Covid in Georgia

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## Intro

Georgia reported their first confirmed case of Covid-19 on 26 February 2020. As of 2 July 2020, only 939 confirmed cases and 15 deaths have been reported. The success of Georgia’s limitation of spread of Covid-19 can be attributed to early non-pharmaceutical interventions including closing schools (early March), closing borders and mandatory quarantining of international arrivals, lockdown of individual affected areas (late March) and full lockdown throughout the country (30 March). Alongside this, they have implemented extensive contact tracing and testing, and all confirmed cases are treated in hospital. There is regular screening of key workers. As of 2 July 2020, 117,701 PCR tests for Covid-19 have been conducted, a rate of more than 30,000 tests per million population.

Here we use test data and contact tracing data collected by the National Center for Disease Control (NCDC) in Georgia to characterize the epidemic in Georgia. This includes demographics of those infected compared to the general population, symptom and death rates by age, and rates of transmission to different types of contacts. We also calculate the secondary attack rate for different types of contacts and overall number of secondary cases per primary case (observed reproductive number).

## Methods

### Data collection

* Types of test used, criteria for testing (case identification).
  + A variety of test types were used, here we present the results from antigen tests confirming current infection, primarily real time PCR tests. [In the database as confirmation test result]
  + Data exported on 24 June 2020 [match numbers in intro to data we have]
* More detail on policies - maybe in a figure
* Ethics waiver from NCDC and registered in Bristol

### Analysis

* De-identified personal IDs used to link contact tracing database to confirmed cases
* Data cut off date 24 June 2020
* Analysis conducted in R
* Can we compare demographics of infection to national demographics? geostat or other source?

## Results

##   
## Attaching package: 'lubridate'

## The following objects are masked from 'package:base':  
##   
## date, intersect, setdiff, union

##   
## Attaching package: 'dplyr'

## The following objects are masked from 'package:stats':  
##   
## filter, lag

## The following objects are masked from 'package:base':  
##   
## intersect, setdiff, setequal, union

##   
## Attaching package: 'gridExtra'

## The following object is masked from 'package:dplyr':  
##   
## combine

## Loading required package: lattice

## Loading required package: survival

## Loading required package: Formula

##   
## Attaching package: 'Hmisc'

## The following objects are masked from 'package:dplyr':  
##   
## src, summarize

## The following objects are masked from 'package:base':  
##   
## format.pval, units

## Warning: Removed 714 row(s) containing missing values (geom\_path).

## Warning: Removed 912 row(s) containing missing values (geom\_path).

## Warning: Removed 509 row(s) containing missing values (geom\_path).

## quartz\_off\_screen   
## 2

## Warning: Removed 3 rows containing missing values (geom\_bar).

## Warning: Removed 912 row(s) containing missing values (geom\_path).

## Warning: Removed 509 row(s) containing missing values (geom\_path).

## quartz\_off\_screen   
## 2

### Table of demographics of those infected (treated database)

Hospitalization data and comorbidities were available for 500 diagnosed patients with hospitalization dates ranging from to -. Could add more detail here on particular symptoms if of interest.

##   
## Attaching package: 'data.table'

## The following objects are masked from 'package:dplyr':  
##   
## between, first, last

## The following objects are masked from 'package:lubridate':  
##   
## hour, isoweek, mday, minute, month, quarter, second, wday, week,  
## yday, year

##   
## Attaching package: 'arsenal'

## The following object is masked from 'package:Hmisc':  
##   
## %nin%

## The following object is masked from 'package:lubridate':  
##   
## is.Date

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  | 0-49 (N=305) | 50+ (N=195) | Total (N=500) | p value |
| **Age (years)** |  |  |  | < 0.001 |
| Mean (SD) | 30.364 (12.586) | 62.087 (9.875) | 42.736 (19.347) |  |
| Median (Q1, Q3) | 33.000 (21.000, 40.000) | 60.000 (54.000, 68.000) | 44.000 (29.000, 56.250) |  |
| Min - Max | 0.000 - 49.000 | 50.000 - 90.000 | 0.000 - 90.000 |  |
| Missing | 0 | 0 | 0 |  |
| **Gender** |  |  |  | < 0.001 |
| Female | 132 (43.3%) | 121 (62.1%) | 253 (50.6%) |  |
| Male | 173 (56.7%) | 74 (37.9%) | 247 (49.4%) |  |
| Missing | 0 | 0 | 0 |  |
| **Disease severity** |  |  |  | < 0.001 |
| critical | 5 (1.6%) | 18 (9.2%) | 23 (4.6%) |  |
| mild | 208 (68.2%) | 67 (34.4%) | 275 (55.0%) |  |
| moderate | 81 (26.6%) | 82 (42.1%) | 163 (32.6%) |  |
| Severe | 11 (3.6%) | 28 (14.4%) | 39 (7.8%) |  |
| Missing | 0 | 0 | 0 |  |
| **Length of hospital stay** |  |  |  | 0.616 |
| Mean (SD) | 21.770 (8.609) | 22.021 (8.511) | 21.868 (8.563) |  |
| Median (Q1, Q3) | 20.000 (16.000, 27.000) | 21.000 (17.000, 26.500) | 21.000 (16.000, 27.000) |  |
| Min - Max | 2.000 - 64.000 | 2.000 - 65.000 | 2.000 - 65.000 |  |
| Missing | 1 | 0 | 1 |  |
| **Outcome** |  |  |  | < 0.001 |
| dead | 1 (0.3%) | 12 (6.2%) | 13 (2.6%) |  |
| recovered | 304 (99.7%) | 183 (93.8%) | 487 (97.4%) |  |
| Missing | 0 | 0 | 0 |  |
| **Diagnosis date** |  |  |  | 0.628 |
| Nmiss | 38 | 27 | 65 |  |
| median | 2020-04-14 | 2020-04-12 | 2020-04-13 |  |
| Min - Max | 2020-03-04 - 2020-05-02 | 2020-03-10 - 2020-05-02 | 2020-03-04 - 2020-05-02 |  |
| **Place of exposure** |  |  |  | 0.024 |
| Europe and Central Asia | 42 (16.0%) | 12 (7.3%) | 54 (12.6%) |  |
| Georgia | 213 (81.0%) | 149 (90.3%) | 362 (84.6%) |  |
| Latin America and the Caribbean | 0 (0.0%) | 1 (0.6%) | 1 (0.2%) |  |
| Middle East and North Africa | 4 (1.5%) | 0 (0.0%) | 4 (0.9%) |  |
| North America | 4 (1.5%) | 3 (1.8%) | 7 (1.6%) |  |
| Missing | 42 | 30 | 72 |  |

###Table of demographics of those tested (tested database)

A total of 96317 antigen (PCR) test results were available, with 67886 unique individual patient IDs recorded.

## [1] "2080-01-21"

## [1] "1902-01-01"

## `summarise()` ungrouping output (override with `.groups` argument)

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | neg (N=66941) | pos SARS-CoV-2 (N=763) | Suspicted (N=156) | Total (N=67860) | p value |
| **Age** |  |  |  |  | 0.065 |
| Mean (SD) | 41.279 (18.929) | 42.733 (19.786) | 42.718 (20.144) | 41.298 (18.942) |  |
| Median (Q1, Q3) | 40.000 (28.000, 55.000) | 44.000 (28.000, 57.000) | 40.500 (28.750, 57.250) | 40.000 (28.000, 55.000) |  |
| Min - Max | 0.000 - 102.000 | 0.000 - 91.000 | 0.000 - 90.000 | 0.000 - 102.000 |  |
| Missing | 9 | 0 | 0 | 9 |  |
| **Gender** |  |  |  |  | 0.547 |
| Female | 34552 (51.6%) | 389 (51.0%) | 74 (47.4%) | 35015 (51.6%) |  |
| male | 32389 (48.4%) | 374 (49.0%) | 82 (52.6%) | 32845 (48.4%) |  |
| Missing | 0 | 0 | 0 | 0 |  |
| **Fever** |  |  |  |  | 0.006 |
| No | 6003 (32.6%) | 30 (21.0%) | 7 (21.9%) | 6040 (32.5%) |  |
| Yes | 12405 (67.4%) | 113 (79.0%) | 25 (78.1%) | 12543 (67.5%) |  |
| Missing | 48533 | 620 | 124 | 49277 |  |
| **Cough** |  |  |  |  | 0.855 |
| No | 12227 (68.9%) | 88 (66.7%) | 21 (67.7%) | 12336 (68.9%) |  |
| Yes | 5527 (31.1%) | 44 (33.3%) | 10 (32.3%) | 5581 (31.1%) |  |
| Missing | 49187 | 631 | 125 | 49943 |  |
| **Risk Group** |  |  |  |  | < 0.001 |
| Contact of confirmed case | 4057 (7.4%) | 50 (6.7%) | 13 (9.2%) | 4120 (7.4%) |  |
| Healthcare worker | 11719 (21.4%) | 7 (0.9%) | 13 (9.2%) | 11739 (21.1%) |  |
| Non-covid patient | 1460 (2.7%) | 1 (0.1%) | 2 (1.4%) | 1463 (2.6%) |  |
| Other | 2904 (5.3%) | 5 (0.7%) | 1 (0.7%) | 2910 (5.2%) |  |
| Symptomatic | 20045 (36.6%) | 634 (84.8%) | 63 (44.7%) | 20742 (37.3%) |  |
| Transport/other essential worker | 3064 (5.6%) | 8 (1.1%) | 7 (5.0%) | 3079 (5.5%) |  |
| Under quarantine/isolation | 11540 (21.1%) | 43 (5.7%) | 42 (29.8%) | 11625 (20.9%) |  |
| Missing | 12152 | 15 | 15 | 12182 |  |
| **IsForeigner** |  |  |  |  | 0.515 |
| FALSE | 4323 (99.5%) | 253 (100.0%) | 8 (100.0%) | 4584 (99.5%) |  |
| TRUE | 22 (0.5%) | 0 (0.0%) | 0 (0.0%) | 22 (0.5%) |  |
| Missing | 62596 | 510 | 148 | 63254 |  |

