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## Influenza Resurgence after Relaxation of Public Health and Social Measures, Hong Kong, 2023

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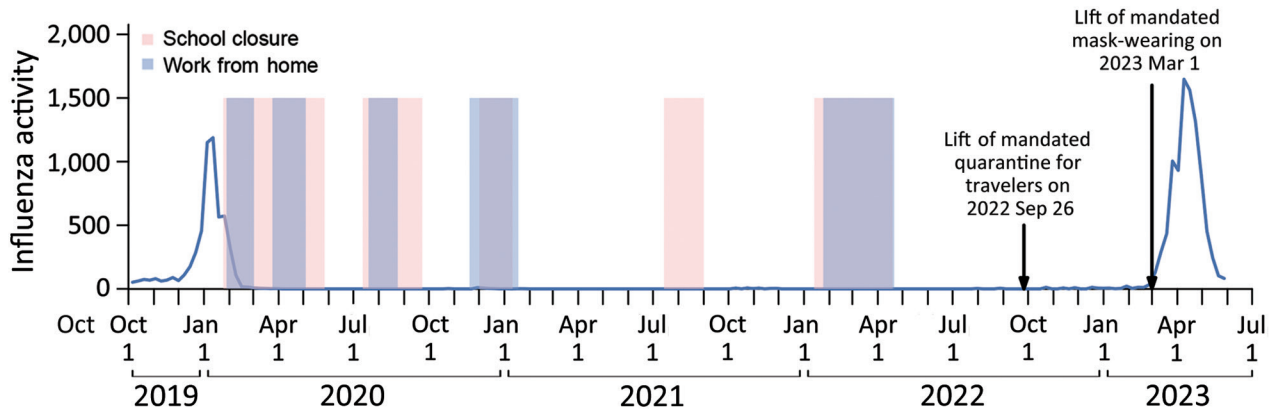
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Soon after a mask mandate was relaxed (March 1, 2023), the first post-COVID-19 influenza season in Hong Kong lasted 12 weeks. After other preventive measures were accounted for, mask wearing was associated with an estimated 25% reduction in influenza transmission. Influenza resurgence probably resulted from relaxation of mask mandates and other measures.

To control COVID-19, Hong Kong, China, put in place several public health and social measures (PHSMs), including mandatory mask wearing, school closures, hand hygiene, and avoidance of gatherings. In early 2020, those measures also reduced influenza transmission (1), and according to laboratory surveillance records, influenza virus did not circulate in the community for 3 years (2). From mid-2022 through 2023, PHSMs were progressively relaxed, and on March 1, 2023, the local mask mandate was lifted. We investigated the effects of PHSMs on influenza transmission in Hong Kong.

We collected weekly influenza-like illness consultation rates reported by private general practitioners and the weekly proportion of sentinel respiratory specimens that tested positive for influenza virus in Hong Kong during October 2010–May 2023. We established a proxy for influenza virus activity by multiplying rates of influenza-like illness by the proportion of influenza-positive samples following previous studies (3,4) (Appendix, <https://wwwnc.cdc.gov/EID/article/29/12/23-0937-App1.pdf>). We found that weekly influenza activity had decreased to almost zero since March 2020, when PHSMs against COVID-19 began (Figure). Before mandatory on-arrival quarantine of travelers started on September 26, 2022, only sporadic influenza-positive samples were detected by surveillance, all from travelers or children who had recently received live-attenuated influenza vaccine (5). After travel restrictions were removed, sporadic influenza detections increased, but overall



**Figure.** Weekly influenza activity and several preventive measures against COVID-19 in Hong Kong during 2020–2023. The blue line represents the weekly influenza activity, measured by the product of influenza-like illness rates and laboratory detections of influenza.

activity remained low. After mandatory indoor and outdoor mask wearing restrictions were lifted on March 1, 2023, influenza transmission increased substantially; the first influenza season after COVID-19 in Hong Kong started and peaked on April 9, ended on May 25, and lasted for 12 weeks (6).

Because various other PHSMs were implemented concurrently with the mask mandate, resurgence of influenza activity could not be attributed to relaxation of the mask mandate alone. Therefore, we used a previous approach that estimated the time-varying effective reproductive number ( $R_t$ ) (7) and a multivariable log-linear regression model on  $R_t$  that could allow for adjustment of other factors affecting influenza transmission, including depletion of susceptible persons, seasonal differences, and meteorologic predictors and preventive measures (Appendix). Because the

predominating influenza strain in 2023 was influenza A(H1N1)pdm09, we identified previous influenza A(H1N1)pdm09 epidemics that had occurred during 2010–2020. To construct a preventive score, we used data from cross-sectional telephone surveys among the general adult population in Hong Kong from 2020 to 2023 as a proxy for the intensity of preventive measures, other than mask wearing, against COVID-19 (1). The preventive score included the average proportion of persons who avoided visiting crowded places, avoided going to healthcare facilities, avoided touching public objects, or used protective measures when touching public objects, and washed hands immediately after going out. Before 2020, the proportion of those preventive measures was established as baseline. When constructing a preventive score, we compared the Akaike information criterion of 4

**Table.** Effects of public health and social measures to protect against COVID-19 on  $R_t$  for influenza, Hong Kong, 2010–2023\*

