# DapR1: Notes on the Live R Session, Week 8

This week, we will be picking up where we left off the last time. We'll continue working with probability using operators. We'll also talk about writing a clean report using Markdown, including learning about in-line R coding and notation.

First, let's load the tidyverse and create our sample space. We'll also recreate our sample data from last week. Because we used the *set.seed()* function last week, we can use the same seed value to make R generate the same data we used last week.

```
library(tidyverse)
skittles <- c('red', 'green', 'yellow', 'purple', 'orange')
set.seed(2210)
s5 <- tibble(Obs = sample(skittles, 5, replace = T))
s100 <- tibble(Obs = sample(skittles, 100, replace = T))</pre>
```

Although we're working with character data in this example, you can also create sample data with numeric values. Consider an example where you want to calculate the probability of rolling an even number on a die. You could use seq() function, which allows you to generate sequences of numbers, to create your sample space and your events of interest:

dSp <- seq(1, 6) dEv <- seq(2, 6, by = 2)

## Joint Probability

With joint probability, we're interested in looking at the probability of multiple outcomes. For example, if we were to select a skittle from the bag, what's the likelihood we would select a yellow **OR** a purple?

Because these events are mutually exclusive, we would expect their joint probability to be the sum of their individual probabilities.

 $p(yellow \cup purple) = .20 + .20 = .40$ 

Here, we'll review two ways you can specify multiple events of interest and use these methods to compute joint probability in your sample.

#### The AND and OR operators

Two operators that may be used for specifying multiple conditions are & and |. As you might expect, & refers to 'and'. It allows you to specify values that meet all given conditions. The | operator stands for 'or' and allows you to specify values that meet any of the given conditions.

```
successes <- s100 == 'yellow'|s100 == 'purple'
failures <- s100 != 'yellow' & s100 != 'purple'</pre>
```

```
sum(successes)/(sum(successes) + sum(failures))
```

## [1] 0.4

## %in% or is.element()

You can get the same results using the %in% operator. This operator can be used to determine whether a given element (or values from a vector of elements) is found in a dataset or vector. Also, note the use of another operator. The ! operator stands for 'not' and allows you to select values that do not meet a specified condition.

```
events <- c('purple', 'yellow')
successes <- s100$Obs %in% events
failures <- !s100$Obs %in% events</pre>
```

```
sum(successes)/(sum(successes) + sum(failures))
```

## [1] 0.4

You can also use the **is.element()** function in the same way. *is.element(x, y)* is identical to  $x \ \% in \% y$ 

```
is.element(s100$0bs, events)
```

```
##
     [1] FALSE
                TRUE TRUE FALSE FALSE
                                        TRUE FALSE
                                                    TRUE
                                                          TRUE FALSE
                                                                       TRUE
                                                                            TRUE
##
    [13] FALSE
               TRUE FALSE FALSE FALSE
                                        TRUE FALSE
                                                    TRUE
                                                          TRUE FALSE
                                                                       TRUE FALSE
    [25] FALSE FALSE FALSE
                           TRUE FALSE
                                        TRUE FALSE
                                                    TRUE
                                                          TRUE
                                                                       TRUE FALSE
##
                                                                 TRUE
         TRUE FALSE FALSE
                            TRUE FALSE FALSE FALSE FALSE FALSE
##
   [37]
                                                                 TRUE
                                                                       TRUE TRUE
##
    [49] FALSE FALSE FALSE
                            TRUE
                                 TRUE FALSE
                                              TRUE FALSE
                                                          TRUE FALSE
                                                                       TRUE FALSE
##
    [61] FALSE
               TRUE FALSE FALSE FALSE
                                              TRUE
                                                    TRUE FALSE FALSE
                                                                       TRUE TRUE
                                        TRUE
##
   [73]
         TRUE
               TRUE FALSE FALSE FALSE FALSE FALSE FALSE
                                                          TRUE FALSE FALSE FALSE
##
   [85] FALSE FALSE FALSE FALSE FALSE
                                        TRUE FALSE FALSE
                                                          TRUE FALSE FALSE FALSE
    [97] FALSE
               TRUE FALSE FALSE
##
successes <- is.element(s100$0bs, events)</pre>
```

```
failures <- !is.element(s100$0bs, events)
sum(successes)/(sum(successes) + sum(failures))</pre>
```

## [1] 0.4

Now, let's compute the probability of selecting a yellow **AND** a purple skittle in two selections.

