Study protocol on advance care planning in multiple sclerosis (ConCure-SM): intervention construction and multicentre feasibility trial

Ludovica De Panfilis,1 Simone Veronese,2 Michela Bruzzone,3 Marta Cascioli,4 Alberto Gajofatto,5,6 Maria Grazia Grasso,7 Paola Kruger,8 Alessandra Lugaresi,9,10 Leigh Manson,11 Sara Montepietra,12 Francesco Patti,13 Eugenio Pucci,14 Claudio Solaro,15 Andrea Giordano,16 Alessandra Solari16

ABSTRACT

Introduction. Multiple sclerosis (MS) is the most common cause of progressive neurological disability in young adults. The use of advance care planning (ACP) for people with progressive MS (pwPMS) remains limited. The ConCure-SM project aims to assess the effectiveness of a structured ACP intervention for pwPMS. The intervention consists of a training programme on ACP for healthcare professionals caring for pwPMS, and a booklet to be used during the ACP conversation. Herein, we describe the first two project phases.

Methods. In phase 1 we translated and adapted, to the Italian legislation and MS context, the ACP booklet of the National ACP Programme for New Zealand. Acceptability, comprehensibility and usefulness of the booklet were assessed via 13 personal cognitive interviews with pwPMS and significant others (SOs), and one health professional focus group. Based on these findings, we will revise the booklet. In phase 2 we will conduct a single-arm pilot-feasibility trial with nested qualitative study. Participants will be 40 pwPMS, their SOs, health professionals from six MS and rehabilitation centres in Italy. In the 6 months following the ACP conversation, we will assess completion of an advance care plan document (primary outcome), as well as safety of the intervention. Secondary outcomes will be a range of measures to capture the full process of ACP; patient-carer congruence in treatment preferences; quality of patient-clinician communication and caregiver burden. A qualitative process evaluation will help understand the factors likely to influence future implementation and scalability of the intervention.

Ethics and dissemination. The project is co-led by a neurologist and a bioethicist. Phase 1 has received ethical approvals from each participating centre, while phase 2 will be submitted to the centres in May 2021. Findings from both phases will be disseminated widely through peer-reviewed publications, conferences and workshops. Trial registration number: ISRCTN48527663; Pre-results.

INTRODUCTION

With a lifetime risk of 1 in 400, multiple sclerosis (MS) is the most common cause of progressive neurological disability in young adults. Approximately 2.3 million people worldwide have MS, with Canada, the USA and some European countries, including Italy, having the highest prevalence rates.1 Around 15% of people with MS have a primary progressive course at diagnosis, and a further 35% develop secondary progressive disease after 15 years.2 A mean reduction in life expectancy by 7–14 years has been reported in people with MS, with improved figures over the last two decades.3–5 Few treatment options are currently available to delay or prevent further clinical worsening of people with primary or secondary progressive MS (pwPMS). They may live for many years experiencing a wide range of symptoms, impairments (including cognitive impairment which affects 40%–70% of sufferers6) and comorbidities.5 7–10

Advance care planning (ACP) is a process that ‘enables individuals who have decisional capacity to identify their values, to reflect upon the meanings and consequences of serious illness scenarios, to define goals and preferences for future medical treatment and care, and to discuss these with family and healthcare professionals (HPs)’.11
Consistently with the Shared Decision Making model, ACP involves both the patient and his/her HPs. Together, they make informed decisions about the patient’s (future) care. Also, the family can be involved in the process, if the patient wishes. ACP differs from general medical decision-making in that it is based on an anticipated deterioration in the health of a patient. It includes a focus on the person’s wishes and preferences for the time when they lose decisional capacity. In fact, it aims to align evidence-based practice and person-centred care using a bioethical focus to identify the patient’s values, preferences and desires. The planning process helps the patient to identify his/her personal values and goals, understand his/her health status, and the treatment and healthcare options available. Finally, ACP encourages discussion around end-of-life (EOL) care (a subject that is generally not considered part of healthcare planning, and one that can be avoided by both patients and HPs). It is up to the patient to determine the occurrence and content of any ACP discussion: if the patient does not wish to engage in conversations about his/her future care, this preference should be respected. The ACP process may result in the patient choosing to write an advance care plan document and to appoint a trustee (or else).

On 22 December 2017, the Italian Parliament approved the first law on EOL: ‘Provisions for informed consent and advance directives’ (L. 219/2017; http://www.trovanorme.salute.gov.it/norme/dettaglioAtto?id=626663). This law regulates advance directives (AD; Article 4) and ACP (Article 5), and a number of rights citizens have regarding healthcare issues, including the right to: be fully informed about one’s health status and to give consent (or dissent) to treatment; withhold consent to lifesaving treatments; be assisted until death. Moreover, the law states that the physician has a duty to respect the patient’s wishes. In a recent Italian survey, 88% (1752/2000) of citizens considered the Law 219/2017 as quite or very important, and 76% had a positive attitude towards making/registering AD or ACP. Importantly, this Law triggers HPs and healthcare authorities in promoting educational programmes on the topic, as well as programmes to implement ACP in daily clinical practice.

