COVID-19 data analysis and forecasting for India and Germany using SIRD model

Dhwanit, Shristi, Akksay and Bao

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1. Introduction

Coronaviruses are a family of human pathogens and have been responsible for a variety of respiratory diseases in humans ranging from common cold to dangerous ones like SARS (Severe Acute Respiratory) and MERS (Middle East Respiratory Syndrome). At the end of 2019, a novel strain of coronavirus was identified as a casue of pneumonia-type illness in Wuhan, Hubei in China. The virus was highly contagious and rapidly spread throughout the city. Though the Chinese government ordered lockdown of the city, the virus had spread to different parts of the world resulting in a global pandemic of a massive scale in February 2020. The virus has been named as 2019-nCov and the disease has been termed as COVID-19 (Coronavirus disease 2019).

The understanding of the virus is evolving and several models have been proposed to find the basic reproducibility number (denoted by R_0), case fatality rate and case recovery rate. In this study, we use epidemic transmission dynamics based compartmental model called $SIRD\ model$ to study the progression of the disease for different countries (in particular, Germany and India). We use the discrete approximations as in [1] to estimate the R_0 , case mortality and case recovery rates and analyse the changes as time progresses. Furthermore, we fit the model to the epidemic curves of these countries to make forecasts about the peak of the epidemic and the flattening of the curve.

2. Data visualization and exploratory analysis

The COVID-19 data used in this analysis is available on the public git repository COVID. The data has the time series of confirmed, recovered and death cases of COVID-19 for various countries. The first five rows of confirmed cases database look like this:

```
confirmed <- read.csv("data/time_series_covid19_confirmed_global.csv")

deaths <- read.csv("data/time_series_covid19_deaths_global.csv")

recovered <- read.csv("data/time_series_covid19_recovered_global.csv")

head(confirmed, 5)</pre>
```

```
##
     Province.State Country.Region
                                          Lat
                                                  Long X1.22.20 X1.23.20 X1.24.20
## 1
                        Afghanistan
                                      33.0000 65.0000
                                                               0
                                                                         0
                                                                                  0
## 2
                                      41.1533 20.1683
                                                               0
                                                                         0
                            Albania
                                                                                  0
                            Algeria
## 3
                                      28.0339
                                               1.6596
                                                               0
                                                                         0
                                                                                  0
                                                                         0
## 4
                            Andorra
                                     42.5063
                                               1.5218
                                                                                  0
## 5
                              Angola -11.2027 17.8739
                                                                                  0
##
     X1.25.20 X1.26.20 X1.27.20 X1.28.20 X1.29.20 X1.30.20 X1.31.20 X2.1.20
                                0
                                                   0
## 1
            0
                      0
                                         0
                                                             0
                                                                      0
                                                                               0
            0
                      0
                                0
                                         0
                                                   0
                                                             0
## 2
                                                                      0
                                                                               0
```

