Supplement 2

Sites of pain

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1	In	nport and c	eck data	
da		read_rds('data read_rds('data	eaned/data-pain-sites.rds') eaned/data-demographics.rds	')
##		sites		
##	[1]	596 21		
na	mes(d	ata)		
## ## ## ## ##	[4] [7] [10] [13] [16]	"ID" "Shoulder" "Wrists.Hands" "Lower_back" "Thoracic_spine" "Hips" "Ankles.Feet"	"Arms" "Elb	es"
gl	impse	(data)		
## ## ## ##	Colu \$ ID \$ He \$ Th \$ Sh	ad roat oulder	hr> "No", "No", "No", "Yes" hr> "No", "No", "No", "No", hr> "No", "No", "No", "No",	5", "RPB76", "RPB77", "RPB , "Yes", "No", "No", "No", "No", "No", "No", " "No", "No", "No", "
##	\$ Ar \$ El \$ Wr		hr> "No", "No", "No", "No",	"No", "No", "No", "No", " "No", "No", "No", "No", " "No", "No", "No", "
## ##	\$ Lo	per_back wer_back	hr> "No", "No", "No", "No", hr> "No", "No"	, "No", "No", "No", "No", "No", "No", "No", "No", " "No", "No", "No", "No", "
## ##	\$ Ce \$ Th	domen rvical_spine oracic_spine mbosacral_spine	hr> "No", "No", "No", "No", hr> "No", "No"	", "No", "No", "Yes", "No" "No", "No", "No", " "No", "No", "No", " "No", "No", "No", "
##	\$ Gr \$ Hi	roin	hr> "No", "No", "No", "No",	"No", "No", "No", " "No", "No", "No", "Yes",

```
<chr> "No", "No", "No", "No", "Yes", "No", "Yes",...
## $ Legs
                       <chr> "No", "No", "No", "No", "Yes", "No", "Yes",...
## $ Knees
                       <chr> "No", "No", "No", "No", "Yes", "No", "Yes",...
## $ Ankles.Feet
## $ Buttocks
                       <chr> "No", "No", "No", "No", "No", "No", "No", "Yes", ...
                       <chr> "RP", "RP", "RP", "RP", "RP", "RP", "RP", "RP", "RP", "...
## $ Site
## Demographics
dim(demo)
## [1] 596
names (demo)
## [1] "ID"
                           "Site"
                                               "Sex"
## [4] "Age"
                           "Employment_status" "CD4_recent"
## [7] "ART_currently"
                           "Education"
glimpse(demo)
## Rows: 596
## Columns: 8
## $ ID
                       <chr> "RPB73", "RPB74", "RPB75", "RPB76", "RPB77", "RPB...
                       <chr> "RP", "RP", "RP", "RP", "RP", "RP", "RP", "RP", "RP", "...
## $ Site
## $ Sex
                       <chr> "Female", "Female", "Female", "Female", "Female", ...
                       <dbl> 36, 27, 39, 36, 31, 32, 28, 37, 31, 25, 31, 24, 3...
## $ Age
## $ Employment status <chr> "Other", "Unemployed", "Other", "Unemployed", "Un...
## $ CD4_recent <dbl> 391, 571, 591, 207, 126, 225, 543, 410, 74, 212, ...
                     <chr> "Yes", "Yes", "Yes", "Yes", "Yes", "Yes", "Yes", ...
## $ ART_currently
## $ Education
                      <chr> "Tertiary", "Secondary", "Secondary", "Primary", ...
```

2 Basic descriptive statistics

2.1 Pain sites

```
data %>%
   select(-ID, -Site) %>%
   mutate_if(is.character, factor) %>%
   skim()
```

Table 1: Data summary

Name	Piped data
Number of rows	596
Number of columns	19
Column type frequency: factor	19
Group variables	None

Variable type: factor

$skim_variable$	$n_missing$	$complete_rate$	n _unique	top_counts
Head	0	1	2	No: 401, Yes: 195
Throat	0	1	2	No: 577, Yes: 19
Shoulder	0	1	2	No: 549, Yes: 47
Arms	0	1	2	No: 571, Yes: 25
Elbows	0	1	2	No: 574, Yes: 22
Wrists.Hands	0	1	2	No: 561, Yes: 35
Chest	0	1	2	No: 479, Yes: 117
$Upper_back$	0	1	1	No: 596
Lower_back	0	1	2	No: 554, Yes: 42
Abdomen	0	1	2	No: 437, Yes: 159
Cervical_spine	0	1	2	No: 566, Yes: 30
Thoracic_spine	0	1	2	No: 522, Yes: 74
Lumbosacral_spine	0	1	2	No: 504, Yes: 92
Groin	0	1	2	No: 542, Yes: 54
Hips	0	1	2	No: 556, Yes: 40
Legs	0	1	2	No: 497, Yes: 99
Knees	0	1	2	No: 512, Yes: 84
Ankles.Feet	0	1	2	No: 412, Yes: 184
Buttocks	0	1	2	No: 577, Yes: 19

2.2 Demographics

```
demo %>%
    select(-ID, -Site) %>%
    mutate_if(is.character, factor) %>%
    skim()
```

Table 3: Data summary

Name Number of rows Number of columns	Piped data 596 6
Column type frequency: factor numeric	4 2
Group variables	None

Variable type: factor

skim_variable	$n_{missing}$	$complete_rate$	n_unique	top_counts
Sex	0	1.00	2	Fem: 481, Mal: 115
Employment_status	49	0.92	4	Une: 330, Ful: 131, Par: 52, Oth: 34
ART_currently	5	0.99	2	Yes: 460, No: 131
Education	37	0.94	3	Sec: 395, Pri: 99, Ter: 65

Variable type: numeric

skim_variable	n_missing	complete_rate	mean	sd	p0	p25	p50	p75	p100
Age	8	0.99	37.28	9.06	19	31	36	42	76
CD4_recent	99	0.83	320.71	238.92	1	155	261	432	1232

2.3 Add acute/chronic pain column to demographic data

Table 6: Data summary

Name	Piped data
Number of rows	596
Number of columns	1
Column type frequency:	
factor	1
Group variables	None

Variable type: factor

$skim_variable$	$n_missing$	$complete_rate$	n_unique	top_counts
Pain_def	0	1	2	Acu: 387, Chr: 209

2.4 Boostrap functions

```
# Proportion
prop_func <- function(d, i){
    dat <- d[i, ]
    dat_vec <- dat[[1]]
    dat_prop <- mean(dat_vec == 'Yes', na.rm = TRUE)
    dat_prop
}
# Median
median_func <- function(d, i){</pre>
```

```
dat <- d[i, ]
  dat_vec <- dat[[1]]
  dat_median <- median(dat_vec, na.rm = TRUE)
  dat_median
}</pre>
```

3 Proportion point estimates with 95% CIs

3.1 Process data

```
# Set seed
set.seed(2020)
# Remove ID and upper_back (only one outcome -- no pain) columns
prop <- data[, !(names(data) %in% c('ID', 'Upper_back'))]</pre>
# Bootstrap CIs
prop_boot <- prop %>%
    # Remove Site column
   select(-Site) %>%
    # Pivot to long format
   pivot_longer(cols = everything(),
                 names_to = 'body_site',
                 values_to = 'pain_present') %>%
    # Add body regions
   mutate(region = case_when(
        body_site == 'Chest' |
            body_site == 'Head' |
            body_site == 'Throat' |
            body_site == 'Shoulder' ~ 'Head and upper torso',
        body_site == 'Lower_back' |
            body_site == 'Abdomen' |
            body_site == 'Hips' |
            body_site == 'Buttocks' |
            body_site == 'Groin' ~ 'Lower torso',
        body_site == 'Legs' |
            body_site == 'Knees' |
            body_site == 'Ankles.Feet' ~ 'Lower limbs',
        body_site == 'Arms' |
            body_site == 'Elbows' |
            body_site == 'Wrists.Hands' ~ 'Upper limbs',
        body_site == 'Cervical_spine' |
            body_site == 'Thoracic_spine' |
            body_site == 'Lumbosacral_spine' ~ 'Spinal column',
        TRUE ~ 'other'
    # Nest by body region and body site
   group_by(region, body_site) %>%
   nest() %>%
    # Boostrap data
   mutate(boot = map(.x = data,
```

```
~ boot(data = .x,
                             statistic = prop_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
   mutate(ci = map(.x = boot,
                   ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                           ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
    select(-data, -boot, -ci) %>%
    # Unnest
   unnest(cols = c(point_est, lower_ci, upper_ci))
# Re-nest by body region and generate figures and tables
prop_boot2 <- prop_boot %>%
   group_by(region) %>%
   nest() %>%
    # Fix site labels
   mutate(data = map(.x = data,
                      ~ .x %>%
                          mutate(body_site = str_replace_all(body_site,
                                                         pattern = '_',
                                                         replacement = ' '),
                                 body_site = str_replace_all(body_site,
                                                         pattern = '\\.',
                                                         replacement = ' & '),
                                 body_site = str_replace_all(body_site,
                                                         pattern = 'Lower back',
                                                         replacement = 'Flank')))) %>%
    # Re-order sites by point_est
   mutate(data = map(.x = data,
                      ~ .x %>%
                          mutate(body_site = fct_reorder(body_site,
                                                    point_est)))) %>%
    # Plot data
   mutate(plots = map2(.x = data,
                        y = region,
                       ~ .x %>%
                           ggplot(data = .) +
                           aes(x = body_site,
                               y = point_est,
                               ymin = lower_ci,
                               ymax = upper_ci) +
                           geom_hline(yintercept = 0.1,
                                      linetype = 2) +
```

```
geom_pointrange(size = 1) +
                       coord_flip() +
                        labs(title = .y,
                             subtitle = '(Point estimate with 95%CI)',
                             y = 'Proportion with pain') +
                        scale_y_continuous(limits = c(0, 0.4)) +
                        theme_minimal(base_size = 18) +
                        theme(plot.title = element_text(size = 18),
                              plot.subtitle = element_text(size = 12),
                              axis.title.y = element_blank(),
                              panel.grid = element_blank(),
                              axis.text = element_text(colour = '#000000'),
                              axis.line = element_line(size = 0.5),
                              axis.ticks = element_line(size = 0.5)))) %>%
# Tabulate data
mutate(tables = map2(.x = data,
                     .y = region,
                     ~ .x %>%
                         kable(caption = .y,
                               digits = 2)))
```

3.2 Tabulated proportions (with 95% CIs), by body region

```
walk(prop_boot2$tables, ~ print(.x))
```

Table 8: Head and upper torso

body_site	point_est	lower_ci	upper_ci
Head	0.33	0.29	0.37
Throat	0.03	0.02	0.05
Shoulder	0.08	0.06	0.10
Chest	0.20	0.17	0.23

Table 9: Upper limbs

body_site	point_est	lower_ci	upper_ci
Arms	0.04	0.03	0.06
Elbows	0.04	0.02	0.05
Wrists & Hands	0.06	0.04	0.08

Table 10: Lower torso

body_site	point_est	lower_ci	upper_ci
Flank	0.07	0.05	0.09
Abdomen	0.27	0.23	0.30
Groin	0.09	0.07	0.11
Hips	0.07	0.05	0.09
Buttocks	0.03	0.02	0.05

Table 11: Spinal column

body_site	point_est	lower_ci	upper_ci
Cervical spine	0.05	0.03	0.07
Thoracic spine	0.12	0.10	0.15
Lumbosacral spine	0.15	0.13	0.18

Table 12: Lower limbs

body_site	point_est	lower_ci	upper_ci
Legs	0.17	0.14	0.20
Knees	0.14	0.11	0.17
Ankles & Feet	0.31	0.27	0.35

3.3 Plotted proportions (with 95% CIs), by body region

```
upper <- prop_boot2$plots[[1]] +
    theme(axis.title.x = element_blank())

arm <- prop_boot2$plots[[2]] +
    theme(axis.title.x = element_blank())

lower <- prop_boot2$plots[[3]] +
    theme(axis.title.x = element_blank())

spine <- prop_boot2$plots[[4]]

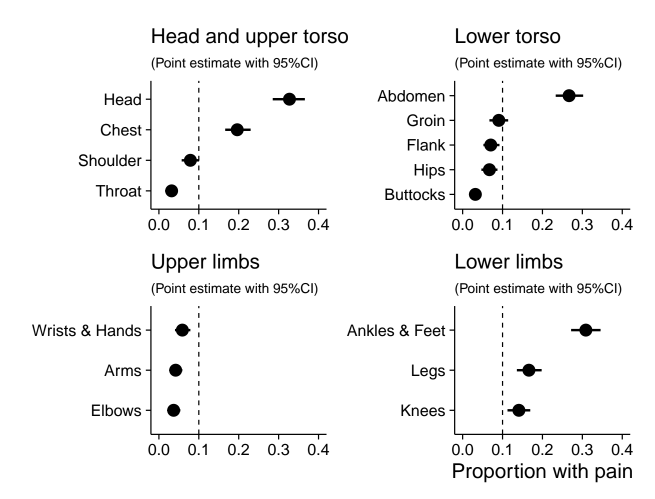
leg <- prop_boot2$plots[[5]]

# Patchwork

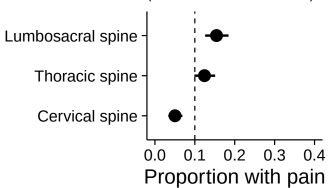
plot_prop <- upper + lower + arm + leg + spine +
    plot_layout(ncol = 2)

# Output

plot_prop</pre>
```



Spinal column



4 By sex

4.1 Process data

```
# Set seed
set.seed(2020)
# Select sex data
sex <- demo[, c('ID', 'Sex')]</pre>
# Join to boot_data & remove ID, site, and upper_back (only one outcome -- no pain)
sex <- left_join(data, sex) %>%
    select(-ID, -Site, -Upper_back)
# Bootstrap CIs
sex_boot <- sex %>%
    # Pivot to long format
   pivot_longer(cols = -Sex,
                 names_to = 'body_site',
                 values_to = 'pain_present') %>%
    # Add body regions
   mutate(region = case_when(
        body_site == 'Chest' |
            body_site == 'Head' |
            body_site == 'Throat' |
            body_site == 'Shoulder' ~ 'Head and upper torso',
        body_site == 'Lower_back'
            body_site == 'Abdomen' |
            body_site == 'Hips' |
            body_site == 'Buttocks' |
            body_site == 'Groin' ~ 'Lower torso',
        body_site == 'Legs' |
            body site == 'Knees' |
            body_site == 'Ankles.Feet' ~ 'Lower limbs',
        body_site == 'Arms' |
            body_site == 'Elbows' |
            body_site == 'Wrists.Hands' ~ 'Upper limbs',
        body_site == 'Cervical_spine' |
            body_site == 'Thoracic_spine' |
            body_site == 'Lumbosacral_spine' ~ 'Spinal column',
        TRUE ~ 'other'
   )) %>%
    # Nest by body region and body site
    group_by(Sex, region, body_site) %>%
   nest() %>%
    # Boostrap data
   mutate(boot = map(.x = data,
                      ~ boot(data = .x,
                             statistic = prop_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
   # Get CI
```

```
mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
    mutate(point_est = map(.x = ci,
                          ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
   select(-data, -boot, -ci) %>%
    # Unnest
   unnest(cols = c(point_est, lower_ci, upper_ci))
# Re-nest by body region and generate figures and tables
sex_boot2 <- sex_boot %>%
   group_by(region) %>%
   nest() %>%
    # Fix site labels
   mutate(data = map(.x = data,
                      ~ .x %>%
                          mutate(body_site = str_replace_all(body_site,
                                                        pattern = '_',
                                                        replacement = ' '),
                                 body_site = str_replace_all(body_site,
                                                        pattern = '\\.',
                                                        replacement = ' & ')))) %>%
    # Re-order sites by point_est
   mutate(data = map(.x = data,
                      ~ .x %>%
                          mutate(body_site = fct_reorder(body_site,
                                                    point_est)))) %>%
    # Plot data
   mutate(plots = map2(.x = data,
                        .y = region,
                       ~ .x %>%
                           ggplot(data = .) +
                           aes(x = body_site,
                               y = point_est,
                               ymin = lower_ci,
                               ymax = upper_ci,
                               fill = Sex) +
                           geom_linerange(position = position_dodge2(width = 0.6),
                                          size = 1,
                                          colour = '#000000') +
                           geom_point(shape = 21,
                                      colour = '#000000',
                                      position = position_dodge2(width = 0.6),
                                      size = 6,
                                      stroke = 1) +
                           coord_flip() +
                           labs(title = .y,
                                subtitle = '(Point estimate with 95%CI)',
```

```
y = 'Proportion with pain') +
                       scale_y_continuous(limits = c(0, 1)) +
                       scale_fill_manual(values = c('#000000', '#FFFFFF')) +
                       theme_minimal(base_size = 18) +
                       theme(plot.title = element_text(size = 18),
                             plot.subtitle = element_text(size = 12),
                             legend.title = element_blank(),
                             legend.position = 'top',
                             axis.title.y = element_blank(),
                             panel.grid = element_blank(),
                             axis.text = element_text(colour = '#000000'),
                             axis.line = element_line(size = 0.5),
                             axis.ticks = element_line(size = 0.5)))) %>%
# Tabulate data
mutate(tables = map2(.x = data,
                     .y = region,
                     ~ .x %>%
                         kable(caption = .y,
                               digits = 2)))
```

4.2 Tabulated proportions (with 95% CIs), by age and body region

```
walk(sex_boot2$tables, ~ print(.x))
```

Table 13: Head and upper torso

Sex	body_site	point_est	lower_ci	upper_ci
Female	Head	0.38	0.33	0.42
Female	Throat	0.03	0.01	0.04
Female	Shoulder	0.07	0.05	0.10
Female	Chest	0.20	0.16	0.23
Male	Head	0.12	0.06	0.18
Male	Throat	0.04	0.01	0.09
Male	Shoulder	0.11	0.06	0.18
Male	Chest	0.20	0.13	0.27

Table 14: Upper limbs

Sex	body_site	point_est	lower_ci	upper_ci
Female	Arms	0.04	0.03	0.06
Female	Elbows	0.03	0.02	0.05
Female	Wrists & Hands	0.06	0.04	0.09
Male	Arms	0.03	0.01	0.07
Male	Elbows	0.05	0.02	0.10
Male	Wrists & Hands	0.03	0.01	0.08

Table 15: Lower torso

Sex	body_site	point_est	lower_ci	upper_ci
Female	Lower back	0.07	0.05	0.10
Female	Abdomen	0.28	0.24	0.32
Female	Groin	0.10	0.07	0.12
Female	Hips	0.06	0.04	0.09
Female	Buttocks	0.03	0.01	0.04
Male	Lower back	0.05	0.02	0.10
Male	Abdomen	0.19	0.12	0.27
Male	Groin	0.07	0.03	0.12
Male	Hips	0.10	0.05	0.15
Male	Buttocks	0.04	0.01	0.09

Table 16: Spinal column

Sex	body_site	point_est	lower_ci	upper_ci
Female	Cervical spine	0.06	0.04	0.08
Female	Thoracic spine	0.14	0.11	0.17
Female	Lumbosacral spine	0.16	0.12	0.19
Male	Cervical spine	0.03	0.00	0.06
Male	Thoracic spine	0.07	0.03	0.12
Male	Lumbosacral spine	0.14	0.08	0.20

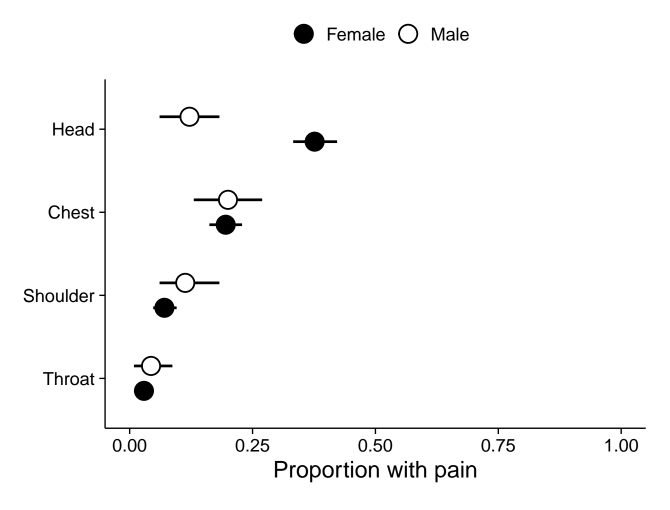
Table 17: Lower limbs

Sex	body_site	point_est	lower_ci	upper_ci
Female	Legs	0.15	0.12	0.18
Female	Knees	0.13	0.10	0.16
Female	Ankles & Feet	0.28	0.25	0.32
Male	Legs	0.23	0.16	0.31
Male	Knees	0.19	0.12	0.26
Male	Ankles & Feet	0.42	0.33	0.50

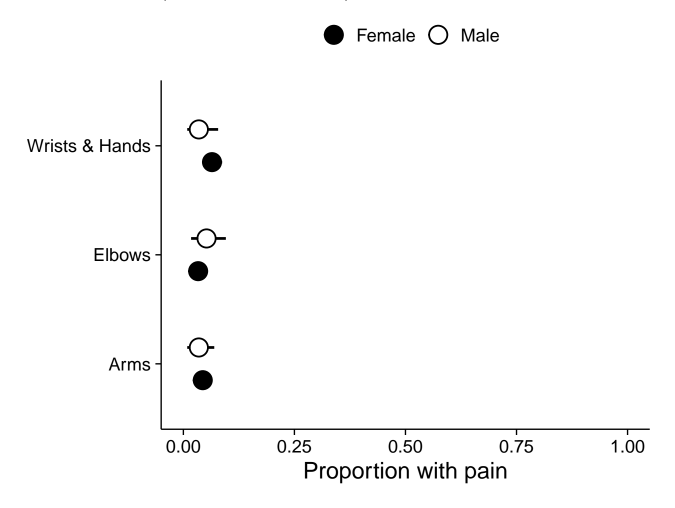
4.3 Plotted proportions (with 95% CIs), by age and body region

walk(sex_boot2\$plots, ~ print(.x))

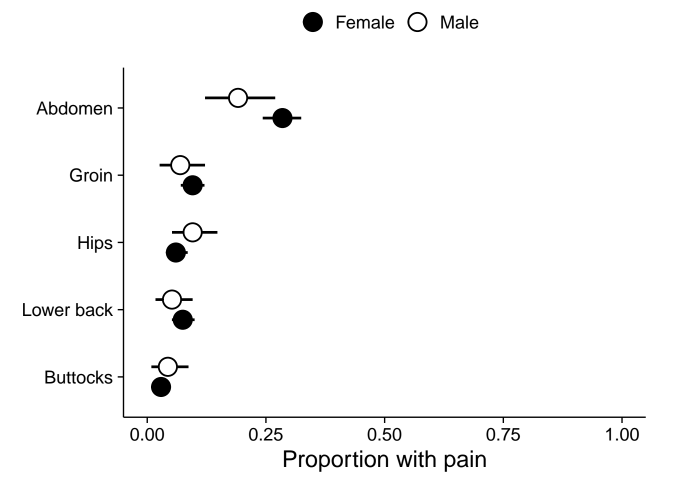
Head and upper torso



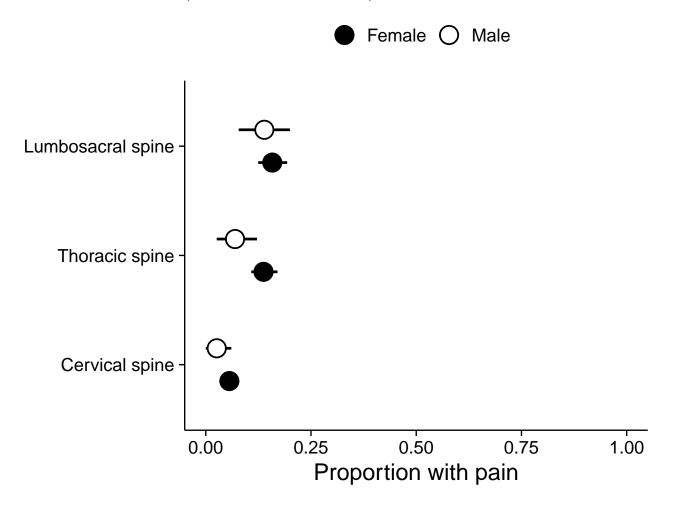
Upper limbs



Lower torso



Spinal column



Lower limbs

(Point estimate with 95%CI)



5 By age

For ease of tabulation and plotting in this section of the supplement, I divided age into seven age categories. However for data analysis (see: Logistic regression, section 7), I analysed age as a continuous variable.

