

Situational awareness and forecasting for Norway

FHI COVID-19 modelling team

Week 2, 13 January 2021

Highlights:

• National epidemiological situation: Our models evaluate the present situation as unstable, following a clear worsening during the first weeks of December. The effective reproduction number R_{11} acting in our changepoint model from 27 December is estimated to be 1.24 (median, 95% CI 0.88-1.61). The estimated probability that R_{11} is larger than 1 is 89%. There is a slight reduction of this effective reproduction number, compared to the estimate reported on January 6. These estimates do not likely capture all the effects of the interventions introduced in Norway on January 4, because of the time delay between infections and testing. However, they might be affected by the reduced contacts during the Christmas vacations. The SMC model estimates the 7-days averaged effective reproduction number one week ago to be 1.04 (95% CI 0.83-1.25). In this model, the estimated probability that the daily reproduction number one week ago was above 1 is 63%. The SMC model indicates a tendency to a reduction of the averaged effective reproduction number in the days since after Christmas. We conclude that the reproduction number is currently around 1.1-1.2, slightly improving at national aggregated level.

Since the start of the epidemic, we estimate that in total 106.000 (95% CI 91.000- 122.000) persons in Norway have been infected. The current estimate of the detection probability is around $\sim 65\%$.

- National forecasting: In one week, we estimate ca 1.500 new cases per day (median; 95%CI 640-3.000), and a prevalence (total number of infected people in Norway) of 8.700 (median; 95% CI 4.400-15.600). Hospitalisations in one week are estimated to be 210 (median 95% CI 142-300) and patients on ventilator treatment are estimated to be 29 (median 95% CI 17-43); the corresponding three-week projections are (95% CI 130 650) and (95% CI 18 80). The probability that the surge capacity will exceed 500 ventilator beds is estimated at 54%.
- Regional epidemiological situation and forecasting: We estimate the reproduction number since 21 December in Oslo to be 1.48 (mean, 95% CI 1.05-1.96) and in Rogaland 1.43 (mean, 95% CI 0.40-2.38) Viken 1.05 (mean, 95% CI 0.77 1.36), Trøndelag 1.21. (0.3 2.2). The other counties have reproductions numbers around 1. Our estimates have high uncertainty. Compared to a week ago, the estimates of the effective reproduction numbers in Oslo, and Rogaland increased, the ones in Viken and Trøndelag were roughly stable. As usual, estimates of the effective reproduction numbers exclude imported cases, which are generated elsewhere. Oslo: The number of new cases per day is estimated to be 164 (median, 95% CI 75 335) on 17 January, and on 24 January (95% CI 86-560). Hospital prevalence in one week is estimated to be 26 (median; 95% CI 10-46).
- Telenor mobility data, local mobility and foreign visitors: Inter-municipality mobility, measured as outgoing mobility of mobile phones from each municipality has declined over the last weeks to very low levels. There was a huge peak in the first weeks of December. In the last days we see a slight increase in mobility, in connection with the new year. The analysis of foreign roamers has been updated to show visitors from the nine most-visiting nationalities to Norway. After the drop during Christmas, there is now an increasing trend in January 2021 in foreign visitors coming to Norway, from Poland in particular. We present new plots with number of visitors for each county.



What this report contains:

This report presents results based on a mathematical infectious disease model describing the geographical spread of COVID-19 in Norway. The model consists of three layers:

- Population structure in each municipality.
- Mobility data for inter-municipality movements (Telenor mobile phone data).
- Infection transmission model (SEIR-model)

The model produces estimates of the current epidemiological situation at the municipality, county (fylke), and national levels, a forecast of the situation for the next three weeks, and a long term prediction. We run three different models built on the same structure indicated above: (1) a national changepoint model, (2) a regional changepoint model and (3) a national Sequential Monte Carlo model, named SMC model.

How we calibrate the model: The national changepoint model is fitted to Norwegian COVID-19 hospital incidence data from March 10 until yesterday, and data on the laboratory-confirmed cases from May 1 until yesterday. We do not use data before May 1, as the testing capacity and testing criteria were significantly different in the early period.

Note that the results of the national changepoint model are not a simple average or aggregation of the results of the regional changepoint model because they use different data. The estimates and predictions of the regional model are more uncertain than those of the national model. The regional model has more parameters to be estimated and less data in each county; lack of data limits the number of changepoints we can introduce in that model. In the regional changepoint model, each county has its own changepoints and therefore a varying number of reproduction numbers. Counties where the data indicate more variability, have more changepoints.

The national SMC model is also calibrated both to the hospitalisation incidence data (same data as described above) and the laboratory-confirmed cases.

Telenor mobility data: The mobility data account for the changes in the movement patterns between municipalities that have occurred since the start of the epidemic.

How you should interpret the results: The model is stochastic. To predict the probability of various outcomes, we run the model many times in order to represent the inherent randomness.

We present the results in terms of mean values, 95% confidence intervals, medians, and interquartile ranges. We emphasise that the confidence bands might be broader than what we display, because there are several sources of additional uncertainty which we currently do not fully explore: firstly, there are uncertainties related to the natural history of SARS-CoV-2, including the importance of asymptomatic and presymptomatic infection. Secondly, there are uncertainties related to the timing of hospitalisation relative to symptom onset, the severity of the COVID-19 infections by age, and the duration of hospitalisation and ventilator treatment in ICU. We continue to update the model assumptions and parameters in accordance with new evidence and local data as they become available. A full list of all updates can be fount at the end of this report.

Estimates of all reproduction numbers are uncertain, and we use their distribution to assure appropriate uncertainty of our predictions. Uncertainties related to the model parameters imply that the reported effective reproductive numbers should be interpreted with caution.

When we forecast beyond today, we use the most recent reproduction number for the whole future, if not explicitly stated otherwise.

In this report, the term patient in ventilator treatment includes only those patients that require either invasive mechanical ventilation or ECMO (Extracorporeal membrane oxygenation).



1 Estimated national reproduction numbers

Calibration of our national changepoint model to hospitalisation incidence data and test data leads to the following estimates provided in table 1. Figure 1 shows the estimated daily number of COVID-19 patients admitted to hospital (1a) and the estimated daily number of laboratory-confirmed SARS-CoV-2 cases (1b), with blue medians and interquantile bands, which are compared to the actual true data, provided in red. The uncertainty captures the uncertainty in the calibrated parameters in addition to the stochastic elements of our model and the variability of other model parameters.

Reff	Period
3.11/3.12(2.31-3.95)	From Feb 17 to Mar 14
0.52/0.52(0.39-0.63)	From Mar 15 to Apr 19
0.66/0.66(0.21-1.19)	From Apr 20 to May 10
0.58/0.58(0.07-1.15)	From May 11 to Jun 30
0.77/0.75(0.06-1.52)	From Jul 01 to Jul 31
1.01/1.01(0.63-1.39)	From Aug 01 to Aug 31
0.97/0.97(0.68-1.26)	From Sep 01 to Sep 30
1.24/1.24(0.94-1.54)	From Oct 01 to Oct 25
1.46/1.47(1.1-1.91)	From Oct 26 to Nov 04
0.84/0.84(0.74-0.91)	From Nov 05 to Nov 30
1.06/1.06(0.96-1.16)	From Dec 01 to Dec 26
1.24/1.25(0.88-1.61)	From Dec 27
/	

Median/Mean (95% credible intervals)

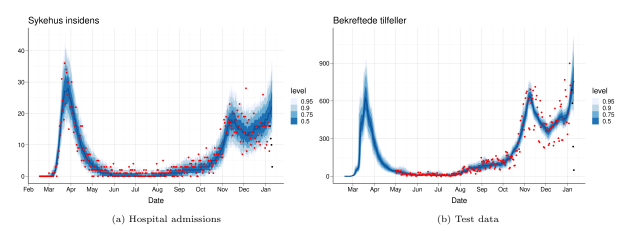


Figure 1: A comparison of true data (red) and predicted values (blue) for hospital admissions and test data. The last four data points (black) are assumed to be affected by reporting delay. B) Comparison of our simulated number of positive cases, with blue median and interquartile bands to the actual true number of positive cases, provided in red. The uncertainty captures the uncertainty in the calibrated parameters, in addition to the stochastic elements of our model and the variability of other model parameters. Note that we do not capture all the uncertainty in the test data–our blue bands are quite narrow. This is likely because we calibrate our model parameters on a 7-days moving average window of test data, instead of daily. This is done to avoid overfitting to random daily variation. Moving averages over 7 days are less variable than the daily data.



In figure 2, we show how our national model fits the national hospital prevalence data (2a) and the daily number of patients receiving ventilator treatment (2b). Those data sources are not used to estimate the parameters, and can therefore be seen as a validation of the model assumptions.

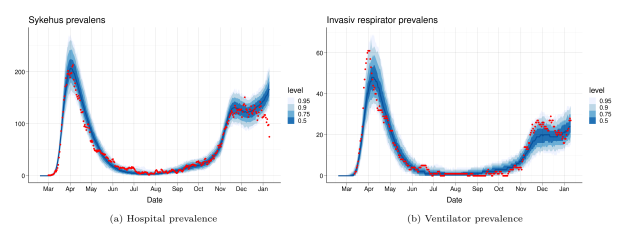


Figure 2: A comparison of true data (red) and predicted values (blue) for hospital and respirator prevalence.

1.1 National SMC-model: Estimated daily reproduction numbers

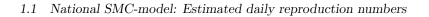
In the SMC-model, we allow for estimation of a different reproduction number for each day t. To reduce spurious fluctuation, we report a 7-days moving average, R(t), representing the average reproduction number for the whole week before day t. However, until March 8 we keep the reproduction number constant. By assuming a time varying reproduction number R(t), we can detect changes without introducing explicit changepoints. Thus, we can easier detect unexpected changes.

The SMC model uses the daily number of new admissions to hospital and the daily number of positive and negative lab-confirmed tests, to estimate all its parameters. Because of the time between infection and the possibility to be detected as positive by a test, and because if a delay in reporting tests, the data contain information on the transmissibility until a week before the end of the data (today).

The parameters π_0 and π_1 related to the probability to detect a positive case by testing are estimated off-line.

The figure below shows the SMC estimate of the 7-day-average daily reproduction number R(t) from the start of the epidemic in Norway and until today. In the figure we plot the 95% confidence interval and quantiles of the estimated posterior distribution of R(t).





