What can Safety Cases offer for patient safety? A multisite case study

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ABSTRACT

Background The Safety Case is a regulatory technique that requires organisations to demonstrate to regulators that they have systematically identified hazards in their systems and reduced risks to being as low as reasonably practicable. It is used in several high-risk sectors, but only in a very limited way in healthcare. We examined the first documented attempt to apply the Safety Case methodology to clinical pathways.

Methods Data are drawn from a mixed-methods evaluation of the Safer Clinical Systems programme. The development of a Safety Case for a defined clinical pathway was a centrepiece of the programme. We base our analysis on 143 interviews covering all aspects of the programme and on analysis of 13 Safety Cases produced by clinical teams.

Results The principles behind a proactive, systematic approach to identifying and controlling risk that could be curated in a single document were broadly welcomed by participants, but was not straightforward to deliver. Compiling Safety Cases helped teams to identify safety hazards in clinical pathways, some of which had been previously occluded. However, the work of compiling Safety Cases was demanding of scarce skill and resource. Not all problems identified through proactive methods were tractable to the efforts of front-line staff. Some persistent hazards, originating from institutional and organisational vulnerabilities, appeared also to be out of the scope of control of even the board level of organisations. A particular dilemma for organisational leadership was whether to prioritise fixing the risks proactively identified in Safety Cases over other pressing issues, including those that had already resulted in harm.

Conclusions The Safety Case approach was recognised by those involved in the Safer Clinical Systems programme as having potential value. However, it is also fraught with challenge, highlighting the limitations of efforts to transfer safety management practices to healthcare from other sectors.

INTRODUCTION

Patient safety remains a major challenge for healthcare, despite more than two decades of sustained policy, practice and research attention.1 2 The initial enthusiasm for borrowing practices and methods from other safety-critical industries (such as aviation) at the outset of the patient safety movement3–5 has been tempered by experience.6–12 It is now widely recognised that attempts to
Box 1 Typical features of safety cases

Safety Cases are developed to ‘make the case’ that risk has been reduced to a level ‘as low as reasonably practicable’ (ALARP). To do so, Safety Cases integrate various forms of prospective risk management analysis, based on the idea that operators are better placed than external regulators to assess risks in their own systems. The core of the Safety Case is typically a risk-based argument and corresponding evidence to demonstrate that all risks associated with a particular system have been identified, that appropriate risk controls have been put in place, and that there are appropriate processes in place to monitor the effectiveness of the risk controls and the safety performance of the system on an ongoing basis.23

Safety cases typically contain:
⇒ A description of the system and its operational context;
⇒ How safe the system is claimed to be and the criteria by which safety is assessed;
⇒ How hazards have been identified and how the risks they pose have been assessed;
⇒ What kind of risk control measures have been put into place and why they are effective; and
⇒ Why the residual level of risk is acceptable.23

Safety Cases are typically reviewed and assessed by an external regulator, for example, in the nuclear or petrochemical industries in the UK. However, some industrial sectors have also deployed the approach outside of a regulatory requirement. For example, the automotive industry uses Safety Cases that are part of the ISO26262 standard, but this is not mandated by regulators.17 18

transfer approaches between contexts require care and caution, and should be supported by theory and empirical evaluation.13–15 This paper seeks to contribute to addressing this need through examination of an attempt to introduce into healthcare a specific safety approach—the Safety Case—that is already used in other industries (including oil, transport and mining) both as a regulatory technique,16 and, more rarely, as a quality management approach without regulatory mandate (eg, in the automotive industry).17 18

The specifics of the Safety Case approach vary between sectors and regulators,19 but the general principles are listed in box 1. In brief, a claim to operational safety is justified through a series of linked arguments that explain how safety has been secured, with supporting evidence, including the processes in place to control risk. Where used as a regulatory technique, Safety Cases are produced by organisations to ‘make the case’ to the relevant regulator that they have put in place adequate measures to reduce risks in a product or system to a level ‘as low as reasonably practicable’ (often abbreviated as ALARP). The regulator then reviews the Safety Case and either grants the organisation licence to operate, or may require further risk assessments, justification of the measures proposed or additional risk mitigations.20

As an approach requiring organisations to proactively describe what procedures and actions they are putting in place to control risk, Safety Cases can be contrasted with prescriptive, compliance-oriented approaches, where organisations are required to show that they have met externally imposed safety standards.21 Because they are written for a specific system and its context of use, they are intended to be more adaptable to specific situations than generic safety standards, and also more responsive to rapid change in technologies or practices.22