## Contact networks

Index cases summary

contacts$missingPatID <- is.na(contacts$`Patient unique ID`)  
contacts$missingPatID[contacts$missingPatID] <- 1:sum(contacts$missingPatID)  
contacts$newPatID <- ifelse(is.na(contacts$`Patient unique ID`),paste("missingP",contacts$missingPatID,sep=""),contacts$`Patient unique ID`)  
  
contacts$missingCID <- is.na(contacts$`Contact ID`)  
contacts$missingCID[contacts$missingCID] <- 1:sum(contacts$missingCID)  
contacts$newCID <- ifelse(is.na(contacts$`Contact ID`),paste("missingC",contacts$missingCID,sep=""),contacts$`Contact ID`)  
  
contacts$case <- as.factor(contacts$`Confirmed Case`)  
levels(contacts$case) <- c(FALSE,NA,TRUE)  
  
contacts$Age[contacts$Age=="\*"] <- NA  
contacts$Age <- as.numeric(contacts$Age)  
contacts$Gender[contacts$Gender=="male"] <- "Male"  
# rate of transmission by type of contact  
  
contactsType <- contacts %>% filter(`Contact type`!="unknown")  
contactsType$Coworker <- contactsType$`Contact type`=="Co-worker"  
contactsType$Family <- contactsType$`Contact type`=="Family member"  
contactsType$Friend <- contactsType$`Contact type`=="Friend"  
contactsType$Medical <- contactsType$`Contact type`=="Medical Personnel"  
contactsType$Neighbour <- contactsType$`Contact type`=="Neighbour"  
  
mType <- glm(data=contactsType,case ~ `Contact type`,family=binomial)  
summary(mType)

##   
## Call:  
## glm(formula = case ~ `Contact type`, family = binomial, data = contactsType)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.9628 -0.9628 -0.7306 1.4084 2.1964   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -1.1844 0.1185 -9.995 < 2e-16 \*\*\*  
## `Contact type`Family member 0.6560 0.1283 5.112 3.19e-07 \*\*\*  
## `Contact type`Friend -0.2709 0.2410 -1.124 0.261   
## `Contact type`Medical Personnel -15.3816 302.3142 -0.051 0.959   
## `Contact type`Neighbour -1.1338 0.2681 -4.230 2.34e-05 \*\*\*  
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3032.8 on 2581 degrees of freedom  
## AIC: 3042.8  
##   
## Number of Fisher Scoring iterations: 15

mType2 <- glm(data=contactsType,case ~ Family,family=binomial)  
summary(mType2)

##   
## Call:  
## glm(formula = case ~ Family, family = binomial, data = contactsType)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.9628 -0.9628 -0.6119 1.4084 1.8802   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -1.58045 0.09281 -17.03 <2e-16 \*\*\*  
## FamilyTRUE 1.05207 0.10507 10.01 <2e-16 \*\*\*  
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3078.7 on 2584 degrees of freedom  
## AIC: 3082.7  
##   
## Number of Fisher Scoring iterations: 4

mType3 <- glm(data=contactsType,case ~ Coworker,family=binomial)  
summary(mType3)

##   
## Call:  
## glm(formula = case ~ Coworker, family = binomial, data = contactsType)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.8794 -0.8794 -0.8794 1.5082 1.7037   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.75058 0.04579 -16.391 < 2e-16 \*\*\*  
## CoworkerTRUE -0.43385 0.12704 -3.415 0.000638 \*\*\*  
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3179.0 on 2584 degrees of freedom  
## AIC: 3183  
##   
## Number of Fisher Scoring iterations: 4

mType4 <- glm(data=contactsType,case ~ Friend,family=binomial)  
summary(mType4)

##   
## Call:  
## glm(formula = case ~ Friend, family = binomial, data = contactsType)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.8692 -0.8692 -0.8692 1.5208 1.8248   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.77869 0.04361 -17.854 <2e-16 \*\*\*  
## FriendTRUE -0.67660 0.21436 -3.156 0.0016 \*\*   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3180.0 on 2584 degrees of freedom  
## AIC: 3184  
##   
## Number of Fisher Scoring iterations: 4

mType5 <- glm(data=contactsType,case ~ Medical,family=binomial)  
summary(mType5)

##   
## Call:  
## glm(formula = case ~ Medical, family = binomial, data = contactsType)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.87 -0.87 -0.87 1.52 1.52   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.77638 0.04286 -18.116 <2e-16 \*\*\*  
## MedicalTRUE -15.78969 302.31422 -0.052 0.958   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3144.3 on 2584 degrees of freedom  
## AIC: 3148.3  
##   
## Number of Fisher Scoring iterations: 15

mType6 <- glm(data=contactsType,case ~ Neighbour,family=binomial)  
summary(mType6)

##   
## Call:  
## glm(formula = case ~ Neighbour, family = binomial, data = contactsType)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.8898 -0.8898 -0.8898 1.4954 2.1964   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.72236 0.04375 -16.509 < 2e-16 \*\*\*  
## NeighbourTRUE -1.59590 0.24434 -6.532 6.51e-11 \*\*\*  
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3128.4 on 2584 degrees of freedom  
## AIC: 3132.4  
##   
## Number of Fisher Scoring iterations: 4

####  
# age and gender of index case  
mAS <- glm(data=contacts,case ~ Age + Gender, family=binomial)  
summary(mAS)

##   
## Call:  
## glm(formula = case ~ Age + Gender, family = binomial, data = contacts)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.9192 -0.8914 -0.8190 1.4782 1.6140   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.857966 0.111157 -7.718 1.18e-14 \*\*\*  
## Age -0.001415 0.002191 -0.646 0.5186   
## GenderMale 0.217877 0.086238 2.526 0.0115 \*   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3183.9 on 2583 degrees of freedom  
## (296 observations deleted due to missingness)  
## AIC: 3189.9  
##   
## Number of Fisher Scoring iterations: 4

mA <- glm(data=contacts,case ~ Age , family=binomial)  
summary(mA)

##   
## Call:  
## glm(formula = case ~ Age, family = binomial, data = contacts)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.8883 -0.8645 -0.8465 1.5139 1.5847   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.724045 0.096973 -7.466 8.24e-14 \*\*\*  
## Age -0.002181 0.002164 -1.008 0.314   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3190.2 on 2584 degrees of freedom  
## (296 observations deleted due to missingness)  
## AIC: 3194.2  
##   
## Number of Fisher Scoring iterations: 4

mS <- glm(data=contacts,case ~ Gender, family=binomial)  
summary(mS)

##   
## Call:  
## glm(formula = case ~ Gender, family = binomial, data = contacts)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.9005 -0.9005 -0.8195 1.4823 1.5840   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.91881 0.05936 -15.478 < 2e-16 \*\*\*  
## GenderMale 0.22566 0.08540 2.642 0.00823 \*\*   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3184.3 on 2584 degrees of freedom  
## (296 observations deleted due to missingness)  
## AIC: 3188.3  
##   
## Number of Fisher Scoring iterations: 4

### merge contacts with screening data  
testsub <- testingU %>% filter(!is.na(PersonalID)) %>% dplyr::select(PersonalID,Gender,Age)  
# testdate <- testingU %>% dplyr::select(PersonalID,Date) ### need to start from full PCR testing database for this  
testdate <- testingF %>% dplyr::select(PersonalID,`Test Date`,`Confirmation Test Result`) %>% filter(!is.na(PersonalID))  
cases <- contacts$`Patient unique ID`  
cases2 <- testdate %>% filter(PersonalID %in% cases)  
cases.firstpos <- cases2 %>% group\_by(PersonalID) %>% filter(`Confirmation Test Result` =="pos SARS-CoV-2") %>% summarise(Date=min(`Test Date`))

