Enhancing trust in clinical decision support systems: a framework for developers

Caroline Jones, James Thornton, Jeremy C Wyatt

INTRODUCTION

Systematic reviews show that clinical decision support systems (CDSSs) can improve the quality of clinical decisions and healthcare processes and patient outcomes, although caution has been expressed as to balancing the risks of using CDSSs (e.g., alert fatigue) when only small or moderate improvements to patient care have been shown. Yet, despite the potential benefits, studies indicate that uptake of these tools in clinical practice is generally low due to a range of factors. The well-funded National Health Service (NHS) PRODIGY programme is an example of a carefully developed CDSS - commissioned by the Department of Health to support GPs - which failed to influence clinical practice or patient outcomes, with low uptake by clinicians in a large-scale trial. A subsequent qualitative study revealed that, among other issues — such as the timing of the advice — trust was an issue: ‘I don’t trust … practising medicine like that … I do not want to find myself in front of a defence meeting, in front of a service tribunal, a court, defending myself on the basis of a trial of computer guidelines’.

Another qualitative study exploring factors hindering CDSSs’ uptake in hospital settings found that clinicians perceive that CDSSs ‘may reduce their professional autonomy or may be used against them in the event of medical-legal controversies’. Thus, CDSSs may be ‘perceived as limiting, rather than supplementing, physicians’ competencies, expertise and critical thinking’, as opposed to a working tool to augment professional competence and encourage interdisciplinary working in healthcare settings. Similarly, a recent survey carried out by the Royal College of Physicians revealed that senior physicians had serious concerns about using CDSSs in clinical practice, with trust and trustworthiness being key issues (see examples below).

Trust is an important foundation for relationships between the developers of information systems and users, and is a contemporary concern for policymakers. It has, for example, been highlighted in the House of Lords Select Committee on Artificial Intelligence (AI) report; the Topol Review; a number of European Commission communications; reports; and most recently a White Paper on AI; and investigated in the context of knowledge systems, for example, for Wikipedia. Although it is an important concept, it is not always defined; rather, its meaning may be inferred. For example, the House of Lords Select Committee used the phrase ‘public trust’ eight times, but the core concern appeared to be about confidence over the use of patient data, rather than patient perceptions regarding the efficacy (or otherwise) of the AI in question. Such documents appear to take an implicit or one-directional approach to what is meant by ‘trust’.

Notably, the Guidelines of the High-Level Expert Group on AI outline seven key requirements that might make AI systems more trustworthy; whereas the White Paper focuses on fostering an ‘ecosystem of trust’ through the development of a clear European regulatory framework with a risk-based approach. Therefore, in keeping with the drive for promoting clinical adoption of AI and CDSSs while minimising the potential risks, here we apply Onora O’Neill’s multidirectional trust and trustworthiness framework to explore key issues underlying clinician (doctor, nurse or therapist) trust in and the use (or non-use) of AI and CDSS tools for advising them about patient management, and the implications for CDSS developers. In doing so, we do not seek to examine particular existing CDSSs’ merits and flaws in-depth, nor do we address the merits of the deployment process itself. Rather, we focus


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on generic issues of trust that clinicians report having about CDSSs’ properties, and on improving clinician trust in the use and outputs of CDSSs that have already been deployed.

Two points merit attention at this stage. First, O’Neill’s25 framework is favoured as—in the words of Karen Jones—O’Neill ‘has done more than anyone to bring into theoretical focus the practical problem that would-be trusters face: how to align their trust with trustworthiness.’26 Second, some nuance is required when determining who or what is being trusted. For example, Annette Baier makes clear that her own account of trust supposes: that the trusted is always something capable of good or ill will, and it is unclear that computers or their programs, as distinct from those who designed them, have any sort of will. But my account is easily extended to firms and professional bodies, whose human office-holders are capable of minimal goodwill, as well as of disregard and lack of concern for the human persons who trust them. It could also be extended to artificial minds, and to any human products, though there I would prefer to say that talk of trusting products like chairs is either metaphorical, or is shorthand for talk of trusting those who produced them.27

Similarly, Joshua James Hatherley ‘reserve[es] the label of ‘trust’ for reciprocal relations between beings with agency’.28 Accordingly, our focus is on the application of O’Neill’s25 framework to CDSS developers as ‘trusted’ agents, and measures they could adopt to become more trustworthy.

