COVID-19: a pandemic experience that illuminates potential reforms to health research

Hannah N Kozlowski1,2, Michael E Farkouh3, Meredith S Irwin4, Laszlo G Radvanyi5, Aaron D Schimmer6, Uri Tabori4 & Norman D Rosenblum4,*

EMBO Mol Med (2020) 12: e13278

COVID-19 has halted research around the globe and forced researchers out of their laboratories. Non-emergency medical appointments were canceled. Ongoing clinical trials were challenged to create new modes of operation while public pressure mounted to find therapeutic options against COVID-19. Yet, the inability to conduct research during COVID-19 was overcome with cooperation, resource sharing, and compassion, which provides important lessons on how to improve health related research as we enter a new normal.

To meet the COVID-19 challenge, the global scientific community focused human and material resources to develop new drugs, therapies, vaccines, diagnostics, and so on. However, it takes 10–20 years to turn a basic discovery into a new drug via the typical translational path (Paul et al, 2010; Mohs & Greig, 2017), which would not be compatible with the goal of rapidly developing diagnostics and therapeutics for COVID-19. This situation created a need to overcome the many hurdles along the translational research pathway—financing, business development, regulation, product development, commercialization, and different cultures—in order to satisfy an urgent societal need. Here, we discuss the lessons learned from the Toronto Academic Health Sciences Network’s (TAHSN) response to COVID-19. We highlight the critical importance of cooperation and compassion in maintaining productivity during the pandemic and identify opportunities for shaping a new era of research with increased patient engagement, fewer silos, and a shared goal of improving health and decreasing disease burden.

Themes emerging from the COVID-19 pandemic

The development of ready-for-use clinical products during the COVID-19 pandemic has engendered cooperativity, engagement, and investment in a common goal (Fig 1), enabled by a shared view of the problem and a genuine desire to contribute to its solution. The fruits of these efforts can be seen around the world as pharmaceutical and biotech companies, and research groups team up to develop, test, and manufacture vaccines (Liu, 2020). This created a real “open science” environment, in which equipment, manufacturing capabilities, patient cohorts, and results were shared in real time (Rouleau, 2020). Prior barriers that obstructed cooperation and sharing were dismantled, facilitating the development of novel ideas into solutions that benefit patients.

Sharing expertise and resources increased clinical capacity and catalyzed scientific discovery

To expand COVID-19 testing capacity and reduce redundancy, four medical research institutions in Toronto agreed to a resource-sharing approach. This is not necessarily new: many philanthropic organizations including the Gates Foundation and the Wellcome Trust have already made open data sharing mandatory for their grants (Kiley et al, 2017). In Toronto, this new sharing agreement went further to include samples and reagents to allow free movement between institutes without any attendant intellectual property (IP) agreements.

Two major lessons emerged. First, focusing on the shared goal, rather than first negotiating IP rights, increased productivity. Early conversations around IP are expensive, slow negotiations between institutes and have stymied innovation. Second, sharing resources such as reagents, equipment, staff, and expertise increased the number of problems that could be solved. Without such agreements, hospitals would only test their own patients and the majority of these basic tests indicated only viral load. Instead, cooperation leveraged equipment, protocols, and expertise external to hospitals. The Ontario Institute of Cancer Research, normally a

1 Biomedical Engineering, University of Toronto, Toronto, ON, Canada
2 MD/PhD Program, Temerty Faculty of Medicine, University of Toronto, Toronto, ON, Canada
3 Peter Munk Cardiac Centre, Heart and Stroke Richard Lewar Centre, University of Toronto, Toronto, ON, Canada
4 Department of Paediatrics, The Hospital for Sick Children, University of Toronto, Toronto, ON, Canada
5 Ontario Institute for Cancer Research, Toronto, ON, Canada
6 Princess Margaret Cancer Centre, University of Toronto, Toronto, ON, Canada

*Corresponding author: E-mail: norman.rosenblum@sickkids.ca
DOI 10.15252/emmm.202013278 | EMBO Mol Med (2020) 12: e13278 | Published online 19 October 2020
cancer-focused genomics institute, sequenced indeterminate and positive COVID-19 samples. They shared the sequences in real time to further the scientific community’s understanding of COVID-19, which exemplifies how resource sharing between institutions can increase testing capacity and support scientific discovery.

