

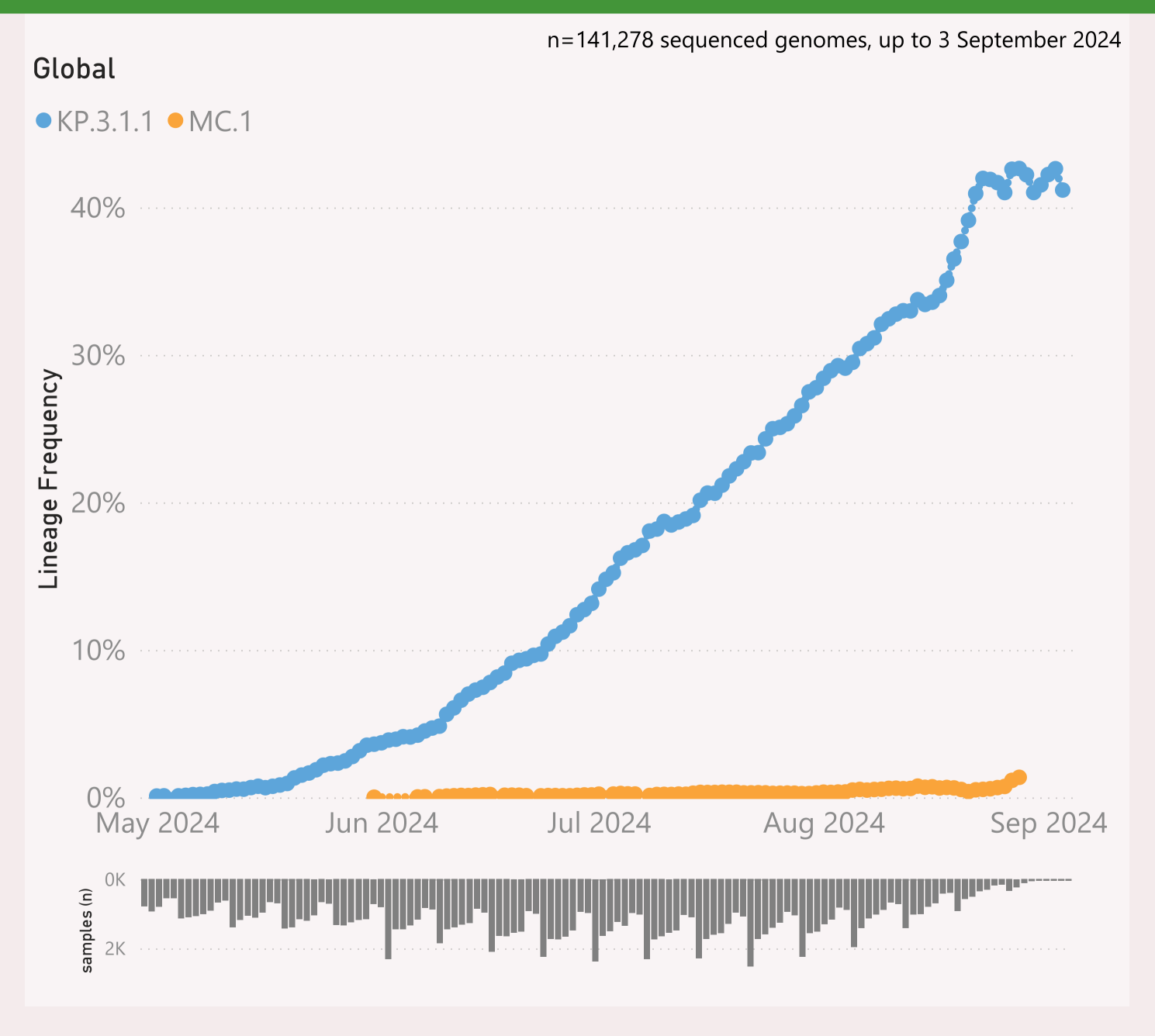
This page shows the frequency of the top 6 "L2" lineages, across recent months.

The detailed Lineage classifications are provided by Nextclade. I roll those up into "L2" groups, which roughly follow the WHO Variant definitions. For example, my "BA.2.86.\*" group includes BA.2.86 and all it's descendants, e.g. the JN.\* lineages.

The detailed Lineage classifications are quite numerous and dynamic, so the "Lineage L2" groups give a simpler and more stable basis for analysis and comparison.