To optimise the alignment between patient preferences and values and the care they receive, HPs should integrate best ACP practices in the care of pwPMS. A recent guideline on palliative care in MS found no evidence of the effects of ACP in pwPMS. However, there is some evidence from non-neurological progressive and life-threatening illnesses that ACP decreases the use of life-sustaining treatment, increases hospice/palliative care, reduces hospitalisations and increases alignment with patients’ EOL wishes. Furthermore, there is evidence that patients with MS and caregivers often would like to discuss the issues of death and dying and HPs should acknowledge and encourage these discussions. However, often HPs leave EOL discussions until the later stages of progression in MS, and caregivers may be left having to take difficult decisions. A realist review identified two main barriers for ACP discussions taking place: the long and uncertain MS trajectory, with periods of stability punctuated by crisis; and lack of ACP communication skills and confidence of HPs.

ConCure-SM is a project aimed to set up and evaluate the efficacy of an ACP intervention for pwPMS in Italy. The Shared Decision Making model described earlier is the theoretical framework of the project. The Medical Research Council framework for developing and evaluating complex interventions is the methodological framework of the project. The framework has a phased approach, from a preclinical research phase to a final phase in which the intervention is introduced into the health service, leading to a theory-driven intervention: a ‘bottom-up’ development which guarantees to enter a phase III trial with an appropriate theory and pilot work. Furthermore, both quantitative and qualitative methods are integrated within the framework, in order to better appraise the effects of the (complex) intervention both as a whole and on its components.

Our study hypotheses are that the intervention will produce: higher completion of an advance care plan document; increased congruence in treatment preferences between pwPMS and their carers; increased quality communication about EOL care.

METHODS AND ANALYSIS
The study protocol (FISM 2020/R-Multi/024; V.1.0; 15 March 2021) was designed following the Standard Protocol Items: Recommendations for Interventional Trials (SPIRIT) guidelines (online supplemental appendix 1) and the SPIRIT-PRO Extension. The pilot/feasibility study follows the Consolidated Standards of Reporting Trials (CONSORT) guidance for trials on social and psychological interventions (CONSORT-SP1 2018). It was registered on the ISRCTN registry on 30 March 2021. Qualitative data will be reported following the Consolidated criteria for Reporting Qualitative research checklist.

Figure 1 outlines the two project phases and inscribed actions. The red dot identifies the current advancement status.

Phase 1
The first project phase involves production of the ACP booklet (figure 1).

Provisional booklet
Early in 2020, an interdisciplinary panel translated into Italian and adapted to the MS context and to the Italian legislation, the ACP booklet of the Health Quality & Safety Commission’s New Zealand National ACP Programme (https://www.myACP.org.nz). The panel was made of five neurologists, one palliative care physician, one palliative care nurse, one psychologist, one bioethicist, one expert patient, one representative of the Italian MS Society and...
the author of the original booklet. The resulting booklet in its provisional version (online supplemental appendix 2) consists of an introduction, a ‘guidance’ (the odd pages in most instances) and the advance care plan document (the even pages) to be completed electronically or manually by the pwPMS together with his/her referring physician. A significant other (SO), such as a family member, can participate in the process if requested by the pwPMS. The introduction explains the concepts of ACP and AD according to the Italian Law 219/2017, and describes why ACP is important in MS. Ten sections follow: ‘My Advance Care Plan’; ‘What matters to me’; ‘What worries me’; ‘Why I’m making an Advance Care Plan’; ‘How I make decisions’; ‘If I were no longer able to make decisions: my trustee’; ‘Thinking about my EOL’; ‘My treatment and care choices’; ‘Signatures’; ‘Acronyms’. If the advance care plan document is completed, the pwPMS (and, when applicable, the pwPMS trustee) sign on page 29; the document is then scanned and stored, together with the completed booklet, in the (electronic) medical record.

Users’ assessment and revision
Between September and November 2020, the acceptability (contents, format, envisaged administration procedure), comprehensibility and usefulness of the provisional booklet were assessed by conducting 13 personal cognitive interviews with pwPMS, pwPMS’ SOs and a focus group meeting (FGM) with HPs. Due to the COVID-19 pandemic, all the interviews and the FGM were held using digital platforms. Results of the qualitative (thematic) analysis and the revision of the booklet are underway.

Phase 2
The second project phase will be dedicated to the conduction of the multi-centre, pilot and feasibility single-arm trial with a nested qualitative study. This phase (to be accomplished from May 2021 to November 2022) has three inscribed actions: intervention set up; pilot trial and qualitative study (figure 1).

Intervention set up
Training programme
The goal of this intervention is to prime HPs to discuss goals of care and ACP. To achieve this, HPs will attend a training programme (called Train-ConCure-SM) that will be Continuing Medical Education accredited, residential, and last one-and-half days (12 hours). The programme aims to: improve the HP knowledge, competencies and skills in ACP based on up-to-date scientific evidence; support and guide HPs in the ACP embedding in clinical practice; improve the communication between HPs and patient promoting an effective patient–practitioner partnership in decision-making.

The training will be interactive in style. Its residential nature and the use of role-playing exercises aim at supporting group discussion and the exchange of experiences between participants.

It will consist of the following: one 2.5-hour theoretical session on the clinical, ethical and statutory principles of Shared Decision Making and ACP; two 4-hour empirical sessions (one on each day) on conducting ACP conversations in various clinical scenarios using the ConCure-SM booklet through guided role play exercises; two 45-min self-evaluation sessions (at the beginning and at the end of the training programme).

