# Supplementary Text: Learning Mutational Patterns at Scale to Analyze Sequence Divergence in Novel **Pathogens**

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#### SUPPLEMENTARY METHODS: NOTES ON Q-DISTANCE & SUPPORTING RESULTS

The *q-distance* is a pseudo-metric since distinct sequences can induce the same distributions over each index, and thus evaluate to have a zero distance. This is actually desirable; we do not want our distance to be sensitive to changes that are not biologically relevant. The intuition is that not all sequence variations brought about by substitutions are equally important or likely. Even with no selection pressure, we might still see random variations at an index if such variations do not affect the replicative fitness. Under that scenario, the corresponding  $\Phi_i$  will predict a flat distribution no matter what the input sequence is, thus contributing nothing to the overall distance. And even if two strains x, y have the same entry at some index i, the remaining residues might induce different distributions  $\Phi_i$  based on the remote dependencies, i.e., the entries in  $x_{-i}, y_{-i}$ . Also, it matters if the sequences come from two different background populations P,Q, i.e., if the induced Qnets  $\Phi^P, \Phi^Q$  are different. Thus, if we construct Qnets for H1N1 Influenza A separately for the collection years 2008 and 2009, then the same exact sequence collected in the respective years might have a non-zero distance between them, reflecting the fact that the background population the sequences arose from are different, inducing possibly different expected mutational tendencies (See SI-Table 1).

Next, we induce q-distance between a sequence and a population and between two populations.

**Definition 1** (Pseudo-metric between populations). Using the notion of Hausdorff metric between sets:

$$\forall x \in P, y \in Q,$$

$$\theta(x,Q) = \min_{y \in Q} \theta(x,y)$$
(1)

$$\theta(x,Q) = \min_{y \in Q} \theta(x,y)$$

$$\theta(P,Q) = \max \left\{ \max_{x \in P} \theta(x,Q), \max_{y \in Q} \theta(y,P) \right\}$$
(2)

#### In-silico Corroboration of Qnet Constraints

We carry out in-silico experiments to corroborate that the constraints represented within an inferred Qnet are indeed reflective of the biology in play. We compare the results of simulated mutational perturbations to sequences from our databases (for which we have already constructed Qnets), and then use NCBI BLAST (https://blast.ncbi.nlm.nih.gov/Blast.cgi) to identify if our perturbed sequences match with existing sequences in the databases (See SI-Fig. 1). We find that in contrast to random variations, which rapidly diverge the trajectories, the Qnet constraints tend to produce smaller variance in the trajectories, maintain a high degree of match as we extend our trajectories, and produces matches closer in time to the collection time of the initial sequence suggesting that the Qnet does indeed capture realistic constraints.

#### Significance Test for Population Membership

For our modeling to be reliable, we need a quantitative test of how well the Qnet represents the data. Here, we formulate an explicit membership test to ascertain if individual samples may indeed be generated by the Qnet with sufficiently high probability.

**Definition 2** (Membership probability of a sequence). Given a population P inducing the Qnet  $\Phi^P$  and a sequence x, we can compute the membership probability of x:

$$\omega_x^P \triangleq Pr(x \in P) = \prod_{j=1}^N \left( \Phi_j^P(x_{-j})|_{x_j} \right) \tag{3}$$

 $x_j$  is the  $j^{th}$  entry in x, and is thus an element in the set  $\Sigma_j$ . Since we are mostly concerned with the case where  $\Sigma_j$  is a finite set,  $\Phi_j^P(x_{-j})|_{x_j}$  is the entry in the probability mass function corresponding to the element of  $\Sigma_j$  which appears at the  $j^{th}$  index in sequence x.

We can carry out this calculation for a sequence x known to be in the population P as well, which allows us to define the membership degree  $\omega_x^P$ .

**Definition 3** (Membership degree). Let X be a random field representing a population P, i.e., X = x is a randomly drawn sequence from P. Then the membership degree  $\omega^P$  is a function of the random variable X:

$$\omega^{P}(X) \triangleq \prod_{j=1}^{N} \left( \Phi_{j}^{P}(X_{-j})|_{X_{j}} \right) \tag{4}$$

Note that  $\omega^P$  takes values in the unit interval [0,1], and the probability x is a member of the population P is  $\omega^P(X=x)$ , denoted briefly as  $\omega^P_x$  or  $\omega_x$  if P is clear from context.

Since  $\omega^P(X)$  is a random variable, we can now compute sets of sequences that better represent the population P, and ones that are on the fringe. We can also evaluate using a pre-specified significance-level if a particular sequence is not from the population P, thus identifying if we need to recompute the predictors  $\Phi$ , or split the base population. We can set up a hypothesis testing scenario to determine if sequences are indeed from a test population, as follows:

Given a population P, inducing a Qnet  $\Phi^P$ , and a sequence x, we assume the null hypothesis is  $x \notin P$ . We reject the null hypothesis at a pre-specified significance  $\alpha$ , if

$$Pr(\omega^P(X) \ge \omega^P(X = x)) \le \alpha$$
 (5)

The fraction of newly observed sequences that do not reject the null hypothesis can then be used as an estimate of the species-specific divergence in population characteristics.

### **Proof of Probability Bounds**

**Theorem 1** (Probability bound). Given a sequence x of length N that transitions to a strain  $y \in Q$ , we have the following bounds at significance level  $\alpha$ .

$$\omega_y^Q e^{\frac{\sqrt{8}N^2}{1-\alpha}\theta(x,y)} \ge Pr(x \to y) \ge \omega_y^Q e^{-\frac{\sqrt{8}N^2}{1-\alpha}\theta(x,y)}$$
(6)

where  $\omega_y^Q$  is the membership probability of strain y in the target population Q (See Def. 2), and  $\theta(x,y)$  is the q-distance between x,y (See Def. 2 in Qnet Framework).

*Proof.* Using Sanov's theorem<sup>1</sup> on large deviations, we conclude that the probability of spontaneous jump from strain  $x \in P$  to strain  $y \in Q$ , with the possibility  $P \neq Q$ , is given by:

$$Pr(x \to y) = \prod_{i=1}^{N} \left( \Phi_i^P(x_{-i})|_{y_i} \right) \tag{7}$$

Writing the factors on the right hand side as:

$$\Phi_i^P(x_{-i})|_{y_i} = \Phi_i^Q(y_{-i})|_{y_i} \left( \frac{\Phi_i^P(x_{-i})|_{y_i}}{\Phi_i^Q(y_{-i})|_{y_i}} \right)$$
(8)

we note that  $\Phi_i^P(x_{-i})$ ,  $\Phi_i^Q(y_{-i})$  are distributions on the same index i, and hence:

$$|\Phi_i^P(x_{-i})_{y_i} - \Phi_i^Q(y_{-i})_{y_i}| \le \sum_{y_i \in \Sigma_i} |\Phi_i^P(x_{-i})_{y_i} - \Phi_i^Q(y_{-i})_{y_i}|$$
(9)

