Complex advance care planning interventions for chronic serious illness: how do they work: a scoping review

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ABSTRACT
Context Advance care planning (ACP) interventions have the potential to improve outcomes for patients with chronic serious illness. Yet the rationale for outcome choices and the mechanisms by which outcomes are achieved are not always clear.

Objectives To identify and map proposed mechanisms on how complex ACP interventions can impact outcomes for patients with chronic serious illness and to explore factors that might explain intervention outcomes.

Methods This is a scoping review of randomised controlled trials of complex ACP interventions for patients with chronic serious illness which explicitly stated the mechanism(s) by which the intervention was thought to work. We searched six databases and hand-searched key journals and reference lists.

Results Inclusion yielded 16 articles. Inclusion procedures and mapping of mechanisms and outcomes indicated that causality between components and outcomes was not always clearly described. Tailoring intervention content to patients’ needs was linked to the greatest number of different outcome categories, while promoting competence and confidence to engage in ACP was most often explicitly linked to a primary outcome. Three main factors which might have affected intended outcomes were identified: participant characteristics, such as illness experience or cultural differences; the setting of implementation; or methodological limitations of the study.

Conclusion Findings highlighted two main points of consideration for future ACP intervention studies: the need for clearly stated logic in how interventions are expected to impact primary outcomes and the importance of considering how an intervention may function for patients with chronic serious illnesses within a specific setting.

Key messages
What was already known?
► Examining how advance care planning (ACP) interventions are currently proposed to affect outcomes can guide expectations of what ACP can achieve.

What are the new findings?
► Building skills and confidence was most often used to explicitly state how the intervention would affect outcomes.
► Contextual factors considered a priori mainly related to sociocultural factors and racial disparities, and other factors were considered in light of the results obtained.

What is their significance?
► Intervention rationales are identified, but further clarity is needed regarding how components operate.
► Participants, setting and study design effects should be considered a priori.

INTRODUCTION
Advance care planning (ACP) refers to a process which ‘enables individuals to define goals and preferences for future medical treatment and care, to discuss these goals and preferences with family and health-care providers, and to record and review these preferences if appropriate’. 1 This can, but not necessarily must, entail or result in the completion of an advance directive (AD) specifying care wishes, which can be used in the event the patient loses decisional capacity. It can also involve the appointment of a surrogate decision maker (SDM) who makes decisions about medical care in the patient’s stead. By communicating with health professionals, informal caregivers, family members and loved ones, patients engaging in ACP can make their wishes...
for future health care known. Additionally, ACP can help patients and their SDM be better prepared to make the best possible in-the-moment care decisions together with clinicians.  

Studies of interventions designed to implement or improve ACP, as well as reviews of the ACP research literature, demonstrate a large degree of heterogeneity in outcomes and evidence of effectiveness, and suggest that complex interventions may be more effective in meeting patient preferences for end-of-life care. The spectrum of outcomes ranges from proximal outcomes, such as ACP engagement, to distal outcomes such as concordance of care with patient wishes. For many of these outcomes, reviews report mixed findings. Most recently, a scoping review of ACP randomised controlled trials (RCTs) mapped the outcomes of interventions according to a standardised ACP Outcome Framework. This review found a greater percentage of positive results for outcome domains related to processes (such as knowledge) and actions (such as completion of an AD) than for quality of care, health status and healthcare utilisation. As a result, the reviewers recommended further research to tailor interventions to specific contexts and to set appropriate expectations of ACP outcomes. Taken together, this highlights the current state of the scientific discourse around ACP in terms of its conceptualisation and its goals. Indeed, questions have been posed about whether ACP has the capacity to address what has been defined by a Delphi panel as its most important outcome, namely goal-concordant care, leading to discussions regarding what the focus of ACP research ought to be. The workings of ACP as a process are complex, but this complexity may not always be addressed in the research literature and may be overlooked when interventions and outcomes are considered discretely. To set appropriate expectations for ACP interventions, it is important to consider not only the content of an intervention, but also how the intervention is expected to lead to the outcomes of interest. A recent review of ACP interventions for patients with cancer indicated that studies propose different mechanisms by which ACP interventions are expected to affect outcomes, but that not every paper does so in equal detail. This implies that adequate attention for the rationale of ACP interventions may be lacking, at least in the reporting of these interventions. In the middle of a debate where the benefit of continuing ACP research is coming into question, this is especially concerning. In a response to Dr Morrison’s notes from the editor, call for further work to develop evidence-based conceptual models of ACP. This response further notes that the intervention logic must also be considered within the system where the intervention is implemented. To our knowledge, however, no review of ACP interventions exists for chronic serious illness that specifically investigates not only intervention components and outcomes, but also the mechanisms by which the intervention is hypothesised to affect said outcomes—that is, what do authors expect the intervention will achieve and how? Neither has a review examined whether and how the authors of these articles explain the study findings that were obtained—for example, if the intervention did not achieve the expected results, is this due to factors intrinsic to the intervention; do systemic factors preclude intervention impact; or should the hypothesised mechanism be re-evaluated to more accurately reflect what ACP can accomplish?

