### Delta variants, C.1.2, Mu, P.1

Sept. 11, 2021 GISAID data feed provided to cov.lanl.gov Bette Korber and James Theiler

#### "Delta variant" variants

We have highlighted 52 variants of Delta that seemed of particular interest as they had at least one or a combination of the following features:

- Common
- Consistently increasing in frequency compared to baseline Delta in multiple geographic regions
  - Slide 3 shows a mutation that shows such potential for positive selection
  - Slide 4 shows a counter example, a common mutation that is not consistently increasing, so its relatively high sampling is more likely to be founder effects or sampling artifacts.
  - Many of the variants listed are just found in a small number of countries, and will continue to monitor them.
- In an interesting region: the RBD, NTDss, or potentially related to furin cleavage

Note: Site Spike 142 in Delta. As a cautionary note: It has become apparent through various analyses that the G142 ancestral mutation of the spike in recent lineages, such as Delta, could be an artefact of sequencing protocols. Almost every new mutation that is common in Delta (found over 300 times in the 5/26/2021 – 8/24/2021 GISAID sample from 8/24/2021) comes in two flavors of Spike, some with the G142D mutation, and some with the ancestral G142 in this position. A slide illustrating examples of this is shown in slide 5. Given this, we are at least for the present, "setting aside" the G142 ancestral versions of Delta variants from our list, and unless a mutational variant is reported with only the ancestral form of G142 associated with it, we choose the G142D version of the variant Spike as the preferred form of the Delta variant.

In our current counts for a variant forms in Delta, we do not split the data by 142 status, and so do not risk undercounting new variant forms in our tallies. For example: in the last 60 days we find the form of Spike with this basic Delta backbone pattern: T19R,**G142D**,E156G,F157-,R158-,L452R,T478K,D614G,P681R,D950N exactly 63571 times, and the form of Spike without G142D: T19R,E156G,F157-,R158-,L452R,T478K,D614G,P681R,D950N exactly 7467 times, and sp the count of the basic form to be ~ 63,571 + 7,467 = 71,038 times; the G142D toggling is not considered.

With thanks to Elodie Ghedin for leading the NIH working group discussion on Spike site 142, and to Jim Davis for sharing his data and a very helpful talk.

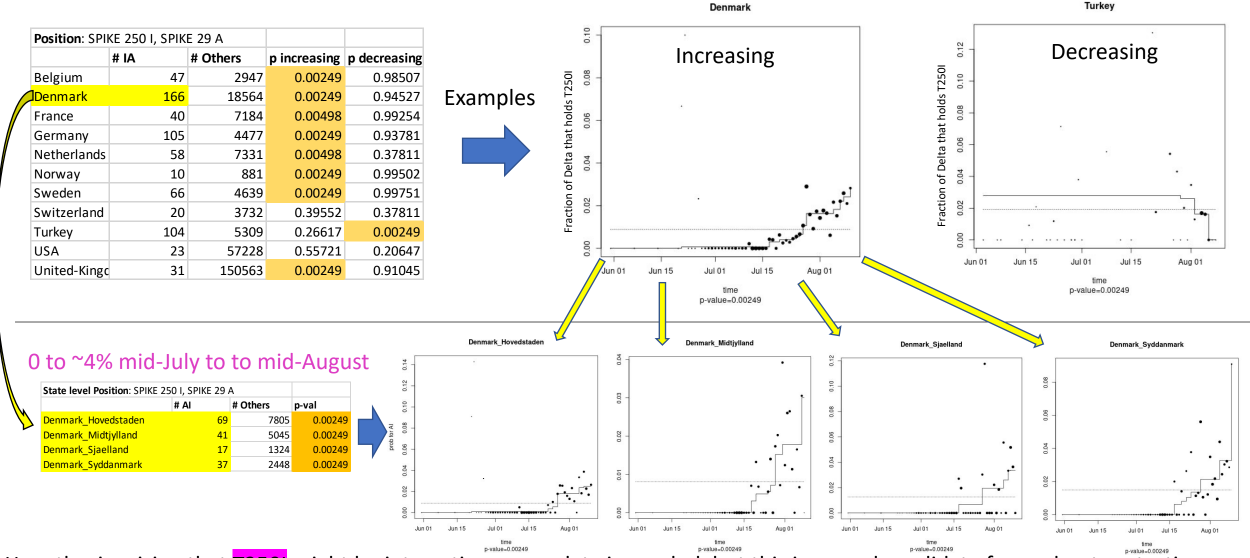
#### Example of a mutational pattern suggesting possible increase compared to other Deltas: T250I

