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Lung cancer and end-of-life care: a systematic review and thematic synthesis of aggressive inpatient care

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ABSTRACT

Objectives Despite recent advances in thoracic oncology, most patients with metastatic lung cancer die within months of diagnosis. Aggressiveness of their end-of-life (EOL) care has been the subject of numerous studies. This study was undertaken to evaluate the literature on aggressive inpatient EOL care for lung cancer and analyse the evolution of its aggressiveness over time.

Methods A systematic international literature search restricted to English-language publications used terms associated with aggressiveness of care, EOL and their synonyms. Two independent researchers screened for eligibility and extracted all data and another a random 10% sample of the abstracts. Electronic Medline and Embase databases were searched (2000-20 September 2018). EOLcare aggressiveness was defined as follows: 1) chemotherapy administered during the last 14 days of life (DOL) or new chemotherapy regimen during the last 30 DOL; 2) >2 emergency department visits; 3) >1 hospitalisation during the last 30 DOL; 4) ICU admission during the last 30 DOL and 5) palliative care started <3 days before death.

Results Among the 150 articles identified, 42 were retained for review: 1 clinical trial, 3 observational cohorts, 21 retrospective analyses and 17 administrative data-based studies. The percentage of patients subjected to aggressive therapy seems to have increased over time. Early management by palliative care teams seems to limit aggressive care.

Conclusions Our analysis indicated very frequent aggressive EOL care for patients with lung cancer, regardless of the definition used. The extent of that aggressiveness and its impact on healthcare costs warrant further studies.

INTRODUCTION

Lung cancer is the most common cancer worldwide, with 1.8 million new cases reported in 2012. Tobacco smoking, including second-hand and third-hand smoke exposure, remains the major risk factor for lung cancer. 1 2 The lung cancer prognosis is generally poor, with 5-year overall survival (OS) estimated at 10%–15% worldwide. In France, respective 5-year and 10-year survival rates in 2015 were 17% and 10%. Lung cancer's poor prognosis can be explained by its diagnosis at an advanced stage, despite recent progress made in screening. In 2012, about 1.6 million lung cancer deaths were reported worldwide, making it the leading cause of death due to cancer for men and the second for women after breast cancer.³⁻⁶ Over the past 10 years, management of patients with lung cancer, especially non-small cell lung cancer (NSCLC), benefited from the development of targeted therapies for patients with addictive mutation(s) and immunecheckpoint inhibitors. 7-10 Although those advances prolonged OS, the large majority of patients with lung cancer will die of the disease during the 5 years following its diagnosis.

However, those new treatments, which have increased the number of therapeutic options, render management of the end-of-life (EOL) period more complex. 11 Over the past few years, several teams examined the EOL period, specifically focusing on the factors that can negatively impact it. 11-14 Appropriate and reasonable use of active agents against solid tumours during this period poses numerous problems in clinical practice: no



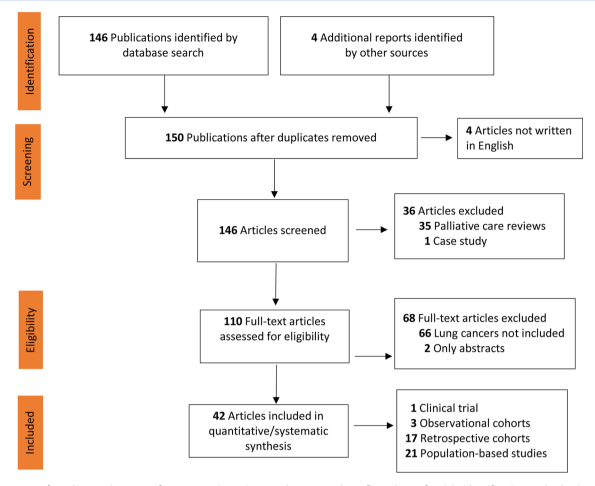


Figure 1 Preferred Reporting Items for Systematic Reviews and Meta-Analyses flow chart of article identification and selection. Other sources: references of selected articles identified four studies written in Spanish or Japanese.

precise definition of the EOL period exists, it is often difficult to accurately evaluate an individual's life expectancy and defining the concept of aggressiveness-of-care is not easy.

During the 2000s, Earle *et al*¹⁵ proposed a series of claims-based quality indicators that measure the aggressiveness of cancer care, including: intensive care unit (ICU) admission during the last months of life, repeated hospitalisations, late or no referral to palliative care and patients given chemotherapy during the last 14 days of life (DOL). The National Quality Forum (NQF) and American Society of Clinical Oncology (ASCO) jointly validated those indicators. ¹¹ 15-20 The NQF-ASCO EOL-care items have been integrated into ASCO's Quality Oncology Practice Initiative. ¹⁸ ²¹ However, data on those indicators of EOL-care aggressiveness vary widely from one country to another, one type of cancer to another and the organisation of that care.