Model†	PHSM description	% Change in $R_t$ (95% CI)	$\Delta$ AIC‡
Model 1			3.62
Mask		-25 (-43 to -1)	
Preventive score 1	Avoid social gatherings. Wash hands after being outside. Avoid touching or use protective measures with shared objects.	-82 (-91 to -63)	
Model 2			9.55
Mask		-26 (-44 to -2)	
Preventive score 2	Avoid going out as much as possible. Wash hands after being outside. Avoid touching or use protective measures with shared objects.	-80 (-91 to -55)	
Model 3 (main model)			0
Mask		-25 (-43 to -1)	
Preventive score 3	Avoid going to crowded places. Avoid going to healthcare facilities. Avoid touching or use protective measures with shared objects.	-77 (-88 to -60)	
Model 4			2.79
Mask		-24 (-43 to 0)	
Preventive score 4	Avoid going to crowded places. Avoid going to healthcare facilities. Avoid touching or use protective measures with shared objects. Wash hands after being outside.	-81 (-90 to -62)	

\*AIC, Akaike information criterion;  $R_t$ , time-varying effective reproductive number.

†Models were adjusted for depletion of susceptible persons, between-season effects, and absolute humidity.

‡ $\Delta$ AIC<sub>model*i*</sub> = AIC<sub>model*i*</sub> - AIC<sub>min</sub>, AIC<sub>min</sub> = min(AIC<sub>model*i*</sub>),  $i = 1, \dots, 4$ .

combinations of those protective measures. Meteorologic variables provided by the Hong Kong Observatory (<http://hko.gov.hk>) were temperature, wind speed, and relative and absolute humidity. To quantify the effects of meteorologic variables, we fitted the models to data before the COVID-19 pandemic.

Among the 9 epidemics of 2010–2023, the estimated  $R_t$  varied from 0.62 to 1.38 (median 1.02) (Appendix Figure 1). The estimated  $R_t$  showed a decreasing pattern in each season, ranging from  $\approx 1.2$  at the beginning of an epidemic period to 0.8 at the end of an epidemic period. After model selection (Appendix), we found that a model of absolute humidity, mask wearing, and preventive score 3 (Table) explained 92% of the observed variance in estimated  $R_t$  (Appendix Table 1). Changes in absolute humidity (Appendix Figure 2, panel A), the proportion of mask wearing, and preventive score 3 (Appendix Figure 2, panel B) strongly correlated with changes in  $R_t$ . After adjusting for other factors, such as depletion of susceptible persons, between-season effects, and absolute humidity, we found that mask wearing was associated with a 25% (range 1%–43%) reduction in  $R_t$  and that other preventive measures (combined) were associated with a 77% (range 60%–88%) reduction (Table).

We found that that influenza increased after PHSMs were relaxed and influenza transmission increased shortly after the mask mandate was relaxed. Our results are consistent with those of several studies that found that PHSMs against COVID-19 may reduce influenza transmission (8) and that mask wearing may have a low to moderate protective effect against influenza virus transmission in the community (9,10).

A limitation of our analysis was that we used results of survey reports to generate a proxy of intensity of implemented PHSMs over time, which may not be accurate. Also, we used a proxy measure of influenza activity based on surveillance data, and the reliability of our analysis depended on the accuracy of this proxy. In addition, influenza vaccination coverage (Appendix Figure 5) was not included in the model because our model included the effect of vaccination via season-specific intercept. Nevertheless, our study results suggest that the resurgence of influenza after relaxation of PHSMs was most likely affected by the lifting of mask mandate and other PHSMs.

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## SARS-CoV-2 Variants BQ.1 and XBB.1.5 in Wastewater of Aircraft Flying from China to Denmark, 2023

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We analyzed wastewater samples from 14 aircraft arriving in Denmark directly from China during January 9–February 12, 2023. Wastewater from 11 aircraft was SARS-CoV-2–positive by PCR; 6 predominantly contained BQ.1 and XBB.1 subvariants. Wastewater-based surveillance can contribute to public health monitoring of SARS-CoV-2 and other emerging infectious agents.

Relaxation of China's zero-COVID policy in December 2022 led the European Centre for Disease Prevention and Control to recommend several nonpharmaceutical interventions to curb COVID-19 spread and monitor any emerging SARS-CoV-2 variants; those interventions included wastewater-based surveillance (1). We report results of subsequent wastewater surveillance of aircraft arriving at Copenhagen Airport in Copenhagen, Denmark, directly from Beijing or Shanghai, China.

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During weeks 2–6 of 2023 (January 9–February 12), a total of 14 aircraft arrived at Copenhagen Airport from China. A service truck extracted waste from the aircraft by using vacuum pressure, after which a rinsing program was performed, and the disinfectant Idu-Flight (Brenntag Nordic A/S, <https://www.brenntag.com>) was added to the waste tank. Wastewater samples were collected as grab samples from the service truck and immediately transported to Statens Serum Institut in Copenhagen for analysis.

The pH value of the sample material ranged from 9–10 because of the addition of Idu-Flight. Idu-Flight contains the active ingredients glutaraldehyde and benzalkonium chloride; the disinfectant is expected to negatively affect the stability of virus particles and hinder amplification of RNA sequences. We adjusted the samples to pH 7.5–8.5 by using HCl and homogenized them by vigorous vortexing. We split the 14 samples into a total of 43 aliquots and then centrifuged those at either 4,000 × g or 10,000 × g for 10 min to pellet solid material. For the first aliquot from aircraft AC1, we analyzed 10 mL of sample material without any centrifugation; for all other samples, we analyzed 10 mL of supernatant after centrifugation. We purified viruses by using NanoTrap Microbiome A particles (Ceres Nanosciences Inc., <https://www.ceresnano.com>) and RNA by using Maxwell RSC Cartridges (Promega Corporation, <https://www.promega.com>). We performed quantitative reverse transcription PCR (qRT-PCR) in technical triplicate by using the GoTaq Enviro kit (Promega) and the US Centers for Disease Control and Prevention N2 primer/probe for SARS-CoV-2 detection (Table; Appendix Table, <https://wwwnc.cdc.gov/EID/article/29/12/23-0717-App1.pdf>).

