# Multiway Frequency Tables

# **Packages**

library(tidyverse )

## Multi-way frequency analysis

➤ A study of gender and eyewear-wearing finds the following frequencies:

```
gender contacts glasses none female 121 32 129 male 42 37 85
```

- Is there association between eyewear and gender?
- Normally answer this with chisquare test (based on observed and expected frequencies from null hypothesis of no association).
- Two categorical variables and a frequency.
- We assess in way that generalizes to more categorical variables.

#### The data file

```
gender contacts glasses none female 121 32 129 male 42 37 85
```

- This is not tidy!
- Two variables are gender and eyewear, and those numbers all frequencies.

```
# A tibble: 2 x 4
gender contacts glasses none
<chr> <dbl> <dbl> <dbl> <dbl> 1 female 121 32 129
2 male 42 37 85
```

# Tidying the data

3 female none

4 male contacts

5 male glasses 6 male none

```
eyewear %>%
  pivot longer(contacts:none, names to="eyewear",
              values to="frequency") -> eyes
eyes
# A tibble: 6 x 3
  gender eyewear frequency
  <chr> <chr> <dbl>
1 female contacts
                       121
2 female glasses
                  32
```

129

85

42 37

# Making tidy data back into a table

- use pivot\_wider
- or this (we use it again later):

```
xt <- xtabs(frequency ~ gender + eyewear, data = eyes)
xt</pre>
```

```
eyewear
gender contacts glasses none
female 121 32 129
male 42 37 85
```

## Modelling

- Predict frequency from other factors and combos.
- plm with poisson family.

```
eyes.1 <- glm(frequency ~ gender * eyewear,
  data = eyes,
  family = "poisson"
)</pre>
```

Called log-linear model.

# What can we get rid of?

```
drop1(eyes.1, test = "Chisq")
Single term deletions
Model:
frequency ~ gender * eyewear
              Df Deviance AIC LRT Pr(>Chi)
<none>
                   0.000 47.958
gender:eyewear 2 17.829 61.787 17.829 0.0001345 ***
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
nothing!
```

#### Conclusions

- drop1 says what we can remove at this step. Significant = must stay.
- Cannot remove anything.
- ► Frequency depends on gender-wear combination, cannot be simplified further.
- Gender and eyewear are associated.
- Stop here.

#### prop.table

#### Original table:

xt

#### eyewear

```
gender contacts glasses none
female 121 32 129
male 42 37 85
```

Calculate eg. row proportions like this:

```
prop.table(xt, margin = 1)
```

#### eyewear

```
gender contacts glasses none
female 0.4290780 0.1134752 0.4574468
male 0.2560976 0.2256098 0.5182927
```

#### Comments

- margin says what to make add to 1.
- ▶ More females wear contacts and more males wear glasses.

#### No association

#### Suppose table had been as shown below:

#### eyewear

```
gender contacts glasses none female 0.5000000 0.1000000 0.4000000 male 0.4901961 0.1045752 0.4052288
```

#### Comments

- Females and males wear contacts and glasses in same proportions
  - though more females and more contact-wearers.
- No association between gender and eyewear.

### Analysis for revised data

```
eyes.2 <- glm(frequency ~ gender * eyewear,
  data = eves2,
  family = "poisson"
drop1(eyes.2, test = "Chisq")
Single term deletions
Model:
frequency ~ gender * eyewear
               Df Deviance AIC LRT Pr(>Chi)
<none>
                  0.000000 47.467
gender:eyewear 2 0.047323 43.515 0.047323 0.9766
No longer any association. Take out interaction.
```

#### No interaction

```
eyes.3 <- update(eyes.2, . ~ . - gender:eyewear)</pre>
drop1(eyes.3, test = "Chisq")
Single term deletions
Model:
frequency ~ gender + eyewear
       Df Deviance AIC LRT Pr(>Chi)
<none> 0.047 43.515
gender 1 48.624 90.091 48.577 3.176e-12 ***
eyewear 2 138.130 177.598 138.083 < 2.2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
 More females (gender effect)
 more contact-wearers (eyewear effect)
 no association (no interaction).
```

# Chest pain, being overweight and being a smoker

- ▶ In a hospital emergency department, 176 subjects who attended for acute chest pain took part in a study.
- ► Each subject had a normal or abnormal electrocardiogram reading (ECG), were overweight (as judged by BMI) or not, and were a smoker or not.
- ▶ How are these three variables related, or not?

