Responsible AI Checklist (RAIC) for Health AI *Responsible AI Checkpoint Three Large Scale and Longer-Term Impacts*

Coalition for Health Al

June 26, 2024

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Checklist Document Versions

As this checklist is passed back and forth between different Reporters and Reviewers, Table 1 will help track versions of the document. Italicized information in the checklist serve as examples and should be replaced during use.

Table 1. Checklist Document Versions

			Versio	ns		
Document Version	Application & Model Version	Content Description	Reporter or Reviewer Name	Contact Information and Role	Organization	Date
<1.0>	Exacerbation Risk version 1.0	<documentation and="" evidence<br="">provided by implementer and development teams/specific departments from Mayo Clinic></documentation>	<name></name>	<reporter 1=""> E-mail: Phone: Title:</reporter>	<mayo clinic=""></mayo>	<may 1,="" 2024=""></may>
<2.0>	Exacerbation Risk version 1.0	<documentation and="" evidence<br="">related to use and human- factors considerations provided by external consultant at ideas42></documentation>	<name></name>	<reporter 2=""> Email: Phone: Title:</reporter>	<ideas42></ideas42>	<may 2024="" 5,=""></may>
<3.0>	Exacerbation Risk version 1.0	<summary and<br="" findings="" of="">review of documentation and evidence provided by development and implementer teams at Mayo and consultants from ideas42></summary>	<name></name>	<reviewer 1=""> Email: Phone: Title</reviewer>	<mayo clinic=""></mayo>	<may 2024="" 7,=""></may>

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1 Purpose and Use

1.1. Purpose

The Responsible AI Checklist (RAIC) is intended to guide the development and evaluation of a complete **AI solution** and **system** against CHAI content for trustworthy AI¹. This tool is intended first for self-reporting and self-review, as well as a tool for self-reporting for independent review. The goal of the RAIC is to ensure that AI solutions and systems fulfill all five key, principle-based areas for trustworthy AI: 1. Usefulness, Usability, and Efficacy; 2. Fairness; 3. Safety; 4. Transparency and Intelligibility; 5. Privacy and Security. In alignment with these areas, the RAIC translates best practice considerations (detailed in the Responsible AI Guide) that meet core ethical and quality principles into detailed yes/no questions, or evaluation criteria, to determine whether best practices are met (see accompanying Responsible AI Guide). The relationship between evaluation criteria and their original considerations, as well as criteria that have been combined across multiple areas and considerations are mapped in a Traceability Matrix located in the Appendix of this document. The RAIC encourages a holistic understanding of AI solutions in context, encompassing the interplay of human-factors, data, algorithms, infrastructure, and real-world workflows, facilitating conversations across developer and implementer teams, and As a self-review tool for developer and implementation teams, this iteration of the RAIC also serves as a starting point for facilitating conversation and alignment on best practices across the full AI lifecycle.

A secondary purpose of this version of the tool is to guide an understanding of the state of trustworthy AI in healthcare and the needs of representative stakeholders and healthcare organizations by stress-testing the checklist in the real-world. Specifically, utilization of this tool and feedback on existing end-to-end capabilities and practices will aid both in improving and iterating on the RAIC and its subsequent versions, as well as an understanding of the challenges that may influence the feasibility of best practices.

1.2. Intended Users

Intended users of the RAIC are developer and implementation teams within or outside of health systems with accountable Reporters from teams providing documentation and summaries for executive review. Multiple stakeholders (see section 3.3 in the Appendix and section 3.2 in the Responsible AI Guide) may be involved in the selection, procurement, development, and deployment process of an AI solution. This iteration of the RAIC does not prescribe roles and responsibilities, however it outlines usage for those completing and reviewing the document (see Responsible AI Guide, pg. 2 for further details on this and plans

¹ The RAIC was developed by forming expert workgroups for each principle area. Workgroups conducted a full landscape analysis and synthesized findings into a series of considerations and criteria for each lifecycle stage for their specific principle-based focus areas. These considerations and criteria were then compiled into a survey sent out to the broader CHAI community to gain multi-stakeholder feedback and ratings as part of a modified Delphi-process to gain consensus across multiple stakeholders. Results were then reviewed during the Fall convening and discussed further. Considerations that were rated as "Extremely Important" by at least 50% of the respondents, and/or were deemed extremely important following the second round of discussions, were included in this version of the Responsible AI Guide and Checklist. Additional considerations and criteria that were rated as either "Extremely Important" or "Very important" by at least 65% of survey respondents are included in the Traceability Matrix but not in this version of the Responsible AI Guide or Checklist.

for future versions). Developer and implementer teams may be entirely or in part internal or external to the healthcare organization looking to develop, procure, or implement an AI solution. As such, this tool may also be used as part of a collaborative process across developer and implementer teams to foster trust and alignment on best practices.

This checklist is most appropriate for products or devices that are themselves AI software (predictive or generative) or those that are AI assisted/AI enabled. At this point in time, AI tools often used in drug discovery and development (e.g. target selection or antibody design) in the pharmaceutical industry fall outside the targeted scope of the RAIC.

AI software examples: Payer/provider risk stratification or prediction, diagnostic algorithms, automated EHR coding, provider decision or administrative support, patient decision support, patient or provider facing chatbot used for education or assistance

AI assisted/AI enabled examples: AI enabled medical devices, AI assisted surgical robots, radiological technologies that are AI assisted or AI enabled for clinical (diagnostic/risk prediction) or nonclinical purposes (automated image quality enhancement.)

The **Reporter** is the individual tasked to gather responses and documentation from appropriate "**Providers of Evidence**," or experts in the areas pertaining to RAIC items. The **Reviewer** can either be an internal executive responsible for checking the completeness and appropriateness of the explanations and documentation to guide the development, procurement, and/or implementation of an AI solution based on best practices, or an external independent Reviewer who will evaluate the overall AI system for alignment with best practices. Note that there may be multiple Reporters, Providers of Evidence, and Reviewers. For smaller organizations or health systems there may be fewer stakeholders available, or the need to consult with external experts to ensure best practices in specific areas. We do not expect that all best practices are feasible at this point and aim to further understand feasibility as they are stress-tested in the real world. Examples of user personas and scenarios are provided in the Appendix (section 3.4).

