

Next-Generation Analysis (NGA): case study for Portugal

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This report presents the results of the paper by Caetano et al. (2024), titled “Identifying the Main Drivers of Transmission in the Early Phase of the COVID-19 Pandemic in Portugal” in which we analysed how age-specific contact patterns, susceptibility, and infectivity, affect the basic reproduction number R_0 while considering transmission heterogeneity. To understand each age group’s impact on overall transmission, we calculated sensitivity measures associated with R_0 . Additionally, we introduced a novel indicator designed to estimate the expected change in new infections following public health interventions. This methodology offers valuable tools for strategizing the distribution of mitigation measures during infectious disease outbreaks. Its adaptability enables easy integration of population-specific data, supporting comparative evaluations of epidemic control strategies across various countries. All code is open-source, but please ensure you cite Caetano et al. (2024) when building upon this work.

In this document, we demonstrated the methodology through an analysis of COVID-19 transmission dynamics in Portugal during March 2020, as published in Caetano et al. (2024). The results highlighted that individuals aged 30–59 played the most significant role in driving transmission. Furthermore, by evaluating the effects of changes in susceptibility and infectivity, we found that reducing susceptibility by 10% in the 30–39 age group led to a 17% decrease in incidence in this group after three generations, along with an indirect reduction of 4–6% in other age groups.

Load code

If the `curl` package is not installed yet, you can do so by

```
install.packages(curl) # install the curl library (to easily download files from GitHub)
```

Please load the functions we prepared on the SOCRATES GitHub repository.

```
# load package 'curl'
library('curl')

# download R files with NGA functions
curl_download("https://github.com/lwillem/socrates_rshiny/blob/master/R/next_gen_lib.R",
              destfile = "next_gen_lib.R")
curl_download("https://github.com/lwillem/socrates_rshiny/blob/master/R/plot_next_gen.R",
              destfile = "plot_next_gen.R")

# load the functions
source('next_gen_lib.R')
source('plot_next_gen.R')
```

Load data

Load social contact data for Portugal based on Prem et al (2021, Plos Comput Biol).

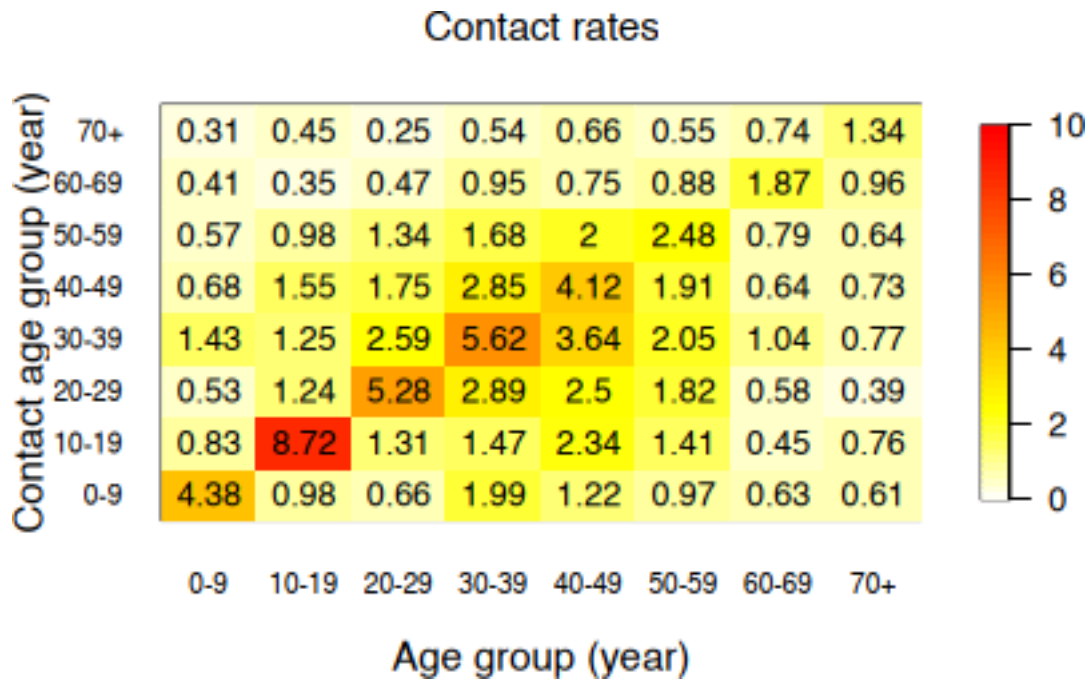
```
# download rds file with social contact data
curl_download("https://github.com/lwillem/socrates_rshiny/blob/master/data/CM_Portugal_Prem2021.rds",
              destfile = "CM_Portugal_Prem2021.rds")

mij <- readRDS(file = '../data/CM_Portugal_Prem2021.rds')

# set contact matrix into SOCRATES format
mij <- t(mij)
```

