Objectives: Unintended events (UEs) are prevalent in healthcare facilities, and learning from them is key to improve patient safety. The Prevention and Recovery Information System for Monitoring and Analysis (PRISMA)-method is a root cause analysis method used in healthcare facilities. The aims of this systematic review are to map the use of the PRISMA-method in healthcare facilities worldwide, to assess the insights that the PRISMA-method offers, and to propose recommendations to increase its usability in healthcare facilities.

Methods: PubMed, EMBASE.com, CINAHL, and The Cochrane Library were systematically searched from inception to February 26, 2020. Studies were included if the PRISMA-method for analyzing UEs was applied in healthcare facilities. A quality appraisal was performed, and relevant data based on an appraisal checklist were extracted.

Results: The search provided 2773 references, of which 25 articles reporting 10,816 UEs met our inclusion criteria. The most frequently identified root causes were human-related, followed by organizational factors. Most studies took place in the Netherlands (n = 20), and the sample size ranged from 1 to 2028 UEs. The study setting and collected data used for PRISMA varied widely. The PRISMA-method performed by multiple persons resulted in more root causes per event.

Conclusions: To better understand UEs in healthcare facilities and formulate optimal countermeasures, our recommendations to further improve the PRISMA-method mainly focus on combining information from patient files and reports with interviews, including multiple PRISMA-trained researchers in an analysis, and modify the Eindhoven Classification Model if needed.

Key Words: PRISMA, root cause analysis, root cause, unintended event, incident, error, Eindhoven Classification Model, patient safety

Abbreviations: ECM = Eindhoven Classification Model, ED = emergency department, GLD = gastroenterology and liver disorders, ICU = intensive care department, MERS-TM = medical event reporting system for transfusion medicine, NICU = neonatal intensive care unit, NHLBI, NIH = National Heart, Lung, and Blood Institute Quality Assessment Tool for Observational Cohort and Cross-Sectional Studies, PICU = pediatric intensive care unit, PRISMA = Prevention and Recovery Information System for Monitoring and Analysis, RCA = root cause analysis, UE = unintended event

One of the key elements to improve patient safety is learning from adverse events. Especially unintended events (UEs) can be a valuable source of information to improve patient safety. Unintended events are a broad group of events, including near misses, that might but do not necessarily result in patient harm. They can be both preventable and unexpected. In clinical practice, there is an increasing interest in the development of systems to systematically evaluate UEs, because several large studies showed that medical errors result in patient harm, leading to temporary or permanent disability or death. To gain insight into the quality of care, it is necessary to know more about the types of UEs and their root causes, which will simultaneously give insight into possibilities for prevention of UEs. However, there is no standard method to systematically collect, analyze, and compare data related to UEs and their causes among healthcare facilities.

A frequently used approach to improve patient safety is to thoroughly study and analyze a UE to find contributing factors and root causes, and formulate corrective measures. A number of analysis tools are brought together under the term root cause analysis (RCA), for example, fishbone diagrams, cause-effect charts, and “five whys.” A comprehensive RCA-method that is frequently used in Dutch healthcare facilities is the Prevention and Recovery System for Monitoring and Analysis (PRISMA)-method. This method examines the relative contributions of latent factors (technical and organizational root causes), active failures (human behavior-related root causes), and other factors (patient-related and other root causes). The main goal of the PRISMA-method is to build a quantitative database of UEs and process deviations, to facilitate the development and evaluation of system-based preventive strategies.

The PRISMA-method consists of 3 important pillars: incident description, root cause classification, and the translation to structural measures. First, the incident is described using a causal tree that provides a visual interpretation of the chronological chain of critical activities and decisions leading to an UE. At the top of the
tree, the top event, a brief description of the event, is formulated, which is the start of the analysis. By continuously asking “why” the event occurred, all direct causes are identified. When no further objective causes can be identified, the last indirect cause is considered as the root cause. All root causes are logically related to each other and incorporated into the tree (Appendix 1, http://links.lww.com/JPS/A433, and Fig. 1). After compiling the causal tree, the root causes, which form the bottom layer of the causal tree, are classified using the Eindhoven Classification Model (ECM; Appendix 2, http://links.lww.com/JPS/A434). The ECM distinguishes 4 main categories, namely, technical, organizational, human, and other factors, including patient-related factors and unclassifiable factors. These main categories can be subdivided into 20 subcategories. After root cause classification, information on the event and the identified root causes are placed in a database. Prevention strategies can then be directed at these most frequently occurring (combinations of) root causes.

