

Bad data & Outliers

Paul Schmidt

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Cleaning data, dealing with missing data and comparing results for correlation and regression before vs. after removing an outlier from the data.

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```
# (install &) load packages
pacman::p_load(
  broom,
  conflicted,
  here,
  janitor,
  naniar,
  readxl,
  tidyverse
```

```
)

# handle function conflicts
conflicts_prefer(dplyr::filter)
conflicts_prefer(dplyr::select)
```

There are two download links:

- Download the **original** excel file [here](#).
- Download the **formatted** excel file [here](#).

1 Data

Imagine that this dataset was obtained by you. You spent an entire day walking around the campus of a university and asked a total of 29 people for things like how old they are and you also tested how well they could see on a scale of 1-10.

1.1 Import

Assuming you are working in a [R-project](#), save the formatted file somewhere within the project directory. I have saved it within a sub folder called **data** so that the relative path to my file is `data/vision_fixed.xls`.

```
path <- here("data", "vision_fixed.xls")
dat <- read_excel(path)
```

```
dat
```

```
# A tibble: 29 x 9
  Person    Ages Gender `Civil state` Height Profession Vision Distance PercDist
  <chr>    <dbl> <chr>   <chr>          <dbl> <chr>      <dbl>    <dbl>    <dbl>
1 Andrés    25 M     S              180 Student      10      1.5      15
2 Anja      29 F     S              168 Professio~  10      4.5      45
3 Armando  31 M     S              169 Professio~   9      4.5      50
4 Carlos   25 M     M              185 Professio~   8       6      75
5 Cristi~  23 F     <NA>          170 Student      10       3      30
6 Delfa    39 F     M              158 Professio~   6      4.5      75
7 Eduardo  28 M     S              166 Professio~   8      4.5     56.2
8 Enrique   NA <NA>   <NA>           NA Professio~  NA       6      NA
```

```

  9 Fanny      25 F      M      164 Student      9      3      33.3
10 Franci~    46 M      M      168 Professio~    8      4.5    56.2
# i 19 more rows

```

This is optional, but we could argue that our column names are not in a desirable format. To deal with this, we can use the `clean_names()` functions of `{janitor}`. This package has several more handy functions for cleaning data that are worth checking out.