New partnerships were built on a foundation of resource-sharing initiatives and the motivation to contribute during the crisis. In Toronto, new collaborations between academia and industry resulted in multicenter clinical trials that transitioned from idea to approval in as few as 14 days. The scale and speed of these agreements were only possible because academic researchers and private industry worked together with a shared understanding of the problem and jointly identified their collective goals, expertise, available resources, and clinical networks. In one instance, a conversation between two colleagues on the use of immunomodulators to protect patients with cancer from the severe side effects of COVID-19 infection became a phase three, multicenter clinical trial. The total time from initial conception to protocol drafting to approval by Health Canada was only 6 weeks. We have also seen international pharmaceutical companies share their manufacturing facilities, supply chains, and expertise in scaling (Liu, 2020; The Association of the British Pharmaceutical Industry, 2020). These joint commitments led to further cooperation between researchers, private industry, research ethics boards, and national regulators.

**Cooperation reduced inefficiencies associated with clinical translation**

During COVID-19, regulators worked closely with scientists to quickly translate scientific discoveries into medical interventions. Health Canada released guidance documents for approval of COVID-19 medical devices, including diagnostic tests, to simplify and streamline the application process, without impacting post-market safety standards (Reid, 2020). Similarly, the US FDA issued an emergency use authorization that expedited the review of COVID-19 diagnostics (Reid, 2020). This sort of response by regulators around the world led to increased cooperation with researchers, which boosted productivity.

**Compassion and support for the people behind the research fostered sustained productivity**

The research community also realized that supporting research meant supporting each other. Behind each study are researchers with unique values, expectations and experience. Attention to individuals and their circumstances—delays in student graduation, personal comfort in the workplace, safe commuting, family responsibilities, challenges associated with working from home, and uncertainty due to job security—became central to discussions on restarting research after COVID-19 restrictions are loosened. For example, flexible working times and shift work allowed for physical distancing, avoiding rush hours, and benefited families with working parents. Beyond any particular research program and institution, TAHSN provided virtual resources, seminars, social interactive space, and training to promote support research staff and foster a positive working environment, all of which strengthened the research community and sustained productivity.

During the pandemic, several non-COVID-19 trials were also forced to adapt. Running these trials without compromising rigor or the safety of patients required cooperation between researchers, local research ethics boards (REBs), patients, and patient advocates to address patient anxieties and clinical priorities by developing new protocols. In Toronto and medical clinics worldwide, clinicians adapted by moving medical visits and clinical trial appointments online. Using video calling, physicians saw their patients in their home environment as well as other family members who joined the call. Although these interactions provided a broader view of the patient, their health, and their environment, careful retrospective analyses of the safety, reliability, and effectiveness of virtual patient assessments are still needed to determine if this approach is sustainable.

**A framework for the “new normal”**

Harnessing lessons learned during COVID-19 to improve health research requires joint action to further improve research...
To foster cooperation, we need to assemble team members of the broader research and translation community around a shared goal. During COVID-19, researchers, industry professionals, and regulators were each determined to find solutions that benefit patients right away. Maintaining this shared focus going forward requires that we rebuild activities to support cooperation. In Toronto, we started by restructuring hospital grand rounds, the institute-wide clinical seminars, to educate the community on a relevant medical topic. Clinicians and researchers from different fields now give joint talks to highlight the larger problem and possible solutions from their perspective. This was particularly helpful during COVID-19 as infectious-disease clinicians discussed the newfound pediatric presentation of COVID-19 and immunologists explained the immune response and how that informs vaccine development. Moving forward, collaborative research should become a model for trainees and other researchers to strive for as they see the fruitful outputs of teamwork.