On the face of it, the Safety Case would appear to have value as an approach to safety management in healthcare, particularly in its potential for prospective identification and control of risk. However, the Safety Case approach has only rarely been used in healthcare, and only in a very limited number of applications (eg, development of information systems and medical devices).23 24 In this article, we develop an analysis of the application the Safety Case approach within the UK National Health Service (NHS) using a case study of the first documented attempt to apply the principles of the methodology to clinical pathways. As the approach was deployed outside a regulatory context, our analysis focuses on the transferability of an approach to risk management that is proactive, structured, and tailored in nature and that presents evidence about the safety of specific clinical systems and existing mitigations in a single ‘case’ document.

METHODS

Case study: the Safer Clinical Systems programme

Our analysis draws on an evaluation we conducted of a programme known as Safer Clinical Systems, which is designed to improve the safety and reliability of clinical pathways based on learning adapted from a range of hazardous industries. It seeks to enable organisations to make improvements to local clinical systems and pathways through a structured methodology for identifying risks and re-engineering systems to control risk and enhance resilience.25 26 Use of the principles of the Safety Case approach is a centrepiece of the Safer Clinical Systems programme, although outside a regulatory context.

Funded by the Health Foundation, the Safer Clinical Systems programme was developed by a team at Warwick University and tested over a number of phases. Following initial development, a ‘testing phase’ involving eight NHS hospital sites (seven in England, one in Scotland) ran from 2011 to 2014. An ‘extension phase’ (2014 to 2016) involved further work by five of these sites and one new site.

Each participating hospital site (table 1) was required to establish a multidisciplinary clinical team. Sites in...
the testing phase were advised by a support team of clinicians and experts, received inperson training, had access to other resources (such as a reference manual and telephone support) and were required to report their progress regularly. Sites in the extension phase had less bespoke support and were expected instead to build on their previous learning.

A requirement of participating teams was that they use the Safer Clinical Systems approach to proactively assess risks and hazards in their clinical pathways and that they produce Safety Cases at the end of their projects describing the risks and how they were being mitigated. The Safety Cases were expected to be similar in format to those used in other sectors, comprising a description of the clinical pathway covered, the key hazards identified through structured analysis using prescribed tools, the risk controls implemented, and, critically, a ‘safety claim’ and associated ‘confidence argument’—a pronouncement on the current safety of the system concerned, and a statement explaining how risks had been made ALARP. Rather than being presented to an external regulator, as would be the case if the Safety Case were being used as a regulatory technique, the principal intended audience in this programme was the senior leadership (executive and board level) within organisations.

**Evaluation methods**

To study the testing and extension phases of the Safer Clinical Systems programme, we used a mixed-methods, longitudinal design, involving interviews, ethnographic observations, and documentary analysis across the nine participating sites. The analysis we report here is based primarily on interviews and documentary analysis. Ethnographic observations (over 850 hours) provided valuable data on how clinical teams carried out their Safer Clinical Systems projects in practice in the context of existing and competing demands, but are not reported in detail here.

Across the nine sites, we conducted 89 semistructured interviews in the testing phase and 39 in the extension phase with participating clinical team members and programme leaders. Sampling at the sites sought to purposefully include a range of different roles in the programme, including the clinical leaders of each project and others. We also conducted 5 semistructured interviews in the testing phase, followed by 10 in the extension phase, with organisational senior leadership, comprising executive team/board members. Interviews explored general experiences of the programme as well as specific exploration of using the Safety Case approach. Participants were informed of the aims and commissioners of the evaluation. All interviews were conducted by experienced social scientists using topic guides (online supplemental material 1). Interviews were conducted either in person or by telephone, between November 2012 and June 2016, and were digitally audio recorded and then transcribed for analysis.

