A prospective cohort study to evaluate the incidence of febrile neutropenia in patients receiving pegfilgrastim on-body injector versus other options for prophylaxis of febrile neutropenia: breast cancer subgroup analysis

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Abstract

Background Breast cancer chemotherapy often carries a high risk of febrile neutropenia (FN); guidelines recommend prophylaxis with granulocyte colony-stimulating factor (G-CSF), such as pegfilgrastim. Neulasta® Onpro® on-body injector (OBI) is a delivery device administering pegfilgrastim approximately 27 h after application.

Methods This prospective study examined patients with breast cancer who received chemotherapy with a high risk of FN, receiving OBI (“OBI”) or other options (other G-CSF or none; “other”). The primary endpoint was FN incidence; secondary endpoints included chemotherapy delivery, adherence (G-CSF in all cycles), compliance (G-CSF day after chemotherapy), and FN incidence in patients receiving curative or palliative treatment.

Results A total of 1776 patients with breast cancer were enrolled (OBI, n = 1196; other, n = 580). Across all cycles, FN incidence was lower for OBI (4.4% [95% CI, 3.3–5.6%]) than other (7.4% [5.3–9.6%]). For curative treatment, the FN incidence across all cycles was lower for OBI (4.6% [3.4–5.8%]) than for other (7.1% [5.0–9.3%]). For palliative treatment (OBI, n = 33; other, n = 20), 3 patients (15%) in the other and none in the OBI group had FN. After adjusting for baseline covariates, FN incidence remained lower for OBI (4.6% [3.5–6.1%]) versus other (7.8% [5.7–10.5%]). Adherence was higher for OBI (93.8%) than for other G-CSF (69.8%), as was compliance (90.5 and 53.2%, respectively). Chemotherapy dose delays/reductions were similar for OBI (4.7%/32.3%, respectively) and other (4.7%/30.0%) groups.

Conclusion Pegfilgrastim OBI was associated with a lower FN incidence in patients with breast cancer compared to other options for FN prophylaxis.

Trial registration www.clinicaltrials.gov, NCT02178475, registered 30 June, 2014

Keywords Pegfilgrastim · Febrile neutropenia · Breast cancer · Compliance · Chemotherapy
Introduction

Febrile neutropenia (FN) is a potentially life-threatening complication that can occur in patients with breast cancer who are receiving myelosuppressive chemotherapy and is associated with significant morbidity, mortality, healthcare resource utilization, and costs [1, 2]. In a recent analysis, the incidence of FN in the USA was as high as 20.6% in patients with breast cancer receiving chemotherapy without prophylaxis for FN [3]. According to real-world data from the USA, the rate of hospitalization for FN in patients with breast cancer was 13.9%, and the mortality rate in hospitalized patients with FN was estimated to be 2.0–2.6%; the mean length of stay ranged from 4.1 to 5.7 days and mean hospital costs ranged from $16,940 to $37,087 USD [1, 2]. Among patients with metastatic breast cancer, resource utilization and mortality rate were even higher: FN-associated hospitalization rate was 89%, and the mortality rate in hospitalized patients was 7.3% [4]. Additionally, FN may necessitate chemotherapy dose delays or reductions, which are significantly associated with increased mortality [5, 6]. Among patients with breast cancer receiving standard myelosuppressive chemotherapy, up to 60% and up to 96% experienced dose delays and dose reductions, respectively [1, 7, 8].

The risk of FN varies based on chemotherapy regimen and comorbidities, and National Comprehensive Cancer Network (NCCN) guidelines recommend prophylaxis with granulocyte colony-stimulating factors (G-CSFs) in patients receiving chemotherapy regimens with a high risk (>20%) of FN and intermediate-risk (10–20%) of FN with ≥1 risk factor for FN [9, 10]. The NCCN recommends following the FDA-approved dosing schedule, which is G-CSF prophylaxis administered the day following chemotherapy [9, 10]. Pegfilgrastim is a long-acting G-CSF that has been shown to reduce the risk of FN by 94% with first-cycle use in patients with breast cancer receiving docetaxel 100 mg/m² every 3 weeks [9, 11]. The Neulasta® Onpro® on-body injector (OBI) is a delivery device that administers pegfilgrastim 6 mg approximately 27 h after application in accordance with NCCN guidelines and FDA recommendations, thus eliminating the need for an office visit the day after chemotherapy administration [12]. In a retrospective study of twenty-eight patients who received chemotherapy and pegfilgrastim OBI between 2016 and 2018, there were no hospitalizations due to FN or chemotherapy dose delays or reductions; however, there was no comparator arm in this study, necessitating additional research [13].

