Measuring the Quality of Palliative Care for Patients with End-Stage Liver Disease

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Abstract
Background/Aims We examined the quality of palliative care received by patients with decompensated cirrhosis using an explicit set of palliative care quality indicators (QIs) for patients with end-stage liver disease (PC-ESLD).

Methods We identified patients newly diagnosed with decompensated cirrhosis at a single veterans health center and followed up them for 2 years or until death. We piloted measurement of PC-ESLD QIs in all patients confirmed to have ESLD using a chart abstraction tool.

Results Out of 167 patients identified using at least one sampling strategy, 62 were confirmed to meet ESLD criteria with chart abstraction. Ninety-eight percent of veterans in the cohort were male, mean age at diagnosis was 61 years, and 74% were White. The overall QI pass rate was 68% (64% for information care planning QIs and 76% for supportive care QIs). Patients receiving specialty palliative care consultation were more likely to receive information care planning QIs (67% vs. 37%, \( p = 0.02 \)). The best performing sampling strategy had a sensitivity of 62% and specificity of 60%.

Conclusion Measuring the quality of palliative care for patients with ESLD is feasible in the veteran population. Our single-center data suggest that the quality of palliative care is inadequate in the veteran population with ESLD, though patients offered specialty palliative care consultation and those affected by homelessness, drug, and alcohol abuse may receive better care. Our combination of ICD-9 codes can be used to identify a cohort of patients with ESLD, though better sensitivity and specificity may be needed.

Keywords Palliative care · Cirrhosis · Healthcare quality · Veterans · End of life

Abbreviations
ICD-9 CM International Classification of Diseases, 9th Revision, Clinical Management
CDW Corporate data warehouse

Introduction
End-stage liver disease (ESLD) is a morbid, costly condition characterized by advanced fibrosis of the liver and limited residual hepatocyte function [1]. Fifty percent of patients with ESLD will die within 2 years of their index decompensation, and while liver transplantation is a definitive cure, it is available to a limited number of patients [2]. For example, there were 38,170 reported deaths from ESLD in the USA in 2014, while only 7237 liver transplants were performed for all forms of liver disease [3, 4]. The supportive needs of patients with ESLD are also high. Their symptom burden is comparable to patients with advanced lung and colon cancer in the last 6 months prior to death, and Medicare beneficiaries with cirrhosis also report greater functional disability and use of healthcare utilization compared to age-matched controls and patients with heart failure [5, 6].

Despite the immense morbidity and high burden of palliative issues in patients with ESLD, a growing body of literature suggests that referral to services that can provide adequate symptom control and coordination of end-of-life care for this population is generally poor. Only 29% of Medicare beneficiaries from 2007 to 2011 with ESLD were referred to hospice within 3 years of an index hospitalization despite the fact that 77% of the same cohort were re-hospitalized and 56% died in the same follow-up period [7]. In a national study examining in-hospital deaths of patients with ESLD...
from 2009 to 2013, use of invasive life-sustaining treatments was high with 56% percent of patients receiving mechanical ventilation, 16% receiving hemodialysis, and 6% receiving cardiopulmonary resuscitation. However, only 30% of patients received palliative care, which was associated with reduced procedure burden, lower costs, and lower lengths of stay [8].

A lack of clear guidelines to help healthcare teams deliver high-quality palliative care and coordinate end-of-life care for patients with ESLD is evident, though these have long been a priority for subspecialty organizations treating cancer and other non-chronic conditions such as heart failure, chronic obstructive pulmonary disease, and end-stage renal disease [9–12]. Nineteen quality indicators (QIs) for palliative care in ESLD were recently selected using the RAND/UCLA appropriateness method (RAM) [13]. Our study’s primary aim was to pilot the use of these QIs to measure the quality of palliative care delivered to veterans with ESLD. The Veterans Affairs (VA) Health System is the largest provider of care for patients with hepatitis C, and given the rising incidence of ESLD in this population, optimizing palliative care in this population is particularly crucial [14–16]. Our secondary aim is to compare the sensitivity and specificity of three different sampling strategies used in previous research to identify patients with ESLD within the veteran population.