## `summarise()` ungrouping output (override with `.groups` argument)

contacts2 <- left\_join(contacts,cases.firstpos,by=c("Patient unique ID"="PersonalID"))  
  
# contacts.results <- testdate %>% filter(PersonalID %in% contacts2$`Contact ID`) # don't need this after all  
  
contact.merge <- left\_join(contacts2,testsub,by=c("Contact ID"="PersonalID")) # don't need date of follow up test  
contactsType.m <- contact.merge %>% filter(`Contact type`!="unknown")  
contactsType.m$Coworker <- contactsType.m$`Contact type`=="Co-worker"  
contactsType.m$Family <- contactsType.m$`Contact type`=="Family member"  
contactsType.m$Friend <- contactsType.m$`Contact type`=="Friend"  
contactsType.m$Medical <- contactsType.m$`Contact type`=="Medical Personnel"  
contactsType.m$Neighbour <- contactsType.m$`Contact type`=="Neighbour"  
  
### numbers in each time period  
length(unique(contact.merge$newPatID[!is.na(contact.merge$Date)]))

## [1] 760

contact.merge$TimePeriod <- NA  
contact.merge$TimePeriod <- with(contact.merge,ifelse(Date<="2020-03-29","Pre-lockdown",ifelse(Date>"2020-03-29" & Date <= "2020-04-27","Lockdown","Restrictions easing")))  
  
contact.merge %>% group\_by(TimePeriod) %>%   
 summarise(cases=length(unique(newPatID)),.groups="keep") # cases in each time period

## # A tibble: 4 x 2  
## # Groups: TimePeriod [4]  
## TimePeriod cases  
## <chr> <int>  
## 1 Lockdown 352  
## 2 Pre-lockdown 66  
## 3 Restrictions easing 342  
## 4 <NA> 157

contact.merge %>% group\_by(TimePeriod,newPatID) %>%   
 summarise(contacts=n(),.groups="keep") %>% group\_by(TimePeriod) %>% summarise(mean=mean(contacts),min=min(contacts),max=max(contacts),median=median(contacts)) # number of contacts in each time period

## `summarise()` ungrouping output (override with `.groups` argument)

## # A tibble: 4 x 5  
## TimePeriod mean min max median  
## <chr> <dbl> <int> <int> <dbl>  
## 1 Lockdown 3.68 1 23 3  
## 2 Pre-lockdown 2.73 1 17 2  
## 3 Restrictions easing 3.06 1 30 2  
## 4 <NA> 2.28 1 23 1

contact.merge %>% filter(`Contact type`!="unknown") %>% group\_by(TimePeriod,newPatID) %>%   
 summarise(contacts=n(),.groups="keep") %>% group\_by(TimePeriod) %>% summarise(mean=mean(contacts),min=min(contacts),max=max(contacts),median=median(contacts)) # number of contacts in each time period

## `summarise()` ungrouping output (override with `.groups` argument)

## # A tibble: 4 x 5  
## TimePeriod mean min max median  
## <chr> <dbl> <int> <int> <dbl>  
## 1 Lockdown 4.38 1 23 4   
## 2 Pre-lockdown 3.28 1 17 2.5  
## 3 Restrictions easing 3.97 1 30 3   
## 4 <NA> 3.80 1 23 3

contact.merge %>% group\_by(TimePeriod) %>%   
 summarise(contacts=n(),newcase=sum(as.logical(case),na.rm=TRUE),.groups="keep")

## # A tibble: 4 x 3  
## # Groups: TimePeriod [4]  
## TimePeriod contacts newcase  
## <chr> <int> <int>  
## 1 Lockdown 1296 383  
## 2 Pre-lockdown 180 38  
## 3 Restrictions easing 1048 297  
## 4 <NA> 358 77

contact.merge %>% group\_by(TimePeriod,`Contact type`) %>%   
 summarise(contacts=n(),newcase=sum(as.logical(case),na.rm=TRUE),.groups="keep") %>% View()  
  
contact.merge %>% group\_by(`Contact type`) %>%   
 summarise(contacts=n(),newcase=sum(as.logical(case),na.rm=TRUE),.groups="keep")

## # A tibble: 6 x 3  
## # Groups: Contact type [6]  
## `Contact type` contacts newcase  
## <chr> <int> <int>  
## 1 Co-worker 397 93  
## 2 Family member 1766 655  
## 3 Friend 148 28  
## 4 Medical Personnel 63 0  
## 5 Neighbour 212 19  
## 6 unknown 296 0

m2 <- glm(data=contact.merge,case ~ Age.x,family=binomial)  
summary(m2)

##   
## Call:  
## glm(formula = case ~ Age.x, family = binomial, data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.8883 -0.8645 -0.8465 1.5139 1.5847   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.724045 0.096973 -7.466 8.24e-14 \*\*\*  
## Age.x -0.002181 0.002164 -1.008 0.314   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3190.2 on 2584 degrees of freedom  
## (296 observations deleted due to missingness)  
## AIC: 3194.2  
##   
## Number of Fisher Scoring iterations: 4

require(MASS)

## Loading required package: MASS

##   
## Attaching package: 'MASS'

## The following object is masked from 'package:dplyr':  
##   
## select

exp(cbind(coef(m2), confint(m2)))

## Waiting for profiling to be done...

## 2.5 % 97.5 %  
## (Intercept) 0.4847872 0.4004405 0.585706  
## Age.x 0.9978212 0.9935923 1.002060

m3 <- glm(data=contact.merge,case ~ Age.x + Age.y + Gender.x + Gender.y + `Contact type`,family=binomial)  
summary(m3)

##   
## Call:  
## glm(formula = case ~ Age.x + Age.y + Gender.x + Gender.y + `Contact type`,   
## family = binomial, data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -1.3708 -1.0307 -0.7387 1.2221 2.1258   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -1.295e+00 2.011e-01 -6.442 1.18e-10 \*\*\*  
## Age.x -8.324e-04 2.400e-03 -0.347 0.728743   
## Age.y 1.116e-02 2.429e-03 4.596 4.31e-06 \*\*\*  
## Gender.xMale 1.632e-01 9.686e-02 1.685 0.091947 .   
## Gender.ymale -9.995e-02 9.662e-02 -1.034 0.300957   
## `Contact type`Family member 6.626e-01 1.423e-01 4.657 3.22e-06 \*\*\*  
## `Contact type`Friend -3.569e-01 2.602e-01 -1.371 0.170226   
## `Contact type`Medical Personnel -1.479e+01 2.290e+02 -0.065 0.948517   
## `Contact type`Neighbour -1.026e+00 2.866e-01 -3.581 0.000342 \*\*\*  
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 2616.8 on 1963 degrees of freedom  
## Residual deviance: 2466.2 on 1955 degrees of freedom  
## (918 observations deleted due to missingness)  
## AIC: 2484.2  
##   
## Number of Fisher Scoring iterations: 14

m4 <- glm(data=contactsType.m,case ~ Age.x + Age.y + Gender.x + Gender.y + Coworker + Family + Friend + Medical + Neighbour,family=binomial)  
summary(m4)

##   
## Call:  
## glm(formula = case ~ Age.x + Age.y + Gender.x + Gender.y + Coworker +   
## Family + Friend + Medical + Neighbour, family = binomial,   
## data = contactsType.m)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -1.3708 -1.0307 -0.7387 1.2221 2.1258   
##   
## Coefficients: (1 not defined because of singularities)  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -2.321e+00 3.037e-01 -7.643 2.12e-14 \*\*\*  
## Age.x -8.324e-04 2.400e-03 -0.347 0.728743   
## Age.y 1.116e-02 2.429e-03 4.596 4.31e-06 \*\*\*  
## Gender.xMale 1.632e-01 9.686e-02 1.685 0.091947 .   
## Gender.ymale -9.995e-02 9.662e-02 -1.034 0.300957   
## CoworkerTRUE 1.026e+00 2.866e-01 3.581 0.000342 \*\*\*  
## FamilyTRUE 1.689e+00 2.606e-01 6.482 9.08e-11 \*\*\*  
## FriendTRUE 6.693e-01 3.400e-01 1.969 0.049010 \*   
## MedicalTRUE -1.376e+01 2.290e+02 -0.060 0.952086   
## NeighbourTRUE NA NA NA NA   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 2616.8 on 1963 degrees of freedom  
## Residual deviance: 2466.2 on 1955 degrees of freedom  
## (622 observations deleted due to missingness)  
## AIC: 2484.2  
##   
## Number of Fisher Scoring iterations: 14

m5 <- glm(data=contactsType.m,case ~ Age.y + Coworker + Family + Friend + Medical,family=binomial)  
summary(m5)

