Priority of Treatment Outcomes for Caregivers and Patients with Mild Cognitive Impairment: Preliminary Analyses

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ABSTRACT

Introduction: The patient-centered movement advocates for greater attention to the outcomes that matter most to patients and their families. In neurodegenerative disease, determination of patient and caregiver priorities has received scant attention in part because dementia patients are deemed unreliable reporters. However, people

with mild cognitive impairment (MCI) likely retain capacity to report their preferences.

Methods: In two separate MCI cohorts, we conducted preliminary analyses of patient and caregiver priorities among seven patient and five caregiver outcomes of the HABIT® Healthy Action to Benefit Independence & Thinking program (Mayo Clinic, Rochester, MN, USA).

Results: Via interview and paper-and-pencil reporting both patient and caregiver respondents’ ranked patient and caregiver quality of life and patient self-efficacy as highest priorities, ranking them ahead of patient and caregiver mood, patient functional status, patient distressing behaviors and caregiver burden. Patients and caregivers tended to value the outcomes for their loved ones higher than their own outcomes.

Conclusion: Caregivers appeared to be reasonable, but not perfect, proxies for patient reports. Additional research with larger cohorts and a more comprehensive range of outcomes is needed.

Keywords: Alzheimer’s disease; Caregivers; Mild cognitive impairment; Patient preference; Quality of life
INTRODUCTION

The patient-centered outcomes research movement has been galvanized and promoted by the Patient-Centered Outcomes Research Institute. Patient-centered outcomes research includes helping people and their caregivers consider their full range of healthcare options in terms of the risk and benefits. Patient centeredness promotes patients and caregivers making and communicating healthcare decisions informed by their personal characteristics, conditions, and preferences. One set of key preferences involves specifying outcomes that matter most to patients and caregivers [1].

Bringing patient centeredness to care and research in the area of central nervous system diseases, including neurodegenerative diseases, is complicated by concerns about patients’ capacity to state their preferences in a fully informed fashion [2]. For this reason, some have argued that using patient and caregiver-reported outcomes, such as quality of life (QoL) [3] measures, is ill-suited to research in Alzheimer’s disease (AD) or related conditions [4]. However, modern nomenclature of neurodegenerative disease diagnosis recognizes an initial phase of the illness characterized by concerning, but mild cognitive impairments (MCI) with retained functional capacities. These retained functional capacities enable an individual to establish health care preferences and participate in health care decision-making [5]. This phase of illness is labeled MCI [6–8] or mild neurocognitive disorder [9]. Patients with MCI are at high risk to progress to dementia, most often related to AD. But patients with MCI do not yet have the impairments in medical decision-making capacity seen in patients with mild dementia due to AD [5]. This status affords a special opportunity to consider outcome preferences for patients and their caregivers, including the opportunity to determine whether patient and caregiver preferences are generally consistent across the couple and whether they change with clinical intervention.

In the present pair of preliminary studies, our aim was to explore the most important behavioral outcomes for caregivers and patients with MCI.

METHODS

We used convenience cohorts assembled from the HABIT® Healthy Action to Benefit Independence and Thinking program (Mayo Clinic, Rochester, MN [10]). HABIT is a multi-component behavioral intervention program for persons with MCI and their partners. In the HABIT program, the patients are referred to as the participants. The caregivers are referred as ‘partners’ because in many cases they are not yet providing much care. Patient/participant and caregiver/partner labels are used interchangeably herein. Data collection for this program focuses on an array of patient- and caregiver-reported outcome measures (see Table 1). Patients and caregivers complete the measures listed in Table 1 just before, immediately after and at one-year intervals following the HABIT program. They are thus familiar with the constructs we were asking them to prioritize, at least as measured by these instruments. This gave them the opportunity to associate the outcomes we were asking them to rank (e.g., patient’s memory-based daily function) to a specific measure (The Everyday Cognition scale, [11]) that the patients or their caregivers had previously completed.
Table 1 Twelve outcome areas measured in Healthy Action to Benefit Independence & Thinking (HABIT)