Using a standard refinement of Pinsker's inequality<sup>2</sup>, and the relationship of Jensen-Shannon divergence with

total variation, we get:

$$\theta_{i} \ge \frac{1}{8} |\Phi_{i}^{P}(x_{-i})_{y_{i}} - \Phi_{i}^{Q}(y_{-i})_{y_{i}}|^{2} \Rightarrow \left| 1 - \frac{\Phi_{i}^{Q}(y_{-i})_{y_{i}}}{\Phi_{i}^{P}(x_{-i})_{y_{i}}} \right| \le \frac{1}{a_{0}} \sqrt{8\theta_{i}}$$

$$\tag{10}$$

where  $a_0$  is the smallest non-zero probability value of generating the entry at any index. We will see that this parameter is related to statistical significance of our bounds. First, we can formulate a lower bound as follows:

$$\log \left( \prod_{i=1}^{N} \frac{\Phi_{i}^{P}(x_{-i})|_{y_{i}}}{\Phi_{i}^{Q}(y_{-i})|_{y_{i}}} \right) = \sum_{i} \log \left( \frac{\Phi_{i}^{P}(x_{-i})|_{y_{i}}}{\Phi_{i}^{Q}(y_{-i})|_{y_{i}}} \right) \ge \sum_{i} \left( 1 - \frac{\Phi_{i}^{Q}(y_{-i})_{y_{i}}}{\Phi_{i}^{P}(x_{-i})_{y_{i}}} \right) \ge \frac{\sqrt{8}}{a_{0}} \sum_{i} \theta_{i}^{1/2} = -\frac{\sqrt{8}N}{a_{0}} \theta$$
 (11)

Similarly, the upper bound may be derived as:

$$\log \left( \prod_{i=1}^{N} \frac{\Phi_{i}^{P}(x_{-i})|_{y_{i}}}{\Phi_{i}^{Q}(y_{-i})|_{y_{i}}} \right) = \sum_{i} \log \left( \frac{\Phi_{i}^{P}(x_{-i})|_{y_{i}}}{\Phi_{i}^{Q}(y_{-i})|_{y_{i}}} \right) \leq \sum_{i} \left( \frac{\Phi_{i}^{Q}(y_{-i})_{y_{i}}}{\Phi_{i}^{P}(x_{-i})_{y_{i}}} - 1 \right) \leq \frac{\sqrt{8}N}{a_{0}} \theta$$
(12)

Combining Eqs. 11 and 12, we conclude:

$$\omega_y^Q e^{\frac{\sqrt{8}N}{a_0}\theta} \ge Pr(x \to y) \ge \omega_y^Q e^{-\frac{\sqrt{8}N}{a_0}\theta} \tag{13}$$

Now, interpreting  $a_0$  as the probability of generating an unlikely event below our desired threshold (*i.e.* a "failure"), we note that the probability of generating at least one such event is given by  $1 - (1 - a_0)^N$ . Hence if  $\alpha$  is the pre-specified significance level, we have for N >> 1:

$$a_0 \approx (1 - \alpha)/N \tag{14}$$

Hence, we conclude, that at significance level  $\geq \alpha$ , we have the bounds:

$$\omega_y^Q e^{\frac{\sqrt{8}N^2}{1-\alpha}\theta} \ge Pr(x \to y) \ge \omega_y^Q e^{-\frac{\sqrt{8}N^2}{1-\alpha}\theta}$$
(15)

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**Remark 1.** This bound can be rewritten in terms of the log-likelihood of the spontaneous jump and constants independent of the initial sequence x as:

$$|\log Pr(x \to y) - C_0| \le C_1 \theta \tag{16}$$

where the constants are given by:

$$C_0 = \log \omega_y^Q \tag{17}$$

$$C_1 = \frac{\sqrt{8}N^2}{1 - \alpha} \tag{18}$$

#### Multivariate Regression to Identify Factors in Strain Prediction

We investigate the key factors that contribute to our successful prediction of the dominant strain in the next season. We carry out a multivariate regression with data diversity, the complexity of inferred Qnet and the edit distance of the WHO recommendation from the dominant strain as independent variables. Here we define data diversity as the number of clusters we have in the input set of sequences, such that any two sequences five or less mutations apart are in the same cluster. Qnet complexity is measured by the number of decision nodes in the component decision trees of the recursive forest.

We select several plausible structures of the regression equation, and in each case conclude that data diversity has the most important and statistically significant contribution (See SI-Tab. 17).

#### REFERENCES

- [1] Cover TM, Thomas JA. Elements of Information Theory (Wiley Series in Telecommunications and Signal Processing). New York, NY, USA: Wiley-Interscience; 2006.
- [2] Fedotov AA, Harremoës P, Topsoe F. Refinements of Pinsker's inequality. IEEE Transactions on Information Theory. 2003;49(6):1491–1498.

## SUPPLEMENTARY FIGURES & TABLES

SI Tab. 1

EXAMPLES: QNET INDUCED DISTANCE VARYING FOR FIXED SEQUENCE PAIR WHEN BACKGROUND POPULATION CHANGES (ROWS 1 -5), SEQUENCES WITH SMALL EDIT DISTANCE AND LARGE Q-DISTANCE, AND THE CONVERSE (ROWS 6-9)

	Edit dist.	Sequence A	Sequence B	Q-dist.	Year A*	Year B*
1	18	A/Singapore/23J/2007	A/Tennessee/UR06-0294/2007	0.0111	2007	2007
2	18	A/Singapore/23J/2007	A/Tennessee/UR06-0294/2007	0.0094	2008	2008
3	18	A/Singapore/23J/2007	A/Tennessee/UR06-0294/2007	0.0027	2009	2009
4	18	A/Singapore/23J/2007	A/Tennessee/UR06-0294/2007	0.0025	2010	2010
5	18	A/Singapore/23J/2007	A/Tennessee/UR06-0294/2007	0.6163	2007	2010
6	11	A/Naypyitaw/M783/2008	A/Singapore/201/2008	0.8852	2008	2008
7	15	A/Cambodia/W0908339/2012	A/Singapore/DMS1233/2012	0.2737	2012	2012
8	126	A/South Dakota/03/2008	A/Singapore/10/2008	0.3034	2008	2008
9	141	A/Jodhpur/3248/2012	A/Cambodia/W0908339/2012	0.2405	2012	2012

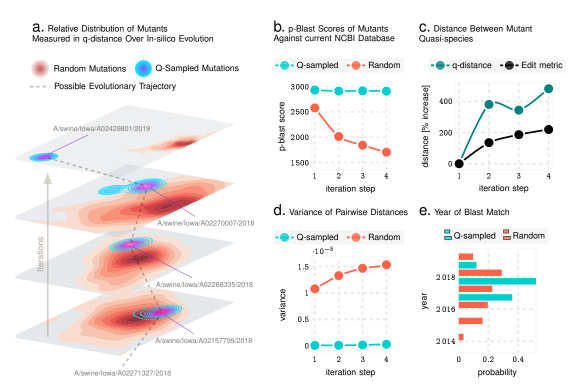
<sup>\*</sup>Year A and year B correspond to the assumed collection years for sequences A and B respectively for the purpose of this example. Sequence A in row 1 is collected in 2007, but is assumed to be from different years in rows 2-4 to demonstrate the change in q-distance from sequence B, arising only from a change in the background population.

