Exploring crowdsourced self-care techniques

A study on Parkinson's disease

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Exploring crowdsourced self-care techniques: A study on Parkinson’s disease

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A B S T R A C T
Living with Parkinson’s Disease introduces a range of significant challenges into one’s daily life. While medical interventions exist to overcome some of these challenges, patient self-care techniques often form an essential complement to the treatments recommended by medical doctors. Knowledge on these self-care techniques often originates from those living with Parkinson’s themselves or their close caregivers, as they have the knowledge and experience required to assess self-care techniques. This so-called ‘patient knowledge’ is usually exchanged in peer meetings or discussion forums. Although vital to the Parkinson’s Disease community, this information is often difficult to access due to its unstructured format and the difficulty of navigating through online forums. We present an online tool that allows for contributing, assessing, and finally discovering Parkinson’s Disease self-care techniques. The custom discovery tool was populated with self-care knowledge by over 300 people with Parkinson’s and dozens of their carers, spanning areas such as daily well-being and using assistive equipment. Then, we invited patients to explore the discover features in a smaller scale trial. While well-received, our deployment highlighted several challenges that we further discuss in this paper. Overall, our study contributes to crowdsourced digital health solutions and provides both design and research implications to this challenging domain with a vulnerable user group.

1. Introduction
Parkinson’s disease (PD) is a chronic neurological disorder caused by progressive loss of neurons in the central nervous system (Jankovic and Tolosa, 2007). That leads to decrease in dopamine levels, which manifests as a wide variety of both motor and non-motor symptoms (Jankovic and Tolosa, 2007). Motor symptoms, such as tremors, bradykinesia, rigidity, and postural instability, impact daily activities (Jankovic, 2008). For example, moving around, eating or getting dressed may be difficult (Hariz and Forsgren, 2011). Non-motor symptoms include e.g. autonomic dysfunction, cognitive impairment, and depression (Weintraub et al., 2008).

Often, PD affects negatively one’s quality of life (QoL). Andrejack and Mathur (2020) describe that QoL of those living with PD is influenced by three main elements. These are the availability and quality of professional health care, support from family, friends and peers, and the ability to engage in self-care, i.e. modifying one’s lifestyle to cope with the condition as well as actively seeking to mitigate any potential symptoms through various means, typically at home (Andrejack and Mathur, 2020).

The symptoms challenge daily life, and living with these challenges issues various forms of self-care (Nunes and Fitzpatrick, 2018). Further, the effectiveness of self-care techniques is subjective and can thus vary a lot between subjects. Typical sources for self-care information are online forums, peer group discussions, or practically any information media that allow people to exchange ideas and experiences (Attard and Coulson, 2012; Preece, 1999). To this end, the best experts on self-care are people living with a given condition: the patients and their direct personal caregivers and family members. These individuals possess valuable ‘patient expertise’, practices, and methods that outline how to best manage the condition. However, the information as collected in current online sources is often presented in an unorganized structure.
In the case of chronic diseases, the outcome of self-care is maintaining the quality of life and the perceived control of the disease (Riegel et al., 2012). It may also reduce mortality, hospitalization, and healthcare costs (Iovino et al., 2020). Self-care is often a collaborative endeavor between patients and carers (Nunes and Fitzpatrick, 2015), and for example, in the case of PD, most of the patients have an informal, dedicated caregiver (Schrag et al., 2006). Caregiver participation facilitates self-care (Iovino et al., 2020), and family members may often have different roles in self-care (Pina et al., 2017).

Each chronic condition has unique self-care techniques. Nunes and Fitzpatrick (2018) identified four main types of self-care activities in PD, namely: taking medication, exercise, adapting lifestyle, and accepting the disease’s consequences. Medication plays a crucial role in managing daily life for many people with PD. Participants had different strategies for maintaining their medication schedule, such as using alarms or using the placement of the medication as a reminder. Exercising was seen as another important aspect of self-care, enhancing movement control and wellbeing. Adapting lifestyle refers to adjustments due to a worsening condition, either permanently (e.g., lose driver’s license) or temporarily (e.g., cancel a visit to a friend) due to daily symptom fluctuations.

Pina et al. (2017) identified several roles in self-care (Pina et al., 2017). Caregiver participation facilitates self-care (Iovino et al., 2020), and family members may often have different roles in self-care. This highlights the daily life context related to self-care activities.

2. Related work

2.1. Self-care of a chronic condition

Self-care is defined e.g. by the UK Department of Health as "the activities that enable people to deal with the impact of a long term condition on their daily lives, dealing with the emotional changes, adherence to treatment regimes, and maintaining those things that are important to them – work, socialising, family", and improving self-care is seen as important in supporting people with chronic conditions (Department of Health, 2006). The Middle-Range Theory of Self-Care of Chronic Illness (Riegel et al., 2012) defines self-care as "a process of maintaining health through health-promoting practices and managing illness". The theory divides self-care into self-care maintenance, self-care monitoring, and self-care management. Self-care maintenance targets maintaining physical and emotional stability with the chronic illness, including actions such as lifestyle changes and adherence to the treatment, e.g. medication adherence (Riegel et al., 2012). Self-care monitoring focuses on recognizing changes to a condition, with monitoring body functions, feelings, and symptoms (Riegel et al., 2012). Self-care management is reacting to these results, determining if actions are needed. For example, the patient may decide whether to take extra medication when their symptoms worsen (Riegel et al., 2012).

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It is notable, however, that more than half of PD patients consider the various online information sources as very helpful in dealing with their medical situation (Stocchi and Bloem, 2013).

In a study relevant to our paper, Riggare et al. (2019) studied knowledge acquisition among PD patients in Sweden. Of the 346 persons with PD who responded to the online survey, 36% indicated to find disease-specific knowledge by themselves through the internet, 29% received information via patient organizations, 25% from healthcare professionals, and 5% from other patients. A total of 91% of surveyed patients felt that obtaining knowledge about PD is important for them, although just over half (55%) of respondents expressed that they had been able to obtain the information they require. A recent study among 190 PD patients by Marxreiter et al. (2020) found out that approximately 75% of the respondents use the internet for accessing disease-related information in Germany.

Fox and Brenner (2012) studied internet use by caregivers of chronic conditions, they found that 79% of caregivers have access to the internet and that the large majority of them (88%) search for health-related information on the internet. Caregivers are furthermore more likely to use both in-person and online social relationships for acquiring health-related information than age matched non-caregivers (Fox and Brenner, 2012).

These prior works highlight the value of online approaches in providing people living with PD with practical information on coping with PD on a daily basis. In this paper, we build on existing knowledge within the PD community to provide a structured way to discover relevant self-care techniques.

2.3. Peer-provided health data in online communities

A variety of eHealth solutions have been designed for exploring peer-generated content in different contexts, e.g. weight loss (Hosio et al., 2020), different chronic illness (Berkanish et al., 2022; Hos-sain et al., 2021), mental health (Fortuna et al., 2020) and hospital admission (Haldar et al., 2020). Online communities have emerged which help patients to increase their understanding of their treatment through sharing insights (Johnson and Ambrose, 2006). In such support groups, patients often share and find operationalizations of coping mechanisms, including instructions and routines, to better manage their condition (Huh and Ackerman, 2012). Charlton and Barrow (2002) studied self-help groups for PD patients and found that participation in such a support group may help both in accepting the disease as well as in finding ways to better cope with it. Other work shows that participation in an online PD support group can improve patient quality of life (Lieberman et al., 2005, 2006). Online forums may serve as an important source of disease-specific information in areas where medical personnel might have limited knowledge or experience with the condition (Zhou et al., 2014).

Hartzler and Pratt (2011) studied how patients’ expertise differs from clinicians’ expertise. Patient expertise is defined by personal experiential knowledge as obtained through daily management of one’s personal condition, whereas the source of clinician expertise is education and practical experiment. Hartzler and Pratt (2011) identify that patient expertise focuses on daily coping strategies and offers actionable advice, whereas the clinical expertise is more knowledge-oriented and prescriptive, focusing on medical topics. Hence, patients can offer important information that substantially differs from knowledge typically provided by clinicians, with a particular focus on personal issues experienced in the daily life context (Hartzler and Pratt, 2011). Prior work has shown how patient knowledge can be collected via ethnographic studies and by collecting techniques of daily practices (Pols, 2014; Attard and Coulson, 2012).

