Quality of life measurement for patients undergoing treatment for lung cancer

R J Fergusson, A Cull

The outlook for most patients with lung cancer remains bleak, with an overall five year survival rate of less than 10%. Surgery still offers the best chance of cure in the small group of patients with resectable lesions. The life span of patients with small cell tumours has been extended by combination chemotherapy but long term survival is rare. Nearly all patients are therefore treated with palliative intent.

The outcome of treatment has traditionally been measured in terms of the extent and duration of tumour response and the patient’s survival time. In the absence of major therapeutic advances in the past decade, differences between treatments as measured by these biological indices have been small and it has become increasingly relevant to compare the cost (in terms of morbidity) at which any gain is achieved. The psychosocial issues raised by the disease and its treatment have recently been comprehensively reviewed.1

Scales for monitoring toxicity related to treatment2 and performance status3-6 are widely used but inadequate for assessing some key aspects of patients’ experience—for example, pain and nausea—that are highly subjective. Furthermore, clinical experience shows that patients with apparently similar performance status and toxicity may experience substantially different quality of life. Although good clinicians consider quality of life as an important variable, most also regard it as a matter of clinical judgement: the “art” as opposed to the “science” of medicine. These assessments are unsystematic and subjectively biased and more reproducible methods are required for use in (1) auditing clinical practice, (2) evaluating treatment outcome in clinical trials, (3) informed decision making in the care of individual patients, (4) justifying needs for supportive services, and (5) allocating resources for medical services.

Vigorous research efforts by clinicians and social scientists over the past decade have resulted in a bewildering array of quality of life measures, but only by sustained collaboration can the reliability and validity of these instruments be ascertained and common problems in data collection and analysis overcome. Those unfamiliar with developments in this subject may view the task of accumulating “soft” data as cumbersome and time consuming, but considerable progress has been made

What is “quality of life”?
The concept of quality of life may be accepted as a basis for clinical decision making but it is often used with so comprehensive a meaning that it defies precise definition. There is now general agreement7 that quality of life in the context of health is a multidimensional concept concerned with the impact of physical symptoms and side effects of treatment on patients’ functioning and psychosocial well-being. Specific research questions may justify a narrower focus but studies sampling only one of these domains cannot adequately reflect the patient’s quality of life. A second point of consensus is that the emphasis should be placed on assessing the subjective experience of the person whose quality of life is in question.8

Can quality of life be measured?
It follows from this definition that indirect indicators such as time in hospital or days off work may provide valuable information about the outcome of treatment but are inadequate as measures of quality of life. What is required is an instrument that measures relevant symptoms and side effects and their impact on the patient’s physical functioning, emotional state, and social activity. The ideal method should be short, easy to administer, and easy to interpret and should have validity, adequate reliability, and responsiveness to change over time. The choice of a specific test always depends on the particular question being asked and any one “gold standard” test is unlikely to be applicable to all circumstances. Some generic health measures9,10 have been validated for use with patients suffering from cancer, but several new tools have now been developed specifically for assessing quality of life in these patients. Several excellent reviews of available instruments have been published recently.11-13 Selection of a particular method raises several questions for consideration.

Performance indices and doctors’ ratings
Patients’ performance status and the response
to treatment and its toxic effects are generally assessed by the physician. Examples of observer scales are listed in table 1. The Karnofsky performance index has been in use longest and is still widely favoured as a prognostic indicator, though inter-rater reliability is unsatisfactory and is related to the experience of the assessor. Scores are also influenced by whether the evaluation is performed at home or in hospital. The World Health Organisation and Eastern Cooperative Oncology Group (ECOG) scales share many of the same limitations. Although reliability can be improved by training raters, these instruments all neglect psychosocial variables and to that extent remain unsatisfactory as measures of quality of life.

The Carlens Vitagram was specifically developed for use with patients with lung cancer. It is based on a points system that reflects working capacity and time in hospital as well as physical symptoms and functional capacity. It is therefore a performance index. The Spitzer scale is a quick and easy measure that addresses a more comprehensive range of quality of life issues, but it has too few items in each domain for sensitive scaling.

