Dialogue

In this document, the dialogue with the AI is described, including all control questions. Of each question, the expected answer is calculated.

```
library(testthat)
```

Expectations

A paper does not typically hold code (like an R markdown document). Hence, in the end, all the constants are copy-pasted into a paper and then left unchecked.

To make it easy to compare the final paper with the constants we've copy-pasted, here are all the constants:

```
total n fish <- 300
n_fish_control <- 60
n_fish_transplanted <- 240
origin_values <- c("Lake", "Stream")</pre>
transplanted_values <- c("Lake", "Stream", "Control")</pre>
lowest pre mass <- 0.5
highest_pre_mass <- 3.6
n_survived <- 151</pre>
n_died <- 89
n_enclosures <- 80</pre>
n_enclosures_with_2_fish <- 3
n_enclosures_with_3_fish <- 74
n_enclosures_with_4_fish <- 3
anomous_enclosures <- c("L20", "L25", "L6", "L7", "S22", "S24")
cage_mass_mean_for_l1 <- 1.256667</pre>
cage_mass_stdev_for_l1 <- 0.4532475</pre>
cage_mass_mean_deviation_sd_for_l1 <- c(1.1546303, -0.5662836, -0.5883466)
abs_cage_mass_mean_deviation_sd_for_l1 <- abs(cage_mass_mean_deviation_sd_for_l1)
# Check consistency between these variables
expect_equal(
 total_n_fish,
  n_fish_control + n_fish_transplanted
expect_equal(
 n_fish_transplanted,
  n_survived + n_died
expect_equal(
 n_enclosures,
  n_enclosures_with_2_fish + n_enclosures_with_3_fish + n_enclosures_with_4_fish
expect_equal(
 length(anomous enclosures),
 n_enclosures_with_2_fish + n_enclosures_with_4_fish
```

)

All solutions to the dialog are written here, and tested below, during the dialogue.

D1. Ask an AI to read the dataset

D1.1: read data

• The dialogue starts by uploading the data for the paper at here.

```
t_all <- read.csv("Bolnick_traits.txt", sep = " ")</pre>
```

D1.2: show first rows

Ask the AI:

Could you show the first rows of the data?

knitr::kable(head(t_all))

sampl	e lfis hII	O origin enclo	osurteranspla	ndtate_inpre_	_mapssele	ng th st_	_mapsesst	lengeth	bd	gw	grn	grl	sex	survived
118	118	StreamS2	Stream	06/03/11.75	4.9	NA	NA	NA	NA	NA	NA	NA	NA	0
12	12	StreamL8	Lake	06/02/11.13	4.6	NA	NA	NA	NA	NA	NA	NA	NA	0
122	122	Lake S2	Stream	06/03/12.37	5.6	NA	NA	NA	NA	NA	NA	NA	NA	0
123	123	Lake S3	Stream	06/03/11.71	5.2	NA	NA	NA	NA	NA	NA	NA	NA	0
124	124	Lake S4	Stream	06/03/12.45	5.4	NA	NA	NA	NA	NA	NA	NA	NA	0
127	127	Lake S7	Stream	06/03/10.85	4.0	NA	NA	NA	NA	NA	NA	NA	NA	0

We select only the relevant data here, so that the analysis can be checked by humans more easily:

```
t <- t_all |> dplyr::select(fishID, origin, enclosure, pre_mass, transplant, survived) |> dplyr::arran,
knitr::kable(head(t))
```

	fishID	origin	enclosure	pre_mass	transplant	survived
27	181	Lake	L1	1.00	Lake	0
28	182	Lake	L1	0.99	Lake	0
149	1	Stream	L1	1.78	Lake	1
16	16	Stream	L10	1.22	Lake	0
36	193	Lake	L10	1.71	Lake	0
164	17	Stream	L10	2.06	Lake	1

D2. Describe the dataset

D2.1: count fish

Ask the AI:

The `fishID` column denotes the ID of a fish. How many fish are in this dataset?

expect_equal(total_n_fish, length(unique(t\$fishID)))

The expected answer is 300.

D2.2: get the 'origin' values

Ask the AI:

The `origin` column denotes the location where each fish comes from. What are the locations the fish come from?

```
expect_equal(unique(t$origin), origin_values)
```

The correct answers is Lake, Stream.