As the online communities are often driven by patients rather than medical professionals, the accuracy of the information has sometimes been raised as a concern. To overcome this potential issue, Attard and Coulson (2012) proposed that health care professionals could ensure the accuracy of the information in online support groups. However, Visser et al. (2016) pointed out that there exist norms for behavior in communities in which both professionals and patients co-exist that would hinder the openness of such a system. Visser et al. (2016) argue that patients would not want to ask too many questions, and also that the type of questions asked would change. Further, users would not want to complain too much or make knowledge claims about their own disease in their contributions.

Lederman et al. (2014) studied how users evaluate the credibility of information in online health forums. They stated that three types of information can be found from these forums, experiential (experiences with a condition), scientific (e.g., research-based information about the condition, treatment), and non-medical factual information (e.g., insurance policy regarding the condition). They further focused on users’ credibility evaluation on the experiential and scientific information. They identified five criteria for the credibility assessment; reference credibility (credibility of cited external sources), argument quality (logical soundness based on common sense), verification (confirming from several sources), contributor’s literacy competence (the way that a message is written), and crowd consensus (group opinion regarding the validity of an experiential statement). The used criteria depend on information type; argument quality, verification, and contributor’s literacy competence are used in forming an assessment of both experiential and scientific information. In addition, in validation of scientific information, reference credibility matters, and in validation of experiential information the crowd consensus matters.

Despite the benefits of these patient-driven message boards, some challenges also arise. Information can easily get lost in threads, especially if the volume of posts is large (Hartzler and Pratt, 2011). Another issue is the aforementioned correctness of the information, with prior work highlighting that the trustworthiness in online portals increases by the size of the community through self-correction (Bloem and Stocchi, 2015). Further, new users need to build a relationship with other forum members in order to get familiar with other users’ characteristics and their expertise level, a process that might be time-consuming (Mamykina et al., 2015). In the tool presented in this paper, we allow users to discover self-care techniques via pre-defined criteria, thereby allowing both recent and older contributions to the surface based on their relevance to the user’s query.

2.4. Crowdworking in healthcare

Crowdworking has been utilized in a range of healthcare-related applications, including diagnosis, surveillance, nutrition, and education (Wazny, 2018; Créquit et al., 2018; Pripp, 2017). In a review on the use of crowdworking in health and medicine, Ranard et al. (2014) classify crowdworking tasks into four groups according to the type of task: problem-solving, data processing, surveillance/monitoring, and surveying.

In our work, we focus specifically on manual crowd contributions, in which individuals contribute knowledge of their illnesses to support others. Hosio et al. (2016) contributed an online decision support tool, named AnswerBot, for this specific goal. AnswerBot enables the collection and assessment of solutions for a variety of problems and has been successfully utilized for the identification of suitable treatments for lower back pain (Hosio et al., 2018) as well as weight loss methods (Hosio et al., 2020). Lower back pain and weight loss as a health issue have similarities, as there is no ‘golden solution’ that works for all. Similarly, coping with PD is also a process in which patients need to find self-care techniques that work appropriately for them (Nunes and Fitzpatrick, 2018).

Crowdworked healthcare solutions also extend beyond the academic realm. Two popular platforms in this area are PatientsLikeMe1.
and StuffThatWorks\textsuperscript{2}, in which users can explore information statistics on several conditions based on community-contributed data. Through aggregate data, PatientsLikeMe allows users to find other patients with similar conditions or treatments, and subsequently to share their experiences (Brubaker et al., 2010). Users of the website experienced an increase in their knowledge of their symptoms, a better understanding of treatment side effects, and experienced the peer support as beneficial (Wicks et al., 2010). The StuffThatWorks forum\textsuperscript{2} collects data about the treatments and symptoms of different conditions, intending to utilize machine learning to identify the most effective treatments. The website’s data is currently not available for the general public and is shown only to registered users with a diagnosed condition in their profiles. In this paper, we also focus on the collection of crowdsourced data. In contrast to the discussed prior works, which focus on treatment and symptoms, our solution aims to offer actionable information for people with PD and their caregivers in a daily life context through self-care techniques. Using this approach, we hope to reduce the required effort for the user in identifying what actionable steps they can take as based on information that is typically presented to users in other sources.

3. Overview of the study

This study was conducted in two consecutive phases. In the first phase, the self-care data was collected and assessed. In the second phase, the participants discovered these self-care techniques through the established tool. The whole study was carried out in English, and due to that the recruitment activities focused mainly on English-speaking countries. The study procedures follow the guidance of our university and the Finnish National Advisory Board on Research Ethics, according to which a formal ethical review for this study (both phases) is not required (National Advisory Board on Research Ethics, 2019). In practice, this is due to the study not being classified as invasive and has no real medical intervention, focusing on information discovery. Next, Sections 4 and 5 describe the study’s phases in detail, covering the apparatuses, recruitment, participants and the results. Lastly, we discuss the findings of both phases in Section 6.

4. Phase 1: Self-care data collection

4.1. Apparatus

We built an online tool for collecting and assessing PD self-care techniques from the global PD community. The technical solution is based on AnswerBot, a flexible decision support tool that has been deployed across a number of problem domains (Hosio et al., 2016). The tool was implemented using HTML, JavaScript, PHP, and MySQL. Participants were directed to a website in which the study was introduced and, after consent was received, provided us with their demographic details as well as their connection to PD (patient, caregiver). Following this, participants were asked to share their daily self-care techniques and practices, and to assess the techniques submitted by others based on the pre-set criteria, namely: sociality, affordability, effectiveness, and familiarity. These four criteria were selected as a result of a workshop among the research team, as based on the disease’s impact on life and informed by prior work on crowdsourcing treatments for low back pain Hosio et al. (2018). PD may cause stigma (Nunes and Fitzpatrick, 2015) and bring discomfort in social circumstances (Hariz and Forsgren, 2011), as such the level of ‘sociality’ required by the technique is relevant. PD, similarly to other chronic illnesses, causes financial costs for the patient, e.g. medication, assistive equipment or rehabilitation (Wong et al., 2014). As such, financial resources might be strained — pointing to the importance of a solution’s affordability. When considering patient treatments or self-care solutions, effectiveness is of obvious importance. Finally, as PD patients are often presented with a range of techniques throughout their life, familiarity was added as a final criterion.

Fig. 1 shows the specific look and feel of the user interface (UI) as well as the detailed descriptions of these criteria. The UI displays the list of submitted techniques sorted in descending order with the techniques with the least accumulated ratings on top of the list. Participant assessment was collected via a slider input (0-100), and the user may choose to assess 1–4 of the criteria. The use of such sliders for assessing arbitrary options has been shown as highly intuitive and easy to use in other contexts (Hosio et al., 2016). Further, we followed the suggestions by Matejka et al. (2016) in the slider implementation: The value is shown in real-time when moving the slider handle, and there are no tick marks on the axis. In addition, the participant was asked which symptom or PD-related issue the technique is suitable for through an open-ended text field. The interfaces for the collection and subsequent assessment of self-care techniques are shown in Fig. 1. When submitting a technique, participants provided a title and description, indicated whether the technique could be practiced at home, and provided an optional hyperlink to provide additional details. The participants were allowed to submit as many techniques as they wanted. In order to bootstrap the system and to avoid the cold-start problem, we manually inserted five self-care techniques that we obtained from Parkinson’s UK and Parkinson’s Foundation web site before offering the tool to the general public.

