# Two numeric explanatory variables

INTERMEDIATE REGRESSION IN R

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#### Visualizing 3 numeric variables

- 3D scatter plot
- 2D scatter plot with response as color

#### Another column for the fish dataset

species	mass_g	length_cm	height_cm
Bream	1000	33.5	18.96
Bream	925	36.2	18.75
Roach	290	24.0	8.88
Roach	390	29.5	9.48
Perch	1100	39.0	12.80
Perch	1000	40.2	12.60
Pike	1250	52.0	10.69



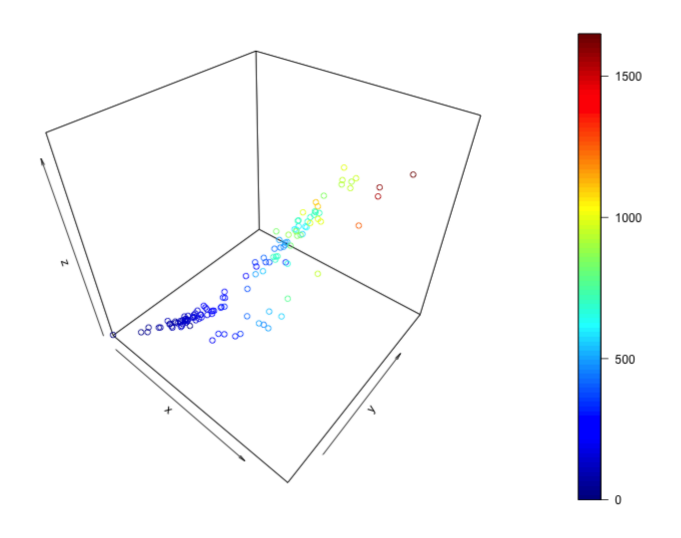
#### 3D scatter plot

```
library(plot3D)
scatter3D(fish$length_cm, fish$height_cm, fish$mass_g)
library(plot3D)
library(magrittr)
fish %$%
  scatter3D(length_cm, height_cm, mass_g)
```

#### 3D scatter plot

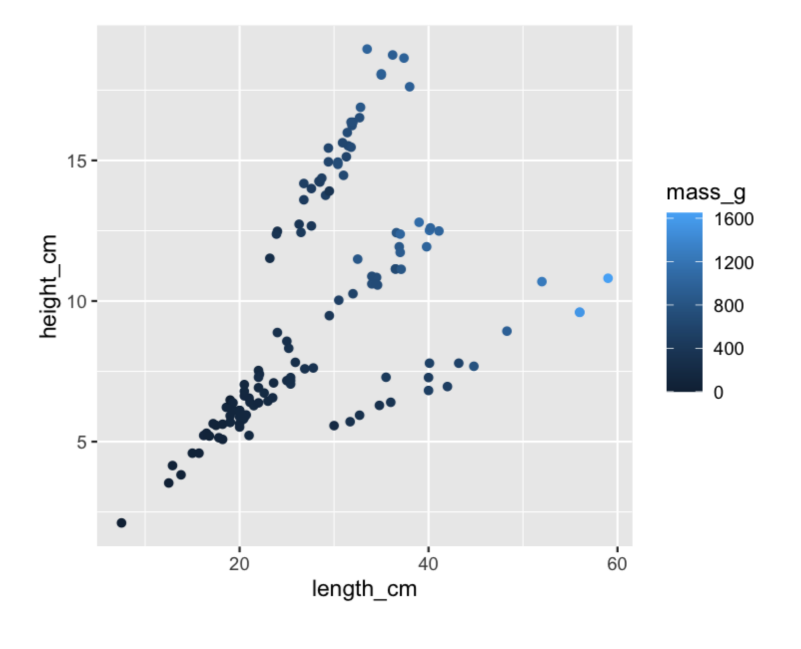
```
library(plot3D)
library(magrittr)

fish %$%
  scatter3D(length_cm, height_cm, mass_g)
```



