Self-management programme of activity coping and education—SPACE for COPD(C)—in primary care: a pragmatic randomised trial

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ABSTRACT

Introduction We have previously developed a supported self-management programme (SMP): Self-management Programme of Activity, Coping and Education for chronic obstructive pulmonary disease (COPD), which was successfully delivered on an individual basis. Patients expressed an interest in delivering the intervention in groups.

Aim To explore the feasibility, acceptability and clinical effectiveness of the intervention delivered and supported by healthcare professionals (HCPs) in groups within primary care.

Methods A prospective, single-blinded randomised controlled trial was conducted, with follow-up at 6 and 9 months. Participants were randomly assigned to control (usual care) or intervention (a six-session, group-based SMP delivered over 5 months). The primary outcome was change in COPD Assessment Test (CAT) at 6 months. Semistructured focus groups were conducted with intervention participants to understand feasibility and acceptability. A focus group was conducted with HCPs who delivered the intervention to gain insight into any potential facilitators/barriers to implementing the intervention in practice. All qualitative data were analysed thematically.

Results 193 participants were recruited, (median Medical Research Council (MRC) grade 2). There was no significant difference between the intervention and control group for the primary outcome (CAT). However, an improvement in self-reported patient activation (at 6 and 9 months), knowledge (at 6 months), mastery (at 6 and 9 months) and fatigue (at 6 months), in the intervention group compared with usual care was demonstrated. Qualitative results indicated that the intervention was acceptable to patients who took part in the intervention and HCPs valued the intervention, suggesting it might be best delivered early in the disease process.

Conclusions A supported self-management intervention is feasible and acceptable when delivered as a group-based intervention, by HCPs in the community.

INTRODUCTION

Chronic obstructive pulmonary disease (COPD) is the third-leading cause of death worldwide and is associated with considerable disability, impaired quality of life and high utilisation of healthcare resources.1 Symptoms and manifestations of the disease can be modified by adopting appropriate health behaviours including, but not limited to, exercise, physical activity, smoking cessation, anxiety management, breathing control, medication adherence and exacerbation management.2

National and international guidelines for the management of COPD suggest that self-management support should be provided for people with COPD.3 Interventions to support self-management in patients with COPD have been shown to be effective in improving health related quality of life, reducing hospital
admissions and reducing COPD specific health costs. However, trials have mainly recruited people from secondary care and excluded those with mild disease despite those with mild dyspnoea representing 38%–54% of diagnosed patients in primary care. Lahham et al explored the impact of home-based pulmonary rehabilitation (PR) in this population, but found no consistent benefits on exercise capacity, symptoms or quality of life compared with standard care. However, standard care comprised attention control, weekly social telephone calls and the provision of the ‘better living with COPD’ book which may go some way to explain similar improvements in both groups, with authors suggesting PR is not well matched to the needs of a milder group. A lighter touch self-management strategy may be beneficial. Furthermore, the support from healthcare professionals (HCPs) seems to be key in patients’ maintaining their engagement in self-management activities.

We have previously developed and described SPACE (Self-management Programme of Activity, Coping and Education) for COPD which is a structured programme of exercise, education and psychosocial support. The intervention was underpinned by social cognitive theory and motivational interviewing techniques. This was developed by our institution as a collaboration between experts, patients and carers and has been awarded a Crystal Mark for Clarity by the Plain English Campaign. In discussion with payers, there was interest in exploring the delivery of the SPACE for COPD programme as a group-based intervention in the community rather than on a one-to-one basis. The programme has also been used as remote delivery option for PR services throughout the COVID-19 pandemic.

Objectives were to address the following, using quantitative and qualitative methods:

1. Examine whether group-based delivery of SPACE for COPD, with sustained support, improves patient outcomes following the intervention compared with a control group.
2. Explore feasibility, acceptability and efficacy of the intervention to be delivered and supported by HCPs. This will be done by:
   a. Exploring HCP’s experiences of delivering the intervention and identify any barriers to delivery in practice.
   b. Understanding, from the participants’ perspective, the feasibility and acceptability of the SPACE for COPD intervention delivered by HCPs in a group-based community setting.

**METHODS**

**Trial design**

Prospective, single-blinded, randomised controlled trial. We used a mixed-methods design to determine the effectiveness and acceptability of a self-management support intervention for people with COPD, managed in primary care. A separate health economic evaluation of the intervention will be reported elsewhere. Details of the study protocol have previously been published. The trial was sponsored by the University Hospitals of Leicester National Health Service (NHS) Trust (study number 152408).

After publication of the protocol, we amended the procedure to allow us to send out questionnaires or for one of the study team to go a participant’s home to complete the primary outcome measure in order to minimise data loss of the primary outcome. This amendment came into place after recruitment started. Prior to this, all patients had to complete questionnaires within in person visits. In addition, a second Endurance Shuttle Walk Test (ESWT) was carried out at all time points if the patient walked for more than 8 min on the initial prescribed speed from the Incremental Shuttle Walk Test (ISWT).

**Participants**

Participants were recruited from seven general practices (GPs) and from those who responded to poster advertisements displayed at GP and hospitals within Leicester, Leicestershire and Rutland. Participants were also recruited from the NIHR Biomedical Research Centre—Respiratory at University Hospitals of Leicester NHS Trust.

