An Exploratory Step Toward Measuring the “Meaning of Life” in Patients with Tinnitus and in Cochlear Implant Users

Richard Tyler1  Ann Perreauf2  Anne-Mette Mohr3  Helena Ji1  Patricia C. Mancini4

1Department of Otolaryngology-Head and Neck Surgery, University of Iowa, Iowa City, IA
2Department of Communication Sciences and Disorders, Augustana College, Rock Island, IL
3Psykologcentret NV, Copenhagen NV, Denmark
4Department of Speech-Language Pathology and Audiology, Universidade Federal de Minas Gerais, Belo Horizonte, Brazil

Abstract

Background Many questionnaires attempt to quantify the “quality of life.” However, we believe understanding the quality of life is complex, and many widely used questionnaires do not capture the broad range of factors that we believe are important. Many do not include questions about communicating.

Purpose We developed a preliminary questionnaire designed to measure “The Meaning of Life” from a broader perspective.

Research Design We reviewed other scales and sought input from individuals with disabilities and developed an initial 23-item questionnaire.

Study Sample As a first step, we sampled 116 adults with tinnitus and 196 with cochlear implants (CIs). Individuals who were participating in our CI or tinnitus research programs participated.

Data Collection and Analysis To compare differences between the two participant groups, independent sample t-tests were completed for specific items on the questionnaire and for the total score. We compared age and gender differences across all participants using t-tests. Statistical significance was defined as p < 0.05. An exploratory factor analysis was conducted to examine the relationship among the questionnaire items using oblique rotation to produce correlated factors. Extracted factors with an eigenvalue >1.0 were retained according to the Kaiser-Guttman rule.

Results Four factors were prominent in this initial sample, which we labeled (1) friendship and positive outlook, (2) physical health, (3) hearing and mental health, and (4) satisfaction with life. Participants with tinnitus reported more trouble sleeping than participants with CI, whereas both groups had lower scores on hearing. Older patients reported more difficulty with remembering things but were more satisfied with their financial situation. Female participants reportedly had more hobbies and were more satisfied with their sex lives than male participants.

Conclusions This exploratory study intended to take a broader look at quality of life scales. Further work is needed with a larger sample including younger and older participants with and without disabilities.
Introduction

Quality of life scales are developed to measure individuals’ feelings of their own health and well-being. The first scale was developed by the American psychologist John Flanagan and has been adapted for use in chronic illness groups (Flanagan, 1978;10 Burckhardt and Anderson, 20033). There are currently several scales intended to measure functional ability, health status, psychological well-being, social networks and social support, and life satisfaction and morale (EuroQol Group, 1990;27 Ware and Sherbourne, 1992;33 Brazier et al, 20022). For example, the Medical Outcomes Study 36-Item Short-Form Health Survey (SF-36) was designed for use in clinical practice and research, health policy evaluations, and general population surveys (Ware and Sherbourne, 199223). This widely used questionnaire relies on patient self-reporting on eight dimensions of health: vitality, physical functioning, bodily pain, general health perceptions, physical role functioning, emotional role functioning, social role functioning, and mental health (using a magnitude estimation interval scale from 0 to 100). The SF-36 can be applied to over 200 diseases and has been translated into many languages (Garrat et al, 200211).

Derived from a selection of SF-36 items, the SF-6D is a classification for describing health (Brazier et al, 20022). It is composed of six multi-dimensional levels and describes 18,000 health states in all. The SF-6D provides a means for using the SF-36 in economic evaluation by estimating a preference-based single index measure for cost-effectiveness of health intervention analyses. Another widely used scale is the EQ-5D, a standardized instrument developed in the United Kingdom that is based on items from the SF-36 and measures health-related quality of life (EuroQol Group, 199027). This scale assesses how people perform on five dimensions of health: mobility, self-care, usual activities, pain/discomfort, and anxiety/depression (Brazier et al, 20022).

In practical terms, what exactly does “quality of life” mean? How happy are you? Can you take care of yourself or not? Do you have disabilities? Is it straightforward to compare one person’s quality of life to another? Are we not all different; do we not have different needs and expectations? Does the public know what it is like to be deaf, or unable to talk, or see clearly? Would not your expectations and interpretation of your quality of life be influenced if you were retired or not, if you lived alone or not?

Health-related quality of life scales are very important as they are used to determine the effectiveness of a treatment and to compare treatments. In addition, in some countries, governments appropriate financial resources for different health-care problems depending on the relative benefit of treating different diseases.