D2.3: get the 'transplant' values

Ask the AI:

The `transplant` column denotes the location where each fish is transplanted to.

What are the locations the fish are transplanted to?

```
expect_equal(unique(t$transplant), transplanted_values)
```

The correct answers is Lake, Stream, Control.

D2.4: remove the controls and count the fish

Ask the AI:

Remove the rows for which the `transplant` value is 'Control'. How many fish are left?

```
t_1 <- t |> dplyr::filter(transplant != "Control")
expect_equal(n_fish_transplanted, length(unique(t_1$fishID)))
```

The correct answer is 240.

D2.5: get the range of 'pre_mass'

Ask the AI:

The `pre_mass` column denotes the mass of a fish before the transplantation.

What is the range of the values in this column?

```
expect_equal(lowest_pre_mass, min(t_1$pre_mass))
expect_equal(highest_pre_mass, max(t_1$pre_mass))
```

The correct answer is 0.5 to 3.6.

D2.6: count the number of dead fish and fish that survived

Ask the AI:

The `survived` column denotes if the fish survived the experiment, where 0 means it died and 1 means that it survived. How many fish died? And how many survived?

```
expect_equal(n_survived, sum(t_1$survived == TRUE))
expect_equal(n_died, sum(t_1$survived == FALSE))
```

The correct answers are 89 died and 151 survived.

D2.7: count the number of enclosures

Ask the AI:

```
The `enclosure` column denotes the ID of an enclosure.

How many enclosures are in this dataset?

expect_equal(n_enclosures, length(unique(t_1$enclosure)))
```

The correct answer is 80 enclosures.

D2.8: count the number of fish in each enclosure

Ask the AI:

How many fish are in each enclosure?

```
n_fish_per_enclosure <- dplyr::count(t_1, enclosure)
expect_equal(n_enclosures_with_2_fish, sum(n_fish_per_enclosure$n == 2))
expect_equal(n_enclosures_with_3_fish, sum(n_fish_per_enclosure$n == 3))
expect_equal(n_enclosures_with_4_fish, sum(n_fish_per_enclosure$n == 4))
knitr::kable(head(n_fish_per_enclosure))</pre>
```

n
3
3
3
3
3
3

The correct answer is:

- 3 enclosures have 2 fish
- 74 enclosures have 3 fish
- 3 enclosures have 4 fish

As a note to self, these are the anomalies:

```
expect_equal(anomous_enclosures, n_fish_per_enclosure[n_fish_per_enclosure$n != 3, ]$enclosure)
knitr::kable(n_fish_per_enclosure[n_fish_per_enclosure$n != 3, ])
```

	enclosure	n
13	L20	4
18	L25	2
37	L6	2
38	L7	4
55	S22	4
57	S24	2

Q0. Describe the conclusion

• A conclusion drawn from the data is that the extreme body masses are likelier to survive. Would you agree that the data supports this claim?

D4. Reproduce the results in that paper

D4.1: calculate 'cage_mass_mean'

Ask the AI:

To make a better comparison, we are going to standarize body masses per enclosure.

Add a column to the data called `cage_mass_mean` which holds the average `pre_mass` within the enclosure each fish is in.

Could you show me the data for enclosure L1?

```
cage_mass_mean_per_enclosure <- t_1 |>
  dplyr::select(enclosure, pre_mass) |>
  dplyr::group_by(enclosure) |>
  dplyr::summarise(cage_mass_mean = mean(pre_mass))
t_2 <- merge(t_1, cage_mass_mean_per_enclosure)
knitr::kable(head(t_2))</pre>
```

enclosure	fishID	origin	pre_mass	transplant	survived	cage_mass_mean
L1	181	Lake	1.00	Lake	0	1.256667
L1	182	Lake	0.99	Lake	0	1.256667
L1	1	Stream	1.78	Lake	1	1.256667
L10	16	Stream	1.22	Lake	0	1.663333
L10	193	Lake	1.71	Lake	0	1.663333
L10	17	Stream	2.06	Lake	1	1.663333

```
knitr::kable(t_2[t_2$enclosure == "L1", ])
```

enclosure	fishID	origin	pre_mass	transplant	survived	$cage_mass_mean$
L1	181	Lake	1.00	Lake	0	1.256667
L1	182	Lake	0.99	Lake	0	1.256667
L1	1	Stream	1.78	Lake	1	1.256667

```
expect_equal(
  t_2[t_2$enclosure == "L1", ]$cage_mass_mean,
  rep(cage_mass_mean_for_l1, 3),
  tolerance = 1.0e-6
)
```

For enclose L1, the expected cage_mass_mean for each of the fish is 1.256667.

