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Advancing precision medicine for acute respiratory distress syndrome

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Acute respiratory distress syndrome (ARDS) is a heterogeneous clinical syndrome. Understanding of the complex pathways involved in lung injury pathogenesis, resolution, and repair has grown considerably in recent decades. Nevertheless, to date, only therapies targeting ventilation-induced lung injury have consistently proven beneficial, and despite these gains, ARDS morbidity and mortality remain high. Many candidate therapies with promise in preclinical studies have been ineffective in human trials, probably at least in part due to clinical and biological heterogeneity that modifies treatment responsiveness in human ARDS. A precision medicine approach to ARDS seeks to better account for this heterogeneity by matching therapies to subgroups of patients that are anticipated to be most likely to benefit, which initially might be identified in part by assessing for heterogeneity of treatment effect in clinical trials. In October 2019, the US National Heart, Lung, and Blood Institute convened a workshop of multidisciplinary experts to explore research opportunities and challenges for accelerating precision medicine in ARDS. Topics of discussion included the rationale and challenges for a precision medicine approach in ARDS, the roles of preclinical ARDS models in precision medicine, essential features of cohort studies to advance precision medicine, and novel approaches to clinical trials to support development and validation of a precision medicine strategy. In this Position Paper, we summarise workshop discussions, recommendations, and unresolved questions for advancing precision medicine in ARDS. Although the workshop took place before the COVID-19 pandemic began, the pandemic has highlighted the urgent need for precision therapies for ARDS as the global scientific community grapples with many of the key concepts, innovations, and challenges discussed at this workshop.

Introduction

Acute respiratory distress syndrome (ARDS) occurs in a quarter of all critically ill patients who require mechanical ventilation. Despite considerable gains in preventing ventilation-induced lung injury, ARDS is still associated with substantial mortality and long-term morbidity among survivors. 1–3 To date, no specific pharmacotherapy has proven effective against ARDS. Countless agents that showed promise in preclinical studies have been ineffective in human trials, a gap attributed in part to clinical and biological heterogeneity in human ARDS. 4 A precision medicine approach is intended to address explicitly how such underlying heterogeneity influences response to therapy among different patients with the same diagnosis. 5

A limited ability to rigorously identify potential sources of heterogeneity has hampered feasibility of a precision medicine approach to ARDS. Growing evidence suggests that subsets of patients, identified through combined clinical–molecular multivariable phenotyping, might exhibit differential responses to therapies deemed ineffective for the overall population, 6–9 renewing hope for the application of a precision medicine approach to ARDS.

Motivated by these advances, the Division of Lung Diseases within the US National Heart, Lung, and Blood Institute (NHLBI) convened a workshop to establish research opportunities and explore potential roadblocks for accelerating precision medicine in ARDS. Invited experts in preclinical studies, human translational research, and clinical trials of ARDS were joined by experts in precision medicine and adaptive trial design outside the field of ARDS. This Position Paper summarises presentations and discussions from the group, weighing the current state of research relevant to advancing precision medicine in ARDS and proposing strategic coordination of future research from bench to bedside.

Key messages

- Mortality and morbidity from acute respiratory distress syndrome (ARDS) remain high; phenotypic heterogeneity (clinical and biological) is inherent in ARDS and is likely to influence response to therapy, yet remains poorly understood
- Preclinical models are invaluable for understanding specific biological processes and identifying targetable nodes in pro-injury and pro-resolution pathways, yet cannot fully replicate ARDS heterogeneity
- Multicentre observational cohort studies that collect clinical, physiological, radiological, and biological data and samples in a harmonised and pragmatic manner will facilitate deep phenotyping of patients and enable identification of mechanistic pathways for subsequent interrogation with preclinical models (ie, reverse translation) and for potential targeted intervention
- Development of rapid, practical diagnostic assays that can yield results within minutes to a few hours to support predictive and prognostic enrichment approaches will be important for molecular signature-guided therapies in trials or clinical practice
- Platform trials with a master protocol, with or without adaptive features, could facilitate efficient simultaneous or sequential testing of multiple candidate therapies, accelerate detection of treatment-responsive subgroups, and create a discovery pipeline for ARDS pharmacotherapies drawing from repurposed or novel drugs
- Discovery is accelerated by collaboration and coordination among key stakeholders, including sponsors, regulatory agencies, industry, academia, patient advocates, and the broader medical community

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Panel 1: Summary statements developed by workshop participants: rationale for precision medicine in ARDS

- Phenotypic heterogeneity (clinical and biological) is inherent in ARDS and is likely to influence response to therapy, yet remains poorly understood; better understanding of heterogeneity is essential for identifying treatments and minimising harm.
- Current proven treatments involve lung-protective ventilation and optimisation of supportive care, but mortality and morbidity from ARDS remain high; pathways involved in pathogenesis, resolution, and repair include multiple potential targets for therapeutic development.
- ARDS is a clinical diagnosis; clinical features alone might not distinguish biological heterogeneity with sufficient precision to match targeted therapies to patients in whom the relevant pathway is most active.

RESUS—acute respiratory distress syndrome.