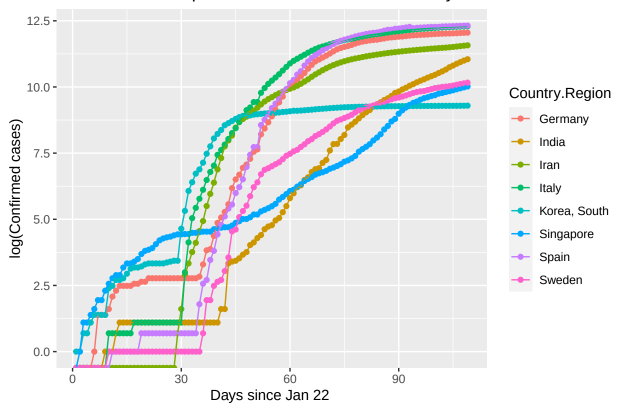
##	3	0	0	0	0	0	0	0	0	
##	4	0	0	0	0	0	0	0	0	
##	5	0	0	0	0	0	0	0	0	
##			X2.3.20 X2			6.20 X2.7.				
##	1	0	0	0	0	0	0	0 0		
##	2	0	0	0	0	0	0	0 0	_	
##	3	0	0	0	0	0	0	0 0	0	
##	4	0	0	0	0	0	0	0 0	0	
##	5	0	0	0	VO 14 00	0	0	0 0	VO 10 00	
##	1	X2.11.20	X2.12.20		X2.14.20					
##	1 2	0	0	0	0	0	0	0	0	
##	2	0	0	0	0	0	0	0	0	
##	<i>ا</i>	0	0	0	0	0	0	0	0	
##	5	0	0	0	0	0	0	0	0	
##	J	X2.19.20	·	·	X2.22.20	X2.23.20	X2.24.20	·	X2.26.20	
##	1	0	0	0	0	0	1	1	1	
##	2	0	0	0	0	0	0	0	0	
##	3	0	0	0	0	0	0	1	1	
##	4	0	0	0	0	0	0	0	0	
##	5	0	0	0	0	0	0	0	0	
##		X2.27.20	X2.28.20	X2.29.20	X3.1.20	X3.2.20 X3	3.3.20 X3	.4.20 X3.5	.20 X3.6.2	20
##	1	1	1	1	1	1	1	1	1	1
##	2	0	0	0	0	0	0	0	0	0
##	3	1	1	1	1	3	5	12	12 1	۱7
##	4	0	0	0	0	1	1	1	1	1
##	5	0	0	0	0	0	0	0	0	0
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##		X3.7.20 X	X3.8.20 X3	3.9.20 X3		.11.20 X3.				15.20
## ##	1	X3.7.20 X	4	4	.10.20 X3	7	7	7	11	15.20 16
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	1 2 3	1 0 17	4 0 19	4 2 20	5 10 20	7 12 20	7 23 24	7 33 26	11 38 37	15.20 16 42 48
## ## ##	2 3 4	1 0 17 1	4 0 19 1	4 2 20 1	5 10 20 1	7 12 20 1	7 23 24 1	7 33 26 1	11 38 37 1	15.20 16 42
## ## ## ##	1 2 3 4 5	1 0 17 1 0	4 0 19 1 0	4 2 20 1 0	5 10 20 1 0	7 12 20 1 0	7 23 24 1 0	7 33 26 1 0	11 38 37 1 0	15.20 16 42 48
## ## ##	2 3 4	1 0 17 1 0 X3.16.20	4 0 19 1 0 X3.17.20	4 2 20 1 0 X3.18.20	5 10 20 1 0 X3.19.20	7 12 20 1 0 X3.20.20	7 23 24 1 0 X3.21.20	7 33 26 1 0 X3.22.20	11 38 37 1 0 X3.23.20	15.20 16 42 48
## ## ## ##	2 3 4	1 0 17 1 0 X3.16.20 21	4 0 19 1 0 X3.17.20 22	4 2 20 1 0 X3.18.20 22	5 10 20 1 0 X3.19.20	7 12 20 1 0 X3.20.20 24	7 23 24 1 0 X3.21.20	7 33 26 1 0 X3.22.20 40	11 38 37 1 0 X3.23.20 40	15.20 16 42 48
## ## ## ##	2 3 4 5	1 0 17 1 0 X3.16.20 21 51	4 0 19 1 0 X3.17.20 22 55	4 2 20 1 0 X3.18.20 22 59	5 10 20 1 0 X3.19.20 22 64	7 12 20 1 0 X3.20.20 24 70	7 23 24 1 0 X3.21.20 24 76	7 33 26 1 0 X3.22.20 40 89	11 38 37 1 0 X3.23.20 40 104	15.20 16 42 48
## ## ## ## ## ##	2 3 4 5 1 2 3	1 0 17 1 0 X3.16.20 21 51 54	4 0 19 1 0 X3.17.20 22 55 60	4 2 20 1 0 X3.18.20 22 59 74	5 10 20 1 0 X3.19.20 22 64 87	7 12 20 1 0 X3.20.20 24 70 90	7 23 24 1 0 X3.21.20 24 76 139	7 33 26 1 0 X3.22.20 40 89 201	11 38 37 1 0 X3.23.20 40 104 230	15.20 16 42 48
## ## ## ##	2 3 4 5 1 2 3 4	1 0 17 1 0 X3.16.20 21 51	4 0 19 1 0 X3.17.20 22 55 60 39	4 2 20 1 0 X3.18.20 22 59 74 39	5 10 20 1 0 X3.19.20 22 64 87 53	7 12 20 1 0 X3.20.20 24 70 90 75	7 23 24 1 0 X3.21.20 24 76 139 88	7 33 26 1 0 X3.22.20 40 89 201 113	11 38 37 1 0 X3.23.20 40 104 230 133	15.20 16 42 48
## ## ## ## ## ## ##	2 3 4 5 1 2 3 4 5	1 0 17 1 0 X3.16.20 21 51 54 2	4 0 19 1 0 X3.17.20 22 55 60 39 0	4 2 20 1 0 X3.18.20 22 59 74 39 0	5 10 20 1 0 X3.19.20 22 64 87 53	7 12 20 1 0 X3.20.20 24 70 90 75	7 23 24 1 0 X3.21.20 24 76 139 88 2	7 33 26 1 0 X3.22.20 40 89 201 113 2	11 38 37 1 0 X3.23.20 40 104 230 133 3	15.20 16 42 48
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## ## ## ## ## ## ## ## ## ## ## ## ##	2 3 4 5 1 2 3 4 5 1 2	1 0 17 1 0 X3.16.20 21 51 54 2 0 X3.24.20	4 0 19 1 0 X3.17.20 22 55 60 39 0 X3.25.20 84 146	4 2 20 1 0 X3.18.20 22 59 74 39 0 X3.26.20 94	5 10 20 1 0 X3.19.20 22 64 87 53 0 X3.27.20 110 186	7 12 20 1 0 X3.20.20 24 70 90 75 1 X3.28.20 110 197	7 23 24 1 0 X3.21.20 24 76 139 88 2 X3.29.20 120	7 33 26 1 0 X3.22.20 40 89 201 113 2 X3.30.20 170	11 38 37 1 0 X3.23.20 40 104 230 133 3 X3.31.20	15.20 16 42 48
## ## ## ## ## ## ## ## ## ## ## ## ##	2 3 4 5 1 2 3 4 5 1 2 3	1 0 17 1 0 X3.16.20 21 51 54 2 0 X3.24.20 74 123	4 0 19 1 0 X3.17.20 22 55 60 39 0 X3.25.20 84 146 302	4 2 20 1 0 X3.18.20 22 59 74 39 0 X3.26.20 94 174	5 10 20 1 0 X3.19.20 22 64 87 53 0 X3.27.20 110 186	7 12 20 1 0 X3.20.20 24 70 90 75 1 X3.28.20 110 197 454	7 23 24 1 0 X3.21.20 24 76 139 88 2 X3.29.20 120 212	7 33 26 1 0 X3.22.20 40 89 201 113 2 X3.30.20 170 223	11 38 37 1 0 X3.23.20 40 104 230 133 3 X3.31.20 174 243	15.20 16 42 48
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######################	2 3 4 5 1 2 3 4 5 1 2 3 4 5 1 2 3	1 0 17 1 0 X3.16.20 21 51 54 2 0 X3.24.20 74 123 264 164 3 X4.1.20 237 259 847	4 0 19 1 0 X3.17.20 22 55 60 39 0 X3.25.20 84 146 302 188 3 X4.2.20 X4 273 277 986	4 2 20 1 0 X3.18.20 22 59 74 39 0 X3.26.20 94 174 367 224 4 4.3.20 X4 281 304 1171	5 10 20 1 0 X3.19.20 22 64 87 53 0 X3.27.20 110 186 409 267 4.20 X4. 299 333 1251	7 12 20 1 0 X3.20.20 24 70 90 75 1 X3.28.20 110 197 454 308 5 5.20 X4.6. 349 361 349 361 31320 14	7 23 24 1 0 X3.21.20 24 76 139 88 2 X3.29.20 120 212 511 334 7 20 X4.7.2 667 42 677 38	7 33 26 1 0 X3.22.20 40 89 201 113 2 X3.30.20 170 223 584 370 7 20 X4.8.20 23 444 33 400 68 1572	11 38 37 1 0 X3.23.20 40 104 230 133 3 X3.31.20 174 243 716 376 7 X4.9.20 484 409 1666	15.20 16 42 48
########################	2345 12345 12345 1234	1 0 17 1 0 X3.16.20 21 51 54 2 0 X3.24.20 74 123 264 164 3 X4.1.20 237 259 847 390	4 0 19 1 0 X3.17.20 22 55 60 39 0 X3.25.20 84 146 302 188 3 X4.2.20 X4 273 277 986 428	4 2 20 1 0 X3.18.20 22 59 74 39 0 X3.26.20 94 174 367 224 4 4.3.20 X4 281 304 1171 439	5 10 20 1 0 X3.19.20 22 64 87 53 0 X3.27.20 110 186 409 267 4.4.20 X4. 299 333 1251 466	7 12 20 1 0 X3.20.20 24 70 90 75 1 X3.28.20 110 197 454 308 5 5.20 X4.6. 349 361 349 361 31320 14 501 5	7 23 24 1 0 X3.21.20 24 76 139 88 2 X3.29.20 120 212 511 334 7 20 X4.7.2 667 42 677 38 623 146 625 54	7 33 26 1 0 X3.22.20 40 89 201 113 2 X3.30.20 170 223 584 370 7 20 X4.8.20 23 444 33 400 68 1572 45 564	11 38 37 1 0 X3.23.20 40 104 230 133 3 X3.31.20 174 243 716 376 7 X4.9.20 484 409 1666 583	15.20 16 42 48
########################	2345 12345 12345 12345	1 0 17 1 0 X3.16.20 21 51 54 2 0 X3.24.20 74 123 264 164 3 X4.1.20 237 259 847 390 8	4 0 19 1 0 X3.17.20 22 55 60 39 0 X3.25.20 84 146 302 188 3 X4.2.20 X4 273 277 986 428 8	4 2 20 1 0 X3.18.20 22 59 74 39 0 X3.26.20 94 174 367 224 4 4.3.20 X4 281 304 1171 439 8	5 10 20 1 0 X3.19.20 22 64 87 53 0 X3.27.20 110 186 409 267 4 .4.20 X4. 299 333 1251 466 10	7 12 20 1 0 X3.20.20 24 70 90 75 1 X3.28.20 110 197 454 308 5 5.20 X4.6. 349 361 3361 31320 14 501 5	7 23 24 1 0 X3.21.20 24 76 139 88 2 X3.29.20 120 212 511 334 7 20 X4.7.2 667 42 677 38 625 54 16	7 33 26 1 0 X3.22.20 40 89 201 113 2 X3.30.20 170 223 584 370 7 20 X4.8.20 23 444 33 400 68 1572 45 564 17 19	11 38 37 1 0 X3.23.20 40 104 230 133 3 X3.31.20 174 243 716 376 7 X4.9.20 484 409 1666 583 19	15.20 16 42 48
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########################	-2345 12345 12345 1	1 0 17 1 0 X3.16.20 21 51 54 2 0 X3.24.20 74 123 264 164 3 X4.1.20 237 259 847 390 8	4 0 19 1 0 X3.17.20 22 55 60 39 0 X3.25.20 84 146 302 188 3 X4.2.20 X4 273 277 986 428 8	4 2 20 1 0 X3.18.20 22 59 74 39 0 X3.26.20 94 174 367 224 4 4.3.20 X4 281 304 1171 439 8	5 10 20 1 0 X3.19.20 64 87 53 0 X3.27.20 110 186 409 267 4.4.20 X4. 299 333 1251 466 10 X4.13.20	7 12 20 1 0 X3.20.20 24 70 90 75 1 X3.28.20 110 197 454 308 5 5.20 X4.6. 349 361 349 361 31320 14 501 5 14 X4.14.20 714	7 23 24 1 0 X3.21.20 24 76 139 88 2 X3.29.20 120 212 511 334 7 20 X4.7.2 667 42 677 38 625 54 16	7 33 26 1 0 X3.22.20 40 89 201 113 2 X3.30.20 170 223 584 370 7 20 X4.8.20 23 444 33 400 68 1572 45 564 17 19	11 38 37 1 0 X3.23.20 40 104 230 133 3 X3.31.20 174 243 716 376 7 X4.9.20 484 409 1666 583 19	15.20 16 42 48

