Prioritizing clinical research studies during the COVID-19 pandemic: lessons from New York City

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Coronavirus disease 2019 (COVID-19), caused by SARS-CoV-2, was first recognized in Wuhan, China in December 2019 (1) and rapidly spread through Asia and Europe, with the first US case identified in January 2020. The first New York case was reported in early March 2020 and by the first week in April, the New York Presbyterian Hospital system, including Columbia and Weill Cornell, had close to 2500 in-patients with over 700 patients on ventilators. The standard of care for COVID-19 was supportive care only and with significant morbidity and mortality recognized, there was a desperate need to identify effective treatments. Launching clinical research studies during this public health crisis required communication, collaboration, prioritization, keeping current with rapidly changing evidence, and maintaining high standards of scientific integrity and participant safety. An emerging crisis without proven treatments COVID-19 is a mild illness in more than 80% of people infected, but about 15% will require hospitalization and about 5% intensive care unit support (2). Patients most commonly present with symptoms of a viral respiratory illness, but the infection can provoke an intense immune response that leads to cytokine storm, coagulopathy, respiratory failure, and end-organ disease. Some patients progress to acute respiratory distress syndrome (ARDS) and require prolonged mechanical ventilation, sometimes with associated complications. Early reports from China and Europe documented wide-ranging approaches for COVID-19 […]

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An emerging crisis without proven treatments

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With a multitude of potential options, how does one pursue the most promising clinical trials? Although they are separate medical schools, Columbia and Weill Cornell share our major affiliated hospital, NewYork-Presbyterian. This structure facilitated communication and collaboration from the beginning, and consequently, the leaders of both Departments of Medicine and Divisions of Infectious Disease set up multidisciplinary committees to evaluate current data to prioritize and recommend initiation of COVID-19 clinical studies. The committees broadly represented the relevant medical disciplines and expertise in clinical trials at each institution. Furthermore, the deliberations and decisions made by each of the committees was communicated across the institutions to ensure that parallel trials were opened and that potentially harmful therapies were not pursued. Both committees received hundreds of study ideas — randomized clinical trials; single-arm pilot studies; expanded-access programs; compassionate-use studies, both multicenter and single-patient emergency use; and retrospective observational studies from a variety of sources — international, US government, pharmaceutical companies, and investigator initiated. At the same time, desperate patients, their families, and providers exerted pressure to use specific therapies and strategies. The support of the leadership of the medical schools and the hospital to focus on the most promising clinical studies across institutions was critical to standardizing approaches and advancing the science in service of our patients.

Testing the most promising drug candidates

Following an early review of available data, we prioritized studies of the investigational antiviral drug remdesivir, the repurposed drug hydroxychloroquine, the immunomodulator sarilumab, and convalescent plasma. Remdesivir demonstrated in vitro activity against SARS-CoV-2 (3) and a favorable safety record from prior clinical studies of Ebola virus. We pursued industry-sponsored randomized phase III multicenter studies of remdesivir in moderate and severe COVID-19 as well as parallel compassionate-use, and later, expanded-access, programs. Ultimately, we prioritized the randomized controlled clinical trials and contributed efficacy and safety data that demonstrated the clinical benefits of remdesivir in COVID-19 (4). The subsequent US FDA emergency use authorization (EUA) of remdesivir, albeit with a limited supply, posed additional challenges in determining which patients received treatment. To address this issue, we convened an expert group who reviewed the available data and developed a hospital-wide policy for remdesivir use.

Hydroxychloroquine, an FDA-approved drug for malaria and certain autoimmune diseases, demonstrated in vitro activity against SARS-CoV-2 (5) and was readily available. With early clinical trial data from China and Europe (6), no other available COVID-19 treatment, and provider and community pressure, we initially recommended hydroxychloroquine off label for hospitalized patients with COVID-19, along with a commitment to collect our data (7, 8). Ultimately, prospective clinical trials failed to demonstrate benefit (9), and the use of hydroxychloroquine and further studies were abandoned.

The cytokine storm associated with severe COVID-19 prompted suggestions for investigation of a host of immunomodulatory drugs — both agonists and antagonists directed at multiple steps of the immune cascade. We focused efforts on an early randomized, controlled clinical trial of sarilumab, an IL-6 inhibitor. Another IL-6 inhibitor, tocilizumab, was available off label and there was some tension...
Figure 1. Course of SARS-CoV-2 infection/COVID-19: clinical stages and potential interventions. Displayed pictorially are the clinical stages of disease resulting from SARS-CoV-2 infection and coronavirus infectious disease 2019 (COVID-19). Prior to exposure, preventive strategies may help avoid or abrogate infection. After infection, antivirals may reduce viral replication and resultant complications. If viral replication continues and an inflammatory response develops, immunomodulators could prevent or dampen an exaggerated immune response. If further progression with tissue destruction occurs, cellular therapies could promote tissue repair. Importantly, with novel preventive and therapeutic strategies, recovery from SARS-CoV-2 infection could improve at all stages. Figure adapted with permission from BioCentury (https://www.biocentury.com/trib-timeline).

Insights for future efforts

Acknowledging both accomplishments and missteps, what lessons can be learned from this extraordinary time? First and foremost, initiate broad communication and engagement of everyone concerned on a regular and ongoing basis: patients and their families; providers, scientists, and colleagues; research clinicians and staff; research administration at the IRB and contracts offices; medical school and hospital leaders; public affairs; and the media. Second, reach out to colleagues across disciplines to foster new ideas and initiate new collaborations. One example was to engage our transfusion medicine service and blood banks to establish a study to collect convalescent plasma from people who recovered from COVID-19 for later use in randomized clinical trials.

We initiated virtual scientific forums and town halls for basic, translational, and clinical researchers, and clinicians across multiple disciplines to come together, exchange knowledge, and promote discussion and plans. Also, we acknowledged and accepted conflict among stakeholders, but relied on our prioritization committees to advance what we thought were the best clinical trial ideas.

Third, use this broad dialog to evaluate and prioritize research plans and institute an “all hands on deck” strategy — engage research laboratories and clinical research units and redeploy research clinicians and staff as needed from other areas. Fourth, keep current with the challenging and rapidly changing field. Standing virtual meetings between the Depart-