1.3. Usage

Usage of the RAIC is guided by the AI Lifecycle (Figure 1). The AI Lifecycle can be an iterative and non-linear/agile outline of the processes required for effective and trustworthy design, development, and use of a health AI system from end-to-end. To facilitate the agile process, we have identified a **planning checkpoint** and several **responsible AI checkpoints** that aim to help teams ensure that the necessary steps have been taken prior to moving a tool into real-world use. The four checkpoints are summarized below. Examples of user personas and scenarios are provided in the Appendix (section 3.4).

- 1. The **planning checkpoint** follows Stage 1, where both developer and implementer teams (independently or together) are asked to define the specific problem and plan adequately for a potential AI solution. This checkpoint primarily helps teams:
 - a. Appropriately consider the risks, benefits, costs, and needs for an AI solution both at the clinical and population levels
 - b. Consider the risks, benefits, costs, and needs around purchasing or developing an AI solution in house
 - c. Gain multi-stakeholder insights to help guide human-centered AI solution design, development (or purchasing) and downstream needs to maximize real-world effectiveness and trust
- 2. **Responsible AI checkpoint one** appears when progressing from iterations through design, development, and assessment processes, to the small-scale pilot phase. The goal of this checkpoint is to address readiness for piloting and to prepare for real-world risks and needs. Any updates to clinical and population risk summaries should be made based on new insights from the design, development, and silent-evaluation process. An important note is that this checkpoint is not only meant for developer organizations. There are items that assess for readiness for the implementer/purchasing organization, items to guide conversations around responsibilities between developer and implementer organizations, items that speak to the larger AI system design and development (e.g. safety, privacy, security, and

monitoring planning), and items that a purchasing/implementing organization may use to understand vendor best practices. An organization or health system acquiring or purchasing an AI solution may choose to use this checkpoint as part of their procurement process. For example, they may require developer organizations to provide relevant evidence in support of best practices during design, development, and evaluation to help make purchasing decisions to foster transparency. It is also recommended that purchasing/implementing organizations review the planning checkpoint items alongside the developer organization to ensure appropriate planning, risk determination, and usability for the broader AI system (beyond the AI solution alone).

- 3. **Responsible AI checkpoint two** appears when progressing from piloting to at-scale deployment of the AI system, which requires evaluation of readiness and preparation for the broader needs and wider scope of risk. Any updates to clinical and population risk summaries should be made based on new insights from initial real-world piloting.
- 4. **Responsible AI checkpoint three** appears following full scale deployment to evaluate for longer-term readiness for monitoring, managing, and updating the AI system. This checkpoint is repeated throughout regular monitoring of the AI solution, at appropriately timed intervals depending on the use case, and as dictated by the developer and/or implementer organization. As in previous checkpoints, updates should be made to clinical and population risk summaries based on insights gained from regular monitoring of AI solutions and systems.

Within each checkpoint checklist, relevant evaluation criteria are listed and given an identifier. The color coded Evaluation Criteria Identifier (EC Identifier) links each criterion to the original consideration as defined within principle area workgroups (see Traceability Matrix in the Appendix 3.1; See Section 1.5 for further details.)

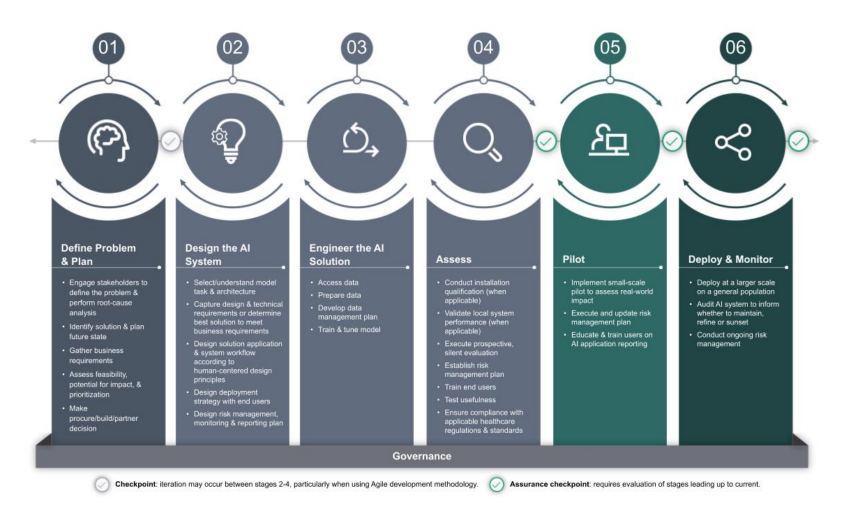


Figure 1: The CHAI AI Lifecycle Framework. Derived from CHAI Responsible AI Guide. The gray checkmark represents the Planning Checkpoint, while the green checkmarks correspond to Responsible AI Checkpoints 1-3.

1.4 How to complete this checklist

1.4.1 General

Who Should Complete This Checklist?

Each checkpoint checklist should first be completed by at least one **Reporter.** While there may be multiple stakeholders involved in sharing evidence necessary to respond to criteria, the Reporter is the individual responsible for requesting this information (if available), making sure available evidence is clearly documented for relevant evaluation criteria in the checklist, and indexing it in a centralized place for ease of Reviewer access. They will also provide a summary at the end of each checkpoint that provides reviewers with a broad overview of the potential or observed benefits, costs, risks, and/or adverse events associated with that checkpoint. Example roles, professions, and representative organizations are shown in Table 3.3 in the Appendix and described in more detail in the CHAI Responsible AI Guide.

Reporters will then pass the checklist off to at least one **Reviewer** who is internal to either developer and/or implementer organizations (such as an area specific executive). Ideally, organizations will also pursue independent and external third-party review. The Reviewer will go over the responses to evaluation criteria and evidence, and indicate whether best practices for each criteria have been met. They will also provide a summary of findings based on the available evidence and any observed gaps. This feedback can be used to improve processes, help guide teams on next steps, or help build/design solutions to fill gaps in best practices.

For **Responsible AI Checkpoints 1-3** the following steps are required.