Participants were eligible for the trial if they had an established diagnosis of COPD as defined by The Global Initiative for Chronic Obstructive Lung Disease (GOLD) criteria.

Patients were excluded from participating in the trial if they were unable to participate in the exercise component of the SPACE for COPD programme due to neurological, locomotive, psychiatric disability, or other comorbidities where exercise would be a contraindication (eg, unstable angina; unable to read/write English to the level of an 8 years old; unwilling to be randomised; previous participants of PR or have received the SPACE for COPD manual in the previous 12 months.

Baseline assessment: Participants who expressed an interest in the study were telephoned by a researcher and invited to a recruitment assessment. Following informed consent, spirometry was undertaken, demographic details were collected and the participant was asked to complete baseline questionnaires. A SenseWear accelerometer was given to them to wear on their dominant arm for 7 days, and pick up was arranged. Participants were also randomised and shuttle walk tests (SWTs) were carried out if the participant was randomised to the intervention group.

**Intervention: SPACE for COPD group-based self-management programme**

Participants in the intervention group received a SPACE for COPD manual and attended the SPACE for COPD group-based self-management programme (SMP) usually within 1 month of their baseline appointment. The programme was structured around the SPACE for COPD...
manual, which combines both generic self-management skills and disease-specific tasks. The programme was facilitated by two trained HCPs (eg, physiotherapist, respiratory specialist nurse, occupational therapist, health psychologist) to groups of up to 10 participants and delivered through six 2-hour sessions, over a 5-month period. Earlier sessions were delivered closer together in time, allowing group cohesion to take place, an important factor in optimising group dynamics.15 As much as possible, the same facilitators delivered all sessions for each group. Sessions were held at community venues, at times and locations to suit group participants to increase retention and engagement in the intervention. The content of the intervention and accompanying self-management components were described in the published protocol.17 Participants were provided with a contact number for at least one of the facilitators in case they had additional queries/were unable to attend any sessions. The exercise component of the manual was completed by participants at home. A full description of the rationale, development and efficacy of the work underpinning the SPACE for COPD manual is detailed elsewhere.13,14 Twelve HCPs attended a 1-day training course and were given an HCP delivery manual to guide the content of the sessions.

Usual care
Participants in the control group continued with any usual check-ups/reviews—no additional care was provided or removed from their current access. If participants were referred to PR in the duration of their time in the study, they were not denied access to the programme. No additional advice, information or recommendations were provided to participants in this group.

Outcomes
The primary outcome measure was health status at 6 months postbaseline assessment, measured using the COPD Assessment Test (CAT). Scores range from 0 to 40 with higher scores indicating greater clinical impact.

Secondary outcomes were COPD knowledge (using the Bristol COPD Knowledge Questionnaire, BCKQ), health-related quality of life (using the EuroQol 5 dimension, 5 level tool), disease-specific quality of life (using the Chronic Respiratory Questionnaire: CRQ), anxiety and depression (using the Hospital anxiety and Depression Score; HADS), patient motivation (using the patient activation measure: PAM) and exercise capacity (using the ISWT and ESWT—at baseline this was only performed in the intervention group to avoid the test itself acting as an intervention). Outcome measures and details of study clinical assessments and outcome measures at all appointments have previously been described in the study protocol.17 Participants were classed as completers if they attended four out of the six self-management group sessions. Intervention fidelity was also measured (online supplemental appendix 1). Intervention facilitators completed checklists at the end of each self-management group session, and one of the trainers observe one session per self-management group, completing their own checklist.

Any serious adverse events were reported to the sponsor. A serious adverse event was defined as an acute exacerbation of COPD that resulted in a hospital admission.

Qualitative outcomes
Qualitative focus groups were conducted by SC (see acknowledgements) at the end of the SPACE for COPD intervention and aimed to understand participants’ experiences of the group-based SMP. Those participants allocated to the intervention group were invited to take part in qualitative focus groups at the end of the intervention period. Focus groups were conducted with each SMP group, with between 3 and 10 participants (number dependent on each group size).

HCPs delivering the SPACE for COPD SMP were also invited to participate in a meeting to discuss aspects of feasibility and acceptability, such as gaining insight into any potential facilitators/barriers to implementing the intervention in practice (and derive practical recommendations for doing so). Sample questions are provided in the online supplement.

Acceptability findings from both the focus groups and HCP discussion will be reported here. Additional qualitative findings which may additionally contribute to the design and implementation of this programme moving forward may be reported separately.

Sample size
The power calculation was based on the primary outcome at 6 months.19,20 To detect a mean±SD between-group difference of 2.5±5.0 in the change in CAT with 80% power, 64 people per group are required (α=0.05, two tailed). In anticipation of a possible 25% attrition rate, the total sample size was increased to 86 per group (172 in total). Sample size calculations were carried out using software developed by Machin et al https://onlinelibrary.wiley.com/doi/book/10.1002/9781444300710

Randomisation and blinding
Participants were randomised by an unblinded member of the study team using an online randomisation tool (sealed envelope).21 Individuals were randomised (1:1) to the control group or the intervention group. The system randomises patients in random permuted blocks. This allows for the 1:1 ratio, but due to the random permuted blocks of 2, 4 or 6 ensures full randomisation. Participants were immediately informed of their allocated treatment by an un-blinded member of the study team. Data collection at 6 and 9 months was carried out.
by the blinded member of the study team, except the ESWT, which was carried out by an un-blinded member of the study team as only intervention participants had a prescribed ESWT speed at baseline.

