Developing a cancer-specific trigger tool to identify treatment-related adverse events using administrative data

Saul N. Weingart1,2,3 | Jason Nelson4 | Benjamin Koethe4 | Omar Yaghi1 | Stephan Dunning3 | Albert Feldman3 | David M. Kent1,2,4 | Allison Lipitz-Snyderman5

1Tufts Medical Center, Boston, MA, USA
2Department of Medicine, Tufts University School of Medicine, Boston, MA, USA
3OptumLabs, Cambridge, MA, USA
4Predictive Analytics and Comparative Effectiveness Center, Tufts University School of Medicine, Boston, MA, USA
5Memorial Sloan Kettering Cancer Center, New York, NY, USA

Correspondence
Saul N. Weingart, Tufts Medical Center, 800 Washington St., Boston, MA 02111, USA.
Email: sweingart@tuftsmedicalcenter.org.

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Abstract

Background: As there are few validated tools to identify treatment-related adverse events across cancer care settings, we sought to develop oncology-specific “triggers” to flag potential adverse events among cancer patients using claims data.

Methods: 322,887 adult patients undergoing an initial course of cancer-directed therapy for breast, colorectal, lung, or prostate cancer from 2008 to 2014 were drawn from a large commercial claims database. We defined 16 oncology-specific triggers using diagnosis and procedure codes. To distinguish treatment-related complications from comorbidities, we required a logical and temporal relationship between a treatment and the associated trigger. We tabulated the prevalence of triggers by cancer type and metastatic status during 1-year of follow-up, and examined cancer trigger risk factors.

Results: Cancer-specific trigger events affected 19% of patients over the initial treatment year. The trigger burden varied by disease and metastatic status, from 6% of patients with nonmetastatic prostate cancer to 41% and 50% of those with metastatic colorectal and lung cancers, respectively. The most prevalent triggers were abnormal serum bicarbonate, blood transfusion, non-contrast chest CT scan following radiation therapy, and hypoxemia. Among patients with metastatic disease, 10% had one trigger event and 29% had two or more. Triggers were more common among older patients, women, non-whites, patients with low family incomes, and those without a college education.

Conclusions: Oncology-specific triggers offer a promising method for identifying potential patient safety events among patients across cancer care settings.

Keywords: adverse event, epidemiology, oncology, patient safety, quality of care, trigger tool

1 | INTRODUCTION

Oncology care is an extraordinarily high-risk activity, given the nature of the disease and its toxic therapies. While medical oncology was at the epicenter of the patient safety revolution with the 1994 overdose of Dana-Farber Cancer Institute patient Betsy Lehman, there has been surprisingly little reliable and consistent information about patient safety in cancer care. Only three high-quality studies of chemotherapy errors have been published to date, and virtually all patient safety-related research performed in oncology settings has been conducted in regional or national referral centers. A literature review
examing high-quality studies of medication errors related to chemotherapy concluded that our ability to measure errors and injuries across the continuum of cancer care is poor at best. A variety of factors account for the dearth of robust research studies in cancer patient safety, including the physiologic vulnerability of cancer patients and the expected toxicities of many cancer-directed therapies. Though successful in various medical settings and in flagging potential diagnostic delays, attempts to identify treatment-related complications using so-called “trigger tools” have worked poorly in cancer care. An oncology trigger tool piloted in the UK National Health Service showed poor performance characteristics, a rigorous French study examining a 22-item trigger tool for adverse drug events showed low positive predictive values (PPVs), and a Danish cancer center study showed disappointing interrater agreement, even with use of expert chart reviewers.

Without a robust measurement approach to patient safety in oncology that works across the continuum of oncology care, it is difficult to advise patients and their clinicians about the likely toxicities of therapy, the risk of treatment-related errors, or the best site of care for their disease. Better measurement of adverse events (AEs) and medical errors could help medical and cancer center leaders to identify opportunities for improvement and inform programmatic priorities for policy makers. Most health-care organizations use quality metrics appropriate for general medical patients to describe the quality of oncology care, but the applicability of commonly used metrics such as infection rates and readmissions apply poorly to oncology care. Efforts to assess cancer programs based on cancer registry data are limited to a small subset of analytic cases and outdated information. Creating a more streamlined and accessible approach to patient safety measurement for oncology would develop significant social value.

To address this problem, a team of researchers, oncology practitioners, and quality measurement and patient safety experts developed a set of oncology-specific triggers using clinical data from patients at Memorial Sloan Kettering Cancer Center (MSK) undergoing an initial course of cancer-directed therapy. Trigger tools use indicators, such as antidote medications, abnormal laboratory parameters, “stat” medication orders, and changes in the level of care, to signal the presence of a medical error or iatrogenic injury. Unlike previously published studies that failed to validate oncology-specific trigger tools, the MSK team identified 49 high-value oncology triggers with an overall PPV of 0.48 for AEs and 0.18 for preventable events using physician chart review as the gold standard.

