Risk and Healthcare Costs of Chemotherapy-Induced Neutropenic Complications in Women with Metastatic Breast Cancer

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Introduction

Breast cancer is the most common form of cancer among women in the US, and the second leading cause of cancer death [1]. It is estimated that 1 in every 8 women will develop breast cancer during her lifetime [1]. In 2009, an estimated 192,370 women were diagnosed with breast cancer, and 40,170 women died from the disease [1]. Approximately 6% of women with breast cancer have metastatic disease at initial presentation and an additional 20–40% of breast cancer patients develop metastatic disease at some point following diagnosis, often many years later. Median survival in women with metastatic breast cancer (MBC) is about 18–24 months, but many patients survive several years [2].

Key Words

Metastatic breast cancer • Febrile neutropenia • Costs and cost analysis

Abstract

Background: The burden of chemotherapy-induced neutropenic complications (CINC) in women with metastatic breast cancer (MBC) is largely unknown and may differ across cancer populations due to variation in the characteristics of patients, their disease and their treatment. Methods: This study employed a retrospective cohort design and US healthcare claims data (2003–2009). For each woman in the study database who received myelotoxic chemotherapy for MBC, the first observed course and each cycle within the course were characterized. Risk and healthcare costs of CINC – by care setting – were descriptively analyzed on an overall basis by chemotherapy cycle and chemotherapy regimen. Results: Among 2,620 study subjects, most received chemotherapy with cyclophosphamide/doxorubicin (25%), docetaxel (20%) or paclitaxel (12%). Thirty-one percent of subjects received colony-stimulating factors (CSF) prophylactically in their first chemotherapy cycle and an additional 13% first received CSF prophylaxis after cycle one. CINC developed in 11% of subjects; among these subjects, 88% required inpatient care and 45% experienced CINC in the first cycle of chemotherapy. For CINC requiring inpatient care, costs averaged USD 12,869 (95% CI: USD 12,622–13,116), and for CINC requiring outpatient care only, USD 2,030 (CI: USD 1,925–2,135). Conclusion: CINC is a clinically and economically important threat among women with MBC, and should be an important consideration in the treatment of this population.

cycle of chemotherapy. For CINC requiring inpatient care, costs averaged USD 12,869 (95% CI: USD 12,622–13,116), and for CINC requiring outpatient care only, USD 2,030 (CI: USD 1,925–2,135). Conclusion: CINC is a clinically and economically important threat among women with MBC, and should be an important consideration in the treatment of this population.

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Neutropenia is a common side effect of myelosuppressive chemotherapy. Neutropenia increases the risk of infection, which is typically signaled by fever. When neutropenic patients develop fever, i.e. febrile neutropenia (FN), the likelihood of infection and serious consequences often necessitates hospitalization for urgent evaluation, ongoing monitoring and administration of intravenous antibiotics [10, 11]. FN, as well as severe or prolonged neutropenia, can lead to dose delays, dose reductions, and/or chemotherapy discontinuations, interfering with the delivery of optimal treatment and possibly adversely affecting patient outcomes [11–14].

In several recent randomized trials of the efficacy and safety of combination versus sequential-agent chemotherapy in women with MBC, the risk of FN has been reported to be as high as 48%, although for many chemotherapy regimens, the risk of FN has been reported to be much lower [15–18]. Because the intent of chemotherapy is typically palliative in this patient population, the incidence of toxicities such as FN is an important consideration in the selection of chemotherapy regimens and, along with overall survival, should be considered as one of the key endpoints in making treatment decisions [15]. Moreover, because the benefits of chemotherapy for MBC – in terms of both extensions in life expectancy and an enhanced quality of life – are typically limited, the cost of FN and other associated toxicities is an especially important consideration in an era of increased emphasis on achieving an acceptable balance between the costs and benefits of medical interventions [16–20].
Chemotherapy Courses, Cycles and Regimens

For each patient in the source population, each unique cycle within the course of chemotherapy was identified. The first cycle was defined as beginning with the date of initiation of chemotherapy (i.e. the index date) and ending with the first service date for the next administration of chemotherapy (as evidenced by a medical claim with a corresponding HCPCS, ICD-9-CM or UB-92 code), occurring at least 14 days but no more than 59 days after the index date. If a second chemotherapy cycle did not commence prior to day 60 or if there was evidence of the receipt of radiation therapy (based on medical claims with relevant HCPCS, ICD-9-CM or UB-92 codes) during this period, both the first cycle and course of chemotherapy were considered to have been completed 30 days after the beginning of the cycle or on the day prior to initiation of radiation therapy, whichever occurred first. The second and all subsequent cycles of chemotherapy were similarly defined.

