Lunch and Learn and Quarto

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1. Lunch and Learn and Quarto

This is an example Quarto project made for RStudio's Lunch and Learn on 8/3/2021. To learn more about Quarto visit https://quarto.org.

Part I. Section title

2. First Quarto

This is an example Quarto document. Note the qmd extension - this tells Quarto that this is a Markdown files that contains computations.

Since Quarto based on Markdown, we can **bold** and *italicize* text. We can also make headers.

3. Let's make a table

Meal	Food
Breakfast	Coffee
Lunch Dinner	Leftovers Spam Musubi

4. Example images



Figure 4.1.: My support system



Figure 4.2.: Lunch: Leftovers

Figure 4.3.: Dinner: Spam musubi

5. Code and chunk options

Quarto is based on RMarkdown, so you can do all the R stuff you're used to as well.

```
library(tidyverse)
standard_curves <- readxl::read_xlsx('std_curve.xlsx', sheet = "everything") %>%
  janitor::clean_names() %>%
  filter(amoa < 40)
lm_eqn = function(df){
    m = lm(log_qty \sim ct, df);
    data.frame(
      a = format(as.numeric(coef(m)[1]), digits = 2),
      b = format(as.numeric(coef(m)[2]), digits = 2),
      r2 = format(summary(m)$r.squared, digits = 3)
}
st_splits <- standard_curves %>%
  group_by(amoa, run) %>%
  group_split()
eqs <- st_splits %>%
  lapply(., lm_eqn) %>%
  bind_rows()
labels <- lapply(st_splits, slice_head, n = 1) %>%
  bind_rows() %>%
  select(amoa, run) %>%
  bind cols(eqs) %>%
  mutate(amoa = paste0("amoA_AOB_p", amoa)) %>%
  mutate(eq_label = paste0("y = ", a, " - ", abs(as.numeric(b)), "x<br>r^2 = ", r2))
standard curves %>%
  mutate(amoa = paste0("amoA_AOB_p", amoa)) %>%
```

```
ggplot(aes(log_qty, ct)) +
geom_point() +
facet_grid(run ~ amoa, scales = "free") +
 panel.border = element rect(color = "black", size = 1, fill = NA),
 panel.grid.minor.x = element_blank(),
 panel.grid.minor.y = element_blank(),
 panel.grid.major.x = element_line(color = "gray", size = 0.5, linetype = "dashed"),
 panel.grid.major.y = element line(color = "gray", size = 0.5, linetype = "dashed"),
 panel.spacing = unit(0.5, "lines"),
 panel.background = element_blank(),
 strip.background = element_rect(color = "black", size = 1, fill = NA),
) +
labs(
 x = "Log(gene copies per reaction)",
 y = "Ct"
) +
scale_x continuous(limits = c(0, 7), breaks = seq(0, 7, 1), expand = c(0, 0)) +
scale_y_continuous(limits = c(0, 25)) +
geom_smooth(aes(group=1), method="lm", se=FALSE) +
ggtext::geom_richtext(data = labels, aes(x = 3, y = 5, label = eq_label),
                      size = 4, fontface = "bold", inherit.aes = FALSE)
```

We can also throw some Python in here:

```
xs = [x for x in range(10)]
print(*(f'{x} squared is {x ^ 2}.' for x in xs), sep='\n')

0 squared is 2.
1 squared is 3.
2 squared is 0.
3 squared is 1.
4 squared is 6.
5 squared is 7.
6 squared is 4.
7 squared is 5.
8 squared is 10.
9 squared is 11.
```

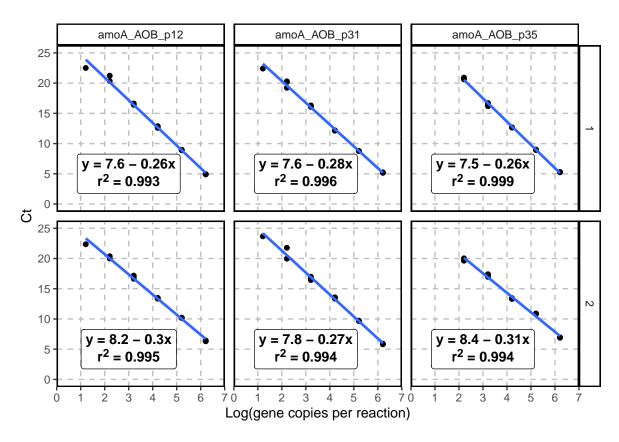


Figure 5.1.: Standard curves for the LAMPS crop priming experiment

```
import matplotlib.pyplot as plt
import numpy as np

Z = np.random.rand(6, 10)
x = [x + 0.5 for x in xs]
y = np.arange(4.5, 11, 1)

fig, ax = plt.subplots();
ax.pcolormesh(x, y, Z)
```