counts

182 [T19R, T29A, E156-, F157-, R158G, T250I, L452R, T478K, D614G, P681R, D950N]

380 [T19R, T29A, G142D, E156-, F157-, R158G, T250I, L452R, T478K, D614G, P681R, D950N]

T250I had a significant presence in 11 countries, it was increasing in 8, decreasing in just 1, relative to other Delta's.



Hypothesis raising that T250I might be interesting, more data is needed, but this is a good candidate for exploratory testing

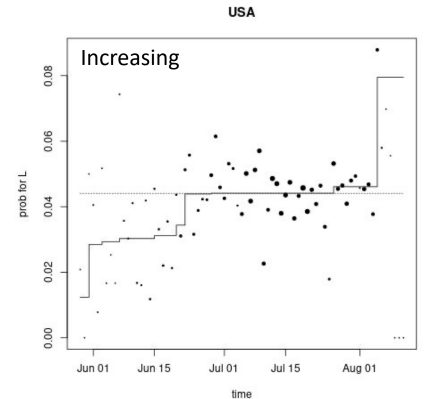
V1264 is a common mutation, but is likely founder/sampling effects, as it sometimes increases, sometimes decreases, and often is just maintained relative to other Deltas in the 19 countries it was found more than 10 times.

COUNTS	<mark>42D</mark> ,E156-,F157-,R158G,	L452R,T478K,D614G,P681R,D950N,	V1264L]
-			
213 [T19R,	E156-,F157-,R158G,	L452R,T478K,D614G,P681R,D950N,	V1264L]
744 [T19R, <mark>G1</mark>	<mark>42D</mark> ,E156-,F157-,R158G, <mark>A2</mark>	22V,L452R,T478K,D614G,P681R,D950N,	V1264L
346 [T19R,	The state of the s	22V,L452R,T478K,D614G,P681R,D950N,	V1264L
[,	, , , ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	, , , , , , , , , , , , , , , , , , , ,	,
		200V   450D T450V D4446 D404D D050V W4	0746 140641

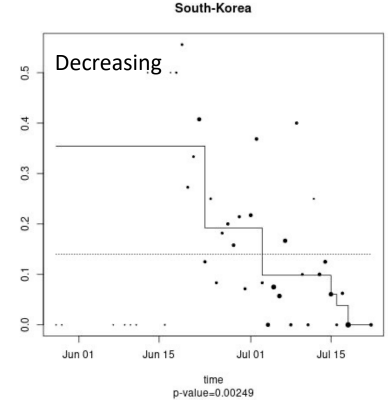
#### Countries

569 [T19R,<mark>G142D</mark>,E156-,F157-,R158G,A222V,L452R,T478K,D614G,P681R,D950N,<mark>N1074S,V1264L</mark>] 318 [T19R, E156-,F157-,R158G,A222V,L452R,T478K,D614G,P681R,D950N,<mark>N1074S,V1264L</mark>]