Therefore, research focused on health-related quality of life is exceedingly important because of the implications one’s quality of life can have on the effectiveness of current and future treatments, and on health care in a more general sense. In the development of quality of life scales, these measures are intended to reflect real-life experiences and challenges faced by everyone.

Typically, questionnaires are determined to be valid if (a) they measure what they state they are measuring, (b) they are reliable, and (c) they are sensitive (Kuk et al, 199016; Edwards, 201015). When a new questionnaire is developed, it is typically correlated to a previously published “gold standard” questionnaire to establish its validity.

Interestingly, many of the widely used scales contain no questions about hearing. Several have been applied to patients with hearing loss (Morgan et al, 200222; Stark and Hickson, 200426 Barton et al, 20051; Chia et al, 20077; Gopinath et al, 201212) and cochlear implants (CIs) (Hinderink et al, 200014; Conterra et al, 201616). Concerns from previous studies have evolved from these quality of life scales (e.g., EQ-5D and SF-6D) being unlikely to detect the expected quality of life deficits associated with hearing impairment (Morgan et al, 200222 Barton et al, 20051), for example. Therefore, some quality of life questionnaires have been developed to include questions on hearing. For example, the Health Utilities Index Mark 3 (Feeny et al, 19955) is part of a health status classification system that evaluates health on eight attributes: vision, hearing, speech, ambulation, dexterity, emotion, cognition, and pain. The World Health Organization Disability Assessment Schedule 2.0 (WHO, 201035) is a generic assessment instrument for health and disability that covers six domains of functioning: cognition, mobility, self-care, getting along, life activities, and participation. Under the cognition domain, understanding and communication are evaluated, but we note that there is only one question assessing the “general understanding of what people say” on this instrument.

Our background in audiology has led us to focus on hearing-related issues. The changes in quality of life and satisfaction after cochlear implantation have been documented before the generic health-related quality of life scales were developed (Wexler et al, 198234 Tyler and Kelsay, 199032 Maillet et al, 199518). Hinderink et al (200014) developed a handicap scale, the Nijmegen Cochlear Implantation Questionnaire, to determine quality of life changes for CI users. Three principal domains were distinguished: physical, psychological, and social (see also McRackan et al, 201721). Indeed, the National Institute on Deafness and Other Communication Disorders 2017–2021 Strategic Plan (NIH, 200823) states that there is a need for a universally accepted quality of life measure, specifically for use with CI patients, as research in quality of life has yet to produce a valid questionnaire.

This report raised our interest regarding what constitutes a valid quality of life measure. Quality of life is a broad concept that is affected by many events and circumstances. Quality of life was defined by the WHOQOL Group (199528) as:

Individuals’ perception of their position in life in the context of the culture and value systems in which they live and in relation to their goals, expectations, standards and concerns. It is a broad ranging concept affected in a complex way by the persons’ physical health, psychological state, level of independence, social relationships and their relationship to salient features of their environment.

This definition led us to believe that a quality of life questionnaire must address a greater variety of real-life issues. It is important to consider that our needs in life change
depending on our circumstances, that is, as we get older, if we
do or do not have a job, or if we have or do not have a caregiver.
Indeed, we are all different with different life experiences. Our
perception of the quality of our life might depend on our age,
our expectations, our gender, our employment status, and our
personal interests, to name just a few examples.
In this article, as an initial step, we have attempted to
address a broader range of activities related to everyday life,
for all ages. We appreciate that this is not the final product
but wanted to share our concerns about present scales and
welcome input from others. Again, with our audiology
background, we wanted to ensure the everyday difficulties
associated with hearing loss are considered.

Methods
To determine the items for inclusion in the questionnaire, we
discussed the meaning of “qualify of life” with research and
clinical professionals (audiologists, physicians, and psychol-
gists) involved in health care, and people who have a variety
of handicaps (hearing, vision, and mobility). We asked them
what areas they felt were important in their lives, and what
questions would be relevant to capture things they felt were
important in enjoying life, and what questions would best
reflect the handicaps experienced by people with disabili-
ities. As a first step, we explored these questions with
individuals who were participating in our CI or tinnitus
research programs at the University of Iowa. The preliminary
questionnaire contained 23 items (see ◄ Table 1).

The sample consisted of 312 adults, 116 of whom had
tinnitus and 196 of whom had CIs. Of the 312 participants,
149 were females (mean age = 64.9 years) and 163 were males
(mean age = 65.8 years). Participants’ ages ranged from 23 to
88 years (average age = 65.4 years); 231 were >60 years and 81
were 60 years old or younger. The mean age for the tinnitus
group was 67.6 years (standard deviation = 5.6) and for the CI
group, the mean age was 64.1 years (standard deviation = 13.5).