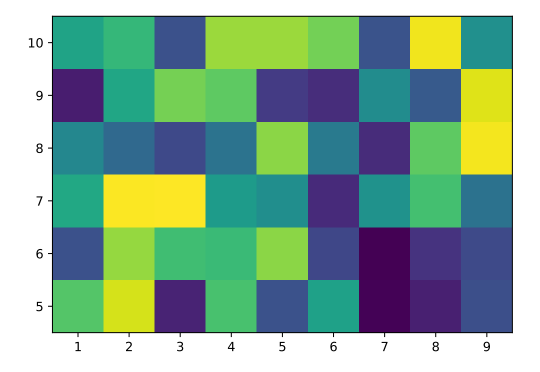


Figure 5.2.: That's a heatmap, baby!

5.1. Math stuff

We can also write math stuff! For example, here is a definition:

Definition 5.1 (Continuity). The function $f: \mathbb{R}^n \to \mathbb{R}^m$ is continuous at a point $x \in \mathbb{R}^n$ if for all $\varepsilon > 0$ there exists $\delta > 0$ such that if $|x - x_0| < \delta$, then $|f(x) - f(x_0)| < \varepsilon$. If this is true for all such x in the domain of f, we say that f is a continuous function.

5.1.0.1. Example

Define $f: \mathbb{R} \to \mathbb{R}$ by:

$$f(x) = \begin{cases} 1, x \in \mathbb{Q}, \\ 0, x \notin \mathbb{Q} \end{cases}$$

Prove that f is not a continuous function.

Proof. Let $\varepsilon = \frac{1}{2}$ and choose any $x \in \mathbb{Q}$. For any $\delta > 0$, we can find some $c \notin \mathbb{Q}$ such that $|x - c| < \delta$ since the irrationals are dense in \mathbb{R} . But then $|f(x) - f(c)| = |1 - 0| = 1 > \frac{1}{2}$, showing that f is not continuous at x.

5.2. Adding references

We can also add references. For instance, the following definition of k-partially colored comes from this paper: (Blair et al. 2020)

Definition 5.2 (k-partially colored). Let D be a diagram of a link L with n crossings. We call D k-partially colored if we have specified a subset A of the strands of D and a function $f : \to \{1, 2, ..., k\}$. We refer to this partial coloring by the tuple (A, f). Given k-partial colorings (A_1, f_1) and (A_2, f_2) of D, we say (A_2, f_2) is the result of a coloring move on (A_1, f_1) if

- 1. $A_1 \subset A_2$ and $A_2 A_1 = \{s_i\}$ for some strand s_i in D;
- 2. $f_2|_{A_1}=f_1$;
- 3. s_i is adjacent to s_i at some crossing $c \in v(D)$, and $s_i \in A_1$;
- 4. the over-strand s_k at c is an element of A_1 ;
- 5. $f_1(s_i) = f_2(s_j)$.

5.3. Cross-references

Along the way, we've been giving each of the items above labels. The Visual Editor knows about these labels and we can call them up for cross referencing. For example:

- \bullet We were pretty happy about the standard curves in fig. 5.1
- $\bullet\,$ I love me some heatmaps like fig. $5.2\,$
- def. 5.2 is trivially true for the unknot.

6. Bibliography

7. Second: A Pure Python qmd

This is a pure Python qmd document. Since there are no R code chunks, it is executed via the Jupyter kernel.

7.1. Adding days per month from date range to a dataframe

Suppose you have a dataset with a column of start dates and column of end dates. For example:

```
import pandas as pd
import calendar

date_df = pd.DataFrame({
    "START_TM": ['2/15/2010', '2/15/2010', '3/16/2010'],
    "END_TM": ['4/18/2010', '2/18/2010', '5/20/2010']
})
date_df["START_TM"] = date_df["START_TM"].astype('datetime64')
date_df["END_TM"] = date_df["END_TM"].astype('datetime64')
date_df
```

	START_TM	END_TM
0	2010-02-15	2010-04-18
1	2010-02-15	2010-02-18
2	2010-03-16	2010-05-20

Our goal is to count the number of days in each month this range of dates falls over.