SI Tab. 2 CORRELATION BETWEEN Q-DISTANCE AND EDIT DISTANCE BETWEEN SEQUENCE PAIRS

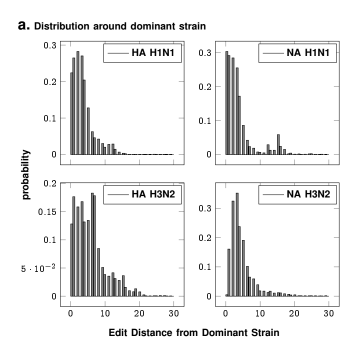
Phenotypes	Correlation
Influenza H1N1 HA	0.76
Influenza H1N1 NA	0.74
Influenza H3N2 HA	0.85
Influenza H3N2 NA	0.79

SI Tab. 3
NUMBER OF SEQUENCES COLLECTED FROM PUBLIC DATABASES

Database	Strain	No. of Sequences
NCBI	Influenza H1N1 HA	17,894
NCBI	Influenza H1N1 NA	16,637
NCBI	Influenza H3N2 HA	18,265
NCBI	Influenza H3N2 NA	14,699
GISAID	Influenza H1N1 HA	1,528
GISAID	Influenza H1N1 NA	1,490
GISAID	Influenza H3N2 HA	13,975
GISAID	Influenza H3N2 NA	13,811
Total		98,299



SI Fig. 1. Q-distance validation in-silico using Influenza A sequences from NCBI database. Panel a illustrates that the Qnet induced modeling of evolutionary trajectories initiated from known haemagglutinnin (HA) sequences are distinct from random paths in the strain space. In particular, random trajectories have more variance, and more importantly, diverge to different regions of the landscape compared to Qnet predictions. Panels b-e show that unconstrained Q-sampling produces sequences maintain a higher degree of similarity to known sequences, as verified by blasting against known HA sequences, have a smaller rate of growth of variance, and produce matches in closer time frames to the initial sequence. Panel c shows that this is not due to simply restricting the mutational variations, which increases rapidly in both the Qnet and the classical metric.



SI Fig. 2. **No. of mutations from the seasonal dominant strain over the years** The quasispecies that circulates each season for each sub-type is tightly distributed around the dominant strain on average.

SI Tab. 4 H1N1 HA NORTHERN HEMISPHERE

Year	WHO Recommendation	Dominant Strain	Qnet Recommendation	WHO Error	Qnet Error
2001-02	A/New Caledonia/20/99	A/Canterbury/41/2001	A/Dunedin/2/2000	4	6
2002-03	A/New Caledonia/20/99	A/Taiwan/567/2002	A/New York/241/2001	3	1
2003-04	A/New Caledonia/20/99	A/Memphis/5/2003	A/New York/291/2002	5	2
2004-05	A/New Caledonia/20/99	A/Thailand/Siriraj-Rama-TT/2004	A/New York/222/2003	7	4
2005-06	A/New Caledonia/20/99	A/Niedersachsen/217/2005	A/Canterbury/106/2004	8	10
2006-07	A/New Caledonia/20/99	A/India/34980/2006	A/Auckland/619/2005	6	1
2007-08	A/Solomon Islands/3/2006	A/Norway/1701/2007	A/New York/8/2006	8	11
2008-09	A/Brisbane/59/2007	A/Pennsylvania/02/2008	A/Kentucky/UR06-0476/2007	2	2
2009-10	A/Brisbane/59/2007	A/Singapore/ON1060/2009	A/Hong Kong/549/2008	119	119
2010-11	A/California/7/2009	A/England/01220740/2010	A/New York/14/2009	5	1
2011-12	A/California/7/2009	A/Punjab/041/2011	A/Kansas/01/2010	7	2
2012-13	A/California/7/2009	A/British Columbia/001/2012	A/Moscow/WRAIR4308T/2011	11	4
2013-14	A/California/7/2009	A/Moscow/CRIE-32/2013	A/Helsinki/1199/2012	10	2
2014-15	A/California/7/2009	A/Thailand/CU-C5169/2014	A/Maryland/02/2013	12	0
2015-16	A/California/7/2009	A/Georgia/15/2015	A/Utah/3691/2014	14	2
2016-17	A/California/7/2009	A/Hawaii/21/2016	A/Adana/08/2015	16	0
2017-18	A/Michigan/45/2015	A/Michigan/291/2017	A/Beijing-Huairou/SWL1335/2016	5	4
2018-19	A/Michigan/45/2015	A/Washington/55/2018	A/India/C1721549/2017	6	1
2019-20	A/Brisbane/02/2018	A/Kentucky/06/2019	A/New Jersey/01/2018	5	1
2020-21	A/Hawaii/70/2019	A/Togo/905/2020	A/ltaly/8949/2019	4	8
2021-22	A/Victoria/2570/2019	A/Ireland/20935/2022	A/Togo/45/2021	9	3
2022-23	-1	-1	A/Netherlands/00068/2022	-1	-1

<sup>\*</sup> Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric

SI Tab. 5 H1N1 HA SOUTHERN HEMISPHERE

Year	WHO Recommendation	Dominant Strain	Qnet Recommendation		Qnet Error
2001-02	A/New Caledonia/20/99	A/Canterbury/41/2001	A/South Canterbury/50/2000	4	6
2002-03	A/New Caledonia/20/99	A/Taiwan/567/2002	A/Canterbury/41/2001	3	1
2003-04	A/New Caledonia/20/99	A/Memphis/5/2003	A/New York/291/2002	5	2
2004-05	A/New Caledonia/20/99	A/Thailand/Siriraj-Rama-TT/2004	A/Memphis/5/2003	7	4
2005-06	A/New Caledonia/20/99	A/Niedersachsen/217/2005	A/Canterbury/106/2004	8	10
2006-07	A/New Caledonia/20/99	A/India/34980/2006	A/Niedersachsen/217/2005	6	2
2007-08	A/New Caledonia/20/99	A/Norway/1701/2007	A/Thailand/CU68/2006	14	6
2008-09	A/Solomon Islands/3/2006	A/Pennsylvania/02/2008	A/Kentucky/UR06-0476/2007	9	2
2009-10	A/Brisbane/59/2007	A/Singapore/ON1060/2009	A/Belem/241/2008	119	119
2010-11	A/California/7/2009	A/England/01220740/2010	A/Singapore/ON1060/2009	5	1
2011-12	A/California/7/2009	A/Punjab/041/2011	A/England/01220740/2010	7	2
2012-13	A/California/7/2009	A/British Columbia/001/2012	A/Punjab/041/2011	11	4
2013-14	A/California/7/2009	A/Moscow/CRIE-32/2013	A/India/P122045/2012	10	5
2014-15	A/California/7/2009	A/Thailand/CU-C5169/2014	A/Jiangsuhailing/SWL1382/2013	12	4
2015-16	A/California/7/2009	A/Georgia/15/2015	A/Thailand/CU-C5169/2014	14	2
2016-17	A/California/7/2009	A/Hawaii/21/2016	A/Georgia/15/2015	16	2
2017-18	A/Michigan/45/2015	A/Michigan/291/2017	A/Beijing-Huairou/SWL1335/2016	5	4
2018-19	A/Michigan/45/2015	A/Washington/55/2018	A/Michigan/291/2017	6	1
2019-20	A/Michigan/45/2015	A/Kentucky/06/2019	A/Washington/55/2018	7	1
2020-21	A/Brisbane/02/2018	A/Togo/905/2020	A/Italy/8451/2019	10	8
2021-22	A/Victoria/2570/2019	A/Abidjan/457/2021	A/Togo/35/2021	9	4
2022-23	-1	-1	A/Cote_D'Ivoire/1270/2021	-1	-1

