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Delirium detection tools show varying completion rates and positive score rates when used at scale in routine care

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











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REVIEW ARTICLE

Delirium detection tools show varying completion rates and positive score rates when used at scale in routine practice in general hospital settings: A systematic review

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Abstract

Background: Multiple short delirium detection tools have been validated in research studies and implemented in routine care, but there has been little study of these tools in real-world conditions. This systematic review synthesized literature reporting completion rates and/or delirium positive score rates of detection tools in large clinical populations in general hospital settings.

Methods: PROSPERO (CRD42022385166).

Medline, Embase, PsycINFO, CINAHL, and gray literature were searched from 1980 to December 31, 2022. Included studies or audit reports used a validated delirium detection tool performed directly with the patient as part of routine care in large clinical populations ($n \geq 1000$) within a general acute hospital setting. Narrative synthesis was performed.

Results: Twenty-two research studies and four audit reports were included. Tools used alone or in combination were the Confusion Assessment Method (CAM), 4 'A's Test (4AT), Delirium Observation Screening Scale (DOSS), Brief CAM (bCAM), Nursing Delirium Screening Scale (NuDESC), and Intensive Care Delirium Screening Checklist (ICDSC). Populations and settings varied and tools were used at different stages and frequencies in the patient journey, including on admission only; inpatient, daily or more frequently; on admission and as inpatient; inpatient post-operatively. Tool completion rates ranged from 19% to 100%. Admission positive score rates ranged from: CAM 8%–51%; 4AT 13%–20%. Inpatient positive score rates ranged from: CAM 2%–20%, DOSS

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6%–42%, and NuDESC 5–13%. Postoperative positive score rates were 21% and 28% (4AT). All but two studies had moderate–high risk of bias.

Conclusions: This systematic review of delirium detection tool implementation in large acute patient populations found clinically important variability in tool completion rates, and in delirium positive score rates relative to expected delirium prevalence. This study highlights a need for greater reporting and analysis of relevant healthcare systems data. This is vital to advance understanding of effective delirium detection in routine care.

KEYWORDS

delirium, detection, geriatric assessment, hospitals, older people, routine data, systematic review

INTRODUCTION

Delirium is an acute, severe neuropsychiatric syndrome with an overall occurrence of 23% in adult medical hospital inpatients.¹ It is associated with adverse outcomes, including higher mortality, new institutionalization, and higher risk of future dementia.^{2–4}

Delirium detection is essential to deliver effective care. A delirium diagnosis prompts access to specific management pathways and facilitates accurate communication with patients and relatives. Several national guidelines and care standards recommend routine delirium detection using validated tools.^{5–10} Yet delirium remains under-detected in routine care, with studies comparing clinically-documented rates of delirium with reference-standard assessments demonstrating widespread under-detection.^{11–13} Hospital discharge administrative data also demonstrate under-detection and under-coding.¹⁴

Multiple delirium detection tools have been developed with high sensitivity and specificity in diagnostic test accuracy studies.^{15,16} Yet good tool performance in validation studies, where assessments have generally been performed by research teams, does not guarantee that tools will function well in usual care. Indeed, some studies of real-world practice suggest that delirium tools are commonly not completed, and that tools may produce lower delirium positive score rates than expected.^{12,17–20}

Suboptimal completion rates could indicate difficulties in performing the particular tool by clinical staff, or the tool not being attempted for various reasons including culture, staff awareness, stage of tool implementation within the given healthcare system, timepoint in the patient journey the tool is used, and so forth.^{17,21–23} Considering completed tools, if the positive score rates are much lower than the expected delirium prevalence in the target population, this suggests that the sensitivity of the tool in practice might be lower than the sensitivity shown in research settings.^{12,18–20}

Key points

- We found 22 research studies and 4 audit reports examining 6 different validated delirium detection tools in acute care
- Tool completion and positive score rates varied widely in diverse populations
- Some studies reported positive score rates far below expected for the setting, despite high tool completion

Why does this paper matter?

Delirium underdetection is of immense concern. This study found variable disparities between tool performance in diagnostic test accuracy studies and in performance in real-world practice. Analyzing and reporting routine healthcare systems data is vital for effective delirium detection at scale.

Lower real-world sensitivity could reflect multiple factors including staff training, clinical time constraints, timepoint or frequency of testing in the patient journey, or the operational method of tool completion. Evidence on these two metrics of *tool completion rates* and *delirium positive score rates* is crucial for clinicians and policymakers aiming to improve delirium detection because it can inform the choice of tool appropriate for the setting and clinical population.

We performed a systematic review of large-scale ($N \geq 1000$) studies and audit reports examining delirium detection tool metrics when tools are used in routine clinical practice in general hospital settings (excluding critical care). We report: (a) tool completion rate (where available), and (b) delirium positive score rate. We report contextual factors including the clinical population, stage

of the patient journey, the staff discipline(s) using the tool and, where reported, information on factors that might affect tool completion.

METHODS

The systematic review is reported according to Preferred Reporting of Items in Systematic Reviews and Meta-analysis (PRISMA) guidelines (Supplementary Table 1).²⁴ The protocol was prospectively registered with PROSPERO (CRD42022385166).

Search strategy and data sources

The search strategy was developed with academic librarian support and included three main concepts: “delirium”, “assess*” (and synonyms), and “clinical practice” (and synonyms). Search strategies were developed for Medline, Embase, and PsycINFO (all via Ovid), and CINAHL (via EBSCO) (Supplementary Table 2). All databases were searched from 1980 when delirium was first coded in the Diagnostic and Statistical Manual of Mental Disorders (DSM) III,²⁵ to December 31, 2022. Study eligibility criteria are below.

Inclusion and exclusion criteria

Studies using any peer-reviewed methodology were included if they included adults aged ≥ 18 years admitted to an acute hospital setting, described the use of a specified delirium assessment tool in study participants, reported tool use in an unselected population of ≥ 1000 patients, and were published in English, or translatable to English using online tools. No restrictions were placed on the study type or aims, or geographical location.

