

National Wastewater Surveillance Programme - COVID-19

Week 27 2022 (Week Ending 10 July 2022)

Report prepared on 13 July 2022

Overview

SARS-CoV-2, the virus that causes COVID-19 disease, is shed in the faeces of people that are infected and so the viral RNA can be detected in wastewater. As such, testing wastewater for SARS-CoV-2 RNA is an efficient population-based COVID-19 surveillance tool. Based on national and international data, this method has been shown to be an indicator of increasing and decreasing cases (i.e., early warning system) and complements other surveillance tools. A national wastewater COVID-19 surveillance programme was established in 2021 by the Institute of Environmental Science and Research (ESR). This work is funded by the New Zealand Ministry of Health and is part of New Zealand's COVID-19 response.

Wastewater samples are collected from wastewater treatment plants across both the North and South Island of New Zealand. Most sites are sampled at least weekly between Monday and Thursday of any given week. The number of sites and frequency of collection varies over time.

Approach

Samples are sent from each wastewater treatment plant to ESR. Processing involves the concentration of virus and extraction of viral RNA. The presence of SARS-CoV-2 RNA in the sample is then determined using RT-qPCR.

A result of not detected means that SARS-CoV-2 RNA is either absent from the sample, or at a level too low to be detected. When SARS-CoV-2 RNA is detected, the concentration in the sample can be calculated. Low amounts of SARS-CoV-2 RNA in a sample may not be able to be accurately quantified and are recorded as less than the limit of quantitation. For quantitation, the raw concentration data (i.e., genome copies per L) is converted to a viral load of genome copies per day per person. This calculation considers the flow rate of wastewater entering the wastewater treatment plant and the population in the catchment. This is the population-normalised viral load. Currently, sequencing is performed using a S-gene (spike) barcoding assay able to call variants.

Key Points & Limitations

- SARS-CoV-2 RNA concentrations should not be compared between wastewater catchments.
- Day to day variability in SARS-CoV-2 RNA concentrations, especially in smaller catchments, is to be expected. Greater variability is expected with grab samples.
- Generally, increasing viral loads are associated with increasing numbers of people with SARS-CoV-2 infection and vice versa (decreasing concentrations indicating decreasing cases). However, there are a number of factors that affect the amount of viral RNA detected and so data from wastewater surveillance cannot indicate the exact number of COVID-19 cases in the catchment area.
- The number of COVID-19 cases reported via individual testing are reported for each region to provide a comparison to the wastewater results. The cases in each catchment area are an estimate of the number of people in that wastewater catchment area that have reported a positive test. However, because the wastewater catchments do not exactly align with regional boundaries, the number of cases estimated by region and by water catchment area may be different.
- Data are provisional and may be subject to change by location.
- As septic tank systems are not connected to wastewater treatment plants, the wastewater from these households will not be represented in the data.

Results For Week 27, 2022

In the week ending 10 July 2022, 158 samples were collected from 109 locations in New Zealand. Analysis of some samples is in progress. Of the 109 locations sampled, 103 have at least one result available for the week. Whangarei, Cambridge, Te Awamutu, Eketahuna, Richmond and Hokitika are the sites without results (as of 13 July).

SARS CoV-2 RNA was detected at all sites except for a sample collected on 5 July from Kaponga (Taranaki) where SARS-CoV-2 was not detected (Figure 1, Figure 2, Table 1).

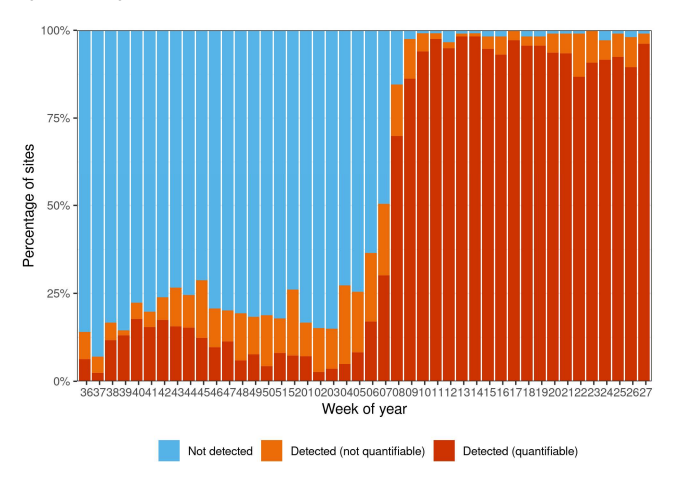


Figure 1 below gives the proportion of sites with positive results for this and previous weeks.

Figure 1. Results for SARS-CoV-2 RNA in wastewater collected across New Zealand.

There are **increases** in SARS-CoV-2 levels in wastewater across almost all sites (Figure 2). Using a threshold of an increase of 30% or more (Figure 2), 77% of sites have an increase compared with the previous week, 84% of sites have increased from levels measured two weeks ago, and 88% of sites have increased compared with one month ago. Compared with levels one month ago, a third of sites have levels of SARS-CoV-2 in wastewater which have increased 5-fold or more. Examples include Gisborne, Queenstown, Mt Maunganui, North Shore (Rosedale), Napier and Paraparaumu (See the section with individual graphs).



Figure 2. Comparison of SARS-CoV-2 levels for the week ending 10 July 2022, compared with the levels measured 4 weeks previously. Only sites with results for both time points are included. When the viral quantity is 30% or more higher this is labelled as increased (red up arrow on map). When the viral quantity is 30% or more lower this is labelled as decreased (blue down arrow on map). If viral levels have changed less than this in the two compared weeks, this is labelled as no change (yellow circle on map).

Regional summaries indicate an **increase** in SARS-CoV-2 levels (genome copies per person per day) across the country, week ending 10 July 2022 (Figure 3, scale is log₁₀).

