Effectiveness of acute geriatric unit care on functional decline and process outcomes among older adults admitted to hospital with acute medical complaints: a protocol for a systematic review

Íde O’Shaughnessy, Katie Robinson, Margaret O’Connor, Mairéad Conneely, Damien Ryan, Fiona Steed, Leonora Carey, Aoife Leahy, Rose Galvin

ABSTRACT
Introduction Older adults are clinically heterogeneous and are at increased risk of adverse outcomes during hospitalisation due to the presence of multiple comorbid conditions and reduced homeostatic reserves. Acute geriatric units (AGUs) are units designed with their own physical location and structure, which provide care to older adults during the acute phase of illness and are underpinned by an interdisciplinary comprehensive geriatric assessment model of care. This review aims to update and synthesise the totality of evidence related to the effectiveness of AGU care on clinical and process outcomes among older adults admitted to hospital with acute medical complaints.

Design Updated systematic review and meta-analysis

Methods and analysis MEDLINE, Cumulative Index of Nursing and Allied Health Literature, Controlled Trials in the Cochrane Library and Embase electronic databases will be systematically searched from 2008 to February 2021. Trials with a randomised design that deliver an AGU intervention to older adults admitted to hospital for acute medical complaints will be included. The primary outcome measure will be functional decline at discharge from hospital and at follow-up. Secondary outcomes will include length of stay, cost of index admission, incidence of unscheduled hospital readmission, living at home (the inverse of death or institutionalisation combined; used to describe someone who is in their own home at follow-up), mortality, cognitive function and patient satisfaction with index admission. Title and abstract screening of studies for full-text extraction will be conducted independently by two authors. The Cochrane risk of bias 2 tool will be used to assess the methodological quality of the included trials. The quality of evidence for outcomes reported will be assessed using the Grading of Recommendations Assessment, Development and Evaluations framework. A pooled meta-analysis will be conducted using Review Manager, depending on the uniformity of the data.

Ethics and dissemination Formal ethical approval is not required as all data collected will be secondary data and will be analysed anonymously. The authors will present the findings of the review to a patient and public involvement stakeholder panel of older adults that has been established at the Ageing Research Centre in the University of Limerick. This will enable the views and opinions of older adults to be integrated into the discussion section of the paper.

PROSPERO registration number CRD42021237633.

INTRODUCTION
With population ageing worldwide, the number of older adults attending the emergency department (ED) continues to increase, with evidence of growth in attendances over the past decade beyond that expected from international demographic changes. It is anticipated that the number of adults aged ≥65 years will increase from 1 billion in 2019 to 1.4 billion by 2030 and further increase to 2.1 billion by 2050. The ED is often the main portal of entry to unscheduled care for older adults who account for up to 25% of all attendances. This disproportionate level of ED use by the
older population has significant ramifications related to ED overcrowding\(^1\)\(^4\) and is considered a major patient safety concern as well as being associated with reduced quality of patient care.\(^5\) Older adults are clinically heterogeneous and often present to the ED and acute care services with a non-specific complaint or with classical frailty syndromes, which are often triggered by a minor stressor event\(^6\) such as an infection or a complication associated with introduction of a new medication. Their complex medical and psychosocial needs may complicate their ED care\(^7\)\(^8\) and thus increase their susceptibility to adverse outcomes. Up to 60% of older adults who present to the ED are admitted for inpatient care as demonstrated in a retrospective cohort study of 550 older adults.\(^9\)

There is broad agreement within the literature that hospital admission poses a significant risk for older adults\(^10\) and that system wide reform is required to manage the associated deterioration that occurs in vulnerable older adults due to prolonged exposure to such an environment. Loyd et al reported a 30% (95% CI 24% to 33%) prevalence rate of hospital-associated disability among older adults in their meta-analysis of 15 longitudinal studies of older adults hospitalised in acute care.\(^11\) The reasons underlying older adults higher rate of adverse outcomes during hospitalisation are multifaceted; however, the most potent intrinsic factor is the clinical condition of frailty.\(^12\)\(^13\) The concept of frailty suggests that the accumulation of health deficits is more significant in contributing to vulnerability than specific conditions or physical limitations.\(^14\) The presence of diminished homeostatic reserves and multiple comorbid conditions leaves older adults more vulnerable to functional decline and serious sequelae.\(^15\)