##   
## Call:  
## glm(formula = case ~ Age.y + Coworker + Family + Friend + Medical,   
## family = binomial, data = contactsType.m)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -1.3605 -1.0376 -0.7534 1.2162 2.1196   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -2.350536 0.276946 -8.487 < 2e-16 \*\*\*  
## Age.y 0.011369 0.002416 4.706 2.52e-06 \*\*\*  
## CoworkerTRUE 1.050575 0.286130 3.672 0.000241 \*\*\*  
## FamilyTRUE 1.702670 0.260365 6.540 6.17e-11 \*\*\*  
## FriendTRUE 0.707827 0.339150 2.087 0.036883 \*   
## MedicalTRUE -13.690481 229.079091 -0.060 0.952344   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 2616.8 on 1963 degrees of freedom  
## Residual deviance: 2470.5 on 1958 degrees of freedom  
## (622 observations deleted due to missingness)  
## AIC: 2482.5  
##   
## Number of Fisher Scoring iterations: 14

t1 <- table(contact.merge$`Contact type`,contact.merge$TimePeriod)  
chisq.test(t1)

## Warning in chisq.test(t1): Chi-squared approximation may be incorrect

##   
## Pearson's Chi-squared test  
##   
## data: t1  
## X-squared = 135.3, df = 10, p-value < 2.2e-16

ncontacts <- contact.merge %>% group\_by(TimePeriod,newPatID) %>% summarise(n=n())

## `summarise()` regrouping output by 'TimePeriod' (override with `.groups` argument)

kruskal.test(ncontacts$n,ncontacts$TimePeriod)

##   
## Kruskal-Wallis rank sum test  
##   
## data: ncontacts$n and ncontacts$TimePeriod  
## Kruskal-Wallis chi-squared = 17.609, df = 2, p-value = 0.00015

Rates of transmission by type of contact, gender and age of index case - table as in Bi

Contact by age groups and type of contact

##   
## 0-9 10-19 20-29 30-39 40-49 50-59 60-69  
## 0-9 0.8461538 0.2857143 0.2105263 0.5555556 0.4117647 0.3636364 0.5000000  
## 10-19 0.2666667 0.4242424 0.3684211 0.3939394 0.6486486 0.5151515 0.4642857  
## 20-29 0.4000000 0.3750000 0.1785714 0.1481481 0.1794872 0.4074074 0.3500000  
## 30-39 0.3414634 0.3333333 0.3714286 0.3797468 0.2439024 0.3902439 0.4285714  
## 40-49 0.2500000 0.3939394 0.4444444 0.3863636 0.4411765 0.5600000 0.4545455  
## 50-59 0.3636364 0.4583333 0.2857143 0.2698413 0.4193548 0.5000000 0.5625000  
## 60-69 0.2500000 0.3333333 0.3750000 0.2750000 0.4000000 0.3750000 0.3571429  
## 70+ 0.3684211 0.4000000 0.3888889 0.2307692 0.2631579 0.5600000 0.3636364  
##   
## 70+  
## 0-9 0.4285714  
## 10-19 0.6000000  
## 20-29 0.2142857  
## 30-39 0.3750000  
## 40-49 0.4285714  
## 50-59 0.5500000  
## 60-69 0.6000000  
## 70+ 0.7000000

##   
## FALSE TRUE  
## Female 995 397  
## Male 796 398

##   
## Call:  
## glm(formula = case ~ Gender.x, family = binomial, data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.9005 -0.9005 -0.8195 1.4823 1.5840   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.91881 0.05936 -15.478 < 2e-16 \*\*\*  
## Gender.xMale 0.22566 0.08540 2.642 0.00823 \*\*   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3184.3 on 2584 degrees of freedom  
## (296 observations deleted due to missingness)  
## AIC: 3188.3  
##   
## Number of Fisher Scoring iterations: 4

## Waiting for profiling to be done...

## 2.5 % 97.5 %  
## (Intercept) 0.398995 0.3548142 0.4478065  
## Gender.xMale 1.253149 1.0600334 1.4816015

##   
## FALSE TRUE  
## Female 587 389  
## male 622 366

##   
## Call:  
## glm(formula = case ~ Gender.y, family = binomial, data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -1.008 -1.008 -0.962 1.356 1.409   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.41145 0.06538 -6.293 3.11e-10 \*\*\*  
## Gender.ymale -0.11886 0.09281 -1.281 0.2   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 2616.8 on 1963 degrees of freedom  
## Residual deviance: 2615.1 on 1962 degrees of freedom  
## (918 observations deleted due to missingness)  
## AIC: 2619.1  
##   
## Number of Fisher Scoring iterations: 4

## Waiting for profiling to be done...

## 2.5 % 97.5 %  
## (Intercept) 0.6626917 0.5826406 0.7529031  
## Gender.ymale 0.8879310 0.7401554 1.0650244

##   
## FALSE TRUE  
## 0-9 91 62  
## 10-19 157 107  
## 20-29 329 80  
## 30-39 287 116  
## 40-49 260 144  
## 50-59 344 146  
## 60-69 188 75  
## 70+ 135 65

##   
## Call:  
## glm(formula = case ~ relevel(as.factor(AgeGroup.X), ref = "50-59"),   
## family = binomial, data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -1.0195 -0.8411 -0.8194 1.3441 1.8065   
##   
## Coefficients:  
## Estimate Std. Error z value  
## (Intercept) -0.85704 0.09877 -8.677  
## relevel(as.factor(AgeGroup.X), ref = "50-59")0-9 0.47331 0.19203 2.465  
## relevel(as.factor(AgeGroup.X), ref = "50-59")10-19 0.47362 0.15960 2.968  
## relevel(as.factor(AgeGroup.X), ref = "50-59")20-29 -0.55700 0.15905 -3.502  
## relevel(as.factor(AgeGroup.X), ref = "50-59")30-39 -0.04886 0.14786 -0.330  
## relevel(as.factor(AgeGroup.X), ref = "50-59")40-49 0.26617 0.14334 1.857  
## relevel(as.factor(AgeGroup.X), ref = "50-59")60-69 -0.06192 0.16855 -0.367  
## relevel(as.factor(AgeGroup.X), ref = "50-59")70+ 0.12615 0.18041 0.699  
## Pr(>|z|)   
## (Intercept) < 2e-16 \*\*\*  
## relevel(as.factor(AgeGroup.X), ref = "50-59")0-9 0.013709 \*   
## relevel(as.factor(AgeGroup.X), ref = "50-59")10-19 0.003002 \*\*   
## relevel(as.factor(AgeGroup.X), ref = "50-59")20-29 0.000462 \*\*\*  
## relevel(as.factor(AgeGroup.X), ref = "50-59")30-39 0.741070   
## relevel(as.factor(AgeGroup.X), ref = "50-59")40-49 0.063330 .   
## relevel(as.factor(AgeGroup.X), ref = "50-59")60-69 0.713347   
## relevel(as.factor(AgeGroup.X), ref = "50-59")70+ 0.484414   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3141.0 on 2578 degrees of freedom  
## (296 observations deleted due to missingness)  
## AIC: 3157  
##   
## Number of Fisher Scoring iterations: 4

## Waiting for profiling to be done...

## 2.5 %  
## (Intercept) 0.4244186 0.3487783  
## relevel(as.factor(AgeGroup.X), ref = "50-59")0-9 1.6052988 1.0992944  
## relevel(as.factor(AgeGroup.X), ref = "50-59")10-19 1.6057936 1.1739555  
## relevel(as.factor(AgeGroup.X), ref = "50-59")20-29 0.5729275 0.4181990  
## relevel(as.factor(AgeGroup.X), ref = "50-59")30-39 0.9523173 0.7120949  
## relevel(as.factor(AgeGroup.X), ref = "50-59")40-49 1.3049526 0.9853291  
## relevel(as.factor(AgeGroup.X), ref = "50-59")60-69 0.9399592 0.6735959  
## relevel(as.factor(AgeGroup.X), ref = "50-59")70+ 1.1344495 0.7939630  
## 97.5 %  
## (Intercept) 0.5138561  
## relevel(as.factor(AgeGroup.X), ref = "50-59")0-9 2.3359392  
## relevel(as.factor(AgeGroup.X), ref = "50-59")10-19 2.1955482  
## relevel(as.factor(AgeGroup.X), ref = "50-59")20-29 0.7805830  
## relevel(as.factor(AgeGroup.X), ref = "50-59")30-39 1.2718017  
## relevel(as.factor(AgeGroup.X), ref = "50-59")40-49 1.7287973  
## relevel(as.factor(AgeGroup.X), ref = "50-59")60-69 1.3050672  
## relevel(as.factor(AgeGroup.X), ref = "50-59")70+ 1.6117746

##   
## FALSE TRUE  
## 0-9 102 59  
## 10-19 99 63  
## 20-29 231 104  
## 30-39 249 117  
## 40-49 192 121  
## 50-59 171 148  
## 60-69 107 85  
## 70+ 58 58

