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Quality of life during chemotherapy in lung cancer patients: results across different treatment lines

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Background: Most lung cancer patients are diagnosed at an advanced disease stage and predominantly receive palliative treatment, which increasingly consists of several chemotherapy lines. We report on patients' quality of life (QOL) to gain knowledge on QOL during and across multiple lines of chemotherapy. This includes patients with (neo)adjuvant therapy up to 3rd or above line palliative chemotherapy.

Methods: Lung cancer patients receiving outpatient chemotherapy at the Kufstein County Hospital completed an electronic version of the EORTC QLQ-C30. Linear mixed models were used for statistical analysis.

Results: One hundred and eighty seven patients were included in the study. Surprisingly, irrespective of the chemotherapy line patients reported stable QOL scores during treatment. None of the calculated monthly change rates attained clinical significance, referring to established guidelines that classify a small clinical meaningful change as 5 to 10 points. According to treatment line, 3rd or above line palliative chemotherapy was associated with the worst QOL scores, whereas patients undergoing (neo)adjuvant or 1st line palliative chemotherapy reported fairly comparable QOL.

Conclusion: The essential finding of our study is that all QOL aspects of the EORTC QLQ-C30 questionnaire remained unchanged during each chemotherapy line in an unselected population of lung cancer patients. Between treatment lines pronounced differences were found, indicating that later palliative chemotherapy lines are associated with higher QOL impairments. These changes in QOL may not primarily be related to the treatment, but rather refer to impairments due to disease progression and may be partly due to a consequence of the prior therapies.

Worldwide, lung cancer is one of the most commonly diagnosed oncological diseases and the leading cause of cancer-related death in men. In women, lung cancer ranks number four with regard to incidence but number two in terms of mortality (Ferlay *et al*, 2010). Most patients are diagnosed at an advanced stage without curative treatment options. In this situation, systemic palliative treatment has only limited effect on survival. Consequently to maintain or improve patients' quality of life (QOL) represents a main treatment goal (Petrosyan *et al*, 2012).

Owing to enhanced treatment possibilities, a growing percentage of patients nowadays are referred to multiple treatment lines. Several trials provide evidence that chemotherapy (CT) offers a benefit in advanced lung cancer patients with respect to survival and QOL (Shepherd *et al*, 2000; O'Brien *et al*, 2006; Ciuleanu *et al*, 2009). Subjectively reported QOL data favours CT over best supportive care alone, as six out of eight reviewed studies were associated with a more favourable outcome in terms of better QOL (four studies) and improved symptom control (two studies)

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(Pat *et al*, 2008). Even studies that primarily did not report a more favourable QOL outcome for CT showed positive effects of CT on specific QOL domains like functional activity (Ranson *et al*, 2000) and pain (Brown *et al*, 2005).

Although there is some knowledge about lung cancer patients' QOL undergoing palliative treatment, it has to be kept in mind that most of the studies investigated established treatment options in terms of superiority concerning patients' QOL (Dancey *et al*, 2004; Gebbia *et al*, 2010; Thongprasert *et al*, 2011) in a clinical trial setting with a highly selected patient population. Furthermore, information about patients' QOL is mostly limited to one line of treatment (predominantly 1st (Belani *et al*, 2006; Gebbia *et al*, 2010) and 2nd line CT (Dancey *et al*, 2004; Tassinari *et al*, 2009)). Although multiple lines of CT are often part of a comprehensive treatment concept, changes in patients' health status from one line to another are rarely investigated (Socinski *et al*, 2002).

Routine QOL assessments are not yet incorporated in clinical practice, although they are generally recommended (Valderas *et al*, 2008; Engelen *et al*, 2012) and known as a more reliable collection method for QOL data, as there is a considerable incongruity between clinicians' ratings and patients' self-report (Pakhomov *et al*, 2008; Basch, 2010). Hence, as part of the implementation of a computer-based QOL monitoring, our study offers important insights into QOL of lung cancer patients undergoing CT, as it presents longitudinal data with respect to multiple CT lines, assessed within the daily clinical routine of the Kufstein County Hospital. A broad range of patients receiving (neo)adjuvant and/or multiple lines of palliative CT was extensively monitored. Besides the comparison between CT lines, the pattern of changes in functioning and the course of QOL during CT were examined.

