A qualitative analysis of patient-identified adaptive behaviour changes following interdisciplinary Acceptance and Commitment Therapy for chronic pain

M. Thompson\textsuperscript{1}, K.E. Vowles\textsuperscript{2,3}, G. Sowden\textsuperscript{3}, J. Ashworth\textsuperscript{3}, J. Levell\textsuperscript{3}

\textsuperscript{1} Faculty of Health and Applied Sciences, University of the West of England, Bristol, UK
\textsuperscript{2} Department of Psychology, University of New Mexico, Albuquerque, USA
\textsuperscript{3} IMPACT Pain Service, Staffordshire and Stoke-on-Trent NHS Partnership Trust, Stoke-on-Trent, UK

Abstract

**Background:** Interdisciplinary treatment programmes for chronic pain have strong evidence of treatment effect both immediately after treatment and at follow-up. However, despite strong outcome evidence, it is less clear which specific changes in behaviour are most relevant to patients or to outcomes. Indeed, it is not unknown for clinicians and patients to have different views with regard to goals of treatment. This study sought to evaluate the patients’ perspective regarding important behavioural changes that occurred while they were enrolled in a 4-week interdisciplinary programme of Acceptance and Commitment Therapy (ACT) for chronic pain.

**Methods:** Qualitative data were collected during a treatment session towards the end of treatment. In total, 104 completers from 16 consecutive treatment groups contributed to a data set consisting of 315 unique qualitative comments.

**Results:** Thematic analysis resulted in a theme hierarchy including overarching themes, midlevel themes and subthemes. Three overarching themes were identified as follows: (1) interacting with self – describing an interplay between various aspects of the individual, (2) activity – concerning how individuals practically and sustainably undertook activities and (3) interacting with others – exploring relationships with other people. The results section further describes the midlevel and subthemes that cluster under the overarching themes.

**Conclusions:** These data provide initial insights into the patient’s perspective of adaptive behavioural changes gained as part of an interdisciplinary programme of chronic pain rehabilitation. Overall, the data suggest the importance of a mix of both ACT-specific and more universal coping/pain rehabilitation elements. Future research may examine how these processes relate more directly to treatment outcome.

**Significance:** This study provides new qualitative insights into the patient’s perspective of adaptive behavioural changes gained as part of interdisciplinary pain rehabilitation. This and future work may help provide a more detailed understanding of the processes and behaviours that result in successful rehabilitation outcomes.
1. Introduction

Interdisciplinary treatment programmes for chronic pain have strong evidence of treatment effect (Fordyce, 1976; Turk and Burwinkle, 2005; Gatchel and Okifuji, 2006; Gatchel et al., 2014; Main et al., 2015). These interdisciplinary programmes typically include a mix of professional disciplines (e.g. psychology, physical and occupational therapy, physicians) and offer an intensive and time-limited dose of treatment, typically several hours per day, often on multiple days each week for several weeks (British Pain Society, 2013; Gatchel et al., 2014). Efficacy and effectiveness evidence suggest treatment completers reliably experience significant reductions in distress and disability, successful return to work, and renewed engagement in meaningful activities of daily living (Flor et al., 1992; Turk, 2002; Turk and Burwinkle, 2005; Gatchel and Okifuji, 2006; Gatchel et al., 2014; Kamper et al., 2014, 2015). Long-term data further indicate good maintenance of treatment gains through follow-ups of 3 to 7 years (Jensen et al., 2005; Hatten et al., 2006; Bergström et al., 2010; Vowles et al., 2011).

While the outcome evidence in support of interdisciplinary rehabilitation for chronic pain is robust, it is not yet clear which specific changes in behaviour are most relevant to that outcome. In other words, while outcome evidence is strong, the process evidence suggesting how these outcomes are achieved is weaker. Similar messages can be found in the 2012 Cochrane review of evidence for adults with pain and disability, which called for less general randomized controlled trials and instead for different types of studies to help understand what components work, for which patient, on which outcome and why (Williams et al., 2012). There are numerous ways of pursuing a more process-orientated agenda, one is to try and assess the perspective of patients themselves. After all, it is not uncommon for clinicians and patients to have different views with regard to the goals of treatment (Thorne and Morley, 2009; O’Brien et al., 2010). Thus, it may be useful to assess and evaluate important behavioural changes from the patient perspective. It may be that mixed method analytic approaches, which include both qualitative and quantitative evaluation of patient data, may allow for more informed and nuanced insights into the clinical material that patients find most compelling or useful. These methods may, in time, allow researchers to form a more informed understanding of the specific processes and behaviours to target for more successful outcomes within chronic pain rehabilitation programmes.

