

1 Susceptible host dynamics explain pathogen resilience  
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## 5 **Abstract**

6 Major priority for epidemiological research in the time of anthropogenic change is  
7 understanding how infectious disease dynamics respond to perturbations. Interven-  
8 tions to slow the spread of COVID-19 significantly disrupted the transmission of other  
9 human pathogens, providing unique opportunities to learn about pathogen charac-  
10 teristics from spatiotemporal variation in re-emergence patterns. As interventions  
11 lifted, a key question of whether and when respiratory pathogens would eventually  
12 return to their pre-pandemic dynamics remains to be answered. To address this  
13 gap, we develop a framework for estimating pathogen resilience based on how fast  
14 epidemic patterns return to their pre-pandemic, endemic cycles. Our analysis re-  
15 veals a possibility that some pathogens may have settled to endemic cycles that are  
16 different from their pre-pandemic patterns. Finally, we show that heterogeneity in  
17 pathogen resilience can be understood in terms of how fast a susceptible host popula-  
18 tion becomes replenished. Our framework offers a novel perspective to characterizing  
19 epidemic dynamics of endemic pathogens and measuring epidemic time scales.

Understanding how ecological systems respond to perturbations is a fundamental challenge in predicting species persistence and extinction (Bender et al., 1984; Ives and Carpenter, 2007; Scheffer et al., 2009). These responses are often characterized in terms of resilience, which captures how fast a system returns to its stable, reference state following a perturbation (Pimm, 1979; Neubert and Caswell, 1997; Gunderson, 2000; Dakos and Kéfi, 2022). Both theoretical and empirical efforts to quantify resilience of ecological systems have provided key insights for understanding the dynamics of complex systems and linking these findings to actionable strategies for species conservation. However, despite rich literature on ecological resilience, there have been limited applications to measuring the resilience of host-pathogen systems, especially for human pathogens.

Non-pharmaceutical interventions (NPIs) to slow the spread of COVID-19 disrupted the transmission of other human pathogens, providing large-scale natural experiments for understanding how various host-pathogen systems respond to perturbations (Baker et al., 2020; Gomez et al., 2021; Koltai et al., 2022; Park et al., 2024). In particular, as interventions lifted, large heterogeneities in outbreak dynamics were observed across different pathogens in different countries (Figure 1), likely reflecting differences in NPI patterns, pathogen characteristics, immigration/importation from other countries, and pre-pandemic pathogen dynamics. Even though more than four years have already passed since the emergence of COVID-19, current circulation patterns for many respiratory pathogens appear to be different from their pre-pandemic, seasonal patterns, especially in Hong Kong and Korea: some pathogens, such as human metapneumovirus and bocavirus in Korea, are circulating at lower levels, whereas other pathogens, such as RSV in Korea, seem to exhibit different seasonality (Figure 1). These observations pose two fundamental questions for current and future infectious disease dynamics: (1) can we learn about underlying pathogen characteristics, such as their transmissibility or duration of immunity, from re-emergence patterns? and (2) can we predict whether and when other respiratory pathogens will eventually return to their pre-pandemic dynamics?

To address this question, we propose a framework for characterizing the resilience of a host-pathogen system based on how fast the system recovers from perturbation. We begin by laying out a few example scenarios that represent the expected outbreak dynamics following COVID-19 interventions and illustrating how resilience can be measured by comparing the post- and pre-pandemic dynamics of susceptible and infected hosts. In practice, the dynamics of susceptible hosts are often unavailable, and traditional susceptible reconstruction approaches often require long-term endemic time series, which cannot be applied due to disruptions in epidemic patterns caused by COVID-19 NPIs. Instead, we utilize Takens’ embedding theorem to reconstruct empirical attractors from data and further measure the distance from this empirical attractor, which allows us to characterize the rate at which this distance decreases. Analyses of pathogen surveillance data for a wide array of respiratory pathogens in Canada, Hong Kong, and Korea suggest ... Finally, we show that variation in resilience estimates can be understood in terms of susceptible host dynamics.