Although this method is accepted in healthcare facilities and adopted by the World Health Organization,11,12 little is known about the application (type of event, study setting) and implementation (data collection, study size, execution of analysis) of the PRISMA-method for analyzing UEs in healthcare facilities. This knowledge gap hinders our ability to learn from insights the PRISMA-method has already provided, and makes it difficult to further improve the application of PRISMA to prevent UEs in healthcare facilities worldwide.

The aims of this study are therefore to give a systematic literature overview of the use of the PRISMA-method in healthcare facilities and to assess the overall insights the PRISMA-method offers to better understand the causes of the UEs. We are particularly interested in the application and implementation of the PRISMA-method and the most common root causes contributing to UEs in healthcare facilities. Moreover, we intend to provide recommendations for further improvement of the method.

METHODS

To achieve this aim, we performed a systematic literature review. Two researchers (B.E.J.M.D., R.O.) used the structure and recommendations from the Preferred Reporting Items for Systematic Review and Meta-Analysis checklist (PRISMA statement, www.prisma-statement.org) as a guideline to conduct the literature search.13 We performed systematic searches to find all relevant publications about the “Prevention and Recovery Information System for Monitoring and Analysis” (PRISMA) in the analysis of medical incidents and events searching the bibliographic databases PubMed, EMBASE.com, CINAHL (via EBSCO), and The Cochrane Library (via Wiley) from inception to February 26, 2020. Search terms included controlled terms (e.g., MeSH in PubMed and Emtree in Embase) as well as free-text terms. We used free-text terms only in The Cochrane Library. Search terms expressing “PRISMA” were used in combination with terms comprising “medical events.” In the search, all languages were accepted.

The references of the identified articles were searched for relevant publications. The full search strategies for all databases can be found in Appendix 3, http://links.lww.com/JPS/A435.

Subsequently, all search results were uploaded to Endnote, to facilitate organizational and removal of duplicates. Second, results (duplicates excluded) were imported in Rayyan, an Internet-based software program that facilitates collaboration among reviewers during the study selection process. Two reviewers (B.E.J.M.D., M.B.) independently screened titles and abstracts and assessed these for eligibility. Ineligible records were excluded (reasons were recorded), and full texts of the remaining studies were assessed independently for eligibility. Results of these assessments were compared, and in case of uncertainty about inclusion, the articles were discussed with a third researcher (H.M.). English- or Dutch-written articles were included in the study if they (1) published original findings using PRISMA analysis in healthcare facilities, (2) classified the root causes using the ECM, and/or (3) used an observational study design, which included prospective and retrospective cohort studies, case-control studies, cross-sectional studies, case series, and case reports. Studies that applied PRISMA outside of healthcare facilities, articles only focusing on elaborating PRISMA, and articles using the ECM in combination with an unknown RCA or RCA other than PRISMA were excluded. When there was no full text available, articles were excluded. There were no restrictions by publication year, type of setting within healthcare facilities, or follow-up.

![FIGURE 1. Example of root causal tree.](https://example.com/root_causal_tree.png)
Quality Appraisal and Data Synthesis

Two authors (B.E.J.M.D., M.B.) independently assessed the quality of included studies using the National Heart, Lung, and Blood Institute Quality Assessment Tool for Observational Cohort and Cross-Sectional Studies to ensure an objective evaluation process. This tool includes 14 questions assessing the internal validity of the studies resulting in an overall quality rating of "good," "fair," or "poor." The tool considers information such as specification of study population, participation rate, measurement of exposure(s) of interest, time frame, and blinding of outcome assessors.