Chemotherapy agents were ascertained on a cycle-specific basis based on a review of all HCPCS Level II codes on medical claims with service dates within 6 days of the start of each cycle of chemotherapy. Regimens were categorized based on the combination of agents received in the first cycle. For patients in the source population with multiple courses of chemotherapy during the period of interest, only the first one was considered in analyses.

Exclusionary Criteria

Patients were excluded from the source population if they had: (1) ≥2 medical encounters at least 30 days apart with a diagnosis of another malignant neoplasm (ICD-9-CM 140.xx–208.xx), excluding malignant neoplasm of breast (174.xx), local or distant secondary malignant neoplasm (196.xx–199.xx), malignant neoplasm of skin other than melanoma (173.xx) or another malignant neoplasm if it was the same site as the metastases (e.g. malignant neoplasm of liver [155.0] if there was evidence of metastasis to liver [197.7], as defined above), (2) any gaps in their eligibility for comprehensive medical and drug benefits during the 1-year (‘pretreatment’) period prior to the initiation of chemotherapy, (3) evidence of hematopoietic stem-cell transplantation prior to or during receipt of chemotherapy or (4) any pharmacy claims for myelotoxic chemotherapy drugs or colony-stimulating factors (CSF) agents (precise dates of administration could not be ascertained for drugs dispensed through retail pharmacies) during the 30-day period before or during the receipt of chemotherapy.

Neutropenic Complications

Episodes

Neutropenic complications comprised encounters for the treatment of neutropenia (ICD-9-CM 288.0) as well as encounters for the treatment of related conditions. The latter comprised infections (list available from the authors upon request) and fever (ICD-9-CM 780.6).

Identifying hospitalization for CINC was based on admission for inpatient care with a diagnosis (principal or secondary) of neutropenia, fever or infection. Hospitalizations were identified on a cycle-specific basis using facility inpatient claims with admission dates any time between day 6 and the last day of the chemotherapy course.
Chemotherapy cycle. Hospitalizations for any reason (i.e. all-cause) were also identified.

Ascertaining outpatient care for CINC was based on an ambulatory encounter with a diagnosis of neutropenia, fever or infection and an HCPCS Level I (i.e. CPT) code for IV administration of antimicrobial therapy. Such encounters that preceded a hospitalization during the same cycle of chemotherapy were not considered as outpatient episodes. Outpatient care episodes were identified between day 6 and the last day of the chemotherapy cycle.

**Burden**

Characterizing the clinical and economic burdens of CINC was based on the care setting. For such complications requiring inpatient care, the burden was characterized in terms of total days in hospital, total inpatient costs, total outpatient costs incurred after the hospitalization but within same cycle and total overall costs (i.e. total inpatient plus total outpatient costs).

For complications treated on an outpatient basis, the burden was characterized in terms of total outpatient costs from the date of the event to the end of the cycle. Outpatient costs comprised encounters with a diagnosis of neutropenia, fever or infection, as well as use/prescriptions for CSF agents and antimicrobial therapy. Direct healthcare costs were estimated by tallying total allowed or paid amounts on medical and pharmacy claims for encounters with a diagnosis of neutropenia, fever or infection as well as claims for selected drugs (i.e. CSF agents and antibiotic therapy) from the date of the event to the end of the cycle.

**Patient and Treatment Characteristics**

Characteristics of women with MBC were evaluated including: age, the presence of selected comorbidities (diabetes, cardiovascular, liver and renal disease), history of blood disorders (anemia, neutropenia or other), infection, hospitalization, radiation treatment and other treatment modalities, the site of metastases, the use of CSF, the chemotherapy regimen and the year in which it took place (post-MBC diagnosis).