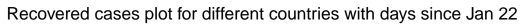
##	3	1761	1825	1914	1983	2070	2160	2268	241	8
##	4	601	601	638	646	659	673	673	69	6
##	5	19	19	19	19	19	19	19	1	9
##		X4.18.20	X4.19.20	X4.20.20	X4.21.20	X4.22.20	X4.23.20	X4.24.20	X4.25.2	0
##	1	933	996	1026	1092	1176	1279	1351	146	3
##	2	548	562	584	609	634	663	678	71	2
##	3	2534	2629	2718	2811	2910	3007	3127	325	6
##	4	704	713	717	717	723	723	731	73	8
##	5	24	24	24	24	25	25	25	2	5
##		X4.26.20	X4.27.20	X4.28.20	X4.29.20	X4.30.20	X5.1.20	X5.2.20 X	5.3.20 X	5.4.20
##	1	1531	1703	1828	1939	2171	2335	2469	2704	2894
##	2	726	736	750	766	773	782	789	795	803
##	3	3382	3517	3649	3848	4006	4154	4295	4474	4648
##	4	738	743	743	743	745	745	747	748	750
##	5	26	27	27	27	27	30	35	35	35
##		X5.5.20	X5.6.20 X	5.7.20 X5	.8.20 X5.	9.20				
##	1	3224	3392	3563	3778	4033				
##	2	820	832	842	850	856				
##	3	4838	4997	5182	5369	5558				
##	4	751	751	752	752	754				
##	5	36	36	36	43	43				