Changes to organisational structure and processes are therefore required to better meet the needs of older adults within acute care.\(^16\) Underpinning all the innovative and evidenced based changes in the way care is delivered is comprehensive geriatric assessment (CGA), which is considered more effective than usual care for frail older adults.\(^17\) CGA was first coined by Rubenstein et al and is defined as a ‘multidimensional, interdisciplinary diagnostic process to determine the medical, psychological and functional capabilities of a frail older person in order to develop a co-ordinated and integrated plan for treatment and long-term follow-up.’\(^18\) Social and nutritional assessments have recently been recognised as important dimensions of CGA.\(^19\) CGA is both therapeutic and diagnostic and emphasises improving the quality of life and functional status of frail older adults and at the same time, improving prognosis and outcomes.\(^16\) Ellis et al conducted a Cochrane review of 29 trials and 13 766 participants, which synthesised the evidence on the effectiveness and resource use of CGA for older adults admitted to hospital.\(^17\) Older adults who receive CGA are more likely to be alive and in their own homes after an emergency admission to hospital and are less likely to be admitted to a nursing home at discharge or at three to 12 months’ follow-up. However, the authors were unable to determine whether the results show a difference in effect between discrete specialised wards and mobile multidisciplinary teams across several wards as the analysis was underpowered. Therefore, greater understanding of the the specific impact and organisational forms of CGA delivery in acute services are therefore required to advance the evidence base. Dedicated acute geriatric units (AGU), are units designed with their own physical location and structure, which provide care to older adults during admission to hospital for an acute medical illness including acute exacerbations of chronic diseases.\(^20\) While variations in the definition exist\(^21\) all are based on the seminal research on AGUs.\(^22\)\(^23\) Furthermore, inclusive of all definitions are specialised interdisciplinary teams who embed geriatric competencies into their practice.

The effectiveness of AGU care for older adults admitted to hospital with acute medical disorders was previously examined in a systematic review and meta-analysis of five RCT, four non-randomised trials and two case–control studies.\(^24\) The five RCTs dated from 1985 to 2000 and found that admission to an AGU conferred a lower risk of functional decline at discharge and older adults were more likely to be living at home after discharge when compared with conventional care units. However, only two of the five RCTs reported on functional decline at discharge; one RCT presented results at 3-month follow-up, with no differences in the incidence of functional decline reported between groups. Meta-analysis of case fatality either in hospital or at 3-month follow-up did not show any significant differences between groups. Furthermore, authors were unable to draw firm conclusions on length of stay (LoS) between groups due to heterogeneity among trials that reported on this outcome. Further studies on caring for older adults in an AGU have shown lower incidence of adverse outcomes such as delirium\(^24\) and association with trends of lower LoS and greater cost effectiveness when compared with conventional care units.\(^25\)\(^27\) More recently, the impact of AGU care has focused on health-related quality of life (HRQoL) and functional decline after discharge from hospital.\(^24\) A prospective controlled trial where an AGU intervention was provided to 206 frail older adults, aged ≥75 years, in need of acute in-patient treatment when compared with routine care (n=202) found that older adults in the intervention group were less likely to present with decline in HRQoL and in activities of daily living (ADL) at 3-month follow-up.

Our review aims to update and synthesise the totality of recent research evidence related to the effectiveness of AGUs among older adults admitted to hospital with acute medical complaints. We hypothesise that older adults admitted to an AGU experience less functional decline and more favourable process outcomes when compared with conventional care units.

METHODS AND ANALYSIS

Study design

This protocol for a systematic review will be conducted in accordance with the Preferred Reporting of Items for Systematic Reviews and Meta-Analyses (PRISMA) Protocols guidelines.\(^28\) A template will be constructed outlining...
the essential items and components of the protocol as provided in online supplemental file 1. The systematic review and meta-analysis will comply with the reporting guidance outlined in 27-item PRISMA checklist. The methodology for the review will be underpinned by the Cochrane handbook for systematic reviews of interventions. To ensure scientific rigour when assessing intervention effects, we have chosen to only include trials with a randomised design. Randomised controlled trials (RCTs) including cluster trials and quasi-RCTs will also be included in the systematic review.