The frequency shown at each point is based on the 7-day rolling average across all lineages.

The grey column chart across the bottom shows the volume of sequences available by date. As there can be long sample and data processing times, it is quite routine for recent dates to show lower sample sizes.

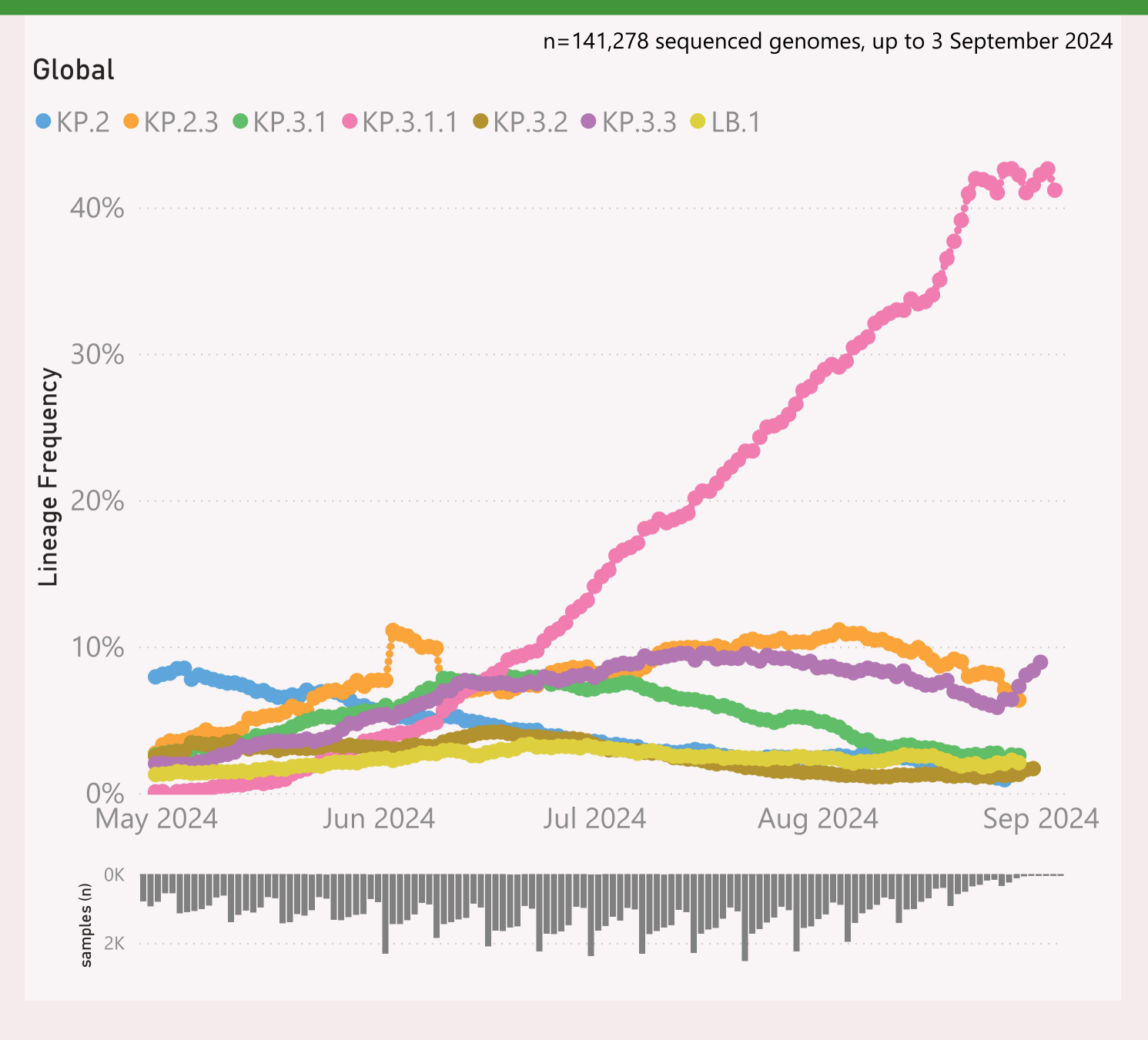


This page shows the frequency of the top 7 lineages, across recent months. The lineages are filtered for a "Lineage L2" group of interest, currently "JN.1.\* + DeFLuQE".

