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Measuring Disease-Free Survival and Cancer Relapse Using Medicare Claims From CALGB Breast Cancer Trial Participants (Companion to 9344)

Elizabeth B. Lamont, James E. Herndon II, Jane C. Weeks, I. Craig Henderson, Craig C. Earle, Richard L. Schilsky, and Nicholas A. Christakis For the Cancer and Leukemia Group B

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Abstract

To determine the accuracy with which Medicare claims data measure disease-free survival in elderly Medicare beneficiaries with cancer, we performed a criterion validation study. We merged gold-standard clinical trial data of 45 elderly patients with node-positive breast cancer who were treated on the Cancer and Leukemia Group B (CAL-GB) adjuvant breast trial 9344 with Centers for Medicare and Medicaid Services (CMS) data files and compared the results of a CMS-based algorithm with the CALGB disease-free survival information to determine sensitivity and specificity. For 5-year disease-free survival, the sensitivity of the CMS-based algorithm was 100% (95% confidence interval [CI] = 81% to 100%), the specificity was 97% (95% CI = 83% to 100%), and the area under the receiver operator curve was 97% (95% CI = 90% to 100%). For 2-year disease-free survival, the test characteristics were less favorable: sensitivity was 83% (95% CI = 36% to 100%), specificity was 95% (95% CI = 83% to 100%), and area under the receiver operator curve was 84% (95% CI = 66% to 100%).

The elderly are numerically under-represented (1–3) and possibly physiologically misrepresented (4) in clinical trials of cancer chemotherapy such that trial participants are potentially younger and healthier than the average elderly cancer patient. The general population of elderly Americans therefore may not experience the same benefits and toxicities of chemotherapy as trial participants. Nevertheless, clinicians need information about the expected benefits and toxicities of chemotherapy in the elderly. A solution to this problem is to supplement clinical trial results with observational results. Prior research has shown that Centers for Medicare and Medicaid Services (CMS) data can be used to accurately measure chemotherapy use (5,6), but the extent to which it can be used to measure outcomes that are traditionally reported in clinical trials, including conventional survival and toxicity endpoints, is unknown.

Disease-free survival is one of the most common survival metrics in adjuvant chemotherapy trials and is defined as the time from enrollment to the first of two events: cancer relapse or death from any cause. Disease-free survival incorporates the morbidity associated with recurrent disease that overall survival does not and is thus most relevant to those cancers with long periods between recurrence and death (e.g., local or regionally advanced cancers of the breast, prostate, colon, and rectum). Indeed, the Food and Drug Administration has accepted disease-free survival as a regulatory endpoint that demonstrates clinical benefit for adjuvant therapy.

We are unaware of previous attempts to measure disease-free survival through Medicare claims. This absence is not due to an inability to measure survival—the accuracy of CMS...
vital status information is well established (7). Rather, it is the result of ambiguity regarding the ability of claims to capture cancer recurrence. As a preliminary study to determine the accuracy with which Medicare claims data capture disease-free survival in elderly breast cancer patients, we developed an algorithm for measuring both disease recurrence and death using Medicare data files and compared the results with an external gold-standard measure of disease-free survival, Cancer and Leukemia Group B (CALGB) clinical trial data.

The model cohort included all patients aged ≥65 years (n = 52) who consented, enrolled, and were treated on CALGB trial 9344, “Doxorubicin dose escalation, with or without Taxol, as part of the CA adjuvant regimen for node positive breast cancer” between January 1, 1995, and December 31, 1997. These 52 patients represented only 5.5% (52 of 944) of the full enrollment during this period. Despite what seems like a small percentage of elderly enrollees, the trial was chosen because the absolute value of enrolled elderly was among the largest of all phase 3 trials during that period.

We then carefully linked the patients’ CALGB clinical trial data (e.g., disease-free survival information) to their CMS Medicare claims files (i.e., denominator, Carrier, Outpatient, and MedPAR files) from enrollment through December 31, 2000 (8). We were able to match 51 (98%) of the 52 patients to Medicare files, a rate consistent with previous studies (9). Six patients were excluded because they were enrolled in health maintenance organizations and their claims were not processed through CMS (n = 3) or because they were not enrolled in Medicare part B (n = 3). Thus, the final analytic sample contained 45 patients.