4.2. Recruitment

We invited both people with PD and their personal caregivers to join the study. There was no reward for participating, but the participants were offered a chance to join a raffle of five Amazon gift cards ($100). In order to recruit participants, we engaged in extensive contact with several organizations and foundations related to PD, supplemented with Facebook groups and bloggers. The European Parkinson’s Disease Association forwarded our message to their member organizations and to their online forum. Parkinson’s UK added our study to their online ‘Take Part in Research’ section and sent the message to their ‘Take Part Hub’. The Davis Phinney Foundation shared the invitation in their email list and Parkinson Society British Columbia added our study to the Research Volunteer Recruitment page. ParkinsonsDisease.net added a post to their Facebook group. Parkinson Association of the Rockies, Parkinson Society Newfoundland and Labrador, the Cure Parkinson’s Trust, Parkinson Wellness Project, and The Finnish Parkinson Association agreed to share the information with their members. We created a profile for advertising the study on Parkinson’s Resource Organization’s platform. Additionally, we added a post to Parkinson’s UK discussion forum on their ‘Research Opportunities’ page and shared our post through several social media accounts.

4.3. Participant profiles

The data collection started in January 2021 and is ongoing. For the analyses of the collected data as presented in this paper (Phase 1), we took a snapshot of the database at the end of May 2021, when Phase 2 started. A total of 320 participants had submitted and/or assessed one or more self-care techniques through the website (see Fig. 1), along with the background demographics questionnaire. The majority of our participants were from the UK and USA, with 11 countries of origin reported. Of this sample, 288 participants were diagnosed with PD, and 31 acted as a caregiver of someone with PD. We provide an overview of participant demographics in Table 1.

In the questionnaire, we asked the participants to select their symptoms from a list based on (Massano and Bhatia, 2012). Table 2 presents...
Fig. 1. On the left, the interface for assessing a self-care technique. When submitting, the text in the top-right box is shown to the user, and the technique is submitted with the form on the bottom-right box.

### Table 1

Participant details. Note, in the case of caregivers the disease-specific data refers to the person for which they act as caregiver. The categories for education level are primary education, secondary education, vocational degree, bachelor’s degree, master’s degree, PhD, or above. The categories for work status are work full time, work part-time, unemployed, and retired.

<table>
<thead>
<tr>
<th>Metrics</th>
<th>PD Patients</th>
<th>Caregivers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of participants</td>
<td>288</td>
<td>31</td>
</tr>
<tr>
<td>Mean age (range min-max)</td>
<td>67.3 (38–86)</td>
<td>65.7 (44–86)</td>
</tr>
<tr>
<td>Mean years since PD diagnosis</td>
<td>5.7</td>
<td>9</td>
</tr>
<tr>
<td>Mean years since onset of symptoms</td>
<td>8.8</td>
<td>11</td>
</tr>
<tr>
<td>Women/Men/Non-binary/ Prefer not to disclose</td>
<td>57%/43%/0%/0%</td>
<td>74%/26%/0%/0%</td>
</tr>
<tr>
<td>Education: Prim./Sec./ Voc./Bac./Mas./PhD</td>
<td>1%/27%/21%/27%/18%/6%</td>
<td>0%/19%/35%/23%/16%/6%</td>
</tr>
<tr>
<td>Work status: Full time/ part time/unemp./retired</td>
<td>8%/9%/6%/77%</td>
<td>23%/13%/16%/48%</td>
</tr>
</tbody>
</table>

### Table 2

The frequency of reported symptoms (as percentage). The three largest frequencies are shown in bold.

<table>
<thead>
<tr>
<th>Symptom</th>
<th>PD %</th>
<th>Caregivers %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Motor symptoms</td>
<td>69.8</td>
<td>64.5</td>
</tr>
<tr>
<td>Bradykinesia (i.e. slowness of movements, decrease of amplitude or speed in consecutive movements)</td>
<td>61.8</td>
<td>58.1</td>
</tr>
<tr>
<td>Tremor</td>
<td>58.7</td>
<td>45.2</td>
</tr>
<tr>
<td>Rigidity (stiff or inflexible muscles)</td>
<td>59.7</td>
<td>80.6</td>
</tr>
<tr>
<td>Postural and gait impairment</td>
<td>31.9</td>
<td>35.5</td>
</tr>
<tr>
<td>Mood and cognition related symptoms</td>
<td>35.8</td>
<td>54.8</td>
</tr>
<tr>
<td>Anxiety, panic attacks</td>
<td>9.0</td>
<td>32.3</td>
</tr>
<tr>
<td>Mood disorders, e.g. apathy or depression</td>
<td>20.1</td>
<td>54.8</td>
</tr>
<tr>
<td>Hallucinations, illusions, delusions</td>
<td>1.0</td>
<td>9.7</td>
</tr>
<tr>
<td>Cognitive impairment</td>
<td>18.4</td>
<td>22.6</td>
</tr>
<tr>
<td>Dementia</td>
<td>7.0</td>
<td>7.1</td>
</tr>
<tr>
<td>Sensory dysfunction and sensory symptoms</td>
<td>49.3</td>
<td>58.1</td>
</tr>
<tr>
<td>Orthostatic hypotension (drop in blood pressure when getting up)</td>
<td>16.0</td>
<td>25.8</td>
</tr>
<tr>
<td>Constipation</td>
<td>26.0</td>
<td>25.8</td>
</tr>
<tr>
<td>Urinary dysfunction (urgency, retention)</td>
<td>17.7</td>
<td>16.1</td>
</tr>
<tr>
<td>Sexual issues</td>
<td>20.9</td>
<td>38.7</td>
</tr>
<tr>
<td>Sweating problems</td>
<td>39.2</td>
<td>41.9</td>
</tr>
<tr>
<td>Sleep disorders (e.g. insomnia, REM behavior disorder, daytime sleepiness, restless legs)</td>
<td>43.4</td>
<td>41.9</td>
</tr>
<tr>
<td>Pain</td>
<td>80.6</td>
<td>80.6</td>
</tr>
</tbody>
</table>
Table 3
The stages of the Modified Hoehn & Yarn scale, used to describe the progression of PD (Goetz et al., 2004), and the distribution in our sample. Note, in the case of caregivers the disease-specific data refers to the person for which they act as caregiver.

<table>
<thead>
<tr>
<th>Stage</th>
<th>Description</th>
<th>PD %</th>
<th>Caregivers %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stage 0</td>
<td>No signs of disease.</td>
<td>1.7</td>
<td>0</td>
</tr>
<tr>
<td>Stage 1</td>
<td>Symptoms on one side only (unilateral).</td>
<td>26.4</td>
<td>9.7</td>
</tr>
<tr>
<td>Stage 1.5</td>
<td>Symptoms unilateral and also involving the neck and spine.</td>
<td>10.1</td>
<td>6.5</td>
</tr>
<tr>
<td>Stage 2</td>
<td>Symptoms on both sides (bilateral) but no impairment of balance.</td>
<td>14.9</td>
<td>0</td>
</tr>
<tr>
<td>Stage 2.5</td>
<td>Mild bilateral symptoms with recovery when the ‘pull’ test is given (the doctor stands behind the person and asks them to maintain their balance when pulled backward).</td>
<td>14.2</td>
<td>19.4</td>
</tr>
<tr>
<td>Stage 3</td>
<td>Balance impairment. Mild to moderate disease. Physically independent.</td>
<td>28.8</td>
<td>38.7</td>
</tr>
<tr>
<td>Stage 4</td>
<td>Severe disability, but still able to walk or stand unassisted.</td>
<td>3.5</td>
<td>19.4</td>
</tr>
<tr>
<td>Stage 5</td>
<td>Needing a wheelchair or bedridden unless assisted.</td>
<td>0.3</td>
<td>6.5</td>
</tr>
</tbody>
</table>

Table 4
Distribution of impact of PD on daily life as based on the Parkinson’s Disease Activities of Daily Living Scale (PADLS) (Hobson et al., 2001). The PADLS is used in assessing how PD affects to patient’s day-to-day activities (e.g. housework, shopping, dressing, eating) in the last month. Note, in the case of caregivers the disease-specific data refers to the person for which they act as caregiver.