**Statistical methods**

Analysis was carried out on an intention-to-treat basis. The main analysis compared primary and secondary outcome measures between treatment groups at 6 months post-baseline to assess the effect of the group-based SMP. Data were also analysed at 9 months to assess the longer-term effect. Data were checked for normality of distribution and multivariate outliers.

Missing data were imputed in Stata (V.15; StataCorp) using multiple imputed chained equations.

Full details of statistical tests are given in online supplementary appendix 2.

**Qualitative methods**

Focus groups were analysed using Thematic Analysis\(^\text{22}\) supported by NVivo software (V.9). This approach follows six distinct stages: familiarisation with data; generating initial codes; searching for themes; reviewing themes; defining and naming themes and producing the report. Authors CB and PP carried out initial coding and a sample of interviews were coded by another member of the team to ensure consistency and to enhance interpretive authenticity. Throughout the data analysis, the team met to discuss and review emerging themes and search for accounts that provided contesting views of the same phenomena or identified different phenomena. Our patient representatives were invited to comment on our (anonymised) findings throughout the analysis process to ensure any interpretations made stayed close to the direct experience of patients.\(^\text{23}\) Finally, member-checking occurred as part of a dissemination event for intervention participants—no changes were made to the acceptability results presented here. A note-based analysis using a constant comparative method was used to analyse notes taken during the HCP discussion.\(^\text{24}\)

**RESULTS**

**Participant flow**

Figure 1 shows the Consolidated Standards of Reporting Trials flow diagram of participant enrolment, allocation, follow-up and analysis. COPD, chronic obstructive pulmonary disease; PR, pulmonary rehabilitation; SPACE, Self-management Programme of Activity, Coping and Education.

**Sample size**

During initial stages of the study, 19 patients were deemed to be ineligible. There were two important implications of this: the overall sample size was lower than expected, and the group sizes had become unequal. As a consequence, the sample size was reassessed and additional participants recruited. The allocation ratio of 1:1 was maintained and as such recruitment was extended to 193 patients (of which 174 would be eligible) in order to maintain 80% power. This number (174) allows for a slightly higher level of attrition, as well as an inequality in the numbers allocated to the two groups.
Participants were recruited between March 2015 and October 2016. One hundred and ninety-three participants were randomised. At the end of the trial, data were available for the primary outcome in 65% (n=52) of the intervention group and 93% (n=70) of the usual care group. Missing data rates across variables ranged from 5/193 (2.6%; age/gender) to 101/193 (52.3%; ESWT at 9 months). This large volume of missing data for shuttle walking tests occurred when some participants were unable to attend 9-month appointments. In such circumstances questionnaires were sent by post.

Eighteen serious adverse events were reported by participants; 11 in the intervention arm and 7 in the usual care arm. None were considered to be related to the intervention. No deaths occurred.

### Outcomes and estimation

#### Primary outcome

At 6 months, there was no significant difference between groups in the total CAT score (mean difference 1.70, 95% CI −0.53 to 3.94, p=0.135). When the total CAT score at 9 months was added to the model, there was still no significant difference between groups over time (mean difference 0.511, 95% CI −1.28 to 2.30, p=0.575) (see figure 2).

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**Table 1** Baseline characteristics of all participants