5.1 Process data

```
# Set seed
set.seed(2020)

# Select age data
age <- demo[, c('ID', 'Age')]

# Join to boot_data & remove ID, study site, and upper_back
# (only one outcome -- no pain)
age <- left_join(data, age) %>%
    select(-ID, -Site, -Upper_back)
```

```
# Get complete cases
age <- age[complete.cases(age), ]</pre>
# Pivot and add age group categories (10 year periods)
age_boot <- age %>%
    # Pivot to long format
    pivot_longer(cols = -Age,
                 names_to = 'body_site',
                 values_to = 'pain_present') %>%
    # Add age categories
    mutate(age_group = case_when(
        Age < 28 \sim '18-27',
        Age >= 28 & Age < 38 ~ '28-37',
        Age >= 38 & Age < 48 ~ '38-47',
        Age >= 48 \& Age < 58 ~ '48-57',
        Age >= 58 & Age < 68 ~ '58-67',
        Age >= 68 & Age < 78 ~ '68-77',
        Age >= 78 & Age < 88 ~ '78-87'
    ))
# Print count per age group
age_boot %>%
    group_by(body_site, age_group) %>%
    summarise(count = n()) %>%
    filter(body_site == 'Abdomen') %>%
    ungroup() %>%
    select(-body_site) %>%
    kable(caption = 'Participant count per age group')
```

Table 18: Participant count per age group

age_group	count
18-27	65
28-37	283
38-47	160
48-57	58
58-67	20
68-77	2

```
R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
   mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                          ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
    select(-data, -boot, -ci) %>%
    # Unnest
    unnest(cols = c(point_est, lower_ci, upper_ci)) %>%
   ungroup()
# Re-nest by body region and generate figures and tables
age_boot2 <- age_boot2 %>%
    # Fix site labels
   mutate(body_site = str_replace_all(body_site,
                                  pattern = '_',
                                  replacement = ' '),
           body_site = str_replace_all(body_site,
                                  pattern = '\\.',
                                  replacement = ' & ')) %>%
    # Group and nest
   group_by(body_site) %>%
   nest() %>%
    # Arrange age groups
    # Plot data
   mutate(plots = map2(.x = data,
                        .y = body_site,
                       ~ .x %>%
                           ggplot(data = .) +
                           aes(x = age\_group,
                               y = point_est,
                               ymin = lower_ci,
                               ymax = upper_ci) +
                           geom_linerange(size = 1,
                                          colour = '#000000') +
                           geom_point(colour = '#000000',
                                      size = 6) +
                           labs(title = .y,
                                subtitle = '(Point estimate with 95%CI)',
                                caption = 'Age group 68-77 years removed because n = 2',
                                x = 'Age group (Years)',
                                y = 'Proportion with pain') +
                           scale_y_continuous(limits = c(0, 1)) +
                           coord_flip() +
```

5.2 Tabulated proportions (with 95% CIs), by age group and body site

```
walk(age_boot2$tables, ~ print(.x))
```

Table 19: Head (Age group 68-77 years removed because n=2)

age_group	point_est	lower_ci	upper_ci
18-27	0.45	0.32	0.57
28-37	0.34	0.28	0.39
38-47	0.31	0.24	0.39
48-57	0.17	0.09	0.28
58-67	0.35	0.15	0.55

Table 20: Throat (Age group 68-77 years removed because n = 2)

age_group	point_est	lower_ci	upper_ci
18-27	0.05	0.00	0.11
28-37	0.03	0.01	0.05
38-47	0.03	0.01	0.06
48-57	0.05	0.00	0.12
58-67	0.00	0.00	0.00

Table 21: Shoulder (Age group 68-77 years removed because n=2)

age_group	$point_est$	$lower_ci$	upper_ci
18-27	0.12	0.05	0.22
28-37	0.07	0.04	0.10
38-47	0.09	0.05	0.14
48-57	0.07	0.02	0.14
58-67	0.05	0.00	0.15

Table 22: Arms (Age group 68-77 years removed because n=2)

age_group	point_est	lower_ci	upper_ci
18-27	0.02	0.00	0.05
28-37	0.04	0.02	0.07
38-47	0.06	0.02	0.09
48-57	0.05	0.00	0.12
58-67	0.00	0.00	0.00

Table 23: Elbows (Age group 68-77 years removed because n=2)

age_group	point_est	lower_ci	upper_ci
18-27	0.00	0.00	0.00
28-37	0.03	0.01	0.06
38-47	0.04	0.01	0.07
48-57	0.05	0.00	0.10
58-67	0.15	0.00	0.30

Table 24: Wrists & Hands (Age group 68-77 years removed because $n\,=\,2)$

age_group	point_est	lower_ci	upper_ci
18-27	0.02	0.00	0.05
28-37	0.07	0.04	0.10
38-47	0.04	0.01	0.07
48-57	0.07	0.02	0.14
58-67	0.15	0.00	0.30

Table 25: Chest (Age group 68-77 years removed because n=2)

age_group	point_est	lower_ci	upper_ci
18-27	0.17	0.08	0.26
28-37	0.20	0.15	0.25
38-47	0.20	0.14	0.26
48-57	0.19	0.09	0.29
58-67	0.20	0.05	0.40

Table 26: Lower back (Age group 68-77 years removed because n = 2)

age_group	point_est	lower_ci	upper_ci
18-27	0.09	0.03	0.17
28-37	0.07	0.04	0.10
38-47	0.05	0.02	0.09
48-57	0.10	0.03	0.19
58-67	0.10	0.00	0.25

Table 27: Abdomen (Age group 68-77 years removed because n=2)

age_group	point_est	lower_ci	upper_ci
18-27	0.23	0.14	0.34
28-37	0.31	0.26	0.37
38-47	0.29	0.22	0.37
48-57	0.10	0.03	0.19
58-67	0.05	0.00	0.15

Table 28: Cervical spine (Age group 68-77 years removed because n=2)

age_group	$point_est$	lower_ci	upper_ci
18-27	0.12	0.05	0.20
28-37	0.05	0.02	0.08
38-47	0.04	0.02	0.08
48-57	0.00	0.00	0.00
58-67	0.00	0.00	0.00

Table 29: Thoracic spine (Age group 68-77 years removed because n=2)

age_group	point_est	lower_ci	upper_ci
18-27	0.18	0.09	0.28
28-37	0.14	0.11	0.19
38-47	0.07	0.03	0.11
48-57	0.09	0.02	0.17
58-67	0.25	0.05	0.45

Table 30: Lumbosacral spine (Age group 68-77 years removed because $\mathbf{n}=2$)

age_group	point_est	lower_ci	upper_ci
18-27	0.18	0.09	0.29
28-37	0.15	0.11	0.19
38-47	0.12	0.07	0.17
48-57	0.16	0.07	0.26
58-67	0.40	0.20	0.60

Table 31: Groin (Age group 68-77 years removed because n=2)

age_group	point_est	lower_ci	upper_ci
18-27	0.09	0.03	0.17
28-37	0.10	0.07	0.14
38-47	0.09	0.05	0.13
48-57	0.05	0.00	0.12

age_group	point_est	lower_ci	upper_ci
58-67	0.05	0.00	0.15

Table 32: Hips (Age group 68-77 years removed because n=2)

age_group	point_est	lower_ci	upper_ci
18-27	0.03	0.00	0.08
28-37	0.05	0.02	0.07
38-47	0.09	0.05	0.14
48-57	0.14	0.05	0.24
58-67	0.10	0.00	0.25

Table 33: Legs (Age group 68-77 years removed because n=2)

age_group	$point_est$	lower_ci	upper_ci
18-27	0.12	0.05	0.22
28-37	0.13	0.09	0.17
38-47	0.21	0.15	0.28
48-57	0.29	0.17	0.41
58-67	0.20	0.05	0.40

Table 34: Knees (Age group 68-77 years removed because n=2)

age_group	point_est	lower_ci	upper_ci
18-27	0.11	0.05	0.18
28-37	0.10	0.07	0.14
38-47	0.17	0.11	0.23
48-57	0.22	0.12	0.34
58-67	0.30	0.10	0.50

Table 35: Ankles & Feet (Age group 68-77 years removed because $n\,=\,2)$

age_group	point_est	lower_ci	upper_ci
18-27	0.22	0.12	0.32
28-37	0.24	0.19	0.29
38-47	0.37	0.29	0.44
48-57	0.52	0.40	0.64
58-67	0.55	0.35	0.75

Table 36: Buttocks (Age group 68-77 years removed because n=2)

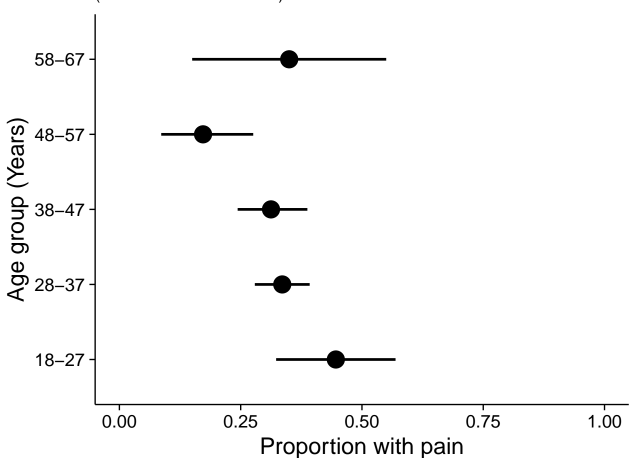
age_group	point_est	lower_ci	upper_ci
18-27	0.03	0.00	0.08

age_group	point_est	lower_ci	upper_ci
28-37	0.04	0.02	0.06
38-47	0.02	0.01	0.05
48-57	0.02	0.00	0.05
58-67	0.05	0.00	0.15

5.3 Plotted proportions (with 95% CIs), by age group and body site

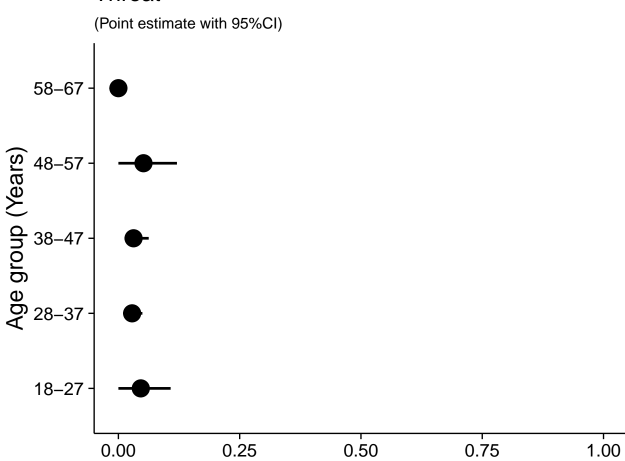
walk(age_boot2\$plots, ~ print(.x))

Head



Age group 68-77 years removed because n = 2



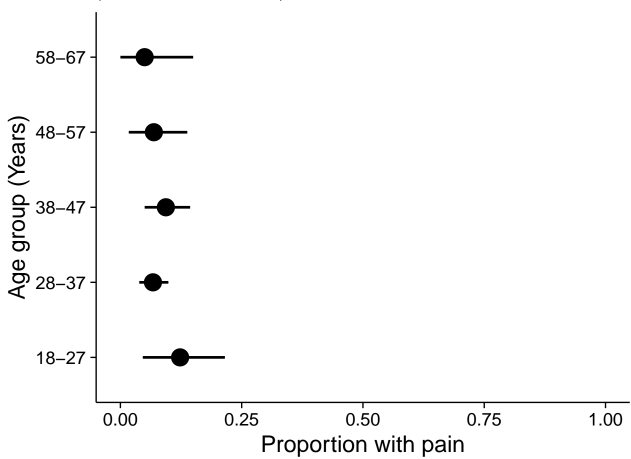


Proportion with pain

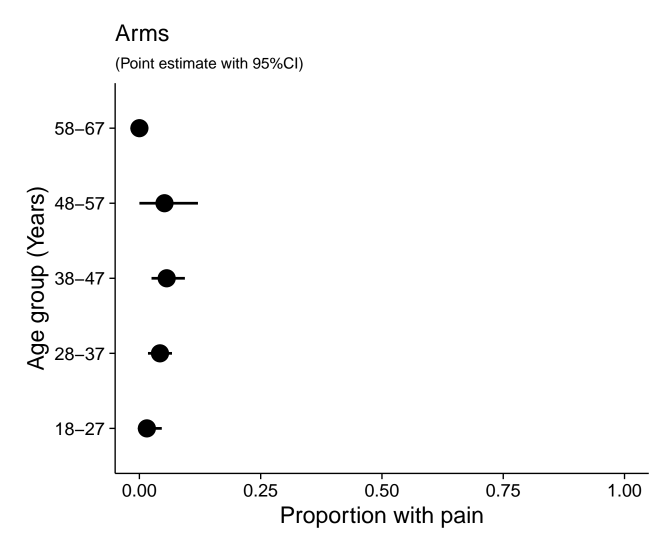
Age group 68-77 years removed because n = 2

Shoulder

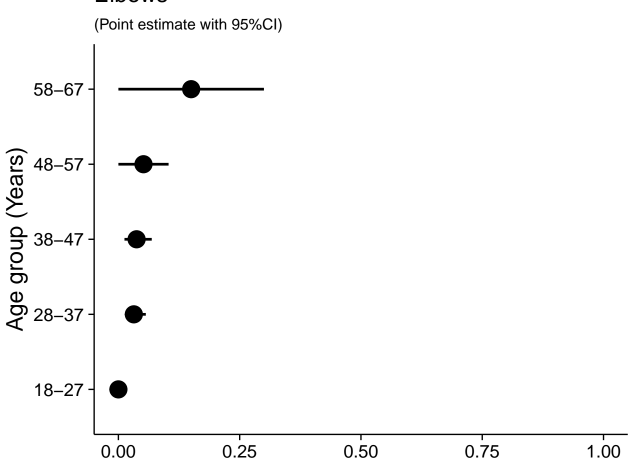




Age group 68-77 years removed because n = 2



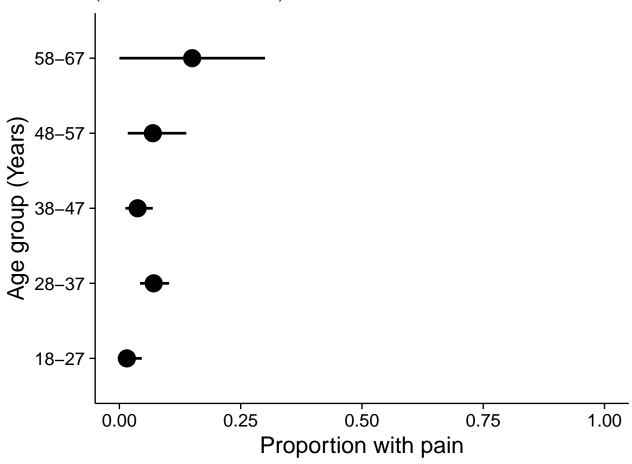
Elbows



Proportion with pain

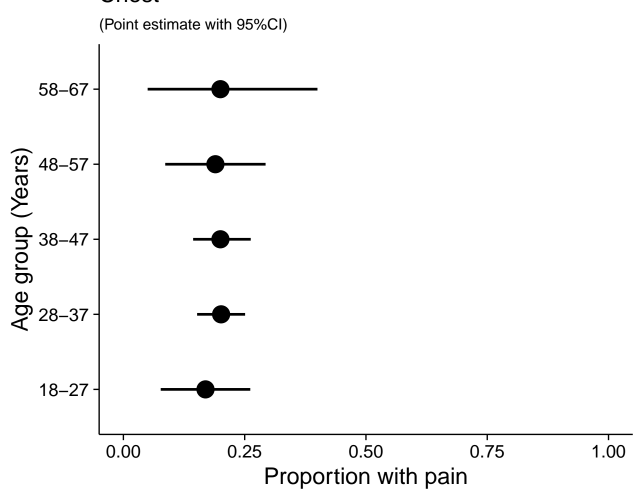
Age group 68-77 years removed because n = 2

Wrists & Hands



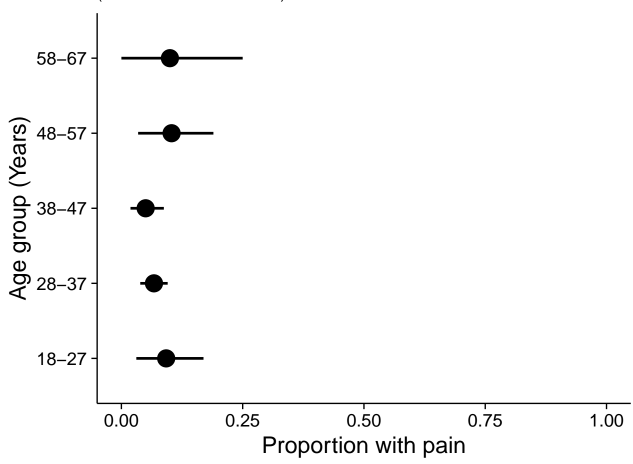
Age group 68-77 years removed because n = 2

Chest



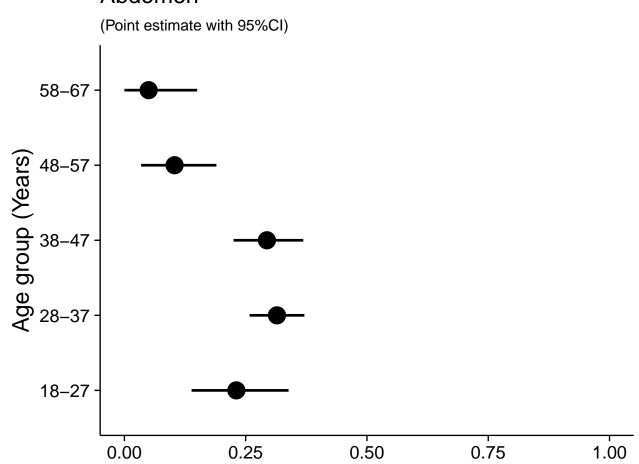
Age group 68-77 years removed because n = 2

Lower back



Age group 68-77 years removed because n = 2

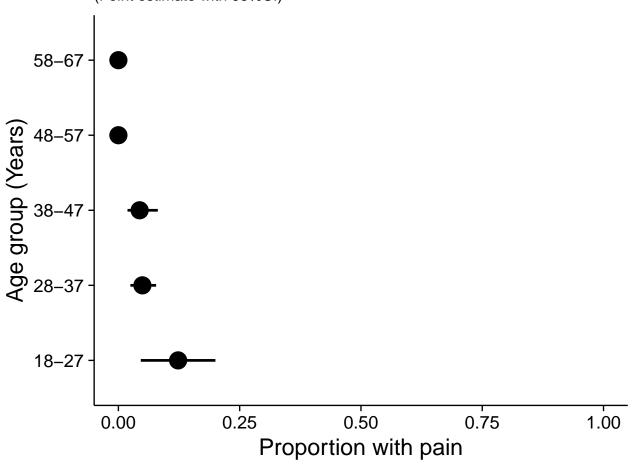
Abdomen



Proportion with pain

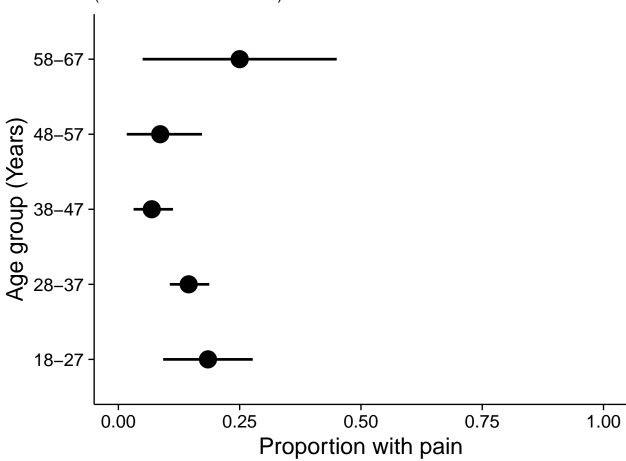
Age group 68-77 years removed because n = 2

Cervical spine



Age group 68-77 years removed because n = 2

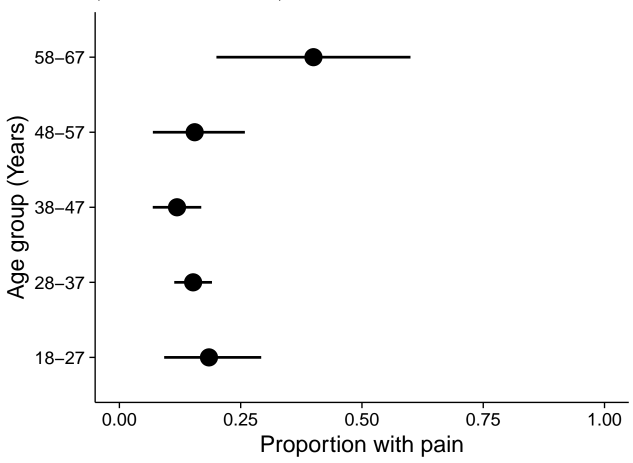
Thoracic spine



Age group 68-77 years removed because n = 2

Lumbosacral spine

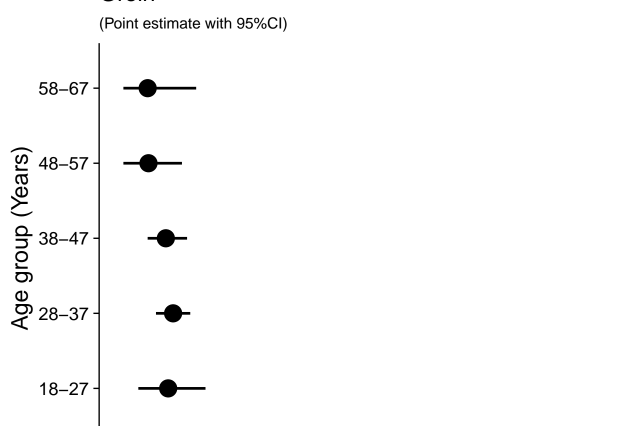
(Point estimate with 95%CI)



Age group 68-77 years removed because n = 2



0.00



0.50

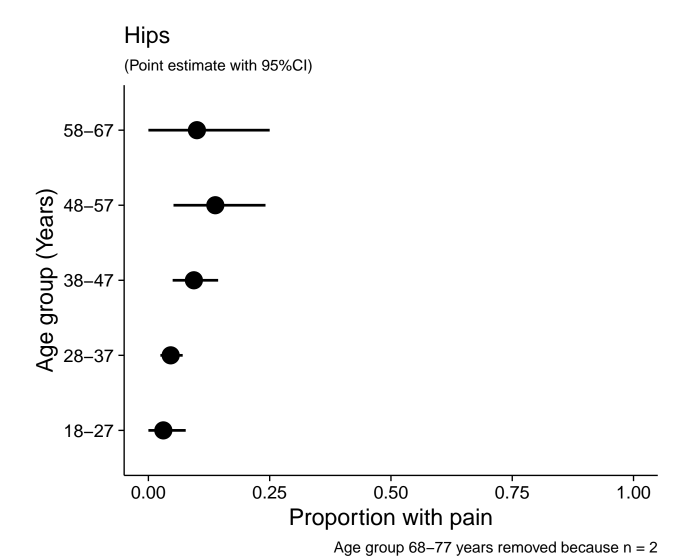
Proportion with pain

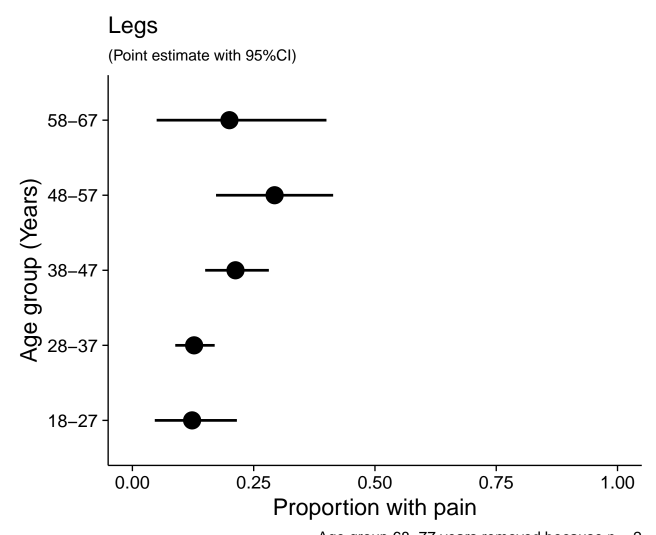
0.25

Age group 68-77 years removed because n = 2

0.75

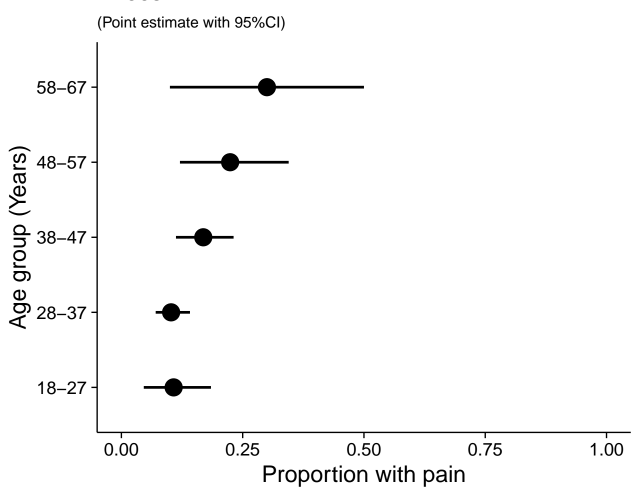
1.00





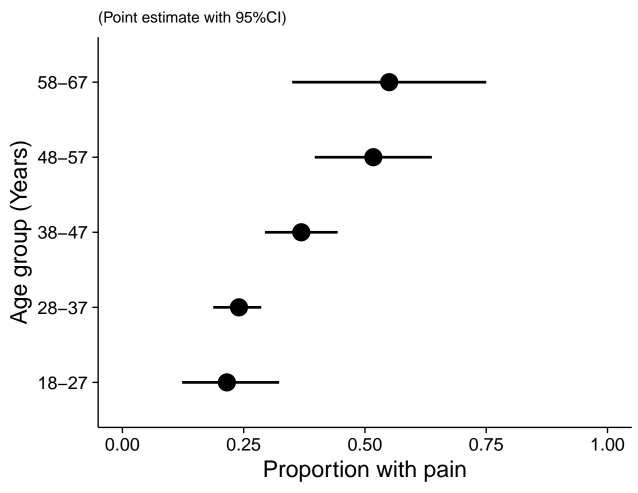
Age group 68-77 years removed because n=2

Knees



Age group 68-77 years removed because n = 2

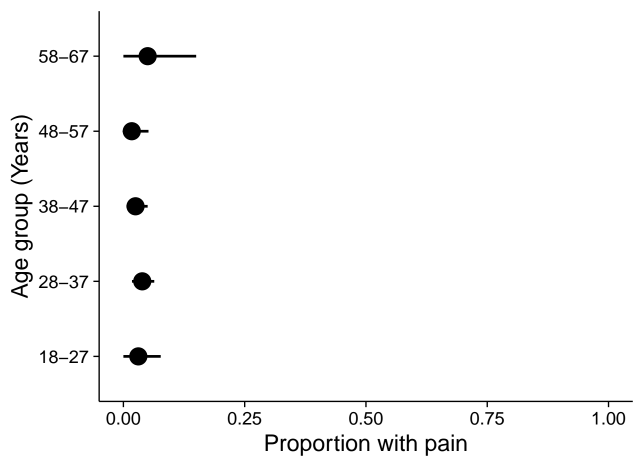
Ankles & Feet



Age group 68-77 years removed because n = 2

Buttocks





Age group 68-77 years removed because n = 2

6 By most recent CD4 T-cell count

For ease of tabulation and plotting in this section of the supplement, I divided the most recent CD4 T-cell count into six categories. However for data analysis (see: Logistic regression, section 7), I analysed CD4 T-cell count as a continuous variable.