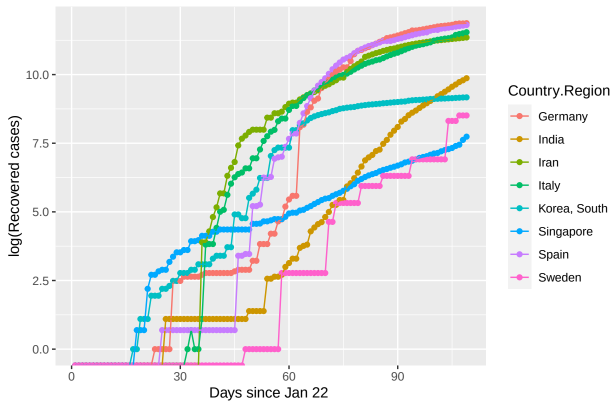
We can see the data starts from January 22, 2020 and gives us the number of confirmed cases on each date since then till now.

Below we plot the progression of the disease (confirmed, recovered, deaths) in a few selected countries.

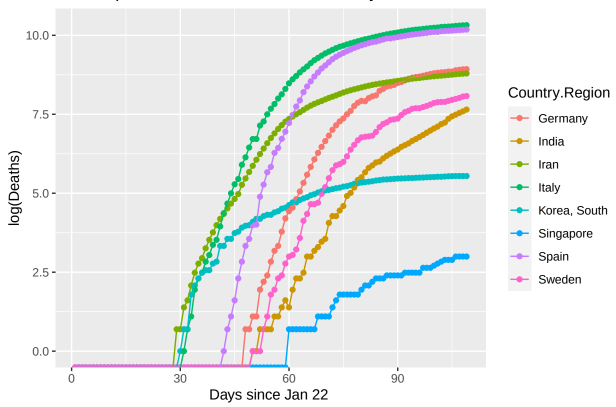
Confirmed cases plot for different countries with days since Jan 22







Deaths plot for different countries with days since Jan 22



We can see from the plots that South Korea has successfully flattened the curve and countries like Germany and Italy have started flattening. On the other hand, India and Singapore are still far from flattening the curve.

3. Formulation of SIRD model

In this section, we discuss the basics of the SIRD model to simulate epidemic progression with time.

3.1 Discrete model

Within this model of the evolution of an epidemic outbreak, people can be divided into different classes. In the susceptible (S), infected (I), recovered (R), dead (D) scheme (SIRD), any individual in the fraction of the overall population that will eventually get sick belongs to one of the aforementioned classes. Let N be the size of the initial population of susceptible people. The discrete SIRD model can be written as follows:

$$S(t) - S(t-1) = -\frac{\alpha}{N}S(t-1)I(t-1),$$

$$I(t) - I(t-1) = \frac{\alpha}{N}S(t-1)I(t-1) - \beta I(t-1) - \gamma I(t-1),$$

$$R(t) - R(t-1) = \beta I(t-1),$$

$$D(t) - D(t-1) = \gamma I(t-1),$$

The basic reproduction number R_0 is then defined as

$$R_0 := \frac{\alpha}{\beta + \gamma}.$$

Since the number of susceptible people is hard to determine and depends on the population, lockdown measures, social distancing etc, we take a different approach to estimate R_0 as mentioned in Ref 1. Let us denote $\Delta X(t) := X(t) - X(t-1)$ for X = I, R, D. Now we define,

$$C\Delta X(T) := \sum_{t=1}^{T} \Delta X(t), \ and$$

$$\mathbf{C}\Delta \mathbf{X}(T) := \left[C\Delta X(1), C\Delta X(2), ..., C\Delta X(T)\right]^{T}.$$

Here C stands for cumulative. Using the approximation $S(t-1) \approx N$ (true if susceptible population is much less than the population of the country), we can get

$$R_0 = \frac{\alpha}{\beta + \gamma} = \frac{I(t) - I(t-1) + R(t) - R(t-1) + D(t) - D(t-1)}{R(t) - R(t-1) + D(t) - D(t-1)}.$$

Summing this equation over time we get,

$$\frac{C\Delta I(t) + C\Delta R(t) + C\Delta D(t)}{C\Delta R(t) + C\Delta D(t)} = R_0.$$

Based on this, we can get a coarse estimate for R_0 by finding a least squares solution to the following regression problem:

$$C\Delta I(t) + C\Delta R(t) + C\Delta D(t)t = [C\Delta R(t) + C\Delta D(t)]R_0,$$

with solution given by

$$\hat{R}_0 = ([\mathbf{C}\Delta\mathbf{R}(\mathbf{t}) + \mathbf{C}\Delta\mathbf{D}(\mathbf{t})]^T [\mathbf{C}\Delta\mathbf{R}(\mathbf{t}) + \mathbf{C}\Delta\mathbf{D}(\mathbf{t})])^{-1} [\mathbf{C}\Delta\mathbf{R}(\mathbf{t}) + \mathbf{C}\Delta\mathbf{D}(\mathbf{t})]^T [\mathbf{C}\Delta\mathbf{I}(\mathbf{t}) + \mathbf{C}\Delta\mathbf{R}(\mathbf{t}) + \mathbf{C}\Delta\mathbf{D}(\mathbf{t})].$$