Analysis, conducted by EL and guided by the wider team, was based on the constant comparative method combining inductive and deductive approaches. We coded interviews and observations using an inductive approach, deriving codes directly from each interview and then progressively clustering codes in higher order categories and themes. To strengthen explanatory
power, this inductive strategy was complemented by theoretical concepts drawn from the wider literature. GL and EL conducted a documentary analysis of the Safety Cases prepared by the clinical teams (table 2). We used recommendations and guidelines for writing and maintaining safety cases in other sectors,29–31 to organise the Safety Cases’ content thematically, and identified their main strengths and weaknesses in terms of completeness, presence of appropriate evidence and analyses to support the claims, consistency with the site’s safety improvement objectives, readability, and presence of a safety claim and confidence argument. Finally, we organised our higher order themes and overall reflections using concepts and themes proposed by recent works on the topic.19 32 Regular team meetings and correspondence provided oversight of the analytical approach, consistency and adequacy of codes, and reporting. Given the nature of the programme, we did not undertake a formal test for theoretical saturation for the interviews or the Safety Cases.

Findings
Across the testing and extension phases of Safer Clinical Systems, we undertook 143 interviews with participants across programme leadership, clinical teams and organisational leadership. We analysed 13 submitted Safety Cases; although 14 should have been developed, one site from the extension phase struggled to implement the programme in full and did not produce a Safety Case. In presenting our analysis below, we consider, first, participants’ views on the Safety Case as a novel approach to understanding and managing safety risk in healthcare, and second, the work that went into developing Safety Cases. We then turn to the analysis of Safety Cases themselves.

Views on the value of safety cases
By the end of the programme, members of the project teams and senior leadership in the participating organisations had largely come to see the Safety Case as a valuable approach, with the potential to make hazards visible in an accountable, systematic and scientific way. The analytical steps required to compile a Safety Case, such as process mapping the patient pathway, were seen to be particularly useful in proactively identifying threats to safety, rather than reactively managing incidents once they had happened. The role of Safety Cases in enabling an overarching, system-wide view of the hazards, rather than focusing on what happens in particular segments of the pathway, was also welcomed. Broadly, teams valued the possibilities of new ways of thinking about risk.

I like the idea that you just have one document that you can hand to somebody and say how safe is your system. I like the concept that you can say ‘Well this is what our system is like just now’. (Project participant)

Some organisational senior leaders agreed, at least in principle, that Safety Cases could offer value, and recognised the importance of a prospective approach to safety.

We have immensely complex systems which could be simplified and therefore made a bit more reliable. […] So something which looks at that could certainly be a useful thing, because it’s saying ‘Well actually here is a little nest of complexity which you can reduce, but it’s also a significant risk to the patient, because you’re missing information or you’re hurrying things through.’ […] (Senior leader)

Other senior leaders, however, were not always clear on the practicalities of the approach, and some found it difficult to identify the added value of Safety Cases. They suggested, for example, that existing risk management tools performed very similar functions.

Table 2 Format and content of 13 Safety Cases reviewed

<table>
<thead>
<tr>
<th></th>
<th>Median (min-max)</th>
<th>Number of Safety Cases with available data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reporting (number of pages)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Safety Case (without executive summary and appendices)</td>
<td>20 (14–40)</td>
<td>13</td>
</tr>
<tr>
<td>Executive summary</td>
<td>2.5 (1–7)</td>
<td>8</td>
</tr>
<tr>
<td>Total length of submitted document (including executive summary and appendices)</td>
<td>43 (14–360)</td>
<td>13</td>
</tr>
<tr>
<td>Identification of hazards in the clinical pathway</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of hazards or failure modes identified</td>
<td>22 (5–99)</td>
<td>11</td>
</tr>
<tr>
<td>Number of ‘high-risk’ hazards identified</td>
<td>7.5 (4–36)</td>
<td>8</td>
</tr>
<tr>
<td>Choice of risk controls</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of interventions selected</td>
<td>7 (2–40)</td>
<td>11</td>
</tr>
<tr>
<td>Measurement of progress</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of measures in ‘safety set’</td>
<td>5.5 (3–13)</td>
<td>12</td>
</tr>
</tbody>
</table>
delineation is between Safety Case and risk register. (Senior leader)

Some project teams saw the Safety Case as useful for a secondary reason: that of securing the attention and interest of senior leaders in their organisations. Their hope was that, by providing new evidence and analysis of the riskiness of clinical systems, senior management attention, support, and resources might be solicited.

So they’ve [senior management] actually kind of bought into it, so I think they will feel pressure to deliver. (Project participant)

However, as we explain below, the exact fit of Safety Cases into the existing ecology of tools and documents in healthcare was not clear to all participants.