Reporter Responsibilities for Completion (Responsible AI Checkpoints 1-3)

- 1. All Reporter required sections of the checklist or summaries are denoted with dark blue coloring.
- 2. Provide existing (from prior checkpoints) and updated clinical risk classification and Population Impact information in the "Clinical Risk and Population Impact Summaries" table at the start of each Checkpoint (Review Clinical Risk and Population Impact Tools in sections 1.4.2 and 1.4.3 respectively for any necessary updates).
- 3. The Reporter will then complete the relevant Responsible AI Checkpoint Checklist providing a brief explanation and document code in the "Evidence and Explanation & Metadata/Documentation Code" column of the checklist, with supporting evidence indexed within the "Evidence & Explanation Metadata Table" (see Section 2.5 for further instructions and Table).
- 4. The Reporter will complete the "Executive Summary of Anticipated and Observed Benefits, Risks, and Limitations" section (Section 2.3) for the relevant Responsible AI Checkpoint.
- 5. Reporter responsibilities for each Responsible AI Checkpoint Checklist will end by updating the document version table (Page 2) and up-versioning the document header, prior to sending the checklist and associated evidence to the appropriate Reviewer.

Reviewer Responsibilities for Completion (Responsible AI Checkpoints 1-3)

1. All Reviewer required sections of the checklist or summaries are denoted with light blue coloring.

- 2. The Reviewer will go through information provided in the checklist by the Reporter along with accompanying documentation listed in Evidence and Explanation Metadata table.
- 3. Reviewers will then complete the Summary of Findings table (Section 2.4), summarizing findings provided in the checklist by the Reporter in the context of anticipated and observed benefits, risks, and limitations of the AI solution.
- 4. Reviewers will then update the document version table on Page 2 and up-version the document header.

Example Reporter Role Responses

EC Identifier	Evaluation Criteria	Evidence and Explanation Metadata/Document Code	N/A or Cannot Complete (CC): Describe in Comment	Reporter Initials & Date	Evidence & Explanations (Yes/No/Partial/NA)	Limitations or Adverse Outcomes	Criteria Met (Yes/No/Partial/NA)	Reviewer Initials & Date
LS2.F.C1.EC2	Will the real-world/clinical outcome measure be available for evaluation within an adequate time frame and in a manner that accurately represents the target population?	Evidence and explanation: Real-world retrospective data was used for evaluation of model performance and comparable to target population. Metadata/Document Location: <insert link<br="">to bias assessment document and relevant data showing summary of real-world retrospective data population descriptives and demographics and comparison to target population descriptives and demographics.)</insert>		M.G. 05/06/2024				
LS2.F.C1.EC3	Will real-world/clinical outcomes be systematically compared for across all relevant socio-demographic subgroups, ensuring fairness and addressing potential bias?	Evidence and explanation: Overall ER admission rates are lower following use of the AI solution. Clinical outcomes are similar for all subgroups except for Black Patients, who show higher ER admissions following discharge at the same population level risk threshold compared to the sample majority group and compared to the population mean. Metadata/Document Location: <insert link<="" td=""><td></td><td>M.G. 05/06/2024</td><td></td><td></td><td></td><td></td></insert>		M.G. 05/06/2024				

to bias assessment document and relevant data showing likelihood of ER admissions following discharge (as measure of clinical outcomes that AI solution aimed to impact)					
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Example Reviewer Responses

EC Identifier	Evaluation Criteria	Evidence and Explanation Metadata/Document Code	N/A or Cannot Complete (CC): Describe in Comment	Reporter Initials & Date	Evidence & Explanations (Yes/No/Partial/NA)	Limitations or Adverse Outcomes	Criteria Met (Yes/No/Partial/NA)	Reviewer Initials & Date
LS2.F.C1.EC2	Will the real-world/clinical outcome measure be available for evaluation	Evidence and explanation: Real- world retrospective data for ER admission rates are available and will be used for evaluation of model's impact on clinical outcomes. Data is comparable to target population. Metadata/Document Location: <insert assessment<br="" bias="" link="" to="">document and relevant data showing summary of real-world retrospective data population descriptives for measure and demographics and comparison to target population descriptives for measure and demographics of sample.)</insert>		M.G. 05/06/2024	Yes	No, None stated	Partial, Provide justification for why this clinical outcome was selected.	N.E. 05/10/2024
LS2.F.C1.EC3	Will real-world/clinical outcomes be systematically compared for	Evidence and explanation: Overall ER admission rates are lower following use of the AI solution. Clinical outcomes are similar for all subgroups except for Black Patients, who show higher ER admissions following discharge at the same population level risk threshold compared to the sample majority group and compared to the population mean.		M.G. 05/06/2024	Partial, provide information on what threshold was selected and why.	Yes, Black patients have poorer outcomes at the chosen threshold	Partial	N.E. 05/10/2024

Metadata/Document Location:			
<insert assessment<="" bias="" link="" td="" to=""><td></td><td></td><td></td></insert>			
document and relevant data			
showing likelihood of ER			
admissions following discharge			
(as measure of clinical outcomes			
that AI solution aimed to impact)			

1.4.2 Clinical Risk Evaluation

Risk should be assessed from both the **clinical** and **population** perspective. For **clinical risk**, we adopt the International Medical Device Forum's (IMDRF's) categorization system for assessment of clinical risk (See Table 2). This should be done by a licensed clinician based on the FDA IMDRF guidance.

Table 2. Assessment criteria for clinical risk level. Levels are described in detail in <u>"Software as a Medical Device": Possible Framework for Risk Categorization and</u> <u>Corresponding Considerations</u>" by IMDRF Software as a Medical Device (SaMD) Working Group (2014).

Clinical Risk Classification									
	Signifi	cance of information provided to healthc	are decision						
State of Healthcare situation or condition	Treat or diagnosis	Treat or diagnosis Drive clinical management							
Non-Serious	Ш	Ι	Ι						
Serious	III	П	Ι						
Critical	IV	III	II						

Clinical risk classification and summaries should be provided in Section 2.1, Table 3. Clinical Risk and Population Impact Evaluation Summaries

1.4.3 Population Impact Evaluation Tool:

Population risk refers to how systemic, individual, and group-level tendencies when combined with decision-making demands across the AI lifecycle, can impact health and well-being for entire subgroups and over longer periods.

