



Federal Office of Public Health FOPH Public Health Directorate Communicable Diseases Division

Schwarzenburgstrasse 157 3003 Bern Switzerland

Geneva, September 26, 2022

## Swiss national SARS-CoV-2 genomic and variants surveillance program: report of the month of August

Geneva Centre for Emerging Viral Diseases

Division of Infectious Diseases

**Department of Medicine** 

Laboratory of virology

Division of Laboratory Medicine

Diagnostic Department

### 1. Summary

In August, COVID-19 cases numbers remained low and relatively stable in Switzerland, with the vast majority of cases still caused by BA.5.

Approximately 2.5% of the total number of cases identified in Switzerland in August were sequenced by the Surveillance program, yielding over 1,900 sequences. The vast majority of the sequences in Switzerland belong to BA.5, but BA.2.75 has been seen as well.

The currently circulating BA.4/5 variants are at least partially resistant to all the monoclonal antibody therapies used in Switzerland. In particular they are resistant to Evusheld®, which is composed of tixagevimab (complete resistance) and cilgavimab (partial resistance); and sotrovimab, for which the clinical effectiveness is unclear. BA.2.75, in contrast, displays high resistance to cilgavimab, but is susceptible to tixagevimab as lacks a spike 486 mutation present in BA.4/5 that leads to complete escape.

BA.4 and 5 sublineages with Spike 346 mutations enabling complete escape from cilgavimab are growing in frequency. Similarly, BA.2.75 sublineages (such as BA.2.75.2) have acquired not only the 346 mutations enabling complete escape from cilgavimab, but also the 486 mutations enabling complete escape from tixagevimab. This means that these variants completely escape Evusheld® combination treatment. At the end of August, they represented approximately 7% of the sequences identified by the national genomic surveillance program. Unfortunately, while there is still considerable uncertainty, one of these lineages displaying complete escape from Evusheld® is likely to soon be the next dominant variant.

The escape from immunity derived from vaccination or previous exposure by these subvariants is likely to be much weaker than the escape seen against the monoclonal antibodies. While these variants substantially escape immunity from pre-Omicron vaccination and prior infections, updated vaccines or previous infection with an Omicron lineage will improve protection. Most importantly these subvariants have so far not resulted in a large increase of hospitalizations.

# 2. <u>Description of the Swiss national SARS-CoV-2 genomic and variants surveillance program.</u>

The overall goal of the program is to provide epidemiological trends and to highlight meaningful observations.

Because greater transmissibility and/or immune escape potential of the different VOCs and VOIs can result in new surges in COVID-19 numbers despite the vaccination campaign, this program aims to closely monitor each variant displaying mutations known to be linked with either increased transmissibility or immune escape potential.

Currently, 12 diagnostic laboratories are participating in the program, including university hospital centres in Switzerland (Geneva, Lausanne, Bern, Basel, Zurich, Ticino), in addition to private laboratories (Viollier; Medisupport including Dianalabs, Polyanalytic, Dianalabs Valais, Proxilab and Bioanalytica; Labor Team W, Risch), cantonal-based laboratories (Hôpital du Valais – Institut Central), Spital Region Oberaargau (Bern, Solothurn, Aargau, Luzern), and 2 high-throughput sequencing platforms (Health 2030 Genome Centre in Geneva, Genomics Facility Basel run by ETH Zürich and University of Basel).

Processed sequencing data are shared openly within 14 days from positive PCR result through the GISAID platform (<a href="https://www.gisaid.org">https://www.gisaid.org</a>) and eventually through the Swiss Pathogen Surveillance Platform (SPSP). The centralized analysis of this National Surveillance will be performed by the groups of Pr. Neher, Pr. Stadler and Dr. Althaus, where variants of concern are counted, analyzed and all sequences scanned for new variants with potential changes in antibody-Spike interactions (<a href="https://nextstrain.org/groups/swiss">https://covariants.org/per-country</a>, <a href="https://covariants.org/per-country">https://covariants.org/per-country</a>, <a href="https://covariants.org/per-country">https://covariants.org/per-countr

In order to complement the genomic surveillance based on patient samples, sequencing of SARS-CoV-2 in wastewater samples is also performed. Samples are collected daily in six wastewater treatment plants (WWTP), under the coordination of Eawag. Up to 50 samples per week over the first 26 weeks have been performed. The sequencing and analysis of these samples, including detection of variants, is done under the coordination of Prof Niko Beerenwinkel. It started in December 2020 for Lausanne and Zurich, and in February 2021 for all six WWTP (https://bsse.ethz.ch/cbg/research/computational-virology/sarscov2-variants-wastewater-surveillance.html).

Immunological characterization of the variants within the surveillance program is coordinated by Professor Trono's team at EPFL.

This report has been produced by <u>Erik Boehm</u>, <u>Pauline Vetter</u>, <u>Marc Friedli</u>, Samuel Cordey, Richard Neher, Christian Althaus, Martina Reichmuth, Cornelius Römer, Niko Beerenwinkel, David Dreifuss, Chaoran Chen, Tanja Stadler, Priscilla Turelli, Didier Trono, Emma Hodcroft, Silvan Heeb, Anna Fesser, and Laurent Kaiser. The list of the participants and collaborators of the program can be found at the end of this report in the appendix.

This report covers the period of 1 August to 28 August, 2022 (weeks 31, 32, 33, 34). All data presented in this report are based on the sampling date.

# 3. <u>Variants of concern (VOCs)</u>, variant of interest (VOI) and other surveilled variants: brief summary and special focus

Five variants and their sub-lineages are considered VOCs by the WHO, B.1.1.7 (Alpha), B.1.351 (Beta), P.1 (Gamma), B.1.617.2 (Delta), and B.1.1.529 (Omicron). Worldwide, all VOCs except Omicron have essentially disappeared from samples collected within the last 30 days (<a href="https://www.who.int/publications/m/item/weekly-epidemiological-update-on-covid-19---24-august-2022">https://www.who.int/publications/m/item/weekly-epidemiological-update-on-covid-19---24-august-2022</a>).