Preparing safety cases

Project teams were required to learn new techniques to prepare the Safety Cases, including use of systematic methods to identify and assess risks in their clinical pathways, to propose risk controls and to identify metrics that could be used to monitor systems. Production and communication of Safety Cases also required skills in making persuasive claims, structuring arguments and presenting evidence compellingly. The participating teams were, understandably, unfamiliar with many of these skills, and expressed uncertainties about the expected structure, content and style of the Safety Case itself, especially in terms of what issues to emphasise and how to evidence them. Participants described compiling and drafting the Safety Case as labour-intensive and difficult.

I think the other bit that we have been challenged by is the actual writing of the Safety Case and again it is because it is fairly new to healthcare in general. I think we are going to go through a few reiterations before we fully understand what it is and how to use it. (Project participant)

Notwithstanding the training and support received in the ‘testing’ phase, teams continued to report difficulties with preparing and drafting Safety Cases well into the extension phase. A recurrent source of ambiguity related to the size and scope of the clinical system that the Safety Cases should target. The first, diagnostic, step in the Safer Clinical Systems process involved defining the clinical pathway of focus. However, determining the boundaries of the pathway was far from straightforward. Furthermore, clinical pathways typically involved dozens of technological systems (eg, infusion pumps, IT systems) and socio-technical processes (eg, guidelines, multidisciplinary meetings). Each might be amenable to risk assessment and management individually, but making sense of their connections, aggregate risks and potential interactions was a much more complex task.

It’s not a linear process and you do go back trying to understand another bit of the process that you thought you understood, but actually didn’t as (…) you had hoped. (Project participant)

Once the pathways and their components had been determined (or at least approximated), project teams used a range of methods recommended by the Safer Clinical Systems programme, mostly derived from similar activities in other industries, to assess hazards and risks. The teams found the processes often challenging and time-consuming, with much discussion about the relative merits of different sources of data and evidence. Despite the challenges, teams generally concluded that conducting a systematic risk assessment using structured tools offered important new insights about clinical pathways.

What I’ve loved doing is, is talking to the staff and actually understanding what goes on, because it’s only when you understand what goes on that you can put it right… You’ve worked in the hospital for years and there’s still things you didn’t realise actually went on and things that people did that you didn’t realise that they actually did. That was quite an eye-opener. (Project participant)

This new understanding through structured risk assessment enabled teams to identify multiple short-comings that had potential to harm patients. The hazards they unearthed varied greatly in scale, level of risk posed and tractability to intervention. Some problems identified were amenable to resolution by the project teams, typically those with their roots in suboptimal service planning and pathway design, failures in communication among staff, or unclear distribution of responsibility or ownership of key processes. In response to these, most, but not all, sites designed or implemented some risk controls and documented them in their Safety Cases.

[Staff are] given the freedom and the autonomy to go ahead and do whatever things they think might be necessary to make things better. And that’s what people do, there is very much a culture of promoting change there, so they talked about small cycles of change, doing PDCA [Plan Do Study Act] cycles, and there’s a number of different projects that are running. (Observation notes)

The extent to which these risk control interventions were consistent with the principles of the Safer Clinical Systems programme varied by site. Some project teams were able to draw on extensive experience, while others foundered at this stage. Common to all sites, however, was the identification of issues that were well beyond the scope of control of the front-line teams themselves. These vulnerabilities tended to originate from deep-rooted institutional and organisational pathologies or constraints. The importance of these problems, including, for example, staffing levels, was beyond doubt. Exactly what to do about them was less clear. Some project teams made valiant attempts to at least mitigate the risks through local work, but others
appeared to accept that standard quality improvement efforts would not solve the issues. Some teams described the ongoing failure to mitigate the risks in their Safety Cases, in part, as noted above, in the hope that action from senior level might be provoked.

There were other things that were discussed at the [meeting] that they thought would be good as a team to change... but with some of them, they just knew it would be impossible to do so, so actually they didn't even bother to write them down. (Observation notes)

And the team very bravely went to the board and said, you know, our Safety Case is showing and we're telling you that our processes are unsafe, so it alerted people to the issues. [...] So that was the strength of it. (Project participant)

However, as we now describe, for senior organisational leaders, both the imperative offered by the Safety Case and their own ability to act were less clear.