Insights in these fields will aid the development of new interventions to facilitate ACP or the refining of existing initiatives. This is made possible by first identifying pathways from interventions to outcomes via hypothesised mechanisms. Then we can assess which pathways yield positive results, while also highlighting contextual and implementation factors that should be considered. Altogether, this allows us to identify gaps in the current research at different ‘links’ within this ‘chain’.

The purpose of the present scoping review was to identify which mechanisms are proposed to explain how complex ACP interventions, tested for effectiveness through an RCT, are expected to impact outcomes for patients with serious illness, to establish the factors authors refer to in order to explain the study findings, and to map the available evidence.

The research questions can be summarised as follows:

- What are the core components of the intervention?
- Which primary outcome(s) is/are chosen to evaluate the effectiveness of the intervention?
- What is the mechanism by which authors propose the intervention will work to generate change in outcomes?
- What are the results for the primary outcome(s)?
- Which factors contributed to intervention success or failure to affect the primary outcome(s), according to the authors?

**METHODS**

**Study design**

To address the aims of this study, a scoping review design was considered most appropriate. A scoping review is a process of mapping and describing the existing literature, which can be undertaken when the area of study in question is complex and/or heterogeneous in methods or in discipline. This scoping review was conducted according to the methodological framework set forth by Arksey and O’Malley and the additional recommendations by Levac et al. The framework recommends the following steps: (1) identifying the research question; (2) identifying relevant studies; (3) study selection; (4) charting the data; and (5) collating, summarising and reporting the results. The Preferred Reporting Items for Systematic Reviews and Meta-Analyses extension for Scoping Reviews...
checklist (PRISMA-ScR)\textsuperscript{26} was followed for writing the present review report.

Search strategy
We conducted an online search of the following databases: PubMed, PsycINFO, MEDLINE, Embase, Cochrane Central Register of Controlled Trials (CENTRAL) and the Cumulative Index to Nursing and Allied Health Literature (CINAHL). Key terms, Medical Subject Headings (MeSH), Embtree headings and CINAHL headings were used in their applicable databases. The search was first conducted in March 2020 using a search strategy informed by terms and keywords used in existing systematic reviews of ACP.\textsuperscript{3–6} This strategy underwent piloting to ensure the search was sensitive enough to capture key articles, yet specific enough to avoid capturing large numbers of irrelevant articles (see online supplemental figure 1 for an example search strategy). The search was last repeated on 18 November 2020 to identify additional articles that may have been published since the initial search. We included studies that were published in the English language from January 2010 to the date of the last search. Filters were applied for publication type (journal articles, articles in press) where possible.

The references of key articles were screened for additional studies to include. We also hand-searched the most recent issues of the following key journals: Journal of Palliative Care, Journal of Palliative Medicine, Journal of Pain and Symptom Management, Journal of the American Medical Association, BMJ Supportive & Palliative Care, and Palliative Medicine.