6.1 Process data

```
# Set seed
set.seed(2020)

# Select CD4 recent data
cd4 <- demo[, c('ID', 'CD4_recent')]

# Join to boot_data & remove ID, study site, and upper_back
# (only one outcome -- no pain)
cd4 <- left_join(data, cd4) %>%
```

```
select(-ID, -Site, -Upper_back)
# Get complete cases
cd4 <- cd4[complete.cases(cd4), ]</pre>
# Pivot and add CD4 recent group categories (counts of 100)
cd4_boot <- cd4 %>%
    # Pivot to long format
   pivot_longer(cols = -CD4_recent,
                 names_to = 'body_site',
                 values_to = 'pain_present') %>%
    # Add CD4 recent categories
   mutate(cd4_group = case_when(
        CD4_recent < 100 ~ '0-99',
        CD4_recent >= 100 & CD4_recent < 200 ~ '100-199',
        CD4_recent >= 200 & CD4_recent < 300 ~ '200-299',
        CD4_recent >= 300 & CD4_recent < 400 ~ '300-399',
        CD4_recent >= 400 & CD4_recent < 500 ~ '400-499',
        CD4_recent >= 500 ~ '500+'
   ))
# Print count per CD4 recent group
cd4_boot %>%
   group_by(body_site, cd4_group) %>%
    summarise(count = n()) %>%
   filter(body_site == 'Abdomen') %>%
   ungroup() %>%
   select(-body_site) %>%
   kable(caption = 'Participant count per CD4 group')
```

Table 37: Participant count per CD4 group

cd4_group	count
0-99	68
100-199	106
200-299	108
300-399	77
400-499	46
500+	92

```
stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
   mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                         ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
    select(-data, -boot, -ci) %>%
    # Unnest
   unnest(cols = c(point_est, lower_ci, upper_ci)) %>%
    ungroup()
# Re-nest by body region and generate figures and tables
cd4_boot2 <- cd4_boot2 %>%
    # Fix site labels
   mutate(body_site = str_replace_all(body_site,
                                  pattern = '_',
                                  replacement = ' '),
           body_site = str_replace_all(body_site,
                                  pattern = '\\.',
                                  replacement = ' & ')) %>%
    # Group and nest
   group_by(body_site) %>%
   nest() %>%
    # Arrange CD4 recent groups
    # Plot data
   mutate(plots = map2(.x = data,
                        .y = body_site,
                       ~ .x %>%
                           ggplot(data = .) +
                           aes(x = cd4\_group,
                               y = point_est,
                               ymin = lower_ci,
                               ymax = upper_ci) +
                           geom_linerange(size = 1,
                                          colour = '#000000') +
                           geom_point(colour = '#000000',
                                      size = 6) +
                           labs(title = .y,
                                subtitle = '(Point estimate with 95%CI)',
                                x = expression('CD4 group (cells.mm'^-3*')'),
                                y = 'Proportion with pain') +
                           scale_y_continuous(limits = c(0, 1)) +
                           coord_flip() +
                           theme_minimal(base_size = 18) +
                           theme(plot.title = element_text(size = 18),
```

6.2 Tabulated proportions (with 95% CIs), by CD4 (recent) group and body site

```
walk(cd4_boot2$tables, ~ print(.x))
```

Table 38: Head

cd4_group	point_est	lower_ci	upper_ci
0-99	0.26	0.16	0.37
100-199	0.40	0.30	0.49
200-299	0.32	0.24	0.42
300-399	0.36	0.25	0.48
400-499	0.30	0.17	0.43
500+	0.25	0.16	0.34

Table 39: Throat

cd4_group	point_est	lower_ci	upper_ci
0-99	0.07	0.01	0.15
100-199	0.02	0.00	0.05
200-299	0.03	0.00	0.06
300-399	0.01	0.00	0.04
400-499	0.04	0.00	0.11
500+	0.00	0.00	0.00

Table 40: Shoulder

cd4_group	point_est	lower_ci	upper_ci
0-99	0.07	0.01	0.15
100-199	0.10	0.05	0.17
200-299	0.06	0.02	0.11
300-399	0.05	0.01	0.10
400-499	0.15	0.04	0.26
500+	0.08	0.02	0.13

Table 41: Arms

cd4_group	point_est	lower_ci	upper_ci
0-99	0.06	0.01	0.12
100-199	0.08	0.03	0.14
200-299	0.00	0.00	0.00
300-399	0.04	0.00	0.09
400 - 499	0.02	0.00	0.07
500+	0.05	0.01	0.11

Table 42: Elbows

$cd4_group$	$point_est$	$lower_ci$	upper_ci
0-99	0.01	0.00	0.04
100-199	0.05	0.01	0.09
200-299	0.02	0.00	0.06
300-399	0.04	0.00	0.08
400-499	0.07	0.00	0.15
500+	0.07	0.02	0.12

Table 43: Wrists & Hands

$cd4_group$	$point_est$	$lower_ci$	upper_ci
0-99	0.01	0.00	0.04
100-199	0.06	0.02	0.10
200-299	0.06	0.02	0.10
300-399	0.06	0.01	0.12
400-499	0.07	0.00	0.13
500+	0.14	0.08	0.22

Table 44: Chest

cd4_group	point_est	lower_ci	upper_ci
0-99	0.34	0.22	0.46
100-199	0.23	0.15	0.31
200-299	0.17	0.10	0.23
300-399	0.17	0.09	0.26
400-499	0.17	0.09	0.28
500+	0.11	0.05	0.17

Table 45: Lower back

$cd4_group$	$point_est$	lower_ci	upper_ci
0-99	0.06	0.01	0.12
100-199	0.08	0.04	0.13
200-299	0.06	0.02	0.10
300-399	0.06	0.01	0.13

cd4_group	point_est	lower_ci	upper_ci
400-499	0.11	0.02	0.20
500+	0.10	0.04	0.16

Table 46: Abdomen

$cd4_group$	$point_est$	lower_ci	upper_ci
0-99	0.34	0.22	0.46
100-199	0.29	0.21	0.38
200-299	0.27	0.19	0.36
300-399	0.23	0.14	0.32
400-499	0.30	0.17	0.46
500+	0.22	0.13	0.30

Table 47: Cervical spine

$cd4_group$	$point_est$	$lower_ci$	upper_ci
0-99	0.04	0.00	0.10
100-199	0.09	0.04	0.15
200-299	0.02	0.00	0.05
300-399	0.06	0.01	0.13
400-499	0.09	0.02	0.17
500+	0.03	0.00	0.08

Table 48: Thoracic spine

cd4_group	point_est	lower_ci	upper_ci
0-99	0.13	0.06	0.22
100-199	0.11	0.06	0.18
200-299	0.10	0.05	0.16
300-399	0.16	0.08	0.25
400-499	0.22	0.11	0.35
500+	0.14	0.08	0.22

Table 49: Lumbosacral spine

cd4_group	point_est	$lower_ci$	upper_ci
0-99	0.09	0.03	0.16
100-199	0.14	0.08	0.22
200-299	0.15	0.08	0.22
300-399	0.18	0.10	0.29
400-499	0.30	0.17	0.43
500+	0.21	0.13	0.28

Table 50: Groin

cd4_group	point_est	lower_ci	upper_ci
0-99	0.10	0.03	0.18
100-199	0.12	0.07	0.19
200-299	0.09	0.05	0.16
300-399	0.12	0.05	0.19
400-499	0.11	0.02	0.20
500+	0.08	0.02	0.13

Table 51: Hips

$cd4_group$	$point_est$	lower_ci	upper_ci
0-99	0.01	0.00	0.06
100-199	0.06	0.02	0.10
200 - 299	0.05	0.01	0.09
300-399	0.05	0.01	0.10
400-499	0.11	0.02	0.20
500+	0.14	0.08	0.22

Table 52: Legs

cd4_group	point_est	lower_ci	upper_ci
0-99	0.13	0.06	0.22
100-199	0.11	0.06	0.18
200-299	0.15	0.08	0.21
300-399	0.18	0.10	0.27
400-499	0.20	0.09	0.30
500+	0.23	0.15	0.32

Table 53: Knees

$cd4_group$	$point_est$	$lower_ci$	upper_ci
0-99	0.12	0.04	0.19
100-199	0.11	0.06	0.18
200-299	0.15	0.08	0.21
300-399	0.12	0.05	0.19
400-499	0.17	0.07	0.28
500+	0.22	0.14	0.30

Table 54: Ankles & Feet

cd4_group	point_est	lower_ci	upper_ci
0-99	0.26	0.16	0.37
100-199	0.30	0.22	0.40
200-299	0.31	0.23	0.41
300-399	0.25	0.16	0.34

cd4_group	point_est	lower_ci	upper_ci
400-499	0.26	0.15	0.39
500+	0.35	0.25	0.45

Table 55: Buttocks

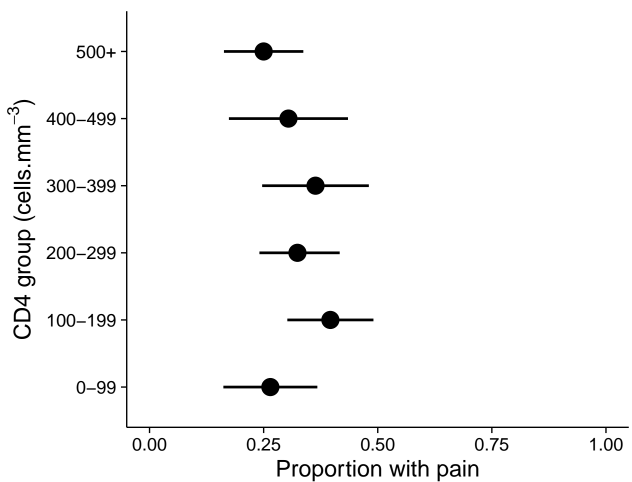
$cd4_group$	$point_est$	$lower_ci$	upper_ci
0-99	0.04	0.00	0.10
100-199	0.01	0.00	0.03
200-299	0.03	0.00	0.06
300-399	0.05	0.01	0.10
400-499	0.04	0.00	0.11
500+	0.07	0.02	0.12

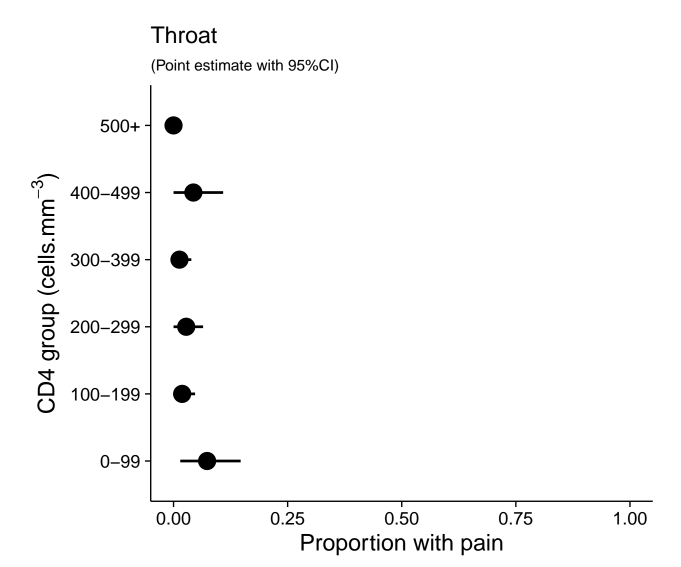
6.3 $\,$ Plotted proportions (with 95% CIs), by CD4 (recent) group and body site

walk(cd4_boot2\$plots, ~ print(.x))

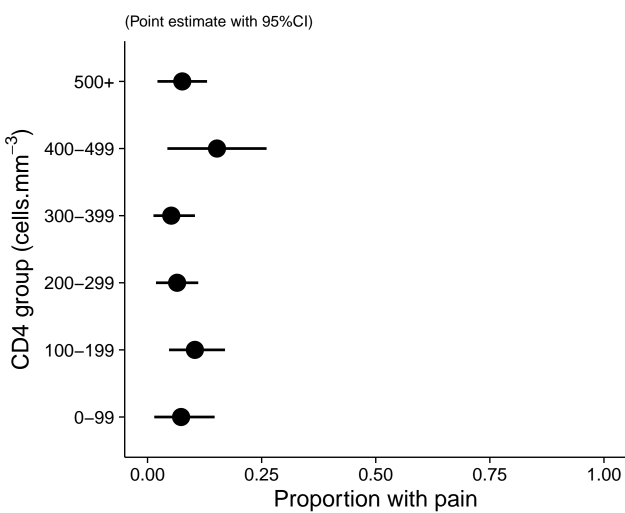
Head

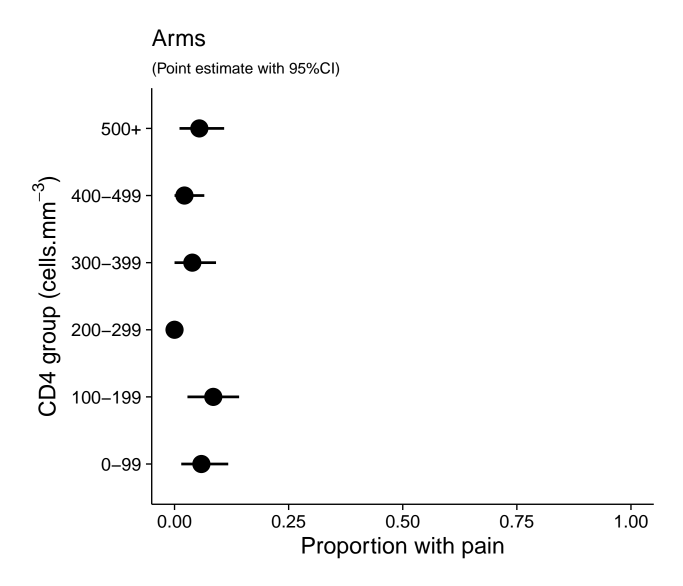
(Point estimate with 95%CI)