Age was assessed as of the first day of the first cycle of chemotherapy. Chronic comorbidities and a history of blood disorders, infections, hospitalization (all-cause) and radiation treatment were assessed from the beginning of the 1-year pretreatment period to the first day of the first cycle of chemotherapy. Chronic comorbidities were identified on the basis of ≥1 diagnosis codes on inpatient claims, ≥2 diagnosis codes on outpatient claims (excluding those for laboratory services) on different days, ≥1 procedure codes, and ≥1 drug codes, as appropriate (online supplement). Blood disorders and infections were identified on the basis of ≥1 diagnosis codes (in inpatient and/or outpatient claims) and ≥1 drug codes, as appropriate.

Other treatment modalities included the receipt of endocrine therapy [presumably among women with tumors that are hormone-receptor positive (i.e. that express estrogen and/or progesterone receptors)] and trastuzumab/lapatinib (presumably among women with tumors that express HER-2). Receipt of such therapies was ascertained from the beginning of the 1-year pretreatment period to the first day of the first cycle of chemotherapy using corresponding drug codes.

Use of CSF (i.e. filgrastim, pegfilgrastim or sargramostim) was characterized in terms of the reason, timing and duration of administration (daily agents only), on a cycle-specific basis, among all subjects in the study population. Characterizing the reason for the use of these agents was based on the timing of their administration within the course and cycles of chemotherapy, as follows: primary prophylaxis – receipt on or before day 5 of cycle one, secondary prophylaxis – first receipt (as prophylaxis) on or before day 5 of cycle two or a subsequent cycle and treatment – first receipt (in a given cycle) any time after cycle day 5 or receipt following the end of prophylaxis (defined as a gap of ≥3 days). The timing of the use of these agents – as prophylaxis and treatment – was depicted based on the day of first receipt during the course and cycle of administration. The duration of CSF use – as prophylaxis and treatment – was characterized based on temporal patterns of administration. Receipt of CSF was identified based on medical claims with relevant codes: filgrastim (J1440, J1441), pegfilgrastim (C9119, S0135, J2505) or sargramostim (J2820).

**Statistical Analyses**

Characteristics of MBC patients and their cancer and treatment were described on an overall basis and stratified by chemotherapy regimen. Characteristics of the chemotherapy regimens were described, including the number and duration of cycles. The use of CSF was described on an overall basis and among selected common chemotherapy regimens, in terms of the frequency of administration as primary prophylaxis, secondary prophylaxis and treatment, respectively, as well as the patterns and levels of use within the course and cycles among those receiving CSF. Categorical variables were reported as numbers and percentages; for continuous variables, means (standard deviations), ranges, medians and percentiles (25th and 75th) were reported.

The crude incidence of CINC requiring inpatient care and outpatient care only was descriptively analyzed over the entire chemotherapy course as well as on a cycle-specific basis. Risks were analyzed for all patients in the study population and patients receiving selected chemotherapy regimens. The burden of CINC was descriptively analyzed among patients developing this condition, and was summarized in terms of levels of the associated use of resources (e.g., hospitalizations and outpatient encounters) and expenditures during the cycle, considering all episodes collectively and by care setting. Risks and burden were estimated in a non-comparative fashion, and were summarized using means and percentages along with corresponding 95% CIs (estimated using normal and binomial distributions, as appropriate).

**Results**

**Patient Characteristics**

A total of 130,726 adult women in the study database met our criteria for a diagnosis of breast cancer (≥30 days apart). Among these women, 25,247 (19%) met our criteria for a diagnosis of MBC and 4,695 (4%) also initiated a course of myelotoxic chemotherapy (with a minimum first-cycle length of 14 days). The final study population included 2,620 women with MBC who met all remaining selection criteria (fig. 1).

Mean (± SD) age of study subjects was 55 (±10) years, and the prevalence of each of the chronic comorbidities
was less than 10% (table 1). Seventeen percent of subjects had evidence of prior receipt of chemotherapy (i.e. during the 1-year history period), and 10% had evidence of prior radiation therapy. Other treatment modalities received by study subjects – prior to the index course of chemotherapy – included endocrine therapy (35%) and trastuzumab/lapatinib (3.5%). The most common site of metastases was the bone (45%), followed by the lung (23%), liver (19%), and brain/spinal cord (7%).