We start by adding columns for each month:

```
months = {calendar.month_name[i]:[0 for _ in range(date_df.shape[0])] for i in range(1, 13)}
for m in months:
    date_df[m] = [0 for _ in range(date_df.shape[0])]
date_df
```

	START_TM	END_TM	January	February	March	April	May	June	July	August	Septemb
0	2010-02-15	2010-04-18	0	0	0	0	0	0	0	0	
1	2010-02-15	2010-02-18	0	0	0	0	0	0	0	0	
2	2010-03-16	2010-05-20	0	0	0	0	0	0	0	0	

7.1.1. Helper functions

```
def insert_days_per_month(outer_row):
    dpm = days_per_month(outer_row)
    for index, inner_row in dpm.iterrows():
        outer_row[inner_row['Month']] = inner_row['NumDays']
    return(outer_row)

def days_per_month(row):
    s = pd.Series(index = pd.date_range(row[0], row[1]))[1: ]
    days_in_month = s.resample('MS').size().to_period('m').\
    rename_axis('Month').reset_index(name = 'NumDays')
    days_in_month['Month'] = days_in_month['Month'].apply(
        lambda x: calendar.month_name[x.month])
    return(days_in_month)
```

We can get the desired result with apply:

```
date_df = date_df.apply(lambda x: insert_days_per_month(x), axis = 1)
date_df
```

DeprecationWarning:

The default dtype for empty Series will be 'object' instead of 'float64' in a future version

	START_TM	END_TM	January	February	March	April	May	June	July	August	Septemb
0	2010-02-15	2010-04-18	0	13	31	18	0	0	0	0	
1	2010-02-15	2010-02-18	0	3	0	0	0	0	0	0	
2	2010-03-16	2010-05-20	0	0	15	30	20	0	0	0	

7.2. EXTRA EXTRA READ ALL ABOUT IT HOT OFF THE PRESSES

Quarto has support for interactive documents. Support formats include:

- JavaScript: Observable JS
- R: Shiny
- Python: Jupyter Widgets are all supported, such as IPyLeaflet and Plotly

7.3. IPyLeaflet

```
from ipyleaflet import Map, Marker

csg_loc = (33.772819, -117.9694484)

cham_soot_gol = Map(center=csg_loc, scroll_wheel_zoom=True)
cham_soot_gol.add_layer(Marker(location=csg_loc, title="Cham Soot Gol"))
cham_soot_gol
```

 ${\tt Map(center=[33.772819, -117.9694484], controls=(ZoomControl(options=['position', 'zoom_in_tentoric and options=['position', 'zoom_in_tentoric and opt$

Can do everything you're used to with Python but with the awesome Visual Editor stuff:

7.4. Plotly

```
Unable to display output for mime type(s): text/html
Unable to display output for mime type(s): text/html
```

8. Example Jupyter Notebook

The editing experience with Jupyter + Quarto is very similar to the RStudio editing experience.

I'm not lying!

When we make changes and save here, the preview will update. Here's some code:

```
for x in range(10):
    print(f'{x} squared is {x ^2}.')
```

```
O squared is 2.

1 squared is 3.

2 squared is 0.

3 squared is 1.

4 squared is 6.

5 squared is 7.

6 squared is 4.

7 squared is 5.

8 squared is 10.

9 squared is 11.
```

Here's a figure.

```
import numpy as np
import matplotlib.pyplot as plt

r = np.arange(0, 2, 0.01)
theta = 2 * np.pi * r
fig, ax = plt.subplots(subplot_kw={'projection': 'polar'})
ax.plot(theta, r)
ax.set_rticks([0.5, 1, 1.5, 2])

ax.grid(True)
plt.show()
```

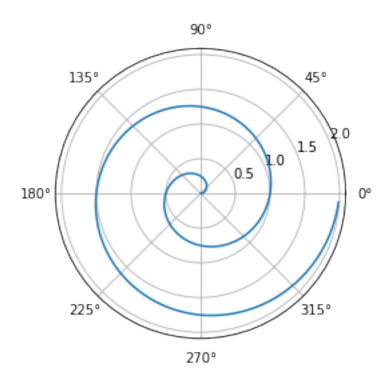


Figure 8.1.: ?(caption)

And we can add chunk options just like we did in RStudio.

See fig. 8.1 for an example of a projection of a straight line into polar coordinates.

