Vaccine Effectiveness in Catalonia

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16 12 2021

Introduction

In this work data from the COVID19 pandemic in Catalonia from 2021 will be analysed. Specifically, the relationship between the disease severity and vaccine status will be investigated. Approximately in the summer of 2021 European governments started to vaccinate persons with higher risk of a severe COVID19 infection. Since this data is purely observatory, this constitutes a study of vaccine effectiveness compared to vaccine efficacy, which was done in clearly designed randomized-control studies done for clinical admission.

Data Preparation

Since this is a real data set it naturally does not come precisely in the format our analysis expects it to be. Therefore, we first have to prepare and also translate it.

```
data = read.csv("Impacte_del_COVID.csv", encoding="UTF-8")
colnames(df) = c("sex", "age", "date", "event", "vaccinated", "count")
df$sex = as.factor(df$sex)
                                         # male, female, NaN
df$age = as.factor(df$age)
                                         # discretizied for some reason
df$event = as.factor(df$event)
                                         # positive, hospitalized, critical (ICU)
df$date = as.Date(df$date,"%d/%m/%Y")
                                         # now in format Year, month, day
df$vaccinated = as.factor(df$vaccinated) # none, partial, full
df$count = as.numeric(df$count) # this means #count people with these characteristics have been observ
# the task asks us to only analyze people older than 30
df = subset(df, age=="30 a 39" | age == "40 a 49" | age == "50 a 59" |
         age == "60 a 69" | age == "70 a 79" | age == "80 o més")
# also only between 1st March 2020 and 12. December 2021
df = subset(df, date>"2021-03-01" | date > "2021-12-12")
dim(df)
```

[1] 17059 6

The dataset captures these four event types.

```
table(df$event) # dependent variable
```

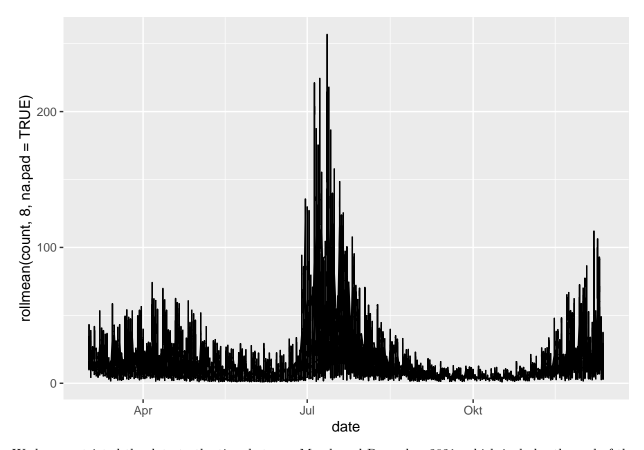
And shows the relationship to vaccine status.

table(df\$vaccinated)

```
## Completa No iniciada Parcial ## 5614 7981 3464
```

Descriptive Analysis

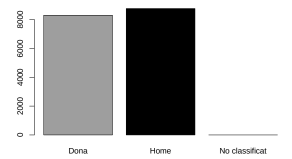
Firstly, let's plot the sum of all captured events as a function of time.

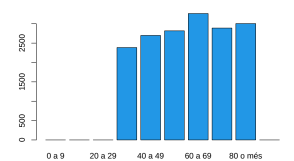


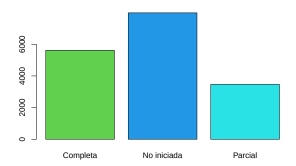
We have restricted the data to the time between March and December 2021, which includes the end of the Spring's wave but especially the big wave of infections in July as well as the beginning of the winter wave starting in November.

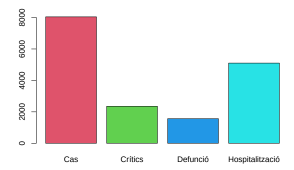
Exploratory Plots

```
plot(df$sex,col=8:9)
plot(df$age,col=4)
plot(df$vaccinated,col=3:5)
plot(df$event,col=2:6)
```









Slightly more male persons were observed than females. Since we restricted the data only to individuals older than 30, the age groups are quite balanced, with older people occurring slightly more often. People with a COVID related event were most of the time not vaccinated at all. The least common case was being partially vaccinated, which makes sense, since this status only held for around two months. Events captured in the dataset occurred by their severity: quarantine, hospitalization, critical (ICU), and death.