Consequently, the aims of this study are as follows:

- To compare lung cancer patients' QOL across multiple lines of CT.
- To investigate the course of patients' QOL within each line of CT.
- To compare the changes of QOL across CT lines in younger (<70 years) and older patients (≥70 years).

MATERIALS AND METHODS

Sample and procedure. At the Department of Internal Medicine at the Kufstein County Hospital, lung cancer patients were approached at the time of diagnosis to participate in routine QOL monitoring. At each visit, outpatients receiving CT were addressed to complete the QOL questionnaire EORTC QLQ-C30 (Aaronson *et al*, 1993), using a tablet PC for autonomous data entry. In case of arising questions, a study nurse was available to provide assistance. Quality of life assessment was carried out electronically by means of the Computer-based Health Evaluation System (CHES), which did not only collect and store patient-entered data but also provided the authorised doctor immediately a QOL profile for each patient. Sociodemographic and clinical data were gathered from the hospital record. The study was approved by the ethics committee of Innsbruck Medical University.

Assessment instrument

EORTC QLQ-C30. All patients completed the EORTC QLQ-C30, which is an internationally validated and widely used cancer-targeted QOL questionnaire. It includes five Functioning Scales (Physical, Social, Role, Cognitive, and Emotional Functioning), a scale for Global QOL, and nine Symptom Scales (Fatigue, Pain, Nausea/Vomiting, Dyspnoea, Appetite Loss, Sleep Disturbance,

Constipation, Diarrhoea, and Financial Difficulties). Referring to taste alterations, the QLQ-C30 was supplemented with two additional items taken from the EORTC QOL Group item bank. These items were summed up to an already previously used taste alteration subscale (higher values indicating more severe taste alterations) (Zabernigg *et al*, 2010; Giesinger *et al*, 2011; Gamper *et al*, 2012).

Statistical analysis. Patient characteristics are presented as means, standard deviations, and percentages. As linear mixed models allow data modelling with a varying number of assessments per patient and time-varying covariates (such as e.g. CT line), this modelling approach was used to compare the symptom burden and functioning between patients undergoing different CT lines. The following terms were included in the model: a random baseline, a first-order autocorrelation covariance matrix, and a fixed-effect patient group (CT line).

In a secondary analysis, we investigated physical and psychosocial symptom trajectories in terms of monthly change rates within CT lines, further including a focus on differences between younger (<70 years) and older patients (≥70 years). For interpretation of these change rates, the thresholds for minimal important change should be used. Osoba *et al* (1998) advice to use the following thresholds for the QLQ-C30: a change of 5–10 score points indicates a small clinical change, 10–20 points can be interpreted as a moderate change, and above 20 points marks a large change.

RESULTS

Patient characteristics. About 220 patients diagnosed with lung cancer and treated at the outpatient unit of the Department of Internal Medicine of the Kufstein County Hospital were approached to participate in routine PRO collection. In total, 187 patients allocated to outpatient CT were included in regular QOL assessments at each hospital visit (inclusion rate of 85%) with a total number of 996 PRO assessments. Reasons for refusal of routine QOL monitoring were severely impaired lung function, treatment only with surgery, rejection of CT by the patient, and in very few instances rejection of PRO assessment or basic language problems.

Mean age at first assessment was 69 years (s.d. 9.9), 68.5% were male, 78.7% of the patients were suffering from NSCLC, and 21.3% from SCLC. At the time of study inclusion, 16.3% of the patients received adjuvant CT, 50.6% received 1st line palliative CT, 21.9% 2nd line palliative CT, and 11.2% 3rd + line palliative CT.

In Table 1 patient characteristics are given separately for patients during different CT lines including applied treatments. During the study period, 69.5% of the patients received one CT line, 23.3% received two CT lines, and 7.2% received three or more CT lines.

At the time of study inclusion, 78.7% of patients were diagnosed with NSCLC, of which 18.4% received (neo)adjuvant CT, 74.8% 1st, 4.1% 2nd, and 2.7% 3rd line palliative CT. Regarding SCLC patients, 32.5% were undergoing (neo)adjuvant CT, 60.0% 1st, 5.0% 2nd, and 2.5% 3rd line palliative CT. Within the study period, 7.5% of the patients passed from (neo)adjuvant to 1st line palliative CT, 22.9% passed from 1st line to 2nd line palliative CT, and 7.5% from 2nd to 3rd + line palliative CT (see Figure 1).