#### The data

In modelling-friendly format:

ecg bmi smoke count abnormal overweight yes 47 abnormal overweight no 10 abnormal normalweight yes 8 abnormal normalweight no 6 normal overweight yes 25 normal overweight no 15 normal normalweight yes 35 normal normalweight no 30

#### First step

```
my_url <- "http://ritsokiguess.site/datafiles/ecg.txt"
chest <- read_delim(my_url, " ")
chest.1 <- glm(count ~ ecg * bmi * smoke,
    data = chest,
    family = "poisson"
)
drop1(chest.1, test = "Chisq")</pre>
```

That 3-way interaction comes out.

#### Removing the 3-way interaction

```
chest.2 <- update(chest.1, . ~ . - ecg:bmi:smoke)</pre>
drop1(chest.2, test = "Chisq")
Single term deletions
Model:
count ~ ecg + bmi + smoke + ecg:bmi + ecg:smoke + bmi:smoke
         Df Deviance AIC LRT Pr(>Chi)
<none> 1.3885 53.096
ecg:bmi 1 29.0195 78.727 27.6310 1.468e-07 ***
ecg:smoke 1 4.8935 54.601 3.5050 0.06119 .
bmi:smoke 1 4.4689 54.176 3.0803 0.07924 .
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
At \alpha = 0.05, bmi:smoke comes out.
```

#### Removing bmi:smoke

```
chest.3 <- update(chest.2, . ~ . - bmi:smoke)</pre>
drop1(chest.3, test = "Chisq")
Single term deletions
Model:
count ~ ecg + bmi + smoke + ecg:bmi + ecg:smoke
         Df Deviance AIC LRT Pr(>Chi)
<none> 4.469 54.176
ecg:bmi 1 36.562 84.270 32.094 1.469e-08 ***
ecg:smoke 1 12.436 60.144 7.968 0.004762 **
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
ecg:smoke has become significant. So we have to stop.
```

# Understanding the final model

- ▶ Thinking of ecg as "response" that might depend on anything else.
- ▶ What is associated with ecg? Both bmi on its own and smoke on its own, but *not* the combination of both.
- ecg:bmi table:

```
xtabs(count ~ ecg + bmi, data = chest)
```

```
bmi
ecg normalweight overweight
abnormal 14 57
normal 65 40
```

▶ Most normal weight people have a normal ECG, but a majority of overweight people have an abnormal ECG. That is, knowing about BMI says something about likely ECG. ecg:smoke

ecg:smoke table:

```
xtabs(count ~ ecg + smoke, data = chest)
```

```
smoke
ecg no yes
abnormal 16 55
normal 45 60
```

- ► Most nonsmokers have a normal ECG, but smokers are about 50–50 normal and abnormal ECG.
- ▶ Don't look at smoke: bmi table since not significant.

# Simpson's paradox: the airlines example

	Alaska	Airlines	America	West
Airport	On time	Delayed	On time	Delayed
Los Angeles	497	62	694	117
Phoenix	221	12	4840	415
San Diego	212	20	383	65
San Francisco	503	102	320	129
Seattle	1841	305	201	61
Total	3274	501	6438	787

Use status as variable name for "on time/delayed".

- Alaska: 13.3% flights delayed (501/(3274 + 501)).
- America West: 10.9% (787/(6438 + 787)).
- ► America West more punctual, right?

### Arranging the data

➤ Can only have single thing in columns, so we have to construct column names like this:

airport	aa_ontime	aa_delayed	aw_ontime	aw_delayed
LosAngeles	497	62	694	117
Phoenix	221	12	4840	415
SanDiego	212	20	383	65
SanFrancis	co 503	102	320	129
Seattle	1841	305	201	61

Read in:

```
my_url <- "http://ritsokiguess.site/datafiles/airlines.txt"
airlines <- read_table(my_url)</pre>
```

# Tidying

➤ Some tidying gets us the right layout, with frequencies all in one column and the airline and delayed/on time status separated out. This uses one of the fancy versions of pivot\_longer:

# The data frame punctual

A tibble:  $20 \times 4$ airline status airport freq <chr> <chr>> <chr> <dbl>1 LosAngeles aa ontime 497 2 LosAngeles 62 delayed aa 3 LosAngeles ontime 694 aw 4 LosAngeles aw delayed 117 Phoenix ontime 221 ลล Phoenix delayed 12 ลล Phoenix ontime 4840 aw 8 Phoenix delayed 415 aw SanDiego ontime 212 aa SanDiego delayed 20 aa SanDiego ontime 383 aw 12 SanDiego delayed 65 aw SanFrancisco aa ontime 503 14 SanFrancisco aa delayed 102 15 SanFrancisco aw ontime 320

### Proportions delayed by airline

► Two-step process: get appropriate subtable:

```
xt <- xtabs(freq ~ airline + status, data = punctual)
xt</pre>
```

```
status
airline delayed ontime
aa 501 3274
aw 787 6438
```

▶ and then calculate appropriate proportions:

```
prop.table(xt, margin = 1)
```

```
status
airline delayed ontime
aa 0.1327152 0.8672848
aw 0.1089273 0.8910727
```

▶ More of Alaska Airlines' flights delayed (13.3% vs. 10.9%).

#### Proportion delayed by airport, for each airline

```
xt <- xtabs(freq ~ airline + status + airport, data = punctual)
xp <- prop.table(xt, margin = c(1, 3))
ftable(xp,
   row.vars = c("airport", "airline"),
   col.vars = "status"
)</pre>
```

		status	delayed	ontime
airport	${\tt airline}$			
LosAngeles	aa		0.11091234	0.88908766
	aw		0.14426634	0.85573366
Phoenix	aa		0.05150215	0.94849785
	aw		0.07897241	0.92102759
SanDiego	aa		0.08620690	0.91379310
	aw		0.14508929	0.85491071
SanFrancisco	aa		0.16859504	0.83140496
	aw		0.28730512	0.71269488
Seattle	aa		0.14212488	0.85787512
	aw		0.23282443	0.76717557

## Simpson's Paradox

Airport	Alaska	America West
Los Angeles	11.4	14.4
Phoenix	5.2	7.9
San Diego	8.6	14.5
San Francisco	16.9	28.7
Seattle	14.2	23.2
Total	13.3	10.9

- America West more punctual overall,
- but worse at *every single* airport!
- ► How is that possible?
- Log-linear analysis sheds some light.

#### Model 1 and output

```
punctual.1 <- glm(freq ~ airport * airline * status,</pre>
 data = punctual, family = "poisson"
drop1(punctual.1, test = "Chisq")
Single term deletions
Model:
freq ~ airport * airline * status
                       Df Deviance AIC LRT Pr(>Chi)
<none>
                            0.0000 183.44
airport:airline:status 4 3.2166 178.65 3.2166
                                                   0.5223
```

#### Remove 3-way interaction

```
punctual.2 <- update(punctual.1, ~ . - airport:airline:status)</pre>
drop1(punctual.2, test = "Chisq")
Single term deletions
Model:
freq ~ airport + airline + status + airport:airline + airport:status +
   airline:status
               Df Deviance ATC LRT Pr(>Chi)
                       3.2 178.7
<none>
airport:airline 4 6432.5 6599.9 6429.2 < 2.2e-16 ***
                4 240.1 407.5 236.9 < 2.2e-16 ***
airport:status
airline:status 1 45.5 218.9 42.2 8.038e-11 ***
               0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
Stop here.
```

# Understanding the significance

airline:status:

```
xt <- xtabs(freq ~ airline + status, data = punctual)
prop.table(xt, margin = 1)</pre>
```

```
status
airline delayed ontime
aa 0.1327152 0.8672848
aw 0.1089273 0.8910727
```

- More of Alaska Airlines' flights delayed overall.
- Saw this before.

# Understanding the significance (2)

airport:status:

```
xt <- xtabs(freq ~ airport + status, data = punctual)
prop.table(xt, margin = 1)</pre>
```

#### status

```
airport delayed ontime
LosAngeles 0.13065693 0.86934307
Phoenix 0.07780612 0.92219388
SanDiego 0.12500000 0.87500000
SanFrancisco 0.21916509 0.78083491
Seattle 0.15199336 0.84800664
```

- Flights into San Francisco (and maybe Seattle) are often late, and flights into Phoenix are usually on time.
- Considerable variation among airports.