This study was approved by the University of Chicago and Massachusetts General Hospital institutional review boards and conducted in compliance with their regulations. Data quality was ensured by careful review of data by CALGB Statistical Center staff and by the study chairperson. Statistical analyses were approved and confirmed by CALGB statisticians. All analyses were two-sided and performed using STATA version 8 SE; P<.05 was considered statistically significant.

We developed and ultimately refined a clinically intuitive algorithm to measure disease-free survival that required screening patients’ Medicare claims from the calendar date of their enrollment on CALGB 9344 forward through their last Medicare claim up to December 31, 2000 (the last day of available claims files), for evidence of cancer relapse or death. In clinical medicine, cancers are described according to primary anatomic sites (i.e., where the tumor originates) and secondary anatomic sites (i.e., location of tumor spread). The primary anatomic site is well documented in Surveillance, Epidemiology, and End Results (SEER) (10) and may be reasonably well documented in CMS files (11). However, whether (or when) the tumor has spread to anatomically distinct sites is not reported or measured by SEER, and it is not known whether such events are captured reliably in CMS claims. Because CMS claims use International Classification of Diseases 9th Revision–Clinical Modification (ICD-9-CM) diagnostic codes that do have distinct values for primary and secondary cancer sites, it is at least theoretically possible that CMS claims distinguish primary anatomic malignancy sites from secondary sites of primary cancer spread.
In CMS files, we identified cancer relapse by evaluating MedPAR, Carrier, and Outpatient files for ICD-9 diagnostic codes indicating secondary sites of cancer (Table 1) and then dated the relapse by “claim through date.” From the universe of ICD-9-CM secondary sites of cancer codes, we omitted the code 198.2 indicating “secondary malignant neoplasm of the breast” to avoid erroneous overcoding of patients’ primary breast cancer as “relapse” and ultimately omitted the code 198.89 “secondary malignant neoplasm of unspecified site” because of its high false-positive rate. Date of death was obtained from CMS denominator files. The first of the two possible events (i.e., relapse or death) was chosen to represent disease-free survival according to CMS claims. Individuals without either event were censored at the date of their last CMS claim with a maximum date of December 31, 2000.

We compared CMS disease-free survival information with the gold standard of CALGB disease-free survival information (censored at December 31, 2000) to calculate sensitivity, specificity, and area under the receiver operator curve of the disease-free survival indicator during the entire follow-up period and at traditional trial–censoring points (i.e., 2 years and 5 years). We defined sensitivity as the proportion of the patients known (according to CALGB data) to have either died or relapsed who were correctly identified through CMS claims as such. We defined specificity as the proportion of the patients known (according to CALGB data) to be both alive and without disease relapse who were correctly identified through CMS claims as such. The global performance of the measures was summarized by the area under the receiver operator curve (graph of sensitivity versus 1 – specificity); the greater the area under the receiver operator curve (maximum 1.00), the better the discriminatory accuracy of the measure.

The sample was exclusively female, the mean age was 69.6 years (standard deviation ± 4.2 years), and 82% were white, 11% black, and 7% Hispanic. During the entire follow-up period (i.e., a maximum of 2162 days in the CALGB data source and 2142 days in the CMS data source), 15 (33%) of the 45 patients had relapsed or had died according to CMS claims. By comparison, the gold-standard source of survival information noted, as a first event, relapse or death in 14 (31%) of 45 ($P = .84$) (Supplementary Figure, available at http://jncicancerspectrum.oxfordjournals.org/jnci/content/vol98/issue18). The 5-year disease-free survival sensitivity (100%, 95% confidence interval [CI] = 83% to 100%), specificity (97%, 95% CI = 87% to 100%), and area under the receiver operator curve (97%, 95% CI = 90% to 100%) were favorable, each being more favorable than 2-year disease-free survival (sensitivity = 83%, 95% CI = 36% to 100%; specificity = 95%, 95% CI = 83% to 100%; and area under the receiver operator curve = 84%, 95% CI = 66% to 100%) (Table 2).