We undertook the present project in order to further develop the use of oncology-specific triggers to identify treatment-related AEs. Our project had three specific aims: (a) to construct a claims-based trigger tool capturing the MSK triggers as International Classification of Diseases (ICD) and Current Procedural Terminology (CPT) codes, and (b) to examine the prevalence of trigger events among a commercial claims cohort. We hypothesized that it would be feasible to create a cancer-specific claims-based trigger tool, and that the prevalence of trigger events would vary by cancer type and metastatic status.

2 METHODS

2.1 Subjects

We selected a cohort of patients undergoing an initial course of cancer-directed therapy for breast, lung, colorectal, and prostate cancers using the OptumLabs® Data Warehouse (OLDW). OLDW includes de-identified administrative claims and electronic health record (EHR) data on over 200 million patients, including claims for inpatient and ambulatory care for commercial and Medicare Advantage enrollees. It includes limited patient demographic information drawn from enrollment records. Socioeconomic status information in OLDW, including race/ethnicity, household income, and educational attainment, are imputed variables sourced from a national supplier of consumer marketing data. Mortality status is ascertained in OLDW through multiple sources including the Social Security Death Index, inpatient discharge status, and electronic medical records. We used ICD and CPT codes to select patients with cancer diagnoses who received cancer-specific therapies including surgery, radiation therapy, or chemotherapy (infusion as well as oncolytic or hormonal therapies). Inclusion criteria included a new cancer diagnosis of breast, lung, colorectal, and prostate cancers from 1 January 2008 through 31 December 2014, with initiation of a cancer-specific surgery, radiation, or chemotherapy during that period. To ensure a new cancer diagnosis, subjects with cancer diagnoses or treatments in 2005-2007 and those with a cancer recurrence code were excluded.

We abstracted sociodemographic characteristics (including age, gender, race/ethnicity, insurance (commercial, Medicare managed care), household income, and educational attainment), cancer diagnosis, and cancer-specific therapies from the claims database, excluding cases of male breast cancers and subjects under age 18. We used a modified algorithm that excludes cancer as a comorbidity to calculate each patient’s Charlson comorbidity index, an algorithm developed by Whyte and colleagues to classify cancer metastatic status, and the number of unplanned hospital admissions and inpatient days as an additional indicator of individuals at high-risk of harm. We abstracted the dates associated with diagnosis and treatment codes, hospitalizations, and vital status.

2.2 Measurements

To define a set of oncology-specific triggers, we identified ICD and CPT codes corresponding to 16 of the 23 highest
PPV triggers from the MSK developmental study (Table 1). Triggers included events such as neutropenic fever, abnormal serum potassium or bicarbonate, return to the operating room or interventional suite within 30 days of surgery, initiation of therapeutic anticoagulation, and nephrology consultation. We recognized some inherent ambiguity in the use of ICD codes, as certain codes denote nonspecific laboratory abnormalities (eg, 790.6).

To distinguish between complications related to a cancer-specific treatment rather than the patient’s cancer or non-cancer comorbidities, we required a logical and temporal relationship between each trigger and its likely cause. We assumed that each trigger event would be temporally related to a specific exposure and that it would persist for a limited period of time. For example, neutropenic fever was associated with chemotherapy but not surgery or radiation. We assumed that neutropenic fever would follow within 30 days of chemotherapy and expect to persist for no more than 30 days.

Recognizing the diversity of therapies and therapeutic regimens, we consulted with oncology clinicians to make generic assumptions about the most common and likely relationship of triggers and exposures, as shown in the Appendix.

### 2.3 Analyses

We characterized the cohort by sociodemographic and clinical characteristics and cancer-specific treatments (surgery, radiation, and/or chemotherapy), stratified by cancer type (breast, colorectal, lung, prostate) and metastatic status.

We then tabulated the number and percent of patients with each AE trigger during a 1-year period beginning with the date of the initial cancer-directed therapy. We tabulated the number and percent of patients with no trigger events, one event, and two or more events. We performed separate analyses by cancer type and metastatic status. We