Some items address specific abilities related to physical
health, whereas other items relate to emotional well-being,
social interactions, and thoughts and emotions. Mean scores
were calculated for each item, and an average total score was
calculated from the average across the 23 items. To compare
differences between the two participant groups using CIs
and tinnitus, independent sample t-tests were completed for
specific items on the questionnaire and for the total score. Also, we
compared age and gender differences across all participants
using t-tests. Data were analyzed using the Statistical Package
for the Social Sciences (IBM SPSS Statistics for Windows,
Version 24.0; IBM Corp., Armonk, NY) and Microsoft® Office
Excel® 2007 (Microsoft Press, Redmond, WA). For all tests,
statistical significance was defined as \( p < 0.05 \). Finally, an
exploratory factor analysis was conducted to examine the
relationship among the questionnaire items using oblique
rotation to produce correlated factors. Extracted factors with
an eigenvalue >1.0 were retained according to the Kaisers-
Guttman rule; however, we first calculated the confidence
interval for eigenvalues and confirmed that the entire confi-
dence interval for these eigenvalues was >1.0.

Table 1 Meaning of Life Questionnaire. Participants Were Asked
to “Please Indicate Your Agreement with Each Statement on a
Scale from 0 (Completely Disagree) to 100 (Completely Agree)”

<table>
<thead>
<tr>
<th>Item</th>
<th>Statement</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>I hear well in any situation.</td>
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<tr>
<td>2.</td>
<td>I see well in any situation.</td>
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<tr>
<td>3.</td>
<td>I walk easily in any situation.</td>
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<tr>
<td>4.</td>
<td>I talk well and am easily understood.</td>
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<tr>
<td>5.</td>
<td>I sleep well.</td>
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<tr>
<td>6.</td>
<td>I manipulate things well with my hands.</td>
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<tr>
<td>7.</td>
<td>I concentrate and focus well.</td>
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<tr>
<td>8.</td>
<td>I eat and drink with ease.</td>
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<tr>
<td>9.</td>
<td>I have many friends that I socialize with.</td>
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<tr>
<td>10.</td>
<td>I always remember things.</td>
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<tr>
<td>11.</td>
<td>I have many hobbies.</td>
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<tr>
<td>12.</td>
<td>I have emotional support from many others.</td>
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<tr>
<td>13.</td>
<td>I participate in several recreational activities.</td>
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<tr>
<td>15.</td>
<td>I am satisfied with my sex life.</td>
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<tr>
<td>16.</td>
<td>I am satisfied with my financial situation.</td>
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<tr>
<td>17.</td>
<td>I feel good about my self-image.</td>
</tr>
<tr>
<td>18.</td>
<td>I am very healthy.</td>
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<tr>
<td>19.</td>
<td>I have close friends or family that I can confide in.</td>
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<tr>
<td>20.</td>
<td>In general, I get all the pleasure I want out of life.</td>
</tr>
<tr>
<td>21.</td>
<td>I think the future looks very bright.</td>
</tr>
<tr>
<td>22.</td>
<td>I never feel depressed, sad, or anxious.</td>
</tr>
<tr>
<td>23.</td>
<td>I never experience pain or discomfort.</td>
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</table>