##   
## Call:  
## glm(formula = case ~ relevel(as.factor(AgeGroup.Y), ref = "50-59"),   
## family = binomial, data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -1.1774 -0.9886 -0.8622 1.2766 1.5295   
##   
## Coefficients:  
## Estimate Std. Error z value  
## (Intercept) -0.14445 0.11227 -1.287  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")0-9 -0.40298 0.19839 -2.031  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")10-19 -0.30753 0.19641 -1.566  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")20-29 -0.65358 0.16294 -4.011  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")30-39 -0.61083 0.15864 -3.850  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")40-49 -0.31725 0.16149 -1.965  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")60-69 -0.08573 0.18362 -0.467  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")70+ 0.14445 0.21700 0.666  
## Pr(>|z|)   
## (Intercept) 0.198222   
## relevel(as.factor(AgeGroup.Y), ref = "50-59")0-9 0.042225 \*   
## relevel(as.factor(AgeGroup.Y), ref = "50-59")10-19 0.117410   
## relevel(as.factor(AgeGroup.Y), ref = "50-59")20-29 6.04e-05 \*\*\*  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")30-39 0.000118 \*\*\*  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")40-49 0.049461 \*   
## relevel(as.factor(AgeGroup.Y), ref = "50-59")60-69 0.640589   
## relevel(as.factor(AgeGroup.Y), ref = "50-59")70+ 0.505613   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 2616.8 on 1963 degrees of freedom  
## Residual deviance: 2584.5 on 1956 degrees of freedom  
## (918 observations deleted due to missingness)  
## AIC: 2600.5  
##   
## Number of Fisher Scoring iterations: 4

## Waiting for profiling to be done...

## 2.5 %  
## (Intercept) 0.8654971 0.6939851  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")0-9 0.6683227 0.4514796  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")10-19 0.7352580 0.4989521  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")20-29 0.5201825 0.3773293  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")30-39 0.5429013 0.3972843  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")40-49 0.7281461 0.5301109  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")60-69 0.9178454 0.6397592  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")70+ 1.1554054 0.7547533  
## 97.5 %  
## (Intercept) 1.0781377  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")0-9 0.9835125  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")10-19 1.0784814  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")20-29 0.7149624  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")30-39 0.7401734  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")40-49 0.9986995  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")60-69 1.3148173  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")70+ 1.7692870

##   
## FALSE TRUE  
## Co-worker 304 93  
## Family member 1111 655  
## Friend 120 28  
## Medical Personnel 63 0  
## Neighbour 193 19  
## unknown 0 0

##   
## Call:  
## glm(formula = case ~ relevel(as.factor(`Contact type`), ref = "Family member"),   
## family = binomial, data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.9628 -0.9628 -0.7306 1.4084 2.1964   
##   
## Coefficients:  
## Estimate  
## (Intercept) -0.52838  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker -0.65605  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend -0.92691  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel -16.03769  
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour -1.78987  
## Std. Error  
## (Intercept) 0.04926  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker 0.12833  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend 0.21558  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel 302.31422  
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour 0.24544  
## z value  
## (Intercept) -10.726  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker -5.112  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend -4.300  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel -0.053  
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour -7.293  
## Pr(>|z|)  
## (Intercept) < 2e-16  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker 3.19e-07  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend 1.71e-05  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel 0.958  
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour 3.04e-13  
##   
## (Intercept) \*\*\*  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker \*\*\*  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend \*\*\*  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel   
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour \*\*\*  
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3032.8 on 2581 degrees of freedom  
## (296 observations deleted due to missingness)  
## AIC: 3042.8  
##   
## Number of Fisher Scoring iterations: 15

## Waiting for profiling to be done...

## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred

##   
## (Intercept) 5.895590e-01  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker 5.188982e-01  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend 3.957761e-01  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel 1.083729e-07  
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour 1.669818e-01  
## 2.5 %  
## (Intercept) 5.350715e-01  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker 4.017780e-01  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend 2.546915e-01  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel 3.299302e-54  
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour 9.996608e-02  
## 97.5 %  
## (Intercept) 0.64907297  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker 0.66478107  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend 0.59483413  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel 0.01123575  
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour 0.26303306

##   
## Call:  
## glm(formula = case ~ relevel(as.factor(AgeGroup.Y), ref = "50-59") +   
## relevel(as.factor(AgeGroup.X), ref = "50-59") + relevel(as.factor(`Contact type`),   
## ref = "Family member") + Gender.x + Gender.y, family = binomial,   
## data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -1.4315 -1.0107 -0.6933 1.1916 2.1920   
##   
## Coefficients:  
## Estimate  
## (Intercept) 0.18402  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")0-9 -0.59586  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")10-19 -0.48623  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")20-29 -0.63050  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")30-39 -0.55083  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")40-49 -0.25121  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")60-69 -0.12486  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")70+ 0.04151  
## relevel(as.factor(AgeGroup.X), ref = "50-59")0-9 0.07113  
## relevel(as.factor(AgeGroup.X), ref = "50-59")10-19 0.16417  
## relevel(as.factor(AgeGroup.X), ref = "50-59")20-29 -0.49689  
## relevel(as.factor(AgeGroup.X), ref = "50-59")30-39 -0.30459  
## relevel(as.factor(AgeGroup.X), ref = "50-59")40-49 0.16928  
## relevel(as.factor(AgeGroup.X), ref = "50-59")60-69 -0.27759  
## relevel(as.factor(AgeGroup.X), ref = "50-59")70+ -0.09651  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker -0.63193  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend -0.95827  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel -16.20272  
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour -1.66316  
## Gender.xMale 0.18507  
## Gender.ymale -0.08229  
## Std. Error  
## (Intercept) 0.17011  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")0-9 0.20489  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")10-19 0.20529  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")20-29 0.16948  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")30-39 0.16539  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")40-49 0.16944  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")60-69 0.19212  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")70+ 0.22574  
## relevel(as.factor(AgeGroup.X), ref = "50-59")0-9 0.21555  
## relevel(as.factor(AgeGroup.X), ref = "50-59")10-19 0.18062  
## relevel(as.factor(AgeGroup.X), ref = "50-59")20-29 0.18203  
## relevel(as.factor(AgeGroup.X), ref = "50-59")30-39 0.16634  
## relevel(as.factor(AgeGroup.X), ref = "50-59")40-49 0.16628  
## relevel(as.factor(AgeGroup.X), ref = "50-59")60-69 0.19216  
## relevel(as.factor(AgeGroup.X), ref = "50-59")70+ 0.20871  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker 0.15204  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend 0.23603  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel 372.52862  
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour 0.26313  
## Gender.xMale 0.09810  
## Gender.ymale 0.09781  
## z value  
## (Intercept) 1.082  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")0-9 -2.908  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")10-19 -2.369  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")20-29 -3.720  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")30-39 -3.331  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")40-49 -1.483  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")60-69 -0.650  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")70+ 0.184  
## relevel(as.factor(AgeGroup.X), ref = "50-59")0-9 0.330  
## relevel(as.factor(AgeGroup.X), ref = "50-59")10-19 0.909  
## relevel(as.factor(AgeGroup.X), ref = "50-59")20-29 -2.730  
## relevel(as.factor(AgeGroup.X), ref = "50-59")30-39 -1.831  
## relevel(as.factor(AgeGroup.X), ref = "50-59")40-49 1.018  
## relevel(as.factor(AgeGroup.X), ref = "50-59")60-69 -1.445  
## relevel(as.factor(AgeGroup.X), ref = "50-59")70+ -0.462  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker -4.156  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend -4.060  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel -0.043  
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour -6.321  
## Gender.xMale 1.887  
## Gender.ymale -0.841  
## Pr(>|z|)  
## (Intercept) 0.279352  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")0-9 0.003635  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")10-19 0.017859  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")20-29 0.000199  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")30-39 0.000867  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")40-49 0.138184  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")60-69 0.515766  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")70+ 0.854098  
## relevel(as.factor(AgeGroup.X), ref = "50-59")0-9 0.741393  
## relevel(as.factor(AgeGroup.X), ref = "50-59")10-19 0.363376  
## relevel(as.factor(AgeGroup.X), ref = "50-59")20-29 0.006339  
## relevel(as.factor(AgeGroup.X), ref = "50-59")30-39 0.067092  
## relevel(as.factor(AgeGroup.X), ref = "50-59")40-49 0.308678  
## relevel(as.factor(AgeGroup.X), ref = "50-59")60-69 0.148585  
## relevel(as.factor(AgeGroup.X), ref = "50-59")70+ 0.643785  
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker 3.23e-05  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend 4.91e-05  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel 0.965308  
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour 2.60e-10  
## Gender.xMale 0.059224  
## Gender.ymale 0.400199  
##   
## (Intercept)   
## relevel(as.factor(AgeGroup.Y), ref = "50-59")0-9 \*\*   
## relevel(as.factor(AgeGroup.Y), ref = "50-59")10-19 \*   
## relevel(as.factor(AgeGroup.Y), ref = "50-59")20-29 \*\*\*  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")30-39 \*\*\*  
## relevel(as.factor(AgeGroup.Y), ref = "50-59")40-49   
## relevel(as.factor(AgeGroup.Y), ref = "50-59")60-69   
## relevel(as.factor(AgeGroup.Y), ref = "50-59")70+   
## relevel(as.factor(AgeGroup.X), ref = "50-59")0-9   
## relevel(as.factor(AgeGroup.X), ref = "50-59")10-19   
## relevel(as.factor(AgeGroup.X), ref = "50-59")20-29 \*\*   
## relevel(as.factor(AgeGroup.X), ref = "50-59")30-39 .   
## relevel(as.factor(AgeGroup.X), ref = "50-59")40-49   
## relevel(as.factor(AgeGroup.X), ref = "50-59")60-69   
## relevel(as.factor(AgeGroup.X), ref = "50-59")70+   
## relevel(as.factor(`Contact type`), ref = "Family member")Co-worker \*\*\*  
## relevel(as.factor(`Contact type`), ref = "Family member")Friend \*\*\*  
## relevel(as.factor(`Contact type`), ref = "Family member")Medical Personnel   
## relevel(as.factor(`Contact type`), ref = "Family member")Neighbour \*\*\*  
## Gender.xMale .   
## Gender.ymale   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 2616.8 on 1963 degrees of freedom  
## Residual deviance: 2436.6 on 1943 degrees of freedom  
## (918 observations deleted due to missingness)  
## AIC: 2478.6  
##   
## Number of Fisher Scoring iterations: 15