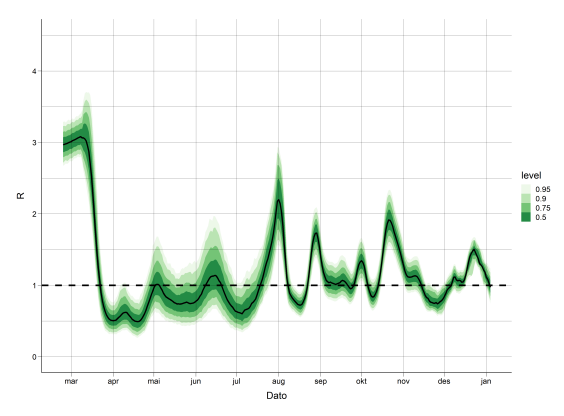


Figure 3: R(t) estimates using a Sequential Monte Carlo approach calibrated to hospitalisation incidence and test data. The large uncertainty during the last 7 days reflects the lack of available data due to the transmission delay, test delay, time between symptoms onset and hospitalisation. The green band shows the 95% posterior credibility interval. As we use test data only from 1 August, the credibility interval becomes more narrow thereafter.



2 National estimate of cumulative (total) number of infections

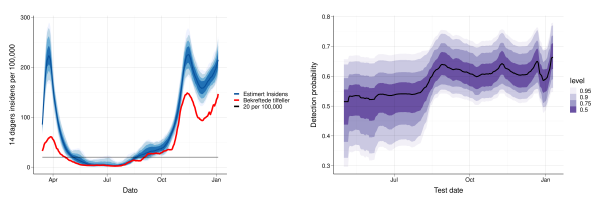
The national changepoint model estimates the total number of infections and the symptomatic cases that have occurred (Table 2).

Figure 4a shows the modelled expected daily incidence (blue) and the observed daily number of laboratoryconfirmed cases (red). When simulating the laboratory-confirmed cases, we also model the detection probability for the infections (both symptomatic, presymptomatic and asymptomatic), Figure 4b. There are two differences between this estimate of the detection probability and the previous one provided in figure 4a. In figure 4b, we calibrate our model to the true number of positive cases, instead of using the test data directly. Furthermore, in figure 4a we use a parametric model to estimate the detection probability that depends on the true total number of tests performed.

Table 2: Estimated cumulative number of infections, 2021-01-10

Region	Total	No. confirmed	Fraction reported	Min. fraction
Norway	$106379 \ (91308; \ 122135)$	55473	52%	45%

Fraction reported=Number confirmed/number predicted; Minimal fraction reported=number confirmed/upper CI



(a) Number of laboratory-confirmed cases vs model-based esti-(b) Estimated detection probability for an infected case per calmated number of new infected individuals endar day

Figure 4



3 National 3-week predictions: Prevalence, Incidence, Hospital beds and Ventilator beds

The national changepoint model estimates the prevalence and daily incidence of infected individuals (asymptomatic, presymptomatic and symptomatic) for the next three weeks, aggregated to the whole of Norway (table 3). In addition, the table shows projected national prevalence of hospitalised patients (hospital beds) and prevalence of patients receiving ventilator treatment (ventilator beds). The projected epidemic and healthcare burden are illustrated in figure 5.

Table 3: Estimated national prevalence, incidence, hospital beds and ventilator beds. Median/Mean (CI)

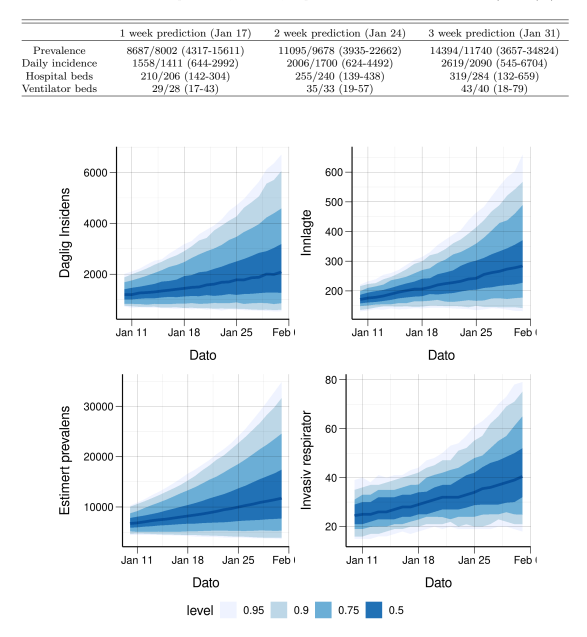


Figure 5: National 3 week predictions for incidence (top left), prevalence (bottom left), hospital beds (top right) and ventilator beds (bottom right)



4 National long-term scenarios: Prevalence, Hospital beds and Ventilator beds

Results from 12-month scenario of the calibrated national changepoint model, showing expected prevalence (Figure 6a), hospital beds (Figure 6b) and ventilator beds (Figure 6c), in the case where the transmissibility stays the same as today. The figures are made using the 200 candidate models, where the reproductive numbers are varying according to their estimated uncertainty as of today. The confidence intervals shown in the plots are two-tailed around the median, and therefore the upper 95% level shows the 97.5% boundary. Note that age-specific attack rate after 21 days of projection is assumed to follow the demography in each county, instead of being informed by the current age-distribution of the laboratory-confirmed cases.

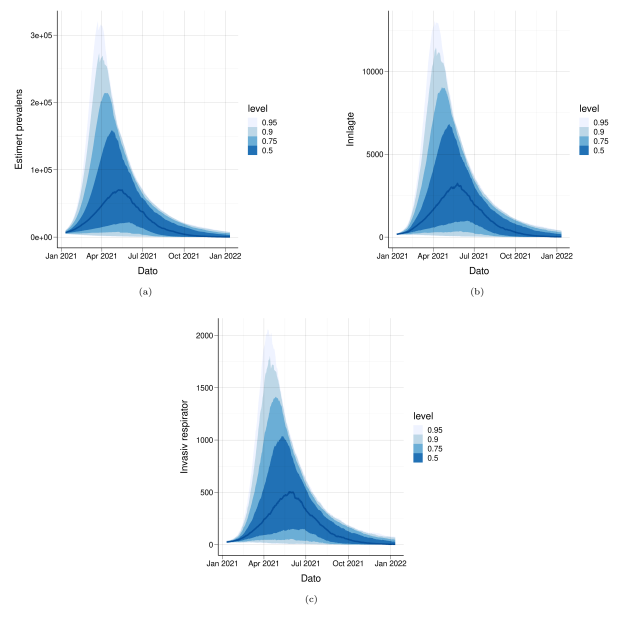


Figure 6: Long-term predictions for prevalence (a), hospital beds (b) and ventilator beds (c)

The probability of a surge capacity need above 500 ICU ventilator beds is 54 %. We estimate the



probability of a surge capacity need above 1000~ICU beds to be equal to 30~%.



5 National scenario-based long-term predictions: Hospital beds and Ventilator beds

Here we show how the epidemic estimated from the national changepoint model will develop under three assumed epidemiological scenarios, by fixing the effective reproduction number to be 1.2, 1.3 or 1.4, from today. We show the daily number of COVID-19 patients in hospital, including patients receiving ventilator treatment, (Figure 7, and the daily number of patients on ventilator treatment, figure 8. Note that age-specific attack rate after 21 days of projection is assumed to follow the demography in each county, instead of being informed by the current age-distribution of the laboratory-confirmed cases. Additional information about the total attack rate (cumulative incidence) and healthcare burden and surge capacity for these scenarios are provided in Table 4.

Table 4: Predicted numbers of total infected, total number of hospitalisations, total number needing ventilator treatment, and the predicted peak number in ward (not in respirator), hospitalised (both with and without ventilator treatment) and ventilated treatments based on three different scenarios with R effective equal to 1.1, 1.2 and 1.3.

	Reff=1.2	Reff=1.3	Reff=1.4
Total:			
Attack rate (infected)	1.640.000(1.620.000 - 1.660.000)	2.230.000(2.210.000 - 2.240.000)	2.690.000(2.680.000 - 2.700.000)
Hospitalisations	51.000(50.100 - 51.600)	69.500(68.800 - 70.100)	84.000(83.400 - 84.600)
Patients on ventilator	4.390(4.270 - 4.520)	5.970(5.820 - 6.080)	7.210 (7.060 - 7.350)
At peak			
Hospital beds, excl. vent.	1.980(1.850 - 2.110)	3.880(3.740 - 4.040)	5.980(5.810 - 6.130)
Hospital beds, incl. vent.	2.330(2.190 - 2.500)	4.580(4.430 - 4.760)	7.060(6.870 - 7.210)
Ventilator beds	377(349 - 411)	728(686 - 771)	1.110(1.060 - 1.160)

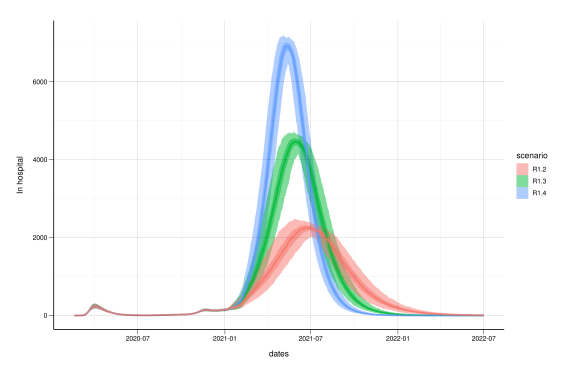


Figure 7: Predicted number of COVID-19 patients in hospital based on three different scenarios with R effective equal to 1.1, 1.2 and 1.3. Shaded areas show interquartile range and 95% confidence interval around the median.



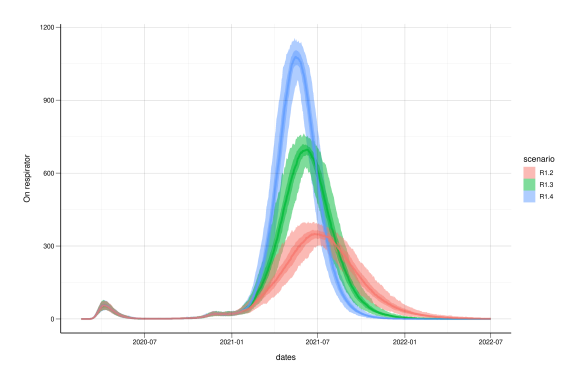


Figure 8: Predicted number of COVID-19 patients needing ventilator treatment based on three different scenarios with R effective equal to 1.1, 1.2 and 1.3. Shaded areas show interquartile range and 95% confidence interval around the median.



6 Estimated regional reproduction numbers

Calibration of our regional changepoint model to hospitalisation incidence data and test data leads to the following estimates for current regional reproduction numbers by county (Table 5). A full list of all regional reproduction numbers can be found at the end of the report.

Below we show the estimated daily number of COVID-19 patients admitted to hospital and the estimated daily number of laboratory-confirmed SARS-CoV-2 cases for each county. Model estimates are shown with blue medians and interquantile bands, which are compared to the actual true data, provided in red. The blue bands describe the uncertainty in the calibrated parameters, in addition to the stochastic elements of our model. Last four data points are shown in black as they may be affected by reporting delay.