O’Neill’s trust and trustworthiness framework: A summary

O’Neill notes that ‘trust is valuable when placed in trustworthy agents and activities, but damaging or costly when (mis)placed in untrustworthy agents and activities’.25 She usefully disaggregates trust into three core but related elements:

1. Trust in the truth claims made by others, such as claims about a CDSS’s accuracy made by its developer. These claims are empirical, since their correctness can be tested by evaluating the CDSS.29

Trust in others’ commitments or reliability to do what they say they will, such as clinicians trusting a developer to maintain and update their CDSS products. This is normative: we use our understanding of the world and the actors in it to judge the plausibility of a specific commitment, such as our bank honouring its commitment to send us statements.

2. Trust in others’ competence or practical expertise to meet those commitments. This is again normative: we use our knowledge of the agent in whom we place our trust and our past experience of their actions to judge their competence, such as trust in our dentist’s ability to extract our tooth and the ‘skill and good judgement she brings to the extraction’.25

This approach utilises two ‘directions of fit’: the empirical element (1) in one direction (does the claim ‘fit’ the world as it is?), and the two normative elements (2-3) in another (does the action ‘fit’ the claim?).25 Relatedly, O’Neill has written on the concept of ‘judgement’, drawing a distinction between judgement in terms of looking at the world and assessing how it measures up (or ‘fits’) against certain standards (normative), versus an initial factual judgement of what a situation is, which ‘has to fit the world rather than to make the world fit or live up to a principle (empirical).30

In deciding whether to trust and use a CDSS, a user is similarly also making judgements about it. O’Neill’s threefold framework may therefore provide a helpful way to examine the issues in this context. In the following sections we discuss how CDSS developers can use each component of this framework to increase their trustworthiness, and conclude with suggestions on how informaticians might fruitfully apply this framework more widely to understand and improve user–developer relationships. Inevitably, this theoretical approach cannot address every potential issue, but it is used here as a means of organ-ising diverse concerns around trust issues into a coherent framework.

TRUSTING THE TRUTH CLAIMS MADE BY DEVELOPERS

CDSS developers might assume that their users are interested in the innovative machine learning or knowledge representation method used, or how many lines of code the CDSS incorporates. However, Petkus et al’s31 recent survey of the views and experience of 19 senior UK physicians representing the views of a variety of specialties provides some evidence of what a body of senior clinicians expect from CDSSs, that developers can use to shape their truth claims and build clinical trust. While this is not generalisable/representative of all clinicians it does provide a useful illustration of clinical concerns, and our intent is to demonstrate how applying O’Neill’s trust/trustworthiness framework might help our understanding of how to mitigate these issues. Table 1 shows the six clinical concerns about CDSSs which scored highest in the analysis. The score combines both the participant-rated

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Concerns about CDSS quality in Petkus et al survey</th>
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<tbody>
<tr>
<td>Concerns about CDSS quality</td>
<td>Score</td>
</tr>
<tr>
<td>The accuracy of advice may be insufficient for clinical benefit</td>
<td>15.5</td>
</tr>
<tr>
<td>How extensively was clinical effectiveness of CDSS tested</td>
<td>15</td>
</tr>
<tr>
<td>Whether CDSSs are based on the latest evidence</td>
<td>14.5</td>
</tr>
<tr>
<td>CDSSs can interrupt clinical workflow or disrupt consultations</td>
<td>14.5</td>
</tr>
<tr>
<td>CDSSs can ignore patient preferences</td>
<td>12.5</td>
</tr>
<tr>
<td>Whether the CDSS output is worded clearly</td>
<td>11.5</td>
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</table>

CDSS, clinical decision support systems.
severity of the concern and its frequency in the responses; the maximum score on this scale was 19. The greatest concerns here relate to O’Neill’s concept (or direction of fit) of empirical trust. Whether the advice provided by a CDSS is correct, has strong evidence for its clinical effectiveness from testing etc. ultimately concerns whether its advice ‘fits’ or matches (eg, in diagnosis) the patient’s actual condition. Can it (and/or the people that designed/made it) be trusted in an empirical sense of being factually correct?

What kind of truth claims may appeal to clinicians?