Eligibility criteria
Papers were considered for inclusion based on the criteria specified in table 1.

A primary determinant of intervention complexity was the number of components in the intervention, which could be delivered separately or as a package. Interventions with two or more components were considered complex. Interventions consisting of one facilitated discussion were also considered complex due to the flexibility permitted and the training required of the person facilitating the discussion.\textsuperscript{20}

The study introduction and methods sections were read closely multiple times to ascertain whether the study explicitly states or describes the mechanism(s) by which the intervention was expected to generate changes in outcomes.

Methodological quality was not considered an exclusion criterion as we sought to provide an overview from RCTs of complex ACP interventions for chronic serious illness.

Selection of sources of evidence
All retrieved articles were uploaded to Zotero reference manager. Duplicates of retrieved records were removed. Titles and abstracts of all identified articles were screened against the eligibility criteria by JS using a standardised form. Articles identified as potentially relevant were retrieved in full. Full-text screening against the eligibility criteria was completed by JS. During both steps of the screening process, cases of doubt were checked by a second researcher (ADV). During title and abstract screening, if there was unresolved ambiguity, the article was retained for full-text review. In the case of ambiguity in the full-text screening step, a third researcher (KP) was consulted for discussion to achieve consensus.

Data charting process
The following characteristics of the study were extracted using a standardised data charting sheet, which was developed through iterative discussions between the authors: study authors, publication year, country, study setting and sample characteristics. Data extracted regarding the study design included conceptual model or theory used (if any), core intervention components, control condition, duration of the intervention, primary outcome(s), mechanisms and primary findings. Each article was examined for the implications of the results for the proposed mechanism(s) and/or factors which were considered to have impacted the results (eg, contextual factors). Relevant text excerpts were extracted. Two authors (JS and ADV) cross-checked the data from a small sample of studies to achieve consensus regarding the extracted text, after

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\hline
\textbf{Table 1 Inclusion and exclusion criteria} \\
\hline
\textbf{Inclusion criteria} & \textbf{Exclusion criteria} \\
\hline
\begin{itemize}
\item Concerns a peer-reviewed article. \\
\item Reported in English. \\
\item Of primary research. \\
\item Reporting on the quantitative primary outcome(s). \\
\item Of a randomised controlled trial. \\
\item Of a complex advance care planning intervention. \\
\item For chronic serious illness. \\
\item In a sample of adults (>18 years old). \\
\item Where the mechanism(s) by which the intervention is expected to generate changes in outcomes is/are described. \\
\item Conducted in any setting. \\
\end{itemize} & \\
\hline
\begin{itemize}
\item Published prior to 1 January 2010. \\
\item Does not state which outcome is considered the primary outcome. \\
\item Reports feasibility and/or satisfaction outcomes only. \\
\item Reports on a secondary analysis of a randomised controlled trial only. \\
\item Reports on a trial where advance care planning (ACP) is embedded in a broader intervention, such as palliative care interventions that also include pain or symptom management, so that it is not possible to distinguish the discrete effects of the ACP component. \\
\item Reports on psychiatric advance directives and/or crisis planning. \\
\item Reports a study protocol only. \\
\end{itemize} & \\
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which one author (JS) proceeded with the remaining charting.

Synthesis of results
After data extraction, we used descriptive statistics to describe the included studies. Relevant excerpts were imported into NVivo for qualitative analysis. In the first step, open codes were applied to the extracted text regarding intervention components, mechanisms and factors impacting the effect of the intervention on the primary outcome(s). The codes were then combined into descriptive themes. Mechanisms and the outcomes they were expected to generate were mapped in order to indicate which retained articles specified links between mechanisms and outcomes.

RESULTS
Of 1535 articles identified, 16 met the eligibility criteria and were retained for analysis (figure 1).

Study characteristics
The characteristics of the retained studies can be found in online supplemental table 1.