Differences in symptom burden between CT lines. We compared the estimated patient-reported QOL scores across CT lines. As one would expect, we found a significant association between CT line and all Functioning scales of the QLQ-C30 (Physical, Role, Social, Emotional, and Cognitive Functioning), Global QOL,

Table 1. Patient characteristics

Number of patients	(Neo) adjuvant CT n = 46	1st pall. CT n = 146	2nd pall. CT n = 55	3rd + pall. CT n = 22
Age				
Mean (s.d.)	64.3 (8.7)	68.3 (10.4)	67.6 (9.1)	66.2 (8.7)
Sex				
Men	64.1%	63.9%	64.1%	47.9%
Women	35.9%	36.1%	35.9%	52.1%
Diagnosis				
NSCLC	78.8%	84.3%	90.0%	90.4%
SCLC	21.2%	15.7%	10.0%	9.6%
Time since diagnosis^a				
Mean (s.d.)	2.2 (1.3)	4.6 (7.5)	17.8 (17.0)	20.4 (13.5)
Tumour stage				
I	19.8%	3.4%	3.1%	0%
II	17.7%	0%	4.1%	0%
III	51.0%	27.5%	22.4%	7.2%
IV	11.5%	69.1%	70.4%	92.8%
Previous surgery				
Yes	43.5%	22.1%	32.1%	30.9%
Metastasis				
Yes	42.8%	69.9%	82.0%	73.6%
Chemotherapy regimen				
NSCLC monotherapy	12.3%	20.7%	82.4%	84.9%
Gemcitabine, Docetaxel, Vinorelbine, Palitaxel, Pemetrexed				
NSCLC platin combination therapy	64.4%	63.6%	7.1%	7.5%
Vinorelbine (+ antibodies), Gemcitabine Pemetrexed,				
Docetaxel, respectively + platines				
SCLC platin combination therapy	23.3%	12.8%	0.0%	3.2%
Etoposide + platines, Cisplatin/Irinotecan				
SCLC non-platin therapy	0.0%	2.9%	10.6%	4.3%
CAV, Topotecan				

Abbreviations: CT = chemotherapy; NSCLC = non-small-cell lung carcinoma; pall. = palliative; SCLC = small-cell lung carcinoma. Total N = 187. Sixty three patients were accounted for two, and 14 patients for three or more chemotherapy lines because they passed from one line to another.

^aNumber of months that passed since diagnosis, averaged across all assessments within a CT line.

Fatigue, Pain, Dyspnoea, and Appetite Loss with worse outcomes for patients in later treatment lines (see Table 2, Figure 2, and Figure 3). Interestingly, in pairwise comparisons only minor differences were detectable between patients during (neo)adjuvant and 1st line palliative treatments, respectively. Concerning Role, Social, and Cognitive Functioning, Global QOL, Fatigue, Pain, and Dyspnoea, no differences between (neo)adjuvant or 1st line palliative CT were found. In contrast, patients undergoing 2nd or 3rd + line palliative CT showed significantly worse outcomes. Specifically with regard to Physical Functioning, 3rd + line CT patients showed the most severe impairments. All in all, patients undergoing 2nd and 3rd + line palliative CT suffered from high levels of symptom burden in the majority of the QLQ-C30 subscales. However, no significant differences between CT lines were found with regard to Nausea/Vomiting, Sleeping Disturbances, Constipation, Diarrhoea, Financial Impact, and Taste Alteration.

Symptom trajectories within CT lines. According to the minimal important difference thresholds for the QLQ-C30 reported by Osoba *et al* (1998), all estimated changes for a period of on average 7.6 months (s.d. 7.3, minimum 0 and maximum 42) that reached statistical significance did not come within the category of small clinical changes (difference between 5 and 10 points) in terms of clinical significance. Social Functioning and Taste significantly worsened during adjuvant CT and 2nd line palliative CT, respectively. During adjuvant, 1st, and 2nd line palliative CT, Cognitive Functioning improved, whereas during 3rd + line palliative CT, this function area showed deterioration (for detailed numbers please see Table 3).