Similarly, the case fatality rate $(\hat{\beta})$ and case recovery rate $(\hat{\gamma})$ can be estimated as:

$$\hat{\beta} = ([\mathbf{C}\Delta\mathbf{I}(\mathbf{t})]^T [\mathbf{C}\Delta\mathbf{I}(\mathbf{t})])^{-1} [\mathbf{C}\Delta\mathbf{I}(\mathbf{t})]^T [\mathbf{C}\Delta\mathbf{R}(\mathbf{t})],$$

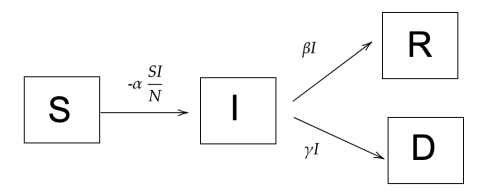
$$\hat{\gamma} = ([\mathbf{C}\Delta\mathbf{I}(\mathbf{t})]^T [\mathbf{C}\Delta\mathbf{I}(\mathbf{t})])^{-1} [\mathbf{C}\Delta\mathbf{I}(\mathbf{t})]^T [\mathbf{C}\Delta\mathbf{D}(\mathbf{t})].$$

3.2. Continuous model

In the continuous, the number of people in each class is a function of conitnuous time. So S(t) denotes the susceptible people at a time t. The mean-field kinetics of the SIRD epidemic evolution is described by the following system of differential equations:

$$\begin{split} \frac{dS}{dt} &= -\frac{\alpha}{N} S(t) I(t), \\ \frac{dI}{dt} &= \frac{\alpha}{N} S(t) I(t) - \beta I(t) - \gamma I(t), \\ \frac{dR}{dt} &= \beta I(t), \\ \frac{dD}{dt} &= \gamma I(t), \end{split}$$

with initial condition $[S(t_0), I(t_0), R(t_0), D(t_0)]$ for some initial time t_0 . The parameter α is the infection rate, i.e. the probability per unit time that a susceptible individual contracts the disease when entering in contact with an infected person. The parameters β and γ denote, respectively, the recovery and death rates. This scheme has good chances to capture at least the gross features of the full time course of the outbreak.



{Above

figures shows SIRD model classes and change per unit time shown above arrows.}

4 Results

In this section we present our results for basic reproduction number, case fatality rate, case recovery ratios and time series forecasting for Germany and India.

4.1 Results for Germany

We first use the discrete model to find these parameters and predict time series for Germany. We use the time series data available and for the required vectors $\mathbf{C}\Delta\mathbf{X}(T)$ for X=I,R,D.

```
country = "Germany" #Country chosen

#Extract country data from countries data
germany_cnf_melt = confirmed_countries_melt[which(confirmed_countries_melt$Country.Region == country),

#Extract confirmed number as a vector
germany_cnf = germany_cnf_melt$value

#Extract delta infected vector delta_cnf(t) = I(t) - I(t-1)
germany_delta_cnf = diff(germany_cnf)

#Cumulative sum of delta_cnf
germany_cum_delta_cnf = cumsum(germany_delta_cnf)

germany_deaths_melt = deaths_countries_melt[which(deaths_countries_melt$Country.Region == country),]

#Extract deaths number as a vector
germany_deaths = germany_deaths_melt$value

#Extract delta deathsvector delta_deaths(t) = D(t) - D(t-1)
```

```
#Cumulative sum of delta_deaths
germany_cum_delta_deaths = cumsum(germany_delta_deaths)
#Extract germany data from countries recovered data
germany_recovered_melt = recovered_countries_melt[which(recovered_countries_melt$Country.Region == coun
#Extract recovered number as a vector
germany_recovered = germany_recovered_melt$value
#Extract delta recovered vector delta_recovered(t) = R(t) - R(t-1)
germany_delta_recovered = diff(germany_recovered)
#Cumulative sum of delta_recovered
germany_cum_delta_recovered = cumsum(germany_delta_recovered)
### Caluclating infected numbers from confirmed cases
#Extract infected number cases as a vector
germany_inf = germany_cnf - germany_recovered - germany_deaths
#Extract delta infected vector delta_inf(t) = I(t) - I(t-1)
germany_delta_inf = diff(germany_inf)
#Cumulative sum of delta_inf
germany_cum_delta_inf = cumsum(germany_delta_inf)
We then use different time windows to estimate the time evolution of these parameters.
###ESTIMATING CASE FATALITY RATIO
#Making data frame of cumulative data
germany_cum_data_full = data.frame(delta_inf= germany_delta_inf, cum_delta_inf=germany_cum_delta_inf, d
##VARY THE NUMBER OF DAYS CHOSEN FOR ANALYSIS
ndays = 65:97
gamma_data <- data.frame(matrix(ncol = 3, nrow = 0))</pre>
x <- c("est", "lwr", "upr")
colnames(gamma data) <- x</pre>
beta_data <- data.frame(matrix(ncol = 3, nrow = 0))</pre>
x <- c("est", "lwr", "upr")</pre>
colnames(beta_data) <- x</pre>
RO_data <- data.frame(matrix(ncol = 3, nrow = 0))</pre>
x <- c("est", "lwr", "upr")
```

