Supplementary Material 1

Database Searched	Date Searched	PICO 1	PICO 2
MEDLINE (Ovid)	03/25/2021	560	890
Cochrane Central Register of Controlled Trials	03/25/2021	75	100
Embase (Ovid)	03/25/2021	834	1296
CINAHL (Ebsco)	03/25/2021	239	323
APA PsycINFO (Ovid)	03/25/2021	85	107
PubMed Central	03/25/2021	127	182
Web of Science	03/25/2021	259	361
Total		2160	3259
After deduplication: Yale De-duplicator		1464	2173
After deduplication: Covidence De-duplicator		1456	2166

Working Search Methods

The review team collaborated with a research librarian (A.L.B.) to develop and execute a comprehensive search of the literature. The search was created in partnership with several librarians and project team members from the larger GEAR 2.0 effort to conduct several scoping reviews on various topics related to dementia care in the field of emergency medicine. This search combined controlled vocabulary and title/abstract terms related to the accuracy and feasibility of dementia screening tools in the emergency department. The search was adapted from a GEAR 2.0 baseline search to fit the needs of the specific project question and translated for the following databases: MEDLINE (Ovid), Cochrane Central Register of Controlled Trials (CENTRAL), Embase (Ovid), CINAHL (Ebsco), APA PsycINFO (Ovid), PubMed Central, and Web of Science (Clarivate). All searches were performed on March 25, 2021. An exclusion filter from McGill University was used to focus on adult patient populations. No other publication type, language, or date filters were applied. Results were downloaded to a citation management software (EndNote) and underwent automated deduplication using a system at the Cushing/Whitney Medical Library at Yale University. Unique records were uploaded to a screening platform (Covidence) for independent review by several project team members using predetermined inclusion and exclusion criteria.

Used age filter from: https://libraryguides.mcgill.ca/knowledgesyntheses/search-tools

Ovid MEDLINE(R) ALL <1946 to March 24, 2021>

1 exp Emergency Medical Services/ 146955

2 Emergency Medicine/ 13903

3 (emergicenter* or triage* or unscheduled-acute-care).ti,ab. 20523

4 ((ED or EMS or ER) adj1 (care* or visit* or stay* or admit* or admission* or evaluation* or assess*)).ti,ab. 12310

5 (trauma adj1 (care* or support* or center* or centre* or department* or unit* or room* or ward* or service*)).ti,ab. 23093

6 ((Emergency or emergencies) adj2 (admit* or admission* or care* or treatment* or service* or dispatch* or department* or unit* or ward* or room* or center* or centre* or system* or personnel* or physician* or provider* or doctor* or nurs* or patient*)).ti,ab. 169983 7 or/1-6 273992

8 exp Dementia/ 172364

9 (dementia* or amentia* or demention* or CADASIL or Alzheimer* or Creutzfeldt-Jakob or Huntington* or Lewy-Bod*).ti,ab. 248697

10 Cognitive Dysfunction/ 20913

11 Cognition Disorders/ 64841

12 ((cognit* or neurocognit* or frontotemporal) adj2 (disorder* or defect* or deficit* or decline* or deteriorat* or disabilit* or dysfunction* or disfunction* or impaired or impairment* or interference*)).ti,ab. 138648

13 or/8-12 389398

14 exp "Diagnostic Techniques and Procedures"/ 7441551

15 di.fs. 2635879

16 exp "Mental Status and Dementia Tests"/ 8157

17 exp Geriatric Assessment/ 28853

18 exp Neuropsychological Tests/ 181403

19 ((mental* or cognitive* or cognition* or orientation* or agitation* or memory* or concentration* or dementia* or mini-cog* or mini-mental* or neurocognit*) adj3 (assess* or test* or eval* or screen* or guestion* or exam* or scale* or calculator*)).ti,ab. 190997

20 Montreal cognitive assessment.ti,ab. 3256

21 MOCA.ti,ab. 2881

22 Mini-Mental Status Examination.ti,ab. 1037

23 MMSE.ti,ab. 11543

24 Saint Louis University Mental Status.ti,ab. 37

25 SLUMS.ti,ab. 1701

26 "AD8".ti,ab. 228

27 Quick Dementia Rating System.ti,ab. 7

28 ODRS.ti.ab. 7

29 or/14-28 8937325

30 7 and 13 and 29 1041

31 [Accuracy Outcome Concept–For PICO 1] 0

32 exp "Sensitivity and Specificity"/ 601726

33 exp "Models, Theoretical"/ 1818464

34 exp "Reproducibility of Results"/ 414121

35 (accurac* or accurate* or reproducib* or specificit* or sensitivity* or likelihood* or like-lihood* or statistic* or analysis* or analyses* or analyze* or mathematic* or calculation* or ratio* or probabilit* or estimat* or false-positive* or false-negative* or truepositive* or true-negative* or concept* or theoretical*).ti,ab. 9626627 36 or/32-35 10701541

37 [Pragmatic Concept-For PICO 2] 0

38 (pragmati* or practical* or feasibilit* or usabilit* or acceptabilit*

or acceptance*).ti,ab. 598307

39 ease-of-use.ti,ab. 10062

40 (organization* or organisation* or administration* or method* or standard* or instrument* or tool*).ti,ab. 8787459

41 exp "Organization and Administration"/ 1530624

42 og.fs. 492823

43 exp Methods/ 692757

44 mt.fs. 3938387

45 st.fs. 742607

46 is.fs. 673805

47 (education* or training* or learn* or simulation*).ti,ab. 1587309

48 exp Education/ 828249

49 ed.fs. 285229

50 exp "Task Performance and Analysis"/ 36465

51 time/ or time factors/ 1216614

52 (time* or timing*).ti,ab. 3964012

53 Automation/ 18746

54 (automated* or automation*).ti,ab. 132404

55 or/38-54 15388882

56 [PICO 1: Combined & filtered] 0

57 30 and 36 579

58 (exp infant/ or exp child/ or adolescent/) not exp adult/ 1920361 59 57 not 58 560

60 [PICO 2: Combined & filtered] 0

61 30 and 55 936

62 (exp infant/ or exp child/ or adolescent/) not exp adult/ 1920361 63 61 not 62 890

Embase <1974 to 2021 March 24>

1 exp Emergency Health Service/ 108608

2 Emergency Medicine/ 42258

3 exp Emergency Ward/ 160076

4 exp emergency physician/ 13158

5 exp Emergency Nurse Practitioner/ 337

6 exp Emergency Nursing/ 6697

7 exp Emergency Patient/ 3803

8 (emergicenter* or Triage* or unscheduled-acute-care).ti,ab. 32081

9 ((ED or EMS or ER) adj1 (care* or visit* or stay* or admit* or admission* or evaluation* or assess*)).ti,ab. 25141

10 (trauma adj1 (care* or support* or center* or centre* or department* or unit* or room* or ward* or service*)).ti,ab. 29792

11 ((Emergency or emergencies) adj2 (admit* or admission* or care* or treatment* or service* or dispatch* or department* or unit* or ward* or room* or center* or centre* or system* or personnel* or physician* or provider* or doctor* or nurs* or patient*)).ti,ab. 258392

12 or/1-11 404652

13 exp Dementia/ 377840

14 (dementia* or amentia* or demention* or CADASIL or Alzheimer* or Creutzfeldt-Jakob or Huntington* or Lewy-Bod*).ti,ab. 347273

15 exp Cognitive Defect/ 516020

16 ((cognit* or neurocognit* or frontotemporal) adi2 (disorder* or defect* or deficit* or decline* or deteriorat* or disabilit* or dysfunction* or disfunction* or impaired or impairment* or interference*)).ti,ab. 213817

17 or/13-16 617546

18 clinical assessment/ 171905

19 exp dementia assessment/ 50912

20 di.fs. 3232060

21 exp geriatric assessment/ 17751

22 ((mental* or cognitive* or cognition* or orientation* or agitation* or memory* or concentration* or dementia* or mini-cog* or mini-mental* or neurocognit*) adj3 (assess* or test* or eval* or screen* or question* or exam* or scale* or calculator*)).ti,ab. 266911

23 montreal cognitive assessment/7784

24 Montreal cognitive assessment.ti,ab. 6548

25 MOCA.ti.ab. 7287

26 exp Mini Mental State Examination/ 42152

27 Mini-Mental Status Examination.ti,ab. 1807

28 MMSE.ti.ab. 24764

29 Saint Louis University Mental Status.ti,ab. 70

30 SLUMS.ti,ab. 1925

31 "AD8".ti,ab. 353

32 Quick Dementia Rating System.ti,ab. 7

33 QDRS.ti,ab. 10

34 or/18-33 3642188

35 [Accuracy Outcome Concept-For PICO 1] 0

36 exp diagnostic test accuracy study/ or exp diagnostic accuracy/ or exp accuracy/ 528272

37 exp "sensitivity and specificity"/ 388710

38 exp statistical analysis/ 2597960

39 exp conceptual framework/ 29130

40 (accurac* or accurate* or reproducib* or specificit* or sensitivity* or likelihood* or like-lihood* or statistic* or analysis* or analyses* or analyze* or mathematic* or calculation* or ratio* or probabilit* or estimat* or false-positive* or false-negative* or truepositive* or true-negative* or concept* or theoretical*).ti,ab. 12457155 41 or/36-40 13240261

42 [Pragmatic Concept–For PICO 2] 0

43 (pragmati* or practical* or feasibilit* or usabilit* or acceptabilit* or acceptance*).ti,ab. 767362

44 ease-of-use.ti,ab. 14290

45 (organization* or organisation* or administration* or method* or standard* or instrument* or tool*).ti,ab. 12490561

46 exp "organization and management"/ 2101767

47 (education* or training* or learn* or simulation*).ti,ab. 1980963

48 exp education/ 1482404 49 task performance/ 145790

50 exp time/ 635316 51 (time* or timing*).ti,ab. 5368545

52 automation/ or exp autoanalysis/ 66802 53 (automated* or automation*).ti,ab. 188393

54 or/43-53 17314801

55 [PICO 1 Combined & filtered] 0

56 12 and 17 and 34 and 41 834

57 56 not (exp juvenile/ not exp adult/) 815

58 [PICO 2 Combined & filtered] 0

59 12 and 17 and 34 and 54 1337

60 59 not (exp juvenile/ not exp adult/) 1296

APA PsycInfo <1806 to March Week 3 2021>

1 exp Emergency Medicine/ 357

2 exp Emergency Personnel/ 11799

3 (emergicenter* or Triage* or unscheduled-acute-care).ti.ab. 1651 4 ((ED or EMS or ER) adj1 (care* or visit* or stay* or admit* or admission* or evaluation* or assess*)).ti,ab. 1667

5 (trauma adj1 (care* or support* or center* or centre* or department* or unit* or room* or ward* or service*)).ti,ab. 1510

6 ((Emergency or emergencies) adj2 (admit* or admission* or care* or treatment* or service* or dispatch* or department* or unit* or ward* or room* or center* or centre* or system* or personnel* or physician* or provider* or doctor* or nurs* or patient*)).ti,ab. 17905

7 or/1-6 32107

8 exp Dementia/ 80210

9 (dementia* or amentia* or demention* or CADASIL or Alzheimer* or Creutzfeldt-Jakob or Huntington* or Lewy-Bod*).ti,ab. 106861

10 ((cognit* or neurocognit* or frontotemporal) adj2 (disorder* or defect* or deficit* or decline* or deteriorat* or disabilit* or dysfunction* or disfunction* or impaired or impairment* or interference*)).ti,ab. 80069

11 or/8-10 161116

12 exp Diagnostic Criteria/ 3425

13 exp Geriatric Assessment/ 1063

14 neuropsychological assessment/ 15227

15 ((mental* or cognitive* or cognition* or orientation* or agitation* or memory* or concentration* or dementia* or mini-cog* or mini-mental* or neurocognit*) adj3 (assess* or test* or eval* or screen* or question* or exam* or scale* or calculator*)).ti,ab. 126181

16 Montreal cognitive assessment.ti,ab. 1263

17 MOCA.ti,ab. 1034

- 18 mini mental state examination/777
- 19 Mini-Mental Status Examination.ti,ab. 536

20 MMSE.ti,ab. 6091

21 Saint Louis University Mental Status.ti,ab. 16

22 SLUMS.ti,ab. 472

23 "AD8".ti,ab. 44

24 or/12-23 141849

25 7 and 11 and 24 146

26 [Accuracy Outcome Concept-For PICO 1] 0

27 test sensitivity/ 308

28 models/ 70462

29 test validity/ or clinical validity/ 80831

30 (accurac* or accurate* or reproducib* or specificit* or sensitivity* or likelihood* or like-lihood* or statistic* or analysis* or analyses* or analyze* or mathematic* or calculation* or ratio* or probabilit* or estimat* or false-positive* or false-negative* or truepositive* or true-negative* or concept* or theoretical*).ti,ab. 2064410

31 or/27-30 2122572

32 [Pragmatic Concept-For PICO 2] 0

33 pragmatics/ 5210

1314.e41

34 (pragmati* or practical* or feasibilit* or usabilit* or acceptabilit* or acceptance*).ti,ab. 229448 35 ease-of-use.ti,ab. 2901 #6 36 test administration/ 3277 37 (organization* or organisation* or administration* or method* or standard* or instrument* or tool*).ti,ab. 1528305 38 exp Testing Methods/ 14401 39 exp training/ 78601 40 exp Time/ 20131 41 (time* or timing*).ti,ab. 737235 42 exp Automation/ 2457 #5 43 (automated* or automation*).ti,ab. 14229 44 or/33-43 2160738 45 [PICO 1: Combined only] 0 46 25 and 31 85 47 [PICO 2: Combined only] 0 48 25 and 44 107 #4 Web of Science PICO 1 # 13 259 #12 #3 Indexes=SCI-EXPANDED, CPCI-S, ESCI Timespan=All years # 12 264 #11 AND #10 AND #7 AND #4 Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years # 11 21.509.659 #2 TS=(accurac* or accurate* or reproducib* or specificit* or sensi-25.620 tivity* or likelihood* or like-lihood* or statistic* or analysis* or analyses* or analyze* or mathematic* or calculation* or ratio* or probabilit* or estimat* or false-positive* or false-negative* or true-positive* or true-negative* or concept* or theoretical*) #1 Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, 14.077 BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years # 10 324,054 #9 OR #8 Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years PICO 2 #9 # 13 35,680 361 TS=(Montreal cognitive assessment OR MOCA OR Mini-Mental #12 Status Examination OR MMSE OR Saint Louis University Mental Status OR SLUMS OR "AD8" OR Quick Dementia # 12 Rating System OR QDR) 375 Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years # 8 303.271 # 11 TS=((mental* or cognitive* or cognition* or orientation* or agitation* or memory* or concentration* or dementia* or mini-cog* or mini-mental* or neurocognit*) NEAR/3 (assess* or test* or eval* or screen* or question* or exam* or scale* or calculator*)) Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years #7 490,311 # 10 #6 OR #5 324,054

Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years 194,671 TS=((cognit* or neurocognit* or frontotemporal) NEAR/2 (disorder* or defect* or deficit* or decline* or deteriorat* or disabilit* or dysfunction* or disfunction* or impaired or impairment* or interference*)) Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years 381,017 TS=(dementia* or amentia* or demention* or CADASIL or Alzheimer* or Creutzfeldt-Jakob or Huntington* or Lewy-Bod*) Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years 206.166 #3 OR #2 OR #1 Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years 183,526 TS=((Emergency or emergencies) NEAR/2 (admit* or admission* or care* or treatment* or service* or dispatch* or department* or unit* or ward* or room* or center* or centre* or system* or personnel* or physician* or provider* or doctor* or nurs* or patient*)) Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years TS=(trauma NEAR/1 (care* or support* or center* or centre* or department* or unit* or room* or ward* or service*)) Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years TS=((ED or EMS or ER) NEAR/1 (care* or visit* or stay* or admit* or admission* or evaluation* or assess*)) Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years Web of Science Indexes=SCI-EXPANDED, CPCI-S, ESCI Timespan=All years #11 AND #10 AND #7 AND #4 Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years 24.214.287 TS=(pragmati* or practical* or feasibilit* or usabilit* or acceptabilit* or acceptance* or ease-of-use or organization* or organisation* or administration* or method* or standard* or instrument* or tool* or education* or training* or learn* or simulation* or time* or timing* or automated* or automation*) Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years

#9 OR #8

Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years

9 35,680

TS=(Montreal cognitive assessment OR MOCA OR Mini-Mental Status Examination OR MMSE OR Saint Louis University Mental Status OR SLUMS OR "AD8" OR Quick Dementia

Rating System OR ODR)

Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years

8 303,271

TS=((mental* or cognitive* or cognition* or orientation* or agitation* or memory* or concentration* or dementia* or mini-cog* or mini-mental* or neurocognit*) NEAR/3

(assess* or test* or eval* or screen* or question* or exam* or scale* or calculator*))

Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years

#7

490,311

#6 OR #5

Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years

#6

194,671

TS=((cognit* or neurocognit* or frontotemporal) NEAR/2 (disorder* or defect* or deficit* or decline* or deteriorat* or disabilit* or dysfunction* or disfunction* or impaired or impairment* or interference*))

Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years

5

381,017

TS=(dementia* or amentia* or demention* or CADASIL or Alzheimer* or Creutzfeldt-Jakob or

Huntington* or Lewy-Bod*)

Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years

#4

206.166

#3 OR #2 OR #1

Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years

3

J 183,526

TS=((Emergency or emergencies) NEAR/2 (admit* or admission* or care* or treatment* or service* or dispatch* or department* or unit* or ward* or room* or center* or centre* or system* or personnel* or physician* or provider* or doctor* or nurs* or patient*))

Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years

2

25,620

TS=(trauma NEAR/1 (care* or support* or center* or centre* or department* or unit* or room* or ward* or service*))

Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years

#1

14,077

TS=((ED or EMS or ER) NEAR/1 (care* or visit* or stay* or admit* or admission* or evaluation* or assess*))

Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, BKCI-S, BKCI-SSH, ESCI, CCR-EXPANDED, IC Timespan=All years

PubMed CENTRAL (PUuMed, Medline NOT Medline) PICO 1, 127 results

(("emergency care*"[Title/Abstract] OR "emergency treatment*"[-Title/Abstract] OR "emergency service*"[Title/Abstract] OR "emergency dispatch*"[Title/Abstract] OR "emergency department*"[Title/ Abstract] OR "emergency unit*"[Title/Abstract] OR "emergency ward*"[Title/Abstract] OR "emergency room*"[Title/Abstract] OR "emergency center*"[Title/Abstract] OR "emergency centre*"[Title/ Abstract] OR "emergency system*"[Title/Abstract] OR "emergencypersonnel"[Title/Abstract] OR "emergency physician*"[Title/Abstract] OR "emergency provider*"[Title/Abstract] OR "emergency doctor*"[-Title/Abstract] OR "emergency nurs*"[Title/Abstract] OR "emergency patient*"[Title/Abstract] OR "emergency admission*"[Title/Abstract] OR "emergency admit*"[Title/Abstract] OR "trauma care*"[Title/Abstract] OR "trauma treatment*"[Title/Abstract] OR "trauma service*"[Title/Abstract] OR "trauma dispatch*"[Title/Abstract] OR "trauma department*"[Title/Abstract] OR "trauma unit*"[Title/Abstract] OR "trauma ward*"[Title/Abstract] OR "trauma room*"[Title/ Abstract] OR "trauma center*"[Title/Abstract] OR "trauma centre*"[Title/Abstract] OR "trauma system*"[Title/Abstract] OR "trauma service*"[Title/Abstract] OR "trauma-personnel"[Title/Abstract] OR "trauma physician*"[Title/Abstract] OR "trauma provider*"[Title/Abstract] OR "trauma doctor*"[Title/Abstract] OR "trauma nurs*"[Title/ Abstract] OR "trauma patient*"[Title/Abstract] OR "emergicenter"[Title/Abstract] OR "unscheduled-acute-care"[Title/Abstract] OR "ED-care"[Title/Abstract] OR "ed visit*"[Title/Abstract] OR "ed stay*"[Title/Abstract] OR "ed admit*"[Title/Abstract] OR "ed admission*"[Title/Abstract] OR "ed evaluation*"[Title/Abstract] OR "ed assess*"[Title/Abstract] OR "ER-care"[Title/Abstract] OR "er visit*"[Title/Abstract] OR "er stay*"[Title/Abstract] OR "er admission*"[-"er evaluation*"[Title/Abstract] OR Title/Abstract] OR "er assess*"[Title/Abstract] OR "EMS-care"[Title/Abstract] OR "ems evaluation*"[Title/Abstract] OR "ems assess*"[Title/Abstract]) AND ("dementia*"[Title/Abstract] OR "amentia*"[Title/Abstract] OR "demention*"[Title/Abstract] OR "CADASIL"[Title/Abstract] OR "alzheimer*"[Title/Abstract] OR "Creutzfeldt-Jakob"[Title/Abstract] OR "huntington*"[Title/Abstract] OR "lewy bod*"[Title/Abstract] OR "cognitive disorder*"[Title/Abstract] OR "cognitive defect*"[Title/Abstract] OR "cognitive deficit*"[Title/Abstract] OR "cognitive decline*"[Title/Abstract] OR "cognitive deteriorat*"[Title/Abstract] OR "cognitive disabilit*"[Title/Abstract] OR "cognitive dysfunction*"[Title/ Abstract] OR "cognitive disfunction*"[Title/Abstract] OR "cognitiveimpaired"[Title/Abstract] OR "cognitive impairment*"[Title/Abstract] OR "cognitive interference*"[Title/Abstract] OR "neurocognitive disorder*"[Title/Abstract] OR "neurocognitive defect*"[Title/Abstract] OR "neurocognitive deficit*"[Title/Abstract] OR "neurocognitive decline*"[Title/Abstract] OR "neurocognitive deteriorat*"[Title/Abstract] OR "neurocognitive disabilit*"[Title/Abstract] OR "neurocognitive dysfunction*"[Title/Abstract] OR "neurocognitive impairment*"[Title/ Abstract] OR "frontotemporal disorder*"[Title/Abstract] OR "frontotemporal defect*"[Title/Abstract] OR "frontotemporal dysfunction*"[-Title/Abstract] OR "frontotemporal impairment*"[Title/Abstract] OR "impaired cognit*"[Title/Abstract] OR "impaired neurocogn*"[Title/ Abstract]) AND ((("mental*"[Title/Abstract] OR "cognitive*"[Title/Abstract] OR "cognition*"[Title/Abstract] OR "orientation*"[Title/Abstract] OR "agitation*"[Title/Abstract] OR "memory*"[Title/Abstract] OR "concentration*"[Title/Abstract] OR "dementia*"[Title/Abstract] OR "mini cog*"[Title/Abstract] OR "mini mental*"[Title/Abstract] OR "neurocognit*"[Title/Abstract]) AND ("assess*"[Title/Abstract] OR "test*"[Title/Abstract] OR "eval*"[Title/Abstract] OR "screen*"[Title/

Abstract] OR "question*"[Title/Abstract] OR "exam*"[Title/Abstract] OR "scale*"[Title/Abstract] OR "calculator*"[Title/Abstract])) OR ("montreal cognitive assessment"[Title/Abstract] OR "MOCA"[Title/ Abstract] OR "mini mental status examination"[Title/Abstract] OR "MMSE"[Title/Abstract] OR "saint louis university mental status"[-Title/Abstract] OR "SLUMS"[Title/Abstract] OR "AD8"[Title/Abstract] OR "quick dementia rating system"[Title/Abstract] OR "QDR"[Title/ Abstract])) AND ("accurac*"[Title/Abstract] OR "accurate*"[Title/Abstract] OR "reproducib*"[Title/Abstract] OR "specificit*"[Title/Abstract] OR "sensitivity*"[Title/Abstract] OR "likelihood*"[Title/ Abstract] OR "like lihood*"[Title/Abstract] OR "statistic*"[Title/Abstract] OR "analysis*"[Title/Abstract] OR "analyses*"[Title/Abstract] OR "analyze*"[Title/Abstract] OR "mathematic*"[Title/Abstract] OR "calculation*"[Title/Abstract] OR "ratio*"[Title/Abstract] OR "probabilit*"[Title/Abstract] OR "estimat*"[Title/Abstract] OR "false positive*"[Title/Abstract] OR "false negative*"[Title/Abstract] OR "true positive*"[Title/Abstract] OR "true negative*"[Title/Abstract] OR "concept*"[Title/Abstract] OR "theoretical*"[Title/Abstract])) NOT (("emergency care*"[Title/Abstract] OR "emergency treatment*"[Title/ Abstract] OR "emergency service*"[Title/Abstract] OR "emergency dispatch*"[Title/Abstract] OR "emergency department*"[Title/Abstract] OR "emergency unit*"[Title/Abstract] OR "emergency ward*"[Title/Abstract] OR "emergency room*"[Title/Abstract] OR "emergency center*"[Title/Abstract] OR "emergency centre*"[Title/ Abstract] OR "emergency system*"[Title/Abstract] OR "emergencypersonnel"[Title/Abstract] OR "emergency physician*"[Title/Abstract] OR "emergency provider*"[Title/Abstract] OR "emergency doctor*"[-Title/Abstract] OR "emergency nurs*"[Title/Abstract] OR "emergency patient*"[Title/Abstract] OR "emergency admission*"[Title/Abstract] OR "emergency admit*"[Title/Abstract] OR "trauma care*"[Title/Abstract] OR "trauma treatment*"[Title/Abstract] OR "trauma service*"[Title/Abstract] OR "trauma dispatch*"[Title/Abstract] OR "trauma department*"[Title/Abstract] OR "trauma unit*"[Title/Abstract] OR "trauma ward*"[Title/Abstract] OR "trauma room*"[Title/ Abstract] OR "trauma center*"[Title/Abstract] OR "trauma centre*"[Title/Abstract] OR "trauma system*"[Title/Abstract] OR "trauma service*"[Title/Abstract] OR "trauma-personnel"[Title/Abstract] OR "trauma physician*"[Title/Abstract] OR "trauma provider*"[Title/Abstract] OR "trauma doctor*"[Title/Abstract] OR "trauma nurs*"[Title/ Abstract] OR "trauma patient*"[Title/Abstract] OR "emergicenter"[Title/Abstract] OR "unscheduled-acute-care"[Title/Abstract] OR "ED-care" [Title/Abstract] OR "ed visit*" [Title/Abstract] OR "ed stay*"[Title/Abstract] OR "ed admit*"[Title/Abstract] OR "ed admission*"[Title/Abstract] OR "ed evaluation*"[Title/Abstract] OR "ed assess*"[Title/Abstract] OR "ER-care"[Title/Abstract] OR "er visit*"[Title/Abstract] OR "er stay*"[Title/Abstract] OR "er admission*"[-Title/Abstract] OR "er evaluation*"[Title/Abstract] "er OR assess*"[Title/Abstract] OR "EMS-care"[Title/Abstract] OR "ems evaluation*"[Title/Abstract] OR "ems assess*"[Title/Abstract]) AND ("dementia*"[Title/Abstract] OR "amentia*"[Title/Abstract] OR "demention*"[Title/Abstract] OR "CADASIL"[Title/Abstract] OR "alzheimer*"[Title/Abstract] OR "Creutzfeldt-Jakob"[Title/Abstract] OR "huntington*"[Title/Abstract] OR "lewy bod*"[Title/Abstract] OR "cognitive disorder*"[Title/Abstract] OR "cognitive defect*"[Title/Abstract] OR "cognitive deficit*"[Title/Abstract] OR "cognitive decline*"[Title/Abstract] OR "cognitive deteriorat*"[Title/Abstract] OR "cognitive disabilit*"[Title/Abstract] OR "cognitive dysfunction*"[Title/ Abstract] OR "cognitive disfunction*"[Title/Abstract] OR "cognitiveimpaired"[Title/Abstract] OR "cognitive impairment*"[Title/Abstract] OR "cognitive interference*"[Title/Abstract] OR "neurocognitive disorder*"[Title/Abstract] OR "neurocognitive defect*"[Title/Abstract] OR "neurocognitive deficit*"[Title/Abstract] OR "neurocognitive decline*"[Title/Abstract] OR "neurocognitive deteriorat*"[Title/Abstract] OR "neurocognitive disabilit*"[Title/Abstract] OR "neurocognitive dysfunction*"[Title/Abstract] OR "neurocognitive impairment*"[Title/

Abstract] OR "frontotemporal disorder*"[Title/Abstract] OR "frontotemporal defect*"[Title/Abstract] OR "frontotemporal dysfunction*"[-Title/Abstract] OR "frontotemporal impairment*"[Title/Abstract] OR "impaired cognit*"[Title/Abstract] OR "impaired neurocogn*"[Title/ Abstract]) AND ((("mental*"[Title/Abstract] OR "cognitive*"[Title/Abstract] OR "cognition*"[Title/Abstract] OR "orientation*"[Title/Abstract] OR "agitation*"[Title/Abstract] OR "memory*"[Title/Abstract] OR "concentration*"[Title/Abstract] OR "dementia*"[Title/Abstract] OR "mini cog*"[Title/Abstract] OR "mini mental*"[Title/Abstract] OR "neurocognit*"[Title/Abstract]) AND ("assess*"[Title/Abstract] OR "test*"[Title/Abstract] OR "eval*"[Title/Abstract] OR "screen*"[Title/ Abstract] OR "question*"[Title/Abstract] OR "exam*"[Title/Abstract] OR "scale*"[Title/Abstract] OR "calculator*"[Title/Abstract])) OR ("montreal cognitive assessment"[Title/Abstract] OR "MOCA"[Title/ Abstract] OR "mini mental status examination"[Title/Abstract] OR "MMSE"[Title/Abstract] OR "saint louis university mental status"[-Title/Abstract] OR "SLUMS"[Title/Abstract] OR "AD8"[Title/Abstract] OR "guick dementia rating system"[Title/Abstract] OR "ODR"[Title/ Abstract])) AND ("accurac*"[Title/Abstract] OR "accurate*"[Title/Abstract] OR "reproducib*"[Title/Abstract] OR "specificit*"[Title/Abstract] OR "sensitivity*"[Title/Abstract] OR "likelihood*"[Title/ Abstract] OR "like lihood*"[Title/Abstract] OR "statistic*"[Title/Abstract] OR "analysis*"[Title/Abstract] OR "analyses*"[Title/Abstract] OR "analyze*"[Title/Abstract] OR "mathematic*"[Title/Abstract] OR "calculation*"[Title/Abstract] OR "ratio*"[Title/Abstract] OR "probabilit*"[Title/Abstract] OR "estimat*"[Title/Abstract] OR "false positive*"[Title/Abstract] OR "false negative*"[Title/Abstract] OR "true positive*"[Title/Abstract] OR "true negative*"[Title/Abstract] OR "concept*"[Title/Abstract] OR "theoretical*"[Title/Abstract]) AND "medline"[Filter])

PubMed CENTRAL (PUuMed, Medline NOT Medline)

PICO 2, 182 results

((((emergency-care*[tiab] OR emergency-treatment*[tiab] OR emergency-service*[tiab] OR emergency-dispatch*[tiab] OR emergency-department*[tiab] OR emergency-unit*[tiab] OR emergencyward*[tiab] OR emergency-room*[tiab] OR emergency-center*[tiab] OR emergency-centre*[tiab] OR emergency-system*[tiab] OR emergency-personnel[tiab] OR emergency-physician*[tiab] OR emergency-provider*[tiab] OR emergency-doctor*[tiab] OR emergencynurs*[tiab] OR emergency-patient*[tiab] OR emergency-admission* [tiab] OR emergency-admit*[tiab] OR trauma-care*[tiab] OR traumatreatment*[tiab] OR trauma-service*[tiab] OR trauma-dispatch*[tiab] OR trauma-department*[tiab] OR trauma-unit*[tiab] OR traumaward*[tiab] OR trauma-room*[tiab] OR trauma-center*[tiab] OR trauma-centre*[tiab] OR trauma-system*[tiab] OR trauma-service* [tiab] OR trauma-personnel[tiab] OR trauma-physician*[tiab] OR trauma-provider*[tiab] OR trauma-doctor*[tiab] OR trauma-nurs* [tiab] OR trauma-patient*[tiab] OR emergicenter[tiab] OR unscheduled-acute-care[tiab] OR ED-care[tiab] OR ED-visit*[tiab] OR ED-stay*[tiab] OR ED-admit*[tiab] OR ED-admission*[tiab] OR EDevaluation*[tiab] OR ED-assess*[tiab] OR ER-care[tiab] OR ER-visit* [tiab] OR ER-stay*[tiab] OR ER-admission*[tiab] OR ER-evaluation* [tiab] OR ER-assess*[tiab] OR EMS-care[tiab] OR EMS-evaluation* [tiab] OR EMS-assess*[tiab]) AND (dementia*[tiab] OR amentia*[tiab] OR demention*[tiab] OR CADASIL[tiab] OR Alzheimer*[tiab] OR Creutzfeldt-Jakob[tiab] OR Huntington*[tiab] OR Lewy-Bod*[tiab] OR cognitive-disorder*[tiab] OR cognitive-defect*[tiab] OR cognitivedeficit*[tiab] OR cognitive-decline*[tiab] OR cognitive-deteriorat** [tiab] OR cognitive-disabilit*[tiab] OR cognitive-dysfunction*[tiab] OR cognitive-disfunction*[tiab] OR cognitive-impaired[tiab] OR cognitive-impairment*[tiab] OR cognitive-interference*[tiab] OR neurocognitive-disorder*[tiab] OR neurocognitive-defect*[tiab] OR neurocognitive-deficit*[tiab] OR neurocognitive-decline*[tiab] OR neurocognitive-deteriorat**[tiab] OR neurocognitive-disabilit*[tiab] OR neurocognitive-dysfunction*[tiab] OR neurocognitiveimpairment*[tiab] OR frontotemporal-disorder*[tiab] OR frontotemporal-defect*[tiab] OR frontotemporal-dysfunction*[tiab] OR frontotemporal-impairment*[tiab] OR impaired-cognit*[tiab] OR impaired-neurocogn*[tiab])) AND (((mental*[Title/Abstract])) OR cognitive*[Title/Abstract] OR cognition*[Title/Abstract] OR orientation*[Title/Abstract] OR agitation*[Title/Abstract] OR memory*[Title/ Abstract] OR concentration*[Title/Abstract] OR dementia*[Title/Abstract] OR mini-cog*[Title/Abstract] OR mini-mental*[Title/Abstract] OR neurocognit*[Title/Abstract]) AND (assess*[Title/Abstract] OR test* [Title/Abstract] OR eval*[Title/Abstract] OR screen*[Title/Abstract] OR guestion*[Title/Abstract] OR exam*[Title/Abstract] OR scale*[Title/ Abstract] OR calculator*[Title/Abstract])) OR (Montreal cognitive assessment[Title/Abstract] OR MOCA[Title/Abstract] OR Mini-Mental Status Examination[Title/Abstract] OR MMSE[Title/Abstract] OR Saint Louis University Mental Status[Title/Abstract] OR SLUMS[Title/ Abstract] OR "AD8" [Title/Abstract] OR Quick Dementia Rating System [Title/Abstract] OR ODR[Title/Abstract]))) AND (pragmati*[Title/Abstract] OR practical*[Title/Abstract] OR feasibilit*[Title/Abstract] OR usabilit*[Title/Abstract] OR acceptabilit*[Title/Abstract] OR acceptance*[Title/Abstract] OR ease-of-use[Title/Abstract] OR organization* [Title/Abstract] OR organisation*[Title/Abstract] OR administration* [Title/Abstract] OR method*[Title/Abstract] OR standard*[Title/Abstract] OR instrument* [Title/Abstract] OR tool*[Title/Abstract] OR education*[Title/Abstract] OR training*[Title/Abstract] OR learn*[Title/ Abstract] OR simulation*[Title/Abstract] OR time*[Title/Abstract] OR timing*[Title/Abstract] OR automated*[Title/Abstract] OR automation* [Title/Abstract])) NOT ((((emergency-care*[tiab] OR emergencytreatment*[tiab] OR emergency-service*[tiab] OR emergencydispatch*[tiab] OR emergency-department*[tiab] OR emergency-unit* [tiab] OR emergency-ward*[tiab] OR emergency-room*[tiab] OR emergency-center*[tiab] OR emergency-centre*[tiab] OR emergencysystem*[tiab] OR emergency-personnel[tiab] OR emergency-physician*[tiab] OR emergency-provider*[tiab] OR emergency-doctor*[tiab] OR emergency-nurs*[tiab] OR emergency-patient*[tiab] OR emergency-admission*[tiab] OR emergency-admit*[tiab] OR trauma-care* [tiab] OR trauma-treatment*[tiab] OR trauma-service*[tiab] OR trauma-dispatch*[tiab] OR trauma-department*[tiab] OR traumaunit*[tiab] OR trauma-ward*[tiab] OR trauma-room*[tiab] OR traumacenter*[tiab] OR trauma-centre*[tiab] OR trauma-system*[tiab] OR trauma-service*[tiab] OR trauma-personnel[tiab] OR trauma-physician*[tiab] OR trauma-provider*[tiab] OR trauma-doctor*[tiab] OR trauma-nurs*[tiab] OR trauma-patient*[tiab] OR emergicenter[tiab] OR unscheduled-acute-care[tiab] OR ED-care[tiab] OR ED-visit*[tiab] OR ED-stay*[tiab] OR ED-admit*[tiab] OR ED-admission*[tiab] OR EDevaluation*[tiab] OR ED-assess*[tiab] OR ER-care[tiab] OR ER-visit* [tiab] OR ER-stay*[tiab] OR ER-admission*[tiab] OR ER-evaluation* [tiab] OR ER-assess*[tiab] OR EMS-care[tiab] OR EMS-evaluation* [tiab] OR EMS-assess*[tiab]) AND (dementia*[tiab] OR amentia*[tiab] OR demention*[tiab] OR CADASIL[tiab] OR Alzheimer*[tiab] OR Creutzfeldt-Jakob[tiab] OR Huntington*[tiab] OR Lewy-Bod*[tiab] OR cognitive-disorder*[tiab] OR cognitive-defect*[tiab] OR cognitivedeficit*[tiab] OR cognitive-decline*[tiab] OR cognitive-deteriorat** [tiab] OR cognitive-disabilit*[tiab] OR cognitive-dysfunction*[tiab] OR cognitive-disfunction*[tiab] OR cognitive-impaired[tiab] OR cognitive-impairment*[tiab] OR cognitive-interference*[tiab] OR neurocognitive-disorder*[tiab] OR neurocognitive-defect*[tiab] OR neurocognitive-deficit*[tiab] OR neurocognitive-decline*[tiab] OR neurocognitive-deteriorat**[tiab] OR neurocognitive-disabilit*[tiab] OR neurocognitive-dysfunction*[tiab] OR neurocognitive-impairment*[tiab] OR frontotemporal-disorder*[tiab] OR frontotemporaldefect*[tiab] OR frontotemporal-dysfunction*[tiab] OR frontotemporal-impairment*[tiab] OR impaired-cognit*[tiab] OR impairedneurocogn*[tiab])) AND (((mental*[Title/Abstract] OR cognitive*[Title/ Abstract] OR cognition*[Title/Abstract] OR orientation*[Title/Abstract] OR agitation*[Title/Abstract] OR memory*[Title/Abstract] OR