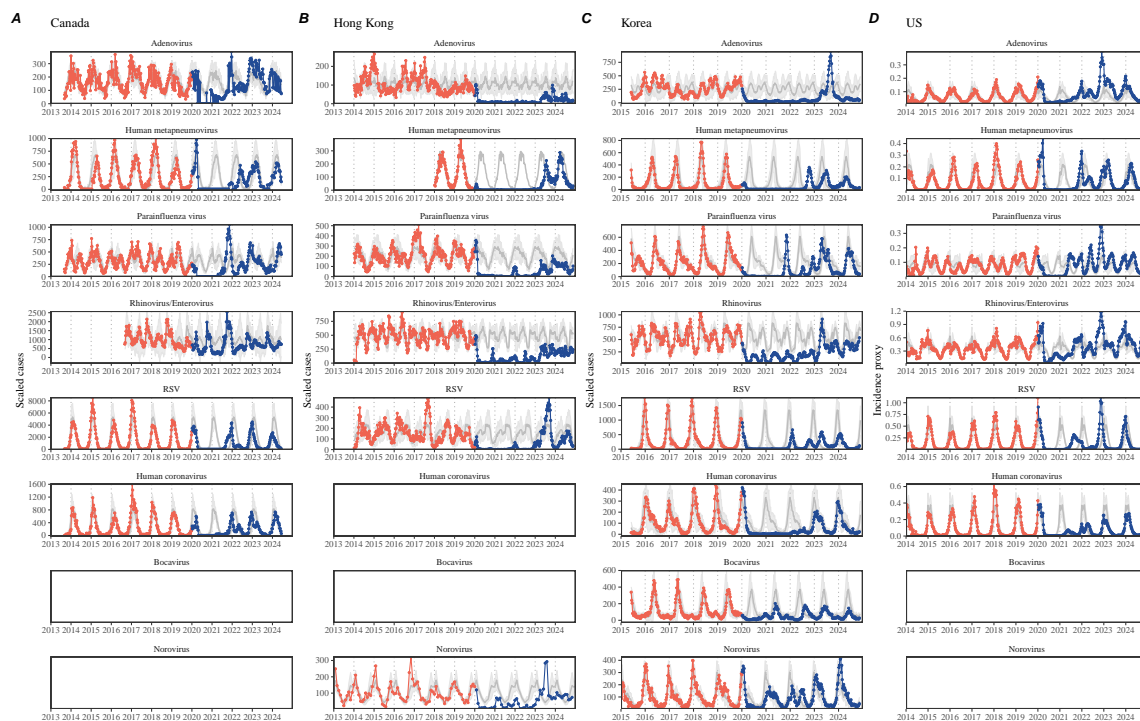


Figure 1: **Observed heterogeneity in responses to COVID-19 pandemic across respiratory pathogens in (A) Canada, (B) Hong Kong, (C) Korea, and (D) US.** Red points and lines represent data before 2020. Blue points and lines represent data since 2020. Gray lines and shaded regions represent the mean seasonal patterns and corresponding 95% confidence intervals based on the observed outbreak patterns before 2020.

63 [SWP: Revisit.]

## 64 Conceptual introduction to pathogen resilience

65 In classical ecological literature, resilience of an ecological system is measured by the  
 66 rate at which the system returns to its reference state following a perturbation. This  
 67 rate corresponds to the largest real part of the eigenvalues of the linearized system  
 68 near equilibrium—here, we refer to this value as the *intrinsic* resilience of the system,  
 69 which represents the expected rate of return from perturbed states. However, res-  
 70 piratory pathogens often exhibit seasonal variation in transmission, meaning that the  
 71 intrinsic resilience of a host-pathogen system varies across season. Nonetheless, we  
 72 can still measure the *empirical* resilience of a host-pathogen system by looking at how  
 73 fast the system returns to the pre-pandemic, endemic dynamics after interventions  
 74 are lifted.