Data Extraction

The following data were extracted from each included study: title, first author, publication year, language, aim, design, study period, study population, hospital setting, department setting, data collection, PRISMA analysis, intervention, type of ECM used, follow-up, outcomes, limitations, future research, and implications. In addition, specific data regarding characteristics and application of the PRISMA-method were extracted, such as number of researchers conducting the PRISMA, training/experience in PRISMA of researchers, ECM-classification profiles, and types of suggested measures for improvement. During the whole process, uncertainties were resolved through team discussions (B.E.J.M.D., M.B., and H.M.) and consensus.

RESULTS

Study Selection

A total of 3926 studies including duplicates were retrieved from all databases after the initial search. After removal of 1153 duplicates, 2773 unique studies remained for the title and abstract review. Next, 2150 studies were excluded after screening titles and abstracts. Main reasons for exclusion were "Descriptive," "Use of causal tree, no PRISMA," "Not in Healthcare facility," "No RCA," and "Other type of RCA." From the resulting 623 articles, there was no access to a full-text version of 72 of the studies, and 17 articles were poster or congress abstracts. Another 509 studies were excluded after full-text evaluation because the study applied a different type of analysis from the PRISMA-method. Eventually, 25 studies were included in the final review. Figure 2 outlines the article selection process.

Quality Assessment

Overall, 3 studies were rated as "poor," 8 studies as "fair," and 14 studies as "good" quality (Table 1). The articles with a "poor" quality rating lacked information concerning research question, study design, study population, and description of implication of the PRISMA-method. Because of the aim of the study was to give a comprehensive overview of the application of PRISMA, we also included these "poor"-rated studies in our sample. Overall agreement on methodological quality score between researchers was 88%.

Study Characteristics

The characteristics of the included articles are presented in Table 2. The publication year of the 25 articles ranged from 1998 to 2019, and most of the studies (n = 20) took place in the Netherlands. The sample size ranged from a single case study to a multicenter study analyzing 2028 UEs. All studies were presented as observational cohort studies, with a retrospective (n = 23) and prospective (n = 1) design. Van Dulmen et al combined a retrospective patient record review with prospective event reporting. The design was not clearly mentioned in 5 studies but interpreted as observational by the researchers.

Study Population

Variations in definitions, type of event, and measurement of the study population were evident. In 6 studies, events without deviations were analyzed. The study objects in these articles were event reports in transfusion medicine, readmissions, unplanned intensive care department admissions, and an emergency department (ED) length of stay >6 hours.
In 19 studies, events with deviations were analyzed, including incidents, 16,24,26,30–32,34,36 UEs, 22,27,28,33,35 and adverse events. 23,29,37 In almost all (n = 7) of the studies analyzing incidents, a clear definition of “an incident” was mentioned. For example: “A critical incident was defined as an event that had actual or potential harmful effects on the outcome of the management of a patient or group.” 33 Unintended events were defined as “a broader group of events—including near misses—that do not necessarily result in patient harm.” 22,27,28,32,35 The adverse events in the study by Zegers et al. 37 and Merten et al. 23 were defined as “unintended injuries among patients that results in disability, death or prolonged hospital stay, and is caused by healthcare management.”

### TABLE 1. NHLBI, NIH: National Heart, Lung, and Blood Institute Quality Assessment Tool for Observational Cohort and Cross-Sectional Studies 13

| Reference | Author | Score | Missing Information in Description of: |
|-----------|--------|-------|-----------------------------------------|
| 12        | Battles and Shea | Fair | Study population Exposure measures and assessment |
| 14        | Cooksley et al | Fair | Study population Exposure measures and assessment |
| 15        | Driesen et al | Good | |
| 16        | Van Dulmen et al | Fair | Sample size justification Exposure measures and assessment |
| 17        | Fluitman et al | Good | |
| 18        | Van Galen et al | Good | |
| 19        | Van Galen et al | Good | |
| 20        | Kaplan et al | Poor | Research question Participation rate Time frame |
| 21        | Klemt-Kropp et al | Poor | Research question Exposure measures and assessment Outcome measures |
| 22        | Lubberding et al | Good | |
| 23        | Merten et al | Good | |
| 24        | Van Noord et al | Good | |
| 25        | Rodrigues et al | Poor | Research question Participation rate Time frame Exposure measures and assessment Outcome measures |
| 26        | Sasaki et al | Fair | Study population Blinding of outcome assessors |
| 27        | Smits et al | Good | |
| 28        | Smits et al | Good | |
| 29        | Smits et al | Fair | Study population Groups recruited from the same population |
| 30        | Snijders et al | Good | |
| 31        | Snijders et al | Fair | Study population |
| 32        | Thomas and Mackway-Jones | Good | |
| 33        | De Vries et al | Fair | Sample size Blinding of outcome assessors |
| 34        | Wagner et al | Good | |
| 35        | Van Wagendonk et al | Good | |
| 36        | Wubben et al | Fair | Sample size justification Time frame |
| 37        | Zegers et al | Good | |