R	Parameter	County	From	$\Pr(R>1)$
1.48(1.05 - 1.96)	$\mathbf{R7}$	Oslo	2020-12-21	0.98
1.43 (0.4-2.38)	R6	Rogaland	2020-12-21	0.79
0.99(0.11-2.16)	R6	Møre og Romsdal	2020-12-21	0.46
0.93(0.08-2.12)	R6	Nordland	2020-12-21	0.42
1.05(0.77 - 1.36)	R7	Viken	2020-12-21	0.62
0.69(0.04-1.59)	R6	Innlandet	2020-12-21	0.23
0.99(0.13-2)	R6	Vestfold og Telemark	2020-12-21	0.46
0.9(0.07-2.1)	R6	Agder	2020-12-21	0.4
0.92(0.1-1.88)	R7	Vestland	2020-12-21	0.41
1.21(0.29-2.23)	R6	Trøndelag	2020-12-21	0.64
0.86(0.05-1.95)	R6	Troms og Finnmark	2020-12-21	0.38

Table 5: Estimated current regional reproduction numbers

Mean and 95% credible intervals

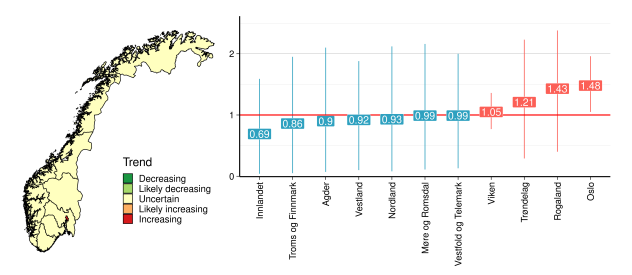
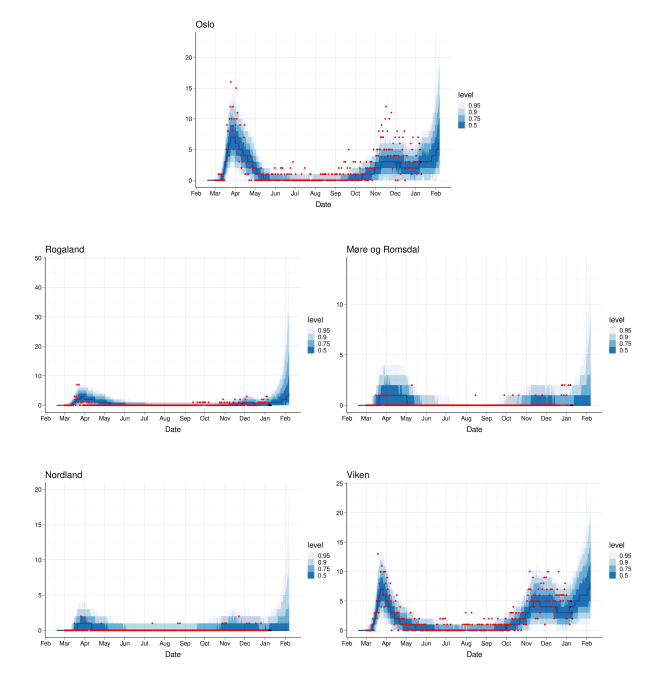


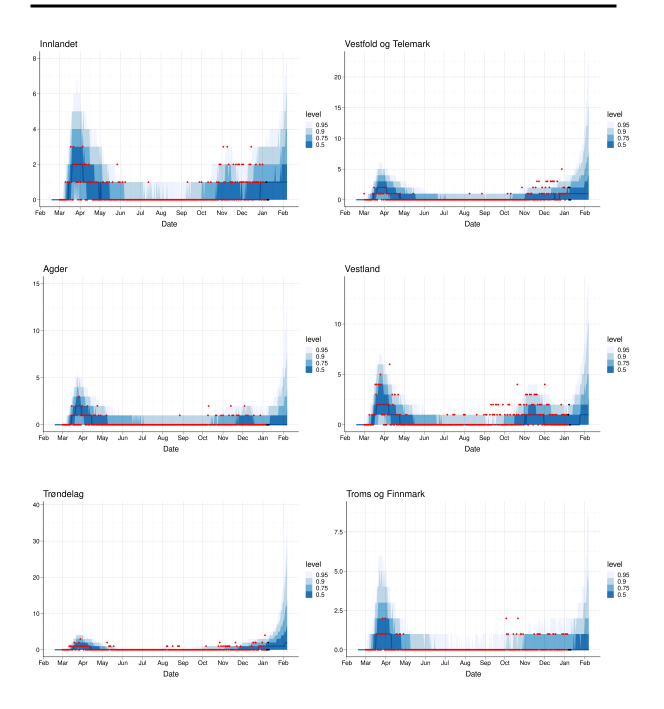
Figure 9: The map shows the direction of the trend in incidence in the counties based on the latest reproduction numbers shown in the other chart. The trend is increasing if the probability that the latest reproduction number is above one is above 95%, the trend is likely increasing if this probability is between 80% and 95%, the trend is uncertain if the probability is between 20% and 80%, the trend is likely decreasing if the probability is between 5% and 20% and is decreasing if the probability that the latest R is above one is less than 5%.



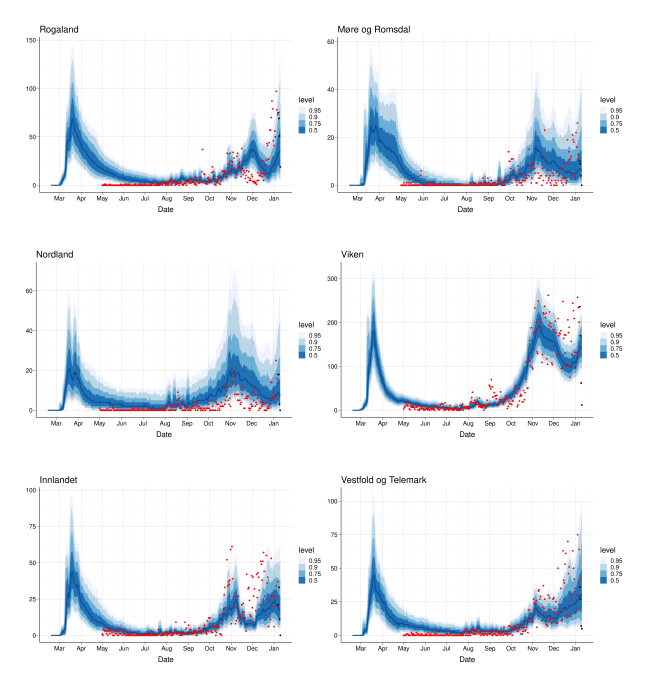


Estimated vs observed hospital incidence data by county:



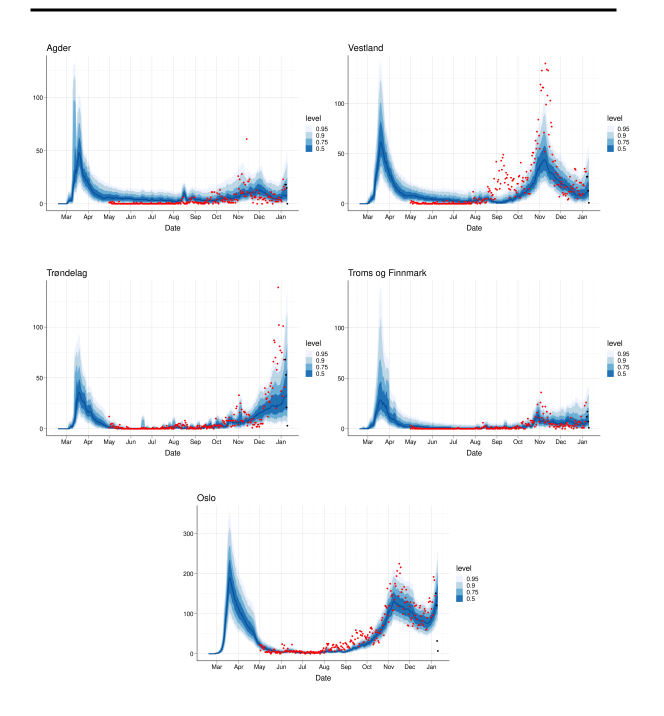






Estimated and observed lab-confirmed test data by county:







7 Regional 3-week predictions: Cumulative (total) incidence and Prevalence

Below is shown the estimated short-term forecasting of total incidence of infected individuals (table 6), daily incidence (table 7) and prevalence (table 8) for each county.

Region	Total	No. confirmed	Fraction reported	Min. fraction
Oslo	23010 (17133; 31277)	15181	66%	49%
Rogaland	7852 (5635; 10869)	2959	38%	27%
Møre og Romsdal	3197 (1899; 5333)	1060	33%	20%
Nordland	3832(1973;6955)	795	21%	11%
Viken	26549 (20174; 34494)	17701	67%	51%
Innlandet	4873 (3418; 7015)	3012	62%	43%
Vestfold og Telemark	5347 (3818; 7316)	2619	49%	36%
Agder	4164 (2720; 6644)	1331	32%	20%
Vestland	6544 (4402 ; 10174)	5892	90%	58%
Trøndelag	3870 (2505; 6170)	3078	80%	50%
Troms og Finnmark	2663(1469;5539)	1131	42%	20%

Table 6: Estimated cumulative number of infections, 2021-01-10

Fraction reported=Number confirmed/number predicted; Minimal fraction reported=number confirmed/upper CI

Table 7: Predicted incidence per day: Median/Mean (CI)

Region	1 week prediction (17 Jan)	2 weeks prediction (24 Jan)	3 weeks prediction (31 Jan)
Agder	13/24 (1-123)	14/37 (0-232)	19/63 (1-453)
Innlandet	29/36 (8-103)	29/41 (6-152)	35/53 (7-207)
Møre og Romsdal	12/25 (1-115)	12/40 (0-205)	12/70(0-418)
Nordland	14/27 (1-141)	12/41 (0-287)	12/68 (0-538)
Oslo	164/174 (74-335)	220/246 (86-564)	288/338 (94-927)
Rogaland	71/103 (7-380)	98/178 (5-768)	140/318(3-1615)
Troms og Finnmark	9/17 (1-83)	8/23 (0-154)	8/35 (0-293)
Trøndelag	53/81 (6-303)	64/126(2-602)	74/208 (2-1166)
Vestfold og Telemark	39/58 (7-223)	40/80 (5-374)	45/117 (5-682)
Vestland	29/42 (5-167)	28/57 (2-277)	33/85 (2-460)
Viken	259/273 (131-485)	266/295 (126-593)	297/336 (116-757)

Table 8: Predicted prevalence. Number of infectious individuals (asymptomatic plus pre-symptomatic plus symptomatic) per day. Median/Mean and 95 perc. CI for three weeks prediction.