Results

Ratings for each of the 23 items and the total score for the two
groups are presented in ◄ Figure 1. The total score revealed an
overall mean of 76.8% (standard error = 1.16) for the partici-
pants with CI and 76.9% (standard error = 1.39) for the partici-
pants with tinnitus. Item 8 (“I eat and drink with ease”) received
the highest ratings from both groups (94% for participants
with CI and 95% for participants with tinnitus), followed by
items 6 (“I manipulate things well with my hands”) and 19 (“I
have close friends or family that I can confide in”). The lowest
mean rating (48% for both groups) was found on item 1 (“I hear
well in any situation”). Items 22 (“I never feel depressed, sad, or
anxious”) and 23 (“I never experience pain or discomfort”) also
had a low mean score of 260% for participants with CI and
tinnitus. In addition, item 5 (“I sleep well”) produced the
greatest difference in scores between participants with CI
and tinnitus, with lower scores for the tinnitus group. Indepen-
dent sample t-tests were performed to determine significant
differences between participants with CI and tinnitus across
the 23 items. Ratings were significantly different (\( p < 0.05 \)) in
items 5 (“I sleep well”), 10 (“I always remember things”), 12
(“I have emotional support from many others”), 17 (“I feel good
Figure 2 shows the ratings grouped by age. Total scores averaged 76.3% for participants aged >60 years (N = 231) and 78.2% for those aged 60 years or younger (N = 81). The highest mean ratings for all ages were observed on items 8 (“I eat and drink with ease”) and 6 (“I manipulate things well with my hands”), and the lowest ratings were identified on items 1 (“I hear well in any situation”) and 23 (“I never experience pain or discomfort”). We note that the greatest difference on the mean ratings between groups was observed for item 1 (“I hear well in any situation”); the mean rating of participants aged >60 years was 44%, whereas 57% was found for those aged ≤60 years. Independent sample t-tests were performed to determine significant differences between young and old participants on the 23 items. Ratings were significantly different (p < 0.05) on item 1 (“I hear well in any situation.”), 4 (“I talk well and am easily understood”), 6 (“I manipulate things well with my hands”), 7 (“I concentrate and focus well”), 8 (“I eat and drink with ease”), 10 (“I always remember things”), 11 (“I have many hobbies”), and 16 (“I am satisfied with my financial situation”).

We also compared the ratings across male (N = 163) and female (N = 149) participants, combining CI and tinnitus...
participant groups. Figure 3 shows the mean ratings for males and females across the 23 items and for the total score. In general, females exhibited higher mean ratings than males for all items. The total scores averaged 74.8% for males and 79.0% for females. The highest mean rating for both genders was observed for item 8 (“I eat and drink with ease”), in which the mean rating for males was 93.5% and for females was 95.6%. The lowest ratings were identified for items 1 (44.7%; “I hear well in any situation”) and 23 (54.3; “I never experience pain or discomfort”). Independent sample t-tests were performed to determine if these differences between male and female participants were statistically significant. Differences were highly significant ($p < 0.05$) for items 9 (“I have many friends that I socialize with”), 11 (“I have many hobbies”), 15 (“I am satisfied with my sex life”), 20 (“In general, I get all the pleasure I want out of life”), and 21 (“I think the future looks very bright”), and for the total score.

Individual ratings for the total score were plotted against age, as shown in Figure 4. In general, lower mean ratings were found for participants aged between 55 and 75 years and higher mean ratings were reported for participants aged between 60 and 80 years.

**Factor Analysis**

A factor analysis was used to examine the shared features among items of the Meaning of Life Questionnaire. Initially, to determine whether factor analysis was adequate for the data, the Kaiser-Meyer-Olkin (KMO; SPSS v. 24.0) measure of sampling adequacy was computed. The KMO determines the proportion of variance among variables that might be

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**Fig. 3** Mean ratings of participants grouped by gender (female = gray bars; male = white bars).

**Fig. 4** Age of participants (x-axis) and individual mean ratings of total score in the Meaning of Life Questionnaire (y-axis).
common variance. Factor analysis is deemed appropriate when the KMO ratio among the observed variables is close to 1. A ratio of 0.91 was found for the set of 23 items, indicating that factor analysis was appropriate to explore the underlying factors of the questionnaire.

To further evaluate the structure of the questionnaire, an exploratory factor analysis with oblique rotation assuming correlated factors was conducted. The factor analysis revealed four factors with eigenvalues (and confidence intervals) greater than one. The first factor explained 42.3% of the total variance (eigenvalue = 9.7); the second factor explained 7.4% of the total variance (eigenvalue = 1.7); the third factor explained 6.7% of the total variance (eigenvalue = 1.5); and the fourth factor explained 4.6% of the total variance (eigenvalue = 1.1). This suggests that the 23 original items can be reduced to four underlying factors. The four factors explained 61% of the total variance in the responses from all 312 participants with CI and tinnitus.

The structure matrix contains estimates of the correlations between each of the variables and the estimated factors. To determine the shared features among the items, the proportion of variance in each item (or communality values) was first examined. Values >0.50 indicate high correlations between the items and the factors. The communality values of each variable explained by these four factors ranged from 0.55 to 0.86 (see Table 2). Analyzing the four separate factors, seven items loaded on factor 1, nine items loaded on factor 2, 5 items loaded on factor 3, and two items loaded on factor 4. Further examination of these factors revealed that items from factor 1 (9, 11, 12, 13, 19, 20, and 21) relate to friendship with others and positive outlook on one's life. For factor 2, items 2, 3, 4, 6, 7, 8, 10, 17, and 18 relate to one's physical health. For factor 3, items 1, 5, 14, 22, and 23 relate to one's hearing and mental health. For factor 4, items 15 and 16 relate to the degree of satisfaction in one's life.