A further analysis considering age groups (younger patients < 70 years and older patients ≥ 70 years) and CT line yielded only one statistically significant change that was associated with an calculated difference approximating the lower limit of clinical significant changes of five points (Osoba *et al*, 1998). The monthly

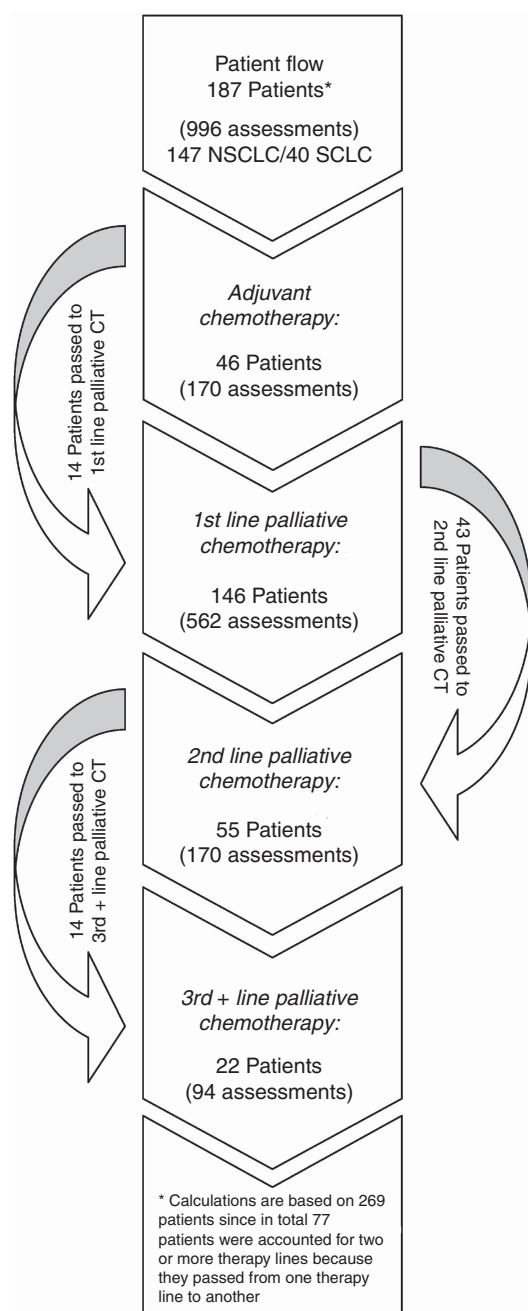


Figure 1. Patient flow. Abbreviation: CT = chemotherapy.

change rate of self-reported Constipation in older patients receiving adjuvant CT was about 3.4 points higher than that of younger patients (data not shown).

Altogether, only minor changes were detectable, which, although statistically significant, were below the usually accepted threshold of clinical significance.

Pattern of change in Functioning and Symptom Scales within age groups across CT lines. Further analyses taking age groups (younger patients <70 years and older patients ≥70 years) into account showed differences in the trend of Physical Functioning and Appetite Loss between CT groups according to their age (see Table 4). In younger patients, Physical Functioning is quite the same during (neo)adjuvant and 1st line palliative CT (mean 71.8 and 72.8 points, respectively). During 2nd and 3rd line CT, it is significantly lower than during previous lines (mean 56.3 and

53.4 points, respectively). Older patients seem to have a relatively stable Physical Functioning until a drop at the time of 3rd + line CT. A comparable age-related difference in symptom trajectory could also be true for Pain, as PRO scores show a similar pattern ($P=0.053$, narrowly missing statistical significance). Concerning Appetite Loss, we observed a substantial deterioration in younger patients between 1st and 2nd line CT with similar levels in previous (mean 16.6 points during (neo)adjuvant and 17.8 points during 1st line palliative CT) and following lines (mean 36.0 points during 2nd and 40.3 points during 3rd + line palliative CT). However in older patients, Appetite Loss decreased from 2nd to 3rd line CT, again being stable in previous lines. This particular finding may be explained by the fact that only a few patients received 3rd + line CT and therefore single ratings unduly biased the overall results.

