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Results. Analysis of transcripts resulted in three superordinate themes: (1) The thirst for comparative friendship; (2) Conjecture and the imminent choice; (3) Progressive and prospective pain management. These themes reflect a desire for empathic, socially comparative friendships and the search for a forum in which to enhance pain self-management strategies, yet also internal conflict over initial CPSG attendance.

Conclusion. Social support and associated friendships are attractive to prospective CPSG members and are conceptualised as opportunities to engage in social comparison and nurture self-care. The first visit to the support group presents a significant hurdle, but can be facilitated by managing the transition between therapeutic care and CPSG attendance. Clinicians can challenge preconceptions, foster positive viewpoints regarding the group and support collective decision-making to attend. Following initial attendance, psychosocial wellbeing was enhanced.
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Running head: Decisional balance and the pain support group

The decisional balance: an interpretative phenomenological analysis reflecting the transition from pain management services to chronic pain support group attendance

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Statement of contribution

What is already known on this subject?
- Social support functions as a protective buffer against declining health
- Joining a peer-led support group can be initially intimidating and the first visit presents a significant hurdle

What does this study add?
- Participants are predominantly attracted to support groups due to the opportunity to develop new friendships
- Health-related peer groups function as forums for social comparison, enhancing self-esteem and self-efficacy
- Experience of pain management programmes primes willingness to attend support groups
- The initial decision to attend is difficult but facilitated by collective, group decision-making processes
- Health-care professionals dynamically prime the transition towards peer support structures
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Chronic pain is a significant and widespread issue, causing high levels of distress and disability (Williams, Eccleston, & Morley, 2012). Coping with chronic pain represents a major challenge for the patients who experience this long-term condition and the healthcare professionals (HCPs) who work to support them. Structured interventions for managing chronic pain have become recommended practice across primary and secondary care (British Pain Society, 2013), and treatment guidelines are multidisciplinary, covering a wide-range of educational, medical and psychosocial dimensions within formalised Pain Management Programmes (PMPs).

**Challenges to multidisciplinary pain management**

PMPs incorporating Cognitive Behavioural Therapy generate a moderate impact on measures of disability, mood and pain catastrophizing, and ‘third-wave’ acceptance-based programmes have engendered positive functional and psychological outcomes (Fedoroff, Blackwell, & Speed, 2014). However, such outcomes may diminish with time (Ehde, Dillworth, & Turner, 2014). A recent systematic review concluded that the only persisting outcome at 12-months following PMP completion, when compared with a wait-list control, was mood (Williams et al., 2012). Similarly, when PMPs were compared against active controls, only disability was reduced at 12-month follow-up. With low back pain, pain intensity was reduced immediately after PMP, but quality of life and depression were not improved (Hoffman, Papas, Chatkoff, & Kerns, 2007). To enhance the longevity of PMP outcomes, there is a clear need for additional care beyond the programme end.

It is increasingly evident that an individualistic treatment-focus is limited: pain is experienced in a social world and coping is often undertaken in the context of wider support structures (Riekert, Ockene, & Pbert, 2014). Responding to pain is a function of social learning, with pain-related behaviours, beliefs and attitudes formed through childhood and interpersonal modelling (Goubert, Vlaeyen, Crombez, & Craig, 2011). The lived experience of facing a chronic health condition is resolutely social; patients develop an understanding of their condition and expectations for their future through interpersonal exchanges with healthcare professionals (HCPs) and wider social relationships (Davison, Pennebaker, & Dickerson, 2000). Therapeutic interventions isolated from social environments fail to reflect the reality of living with chronic
pain in a social world (Sullivan, 2012). Treatment failure or limited outcomes may reflect difficulty in transferring learned skills from a clinical setting to social environments (Linton, Helsing, & Larsson, 1997), therefore there is a need for research that approaches pain management through the context of social support.