The present analyses sought to perform such an analysis using data collected during a treatment session that took place at the end of an 8-day interdisciplinary programme of rehabilitation for chronic pain. The purpose of this session was to collect and share patient perspectives on adaptive behavioural changes that they felt had been made over the course of treatment. The data from patient responses allowed for a qualitative analysis aided by descriptive statistics to help elucidate the patient-identified adaptive behaviour changes that had taken place over the course of treatment.

2. Method

2.1 Participants

Participants included 104 completers (65.4% female) of an interdisciplinary programme of chronic pain rehabilitation located in the Midlands of the United Kingdom. Average age was 48.0 years ($SD = 11.8$), and the sample was predominately White European in ethnicity (96.1%). Most patients were married or cohabitating (64.4%; single: 19.3%; divorced: 11.5%; widowed: 4.8%) and unemployed at the time of treatment onset (65.0% not working; 21.2% working full-time or part-time; 3.8% working as a ‘homemaker’; 11.5% no employment status identified). Modal pain duration was 6.7 years (range: 0.5–27.6 years). The most frequently identified primary site of pain was low back: 49.0%, followed by full body: 15.4%, neck: 11.5%, lower limb: 7.6%, shoulder/upper limb: 5.7%, midback: 4.9%, abdominal: 4.8%, and head: 1.1%. Most individuals, 60.6%, reported at least one secondary pain site.

Treatment was provided in a group format with the data of 16 consecutive groups included in the present analyses. Average group size was 6.9 ($SD = 1.5$) individuals. Previously published quantitative work has documented the effectiveness of the treatment programme at the end of treatment and at a 3-month follow-up appointment (Vowles et al., 2014, 2017). The material detailed in the present report has not previously been published.

2.2 Treatment details

The interdisciplinary pain rehabilitation programme included 2 days of treatment each week over a 4-week period. Each treatment day included approximately 5.5 h of active intervention provided by an
interdisciplinary team. Specifically, each day included 1 h of physical conditioning training and exposure to avoided movements and 1 h of physical activity (e.g. mindful movement, Tai Chi, exposure to novel activities; activity planning). In addition, each day included 1.5 h of psychological content, which included 30 min of mindfulness training, 1 h of activity management (e.g. values clarification, goal setting) and 1 h of educational or skills training content (e.g. pain physiology, effective communication, relapse preparation and prevention). The treatment team consisted of two physiotherapists, one clinical psychologist, one nurse and one physician.

Treatment was based on the therapeutic model of Acceptance and Commitment Therapy (ACT), an intervention with ‘strong’ empirical support, as graded by the American Psychological Association’s Society of Clinical Psychology (Society of Clinical Psychology, 2011). The overall focus of ACT for chronic pain involves increasing adaptive responses to pain, particularly in patient-identified values, defined as domains of activity that bring meaning, importance and vitality to the patient’s life. To date, evidence suggests that ACT for chronic pain is associated with significant and sustained reductions in disability, improvements in overall emotional and physical functioning, greater engagement with valued activities and fewer ineffective struggles for pain control (Vowles and Thompson, 2011; McCracken and Vowles, 2014; Veehof et al., 2016).

2.3 Data collection procedure

Data were collected during the final treatment hour on the seventh day of the 8-day treatment programme. The purpose of the session was to operationally define patient-identified adaptive behaviour changes that had occurred over the course of treatment. Colloquially, the treatment team referred to this session as the ‘survival guide’ session in that its purpose was to derive a discrete and specific list of adaptive behaviour changes that would aid patients in ‘surviving’ effectively with chronic pain once treatment had ended.

The session was led by a clinical psychologist (KEV) and began by asking patients to identify actions taken since the beginning of treatment that had contributed to improvement or had been helpful in achieving improved values-based action. The session leader then wrote down the actions identified by patients on a visual display (e.g. whiteboard, flip-chart). A primary task of the session leader was to aid patients in defining discrete behavioural actions that had been taken, as opposed to outcomes that had been achieved by these actions. For example, if a patient noted that he or she was ‘doing better’, then the patient was asked to clarify what had been done to occasion that improvement (e.g. ‘spend more time with family’, ‘ask for help when I need it’, ‘practice mindfulness’). Over the course of the 60-min session, a list was generated by the group. The psychologist periodically asked patients whether the list was ‘complete’, in the sense that engagement in the identified actions would aid in getting functioning back on track in relation to improved values in the event of a setback or that continued engagement in these actions would keep engagement in values at an adequate level for that patient. This question was used to aid in the generation of a more complete list and to aid in reinforcing a primary message in treatment, which was that treatment success involves effective responding to pain such that it does not needlessly limit one’s quality of life. At the end of the session, all identified actions were typed out and distributed to patients so that the survival guide could be taken home by patients and used to help guide future behaviour.