Studies of patients admitted to critical care or community settings were excluded, as were studies where the assessment was not performed directly with the patient (for example, studies using Electronic Health Record (EHR) data to develop a prediction model retrospectively). Studies only reporting on tools for rating delirium severity or resolution (rather than detection) or studies where tools were not used in routine clinical practice were excluded, as were systematic reviews, meta-analyses, abstracts, letters to editors, and opinion pieces.

Title and abstract screening

All searches were performed on December 31, 2022, with study de-duplication using Covidence software (Veritas

Health Innovation, Melbourne, Australia), with manual checking by RSP of software-identified duplicates. Title and abstract screening were independently performed in Covidence by two reviewers (RSP and CS). Discrepancies between reviewers at the title/abstract screening stage were resolved through discussion, and articles with persisting discrepancies were included in a full-text review.

Full-text review

Full-text review was performed independently in Covidence by two reviewers (RSP and AAng). Excluded full-text articles were assigned a single primary reason for exclusion (Figure 1). Disagreement between reviewers was resolved through discussion involving an additional senior reviewer (AMJM) for persisting disagreement.

For any identified abstracts (e.g., published conference abstracts), we searched the databases above for a related full-text publication. We used forward citation on all included studies to identify relevant peer-reviewed publications that may have been missed in the search algorithm.

We scoped gray literature to identify additional relevant publications, including audit reports, conference abstracts and papers not picked up in the systematic search.

Risk of bias

Two reviewers (RSP and AAng) independently assessed studies for risk of bias (RoB) using an adapted version of the ROBINS-E quality assessment tool for observational studies.²⁶ Discrepancies were resolved through discussion, with an additional reviewer available if required (AMJM). Studies were assessed as having low RoB, some concerns, high RoB, or non-applicable across seven domains (confounding, measurement of exposure, participant selection, post-exposure interventions, missing data, measurement of the outcome, and selection of the reported result), with an overall judgment assigned.

Data extraction and synthesis

Two independent reviewers (RSP and AAng) extracted data for each study on the assessment tool used, the timepoint(s) and frequency of assessment in the patient journey, and whether the study reported on the entire eligible population based on the inclusion criteria or only on patients with a completed delirium assessment. For studies reporting on the entire eligible population, we

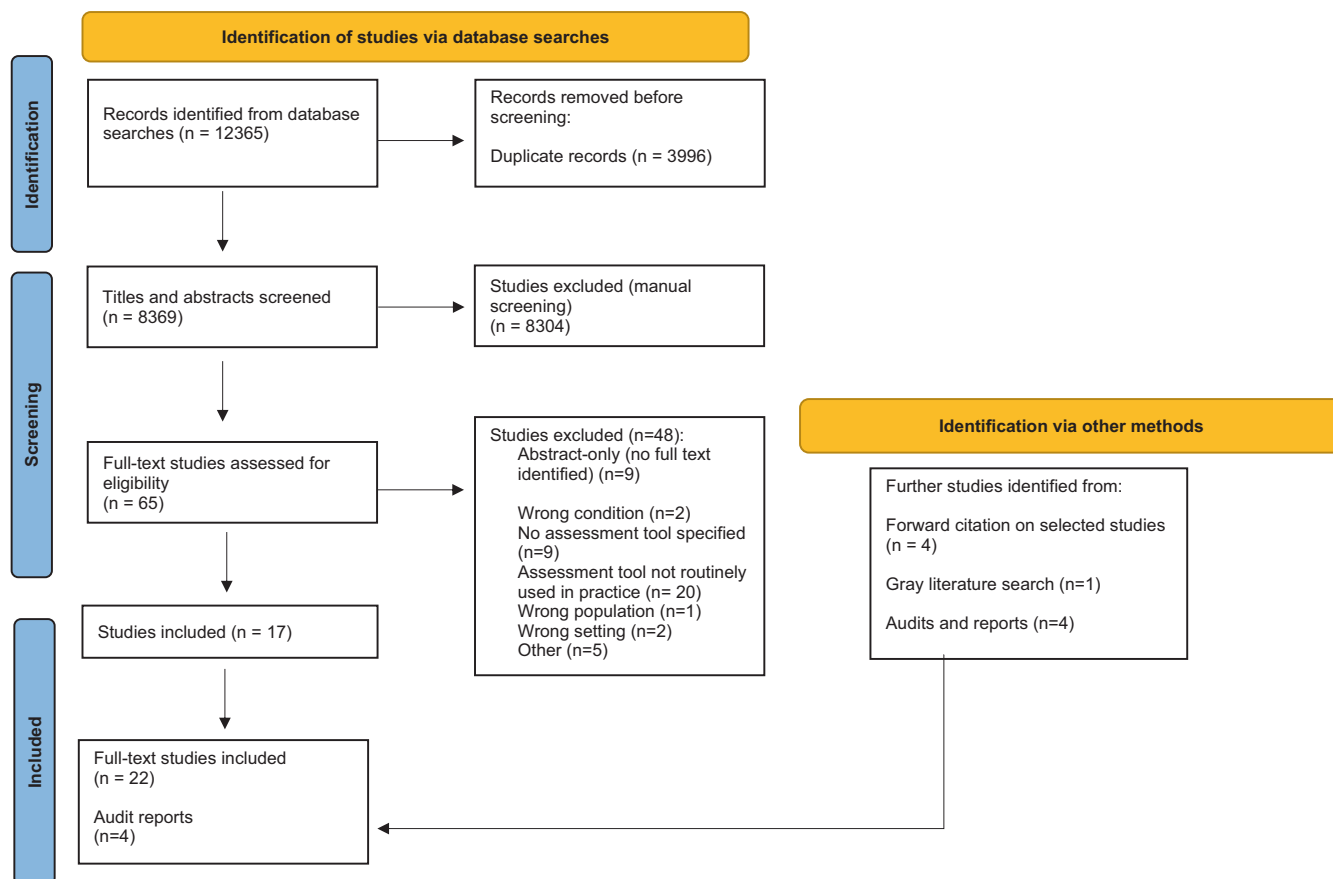


FIGURE 1 Study inclusion diagram.

extracted or calculated (if not explicitly reported) the proportion of eligible patients assessed using the tool (tool completion rate). We also extracted or calculated (if not explicitly reported) the positive score rate. If calculated, this was the number of patients with a positive score or number of positively scored observations divided by the total number of patients assessed or total number of observations.