	p increasing	p-decreasing
Australia	0.00249	0.80348
Belgium	0.25622	0.92289
Denmark	0.05224	0.06219
France	0.84826	0.00249
Germany	0.00746	0.68657
Hong-Kong	0.93781	0.00249
India	0.57463	0.16169
Indonesia	0.0199	0.90796
Israel	0.16418	0.16667
Japan	0.00249	0.02488
Mexico	0.99751	0.00249
Netherlands	0.43781	0.6393
Singapore	0.00249	0.99502
South-Korea	0.6194	0.00249
Sweden	0.02239	0.53234
Switzerland	0.83085	0.00498
Turkey	0.00249	0.00498
USA	0.00249	0.92786
United-Kingdom	0.00249	0.88308



p-value=0.00249



This suggests that V1264 is a less interesting candidate for exploratory testing

Small alignment of the most variable positions in Spike, position numbers are written vertically.

#### To represent these variants, we would choose the version with G142D present

Few NTDss and RBD common changes

NTDss RBD 11111111 11111111222223444555666677788901112222

2225667799134455582455580145047178801957570261356 51791670757282567812801398762172370829109044429724 XSpike run on just Deltas: the most common forms with 195T The most common forms of Spike, those found > 300 time in the past 3 months. G142D toggles in association with almost all of them... only one exception, D138H.

Global <u>cseqs</u> . 246849 LRATTHAVKTKSDGYEFRGAYTPDVVKGLNTTQQSAETTIADNVGPGMSV	Global 246849	UK 98889		NAmer 53631	Asia Afr 9404		SAmer 4724					Pct [Context]
001IDGRN 002IGRN	83021 6973	69888 706	8841 4275	3093 1298	727 183	1 0	59 10	412 501				84% [T19R.T951.G142D.E156F157R158G.L452R.T478K.D614G.P681R.D950N] How to read this 82% [T19R,T951,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N]
003N	1102	1043	31	26	2	0	0	0	1102	825	74%	74% [T19R,T95I,D138H,G142D,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N] *
004I <mark>D</mark> GR	96 435	1 41	82 248	5 129	6 6	0 0	0 2	2 9	73 423			78% [T19R,T95I,G142D,E156-,F157-,R158G,L452R,T478K,D614G,P681R] 81% [T19R,T95I,E156-,F157-,R158G,L452R,T478K,D614G,P681R]
006I <mark>D</mark> GRN 007IGRIN.	1077 627	19 12	1034 610	15 5	1 0	0 0	8 0	0	1075 627			93% [T19R,T95I,G142D,E156-,F157-,R158G,L452R,T478K,D614G,P681R, <mark>T719I</mark> ,D950N] 94% [T19R,T95I,E156-,F157-,R158G,L452R,T478K,D614G,P681R, <mark>T719I</mark> ,D950N]
008IDGRNS 009IGRNS	771 14	758 7	9 6	1 0	3 1	0	0 0	9	771 14			93% [T19R,T95I,G142D,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N, <mark>P11625</mark> ] 00% [T19R,T95I,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N, <mark>P11625</mark> ]
010IDGRN 011IGRN	618 7	609 6	9 1	0 0	9 9	0	0 0	0	618 7			92% [T19R, <mark>T511</mark> ,T951,G142D,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N] 80% [T19R, <mark>T511</mark> ,T951,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N]
012	544 9	536 3	7 5	0	1	0	0 0	0	544 9			90% [T19R, <mark>V70F</mark> ,T95I,G142D,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N] 80% [T19R, <mark>V70F</mark> ,T95I,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N]
014IDGRN 015IGRRN	474 14	399 4	74 10	0 0	1 0	0	0	0	474 14			88% [T19R,T95I,G142D,E156-,F157-,R158G, <mark>Y248H</mark> ,L452R,T478K,D614G,P681R,D950N] 71% [T19R,T95I,E156-,F157-,R158G, <mark>Y248H</mark> ,L452R,T478K,D614G,P681R,D950N]
016IDGRN 017IGRN	458 27	397 1	38 16	20 8	3 2	0	0 0	0	458 27			86% [ <mark>L5F</mark> ,T19R,T95I,G142D,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N] 66% [ <mark>L5F</mark> ,T19R,T95I,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N]
018I <mark>D</mark> GRN 019IGRN	457 386	86 19	76 102	274 261	19 3	0	2 0	0 1	457 386			80% [T19R,T95I,G142D,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N, <mark>V1104L</mark> ] 68% [T19R,T95I,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N, <mark>V1104L</mark> ]
020IDGRN 021IGRN	342 68	7 11	323 57	1 0	0 0	11 0	0 0	0	342 68			88% [T19R, <mark>H66Y</mark> ,T95I,G142D,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N] 85% [T19R, <mark>H66Y</mark> ,T95I,E156-,F157-,R158G,L452R,T478K,D614G,P681R,D950N]
022I <mark>D</mark> GRN 023IGRN	339 188	0	318 180	16 6	5 2	0	0 0	0 0	339 188			81% [T19R,T95I,G142D,E156-,F157-,R158G,L452R,T478K,D614G,P681R, <mark>I850L</mark> ,D950N] 49% [T19R,T95I,E156-,F157-,R158G,L452R,T478K,D614G,P681R, <mark>I850L</mark> ,D950N]
024IDGRN 025IGRN	301 231	62 4	199 218	31 4	7 0	0	2 5	9	284 231			73% [T19R,T95I,G142D,E156-,F157-,R158G,L452R,T478K,D614G, <mark>Q677H</mark> ,P681R,D950N] 74% [T19R,T95I,E156-,F157-,R158G,L452R,T478K,D614G, <mark>Q677H</mark> ,P681R,D950N]
026 · · · · · · D · · · · · · · · · · · ·	63 365	0 1	20 196	6 32	37 136	0 0	0 0	9	61 296			31% [T19R,G142D,L452R,T478K,D614G,P681R,D950N] ** 73% [T19R,L452R,T478K,D614G,P681R,D950N]
Local Exact Pct [Cont	ext	]										
6064 5747 929 TT10	р т	SET	<b>E</b> 1	E 6	C10	-7	D	1 5 0	G 1	16	20	DD TATON D6146 D691D D050N]