### Table 1 Coding algorithm for selected oncology-specific triggers

| Trigger | Coding algorithm* |
|---------|-------------------|
| General care |                     |
| Pressure ulcer | ICD9 707.x (exclude 707.21 and 707.22) |
| Return to the operating room or interventional radiology within 30 d of surgery | See Osborne NH, et al |
| Vital signs |                     |
| Low oximetry results (SaO2 < 88%) | ICD9 799.02 |
| Fever (> 38.2°C) | ICD9 780.6 and ICD9 288.x |
| Orders |                     |
| Blood transfusion | ICD9 V58.2, CPT 36 430 |
| Contact precautions/order for isolation | ICD9 V07.x |
| Nasogastric tube (not in operating room) | ICD9 96.07, CPT 43 753 |
| Non-contrast chest CT after radiation to the chest | CPT 71 250 |
| Percutaneous drain placement | ICD9 54.91, CPT 32 557 |
| Laboratories |                     |
| Abnormal serum bicarbonate (< 18, > 36 mEq/L) | ICD9 790.6, 276.2, 276.3 |
| Abnormal serum potassium (> 6, < 2.5 mEq/L) | ICD9 276.8, 276.7 |
| Clostridium difficile toxin positive | ICD9 008.45 |
| Elevated creatinine > 1 mg/dL and 50% greater than baseline | ICD9 584.9 acute kidney injury (not present on admission) |
| Positive blood culture without contaminant | ICD9 790.7 bacteremia, CPT 87 040, 87 103 |
| Medication-related |                     |
| Initiation of therapeutic anticoagulation | medications: warfarin, enoxaparin, apixaban, rivaroxaban, dabigatran, fondaparinux, edoxaban |
| Consultations | E&M visit (outpatient CPT 99241-99245, inpatient CPT 99251-99255) and provider_specialty = ‘nephrologist’ |

*ICD9 codes mapped to ICD10

### Note:
Osborne NH, Nicholas LH, Ryan AM, Thumma JR, Dimick JB. Association of hospital participation in a quality reporting program with surgical outcomes and expenditures for Medicare beneficiaries. *JAMA* 2015; 313:496-504.

Abbreviations: CPT, Current Procedure Terminology; CT, computed tomography; ICD, International Classification of Diseases.
examined the prevalence of trigger events by sociodemographic and clinical characteristics, using the Chi-square and Wilcoxon rank-sum test for categorical and continuous variables, respectively.

Analyses used SAS 9.4 for Windows (SAS Institute) and R 3.4.3 (The R Foundation). The study protocol was reviewed in advance by the Tufts Health Sciences Institutional Review Board (IRB) and determined to be exempt from human subjects review due to the use of a de-identified dataset.

3 | RESULTS

3.1 | Cohort characteristics

The study cohort included 322,887 unique subjects with breast, colorectal, lung, and prostate cancers (Table 2). The mean age was 64, consistent with a commercially insured patient population. Males comprised a greater percentage of patients with colorectal and lung cancers than women. While the majority of patients were white, Asians, blacks, and Hispanics were also present. There was missing data regarding race/ethnicity, household income, and education for at least one-third of the cohort.

Overall, 27% of patients had metastatic cancer. The percent of patients with metastatic disease varied from 22% for breast cancer to 59% for lung cancer. The cancer-specific Charlson index suggested a moderate burden of comorbid non-cancer illness. Forty-three percent of patients received multimodality cancer therapy.

3.2 | Trigger prevalence

Cancer-specific trigger events were common, affecting 19% of patients over the initial 1-year course of therapy (Table 3). The trigger burden varied by disease and metastatic status. Among patients with nonmetastatic disease, the prevalence of trigger events was greatest among patients with lung (33%) and colorectal (19%) cancers and least among those with prostate (6%) and breast (10%) cancers—likely a reflection of treatment types and toxicities as well as patients’ underlying physiologic reserve. There was a similar, but amplified, pattern among patients with metastatic disease. There was a particularly heavy burden of trigger events among those with lung (50%) and colorectal (41%) cancers, although patients with metastatic breast (31%) and prostate (25%) cancers also experienced significant treatment-related morbidity. The most prevalent triggers were abnormal serum bicarbonate, blood transfusion, non-contrast chest CT scan following radiation therapy, hypoxemia, contact precautions, neutropenic fever, and abnormal serum potassium.

3.3 | Multiple triggers

Certain patients experienced a particularly high number of trigger events, although it is important to note that a single adverse event could give rise to multiple triggers. As shown in Table 4, 19% of patients had at least one event. Among patients with nonmetastatic disease, 4% had one trigger event over the course of the year and 8% had two or more. Among those with metastatic disease, 10% had one trigger event and 29% had two or more. Individual patients with lung cancer had a particularly high burden of trigger events; one-quarter of patients with nonmetastatic disease and one-third of those with advanced disease experienced multiple triggers.

3.4 | Risk factors associated with triggers

Table 5 displays trigger prevalence by subject characteristics, stratified by cancer type. Triggers were less prevalent among young patients, men, whites, families with incomes over $150,000 per year, and patients with some college education. These differences were small but statistically significant ($P < .001$), perhaps reflecting certain patients’ better access to care, earlier cancer detection, and lower intensity therapy.