The Lineage classifications are provided by Nextclade. The colour assignments are random.

The frequency shown at each point is based on the 7-day rolling average across all lineages.

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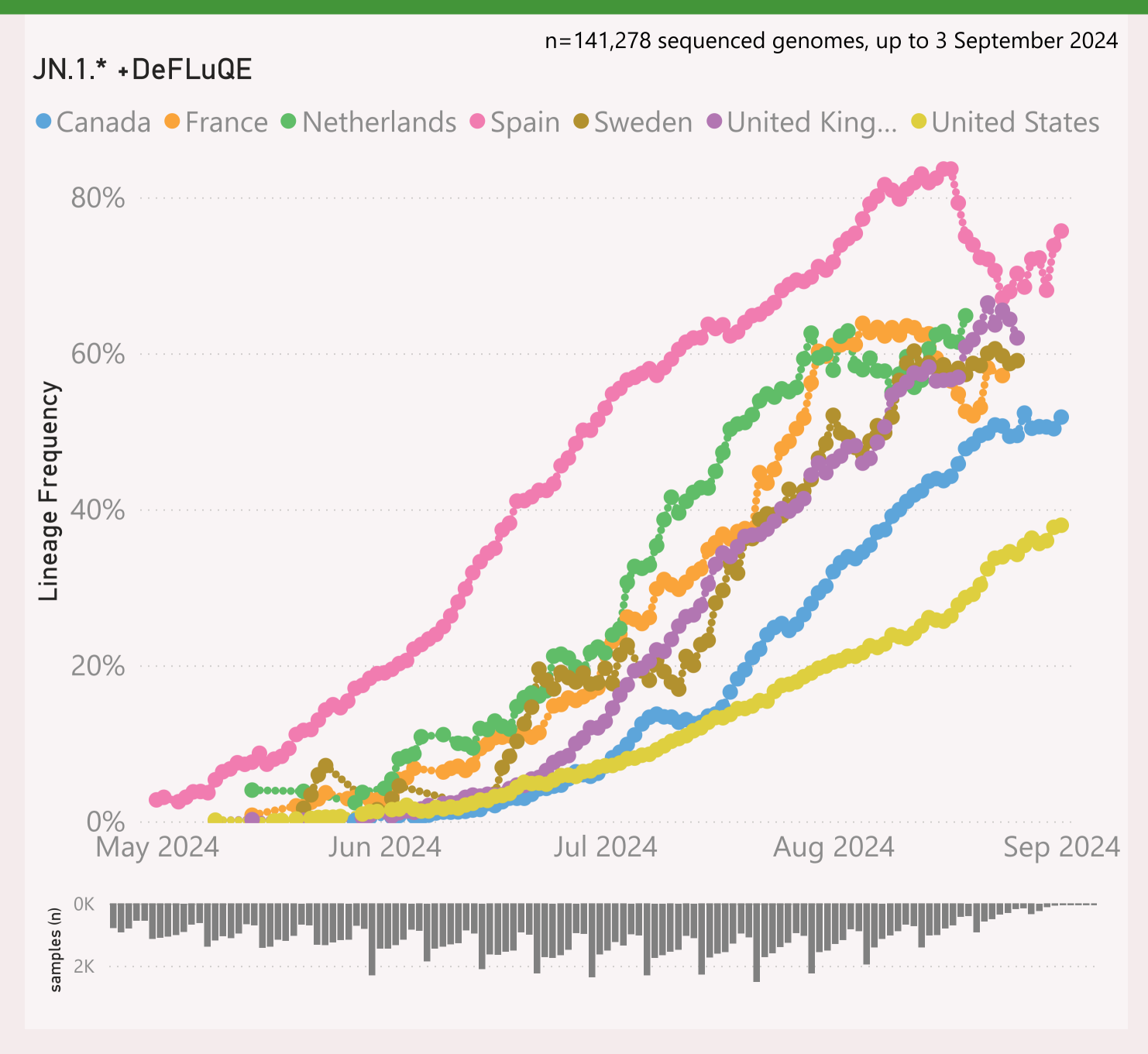