Chemotherapy that carries a high risk of febrile neutropenia is commonly used in breast cancer [14]; however, there is limited evidence of clinical benefits of adherence and compliance to OBI in this patient population. Here, we report analyses of the subgroup of patients with breast cancer with the objective to estimate the incidence of FN in patients with breast cancer who received myelosuppressive chemotherapy and G-CSF prophylaxis with pegfilgrastim OBI or other options; chemotherapy delivery, adherence, and compliance were also evaluated.

Methods

Study design and population

The primary analysis of this multicenter, prospective, observational cohort study in patients receiving myelosuppressive chemotherapy and at high risk for developing FN between November 7, 2018, and April 9, 2020, was previously reported [15]. Briefly, in the primary analysis adults were eligible if they had breast, lung, or prostate cancer or NHL, a life expectancy of >6 months, ≥4 planned chemotherapy cycles administered every 3 or 4 weeks, and received a chemotherapy regimen with high FN risk (>20%) or intermediate FN risk (10–20%) and ≥1 risk factor for FN as defined by NCCN guidelines at that time [10]. In this analysis, only patients with breast cancer were included. Patients who received radiation <2 weeks before enrollment or had planned chemotherapy dose reduction for cycle 1, concurrent primary cancers (except non-melanoma skin cancer or adequately treated carcinoma in situ), or significant laboratory abnormalities per the investigator were excluded. Demographic and clinical characteristics, including tumor type, chemotherapy regimen, type of G-CSF prophylaxis received and timing, age, sex, laboratory measurements, comorbidities, and history of other malignancies, were collected. Patients were followed from study enrollment until death, discontinuation of chemotherapy, withdrawal of consent, lose to follow-up, or end of the study.

Endpoints

The primary endpoint was the incidence of FN during the study period. FN was defined as an absolute neutrophil count (ANC) <1000 × 10⁹/L and occurrence of 1 of the following within 24 h of decreased ANC: temperature >38 °C, use of oral antibiotics (i.e., ciprofloxacin, levofloxacin, moxifloxacin, or amoxicillin-clavulanate), or any intravenous antibiotics. Secondary endpoints included the incidence of FN in patients who received treatment with curative or palliative intent, chemotherapy delivery, adherence, and compliance. Adherence was defined as G-CSF support received in all chemotherapy cycles irrespective of G-CSF administration timing. G-CSF support was defined as the administration of a long-acting G-CSF (pegfilgrastim...
OBI, pegfilgrastim, or biosimilar pegfilgrastim (pegfilgrastim-jmdb, pegfilgrastim-cbqv, pegfilgrastim-bmez) or ≥10 administrations of a short-acting G-CSF (filgrastim, biosimilar filgrastim [filgrastim-sndz or filgrastim-aafi], tbo-filgrastim, or sargramostim) in each cycle. Compliance with pegfilgrastim was defined as pegfilgrastim OBI, pegfilgrastim, or biosimilar pegfilgrastim administered the day after chemotherapy completion in all cycles in which pegfilgrastim was administered.