6964 | T19R, T95I, E156-, F157-, R158G, L452R, T478K, D614G, P681R, D950N |

This mutation list refers to an exact from of Spike that carries this pattern of mutations. It was found exactly 5747 times, and accompanied by additional mutations 6964 times.

Range of dates: 2021-05-26 - 2021-08-24

Pango linage designation in GISAID (version: 2021-07-28): AY.1, AY.10, AY.11, AY.12, AY.2, AY.3, AY.3.1, AY.4, AY.5, AY.6, AY.7, AY.8, AY.9, B.1.617.2

## C.1.2 Which forms to use?

It is still quite rare and over half the Spike sequence have large gaps.

Bette Korber & James Theiler
Sept 1 GISAID data

Please see the new paper from the South African group regarding their discovery of this variant:

Scheepers et al. The continuous evolution of SARS-CoV-2 in South Africa: a new lineage with rapid accumulation of mutations of concern and global detection. https://www.medrxiv.org/content/10.1101/2021.08.20.21262342v2

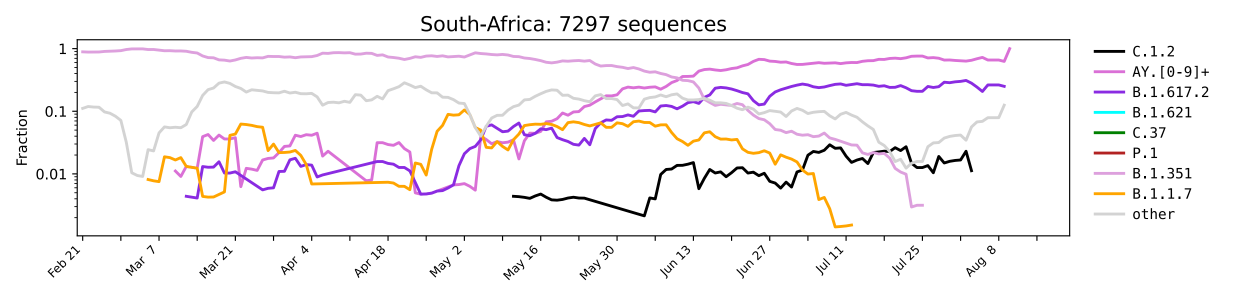
The mutations included in the listing to represent C.1.2 were based on very early sampling, as people were requesting them. It is very likely the Spike mutational listing will need updating as more data accrues. We were asked to select two for the NIH variant working group, so we picked the most common form in South Africa, and a still very rare variant found in the UK and Europe that had interesting amino acid substitutions.