# Understanding the significance (3)

airport:airline:

```
xt <- xtabs(freq ~ airport + airline, data = punctual)
prop.table(xt, margin = 2)</pre>
```

#### airline

aa	aw
0.14807947	0.11224913
0.06172185	0.72733564
0.06145695	0.06200692
0.16026490	0.06214533
0.56847682	0.03626298
	0.14807947 0.06172185 0.06145695 0.16026490

- ▶ What fraction of each airline's flights are to each airport.
- Most of Alaska Airlines' flights to Seattle and San Francisco.
- Most of America West's flights to Phoenix.

#### The resolution

- Most of America West's flights to Phoenix, where it is easy to be on time.
- Most of Alaska Airlines' flights to San Francisco and Seattle, where it is difficult to be on time.
- Overall comparison looks bad for Alaska because of this.
- ▶ But, comparing like with like, if you compare each airline's performance to the same airport, Alaska does better.
- Aggregating over the very different airports was a (big) mistake: that was the cause of the Simpson's paradox.
- Alaska Airlines is more punctual when you do the proper comparison.

## Ovarian cancer: a four-way table

- Retrospective study of ovarian cancer done in 1973.
- Information about 299 women operated on for ovarian cancer 10 years previously.
- Recorded:
  - stage of cancer (early or advanced)
  - type of operation (radical or limited)
  - X-ray treatment received (yes or no)
  - ▶ 10-year survival (yes or no)
- Survival looks like response (suggests logistic regression).
- Log-linear model finds any associations at all.

### The data

#### after tidying:

```
stage operation xray survival freq
early radical no no 10
early radical no yes 41
early radical yes no 17
early radical yes yes 64
early limited no no 1
early limited no yes 13
early limited yes no 3
early limited yes yes 9
advanced radical no no 38
advanced radical no yes 6
advanced radical yes no 64
advanced radical yes yes 11
advanced limited no no 3
advanced limited no yes 1
advanced limited yes no 13
advanced limited yes yes 5
```

# Reading in data

```
my_url <- "http://ritsokiguess.site/datafiles/cancer.txt"
cancer <- read_delim(my_url, " ")
cancer %>% slice(1:6)
# A tibble: 6 x 5
```

```
stage operation xray survival
                              freq
 <chr> <chr> <chr> <chr>
                             dbl>
                                10
1 early radical no
                     no
2 early radical no yes
                               41
3 early radical yes
                               17
                     no
4 early radical yes
                               64
                   yes
5 early limited
              no
                     nο
6 early limited
                                13
               no
                     yes
```

### Model 1

```
hopefully looking familiar by now:
```

```
cancer.1 <- glm(freq ~ stage * operation * xray * survival,
  data = cancer, family = "poisson"
)</pre>
```

### Output 1

### See what we can remove:

Non-significant interaction can come out.

#### Model 2

```
cancer.2 <- update(cancer.1, . ~ . - stage:operation:xray:survival)</pre>
drop1(cancer.2, test = "Chisq")
Single term deletions
Model:
freq ~ stage + operation + xray + survival + stage:operation +
    stage:xray + operation:xray + stage:survival + operation:survival +
   xray:survival + stage:operation:xray + stage:operation:survival +
   stage:xray:survival + operation:xray:survival
                        Df Deviance AIC LRT Pr(>Chi)
                            0.60266 96.732
<none>
stage:operation:xray 1 2.35759 96.487 1.75493 0.1853
stage:operation:survival 1 1.17730 95.307 0.57465 0.4484
stage:xray:survival
                    1 0.95577 95.085 0.35311 0.5524
operation:xray:survival 1 1.23378 95.363 0.63113 0.4269
```

Least significant term is stage:xray:survival: remove.

# Take out stage:xray:survival

```
cancer.3 <- update(cancer.2, . ~ . - stage:xray:survival)</pre>
drop1(cancer.3, test = "Chisq")
Single term deletions
Model:
freq ~ stage + operation + xray + survival + stage:operation +
    stage:xray + operation:xray + stage:survival + operation:survival +
    xray:survival + stage:operation:xray + stage:operation:survival +
    operation:xray:survival
                        Df Deviance AIC LRT Pr(>Chi)
                            0.95577 95.085
<none>
stage:operation:xray 1 3.08666 95.216 2.13089 0.1444
stage:operation:survival 1 1.56605 93.696 0.61029 0.4347
operation:xray:survival 1 1.55124 93.681 0.59547 0.4403
```

operation:xray:survival comes out next.