<table>
<thead>
<tr>
<th></th>
<th>Intervention (n=97)</th>
<th>Control (96)</th>
<th>Overall (193)</th>
</tr>
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<tbody>
<tr>
<td>Gender: male, n (%)</td>
<td>52 (53.6)</td>
<td>53 (55.2)</td>
<td>105 (54.4)</td>
</tr>
<tr>
<td>Age: mean (SD)</td>
<td>69.6 (8.1)</td>
<td>70.5 (8.4)</td>
<td>70.1 (8.2)</td>
</tr>
<tr>
<td>Ethnicity: white, n (%)</td>
<td>86 (88.7)</td>
<td>89 (92.7)</td>
<td>175 (90.7)</td>
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<td>BMI: mean (SD)</td>
<td>27.1 (5.0)</td>
<td>27.7 (5.8)</td>
<td>27.5 (5.4)</td>
</tr>
<tr>
<td>Marital status: married, n (%)</td>
<td>51 (52.6)</td>
<td>53 (55.2)</td>
<td>104 (53.9)</td>
</tr>
<tr>
<td>Employment status: n (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Full-time</td>
<td>12 (12.4)</td>
<td>9 (9.4)</td>
<td>21 (10.9)</td>
</tr>
<tr>
<td>Part-time</td>
<td>7 (7.2)</td>
<td>7 (7.3)</td>
<td>14 (7.3)</td>
</tr>
<tr>
<td>Retired</td>
<td>66 (68)</td>
<td>71 (74)</td>
<td>137 (71)</td>
</tr>
<tr>
<td>FEV1: mean (SD)</td>
<td>1.7 (0.6)</td>
<td>1.7 (0.6)</td>
<td>1.7 (0.6)</td>
</tr>
<tr>
<td>FVC: mean (SD)</td>
<td>2.9 (0.8)</td>
<td>2.8 (0.9)</td>
<td>2.9 (0.8)</td>
</tr>
<tr>
<td>FEV1/FVC: mean (SD)</td>
<td>0.6 (0.1)</td>
<td>0.6 (0.1)</td>
<td>0.6 (0.1)</td>
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<tr>
<td>Gold stage: n (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>19 (19.6)</td>
<td>19 (19.8)</td>
<td>38 (19.7)</td>
</tr>
<tr>
<td>II</td>
<td>53 (54.6)</td>
<td>57 (59.4)</td>
<td>110 (57)</td>
</tr>
<tr>
<td>III</td>
<td>15 (15.5)</td>
<td>15 (15.6)</td>
<td>30 (15.5)</td>
</tr>
<tr>
<td>IV</td>
<td>3 (3.1)</td>
<td>2 (2.1)</td>
<td>5 (2.6)</td>
</tr>
<tr>
<td>Medical Research Council (MRC) Grade: median (IQR)</td>
<td>2 (2–3)</td>
<td>2 (2–3)</td>
<td>2 (2–3)</td>
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<tr>
<td>Smoking status: n (%)</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Current smoker</td>
<td>13 (13.4)</td>
<td>15 (15.6)</td>
<td>28 (14.5)</td>
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<tr>
<td>Ex-smoker</td>
<td>67 (69.1)</td>
<td>68 (70.8)</td>
<td>135 (69.9)</td>
</tr>
<tr>
<td>Never smoked</td>
<td>4 (4.1)</td>
<td>8 (8.3)</td>
<td>12 (6.2)</td>
</tr>
<tr>
<td>Pack years: mean (SD)</td>
<td>39.5 (25.7)</td>
<td>34.4 (27.0)</td>
<td>36.8 (26.4)</td>
</tr>
<tr>
<td>Comorbidities: n (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ischaemic heart disease</td>
<td>7 (7.2)</td>
<td>10 (10.4)</td>
<td>17 (8.8)</td>
</tr>
<tr>
<td>Hypertension</td>
<td>41 (42.3)</td>
<td>46 (47.9)</td>
<td>87 (45.1)</td>
</tr>
<tr>
<td>Diabetes</td>
<td>13 (13.4)</td>
<td>13 (13.5)</td>
<td>26 (13.5)</td>
</tr>
<tr>
<td>Heart failure</td>
<td>1 (1.0)</td>
<td>3 (3.1)</td>
<td>4 (2.1)</td>
</tr>
<tr>
<td>Arthritis</td>
<td>24 (24.7)</td>
<td>35 (36.5)</td>
<td>59 (30.6)</td>
</tr>
<tr>
<td>Other</td>
<td>21 (21.6)</td>
<td>22 (22.9)</td>
<td>43 (22.3)</td>
</tr>
<tr>
<td>None</td>
<td>13 (13.4)</td>
<td>11 (11.5)</td>
<td>24 (12.4)</td>
</tr>
</tbody>
</table>

BMI, body mass index; FEV1, forced expiratory volume in 1 s; FVC, forced vital capacity.
Table 2  Comparison of primary and secondary outcomes

| Characteristic | Baseline | | | 6months | | | 9months | | |
|---------------|----------|----------|----------|----------|----------|----------|----------|----------|
|               | Intervention | Control | Intervention | Control | Intervention | Control | Intervention | Control |
| Mean (95% CI) | Mean (95% CI) | Mean (95% CI) | Mean (95% CI) | Mean (95% CI) | Mean (95% CI) | Mean (95% CI) | Mean (95% CI) | Mean (95% CI) |
| CAT | 15.5 (14.0 to 17.1) | 14.0 (12.5 to 15.5) | 16.9 (12.0 to 21.7) | 15.9 (14.4 to 17.4) | 15.8 (12.8 to 18.7) | 14.7 (13.2 to 16.3) |
| Dyspnoea | 3.4 (3.0 to 3.7) | 3.3 (3.0 to 4.2) | 3.9 (3.5 to 4.2) | 3.8 (3.5 to 4.2) | 4.0 (3.6 to 4.4) | 3.9 (3.6 to 4.2) |
| Fatigue | 4.6 (4.3 to 4.8) | 4.6 (4.4 to 4.9) | 4.8 (4.4 to 5.1) | 4.4 (4.2 to 4.7) | 4.6 (4.3 to 5.0) | 4.5 (4.2 to 4.8) |
| Emotion | 5.3 (5.0 to 5.5) | 5.3 (5.1 to 5.6) | 5.3 (5.0 to 5.6) | 5.2 (5.0 to 5.5) | 5.3 (5.0 to 5.6) | 5.2 (5.0 to 5.5) |
| Mastery | 5.6 (5.3 to 5.9) | 5.9 (5.6 to 6.1) | 5.9 (5.6 to 6.3) | 5.8 (5.5 to 6.0) | 6.0 (5.7 to 6.3) | 5.8 (5.6 to 6.1) |
| Anxiety | 4.8 (4.0 to 5.7) | 5.6 (4.9 to 6.4) | 4.8 (3.8 to 5.7) | 5.5 (4.7 to 6.3) | 4.8 (3.8 to 5.7) | 5.3 (4.5 to 6.1) |
| Depression | 3.4 (2.9 to 4.0) | 3.6 (3.0 to 4.1) | 3.2 (2.5 to 3.9) | 3.9 (3.3 to 4.5) | 3.6 (2.9 to 4.3) | 3.8 (3.2 to 4.4) |
| BCKQ Score | 35.2 (33.5 to 36.9) | 32.7 (31.1 to 34.4) | 38.6 (36.8 to 40.5) | 33.7 (31.9 to 35.4) | 39.7 (37.7 to 41.8) | 35.4 (33.7 to 37.1) |
| PAM Score | 59.2 (56.3 to 62.1) | 59.8 (57.0 to 62.6) | 64.4 (60.7 to 68.1) | 59.3 (56.3 to 62.3) | 65.3 (61.7 to 69.0) | 59.2 (56.4 to 62.1) |
| PAM Level | 2.6 (2.4 to 2.8) | 2.7 (2.5 to 2.9) | 2.9 (2.6 to 3.1) | 2.6 (2.4 to 2.9) | 3.0 (2.8 to 3.3) | 2.6 (2.4 to 2.8) |
| ISWT m | 400.7 (362.8 to 438.7) | – | 424.9 (380.8 to 469.0) | 395.1 (352.8 to 437.3) | 401.1 (359.0 to 443.1) | 376.3 (335.8 to 416.7) |
| ESWT s | 432.0 (342.7 to 521.3) | – | 596.6 (491.1 to 702.0) | 467.0 (376.2 to 557.7) | 524.4 (417.4 to 631.4) | 483.8 (389.3 to 578.2) |