For studies reporting on patients in both acute hospitals and excluded settings (e.g., critical care and rehabilitation facilities), we extracted data only for patients admitted to acute hospitals where possible. It is specified if the study did not report separately for the different settings.

Narrative data synthesis was performed due to anticipated heterogeneity in study populations and methodologies. Results are reported in tables and summary figures (Integrated Development for R. Rstudio, PBC, Boston, MA, USA).

RESULTS

Search results

After deduplication, 8369 articles were screened and 65 further evaluated. Nine articles were conference

abstracts for which no related full-text study could be identified (Supplementary Table 3). Of the 56 full-text studies screened, 17 (30%) met the inclusion criteria. There was title-abstract agreement between reviewers in 8271/ 8369 (99%) of cases and 50/56 (89%) of cases at full-text review. Studies excluded after full-text review are summarized in Supplementary Table 4. No studies were excluded for language reasons. A further four papers were identified using forward citation on included full-text studies, and one paper during gray literature search.

Study characteristics and patient populations

Twenty-two studies were included, representing over 500,000 patients (Table 1). Study denominators were either number of patients or total number of observations, precluding overall summary statistics. Studies were conducted in the United States (7), United Kingdom (5), Switzerland (5), Brazil (2), Colombia (1), New Zealand (1) and Italy (1). The average age of study participants ranged from 55–86 years. Fifteen (68%) studies only included adults at least ≥ 60 years; one included patients ≥ 50 years and six included all patients ≥ 18 years. Most

TABLE 1 Summary of all included studies and patient populations.

Author, year	Country	Study design	Type of hospital/setting	Patient population	Total number of patients in study	Study inclusion criteria (including age and patient consent)	Study exclusion criteria	Age (mean (SD))/median (IQR))	Sex F (%)
Alhairadi et al., 2021	NZ	Retrospective analysis of routine EHR data	Acute hospitals, two sites	Acute medical and geriatrics admissions	7799	≥75 years	None specified	84 (6)	58
Anand et al., 2022	UK	Retrospective analysis of routine EHR data	Acute hospitals, two sites	Acute medical and surgical admissions (Salford only)	82,770 admissions (31,266 patients)	≥65 years	Not resident in Scotland (Lothian data only)	79 (8)	54
Avelino-Silva et al., 2017	Brazil	Prospective cohort	Acute hospital	Geriatric ward ^a	1409	≥60 years, acute illness, patient consent	End of life care, incomplete data, LOS < 48 hours	80	61
Corradi et al., 2016	US	Retrospective analysis of routine EHR data	Acute hospital	All inpatients	88,206 encounters	≥18 years with CAM scores between Sept 2012 and June 2015	No CAM score	NA	NA
Di Bari et al., 2022	Italy	Cross-sectional cohort	Academic tertiary and community hospitals	Patients admitted via the ED	3358	≥75 years	Out of area, ophthalmological problems, software issue	83	56
Dulin et al., 2022	US	Retrospective cohort	Academic tertiary hospital	Patients admitted from home	93,388 admissions	≥18 years; hospitalized through office, clinic or ED; discharged alive	Admitted not from home, admitted to ICU, died pre-discharge	57	48
Fuchs et al., 2020	Switzerland	Prospective cohort	Acute university hospital	Discharged from acute medical and surgical units	10,261	≥65 years	LOS < 1 day, missing data	77	41
Garcez et al., 2020	Brazil	Retrospective cohort	Tertiary university hospital	Geriatric ward ^a	1554	≥65 years; nonelective; has admission arousal assessment	Discharged or died <48 h	81	61
Han et al., 2021	UK	Prospective cohort	Acute hospital	Patients with an acute hip fracture	1082	≥60 years	None specified	84 (9)	73
LaHue et al., 2021	US	Retrospective cohort	Acute hospital	Admitted to specified medical and surgical wards	22,708	≥50 years, hospitalized >1 day	Not admitted to and discharged from same unit	67 (11)	48

(Continues)

TABLE 1 (Continued)

Author, year	Country	Study design	Type of hospital/setting	Patient population	Total number of patients in study	Study inclusion criteria (including age and patient consent)	Study exclusion criteria	Age (mean (SD))/median (IQR))	Sex F (%)
Lee et al., 2019	US	Retrospective cohort	Academic medical centre	All inpatients	12,082 episodes (9017 unique patients)	≥65 years, admitted via ED, at least one DOSS score, no missing data	None specified	76	50
Lee et al., 2022	US	Retrospective cohort	Academic medical centre	Patients admitted from ED	7927	≥65 years, at least one DOSS score	None specified	76	52
Marquetand et al., 2021	Switzerland	Pragmatic prospective cohort, secondary analysis	Large health care system	Acute medical and acute surgical admissions	3076	≥80 years	LOS <24 h, missing data, unknown division, admission or discharge destination	86	51
Matharu et al., 2022	UK	Prospective cohort	Acute hospitals, multi-site	Patients with an acute hip fracture	135,685	≥60 years	Missing data for ≥1 covariate	83 (9)	71
Peralta-Cuervo et al., 2021	Colombia	Retrospective observational	Acute hospital	Acute medical admissions	1599	≥75 years or ≥60 years with risk factor	No CAM score	86 (9)	57
Reynish et al., 2017	UK	Prospective cohort	Acute hospital	Acute medical admissions ^a	10,014	≥65 years of age	None specified	79	57
Rohatgi et al., 2019	US	Retrospective cohort study	Acute hospital	Discharged from acute medical and surgical units	105,455 encounters	≥18 years, admitted to any medical-surgical unit ≥24 h	>1 day in ICU, in short-stay areas, or in inpatient psychiatry unit	60 (46–71)	52
Schubert et al., 2018	Switzerland	Prospective cohort	Acute university hospital	Patients discharged from acute medical and surgical units	29,278	≥18 years, discharged Jan –Dec 2014	LOS <1 day	55 (19)	52
Spiller et al., 2022	Switzerland	Prospective cohort	Acute hospital	All hospitalized patients	29,967	≥65 years or <65 years with “delirious symptoms” or cognitive impairment	Missing screening data	71 (12)	41
Tyas et al., 2021	UK	Retrospective analysis of routine data	Acute hospitals, multi-site	Patients with acute hip fracture	21,274 (2020) 22,098 (2019)	≥60 years	None specified	NA	NA

