The impact of psychosocial need and needs resolution on quality of life in patients with brain tumors

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The diagnosis of cancer of the central nervous system (CNS) is often the diagnosis of an incurable, progressive disease with devastating effects on the physical, psychosocial, and cognitive functioning of patients. Because many of the treatment options are noncurative in nature, issues related to quality and quantity of life become paramount.

The purpose of the authors' research was to explore the prevalence of psychosocial needs in this cancer population and to determine whether these needs and their resolution impact on quality of life (QOL).

Telephone interviews were conducted with 75 patients in whom primary CNS cancer was diagnosed and who were able to pass the Mini-Mental Health Status Examination. Analysis of results indicated that the majority of patients (97.3%) had at least one concern; concerns over treatment side effects, controlling uncertainty, having a meaningful existence, self and body image, and family concerns were among the five most frequently cited need domains. Most patients (91.8%) received help. However, 75.3% reported needing additional help. The number of needs reported and the severity of fatigue most significantly impacted QOL.

The study identified the needs and experiences of the patient with CNS cancer. Quality of life and needs assessment information can be used to screen patients for distress and to measure the outcome of medical and psychosocial care and ultimately to ease the burden of illness.

Key Words * brain tumor * quality of life * psychosocial need
decline, punctuated by periodic crisis, toward death. Like those with progressive neurodegenerative
diseases, individuals with cancers of the CNS have to cope with cognitive impairment, compromised
performance status, neurological deterioration, impaired communication, and personality change. These
patients share with other cancer patients the experiences of surgery, radio- and chemotherapy.

Care for these patients may include a mix of treatment for cure, for life prolongation, rehabilitative
treatment, and palliative care in varying proportions. Therapeutic interactions, when cure is not
promising, are aimed at allowing a patient to live meaningfully for as long as possible. As brain tumor
patients "struggle to relate to their environment"[21] and cope with the many devastating changes
brought on by this disease, quality of life (QOL) becomes an important concern for them.

To alleviate the burdens created by this life-threatening illness, it is critical to have information on
patients' and families' needs and QOL. The care that patients with cancer of the CNS require includes
addressing the psychosocial consequences of the disease. Although we know that psychosocial care can
be effective,[15,21,25,57,62,63] without data on needs and QOL, effective interventions are difficult to
design and evaluate. Psychosocial data are particularly useful as an outcome of care. In direct application
to medical practice, the impact of symptoms and treatment options on QOL can aid in the
decision-making process for patients, families, and healthcare providers. As well, QOL is a vital quality
assurance issue in measuring the toxic effects on the target organ.[13]

The purpose of our research is to explore the impact of the burden of psychosocial need on QOL.
Specifically, this paper will address two research questions. First, what is the prevalence of psychosocial
needs in this population of cancer patients and what are the most prominent needs? Second, do
psychosocial needs and problems in resolving need impact on QOL and, if so, what is the path of their
influence on QOL?

REVIEW OF THE LITERATURE

It is well documented in the literature that CNS cancer patients suffer from short-term memory loss,
declining strength, fatigue, loss of hearing and eyesight, seizures, cognitive deficits, altered facial
appearance, motor alterations, and changes in speech.[2,5-8,10,11,20,21,24,39,43-45,51,61] These
changes are associated with psychological and social distress that includes anxiety, depression, guilt,
uncertainty, difficult interpersonal relationships, social isolation, loss of control, loss of independence,
and negative affect.[2,4,8,11,20,23,31,40,43,45,51,60-62] These patients live a "double existence;" they
must simultaneously cope with their concerns about dying and their concerns about how long they can
continue to maintain function.[43] Many of the studies that examine the psychosocial consequences of
CNS are not empirically based. There is a rich clinical literature that describes the deleterious impact of
this disease on patients and families, often providing practical guidelines or describing psychosocial
interventions.[2,5-8,11,20,21,26,35,36,43,45,49,53,58]