**Statistical analysis**

Patients with breast cancer were categorized by the type of G-CSF prophylaxis received in the first cycle of chemotherapy: pegfilgrastim OBI (OBI) or other option (other), which included pegfilgrastim pre-filled syringe (PFS), pegfilgrastim biosimilar PFS, filgrastim, tbo-filgrastim, filgrastim-sndz, or no G-CSF prophylaxis selected at the physician's discretion. Patients remained in the originally-assigned group even if a different type of G-CSF prophylaxis was administered in a subsequent cycle. The incidence of FN was calculated as the percentage of patients who experienced FN, and 95% confidence intervals (CIs) were calculated using a normal approximation method. As reported in the primary analysis, the incidence of FN was adjusted using a standardized log-binomial model to account for confounding variables [15]. These included prior surgery within 6 months before study.

### Table 1 Patient demographics and baseline characteristics

|                        | On-body injector (N = 1196) | Other physician choice (N = 580) | All patients with breast cancer (N = 1776) |
|------------------------|------------------------------|----------------------------------|--------------------------------------------|
| Sex, n (%)             |                              |                                  |                                            |
| Male                   | 3 (0.3)                      | 5 (0.9)                          | 8 (0.5)                                    |
| Female                 | 1193 (99.7)                  | 575 (99.1)                       | 1768 (99.5)                                |
| Age, years             |                              |                                  |                                            |
| Median (IQR)           | 60 (49–68)                   | 58 (48–67)                       | 59 (49–67)                                 |
| ECOG performance status, n (%) |                        |                                  |                                            |
| 0–1                    | 1174 (98.2)                  | 574 (99.0)                       | 1748 (98.4)                                |
| ≥2                     | 14 (1.2)                     | 4 (0.7)                          | 18 (1.0)                                   |
| Missing                | 8 (0.7)                      | 2 (0.3)                          | 10 (0.6)                                   |
| Number of comorbidities, n (%) |                     |                                  |                                            |
| ≥2                     | 173 (14.5)                   | 82 (14.1)                        | 255 (14.4)                                 |
| ≤2                     | 1023 (85.5)                  | 498 (85.9)                       | 1521 (85.6)                                |
| History of any other malignancy, n (%) |                     |                                  |                                            |
| Yes                    | 51 (4.3)                     | 34 (5.9)                         | 85 (4.8)                                   |
| No                     | 1145 (95.7)                  | 546 (94.1)                       | 1691 (95.2)                                |
| Antibiotic use 0–7 days prior to initiation of chemotherapy, n (%) |                     |                                  |                                            |
| Yes                    | 35 (2.9)                     | 29 (5.0)                         | 64 (3.6)                                   |
| No                     | 1161 (97.1)                  | 551 (95.0)                       | 1712 (96.4)                                |
| Prior surgery, n (%)   |                              |                                  |                                            |
| Yes                    | 980 (81.9)                   | 402 (69.3)                       | 1382 (77.8)                                |
| No                     | 216 (18.1)                   | 178 (30.7)                       | 394 (22.2)                                 |
| Prior chemotherapy, n (%) |                          |                                  |                                            |
| Yes                    | 2 (0.2)                      | 0 (0.0)                          | 2 (0.1)                                    |
| No                     | 1194 (99.8)                  | 580 (100.0)                      | 1774 (99.9)                                |
| Prior radiotherapy, n (%) |                        |                                  |                                            |
| Yes                    | 19 (1.6)                     | 10 (1.7)                         | 29 (1.6)                                   |
| No                     | 1177 (98.4)                  | 570 (98.3)                       | 1747 (98.4)                                |
| Intent of treatment, n (%) |                          |                                  |                                            |
| Curative               | 1424 (87.7)                  | 606 (83.8)                       | 2030 (86.5)                                |
| Palliative             | 200 (12.3)                   | 117 (16.2)                       | 317 (13.5)                                 |

*aExcluding nonmelanoma skin cancer. bWithin 6 months prior to study enrollment. ECOG, Eastern Cooperative Oncology Group; IQR, interquartile range"
enrollment, antibiotic use 0-7 days before initiation of chemotherapy, and FN risk of the chemotherapy regimen. Adjusted incidences of FN, relative risks, and associated p-values were calculated using a standardized log-binomial model in which weight was assigned to each patient equal to the inverse probability of exposure conditional on that patient’s confounder information [16, 17]. Associated 95% CIs for adjusted incidences and relative risk were calculated using bootstrap methods. Patient characteristics, chemotherapy delivery, adherence, and compliance were summarized using descriptive analyses. Missing data were not imputed.