Of the 43 qRT-PCR reactions, 31 (72%) were positive for SARS-CoV-2, representing 11 aircraft. We conducted whole-genome sequencing of samples from those 11 aircraft by using the Illumina MiSeq platform (<https://www.illumina.com>) according to the ARTIC protocol; we generated 2 × 150-bp paired-end reads by using the ARTIC 4.1 primer scheme (2). Wastewater raw reads are available from the European Nucleotide Archive (<https://www.ebi.ac.uk/ena>; accession no. PRJEB66221). We trimmed reads by using Trim Galore with default settings (3; <https://zenodo.org/record/5127899>). We removed human sequence reads by using the BWA-MEM alignment algorithm with default settings (H. Li, unpub. data, <http://arxiv.org/abs/1303.3997>) and the human genome reference build GRCh38. We then used BWA-MEM with default settings to map SARS-CoV-2 reads to the SARS-CoV-2 wild-type reference genome (GenBank accession no. MN908947.3). We performed

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## Appendix

### 1 Description of the Time Series

#### 1.1 The time series for ILI

We collected the weekly consultation rates of influenza-like illness (ILI) reported by Private Medical Practitioner (PMP) Clinics ([chp.gov.hk](http://chp.gov.hk)) and the weekly proportion of sentinel respiratory specimens that tested positive for influenza viruses in Hong Kong from October 2010 through May 2023.

To measure the influenza virus activity, we multiplied the ILI rates with the proportions of influenza-positive specimens together to obtain an influenza proxy ( $I$ ). This influenza proxy shows a stronger correlation with the incidence of influenza virus infections in the community than either influenza-like illness rates or laboratory detection rates alone. We then first multiplied the weekly ILI rates by a constant 70, based on the previous record number of general practitioners in the surveillance. In addition, we divided it by 0.9 as a health seeking (HS) proportion for ILI symptoms in Hong Kong (2) and divided it by 0.3 as 30% of influenza cases have ILI symptoms (3). To ensure consistency with expected population-level infection rates, we used the constant to scale up the proxy values (4,5). Finally, we used flexible cubic splines to interpolate daily influenza proxy values from the weekly data.

To identify influenza epidemics, we defined each season's influenza epidemic as a period of at least 12 or more consecutive weeks during which the epidemic baseline was exceeded. The epidemic baseline was determined as 40% quantile of all the non-zero weekly influenza proxy for each influenza season (6).

## **1.2 Time series of meteorological data**

We retrieved 10 meteorological predictors provided by Hong Kong Observatory ([hko.gov.hk](http://hko.gov.hk)), including pressure, temperature, relative humidity, amount of cloud, rainfall, number of hours of reduced visibility, total bright sunshine, global solar radiation, evaporation, and wind speed. Due to high correlations among these variables, we selected temperature, wind speed, and absolute humidity based on previous literature (5,7,8). We derived the daily mean absolute humidity from the mean relative humidity and mean temperature (7,9), and then obtained the daily and weekly absolute humidity.

## **1.3 Time series of preventive measures**

We did cross-sectional telephone surveys among the general adult population in Hong Kong from 2020 to 2023 (10). The methods and survey instruments used were similar to those used for surveys during the SARS epidemic in 2003 (11,12) the influenza A H1N1 pandemic in 2009 , and the influenza A H7N9 outbreak in China in 2013 (13). Participants were recruited using random-digit dialling of both landline and mobile telephone numbers. Telephone numbers were randomly generated by a computer system. Calls were made during both working and non-working hours by trained interviewers to avoid over-representation of non-working groups. Respondents were required to be at least 18 years old and able to speak Cantonese or English. New respondents were recruited for each survey round. Within each household, an eligible household member with the nearest birthday was invited to participate in the survey, which was not necessarily the person that initially answered the telephone. Survey items included measures of risk perception, attitudes towards COVID-19, and preventive measures taken against contracting COVID-19, including hygiene, face masks, and reduction of social contact. All participants gave verbal informed consent. The prevalence of those preventive measures prior to 2020 was set to be the baseline prevalence.

To proxy the intensity of preventive measures against COVID -19 other than mask wearing, we used data from the survey to construct a preventive score (e.g., the average of proportions of people avoiding visiting crowded places, avoiding touching public objects or using protective measures when touching public objects, and washing hands immediately after going out).

## 2 Estimation of time-varying effective reproductive number

### 2.1 Model details

We used the framework in Cori et al (14) to estimate the  $R_t$  from real data. In brief, it assumes that the distribution of infectiousness through time after infection is independent of calendar time. Transmission then is modelled by using a Poisson process. Denote  $w_s$  a probability distribution of the infectiousness profile since infection, therefore the rate for infection at time step  $t-s$  generates new infections in time step  $t$  is equal to  $R_t w_s$ , where  $R_t$  is the instantaneous reproductive number at  $t$ . Also, the incidence at time  $t$  is Poisson distributed with mean  $R_t \sum_{s=1}^t I_{t-s} w_s$ .