Drawing on the evidence in Table 1, developers should report to clinicians: the accuracy of the advice or risk estimates; CDSS effectiveness (impact on patients, decisions and the NHS); whether the CDSS content matches current best evidence (see ‘Guidelines: codes and standards frameworks’ below); its usability and ease of use in clinical settings; whether its output is worded clearly, and if takes account of patient preferences. These claims should be phrased in professional language, avoiding the extravagant claims about AI often seen in the press. Instead of different developers adopting a range of metrics for reporting study results there is a need for a standard CDSS performance reporting ‘label’ for these assessments, to help clinicians identify, compare and judge the empirical claims being made about competing CDSSs. This is by analogy with European Union (EU) consumer regulations dictating how, for example, tyre manufacturers report on road noise, braking performance and fuel economy for their tyres (figure 1), and EU plans for a health app label.

Ensuring that the truth claims can be verified

First, CDSS developers should be aware of the ‘evidence-based medicine’ culture, reflected in the top three concerns in Table 1. This means that, before clinicians make decisions such as how to treat a patient or which CDSS to use, they look for well designed, carefully conducted empirical studies in typical clinical settings using widely accepted outcomes that answer well-structured questions. This entails a ‘critical appraisal’ process to identify and reject studies that are badly designed or conducted, or from settings or with patients that do not resemble those where the CDSS will be used. So, it has long been established that empirical evaluation and the evidence it generates are crucial to generating trust. However, a systematic review of empirical research has shown that, when CDSS developers themselves carried out the study, they were three times as likely to generate positive results as when an independent evaluator did so. Therefore, studies that establish these truth claims should be carried out by independent persons or bodies. To counter suspicions of bias or selective reporting, the full study protocol and results should be made openly available, for example, by publication. Again, there is

Reading the tyre label

Your tyres will come with a label divided into three sections with information on:

1. Fuel Efficiency
   Depending on the tyre’s rolling resistance, its Fuel Efficiency Class will range from:
   - A is the most efficient tyre and will save you fuel;
   - G is the least efficient tyre and will use up the most fuel.

2. Wet Grip
   The Wet Grip rating tells you how well the tyres perform in wet conditions on a scale from A (safest) to G (worst performing tyre).

3. Noise
   A tyre’s noise level is measured in decibels (dB) using a three wave scale
   A brand space provides the manufacturer’s details, including the trade name/mark, tyre line, tyre dimensions, load index, speed rating, etc.

Figure 1 Example of an EU tyre label and how to interpret it.
Concerns about professional practice, ethics and liability

<table>
<thead>
<tr>
<th>Concerns about professional practice, ethics and liability</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>The legal liability of doctors who rely on CDSS advice is unclear</td>
<td>17.5</td>
</tr>
<tr>
<td>Some CDSSs act like a ‘black box’, with no insight possible for the user about how they arrived at their advice or conclusions</td>
<td>15</td>
</tr>
<tr>
<td>Doctors may follow incorrect CDSS advice, even if they would make correct decisions without it</td>
<td>13.5</td>
</tr>
<tr>
<td>CDSSs can embed unconscious bias, with some patient groups receiving unfair care as a result</td>
<td>12.5</td>
</tr>
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CDSS, clinical decision support systems.

In terms of potential legal liability, the situation does indeed appear to be unclear. Searches we carried out in legal databases (Lexis Library, Westlaw, BAILII), and PubMed, for terms around CDSS (adviser, expert system, risk score, algorithm, flowchart, automated tool, etc) turned up blank; nor have other researchers been able to locate published decisions in the UK, Europe or USA. However, it is well established that clinicians are legally responsible for the medical advice and treatment given to their patients, irrespective of the use of a CDSS. They must still reach the standard of the reasonable clinician in the circumstances. This makes it all the more important, if clinical uptake is to be improved, that clinicians have reasons to trust the CDSS developers and in turn their products/services.

How can CDSS developers facilitate this trust?

While developers cannot fix an uncertain legal framework, there are several steps they can take to nurture trust in this area. Most obviously, to ensure that the way the CDSS works and comes to its conclusions are made as clear as possible to users. It may not be realistic to do so completely, particularly as CDSS software becomes more complex via machine learning. However, giving—where possible—some account of the mechanism for how decisions are arrived at; the quality, size and source of any data-sets relied on; and assurance that standard guidelines for training the algorithm were followed (as well as monitoring appropriate learning diagnosties) will probably assuage some clinicians’ concerns.

