

Respiratory Viruses in Luxembourg (ReViLux)

Weekly report - 28 March 2023

Executive summary

The sentinel network detected a moderate epidemic activity over the last week (2023/12), based on 7.6% of consultations being associated with influenza-like illness. Among the specimen collected by the sentinel network over the last two weeks, the percentage of positive tests for Influenza type B was 34.8%, followed by SARS-CoV-2 (13.3%) and Human rhinovirus (11.0%). Overall, during this season, Influenza A (H3) with 89.4% is the most frequent detected strain in submitted specimens. However, in recent weeks, increased circulation of Influenza A (H1) pdm and Influenza B have been observed compared to H3.

Concerning SARS-CoV-2 genomic surveillance, the recombinant sub-variant XBB.1.5 has been the most frequent detected variant for the past few weeks with 50% (95%CI: 43.8 - 56.2%), followed by XBB.1.9 with 24.6% (95%CI: 19.3 - 29.9%).

IN THE SPOTLIGHT

XBB.1.5 like sub-lineages

XBB.1.5 is a subtype with additional spike RBD mutation F486P. The lineage was first detected in October 2022 in the USA and is estimated to have a large growth advantage to other previously circulating lineages. Currently, there are no signals that the infection severity of XBB.1.5 is different from that of other circulating Omicron sub-lineages. ECDC categorizes lineages that have similar Spike proteins as **VOI XBB.1.5 lineages** (XBB.1.5-like), which include XBB.1.5, XBB.1.9.1*, XBB.1.9.2* and XBB.1.16.

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. The main monitoring activities are:

- The Sentinel Surveillance. It provides a broad picture of respiratory diseases affecting the Luxembourgish population, based on clinical and laboratory data.
- The SARS-CoV-2 Genomic Surveillance. It enables detailed observation of SARS-CoV-2 variants and mutations through time and space, as well as studying specific groups of interest.

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

Sentinel Surveillance Network

The Sentinel Surveillance aims at monitoring the circulating respiratory viruses, from traditional ones like Influenza to more recent ones like SARS-CoV-2, and hence underpin public health actions. The Sentinel Network is a group of general practitioners and paediatricians spread over the country. They report the weekly number of patients showing symptoms suggestive of acute respiratory infection (ARI) and influenza-like illness (ILI), and those patients are then sampled and tested for a panel of respiratory viruses. The circulation of respiratory viruses in the north hemisphere is generally monitored by seasons that go from week 40 to week 20. The period between weeks 20 and 40 is usually called interseason.

Clinical results

Last week, 7.6% of consultations were reported as ILI, which represents a moderate epidemic activity for Luxembourg, according to ECDC and the Moving Epidemic Method. 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.

Laboratory results

Over the last two weeks, the most frequently detected viruses (according to positivity rates) were Influenzavirus B (34.8%), SARS-CoV-2 (13.3%) and Human rhinovirus (11%). An overview of the viral circulation during the current season is displayed in Figure 2 and Table 2.

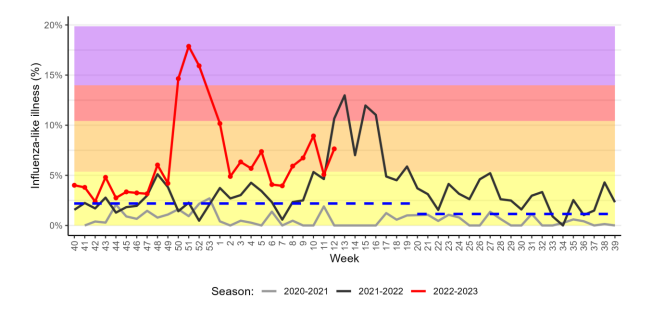


Figure 1. Percentage of patients with influenza-like illness over the last three seasons Background colours according to intensity of circulation: low, medium, high, very high.

Table 1. Syndromic surveillance over the last 4 weeks

Week -	ARI		ILI		Total
	N	%	N	%	consultations
2023/9	79	20.47	26	6.74	386
2023/10	56	16.67	30	8.93	336
2023/11	60	16.04	19	5.08	374
2023/12	54	17.94	23	7.64	301

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

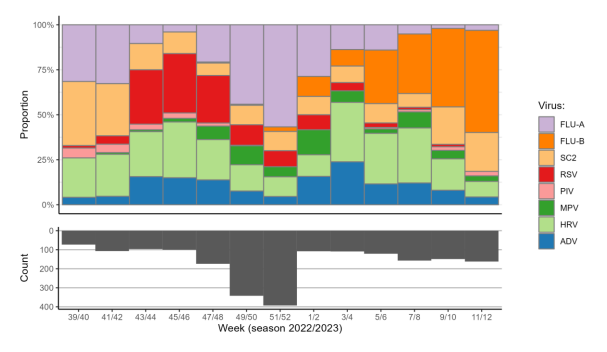


Figure 2. Distribution of respiratory viruses detected within the Sentinel Network, by two-week periods. Results from last weeks are not yet consolidated.