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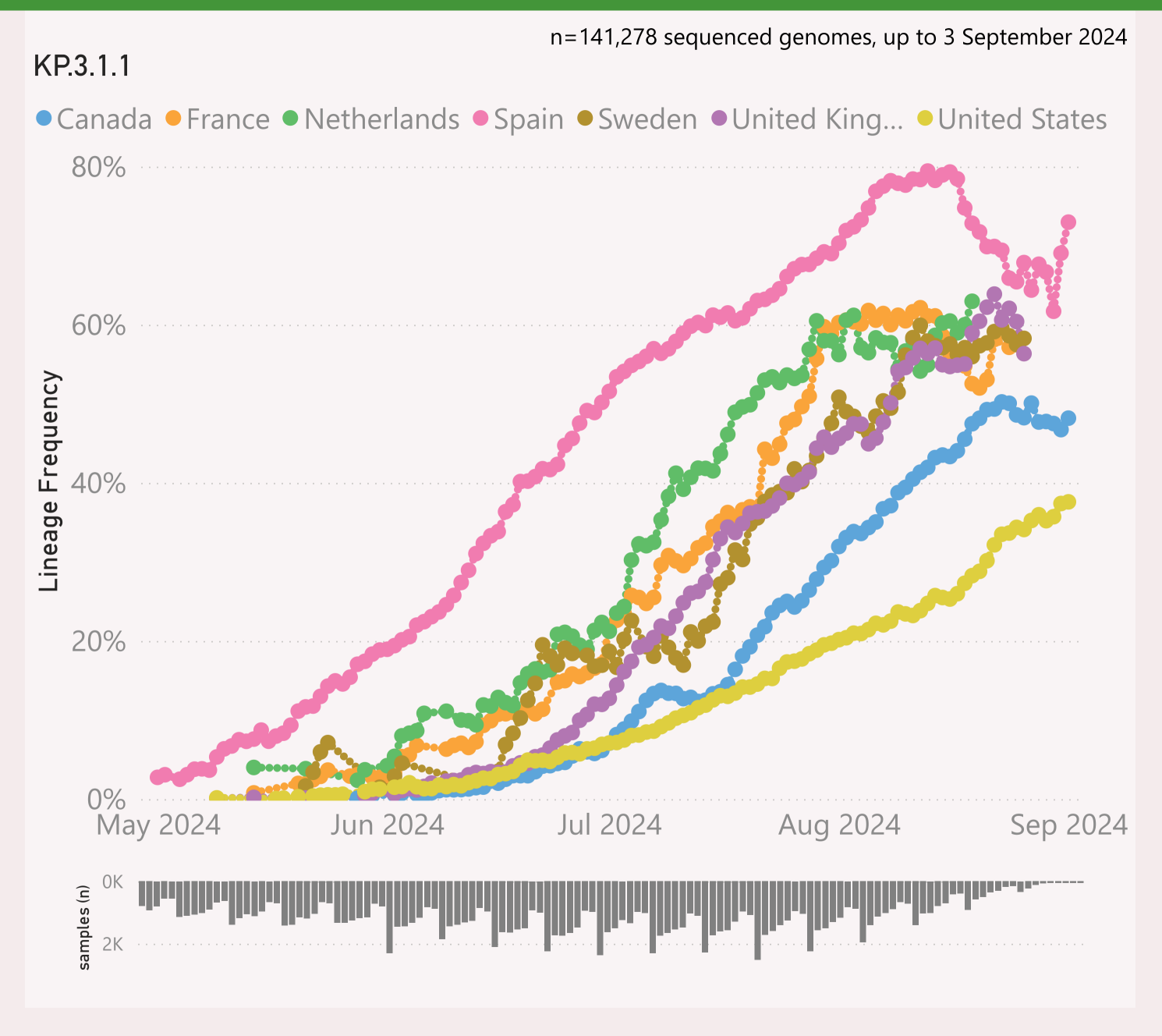
This page shows the frequency of a selected "Lineage L2" group of interest, for the 7 countries reporting the most samples over recent months.

The detailed Lineage classifications are provided by Nextclade. I roll those up into "L2" groups, which roughly follow the WHO Variant definitions. For example, my "JN.1.\* +FLiRT" group includes the descendants of JN.1.\* with the mutations: F456L & R346T.

The detailed Lineage classifications are quite numerous and dynamic, so the "Lineage L2" groups give a simpler and more stable basis for analysis and comparison.

The frequency shown at each point is based on the 7-day rolling average across all lineages, for that country.