After examining the statements within each factor, we propose the following labels: (a) friendship and positive outlook, (b) physical health, (c) hearing and mental health, and (d) satisfaction.

### Discussion

The purpose of this investigation was to explore the “quality of life” with a broader set of questions relevant to day-to-day activities. We considered this as a first step. With our interest in audiology, we want to ensure that the broad consequences of hearing loss are represented and that these items were sensitive to changes in hearing function. The comparison

<table>
<thead>
<tr>
<th>Factor Item</th>
<th>Statement</th>
<th>Factor Matrix</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (friendship and positive outlook)</td>
<td>9</td>
<td>I have many friends that I socialize with.</td>
</tr>
<tr>
<td></td>
<td>11</td>
<td>I have many hobbies.</td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>I have emotional support from many others.</td>
</tr>
<tr>
<td></td>
<td>13</td>
<td>I participate in several recreational activities.</td>
</tr>
<tr>
<td></td>
<td>19</td>
<td>I have close friends or family that I can confide in.</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>In general, I get all the pleasure I want out of life.</td>
</tr>
<tr>
<td></td>
<td>21</td>
<td>I think the future looks very bright.</td>
</tr>
<tr>
<td>2 (physical health)</td>
<td>2</td>
<td>I see well in any situation.</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>I walk easily in any situation.</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>I talk well and am easily understood.</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>I manipulate things well with my hands.</td>
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<td></td>
<td>8</td>
<td>I eat and drink with ease.</td>
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<tr>
<td></td>
<td>10</td>
<td>I always remember things.</td>
</tr>
<tr>
<td></td>
<td>17</td>
<td>I feel good about my self-image.</td>
</tr>
<tr>
<td></td>
<td>18</td>
<td>I am very healthy.</td>
</tr>
<tr>
<td>3 (hearing and mental health)</td>
<td>1</td>
<td>I hear well in any situation.</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>I sleep well.</td>
</tr>
<tr>
<td></td>
<td>14</td>
<td>In general, I feel very relaxed.</td>
</tr>
<tr>
<td></td>
<td>22</td>
<td>I never feel depressed, sad, or anxious.</td>
</tr>
<tr>
<td></td>
<td>23</td>
<td>I never experience pain or discomfort.</td>
</tr>
<tr>
<td>4 (satisfaction)</td>
<td>15</td>
<td>I am satisfied with my sex life.</td>
</tr>
<tr>
<td></td>
<td>16</td>
<td>I am satisfied with my financial situation.</td>
</tr>
</tbody>
</table>
between CI users and participants with tinnitus showed lowest mean ratings for both groups on item 1 (“I hear well in any situation”). Because all participants presented hearing disorders, hearing may be considered an important aspect of the meaning of life for this sample. As reported by many researchers, CIs result in a very perceptible improvement in communication and also in patients’ perception of their quality of life (Tyler and Kelsay, 1990; Maillet et al., 1995; Lin et al., 2012; Contrera et al., 2016). However, even after patients receive a CI, they often continue to report difficulties in everyday activities, such as hearing in the presence of background noise, listening to music, and using the telephone (Tyler and Kelsay, 1990; Maillet et al., 1995; Pisoni, 2000; Firszt et al., 2004). Lower ratings were also observed on items 22 (“I never feel depressed, sad, or anxious”) and 23 (“I never experience pain or discomfort”).

We know that hearing loss can contribute to social isolation, loneliness, frustration, and dependence on a caregiver. However, after patients receive a CI, they often report an improvement in these aspects that contribute to one’s quality of life (Tyler and Kelsay, 1990; Maillet et al., 1995). Maillet et al. (1995) evaluated the changes in the quality of life of CI users between preimplantation and 24 months postimplantation. They also examined the relationship between age, years of deafness, and speech recognition ability, and their satisfaction in life. The results showed that after cochlear implantation, patients had a significant and positive change in the quality of life and in their ability to communicate. In addition, the degree of perceived improvement in the quality of life was a function of the years that the patients had been deaf: patients who were deaf for #10 years had significantly greater improvement after receiving the CI than those who were deaf for $20 years (Maillet et al., 1995).