The rationale for precision medicine in ARDS

The concept of precision medicine encapsulates what practising clinicians strive to do every day: deliver care tailored to the individual patient that maximises potential benefit and minimises risks. The extent to which this approach is attainable in practice might depend partly on the ability to prospectively characterise patients according to likely treatment responsiveness. Other fields that pair molecular diagnostics with targeted therapies (e.g., BRAF inhibitors for melanoma or targeting of receptors for oestrogen, progesterone, or human epidermal growth factor receptor 2 in breast cancer11,12) or that identify and refine biomarker-defined subgroups for targeted therapies through rigorous cohort characterisation (e.g., subphenotyping severe asthma13) provide important examples of the value of identifying and targeting treatment-responsive subphenotypes to improve patient outcomes. ARDS by definition is an inherently heterogeneous clinical syndrome. Therapeutic discovery is likely to be accelerated by subphenotyping patients with ARDS according to mechanistic drivers that can be made clinically accessible and actionable.

Clinically overt sources of heterogeneity

ARDS is a diffuse lung injury characterised by alveolar inflammation and disruption of the alveolar–capillary barrier.1 In practice, however, neither alveolar inflammation nor barrier function is measured routinely owing to a lack of well validated, widely available tools for measurement. Instead, ARDS diagnosis has broad syndromic criteria and reflects the collective views of the multidisciplinary expert author group.

From Oct 22 to 23, 2019, multidisciplinary experts in ARDS, as well as in precision medicine and adaptive trial design, convened in Bethesda, Maryland, for the NHLBI workshop. The workshop, entitled Precision Medicine in ARDS, was developed and chaired by CSC, BTT, and NRA. The workshop chairs, with input from the session moderators (JCM, RMB, MNG, TRM, JAB, TJS, MM) through a series of pre-workshop conference calls, organised the workshop into three sessions to discuss thematic questions that broadly defined current and future research directions to advance precision medicine in ARDS. The resulting thematic questions concerned:

1. the state of ARDS research relevant to advancing precision medicine in ARDS;
2. what can be learned from other disease areas that have made advances in precision medicine; and
3. pressing considerations that need to be addressed to design future clinical trials to advance precision medicine in ARDS.

The workshop’s chairs and session moderators assigned participants topics to present according to their expertise. Before the workshop, participants prepared briefs summarising the material to be presented, and slides were submitted to the chairs for feedback. At the workshop, presentations were followed by group discussions aimed at synthesising key messages for each content area. A summary discussion at the end of the workshop attended by all participants and led by the co-chairs resulted in draft consensus statements prepared by JRB and submitted to all participants for revision, which was achieved through electronic communication after the meeting, to arrive at the final summary statements presented here (panels 1, 2).

The content of this Position Paper was developed from presentations and discussions during the workshop and revised with input from all authors, and reflects the collective views of the multidisciplinary expert author group.
fractional concentration of oxygen in inspired air (FiO₂) correlate with higher incidence of diffuse alveolar damage on autopsy, but the PaO₂/FiO₂ ratio is influenced substantially by ventilator settings and does not consistently differentiate treatment responsiveness. Stratification of ARDS by PaO₂/FiO₂ ratios has yielded some success, however, with positive trial results for a proning strategy in patients who have a PaO₂/FiO₂ ratio of less than 150; whether patients with higher PaO₂/FiO₂ ratios might respond to the same interventions is less clear.

Ventilation-induced lung injury might be an important modifier of treatment responsiveness, but the differential effect of ventilator strategies on lung injury is not readily detectable at the bedside. Instead, global respiratory mechanics and the degree of ventilator support required are routinely used to characterise patients with ARDS. Distribution of lung injury (sometimes categorised radiographically as focal or diffuse ARDS) signifies regional mechanical heterogeneity, correlates with biomarkers of alveolar epithelial injury, and has prognostic value, but is of unclear value in predicting treatment response and is not consistently characterised in practice.

ARDS has often been characterised as direct (pulmonary) or indirect (extrapulmonary) on the basis of whether the injurious risk factor originated in the lungs (eg, pneumonia, aspiration) or elsewhere (eg, abdominal sepsis, pancreatitis, transfusion-related). Direct ARDS is associated with more alveolar epithelial injury, less endothelial injury, and greater risk of progression to fibrosis than indirect ARDS. Clinical predictors of mortality might also differ between direct and indirect ARDS. However, identifying the cause and origin of lung injury is fraught with imprecision. Patients often have no identifiable risk factor, and direct and indirect injury often occur together. Additionally, categorisation by direct or indirect precipitant does not address whether the injurious stimulus is transient (eg, transfusion, trauma, aspiration) or sustained over days (eg, infection, pancreatitis).

Other clinically overt factors could influence disease course and therapeutic responsiveness in ARDS.
Supportive therapies including differing approaches to fluid management and sedation contribute additional clinical heterogeneity that might influence both patient outcomes and therapeutic responsiveness.27,28 Some commonly prescribed medications used to treat underlying diseases, including β blockers and β agonists, statins, and inhaled and systemic corticosteroids,3,23-31 might directly influence host response to lung injury. Environmental factors such as smoking, alcohol use, and ambient air quality might modify the biological response to pulmonary insults,30-33 In infection-associated ARDS, different pathogen species and strains precipitate lung injury via different virulence mechanisms and pathogen-host interactions.34 Although evaluating heterogeneity in respiratory mechanics has proven useful to guide interventions targeting ventilation-induced lung injury, clinically obvious heterogeneity has been less helpful for therapies targeting other pathways of ARDS pathogenesis.35

**Clinically occult sources of heterogeneity**

Early studies of the molecular biology of ARDS indicated that substantial heterogeneity exists between patients.36 Biomarkers of key pathways including inflammation, coagulation, alveolar epithelial injury, and vascular endothelial activation can differ considerably between patients,30-32 suggesting a role for molecular subphenotyping in unravelling heterogeneity.