75 As an example, consider an intervention that reduce transmission by 50% for 6  
 76 months starting in 2020, which causes epidemic patterns to deviate from its original

stable annual cycle for a short period of time and eventually come back (Figure 2A). To measure the empirical resilience of this system, we first need to be able to measure the distance from its pre-pandemic attractor. There are many different ways we can measure the distance from attractor, but for illustrative purposes, we choose one of the most parsimonious approach: that is, we look at how the susceptible (S) and infected (I) populations change over time and measure the distance on the SI phase plane (Figure 2B). In this simple case, the locally estimated scatterplot smoothing (LOESS) fit indicates that the distance from attractor decreases linearly on average (Figure 2C). Furthermore, the overall rate of return matches the intrinsic resilience of the seasonally unforced system (Figure 2C).

Alternatively, NPIs can permanently change our behavior and have persisting impact on the pathogen dynamics; as an example, we consider a scenario in which a 10% reduction in transmission persists even after the NPIs are lifted (Figure 2D–F). In such cases, we cannot know whether the pathogen will return to its original cycle or a different cycle until many years have passed after the NPIs are lifted, meaning that we cannot measure the distance against the new attractor that the system will eventually approach. Nonetheless, we can still measure the distance against the original, pre-pandemic attractor and ask how the distance changes over time (Figure 2E). The LOESS fit suggests that the distance from the attractor will initially decrease exponentially on average (equivalently, linearly on a log scale) and eventually plateau (Figure 2F). Here, a permanent 10% reduction in transmission rate slows down the system, which causes the distance from the attractor to decrease at a slower rate (Figure 2F) than it would have otherwise in the absence of permanent transmission reduction (Figure 2C). This example shows that resilience is not necessarily an intrinsic property of a specific pathogen. Instead, pathogen resilience is a property of a specific attractor that a host-pathogen system approaches, which depends on both pathogen and host characteristics. *[SWP: Add discussion about observation error, e.g., under-reporting.]*

Finally, transient phenomena can also complicate the picture (Figure 2G–I). For example, a stage-structured model for RSV initially exhibits a stable annual cycle, but perturbations from NPIs cause the epidemic to exhibit biennial cycles (Figure 2G). In this case, the distance from the attractor initially decreases exponentially at a rate that is consistent with intrinsic resilience, but the rate of decrease slows down as the epidemic exhibits a biennial cycle (Figure 2I). In classical ecological theory, this behavior is also referred to as a ghost attractor, which causes long transient dynamics and slow transitions.

These observations indicate three possibilities. First, we can directly estimate the empirical resilience of a host-pathogen system by looking at how fast the system approaches an attractor, provided that we have a reasonable way of reconstructing the attractor and calculating the distance from the reconstructed attractor. The empirical approach to estimating pathogen resilience is particularly convenient because it does not require us to know the true underlying model. As we show in Supplementary Materials, model misspecification can lead to severe biases in estima-