Explore social contact data

```
# plot matrix
matrix_plot(mij)
```



Run next-generation analysis

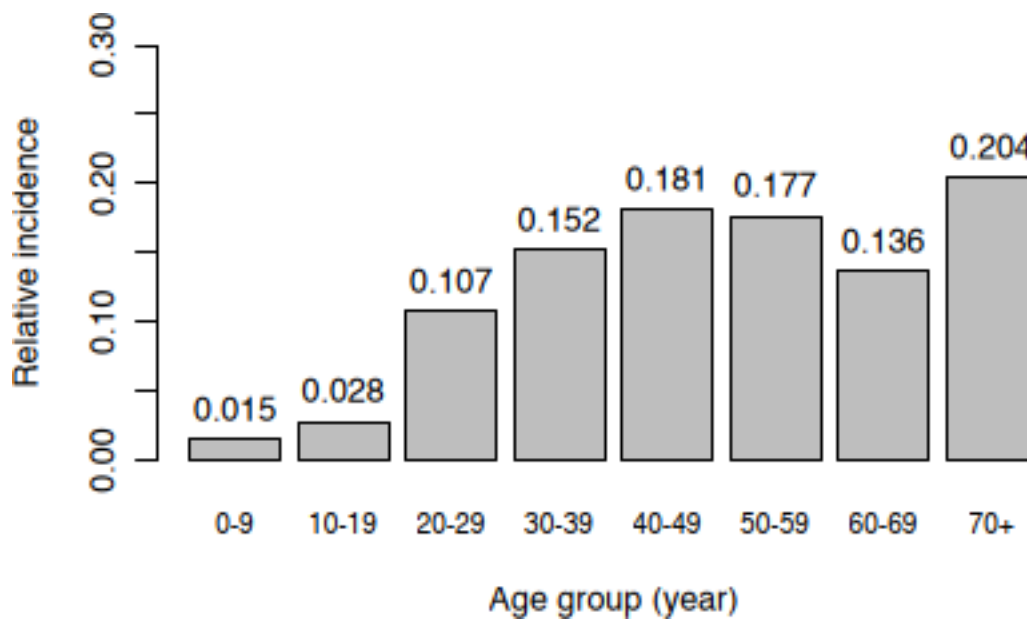
```
# parameters
qi      = c(0.545,0.555,0.590,0.700,0.760,0.900,0.990,0.990)
qs      = c(0.069,0.081,0.223,0.221,0.264,0.353,0.439,0.703)
q       = 1
delta_p = -0.10
nr_gen  = 3

# calculate relative incidence
next_generation_matrix = NGM_SIR(q=q,a=qs,M=t(mij),h=qi)
relative_incidence     = standardize_RI(eigen(next_generation_matrix)$vectors[,1])
names(relative_incidence) = colnames(mij)

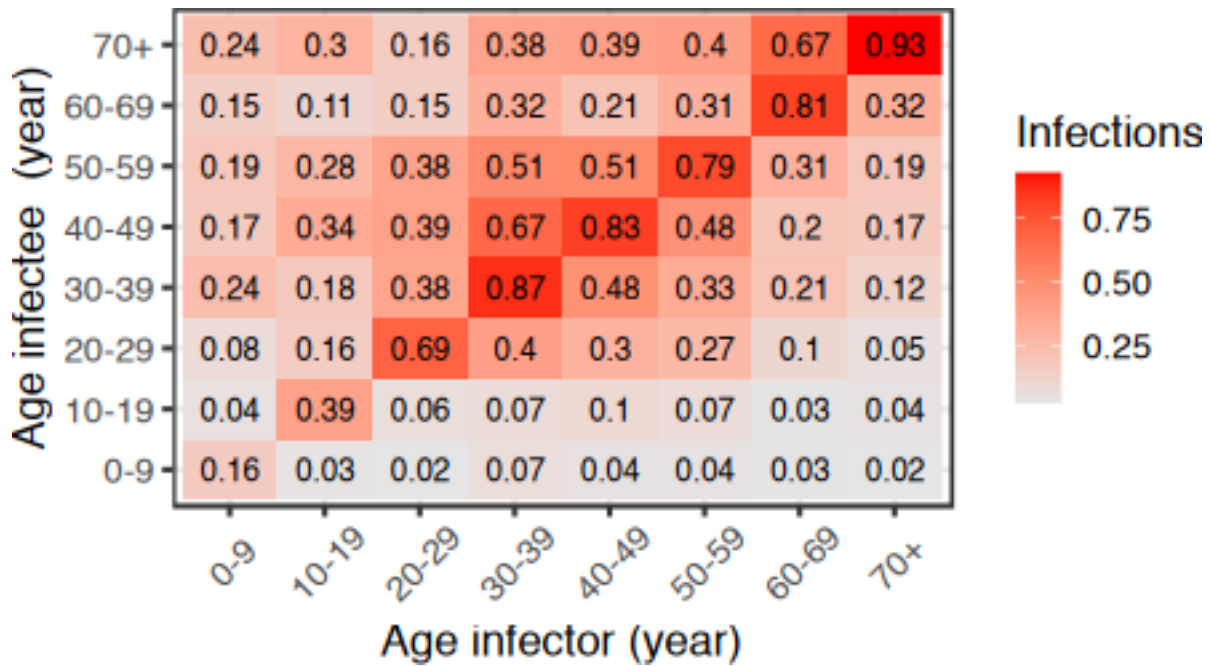
# run full next-generation analysis according to Ceatano et al (2024).
NGA = run_NGA(M=mij,a=qs,h=qi,q=q,p=delta_p,nr_gen=nr_gen)
```