Content of, and responses to, safety cases
Our documentary review showed that submitted Safety Cases were highly variable in format and length (table 2). Some were highly structured, clearly written and precise in the use of evidence; others were harder to follow, lacking in clarity and less well organised. Our review also found that the descriptive elements (analysis of risk and hazards) were much better achieved than the assurance components (the safety claim and the confidence argument). Indicative, perhaps, of the intractability to local-level intervention of some of the hazards uncovered, or the lack of expert safety science input in the project teams, most Safety Cases focused more on what had been done to determine the risk than on the level of safety that had been achieved in mitigating it. The documents also varied in the extent to which they reported the residual risks—those that remained despite the implementation of risk controls—in a clear and transparent way. For instance, one Safety Case noted that the diagnostic process had found 99 ways in which the pathway could fail, that the level of reliability in the microsystem remained lower than acceptable, and that radical re-design was needed. Others were more circumspect. Accordingly, while they documented sometimes-extensive mitigations, none of the Safety Cases could make an unambiguous safety claim supported by a powerful confidence argument. Some teams were not clear about how the evidence gathered and analyses conducted would contribute to the safety claim. Some sites listed project activities in lieu of offering an actual safety claim, reporting what they had done rather than the level of safety they had reached.

It was a useful, [...] a really good repository for all the stuff we’ve done in the project, which I find really good. And has been good when people ask ‘What did you do?’ then you can say that this is what we did, so that’s useful. I’m not sure about whether people use it for what it is meant to be, which is to prove the pathway is now safe, I’m not sure whether it is used for that really. (Project participant)

Sometimes, safety claims were reported for each identified hazard (comparing levels of risk before and after the interventions they had implemented) rather than at the level of the clinical system. No site explicitly discussed whether risks had been reduced ‘as low as reasonably practical’. Some sites claimed improvements as a result of the interventions they had implemented, but these did not always stand up to statistical scrutiny.33

The response of senior leadership to the Safety Cases submitted by teams varied. Some focused on the potential of the Safety Case for supporting organisational-level decision making in relation to risk reduction, resource allocation and strategic prioritisation.

I think it would be easier to respond to a Safety Case rather than more so the [other quality and safety] data I get. Because it’s back to first principles, what are we actually here to do... Then if we have an unsafe system everything else needs to fall in behind that, no matter cost pressures, no matter personal opinion, no matter all the other complexities in a big system. If an element is at risk, then that will always be made a priority. (Senior leader)

Not all senior leaders, however, were so confident that the insight offered by Safety Cases would or should inevitably lead to action. Some of the issues identified in the Safety Cases were beyond the ability not only of front-line teams to solve, but also of organisational leaders. Issues such as staffing levels, IT interoperability, and securing timely discharge required at least interorganisational coordination, resourcing, coordination, and support across the whole healthcare system. Additionally, the prevailing approach to risk management, and the perceived unavoidability of risks in the complex systems of healthcare, meant that the insights offered by a Safety Case might be unwelcome or not necessarily candidates for priority attention. In a system that relied primarily on retrospective risk management approaches, such as incident reporting and investigations, the need to tackle risks of recurrence (where problems had already manifested as serious incidents or ‘near misses’, and might do again) could easily take precedence over addressing seemingly ‘theoretical’ risks (problems identified through a detailed prospective analysis but yet to occur).

Because you’re saying actually ‘That was a potential harm on our risk management system, and we knew about it, and we were accepting that we don’t have enough money to address all of these issues at one time’. So there is, if you like, a prioritisation and rationing of where we put money according to the level of risk. [...] It’s a bit like county councils putting crossings on roads, or a zebra crossing. You’re waiting...
Some feared that, given the legal obligation of boards to take action in response to safety risks that were revealed to them, an unintended consequence of the Safety Case approach might be to distract organisational focus from areas that were at least as worthy of attention but lacked the spotlight offered by the Safety Case. There was a perception that to have a Safety Case for every pathway or area of practice would likely be impossible, and that too many Safety Cases would be overwhelming.

The complexity of health care is such that there are hundreds of complex connected pathways that patients are on and so... You in theory could write hundreds [of Safety Cases] and that would then become meaningless because if you write hundreds no one would ever read them. So, I think it might be helpful in some specific examples... Rather than being something that could cover everything that we do to patients. (Senior leader)

Consequently, Safety Cases might serve not to assure about control of risks, but to unnerve—and unnerve leaders who were not always well placed to act, given the scope of their control and the other priorities they faced. In a system where Safety Cases were new, without an established function in safety management, and covering only a small proportion of safety-critical activity, the information they provided was not always readily actionable from a managerial perspective and, moreover, had potential to create uncontrolled reputational risk.