<sup>\*</sup> Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric

SI Tab. 6 H1N1 NA NORTHERN HEMISPHERE

Year	WHO Recommendation	Dominant Strain	Qnet Recommendation	WHO Error	Qnet Error
2001-02	A/New Caledonia/20/99	A/New York/447/2001	A/Memphis/15/2000	4	4
2002-03	A/New Caledonia/20/99	A/Paris/0833/2002	A/New York/341/2001	1	5
2003-04	A/New Caledonia/20/99	A/Memphis/5/2003	A/New York/291/2002	3	5
2004-05	A/New Caledonia/20/99	A/Singapore/14/2004	A/New York/223/2003	2	3
2005-06	A/New Caledonia/20/99	A/Taiwan/5524/2005	A/Florida/3e/2004	3	0
2006-07	A/New Caledonia/20/99	A/Massachusetts/08/2006	A/Sofia/361/2005	4	2
2007-08	A/Solomon Islands/3/2006	A/Tennessee/UR06-0106/2007	A/Sofia/490/2006	9	2
2008-09	A/Brisbane/59/2007	A/Sendai/TU66/2008	A/Maryland/04/2007	0	3
2009-10	A/Brisbane/59/2007	A/Thailand/SR08021/2009	A/Paris/910/2008		87
2010-11	A/California/7/2009	A/Finland/2460N/2010	A/Rome/709/2009	2	9
2011-12	A/California/7/2009	A/Tula/CRIE-GSYu/2011	A/Oman/SQUH-40/2010	4	2
2012-13	A/California/7/2009	A/Bangalore/697-32/2012	A/Nizhnii Novgorod/CRIE-ZCA/2011	4	0
2013-14	A/California/7/2009	A/Jiangsugusu/SWL1824/2013	A/LongYan/SWL33/2013	5	3
2014-15	A/California/7/2009	A/LongYan/SWL2457/2014	A/Utah/06/2013	9	3
2015-16	A/California/7/2009	A/Michigan/45/2015	A/Maryland/02/2014	14	4
2016-17	A/California/7/2009	A/Mexico/4436/2016	A/India/Pun151245/2015	14	0
2017-18	A/Michigan/45/2015	A/Illinois/37/2017	A/Utah/02/2016	3	3
2018-19	A/Michigan/45/2015	A/Kenya/47/2018	A/Maine/24/2017	4	0
2019-20	A/Brisbane/02/2018	A/Texas/7939/2019	A/Missouri/03/2018	1	0
2020-21	A/Hawaii/70/2019	A/Togo/897/2020	A/Texas/112/2019	0	5
2021-22	A/Victoria/2570/2019	A/Cote_d'Ivoire/3729/2021	A/Togo/0071/2021	1	5
2022-23	-1	-1	A/Lyon/820/2021	-1	-1

<sup>\*</sup> Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric

SI Tab. 7 H1N1 NA SOUTHERN HEMISPHERE

Year	WHO Recommendation	Dominant Strain	Qnet Recommendation		Qnet Error
2001-02	A/New Caledonia/20/99	A/New York/447/2001	A/Canterbury/37/2000	4	6
2002-03	A/New Caledonia/20/99	A/Paris/0833/2002	A/New York/447/2001	1	5
2003-04	A/New Caledonia/20/99	A/Memphis/5/2003	A/New York/291/2002	3	5
2004-05	A/New Caledonia/20/99	A/Singapore/14/2004	A/Memphis/5/2003	2	3
2005-06	A/New Caledonia/20/99	A/Taiwan/5524/2005	A/Canterbury/106/2004	3	6
2006-07	A/New Caledonia/20/99	A/Massachusetts/08/2006	A/Sofia/361/2005	4	2
2007-08	A/New Caledonia/20/99	A/Tennessee/UR06-0106/2007	A/Thailand/RMSC-UDN-20/2006	4	8
2008-09	A/Solomon Islands/3/2006	A/Sendai/TU66/2008	A/Tennessee/UR06-0151/2007	15	13
2009-10	A/Brisbane/59/2007	A/Thailand/SR08021/2009	A/Nebraska/07/2008	87	87
2010-11	A/California/7/2009	A/Finland/2460N/2010	A/Rome/709/2009	2	9
2011-12	A/California/7/2009	A/Tula/CRIE-GSYu/2011	A/Finland/2460N/2010	4	2
2012-13	A/California/7/2009	A/Bangalore/697-32/2012	A/Tula/CRIE-GSYu/2011	4	0
2013-14	A/California/7/2009	A/Jiangsugusu/SWL1824/2013	A/Oman/SQUH-63/2012	5	4
2014-15	A/California/7/2009	A/LongYan/SWL2457/2014	A/NanPing/SWL1640/2013	9	6
2015-16	A/California/7/2009	A/Michigan/45/2015	A/LongYan/SWL2457/2014	14	5
2016-17	A/California/7/2009	A/Mexico/4436/2016	A/Michigan/45/2015	14	0
2017-18	A/Michigan/45/2015	A/Illinois/37/2017	A/Mexico/4436/2016	3	3
2018-19	A/Michigan/45/2015	A/Kenya/47/2018	A/Kentucky/26/2017	4	2
2019-20	A/Michigan/45/2015	A/Texas/7939/2019	A/Kenya/47/2018	4	0
2020-21	A/Brisbane/02/2018	A/Togo/897/2020	A/Texas/7939/2019	6	5
2021-22	A/Victoria/2570/2019	A/Cote_D'Ivoire/1496/2021	A/Togo/0155/2021	1	7
2022-23	-1	-1	A/Dakar/35/2021	-1	-1

<sup>\*</sup> Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric

SI Tab. 8 H3N2 HA NORTHERN HEMISPHERE

Year	WHO Recommendation	Dominant Strain	Dominant Strain Qnet Recommendation		Qnet Error
2005-06	A/California/7/2004	A/Denmark/195/2005	A/Tairawhiti/369/2004		2
2006-07	A/Wisconsin/67/2005	A/New York/5/2006	A/South Australia/22/2005	5	4
2007-08	A/Wisconsin/67/2005	A/Tennessee/11/2007	A/Colorado/05/2006	8	5
2008-09	A/Brisbane/10/2007	A/Massachusetts/13/2008	A/Virginia/UR06-0021/2007	3	2
2009-10	A/Brisbane/10/2007	A/Hawaii/14/2009	A/Manhean/03/2008	7	6
2010-11	A/Perth/16/2009	A/Utah/12/2010	A/Philippines/5/2009	8	7
2011-12	A/Perth/16/2009	A/Piaui/14202/2011	A/Singapore/C2010.310/2010	4	4
2012-13	A/Victoria/361/2011	A/Alborz/927/2012 A/Tehran/895/2012		4	3
2013-14	A/Victoria/361/2011	A/Delaware/01/2013 A/Singapore/H2012.934/2012		4	1
2014-15	A/Texas/50/2012	A/Alborz/72205/2014 A/Nebraska/03/2013		10	9
2015-16	A/Switzerland/9715293/2013	A/Parma/471/2015	A/Ontario/01/2014	10	0
2016-17	A/Hong Kong/4801/2014	A/Guangdong/12/2016	A/Oregon/02/2015	0	0
2017-18	A/Hong Kong/4801/2014	A/Maryland/25/2017	A/New York/03/2016	3	1
2018-19	A/Singapore/INFIMH-16-0019/2016	A/Vermont/04/2018	A/Ontario/038/2017	8	5
2019-20	A/Kansas/14/2017	A/Kentucky/27/2019	A/California/7330/2018	16	12
2020-21	A/Hong Kong/2671/2019	A/India/Pun-NIV289524/2021_Jan	A/California/NHRC- OID_FDX100215/2019		14
2021-22	A/Cambodia/e0826360/2020	A/Human/New_York/PV60641/2022	A/India/Pun-NIV291000/2021_Jan	14	5
2022-23	-1	-1	A/Denmark/370/2022		-1