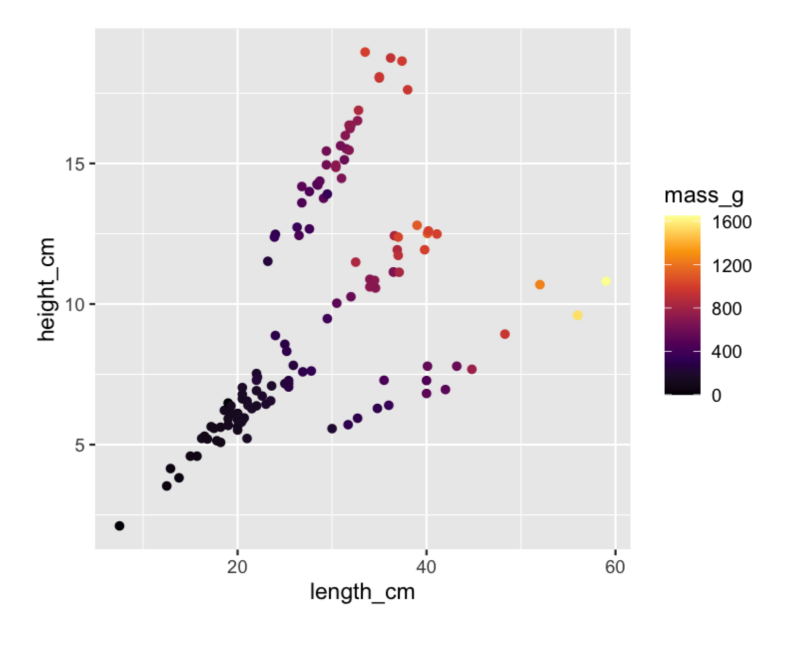
#### 2D scatter plot, color for response

```
ggplot(
  fish,
  aes(length_cm, height_cm, color = mass_g)
) +
  geom_point()
```



#### Viridis color scales

```
ggplot(
  fish,
  aes(length_cm, height_cm, color = mass_g)
) +
  geom_point() +
  scale_color_viridis_c(option = "inferno")
```



#### Modeling with 2 numeric explanatory variable

```
mdl_mass_vs_both <- lm(mass_g \sim length_cm + height_cm, data = fish)
```

```
Call:

lm(formula = mass_g ~ length_cm + height_cm, data = fish)

Coefficients:

(Intercept) length_cm height_cm

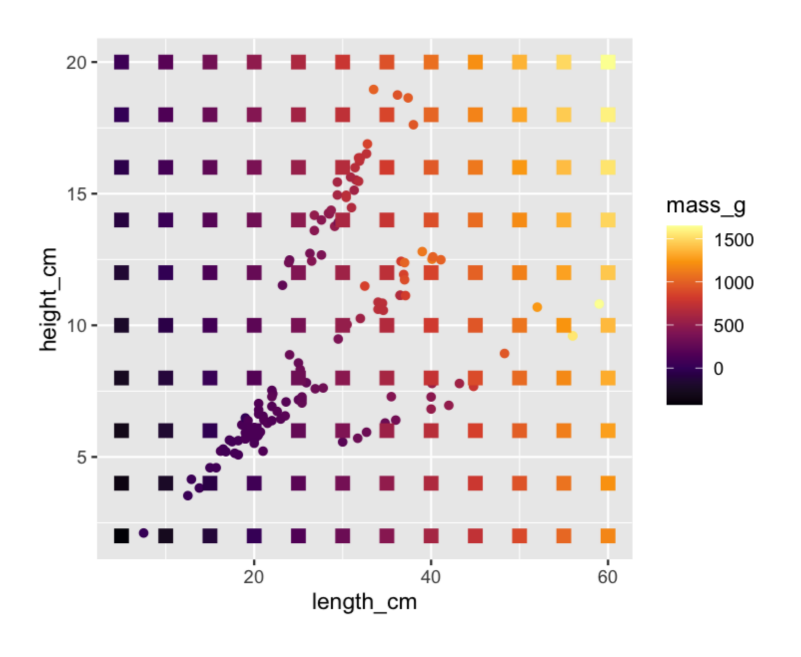
-622.16 28.97 26.34
```