Trainees will be physicians and other HPs from the six enrolling centres. The Italian Law 219/2017 prescribes that ACP involves the patient, his/her referring physician and (when applicable) the trustee. We decided to train HPs other than physicians in order to promote ACP knowledge within the caring team. Each centre will provide 1–3 physicians, plus one HP from the following: MS nurse, therapist, psychologist or social worker. Thus, there will be 12–24 participants overall (2–4 from each centre). Trainers will be a panel of neurologists, psychologists, a palliative care physician, a palliative care nurse and a bioethicist. All have consolidated experience in orchestrating training courses and workshops on patient–clinician communication and Shared Decision Making, and four on ACP and EOL conversations. These four researchers will support physicians at the centres for issues concerning the conduction of the ACP conversation during the pilot trial.

Web platform
As part of the intervention set up action, a web-based trial platform will be created containing the pseudo-anonymised trial case record form (eCRF) and the outcome measures. The platform will be ID/password protected, with dedicated accesses based on the stakeholder (pwPMS, SO, HPs, centre principal investigator (PI), interviewer, data manager) and operation (completion, consultation).
Linguistic validation of measures

Two outcome measures not available in Italian will be translated and cross-culturally adapted, following accepted guidelines\(^3^9\)\(^3^0\): the 4-item ACP Engagement questionnaire (4-item ACP-E)\(^3^1\), and the Quality of Communication Questionnaire (QOC).\(^3^2\) The main steps in this process are the following:

- **Forward translation**: two qualified translators, both living in Italy, will produce two independent translations. A panel consisting of the translators, two MS HPs and two lay persons will review the forward translations and a consensus version will be produced.

- **Backward translation**: the consensus translation generated in step 1 will be independently translated back into the source language by a third qualified translator, living in the target country. The backward translation will be produced without access to the original version and without consulting the other translators.

- **Translation refinement**: in a meeting between those participating in step 1 and the backward translator, the backward translation will be compared with the original, and further refinements to the Italian version will be made. Differences will be resolved by discussion.

- Each translated questionnaire will be proofread, and then administered to/debriefed with 5–10 patients.

Pilot and feasibility trial

The six centres involved in the pilot trial are located in northern (four centres), central and southern Italy (one centre each). Two of the centres are rehabilitation hospitals (one of which a research hospital), three are MS centres (two university hospitals, one research hospital) and one is a rehabilitation and MS centre from a research hospital. Recruitment will be competitive, with no prespecified minimum number of enrolled subjects per centre. The maximum number of enrolled subjects per centre is 12.

There will be a baseline assessment (T0), an ACP conversation taking place within 1 month from the baseline assessment, and a follow-up assessment within 1 week of the ACP conversation (T1) and 6 months (T2) thereafter. The baseline and follow-up assessments will be performed via the web-based ConCure-SM platform. The physician will record on the platform subsequent ACP conversations that should occur during follow-up. Participants (pwPMS, SOs) will be free to withdraw from the study at any time, without giving reasons and with no risk of prejudicing future care. Study personnel will make every effort to obtain, and record, information about the drop out reasons.

The objectives of the pilot and feasibility trial are reported in box 1. trial procedures are summarised in figure 2.

Eligibility and screening

PwPMS (inpatients or outpatients) will be included if they are: ≥18 years of age; diagnosed with primary or secondary PMS\(^3^3\) one or more years before inclusion; able to communicate in Italian and gave written consent. In addition, one or more of the following conditions that would make ACP relevant must be present: expressed desire for ACP; questions about own future; thoughts about hastening death or medically assisted suicide; high risk for death within 2 years using the ‘Surprise Question’\(^3^4\); high risk for development of severe cognitive compromise/dementia within 2 years; high risk for development of impairments preventing communication within 2 years; significant suffering (eg, uncontrolled physical symptoms, psychosocial or existential issues). PwPMS will be excluded if they have one or more of the following: severe cognitive compromise (MMSE <19) or impairments preventing communication; psychosis or other serious psychiatric conditions; advance care plan document completed.

PwPMS are recruited prospectively by the ACP-trained physicians involved in their care, when the potentially eligible pwPMS attends the centre for an outpatient visit or hospitalisation. PwPMS who show interest in participating receive full verbal and written information about the study purpose and procedures.

Baseline assessment (T0)

The ACP-trained physician makes an appointment with pwPMS who provided initial verbal consent to participate in the study, and checks all eligibility criteria. A written, signed informed consent is obtained, according to the Declaration of Helsinki and to the Good Clinical Practice (GCP) guidelines of the EU. The informed consent is kept on file by the study personnel, and is available for inspection by regulatory authorities or authorised persons.

Then, the physician gives the pwPMS the credentials to the trial platform, so that the pwPMS completes the baseline set of questionnaires/instruments (completion time around 40 min). If the pwPMS has difficulties in using the trial platform, a telephone interview is scheduled within...
Eligible pwPMS:  
- Age ≥ 18 years  
- At least one out of seven conditions that would make ACP relevant  
- Able to communicate in Italian  
- Adequate cognitive and communicative ability to participate  
- No serious psychiatric conditions  
- No previous advance care plan document completed

Participant screening:  
- Confirm eligibility  
- Obtain name/contact of significant other (if applicable) and permission to contact

Baseline assessment (T0):  
- PwPMS:  
  - HADS  
  - eCPS  
  - 4-item ACP Engagement  
  - MSQOL-29