### Study Setting

The majority of the studies (n = 23) were conducted in a hospital setting. In the 2 other studies, patients were included from 3 allied healthcare disciplines (physical therapy, occupational therapy, and Cesar-Mensendieck exercise therapy) 39 and from a blood and transfusion center. 22 The majority of studies (n = 16) limited the study population to one department or specialty, whereas others included cases from various hospital departments. 19,29,35,36–39

### Data Collection and Implementation of PRISMA

The collected data used for the PRISMA analysis were gathered using different sources. In 9 studies, the information for the PRISMA analysis was based on patient files, 14–16,23,24,29,35 and in 5 studies, the data were collected from a report form. 20,26,30,31,33 In 3 studies, the report form was based on voluntary, nonpunitive reporting. 26,30,31 In some studies, the data were collected from a combination of sources; 6 studies used a combination of a report form and an interview. 19,22,27,28,34,35 and 2 studies used a combination of a report form, an interview, and a patient record. 22,26 In 3 studies, the sources underlying the data used for analysis are unknown. 12,21,25

Researchers received a PRISMA-training in almost half (n = 12) of the included studies. 15,17,18,20,22,24,27,30,31,34,35 This training ranged from a 3-hour workshop 20 to a 2-day course. 30,31 In the majority of studies (n = 14), the PRISMA analysis was performed by 2 persons or a team (independent from each other). This team could consist of “analysts,” 15,17,18,22,34 multidisciplinary safety commission consisting of at least 1 physician and 3 nurses, 36,37 multiple doctors, 12,23,26,33,37 or physical therapists. 16 The mean root causes per event were 2.5 in the articles where the analysis was performed by multiple persons (range, 1.4–6.3). In 8 studies, the PRISMA analysis was conducted by a single person 19,20,24,28,29,32,35,36; here the mean root causes per event were 1.8 (range, 1.0–2.9). In 3 articles, it is unknown who performed the PRISMA. 14,20,22 Furthermore, interobserver agreement on root causes and ECM classification was studied in 10 of the studies, 16,24,27,33,37 with scores ranging from “moderate” to “good.”

### Root Causes and ECM

Human-healthcare worker–related root causes were the most frequently found ECM-classified root causes in the majority of the studies (n = 14). 16,18,20,22,24,26,30,31,33–37 In other studies, the main category of root causes was organizational factors (n = 5). 15,25,21,26,32 The adverse events in the study by Zegers et al. 37 and Merten et al. 23 were defined as “unintended injuries among patients that results in disability, death or prolonged hospital stay, and is caused by healthcare management.”