```

dat <- dat %>% clean_names()
dat

# A tibble: 29 x 9
  person      ages gender civil_state height profession vision distance perc_dist
  <chr>    <dbl> <chr>   <chr>      <dbl> <chr>      <dbl>    <dbl>    <dbl>
1 Andrés      25 M      S        180 Student      10      1.5      15
2 Anja        29 F      S        168 Professio~    10      4.5      45
3 Armando     31 M      S        169 Professio~     9      4.5      50
4 Carlos      25 M      M        185 Professio~     8      6        75
5 Cristina    23 F      <NA>     170 Student      10      3        30
6 Delfa       39 F      M        158 Professio~     6      4.5      75
7 Eduardo     28 M      S        166 Professio~     8      4.5     56.2
8 Enrique     NA <NA>   <NA>      NA Professio~    NA      6        NA
9 Fanny       25 F      M        164 Student      9      3      33.3
10 Francis~   46 M      M        168 Professio~     8      4.5     56.2
# i 19 more rows

```

1.2 Goal

Very much like in the previous chapter, our goal is to look at the relationship of two numeric variables: `ages` and `vision`. What is new about this data is, that it (i) has missing values and (ii) has a potential outlier.

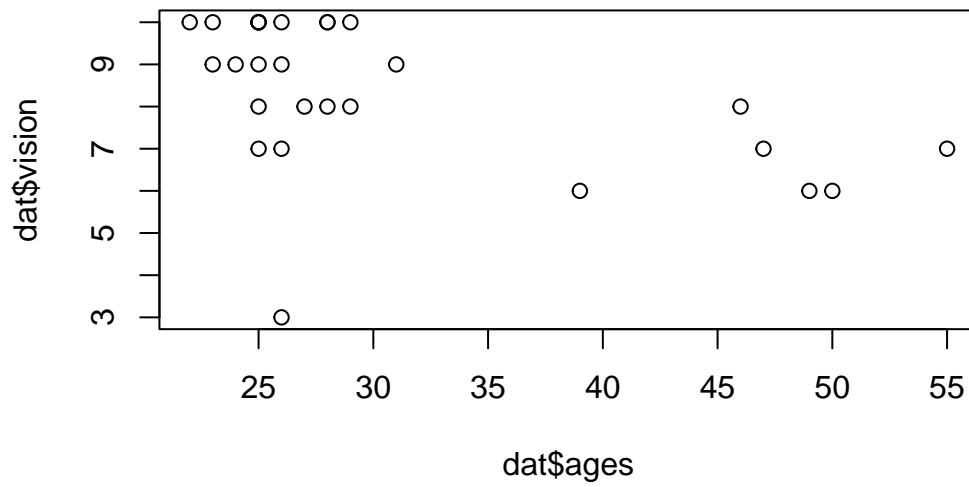
1.3 Exploring

To quickly get a first feeling for this dataset, we can use `summary()` and draw a plot via `plot()` or `ggplot()`.

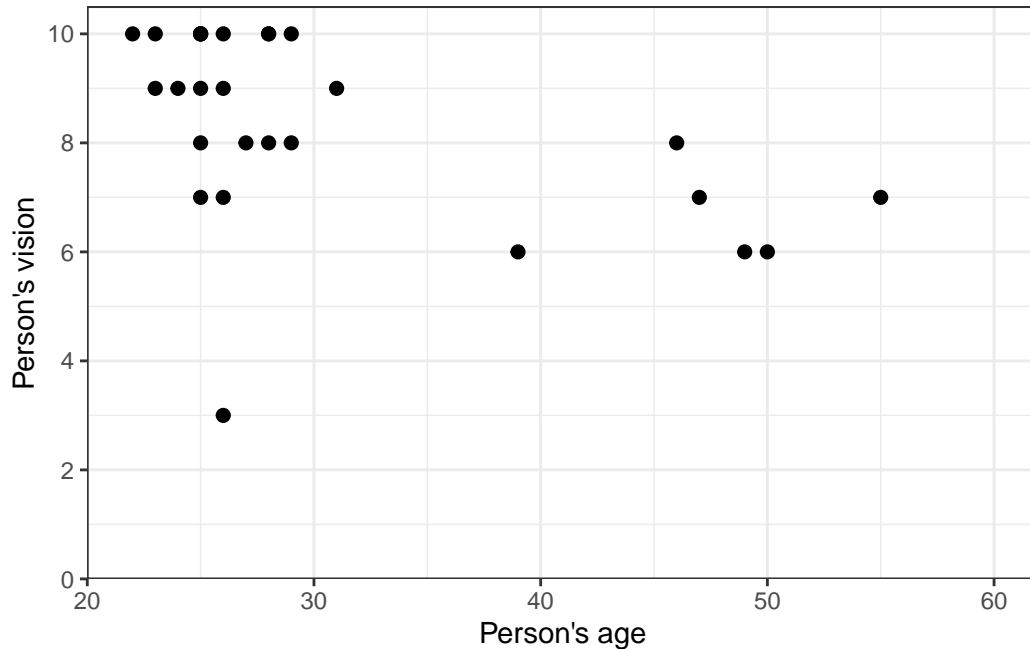
```
summary(dat)
```

person	ages	gender	civil_state
Length:29	Min. :22.00	Length:29	Length:29
Class :character	1st Qu.:25.00	Class :character	Class :character
Mode :character	Median :26.00	Mode :character	Mode :character
	Mean :30.61		
	3rd Qu.:29.50		
	Max. :55.00		
	NA's :1		
height	profession	vision	distance
Min. :145.0	Length:29	Min. : 3.000	Min. :1.500
1st Qu.:164.8	Class :character	1st Qu.: 7.000	1st Qu.:1.500
Median :168.0	Mode :character	Median : 9.000	Median :3.000
Mean :168.2		Mean : 8.357	Mean :3.466
3rd Qu.:172.8		3rd Qu.:10.000	3rd Qu.:4.500
Max. :190.0		Max. :10.000	Max. :6.000
NA's :1		NA's :1	
perc_dist			
Min. : 15.00			
1st Qu.: 20.24			
Median : 40.18			
Mean : 45.45			
3rd Qu.: 57.19			
Max. :150.00			
NA's :1			