BCKQ, Bristol COPD Knowledge Questionnaire; CAT, COPD Assessment Test; COPD, chronic obstructive pulmonary disease; CRQ-SR, Chronic Respiratory Questionnaire; ESWT, Endurance Shuttle Walking Test; HADS, Hospital Anxiety and Depression Scale; ISWT, Incremental Shuttle Walking Test; PAM, Patient Activation Measure.
Secondary outcomes
The mean difference in PAM score was significant at 6 months, in favour of the intervention group (mean difference −5.66, 95% CI −10.03 to −1.29, p=0.011). This difference was maintained at 9 months (mean difference −6.70, 95% CI −10.85 to −2.56, p=0.002) (see figure 3). The mean difference in PAM level was also significant at 6- (mean difference −0.36, 95% CI −0.69 to −0.03, p=0.031) and 9 months (mean difference −0.57, 95% CI −0.90 to −0.23, p=0.001), in favour of the intervention group (see figure 4).

The mean difference in BCKQ score was significant at 6 months, in favour of the intervention group (mean difference −2.53, 95% CI −4.59 to −0.47, p=0.016), this difference in both groups was sustained at 9 months but was not significantly different between groups (mean difference −1.88, 95% CI −4.16 to 0.40, p=0.106) (see figure 5).

The mean difference in the fatigue (mean difference −0.42, 95% CI −0.81 to −0.03, p=0.035) and mastery (mean difference −0.44, 95% CI −0.78 to −0.10, p=0.011) domains of the CRQ were significant at 6 months, and for the mastery domain this difference was sustained at 9 months (mean difference −0.44, 95% CI −0.78 to −0.10, p=0.011) (see figure 6).

There are no significant differences in the CRQ Emotion and Dyspnoea subscales, HADS, ISWT or ESWT at 6 months or 9 months (table 2 and online supplemental appendix 3).

Qualitative results: acceptability of the programme: participants
Data are discussed below; covering delivery practicalities and integration of the SPACE for COPD manual within the SMP (see table 3 for example quotes).

Were the venues suitable?
Overall, venues and locations were well received. Parking was also important for those driving to venues. For some, venue proximity was more important than venue quality. For others, proximity wasn’t a problem—they could get a lift or share lifts with other group members.

What did they think of the session timings?
Generally, participants were happy with the time of day groups were run, with only a couple saying they would have preferred a different time of day (eg, morning instead of afternoon). Mainly, participants were happy with session length—only one smaller group thought session could be condensed. Finally, everyone was satisfied with length of time between sessions—though one group suggested a month as the longest time between sessions, stating 6 weeks might result in a slight decrease in motivation.

How was the SPACE for COPD manual used?
Everyone thought that the SPACE for COPD manual complemented group activities in sessions. Positive
aspects of the manual included: good reference point for the future, informative, encompasses everyone and did not ‘preach’. Aspects most liked were: exercise charts, breathing techniques and overall advice given. Learning points included: expensive resource that some might not look at again in the future, could be condensed, strength training could be clearer.

**Discussion with HCPs**

Main themes arising from the HCP discussion are outlined in table 4. These cover learning gained from delivering the SPACE for COPD group-based SMP, how this learning has resulted in a general change in their professional practice and thoughts about future programme implementation.

**DISCUSSION**

We developed and delivered a novel SMP that was acceptable (to participants and HCPs) in a group and community-based format delivered by those HCPs likely to deliver the intervention in practice. The novelty of our approach included group sessions being spread out further across a longer period than traditional SMPs, with earlier sessions being closer together in time to allow group cohesion to take place.