#### Omicron

The Omicron VOC (B.1.1.529) is characterized by a high divergence in the spike protein, which has allowed it to substantially escape immunity conferred by vaccination (using the original Wu-1 sequence) and prior infection with pre-Omicron variants. This VOC currently has 5 sublineages: BA.1-5, many of which now have further "sub-sublineages". Despite all being considered "Omicron", some BA.X sublineages differ from each other (in terms of mutation counts) more than the earlier VOCs differed from the original Wu-1 strain (although BA.4 and 5 had the same spike when they first appeared). These sublineages have successively replaced each other, with BA.5 currently being dominant.

Notably, this is the first VOC to have subvariants causing multiple successive waves, arguing for treating Omicron subvariants as distinct VOCs.

While BA.5 replaced BA.2, highly derived BA.2 sublineages have appeared, in particular, BA.2.75. Growth signals outside of India were initially small, but further observation has confirmed that this variant seems to outcompete BA.4 and 5.

Subvariants of BA.4 and BA.5 containing additional mutations, particularly at position 346 of the spike protein (such as BF.7 and BA.4.6), are also cause for concern as they seem to confer a growth advantage and enable complete escape from cilgavimab, rendering it fully resistant to Evusheld®. In particular, BA.4.6 with the R346T mutation has signs of growth in many western countries. Some of these BA.4 and 5 subvariants seem to be competitive with BA.2.75.

BA.2.75 is likewise spawning subvariants, in particular BA.2.75.2, which has a spike 486 mutation enabling complete escape from tixagevimab. BA.2.75 subvariants also seem to be picking up 346 mutations enabling complete escape from cilgavimab.

## Detection

All sub-lineages are still detected by RT-PCR tests, but BA.1,3,4, and 5 (but not BA.2) exhibit S-gene target failure with some assays that can be used as a proxy prior to sequencing, as seen with VOC Alpha (and due to the same deletion as found in Alpha). Given the current virus circulating, the absence of S-gene target failure is currently a good proxy for BA.2.75 infection. Likewise, its presence is indicative of a likely BA.5 (or BA.5 subvariant) infection. Antigenic tests are still able to detect these variants, although a recent study suggested that the tests were significantly less sensitive to Omicron BA.2, the drop in sensitivity relative to the currently circulating BA.5 was minor. There is some evidence that sensitivity may decrease depending on the patient's immune status, which may confound results.

## Immune escape

Extensive data demonstrates that Omicron variants are substantially able to evade neutralizing antibodies (nAbs) from non-Omicron infections and after 2-3 doses of vaccine. Escape from monoclonal antibodies is extensive and is covered by the "Therapeutic intervention effectiveness" section.

Data suggests low neutralization of BA.2.75 and BA.4/5 by sera from BA.1 and BA.2 vaccine-breakthrough infections. A recent publication found that BA.4 or BA.5 convalescent sera has reduced neutralization of (a 2.4 to 2-6 fold reduction) the BA.4.6, 4.7, and 5.9 sublineages – all of which contain either R346T or R346S mutations. Similarly, the BA.2.75.2 sublineage neutralization titers were 6.5x lower than the neutralization titers for BA.5. Furthermore these sublineages completely escape neutralization by Evusheld® and have substantially reduced neutralization by sotrovimab.

While all Omicron sublineages largely escape humoral immunity, cell mediated immunity remains mostly intact.

A  $3^{rd}$  vaccine dose or a combination of previous infection and 2 vaccine doses results in poor neutralizing titers against the circulating BA.4 and 5, and a corresponding poor efficacy at preventing infection. Efficacy preventing hospitalization and death is reduced but remains high. The efficacy of four doses at preventing symptomatic disease is relatively high (>60%), but remains poor (~30%) at preventing any infection.

Updated bivalent vaccines using the Omicron BA.1 sequence performed much better against the currently circulating BA.4/5 and sublineages, but neutralization was still reduced by approximately 3-fold relative to BA.1. Similarly, bivalent vaccines with BA.1 and in particular BA.5 did improve neutralization of BA.2.75, although the neutralization was still reduced relative to BA.1 and BA.5 respectively. Unlike the monoclonal antibody situation, neutralization capacity of patient sera against new variants is expected to gradually decline rather than suffer sharp drops in or complete loss of efficacy. Still, it is clear that additional updates would be very beneficial.

### Severity

A multitude of clinical, *in vitro* and *in vivo* studies indicate that Omicron BA.1 and BA.2 cause intrinsically milder disease. Importantly, BA.1,2, and 3 use a TMPRSS2-independent entry pathway, and exhibit less cleavage of the spike protein and corresponding cell to cell fusion. This suggests an altered tropism that may favor replicating in the upper rather than lower respiratory tract, and is a possible explanation for observations of the infections being less severe. BA.4 and BA.5 have apparently regained the ability to use TMPRSS2 mediated entry pathways and exhibit heightened cell to cell fusion relative to BA.1&2 (although it is still not quite as high as that of earlier variants), suggesting that their intrinsic severity may be closer to that of previous variants.

There are conflicting animal studies with regards to the question of intrinsic severity, and severity determinations are complicated by an over representation of reinfections/vaccine breakthroughs, which are expected to be mild due to the protective effect of prior vaccination/exposure.

Significant protection is afforded by vaccination and past exposure, which may mask differences in intrinsic immunity, and render such differences in naïve populations largely irrelevant from a public health perspective.

## Therapeutic intervention effectiveness

All sublineages display complete escape from combination of casirivimab/imdevimab. A matched cohort study found a noticable clinical benefit of sotrovimab treatment during a BA.1 wave. Both *in vitro* and *in vivo* data suggests that sotrovimab is even less effective against BA.2, 4 and 5. Indeed a study in Qatar failed to find any beneficial effect of sotrovimab treatment in the context of BA.2 infections. In contrast a study done in the USA found a beneficial effect for both BA.1 and BA.2, although it had significant limitations specifically regarding its BA.2 conclusions. Currently, there is conflicting data regarding the efficacy of sotrovimab against BA.2.75, with reports variously indicating that its sensitivity is relatively unchanged or decreased relative to BA.5. Further studies are needed to clarify.