To further support cooperation and build partnerships, we need to invest in research clusters that are grouped by disease or clinical question instead of methods. Each cluster would include basic science, health services, implementation, engineering, and clinical expertise. Seminars within each cluster would describe different facets of the problem, as per the expertise of the speaker. This provides researchers and trainees a shared view of the big picture so that they can readily come together to solve the problem. These collaborative clusters should also include patient advocates and industry to open communication and help different groups to understand each other and facilitate a shared culture. In such a disease-focused research cluster, the role of each member will be clearly understood, reducing the number of barriers to producing research that benefits patients.

Table 1. Summary of changes implemented during COVID-19 that benefited the advancement of medical discoveries. Further benefits can be seen if long-term recommendations are implemented.

| Themes                      | Short-term changes to adjust for the impact of COVID-19                                                                 | Long-term changes inspired by the changing COVID-19 environment                                                                 |
|-----------------------------|-------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------|
| Sharing resources and expertise | Created new data-sharing agreements without extended IP negotiations                                                 | Keep focused on producing research that benefits patients                                                                      |
| Cooperation                 | Regulators released clear guidance documents to streamline regulatory approval applications                          | Create research clusters that promote sharing and cooperation between basic science, health service, implementation, engineering, and clinical researchers; patient advocates and private industry |
|                             | REB members learned from non-REB scientists to efficiently and thoroughly review ethics applications                   | Recognize researchers for collaborative or middle author roles in order to promote team science                                  |
|                             | REBs, researchers and patients worked together to revise clinical trial protocols                                        | Reward non-publication achievements (i.e., patient enrollment, clinical guidelines)                                               |
|                             | Clinicians and researchers conducted medical appointments over the phone or online                                       | Create government-industry grants that support the long-term development of scientific discoveries into clinical impact        |
| Humanizing research         | Acknowledged and discussed researchers’, staff, and students’ unique values and circumstances                            | Continue thinking about health and personal circumstances of all people involved in research                                     |
|                             | Increased resources for mental health and community building                                                           | Continue virtual and in-person resource sharing to promote good mental health and foster community                             |
|                             | Enhanced researcher consultation with patients to understand their disease, their experience, and what matters to them | Continue to engage patients from inception and throughout a project                                                            |
|                             |                                                                                                                         | Develop pathways for students to learn from patients and experience medical clinics                                             |

To encourage resource sharing and to move research discoveries closer to patients, research institutions need to value and reward teamwork. Yet, the current academic structure does not always reward team science or the building of collaborative networks. In the medium-term, we should change how we evaluate scientific contributions and create promotional metrics that recognize team science. In Toronto, we acknowledge that success may not always result in a publication but in the adoption of a new policy, better clinical guidelines, a new partnership with industry or the clinical validation and implementation of a diagnostic test. In the long term, we should provide more grants that promote collaboration akin to the European Union’s Research Programmes that provide long-term funding for collaborations between multinational research groups.

Supporting long-term development relieves the pressure to publish quickly and often and encourages teams to tackle big problems. Given the time and resource needed to bring new interventions to the clinic, the funds for these grants should come from government-industry partnerships, and there should be an option of applying for additional funding if pre-specified milestones are met (van Dijk et al, 2019).

Producing research that is meaningful to patients also requires that we humanize research. During COVID-19, there was a shift from thinking about research as something that eventually trickles down to a “patient”, to understanding that we all may benefit. This makes it more natural to think of patients as the reason and focus of our scientific inquiries. In Toronto, we previously started shifting toward patient-
centered research by involving patients and families in advisory committees, ethics boards, and research teams as charitable foundations have been doing for years (Stevens, 2019).