Denote  $Y_k$  the actual (but unobserved) number of new local cases infected on day  $k$ . Then, we have:

$$Y_t \sim \text{Poisson}\left\{R_t \sum_{k=1}^{t-1} Y_k w_{t-k}\right\}$$

where  $R_t$  are the time-varying effective reproductive number at time  $t$  respectively.

### 2.2 Likelihood function

We used the smoothing method as in Cori et al., assuming that the transmissibility is constant over a time period  $[t - \tau + 1, t]$ , where  $\tau$  is the smoothing parameter. Hence likelihood at a time period  $t$  is

$$P(Y_t, \dots, Y_{t-\tau+1} | Y_1, \dots, Y_{t-\tau}) = \prod_{s=t-\tau+1}^t \frac{(R_t^\tau \phi(s))^{Y_s} e^{-R_t^\tau \phi(s)}}{Y_s!}$$

where  $\phi(t) = \sum_{k=1}^{t-1} Y_k w_{t-k}$ . The total likelihood is the product of individual likelihood at each time  $t$  in the observed data. The first  $\tau - 1$  days were excluded due to  $\tau$ -day smoothing.

### 2.3 Priors

We assumed the prior for  $R_t$  is Gamma(1,1.5) with mean and standard deviation equal to 1.5.

### 2.4 Estimation of model parameters

We conducted our analysis in a Bayesian framework and used a Markov chain Monte Carlo (MCMC) algorithm to estimate model parameters. At each MCMC step  $k$ , we update the

model parameters  $\theta$  by using random walk Metropolis-Hastings algorithm (15). The step size of the proposal was adjusted to have acceptance rate for 20-30%.

### **2.5 Assumption on input parameter in data analysis**

We use the estimated distribution with mean 2.7 days (16) for serial interval. The empirical distribution of reporting delay would be used for a deconvolution approach by Miller et al. (17) to obtain the epidemic curve by infection time, which was achieved by using the ‘fit\_incidence’ function in the ‘incidental’ package in R.

We analyse the epidemic curve up to 31 May 2023, and take  $\tau = 14$  in our analysis, to avoid unstable estimates for time-varying reproductive number.

### **2.6 Inference**

After obtaining the epidemic curve by infection time, we use the model in Section 2.1 to estimate  $R_t$ . We use a Markov chain Monte Carlo approach to estimate the model parameter, as stated in Section 2.4.

We accounted for the uncertainty of input parameters, including incubation period and infectiousness profile to obtain the final estimates of  $R_t$  in addition to model parameter uncertainty as follows:

We followed the bootstrap approach (18,19) to account for the uncertainty of input parameters, including incubation period, to obtain the final estimates of  $R_t$  in addition to uncertainty of model parameters. In each iteration, we use the above deconvolution approach to reconstruct the epidemic curve by infection dates. Then we use above approach to estimate  $R_t$ . We presented the mean, 2.5% and 97.5% quantiles for those  $R_t$  estimates for each time point across the 200 bootstrap iterations.

## **3 Construction of Multivariable Regression Models**

We used multivariable log-linear regression models to investigate the underlying association between the transmissibility of influenza and different driving factors.

For meteorological factors, we tested different regression forms to investigate the underlying association between the transmissibility of influenza and different plausible driving forces. We compared linear form (*i. e.*  $f_{kt}$ , where  $f_{kt}$  are the  $k$  – th drivers), exponential form



(i. e.  $\varphi(f_{kt})$ , where  $\varphi(f_{kt}) = \exp(f_{kt})$ ), power form (i. e.  $\varphi(f_{kt}) = f_{kt}^2$ ) of associations across all the meteorological drivers with influenza transmissibility (Table S1).

Following the epidemic model theory and the from aforementioned results, we construct a general multivariable nonlinear regression model described by te Beest et al (20). Consider the  $S_{0j}$  is the susceptibles (fraction) at the start of  $i$  th weeks of  $j$  th epidemic and  $R_0$  is the basic reproduction number. Therefore, the instantaneous reproduction number  $R_{ij}$  can be written as

$$\log(R_{ij}) = \log(R_0 S_{0j}) + z_j h_{ij} + \sum_k \beta_k \log(d_{ijk}) + \sum_l \varphi(f_{ijl}) + \epsilon_{ij}$$

$R_{ij}$  is the time varying instantaneous reproduction number on day  $i$  of epidemic  $j$ .  $S_{0j}$  represents the initial fraction of susceptibles at the start of season  $j$ ,  $h_{ij}$  is the observed cumulative incidence of general practitioner consultations by patients with ILI up to week  $i-1$  of season  $j$ , and  $z_j$  is a seasonal effect that adjusts  $h_{ij}$  to the season-effect. In addition, the season-specific intercept could capture the pre-season influenza vaccine effect. The effect of the driving factors ( $d_{ijk}$  or  $f_{ijl}$ ) during week  $i$  for the epidemic  $j$ , is determined by the respective coefficients. We treated the parameters  $\log(R_0 S_{0j})$  and  $z_j$  as the nuisance parameters. The coefficient  $\beta_k$  represents the association between  $R_{ij}$  and  $d_{ijk}$ .  $\epsilon_{ij} \sim N(0, \sigma^2)$  is the error term.