**Search strategy**

Searches were carried out in the following electronic databases—MEDLINE, Cumulative Index of Nursing and Allied Health Literature, Cochrane Central Register of Controlled Trials in the Cochrane Library and Embase. A search string was developed using the keywords in the systematic review and meta-analysis published in 2009. MeSH terms and associated keywords covering the topics of older adults, AGU, CGA and RCT were applied to relevant databases outlined above. An example search strategy from MEDLINE in EbSCO are presented in online supplemental file 2. Studies were limited from the year 2008 onwards as this review aims to update a previous meta-analysis. Trials included in the previous version of the review will be integrated into the new evidence found. The reference lists of all potentially eligible studies will be hand searched for additional papers. Studies in all languages will be included; the authors will seek translation for studies published in languages other than English.

**Eligibility criteria**

Trials will be included that meet the following inclusion criteria.

**Population**

Older adults (>65 years) admitted to the acute care setting for medical reasons.

**Intervention**

In-keeping with the definition used by Baztán et al in the previous meta-analysis, the intervention comprises an AGU intervention delivered by interdisciplinary teams during the acute phase of illness to prevent functional decline and related complications in older adults admitted to the acute care setting. The 2017 Cochrane report by Ellis et al will be used as the reference standard when describing the components of CGA across the included trials. In their report they outline the following components: clinical leadership, structured assessment, multidisciplinary team meetings, goal setting, involving patients and carers in goal setting, outpatient follow-up, ward environment, adequate time, specialist knowledge, experience, competence and tailoring treatment plans to the individual.

**Comparison**

Usual care, other non-AGU interventions such as admission to acute medical wards.

**Exclusion criteria**

Trials will be excluded if their population is <65 years and if the intervention is aimed at specific medical or surgical complaints or specialty units such as stroke or orthogeriatrics. To ensure we did not include trials that evaluate interventions in the sub-acute phase, exclusions will apply where patients have been transferred from other specialty units such as intensive care to an AGU or admitted to an AGU three or more days after a hospital admission.

**Outcome variables**

The primary outcome measure will be functional decline on discharge from hospital and at follow-up. Functional decline will be defined as loss of independence in one or more ADLs compared with the status prior to admission. Various ADL indices such as the Katz Index of Independence in ADL and Barthel Index, which use a numerical scale to measure performance in ADLs will be included in our meta-analysis. Secondary outcomes will include LoS, living at home (the inverse of death or institutionalisation combined; used to describe someone who is in their own home at follow-up), mortality, cost of index admission, incidence of unscheduled hospital readmission, cognitive function and patient satisfaction with the index admission. For the outcome living at home, we will not measure the proportion of patients in receipt of personal assistance and/or services. All secondary outcomes will be recorded at discharge from hospital and at follow-up periods reported in trials.

**Study selection and data extraction**

Screening

References generated from the search strategy will be exported into Endnote software and duplicates deleted. Two authors (IO’S and RG) will independently screen relevant studies by title and abstract for eligibility. Studies that are selected by the reviewers as meeting the inclusion criteria will undergo a full text review. If a disagreement about eligibility arises, both authors will meet to come to a consensus. Where consensus cannot be reached, third and fourth authors will be consulted (KR and MO'C).

Data synthesis and analysis

Data will be independently extracted from the relevant trials by two reviewers (IO’S and RG); data from trials in the previous version of the review will also be extracted. The information compiled will include trial authors, year of publication, study population, sample size, interventions provided, controls provided, outcomes measured and duration of follow-up. Data will be gathered into a preprepared Microsoft Excel document. A pooled meta-analyses will be carried out where the data are homogenous, which will be determined by the outcomes measured and the time points accessed across the included trials. For the primary outcome of functional decline, we will calculate risk ratios with a 95% CI to determine the intervention effect. The same approach will be applied for all dichotomous secondary outcome measures. For
continuous outcomes (length of hospital stay and cost of index admission) we will calculate the intervention effect using mean differences (MD) and 95% CI where trials all used the same method of measurement. Standardised MD and 95% CI will be applied where trials used different methods of measurement. The median and IQR will be used in the event that the mean and SD are not reported. Authors will be contacted where data are not available. Data for the meta-analyses will be analysed using Review Manager V.5.4. (Cochrane Collaboration).