<sup>\*</sup> Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric

SI Tab. 9 H3N2 HA SOUTHERN HEMISPHERE

Year	WHO Recommendation	Dominant Strain Qnet Recommendation		WHO Error	Qnet Error
2005-06	A/Wellington/1/2004	A/Denmark/195/2005	A/Waikato/21/2004	3	3
2006-07	A/California/7/2004	A/New York/5/2006	A/South Australia/22/2005	12	4
2007-08	A/Wisconsin/67/2005	A/Tennessee/11/2007	A/New York/923/2006	8	5
2008-09	A/Brisbane/10/2007	A/Massachusetts/13/2008	A/Tennessee/11/2007	3	2
2009-10	A/Brisbane/10/2007	A/Hawaii/14/2009	A/Manhean/03/2008	7	6
2010-11	A/Perth/16/2009	A/Utah/12/2010	A/Hawaii/14/2009	8	7
2011-12	A/Perth/16/2009	A/Piaui/14202/2011	A/Utah/12/2010	4	4
2012-13	A/Perth/16/2009	A/Alborz/927/2012	A/Alborz/927/2012 A/Piaui/14202/2011		4
2013-14	A/Victoria/361/2011	A/Delaware/01/2013	A/Callao/IPE00830/2012	4	7
2014-15	A/Texas/50/2012	A/Alborz/72205/2014	A/Delaware/01/2013	10	7
2015-16	A/Switzerland/9715293/2013	A/Parma/471/2015	A/Alborz/72205/2014	10	0
2016-17	A/Hong Kong/4801/2014	A/Guangdong/12/2016	A/Parma/471/2015	0	0
2017-18	A/Hong Kong/4801/2014	A/Maryland/25/2017	A/Ontario/196/2016	3	4
2018-19	A/Singapore/INFIMH-16-0019/2016	A/Vermont/04/2018	A/Texas/279/2017	8	5
2019-20	A/Switzerland/8060/2017	A/Kentucky/27/2019 A/Santa Catarina/1200/2018		13	12
2020-21	A/South Australia/34/2019	A/India/Pun-NIV289524/2021_Jan	A/Kentucky/27/2019	12	14
2021-22	A/Hong Kong/2671/2019	A/Darwin/9a/2021	A/India/PUN-NIV301718/2021	19	1
2022-23	022-23 -1 -1 A/Saint-Martin/00754/2022		A/Saint-Martin/00754/2022	-1	-1

 $<sup>^{\</sup>star}$  Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric

SI Tab. 10 H3N2 NA NORTHERN HEMISPHERE

Year	WHO Recommendation	Dominant Strain	Dominant Strain Qnet Recommendation		Qnet Error
2003-04	A/Moscow/10/99	A/Denmark/107/2003	A/New York/100/2002	13	3
2004-05	A/Fujian/411/2002	A/Hyogo/36/2004	A/New York/20/2003	3	16
2005-06	A/California/7/2004	A/Denmark/203/2005	A/Hong Kong/HKU20/2004	4	0
2006-07	A/Wisconsin/67/2005	A/Berlin/32/2006	A/Mexico/InDRE2227/2005	1	1
2007-08	A/Wisconsin/67/2005	A/Brazil/80/2007	A/Baden-Wuerttemberg/17/2006	8	7
2008-09	A/Brisbane/10/2007	A/Missouri/05/2008	A/Washington/01/2007	3	2
2009-10	A/Brisbane/10/2007	A/Oklahoma/09/2009	A/Wisconsin/24/2008	3	1
2010-11	A/Perth/16/2009	A/California/17/2010	A/California/17/2010 A/New York/70/2009		3
2011-12	A/Perth/16/2009	A/Texas/14/2011 A/California/14/2010		3	2
2012-13	A/Victoria/361/2011	A/New York/02/2012 A/Singapore/C2011.493/2011		4	1
2013-14	A/Victoria/361/2011	A/Michigan/02/2013	A/New York/01/2012	3	1
2014-15	A/Texas/50/2012	A/Tehran/69634/2014	A/Boston/DOA2-176/2013	3	1
2015-16	A/Switzerland/9715293/2013	A/Parma/471/2015	A/Thailand/CU-B10520/2014	3	0
2016-17	A/Hong Kong/4801/2014	A/North Carolina/62/2016	A/Delaware/02/2015	7	2
2017-18	A/Hong Kong/4801/2014	A/Texas/277/2017	A/New York/03/2016	8	0
2018-19	A/Singapore/INFIMH-16-0019/2016	A/Japan/NHRC_FDX70352/2018	A/Colorado/11/2017	4	3
2019-20	A/Kansas/14/2017	A/Washington/9757/2019 A/Guangxi-Fangcheng/54/2019		3	11
2020-21	A/Hong Kong/2671/2019	A/Bangladesh/1004005/2020	A/Maryland/02/2019	3	13
2021-22	A/Cambodia/e0826360/2020	A/Stockholm/10/2022	A/Darwin/9/2021	2	2
2022-23	-1	-1	A/Michigan/UOM10042819294/2021	-1	-1