4 | DISCUSSION

In this retrospective cohort study of 322,887 patients with breast, lung, colorectal, and prostate cancer treated for an initial course of cancer-directed therapy, we found that one in five patients had a “trigger” event that indicated a likely treatment-related AE. The most common triggers included laboratory abnormalities of bicarbonate and potassium, need for blood transfusion, hypoxemia, neutropenic fever, and contact precautions. The burden of event triggers fell disproportionately on patients with lung and colorectal cancer compared to those with breast or prostate cancer, and among those with metastatic disease. The prevalence of trigger events among patients with metastatic disease was more than triple the rate among those with nonmetastatic disease (39.1% vs 12.0%), and as high as 50.2% in patients with metastatic lung cancer. Nearly three in four patients with a trigger had two or more such events.

Triggers are clinical indicators that signal the possibility of treatment-related injury, and therefore, the trigger rate needs to be adjusted by the probability that the trigger denoted an actual harm event. The general medicine literature describes PPVs of 17%-45% based on physician chart review as the gold standard.9,10,27-30 Since the number of trigger events may overestimate the number of actual AEs,
we estimated the incidence of AEs and preventable AEs present in the cohort using PPVs calculated in the original MSK medical record review-based developmental study. Using the MSK trigger-specific PPVs, the present study found an estimated 97,521 AEs and 24,915 preventable AEs affecting 30.2% and 7.7% of cancer patients in this cohort, respectively.

Direct comparison of our findings with research on AEs in cancer care is challenging, given the use of inconsistent and disparate research methods. Comparison with toxicity rates in

### TABLE 2  Cohort characteristics

| Characteristic                        | Breast | Colorectal | Lung   | Prostate | Overall |
|--------------------------------------|--------|------------|--------|----------|---------|
| N                                    | 124,253| 52,383     | 51,311 | 94,940   | 322,887 |
| Age [mean (SD)]                      | 59.5 (12.1) | 63.2 (12.5) | 67.1 (10.5) | 66.9 (9.1) | 63.5 (11.6) |
| Sex [n (%)]                          | Male   | 27,616 (52.7%) | 27,170 (53.0%) | 94,940 (100.0%) | 149,726 (46.4%) |
|                                      | Female | 124,253 (100.0%) | 24,767 (47.3%) | 24,141 (47.0%) | 173,161 (53.6%) |
| Race/Ethnicity [n (%)]               |        |            |        |          |         |
| Missing/Unknown                      | 37,198 (29.9%) | 18,961 (36.2%) | 19,432 (37.9%) | 32,741 (34.5%) | 108,332 (33.6%) |
| Asian                                | 2398 (1.9%) | 862 (1.6%)   | 634 (1.2%)   | 1019 (1.1%)   | 4913 (1.5%)   |
| Black                                | 9,543 (7.7%) | 3,847 (7.3%)  | 3,538 (6.9%)  | 7,920 (8.3%)  | 24,848 (7.7%) |
| Hispanic                             | 5,491 (4.4%) | 2,360 (4.5%)  | 1,271 (2.5%)  | 3,463 (3.6%)  | 12,585 (3.9%) |