In the 14 studies where human–healthcare–related root causes were most prominent, the majority of the studies analyzed incidents. 16,25,30,34,36 Furthermore, the analysis was limited to one department, 16,20,22,24,26,33,35–37 and the PRISMA was done by multiple persons. 16,18,22,23,30,32,34,37 in most of the cases. Six
| Reference | Author | Country, Year of Publication | Type of Event | Setting Disorder | Data Collection | No. Root Causes | Root Cause Per Event | ECM Modification | Most Reported ECM Classification |
|-----------|--------|-----------------------------|---------------|-----------------|----------------|-----------------|---------------------|-----------------|----------------------------------|
| 12        | Battles and Shea | United States and the Netherlands, 2001 | Error (n = 3) | ED and IC | NR | 19 | 6.3 | Original | Organizational, n = 11 (59%) |
| 14        | Cooksley et al | United Kingdom, 2015 | Readmissions (event) (n = 50) | Oncology | Patient file | NR | NR | Original | Patient, n = 50 (100%) disease related |
| 15        | Driesen et al | The Netherlands, 2018 | Increased ED-LOS (event; n = 74) | ED | Patient file | 276 | 3.3 | Modified • Disease-related factor added | Organizational, n = 216 (78%) |
| 16        | Van Dulmen et al | The Netherlands, 2011 | Incidents (n = 18) | Physiotherapy, occupational therapy, and cesar-mensendieck therapy | Patient file | 55 | 3.1 | Original | Disease related, n = 40 (73%) |
| 17        | Fluitman et al | The Netherlands, 2016 | Readmissions (event) (n = 50) | Internal medicine | Patient file | 100 | 2 | Modified • Disease-related factor added | Disease related, n = 46 (46%) |
| 18        | Van Galen et al | The Netherlands, 2016 | Unplanned admissions (event; n = 49) | IC | Patient file | 155 | 3.2 | Modified • Disease-related factor added | Human, n = 71 (46%) |
| 19        | Van Galen et al | The Netherlands, 2018 | Readmissions (event; n = 237) | Surgery | Report form, interview | 271 | 1.1 | Modified • Disease-related factor added | Disease related, n = 218 (87%) |
| 20        | Kaplan et al | United States, 1998 | Events/process (n = 506) | Blood center Transfusion center | Report form | 1238 | 2.9 | Modified • Content adopted for MERS-TM, no change in (sub)categories | Human, n = 569 (46%) |
| 21        | Klemt-Kropp et al | The Netherlands, 2011 | Error (n = 1) | MDL | NR | 8 | 8 | Original | Organizational, n = 5 (63%) |
| 22        | Lubberding et al | The Netherlands, 2011 | UEs (n = 625) | Internal medicine | Report form, interview | 920 | 1.5 | Original | Human, n = 706 (77%) |
| 23        | Merten et al | The Netherlands, 2013 | Adverse events (n = 663) | All departments, excluding psychiatry, obstetrics, children<1 y | Patient file | 1278 | 1.9 | Modified • Human violation added • Human-other added | Human, n = 808 (63%) |
| 24        | Van Noord et al | The Netherlands, 2010 | Malpractice claims (incident; n = 50) | ED | Patient file | 114 | 2.3 | Modified • Organizational x-ray results (OR) added • Organizational no contact with supervisor (OS) added | Human, n = 76 (67%) |
|   | Study Authors | Country, Year | Type (n =) | Department(s) | Method | n | Original/Modified | Case Details |
|---|---------------|---------------|------------|---------------|--------|---|-----------------|-------------|
| 25 | Rodrigues et al | Portugal, 2011 | Error (n = 79) | Radiology | NR | NR | NR | Modified |
|     |               |               |            |               |        |    |     | Content adopted for medical imaging field, no change in (sub) categories |
| 26 | Sasaki et al   | Japan, 2018   | Incidents (n = 95) | Allergology | Report form | 220 | 2.3 | Modified |
|     |               |               |            |               |        |    |     | Home/facility category instead of organizational home facility, n = 74 (34%) |
| 27 | Smits et al    | The Netherlands, 2009 | UEs (n = 2028) | ED, internal medicine, surgery | Report form interview | 3015 | 1.4 | Original |
|     |               |               |            |               |        |    |     | Human, n = 507 (60%) |
| 28 | Smits et al    | The Netherlands, 2009 | UEs (n = 522) | ED | Report form interview | 845 | 1.6 | Original |
|     |               |               |            |               |        |    |     | Human, n = 293 (55%) |
| 29 | Smits et al    | The Netherlands, 2019 | Adverse events (n = 571) | All departments, excluding psychiatry, obstetrics, children<1 y | Patient file | 588 | 1.0 | Modified |
|     |               |               |            |               |        |    |     | Violation added, Human skills added, Other added |
| 30 | Snijders et al | The Netherlands, 2008 | Incidents (n = 981) | NICU and PICU | Report form | 2313 | 2.4 | Original |
|     |               |               |            |               |        |    |     | Human, n = 1480 (64%) |
| 31 | Snijders et al | The Netherlands, 2010 | Incidents (n = 533) | NICU and PICU | Report form | 1233 | 2.3 | Original |
|     |               |               |            |               |        |    |     | Human, n = 293 (55%) |
| 32 | Thomas and Mackway-Jones | United Kingdom, 2007 | Incidents (n = 349) | ED | Report form, patient file, interview | 852 | 2.4 | Modified |
|     |               |               |            |               |        |    |     | Technical-other instead of Technical-material, Human execution instead of human intervention, Multiple patients added |
| 33 | De Vries et al | The Netherlands, 2015 | UE (n = 41) | Urology | Report form | NR | NR | Original |
|     |               |               |            |               |        |    |     | Human, estimated 65% |
| 34 | Wagner et al   | The Netherlands, 2016 | Incidents (n = 2028) | ED, Internal medicine, Surgery | Report form interview | 3015 | 1.5 | Modified |
|     |               |               |            |               |        |    |     | Patient related and other combined in one category |
| 35 | Van Wagendonk et al | The Netherlands, 2010 | UEs (n = 881) | Surgery | Report form interview | 1250 | 1.4 | Original |
|     |               |               |            |               |        |    |     | Human, n = 904 (73%) |
| 36 | Wubben et al   | The Netherlands, 2010 | Incidents (n = 15) | Operation room | Report form, patient file, interview | 40 | 2.7 | Original |
|     |               |               |            |               |        |    |     | Human, n = 16 (40%) |
| 37 | Zegers et al   | The Netherlands, 2011 | Adverse events (n = 367) | Surgery | Patient file | 653 | 1.8 | Original |
|     |               |               |            |               |        |    |     | Human, n = 426 (66%) |