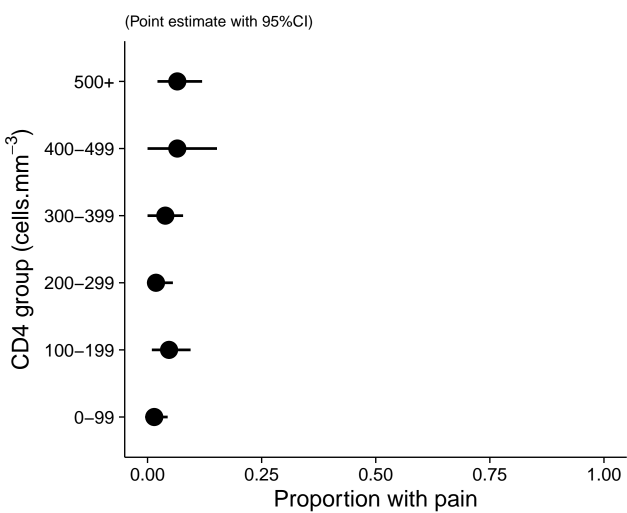






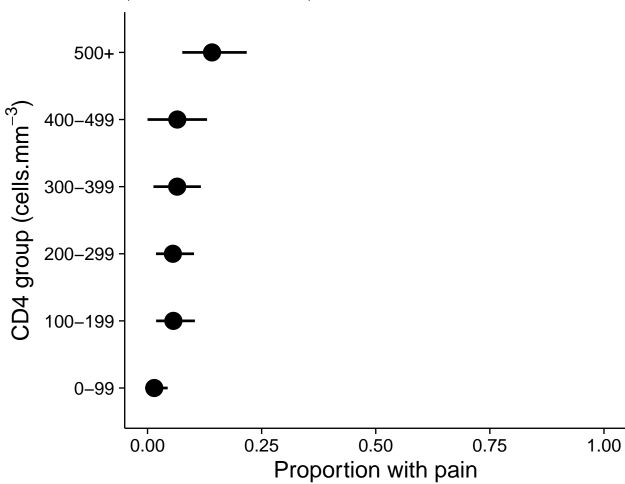


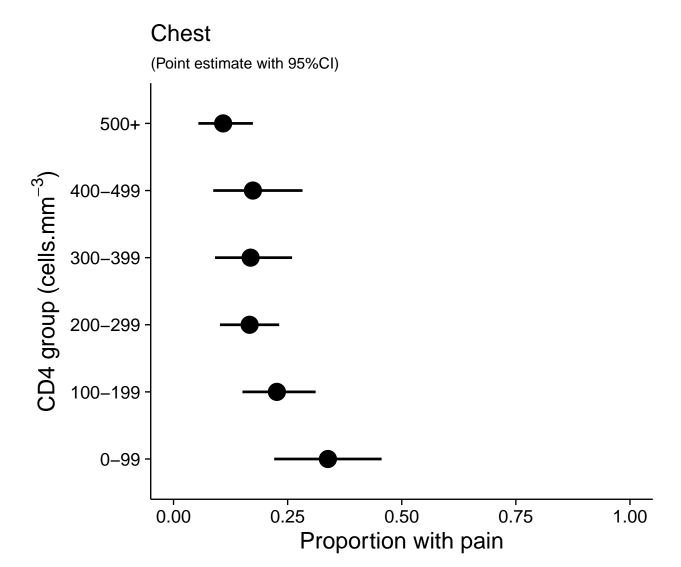


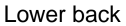


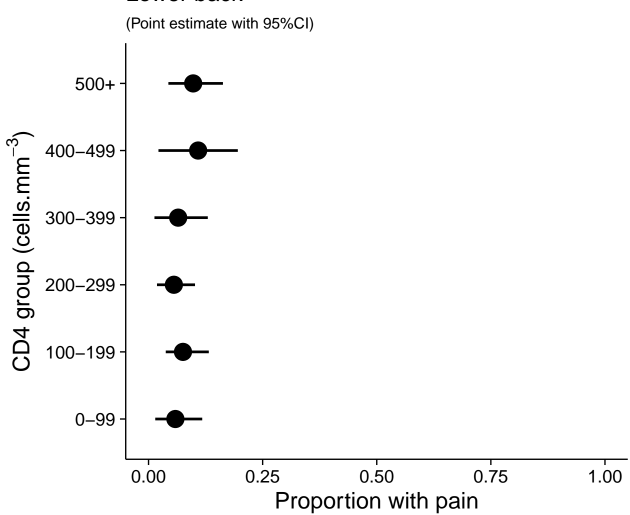


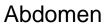


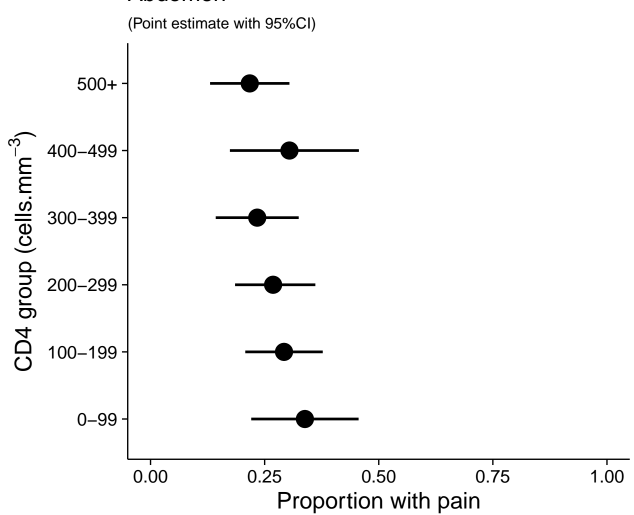




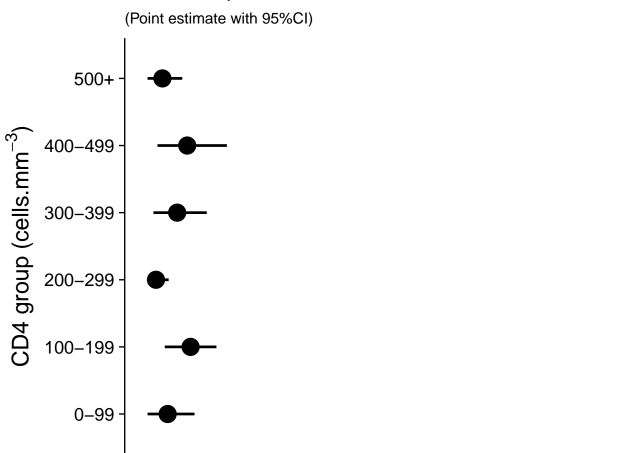








Cervical spine



0.50

Proportion with pain

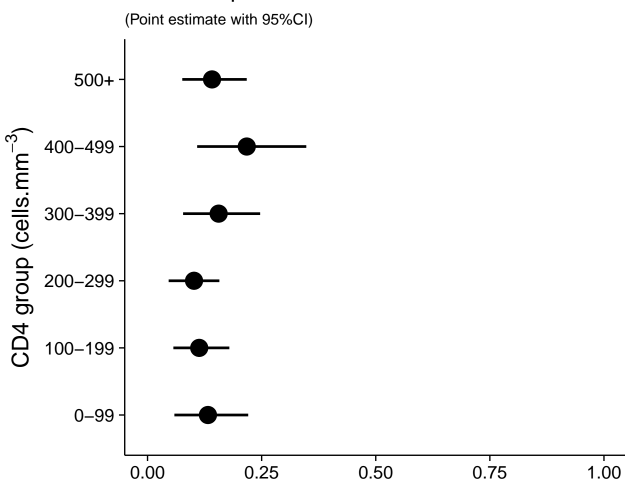
0.75

1.00

0.25

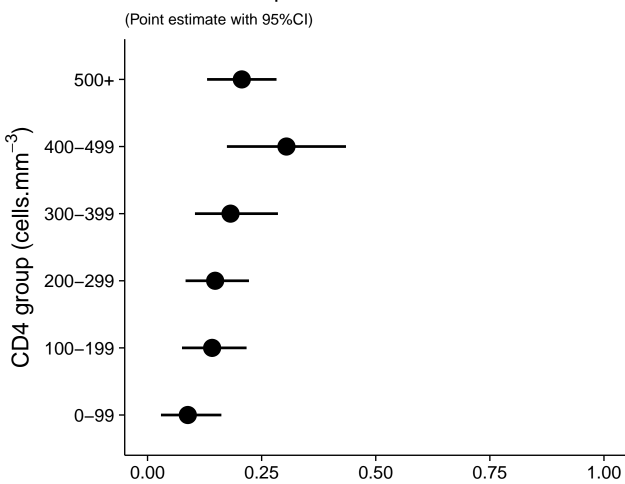
0.00

Thoracic spine



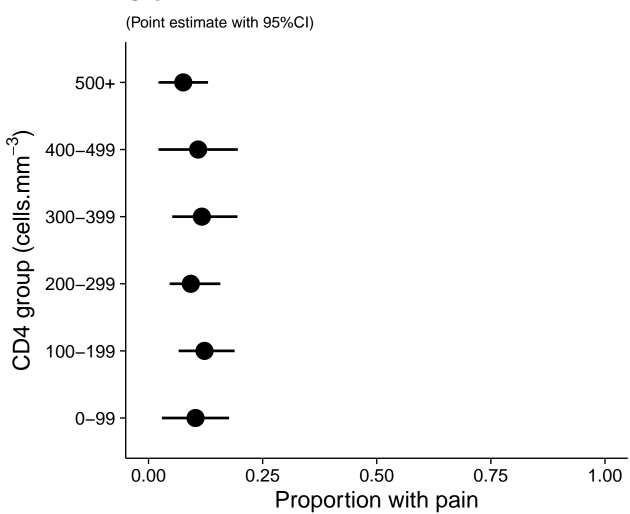
Proportion with pain

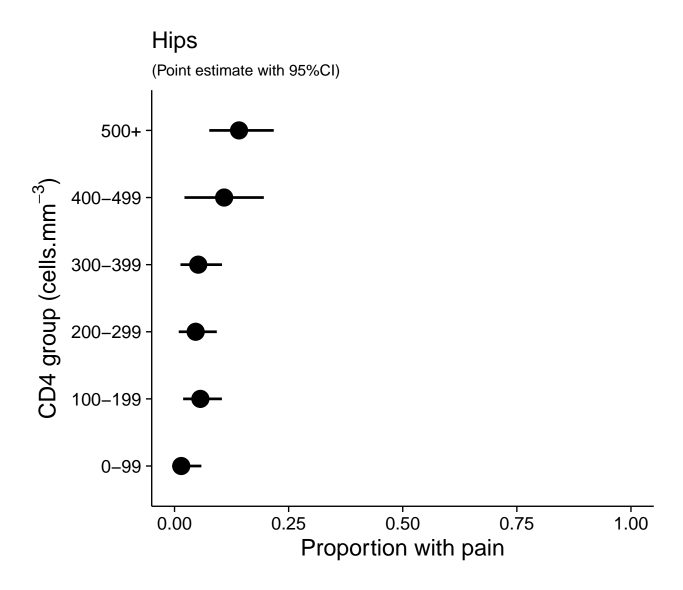
Lumbosacral spine

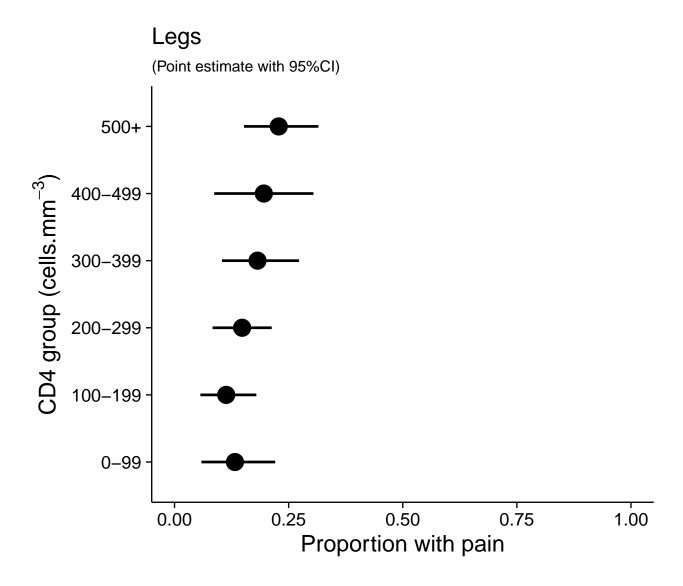


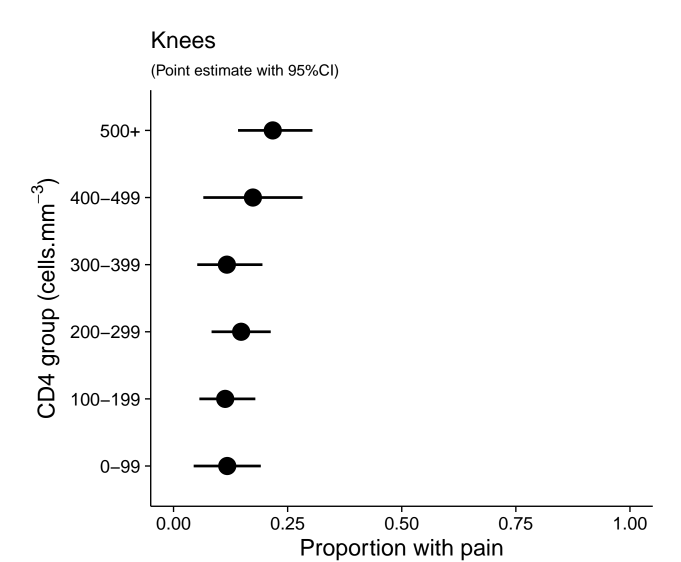
Proportion with pain



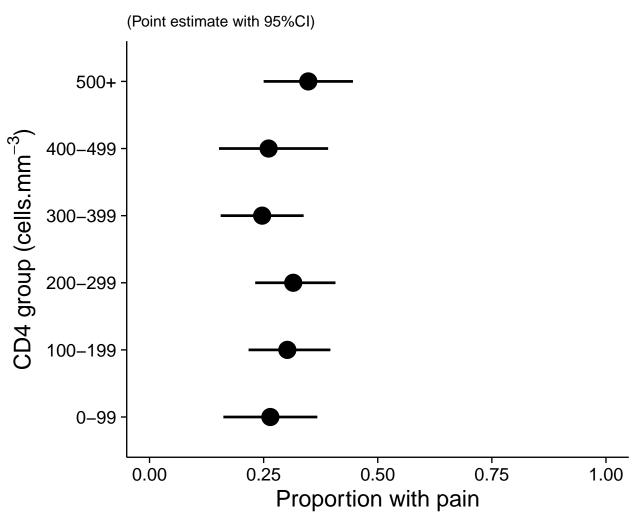






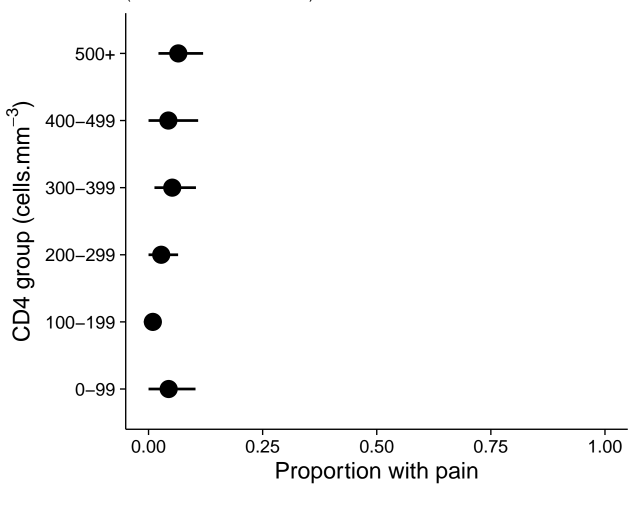






Buttocks





7 By pain definition

Pain in the last week or month (acute) versus persistent pain for the last 3 months (chronic)

7.1 Process data

```
# Set seed
set.seed(2020)

# Select pain data
pain <- demo[, c('ID', 'Pain_def')]

# Join to boot_data & remove ID, site, and upper_back (only one outcome -- no pain)
pain <- left_join(data, pain) %>%
    select(-ID, -Site, -Upper_back)

# Bootstrap CIs
```

```
pain_boot <- pain %>%
    # Pivot to long format
    pivot_longer(cols = -Pain_def,
                 names_to = 'body_site',
                 values_to = 'pain_present') %>%
    # Add body regions
   mutate(region = case_when(
        body_site == 'Chest' |
            body_site == 'Head' |
            body_site == 'Throat' |
            body_site == 'Shoulder' ~ 'Head and upper torso',
        body_site == 'Lower_back' |
            body_site == 'Abdomen' |
            body_site == 'Hips' |
            body_site == 'Buttocks' |
            body_site == 'Groin' ~ 'Lower torso',
        body_site == 'Legs' |
            body_site == 'Knees' |
            body_site == 'Ankles.Feet' ~ 'Lower limbs',
        body_site == 'Arms' |
            body_site == 'Elbows' |
            body_site == 'Wrists.Hands' ~ 'Upper limbs',
        body_site == 'Cervical_spine' |
            body_site == 'Thoracic_spine' |
            body_site == 'Lumbosacral_spine' ~ 'Spinal column',
       TRUE ~ 'other'
   )) %>%
    # Nest by body region and body site
   group_by(Pain_def, region, body_site) %>%
   nest() %>%
    # Boostrap data
    mutate(boot = map(.x = data,
                      ~ boot(data = .x,
                             statistic = prop_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
    mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                           ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
    select(-data, -boot, -ci) %>%
    # Unnest
   unnest(cols = c(point_est, lower_ci, upper_ci))
```

```
# Re-nest by body region and generate figures and tables
pain_boot2 <- pain_boot %>%
   group_by(region) %>%
   nest() %>%
    # Fix site labels
   mutate(data = map(.x = data,
                      ~ .x %>%
                          mutate(body_site = str_replace_all(body_site,
                                                         pattern = '_',
                                                         replacement = ' '),
                                 body_site = str_replace_all(body_site,
                                                        pattern = '\\.',
                                                        replacement = ' & ')))) %>%
    # Re-order sites by point_est
   mutate(data = map(.x = data,
                      ~ .x %>%
                          mutate(body_site = fct_reorder(body_site,
                                                    point_est)))) %>%
    # Plot data
   mutate(plots = map2(.x = data,
                        .y = region,
                       ~ .x %>%
                           ggplot(data = .) +
                           aes(x = body_site,
                               y = point_est,
                               ymin = lower_ci,
                               ymax = upper ci,
                               fill = Pain_def) +
                           geom_linerange(position = position_dodge2(width = 0.6),
                                          size = 1,
                                          colour = '#000000') +
                           geom_point(shape = 21,
                                      colour = '#000000',
                                      position = position_dodge2(width = 0.6),
                                      size = 6,
                                      stroke = 1) +
                           coord_flip() +
                           labs(title = .y,
                                subtitle = '(Point estimate with 95%CI)',
                                y = 'Proportion with pain') +
                           scale_y_continuous(limits = c(0, 1)) +
                           scale_fill_manual(values = c('#000000', '#FFFFFF')) +
                           theme_minimal(base_size = 18) +
                           theme(plot.title = element text(size = 18),
                                 plot.subtitle = element_text(size = 12),
                                 legend.title = element_blank(),
                                 legend.position = 'top',
                                 axis.title.y = element_blank(),
                                 panel.grid = element_blank(),
                                 axis.text = element_text(colour = '#000000'),
                                 axis.line = element_line(size = 0.5),
                                 axis.ticks = element_line(size = 0.5)))) %>%
    # Tabulate data
```

7.2 Tabulated proportions (with 95% CIs), by age and body region

```
walk(pain_boot2$tables, ~ print(.x))
```

Table 56: Head and upper torso

Pain_def	$body_site$	point_est	lower_ci	upper_ci
Acute	Head	0.38	0.33	0.43
Acute	Throat	0.03	0.01	0.05
Acute	Shoulder	0.08	0.05	0.11
Acute	Chest	0.24	0.20	0.28
Chronic	Head	0.23	0.18	0.30
Chronic	Throat	0.03	0.01	0.06
Chronic	Shoulder	0.08	0.04	0.11
Chronic	Chest	0.12	0.08	0.17

Table 57: Upper limbs

Pain_def	body_site	point_est	lower_ci	upper_ci
Acute	Arms	0.05	0.03	0.07
Acute	Elbows	0.03	0.02	0.05
Acute	Wrists & Hands	0.05	0.03	0.07
Chronic	Arms	0.02	0.00	0.05
Chronic	Elbows	0.05	0.02	0.08
Chronic	Wrists & Hands	0.08	0.04	0.11

Table 58: Lower torso

Pain_def	body_site	point_est	lower_ci	upper_ci
Acute	Lower back	0.07	0.04	0.09
Acute	Abdomen	0.30	0.26	0.35
Acute	Groin	0.11	0.08	0.15
Acute	Hips	0.02	0.01	0.04
Acute	Buttocks	0.03	0.02	0.05
Chronic	Lower back	0.08	0.04	0.11
Chronic	Abdomen	0.20	0.15	0.26
Chronic	Groin	0.05	0.02	0.08
Chronic	Hips	0.15	0.11	0.20
Chronic	Buttocks	0.03	0.01	0.06

Table 59: Spinal column

Pain_def	body_site	point_est	lower_ci	upper_ci
Acute	Cervical spine	0.07	0.05	0.10
Acute	Thoracic spine	0.14	0.11	0.18
Acute	Lumbosacral spine	0.14	0.11	0.18
Chronic	Cervical spine	0.01	0.00	0.02
Chronic	Thoracic spine	0.10	0.06	0.13
Chronic	Lumbosacral spine	0.18	0.13	0.23

Table 60: Lower limbs

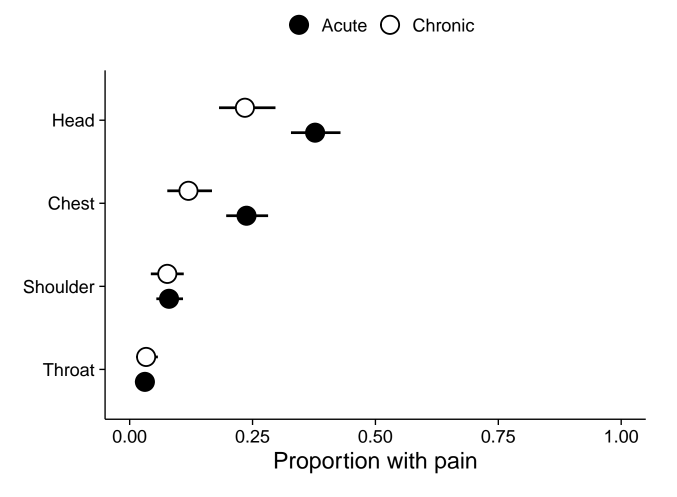
Pain_def	body_site	point_est	lower_ci	upper_ci
Acute	Legs	0.12	0.09	0.15
Acute	Knees	0.10	0.07	0.13
Acute	Ankles & Feet	0.23	0.19	0.27
Chronic	Legs	0.25	0.19	0.32
Chronic	Knees	0.22	0.16	0.27
Chronic	Ankles & Feet	0.46	0.39	0.53

7.3 Plotted proportions (with 95% CIs), by age and body region

walk(pain_boot2\$plots, ~ print(.x))

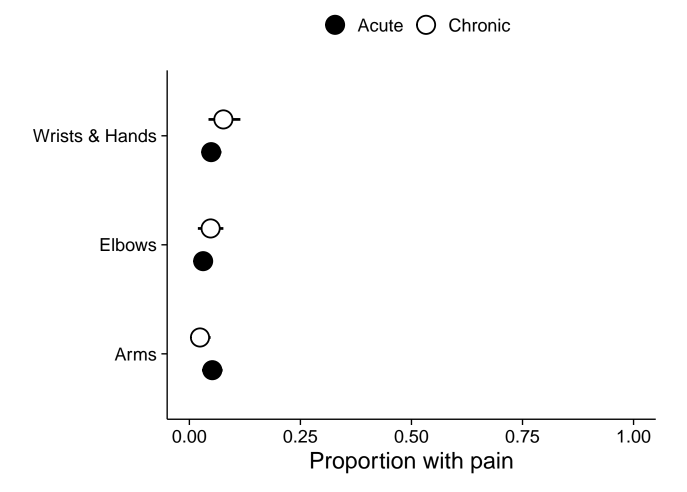
Head and upper torso

(Point estimate with 95%CI)



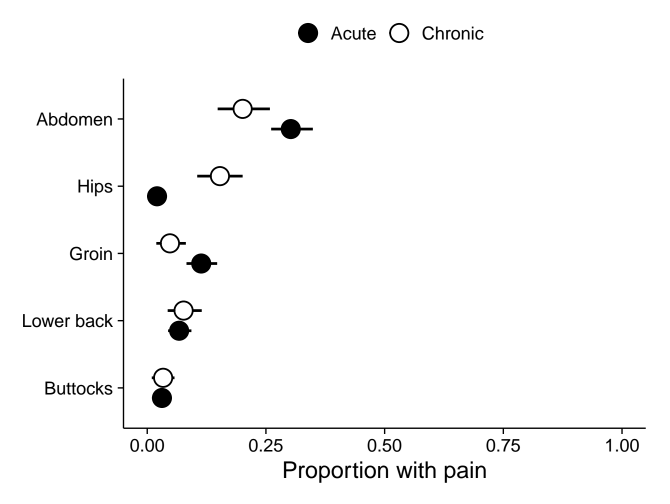
Upper limbs

(Point estimate with 95%CI)



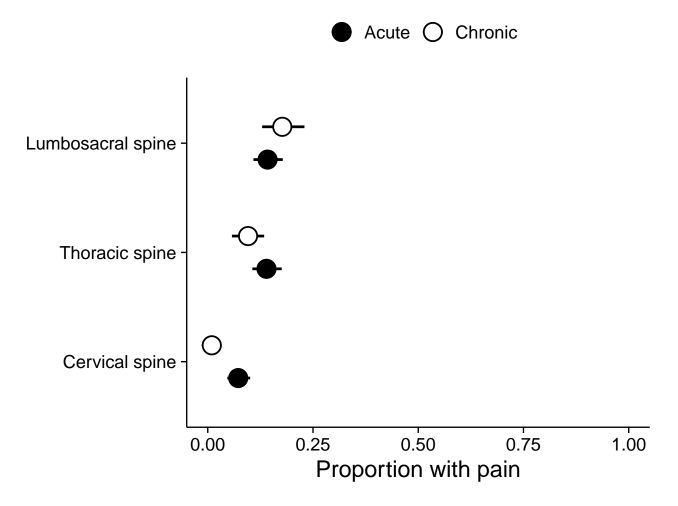
Lower torso

(Point estimate with 95%CI)



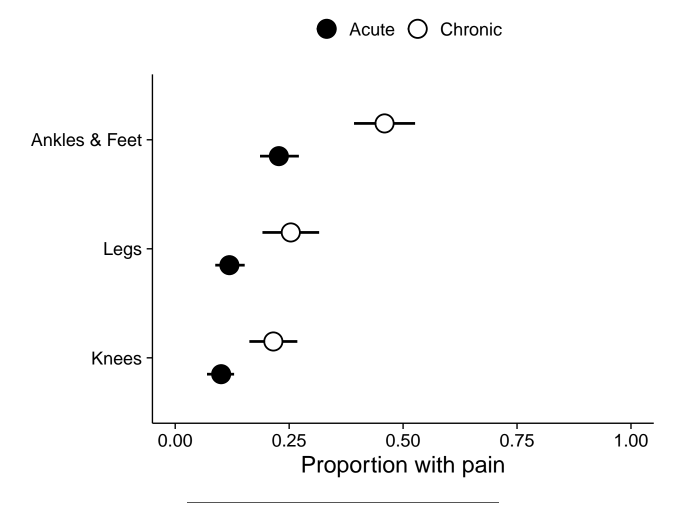
Spinal column

(Point estimate with 95%CI)



Lower limbs

(Point estimate with 95%CI)



8 Logistic regression

Note: Age and CD4 T-cell count treated as a continuous variables.

Note: Age and CD-4 T-cell count were scaled and centered.

8.1 Process data

Extract and process data on study site, participant ID, CD4 T-cell count, age, sex, and pain definition.

8.2 Family-wise error correction

Models:

Alpha threshold for significance was corrected to control the family-wise error rate. The correction was applied to the results of the likelihood ratio tests comparing null models to full models. We chose the conservative Dunn-Bonferroni correction.

8.3 Run glmer model for every body site

Note: p-values reported for the likelihood ratio tests below are uncorrected for multiple comparisons.

```
#-- Head --#
head <- data_logit.long %>%
    # Filter by pain site
    filter(Pain_site == 'Head') %>%
    filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = head,
              family = binomial())
## Full model
head_mod <- glmer(Pain_present ~
                  Age +
                  Sex +
                  CD4_recent +
                  Pain def +
                   (1|Site),
              data = head,
              family = binomial())
## Compare models
anova(null_mod, head_mod, test = 'LRT')
## Data: head
```

```
## null_mod: Pain_present ~ 1 + (1 | Site)
## head_mod: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
                         BIC logLik deviance Chisq Df Pr(>Chisq)
           npar AIC
              2 599.13 607.53 -297.56
                                       595.13
## null_mod
## head mod
              6 589.86 615.06 -288.93
                                      577.86 17.27 4 0.001713 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Significant
## (family-wise error correction, alpha = 0.0024)
## Print model
summary(head mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
     Data: head
##
##
       AIC
                BIC logLik deviance df.resid
##
     589.9
              615.1
                     -288.9
                                577.9
##
## Scaled residuals:
      Min
               1Q Median
                               3Q
## -1.0014 -0.7480 -0.4873 1.0605 2.6634
##
## Random effects:
                      Variance Std.Dev.
## Groups Name
## Site
          (Intercept) 0.1263 0.3553
## Number of obs: 493, groups: Site, 5
##
## Fixed effects:
                  Estimate Std. Error z value Pr(>|z|)
                             0.28339 -1.385 0.16599
## (Intercept)
                  -0.39255
## Age
                   0.03543
                              0.12234
                                      0.290 0.77213
## SexMale
                  -1.34687
                              0.38448 -3.503 0.00046 ***
## CD4_recent
                              0.11511 -0.967 0.33344
                  -0.11133
## Pain_defChronic -0.54907
                              0.42960 -1.278 0.20121
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
                            SexMal CD4_rc
              (Intr) Age
              0.186
## Age
              -0.094 -0.103
## SexMale
              0.068 -0.071 0.081
## CD4 recent
## Pan_dfChrnc -0.685 -0.238 -0.042 -0.203
#-- Throat --#
throat <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Throat') %>%
   filter(complete.cases(.))
## Null
```

```
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
             data = throat,
             family = binomial())
## Full model
thrt_mod <- glmer(Pain_present ~</pre>
                 Age +
                 Sex +
                 CD4 recent +
                 Pain def +
                  (1|Site),
             data = throat,
             family = binomial())
## Fit is singular, print model to check SD of random effect
## Print model
summary(thrt_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
     Data: throat
##
##
       AIC
                BIC
                      logLik deviance df.resid
##
     125.0
              150.2
                       -56.5
                                113.0
                                           487
##
## Scaled residuals:
##
               1Q Median
                               3Q
## -0.3660 -0.1799 -0.1466 -0.1094 8.5792
##
## Random effects:
## Groups Name
                      Variance Std.Dev.
## Site
          (Intercept) 0
## Number of obs: 493, groups: Site, 5
##
## Fixed effects:
##
                  Estimate Std. Error z value
                                                         Pr(>|z|)
                              ## (Intercept)
                  -4.12952
## Age
                   0.07201
                              0.33111
                                       0.217
                                                           0.8278
## SexMale
                   0.91911
                              0.62751
                                        1.465
                                                           0.1430
## CD4_recent
                  -0.78730
                              0.45238 - 1.740
                                                           0.0818 .
## Pain_defChronic 0.19658
                              0.68881
                                       0.285
                                                           0.7753
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
              (Intr) Age
                            SexMal CD4_rc
## Age
               0.160
## SexMale
              -0.330 -0.224
              0.512 -0.132 0.168
## CD4_recent
## Pan_dfChrnc -0.475 -0.240 -0.211 -0.268
## convergence code: 0
## boundary (singular) fit: see ?isSingular
```

```
#-- Shoulder --#
shoulder <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Shoulder') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = shoulder,
              family = binomial())
## Full model
shdr_mod <- glmer(Pain_present ~</pre>
                  Age +
                  Sex +
                  CD4_recent +
                  Pain_def +
                  (1|Site),
              data = shoulder,
              family = binomial())
## Compare models
anova(null_mod, shdr_mod, test = 'LRT')
## Data: shoulder
## Models:
## null_mod: Pain_present ~ 1 + (1 | Site)
## shdr mod: Pain present ~ Age + Sex + CD4 recent + Pain def + (1 | Site)
           npar AIC BIC logLik deviance Chisq Df Pr(>Chisq)
## null mod
            2 286.42 294.82 -141.21
                                         282.42
            6 292.25 317.45 -140.12
                                       280.25 2.1705 4
## shdr_mod
                                                              0.7044
## No significant difference between null and full model
## Print model
## Print model
summary(shdr mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
   Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
     Data: shoulder
##
##
##
       AIC
                BIC
                       logLik deviance df.resid
                     -140.1
##
      292.2
              317.5
                                 280.2
                                            487
##
## Scaled residuals:
      Min 1Q Median
                               ЗQ
## -0.4264 -0.3111 -0.2893 -0.2690 4.2089
##
## Random effects:
## Groups Name
                      Variance Std.Dev.
## Site (Intercept) 0.01967 0.1403
## Number of obs: 493, groups: Site, 5
##
```

```
## Fixed effects:
##
                   Estimate Std. Error z value
                                                          Pr(>|z|)
## (Intercept)
                             0.25193 -10.187 <0.0000000000000000 ***
                  -2.56646
                   -0.14999
                               0.20336 -0.738
## Age
                                                             0.461
## SexMale
                   0.53246
                               0.43017
                                        1.238
                                                             0.216
## CD4 recent
                   -0.06105
                               0.18801 -0.325
                                                             0.745
## Pain_defChronic 0.10528
                               0.46037
                                       0.229
                                                             0.819
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
              (Intr) Age
                             SexMal CD4_rc
               0.362
## Age
              -0.293 -0.137
## SexMale
## CD4_recent 0.111 -0.131 0.122
## Pan_dfChrnc -0.529 -0.368 -0.252 -0.199
#-- Arm --#
arms <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Arms') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = arms,
              family = binomial())
## Full model
arms_mod <- glmer(Pain_present ~</pre>
                  Age +
                  Sex +
                  CD4_recent +
                  Pain_def +
                  (1|Site),
              data = arms,
              family = binomial())
## Fit is singular, print model to check SD of random effect
## Print model
summary(arms_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
     Data: arms
##
##
        AIC
                 BIC
                       logLik deviance df.resid
##
      189.1
               214.3
                        -88.6
                                 177.1
                                            487
##
## Scaled residuals:
      Min
                1Q Median
                                ЗQ
## -0.4341 -0.2334 -0.2115 -0.1835 5.5338
##
```

```
## Random effects:
## Groups Name
                     Variance Std.Dev.
## Site (Intercept) 0
## Number of obs: 493, groups: Site, 5
## Fixed effects:
                 Estimate Std. Error z value
                                                       Pr(>|z|)
                ## (Intercept)
                                     1.417
## Age
                  0.30866 0.21783
                                                          0.156
                 -0.05672 0.59049 -0.096
## SexMale
                                                          0.923
## CD4_recent
                  -0.10059 0.25597 -0.393
                                                          0.694
## Pain_defChronic -0.62336 0.59010 -1.056
                                                          0.291
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
             (Intr) Age
                           SexMal CD4_rc
## Age
              0.144
## SexMale
             -0.310 -0.207
## CD4 recent 0.222 -0.093 0.123
## Pan_dfChrnc -0.468 -0.297 -0.101 -0.317
## convergence code: 0
## boundary (singular) fit: see ?isSingular
#-- Elbow --#
elbow <- data_logit.long %>%
   # Filter by pain site
   filter(Pain site == 'Elbows') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
             data = elbow,
             family = binomial())
## Full model
elbw_mod <- glmer(Pain_present ~
                 Age +
                 Sex +
                 CD4 recent +
                 Pain_def +
                 (1|Site),
             data = elbow,
             family = binomial())
## Fit is singular, print model to check SD of random effect
## Print model
summary(elbw_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
     Data: elbow
##
```

```
AIC
##
                BIC
                      logLik deviance df.resid
##
      172.1
              197.3
                      -80.1
                                160.1
                                            487
##
## Scaled residuals:
      Min
              1Q Median
                               3Q
## -0.4574 -0.2127 -0.1764 -0.1533 6.7009
## Random effects:
## Groups Name
                       Variance Std.Dev.
## Site
         (Intercept) 0
## Number of obs: 493, groups: Site, 5
## Fixed effects:
##
                  Estimate Std. Error z value
                                                          Pr(>|z|)
## (Intercept)
                  -3.38921
                              0.33556 -10.100 <0.0000000000000000 ***
## Age
                   0.42521
                              0.21018
                                       2.023
                                                            0.0431 *
## SexMale
                              0.59159 -0.281
                  -0.16645
                                                            0.7784
## CD4 recent
                   0.04229
                              0.23571
                                       0.179
                                                            0.8576
## Pain_defChronic 0.45074
                              0.53643
                                       0.840
                                                            0.4008
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
               (Intr) Age
##
                            SexMal CD4 rc
## Age
               0.023
## SexMale
              -0.244 -0.138
## CD4_recent 0.103 -0.100 0.138
## Pan_dfChrnc -0.595 -0.316 -0.115 -0.352
## convergence code: 0
## boundary (singular) fit: see ?isSingular
#-- Wrist & hand --#
hand <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Wrists.Hands') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
             data = hand,
             family = binomial())
## Full model
hand_mod <- glmer(Pain_present ~
                 Age +
                 Sex +
                 CD4 recent +
                 Pain_def +
                  (1|Site),
              data = hand,
              family = binomial())
## Fit is singular, print model to check SD of random effect
## Print model
```

```
summary(hand_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
      Data: hand
##
##
       AIC
                       logLik deviance df.resid
                BIC
##
      243.3
               268.5
                       -115.6
                                 231.3
##
## Scaled residuals:
##
      Min
               1Q Median
                                ЗQ
                                       Max
## -0.6343 -0.2735 -0.2284 -0.2038 6.8403
##
## Random effects:
## Groups Name
                       Variance Std.Dev.
          (Intercept) 0
## Site
## Number of obs: 493, groups: Site, 5
## Fixed effects:
##
                  Estimate Std. Error z value
                                                          Pr(>|z|)
## (Intercept)
                   -2.7754
                             0.2588 -10.722 <0.0000000000000000 ***
## Age
                     0.2316
                                0.1805
                                        1.283
                                                             0.199
                                                             0.251
## SexMale
                    -0.6507
                                0.5663 - 1.149
## CD4_recent
                     0.2766
                                0.1711
                                        1.616
                                                             0.106
## Pain defChronic
                     0.4408
                                0.4376
                                       1.007
                                                             0.314
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
              (Intr) Age
                             SexMal CD4_rc
## Age
               0.191
## SexMale
              -0.208 -0.145
## CD4_recent -0.005 -0.020 0.126
## Pan_dfChrnc -0.601 -0.377 -0.100 -0.374
## convergence code: 0
## boundary (singular) fit: see ?isSingular
#-- Chest --#
chest <- data_logit.long %>%
    # Filter by pain site
    filter(Pain_site == 'Chest') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = chest,
              family = binomial())
## Full model
```