Analyses of clinical and biological data from trials and observational cohorts have identified two subphenotypes, distinguished in part by degree of inflammation, circulatory shock, and multiorgan failure, which appear to have differential responses to multiple treatments in secondary analyses of clinical trials.32,40,41 Parsimonious prediction models based on these analyses42 warrant prospective testing.

Better understanding of the drivers of molecular heterogeneity might help to identify new therapeutic targets. Clinical heterogeneity undoubtedly accounts for some biological variation, but not all. Genetic predisposition helps to shape the response to infectious and sterile pro-inflammatory stimuli that might precipitate ARDS, yet also influences resolution and repair.43 For certain lung infections, genetic polymorphisms influence the pathogen-host interaction directly.44 Genetically predicted plasma concentrations of sRAGE (soluble receptor for advanced glycation end products, an inflammatory mediator and alveolar epithelial injury marker) and angiopoietin 2 (a marker of endothelial activation) are associated with risk of ARDS during sepsis.45,46 Polymorphisms in genes that encode cell matrix proteins, vascular endothelial growth factor receptor 1, and haptoglobin also correlate with ARDS risk.47-50 Together, these findings suggest that genetic polymorphisms could be an important source of biological heterogeneity in lung injury, resolution, and repair.

Differences in the lung and gut microbiomes might also contribute to ARDS heterogeneity. Although proximal airways are lined with commensal bacteria, healthy alveoli are resistant to bacterial growth, providing few nutrient substrates and containing bactericidal surfactant.33 However, as lung injury develops, alveolar flooding by protein-rich liquid and surfactant inactivation together create a more favourable environment for bacterial growth, which might drive further inflammation and injury in a positive feedback loop.51 Lung bacterial overgrowth might occur, and translocated gut bacteria and related pathogen-associated molecular patterns could enter the alveolus.52 Therapeutic implications of this dysbiosis are not well understood.

**Challenges for precision medicine in ARDS**

Workshop participants agreed that advancing precision medicine for ARDS holds considerable appeal and has a strong rationale (panel 1), but also faces numerous hurdles. Major obstacles include limited understanding of the key nodal points in the multiple pathways that mediate acute lung injury, and the difficulty of prompt identification of patients in whom specific pathways are deranged. The pathogenesis of acute lung injury involves several complex steps, including increased endothelial permeability, alveolar epithelial cell death and dysfunction, loss of surfactant function, activation of coagulation cascades, and triggering of complex innate immunity pathways in the lungs. The key nodes of these pathways that govern pivotal downstream events towards resolution and repair remain uncertain, and the importance of a specific pathway might differ between patients depending on cause of lung injury, patient predisposition, and other factors.

Adding to the challenge, the timeframe for characterising patients is short because ARDS develops rapidly, and treatments might be most effective when initiated early. For example, current precision medicine approaches to staging and molecular classification of breast and lung cancer require days to weeks, yet in a recent trial of ARDS, 48-h mortality was 10% and 96-h mortality was 16%.55 Rapid assays will need to be developed to enable prompt biological phenotyping within minutes to a few hours.56 Proximate biospecimens, such as those from bronchoalveolar lavage, require additional invasive procedures and have been obtained in a minority of large-scale human studies; even when collected, specimen acquisition and analysis protocols often differ between studies. Tissue samples are rarely obtained, both because of procedural risk and because the finding of diffuse alveolar damage does not substantially change treatment.57,58 Acute and chronic comorbidities vary and can influence the likelihood of developing and surviving ARDS.59 Even if comorbidities do not modify biological treatment responsiveness, they undoubtedly influence risk of death attributable to ARDS. To add further complexity, there is no widely
accepted surrogate endpoint for clinical trials in ARDS, complicating early-phase clinical studies.

Adapting preclinical investigations to precision medicine

Roles for rigorous preclinical models

Animal models will never fully replicate the heterogeneity of human ARDS. Nevertheless, they remain essential for understanding more deeply the cellular and molecular drivers of disease and have an important role in the development of a precision medicine strategy for ARDS (panel 2). Animal studies can be highly effective in modelling a specific biological feature of lung injury or treatable trait (eg, endothelial barrier disruption), particularly when coupled with careful observation in human studies that can feasibly identify biomarkers of those targeted pathways.

Among the most powerful roles of animal models is in helping to define key nodes of mechanistic pathways that have been identified in rigorous human studies, a bedside-to-bench approach termed reverse translation (figure). Once identified, these pathways and nodes could inform targets for drug development to be tested in subsequent trials.