```
plot(y = dat$vision, x = dat$ages)
```



```
ggplot(data = dat) +
  aes(x = ages, y = vision) +
  geom_point(size = 2) +
  scale_x_continuous(
    name = "Person's age",
    limits = c(20, 60),
    expand = expansion(mult = c(0, 0.05))
  ) +
  scale_y_continuous(
    name = "Person's vision",
    limits = c(0, NA),
    breaks = seq(0, 10, 2),
    expand = expansion(mult = c(0, 0.05))
  ) +
  theme_bw()
```



Apparently, most people are in their 20s and can see quite well, however some people are older and they tend to have a vision that's a little worse.

2 Missing Data?

While the data has 29 rows, it actually only holds **vision** and **ages** information for 28 people. This is because instead of values, there are NA (Not Available) for one person. Note that NA as missing values are treated somewhat special in R. As an example: If you want to filter for missing values, you cannot write `value == NA`, but must instead write `is.na(value)`:

```
dat %>%
  filter(is.na(vision))
```

```
# A tibble: 1 x 9
  person  ages gender civil_state height profession vision distance perc_dist
<chr>   <dbl> <chr>   <chr>         <dbl> <chr>         <dbl>   <dbl>   <dbl>
1 Enrique  NA <NA>   <NA>         NA Professional  NA       6      NA
```

Moreover, if you want to count the missing observations (per group) in a dataset, the most basic way of doing it is `sum(is.na(values))` (or for not-missing: `sum(!is.na(values))`). However,

if you are dealing with missing values a lot, you may also want to check out [{naniar}](#), which provides *principled, tidy ways to summarise, visualise, and manipulate missing data*.

```
# standard functions
dat %>%
  group_by(profession) %>%
  summarise(
    n_rows = n(),
    n_NA = sum(is.na(vision)),
    n_notNA = sum(!is.na(vision))
  )
```

```
# A tibble: 2 x 4
  profession    n_rows  n_NA n_notNA
  <chr>          <int> <int>   <int>
1 Professional     18     1     17
2 Student          11     0     11
```

```
# naniar functions
dat %>%
  group_by(profession) %>%
  summarise(
    n_rows = n(),
    n_NA = n_miss(vision),
    n_notNA = n_complete(vision)
  )
```

```
# A tibble: 2 x 4
  profession    n_rows  n_NA n_notNA
  <chr>          <int> <int>   <int>
1 Professional     18     1     17
2 Student          11     0     11
```

3 Corr. & Reg.

Let's estimate the correlation and simple linear regression and look at the results in a tidy format:

```
cor <- cor.test(dat$vision, dat$ages)
tidy(cor)
```

A tibble: 1 x 8

	estimate	statistic	p.value	parameter	conf.low	conf.high	method	alternative
	<dbl>	<dbl>	<dbl>	<int>	<dbl>	<dbl>	<chr>	<chr>
1	-0.497	-2.92	0.00709	26	-0.734	-0.153	Pearson's~	two.sided

```
reg <- lm(vision ~ ages, data = dat)
tidy(reg)
```

A tibble: 2 x 5

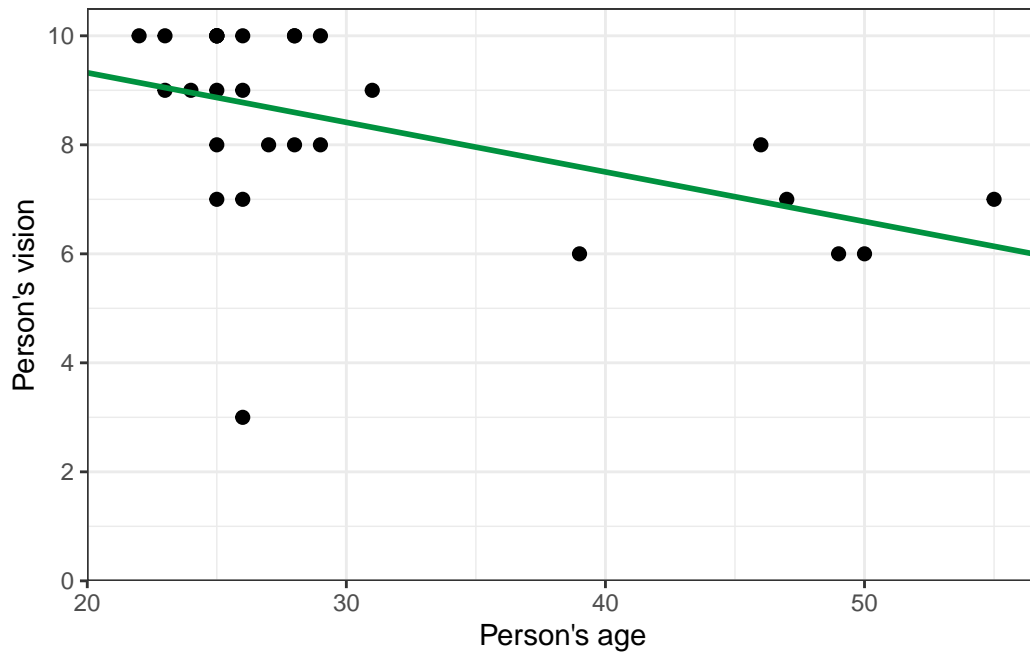
	term	estimate	std.error	statistic	p.value
	<chr>	<dbl>	<dbl>	<dbl>	<dbl>
1	(Intercept)	11.1	0.996	11.2	1.97e-11
2	ages	-0.0910	0.0311	-2.92	7.09e- 3

Thus, we have a moderate, negative correlation of -0.497 and for the regression we have *vision* = 11.14 + -0.09 *ages*. We can plot the regression line, too:

```
ggplot(data = dat) +
  aes(x = ages, y = vision) +
  geom_point(size = 2) +
  geom_abline(
    intercept = reg$coefficients[1],
    slope = reg$coefficients[2],
    color = "#00923f",
    linewidth = 1
  ) +
  scale_x_continuous(
    name = "Person's age",
    limits = c(20, 55),
    expand = expansion(mult = c(0, 0.05))
  ) +
  scale_y_continuous(
    name = "Person's vision",
    limits = c(0, NA),
    breaks = seq(0, 10, 2),
    expand = expansion(mult = c(0, 0.05))
  ) +
```



```
theme_bw()
```



4 Outlier?

Looking at the plot, you may find one data point to oddly stick out from all others: Apparently there was one person in their mid-20s who had a vision score of only 3, which is the lowest by far.

i Note

Here, we manually and thus subjectively identified a data point as a potential outlier. We do not discuss automatic and thus objective approaches for outlier detection, but see e.g. [here](#) or [here](#).

4.1 Step 1: Investigate

In such a scenario, the first thing you should do is find out more about this suspicious data point. In our case, we would start by finding out the person's name. One way of doing this is by simply filtering the data:

```
dat %>%
  filter(vision == 3)

# A tibble: 1 x 9
  person    ages gender civil_state height profession vision distance perc_dist
  <chr>    <dbl> <chr>   <chr>         <dbl> <chr>         <dbl>    <dbl>    <dbl>
1 Rolando    26 M      M             180 Professional      3        4.5      150
```

We find that it was 26 year old Rolando who supposedly had a vision score of only 3.

4.2 Step 2: Act

Since we pretend it is you who collected the data, you should now

- think back if you can actually remember Rolando and if he had poor vision and/or
- find other documents such as your handwritten sheets to verify this number and make sure you did not make any typos transferring the data to your computer.

This may reaffirm or correct the suspicious data point and thus end the discussion on whether it is an outlier that should be removed from the data. However, you may also decide to delete this value. Yet, it must be realized, that deleting one or multiple values from a dataset almost always affects the results from subsequent statistical analyses - especially if the values stick out from the rest.

5 Corr. & Reg. - again

Let us estimate correlation and regression again, but this time excluding Rolando from the dataset. Note that there are multiple ways of obtaining such a subset - two are shown here:

```
dat_noRo <- dat %>%
  filter(person != "Rolando")
```

```
dat_noRo <- dat %>%
  filter(vision > 3)
```

We now apply the same functions to this new dataset:

```
cor_noRo <- cor.test(dat_noRo$vision, dat_noRo$ages)
reg_noRo <- lm(vision ~ ages, data = dat_noRo)
```

and directly compare these results to those from above:

```
tidy(cor) %>%
  select(1, 3, 5, 6)

# A tibble: 1 x 4
  estimate p.value conf.low conf.high
  <dbl>    <dbl>    <dbl>    <dbl>
1  -0.497 0.00709  -0.734  -0.153

tidy(reg) %>%
  select(1, 2, 3)