Region	17 Jan	24 Jan	31 Jan	low CI, 31 Jan	high CI, 31 Jan
Agder	82/134	98.5/201	148/343	22	2159
Innlandet	190/220	190.5/248	257/337	70	1044
Møre og Romsdal	74/136	76.5/206	92.5/347	12	1899
Nordland	90/149	86/212	89/342	11	2381
Oslo	871.5/924	1190.5/1304	1543.5/1779	539	4578
Rogaland	382.5/526	519/875	719/1515	35	7295
Troms og Finnmark	58/98	55/128	58/192	9	1414
Trøndelag	304.5/429	366/646	448.5/1040	36	5259
Vestfold og Telemark	233.5/322	253/432	287.5/620	51	3335
Vestland	179/244	186.5/322	246/488	41	2329
Viken	1568/1645	1610.5/1742	1721.5/1919	746	4071



8 Regional 3-week predictions: Hospital beds and ventilator beds

Below is shown the estimated short-term forecasting of expected hospital prevalence (table 9) and patients on ventilator treatment for each county (table 10).

Region	1 week prediction (17 Jan)	2 weeks prediction (24 Jan)	3 weeks prediction (31 Jan)
Agder	2/3 (0-12)	2/4 (0-19)	2/6 (0-32)
Innlandet	6/7 (1-19)	7/8 (1-21)	7/9 (1-29)
Møre og Romsdal	2/3 (0-12)	2/4 (0-18)	2/6 (0-30)
Nordland	2/4 (0-16)	3/5 (0-25)	3/7(0-40)
Oslo	26/26(10-46)	27/28 (9-54)	31/34(11-72)
Rogaland	7/9 (1-26)	10/13(1-46)	14/22(1-91)
Troms og Finnmark	2/3 (0-11)	2/3 (0-15)	2/4 (0-20)
Trøndelag	8/10 (1-28)	9/13 (1-43)	11/18 (1-77)
Vestfold og Telemark	8/9 (1-25)	8/11 (1-34)	9/14 (1-57)
Vestland	3/5 (0-16)	5/6 (0-24)	6/9 (0-38)
Viken	38/39 (19-64)	44/46 (20-80)	50/53 (23-98)

Table 9: Number of hospitalisation beds occupied by Covid-19 patients: Median/Mean (CI)

Table 10: Number of ICU beds occupied by Covid-19 patients: Median/Mean (CI)

Region	1 week prediction (17 Jan)	2 weeks prediction (24 Jan)	3 weeks prediction (31 Jan)
Agder	0/0 (0-2)	0/1 (0-3)	0/1 (0-4)
Innlandet	1/1 (0-4)	1/1 (0-4)	1/1 (0-5)
Møre og Romsdal	0/0 (0-2)	0/1 (0-3)	0/1 (0-4)
Nordland	0/1 (0-3)	0/1 (0-4)	0/1 (0-5)
Oslo	4/4 (0-8)	4/4 (0-9)	4/5(1-11)
Rogaland	1/1(0-4)	1/2(0-6)	2/2(0-10)
Troms og Finnmark	0/0(0-2)	0/0 (0-3)	0/1(0-3)
Trøndelag	1/1(0-5)	1/2(0-6)	1/2 (0-10)
Vestfold og Telemark	1/1 (0-5)	1/2 (0-5)	1/2(0-7)
Vestland	0/1 (0-3)	1/1 (0-4)	1/1 (0-5)
Viken	5/5(1-11)	6/6 (1-12)	7/7 (2-15)



9 14-day trend analysis of confirmed cases and hospitalisations by county

To estimate recent trends in hospitalisation and number of positive tests, we present results in table 11 based on a negative binomial regression where we account for weekend effects. We exclude the last three days to avoid problems of reporting delay and fit the model using data from 17 days to 3 days before the current date. We fit a separate trend model for confirmed cases and for hospital incidence. We only fit a trend model if there has been more than 5 cases or hospitalisations in the 14-day period.

	Average daily inc	rease last 14 days	Doubling Time (days)		
County	Hospitalisations	Cases	Hospitalisations	Cases	
Agder	Not enough data	-12.6 (-18.1, -7) %	Not enough data	-5.1 (-3.5, -9.5)	
Innlandet	-2.5 (-14.1, 10.3) %	4.4 (-1.4, 10.7) %	-27.2(-4.6, 7.1)	15.9(-48.2, 6.8)	
Møre og Romsdal	Not enough data	-0.8 (-11.5, 11.4) %	Not enough data	-86.1 (-5.7, 6.4)	
Nordland	Not enough data	-6.1 (-13.3, 1.5) %	Not enough data	-11 (-4.9, 45.1)	
Norge	-0.1 (-3.4, 3.4) %	-3.9 (-5, -2.8) %	-1155.4 (-20, 20.8)	-17.4 (-13.4, -24.5)	
Oslo	-3.1 (-9.5 , 3.7) %	-4.4 (-5.6, -3.2) %	-21.9(-6.9, 19.3)	-15.4 (-12, -21.4)	
Rogaland	17.3 (-6.2, 56.1) %	-9.5 (-14.3, -4.6) %	4.3 (-10.9, 1.6)	-6.9 (-4.5, -14.8)	
Troms og Finnmark	Not enough data	-12.8 (-19.7, -5.8) %	Not enough data	-5 (-3.2, -11.6)	
Trøndelag	Not enough data	-8.8 (-13.6, -3.9) %	Not enough data	-7.5 (-4.7, -17.3)	
Vestfold og Telemark	15.8 (-3.9, 45.2) %	-5.3 (-9.8, -0.7) %	4.7 (-17.4, 1.9)	-12.7 (-6.7, -106.1)	
Vestland	-10.7 (-20.6, -0.6) %	-11 (-13.1, -8.9) %	-6.1 (-3, -124.7)	-5.9(-4.9, -7.5)	
Viken	4 (-2.6, 11.1) %	-1.2 (-2.3, -0.1) %	17.8(-26.7, 6.6)	-56.6 (-29.6, -650.5	

Table 11: T	rend anal	ysis for	the	last	14	days
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10 Scenario-based short-term predictions for Oslo:

Oslo has experienced increasing infection levels in the last months. Rising case numbers can lead to less efficient contact tracing due to a lack of resources. This, in turn, can cause the reproductive number to increase. To explore the short-term consequences of a less effective contact tracing in Oslo, we compare projections of the regional changepoint model, where the current reproduction number in Oslo is increased to 1.55; 1.60 and 1.65 from today, respectively. In these scenarios we assume no change to the reproductive numbers in the other counties. Table 12 and Figure compares these projected scenarios with a projection of the current epidemiological situation in Oslo.

Table 12: 4 week predictions in Oslo: Prevalence and Incidence (mean/median(CI))

Scenario	Prevalence	Incidence
Current	2063/1946 (1075-3496)	389/358 (207-673)
R = 1.55	2225/2104 (1040-3832)	424/402 (192-738)
R = 1.60	2428/2299 (1257-4260)	466/438 (232-831)
R = 1.65	2640/2486 (1318-4574)	511/488 (241-882)

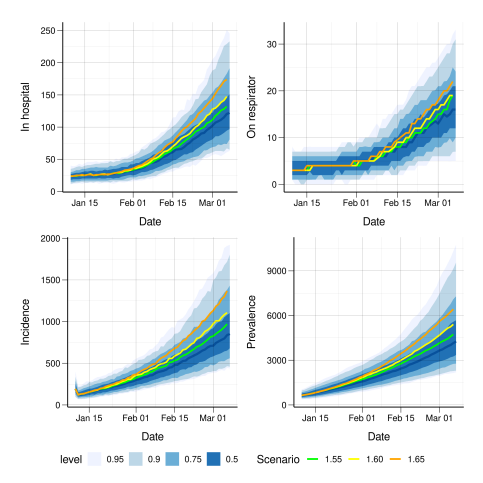


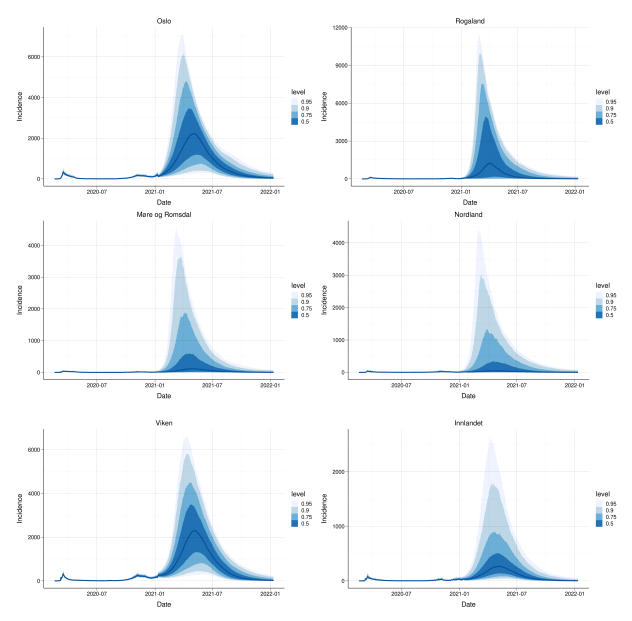
Figure 20: Future predictions for Oslo assuming the reproductive number will remain constant vs alternative scenarios. Confidence intervals correspond to "current scenario". Parameters showed are: Hospitalisations (top left), On respirator (top right), Incidence (bottom left) and Prevalence (bottom right).



11 Regional long-term predictions

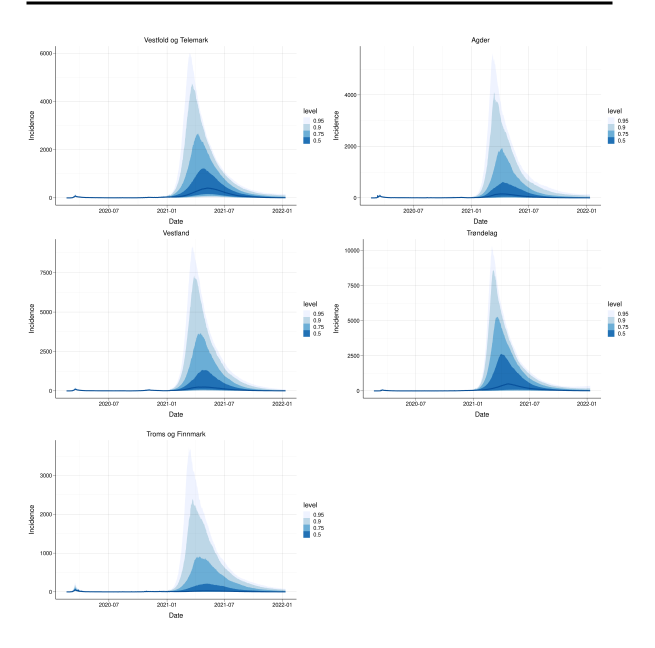
11.1 Incidence

Predicted incidence (asymptomatic, pre-symptomatic and symptomatic) of the calibrated regional changepoint model for each county per day, with confidence intervals.