Similarly, *in vitro* data suggests that both antibody components of Evusheld® (tixagevimab and cilgavimab) will have significantly reduced neutralization against BA.4/5, but one component (cilgavimab) should retain efficacy against BA.2. Additional spike 346 mutations seen in BA.4/5 sublineages lead to complete escape from cilgavimab, and these appear it is likely that cilgavimab will lose any efficacy that it still had as these sublineages become the dominant BA.4/5 sublineages.

Interestingly, while BA.2.75 contains an escape mutation against cilgavimab associated with up to a 12 fold reduction in neutralization, it lacks the spike 486 escape mutation of BA.4/5 and is susceptible to tixagevimab. The overall effect is only a modest reduction in neutralizing activity by Evusheld® against BA.2.75 relative to BA.2, and a substantial improvement relative to BA.4/5.

Unfortunately, BA.4, BA.5, and BA.2.75 are forming sublineages that are outcompeting their parent strains, which contain additional escape mutations. Although BA.2.75 is susceptible to tixagevimab, BA.2.75.2 in particular has mutations resulting in a complete escape from tixagevimab. Similarly spike 346

mutations, which confer complete escape from cilgavimab in the BA.4 and 5 sublineages, are seen in BA.2.75, BA.4, and BA.5 sublineages. Notably, these lineages include BQ.1 (a BA.5 sublineage), and BA.2.75 – both these lineages are expected to completely escape Evusheld, and are outcompeting their parent lineages.

As sotrovimab and tixagevimab/cilgavimab are still being used, additional mutations causing escape from them will thus be closely followed (Table 1, and section 7).

In addition to mAbs, there are a number of other antiviral treatments under development, such as 3CL like protease inhibitors like Paxlovid® (PF-07321332, nirmatrelvir/ritonavir) or RNA nucleotide analogues (which interfere with replication of the viral genome, such as molnupiravir). Patients taking this drug have been noted to have a higher "rebound" frequency than those who do not take the drug. Importantly, while viral loads may go up when the drug is discontinued, the drug still efficiently limits viral replication when given. Resistance mutations to Paxlovid® have been noted, in particular L50F and E166Q. Notably, E166V has been observed to occur at higher frequencies in Paxlovid® treated groups during clinical trials. Given the low proportion of the population that receives Paxlovid®, it is unclear if there is enough selection pressure for these mutations to become common. Indeed, known escape mutations against Paxlovid® are currently very rare.

Resistance to molnupiravir comes at the cost of significantly reduced viral fitness, so limited use is unlikely to lead to resistant strains. Preliminary data confirms that molnupiravir, Paxlovid®, and remdesivir all retain full *in vitro* efficacy against Omicron sub-lineages.

AA position	World	Europe	Switzerland	Mutations
Sotrovimab				
337	16	6	1	R/L/H/T
340	78	55	6	K/A/G/Q/V
356	357	83	2	Т
377	0	0	0	K
Cilgavimab				
346	19 769	7 154	124	I/K/S/T
444	1 014	350	7	E/Q/R
445	760	247	7	Α
446	3 127	601	18	S/V
450	0	0	0	K
Paxlovid®				
144	0	0	0	M/F/A/G/Y
165	1	1	0	Т
166	0	0	0	Q/V
172	0	0	0	Q/F
192	0	0	0	T/S/V

Table 1: Positions where mutations are known to result in escape from sotrovimab, cilgavimab, or Paxlovid®, their prevalence, and the specific amino acid mutations known to result in escape, August 2022. Note: defining mutations of the currently dominant Omicron lineages and are not shown. Only mutations enhancing escape in sublineages are shown.

The circulation of specific mutations leading to decreased effectiveness of known therapeutics available for clinical use in Switzerland will be surveilled (see Section 7), but their prevalence remains low.

Notably, there has been a substantial and worrying increase in Europe and the rest of the world in mutations

leading to escape from Cilgavimab (particularly at position 346) although, aside from the 346T mutation, their presence in Switzerland remained low during the month of August. While the overall prevalence is low, it is worth noting that the number of sequences bearing a 346 mutation in Switzerland more than doubled in August relative to July.

## 4. Epidemiology in Switzerland and number and origin of sequences produced through the program during the surveilled period

Data in this report comes from 3 sources: 1) The publicly available data on COVID-19 as reported by the FOPH (https://www.covid19.admin.ch), including data that is declared to the FOPH by the different laboratories in Switzerland; 2) data originating from laboratories participating in the surveillance program; and 3) sequences submitted to GISAID, for which the corresponding infected person was in Switzerland (resident or recent travel history to Switzerland).

General caveat: the numbers and denominators are fluid and variable over time; and are subject to change depending notably on the different databases used, and variable declaration delays. All data generated by this program is also submitted to SPSP.

Data will be presented here by regions, using the same region definitions that are used for the influenza sentinel surveillance system in Switzerland. Data are presented according to residency post-code.



Region 1 includes the cantons of Geneva, Neuchatel, Vaud and Wallis

Region 2 includes the cantons of Bern, Fribourg and Jura Region 3 includes the cantons of Aargau, Basel (Basel-Stadt and Basel-Land) and Solothurn

Region 4 includes the cantons of Luzern, Unterwalden (Obwalden and Niedwalden), Schwitz, Uri and Zug

Region 5 includes the cantons of Appenzell (Appenzell Ausserrhoden and Appenzell Innerrhoden), Glarus, Sankt Gallen, Schaffhausen, Thurgau and Zurich.

Region 6 includes the cantons of Graubünden and Ticino.