Methods

Cohort Identification

We identified a cohort of veterans from a single VA health system, the VA Greater Los Angeles (GLA) Healthcare System, that had been newly diagnosed with ESLD in the year 2012 using VA Corporate Data Warehouse (CDW). We first included patients using any of the following three different sampling strategies reported in the literature: (1) at least one International Classification of Diseases, 9th edition (ICD-9) code for cirrhosis plus at least one ICD-9 code for liver decompensation [17], (2) at least one ICD-9 code for liver decompensation during an inpatient hospitalization (limited to ascites, spontaneous bacterial peritonitis (SBP), or variceal hemorrhage), and (3) at least two ICD-9 codes for liver decompensation (limited to ascites, SBP, or variceal hemorrhage) [18]. We only included patients with at least one outpatient visit or one hospitalization at the GLA VA. In order to capture a population of patients that were newly decompensated, patients with any ICD-9 code for liver decompensation in 2011 were excluded (Table 1).

Next, we used a chart abstraction tool to confirm clinical documentation of a diagnosis of cirrhosis in the medical record as well as criteria for ESLD (MELD score greater than or equal to 20 or a Child–Turcotte–Pugh (CTP) score of 12 or greater from 2012 to 2013). To ensure a minimum amount of time to measure the quality of palliative care, only patients alive for at least 30 days following initial diagnosis of ESLD were included in the final sample. Interrater agreement between 2 chart abstractors used for this study was determined by randomly selecting 10% of patient charts to determine how frequently both abstractors agreed on the criteria for screening in or screening out patients for inclusion in our final cohort.

Development of Quality Indicators for Palliative Care in ESLD

A previous publication highlighted the procedure involved in developing and validating these QIs developed to be used with medical record abstraction using the RAM [13]. This process involved multiple steps, including review of guidelines, development of candidate quality indicators, and convening of an expert multidisciplinary panel for review of these indicators to determine appropriateness using a modified Delphi approach. From this analysis, 13 information and care planning and 6 supportive care QIs were chosen.

Full Medical Record Abstraction of Quality Indicators

Medical records were abstracted from time of diagnosis for 2 years or until death. A full chart abstraction tool

| Table 1 Sampling strategies used for identification of ESLD |
|-----------------------------------------------------------|
| 1  | At least 1 ICD-9 code for cirrhosis\(^a\) and at least 1 ICD-9 code for liver decompensation\(^b\) |
| 2  | At least 1 ICD-9 code for liver decompensation during an inpatient admission (limited to only ascites, spontaneous bacterial peritonitis, and variceal hemorrhage)\(^c\) |
| 3  | At least 2 ICD-9 codes for liver decompensation (limited to ascites, spontaneous bacterial peritonitis, or variceal hemorrhage)\(^c\) |

\(^a\)ICD-9 codes for cirrhosis include 571.2, 571.5, 571.6

\(^b\)ICD-9 codes for liver decompensation include 789.5, 789.59, 572.3, 456.0, 456.20, 456.2, 456.1, 456.21, 567.0, 567.2, 567.21, 567.23, 567.29, 567.8, 567.89, 567.9, 572.2, 572.4

\(^c\)ICD-9 codes for liver decompensation (limited to ascites, spontaneous bacterial peritonitis, or variceal hemorrhage) include 789.5, 789.59, 572.3, 456.0, 456.20, 456.2, 456.1, 456.21, 567.0, 567.2, 567.21, 567.23, 567.29, 567.8, 567.89, 567.9
with guidelines and training materials was developed to measure all 19 palliative care quality indicators for ESLD, as well as criteria used to determine numerators and denominators for each QI. A patient who was eligible for a QI received a score of 1 if the recommended process measure was received; otherwise, they received a score of 0. Pass rates were calculated by dividing the number of times the QIs were passed over the number of eligible patients for that QI. We abstracted information on patient demographics, homelessness, history of substance abuse, history of viral hepatitis C, history of hepatocellular cancer, documentation pertaining to liver transplant evaluation, and receipt of specialty palliative care services. We also calculated overall quality scores by domain as an average across the domain measures and stratified scores based on variables of interest to explore how different patient characteristics are associated with the quality of palliative care provided. Two-sided Fisher’s exact tests were used to compare the categorization of domain-level quality scores by patient characteristics. Interrater reliability on receipt of quality indicators was determined by calculating the kappa coefficient for each numerator and denominator for a 20% subset of our final sample that were abstracted separately by two abstractors. The confidence intervals for sensitivity and specificity are exact binomial confidence intervals constructed using the method of Clopper and Pearson. [19].

Comparison of Sampling Strategies

In order to compare the various sampling strategies used to identify patients with ESLD, we calculated the sensitivity and specificity of each of the strategies separately against a gold standard of the screening abstraction tool.