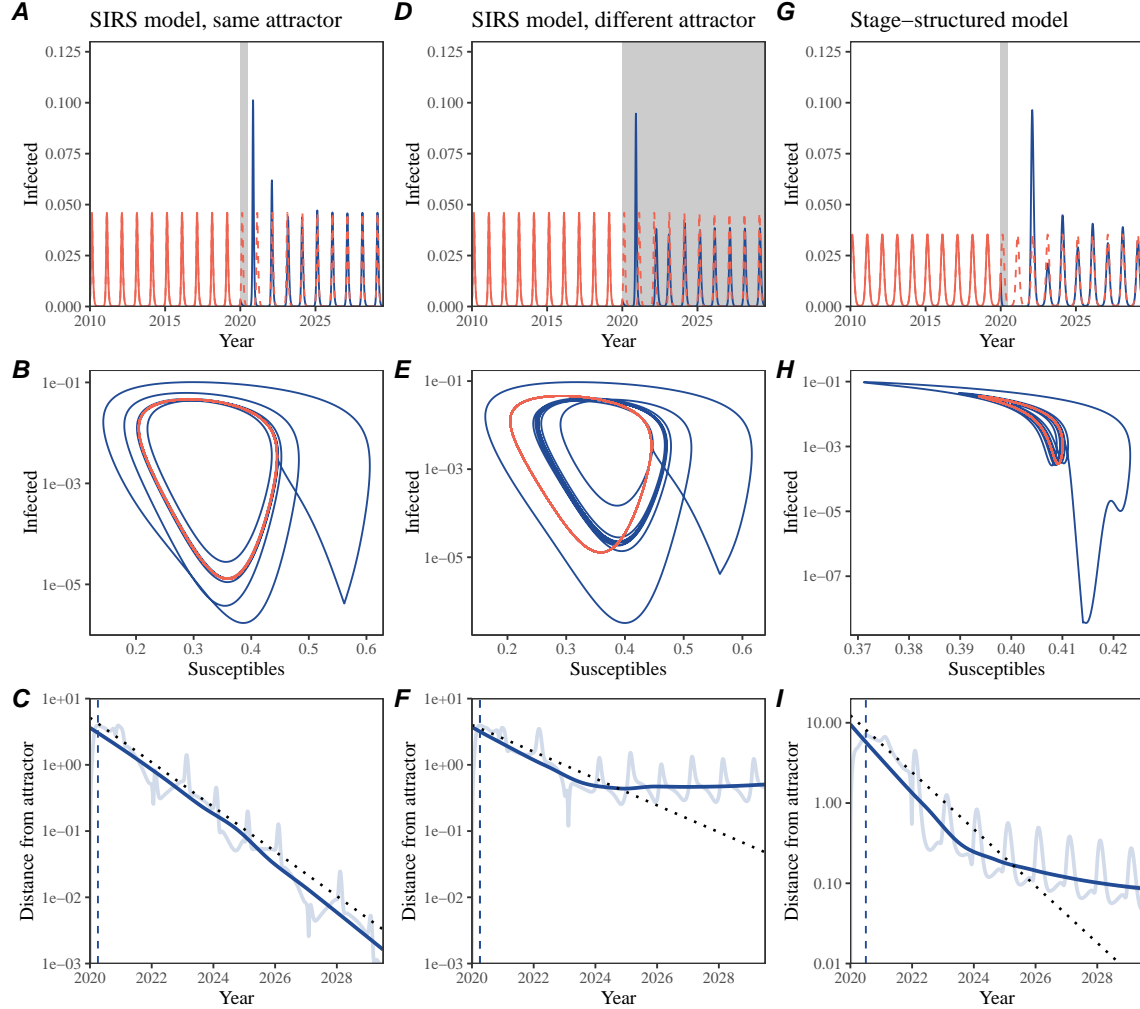


Figure 2: **Conceptual framework for measuring pathogen resilience following NPIs across different scenarios.** (A, D, G) Simulated epidemic trajectories across various models. Red and blue solid lines represent epidemic dynamics before and after interventions are introduced, respectively. Red dashed lines represent counterfactual epidemic dynamics in the absence of interventions. Gray regions indicate the duration of interventions. (B, E, H) Phase plane representation of the corresponding model. Red and blue solid lines represent epidemic trajectories on an SI phase plane before and after interventions are introduced, respectively. (C, F, I) Changes in logged distance from attractor over time. Transparent solid lines represent the logged distance from attractor. Non-transparent solid lines represent the locally estimated scatterplot smoothing (LOESS) fits to the logged distance from attractor. Dotted lines are superimposed as a comparison to have the same slope as the intrinsic resilience of the system.

120 tion of intrinsic resilience, suggesting that empirical approaches can be more reliable

121 ([SWP: TODO]). Second, resilience estimates allow us to make phenomenological  
 122 predictions about the dynamics of a host-pathogen system following a perturbation  
 123 assuming an exponential decrease in the distance from the attractor; this in turn  
 124 can be used to estimate when the system will reach the attractor of interest. Fi-  
 125 nally, deviation from an exponential decrease in the distance from attractor can  
 126 provide information about whether the system has approached an attractor, or a  
 127 ghost attractor, that is different from the original attractor before perturbation was  
 128 introduced.