2.4 Transcription, overview of data set and data analysis

As described above, feedback from patient groups was transcribed by the session leader during the session. The comments generated by all the groups formed the data analysed in this study. In total, the 16 treatment groups provided 315 unique comments. The number of comments collected from each group ranged in total from 13 to 35 ($M = 19.7, SD = 6.7$). The number of characters in each comment ranged from 4 to 141 ($M = 36.0, SD = 20.6$). The average comment was about as long as: ‘Worry less about what others think’.

Given the nature of the data set above, it was decided to analyse the comments using a mixed methods approach, that is both qualitative data analysis and simple, descriptive statistics. More specifically, thematic analysis (TA) was used to analyse the data. This method is regarded as a robust analytical technique that has flexibility in terms of approach, theory and epistemology (see Braun and Clarke, 2006; Clarke and Braun, 2013). Other, well-regarded qualitative methods were also considered, but ultimately not used as it was felt that these methods were better suited to different contexts. For example, Interpretative Phenomenological Analysis (IPA; Smith and Osborn, 2008) tends to collect data in
greater detail, from a smaller number of participants, often through semi-structured interviews. In addition, a Grounded Theory approach was considered but also not used as some argue that the end results need to involve the production of a ‘plausible/useful theory’ (see Braun and Clarke, 2006, p. 81) – which this research did not set out to produce. Because this data set consisted of hundreds of single, pre-existing comments, provided by individual patients within the context of a group treatment, it was felt that TA was the most appropriate method to use.

According to Braun and Clarke (2006), TA involves ‘searching across a data set... to find repeated patterns of meaning (p. 86)’. Braun and Clarke describe six main phases to this process: transcribing the data set, familiarizing oneself with the data set, initial coding, searching for themes, reviewing and refining themes, and reporting the analysis. More specifically, in line with these guidelines, during the initial coding of the survival guide data, a representative specific theme was fitted to each comment, from each group. Each specific theme represented a short, subjective, descriptive label that aimed to summarize each comment. Generally speaking, only one code was allocated per comment. However, in certain instances, more than one code was needed, for example, when compound comments were analysed.

As coding progressed, themes were identified, reviewed and refined. Refinement of the themes continued until the structure was perceived to adequately capture the observed data. Notes were used as a way of recording and exploring the process of allocating themes, helping to document the analytic process and achieve consensus across the two coders. The initial coding of the data set was conducted by one coder (MT). The other coder (KEV) initially provided feedback and suggestions, and then independently rated comments to provide an index of inter-rater reliability with regard to the identified themes. The initial inter-rater reliability (Kappa) for all specific theme groups was 0.74, indicating good initial agreement. The analysis resulted in a three-level theme hierarchy: (1) overarching theme (the broadest category), (2) midlevel theme and (3) subtheme (the finest level of theme detail). In some areas, all three theme levels were needed to adequately capture the data; in others, two levels sufficed.

Finally, in the spirit of reflexivity (see Braun and Clarke, 2013, p. 303), it is worth being explicit about the context, standpoints and perspective of the authors on this study. As already noted, the interdisciplinary treatment programme the data were collected from follows an ACT approach. Moreover, many of the authors of this study have worked and published within the ACT domain for many years. Potential limitations of this perspective are more fully expanded in this discussion.

3. Results

For each of the overarching, midlevel and subthemes, representative quotes are provided to illustrate specific themes. In addition, simple, descriptive quantitative information is provided to order the themes in terms of decreasing prevalence across groups. We elected to provide more detail on themes that occurred in at least nine of the groups (>50% of groups). Themes that occur with less frequency are described in less detail.

In descending order of frequency, the three overarching themes that emerged were as follows: (1) Interacting with self (140 comments; 44% of total), (2) Activity (94 comments; 30%) and (3) Interacting with others (69 comments, 22%). Of the total data set, a small number of comments (n = 13) could not be clearly allocated as they fitted with multiple themes (4 comments; 1%) or did not fit well within any of the identified overarching themes (9 comments, 3%). These 13 comments were not explored further.

3.1 Overarching theme 1: Interacting with self

The 140 comments within this theme highlighted the interplay between various aspects of the individual. Each of the 16 groups had at least one comment related to this overarching theme. Four midlevel themes are described below and shown in Fig. 1.

3.1.1 Midlevel theme 1.1: Relationship with private events

The most prevalent midlevel theme in this area was Relationship with private events (67 comments, 16 groups). The term ‘private events’ refers to an individual’s thoughts, feelings, memories, body sensations and urges and is derived from the radical behavioural approach that underlies ACT. It refers to behavioural events that are only observable to the individual who is behaving (Skinner, 1953; Hayes and Wilson, 1995; Moore, 2007). As described elsewhere, ACT does not seek to alter the form or frequency that private events take, but instead seeks to alter the ways that these private events influence subsequent action. Every group made at least one comment related to this area. Fig. 1 outlines the four...
subthemes within this midlevel theme, as well as the frequency details.