## Waiting for profiling to be done...

## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
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## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred  
  
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred

##   
## Call:  
## glm(formula = case ~ TimePeriod, family = binomial, data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.8772 -0.8772 -0.8672 1.5109 1.7101   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.78418 0.06167 -12.716 <2e-16 \*\*\*  
## TimePeriodPre-lockdown -0.41452 0.19508 -2.125 0.0336 \*   
## TimePeriodRestrictions easing 0.02744 0.09354 0.293 0.7693   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 2867.7 on 2315 degrees of freedom  
## Residual deviance: 2862.3 on 2313 degrees of freedom  
## (566 observations deleted due to missingness)  
## AIC: 2868.3  
##   
## Number of Fisher Scoring iterations: 4

## Waiting for profiling to be done...

## 2.5 % 97.5 %  
## (Intercept) 0.4564958 0.4041414 0.5146946  
## TimePeriodPre-lockdown 0.6606573 0.4455560 0.9591360  
## TimePeriodRestrictions easing 1.0278173 0.8553913 1.2343759

##   
## Call:  
## glm(formula = case ~ TimePeriod + family + contactage40, family = binomial,   
## data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -1.2626 -1.0516 -0.7303 1.1952 1.9269   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.30323 0.09113 -3.327 0.000877 \*\*\*  
## TimePeriodPre-lockdown -0.16064 0.22088 -0.727 0.467062   
## TimePeriodRestrictions easing -0.23999 0.10475 -2.291 0.021960 \*   
## familyOther -1.14347 0.12411 -9.214 < 2e-16 \*\*\*  
## contactage40Contact ≥ 40 0.50136 0.10147 4.941 7.78e-07 \*\*\*  
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 2354.9 on 1764 degrees of freedom  
## Residual deviance: 2235.9 on 1760 degrees of freedom  
## (1117 observations deleted due to missingness)  
## AIC: 2245.9  
##   
## Number of Fisher Scoring iterations: 4

## Waiting for profiling to be done...

## 2.5 % 97.5 %  
## (Intercept) 0.7384260 0.6172424 0.8824213  
## TimePeriodPre-lockdown 0.8515979 0.5484531 1.3066016  
## TimePeriodRestrictions easing 0.7866323 0.6403448 0.9655930  
## familyOther 0.3187118 0.2490004 0.4051700  
## contactage40Contact ≥ 40 1.6509624 1.3537543 2.0152592

##   
## Call:  
## glm(formula = case ~ family, family = binomial, data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -0.9628 -0.9628 -0.6119 1.4084 1.8802   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.52838 0.04926 -10.73 <2e-16 \*\*\*  
## familyOther -1.05207 0.10507 -10.01 <2e-16 \*\*\*  
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 3191.3 on 2585 degrees of freedom  
## Residual deviance: 3078.7 on 2584 degrees of freedom  
## (296 observations deleted due to missingness)  
## AIC: 3082.7  
##   
## Number of Fisher Scoring iterations: 4

## Waiting for profiling to be done...

## 2.5 % 97.5 %  
## (Intercept) 0.5895590 0.5350716 0.6490729  
## familyOther 0.3492142 0.2834018 0.4279470

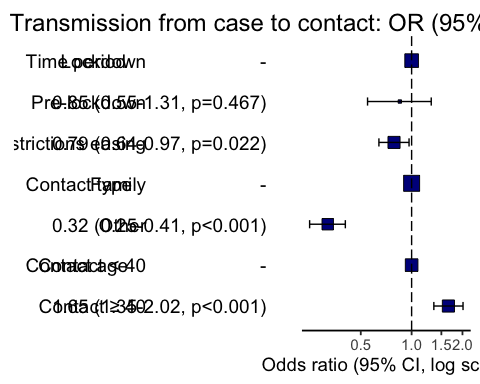
##   
## Call:  
## glm(formula = case ~ contactage40, family = binomial, data = contact.merge)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -1.0740 -1.0740 -0.9032 1.2844 1.4790   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) -0.68583 0.06621 -10.358 < 2e-16 \*\*\*  
## contactage40Contact ≥ 40 0.43776 0.09330 4.692 2.71e-06 \*\*\*  
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 2616.8 on 1963 degrees of freedom  
## Residual deviance: 2594.6 on 1962 degrees of freedom  
## (918 observations deleted due to missingness)  
## AIC: 2598.6  
##   
## Number of Fisher Scoring iterations: 4

## Waiting for profiling to be done...

## 2.5 % 97.5 %  
## (Intercept) 0.5036711 0.4419412 0.5729586  
## contactage40Contact ≥ 40 1.5492314 1.2906599 1.8607125

## Note: dependent includes missing data. These are dropped.  
## Waiting for profiling to be done...  
## Waiting for profiling to be done...  
## Waiting for profiling to be done...