**The role of social support**

Social support has been conceptualised as emotional, informational and instrumental assistance provided by significant others (House, Kahn, McLeod, & Williams, 1985). It is a multifaceted phenomenon, promoting personal and societal well-being. Social support can be activated through; (i) primary, enduring, informal support structures (e.g. family and close friends), or (ii) secondary, larger, more formalised groups (e.g. hobby groups or work colleagues) (Thoits, 2011). Irrespective of type of social support, outcomes are almost uniformly positive; for example, reduced stress-related reactivity (Lovell, Moss, & Wetherell, 2012), enhanced psychological well-being (Taylor & Stanton, 2007), reduced risk of cardiovascular disease (Compare et al., 2013; Gallagher, Luttik, & Jaarsma, 2011), increased longevity (Uchino, 2009), heightened quality of life, coping and adjustment (Thoits, 2011). Where chronic illness represents a stressor, social support functions as a protective buffer, insulating the individual from decline in physical/mental health (Uchino, Carlisle, Birmingham, & Vaughn, 2011; Uchino, 2009). Such social support structures are potentially invaluable as patients often report a contraction in social opportunities as their condition persists (Rodham, McCabe & Blake, 2009).

**Chronic Pain Support Groups (CPSGs)**

To counteract social isolation, a wide variety of CPSGs exist, providing predominantly peer-to-peer support structures (Embuldeniya et al., 2013). CPSGs are usually volunteer-led, socially active and well-attended by chronic pain patients from wide-ranging social and cultural backgrounds (Cowan, 2013; Subramaniam, Stewart, & Smith, 1999). CPSGs are ordinarily extra-medical, typically providing peer networking, social events and speakers in an informal meeting structure (Cowan, 2013). Following discharge from PMP, it is recommended that additional support is sought from regional CPSGs (British Pain Society, 2013). However, minimal research has investigated the transition from PMP to CPSG and it is not known why people decide to
engage with CPSGs. Joining a CPSG represents a patient-centred, considered decision to seek out social support and engage with self-care (Kitson, Marshall, Bassett, & Zeitz, 2013). Research is needed to investigate the rationale for CPSG membership, identifying what encourages attendance and which factors consistently influence the decision-making process.

Current study
This study aimed to investigate issues surrounding participants’ initial decision to attend a CPSG after PMP completion, through retrospective reflection. Interpretative Phenomenological Analysis (IPA) was used to explore the decisional balance (Prochaska et al., 1994), rationale and motivations for CPSG attendance as it offers opportunity for insight into the internal world and the self as a sense-making agent (Smith, Flowers & Larkin, 2009). Such an idiographic stance is considered crucial for representing the unique pressures surrounding decision-making processes (Ando et al., 2015).

Method
Participants
Twelve participants (8 females, 4 males) were purposively sampled (following Smith, Flowers, & Larkin, 2009) from Milton Keynes CPSG. Potential participants were invited to take part if they suffered from chronic, non-malignant pain, had previously completed a six-week PMP through Milton Keynes General Hospital, had attended the CPSG for two or more years, had attended a minimum of 50% of meetings in the past 12 months and were fluent English speakers. All participants were currently retired or unable to work due to their health status. Demographic characteristics are shown in Table 1. All identifying information has been changed.

Procedures
The current study was reviewed and ethical approval granted by the University of [Redacted] Department of Psychology Research Ethics Committee. Participants were approached, following permission from the CPSG Committee, via their newsletter and through a presentation at a support group meeting. Prospective participants were given the study information sheet and were asked to contact the researcher if they wished to take part. Written, informed consent was
obtained prior to commencement of interviews, which were conducted in participants’ homes (N = 11) or at the University of (N = 1) and lasted between 45 and 120 minutes. All participants were debriefed, received a £30 gift voucher for their time and travel costs reimbursed.

**Data collection and analysis**

To facilitate openness within interviews and encourage participants to discuss issues central to their decisions surrounding joining the CPSG, an open-ended semi-structured interview style was used (Smith et al., 2009). Participants were given freedom to lead the interview and make sense of their experiences through highlighting issues important to them from their own perspectives, whilst still allowing the researcher freedom to probe further on topics related to the aims of the study (Smith, 2011; see Table 2 for interview schedule). Interviews were all conducted by the first author, audio-recorded and transcribed verbatim.