<sup>\*</sup> Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric

SI Tab. 11 H3N2 NA SOUTHERN HEMISPHERE

Year	WHO Recommendation	Dominant Strain	Qnet Recommendation	WHO Error	Qnet Error
2003-04	A/Moscow/10/99	A/Denmark/107/2003	A/New York/101/2002		3
2004-05	A/Fujian/411/2002	A/Hyogo/36/2004	A/New York/20/2003	3	16
2005-06	A/Wellington/1/2004	A/Denmark/203/2005	A/Wellington/1/2004	2	2
2006-07	A/California/7/2004	A/Berlin/32/2006	A/Mexico/InDRE2227/2005	3	1
2007-08	A/Wisconsin/67/2005	A/Brazil/80/2007	A/Ohio/06/2006	8	10
2008-09	A/Brisbane/10/2007	A/Missouri/05/2008	A/Brazil/80/2007	3	2
2009-10	A/Brisbane/10/2007	A/Oklahoma/09/2009	A/Wisconsin/24/2008	3	1
2010-11	A/Perth/16/2009	A/California/17/2010	A/California/17/2010 A/New York/70/2009		3
2011-12	A/Perth/16/2009	A/Texas/14/2011	A/Texas/14/2011 A/Virginia/05/2010		2
2012-13	A/Perth/16/2009	A/New York/02/2012	A/Texas/14/2011	4	1
2013-14	A/Victoria/361/2011	A/Michigan/02/2013	A/New York/02/2012	3	3
2014-15	A/Texas/50/2012	A/Tehran/69634/2014	A/Michigan/02/2013	3	1
2015-16	A/Switzerland/9715293/2013	A/Parma/471/2015	A/Tehran/69634/2014	3	2
2016-17	A/Hong Kong/4801/2014	A/North Carolina/62/2016	A/Parma/471/2015	7	2
2017-18	A/Hong Kong/4801/2014	A/Texas/277/2017	A/Guangdong/264/2016	8	0
2018-19	A/Singapore/INFIMH-16-0019/2016	A/Japan/NHRC_FDX70352/2018	A/Texas/277/2017	4	3
2019-20	A/Switzerland/8060/2017	A/Washington/9757/2019			10
2020-21	A/South Australia/34/2019	A/Bangladesh/1004005/2020	A/Washington/9757/2019	1	13
2021-22	A/Hong Kong/2671/2019	A/India/PUN-NIV301718/2021	A/Darwin/11/2021	6	1
2022-23	-1	-1	A/Texas/12723/2022		-1

<sup>\*</sup> Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric

SI Tab. 12 H1N1 NA NORTHERN HEMISPHERE (MULTI-CLUSTER)

Year	WHO Recommendation	WHO Error	Qnet Error 1	Qnet Error 2	Qnet Recommendation 1	Qnet Recommendation 2
2001-02	A/New Caledonia/20/99	4	1	6	A/New South Wales/26/2000	A/Canterbury/37/2000
2002-03	A/New Caledonia/20/99	1	0	5	A/Wellington/1/2001	A/New York/447/2001
2003-04	A/New Caledonia/20/99	3	2	8	A/Paris/0833/2002	A/Taiwan/141/2002
2004-05	A/New Caledonia/20/99	2	3	4	A/Memphis/5/2003	A/Hanoi/1004/2003
2005-06	A/New Caledonia/20/99	3	0	1	A/Denmark/130/2004	A/Paris/650/2004
2006-07	A/New Caledonia/20/99	4	2	8	A/Sofia/361/2005	A/Wellington/11/2005
2007-08	A/Solomon Islands/3/2006	9	4	8	A/Sofia/246/2006	A/New York/8/2006
2008-09	A/Brisbane/59/2007	0	13	19	A/Tennessee/UR06-0151/2007	A/Ohio/UR06-0178/2007
2009-10	A/Brisbane/59/2007	87	88	90	A/Sendai/TU66/2008	A/Japan/618/2008
2010-11	A/California/7/2009	2	1	6	A/South Carolina/WRAIR1645P/2009	A/Wisconsin/629-D00809/2009
2011-12	A/California/7/2009	4	1	3	A/England/21680633/2010	A/Hangzhou/178/2010
2012-13	A/California/7/2009	4	1	22	A/Joshkar-Ola/CRIE-BLP/2011	A/Rio Grande do Sul/578/2011
2013-14	A/California/7/2009	5	4	13	A/Thailand/MR10580/2012	A/Mexico/INMEGEN-INER 15/2012
2014-15	A/California/7/2009	9	3	7	A/Minnesota/02/2013	A/Helsinki/430/2013
2015-16	A/California/7/2009	14	4	7	A/Helsinki/808M/2014	A/Virginia/NHRC430739/2014
2016-17	A/California/7/2009	14	0	3	A/Michigan/45/2015	A/Colorado/30/2015
2017-18	A/Michigan/45/2015	3	3	8	A/Mexico/4436/2016	A/Arizona/03/2016
2018-19	A/Michigan/45/2015	4	0	4	A/California/NHRC_QV11073/2017	A/Minnesota/35/2017
2019-20	A/Brisbane/02/2018	1	0	2	A/Kenya/47/2018	A/Colorado/7682/2018
2020-21	A/Hawaii/70/2019	0	3	8	A/California/NHRC-OID_BOX-ILI- 0012/2019	A/Indiana/30/2019
2021-22	A/Victoria/2570/2019	1	5	51	A/Togo/0071/2021	A/Yunnan-Mengzi/1462/2020
2022-23	-1	-1	-1	-1	A/Netherlands/10646/2022	A/Sydney/234/2022

<sup>\*</sup> Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric

SI Tab. 13 H1N1 NA SOUTHERN HEMISPHERE (MULTI-CLUSTER)

Year	WHO Recommendation	WHO Error	Qnet Error 1	Qnet Error 2	Qnet Recommendation 1	Qnet Recommendation 2
2001-02	A/New Caledonia/20/99	4	1	6	A/New South Wales/26/2000	A/Canterbury/37/2000
2002-03	A/New Caledonia/20/99	1	0	5	A/Wellington/1/2001	A/New York/447/2001
2003-04	A/New Caledonia/20/99	3	2	8	A/Paris/0833/2002	A/Taiwan/141/2002
2004-05	A/New Caledonia/20/99	2	3	4	A/Memphis/5/2003	A/Hanoi/1004/2003
2005-06	A/New Caledonia/20/99	3	0	1	A/Denmark/130/2004	A/Paris/650/2004
2006-07	A/New Caledonia/20/99	4	2	8	A/Sofia/361/2005	A/Wellington/11/2005
2007-08	A/New Caledonia/20/99	4	4	8	A/Sofia/246/2006	A/New York/8/2006
2008-09	A/Solomon Islands/3/2006	15	13	19	A/Tennessee/UR06-0151/2007	A/Ohio/UR06-0178/2007
2009-10	A/Brisbane/59/2007	87	88	90	A/Sendai/TU66/2008	A/Japan/618/2008
2010-11	A/California/7/2009	2	1	6	A/South Carolina/WRAIR1645P/2009	A/Wisconsin/629-D00809/2009
2011-12	A/California/7/2009	4	1	3	A/England/21680633/2010	A/Hangzhou/178/2010
2012-13	A/California/7/2009	4	1	22	A/Joshkar-Ola/CRIE-BLP/2011	A/Rio Grande do Sul/578/2011
2013-14	A/California/7/2009	5	4	13	A/Thailand/MR10580/2012	A/Mexico/INMEGEN-INER 15/2012
2014-15	A/California/7/2009	9	3	7	A/Minnesota/02/2013	A/Helsinki/430/2013
2015-16	A/California/7/2009	14	4	7	A/Helsinki/808M/2014	A/Virginia/NHRC430739/2014
2016-17	A/California/7/2009	14	0	3	A/Michigan/45/2015	A/Colorado/30/2015
2017-18	A/Michigan/45/2015	3	3	8	A/Mexico/4436/2016	A/Arizona/03/2016
2018-19	A/Michigan/45/2015	4	0	4	A/California/NHRC_QV11073/2017	A/Minnesota/35/2017
2019-20	A/Michigan/45/2015	4	0	2	A/Kenya/47/2018	A/Colorado/7682/2018
2020-21	A/Brisbane/02/2018	5	2	7	A/California/NHRC-OID_BOX-ILI- 0012/2019	A/Indiana/30/2019
2021-22	A/Victoria/2570/2019	1	7	58	A/Togo/0155/2021	A/Shandong/00204/2021
2022-23	-1	-1	-1	-1	A/Switzerland/86136/2022	A/Wisconsin/04/2021

