INTRODUCTION

Machine learning for healthcare (MLHC) is at the juncture of leaping from the pages of journals and conference proceedings to clinical implementation at the bedside. Succeeding in this endeavour requires the synthesis of insights from both the machine learning and healthcare domains, in order to ensure that the unique characteristics of MLHC are leveraged to maximise benefits and minimise risks. An important part of this effort is establishing and formalising processes and procedures for characterising these tools and assessing their performance. Meaningful progress in this direction can be found in recently developed guidelines for the development of MLHC models, guidelines for the design and reporting of MLHC clinical trials, and protocols for the regulatory assessment of MLHC tools.

But while such guidelines and protocols engage extensively with relevant technical considerations, engagement with issues of fairness, bias and unintended disparate impact is lacking. Such issues have taken a place of prominence in the broader ML community, with recent work highlighting issues such as racial disparities in the accuracy of facial recognition and gender classification software, gender bias in the output of natural language processing models and racial bias in algorithms for bail and criminal sentencing. MLHC is not immune to these concerns, as seen in disparate outcomes from algorithms for allocating healthcare resources, bias in language models developed on clinical notes and melanoma detection models developed primarily on images of light-coloured skin. Within this paper, we will examine the inclusion of fairness in recent guidelines for MLHC model reporting, clinical trials and regulatory approval. We highlight opportunities to ensure that fairness is made fundamental to MLHC, and examine ways how this can be operationalised for the MLHC context.

FAIRNESS AS AN AFTERTHOUGHT?

Model development and trial reporting guidelines

Several recent documents have attempted, with varying degrees of practical implication, to enumerate guiding principles for MLHC. Broadly, these documents do an excellent job of highlighting artificial intelligence (AI)-specific technical and operational concerns, such as how to handle human-AI interaction, or how to account for model performance errors. Yet as outlined in table 1, references to fairness are either conspicuously absent, made merely in passing, or relegated to supplemental discussion.

Notable examples are the recent the Standard Protocol Items: Recommendations for Interventional Trials-AI (SPIRIT-AI) and Consolidated Standards of Reporting Trials-AI (CONSORT-AI) extensions, which expand prominent guidelines for the design and reporting of AI clinical trials to include concerns relevant to AI. While the latter states in the discussion that ‘investigators should also be encouraged to explore differences in performance and error rates across population subgroups’, there is no more formal inclusion of the concept into the guideline itself. Similarly, the announcement papers for the upcoming Transparent Reporting of a multivariable prediction model for Individual Prognosis or Diagnosis-Machine Learning (TRIPOD-ML) andStandards for Reporting of Diagnostic Accuracy Studies AI Extension (STARD-AI) guidelines for model reporting do not allude to these issues (though we wait in anticipation for their potential inclusion in the final versions of these guidelines). While recently published guidelines from the editors of journals and conference proceedings
Respiratory, sleep and critical care medicine journals engage with the concept in an exemplary fashion, the depth of their discussion is relegated to a supplementary segment of the paper.1

Regulatory guidance

Broadly, the engagement of prominent regulatory bodies with MLHC remains at a preliminary stage, and engagement with fairness tends to be either minimal or vague. The Food and Drug Administration in the USA has made significant strides towards modernisation of its framework for the approval and regulation of software-based medical interventions, including MLHC tools.5 Their documents engage broadly with technical concerns, and criteria for effective clinical evaluation, but entirely lack discussion of fairness or the relationship between these tools and the broader health equity context.20 The Canadian Agency for Drugs and Technologies in Health has explicitly highlighted the need for fairness. However, moving from vague commitments of fairness to practical and effective guidance is far from a trivial task. As work in the machine learning community has demonstrated, fairness has multiple definitions which can occasionally be incompatible,7 and bias can arise from a complex range of sources.30 Operationalisation of fairness must be context-specific, and embeds the relevant values in a field. We call for concerted effort from the MLHC community, and in particular the groups responsible for the development and propagation of guidelines, to affirm a commitment to fairness in an explicit and operationalised fashion. Similarly, we call on the various regulatory agencies to establish clear minimum standards for AI fairness. In box 1, we highlight a non-exhaustive series of recommendations that are likely to be beneficial as the MLHC community engages in this endeavour.

## OPERATIONALISING FAIRNESS IN MLHC PRACTICE

If fairness is an afterthought in the design and reporting of MLHC papers and trials, as well as regulatory processes, it is likely to remain an afterthought in the development and implementation of MLHC tools. If MLHC is going to prove effective for—and be trusted by—a diverse range of patients, fairness cannot be a post-hoc and after-the-fact consideration. Nor is it sufficient for fairness to be a vague abstraction of academic importance but ineffectual consequence. The present moment affords a tremendous opportunity to define MLHC such that fairness is integral, and to ensure that this commitment is reflected in model reporting guidelines, clinical trial guidelines and regulatory approaches.

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### Recommendations for operationalising fairness

- Engage members of the public and in particular members of marginalised communities in the process of determining acceptable fairness standards.
- Collect necessary data on vulnerable protected groups in order to perform audits of model function (eg, on race, gender).
- Analyse and report model performance for different intersectional subpopulations at risk of unfair outcomes.
- Establish target thresholds and maximum disparities for model function between groups.
- Be transparent regarding the specific definitions of fairness that are used in the evaluation of a machine learning for healthcare (MLHC) model.
- Explicitly evaluate for disparate treatment and disparate impact in MLHC clinical trials.
- Commit to postmarketing surveillance to assess the ongoing real-world impact of MLHC models.
CONCLUSION
Values are embedded throughout the MLHC pipeline, from the design of models, to the execution and reporting of trials, to the regulatory approval process. Guidelines hold significant power in defining what is worthy of emphasis. While fairness is essential to the impact and consequences of MLHC tools, the concept is often conspicuously absent or ineffectually vague in emerging guidelines. The field of machine MLHC has the opportunity at this juncture to render fairness integral to the identity field. We call on the MLHC community to commit to the project of operationalising fairness, and to emphasise fairness as a requirement in practice.

Author affiliations
1Department of Radiology & Imaging Sciences, Emory University, Atlanta, Georgia, USA
2Fogarty International Center, National Institutes of Health (NIH), Bethesda, Maryland, USA
3Temerty Faculty of Medicine, University of Toronto, Toronto, Ontario, Canada
4Laboratory for Computational Physiology, Harvard-MIT Division of Health Sciences and Technology, Cambridge, Massachusetts, USA
5Division of Pulmonary Critical Care and Sleep Medicine, Beth Israel Deaconess Medical Center, Boston, Massachusetts, USA
6Department of Biostatistics, Harvrd T.H. Chan School of Public Health, Boston, Massachusetts, USA
7Department of Computer Science, University of Toronto, Toronto, Ontario, Canada
8Department of Medicine, University of Toronto, Toronto, Ontario, Canada
9Vector Institute for Artificial Intelligence, Toronto, Ontario, Canada

Twitter Judy Wawira Gichoya @judywawira, Lian G McCoy @liamgmcocoy and Leo Anthony Cell @MITCriticalData

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ORCID iD Leo Anthony Cell http://orcid.org/0000-0001-6712-6626

REFERENCES