Comparison of doctors and patients' ratings of the patient's quality of life based on several of these instruments showed poor agreement, suggesting that physicians could not accurately determine what patients felt. Furthermore, inter-rater reliability between health professionals was poor. Where possible, patients' "self report" measures should be included in the assessment of quality of life.

### Is quality of life assessment by patients really feasible?

Clinical experience and published data suggest that the overwhelming majority of patients with cancer welcome the opportunity to report their experience even within the confines of a formal research study, but care should be taken that all the questions are easy to read and interpret. Visual analogue scales have commonly been used but may be difficult for patients to understand and are time consuming to score. A questionnaire should specify the time period to which the questions refer (for example, the past 24 hours, the past three days, etc) and it is important to consider carefully how often measures need to be repeated. There may be considerable loss of compliance if the cumulative burden on the patient is too great, particularly when performance is deteriorating. With this in mind the Medical Research Committee has favoured the diary card developed for use in lung cancer as a means of obtaining frequently repeated measures of a limited number of variables over time.

The most prevalent problem reported in conducting quality of life studies has been in obtaining the cooperation of medical staff. Where it can be shown that questions relevant to the clinician can be addressed by procedures that can be accommodated in clinical practice scepticism can be overcome, but some commitment is required by a member of staff to ensure the quality control of the data collected.

### Which self report measure should be used?

Although the standardised interview conducted by a trained interviewer may yield the best data, this is expensive, time consuming, and impractical for most clinical studies, where large numbers of patients may be participating. Attention has largely focused on data collected by questionnaire and diary card.

Measures designed for use across a wide range of chronic diseases are often lengthy and hence unsuitable for studies requiring repeated testing, particularly among debilitated patients. The Psychological Adjustment to Illness Scale was originally published as a semi-structured interview with normative data for lung cancer patients covering seven quality of life domains. It is now available as a self report questionnaire with good reliability and validity but it has the disadvantage that it is copyright and expensive to obtain and, in common with other generic measures, it fails to assess specific physical symptoms or side effects.

There has been a great temptation for researchers to develop ad hoc, study specific measures with the problem that reliability and validity testing are neglected and cross study comparisons become impossible. The recommendation at this stage would be to use the best available measure rather than to seek to develop a new one. Some of the most widely used questionnaires that have been applied to quality of life assessment in patients with lung cancer are shown with brief details in table 2. The Linear Self Assessment System (LASA) appears to detect change over time but is subject to all the criticisms levelled at visual analogue scales—that is, it is time consuming to score, scores suggest a spurious degree of accuracy, and in group data patients showing equal changes in scores may not show the same evidence of improvement or deterioration. The Functional Living Index—Cancer (FLIC) is widely used but there have been problems with its format, which imposes seven equal intervals on the analogue scale. It is sometimes now presented as a seven point scale. It has relatively few items referring to symptoms and has been found somewhat insensitive to change over time. The Cancer Rehabilitation Evaluation System (CARES) is a well developed and comprehensive instrument with a short version that is more practical for research purposes. It focuses on problems amenable to rehabilitative effort and patients can identify areas where they
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Table 2  Multidimensional quality of life measures used with patients with lung cancer

<table>
<thead>
<tr>
<th>Name and reference</th>
<th>No of items and format</th>
<th>Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linear Self Assessment System (LASA)39</td>
<td>25 linear analogue</td>
<td>Symptoms, physical activity, mood, social interaction</td>
</tr>
<tr>
<td>Functional Living Index—Cancer (FLIC)²²</td>
<td>22 linear analogue</td>
<td>Symptoms, physical activity, mood, social interaction</td>
</tr>
<tr>
<td>Career Rehabilitation Evaluation System (CARES)²⁹</td>
<td>7 intervals</td>
<td>Physical, psychosocial, marital, sexual, medical, social interaction</td>
</tr>
<tr>
<td>Rotterdam Symptom Checklist (RSCL)³⁰</td>
<td>139 or 59 4 point scale + yes/no</td>
<td>Physical symptoms, psychological distress, 8 item physical function scale</td>
</tr>
<tr>
<td>EORTC core questionnaire and lung cancer module ¹¹ ¹²</td>
<td>30 + 13 4 point scale + yes/no</td>
<td>Functional state, symptoms, psychological distress, social interaction, financial state, global health and quality of life</td>
</tr>
</tbody>
</table>