#### The prediction flow

```
explanatory_data <- expand_grid(</pre>
  length_cm = seq(5, 60, 5),
  height_cm = seq(2, 20, 2)
prediction_data <- explanatory_data %>%
  mutate(
    mass_g = predict(mdl_mass_vs_both, explanatory_data)
```

#### Plotting the predictions

```
ggplot(
  fish,
  aes(length_cm, height_cm, color = mass_g)
) +
  geom_point() +
  scale_color_viridis_c(option = "inferno") +
  geom_point(
    data = prediction_data, shape = 15, size = 3
)
```



#### Including an interaction

```
mdl_mass_vs_both_inter <- lm(mass_g \sim length_cm * height_cm, data = fish)
```

```
Call:

lm(formula = mass_g ~ length_cm * height_cm, data = fish)

Coefficients:

(Intercept) length_cm height_cm length_cm:height_cm

159.1144 0.3001 -78.1234 3.5455
```

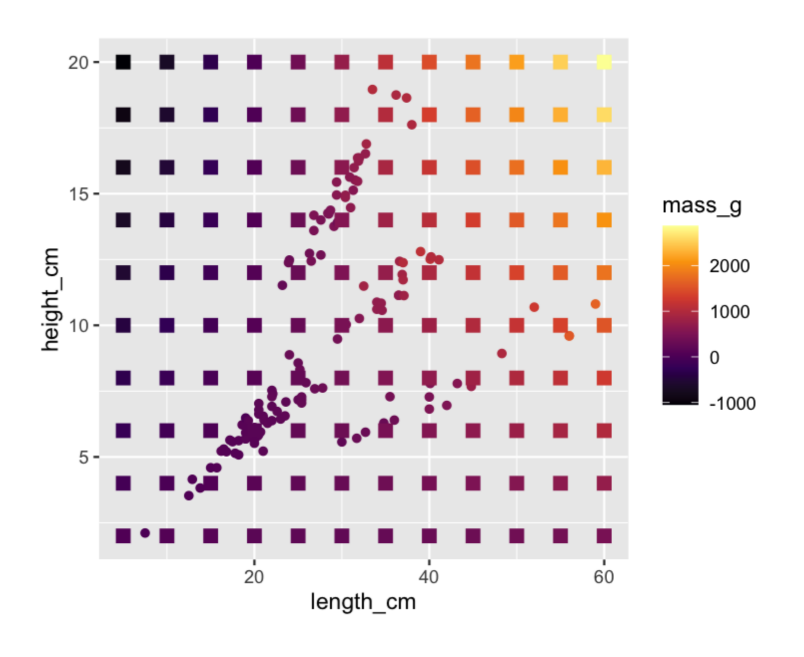


#### The prediction flow again

```
explanatory_data <- expand_grid(</pre>
  length_cm = seq(5, 60, 5),
  height_cm = seq(2, 20, 2)
prediction_data <- explanatory_data %>%
  mutate(
    mass_g = predict(mdl_mass_vs_both_inter, explanatory_data)
```

#### Plotting the predictions

```
ggplot(
  fish,
  aes(length_cm, height_cm, color = mass_g)
) +
  geom_point() +
  scale_color_viridis_c(option = "inferno") +
  geom_point(
    data = prediction_data, shape = 15, size = 3
)
```



## Let's practice!

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# More than 2 explanatory variables