In addition, even if some ‘black-box’ elements are unavoidable, clinicians’ anxieties regarding the dependability or commitment aspects of O’Neill’s trust framework may be alleviated by ensuring that frequent updates, fixes and support are all available. This should help clinicians feel more confident that the CDSS is likely to be reliable, and gives them something concrete to point to later to evidence their diligence and reasonableness, for example if they appear in court or at a professional conduct hearing.
Box 1  Developer actions that suggest competence and commitment to producing high quality clinical decision support systems (CDSSs)

► Recruit and retain a good development team with the right skills.58
► Use the right set of programming tools and safety-critical software engineering processes and methods, for example, HAZOP (Hazard and Operability Analysis) to understand and limit the risks of CDSSs.17 60
► Carry out detailed user research for example, user-centred design workshops; establish an online user community and monitor it for useful insights; or form a multidisciplinary steering group of key stakeholders.13 61
► Obtain the best quality, unbiased data to train the algorithm; use the right training method and diagnostics to monitor the learning process.46
► Implement relevant technical standards, obtain a CE mark (Conformité Européenne: the EU’s mandatory conformity mark by which manufacturers declare that their products comply with the legal requirements regulating goods sold in the European Economic Area) for their CDSS as a medical device.45
► Publish an open interface to their software; carry out interoperability testing.62
► Build on a prior track record of similar products that appeared safe.58
► Follow relevant codes of practice for artificial intelligence and data-based technologies.46 47
► Implement continuing quality improvement methods, for example, log and respond to user comments and concerns60; deliver updates to the CDSS regularly65; seek to become certified as ISO 9000 compliant.

TRUSTING OTHERS’ COMPETENCE

O’Neill23 suggests that we ask whether others’ actions meet, or will meet, the relevant standards or norms of competence. Factors that may impact positively on improving clinician trust include, but are not limited to, those listed in box 1.

In this section, we focus on the technical standards,45 and current codes of practice and development standards frameworks potentially applicable to CDSSs.46 47 Much more could be said about these approval processes than space permits here. However, the point is not to analyse the merits of the approval processes, but to illustrate how O’Neill’s framework helps to highlight their additional importance (beyond being strictly required) as a way to enhance (normative) trust.

UK and EU Regulation on Medical Devices

The initial question is whether CDSSs are medical devices? Classification as a medical device means that a CDSS will be subject to the EU Regulation on Medical Devices.45 The European Medicines Agency (the agency responsible for the evaluation and safety monitoring of medicines in the EU) states that ‘medical devices are products or equipment intended generally for a medical use’.48 Article 1 stipulates that ‘medical devices’, manufactured for use in human beings for the purpose of, inter alia, diagnosis, prevention, monitoring, treatment or alleviation of disease, means: ‘any instrument, apparatus, appliance, software, material or other article, whether used alone or in combination, including the software intended by its manufacturer to be used specifically for diagnostic and/or therapeutic purposes and necessary for its proper application’.45

In the UK, the Medicines and Healthcare Products Regulatory Agency (MHRA) has indicated that a CDSS is ‘usually considered a medical device when it applies automated reasoning such as a simple calculation, an algorithm or a more complex series of calculations. For example, dose calculations, symptom tracking, clinicians (sic) guides to help when making decisions in healthcare’.49 Hence, although some CDSSs may fall outside this definition (eg, by providing information only), our analysis is directed at those that do fall within the meaning of medical devices.

Accordingly, developers must adhere to the requirements of the EU Medical Devices Regulation45 and post-Brexit, under domestic legislation, namely the Medicines and Medical Devices Act 2021.50 These requirements include passing a conformity assessment carried out by an EU-recognised notified body (for medical devices for sale in both Northern Ireland and the EU), or a UK approved body (for products sold in England, Wales and Scotland)51 to confirm that the CDSS meets the essential requirements (the precise assessment route depends on the classification of the device).52 The focus of this testing is safety. Following confirmation that the device meets the essential requirements, a declaration of conformity must be made and a CE mark must be visibly applied to the device prior to it being placed on the market53 (from 1 January 2021 the UKCA (UK Conformity Assessed) mark has been available for use in England, Wales and Scotland,54 and the UKNI (UK Northern Ireland) mark for use in Northern Ireland).55 The general obligations of manufacturers are provided in Article 10 of the EU Medical Devices Regulation, including risk management, clinical evaluation, postmarket surveillance and processes for reporting and addressing serious incidents45; see also the ‘yellow card’ scheme operated by the MHRA which allows clinicians or members of the public to report issues with medical devices.56 Clinical users will rightly mistrust any CDSS developer who is unaware of these regulations, or fails to follow them carefully.