- Significant other:  
  - General data  
  - ZBI

- Physician:  
  - PwPMS general and clinical data (EDSS [Kurtzke 1983], Barthel Index [Mahoney 1965])  
  - Physician’s general data

First ACP conversation:  
- OPTION scale (physician’s competences)

Follow-up assessment (T1):  
- PwPMS:  
  - HADS  
  - QOC  
  - 4-item ACP

- Significant other:  
  - ZBI  
  - QOC-SO

- Physician:  
  - QOC-Doc

Follow-up assessment (T2):  
- PwPMS:  
  - HADS  
  - 4-item ACP Engagement  
  - MSQOL-29

- Significant other:  
  - ZBI

- Physician:  
  - PwPMS clinical/ACP update

Figure 2  Summary of trial procedures. ACP, advance care planning; ACP-E, ACP Engagement; eCPS, Control Preference Scale, electronic; EDSS, Expanded Disability Status Scale; HADS, Hospital Anxiety and Depression Scale; MSQOL-29, 29-item Multiple Sclerosis Quality of Life; OPTION, Observing Patient Involvement in Shared Decision Making; QOC, Quality of Communication; SO, significant other; ZBI, Zarit Burden Interview. Kurtzke, Mahoney and Barthel.

a week with an independent, trained interviewer who will administer the questionnaires/instruments.

The ACP conversation is scheduled at the centre, within a month. It is the starting point of a process that is followed up during the study. However, for feasibility reasons and to adapt to participant needs, subsequent conversations are recorded, but not scheduled a priori. The pwPMS is invited to involve his/her SO (family member, relative or friend, who is next of kin or is key decision-maker as designated by the pwPMS and with whom the pwPMS shares his/her life). If the pwPMS agrees on involving his/her SO, the SO is contacted by a study researcher to confirm eligibility, explain the study and obtain verbal consent. Consenting SOs receive credentials to access the trial platform and complete the baseline set of questionnaires (completion time about 15 min). If the SO has difficulty
in using the trial platform, a telephone interview is scheduled within a week with an independent, trained interviewer who will administer the questionnaires.

Finally, the physician completes the eCRF via the trial platform.

Each centre will collect information on the number of pwPMS and SOs approached, screened and eligible prior to enrolment, with reasons for non-enrolment.

**ACP conversation**

The conversation involves the pwPMS, the ACP-trained physician involved in his/her care and, when applicable, the SO. In addition, if the pwPMS agrees, the non-physician ACP-trained HP at the centre will participate. The first conversation takes place in a dedicated room at the centre, and is audio-recorded. At MS centres and rehabilitation centres, physician time and space are at premium, particularly for outpatient care. For this reason, 1-hour slot is reserved for the conversation. In the case an SO participates it is envisaged that there will be a session closed to the SO, followed by an open session.

About 1 week before the scheduled ACP conversation, reminder emails (or telephone calls) are sent to pwPMS/ SOs. At the end of the ACP conversation, the physician invites the pwPMS (when applicable the SO) to complete the T1 follow-up assessment within 1 week. The physician completes the QOC-Doc immediately after the ACP conversation ends.

**Follow-up assessments (T1, T2)**

The pwPMS completes the questionnaires by 1 week (T1, assessment time of about 20 min) and 6 months (T2, assessment time of about 30 min) after the first ACP conversation using the trial platform. The SO completes the questionnaires/instruments (T1, assessment time of about 20 min) using the trial platform. In the event the pwPMS/SO have difficulties in using the trial platform, a telephone interview is scheduled with an independent, trained interviewer who will administer the questionnaires/instruments.

About 1 week before the T2 assessment, reminder emails (or telephone calls) are sent to pwPMS. The physician completes the questionnaire (T1, QOC-Doc) and the eCRF using the trial platform. He/she records on the platform the date, duration, participants and mode (face to face, teleconference or on the telephone) of subsequent ACP conversations that occur during follow-up.

**Outcome measures**

A range of measures will be collected to capture the full process of ACP and whether the ConCure-SM intervention has any effect on completion of an advance care plan document (primary outcome measure), congruence in treatment preferences between pwPMS and their carers, quality of patient–clinician communication and caregiver burden (table 1). In addition, since a study-related increase in emotional burden can not be excluded, serious adverse events (admission to psychiatric ward, suicide attempt, death) will be monitored by the independent Data and Safety Monitoring Committee (DSMC).

We will use the published Italian version of the following inventories: Control Preference Scale (CPS)35; Hospital Anxiety and Depression Scale (HADS)36; Observing Patient Involvement in Decision Making (OPTION)37; 29-item Multiple Sclerosis Quality of Life (MSQOL)38; Zarit Burden Interview (ZBI).39 The 4-item ACP-E and the QOC inventories will be translated/culturally adapted from the source language (see previously).

**ACP engagement**

The ACP process will be assessed using the 4-item ACP-E questionnaire.31 Originally developed and validated to measure the complex behaviour of ACP, the questionnaire is available in four versions (55-item, 34-item, 9-item, 4-item). In this study, we will use the 4-item version which focuses on the readiness behaviour change construct.

