One Health, “Disease X” & the challenge of “Unknown” Unknowns

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The emergence of SARS-CoV-2 and its rapid spread globally emphasizes the ever-present threat of emerging and re-emerging infectious diseases. In this review, the pathogen pyramid framework was utilized to identify the “unknown unknowns” associated with the emergence and rapid transmission of novel infectious disease agents. Given that the evolutionary origin of most of the emerging infectious disease agents can be traced to an animal source, we argue the need to integrate the “One Health” approach as a part of surveillance activities. The need for focusing on undertaking global and regional mapping activities to identify novel pathogens is discussed, given that there are an estimated 1.67 million unknown viruses, of which around 631,000 to 827,000 unknown viruses have the capacity to infect human beings. The emerging risks due to the ever-expanding interface between human, animals, both domestic and wildlife, and the environment are highlighted, these are largely driven by the need for safe habitation, growing food, developing infrastructure to support the increasing human population and desire for economic growth. The One Health approach provides a holistic way to address these cross-sectoral issues, by bridging institutional gaps, enumerating priority risk areas and pathogens, and highlighting putative risk factors for subsequent spillover events involving emerging and re-emerging infectious disease pathogens at the human-animal-environment interface.

Key words COVID-19 - emerging infectious diseases - One Health - pandemics - pathogen pyramid - surveillance

The default response to infectious disease threats has been the development of preventive or curative options post facto, often in the face of rising disease burden, human suffering, and costs. Though the value of timely identification of emerging infectious diseases has been well-established in the context of transmission containment, even with the most sensitive of surveillance systems, it is difficult to identify a novel pathogen¹. The evolutionary advantage held by microbes enables them to adapt rapidly to host species, and eventually spill over. In the 25 families of viruses that can potentially infect human beings, there are an estimated 1.67 million unknown viruses, of which, an estimated 631,000 to 827,000 unknown viruses have
the capacity to infect human beings\textsuperscript{2,3}. Given this wide array of potential spillover threats, and the absence of a risk stratified global virus atlas, it becomes important to improve disease surveillance, especially at the interfaces with the highest risk of novel pathogen emergence.

The most recent cross-species spillover of a novel coronavirus, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), has assumed pandemic proportions, resulting in almost 2.5 million deaths from the novel coronavirus disease (COVID-19) globally by the end of February 2021\textsuperscript{4}. This is the third instance of the emergence of a novel coronavirus, after the severe acute respiratory syndrome (SARS) in 2003 and Middle East respiratory coronavirus (MERS CoV) in 2012\textsuperscript{5-7}. The consistency with which these novel pathogens have transcended species and international borders, while a cause of grave concern, is also an indicator of a complex interplay of various factors at the human-animal-environment interface, through which the pathogens continue to expand their host-tropism. While some efforts at using computational approaches have shown encouraging signs, accurate prediction models, which can predict the potential spillover threat, as well as its impact on human health, are yet to be developed\textsuperscript{8-10}. Although SARS-CoV-1, MERS-CoV and SARS-CoV-2 all share similar phylogenetic roots, yet the considerable heterogeneity between the pathogen transmission patterns, disease severity and case fatality rates gives rise to varying magnitudes of global health threats. What makes SARS-CoV-2 a truly unique threat is its high transmissibility, and ability to cause adverse health outcomes (severe disease requiring hospitalization, and deaths), especially in vulnerable population groups, such as the elderly or those with co-morbidities, while simultaneously keeping overall case fatality rates low enough to enable further spread\textsuperscript{11-17}.

**Using the pathogen pyramid to explore emerging infectious diseases**

These novel microbes are new entrants in a long emerging pattern. Between 1940 and 2004, 335 pathogens have emerged, with 60 per cent having a zoonotic source, of which 71 per cent originate from wildlife\textsuperscript{18}. Prediction models have identified emerging infectious disease (EID) hotspots in Africa, Latin America and South Asia\textsuperscript{19}. Most of these models take into account the current understanding of disease dynamics, and assumptions with respect to some of the “known unknown” determinants. The “unknown unknowns”, which form a large part of the problem at hand, remain in the blind spot. As the rapidly expanding pandemic of COVID-19 has adequately proved, this is an oversight one cannot afford.

One of the approaches used to identify the origins of zoonotic diseases utilizes the ecologic “pathogen pyramid”. This approach identifies the pathogens’ transitions from being an environment- or animal-specific agent into a human-infecting one\textsuperscript{20,21}. This approach identifies intermediate levels of adaptation, through which zoonotic microbes evolve into efficient human pathogens over time (summarized in the Figure).

At level 1 exposure, humans are exposed to a wide range of pathogens of environmental or animal origin, which may leave an immunologic trace, that are either unlikely, or not known to be disease causing pathogens. Naturally acquired infection with non-human simian retroviruses is a typical example\textsuperscript{22}. Given the number of unknown viruses which exist at the human-animal-environment interface without crossing the species barrier effectively, or without adapting to a pathogenic role in human hosts, it is difficult to estimate the number and types of viruses which may be included in this level\textsuperscript{22}.