Recent efforts to incorporate precision medicine in paediatric sepsis care offer an important example of how animal models could inform development of precision medicine for ARDS. Through a series of cohort studies, investigators developed a biomarker panel to predict mortality from paediatric sepsis in humans and subsequently found similar biomarkers were predictive of mortality in murine models of sepsis. The murine models were then used to test therapies targeting the predictive biomarkers to determine whether they were causally related to outcomes and to identify other biological features (eg, higher bacterial burden) of the group with high predicted mortality. This conservation of findings from bedside to bench might permit initial testing of candidate therapies in preclinical models to understand the implications of biological heterogeneity and potentially inform trial design.

Reverse translation might also have an important role in understanding why some promising candidate treatments are not beneficial in human clinical trials. For example, heterogeneity of treatment effect by latent class determinants of a differential response to simvastatin but not in a related trial of rosuvastatin for ARDS. Taking this clinical observation back to relevant preclinical models could help to identify pivotal mechanistic determinants of a differential response to simvastatin when compared with rosuvastatin for clinically defined ARDS subphenotypes, and also potentially identify novel, mechanistically driven biomarkers that better predict a beneficial treatment response to one statin versus another.

Modelling sources of human heterogeneity

Introducing heterogeneity, inherent in human ARDS, into preclinical models might contribute to understanding of downstream biological consequences and enable identification of potentially novel nodes for therapeutic
targeting. This strategy can take many forms, including but not limited to modifying host susceptibility, varying insult type or intensity, and altering environmental or supportive care factors. Crossbreeding of mice to introduce genetic diversity might inform understanding of genetically regulated complex traits ranging from inflammation to wound repair. The Collaborative Cross model for complex trait analysis is one such example, a collection of roughly 200 distinct recombinant inbred strains of mice created by reciprocal inter-cross breeding of eight founder strains. This model could be integrated with human genome-wide association studies to dissect a genetic basis for ARDS heterogeneity.

Adding variation to mechanism and intensity of the lung injury stimulus might also help to elucidate differential activation of mechanistic pathways and resulting effects on treatment responsiveness. For example, in the caecal ligation and puncture model of experimental sepsis, caecal puncture of random severity can be performed in outbred mice to yield a heterogenous insult in genetically heterogeneous animals, facilitating interrogation of biological pathways that might influence heterogeneity of host response and treatment effect. A study using a caecal ligation and puncture model with variable severity in outbred mice identified two distinct sepsis phenotypes: a high-mortality variant characterised by early onset of cardiogenic shock; and a low-mortality variant with intact cardiac function, which exhibited differential responses to treatment with hydrocortisone, ascorbic acid, and thiamine.

Multisystem crosstalk and modifiable risk
Acute and long-term extrapulmonary morbidity, including physical, cognitive, and psychiatric disability, are common in ARDS. Bidirectional pathogenic interactions of the lungs with the kidneys, brain, and other vital organ systems have been discovered. For example, acute kidney injury might predispose the lungs to a secondary inflammatory insult, precipitating lung injury that in turn could exacerbate renal injury. Similarly, endothelial activation and neuroinflammatory signalling from brain injury might predispose to lung injury, which in turn could exacerbate brain injury. Many pathways of multiorgan crosstalk are similar across critical illness syndromes and thus might be considered clinically in terms of treatable traits rather than being unique to any one particular diagnosis. Deciphering mechanisms of multisystem interactions, with preclinical models and human data, could aid development of interventions that attenuate such multiorgan positive feedback crosstalk loops and associated morbidity in various critical illness syndromes, including ARDS.

Human ex-vivo and in-vivo models
Human experimental models of lung injury might have a unique role in improving understanding of ARDS pathophysiology. Ex-vivo human lung models, using organs declined for transplant, allow experimental manipulation and serial evaluation of distal lung tissue that otherwise is typically inaccessible in human ARDS. In-vivo human models range from subclinical lung inflammation in healthy volunteers induced by inhaled lipopolysaccharide to surgeries such as cardiopulmonary bypass or intra-operative single-lung ventilation that induce lung inflammation or ischaemia reperfusion injury in a more regulated setting. As an example of potential relevance to ARDS heterogeneity, differential inflammatory responses to lipopolysaccharide have been linked to genetic polymorphisms that modulate innate immunity in healthy volunteers. However, human models have important limitations. For instance, ex-vivo models fail to capture multisystem crosstalk and are technically difficult to develop and maintain. In-vivo models yield low-level regulated inflammation, which might be informative but contrasts with the dysregulated inflammation, physiological derangements, and severity of lung injury characteristic of ARDS.

Clinical cohorts to advance precision medicine
Large observational studies
Rigorous cohort studies are essential to understand the entire spectrum of ARDS pathogenesis and recovery in usual care. These natural history experiments serve as a crucial link in the translational science continuum, informing research directions in preclinical studies and clinical trials (panel 2). Key aims for ARDS cohort studies include the following: to identify, validate, and refine clusters of heterogeneity (subphenotypes) and factors associated with treatment responsiveness; to develop rapid diagnostics in support of such subphenotyping; to explore underlying mechanisms and potential drivers of biological heterogeneity; to identify populations at greater risk of ARDS-attributable outcomes for clinical trial prognostic enrichment; to evaluate generalisability of clinical trial findings to less selective populations; and to identify novel interventions for testing in preclinical models and clinical trials. Cohort studies could also enrol patients at risk of ARDS to determine why some patients progress to severe lung injury whereas others do not.