chst_mod <- glmer(Pain_present ~</pre>

Age + Sex +

```
CD4_recent +
                 Pain_def +
                 (1|Site),
             data = chest,
             family = binomial())
## Compare models
anova(null_mod, chst_mod, test = 'LRT')
## Data: chest
## Models:
## null_mod: Pain_present ~ 1 + (1 | Site)
## chst_mod: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
                 AIC
                          BIC logLik deviance Chisq Df Pr(>Chisq)
           npar
              2 479.88 488.28 -237.94
                                       475.88
## null mod
                                       468.58 7.3024 4
              6 480.58 505.78 -234.29
## chst mod
                                                            0.1207
## No significant difference in null and full models
## Print model
## Print model
summary(chst mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
     Data: chest
##
##
       AIC
                BIC
                      logLik deviance df.resid
##
     480.6
              505.8 -234.3
                               468.6
                                          487
##
## Scaled residuals:
##
      Min
           1Q Median
                              3Q
## -0.7467 -0.5449 -0.4222 -0.3084 4.7140
## Random effects:
## Groups Name
                      Variance Std.Dev.
## Site (Intercept) 0.04797 0.219
## Number of obs: 493, groups: Site, 5
## Fixed effects:
##
                  Estimate Std. Error z value
                                                  Pr(>|z|)
## (Intercept)
                  0.680
## Age
                   0.09422
                             0.13852
                                                    0.4964
## SexMale
                   0.04311
                             0.32631
                                      0.132
                                                    0.8949
## CD4_recent
                  -0.27859
                             0.14976 -1.860
                                                    0.0628
## Pain_defChronic -0.75437
                             0.38896 -1.939
                                                    0.0524 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
              (Intr) Age
                           SexMal CD4_rc
## Age
              0.254
              -0.150 -0.107
## SexMale
## CD4_recent 0.132 -0.151 0.064
```

```
## Pan_dfChrnc -0.562 -0.253 -0.118 -0.213
#-- Abdomen --#
abdomen <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Abdomen') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = abdomen,
              family = binomial())
## Full model
abdm_mod <- glmer(Pain_present ~
                  Age +
                  Sex +
                  CD4_recent +
                  Pain_def +
                  (1|Site),
              data = abdomen,
              family = binomial())
## Fit is singular, print model to check SD of random effect
## Print model
summary(abdm_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
     Data: abdomen
##
       AIC
##
                 BIC
                       logLik deviance df.resid
##
      581.2
               606.4
                      -284.6
                                 569.2
                                            487
##
## Scaled residuals:
      Min
                1Q Median
                                ЗQ
## -0.8003 -0.6690 -0.5480 1.3841 2.3391
##
## Random effects:
## Groups Name
                       Variance Std.Dev.
## Site
           (Intercept) 0
## Number of obs: 493, groups: Site, 5
##
## Fixed effects:
##
                   Estimate Std. Error z value
                                                     Pr(>|z|)
## (Intercept)
                   -0.8813
                                0.1365 -6.458 0.000000000106 ***
                    -0.1613
                                0.1219 -1.323
                                                        0.186
## Age
## SexMale
                    -0.3072
                                                        0.308
                                0.3015 -1.019
## CD4_recent
                    -0.1145
                                0.1154 -0.993
                                                        0.321
## Pain_defChronic -0.2542
                                0.2706 -0.940
                                                        0.347
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
```

```
## Correlation of Fixed Effects:
                             SexMal CD4_rc
##
              (Intr) Age
## Age
               0.403
              -0.275 -0.183
## SexMale
## CD4_recent 0.170 -0.038 0.157
## Pan_dfChrnc -0.537 -0.333 -0.143 -0.324
## convergence code: 0
## boundary (singular) fit: see ?isSingular
#-- Lower back (flank) --#
low_back <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Lower_back') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = low_back,
              family = binomial())
## Full model
back_mod <- glmer(Pain_present ~</pre>
                  Age +
                  Sex +
                  CD4_recent +
                  Pain def +
                  (1|Site),
              data = low_back,
              family = binomial())
## Fit is singular, print model to check SD of random effect
## Print model
summary(back_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
     Data: low_back
##
        AIC
##
                 BIC
                       logLik deviance df.resid
                       -130.1
##
      272.1
               297.4
                                 260.1
##
## Scaled residuals:
       Min 1Q Median
                              3Q
## -0.4014 -0.2965 -0.2655 -0.2528 4.3683
##
## Random effects:
## Groups Name
                       Variance
                                             Std.Dev.
           (Intercept) 0.00000000000001446 0.0000001202
## Number of obs: 493, groups: Site, 5
##
## Fixed effects:
                   Estimate Std. Error z value
                                                          Pr(>|z|)
## (Intercept)
                  -2.59625
                               0.23601 -11.001 <0.00000000000000000 ***
```

```
## Age
                   0.10656
                               0.18068
                                       0.590
                                                             0.555
## SexMale
                   -0.23426
                               0.48221 -0.486
                                                             0.627
## CD4 recent
                                                             0.619
                   0.08667
                               0.17452
                                       0.497
## Pain_defChronic 0.33885
                               0.41624
                                       0.814
                                                             0.416
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
              (Intr) Age
                            SexMal CD4_rc
## Age
               0.259
## SexMale
              -0.246 -0.160
## CD4_recent 0.080 -0.060 0.161
## Pan_dfChrnc -0.591 -0.357 -0.141 -0.355
## convergence code: 0
## boundary (singular) fit: see ?isSingular
#-- Groin --#
groin <- data_logit.long %>%
    # Filter by pain site
    filter(Pain_site == 'Groin') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
             data = groin,
             family = binomial())
## Full model
groin_mod <- glmer(Pain_present ~</pre>
                   Age +
                   Sex +
                   CD4_recent +
                  Pain_def +
                   (1|Site),
               data = groin,
               family = binomial())
## Compare models
anova(null_mod, groin_mod, test = 'LRT')
## Data: groin
## Models:
## null_mod: Pain_present ~ 1 + (1 | Site)
## groin_mod: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
                   AIC BIC logLik deviance Chisq Df Pr(>Chisq)
##
            npar
                2 304.54 312.94 -150.27
                                          300.54
## null mod
               6 308.23 333.43 -148.12
                                          296.23 4.3039 4
## groin_mod
                                                               0.3664
## No significant difference between null and full model
## Print model
summary(groin_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial (logit)
```

```
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
     Data: groin
##
##
        AIC
                       logLik deviance df.resid
                 BIC
##
      308.2
               333.4
                       -148.1
                                 296.2
##
## Scaled residuals:
##
       Min
                1Q Median
                                3Q
                                       Max
## -0.7049 -0.4182 -0.2169 -0.1479 7.0102
##
## Random effects:
## Groups Name
                       Variance Std.Dev.
## Site
          (Intercept) 0.9946 0.9973
## Number of obs: 493, groups: Site, 5
##
## Fixed effects:
##
                   Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                   -2.57300
                              0.77458 -3.322 0.000894 ***
## Age
                   -0.05428
                               0.16923 -0.321 0.748400
## SexMale
                   -0.63053
                               0.44135 -1.429 0.153105
## CD4_recent
                    0.20593
                               0.16740
                                        1.230 0.218616
## Pain_defChronic -0.41363
                               1.05390 -0.392 0.694708
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
               (Intr) Age
                             SexMal CD4_rc
                0.108
## Age
## SexMale
               -0.053 -0.162
## CD4_recent 0.001 -0.081 0.066
## Pan_dfChrnc -0.727 -0.126 -0.019 -0.141
#-- Buttocks --#
buttocks <- data_logit.long %>%
    # Filter by pain site
    filter(Pain_site == 'Buttocks') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = buttocks,
              family = binomial())
## Full model
butt_mod <- glmer(Pain_present ~</pre>
                  Age +
                  Sex +
                  CD4_recent +
                  Pain_def +
                  (1|Site),
              data = buttocks,
              family = binomial())
## Fit is singular, print model to check SD of random effect
```

```
summary(butt_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
     Data: buttocks
##
##
       AIC
                BIC
                      logLik deviance df.resid
     170.7
              195.9
                      -79.4
                               158.7
##
##
## Scaled residuals:
      Min
               1Q Median
                              ЗQ
                                     Max
## -0.3403 -0.2151 -0.1860 -0.1710 6.1831
##
## Random effects:
## Groups Name
                      Variance Std.Dev.
          (Intercept) 0
## Site
## Number of obs: 493, groups: Site, 5
##
## Fixed effects:
##
                  Estimate Std. Error z value
                                                        Pr(>|z|)
## (Intercept)
                  -0.12939
                             0.26949 -0.480
## Age
                                                          0.631
## SexMale
                   0.65435
                             0.57735
                                      1.133
                                                          0.257
## CD4 recent
                   0.23689
                                      1.039
                                                          0.299
                             0.22789
## Pain_defChronic 0.09589
                             0.57052 0.168
                                                          0.867
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
             (Intr) Age
                           SexMal CD4_rc
## Age
              0.379
              -0.386 -0.244
## SexMale
## CD4_recent -0.038 -0.077 0.219
## Pan_dfChrnc -0.512 -0.318 -0.169 -0.344
## convergence code: 0
## boundary (singular) fit: see ?isSingular
#-- Hips --#
hips <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Hips') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
             data = hips,
             family = binomial())
## Full model
hips_mod <- glmer(Pain_present ~
                 Age +
```

Print model

```
Sex +
                 CD4_recent +
                 Pain_def +
                 (1|Site),
             data = hips,
             family = binomial())
## Compare models
anova(null_mod, hips_mod, test = 'LRT')
## Data: hips
## Models:
## null_mod: Pain_present ~ 1 + (1 | Site)
## hips_mod: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
           npar
                         BIC logLik deviance Chisq Df Pr(>Chisq)
                 AIC
              2 224.25 232.65 -110.12
                                       220.25
## null_mod
## hips_mod
              6 225.28 250.49 -106.64
                                       213.28 6.9633 4
                                                            0.1378
## No significant difference between null and full model
## Print model
summary(hips_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
     Data: hips
##
       AIC
##
                BIC
                      logLik deviance df.resid
     225.3
              250.5 -106.6
##
                               213.3
##
## Scaled residuals:
      Min 1Q Median
                              30
## -0.6294 -0.3457 -0.1553 -0.1426 7.1926
## Random effects:
## Groups Name
                      Variance Std.Dev.
          (Intercept) 0.06768 0.2601
## Number of obs: 493, groups: Site, 5
## Fixed effects:
                  Estimate Std. Error z value
                                                         Pr(>|z|)
## (Intercept)
                  -3.74770
                             0.292
## Age
                   0.05564
                             0.19046
                                                        0.770190
## SexMale
                   0.31715
                             0.43874
                                      0.723
                                                        0.469760
## CD4 recent
                   0.12278
                             0.17504
                                      0.701
                                                        0.483039
## Pain_defChronic 1.88246
                             0.55730
                                      3.378
                                                         0.000731 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
              (Intr) Age
                           SexMal CD4_rc
## Age
              0.155
              -0.157 -0.117
## SexMale
```

```
## CD4 recent
              0.006 -0.110 0.243
## Pan_dfChrnc -0.717 -0.284 -0.115 -0.217
#-- Legs --#
legs <- data_logit.long %>%
    # Filter by pain site
    filter(Pain_site == 'Legs') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = legs,
              family = binomial())
## Full model
legs_mod <- glmer(Pain_present ~</pre>
                  Age +
                  Sex +
                  CD4_recent +
                  Pain_def +
                  (1|Site),
              data = legs,
              family = binomial())
## Fit is singular, print model to check SD of random effect
## Print model
summary(legs_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
      Data: legs
##
##
        AIC
                 BIC
                       logLik deviance df.resid
##
      431.0
               456.2
                      -209.5
                                 419.0
                                            487
##
## Scaled residuals:
##
      Min
                1Q Median
                                ЗQ
## -0.8089 -0.4651 -0.3498 -0.3251 3.2001
##
## Random effects:
## Groups Name
                       Variance Std.Dev.
## Site (Intercept) 0
## Number of obs: 493, groups: Site, 5
## Fixed effects:
                   Estimate Std. Error z value
                                                          Pr(>|z|)
                                0.1812 -11.170 <0.0000000000000000 ***
## (Intercept)
                   -2.0238
## Age
                     0.1117
                                0.1305
                                        0.856
                                                            0.3920
                                0.3013 2.020
                                                            0.0433 *
## SexMale
                     0.6088
## CD4_recent
                     0.1644
                                0.1252
                                       1.313
                                                            0.1892
## Pain_defChronic
                     0.6567
                                0.2903
                                        2.262
                                                            0.0237 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

```
##
## Correlation of Fixed Effects:
##
               (Intr) Age
                             SexMal CD4_rc
               0.228
## Age
## SexMale
               -0.325 -0.184
## CD4 recent 0.029 -0.088 0.209
## Pan_dfChrnc -0.581 -0.323 -0.148 -0.332
## convergence code: 0
## boundary (singular) fit: see ?isSingular
#-- Knees --#
knees <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Knees') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = knees,
              family = binomial())
## Full model
knee_mod <- glmer(Pain_present ~</pre>
                  Age +
                  Sex +
                  CD4_recent +
                  Pain_def +
                  (1|Site),
              data = knees,
              family = binomial())
## Fit is singular, print model to check SD of random effect
## Print model
summary(knee_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
      Data: knees
##
##
       AIC
                 BIC
                       logLik deviance df.resid
      400.2
               425.4
                       -194.1
                                 388.2
##
                                             487
##
## Scaled residuals:
                1Q Median
                                3Q
## -0.8381 -0.4380 -0.3233 -0.2914 3.8153
##
## Random effects:
                       Variance Std.Dev.
## Groups Name
           (Intercept) 0
## Site
## Number of obs: 493, groups: Site, 5
## Fixed effects:
##
                   Estimate Std. Error z value
                                                            Pr(>|z|)
```

```
0.1932 -11.314 < 0.0000000000000000 ***
## (Intercept)
                   -2.1860
## Age
                     0.2308
                                0.1323
                                         1.745
                                                            0.08100 .
                     0.4132
                                         1.313
## SexMale
                                0.3147
                                                            0.18928
## CD4_recent
                     0.0948
                                0.1320 0.718
                                                            0.47273
## Pain_defChronic
                    0.8155
                                0.3015
                                       2.705
                                                            0.00683 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
              (Intr) Age
                             SexMal CD4_rc
## Age
               0.160
              -0.295 -0.163
## SexMale
## CD4_recent 0.059 -0.096 0.197
## Pan_dfChrnc -0.604 -0.309 -0.150 -0.336
## convergence code: 0
## boundary (singular) fit: see ?isSingular
#-- Ankles.Feet --#
ankles <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Ankles.Feet') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = ankles,
              family = binomial())
## Full model
ankl_mod <- glmer(Pain_present ~
                  Age +
                  Sex +
                  CD4_recent +
                  Pain_def +
                  (1|Site),
              data = ankles,
              family = binomial())
## Fit is singular, print model to check SD of random effect
## Print model
summary(ankl_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
     Data: ankles
##
##
       AIC
                BIC
                      logLik deviance df.resid
##
      570.2
              595.4
                     -279.1
                                 558.2
                                            487
##
## Scaled residuals:
                1Q Median
                                ЗQ
## -1.4370 -0.5803 -0.4904 0.9236 2.3929
##
```

```
## Random effects:
                      Variance Std.Dev.
## Groups Name
## Site (Intercept) 0
## Number of obs: 493, groups: Site, 5
## Fixed effects:
                  Estimate Std. Error z value
                                                          Pr(>|z|)
                            0.14377 -8.816 < 0.0000000000000000 ***
## (Intercept)
                  -1.26753
                                       2.473
## Age
                   0.27385
                              0.11072
                                                            0.0134 *
## SexMale
                   0.38509
                              0.26289 1.465
                                                            0.1430
## CD4_recent
                  -0.08338
                              0.11100 -0.751
                                                            0.4526
## Pain_defChronic 0.96276
                            0.24465 3.935
                                                        0.0000831 ***
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
              (Intr) Age
                            SexMal CD4_rc
## Age
              0.216
              -0.294 -0.177
## SexMale
## CD4 recent 0.142 -0.093 0.177
## Pan_dfChrnc -0.584 -0.295 -0.135 -0.356
## convergence code: 0
## boundary (singular) fit: see ?isSingular
#-- Cervical spine --#
cervical <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Cervical_spine') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
             data = cervical,
             family = binomial())
## Full model
neck_mod <- glmer(Pain_present ~</pre>
                 Age +
                 Sex +
                 CD4_recent +
                 Pain_def +
                  (1|Site),
              data = cervical,
             family = binomial())
## Compare models
anova(null_mod, neck_mod, test = 'LRT')
## Data: cervical
## Models:
## null_mod: Pain_present ~ 1 + (1 | Site)
## neck_mod: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
                          BIC
                                logLik deviance Chisq Df Pr(>Chisq)
           npar AIC
## null_mod
            2 205.43 213.83 -100.716
                                         201.43
## neck_mod 6 206.19 231.40 -97.096
                                         194.19 7.2402 4
                                                              0.1237
```

```
## No significant difference between null and full models
## Print model
summary(neck_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
     Data: cervical
##
                     logLik deviance df.resid
##
       AIC
                BIC
##
      206.2
               231.4
                       -97.1
                                 194.2
                                            487
##
## Scaled residuals:
      Min
               1Q Median
                                3Q
## -0.3705 -0.3024 -0.2546 -0.0844 11.3198
## Random effects:
                       Variance Std.Dev.
## Groups Name
         (Intercept) 0.08489 0.2914
## Number of obs: 493, groups: Site, 5
##
## Fixed effects:
                  Estimate Std. Error z value
                              0.31368 -7.577 0.000000000000354 ***
                  -2.37663
## (Intercept)
                   0.14746
                               0.26428
                                       0.558
## Age
                                                            0.577
                   -0.60510
## SexMale
                              0.80413 - 0.752
                                                            0.452
## CD4_recent
                   0.06598
                               0.24090
                                       0.274
                                                            0.784
## Pain_defChronic -2.65214
                              1.11087 -2.387
                                                            0.017 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
              (Intr) Age
                             SexMal CD4_rc
## Age
               0.267
              -0.243 -0.106
## SexMale
## CD4_recent 0.102 -0.145 -0.016
## Pan_dfChrnc -0.310 -0.224 -0.046 -0.143
#-- Thoracic spine --#
thoracic <- data_logit.long %>%
    # Filter by pain site
    filter(Pain_site == 'Thoracic_spine') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = thoracic,
              family = binomial())
## Full model
thrx_mod <- glmer(Pain_present ~
                  Age +
                  Sex +
```

```
CD4_recent +
                 Pain_def +
                 (1|Site),
             data = thoracic,
             family = binomial())
## Compare models
anova(null_mod, thrx_mod, test = 'LRT')
## Data: thoracic
## Models:
## null_mod: Pain_present ~ 1 + (1 | Site)
## thrx_mod: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
                         BIC logLik deviance Chisq Df Pr(>Chisq)
           npar AIC
              2 392.73 401.13 -194.36
                                       388.73
## null_mod
                                       384.75 3.9765 4
              6 396.75 421.95 -192.38
                                                            0.4092
## thrx mod
## No significant difference between null and full models
## Print model
summary(thrx_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
     Data: thoracic
##
##
       AIC
                BIC
                      logLik deviance df.resid
     396.7
##
              422.0
                     -192.4
                               384.7
                                          487
##
## Scaled residuals:
      Min
              1Q Median
                              3Q
## -0.5846 -0.4324 -0.3798 -0.2759 3.8233
##
## Random effects:
## Groups Name
                      Variance Std.Dev.
## Site
          (Intercept) 0.05601 0.2367
## Number of obs: 493, groups: Site, 5
## Fixed effects:
                  Estimate Std. Error z value
                                                     Pr(>|z|)
                  ## (Intercept)
## Age
                  -0.01341
                             0.15894 -0.084
                                                        0.933
                             0.47408 - 1.398
## SexMale
                  -0.66287
                                                        0.162
## CD4_recent
                   0.14524
                             0.14100 1.030
                                                       0.303
## Pain_defChronic -0.34346
                             0.42404 -0.810
                                                       0.418
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
                           SexMal CD4_rc
##
              (Intr) Age
## Age
               0.281
## SexMale
              -0.172 -0.101
## CD4 recent 0.055 -0.079 0.067
```

Pan_dfChrnc -0.602 -0.335 -0.108 -0.264

```
# Lumbosacral spine
lumbar <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Lumbosacral_spine') %>%
   filter(complete.cases(.))
## Null
null_mod <- glmer(Pain_present ~ 1 + (1|Site),</pre>
              data = lumbar,
              family = binomial())
## Full model
lmbr_mod <- glmer(Pain_present ~</pre>
                  Age +
                  Sex +
                  CD4 recent +
                  Pain_def +
                  (1|Site),
              data = lumbar,
              family = binomial())
## Compare models
anova(null_mod, lmbr_mod, test = 'LRT')
## Data: lumbar
## Models:
## null_mod: Pain_present ~ 1 + (1 | Site)
## lmbr mod: Pain present ~ Age + Sex + CD4 recent + Pain def + (1 | Site)
           npar AIC BIC logLik deviance Chisq Df Pr(>Chisq)
## null mod
            2 451.68 460.08 -223.84
                                         447.68
              6 456.11 481.32 -222.06
## lmbr_mod
                                        444.11 3.567 4
                                                             0.4678
## No significant difference between null and full models
## Print model
summary(lmbr_mod)
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial ( logit )
## Formula: Pain_present ~ Age + Sex + CD4_recent + Pain_def + (1 | Site)
##
     Data: lumbar
##
##
       AIC
                BIC logLik deviance df.resid
              481.3 -222.1
##
      456.1
                                 444.1
##
## Scaled residuals:
      Min
              1Q Median
                                3Q
## -0.7060 -0.4701 -0.4167 -0.3660 2.8857
## Random effects:
                      Variance Std.Dev.
## Groups Name
         (Intercept) 0.04987 0.2233
## Number of obs: 493, groups: Site, 5
## Fixed effects:
```

```
##
                   Estimate Std. Error z value
                                                       Pr(>|z|)
## (Intercept)
                   -1.59777
                               0.22763 -7.019 0.00000000000223 ***
                    0.02967
                               0.14127
                                         0.210
## Age
                                                         0.8336
                                                         0.9242
## SexMale
                    0.03293
                               0.34633
                                         0.095
## CD4 recent
                    0.22649
                               0.12388
                                         1.828
                                                         0.0675
## Pain defChronic -0.05838
                                                         0.8774
                               0.37842 - 0.154
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
               (Intr) Age
                             SexMal CD4_rc
                0.250
## Age
## SexMale
               -0.199 -0.130
                0.009 -0.108 0.134
## CD4_recent
## Pan_dfChrnc -0.609 -0.335 -0.147 -0.232
```

Body sites with models that generated an "is singular" error included:

- Throat
- Arms
- Elbows
- Wrists & hands
- Abdomen
- Lower back (flanks)
- Buttocks
- Legs
- Knees
- Ankles & feet

All these models had a random error variance of 0 (or tending to 0), indicating that **site** was not contributing to the model and could be removed. That is, there is not enough additional site-level variation to warrant adding a random effect to explain all the observed variation. Analyzed these using glm, not glmm, and check for issues.

8.4 Run glm for selected body sites

Note: p-values reported for the likelihood ratio tests below are uncorrected for multiple comparisons.