### Results

#### Patient disposition and baseline characteristics

Of the 2347 patients enrolled between November 7, 2018, and April 9, 2020, 1776 had breast cancer (OBI, 1196; other, 580). Baseline characteristics were generally well balanced across groups (Table 1). Patient characteristics were comparable between groups for sex, age, ECOG performance status, number of comorbidities, history of any other malignancy, and prior antibiotic use, chemotherapy, or radiotherapy. A higher percentage of patients in the OBI

| Table 2 Baseline febrile neutropenia risk and chemotherapy regimens |
|---------------------------------------------------------------|
| FN risk of chemotherapy regimen, n (%)                       |
|                                                                 |
| High                                                          |
| 1069 (89.4)                                                   |
| Intermediate                                                   |
| 127 (10.6)                                                    |
| Chemotherapy regimen, n (%)                                    |
| High risk for FN (>20%)                                        |
| TCb                                                          |
| 573 (47.9)                                                    |
| TCHPc                                                        |
| 392 (32.8)                                                    |
| TCHd                                                         |
| 81 (6.8)                                                      |
| TACE                                                          |
| 23 (1.9)                                                      |
| Intermediate risk for FN (10–20%)                             |
| AC                                                           |
| 61 (5.1)                                                      |
| AC→T                                                          |
| 47 (3.9)                                                      |
| Docetaxel                                                     |
| 8 (0.7)                                                       |
| TH                                                            |
| 5 (0.4)                                                       |
| Paclitaxel                                                    |
| 3 (0.3)                                                       |
| CMF classic                                                   |
| 2 (0.2)                                                       |
| Bendamustine and rituximabf                                   |
| 1 (<0.1)                                                      |
| Carboplatin and paclitaxeleb                                  |
| 0                                                             |
| Cisplatin and etoposideb                                      |
| 0                                                             |

*Other physician choice includes long-acting G-CSF (n = 427; pegilgrastim pre-filled syringe, pegfilgrastim biosimilar pre-filled syringe), short-acting G-CSF (n = 45; filgrastim, tbo-filgrastim, filgrastim-sndz), or no G-CSF prophylaxis (n = 108); selected at the physician’s discretion. *4 Docetaxel 75 mg/m² and cyclophosphamide 600 mg/m² every 3 weeks for 4 cycles. *5 Docetaxel 75 mg/m², carboplatin (AUC 6), trastuzumab (day 1), and pertuzumab (day 1) every 3 weeks for 6 cycles; trastuzumab dosing weekly and every 3 weeks was allowed. *6 Docetaxel 75 mg/m², carboplatin (AUC 6), and trastuzumab (day 1) every 3 weeks for 6 cycles; trastuzumab dosing weekly and every 3 weeks was allowed. *7 Docetaxel (75 mg/m² day 1), doxorubicin (50 mg/m² day), and cyclophosphamide (500 mg/m² day) every 3 weeks for 6 cycles. *8 One patient received 4 cycles of bendamustine and rituximab for follicular lymphoma that was incorrectly coded as tumor type “breast” by the site and was included in the analysis. *9 One patient received carboplatin and paclitaxel, which is not a common chemotherapy regimen for breast cancer. We were unable to obtain clarification from the site regarding the use of this regimen. *10 One patient received cisplatin and etoposide for poorly differentiated neuroendocrine carcinoma considered to be small cell carcinoma of the breast. AC, doxorubicin cyclophosphamide; AC→T, doxorubicin, cyclophosphamide→docetaxel; AUC, area under the curve; CMF, cyclophosphamide, methotrexate, fluorouracil; FN, febrile neutropenia; G-CSF, granulocyte colony-stimulating factor; TAC, docetaxel, doxorubicin, cyclophosphamide; TC, docetaxel, cyclophosphamide; TCH, docetaxel, carboplatin, trastuzumab; TCHP, docetaxel, carboplatin, trastuzumab, pertuzumab; TH, docetaxel, trastuzumab.
group had prior surgery within 6 months of enrollment compared with patients in the other group (81.9 versus 69.3%, respectively). Most patients received chemotherapy with curative intent (87.7% in the OBI group; 83.8% in the other group).