### Table 1 Secondary outcome measures of the trial (in alphabetical order)

| Scale name | Assessor | Construct | Author | Italian version | Timing |
|------------|----------|-----------|--------|-----------------|--------|
| 4-item ACP-E | Patient | ACP engagement | Sudore et al31 | – | T0/T1/T2 |
| eCPS | Patient | Role preferences | Degner et al40 | Solari et al42 | T0 |
| HADS | Patient | Mood symptoms | Zigmond and Snith44 | Costantini et al36 | T0/T1/T2 |
| MSQOL-29 | Patient | Health-related QOL | Rosato et al38 | Rosato et al38 | T0/T2 |
| OPTION | Third observer | Shared Decision Making (physician’s skills) | Elwyn et al43 | Goss et al37 | – |
| QOC | Patient | Communication quality (physician’s skills) | Engelberg et al42 | – | T1 |
| QOC-Doc | Physician | Communication quality (physician’s skills) | – | – | T1 |
| QOC-SO | SO | Communication quality (physician’s skills) | – | – | T1 |
| ZBI | SO | Caregiver burden | Hérbert et al46 | Chattat et al39 | T0/T1/T2 |

ACP-E, Advance Care Planning Engagement; eCPS, Control Preference Scale, electronic; HADS, Hospital Anxiety and Depression Scale; MSQOL-29, 29-item Multiple Sclerosis Quality of Life; OPTION, Observing Patient Involvement in Shared Decision Making; QOC, Quality of Communication; SO, significant other; ZBI, Zarit Burden Interview.
within the quality of life ACP domain. Responses are on a 5-point Likert scale (1) ‘I have never thought about it’; (2) ‘I have thought about it, but I am not ready to do it’; (3) ‘I am thinking about doing it in the next 6 months’; (4) ‘I am definitely planning to do it in the next 30 days’; (5) ‘I have already done it’.

Role preferences

The CPS is the most used instrument to assess patient preferences for involvement in decisions about their health. It consists of five ‘cards’ on a board, each illustrating a different role in decision-making by means of a cartoon and short descriptive statement. In its original version, administration requires a trained examiner, who asks the patient to choose the preferred card, which is then covered up. The procedure continues (four choices) until one card is left. If the second preference is incongruent with the first (non-adjacent pairing, such as card A with card C), the test is explained again, and immediately readministered. In the event of a further incongruence, the test is not readministered, and a preference is not assigned. Six scores are possible based on the subject’s two most preferred roles: active–active, active–collaborative, collaborative–active, collaborative–passive, passive–collaborative and passive–passive. These scores are grouped as: active (active–active or active–collaborative), collaborative (collaborative–active or collaborative–passive) or passive (passive–collaborative or passive–passive). We will use the electronic, Italian self-administered CPS.

Quality of the conversation

We will assess the quality of the first ACP conversation considering three perspectives: an independent observer, the pwPMS, and the physician. Each conversation will be unobtrusively audio-taped and transcribed verbatim; subsequently, a specially trained third observer will evaluate the behaviour of the physician in terms of patient involvement in decision-making using the OPTION (http://www.glynelwyn.com/observer-option-instrument.html). The OPTION consists of 12 items, each rated on a 5-point Likert scale ranging from 0 (behaviour not observed) to 4 (behaviour observed to high standard). A total score (range 0–48) is obtained by adding the scores of each item. After the ACP conversation, pwPMS will complete the QOC; SOs will complete the SO version (QOC-SO), and physicians the physician version (last two items) of the QOC. Developed from qualitative studies with patients, families and clinicians, the QOC consists of 19 items measuring general communication (9 items) and communication about EOL care (8 items), each rated on a scale from 0 (‘very worst I can imagine’/‘not at all’) to 10 (‘very best I can imagine’/‘extremely’), or identified as something the clinician did not do. The 0/10 ratings are recoded to 1/11, with 0 imputed for ‘did not do’ (http://depts.washington.edu/eolcare/products/instruments/).

Other outcome measures

PwPMS quality of life will be assessed using the electronic version of the MSQOL-29, which is the shortened form of the MSQOL-54. MSQOL-29 includes 25 items forming 7 subscales and 4 single items, and 1 filter question for 3 ‘sexual function’ items. Mood symptoms will be assessed with the HADS, a self-assessed questionnaire consisting of 14 multiple-choice (0–3 Likert scale) items probing symptoms of anxiety (7 items) and depression (7 items). HADS anxiety (HADS-A) and depression (HADS-D) scores range from 0 (no symptoms) to 21 (most severe symptoms). A cut-off score of 8 or above was recommended for patients with MS, since it was found to be an accurate indicator of major depression (90% sensitivity, 87% specificity) and generalised anxiety disorder (88.5% sensitivity; 81% specificity) in this population.

Finally, SO burden will be assessed using the ZBI, a 22-item self-report measure of subjective burden among caregivers addressing functional and behavioural impairments as well as the home care circumstances. A total 0 (low burden) to 88 (high burden) score is obtained by summing item responses, each scored on a 5-point Likert scale ranging from 0 (never) to 4 (nearly always present).

Meetings

There will be two study meetings (teleconferences): the investigators’ meeting will be held before enrolment starts. Participants will be the Steering Committee, the centre PIs and the HPs who participated in the training programme. The aim of this meeting is to provide clear information on the study procedures, and to train HPs on the use of the trial platform. A second meeting will be run about 2 months after enrolment starts, in order to monitor possible difficulties, top-up centres’ motivation and provide a safe place for peer discussion on the implementation of the intervention. Both meetings will last about 2 hours. Additional meetings will be organised whenever needed. In addition, the study PIs and the Training Panel will be available for inquiries about the implementation of the intervention at the participating centres.