9. DNA String Stuff

Here are some functions to do some basic DNA string calculations.

```
import pandas as pd
def reverse_complement(nuc_sequence: str) -> str:
   Returns the reverse complement of a nucleotide sequence.
   >>> reverse_complement('ACGT')
   'ACGT'
   >>> reverse_complement('ATCGTGCTGTCGTCAAGAC')
    'GTCTTGACGACAGCACCACT'
   >>> reverse_complement('TGCTAGCATCGAGTCGATCGATATATTTAGCATCAGCATT')
    'AATGCTGATGCTAAATATATCGATCGACTCGATGCTAGCA'
    11 11 11
    complements = {
        "A": "T",
        "C": "G",
        "G": "C".
        "T": "A"
   rev_seq = "".join([complements[s] for s in nuc_sequence.upper()[::-1]])
   return rev seq
def gc_content(nuc_sequence: str) -> float:
   Calculates the GC content of a nucleotide sequence.
   >>> gc_content('ACGT')
   0.5
    11 11 11
   gc_tally = 0
   for nuc in nuc_sequence.lower():
        if nuc == 'g' or nuc == 'c':
            gc_tally += 1
    return gc_tally / len(nuc_sequence)
def random_dna_string(seq_length: int = 10) -> str:
```

```
Generates a random DNA string seq_length bp long
    >>> len(random_dna_string())
   >>> len(random_dna_string(20))
    20
    11 11 11
   from random import choice
    dna string = ""
    for _ in range(seq_length):
        dna_string += choice("ACGT")
    return dna_string
def make_strings_df(num_strings: int = 10, str_length: int = 10) -> pd.DataFrame:
   Generates a pandas dataframe with num strings DNA sequences of length str length with
   columns "Sequence", "GC Content", "Reverse Complement"
   >>> df = make_strings_df(100, 37)
   >>> df.shape
    (100, 3)
   >>> len(df['Sequence'][0])
    37
    dna_strings_list = [random_dna_string(str_length) for _ in range(num_strings)]
    strings_df = pd.DataFrame({
        "Sequence": dna_strings_list
    })
    strings_df['GC Content'] = strings_df['Sequence'].apply(gc_content)
    strings_df['Reverse Complement'] = strings_df['Sequence'].apply(reverse_complement)
    return strings_df
import doctest
doctest.testmod(verbose=0)
```

TestResults(failed=0, attempted=9)

But that's a lot of function definitions and code testing that a lot of people probably don't care about. Let's set fold and summary to hide this chunk.

Let's use the function and create a histogram of the GC contents for the simulated sequences.

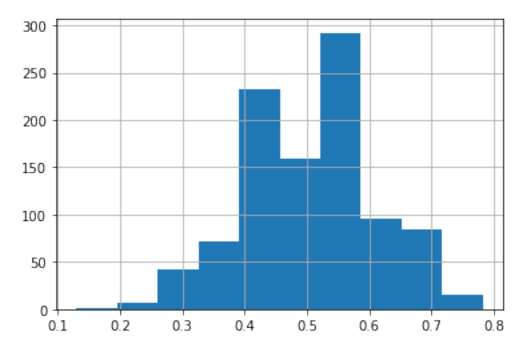
```
strings_df = make_strings_df(1000, 23)
print(f'strings_df has {strings_df.shape[0]} rows and {strings_df.shape[1]} columns.')
```

strings_df has 1000 rows and 3 columns.

strings_df.head(10)

	Sequence	GC Content	Reverse Complement
0	TAATAATGGGCTAAACTATGTTT	0.260870	AAACATAGTTTAGCCCATTATTA
1	GACCGTGACCCAAGGCAGATGGG	0.652174	CCCATCTGCCTTGGGTCACGGTC
2	TAGGGTTGTGCTTTACCTTACAT	0.391304	ATGTAAGGTAAAGCACAACCCTA
3	GCAAGGCCGGATACGCGTATAAT	0.521739	ATTATACGCGTATCCGGCCTTGC
4	ACCACTCCTCAAACGTTACTGAT	0.434783	ATCAGTAACGTTTGAGGAGTGGT
5	CCTCGTCAGTTGTCACTTCTATG	0.478261	CATAGAAGTGACAACTGACGAGG
6	ACAATGATCGCAGCCGAGGTATA	0.478261	TATACCTCGGCTGCGATCATTGT
7	GTTGGATATTCCGCAGCAGAGGA	0.521739	TCCTCTGCTGCGGAATATCCAAC
8	CGCTTAAAATCCCTGCATAGACC	0.478261	GGTCTATGCAGGGATTTTAAGCG
9	AGACCACTACTGGGTGGAGACGG	0.608696	CCGTCTCCACCCAGTAGTGGTCT

strings_df['GC Content'].hist();



Blair, R., A. Kjuchukova, R. Velazquez, and P. Villanueva. 2020. "Wirtinger Systems of