D4.2: calculate 'cage_mass_stdev'

Ask the AI:

To make a better comparison, we are going to standarize body masses per enclosure. Add a column to the data called `cage_mass_stdev` which is the standard deviation of the `pre_mass` distribution of each enclosure each fish is in.

Could you show me the data for enclosure L1?

```
cage_mass_stdev_per_enclosure <- t_2 |>
  dplyr::select(enclosure, pre_mass) |>
  dplyr::group_by(enclosure) |>
  dplyr::summarise(cage_mass_stdev = sd(pre_mass))
t_3 <- merge(t_2, cage_mass_stdev_per_enclosure)
knitr::kable(head(t_3))</pre>
```

enclosure	fishID	origin	pre_mass	transplant	survived	cage_mass_mean ca	ge_mass_stdev
$\overline{\text{L1}}$	181	Lake	1.00	Lake	0	1.256667	0.4532475
L1	182	Lake	0.99	Lake	0	1.256667	0.4532475
L1	1	Stream	1.78	Lake	1	1.256667	0.4532475
L10	16	Stream	1.22	Lake	0	1.663333	0.4219400
L10	193	Lake	1.71	Lake	0	1.663333	0.4219400
L10	17	Stream	2.06	Lake	1	1.663333	0.4219400

```
knitr::kable(t_3[t_3$enclosure == "L1", ])
```

enclosure	fishID	origin	pre_mass	transplant	survived	cage_mass_mean ca	age_mass_stdev
L1	181	Lake	1.00	Lake	0	1.256667	0.4532475
L1	182	Lake	0.99	Lake	0	1.256667	0.4532475
L1	1	Stream	1.78	Lake	1	1.256667	0.4532475

```
expect_equal(
  t_3[t_3$enclosure == "L1", ]$cage_mass_stdev,
  rep(cage_mass_stdev_for_l1, 3),
  tolerance = 1.0e-6
)
```

D4.3: calculate 'cage_mass_mean_deviation_sd'

Ask the AI:

```
To make a better comparison, we are going to standarize body masses per enclosure. Add a column to the data called `cage_mass_mean_deviation_sd`. Its values are calculated per fish. Each fish its `cage_mass_mean_deviation_sd` equals the absolute difference between its `pre_mass` and its enclosure's `cage_mass_mean`, divided by the `cage_mass_stdev` of its enclosure. Could you show me the data for enclosure L1?
```

```
cage_mass_mean_deviation_sd_per_fish <- t_3 |>
   dplyr::select(fishID, pre_mass, cage_mass_mean, cage_mass_stdev) |>
   dplyr::group_by(fishID) |>
   dplyr::mutate(
        cage_mass_mean_deviation_sd =
            abs(pre_mass - cage_mass_mean) / cage_mass_stdev
   ) |>
   dplyr::select(fishID, cage_mass_mean_deviation_sd)
   expect_equal(
        nrow(cage_mass_mean_deviation_sd_per_fish),
```

```
n_fish_transplanted
)

expect_equal(
    nrow(t_3),
    n_fish_transplanted
)

t_4 <- merge(t_3, cage_mass_mean_deviation_sd_per_fish)

expect_equal(
    nrow(t_4),
    n_fish_transplanted
)

knitr::kable(head(t_4))</pre>
```

fishID	enclosure	origin	pre_mas	sstransplant	survived	cage_mass_	_meage_mass_stdeage	_mass_mean_deviation_sd
1	L1	Stream	1.78	Lake	1	1.256667	0.4532475	1.1546303
2	L2	Stream	0.65	Lake	1	1.016667	0.5589574	0.6559832
3	L2	Stream	1.66	Lake	1	1.016667	0.5589574	1.1509524
4	L3	Stream	2.26	Lake	1	1.453333	0.8309834	0.9707375
5	L3	Stream	1.50	Lake	0	1.453333	0.8309834	0.0561584
6	L4	Stream	1.89	Lake	1	1.723333	0.9016282	0.1848508

```
knitr::kable(t_4[t_4$enclosure == "L1", ])
```