Results

Descriptive Statistics of Patient Cohort

We identified 167 patients from the VA CDW that met at least one of our three sampling strategies in 2012. Out of this sample, 71 patients were identified to be true positives using the screening chart abstraction tool, while 96 were screened out because either MELD and Child–Pugh cutoffs were too low or laboratory data were missing. Since 9 patients of the included cohort died within 30 days of diagnosis, our final sample contained 62 patients. Our interrater agreement for screening for a 10% sample (N = 17) was 100% (Fig. 1). Our final sample had a mean age at diagnosis of 61 years; all but one patient were male, 74% were White, and 53% were either single, separated, divorced, or widowed. Sixty-six percent of patients had died by the end of the study period. Seventy-six percent of patients were exempt from co-pay, 36% were homeless, and 10% lived in supportive housing. Seventy-nine percent of patients had a history of tobacco use, and 66% reported a history of drug or alcohol abuse. Sixteen percent of patients had a diagnosis of liver cancer.

![Fig. 1 Flow diagram for cohort identification process](image-url)
and 60% of patients had a diagnosis of hepatitis C. Forty-two percent of patients had no documentation of whether they were being considered for liver transplant, 48% were not being considered for transplant, and 10% were being considered for transplant at the end of the study period. Forty-five percent received specialty palliative care consultation (Table 2).

**Receipt of Quality Indicators**

Our cohort of 62 patients triggered 245 information care planning QIs, of which 157 (64%) were passed, and 118 supportive care QIs, of which 90 (76%) were passed. The overall pass rate was 68%. Seventeen out of 19 QIs were triggered at least twice with a denominator range of 2–62. Interrater agreement was good to excellent with denominator simple kappa coefficients ranging from 0.75 to 1 and numerator kappa scores ranging from 0.56 to 1.

Of all patients with ESLD who died during follow-up, 63% of patients were offered specialty palliative care or hospice within 6 months of death. Out of patients being considered for transplantation, only 55% were offered advance care planning (ACP) within 3 months of consideration, while only 50% of patients not considered for transplant were offered ACP. Of the patients not considered for transplant who were initiated on hemodialysis or received an intracardiac device, only 14% had timely preferences or goals of care documented. Fifty-eight percent of patients newly diagnosed with hepatic encephalopathy had ACP documented in their chart within a month of that diagnosis, while 91% percent of patients who died an expected death had evidence of ACP performed within 6 months of death. All patients with ESLD admitted to a hospital or nursing home had the name of their surrogate decision maker identified (or documentation to identify a surrogate decision maker) in the electronic health record.

When receiving treatment in the intensive care unit, only 32% of patients who survived 48 h after admission had their care preferences documented and none of the 5 patients receiving mechanical ventilation for 48 h had goals of care or preferences for mechanical ventilation documented. Only 19% of patients with advance directive or DNR orders prior to being admitted to a second venue had the DNR present in their medical record at the second venue, but all patients with orders to withhold life-sustaining treatment had their preferences followed.

With regard to treatment of pain, 39% of patients in the full sample were inappropriately prescribed NSAIDs, while only 7% had total dosages of Tylenol exceeding 3 grams. Ninety-one percent of patients with symptomatic ascites had paracentesis offered within 48 h of admission, and none of the 5 patients experiencing dyspnea in the last 3 days had their care of dyspnea documented (Table 3).

**Stratifying Quality Indicator Receipt by Variables of Interest**

Patients with the highest overall QI pass rates included those who were homeless (75%), with a history of HCC (71%), and patients receiving specialty palliative care services (66%). Those with the lowest rates included patients who did not receive palliative care (44%) and without history of HCC (57%). Differences in overall pass proportions by clinical characteristics were not statistically significant.

The highest pass proportions for information planning QIs were observed in patients who were not candidates for liver transplant (62%), had a history of HCC (74%), and who were homeless (72%). The lowest pass proportions for information planning were observed in patients who did not receive palliative care services (37%) and those that did not have HCC (55%). The difference in information planning QI