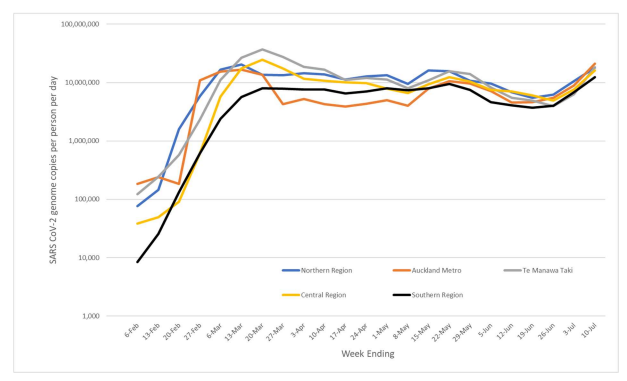
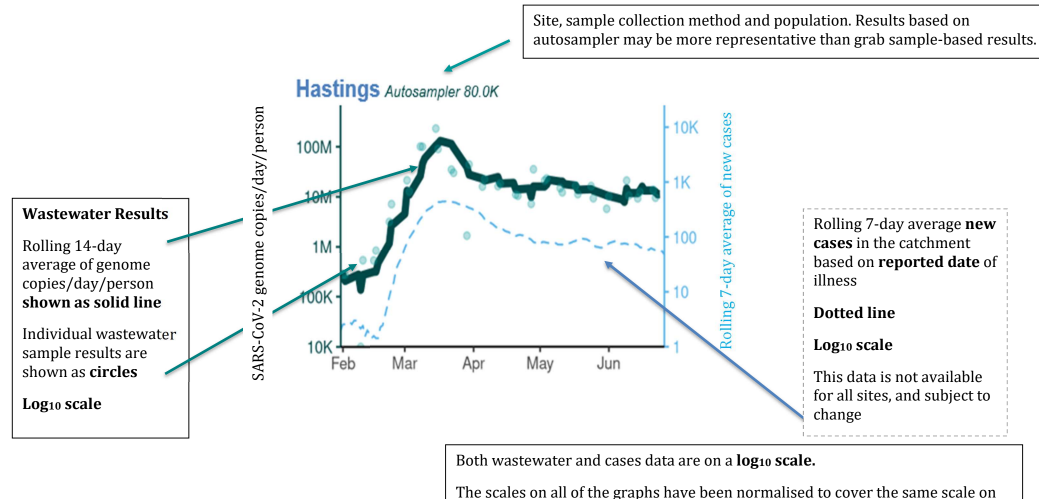


Figure 3. Regional trends in SARS-CoV-2 levels in wastewater based on genome copies per person per day. Northern Region excludes the Auckland metro sites (Eastern, Western, Southwestern and Rosedale- North Shore).

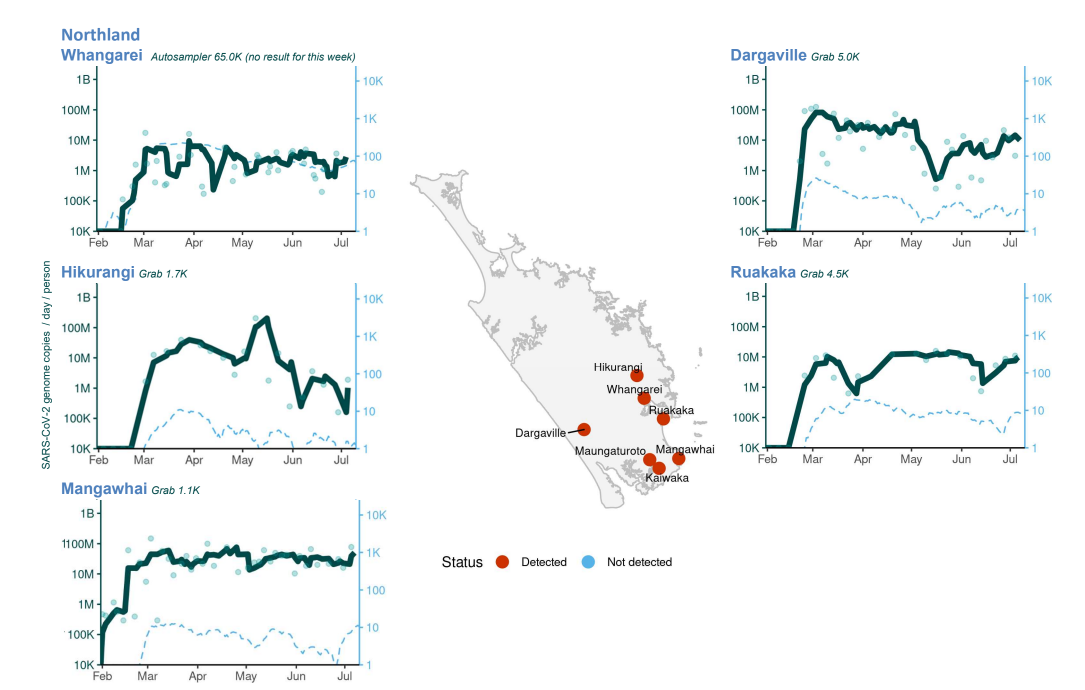
Consistent with results from sequencing of clinical cases, BA.4/5 and BA.2 group (containing BA.2 and BA.2.12.1) were detected in wastewater from all sentinel sites. Overall, these variants were detected in approx equal percentages (50 % BA.4/5 & 50% BA.2 group, but noting that level of precision and sensitivity in the percentage estimates can be uncertain). The BA.2.75 (sub)variant was not detected.