	fishID	enclosure	origin	pre_mas	stranspla	ant survived	cage_mass_	_m cage _mass_	stdæge_mass_mean_deviation_sd
1	1	L1	Stream	1.78	Lake	1	1.256667	0.4532475	1.1546303
181	181	L1	Lake	1.00	Lake	0	1.256667	0.4532475	0.5662836
182	182	L1	Lake	0.99	Lake	0	1.256667	0.4532475	0.5883466

```
expect_equal(
  t_4[t_4$enclosure == "L1", ]$cage_mass_mean_deviation_sd,
  abs_cage_mass_mean_deviation_sd_for_l1,
  tolerance = 1.0e-6
)
```

The values of cage_mass_mean_deviation_sd for enclosure L1 are expected to be 1.1546303, 0.5662836, 0.5883466.

D4.4: reproduce the plot

Ask the AI:

Create a scatter plot of this data:

- Each point is a fish.
- Use a blue color for fish

that have 'Lake' as their origin.

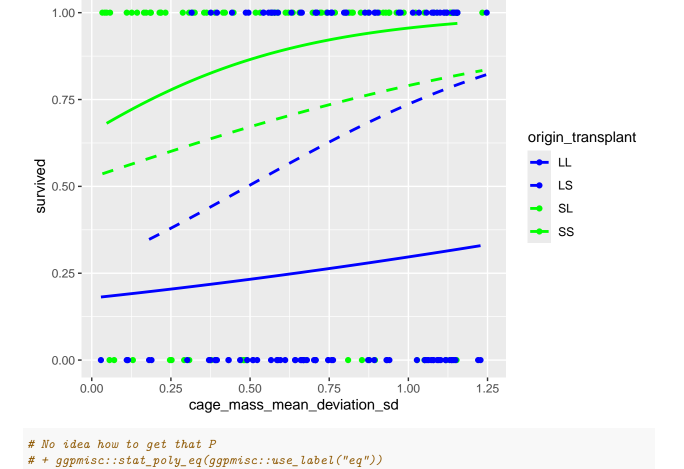
- Use a green color for fish

- that have 'Stream' as their origin.
- On the X axis, put the values of `cage_mass_mean_deviation_sd` of each fish
- On the Y axis, put the `survived` of each fish.

```
Put the fish in four categories:
Fish that have an `origin` of 'Lake' and
a `transplant` of 'Lake' are in category 'LL'
Fish that have an `origin` of 'Lake' and
a `transplant` of 'Stream' are in category 'LS'
Fish that have an `origin` of 'Stream' and
a `transplant` of 'Lake' are in category 'SL'
Fish that have an `origin` of 'Stream' and
a `transplant` of 'Stream' are in category 'SS'
Show a trendline for each binomal fit on each of these 4 categories.
Use blue lines for categories that originate from a lake
Use green lines for categories that originate from a stream
Use solid lines for categories 'LL' and 'SS'
Use dashed lines for categories 'LS' and 'SL'
```

This should reproduce the plot in the paper:

```
t_4$origin_transplant <- paste0(</pre>
  stringr::str_sub(t_4$origin,1,1),
  stringr::str_sub(t_4$transplant,1,1)
ggplot2::ggplot(
 data = t_4,
  ggplot2::aes(
   x = cage_mass_mean_deviation_sd,
   y = survived,
   color = origin_transplant,
   lty = origin_transplant
) + ggplot2::scale_color_manual(
   values = c("blue", "blue", "green", "green")
  ggplot2::scale_linetype_manual(
   values = c("solid", "dashed", "dashed", "solid")
  ) +
  ggplot2::geom_point() +
  ggplot2::geom_smooth(
   method = "glm",
   method.args = list(family = "binomial"),
   se = FALSE
\# `geom_smooth()` using formula = 'y ~ x'
```



Q1: Ask if the conclusion is correct

• A conclusion drawn from the data is that the extreme body masses are likelier to survive. Would you agree that the data supports this claim?