This systematic review was conducted to synthesise all available information about aggressiveness of EOL care for those with cancer, focusing on lung cancer. The economic impact of aggressive EOL care was also evaluated.

METHODS

A systematic review was conducted and reported in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA).²²

Search strategy

The preplanned search strategy is delineated in the PRISMA flow chart (figure 1). Medline and Embase (both 2000-September 2018) databases were subjected to an exhaustive search for publications using the following terms: (aggressiveness), (Aggressive) (End of Life), (Lung cancer), (Lung neoplasm), (NSCLC), (Lung tumors), (palliative care) and (supportive care). Each full abstract was initially read and only those analysing EOL-care aggressiveness for patients with cancer were retained. That step identified 150 references. The second step was restricted to articles analysing—at least in part—lung cancers, excluding case reports and general reviews; it found 42 publications examining one or several indicators of EOL-care aggressivity of patients with lung cancer that were read in their entirety. 11 14 16 23-65

Study selection

Study abstracts, titles and full texts, when appropriate, were screened independently by two reviewers (OB

and CC) for conformity with the inclusion and exclusion criteria. A third author (FG) redid the search and 10% of the abstracts were randomly selected to see if he selected the same studies as the other two.

Inclusion criteria were: patients with lung cancer enrolled in the study and at least one of the six aggression factors validated by Earle *et al*¹⁵ was assessed. Exclusion criteria were: any literature review, congress abstracts without subsequent publications, studies that did not evaluate aggression factors and articles not published in English.

Any discrepancies unresolved by discussion between reviewers were adjudicated by a third reviewer (MD). Studies satisfying the selection criteria were retrieved and their full texts were analysed.

Data extraction

One reviewer (OB) recorded all information and available data on a dedicated data-extraction form.

Analysis

A summary, including descriptions, comparisons and limitations of each EOL-care indicator, was made. Study heterogeneity concerning populations, interventions, comparators and outcomes proscribed meta-analysis of combined data. Should another criterion of EOL-care aggressiveness be identified and reported by a substantial number of articles, it will be included in our systematic review.

RESULTS

Selected studies

Among 146 identified titles, 42 papers (1 clinical trial, 3 observational cohorts, 21 retrospective analyses and 17 administrative data-based studies) meeting the eligibility criteria are summarised in online supplementary appendix table S1.

Chemotherapy during the last 30 DOL

Because this indicator was the most studied, its impact on management is the best known (table 1). It was evaluated according to three modalities 11 15 16: the percentages of patients receiving chemotherapy during the last 14 or 30 DOL or starting a new chemotherapy line during the last 30 DOL. During our systematic search, we identified the criterion of chemotherapy during the last 30 days, which was frequently mentioned in the most recent studies, and added it to our criteria. That situation was examined in one study from 1996¹¹ on patients with solid tumours: 4.9% of patients in 1993 and 5.7% in 1996 started a new treatment line during the last 30 DOL. For this item, most publications were based on retrospective populations or analyses of administrative data. Nine papers concerned only patients with lung cancer; the others had cancers affecting various organs.

The percentages of patients given chemotherapy during the last 14 DOL ranged from 1% to 23.8%

and 6.4% to >50% received it during the last 30 DOL. Those rates also varied widely among countries with similar healthcare systems for patients with lung cancer during the last 30 DOL: 5.7% in Norway, 5.9% in Canada, 12.1% in the USA and >16% in the Netherlands and Germany. Those variations can be explained by the different healthcare systems, payment organisations and cultural and societal considerations, notably in northern Europe.

Factors associated with administering EOL chemotherapy

Young age was identified among the factors explaining a high EOL-chemotherapy rate, regardless of tumour type, in several studies. ⁵³ ⁶⁵⁻⁶⁹ By contrast, no consistent association was found among these studies between sex and tendency to administer EOL chemotherapy. ⁶⁶ ⁷⁰ Considerable differences associated with the doctors managing these patients were also found ⁶⁹ ⁷¹; the feeling of professional failure, uselessness and frustration and stress linked to their private lives impacted the pursuance of active EOL treatments. Active EOL therapies were also administered to a higher percentage of patients with lung cancer without comorbidities, with high socioeconomic status and for those with anxiety or depressive profiles. ²⁹ ⁴⁰ ⁷²

Some factors were also associated with the organisation of healthcare systems.⁴⁸ Early introduction of palliative care increased the median time between the last chemotherapy dose and death (64 vs 40.5 days; p=0.02), and patients received less chemotherapy during the last 60 DOL (52.5% vs 70.1%; p=0.05), after adjustment for age, sex and Eastern Cooperative Oncology Group performance status (ECOG PS).³⁴