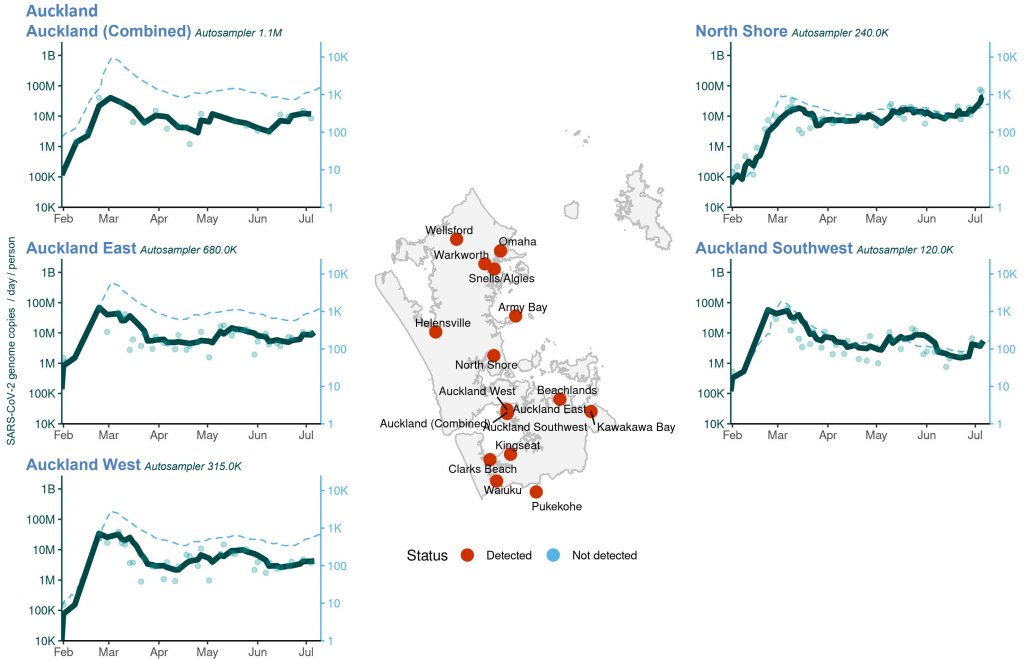
Interpreting site graphs



every graph.

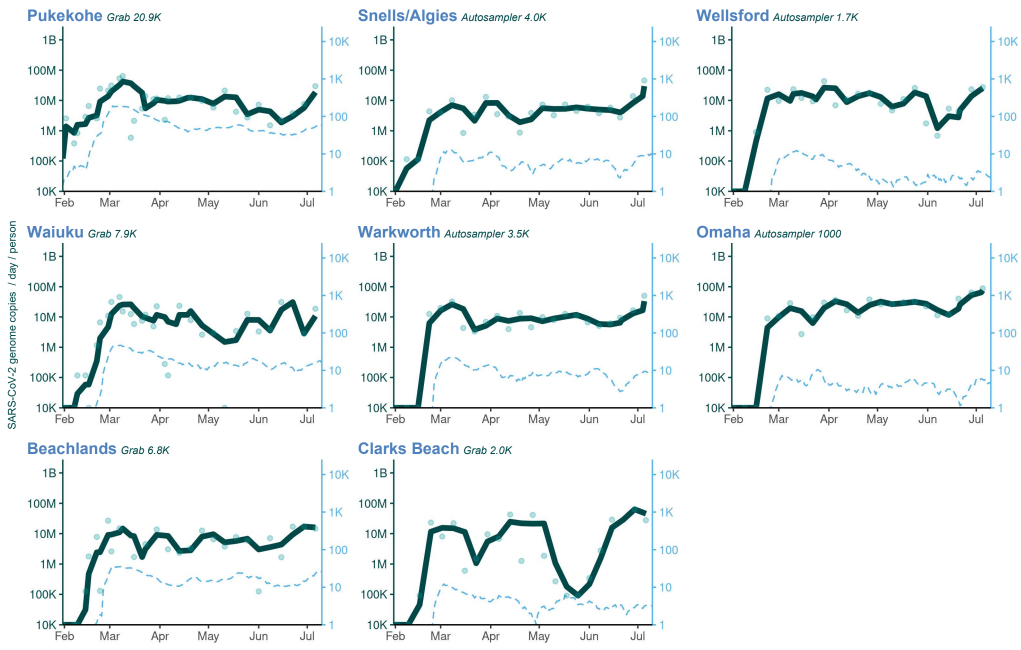
Care should be taken when interpreting the data.



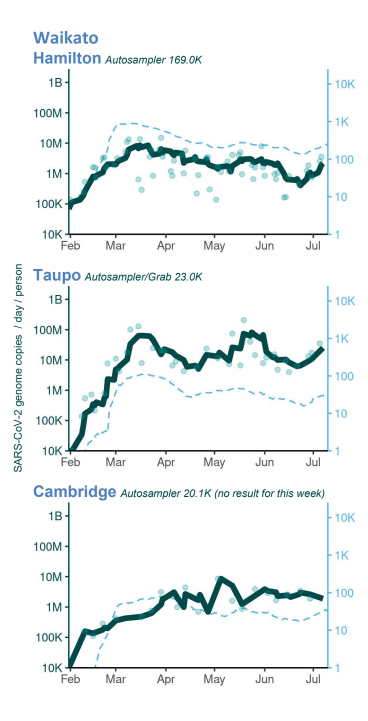


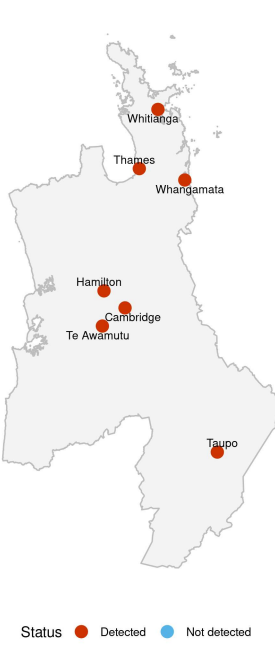
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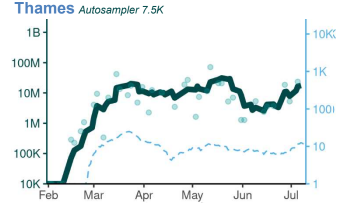
Cases - 7 day rolling average



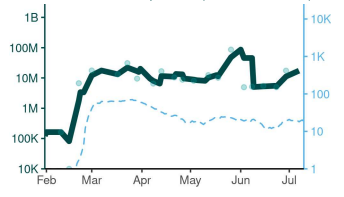
Cases - 7 day rolling average







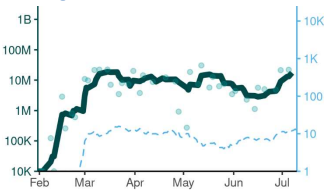
Te Awamutu Autosampler 13.1K (no result for this week)