GLD, gastroenterology and liver disorders; IC, intensive care department; MERS-TM, medical event reporting system for transfusion medicine; NICU, neonatal intensive care unit; NR, not reported; PICU, pediatric intensive care unit.
studies used a combination of data sources for the PRISMA-method. Patient interviews were used in 5 studies. The number of root causes per event ranged from 1.45 to 3.2.

In 13 studies, the original ECM was used, and in 12 studies, the ECM was modified because the current ECM did not sufficiently cover the observed types of failures (Table 2). In 4 studies, a new DF category was added. The disease-related root cause was first described by Fluitman et al in 2016 and defined as “failures related to the natural progress of disease which are beyond control of patients, their carers and staff.” In these 4 studies, the DF refers to readmissions and unplanned admissions of patients. The authors assumed that progression of disease would be identified as root cause in many readmissions without other factors contributing. Other modifications included adaptation of the content of categories to the study population and addition of new subcategories, for example, human violation and organizational x-ray results.

**DISCUSSION**

This systematic review identified 25 original studies using the PRISMA-method for RCA. The results show that the PRISMA-method is applicable in a wide variety of healthcare settings. Most studies analyzed UEs, predominantly incidents. Furthermore, most studies were limited to one department or specialty and used patient files or report forms for data collection, and the analysis was done by multiple persons. Overall, human-healthcare worker-related root causes were the most common root causes, followed by organizational factor-related root causes.

Based on descriptions of various RCA methods in the literature and their limitations, the results of this review, we believe that PRISMA has some important features that make it one of the most valuable RCA methods. First, PRISMA is suitable for events varying in seriousness and complexity, whereas other methods are designed to investigate more serious adverse events. Another advantage of PRISMA medical is that it intends to build a database of multiple events to find reoccurring patterns of root causes. Based on these patterns, it might be possible to formulate more effective corrective measures. Because both active failures (human failures) and latent failures (technical and organizational failures) are discovered through PRISMA, the total profile of root causes provides a more realistic view of how the system is actually working. The standardized root cause classification improves the comparability of investigated events between healthcare facilities as well as in the identification of common problems and concerns that exist throughout the medical field.

Most studies reported events where the reporting system and analysis was limited to a single department or service within the hospital. Local, department-based reporting systems can help to get faster and more detailed insight into unit-specific safety issues. The assumption is that incidents differ between departments and that incidents reported and analyzed by the department where the incident happened will create a greater sense of urgency and willingness to change practice. Wagner et al. found significant differences in incident types and frequencies in root causes between emergency, surgery, and internal medicine departments, therefore recommending department-based reporting. A limitation of department-based analysis is that causes occurred in other departments are classified as external. It is possible that an external factor in fact had more influence on the system.

