Ethics of controlled human infection to address COVID-19

High social value is fundamental to justifying these studies


SUFFICIENT SOCIAL VALUE

CHIs have a long, complicated history. They have contributed to substantial improvements in clinical and public health practice, including the recent licensure of two vaccines (5), but also involved some unethical research (3). The first step in justifying SARS-CoV-2 CHIs, especially as they would involve major uncertainty and controversy, is to demonstrate their high social value. Crucially, SARS-CoV-2 CHIs should address relevant, unresolved scientific questions in rigorously designed and conducted experiments.

SARS-CoV-2 CHIs could have high social value in several ways. For example, they could help prioritize among the almost 100 investigational vaccines and over 100 experimental treatments for COVID-19 currently in development. CHIs could help identify the most promising agents, which would inform the design of larger trials, guide decisions to scale up manufacturing early, and thereby accelerate product development and implementation. If they saved even a few months of vaccine development (1), SARS-CoV-2 CHIs would contribute to faster control of the pandemic and reduce the need for, and associated costs of, physical distancing measures, providing substantial benefits for much of the world’s population (including the most vulnerable).

To achieve high social value in this way, coordination of stakeholders is essential. Sponsors of SARS-CoV-2 CHIs should delineate a credible path forward from CHIs to rigorous field studies, and eventually toward scaled-up production. This is a considerable challenge given the rapidly evolving research response to the pandemic; many approaches to accelerating product development are already appropriately being pursued in parallel. It is therefore essential to plan and evaluate SARS-CoV-2 CHIs as a complement, not an alternative, to these other approaches and ensure that CHI results are integrated into the dynamic COVID-19 research landscape. For example, the World Health Organization is convening sponsors of SARS-CoV-2 CHIs to increase transparency and promote coordination. Research sponsors should lead by establishing and enforcing standards for rapid data collection, dissemination, and sharing that permit aggregation of results across CHIs. Medical journals should require compliance with these standards before accepting manuscripts. Regulatory agencies should collaborate with sponsors, researchers, and policy-makers to define how CHI data will inform or modify larger trials, licensure, and manufacturing. Finally, sponsors and governments should implement mechanisms to ensure widespread, equitable access to proven products whose development was accelerated by SARS-CoV-2 CHIs. Such wide-ranging stakeholder coordination is difficult but important to demonstrate high social value. Though not achieved for proposed Zika virus CHIs during the 2015–2016 epidemic, it did occur later (6).

SARS-CoV-2 CHIs could have high social value in other ways, and individual CHIs could address multiple scientific questions. For example, CHIs could clarify dynamics of infection, viral pathogenesis, and risk of vaccine pathogenesis or identify correlates of protection—all of which could inform the development and implementation of vaccines. CHIs could also illuminate poorly understood parameters for modeling the pandemic and public health responses, including who is infectious and when and how infections occurred. This information is difficult to collect by observation alone, and existing animal models do not fully replicate clinical disease seen in humans. Additionally, if the pandemic wanes before larger trials are completed, SARS-CoV-2 CHIs could be critical for advancing research until the next outbreak, as with Zika virus (6). All of these paths to high social value would require similar, extensive coordination with relevant stakeholders.

SARS-CoV-2 CHIs admittedly have limited generalizability, as they would need to be conducted with low-risk populations (see below) with a non-natural mode of infection. Therefore, although some propose replacing efficacy trials with SARS-CoV-2 CHIs (7), it is more likely that CHIs accelerate vaccine or treatment development by informing larger trials, not by making such trials redundant. Yet almost all disease models or trial designs require some extrapolation or further testing. For example, field trials with frontline workers could also accelerate vaccine development, but they would not include older, retired individuals.