The empirically based psychosocial literature on CNS cancers center on performance status and survival
as indicators of QOL[1,3,22,27,28,32,33,38,41,44,46,52,54,64] Performance status is most often assessed
using the Karnofsky Performance Scale (KPS) and cognitive functioning as indicators of QOL, which are
proxies that view QOL from a single perspective. There is, however, consensus that QOL is a
multidimensional concept that is best measured from the patient's perspective. In fact, the few available
studies that have examined the association between the KPS and QOL have found a lack of
 correspondence between the two.[41,47]
Only a few investigators have addressed subjective QOL either as a multidimensional concept or as a component of the broader concept.[30,41,42,47,48,55,59-61,66,67] Aiken[1] has stated in his review of QOL in malignant astrocytic cancers that "despite two decades of modern organized clinical trials for the treatment of malignant glioma, there is scant literature about the QOL of these patients." Two QOL questionnaires have been developed specifically for the brain tumor population that have been shown to be valid and reliable.[47,67] Both of these measures have modules attached to existing well-known QOL measures. Osoba and colleagues[47] have developed a brain tumor module to accompany the European Organization for Research and Treatment of Cancer QOL questionnaire and Weitzner and colleagues[67] have similarly developed a module to accompany the Functional Assessment of Cancer Therapy (FACT). Other measures of QOL developed specifically for brain tumor patients such as the PRESTON Profile,[40] the Aachen Life Quality Inventory,[30] and a questionnaire developed by Mackworth, Fobair, and Prados[41] have not been subjected to rigorous testing.

Although QOL has been described using a number of parameters, only two studies, to our knowledge, have used standardized QOL questionnaires that have been sufficiently evaluated. Weitzner and colleagues,[66] using the FACT, found that QOL was most affected by the extent of tumor involvement, bilateral involvement, poor performance status, aggressive treatment, by being female, divorced, and by an inability to work. Osoba and colleagues,[48] using the European Organization for Research and Treatment of Cancer QOL questionnaire, documented that recurrence, dysphasia, confusion, and motor deficit had detrimental effects on QOL and that although all were associated with decreased physical functioning, they otherwise differentially impacted on the specific dimensions of QOL. For example, confusion was associated with decreases in role-cognitive function, whereas dysphasia was associated with decreases in role and social functioning.

Other studies based their evaluations of QOL on a component of QOL such as activities of daily living (ADL)[14] or mood states.[46,61] For example, Coucair and colleagues[14] identified that ADL were associated with gender, performance status, mental status, speech impairment, motor deficit, cranial nerve deficit, and corticosteroid usage. Mackworth, et al.,[41] examined factors affecting QOL by using the well-being subscale of their own instrument as the dependent variable, regressing it on the other dimensions of the instrument. Including both children and adults, they discovered that depression and socializing had the largest impact on well-being whereas energy and reduced symptoms had less of an effect.

Another perspective on QOL has been to use a more qualitative approach. Taphoorn and colleagues[59,61] based their assessment of QOL on established mood scales and a structured interview. In a study that compared patients with a low-grade glioma and survivors of the disease, they documented that patients reported more disturbances in affective status, had more complaints about their physical condition, and were less satisfied with their overall sense of well-being.

Both the clinical and research literature illustrates the detrimental effects of CNS cancer on components of QOL. Only a few investigators have introduced standardized measures that represent a multidimensional and subjective perspective of QOL. Similarly, we were unable to find any reported literature that explicitly assessed patients' concerns and needs. Because the psychosocial aspects of CNS cancers are just beginning to be explored, there is much to learn about the QOL of these patients, as well as the need to apply this knowledge to clinical practice.

**CLINICAL MATERIAL AND METHODS**
Design and Study Population

Because an objective of the research was to verify the reliability and validity of an inventory of patients' psychosocial needs, the design was a cross-sectional one in which data were collected at two points, 1 week apart. The study population consisted of outpatients at the Princess Margaret Hospital in Toronto, Ontario, Canada, in whom a primary brain tumor had been diagnosed. Inclusion criteria were 1) age of 18 years and older, 2) ability to understand and speak English, and 3) ability to provide informed written consent. Exclusion criteria were 1) hearing problems, 2) first visit to the clinic, or 3) cognitive impairment. Cognitive impairment was assessed using the Mini-Mental Health Status Examination, and patients who scored below 25 were excluded from the study. Physicians and nurses in the clinic identified appropriate patients to approach.