| White                                | 69,623 (56.0%) | 26,353 (50.3%) | 26,436 (51.5%) | 49,797 (52.5%) | 172,209 (53.3%) |
| Annual household income [n (%)]      |        |            |        |          |         |
| Unknown                              | 44,997 (36.2%) | 22,584 (43.1%) | 23,569 (45.9%) | 37,349 (39.3%) | 128,499 (39.8%) |
| <$25K                                | 15,627 (12.6%) | 6,713 (12.8%)  | 8,845 (17.2%)  | 10,733 (11.3%) | 41,918 (13.0%) |
| $24K - $149K                         | 19,793 (15.9%) | 8,511 (16.2%)  | 8,488 (16.5%)  | 15,694 (16.5%) | 52,416 (16.2%) |
| $150K - 249K                         | 23,057 (18.6%) | 8,367 (16.0%)  | 6,551 (12.8%)  | 17,314 (18.2%) | 55,289 (17.1%) |
| $250K - $499K                        | 12,336 (9.9%) | 3,964 (7.6%)   | 2,553 (5.0%)   | 8,514 (9.0%)   | 27,367 (8.5%) |
| $500K+                               | 8,513 (6.9%) | 2,244 (4.3%)   | 1,305 (2.5%)   | 5,336 (5.6%)   | 17,398 (5.4%) |
| Education [n (%)]                    |        |            |        |          |         |
| Missing/Unknown                      | 34,322 (27.6%) | 17,925 (34.2%) | 18,449 (36.0%) | 30,698 (32.3%) | 101,394 (31.4%) |
| Less than 12th grade                 | 217 (0.2%) | 141 (0.3%)   | 89 (0.2%)     | 149 (0.2%)     | 596 (0.2%)     |
| High school diploma                 | 20,749 (16.7%) | 9,891 (18.9%)  | 10,987 (21.4%) | 16,468 (17.3%) | 58,095 (18.0%) |
| Less than bachelor degree           | 48,901 (39.4%) | 18,423 (35.2%) | 17,365 (33.8%) | 35,020 (36.9%) | 119,709 (37.1%) |
| Bachelor degree plus                | 20,064 (16.1%) | 6,003 (11.5%)  | 4,421 (8.6%)  | 12,605 (13.3%) | 43,093 (13.3%) |
| Insurance type [n (%)]              |        |            |        |          |         |
| Private insurance                   | 99,932 (80.4%) | 42,456 (81.0%) | 37,196 (72.5%) | 69,486 (73.2%) | 249,070 (77.1%) |
| Medicare Advantage                  | 24,321 (19.6%) | 9,927 (19.0%)  | 14,115 (27.5%) | 25,454 (26.8%) | 73,817 (22.9%) |
| Clinical characteristics             |        |            |        |          |         |
| Metastatic disease [n (%)]          | 26,791 (21.6%) | 18,671 (35.6%) | 30,169 (58.8%) | 10,800 (11.4%) | 86,431 (26.8%) |
| Charlson index* [mean (SD)]         | 2.0 (1.6) | 2.8 (1.9)   | 3.6 (1.8)    | 2.8 (1.4)    | 2.6 (1.7)    |
| Rehospitalized w/in 1 y [n (%)]     | 36,780 (29.6%) | 33,047 (63.1%) | 32,092 (62.5%) | 37,798 (39.8%) | 139,717 (43.3%) |
| Hospital days [mean (SD)]           | 5.1 (8.8) | 10.4 (13.9) | 10.4 (12.9) | 4.1 (7.9) | 7.3 (11.4) |
| Treatment type [n (%)]              |        |            |        |          |         |
| Chemotherapy only                   | 18,357 (14.8%) | 8,724 (16.7%) | 11,501 (22.4%) | 18,830 (19.8%) | 57,412 (17.8%) |
| Radiation only                      | 11,407 (9.2%) | 1273 (2.4%)  | 7,260 (14.1%) | 21,253 (22.4%) | 41,193 (12.8%) |
| Surgery only                        | 22,952 (18.5%) | 22,770 (43.5%) | 6,818 (13.3%) | 31,598 (33.3%) | 84,138 (26.1%) |
| Multimodality                       | 71,537 (57.6%) | 19,616 (37.4%) | 25,732 (50.1%) | 23,259 (24.5%) | 140,144 (43.4%) |