Thus, there are many potential ways in which SARS-CoV-2 CHIs could have high social value. Before their initiation, it is essential that the given social value is judged as compelling enough to justify its pursuit.
Ethical framework for SARS-CoV-2 controlled human infection studies (CHIs)

**SUFFICIENT SOCIAL VALUE**
Identify and address relevant, unresolved scientific questions in rigorously designed and conducted experiments
- Use rigorous methods to develop CHI models, including high-quality manufacturing and process of challenge strains
- Define and regularly review priority scientific questions, e.g., selecting the most promising vaccine and treatment candidates; identifying correlates of protection; clarifying infection dynamics, mechanisms of disease, and possible vaccine pathogenesis
- Coordinate with stakeholders to ensure SARS-CoV-2 CHI results will affect future research, clinical, or public health practice and delineate a credible path forward for CHI results to make an impact
- Establish and enforce standards for data collection in SARS-CoV-2 CHIs
- Share data, samples, and challenge strains appropriately
- Disseminate SARS-CoV-2 CHI results quickly through open-access publication

Realize benefits by facilitating equitable access to proven safe and effective products
- Use mechanisms such as compulsory licensure under Trade-Related Aspects of Intellectual Property Rights (TRIPS) agreement, march-in rights against patents under U.S. Bayh-Dole Act, U.S. Food and Drug Administration’s priority review voucher program

**CONTEST-SPECIFIC STAKEHOLDER ENGAGEMENT**
Engage public
- Create community advisory boards; use media to inform and engage
- Gather public input through informed opinion surveys
- Adapt engagement strategies to physical distancing as needed

Coordinate with international research, clinical, public health community
- Engage researchers, sponsors, regulators, ministries of health, etc. before, during, and after study implementation

**SUITABLE SITE SELECTION**
Consider feasibility of recruitment, risk, generalizability, availability of infrastructure, potential effects on local health care system

- Select location with available expertise
- Bring in extra resources so as not to unduly compromise pandemic response

**FAIR PARTICIPANT SELECTION**
Enroll low-risk groups with capacity to provide voluntary informed consent
- Enroll younger, adult participants without comorbidities

**ROBUST INFORMED CONSENT**
Ensure participant understanding
- Use evidence-based, context-specific consent materials
- Test participants on understanding of key criteria: deliberate infection; risks and burdens; potential social value; study purpose; uncertainty; restrictions on liberty to protect others

**PROPORTIONATE PAYMENT**
Avoid undue influence, exploitation, incentives to withhold information
- Compensate participants for their time to avoid exploitation and inequities in access to CHIs
- Set objective, verifiable eligibility criteria (in case money tempts participants to withhold disqualifying information)

**REASONABLE RISK–BENEFIT PROFILE**
Identify and reduce risks
- Enroll younger, adult participants without comorbidities; refine and update eligibility criteria in light of new evidence
- Monitor closely; provide prompt, free treatment and compensation for research-related injury
- Confine participants in in-patient isolation for at least 14 days
- Inform public health officials about study in advance

Ensure reduced risks are ethically acceptable
- Risks should not exceed upper limits
- Risks should be reasonable in relation to social value
adults, there is substantial consensus that risks to participants should not exceed an absolute upper limit. Regulations and ethics guidance do not clearly delineate this limit. Some commentators have argued that it should not exceed a 1% risk of death or the risks posed by activities that, like research, expose some people to risk to benefit others, such as living organ donation (8, 10). Although these are imperfect analogies to research, they provide helpful context for evaluating limits of acceptable research risk.

Current data on SARS-CoV-2 infection come from relatively small samples with missing data points and are still being scrutinized. Data suggest that 20- to 44-year-olds with diagnosed infection—including those with underlying conditions—have a mortality risk less than 0.2% (11). But diagnostic testing has been limited, making the number of undiagnosed infections unknown. One attempt to account for these limitations estimates that healthy adults aged 20 to 29 have a 0.03% risk of death and a 1.1% risk of hospitalizations (9). These risks could be further reduced by refining eligibility criteria based on emerging data. Recognizing the uncertainties, risks from SARS-CoV-2 CHIs appear comparable to the risks from some other research and activities similar to research (table S1). They also seem to fall below the upper risk limits proposed for research.

CURRENT CONTEXT SPECIFIC STAKEHOLDER ENGAGEMENT

CHIs have a checkered history (3), and it can be counterintuitive for the public that researchers would infect people with disease-causing pathogens. Although the current pandemic context with widespread physical distancing might complicate public engagement, it remains important and feasible as SARS-CoV-2 CHIs are developed. For example, public opinion surveys could identify concerns and information deficits, and researchers could engage the media or convene virtual advisory groups. Maintaining transparency and accountability to diverse communities is important for mitigating potential mistrust, especially in a pandemic (12). As noted above, engagement with stakeholders in the research community, health professionals, and policy-makers is also critical for ensuring that the results from SARS-CoV-2 CHIs translate into social benefits.