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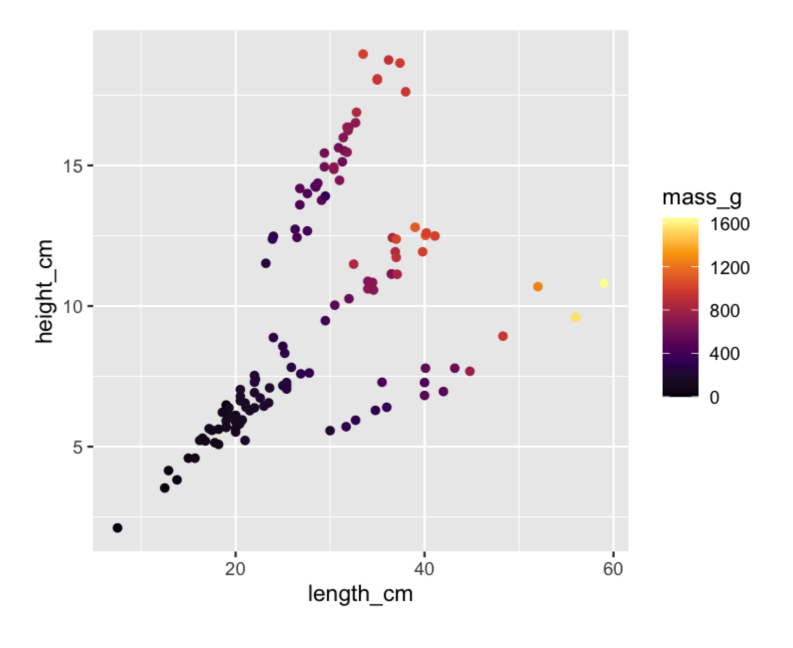
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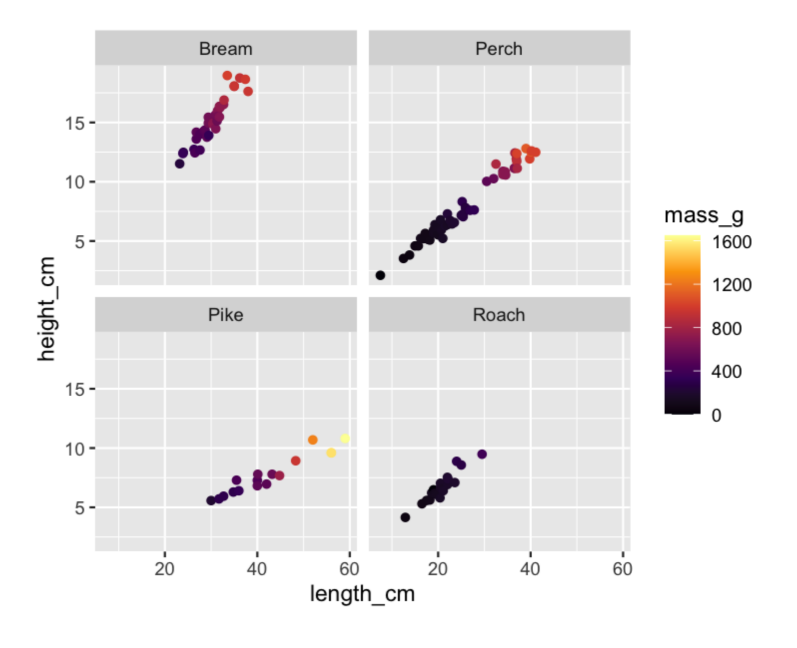
#### From last time

```
ggplot(
  fish,
  aes(length_cm, height_cm, color = mass_g)
) +
  geom_point() +
  scale_color_viridis_c(option = "inferno")
```



#### Faceting by species

```
ggplot(
  fish,
  aes(length_cm, height_cm, color = mass_g)
) +
  geom_point() +
  scale_color_viridis_c(option = "inferno") +
  facet_wrap(vars(species))
```



#### Different levels of interaction

No interactions

```
lm(mass_g ~ length_cm + height_cm + species + 0, data = fish)
```

2-way interactions between pairs of variables

```
lm(
   mass_g ~ length_cm + height_cm + species + length_cm:height_cm + length_cm:species + height_cm:species + 0,
   data = fish
)
```

3-way interaction between all three variables

```
lm(
   mass_g ~ length_cm + height_cm + species + length_cm:height_cm + length_cm:species + height_cm:species + length_cm:height_cm:species + 0,
   data = fish
)
```

#### All the interactions

```
lm(
   mass_g ~ length_cm + height_cm + species + length_cm:height_cm + length_cm:species + height_cm:species + length_cm:height_cm:species + 0,
   data = fish
)
```

```
lm(
   mass_g ~ length_cm * height_cm * species + 0,
   data = fish
)
```

#### Only 2-way interactions

```
1m(
  mass_g ~ length_cm + height_cm + species + length_cm:height_cm + length_cm:species + height_cm:spec
  data = fish
1m(
  mass_g \sim (length_cm + height_cm + species) ^2 + 0,
  data = fish
lm(
```

```
lm(
  mass_g ~ I(length_cm) ^ 2 + height_cm + species + 0,
  data = fish
)
```