Level 2 reflects the stage of infection, where zoonotic viruses can overcome the species barrier, invade human cells, and cause disease. However, the risk of human-to-human transmission, while theoretically possible, remains very low, as is seen in the case of diseases like Japanese encephalitis or rabies\textsuperscript{23-26}. Another example is the influenza A(H5N8) virus, which is not known to exhibit human-to-human transmission, and does not show efficient transmission between ferrets, the animal model of choice for human influenza infections in human beings\textsuperscript{27-31}.

At level 3, in addition to causing human infections, the zoonotic viral pathogens can cross the human-to-human transmission barrier and can establish limited transmission chains, resulting in outbreaks, which flare up and then eventually die down. This is the case for diseases like the plague, Nipah, Ebola or Marburg virus diseases, which are rapidly fatal.

At level 4, the pathogen can establish efficient human-to-human transmission chains, which can result in disease burden of epidemic or pandemic proportions, as has been observed for the ongoing COVID-19 pandemic, its predecessors MERS-CoV,
SARS, or influenza. HIV can also be considered to be a pathogen at this level of the pathogen pyramid. Levels 3 and 4 provide opportunities for interventions to disrupt transmission chains of emerging infectious diseases. This structure is referred to as a pyramid owing to the tapering number of pathogens from level 1 to level 4, with fewer pathogen members at each incremental level. **Emerging infectious diseases, zoonoses and “Disease X”**

Tracing this ecological chain of uncertainties almost always leads to an animal source. The diseases identified by the WHO’s priority research and development (R&D) blueprint in all zoonotic in nature and have been responsible for outbreaks in the recent past. COVID-19 is the latest entrant in the list, having graduated from the category of Disease “X”. This ominously named category represents the currently unknown pathogens, which may emerge from unidentified spillover events, and result in outbreaks that may pose international threats. The risk of the emergence of these yet to be identified threats grow larger with increasing human population, loss of biodiversity, changing climates, aggressive land use for human habitation and agriculture, all of which contribute to the expanding interfaces between humans, animals and the environment. The epidemiology of the H1N1 influenza virus outbreak in 2009, which contained genetic material from human, avian and swine origin, involved wildlife, pig farming, animal movement and farm workers.

The specific animal-to-human spillover event for SARS-CoV-2 infection in man remains poorly understood. Hypothesized to have originated from horseshoe bats, the role and identity of any intermediate hosts in the transmission chain remains unclear. Genetic studies propose two pathways of transmission: viral mutations in human after zoonotic transmission; or mutations in animal reservoirs before zoonotic transmission to human. While the current evidence base is not substantive enough to distinguish which of these two trajectories was taken by SARS-CoV-2, the latter represents a more efficient transmission pathway, which is reflective of the threat posed by the “unknown unknowns”.

As human residence, agriculture, and consumption patterns continue to invade wildlife habitat, the pathophysiological aspects of these changing equations are often overshadowed by the economic imperative or cost-benefit rationalizations. Human encroachments into animal well-being and territory are usually considered under the framework of biodiversity and conservation, and the disease risk perspective often remains unexplored and undervalued. The role and risk of densification of animals through factory farming, animal markets, and wildlife trade should be evaluated, and disease surveillance conducted in these high-risk contexts. Densification further enhances the opportunities for viral replication and mutation.

**One Health approach and emerging infectious disease threats**

Despite the emergence of some structured efforts at undertaking cross-sectoral approaches to build...
could increase by 10, 20 and 36 per cent, respectively over a five-year period\textsuperscript{47}. Other estimates computed early in the COVID-19 pandemic revealed that even modest levels of reduction in coverage of maternal and child health services in LMICs were likely to result in over 250,000 additional child deaths and over 12,000 maternal deaths\textsuperscript{48}. The Global Financing Facility estimated that childhood vaccinations were likely to drop by 50 per cent or more in many LMICs facing COVID-19 related lockdowns or service restrictions\textsuperscript{49}. Another modelling effort showed that in a high-impact setting in Africa, for every COVID-19 death that could be attributed to SARS-CoV-2 infection through person-to-person transmission during routine childhood vaccinations, 84 under-five deaths due to vaccine preventable diseases, such as measles, could be avoided\textsuperscript{50}.

The evolution of the COVID-19 threat has given us a unique opportunity to introspect on the narrative of reactive response driven epidemic management \textit{vis-à-vis} the need to develop proactive prediction systems which can pre-empt novel spillovers and emergence or re-emergence of dangerous pathogens\textsuperscript{51}. Further, the trajectory at which the pandemic has progressed in different countries has been slightly different, which also drives the issue of identifying and contextualizing evidence to base policy on. As identified by WHO, a one-size-fits-all approach to manage COVID-19 is likely to fail, given the wide variance in the epidemiology of COVID-19 globally\textsuperscript{52}.

