

Respiratory Viruses in Luxembourg (ReViLux)

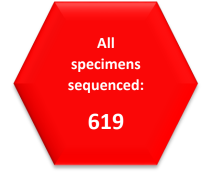
Weekly report (4 - 10 July 2022)

Executive summary

The sentinel network reported 5.2% consultations for influenza-like illness, thus exceeding the baseline circulation threshold, according to the European Centre for Disease Prevention and Control (ECDC). Within the specimens collected by the sentinel network, SARS-CoV-2 was the most frequently detected virus over the last four weeks (52.1%), followed by Human rhinovirus (22.5%) and Parainfluenzavirus (11.3%).

Regarding the SARS-CoV-2 genomic surveillance, LNS sequenced 565 specimens from residents in Luxembourg in week 27 (of 5616 total cases in the Grand Duchy of Luxembourg; 10.1%). This exceeds the ECDC recommendations to detect emerging variants.

The Omicron variant remains the only one detected in the representative sample. The Omicron BA.5 lineage is the most frequent one (89.2%, confidence interval: 86.4 - 92.0%), followed by Omicron BA.4 (5.6%, confidence interval: 3.5 - 7.7%).







Introduction

The Laboratoire national de santé, as **National Reference Laboratory for Acute Respiratory Infections in Luxembourg**, performs close surveillance on respiratory viruses, with a special focus on SARS-CoV-2. There are currently two active projects:

- The Sentinel Surveillance Network. It provides a broad picture of respiratory diseases affecting the Luxembourgish population, based on its double monitoring system (syndromic and virological).
- The National SARS-CoV-2 Genomic Surveillance Program. It enables detailed observation of SARS-CoV-2 mutations and variants through time and space, and also monitoring specific groups of interest.

The objective of the ReViLux is to inform public health actions in Luxembourg.

Sentinel Surveillance Network

The **Sentinel Surveillance Network** aims at monitoring the circulating respiratory viruses, including SARS-CoV-2, and hence underpin public health actions. Following the World Health Organization (WHO) and European Centre for Disease Prevention and Control (ECDC) guidance, it focuses on cases of acute respiratory infection (ARI) and influenza-like illness (ILI).

The season 2021/2022 ended by week 20, and it is currently the interseason period. The history of ILI consultations is displayed in Figure 1, and a detailed summary of the number of ARI and ILI cases during the last four weeks is included in Table 1. In the week of study, 5.2% of consultations were reported as ILI, thus exceeding the threshold for baseline circulation during the interseason, according to ECDC (1.14%).

Table 1. Syndromic surveillance over the last 4 weeks

Week •	ARI		ILI		Total	
	N	%	N	%	consultations	
2022/24	55	15.80	11	3.16	348	
2022/25	54	14.17	10	2.62	381	
2022/26	46	12.47	17	4.61	369	
2022/27	50	15.34	17	5.21	326	

ARI: Acute Respiratory Infections; ILI: Influenza-Like Illness.

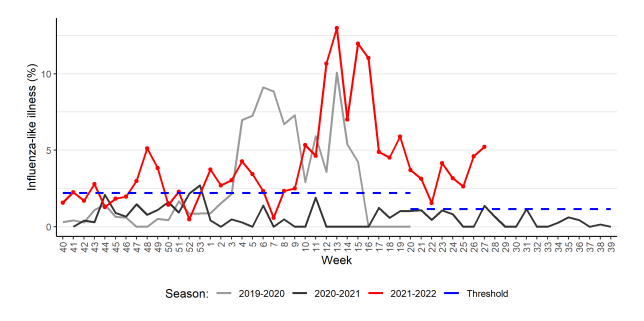


Figure 1. Percentage of patients with influenza-like illness over the last three seasons

Additionally, a selection of sentinel cases is further studied in order to monitor the circulation of respiratory viruses in the country, as shown in Figure 2. Over the last 4 weeks, the positivity rate was at 61.5%, and the most frequently detected viruses were SARS-CoV-2 (52.1%), Human rhinovirus (22.5%) and Parainfluenzavirus (11.3%). Co-infections were detected in 3 specimens, including 1 with SARS-CoV-2. These results are displayed more in detail in Table 2.