Patients who met the entry criteria were enrolled between July 1996 and April 1998 and they were given verbal and written explanations prior to signing a consent form. Data were collected by means of a telephone interview that took approximately 45 minutes to complete. The number of patients who were interviewed at both time points was 75. Of the 95 eligible patients originally accrued, 15 patients dropped out of the study before the first interview. Of the remaining 80 patients, five were interviewed at Time 1 only. Of the 20 patients who either dropped out or completed Time 1 only, 45% (nine patients) were too ill to be interviewed, 25% (five patients) could not be contacted by telephone, 5% (one patient) refused to complete Time 2, and 25% (five patients) had other reasons.

Measurement Instruments

The questionnaire package consisted of five instruments and a section on demographics. The Coping in Stressful Situations scale is designed to measure three major types of coping styles: task-oriented, emotion-oriented, and avoidance coping. The Coping in Stressful Situations scale is a 48-item inventory in which individuals are asked to indicate how much they engage in specific activities, representing these three coping strategies, when they encounter a difficult or stressful situation.[17,50] Functional Assessment of Cancer Therapy with the brain tumor module is designed to measure QOL in patients with cancer.[12,67] This measure consists of the following subscales: physical, functional, social/family, and emotional well-being and relationship with doctor as well as the total score of these subscales. The relationship with doctor subscale was not used for this study because of the lack of variance in the responses to the two items that comprise this subscale.

The Fatigue Severity Scale is a nine item scale that measures the degree to which individuals with chronic conditions, including cancer, experience fatigue.[34] The Life Event Survey is designed to measure life event stress by assessing both the extent to which an event is desirable and the degree of personal impact.[56] Individuals are asked to identify events that occurred in the last 6 months and indicate the extent to which the event had a negative or positive impact on their lives.

The Princess Margaret Hospital Needs Assessment Inventory, developed by two of the investigators, is designed to identify psychosocial needs, difficulty resolving need, availability and sources of support, the need for help, barriers to need resolution, and service preferences. A qualitative methodology was used to identify a total of 58 specific needs that were then grouped into 12 domains of need: healthcare system, treatment side effects, maintenance of ADL, controlling uncertainty, self and body image, nutrition, finances and employment, family, belonging, social support, emotional distress, and meaningful existence. Standard psychometric procedures were used to test for validity and reliability. All of the instruments have established reliability and validity.
Using the literature and clinical experience as the basis for the hypothesized model, the following hypotheses are brought forward to clarify the paths of the determinants of global QOL (that is, the total score on the FACT). A diagram of the hypothesized model is shown in Fig. 1, depicting how the independent variables interact with one another and their paths of influence on QOL. For simplicity and clarity, the factors thought to influence QOL have been grouped as follows: demographic and medical characteristics, problems resolving need, burden of need, availability of resources, fatigue, and life event stress. Problems resolving need include coping strategies used to problem solve, difficulty resolving need, and number of barriers to needs resolution. Burden of need refers to number of needs and number of domains in which an individual requires assistance.

Demographic variables and life event stress are hypothesized to be contextual factors that have an indirect impact on QOL. Their influence is felt through their relationships with problems resolving need, availability of resources, and fatigue. For example, income can be seen as a means to acquire the resources necessary to decrease the burden of need and thus increase QOL. Marital status and being a member of a family unit are expected to impact on availability of support by providing the necessary help.

This disease and its side effects are believed to have both a direct and indirect impact on QOL. Indirectly, the severity of the disease will impact on fatigue, problems resolving need, and burden of need. As the disease progresses, the patient is likely to suffer more fatigue and encounter more difficulties coping that will generate a greater number of needs. Severity of disease will also have a direct effect on QOL, particularly on physical and functional well-being, which are dimensions of QOL.