DISCUSSION

In patients with advanced stage lung cancer, a main goal of systemic treatment is to maintain or improve QOL. Palliative CT offers the possibility to control or decrease cancer-associated symptoms (Hickish *et al*, 1998). Maintaining or improving QOL is also one of the patients' major concerns, as in burdened patients the wish for symptom relief even exceeds the wish for survival (Silvestri *et al*, 1998). This general preference in patients' treatment wishes remains basically unchanged, even though the therapeutic options considerably advanced within the past decades (e.g. emergence of maintenance CT) (Gerber *et al*, 2012). Current knowledge on lung cancer patients' QOL during CT is generally based on data provided by clinical trials including selected patient populations. Such kind of studies is mostly investigating only a single line of CT (Claassens *et al*, 2011; Damm *et al*, 2013).

In contrast, the data reported here was collected in an outpatient unit of the Kufstein County Hospital including unselected patients receiving CT for lung cancer, for what reason a broad spectrum of patients is mirrored. The most important result of our analyses is that systemic therapies on average are mainly associated with a stable QOL over time irrespective of treatment line and extent of already experienced QOL impairments. Consequently, CT itself seems not to deteriorate patients' QOL as a drop in scores mostly occurs between treatment lines. These differences between CT lines may primarily be related to impairments because of comorbidities, longer time since diagnosis, worse tumour stage (Lee *et al*, 2011), and especially disease progression.

Our data offer additional evidence that adjuvant CT has only limited negative impact on QOL, which is in line with data from a previous randomised trial (Bezjak *et al*, 2008). Only the monthly change rate of Constipation in older patients undergoing adjuvant CT approximated the lower range for a clinical significant change (Osoba *et al*, 1998) (data not shown), which is consistent with recent findings of Park *et al* (2013). As a consequence, the assumption that adjuvant CT would negatively impact the QOL in elderly patients with lung cancer needs to be questioned.

For patients receiving different lines of palliative treatment, it is necessary to keep in mind that they might suffer from an inadequate QOL deterioration in the long run. Unrecognised burden may especially occur if symptoms and QOL are not continuously monitored. Besides the encouragement of the need for close-meshed routine QOL monitoring, the data of our study also shows that according to age patients may have different needs. In patients aged 70 and younger, there seems to occur a pronounced aggravation of impairment of Physical Functioning,

Table 2. Differences between CT lines					
	Adjuvant CT (0)	1st pall. CT (1)	2nd pall. CT (2)	3rd + pall. CT (3)	P-value
Physical Functioning	69.1^{2,3}	66.4^{2,3}	55.8^{0,1}	48.0^{0,1,2}	<0.001
Role Functioning	59.8^{2,3}	55.1^{2,3}	40.0^{0,1}	34.7^{0,1}	<0.001
Social Functioning	76.9^{2,3}	76.3^{2,3}	65.5^{0,1}	59.2^{0,1}	<0.001
Emotional Functioning	71.4²	71.7²	64.5^{0,1}	68.7	0.030
Cognitive Functioning	88.2^{2,3}	86.5^{2,3}	80.2^{0,1}	77.1^{0,1}	0.001
Global QOL	57.0^{2,3}	57.6^{2,3}	48.1^{0,1}	48.7^{0,1}	<0.001
Fatigue	42.1^{2,3}	45.0^{2,3}	55.1^{0,1}	56.0^{0,1}	<0.001
Pain	21.9^{2,3}	25.0^{2,3}	36.2^{0,1}	42.7^{0,1}	<0.001
Nausea/Vomiting	8.0	10.8	12.8	10.5	0.389
Dyspnoea	33.2^{2,3}	37.2^{2,3}	50.7^{0,1}	48.8^{0,1}	<0.001
Appetite Loss	18.7^{2,3}	22.9²	36.6^{0,1}	31.7⁰	<0.001
Sleeping Disturbances	28.7	27.2	33.3	37.8	0.098
Constipation	11.8	18.9	18.1	17.7	0.249
Diarrhoea	4.9	5.8	10.4	8.4	0.059
Financial Impact	17.4	14.0	19.3	17.2	0.168
Taste Alterations	17.8	16.4	20.2	16.7	0.570

Abbreviations: CT = chemotherapy; pall. = palliative; QOL = quality of life. Statistically significant ($P < 0.05$) differences between CT lines are marked in bold, and superscript numbers indicate significant differences to CT line.