| Outcomes patient reports about him or herself | Title of measure | Example of a question from measure |
|-----------------------------------------------|------------------|-----------------------------------|
| Depression                                    | Center for Epidemiological Studies-Depression (CES-D) [16] | I felt depressed                  |
| Quality of life                               | Quality of life [17] | Rate your life as whole           |
| Self-efficacy                                 | Self-efficacy in Mild Cognitive Impairment [18] | How confident are you that you can get your errands done despite your memory/cognitive difficulties |
| Anxiety                                       | Reach Anxiety Inventory Form [19] | I was worried                     |

| Outcomes caregiver reports about patient      | Title of measure | Example of a question from measure |
|-----------------------------------------------|------------------|-----------------------------------|
| Daily functioning                             | Functional Assessment Questionnaire (FAQ) [20] | Has problems functioning outside familiar environments |
| Memory-based activities of daily living      | Everyday Cognition (ECOG) [11] | Ability to remember things that happened recently |
| Distressing behaviors                         | The Neuropsychiatric Inventory (NPI) [21] | Frequency of agitation/aggression |

| Outcomes caregiver reports about her or himself | Title | Example of question |
|------------------------------------------------|-------|---------------------|
| Burden                                         | Zarit Burden-Short Form [22] | Do you feel that your relative asks for more help than he/she needs? |
| Self-efficacy                                  | Pearlin Mastery [23] | I can do just about anything if I really set my mind to it |
| Anxiety                                        | Reach Anxiety Inventory Form [19] | I was worried |
| Depression                                     | CES-D [16] | I felt depressed |
| Quality of life                                | Quality of life [17] | Rate your life as whole |
The first study was intended to pilot our forced-choice ranking method, so we limited the cohort to cognitively intact caregivers that had previously completed the HABIT program with their participants who had MCI. These caregivers were completing the outcome measures as part of on-going longitudinal follow-up of program outcomes. The caregivers prioritized the importance of the various constructs the program evaluation measures were intended to assess. This analysis had the additional benefit of addressing the problem that caregivers’ preferences are generally not considered when addressing priorities for, and impacts of, treatments for dementia [12]. In the second study, we sought to obtain the perspective of the patient him or herself, examine the concordance between patient and caregiver rankings, and to explore whether there were shifts in reported priorities from pre- to post-treatment. In the second study, patients with MCI and their respective caregivers (different from the first study) were surveyed. In study two, the patients and the caregivers were surveyed immediately before and immediately after completion of the HABIT program. In both studies data were stored and statistics performed using Microsoft Excel for Mac Version 14.5 (Microsoft Corp, Seattle, Washington, USA).

Study 1

Participants
Participants in the HABIT program are generally referred from the Behavioral Neurology or Neuropsychology practices at the Mayo Clinic though couples whom learn of the program via the internet can be self-referred. Patients are required to have a medical diagnosis of MCI. Patients generally met standard Mayo diagnostic criteria for MCI [7, 8]. Prior to initiating the program cognitive status is verified by administration of Dementia Rating Scale (DRS) [13]. Patients must score >115 on the DRS to participate in the program. All 36 caregiver alumni that completed the Mayo HABIT program at Mayo Clinic Minnesota or May Clinic Arizona during calendar year 2012 were candidates for the study. Three could not be reached by phone. The remaining 33 contacted via telephone consented to participate in in-person or telephone interviews to follow-up on perceived outcomes from the program. All caregivers had previously completed the set of 12 patient/caregiver-centered outcomes measures (See Table 1) routinely used to assess HABIT clinical outcomes and evaluate the HABIT program. Caregivers for the HABIT program are required to have a Mini Mental State Examination (MMSE) scores in the normal range (≥ 27) and to have regular (at least once weekly) contact with the person with MCI.

Interview
The primary author obtained consent and completed a structured interview with all caregivers. In the interview, she used the method of pairwise comparisons for rank ordering on a scale of 1 = most important to 12 = least important (see S1). It typically required more than 30 paired comparisons to determine the rank order.

Study 2

Participants
Couples attending the summer 2014 HABIT sessions at Mayo Clinic Rochester agreed to participate in the study. Rank prioritization was again on scale from 1 = most important to 12 = least important. However, for this study data were collected via paper forms rather than
in-person interview (see S2). Inclusion and Exclusion criteria were the same as study 1. Again all patients were required to have a caregiver with at least once weekly contact with the patient and MMSE ≥ 27.