Fatigue is hypothesized to have a "cloud effect." That is, as fatigue increases, an individual will have more problems resolving need, a greater burden of need, and experience a lower QOL. Studies in the literature document a substantial relationship between fatigue and QOL.[9,19,29,65] Experiencing problems resolving need is hypothesized as impacting on the burden of need and the availability of support. The more dependence on emotion-oriented and avoidance coping strategies, the greater the...
burden of need. However, the greater the use of task-oriented strategies, the less the burden of need. The more difficulties coping and barriers to needs resolution encountered by the patient, the greater the burden of need. Burden of need is hypothesized to have a direct relationship to QOL: the more needs a patient has, the lower the QOL. Last, the provision of support is hypothesized to reduce the number of needs by supplying the resources necessary to meet these needs.

RESULTS

Demographic and Medical Characteristics

As noted in Table 1, the average age of the patients was 43.6 years (range 22-76 years), there were slightly more women than men, most patients were married, and slightly more that 77.3% had been born in Canada. Although one-third of the patients were currently employed, slightly more than half were either on sick leave or long-term disability. This was a highly educated population; one-third of the patients had an university degree.

<table>
<thead>
<tr>
<th>TABLE 1</th>
<th>SUMMARY OF DEMOGRAPHIC AND MEDICAL CHARACTERISTICS IN 75 PATIENTS*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Characteristic</td>
<td>No. of Patients</td>
</tr>
<tr>
<td>age (yrs)</td>
<td>mean 43.6 SD 12.3 range 22-76</td>
</tr>
<tr>
<td>sex</td>
<td>F</td>
</tr>
<tr>
<td></td>
<td>M</td>
</tr>
<tr>
<td>married,living w/partner</td>
<td>55</td>
</tr>
<tr>
<td>living arrangements</td>
<td>spouse/partner</td>
</tr>
<tr>
<td></td>
<td>spouse/partner &amp; children</td>
</tr>
<tr>
<td></td>
<td>other nuclear,extended family</td>
</tr>
<tr>
<td></td>
<td>friend</td>
</tr>
<tr>
<td></td>
<td>alone</td>
</tr>
<tr>
<td>ages of children (yrs)</td>
<td>children &lt; 18</td>
</tr>
<tr>
<td></td>
<td>children &gt; 19</td>
</tr>
<tr>
<td></td>
<td>no children</td>
</tr>
<tr>
<td>born in Canada, yes</td>
<td>58</td>
</tr>
<tr>
<td>education</td>
<td>no diploma</td>
</tr>
<tr>
<td></td>
<td>high school diploma</td>
</tr>
<tr>
<td></td>
<td>some university/complete community college</td>
</tr>
<tr>
<td></td>
<td>university/postgraduate degree</td>
</tr>
<tr>
<td>employment status</td>
<td>paid employment</td>
</tr>
<tr>
<td></td>
<td>homemaker</td>
</tr>
<tr>
<td></td>
<td>sick leave</td>
</tr>
<tr>
<td></td>
<td>long-term disability leave</td>
</tr>
<tr>
<td>income (69 patients)</td>
<td>&lt; 29,999</td>
</tr>
<tr>
<td></td>
<td>30,000-49,999</td>
</tr>
<tr>
<td></td>
<td>50,000-79,999</td>
</tr>
<tr>
<td></td>
<td>80,000-99,999</td>
</tr>
<tr>
<td></td>
<td>≥100,000</td>
</tr>
<tr>
<td>tumor type (74 patients)</td>
<td>astrocytoma</td>
</tr>
<tr>
<td></td>
<td>glioblastoma</td>
</tr>
<tr>
<td></td>
<td>oligodendrogloma</td>
</tr>
<tr>
<td></td>
<td>medulloblastoma</td>
</tr>
</tbody>
</table>
On average, the length of time since diagnosis was 35.4 months. As shown in Table 1, astrocytoma was the most common tumor type, followed by glioblastoma and oligodendroglioma. Tumors were graded as either of low or high severity by one the investigators (J.M.); patients were almost evenly divided between the two grades.