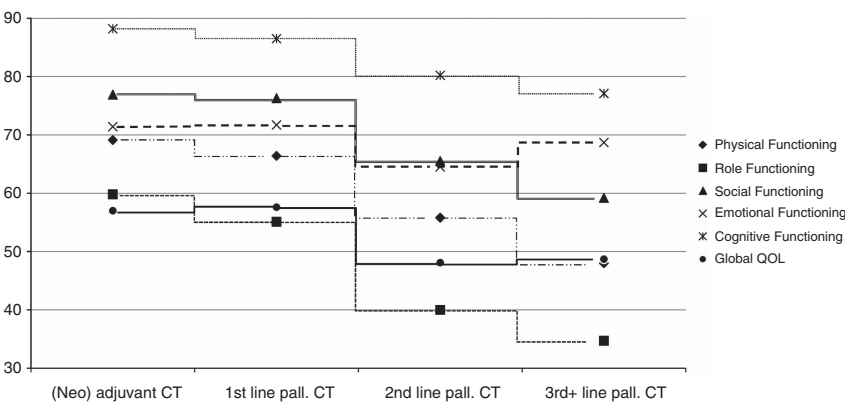


Figure 2. Differences in functioning and Global QOL between CT lines. Abbreviations: CT = chemotherapy; pall. = palliative.

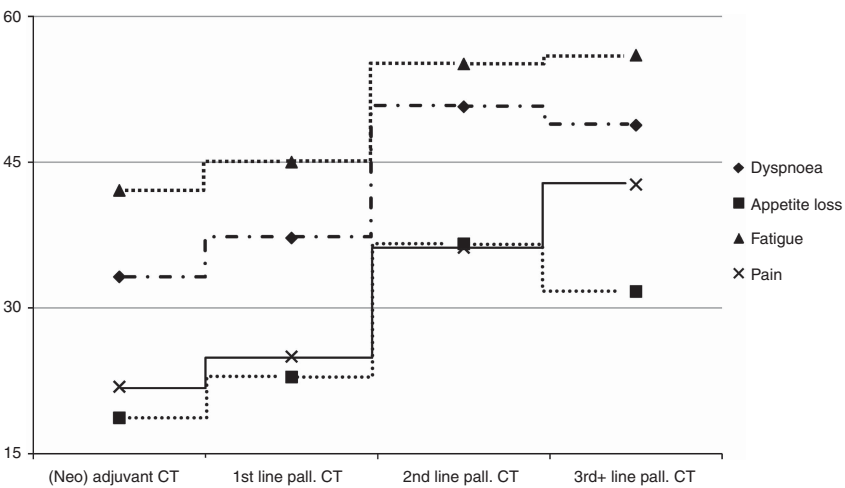


Figure 3. Differences in symptoms between CT lines. Abbreviations: CT = chemotherapy; pall. = palliative.

Pain, and Appetite Loss from adjuvant or 1st line palliative treatment to 2nd and 3rd + line palliative CT. On the other hand, patients older than 70 years appear to experience quite similar Physical Functioning and Pain during adjuvant, 1st, and 2nd line palliative CT until an increasing deterioration at 3rd + line palliative CT.

Thus, close-meshed, longitudinal QOL assessment across CT treatment lines should enable early recognition of arising symptom aggravation and consequently facilitates timely intervention in terms of drug dose modifications, offering of additional medication or psychosocial intervention. Regular QOL monitoring even outside the hospital setting has also been suggested by the results of different studies investigating the concept of maintenance treatment in 1st line therapy of patients with metastatic NSCLC. This goal of patients' QOL monitoring at home may be reached by using specialised software, as we did within our study, that can also be applied via web on several devices (computers, smart phones, tablets). Previous studies support the feasibility of web-based PROs as well as their potential to facilitate a more comprehensive medical care at hospital-free intervals at home. Web-based PRO assessments

enhance patient–clinician communication in terms of a numerically increased and more targeted discussion of symptoms (Basch *et al*, 2007a,b; Berry *et al*, 2011; Snyder *et al*, 2013).