Here, participants with tinnitus were found to be more severely affected by sleeping problems than participants with CI. This result was expected because sleep disturbance has long been recognized as the single most important complaint among adults with tinnitus (Tyler and Baker, 1983; Mckenna, 2000; Mckenna and Daniel, 2006). On the item “I hear well in any situation,” participants with tinnitus scored higher than CI recipients. Tyler and Kelsay (1990) explored the disadvantages of cochlear implantation for 53 of some of the better CI patients. In this study, the main disadvantages were found to be environmental sound perception (reported by 47% of the sample), and speech perception when speechreading may be used (17%). Therefore, many CI patients have difficulty with speech perception, especially in noisy environments. Tyler and Baker (1983; see also Henry et al., 2015) reported that it is difficult for some patients with tinnitus to distinguish hearing difficulties caused by hearing loss from difficulties caused by tinnitus. This is an issue that must be addressed in individual counseling for patients with tinnitus, as recommended by Tinnitus Activities Treatment (Tyler et al., 2006; 2007). Providing a general understanding to patients with tinnitus on hearing, hearing loss, and tinnitus can help in many ways: (a) it helps patients realize they are not alone in having tinnitus, (b) it removes misperceptions and some of the fear patients may have of the unknown, and (c) it assists patients in being able to develop realistic expectations with regard to what is likely to change during tinnitus treatment (Tyler et al., 2006; 2007).

We noted that age had an influence on the ratings given by participants, especially for items 10 (“I always remember things”) and 16 (“I am satisfied with my financial situation”). Participants who were aged >60 years had lower ratings for item 10 on memory, in agreement with the common knowledge on the association between age and memory loss. Certain brain changes are observed with age, and 25% of people in the community aged >50 years report complaints of forgetfulness (Koivisto et al., 1995). On the other hand, higher scores were reported by the older age group for the latter statement (“I am satisfied with my financial situation”). One reason may rely on the fact that young people are grappling with choosing and establishing careers, finding life partners, handling the emotional stress of building a stable life, and navigating financial issues, whereas older individuals have progressed through these changes in the later years of their life.

The questionnaire showed also that men and women are different regarding their feelings of satisfaction and happiness. Females presented with higher ratings in all statements, and significant differences were observed on items from factors labeled friendship and positive outlook, and satisfaction. A number of studies have suggested that women may be more empathetic than men, and a model for gender differences in empathy has been proposed (Decety and Jackson, 2004; Rueckert and Naybar, 2008). Therefore, our study showed the contingent nature of the perception of quality of life by gender.

Questionnaires of quality of life should provide information about the impact of a disability and its treatment. We were concerned that many of the present widely used scales are not sufficiently broad, and many do not include or are sensitive to the consequences of hearing loss. We consider this only the first step and welcome comments to enhance this questionnaire. In future studies, we aim to administer the questionnaire to individuals with vision and mobility challenges. We also think it is important to consider other aspects of diversity, such as ethnicity, socioeconomic status, employment status, and interest in hobbies, which are likely to influence the questionnaire responses. For example, hearing ability is particularly important for musicians. Because musicians depend on good hearing to perform and practice, it should be reasonable to assume that hearing is an important part of musicians’ definition of their “quality of life.” If a musician has hearing loss, would this not have a greater impact on his or her quality of life than on others who are not musicians? In this way, we maintain that hearing is likely to affect one’s quality of life and quality of life assessments that are sensitive to changes in hearing function and communication abilities should be implemented in research and clinical practice.

Finally, there are some shortcomings to our study. It is likely that some CI respondents also had tinnitus.
Therefore, there is no definite distinction between the CI and the tinnitus groups. In a future study, we will explore differences between CI recipients with and without tinnitus. Overall, our research indicated that hearing ability deeply impacts the quality of life of individuals with hearing disorders. Also, differences in specific areas of quality of life were observed in the responses given by the participants when grouped by those with CIs and tinnitus, and older and younger adults. These differences demonstrate that the meaning of “quality of life” is an individual concept which depends on many circumstances. The contingent nature of quality of life evaluation makes it difficult to construct one scale that is suitable for a diverse range of people. Perhaps it is necessary to adjust the quality of life based on age, gender, and vocation.

Our purpose was to create an opportunity to investigate quality of life in a different manner and to provide a starting point to proceed in a different direction than previously done. We also hope to raise some cautionary note for all who investigate the quality of life. Indeed, we do believe that the complexity and diversity of this concept have not been fully appreciated.

**Abbreviations**

CI: cochlear implants  
KMO: Kaiser-Meyer-Olkin

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**References**