**Treatment and Supportive Care**

The top five chemotherapy regimens – based on agents administered in cycle 1 – were received by approximately 70% of all patients, and included doxorubicin/cyclophosphamide (24%), docetaxel alone (20%), paclitaxel alone (12%), carboplatin/paclitaxel (6%), and carboplatin/docetaxel (5%) (table 2). From 2004 to 2008, use of doxorubicin/cyclophosphamide decreased from 33 to 27%, while use of docetaxel alone decreased from 35 to 24%; use of paclitaxel alone increased from 10 to 21% over this same period. Mean (±SD) duration of chemotherapy, on an overall basis, was 106 (± 55) days, mean number of cycles per course was 5.5 (± 2.7), and mean duration of cycles was 19 days (± 8.4). Duration of chemotherapy was largely comparable among the most common regimens.

Nearly 50% of all patients received at least one administration of a CSF agent – for any reason – during their chemotherapy course; 44% received a CSF as prophylaxis and 17% received a CSF as treatment (thus, about 11% [44 + 17 – 50%] received a CSF as prophylaxis and, at another time during their chemotherapy course, received a CSF as treatment). Pegfilgrastim was the most commonly used CSF agent as primary prophylaxis (91%) and secondary prophylaxis (75%), followed by filgrastim (6 and 26%, respectively). Filgrastim was the most commonly used CSF as treatment (68%), followed by pegfilgrastim (30%). Among patients who received CSF prophylactically, mean number of cycles with prophylaxis was 3.7 (± 2.2). Prophylaxis was first administered on day 1 of the cycle (i.e., on the same day chemotherapy was first administered in the cycle) for 18% of patients, day 2 for 62%,
## Table 2. Use of CSF in study population