To do this, we would use we use the following formula:

 $p(yellow \cap purple) = .20 * .20 = .04$ 

## 6 yellow purple

Let's check to see if our data follow the expected result. To do this, we'll need to add a new column that represents the second skittle selection:

```
s100$Obs2 <- sample(skittles, 100, replace = T)
head(s100)
## # A tibble: 6 x 2
## Obs Obs2
## <chr> <chr> ## 1 green orange
## 2 yellow orange
## 3 yellow orange
## 4 orange purple
## 5 red purple
```

```
successes <- (s100$0bs=='purple'&s100$0bs2=='yellow')|(s100$0bs=='yellow'&s100$0bs2=='purple')
failures <- !successes</pre>
```

```
sum(successes)/(sum(successes) + sum(failures))
```

## [1] 0.05

You can also look at the proportion table for multiple events:

```
s100 %>%
table %>%
prop.table()
```

## Obs2 ## Obs green orange purple red yellow ## 0.01 0.05 0.01 0.05 0.06 green 0.06 0.02 orange 0.04 0.05 0.03 ## ## purple 0.03 0.08 0.06 0.02 0.02 ## red0.06 0.04 0.05 0.04 0.03 ## yellow 0.06 0.06 0.03 0.03 0.01

Note that you can sum the appropriate locations in the proportion table to get the corresponding probability.

```
pTab <- s100 %>%
  table() %>%
  prop.table()
pTab['purple', 'yellow'] + pTab['yellow', 'purple']
```

## [1] 0.05

You can also compute the summed values for each row or each column using the addmargins function:

```
pTab %>%
    addmargins()
```

##	(	]bs2					
##	Obs	green	orange	purple	red	yellow	Sum
##	green	0.01	0.05	0.01	0.05	0.06	0.18
##	orange	0.04	0.05	0.06	0.02	0.03	0.20
##	purple	0.03	0.08	0.06	0.02	0.02	0.21
##	red	0.06	0.04	0.05	0.04	0.03	0.22
##	yellow	0.06	0.06	0.03	0.03	0.01	0.19
##	Sum	0.20	0.28	0.21	0.16	0.15	1.00

sum(pTab[1,])

## [1] 0.18

sum(pTab[,2])

### ## [1] 0.28

However, the table outputs in R are not appropriate for a formal report. If you want to include a table in your report, you should use the kbl() function to produce a formal table to display your data:

```
library(kableExtra)
```

```
pTab %>%
kbl(booktabs = T, digits = 2, caption = 'Skittles Experiment Data') %>%
kable_styling(latex_options = c('hold_position', 'striped')) %>%
column_spec(1, bold = T) %>%
row_spec(0, bold = T)
```

	green	orange	purple	$\mathbf{red}$	yellow
green	0.01	0.05	0.01	0.05	0.06
orange	0.04	0.05	0.06	0.02	0.03
purple	0.03	0.08	0.06	0.02	0.02
red	0.06	0.04	0.05	0.04	0.03
yellow	0.06	0.06	0.03	0.03	0.01

Table 1: Skittles Experiment Data

## Conditional Probability Data & Write-Up

Now let's import some data that we can use to demonstrate conditional probability. We'll also write up our results neatly and talk about in-line R coding.