129 [SWP: *Multi-strain system to be discussed in the supp after some more investi-*  
 130 *gation.*]

## 131 Inferring pathogen resilience from real data

132 Based on these observations, we now set out to infer pathogen resilience from real  
 133 data. To do so, we first reconstruct an empirical attractor by utilizing Takens’  
 134 theorem, which states that an attractor of a nonlinear multidimensional system can  
 135 be mapped onto a delayed embedding. Here, we use delayed copies of logged values  
 136 of pre-pandemic cases,  $C(t)$ , to reconstruct the attractor:

$$\langle \log(C(t) + 1), \log(C(t - \tau) + 1), \dots, \log(C(t - (M - 1)\tau) + 1) \rangle, \quad (1)$$

137 where the delay  $\tau$  and embedding dimension  $M$  is determined based on autocor-  
 138 relations and false nearest neighbors. This allows us to measure nearest neighbor  
 139 distance between the current state of the system and the empirical attractor at any  
 140 given point in time, from which we can quantify how fast this distance decreases by  
 141 fitting a linear regression on a log scale. Then, the resulting estimate of the slope  
 142 of linear regression corresponds to pathogen resilience. Estimates of distance from  
 143 attractor and linear regressions fits are presented in [SWP: TODO: *Supplementary*  
 144 *Materials.*]

145 For most pathogens, resilience estimates are consistent across different countries  
 146 with a few exceptions (Figure 3A). For example, resilience estimates for adenovirus  
 147 for Hong Kong and Korea are orders of magnitudes lower than the corresponding  
 148 estimate for Canada. Another exception is human metapneumovirus, where the  
 149 estimated resilience for Hong Kong is considerably higher than in other countries.  
 150 Otherwise, we find that resilience estimates range between 1–3/years for most other  
 151 pathogens, suggesting a rather fast return to their regular outbreak cycles.

152 Using resilience estimates, we now predict when each pathogen will return to their  
 153 original pre-pandemic cycles. Specifically, we extend our linear regression fits to em-  
 154 pirical estimates of distance from attractor and ask when the predicted regression line  
 155 will cross a threshold value, which we set to a mean of pre-pandemic distances. Sur-  
 156 prisingly, we find that most pathogens should have returned to their original cycles  
 157 by the end of 2024 (Figure 3B). While these predictions are consistent with several  
 158 pathogens (e.g., seasonal coronavirus epidemics in Canada and Korea in Figure 1),