### Table 2

| Characteristic                                | Mean (SD) or N (%) | Total = 62 |
|-----------------------------------------------|--------------------|------------|
| Age at diagnosis                              | 61 (7)             |            |
| Sex                                           |                    |            |
| Male                                          | 61 (98%)           |            |
| Female                                        | 1 (2%)             |            |
| Race                                          |                    |            |
| White                                         | 46 (74%)           |            |
| African-American                              | 9 (15%)            |            |
| Asian                                         | 1 (2%)             |            |
| Other                                         | 4 (7%)             |            |
| No data                                       | 2 (3%)             |            |
| Marital status                                |                    |            |
| Married                                       | 21 (34%)           |            |
| Lives with significant other                  | 2 (3%)             |            |
| Single, separated, divorced, or widowed       | 32 (53%)           |            |
| No data                                       | 6 (10%)            |            |
| Co-pay exempt                                 | 47 (76%)           |            |
| Homeless                                      | 22 (36%)           |            |
| Lives in supportive living                    | 6 (10%)            |            |
| Smoker                                        | 49 (79%)           |            |
| History of drug or alcohol use                | 41 (66%)           |            |
| Hepatitis C                                   | 37 (60%)           |            |
| Hepatocellular carcinoma (HCC)                | 10 (16%)           |            |
| Transplantation status                        |                    |            |
| No documentation                              | 26 (42%)           |            |
| Not being considered for transplant           | 30 (48%)           |            |
| Still being considered                        | 6 (10%)            |            |
| Died                                          | 41 (66%)           |            |
| Received specialty palliative care consultation| 28 (45%)           |            |
| Domain                          | Quality indicator (QI)                                                                 | Eligible events (No.) | QIs passed (%) |
|--------------------------------|----------------------------------------------------------------------------------------|-----------------------|----------------|
| Information care planning      | IF a patient with advanced ESLD and HCC with BCLC Stage D is not a candidate for transplant, THEN the patient should be offered palliative care and/or hospice services or a reason why should be documented BECAUSE survival is less likely than 6 months and hospice provides additional support for patients and caregiver/family. | 245                  | 64             |
|                                | IF a patient has advanced end-stage liver disease and is diagnosed with hepatorenal syndrome and is not a liver transplant candidate, THEN the patient should be offered palliative and/or hospice services BECAUSE these patients have a poor prognosis and would benefit from the additional supportive services | 5                    | 80             |
|                                | IF a patient with advanced end-stage liver disease dies, THEN there should be evidence that he or she has been offered palliative care or hospice within 6 months before death BECAUSE palliative care teams can improve a range of patient and caregiver outcomes | 41                   | 63             |
|                                | IF a patient has advanced end-stage liver disease and is being considered for transplant, THEN the patient should be offered advance care planning within 3 months of consideration of transplant or a reason why such a discussion did not occur BECAUSE effective communication about goals of care is important to ensure that patient preferences are followed during the course of their illness | 20                   | 55             |
|                                | IF a patient is newly diagnosed with advanced end-stage liver disease and is not considered for transplant, THEN the patient should be offered advance care planning within 3 months or a reason why such a discussion did not occur BECAUSE effective communication about goals of care is important to ensure that patient preferences are followed during the course of their illness | 16                   | 50             |
|                                | IF a patient with advanced end-stage liver disease dies an expected death, THEN there should be documentation of advance care planning in the medical record within 6 months before death BECAUSE a patient’s goals and values should guide treatment and life-sustaining care decisions | 22                   | 91             |
|                                | IF a patient with advanced end-stage liver disease is admitted to a hospital or nursing home, THEN within 48 h of admission the medical record should contain the name of the patient’s surrogate decision maker, or documentation of a discussion to identify/search for a surrogate decision maker BECAUSE patient’s values and preferences should guide life-sustaining care and patients with end-stage liver disease often cannot make decisions for themselves | 54                   | 100            |
|                                | IF a patient with advanced end-stage liver disease is newly diagnosed with hepatic encephalopathy and does not have advance care planning previously documented, THEN the medical record should document advance care planning or lack of ability to do so within 1 month BECAUSE patients with a history of hepatic encephalopathy are at higher risk of lacking capacity to make decisions for themselves and care should be guided by their goals and preferences | 12                   | 58             |
|                                | IF a patient with advanced end-stage liver disease is admitted to the ICU and survives 48 h, THEN within 48 h, the medical record should document that the patient’s preferences for care have been considered or an attempt was made to identify them BECAUSE patients with end-stage liver disease are at a high risk of requiring life-sustaining treatment decisions | 31                   | 32             |
|                                | IF a patient with advanced end-stage liver disease requires mechanical ventilation for greater than 48 h, THEN within 48 h of the initiation of mechanical ventilation, the medical record should document the goals of care and the patient’s preference for mechanical ventilation or why this information is not available BECAUSE patients’ values and preferences should guide life-sustaining care | 5                    | 0              |
|                                | IF a patient with advanced end-stage liver disease has orders in the hospital or nursing home to withhold or withdraw a life-sustaining treatment (e.g., DNR, no tube feeding, no hospital transfer), THEN these treatment preferences should be followed BECAUSE patients’ values and preferences should guide end-of-life care | 9                    | 100            |
pass rate by palliative care receipt was statistically significant (67% vs. 37%, \(p=0.02\)). The highest pass proportions for supportive care QIs were observed in patients with a history of homelessness (87%) and in patients with history of drug or alcohol abuse (83%). The difference in supportive care QI pass proportions by homelessness (87% vs. 58%, \(p=0.02\)) and history of drug or alcohol abuse (83% vs. 58%, \(p=0.05\)) were significantly different (Table 4).