SUITABLE SITE SELECTION

Selecting suitable sites for SARS-CoV-2 CHIs requires considering risks to participants, study personnel, and third parties; feasibility of recruitment; availability of necessary infrastructure; and potential effects on local pandemic responses. Sites should be selected for sound scientific reasons while avoiding especially vulnerable populations. For example, performing CHIs in locations with high community spread of SARS-CoV-2 could be an acceptable way to reduce relative risks for participants, provided that high transmission is not due to underlying injustices. Given that participants would require testing, medical attention, and treatment, and research personnel would require personal protective equipment, sponsors should also demonstrate to ethics review boards or public health authorities that CHIs will not unduly compete for scarce resources and thereby compromise the local pandemic response. All sites should have sufficient capacity to conduct rigorous studies, provide high-quality care to participants, and minimize research risks. Sites experienced with conducting CHIs might be favored to ensure that studies and local public engagement can be launched quickly, effectively, and responsibly.

FAIR PARTICIPANT SELECTION

Selecting participants fairly for SARS-CoV-2 CHIs primarily requires considering fair distribution of research risks and burdens. Because of the uncertainty and potential high risk involved, participants who are at relatively low risk of serious and irreversible harm and have capacity to give their own consent should be selected (i.e., young, healthy and competent adults).

ROBUST INFORMED CONSENT

There is widespread consensus on obtaining high-quality informed consent for CHIs and using rigorous procedures to maximize participant understanding. Evidence-based approaches to consent include requiring participants to pass a test on key study information (14). Ongoing informed consent will be important as new data emerge, notably on the risks of SARS-CoV-2 infection.

PROPORTIONATE PAYMENT

Members of our group disagree about the ethical permissibility of offering payment to CHI participants, and there may be relevant regulatory limits in different jurisdictions. Nevertheless, as SARS-CoV-2 CHIs require confinement and follow-up, fairness seems to demand offering participants compensation for their time. This may total several thousand dollars in the United States, assuming compensation at a fair minimum wage for unskilled labor, as in other CHIs. By contrast, incentives beyond compensation could be avoided, given the number of people already indicating willingness to participate. Concerns that the undue influence of monetary compensation compromises risk judgments are unsupported by the available data, as financial motivations are associated with greater attention to risk (15). Moreover, a rigorous informed consent process could maximize understanding. In case payment tempts participants to withhold disqualifying information, eligibility criteria should be objectively verifiable.

CONCLUSION

Given the extraordinary nature of the pandemic, our framework and analysis support laying the groundwork for SARS-CoV-2 CHIs—for example, by developing a challenge strain, drafting consensus protocols that address ethical concerns, and engaging stakeholders to enhance their social value, minimize risks, and build public trust.

REFERENCES AND NOTES

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ACKNOWLEDGMENTS

The opinions expressed in the article are the authors’ and do not reflect the views of organizations with which the authors have affiliations, including the National Institutes of Health, the Department of Health and Human Services, or the United States government. This work was primarily supported by a Making a Difference Grant from the Greenwall Foundation (S.K.S., A.R., R.P., D.D.) along with support from the Wellcome Trust (S.K.S., E.J., D.K., M.K., M.J.S., V.V.), Brocher Foundation (S.K.S., A.R., R.P., D.D., T.C.D., H.F.L., E.J., N.S.J., D.K., J.K., D.M., S.C.M., T.L.R., M.R., A.S., M.J.S., V.V.), and NIH Clinical Center Department of Bioethics (A.R.). The authors also thank C. Chui, K. Littler, P. Pittsutthithum, and M. Yu for their contributions, and M. Danis, C. Grady, M. Nicolini, J. Ochoa, and H. Taylor for helpful discussion.

SUPPLEMENTARY MATERIALS

Published online 7 May 2020 10.1126/science.abc1076
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Science 368 (6493), 832-834.
DOI: 10.1126/science.abc1076 originally published online May 7, 2020

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