A ‘bottom-up’, idiographic stance was adopted for first-stage analyses, with each transcript analysed independently in full, to reflect the experience of each individual in its own light and allow space for researcher reflection (Smith et al., 2009). Each transcript was analysed recursively, constantly moving between themes and transcript to ensure the analyses were grounded in the data. After an initial reading for familiarity, transcripts were read in-depth for descriptive, linguistic and conceptual content, which collectively formed emergent themes in conjunction with the researcher’s reflective notes. A cross-case analysis was then undertaken, abstracting super-ordinate and clustered subordinate themes in order to demonstrate relationships, convergences and divergences, developed and grounded in the data. Quotations presented in results represent key extracts, chosen to best highlight the theme under discussion (Smith et al., 2009).

***Insert table 2 about here***

**Quality and rigour**

Efforts were made throughout the analyses to maintain a stance of sensitivity, transparency, coherence and rigour through idiographic, iterative practice (Yardley, 2007). An independent auditor with expertise in IPA examined the emergent themes, triangulating clustered themes. The auditor worked closely with transcripts to ensure fidelity to original interviews was
maintained. Interpretations of quotations were discussed to ensure themes were an accurate representation of the data. Due to the interpretative nature of the analysis, it is recognised that the researcher's own views may impact upon the representations of interviewees' thoughts, however efforts were made to 'bracket-off' assumptions and pre-conceptions through the use of a reflective log (following Smith et al., 2009). It is recognised that the 'double-hermeneutic' is part of the active sense-making process by the researcher when conducting IPA; as such, the current interpretation may not be the sole interpretation of accounts. The current research therefore focuses on expression of the experience of participants rather than an application of 'top-down' theory to transcript data (Vangeli & West, 2012).

***Insert table 3 about here***

Results
Analysis of transcripts resulted in three superordinate themes: (1) The thirst for comparative friendship; (2) Conjecture and the imminent choice; (3) Progressive and prospective pain management. Super-ordinate and subordinate themes are presented in Table 3.

The thirst for comparative friendship
The first super-ordinate theme was evident across all participants and represented the powerful role of present and future interpersonal relationships in deciding to invest in CPSG membership.

Catching the lifeline of continued support. During the PMP, participants reported hearing about the CPSG through a presentation by current members. Hearing about the group caused feelings of anxiety and frustration, in particular because it represented discharge from hospital-based pain services and yet seemed a great opportunity, therefore participants felt conflicted between their desire for support yet concern over ‘losing’ their close clinical supervision by pain specialists. This conflict was clearly expressed by Nicole as she felt that the group would have been useful for her before the end of the PMP:

"In my head at that time I was thinking ‘why wasn’t I aware of this before’? Because I would’ve loved to go ‘cause it was an extension of what we were doing.” (Nicole).
For the majority, the conflict was short-lived and the decision to attend was rapid and natural, with their decision-making assisted through the information provided at the PMP: they felt like the CPSG would offer a lifeline, opportunity and a wise choice. Engaging with the group was perceived as a way of ensuring that there was a safety net:

“It is my lifeline. If I didn’t have this, the pain would overtake me.” (Laura).

The majority of participants demonstrated a strong, predominant opinion that joining the group was not optional, but a compulsory and non-negotiable step, a metaphorical safety parachute to ensure adequate support provision in the future:

“And just knowing that support’s there (.) you may never, ever use it, it’s like flying... you have a parachute on, you may never ever use it, but it’s there. It’s compulsory.” (Ron).

The recommendation that PMP completers join the CPSG was not viewed neutrally as an optional decision, but instead, because it had been strongly recommended by HCPs on the PMP, participants felt they needed to adhere to this advice. The need for adherence was not, however, seen as pressure from clinicians, but was a valued recommendation arising from trust built through a successful PMP:

“I was thinking that I’d been told it was a good idea. I trusted what they said because I’d found the actual group, the programme, very useful. I thought ok, they recommend this, I’ll give it a go.” (Jane).

