

# 4. Model fitting

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Two of the most important R commands;

- lm(): fits Linear Models
- glm(): fits Generalized Linear Models

(If you've used SAS, its glm is **not** the same as R's)

'Linear Regression' and 'Logistic Regression' are special cases.

There's a lot to learn here – entire graduate courses! – so the help files are huge. How are lm(), glm() used in genetics?

Many analyses fit the 'additive model'

 $y = \beta_0 + \beta \times \#$ minor alleles



An alternative is the 'dominant model';



 $y = \beta_0 + \beta \times (G \neq AA)$ 

or the 'recessive model';



 $y = \beta_0 + \beta \times (G = = AA)$ 

Finally, the 'two degrees of freedom model';





The lm() command fits all of these, in the same way. Formally,

```
lm(outcome \sim genetic.predictor, [...] )
```

estimates the association between outcome and predictor

The **optional** arguments [...] might be

- data = my.data your dataset
- subset = race=="CEPH" use partial data
- weights = for advanced analyses

#### Use of lm() in genetics

How to make the genetic.predictor variable? Note that when R meets FALSE or TRUE in a 'math' setting, it will **coerce** them to be zero or one. So 1 + 2\*TRUE is 3, TRUE + 2\*FALSE is 1, etc

Suppose you had genotypes stored in vector g, as character strings "AA"/"Aa"/"aa". You might use these commands;

Chosen Model	genetic.predictor <-
Additive	(g=="Aa") + 2*(g=="aa")
Dominant	(g=="Aa")   (g=="aa")
Recessive	g=="aa"
2 degrees of freedom	factor(g)

- There are many other ways to do this! Use table(g, genetic.predictor) to check what you did
- Often, genotypes may be stored as 0/1/2. This is easier to work with in R but makes it harder to decide if A/C/G/T is the risk allele

## lm(): Estimates, Intervals, p-values

lm() produces point estimates for your model;

- also available via my.lm\$coefficients Or coef(my.lm).

The coefficients in the output tell you the **additive increase** in outcome associated with a **one-unit** difference in the genetic predictor.

The coefficient for predictor is in units of cholesterol per 'a' allele

## lm(): Estimates, Intervals, p-values

You will also want **confidence intervals**;

Remember to **round these numbers** to an appropriate number of significant figures! (2 or 3 is usually enough)

We are **seldom** interested in the Intercept

## lm(): Estimates, Intervals, p-values

Two-sided **p-values** are also available;

In this data, we have **strong evidence** of an **additive effect** of the minor allele on cholesterol

summary(my.lm) gives many other details - ignore for now

Confidence intervals are just Estimate  $\pm$  2×Std.Error

## Use of glm() in genetics

Logistic regression is the 'default' analysis for binary outcomes

Outcome	Туре	Regression	Scale
Cholesterol			
Blood Pressure	Continuous	Linear	Difference in Outcome
BMI			
Death Stroke BMI>30	Binary	Logistic	Ratio of odds

What are **odds**? Really just **probability**...

Odds are a [gambling-friendly] measure of chance;



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- so what are odds ratios?

Using the data from the bar charts;

```
> genpred2 <- factor(g) # the 2df model</pre>
```

```
> glm1 <- glm( dead10yrs ~ genpred2, family=binomial)</pre>
```

```
> coef(glm1)
```

pred2Aa pred2aa 0.6931 2.8904

These are log odds ratio estimates; to transform to OR, use  $e^{0.6931} = 2, e^{2.8904} = 18$ 

They are given **relative to the baseline group** – 'AA' in this case

Don't forget the family=binomial argument!

## Use of glm() in genetics

Confidence intervals, p-values as with lm(), for the log odds ratios;

Use exp() to get odds ratio estimates, intervals; p-values are scale-independent

#### The formula syntax

We fit  $lm(y \sim genetic.predictor)$  and  $glm(y \sim genpred2)$ . To see how phenotype depends on *several* covariates, we specify e.g.

```
y \sim genotype.pred + age + sex
```

- formally, this gives *multivariate regression*; the genotype.pred coefficients reflect the genotype effects *adjusted for age and sex* 

- Separate covariates with '+'. This is *not* addition!
- For now, make predictor variables first, then do regression. It's possible to do everything in one step, but use of e.g. '+' will confuse R – unless you're careful.
- For keen people; in the formula syntax, \* indicates that interactions should be fitted, I() insulates mathematical operations, -1 removes the intercept... see ?formula
- For very keen people; vcovHC() in the sandwich package provides 'robust' standard errors; coeftest() in the lmtest package uses them to give 'robust' tests.