TABLE 1 (Continued)

Author, year	Country	Study design	Type of hospital/setting	Patient population	Total number of patients in study	Study inclusion criteria (including age and patient consent)	Study exclusion criteria	Age (mean (SD))/median (IQR)) Sex F (%)
Wong et al., 2018	US	Retrospective cohort	Academic medical centre	Acute medical admissions	18,223	≥18 years discharged Jan 1 2016–Nov 30 2017, completed assessment	Admitted with delirium, altered mental status, or requiring ICU admission	35.9% >64 years 52
Zipser et al., 2022	Switzerland	Pragmatic prospective cohort	Large health care system	Acute medical and surgical wards	17,158 (surgical) 8214 (medical)	≥18 years old, stay at least 24 h	Missing data, admission/ discharge not obtained	56 (37–70) 52

Abbreviations: CAM, Confusion Assessment Method, DOSS, Delirium Observation Screening Scale, ED, Emergency Department, ICU, Intensive Care Unit, LOS, length of stay; EHR, electronic health record.
^aAssessments performed by specialist trained nurses.

were observational cohort studies. Four studies self-described as retrospective analyses of prospectively collected routine EHR data. Five were multi-center. All were conducted in acutely hospitalized medical and/or surgical patients, with some in specific sub-populations, such as patients admitted to a single geriatrics unit (two studies) or hospitalized with an acute hip fracture (three studies).

Assessment tools used

Studies reported the use of six different validated assessment tools (Table 2 and Figure 2): the Confusion Assessment Method (CAM; six studies), brief-CAM (bCAM; one study), the 4 ‘A’s Test (4AT; six studies), the Delirium Observation Screening Scale only (DOSS; four studies), DOSS or Intensive Care Delirium Screening Checklist (ICDSC; one study), the Nursing Delirium Screening Scale (NuDESC; two studies) and, in two studies, the DOSS combined with the ICDSC or electronic Patient Assessment-Acute Care (ePA-AC), a DSM-based nursing construct including functional domains shown to be associated with delirium but not validated as a detection tool.²⁷ Studies using tool combinations did not always report the number of observations using each tool.

Timepoint(s) of Tool Use in Patient Journey

Delirium assessments were done at different timepoints during the patient journey. These can be classified into four categories (Table 2 and Supplementary Figure 1): admission assessment only (five studies^{21,22,28–30}); on admission and as inpatient, daily or more frequently (two studies^{17,31}); inpatient, daily or more frequently (11 studies^{12,18–20,32–38}); inpatient post-operatively (three studies^{39–41}). In one study, a single assessment was performed on ED attendance but only 31.5% of the study population were hospitalized.⁴²

Tool completion rate

Six studies reported data only on patients with a completed delirium assessment without providing a denominator (three CAM studies,^{18,28,29} one DOSS,³⁶ and both using the NuDESC^{19,33}) (Table 2). In the 16 studies reporting on the whole population eligible for assessment, the tool completion rate ranged from 19% (all inpatients ≥18 years, bCAM, although the center reported early data from a staged implementation process led by nursing champions and the completion rate was 63%

TABLE 2 Tool name, tool characteristics, tool completion rate (proportion of patients with a completed tool assessment), delirium positive score rate (proportion of assessed patients with a positive score or positively scored assessments) and proportion of patients *unable to assess* for all included studies.

Author, year	Tool name	Timepoint(s) of assessment(s) in patient journey	Frequency of assessment(s)	Time from admission to first assessment	Tool completion rate: No. of patients or assessments performed/no. reported as eligible for assessment (%)	Positive score rate: No. of patients with a positive score/ no. of patients or assessments (%)	Number (proportion) of patients classed as <i>unable to assess</i> (%)
Alhaidari et al., 2021	4AT	Admission only	One assessment only	Not specified	6492/7799 (83)	1154/7799 (15)	875/7799 (11%)* **"meaningful reason for omission"
Anand et al., 2022	4AT	Admission only	One assessment only	Not specified	52,965/ 82,770 (64)	10,685/52965 (20)	0
Avelino-Silva et al., 2017	CAM	Admission and Inpatient	On admission and Daily	<24 h of admission	1409/1522 (93)	379/1409 (27) (admission) 278/1409 (20) (inpatient)	Not specified
Corradi et al., 2016	CAM	Inpatient	3 × /day	Not specified	Only patients assessed using tool included	6926/88206 (8)	15,094/88206 encounters (17)
Di Bari et al., 2022	4AT	Admission only	One assessment only	Not specified	3188/3358 (95)	422/3188 (13)	Not specified
Dulin et al., 2022	bCAM	Admission and Inpatient	On admission and every nursing shift	Not specified	17,769/93,388 (19)* <i>*early in staged implementation, 63% by end of process. Range 12% (labor & delivery)-98% (medical wards)</i>	2593/17,769 (15)	Not specified
Fuchs et al., 2020	DOSS; ICDSC	Inpatient	3 × /day	Not specified	10,261/10733 (96)	3285/ 10,261 (32)* <i>*may include patients admitted to ICU assessed using ICDSC</i>	0
Garcez et al., 2020	CAM	Admission only	One assessment only	Not specified	Only patients assessed using tool included	445/1554 (29)	0
Han et al., 2021	4AT	Inpatient (post-op)	One assessment only	Post-op (<24 h)	1082/1082 (100)	222/1082 (21)	0
LaHue et al., 2021	NuDESC	Inpatient	2 × /day	Not specified	Not specified	13.0% during first epoch post intervention; 12.0%, 11.7%, and 13.0% in subsequent epochs	Not specified
Lee et al., 2019	DOSS	Inpatient	2 × /day	<24 h of admission	14,527/23821 (61)	2582/12082 (21)	Not specified
Lee et al., 2022	DOSS	Inpatient	2 × /day	<24 h of admission	7927/12082 (66)	2008/7927 (25)	0