<sup>\*</sup> Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric

SI Tab. 14 H3N2 NA NORTHERN HEMISPHERE (MULTI-CLUSTER)

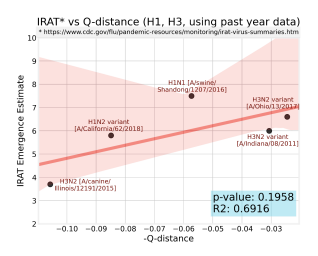
Year	WHO Recommendation	WHO Error	Qnet Error 1	Qnet Error 2	Qnet Recommendation 1	Qnet Recommendation 2
2003-04	A/Moscow/10/99	13	4	5	A/Auckland/612/2002	A/New York/87/2002
2004-05	A/Fujian/411/2002	3	16	18	A/New York/20/2003	A/New York/12/2003
2005-06	A/California/7/2004	4	1	7	A/New York/358/2004	A/Singapore/36/2004
2006-07	A/Wisconsin/67/2005	1	3	8	A/Macau/557/2005	A/Hong Kong/HKU53/2005
2007-08	A/Wisconsin/67/2005	8	0	10	A/Wisconsin/42/2006	A/Wisconsin/44/2006
2008-09	A/Brisbane/10/2007	3	4	10	A/Missouri/06/2007	A/Japan/72/2007
2009-10	A/Brisbane/10/2007	3	1	7	A/Wisconsin/24/2008	A/Mississippi/UR07-0042/2008
2010-11	A/Perth/16/2009	2	3	8	A/New York/70/2009	A/Japan/883/2009
2011-12	A/Perth/16/2009	3	2	2	A/California/19/2010	A/Virginia/05/2010
2012-13	A/Victoria/361/2011	4	1	12	A/Texas/14/2011	A/Singapore/GP1684/2011
2013-14	A/Victoria/361/2011	3	1	5	A/Idaho/38/2012	A/Pavia/135/2012
2014-15	A/Texas/50/2012	3	1	1	A/Nevada/05/2013	A/Michigan/02/2013
2015-16	A/Switzerland/9715293/2013	3	0	4	A/Nicaragua/6866_14/2014	A/Iran/91244/2014
2016-17	A/Hong Kong/4801/2014	7	1	25	A/New Jersey/13/2015	A/California/NHRC_BRD41056N/2015
2017-18	A/Hong Kong/4801/2014	9	1	4	A/Guangdong/264/2016	A/Victoria/668/2016
2018-19	A/Singapore/INFIMH-16- 0019/2016	3	2	4	A/Netherlands/3530/2017	A/Washington/17/2017
2019-20	A/Kansas/14/2017	3	4	10	A/England/538/2018	A/California/BRD12490N/2018
2020-21	A/Hong Kong/2671/2019	3	1	13	A/England/9738/2019	A/Washington/9757/2019
2021-22	A/Cambodia/e0826360/2020	2	3	7	A/Laos/527/2021	A/Michigan/UOM10045655748/2020
2022-23	-1	-1	-1	-1	A/Maine/02/2022	A/Michigan/UOM10042819294/2021

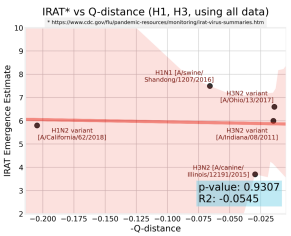
<sup>\*</sup> Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric

SI Tab. 15 H3N2 NA SOUTHERN HEMISPHERE (MULTI-CLUSTER)

Year	WHO Recommendation	WHO Error	Qnet Error 1	Qnet Error 2	Qnet Recommendation 1	Qnet Recommendation 2
2003-04	A/Moscow/10/99	13	4	5	A/Auckland/612/2002	A/New York/87/2002
2004-05	A/Fujian/411/2002	3	16	18	A/New York/20/2003	A/New York/12/2003
2005-06	A/Wellington/1/2004	2	1	7	A/New York/358/2004	A/Singapore/36/2004
2006-07	A/California/7/2004	3	3	8	A/Macau/557/2005	A/Hong Kong/HKU53/2005
2007-08	A/Wisconsin/67/2005	8	0	10	A/Wisconsin/42/2006	A/Wisconsin/44/2006
2008-09	A/Brisbane/10/2007	3	4	10	A/Missouri/06/2007	A/Japan/72/2007
2009-10	A/Brisbane/10/2007	3	1	7	A/Wisconsin/24/2008	A/Mississippi/UR07-0042/2008
2010-11	A/Perth/16/2009	2	3	8	A/New York/70/2009	A/Japan/883/2009
2011-12	A/Perth/16/2009	3	2	2	A/California/19/2010	A/Virginia/05/2010
2012-13	A/Perth/16/2009	4	1	12	A/Texas/14/2011	A/Singapore/GP1684/2011
2013-14	A/Victoria/361/2011	3	1	5	A/Idaho/38/2012	A/Pavia/135/2012
2014-15	A/Texas/50/2012	3	1	1	A/Nevada/05/2013	A/Michigan/02/2013
2015-16	A/Switzerland/9715293/2013	3	0	4	A/Nicaragua/6866_14/2014	A/Iran/91244/2014
2016-17	A/Hong Kong/4801/2014	7	1	25	A/New Jersey/13/2015	A/California/NHRC_BRD41056N/2015
2017-18	A/Hong Kong/4801/2014	9	1	4	A/Guangdong/264/2016	A/Victoria/668/2016
2018-19	A/Singapore/INFIMH-16- 0019/2016	3	2	4	A/Netherlands/3530/2017	A/Washington/17/2017
2019-20	A/Switzerland/8060/2017	10	4	10	A/England/538/2018	A/California/BRD12490N/2018
2020-21	A/South Australia/34/2019	1	1	13	A/England/9738/2019	A/Washington/9757/2019
2021-22	A/Hong Kong/2671/2019	6	1	49	A/Darwin/11/2021	A/Hawaii/28/2020
2022-23	-1	-1	-1	-1	A/Congo/313/2021	A/Texas/12723/2022

<sup>\*</sup> Dominant strain is calculated as the one closest to the centroid in the strain space that year in the edit distance metric





SI Fig. 3. **IRAT vs. Q-distance relationship for H1- and H3- sub-types, using past year data vs. using all data.** On the result when computing average q-distance between the target strain and the circulating human strains from the past year, and on the right is the result when using all available human strains of that sub-type. Evidently, the former has a much higher correlation, since a strain being "close" to humans at some point does not necessarily mean being close now.