| Chemotherapy course | Overall (n = 2,620) | Doxorubicin/cyclophosphamide (n = 641) | Docetaxel (n = 531) | Paclitaxel (n = 316) | Carboplatin/paclitaxel (n = 160) | Carboplatin/docetaxel (n = 133) |
|---------------------|---------------------|----------------------------------------|---------------------|---------------------|----------------------------------|----------------------------------|
| Duration, days      |                     |                                        |                     |                     |                                  |                                  |
| Mean ± SD           | 106.0 ± 55.2        | 105.5 ± 46.4                           | 110.0 ± 61.2        | 92.5 ± 59.5         | 107.7 ± 60.4                    | 114.1 ± 51.1                    |
| Min/med/max         | 14/105/364          | 14/103/280                             | 14/106/295          | 14/91/274           | 14/105/364                      | 14/112/269                      |
| Chemotherapy cycles |                     |                                        |                     |                     |                                  |                                  |
| Number per course   |                     |                                        |                     |                     |                                  |                                  |
| Mean ± SD           | 5.5 ± 2.7           | 6.0 ± 2.5                               | 5.6 ± 2.9           | 5.3 ± 3.1           | 5.4 ± 2.6                       | 5.3 ± 2.2                       |
| Min/med/max         | 1/5/9               | 1/7/9                                   | 1/6/9               | 1/5/9               | 1/5/9                           | 1/5/9                           |
| Duration, days      |                     |                                        |                     |                     |                                  |                                  |
| Mean ± SD           | 18.8 ± 8.4          | 19.8 ± 6.7                              | 19.5 ± 8.2          | 15.0 ± 7.9          | 19.0 ± 8.6                      | 21.4 ± 7.5                      |
| Min/med/max         | 7/21/60             | 7/21/60                                 | 7/21/60             | 7/14/58             | 7/21/58                         | 7/21/59                         |
| Duration, in weeks, % |                   |                                         |                     |                     |                                  |                                  |
| Weekly              | 17.9                | 1.0                                     | 16.8                | 34.7                | 22.5                            | 8.3                             |
| Biweekly            | 21.7                | 38.9                                    | 13.2                | 30.0                | 12.3                            | 8.5                             |
| Triweekly           | 44.7                | 46.9                                    | 54.6                | 26.5                | 47.1                            | 64.2                            |
| Other               | 15.7                | 13.2                                    | 15.4                | 8.8                 | 18.1                            | 19.0                            |
| Use of prophylaxis during course, % | 49.3 | 60.1                                    | 36.0                | 30.1                | 52.5                            | 65.4                            |
| CSF agent, %        | 44.3                | 55.2                                    | 28.4                | 25.9                | 47.5                            | 60.9                            |
| Pegfilgrastim       | 87.2                | 92.1                                    | 73.0                | 86.8                | 81.0                            | 78.3                            |
| Filgrastim          | 15.3                | 10.5                                    | 10.0                | 10.5                | 11.0                            | 21.7                            |
| Sargramostim        | 3.7                 | 2.8                                     | 4.0                 | 3.9                 | 2.0                             | 4.3                             |
| No. of cycles with prophylaxis, mean ± SD | 3.7 ± 2.2 | 4.0 ± 2.3                               | 3.5 ± 2.4           | 3.1 ± 2.5           | 3.3 ± 2.1                       | 4.1 ± 2.1                       |
| Cycle day of first receipt, % | 18.0 | 19.2                                    | 18.3                | 20.2                | 17.5                            | 25.9                            |
| 2                   | 62.1                | 65.9                                    | 56.7                | 63.6                | 62.7                            | 55.3                            |
| ≥3                  | 19.9                | 14.8                                    | 25.0                | 16.3                | 19.7                            | 18.8                            |
| Duration of prophylaxis, mean ± SD | 4.2 ± 4.0 | 6.2 ± 5.7                               | 3.1 ± 2.4           | 3.6 ± 4.2           | 4.1 ± 2.2                       | 5.0 ± 4.2                       |
| Use of primary prophylaxis, % | 31.1 | 43.8                                    | 16.0                | 15.8                | 26.3                            | 45.1                            |
| CSF agent, %        | 31.1                | 43.8                                    | 16.0                | 15.8                | 26.3                            | 45.1                            |
| Pegfilgrastim       | 91.4                | 92.9                                    | 85.9                | 94.0                | 97.6                            | 91.7                            |
| Filgrastim          | 6.1                 | 5.3                                     | 12.9                | 2.0                 | 2.4                             | 3.3                             |
| Sargramostim        | 3.1                 | 2.1                                     | 1.2                 | 6.0                 | 0.0                             | 5.0                             |
| Cycle day of first receipt, % | 16.7 | 18.9                                    | 20.2                | 14.6                | 19.0                            | 17.5                            |
| 2                   | 65.7                | 68.4                                    | 65.5                | 68.8                | 69.0                            | 54.4                            |
| ≥3                  | 17.7                | 12.7                                    | 14.3                | 16.7                | 11.9                            | 28.1                            |
| Duration of prophylaxis, mean ± SD | 4.5 ± 4.1 | 5.3 ± 5.1                               | 2.3 ± 2.6           | 5.0 ± 3.3           | 7.0 ± 0.0                       | 8.2 ± 5.8                       |
| Use of secondary prophylaxis, % | 13.2 | 11.4                                    | 12.4                | 10.1                | 21.2                            | 15.8                            |
| CSF agent, %        | 13.2                | 11.4                                    | 12.4                | 10.1                | 21.2                            | 15.8                            |
| Pegfilgrastim       | 75.4                | 82.2                                    | 75.8                | 81.3                | 73.5                            | 81.0                            |
| Filgrastim          | 26.4                | 19.2                                    | 22.7                | 28.1                | 17.6                            | 23.8                            |
| Sargramostim        | 4.1                 | 5.5                                     | 3.0                 | 0.0                 | 8.8                             | 0.0                             |
| No. of cycles with prophylaxis, mean ± SD | 3.4 ± 1.9 | 3.7 ± 1.9                               | 3.4 ± 2.2           | 3.3 ± 2.3           | 3.1 ± 1.8                       | 3.7 ± 1.7                       |
| Cycle day of first receipt, % | 18.3 | 19.4                                    | 18.0                | 20.8                | 17.2                            | 27.6                            |
| 2                   | 61.3                | 65.0                                    | 55.4                | 63.0                | 61.3                            | 55.5                            |
| ≥3                  | 20.4                | 15.6                                    | 26.6                | 16.2                | 21.5                            | 17.0                            |
| Duration of prophylaxis, mean ± SD | 4.1 ± 4.0 | 6.6 ± 6.0                               | 3.2 ± 2.4           | 3.5 ± 4.3           | 4.0 ± 2.1                       | 4.3 ± 3.6                       |