The many sources of clinically overt heterogeneity in ARDS necessitate harmonised clinical data and biospecimen collection and sufficiently large sample sizes to ensure statistical power for dissecting clinically occult heterogeneity. An international period prevalence study suggested that, before the COVID-19 pandemic, the typical 12-bed intensive care unit might have seen approximately 65 patients with ARDS per year (5·5 cases per bed per year). Multicentre cohort studies are well suited to ensuring timely accrual and diversity (ethnic, phenotypic, and biological) that is representative...
of the total population of patients with ARDS, as required for discovery and validation of subphenotypes. Encouraging broader patient participation in clinical research studies, with the involvement of patient advocates, could further enhance development of these larger datasets.

Harmonising data and specimen collection
Observational research networks in other fields provide a template for building synergistic research teams and comprehensive cohorts to advance precision medicine. For example, the NHLBI Severe Asthma Research Program is a uniquely structured multicentre collaborative network of investigators who lead independently funded mechanistic studies but are required to recruit patients and obtain samples that are shared across the network. The result—a rigorously phenotyped cohort of more than 700 patients with pulmonary function measures, imaging, and multiple biological samples—enabled identification of several asthma subphenotypes and key mechanistic nodes that are now being targeted in clinical trials.

The Severe Asthma Research Program offers several lessons that are relevant to ARDS. The funding mechanism explicitly mandated a team science approach, wherein multiple investigators contributed to a shared cohort with individual mechanistic studies championed by a specific investigator and embedded within the cohort. This shared cohort model required that all investigations have a common longitudinal protocol, with uniform data acquisition and analysis procedures, while allowing some highly specialised studies to involve only a subset of sites.

Development of rapid diagnostics
Because ARDS develops and evolves quickly, the development of rapid diagnostics that can be deployed on-site (ie, at the bedside or in the hospital clinical laboratory) will be crucial for timely biological phenotyping. Assay platforms will need to generate actionable data within minutes to a few hours, rather than days, to be implemented in precision medicine trials. Given current technologies and the existing body of evidence, protein-based enrichment strategies appear closest to clinical application, when compared with gene expression (mRNA)-based enrichment strategies. The development of these assay platforms will be most efficient via collaborations with industry and with early involvement of regulatory agencies.

Clinical trials to advance precision medicine
Predictive and prognostic enrichment
Enrichment strategies are essential to enable precision trials among critically ill patients. Enrichment broadly refers to the selection of a patient cohort in whom an experimental intervention is more likely to be of benefit, when compared with an unselected cohort. Prognostic enrichment entails selecting patients who are more likely to have a disease-related event—eg, those at higher risk of ARDS-related death. Prognostic enrichment decreases the sample size required to detect relative differences in a trial endpoint for a given desired power. However, it does not address heterogeneity in treatment response. Predictive enrichment entails selecting patients anticipated to have an increased likelihood of responding to an intervention on the basis of clinical or biological characteristics and mechanism of action of the intervention. The challenge for predictive enrichment is that it is difficult to prospectively identify subgroups most likely to benefit from specific therapies. When treatment response heterogeneity is plausible, trials should be designed to allow for that possibility.

Lessons from other fields: innovative trial designs
Innovative trial designs might help to accelerate precision therapeutic discovery, in part by embracing disease heterogeneity in the trial design. Leveraging of disease heterogeneity in the trial design
Drug effects are assessed in each of ten prespecified biomarker signatures

Use of analyses that incorporate biological subtypes
The main outcome is Bayesian probability of success in a subsequent confirmatory phase 3 trial for each prespecified biomarker signature

Use of efficient platform trial design
A master protocol with continuous patient enrolment is used, enabling simultaneous evaluation of up to five candidate therapies; therapies can be added to or removed from the protocol without interrupting enrolment

Personalisation of therapy within a trial to maximise benefit to each patient
Therapy can be escalated or deescalated on the basis of each individual’s response to therapy, according to the pathological complete response

Early and transparent collaboration with multiple drug companies in a single platform trial
Investigators determine the most promising drug candidates and work with Quantum Leap Healthcare Collaborative, a non-profit organisation, to secure drugs from multiple companies

Predictive enrichment

| I-SPY2 trial<sup>a</sup> for breast cancer | PrecISE trial<sup>b</sup> for severe asthma |
|-----------------------------------------|------------------------------------------|
| Leveraging of disease heterogeneity in the trial design | Drug effects are assessed in each of ten prespecified biomarker signatures |
| Use of analyses that incorporate biological subtypes | The main outcome is Bayesian probability of success in a subsequent confirmatory phase 3 trial for each prespecified biomarker signature |
| Use of efficient platform trial design | A master protocol with continuous patient enrolment is used, enabling simultaneous evaluation of up to five candidate therapies; therapies can be added to or removed from the protocol without interrupting enrolment |
| Personalisation of therapy within a trial to maximise benefit to each patient | Therapy can be escalated or deescalated on the basis of each individual’s response to therapy, according to the pathological complete response |
| Early and transparent collaboration with multiple drug companies in a single platform trial | Investigators determine the most promising drug candidates and work with Quantum Leap Healthcare Collaborative, a non-profit organisation, to secure drugs from multiple companies |

Table: Key principles shaping precision medicine trials in breast cancer and severe asthma