The danger is that what you have is a legal requirement to spend money on a Safety Case that actually is of low, relative risk to harms that are occurring in the absence of Safety Cases. So what you get is a spurious diversion of money to a wheel that has been made very squeaky, but actually isn’t causing harm... There’s the risk of diversion to get a perfect patch in one part of the system while everything else is actually terrible. (Senior leader)

(A danger) is, you know, if it does get into the wrong hands, particularly with the media, because there’s not the openness and the ability to manage some of this data, which needs explanation. But we do pride ourselves on being a very open and transparent board. (Senior leader)

**DISCUSSION**

Our examination of an attempt to introduce the principles and methodologies of the Safety Case approach into healthcare suggests that the approach was broadly welcomed by participants in our study, but was fraught with challenge. In other sectors, the Safety Case rests on the ALARP principle. While the Safety Cases produced by participating teams in the Safer Clinical Systems programme did present proactive analyses of risks, they did not show that the risks in clinical pathways on which they focused had been reduced as far as reasonably possible. Instead, teams identified multiple residual risks that had resisted efforts at control and mitigation by the teams themselves. These findings emphasise the importance of careful consideration of context and implementation when transferring safety management approaches from one setting to another. The evidence underlying other industrial risk management techniques (e.g., Failure Modes and Effects Analysis, ‘5 Whys’ or Root Cause Analysis) is also weak, but the regulatory function of Safety Cases warrants specific caution. Sujan et al’s review of various sectors nonetheless concluded that even with the differences in regulatory context, healthcare organisations could benefit from using the Safety Case approach to develop understanding and exposition of their current levels of risk.

An important feature of the programme we examined—essentially a feasibility study—was that the Safety Case approach was being used outside the regulatory frameworks and infrastructures characteristic of use of the technique in most other sectors. Without an external regulatory requirement to satisfy, participating organisations in the Safer Clinical Systems programme may not have felt a strong imperative to make the responses that might otherwise be expected; absent the spectre of regulatory action, senior leadership may not have felt compelled to reduce the risks ALARP. However, even when Safety Cases are part of a regulatory framework, they are not always rigorous or successful in controlling risk or showing they have been reduced ALARP. While our study does not allow conclusions to be drawn about what might happen if Safety Cases were included in a regulatory regime in healthcare, it does allow insights into the nature of the challenges that might be anticipated should regulators consider introducing the approach in healthcare settings.

Some of the challenges we identified arose from the mismatch between the complexity and interdependencies of clinical pathways, with their often unbounded character, and the more tightly defined (and often more mechanical or technical) applications of the approach in other industries. Future research might usefully clarify whether and how the scope of a Safety Case could best be defined for healthcare settings, noting that the highly dynamic and interdependent nature of multiple subsystems of care may defy attempts to impose clear boundaries. These kinds of questions are becoming increasingly prominent in safety science as recognition grows that the development of networked
complex systems (eg, unmanned aircraft systems) requires a shift from relatively static prelaunch assessment to a dynamic approach that can accommodate changes in the system’s properties and behaviour during its life-cycle.\textsuperscript{41,42}

Other challenges arose in the demanding nature of the expertise, skill and time commitment required to engage in the tasks of conducting safety analyses, identifying and testing risk controls, and compiling a Safety Case. The variable quality of the Safety Cases submitted by clinical teams in this programme is likely to be linked to variable competencies and available capacity. In contrast, in safety-critical industries where these risk assessment techniques originated, the design of effective risk controls is the responsibility of safety/reliability engineers with extensive training and expertise. For healthcare, use of the Safety Case approach will require additional resource and new dedicated roles with specific expertise, rather than relying on making further demands of existing clinical teams.\textsuperscript{40,43}

The resourcing implications of a wholesale effort to shift the regulatory system and culture of an entire sector could, however, be enormous, especially given the volume and complexity of activity in healthcare and the number of diverse clinical pathways.

An additional set of challenges was more cultural in character, and related to the revelatory potential of the Safety Case. On one hand, participants—especially clinical teams—appreciated the value of the Safety Case in offering a proactive, prospective and rigorous approach to identifying safety risks. Some also saw it as a means of attracting managerial attention and obtaining resources.\textsuperscript{44} But leaders in organisations were not always convinced that the approach offered much that was new, suggesting that more evidence would be needed to demonstrate the added value of Safety Cases—especially in moving beyond description to solution,\textsuperscript{45} and adding value over current approaches such as risk registers. A further concern at the leadership level was that it was unclear whether areas that did have a Safety Case should be considered to have a stronger warrant for action than those that did not. A framework for supporting prioritisation of risks is likely to be helpful in any future use of Safety Cases. However, current tools, such as risk matrices, may be flawed,\textsuperscript{46,47} so better tools should be investigated.