Nevertheless, NHSX (the organisation tasked with setting the overall strategy for digital transformation in the NHS) is seeking to ‘streamline’ the assurance process of digital health technologies.57 Similarly, in the USA, the Food and Drug Administration (FDA) is piloting an approach where developers demonstrating ‘a culture of quality and organisational excellence based on objective criteria’ could be precerti-fied.58 Such ‘trusted’ developers could then benefit from less onerous FDA approval processes for their future products due to their demonstrable competence.25

Guidelines: codes and standards frameworks

In addition to the generic Technology Code of Practice59 which should inform developers’ practices, there are two sets of guidance specifically focused on the development and use of digital health tools, including data-derived AI tools for patient management – one issued by the Department of Health and Social Care (DHSC) and NHS...
England, and the second by the National Institute for Health and Care Excellence (NICE). The DHSC and NHS England code of conduct aims to complement existing frameworks, including the EU Regulation and CE mark process, to ‘help to create a trusted environment’, supporting innovation that is safe, evidence based, ethical, legal, transparent and accountable. It refers to the ‘Evidence standards framework for digital health technologies’ developed by NICE in conjunction with NHS England, NHS Digital, Public Health England, MedCity and others. The aim of this standards framework is to facilitate better understanding by developers (and others) as to what ‘good levels of evidence for digital healthcare technologies look like’ and is applicable to technologies using AI with fixed algorithms; whereas those using adaptive algorithms are instead governed by the DHSC code (see Principle 7). Visible and/or certified compliance with these codes and standards would provide developers with normative objective standards to meet, and point clinical users of CDSSs to evidence of their competence. Having confidence in the professionalism of the developers should go some way towards reassuring clinicians as to the safety, accuracy and efficacy of CDSSs, thus potentially fostering greater uptake in practice.

CONCLUSION
O’Neill’s approach to trust and trustworthiness, focusing on empirical trust in developers’ truth claims and normative trust in their commitment and competence to meet those claims, has proved a useful framework to analyse and identify ways that developers can improve user trust in them, and in turn—it is suggested—the CDSSs they produce. That is, of course, not to suggest that developers are necessarily at fault in any way. It may be that they are unfairly distrusted by (potential) users. We suggest the application of O’Neill’s framework has helped to identify ways to facilitate and enhance trust in developers, and by extension, their CDSSs.

In summary, developers should:
- Make relevant claims about system content, performance and impact framed in professional language, preferably structured to a standard that allows clinicians to compare claims about competing CDSSs. These claims need to be supported by well-designed empirical studies, conducted by independent evaluators.
- Minimise ‘black box’ elements, ensure that internal mechanisms are—so far as possible—explained to users, and that CDSS software comes with a comprehensive update and support package. This could help clinicians gain a sense of control over the CDSS, and thus perceive the technology as a valuable working tool that complements their own skills and expertise.
- Comply with all relevant legal and regulatory (codes and standards) frameworks. Having confidence in the professionalism and competence of the developers should go some way towards reassuring clinicians as to the safety, accuracy and efficacy of CDSSs, thus potentially fostering greater uptake in their use.

The benefit of applying O’Neill’s framework is that it requires us to consider issues associated with different facets of both trust and trustworthiness, maximising the possibilities for enhancing trust and trustworthiness once such concerns or objections are overcome. An implicit or one-directional understanding of trust might result in a narrower conclusion, focused on just one element of O’Neill’s framework. For example, an understanding solely based on normative competence might focus on the importance of complying with the regulations (not only to avoid sanctions, but to enhance trust); this is important, but O’Neill’s framework demands consideration of different, equally useful, elements of trustworthiness.

This analysis is focused on clinician use of decision support tools, but we believe that a similar analysis would generate useful insights had we looked at other users and information systems, such as the public use of risk assessment apps, or professional use of electronic referral or order communication system advisory tools. The principles of examining the empirical truth claims of the software and the evidence on which they are based, then the competence of the supplier to match these claims and their commitment to do so, seems to generate useful insights no matter who the users are or what digital service is being trusted. Thus, we suggest that O’Neill’s framework is considered by health and care informaticians—both those developing and evaluating digital services—as a useful tool to help them explore and expand user trust in these products and services.

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45. EU Regulation on Medical Devices 2017/745.