11.2 Hospitalisations

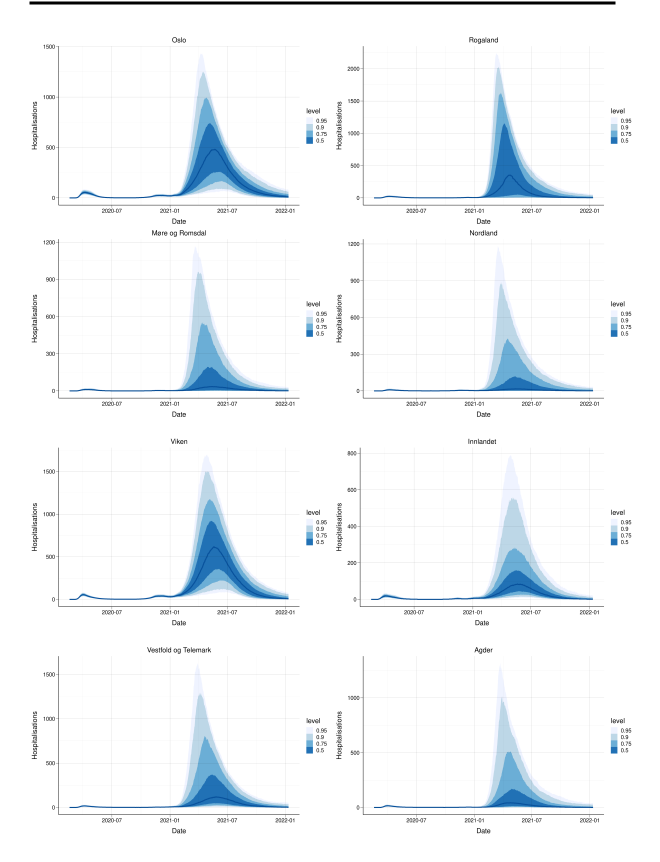


11.2 Hospitalisations

Estimated prevalence of COVID-19 patients in hospital, including patients receiving ventilator treatment.

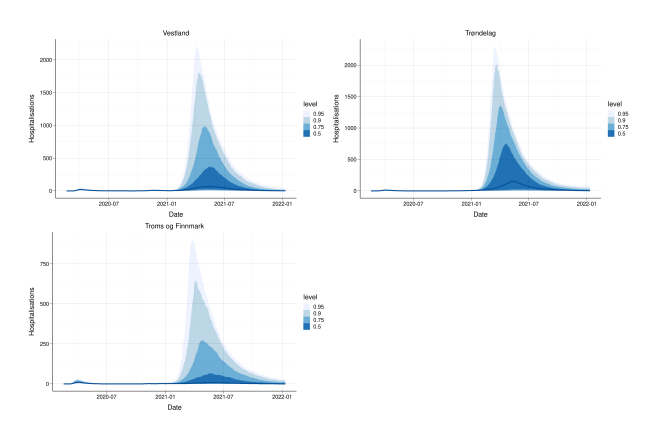


11.2 Hospitalisations





11.2 Hospitalisations





12 Mobility data

Number of trips out from each municipality during each day is based on Telenor mobility data. We observed a large reduction in inter-municipality mobility in March (with minimum reached on Tuesday 17 March), and thereafter we see an increasing trend in the mobility lasting until vacation time in July. The changes in mobility in July coincides with the three-week "fellesferie" in Norway, and during August the mobility resumes approximately the same levels as pre-vacation time. There is however a significant local variation.

The reference level is set to 100 on March 2nd 2020 for all the figures in this section, and we plot the seven-day, moving average of the daily mobility. Figure 21 shows an overview of the mobility since March for the largest municipalities in each county, and Figure 22 shows the total mobility out from all municipalities in each county, including Oslo. Figure 23 and 24, zooms in on mobility from August 31, for municipalities and counties, respectively.

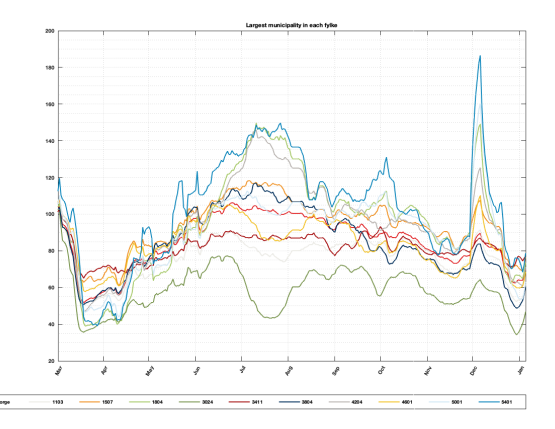


Figure 21: Mobility for selected municipalities for all of 2020: Nationally (Norge), Stavanger (1103), Ålesund (1507), Bodø (1804), Bærum (3024), Ringsaker (3411), Sandefjord (3804), Kristiansand (4204), Bergen (4601), Trondheim (5001), Tromsø (5401).



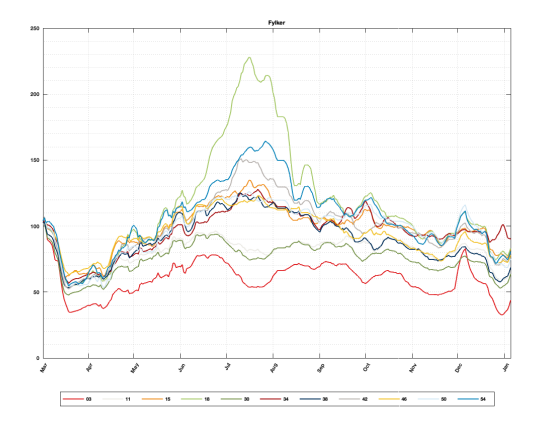


Figure 22: Mobility for fylker for all of 2020: Oslo (03), Rogaland (11), Møre og Romsdal (15), Nordland (18), Viken (30), Innlandet (34), Vestfold og Telemark (38), Agder (42), Vestland (46), Trøndelag (50), Troms og Finmark (54).



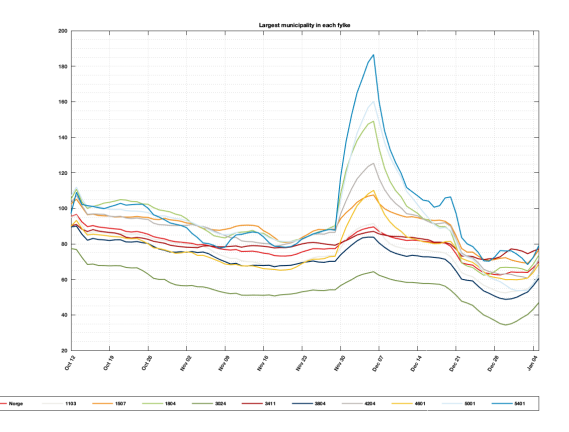


Figure 23: Zoom: Mobility from August 31 and onwards: Nationally (Norge), Stavanger (1103), Ålesund (1507), Bodø (1804), Bærum (3024), Ringsaker (3411), Sandefjord (3804), Kristiansand (4204), Bergen (4601), Trondheim (5001), Tromsø (5401).



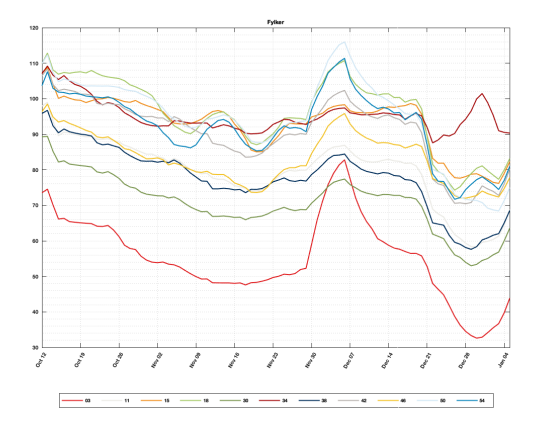


Figure 24: Zoom: Mobility from August 31 and onwards, per fylker: Oslo (03), Rogaland (11), Møre og Romsdal (15), Nordland (18), Viken (30), Innlandet (34), Vestfold og Telemark (38), Agder (42), Vestland (46), Trøndelag (50), Troms og Finnmark (54).



	51	52	53	2	3
Norge	81.9	74.4	62.7	66.5	62.7
Stavanger	77.1	68.5	53.9	57.6	53.5
${ m \AA}{ m lesund}$	94.7	84.5	71.2	72.2	68.3
\mathbf{Bod} ø	97.2	78.7	64.4	68.5	67.5
Bærum	58.0	51.1	36.9	43.1	43.6
Ringsaker	83.1	77.0	71.7	75.9	68.7
Sandefjord	73.2	64.7	50.6	56.4	53.8
Kristiansand	95.8	82.5	63.7	65.0	61.8
Bergen	82.3	77.5	61.1	63.8	62.0
Trondheim	102.0	81.8	59.9	57.3	58.5
Tromsø	107.0	97.0	72.6	72.3	73.1

Table 13: Municipalities

	51	52	53	2	3
Oslo	58.8	52.8	34.5	39.7	40.4
Rogaland	83.0	74.1	59.3	64.6	60.3
Møre og Romsdal	97.7	89.6	78.1	79.0	73.1
Nordland	101.4	89.3	77.2	79.9	77.1
Viken	73.0	66.2	53.9	60.0	57.4
Innlandet	95.9	91.9	94.3	90.5	77.2
Vestfold og Telemark	79.1	69.6	58.1	65.0	61.5
Agder	95.4	84.0	70.5	76.0	73.1
Vestlandet	87.5	82.5	72.0	74.7	70.5
Trøndelag	99.4	87.1	71.7	71.3	68.6
Troms og Finnmark	97.1	87.0	74.4	77.0	74.7

Table 14: Counties

Weekly mobility for Norway and selected municipalities is displayed in Tables 13 and mobility for counties is displayed in 14. The percentages in the tables are to be interpreted towards the reference level of 100% for week 10 in March 2020. The color-coding encodes the following: 'Green' monotonic decrease in mobility, 'Yellow' almost monotonic decrease or flat mobility trend, 'Red' increasing mobility.

12.1 Foreign roamers on Telenor's network in Norway

An analysis of foreign roamers in Norway from January 2020 has been carried out, to better understand the potential virus importation. In Figure 25 the total number of roamers per day per county are displayed. We can see an approximate 40% drop in the number of visiting roamers after the lock-down in March. The number of visiting roamers recover during the Summer, and there is a spike of visitors in August followed by a drop again. During October and November the levels of visiting, foreign roamers to Norway have reached quite high levels, just 10% short of the all-year high for 2020, and Oslo and Viken have seen big increases in visitors. There is a reduction in visitors during Christmas, and in January 2021 we see an increasing trend again.