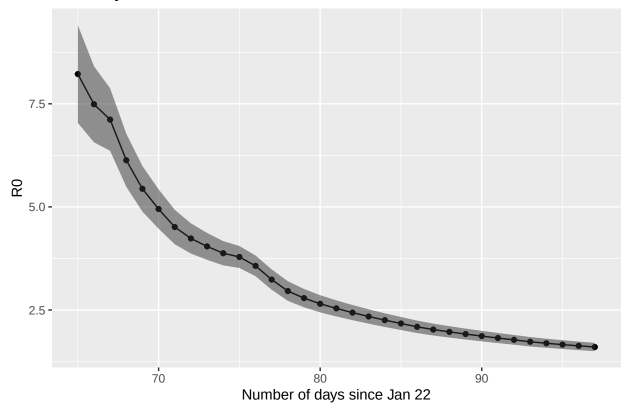
germany_delta_deaths = diff(germany_deaths)

```
colnames(RO_data) <- x</pre>
#loop over days window chosen
for (days in ndays) {
germany_cum_data = germany_cum_data_full[1:days, ]
#View(germany_cum_data)
#fitting a linear model for case fatality ratio
germany_gamma <- lm(cum_delta_deaths ~ cum_delta_inf -1 , data=germany_cum_data) # build linear regr
###ESTIMATING CASE RECOVERY RATIO
cor(germany_cum_delta_inf, germany_cum_delta_recovered)
#high correlation
#fitting a linear model for case recovery ratio
germany_beta <- lm(cum_delta_recovered ~ cum_delta_inf -1 , data=germany_cum_data) # build linear regr
###ESTIMATING RO
#fitting a linear model for case basic reproducibility number RO
germany_R0 <- lm(cum_delta_deaths + cum_delta_recovered + cum_delta_inf ~ I(cum_delta_recovered + cum_
##Storing estimations and conf intervals
conf = confint(germany_gamma)
gamma_row <- list(est = summary(germany_gamma)$coefficients[1], lwr = conf[1], upr = conf[2])</pre>
gamma_data = rbind(gamma_data, gamma_row)
conf = confint(germany_beta)
beta_row <- list(est = summary(germany_beta)$coefficients[1], lwr = conf[1], upr = conf[2])
beta_data = rbind(beta_data, beta_row)
conf = confint(germany_R0)
RO_row <- list(est = summary(germany_RO)$coefficients[1], lwr = conf[1], upr = conf[2])</pre>
RO_data = rbind(RO_data, RO_row)
}
```

We plot plot the estimates and the corresponding 99% confidence intervals for $R_0, \hat{\beta}, \hat{\gamma}$ as below.

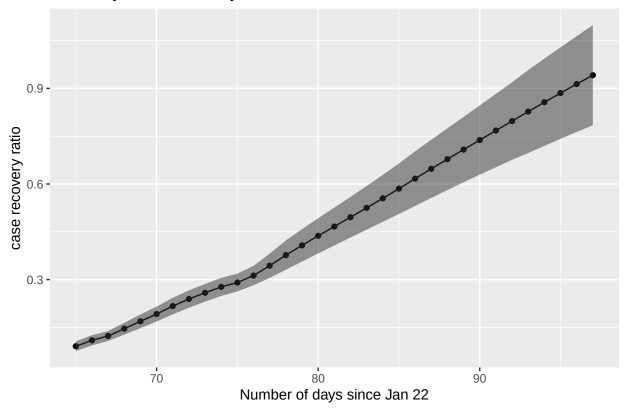
ggplot(RO_data, aes(ndays, est)) + geom_point() + geom_line(aes(ndays, est)) + geom_ribbon(aes(ymin=lwr

Germany: R0 estimate evolution

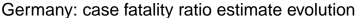


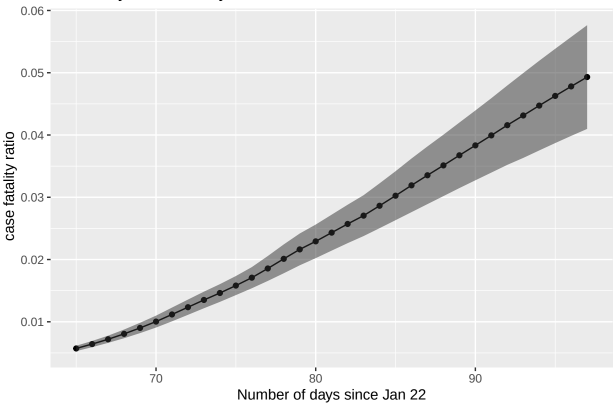
ggplot(beta_data, aes(ndays, est)) + geom_point() + geom_line(aes(ndays, est)) + geom_ribbon(aes(ymin=l

Germany: case recovery ratio estimate evolution



ggplot(gamma_data, aes(ndays, est)) + geom_point() + geom_line(aes(ndays, est)) + geom_ribbon(aes(ymin=

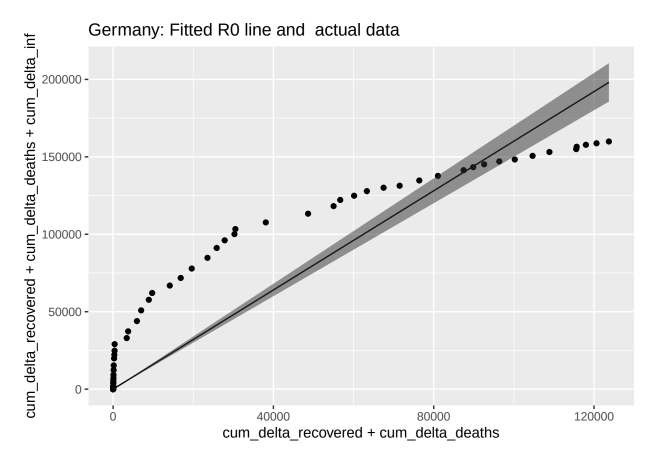




From these, we can see that the R_0 for Germany has come down from 8.2 to 1.6 (99% CI : [1.5 1.7]). This is good news and indicates flattening of the curve. Note that if $R_0 < 1$, the disease stops spreading. The case recovery ratio is going up and has reached approximately 0.9 while the case fatality ratio is about 0.049. We note here that since the estimate for case recovery and fatality only includes infected cases, hence the estimates for $\hat{\beta}$ and $\hat{\gamma}$ are only relibate for the early stages of the epidemic.