D5 Reproduce the results that show a flaw in the reasoning

D5.1 Add relative standarized body mass

Ask the AI:

```
To make a better comparison, we are going to standarize body masses per enclosure. Add a column to the data called `cage_mass_mean_deviation_sd_rel`.

Its values are calculated per fish.

Each fish its `cage_mass_mean_deviation_sd_rel` equals the difference between its `pre_mass` and its enclosure's `cage_mass_mean`, divided by the `cage_mass_stdev` of its enclosure.

Could you show me the data for enclosure L1?

cage_mass_mean_deviation_sd_per_fish_rel <- t_4 |> dplyr::select(fishID, pre_mass, cage_mass_mean, cage_mass_stdev) |> dplyr::group_by(fishID) |>
```

```
dplyr::mutate(
    cage_mass_mean_deviation_sd_rel =
        (pre_mass - cage_mass_mean) / cage_mass_stdev
) |>
    dplyr::select(fishID, cage_mass_mean_deviation_sd_rel)
expect_equal(
    nrow(cage_mass_mean_deviation_sd_per_fish_rel),
    n_fish_transplanted
)

expect_equal(
    nrow(t_4),
    n_fish_transplanted
)

t_5 <- merge(t_4, cage_mass_mean_deviation_sd_per_fish_rel)
expect_equal(
    nrow(t_5),
    n_fish_transplanted
)

knitr::kable(head(t_5))</pre>
```

fishII	enclo	surerigin pre_n	natssanspl	la nt rvivo	edage_mas	s <u>cagoea</u> mass	s <u>cægtele</u> vnass_r	mea n ri gin	<u>viattionsepakahnt</u> mass_mean_d	$ m leviation_s$
1	L1	Stream1.78	Lake	1	1.256667	0.4532475	1.154630	3 SL	1.1546303	
2	L2	Stream 0.65	Lake	1	1.016667	0.5589574	0.655983	2 SL	-0.6559832	
3	L2	Stream1.66	Lake	1	1.016667	0.5589574	1.150952	4 SL	1.1509524	
4	L3	Stream2.26	Lake	1	1.453333	0.8309834	0.970737	5 SL	0.9707375	
5	L3	Stream1.50	Lake	0	1.453333	0.8309834	0.056158	4 SL	0.0561584	
6	L4	Stream1.89	Lake	1	1.723333	0.9016282	0.184850	8 SL	0.1848508	

```
knitr::kable(t_5[t_5$enclosure == "L1", ])
```

	fishII	Dencl	osu m rigin pre_n	n asa nspl	a sıt rviv	edage_mas <u>eageean</u> ass	a gt demass_m	eaor <u>ig</u> ibe <u>v</u>	vitatinspysyltmass_mean_deviation_se
1	1	L1	Stream.78	Lake	1	1.256667 0.4532475	1.1546303	SL	1.1546303
181	181	L1	Lake 1.00	Lake	0	$1.256667 \ 0.4532475$	0.5662836	LL	-0.5662836
182	182	L1	Lake 0.99	Lake	0	$1.256667 \ 0.4532475$	0.5883466	LL	-0.5883466

```
expect_equal(
  t_5[t_5$enclosure == "L1", ]$cage_mass_mean_deviation_sd_rel,
  cage_mass_mean_deviation_sd_for_l1,
  tolerance = 1.0e-6
)
```

D5.2 Plot relative standarized body mass with same fit

Ask the AI:

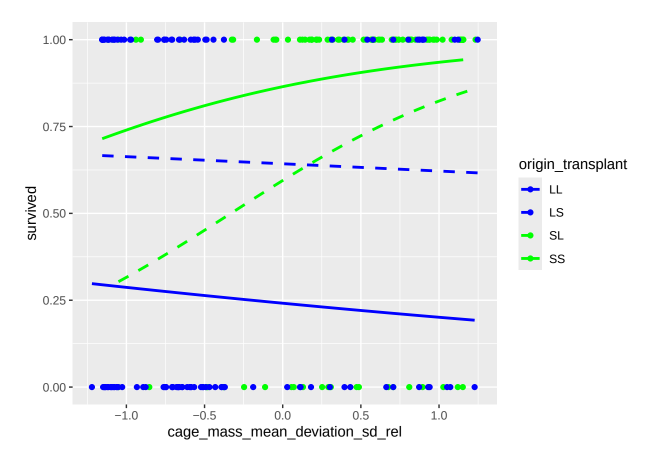
```
Create a scatter plot of this data:
- Each point is a fish.
```