concentration*[Title/Abstract] OR dementia*[Title/Abstract] OR minicog*[Title/Abstract] OR mini-mental*[Title/Abstract] OR neurocognit* [Title/Abstract]) AND (assess*[Title/Abstract] OR test*[Title/Abstract] OR eval*[Title/Abstract] OR screen*[Title/Abstract] OR guestion*[Title/ Abstract] OR exam*[Title/Abstract] OR scale*[Title/Abstract] OR calculator*[Title/Abstract])) OR (Montreal cognitive assessment[Title/ Abstract] OR MOCA[Title/Abstract] OR Mini-Mental Status Examination[Title/Abstract] OR MMSE[Title/Abstract] OR Saint Louis University Mental Status[Title/Abstract] OR SLUMS[Title/Abstract] OR "AD8"[Title/Abstract] OR Quick Dementia Rating System[Title/Abstract] OR QDR[Title/Abstract]))) AND (pragmati*[Title/Abstract] OR practical*[Title/Abstract] OR feasibilit*[Title/Abstract] OR usabilit*[Title/Abstract] OR acceptabilit*[Title/Abstract] OR acceptance*[Title/ Abstract] OR ease-of-use[Title/Abstract] OR organization*[Title/Abstract] OR organisation*[Title/Abstract] OR administration*[Title/Abstract] OR method*[Title/Abstract] OR standard*[Title/Abstract] OR instrument* [Title/Abstract] OR tool*[Title/Abstract] OR education* [Title/Abstract] OR training*[Title/Abstract] OR learn*[Title/Abstract] OR simulation*[Title/Abstract] OR time*[Title/Abstract] OR timing* [Title/Abstract] OR automated*[Title/Abstract] OR automation*[Title/ Abstract]) AND (medline[Filter]))

Cochrane CENTRAL (trials)

ID Search Hits

#1 (emergicenter* or Triage* or unscheduled-acute-care):ti,ab,kw 1755

#2 ((ED or EMS or ER) near/1 (care* OR visit* or stay* or admit* or admission* or evaluation* OR assess*)):ti,ab,kw 1914

#3 ((trauma) near/1 (care* or support* or center* or centre* or department* or unit* or room* or ward* or service*)):ti,ab,kw 1400

#4 ((Emergency or emergencies) near/2 (admit* or admission* or care* or treatment* or service* or dispatch* or department* or unit* or ward* or room* or center* or centre* or system* or personnel or physician* or provider* or doctor* or nurs* or patient*)):ti,ab,kw 19318 #5 #1 OR #2 OR #3 OR #4 21387

#6 (dementia* or amentia* or demention* or CADASIL or Alzheimer* or Creutzfeldt-Jakob or Huntington* or Lewy-Bod*):ti,ab,kw 20470

#7 ((cognit* or neurocognit* or frontotemporal) near/2 (disorder* or defect* or deficit* or decline* or deteriorat* or disabilit* or dysfunction* or disfunction* or impaired or impairment* or interference*)):ti.ab.kw 20493

#8 #6 OR #7 35214

#9 (mental* or cognitive* or cognition* or orientation* or agitation* or memory* or concentration* or dementia* or mini-cog* or mini-mental* or neurocognit*) NEAR/3

(assess* or test* or eval* or screen* or question* or exam* or scale* or calculator*):ti,ab,kw 35794

#10 (Montreal cognitive assessment OR MOCA OR Mini-Mental Status Examination OR MMSE OR Saint Louis University Mental Status OR SLUMS OR AD8 OR Quick Dementia Rating System OR QDR):ti,ab,kw 6749

#11 #9 OR #10 37101

#12 #5 AND #8 AND #11 106

#13 (accurac* or accurate* or reproducib* or specificit* or sensitivity* or likelihood* or like-lihood* or statistic* or analysis* or analyses* or analyze* or mathematic* or calculation* or ratio* or probabilit* or estimat* or false-positive* or false-negative* or truepositive* or true-negative* or concept* or theoretical*):ti,ab,kw 752666

#14 #12 AND #13 76

#15 (pragmati* or practical* or feasibilit* or usabilit* or acceptabilit* or acceptance* or ease-of-use or organization* or organisation* or administration* or method* or standard* or instrument* or tool* or education* or training* or learn* or simulation* or time* or timing* or automated* or automation*):ti,ab,kw 1215948

#16 #12 AND #15 101 #17 MeSH descriptor: [Infant] explode all trees 32413 #18 MeSH descriptor: [Child] explode all trees 56688 #19 MeSH descriptor: [Adolescent] explode all trees 104818 #20 #17 or #18 or #19 149861 #21 MeSH descriptor: [Adult] explode all trees 467867 #22 #20 NOT #21 58997 #23 #14 NOT #22 in Trials 75 #24 #16 NOT #22 in Trials 100 **CINAHL** Complete S1 MH "Emergency Medical Services+" Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 107,911 S2 MH "Emergency Medicine" Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 12,809 S3 MH "Physicians, Emergency" Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 4,445 S4 MH "Emergency Nurse Practitioners" Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 662 S5 MH "Emergency Nursing+" Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 15,518 S6 MH "Emergency Patients" Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 8,329 S7 TI (emergicenter* or Triage* or unscheduled-acute-care) OR AB (emergicenter* or Triage* or unscheduled-acute-care) Expanders -Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 11.300 S8 TI (("ED" or "EMS" or "ER") N1 (care* or visit* or stay* or admit* or admission* or evaluation* or assess*)) OR AB (("ED" or "EMS" or "ER") N1 (care* or visit* or stay* or admit* or admission* or evaluation* or assess*)) Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 9,224 S9 TI (trauma N1 (care* or support* or center* or centre* or department* or unit* or room* or ward* or service*)) OR AB (trauma N1 (care* or support* or center* or centre* or department* or unit* or room* or ward* or service*)) Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 14,368

S10 TI ((Emergency or emergencies) N2 (admit* or admission* or care* or treatment* or service* or dispatch* or department* or unit* or ward* or room* or center* or centre* or system* or personnel* or physician* or provider* or doctor* or nurs* or patient*)) OR AB ((Emergency or emergencies) N2 (admit* or admission* or care* or treatment* or service* or dispatch* or department* or unit* or ward* or room* or center* or centre* or system* or personnel* or physician* or provider* or dispatch* or department* or unit* or ward* or room* or center* or centre* or system* or personnel* or physician* or provider* or doctor* or nurs* or patient*)) Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 95,158

S11 S1 OR S2 OR S3 OR S4 OR S5 OR S6 OR S7 OR S8 OR S9 OR S10 Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 185,675

S12 (MH "Dementia+") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 76,007

S13 TI (dementia* or amentia* or demention* or CADASIL or Alzheimer* or Creutzfeldt-Jakob or Huntington* or Lewy-Bod*) OR AB (dementia* or amentia* or demention* or CADASIL or Alzheimer* or Creutzfeldt-Jakob or Huntington* or Lewy-Bod*) Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 81,565

S14 (MH "Cognition Disorders") OR (MH "Mild Cognitive Impairment") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 31,167

S15 TI ((cognit* or neurocognit* or frontotemporal) N2 (disorder* or defect* or deficit* or decline* or deteriorat* or disabilit* or dysfunction* or disfunction* or impaired or impairment* or interference*)) OR AB ((cognit* or neurocognit* or frontotemporal) N2 (disorder* or defect* or deficit* or decline* or deteriorat* or disabilit* or dysfunction* or disfunction* or impaired or impairment* or interference*)) Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 48,748

S16 S12 OR S13 OR S14 OR S15 Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 141,321

S17 (MH "Clinical Assessment Tools") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 169,223

S18 (MH "Mental Status/EV") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 615

S19 (MH "Diagnostic Tests, Routine") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 4,591

S20 (MH "Geriatric Assessment+") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 17,266

S21 (MH "Neuropsychological Tests+") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 36,416

S22 TI ((mental* or cognitive* or cognition* or orientation* or agitation* or memory* or concentration* or dementia* or mini-cog* or mini-mental* or neurocognit*) N3 (assess* or test* or eval* or screen* or question* or exam* or scale* or calculator*)) Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 9,268

S23 AB ((mental* or cognitive* or cognition* or orientation* or agitation* or memory* or concentration* or dementia* or mini-cog* or mini-mental* or neurocognit*) N3 (assess* or test* or eval* or screen* or question* or exam* or scale* or calculator*)) Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 61,663

S24 TI (Montreal cognitive assessment OR MOCA OR Mini-Mental Status Examination OR MMSE OR Saint Louis University Mental Status OR SLUMS OR "AD8" OR Quick Dementia Rating System OR QDR) Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 865

S25 AB (Montreal cognitive assessment OR MOCA OR Mini-Mental Status Examination OR MMSE OR Saint Louis University Mental Status OR SLUMS OR "AD8" OR Quick Dementia Rating System OR QDR) Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 6,756

S26 S17 OR S18 OR S19 OR S20 OR S22 OR S23 OR S24 OR S25 Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search.

Database - CINAHL Complete 247,110

S27 S11 AND S16 AND S26 Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 440

S28 (MH "Reliability and Validity+") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 258,799

S29 TI (accurac* or accurate* or reproducib* or specificit* or sensitivity* or likelihood* or like-lihood* or statistic* or analysis* or analyses* or analyze* or mathematic* or calculation* or ratio* or probabilit* or estimat* or false-positive* or false-negative* or true-positive* or true-negative* or concept* or theoretical*) Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 326,037

S30 AB (accurac* or accurate* or reproducib* or specificit* or sensitivity* or likelihood* or like-lihood* or statistic* or analysis* or analyses* or analyze* or mathematic* or calculation* or ratio* or probabilit* or estimat* or false-positive* or false-negative* or true-positive* or true-negative* or concept* or theoretical*) Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 1,596,933

S31 S29 OR S30 Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 1,735,631

S32 MH "Task Performance and Analysis+") OR (MH "Time Management") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 22,887

S33 (MH "Time+") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 197,095

S34 (MH "Education+") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 951,735

S35 TI ((pragmati* or practical* or feasibilit* or usabilit* or acceptabilit* or acceptance* or ease-of-use or organization* or organisation* or administration* or method* or standard* or instrument* or tool* or education* or training* or learn* or simulation* or time* or timing* or automated* or automation*) Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

Search Screen - Advanced Search

Database - CINAHL Complete 578,579

S36 AB ((pragmati* or practical* or feasibilit* or usabilit* or acceptabilit* or acceptance* or ease-of-use or organization* or organisation* or administration* or method* or standard* or instrument*

or tool* or education* or training* or learn* or simulation* or time* or timing* or automated* or automation*) Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 2,116,284 S37 S32 OR S33 OR S34 OR S35 OR S36 Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 2,991,525 S38 S27 AND S31 Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 244 S39 NOT ((MH "Child+") or (MH "Adolescence")) NOT (MH "Adult+") Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 623,765 S40 S38 NOT S39 Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 239 S41 S27 AND S37 Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 330 S42 NOT ((MH "Child+") or (MH "Adolescence")) NOT (MH "Adult+") Expanders - Apply equivalent subjects

Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases

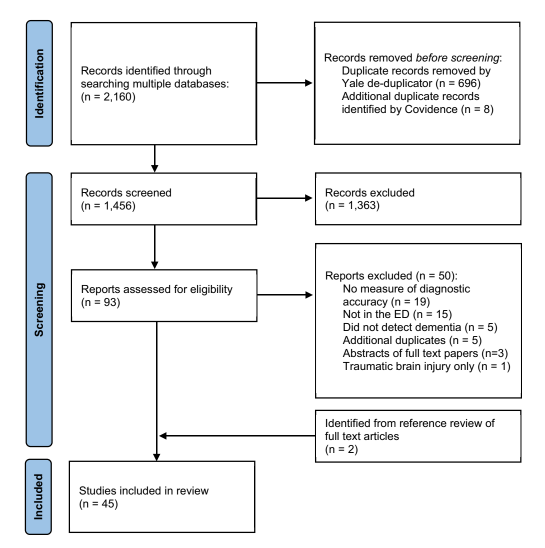
Search Screen - Advanced Search Database - CINAHL Complete 623,765 S43 S41 NOT S42 Expanders - Apply equivalent subjects Search modes - Boolean/Phrase Interface - EBSCOhost Research Databases Search Screen - Advanced Search Database - CINAHL Complete 323

DEDUPLICATION

PICO 1

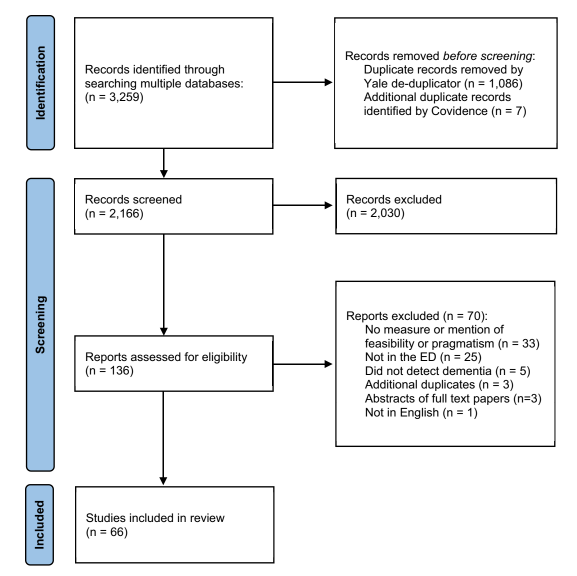
Summary: RIS data file: PICO 1_GEAR2.txt (2160 references) Total references considered: 2160 Sensitivity level: medium Total references removed: 696 Remaining references: 1464 Duration: 27 seconds (Uploading: 19 seconds|Processing: 8 seconds) Start time: March 25, 2021, 1651-0400 h Code version: 1e70a5 (January 25, 2021) PICO 2

Summary: RIS data file: PICO 2_GEAR2.txt (3259 references) Total references considered: 3259 Sensitivity level: medium Total references removed: 1086 Remaining references: 2173 Duration: 40 seconds (Uploading: 28 seconds|Processing: 12 seconds) Start time: March 25, 2021, 1658-0400 h Code version: 1e70a5 (January 25, 2021)



Supplementary Fig. 1. PRISMA-ScR flow diagram PICO 1.





Supplementary Fig. 2. PRISMA-ScR flow diagram PICO 2.