The friendship agenda. All participants expressed sadness, but a sense of inevitability about their contracting social circles since developing chronic pain. They saw chronic pain as the prime reason for losing their so-called ‘normal friends’: they felt they were now perceived to be ‘hard work’ and old friends were not willing to fight to continue with friendships, causing a mutual lack of motivation to maintain such friendships. This was a vicious cycle that Stuart discussed:

“I suppose the best way to put it is that people who were close and regular friends are no longer [...]. So from a circle of friends that was yay big, it just goes ‘whoosh’. And it’s very easy to get into a cycle where I’ve got nobody to see so I’m not going to go out.” (Stuart).
Consequently, all participants expressed a key desire to build friendships and were motivated to see the CPSG as an opportunity to put this into action. For Charlotte, it was the PMP itself that highlighted this need for friends: doing the PMP in small groups fostered close relationships and a sense of openness and emphasised the need for new friendship groups:

"And that [the PMP] I think brought me out of myself quite a bit and made me want to see if I could go and find somewhere that, y’know, some sort of group of people that I could spend time with.” (Charlotte).

Participants all articulated the difference between friends with pain and friends without; friends who understood pain were conceptualised as people enabling you to move forwards and stretch your capacity for coping, therefore promoting greater adjustment to living with chronic pain. The desire to build friendships was expressed cautiously, suggesting a sense of insecurity as a result of pre-CPSG loss of friends but also elements of self-doubt and self-preservation as participants perhaps did not want to invest too much hope in prospective but hypothetical friendships:

“So it was like I’ll be able to find out more information and I’ll be able to hopefully make a few friends, and hopefully then be able to ask things and maybe even feel able to help as well.” (Nicole).

*The freedom of social referencing.* Having attended the PMP, participants spoke about finally having met people who understood their experience of pain. This translated into a motivation to seek out further empathic understanding at the CPSG, and the mental freedom, respite and psychological bolstering that this offers. The struggle to explain the atypical freedom the CPSG offers from being judged ‘different’, led participants to perceive the CPSG as a radical change in culture, moving from isolation to integration:

"I dunno, the only way you could describe it really is being a foreigner in a foreign country and suddenly finding somebody who spoke your own language and you could relate.” (Ron).

Being surrounded by people who also suffered with pain also facilitated downward social comparison, which elevated the self-esteem of group members. Seeing others suffering helped participants reconceptualise their health status more positively, reviving new depths of coping
and interpersonal learning. For many participants, this social referencing was activated through self-talk, seeking to encourage the self to re-evaluate and re-orientate:

“...I looked at some of my other friends in the group and I’m sitting there thinking I’m slightly better and a few of the members are genuinely really, really sick. And I said to myself ‘ah get your backside off your chair’.” (Alex).

**Conjecture and the imminent choice**

The second superordinate theme, voiced by eleven participants, represented internal conflict surrounding making the initial decision to attend. Such conflict was the result of balancing personal anxieties and self-esteem with the need to make a rational and logical decision to attend, either alone or with colleagues from the PMP.

*Now or never*. On completing the PMP, participants discussed facing the reality that they needed to make an active choice to continue to utilise the techniques and clinical recommendations they had received, and that if time passed, the impetus for change would diminish. Consequently the sense of urgency was palpable for six participants who therefore made a rapid and immediate choice, rationalised as an essential decision in the context of the post-PMP need for dynamic action without delay. Ron argued the decision was pre-weighted towards acceptance of CPSG membership:

“If the PMP hadn’t been quite so successful, I might have felt ‘what’s the point because nobody can help me’... it just opened my eyes so much to the fact that there was still life there so I thought ‘go for it’.” (Jasmine).