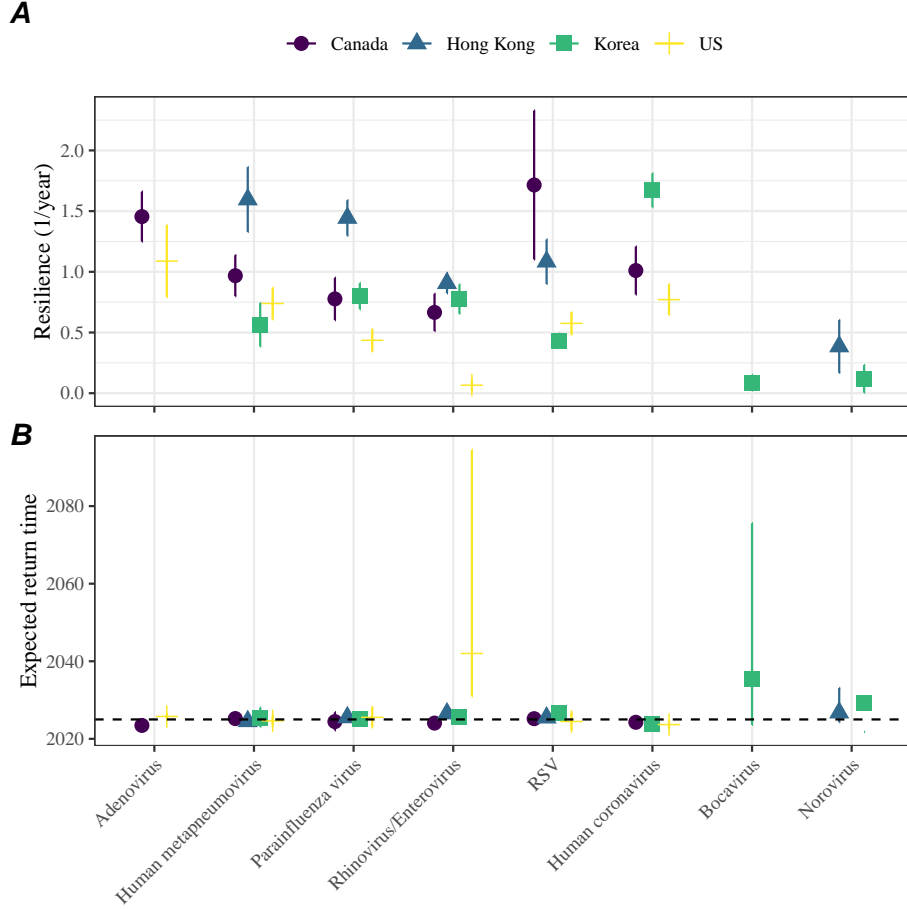


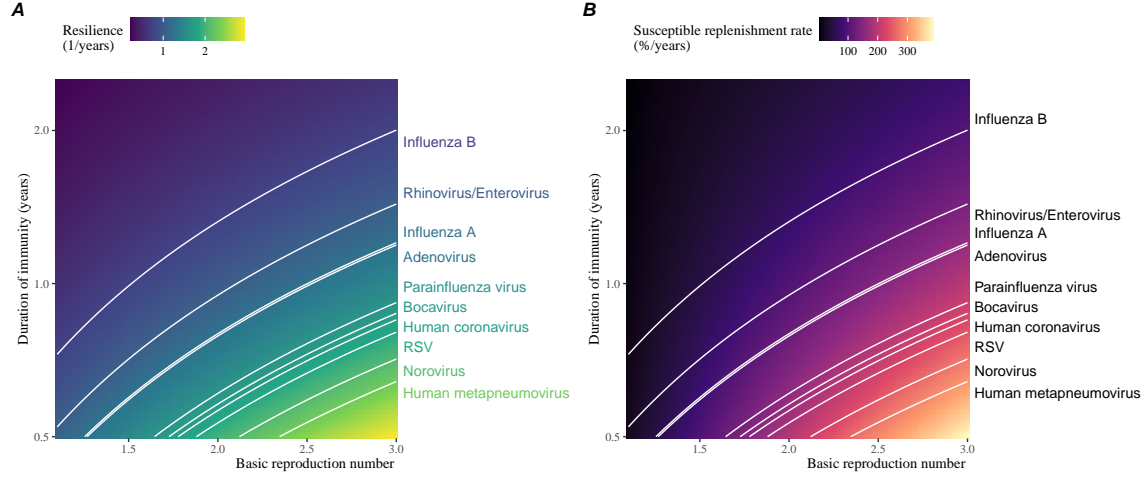
Figure 3: **Summary of resilience estimates.** (A) Estimated pathogen resilience. (B) Predicted timing of when each pathogen will return to their pre-pandemic cycles. The dashed line in panel B indicates the end of 2024 (current observation time). Error bars represent 95% confidence intervals.

there are a few inconsistencies. For example, we predict that bocavirus epidemics in Korea should have returned by the end of 2023 (Figure 3B), but the observed patterns indicate clear deviation from pre-pandemic cycles (Figure 1). These deviations, alongside plateaus in the estimated distance from attractor, suggest that some of these pathogens, such as bocavirus, may have converged to different attractors.