<sup>a</sup> For the Severe Asthma Research Program see www.severeasthma.org
heterogeneity and uncertainty around matching the right therapy to the right patient. Platform trials evaluate multiple therapies for a disease simultaneously against a single control group.8 Platform trials typically are designed to permit the addition and removal of therapies without stopping the trial, according to predefined rules regarding efficacy and futility in the overall trial population or particular subgroups. The I-SPY2 and PrecISE trials are examples of adaptive platform trials (table).79,80 I-SPY2 has been designed to evaluate neoadjuvant therapies for high-risk, early stage breast cancer,11 while therapies for severe and exacerbation-prone asthma are being assessed in PrecISE.81 Both are signal-finding phase 2 trials that aim to rapidly evaluate multiple candidate therapies for their probability of success in phase 3 trials.

I-SPY2 and PrecISE offer several lessons that might be relevant to ARDS. As platform trials, they enrol continuously using a single master protocol and evaluate multiple treatments simultaneously.79,80 A given therapy could be eliminated from the trial and replaced by a new candidate therapy without interrupting enrolment, so that more therapies are tested in a time-efficient and cost-effective manner. Both trials embrace a team science approach and engaged key collaborators and stakeholders early in trial formulation, including sponsors, regulatory agencies, pharmaceutical and biotechnology industries, academia, patient advocates, and the broader medical community.

Both I-SPY2 and PrecISE leverage the inherent biological heterogeneity of their respective targeted phenotypes. The main outcome of I-SPY2 is the predictive probability of success in a confirmatory phase 3 trial within each of ten prespecified biomarker signatures; a therapy is deemed sufficiently promising to progress if there is an 85% Bayesian predicted probability of success in a 300-patient, 1:1 randomised, confirmatory neoadjuvant phase 3 trial for any of the prespecified biomarker signatures.12 The trial design includes the use of standard (US Food and Drug Administration [FDA]-approved), qualifying (highly promising, and prespecified for FDA validation), and exploratory (hypothesis-generating) biomarkers,7 and uses incoming data to identify the best biomarkers for stratifying patients for each of the therapies under investigation. Over a decade, more than 20 agents from multiple pharmaceutical companies have been evaluated in I-SPY2, several agents have progressed to phase 3 trials and have been validated, spanning a range of biomarker signatures.81 In addition, the prognostic importance of an early endpoint—pathological complete response to therapy regardless of subtype or treatment delivered—has been established,82 and new tumour classifiers have been identified that enhance the ability to target therapies and improve outcomes.

In PrecISE, novel candidate therapies are chosen according to their hypothesised effectiveness for biological subtypes of severe asthma based on a priori beliefs about underlying mechanisms. A crossover design allows each patient to serve as his or her own control and, through repeated randomisations, receive multiple therapies during the study. Randomisation at every step is weighted (initially in a 2:1 ratio) so that patients in the biomarker subgroup targeted by a particular intervention are more likely to receive that intervention, and definitions of the biomarker-defined subgroups are updated over time as trial data accumulate.83 This approach allows for detection of drug effects in subgroups not anticipated to benefit from the drug as well as confirmation of non-responders, which are essential for targeted treatments. At the conclusion of this phase 2 signal-finding trial, novel therapies demonstrating efficacy across three dimensions of asthma severity (lung function, symptom control, and exacerbations), and the optimally defined disease subtype that each therapy should target, will be identified for further study in a phase 3 confirmatory trial.

Although I-SPY2 and PrecISE are phase 2 trials, platform trials similarly might improve efficiency when multiple candidate therapies amenable to the same platform are ready for confirmatory phase 3 testing.

Early lessons from COVID-19

After the October 2019 workshop, the SARS-CoV-2 virus, which causes the clinical syndrome of COVID-19 with severe ARDS, has spread across the globe. Spurred by the urgent need for novel therapies, many clinical trials groups have embraced platform trials to accelerate scientific discovery. The RECOVERY platform trial, run through the UK’s National Health Service, was the first to demonstrate a survival benefit with corticosteroids for severe COVID-19,84 a finding since replicated in several subsequent trials.85 In addition, the RECOVERY platform design rapidly evaluated and discarded several other repurposed therapies (eg, hydroxychloroquine, lopinavir–ritonavir) as ineffective in the population studied.

In the USA, the Accelerating COVID-19 Therapeutic Interventions and Vaccines (ACTIV) and Collaborating Network of Networks for Evaluating COVID-19 and Therapeutic Strategies (CONNECTS) public–private partnerships with government, academia, and industry aim to rapidly assess promising candidate therapies for COVID-19 with a series of federally funded master protocols and platform trials that use National Institutes of Health-supported research networks to test prioritised agents, including immune modulators, monoclonal antibodies, anti-thrombotics, and other therapeutics.86,87 Growing out of this NHLBI workshop, the I-SPY COVID phase 2 platform trial has fostered novel collaborations, pairing the precision medicine expertise of oncologists with the content expertise of ARDS clinical trialists, with repurposed and investigational agents under evaluation that target several mechanistic pathways in severe COVID-19.88

Worldwide trial efforts have been coordinated by the WHO Solidarity Trial Consortium90 and the REMAP-CAP
Trial investigators. REMAP-CAP was planned before the COVID-19 pandemic as a perpetual enrolment adaptive platform trial studying treatments for community-acquired pneumonia. Its unique platform design was intended for rapid adaptation to respiratory pandemics and illustrates successful implementation of such trial design principles on a global scale.