Such rapid decision-making clearly demonstrated that half of participants felt the offer of CPSG membership forced a choice on the approach-avoidance coping continuum:

“So it’s a case of I won’t know until I go. And if I don’t like it, I won’t go. I’ve got to give it a try [...] I didn’t flinch, I thought ‘no, I’m gonna go’.” (Neil).

*Dancing around attendance*. The remaining half of participants discussed wrestling with the decision to join the CPSG and their ambivalence about this choice. Some participants expressed
decision-making difficulties related to readiness to change and fledgling acceptance of their pain condition. The rationale for the difficulty of the decision was typically the disruptive nature of chronic pain and the emotional impact of living with long-term pain. Pain caused internal conflict as participants struggled to overcome their pain-related desire to disengage and instead assert authority over pain:

"I was in two minds whether to go, but I ended up, I said 'och, bite the bullet, just go'. You're gonna sit here in pain, so you're as well sitting in the group in pain and chatting."

(Alex).

Not all participants found such self-talk possible, and many reported passive decision-making, choosing to attend only because colleagues from the PMP agreed to attend. However anger and disappointment was evident in all discussions of collective decision-making; frustration that promises to attend together were broken or that colleagues initially came but did not return:

"So we all [PMP group] decided we'd meet up there [the CPSG]. And that's where we all would have a reunion every month and that's how we'll keep together [...]. (...) It's only me turned up." (Ron).

Whether the initial decision or motivation to attend was made independently or collectively, five participants reported appreciating that attendance was never pressurised:

"No pressure. The last person someone needs that's in pain is pressure. If you're pressurised, you back away from it 100%." (Rebecca).

Evident in their discussions surrounding the lack of external pressure, was the increased propensity to attend; reducing pressure precipitated autonomy in decision-making. Without pressure, they were able to take ownership of their choice and continue to go with or without their colleagues, and the sense of an enhanced internal locus of control was apparent across participants.

_The confidence continuum._ All participants expressed surprise and satisfaction at their first visits to the CPSG, such positive experiences acting as validation of their choice to attend, imbuing
attendees with pride. This enhanced self-confidence and self-efficacy, and participants were encouraged by their own ability to succeed in new environments:

"And it were like a relief now because the relief that I’ve got there and I did it by myself y’know, so I were sorta proud of myself." (Alison)

Crises of confidence before attending were common, but in those participants who decided collectively with colleagues to attend, confidence levels were significantly higher, evident through their use of positive language. Motivation and momentum were enhanced through joint decision-making:

“So there was a gap of a few weeks I think where I sort of plucked up courage and like I said at first, I went with a couple of friends. They decided they wanted to go along and y’know we’d all go and see what it was like and from that it sort of gave me the incentive to go on my own the next time.” (Charlotte).

By contrast, those attending alone appeared much more tentative, hesitant and concerned over whether they had made the correct decision:

“The way Ron and this other lady were speaking, it was like it sounds really good. But I was also really hesitant because it would be another lot of strangers that I didn’t know.” (Nicole).

**Progressive and prospective pain management**

The third superordinate theme, representative of the views of all participants, addresses issues raised through a need for continuity in care and the transition from PMP to CPSG membership. There was a clear desire to extend the health provisions that participants had received, to build on the practical knowledge of pain management strategies and enhance quality of life in the future.

*The need for continuity.* Positive perceptions of the PMP were extremely important to all participants and strong drivers in the decision to attend the CPSG. The PMP primed CPSG
attendance on multiple levels; through HCP referrals, the need to continue with learned techniques and the ease of contact with healthcare services. Nine participants saw the CPSG as a simple extension of the PMP. Viewing it so enabled them to reduce worries about attendance as the decision was, to some extent, nullified. They attended because the PMP had finished and they were moving on to whatever came next:

"I'll be honest with you, I didn't want the six weeks [PMP] to end. I didn't, honestly, I loved going to the church and doing all the things we done. And so I went 'the monthly group's gonna hopefully be something the same'." (Alex).