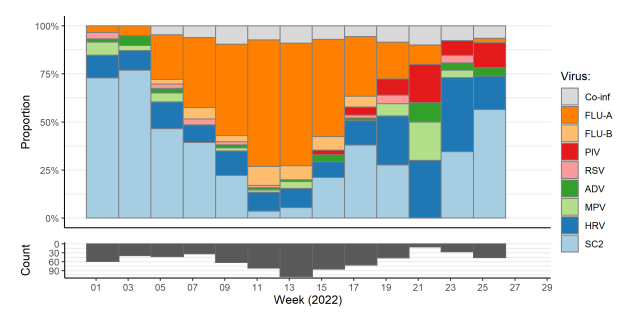


Figure 2. Distribution of respiratory viruses detected within the Sentinel Network, by two-week periods. Results from last weeks are not yet consolidated.

Co-inf: co-infection; FLU-A: influenzavirus A; FLU-B: influenzavirus B; PIV: parapneumovirus; RSV: respiratory syncytial virus; ADV: adenovirus; MPV: metapneumovirus; HRV: human rhinovirus; SC2: SARS-CoV-2.

Table 2. Distribution of respiratory viruses detected within the Sentinel Network over the last 4 weeks and during the interseason period.

Vince	Last 4	4 weeks	Interseason	
Virus	N*	%	N*	%
SARS-CoV-2	37	52.1	45	49.5
Human rhinovirus	16	22.5	24	26.4
Parainfluenzavirus	8	11.3	9	9.9
Adenovirus	5	7.0	6	6.6
Respiratory syncytial virus	2	2.8	3	3.3
Metapneumovirus	2	2.8	3	3.3
Influenzavirus A	1	1.4	1	1.1
Influenzavirus B	0	0.0	0	0.0
Total	71	100.0	91	100.0

Co-infection cases counted once for each virus detected.

SARS-CoV-2 Genomic Surveillance

The current sequencing strategy

The National Reference Laboratory for Acute Respiratory Infections at LNS receives SARS-CoV-2 positive samples (nasopharyngeal or oropharyngeal swabs analysed by RT-PCR) from the national network of laboratories and proceeds as follows:

- 1. Sequencing a representative sample of specimens.
- 2. Sequencing specimens from target groups (i.e. hospital cases and post-vaccination cases).
- 3. Sequencing specimens from clusters with high transmission.

The representative sample of specimens is a systematic selection from all SARS-CoV-2 positive cases registered in Luxembourg to detect emerging variants and early increases in their incidence and transmission within the community in Luxembourg. This sample is selected according to the ECDC guidelines.

A screening by targeted PCR tests is also carried out when a new variant emerges in order to enable earlier evaluation of its spread in the population.

SARS-CoV-2 lineages have been assigned based on Rambaut et al. using the Phylogenetic Assignment of Named Global Outbreak LINeages (pangolin) software (4.1.1, designation v1.11). The Pango nomenclature is used in addition to the WHO nomenclature to enable easier visualization of links between any evolving variants and their ancestor.

Specimens 619 specimens 565 specimens from national cases 10.1% sequencing coverage

Figure 3. Flowchart of specimens collected during week 27/2022

In week 27, 5616 new cases were registered in Luxembourg; hence, the minimum sample size required to detect emerging variants at a 2.5% incidence is estimated to be 542 specimens (9.7%).

As shown in Figure 3, the microbial genomics unit at the LNS sequenced 619 specimens from the week of study, including 565 national ones. The weekly sequencing coverage remains at 10.1% (out of 5616 cases registered in Luxembourg; see coverage trend in Figure 4), which exceeds the recommended sample size.

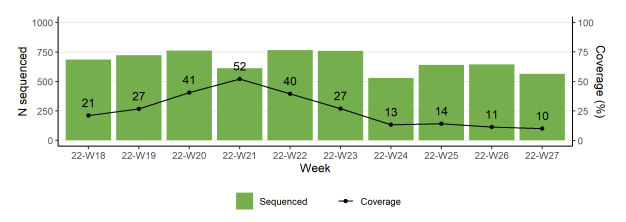


Figure 4. Number of specimens sequenced and national coverage based on weekly number of positive cases in Luxembourg. The coverage from the last two weeks is not consolidated yet.