Access to palliative care near the patient's home or in the hospital where the patient is managed was associated with a lower rate of active treatments during the last 30 DOL.^{66 73} However, not being provided information on palliative care units 2 weeks before death was an independent risk factor for a shorter interval between the end of chemotherapy and death.⁷⁰

Other parameters were described less frequently as being associated with near-death chemotherapy: symptomatic disease, poor ECOG PS, dying in a small hospital (<1000 beds). 66 70 On the other hand, to be included in a therapeutic trial did not seem to influence the interval between chemotherapy and death.⁵⁹ Less invasive oral treatments (targeted therapies, chemotherapies), often better tolerated, can easily be maintained at the EOL.⁷⁴ Notably, according to a prospective analysis of patients with lung cancer, oral chemotherapy was the only factor associated with the probability of receiving chemotherapy during the last 14 DOL.⁷⁵ The EOL-chemotherapy rate did not differ significantly according to treatment line⁵¹: 39% for first line, 28% for second line and 33% for third or more line.⁵¹

The impact of EOL treatment on the quality of care was examined in several studies.³⁴ 41 51 76 77 One

 Table 1
 Reports of studies on chemotherapy during the last 30 days of life (DOL)

| | | | | % Receiving chemotherapy during | |
|---------------------------------------|-------------|----------------------|------------------------------|---------------------------------|-------------|
| Study | Patients, N | Type of cancer | Inclusion criteria | Last 30 DOL | Last 14 DOL |
| Rautakorpi <i>et al</i> ²⁵ | 38 982 | All | Any | 16 | 7.2 |
| Falchook <i>et al</i> ³² | 12 765 | Lung | Patients<65 years | 12.6 | |
| Ersek <i>et al</i> ⁴² | 847 | NSCLC | Any | 30.6 | |
| Wright <i>et al</i> ¹⁴ | 1146 | All | Any | | 6.5 |
| Bekelman <i>et al</i> ⁴⁹ | 389 073 | All | USA | 10.6 | |
| | | | The Netherlands | 10.6 | |
| | | | Belgium | 12.7 | |
| | | | Canada | 8.8 | |
| | | | UK | NR | |
| | | | Germany | 10.5 | |
| | | | Norway | 4.8 | |
| Sano and Fushimi ⁴⁵ | 26 012 | All | Palliative care consultation | | 1.7 |
| | | | SOC | | 4.0 |
| Parekh <i>et al</i> ⁵¹ | 242 | NSCLC | 2009–10 Prepalliative care | | 17.0 |
| | | | 2003–14 Postpalliative care | | 7.0 |
| Pacetti <i>et al</i> ³⁹ | 2164 | All | Any | 24.3 | , |
| Nang <i>et al</i> ⁵² | 82 242 | All | 2005 | 21.5 | 4.4 |
| rrang ct ui | 02 272 | 7.111 | 2009 | | 4.4 |
| Choi <i>et al</i> ³⁰ | 263 | All | 2009 | 14.0 | 3.5 |
| CHOI Et al | 203 | All | 2002 | 42.7 | 23.8 |
| Cheung <i>et al</i> ³¹ | 107 253 | All | Any | 42.7 | 3.6 |
| Lee <i>et al³⁸</i> | 2345 | All | 2000 | 25.7 | 3.0 |
| Lee et al | 2343 | All | 2005 | | |
| | | | | 32.7 | |
| (-1+ -1 ² 7 | 200 | NCCLC | 2010 | 44.2 | F 0 |
| Kok <i>et al²⁷</i> | 266 | NSCLC | 2008–2009 | | 5.0 |
| | 440 | NICCLO | 2011–2012 | 27 | 1.0 |
| Zhang <i>et al</i> ⁶³ | 118 | NSCLC | Any | 27 | 7.6 |
| Adam <i>et al³³</i> | 120 | All | Any | 11.7 | 7.6 |
| Sesé <i>et al</i> ³⁵ | 94 | Lung | Any 55 | | 22 |
| Amano <i>et al⁵⁸</i> | 266 | All | Early palliative care | 17 | |
| 20 | | | SOC | 23 | |
| Greer <i>et al</i> ²⁹ | 151 | NSCLC | SOC | 43.3 | 23.9 |
| | | | Early palliative care | 30.0 | 13.6 |
| Mack <i>et al</i> ⁴⁶ | 1231 | NSCLC and colorectal | Any | | 16.0 |
| Näppä <i>et al⁴⁰</i> | 374 | All | Any | 23 | |
| Gonsalves <i>et al</i> ⁴⁴ | 200 | All | 2002 | 5 | |
| | | | 2008 | 18 | |
| Ho <i>et al³⁷</i> | 227 161 | All | 1993 | | 2.02 |
| | | | 2003 | | 2.9 |
| Andreis <i>et al⁵⁹</i> | 102 | All | Any | 16 | 6 |
| Warren <i>et al</i> ⁵³ | 21 633 | All | Patients with OS <6 months | 14.8 | 9 |
| | | | Patients with OS >6 months | 10.2 | 6.4 |
| Keating <i>et al</i> ⁴⁷ | 5826 | NSCLC and colorectal | VHA cohort | | 4.6 |
| - | | | SEER cohort | | 7.5 |
| Kao <i>et al⁶²</i> | 747 | All | Any | 18 | 8.0 |
| Hashimoto <i>et al</i> ⁵⁵ | 255 | All | Any | 12.6 | 3.10 |
| Tang <i>et al²⁸</i> | 242 530 | All | Trend 2000–2006 | 16.2 | |
| Barbera <i>et al</i> ³⁴ | 5855 | All | Any | | 4.6 |
| Earle <i>et al</i> ¹¹ | 215 484 | All | 1993 | | 9.70 |
| | | | 1999 | | 11.6 |
| Temel <i>et al</i> ³⁶ | 46 | NSCLC | Integrated palliative care | 40 | 23 |