SI Tab. 16
INFLUENZA A STRAINS EVALUATED BY IRAT AND CORRESPONDING QNET COMPUTED RISK SCORES

		1	IDAT	1		1	1	I	1
Influenza Virus	Subype	IRAT Date	IRAT Emer- gence Score	IRAT Impact Score	HA Qnet Sample	NA Qnet Sample	HA Avg. Q-dist	NA Avg. Q-dist.	Both Avg. Q-dist.
A/swine/Shandong/1207/2016	H1N1	Jul 2020	7.5	6.9	1000	1000	0.094135	0.020530	0.057332
A/duck/New York/1996	H1N1	Nov 2011	2.3	2.4	1000	1000	-1	-1	-1
A/California/62/2018	H1N2	Jul 2019	5.8	5.7	55	55	0.108902	0.060951	0.084926
A/Ohio/13/2017	H3N2	Jul 2019	6.6	5.8	1000	1000	0.018431	0.030623	0.024527
A/Indiana/08/2011	H3N2	Dec 2012	6.0	4.5	1000	1000	0.052311	0.009103	0.030707
A/canine/Illinois/12191/2015	H3N2	Jun 2016	3.7	3.7	1000	1000	0.060665	0.150856	0.105761
A/American wigeon/South Carolina/AH0195145/2021	H5N1	Mar 2022	4.4	5.1	335	323	0.172180	0.511387	0.337368
A/American green-winged teal/Washington/1957050/2014	H5N1	Mar 2015	3.6	4.1	326	314	0.191127	0.448228	0.316856
A/Vietnam/1203/2004	H5N1	Nov 2011	5.2	6.6	258	246	0.167254	0.011074	0.091342
A/Northern pintail/Washington/40964/2014	H5N2	Mar 2015	3.8	4.1	-1	-1	-1	-1	-1
A/Sichuan/06681/2021	H5N6	Oct 2021	5.3	6.3	45	45	0.361591	0.051794	0.206692
A/Yunnan/14564/2015	H5N6	Apr 2016	5.0	6.6	16	16	-1	-1	-1
A/Astrakhan/3212/2020	H5N8	Mar 2021	4.6	5.2	-1	-1	-1	-1	-1
A/gyrfalcon/Washington/41088/2014	H5N8	Mar 2015	4.2	4.6	-1	-1	-1	-1	-1
A/Netherlands/219/2003	H7N7	Jun 2012	4.6	5.8	46	46	0.275671	0.352060	0.313455
A/turkey/Indiana/1573-2/2016	H7N8	Jul 2017	3.4	3.9	-1	-1	-1	-1	-1
A/chicken/Tennessee/17-007431-3/2017	H7N9	Oct 2017	3.1	3.5	496	495	0.102686	0.256855	0.179693
A/chicken/Tennessee/17-007147-2/2017	H7N9	Oct 2017	2.8	3.5	496	495	0.209532	0.254090	0.231788
A/Hong Kong/125/2017	H7N9	May 2017	6.5	7.5	437	437	0.029599	0.005775	0.017687
A/Shanghai/02/2013	H7N9	Apr 2016	6.4	7.2	178	178	0.005495	0.003556	0.004525
A/Bangladesh/0994/2011	H9N2	Feb 2014	5.6	5.4	13	12	-1	-1	-1
A/Anhui-Lujiang/39/2018	H9N2	Jul 2019	6.2	5.9	31	30	0.029024	0.168090	0.098557
A/Jiangxi-Donghu/346/2013	H10N8	Feb 2014	4.3	6.0	-1	-1	-1	-1	-1

 <sup>\* -1</sup> indicates missing data, either from lack of human sequence data available for that virus sub-type (less than 30 strains) or missing IRAT sequence data (in the case of A/duck/New York/1996)

SI Tab. 17
GENERAL LINEAR MODEL FOR EVALUATING EFFECT OF DATA DIVERSITY ON QNET PERFORMANCE

Variable Name	Description
qnet_complexity	Cumulative number of nodes in all predictors in the corresponding Qnet
data_diversity	Number of clusters in set of input sequence where each sequence in a specific cluster is separated by at least 5 mutations from sequences not in the cluster
ldistance_WHO	Deviation of WHO predicted strain from the dominant strain

=============	===========		
Dep. Variable:	der	No. Observations:	235
Model:	GLI	M Df Residuals:	230
Model Family:	Gaussian	n Df Model:	4
Link Function:	identity	Scale:	23.214
Method:	IRL		-700.43
Date:	Thu, 11 Jun 2020	Deviance:	5339.2
Time:	16:45:40	Pearson chi2:	5.34e+03
No. Iterations:	;	Covariance Type:	nonrobust

	coef	std err		P> z	[0.025	0.975]
Intercept qnet_complexity data_diversity qnet_complexity:data_diversity ldistance_WHO	-0.1116	1.090	-0.102	0.918	-2.248	2.025
	0.0005	0.000	1.075	0.282	-0.000	0.001
	0.3197	0.126	2.531	0.011	0.072	0.567
	-6.932e-05	5.01e-05	-1.383	0.167	-0.000	2.89e-05
	-0.0348	0.035	-1.007	0.314	-0.102	0.033

Dep. Variable:		dev	No. Observations:	235
Model:		GLM	Df Residuals:	231
Model Family:		Gaussian	Df Model:	3
Link Function:		identity	Scale:	23.306
Method:		IRLS	Log-Likelihood:	-701.41
Date:	Thu,	11 Jun 2020	Deviance:	5383.6
Time:		16:45:47	Pearson chi2:	5.38e+03
No. Iterations:		3	Covariance Type:	nonrobust

	coef	std err	Z	P> z	[0.025	0.975]
Intercept qnet_complexity data_diversity ldistance_WHO	1.0841 -4.12e-05 0.1788 -0.0695	0.665 0.000 0.075 0.024	1.630 -0.156 2.392 -2.930	0.103 0.876 0.017 0.003	-0.219 -0.001 0.032 -0.116	2.387 0.000 0.325 -0.023

SI Tab. 18 Numbering Conversion to PDM09 and H3 Schemes

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81         64         73           82         65         74           83         66         75           84         67         76           85         68         77           86         69         72           86         77         70         75           88         71         80           89         72         81         72           80         77         83         82           80         76         84         84           84         77         85         85           85         78         86         79         87           80         88         81         88         81         88           86         79         87         80         88         89         86         93         90         90         90         97         90         90         97         90         93         96         93         96         93         96         99         99         99         99         99         99         99         99         99         99         99         99         99         99         <	33 34 35 35 37 37 37 37 37 37 37 37 37 37 37 37 37
83         66         75           34         67         76           34         67         76           35         68         77           36         69         78           37         70         79           38         71         80           39         72         81           39         72         81           30         73         82           31         74            32         75         83           33         76         84           34         77         85           36         79         87           86         79         87           87         86         79           87         88         86           39         82         90           100         83         91           101         84         92           102         85         -           103         86         93           104         87         94           105         88         95           106         89         99	3 3 3 3 3 3 4 5 5 7 7 3 3 4 5 7 7 7 3 3 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9
84         67         76           85         68         77           86         69         76           86         69         77           86         69         77           86         77         70           87         70         75           88         71         80           99         72         81           90         73         82           91         74         -           92         75         83           93         76         84           94         77         85           95         78         86           96         79         87           97         80         88           88         81         88           98         81         82           900         83         91           902         85         -           103         86         93           104         87         94           105         88         99           106         89         96           107         90         97<	33 39 31 33 34 35 35 37 37 38 39 30 30 30 30 30 30 30 30 30 30 30 30 30
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Query	H1N1pdm	НЗ
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