Even less tractable was what to do about some of the problems reported in the Safety Cases. Clinical teams had done their best to implement risk controls where they could, but they did not have sufficient power and access to resources to address those that were institutional or structural in character. They therefore often fell back on weaker administrator measures, like training or procedures.\textsuperscript{8} Yet organisational leaders were often similarly challenged, given their limited capacity and resources for radical systems re-design, improved staffing, IT infrastructure, or other major re-engineering or influencing of activities outside the organisation itself. These findings are indicative of broader problems with the selection of risk controls in health services.\textsuperscript{44,48} What may need to be addressed before Safety Cases could achieve their potential.

Our study has a number of strengths, including its in-depth, mixed-methods, longitudinal design with engagement both with the project teams and senior leaders in organisations. It was limited in its ability to assess the impact of the Safety Case approach in improving safety, not least because of issues with data on processes and outcomes.\textsuperscript{33}

CONCLUSIONS

The Safety Case approach offers promise in principle as a safety management approach in healthcare, but substantial challenges need to be addressed before further deployment, particularly in regulation. Further experimentation with the use of Safety Cases in healthcare might therefore more profitably focus on how to make the most of their assets—including the new insights offered by prospective, system-wide risk analysis—while managing their potential unintended consequences.

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SCS Evaluation Topic Guide
Site Level

TOPIC GUIDE

Evaluation of Safer Clinical Systems – Testing phase

Interviews with site teams about how the Safer Clinical Systems approach is expected to improve patient safety

What is your present job title?

What is your role on the Safer Clinical Systems Phase 2 project within your organisation?

What motivated your organisation to get involved in this project?

What motivated you to join the Safer Clinical Systems team in your organisation?

What did you find appealing about the Safer Clinical Systems approach when you first heard about it? Was there anything off-putting?

Which elements of the Safer Clinical Systems approach did you find the most useful during Step 1 (Pathway and Context)? Which were the least useful?

Which elements of the Safer Clinical Systems approach did you find the most useful during Step 2 (System Diagnosis)? Which were the least useful?

What elements or tools have most helped your work on the project to date?

What have been the main challenges so far?

How will you know if Safer Clinical Systems has been successful in improving your selected pathway?

Do you think the Safer Clinical Systems approach will work in practice? Can you describe what you think are the mechanisms through which taking part in Safer Clinical Systems will deliver improvements in patient safety in your organisation?

What are the risks associated with this approach?

Are there any other aspects of Safer Clinical Systems and your experiences to date that you would like to discuss?
TOPIC GUIDE

Evaluation of Safer Clinical Systems – Extension phase

Interviews with site teams about how the Safer Clinical Systems approach is expected to improve patient safety

Tell me about your current project. What motivated your organisation to do this one? What are you trying to achieve with it?

Can you tell me how your team is set up. Who is in the team and why those people? What skills and what roles did you feel were essential?

Why was it felt that the Safer Clinical Systems approach was a good way of achieving those aims?

Which are the really essential parts of the Safer Clinical Systems approach from your point of view?

Which elements of the Safer Clinical Systems approach are you planning to use/have used so far on the project? If you have decided to drop some bits of the approach, why is that?

Can you tell us about the sequence in which you have use or will be using the tools and techniques from Safer Clinical Systems?

If you are planning to use any of the diagnostic tools, can you tell me a bit more about that. Why is it important to use the ones you have chosen and what do you expect the benefits will be?

Tell me more about how you are planning to address any hazards you identify in your clinical systems. What kind of interventions or risk controls are you planning to introduce? How have you (or will you) gone about choosing these? How are you expecting that these interventions or risk controls will address the problems you have found? How much time have you been able to spend looking at the published evidence on these kinds of interventions? What challenges do you expect in implementing the interventions?

Can you tell me about the improvement methods you will be using. Are you planning to use PDSA cycles? If so, why is that the best way of tackling the problems you have identified? What other improvement methods did you consider? If you can’t get the change you need using your existing methods, what would you do?