Divisions of the different regions, from <a href="https://covariants.org/per-country">https://covariants.org/per-country</a>

## Number of cases processed by the laboratories participating in the surveillance program

During August (1 August to 28 August), the FOPH reported a total of 70'550 confirmed SARS-CoV-2 cases in Switzerland, representing a decrease from July. Supplementary Table 1 provides an overview of the number and incidence of confirmed cases, the effective reproduction number  $R_{\rm e}$ , the number and incidence of tests, test positivity, the number and proportion of sequenced samples, and the number and proportion of VOCs by canton, region and for Switzerland overall.

As of the writing of this report, the laboratories participating in this program reported 21'781 positive tests during the surveilled program, which represents about 30% of the total number of cases reported in Switzerland (including both PCR and antigen-based tests). Detailed data regarding the total number of tests performed each week by the laboratories participating in the surveillance program (including negative and positive tests numbers, and the number of the positive tests that have been sequenced) are available in supplementary Table 3.

## Number of declared SARS-CoV-2 sequences produced through the surveillance program (presented by submission date, further declarations are still ongoing)

A total of 1'930 SARS-CoV-2 sequences have been declared to have been submitted to GISAID during this period. There are 2'272 sequences are available for this period on GISAID as of 23 September 2022, and the difference may be explained by reporting delays and differences between collection/submission dates.

This represents around 2.5% of all cases detected in Switzerland during the surveilled period (see Supplementary Table 2 and 3 in the Appendix for details). Of note, this number includes sequences from samples received from other laboratories at the request of the Cantonal physician teams.

am during the survein	ed period by calerida	i week (data is incomplete).
Week	Date	Number of sequences declared and successfully submitted to GISAID during the surveilled period, by all laboratories in the program
31	Aug. 1 to Aug. 7	540
32	Aug. 8 to Aug. 14	453
33	Aug. 15 to Aug. 21	560
34	Aug. 22 to Aug. 28	377
	Total	1 930

Table 2 shows the number of sequences successfully submitted to GISAID through the surveillance program during the surveilled period by calendar week (data is incomplete).

Table 2: number of sequences submitted to GISAID through the surveillance program. Note these data are not by sampling date but rather by submission to GISAID date.

The total number of SARS-CoV-2 sequences declared and submitted to GISAID by each laboratory during this period is available in Supplementary Table 3 in the appendix.

<u>Covering of sequencing in Switzerland and contribution of the national SARS-CoV-2 surveillance</u> sequencing program

As shown in Figure 1, the total number of SARS-CoV-2 sequences submitted per week generally remained stable during August (weeks 31-34), while the fraction sequenced was generally stable as well. Since the beginning of this program, almost all of the sequences available, and on which the surveillance is conducted, come from the national surveillance program.

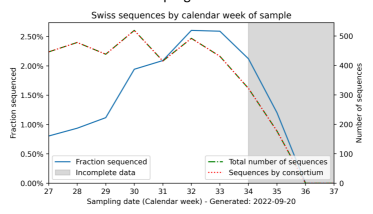


Figure 1: Number of SARS-CoV-2 sequences available for Switzerland (total available Swiss sequences in GISAID in green, Swiss sequences submitted through the program in dotted orange) and fraction of the total number of positive cases declared to the FOPH that have been sequenced (blue curve).

During the surveilled period, the absolute number of sequences generated remained fairly high, at hundreds of sequences per week. The total proportion of positive sequenced cases was approximately 2-2.5% during the month, with the higher percentages in weeks 32 and 33. These sequences include those from sites with hospitalized patients, and thus should be adequate for surveillance.

Figure 2 displays the number and fraction of SARS-CoV-2 cases sequenced for each Swiss region. Regions 4 and 6 had the lowest total number of sequences, while regions 4 and 3 had the lowest fraction of cases sequenced.

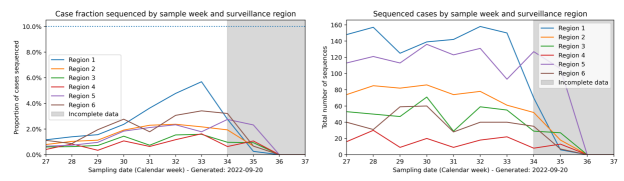


Figure 2: Sequencing coverage among the different Swiss regions per week, presented by fraction of cases sequenced (left) and by number of sequences (right).

## 5. Recently circulating variants in Switzerland as of August 2022

Determination of the proportion of total number of sequences over time falling into defined variant groups is done by Emma Hodcroft's team and displayed on the CoVariant website (<a href="https://covariants.org/per-country">https://covariants.org/per-country</a>). Those results are based on the total number of sequences submitted to GISAID over the time period for Switzerland. Those data mainly, but not exclusively, come from the national genomic surveillance program since its beginning (see Figure 1).

Omicron BA.5 was by the most commonly retrieved lineage in August (Figures 3-5, Table 3). No BA.3 was found during the month of August. Delta an Omicron BA.1 have essentially disappeared, with 1 sequence each sequence found in August. An estimate of the total number of VOCs circulating in Switzerland, corrected by taking in account the fraction of sequencing in Switzerland is available through the covSPECTRUM program, developed at ETHZ, at <a href="https://cov-spectrum.ethz.ch/explore/Switzerland">https://cov-spectrum.ethz.ch/explore/Switzerland</a>. Notably, 15 BA.2.75 sequences were found in Switzerland in the month of August.

Region	BA.1	BA.2	BA.2.12.1	BA.4	BA.5	Delta	None	other	sequences	cases	% sequenced
All	2	39	1	55	1370	0	5	188	1660	70550	2.35%
1	0	7	1	20	434	0	0	58	520	12382	4.20%
2	2	5	0	10	216	0	1	31	265	12008	2.20%
3	0	6	0	5	150	0	1	10	172	14218	1.21%
4	0	3	0	2	45	0	0	7	57	5554	1.03%
5	0	14	0	13	385	0	2	60	474	21257	2.23%
6	0	4	0	5	113	0	1	20	143	5131	2.79%

Table 3: number of sequences corresponding to selected variants in each region of Switzerland from 1 August 2022 to 28 August 2022, according to data received by 22 September, 2022. Sequences with poor coverage where lineages could not be assigned are listed as "None".