Critical Patients over time

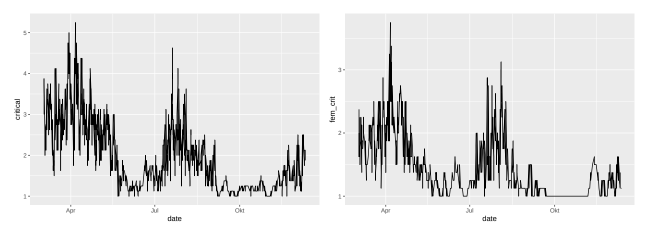
```
# males
df_critial_male = df[df$sex=="Home" & df$event=="Critics",]
critical = rollmean(df_critial_male$count,8,na.pad=TRUE)
ggplot( data = df_critial_male, aes( date, critical, vaccinated )) + geom_line()
```

Warning: Removed 4 row(s) containing missing values (geom_path).

```
#females
df_critial_female = df[df$sex=="Dona" & df$event=="Critics",]
```

```
fem_crit = rollmean(df_critial_female$count,8,na.pad=TRUE)
ggplot( data = df_critial_female, aes( date, fem_crit, vaccinated )) + geom_line()
```

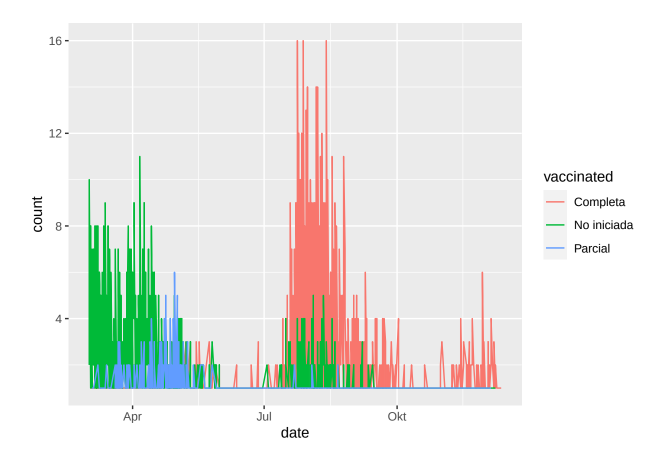
Warning: Removed 2 row(s) containing missing values (geom_path).



On the left we see the development over time of male persons with a critical case, and on the right females on the ICU. To make the plot smoother amoving average over 8 days was calculated to tackle day-of-the-week effects. The three waves are clearly visible.

In the following plot we can see the development in dependency on the vaccine status.

```
df_deaths = df[df$event=="Defunció",]
ggplot(df_deaths,aes(x=date,y=count, color=vaccinated)) + geom_line()
```



Data Analysis

Contingency Table

A contingency table holds a count variable dependent on two (multilevel) factors. To build it we loop over the dataset, check the variables of interest and sum up the count variable and save it in a new dataframe.

Vaccine Status and Infection Gravity

Firstly, we investigate the relationship between vaccine status and the gravity of a coronavirus infection.

```
# init dataframe for count sums
table_vaccine = data.frame(matrix(0,ncol=3,nrow=4))
colnames(table_vaccine) = c("none","partial","full")  # vaccine status
rownames(table_vaccine) = c("casa","hospital","critical","dead")  # event type
for (i in 1:dim(df)[1]){
   content_i = df[i,]

   age = content_i[2]
   vacc = content_i[5]
   event = content_i[4]
   count = content_i[6]
```

```
# define row and col variable, where count should be increased
if (event=="Cas") row=1
if (event=="Hospitalització") row = 2
if (event=="Crítics") row = 3
if (event=="Defunció") row = 4

if (vacc=="No iniciada") col =1
if (vacc=="Parcial") col =2
if (vacc=="Completa") col =3

table_vaccine[row,col] = count + table_vaccine[row,col]
}
table_vaccine
```

```
## casa 138487 24429 88167
## hospital 14323 1842 6243
## critical 3228 342 827
## dead 1513 336 1391
```

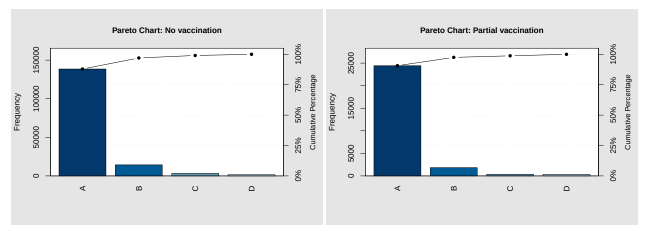
Now that we have the contingency table for event type by vaccine status, we can perform a χ^2 test testing the independence of both factors.