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Mean duration of prophylaxis with the daily agents (i.e. filgrastim and sargramostim) was 4.2 (± 4.0) days. Thirty-one percent of all subjects (70% of subjects receiving prophylaxis at anytime) received CSF prophylaxis in their first chemotherapy cycle, and an additional 13% of all subjects (30% of subjects receiving prophylaxis at any time) first received CSF prophylaxis after cycle one. Receipt of primary prophylaxis was highest among patients receiving chemotherapy with carboplatin/docetaxel (45%) and lowest among those receiving chemotherapy with docetaxel alone or paclitaxel alone (16%); receipt of secondary prophylaxis was largely comparable across regimens.

**Risk and Burden of Neutropenic Complications**

Over their course of chemotherapy, 286 (11.0% [95% CI 9.8–12.2] of all subjects) patients experienced CINC, the large majority (88%) of whom required inpatient care (9.7% [8.6–10.9] of all subjects) (table 3). Risk of CINC was 4.9% [4.1–5.8] in cycle 1 (45% of all subjects experiencing complications), and was 2.3% (1.8–2.9) in cycle 2 (21% of all subjects experiencing complications). In all subsequent cycles, 4.9% (4.1–5.8) of patients experienced CINC requiring inpatient care or outpatient care only. Overall, 16.6% of patients were hospitalized for any reason during their chemotherapy course.

Among the patients who developed CINC requiring inpatient care, mean (95% CI) associated expenditures were USD 12,869 (USD 12,622–13,116), comprising USD 12,554 (USD 12,307–12,801) for hospital-related care and USD 315 (USD 247–383) for subsequent outpatient care during the same cycle (table 4). Mean number of days in hospital was 6.6 (1.8–11.3). Among the patients who developed CINC requiring outpatient care only, mean associated expenditures were USD 2,030 (USD 1,925–2,135), comprising USD 1,249 (USD 1,146–1,351) for outpatient visits for reasons other than the receipt of CSF or antimicrobial therapy, USD 443 (USD 372–513) for receipt of CSF and USD 338 (USD 295–381) for receipt of antimicrobial therapy. The mean number of outpatient encounters for CINC was 2.9 (95% CI 0.0–5.8).

**Discussion**

Using data from 2 large U.S. private healthcare claims databases, we examined the risk and burden of CINC among women with MBC. Not surprisingly, our study population comprised women who had additional risk factors for complications, including a history of chemotherapy or radiation therapy, chronic comorbidities and/or bone marrow involvement. We found that over 10% of these women experienced CINC during their chemotherapy course, most commonly requiring inpatient care and during the first cycle. We also found that the economic consequences were substantial for the patients experiencing neutropenic events, with mean associated costs ap-
approaching USD 13,000 for those requiring inpatient care and totaling over USD 2,000 for those requiring outpatient care only.

Our findings on the risk of CINC among women with MBC are largely comparable to published data from clinical trials, despite the differences in study designs and case-ascertainment criteria. For example, among women in our study population who received chemotherapy with doxorubicin/cyclophosphamide – the most common regimen – the number requiring inpatient care for CINC (9.7%) was similar to that (9 and 10%, respectively) in two randomized controlled trials of the same combination of chemotherapy agents as first-line treatment in MBC [27, 28]. Our findings also are largely comparable to those from a recent retrospective cohort study focusing on patients with advanced non-small cell lung cancer. In the Stokes et al. study [21], 10% of the subjects were classified as having developed febrile neutropenia. Of these, 72% required hospitalization; corresponding estimates from our study were 11 and 88%. The percentage of MBC patients developing CINC who required inpatient care in our study also is consistent with data from another published study of CSF use in cancer patients receiving chemotherapy [29, 30]. In contrast with rates for lung cancer and breast cancer, a population-based examination of the hospitalization rate for chemotherapy-associated neutropenia among older adults with non-Hodgkin's lymphoma in the US reported that 22% of subjects were hospitalized [31].