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 2 weeks compared to the whole current season.

Vince	Last 2 weeks			Season 2022/2023		
Virus	N*	Positivity rate (%)	N*	Positivity rate (%)		
Influenzavirus B	92	34.8	280	11.9		
SARS-CoV-2	35	13.3	278	17.5		
Human rhinovirus	14	11.0	391	19.0		
Adenovirus	7	5.5	200	9.7		
Metapneumovirus	5	3.9	126	6.1		
Parainfluenzavirus	4	3.1	28	1.4		
Influenzavirus A	5	1.9	404	17.2		
Respiratory syncytial virus	0	0.0	208	10.1		
Total	162		1915			

^{*}Co-infection cases counted once for each virus detected.

Genotyping analyses

In order to assess the circulation of viral strains, genotyping analyses were also performed for Influenzavirus A and Respiratory syncytial virus (RSV) (Figure 3). During the current season, the most frequently detected strain for Influenzavirus A was H3 (89.4%), and for RSV it was type B (74.7%).



Figure 3. Detection of influenzavirus-A (Flu-A) and Respiratory syncytial virus (RSV) strains in sentinel samples during the current season.

SARS-CoV-2 Genomic Surveillance

SARS-CoV-2 has posed a serious threat to the world population since 2020, and it is still a challenge for health systems. For this reason, the National Reference Laboratory for Acute Respiratory Infections at LNS continues to monitor the genomic characteristics of the SARS-CoV-2 virus detected in Luxembourg.

LNS receives positive specimens (nasopharyngeal or oropharyngeal swabs analysed by RT-PCR) from the national network of laboratories. All hospital specimens are sequenced, as well as a representative selection of community specimens. Illumina and Nanopore devices are used respectively. When needed, an additional screening by targeted PCR is also performed to enable earlier evaluation of the spread of newly emerging variants.

Bioinformatic analyses are based on a standardised pipeline, and lineage assignment is performed through the PANGOLin software (4.2, pango-data 1.18.1.1, mode UShER). In order to allow easier visualisation, WHO and Nextstrain categorisations are also used.

Sequencing activity

In week 2023/11, 1796 new cases were registered in Luxembourg; hence, the minimum sample size required to detect emerging variants at a 5% incidence is estimated to be 252 specimens (14%). The microbial genomics unit at LNS sequenced 359 specimens from the week of study, including 318 national ones (sequencing coverage: 17.7%; see Figure 4). This reaches the recommended sample size.

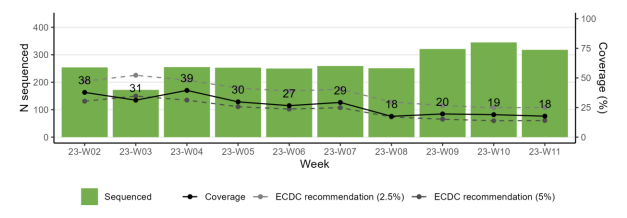


Figure 4. Number of specimens sequenced and national coverage based on weekly number of positive cases in Luxembourg. The coverage from the last two weeks is not consolidated yet.

Variant circulation

The XBB.1.5 like variant remains the main one detected within the representative sample, the most frequent subvariant being XBB.1.5 (50%), followed by XBB.1.9 (24.6%).

An overview of the variants and lineages circulating over the last weeks is displayed in Figure 5, and further details are shown in Table 3. The history of the circulation of each variant since January 2021 is displayed in Figure 6.

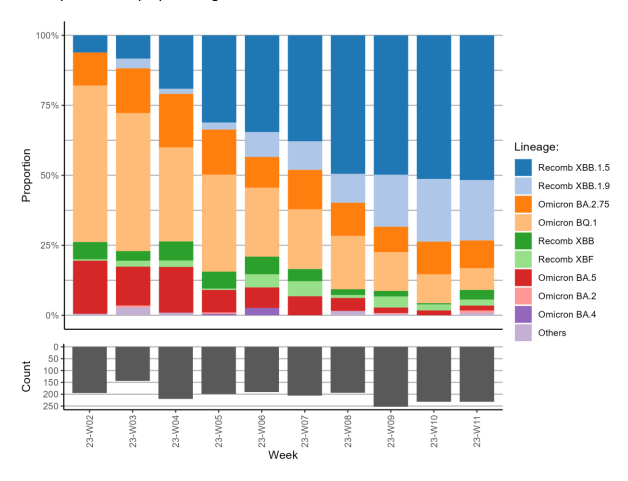


Figure 5. Distribution of lineages within the representative sample during the last 10 weeks.

Table 3. 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.