We finally define a baseline model based on intrinsic factors (depletion of susceptibles over time and between-season effects) only.

i.e.  $\log(R_0 S_{0j}) + z_j h_{ij} + \epsilon_{ij}$ .

Improved models including other significant factors were then created (Table S2). R-squared ( $R^2$ ) was used to quantify the effects of each factor. Therefore, these  $\Delta R^2$  measures (comparing the R-square values of these models) indicate the variance in transmissibility explained by respective drivers.

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**Appendix Table 1.** AIC values for the model incorporating intrinsic factors and different forms of meteorological drivers to identify the best form of association of  $R_t$  and meteorological drivers\*

Forms of Association	$\Delta AIC$ for associations of influenza and drivers		
	Mean temperature	Mean wind speed	Mean absolute humidity
Linear	6.89	0.0047	1.39
Exponential	3.08	0.0036	0.43
Power	0	0	0

\*  $\Delta AIC_i = AIC_i - AIC_{min}$ ,  $AIC_{min} = \min(AIC_{Linear}, AIC_{Exponential}, AIC_{Power})$ ,  $i = Linear, Exponential, Power$ .

**Appendix Table 2.** Variance Explained by the Driving Factors of Influenza Transmission in the Hong Kong, 2010 – 2023

Driving Factor	Regression Terms $M^*$	$R^{2†}$	$\Delta R^{2‡}$	df	$R_{adj}^2 §$	P value
Depletion of susceptibles	$zh_{ij}$	0.461	0.461	158	0.458	< 0.001
Between-season effect	$\log(R_0 S_{0j}) + z_j h_{ij}$	0.899	0.438	142	0.887	< 0.001
Absolute humidity	$\log(R_0 S_{0j}) + z_j h_{ij} + \beta_1 \varphi(d_{ij1}) ¶$	0.903	0.004	141	0.891	0.021
Mask	$\log(R_0 S_{0j}) + z_j h_{ij} + \beta_2 d_{ij2}$	0.902	0.003	141	0.900	0.044
Preventive score 3	$\log(R_0 S_{0j}) + z_j h_{ij} + \beta_3 d_{ij3}$	0.912	0.013	141	0.901	1.009e-05
Final model	$\log(R_0 S_{0j}) + z_j h_{ij} + \beta_1 \varphi(d_{ij1}) ¶$ $+ \beta_2 d_{ij2} + \beta_3 d_{ij3}$	0.920	0.008	139	0.908	0.001
School holidays	$\log(R_0 S_{0j}) + z_j h_{ij} + \beta_4 d_{ij4}$	0.902	0.003	141	0.889	0.057
Temperature	$\log(R_0 S_{0j}) + z_j h_{ij} + \beta_5 \varphi(d_{ij5}) ¶$	0.909	0.010	141	0.897	0.0001

\*All fitted regression models take the form  $\log(R_{ij}) = M + \varepsilon$ , and the regression terms  $M$  differ for each model.

†  $R^2$  is the variance of the influenza reproduction numbers that is explained by each model.

‡  $\Delta R^2$  is the proportion of the variance explained by a specific driving factor.

§  $R_{adj}^2$  provides a measure of parsimony for each model.

¶  $\varphi$  is the power form association.

**Appendix Table 3.** Estimates of the Strength of Driving Factors of Final Model on Influenza Transmission in the Hong Kong, 2010–2023

Driving Factor	Variable	Estimate	95% CI	P value
Between-season intercept	Median $[S_{0j}]^*$	0.833	0.635, 1.085	
Between-season depletion of susceptibles	Median $[z_j]^*$	-0.176	-0.222, -0.153	
Absolute humidity	$\beta_1$	-0.051	-0.083, -0.018	0.003
Mask	$\beta_2$	-0.288	-0.563, -0.01	0.039
Preventive score 3	$\beta_3$	-1.497	-2.086, -0.908	<0.001

\* For regression coefficients that were specific for each season, we estimated the range of  $R_0 S_{0j}$  to be 0.794-0.993 and the range of  $z_j$  to be -1.622 to -0.086 .

**Appendix Table 4.** Several non-pharmaceutical interventions (NPIs) included in the study

Control measure	Description	Time period	Notes
Mandate quarantine	Departure location	Compulsory quarantine is required for inbound travellers from specific or all overseas countries/regions.	14 days
			21d ays
			14 or 7 days with two consecutive RAT negatives
			3days
Contact quarantined	Close contacts of confirmed cases are required to be quarantined in a quarantine centre or hotel, regardless of their infection status.	01/01/2020 – 02/07/2022	Camp
		02/08/2022 – 12/28/2022	Camp/home
Case isolated	Confirmed cases are required to be isolated at hospital or isolation facility upon testing positive, regardless of their symptoms.	01/23/2020 – 02/07/2022	Mandate hospital isolation
		02/08/2022 – 01/30/2023	Hospital/isolation facility/home
Community-based	All kindergartens, primary and secondary schools and private schools in Hong Kong should suspend face-to-face classes and all on-campus activities	01/25/2020 – 05/26/2020	Closed
School closure		07/13/2020 – 09/22/2020	
		12/02/2020 - 01/10/2021	
		07/15/2021 – 08/31/2021	
Work-from-home	Special work arrangement for civil servants. Private business was encouraged to follow the work at home arrangements.	01/14/2022 – 04/18/2022	Flexible work, except for essential service
		01/29/2020 – 03/01/2020	
		03/25/2020 – 05/03/2020	
		07/20/2020 – 08/23/2020	
		11/20/2020 – 01/17/2021	
Mask mandate	The mandatory mask-wearing requirement stipulates a person must wear a mask all the time when the person is entering or present in a specified indoor or outdoor public place.	01/25/2022 – 04/20/2022	public indoors public indoors and outdoors
		07/15/2020 – 07/28/2020	
		07/28/2020 - 03/01/2023	