#### This lineage is only sampled much in S. Africa, and even there still quite rare.

But it has also been detected in the UK, Portugal, Switzerland, Mauritius, Zimbabwe, Botswana, the DRC, Shenzhen.

In South Africa, C.1.2 sampling has come up very slightly. These plots of the dynamics are based on the Pango associated names in GISAID and inclusive of all data as of Sept. 1. 2021.

Beta (B.1.351) prevalence in S. Africa transitioned to Delta prevalent (B.1.617.2 and AY's) in early June. C.1.2 (black) has become more frequent over the summer, but is still quite rare (black). The plot is weekly average of sampling frequency, log scale, and highlights some forms we are tracking. We just split Delta on B.1.617.2 and AY's for convenience.



The Spike region between 150 and 410 is often missing data in C.1.2 lineage sequences sampled from GISAID 9/1/2021. The light blue X's are unresolved sequence in this amino acid alignment. We factor in only called bases to arrive at the consensus form, and then picked a variant that matched the consensus to try to be representative. --bk.

# Snapshot from C.1.2 Spike protein alignment 410 150

Only 7 of the 126 C.1.2 at the time sampling (9/1/2021) in GISAID passed LANL quality control in the complete genome, intact through the whole genome. Here are their Spikes. We choose the full genome from among these, picking the relevant Spike based on the complete alignment of all 126 C.1.2 available from GISAID on Sept 1.

#### The 7 QC passed C.1.2 have these 5 forms of Spike:

Form 3 is the backbone of C.1.2, both the Spike consensus and the most common form. GISAID example: EPI ISL 3132623

Form 1 has an additional mutation in the RBD, and is found internationally, UK, Portugal, Mauritius. GISAID example: EPI ISL 2989113

It also has N440K and has switched out N679K for the more common P681H, both of these mutations may augment Furin cleavage, and the N679K is the more common in C.1.2. All C.1.2 have one or the other, not both.

Notes on frequencies of particular Spike mutations in 126 C.1.2 that arise in addition to form 3 above:

P25LP L 49%

N440K K is rare 5%, but is international (UK, Portugal, as well as South Africa

T478K K is occasional, 20%

L585F is occasional, 19%

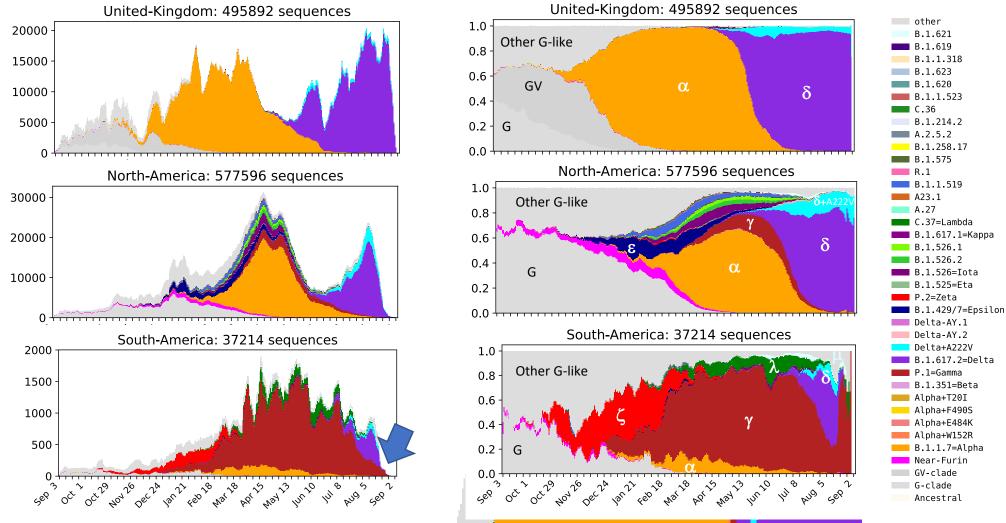
879T T is rare, 9%

D936H H is rare, 6%, only one N

P384L L is rare

P681H rare N679K is much more common, it is one or the other

#### Mu and Gamma seem to be sampled with increasing frequency recently in South America.

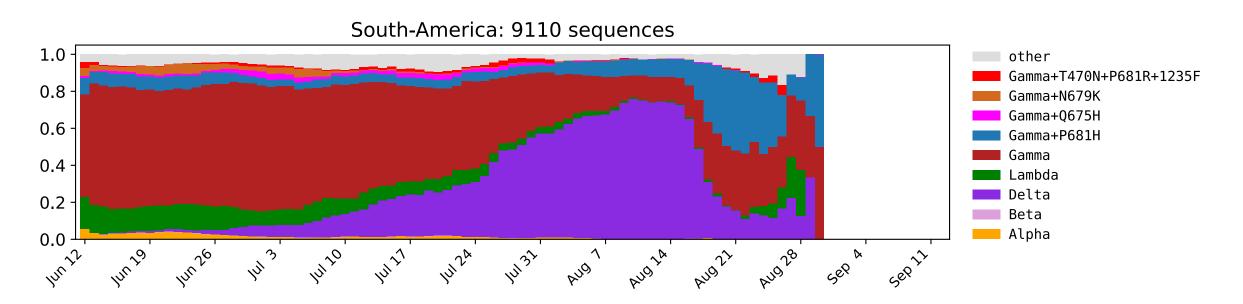


This may be an artifact due to very limited sampling the most recent weeks (see blue arrow), more time is needed to resolve this, but this situation merits continued monitoring to see if sampling resolves to increases in Delta.

The next few slides take a closer look at the Mu and Gamma variants that are persisting in Chile, Columbia, and Brazil. Interesting variant forms of Mu and Gamma are available in the 9/11/2021 listing.

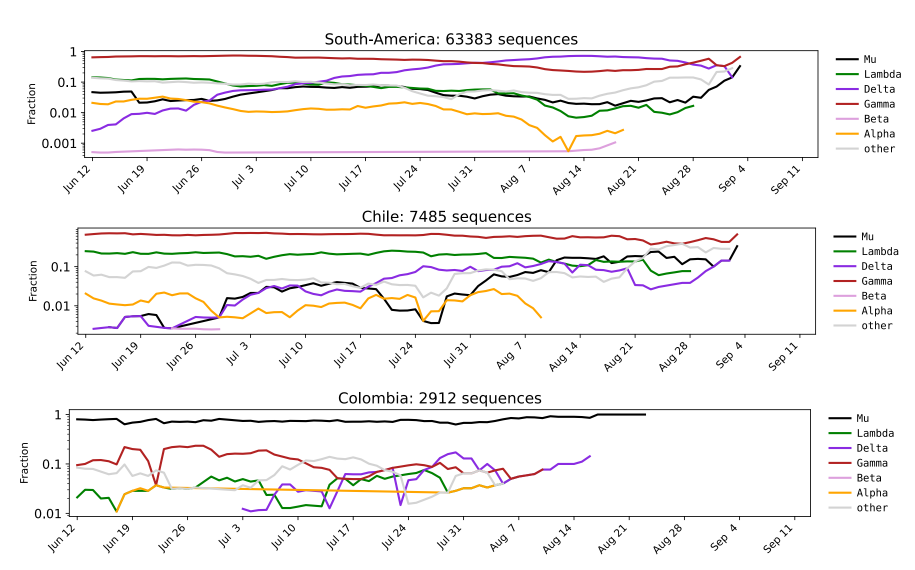
Variants of P.1 (Gamma) that carry increasing positive charge relative to base line P.1, particularly that have added P681H, but Q675H, N679K and 681R as well, are becoming increasing common among recent Brazilian samples, and persisting along with Delta. The numbers are still very small in recent sampling, and these Gamma mutations have some regional associations within Brazil, so the overall frequencies in each time window of each Gamma form are biased by local sampling issues.