TABLE 2 (Continued)

Author, year	Tool name	Timepoint(s) of assessment(s) in patient journey	Frequency of assessment(s)	Time from admission to first assessment	Tool completion rate: No. of patients or assessments performed/no. reported as eligible for assessment (%)	Positive score rate: No. of patients or assessments with a positive score/ no. of patients or assessments (%)	Number (proportion) of patients classed as <i>unable to assess</i> (%)
Marquetand et al., 2021	DOSS; ePA-AC; ICDSC	Inpatient	3×/day	Not specified	Only patients assessed using tool included	1285/3076 (42)* *may include patients admitted to ICU assessed using ICDSC	Not specified
Matharu et al., 2022	4AT	Inpatient (post-op)	One assessment only	Post-op (<1 week)	114,886/124960 (92)	32,244/114,886 (28)	Not specified
Peralta-Cuervo et al., 2021	CAM	Admission only	One assessment only	Not specified	Only patients assessed using tool included	816/ 1599 (51)	Not specified
Reynish et al., 2017	CAM	Admission only	One assessment only	Not specified	10,014/ 12,673 (79)	765/ 10,014 (8)	0
Rohatgi et al., 2019	CAM	Inpatient	Every nursing shift	Not specified	99	5333/105455 (2)	Not specified
Schubert et al., 2018	DOSS	Inpatient	Every 24 h	Not specified	10,906/ 29,278 (37)* *29278 “eligible patients ≥ 18 years” included in the study but assessment only recommended for patients at “high risk” (≥65 years or < 65 years and delirious symptoms)	3069/ 10,906 (28)	Not specified
Spiller et al., 2022	DOSS	Inpatient	3x/day	Not specified	29,967/ 48,840 (61)	21,612/345,662 (6) of observations	Not specified
Tyas et al., 2021	4AT	Inpatient (post-op)	One assessment only	Post op (<72 h)	18,511/21274 (87) 20,601/22098 (93)	Not specified Not specified	Not specified
Wong et al., 2018	NuDESC	Inpatient	2×/day	Not specified	Only patients assessed using tool included	878/18223 (5) of observations	Not specified
Zipser et al., 2022	DOSS; ePA-AC; ICDSC	Inpatient	3×/day	Not specified	98	Surgical: 2093/17,158 (12) Medical: 2237/8214 (27)	Not specified

Abbreviations: 4AT, the 4 A's Test; (b)CAM, (brief) Confusion Assessment Method; DOSS, Delirium Observation Screening Scale; ePA-AC, Electronic Patient Assessment-Acute Care; ICDSC, Intensive Care Delirium Screening Checklist; NuDESC, Nursing Delirium Screening scale.