While it is common to refer to systemic, individual, and group-level tendencies as "biases"—it is important to note that they are often the result of things like:

- Historical Norms/policies
- Current Societal Norms/policies
- Scope of Skills/Responsibilities
- Natural limitations/variability in cognitive resources/awareness
- The burden of increasing clinical/administrative demands
- Role specialization (and therefore less insight into other roles or expertise)

It is normal for us to:

- Not have all knowledge about a topic
- To want to use data that is readily available or easily accessible
- To be focused on our role-specific responsibilities and not aware of the roles/responsibilities of others
- To focus on resolving a specific problem (e.g. sepsis prediction), without considering how it might unintentionally harm a subgroup of individuals due to bias in data/measurement
- To want to follow shortcuts

The following questions will help stakeholders involved in purchasing or developing an AI solution, together with other relevant stakeholders (see Section 3.3 in the Appendix) to evaluate population risk and impact in a way that will improve current practices and minimize population-level harm across several domains. This will allow teams to leverage the power of health AI to positively impact patients and providers and reduce healthcare gaps and inequities, rather than perpetuate or prolong them. These questions are best explored with patient advocacy/population health and medical area experts present or consulted. Given that bias in AI is unavoidable, this tool will also help organizations evaluate and prioritize bias mitigation efforts towards algorithms with greater risk and/or those that may be impacted by ethical/legal guidelines. Using this tool aims to improve current practices and minimize population-level harm. (Tool adapted to health-specific context in part from ethicstoolkit.ai)

Identify who will be impacted by the AI system:

Primary Impacted: Who or what may be or is directly impacted based on the objectives of the AI system? (e.g. patients, family caretakers, physicians, nursing, organization, business operations, etc.)

Secondary: Who or what may be or is impacted downstream based on those primarily impacted? (e.g. if physicians and their clinical workflows are primarily impacted, downstream effects may be experienced by nursing staff, or radiology technicians)

Unexpected/Unintended: Who or what may be impacted unexpectedly/unintentionally at the population or location level? Examples may include:

- Patients who do not speak English or their children
- Physicians working in community hospitals vs. academic medical centers
- Patients without insurance
- Acquired hospitals that use a different (non-integrated) electronic medical record system
- Members of a specific socio-demographic subgroup
- Individuals with visible or invisible disabilities

Select the types of impact that the AI system may have on PATIENTS and the degree, scale, and direction of impact for each type:

• Access to Health Goods/Benefits:

Algorithms that impact who, what, where, or how someone does/does not have access health goods or benefits (ability to track health status, ability to access test results, disease management, advanced care management services)

Select Degree: Minor Impact | Moderate Impact | Major Impact

Select Scale: Small Proportion | Substantial Proportion OR Primarily one or more Vulnerable Subpopulations | Nearly Every Person OR Majority of one or more Vulnerable Subpopulations Select Direction: Positive Impact | Mostly Positive Impact | Mostly Negative Impact | Negative Impact

Access to Direct Health Services/Healthcare: Algorithms that impact who or how someone does/does not have access to necessary direct health care services (transportation coordination, medicine or health service approval, preventative care appointments, specialty care services, diagnostic testing, mental health screening, etc.)
 Select Degree: Minor Impact | Moderate Impact | Major Impact
 Select Scale: Small Proportion | Substantial Proportion OR Primarily one or more Vulnerable Subpopulations | Nearly Every Person OR Majority of one or more Vulnerable Subpopulations

Select Direction: Positive Impact | Mostly Positive Impact | Mostly Negative Impact | Negative Impact

Emotional Health/Well Being: These algorithms impact the emotional health or well-being of an individual or group. (Time waiting for health services/benefits, effort required to arrange for services) Select Degree: Minor Impact | Moderate Impact | Major Impact Select Scale: Small Properties | Substantial Properties OP Primarily one or more Vulnerable Subpopulations | Nearly Every Person OP Majority of one or more Vulnerable Subpopulations

Select Scale: Small Proportion | Substantial Proportion OR Primarily one or more Vulnerable Subpopulations | Nearly Every Person OR Majority of one or more Vulnerable Subpopulations Select Direction: Positive Impact | Mostly Positive Impact | Mostly Negative Impact | Negative Impact

Life/Safety: These algorithms directly impact individual or group safety or life (e.g. diagnostic, treatment, recommended treatments)
 Select Degree: Minor Impact | Moderate Impact | Major Impact
 Select Scale: Small Proportion | Substantial Proportion OR Primarily one or more Vulnerable Subpopulations | Nearly Every Person OR Majority of one or more Vulnerable Subpopulations
 Select Direction: Positive Impact | Mostly Positive Impact | Mostly Negative Impact | Negative Impact

- Financial: These algorithms impact the costs associated with healthcare for individuals, groups, or in specific areas. (e.g. health plan premiums, cost of care)
 Select Degree: Minor Impact | Moderate Impact | Major Impact
 Select Scale: Small Proportion | Substantial Proportion OR Primarily one or more Vulnerable Subpopulations | Nearly Every Person OR Majority of one or more Vulnerable Subpopulations
 Select Direction: Positive Impact | Mostly Positive Impact | Mostly Negative Impact | Negative Impact
- Privacy: These algorithms impact the privacy of personal health information for an individual or group.
 Select Degree: Minor Impact | Moderate Impact | Major Impact
 Select Scale: Small Proportion | Substantial Proportion OR Primarily one or more Vulnerable Subpopulations | Nearly Every Person OR Majority of one or more Vulnerable Subpopulations
 Select Direction: Positive Impact | Mostly Positive Impact | Mostly Negative Impact | Negative Impact

- Trust: These algorithms impact the trust that an individual or group may have in the healthcare system, clinician(s), or other healthcare professional.
 Select Degree: Minor Impact | Moderate Impact | Major Impact
 Select Scale: Small Proportion | Substantial Proportion OR Primarily one or more Vulnerable Subpopulations | Nearly Every Person OR Majority of one or more Vulnerable Subpopulations
 Select Direction: Positive Impact | Mostly Positive Impact | Mostly Negative Impact | Negative Impact
- Freedom/Agency/Rights: These algorithms impact an individual's freedom/agency/rights as it pertains to their healthcare or health information.
 Select Degree: Minor Impact | Moderate Impact | Major Impact
 Select Scale: Small Proportion | Substantial Proportion OR Primarily one or more Vulnerable Subpopulations | Nearly Every Person OR Majority of one or more Vulnerable Subpopulations
 Select Direction: Positive Impact | Mostly Positive Impact | Mostly Negative Impact | Negative Impact

Is it possible that the degree or scale of impact could vary by context (population subgroup or location implemented).