The first subtheme, labelled Flexibility/Willingness (34 comments, 15 groups), related to both flexibility in responding to and willingness to experience difficult private events. Some comments encapsulated this idea in a very succinct way: ‘Be flexible’ and ‘Struggle less’. Other longer comments described it as follows: ‘Control the things you can control, leave the things you cannot control alone’ and ‘See if you can be fully accepting and willing to be who you are right now’. Some comments referred specifically to metaphors and experiential tasks used within the treatment programme to help patients respond more flexibly and adaptively to private events. For example, some common ACT exercises ask patients to identify the ways that private events are unhelpfully influencing behaviour and to practice responding to them differently to maximize valued living. Reference to these exercises was made in several comments, such as ‘Spot the passengers and notice them pushing you around’ and ‘Don’t let the passengers drive your life’. This refers to the passengers on the bus exercise where ‘passengers’ are thoughts and other private events on a metaphorical bus being driven through one’s life (see Hayes et al., 2011a, p. 250).

The second subtheme, Paying attention/Being aware (20 comments, 13 groups), included comments highlighting the importance of being able to monitor experience occurring in the present moment. As noted, each treatment day included 30 min of mindfulness exercises, which were designed specifically to augment present-focused awareness and effective responding to present experiences. The phrase ‘Be mindful’ appeared in seven of the 20 comments in this group. Other comments did not directly refer to mindfulness but referred to other aspects of awareness and directed attention, for example ‘Be aware of your choices’ and ‘Be in the here & now, not the there & then’.

The third subtheme was labelled, ‘Jump’/Take calculated risks (11 comments, 10 groups). This label highlights the notion of taking calculated risks to pursue values. ‘Jump’ refers to an experiential exercise targeting willingness from the ACT tradition,
related to the Zen saying: ‘You cannot jump a canyon in two steps’ (see Hayes et al., 1999, p. 240). The exercise suggests that while the ‘quantity’ or height of any jump can be modified, the essential ‘quality’ of jumping – that is putting yourself out into space and letting gravity do the rest (Hayes et al., 1999, p. 241) remains vital and unchanged whatever the height. Comments ranged from the succinct ‘Jump’ to the longer ‘Take a chance when it matters (Jump!)’.

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They also included other comments which appeared to capture the same sentiment without using the term jump: ‘Take risks in a controlled, sensible way’. More specifically, of the 11 comments, eight used the term ‘Jump’, while another three mentioned taking risks. This sense of jumping/taking risks is another illustration of being flexible and willing (see earlier) in the pursuit of values-based activities; however, the clustering of comments around these two terms seemed to warrant its own specific theme.

One additional subtheme was present but was recorded in only two groups. It was labelled Responding to barriers (two groups, two comments). One of these two comments, for example, was, ‘Stop using barriers as excuses’.

3.1.2 Midlevel theme 1.2: Relationship with self

Relationship with self was the second most popular midlevel theme under the Interacting with self overarching theme (37 comments, 26% of main theme total; 16 groups). Again, every group made comments related to this area. The comments at this midlevel related to the individual’s responses and reactions to him or herself. There were six subthemes in this area; however, only one occurred in the majority of groups.

The largest subtheme was Being gentle/Not beating one’s self up (17 comments, 14 groups), and the content was closely related to the theme’s title, for example ‘Be gentle with yourself’, ‘It is not necessary to beat yourself up’ and ‘Don’t beat yourself up when it doesn’t achieve anything’. In many ways, this subtheme shares some similarities to earlier subthemes that captured awareness of and flexibility with private events. However, in this particular subtheme, the focus was more on enhancing flexibility towards one’s self more generally. As such, it seemed to warrant its own specific theme.

The remaining five subthemes only occurred in a minority of groups. Give credit to yourself (seven comments, six groups), Laugh (five comments, five groups), Be honest with yourself (four comments, four groups) and Trust/Believe in yourself (two comments, two groups). Comments from these subthemes closely mirrored the theme titles.

3.1.3 Midlevel theme 1.3: Values selection and awareness

The theme, labelled Values selection and awareness, was composed of 25 comments across 11 groups, representing 18% of the comments within the Interacting with self overarching theme. No subthemes were necessary under this midlevel theme as all comments seemed to cohere sufficiently at the midlevel theme.