Figure 26 showcases the levels of roamers from the following countries: Poland, Lithuania, Sweden, Netherlands, Denmark, Latvia, Germany, Spain, Finland and the rest of the world. These levels represent the total number of foreign, visiting roamers from each of the countries per day in Norway, since October 2020.

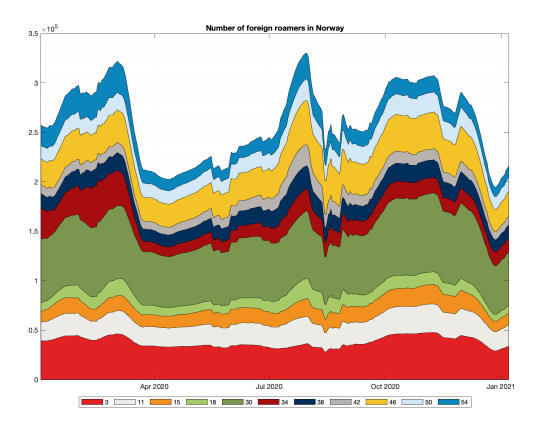


Figure 25: The total number of foreign roamers in Norway broken down on different fylker: Oslo (3), Rogaland (11), Møre og Romsdal (15), Nordland (18), Viken (30), Innlandet (34), Vestfold og Telemark (38), Agder (42), Vestland (46), Trøndelag (50), Troms og Finmark (54).



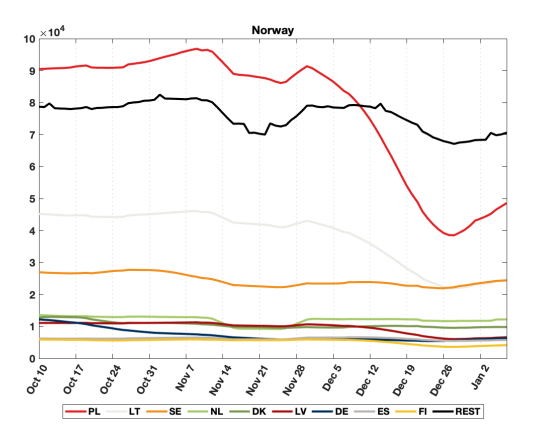
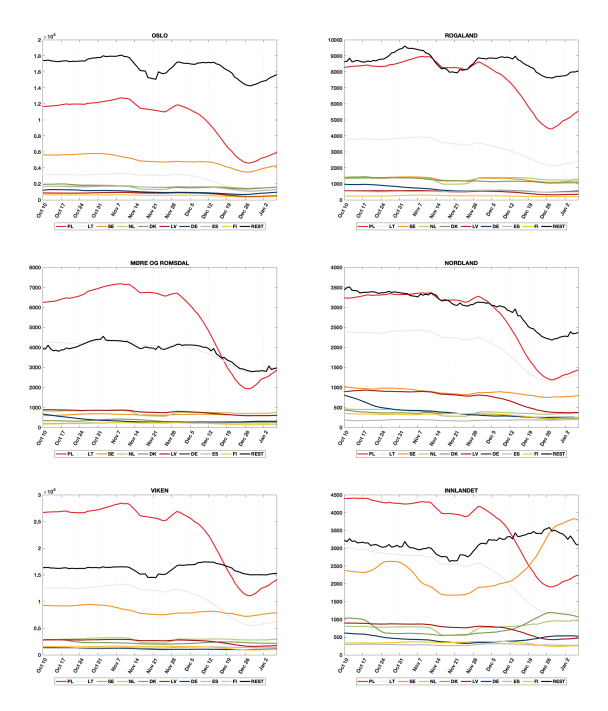


Figure 26: National overview of total number of foreign, visiting roamers from Poland, Lithuania, Sweden, Netherlands, Denmark, Latvia, Germany, Spain, Finland and the rest of the world.

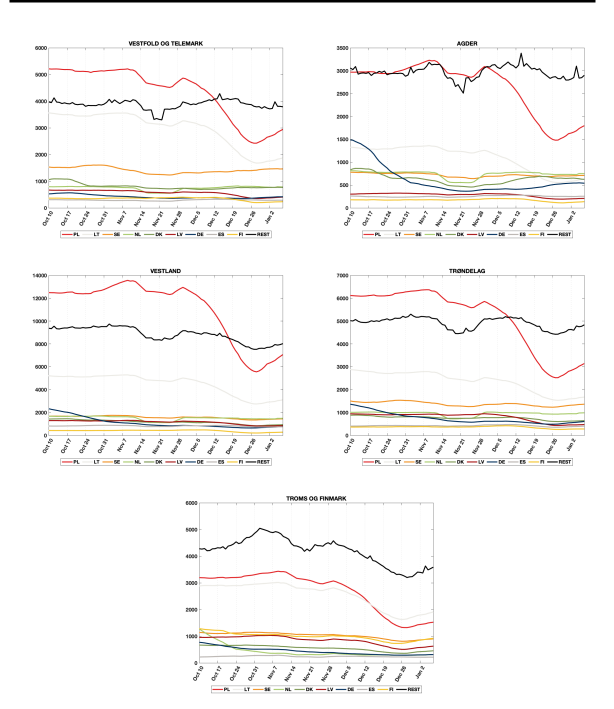


12.2 Foreign roamers per county (fylke) in Norway





12.2 Foreign roamers per county (fylke) in Norway





13 Methods

13.1 Model

We use a metapopulation model to simulate the spread of COVID-19 in Norway in space and time. The model consists of three layers: the population structure in each municipality, information about how people move between different municipalities, and local transmission within each municipality. In this way, the model can simulate the spread of COVID-19 within each municipality, and how the virus is transported around in Norway.

13.1.1 Transmission model

We use an SEIR (Susceptible-Exposed-Infected-Recovered) model without age structure to simulate the local transmission within each area. Mixing between individuals within each area is assumed to be random. Demographic changes due to births, immigration, emigration and deaths are not considered. The model distinguishes between asymptomatic and symptomatic infection, and we consider presymptomatic infectiousness among those who develop symptomatic infection. In total, the model consists of 6 disease states: Susceptible (S), Exposed, infected, but not infectious (E₁), Presymptomatic infected (E₂), Symptomatic infected (I), Asymptomatic infected (I_a), and Recovered, either immune or dead (R). A schematic overview of the model is shown in figure 29.

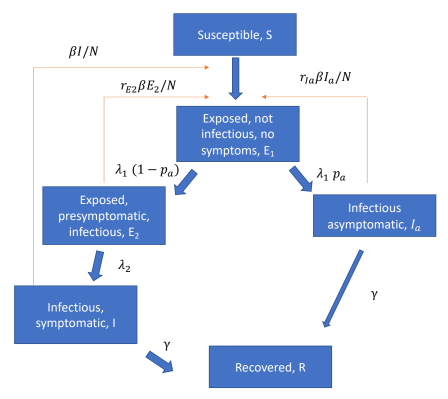


Figure 29: Schematic overview of the model.

13.2 Movements between municipalities:

We use 6-hourly mobility matrices from Telenor to capture the movements between municipalities. The matrices are scaled according to the overall Telenor market share in Norway, estimated to be 48%. Since week 8, we use the actual daily mobility matrices to simulate the past. In this way, alterations in the mobility pattern will be incorporated in our model predictions. To predict future movements, we use the



latest weekday measured by Telenor, regularised to be balanced in total in- and outgoing flow for each municipality.

13.3 Healthcare utilisation

Based on the estimated daily incidence data from the model and the population age structure in each municipality, we calculated the hospitalisation using a weighted average. We correct these probabilities by a factor which represents the over or under representation of each age group among the lab confirmed positive cases. The hospitalisation is assumed to be delayed relative to the symptom onset. We calculate the number of patients admitted to ventilator treatment from the patients in hospital using age-adjusted probabilities and an assumed delay.

13.4 Seeding

At the start of each simulation, we locate 5.367.580 people in the municipalities of Norway according to data from SSB per January 1. 2020. All confirmed Norwegian imported cases with information about residence municipality and test dates are used to seed the model, using the data available until yesterday. For each case, we add an additional random number of infectious individuals, in the same area and on the same day, to account for asymptomatic imported cases who were not tested or otherwise missed. We denote this by the amplification factor.

13.5 Calibration

Estimation of the parameters of the model: the reproduction numbers, the amplification factor for the imported cases, the parameters of the detection probability and the delay between incidence and test, is done using Sequential Monte Carlo Approximate Bayesian Computation (SMC-ABC), as described in Engebretsen et al. (2020): https://royalsocietypublishing.org/doi/10.1098/rsif.2019.0809, where the algorithm can be found in the supplement.

The idea behind ABC is to try out different parameter sets, simulate using these, then compare how much the simulations deviate from the observations in terms of summary statistics. We thus test millions of combinations of $R_0, R_1, R_2, R_3, R_4, R_5, R_6, R_7, R_8, R_9, R_{10}, R_{11}$, the amplification factor, and the parameters for the positive tests, to determine the ones that lead to the best fits to the true number of hospitalised individuals, from March 10 until the last available data point, and the laboratory-confirmed COVID-19 cases from May 1 until the latest available data point.

In the ABC procedure we thus use two summary statistics, one is the distance between the simulated hospitalisation incidence and the observed incidence, and the other is the distance between the observed number of laboratory-confirmed cases and the simulated ones. As the two summary statistics are not on the same scale, we use two separate tolerances in the ABC-procedure, ensuring that we get a good fit to both data sources.

13.5.1 Calibration to hospitalisation data

In order to calibrate to the hospitalisation data, we need to simulate hospital incidence. The details on how we simulate hospitalisations are described in Section 13.3, using the parameters provided in Section 14, which are estimated from individual-level Norwegian data, and updated regularly. As our distance measure, we calculate the squared distance over each time point and each county.

13.5.2 Calibration to test data

We include the laboratory-confirmed cases in the calibration procedure, as these contain additional information about the transmissibility, and the delay between transmission and testing is shorter than the delay between transmission and hospitalisation. Therefore, we simulate also the number of detected positive cases in our model. We assume that the number of detected positive cases can be modelled as a binomial process of the simulated daily total incidence of symptomatic and asymptomatic cases, with a



success probability π_t , which changes every day. We also assume a delay d between the day of test and the day of transmission.