Nested qualitative study

We will perform one-on-one semi-structured interviews with pwPMS and SOs, chosen using a maximum variation strategy, and FGMs of HPs involved in intervention delivery. For pwPMS and SOs interviews were considered most appropriate to limit interview burden and hopefully make it easier for participants to express their feelings, and recount their experiences of the intervention. For the patient referring physicians and the other HPs we chose FGMs as they promote interaction and exchange of ideas. A minimum of 10 interviews (five with pwPMS and five with SOs) and two FGMs will be held, the final number depending on the achievement of ‘data saturation’. Interviews and FGMs will be run via video teleconference, which will ease participation of pwPMS with severe disability and SOs with caregiving commitments, as...
well as HPs. If the pwPMS and/or SOs have no access to internet using personal computer or other devices, such participants will be interviewed on the telephone.

The interviews aim to provide important feedback on participant perception of the quality of the intervention provided, and will serve as a process measure. Insights from this qualitative analysis will serve to inform fine-grain intervention refinement. They will take place within 2 months of trial completion, and last no more than an hour. To reduce social desirability response bias, the interviewers will be researchers not involved in the ConCure-SM intervention delivery. Before starting, interviewees will be informed of study aims and requirements, and provide written consent. The interviewer will then explain that the aim of the interview is to obtain participant feedback on experience of the pilot study and stress that positive and negative experiences of, and feelings about, the intervention are welcome. Participants will be assured that the interviews are confidential, and that the audio recordings and subsequent transcripts will be fully anonymised. The interviewer will then pose each question in turn, neutrally (so as to not suggest any particular reply) and in an open-ended fashion (to allow many possible replies). As each question is discussed, follow-up questions will clarify and explore participant responses. Participants will be also encouraged to elaborate on any pertinent themes or views that emerge. The interviewer will note any potentially informative non-verbal gestures. At the end of the interview, the interviewer will verbally summarise the key points and ask the participant if the summary is full and correct.

The FGMs aim to collect insights and living experiences about the intervention and to identify possible barriers to its implementation; they will provide important feedback on the intervention and on factors that can enable its implementation and adoption. For this reason, HPs other than the physicians involved in the ACP conversation will be involved. Each FGM (teleconference) will last about 2 hours; participants will be 6–10 physicians who delivered the intervention and HPs from the participating centres. All participants will provide written informed consent prior to the FGM, that will be conducted by two psychologists specifically trained in qualitative research. One will be the facilitator, whose job is to engage all participants, promote exchange, moderate conflicts, ensure that all prespecified topics will be adequately covered, and allow exploration of any pertinent issues that arise. He/she will first explain the purpose of the meeting and ask participants to introduce themselves. He/she will then introduce each topic in turn, in an open-ended fashion. At any point, the facilitator can probe for further information and ask follow-up questions to stimulate further discussion. After all prespecified topics are fully discussed, the facilitator will summarise the main points, and ask for further feedback and whether all concerns have been fully aired. The co-moderator will take notes and oversee the audio recording. Subsequently, they will produce a report from the audio recordings/transcript and field notes, which will be submitted to participants for review (respondent validation).

Data analysis

Study power

As this is a pilot and feasibility study, a formal sample size calculation is not required. We aim to recruit at least 40 pwPMS from six centres to assess feasibility across a diverse range of participants including those with different care needs and living conditions. There are no data available on the occurrence of ACP in pwPMS: by hypothesising a proportion in the pwPMS population of 10%, a sample size of 35 subjects achieves a power of 90%, assuming a type I error of 5%, to detect a proportion of ACP documentation of 30%. By hypothesising a proportion in the pwPMS population of 8%, a sample size of 35 subjects achieves a power of 95%, assuming a type I error of 5%, to detect a proportion of ACP documentation of 30%. By adding 15% of drop outs or incomplete data, 40 pwPMS should be recruited.

Statistics

Descriptive statistics will be calculated for general and clinical variables. Specifically, continuous variables will be summarised by their mean and SD, or median and IQR; categorical variables will be summarised as numbers and percentages. Categorical variables will be compared using the χ² test. The normality assumption of continuous variables will be tested with the Shapiro-Wilk test. Depending on data distribution, between-group comparisons will be carried out using either the two-sided unpaired t-test or the Wilcoxon two-sided two sample test; within-group comparisons will be carried out using either the paired t-test or the Wilcoxon signed-rank test; correlations will be computed using Pearson’s or Spearman’s coefficients.

Our primary end-point is the proportion of pwPMS completing an ACP during the 6-month period. Change in the secondary outcome measures will be also calculated. In addition, we will calculate the following feasibility outcomes: recruitment rate (enrolment per month; reasons for non-eligibility, non-enrolment); retention rate (proportion completing the intervention and study follow-up); missing data (proportion fully completed, for each scale, at each time point). Data will be analysed according to the intention-to-treat principle. Multiple imputation of missing values will employ Rubin’s approach. A p value less than 0.05 will be considered statistically significant. No correction for multiple comparisons will be applied. All analyses will be performed using STATA V.16. Assumptions in determining the sample size of the main trial will be checked.