## Susceptible host dynamics explain variation in pathogen resilience

So far, we focused on quantifying pathogen resilience from the observed patterns of pathogen re-emergence following COVID-19 interventions. But what factors deter-

mine how resilient a host-pathogen system is? Here, we use a standard Susceptible-Infected-Recovered-Susceptible (SIRS) model to show that susceptible host dynamics explain variation in pathogen resilience. To do so, we vary the basic reproduction number  $\mathcal{R}_0$ , which represents the average number of secondary infections caused by a newly infected individual in a fully susceptible population, and the duration of immunity and compute intrinsic resilience for each parameter.



**Figure 4: Linking pathogen resilience to epidemiological parameters and susceptible host dynamics.** (A) The heat map represents intrinsic resilience as a function of the basic reproduction number  $\mathcal{R}_0$  and the duration of immunity. (B) The heat map represents per-capita susceptible replenishment rate as a function of the basic reproduction number  $\mathcal{R}_0$  and the duration of immunity. The standard SIRS model is used to compute intrinsic resilience and per-capita susceptible replenishment rate. Lines correspond to a set of parameters that are consistent with mean resilience estimates for each pathogen. Pathogens are ranked based on their mean resilience estimates, averaged across different countries.

We find that pathogen resilience increases with higher  $\mathcal{R}_0$  and shorter duration of immunity (Figure 4A). These variations can be understood in terms of the susceptible host dynamics, where faster per-capita susceptible replenishment rate causes the system to be more resilient (Figure 4B). This rate can be expressed as a ratio between absolute rate at which new susceptibles enter the population and the equilibrium number of susceptible individuals in the population,  $\bar{S}$ . Therefore, both higher  $\mathcal{R}_0$  and shorter duration of immunity can drive faster per-capita susceptible replenishment rate (Figure 4B), especially because higher  $\mathcal{R}_0$  leads to lower  $\bar{S}$ .

Finally, we can now rank different pathogens based on their empirical resilience. For simplicity, we take the average of resilience estimates across countries to determine the ranking, which shows that influenza B is least resilient (Figure 4A). This observation is consistent with the extinction of influenza B/Yamagata, which



indicates the persistence of a pathogen should be associated with its resilience. Similarities in resilience ranking between RSV and Human metapneumovirus is also interesting, given the apparent coupling of their dynamics through cross immunity. These rankings further allow us to map each pathogen onto a set of parameters that are consistent with its empirical resilience (Figure 4A) and obtain a plausible range of susceptible replenishment rates for each pathogen (Figure 4B).

## Discussion

The COVID-19 interventions have caused major disruptions to circulation patterns of both respiratory and non-respiratory pathogens, adding challenges to predicting their future dynamics. On the other hand, these interventions offer large-scale natural experiments for understanding how different pathogens respond to perturbations. In this study, we show that pathogen re-emergence patterns following COVID-19 interventions can be characterized through the lens of ecological resilience. Traditionally, ecological resilience measures how fast a system returns to a reference state following a perturbation. In the context of respiratory pathogens, resilience measures how fast epidemics return to their endemic cycles after interventions are lifted.

We use an attractor reconstruction approach to quantify how distance from attractor changes over time for each pathogen. By fitting a linear regression to log distances, we can estimate pathogen resilience and further predict when each pathogen will return to their endemic cycles. Consistency in resilience estimates across countries is particularly surprising given that each country imposed different intervention measures; this consistency provides robustness to our estimates. The ability to predict future epidemic patterns from resilience estimates also offers a new paradigm for epidemic forecasting. While this approach cannot predict the exact timing of outbreaks or epidemic patterns, it is nonetheless useful for predicting when epidemics will settle down to regular cycles after a large perturbation, such as COVID-19 interventions.