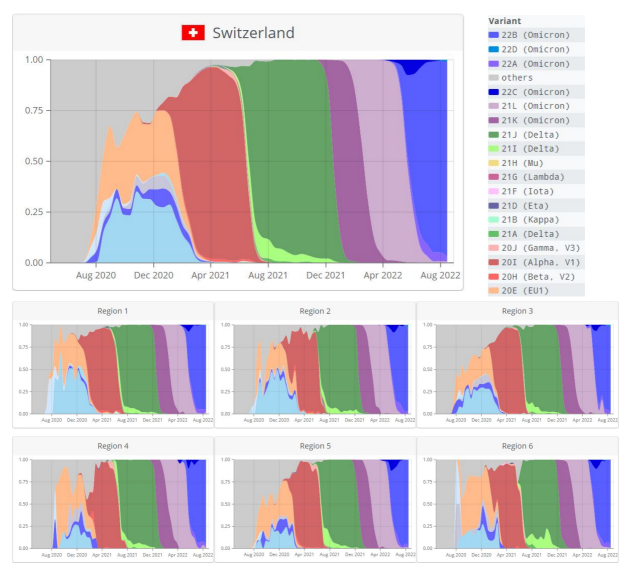


Figure 3: proportion of the total number of sequences (not cases), over time, that fall into defined variant groups, for Switzerland. Screenshot from CoVariants website. Dynamic navigation is available at <a href="https://covariants.org/per-country">https://covariants.org/per-country</a>. Dark green indicates the B.1.617.2 (Delta) lineage or its sub-lineages. Dark Red indicates B.1.1.7 (Alpha). Purple/21K indicates Omicron BA.1, Light purple/21L indicates Omicron BA.2. Dark blue/22C indicates Omicron BA.2.12.1, while a faint blue/22B indicates Omicron BA.5 and a blueish-purple/22A indicates Omicron BA.4. Cyan/22D indicates BA.2.75

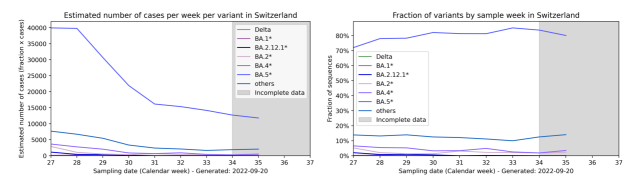


Figure 4: (Left): Percentage of circulating VOCs and VOIs in Switzerland by week, up to the 34 first weeks of 2022 (total number of BA.2, BA.2.12.1, BA.4, and BA.5 sequences from Switzerland and successfully submitted to GISAID are shown here). Note the grey shaded area indicates a period of incomplete data. (Right): Estimated number of sequences BA.2, BA.2.12.1, BA.4, and BA.5 retrieved during the surveilled period.

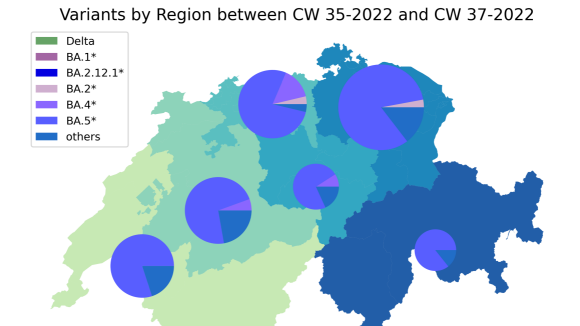


Figure 5: Distribution of variants per region, for the Start of September 2022 shown on a map. The size of the pie chart corresponds to the total number of sequences. Note the large share of the share of the BA.5 lineage, in all regions as of the time of this report.

Pie area represents number of cases

#### 6. Assessment of the competition between the different variants in Switzerland

The competition between different SARS-CoV-2 variants can be modelled using multinomial logistic regression. The analysis by Dr. Althaus' group is based on sequences retrieved from CovSPECTRUM. The results correctly predicted that Omicron would be the dominant SARS-CoV-2 variant in Switzerland. These models also correctly predicted that the BA.1 sub-lineage would be replaced by the BA.2 sublineage (Figure 6). In May, the analysis predicted that BA.2 would be displaced by other variants (mainly BA.5), which was indeed correct. The current estimates suggest that BA.5 prevalence will decline, with other lineages starting to displace it, although which specific lineage is not clear.

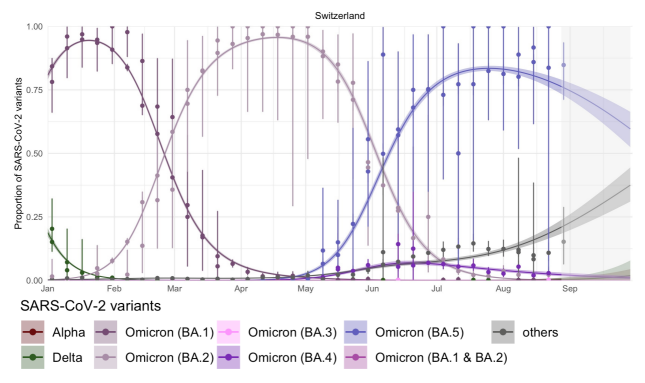


Figure 6: Observed and modeled proportion of SARS-CoV-2 variants over time in Switzerland. Left: graphed by linear proportions. Right: graphed by Log<sub>10</sub> proportions. Since early 2021, multiple successive variants have been observed to displace all previous variants (often achieving a dominance of >99.9% of the circulation). Alpha was replaced by Delta, followed by Omicron BA.1, then Omicron BA.2, and now Omicron BA.5. Model fits are based on a multinomial logistic regression with splines.