The grey column chart across the bottom shows the volume of sequences available by date. As there can be long sample and data processing times, it is quite routine for recent dates to show lower sample sizes.



This page shows the frequency of a selected Lineage of interest, for the 7 countries reporting the most samples over recent months.

The Lineage classifications are provided by Nextclade.

The frequency shown at each point is based on the 7-day rolling average across all lineages, for that country.

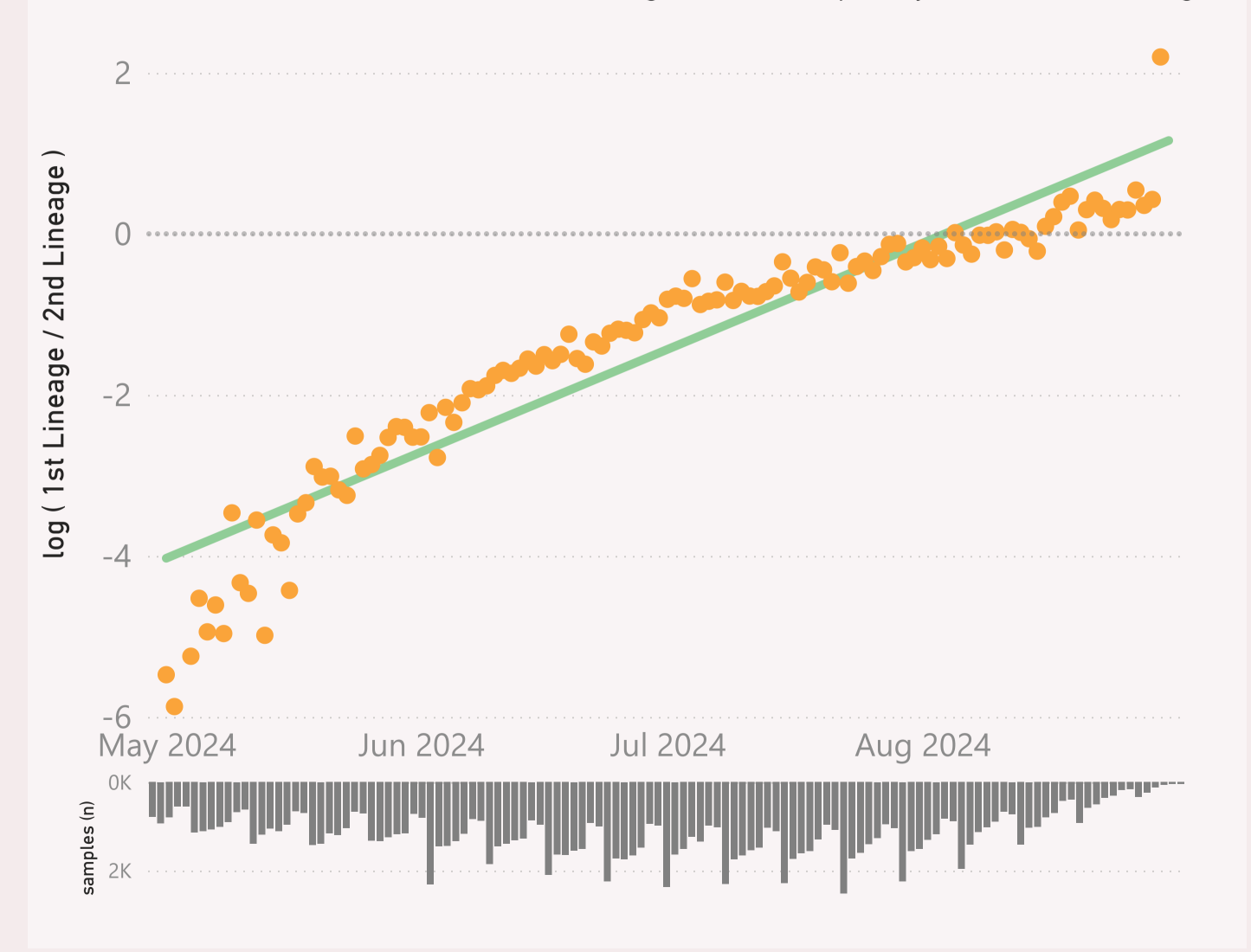
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n=141,249 sequenced genomes, up to 31 August 2024