EORTC—European Organisation for Research on the Treatment of Cancer

would like to have help. The main obstacle to its use in Britain is its cost. The Rotterdam Symptom Checklist (RSCL)³⁰ is a particularly clear questionnaire, which is increasingly popular because it is widely applicable and easy for patients to complete. It was recommended by the Medical Research Council’s working party on the quality of life.¹¹ Its particular advantage is its flexibility, which allows extra items to be incorporated if additional illness or treatment related variables need to be assessed.

Attention is, however, drawn to the measure developed by the quality of life study group of the European Organisation for Research on the Treatment of Cancer (EORTC). International cooperation over several years has resulted in a core questionnaire covering the generic impact of cancer and its treatment¹³ and a 13 item module¹² of items specifically relevant to patients with lung cancer—for example, cough, dyspnoea, pain, and treatment related side effects. This scale has been very carefully developed and promises to be a useful instrument for measuring quality of life in these patients. Full details of the psychometric properties of the final version of the scale are soon to be published.

Quality of life measurement in lung cancer

Before we consider the effect of treatment on quality of life it is important to remember that left untreated lung cancer has a substantial and progressive impact on patients’ quality of life. Poorer performance and more extensive disease are associated with increased psychological distress,¹⁰ and depressive illness has been reported more commonly among patients who did not have active treatment.²⁴

There are surprisingly few reports of quality of life measurements in patients undergoing treatment for lung cancer. The increasing use of chemotherapy for small cell tumours and more recently for non-small cell lung cancer has stimulated most interest in this subject.

CHEMOTHERAPY

Many early studies of chemotherapy in small cell lung cancer included assessment of functional performance as an attempt to measure the quality of life.²⁵⁻³¹ Most reported an association between improved scores and response to treatment, though one study²⁹ suggested that subsequent maintenance treatment was accompanied by a fall in performance scores. Coates et al²⁹ also attempted to assess quality of life by means of a linear analogue self-assessment scale, measuring general wellbeing and other specific factors (mood, pain, nausea, vomiting, appetite, breathlessness, and physical activity) and showed a good correlation with performance ratings on the ECOG scale.

The intensity of performance scores for detecting changes in quality of life in lung cancer was highlighted in a recent study²⁸ that compared three different instruments (diary cards, the EORTC questionnaire, and the quality of life index of Spitzer et al) in a small group of patients enrolled in a randomised trial of duration of chemotherapy in small cell lung cancer. The diary cards showed a worsening of quality of survival as treatment continued, which was not seen with the Karnofsky scores. The comparisons between the three quality of life instruments showed the appropriate convergent and divergent validity and showed that the diary cards were more sensitive to short term changes.

Non-small cell lung cancer is a less chemosensitive disease, but for the small proportion of patients who might benefit from this form of treatment would seem a rational choice as most patients have metastatic disease at presentation. In a recent review of the role of chemotherapy in advanced non-small cell lung cancer Splinter commented that few of the 142 published studies attempted to measure quality of life. In most, the response to treatment correlated with improvement in physical performance, but in others no change was seen and in some quality of life as measured by Karnofsky index fell during treatment. The study reported by Bakker et al²⁴ deserves comment. It assessed the effects of three drugs (vinodesine, cisplatin, and bleomycin) in an uncontrolled group of relatively fit patients. The authors reported a high response rate (48%), with a median survival of 47 weeks in those responding, but concluded that the fall in performance during treatment offset the benefits of treatment.