### 7. Surveillance of mutations associated with reduced mAb treatment efficacy

A number of mutations have been reported to result in significantly reduced *in vitro* effectiveness of the monoclonal antibodies used to treat patients in Switzerland. Notably, all Omicron lineages are completely resistant to neutralization by casirivimab/imdevimab. Sotrovimab retains substantial neutralizing activity against BA.1, but this neutralization is significantly decreased for BA.2/4/5 (and in particular, some of their sublineages). The escape is however not complete, and sotrovimab retains significant activity at higher titers – thus the frequencies of mutations (aside from lineage defining mutations) reported to escape neutralization by sotrovimab are still being followed (Table 4).

	337H		337L		337R		337T	
Dates	Global	Switzerland	Global	Switzerland	Global	Switzerland	Global	Switzerland
1.8 to 4.9	6	0 (Eur:3)	3	1 (Eur: 2)	5	0 (Eur: 1)	2	0

		340A		340K		340G		340Q	
Γ	Dates	Global	Switzerland	Global	Switzerland	Global	Switzerland	Global	Switzerland
	1.8 to 4.9	6	1 (Eur: 3)	33	4 (Eur: 22)	4	0 (Eur: 1)	28	1 (Eur: 26)

	340V	340V		356T		
Dates	Global	Switzerland	Global	Switzerland	Global	Switzerland
1.8 to 4.9	7	0 (Eur: 3)	357	2 (Eur: 83)	0	0

Table 4: Global and Swiss counts of sequences bearing escape mutations from sotrovimab for August 2022 (1 August to 28 August)

Evusheld®, which consists of a cocktail of 2 mabs, tixagevimab and cilgavimab, is also used in Switzerland. Tixagevimab is not effective against BA.2/4/5, although the BA.2.75 sublineage is neutralized by tixagevimab, likely due to a reversion mutation. Cilgavimab is able to neutralize BA.2/4/5, although its neutralization is reduced against BA.4/5, and eliminated in the BA.4.6/4.7/5.9 sublineages.

Mutations resulting in partial escape from cilgavimab are shown in table 5. As seen in tables 4&5, known mutations enabling escape from sotrovimab and Evusheld® have been detected in Switzerland, and are growing in frequency. During August, variants expected to be completely resistant to Evusheld® (cilgavimab and tixagevimab) made up nearly 7% of the sequences. The 346T mutations are also associated with escape from humoral immunity, thus it is not expected to be seen only in response to mAb treatment.

_	3461		346K		346S		346T	
Dates	Global	Switzerland	Global	Switzerland	Global	Switzerland	Global	Switzerland
1.8 to 4.9	1 498	33 (Eur: 1133)	233	1 (Eur: 118)	971	7 (Eur: 491)	20 216	107(Eur:6973)

	444E		444Q		444R		445A	
Dates	Global	Switzerland	Global	Switzerland	Global	Switzerland	Global	Switzerland
1.8 to 4.9	7	0 (Eur: 4)	15	0 (Eur: 5)	1119	7 (Eur: 395)	899	7 (Eur: 327)

	446S		446V		450K		
Dates	Global Switzerland		Global	Switzerland	Global	Switzerland	
1.8 to 4.9	3 498	14 (Eur: 731)	83	0 (Eur: 34)	0	0	

Table 5: Global and Swiss counts of sequences bearing escape mutations from cilgavimab for August 2022(1 August to 28 Augustr). Note: defining mutations of the Dominant BA.4/5 Omicron lineages and are not shown.

Paxlovid®, which inhibits the main viral protease encoded by the viral gene nsp5 is also used in Switzerland. In the absence of any treatment with Paxlovid®, escape mutations are not expected to produce any benefit, and the mutations are not linked to general antigenic shift like the escape mutations to the therapeutic mAbs (ie: sotrovimab and Evusheld®). This likely explains the scarcity of

escape mutations against Paxlovid®. Notably, the only escape mutation found worldwide during this period belong to a BA.2.75.1 sequence. Mutations resulting in partial escape from Paxlovid® are shown in table 6.

	144M		144F		144A		144G	
Dates	Global	Switzerland	Global	Switzerland	Global	Switzerland	Global	Switzerland
4.7 to 31.7	0	0	0	0	0	0	0	0

	144Y		165T		166Q/V		172Q	
Dates	Global	Switzerland	Global	Switzerland	Global	Switzerland	Global	Switzerland
4.7 to 31.7	0	0	1	0 (Eur: 1)	0	0	0	0

	172F		192T		192S		192V	
Dates	Global	Switzerland	Global	Switzerland	Global	Switzerland	Global	Switzerland
4.7 to 31.7	0	0	0	0	0	0	0	0

Table 6: Global and Swiss counts of sequences bearing escape mutations from Paxlovid®: Sequenced escape mutations remained extremely rare worldwide during the month of August (1 August to 28 August)

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#### 8. Wastewater surveillance program

Sequences are obtained from six wastewater treatment plants (WWTPs) that are tested on a daily basis. During the month of August, BA.5 kept its major variant position, making up 93% - 99% of the sequenced SARS-CoV-2 RNA depending on the location (Figures 7, 8). After rising while BA.2 was still the dominant variant, the BA.4 variant has now declined and is either almost extinct, or makes up no more than 5% of the sequenced RNA depending on the location.

BA.2.75 was first detected in samples from Geneva on June 22nd, and on the June 23rd in samples from Zürich. Since then, detections of the variant have been common occurrence, but a consistent rise in relative abundance has not been observed.

Quantification of BA.4 and BA.5 in sewage has exhibited some particular challenges. Those variants share most of their defining mutations with BA.2, which was the dominating variant at the time of their introduction and spread. The very limited number of discriminating mutations thus requires particular care in the wet lab protocols and computational methods used. However, although challenging, quantification is still possible.