Since R_0 is estimated by a linear model, for curiosity we carry out some diagnostics to see how if the data actually fits to the linear plot. We the plot out fitted slope (which is equal to R_0) and observe that a rather poor fit which is expected because R_0 changes with time.

```
##Predicted RO using model
# add 'fit', 'lwr', and 'upr' columns to dataframe (generated by predict)
RO_predict <- cbind(germany_cum_data, predict(germany_RO, interval = 'confidence'))
RO_prediction = predict(germany_RO)
# plot the points (actual observations), regression line, and confidence interval
p <- ggplot(RO_predict, aes(cum_delta_recovered + cum_delta_deaths, cum_delta_recovered + cum_delta_deaths
p <- p + geom_point()
p <- p + geom_line(aes(cum_delta_deaths + cum_delta_recovered, RO_prediction))
p <- p + geom_ribbon(aes(ymin=lwr,ymax=upr), alpha=0.5) + ggtitle("Germany: Fitted RO line and actual operation)</pre>
```



Now, we solve the actualy differential equation to fit the evolution of the disease and forecast the time series. We intialize the values below. We ignore the initial data because of noisiness and low scale testing.

```
library(deSolve)
library(RColorBrewer)

Infected <- germany_inf[50:97]
Recovered <- germany_recovered[50:97]
Deaths <- germany_deaths[50:97]
Confirmed <- germany_cnf[50:97]
day <- 0:(length(Infected)-1)
N <- 830000

###edit 1: use different boundary condition
###init <- c(S = N-1, I = 1, R = 0)
init <- c(S = N-Infected[1] - Recovered[1] - Deaths[1], I = Infected[1], R = Recovered[1], D = Deaths[1]</pre>
```

Then, we define the differential changes in the quantities with respect to time.

```
SIR <- function(time, state, parameters) {
  par <- as.list(c(state, parameters))
  ####edit 2; use equally scaled variables
  with(par, { dS <- -alpha * (S/N) * I
  dI <- alpha * (S/N) * I - beta * I - gamma * I
  dR <- beta * I
  dD <- gamma * I</pre>
```

```
list(c(dS, dI, dR, dD))
})
}
```

Then we define an optimizer to find the optimum parameters to fit the confirmed cases curve with an initial guess. For this we also define a misfit function which is a simple L2 error function. The code is given below.

```
RSS.SIR <- function(parameters) {
    names(parameters) <- c("alpha", "beta", "gamma")
    out <- ode(y = init, times = day, func = SIR, parms = parameters)
    fit <- out[, 3] + out[, 4] + out[, 5]
    RSS <- sum((Confirmed- fit)^2)
    return(RSS)
}

lower = c(0, 0, 0)
upper = c(10, 1, 1) ###Limit box for parameters for L-BFGS-B

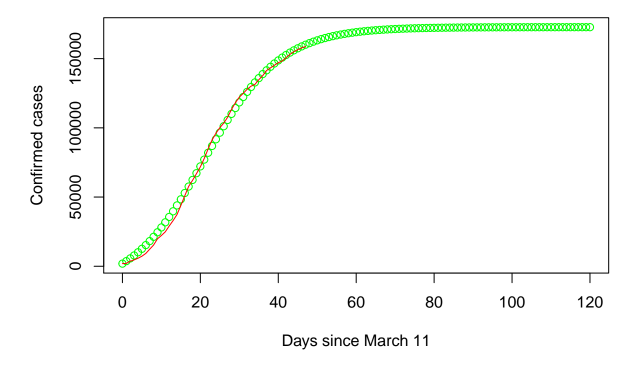
optimsstart <- c(0.7, 0.4, 0.2) #initial guess for parameters

set.seed(12)
Opt <- optim(optimsstart, RSS.SIR, method = "L-BFGS-B", lower = lower, upper = upper, hessian = TRUE)
#Opt$par</pre>
```

Once we have optimized, we can predict the case evolution as follows.

```
Opt_par <- Opt$par
names(Opt_par) = c("alpha", "beta", "gamma")
t <- 0:120
fit <- data.frame(ode(y = init, times = t, func = SIR, parms = Opt_par))
predict <- fit$I + fit$D + fit$R
plot(t, predict, col="green", xlab="Days since March 11", ylab="Confirmed cases")
lines(day, Confirmed, col="red")
title("Germany: Green is confirmed cases predicted by our model, red is actual data.")</pre>
```

ermany: Green is confirmed cases predicted by our model, red is actua



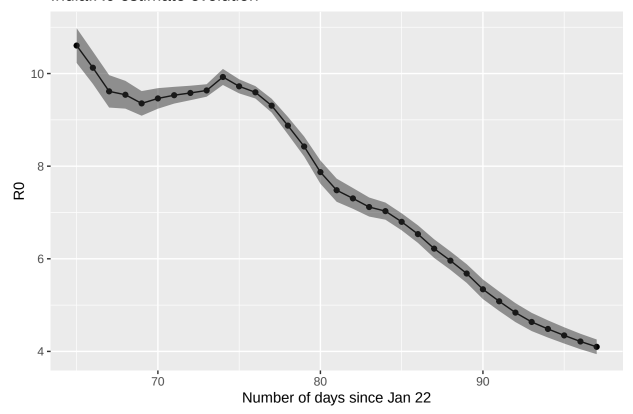
The results indicate that by the end of June, the confirmed cases will peak and the epidemic will end in Germany.