Most studies of chemotherapy in non-small
cell lung cancer show only a small survival advantage in responding patients and very few have compared active treatment with the best supportive care. As only palliation can be expected there would appear to be a strong case for including measurements of quality of life in any future studies.

**RADIOTHERAPY**

Although radiotherapy is recognised as being capable of effectively palliating symptoms in patients with lung cancer, little is known of its impact on other aspects of a patient's life. In two studies of combined chemotherapy and radiotherapy in small cell lung cancer the periods of radiotherapy were associated with a deterioration in the quality of life. Mine et al randomised 81 patients with inoperable non-small cell lung cancer to receive either radiotherapy plus chemotherapy or radiotherapy alone and used the Karnofsky index to assess the quality of survival. No difference in score between the groups was seen and there was no actual survival advantage for either treatment.

Similar results were reported by Kaasa et al and Kaasa and Masteekaas in a study of 95 patients with non-small cell lung cancer randomised to receive chemotherapy (cisplatin and etoposide) or radiography (2.8 Gy x 15). Quality of life was assessed with a locally developed questionnaire covering psychosocial wellbeing, disease and treatment symptoms, physical function, and everyday activity. The tumour response rate in the radiotherapy group (42%) was double that in the chemotherapy arm but overall survival in the two groups was identical. After two weeks of treatment there was a significant drop in performance for the patients having chemotherapy, presumably reflecting the toxicity of treatment. Subsequently there were no differences in quality of life between the two groups. Of interest was the fact that psychosocial wellbeing correlated closely with disease related symptoms (anorexia, tiredness, pain) but poorly with treatment related effects (nausea, vomiting, alopecia, dysphagia). Thus patients recognised the signs of improvement and worsening of their disease and this was reflected in their wellbeing; but they may have accepted side effects of treatment as the price to pay for a chance of overcoming their cancer, and consequently this had less impact on their overall quality of life. Coates et al reported similar results in patients with small cell lung cancer.

This important finding perhaps should be considered when we decide to withhold treatments with low activity from patients in the belief that we are protecting them from harm when there is little chance of benefit. It emphasises three important points: firstly, that quality of life measures may have a role in decisions about treatment options; secondly, that insensitive instruments, such as performance status, that only cover certain aspects of wellbeing may miss differences in the overall effectiveness of treatments; and, thirdly, that assessment of quality of life must include input from the patient.

**SURGERY**

Resection still offers the best chance of cure in patients with lung cancer, but what is their quality of survival and can the surgeon provide any palliation for patients found to be unresectable at thoracotomy? These were the questions addressed by the one major report on quality of life in patients having surgery. The authors used the Carlens Vitagram to assess the quality of survival. This method has been validated in lung cancer. The quality of survival in patients cured by resection was excellent. The patients who had an operation but subsequently died of the disease did not have a better quality of survival than non-surgically treated patients with the same stage of disease. The authors concluded that an operation had no palliative effect and the possible benefits of "reducing the tumour burden" could be dismissed. This important finding requires further verification by the more sophisticated methods now available for assessing quality of life.

**Conclusions**

Most patients with lung cancer are treated with palliative intent, where, by definition, the focus is on the quality rather than merely the duration of survival, yet relatively few studies report data on quality of life.

Substantial progress has been made in defining the concept of health related quality of life to allow agreement about what is to be assessed. An impressive range of practically useful measures has now been developed for collecting data about patients' subjective experience of disease and treatment in a reliable and valid way. Of particular interest is the modular assessment strategy, whereby a generic measure for patients with cancer can be supplemented by a standardised scale specifically relevant to lung cancer.

Work is continuing to show how quality control in collection of data on quality of life can best be achieved and to address problems that can arise in data analysis in longitudinal studies where there are problems of attrition. Sufficient progress has been made to suggest that assessment of quality of life should be included in the audit of clinical practice and evaluating treatment outcomes in clinical trials. The data obtained could provide an objective basis for informing decision making for individual patients and for making the case for allocation of appropriate resources to medical and supportive services.

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