The number of root causes per event ranged from 1.45 to 3.2.

The patient file method for data collections was the most used, but this may not be the most accurate method. Technical and organizational causal factors are less often reported in patient files and may therefore be underestimated. An organizational factor could, for example, be a shortage of hospital staff. However, such causes are not typically written down in patient files and therefore remain invisible to the researchers. This is in line with our results; in the studies where the patient file was the only data source, the majority of the articles had a human factor as most common root cause. In contrast to other data sources, multiple studies suggested that the patient file or a report, an interview with those involved at a local level can lead to a deeper understanding. The interview depends on the recall of the reporters, and UEs should be discussed within a few days. This is in line with our results showing that when the data were based on interviews only, organizational root causes were most prominent.

On average, there were more root causes per event in the studies where the PRISMA analysis was done by multiple persons (2.5 versus 1.8). An explanation could be that multiple persons have different perspectives, especially when there is a multidisciplinary team including different healthcare workers, for example, doctors, nurses, or people without direct patient healthcare experience or contact, for example, experts in the field of human factors/ergonomics, clinical technologists, or PhD researchers. This could also contribute to finding more technical and organizational root causes. Furthermore, based on our practical experience, we believe it is important for the reliability of the PRISMA-method that the researchers are trained in PRISMA and that they have knowledge of the department and its processes. In this study, almost half of the articles mentioned that the researchers received training. Only in a small part of the studies the content and the duration of the training were mentioned, and therefore, it was difficult to substantiate this hypothesis. Overall, human-healthcare worker-related root causes were most prominent in the included studies. These root causes can be defined as “errors and violations committed by those in direct contact with the patient—system interface.”

Our finding that the root causes were mainly human-related is in line with other studies investigating (adverse) events that also found that causes related to the people involved, such as human behavior, cognitive biases, and poor communication, were frequently or dominantly reported. However, as mentioned before, the type of root cause can be dependent on the source of the data collection and the level at which the analysis was done.

**Strengths**

This is the first systematic literature review giving an overview of the use of the PRISMA-method in healthcare facilities. An experienced librarian (R.O.) assisted in the literature search, and the screening of the articles, the data collection, and the quality assessment were independently done by 2 researchers (B.E.J.M.D. and M.B.).

**Limitations**

First, 3 studies were rated “poor quality” because information on several study characteristics was not reported. However, we included these studies because the aim of our study is to provide an overview, and therefore, the lacking information does not bias our
results or discussion. Second, 20 of the 25 included studies were concerned with Dutch healthcare facilities. This may be the result of the popularity of the PRISMA analysis in the Netherlands, which in turn could be explained by the Dutch origin of the method and its founders.

**Recommendations**

To get a complete view of all contributing root causes of UEs, we recommend combining patient files and reports with interviews when conducting a PRISMA analysis, thereby striving for a small time lag between the occurrence and the assessment of the UE to decrease the likelihood of memory failure.35 In that case, involved healthcare providers can be interviewed to provide more information on organizational and technical factors that might have contributed to the UE. Preferably, the PRISMA should be performed by multiple researchers representing various disciplines, who should get training in making causal trees and classifying root causes, as training of researchers increases reliability.25 When using department-based analyzing, it is advisable to modify the ECM to gain more insight into the causes outside of the department, for example, the addition of the DRF.

**CONCLUSIONS**

This study outlines the current variety of the use of the PRISMA-method in healthcare facilities worldwide. Human-healthcare worker–related root causes were found to be the most commonly reported root causes, followed by organizational factor–related root causes. The selected studies showed that the PRISMA-method is a useful tool with added value for analyzing UEs in healthcare facilities. To improve the usability of PRISMA, we suggest combining information from patient files and reports with interviews, including multiple PRISMA-trained researchers representing various disciplines in an analysis, and modify the ECM if needed, to make care safer and more efficient by learning from UEs in healthcare processes.

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