The percentage of patients who received high FN risk chemotherapy regimens was comparable between OBI and other groups (89.4 and 82.2%, respectively; Table 2). The most common chemotherapy regimen with high FN risk was docetaxel and cyclophosphamide (OBI, 47.9%; other, 43.8%) followed by docetaxel, carboplatin, trastuzumab, and pertuzumab (OBI, 32.8%; other, 31.4%). The most common chemotherapy regimen with intermediate FN risk was doxorubicin and cyclophosphamide (OBI, 5.1%; other, 8.1%).

Among patients in the other group during cycle 1, 73.6 and 7.8% of patients received long- and short-acting G-CSF, respectively, and 18.6% received no G-CSF support.

### Incidence of febrile neutropenia

Across all cycles, the incidence of FN was lower in patients who received pegfilgrastim OBI (4.4% [95% CI, 3.3–5.6%]) compared with patients who received other options (7.4%)

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**Fig. 1** Incidence of FN in patients with breast cancer who received pegfilgrastim OBI or other options. **a** Incidence of FN; percent plus 95% CI. **b** Incidence of FN in patients who received pegfilgrastim OBI in every cycle; percent plus 95% CI. **c** Relative risk of FN. CI, confidence interval; FN, febrile neutropenia; OBI, on-body injector; Other, other physician choice options; RR, relative risk
The OBI group had a lower incidence of FN at each cycle. Similar trends were observed in patients who received pegfilgrastim OBI in every cycle; across all cycles, the incidence of FN in patients who received pegfilgrastim OBI in every cycle was 4.0% (95% CI, 2.8−5.2%; Fig. 1b). After adjusting for covariates (prior surgery within 6 months prior to study enrollment, antibiotic use 0−7 days prior to initiation of chemotherapy, and FN risk of chemotherapy regimen), the estimated incidence of FN was 4.6% (95% CI, 3.5−6.1%) for the OBI group compared with 7.8% (95% CI, 5.7−10.5%) for the other group. The risk of developing FN was significantly reduced for the OBI group versus the other group (RR, 0.60 [95% CI, 0.40−0.88]; \( P = 0.011 \); Fig. 1c). The risk of developing FN was further reduced in patients who received pegfilgrastim OBI in every cycle (RR, 0.55 [95% CI, 0.39−0.84]; \( P = 0.005 \)). Similarly, for patients who received treatment with curative intent, the incidence of FN across all cycles was lower in patients who received pegfilgrastim OBI (4.6% [95% CI, 3.4−5.8%]; Fig. 2a) and in patients who received pegfilgrastim OBI in every cycle (4.1% [95% CI, 2.9−5.4%]; Fig. 2b) compared with other options (7.1% [95% CI, 5.0−9.3%]). Of patients treated with palliative intent (OBI, 33; other, 20), FN was only observed during cycle 1 in 3 patients (15%) in the other group (Fig. 3). Two of these patients received docetaxel with no G-CSF in cycle 1; the other patient received docetaxel plus cyclophosphamide plus biosimilar pegfilgrastim in cycle 1.

### Chemotherapy delivery

The percentage of patients who required chemotherapy dose delays or reductions was comparable for patients who received pegfilgrastim OBI or other options. Across all cycles, the percentage of patients with chemotherapy dose delays was 4.7% (95% CI, 3.5−5.9%) for the OBI group and 4.7% (95% CI, 2.9−6.4%) for the other group; at each cycle, the percentage of patients with chemotherapy dose delays was similar between groups. Across all cycles, the percentage of patients...
with chemotherapy dose reductions was 32.3% (95% CI, 29.6–34.9%) for the OBI group and 30.0% (95% CI, 26.3–33.7%) for the other group, and percentages at each cycle were comparable between groups. Almost all chemotherapy dose delays and reductions occurred after the first cycle.