⁻ Use a blue color for fish that have 'Lake' as their origin.

```
- Use a green color for fish
  that have 'Stream' as their origin.
- On the X axis, put the values of `cage_mass_mean_deviation_sd_rel`
  of each fish
- On the Y axis, put the `survived` of each fish.
- Put the fish in four categories:
  - Fish that have an `origin` of 'Lake' and
     a `transplant` of 'Lake' are in category 'LL'
  - Fish that have an `origin` of 'Lake' and
     a `transplant` of 'Stream' are in category 'LS'
  - Fish that have an `origin` of 'Stream' and
     a `transplant` of 'Lake' are in category 'SL'
  - Fish that have an `origin` of 'Stream' and
     a 'transplant' of 'Stream' are in category 'SS'
- Show a trendline for each binomal fit on each of these 4 categories.
  - Use blue lines for categories that originate from a lake
  - Use green lines for categories that originate from a stream
   - Use solid lines for categories 'LL' and 'SS'
   - Use dashed lines for categories 'LS' and 'SL'
```

This should reproduce a suggest plot:

```
ggplot2::ggplot(
  data = t_5,
  ggplot2::aes(
   x = cage_mass_mean_deviation_sd_rel,
   y = survived,
   color = origin transplant,
   lty = origin_transplant
) + ggplot2::scale_color_manual(
   values = c("blue", "blue", "green", "green")
 ) +
  ggplot2::scale_linetype_manual(
   values = c("solid", "dashed", "dashed", "solid")
  ggplot2::geom_point() +
  ggplot2::geom_smooth(
   method = "glm",
   method.args = list(family = "binomial"),
   se = FALSE
#> `geom_smooth()` using formula = 'y ~ x'
```



The only difference is that this plot uses relative body masses. The fit, however, is unfair: the binomial distribution is assumed to be monotonically increasing/decreasing. This means that a binomial distribution cannot be used to fit on data that is shaped like a U (as is assumed in the original paper: the extreme body masses have the highest fitness). Hence a different fit should be used.

Q2: Ask if the conclusion is correct

• A conclusion drawn from the data is that the extreme body masses are likelier to survive. Would you agree that the data supports this claim?

D5.3 Plot relative standarized body mass with parabolic fit

Ask the AI:

Create a scatter plot of this data:

- Each point is a fish.
- Use a blue color for fish that have 'Lake' as their origin.
- Use a green color for fish that have 'Stream' as their origin.
- On the X axis, put the values of `cage_mass_mean_deviation_sd_rel` of each fish
- On the Y axis, put the `survived` of each fish.
- Put the fish in four categories:
 - Fish that have an `origin` of 'Lake' and
 - a `transplant` of 'Lake' are in category 'LL'
 - Fish that have an `origin` of 'Lake' and

```
a `transplant` of 'Stream' are in category 'LS'
Fish that have an `origin` of 'Stream' and
    a `transplant` of 'Lake' are in category 'SL'
Fish that have an `origin` of 'Stream' and
    a `transplant` of 'Stream' are in category 'SS'
Show a trendline for each parabolic fit on each of these 4 categories.
Use blue lines for categories that originate from a lake
Use green lines for categories that originate from a stream
Use solid lines for categories 'LL' and 'SS'
Use dashed lines for categories 'LS' and 'SL'
```

The only change is to use a parabolic fit.

This should reproduce a suggest plot:

```
ggplot2::ggplot(
  data = t_5,
  ggplot2::aes(
   x = cage_mass_mean_deviation_sd_rel,
   y = survived,
   color = origin_transplant,
   lty = origin_transplant
) + ggplot2::scale_color_manual(
   values = c("blue", "blue", "green", "green")
  ggplot2::scale_linetype_manual(
   values = c("solid", "dashed", "dashed", "solid")
  ggplot2::geom_point() +
 ggplot2::geom_smooth(
   method = "lm",
   formula = y \sim x + I(x^2),
   se = FALSE
```