## Warning: Removed 3 rows containing missing values (geom\_errorbarh).



##   
## FALSE TRUE   
## 2244 638

##   
## FALSE TRUE   
## 1041 301

## [1] 477

## [1] "Patient67" "Patient82" "3003058146" "3003059349" "3003141462"   
## [6] "3003148320" "3003172260" "3003207687" "3006018336" "3006024483"   
## [11] "3006029058" "3006044910" "3006053838" "3012024279" "3012025857"   
## [16] "3015124239" "3018025674" "3021046953" "3024000153" "3024023619"   
## [21] "3024040392" "3024053589" "3024056673" "3024069246" "3024112866"   
## [26] "3024120477" "3024136563" "3024152832" "3024163893" "3024168744"   
## [31] "3024191136" "3027009930" "3027035475" "3027046971" "3027057063"   
## [36] "3030019146" "3033013719" "3033153999" "3033220740" "3033259833"   
## [41] "3036044646" "3039012912" "3045021186" "3045044127" "3045049719"   
## [46] "3045050997" "3051004647" "3051058518" "3051136236" "3051151308"   
## [51] "3057025608" "3057040344" "3057044310" "3057044634" "3057059160"   
## [56] "3057061776" "3057077517" "3057085731" "3057089475" "3057090009"   
## [61] "3057139188" "3057153495" "3057155481" "3057171096" "3057187374"   
## [66] "3057203508" "3057221562" "3063001002" "3063001005" "3063031620"   
## [71] "3069020478" "3069029226" "3072026562" "3072038742" "3072040230"   
## [76] "3072056715" "3072059835" "3072105732" "3072118722" "3072129885"   
## [81] "3072132774" "3072174828" "3072209988" "3072211455" "3072214428"   
## [86] "3072218496" "3072226170" "3072237285" "3072241458" "3072258777"   
## [91] "3072259242" "3075004287" "3075013386" "3075038064" "3075046803"   
## [96] "3075060396" "3075062523" "3078001836" "3078031023" "3078036651"   
## [101] "3081014892" "3081051255" "3081092907" "3081124233" "3081175524"   
## [106] "3081189708" "3081216093" "3081222003" "3081262005" "3081268362"   
## [111] "3084011442" "3087018237" "3087022986" "3087023823" "3087034347"   
## [116] "3090050982" "3090090432" "3090101577" "3090117072" "3096006654"   
## [121] "3351184671" "3450404439" "3462061191" "3624193929" "4050010950"   
## [126] "4050062007" "4050243972" "4050469446" "4053017793" "4572275604"   
## [131] "4650032940" "4650065676" "4950350193" "5424204081" "5850450534"   
## [136] "6003013191" "9003000873" "12003008496" "21003020355" "21003047856"   
## [141] "27003029100" "30003007287" "30003007524" "30003015507" "30003021264"   
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## [156] "30003058506" "30003059328" "30003060846" "30003061866" "30003062631"   
## [161] "30003064608" "30003067968" "30003069249" "30003071544" "30003074151"   
## [166] "30003075570" "30003076614" "30003079542" "30003081621" "30003084684"   
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## [186] "30003131436" "30003139374" "30003150873" "30003156855" "30003164442"   
## [191] "30003166635" "30003170373" "30003175545" "30003180099" "30003182256"   
## [196] "30003187521" "30003196587" "30003198507" "30003200334" "30003202710"   
## [201] "30003204366" "30003205533" "30003207693" "30603218280" "30903216966"   
## [206] "30903219567" "30903223851" "31503228171" "31803222882" "31803223722"   
## [211] "31950006540" "31950008547" "32250011487" "33003006675" "36003051501"   
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## [236] "45003075642" "45003075669" "48003023871" "51003027357" "51003044118"   
## [241] "51303105780" "52803105783" "54003019803" "54003061542" "54003219570"   
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## [251] "57003301728" "57003303930" "57903356802" "59103346467" "59403339708"   
## [256] "60003135678" "63003002433" "63003003648" "63003012027" "63003024117"   
## [261] "63003088107" "63003105408" "63003116829" "63003121497" "64050000618"   
## [266] "66003001488" "66003001575" "66003002874" "66003008493" "66003014673"   
## [271] "66003018975" "66003020490" "66003021756" "66003024507" "66003024546"   
## [276] "66003026712" "66003029241" "66003031626" "66003031674" "66003031821"   
## [281] "66003032760" "66003036345" "66003036393" "66003051246" "66003051672"   
## [286] "66003052950" "66003062586" "66003065223" "66003065388" "66003065613"   
## [291] "66003066201" "66003066630" "66003067497" "66003075054" "66603077121"   
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## [301] "78003090681" "81003000198" "81003020229" "81003020580" "84003014235"   
## [306] "84003030930" "84003058593" "84003059784" "84003065943" "84003071850"   
## [311] "84003091161" "84003096513" "84003127812" "84003160578" "84003190896"   
## [316] "84003199683" "84003218454" "84003219792" "84003228684" "84003243798"   
## [321] "84003270375" "84003275748" "84003326538" "84003329757" "84003330054"   
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## [331] "90003027570" "93003086550" "93003145224" "93003152451" "96003031449"   
## [336] "99003101493" "105003082608" "105003100266" "105003136428" "105003187071"  
## [341] "105003349326" "105003367257" "105603414903" "108003040176" "108003155238"  
## [346] "108903164925" "111003024042" "111003173889" "114003022143" "114003067008"  
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## [356] "120003000537" "123003021459" "126003001083" "126003090960" "126003117246"  
## [361] "129003016377" "129003021654" "129003056073" "129003072453" "129003114078"  
## [366] "129003116853" "129003118494" "129003129390" "132003014655" "135003093843"  
## [371] "138003012369" "139503077091" "141003020097" "141003110829" "153003005922"  
## [376] "153003091437" "156003053160" "156003059619" "159003039630" "159003047526"  
## [381] "159003060075" "159003114750" "159003132624" "161103190311" "162003032583"  
## [386] "162003034494" "162003044619" "162303190323" "165003021231" "165003026643"  
## [391] "168003011856" "168003034428" "171003054690" "171003153225" "174003094035"  
## [396] "177003048717" "177003075666" "177003206532" "177003240558" "177003274152"  
## [401] "177003298596" "177003305916" "177003325683" "177003339195" "177003377244"  
## [406] "177006019611" "178650045444" "180003059649" "180003070857" "180003253230"  
## [411] "180003264117" "180003278700" "180003309201" "180003313143" "180003465351"  
## [416] "180006010383" "180009037254" "180750065244" "182850155871" "183003042123"  
## [421] "183003060822" "183003084651" "183003148443" "183003225438" "183009000723"  
## [426] "183009015648" "183012060819" "183012096921" "183012114147" "183012140223"  
## [431] "183012186249" "183012186393" "183012206643" "183012207753" "183012217257"  
## [436] "183015025128" "183018025842" "183018065514" "183018072318" "183018104880"  
## [441] "183018212865" "183018222405" "183018238551" "183021025011" "183027059913"  
## [446] "183027082626" "184953008154" "185412228642" "185850084405" "185850161457"  
## [451] "186003001227" "186003008565" "186003015066" "186003059331" "186003069087"  
## [456] "186003069174" "186003111840" "186009000858" "186009005184" "186009005556"  
## [461] "186009005973" "186009029367" "186009040749" "186009041289" "186009049686"  
## [466] "186009049689" "186012050091" "186012055092" "186015064962" "186018017091"  
## [471] "186018039042" "186018045687" "186018074922" "186018088236" "186018185553"  
## [476] "186021031728" "186021031731" "186306044766" "195024004602" "Patient1"   
## [481] "Patient10" "Patient11" "Patient12" "Patient13" "Patient14"   
## [486] "Patient15" "Patient16" "Patient17" "Patient18" "Patient19"   
## [491] "Patient2" "Patient20" "Patient21" "Patient22" "Patient23"   
## [496] "Patient24" "Patient25" "Patient27" "Patient28" "Patient29"   
## [501] "Patient3" "Patient30" "Patient31" "Patient32" "Patient33"   
## [506] "Patient34" "Patient35" "Patient36" "Patient37" "Patient38"   
## [511] "Patient39" "Patient4" "Patient40" "Patient41" "Patient42"   
## [516] "Patient43" "Patient44" "Patient45" "Patient46" "Patient48"   
## [521] "Patient49" "Patient5" "Patient50" "Patient132" "Patient52"   
## [526] "Patient53" "Patient9" "Patient55" "Patient56" "Patient58"   
## [531] "Patient59" "Patient6" "Patient60" "Patient61" "Patient62"   
## [536] "Patient63" "Patient64" "Patient65" "Patient66" "Patient68"   
## [541] "Patient69" "Patient7" "Patient70" "Patient71" "Patient72"   
## [546] "Patient74" "Patient75" "Patient76" "Patient77" "Patient78"   
## [551] "Patient79" "Patient8" "Patient80" "Patient81" "Patient83"   
## [556] "Patient84" "Patient86" "Patient87" "Patient88" "Patient89"   
## [561] "Patient73" "Patient85" "Patient90" "Patient91" "Patient92"   
## [566] "Patient93" "Patient94" "Patient95" "Patient96" "Patient97"   
## [571] "Patient98" "Patient99" "Patient100" "Patient101" "Patient102"   
## [576] "Patient103" "Patient104" "Patient105" "Patient106" "Patient107"   
## [581] "Patient108" "Patient109" "Patient110" "Patient111" "Patient112"   
## [586] "Patient113" "Patient114" "Patient115" "Patient116" "Patient117"   
## [591] "Patient118" "Patient119" "Patient120" "Patient121" "Patient122"   
## [596] "Patient123" "Patient124" "Patient125" "Patient126" "Patient127"   
## [601] "Patient128" "Patient130" "Patient131" "Patient133" "Patient134"

##   
## 1-sample proportions test with continuity correction  
##   
## data: 301 out of 477, null probability 0.5  
## X-squared = 32.235, df = 1, p-value = 1.366e-08  
## alternative hypothesis: true p is not equal to 0.5  
## 95 percent confidence interval:  
## 0.5857739 0.6741366  
## sample estimates:  
## p   
## 0.6310273

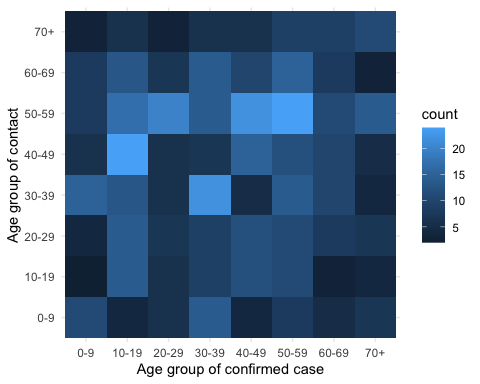
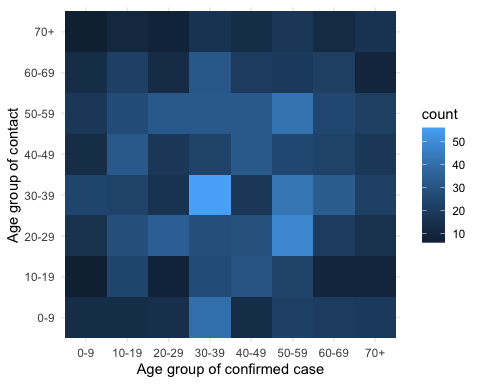
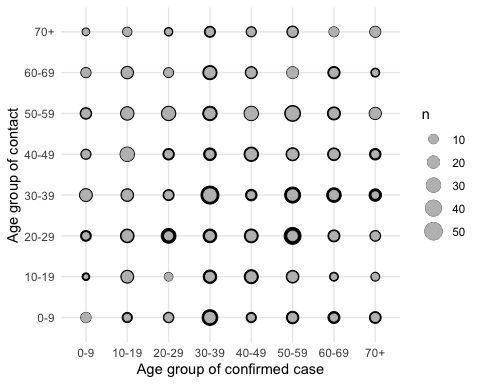
##   
## 1-sample proportions test with continuity correction  
##   
## data: 301 out of 1342, null probability 0.5  
## X-squared = 406.95, df = 1, p-value < 2.2e-16  
## alternative hypothesis: true p is not equal to 0.5  
## 95 percent confidence interval:  
## 0.2024221 0.2477623  
## sample estimates:  
## p   
## 0.2242921