We did not observe a difference in our primary outcome of health status measured by the CAT. Both groups had an improvement of 1.4 and 1.9 points for the intervention and control groups, respectively (a change of 2 is deemed clinically relevant\(^20\)). It may be that the intervention was not intense enough to make a difference in this outcome. However, we did demonstrate an improvement in self-reported patient activation (at 6 and 9 months), knowledge (at 6 months), mastery (at 6 and 9 months) and fatigue (at 6 months), in the intervention group compared with usual care. PAM scores improved more than the minimum important difference (MID) of 4 points\(^25\) at 6 and 9 months, compared with baseline in the intervention group. This would indicate improved activation and increased self-management skills. There were no statistically significant improvements in anxiety and depression, exercise capacity, dyspnoea or emotion at 6 or 9 months. However, clinically relevant improvements were noted for dyspnoea in both groups (more than the minimal important difference of 0.5 at 6 and 9 months\(^26\)) and for the ESWT at 6 months in the intervention group this was approaching the MID of 174 s.\(^27\) Qualitative results demonstrated that the intervention was acceptable to patients who took part in the intervention and that the SPACE for COPD manual was well received. HCPs also valued the intervention, suggesting it might be best delivered earlier on in the disease process.

MRC and CAT scores demonstrated that overall, participants were milder and less symptomatic compared with a typical PR population.\(^28\) The median MRC score was 2, and our median baseline CAT score was 14 (IQR 10–20; mean 14.6). As a comparison, a PR audit conducted around the same time as this study\(^29\) noted a median CAT score was

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**Table 3  Example quotes from intervention participant focus groups**

<table>
<thead>
<tr>
<th>Theme</th>
<th>Example quote</th>
</tr>
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<tbody>
<tr>
<td>Venues</td>
<td>It’s more important to me to be near where I live than have a fantastic venue that’s 20 miles away (Female, Group 5) For me the venue was ideal and it was the thing that concerned me when it was mentioned that they would look at where the members of the group lived and try and make it local and for me in Kirby of course, no problem, and superb, the venue’s good (Male, Group 8)</td>
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<tr>
<td>Session timings</td>
<td>…I don’t know whether it’s because we’re such a small group, but overall my impression was I thought it could be a bit more condensed into perhaps a—and I thought 2 hours is a long time, but that’s only my opinion (Female, Group 4) Well, I think myself it was a good idea, because if you’d have come every week, if you hadn’t developed from the week before you wouldn’t feel very nice, would you, whereas if you went, say, every 3 or 4 weeks as it is you’d see a bit of an improvement and that made you feel a lot better (Male, Group 9)</td>
</tr>
<tr>
<td>SPACE for COPD manual</td>
<td>Yeah because it was like a starting basis for conversations wasn’t it and discussions (Male, Group 3) This and, you know, to take home and then you can digest it a bit longer, you know, the different things. And it’s not preaching at you. You know, it’s more advice rather than actual you’ve got to do it (Female, Group 6) Yeah I think it could be abbreviated, because there’s quite a number of bits of repeats in there (Female, Group 7)</td>
</tr>
</tbody>
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COPD, chronic obstructive pulmonary disease; SPACE, Self-management Programme of Activity, Coping and Education.
Table 4  HCP discussion themes

<table>
<thead>
<tr>
<th>Theme</th>
<th>Discussion points</th>
</tr>
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<tbody>
<tr>
<td>Changes in professional practice</td>
<td>The intervention taught HCP’s to allow patients to talk more and explore their own answers/reasons. HCPs felt that in their profession they wanted to give answers/make suggestions straight away, but have learnt to allow silence and time for patients to explore answers/reasons for themselves.</td>
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<tr>
<td>Where this programme might fit in practice</td>
<td>HCP’s thought group-based COPD could be a good intervention at diagnosis—GP don’t have time to explain COPD, diagnosis, what to expect etc. SPACE could be that introduction as it covers healthy lifestyle, walking, exercising etc. At diagnosis, patients may also feel they are too good for PR, and may not be offered rehab until later on in the disease process.</td>
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<tr>
<td>Benefits of the group-based design</td>
<td>Longevity of the groups allows ownership and responsibility to develop among patients. The time between sessions allows the group to come together, and patients looked forward to coming back. They used the time to work on their goals, join the gym/swimming sessions/walking groups, thus allowing longer time for support.</td>
</tr>
<tr>
<td>Group size and management</td>
<td>Often if the group size is too small it can be a struggle. A good size is approximately 10 people; bigger groups tend to be more respectful than smaller groups and as they require more management it seems easier to keep track of time. It can be tricky to manage patients with such a wide range of MRC grade 1–5, compared with PR when all patients have a limitation of some sort. However, this improved with time and HCP’s felt that they learnt how to manage the diversity of patients’ expectations in the groups.</td>
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</tbody>
</table>
| Miscellaneous | ► SPACE could be used as part of tailor made/personalised medicine style approach where it is a part of a suite of services offered, and the clinician and patient decide together, the best option.  
► The training day was really useful, and vital to delivering the sessions well and using the right techniques such as MI.  
► In the future it could be delivered by practice nurses who are respiratory specialists. Need to ensure they are happy managing groups. |

COPD, chronic obstructive pulmonary disease; GP, general practice; HCP, healthcare professional; PR, pulmonary rehabilitation; SPACE, Self-management Programme of Activity, Coping and Education.

22 (IQR 16–28) and 66% of patients were MRC 3 or 4. These numbers are also comparable to participants in the Kon et al study, determining the MCID for CAT, where the mean CAT score at baseline was 21.4. As our scores are significantly lower, it is possible that we achieved a floor effect in the intervention group whereby change may be unlikely to be achievable. Other studies have also demonstrated similar improvements in intervention and control groups for QoL measures, for example, Chaplin et al. As such, it could be hypothesised that attending a specialist respiratory assessment and being part of a research study could increase the likelihood of patients seeking information about, and take a more active role in, self-management of their COPD (including increased confidence to undertake physical activity or training). Conducting exercise tests as part of this assessment would have further increased confidence to undertake physical activity or training and as such were not included in the baseline assessment for control group participants.