## Global - JN.1.\* +DeFLuQE vs JN.1.\* +FLiRT

● log (1st Lineage / 2nd Lineage ) ● trend

growth of 4.3% per day, crossover on 04-Aug-24

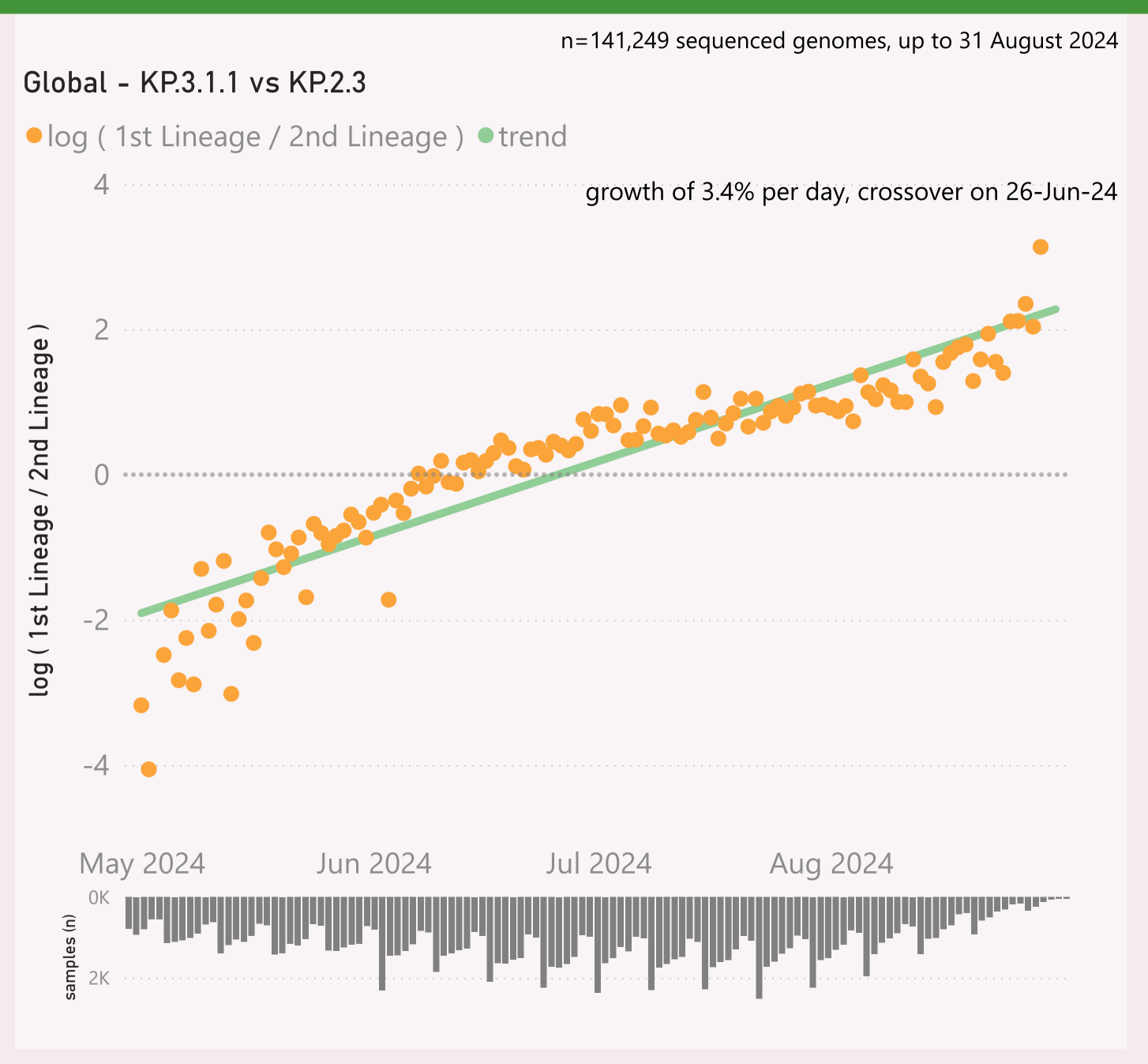


This page compares the relative frequency of 2 selected "Lineage L2" groups, over recent months. A challenging Lineage L2 is selected first, and compared to the incumbent.