Supplementary Table 1 PICO 1 Abstraction Table

Study, Location, Time Frame (* in PICO 1 & PICO 2)	No. of Patients (median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design; Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Gold Standard for Dementia or Cognitive Impairment	Measures of Accuracy, Reliability, Sensitivity, Specificity, Negative LR, Positive LR, Correlation Coefficients, etc
Hospital of the University of Pennsylvania, PA, USA; ED; September 2007 and May 2008	N = 829; 65 y or older; mean age 75.7 ± 7.1 y (65- 105)	Spoke English, were 65 y or older, lived within a 30- mile radius of the ED in the state of Pennsylvania, and lived independently (ie, not in a nursing home)	End-stage disease with prognosis of 6 mo or less, cancer diagnosis with active treatment, known alcohol or drug abuse, history of neurologic disease (eg, cerebral vascular accident with residual effects, multiple sclerosis, etc), a previous medical history of dementia or delirium, or resided in a nursing home	to the ED of a large, urban, tertiary academic health center was conducted to identify rates of impairment among older adults; and identify relationships, if any, between ED environmental factors and presence of cognitive impairment	. ,	N/A	No measure of diagnostic accuracy but identified factors associated with positive cognitive screen tests: Patients were more likely to screen positive for cognitive impairment using the SIS if they were 85 y or older (RR 1.63, $P < .001$), Black (RR 1.85, $P < .001$), and male (RR 1.42, F < .001). Interestingly, only age was significantly associated with screening positive for cognitive impairment in the ED using the CLOX1 (75-84 y: RR 1.35, $P < .001$; \geq 85 y: RR 1.69, $P < .001$)
Carpenter*, 2008							Clinicians should select one population-appropriate primary screening tool and consider others for specific situations. For example, if one has very little time available, the Clock Drawing Test may be the most useful screening tool, whereas the Hopkins Verbal Learning Test may be superior in mildly impaired or highly educated patients. The MMSE has been evaluated most extensively, but current copyright restrictions limit its use, and diagnostic inaccuracy is a problem in relationship to educational levels. High-functioning, educated populations can be tested with instruments demonstrating less ceiling effect, but so far these tools are more time consuming.
Turner, 2012, Washington University in St Louis, St Louis, MO, USA; ED; time frame not specified	N = 170; mean age 74 y; 79% had cognitive impairment by MoCA	English-speaking community-dwelling patients aged ≥65 y		Randomized, single-center, cross sectional, consecutive sampling trial	Brief Alzheimer's Screen (BAS), Short Blessed Test (SBT), caregiver-AD8 (cAD8)	MoCA	BAS: sensitivity 61%, specificity 83%, LR+ 3.6, LR- 0.47, AUC 0.797 cAD8: sensitivity 54%, specificity 78%, LR+ 2.4, LR- 0.59, AUC 0.590 SBT: sensitivity 47%, specificity 89%, LR+ 4.1, LR- 0.60, AUC 0.746
Heidt, 2009, Washington University in St Louis, St Louis, MO, USA; ED; time frame not specified but was done in 5 mo	$\begin{split} N &= 251; \text{ mean age} \\ 76 \text{ y}; 53\% \text{ had} \\ \text{cognitive} \\ \text{impairment (MMSE} \\ \text{score} \leq 23) \end{split}$	English-speaking patients over age 65 y who had not received potentially sedating medications including anti-emetics, sedative-hypnotics, or narcotic-analgesia prior to criterion standard testing.		Prospective, cross-sectional convenience sampling	PMH, emergency physician note, nurse note, inpatient physician note documentation of cognitive impairment	MMSE	Did not document cognitive impairment in patients with abnormal MMSE: PMH: 86%, emergency physician: 72%, emergency nurse: 84%, inpatient physician: 60%

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Study, Location, Time Frame (* in PICO 1 & PICO 2)	No. of Patients (median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design; Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Gold Standard for Dementia or Cognitive Impairment	Measures of Accuracy, Reliability, Sensitivity, Specificity, Negative LR, Positive LR, Correlation Coefficients, etc
Eagles*, 2014, Civic Campus of the Ottawa Hospital; ED; June 17 —August 16, 2013	N = 260; age ≥ 75 y; mean age 83.7 (5.9) y; 38.4% had altered mental status	to the ED Monday to	 Patients who have been previously enrolled on a prior visit within 30 d; patients with known history of cognitive impairment or obviously altered or delirious; patients with communication barriers, including non-English or French speaking, auditory, verbal, or visual impairment severe enough to affect cognitive testing; patients who have a decreased level of consciousness such that they are not able to respond to verbal questioning; (4) patients triage and Acuity Scale level 1 or judged by their attending ED nurse or physician to be too critically ill; and (5) patients from long-term care/nursing homes and transfers from other hospitals. 	0800 and 1600 h	O3DY	Folstein Mini-Mental State Exam	O3DY by nurses had a sensitivity of 84.6% (95% CI 64.3-95.0) and specificity of 54.2% (95% CI 39.3- 68.3). O3DY by physicians had a sensitivity of 78.9% (95% CI 53.9- 93.0) and specificity of 39.4% (95% CI 23.4-57.8)
Carpenter, 2010, urban medical center; ED; 2 mo	N = 111; age > 65 y; mean age 77 y; 35% had cognitive impairment based on MMSE score	English-speaking patients aged ≥65 y who had not received potentially sedating medications		Cross-sectional convenience sampling	Caregiver- administered AD8, BAS, SBT, and the O3DYO3DY	MMSE score \leq 24	BAS: sensitivity 100%, specificity 53 and AUC 0.945 (95% CI 0.905-0.98% SBT: sensitivity 95%, specificity 68% and AUC 0.890 (95% CI 0.816-0.96 O3DY: sensitivity 95% and specificit 51% Caregiver AD8: sensitivity 87%, specificity 67%, and AUC 0.825 (95 CI 0.72% 0.012)
Rodriguez- Molinero, 2010, 4 tertiary university teaching hospitals; ED; July through November 2003	N = 101; undefined mean age or age criteria			Cross-sectional; older adult patients selected at random	(1) Physician recognition of cognitive impairment, (2) cognitive data shown in the patient's medical records	S-IQCODE (Short Form of the Informant Questionnaire on Cognitive Decline in the Elderly)	Cl 0.733-0.917) 1. Concordance between the physicians' impression on the presence of cognitive impairment and the S-IQCODE obtained from family member-carer was 0.26 (95 Cl 0.06-0.45). 2. Concordance between informatio on cognitive decline from medical records and the results of the S- IQCODE was 0.47 (95% Cl 0.05-0.8)

Huff*, 2001, University of Virginia; ED; 7 wk	N = 444; age > 55 y; no mean age reported; % cognitive impairment not reported; care partners not reported	Aged ≥55 y	Head trauma or multisystem trauma, inability to speak English, educational level of ≤7 y, acute medical illness, or contact or droplet isolation, patients that the research assistants felt might be harmed by mental distress or other discomfort by test administration	Prospective comparison of the QCS and the MMSE in a convenience sample; 16 h per day for a total of roughly 80 h per week for a 7-wk period	QCS	MMSE score	QCS scores were significantly correlated ($r = 0.783$) with MMSE scores
Gerson*, 1994, Community teaching hospital; ED; 3 mo (March 1, 1991–May 31, 1992)	$\begin{split} N &= 547; \mbox{ age } > 65 \mbox{ y;} \\ mean \mbox{ age } 76.7 \mbox{ y} \\ (\pm 7.7 \mbox{ SD}); \mbox{ 33.5\%} \\ had \mbox{ cognitive} \\ impairment \mbox{ based} \\ on \mbox{ OMCT} \end{split}$		Refusal to participate, physical condition prevented participation, known dementia, unable to communicate in English	ED social worker enrolled 7 AM to 3 PM Monday to Friday. Medical students enrolled in 3 different blocks: evenings 3 PM-7 PM, weekend days 7 AM -3 PM, nights 11 PM-7 AM. Five shifts per week were randomly selected in a 3:2:1 ratio to approximate patient flow and medical student availability.	Logistic regression model to predict cognitive impairment	6-item OMCT	Predictors of cognitive impairment were age >80 y (adjusted OR 3.68, 95% CI 2.21-6.14) and living in nursing home (adjusted OR 13.8, 95% CI 3.79-50.2)
Zaffarana, 2013, Florence, Italy; ED; January 1, 2010, to December 31, 2010	$N=169;~age>75~y;$ mean age 83 \pm 5.3 y; 18.9% had dementia		Subjects triaged as very low severity ("white" code) or with communication disorders	Retrospective analysis		Reported by a patient's kin or when specific indication and/or therapy were recorded in medical chart	•
Gagne, 2018, CHU de Quebec Hôpital de l'Enfant-Jésus (Quebec City), the CHU de Quebec CHUL (Quebec City), the Hôpital de Trois-Rivières), and the Centre Hospitalier de Lanaudière (Lanaudière); ED; 6-8 wk at each participating center (between February and May 2016)	mean age 76.8 (7.4) y	Patients aged ≥65 y who were independent or semiindependent (able to perform at least 5 activities of daily living), had an 8-h exposure to the ED environment from the time of registration (because of the high frequency of delirium with prolonged periods of stay in the ED), and were admitted (or waiting to be admitted) to a hospital ward	Lived in a nursing home or long-term care facility, had an unstable medical state that could lead to intensive care, could not communicate in French, or were unable to provide consent. Finally, patients with a history of a psychiatric disorder were also excluded.	Prospective comparison of 4AT- F and TICS-m	French version of the 4 A's Test (4AT-F)		Sensitivity 49% (34, 64). specificity 87% (82, 92), PPV 48% (33, 63), NPV 88% (74, 85), positive LR 3.77, negative LR 0.59
Marlow, 2010, Data from National Hospital Ambulatory Medical Care Survey (NHAMCS); ED; 2005-2006	None reported				Orientation to person, place, and time	Patient self-reported reason for visit	Sensitivity: 50.15% (SE = 4.27)
							(continued on next page)

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Supplementary Table 1 (continued)

Study, Location, Time Frame (* in PICO 1 & PICO 2)	No. of Patients (median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design; Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Gold Standard for Dementia or Cognitive Impairment	Measures of Accuracy, Reliability, Sensitivity, Specificity, Negative LR, Positive LR, Correlation Coefficients, etc
Carpenter, 2010, Washington University in St Louis; ED	N = 105; age > 65 y; mean age 77 y; 31% with cognitive impairment; N/A for care partners	0 - 0		Convenience sampling	MMSE administered at home after 3 wk follow-up	MMSE score <24 in ED	Cognitive dysfunction (MMSE < 24) was present in 31% in the ED, including 5% with delirium. At follow-up, 26% had cognitive dysfunction and none had delirium.
Dziedzic*, 1998, University of Virginia School of Medicine, Charlottesville, VA; ED; 2½ wk	N = 31; age ≥ 65 y; mean/median age	65 y or older, absence of recent head or multisystem trauma; able to speak English as a primary language; not be acutely experiencing alcohol or substance intoxication; score of 15 on the Glasgow Coma Scale; and educational level equivalent to 9 y or more		Patient sample was collected in a time period of 2½ wk, during shifts randomly distributed among 3 attending emergency medicine physicians. Shifts during daytime hours were maximized.	Physician perception of cognitive impairment	MMSE score	MMSE findings agreed with the treating physicians' assessments in 21 (67%) cases
Shah, 2009, Monroe County, NY; EMS and ED; June- December 2007	N = 187; age \geq 60 y; mean age 75.6 \pm 9.2 y; 8.6% had a medical history of dementia		Did not speak English or refused transport	Cross-sectional	EMS SIS	ED Mini-Cog, ED CLOX1, ED CLOX2	Compared to Mini-Cog: sensitivity 29% (20%-39%), specificity 96% (88%-99%) Compared to CLOX1: sensitivity 21% (13%-31%), specificity 93% (82%- 98%) Compared to CLOX2: sensitivity 23% (14%-35%), specificity 92% (83%- 97%)
Schofield, 2010, Glasgow, UK; accident and emergency (A&E); February- August 2007	$N = 601$; age ≥ 65 y; mean age 77 y; 37.6% with cognitive impairment (MMSE score ≤ 23); N/A for care partners		Verbal communication categorized as none, or sounds only according to the Glasgow Coma Scale, learning disability, severe hearing disability, unable to speak English and lack of interpreter	Convenience sampling, focusing on periods of high attendance by older patients	AMT10 (different cutoffs), AMT4 (different cutoffs), receiving nurse's judgment	MMSE score ≤24	Nurse's judgment: sensitivity 50.5% (44%-57%), specificity 98.6% (96%- 100%), PPV 97% (92%-99%), NPV 69% (65%-73%) AMT4 cutoff 3/4: sensitivity 80% (75- 85), specificity 88% (84-91), PPV 84% (78%-88%), NPV 85% (81%-89%) AMT10 cutoff 7/10: sensitivity 76% (69%-81%), specificity 93% (90%- 96%), PPV 90% (84%-93%), NPV 83% (79%-87%)
Carpenter*, 2011, Barnes Jewish Hospital, St Louis, MO; ED; July 1, 2008–April 20, 2009	mean age 76 y;	All ED patients aged ≥65 y	Patients who received medications that may have affected their mental status during the testing period (narcotics, benzodiazepines, antiemetics), were too critically ill to participate, as judged by the attending emergency physician, were unable to consent or cooperate with data acquisition, did not speak English, or refused to complete the questioning	Prospective, cross-sectional, convenience sampling. Enrollment occurred weekdays and weekends during equally distributed day, evening, and overnight shifts	SIS, AD8 (caregiver and patient), combined SIS and caregiver AD8 (cAD8; abnormal SIS or abnormal cAD8 result)	Mini-Mental State Examination score ≤24	 (15%-07%) SIS: sensitivity 74% (68%-80%), specificity 77% (74%-80%), positive LR 3.3 (2.5-4.1), negative LR 0.33 (0.25-0.44), AUC 0.83 (0.78-0.87) cAD8: sensitivity 63% (53%-72%), specificity 79% (73%-85%), positive LR 3.0 (1.9-4.6), negative LR 0.44 (0.31-0.62), AUC 0.74 (0.65-0.81) pAD8: sensitivity 37% (28%-46%), specificity 82% (77%-86%), positive LR 2.0 (1.1-3.3), negative LR 0.77 (0.63-0.93), AUC 0.67 (0.60-0.74) SIS or cAD8: sensitivity 89% (80%-95%), specificity 70% (63%-73%), positive LR 3.0 (2.3-3.6), negative LR 0.16 (0.07-0.30)

Carpenter*, 2011, Barnes Jewish Hospital, St Louis, MO; ED; June 10, 2009–March 9, 2010	mean age 78 \pm 8 y;		Patients receiving mental status—altering medications (antiemetics, benzodiazepines, or narcotics) prior to or during the testing period, emergency physician judgment of critical illness precluding informed consent or safe data collection, subject inability to consent or comply with data acquisition, non-English speaking, or refusal to complete the questioning		O3DY, BAS, SBT, and cAD8	MMSE score ≤23	SBT: sensitivity 95% (88%-98%), specificity 65% (61%-67%), positive LR 2.7 (2.2-3.0), negative LR 0.08 (0.03-0.2), AUC 0.930 (0.862-0.971) BAS: sensitivity 95% (88%-98%), specificity 52% (48%-54%), positive LR 2.0 (1.7-2.2), negative LR 0.10 (0.03-0.3), AUC 0.934 (0.867-0.974) O3DY: sensitivity 95% (85%-99%), specificity 51% (46%-53%), positive LR 2.0 (1.6-2.1), negative LR 0.10 (0.03-0.3) cAD8: sensitivity 83% (71%-91%), specificity 63% (55%-68%), positive LR 2.2 (1.6-2.8), negative LR 0.27 (0.1-0.5), AUC 0.816 (0.727-0.886) SBT or cAD8: sensitivity 91% (81%- 97%), specificity 27% (20%-30%), positive LR 1.2 (1.0-1.4), negative LR 0.32 (0.1-0.9) BAS or cAD8: sensitivity 97% (90%- 99%), specificity 11% (6%-12%), positive LR 1.1 (0.9-1.1), negative LR 0.27 (0.04-1.6) O3DY or cAD8: sensitivity 100%, specificity 0% (continued on next page)
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Study, Location, Time Frame (* in PICO 1 & PICO 2)	No. of Patients (median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design; Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Gold Standard for Dementia or Cognitive Impairment	Measures of Accuracy, Reliability, Sensitivity, Specificity, Negative LR, Positive LR, Correlation Coefficients, etc
Huang, 2020, Taipei Veterans General Hospital, Taiwan; ED; August 2018 to February 2019	N = 106; age \geq 75 y; mean age 87.3 \pm 5.2 y; 58.5%	Admission in the observation room of the ED, age of ≥75 y, and willingness to provide written informed consent	(1) Unstable clinical conditions, eg, using high-flow oxygen supplement, inotropic agents with pump, or emergent diseases, eg, acute myocardial infarction, cerebrovascular accident, surgical indication, sepsis; (2) diagnosed with malignant tumors within 3 y who were not in a stable disease state, including the need to receive tumor-related treatment or to receive unacceptable conditions for palliative care; (3) with autoimmune diseases who were not in a stable disease state requiring immunosuppressive agents to reach therapeutic targets; (4) unable to cooperate with blood evaluation or routine physiology test (eg, old stroke with bed- ridden status, aphasia, confusion, or unconsciousness, hemiplegia); (5) unwilling to participate in the trial; (6) unwilling to provide informed consent; (7) unable to cooperate with long-term follow-up assessment; and (8) subjects who had been enrolled in this study		Demographics, handgrip strength, and blood markers as predictors of cognitive impairment	Chinese MMSE score <23	The independent predictor of cognitive impairment was handgri strength (OR 0.86, 95% CI 0.80-0.94 <i>P</i> < .001) and age (OR 1.15, 95% CI 1.02-1.29; <i>P</i> < .05). TNF- <i>a</i> , IL-6, and visfantin were higher in the cognitive impairment group compared to controls, albumin wa lower. IL-6 was higher in the dementia group compared with those in the cognitive impairment — no dementia group.