Continued

| Table 1 Continued | | | | | | |
|---|--------|-------|---------------|---------------------------------|------|--|
| | | | | % Receiving chemotherapy during | | |
| Yun <i>et al</i> ⁵⁴ | 3750 | All | Any | 30.9 | 22.5 | |
| Murillo and Koeller ⁴¹ | 417 | NSCLC | Any | 43 | 20.0 | |
| Earle <i>et al</i> ¹⁶ | 28 777 | All | 1993 | | 13.8 | |
| | | | 1996 | | 18.5 | |
| Giorgi <i>et al</i> ⁵⁷ | 193 | All | Any | 33 | | |
| Emanuel <i>et al</i> ⁴⁸ | 7919 | All | Massachusetts | 9 | | |
| | | | California | 9 | | |
| Aragon-Ching <i>et al</i> ⁵⁶ | 144 | All | Any | 26 | | |

NR, not reported; NSCLC, non-small cell lung cancer; SEER, Surveillance, Epidemiology and End Results; SOC, standard of care; VHA, Veterans Health Administration.

compared three patient groups: no chemotherapy, conventional chemotherapy but not during the last 14 DOL and late chemotherapy including during the last 14 DOL. The late chemotherapy group had significantly shorter survival after starting palliative care, was hospitalised more frequently, a documented decision to stop treatment was received less often and patients died more frequently in the hospital. Although receiving chemotherapy prolonged OS by 2 months (OR 1.26, 95% CI 1.20 to 1.32; p<0.001), compared with not receiving it, no survival difference was found between conventional and late chemotherapy groups. Those authors concluded the absence of any benefit of aggressive EOL management.

Among the factors able to lower the percentage of patients receiving chemotherapy during the last 30 DOL, the impact of early access to a palliative care strategy was explored in several studies. In a monocentric, randomised trial that included 151 patients with advanced NSCLCs over 3 years, early access to palliative care declined non-significantly and lower percentages of patients were given chemotherapy during the last 30 DOL (30% vs 43.3%; p=0.14) and last 14 DOL (13.6% vs 23.6%, p=0.18). The failure to reach significance probably reflects the too small number of patients recruited. Despite the lighter therapeutic pressure during the EOL period in that study, an OS benefit was observed for the early palliative care group (11.6 vs 8.9 months, p=0.02).

Similarly, an analysis of Japanese administrative data from patients with cancer aged >65 years showed that early access to palliative care consultations lowered, from 4% to 1.5%, the number of patients receiving chemotherapy doses during the last 14 DOL. The contrast, again in Japan, the use of an aggregated aggressiveness-of-care criterion at the EOL to compare 54 patients seen in palliative care consultations >3 months before their deaths vs 211 patients without such appointments revealed improved quality of life but did not significantly lower the rate of patients who had received chemotherapy during the last 14 DOL; that lack of significance could also be attributable to the small sample size. See

Repeated emergency department visits during the last 30 DOL

This criterion was analysed in 38% of the articles retained (table 2) and concerned 1.8%–66.9% of the patients. Three studies, one prospective, addressed only patients with lung cancer. ^{36 65 78} Early access to palliative care seemed to lower the number of repeated ED consultations. Temel *et al*, based on 46 patients, evaluating early palliative care management versus standard of care (SOC) found the former (10%) had fewer repeated ED visits. ³⁶ That access in global management seems to allow patients and their entourage time to reflect on the prognosis, disease evolution and EOL organisation.

Factors associated with ED visits during the last 30 DOL

A meta-analysis of ED visits during the last 30 DOL included 30 studies and 1 181 842 patients with cancer (including haemopathies). Factors significantly associated with ED visits were male sex (OR 1.24, 95% CI 1.19 to 1.29), African origin (OR 1.45, 95% CI 1.40 to 1.50), lung cancer (OR 1.17, 95% CI 1.10 to 1.23) and low socioeconomic level (OR 1.15, 95% CI 1.10 to 1.19). In contrast, patients receiving palliative care had fewer ED consultations (OR 0.43, 95% CI 0.36 to 0.51]).