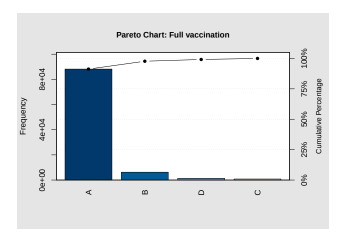
```
(chisq_vac = chisq.test(table_vaccine) )
```

```
##
## Pearson's Chi-squared test
##
## data: table_vaccine
## X-squared = 1332.2, df = 6, p-value < 2.2e-16</pre>
```

The null hypothesis that the type of event and vaccine status are independent can be rejected.

Pareto charts





Age and Infection Gravity

```
table_age = data.frame(matrix(0,ncol=6,nrow=4))
rownames(table_age) = c("casa", "hospital", "critical", "dead")
dd = df
dd$age = as.numeric(dd$age)
for (i in 1:dim(df)[1]){
  row_i = dd[i,]
        = row_i[[2]]
  age
  vacc = row_i[5]
  event = row_i[[4]]
  count = row_i[6]
  if (event=="Cas") row=1
  if (event=="Hospitalització") row = 2
  if (event=="Critics") row = 3
  if (event=="Defunció") row = 4
  table_age[row,age-3] = count + table_age[row,age-3]
  \# the factor starts to count at 4 for some reason
}
table_age
```

```
Х5
##
               X1
                     Х2
                           ХЗ
                                 Х4
                                              Х6
            74392 70147 43811 31794 17442 13497
## hospital 2449 3318
                         3766
                              4489
                                     4041
                                            4345
## critical
              408
                    683
                          873
                               1237
                                       967
                                             229
## dead
               13
                     36
                          125
                                383
                                       701
                                           1982
(chisq_age = chisq.test(table_age) )
```

```
##
## Pearson's Chi-squared test
```

```
##
## data: table_age
## X-squared = 32235, df = 15, p-value < 2.2e-16</pre>
```

Again, the null hypothesis that age and severity of COVID19 infection are independent can be rejected.

We can observe the difference between observed Counts and Expected Counts under null hypothesis.

```
table_age - chisq_age$expected
```

```
##
                    X1
                               X2
                                          ХЗ
                                                      Х4
                                                                 Х5
                                                                             Х6
## casa
             5387.2237
                        3891.2685
                                   427.3572 -2058.19170 -3234.7826 -4412.8752
## hospital -3709.3581 -2595.0185 -105.7901
                                              1467.84798
                                                          2195.6925
## critical
            -800.4211
                        -477.2795
                                   113.2597
                                               644.17577
                                                           604.9053
                                                                       -84.6402
## dead
             -877.4445
                        -818.9705 -434.8268
                                               -53.83205
                                                           434.1847
                                                                     1750.8892
```

#Statistical Inference

Parametric Analysis

Poisson Regression

To further investigate the relationship between vaccine status and event type, we fit a poisson model on the count variable from the contingency table. The model predicts the count variable through the event type and the vaccine status assuming a poisson distribution. We use poisson's canonical link function.

```
table_vaccine
```

##

```
##
              none partial full
## casa
                     24429 88167
            138487
## hospital 14323
                      1842 6243
## critical
              3228
                       342
                             827
## dead
              1513
                       336
                           1391
y = unname(unlist(table_vaccine))
print(y)
                    # y is a vector of the count variable
   [1] 138487
                14323
                         3228
                                1513 24429
                                              1842
                                                      342
                                                              336
                                                                  88167
                                                                           6243
## [11]
           827
                 1391
vaccine_status = as.factor(c(rep(1,4),rep(2,4),rep(3,4)))
event_type = as.factor(c(rep(c(1,2,3,4),3)))
poiss_glm = glm(y~vaccine_status+event_type, family="poisson")
summary(poiss_glm)
##
## Call:
```