- No likelihood of systematic variation in scope of impact by context
- Small likelihood of systematic variation in scope of impact by context, but variability is due to known and validated clinical or social needs
- Small likelihood of systematic variation in scope of impact by context
- Medium likelihood of systematic variation in scope of impact by context, but variability is due to known and validated clinical/social needs
- Medium likelihood of systematic variation in scope of impact by context
- High likelihood of variation in scope of impact by context, but variability is due to known and validated clinical/social needs
- High likelihood of variation in scope of impact by context

1.5 How to interpret this checklist

The checklist is designed not as a binary pass-fail assessment, but rather as a comprehensive tool to evaluate the risk-benefit profile of the AI solution and its associated system and to guide best practices across developer and implementer teams. Given the inherent complexity of each use case and implementation, a nuanced approach is essential. The checklist aims to facilitate transparency and furnish reviewers with substantial evidence, empowering relevant parties to make informed go/no-go decisions. Furthermore, it underscores the importance of additional measures that may be undertaken by the implementation or developer organization. These measures are crucial for preventing and mitigating adverse outcomes, as well as ensuring that the AI solution is employed judiciously in contexts where its limitations are acknowledged and respected.

Throughout the checklist, each evaluation criteria has received one or more coding tags in the left-hand column (example: LS1.U.C1.EC1). These identifiers are designed for traceability to the considerations in the Responsible AI Guide, and they are color-coded by principle area. Some evaluation criteria are based on considerations that space multiple principle areas or span multiple considerations within a principle area. :

• Usefulness, Usability, Efficacy: (Principle Area Denoted with U)

- Fairness: (Principle Area Denoted with F)
- Safety: (Principle Area Denoted with S)
- Transparency, Intelligibility, and Accountability: (Principle Area Denoted with T)
- Privacy and Security: (Principle Area Denoted with PS)

(example: LS1.U.C1.EC1 would denote Lifecycle Stage 1, Usefulness, Usability, and Efficacy Principle Area, Consideration 1, Evaluation Criteria 1.)

Note: once the review of the checklist is complete, we'll be creating more streamlined, sequential tags. For now, the color coding will give you what's most important, as many evaluation criteria reflect overlaps in different principle-based considerations through the lifecycle.

2 Reporting Checklist

Columns and sections to be completed by the Reporter are denoted in dark blue and by Reviewer in light blue.

2.1 Clinical Risk & Population Impact Evaluation Summary

Clinical Risk and Population Impact Evaluation tools are provided in sections 1.4.2 and 1.4.3 respectively. **Reporters** should provide a summary of clinical risk (including classification level) in Table 3 below, and a summary of population impact initially in the Planning Phase (Stage 1). If not completed during the Planning Phase **and** as insights are gained during subsequent Checkpoints, tools in sections 1.4.2 and 1.4.3 should be revisited and information in Table 3 should be updated. **Reviewers** should go over this information to gain context for the information that follows in the checklist (Section 2.3).

Table 3. Clinical Risk and Population Impact Summaries

	Clinical Risk Classification & Population Impact Summaries									
Domain		Reporter Initials and Date								
Clinical Risk Classification & Summary										
Population Impact Summary										

2.2 Checklist Stage 6: Deploy & Monitor

			Checklist: Si	tage 6 Deploy 8	Monitor				
Criterion Number	EC Identifier	Evaluation Criteria	Evidence and Explanation Metadata/Document Code	N/A or Cannot Complete (CC): Describe in comment	Reporter Initials & Date	Evidence & Explanations Provided? (Yes/No/ Partial/NA)	Benefits, Limitations or Adverse Outcomes	Criteria Met? (Yes/No/ Partial/NA)	Reviewer Initials & Date
	Responsible AI (Checkpoint 3: Large-scale and Long	er-term Impacts		1		•		
AC3.CR1	LS6.U.C3.EC2 LS6.F.C2.EC1 LS6.F.C2.EC2 LS6.F.C8.EC1 LS6.PS.C1.EC2	Has a governance plan been established to delineate accountability for monitoring AI performance and security over time, ensuring that relevant personnel are both qualified and trained to communicate incident impacts with stakeholders?							
AC3.CR2	LS6.S.C2.EC1 LS6.S.C2.EC2 LS6.S.C2.EC3 LS6.S.C4.EC1	Is there a plan for tracking and mitigating safety risks by severity and frequency, incorporating an organizational standard for "adverse events" and "serious adverse events"?							
AC3.CR3	LS6.PS.C1.EC1	Are privacy and security incident response plans established, maintained, and tested according to policies for AI systems?							
AC3.CR4	LS6.F.C4.EC3 LS6.F.C4.EC4	Will data and model security undergo regular monitoring, along							

	LS6.PS.C3.EC1	with privacy risk evaluations of the AI system environment according to an agreed-upon schedule?				
AC3.CR5	LS6.S.C1.EC7 LS6.PS.C4.EC1	Does the implementer organization routinely update Standard Operating Procedures (SOPs) for risk management to ensure consistency in decision-making for identified security, privacy, and safety risks, and are policies established for managing legal compliance in these areas?				
AC3.CR6	LS6.U.C2.EC3 LS6.S.C1.EC6 LS6.S.C11.EC2	Does the developer provide a risk management plan outlining key AI- related safety risks that have been identified and mitigated across the supply chain or in other organizations?				
AC3.CR7	LS6.S.C11.EC1 LS6.S.C11.EC3	Are there responsible AI techniques or standards in place to support the developer's supply chain risk management, including guidelines on how safety risks should be reported and managed by both developer and implementer organizations?				
AC3.CR8	LS6.S.C5.EC2 LS6.S.C5.EC3	Is an audit trail accessible to independent reviewers, such that they can identify authorized users, actions on the interface, and the				

		decision-making process based on the output of the AI solution?				
AC3.CR9	LS6.F.C6.EC1	Have potential long-term risks associated with the model's performance (that is, risks not measurable during the pilot stage but potentially arising in deployment) been identified?				
AC3.CR10	LS6.S.C1.EC1 LS6.S.C1.EC5 LS6.S.C4.EC2	Are there established processes and procedures for risk management, including reporting adverse events, safety issues, and their causes to the implementer and developer organizations (when separate), and is information shared with regulatory bodies, as appropriate, if safety concerns meet the defined risk threshold?				
AC3.CR11	LS6.PS.C1.EC3 LS6.PS.C2.EC1	Is there a process aligned with legal, contractual, and regulatory requirements, ensuring that personnel review, analyze, and report privacy and security impacts in the AI environment to external and internal stakeholders?				
AC3.CR12	LS6.S.C4.EC3	Are there established procedures for sharing recalls and corrective actions with other health system implementers?				