Note: Percentages may not add to 100% due to rounding.

*NIH measure for cancer patients’ modification. https://healthcaredelivery.cancer.gov/seermedicare/considerations/calculation.html.
| Trigger                                      | Breast cancer n = 124,253 | Colorectal cancer n = 52,383 | Lung cancer n = 51,311 | Prostate cancer n = 94,940 |
|----------------------------------------------|----------------------------|-------------------------------|------------------------|---------------------------|
|                                               | Nonmetastatic n = 97,462  | Metastatic n = 26,791         | Nonmetastatic n = 33,712 | Metastatic n = 18,671      |
|                                               |                            |                               | Nonmetastatic n = 21,142 | Metastatic n = 30,169      |
|                                               |                            |                               | Nonmetastatic n = 84,140 | Metastatic n = 10,800      |
| Any trigger                                  | 10.2% (99,589/97,462)      | 30.9% (82,751/26,791)         | 19.2% (64,888/33,712)   | 41.4% (77,381/18,671)      |
|                                              | 32.8% (69,301/21,142)      | 50.2% (15,157/30,169)         |                        | 5.9% (49,878/84,140)       |
| Anticoagulation                              | 0.1% (77/87,345)           | 0.2% (56/25,501)              | 0.2% (63/32,893)        | 0.5% (97/18,217)           |
|                                              | 0.3% (49,177/97,799)       | 0.4% (109,26/26,252)          | 0.3% (49,177/97,799)    | 0.1% (49,638/869)          |
| Bacteremia/positive blood culture            | 0.3% (279/87,345)          | 1.4% (361/25,501)             | 1.1% (355/32,893)       | 3.1% (571/18,217)          |
|                                              | 1.6% (279/17,799)          | 2.4% (617/26,252)             | 1.6% (279/17,799)       | 0.3% (183,638/169)         |
| Abnormal serum bicarbonate                   | 1.8% (1575/87,345)         | 5.0% (1285/25,501)            | 3.9% (1287/32,893)      | 9.3% (1700/18,217)         |
|                                              | 4.7% (831/17,799)          | 7.3% (19,172/26,252)          | 1.9% (1239/63,869)      | 6.3% (619,9818)            |
| Blood transfusion                            | 1.9% (1682/87,345)         | 8.8% (2234/25,501)            | 2.6% (854/32,893)       | 9.6% (1740/18,217)         |
|                                              | 12.2% (2172/17,799)        | 23.2% (6087/26,252)           | 0.6% (399/63,869)       | 7.6% (749,9818)            |
| C. difficile positive                         | 0.3% (296/87,345)          | 1.1% (271/25,501)             | 1.1% (362/32,893)       | 2.5% (450/18,217)          |
|                                              | 1.3% (227/17,799)          | 1.8% (480/26,252)             | 0.2% (122/63,869)       | 0.7% (66,9818)             |
| Non-contrast chest CT following XRT          | 1.6% (847/53,272)          | 3.0% (496/16,658)             | 2.1% (128/6022)         | 4.1% (216/5229)            |
|                                              | 10.3% (977/9523)           | 11.3% (2339/20,735)           | 1.1% (422/37,290)       | 4.9% (218,4425)            |
| Elevated creatinine                          | 0.1% (131/97,462)          | 0.5% (136/26,791)             | 0.3% (853/32,893)       | 0.8% (158/18,217)          |
|                                              | 0.6% (125/21,142)          | 1.0% (315/30,169)             | 0.1% (67/84,140)        | 0.4% (43,10800)            |
| Hypoxemia/low oximetry                       | 1.0% (992,97/462)          | 4.0% (1078/26,791)            | 2.8% (937/32,893)       | 5.9% (1103/18,671)         |
|                                              | 12.2% (25,721/21,142)      | 16.3% (4927/30,169)           | 1.2% (105/84,140)       | 4.6% (493,10800)           |
| Contact precautions/isolation                | 6.0% (2630/43,825)         | 9.5% (2147/22,616)            | 4.1% (483/11,816)       | 6.0% (931/15,614)          |
|                                              | 6.2% (732/11,732)          | 8.0% (1976/24,672)            | 1.2% (367/29,511)       | 3.6% (322/8924)            |
| Nasogastric tube                             | < 11 events < 11 events    | < 11 events < 11 events       | < 11 events < 11 events | < 11 events < 11 events    |
| Nephrology consult                           | 0.3% (254/97,462)          | 1.3% (350/26,791)             | 1.4% (485/32,893)       | 3.1% (577/18,671)          |
|                                              | 1.3% (279/21,142)          | 2.1% (645/30,169)             | 0.6% (541/84,140)       | 2.6% (284,10800)           |
| Neutropenic fever                            | 4.0% (1765/43,825)         | 6.8% (1535/22,616)            | 4.5% (528/11,816)       | 5.7% (894/15,614)          |
|                                              | 6.7% (788/11,732)          | 7.4% (1814/24,672)            | 0.6% (173/29,511)       | 2.6% (231,8924)            |
| Percutaneous drain                           | 0% (25/68,318)             | 0.1% (18/15,425)              | 0.8% (206/27,078)       | 2.4% (252/10,600)          |
|                                              | 0.6% (48/8144)             | 0.5% (25/5160)                | 0.3% (96/38,298)        | 0.6% (12,2060)             |
| Abnormal serum potassium                     | 2.5% (2218/87,345)         | 8.7% (2211/25,501)            | 7.3% (2417/32,893)      | 17.7% (3233/18,217)        |
|                                              | 8.1% (1445/17,799)         | 13.9% (3656/26,252)           | 1.8% (1135/63,869)      | 6.9% (674,9818)            |
| Press ulcer                                  | 0.3% (230/68,318)          | 0.6% (85/15,425)              | 1.2% (313/27,078)       | 1.5% (154/10,600)          |
|                                              | 1.0% (78/8144)             | 0.9% (45/5160)                | 0.2% (95/38,298)        | 0.8% (17,2060)             |
| Return to OR or IR                           | 0.1% (63/68,318)           | 0.2% (37/15,425)              | 2.4% (638/27,078)       | 3.7% (392/10,600)          |
|                                              | 0.4% (33/8144)             | 0.4% (21/5160)                | 0.6% (229/38,298)       | 0.7% (14,2060)             |