median age visiting an E The target con cognitive im irrespective etiology. The was based o criteria (vers R, V) made b in geriatric c MMSE were substitute ge	teview when e following different environment than the ED or case- y ion consisted vith a mean or ≥65 y, pD. dition was mpairment of the e diagnosis on the DSM sion III, IV, IV- by a specialist care. CAM and accepted as a old standard heir wide use ractice. t was an to assess the ED. vided tat to	Systematic review and meta- analysis for diagnostic accuracy of tests detecting cognitive impairment (including delirium)	Ten different tests: 4AT, 6-CIT/SBT, AFT, AMT, AMT4, BAS, Mini-Cog, O3DY, SIS, cAD8	Eight studies used the MMSE as reference standard with cutoff values varying from 226 to 24 points, and 1 study used the DSM criteria for dementia.	 O3DY: no. of studies: 3, no. of patients: 518, pooled sensitivity 0.90 (95% CI 0.71-0.97), pooled specificity 0.61 (95% CI 0.47-0.73) 6-CIT/SBT: no. of studies: 3, no. of patients: 685, pooled sensitivity 0.89 (95% CI 0.78-0.95), pooled specificity 0.67 (95% CI 0.56-0.77) cAD8: no. of studies: 2, no. of patients: 482, pooled sensitivity 0.75 (95% CI 0.52-0.89), pooled specificity 0.71 (95% CI 0.52-0.85) SIS: no. of studies: 3, no. of patients: 746, pooled sensitivity 0.72 (95% CI 0.59-0.82), pooled specificity 0.79 (95% CI 0.75-0.83) (continued on next page)
					(comment on new page)

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Study, Location, Time Frame (* in PICO 1 & PICO 2)	No. of Patients (median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design; Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Gold Standard for Dementia or Cognitive Impairment	Measures of Accuracy, Reliability, Sensitivity, Specificity, Negative LR, Positive LR, Correlation Coefficients, etc
Barbic, 2018, Vancouver, Canada (not specifically mentioned); ED; June-November 2016	N = 117; age ≥ 75 y; median age 81.9 y (IQR 77-85); 12.0% (95% CI 6.1%- 17.9%); N/A for caregiver	Aged ≥75 y and presented to the ED	Patients triaged as Canadian Emergency Department Triage and Acuity Scale level 1 (resuscitation); if their condition was deemed too critical for evaluation; patients requiring emergent ED administration of medications that might negatively affect their neurologic and/or executive function (eg, opioids, benzodiazepines); patients with significant communication barriers affecting evaluation (eg, visual, verbal, or auditory impairments); patients with overt hallucinations, agitation, or confusion; patients who did not speak English; patients from nursing homes or long-term care facilities; patients with a previous diagnosis of cognitive impairment (eg, patients already enrolled in the study and patients unable to provide full, written, informed consent in English.		O3DY and SBT	MMSE score ≤24	O3DY: Sensitivity: 71.4% (95% CI 47.8 95.1), specificity: 56.3% (46.7%- 65.9%), AUC: 0.51 (95% CI 0.42- 0.61), positive LR: 1.63 (1.10-2.43) negative LR: 0.51 (0.22-1.18). The O3DY and MMSE scores agreed in 58.1% of cases. SBT: sensitivity: 85.7% (67.4%-99.9%), specificity: 58.3% (48.7%-67.8%), AUC: 0.52 (95% CI 0.43-0.61), positive LR: 0.25 (0.07-0.89). The SBT score agreed with the MMSE score in 61.5% of cases.
Roth, 2015, Pittsburgh, PA, USA; ED	N = 806; age 265 y; 58.2% with cognitive impairment	Aged ≥65 y and presented to the ED	Luguon.	Prospective observational study. Convenience sampling.	Multiple logistic regression model identifying factors predicting cognitive impairment.	SBT score \geq 4	A model of age >85 y (AOR 2.04, 95 CI 1.22-3.13), Black race (AOR 1.8, 95% CI 1.3-2.5), less than high school education (AOR 2.1, 95% CI 1.6-2.9), any fall in past year (AOR 1.8, 95% CI 1.2-2.4), any potentially inappropriate medication (AOR 1.4 95% CI 1.1-1.94) had moderate predictive accuracy for cognitive impairment (AUC = 0.66). A score of 2 would produce a sensitivity of 72.0%, specificity of 51.6%, positive LR of 1.49, negative LR of 0.54.

Eagles*, 2020, Ottawa, Canada; ED; June-August 2013	N = 260; age ≥75 y; mean age 83.7 y	Aged ≥75 y and presented to the ED	Patients who had a known history of cognitive impairment or were obviously cognitively impaired; were non English- or French- speaking patients; had auditory, verbal, or visual impairments severe enough to affect cognitive testing; were critically ill; resided in a long-term care home or were transferred from other hospitals.	Prospective cohort. Monday to Friday between 0800 and 1600 h	O3DY	MMSE score <25	When completed by nurses: (WORLD reversal) Agreement O3DY score and MMSE = 64.9% Sensitivity: 84.6% (95% CI 64.3-95.0) Specificity (95% CI): 54.2% (95% CI 39.3-68.4) When completed by nurses: (Serial 7s) Agreement O3DY score and MMSE = 67.7%. Sensitivity: 81.5% (95% CI 61.3-93.0). Specificity: 57.1% (95% CI 39.5-73.2). When completed by physicians: (WORLD reversal) Agreement O3DY score and MMSE = 53.8% Sensitivity: 78.9% (95% CI 53.9-93.0) Specificity: 39.4% (95% CI 23.4-57.8) When completed by physicians: (Serial 7s) Agreement O3DY score and MMSE = 51.2%. Sensitivity: 70.0% (95% CI 45.7-87.2). Specificity: 34.8% (95% CI 17.2-57.1).
Wilber, 2005, Akron, OH, USA; ED; fall of 2003	$N = 150$; age ≥ 65 y; mean age 75 (\pm 7) y; 23% with cognitive impairment	All patients aged ≥65 y who were able to communicate in English	Unable or unwilling to perform testing, those who were medically unstable, and those who received medications during the study that could affect their mental status	Prospective, randomized, cross- sectional study. Convenience sampling	SIS, Mini-Cog	MMSE score ≤23	SIS: sensitivity 94% (95% CI 73-100), specificity 86% (95% CI 74-94), PPV 68% (95% CI 46-85), NPV 98% (95% CI 89-100), AUC 0.96 (95% CI 0.92-1.0) Mini-Cog: sensitivity 75% (95% CI 48- 93), specificity 85% (95% CI 73-93), PPV 57% (95% CI 34-78), NPV 93% (95% CI 82-98)
Stair*, 2007, Boston, MA, USA; ED; June 2002 —October 2003	$\begin{split} N &= 684 (666 \\ completed \ both \\ MMSE \ and \ QCS); \\ age &> 18 \ y; \ mean \\ age \ 48 \ \pm \ 18 \ y \end{split}$	Age >18 y, ability to speak English or Spanish, and ability to answer questions		Prospective study	QCS	MMSE score \leq 23	Sensitivity 64%, specificity 85% For patients aged >55 y, the sensitivity was 64% and specificity 82%; for those with >8 y of education, the sensitivity was 59% and specificity 86%
Lague*, 2018, Quebec, Canada; ED; March-July 2015	N = 171; age ≥65 y; mean age 76.9 (8.3) y; 22% with cognitive impairment based on TICS-m \leq 27	 (1) Were aged ≥65 y; (2) were independent or semiindependent (can perform 5 of the 7 activities of daily living without any help); (3) spent ≥8 h in the ED; and (4) were admitted to any hospital ward. 	 Were living in a long- term care facility; (2) were unable to consent; were unable to communicate in French or English; (4) were experiencing an unstable medical condition leading to their admission to the intensive care unit (ICU); had a previous diagnosis of severe dementia or any other psychiatric condition; or had delirium during their 8-h ED stay. 	Prospective observational cohort	Bergman-Paris Question (BPQ)	TICS-m score ≤27	Sensitivity 86.5% (95% CI 71.2-95.5), specificity 27.8% (95% CI 20.4-36.3), PPV 25.0% (95% CI 17.8-33.4), NPV 88.1% (95% CI 74.4-96.0), AUC 0.57 (95% CI 0.50-0.64), adjusted AUC for age and sex 0.71 (95% CI 0.62-0.80)
							(continued on next page)

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1314.e58

Study, Location, Time Frame (* in PICO 1 & PICO 2)	No. of Patients (median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design; Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Gold Standard for Dementia or Cognitive Impairment	Measures of Accuracy, Reliability, Sensitivity, Specificity, Negative LR, Positive LR, Correlation Coefficients, etc
Bedard*, 2017, Quebec, Canada; ED; February- May 2016	N = 313; age ≥ 65 y; mean age 76.8 (7.5) y; 27.2% with cognitive impairment, TICS- m score ≤ 27	(1) Age ≥ 65 y; (2) an ED length of stay ≥8 h; (3) awaiting admission to a care unit; and (4) independent or semiindependent for activities of daily living	(1) Had an unstable medical condition that could lead to intensive care; (2) inability to communicate in French; (3) unable to consent; (4) history of a severe psychiatric condition (eg, schizophrenia, severe depression, or bipolar disorder); and (5) were living in a nursing home or another long-term care center		O3DY-F (French) score < 4	TICS-m score <27	Sensitivity 76.2% (66.7%-84.8%), specificity 67.6% (61.0%-73.6%), PPV 46.7% (38.1%-55.4%), NPV 88.4% (82.6%-92.8%), positive LR 2.4, negative LR 0.4
Rodriguez- Molinero, 2010, Madrid, Spain; ED; July- November 2003	$\begin{split} N &= 98; age \geq 65 \; y; \\ mean \; age \; 81.7 \pm \\ 7.3 \; y; \; 48\% \; positive \\ impairment \; with \\ Pfeiffer \; test; \; 66\% \\ were \; women \; and \\ mean \; age \; was \\ 56.1 \; y \; (\pm 12.6 \; y); \\ 64.4\% \; were \; children \\ of \; patients, \; 18.8\% \\ were \; spouses, \; and \\ 15.8\% \; other \; family \\ members; \; 1\% \; of \\ informants \; had \; no \\ family \; relationship \\ with \; the \; patients. \end{split}$	Patients older than 80 y, and patients between 65 and 79 y, provided that the latter had at least 2 comorbid chronic conditions	Those who had no available informant, who failed to sign the informed consent, who had no clinical history of emergencies, or whose physician failed to meet the criteria outlined below: Once a patient had been selected, one of the physicians declaring themselves responsible for the patient was required to participate. The highest ranking physician was selected, with those who had less than 1 year of experience or had already participated in the study in connection with another patient being excluded.	Cross-sectional; weekdays and weekends based on researcher availability	Physician perception of cognitive impairment	IQCODE	Concordance (κ) between IQCODE obtained from the relatives and physicians' perceptions of cognitive impairment was 0.26 (95% CI 0.06- 0.45; power of the comparison, 95%)

Schnitker*, 2015, Australia; ED; May 2012- February 2013	$N=580;$ age ≥70 y; mean age 80.3 \pm 6.7 y	All ED patients aged ≥70 y	Patients who (1) stayed >2 h in ED before the research nurse was available to approach them; (2) were severely ill, which prevented consent; (3) had consented for the study during a previous ED visit; (4) required an interpreter and where no suitable interpreter could be found in a timely manner (2 h); or (5) who were not able to participate in the planned phone follow-up (7 and 28 d post ED visit).	Prospective (?). Weekdays from 8 AM to 5 PM	Physician perception of cognitive impairment	OMCT score \geq 9	Sensitivity 24% (95% CI 17-31; PPV of 88%) and specificity 96% (95% CI 92- 99; NPV of 54%)	1314.e60
Ouellet*, 2016, Quebec, Canada; ED; May 2009 –March 2011	$N=306;$ age ${\geq}65$ y; mean age 77.0 \pm 7.2 y; 62.4% with cognitive impairment based on MoCA ${<}$ 26, 22.9% for MoCA ${<}$ 21	(1) Be 65 y or older, (2) be presenting to the ED specifically for a minor traumatic injury (ie, soft tissue/osseous lesions such as lacerations, contusions, sprains, simple extremity fractures, minor thoracic injuries, or minor head injury), (3) be discharged home within 48 h of the ED visit, and (4) be independent in basic activities of daily living in the month prior to the ED visit	 Injuries leading to admission to any ward, living in long-term care, (3) diagnosis of dementia, (4) delirium or confusion at the ED visit, and (5) inability to give a verbal consent, to communicate in French or English, or to attend follow-up assessments 	Prospective cohort. Any day or time; 24/7 recruitment schedule	Model predicting cognitive impairment	MoCA score < 26	Male sex, age 85 y, higher depression scores, slower walking speed, and self-reported memory problems were predictive of cognitive impairment (continued on next page)	A. Nowroozpoor et al. / JAMDA 23 (2022) 1314.e.

Supplementary Table 1 (continued)

Study, Location, Time Frame (* in PICO 1 & PICO 2)	No. of Patients (median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design; Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Gold Standard for Dementia or Cognitive Impairment	Measures of Accuracy, Reliability, Sensitivity, Specificity, Negative LR, Positive LR, Correlation Coefficients, etc
Carpenter, 2019, 4 studies occurred in the United States, 2 in Canada, 2 in Ireland, and 1 in Scotland; ED; Studies were conducted between 2003 and 2016	N = 2423 patients, N = 9 studies; age ≥ 65 y; a weighted average for dementia prevalence of 31% (range, 12%-43%)	Studies that described adults aged ≥65 y, evaluated in the ED setting with an index test for dementia and compared with an acceptable reference standard for dementia. A priori determinants of acceptable reference standards included the MMSE or more formal neuropsychological evaluation by qualified individuals (psychiatrist, neurologist, geriatrician) using <i>DSM-5</i> criteria. Studies had to provide sufficient detail on the dementia screening test and reference standard to construct 2-by-2 tables.		Systematic review and meta- analysis	AMT-4, cAD8, O3DY, SBT, and the SIS	MMSE, formal neuropsychologic evaluation by qualified individuals (psychiatrist, neurologist, geriatrician) using DSM-V criteria	AMT4: pooled sensitivity 0.74 (0.69- 0.79), pooled specificity 0.88 (0.85 0.91), pooled negative LR 7.69 (3.46 17.10), pooled negative LR 0.31 (0.10-0.90) cAD8: pooled sensitivity 0.72 (0.62- 0.81), pooled specificity 0.72 (0.64 0.79), pooled positive LR 2.53 (1.82 3.51), pooled negative LR 0.39 (0.26-0.59) O3DY: pooled sensitivity 0.92 (0.84- 0.96), pooled specificity 0.63 (0.58 0.68), pooled negative LR 0.17 (0.05-0.66) SBT: pooled sensitivity 0.87 (0.80- 0.92), pooled specificity 0.70 (0.66 0.74), pooled specificity 0.70 (0.66 0.74), pooled negative LR 2.18 (0.09-0.39) SIS: pooled sensitivity 0.69 (0.62- 0.74), pooled specificity 0.81 (0.77 0.84), pooled positive LR 3.53 (2.36 5.29), pooled negative LR 0.39 (0.31-0.50)
Bissig*, 2019, California, USA; ED; second half of 2016	mean age 68 \pm	Patients \geq 45 y old, who communicated in spoken English, and had been in the hospital for less than 24 h		Cross-sectional observational study	SIS	Previously documented cognitive impairment	Sensitivity 86%, specificity 77%

Wilding*, 2016, Ontario, Canada; ED; January –August 2010	N = 238; age ≥ 75 y; mean age 81.9 y; 13.4% with cognitive impairment based on MMSE score <25	Patients ≥75 y old	Patients who (1) were medically unstable (abnormal vital signs, required use of opioids, or those in obvious distress, as determined by initial ED staff or geriatric emergency management nurse assessment); (2) had a preexisting diagnosis of cognitive impairment or were obviously impaired (overtly confused, agitated, or hallucinating); (3) did not live in the city of Ottawa; (4) lived in long-term care; (5) had a primary language other than English or French; or (6) had hearing or visual impairment severe enough to effect cognitive testing	Prospective cohort; convenience sampling; 7 d per week from 8 ам to 4 рм	O3DY and AFT	MMSE score < 25	O3DY/MMSE: agreement 75.6% (95% CI 69.8%-80.7), sensitivity 93.8% (95% CI 77.8%-98.9%), specificity 72.8% (95% CI 66.1%-78.7%), positive LE 3.5, and negative LR 0.08. AFT, cutoff score < 15: AFT/MMSE: agreement 46.2% (95% CI 40.0%-52.6%), sensitivity 90.6% (95% CI 73.8%-97.5%), specificity 39.3% (95% CI 32.7%-46.4%), positive LR 1.5, and negative LR 0.24. AFT cutoff score < 10: AFT/MMSE: agreement 76.1% (95% CI 70.2%-81.0%), sensitivity 62.5% (95% CI 43.7%-78.3%), specificity 78.2% (95% CI 71.8%-83.5%), positive LR 2.9, and negative LR 0.48
Carpenter, 2011, USA; ED	N = 142; age ≥65 y; mean age 77 y; 34% with cognitive impairment based on MMSE score <24	Consenting English- speaking patients aged ≥65 y who had not received potentially sedating medications		Prospective, cross-sectional, convenience sampling	BAS, SBT, cAD8 stratified by education level	MMSE score <24	 In order of total, less than ninth-grade reading level, more than ninth-grade reading level, not graduating high school, and graduating high school BAS: sensitivity 90, 93, 75, 96, 77; specificity 43, 29, 48, 28, 57; positive LR 1.6, 1.3, 1.5, 1.3, 1.8; negative LR 0.24, 0.25, 0.52, 0.16, 0.41 SBT: sensitivity 90, 93, 67, 87, 71; specificity 47, 38, 53, 44, 50; positive LR 1.7, 1.5, 1.4, 1.6, 1.4; negative LR 0.22, 0.20, 0.62, 0.29, 0.59 cAD8: sensitivity 83, 70, 100, 75, 100; specificity 65, 46, 77, 64, 67; positive LR 2.4, 1.3, 4.4, 2.1, 3;
Boyd*, 2008, New Zealand; ED; December 2005 –March 2006	N = 139; age: ≥75 y (65 y for Maori and Pasifika elders); mean age 82.5 y; 35% with cognitive impairment (BRIGHT)	All those aged ≥75 y (65 y for Maori and Pasifika elders) who presented to the ED with a nonurgent complaint (triage level 3- 5)	Patients who were sleeping, undergoing medical procedures, or in distress. Cognitively impaired patients were only enrolled if their family was available to assist in completing the BRIGHT.	Cross-sectional convenience sampling; 4-h time blocks (Monday-Friday, 8 AM—8 PM) over a 12-wk period	BRIGHT	Cognitive performance scales (different cutoffs reported)	negative LR 0.26, 0.65, 0, 0.39, 0 BRIGHT cutoff ≥ 2 : sensitivity 0.81 (0.66, 0.91), specificity 0.34 (0.24, 0.46), positive LR 1.2, negative LR 0.6 BRIGHT cutoff ≥ 3 : sensitivity 0.78 (0.63, 0.89), specificity 0.54 (0.43, 0.66), positive LR 1.7, negative LR 0.4 BRIGHT cutoff ≥ 4 : sensitivity 0.70 (0.54, 0.83), specificity 0.74 (0.62, 0.83), positive LR 2.7, negative LR 0.4 (continued on next page)