<sup>&</sup>lt;sup>1</sup> To square explanatory variables, see "Introduction to Regression in R", Chapter 2, "Transforming variables"

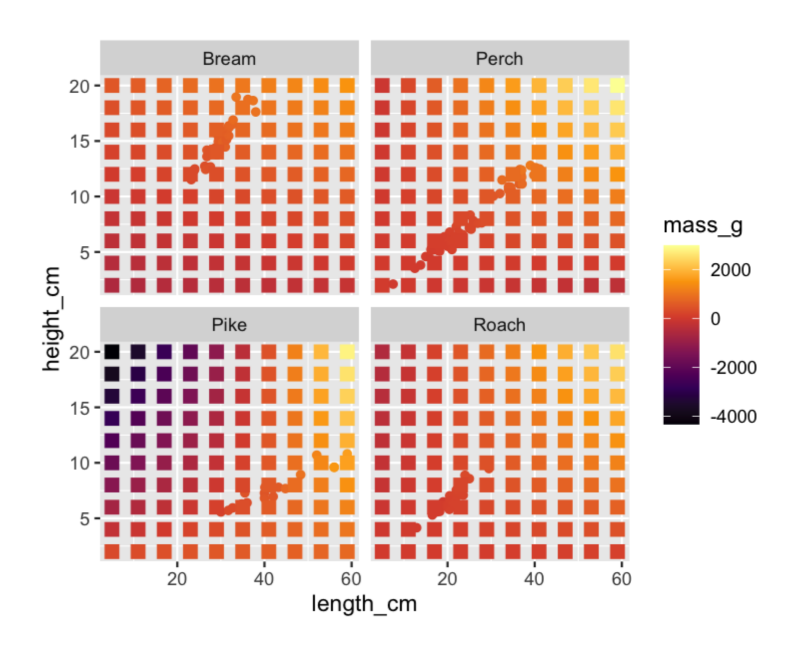


#### The prediction flow

```
mdl_mass_vs_all <-lm(mass_g \sim length_cm * height_cm * species * 0, data = fish)
explanatory_data <- expand_grid(</pre>
  length_cm = seq(5, 60, 6),
  height_cm = seq(2, 20, 2),
  species = unique(fish$species)
prediction_data <- explanatory_data %>%
  mutate(mass_g = predict(mdl_mass_vs_all, explanatory_data))
```

#### Visualizing predictions

```
ggplot(
  fish,
  aes(length_cm, height_cm, color = mass_g)
  geom_point() +
  scale_color_viridis_c(option = "inferno") +
  facet_wrap(vars(species)) +
  geom_point(
    data = prediction_data,
    size = 3, shape = 15
```



## Let's practice!

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# How linear regression works