Ten studies were conducted in the USA, two each in the Netherlands and Australia, and one each in Belgium and Northern Ireland. Participants included patients with cancer, dementia, geriatric frailty, end-stage renal disease, heart failure, HIV and comorbid chronic serious illnesses. Nine studies included dyads of patients and a family carer or SDM.

Intervention components
Of the 16 studies retained (online supplemental table 1), the largest portion (11 studies) made use of an interview or conversation to address the topic of ACP.28–38 Conversations were led by facilitators including (trained) nurses,28 29 32 34–36 38 psychologists,31 32 37 social workers31–33 or graduate students.31 32 Five conversation interventions also included the provision of conversation summaries and/or ADs completed during the conversation to the healthcare provider (general practitioner or treating physician) or placement of documentation in the patient’s health record.28 29 31 32 36

Eight studies made use of an AD or goals of care form, which was presented for the patient to read or completed during an interview or discussion.28 31 32 34 36 39–41 Six studies used informational material in online repository, pamphlet or video format.34 37–41 Four studies used question prompts and conversation openers for patients or communication tips for health providers,29 38 41 42 with two of these studies also including instructions for health providers to facilitate the discussion or endorse use of question prompts.29 38 Three studies used an interactive decision aid in website format for patients.39–41 Two interventions used interactive educational workshops for general practitioners or nursing home staff.42 43 One intervention provided ACP education to participants, but the content of the education was not specified.33

What are the primary outcomes and by which mechanism do authors propose they are achieved?
The outcomes and the mechanisms by which they were proposed to be achieved are mapped in figure 2.

In total, nine types of primary outcome were identified: decisional conflict, dyad congruence, AD or Do-Not-Resuscitate (DNR) order completion, psychospiritual well-being, quality of life, number of conversations/consultations, content of discussions with health providers, patient activation and place of death.

Seven descriptive themes were identified to describe the mechanisms by which interventions were proposed to affect the primary outcome(s): (1) tailoring the intervention to the patient recipient (their coping style, health literacy, desire for information, beliefs and misconceptions) to make the intervention more understandable and acceptable; (2) educating patients and carers/surrogates about the course of illness, treatment options and styles of shared decision making in order to facilitate better understanding of these topics; (3) encouraging active involvement of carers/surrogates and health providers in ACP conversations to promote a shared understanding of the patient’s values and preferences; (4) exploring the patient’s illness beliefs, preferences and values in a systematic way to give the patient an opportunity to reflect on these topics; (5) increasing the degree to which patients, family carers/surrogates and health providers recognise the importance of ACP; (6) addressing the patient’s need for social support in their decision making and ACP
behaviour; and (7) promoting patient and health provider skills, competence and confidence necessary to have ACP conversations.

Studies ranged from referring to a single mechanism (eg, facilitating family carer understanding of the course of dementia and different therapeutic options in order to reduce decisional uncertainty)\(^28\) to several mechanisms (eg, attending to health literacy and other individual differences, building skills to engage in ACP and encouraging the inclusion of family in the process)\(^40\).

Tailoring the delivery style or content of the intervention to the particular needs of the patient recipient, such as by matching intervention delivery to the culture of the patient population, responding to the patient’s coping style or by attending to limited health literacy,\(^30\)\(^39\)\(^40\) was the mechanism linked to the greatest variety of outcome domains. When information was presented in a personalised way, it was anticipated to be more understandable and acceptable to the patient, and maximising the prevalent coping strategies of patients allowed for a strengths-based approach in one intervention.\(^10\) This tailoring was expected to increase the number of conversations patients then conducted about ACP, their completion of ADs,\(^39\)\(^40\) and emotional and health-related outcomes.\(^10\)

Promoting the skills, competence and confidence necessary to participate in ACP conversations, both for patients and health providers, was referred to in the greatest number of studies which stated how the primary outcome was expected to be achieved. It was considered a mechanism for effecting change in the

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**Figure 2** Linkage of mechanisms to outcomes. In this figure, references where linkages between mechanisms and outcomes were made are illustrated using arrows. Where an outcome is described but not how it is expected to be achieved, the reference is placed under the outcome cell. Green outcome cells: positive findings; red outcome cells: negative findings; orange outcome cells: mixed findings. ACP, advance care planning; AD, advance directive; DNR, Do-Not-Resuscitate order.
number of conversations conducted about ACP and end-of-life care, for improving AD completion rates, and for increasing active participation and shared decision making during ACP conversations.