<table>
<thead>
<tr>
<th>Level</th>
<th>PD  %</th>
<th>Caregivers %</th>
</tr>
</thead>
<tbody>
<tr>
<td>No difficulties</td>
<td>16.0</td>
<td>3.2</td>
</tr>
<tr>
<td>Mild difficulties</td>
<td>62.2</td>
<td>32.3</td>
</tr>
<tr>
<td>Moderate difficulties</td>
<td>18.4</td>
<td>51.6</td>
</tr>
<tr>
<td>High levels of difficulties</td>
<td>2.4</td>
<td>9.7</td>
</tr>
<tr>
<td>Extreme difficulties</td>
<td>1.0</td>
<td>3.2</td>
</tr>
</tbody>
</table>

the distribution of participants’ self-reported symptoms. The most common symptoms reported by people with PD were bradykinesia, sleep-related issues, and tremors. On average, the participants with PD reported 6.9 symptoms per participant. The three most common symptoms reported by the caregivers in our sample were postural and gait-related issues, sleep-related issues, and constipation. On average, the caregivers reported 8.5 symptoms per participant. Of note is that our participant sample included all symptoms listed in the survey.

Furthermore, participants self-assessed their PD status (or the PD status of the person for which they act as caregiver) using the Modified Hoehn & Yarn scale (Goetz et al., 2004), see Table 3, as well as the impact of PD on their daily life with Parkinson’s Disease Activities of Daily Living Scale (PADLS) (Hobson et al., 2001), see Table 4.

4.4. Descriptive statistics of submitted techniques and assessments

We first sought to understand both our participant sample and their submitted techniques through descriptive statistics. Our participants submitted in total 115 individual self-care techniques. 24% of all participants submitted one or more techniques, 93.5% of them had PD and 6.5% were caregivers. 64.9% of all submitters were women and 35.1% men. The participants with PD submitted 87.8% of the techniques, and the caregivers the remaining 18.2%.

The total number of times a technique was assessed using one or more criteria was 1052, and the total number of individual ratings was 3876. A large majority of 75.7% of participants with PD, and 80.6% of caregivers assessed one or more techniques (using one or more criteria). On average, participants with PD rated 3.3 techniques per person and caregivers 3.1 techniques per person.

Table 5 shows the best-rated techniques per each criterion. From the caregiver’s point of view, taking a loved one to doctor’s appointments is social, affordable, familiar, and effective. The top-3 of each criterion according to the participants with PD, is more versatile.

During the assessment, participants had the possibility to describe what symptoms or PD-related issues they believe the technique is good for through the use of an open-ended field. We will refer to this input as a participant reflection. 1121 individual reflections were submitted, an average of 9.7 reflections per technique. 69.8% of participants with PD submitted a reflection for one or more techniques, as compared to 61.3% of the caregivers. 38% of the participants providing reflections were men, and 62% were women.

4.5. Qualitative analysis of the techniques

4.5.1. Categorization of the techniques

Next, we set out to categorize the submitted PD self-care techniques in order to obtain a better understanding of participant contributions. Our analysis is based on the reflective thematic analysis framework by Braun and Clarke (2012). First, two of the authors familiarized themselves with all provided techniques and coded the data. Subsequently, they formed nine categories with descriptions (see Table 6) by combining codes, and assigning the techniques to the categories. The techniques were subsequently assigned into these categories independently by two additional authors. Some of the techniques matched to multiple categories, e.g. ‘Walking with hiking sticks’ matches both with the ‘Physical activity and exercise’ and the ‘Equipment, aids and re-organizing’ categories. All authors identified what they believed to be the most suitable category for the technique, and if agreement could not be reached in a subsequent discussion, we allowed techniques to be categorized into multiple categories.

The biggest category was ‘Physical activity and exercise’. Physical exercise may positively impact PD patients’ daily activities, as it maintains physical abilities and alleviates both motor and non-motor symptoms (Lauzé et al., 2016; Murray et al., 2014; Cusso et al., 2016), the versatile impact of exercise was clearly presented in the provided reflections. The exercise was described to alleviate many motor symptoms, such as rigidity and pain, and improve balance, but also to improve mental health and alleviate stress or anxiety. Stress may worsen both the motor and non-motor symptoms (van der Heide et al., 2021), and the second biggest category, ‘Well-being’, offered means for relaxation and symptom management, as well as for keeping a positive mindset. Having interesting hobbies helps in keeping the mind off the disease, and often provides socializing with others. The ‘Social interaction’ category provides techniques for handling social situations.

Diet has also an important role in PD self-care, which is reflected in the number of ‘Diet and nutrition’ category techniques. In addition to symptoms related to autonomic dysfunction, some PD medication might have diarrhoea or constipation as a side-effect (Gage et al., 2011), and some medication regimens suggest low protein intake (Barichella et al., 2009).

‘Medication and treatment’ related techniques highlighted the importance of finding suitable treatment options and encouraging patients to be involved in treatment decisions. ‘Equipment, aids and re-organizing’ category provided practical tips for managing with the symptoms.

4.5.2. The content of reflections

In the reflections, which were analyzed through an inductive coding process, the participants often described which symptoms the technique tackles. For example, ‘going out daily’ was described as mitigating stiffness and pain and improving balance, whereas ‘walking with hiking sticks’ was described as reducing symptoms related to mobility issues.
stability supports the disease. For the technique ‘Be Your Own Best Advocate’ participants described her feelings: ‘’

It is sensible for people our age (68 years) to listen to moans of others! I care for my husband but want any free time to help. Problem was that a very few people end up running activities for others who consider themselves unable to volunteer. It became overwhelming. As a table tennis player described it helps in ‘’

‘’ One participant pointed out that he actively tries to discuss also other things than his PD: ‘’[...] I also try to drive the conversation to their health so that it is not just me and my Parkinson’s.’

Even though many feel participation in a local PD group as a good place for social interaction, some might experience it as overwhelming: ‘’

‘’ A carer described choir singing: ‘’Gives me a break from living with a PD sufferer and the constant caring and emotional turmoil.‘’ One participant pointed out that he actively tries to discuss also other things than his PD: ‘’[...] I also try to drive the conversation to their health so that it is not just me and my Parkinson’s.’

Even though many feel participation in a local PD group as a good place for social interaction, some might experience it as overwhelming: ‘’Have been part of an active group that offered social, physical exercise and vocal help. Problem was that a very few people end up running activities for others who consider themselves unable to volunteer. It became overwhelming. As personal difficulties increase with disease progression I as carer do not want to listen to moans of others! I care for my husband but want any free time and energy to be as far away from Parkinson’s as possible.”

4.6. Participant engagement

The high level of engagement we received from participants in this study was a positive surprise to us. Several participants contacted us
with various questions and suggestions. For example, one of the participants had inputted wrong dosing to a recommendation of using a food supplement, and she asked us how to correct the inputted technique — something that is not currently supported in the user-facing side of the application. Another participant wanted to verify whether their contribution was stored following a page refresh. Another technical challenge was faced by a participant receiving a security warning from the antivirus application in their computer, wondering whether the page was secure. At this time our tool did not force a redirect from http to https, something we have since changed. Finally, one participant reached out to state that he is willing to participate in the study, but he does not have any self-care practices in mind apart from medication. We suggested he join the study without providing new self-care techniques, as providing new data here is not mandatory — instead, he was able to browse and assess the items submitted by others.

5. Phase 2: Discovering self-care techniques using PDCareBox

In the second phase of the study, our goal was to have the community, i.e. people with PD and their caregivers, discover techniques among the knowledgebase built as a result of the first phase.