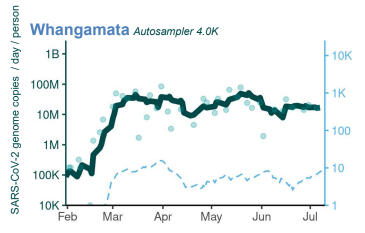


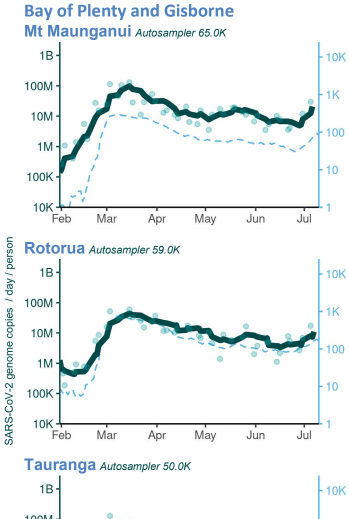
7 day rolling average

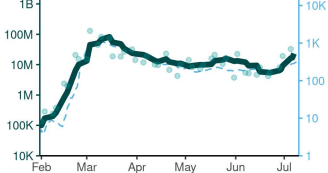
Cases -

Whitianga Autosampler 6.6K





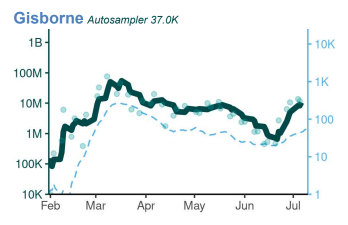




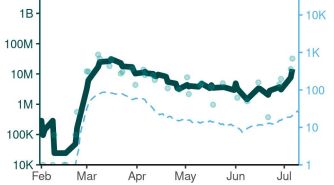


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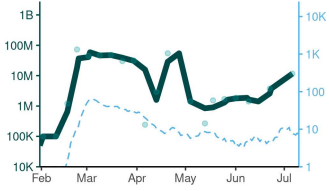
Detected 🔵 Not detected