Within the ACT model, work around values highlights the things that are most important to an individual and seeks to ensure that they guide day-to-day behaviour (Hayes et al., 2011a). In ACT, a distinction is often made between values (a continuous direction of travel, e.g. to be a loving and caring partner) and goals (specific targets, big or small, in that direction of travel, e.g. to plan or attend a social event important to one’s partner). In the patient comments, a number of comments referred to both values and goals, for example ‘Keep values and goals in mind’ and ‘Be guided by your values and goals’. Other comments focused on these same things without specifically using the terms, for example values identification: ‘Have an aim’, ‘Identify what you want’, ‘Choose to do what you want’ and values pursuit: ‘Do things that are worth it’ and ‘Do more of what you want to do’.

3.1.4 Midlevel theme 1.4: Relationship with pain

The fourth midlevel theme was labelled Relationship with pain and consisted of comments describing an individual’s interactions with the pain experience. Again, no further subthemes were needed. Overall, there were 11 comments, across nine groups, representing 8% of the total comments within the Interacting with self theme. Importantly, in keeping with the application of the ACT model in this area (Hayes et al., 2011b; Vowles and Thompson, 2011; McCracken and Vowles, 2014), all comments encapsulated the idea of either struggling less with pain (e.g. ‘Don’t struggle needlessly with pain’ and ‘Be wary of engaging in a struggle with pain. It saps energy that may be best put somewhere else’) or engaging more with life, while also being willing to experience pain (e.g. ‘Live life with pain’, ‘You will have pain regardless, so live your life’).

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3.2 Overarching theme 2: Activity

Activity was the second most prevalent overarching theme (94 comments, 30% of total comments), with relevant comments recorded across all 16 treatment groups. Whereas the preceding midlevel theme of values selection and awareness was related to being aware of what was most important to the individual, the comments in this overarching theme were more concerned with how individuals practically and sustainably move towards their identified values. The two midlevel themes: aspects of activity and specific activity, and their subthemes are described below (Fig. 2).

3.2.1 Midlevel theme 2.1: Approach to activities

The data in this midlevel theme (77 comments, 16 groups) relate to important qualities of overt behaviours and wider approaches to activity. The comments in this theme relate to these qualities and approaches rather than specific activities themselves. Each group provided a comment under this midlevel theme. Three subthemes were identified, all of which occurred in the majority of groups, specifically doing flexibly/pacing; dealing with setbacks/failure; making plans.

The largest subtheme, labelled Doing flexibly/pacing (41 comments, 15 groups), was the most prevalent subtheme in the entire data set. It refers to the spirit and, in some respects, speed at which activity is undertaken. On the one hand, comments repeatedly highlighted the idea of breaking down bigger behavioural objectives into sequential steps. In fact, the word ‘steps’ was mentioned in many comments, for example ‘Start with easy steps’ and ‘Take things one step at a time’. A similar sentiment was also expressed around pacing and slowing down, for example ‘Pace, don’t be reckless in your behaviour’ and ‘Be careful of rushing or pushing through’. Equally, it was noted how it was ‘Ok to stop halfway through an activity and also how it could be good to ‘Experiment’ or, in slightly longer form: ‘Don’t rule anything out and keep your options open’. Ultimately, there was a sense in these comments that ‘Quality matters more than quantity’ and that doing just some of an activity slowly, but sustainably, was better in the long term than forcing oneself to hit a target in an unsustainable way.

The next subtheme, Dealing with setbacks/failure (19 comments, 11 groups), made it clear that patients understood that setbacks could happen, were acceptable, and could even represent an opportunity to learn. For example, comments about the expected nature of setbacks included ‘Expect setbacks’, ‘It is ok to have a bad day’ and ‘Fail (it is ok)’. Equally, other comments focused on the potential to learn.

Figure 2 Overarching, midlevel and subthemes in the ‘Activity’ theme hierarchy.
from setbacks: ‘Learn from your mistakes and learn from your experience’, ‘Learn from your mistakes and try again’, as well as the importance of re-engaging in behaviour: ‘After failing, make sure it matters and start again’.

The final subtheme, Make plans, (17 comments, 10 groups) encouraged individuals to make plans and set goals. The words ‘plan’, ‘goals’ or ‘routine’ regularly featured in specific comments, for example ‘Make and follow plans’, ‘Set realistic goals’, or ‘Establish a routine’.

3.2.2 Midlevel theme 2.2: Specific activities

Comments within the Specific activities midlevel theme focused on the need to actively engage in specific activities. This focus on specific activity was in contrast to the more general approaches to activity captured in the previous midlevel theme. A total of 12 groups provided 17 comments under this midlevel theme. Three specific subthemes were identified, but none of these occurred in the majority of groups.

The first subtheme, labelled Exercise (nine comments, eight groups), placed priority on physical activity such as ‘Exercise’ or ‘Get active/exercise/go to the gym’. The second subtheme referred to more Generally doing activities (five comments, five groups) and highlighted that one should concretely ‘Put things into practice’ and ‘Make a move towards what you want’. The final subtheme listed Specific other activities (three comments, three groups) that an individual could do such as ‘Get out of bed’ or ‘Get out of the house’.