5.1. Apparatus

The users were directed through various recruitment means, much like in the first phase, to the final implementation of PDCareBox at PDCareBox.com. The page contains a short introduction and instructions on how to use the tool on top, and below that the search interface (see Fig. 2). The user specifies the desired level for 1–4 criteria, and clicks ‘Discover best matches’. Operating the search interface brings up the results interface (see Fig. 3). Each result contains a ‘Show what this means’ section which includes possible reflections obtained from participants during the assessment phase (see Fig. 3, E). The reflections are accompanied by the participant’s age and years with Parkinson’s Disease to emphasize the fact that these are reflections left by other people with Parkinson’s or their caregivers. A new search can be started by resetting the sliders, or by simply changing the position of sliders and pressing ‘Discover best matches’. After using the tool for a while (four search rounds), a pop-up reminder about the questionnaire is shown. The link to the questionnaire is also available in the PDCareBox. When entering the questionnaire, the study was first introduced and the consent was confirmed. After that, the questionnaire queried the following topics:

- Background demographics: are they diagnosed with PD, caregiver or someone else, gender, year of birth, years since diagnosis, and PD status using the Parkinson’s Disease Activities of Daily Living Scale (PADLS) (Hobson et al., 2001)
- Self-care information: e.g. what are their typical information sources, do they search information on a specific topic or in general, how does the information from different sources (peer/ professional) differ
- Evaluation of PDCareBox: e.g. benefits, downsides, the usability of the search criteria, improvement ideas

For the second phase, we enhanced the accessibility of the self-care technique discovery interface used in earlier AnswerBot studies (Hosio et al., 2016, 2018, 2020) by following design guidelines for older adults (Consortium, 2021b, a; Nurgalieva et al., 2019, 2021), and also design guidelines for people with Parkinson’s (Nunes et al., 2016).

PD may cause visual disabilities (Nunes et al., 2016), and these hindrances may worsen as people age. To ensure that the text is readable, the text line height of the interface was increased from baseline 1.2 to 1.5 (see Fig. 3, A) as suggested in (Consortium, 2021b; Nurgalieva et al., 2021). We increased the text font size to be 16pt at a minimum across the interface. To ensure that each of the shown techniques is distinguishable from each other, a thematic break was added in between them (see Fig. 3, D).

A high contrast between background elements and foreground text is also needed to support users for with slight visual impairments (Nunes et al., 2016; Nurgalieva et al., 2021). The colors of the criteria slider items and buttons were updated to having a contrast ratio of 4.5:1 (see Fig. 2, B), following (Consortium, 2021b; Nurgalieva et al., 2021).

Standard icons might be unfamiliar to elder users, therefore it is advisable to reinforce these icons with words (Nurgalieva et al., 2021). We replaced an icon for the user-provided URL with the text ‘Learn more’ (see Fig. 3, B). This text link is blue and underscored to make it distinguishable from regular text. As short-term memory loss is commonly accentuated with PD (Nunes et al., 2016), we made non-visited and visited links distinguishable from each other by changing the color of the visited links (Nurgalieva et al., 2021).

The baseline interface showcased the goodness-of-fit of the search as the numerical euclidean distance (0-100) between the search and the participant ratings. To be more informative, we used a more visual indicator (see Fig. 3, C) of a traffic light (red, yellow, or green) and accompanying text (poor, moderate, or close). The interface element next to the criteria name shows the numerical value of the slider position. To highlight the functionality of sliders, we replaced ‘-’ indicating inactive slider with a label ‘Move slider to apply’ (see Fig. 2, A).

The source code of the PDCareBox is publicly available in Github (https://github.com/EetuHuusko/PDCareBox).

5.2. Recruitment

After completing the first phase of the study, the participants had an option to leave their emails to get invitations to the second phase. We utilized this email list in recruiting for the second phase. In addition, we re-contacted those PD organizations that had participated in recruitment in the data collection phase.

The European Parkinson’s Disease Association added our study in a monthly news roundup of surveys and research opportunities. We created a post into the Parkinson’s UK discussion forum to section Research opportunities. The Finnish Parkinson Association added the announcement to their web page’s news section. Parkinson’s Society of Newfoundland and Labrador shared the announcement for their members. A video introducing the study and the tool interface was presented in a Parkinson’s Resource Organization’s support group meeting. We shared the link to the PDCareBox in ParkinsonsDisease.net Facebook page.

5.3. Participant profiles

Thus far, more than 120 people have tried discovering self-care ideas with PDCareBox. 23 persons of those who tried PDCareBox, replied to the provided questionnaire and hence participated in our study. See the overview of participant demographics in Table 7.

<p>| Table 7 |
| Participant details. Note, in case of caregivers the disease specific data refers to the person for which they act as caregiver. PADLS refers to the Parkinson’s Disease Activities of Daily Living Scale (Hobson et al., 2001). |</p>
<table>
<thead>
<tr>
<th>Metrics</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of participants</td>
<td>23</td>
</tr>
<tr>
<td>Mean age (range min-max)</td>
<td>66 (43–80)</td>
</tr>
<tr>
<td>Mean years since PD diagnosis</td>
<td>5.2 (1–13)</td>
</tr>
<tr>
<td>Women/Men/Non-binary/Prefer not to disclose</td>
<td>56.5%/43.5%/0%/0%</td>
</tr>
<tr>
<td>PADLS: None/Mild/Moderate/High/Extreme</td>
<td>21.7%/69.6%/8.7%/0%/0%</td>
</tr>
</tbody>
</table>
5.4. Self-care information

We asked the participants how they typically search for information about self-care, e.g. what information sources they rely on, and do they search for information on a specific topic, or browse for new ideas in general.

Parkinson’s organizations’ websites were the most popular self-care information source, 39% of the participants mentioned the organizations in general or specified which organization’s websites they use. 26% mentioned internet or internet search, including YouTube. One participant specified that when searching information on the internet, you have to evaluate the trustfulness of the information source. 22% said that they rely on information provided by peers. Also, articles, news, and research related to PD were mentioned as information sources by 17% of the participants. 34% told they search for information on a specific topic, 13% said they usually browse in general. One participant mentioned the continuous need for new information due to the disease progression: “I search for specific answers to specific topics. As my Parkinson’s progresses I have new issues to find coping strategies for!”

We asked how the information provided by peers differs from the information provided by professionals. The participants described information provided by professionals to be research-based, more theoretical, and focused on drugs: “The information from peers usually is easy to accept and take on board because of their own experiences and empathy... professionals are more matter of fact, instructive, impersonal.”

The information provided by peers was experienced as practical, and to be relevant in everyday context: “It has more insight into everyday problems that the professionals don’t even think about.”

Peers also understand your situation, as they have experience of living with the condition: “It is often more specific; you have to have the condition in order to really understand what works/doesn’t work; it shows that not all techniques suit all people; it shows that people with Parkinson’s are positively engaged in finding a cure or the easement of symptoms and are willing to share their experiences with others.”

However, few participants brought up the reliability of the peer data. These participants stressed that the observed information might be biased, or even dangerous: “I don’t always trust peer advice, it can range from useful to dangerous.”
5.5. Experiences with PDCareBox

5.5.1. Search statistics

Participants completed between 1–6 searches, with an average of 2.2 searches per participant. Participants could specify the level for 1–4 criteria for each search. On average, participants used 3.1 criteria per search. All four criteria were utilized in the majority of the searches (64.4%), and in 22.2% of the searches only one criterion was specified. Three criteria were used in 11.1% of the searches and two criteria only in 2.2% of the searches. Effectiveness was considered the most important criterion in our study sample, as it was used in 91.1% of searches. Affordability was used in 77.8%, sociality in 75.6%, and community familiarity in 73.3% of the searches. Fig. 4 shows the spread of values of each criterion used in searches.

5.5.2. Adoption of the new search approach

34.7% of the participants described PDCareBox positively, for example, using words interesting, helpful, useful, or novel. A slightly smaller share, 30.4% gave negative feedback. They experienced the tool as not useful, confusing, or bewildering. The same amount of participants (30.4%) gave a neutral response (e.g. do not know or do not have an opinion).

Participants described how identifying a suitable self-care technique with PDCareBox differs from their typical approach. Compared to informal information exchange in e.g. a peer meeting, information search with our tool starts with setting the criteria “I was starting from a set of criteria rather than just hearing spontaneous suggestions from others”. On the other hand, many participants compared the tool to an internet search: “I have always searched for specific topics before”. The new search approach using four criteria was adapted well by some users: “It’s a new way to choose what might be interesting. Makes it easier to narrow the research”, and “Interesting. At first wasn’t sure what to do, but soon got the idea. Useful to hear about other people’s experiences and match them with my own.”. Users described the PDCareBox as quick, patient-specific, and accessible. One user specified that the sliders help in choosing the suitable technique, while another user experienced the sliders as subjective, not knowing e.g. what in practice highly affordable would mean. As a downside, a participant pointed out that finding something specific using the criteria would take a while to locate. As a practical design implication, three participants suggested enhancing the search by filtering the search results by categories: “I would find it useful to have a kind of index of subjects that I could click on, then go to the 4 criteria. Eg: Diet, Muscle stiffness, Swallowing etc””. Few participants felt that better instructions would make understanding the tool easier.