Among symptoms associated with ED visits, dyspnoea and pain were the most common. The percentage of EOL patients consulting the ED seems to have increased over time. In a population-based, retrospective, cohort study using 1993–2003 administrative data from 227 161 patients with cancer, multiple ED visits increased significantly over time from 8.60% to 10.53%. Factors associated with that rise were young age, male sex and rural residence.

Study results also suggest that many emergency medicine specialists feel underqualified to manage these EOL patients, which might explain the high number of ED visits. 82–86

Repeated hospitalisations during the last 30 DOL

This indicator was analysed in 11 (26%) of the 42 articles retained for this synthesis (table 3). As a function of the study and country, 3.6% to >50% of the patients

 Table 2
 Studies reports of repeated end-of-life emergency department (ED) consultation

| Study | Patients, N | Type of cancer | Inclusion criteria | ≥1 ED visit (%) |
|-------------------------------------|----------------|----------------------|-----------------------------------|-----------------|
| Ersek <i>et al</i> ⁴² | 847 | NSCLC | Any | 66.90 |
| Falchook <i>et al</i> ³² | 12 765 | Lung | Patient <65 years | 1.8 |
| Wu et al ²³ | 49 920 | All | Palliative chemotherapy <6 months | 34 |
| | | | Any chemotherapy | 13.7 |
| Henson <i>et al</i> ⁴³ | 681 | All | Any | 29.70 |
| Wright <i>et al</i> ¹⁴ | 1146 | All | Any | 7.90 |
| Wang <i>et al</i> ⁵² | 82 242 | All | 2005 | 34 |
| | | | 2009 | 32.90 |
| Choi <i>et al</i> ³⁰ | 263 | All | 2002 | 22.8 |
| | | | 2012 | 74.8 |
| Cheung <i>et al</i> ³¹ | 107 253 | All | Any | 14.7 |
| Amano <i>et al</i> ⁵⁸ | 266 | All | Early palliative care | 11 |
| | | | SOC | 4.70 |
| Mack et al ⁴⁶ | 1231 | NSCLC and colorectal | Any | 40 |
| Ho <i>et al</i> ³⁷ | 22 761 | All | 1993 | 8.60 |
| | | | 2003 | 10.50 |
| Gonsalves et al ⁴⁴ | 200 | All | 2002 | 6 |
| | | | 2008 | 9 |
| Keating <i>et al</i> ⁴⁷ | 5826 | NSCLC and | VHA cohort | 13.10 |
| | | colorectal | SEER cohort | 14.70 |
| Tang <i>et al</i> ²⁸ | 242 530 | All | Trend 2000– 2006 | 18.3 |
| Temel <i>et al</i> ³⁶ | 46 | NSCLC | Integrated palliative care | 10 |
| Barbera <i>et al</i> ³⁴ | 5855 | All | Any | 32.20 |
| Earle et al ¹⁶ | 28 777 | All | 1993 | 7.20 |
| | | | 1996 | 9.20 |

ED, emergency department; NSCLC, non-small cell lung cancer; SEER, Surveillance, Epidemiology and End Results; SOC, standard of care; VHA, Veterans Health Administration.

were concerned. For example, 59.3% of 12 764 patients with lung cancer aged <65 years were included in analyses based on American medical administrative data.³²

This dimension of the quality of care during the EOL period was also evaluated with patient hospitalisations for >14 days. That rate ranged from 12% in the USA to 42% in Taiwan. However, that indicator is difficult to interpret because it is linked to care facilities and payment modalities.¹¹

ICU admissions during the last 30 DOL

This parameter was analysed in 20/42 (48%) of the retained studies and affected 2%-30% of the subjects included (table 4). That rate seems to have increased over time, ^{16 34 37} rising from 6% in 2002 to 33% in 2008 (p=0.001) in a study comparing 100 patient deaths during each year in the same institution.

Using the Surveillance, Epidemiology and End Results (SEER)-Medicare data from 132 051 patients who died of cancer in 2006–2011, ICU admissions rose from 16.2% to 21.3% (p<0.001).⁵² Again, based on 1992–2005 SEER data, more patients with lung cancer were admitted to the ICU.⁸⁷ Among 175 756 patients with lung cancer entered in the SEER database and hospitalised for a reason other than surgical tumour resection during the study period, 49 373 (28%) had at least one ICU stay. The rate of ICU admissions per 1000 hospitalisations increased over the study period from 140.7 in 1992 to 201.7 in 2005 (p<0.001).