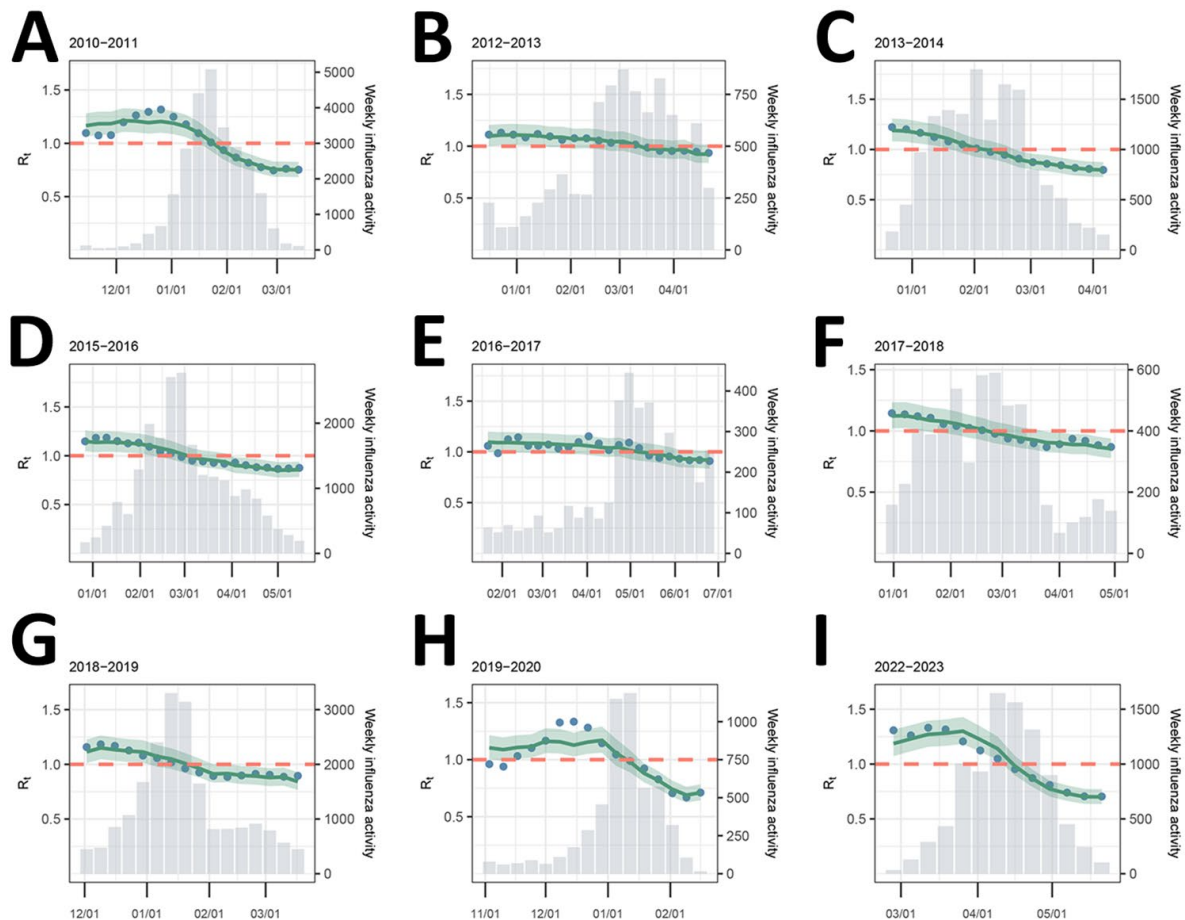
**Appendix Table 5.** Estimates of the strength of driving factors with backward selection on influenza transmission in the Hong Kong, 2010–2023

Driving Factor	Estimate	95% CI	P value
Between-season intercept	0.830	0.641, 1.087	
Between-season depletion of susceptibles	-0.17	-0.19, -0.118	
Temperature	-0.079	-0.115, -0.044	<0.001
Wind speed	-0.031	-0.059, -0.002	0.035
Mask	-0.318	-0.593, -0.045	0.023
Preventive score 3	-1.446	-2.036, -0.855	<0.001