5.5.3. The search criteria

The participants also evaluated the usefulness of each criterion on a five-point scale (‘Not at all useful’ to ‘Extremely useful’), see Fig. 5. This is aligned with the actual search statistics. As stated before, effectiveness was used in 91.1%, affordability in 77.8%, sociality in 75.6%, and community familiarity in 73.3% of the searches.

In addition, we asked if they would like to add some criteria. We got a suggestion of frequency of participation, degree of side effects, the level of required training of the technique, degree of difficulty, and alignment with science. In addition to no responses, a participant specifically stated more criteria would not benefit: “No, the more you have the more complicated it gets.”

5.5.4. Evaluation of the search results

Participants evaluated the self-care techniques provided by PDCareBox on different attributes on a four-point scale from ‘Strongly disagree’ to ‘Strongly agree’, see Fig. 6. The results highlight the positive aspects of the data. We asked with an open-ended field how well the search result matched the required level of criteria. 65% responded positively, 22% gave a negative answer and 13% said they did not know.

One participant described he would be more comfortable if PDCareBox would offer data provided by professionals. However, the variable content with shared experiences was seen also as a benefit: “Wide variety of information provided, easy to understand/ read, comments valuable may discover and learn about new techniques etc” “There may be a wide range of options shown than would usually be suggested. For example, Reiki”. The experience of the disease itself is subjective, and even though the tool would give the techniques matching to the criteria, the user has to evaluate the suitability of the proposed techniques personally: “Parkinson’s is highly Individual. Must judge whether suggestion can be effective for me.” The idea of more data providing better search results was understood: “By continuously adding peoples experience will enrich the data and lead to more options and accurate searches.”

5.5.5. Use of PDCareBox for different self-care functions

According to the Middle-Range Theory of Self-Care of Chronic Illness (Riegel et al., 2012) self-care can be divided into three sections, self-care maintenance, self-care monitoring, and/or self-care management. Referring to this, we asked which of these self-care areas the participants would make use of PDCareBox: (1) Maintaining physical and emotional state, (2) Monitoring and detecting changes in the physical and emotional state, including symptoms, (3) Making conclusions of the physical and emotional state, making decisions on self-care, or (4) none of these. A large majority of 73.9% of the respondents would use it for self-care maintenance (option 1), 26.1% for self-care monitoring (option 2), 39.1% for self-care management (option 3). The remaining 17.4% would not use this tool for any of these purposes (option 4).

6. Discussion

Parkinson’s Disease is a condition with an exceptionally broad range of symptoms that can progress from barely noticeable to extremely severe. In any case, its effect on perceived QoL can be detrimental, and there are no single solutions that work universally for everyone. As such, different self-care techniques are important for PD just as they are important for various other chronic diseases and conditions. To this
end, in our work, we contribute a public online repository of PD self-care techniques, PDCareBox, with a continuously growing database of structured content that is provided and evaluated by people with PD. While we attempted to tailor the UI to be suitable for use by people with PD, we acknowledge that it was still found somewhat confusing by many. Based on this and other feedback from our study participants, we also identify opportunities for future work — both in the development of PDCareBox, as well as in the design of future patient-driven knowledge repositories (see Section 6.4). We publicly release the source code of PDCareBox for use by the (research) community. The longitudinal deployment was not one of the easiest studies to conduct, and we also wish to reflect on our experiences on reaching the vulnerable group of participants and end-users in Section 6.3.

6.1. Self-care techniques providing actions towards coping

Self-care is an essential part of living with a chronic disease, including both daily mundane actions of dealing with the symptoms, as well as dealing with the disease emotionally (Nunes and Fitzpatrick, 2018). We identified nine categories for the collected PD self-care techniques: 'Physical activity and exercise', 'Well-being', 'Equipment, aids and re-organizing', 'Leisure and hobbies', 'Medication and treatment', 'Diet and nutrition', 'Social interaction', 'Knowledge', and 'Monitoring' (see Table 6 for descriptions and frequencies of the categories). According to the user feedback, techniques provide actionable, understandable, and useful information.

Some of the provided self-care techniques were similar to the practical methods identified in Miertová et al. (2014). They collected the activities through Activities in Daily Life scale. The techniques in PDCareBox are collected as open-ended text, hence, the topics of techniques in PDCareBox were more various, spreading to all aspects of daily life. We found similarities also with the categorization of Nunes and Fitzpatrick (2018), with the taking of medication included in 'Medication and treatment', exercising in 'Physical activity and exercise', 'Well-being', and 'Leisure and hobbies', and adapting lifestyle in 'Equipment, aids and re-organizing'. The category accepting the disease consequences is more related psychological coping, and as such does not directly match to our categories, but the reflections of some techniques describe they may help in adapting to a more accepting attitude towards PD. Our approach differs from these earlier papers.

In prior work participants were interviewed and observed, whereas in our study, the participants provided the practices related to self-care as a self-care contribution to the PD community.

There are several positive outcomes related to self-care, such as maintaining quality of life (Riegel et al., 2012), and our participants described the outcomes for the self-care techniques in the open-ended fields during assessment as reflections. Tuijt et al. (2020) investigated PD self-management themes through a literature review, and Haahr et al. (2021) similarly studied categories of PD coping strategies via a metasynthesis. These studies focused on how people with Parkinson’s manage in daily life. In our study, the categories of the self-care techniques have similarities with these studies, but produced more actionable data, highlighting what it is in practice the people living with Parkinson’s do. For example, Tuijt et al. (2020) and Haahr et al. (2021) both defined maintaining independence as a category of coping strategies or self-management. In our study, the independence was an outcome described in the reflections for many practical techniques, such as utilizing assisting equipment, as described in Section 4.5.2.

Inside the theme of maintaining independence, Tuijt et al. (2020) describe the contradiction in receiving help and feeling independent. Similar feelings were described in the reflections by our participants (see Section 4.5.2), on the other hand, they valued techniques supporting independence, and at the same time appreciated the support and help by their families and friends. The psychological strategies category by Tuijt et al. (2020) had similarities to focus to the present from Haahr et al. (2021). We categorized these in techniques for wellbeing. In this category, maintaining normalcy/(pre-Parkinson’s) self was mentioned as a sub-item of focusing on the present strategy by Haahr et al. (2021). Again, in our study, feeling ‘normal’ or having a break from PD was mentioned as an outcome of various techniques in the reflections (see Section 4.5.2).

The Middle-Range Theory of Self-Care of Chronic Illness divides self-care into the categories of self-care maintenance, self-care monitoring, and self-care management (Riegel et al., 2012). A majority of the submitted techniques fall into self-care maintenance, and similarly, most of our participants (73.9%) also indicated that they could imagine using our tool for this purpose. A good quarter (26.1%) would use it for self-care monitoring, and approximately 40% for self-care management.

Participants in our study submitted only a few techniques for monitoring health and symptom status. For us, this was perhaps slightly surprising, as modern mHealth and eHealth solutions are increasingly being used and offered for monitoring various symptoms and body functions (Espay et al., 2016, 2019). We hypothesize that the low number of monitoring techniques submitted is a direct result of the perceived lack of benefit of such tools among the majority of people living with PD, possibly based on prior negative experiences. According to the prior research, less than 40% prepare a list of changes in symptoms for doctor’s appointment (Bloem and Stocchi, 2015). A slightly larger share of the techniques were related to obtaining knowledge and information about PD, which would support decision-making in self-care management. This type of self-care technique is likely to be perceived as more relevant for other types of knowledge gathering — as for example typically found through general search engines.