Patient activation (defined as patients’ knowledge, skills and self-efficacy regarding self-management) is shown to play a central role in COPD self-management behaviours and is a central component in the Chronic Illness Care Mode. It is important to help patients put knowledge and skills into practice in daily life rather than simply telling them what they need to do—we went some way to doing this with the format of the group sessions. Goal setting was carried out between sessions that were gradually spaced further apart so that participants were implementing these goals in their daily lives for longer, still with the promise of feedback to HCPs on returning to a group session. However, qualitative feedback suggests that 6 weeks may have been too long and that 4 weeks may have been more suitable. In the intervention group, PAM scores increased on average by almost five points, similar to findings in other self-management studies. Despite this manuscript describing a lighter touch intervention. This is an important finding as activated patients are more likely to report improved health-related behaviours and clinical outcomes, and partake in collaborative decision-making with HCPs. Furthermore, each point increase in PAM score correlates to a 2% decrease in hospitalisation and 2% increase in medication adherence. Therefore, these data indicate that the intervention has the potential to reduce healthcare use.

Disease-specific knowledge increased postintervention supporting the effect of the intervention and at 3 months post intervention, this knowledge was retained (slight increase in scores between 6 and 9 months in both groups). Reduction of disease-specific knowledge in the months following the completion of SMP and rehabilitation programmes is a common finding, therefore, we were encouraged to see this outcome maintained at 9 months. New international guidelines could offer some guidance when designing the education component of self-management and rehabilitation programmes in the future.
Strengths and limitations

Offering a community-based programme, at venues and times more suitable for participants, went some way to remove common barriers to attendance. A high number of intervention participants could be classed as completers of the intervention (mean number of group sessions attended was 3.88 and 62.5% completed four or more sessions)—this, alongside qualitative findings, seems to indicate that conducting sessions in the community, local to the participant, at times suitable to them, better enabled participants to engage with and complete the SMP. It is hard to assess if the adherence rates to the intervention in this study are comparable to that of one-to-one sessions in Mitchell et al as this was not measured. However, both studies had high completion for the 6 month assessments from participants in the intervention groups (Mitchell et al 80%, this study 83%). Our inclusion criteria allowed for recruitment of a wide range of participants, not just those who would usually be suitable for PR. Indeed, the median MRC of our recruited population was grade 2. This group is often excluded from PR programmes. The intervention was underpinned by social cognitive theory and motivational interviewing techniques. It also included other techniques such as goal setting and solution-focused goal feedback that have been shown to be effective in modifying behaviour. Further, it was delivered at an intensity that was possible to incorporate into routine practice. The way in which the group sessions were delivered maximised support from contact with an HCP and allowed for rapport to develop between participants and HCPs in the initial stages of the intervention, prior to longer breaks between sessions.

For some intervention participants, distance from group venues was a problem, causing withdrawal from the study. This is because they may have had to attend sessions that were outside of their local area as we were recruiting in multiple areas at the same time. For the same reason, other participants were waiting up to a month for group sessions to start. Future studies may wish to consider cluster randomisation over individual participant randomisation to avoid these issues. In practice, this would occur naturally as GP would most likely run SMPs for their patients, ensuring a venue that is suitable for all. While group-based interventions have some advantages in terms of peer-support and likely cost savings; some patients dislike group formats. Also, this study was conducted prior to the start of the COVID-19 pandemic which limited in-person group-based interventions. Future implementation work therefore may need to consider these factors. It is likely that a choice of formats (eg, one to one, group-based, in-person, remote) would help to improve uptake and engagement from both patients receiving the intervention and services providing this.

Results suggest that the CAT may not have been the most suitable primary outcome measure for patients with milder disease. It may be that a general self-management measure (eg, HEI-Q or the PAM are more suitable. These would also measure the important behaviour change aspects of the SMP. While we have not compared the outcomes from this study to those delivering the SPACE for COPD intervention on a one to one basis; we were able to show comparable changes in the CRQ-dyspnoea domain (the primary outcome in the Mitchell et al study) at 6 months in the intervention groups: 0.5 in the current study and 0.60 when delivered 1:1, both meeting the MCID. The Mitchell et al study did not measure the CAT.

Qualitative results indicate acceptability of the intervention on behalf of the patients’ taking part and the HCPs delivering the sessions. Although the focus groups were only completed in those available at the end of the intervention period, all patients were invited to take part regardless of the number of sessions they attended in total. Despite the qualitative results suggesting that the intervention was acceptable, there were a large number of participants who did not respond to the study invitation (81.5%) and a further 36% who declined to take part. Unfortunately, a number of participants were also withdrawn by the study team that fell within the intervention arm of the study. Although the study was powered for the CAT at 6 months, recruitment was uneven between groups. Reasons for withdrawal were not related to the intervention in the vast majority of cases—many were ineligible (did not have a diagnosis of COPD—as a consequence, the sample size was reassessed and additional participants recruited.) For those who were eligible, social problems and other health problems were often cited as reasons for withdrawal. In many cases participants withdrew from the study before commencing the intervention.