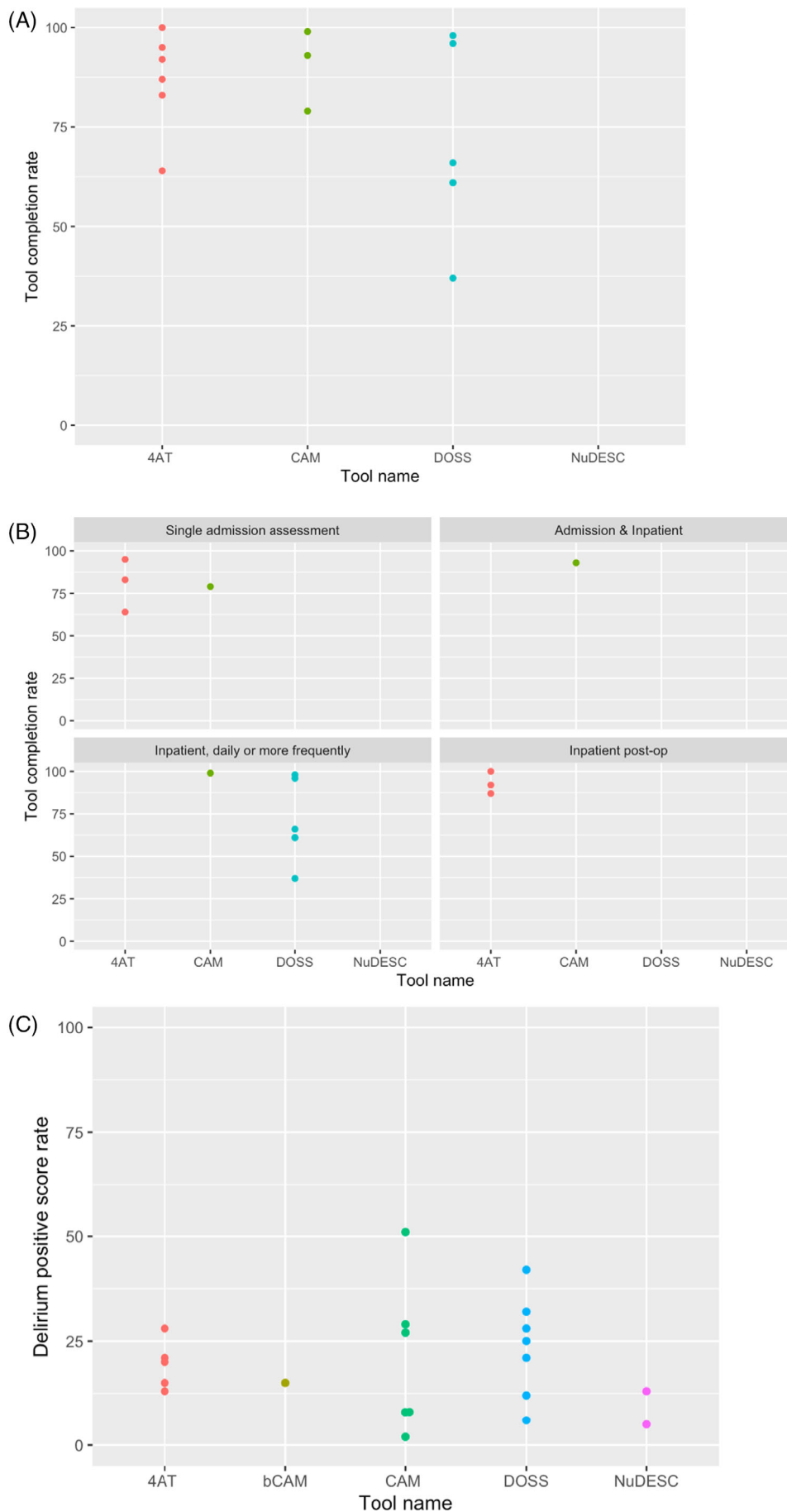


FIGURE 2 Tool completion rates and delirium detection rates in included studies. (A) Tool completion rate by tool name. (B) Tool completion rate by tool name and timepoint(s) of assessment. (C) Delirium positive score rate (proportion of patients or assessments with a positive score) by tool name. 4AT, the 4 'A's Test; (b) CAM, (brief) Confusion Assessment Method; DOSS, Delirium Observation Screening Scale; NuDESC, Nursing Delirium Screening Scale. NB studies using DOSS in addition to the ICDSC and/or ePA-AC reported as DOSS studies.^{32,35} Only one study reported a completion rate using the bCAM and this increased from 19% to 63% during a phased implementation process; this data is not presented in (A) or (B).¹⁸ Not all included studies reported a tool completion and/or delirium detection rate and hence do not appear in both figures.

by the end of implementation) to 100% (single orthopedic center, 4AT post-operatively) (Table 2 and Figure 2).^{17,39} Two of six studies reporting use of the DOSS had completion rates above 75%^{32,38} (overall range 37%³⁷–98%,³⁸ the latter when used in combination with the ICDSC and ePA-AC); all three using the CAM which included all patients^{12,30,31} (range 79%³⁰–99%¹²), and five of six using the 4AT^{21,39–42} (range 64%²²–100%³⁹). A 37% completion rate was calculated in one study using the DOSS, noting the study included all “eligible patients ≥18 years” but assessment was only recommended for patients deemed at high risk of delirium (≥65 years, or <65 years with delirious symptoms) as per hospital protocol.³⁷

Most studies did not report the number of patients without an assessment and only three provided information on the characteristics of patients with no assessment. Corradi et al. reported 15,094 of 88,206 patient encounters (17%) as *unable to assess* using the CAM; these patients had higher mortality rates than patients with a positive delirium score.¹⁸ Alhaidari et al. examined reasons for non-completion of the 4AT in 875 patients (11.2% of the study population), classifying these as follows: reduced patient alertness, communication barriers, pre-existing cognitive disorders, unstructured assessment completed, and prioritization of patient comfort and well-being.²¹ Anand et al. reported a similar mortality rate for 31% of patients without a 4AT completed as for patients with a 4AT score of 1–3 (positive for cognitive impairment but not delirium).²²

Delirium positive score rates

Rates varied by tool and timepoint of assessment.

The positive score rates on a single admission assessment were reported in studies using the CAM and the 4AT (Table 2). CAM-positive rates on admission ranged from 8% in unselected older emergency admissions (specialist nurse assessment)³⁰ to 51% in geriatrics ward (geriatrician assessment) admissions.²⁹ 4AT positive rates on admission were 15% (doctor assessment)²¹ and 20% (assessment by mixed disciplines).²² Di Bari et al. reported data on a single 4AT assessment completed by mixed staff disciplines in older Emergency Department (ED) patients who were discharged (68.5%) or admitted, reporting that overall 13% of patients had a positive score.⁴²

Two studies reported on admission assessment combined with inpatient repeated assessment. One study of the CAM in older geriatrics ward admissions (geriatrician assessment) found that 27% of patients had a positive score on admission and 20% on repeated inpatient assessment.³¹ Another study using the bCAM in adult

emergency admissions performed on admission and twice per day (nurse assessment) reported that 15% of assessments were positive.¹⁷

Eleven studies reported on repeated inpatient assessments with intended tool completion between 1 and 3 times per day (all nurse assessments). There were two studies using the CAM, both in unselected patients aged ≥18 years; overall the positive score rates were 8%¹⁸ and 2%¹² (below expected rates). Three studies using the DOSS in older inpatients reported positive score rates of 21%,³⁴ 25%,³⁵ and 6%,²⁰ with the latter below the expected rate. A study using the DOSS in patients aged ≥18 found a positive score rate of 28%.³⁷ Two studies used the DOSS combined with the ICDSC and the ePA-AC, and reported an overall positive score rate of 42% in patients aged ≥80 years,³⁶ and 12% in surgical patients and 27% in medical patients aged ≥18 years.³⁸ A study using either the DOSS (non-ICU) or ICDSC (ICU) in patients aged ≥65 reported an overall positive score rate of 32%.³² Two studies used the NuDESC. One reported a range of positive score rates in inpatients aged ≥50 across implementation phases, ranging from 5% to 13% post-implementation,³³ another in inpatients aged ≥50 years reported a positive score rate of 5%¹⁹; the rates of 5% are below expected rates.