Factors associated with ICU admission

Factors associated with ICU admission were young age (OR 0.80 for age 70–74 years, 95% CI 0.77 to 0.83 vs OR 1.0 for age <65 years), male sex (OR 1.10, 95% CI 1.07 to 1.12), wealthier patients (OR 1.50, 95% CI 1.42 to 1.57) and localised disease (OR 0.61 for metastatic disease, 95 CI 0.59 to 0.63). Their ICU admissions increased especially for the category 'intermediate ICU', with a risk ratio of 3.09 (95% CI 2.78 to 3.44) between 1992 and 2005.

Those factors were also found for a cohort of 412 946 patients with NSCLC entered in the National Institutes of Health database between 1998 and 2014, with ICU admissions rising from 13.3% in 1998 to 27.9% in 2014 (p<0.01).⁶¹ A multivariate logistic-regression model retained the following factors as independent predictors of ICU care: age <65 years, racial minority (vs white), higher Charlson Comorbidity Index, Medicare or Medicaid insurance (vs private insurance), teaching hospitals and medium or large hospitals (vs small hospitals).

The higher rate of ICU admissions during the last 30 DOL has not been accompanied by an increased number of patients on mechanical ventilation, which has either stabilised or decreased (table 4).²⁸ The chemotherapy administration during the last 30 DOL enhances the risk of ICU admission during that period⁶⁴ can be explained by the appearance of new, more effective, therapeutic strategies, which incite doctors to raise the therapeutic pressure. The identification of pejorative factors of short-term evolution should enable better definition of the prognoses of these patients and limit unnecessary ICU admissions. The impact of these ICU admissions on the patient's entourage was examined in a large study based on >1000 interviews with families that had lost a member to cancer.¹⁴ Their satisfaction was significantly lower when the patient had been admitted to the ICU during the last 30 DOL, than otherwise. In every case, explanation and understanding of medical decisions by the family improved their satisfaction.

The ICU admission rate was lower, ranging from 1% to 9%, when patients were managed in structures that have implemented a policy of early access to palliative care. ²⁶ ⁴⁶ In their prospective randomised trial, Temel *et al* compared standard management with earlier

Table 3 Study reports of repeated end-of-life hospitalisations ≥1 hospital admissions >14 days of hospitalisation Patients, Study Ν Type of cancer Inclusion criteria (%) (%) Falchook et al³² 12 765 Luna Patient <65 years 59.30 Wu et al²³ 49 920 All Palliative chemotherapy <6 months 22.8 42.3 43.2 Any chemotherapy 24.2 Wright et al14 1146 All Any 6.10 Wang et al⁵² 82 242 2005 Αll 14.00 2009 14.80 Cheung et al³¹ 107 253 All Any 6.30 Amano et al⁵⁸ 266 All Early palliative care 3.70 54 9 SOC 8 Gonsalves et al44 200 All 2002 6 2008 16 38 Ho et al³⁷ 227 161 All 1993 8.50 2003 7.50 Tang et al²⁸ Trend 2000-2006 44.4 242 530 All 14.2 Temel et al³⁶ **NSCLC** Integrated palliative care 13 46 Earle et al¹⁶ 28 777 All 1993 12.50 1996 11.60

NSCLC, non-small cell lung cancer; SOC, standard of care.

palliative care and found a significantly lower number of the latter had been admitted to the ICU during the last 30 DOL.^{65 73} Also, in another study comparing management by palliative care teams for >3 months with a shorter time, ICU admissions were 10%–5.6% lower.⁵⁸

Late admissions to EOL palliative care

The absence of palliative care management or its too late initiation, ie <3 days before death, has been poorly investigated. Although this indicator, which concerns 6%–54% of the patients, is more difficult to establish and interpret (table 5), its impact on the degree of entourage satisfaction is high.

Factors associated with EOL palliative care

According to a study using Canadian databases, ¹⁴ the families of patients enrolled in hospice >3 days before death considered EOL-care quality to be excellent more often than those receiving none or <3 days (58.8% vs 43.1%, respectively). Families of patients receiving <3 days of palliative care were less likely to report patients died in their preferred place (40.0% vs 72.8%). Male sex, high Charlson Comorbidity Index, management by oncologists as opposed to 'organ' specialists and management in a teaching hospital were significantly associated with a higher risk of late admission to palliative care. In contrast, the risk of late access to palliative care was lower for African-Americans, single or divorced individuals and when access to it was available. ¹¹

Economic impact of EOL-care aggressiveness

From a societal point of view, patients receiving EOL care considered aggressive have higher healthcare costs than the others (online supplementary appendix table S2).

This overexpenditure was about \$C6000 (\$C18 131 vs \$C12 678) based on Canadian administrative data.³¹ An analysis of EOL costs in seven Western countries found means ranging from US\$3160 in the UK to US\$10 273 in Canada, without EOL lung cancer costs being significantly different from those of other cancers.⁴⁹ EOL care was mainly in hospitals in Belgium, Canada, the UK, Germany and Norway, compared with the Netherlands and the USA. EOL hospital expenditures were higher in the USA, Norway and Canada, intermediate in Germany and Belgium, and lower in the Netherlands and the UK.