glm(formula = y ~ vaccine_status + event_type, family = "poisson")

```
## Deviance Residuals:
##
       Min
                  10
                       Median
                                    30
                                             Max
   -19.271
##
             -6.907
                       -1.291
                                 6.746
                                          15.401
##
##
  Coefficients:
##
                     Estimate Std. Error z value Pr(>|z|)
                                           4555.1
## (Intercept)
                    11.854478
                                0.002602
                                                     <2e-16 ***
## vaccine_status2 -1.765803
                                0.006592
                                           -267.9
                                                     <2e-16 ***
## vaccine_status3 -0.488881
                                0.004086
                                           -119.6
                                                     <2e-16 ***
## event_type2
                    -2.416366
                                0.006972
                                           -346.6
                                                     <2e-16 ***
## event_type3
                    -4.044861
                                0.015212
                                           -265.9
                                                     <2e-16 ***
##
  event_type4
                    -4.350210
                                0.017681
                                           -246.0
                                                     <2e-16 ***
##
                   0 '*** 0.001 '** 0.01 '* 0.05 '. ' 0.1 ' ' 1
## Signif. codes:
##
## (Dispersion parameter for poisson family taken to be 1)
##
##
                                       degrees of freedom
       Null deviance: 647701
                               on 11
## Residual deviance:
                         1382
                               on
                                   6
                                      degrees of freedom
  AIC: 1516.2
##
## Number of Fisher Scoring iterations: 4
```

All parameters are significantly different from zero. Here we can better interpret the vaccination effect because it is estimated in dependency on the event type. Generally, being partially vaccinated is linked to fewer events compared to the baseline (not vaccinated), because the parameter vaccine_status2 has a negative sign. The estimated parameter for fully vaccinated people is also negative, but smaller. However, this does not mean that being partially vaccinated is advantageous. Instead, it is probably due to the fact, that people were only partially vaccinated for a very short time, where there was less chance to be infected.

anova(poiss_glm)

```
## Analysis of Deviance Table
##
## Model: poisson, link: log
##
## Response: y
##
  Terms added sequentially (first to last)
##
##
##
                   Df Deviance Resid. Df Resid. Dev
## NULL
                                       11
                                              647701
                                              545233
## vaccine_status
                    2
                        102469
                                        9
## event_type
                    3
                        543851
                                        6
                                                 1382
```

Multinomial Model

Another option for this data with a categorical would be a multinomial model. Because the model interpretation is difficult and it was not covered in class, it is not further discussed.

```
library(nnet)
multinom_model
                = multinom(event~vaccinated, data=df)
## # weights: 16 (9 variable)
## initial value 23648.795506
## iter 10 value 20325.467825
## final value 20285.735343
## converged
summary(multinom_model)
## Call:
## multinom(formula = event ~ vaccinated, data = df)
##
## Coefficients:
##
                   (Intercept) vaccinatedNo iniciada vaccinatedParcial
## Crítics
                    -1.5905962
                                           0.85955748
                                                             -0.4719066
## Defunció
                    -1.5458369
                                           0.08055976
                                                             -0.6105049
## Hospitalització -0.5151402
                                           0.32111808
                                                             -0.4002018
##
## Std. Errors:
##
                   (Intercept) vaccinatedNo iniciada vaccinatedParcial
                                           0.05565248
                                                             0.07953033
## Crítics
                    0.04603660
## Defunció
                    0.04519186
                                           0.06111938
                                                             0.08133331
## Hospitalització 0.03097434
                                           0.04077959
                                                             0.05117251
## Residual Deviance: 40571.47
## AIC: 40589.47
```

Logistic Regression

In order to fit a simpler, easier to interpret model, we create a binary variable "having a severe infection". Some people might argue, that being infected with the coronavirus is itself not a bad thing. What we are worried about, is having a severe disease, which is considerably worse than the common flu. Therefore we define a severe case as anything worse than being quarantined at home: Being hospitalized, on the ICU, or dying.

```
# define critical case as anything worse than being positive and quarantined ("Cas")
df$severe = as.numeric(! df$event=="Cas")
# now we would need to multiply the rows by count
df_enrolled = df[rep(row.names(df),df$count),]
```

In order to for a logistic Regression, we need to expand the dataset, so that each event constitutes its own row. We realize that by copying each row as often as it's count variable indicates via rep. The sanity check confirms that the resulting dataframe has as many rows as the sum of table_age in the old one.

```
print("sanity check: ")
## [1] "sanity check: "
```

```
dim(df_enrolled)[1] == sum(df$count) # sanity check
```

```
## [1] TRUE
```