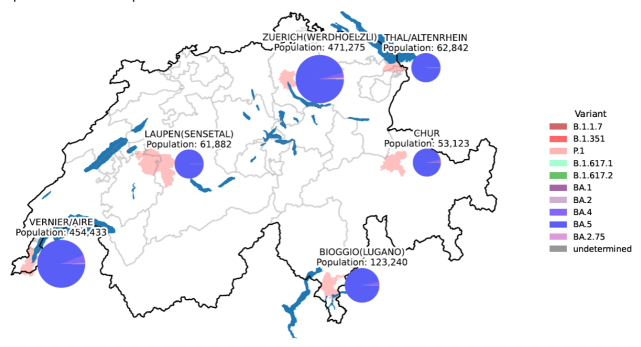


Figure 7:

Overview of the relative abundances of variants of SARS-CoV-2 at the end of August 2022, estimated from wastewater samples collected daily during the month in WWTPs located in 6 different Swiss locations. Pie chart areas are proportional to population connected to the wastewater treatment plants. Pink shaded areas represent catchment areas (boundaries from 2017). Population connected to Vernier (GE) wastewater treatment plant also includes ~44,000 inhabitants from neighbouring French communities.

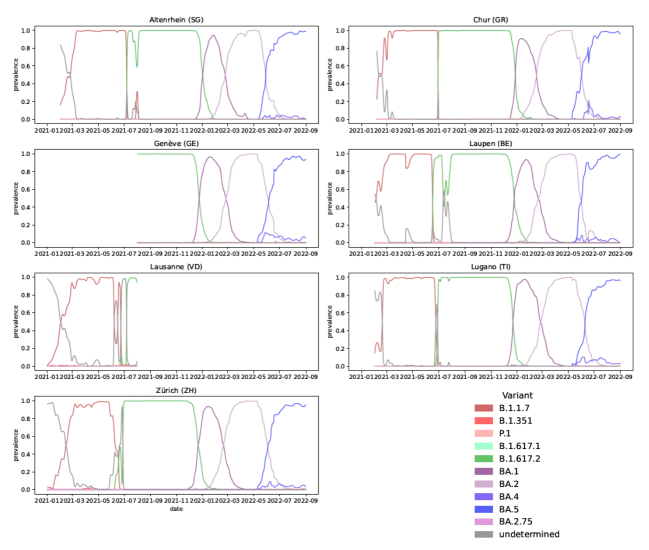


Figure 8: Relative abundances of variants of SARS- CoV-2 estimated from wastewater samples collected daily until August 31, 2022 (except Lausanne: July 31, 2021) in WWTPs located in 7 different Swiss cantons. An online dynamic navigation is available at <a href="https://bsse.ethz.ch/cbg/research/computational-virology/sarscov2-variants-wastewater-surveillance.html">https://bsse.ethz.ch/cbg/research/computational-virology/sarscov2-variants-wastewater-surveillance.html</a>.

### **Acknowledgements:**

https://bsse.ethz.ch/cevo/research/sars-cov-2/swiss-sars-cov-2-sequencing-consortium.html

Meriem Bekliz, Kenneth Adea, Pauline Vetter, Christiane S Eberhardt, Krisztina Hosszu-Fellous, Diem-Lan Vu, Olha Puhach, Manel Essaidi-Laziosi, Sophie Waldvogel-Abramowski, Caroline Stephan, Arnaud G. L'Huillier, Claire-Anne Siegrist, Arnaud M Didierlaurent, Laurent Kaiser, Benjamin Meyer, Isabella Eckerle for the characterization neutralization of Omicron by patient sera.

Erik Boehm, Marc Friedli, Pauline Vetter, Samuel Cordey, Richard Neher, Christian Althaus, Emma Hodcroft, Tanja Stadler, Philippe Lemercier, Ioannis Xenarios, Lorenzo Cerutti, Louis Du Plessis, Silvan Heeb, Anna Fesser, Laurent Kaiser, for the Swiss national SARS-CoV-2 genomic and variants surveillance program coordination committee.

### **Appendix:**

#### SARS-CoV-2 epidemiology in Switzerland:

We used publicly available data on COVID-19 as reported by FOPH (<a href="https://www.covid19.admin.ch">https://www.covid19.admin.ch</a>) and sequence data submitted to GISAID to provide a summary of the SARS-CoV-2 epidemiology in Switzerland. Data presented in this table specifically cover the period from 1 August to 28 August 2022.



sup\_table\_overview\_ Aug.xlsx

<u>Supplementary Table 1:</u> Epidemiological data for Switzerland, its regions and cantons for August: population, number and incidence of confirmed cases, effective reproduction number  $R_e$ , number and incidence of tests, test positivity, number and proportion of sequenced samples, and number and proportion of VOCs.  $R_e$  by region is represented as the median and range of the daily  $R_e$  values for all cantons within a region.

week	Date	Total PCR tests	Positive tests	Sequenced	% positives	% positives sequenced
31	Aug. 1 to Aug. 7	15 812	6 008	540	38.00%	8.99%
32	Aug. 8 to Aug. 14	16 048	5 889	453	36.70%	7.69%
33	Aug. 15 to Aug. 21	14 884	5 139	560	34.53%	10.90%
34	Aug. 22 to Aug. 28	14 282	4 745	377	33.22%	7.95%
	Total	61 026	21 781	1 930	35.69%	8.86%

<u>Supplementary Table 2:</u> Total number of tests performed by the laboratories participating in the surveillance program from 1 August to 28 August 2022.

		Basic Surveillance			Augmented Surveillance					Sentinella Laboratories				
Week	Date	EOC	Labor Team W	Risch	SRO	USB	IFIK	Diana labs	CHUV	UZH	ICH- VS*	HUG	ETH/ Viollier	All
31	Aug. 1 to Aug. 7	20	21	58	13	39	<b>57</b>	57 46	34	92	43	134	29	540
32	Aug. 8 to Aug. 14	30	21	90	6	39	5/		32	94			95	453
33	Aug. 15 to Aug. 21	25	25	25	8	38	73	45	37	50	46	137	96	560
34	Aug. 22 to Aug. 28	18	24	52	11	40	/3	43	35	75	40	137	77	377
·	Total	93	225	38	38	156	130	91	138	311	89	271	297	1930

<u>Supplementary Table 3:</u> number of sequences submitted to GISAID by each laboratory during the surveilled period (1 August to 28 August 2022). \*including sequencing sent to high-throughput platforms.