The data on the number of positive cases are more difficult to use, as the test criteria and capacity have changed multiple times. We take into account these changes by using the total number of tests performed on each day, as a good proxy of capacity and testing criteria. Moreover, we choose not to calibrate to the test data before May 1, because the test criteria and capacity were so different in the early period. The detection probability is modelled as

$$\pi_t = \exp(\pi_0 + \pi_1 \cdot k_t) / (1 + \exp(\pi_0 + \pi_1 \cdot k_t)),$$

where k_t is the number of tests actually performed on day t, and π_0 and π_1 are two parameters that we estimate, assuming positivity of π_1 . We also estimate the delay d. We choose to use a 7-days backwards moving average for the covariate k_t . To calculate the distance between the observed number of positive tests and the simulated ones we also use a 7-days backwards moving average. We do this to take into account potential day-of-the-week-effects. For example, it could well be that the testing criteria are different on weekends and weekdays. However, using instead the number of tests and calibrating on a daily basis would lead to a larger day-to-day variance. This is likely why we find that the uncertainty in the simulated positive cases seems somewhat too low, and that we do not capture all the variance in the daily test data. Moreover, the binomial assumption could be too simple, and a beta-binomial distribution would allow more variance. A limitation of our current model for the detection probability, is that we only capture the changes in the test criteria that are captured in the total number of tests performed.

13.6 Specifications for the national changepoint model

In the national changepoint model, we assume a first reproduction number R_0 until March 14, a second reproduction number R_1 until April 19, a third reproduction number R_2 until May 10, a fourth reproduction number R_3 until June 30, R_4 until July 31, R_5 until August 31, R_6 from September 1 until September 30, R_7 from October 1 until October 26, R_8 until November 4, R_9 from November 5th until November 30th, R_{10} from December 1st until December 20th and a twelfth reproduction number R_{11} from December 21st. This last reproduction number is used for the future. The changepoints follow the changes in restrictions introduced. In the calibration procedure, we obtain 200 parameter sets that we use to represent the distributions of parameters.

After we have obtained the estimated parameters, we run the model with these 200 parameter sets again, from the beginning until today, plus three weeks into the future (or for an additional year). In this way, we obtain different trajectories of the future, allowing us to investigate different scenarios, with corresponding uncertainty.

13.7 Specifications for the regional changepoint model

In the regional changepoint model, each county has its own reproduction numbers, assumed constant in different periods, just like the national changepoint model. As there are more parameters in the regional changepoint model, we obtain 1000 parameter sets in the ABC-SMC.

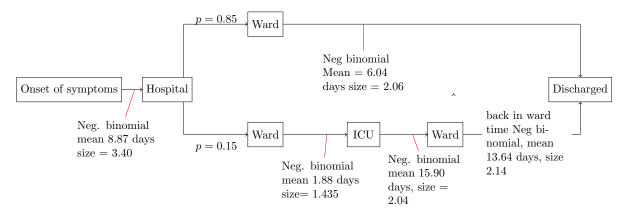
Calibrating regional reproduction numbers is a more difficult estimation problem than calibrating national reproduction numbers, as we have a lot more parameters, and in addition less data in each county. Therefore, we cannot include the same amount of changepoints in the regional model as we can for the national model. Currently we assume five changepoints (six reproduction numbers) for Viken, Oslo, Vestland (the three largest counties of Norway), and four changepoints (five reproduction numbers) in all other counties.

After we have obtained the estimated parameters, we run the model with these 1000 parameter sets again, from the beginning until today, plus three weeks into the future (or for an additional year). In this way, we obtain different trajectories of the future, allowing us to investigate different scenarios, with corresponding uncertainty.



14 Parameters used today

Figures 30 and 31 indicate which assumptions we make in our model, related to hospitalisation. We obtained data from the Norwegian Pandemiregister. These estimates will be regularly updated, on the basis of new data.





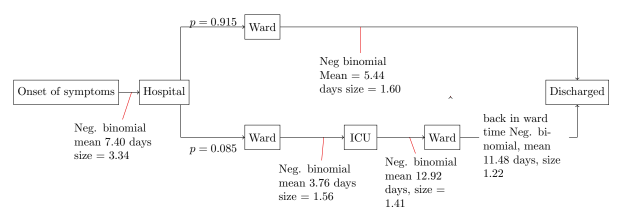


Figure 31: Hospital assumptions and parameters used after 1 August



	Min.	1st Qu.	Median	Mean	3rd Qu.	Max.	Period
R0s	1.944	2.798	3.115	3.122	3.421	4.071	Until March 14
R1s	0.354	0.473	0.517	0.516	0.562	0.646	From March 15 to April 19
R2s	0.024	0.468	0.655	0.662	0.861	1.429	From April 20 to May 10
R3s	0.011	0.374	0.581	0.578	0.767	1.344	From May 11 to June 30
R4s	0.048	0.481	0.768	0.749	1.027	1.756	From July 1 to July 31
R5s	0.535	0.847	1.006	1.009	1.148	1.519	From Aug 1 to Aug 31
R6s	0.598	0.863	0.974	0.975	1.079	1.318	From Sept 1 to Sept 30
m R7s	0.869	1.151	1.236	1.241	1.341	1.571	From Oct 1 to Oct 25
$\mathbf{R8s}$	1.016	1.322	1.463	1.469	1.607	2.024	From Oct 26 to Nov 4
R9s	0.715	0.816	0.842	0.841	0.868	0.923	From Nov 5 to Dec 1
R10s	0.948	1.028	1.058	1.059	1.09	1.233	From Dec 1 to Dec 26
R11s	0.817	1.117	1.241	1.247	1.37	1.969	From Dec 27
AMPs	1.001	1.507	1.77	1.821	2.093	3.087	-
π_0	-1.304	-0.333	0.022	-0.065	0.208	0.767	-
π_1	6.6e-07	2.2e-05	3.8e-05	4.5e-05	6.7 e-05	1.4e-04	-
delays	0	1	2	2.1	3	4	-

Table 15: Estimated parameters

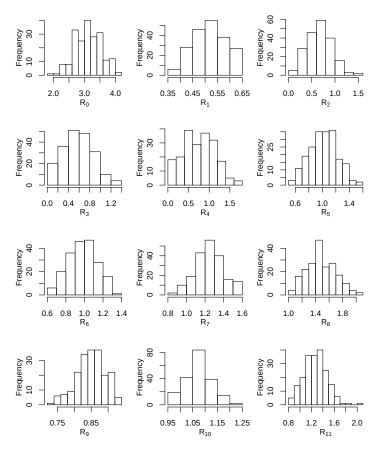


Figure 32: Estimated densities of the reproduction numbers. National model



Table 16

R	Parameter	County	From	То	$\Pr(R>1)$
4.61 (3.81-5.54)	R0	Oslo	2020-02-17	2020-03-14	1
3.13(2.04-4.22)	R0	Rogaland	2020-02-17	2020-03-14	1
3.45(1.72-4.96)	R0	Møre og Romsdal	2020-02-17	2020-03-14	1
3.33 (1.4-5.32)	R0	Nordland	2020-02-17	2020-03-14	1
4(2.87-4.95)	R0	Viken	2020-02-17	2020-03-14	1
3.3(2.03-4.55)	R0	Innlandet	2020-02-17	2020-03-14	1
3.26 (2.11-4.48)	R0	Vestfold og Telemark	2020-02-17	2020-03-14	1
2.79(1.61-3.92)	R0	Agder	2020-02-17	2020-03-14	1
3.68 (2.6-4.74)	R0	Vestland	2020-02-17	2020-03-14	1
4.16(2.68-5.75)	R0	Trøndelag	2020-02-17	2020-03-14	1
3.5(2.67-4.51)	R0	Troms og Finnmark	2020-02-17	2020-03-14	1
0.86(0.73-1.02)	R1	Öslo	2020-03-15	2020-04-19	0.06
0.46(0.11 - 0.72)	R2	Oslo	2020-04-20	2020-06-19	0
0.54(0.21-0.83)	R3	Oslo	2020-06-20	2020-08-31	0
1.39(1.27-1.5)	R4	Oslo	2020-09-01	2020-11-04	1
1.01(0.89-1.17)	R5	Oslo	2020-11-05	2020-11-30	0.49
1.04(0.76-1.31)	R6	Oslo	2020-12-01	2020-12-20	0.64
1.48 (1.05-1.96)	R7	Oslo	2020-12-21		0.98
0.77(0.52 - 1.02)	R1	Rogaland	2020-03-15	2020-04-19	0.04
0.82(0.56-1.01)	R2	Rogaland	2020-04-20	2020-08-31	0.03
0.68(0.48 - 0.92)	R3	Rogaland	2020-09-01	2020-11-04	0.01
1.31 (0.87 - 1.58)	R4	Rogaland	2020-11-05	2020-11-30	0.94
0.56(0.1-1.08)	R5	Rogaland	2020-12-01	2020-12-20	0.05
1.43(0.4-2.38)	R6	Rogaland	2020-12-21		0.79
0.89(0.56-1.21)	R1	Møre og Romsdal	2020-03-15	2020-04-19	0.26
0.61(0.31-0.91)	R2	Møre og Romsdal	2020-04-20	2020-08-31	0
1.13 (0.7 - 1.46)	R3	Møre og Romsdal	2020-09-01	2020-11-04	0.78
0.87(0.4-1.28)	R4	Møre og Romsdal	2020-11-05	2020-11-30	0.26
0.65 (0.07 - 1.42)	R5	Møre og Romsdal	2020-12-01	2020-12-20	0.19
0.99(0.11-2.16)	R6	Møre og Romsdal	2020-12-21		0.46
0.62(0.3-0.95)	R1	Nordland	2020-03-15	2020-04-19	0.01
0.9(0.68-1.13)	R2	Nordland	2020-04-20	2020-08-31	0.19
1.17 (0.91 - 1.42)	R3	Nordland	2020-09-01	2020-11-04	0.91
0.89(0.42 - 1.35)	R4	Nordland	2020-11-05	2020-11-30	0.32
0.58(0.04-1.3)	R5	Nordland	2020-12-01	2020-12-20	0.12
0.93 (0.08-2.12)	R6	Nordland	2020-12-21		0.42
0.37(0.18 - 0.53)	R1	Viken	2020-03-15	2020-04-19	0
0.86(0.59-1.06)	R2	Viken	2020-04-20	2020-06-19	0.11
0.77 (0.31-1.09)	R3	Viken	2020-06-20	2020-08-31	0.13
1.4(1.28-1.53)	R4	Viken	2020-09-01	2020-11-04	1
1 (0.87 - 1.09)	R5	Viken	2020-11-05	2020-11-30	0.5
0.82(0.63-0.99)	R6	Viken	2020-12-01	2020-12-20	0.02
1.05 (0.77-1.36)	R7	Viken	2020-12-21		0.62