Future implications

The importance of self-management is widely acknowledged in people living with COPD and opportunities should be maximised from the time of diagnosis through to more severe disease. However, there is currently no provision for a structured supported SMP in the UK for those with milder disease. Although mild disease was not an eligibility criterion for this study, we did primarily recruit those in GOLD stages I and II with a median MRC grade if 2. SPACE for COPD as a group-based community situated intervention could therefore be offered to newly diagnosed patients with COPD, with less severe problems and those with capacity to manage their disease, to bridge the gap between diagnosis and offering of PR. For those patients with more severe disease, and may have comorbidities, PR would assume greater importance.

This prevention model has only recently been adopted in COPD, with calls for interventions to reduce risk in people with early disease and potential for considerable health and health service gains if we could facilitate self-management support in patients with early disease and...
slow their decline. In order to establish whether this is possible, long-term follow-up studies are required.

Opportunities to improve self-management skills should also be embedded in PR programmes. In the future, there may be an opportunity to explore the value of the SPACE for COPD programme alongside rehabilitation, or indeed, an alternative for those unwilling or unable to attend. Currently the programme is also being tested as a group based intervention for those who have recently completed PR as a maintenance tool. The SPACE for COPD group-based SMP offers patients a choice which may be more convenient for those who may still be working and unable to commit to traditional PR, or for those struggling to travel and are only able to travel in the local area. The nature of the model also allows for patients to access specialist teams for a longer period of time but also encourages the group to meet outside of the scheduled sessions, creating in effect a local COPD community with shared interests. In future, a more intense version of the programme may be warranted to increase gains in the outcomes of interest.

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Contributors LH-W and CB were involved in the development of the intervention, the design and delivery of the trial. CB, PP, LH-W, JB and SS have been involved in drafting the work or revising it critically for important intellectual content and have given the final approval of the version published. JB completed the data analysis. SS is the principal investigator of the study and guarantor of the work.

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Competing interests None declared.

Patient and public involvement Patients and/or the public were involved in the design, or conduct, or reporting, or dissemination plans of this research. Refer to the Methods section for further details.

Patient consent for publication Consent obtained directly from patient(s).

Ethics approval Ethical approval was granted by the Hampshire B Research Ethics Committee (REC reference: 14/SC/1169). Participants gave informed consent to participate in the study before taking part.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available on reasonable request.

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REFERENCES


APPENDIX 1.

Intervention fidelity

Agreement between raters was 71.7% (range 44.4%-100%). Facilitators tended to report enacting more motivational interviewing techniques than observed by the observer. The most common of these were:

- Facilitator circulates regularly during activities to enable them to see how participants are progressing;
- Facilitator asks questions at the end of each activity to bring participants to self-reflect (enabling self-reflection);
- Facilitator uses other group members to solve participant problems before offering own advice.
APPENDIX 2.

Statistical Methods

Analysis was carried out on an intention-to-treat basis. The main analysis compared primary and secondary outcome measures between treatment groups at 6-months post baseline to assess the effect of the group-based SMP. Data were also analysed at 9-months to assess the longer-term effect. Data were checked for normality of distribution and multivariate outliers.

Missing data were imputed in Stata (version 15; StataCorp, College Station, TX, USA) using multiple imputed chained equations.

Pearson correlations (univariable) were carried out to determine which variables appeared to predict outcome variables. In each case, the same 58 candidate predictors were investigated (primary and secondary outcome measures at baseline, 6- and 9-months, age, gender, marital status and FEV_1/FVC ratio). Variables that were highly correlated with the outcome of interest (above 0.7) were not included in the imputation model in order to reduce to likelihood of perfect prediction. Linear regressions were carried out until an optimal analytic model was reached using backwards selection. In all cases, at least 40% of the variance was explained by the imputation model. In total, 22 variables were imputed (primary and secondary outcome measures (including subscales) at 6- and 9-months).

30 imputed datasets were created using primary and secondary outcome variables at baseline, 6-months and 9-months as predictors (CAT, CRQ, HADS, PAM, EQ-5D and BCKQ). Analyses on imputed datasets were carried out using the mi combine command in Stata, which analyses each dataset separately and combines the results. Linear regressions were
performed on 6-month and 9-month data for primary and secondary outcomes, and an additional regression was performed to correct for baseline differences.

Linear regressions and paired-sample t-tests were carried out ‘per protocol’ to validate the un-imputed results. Additionally, per-protocol analyses were performed with the participants who completed the primary outcome measure at 6-months, and the findings were compared with the intention-to-treat analyses. All statistical significance tests were 2-tailed with $\alpha = 0.05$.

**Secondary analyses**

We conducted additional exploratory mixed model analyses, looking at the effects of group and time. 30 imputed datasets were created using primary and secondary outcome variables from the primary analysis as predictors. Models included treatment group, time, group $\times$ time interaction and a random effect for participants.
Appendix 3.

Additional non-significant results:

Figure 1. Hospital Anxiety and Depression Scale – Anxiety Domain

![Graph showing HADS Anxiety Score for Baseline, 6 Months, and 9 Months for Intervention and Usual Care groups.]

Figure 2. Hospital Anxiety and Depression Scale – Depression Domain

![Graph showing HADS Depression Score for Baseline, 6 Months, and 9 Months for Intervention and Usual Care groups.]

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