**Sensitivity and Specificity of Sampling Strategies**

Sensitivities and specificities for the three sampling strategies were calculated and compared. Sampling strategy 1 had the highest sensitivity of 87% and the lowest specificity at 22%. Sampling strategy 2 had the highest specificity at 60% and a sensitivity of 62% (Table 5).

**Discussion**

This pilot study successfully appraised the quality of palliative care delivered to a cohort of decedent veterans who were newly diagnosed with ESLD. The overall receipt of quality palliative care was 68%, which is a low statistic considering the fact that only 10% had documentation that they were being actively considered for liver transplant at the end of the study period and the population had MELD and Child–Pugh score cutoffs of 20 and 12, which correspond to...
20% 3-month and 55% 1-year mortality rates, respectively [20].

Our study found that the receipt of supportive care QIs (76%) was higher than information care planning QIs (64%). A similar discrepancy in receipt of QIs by domain was observed in a study investigating quality of end-of-life care in patients with various serious chronic illnesses hospitalized at a major quaternary academic medical center [21]. In our cohort, only about half of patients were offered ACP, whether they were being considered or not considered for liver transplant. Furthermore, ACP was only offered to 58% of patients with newly diagnosed hepatic encephalopathy, which increases the risk of a patient lacking capacity to make their own decisions. Information care planning practices were even less frequently adopted once patients become hospitalized. Goals of care discussions were documented less than one-third of time and not at all for patients surviving 48 h after ICU admission and after mechanical ventilation, respectively.

There has been increasing interest in improving ACP in patients with ESLD [22], and our findings suggest that greater attention should be paid to understanding these barriers and describing new clinical models of care. In patients with advanced malignancies, for example, shared mental models have been developed to better provide patients with the support needed to guide and document their future choices [23]. Our findings also suggest a statistically higher receipt of ACP in patients who received specialty palliative care consultation. Recently, VA has implemented a Life-Sustaining Treatment Decisions Initiative (LSTDI) nationally aimed at improving goals of care conversations among veterans with serious illness and future work should evaluate whether this has a positive impact on these quality indicators for veterans with ESLD.

The frequency of palliative care consultation for patients with ESLD who are hospitalized at the end of life has been increasing, though these studies do not incorporate a direct measure of specialty services [8]. In patients who are being evaluated for liver transplant, concurrent specialty palliative care consultation was associated with improvement in symptom burden and depression symptoms after 6 months of follow-up [24]. This study further adds to these findings that specialty palliative care services may provide benefits for patients with ESLD. In addition, our study suggests that referral rates to specialty palliative care in ESLD may be lower than those for veterans with

### Table 4 Comparison of quality indicator pass proportions by clinical and demographic characteristics

| Characteristic                              | N     | Overall QI pass proportions | P value | Information care planning QI pass proportion | P value | Supportive care QI pass proportion | P value |
|---------------------------------------------|-------|----------------------------|---------|---------------------------------------------|---------|-----------------------------------|---------|
| Transplant status                           |       |                            |         |                                             |         |                                   |         |
| No documentation or still being considered  | 32    | 64%                        | 0.79    | 58%                                        | 0.80    | 82%                               | 0.09    |
| Not considered or not a candidate           | 30    | 62%                        |         | 62%                                        |         | 59%                               |         |
| Receipt of specialty palliative care services |       |                            |         |                                             |         |                                   |         |
| Yes                                         | 32    | 66%                        | 0.13    | 67%                                        | 0.02    | 60%                               | 0.10    |
| No                                          | 30    | 44%                        |         | 37%                                        |         | 79%                               |         |
| Presence of hepatocellular cancer           |       |                            |         |                                             |         |                                   |         |
| Yes                                         | 10    | 71%                        | 0.50    | 74%                                        | 0.50    | 61%                               | 1.00    |
| No                                          | 52    | 57%                        |         | 55%                                        |         | 60%                               |         |
| History of Active Drug or Alcohol Use       |       |                            |         |                                             |         |                                   |         |
| Yes                                         | 26    | 67%                        | 0.79    | 63%                                        | 1.00    | 83%                               | 0.05    |
| No                                          | 36    | 61%                        |         | 61%                                        |         | 58%                               |         |
| Homelessness                                |       |                            |         |                                             |         |                                   |         |
| Yes                                         | 22    | 75%                        | 0.26    | 72%                                        | 0.41    | 87%                               | 0.02    |
| No                                          | 40    | 59%                        |         | 59%                                        |         | 58%                               |         |