**Global learnings from the pandemic response**

There have been several positive takeaways from the global experiences of dealing with the COVID-19 threat. Global ability to devise technical solutions at a short notice has improved. The whole genome sequence of the virus was examined and rapidly shared on public domains\textsuperscript{43-55}. Publications have come out in pre-print servers, cutting down the time to put evidence to use. This is one of the first instances where the evolving epidemiology in various countries is being studied in real time by a diverse group of researchers from across the world. Access to libraries of pre-tested compounds has accelerated drug development efforts, while computational approaches have helped identify \textit{in silico}, which drugs would be a best fit to repurpose. Innovations in risk-mitigating technologies have allowed businesses, education and healthcare, to transition to the new normal\textsuperscript{56}. The development of COVID-19 vaccines has also taken
place at an unprecedented pace, with global and regional collaborations being developed to produce and disseminate the vaccines globally\(^\text{57,58}\). However, with the inevitable imbalance between demand and supply, we have also been witness to the emergence of the phenomenon of vaccine nationalism, especially in high-income countries, which has threatened the notions of global solidarity\(^\text{59-61}\).

Given the global implications of the spread of COVID-19, these technical solutions may fall short. Diplomatic and international cooperative solutions such as data sharing, using common protocols for rapid product (diagnostic, preventive, therapeutic) testing, and accelerated or common approval systems for deployment of medical products for diagnosis, prevention or management of COVID-19 or other novel spillover events need to be explored and encouraged. Cross-border and regional cooperation remain central to the discussion around developing solutions that work for all. Further, transparent systems, undergoing iterative improvement over time, based on accurate data, collected, and reported through sensitive surveillance systems, would also provide the means to ensure global solidarity in response to emergent infectious disease threats. In our global village, and infectious scourge at any corner of the world is a potential threat to health security for all\(^\text{62}\).

**Using One Health to move from reactive to proactive response**

The holistic view of health encouraged by a One Health perspective supports the inter-sectoral collaboration on human, animal, and environmental health. Novel and emerging infectious diseases is a perfect platform for incorporating this perspective\(^\text{63}\). It is also important to consider the potential for climate change to influence future pandemics\(^\text{64}\). With a change in climate, animal (including vectors) and plant species will shift as the suitability of different areas for specific biomes changes. This will likely change the interaction dynamics at the evolving human-animal-environment interfaces, as different species of animals will come into contact, under a changing ecosystem. Erratic weather patterns, droughts, heat waves and desynchronizations of life cycles of animals and plants may also alter the mix of animals in specific locations. New mixing patterns of different plant and animal species, and pathogens specific to each, has the potential for new cross-over infections between animals, with subsequent human spillovers\(^\text{65}\). Moreover, changes in the distribution of animals, environmental degradation itself, crop failure or livestock death, for example, may alter human interactions with wildlife, bringing them into closer contact or increasing hunting and eating of bushmeat, for instance\(^\text{66}\).

In February 2018, in its blueprint for research on priority diseases, the WHO prophetically added a new potential threat: Disease X\(^\text{35}\). The X stood for something unexpected, a microbe that had the potential to cause a future pandemic. This was based on advocacy from multiple health experts, who pointed out that it was a question of when and not if the next disease with epidemic potential would emerge from a hitherto unknown source, the most likely being a zoonotic one\(^\text{67}\). However, the common consensus was that it would be a pandemic influenza and some nations developed plans to deal with such outbreaks. A regional outbreak that progressed to a public health emergency of international concern and finally to a pandemic in a couple of months, killing millions in less than a year took the world by surprise\(^\text{4}\). Hence, given the continued threat of emerging infectious diseases, the global health community must review the successes, lessons learnt, and mistakes made over the past year to ensure that history does not repeat itself.

COVID-19 is a wakeup call which provides us an opportunity to learn and adapt, so that the impact of future pandemics can be mitigated\(^\text{68}\). It is obvious, that we need to focus on developing more precise early warning systems, which pre-empt outbreaks or spread of epidemics, and enable us to mount an effective response before sustaining losses in human and animal life, and economic well-being\(^\text{69}\). In addition, we must look at investing in strengthening health systems and interdepartmental collaborations; improving the sensitivity, breadth and fidelity of disease surveillance; and empowering communities to react and respond to outbreak threats in a more efficient manner\(^\text{70,71}\). Whilst the immediate focus should be directed towards containing the COVID-19 threat and minimizing the losses, both in terms of human lives and economic security, we must build up the bulwarks before the next novel dangerous pathogen comes calling at our doors. Just as Joshua Lederberg, the Nobel-laureate had stated, “the future of humanity and microbes likely will unfold as episodes of a suspense thriller that could be titled ‘Our Wits Versus Their Genes’”\(^\text{72}\).

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