6.2. Validity of community-contributed techniques

As the sample of our study was recruited through online PD communities, it comes as no surprise that the participants were used to utilizing the internet in obtaining PD-related information. The starting point in using general online search tools, such as Google or Bing, is to identify a theme or a problem area on which the user wishes to obtain more information. The logic behind PDCareBox is different, as the starting point for any search is the selection and weighting of criteria. In online PD communities, peer information is usually exchanged via discussion forums. While often containing rich information, prior work has identified the challenges that arise when trying to find information in this largely unstructured data source (Hartzler and Pratt, 2011). Interestingly, most of our participants did not compare our tool to discussion forums, but rather to internet search.

Aligned with prior research (Hartzler and Pratt, 2011), our participants described the information provided by peers differs from information by professionals in terms of context (daily life vs. medical issues) and form (actionable vs. theoretical/informative), highlighting the practicality of peer provided data. This underlines the importance of sharing (self-care) information as gained through experience with living with PD, to complement the information by professionals. However, the correctness or accuracy of patient-provided health information sometimes raises concerns. In our study, few of our participants described the peer information might be biased, and you have to consider if the techniques provided would be suitable for you, effective, or even safe.

The crowd consensus is an important factor when assessing experience-based information (Lederman et al., 2014), and in online health forums, the common understanding is found via discussion (Huh and Ackerman, 2012). In some platforms, for example, Patient Innovation (Patient Innovation, 2021), the submitted solutions from the users are evaluated by a medical team. While this is suitable in the solution-oriented approach of the Patient Innovation platform, the content of discussion forums might be affected by the presence of professionals (Visser et al., 2016). We will discuss more how these factors shall be taken into account in the future in Section 6.4.

Mamykina et al. (2015) point out the difference in traditional discussion forums and platforms focusing on collective sensemaking,
such as Stack Overflow. Traditional discussion forums allow a diversity of perspectives, and in sites like Stack Overflow the users vote for the ‘best answer’, across one simple criterion of “goodness”. In PDCareBox, the users assess the submitted techniques according to various related criteria, highlighting the different aspects of each technique in a more nuanced, multidimensional fashion. And as evident based on the data log, people also used this functionality when discovering techniques: various combinations of the criteria were used. This approach allows users to search not only for the ‘best’ solution but rather find suggestions that fulfill simultaneously a set of wanted criteria. The reflections provide additional information on the effects of the technique. PDCareBox does not replace discussion forums, which are also a source of empathy and social contacts but act as a novel way for discovering peer-provided information. It also differs from traditional Internet search, as all the data is community contributed and in a structured form.

6.3. Involving the people with PD in online research

6.3.1. Comparing our sample to the general PD population

Prior work has highlighted that internet usage is dependent on age, but there is an indication that having PD does not affect internet use (Marxreiter et al., 2020). Prior work has studied PD burden (Pohar and Jones, 2009) as well as general prevalence of PD in Canada (Wong et al., 2014), allowing us to compare the characteristics of participants in Phase 1 to their respective samples (see Table 8). The mean age of our participants with PD was 67.3 years, slightly lower than in the study by Pohar and Jones (2009), 68.7 years, which reflects also the mean years since diagnosis. We find a clear difference in the education level of the participants with PD, with our sample having a much higher educational attainment as compared to a prior PD sample (Pohar and Jones, 2009). In general, higher education is connected to larger access to the internet and to higher levels of internet use, which may explain the difference (Saboor et al., 2015; Berner et al., 2012). In an online study in Sweden with 346 participants with PD, more than half of the participants with university education (Riggare et al., 2019).

PD is slightly more common with men than with women (Wong et al., 2014; Pohar and Jones, 2009), and men are also more likely to use the internet as compared to women (Morris et al., 2007). In contrast, women are more likely to search for health-related information from the internet (Fox and Fallows, 2003; Riggare et al., 2019). Aligned, the participants with PD in our sample contained more women by a large offset. Similarly, among caregivers, our sample contained more women than reported by Wong et al. (2014) (see Table 8). We hypothesize that women are more likely to share their experiences with PD; in our sample, women were more likely to submit techniques and to provide reflections. This is supported by literature too, Seale et al. (2006) studied internet support groups and interviews of people with cancer by comparing the language of men and women. They noticed that compared to men, women seek and provide emotional support more, and express their feelings and emotions more.

Finally, we find that the caregivers in our sample reported more advanced stages of PD with the Modified Hoehn & Yann scale (Table 3) and higher level of difficulties with the Parkinson’s Disease Activities of Daily Living Scale (Table 4). In addition, on average caregivers reported more symptoms per participant. This is aligned with the fact that the need for support is bigger in the later stages of PD, perhaps enticing participants to contribute to and make use of, the self-care techniques available in our knowledge base.

6.3.2. Experiences with recruiting

Even though our tool was clearly branded as a collaboration between several academic institutions, we note that some users felt unsure about our motives. This is not surprising though, as typical internet safety advice is to be cautious. For example, Age UK instructs people not to click things online without thinking (UK, 2021), and European Parkinson’s Disease association advises to always carefully evaluate the reliability of an online information provider, e.g. is it a commercial website (The European Parkinson’s Disease Association, 2021)? Indeed, we got a few emails from our participants related to security and if there are commercial purposes behind the PDCareBox. When describing self-care information search, our participants indicated that they frequently evaluate the reliability of their online search results. We got one feedback item from a participant that our introduction of the tool seemed like ‘a sales pitch’. This obviously was not our intention, but different people perceive calls to action in a different light.

The various emails in relation to data collection and the deployment of PDCareBox from our participants highlight the high levels of engagement of the PD community members. From these emails, we confirm the need for technological solutions to support those dealing with PD in their lives, which has been previously described in the literature (Wang et al., 2011, 2019; Roupia et al., 2010). Based on our experience, some of our participants were looking for explicit confirmation regarding their eligibility to join, where we would expect more experienced internet users to just try the provided internet link and the functionalities. Clearly communicating the eligibility criteria is therefore critical in the deployment of similar studies.

We tried to get announcements about our study to various private Facebook groups related to PD. Many of these groups had rules denying all kinds of advertising, including research recruiting. In addition, the concept of crowdsourcing was not familiar to most page administrators, few times confused with ‘crowdfunding’, and were not willing to discuss more. This highlights the importance of avoiding jargon when communicating outside of the wider HCI community.

According to our earlier experience on research with people with PD, having a trusted partner, e.g. a well-known PD organization, helps in this effort due to the (implicit) vetting of the study’s reliability. We approached several organizations and succeeded in finding a number of interested partners. Yet, many of the organizations never replied to any of our inquiries. Also, some organizations had decided to only cooperate with medical studies and hence excluded Computer Science-related research. Some of the organizations are staffed by a very small

<table>
<thead>
<tr>
<th>Table 8</th>
<th>Participant characteristics in our study Phase 1 as compared to Pohar and Jones (2009) and Wong et al. (2014).</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants with PD</td>
<td>Our study</td>
</tr>
<tr>
<td>Number of participants</td>
<td>288</td>
</tr>
<tr>
<td>Mean age</td>
<td>67.3</td>
</tr>
<tr>
<td>Mean years since PD diagnosis</td>
<td>5.7</td>
</tr>
<tr>
<td>Women/Men</td>
<td>57%/43%</td>
</tr>
<tr>
<td>Education: Prim./Sec./Post-sec./Univ.</td>
<td>1%/27%/21%/51%</td>
</tr>
</tbody>
</table>

| Caregivers | Our study | Wong et al. (2014) |
| Number of participants | 31 | 274* |
| Mean age | 65.7 | 69 |
| Women/Men | 74%/26% | 62%/38% |
| Work: Full time/part time/not working | 23%/13%/64% | 23%/14%/63% |

*Information is based on 274 responses by people with PD.
Three key design implications:

1. Account for diverse users: The traditional way of attempting to accommodate different types of users from an user experience perspective fall short when designing information search systems for people with complex medical conditions. Issues, such as people's felt comfort when giving advice should be considered in the user interface itself.

2. Design contextually appropriate data quality improvement methods: The intuitive improvements in e.g. text-based communication (fixing grammar, style) are useful but not enough. In our case, it became quickly clear that PD, as a use case, has various other considerations to account for, and for instance the search-ability and different ways of assessing the data are critical.