Table 3. Risk of neutropenic complications

| Risk of neutropenic complications during course, % (95% CI) | Overall (n = 2,620) | Doxorubicin/cyclophosphamide (n = 641) | Docetaxel (n = 531) | Paclitaxel (n = 316) | Carboplatin/paclitaxel (n = 160) | Carboplatin/docetaxel (n = 133) |
|---|---|---|---|---|---|---|
| All events | 11.0 (9.8–12.2) | 9.7 (7.5–12.2) | 11.5 (8.9–14.5) | 7.6 (4.9–11.1) | 5.6 (2.6–10.4) | 9.8 (5.3–16.1) |
| Requiring inpatient care | 9.7 (8.6–10.9) | 8.6 (6.5–11.0) | 9.8 (7.4–12.6) | 7.3 (4.7–10.7) | 5.0 (2.2–9.6) | 7.5 (3.7–13.4) |
| Requiring outpatient care only | 1.6 (1.2–2.2) | 1.2 (0.5–2.4) | 1.7 (0.8–3.2) | 0.6 (0.1–2.3) | 0.6 (0.0–3.4) | 2.3 (0.5–6.5) |

Table 4. Burden of neutropenic complications

| All neutropenic complications (n = 342) | Value (95% CI) |
|---|---|
| Expenditures, USD | 11,094 (10,853–11,335) |
| Events requiring inpatient care (n = 286) | 6.6 (1.8–11.3) |
| Hospital days | 12,554 (12,307–12,801) |
| Expenditures, USD | 315 (247–383) |
| Inpatient | 443 (372–513) |
| Outpatient | 338 (295–381) |
| Total | 12,869 (12,622–13,116) |
| Events requiring outpatient care only (n = 56) | 2.9 (0.0–5.8) |
| Number of encounters | 1,249 (1,146–1,351) |
| Expenditures, USD | 443 (372–513) |
| Encounters | 338 (295–381) |
| CSF | 1,249 (1,146–1,351) |
| Antibiotics | 443 (372–513) |
| Total | 2,030 (1,925–2,135) |

For patients with multiple events, each one was considered separately in these analyses.

Interestingly, while nearly 50% of patients received CSF agents as prophylaxis or treatment during their chemotherapy course – findings comparable to those reported by Stokes et al. [21, 26] – it appears that many of the patients who received prophylaxis in our study were ad-

For patients with multiple events, only the first one during the corresponding period was considered in these analyses.
administered these agents in a manner that is inconsistent with recommendations. Nearly one fifth of all patients receiving prophylaxis (primary or secondary) first received CSF on day one of the cycle, while recommendations specify that prophylaxis should be initiated 24–72 h after completion of chemotherapy and that ‘administration of growth factor on same day as chemotherapy is not recommended’ [26]. Moreover, among patients receiving one of the CSF agents requiring daily administration (filgrastim or sargramostim), the mean duration of prophylaxis was only about 4 days, substantially lower than the 10–11 days typically required for post-nadir absolute neutrophil count (ANC) recovery in two large recent clinical trials [32, 33]. We note, however, that such patients represent only a minority of patients who received prophylaxis as the large majority received prophylaxis with pegfilgrastim, which requires only a single dose per cycle.

Our estimates of the cost of CINC were similar to those previously reported for other patient populations [10, 21, 22, 34]. In two of these studies, which utilized hospital discharge records, the mean cost of hospitalization among patients with solid tumors ranged from USD 8,100 to USD 13,354, and for the subgroup of patients with breast cancer (all stages) mean cost ranged from USD 7,100 to USD 12,372 [10, 22]. In the aforementioned study focusing on patients with advanced non-small cell lung cancer, mean total cost of neutropenia-related care for those treated on an inpatient or outpatient basis was estimated to be USD 12,148, including USD 7,598 for inpatient care [21]. We note that our estimates of the economic burden of CINC may not reflect the full impact of this condition, since downstream-associated (i.e. post-infection) costs were not captured, and such costs have been found to represent a substantial proportion of the total economic burden of this condition [35].