Note: Values shown in the table are prevalence rates and the number of patients with a trigger within an exposure window divided by the number exposed in parentheses.
Table 4  Number of triggers per patient within 1 y, by cancer type and metastatic status

| Cancer type | N       | No trigger | 1 trigger | >1 triggers | No trigger | 1 trigger | >1 triggers | No trigger | 1 trigger | >1 triggers |
|-------------|---------|------------|-----------|-------------|------------|-----------|-------------|------------|-----------|-------------|
| Breast      | 124 253 | 89.8%      | 6.7%      | 3.5%        | 69.1%      | 16.6%     | 14.3%       | 85.3%      | 8.8%      | 5.9%        |
| Colorectal  | 52 383  | 80.8%      | 12.1%     | 7.1%        | 58.6%      | 20.3%     | 21.1%       | 72.8%      | 15.0%     | 12.1%       |
| Lung        | 51 311  | 67.2%      | 18.4%     | 14.4%       | 49.8%      | 23.8%     | 26.4%       | 57.0%      | 21.6%     | 21.4%       |
| Prostate    | 94 940  | 94.1%      | 4.3%      | 1.7%        | 75.4%      | 13.4%     | 11.1%       | 92.0%      | 5.3%      | 2.7%        |
| Total       | 322 887 | 88.0%      | 7.6%      | 4.4%        | 60.9%      | 19.5%     | 19.6%       | 80.7%      | 10.8%     | 8.4%        |

Note: Percentages may not add to 100% due to rounding.

In conclusion, a claims-based oncology-specific trigger tool appears both feasible to construct and instructive in its results. Treatment-related triggers are common in cancer care, suggesting a significant burden of anticipated and potentially unexpected and even preventable AEs. The trigger burden falls unevenly across patients by disease, metastatic status, treatment type, and socioeconomic status, affecting exactly those patients most vulnerable to harm. Additional research is needed to assess the association of cancer triggers with key clinical outcomes such as disease- attributable and overall mortality, resource utilization, and patient-centered outcomes. Oncology-specific triggers offer the opportunity to better understand and characterize the nature and extent of AEs in cancer care, and to inform interventions that may reduce the burden of harm among patients with cancer.

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**DATA AVAILABILITY STATEMENT**

The data that support the findings of this study are available from OptumLabs®. Restrictions apply to the availability of these data, which were used under license for this study.

### TABLE 5 Cohort characteristics by trigger prevalence

| Characteristic                      | No trigger | At least one trigger | Overall  |
|-------------------------------------|------------|----------------------|----------|
| **N**                               | 260,700    | 62,187               | 322,887  |
| **Age [mean(SD)]**                  | 63.3 (11.6)| 64.1 (11.8)          | 63.5 (11.6) |
| **Sex [n (%)]**                     |            |                      |          |
| Male                                | 123,171 (47.3%) | 26,555 (42.7%) | 149,726 (46.4%) |
| Female                              | 137,529 (52.7%) | 35,632 (57.3%) | 173,161 (53.6%) |
| **Race/Ethnicity [n (%)]**          |            |                      |          |
| Missing/Unknown                     | 86,541 (33.2%) | 21,791 (35.0%) | 108,332 (33.6%) |
| Asian                               | 4103 (1.6%) | 810 (1.3%)           | 4913 (1.5%) |
| Black                               | 19,738 (7.6%) | 5110 (8.2%) | 24,848 (7.7%) |
| Hispanic                            | 10,394 (4.0%) | 2191 (3.5%) | 12,585 (3.9%) |
| White                               | 139,924 (53.7%) | 32,285 (51.9%) | 172,209 (53.3%) |
| **Annual household income [n (%)]**|            |                      |          |
| Unknown                             | 101,758 (39.0%) | 26,741 (43.0%) | 128,499 (39.8%) |
| <$25K                               | 32,266 (12.4%) | 9652 (15.5%) | 41,918 (13.0%) |
| $24K-$149K                          | 42,022 (16.1%) | 10,394 (16.7%) | 52,416 (16.2%) |
| $150K-$249K                         | 46,225 (17.7%) | 9064 (14.6%) | 55,289 (17.1%) |
| $250K-$499K                         | 23,263 (8.9%) | 4104 (6.6%) | 27,367 (8.5%) |
| $500K+                              | 15,166 (5.8%) | 2232 (3.6%) | 17,398 (5.4%) |
| **Education [n (%)]**               |            |                      |          |
| Missing/Unknown                     | 80,855 (31.0%) | 20,539 (33.0%) | 101,394 (31.4%) |
| Less than 12th grade                | 456 (0.2%) | 140 (0.2%) | 596 (0.2%) |
| High school diploma                 | 45,256 (17.4%) | 12,839 (20.6%) | 58,095 (18.0%) |
| Less than bachelor degree           | 97,481 (37.4%) | 22,228 (35.7%) | 119,709 (37.1%) |
| Bachelor degree plus                | 36,652 (14.1%) | 6441 (10.4%) | 43,093 (13.3%) |
| **Insurance [n (%)]**               |            |                      |          |
| Private insurance [n (%)]           | 202,110 (77.5%) | 46,960 (75.5%) | 249,070 (77.1%) |
| Medicare Advantage [n (%)]          | 58,590 (22.5%) | 15,227 (24.5%) | 73,817 (22.9%) |
| **Clinical characteristics**        |            |                      |          |
| Metastatic disease [n (%)]          | 52,607 (20.2%) | 33,824 (54.4%) | 86,431 (26.8%) |
| Charlson index* [mean (SD)]         | 2.5 (1.7) | 3.0 (1.9) | 2.6 (1.7) |
| Rehospitalized w/in 1 y [n (%)]     | 97,642 (37.5%) | 42,075 (67.7%) | 139,717 (43.3%) |
| Hospital days [mean (SD)]           | 5.4 (8.6) | 11.7 (15.2) | 7.3 (11.4) |
| **Treatment type [n (%)]**          |            |                      |          |
| Chemotherapy only                   | 42,069 (16.1%) | 15,343 (24.7%) | 57,412 (17.8%) |
| Radiation therapy only              | 38,853 (14.9%) | 2340 (3.8%) | 41,193 (12.8%) |
| Surgery only                        | 78,754 (30.2%) | 5384 (8.7%) | 84,138 (26.1%) |
| Multimodality                       | 101,024 (38.8%) | 39,120 (62.9%) | 140,144 (43.4%) |