Pre and post-intervention rankings
Rankings were made no more than 1 month prior to beginning of the 10-day program. Participants and partners then completed the five components of the program totaling 50 h of intervention: (1) Memory compensation training, (2) Yoga, (3) Computerized cognitive stimulation, (4) Wellness education, (5) Support group therapy. Patients and caregivers completed the rank ordering of the 12 outcomes again on the last day of the program.

Compliance with Ethics
These studies were approved by the Mayo Clinic Institutional Review Board. All procedures followed were in accordance with the ethical standards of the responsible committee on human experimentation (institutional and national) and with the Helsinki Declaration of 1964, as revised in 2013. Informed consent was obtained from all patients for being included in the study.

RESULTS
Study 1
Partner respondents were 91% female with a mean age of 71 years and 54% were college graduates. In this analysis, 87% of caregivers were spouses and the remainder were adult children of the person with MCI. The results of the rank ordering of outcome priorities of caregivers of MCI patients are depicted in Fig. 1. The patient’s QoL has the highest average ranking and was endorsed as the most important treatment outcome by 30% of caregivers. Patients' self-efficacy in handling their MCI ranked second but was comparable to patient QoL. Of note, caregivers did value their own QoL but less than they valued the patients' QoL (p < .05). The patients' daily functioning on memory-based activities was ranked in the upper half of outcomes but not as highly as QoL outcomes. Caregiver depression and burden were of least importance among all treatment outcomes, and ranked last by more than 25% of caregivers.

Study 2
All 16 couples attending the HABIT session agreed to participate. Seventy-five percent of the patient respondents were male. The mean (standard deviation) age of patients was 77.3 (7.1) and mean education was 16.3 (2.5). Caregivers were 81% female. Twelve of the caregivers were spouses, and three more were unmarried romantic partners. One caregiver was a friend. Mean age of the caregiver group was 73.1(7.5) and mean education was 14.9(1.5) years. The pre- and post-intervention average rankings for MCI patient and caregivers are presented in Fig. 2.

Pre-program Rankings
On average patients ranked their own QoL and self-efficacy first and second respectively, followed by caregiver QoL and caregiver mood. Distressing behaviors were least concerning to the patients. Caregivers ranked patient QoL, patient mood, and caregiver QoL as the top three important outcomes, respectively. Fifty percent of caregivers ranked patient QoL as their top priority.
Dependent t tests showed a trend for patients to have changed their prioritization of caregiver burden after participation in the HABIT program. This outcome moved from an average ranking of 6.5 to an average of 4.3 \( (p = .09) \) suggesting increased concern about their partner’s burden after participation in the program As a result, on average, it moved from sixth highest outcome to the highest ranking. No other changes approached statistical significance. None of the caregiver rankings changed significantly from pre- to post-intervention. There was one trend observed. Patient depression trended toward lower priority for caregivers, moving from the second highest to the sixth highest average rank \( (p = .06) \). This suggests that caregivers became less concerned about patient’s mood outcomes from baseline to end of program.

### Post-program Rankings

In comparing patient and caregiver pre-program average rankings only one significant difference emerged. Caregivers appeared more concerned about patient depression than were the patients (mean caregiver rank 4.2, mean patient rank 7.9, \( p < .01 \)). Conversely there was a trend for patients to be more concerned about caregiver burden than are the caregivers themselves (mean patient rank 6.2, mean caregiver rank 8.9, \( p = .08 \)). However, patients and caregivers agree the most important outcomes involve QoL of patients, and that distressing behaviors (hallucinations, delusions, agitation) are one of the least important (at the MCI stage).
DISCUSSION

We examined the relative importance of a variety of patient-reported and caregiver-reported outcomes to patients and caregivers that were reporting on these outcomes. The outcomes for patients included basic and memory-based daily function, anxiety, depression, distressing behaviors, self-efficacy, and QoL. We also inquired about caregiver outcomes including burden, depression, anxiety, self-efficacy, and QoL. The present results suggest that on average at the MCI stage, patient QoL and self-efficacy are most important to both patients with MCI and caregivers. These priorities seemed to hold across methods (face-to-face or phone interview, paper-and-pencil survey). Concerns about more traditional outcomes of anxiety, depression, and daily function were not as highly prioritized. These priorities may emerge in later stages of neurodegenerative disease [14]. At the MCI stage, the caregivers and patients seeking an intensive multi-component behavioral intervention are focused on patients’ opportunities to gain skills that may sustain perceived QoL and enhance self-efficacy regarding their ability to manage memory impairment.