What are the findings for the primary outcomes?
An overview of findings for the primary outcomes can be found in online supplemental table 2. Dyad congruence was the only primary outcome for which a positive result was reported across multiple studies. One study, using a pamphlet and a discussion with a psychologist to help the patient reflect on their values towards end-of-life care, found a reduced likelihood of hospital death.

An intervention including informational leaflets and an ACP conversation during which an AD was completed did not increase patient activation.

For all other outcomes, results were mixed (see online supplemental table 2).

What impacts whether or not the intervention achieves the desired outcome?
After coding the factors through which authors explain intervention findings, as well as the success or failure of the intervention to affect the primary outcome(s), three overarching themes were identified: participant factors, implementation factors and methodological factors.

Participant factors include both individual characteristics as well as possible cultural differences. Five studies evaluated an intervention in diverse or non-white populations; these studies all made a priori considerations of the influence of racial disparities and their impact on access to quality end-of-life care. According to one study, treating the family as an asset to the ACP process through rapport-building communication can help an intervention align with African American participants’ needs. Minority populations such as Latinos may also face language barriers, which limit access to end-of-life communication. One study reported that, additionally, Latinos may not want to burden family members with distressing issues such as end-of-life conversations. Attending to barriers such as those posed by language and health literacy can increase the likelihood of intervention success.

The authors of six studies reflected on the impact of the patient’s illness experience on study outcomes. Physical pain or discomfort from treatments may reduce the likelihood that a patient engages with the intervention. In the case of chronic illness or illnesses with a less-predictable trajectory, patients may see treatment as a fundamental part of their lives or aim to live well within their condition, hindering contemplation of alternative care options. When question prompts are used in an intervention, patients who are not exclusively receiving palliative care may find it difficult to ask questions about the end of life, even after prompting.

In order to engage in ACP, patients must be willing to discuss the topic and desire a role in decision making about future care. When they do not, their engagement with ACP interventions may be limited.

The impact of other patient characteristics was less clear. The effect of age on patient use of online interventions is uncertain. The differences between heterosexual and homosexual patients in ACP interventions for patients with HIV, also merit further consideration.

Implementation refers to how the intervention was realised within the study setting. The brief duration of an intervention (eg, one conversation) was noted by two authors as a possible explanation for an absence of effects, as it may not have provided patients enough time to prepare for decision making.

An overview of findings for the primary outcomes can be found in online supplemental table 2. Two authors as a possible explanation for an absence of effects, as it may not have provided patients enough time to prepare for decision making. The authors of four studies highlighted the importance of engaging the surrogate, caregiver or family; failing to do so may contribute to null results. The authors of four studies emphasised that for ACP interventions involving health providers to achieve the intended outcomes, providers must be trained in ACP and communication skills so that they can initiate conversations and respond to patients.

The authors of two studies noted that their samples were predominantly male, but the impact of gender was not further explored. Implementation of effects, as it may not have provided patients enough time to prepare for decision making, cannot be implemented system-wide and one online intervention was not simple to use.

Methodological factors refer to how the study and its measurements were conducted. These include assessment effects, which may activate patients, contamination of health providers attending to both intervention and control arms, poor outcome fit, lack of fit between construct and measurement, or timing of measurements.

Two studies relying on documentation of ADs and ACP conversations for outcome assessment revealed clinic-level differences in the accessibility of ADs in the patient health record and potential registration bias in the medical file.