Plot relative incidence

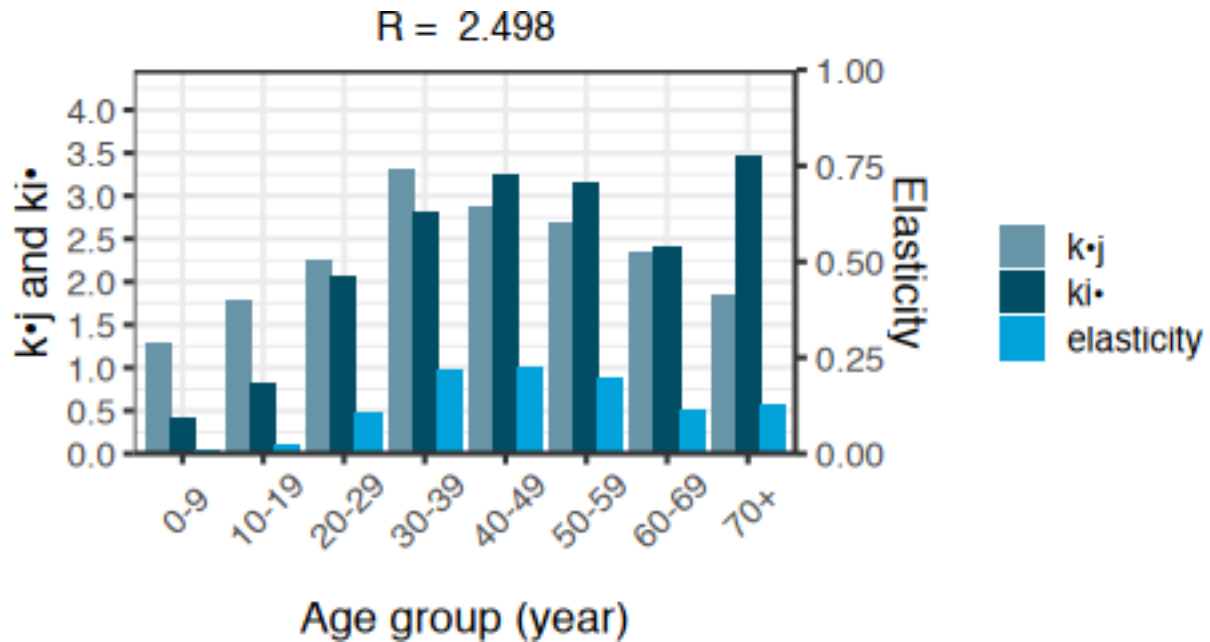
```
# relative incidence
barplot(relative_incidence,
        xlab="Age group (year)",
        ylab="Relative incidence",
        ylim=c(0,0.3),
        cex.names = 0.8) -> bplt
text(x = bplt,
     y = relative_incidence,
     labels = round(relative_incidence,digits=3),
     pos=3)
```



```
# plot next-generation matrix
plot_next_gen_matrix(next_gen_matrix = NGA$next_gen_matrix)
```