3. Provide relatable communications: Crowdsourcing-based systems that collect, organize and transfer information among the community itself have an excellent opportunity to talk to the beneficiaries in their own, relatable style.

We will discuss each of these in detail next.

6.4.1. Crowdsourcing can produce understandable and actionable data

As mentioned earlier, online information is an important source of disease-specific knowledge for people with PD (Riggare et al., 2019). However, in order to turn this available information into something usable, the provided information should be understandable. Fitzsimmons et al. (2010) assessed 100 PD web pages, concluding that none of them complied with current readability guidelines. Understandable health information facilitates shared decision-making (Odisho and Gore, 2017). Riggare et al. (2019) suggested utilizing input from patients to complement the current health information on PD, and Hartzler and Pratt (2011) similarly suggest that tools designed to share patients’ expertise with one another would benefit the users, which is exactly what we set out to do with PDCareBox. Similar to many chronic illnesses and ageing in general, PD may cause cognitive deterioration (Weintraub et al., 2008) and the understanding of complex sentences might be difficult (Goldman and Litvan, 2011). All information in PDCareBox is produced by people with PD and their caregivers, thereby written in the common language without medical jargon. In addition, the descriptions of techniques are short. An overwhelming majority of 88% of the study’s participants considered the information provided by PDCareBox as actionable and 88% as useful.

6.4.2. Improving the crowdsourced data: quality and search-ability

Even though some of our participants pointed out the possible non-reliability of peer-provided data, 78% of participants experienced the techniques provided by our tool as safe. As discussed before, some studies propose a professional moderator for peer-provided information-sharing platforms for ensuring the correctness of the data (Attard and Coulson, 2012). While having a medical professional reviewing the safety and accuracy of provided techniques might affect the user’s behavior (Visser et al., 2016), it would also require resources. In PDCareBox, all techniques are assessed via peer-users, and we could consider adding an option for flagging a technique as a possible safety risk or incorrect information when a moderator could check that technique. Peer-produced data resulted in inconsistencies in the way data is presented, including irregular capitalization and punctuation, which potentially reduces end-user trust and overall UX of PDCareBox (Lederman et al., 2014). Using relatively straightforward string operations, such issues could be reduced in the future.

The sliders in the search were experienced as confusing by some participants, while some adopted the new search strategy well. We analyzed the usability and number of usages in search for each criterion and also asked for feedback about missing criteria. We got varying...
suggestions for new criteria, along with a statement of more criteria complicating the search. In the future, we shall work with the users to find out if some of the current criteria shall be replaced with new, more relevant ones.

Morahan-Martin (2004) studied how internet users search for online health information. People typically use search engines and enter short (frequently misspelled) phrases, with the authors recommending the promotion of more effective search and evaluation techniques (Morahan-Martin, 2004). Relevance and trustworthiness of results are the most important features of a search engine (Pløtneva et al., 2012). Our participants described they often search for information on a specific topic and were hoping to see some of these search functionalities integrated into PDCareBox. Based on these suggestions, we will add an option to filter the search results according to the categories or different parameters (e.g., can be practiced at home), along with a word search (e.g., search techniques with the word ‘tremor’ in the title or description).

6.4.3. Taking into account different user types
Self-care is an integral part of daily life, and it might be difficult to identify the actions as techniques and remember all things when asked to provide self-care techniques. Some might feel uncomfortable taking a role advising others, as only 24% of our participants in Phase 1 submitted a self-care technique. However, describing what a provided technique is good for seemed to be easier, with 75% of our participants having submitted reflections. Prior work has highlighted the different social roles that people play in online health communities characterize, e.g. informational support providers often share information and advice, and informational support seekers ask questions and seek information from others (Yang et al., 2019). In discussion forums, many of the discussion threads start with a question or problem statement (Mamrykina et al., 2015). Future work may consider how different data sharing and seeking behaviors can be supported in one tool, and allow users to submit a specific question or a problem in addition to solutions (in our case, self-care techniques) — allowing for structured responses in the form of rated self-care techniques to problems raised by other users.

We acknowledge that many people with PD have difficulties in using technology, e.g. due to motor symptoms, such as tremor or rigidity, or cognitive impairment (Nunes et al., 2016). Caregivers often provide assistance in this issue, and Piper et al. (2016) suggested solutions that support collaboration and supporting needs in online presence due to symptom fluctuation, such as a ‘family account’. Community-driven tools can offer an easy way for a person with PD and their respective caregiver to sign up to (research) platforms with one account that shares relevant background information. Given these stark differences in the symptoms of people with PD, we may wish to offer users the possibility to filter and assess previously submitted and evaluated self-care techniques based on the profile of the people that have contributed to the self-care solutions.

Providing the best possible user experience for PD patients would require further end-user testing. Following general guidelines in order to enhance the accessibility of a web interface is helpful from the designer’s point of view, but these guidelines cannot take into consideration every variable that comes into play when designing and developing web interfaces. This non-comprehensiveness of the accessibility guidelines has been showcased by Remen and Svanæs (2008), and by Power et al. (2012). Therefore, the more active involvement of people with PD in the design and development of future iterations of PDCareBox is likely to be highly valuable.

6.5. Limitations
We identify several limitations in our work that are critical to consider when interpreting the presented work. First, our participant sample differs from the whole population of people diagnosed with PD and their carers. Based on our sample demographics, we identify a difference in the education level and gender share as compared with the overall PD population (Pohar and Jones, 2009). Second, participant recruitment was carried out with the help of PD organizations. Therefore, our recruitment was biased towards people that are already (active) members of one or multiple PD communities — these individuals might be more intrinsically motivated to share their experiences. Third, our participant sample consists solely of PD patients and carers who have the necessary cognitive and motor ability to participate in our study. Those who suffer from severe symptoms that obstruct internet or computer usage were unable to join without additional help. We identified a difference in the severity of PD when comparing the Modified Hoehn & Yarn scale (Goetz et al., 2004) (Table 3) and the Parkinson’s Disease Activities of Daily Living Scale (Hobson et al., 2001) (Table 4). This suggests that in the later stages of PD, the carers are the ones seeking for information on self-care. Finally, one of the four criteria on which the collected self-care techniques are ranked is ‘affordability’. However, the affordability criterion is heavily dependent on user location, as health care costs vary from country to country. For example, one of the submitted techniques suggested Deep Brain Stimulation – a surgery for alleviating symptoms. The cost for a patient to undergo Deep Brain Stimulation is likely to differ a lot depending on their respective country’s level of social security.

7. Conclusion
Parkinson’s Disease is a neurodegenerative chronic disease with a severe impact on daily life due to a wide possible range of symptoms. ‘Patient knowledge’, as gained through a lived experience of PD, provides actionable and practical information for daily life context that can be as useful in managing PD as suggestions from the medical staff. However, this important information is often available only in an unstructured form, in which actionable information is challenging to find. In this paper, we present a public, accessible, online repository of PD self-care techniques, as contributed and assessed by the global PD community. Through a real-world evaluation, we have identified practical design implications for our tool, as well as opportunities for future research opportunities for community-contributed data repositories. We furthermore identified the characteristics of people with PD who are able and willing to play an active role in the online sharing of PD knowledge, and share our experiences on reaching and interacting with this community. Through our collaboration with the PD community, we wish to further expand PDCareBox to support people in their self-care needs and practices.

CRediT authorship contribution statement

Elina Kuosmanen: Conceptualization, Formal analysis, Investigation, Resources, Writing – original draft. Eetu Huusko: Conceptualization, Software, Formal analysis, Investigation, Data curation, Writing – review & editing. Niels van Berkel: Conceptualization, Formal analysis, Investigation, Writing – review & editing. Francisco Nunes: Conceptualization, Formal analysis, Writing – review & editing. Julio Vega: Conceptualization, Formal analysis, Writing – review & editing. Jorge Goncalves: Writing – review & editing. Mohamed Khamis: Writing – review & editing. Augusto Esteves: Writing – review & editing. Denzil Ferreira: Writing – review & editing. Simo Hosio: Conceptualization, Supervision, Project administration, Funding acquisition, Resources, Investigation, Data curation.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.
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