AC3.CR13	LS6.S.C1.EC3	Is there a mechanism to detect patterns of patient harm associated with the AI solution?				
AC3.CR14	LS6.S.C3.EC1	Are there established and routine processes to assess the clinical relevance of the model and its input variables?				
AC3.CR15	LS6.S.C3.EC2 LS6.S.C3.EC4	Are there processes in place to evaluate the availability of more effective AI methodologies and to determine when it's appropriate to transition to a newer AI solution?				
AC3.CR16	LS6.S.C3.EC3	Is there a defined process to identify incorrect or outdated knowledge or recommendations generated by the AI solution?				
AC3.CR17	LS6.S.C7.EC1 LS6.S.C7.EC2 LS6.S.C12.EC1 LS6.S.C12.EC2	Are all updates to the AI solution documented, detailing version changes and impact testing results to ensure safety and effectiveness?				
AC3.CR18	LS6.F.C2.EC3	Do accountable parties have access to relevant social, ethical, legal, human-factors, and/or clinical stakeholders or advisers in case specific problems arise, and do they have clear procedures for contacting those stakeholders?				
AC3.CR19	LS6.S.C5.EC1	Are adequate access controls implemented for the AI solution to				

		ensure appropriate user permissions?				
AC3.CR20	LS6.U.C3.EC1 LS6.S.C1.EC4	Is there an established feedback loop to consistently identify issues and defects, with a triage process for continuous improvement and monitoring?				
AC3.CR21	LS6.U.C9.EC1	Are there well-defined inclusion and exclusion criteria for the use of the AI solution?				
AC3.CR22	LS6.U.C1.EC1 LS6.T.C1.EC1	Has a preliminary study been conducted to assess both the usability and effectiveness of the AI solution when deployed in the actual environment?				
AC3.CR23	LS6.U.C2.EC1	Has a comprehensive assessment of workflow integration been conducted and documented?				
AC3.CR24	LS6.U.C2.EC2	Does the AI solution accommodate the flow of people and tasks within both physical and digital environments?				
AC3.CR25	LS6.U.C2.EC3	Are there indications that users are disregarding the AI solution in their workflow?				
AC3.CR26	LS6.U.C2.EC4	Is there evidence that users are resorting to workarounds to manage the AI solution's functionalities?				

		Is there evidence that the AI]
	LS6.U.C2.EC5	solution may impede the interaction				
AC3.CR27		between patients and clinicians?				
		Are the tasks involving the use of				
AC3.CR28	LS6.U.C7.EC1	the AI solution adequately supported in the workflow?				
AC3.CK20		Is there evidence that actions taken				
		by users after interacting with the				
	LS6.U.C7.EC2	AI solution differed from what was				
AC3.CR29		originally anticipated?				
		Is there a mechanism for reporting				
		unintended uses of the AI solution,				
	LS6.S.C6.EC1 LS6.S.C6.EC2	including periodic audits to evaluate alignment with its				
	150.5.00.102	intended purpose and identify "off-				
AC3.CR30		label" use?				
		Does the risk management plan				
		explicitly address the potential for				
	LS6.S.C8.EC1	automation bias among end users, including a method to measure and				
	LS6.S.C8.EC2	assess it (e.g., detecting incorrect				
		AI output and its impact on				
AC3.CR31		subsequent decision-making)?				
		Is a plan established to manage user				
	LS6.U.C3.EC3	disagreements with the AI output,				
	LS6.U.C4.EC1 LS6.F.C10.EC2	including a mechanism by which users can override algorithmic				
	LS6.F.C10.EC2	decisions based on clear				
AC3.CR32	L50.1.010.L03	guidelines?				

AC3.CR33	LS6.T.C2.EC2	If the end user is a clinician, is the clinician given adequate guidance on how to explain model output to patients? Do end users utilize the output				
AC3.CR34	LS6.F.C10.EC1	from the AI solution alone to make decisions, without integrating it with other information?				
AC3.CR35	LS6.T.C1.EC2	Is it possible to measure the understanding of end users and key stakeholders, looking at actions taken in response to the AI solution, then to verify the consistency of those actions against the defined limitations and intended use of the model?				
AC3.CR36	LS6.U.C8.EC1 LS6.S.C9.EC3 LS6.S.C9.EC4	Are users provided with clear, non- technical communication about the limitations and clinical implications of the AI solution, covering aspects such as error rates, contraindications, generalizability, reproducibility, and robustness, along with a plain-language explanation of how the AI model was developed, its intended purpose, and its associated safety risks (e.g. model type, dataset description, clinical study results, and representation of				

		subpopulations in training and test sets)?				
AC3.CR37	LS6.S.C9.EC1	Is there a clear explanation provided to clinicians or end users regarding the rationale behind the decisions made or suggested by the AI solution?				
AC3.CR38	LS6.S.C9.EC5	Is there a process in place to regularly update transparency information based on newly discovered limitations observed during local deployment in the implementer environment?				
AC3.CR39	LS6.U.C5.EC1 LS6.U.C5.EC2 LS6.F.C12.EC1 LS6.F.C12.EC2 LS6.F.C12.EC3	Is there a structured and usable process for gathering end user feedback, including feedback on performance, accuracy, and operational challenges, and is a review process in place to address feedback promptly so that existing issues do not escalate or cause harm?				
AC3.CR40	LS6.U.C8.EC2 LS6.F.C11.EC1 LS6.T.C3.EC1 LS6.T.C3.EC2 LS6.S.C9.EC2	Is there a clearly defined method for patients and end users to access documentation about the AI solution, including relevant safety and transparency information, tailored for various levels of expertise and health literacy?				