Lincoro*	V	Veek 10	Week 11		
Lineage*	%	CI %	%	CI %	
Recomb XBB.1.5	50.0	43.8 - 56.2	50.0	43.8 - 56.2	
Recomb XBB.1.9	20.2	15.3 - 25.2	24.6	19.3 - 29.9	
Omicron BQ.1	13.1	8.9 - 17.3	8.3	4.9 - 11.7	
Omicron BA.2.75	11.5	7.6 - 15.4	7.9	4.6 - 11.3	
Recomb XBF	2.8	0.7 - 4.8	2.8	0.7 - 4.8	
Recomb XBB	0.4	0.0 - 1.2	2.8	0.7 - 4.8	
Omicron BA.5	1.6	0.0 - 3.1	1.6	0.0 - 3.1	
Others	0.4	0.0 - 1.2	1.2	0.0 - 2.5	
Omicron BA.2	0.0	-	0.8	0.0 - 1.9	

^{*}Sub-lineages are not included in the count of their parental lineages (e.g. BQ.1 not included in BA.5).

CI: confidence interval at 95%.

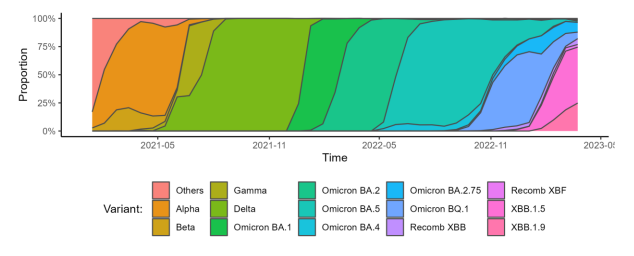


Figure 6. Proportion of each variant circulating in Luxembourg since January 2021.

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 4).

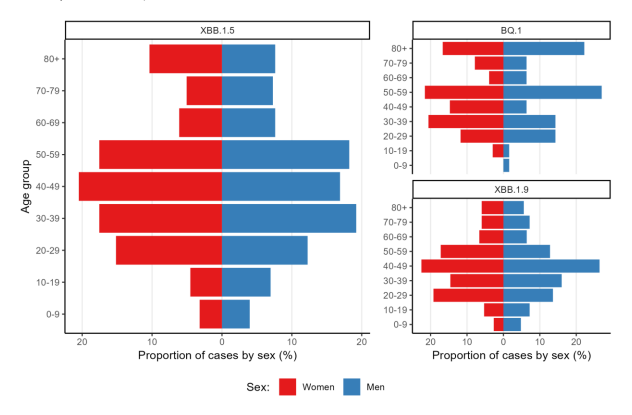


Figure 7. Age group and sex distribution of specimens sequenced over the last 4 weeks, by lineage.

Table 4. Comparison of lineage distribution by sampling setting.

Lincon	_	Hospital				
Lineage	Women	Men	Total	Women	Men	Total
Recomb XBB.1.5	61.8%	63.3%	62.5%	54.2%	56.9%	55.4%
Recomb XBB.1.9	22.1%	24.4%	23.1%	29.2%	28.5%	28.9%
Omicron BQ.1	16.1%	12.2%	14.4%	16.7%	14.6%	15.8%
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, as their early detection might be key to foresee changes in the epidemic evolution. The LNS is currently monitoring mutations to the spike protein, following ECDC guidance, and comparing their prevalence to that observed in Europe (according to GISAID).

Among the specimens collected over the last four weeks, we detected a higher circulation of the following mutations in Luxembourg (compared to Europe):

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Lineage	Mutation	Gene	% in LU	% in EU	Ratio
BQ.1	Q613H	S	14.63	5.0	2.90
BQ.1	F157L	S	26.83	9.4	2.86
BA.5	R346T	S	100.00	48.9	2.04
BA.2.75	K356T	S	18.33	11.2	1.64
BA.2.75	F490S	S	18.33	11.4	1.60
XBB	Q954H	S	100.00	81.2	1.23
XBB	N969K	S	100.00	81.9	1.22
BA.5	Q954H	S	100.00	84.4	1.18
BA.5	N969K	S	100.00	85.8	1.16
BA.2.75	Q954H	S	100.00	85.9	1.16
BQ.1	Q954H	S	100.00	86.2	1.16
BQ.1	N969K	S	100.00	87.7	1.14
BA.2.75	N211V	S	19.67	<5.00	-
BA.2.75	L212I	S	19.67	<5.00	-
BA.5	A1016S	S	33.33	<5.00	-
BA.2.75	N969K	S	100.00	<5.00	-

LU: Luxembourg; EU: Europe

References

Centers for Disease Control and Prevention. SARS-CoV-2 Variant Classifications and Definitions. Retrieved 27 March 2023, from https://www.cdc.gov/coronavirus/2019-ncov/variants/variant-info.html

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. Communicable Disease Threats Report Week https://www.ecdc.europa.eu/en/publications-data/communicable-disease-threats-report-week-12-19-25-march-2023

European Centre for Disease Prevention and Control. SARS-CoV-2 variants of concern. Retrieved 27 March 2023, 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 27 March 2023, 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

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

The Luxembourgish data platform. COVID-19: Rapports hebdomadaires. Retrieved 27 March 2023, from https://data.public.lu/en/datasets/covid-19-rapports-hebdomadaires/