Mean and 95% credible intervals

Table 17

R	Parameter	County	From	То	$\Pr(R>1)$
0.69(0.45 - 1.03)	R1	Innlandet	2020-03-15	2020-04-19	0.03
0.72 (0.57-0.93)	R2	Innlandet	2020-04-20	2020-08-31	0.01
0.93 (0.57-1.26)	R3	Innlandet	2020-09-01	2020-11-04	0.36
0.2(0.08-0.29)	R4	Innlandet	2020-11-05	2020-11-30	0
0.8(0.07 - 1.55)	R5	Innlandet	2020-12-01	2020-12-20	0.34
0.69(0.04-1.59)	R6	Innlandet	2020-12-21		0.23
0.61 (0.26-0.88)	R1	Vestfold og Telemark	2020-03-15	2020-04-19	0
0.86(0.72 - 1.07)	R2	Vestfold og Telemark	2020-04-20	2020-08-31	0.08
0.66 (0.37-0.97)	R3	Vestfold og Telemark	2020-09-01	2020-11-04	0
0.54 (0.12-0.98)	R4	Vestfold og Telemark	2020-11-05	2020-11-30	0.02
0.9(0.14 - 1.57)	R5	Vestfold og Telemark	2020-12-01	2020-12-20	0.42
0.99(0.13-2)	R6	Vestfold og Telemark	2020-12-21		0.46
0.55(0.24-0.87)	R1	Agder	2020-03-15	2020-04-19	0
0.89 (0.69-1.06)	R2	Agder	2020-04-20	2020-08-31	0.12
0.87(0.55-1.23)	R3	Agder	2020-09-01	2020-11-04	0.23
1.02 (0.44-1.48)	R4	Agder	2020-11-05	2020-11-30	0.56
0.46 (0.03-1.06)	R5	Agder	2020-12-01	2020-12-20	0.04
0.9(0.07-2.1)	R6	Agder	2020-12-21		0.4
0.58(0.35-0.83)	R1	Vestland	2020-03-15	2020-04-19	0
0.83 (0.6-1.05)	R2	Vestland	2020-04-20	2020-08-16	0.06
0.52(0.06-1.39)	R3	Vestland	2020-08-17	2020-09-09	0.11
1.26 (1.03-1.49)	R4	Vestland	2020-09-10	2020-11-04	0.98
0.73(0.43-1.11)	R5	Vestland	2020-11-05	2020-11-30	0.07
0.46(0.02 - 1.03)	R6	Vestland	2020-12-01	2020-12-20	0.04
0.92(0.1-1.88)	R7	Vestland	2020-12-21		0.41
0.56(0.28 - 0.84)	R1	Trøndelag	2020-03-15	2020-04-19	0
0.29 (0.03-0.68)	R2	Trøndelag	2020-04-20	2020-08-31	0
0.57(0.15-1.06)	R3	Trøndelag	2020-09-01	2020-11-04	0.05
1.27(0.66-1.7)	R4	Trøndelag	2020-11-05	2020-11-30	0.84
0.98(0.29-1.59)	R5	Trøndelag	2020-12-01	2020-12-20	0.48
1.21 (0.29-2.23)	R6	Trøndelag	2020-12-21		0.64
0.48(0.2-0.78)	R1	Troms og Finnmark	2020-03-15	2020-04-19	0
0.82(0.62-1)	R2	Troms og Finnmark	2020-04-20	2020-08-31	0.03
0.83(0.42 - 1.23)	R3	Troms og Finnmark	2020-09-01	2020-11-04	0.18
0.82(0.31 - 1.34)	R4	Troms og Finnmark	2020-11-05	2020-11-30	0.26
0.67(0.05 - 1.58)	R5	Troms og Finnmark	2020-12-01	2020-12-20	0.2
0.86(0.05-1.95)	R6	Troms og Finnmark	2020-12-21		0.38
1.23(1.01-1.58)	AMP factor	All			-

1.23 (1.01-1.58) AMP factor Mean and 95% credible intervals



Table 18:	Hospitalisation	probabilities	(1/2)
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	Until 2020-05-01	Until 2020-06-01	Until 2020-07-01	Until 2020-08-01
0-9 years	0.0002	0.0005	0.001	0.0005
10 - 19 years	0.001	0.001	0.001	0.001
20 - 29 years	0.006	0.007	0.009	0.010
30 - 39 years	0.014	0.019	0.016	0.014
40 - 49 years	0.018	0.016	0.015	0.013
50 - 59 years	0.040	0.032	0.025	0.030
60 - 69 years	0.053	0.030	0.031	0.035
70 - 79 years	0.075	0.048	0.022	0.041
80 + years	0.326	0.120	0.145	0.040

Table 19: Hospitalisation probabilities (2/2)

	Until 2020-09-01	Until 2020-10-01	Until 2020-11-01	Until 2020-12-01	From 2020-12-01
0-9 years	0.0004	0.0004	0.0004	0.001	0.001
10 - 19 years	0.001	0.001	0.001	0.002	0.001
20 - 29 years	0.015	0.012	0.011	0.007	0.007
30 - 39 years	0.011	0.014	0.014	0.013	0.013
40 - 49 years	0.011	0.013	0.015	0.016	0.016
50 - 59 years	0.022	0.024	0.027	0.027	0.030
60 - 69 years	0.015	0.030	0.031	0.036	0.038
70 - 79 years	0.025	0.033	0.033	0.039	0.049
80 + vears	0.038	0.096	0.076	0.123	0.143



Table 20: Assumptions

Assumptions	Mean	Distribution	Reference
Mobile Mobility Data		Distribution	Telefence
Telenor coverage	48%	1	https://ekomstatistikken.nkom.no/
Data updated	January 9th		https://ekonistatistikken.ikoin.iio/
Data updated Data used in the predictions		Fixed	
Model parameters	January 8th	Fixed	Corrected to preserve population
Exposed period $(1/\lambda_1)$	3 days	Exponential	Feretti et al 2020
Pre-symptomatic period $(1/\lambda_2)$	2 days	Exponential	Feretti et al 2020
Symptomatic infectious period $(1/\gamma)$	5 days	Exponential	Feretti et al 2020
Asymptomatic, infectious period $(1/\gamma)$	5 days	Exponential	Feretti et al 2020
Infectiousness asympt. (r_{I_a})	0.1	Fixed	Feretti et al 2020
Infectiousness presymp (r_{E_2})	1.25	Fixed	guided by Feretti et al 2020
Prob. asymptomatic infection (p_a)	0.4		Feretti et al 2020
Healthcare			
Time sympt. onset to hospitalisation	8.87 days (before August 1st) / 7.40 (After August 1st)	Neg. binomial	
D di la di C di	40%	E' 1	Mizumoto et al 2020
Fraction asymptomatic infections	40%	Fixed	20% for the old population, Diamond Princess
% symptomatic and asymptomatic		1	Saljie et al 2020
infections requiring hospitalization:			corrected for: % of elderly living in
0-9 years	0.1%		elderly homes in Norway (last two age groups)
10 - 19 years	0.1%		and corrected for presence among positive tested since May 1
20 - 29 years	0.5%		Corrected values available in tables 18 and 19
30 - 39 years	1.1%	Fixed	
40 - 49 years	1.4%		
50 - 59 years	2.9%		
60 - 69 years	5.8%		
70 - 79 years	9.3%		
80+ years	22.3%		
% hospitalized patients requiring			
1CU			
Feb - July	15.1%	Fixed	Estimated from "Beredskapsregistret BeredtC19"
August -	8.5%	1 Incu	Estimated from Dereasingsregioner Dereatory
Tugase	0.070		
Probability that an admission has been reported on Monday			
From Sunday	32%		
From Saturday	49%	Fixed	Estimated from "Beredskapsregistret BeredtC19"
From Friday	68%	1 IXCO	Estimated from Deredskapsregistret Deredters
From Thursday	86%		
Probability that an admission has been reported	0070		
From one day before	53%		
From two days before	77%	Fixed	Estimated from "Beredskapsregistret BeredtC19"
From three days before	82%	Fixed	Estimated from Beredskapsregistret Beredic 19
From four days before	91%		
	9170		
Probability that a positive laboratory test has been reported	6 797		
From one day before	6.7%	Fixed	Estimated from MSIS
From two days before	59%	F ixed	Estimated from MS15
From three days before	90%		
From four days before	97%		
Probability that a negative laboratory test has been reported	1.007		
From one day before	16%	E: 1	E.C. MOR
From two days before	74%	Fixed	Estimated from MSIS
From three days before	92%		
From four days before	98%		



Supplementary analysis: EpiEstim estimation of reproduction number based on laboratory-confirmed cases

To complement the results of the metapopulation model, we present estimates of the temporal evolution of the reproduction number in Norway based on an analysis of laboratory-confirmed cases. The primary purpose of this analysis is to provide a more comprehensive perspective on the epidemic situation, taking into account several data sources.

The hospitalisation data are a less biased information source for the number of infections compared to case data because the testing criteria in Norway has changed. For this reason, the present results should be interpreted with caution. During the early part of the period, testing of individuals was mainly based on travel history to areas with an ongoing outbreak. Since the middle of March, testing is recommended for people with an acute respiratory infection. From early May, the testing criteria have been expanded to include less severe symptoms. The analysis of laboratory-confirmed cases does not take into account the effect of imported cases during the early outbreak in Norway; the early results are less reliable than later results when the impact of importations is negligible.

EpiEstim method and assumptions: We estimate the instantaneous reproduction number using the procedure outlined in Thompson et al. (2019). This method, implemented in the EpiEstim R-package, uses a Bayesian approach to estimate the instantaneous reproduction number smoothed over a sliding window of 5 days, see figure 33. For the results to be comparable to those of the metapopulation model, we use the same natural history parameters. We estimate the date of infection for each confirmed case by first estimating the date of symptom onset and then subtracting 5 days for the incubation period. We estimate the date of symptom onset from the empirical delay between onset and testing in the first reported cases. For each case, we draw 100 possible onset dates from the delay distribution; this gives us 100 epi-curves that we use to estimate the reproduction number. The displayed results are the combined results from all these 100 simulated epi-curves. The serial interval was assumed to be 5 days with uncertainty; the serial interval refers to the time between symptom onset between successive cases in a chain of transmission (see https://www.medrxiv.org/content/10.1101/2020.02.03.20019497v2). To account for censoring of observations with onset dates in the last few days we correct the observed data by the mean of a negative binomial distribution with observation probability given by the empirical cumulative distribution of the onset to reporting date distributions. Due to this correction, the results from the last few days are uncertain, as indicated by increasing credible intervals.

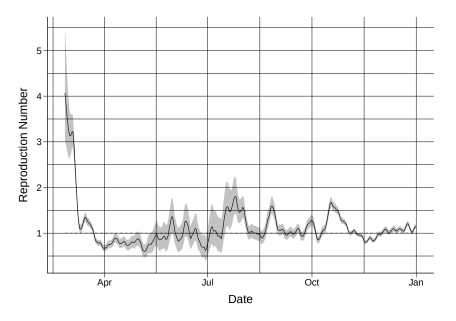


Figure 33: Reproduction number estimated using the R package EpiEstim.



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