## Patient unique ID Date of Birth Age.x Gender.x   
## NA NA NA NA   
## გამოსავალი Contact ID Contact type Confirmed Case   
## NA NA NA NA   
## missingPatID newPatID missingCID newCID   
## NA NA NA NA   
## case Date Gender.y Age.y   
## NA NA NA NA   
## TimePeriod AgeGroup.X AgeGroup.Y family   
## NA NA NA NA   
## contactage40 knownID onwards not\_onwards   
## NA NA NA NA   
## not\_onwards2 Age.of.case Age.of.contact Contact.type   
## NA "Age of case" "Age of contact" "Contact type"   
## Gender.of.case Gender.of.contact   
## "Gender of case" "Gender of contact"

## Note: dependent includes missing data. These are dropped.  
## Waiting for profiling to be done...  
## Waiting for profiling to be done...  
## Waiting for profiling to be done...

## Warning: Removed 5 rows containing missing values (geom\_errorbarh).

## quartz\_off\_screen   
## 2

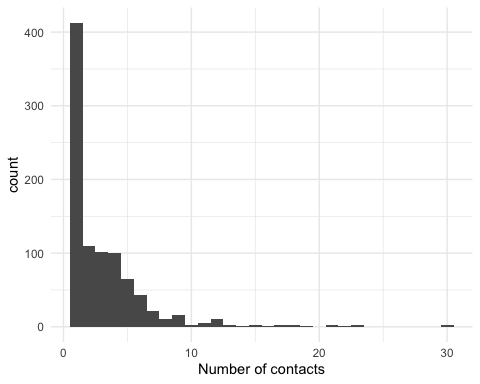
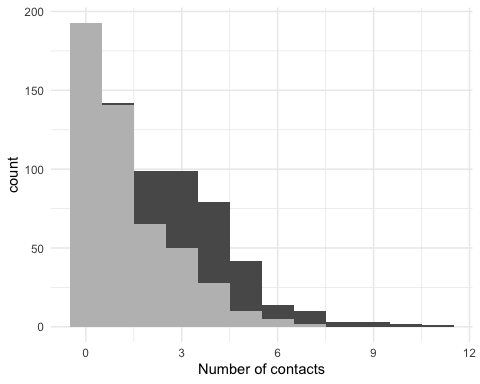
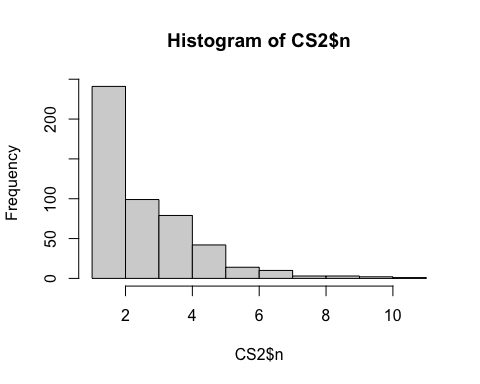


## `summarise()` regrouping output by 'AgeGroup.X' (override with `.groups` argument)

## quartz\_off\_screen   
## 2

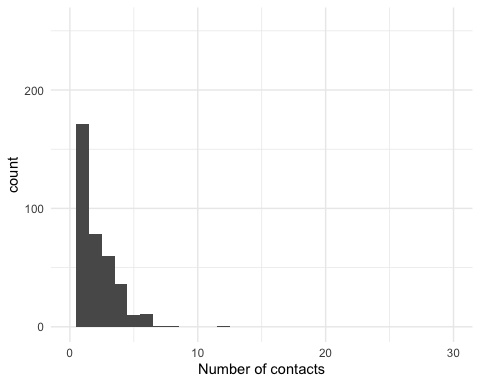
## `summarise()` ungrouping output (override with `.groups` argument)

## `summarise()` ungrouping output (override with `.groups` argument)



## Warning: Removed 291 rows containing non-finite values (stat\_bin).

## Warning: Removed 2 rows containing missing values (geom\_bar).



##   
## Call:  
## glm(formula = case ~ AgeGroup.X \* AgeGroup.Y, family = binomial,   
## data = contact.mergeF)  
##   
## Deviance Residuals:   
## Min 1Q Median 3Q Max   
## -1.3744 -1.0895 -0.8557 1.1586 1.5377   
##   
## Coefficients:  
## Estimate Std. Error z value Pr(>|z|)   
## (Intercept) 0.24741 0.17676 1.400 0.1616   
## AgeGroup.X0-19 0.20458 0.27928 0.733 0.4638   
## AgeGroup.X20-39 -0.45770 0.26412 -1.733 0.0831 .   
## AgeGroup.X60+ -0.40868 0.27842 -1.468 0.1422   
## AgeGroup.Y0-19 -0.66235 0.28050 -2.361 0.0182 \*   
## AgeGroup.Y20-39 -1.06362 0.25608 -4.153 3.28e-05 \*\*\*  
## AgeGroup.Y60+ 0.04027 0.29929 0.135 0.8930   
## AgeGroup.X0-19:AgeGroup.Y0-19 0.42548 0.44463 0.957 0.3386   
## AgeGroup.X20-39:AgeGroup.Y0-19 0.43901 0.40533 1.083 0.2788   
## AgeGroup.X60+:AgeGroup.Y0-19 0.13047 0.45156 0.289 0.7726   
## AgeGroup.X0-19:AgeGroup.Y20-39 0.65608 0.39598 1.657 0.0975 .   
## AgeGroup.X20-39:AgeGroup.Y20-39 0.55829 0.37140 1.503 0.1328   
## AgeGroup.X60+:AgeGroup.Y20-39 0.48131 0.40340 1.193 0.2328   
## AgeGroup.X0-19:AgeGroup.Y60+ -0.13558 0.46613 -0.291 0.7712   
## AgeGroup.X20-39:AgeGroup.Y60+ -0.03970 0.43411 -0.091 0.9271   
## AgeGroup.X60+:AgeGroup.Y60+ 0.25914 0.45293 0.572 0.5672   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1  
##   
## (Dispersion parameter for binomial family taken to be 1)  
##   
## Null deviance: 1924.4 on 1398 degrees of freedom  
## Residual deviance: 1863.5 on 1383 degrees of freedom  
## (367 observations deleted due to missingness)  
## AIC: 1895.5  
##   
## Number of Fisher Scoring iterations: 4

## Waiting for profiling to be done...

## 2.5 % 97.5 %  
## (Intercept) 1.2807018 0.9071674 1.8170074  
## AgeGroup.X0-19 1.2270059 0.7111918 2.1298061  
## AgeGroup.X20-39 0.6327350 0.3758128 1.0599077  
## AgeGroup.X60+ 0.6645293 0.3837559 1.1451644  
## AgeGroup.Y0-19 0.5156371 0.2958559 0.8901902  
## AgeGroup.Y20-39 0.3452055 0.2076507 0.5674368  
## AgeGroup.Y60+ 1.0410959 0.5797955 1.8793717  
## AgeGroup.X0-19:AgeGroup.Y0-19 1.5303222 0.6408140 3.6701764  
## AgeGroup.X20-39:AgeGroup.Y0-19 1.5511730 0.7010542 3.4392010  
## AgeGroup.X60+:AgeGroup.Y0-19 1.1393672 0.4675118 2.7532988  
## AgeGroup.X0-19:AgeGroup.Y20-39 1.9272268 0.8867180 4.1932525  
## AgeGroup.X20-39:AgeGroup.Y20-39 1.7476828 0.8447866 3.6269902  
## AgeGroup.X60+:AgeGroup.Y20-39 1.6181857 0.7327620 3.5685390  
## AgeGroup.X0-19:AgeGroup.Y60+ 0.8732057 0.3500338 2.1823899  
## AgeGroup.X20-39:AgeGroup.Y60+ 0.9610787 0.4091877 2.2482031  
## AgeGroup.X60+:AgeGroup.Y60+ 1.2958212 0.5329579 3.1533394

Transmission by age groups and type of contact

Total number of secondary cases per primary case

##   
## FALSE TRUE   
## 1791 795

Plot of contact network

## Discussion

* main outcomes
* limitations of study
* comparison to other contact tracing studies (Bi, etc.)
* comparison to other countries in the region
* understanding how and why the response was able to be so strong (eg HCV testing and treatment infrastructure)