Circulating lineages detection

The distribution of successfully assigned lineages within the national selection is shown in Figure 6, and it is further detailed in Table 4 (last two weeks). A summary of the VOCs assigned among all specimens sequenced (including non-residents) during the last two weeks and since the beginning of the sequencing activity is shown in Table 5.

The Omicron variant remains the only one detected within the representative sample, the most frequent lineage being Omicron BA.5 (89.2%), followed by Omicron BA.4 (5.6%).

Table 4. Distribution of SARS-CoV-2 lineages detected within the representative sample during the last two weeks. Previously reported cases might be updated by retrospective analysis.

Linoago		Previous	week	Current week			
Lineage	N	% CI %		N	%	CI %	
Omicron BA.5	457	84.5	81.4 - 87.5	430	89.2	86.4 - 92.0	
Omicron BA.4	38	7.0	4.9 - 9.2	27	5.6	3.5 - 7.7	
Omicron BA.2	46	8.5	6.2 - 10.9	25	5.2	3.2 - 7.2	
Total	541	100.0	-	482	100.0	-	

CI: Confidence Interval at 95%.

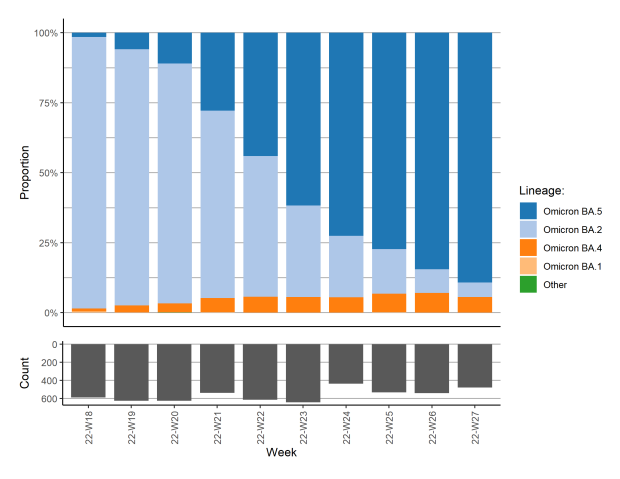


Figure 6. Distribution of lineages within the representative sample during the last 10 weeks.

	Previous week			nt week	·	
Variant -	N	%	N	%	Cumulative count	
Omicron	600	100.0	519	100.0	20 028	
BA.5	218	36.3	227	43.7	1 638	
BA.5.1	88	14.7	37	7.1	280	
BA.2.12.1	10	1.7	6	1.2	220	
BA.4	25	4.2	18	3.5	173	
BA.2.13	0	0.0	0	0.0	43	
BA.2.11	0	0.0	0	0.0	2	
BA.2.75	0	0.0	0	0.0	1	
Delta	0	0.0	0	0.0	13 679	
Others	0	0.0	0	0.0	15 005	
Total	600	100.0	519	100.0	48 712	

Clinical and epidemiological factors

In this section, the lineage distribution of all specimens sequenced over the last month is assessed by demographics (sex and age group, Figure 7) and sampling setting (community vs. hospital, Table 6).

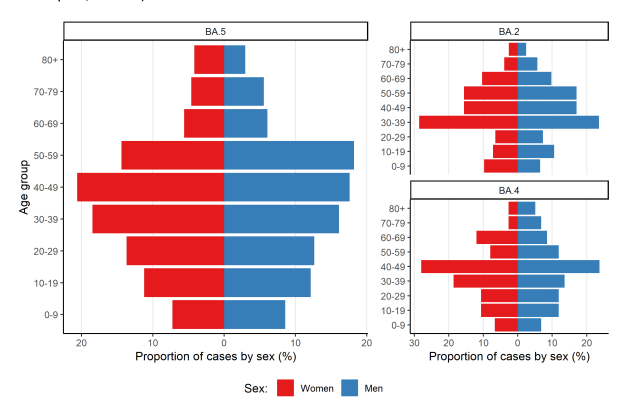


Figure 7. Age group and sex distribution of specimens sequenced over the last 4 weeks, by lineage.

Table 6. Comparison of lineage distribution by sampling setting.