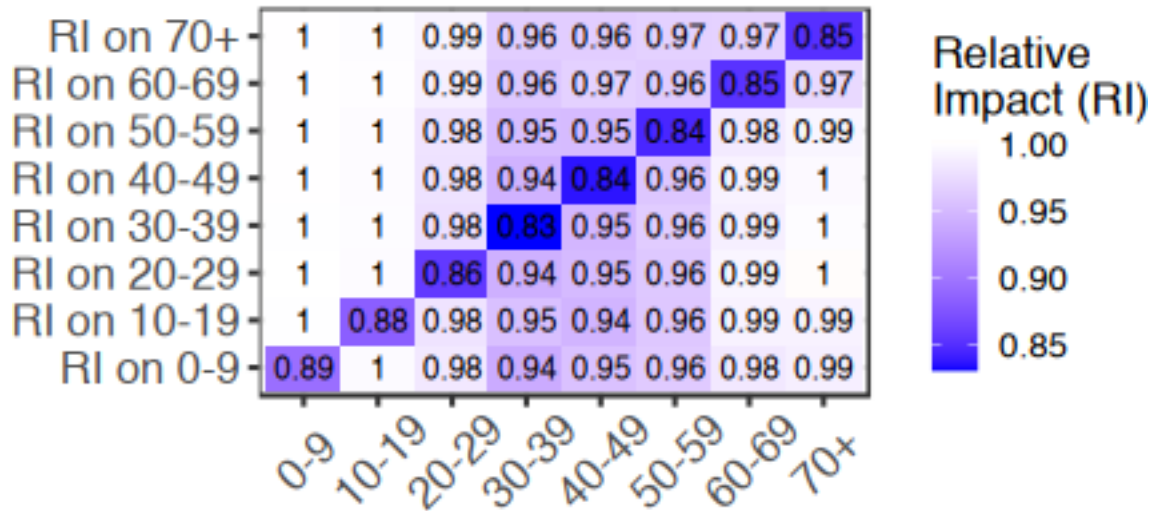


```
# plot elasticity
plot_NGA_elas(R_t = NGA$R_t, elasticity_tbl = NGA$elasticity_tbl)
```



```
# plot relative incidence with regards to susceptibility
plot_NGA_RI(NGA = NGA, delta_p = delta_p, rn_gen = nr_gen,
            bool_susceptibility = TRUE, round_digits = 2)
```

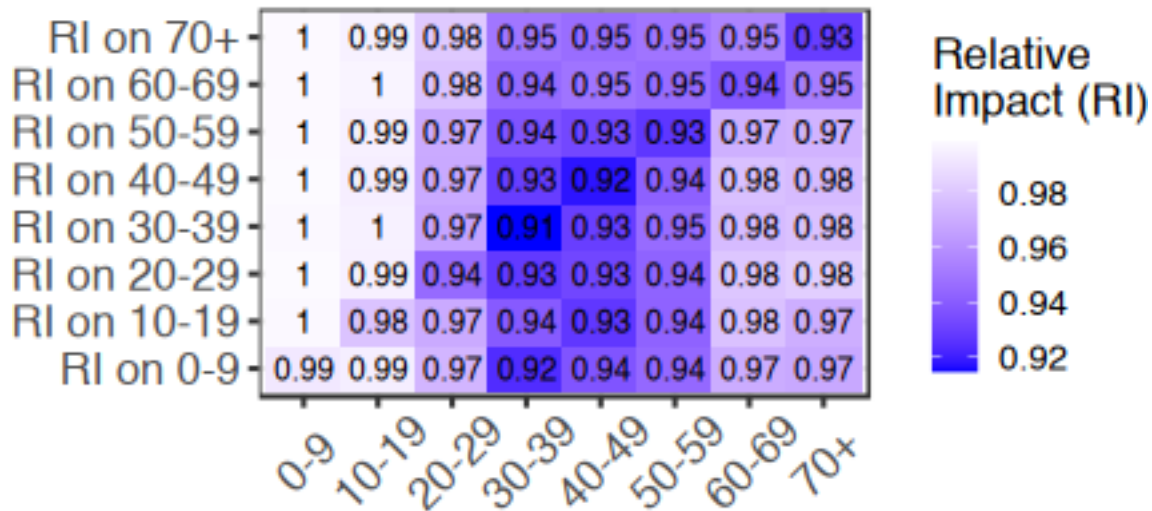
m = 3



Age group (year) with 10% reduction in q-susceptibility

```
# plot relative incidence with regards to infectiousness
plot_NGA_RI(NGA = NGA, delta_p = delta_p, rn_gen = nr_gen,
            bool_susceptibility = FALSE, round_digits = 2)
```

m = 3



Age group (year) with 10% reduction in q-infectivity