## Contact list as of 26.09.22:

Coordination committee mailing list				
Name	e-mail address			
Laurent Kaiser	<u>Laurent.Kaiser@hcuge.ch</u>			
Samuel Cordey	Samuel.Cordey@hcuge.ch			
Marc Friedli	marc.friedli@epfl.ch			
Richard Neher	richard.neher@unibas.ch			
Tanja Stadler	tanja.stadler@bsse.ethz.ch			
Louis Du Plessis	louis.duplessis@bsse.ethz.ch			
Emma Hodcroft	emma.hodcroft@ispm.unibe.ch			
Christian Althaus	christian.althaus@ispm.unibe.ch			
Ioannis Xenarios	ioannis.xenarios@unil.ch			
Philippe Le Mercier	Philippe.Lemercier@sib.swiss			
Pauline Vetter	Pauline.Vetter@hcuge.ch			
Erik Boehm	Erik.Boehm@hcuge.ch			
Lorenzo Cerutti	lorenzo.cerutti@health2030.ch			
Erik Studer	erik.studer@bag.admin.ch			
Oluwaseun Oyewole	oluwaeun.oyewole@bag.admin.ch			

Laboratories mailing list				
Laboratory	Name	e-mail address		
HUG	Laurent Kaiser	Laurent.Kaiser@hcuge.ch		
HUG	Samuel Cordey	Samuel.Cordey@hcuge.ch		
HUG	Pauline Vetter	Pauline.Vetter@hcuge.ch		
HUG	Erik Boehm	Erik.Boehm@hcuge.ch		
CHUV	Gilbert Greub	Gilbert.Greub@chuv.ch		
CHUV	Claire Bertelli	Claire.Bertelli@chuv.ch		
Universtätsspital Basel	Pascal Schlaepfer	pascal.schlaepfer@usb.ch		
Universtätsspital Basel	Karoline Leuzinger	Karoline.Leuzinger@usb.ch		
Universtätsspital Basel	Hans Hirsch	Hans.Hirsch@usb.ch		
IFIK UNIBE	Alban Ramette	alban.ramette@ifik.unibe.ch		
UZH	Alexandra Trkola	trkola.alexandra@virology.uzh.ch		
UZH	Michael Huber	huber.michael@virology.uzh.ch		
EOC Bellinzona	Gladys Martinetti Luchini	Gladys.MartinettiLucchini@eoc.ch		
Stadler group/Viollier laboratories	Louis du Plessis	louis.duplessis@bsse.ethz.ch		
Stadler group/Viollier laboratories	Tanja Stadler	tanja.stadler@bsse.ethz.ch		
Viollier laboratories	Christiane Beckmann	christiane.beckmann@viollier.ch		
Viollier laboratories	Henriette Kurth	Henriette.Kurth@viollier.ch		
Hopitaux du Valais – Institut Central	Alexis Dumoulin	Alexis.Dumoulin@hopitalvs.ch		
Dianalabs	Nadia Liassine	Nadia.liassine@dianalabs.ch		
Dianalabs	Katia Jaton	Katia.jaton@dianalabs.ch		
Dianalabs	Géraldine Jost	Geraldine.jost@dianalabs.ch		
Dianalabs (Genesupport)	Tanguy Araud	Tanguy.araud@genesupport.ch		
Laboratoire Bioanalytica	Michael Naegele	michael.naegele@bioanlytica.ch		
Laboratoire Bioanalytica	Livia Berlinger	livia.berlinger@bioanalytica.ch		
Labor Team W ag	Andreas Lindauer	andreas.lindauer@team-w.ch		
Spital Region Oberaargau	Alexander Imhof	a.imhof@sro.ch		
Laboratory Risch	Nadia Wohlwend	nadia.wohlwend@risch.ch		

BAG mailing list:				
Name	e-mail address			
Erik Studer	erik.studer@bag.admin.ch			
Oluwaseun Oyewole	oluwaeun.oyewole@bag.admin.ch			
Nadia Corazza	Nadia.Corazza@bag.admin.ch			
Anna Fesser	Anna.Fesser@bag.admin.ch			
Ursina Roder	ursina.roder@bag.admin.ch			
Lionel Rohner	lionel.rohner@bag.admin.ch			
Michael Bel	Michael.Bel@bag.admin.ch			
Urs Mayr	urs.mayr@bag.admin.ch			
Damir Perisa	Damir.Perisa@bag.admin.ch			
Katrin Schneider	katrin.schneider@bag.admin.ch			
Martine Bourqui	Martine.Bourqui@bag.admin.ch			
Natalia Krempaska	natalia.krempaska@bag.admin.ch			
Selina Schwegler	Selina.schwegler@bag.admin.ch			
Mirjam Mäusezahl	Mirjam.Mäusezahl@bag.admin.ch			
Tobias Schuster	tobias.schuster@bag.admin.ch			

Sequencing centers:					
Center	Name	e-mail address			
Health 2030 Genome Center	Keith Harshman	keith.harshman@health2030.ch			
Health 2030 Genome Center	Ioannis Xenarios	ioannis.xenarios@health2030.ch			
Genomics Facility Basel-ETH Zurich	Christian Beisel	christian.beisel@bsse.ethz.ch			
The Functional Genomics Center Zurich (FGCZ)- ETHZ and UZH	Ralph Schlapbach	ralph.schlapbach@fgcz.ethz.ch			

Wastewater surveillance program mailing list:				
Name	e-mail address			
Niko Beerenwinkel	niko.beerenwinkel@bsse.ethz.ch			
David Dreifuss	david.dreifuss@bsse.ethz.ch			

Immunological characterization program of the variant mailing list:			
Name e-mail address			
Priscilla Turelli	priscilla.turelli@epfl.ch		
Didier Trono	didier.trono@epfl.ch		