This situation merits continued monitoring, to determine if the transition to Delta will continue in Brazil, or if novel P.1's may be competitive.



#### Mu: B.1.621

- -- Mu is most common in the America's.
- -- Mu was increasing in some parts of the United States through mid-July, but appears to be diminishing in sampling frequency as Delta has increased.
- -- Mu and variants are still common in S. America, particularly in Chile and Columbia.



#### Mu: B.1.621 + N417K and B.1.621.1

- 1) Mu Pango B.1.621 and B.1.621.1 are essentially the same in Spike in their most common form.
- [T95I,+143T,Y144S,Y145N,R346K,E484K,N501Y,D614G,P681H,D950N]
- 2) Within Mu, the mutation N417K is still rare increasing, and is becoming more commonly sampled than B.1.621 where the two are co-circulating (USA, Mexico, and a bit in the UK)

In countries where B.1.621 + N417K is found more than 10 times, the N417K form is increasing relative to other Mu variants.

Within the US: Texas, Washington, and California show an increased sampling of the N417K form

#### **Isotonic Regression Results**

**Isotonic Regression Results** 

Last data update: Sep 12, 2021

Position: SPIKE 417 N

Assumption: Test amino acid form is increasing over time Correlated variant: SPIKE 144 S incl, SPIKE 145 N incl

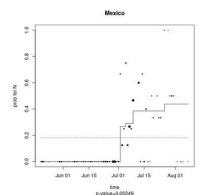
Range of dates: 2021-05-20 - 2021-09-18

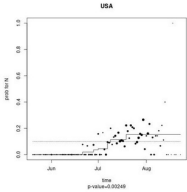
Hosts:

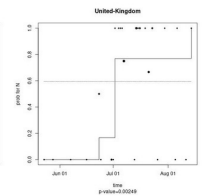
#### Download plots and data

#### Country level

	# N	# Others	total	# days	time window	p-val
Mexico	42	189	231	66	80	0.00249
USA	151	1373	1524	92	95	0.00249
United-Kingdom	22	15	37	26	83	0.00249







Position: SPIKE 417 N

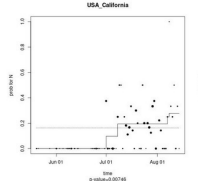
Assumption: Test amino acid form is increasing over time Correlated variant: SPIKE 144 S incl, SPIKE 145 N incl Range of dates: 2021-05-20 - 2021-09-18

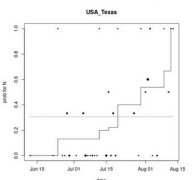
Hosts:

#### Download plots and data

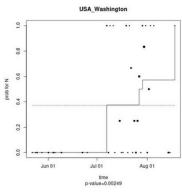
#### State level

	# N	# Others	total	# days	time window	p-val
USA_California	46	236	282	63	86	0.00746
USA_Texas	19	43	62	34	62	0.00746
USA_Washington	28	47	75	38	88	0.00249
United-Kingdom_England	20	12	32	23	80	0.00249





p-value=0.00746



Last data update: Sep 12, 2021