DISCUSSION
This scoping review examined 16 RCTs of complex ACP interventions for patients with chronic illness, finding results that affirm the heterogeneity of intervention components, mechanisms and outcomes identified in the existing literature.

In the retained studies, outcomes across nine categories pertained to ACP actions such as completing an AD, having consultations, patients stating their treatment preferences or SDMs judging the treatment preferences of the patient, or utterances regarding shared decision making and ACP topics during consultations.
Outcomes related to patient well-being included quality of life measures as well as psychospiritual well-being. One study reported on a care utilisation outcome, namely place of death. Additional outcomes included decisional conflict and patient activation. Interventions were expected to influence these outcomes through mechanisms which could be grouped into seven descriptive themes, with an apparent overall emphasis on a tailored approach which facilitates understanding about the patient’s illness and treatment options, encourages the patient to reflect on their personal values, and encourages the patient to talk about these topics with their carers and health providers. In order to proceed with these conversations, interventions may also encourage patients and health providers to see the importance of ACP and should equip them with the skills and confidence necessary to have these discussions effectively. Of these, the mechanism of intervention tailoring stood out as being linked to the greatest variety of outcomes in the included studies. Promoting ACP skills, confidence and competence was the mechanism most often explicitly linked to an outcome.

A finding of the present review which has not yet been fully described elsewhere is that a portion of the studies retained for full-text screening did not specify how the intervention was expected to achieve its primary outcomes: a total of 17 of 111 studies were excluded specifically for this reason.

While the application of the inclusion criteria meant that all included studies described how the intervention was expected to work or outcomes were expected to be achieved, not every study specified the mechanisms for each included component or the mechanism to which a particular outcome was expected to respond. Specifically, the anticipated impact of informational materials, the provision or completion of an AD form, and the provision of the completed AD or conversation summary to a health provider or the upload thereof in the patient’s medical record was most often not explained by the authors. The mechanisms by which decisional conflict, AD/DNR completion, dyad congruence, psychospiritual well-being, and the number of consultations with palliative care were intended to change were likewise not always specified. In the one study to include place of death as a primary outcome, the authors did not state through which mechanism this outcome was expected to change; outcome selection was based on the finding that patients prefer to avoid hospital death.

Outcomes are measured after the implementation of an intervention, in all its complexity, and components of such complex interventions may be synergistic. This makes it difficult to disentangle to which degree individual components contributed to the achieved outcomes, and we cannot make decisive statements regarding which components of ACP do or do not work. This is, however, not the intention of this scoping review. What we do find is that the contributions of intervention components such as informational materials, AD forms and sharing of conversation summaries or ADs with health providers—that is, the ‘how’ and ‘why’ of these components—are often not fully explained in the present literature. The linkage of the identified mechanisms to specific outcomes similarly lacks clarity. This may make it more difficult to ascertain what the ‘active ingredients’ of these complex interventions are and how they work, especially in the absence of a process evaluation.

Which outcomes ACP interventions can or should result in is still a topic of discussion. Even in this sample of studies that is smaller than that of a recent scoping review of ACP trials, we found a heterogeneous selection of primary outcomes. Although a Delphi panel ranked care concordance with patient wishes highest in a list of outcomes that define successful ACP, no included study featured it as a primary outcome. This outcome may require intermediate steps, such as patients being able and willing to talk about ACP, having discussions to make their wishes known to health providers and said wishes being documented. Other outcomes, such as quality of life, care and dying, may be too distant to be achieved by an ACP intervention alone, lying above the ‘ceiling of accountability’ for such interventions. Additionally, when trials examine outcomes pertaining to actions such as the occurrence of a consultation, it may be equally important to consider the quality of such consultations and the attention that is paid during these consultations to patients’ values and concerns.