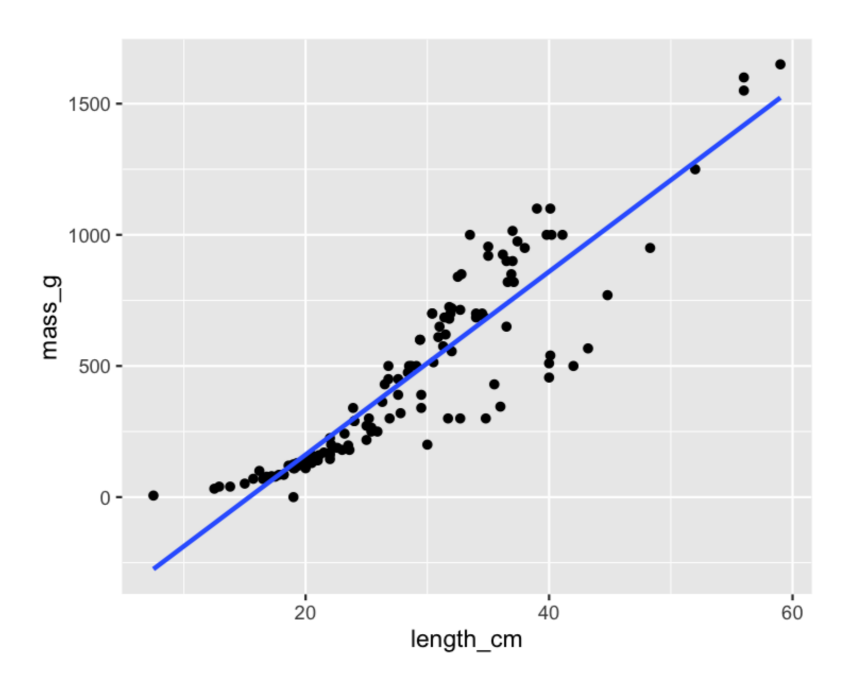
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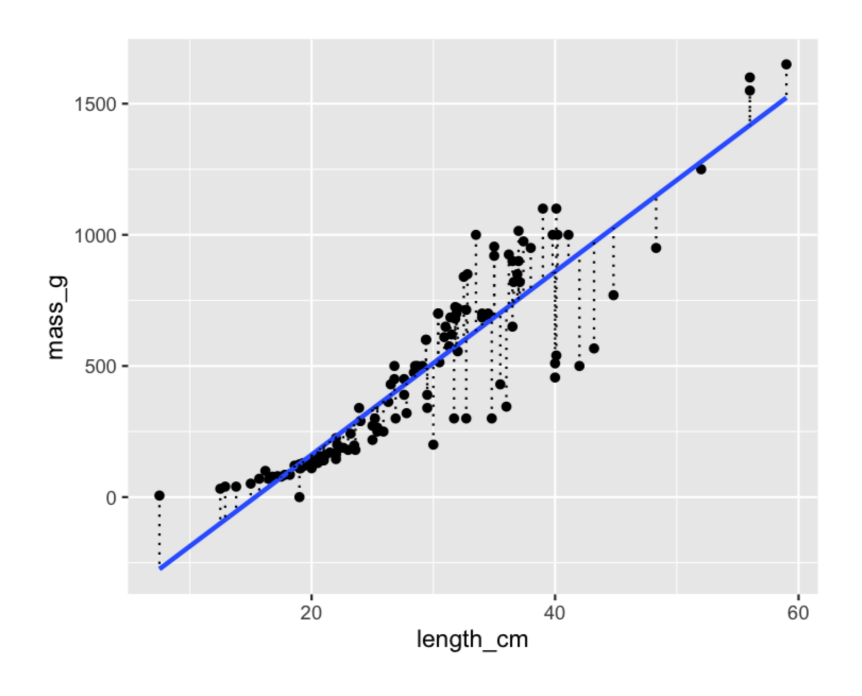


#### The standard simple linear regression plot





#### Visualizing residuals



#### A metric for the best fit

#### The simplest idea (which doesn't work)

- Take the sum of all the residuals.
- Some residuals are negative.

#### The next simplest idea (which does work)

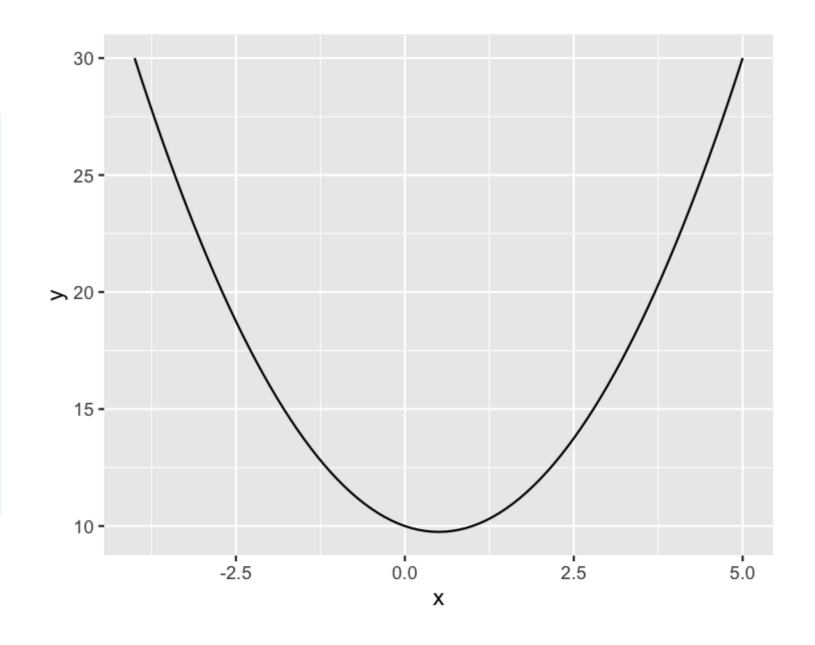
- Take the square of each residual, and add up those squares.
- This is called the sum of squares.