We compared results of the CMS algorithm to the gold-standard data source with respect to the three components of disease-free survival (i.e., relapse, death, and censoring) at 5 years. According to the gold-standard data source, 86% (12 of 14) of the disease-free survival failures were the result of relapsed disease and 14% (2 of 14) were the result of death (Table 3). The ICD-9-CM codes for “secondary malignant neoplasm” applied to CMS data correctly categorized 11 of these 12 relapsed patients in CALGB data (sensitivity = 92%, 95% CI = 66% to 100%) and correctly categorized 31 of the 33 nonrelapsed patients (specificity = 94%, 95% CI = 82% to 99%).

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We measured the patient-level difference in the gold-standard and CMS algorithm time variables. The median difference in CALGB compared with CMS time was 40 days, and the inter-quartile range was 12–86 days. The mean difference was 99 days (95% CI = 4.5 to 193 days).

This preliminary validation study of Medicare claims shows that for elderly Medicare beneficiaries with histories of lymph node–positive breast cancer who were treated on a randomized phase 3 CALGB adjuvant chemotherapy trial, contemporaneous Medicare claims files reflect subsequent 5-year disease-free survival with a high degree of accuracy. The algorithm appears to also distinguish relapse from death within the domain of disease-free survival, which suggests that the secondary anatomic site codes may capture cancer recurrence. Taken with results of prior work documenting the ability of CMS claims to accurately measure certain anticancer therapies (6,12,13), these results suggest that CMS data capture some key elements of the clinical trial paradigm, and thus existing CMS-based data sources (e.g., SEER-Medicare) may be leveraged to yield clinical information regarding the effectiveness of adjuvant cancer treatments in the general population of elderly with solid tumors. Future research will focus on confirming these findings in a larger and more diverse patient sample.

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References

1. Trimble EL, Carter CL, Cain D, Freidlin B, Ungerleider RS, Friedman MA. Representation of older patients in cancer treatment trials. Cancer. 1994; 74(Suppl):2208–14. [PubMed: 8087794]
2. Murthy VH, Krumholz HM, Gross CP. Participation in cancer clinical trials: race-, sex-, and age-based disparities. JAMA. 2004; 291:2720–6. [PubMed: 15187053]
3. Hutchins LF, Unger JM, Crowley JJ, Coltman CA Jr, Albain KS. Underrepresentation of patients 65 years of age or older in cancer-treatment trials. N Engl J Med. 1999; 341:2061–7. [PubMed: 10615079]
4. Lewis JH, Kilgore ML, Goldman DP, Trimble EL, Kaplan R, Montello MJ, et al. Participation of patients 65 years of age or older in cancer clinical trials. J Clin Oncol. 2003; 21:1383–9. [PubMed: 12663731]
5. Lamont EB, Lauderdale DS, Schilsky RL, Christakis NA. Construct validity of medicare chemotherapy claims: the case of 5FU. Med Care. 2002; 40:201–11. [PubMed: 11880793]
6. Lamont EB, Herndon JE 2nd, Weeks JC, Henderson IC, Lilenbaum R, Schilsky RL, et al. Criterion validity of Medicare chemotherapy claims in Cancer and Leukemia Group B breast and lung cancer trial participants. J Natl Cancer Inst. 2005; 97:1080–3. [PubMed: 16030306]
7. Cowper DC, Kubal JD, Maynard C, Hynes DM. A primer and comparative review of major US mortality databases. Ann Epidemiol. 2002; 12:462–8. [PubMed: 12377423]
8. Henderson IC, Berry DA, Demetri GD, Cirrincione CT, Goldstein LJ, Martino S, et al. Improved outcomes from adding sequential Paclitaxel but not from escalating Doxorubicin dose in an adjuvant chemotherapy regimen for patients with node-positive primary breast cancer. J Clin Oncol. 2003; 21:976–83. [PubMed: 12637460]
9. Hatten J. Medicare’s common denominator: the covered population. Health Care Financ Rev. 1980; 2:53–64. [PubMed: 10309331]
10. Zippin C, Lum D, Hankey BF. Completeness of hospital cancer case reporting from the SEER Program of the National Cancer Institute. Cancer. 1995; 76:2343–50. [PubMed: 8635041]
11. Nattinger AB, Laud PW, Bajorunaite R, Sparapani RA, Freeman JL. An algorithm for the use of Medicare claims data to identify women with incident breast cancer. Health Serv Res. 2004; 39(Pt 1):1733–49. [PubMed: 15533184]
12. Virnig BA, Warren JL, Cooper GS, Klabunde CN, Schussler N, Freeman J. Studying radiation therapy using SEER-Medicare-linked data. Med Care. 2002; 40(Suppl):IV-49–54.
13. Cooper GS, Virnig B, Klabunde CN, Schussler N, Freeman J, Warren JL. Use of SEER-Medicare data for measuring cancer surgery. Med Care. 2002; 40(Suppl):IV-43–8.
### Table 1