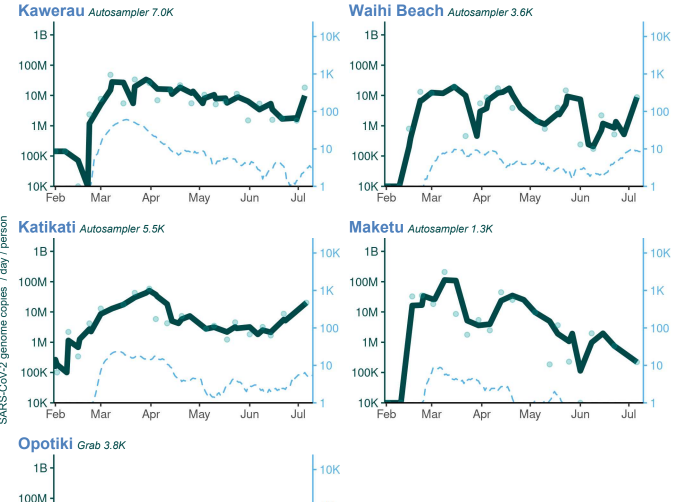




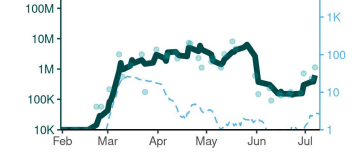


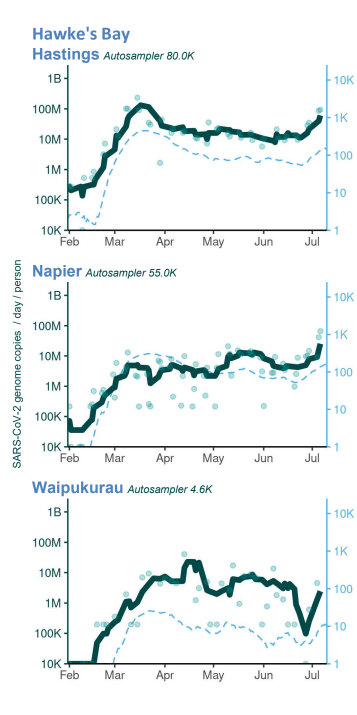


Cases - 7 day rolling average

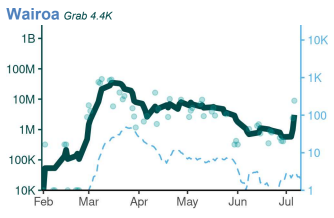




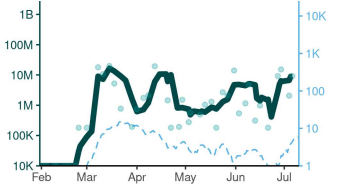




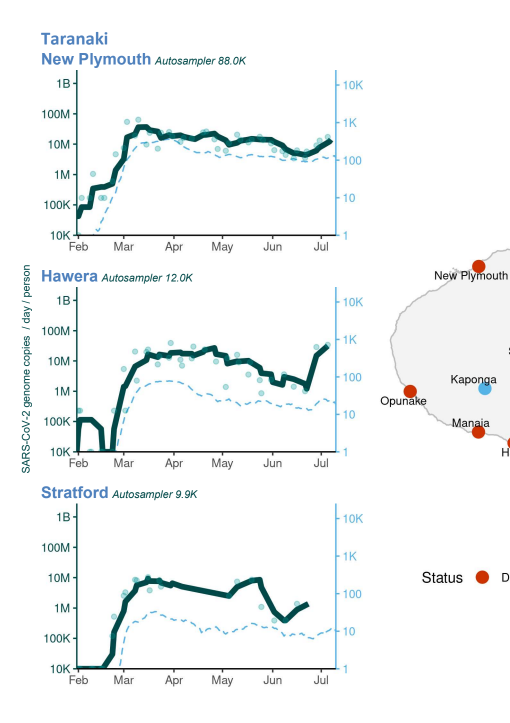


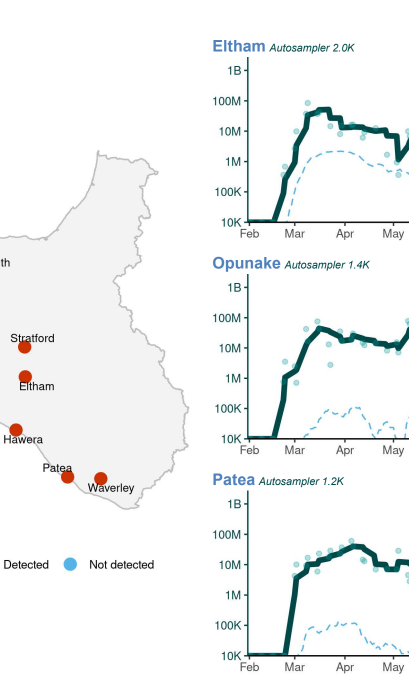


Waipawa Autosampler 2.2K



Cases - 7 day rolling average







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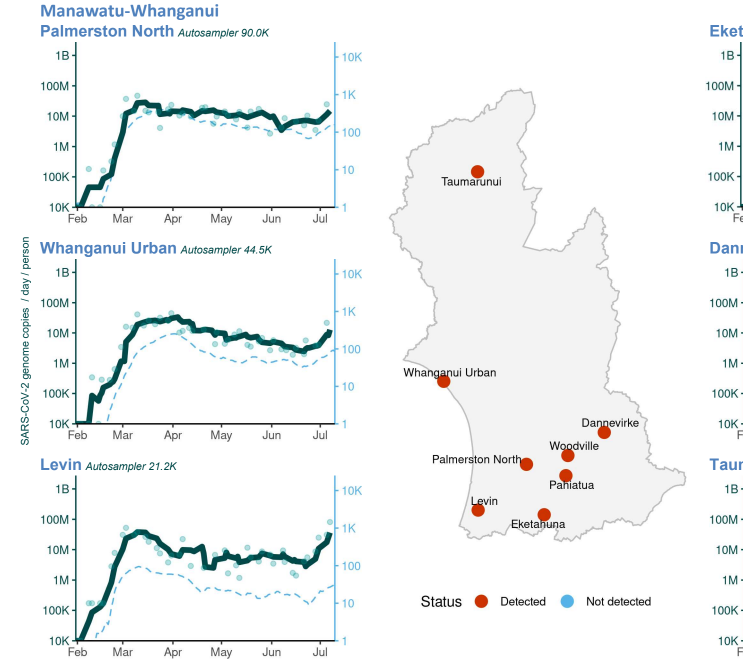
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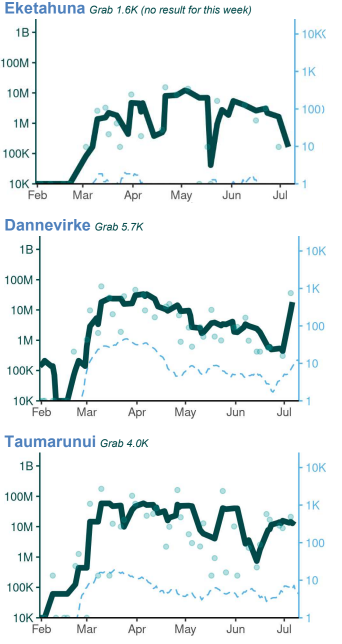
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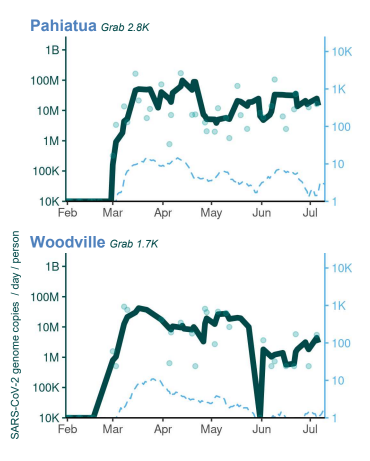
May