For four participants, this decision was in order to maintain close contact with the clinical staff periodically attending the CPSG. They were motivated by the opportunity to access clinical expertise without a formal appointment, and saw this as an advantage that needed to be maximised and a method of preventing health-related decline. There was a clear sense of self-preservation, ensuring they were mentored and looked after, with informal support structures potentially providing assistance or mitigation in case of a health emergency:

"I thought I'll go. Just to keep in touch with pain. Because you don't always get an appointment from them and if they then decide they can't help you any more then you come away from them. You're lost in the crack. So I thought, 'no, I'll go to this pain group' and at least I'll get to see these people. And maybe not all the time, but I'll get to see them and if I do need help, maybe these people will know how to help me." (Laura).

Participants seemed keenly aware of their own fallibility in continuing to use what they had learned, therefore had a thirst for active accountability. They expressed their decision to attend as part of a damage-limitation measure, deliberately initiating an accountability strategy, which would ensure their continued self-management:

"But obviously sorta sometimes you sorta go back to your old ways, you need somebody to say 'get back and do that'. That's what I need, I need a kick" (Alison)

*Refilling the chronic pain toolkit.* Joining the CPSG, for ten participants, represented a technique by which they could self-manage their pain. Three participants explicitly re-used the language
that they had been taught on the PMP about the need for a pain toolkit, and the CPSG was effectively relocated into the toolkit as a strategy to enhance self-care:

“There are days when you try to go to your toolbox for a distraction and there’s none left in so you have try and find something else. And soon the toolbox gets empty and you’ve got to try and replenish it with other things that work.” (Neil).

Refilling the toolkit was also viewed at a deeper level; the CPSG offered new things to go into the toolkit through peer-to-peer learning and information-exchange. Participants respected that the group offered an opportunity to hear from others who had first-hand experience of comparable conditions, therefore their advice became invaluable:

“It’s like we can have a natter amongst ourselves; ‘have you gone and tried this, have you had a word with that, have you spoken to someone about that?’ Trust me, we’ve tried absolutely everything there is.” (Jane).

Through this, they could retain an internal locus of control, learning vicariously through others on an equal level of social comparison, in order to select what would work for their own pain.

_Living well with pain._ Joining the CPSG was a decision that was resolutely future-focused, facilitating renewed quality of life and enhanced coping strategies to continue living well with pain. The support group offered a place where pain would not control the sufferer, but where power could be reasserted, enabling participants to maintain a promotion-focus rather than (re)lapse into learned helplessness:

“It’s my way of making myself get up, get going and go out to this group. It’s like having a special room in the house that you go to get your mind back. That’s the only way I can put it ‘coz it’s so close to where I live. It’s ‘I am going to the pain group, I don’t care what’s going on’.” (Laura).

Participants were therefore determined in their desire to live well with pain and attendance was seen as a contract with the self in order to regain a sense of identity. By talking with others they reaffirmed that what they were doing to manage their pain was correct and they could learn to reengage with life positively:
"It wasn’t so much that I was learning how to cope, it was the fact that I was having what I was doing confirmed as being the right thing”. (Ron).

In essence, the decision to join the CPSG allowed participants to live well with pain, in spite of pain. Participants were able to avoid maladaptive coping through a strong network of interpersonal support, which functioned as a protective barrier, insulating them from further pain-related psychosocial decline.

Discussion

This study investigated the rationale surrounding the decision to attend a CPSG following PMP completion. Results demonstrated three superordinate themes: The thirst for comparative friendship; Conjecture and the imminent choice; and Progressive and prospective pain management. Collectively, these reflected the challenge of living with pain as an individual within a changing social world, and the desire to engage with social support.