# A tibble: 2 x 3
  term          estimate std.error
  <chr>          <dbl>    <dbl>
1 (Intercept)  11.1      0.996
2 ages         -0.0910   0.0311

tidy(cor_noRo) %>%
  select(1, 3, 5, 6)

# A tibble: 1 x 4
  estimate p.value conf.low conf.high
  <dbl>    <dbl>    <dbl>    <dbl>
1  -0.696 0.0000548  -0.851  -0.430

tidy(reg_noRo) %>%
  select(1, 2, 3)

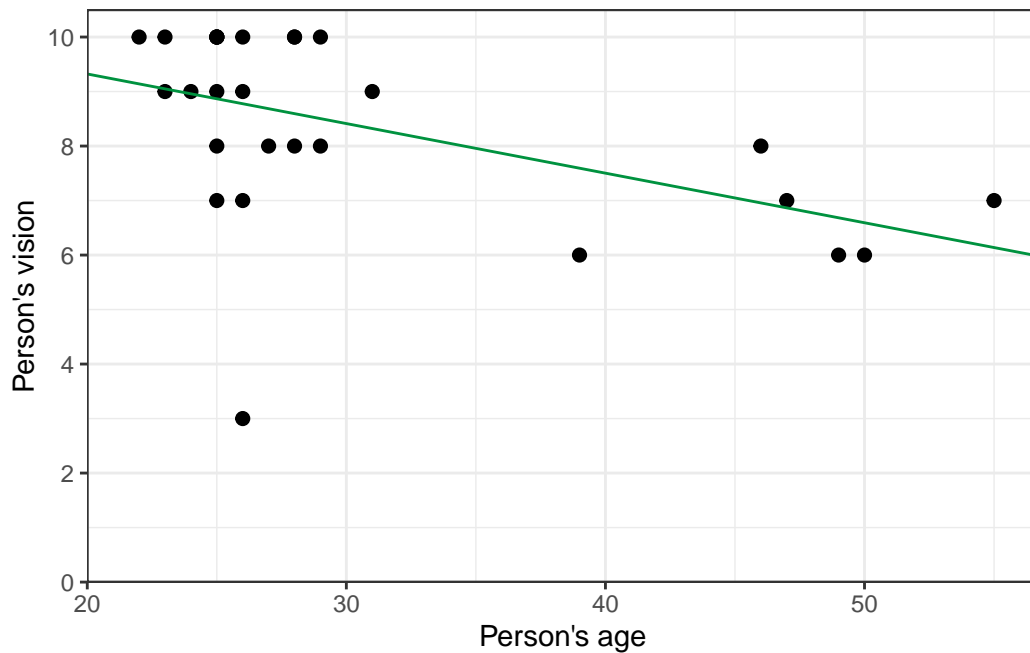
# A tibble: 2 x 3
  term          estimate std.error
  <chr>          <dbl>    <dbl>
1 (Intercept)  11.7      0.679
2 ages         -0.102   0.0211

ggplot(data = dat) +
  aes(x = ages, y = vision) +
  geom_point(size = 2) +
```

```

geom_abline(
  intercept = reg$coefficients[1],
  slope = reg$coefficients[2],
  color = "#00923f",
  linewidth = 1
) +
scale_x_continuous(
  name = "Person's age",
  limits = c(20, 55),
  expand = expansion(mult = c(0, 0.05))
) +
scale_y_continuous(
  name = "Person's vision",
  limits = c(0, NA),
  breaks = seq(0, 10, 2),
  expand = expansion(mult = c(0, 0.05))
) +
theme_bw()