Potential biases with regard to patient (treatment) selection and outcome measurement may impact this study. Most important, the accuracy of the algorithms for identifying patients receiving treatment for MBC and for identifying CINC is unknown, as they have not been validated. Because an ICD-9-CM diagnosis code for CINC (i.e. neutropenia-related fever or infection) does not exist, an algorithm based on codes for neutropenia, fever and infection (and i.v. antibiotic therapy for outpatient care) was employed to identify encounters for the events of interest. While this algorithm has not been formally validated, it (and similar ones) has been utilized in a number of studies published to date [35–37]. Although our algorithm for ascertaining outpatient treatment of CINC required the administration of i.v. antibiotic therapy, we acknowledged that some low-risk FN patients may receive outpatient management with oral (rather than i.v.) antimicrobial therapy in clinical practice. We suspect, however, that the large majority of patients who receive oral antimicrobial drugs on an outpatient basis have either a low ANC (i.e. neutropenia) and are receiving the drugs as prophylaxis against FN, or a normal ANC and are receiving the drugs for a coded infection. To the extent that we may have missed such patients, we may have underestimated the frequency of outpatient FN and may have – to the extent that these patients differ systematically in their economic profile from outpatient FN cases treated with IV agents – mis-estimated economic cost. Finally, while some infection-related encounters during a given cycle may not be related to the neutropenic event (especially those that occur temporally distal, e.g. 2 weeks), it is likely that most of these encounters are a consequence of treatment and the associated neutropenia.

Several generic limitations of retrospective studies based on healthcare claims data also should be noted. All such databases contain errors of omission and commission in coding. In addition, certain biases in coding may exist, such that patients who, for example, are hospitalized for CINC may be more likely to have ‘neutropenia’ (ICD-9-CM 288.0) designated as a secondary (or even primary) diagnosis on future encounters as opposed to patients without a history of these complications, all else being equal. Moreover, information often is not available for one or more clinically important parameters (e.g. in our study, ANC and performance status), and pertinent medical history can be left-censored (e.g. receipt of chemotherapy and neutropenic complications occurring before the time period of the study database are unobservable). Thus, as noted above, although most index courses of chemotherapy probably represent the first course of treatment subsequent to the diagnosis of MBC, we cannot be certain that this was the case for all patients in the study population. The impact of these limitations on our results cannot be assessed within the scope of this study.

In addition to these generic limitations of healthcare claims data, several limitations specific to our study should be noted. First, for some patients, an HCPCS Level I code for the administration of chemotherapy would have been present without any HCPCS Level II codes for specific chemotherapy agents. Not knowing whether or not the agent-specific codes had been omitted erroneously, the HCPCS Level I code had been incorrectly entered or chemotherapy had been obtained from another source (e.g. investigational drug study), we excluded such pa-
patients (i.e. those with missing drug information) from analyses. Second, chemotherapy regimens may be misclassified for patients who received combination therapy with a taxane or anthracycline agent and a new chemotherapy agent during the period in which a unique code is not available. However, only a handful of new myelosuppressive chemotherapy agents were approved for use in the USA during the period of interest, and all such agents were either infrequently used in clinical practice or not indicated for use as combination therapy with a taxane or anthracycline agent in the treatment of MBC. Third, patients obtaining chemotherapy or CSF agents via the retail pharmacy were excluded from analyses since the precise dates of administration cannot be determined in healthcare claims databases. Fourth, our study sample was comprised principally of cancer patients with private healthcare coverage who were less than 65 years old. Our study population may thus not have been representative of the entire population of women with MBC in the USA, which would limit the generalized application of its findings.

In summary, the results of this study suggest that the incidence of CINC among women with MBC is high and that the economic consequences of this condition are considerable. Additional research is needed to identify the subgroups of women with MBC who are at a particularly high risk of CINC, so that they may be targeted for preventive measures.

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