One study used the American SEER-Medicare database to examine the extent to which intensive EOL-care patterns could explain EOL-care costs for dying patients with cancer. To wit, 49.2% received at least one intensive EOL-care intervention, which incurred substantially higher expenditures during the last 30 DOL, compared with not receiving such intensive care (US\$18 700 vs US\$4200, respectively; p<0.001). Among those benefiting from at least one intensive EOL-care intervention, 71.3% of the EOL expenditures were for inpatient services. In contrast, palliative care was the primary expenditure for decedents not receiving any intensive EOL care (66.6%). ICU admission appeared to be the major driver of EOL-spending differences.

DISCUSSION

The results of this systematic review synthesis showed that management of the EOL period differs among countries for patients with lung cancer. EOL-care aggressiveness is less intensive in Canada than the USA. This difference might reflect the differences in the healthcare systems of those two countries, cultural specificities,

Table 4 Studies reports on end-of-life ICU admissions

| Study | Patients, N | Type of cancer | Inclusion criteria | % admitted to ICU |
|--------------------------------------|----------------|----------------------|-----------------------------------|-------------------|
| Mrad <i>et al</i> ⁶¹ | 412 496 | NSCLC | 1998 | 13.3 |
| | | | 2014 | 27.9 |
| Presley <i>et al</i> ²⁶ | 43 654 | NSCLC | Veterans | 0.90 |
| | | | Medicare beneficiaries | NR |
| Ersek <i>et al</i> ⁴² | 847 | NSCLC | Any | 25.40 |
| Falchook <i>et al</i> ³² | 12 765 | Lung | Patient <65 years | 20.60 |
| Wu et al ²³ | 49 920 | All | Palliative chemotherapy <6 months | 31.5 |
| | | | Any chemotherapy | 6.5 |
| Nright <i>et al</i> ¹⁴ | 1146 | All | Any | 13.20 |
| Nang <i>et al⁵²</i> | 82 242 | All | 2005 | 15.30 |
| | | | 2009 | 19.50 |
| Bekelman <i>et al⁴⁹</i> | 389 073 | All | USA | 5.50 |
| | | | The Netherlands | 7 |
| | | | Belgium | 11 |
| | | | Canada | 9.80 |
| | | | UK | NR |
| | | | Germany | 3.50 |
| | | | Norway | NR |
| Choi <i>et al³⁰</i> | 263 | All | 2002 | 1.8 |
| | | | 2012 | 19.9 |
| Cheung <i>et al</i> ³¹ | 107 253 | All | Any | 5.5 |
| Amano <i>et al</i> ⁵⁸ | 266 | All | Early palliative care | 5.60 |
| | | | SOC | 10 |
| Zhang <i>et al</i> ⁶³ | 118 | NSCLC | Any | 25 |
| Mack <i>et al</i> ⁴⁶ | 1231 | NSCLC and colorectal | Any | 9 |
| Ho <i>et al³⁷</i> | 227 161 | All | 1993 | 3.06 |
| | | | 2003 | 5.39 |
| Gonsalves <i>et al</i> ⁴⁴ | 200 | All | 2002 | 6 |
| | | | 2008 | 33 |
| Keating <i>et al</i> ⁴⁷ | 5826 | NSCLC and colorectal | VHA cohort | 12.50 |
| | | | SEER cohort | 19.7 |
| Tang <i>et al</i> ²⁸ | 242 530 | All | Trend 2000–2006 | 11.4 |
| Temel <i>et al</i> ³⁶ | 46 | NSCLC | Integrated palliative care | 2 |
| Barbera <i>et al</i> ³⁴ | 5855 | All | Any | 5.50 |
| Earle <i>et al</i> ¹⁶ | 28 777 | All | 1993 | 7.10 |
| | | | 1996 | 9.40 |

NR, not reported; NSCLC, non-small cell lung cancer; PBS, population-based study; SOC, standard of care; VHA, Veterans Health Administration.

doctors' usual practices. In the USA and Australia, doctors are encouraged to prescribe anticancer therapies, whereas in Canada, the national healthcare system no longer reimburses the cost beyond a certain number of chemotherapy lines.