```
Sex +
               CD4_recent
               Pain_def,
             data = throat,
             family = binomial())
## Compare models
anova(null_mod, thrt_mod, test = 'LRT')
## Analysis of Deviance Table
##
## Model 1: Pain_present ~ 1
## Model 2: Pain_present ~ Age + Sex + CD4_recent + Pain_def
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
          492
                  120.18
          488
                                      0.1257
## 2
                  112.98 4 7.1991
## No significant difference between the null and full models
## Print model
summary(thrt_mod)
##
## Call:
## glm(formula = Pain_present ~ Age + Sex + CD4_recent + Pain_def,
##
      family = binomial(), data = throat)
##
## Deviance Residuals:
      Min
                1Q
                    Median
                                 3Q
                                         Max
## -0.5014 -0.2524 -0.2063 -0.1542
                                      2.9367
##
## Coefficients:
                  Estimate Std. Error z value
                                                       Pr(>|z|)
## (Intercept)
                  ## Age
                  0.07201
                             0.33111 0.217
                                                         0.8278
## SexMale
                   0.91911
                             0.62751
                                     1.465
                                                         0.1430
## CD4_recent
                  -0.78730
                             0.45238 -1.740
                                                         0.0818 .
## Pain defChronic 0.19658
                             0.68880
                                      0.285
                                                         0.7753
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 120.18 on 492 degrees of freedom
## Residual deviance: 112.98 on 488 degrees of freedom
## AIC: 122.98
## Number of Fisher Scoring iterations: 7
#-- Arms --#
arms <- data_logit.long %>%
   # Filter by pain site
   filter(Pain_site == 'Arms') %>%
   filter(complete.cases(.))
## Null
```

```
null_mod <- glm(Pain_present ~ 1,</pre>
                data = arms,
                family = binomial())
## Full model
arms_mod <- glm(Pain_present ~
                Age +
                Sex +
                CD4 recent +
                Pain_def,
              data = arms,
              family = binomial())
## Compare models
anova(null_mod, arms_mod, test = 'LRT')
## Analysis of Deviance Table
## Model 1: Pain_present ~ 1
## Model 2: Pain_present ~ Age + Sex + CD4_recent + Pain_def
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
           492
                   179.82
## 2
           488
                   177.13 4
                                2.686
                                        0.6117
## No significant difference between the null and full models
## Print model
summary(arms_mod)
##
## Call:
## glm(formula = Pain_present ~ Age + Sex + CD4_recent + Pain_def,
##
      family = binomial(), data = arms)
##
## Deviance Residuals:
      Min
                1Q
                     Median
                                   3Q
                                           Max
## -0.5876 -0.3257 -0.2959 -0.2574
                                        2.6283
##
## Coefficients:
                   Estimate Std. Error z value
                                                          Pr(>|z|)
                              0.27466 -10.535 <0.0000000000000000 ***
## (Intercept)
                   -2.89348
## Age
                   0.30866
                               0.21783
                                       1.417
                                                             0.156
## SexMale
                   -0.05672
                               0.59049 -0.096
                                                             0.923
                               0.25597 -0.393
## CD4_recent
                   -0.10059
                                                             0.694
## Pain_defChronic -0.62336
                               0.59010 -1.056
                                                             0.291
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 179.82 on 492 degrees of freedom
## Residual deviance: 177.13 on 488 degrees of freedom
## AIC: 187.13
## Number of Fisher Scoring iterations: 6
```

```
#-- Elbow --#
elbows <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Elbows') %>%
   filter(complete.cases(.))
## Null
null_mod <- glm(Pain_present ~ 1,</pre>
                data = elbows,
                family = binomial())
## Full model
elbw_mod <- glm(Pain_present ~
                Age +
                Sex +
                CD4_recent +
                Pain_def,
              data = elbows,
              family = binomial())
## Compare models
anova(null_mod, elbw_mod, test = 'LRT')
## Analysis of Deviance Table
## Model 1: Pain_present ~ 1
## Model 2: Pain_present ~ Age + Sex + CD4_recent + Pain_def
## Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
           492
                  167.37
## 2
           488
                   160.13 4 7.2361
                                        0.1239
## No significant difference between the null and full models
## Print model
summary(elbw_mod)
##
## glm(formula = Pain_present ~ Age + Sex + CD4_recent + Pain_def,
       family = binomial(), data = elbows)
##
## Deviance Residuals:
      Min
                1Q
                     Median
                                   3Q
                                           Max
## -0.6164 -0.2974 -0.2476 -0.2156
                                        2.7664
##
## Coefficients:
                                                          Pr(>|z|)
##
                   Estimate Std. Error z value
## (Intercept)
                   -3.38921
                              0.33556 -10.100 <0.0000000000000000 ***
## Age
                   0.42521
                               0.21018
                                        2.023
                                                            0.0431 *
## SexMale
                   -0.16645
                               0.59159 -0.281
                                                            0.7784
## CD4 recent
                    0.04229
                               0.23571
                                       0.179
                                                            0.8576
## Pain_defChronic 0.45074
                               0.53643
                                       0.840
                                                            0.4008
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
```

```
##
       Null deviance: 167.37 on 492 degrees of freedom
## Residual deviance: 160.13 on 488 degrees of freedom
## AIC: 170.13
## Number of Fisher Scoring iterations: 6
#-- Wrist & hand --#
wrists <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Wrists.Hands') %>%
   filter(complete.cases(.))
## Null
null_mod <- glm(Pain_present ~ 1,</pre>
                data = wrists,
                family = binomial())
## Full model
wrst_mod <- glm(Pain_present ~</pre>
                Age +
                Sex +
                CD4_recent +
                Pain_def,
              data = wrists,
              family = binomial())
## Compare models
anova(null_mod, wrst_mod, test = 'LRT')
## Analysis of Deviance Table
##
## Model 1: Pain_present ~ 1
## Model 2: Pain_present ~ Age + Sex + CD4_recent + Pain_def
   Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          492
                  242.2
                    231.3 4 10.908 0.02762 *
## 2
           488
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## No significant difference between the null and full models
## (family-wise error correction, alpha = 0.0024)
## Print model
summary(wrst_mod)
##
## glm(formula = Pain_present ~ Age + Sex + CD4_recent + Pain_def,
##
       family = binomial(), data = wrists)
##
## Deviance Residuals:
       Min
                1Q
                     Median
                                   3Q
                                           Max
## -0.8224 -0.3799 -0.3189 -0.2853
                                        2.7809
##
## Coefficients:
                   Estimate Std. Error z value
                                                          Pr(>|z|)
##
```

```
## (Intercept)
                   -2.7754
                               0.2588 -10.723 <0.0000000000000000 ***
## Age
                                       1.284
                               0.1804
                    0.2316
                                                            0.199
                   -0.6507
                                                            0.251
## SexMale
                               0.5662 - 1.149
## CD4_recent
                    0.2766
                               0.1711
                                        1.616
                                                            0.106
## Pain_defChronic 0.4408
                               0.4376
                                       1.007
                                                             0.314
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 242.2 on 492 degrees of freedom
## Residual deviance: 231.3 on 488 degrees of freedom
## AIC: 241.3
##
## Number of Fisher Scoring iterations: 5
#-- Abdomen --#
abdomen <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Abdomen') %>%
   filter(complete.cases(.))
## Null
null_mod <- glm(Pain_present ~ 1,</pre>
               data = abdomen,
                family = binomial())
## Full model
abdm_mod <- glm(Pain_present ~
               Age +
               Sex +
               CD4_recent +
               Pain_def,
              data = abdomen,
             family = binomial())
## Compare models
anova(null_mod, abdm_mod, test = 'LRT')
## Analysis of Deviance Table
## Model 1: Pain_present ~ 1
## Model 2: Pain_present ~ Age + Sex + CD4_recent + Pain_def
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          492
                  578.82
                              9.6645 0.04648 *
## 2
          488
                  569.15 4
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## No significant difference between the null and full models
## (family-wise error correction, alpha = 0.0024)
## Print model
summary(abdm_mod)
##
## Call:
```

```
## glm(formula = Pain_present ~ Age + Sex + CD4_recent + Pain_def,
##
       family = binomial(), data = abdomen)
##
## Deviance Residuals:
      Min
                1Q
                    Median
                                   3Q
                                           Max
## -0.9950 -0.8601 -0.7247
                                        1.9325
                               1.4630
## Coefficients:
##
                   Estimate Std. Error z value
                                                     Pr(>|z|)
## (Intercept)
                   -0.8813
                                0.1365 -6.458 0.00000000106 ***
## Age
                   -0.1613
                                0.1219 -1.323
                                                        0.186
                                0.3015 -1.019
## SexMale
                    -0.3072
                                                        0.308
                    -0.1145
## CD4_recent
                                0.1154 -0.993
                                                        0.321
                                0.2706 -0.940
## Pain_defChronic -0.2542
                                                        0.347
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 578.82 on 492 degrees of freedom
## Residual deviance: 569.15 on 488 degrees of freedom
## AIC: 579.15
##
## Number of Fisher Scoring iterations: 4
#-- Lower back (flank) --#
low_back <- data_logit.long %>%
    # Filter by pain site
    filter(Pain_site == 'Lower_back') %>%
   filter(complete.cases(.))
## Null
null_mod <- glm(Pain_present ~ 1,</pre>
                data = low_back,
                family = binomial())
## Full model
back_mod <- glm(Pain_present ~</pre>
                Age +
                Sex +
                CD4 recent +
                Pain_def,
              data = low_back,
              family = binomial())
## Compare models
anova(null_mod, back_mod, test = 'LRT')
## Analysis of Deviance Table
##
## Model 1: Pain present ~ 1
## Model 2: Pain_present ~ Age + Sex + CD4_recent + Pain_def
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
           492
                   262.78
## 2
           488
                   260.15 4 2.6328
                                         0.621
```

```
## No significant difference between the null and full models
## Print model
summary(back_mod)
##
## Call:
## glm(formula = Pain_present ~ Age + Sex + CD4_recent + Pain_def,
##
      family = binomial(), data = low_back)
##
## Deviance Residuals:
            10 Median
      Min
                                   30
                                           Max
## -0.5466 -0.4105 -0.3690 -0.3520
                                        2.4494
## Coefficients:
##
                   Estimate Std. Error z value
                                                          Pr(>|z|)
                            0.23601 -11.001 <0.00000000000000000 ***
## (Intercept)
                   -2.59625
## Age
                   0.10656
                               0.18068
                                        0.590
                                                             0.555
## SexMale
                   -0.23426
                               0.48221 -0.486
                                                             0.627
## CD4_recent
                    0.08667
                               0.17452
                                        0.497
                                                             0.619
## Pain_defChronic 0.33885
                               0.41624
                                         0.814
                                                             0.416
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 262.78 on 492 degrees of freedom
## Residual deviance: 260.15 on 488 degrees of freedom
## AIC: 270.15
## Number of Fisher Scoring iterations: 5
#-- Buttocks --#
buttocks <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Buttocks') %>%
   filter(complete.cases(.))
## Null
null_mod <- glm(Pain_present ~ 1,</pre>
                data = buttocks,
                family = binomial())
## Full model
butt_mod <- glm(Pain_present ~</pre>
                Age +
                Sex +
                CD4_recent +
                Pain_def,
              data = buttocks,
              family = binomial())
## Compare models
anova(null_mod, butt_mod, test = 'LRT')
```

Analysis of Deviance Table

```
##
## Model 1: Pain_present ~ 1
## Model 2: Pain_present ~ Age + Sex + CD4_recent + Pain_def
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
           492
                   160.99
## 2
           488
                   158.73 4
                               2.2631
                                       0.6875
\#\# No significant difference between the null and full models
## Print model
summary(butt_mod)
## Call:
## glm(formula = Pain_present ~ Age + Sex + CD4_recent + Pain_def,
##
       family = binomial(), data = buttocks)
##
## Deviance Residuals:
       Min
                1Q
                     Median
                                   3Q
                                           Max
## -0.4682 -0.3007 -0.2608 -0.2401
                                        2.7090
##
## Coefficients:
                   Estimate Std. Error z value
##
                                                          Pr(>|z|)
## (Intercept)
                   -3.42677
                               0.33725 -10.161 < 0.0000000000000000 ***
## Age
                   -0.12939
                               0.26949 -0.480
                                                              0.631
## SexMale
                    0.65435
                               0.57735
                                        1.133
                                                              0.257
                                                              0.299
## CD4_recent
                    0.23689
                               0.22789
                                       1.039
## Pain_defChronic 0.09589
                               0.57052
                                        0.168
                                                              0.867
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 160.99 on 492 degrees of freedom
## Residual deviance: 158.73 on 488 degrees of freedom
## AIC: 168.73
##
## Number of Fisher Scoring iterations: 6
#-- Legs --#
legs <- data_logit.long %>%
   # Filter by pain site
   filter(Pain_site == 'Legs') %>%
   filter(complete.cases(.))
## Null
null_mod <- glm(Pain_present ~ 1,</pre>
                data = legs,
                family = binomial())
## Full model
legs_mod <- glm(Pain_present ~</pre>
                Age +
                Sex +
                CD4_recent +
                Pain_def,
```

```
data = legs,
              family = binomial())
## Compare models
anova(null_mod, legs_mod, test = 'LRT')
## Analysis of Deviance Table
##
## Model 1: Pain_present ~ 1
## Model 2: Pain_present ~ Age + Sex + CD4_recent + Pain_def
   Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
          492
                  440.48
## 2
          488
                  419.02 4
                              21.454 0.0002574 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Significant difference between the null and full models
## (family-wise error correction, alpha = 0.0024)
summary(legs_mod)
##
## Call:
## glm(formula = Pain_present ~ Age + Sex + CD4_recent + Pain_def,
      family = binomial(), data = legs)
## Deviance Residuals:
                     Median
      Min
                1Q
                                   3Q
                                           Max
## -1.0034 -0.6258 -0.4805 -0.4483
                                        2.1998
##
## Coefficients:
                  Estimate Std. Error z value
                                                          Pr(>|z|)
                               0.1812 -11.170 <0.0000000000000000 ***
## (Intercept)
                   -2.0238
## Age
                                0.1305 0.856
                     0.1117
                                                            0.3920
                     0.6088
                                0.3013 2.020
                                                            0.0433 *
## SexMale
## CD4_recent
                     0.1644
                                0.1252
                                       1.313
                                                            0.1892
## Pain_defChronic 0.6567
                                0.2903
                                       2.262
                                                            0.0237 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 440.48 on 492 degrees of freedom
## Residual deviance: 419.02 on 488 degrees of freedom
## AIC: 429.02
##
## Number of Fisher Scoring iterations: 4
#-- Knees --#
knees <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Knees') %>%
   filter(complete.cases(.))
## Null
null_mod <- glm(Pain_present ~ 1,</pre>
```

```
data = knees,
                family = binomial())
## Full model
knee_mod <- glm(Pain_present ~</pre>
                Age +
                Sex +
               CD4_recent +
               Pain_def,
              data = knees,
             family = binomial())
## Compare models
anova(null_mod, knee_mod, test = 'LRT')
## Analysis of Deviance Table
## Model 1: Pain_present ~ 1
## Model 2: Pain_present ~ Age + Sex + CD4_recent + Pain_def
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
                  413.48
          492
## 2
          488
                  388.18 4
                              25.298 0.00004382 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Significant difference between the null and full models
## (family-wise error correction, alpha = 0.0024)
summary(knee_mod)
##
## Call:
## glm(formula = Pain_present ~ Age + Sex + CD4_recent + Pain_def,
      family = binomial(), data = knees)
##
## Deviance Residuals:
           1Q Median
      Min
                                  3Q
                                          Max
## -1.0315 -0.5925 -0.4460 -0.4038
                                       2.3429
##
## Coefficients:
##
                  Estimate Std. Error z value
                                                          Pr(>|z|)
## (Intercept)
                   -2.1860
                               0.1932 -11.314 < 0.0000000000000000 ***
## Age
                    0.2308
                               0.1323 1.745
                                                           0.08100 .
## SexMale
                    0.4132
                               0.3147 1.313
                                                           0.18928
                                       0.718
                                                           0.47273
## CD4_recent
                    0.0948
                               0.1320
## Pain_defChronic 0.8155
                               0.3015
                                       2.705
                                                           0.00683 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 413.48 on 492 degrees of freedom
## Residual deviance: 388.18 on 488 degrees of freedom
## AIC: 398.18
##
## Number of Fisher Scoring iterations: 5
```

```
#-- Ankles/feet --#
ankles <- data_logit.long %>%
    # Filter by pain site
   filter(Pain_site == 'Ankles.Feet') %>%
   filter(complete.cases(.))
## Null
null_mod <- glm(Pain_present ~ 1,</pre>
               data = ankles,
               family = binomial())
## Full model
ankl_mod <- glm(Pain_present ~
               Age +
               Sex +
               CD4_recent +
               Pain_def,
             data = ankles,
             family = binomial())
## Compare models
anova(null_mod, ankl_mod, test = 'LRT')
## Analysis of Deviance Table
##
## Model 1: Pain_present ~ 1
## Model 2: Pain_present ~ Age + Sex + CD4_recent + Pain_def
## Resid. Df Resid. Dev Df Deviance
                                         Pr(>Chi)
                  600.78
## 1
          492
## 2
          488
                  558.18 4 42.596 0.00000001255 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Significant difference between the null and full models
## (family-wise error correction, alpha = 0.0024)
summary(ankl_mod)
##
## Call:
## glm(formula = Pain_present ~ Age + Sex + CD4_recent + Pain_def,
##
      family = binomial(), data = ankles)
##
## Deviance Residuals:
      Min
                1Q
                    Median
                                 3Q
                                         Max
## -1.4967 -0.7619 -0.6565
                            1.1107
                                      1.9524
## Coefficients:
                  Estimate Std. Error z value
                                                        Pr(>|z|)
                  ## (Intercept)
                             0.11072 2.473
## Age
                  0.27385
                                                          0.0134 *
                                     1.465
## SexMale
                             0.26289
                  0.38509
                                                          0.1430
## CD4_recent
                  -0.08338
                             0.11100 -0.751
                                                          0.4526
                                                      0.0000831 ***
## Pain_defChronic 0.96276
                             0.24465 3.935
## ---
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
## Null deviance: 600.78 on 492 degrees of freedom
## Residual deviance: 558.18 on 488 degrees of freedom
## AIC: 568.18
##
## Number of Fisher Scoring iterations: 4
```

8.5 GLMM data for tabulation and plotting

Perform GLMM on selected body sites (head, shoulders, chest, hips, cervical spine, thoracic spine, and lumbosacral spine) to extract fixed effect odds ratios (with 95% CIs) to tabulate and to produce forest plots.

```
# Perform GLMM on each selected body sites
glmm_mods <- data_logit.long %>%
    filter(complete.cases(.)) %>%
    # remove body sites with glmm singular fits
    filter(!Pain_site %in% c("Throat", "Arms", "Elbows", "Lower_back",
                             "Wrists. Hands", "Abdomen", "Buttocks",
                             "Legs", "Knees", "Ankles.Feet")) %>%
    # Group and then nest by Pain_site
    group_by(Pain_site) %>%
   nest() %>%
    # Perform logistic regression
   mutate(logistic = map(.x = data,
                          ~ glmer(Pain_present ~
                                       Age +
                                       Sex +
                                      CD4_recent +
                                      Pain_def +
                                       (1|Site),
                                  data = .x,
                                  family = binomial())),
           null = map(.x = data,
                      ~glmer(Pain_present ~ 1 +
                                  (1|Site),
                             data = .x,
                             family = binomial()))) %>%
    # Perform likelihood ratio test
   mutate(LRT = map2(.x = logistic,
                      .y = null,
                      ~ anova(.y, .x, test = 'LRT'))) %>%
    # Extract LRT p-values and correct for multiple comparisons
   mutate(LRT_p = map(.x = LRT,
                       ~ .x[, 8][[2]]),
           LRT_p.corrected = map(.x = LRT_p,
                                  ~ p.adjust(p = .x,
                                             method = 'bonferroni',
                                             n = 21))) %>%
    # Extract effect sizes
    mutate(OR = map(.x = logistic,
                    ~ exp(fixef(.x))),
           CI = map(.x = logistic,
```

```
~ exp(confint.merMod(.x,
                                           method = 'Wald')[2:6, ]))) %>%
ungroup() %>%
mutate(Pain_site = str_replace(Pain_site,
                               pattern = '_',
                               replacement = ' ')) %>%
mutate(Pain_site = str_replace(Pain_site,
                               pattern = '\\.',
                               replacement = ' & ')) %>%
# Bind data into a dataframe and tabulate,
mutate(df = pmap(.1 = list(OR, CI, LRT_p.corrected, Pain_site),
                 ~ cbind(..1, ..2) %>%
                     .[2:5, ] %>%
                     kable(caption = str_glue('{...4}) (GLMM; LRT corrected p-value = {round(...3, 3)}
                           digits = 3,
                           col.names = c('OR', 'Wald lower 95%CI',
                                          'Wald upper 95%CI')))) %>%
# Generate forest plot data
mutate(forest_data = map2(.x = OR,
                          y = CI,
                          # Create the dataframe
                          ~ as.data.frame(cbind(.x, .y)) %>%
                              set_names(nm = c('OR', 'lower_ci',
                                                'upper_ci')) %>%
                              .[-1,] %>%
                              rownames_to_column() %>%
                              mutate(rowname = case_when(
                                  rowname == 'Age' ~ 'Age',
                                  rowname == 'SexMale' ~ 'Sex (male)',
                                  rowname == 'CD4_recent' ~ 'CD4 T-cells',
                                  rowname == 'Pain_defChronic' ~ 'Pain (chronic)'
                                  )) %>%
                              # Round digits to 2 decimal places
                              mutate_if(is.numeric, round, 2) %>%
                              # Paste CI range
                              mutate(ci_range = paste0('(', lower_ci, ' to ',
                                                        upper_ci, ')')) %>%
                              # Cut short confidence interval for plotting
                              mutate(upper_ci = ifelse(upper_ci > 4,
                                                        yes = 4,
                                                        no = upper_ci)))) %>%
# Generate forest plots
mutate(forest_plots = pmap(.1 = list(forest_data, Pain_site, LRT_p.corrected),
                          ~ ..1 %>%
                              ggplot(data = .) +
                              aes(x = OR,
                                  xmin = lower_ci,
                                  xmax = upper_ci,
                                  y = rowname,
                                  label = ci_range) +
                              geom_pointrange(size = 1) +
                              geom_vline(xintercept = 1,
                                         linetype = 2) +
```

```
scale_x_continuous(limits = c(-0.5, 8),
                   breaks = c(0, 1, 2, 3, 4)) +
geom_text(x = 4.5, hjust = 0, size = 5) +
annotate(geom = 'text',
         label = '95% CI',
         size = 5,
         fontface = 2,
         x = 4.5
         y = 4.4,
        hjust = 0) +
labs(title = ..2,
     subtitle = str_glue('(GLMM | LRT corrected p-value = {round(..3
     x = 'Odds ratio') +
theme_minimal(base_size = 18) +
theme(plot.title = element_text(size = 18),
     plot.subtitle = element_text(size = 14),
     panel.grid = element_blank(),
     axis.title.y = element_blank(),
     axis.text = element_text(colour = '#000000'),
      axis.line = element_line(size = 0.5),
      axis.ticks = element_line(size = 0.5))))
```

8.6 GLM data for tabulation and plotting

Perform GLM on selected body sites (throat, arms, elbows, wrists and hands, abdomen, buttocks, legs, knees, and ankles and feet) to extract fixed effect odds ratios (with 95% CIs) to tabulate and to produce forest plots.