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Study, Location, Time Frame (* in PICO 1 & PICO 2)	No. of Patients (median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design; Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Gold Standard for Dementia or Cognitive Impairment	Measures of Accuracy, Reliability, Sensitivity, Specificity, Negative LR, Positive LR, Correlation Coefficients, etc
Wilber*, 2008, Summa Health System's Akron City Hospital and Washington University, Barnes-Jewish Hospital and Cleveland clinic; ED; January 12, 2006–January 14, 2007	$N = 352$; age ≥ 65 y; mean age 77 (± 8) y; 32% with cognitive impairment based on MMSE	ED patients aged ≥65 y who were able to communicate in English	Patients who received medications that may have affected their mental status during the testing period (such as narcotics, antiemetics, or benzodiazepines), were critically ill, were unable to consent or cooperate with data acquisition, were previously enrolled, or refused to complete the questioning	Prospective, cross-sectional, convenience sampling	SIS	MMSE score ≤23	Sensitivity 63% (53, 72), specificity 81% (75, 85), PPV 60% (50, 69), NPV 83% (77, 87), positive LR 3.2 (2.4, 4.3), 0.5 (0.4, 0.6), AUC 0.77 (95% CI 0.72-0.83)
O'Sullivan, 2018, Cork, Ireland; ED; June-November 2015	N = 419; age ≥70 y; median age 77 y; 21.5% with dementia	All ED patients aged ${\geq}70$ y			4AT, 6-CIT (multiple cutoffs reported)	Standardized MMSE, IQCODE, <i>DSM-5</i> criteria	4AT (cutoff 0/1): sensitivity 0.84 (0.74-0.91), specificity 0.63 (0.57- 0.69), PPV 0.39 (0.32-0.46), NPV 0.94 (0.89-0.96), AUC 0.83 6-CIT (cutoff 9/10): sensitivity 0.81 (0.70-0.89), specificity 0.76 (0.71- 0.81), PPV 0.46 (0.37-0.55), NPV 0.94 (0.90-0.97)
Dyer, 2017, Dublin, Ireland; ED; June- August 2014	$\begin{split} N &= 196; \mbox{ age } \geq 70 \mbox{ y}; \\ mean \mbox{ age } 78.5 \ \pm \\ 5.9 \mbox{ y}; \mbox{ 50.1\% had} \\ cognitive \\ impairment \\ (delirium, MCI, or \\ dementia) \end{split}$	All ED patients aged \geq 70 y	Patients who were too unwell, unable to	 Prospective, cross-sectional, convenience sampling. 7 d per week both during and outside of working hours (outside of 0900 and 1700 h and on weekends) 	AMT4	CAM-ICU + AD8 + sMMSE (either positive)	Sensitivity 0.53 (0.42-0.63), specificity 0.96 (0.89-0.99), PPV 94.6% (84.9%-98.8%), NPV 73.33% (65.9%-79.9%), AUC 0.75 (0.68-0.82), positive LR 14.7 (4.7-45.4), negative LR 0.5 (0.4-0.6)
Bedard, 2019, Quebec, Canada; ED; February- May 2016	,	(1) age ≥ 65 y; (2) an ED length of stay ≥8 h; (3) awaiting admission to a care unit; and (4) independent or semiindependent for activities of daily living	(1) had an unstable medical condition that could lead to intensive care; (2) inability to communicate in French; (3) unable to consent; (4) history of a severe psychiatric condition (eg, schizophrenia, severe depression, or bipolar disorder); and (5) were living in a nursing home or another long-term care center	Prospective study	O3DY-F (French) < 4	TICS-m score <27	Sensitivity 76.2% (66.7%-84.8%), specificity 67.6% (61.0%-73.6%), PPV 46.7% (38.1%-55.4%), NPV 88.4% (82.6%-92.8%), positive LR 2.4, negative LR 0.4

Carpenter, 2011, USA; ED	N = 142; age ≥65 y; mean age 77 y; 34% with cognitive impairment based on MMSE score <24	Consenting English- speaking patients aged ≥65 y who had not received potentially sedating medications		Prospective, cross-sectional, convenience sampling	BAS, SBT, cAD8 stratified by whether MMSE was administered first or last	MMSE score <24	In order of total cohort, MMSE first, and MMSE last: BAS: sensitivity (%) 90, 91, 88; specificity (%) 43, 46, 41; positive LR 1.57, 1.67, 1.49; negative LR 0.24, 0.19, 0.30 SBT: sensitivity (%) 90, 86, 94; specificity (%) 47, 48, 48; positive LR 1.70, 1.65, 1.80; negative LR 0.22, 0.29, 0.12 cAD8: sensitivity (%) 83, 89, 78; specificity (%) 65, 68, 64; positive LR 2.37, 2.78, 2.18; negative LR 0.26, 0.16, 0.35
Carpenter, 2011, USA; ED; Han, 2018	N = 142; age ≥65; mean age 77; 34% with cognitive impairment based on MMSE score <24	Consenting English- speaking patients aged ≥65 y who had not received potentially sedating medications		Prospective, cross-sectional, convenience sampling	BAS, SBT, cAD8 in detecting MCI	Detection of MCI defined as normal MMSE score (≥24) but abnormal MoCA score (<26)	0.871) cAD8: sensitivity (%) 40 (34-41), specificity (%) 89 (60-98), positive LR 3.56 (0.84-20.59), negative LR 0.68 (0.60-1.11), AUC 0.506 (0.345- 0.666) SBT: sensitivity (%) 63 (59-65), specificity (%) 63 (59-65), positive LR 5.39 (1.84-19.51), negative LR 0.41 (0.36-0.61), AUC 0.799 (0.692- 0.906) Review of delirium and dementia, including the description of different tests used in detecting
Tong, 2016, Toronto, Ontario, Canada	N = 146; age ≥ 70 y; mean age 80.6 (6.0) y	Participants who were aged ≥70 y and who were present in the ED for a minimum of 4 h	Patients who were (1) critically ill (defined by the Canadian Triage Acuity Scale score of 1), (2) in acute pain (measured using the Numeric Rating Scale with a score ≥ 2 of 10), (3) receiving psychoactive medications, (4) judged to have a psychiatric primary presenting complaint, (5) previously enrolled, (6) blind, or (7) unable to speak English, follow commands, or communicate verbally	Feasibility study, prospective enrollment	Tablet-based serious game (whack-a- mole)	MMSE, MoCA	dementia in the ED Correlation of game response time (RT) and game accuracy: Game RT with MMSE score –0.558, with MoCA –0.339 Game accuracy with MMSE score –0.104 (nonsignificant), with MoCA –0.042 (nonsignificant)

4AT, 4 A's Test; AFT, Animal Fluency Test; AMT, Abbreviated Mental Test; AUC, area under the curve; BAS, Brief Alzheimer's Screen; BRIGHT, Brief Risk Identification for Geriatric Health Tool; cAD8, caregiver-completed Alzheimer's Disease-8; CAM, Confusion Assessment Method; CLOX1, clock-drawing task; *DSM, Diagnostic Statistical Manual of Mental Disorders*; EMS, Emergency Medical Services; LR, likelihood ratio; MCI, mild cognitive impairment; Mini-Cog, mini cognitive; MMSE, Mini-Mental State Examination; MoCA, Montreal Cognitive Assessment; N/A, not applicable; NPV, negative predictive value; O3DY, Ottawa 3DY; OMCT, Orientation, Memory, Concentration Test; PMH, past medical history; PPV, positive predictive value; QCS, Quick Confusion Scale; RR, risk ratio; SBT, Short Blessed Test; 6-CIT, Six-Item Cognitive Impairment Test; SIS, Six-Item Screener; sMMSE, standardized MMSE; TICS-m, Telephone Interview for Cognitive Status–modified.

Supplementary Table 2 PICO 2 Abstraction Table

Study, Location, Time Frame (* in PICO 1 and 2)	No. of Patients (Median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design- Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Measured for Feasibility, Pragmatic Nature, Timing, Efficiency, etc (Yes/No)	If Y: Measures Cited for Feasibility, Preferences, Duration of Instrument, Efficiency	If N: Points Made by Study About Ease of Use, Speed (Quick, Fast, etc), Setting, Integration Into Routine Care, etc
Adler 2019, University Hospital SUNY Downstate (urban tertiary hospital)	No data for ED	Birth year 1978 or earlier (40 y), use of upper extremities, lack of vision or hearing impairments, and English speaking	None	Pilot in ED: Patients in ED were triaged into "fast" and "slow" tracks depending on acuteness; patients in the slow track were approached	Computerized cognitive assessment tool used (the Cognigram) 4-item screen: How old	No		As per text description of study team experience: most adults declined participation, citing that they "had not been seen by a doctor" and "did not feel up to it." Another common reason for refusal was that they would soon be called and not have time to complete the Cognigram. Other observations in the ED included suspicion or nervousness about the Cognigram. Adults were not comfortable with the idea of cognitive testing, even when assured anonymity. They did not accept [the] purpose of evaluating Cognigram implementation, and mentioning that the Cognigram vas used to study dementia or ADRD did not help" Brevity of the test likely to be
					are you? What is your date of birth? What is this place? What year is it?			practical as per the report.
Bedard 2017*, between February and May 2016 in 4 hospitals across the province of Québec	76 y (SD 10.8)	Patients aged ≥65 y, with an 8-h ED stay, admitted on a care unit, independent or semiindependent in their ADL	Patient living in a long- term nursing facility, with an unstable medical condition, preexisting psychiatric condition or severe dementia, a delirium within the 8-h exposure to the ED	Comparison against reference tests	Administration of O3DY	No		None
Bissig 2019*, University of California–Davis Neurology consultation service	N = 100; age 68 y (SD 12); 5 with dementia, 1 MCI	Patient living in a long- term nursing facility, with an unstable medical condition, preexisting psychiatric condition or severe dementia, a delirium within the 8-h exposure to the ED		Integrating screener into ED neurology consultations; administered within 24 h of hospital arrival	SIS	No		None

Blomaard 2021, Leiden University Medical Center, tertiary hospital in the Netherlands; 4 December 2017 until 2 February 2018	N = 953; age 77 y (IQR 73-82)	All patients aged ≥70 y are eligible for screening after routine ED triage	Excluded patients who bypassed triage and patients who were triaged to the immediate urgency level	Before and after implementation of screening program: Implementation: recurring PDSA cycles for implementation, facilitation of program in electronic health record and standard operating procedures	APOP screener (which includes 3 questions on dementia and cognition) followed by interventions: screening older patients for risk of functional decline or mortality and signs of impaired cognition; second, targeted interventions for high-risk patients in the ED; and third, interventions for high-risk patients who are hospitalized or discharged home		Comparison of ED LOS before and after implementation of screener: ED LOS 202 min (IQR 133- 290 min) before vs 196 min (IQR 133- 265 min) after; $P =$.152; hospital admission rate 40% before and 39% after; P = .642	
Boucher 2019, Hôpital de l'Enfant-Jésus (CHU de Québec –Université Laval) between May and July 2018	N = 67; age 75.5 ± 8 y; mild dementia 7/67	Patients aged ≥65 y presenting to the ED of the Hôpital de l'Enfant-Jésus (CHU de Québec –Université Laval) for any medical reason; caregiver, relative, or close friend of a study participant who was present at the time of enrolment	Required resuscitation (CTAS 1); were unable to speak French; were unable to consent; had a physical condition preventing them from using the electronic tablet	comparing tablet	•	Yes	Patient-reported acceptability measure: TAP questionnaire; mean adjusted TAP scores showed no difference: 2.36 for standard RA assessments vs 2.20 for self-assessment using a tablet ($P = .08$); subgroup analysis with age > 85 y showed worse acceptability for tablet or self- assessment	Additional open-ended questions: that assess acceptability and preference of the 2 modes of assessments; comments include liked being able to concentrate and take their time answering the questions on the tablet; the main reason for refusal was fear or dislike of technology
Boyd 2008*, New Zealand ED in Auckland; a 12- wk period between December 2005 and March 2006	N = 139; age 82.5 (±5.4) y	Aged ≥75 y (65 y for Maori and Pasifika elders) who presented to the ED with a nonurgent complaint (triage level 3-5) during a convenience sample of 4-h time blocks (Monday–Friday, 8 AM−8 PM) during the study period; cognitively impaired patient only enrolled if family was available to complete assessment		Cross-sectional study	Comparison of 11-item BRIGHT case-finding tool administered in ED against comprehensive geriatric assessment within 10 d	No		75% of participants had assistance from a visitor or the RA to complete the BRIGHT assessment
Calf 2021*, Systematic review of cognitive screening instruments in ED	,	assessment		Systematic review of diagnostic accuracy of instruments		No		(continued on next page)

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Study, Location, Time Frame (* in PICO 1 and 2)	No. of Patients (Median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design- Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Measured for Feasibility, Pragmatic Nature, Timing, Efficiency, etc (Yes/No)	If Y: Measures Cited for Feasibility, Preferences, Duration of Instrument, Efficiency	If N: Points Made by Study About Ease of Use, Speed (Quick, Fast, etc), Setting, Integration Into Routine Care, etc
Carpenter 2011, level 1 trauma center ED; July 2008–February 2009	21 physicians and 34 nurses (response rate 42%)	Physicians and nurses at a level 1 trauma ED		Cross-sectional survey of staff	8-item survey on ED management of geriatric patients; for the previous 8 mo, older adults were screened by a geriatric technician for cognitive dysfunction (MMSE), falls and function (OARS scale)	Yes	8-item survey regarding geriatric technician role (acceptability and feasibility); 71% of physicians and 85% of nurses found geriatric technician screening as an overall benefit to older patients; 0% of physicians and 18% of nurses thought that geriatric technician screening prolonged the ED length of stay	
Carpenter 2008*, systematic review abstract concerning the practicality and accuracy of brief cognitive screening instruments in primary care		Studies enrolling subjects older than 60 y and which used an acceptable criterion standard to diagnose dementia	Non—English language articles, inpatient or nursing home isolated populations, memory disorder clinic populations without an adequately characterized outside control group, or populations with less than 6 y of median education			Yes	Time needed to administer; reproduction limited to copyright	
Carpenter 2011*, tertiary medical center ED; from July 1, 2008, to April 20, 2009	N = 371; mean age 76 y	e All ED patients aged ≥65 y			SIS, AD8, and MMSE	No		Using both instruments requires more time and training, with the additional need to find consenting caregivers to complete the AD8

questioning

Carpenter 2011*, urban academic university – affiliated medical center between June 2009 and March 2010	$N=$ 169; age = 78 \pm 8 y	All ED patients aged ≥65 y	Patients receiving mental status -altering medications (antiemetics, benzodiazepines, or narcotics) prior to or during the testing period, emergency physician judgment of critical illness precluding informed consent or safe data collection, subject inability to consent or comply with data acquisition, non -English speaking, or refusal to complete the questioning		O3DY, BAS, SBT, cAD8 compared against MMSE	No		
Clevenger 2012, systematic review or scopir review?	Includes 209 articles g pertaining to care for PWD in ED	2		Systematic review or scoping review??	Clinical care for PWD in ED (includes assessment)	ı No		
	os N = 2629; mean age 79 y (IQR 74-	All patients aged ≥70 y	Red triage category (highest acuity) according to the Manchester Triage System (MTS), an unstable medical condition, no permission of nurse or physician to approach the patient, a language barrier and impossibility to obtain informed consent	Multicenter cohort study	APOP screener (which includes 3 questions on dementia/ cognition)	Yes	Mean time to complete the screener was 93 s (SD 29); overall rating of clinical usability was positive, with a mean Likert score of 3.79 (out of 5; SD 0.63)	
Dyer 2017, Irish tertiary urban referral university teaching hospita June-August 201		Patients aged ≥70 y who presented to the ED		Convenience sample; cross- sectional	Informant history; cognitive screeners for delirium (CAM- ICU) and dementia (sMMSE and AD8)	Yes	The length of time to contact informants was $3.1 (\pm 5.8)$ min. In 9.1% (6/66), it took 10 min or longer to contact the informant; brief informant interviewing (mean duration, 6 min); rating of privacy (8.4 ± 1.6/10) and accessibility (8.5 ± 1.47/10)	
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Dziedzic 1998*, academic ED of a university-based hospital	N = 31	Age ≥65 y, absence of recent head or multisystem trauma; able to speak English as a primary language; not be acutely experiencing alcohol or substance intoxication; score of 15 on the GCS; and educational level equivalent to ≥9 y		Cross-sectional	MMSE compared against constructed interview for physician	No		
Eagles 2020*, academic tertiary care hospital ED between June and August 2013		. – .	A known history of cognitive impairment or were obviously cognitively impaired; were non-English or French speaking patients; had auditory, verbal, or visual impairments severe enough to affect cognitive testing; were critically ill; resided in a long-term care home or were transferred from other hospitals.	Prospective cohort	O3DY	Yes	Postimplementation survey of nurses and physicians: 98%, 95%, and 88% of physician respondents judged the O3DY tool to be easy to learn, to use, and to remember, respectively; 97% agreeing that the O3DY tool is easy to learn and use and 94% reporting that it is easy to remember (nurses)	
Eagles 2014, a tertiary-care ED	N = 198; mean age 84.2 y; 31% with evidence of impaired mental status	\geq 75 y of age		Prospective cohort	O3DY	No		Mentioned that it is a feasible tool for ED
Fox 2018, ED (not specified)	N = 785; 81.4 (SD 6.4) y; 9% with dementia diagnosis	Aged \geq 70 y		Prospective randomized double-blind diagnostic accuracy study	4AT	No		Rapid delirium assessment instrument, feasible in routine care
Gerson 1994*, midwestern community teaching hospital; March-May 1992	76.7 y (7.7 SD)	Age ≥65 y treated in the ED	Refused to participate, physical condition prevented participation, had known dementia, unable to communicate in English	Cross-sectional study	Six-item OMCT	Yes	Mean time of 1.9 min (+0.91 SD) was required to complete the test	
Graf 2010, Letter to the editor; commentary, and evidence synthesis			-	Evidence synthesis	QCS described for cognition assessment			QCS, which can be completed more quickly (~2 min) that the MMSE

