REPLY TO KUŚMIERZ AND TOYOIZUMI:

A network-based explanation of why most COVID-19 infection curves are linear

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In a comment on our recent paper (1), the commentators point out that infection curves on small-world (SW) networks are linear only in the vicinity of the critical point (2) and that infection dynamics on complex or SW networks have already been studied, for instance, in ref. 3. In our work (1), we extend the existing literature by showing, in detail, how an explanation of the apparent sustained algebraic growth regimes observed in many countries during the COVID-19 pandemic involves the structure of the underlying contact network of individuals and how a transition from exponential spreading to a power-law behavior happens when the reproduction number takes values in the vicinity of $R \approx 1$.

We are surprised to see that the commentators in ref. 2 try to suggest that, in ref. 1, we demonstrate that the linear behavior should also persist well below the critical reproduction number $R \approx 1$. We do not do that. Throughout the paper and its extensive supporting information, we emphasize that the derivation of the estimates for the critical connectivity $D_c$ (and other arguments) hinges exactly on the assumption that linear behavior can only happen if the reproduction number is close to criticality, that is, $R \approx 1$, or, equivalently, $D \approx D_c$.

The critical values $R \approx 1$ or $D \approx D_c$ are not sharp, and, as we state explicitly in ref. 1, the approximate formula for $D_c$ slightly overestimates the true value. The confusion in ref. 2 might be due to the fact that, at one point (ref. 1, p. 22,687), we state that the critical behavior, that is, linear growth, can be observed when $D$ falls below $D_c$. However, we certainly did not indicate with this sentence that we believe that the critical phase extends to values of $R$ far from $R \approx 1$. Within the context of the entire paper (1) and its supporting information, this is completely clear. It seems that the commentators in ref. 2 took that sentence out of context to create a case that does not exist. For this reason, we strongly object to their statement of having performed a flawed analysis.

In addition, the analysis of close-to-critical spreading on Erdős-Rényi (ER) networks in ref. 2 reveals an important difference between ER and SW networks. A central result in our work (1) is that the critical threshold, above which case numbers grow exponentially in the SW model, is above the levels that are expected from a comparable model with homogeneous mixing of individuals (figure 3A in ref. 1). As seen in figure 2A in ref. 2, the $D_c$ values for ER networks for different settings are clearly below the corresponding $D_c$ for SW networks. This difference arises because of network effects that—due to overlapping neighborhoods of nodes—reduce the spread below those levels expected from homogeneous mixing, an effect that also persists, albeit attenuated, in ER networks.

It is fortunate, however, that this comment gives us the opportunity to cite a work that noted the possibility of linear spreading of diseases on SW networks before (figure 5 in ref. 4) and that we were not aware of at the time of composing ref. 1.

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2 Ł. Kuśmierz, T. Toyoizumi, Infection curves on small-world networks are linear only in the vicinity of the critical point. Proc. Natl. Acad. Sci. U.S.A., 10.1073/pnas.2024297118 (2021).
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https://doi.org/10.1073/pnas.2100906118 | 1 of 1