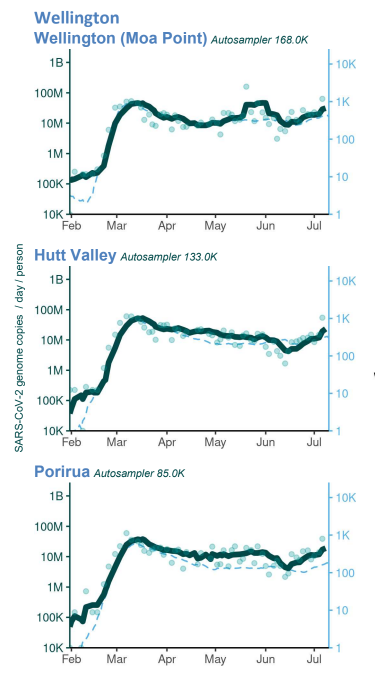




day rolling average

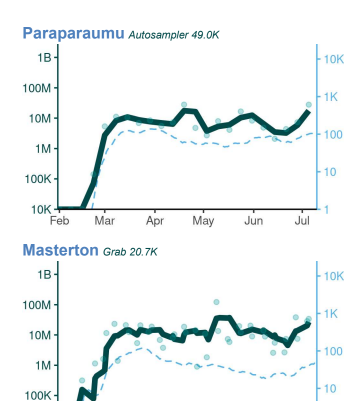
Cases -







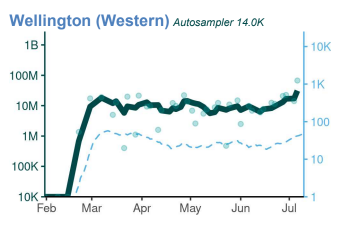
Status 🛑 Detected 🔵 Not detected





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May

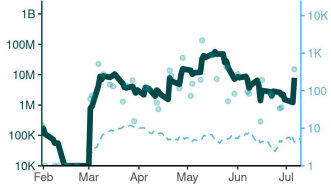
10K -

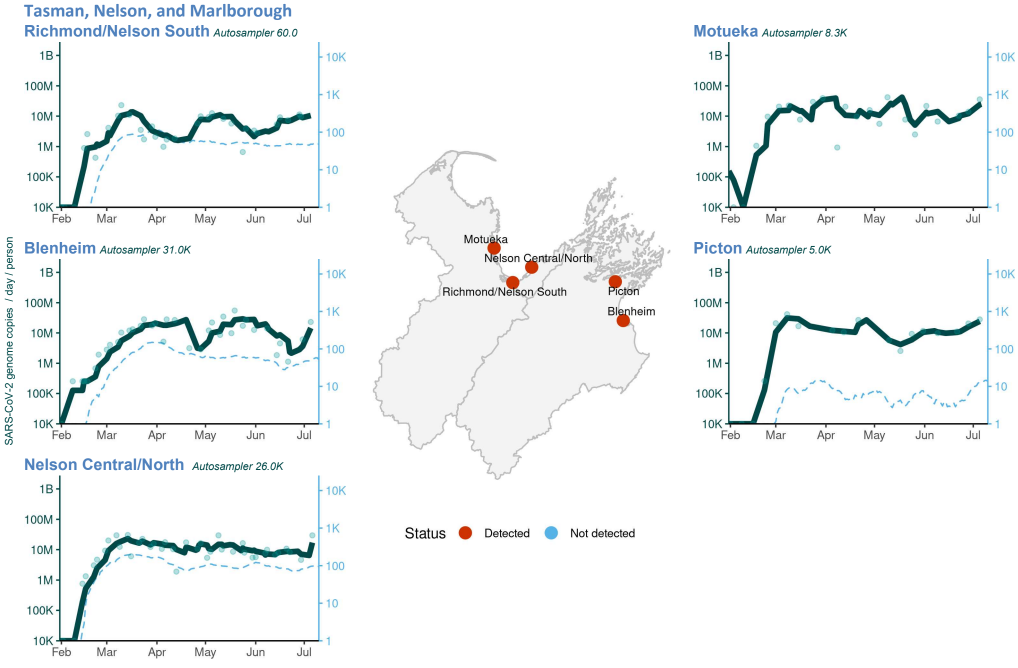
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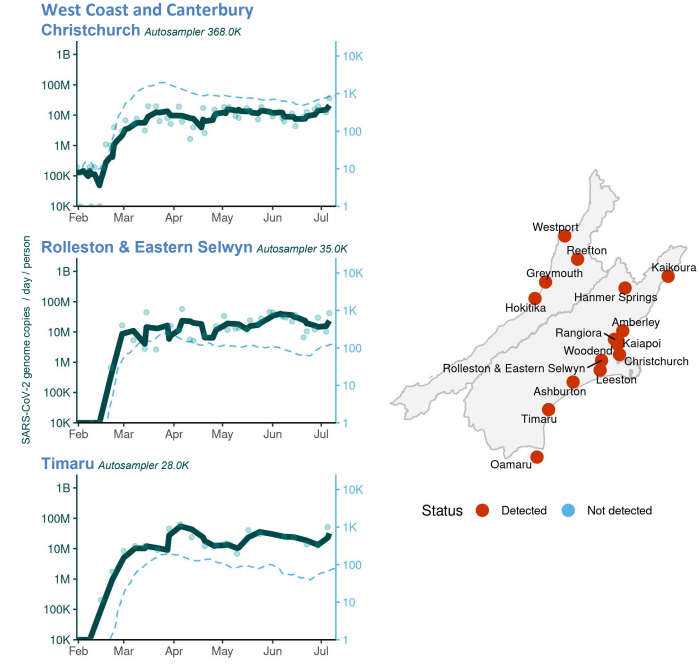
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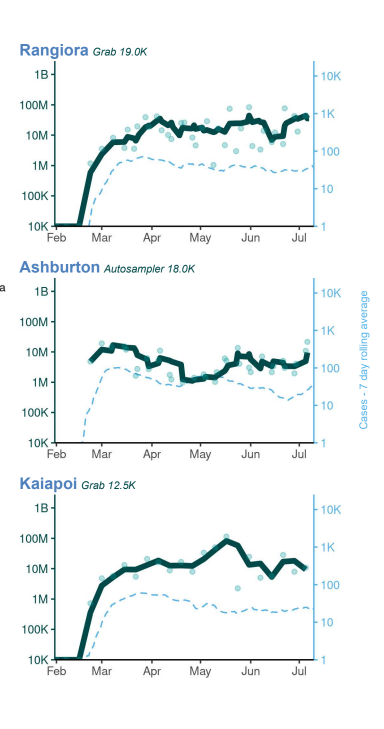