Qualitative data

Interviews and FGMs will be audio-recorded and transcribed verbatim. Data analysis will be conducted by three researchers with experience in qualitative research. Researchers will analyse interviews and FGM data using thematic analysis, with interpretation guided by the four
normalisation process theory (NPT) components (see process evaluation below). Data will be triangulated across sources. The analytical stages can be summarised as follows: (1) each researcher will read the transcriptions and write comments and initial thoughts in a memo. (2) Each researcher will extract portions of the text individually and then share their work to reach an initial agreement. During this stage, they will conduct the thematic analysis inductively providing their insights. (3) Researchers will independently review themes and allocate portions of the text to the newly reconfigured themes. (4) Together, they will redefine themes and rename them to achieve internal consistency. (5) One researcher will extract from the interviews and draft the final report, which will be checked and amended by the other two.

Process evaluation
We will follow the Medical Research Council guidance on process evaluation, which describes three components using a mixed-methods approach: implementation or delivery; mechanisms of impact; contextual factors. We will use NPT to determine if, and in what ways, the ConCure-SM intervention can be successfully 'normalised' (embedded) into clinical practice. At the feasibility and piloting stage, basic quantitative measures of implementation may be combined with in-depth qualitative data to provide detailed understandings of intervention functioning on a small scale. Quantitative measures will include structured observations of audio-recorded ACP conversations. These will be used to examine aspects of fidelity (such as consistency with the Shared Decision Making principles), and dose (the duration of conversations). Qualitative methods will be used to investigate mechanisms of impact and contextual factors, using NPT. NPT identifies four essential determinants of ‘normalising’ complex interventions into common practice: coherence (the extent to which an intervention is understood as being meaningful, achievable and valuable); cognitive participation (the engagement of HPs necessary to deliver the intervention); collective action (the work that brings the intervention into use) and reflexive monitoring (the on-going process of adjusting the intervention to keep it in place). These components are considered to be dynamic and interact within the wider context of the intervention, such as existing organisational structures and procedures. Further, we will use qualitative data to identify required modifications and to develop practical strategies for enabling and sustaining intervention delivery in clinical settings.

Patient and public involvement statement
An expert patient with MS and a representative of the Italian MS Society are part of the Steering Committee of the project and coauthors of the present paper. These same persons were part of the interdisciplinary panel that produced the ACP booklet, which was revised based on the results of a qualitative study with users (pwPMS, SOs and HPs).

Prior to designing and conducting a full trial, the intervention will be pilot tested in a multicentre study involving MS and rehabilitation centres across Italy, and using a mixed-method approach.

We will disseminate key study findings to pwPMS via the Italian MS Society.

Ethics and dissemination
The project is co-led by a neurologist and a bioethicist. Phase 1 has received ethical approvals from each participating centre, while phase 2 will be submitted to the centres in May 2021. Findings from both phases will be disseminated widely through peer-reviewed publications, conferences and workshops. Authorship eligibility will be based on the International Committee of Medical Journal Editors. The final trial (pseudo-anonymised) dataset will be accessed by the study PIs and the data management/analysis team. Details about panels and centres, ethics and administrative considerations, and study management and monitoring are available in online supplemental appendix 3.

DISCUSSION
One of the 10 clinical questions of the European Academy of Neurology (EAN) guideline on palliative care of pwPMS specifically addressed ACP. For this clinical question, formulated with direct patient and caregiver involvement, no evidence was found and two good practice statements were produced: ‘It is suggested that early discussion of the future with ACP is offered to patients with severe MS’; ‘It is suggested that regular communication about the future progression of MS is undertaken with patients and families/caregivers’. To fill this knowledge gap, we conceived the present study, which adheres to the Shared Decision Making model, and to the Medical Research Council framework for developing and evaluating complex interventions. Within this methodological context, the study follows the CONSORT guidance for trials on social and psychological interventions (CONSORT-SPI 2018), as many of the guidance items (excluding items that are specific to the randomisation nature of the study) are relevant for reporting other types of pilot and feasibility studies. This includes the development of the study protocol following the SPIRIT guidance, protocol’s publication and the trial public registration (ISRCTN registry). The consolidated criteria for reporting qualitative research will guide the presentation of findings in the study reports and publications.

To increase generalisability of the study, participants (pwPMS, SOs and HPs) will be enrolled from university hospitals, research hospitals and clinical centres from the different areas of Italy. We will not enrol from primary care practices as in Italy patients with MS are followed in tertiary care centres. Moreover, previous attempts to
involve the family physicians in the care of PwPMS were challenging.55

Personal, semi-structured interviews and FGMs will be run via video teleconference, which will ease participation of pwPMS with severe disability and SOs with caregiving commitments, as well as HPs. If pwPMS and/or SOs have no internet access, using personal computer or other devices, these participants will be interviewed on the telephone. Other measures adopted to minimise bias include: all study personnel will be trained to conform to GCP regulation; electronic version of the study questionnaires/inventories will be used to ensure the data entered is of high quality; an DSMC will monitor and supervise the progress of the trial, and the safety data.

The ConCure-SM intervention (booklet and HP training programme) can be adapted for use in other neurological and non-neurological conditions for which consolidated ACP interventions are not available. The electronic format will ease the incorporation of the advance care plan document (and its updates) in the electronic medical record, that is currently available in some Italian regions and hopefully will be soon available all over Italy.