Two studies reported postoperative positive score rates using the 4AT in older patients (assessment by mixed professionals) of 21%³⁹ and 28%.⁴⁰

Risk of bias

All but two studies had moderate–high risk of bias, mainly due to missing data and selection of the reported result (Supplementary Table 5). Very few studies reported on patients without a completed delirium assessment, making it difficult to determine how bias may affect results. However, given that patients without assessment may be *unable to assess*¹⁸ or have reduced alertness, communication barriers or pre-existing cognitive disorders,²¹ the observed biases may lead to delirium under-detection.

Gray literature

A gray literature search identified 4 eligible reports (Table 3). National hip fracture audit reports from England, Wales, and Northern Ireland (National Hip Fracture Database; NHFD), Scotland, and the Republic of Ireland reported using the 4AT,^{43–45} whilst a national report from Australia and New Zealand reported using the 4AT and CAM.⁴⁶ In the NHFD, over 80% of patients

TABLE 3 Published audits and reports of delirium assessment tool use at scale.

Author, year	Country	Report type	Patient population	Total no. of patients	Tool name	Timepoint of assessment(s) in patient journey	Frequency of assessment	Tool completion rate: No. of patients or assessments performed/no. reported as eligible for assessment (%)	Positive score rate: No. of patients or assessments with a positive score/ patients or assessments (%)
Royal College of Physicians, 2018	England, Wales & Northern Ireland	National audit (2017 data)	Admissions with acute hip fracture	65,958	4AT	Inpatient post-operative	One assessment only	>80%	25%
Public Health Scotland, 2022	Scotland	National audit	Admissions with acute hip fracture	7797	4AT	Admission only	Once on admission to ED, once on admission to ward within 24 hours	72.2%	Not specified
National Report, 2021	Republic of Ireland	National audit	Admissions with acute hip fracture	3806	4AT	Inpatient post-operative	Day 1, day 3, on discharge	41% assessed on day 1, 24% day 3, 35% on day of discharge	Not specified
ANZHR Annual Report, 2022	Australia & NZ	National audit	Admissions with acute hip fracture	15,331	Various including CAM, 4AT	Not specified	Not specified	Australia: 75% NZ: 65%	Australia: 39% NZ: 46%

Abbreviations: 4AT, the 4 A's Test; CAM, Confusion Assessment Method; ED, Emergency Department; NZ, New Zealand.

were assessed post-operatively using the 4AT; of these, 25% had a positive score.⁴³

DISCUSSION

Summary of main findings

This systematic review demonstrates that studies of delirium detection tools implemented in routine care show clinically important variability in the two key metrics of completion rates and delirium positive score rates. Tools were used at different stages of the patient journey, ranging from single ED or post-operative assessments to 1–3 times per day in inpatients. The CAM was successfully implemented in some contexts (e.g., as a single admission assessment in a specialist ward),^{28,31} but in some studies demonstrated lower than expected positive score rates, for example when used multiple times per day on every nursing shift throughout the inpatient stay.^{12,18} The 4AT was implemented as a single assessment in the ED or admission unit,^{21,22,42} or postoperatively^{39–41}; completion rates were high in all studies, except one substudy,²² and positive score rates were broadly aligned with expected delirium prevalence. The DOSS demonstrated low completion rates in studies when used alone,^{20,34,37} but higher overall completion rates when reported in combination with another tool (the ePA-AC or ICDSC)^{32,38}; positive score rates were broadly aligned with expected delirium prevalence, except in one model development study.²⁰ Only two studies used the NuDESC.^{19,33} Neither reported a completion rate; the positive score rate was lower than expected in both studies. Overall, these findings demonstrate that some tools perform poorly in certain routine care contexts, with low completion and/or much lower than expected positive score rates in the target clinical population.

The included studies provided limited information regarding factors that might affect the completion rate. One study reported an overall completion of 19% using the bCAM.¹⁷ This likely reflects the fact that implementation was staged across nursing units, but completion was measured across the whole medical center early in the implementation process; notably, completion was 63% by the end of the four-year process. Furthermore, bCAM assessment was performed in all hospitalized adults, with variation in completion rates from 12% in labor and delivery wards to 98% in medical wards by the end of the staged implementation. It may be that assessment only in higher risk groups, for example, older emergency admissions, may improve completion rates. Three studies reported high 4AT completion rates (92%, 93%, and 100%) in hip fracture patients postoperatively^{39–41}; all were in the UK, where delirium assessment is recommended in national guidelines and data

collected as part of national audits of care (although this information was not reported in the studies).⁴³ Rohatgi et al., reported a CAM completion rate of 99% in the context of a dedicated, highly-resourced delirium prevention initiative.¹² Qualitative studies have suggested that multiple factors influence completion rates, including time constraints, carer availability to provide informant history, tool simplicity, user-friendliness, and staff knowledge of delirium and of the specific tool.^{47–49} These factors could be targeted to improve completion rates.

Some of the variations in completion rates may be a function of the tool, as some allow patients to be classed “unable to assess” (e.g., the CAM and bCAM), whilst others do not (e.g., the 4AT). Inappropriate use of “unable to assess” may mask delirium and these patients often have adverse outcomes.¹⁸ Unfortunately, most studies did not report the proportion of patients recorded as “unable to assess” or did not provide information to determine how these patients were classified in the tool completion rate. Only three studies reported specifically on patients without a completed assessment.^{18,21,22} In a study using the 4AT, patients without an assessment had similar outcomes to patients with probable cognitive impairment,²² whilst in a study using the CAM, patients “unable to assess” had worse outcomes than patients with delirium.¹⁸ This concerning observation highlights the importance of elucidating barriers to delirium assessment.