3.3 Overarching theme 3: Interacting with others

The final overarching theme, Interacting with others, had the fewest comments of the overarching themes \((n = 69, 22\% \text{ of total comments})\). In total, 15 of 16 groups provided comments for this theme. The data in this theme shifted the focus from the relationship with one’s own private events (overarching theme 1) to relationships with others. There were two midlevel themes, expanded below.

3.3.1 Midlevel theme 3.1: Relationship with others

As shown in Fig. 3, The first midlevel theme, Relationship with others (36 comments, 15 groups), identified two subthemes: Socializing and Focusing on self. The first, Socializing (19 comments, 15 groups), encouraged individuals to be around others and to avoid social isolation. For example, ‘Socialize/Don’t isolate yourself’, ‘Don’t withdraw from others, socialize instead’ and ‘Get around people who are like-minded’. The second, Focusing on self (17 comments, 11 groups), encouraged people to do things for themselves. For example, ‘Take time for yourself’, ‘Do something yourself’ and ‘Do things by yourself’.

Also evident within this second specific theme was a focus on patients prioritizing themselves, for example by putting personal needs first. Examples included ‘Take care of your own needs’ and ‘Be selfish and put yourself first when it matters’.

3.3.2 Midlevel theme 3.2: Communication with others

The final midlevel theme, labelled Communication with others (33 comments, 15 groups), reflected aspects of interpersonal communication. Four subthemes were identified.

The largest subtheme was labelled Asking for help, which was the only one observed in the majority of groups with 14 groups providing a total of 20 comments. Comments focused upon individuals asking for what was wanted or needed. Along with variations of ‘Ask for help/Ask for what you want/Ask for what you need’, other comments included ‘Tell people what you want (they cannot read your mind)’, ‘Talk to those who care about you and ask for help if you need it’ and ‘Ask for and accept help when you need it’.

The three remaining specific themes, each of which occurred in a minority of groups only, included a focus on effectively Communicating feelings to others (six comments, six groups), Listening to others (four comments, four groups) and Saying no to requests when that was in one’s best interest (three comments, three groups).

4. Discussion

Over one hundred treatment completers, across 16 consecutive programmes, contributed data to a ‘survival guide’ session. The results provide some initial insights into the patient perspective of salient adaptive behavioural changes gained as part of an interdisciplinary programme of chronic pain rehabilitation. Three overarching themes were identified.

The first theme, ‘Interacting with self’, included details of how patients had learned to interact with internal private events and aspects of their own behaviour. For example, comments indicated increased willingness to experience aversive private
events, such as distress and pain; a broadening out of awareness beyond these experiences to become more aware of individual values, taking calculated risks in the direction of those values; as well as increases in self-compassion. This theme included a number of behavioural interactions consistent with the ACT model of treatment, which itself prioritizes these ways of responding to aversive experiences if it promotes valued living (McCracken, 2005; Hayes et al., 2011a). Previous work using self-report measures in chronic pain has also supported the importance of these response patterns in chronic pain, including acceptance of chronic pain (McCracken, 1998; McCracken et al., 2004; Wicksell et al., 2010), more general willingness and acceptance (McCracken and Zhao-O’Brien, 2010) and values clarity (McCracken and Yang, 2006). Thus, when the patient comments under this theme are considered in the light of what the model itself suggests are adaptive behaviours, a degree of concordance is apparent.

The second overarching theme (‘Activity’), which pertained to patient engagement in activity, included changes in overall approach to activity, such as planning and flexible goal pursuit, adjusting to setbacks, as well as specific activities that were deemed useful to engage in. Overall, the activities within this theme were less ACT-specific and perhaps more in line with what we have previously referred to as ‘traditional’ (McCracken and Vowles, 2007; Vowles and McCracken, 2010) or perhaps ‘more universal’ pain specific coping behaviours. In other words, both ACT and non-ACT interdisciplinary pain rehabilitation place priority on improving strategic engagement in activity, facilitating judicious planning and pacing of activities, and decreasing sedentary or isolating behaviours (Fordyce, 1976; Turk et al., 1983; Jensen et al., 1991; Gatchel et al., 2007; Main et al., 2015).

The final overarching theme described changes in social and interpersonal activity (‘Interacting with others’). Patients noted the importance of improved social functioning, while also prioritizing their own needs. They also noted the importance of clarity in communication. As with the preceding theme, the facilitation of effective communication is often a part of interdisciplinary pain treatments, both ACT and non-ACT, thus its representation in patient comments is unsurprising.