We will explore heterogeneity across the trials by visually inspecting the forest plots and the associated I² statistics. We will consider I² > 50% as significant heterogeneity. If I² is ≤ 50% we will apply a fixed-effects method. If I² is greater than 50%, we will explore the individual trial characteristics to identify potential sources of heterogeneity, using preplanned subgroup analyses. Where there is substantial heterogeneity we will perform a meta-analysis using both fixed-effects and random-effects models and we will present the most conservative outcome. We will conduct separate subgroup analyses after excluding trials that were conducted in the previous version of the review (those published in or before year 2000).

We will conduct a sensitivity analysis to explore the effect of the methodological features on outcomes: (1) randomisation process (data will be reanalysed excluding trials with inadequate or unclear allocation concealment) and (2) bias in the measurement of the outcome (reanalysis of trials without evidence of or with unclear masking of outcome assessor).

Quality assessment
Trials that meet the inclusion criteria, inclusive of trials in the previous version of the review, will be assessed for risk of bias (RoB) using the Cochrane RoB 2 tool. Two independent reviewers (IO'S and RG) will assess each trial’s RoB under the following domains: randomisation process, deviation from intended intervention, missing outcome data, outcome measurement, selective reporting and the overall risk of bias. If a disagreement about RoB arises, both authors will meet to come to a consensus. Where consensus cannot be reached, third and fourth authors will be consulted (KR and MO'C). The Grading of Recommendations, Assessment, Development and Evaluations framework will be used to assess the quality of evidence for outcomes reported and to summarise data narratively. Outcomes will be graded at one of four levels of evidence—very low certainty, low certainty, moderate certainty and high certainty.

Patient and public involvement
The authors will present the findings of the review to a patient and public involvement stakeholder panel of older adults that has been established at the Ageing Research Centre in the University of Limerick https://www.ul.ie/hri/themes/public-and-patient-involvement-ppi. The focus of this session will be to discuss the findings with this group so that the discussion section of the paper can integrate the views and opinions of older adults.

DISCUSSION
This review will update and synthesise the evidence relating to the effectiveness of AGU care on patient and process outcomes for older adults who are admitted to hospital with acute medical complaints. The clinical and social complexity of hospitalised older adults is having an impact on delivery of healthcare services internationally, and related care is considered a complex intervention; therefore, it is proposed that this review will identify the characteristics and components of CGA within an AGU in terms of the intervention, staff profile and resources. The 10 elements of CGA reported by trialists in the 2017 Cochrane review will be used as a reference standard. This will have relevance for clinicians and policy-makers and will enable recommendations to be made regarding current and future AGU establishment following evidence-based research. We chose functional decline as our primary outcome as maintaining independence in the performance of ADLs is an important determinant of quality of life for older adults.

By synthesising the evidence surrounding AGU care for older adults with acute medical complaints, there is potential for a reduction in a patient’s LoS as is the case in orthogeriatric care. Reducing LoS can potentially preserve an older adult’s functional status and thus reduce the risk of increased morbidity and mortality. Subsequently, this can save hospital bed days and overall reduce hospital costs while enabling older adults to live in their community safely for longer.

Ethics and dissemination
Formal ethical approval is not required for the review as all data collected will be secondary data and will be analysed anonymously. The findings of this review will be disseminated through publication in a peer-review journal and presented at relevant conferences.

Study status
Database searches have been completed.