AC3.CR41	LS6.F.C11.EC2 LS6.T.C2.EC1	Is there a defined level of patient awareness regarding the AI solution's use in their care, with reasons identified for informing or not informing patients, considering potential risks and benefits?				
AC3.CR42	LS6.F.C11.EC3	Have human factors or behavioral science experts been consulted to determine the optimal approach for presenting information about the AI solution to patients, aiming to build trust and empower patients?				
AC3.CR43	LS6.F.C5.EC1 LS6.F.C5.EC2 LS6.F.C5.EC3	Is there a clearly defined feedback mechanism for patients or affected groups to report adverse events and express opinions on services related to the AI solution, and is that feedback mechanism unbiased to business interests, compliant with state and federal policies, and equally accessible to all relevant subgroups, with measures in place to prevent exclusion based on language barriers, ability, and so on?				
AC3.CR44	LS6.U.C6.EC1	Have the error rates and response rates improved following the implementation of the AI solution?				

AC3.CR45	LS6.U.C6.EC2 LS6.U.C6.EC3	Has the performance of the AI solution been compared to the standard of care, and is there documented evidence of the relative benefits of the AI solution?				
AC3.CR46	LS6.F.C1.EC1 LS6.F.C1.EC2 LS6.F.C7.EC1	Will model performance and parity, including inputs, outputs, and outcomes, be regularly monitored for significant drift over time across the entire population and relevant socio-demographic subgroups to mitigate unfair or systemic impacts?				
AC3.CR47	LS6.F.C4.EC1 LS6.F.C4.EC2	Will the AI system be regularly monitored to identify drift or bias, and is there a designated timescale for routinely assessing the fairness of the AI solution?				
AC3.CR48	LS6.F.C1.EC3 LS6.F.C3.EC1	Are there technically defined and justified thresholds for "significant" data drift, such that those thresholds enable early detection before potential adverse impacts on a wide scale?				
AC3.CR49	LS6.F.C3.EC3	If manual evaluation of model performance is necessary, have specific time intervals been defined, with sufficient justification provided for these intervals?				

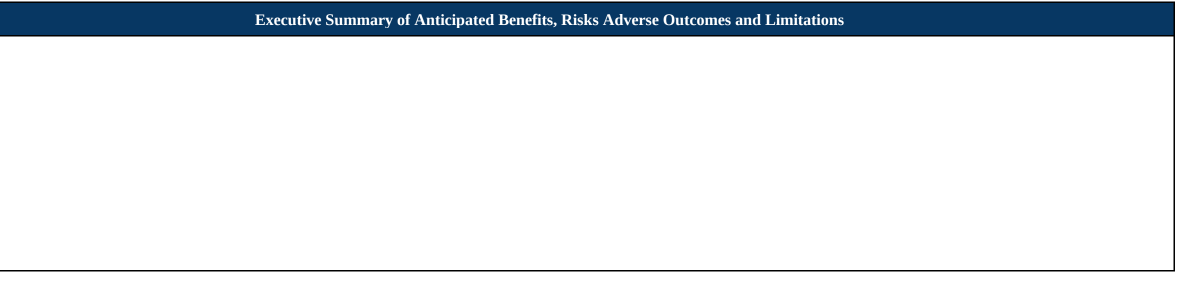
AC3.CR50	LS6.F.C3.EC2	Are there automatic and easily interpretable notifications in place to alert accountable individuals of model performance drift on an ongoing basis?				
AC3.CR51	LS6.F.C7.EC2	Are specific criteria defined for determining the significance of shifts in model performance within subgroups or between subgroups, with sufficient justification provided for the chosen criteria?				
AC3.CR52	LS6.F.C6.EC2	Have impacts of model performance drift been assessed, considering both short- and long- term effects as well as the direction and bias of these impacts?				
AC3.CR53	LS6.U.C9.EC2	Does the AI solution demonstrate varying levels of usefulness for different patient populations (e.g., pregnant women, low-risk patients, patients over age 50)?				
AC3.CR54	LS6.U.C9.EC3	Is there flexibility in the use of the model to accommodate different patient scenarios?				
AC3.CR55	LS6.F.C10.EC4	Are there differences between the pilot and deployment settings or processes that could affect how shared or automated decision- making processes influence fairness				

		and bias (e.g., time constraints,				
		population heterogeneity, patient				
		flow)?				
		Is there variation in model				
		performance based on deployment				
		site (e.g., rural vs. urban,				
		community clinic vs. academic				
	LS6.F.C9.EC1	medical center) or deployment				
		context (e.g., type of device or				
		source of device/assay for specific				
		input data, type of population most				
AC3.CR56		seen)?				
		Does the quality of data differ				
	LS6.F.C9.EC2	across various deployment sites,				
	L50.1.07.L02	and does it affect the performance				
AC3.CR57		of the model?				
		Do issues with data quality				
	LS6.F.C9.EC3	disproportionately impact				
		monitoring efforts or model				
AC3.CR58		performance in certain subgroups?				
		Is there a clear process in place for				
		monitoring and sunsetting AI				
	LS6.S.C10.EC1	solutions that are no longer				
	LS6.S.C10.EC2	supported by the developer or				
	LS6.S.C10.EC3	health system, including				
	LS6.S.C10.EC4	communication with end users				
	250.5.010.204	(contact information for assistance,				
		guidance on transitioning to a new				
AC3.CR59		solution), transition to alternative				

		solutions, handling of patient data (migration, archival, deletion, etc.)?				
AC3.CR60	LS6.S.C1.EC2 LS6.S.C2.EC4	Are Corrective and Preventative Actions (CAPAs) implemented when safety issues or poor outcomes are identified, with a clear process to determine if the AI system needs refinement or discontinuation, and are plans in place for sunsetting and safety investigation?				

2.3 Executive Summary of Anticipated Benefits, Risks, Adverse Outcomes, and Limitations

The **Reporter** should complete this section and provide an overall summary for reviewers based on responses to criteria above.



2.4 Summary of Findings

The **Reviewer** should complete this section and provide an overall summary of findings based on responses, summary, and evidence provided by the Reporter.

Reviewer Summary of Findings

2.5 Evidence & Explanation Metadata

This section should be completed by **Reporters** to list all attached evidence documents and track the source of evidence and explanations listed in the checklist. **Providers of Evidence** include any stakeholders who provided documentation and evidence to the Reporter (See Appendix Section 3.2 for a non-exhaustive list of potential stakeholders that may be involved in providing evidence for various criteria.) The first line is an illustrative example of use.