```
# Perform GLM on each selected body sites
glm_mods <- data_logit.long %>%
    filter(complete.cases(.)) %>%
    # Retain body sites with glmm singular fits
   filter(Pain_site %in% c("Throat", "Arms", "Elbows", "Lower_back",
                            "Wrists. Hands", "Abdomen", "Buttocks",
                            "Legs", "Knees", "Ankles.Feet")) %>%
    # Change lower_back to flank
   mutate(Pain_site = ifelse(Pain_site == 'Lower_back',
                              yes = 'Lower back/flanks',
                              no = Pain_site)) %>%
    # Group and then nest by Pain_site
   group_by(Pain_site) %>%
   nest() %>%
    # Perform logistic regression
   mutate(logistic = map(.x = data,
                          ~ glm(Pain_present ~
                                    Age +
                                    Sex +
                                    CD4 recent +
                                    Pain def,
                                data = .x,
                                family = binomial())),
           null = map(.x = data,
                      ~glm(Pain_present ~ 1,
                             data = .x,
```

```
family = binomial()))) %>%
# Perform likelihood ratio test
mutate(LRT = map2(.x = logistic,
                  .y = null,
                  ~ anova(.y, .x, test = 'LRT'))) %>%
# Extract LRT p-values and correct for multiple comparisons
mutate(LRT_p = map(.x = LRT,
                   -x[, 5][[2]]),
       LRT_p.corrected = map(.x = LRT_p,
                             - p.adjust(p = .x,
                                        method = 'bonferroni',
                                        n = 21))) %>%
# Extract effect sizes
mutate(OR = map(.x = logistic,
                ~ exp(coef(.x))),
       CI = map(.x = logistic,
                      ~ exp(confint.default(.x)))) %>%
ungroup() %>%
mutate(Pain_site = str_replace(Pain_site,
                               pattern = '_',
                               replacement = ' ')) %>%
mutate(Pain_site = str_replace(Pain_site,
                               pattern = '\\.',
                               replacement = ' & ')) %>%
# Bind data into a dataframe and tabulate,
mutate(df = pmap(.1 = list(OR, CI, LRT_p.corrected, Pain_site),
                 ~ cbind(..1, ..2) %>%
                     .[2:5, ] %>%
                     kable(caption = str_glue('{..4} (GLM; LRT corrected p-value = {round(..3, 3)})
                           digits = 3,
                           col.names = c('OR', 'Wald lower 95%CI',
                                         'Wald upper 95%CI')))) %>%
# Generate forest plot data
mutate(forest_data = map2(.x = OR,
                          y = CI,
                          # Create the dataframe
                          ~ as.data.frame(cbind(.x, .y)) %>%
                              set_names(nm = c('OR', 'lower_ci',
                                               'upper_ci')) %>%
                              .[-1,] %>%
                              rownames_to_column() %>%
                              mutate(rowname = case_when(
                                  rowname == 'Age' ~ 'Age',
                                  rowname == 'SexMale' ~ 'Sex (male)',
                                  rowname == 'CD4_recent' ~ 'CD4 T-cells',
                                  rowname == 'Pain_defChronic' ~ 'Pain (chronic)'
                                  )) %>%
                              # Round digits to 2 decimal places
                              mutate_if(is.numeric, round, 2) %>%
                              # Paste CI range
                              mutate(ci_range = paste0('(', lower_ci, ' to ',
                                                       upper_ci, ')')) %>%
                              # Cut short confidence interval for plotting
```

```
mutate(upper_ci = ifelse(upper_ci > 4,
                                                        yes = 4,
                                                        no = upper_ci)))) %>%
# Generate forest plots
mutate(forest_plots = pmap(.1 = list(forest_data, Pain_site, LRT_p.corrected),
                          ~ ..1 %>%
                              ggplot(data = .) +
                              aes(x = OR,
                                  xmin = lower_ci,
                                  xmax = upper_ci,
                                  y = rowname,
                                  label = ci_range) +
                              geom pointrange(size = 1) +
                              geom_vline(xintercept = 1,
                                         linetype = 2) +
                              scale_x_continuous(limits = c(-0.5, 8),
                                                 breaks = c(0, 1, 2, 3, 4)) +
                              geom_text(x = 4.5, hjust = 0, size = 5) +
                              annotate(geom = 'text',
                                       label = '95% CI',
                                       size = 5,
                                       fontface = 2,
                                       x = 4.5,
                                       y = 4.4,
                                       hjust = 0) +
                              labs(title = ...2,
                                   subtitle = str_glue('(GLM | LRT corrected p-value = {round(...3,
                                   x = 'Odds ratio') +
                              theme_minimal(base_size = 18) +
                              theme(plot.title = element_text(size = 18),
                                    plot.subtitle = element_text(size = 14),
                                    panel.grid = element_blank(),
                                    axis.title.y = element_blank(),
                                    axis.text = element_text(colour = '#000000'),
                                    axis.line = element_line(size = 0.5),
                                    axis.ticks = element_line(size = 0.5))))
```

8.7 Tabulated results (fixed effects)

8.7.1 GLMM tabulations

```
# Print OR for fixed effects with 95% CIs and p-values
walk(glmm_mods$df, ~print(.x))
```

Table 61: Head (GLMM; LRT corrected p-value = 0.036)

	OR	Wald lower $95\%CI$	Wald upper 95%CI
Age	1.036	0.815	1.317
SexMale	0.260	0.122	0.553
CD4_recent	0.895	0.714	1.121
Pain_defChronic	0.577	0.249	1.340

Table 62: Shoulder (GLMM; LRT corrected p-value = 1)

	OR	Wald lower 95%CI	Wald upper 95%CI
Age	0.861	0.578	1.282
SexMale	1.703	0.733	3.957
$CD4$ _recent	0.941	0.651	1.360
Pain_defChronic	1.111	0.451	2.739

Table 63: Chest (GLMM; LRT corrected p-value = 1)

	OR	Wald lower 95%CI	Wald upper 95%CI
Age	1.099	0.838	1.442
SexMale	1.044	0.551	1.979
$CD4_recent$	0.757	0.564	1.015
Pain_defChronic	0.470	0.219	1.008

Table 64: Cervical spine (GLMM; LRT corrected p-value = 1)

	OR	Wald lower $95\%CI$	Wald upper 95%CI
Age	1.159	0.690	1.945
SexMale	0.546	0.113	2.640
CD4_recent	1.068	0.666	1.713
$Pain_defChronic$	0.071	0.008	0.622

Table 65: Thoracic spine (GLMM; LRT corrected p-value = 1)

	OR	Wald lower $95\%CI$	Wald upper 95%CI
Age	0.987	0.723	1.347
SexMale	0.515	0.204	1.305
CD4_recent	1.156	0.877	1.524
Pain_defChronic	0.709	0.309	1.628

Table 66: Lumbosacral spine (GLMM; LRT corrected p-value = 1)

	OR	Wald lower 95%CI	Wald upper 95%CI
Age	1.030	0.781	1.359
SexMale	1.033	0.524	2.038
CD4_recent	1.254	0.984	1.599
$Pain_defChronic$	0.943	0.449	1.980

Table 67: Groin (GLMM; LRT corrected p-value = 1)

	OR	Wald lower $95\%CI$	Wald upper 95%CI
Age	0.947	0.680	1.320
SexMale	0.532	0.224	1.264

	OR	Wald lower 95%CI	Wald upper 95%CI
CD4_recent Pain_defChronic	1.229	0.885	1.706
	0.661	0.084	5.217

Table 68: Hips (GLMM; LRT corrected p-value = 1)

	OR	Wald lower 95%CI	Wald upper 95%CI
Age	1.057	0.728	1.536
SexMale	1.373	0.581	3.245
CD4_recent	1.131	0.802	1.593
$Pain_defChronic$	6.570	2.204	19.585

8.7.2 GLM tabulations

Print OR for fixed effects with 95% CIs and p-values
walk(glm_mods\$df, ~print(.x))

Table 69: Throat (GLM; LRT corrected p-value = 1)

	OR	Wald lower $95\%\mathrm{CI}$	Wald upper $95\%CI$
Age	1.075	0.562	2.056
SexMale	2.507	0.733	8.576
CD4_recent	0.455	0.188	1.104
$Pain_defChronic$	1.217	0.316	4.696

Table 70: Arms (GLM; LRT corrected p-value = 1)

	OR	Wald lower 95%CI	Wald upper 95%CI
Age	1.362	0.888	2.087
SexMale	0.945	0.297	3.006
CD4_recent	0.904	0.548	1.493
$Pain_defChronic$	0.536	0.169	1.704

Table 71: Elbows (GLM; LRT corrected p-value = 1)

	OR	Wald lower $95\%\mathrm{CI}$	Wald upper $95\%\mathrm{CI}$
Age	1.530	1.013	2.310
SexMale	0.847	0.266	2.699
CD4_recent	1.043	0.657	1.656
Pain_defChronic	1.569	0.548	4.491

Table 72: Wrists & Hands (GLM; LRT corrected p-value = 0.58)

	OR	Wald lower 95%CI	Wald upper 95%CI
Age	1.261	0.885	1.795

	OR	Wald lower 95%CI	Wald upper 95%CI
SexMale	0.522	0.172	1.583
$CD4$ _recent	1.319	0.943	1.844
$Pain_defChronic$	1.554	0.659	3.663

Table 73: Lower back/flanks (GLM; LRT corrected p-value = 1)

	OR	Wald lower $95\%CI$	Wald upper 95%CI
Age	1.112	0.781	1.585
SexMale	0.791	0.307	2.036
CD4_recent	1.091	0.775	1.535
Pain_defChronic	1.403	0.621	3.173

Table 74: Abdomen (GLM; LRT corrected p-value = 0.976)

	OR	Wald lower 95%CI	Wald upper 95%CI
Age	0.851	0.670	1.081
SexMale	0.736	0.407	1.328
CD4_recent	0.892	0.711	1.118
Pain_defChronic	0.776	0.456	1.318

Table 75: Legs (GLM; LRT corrected p-value = 0.005)

	OR	Wald lower $95\%CI$	Wald upper 95%CI
Age	1.118	0.866	1.444
SexMale	1.838	1.018	3.318
$CD4$ _recent	1.179	0.922	1.506
Pain_defChronic	1.928	1.092	3.407

Table 76: Knees (GLM; LRT corrected p-value = 0.001)

	OR	Wald lower $95\%CI$	Wald upper 95%CI
Age	1.260	0.972	1.632
SexMale	1.512	0.816	2.801
CD4_recent	1.099	0.849	1.424
Pain_defChronic	2.260	1.252	4.081

Table 77: Ankles & Feet (GLM; LRT corrected p-value = 0)

	OR	Wald lower $95\%CI$	Wald upper 95%CI
Age	1.315	1.058	1.634
SexMale	1.470	0.878	2.460
$CD4$ _recent	0.920	0.740	1.144
Pain_defChronic	2.619	1.621	4.230

Table 78: Buttocks (GLM; LRT corrected p-value = 1)

	OR	Wald lower 95%CI	Wald upper 95%CI
Age	0.879	0.518	1.490
SexMale	1.924	0.620	5.965
CD4_recent	1.267	0.811	1.981
Pain_defChronic	1.101	0.360	3.367

8.8 Plotted results (fixed effects)

Includes both GLM and GLMM results.

```
# Extract and process sub-plots from GLMM data
head <- glmm_mods$forest_plots[[1]] +</pre>
    theme(axis.title.x = element_blank())
shoulder <- glmm_mods$forest_plots[[2]] +</pre>
    theme(axis.text.y = element blank(),
           axis.title.x = element_blank())
chest <- glmm_mods$forest_plots[[3]] +</pre>
    theme(axis.text.y = element_blank(),
           axis.title.x = element_blank())
cervical <- glmm_mods$forest_plots[[4]] +</pre>
    theme(axis.text.y = element_blank())
thoracic<- glmm_mods$forest_plots[[5]]</pre>
lumbar <- glmm_mods$forest_plots[[6]] +</pre>
    theme(axis.text.y = element_blank())
groin <- glmm_mods$forest_plots[[7]] +</pre>
    theme(axis.text.y = element_blank(),
           axis.title.x = element blank())
hips <- glmm_mods$forest_plots[[8]] +</pre>
    theme(axis.text.y = element_blank(),
           axis.title.x = element_blank())
# Extract and process sub-plots from GLM data
throat <- glm_mods$forest_plots[[1]] +</pre>
    theme(axis.title.x = element_blank(),
           axis.text.y = element_blank())
arms <- glm_mods$forest_plots[[2]] +</pre>
    theme(axis.text.y = element_blank(),
           axis.title.x = element blank())
elbows <- glm_mods$forest_plots[[3]] +</pre>
    theme(axis.title.x = element_blank())
wrists.hands <- glm_mods$forest_plots[[4]] +</pre>
```

```
theme(axis.text.y = element_blank(),
          axis.title.x = element_blank())
flank <- glm_mods$forest_plots[[5]] +</pre>
    theme(axis.title.x = element_blank())
abdomen <- glm_mods$forest_plots[[6]] +</pre>
    theme(axis.title.x = element blank(),
          axis.text.y = element_blank())
legs <- glm_mods$forest_plots[[7]] +</pre>
    theme(axis.title.x = element_blank())
knees <- glm_mods$forest_plots[[8]] +</pre>
    theme(axis.text.y = element_blank(),
          axis.title.x = element_blank())
ankles.feet <- glm_mods$forest_plots[[9]] +</pre>
    theme(axis.text.y = element_blank())
buttocks <- glm_mods$forest_plots[[10]] +</pre>
    theme(axis.text.y = element_blank(),
          axis.title.x = element_blank())
# Patchwork
log_patch <- head + throat + chest + abdomen + flank + groin +</pre>
    shoulder + arms + elbows + wrists.hands +
    buttocks + hips + legs + knees + ankles.feet +
    cervical + thoracic + lumbar +
    plot_layout(ncol = 4)
# Save
ggsave(filename = 'figures/figure_2.png',
       plot = log_patch,
       width = 16,
       height = 18)
```

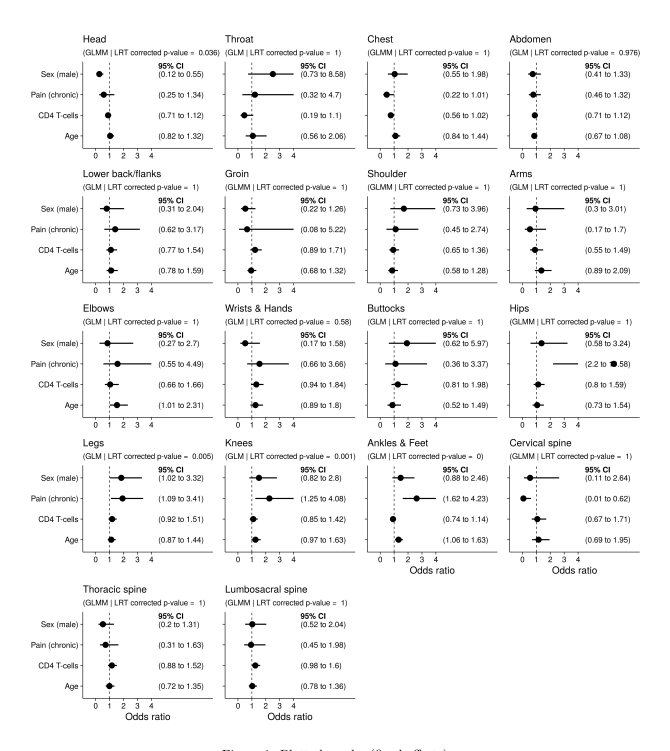


Figure 1: Plotted results (fixed effects)

9 Summary plots

9.1 Body sites with a point estimate pain proportion >10%

```
# Set seed
set.seed(2020)
```

```
# Generate a filter to extract sites with >10% pain prevalence
filter <- data_logit %>%
    select(-Site, -CD4_recent, -Age, -Sex, -Pain_def) %>%
   pivot_longer(cols = everything(),
                 names_to = 'Site',
                 values_to = 'Response') %>%
   group_by(Site, Response) %>%
    summarise(count = n()) %>%
   mutate(total = sum(count),
           prop = count/total) %>%
   filter(Response == 'Yes')
filter_geq10 <- filter(filter, prop >= 0.1) %>%
# Generate new dataset
data_geq10 <- data_logit %>%
   pivot_longer(cols = -c(Site, CD4_recent, Sex, Age, Pain_def),
                 names_to = 'Pain_site',
                 values_to = 'Pain_present') %>%
    # Filter by filter geg10
   filter(Pain_site %in% filter_geq10)
pubs_nofacet <- data_geq10 %>%
    group_by(Pain_site) %>%
   nest() %>%
    # Boostrap data
   mutate(boot = map(.x = data,
                      ~ boot(data = .x[, 'Pain_present'],
                             statistic = prop_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
   mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                           ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
    select(-boot, -ci) %>%
    # Unnest
   unnest(cols = c(data, point_est, lower_ci, upper_ci)) %>%
   ungroup()
plot_nofacet <- pubs_nofacet %>%
    select(-CD4_recent, -Sex, -Age, -Pain_present, -Site, -Pain_def) %>%
   distinct() %>%
```

```
mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '_',
                                   replacement = ' ')) %>%
   mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '\\.',
                                   replacement = ' & ')) %>%
   mutate(Pain_site = factor(Pain_site,
                              levels = c('Head', 'Chest', 'Abdomen',
                                         'Legs', 'Knees', 'Ankles & Feet',
                                         'Thoracic spine', 'Lumbosacral spine'),
                              ordered = TRUE)) %>%
    ggplot(data = .) +
    aes(x = fct rev(Pain site),
       y = point_est,
        ymin = lower_ci,
       ymax = upper_ci) +
    geom_linerange(size = 1) +
    geom_point(size = 5) +
    coord_flip() +
   labs(title = 'Body sites',
         subtitle = '(Point estimate with 95%CI)',
         y = 'Proportion of participants with pain') +
    scale_y_continuous(limits = c(0, 0.6)) +
   theme_minimal(base_size = 18) +
   theme(plot.title = element_text(size = 18),
          plot.subtitle = element_text(size = 12),
          axis.title.y = element blank(),
          panel.grid = element_blank(),
          axis.text = element_text(colour = '#000000'),
          axis.line = element_line(size = 0.5),
          axis.ticks = element_line(size = 0.5))
#-- Body sites by sex --#
pubs_nofacet.sex <- data_geq10 %>%
   group_by(Pain_site, Sex) %>%
   nest() %>%
   # Boostrap data
   mutate(boot = map(.x = data,
                      ~ boot(data = .x[, 'Pain_present'],
                             statistic = prop func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
   mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                          ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
```

```
~ .x$percent[[5]])) %>%
    # Remove columns
    select(-boot, -ci) %>%
    # Unnest
    unnest(cols = c(data, point_est, lower_ci, upper_ci)) %>%
    ungroup()
plot nofacet.sex <- pubs nofacet.sex %>%
    select(-CD4_recent, -Age, -Pain_present, -Site, -Pain_def) %>%
    distinct() %>%
    mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '_',
                                   replacement = ' ')) %>%
    mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '\\.',
                                   replacement = ' & ')) %>%
    mutate(Pain_site = factor(Pain_site,
                              levels = c('Head', 'Chest', 'Abdomen',
                                          'Legs', 'Knees', 'Ankles & Feet',
                                          'Thoracic spine', 'Lumbosacral spine'),
                              ordered = TRUE)) %>%
    ggplot(data = .) +
    aes(x = fct_rev(Pain_site),
        y = point_est,
        ymin = lower_ci,
        ymax = upper_ci,
        fill = Sex) +
    geom_linerange(position = position_dodge2(width = 0.6),
                   size = 1,
                   colour = '#000000') +
    geom_point(shape = 21,
               colour = '#000000',
               position = position_dodge2(width = 0.6),
               size = 4,
               stroke = 1) +
    coord_flip() +
    labs(title = 'Body sites by sex',
         subtitle = '(Point estimate with 95%CI)',
         y = 'Proportion of participants with pain') +
    scale_y_continuous(limits = c(0, 0.6)) +
    scale_fill_manual(values = c('#FFFFFF', '#000000')) +
    theme_minimal(base_size = 18) +
    theme(legend.position = c(0.8, 0.1),
          legend.title = element_blank(),
          plot.title = element_text(size = 18),
          plot.subtitle = element_text(size = 12),
          axis.title.y = element_blank(),
          axis.text.y = element_blank(),
          panel.grid = element_blank(),
          axis.text = element_text(colour = '#000000'),
          axis.line = element_line(size = 0.5),
          axis.ticks = element_line(size = 0.5))
```

```
#-- Body sites by pain chronicity --#
pubs_nofacet.pain_def <- data_geq10 %>%
    group_by(Pain_site, Pain_def) %>%
   nest() %>%
    # Boostrap data
   mutate(boot = map(.x = data,
                      ~ boot(data = .x[, 'Pain_present'],
                             statistic = prop_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
    mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                           ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
   select(-boot, -ci) %>%
    # Unnest
   unnest(cols = c(data, point_est, lower_ci, upper_ci)) %>%
    ungroup()
plot_nofacet.pain_def <- pubs_nofacet.pain_def %>%
    select(-CD4_recent, -Age, -Pain_present, -Site, -Sex) %>%
    distinct() %>%
    mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '_',
                                   replacement = ' ')) %>%
   mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '\\.',
                                   replacement = ' & ')) %>%
   mutate(Pain_site = factor(Pain_site,
                              levels = c('Head', 'Chest', 'Abdomen',
                                          'Legs', 'Knees', 'Ankles & Feet',
                                         'Thoracic spine', 'Lumbosacral spine'),
                              ordered = TRUE)) %>%
    ggplot(data = .) +
    aes(x = fct rev(Pain site),
       y = point_est,
       ymin = lower_ci,
       ymax = upper_ci,
        fill = Pain_def) +
    geom_linerange(position = position_dodge2(width = 0.6),
                   size = 1,
                   colour = '#000000') +
    geom_point(shape = 21,
               colour = '#000000',
```

```
position = position_dodge2(width = 0.6),
               size = 4,
               stroke = 1) +
    coord_flip() +
    labs(title = 'Body sites by pain chronicity',
         subtitle = '(Point estimate with 95%CI)',
         y = 'Proportion of participants with pain') +
    scale y continuous(limits = c(0, 0.6)) +
    scale_fill_manual(values = c('#FFFFFF', '#000000')) +
   theme_minimal(base_size = 18) +
    theme(legend.position = c(0.8, 0.1),
          legend.title = element_blank(),
          plot.title = element_text(size = 18),
          plot.subtitle = element_text(size = 12),
          axis.title.y = element_blank(),
          panel.grid = element_blank(),
          axis.text = element_text(colour = '#000000'),
          axis.line = element_line(size = 0.5),
          axis.ticks = element_line(size = 0.5))
#-- Body sites by age --#
pubs_nofacet.age <- data_geq10 %>%
    select(-CD4_recent, -Site, -Sex, -Pain_def) %>%
    mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '_',
                                   replacement = ' ')) %>%
   mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '\\.',
                                   replacement = ' & ')) %>%
    group_by(Pain_site, Pain_present) %>%
   nest() %>%
    mutate(boot = map(.x = data,
                      ~ boot(data = .x,
                             statistic = median_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
    mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                           ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
    select(-boot, -ci) %>%
    # Unnest
   unnest(cols = c(data, point_est, lower_ci, upper_ci)) %>%
    ungroup() %>%
```