Author affiliations
1School of Allied Health, Faculty of Education and Health Sciences, Ageing Research Centre, Health Research Institute, University of Limerick, Ireland, Limerick, Ireland
2Department of Ageing and Therapeutics, University Hospital Limerick, Dooradoyle, Limerick, Ireland
3Limerick EM Education Research Training (ALERT), Emergency Department, University Hospital Limerick, Dooradoyle, Limerick, Ireland
4Department of Medicine, University Hospital Limerick, Dooradoyle, Limerick, Ireland
5Department of Occupational Therapy, University Hospital Limerick, Dooradoyle, Limerick, Ireland

Contributors IO’S and RG were major contributors in writing the protocol. KR, MO'C and MC contributed to the planning and conceptualisation of the study design, search strategy, and outcome variables. IO’S, RG, KR, MO'C, MC, DR, FS, LC and AL participated in the study design and critically appraised and edited the
# PRISMA-P Reporting Guidelines Checklist

<table>
<thead>
<tr>
<th>Section and topic</th>
<th>Item No</th>
<th>Checklist item</th>
<th>Location where item is reported</th>
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<tbody>
<tr>
<td>ADMINISTRATIVE INFORMATION</td>
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<td>Title:</td>
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<td>Identification</td>
<td>1a</td>
<td>Identify the report as a protocol of a systematic review</td>
<td>Page 1 &amp; 6</td>
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<tr>
<td>Update</td>
<td>1b</td>
<td>If the protocol is for an update of a previous systematic review, identify as such</td>
<td>Page 6 &amp; 7</td>
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<tr>
<td>Registration</td>
<td>2</td>
<td>If registered, provide the name of the registry (such as PROSPERO) and registration number</td>
<td>Page 3</td>
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<td>Authors:</td>
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<tr>
<td>Contact</td>
<td>3a</td>
<td>Provide name, institutional affiliation, e-mail address of all protocol authors; provide physical mailing address of corresponding author</td>
<td>Page 1</td>
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<tr>
<td>Contributions</td>
<td>3b</td>
<td>Describe contributions of protocol authors and identify the guarantor of the review</td>
<td>Page 1 &amp;10</td>
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<td>Amendments</td>
<td>4</td>
<td>If the protocol represents an amendment of a previously completed OR published protocol, identify as such and list changes; otherwise, state plan for documenting important protocol amendments</td>
<td>Page 6</td>
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<td>Support:</td>
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<td>Sources</td>
<td>5a</td>
<td>Indicate sources of financial OR other support for the review</td>
<td>Page 11</td>
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<td>Sponsor</td>
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<td>Provide name for the review funder and/or sponsor</td>
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<td>Role of sponsor OR funder</td>
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<td>Describe roles of funder(s), sponsor(s), and/or institution(s), if any, in developing the protocol</td>
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<tr>
<td>INTRODUCTION</td>
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<td>Rationale</td>
<td>6</td>
<td>Describe the rationale for the review in the context of what is already known</td>
<td>Page 4-6</td>
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<tr>
<td>Objectives</td>
<td>7</td>
<td>Provide an explicit statement of the question(s) the review will address with reference to participants, interventions, comparators, and outcomes (PICO)</td>
<td>Page 6</td>
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<tr>
<td>METHODS</td>
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<tr>
<td>Eligibility criteria</td>
<td>8</td>
<td>Specify the study characteristics (such as PICO, study design, setting, time frame) and report</td>
<td>Page 7</td>
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<td>Section</td>
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<tr>
<td>Information sources</td>
<td>9</td>
<td>Describe all intended information sources (such as electronic databases, contact with study authors, trial registers OR other grey literature sources) with planned dates of coverage</td>
<td>Page 6 &amp; 7</td>
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<tr>
<td>Search strategy</td>
<td>10</td>
<td>Present draft of search strategy to be used for at least one electronic database, including planned limits, such that it could be repeated</td>
<td>Supplementary file 2</td>
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<td>Study records:</td>
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<tr>
<td>Data management</td>
<td>11a</td>
<td>Describe the mechanism(s) that will be used to manage records and data throughout the review</td>
<td>Page 8</td>
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<tr>
<td>Selection process</td>
<td>11b</td>
<td>State the process that will be used for selecting studies (such as two independent reviewers) through each phase of the review (that is, screening, eligibility and inclusion in meta-analysis)</td>
<td>Page 8</td>
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<tr>
<td>Data collection process</td>
<td>11c</td>
<td>Describe planned method of extracting data from reports (such as piloting forms, done independently, in duplicate), any processes for obtaining and confirming data from investigators</td>
<td>Page 8</td>
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<tr>
<td>Data items</td>
<td>12</td>
<td>List and define all variables for which data will be sought (such as PICO items, funding sources), any pre-planned data assumptions and simplifications</td>
<td>Page 7 &amp; 8</td>
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<tr>
<td>Outcomes and prioritization</td>
<td>13</td>
<td>List and define all outcomes for which data will be sought, including prioritization of main and additional outcomes, with rationale</td>
<td>Page 7 &amp; 8</td>
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<tr>
<td>Risk of bias in individual studies</td>
<td>14</td>
<td>Describe anticipated methods for assessing risk of bias of individual studies, including whether this will be done at the outcome OR study level, OR both; state how this information will be used in data synthesis</td>
<td>Page 9</td>
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<tr>
<td>Data synthesis</td>
<td>15a</td>
<td>Describe criteria under which study data will be quantitatively synthesised</td>
<td>Page 8 &amp; 9</td>
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<td>15b</td>
<td>If data are appropriate for quantitative synthesis, describe planned summary measures, methods of handling data and methods of combining data from studies, including any planned exploration of consistency (such as I^2, Kendall’s τ)</td>
<td>Page 8 &amp; 9</td>
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<td>15c</td>
<td>Describe any proposed additional analyses (such as sensitivity OR subgroup analyses, meta-regression)</td>
<td>Page 9</td>
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<td>15d</td>
<td>If quantitative synthesis is not appropriate, describe the type of summary planned</td>
<td>Page 9</td>
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<td>Meta-bias(es)</td>
<td>16</td>
<td>Specify any planned assessment of meta-bias(es)</td>
<td>Page 9</td>
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<td>Confidence in cumulative evidence</td>
<td>Describe how the strength of the body of evidence will be assessed (such as GRADE)</td>
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**MEDLINE Search Strategy**