Supplementary Table 2 (continued)

Graf, 2012	2						No	In ED, screening tools developed to detect these geriatric problems have to be quick, easy to use, and to present a high sensibility.
Groening, 2	2020						No	Though some emergency physician might consider "the old patient" as not exciting, there is a broad consensus that pragmatic geriatric screening tools are required. More practical tools will have to be developed in the future.
ED betwe	reen 15, 2012,	N = 117; mean age 76.4 (±8) y	Aged \geq 65 y		Convenience sample; cross- sectional	- Nurse-administered 6- CIT	No	Noted considerable variation in applicability and successful implementation of the screening instrument between nurses despite training
Han 2018						Table listing screening tests (p. 344 Table 5); AD8, BAS, Mini-Cog, O3DY, SIS, SBT		Mentioned time required for certain tests (Mini-Cog, 10 min; SBT, <5 min; O3DY, <2 min)
Hare 2008, hospital i Western Australia 2007	in	N = 28; mean age 79.2 y; 18% with dementia	Aged \geq 65 y	Did not speak English, unable to speak because of medical condition, critically ill at the time	Quality improvement	АМТ, САМ	No	AMT takes up to 5 min to administer
Hirschman ED of a la		N = 829; age 75.7 \pm 7.1 y	Age \geq 65 y, lived within a 30-mile radius of	Had an end-stage disease with		2 validated screening tools: the SIS and	No	Study measures and analyses controlled for no ED-specific
urban, te academic center; b Septemb 2007, and 2008	ertiary c health oetween oer 6,		the ED in the state of Pennsylvania, and lived independently			CLOX1		environmental variables (eg, crowding, time of triage, triage class, location of screening, wait time, etc) in relation to screening cognitive impairment

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Huff 2001*, University hospital ED	N = 205	Aged 55 y or older	Head trauma or multisystem trauma, inability to speak English, educational level of ≤7 y, acute medical illness, or contact or droplet isolation. Additionally, patients that the research assistants felt might be harmed by mental distress or other discomfort by test administration were excluded		Comparison of QCS against MMSE	Yes	MMSE took significantly longer to administer (311 s mean) than did the QCS (141 s mean; <i>P</i> < .01)	
Irons 2002, University hospital ED; June- August 2000	N = 731; age 18- 25 y (16%); 26- 40 y (30%); 41- 60 y (30%); 61- 75 y (13%); >75 y (8%)	. – .		Prospective, cross-sectional	Validation of QCS against MMSE	No		Average administration and scoring time for the QCS is slightly less than 2½ min; QCS requires no written response from the patient
Keles 2001						No		Standardized tests applied briefly and easily are available and these are beneficial in order to identify and treat cognitive disorders of older adults
Kennelly 2012, ED in urban teaching hospital in Ireland		All medical, surgical, and ED physicians involved in the acute care of older patients in the hospital		Cross-sectional	14-item questionnaire administered to assess knowledge skills and attitudes of physicians toward screening of older patients in ED for cognitive deficits		29% felt they lacked expertise to perform screening; 78% thought screening was important	Clinicians reported several limiting factors that restricted

Supplementary Table 2 (continued)

Kennelly 2013,	All medical,	
urban teaching	surgical, and ED	
hospital, ED,	physicians, 76 of	
January-March	97 completed	
2012	survey	
Koita 2010	N/A	N/A

N/A

Article discusses the No process for conducting a mental status examination on a patient in the ED. It mentions SIS, clock drawing, Mini-Cog, Memory Impairment Screen, Brief Alzheimer Screen, 7min screen, and MMSE as tests for neurologic mental status examinations

N/A

The 7-10 min needed to perform the MMSE and the copyright laws pose further barriers for easy ED use. The 1996 US Preventative Services Task Force literature review found the MMSE, Short Test of Mental Status, the Blessed Orientation Memory Concentration Test, and Functional Activities Questionnaire were all equivalent as a screening tool for detecting dementia. These cognitive tests have not been studied in the ED setting, however, and do not have a defined role in the ED at this time. Wilber and colleagues performed a study in the ED setting comparing the MMSE, SIS, and Mini-Cog. The Mini-Cog consists of 3-item recall and clock drawing; SIS consists of 3-item recall and 3-item temporal orientation (ie, day of week, month, and year). When using a cutoff score of \leq 4 in SIS, the SIS proved to be better than the Mini-Cog. In comparison to the MMSE, the SIS had a sensitivity and specificity of 94% and 86%, respectively, whereas the Mini-Cog had a sensitivity and specificity of 75% and 85%, respectively. Initially, Callahan and colleagues found SIS to perform as well as MMSE, but repeat studies have shown that SIS only had a sensitivity of 63% and specificity of 81%. Cognitive assessment in the ED continues to be an area in need of research. (continued on next page)

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Krupp 2018, Germany; acute geriatric department	N = 165			Patients in an acute geriatric department performed the SIS (4 times), the MMSE (2 times), CDT according to Shulman (2 times), the Regensburg verbal fluency test (2 times), and the Montgomery-Åsberg depression rating scale within a period of 16 d. The overall judgment of a physician blinded to the test results served as the reference standard.		No		The SIS closely correlated with the medical judgment (-0.729). The SIS is a valid, reliable short cognitive test. Using a threshold of 5 points, the SIS detects cognitive deficits relevant to daily living with a higher sensitivity than the MMSE with a threshold of 25. The brevity and simple application of the SIS also enable its application outside geriatric wards.
Lague 2018*, Canada; ED; March-July 2015	N = 171; age ≥ 65 y; 76.9 y (SD 8.3); 2%; 0	Age ≥65 y, independent or semiindependent (can perform 5 of the 7 activities of daily living without any help), spent ≥8 h in the ED, were admitted to any hospital ward	Were living in a long- term care facility, were unable to consent, were unable to communicate in French or English, were experiencing an unstable medical condition leading to their admission to the intensive care unit, had a previous diagnosis of severe dementia or any other psychiatric condition, had delirium during their 8-h ED stay	Participants recruited after being in the ED for at least 8 h	0	Yes	The BPQ had good sensitivity but a low specificity for detecting the 3 geriatric syndromes, cognitive impairment, functional impairment, and frailty. The BPQ could be used to flag patients who would benefit from further screening.	

Laguna 1997, ED N = 536; age ≥ 60 Lanata 2014, Rhode N = 23 resident Island Hospital physicians	y Patients aged ≥60 y No exclusion criteri seen in ED	a To check the reliabilit of the usual medica assessment to detec the cognitive deterioration in old adults attended at HED, compared with that performed systematically by means of an evaluation test of cognitive functions Authors reviewed charts for 100 adult patients admitted to medicine and neurology wards; 23 resident physicians were questioned about their use of cognitive screening tools.	í :t er	Cognitive deterioration was not detected in 111 patients (31.5%); it was mild in 147 (41.8%), moderate in 71 (20.2%), and severe in 23 (6.5%). In patients with moderate-severe deterioration according to the OMCT, such a deterioration was detected by the usual medical evaluation in 7% of cases. The mean time in completing the test was 2.6 \pm 0.9 min. An age \geq 80 y was associated with an increased relative risk for detecting moderate-severe cognitive deterioration (1.98; 95% CI, 1.42-2.78; $P < .001$), whereas the discharge diagnosis of respiratory disease was associated with a decrease of the relative risk (0.41, 95% CI 0.19-0.89; $P < .05$) Authors found 67% and 63% of patients evaluated by attendings in the emergency and medicine departments, respectively, did not receive any form of cognitive testing. In addition, 62% of patients evaluated by neurology attendings received cognitive testing. No physician preformed hierarchical, systematic mental status examinations. The most common reason cited by resident physicians for not using standardized cognitive screening tools was lack of time.
Lucke 2017, Leiden $N = 1632$; age \geq University 70 y MedicalCenter (LUMC) and Alrijne Hospital in the Netherlands; ED; N/R	Patients aged ≥70 y N/R visiting ED	The aim was to investigate 6-CIT, Katz ADL if the 6-CIT is an independent predictor of functional decline and mortality. They compared the 6-CIT score with the Katz ADL and assessed mortality and functional decline 3 mo and 1 y post- ED visit.	No	(continued on next page)

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1314.e74

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Lucke 2015, Leiden University Medical Center (LUMC) and Alrijne Hospital in the Netherlands; ED; N/R	70 y; 78.7 y mean	Patients aged ≥70 y visiting ED		A prospective follow-up study among all patients aged ≥70 y presenting to the ED of a university teaching hospital in the Netherlands. Descriptive data including cognition, measured by the 6-CIT was obtained. Follow-up data consisted of 90- d mortality and 90- d functional decline, defined by 1-point increase in Katz ADL score and/or new institutionalization		No		6-CIT is administered in 2- 3 min and measures cognitiv impairment. Impaired cognition (6-CIT score > 9) was significantly associated with both mortality (OR 3.5 95% CI 1.96-6.27, <i>P</i> value < .001) and functional decline (OR 1.75, 95% CI 1.08-2.82, J value .023) after adjustmen for age, gender, level of education, dementia, numb of different medications use at home, and time of arrival
Maxwell 2013, 2 acute care community hospitals	N = 80; 78.7 y mean; 44%; 27	Patients aged ≥65 y visiting ED with a primary injury		institutionalization The Mini-Cog or Informant Questionnaire on Cognitive Decline in the Elderly (IQCDE) and Vulnerable Elder Survey (VES-13) were administered to patients or surrogates.				Cognitive impairment was present in 36 (44%) of patien (abnormal Mini-Cog: 22%; IQCDE > 3.44: 22%). Injured older adults had higher cognitive and preinjury functional impairment than has been reported in other older populations. A combination of brief screening instruments for u with hospitalized injured older adults or surrogates is useful for risk assessment at
Melady 2018	N/A	N/A	N/A	Not a study				clinical management. This article discusses best practices in the ED for care geriatric patients. Mentions screening for cognitive defects and mentions O3DY and bCAM screening tools. Caregiver history is an essential component of ED evaluation of older adults with functional dependence and/or cognitive impairmer

and/or cognitive impairment.

Meldon 2020, academic ED; October 2019 -May 2020 Initial program N = Patients aged ≥65 y 7718; age ≥ 65 y; visiting ED Program N = 1836; age ≥ 65 y; 75.6 y mean	N/R Implantation of an EMR best practices alert for patients aged ≥65 y. Created an EMR alert for patients aged ≥80 y, fall complaint, history of dementia, polypharmacy (≥10 medications recorded), or high ED utilization (>5 visits in 1 y) in addition to a positive delirium screen. For the first part of the study, ED clinicians educated about these risks and about the EMR alert for comprehensive care assessment. Compared the change in comprehensive geriatric assessment pre- or posttraining.	No	The proportion of geriatric evaluations increased a relative 21% (4.3% - 5.2% , $P =$.09). Authors note that the enhanced period occurred during the beginning of the COVID pandemic.
Morley 2013, N = 35 HIV clinic Clinician in HIV clinic o Ireland; hospital and ED clinicians ED	Surveyed clinicians about cognitive screening tools used and factors limiting cognitive assessments in the clinical setting	No	Participants were asked if an assessment of Orientation in Person, Place and Time (OPPT) was an adequate screening tool for detecting HIV dementias. They were presented with the names of other cognitive screening tools and were asked which they had used previously with HIV-positive patients. MMSE, MoCA screen, the Abbreviated Mental Test (AMT) score, the International HIV Dementia Scale (HIVDS), and the Brief Neurocognitive Screen (BNCS). Thirty-four

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percent (n = 12) of respondents felt that OPPT was a sufficient screening tool for cognitive assessment. Respondents found lack of time, exposed environment, and lack of privacy the most limiting factors when performing cognitive assessment on patients who present acutely to the ED. (continued on next page)

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Ayrstad 2018, Norway; ED; October 2017 —May 2018	N = 111; age ≥65 y, 81 y mean	, Patient seen in ED with suspect infection and admitted		ED nurses screened patients with qSOFA and 4AT (rapid screening of alertness, cognition, attention and fluctuation of symptoms). Time spent on 4AT was recorded	qSOFA and 4AT.	Yes	Median time spent on the assessment with 4AT was 2 min (mean 2.6 min). Among 39 patients with a qSOFA point given for altered mental state, 4AT revealed signs of cognitive impairment in 37 (95%). 4AT revealed signs of cognitive impairment in 26 of 72 patients (36%) where qSOFA did not reveal an altered mental state. 4AT is a rapid assessment of cognitive impairment feasible for use in the ER. 4AT improved the assessment of cognitive impairment in patients aged \geq 65 y with suspected infection.	
Ngian 2008, Australia; teaching hospital; January 2004- April 2006	83 y mean (±6.5)	Patient meeting ASET referral criteria: age \geq 70 y, and 2 of the following 5 criteria required to trigger referral: (1) multiple health problems or >3 regular medications, (2) history of falls or fall- related injury, (3) $>$ 3 presentations to ED in the last 6 mo, (4) problems with memory, or (5) patient or caregiver reports recent functional or behavioral change.		Study objectives were to review discordant cases (using EMR)— older adult patients deemed for discharge by ED but subsequently admitted following ASET review. These cases were examined with regard to clinical outcomes. ASET contribution was also reviewed with respect to assessment of cognitive, functional, and mobility status.		No		Assessment of older adult patients by ASET yielded additional information on functional, mobility and cognitive issues that were overlooked by ED.

This is a review article citing the use of the 6-CIT screen in primary care, outpatient care, and EDs Yes

The 6-CIT has been shown to be a fast, feasible method for screening for cognitive impairment in older adults in the ED, with a mean completion time of 1.9 min. In a USbased study involving 163 ED patients (mean age 78 y), the 6-CIT demonstrated excellent sensitivity at 95% and specificity at 65% (AUC = 0.930) for cognitive dysfunction based on MMSE scores of \leq 23. However, this result was achieved using a lower 6-CIT cutoff of 4/5, and there was no randomization between criterion standard testing and screening. Another US research group used the 6-CIT to screen for cognitive impairment in 271 older patients in an urban teaching hospital ED. The psychometric properties of the instrument were not analyzed; however, the researchers claimed to have discovered 46 from a total of 55 cases of cognitive impairment, where no previous history of cognitive impairment existed.

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Ouellet 2016*, Canada; teaching EDs; May 2009 —March 2011	N = 306; age ≥65 y; mean 77.0 ± 7.2 y; 85.3%	Age ≥65 y, be presenting to the ED specifically for a minor traumatic injury (ie, soft tissue/ osseous lesions such as lacerations, contusions, sprains, simple extremity fractures, minor thoracic injuries, or minor head injury), be discharged home within 48 h of the ED visit, be independent in basic activities of daily living in the month prior to the ED visit.		This study aimed at exploring correlates of global cognitive functioning in older adults being evaluated in the context of a consultation in the ED following a minor traumatic injury.	The MoCA was used to assess cognitive function.	No		Results of multivariate analyse: indicate that the variables most strongly associated with lower MoCA scores are being a man, being 85 y or older, having a lower education, being more depressed, being slower in terms of mobility, and reporting serious memory problems.
Salen 2009, US; ED; N/R	N = 100; age ≥65 y; 9%	Age ≥65 y, English- speaking community- dwelling people seen in a community hospital ED		by an inability to correctly perform a CDT in older adult patients presenting to the ED for reasons other than altered mental status. Also sought to assess whether an ED cognitive impairment screening program as reflected by an abnormal CDT prompted further	CDT	Yes	The CDT seems to be a feasible means for identifying older adult ED patients at risk for cognitive disorders. Routine cognitive screening of older adults with the CDT seems to be well accepted by patients and families, but the sporadic follow-up by PCPs suggests a role for more aggressive ED interventions to delineate the causes of abnormal cognitive screening	
Samaras 2010				older adult patients in the	CAM and SIS	No	examinations.	Mentions the need to identify dementia and delirium
Sanders 1995				ED. This is an editorial regarding a Naughton et al article published in the same issue. The editorial mentions the CAM, a standard OMCT MMSE				
Sanders 2007				MMSE The article is a commentary.				