Although COVID-19 developments post-dated this workshop, they demonstrate the potential for rethinking trial design to accelerate therapeutic discovery for conditions for which few evidence-based treatments exist. They also illustrate the importance of public–private partnerships with broad stakeholder engagement to achieve this vision. COVID-19 exhibits substantial phenotypic heterogeneity despite being precipitated by a single causative organism, highlighting the need for deeper understanding to support a precision medicine approach.

Potential application of platform trial design to ARDS
Therapeutic discovery for ARDS, from COVID-19 or other causes, could be accelerated through platform trials. Building efficient trial machinery with perpetual enrolment to test multiple agents could help to establish a cost-effective drug-development pipeline. Efficiency of the platform design, access to existing trial infrastructure, and input from academic physician-scientists could incentivise pharmaceutical industry collaboration to develop and test new and repurposed drug candidates.

The inherent heterogeneity of ARDS could be leveraged by designing trials to differentiate treatment responsiveness by biomarker signature, potentially to inform predictive enrichment for a subsequent phase 3 trial (figure). Key hurdles include uncertainty about temporal stability of biomarker signatures over time and lack of rapid diagnostics for molecular biomarkers, which would need to be developed and validated before or in parallel with any such trial. An adaptive design could improve efficiency by using data from the ongoing trial to refine subgroup-intervention pairings or eliminate biomarker-defined subgroups from consideration for a particular therapy. Whether response-adaptive randomisation might boost efficiency further depends on the particular construct of the trial. For example, response-adaptive randomisation might improve both statistical efficiency and effectiveness of therapies in a trial when the following conditions are met: at least three arms are available for allocation (including, typically, a standard-of-care control arm); response-adaptive randomisation is not implemented until there is sufficient early data (eg, a burn-in phase) to provide reasonably stable preliminary estimates of treatment effect; and a sufficient fraction of patients are allocated to the control group in an ongoing manner to mitigate risks of bias from changes over time.

Although the efficiencies of platform trials are appealing, some challenges remain. As the eligibility criteria for a platform trial might be broad to capture the range of patients affected by ARDS, careful consideration should be given to whether all treatment arms should be made available to the full range of enrolled participants. Narrower eligibility criteria can be applied for particular candidate therapies within a trial to address specific contraindications, provided that any subgroup of the trial population qualifies for at least two arms so that randomisation can be used. Although best practices might change over time, platform trials can be designed to permit updates to the standard of care across treatment groups as the evidence evolves, and concurrent controls can be used in analyses to reflect the contemporary standard of care. Having multiple interventions in the trial poses challenges for safety monitoring and blinding, which can be mitigated by separating group assignment into a two-step process: unblinded randomisation to one potential intervention in the trial while masking assignment to active or placebo groups for that intervention. This approach preserves the advantages of blinding and allows multiple routes of drug administration to be included in the same master protocol. In some settings such as pragmatic trials, placebo might not be available or masking might not be possible for other reasons; best practices for preserving study integrity with unblinded treatment administration should still be followed. Sustained funding sources are needed to develop and maintain the infrastructure that is necessary to test a series of therapies at multiple sites. These challenges will need to be considered carefully in designing ARDS platform trials.

Endpoint selection
There are no well validated, patient-centred, disease-specific endpoints for ARDS. Mortality remains the most widely accepted endpoint for ARDS trials. Nevertheless, ARDS-attributable risk of death can differ considerably between patients and cohorts, an important factor influencing statistical power of trials and a potential cause of heterogeneity of treatment effect. No validated surrogate endpoint exists for ARDS mortality. Physiological endpoints, such as change in oxygenation or extravascular lung water, have not been consistently correlated with treatment effects on mortality and in some instances have suggested improved lung function for interventions subsequently found to increase mortality.

Endpoints other than mortality must address death as a competing risk. Ventilator-free days, a composite outcome that includes death and time from successful ventilator weaning to day 28, is not clearly patient centred and as originally defined equates death as equivalent to 28 days of ventilator dependence. To overcome this limitation, ventilator-free survival is a ranked composite score that compares each patient to all other patients in the study, first by vital status and then, only if both patients in a pair survive, by duration of ventilation. As an alternative, WHO proposed an ordinal scale for
Panel 3: Unresolved key questions for advancing precision medicine in ARDS

- What is the biological overlap between critical illness syndromes such as ARDS and sepsis? When should trials consider enrolling patients according to treatable traits that might span conventional diagnostic labels?
- Which patients are at highest risk for ARDS-attributable mortality, and how can they be identified early in the disease course? Should ARDS trials restrict enrolment to this subset of patients, and how would this affect the generalisability of results in terms of benefits and risks?
- What are useful surrogate outcome measures for ARDS clinical trials? Should clinical trials focus primarily on mortality or is there a more ARDS-specific endpoint with favourable performance characteristics?
- What is the stability of molecular subphenotypes of ARDS over time, and what implications does this stability have for clinical trials?
- What are the key mechanistic drivers (nodes) of molecular subphenotypes of ARDS?
- What are the contributions of variations in genetics and the microbiome?
- What assays should be prioritised for development into rapid diagnostics to enable future trials?
- What are the highest priority therapeutic candidates for testing in precision medicine trials?