Overall, these data appear to suggest that adaptive patient changes over the course of an interdisciplinary course of ACT for chronic pain were viewed by patients themselves as a mix of both ACT-specific and more universal coping/pain rehabilitation elements. This finding is broadly supported by some of

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**Figure 3** Overarching, midlevel and subthemes in the ‘Interacting with others’ theme hierarchy.
our previous quantitative work which has indicated that both of these two ‘styles’ of responding to pain increased over the course of a similar, but more intensive, ACT-based interdisciplinary treatment (Vowles and McCracken, 2010).

We conceive the separate themes presented above as being related to each other and not completely distinct. This is perhaps illustrated by the values selection and awareness midlevel theme (of the *interacting with self* overarching theme) and the *activity* overarching theme. While it is acknowledged that the two exist in different theme hierarchies, it is also the case that what an individual does (activity) is hopefully influenced by what they personally value (values section and awareness). In this way, themes can be and are related, even when existing in different hierarchies.

Table 1 lists the most numerous themes in the data set. It is interesting that relationship with pain is the last entry in the table and was only explicitly mentioned in nine of 16 groups. Although this may, at first, seem surprising, there are a number of ways of looking at this. Firstly, a lot of content concerning an individual’s changing relationship with pain is included under other more common themes from Table 1 such as flexibility/willingness and paying attention/being aware. In other words, when participants in ACT interdisciplinary treatment increase flexibility with or awareness of private events, this includes thoughts and feelings, and it also explicitly includes body sensations such as pain. In addition, the focus of an ACT interdisciplinary treatment is not necessarily on reducing pain intensity or other symptoms but on increasing daily functioning in the direction of values and goals even when pain reduction is not possible. Indeed, a recent study has even suggested that shifts in pain acceptance (not pain intensity or reduction) may be responsible for successful outcome in a CBT interdisciplinary treatment programme – even when pain acceptance was not explicitly targeted during treatment (see Akerblom et al., 2015). Another recent study has indicated that improvements in pain intensity during treatment do not appear to be a prerequisite for improvements in physical and emotional functioning following interdisciplinary ACT (Vowles et al., 2017).

As qualitative methodology is not used with the same frequency as quantitative methodology in the chronic pain literature, it is worth placing this study in the context of other qualitative pain research. In their review of the qualitative literature, Osborn and Rodham (2010) note that previous chronic pain research has tended to focus on both the ‘lived experience of pain’ and the personal experience of ‘seeking help for pain’. The same review notes a possible discrepancy between qualitative and quantitative chronic pain research. Specifically, that the qualitative literature tends focus less on areas often found in the quantitative literature (i.e. ‘acceptance, fear avoidance and catastrophizing’ [p. 5]). It is noteworthy that the current research, using a qualitative methodology but focused on the end of treatment, primarily resulted in similar themes to those found in the qualitative literature. It seems possible that part of the reason for any discrepancy between qualitative and quantitative chronic pain research. Specifically, that the qualitative literature tends focus less on areas often found in the quantitative literature.

### Table 1 Data on the most prevalent specific themes.

| Theme title                        | Overarching theme | No. of groups | No. of comments |
|------------------------------------|-------------------|---------------|-----------------|
| Doing flexibly/Pacing              | Activity          | 41            | 16              |
| Flexibility/Willingness            | Self              | 34            | 15              |
| Values selection and awareness     | Self              | 25            | 11              |
| Asking for help                    | Others            | 20            | 14              |
| Paying attention/Being aware       | Self              | 20            | 13              |
| Socializing                        | Others            | 19            | 15              |
| Dealing with setbacks/Failure      | Activity          | 19            | 11              |
| Being gentle/Not beating self up   | Self              | 17            | 13              |
| Focusing on self                   | Others            | 17            | 11              |
| ‘Jump’/Take calculated risks       | Self              | 11            | 10              |
| Relationship with pain             | Self              | 11            | 9               |

Italicized rows contain patterns of data found in less than nine groups.

Self = Interacting with self, Others = Interacting with others.

### 4.1 Outcome and process research

As highlighted briefly in the introduction, treatment research can be divided into that which focuses on outcome and that which focuses on process (see Kazdin, 2007). Outcome research concentrates primarily on whether a treatment works or how well it works in comparison with another treatment. Conversely, process research, which is often found within psychotherapy and counselling areas, is more concerned with how a treatment works and what
aspects of a treatment contribute to successful outcome. The work reported here examines treatment process, as it seeks to understand what aspects of treatment patients felt contributed to their behavioural changes. That said, the study also seeks to explore the possibility of linking future process findings to outcome data (see future research, below).

