QTL with GxE or Covariates

- adjust phenotype by covariate
  - covariate(s) = environment(s) or other trait(s)
- additive covariate
  - covariate adjustment same across genotypes
  - “usual” analysis of covariance (ANCOVA)
- interacting covariate
  - address GxE
  - capture genotype-specific relationship among traits
- another way to think of multiple trait analysis
  - examine single phenotype adjusted for others

R/qtl & covariates

- additive and/or interacting covariates
- test for QTL after adjusting for covariates

```r
## Get Brassica data.
library(qtlbim)
data(Bnapus)
Bnapus <- calc.genoprob(Bnapus, step = 2, error = 0.01)

## Scatterplot of two phenotypes: 4wk & 8wk flower time.
plot(Bnapus$pheno$log10flower4, Bnapus$pheno$log10flower8, col = c("blue", "red"), chr = "N2",
     main = "blue = 4wk, red = 8wk")
```
R/qtl & covariates

- additive and/or interacting covariates
- test for QTL after adjusting for covariates

```r
## IM scan of 8wk adjusted for 4wk.
## Adjustment independent of genotype
fl8.4 <- scanone(Bnapus, find.pheno(Bnapus, "log10flower8"),
                 addcov = Bnapus$pheno$log10flower4)

## IM scan of 8wk adjusted for 4wk.
## Adjustment changes with genotype.
fl8.4 <- scanone(Bnapus, find.pheno(Bnapus, "log10flower8"),
                 intcov = Bnapus$pheno$log10flower4)

plot(fl8, fl8.4a, fl8.4, chr = "N2",
     main = "black = 8wk, blue = addcov, red = intcov")
```
scatterplot adjusted for covariate

```r
## Set up data frame with peak markers, traits.
markers <- c("E38M50.133","ec2e5a","wg7f3a")
tmpdata <- data.frame(pull.geno(Bnapus) [,markers])
tmpdata$fl4 <- Bnapus$pheno$log10flower4
tmpdata$fl8 <- Bnapus$pheno$log10flower8

## Scatterplots grouped by marker.
library(lattice)
xyplot(fl8 ~ fl4, tmpdata, group = wg7f3a,
       col = c("blue","red"), cex = 2, type = c("p","r"),
       xlab = "log10(4wk flower time)",
       ylab = "log10(8wk flower time)",
       main = "marker at 47cM")

xyplot(fl8 ~ fl4, tmpdata, group = E38M50.133,
       col = c("blue","red"), cex = 2, type = c("p","r"),
       xlab = "log10(4wk flower time)",
       ylab = "log10(8wk flower time)",
       main = "marker at 80cM")
```

R/qtlbim and GxE

- similar idea to R/qtl
  - fixed and random additive covariates
  - GxE with fixed covariate
- multiple trait analysis tools coming soon
  - theory & code mostly in place
  - properties under study
  - expect in R/qtlbim later this year
  - Samprit Banerjee (N Yi, advisor)