Schnitker 2015*, Australia; ED; 2012-2013	$\begin{array}{l} N=580; age \geq \\ 70 y; 80.3 \pm 6.7 y; \\ 33\% \end{array}$		(1) Stayed >2 h in ED before the research nurse was available to approach them; (2) were severely ill; (3) had consented for the study during a previous ED visit; (4) required an interpreter and where no suitable interpreter could be found in a timely manner (2 h); or (5) who were not able to participate in the planned phone follow-up (7 and 28 d post ED visit)		OMCT	Νο		As it is considered currently, the OMCT is a cognitive screening tool with the most optimal psychometric properties tested (ie, MMSE was used as the reference standard) in the older ED population	1314.e80
Schoenenberger				The article discusses		No			
2014				geriatric screening/ assessment tools: Short					A. N
				blessed test, CAM, Timed					owrc
				up and go, ADL, EGS (discussed in article					Nowroozpoor et
				below)					oor e
Schoenenberger 2014,	N = 1547 (752 control, 795	Age \geq 75 y, ED patient	None	Authors developed a novel multidimensional EGS	The tool met the following	Yes	EGS took <5 min to perform in most (85.8%)		al
Switzerland;	screening); age			tool (has 15 questions).	prerequisites: (1) EGS	;	cases. Of the 70 invited		. / JAMDA
University hospital ED; June	\geq 75 y; 82.8 \pm 5.1 y (control),			ED physicians were trained in its use during	is multidimensional and covers relevant		ED physicians, 41 (64.1%) returned the		MDA
2012–February	$82.7 \pm 5 \text{ y}$			the control period, June-	domains of geriatric		questionnaire that		23
2013	(screening); N/R;			October 2012. October	problems; (2) EGS		asked about their		(202
	N/R			2012—June 2013 was the screening period.	uses validated instruments; and (3)		experience with the EGS. Most responders		(2022) 1314.e31–1314.e88
				sereening periodi	EGS must be feasible		agreed or partially		314.0
					in an ED. The domains were relevant for	;	agreed that EGS domains are suited to		31-
					older ED patients:		detect geriatric		1314
					cognition, falls,		problems: 73.0% agreed		.e88
					mobility, and ADL		or partially agreed for cognition; 77.8%, for		
							falls; 75.0%, for		
							mobility; and 72.2%, for ADL		
								(continued on next page)	

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Shenkin 2019, UK; ED and inpatient; not stated				Participants assessed within 12 h of coming to ED or 96 h as an inpatient	Delirium Rating Scale -Revised-98, CAM, 4AT (www.the4AT. com) 4AT takes <2 min to complete	Yes	Compared the diagnostic accuracy of the 4AT to the other screens for delirium. The 4AT had an AUC of 0.90. The 4AT had specificity of 95% (95% CI 92-97) and sensitivity of 76% (95% CI 61-87). The CAM had specificity of 100% (95% CI 98-100) and sensitivity of 40% (95% CI 26-57). Patients with positive 4AT had longer lengths of stay (median 5 d, IQR 2.0-14.0) than negative 4AT (median 2 d, IQR 1.0-6.0) and higher mortality. Cognitive test items of the 4AT were highly specific (AMT4 score 2:97% 94%-98%); attention score of 2: 98% (96%-99%); but showed lower sensitivity (AMT4 score 2: 47% 32%-62%); attention score of 2: 62% (36%-83%) in detecting existing dementia. Conclusions: The 4AT is a rapid delirium assessment instrument that is feasible in routine care, including with patients with dementia, which has good diagnostic accuracy for delirium for acutely unwell older patients	

The objective was to review published evidence on the Rapid Assessment Interface and Discharge (RAID) service model, examining the strengths and weaknesses of the service design, outcome, and effectiveness. The RAID service has shown quality improvement in the care of older people by reducing their length of stay, avoiding their admission to acute hospital beds, and discharging them in increased numbers back to their original place of residence, rather than an institution or care home. In addition, the RAID model has been shown to reduce the readmission rate after discharge by 65% in comparison with a pre-RAID group. The psychiatric liaison service can support the management of behavioral and psychological symptoms in patients with dementia; an audit of antipsychotic prescriptions for people with dementia has showed a 52% reduction in antipsychotic prescriptions for people with dementia between 2008 and 2011. The RAID service could have contributed to reduced antipsychotic prescriptions, but this was not actually studied as part of the evaluation.

No

The RAID service has shown quality improvement in the care of older people by reducing their length of stay, avoiding their admission to acute hospital beds, and discharging them in increased numbers back to their original place of residence, rather than an institution or care home. In addition, the RAID model has been shown to reduce the readmission rate after discharge by 65% in comparison with a pre-RAID group. The psychiatric liaison service can support the management of behavioral and psychological symptoms in patients with dementia; an audit of antipsychotic prescriptions for people with dementia has showed a 52% reduction in antipsychotic prescriptions for people with dementia between 2008 and 2011. The RAID service could have contributed to reduced antipsychotic prescriptions, but this was not actually studied as part of the evaluation.

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Study, Location, Time Frame (* in PICO 1 and 2)	No. of Patients (Median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design- Timing of Recruitment	Studied	Measured for Feasibility, Pragmatic Nature, Timing, Efficiency, etc (Yes/No)	If Y: Measures Cited for Feasibility, Preferences, Duration of Instrument, Efficiency	If N: Points Made by Study About Ease of Use, Speed (Quick, Fast, etc), Setting, Integration Into Routine Care, etc
Stair 2007*, Urban teaching hospital, ED, June 2002 —October 2003	$N=684,$ age ≥ 18 y, mean 48 ± 18 y, N/R, 0	Age ≥18 y, speak English or Spanish, ability to answer questions		Research assistants would ask the participants, "How many years of school have you completed?" then would flip a coin to determine if MMSE or QCS would be asked first. Time to complete both tests was recorded	MMSE or QCS	Yes	Researchers found that the QCS required less time to complete than the MMSE ($2.7 \pm 1.3 \text{ vs}$ $5.1 \pm 1.9 \text{ mean}$; $P <$.001). Correlation of QCS and MMSE scores was fair, with Pearson r = 0.61 (95% Cl 0.56- 0.66). Conclusions: The QCS can be administered more quickly than the MMSE and is easier to administer in the ED	
Sunkara 2019, NY; ED; March 1–July 1, 2018		ED patients aged ≥75 y likely to be discharged home, English or Spanish speaking		Not reported	Mini-Cog (if participant could answer), IQCODE if participant could not respond	No		Cognitive impairment screening is feasible in the ED and many individuals screen positive. Use of a volunteer workforce may be a feasible interim step to implementing a sustainable program while increasing learners' exposure to positive geriatric care

experiences.

Taylor 2018	 instrument whilst the patient is in any part of the ED setting Clinical assessment tools addressing any aspect of functional ability, and/or cognition assessment The study must include an intervention of any description resulting from the outcome of the instrument administration. There must be a measured outcome as a result of the ED-based 	assessment instrument as the primary outcome •Studies that target an		Scoping review identified 6 measures used for cognitive and delirium screening instruments: AMT, CAM, Blessed Orientation-memory Concentration (BOMC), MMSE, Mini- Cog, Short Portable Status Questionnaire (SPMSQ)			Only 2 of the screens, Mini-Cog and MMSE, have been tested for use in ED. Mini-Cog has a drawing section that is a limitation for use in ED. There was no standard time to administer one of the screens. Authors note that doing them early in a patient's visit would help provide needed information that could impact patient disposition.
Wilber 2006	intervention		Not a study. Article described mental status screening tests		No		MMSE not useful in ED as difficult to preform, patients may have vision, hearing or writing limitation, takes a median of 6 min to do. Article mentioned screens studied for ED use including the
Wilber 2005, 149; age ≥65 y; Summa Health mean age 75 y; System's; ED; fall 23%, 0 2003	Age ≥65 y, English speaking	Unable or unwilling to perform testing, those who were medically unstable, and those who received medications during the study that could affect their mental status.	conducted SIS or Mini- Cog as directed, ≥30 min later an investigator conducted MMSE	SIS, Mini-Cog, MMSE	Yes	SIS agreed with MMSE 88%, and Mini-Cog agreed 83%. Previous study showed patients completed the SIS in <1 min (range, 0.5- 3.5 min) and Mini-Cog took 1.5 min (0.5- 5 min). MMSE takes a median of 5.5 min (range, 3.5-14 min) to complete.	
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Study, Location, Time Frame (* in PICO 1 and 2)	No. of Patients (Median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design- Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Measured for Feasibility, Pragmatic Nature, Timing, Efficiency, etc (Yes/No)	If Y: Measures Cited for Feasibility, Preferences, Duration of Instrument, Efficiency	If N: Points Made by Study About Ease of Use, Speed (Quick, Fast, etc), Setting, Integration Into Routine Care, etc
Wilber 2008*, Summa Health System's Akron City Hospital, Washington University, Barnes-Jewish Hospital, The Cleveland Clinic; ED; January 2006 –January 2007	352 participants, age \geq 65 y, mean age 77 \pm 8; 32%; 0 care partner	Age 65 y; English speaking	Receiving medications that may have affected their mental status (narcotics, antiemetics, or benzodiazepines), were critically ill, were unable to con- sent or cooperate with data acquisition, were previously enrolled, or refused to complete the	At Sites 1 and 3, the SIS was administered first, and the MMSE was administered a minimum of 30 min later. At Site 2, the MMSE was administered first, and the SIS was administered a minimum of 30 min later.	MMSE, SIS	No		Compared sensitivity and specificity of SIS to MMSE. Overall, the SIS was 63% sensitive and 81% specific; the NPV was 83% and the PPV was 60% (Table 1). The overall agreement between the 2 tests was 75%. However, we believe that the SIS, testing temporal orientation and recall, is quick and easy for
			questioning					EPs to incorporate into their physical examination. It provides an objective measure of cognition, as opposed to the unstructured evaluation of cognition by clinical gestalt (often expressed as A&Ox3).

Wilding 2016*, Ontario Canada; ED; January 1, 2010, to August 31, 2010	N = 238; age ≥75 y; mean age 81.9 y; 13.4%; 0	Patients aged ≥75 y with no history of cognitive impairment	cognitively impaired;	MMSE, O3DY, and AFT. MMSE and O3DY were administered followed by AFT	MMSE, O3DY, and AFT	No		The O3DY Scale demonstrated a sensitivity of 93.8% (95% CI 77.8-98.9) and a specificity of 72.8% (95% CI 66.1-78.7). The MMSE and O3DY scale showed agreement in 75.6% of cases. An AFT score <15 demonstrated a sensitivity of 90.6% (95% CI 73.8-97.5) and specificity of 39.3% (95% CI 32.7-46.4). Using a cutoff of <10 for the AFT resulted in a lower sensitivity of 62.5% (95% CI 43.7-78.3) but greater specificity of 78.2% (95% CI 71.8-83.5). The MMSE and the AFT showed agreement in 46.2% and 76.1% of cases with cutoffs of <15 and <10, respectively. The O3DY scale is a feasible screening tool for cognitive impairment in older adult patients presenting to the ED. It is highly practical for use in the time-pressured ED environment, and it does not require paper, pen, or stopwatch. It showed with the MMSE. The AFT did not perform as well, with a much	
Wilkinson 2018, Canada; ED	N = 147; age 70- 94 y			A "Whack-a-mole" style computer game was created to discern inhibition ability in a geriatric population in the ED. The results of the game were then compared to MMSE, MoCA, and CAM evaluations.	had a correlation to MMSE, MOCA, or CAM, determined as,	No	No	diminished specificity. No	
Yamamoto 2019, Japan; hospital ED; October 1, 2014, to September 30, 2015	$\begin{split} N &= 885; \mbox{ age } >50 \ y; \\ mean \ age \ 78.9 \ y; \\ 10\% \ history \ of \\ dementia \ (n = 89, \\ mean \ age \ 85.0 \ \pm \\ 6.53 \ y); \ 0 \ care \\ partner \end{split}$	Non–critically ill patients aged >50 y admitted to the ED	Admitted with critical diseases, receiving sedative medication, unable to consent, or who refused to participate, and those with more than 1 wk of hospitalization	evaluations. Participants approached in ED	Short-term memory recall test (STMT-R) a revised version of the STMT		Short-term memory recall test (STMT-R), The test is normally completed within 2 min, but some participants were unable to complete the questionnaire within 5 min		
								(continued on next page)	

Supplementary	Table 2	(continued)
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Study, Location, Time Frame (* in PICO 1 and 2)	No. of Patients (Median or Mean Age)	Inclusion Criteria	Exclusion Criteria	Study Design- Timing of Recruitment	Intervention: Screening Instrument or Tool Studied	Measured for Feasibility, Pragmatic Nature, Timing, Efficiency, etc (Yes/No)	If Y: Measures Cited for Feasibility, Preferences, Duration of Instrument, Efficiency	If N: Points Made by Study About Ease of Use, Speed (Quick, Fast, etc), Setting, Integration Into Routine Care, etc
Zun 1986, mailed survey	N = 170 Board- certified ED physicians	ED Board-certified physician	N/R	Random sample of 120 of 1174 American Board of Emergency Medicine –certified emergency physicians and a validation group of 50 Board-certified ED physicians were surveyed by questionnaire.	Authors developed a questionnaire to determine the i n d i c a t i o ns, the amount of time necessary to evaluate mental status, the content of the mental status examination (MSE) used, and the ideal characteristics of a short, standardized MSE. The Strub and Black's Composite Mental Status Examination (CMSE) was used as the standard example for answering the	Yes	72% of respondents said they take <5 min on the MSE	No
Zun 1988					questionnaire	No	Ν	Some physicians view the mental status evaluation as a series of odd maneuvers and questions that appear time- consuming and of questionable clinical significance. "Emergency departments are places where physicians hav limited time to examine patients. Texts in emergency medicine have advocated the need to perform formal mental status examinations. However, many physicians find the formal mental statu examination time-consumin and cumbersome." An extensive test is rarely necessary in the ED; rather a short test of cognitive function, such as the Cognitive Capacity Screening Examination or MMSE, may be more appropriate.

4AT, 4 A's Test; AD8, Alzheimer's Disease-8; ADL, activities of daily living; ADRD, Alzheimer's disease and related dementias; AFT, Animal Fluency Test; AMT, Abbreviated Mental Test; APOP, acutely presenting older patient; ASET, Aged Care Service Emergency Teams; AUC, area under the curve; BAS, Brief Alzheimer's Screen; BRIGHT, Brief Risk Identification for Geriatric Health Tool; cAD8, caregiver-completed Alzheimer's Disease-8; CAM-ICU, Confusion Assessment Method—Intensive Care Unit; CDT, Clock-Drawing Test; CFS, clinical frailty scale; ED, emergency department; ECS, Emergency Geriatric Screening; GCS, Glasgow Coma Scale; IQCODE, Informant Questionnaire for Cognitive Decline; LOS, length of stay; MMSE, Mini-Mental State Examination; MoCA, Montreal Cognitive Assessment; N/A, not applicable; NPV, negative predictive value; N/R, not reported; OARS, Older American Resources and Services scale; 03DY, Ottawa 3DY; OMCT, Orientation Memory Concentration Test; PDSA, plan-do-study-act; PPV, positive predictive value; QSPA, Quick Confusion Scale; GOT; Six-Item Cognitive Impairment Test; SIS, Six Item Screener; SMMSE, standardized MMSE; TAP, Treatment Acceptability and Preferences.

Appendix 1. The GEAR 2.0-ADC Network Authors

Names	Degrees
Aggarawal, Neelum	MD
Allore, Heather	PhD
Amy Aloysi	MD, MPH
Belleville, Michael	HS
Bellolio, M Fernanda	MD
Betz, Marian (Emmy)	MD, MPH
Biese, Kevin	MD, MAT
Brandt, Cynthia	MD, MPH
Bruursema, Stacey	LMSW
Carnahan, Ryan	PharmD, MS, BCPP
Carpenter, Christopher	MD, MSC
Carr, David	MD
Chin-Hansen, Jennie	MS, RN, FAAN
Daven, Morgan	MA
Degesys, Nida	MD
Dresden, M Scott	MD, MS
Dussetschleger, Jeffrey	DDS, MPH
Ellenbogen, Michael	AA
Falvey, Jason	DPT, PhD
Foster, Beverley	HS
Gettel, Cameron	MD
Gifford, Angela	MA
Gilmore-Bykovskyi, Andrea	PhD, RN
Goldberg, Elizabeth	MD, ScM
Han, Jin	MD, MSc
Hardy, James	MD
Hastings, S. Nicole	MD
Hirshon, Jon Mark	MD, PhD, MPH
Hoang, Ly	BS
Hogan, Tess	MD
Hung, William	MD, MPH
Hwang, Ula	MD, MPH
Isaacs, Eric	MD
aspal, Naveena	BA
lobe, Deb	BS
Johnson, Jerry Kalha Kathlaan (Katha)	MD
Kelly, Kathleen (Kathy)	MPA
Kennedy, Maura	MD MD BHD
Kind, Amy	MD, PhD
Leggett, Jesseca	BS
Malone, Michael	MD
Moccia, Michelle Moreno. Monica	DNP
· · · · · · · · · · · · · · · · · · ·	BS
Morrow-Howell, Nancy Nowroozpoor, Armin	MSW, PhD MD
Dhuabunwa, Ugochi	MD
Diyemhonian, Brenda	MD, MHSA, MPH
Perry, William	PhD
Prusaczk, Beth	PhD. MSW
	D 4
Resendez, Jason	BA
Rising, Kristen Sano, Mary	MD PhD
Savage, Bob Shah, Manish	HS MD MDH
	MD, MPH
Suyama, Joe	MD, FACEP
Swartzberg, Jeremy	MD
Faylor, Zachary	BS MD MDU
Vaishal, Tolia	MD, MPH
Vann, Allan Wahb, Taraca	EdD
Webb, Teresa Weintraub, Sandra	RN PhD