The trend is shown as a green line and expressed as a daily growth % advantage. If the green line crosses over the 0.0 line, the date when that occurred or is predicted to occur will be shown. At that point the challenging Lineage L2 is considered to have "crossed over" or taken over dominance from the incumbent Lineage L2.

The Lineage classifications are provided by Nextclade. I add the "Lineage L2" groups, typically following common variant groupings, but occasionally being "creative".

The grey column chart across the bottom shows the volume of sequences available by date. As there can be long sample and data processing times, it is quite routine for recent dates to show lower sample sizes.

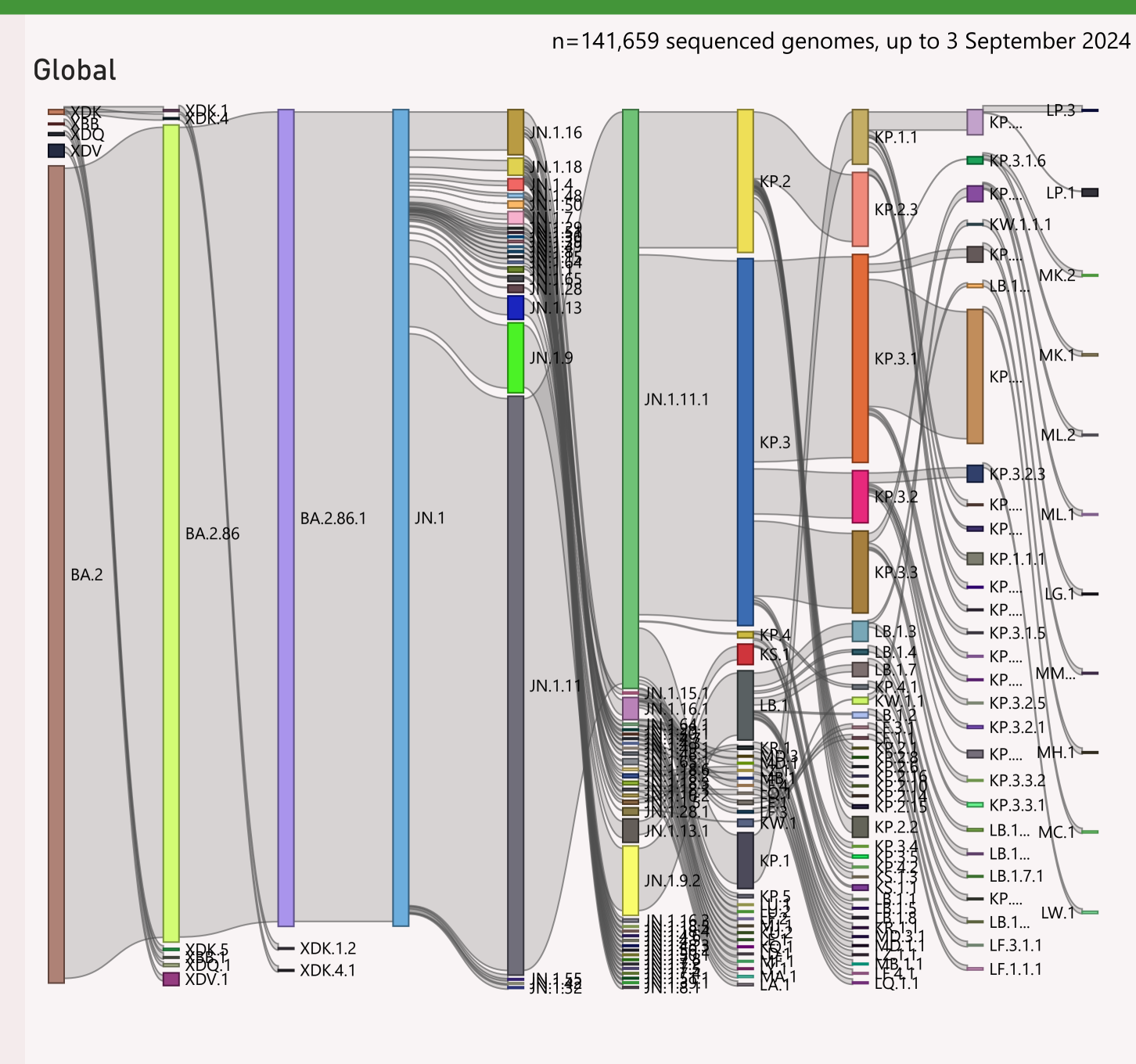


This page compares the relative frequency of 2 selected Lineages, over recent months. A challenging Lineage is selected first, and compared to the incumbent.