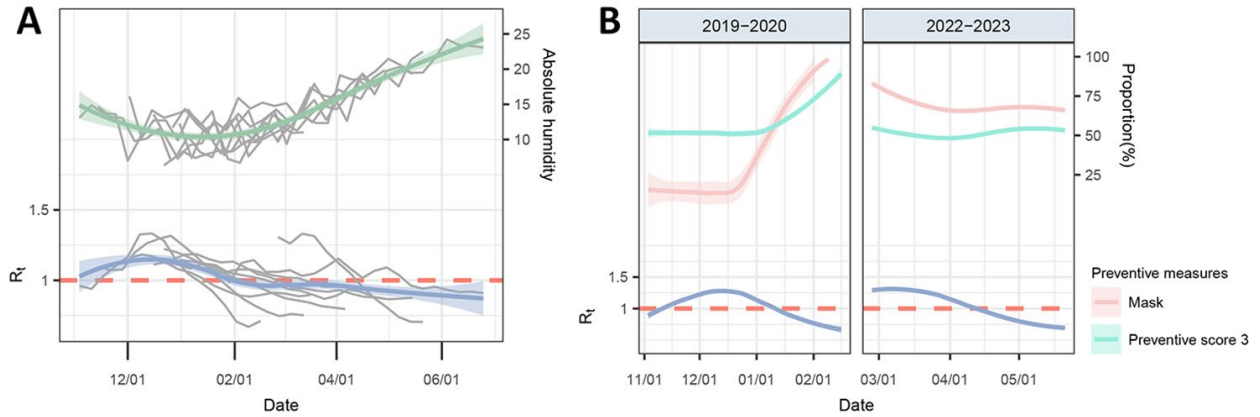
\* For regression coefficients that were specific for each season, we estimated the range of  $R_0S_{0j}$  to be 0.785-1.024 and the range of  $z_j$  to be -1.575 to -0.083 .

**Appendix Table 6.** Effect of public health and social measures against COVID-19 on time-varying reproduction number of influenza with backward selection, Hong Kong, 2010-2023

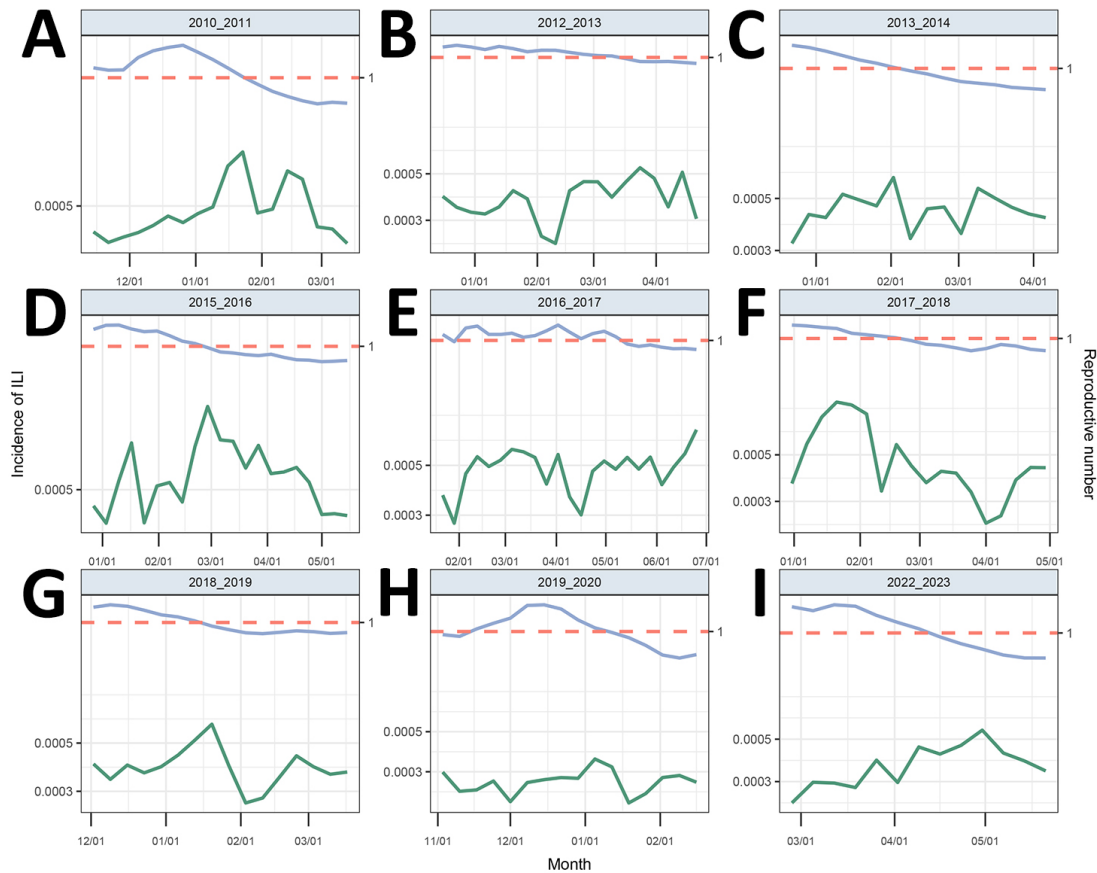
PHSM	Description	% Change in $R_t$ (95% CI)
Mask		-27 (-44 to -4)
Preventive score 3	Avoid going to crowded places Avoid going to health care facilities Avoid touching and protect in public	-76 (-87to -57)



**Appendix Figure 1.** Influenza reproductive number based on the final model. The grey bar represents the weekly influenza activity in 9 epidemic seasons, showing the peak for each season. The green dot represents the reproduction numbers estimated from the weekly influenza proxy series and the green line represents the predicted reproduction numbers. The shaded area represents the 95% predictive interval.