Positive score rates varied widely. This variation likely reflects several factors. One is difference in true delirium occurrence rates in the populations and settings, for example, geriatrics ward versus ED attendances. Some studies reported on all inpatients aged ≥ 18 years,^{12,17–19,37,38} whereas one study only included the oldest old.³⁶ Some studies reported single assessments (prevalent delirium), and others repeated assessments during the inpatient stay (a combination of prevalent and incident delirium). Estimation of real-world sensitivity of the tools for delirium detection, reflecting concordance between positive score and true delirium occurrence rates, could only be done indirectly for all but one study. This study measured delirium prevalence with assessment by trained psychiatrists in a sample of 278 patients, reporting a rate of 17%, compared to a positive score rate of 2% using the CAM in routine practice.¹² For the other studies, estimates of expected rates in the clinical populations concerned can only be made by reference to relevant epidemiologic studies. A systematic review and meta-analysis of delirium occurrence in acute admissions to secondary care aged ≥ 18 years demonstrated a pooled delirium admission prevalence of 15% and cumulative incidence of 9%, with an overall occurrence rate of 23% across 33 studies.¹ In the present review, studies reported positive score rates as low as 8% on admission³⁰ and 2% on repeated inpatient testing¹² using the CAM in

acute medical and/or surgical populations. There was also variation in positive score rates between studies reporting the use of the same tool in comparable clinical populations. Evidence from national audit reports suggests that recommending a specified assessment tool may result in higher rates of delirium detection,^{46,50} although this requires dedicated resources. Provision of training and/or supervision may improve detection rates^{30,31}; for example, specific training in the use of the CAM is recommended for optimum performance.⁵¹

Strengths and limitations

Study strengths include systematic searching and analysis, including of gray literature, by two independent reviewers following a prospectively registered protocol. Studies were screened in all languages, with no restrictions on patient demographic variables such as age, ethnicity, and sex, or study location.

Some limitations should be acknowledged. An a priori decision was made to only include studies reporting on ≥ 1000 patients. This was to increase the likelihood that studies reflected large-scale clinical practice, but it is possible that high-quality smaller studies reporting on tool use in unselected patient populations in routine practice were omitted. We limited the review to published full-text studies and reports only. Although we performed an additional gray literature search to identify audit reports and relevant publications, tools reported in smaller forums may have been missed. For example, nearly one-third of UK units report using the “Single Question in Delirium” (SQiD), but no eligible peer-reviewed publications, abstracts or gray literature reports using the SQiD were identified.⁵² Studies reported completion rates at different stages of the patient journey, but it is possible that tool use was not fully reported in the studies. For example, some “admission only” tools may have been repeated later in the hospitalization as part of routine practice. We excluded studies in ICU settings. Delirium in the ICU is usually considered a different entity, with differences in population, tools, and expected delirium rates. This warrants a separate review. Most included studies were considered to have moderate–high RoB, highlighting the need for higher-quality implementation studies designed to assess tool performance.

Recommendations for future clinical practice

The findings of this systematic review have important implications for clinicians and policymakers seeking to

implement effective delirium assessment into practice. Tool completion rates were variable, and the findings relating to positive score rates demonstrate that certain tools substantially under-detect delirium when used routinely in some clinical contexts. When implementing a delirium assessment tool at scale, healthcare systems should plan to monitor tool completion and positive score rates, to iteratively investigate low completion or unexpectedly high or low positive score rates (choosing qualitative or quantitative methods appropriate to the context), and intervene to improve performance with repeated measurement and improvement. Implementing tools within EHRs could support routine use and measurement. Performance metrics should stimulate detailed evaluation of staff attitudes and barriers to delirium assessment through qualitative studies. Data-sharing and collaboration among healthcare providers and researchers could enable the creation of a comprehensive and up-to-date information repository on real-world assessment tool performance. This could be used to identify factors surrounding tool implementation in different contexts, to evaluate the impact of different strategies, and to inform future research and practice.

Recommendations for future research

Though the studies included in this review provide useful information, we recommend that future studies on delirium tool implementation report a wider set of variables, including a minimum dataset (Supplementary Table 6).

CONCLUSION

Delirium underdetection in routine care remains a major unmet clinical need. This review demonstrates that there is clinically important variability in tool completion and positive score rates relative to expected delirium rates, as well as in the proportion of patients unable to assess (where reported). Our findings are important for clinicians and policymakers seeking to improve delirium detection in acute hospital settings, and to inform future research. We encourage all healthcare systems to analyze and report their relevant data to share learning and work towards effective delirium detection at scale.

AUTHOR CONTRIBUTIONS

All authors were involved in conception and design of the study. RSP registered the protocol to PROSPERO. SDS and ZT provided guidance on systematic review methodology. RSP, CS, and AAng performed title/

abstract screening, full-text review, quality assessment, and risk of bias analysis. AMJM was available to resolve conflicts at title/abstract screening and full-text review. RSP and AMJM drafted the initial manuscript. All authors were involved in drafting the final manuscript. All authors had access to the data, provided feedback, and were involved in the finalization of the manuscript. All authors read and approved the final manuscript.

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CONFLICT OF INTEREST STATEMENT

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

Supplementary Table 1. PRISMA checklist.

Supplementary Table 2. Full search strategies for each database.

Supplementary Table 3. Gray literature: abstract-only and conference presentations.

Supplementary Table 4. List of excluded studies after full-text review.

Supplementary Table 5. Risk of bias in included studies.

Supplementary Table 6. Reporting standards for delirium assessment tool implementation studies.

Supplementary Figure 1. Timepoint(s) of delirium detection tool used in the patient journey.

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