The process literature itself can arguably be divided into two camps: (1) that related to the importance of aspects of patient/provider relationship (also known as therapeutic alliance) and (2) that related to the importance of specific techniques within a given approach (see Castonguay, 1993; Wampold, 2015). A number of researchers interested in process research argue for the primacy of the patient/provider relationship (see Norcross and Lambert, 2011; Norcross and Wampold, 2011). Regular, peer-reviewed, meta-analyses have examined the influence of this relationship on psychotherapeutic outcome and have found a remarkably consistent but relatively small correlation (between 0.22 and 0.28), accounting for about 5% and 8% of the variance (see Vowles and Thompson, 2012). It is noteworthy that such research tends to focus on the working together of an individual therapist and an individual client. Of course, in the chronic pain treatment field, treatment often involves larger groups of patients, who regularly see several different health care professionals concurrently as part of their interdisciplinary treatment.

With this in mind, it is worth noting two things. Firstly, that the influence of the various inter- and intrapatient/provider relationships in group-based pain management programmes might be both significant and complex (see, e.g. Lewis et al., 2010; Williams and Potts, 2010). Secondly, as this specific research only drew upon material from the survival guide session (see methods), such data are only likely to pick up on material related to specific techniques within an approach. It is not likely to highlight material related to aspects of the patient/provider relationship. This of course, does not mean that such relational material was not important, only that these data were not likely to be collected as part of the survival guide session. In some ways, this must be considered a limitation of this research. Further limitations are highlighted below.

4.2 Limitations

As with much research of this nature, the final overarching, midlevel and subthemes were reached through analysis and discussions between a limited number of researchers. Despite reflective checks, it is possible that another team may have categorized the data differently. As noted above, it is important to note that the dividing lines between the final themes are likely in some way to be a reflection of the researchers and their interests, and a different set of researchers may have found other plausible arrangements of the data (see future research below).

The data that make up this data set come from inside a ‘real-life’ chronic pain treatment environment. Equally, similar to previous qualitative research of this type (e.g. Dunford et al., 2014), these data do not come from isolated individuals but from individuals who were part of a treatment group as a whole. As noted in the method, survival guide comments were suggested by patients in the group and discussed as a group, as they were being written up on the white board by the session leader. Naturally, there is a process of filtering and refinement that takes place here, and perhaps some level of influence from both the wider group and the session leader. This may limit the extent to which these comments can be seen to reflect the thoughts of any specific individual within any one group. However, the above notwithstanding, the data still seem to capture potentially interesting insights.

5. Future research and conclusion

In their review, Osborn and Rodham (2010) noted that researchers tended not to ‘accumulate a substantial body of [qualitative] work’ enabling them ‘to develop their arguments’ (p. 5) over time. With this in mind, it is worth considering what future research in this area may look like. It would be interesting if other interdisciplinary treatment programmes were to see the ‘survival guide’ session as a potentially useful clinical tool. If adopted, this could allow for future data collection and analysis that replicates the above work. It would be interesting to see if similar themes emerged from replicated research conducted within treatment programmes from different therapeutic orientations, for example traditional Cognitive Behavioural Therapy (CBT). It seems quite possible that many of the traditional coping behaviours from themes 2 and 3 would be similar given the shared lineage of both ACT and CBT – although, of course, further research would be required to evidence this.

In future work, it will be important to examine the extent to which the themes identified in this work fit with new data or whether revisions are needed. For example, one thing that may happen over time is that some themes that did not feature in
half or more of the groups, may become more (or less) prominent. Moving even more towards a mixed methods approach, it also seems possible that with enough data, future research could examine whether the differences in the existence and the extent of qualitative themes have any relation to the improvements in physical and emotional functioning that are associated with the treatment itself. Thus, more directly attempting to use this qualitative data to examine the patient reported processes that influence quantitative outcome.

In conclusion, this study provides a preliminary, mainly qualitative, exploration into the adaptive behavioural insights gained by patients during an interdisciplinary programme of chronic pain rehabilitation. Importantly, this study provides contemporaneous qualitative evidence of the active ingredients of therapeutic change from the patient point of view; an area that thus far appears to have been underserved by the chronic pain literature. The specific findings of this research highlight the perceived importance of both ACT-specific and more universal coping/pain rehabilitation elements within an ACT-focused interdisciplinary treatment programme. The connections between the findings in this study and findings previously reported in the quantitative assessment and treatment literature may go some way to both help explain and even potentially bridge a gap that has existed between the qualitative and quantitative chronic pain literatures. Importantly, the methodology adopted here may also be adopted by other services, using other frameworks to help continue to pursue process research in this area. Future research following these methods may further investigate the generalizability of these findings and how they translate across programmes of different orientations. Finally, extensions of this work may even have the potential to examine more closely how these potentially relevant processes relate more directly to treatment outcome: marrying the traditions of process and outcome research.

Author contributions
All persons identified as authors participated sufficiently in this research to warrant their authorship.

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