Imagine that we want to investigate the relationship between med school acceptance and academic performance. Specifically we will look at the likelihood of acceptance (yes or no) is related to having higher marks (e.g., above the 75th percentile or greater in our sample; high or...less high. We'll just say low for simplicity's sake).

```
dat <- read.csv('https://uoepsy.github.io/data/MedGPA.csv')
summary(dat)</pre>
```

##	Accept	Acceptance	Sex	BCPM
##	Length:55	Min. :0.000	) Length:55	Min. :2.410
##	Class :characte	r 1st Qu.:0.000	O Class :charact	ter 1st Qu.:3.260
##	Mode :characte	r Median :1.000	O Mode :charact	ter Median :3.530
##		Mean :0.545	5	Mean :3.501
##		3rd Qu.:1.000	C	3rd Qu.:3.755
##		Max. :1.000	C	Max. :4.000
##				
##	GPA	VR	PS	WS
##	Min. :2.720	Min. : 6.000	Min. : 5.000	Min. : 4.000
##	1st Qu.:3.375	1st Qu.: 8.000	1st Qu.: 9.000	1st Qu.: 6.000
##	Median :3.580	Median :10.000	Median :10.000	Median : 8.000

```
##
    Mean
           :3.553
                            : 9.764
                                              : 9.709
                                                                 : 7.148
                     Mean
                                       Mean
                                                         Mean
##
    3rd Qu.:3.770
                     3rd Qu.:11.000
                                       3rd Qu.:10.500
                                                         3rd Qu.: 8.000
##
    Max.
           :3.970
                     Max.
                            :13.000
                                       Max.
                                              :14.000
                                                         Max.
                                                                 :10.000
##
                                                         NA's
                                                                 :1
##
          BS
                           MCAT
                                            Apps
##
   Min.
           : 6.000
                             :18.00
                                              : 1.000
                      Min.
                                       Min.
    1st Qu.: 9.000
                      1st Qu.:34.00
                                       1st Qu.: 5.000
##
                                       Median : 7.000
##
   Median :10.000
                      Median :36.00
##
    Mean
           : 9.782
                      Mean
                             :36.27
                                       Mean
                                              : 8.364
##
    3rd Qu.:11.000
                      3rd Qu.:39.00
                                       3rd Qu.:11.000
##
    Max.
           :14.000
                      Max.
                             :48.00
                                       Max.
                                              :24.000
##
dat$acceptChar <- ifelse(dat$Acceptance == '0', 'Rejected', 'Accepted')</pre>
dat$GPAsplit <- ifelse(dat$GPA >= quantile(dat$GPA)['75%'], 'High', 'Low')
pTab <- table(dat$GPAsplit, dat$acceptChar) %>%
  prop.table() %>%
  round(., 2)
pTab
```

## Accepted Rejected ## High 0.25 0.02 ## Low 0.29 0.44

Remember from yesterday's lecture that we can calculate conditional probability using the following formula:

$$p(A|B) = \frac{p(A \cap B)}{p(B)}$$

Let's compare the probability of being accepted given that marks are high with the probability given that marks are low. In other words, we are comparing an outcome of event A across different levels of event B. If these events were unrelated, we would expect the probability of event A to be generally consistent across both levels of B. To do this, we can plug our variables into the formula above:

$$p(Accepted|High) = \frac{p(Accepted \cap High)}{p(High)}$$
$$p(Accepted|Low) = \frac{p(Accepted \cap Low)}{p(Low)}$$

Remember, we can pull specific values from the probability table we've created.

The probability of being accepted when marks are high:

pTab['High', 'Accepted']/(sum(pTab['High',]))

## [1] 0.9259259

The probability of being accepted when marks are low:

#### ## [1] 0.3972603

Here, we see that the values are quite different, which indicates that the probability of acceptance is likely to be related to overall marks (shocking!). Now let's show an example of how to write up these results.

## **EXAMPLE WRITE-UP**

In this experiment, we collected data on medical school acceptance rates and school performance from 55 participants. Specifically, we investigated whether acceptance to medical school was related to overall performance as measured by GPA (High/Low). We calculated the proportion of participants who fell into each category (see Table 2).

	Accepted	Rejected
High	0.25	0.02
Low	0.29	0.44

Table 2: Medical School Acceptance by GPA

We computed the probability of acceptance at both levels of GPA.  $P(Acceptance|GPA_{high}) = 0.93$ .  $P(Acceptance|GPA_{low}) = 0.4$ . This indicates that the likelihood of being accepted changes at different levels of GPA.

The differences in the probability of acceptance across levels of GPA indicate a possible relationship between school performance and acceptance to medical school.