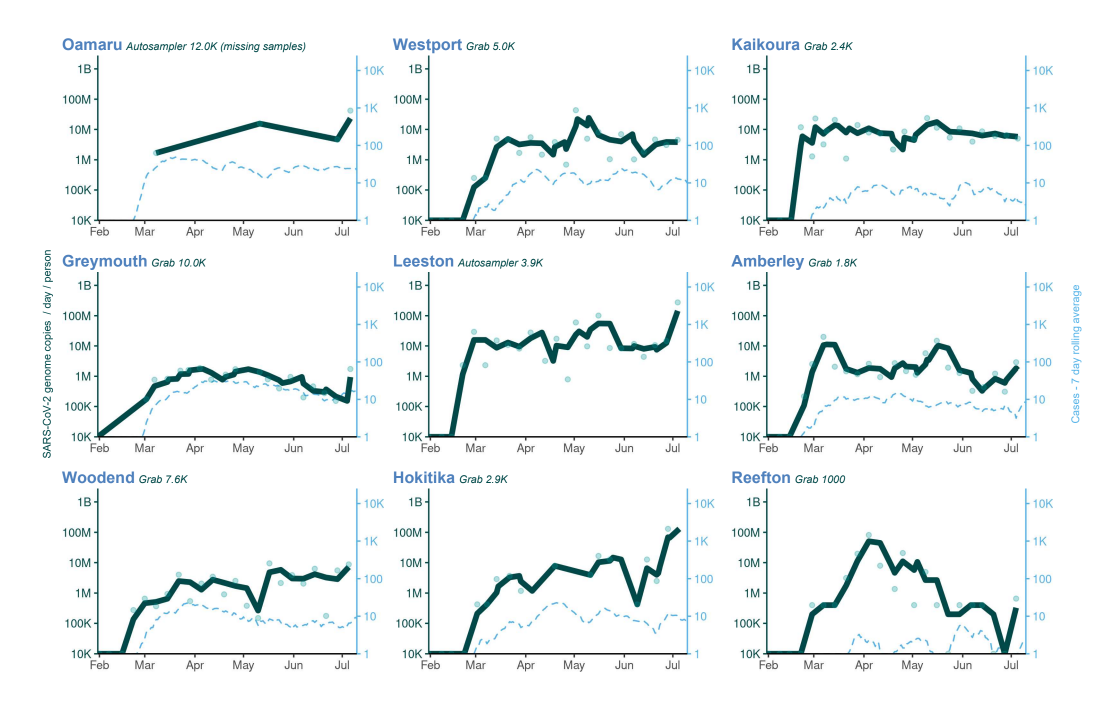


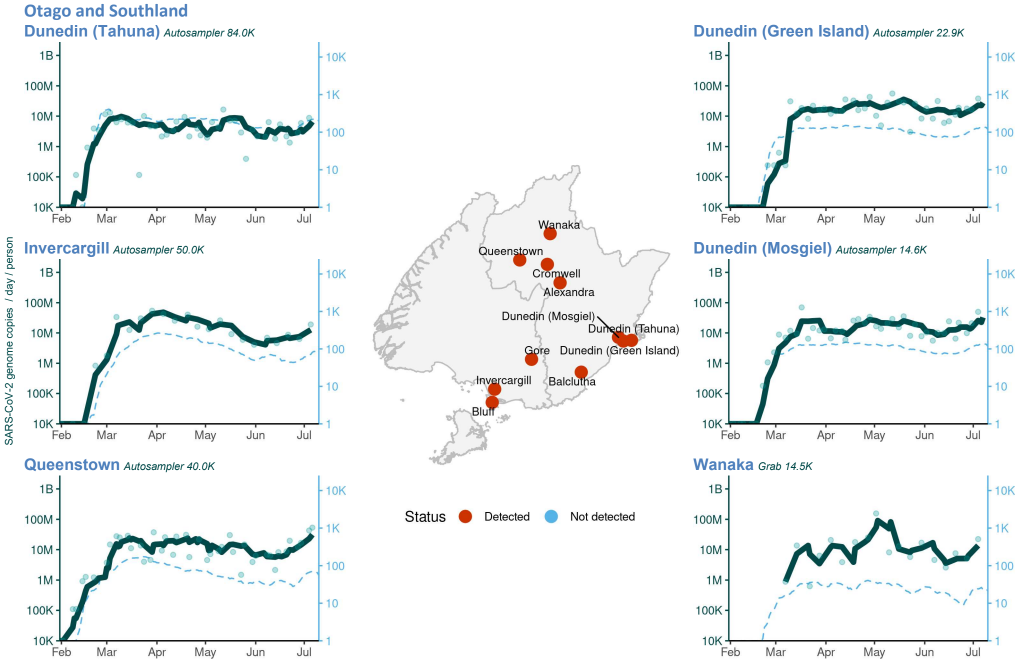


Cases - 7 day rolling average

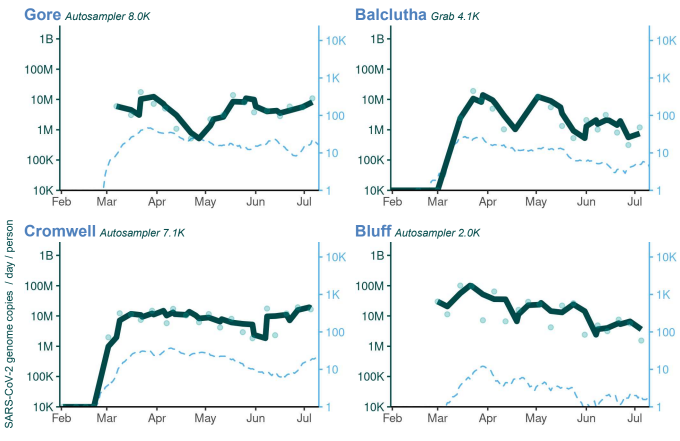








Cases - 7 day rolling average



Alexandra Autosampler 6.2K

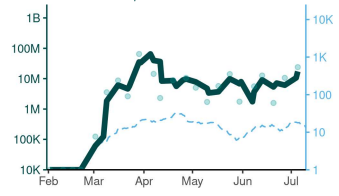


Table 1. Weekly summaryKey: Not Detected (below limit of quantification) Detected (quantifiable))Not sampled.

	2021 2022
Site	36 37 38 39 40 41 42 43 44 45 46 47 48 49 50 51 52 01 02 03 04 05 06 07 08 09 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27
Northland	
Ahipara	
Awanui	
Dargaville	
Haruru	
Hihi	
Hikurangi Kaeo	
Kaikohe	
Kaitaia	
Kaiwaka	
Kawakawa	
Kerikeri	
Kohukohu	
Mangawhai	
Maungaturoto	
Opononi	
Paihia	
Rangiputa	
Rawene	
Ruakaka Russell	
Russell Taipa	
Waipu	
Whangarei	
Whatuwhiwhi	
Auckland	
Army Bay	
Auckland	
(Combined)	
Auckland East Auckland	
Southwest	
Auckland West	
Beachlands	
Clarks Beach	
Helensville	
Kawakawa Bay	
Kingseat	
North Shore	
Omaha Pukekohe	
Snells/Algies	
Waiheke Island	
Waiuku	
Warkworth	
Wellsford	
Waikato	
Cambridge	
Coromandel	
Hamilton	
Huntly	
Kihikihi	
Kinloch	
Mangakino	
Maramarua Matamata	
Matamata	
Meremere	
Morrinsville	
Motuoapa	
Ngaruawahia	
Ngatea	
Omori	
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Paeroa	
Pauanui	
Putaruru	
Raglan –	
Taupo	

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Invercargill																																	

Acknowledgements

This work represents the combined efforts of a large number of individuals and organisations.