We want to model the probability of a person of varying age, vaccination status and sex to have a non-severe or a severe case of Covid19. We are using a logistic model for this problem, because our dependent variable is binary, and we also have multiple explanatory variables. We also consider the explanatory variables as independent. For age and sex this is obviously true, but it might not be true for vaccination status and age, since older people have been vaccinated first.

```
logitReg = glm(severe ~ age + vaccinated + sex, data=df_enrolled, family=binomial(link='logit'))
summary(logitReg)
```

```
##
  glm(formula = severe ~ age + vaccinated + sex, family = binomial(link = "logit"),
##
       data = df_enrolled)
##
## Deviance Residuals:
##
       Min
                 10
                      Median
                                   30
                                            Max
## -1.3018 -0.4767 -0.3305 -0.2504
                                         2.9922
##
## Coefficients:
##
                         Estimate Std. Error z value Pr(>|z|)
                         -4.46513
                                     0.02525 -176.81
                                                        <2e-16 ***
## (Intercept)
## age40 a 49
                          0.55041
                                     0.02515
                                                21.89
                                                        <2e-16 ***
## age50 a 59
                          1.23045
                                     0.02465
                                                49.92
                                                        <2e-16 ***
## age60 a 69
                          1.94236
                                     0.02430
                                                79.93
                                                        <2e-16 ***
## age70 a 79
                                     0.02532
                                             100.17
                          2.53634
                                                        <2e-16 ***
## age80 o més
                          3.16772
                                     0.02616
                                              121.07
                                                        <2e-16 ***
## vaccinatedNo iniciada 1.11729
                                     0.01571
                                                71.12
                                                        <2e-16 ***
## vaccinatedParcial
                          0.49110
                                     0.02552
                                                19.25
                                                        <2e-16 ***
## sexHome
                          0.46780
                                     0.01315
                                                35.56
                                                        <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: 191126 on 281127 degrees of freedom
## Residual deviance: 164713 on 281119 degrees of freedom
## AIC: 164731
##
## Number of Fisher Scoring iterations: 6
```

Again, all parameters are statistically significant having a p-value numerically equal to zero. We can interpret the parameters more easily when we take their exponent, which constitutes the change in the Odds Ratio:

Another meaningful link function for the binary regression is the probit link. This assumes a latent normal process.

```
probitReg = glm(severe ~ age + vaccinated + sex, data=df_enrolled, family=binomial(link='probit'))
summary(probitReg)
```

```
##
## Call:
## glm(formula = severe ~ age + vaccinated + sex, family = binomial(link = "probit"),
       data = df_enrolled)
## Deviance Residuals:
       Min
                 10
                      Median
                                   30
                                           Max
## -1.2505 -0.4832 -0.3408 -0.2330
                                         3.1346
##
## Coefficients:
                          Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                         -2.439663
                                     0.012146 -200.87
                                                        <2e-16 ***
## age40 a 49
                          0.260994
                                     0.011577
                                                22.54
                                                        <2e-16 ***
## age50 a 59
                                                50.75
                          0.599291
                                     0.011808
                                                        <2e-16 ***
## age60 a 69
                                     0.011992
                                                82.43
                          0.988519
                                                         <2e-16 ***
## age70 a 79
                          1.329429
                                     0.012957 102.60
                                                         <2e-16 ***
## age80 o més
                          1.692007
                                     0.013548 124.89
                                                         <2e-16 ***
## vaccinatedNo iniciada 0.614790
                                     0.008353
                                               73.60
                                                         <2e-16 ***
## vaccinatedParcial
                                                20.13
                          0.269894
                                     0.013410
                                                         <2e-16 ***
## sexHome
                          0.239457
                                     0.006912
                                                34.64
                                                         <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 191126 on 281127 degrees of freedom
## Residual deviance: 164497 on 281119 degrees of freedom
## AIC: 164515
##
## Number of Fisher Scoring iterations: 6
The last link function we try is the complementary log log link.
cllReg = glm(severe ~ age + vaccinated + sex, data=df_enrolled, family=binomial(link='cloglog'))
summary(cllReg)
##
## Call:
  glm(formula = severe ~ age + vaccinated + sex, family = binomial(link = "cloglog"),
       data = df_enrolled)
##
##
## Deviance Residuals:
       Min
                 1Q
                      Median
                                   3Q
                                           Max
## -1.3450 -0.4746 -0.3256 -0.2616
                                        2.9406
##
## Coefficients:
                         Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                         -4.31700
                                     0.02357 -183.16
                                                        <2e-16 ***
## age40 a 49
                          0.52397
                                     0.02446
                                               21.42
                                                        <2e-16 ***
## age50 a 59
                          1.17156
                                     0.02369
                                               49.45
                                                        <2e-16 ***
## age60 a 69
                                               79.07
                                                        <2e-16 ***
                          1.81506
                                     0.02296
## age70 a 79
                          2.32566
                                     0.02326
                                               99.97
                                                        <2e-16 ***
## age80 o més
                                     0.02340 121.27
                          2.83725
                                                        <2e-16 ***
```

```
## vaccinatedNo iniciada 0.96158
                                    0.01378
                                              69.80
                                                      <2e-16 ***
## vaccinatedParcial
                         0.42666
                                    0.02299
                                              18.56
                                                      <2e-16 ***
                         0.41778
## sexHome
                                    0.01180
                                              35.41
                                                      <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 191126 on 281127 degrees of freedom
## Residual deviance: 164994 on 281119 degrees of freedom
## AIC: 165012
##
## Number of Fisher Scoring iterations: 6
# function to calculate fisher statistic
fisherStat = function(RegM){
 return(sum(residuals(RegM, type="deviance")^2)/RegM$df.residual)
fisherStat(logitReg)
## [1] 0.58592
fisherStat(probitReg)
## [1] 0.5851513
fisherStat(cllReg)
```