**Psychosocial Needs**

It was the exceptional patient who was without concerns or needs: 97.3% of patients reported at least one concern (Table 2). The average number of needs was 22.4 out of a possible 58. When grouped into the 12 domains, the average number of domains was 7.2; on average patients had needs in more than half of the domains. Treatment side effects, controlling uncertainty, and meaningful existence were cited by at least three-quarters of the patients as a need (Table 2). Controlling uncertainty refers to concerns about how the cancer could impact on the future, recurrence, and death, as well as the patient's current concerns about reacting to the cancer. Meaningful existence refers to concerns over living one's life fully and taking advantage of experiences that could contribute to a meaningful existence. Other domains cited by at least 50% of the study participants included self and body image, family concerns, financial concerns, emotional distress, and maintenance of everyday activities. There was no domain that was cited by less than one-third of the patients. Although most patients with needs were getting help (67 [91.8%] of 73) and evaluated that help as positive, 75.3% of the patients indicated that they required help or desired additional help to meet their needs.
Relationship Between Psychosocial Needs and QOL

The mean scores for the FACT-Brain subscales and the total score indicate that these patients had a moderate QOL. Scores ranged between 2.8 for functional well-being to 3.2 for social/family well-being on a scale where 0 indicates a poor QOL and 4 indicates a high QOL.

In accord with the hypothesized relationships between the variables and QOL, QOL was significantly correlated with many of the hypothesized factors (Table 3). A notable exception was the demographic characteristics of the patients; none were correlated to QOL. Similarly, with the exception of months from diagnosis, none of the medical variables were related to QOL. As the patient's experience with cancer lengthened, QOL declined.
It is evident from the correlations that psychosocial needs and problems resolving those needs had substantial impact on QOL. The total number of needs as well as needs in the domains of maintenance of everyday activities, controlling uncertainty, treatment side effects, meaningful existence, and family concerns were all strongly correlated to QOL in the predicted direction: the greater the number of needs, the lower the QOL. All of the other need domains had statistically significant relationships to QOL that were moderately to somewhat weak in the same direction: the healthcare system, self and body image, nutrition, finances and employment, belonging, and social support.

Of those factors reflecting coping strategies and difficulties resolving need, difficulty resolving needs and number of domains in which help or more help was required were correlated with QOL at a moderate level. The more difficulty a patient had resolving needs and the greater the number of domains in which help was required beyond that already provided, the lower the QOL. Only the use of emotion-oriented coping strategies was correlated to QOL; the more a patient used these strategies as compared to the use of task-oriented or avoidance strategies, the lower the QOL. Number of barriers to needs resolution was also correlated to QOL, although this was a moderately weak correlation. The greater the number of barriers, the lower the QOL. Both the presence and absence of support were correlated to QOL in the expected direction, with equivalent moderate correlations. The evaluation of support, however, had a moderately weak correlation to QOL. The presence and positive appraisal of support was positively associated with QOL; the more help available and the more positive its evaluation, the higher the QOL. The greater the number of domains in which help was required, the lower the QOL.

In addition to needs assessment and demographic/medical factors, life event stress and fatigue were also examined as possible correlates of QOL. Fatigue had a strong correlation to QOL; the more severe the fatigue reported by a patient, the lower the QOL. Life event stress, an indicator of the context in which