*NIH measure for cancer patients’ modification. https://healthcaredelivery.cancer.gov/seermedicare/considerations/calculation.html

Note: Percentages may not add to 100% due to rounding.
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### APPENDIX

#### Oncology triggers, by exposure and timing window

| Trigger | Exposure | Timing (days) | Rationale | Window (days) | Notes |
|---------|----------|---------------|-----------|---------------|-------|
| Anticoagulation | Surgery, chemotherapy | 30 | Treatment-related venous thromboembolism due to immobility, thrombophilia | 180 | Prolonged therapy (3-6 mo) for venous thromboembolism |
| Bacteremia/ positive blood culture | Surgery, chemotherapy | 30 | Post-op or neutropenia-related infection | 30 | Normally 14-d course, but this allows for complicated or device-related infections |
| Abnormal serum bicarbonate | Surgery, chemotherapy | 14 | Perioperative fluid management, chemotherapy-related toxicity | 14 | Same as abnormal serum potassium |
| Blood transfusion (1) | Surgery | 7 | Perioperative bleeding | 7 | Short window post-op |
| Blood transfusion (2) | Chemotherapy | 30 | Chemotherapy-related bone marrow toxicity | 30 | Longer time window |
| C. difficile positive | Surgery, chemotherapy | 30 | Post-antibiotic exposure | 30 | Toxin takes weeks to resolve, at least |
| Non-contrast chest CT following XRT | XRT | 30 | XRT-related inflammation | 30 | |
| Elevated serum creatinine | All | 30 | See Nephrology consult | 30 | See Nephrology consult |
| Hypoxemia/low oximetry (1) | Surgery | 7 | Fluid shifts | 7 | Post-op splinting or fluid shifts |
| Hypoxemia/low oximetry (2) | XRT | 30 | Radiation pneumonitis | 60 | |
| Hypoxemia/low oximetry (3) | Chemotherapy | 90 | Chemotherapy-related toxicity | 90 | Bleomycin toxicity can present late |
| Contact precautions/ isolation | Chemotherapy | 30 | Presumes infection | 30 | See Neutropenic fever |
| Nasogastric tube not placed in OR | Surgery | 7 | Postoperative ileus | 7 | In chemotherapy patients, tube more likely related to disease progression |
| Nephrology consult | All | 30 | Chemotherapy-related renal toxicity or surgery/XRT-related dehydration and azotemia, antibiotic toxicity | 30 | Could argue for shorter window, but debilitation and kidney injury may take time to resolve |

(Continued)
### APPENDIX (Continued)

| Trigger                  | Exposure                  | Timing (days) | Rationale                                    | Window (days) | Notes                                                                 |
|--------------------------|---------------------------|---------------|----------------------------------------------|---------------|----------------------------------------------------------------------|
| Neutropenic fever        | Chemotherapy              | 30            | Chemotherapy-related neutropenia             | 30            | Normally 14-dcourse, but this allows for complicated or device-related infections |
| Percutaneous drain       | Surgery, IR               | 30            | Post-procedural infection                    | 30            | May require extended treatment                                       |
| Abnormal serum potassium | Surgery, chemotherapy     | 14            | See Abnormal serum bicarbonate               | 14            | Shorter time window for surgery and longer for chemotherapy, but fluctuating values during a course of treatment warrant a 2-wk window |
| Pressure ulcer           | Surgery                   | 30            | Post-op pressure ulcer                       | 30            | Assumes that multiple returns to operating room may be needed to address staged procedure or infection |
| Return to OR or IR       | Surgery, interventional radiology | 30            | Post-procedural complication                | 30            |                                                                    |

Abbreviation: CT, computed tomography; IR, interventional radiology; OR, operating room; XRT, radiation therapy.