Lineage		Hospital				
	Women	Men	Total	Women	Men	Total
Omicron BA.5	77.1%	80.9%	78.9%	84.7%	83.1%	84.0%
Omicron BA.2	12.1%	12.4%	12.2%	10.6%	11.3%	10.9%
Omicron BA.4	10.8%	6.7%	8.8%	4.7%	5.6%	5.1%
Total	100.0%	100.0%	100.0%	100.0%	100.0%	100.0%

Mutation surveillance

In addition to the surveillance of SARS-CoV-2 variants, the LNS monitors the occurrence of SARS-CoV-2 mutations reported to have a clinical and epidemiological relevance. Newly acquired mutations may occur and their early detection might be key to expect changes in the epidemic evolution. As each VOC is characterised by a set of defining mutations, which are expected to be highly present, it is interesting to analyse the non-defining ones.

Among the specimens collected over the last four weeks, we did not detect any frequent non-defining mutation (present in at least 1% of any VOC).

References

Centers for Disease Control and Prevention. SARS-CoV-2 Variant Classifications and Definitions. Retrieved 18 July 2022, from https://www.cdc.gov/coronavirus/2019-ncov/variants/variant-info.html

COVID-19 Data Portal - accelerating scientific research through data. (2021). Retrieved 18 July 2022, from https://www.covid19dataportal.org/sequences

Direção-Geral da Saúde. Relatório de Situação. Retrieved 18 July 2022, from https://covid19.min-saude.pt/

European Centre for Disease Prevention and Control. Guidance for representative and targeted genomic SARS-CoV-2 monitoring – 3 May 2021. ECDC: Stockholm; 2021

European Centre for Disease Prevention and Control. SARS-CoV-2 variants of concern. Retrieved 18 July 2022, from https://www.ecdc.europa.eu/en/covid-19/variants-concern

Genomic sequencing of SARS-CoV-2: a guide to implementation for maximum impact on public health. Geneva: World Health Organization; 2021.

GISAID. EpiCoV — Pandemic coronavirus causing COVID-19. Retrieved 5 mai 2022, from https://www.gisaid.org

GitHub - cov-lineages/pangolin: Software package for assigning SARS-CoV-2 genome sequences to global lineages. (2021). Retrieved 18 July 2022, from https://github.com/cov-lineages/pangolin

Hadfield J., Megill C., Bell S., Huddleston J., Potter B., Callender C. et al. (2018). Nextstrain: real-time tracking of pathogen evolution. Bioinformatics, 34(23), 4121-4123. doi: 10.1093/bioinformatics/bty407

Instituto Nacional de Saúde Doutor Ricardo Jorge. Diversidade genética do novo coronavírus SARS-CoV-2 (COVID-19) em Portugal. Retrieved 18 July 2022, from https://insaflu.insa.pt/covid19/

Rambaut A., Holmes E., O'Toole Á., Hill V., McCrone J., Ruis C. et al. (2020). A dynamic nomenclature proposal for SARS-CoV-2 lineages to assist genomic epidemiology. Nature Microbiology, 5(11), 1403-1407. doi: 10.1038/s41564-020-0770-5

Robert Koch Institut. Aktueller Lage-/Situationsbericht des RKI zu COVID-19. Retrieved 18 July 2022, from

https://www.rki.de/DE/Content/InfAZ/N/Neuartiges_Coronavirus/Situationsberichte/Gesamt.html;jsessionid=69BC18053F9591C56EB148E463103DB7.internet101

Santé publique France. Coronavirus : chiffres clés et évolution de la COVID-19 en France et dans le Monde. Retrieved 18 July 2022, from https://www.santepubliquefrance.fr/dossiers/coronavirus-covid-19/coronavirus-chiffres-cles-et-evolution-de-la-covid-19-en-france-et-dans-le-monde

Sciensano. COVID-19 – Bulletin épidémiologique hebdomadaire. Retrieved 18 July 2022, from https://covid-19.sciensano.be/fr/covid-19-situation-epidemiologique

The Luxembourgish data platform. COVID-19: Rapports hebdomadaires. Retrieved 18 July 2022, from https://data.public.lu/en/datasets/covid-19-rapports-hebdomadaires/