Can you tell me about the measurement in this extension project. Are you still using the Safety Set idea? What data are you collecting? Is it all on reliability or are you measuring other things as well? How have you gone about aligning the measurement to the hazards and the risk controls or interventions. What is your experience of setting up the data collection systems and doing the analysis?

I think you will be preparing at least one Safety Case as part of your project. What is the purpose of the Safety Case? What are your views on Safety Cases, including their strengths and weaknesses, based on your previous experience? Where will it be read and by whom? Do you think Safety Cases are a good way of communicating with the senior people in the organisation – such as those at board and executive level? What kinds of problems do these people have in understanding the Safety Case idea? Do you think they feel that the Safety Case makes clear to them what they need to do to make care safer and do they understand what their responsibilities might be in supporting that change?
SCS Evaluation Topic Guide
Site Level

What do you think the Safer Clinical Systems approach will do well? Where do you think it is likely not to work so well?

What difference has it made doing the project on your own this time, without the Support Team’s involvement?

What support and resources have been provided by your organisation?

Who needs to be engaged in this project for it to work? How easy has it been to engage people in the project?

How will you know if this project has been a success?
SCS Evaluation Topic Guide
Site Level

TOPIC GUIDE
Evaluation of Safer Clinical Systems – Extension phase

Interviews with senior leaders about how the Safety Case approach

Can you tell me your job title and give a little background to your role in the organisation?

Can you tell me about any involvement you may have with the Safer Clinical Systems project at your hospital?

Can you tell me a little about the quality and safety data that would normally come to you? How useful do you find it? Do you receive it in a timely way? Is it easy to interpret? How easy do you find it to make that information actionable?

How would you know if there was a hazard in a clinical system? Would you normally expect the evidence about hazards to arise retrospectively – in other words after an incident has happened? What would happen once an incident has occurred to fix the problem?

Can you tell me about any ways you could figure out in advance of an incident happening whether there are hazards in your clinical systems?

How valuable would you find it to have a document that reported an assessment of hazards in a system, quantified the risks, and explained the risk controls in place and whether further action was needed? Would it be a useful addition to what you already receive or should it replace any data you get already?

That kind of a document is called a Safety Case in other industries. If they are going to be used in healthcare, it would be very useful to identify the priorities of senior leaders for format and presentation and that’s where the rest of my questions will be focused.

First I am going to take you through a possible structure for a safety case. To help keep our discussion focused, we will use the example of maternity care, but we are interested in general reflections.

Safety Cases should explain what clinical pathway they cover and describe the facility or physical structures where care is located. Any comments on what might be helpful in terms of the format or presentation of this kind of information?

Safety Cases should then define what must be right and why for this particular pathway and facility. What would you expect to see here?

Safety Cases then document all the things that might go wrong. Usually this will be the outcome of a structured process for identifying the hazards in the system – for example, not having an intensive care unit co-located with a maternity unit might be identified as a hazard. It also quantifies the risks associated with this hazard – for example a 0.5% risk that a patient might experience a catastrophic outcome associated with not having an ICU nearby. How useful would you find this kind of documentation?

Next a Safety Case will identify the risk controls in place, by describing what is being done to mitigate against things going wrong. For example, it might explain the special training staff that have in preventing post-partum haemorrhage so that the risks of a patient requiring ICU care are reduced. The Safety Case should also explain what will happen if it still goes wrong – what risks and consequences would follow, and what the emergency arrangements would be. Again, how useful would you find this?
The Safety Case will explain whether the risks are as low as reasonably practical, and indicate what needs to be done to make the system safer and what the limitations and uncertainties might be. For example, it might recommend that an additional specially trained obstetric anaesthetist is always available. Again, what are your feelings about having this kind of assessment and recommendation? Would it be any more useful than a standard business case? What kind of evidence would you expect to support the claims made in the Safety Case? Do you think you would find it more or less easy to take action in response to a Safety Case compared with the other data you get coming in at the moment?

How would you feel about having a Safety Case from your organisation made publicly available?

In other industries Safety Cases have legal status and are used as the basis of regulating a particular facility. This means that if a particular risk control is identified in the Safety Case as essential to safe operation of the facility, it operates as a legal requirement. How would you feel about this applying in healthcare?

Finally, in terms of presentation: Would you favour having an executive summary at the beginning and a longer technical document that presents the main Safety Case? What’s the maximum length of such a document if it’s going to be useful for you?