		Evidence & Explanation Metadata		
Evidence Document Code	Reporter Name and Role	Provider of Evidence Name(s), Title, Role, & Contact Information	Description	Evidence Archive Location
	<enter name,="" of<br="" reporter="" vp="">Quality></enter>	<enter data="" email@email.com="" engineer,="" name,=""></enter>	Data Management Plan	<link attachment="" document="" or<br="" to=""/> Location>

3 Appendix

3.1 Link to Traceability Matrix

https://docs.google.com/spreadsheets/d/15cJEerA861o3cSV-rzL8n0H_X-65orTBk4uuybdTByg/edit?usp=sharing

3.2 Terms Defined

Al model: A conceptual or mathematical representation of phenomena captured as a system of events, features, or processes. In computationally-based models used in AI, phenomena are often abstracted for mathematical representation, which means that characteristics that cannot be represented mathematically may not be captured in the model. Often used synonymously with "algorithm," though it may be conceptually distinct, prior to the transformation of inputs to outputs.

Al solution: A shorthand for the Al model or algorithm and required technical infrastructure (hardware, software, data warehousing, etc.).

Al system: A fully operational AI use case, including the model, technical infrastructure, and personnel in the workflow.

3.3 Representative roles in health AI industry

The roles of the developer vs. implementer organizations are unique to each AI solution and may vary throughout the lifecycle.

Stakeholder Roles	Example Stakeholder Professions	Example Representative Organizations
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Data Science Developer	Data Scientists, Data Engineers, Data Analysts & Storytellers, Machine Learning Engineers, Product Managers	Academic Medical Centers Community Health systems
Informatics and Information Technology	Biomedical Researchers and Informaticists, Software Developers, Front-End Engineers, Support Engineers, Data engineers, Quality Assurance Analysts, Security & Compliance Experts	Vendors Expert Consultants
Design and Implementation Experts	Implementation Scientists, Human Factors Experts, User Experience Designers, Patient Safety Experts, Clinicians	
End Users	Health Care Providers (e.g. Clinicians and Nurses), Insurers and Payers, Healthcare Operations Workers, Patients and Caregivers	Health Systems such as:
Health System Administration	Health Systems Leadership, Contract Administrators, Vendor Management Specialists	Academic Medical Centers Community Health Systems Integrated Healthcare Systems Primary Care Networks Urgent Care Networks
Clinical Administration	Lab Managers, Nursing Managers, Other Clinical Decision-Makers	Independent Imaging Centers Providers in Private Practice
Impacted Groups	Patients and Caregivers, Patient Advocates	Patient Advocacy Organizations Patient Advisory Boards

Ethics and Regulation & Standards Organizations	Bioethicists, IRB Analysts, IRB Members and Leaders, Lawyers and Legal Advisors, Civil Servants, NGO Decisionmakers, Policy Analysts, Regulatory Experts and Consultants	Federal Government Local Government NGOs Law Firms Standards Organizations Medical and Nursing Societies Medical Device Collaboratives, etc.
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Table 1: Stakeholder Roles, Professions, and Representative Organizations. Derived from CHAI Responsible AI Guide (Link)

3.4 Example User Personas and Scenarios for Development, Procurement, and Implementation

Example 1:

Scenario: A health system or healthcare organization (e.g. payer, EHR company) that has internal developer and implementer teams and are looking to develop a model to predict risk of post-op complications. Example Reporter(s): Chief quality officer is assigned the role of Reporter and project lead and contacts relevant stakeholders who will serve as Providers of Evidence (as appropriate) from the organization (e.g. data, informatics & security, policy/legal, human factors or social & behavioral sciences, clinical area expert, patient advocate). Ideally these individuals work together to complete the planning phase tasks and set a roadmap for the responsible AI checklist tasks and processes. When the model is ready to be piloted, teams and stakeholders will provide evidence to the Reporter for Responsible AI Checkpoint 1. Example Reviewer(s): The Vice President of Quality reviews the evidence and makes a go-no-go decision about moving the project forward to piloting. If no-go decision is made, it may be because modifications and further evidence are required, at which point the AI solution undergoes further iteration. If a go decision is made, the project moves forward to piloting, with relevant stakeholders involved in gathering evidence for the next Responsible AI Checkpoint.

The Reporter and Reviewer for subsequent checkpoints may differ as appropriate for the success of the project and as determined based on expertise required.

Example 2:

Scenario: Health system or healthcare organization purchasing/acquiring an AI solution from an external developer team to assist with imaging diagnostics (mammography), with an internal implementation team. Example Reporter(s): The Chief Medical Officer assigned the role of Reporter from the implementing/purchasing organization to work alongside relevant stakeholders (radiologists, radiology technicians, IT and security, patient privacy) to gather evidence on internal needs, processes, and capabilities to help guide the purchasing decision and design the broader AI system (e.g. end user engagement, operations, security and privacy capabilities, integration capabilities). They also work alongside the developer organization who assigns the Informatics Lead and Product Lead for the AI solution as Reporters from their respective organization, to address some of the Planning Checkpoint items and to gather evidence for best practice criteria in Responsible AI Checkpoint 1.

Example Reviewer(s): The procurement team may assign an internal reviewer (or consult with an external individual if further expertise is required), to review the evidence provided by the developer organization to help make a go-no go decision about purchasing. They may gather information from several potential vendors and use this checkpoint as a way of comparing vendor offerings, model performance, integration capabilities, transparency, privacy/security, etc. to guide the decision around which vendor to purchase from. The reviewer may instead choose to use this checkpoint as a way to select two vendors from which to pilot an AI solution internally, prior to making final purchase decisions. Once the decision to purchase or pilot is made, the implementing/purchasing organization may assign another reporter from the implementer team to help guide the initial pilot (which may lead to another go-no-go decision), or guide a small scale implementation process. Internal implementer and external developer teams will likely continue to collaborate to help troubleshoot problems that may arise during Responsible AI Checkpoint 2 and/or Responsible AI Checkpoint 3.

Additional Notes:

Developer organizations may choose to use the planning and other checkpoint checklists to help guide their development and piloting process, to help prepare for regulatory evaluation, and/or have external expert organizations review or validate the evidence they have provided. They may also choose to summarize the best practice evidence for respective checkpoints to share with potential clients, fostering transparency and trust.

In some cases, such as small community clinics or private practice settings, access to the full list of individuals required for an internal implementation or development team may not be available. In these cases these organziations may look for vendors who are already using best practices or who are willing to be transparent about their development process as outlined in the respective checklists. They may also choose to consult with external experts to help guide them through the purchasing and review processes in a way that is aligned with best practices and criteria defined here.