], HPSse[1,2])T(1-sp[1], )
  sp[1] ~ dbeta(HPSp[1,1], HPSp[1,2])
  se[2] ~ dbeta(HPSse[2,1], HPSse[2,2])T(1-sp[2], )
  sp[2] ~ dbeta(HPSp[2,1], HPSp[2,2])

  #data# Tally, TotalTests, Populations, HPSse, HPSp
  #monitor# prev, prob, se, sp
  #inits# prev, se, sp
}
```

Multiple populations: assumptions

- We typically assume that the sensitivity and specificity *must* be consistent between populations
 - Do you have an endemic and epidemic population?
 - Or vaccinated and unvaccinated?

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- We typically assume that the sensitivity and specificity *must* be consistent between populations
 - Do you have an endemic and epidemic population?
 - Or vaccinated and unvaccinated?
- The populations can be artificial (e.g. age groups) but must not be decided based on the diagnostic test results
 - It helps if the prevalence differs between the populations

Multiple populations: special cases

- A small disease-free group is extremely helpful
 - Contains strong data regarding specificity
 - As long as specificity can be assumed to be the same in the other populations

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- A small disease-free group is extremely helpful
 - Contains strong data regarding specificity
 - As long as specificity can be assumed to be the same in the other populations
- A small experimentally infected group MAY be helpful but it is often dangerous to assume that sensitivity is consistent!

Incorporating populations with known prevalence

Up to now prevalence has been a parameter, but it can also be (partially) observed:

TODO

Other runjags options

There are a large number of other options to runjags. Some highlights:

- The method can be parallel or background or bgparallel
- You can use `extend.jags` to continue running an existing model (e.g. to increase the sample size)
- You can use `coda::as.mcmc.list` to extract the underlying MCMC chains
- Use the `summary()` method to extract summary statistics
 - See `?summary.runjags` and `?runjagsclass` for more information

Using embedded character strings

For simple models we might not want to bother with an external text file. Then we can do:

```
mt <- "  
model{  
  Positives ~ dbinom(prevalence, TotalTests)  
  prevalence ~ dbeta(2, 2)  
  
  #data# Positives, TotalTests  
  #monitor# prevalence  
  #inits# prevalence  
}  
"  
  
Positives <- 7  
TotalTests <- 10  
prevalence <- list(chain1=0.01, chain2=0.99)  
results <- run.jags(mt, n.chains=2)
```

- But I would advise that you stick to using a separate text file!

Setting the RNG seed

If we want to get numerically replicable results we need to add `.RNG.name` and `.RNG.seed` to the initial values, and an additional `#modules#` lecuyer hook to our `basicjags.bug` file:

```
model{
  Positives ~ dbinom(prevalence, TotalTests)
  prevalence ~ dbeta(2, 2)

  #data# Positives, TotalTests
  #monitor# prevalence
  #inits# prevalence, .RNG.name, .RNG.seed
  #modules# lecuyer
}

.RNG.name <- "lecuyer::RngStream"
.RNG.seed <- list(chain1=1, chain2=2)
results <- run.jags('basicjags.bug', n.chains=2)
```

- Every time this model is run the results will now be identical

Practical session 3

Points to consider

1. What are the benefits of including multiple populations?
2. How can we define/obtain these populations?
3. What happens if our fundamental assumptions about consistent Se/Sp are broken?

Summary

- Multiple populations helps to estimate Se and Sp
 - Particularly if the prevalences differ
 - A large number of populations with small N may be better as a random effect
- Populations may be artificial
 - But cannot be based on the result of either test
- But if Se / Sp are inconsistent then we will get misleading results
 - In practice, groups with widely varying prevalence rarely have consistent Se / Sp
 - It is possible to allow Se / Sp to differ between populations, but then there is no benefit of combining the data