Given the ongoing debate regarding ACP as a process and the outcomes which should be used to measure ‘successful’ ACP, these findings, viewed together, are indicative of a gap in how ACP interventions are currently reported and a point of consideration for future intervention research endeavours, from intervention development to implementation and assessment. Here, we echo the statement of Lin et al that identifying causality between the components of an intervention and the outcomes which are chosen to evaluate its effectiveness is a crucial element of publishing intervention research. Greater attention should be paid both to robust intervention development which articulates how the intervention will effect change and to clarity in reporting these mechanisms.

The present scoping review lends insight into the rationales presented for outcome choices in ACP intervention trials, but also identifies potential problems in the implementation and evaluation of interventions. Even if an intervention is clear in how it is proposed to work, it may face problems during implementation: a mismatch between intervention and target population, implementation setting or the methodology of the trial. An absence of significant findings may indicate the intervention does not work as hypothesised or...
may point to ‘teething problems’ in implementation. The impact of ACP on the aggressiveness of care at the end of life, for instance, may be moot in settings where overtreatment or aggressive end-of-life care is not the norm; the impact of an intervention may be limited when patients with chronic life-limiting illness are too unwell to make use of the intervention—or simply do not know how to navigate it. These are, ideally, factors which researchers should visualise before proceeding with an implementation study, for instance through consultation with key stakeholders. Of note are several interventions from the current scoping review which do precisely this, for example by taking into consideration the impact of racial disparities and then tailoring the approach of the intervention to the needs of the target population. Finally, factors which are found in one study to impact intervention results, such as lower patient motivation to engage in ACP behaviours, have the potential to be addressed through components of interventions identified in other studies, which, for example, work by building competence and confidence and by encouraging participants to recognise the relevance of ACP.

Strengths and limitations
This scoping review, to our knowledge, the first review to assess in RCTs of ACP interventions for chronic life-limiting illness the ways in which outcomes are expected to be achieved. By evaluating the linkage between intervention components, mechanisms and outcomes, we were also able to highlight gaps in research reporting, be it in the link between component and mechanism, or mechanism and primary outcome. Taking into account factors described by authors to explain success or failure to achieve the intended outcomes lays bare important considerations that should be made ahead of time when conducting ACP research, such as attention for the target population, the implementation of the intervention in a given setting and methodological factors. This scoping review can serve as a springboard for future realist reviews and evaluations of ACP interventions where cross-sectional, qualitative and grey literature are also included. Data charting for this study was done largely by one author (JS) after cross-checking a sample of extractions with a second author (ADV) and discussing the information to be extracted with all authors. The synthesis of results was performed by one author (JS). Findings were regularly discussed with other members of the research team. The framework for data charting and overall synthesis was discussed during multiple meetings with all authors, and the first and last authors conducted weekly meetings during which findings could be discussed. While we chose to include only RCTs, we did not assess the risk of bias for these studies, consistent with scoping review methodology. Further, we only included trials which involved adult patients with chronic life-limiting illness. Interventions for community-dwelling adults or populations of paediatric patients with chronic life-limiting illness may yield different findings.

CONCLUSION
This scoping review identified components of complex ACP interventions for chronic serious illness, as well as seven mechanisms by which nine outcome categories were proposed to be affected by these interventions. In reporting primary effectiveness in RCT studies, the mechanisms by which the intervention is anticipated to impact the chosen outcomes are, however, frequently unstated or unclearly stated; reporting emphasises ‘what’ is being done but less frequently refers to ‘why’ this is being done. Mechanisms such as promoting ACP skills, confidence and motivation, as well as allowing for intervention tailoring to the patient’s needs, come to the fore most clearly from this review. ACP interventions should take into account patients’ illness experience, willingness to engage in ACP and broader cultural considerations; these should ideally be considered during development to allow for an intervention that is sensitive and responsive to these factors. So too should researchers consider the setting in which the intervention is implemented and how the study and its measurements are conducted. In the light of recent debates regarding what outcomes ACP can be expected to achieve, this review indicates, in sum, ACP interventions do have potential to improve outcomes, but researchers should be clear in how they anticipate these outcomes will be achieved and consider the context in which these interventions are implemented.

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REFERENCES
### Review

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