Search strategy and selection criteria

We searched PubMed for articles published from database inception to Nov 1, 2020, using the terms “acute respiratory distress syndrome” or “acute lung injury” and “precision medicine” without language restrictions to identify potentially relevant publications. This search was supplemented by the authors’ own literature searches for their pre-workshop topic-specific summaries, which were written to inform discussions on the themes developed in the workshop and in this manuscript. When publications with overlapping content were identified, the references deemed most immediately relevant were included in the final citation list.

COVID-19 trials that incorporates post-extubation level of respiratory support, and some COVID-19 trials have used this scale to evaluate time to recovery as the main trial endpoint.

Long-term follow-up of ARDS survivors has revealed protracted physical, cognitive, and psychological morbidity, and core outcome measures have been proposed for studies of ARDS survivorship. These core outcomes were developed with input from both survivors of ARDS and their families, emphasise many domains contributing to quality of life, and have been integrated into recent trials. Measuring long-term outcomes is essential to understand the totality of patient-centred treatment effects and the spectrum of survivorship within trials. To be suitable for consideration as a primary endpoint for trials, long-term outcomes must account for the competing risk of death to retain face validity and should be supported by evidence suggesting that these measures can be modified by candidate interventions. The preferred outcome, or family of outcomes, for a given trial might depend on the particular population enrolled and intervention that is studied.

Response indicators

Response indicators are early markers of therapeutic target engagement and are distinct from trial endpoints. For example, in oncology, tumour shrinkage or change in tumour biomarker serum levels after initial cycles of chemotherapy could indicate drug target engagement but would not themselves be appropriate trial outcomes until unambiguously linked to patient-centred clinical outcomes. The ideal response indicator would be useful early in the treatment course to determine whether the prescribed therapy is likely to be effective for a given patient and thus should be continued in that patient. Response indicators might also be useful to guide dose adjustment for titratable therapies. No universal response indicator exists for ARDS, but intervention-specific response indicators could be tested within trials as they become available.

Candidate interventions

Discovery and development of new drugs is expensive and time-consuming, costing US$1–2 billion or more, and often taking 10–15 years from discovery to regulatory approval. Testing approved drugs and known drug candidates for new indications takes advantage of their established mechanisms of action and known safety and pharmacokinetic profiles, reducing cost and time. Large databases of clinically approved drugs have been established, such as the National Center for Advancing Translational Sciences (NCATS) Pharmaceutical Collection, and efforts have been made to profile these moieties for activity over a wide range of pathways and disease models. Cross-referencing known drug activities with appropriate targets in ARDS could lead to development of new drugs at reduced cost and time than required for entirely new compounds, an approach embraced by the scientific community to fight the COVID-19 pandemic.

Repurposing of drugs has had successes in other pulmonary diseases. For example, sildenafil, a phosphodiesterase type 5 inhibitor, was initially developed as a possible therapy for coronary artery disease, before being tested and approved for erectile dysfunction and subsequently for pulmonary arterial hypertension. In asthma, recognition of the T-helper-2 paradigm provided a pathophysiological basis for new drugs that inhibit the interleukin 5 (IL-5) and IL-4/IL-13 pathways, which were originally developed for several eosinophilic diseases (anti-IL-5 agents) and atopic dermatitis (tralokinumab, an anti-IL-4/IL-13 therapy). Drug discovery efforts for ARDS also should consider whether several different pathways need to be
manipulated simultaneously to maximise therapeutic effect, as in cancer chemotherapy.

Conclusions and future directions

Therapeutic discovery for ARDS requires new approaches that bring phenotypic and biological heterogeneity to the fore to match candidate therapies to resulting subgroups. Concerted coordination across the research community will be required to achieve this precision medicine vision. Mechanistic preclinical studies, translational clinical cohort studies, and randomised trials fulfil intertwined roles in understanding mechanisms, prognostic relevance, and therapeutic implications of ARDS heterogeneity. Clinical trials must be reimagined to build the discovery pipeline, leveraging the efficiencies of novel designs to test multiple candidate therapies simultaneously and match them to the right patient subgroups. Partnership between academia, industry, regulatory agencies, sponsors, and patients must be nurtured.

Several areas of uncertainty remain regarding the best path forward to advance precision medicine in ARDS. These key unanswered questions form a foundation for future areas of research (panel 3). With broad recognition that ARDS heterogeneity modifies therapeutic efficacy, the impetus to develop and advance precision medicine strategies is clear. Deeper understanding of key nodes in mechanistic pathways established through rigorous preclinical studies and well designed observational cohorts, paired with innovative trials designed to test for sources of heterogeneity that influence treatment responsiveness, will accelerate discovery of targeted therapies to reduce morbidity and mortality for patients with ARDS.

Contributors

BTT, NRA, and CSC had the initial idea for the workshop, planned the workshop, invited the participants, and chaired the workshop. All authors attended and contributed intellectual content to the workshop. JRB, BTT, NRA, and CSC wrote the first draft of the manuscript. All authors critically reviewed the manuscript and approved the final version.

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