4.2 Results for India

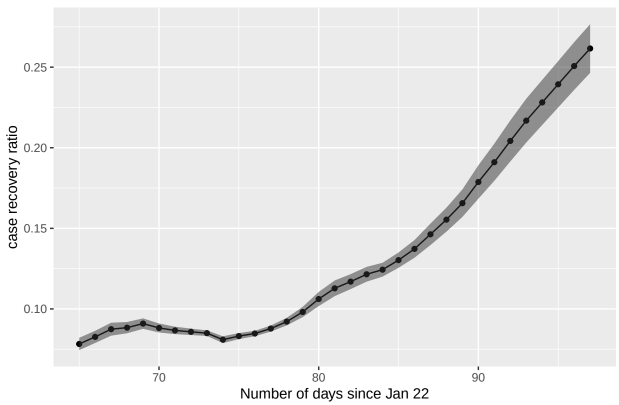
Now we repeat the same analysis for India. Note India is still in the early phase of the disease.

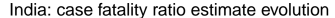
We plot plot the estimates and the corresponding 99% confidence intervals for $R_0, \hat{\beta}, \hat{\gamma}$ as below.

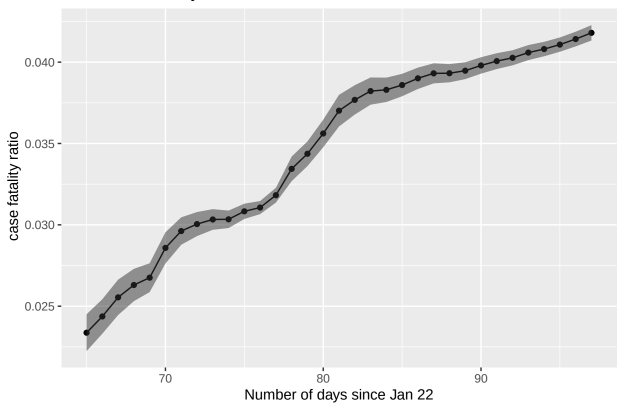
India:R0 estimate evolution



India: case recovery ratio estimate evolution

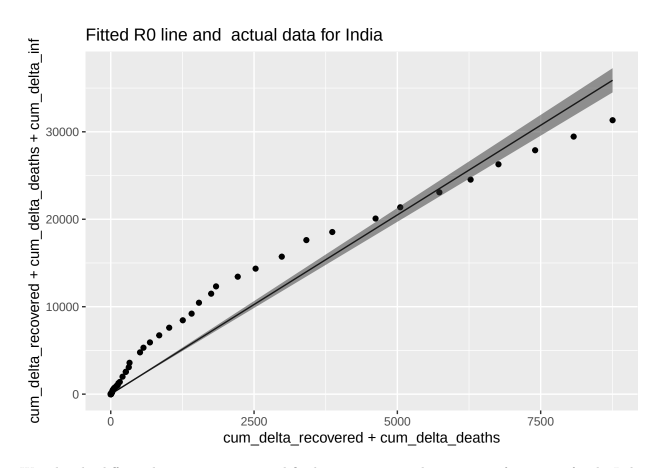






From these, we can see that the R_0 for India has come down from 10.6 to 4.09 (99% CI : [3.94 4.25]). Thus India is quite far away from the end of the epidemic. Note that if $R_0 < 1$, the disease stops spreading. The case recovery ratio is going up and has reached approximately 0.26 while the case fatality ratio is about 0.041.

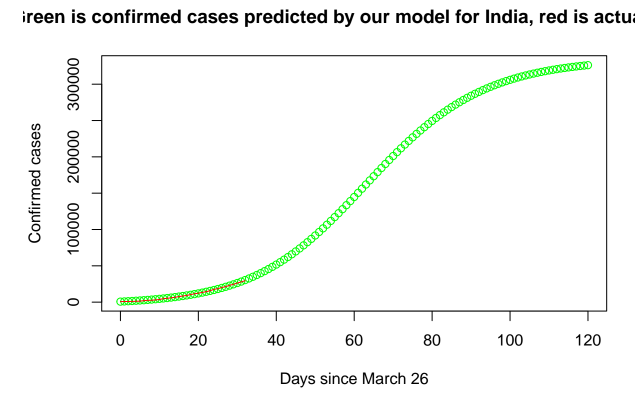
Since R_0 is estimated by a linear model, for curiosity we carry out some diagnostics to see how if the data actually fits to the linear plot. We the plot out fitted slope (which is equal to R_0) and observe that a rather poor fit which is expected because R_0 changes with time.



We solve the differntial equations system and fit the parameters to do time series forecasting for the India data.

Once we have optimized, we can predict the case evolution as follows.

ireen is confirmed cases predicted by our model for India, red is actual



The results indicate that the curve will peak in India by the end of July.

5. Conclusions and Future work

In this study, we used SIRD model commonly used in epidemiology to estimate the evolution of basic reprducibility number, case fatality ratio and case recovery ratio as the disease progresses. We also use it to fit the parameters to the data coming from Germany and India and use it to predict the evolution of disease. Our results indicate that the epidemic should end in germany by the end of June. India might have to wait till the end of July before the peak is reached. The recovery ratios obtained for India (about 26%) are consistent with the government estimates recently reported in business-standard. The prediction of beginning of curve flattening by July end are in line with the recent WHO estimates given WHO envoy interview.

6. Limitations

However, we must state here that there are several limitations to our predictions. First, the model itself is a simplistic model to study the disease as it assumes constant transmission rates among different classes. Also, the model does not take into account the asymptomatic cases which may be contributing to sopreading the disease. Secondly, the data itself might be unreliable as there might be severe underreporting of infected people because of lack of testing. Thirdly, the uplifting of lockdown may accelerate the spread of disease. A more complex model with these factors taken into account would be desirable for a better forecasting.

7. References

[1] Cleo Anastassopoulou "Lucia Russo, Athanasios Tsakris, Constantinos Siettos," Data-based analysis, modelling and forecasting of the COVID-19 outbreak", PLOS ONE (2020) (https://doi.org/10.1371/journal. pone.0230405)