1. exp Aged/
2. exp Aging/
3. (older adults OR elderly OR geriatric* OR aging OR senior* OR older people OR aged 65 OR 65+) .ti,ab.
4. (1. OR 2. OR 3.)
5. exp Geriatric Assessment/
6. Geriatric Assessment .ti,ab.
7. exp Health Services for the Aged/
8. exp Comprehensive Health Care/
9. (5. OR 6. OR 7. OR 8.)
10. (geriatric unit OR specialist geriatric OR acute geriatric) .ti,ab.
11. ((elder* OR older OR geriatric* OR aged) adj3 (unit or ward)) .ti,ab.
12. ((unit* OR ward*) adj3 (geriatric* OR elder* OR aged OR older)) .ti,ab.
13. acute care for elder*. .ti,ab.
15. ( 10. OR 11. OR 12. OR 13. OR 14.)
16. 4 AND 9 AND 15
17. (randomized controlled trial OR controlled clinical trial) .pt.
18. (randomized OR randomised OR randomization OR randomisation) .ti,ab.
19. placebo .ti,ab.
20. random*. .ti,ab.
21. trial .ti,ab.
23. (comparative study OR clinical trial) .pt.
24. clinical trial*. .ti,ab.
25. intervention stud* .ti,ab.
26. follow up .ti,ab.
27. comparative stud*. .ti,ab.
28. (nonrandom OR non-random) .ti,ab.
29. (nonrandomized OR nonrandomised OR non-randomised) .ti,ab.
30. quasiexperiment*. .ti,ab.
31. (quasi-random* OR quasirandom*) .ti,ab.
32. (quasi-control* OR quasicontrol*) .ti,ab.
33. ((controlled) AND (trial OR study)) .ti,ab.
34. (pre-test OR pretest OR post-test OR posttest) .ti,ab.
35. OR/16-33
36. S16 OR S17 OR S18 OR S19 OR S20 OR S21 OR S22 OR S23 OR S24 OR S25 OR S26 OR S27 OR S28 OR S29 OR S30 OR S31 OR S32 OR S33 OR 34
37. 16 AND 36