The results also show that on average the high prioritization of QoL remains after an intensive behavioral intervention. However, the HABIT intervention completed by these patients seemed to sensitize them to the needs of their caregivers, causing their priority for caregiver burden in particular, to rise to a top concern post-treatment. Because we permitted patients and caregivers to rate both patient and caregiver outcomes, asterisked line identifies rankings that differ at $p < .01$. ADLs, activities of daily living; QoL, quality of life; Prt, participant; Self-Eff, self-efficacy.

**Fig. 2** Pre- and post-intervention rankings for patients with MCI and their caregivers (study 2). Rank ordering on a scale of 1 = most important to 12 = least important thus lower rankings equal higher priority. *Asterisked line identifies rankings that differ at $p < .01$. ADLs, activities of daily living; QoL, quality of life; Prt, participant; Self-Eff, self-efficacy.
caregiver outcomes, we seemed to observe a ‘gift of the Magi’ effect where patients and caregivers prioritized their partners outcomes over their own. Participation in the intervention program seemed to amplify this effect. Permitting patients and caregivers to prioritize their partners’ as well as their own outcomes may have led to lower estimates of concordance in this study relative to others (cf. [15]).

This analysis is limited in several respects. First, it focuses only the limited set of outcomes used in our intervention program. We chose to do this because we believed the completion of the outcome measures for the program made the respondents far more familiar with the constructs they were ranking. Had they only been prioritizing the constructs based on their labels (e.g., ‘memory-based daily functions’) there might have been more challenges understanding what the terms meant. Nevertheless, we acknowledge that a variety of traditional clinical trial outcomes including actual cognitive function and a range of biomarkers (brain volumes, amyloid levels, etc.) were not subjected to ranking. Including these traditional outcomes in future examinations of patient and caregiver outcome preferences could be a focus of future analysis.

Second, our method of rank ordering places certain constraints on how these outcomes may relate to each other. We chose this method in deference to other approaches (e.g., likert scaling) as it was very easy for participants to understand. We acknowledge that rank ordering meant different outcomes could not have equal importance to one another. All outcomes were forced to be either more or less important than those of proximal ranking. Patients and caregivers may actually perceive some of these outcomes to be equally important (or unimportant). This approach could have served to magnify differences in the rankings. Conversely, the rank ordering system limited the degree to which proximal items could differ, i.e., by only one rank unit. For example, a person could not declare that his or her highest ranked item was far more important than the second ranked item, which was only slightly more important than the third ranked item. Thus, the rank ordering may have served to diminish the differences in how these outcomes are actually valued. In any event, future studies using different methods for determining patient and caregiver outcome preferences are needed to determine the reliability of the present findings.

A final limitation is the nature of the cohorts used in these studies. They were small convenience samples. A larger sample would have provided more power to identify statistical differences between outcomes, groups, subgroups, or time points. Moreover, the sample is clearly limited in ethnic and educational diversity. It is comprised of people motivated to participate in an intensive multi-component behavioral intervention. This selection factor alone likely biases how different outcomes are viewed. However, from a PCOR research perspective the preferences of this group are exactly the preferences to identify. That is, it is a key to understand the preference of those motivated to engage actively in interventional trials to address MCI. No attempt is made in the present analyses to suggest this is a representative sample of all persons with MCI or their caregivers.

CONCLUSION

The present findings provide preliminary support for the importance of QoL and self-efficacy outcomes to patients with MCI and their caregivers. In addition, the findings
provide preliminary evidence that these preferences are fairly stable even after behavioral intervention. If these findings are supported by future research, they can provide valuable information for the design and focus of programs intended to meet the needs of patient and caregivers impacted by MCI.

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Compliance with Ethics Guidelines. These studies were approved by the Mayo Clinic Institutional Review Board. All procedures followed were in accordance with the ethical standards of the responsible committee on human experimentation (institutional and national) and with the Helsinki Declaration of 1964, as revised in 2013. Informed consent was obtained from all patients for being included in the study.

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