<table>
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<tr>
<th>TABLE 3</th>
<th>Bivariate Correlations with Quality of Life</th>
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<tbody>
<tr>
<td>no. of barriers</td>
<td>0.27*</td>
</tr>
<tr>
<td>difficulty resolving needs</td>
<td>0.48†</td>
</tr>
<tr>
<td>use of emotion-oriented coping</td>
<td>0.37†</td>
</tr>
<tr>
<td>fatigue severity</td>
<td>0.59†</td>
</tr>
<tr>
<td>no. of domains in which help was received</td>
<td>0.40†</td>
</tr>
<tr>
<td>evaluation of help received</td>
<td>0.28</td>
</tr>
<tr>
<td>life event stress</td>
<td>0.31†</td>
</tr>
<tr>
<td>no. of needs</td>
<td>0.69†</td>
</tr>
<tr>
<td>no. of domains requiring help</td>
<td>0.41†</td>
</tr>
<tr>
<td>mos since diagnosis</td>
<td>0.24‡</td>
</tr>
<tr>
<td>need domains</td>
<td></td>
</tr>
<tr>
<td>healthcare system</td>
<td>0.25†</td>
</tr>
<tr>
<td>treatment side effects</td>
<td>0.61†</td>
</tr>
<tr>
<td>maintenance of ADL</td>
<td>0.69†</td>
</tr>
<tr>
<td>controlling uncertainty</td>
<td>0.68†</td>
</tr>
<tr>
<td>self and body image</td>
<td>0.31*</td>
</tr>
<tr>
<td>nutrition</td>
<td>0.35†</td>
</tr>
<tr>
<td>finances &amp; employment</td>
<td>0.31*</td>
</tr>
<tr>
<td>family</td>
<td>0.55†</td>
</tr>
<tr>
<td>belonging</td>
<td>0.31†</td>
</tr>
<tr>
<td>social support</td>
<td>0.34†</td>
</tr>
<tr>
<td>emotional distress</td>
<td>0.55†</td>
</tr>
<tr>
<td>meaningful existence</td>
<td>0.56†</td>
</tr>
</tbody>
</table>

* p ≤ 0.01.
† p ≤ 0.001.
‡ p ≤ 0.05.
the patient experiences cancer, had a weaker association with QOL; the more negatively life event stress in the past 6 months was evaluated, the lower the QOL.

**Determinants of QOL**

Although the bivariate correlations identify those factors associated with QOL, simple bivariate correlations fail to take into account the relative impact of these variables on QOL. By using multiple regression techniques, the relative impact of these variables can be calculated, the resulting model can be compared with the hypothesized model, and a model can be developed to trace the paths of those variables that impact on QOL either directly or indirectly.

The following empirical model was formulated on the basis of theoretical considerations, which are particularly important in the absence of a longitudinal design. Variables were excluded for any of the following reasons: 1) the bivariate correlations with the dependent variable were not significant at a probability level of less than 0.05, 2) multicollinearity between the independent variables, 3) the required assumptions of normal variance and linearity were not met, or 4) a variable lost its explanatory power in the multiple regression analysis.

As shown in Table 4 and the path diagram in Fig. 2, QOL was directly influenced by only two variables: fatigue and number of needs. Taking into account the indirect effects, fatigue was as influential as number of needs in its impact on QOL. Indirectly, fatigue was influential through its impact on number of needs, number of domains in which a patient received support, and emotion-oriented coping. The more severe the fatigue, the greater the number of needs and the fewer the number of domains in which support was received. At the next level of influence--help received--coping difficulties and barriers to needs resolution were influential through their impact on number of needs. The less help received, the more difficulty coping with needs and the greater the number of barriers, the more needs reported by the patient.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Beta Weights</th>
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<tbody>
<tr>
<td></td>
<td>Direct</td>
</tr>
<tr>
<td>fatigue</td>
<td>0.37</td>
</tr>
<tr>
<td>needs</td>
<td>0.54</td>
</tr>
<tr>
<td>marital status</td>
<td>—</td>
</tr>
<tr>
<td>no. of domains in which help was received</td>
<td>—</td>
</tr>
<tr>
<td>difficulty resolving needs</td>
<td>—</td>
</tr>
<tr>
<td>no. of barriers</td>
<td>—</td>
</tr>
<tr>
<td>age</td>
<td>—</td>
</tr>
<tr>
<td>emotion-oriented coping</td>
<td>—</td>
</tr>
<tr>
<td>evaluation of help received</td>
<td>—</td>
</tr>
<tr>
<td>life event stress</td>
<td>—</td>
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</table>
Each of these factors was in turn influenced by a series of variables. These variables, however, because of their greater distance from QOL, had substantially less impact. Age was both correlated to number of domains in which help was received and emotion-oriented coping. Younger patients had more domains in which they received help and were more likely to use emotion-oriented coping strategies than older patients. Emotion-oriented coping was also influenced by fatigue and number of children. However, the total effect of age of children was \( p < 0.05 \) and thus excluded from the model.