The transition into CPSG membership was primed by positive experiences of the PMP and facilitated by peer presentations and self-care recommendations during the PMP. Fostering closer integration between clinical care and support groups therefore enables the PMP to function as a decisional aid, assisting with the decision to attend through peer-to-peer and clinician-initiated information exchange (Edwards & Elwyn, 2009). Clinician recommendation to join the CPSG enabled it to be (re)conceptualised positively, enhancing patient awareness of the benefits of social support (Grande, Arnott, Brundle, & Pilling, 2014). Such decisional-support embodies patient-centred care (NICE, 2012; Stacey et al., 2014) and promoted active decision-making, increasing internal locus of control and agency (Thoits, 2011; Uchino, 2009).

Prospective CPSG membership provoked two responses: an active choice to engage immediately, or a passive decision to attend as a consequence of a collective PMP peer-group trialling the CPSG together. Those who were less confident in their decision demonstrated choice-related internal conflict and lower self-efficacy. The split between confident attenders and those needing support may represent a continuum on the approach-avoidance coping scale (Roth & Cohen, 1986). Approach-coping in this population consisted of positive self-talk, asserting authority over pain, recognising one’s own need for social support as active accountability and
choosing to enable their continued use of PMP learned skills; all essentially positive coping strategies (Lazarus, 1993).

The proportional split between patients attending “now or never” and those “dancing around attendance” suggests that the role of the PMP may be in initiating desire for change and the CPSG in maintaining change. Those who made an active choice to engage with the CPSG demonstrated greater readiness for change, prospectively reflecting a move between the ‘action’ and ‘maintenance’ stages in the Transtheoretical Model (TTM) of health behaviour change (Prochaska & Velicer, 1997), with those collectively attending displaying earlier stages of change (e.g. contemplation). In the latter weeks of a PMP, clinicians could deliberately initiate discussion about propensity to attend a CPSG and if encountering passivity or reticence, could encourage a collective peer-group trial, drawing out debate over the PMP vs. CPSG roles in initiating or maintaining learned skills.

Prospective friendships were a strong attraction: associating with ‘pain friends’ engendered higher levels of empathic understanding and facilitated social integration (Embuldeniya et al., 2013). This was an outcome of downward social-referencing through Social Comparison Theory; seeing others live with comparable or worse pain enhances self-efficacy and self-acceptance (Davison et al., 2000; Festinger, 1954). Current findings model those of Embuldeniya et al. (2013) who found peer support reduces social isolation, encourages shared experiential learning and fosters psychosocial well-being. When advertising a CPSG, the friendship agenda should be emphasised as it is an attractive prospect in the context of pain-related reductions in available social support (Cowan, 2013).

Joining a CPSG was a function of a desire for continued investment in self-care, maintaining informal contact with support structures and developing a repertoire of self-management approaches. This enabled a promotion-focus, pursuing a positive future, as participants felt the CPSG ‘fit’ with their goals and future plans and desire for group attendance was intensified in accordance with Regulatory Fit Theory (Cesario, Higgins, & Scholer, 2008; Higgins, 2000). The group offered an opportunity to consolidate pain management skills, minimising risk of their extinction, in congruence with participants’ desires to live well with pain (Turk et al., 1993). Consequently, enhancing patient perception of the CPSG as an opportunity for
continuity of care and self-management has the potential to load the decisional balance positively in favour of CPSG membership.

This research offers an insight into the rationale for joining a CPSG, yet it is recognised that the participants in this study represent a self-selected sample, retrospectively recalling their decision-making processes. The sample is effectively a feature of the nature of a support group itself: those who invest in a support group and engage in mutual-support are those who have chosen to do so (Embuldeniya et al., 2013). Consequently, working with this population offers an insight into the perspectives and decision-making processes of those who would choose to engage in social support. Future research could aim to consider the perspectives of those who have chosen not to join a CPSG. Due to the nature of the interpretative lens, these analyses are not the only possible interpretation of data (Smith et al., 2009), therefore they are intended as a representation of the dynamic, evolving world expressed by participants at the time of study (Osborn & Smith, 1998).