```
mutate(Pain_site = factor(Pain_site,
                              levels = c('Head', 'Chest', 'Abdomen', 'Groin',
                                          'Legs', 'Knees', 'Ankles & Feet',
                                          'Thoracic spine', 'Lumbosacral spine'),
                              ordered = TRUE)) %>%
    select(-Age) %>%
    distinct()
plot_nofacet.age <- pubs_nofacet.age %>%
    ggplot(data = .) +
    aes(x = fct_rev(Pain_site),
        y = point_est,
        ymin = lower_ci,
       ymax = upper_ci,
        fill = Pain_present) +
    geom_linerange(position = position_dodge2(width = 0.6),
                   size = 1,
                   colour = '#000000') +
    geom_point(shape = 21,
               colour = '#000000',
               position = position_dodge2(width = 0.6),
               size = 4,
               stroke = 1) +
    coord_flip() +
   labs(title = 'Body sites by median age',
         subtitle = '(Point estimate with 95%CI)',
         y = 'Age (years)') +
    scale_fill_manual(values = c('#FFFFFF', '#000000'),
                      labels = c('No pain', 'Pain present')) +
    scale_y_continuous(limits = c(31, 45),
                       breaks = c(31, 35, 39, 43)) +
   theme_minimal(base_size = 18) +
   theme(legend.position = c(0.8, 0.95),
          legend.title = element_blank(),
          plot.title = element_text(size = 18),
          plot.subtitle = element_text(size = 12),
          axis.title.y = element_blank(),
          axis.text.y = element_blank(),
          panel.grid = element_blank(),
          axis.text = element_text(colour = '#000000'),
          axis.line = element_line(size = 0.5),
          axis.ticks = element_line(size = 0.5))
#-- Body sites by CD4 --#
pubs_nofacet.cd4 <- data_geq10 %>%
    select(-Age, -Site, -Sex, -Pain_def) %>%
    mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '_',
                                   replacement = ' ')) %>%
   mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '\\.',
                                   replacement = ' & ')) %>%
   group_by(Pain_site, Pain_present) %>%
```

```
nest() %>%
    mutate(boot = map(.x = data,
                      - boot(data = .x,
                             statistic = median_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
   mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                           ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
    select(-boot, -ci) %>%
    # Unnest
   unnest(cols = c(data, point_est, lower_ci, upper_ci)) %>%
   ungroup() %>%
   mutate(Pain_site = factor(Pain_site,
                              levels = c('Head', 'Chest', 'Abdomen',
                                          'Legs', 'Knees', 'Ankles & Feet',
                                          'Thoracic spine', 'Lumbosacral spine'),
                              ordered = TRUE)) %>%
    select(-CD4_recent) %>%
    distinct()
plot_nofacet.cd4 <- pubs_nofacet.cd4 %>%
    ggplot(data = .) +
    aes(x = fct_rev(Pain_site),
        y = point_est,
        ymin = lower_ci,
        ymax = upper_ci,
       fill = Pain_present) +
    geom_linerange(position = position_dodge2(width = 0.6),
                   size = 1,
                   colour = '#000000') +
   geom_point(shape = 21,
               colour = '#000000',
               position = position_dodge2(width = 0.6),
               size = 4,
               stroke = 1) +
    coord_flip() +
   labs(title = 'Body sites by median CD4 T-cell count',
         subtitle = '(Point estimate with 95%CI)',
         y = expression('CD4 T-cell count (cell.mm'^-3*')')) +
    scale_fill_manual(values = c('#FFFFFF', '#000000'),
                      labels = c('No pain', 'Pain present')) +
    scale_y_continuous(limits = c(100, 500),
```

```
breaks = c(100, 200, 300, 400, 500)) +
    theme_minimal(base_size = 18) +
    theme(legend.position = c(0.8, 0.95),
          legend.title = element_blank(),
          plot.title = element_text(size = 18),
          plot.subtitle = element_text(size = 12),
          axis.title.y = element_blank(),
          panel.grid = element blank(),
          axis.text = element_text(colour = '#000000'),
          axis.line = element_line(size = 0.5),
          axis.ticks = element_line(size = 0.5))
# Patchwork
pub_plot <- plot_nofacet + plot_nofacet.sex + plot_nofacet.pain_def +</pre>
   plot_nofacet.age + plot_nofacet.cd4 +
   plot_layout(ncol = 2)
# Save
ggsave(filename = 'figures/figure_supp1.png',
       width = 14,
       height = 18)
```

9.2 Body sites with a point estimate pain proportion <10%

```
# Set seed
set.seed(2020)
# Generate filter
filter_less10 <- filter(filter, prop < 0.1) %>%
    .$Site
# Generate new dataset
data_less10 <- data_logit %>%
    pivot_longer(cols = -c(Site, CD4_recent, Sex, Age, Pain_def),
                 names_to = 'Pain_site',
                 values_to = 'Pain_present') %>%
    # Filter by filter_geq10
   filter(Pain_site %in% filter_less10)
pubs_nofacet2 <- data_less10 %>%
    group_by(Pain_site) %>%
   nest() %>%
    # Boostrap data
   mutate(boot = map(.x = data,
                      ~ boot(data = .x[, 'Pain_present'],
                             statistic = prop_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
   mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
```

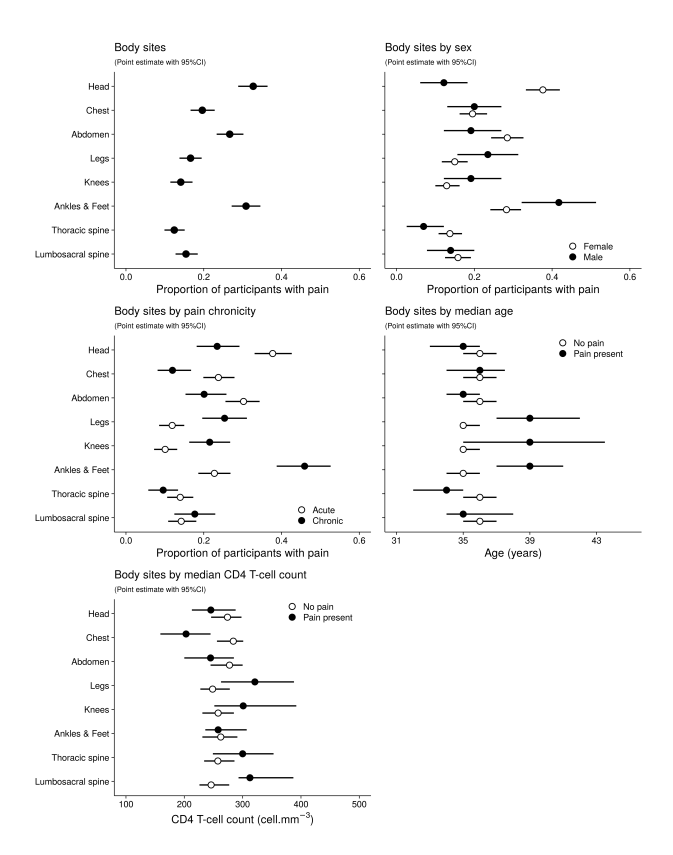


Figure 2: Body sites with a point estimate pain proportion >10%

```
# Extract ci data
   mutate(point_est = map(.x = ci,
                           ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
   select(-boot, -ci) %>%
    # Unnest
   unnest(cols = c(data, point_est, lower_ci, upper_ci)) %>%
    ungroup()
plot_nofacet2 <- pubs_nofacet2 %>%
    select(-CD4_recent, -Sex, -Age, -Pain_present, -Site, -Pain_def) %>%
   distinct() %>%
    mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '_',
                                   replacement = ' ')) %>%
   mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '\\.',
                                   replacement = ' & ')) %>%
   mutate(Pain_site = ifelse(Pain_site == 'Lower back',
                              yes = 'Lower back/flanks',
                              no = Pain_site)) %>%
   mutate(Pain_site = factor(Pain_site,
                              levels = c('Throat', 'Shoulder', 'Arms',
                                          'Elbows', 'Wrists & Hands',
                                          'Lower back/flanks', 'Groin', 'Hips',
                                          'Buttocks', 'Cervical spine'),
                              ordered = TRUE)) %>%
    ggplot(data = .) +
    aes(x = fct_rev(Pain_site),
        y = point_est,
       ymin = lower_ci,
        ymax = upper_ci) +
    geom_linerange(size = 1) +
    geom_point(size = 5) +
   coord_flip() +
   labs(title = 'Body sites',
         subtitle = '(Body sites with <10% pain; Point estimate with 95%CI)',</pre>
         y = 'Proportion of participants with pain') +
   scale y continuous(limits = c(0, 0.3)) +
    theme minimal(base size = 18) +
   theme(plot.title = element_text(size = 18),
          plot.subtitle = element_text(size = 12),
          axis.title.y = element_blank(),
          panel.grid = element_blank(),
          axis.text = element_text(colour = '#000000'),
          axis.line = element_line(size = 0.5),
          axis.ticks = element_line(size = 0.5))
#-- Body sites by sex --#
```

```
pubs_nofacet.sex2 <- data_less10 %>%
    group_by(Pain_site, Sex) %>%
   nest() %>%
    # Boostrap data
   mutate(boot = map(.x = data,
                      ~ boot(data = .x[, 'Pain_present'],
                             statistic = prop_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
   mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                           ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
    select(-boot, -ci) %>%
    # Unnest
    unnest(cols = c(data, point est, lower ci, upper ci)) %>%
   ungroup()
plot_nofacet.sex2 <- pubs_nofacet.sex2 %>%
    select(-CD4_recent, -Age, -Pain_present, -Site, -Pain_def) %>%
   distinct() %>%
   mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '_',
                                   replacement = ' ')) %>%
   mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '\\.',
                                   replacement = ' & ')) %>%
   mutate(Pain_site = ifelse(Pain_site == 'Lower back',
                              yes = 'Lower back/flanks',
                              no = Pain_site)) %>%
   mutate(Pain_site = factor(Pain_site,
                              levels = c('Throat', 'Shoulder', 'Arms',
                                          'Elbows', 'Wrists & Hands',
                                         'Lower back/flanks', 'Groin', 'Hips',
                                          'Buttocks', 'Cervical spine'),
                              ordered = TRUE)) %>%
   ggplot(data = .) +
    aes(x = fct_rev(Pain_site),
        y = point_est,
       ymin = lower_ci,
       ymax = upper_ci,
        fill = Sex) +
    geom_linerange(position = position_dodge2(width = 0.6),
                   size = 1,
```

```
colour = '#000000') +
    geom_point(shape = 21,
               colour = '#000000',
               position = position_dodge2(width = 0.6),
               size = 4,
               stroke = 1) +
    coord_flip() +
   labs(title = 'Body sites by sex',
         subtitle = '(Body sites with <10% pain; Point estimate with 95%CI)',</pre>
         y = 'Proportion of participants with pain') +
    scale_y_continuous(limits = c(0, 0.3)) +
    scale_fill_manual(values = c('#FFFFFF', '#000000')) +
   theme_minimal(base_size = 18) +
   theme(legend.position = c(0.8, 0.1),
          legend.title = element_blank(),
          plot.title = element_text(size = 18),
          plot.subtitle = element_text(size = 12),
          axis.title.y = element_blank(),
          axis.text.y = element_blank(),
          panel.grid = element_blank(),
          axis.text = element_text(colour = '#000000'),
          axis.line = element_line(size = 0.5),
          axis.ticks = element_line(size = 0.5))
#-- Body sites by pain chronicity --#
pubs_nofacet.pain2 <- data_less10 %>%
   group_by(Pain_site, Pain_def) %>%
   nest() %>%
    # Boostrap data
   mutate(boot = map(.x = data,
                      ~ boot(data = .x[, 'Pain_present'],
                             statistic = prop_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
    mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                           ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
    select(-boot, -ci) %>%
    # Unnest
   unnest(cols = c(data, point_est, lower_ci, upper_ci)) %>%
    ungroup()
plot_nofacet.pain2 <- pubs_nofacet.pain2 %>%
```

```
select(-CD4_recent, -Age, -Pain_present, -Site, -Sex) %>%
    distinct() %>%
   mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '_',
                                   replacement = ' ')) %>%
    mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '\\.',
                                   replacement = ' & ')) %>%
   mutate(Pain_site = ifelse(Pain_site == 'Lower back',
                              yes = 'Lower back/flanks',
                              no = Pain_site)) %>%
   mutate(Pain_site = factor(Pain_site,
                              levels = c('Throat', 'Shoulder', 'Arms',
                                          'Elbows', 'Wrists & Hands',
                                         'Lower back/flanks', 'Groin', 'Hips',
                                          'Buttocks', 'Cervical spine'),
                              ordered = TRUE)) %>%
    ggplot(data = .) +
    aes(x = fct_rev(Pain_site),
        y = point_est,
        ymin = lower_ci,
       ymax = upper_ci,
       fill = Pain def) +
    geom_linerange(position = position_dodge2(width = 0.6),
                   size = 1,
                   colour = '#000000') +
    geom_point(shape = 21,
               colour = '#000000',
               position = position_dodge2(width = 0.6),
               size = 4,
               stroke = 1) +
    coord_flip() +
   labs(title = 'Body sites by pain chronicity',
         subtitle = '(Body sites with <10% pain; Point estimate with 95%CI)',</pre>
         y = 'Proportion of participants with pain') +
    scale_y_continuous(limits = c(0, 0.3)) +
    scale_fill_manual(values = c('#FFFFFF', '#000000')) +
   theme minimal(base size = 18) +
   theme(legend.position = c(0.8, 0.1),
          legend.title = element_blank(),
          plot.title = element_text(size = 18),
         plot.subtitle = element_text(size = 12),
          axis.title.y = element_blank(),
          panel.grid = element_blank(),
          axis.text = element_text(colour = '#000000'),
          axis.line = element_line(size = 0.5),
          axis.ticks = element_line(size = 0.5))
#-- Body sites by age --#
pubs_nofacet.age2 <- data_less10 %>%
    select(-CD4_recent, -Site, -Sex, -Pain_def) %>%
   mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '_',
```

```
replacement = ' ')) %>%
    mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '\\.',
                                   replacement = ' & ')) %>%
    group_by(Pain_site, Pain_present) %>%
   nest() %>%
   mutate(boot = map(.x = data,
                      ~ boot(data = .x,
                             statistic = median_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
   mutate(ci = map(.x = boot,
                   " boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                           ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
    select(-boot, -ci) %>%
    # Unnest
   unnest(cols = c(data, point_est, lower_ci, upper_ci)) %>%
   ungroup() %>%
   mutate(Pain_site = ifelse(Pain_site == 'Lower back',
                              yes = 'Lower back/flanks',
                              no = Pain_site)) %>%
   mutate(Pain_site = factor(Pain_site,
                              levels = c('Throat', 'Shoulder', 'Arms',
                                         'Elbows', 'Wrists & Hands',
                                          'Lower back/flanks', 'Groin', 'Hips',
                                          'Buttocks', 'Cervical spine'),
                              ordered = TRUE)) %>%
    select(-Age) %>%
   distinct()
plot_nofacet.age2 <- pubs_nofacet.age2 %>%
    ggplot(data = .) +
    aes(x = fct_rev(Pain_site),
       y = point_est,
       ymin = lower_ci,
       ymax = upper_ci,
        fill = Pain_present) +
    geom_linerange(position = position_dodge2(width = 0.6),
                   size = 1,
                   colour = '#000000') +
    geom_point(shape = 21,
               colour = '#000000',
               position = position_dodge2(width = 0.6),
```

```
size = 4,
               stroke = 1) +
    coord_flip() +
    labs(title = 'Body sites by median age',
         subtitle = '(Body sites with <10% pain; Point estimate with 95%CI)',</pre>
         y = 'Age (years)') +
    scale_fill_manual(values = c('#FFFFFF', '#000000'),
                      labels = c('No pain', 'Pain present')) +
    scale_y_continuous(limits = c(27, 47),
                       breaks = c(27, 31, 35, 39, 43, 47)) +
   theme_minimal(base_size = 18) +
    theme(legend.position = c(0.8, 0.1),
          legend.title = element_blank(),
          plot.title = element text(size = 18),
          plot.subtitle = element_text(size = 12),
          axis.title.y = element_blank(),
          axis.text.y = element_blank(),
          panel.grid = element_blank(),
          axis.text = element_text(colour = '#000000'),
          axis.line = element_line(size = 0.5),
          axis.ticks = element_line(size = 0.5))
#-- Body sites by CD4 --#
pubs_nofacet.cd42 <- data_less10 %>%
    select(-Age, -Site, -Sex, -Pain_def) %>%
   mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '_',
                                   replacement = ' ')) %>%
    mutate(Pain_site = str_replace(Pain_site,
                                   pattern = '\\.',
                                   replacement = ' & ')) %>%
   group_by(Pain_site, Pain_present) %>%
   nest() %>%
   mutate(boot = map(.x = data,
                      ~ boot(data = .x,
                             statistic = median_func,
                             R = 999,
                             stype = 'i',
                             parallel = 'multicore',
                             ncpus = 4))) %>%
    # Get CI
   mutate(ci = map(.x = boot,
                    ~ boot.ci(.x, type = 'perc'))) %>%
    # Extract ci data
   mutate(point_est = map(.x = ci,
                           ~ .x$t0),
           lower_ci = map(.x = ci,
                          ~ .x$percent[[4]]),
           upper_ci = map(.x = ci,
                          ~ .x$percent[[5]])) %>%
    # Remove columns
    select(-boot, -ci) %>%
    # Unnest
```

```
unnest(cols = c(data, point_est, lower_ci, upper_ci)) %>%
    ungroup() %>%
   mutate(Pain_site = ifelse(Pain_site == 'Lower back',
                              ves = 'Lower back/flanks',
                              no = Pain_site)) %>%
    mutate(Pain site = factor(Pain site,
                              levels = c('Throat', 'Shoulder', 'Arms',
                                          'Elbows', 'Wrists & Hands',
                                          'Lower back/flanks', 'Groin', 'Hips',
                                          'Buttocks', 'Cervical spine'),
                              ordered = TRUE)) %>%
    select(-CD4_recent) %>%
    distinct()
plot_nofacet.cd42 <- pubs_nofacet.cd42 %>%
    ggplot(data = .) +
    aes(x = fct_rev(Pain_site),
       y = point_est,
       ymin = lower_ci,
       ymax = upper_ci,
        fill = Pain_present) +
    geom_linerange(position = position_dodge2(width = 0.6),
                   size = 1,
                   colour = '#000000') +
    geom point(shape = 21,
               colour = '#000000',
               position = position_dodge2(width = 0.6),
               size = 4,
               stroke = 1) +
    coord_flip() +
   labs(title = 'Body sites by median CD4 T-cell count',
         subtitle = '(Body sites with <10% pain; Point estimate with 95%CI)',
         y = expression('CD4 T-cell count (cell.mm'^-3*')')) +
    scale_fill_manual(values = c('#FFFFFF', '#000000'),
                      labels = c('No pain', 'Pain present')) +
    scale_y_continuous(limits = c(50, 600),
                       breaks = c(100, 200, 300, 400, 500, 600)) +
   theme minimal(base size = 18) +
    theme(legend.position = c(0.8, 0.95),
          legend.title = element_blank(),
          plot.title = element_text(size = 18),
          plot.subtitle = element_text(size = 12),
          axis.title.y = element_blank(),
          panel.grid = element_blank(),
          axis.text = element_text(colour = '#000000'),
          axis.line = element_line(size = 0.5),
          axis.ticks = element_line(size = 0.5))
# Patchwork
pub_plot2 <- plot_nofacet2 + plot_nofacet.sex2 + plot_nofacet.pain2 +</pre>
   plot_nofacet.age2 + plot_nofacet.cd42 +
   plot_layout(ncol = 2)
# Save
```

```
ggsave(filename = 'figures/figure_supp2.png',
    width = 14,
    height = 18)
```

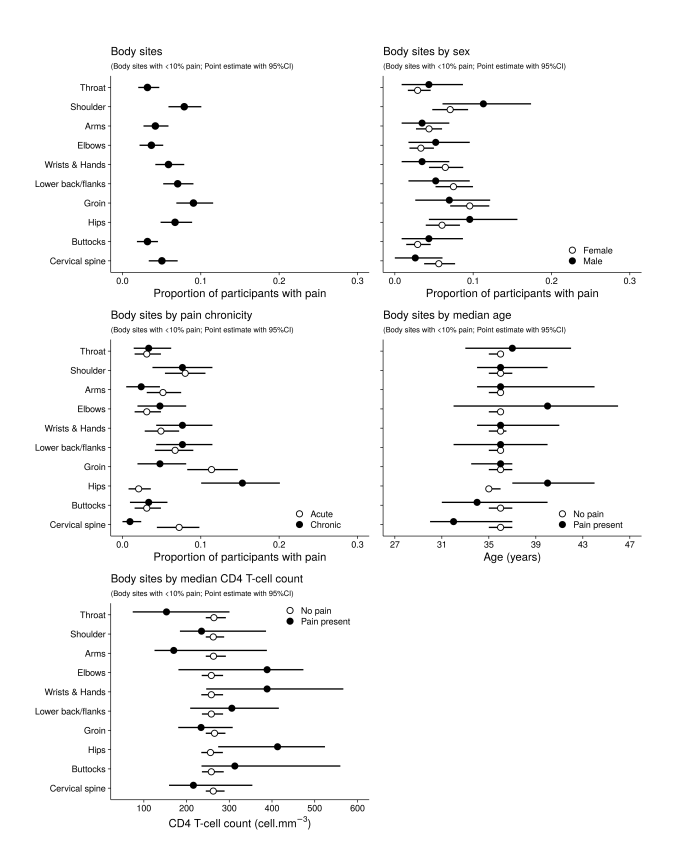


Figure 3: Body sites with a point estimate pain proportion <10%

10 Session information

sessionInfo()

```
## R version 3.6.3 (2020-02-29)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Debian GNU/Linux 10 (buster)
##
## Matrix products: default
## BLAS/LAPACK: /usr/lib/x86_64-linux-gnu/libopenblasp-r0.3.5.so
## locale:
   [1] LC_CTYPE=en_US.UTF-8
                                   LC NUMERIC=C
##
  [3] LC TIME=en US.UTF-8
                                   LC_COLLATE=en_US.UTF-8
   [5] LC_MONETARY=en_US.UTF-8
                                   LC MESSAGES=C
##
   [7] LC_PAPER=en_US.UTF-8
                                   LC_NAME=C
  [9] LC_ADDRESS=C
                                   LC_TELEPHONE=C
##
## [11] LC MEASUREMENT=en US.UTF-8 LC IDENTIFICATION=C
## attached base packages:
## [1] stats
                 graphics grDevices utils
                                               datasets methods
                                                                    base
##
## other attached packages:
   [1] patchwork_1.0.0 lme4_1.1-23
                                        Matrix_1.2-18
                                                         boot 1.3-24
##
   [5] knitr_1.28
                        skimr_2.1.1
                                        forcats_0.5.0
                                                         stringr_1.4.0
                                                         tidyr_1.0.2
  [9] dplyr_0.8.5
                        purrr_0.3.4
                                        readr_1.3.1
## [13] tibble_3.0.1
                        ggplot2_3.3.0
                                        tidyverse_1.3.0
##
## loaded via a namespace (and not attached):
                                          lattice 0.20-38 utf8 1.1.4
  [1] Rcpp 1.0.4.6
                         lubridate 1.7.8
   [5] assertthat_0.2.1 digest_0.6.25
                                          R6_2.4.1
                                                            cellranger_1.1.0
## [9] repr_1.1.0
                         backports_1.1.6
                                          reprex_0.3.0
                                                            evaluate_0.14
                                                            rlang_0.4.5
## [13] highr_0.8
                         httr_1.4.1
                                          pillar_1.4.3
## [17] readxl_1.3.1
                         rstudioapi_0.11
                                          minqa_1.2.4
                                                            nloptr_1.2.2.1
## [21] rmarkdown 2.1
                         labeling 0.3
                                          splines 3.6.3
                                                            statmod 1.4.34
## [25] munsell_0.5.0
                                                            modelr_0.1.6
                         broom_0.5.6
                                          compiler_3.6.3
## [29] xfun 0.13
                         pkgconfig 2.0.3
                                          base64enc 0.1-3
                                                           htmltools 0.4.0
## [33] tidyselect_1.0.0 fansi_0.4.1
                                          crayon_1.3.4
                                                            dbplyr_1.4.3
## [37] withr_2.2.0
                         MASS_7.3-51.5
                                          grid_3.6.3
                                                            nlme_3.1-144
## [41] jsonlite_1.6.1
                         gtable_0.3.0
                                          lifecycle_0.2.0 DBI_1.1.0
## [45] magrittr_1.5
                         scales_1.1.0
                                          cli_2.0.2
                                                            stringi_1.4.6
## [49] farver_2.0.3
                         fs_1.4.1
                                          xm12_1.3.2
                                                            ellipsis_0.3.0
## [53] generics_0.0.2
                         vctrs_0.2.4
                                          tools_3.6.3
                                                            glue_1.4.0
## [57] hms_0.5.3
                         parallel_3.6.3
                                          yaml_2.2.1
                                                            colorspace_1.4-1
## [61] rvest_0.3.5
                         haven_2.2.0
```