We can further strengthen this partnership by creating pathways for students, the researchers-of-tomorrow, to learn from patients directly. In these pathways, students will spend more time in the clinic, meet patients and caregivers, write REB proposals, analyze the flow of information between health professionals, and observe medical procedures (DelNero & McGregor, 2017). For example, asking patients why they continue/discontinue their treatments and what quality of life indicators/outcomes matter to them can inform study design and give students a sense that they are contributing to improve patient health. This experience also breaks down barriers between research and clinic. For researchers, it provides a broader understanding of a disease and the challenges of implementation. For patients and caregivers, it is an opportunity to share their wisdom and learn about the research process.

Change should happen now

Now is the time to harness the lessons learned from COVID-19 and shape a new era of innovation that more effectively brings the benefits of research to patients. We cannot let the positive changes be lost. First, we need to continue to create joint goals between all members of the translational path. Sharing must continue to extend beyond reagents and funding. These actions can be strengthened by restructuring our institutions to reinforce “open science”, developing broad data-sharing agreements and creating collaborative work spaces. Second, we need to value and reward teamwork. This involves creating new measures of academic success and policy makers in order to generate solutions that are used in the clinic.

Acknowledgements

The experiences and framework discussed in this commentary were based on a town hall given by Medical Innovations Toronto (MiTO), a translational research hub within the TAHSN. In this discussion TAHSN leaders discussed how the pandemic changed the research landscape in Ontario, the lessons they learned from the response to COVID-19 and the implications their experiences will have on a new era of medical research.

Author contributions

H.N.K. and N.D.R. conceived the idea, wrote and edited the manuscript; all authors contributed content and reviewed the final manuscript.

Conflict of interest

ADS has received research funding from Takeda Pharmaceuticals and Medivir AB and consulting fees/honorarium from Novartis, Jazz, and Otsuka Pharmaceuticals. ADS also holds stock in Abbvie, UT, LR, MF, MI, HK, and NDR have no relevant relationships to disclose.

References

DelNero P, McGregor A (2017) From patients to partners. Science 358: 414
van Dijk SJ, Domenighetti AA, Gomez-Ospina N, Hunter P, Lindemans CA, Melotte V, van Rossum AMC, Rosenblum ND (2019) Building a professional identity and an academic career track in translational medicine. Front Med 6: 1–6
Kiley R, Peatfield T, Hansen J, Reddington F (2017) Data sharing from clinical trials – a research funder’s perspective. N Engl J Med 377: 1990–1992
Liu A (2020) Pfizer offers up talents, tools and manufacturing capabilities in call for wide COVID-19 collabs. Available at: https://www.fiercepharma.com/pharma/pfizer-offers-up-tale nts-tools-and-manufacturing-capabilities-call-for-wide-covid-19-collabs
Mohs RC, Greg NH (2017) Drug discovery and development: role of basic biological research. Alzheimers Dement Transl Res Clin Inter 3: 651–657
Paul SM, Mytelka DS, Dunwiddie CT, Persinger CC, Munos BH, Lindborg SR, Schacht AL (2010) How to improve R&D productivity: the pharmaceutical industry’s grand challenge. Nat Rev Drug Discov 9: 203–214
Reid A (2020) How regulators are addressing COVID-19 related Medical Device approvals. Available at: https://starfishmedical.com/blog/covid-19-related-medical-device-approvals/
Rouleau G (2020) The key to finding a cure for COVID-19: Open science. Globe Mail Available at: https://www.theglobeandmail.com/opinion/article-the-key-to-finding-a-cure-for-covid-19-open-science/?utm_source=Shared+Article+Sharing+Email+Alerts+/+E-Blasts+/+etc.&utm_campaign=Shared+Web+Article+Links
Stevens ML (2019) Medical philanthropy pays dividends. EMBO Rep 20: 1–6
The Association of the British Pharmaceutical Industry (2020) What are pharmaceutical companies doing to tackle COVID-19? Available at: https://www.abpi.org.uk/medicine-discovery/covid-19/what-are-pharmaceutical-companies-doing-to-tackle-the-disease/

License: This is an open access article under the terms of the Creative Commons Attribution 4.0 License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.