The trend is shown as a green line and expressed as a daily growth % advantage. If the green line crosses over the 0.0 line, the date when that occurred or is predicted to occur will be shown. At that point the challenging Lineage is considered to have "crossed over" or taken over dominance from the incumbent Lineage

The Lineage classifications are provided by Nextclade.

The grey column chart across the bottom shows the volume of sequences available by date. As there can be long sample and data processing times, it is quite routine for recent dates to show lower sample sizes.



This page shows the hierarchy of the significant Lineages, over recent months.

The hierarchy can be read from left to right, starting with the earliest/highest Lineages being broken down into more detailed child Lineages.

The vertical height of each bar segment represents the relative volume of all the samples of that specific Lineage, as well as all it's descendants.

The full picture is typically quite busy, so insignificant Lineages (with few samples, or at the extreme top or bottom of the hierarchy) are not shown.

The Lineage classifications are provided by Nextclade.

## Data Submitted in the last 8 weeks

Country	# Samples Sequenced	Latest Collection date	by Collection date	Latest Submission date	by Submission date
United States	34,169	9/2/2024		9/4/2024	au nit nal alli rataalii kira i
	9,340	9/2/2024		9/4/2024	na la basta mota da di
	5,816	8/27/2024		9/4/2024	alti lakitin saat
⊞ Japan	5,805	9/1/2024		9/4/2024	and a property like the of
⊞ Spain	5,691	9/1/2024		9/4/2024	الطور ووورا المراطأت
	3,346	8/24/2024	labibit	9/4/2024	addithed by
⊕ China	2,108	8/30/2024	الأليانية	9/4/2024	وأراء ويستجارها والمارية
	1,973	8/19/2024	عالله	9/2/2024	ير الرياري
⊕ Australia	1,897	8/25/2024	ditalig.	9/4/2024	المناب ومملك المما
	1,711	8/26/2024	. أألله من الأسلام المناطقة ا	9/3/2024	and the second
⊞ New Zealand	1,670	8/25/2024		9/2/2024	- L
<b>H</b> Germany	1,609	8/23/2024		9/4/2024	الليانا والمارسرانية
⊕ Denmark	1,511	8/26/2024	ال ب	9/4/2024	agha a ha ca
	1,327	9/3/2024	ع برانالله د	9/4/2024	المناهم والمنطم المطاعد
⊞ South Korea	1,315	8/29/2024		9/4/2024	1111
	1,109	8/30/2024	والتراباليون والمرابع	9/4/2024	ages to distinct and
⊕ Brazil	896	8/24/2024	عماناه بنشاء الألب	9/4/2024	والمنتس الإسطا
⊕ Russia	883	8/6/2024	alt - dalidijla i	8/19/2024	
<b>E</b> Singapore	866	8/28/2024	Hytaa	9/2/2024	1, 1,
∃ Israel	853	8/17/2024	, allian	8/22/2024	Junu co
⊕ Puerto Rico	690	8/10/2024	يد عاملان	9/3/2024	Jahren der
Portugal	602	7/30/2024		8/12/2024	
	525	8/20/2024	lataran taadiitiga n	8/30/2024	ا بات
⊕ Guatemala	394	8/13/2024	التأوات.	9/2/2024	
⊞ India	378	8/4/2024	. daan ridgalee	8/29/2024	
⊕ Belgium	368	8/18/2024	. In a lagnik	9/2/2024	radional district
⊕ Switzerland	343	8/4/2024		9/4/2024	.11 1.1
	336	7/29/2024	. atrodaļait	9/2/2024	
Total	91,740	9/3/2024		9/4/2024	nn.hat.lb.l.dbl.bat.atba.bnt.a

This page shows the volume and currency/timeliness of the genomic sequencing data shared via GISAID, over the last 8 weeks, for the countries sharing the most samples.

Each sample shared comes with a Collection date - when the PCR test for that sample was collected. The GISAID system also records a Submission date for each sample, which is typically the date that sample was uploaded.

The latest date of each type is shown, along with "sparkline"-style mini charts to give a flavour for the spread of recent data by Collection date and by Submission date.