### Remove operation:xray:survival

```
cancer.4 <- update(cancer.3, . ~ . - operation:xray:survival)</pre>
drop1(cancer.4, test = "Chisq")
Single term deletions
Model:
freq ~ stage + operation + xray + survival + stage:operation +
   stage:xray + operation:xray + stage:survival + operation:survival +
   xray:survival + stage:operation:xray + stage:operation:survival
                        Df Deviance AIC LRT Pr(>Chi)
<none>
                             1.5512 93.681
xray:survival
                         1 1.6977 91.827 0.1464 0.70196
stage:operation:xray 1 6.8420 96.972 5.2907 0.02144 *
stage:operation:survival 1 1.9311 92.061 0.3799 0.53768
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

#### Comments

- stage:operation:xray has now become significant, so won't remove that.
- Shows value of removing terms one at a time.
- There are no higher-order interactions containing both xray and survival, so now we get to test (and remove) xray:survival.

### Remove xray:survival

```
cancer.5 <- update(cancer.4, . ~ . - xray:survival)</pre>
drop1(cancer.5, test = "Chisq")
Single term deletions
Model:
freq ~ stage + operation + xray + survival + stage:operation +
   stage:xray + operation:xray + stage:survival + operation:survival +
   stage:operation:xray + stage:operation:survival
                        Df Deviance AIC LRT Pr(>Chi)
<none>
                             1.6977 91.827
stage:operation:xray 1 6.9277 95.057 5.2300 0.0222 *
stage:operation:survival 1 2.0242 90.154 0.3265 0.5677
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

### Remove stage:operation:survival

```
cancer.6 <- update(cancer.5, . ~ . - stage:operation:survival)</pre>
drop1(cancer.6, test = "Chisq")
Single term deletions
Model:
freq ~ stage + operation + xray + survival + stage:operation +
   stage:xray + operation:xray + stage:survival + operation:survival +
   stage:operation:xray
                   Df Deviance AIC LRT Pr(>Chi)
<none>
                         2.024 90.154
stage:survival 1 135.198 221.327 133.173 <2e-16 ***
operation:survival 1 4.116 90.245 2.092 0.1481
stage:operation:xray 1 7.254 93.384 5.230 0.0222 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

### Last step?

Finally done!

```
Remove operation: survival.
cancer.7 <- update(cancer.6, . ~ . - operation:survival)</pre>
drop1(cancer.7, test = "Chisq")
Single term deletions
Model:
freq ~ stage + operation + xray + survival + stage:operation +
   stage:xray + operation:xray + stage:survival + stage:operation:xray
                    Df Deviance AIC LRT Pr(>Chi)
                         4.116 90.245
<none>
stage:survival 1 136.729 220.859 132.61 <2e-16 ***
stage:operation:xray 1 9.346 93.475 5.23 0.0222 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

#### Conclusions

- What matters is things associated with survival (survival is "response").
- Only significant such term is stage:survival:

```
xt <- xtabs(freq ~ stage + survival, data = cancer)
prop.table(xt, margin = 1)</pre>
```

#### survival

```
stage no yes
advanced 0.8368794 0.1631206
early 0.1962025 0.8037975
```

- Most people in early stage of cancer survived, and most people in advanced stage did not survive.
- ➤ This true regardless of type of operation or whether or not X-ray treatment was received. These things have no impact on survival.

### What about that other interaction?

```
xt <- xtabs(freq ~ operation + xray + stage, data = cancer)
ftable(prop.table(xt, margin = 3))</pre>
```

```
    stage
    advanced
    early

    operation
    xray

    limited
    no
    0.02836879
    0.08860759

    yes
    0.12765957
    0.07594937

    radical
    no
    0.31205674
    0.32278481

    yes
    0.53191489
    0.51265823
```

- Out of the people at each stage of cancer (since margin=3 and stage was listed 3rd).
- ▶ The association is between stage and xray only for those who had the limited operation.
- For those who had the radical operation, there was no association between stage and xray.
- This is of less interest than associations with survival.

# General procedure

- Start with "complete model" including all possible interactions.
- drop1 gives highest-order interaction(s) remaining, remove least non-significant.
- Repeat as necessary until everything significant.
- Look at subtables of significant interactions.
- Main effects not usually very interesting.
- Interactions with "response" usually of most interest: show association with response.