Adherence and compliance

Adherence to G-CSF (1 long-acting G-CSF or 10 short-acting G-CSF per chemotherapy cycle, regardless of timing) was higher in patients who received pegfilgrastim OBI (93.8% [95% CI, 92.5–95.2%]) compared with patients who received other options (69.8% [95% CI, 66.1–73.6%]; Fig. 4a). Of patients who received pegfilgrastim, compliance (a receipt of long-acting G-CSF the day after the last day of chemotherapy) was higher in patients in the pegfilgrastim OBI group (90.5% [95% CI, 88.8–92.1%]) compared with patients in the other group (n = 462; 53.2% CI [48.7–57.8%]; Fig. 4b).

Discussion

This sub-analysis of a prospective observational study evaluated clinical outcomes in patients with breast cancer who received myelosuppressive chemotherapy and G-CSF support as pegfilgrastim OBI or other options for prophylaxis of FN. The overall incidence of FN was lower in patients who received pegfilgrastim OBI in the first chemotherapy cycle compared with other options, regardless of the use of pegfilgrastim OBI in subsequent cycles. The risk of developing FN was reduced by 40% with pegfilgrastim OBI compared with other options. The incidence of dose delays and reductions was similar among patients who received pegfilgrastim OBI or other options. Adherence with G-CSF prophylaxis and compliance to pegfilgrastim were higher in patients who received pegfilgrastim OBI compared with patients who received other options.

Our findings are consistent with retrospective studies of patients receiving myelosuppressive chemotherapy for non-metastatic solid tumors or NHL. In a study of patients
with non-metastatic breast cancer or NHL receiving high or intermediate FN risk chemotherapy regimens, the incidence of FN in the first cycle was 4–9% in patients who received G-CSF prophylaxis [18]. In another retrospective study of patients with non-metastatic solid tumors (including breast) or NHL receiving high or intermediate FN risk chemotherapy regimens, the incidence of FN in the first cycle ranged from 2.2–3.8% in patients who received G-CSF prophylaxis [19]. The incidence of FN reported in a retrospective study of patients with metastatic breast cancer was higher than observed in the present study; the overall incidence of FN was 15.8%, and the rate of G-CSF prophylaxis was 16.7% in this population [4]. However, treatment in the metastatic setting is typically palliative; the American Society of Clinical Oncology guidelines did not find the demonstrable benefit of G-CSF prophylaxis in patients with metastatic disease, which is consistent with the small number of patients who received G-CSF prophylaxis in the palliative intent subgroup of the present study [6]. In this study, the risk of FN in patients receiving “other” treatment with palliative intent appears to be twofold higher for the patients receiving “other” treatment with curative intent; however, the small patient numbers in these groups mean these data should be interpreted with caution.

The decreased incidence of FN observed with pegfilgrastim OBI in this study may be associated with increased adherence and compliance in this group relative to patients who received other options [20]. Adherence was 34.4% higher, and compliance was 70.1% higher in patients who received pegfilgrastim OBI compared with patients who received other options.

In prospective and retrospective studies, receipt of pegfilgrastim at least 24 h after myelosuppressive chemotherapy administration resulted in improved adherence and patient outcomes [15, 21, 22]. Pegfilgrastim OBI allows for next-day administration without a clinic visit, therefore reducing the patient travel burden, reducing noncompliance because of the travel burden, and optimizing outcomes [23, 24].