Algorithm for measuring disease-free survival in Centers for Medicare and Medicaid Services (CMS) claims files

| Code type | Value | Description |
|-----------|-------|-------------|
| ICD-9 Dx* | 197.0 | Secondary malignant neoplasm of the lung |
|           | 197.1 | Secondary malignant neoplasm of the mediastinum |
|           | 197.2 | Secondary malignant neoplasm of the pleura |
|           | 197.3 | Secondary malignant neoplasm of other respiratory organs |
|           | 197.4 | Secondary malignant neoplasm of the small intestine, including duodenum |
|           | 197.5 | Secondary malignant neoplasm of the large intestine and rectum |
|           | 197.6 | Secondary malignant neoplasm of the retroperitoneum and peritoneum |
|           | 197.7 | Secondary malignant neoplasm of the liver |
|           | 197.8 | Secondary malignant neoplasm of the other digestive organs and spleen |
|           | 198.0 | Secondary malignant neoplasm of the kidney |
|           | 198.1 | Secondary malignant neoplasm of other urinary organs |
|           | 198.2 | Secondary malignant neoplasm of the skin |
|           | 198.3 | Secondary malignant neoplasm of the brain and spinal cord |
|           | 198.4 | Secondary malignant neoplasm of the other parts of the nervous system |
|           | 198.5 | Secondary malignant neoplasm of the bone and bone marrow |
|           | 198.6 | Secondary malignant neoplasm of the ovary |
|           | 198.7 | Secondary malignant neoplasm of the adrenal gland |
|           | 198.8 | Secondary malignant neoplasm of other sites |
|           | 198.82| Secondary malignant neoplasm of the genital organs |

Death indicator† | Date of death | Field contains date of death of Medicare beneficiary |

*ICD-9 = International Classification of Diseases 9th Revision codes applied to CMS ambulatory and hospital files (i.e., Carrier, Outpatient, MedPAR).

†CMS denominator file.
Table 2
Test characteristics of the Centers for Medicare and Medicaid Services (CMS) disease-free survival algorithm

| Censoring Period | Sensitivity (95% CI) | Specificity (95% CI) | Area under ROC (95% CI) |
|------------------|----------------------|----------------------|------------------------|
| 2 y              | 83% (36% to 100%)    | 95% (83% to 100%)    | 84% (66% to 100%)      |
| 5 y              | 100% (81% to 100%)   | 97% (83% to 100%)    | 97% (90% to 100%)      |

* The test characteristics were estimated by comparing CMS files with gold-standard clinical trial data pertaining to 45 elderly women with lymph node–positive breast cancer treated on the adjuvant chemotherapy trial CALGB 9344. CI = confidence interval; ROC = receiver operator curve.
Table 3

Five-year disease-free survival (disease-free survival) censoring proportions for Cancer and Leukemia Group B (CALGB) versus Centers for Medicare and Medicaid Services (CMS) data

| Censoring category | CMS data | CALGB gold-standard data |
|--------------------|----------|--------------------------|
| disease-free survival = 1 |
| Relapsed           | 13       | 12                       |
| Dead               | 2        | 2                        |
| disease-free survival = 0 |
| Censored           | 30       | 31                       |
| Total              | 45       | 45                       |