[1] 0.5869195

All three Fisher Statistics are very similar between 0.585-0.586 indicating Underdispersion.

Comparing the Models In order to compare the three binary regressions and to check which link function results in the best fit, we compare the log-Likelihood and the Akaike Information Coefficient. The Poisson Model cannot be compared with binary regressions because the predicted y is of different nature.

```
round(c(logitReg$aic,probitReg$aic,cllReg$aic),5)
```

```
## [1] 164731.2 164515.2 165012.2
```

With AIC smaller values are considered to show a better model fit.

```
logLik(logitReg)
```

```
## 'log Lik.' -82356.62 (df=9)
```

logLik(probitReg)

```
## 'log Lik.' -82248.58 (df=9)
```

logLik(cllReg)

```
## 'log Lik.' -82497.11 (df=9)
```

With the log-Likelihood bigger values indicate better model fit. So for both metrics the probit-Regression model has the best value.

Model Interpretation However, the values of the logistic Regression are easier to interpret because the estimated parameters can be seen as the log(Odds). Since the models are of similar quality, let's interpret these parameters. Taking the exponential of the parameter, gives the Odds Ratio relative to the baseline group.

round(exp(logitReg\$coef),2)

##	(Intercept)	age40 a 49	age50 a 59
##	0.01	1.73	3.42
##	age60 a 69	age70 a 79	age80 o més
##	6.98	12.63	23.75
##	vaccinatedNo iniciada	${\tt vaccinatedParcial}$	sexHome
##	3.06	1.63	1.60

All age categories have a Odds Ratios (OR) bigger than 1. This means people of all ages have a higher risk to get a severe COVID19 infection than the baseline defined to be individuals between 30 and 40. Also the OR increases with age, which means that the Odds increase the older an individual is. The OR for persons being 80 or more is estimated as 23.75, which is the biggest value in this model. This indicates, that age played a bigger role than vaccine status (in 2021 in Catalonia). The Odds for male individuals are estimated to be 60% to suffer from a severe COVID19 disease. For vaccine status, the fully vaccinated people are taken as baseline group. Non-vaccinated people of the same age group and sex are estimated to have approximately 3 times the Odds of having a severe Coronavirus infection. Also being partially vaccinated increases the odds by around 63% given that other covariates remain the same.

Conclusion

This work studying vaccine effectiveness shows the issues of non-randomized data. Most of the issues, we had to wrap our heads around would naturally dissolve with a randomized control study, which is why it is the gold standard for pharmaceutical studies.

One issue present is that older people were vaccinated first, but still were at high risk for severe Covid infections. Therefore the age necessarily has to be included in the models. Another issue is the baseline fallacy, that at some point more people were vaccinated than not, which automatically shifts the number of events towards vaccinated individuals.

Another observation was that all p-values displayed were strikingly small. This is because both the effect sizes and the sample size of this data set were very large, which naturally influence the statistical inference.

One conclusion, which we were not aware of before, is that male persons suffered of more severe infections. This might be due to statistical link between males and less healthy lifestyles like smoking resulting in cardiovascular diseases, which are a negative predictor for survival of COVID19.

Regarding the main analysis, we can confirm what others have found before:

- Age plays a major role for severe COVID19 diseases. Older people are at much higher risk. In this data the Odds for a severe infection were estimated to be 23 times higher for individuals older than 80 than those between 30 and 40.
- Being (fully) vaccinated clearly lowers the risk for a severe Coronavirus infection. Not being vaccinated is estimated to result in a three times increased risk for a severe COVID19 disease progression.