Our analyses suggest a possibility that several pathogens may have converged to different endemic cycles compared to their pre-pandemic epidemic patterns. Key examples include human metapneumovirus, RSV, and bocavirus in Korea as well as RSV in Hong Kong. These changes may reflect permanent changes in behavior since 2020 or a shift in population-level immunity. However, it seems unlikely that permanent changes in behavior would only affect a few pathogens and not others. A shift in population-level immunity is plausible, as the emergence of SARS-CoV-2 and extinction of influenza B/Yamagata likely caused major changes in immune landscapes; however, we currently do not know how immunity, or lack thereof, from these pathogens would affect infection from other pathogens. Future studies should use detailed mechanistic models, coupled with behavioral and immunological data, to test these hypotheses and better understand post-pandemic dynamics of endemic pathogens.

226 We show that susceptible host dynamics shape pathogen resilience, where faster  
 227 replenishment of the susceptible population causes the pathogen to be more resilient.  
 228 For simplicity, we focus on waning immunity and birth as a main driver of the suscep-  
 229 tible host dynamics but other mechanisms can also contribute to the replenishment  
 230 of the susceptible population. In particular, pathogen evolution, especially the emer-  
 231 gence of antigenically novel strains, can cause effective waning of immunity in the  
 232 population; therefore, we tentatively hypothesize that faster rates of antigenic evo-  
 233 lution can also cause a pathogen to be more resilient. Future studies should explore  
 234 the relationship between the rate of evolution and resilience for antigenically evolving  
 235 pathogens.

236 Quantifying pathogen resilience also offers novel approaches to validating epi-  
 237 demiological models. So far, the majority of model validation in epidemiology is  
 238 based on the ability of a model to reproduce the observed epidemic dynamics and to  
 239 predict future dynamics. However, there can be plethora of models that meet these  
 240 criteria. For example, two major RSV models have been proposed so far to explain  
 241 biennial epidemic patterns: (1) a stage- and age-structured model that allows for  
 242 disease severity to vary with number of past infections and age of infection and (2)  
 243 a pathogen-interaction model that accounts for cross immunity between RSV and  
 244 human metapneumovirus. Since both models can accurately reproduce the observed  
 245 epidemic patterns, standard criteria for model validation do not allow us to distin-  
 246 guish between these two models. Instead, we can measure the empirical resilience  
 247 of each model by simulating various perturbations and compare them to estimates  
 248 of empirical resilience from data, using COVID-19 interventions as an opportunity.  
 249 Future studies should further investigate using pathogen resilience for validating epi-  
 250 demic models.

251 There are several limitations to our work. In particular, our estimates of pathogen  
 252 resilience and the associated ranking are necessarily crude. [SWP: *Limitation TBD.*]  
 253 Nonetheless, our study illustrates the utility of quantifying pathogen resilience for  
 254 understanding how different pathogens respond to perturbations.

255 [SWP: *Conclusion paragraph TBD.*]

## 256 Materials and Methods

### 257 Data

258 We gathered time series on respiratory infections from four different countries: Canada,  
 259 Hong Kong, Korea, and United States (US). As a reference, we also included time  
 260 series data on norovirus infections for available countries—in contrast to respiratory  
 261 pathogens, we expect gastrointestinal viruses, such as norovirus, to be less affected  
 262 by COVID-19 intervention measures.

263 Weekly time series of respiratory infection cases in Canada come from the Res-  
 264 piratory Virus Detection Surveillance System, which collect data from select labo-  
 265 ratories across Canada. We extracted the data from <https://www.canada.ca/en/>

266 `public-health/services/surveillance/respiratory-virus-detections-canada.`  
267 `html.`

268     Weekly time series of respiratory infection cases in Hong Kong came from the  
269     Centre for Health Protection, Department of Health. We extracted the data from  
270     <https://www.chp.gov.hk/en/statistics/data/10/641/642/2274.html>.

271     Weekly time series of respiratory infection cases in Korea came from Korea Dis-  
272     ease Control and Prevention Agency. We extracted the data from [https://dportal.](https://dportal.kdca.go.kr/pot/is/st/ari.do)  
273     [kdca.go.kr/pot/is/st/ari.do](https://dportal.kdca.go.kr/pot/is/st/ari.do).

274     Finally, weekly time series of respiratory infection cases in the US comes from  
275     the National Respiratory and Enteric Virus Surveillance System.

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