We continue to be indebted to the teams across the country who are collecting the wastewater that underpins this work.

The wastewater analysis has been undertaken by ESR by a team which may on any given week include contributions from: Joanne Chapman, Dawn Croucher, Joanne Hewitt, Joycelyn Ho, Anower Jabed, Susan Lin, Olivia Macrae, Ashley McDonald, Andrew Ng, Ashley Orton, Paula Scholes and Fatiha Sulthana. Data science analysis, visualisation and reporting is the result of team effort from: Franco Andrews, Bridget Armstrong, Raewyn Campbell, Gerhard de Beer, Richard Dean, Brent Gilpin, Joanne Hewitt, Dawen Li, Lillian Lu, Helen Morris and Bindu Priya. Ongoing support for this work from the Ministry of Health and ESR management is appreciated.

Notes

Sites and frequency of sample collection: The catchment population sites selected for the surveillance range from approximately 100 to over 1,000,000 individuals. The sites cover all regions of the country. Most major towns and all cities, as well as many smaller communities, are included. In early 2022, the wastewater catchment areas cover over 80% of the population connected to wastewater treatment plants. The sites from which samples have been collected have varied over the last 12 months. New sites may be added over time, and/or sampling may reduce in frequency or cease for other sites. The selection and frequency of sampling vary depending on the local population, access to wastewater collection points, staff availability to collect samples and risk factors. When included, samples are collected at least weekly, with twice weekly sampling being common. A number of samples have also been collected from non-WWTP sites (manholes and pump stations- mostly in Auckland).

Sampling method: The preferred option is to automatically collect a 24 hour 'composite' sample. This is where a pump automatically collects a small volume of wastewater every 15 minutes over 24 hours using a composite sampler. These samplers are available in some wastewater treatment plants. When composite samplers are not available, 'grab' samples are collected. These range from a sample being taken at a single point in time, to 3 samples taken over 30 minutes, to samples collected over a day. Grab samples represent only the composition of the source at that time of collection and may not be as representative as a 24-hour composite sampler. More variation may be expected with grab samples.

Laboratory analysis of wastewater samples: Samples are sent from each wastewater treatment plant to ESR. Processing of each sample commences within an hour or two of receipt. Processing involves the concentration of virus from 250 mL sample to approx. 1 mL using centrifugation and polyethylene glycol. Viral RNA is then extracted from a small volume of 0.2 mL concentrate to give a final volume of 0.05 mL The presence of SARS-CoV-2 RNA is determined using RT-qPCR. SARS-CoV-2 is considered detected when any of the RT-qPCR replicates are positive.

RT-qPCR: Reverse transcription (RT) to convert RNA to complementary DNA (cDNA), followed by quantitative PCR (qPCR). RT-qPCR is used for detection and quantification of viral RNA.

Method sensitivity: The protocol used to concentrate SARS-CoV-2 from wastewater allows for the sensitive detection of SARS-CoV-2 by RT-qPCR. ESR has shown that when 10 individuals are actively shedding SARS-CoV-2 RNA in a catchment of 100,000 individuals, there was a high likelihood of detecting viral RNA in wastewater (https://doi.org/10.1016/j.watres.2021.118032). Shedding by one individual may be detected in wastewater, but it does depend on many factors including the amount and duration of shedding. Very low levels in wastewater may be not able to be quantified (i.e., less than the limit of quantification- see below).

SARS-CoV-2 RNA detected (positive result): A positive detection in the wastewater indicates that at least one person has been shedding SARS-CoV-2 into the wastewater at some point during the time period that the sample was being collected. In some cases, detections could also be due to the shedding of low levels of SARS-CoV-2 RNA by a recently recovered case. The detection of SARS-CoV-2 RNA does not indicate that infectious virus is present.

SARS-CoV-2 RNA not detected (negative result): A negative result can occur because there are no active 'shedding' cases in the catchment or because the SARS-CoV-2 RNA concentration is too low to be detected, most likely because there are a very low number of cases in the wastewater catchment. Therefore, negative finding does not necessarily guarantee the absence of COVID-19 in the community.

Viral loads and normalisation: When detected, the SARS-CoV-2 RNA concentration is calculated as genome copies per L of wastewater. This is then converted to a viral load of genome copies/day/person. This conversion takes into account the flow rate of wastewater entering the treatment plant (the influent) and the population in the catchment. The flow rate is the total volume (m3 per day) recorded at the inlet of the wastewater treatment plant over 24 hours. This is a population-normalised viral load. Currently, the flow rate is the average annual flow rate, but will be replaced with daily flow rate when available (note that rainfall may significantly increase the flow rate at the inlet, diluting the sample, and may result in lower concentrations and a false negative result).

In future, SARS-CoV-2 RNA concentrations will also be normalised by testing for the presence of pepper mild mottled virus (PMMoV). PMMoV is a virus that infects peppers but not humans. Consumption of peppers or pepper products, such as chilli sauce, means that PMMoV is detected in wastewater – normally at very high concentrations. Therefore, PMMoV has been found to be a useful proxy for the amount of faecal material in a wastewater sample. For normalisation, the concentration of SARS-CoV-2 RNA is divided by that of PMMoV in each sample. Different normalisation methods may result in changes to some data points, but trends are unlikely to change significantly.

Limit of quantification: The lowest concentration of the target that can be reliably quantified is referred to as the limit of quantification. For those samples where SARS-CoV-2 is detected but cannot be quantified, a value of 5 genome copies/mL wastewater is used. While a standard method is being used, virus recovery can vary from sample to sample, and this may affect the quantitation.

Data subject to change: Data generated for the New Zealand Wastewater COVID-19 Surveillance Programme should be considered provisional and may be subject to change. Data may be incomplete for the most recent 2-week period due to processing, testing and reporting delays.

Data not shown:

- Data from 'ad hoc' sampling locations including from individual facilities/building (e.g., workplaces, prisons, MIQs) are not included.
- Results from certain samples may not be shown, as the result was either deemed invalid, or the sample could not be tested (e.g., leaked in transit, not labelled).

For further information please contact:

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END OF REPORT