Number of barriers to needs resolution was influenced by the strength of ethnic ties, income, and evaluation of help received. Neither strength of ethnic ties nor income had sufficient impact on QOL to remain in the model. However, the stronger the tie to an ethnic group, the greater the number of reported barriers, and the higher the income, the fewer the number of reported barriers. The evaluation of help received to meet needs was of sufficient strength to remain in the model. Patients who rated the help they received as positive were more likely to indicate that they had experienced fewer barriers to needs resolution.

Lastly, life event stress was a determinant of both the number of domains in which help was received as well as barriers to needs resolution. Life event stress presents interesting results. The more negative the stress of life events was perceived to be, the fewer the barriers to needs resolution. On the other hand, the more negative the appraisal of life event stress, the less help received to meet needs.

**DISCUSSION**

In summary, with two exceptions, all of the study participants had psychosocial needs (an average of
Quality of life was significantly correlated to a number of factors. However, demographic and medical characteristics were not often correlated to QOL. This lack of correlation is substantiated in the literature. Weitzner and colleagues[66] found that the FACT scores correlated only to gender, marital status, inability to work, and extent of tumor involvement. Giovagnoli, et al.,[23] documented the lack of correlation of demographic variables and tumor involvement to QOL using the Functional Living Index--Cancer. Similarly, Mackworth, et al.,[41] found a lack of correlation between age and QOL. Although age is usually considered to be an important prognostic factor in surviving a brain tumor,[37] it is not a predictor of QOL.

When multiple regression techniques were applied, fatigue and number of needs were the only significant predictors of QOL with an r of 0.77 and an r² of 0.59. Because fatigue had substantial indirect effects, its impact on QOL was equal to that of the extent of psychosocial need. At the next level of influence, help received, difficulty coping with needs, and barriers to needs resolution had a moderate effect on QOL. The factors of importance focus on burden of need, fatigue, availability of resources, and difficulties coping. Demographic and medical characteristics play an insignificant role in influencing QOL.

When compared with the hypothesized model, the empirical model differed little from the conceptual model. In both, fatigue and burden of need were direct determinants of QOL. However, whereas medical characteristics, specifically grade of tumor, were hypothesized to be direct determinants of QOL, this did not prove to be the case in the empirical model. In addition, the hypothesis concerning the relationship of the demographic variables to the variables in the model was not borne out.

A limitation of the study is the overemphasis on patients whose cancer is not sufficiently advanced to preclude participation in the study. This would likely impact on severity of disease. Although the grade of the tumor was evenly distributed between high- and low-grade tumors, patients with serious cognitive impairment were excluded from the study. The focus of this research was on 1) the identification and magnitude of psychosocial needs, 2) the resolution of need, 3) the role fatigue plays in determining the level of need and difficulties in resolving need, and 4) the impact of these factors on QOL. This meant that a number of factors known to be determinants of QOL have not been included in this study. These include, for example, symptom severity, optimism, and locus of control.

In a population where cure or long-term remission are so often elusive, QOL becomes a major focus of care. It is important to know the patient's experience of illness, the burden, changes in lifestyle and perspective, and the positive and the negative variables. We can know this best by asking patients to tell us about their needs and QOL. As Lesley Fallowfield said, "imagining what one's quality of life might be in different states is not the same as experiencing the state."[18]

CONCLUSIONS

When needs and QOL assessment are integrated into the care of patients, QOL also becomes a goal of care. No longer is the primary focus on survival alone. Psychosocial care can be effective in helping
patients cope with the diagnosis and treatment of cancer. However, research on the needs and QOL of brain tumor patients is essential to the psychosocial care of patients, and psychosocial care is essential to the well-being of these patients. With needs assessment data, specific needs of patients can be pinpointed for intervention. In our study, treatment side effects, controlling uncertainty and meaningful existence were cited by patients as areas of need requiring intervention. Quality of life can be used as both a source of information for health care providers and as an outcome measure of medical and psychosocial care. Maintaining QOL requires a balance between the burden of illness and making resources accessible to patients to help them reduce that burden.

References