The decision to attend a support group is a significant issue in light of expected costs and benefits of engaging. The decisional balance is positively weighted by the opportunity for continued pain management as an individual surrounded by strong social support structures. CPSG appeal is multifactorial, but is principally influenced through the offer of friendship, representing a collective point of social reference and an opportunity to act on the advice of healthcare professionals. Deciding to attend is enabled through active decision-making processes and can be facilitated through a collective choice to attend, primed during the PMP. Social support is an invaluable resource which promotes healthy outcomes and continued self-care, therefore offering an opportunity for both patients and clinicians alike.
References


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http://doi.org/10.1037/a0035623


Vangeli, E., & West, R. (2012). Transition towards a “non-smoker” identity following smoking cessation: An interpretative phenomenological analysis: Transition towards a "non-


<table>
<thead>
<tr>
<th>Participant</th>
<th>Gender</th>
<th>Age (years)</th>
<th>Cause of Pain</th>
<th>Pain Location(s)</th>
<th>Yrs with CP</th>
<th>Yrs since PMP</th>
<th>SG Attendance</th>
<th>MPS</th>
<th>MPI</th>
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<tr>
<td>Alison</td>
<td>Female</td>
<td>50</td>
<td>OA</td>
<td>Spine, neck, knees, shoulders, hands</td>
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<td>9</td>
<td>6</td>
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<tr>
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<td>68</td>
<td>Accident</td>
<td>Lower back, legs, neck</td>
<td>13 years</td>
<td>9 years</td>
<td>11 times</td>
<td>7</td>
<td>7</td>
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<tr>
<td>Charlotte</td>
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<td>OA, FM</td>
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<td>6 years</td>
<td>10 times</td>
<td>5</td>
<td>5</td>
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<tr>
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<td>Lower back, legs, neck</td>
<td>3 years</td>
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<td>12 times</td>
<td>8</td>
<td>6</td>
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<tr>
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<td>56</td>
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<td>Lower back, neck, shoulders, hands</td>
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<td>8 times</td>
<td>4</td>
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<tr>
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<td>Female</td>
<td>45</td>
<td>Accident</td>
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<td>6 times</td>
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<tr>
<td>Louise</td>
<td>Female</td>
<td>45</td>
<td>Accident</td>
<td>Lower back, neck, shoulders</td>
<td>2 years</td>
<td>10 years</td>
<td>9 times</td>
<td>7</td>
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<tr>
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<tr>
<td>Neil</td>
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<td>9</td>
<td>6</td>
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<tr>
<td>Ron</td>
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</tbody>
</table>

Notes: Pain-related variables are scored using the Brief Pain Inventory (Cleeland, 1992) from 0-10. MPS = Mean Pain Severity (max. 10); MPI = Mean Pain Interference (max. 10); Al = Mean Attendance (max. 12). All names are pseudonyms.
Table 2

Interview Schedule

1. Can you tell me about how you came to experience chronic pain?
2. Remembering back to when you joined the CPSG, can you tell me about that time?
3. How did you decide to join the CPSG?
4. Can you tell me about what it’s like to be a member of a CPSG?
5. What does the CPSG mean to you?
Table 3

*Overview of thematic results*

<table>
<thead>
<tr>
<th>Superordinate Themes</th>
<th>Subordinate Themes</th>
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<td>The Thirst for Comparative Friendship</td>
<td>Catching the Lifeline of Continued Support</td>
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<td>The Friendship Agenda</td>
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<tr>
<td></td>
<td>The Freedom of Social Referencing</td>
</tr>
<tr>
<td>Conjecture and the Imminent Choice</td>
<td>Now or Never</td>
</tr>
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<td></td>
<td>Dancing Around Attendance</td>
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<td>The Confidence Continuum</td>
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<td>Facing Labels and Pre-conceptions</td>
</tr>
<tr>
<td>Progressive and Prospective Pain Management</td>
<td>The Need for Continuity</td>
</tr>
<tr>
<td></td>
<td>Refilling the Chronic Pain Toolkit</td>
</tr>
<tr>
<td></td>
<td>Living Well with Pain</td>
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</tbody>
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