#### A detour into numerical optimization

A line plot of a quadratic equation

```
xy_data <- tibble(
  x = seq(-4, 5, 0.1),
  y = x ^ 2 - x + 10
)

ggplot(xy_data, aes(x, y)) +
  geom_line()</pre>
```



#### Using calculus to solve the equation

$$y = x^2 - x + 10$$

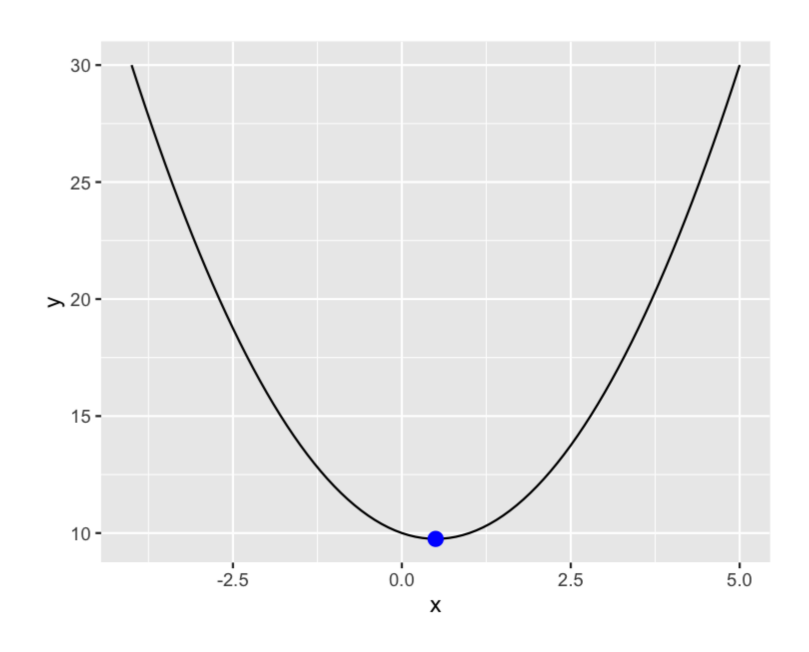
$$\frac{\partial y}{\partial x} = 2x - 1$$

$$0 = 2x - 1$$

$$x = 0.5$$

$$y = 0.5^2 - 0.5 + 10 = 9.75$$

- Not all equations can be solved like this.
- You can let R figure it out.



#### optim()

```
calc_quadratic <- function(x) {
  x ^ 2 - x + 10
}</pre>
```

```
optim(par = 3, fn = calc_quadratic)
```

```
$par
[1] 0.4998047
$value
[1] 9.75
$counts
function gradient
      30
               NA
$convergence
[1] 0
$message
NULL
```

#### Slight refinements

```
calc_quadratic <- function(coeffs) {
  x <- coeffs[1]
  x ^ 2 - x + 10
}</pre>
```

```
optim(par = c(x = 3), fn = calc_quadratic)
```

```
$par
        Χ
0.4998047
$value
[1] 9.75
$counts
function gradient
      30
               NA
$convergence
[1] 0
$message
NULL
```

#### A linear regression algorithm

- 1. Define a function to calculate the sum of squares metric.
- 2. Call optim() to find coefficients that minimize this function.

```
calc_sum_of_squares <- function(coeffs) {
  intercept <- coeffs[1]
  slope <- coeffs[2]
  # More calculation!
}</pre>
```

```
optim(
  par = ???,
  fn = ???
)
```

## Let's practice!

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