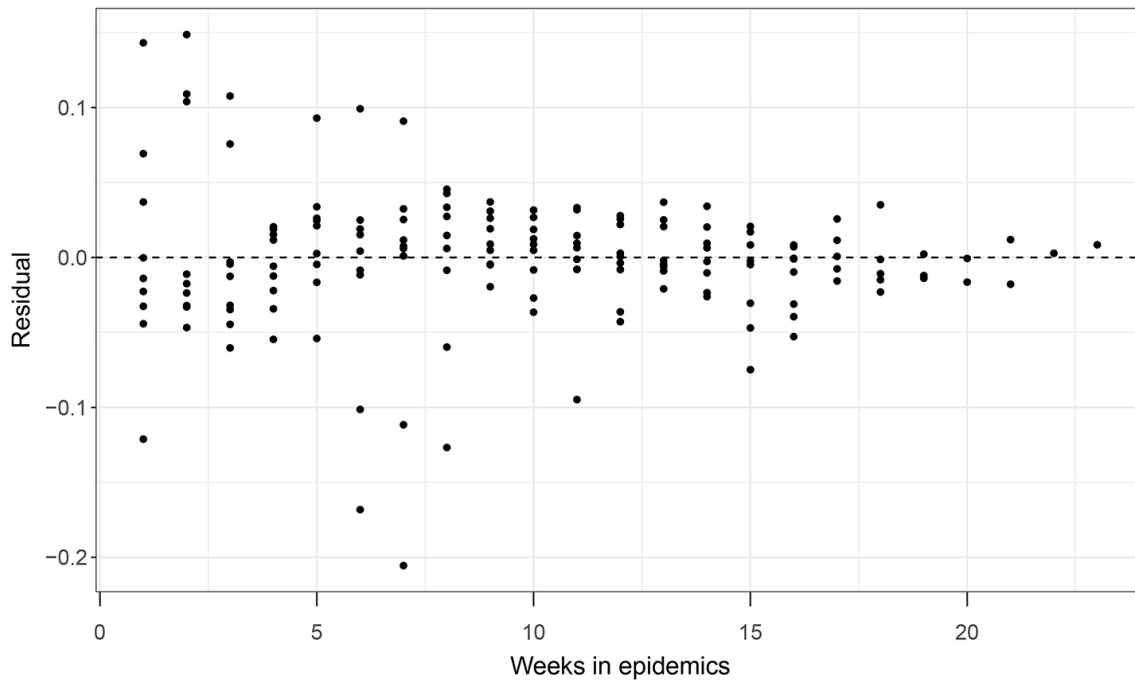


**Appendix Figure 2.** Estimated reproductive number and driving factors. A) The gray lines represent absolute humidity and reproductive number during each season. The green and blue lines represent the smoothed conditional mean of absolute humidity and reproductive number respectively. B) Estimated reproductive number and the proportion of mask wearing and preventive scores during the 2019–20 and 2022–23 influenza seasons.

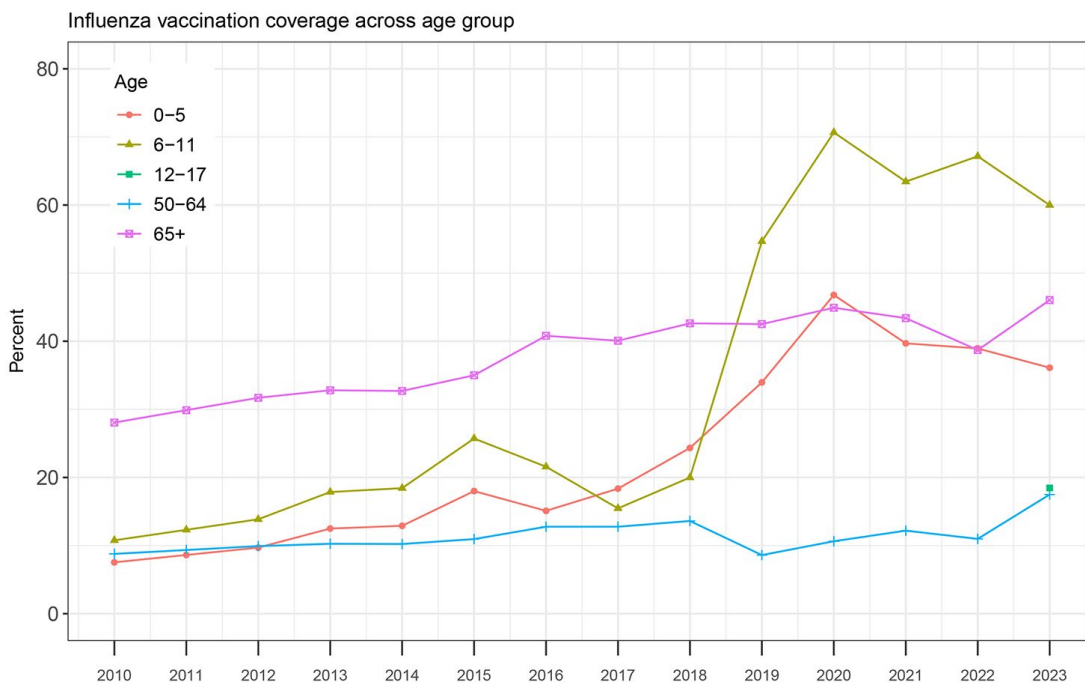


**Appendix Figure 3.** Time series of incidence of influenza-like-illness (ILI) in Hong Kong over 2010–2023 (bottom) and the reproduction number (top) calculated from the ILI proxy.





**Appendix Figure 4.** The residuals of the final model, incorporating depletion of susceptibles over time and between-season effects, absolute humidity, mask and the preventive score, indicating goodness of fit.



**Appendix Figure 5.** Pre-season influenza vaccine coverage by age group in Hong Kong during the study period.