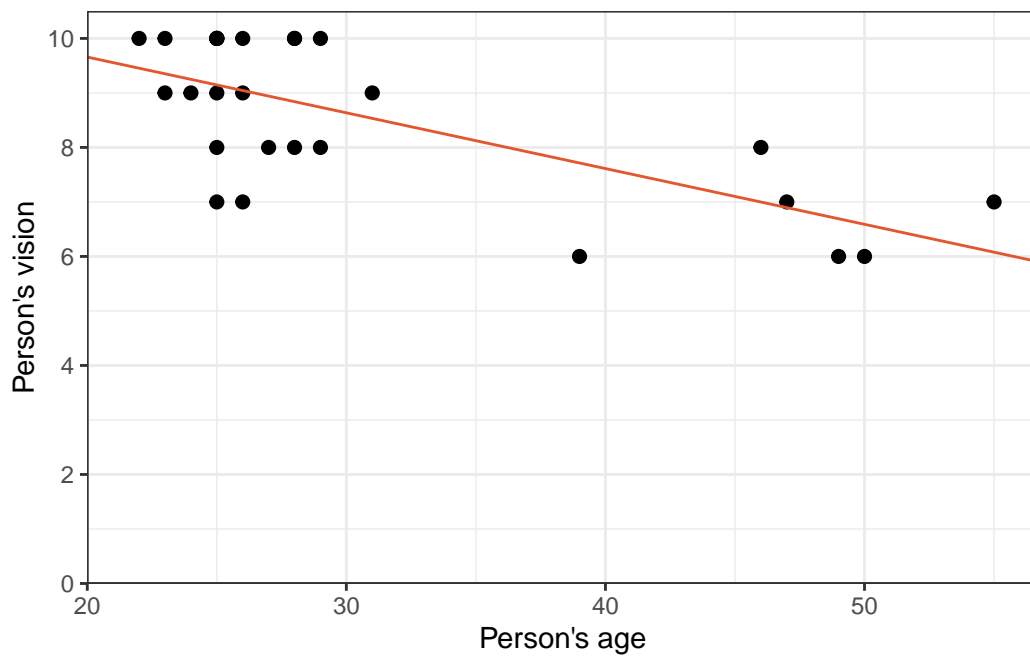
```



```

ggplot(data = dat_noRo) +
  aes(x = ages, y = vision) +
  geom_point(size = 2) +
  geom_abline(
    intercept = reg_noRo$coefficients[1],
    slope = reg_noRo$coefficients[2],
    color = "#e4572e",
    linewidth = 1
  ) +
  scale_x_continuous(
    name = "Person's age",
    limits = c(20, 55),
    expand = expansion(mult = c(0, 0.05))
  ) +
  scale_y_continuous(
    name = "Person's vision",
    limits = c(0, NA),
    breaks = seq(0, 10, 2),
    expand = expansion(mult = c(0, 0.05))
  ) +
  theme_bw()

```



As we can see, removing Rolando from the dataset changed the correlation quite a bit from -0.5 to -0.7. Furthermore, it's p-value became notably smaller. While it was already < 0.05 and thus statistically significant in this case, it must be realized that in other cases removing a single data point can indeed make the difference between a p-value larger and smaller 0.05.

Yet, regarding the parameter estimates - intercept (a) and slope (b) - of the simple linear regression, the changes are not as striking. Even with a visual comparison of the two regression lines, one must look closely to spot the differences.

5.1 R^2 - Coeff. of det.

Nevertheless, it is clear that the red line has a much better fit to the remaining data points, than the green line has - simply because Rolando's data point sticks out so much. One way of measuring how well a regression fits the data is by calculating the coefficient of determination R^2 , which measures the proportion of total variation in the data explained by the model and can thus range from 0 (=bad) to 1 (=good). It can easily be obtained via `glance()`, which is another function from `{broom}`:

```
glance(reg)
```

```
# A tibble: 1 x 12
  r.squared adj.r.squared sigma statistic p.value    df logLik   AIC   BIC
  <dbl>      <dbl> <dbl>     <dbl>   <dbl> <dbl> <dbl> <dbl> <dbl>
1    0.247        0.218  1.54      8.54 0.00709     1 -50.9  108.  112.
# i 3 more variables: deviance <dbl>, df.residual <int>, nobs <int>
```

```
glance(reg_noRo)
```

```
# A tibble: 1 x 12
  r.squared adj.r.squared sigma statistic  p.value    df logLik   AIC   BIC
  <dbl>      <dbl> <dbl>     <dbl>   <dbl> <dbl> <dbl> <dbl> <dbl>
1    0.485        0.464  1.04     23.5 0.0000548     1 -38.4  82.7  86.6
# i 3 more variables: deviance <dbl>, df.residual <int>, nobs <int>
```

Finally, we find that removing Rolando from the dataset increased the R^2 for the simple linear regression from 25% to 49%.