This synthesis shows how difficult it is to compare the quality of EOL-care management because no global quality indicator exists. The most used criteria were developed by Earle *et al.*¹⁵ ¹⁶ Among them, the most frequently analysed were recourse to chemotherapy during the last 14 or 30 DOL and ED visits during the last 30 DOL. A growing body of data suggests that less use of palliative versus active chemotherapy for patients recognised as having short life expectancy or more frequent EOL discussions with them might reduce

intensive EOL care and promote earlier access to palliative care, thereby improving the quality of EOL care of patients with advanced cancer. 14-16 23 29 31 49 61 64 Moreover, it was shown that EOL discussions might be particularly important for patients receiving palliative chemotherapy, who should be informed by substantiated scientific data of the likely outcomes associated with its use. 64 88 The EOL period should be prepared beforehand with the patient and his/her entourage. It was shown that family members' perceptions of EOL care of elderly patients who died of lung or colorectal cancer were associated with earlier palliative care enrolment, avoidance of ICU admissions during the last 30 DOL and death outside the hospital. 88

Table 5 Studies on the transition to palliative care during the last 3 days of life (DOL)

| Study | Patients, | Type of cancer | Inclusion criteria | % Without palliative care during the last 3 DOL |
|----------------------------------|-----------|----------------------------|----------------------------|---|
| Mrad et al ⁶¹ | 412 496 | NSCLC | 1998 | 8.7 |
| | | | 2014 | 53.0 |
| Falchook et al ³² | 12 765 | Lung | Patients<65 years | 54 |
| Wright et al ¹⁴ | 1146 | All | Any | 47.7 |
| Wang <i>et al</i> ⁵² | 82 242 | All | 2005 | 7.2 |
| | | | 2009 | 8.3 |
| Mack <i>et al</i> ⁴⁶ | 1231 | NSCLC and colorectal | Any | 42 |
| Tang <i>et al</i> ²⁸ | 242 530 | All | Trend 2000– 2006 | 17.9 |
| Temel <i>et al</i> ³⁶ | 46 | NSCLC | Integrated palliative care | 35 |
| Earle <i>et al</i> ¹⁶ | 28 777 | All | 1993 | 14.3 |
| | | | 1996 | 17 |

NSCLC, non-small cell lung cancer.

Limitations of this study

Limitations of this systematic review are those inherent with the methods used and the lack of data on this relatively new field of interest. In spite of the search strategy used, key papers may have been omitted. The two researchers extracted data only from the few available studies. A systematic review depends on existing and accessible evidence. When these are unavailable, insufficiently reported, biassed or of very poor quality, it may not be possible to draw definitive conclusions. Similarly, certain stages of systematic reviews imply a degree of subjectivity (selection of studies, evaluation of quality, narrative synthesis of results). In our case, very few prospective or interventional studies have been conducted. As of 29 May 2019, no study has been registered in ClinicalTrials.gov on EOL-care aggressiveness for patients with lung cancer. Because lung cancer management has changed dramatically over the last 15–20 years, we limited ourselves to studies published since 2000, which concerned patients treated during 1990-1995. Data from years prior to 1995 no longer correspond to current reality and we wanted to avoid biassing our analysis with no longer relevant findings. Similarly, grey literature was not included in our analysis because there are so few prospective studies or cohorts that the grey reports could be misleading with data not published in peer-reviewed journals.

Because the included studies were from resource-rich countries, with well-developed palliative care systems, generalisability of our findings to services in resource-poor countries is limited. The types of studies retained for this analysis represent another limitation. Most of the data on the quality of EOL-care management come

from medical administrative databases or retrospective studies. Few data are available from prospective studies or cohorts, which can lead to biassed analyses. In addition, only Earle *et al*¹⁶ considered all six validated criteria of EOL-care quality, while the other studies focused on only one to four of those indicators.

Among the other limitations of our review, we did not evaluate the studies for the risk of bias. Unfortunately, it is difficult to analyse the biases of each study because of the variety of types of studies retained. We also did not contact the authors to obtain individual lung cancer data for several reasons: first, we are not sure that they had analysed their data by type of cancer, and difficulty of obtaining individualised results could alter our results but it is known that lung cancer was included in most studies, and breast cancer is the most aggressively treated. The mix with other cancers probably weakens our results rather than strengthening them.

Implications for clinical practice and future research

Over time, a progressive trend towards higher percentages of patients receiving EOL care deemed aggressive was seen, probably because of the availability of enhanced numbers of therapeutic options and better tolerance of them. Those aggressive EOL treatments raised EOL-care costs, regardless of the healthcare system examined, without any significant benefit for the patient or his/ her family in terms of improved quality of life.⁸⁹ Early access to palliative care lowered the number of patients given aggressive treatments.90 Unfortunately, in most healthcare systems, early access to palliative care during the course of these cancers remains limited. National or international research programmes should contribute to earlier and better identification of the factors supporting aggressive EOL-treatment interventions, and those favouring better palliative management of these patients.

CONCLUSION

Management of patients with lung cancer is complex, notably during the EOL period. The percentage of patients receiving aggressive EOL care continues to rise and it carries a non-negligible impact on healthcare costs. Prospective complementary studies are warranted to better discern the factors supporting aggressive EOL care and to evaluate early palliative care programmes.

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