Study limitations
Three study limitations are noted. We used a single-arm design for the pilot trial. This decision was taken as ACP is currently at premium in MS17 23 and designing a randomised (cluster) trial with standard care or any ‘low intensity’ intervention as a comparator was considered ethically and practically unviable. Another limitation is that our training programme was for HPs only. A multiple-component intervention that targets clinicians and patients simultaneously has been suggested in other disciplines.56 In the current situation regarding ACP, we preferred to have a clear focus on enhancing HP competencies.17 23 Finally, our pilot trial lacks long-term outcomes, chiefly the concordance between preferred and received EOL care and treatments.37 However, the MS trajectory further challenges the collection of this outcome in the typical timeframe of a clinical trial. In line with the principles of ACP, we agreed not to narrow the inclusion criteria only to pwPMS in the late stage of the disease, deserving this relevant outcome to future studies.

Author affiliations
1Bioethics Unit, Azienda USL - IRCCS di Reggio Emilia, Reggio Emilia, Italy
2Fondazione FARO, Turin, Italy
3The Italian Multiple Sclerosis Society, Genoa, Italy
4Hospice “La Torre sul Colle”, USL Umbria 2, Spoleto (PG), Italy
5Department of Neuroscience, Biomedicine, and Movement Sciences, University of Verona, Verona, Italy
6Unit of Neurology, Borgo Roma Hospital, Azienda Ospedaliera Universitaria Integrata Verona, Verona, Italy
7Multiple Sclerosis Unit, IRCCS S. Lucia Foundation, Rome, Italy
8EUPATI Fellow (European Patients Academy for Therapeutic Innovation) Italy, Roma, Italy
9UOSI Riabilitazione Sclerosi Multpla, IRCCS Istituto delle Scienze Neurologiche di Bologna, Bologna, Italy
10Dipartimento di Scienze Biomediche e Neuromotorie, Università di Bologna, Bologna, Italy
11New Zealand Health and Safety Commission, Nelson, New Zealand
12Multiple Sclerosis Center, Azienda USL-IRCCS di Reggio Emilia, Reggio Emilia, Italy
13Multiple Sclerosis Center, University Hospital Policlínico Vittorio Emanuele, Catania, Italy
14UOC Neurologia, ASUR Marche, Fermo, Italy
15Department of Rehabilitation, M.L. Novarese Hospital, Moncivello (VC), Italy
16Unit of Neuroepidemiology, Fondazione IRCCS Istituto Neurologico Carlo Besta, Milan, Italy
17UOSI Riabilitazione Sclerosi Multpla ed altre Malattie Neurologiche, IRCCS Istituto delle Scienze Neurologiche di Bologna; Dipartimento di Scienze Biomediche e Neuromotorie, Università di Bologna, Italy
18Qualitative Analysis Panel: LDP, SV, MC, MJ, EP, CS, AGiordano, AS. Data Safety and Monitoring Committee: Kevin Brazil, School of Nursing and Midwifery, Queen’s University of Belfast, Belfast, Northern Ireland, UK; Bobbie Farsides, Brightton; Luciano Orsi, The Italian Society of Palliative Care (SICP), Milan, Italy; Carlo Perin, Unit of Biomedical Sciences, University of Milan, Milan, Italy; and David Oliver, The Tizard Centre, University of Kent, Canterbury, UK (Chair). Data Management and Analysis Committee: AGiordano, Mariangela Farinotti, Unit of Neuroepidemiology, Fondazione IRCCS Istituto Neurologico Carlo Besta, Milan, Italy. Qualitative Analysis Panel: LDP, SV, MC, Luca Ghirotto, Qualitative Research Unit, Azienda USL-IRCCS di Reggio Emilia, Reggio Emilia, Italy; Katia Mattarozzi, Department of Experimental, Diagnostic and Specialist Medicine, School of Medicine, Alma Mater Studiorum University of Bologna; Marta Perin, Unit of Bioethics, Azienda USL-IRCCS di Reggio Emilia, Reggio Emilia, Italy. HP Training Panel: LDP, SV, MC, KM, EP, Michela Rimondini, Section of Clinical Psychology, Department of Neuroscience, Biomedicine and Movement Sciences, University of Verona, Policlinico G.B. Rossi, Verona, Italy; AS. Linguistic Validation Panel: MF, PK, SV, AGiordano, AS. Enrolling Centers and Investigators: Department of Neuroscience, Biomedicine and Movement Sciences, University of Verona; Unit of Neurology, Borgo Roma Hospital, Azienda Ospedaliera Universitaria Integrata Verona, Verona, Italy; AGiordato, Francesca Gobbi, Riccardo Orlandi. Department of Rehabilitation M. L. Novarese Hospital, Moncivello, Vercelli: CS, Enrica Grange. Multiple Sclerosis Center, Azienda USL-IRCCS di Reggio Emilia, Reggio Emilia, Italy; SM, Francesca Sireci. UOSI Riabilitazione Sclerosi Multpla, IRCCS Istituto delle Scienze Neurologiche di Bologna; Dipartimento di Scienze Biomediche e Neuromotorie, Università di Bologna, Bologna; AL, Loredana Sabbatini, Cinzia Scandellari, Elisa Ferrari. Fondazione IRCCS Santa Lucia, Roma: MGG, Giorgia Precise. University Hospital Polyclínico Vittorio Emanuele, Catania; FP, Clara Grazia Chiaseri, Simona Toscano.

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ORCID iD

Alessandra Solari http://orcid.org/0000-0001-9930-7579

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