The longitudinal assessment of QOL provides insight into patients' perception of their medical condition and associated treatment over time. Certainly, many components influence patients' reports on their QOL, as they experience a broad range of distressing symptoms depending not only on the type of cancer and its treatment but also on patients' age and sex (Deshields *et al*, 2011). Furthermore, conceptual changes due to response shift, the psychosocial situation of the patient and other factors not obvious to others but nevertheless relevant for the individual have an important role on how patients appraise their health condition and well-being. The given medical condition of a patient before the start of cytotoxic therapy influences the efficacy and patients' perception of treatment as well. Small-cell lung cancer patients who suffered from weight loss, extensive disease, and low performance status before the start of CT reported significantly worse QOL, whereas they experienced a relatively higher gain in QOL, although they did not reach QOL levels of patients with limited disease, without weight loss, and better performance status (Bernhard *et al*, 1996). Nevertheless, how valuable achieved QOL improvements are can only be determined by the patient him-/herself, and this also indicates as to why QOL needs to be assessed and discussed in clinical routine for treatment evaluation.

Our study suggests that regular QOL assessments can be effectively conducted within the busy routine of an outpatient setting. Also, patients at an advanced disease stage and with higher age contributed to the data and proved the feasibility of the used approach.

There are some limitations of the study that should be noted: related to the unselected patient group including NSCLC and SCLC patients, the shrinking sample size across CT lines, and the variety of administered cytotoxic agents, especially the reported results concerning 3rd + line CT need to be interpreted with caution. Owing to smaller groups of later treatment lines and the fact that several patients passed from one line to another, especially from 2nd to 3rd line and above, the PROs of some few patients may be over-represented and bias the analysis (e.g. as discussed within the Results section for changes in symptom pattern according to age). The decrease in the number of patients across treatment lines, however, reflects the dwindling of patients in a real-world setting, as patients receiving more than three CT lines generally are an exception. Furthermore, as it is not clear how and if the progression of patients is linked to specific treatments, the variety of CT regimen may be an additional confounder. As this study reports longitudinal QOL data across several CT lines, these constraints take a backseat, but should be considered in future studies to refine the reported results.

Table 3. Monthly change rates during chemotherapy

	Adjuvant CT	1st pall. CT	2nd pall. CT	3rd + pall. CT
Physical Functioning	− 0.5	− 0.1	− 0.3	− 0.3
Role Functioning	− 1.8	0.0	− 0.3	0.1
Social Functioning	− 1.7	0.2	− 1.1	− 0.2
Emotional Functioning	1.3	0.3	− 0.2	0.1
Cognitive Functioning	0.3	0.1	0.1	− 0.6
Global QOL	− 0.4	0.1	− 0.1	− 0.1
Fatigue	2.8	− 0.1	0.3	0.2
Pain	− 0.1	− 0.3	0.1	− 0.2
Nausea/Vomiting	2.0	0.0	0.1	0.0
Dyspnoea	2.6	0.6	0.3	0.3
Appetite Loss	0.3	− 0.3	− 0.2	− 0.3
Sleeping Disturbances	− 1.8	0.1	0.0	− 0.1
Constipation	3.0	0.1	− 0.1	− 0.3
Diarrhoea	0.0	0.2	− 0.1	0.0
Financial Impact	0.1	0.0	0.4	0.2
Taste Alterations	1.1	− 0.1	0.4	− 0.4

Abbreviations: CT = chemotherapy; pall. = palliative; QOL = quality of life. Statistically significant ($P < 0.05$) change rates within CT lines are marked in bold.

Table 4. Differences in change patterns of QLQ-C30 subscales according to age

	P-value	Age	(Neo) adjuvant CT	1st pall. CT	2nd pall. CT	3rd + pall. CT
Physical Functioning	0.008	<70	71.8	72.8	56.3	53.4
		≥70	65.2	58.7	57.5	42.4
Pain	0.053	<70	22.5	26.5	43.8	42.9
		≥70	19.9	23.6	25.8	43.1
Appetite Loss	0.027	<70	16.6	17.8	36.0	40.3
		≥70	24.0	29.1	36.8	22.3

Abbreviations: CT = chemotherapy; pall. = palliative; QOL = quality of life.

CONCLUSION

As there were no strict exclusion criteria, this sample represents the daily clinical practice in lung cancer patients receiving outpatient CT at the Kufstein county hospital. Although QOL significantly deteriorated between treatment lines, most QOL aspects remained unchanged during CT, irrespective of CT line. This means that palliative treatment *per se* did not negatively impact QOL. This information could support patients and their physician to better understand benefits and harms of the treatment. Close-meshed QOL monitoring using computer-based assessment methods offer the possibility to recognise changes in QOL at an early stage, enabling the treating physician to improve the therapeutic management.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

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