*P value for two-sided Fisher’s exact test to compare proportions

### Table 5 Sensitivity and specificity of various sampling strategies

|                          | 1                          | 2                          | 3                          |
|--------------------------|-----------------------------|-----------------------------|-----------------------------|
| Sensitivity (95% CI)     | 87% (77.3–94.0%)            | 62% (49.7–73.2%)            | 49% (37.2–61.4%)            |
| Specificity (95% CI)     | 22% (14.1–31.5%)            | 60% (48.9–69.3%)            | 57% (46.8–67.3%)            |
advanced cancer, highlighting potential room for improvement [25].

Supportive care QIs were more frequently met, likely owing to the fact that avoidance of high doses of acetaminophen and provision of paracetamol for ascites are well-established practices in the management of all patients with ESLD [26]. The avoidance of NSAIDs, however, was not followed 39% of the time, which may reflect the tension that providers face when only limited options are present to alleviate pain in patients with ESLD, who are at higher risk of adverse events from available pain relief strategies, such as narcotics [27]. Patients with a history of homelessness and history of drug and alcohol abuse were more likely to receive supportive QIs. This runs counter to previous studies that have shown more challenges in access and provision of palliative care in these patient populations but may reflect the high commitment to providing these services in the veteran Health Administration [28–30].

Lastly, a combination of one ICD-9 code for cirrhosis and one ICD-9 code for liver decompensation (limited to ascites, SBP, and variceal hemorrhage) had a sensitivity of 62% and specificity of 60% for identifying patients from our total cohort. These moderate values likely reflect the fact that our cohort was focused on a particularly sick population (identified by a MELD score ≥ 20 and Child–Pugh score ≥ 12).

There are several strengths to our study. Our hierarchical method of identifying patients with ESLD using three different sampling strategies and a chart abstraction tool to confirm diagnosis was highly reliable given the 100% interrater agreement between abstractors. Our chart abstraction tool to identify QIs for palliative care in patients with ESLD was also both feasible and reliable, given that interrater agreements were good to excellent. Our findings also highlight significant gaps in the provision of palliative care to patients with ESLD, which are consistent with the currently available literature, suggesting validity of our results.

A few limitations should be highlighted. The findings of this study were performed at a single VA medical center and as such, reported results from a small sample size that cannot be generalized. Certain stratified trends in QI receipt that were not significant using our cohort may be more apparent using a larger cohort of patients. Future work should explore natural language processing and machine learning as potential methods to improve identification of a denominator with higher sensitivity and specificity. The perspectives of veterans with ESLD and caregivers on receipt of palliative care and hospice services should also be collected to help guide future quality improvement work in this field. Our findings demonstrate that it is feasible to measure the quality of palliative care for a veteran cohort of patients with ESLD with good reliability. Our identification of a seriously ill group of patients with ESLD using a combination of ICD-9 codes can be used for future cohort studies in the VA system that are looking to investigate their clinical care and outcomes. Our preliminary findings from this study highlight the need for better information care planning in patients with ESLD, particularly as patients’ care becomes more acute, and suggest some benefit in involving specialty palliative care services in improving this goal. We believe our methods for measuring quality of palliative care can be applied to the national veteran population, which can further add to our knowledge of ways to better improve the care of this vulnerable group.

Author’s contribution FK, SA, KL, and AW were involved in study concept and design; all authors were involved in data acquisition, analysis, and interpretation; AP, AA, and AW were involved in drafting of manuscript; all authors were involved in critical revision of manuscript; FK, SA, KL, and AW were involved in study supervision; AW was involved in administrative, technical, or material support; and AA, ML, and AW were involved in statistical analysis. All authors approved the final submitted manuscript.

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Compliance with Ethical Standards

Conflict of interest The authors declare that they have no conflict of interests.

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