Limitations

The definition of FN used in this study uses an ANC that may be higher than that used to define FN in some clinical settings [25]. However, as the definition was used across all groups, it is not expected to affect the generalizability of the results. Selection bias may exist because of patients being lost to follow-up after enrollment (OBI group, n = 16; 1.3%; other group, n = 10; 1.7%), and therefore it would not be possible to evaluate the risk of FN among these patients. Additionally, these results may be confounded by indication because patients who were perceived to be at higher risk by the investigator were more likely to receive G-CSF prophylaxis. In the analysis of adherence, patients with a better prognosis or lower ECOG may have been prescribed fewer doses of short-acting G-CSF prophylaxis and, thus, may have been inappropriately classified as non-adherent. In addition, reasons for lack of adherence were not captured to the extent required for further analysis. Reasons for lack of compliance were not collected. Patients receiving same-day G-CSF could have been considered non-compliant; however, the percentage of patients receiving same-day G-CSF in the overall study population (~1%; n = 28/2575) was small enough that any difference in the incidence of FN in this subgroup would be unlikely to bias the study results. Factors associated with possible sources of confounding by indication were included in the standardized log-binomial regression, but bias...
can result from unmeasured and unknown risk factors. Finally, the study closed earlier than planned due to the COVID-19 pandemic, and the target sample size was not reached, thus it was not possible to compare patients with curative versus palliative treatment intent.

Conclusions

Patients with breast cancer at high risk of developing FN who received pegfilgrastim OBI had a lower incidence of FN compared with patients who received other options. The decreased incidence of FN in the OBI group may be a result of increased adherence and compliance to G-CSF prophylaxis compared with other options.

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Data availability Qualified researchers may request data from Amgen clinical studies. Complete details are available at http://www.amgen.com/datasharing.

Code availability Not applicable.

Declarations

Ethics approval Approval was obtained from the Institutional Review Boards for various centers involved in the study. The procedures used in this study adhere to the tenets of the Declaration of Helsinki.

Consent to participate Written informed consent was obtained from all participants included in the study.

Consent for publication Not applicable.

Conflict of interest Reshama L. Mahtani reports consulting fees from Biotheranostics, Amgen, Agenda, Eisai, Immunomedics, Genentech, Lilly, Novartis, Merck, Pfizer, Puma, Sanofi, Seagen, AstraZeneca, Daiichi, and participation on an advisory board for AstraZeneca, Amgen, Agenda, Biotheranostics, Eisai, Immunomedics, Lilly, Novartis, Merck, Pfizer, Puma, Sanofi, Seagen, and Daiichi; Rajesh Belani reports employment with Poseida Therapeutics and stock/stock options in Amgen; Jeffrey Crawford reports consulting fees from GlaxoSmithKline, G1 Therapeutics, Merck, Pfizer, Spectrum, and participation on a data safety monitoring board or advisory board for Beyond Spring, G1 Therapeutics, Merrimack, Mylan and Roche; David Dale reports consulting fees from Amgen, and contracted research support from Amgen; Lucy DeCosta reports employment with Amgen Inc.; Prasad L. Gawade reports employment and stockholding with Amgen Inc.; Chanh Huynh has no disclosures to report; William W. MacLaughlin and Mohit Narang have no disclosures to report; Robert Rifkin reports participation in advisory boards for Amgen, BMS (Celgene), Coherus, Fresenius-Kabi, GSK, Janssen, and Pfizer.

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References

1. Kawatkar AA, Farias AJ, Chao C, Chen W, Barron R, Vogl FD, Chandler DB (2017) Hospitalizations, outcomes, and management costs of febrile neutropenia in patients from a managed care population. Support Care Cancer 25:2787–2795. https://doi.org/10.1007/s00520-017-3692-x
2. Pathak R, Giri S, Aryal MR, Karmacharya P, Bhatt VR, Martin MG (2015) Mortality, length of stay, and health care costs of febrile neutropenia-related hospitalizations among patients with breast cancer in the United States. Support Care Cancer 23:615–617. https://doi.org/10.1007/s00520-014-2553-0
3. Averin A, Silvia A, Lamerato L, Richert-Boe K, Kaur M, Sundaresan D, Shah N, Hatfield M, Lawrence T, Lyman GH, Weyercker D (2021) Risk of chemotherapy-induced febrile neutropenia in patients with metastatic cancer not receiving granulocyte colony-stimulating factor prophylaxis in US clinical practice. Support Care Cancer 29:2179–2186. https://doi.org/10.1007/s00520-020-05715-3
4. Weyercker D, Li X, Edelsberg J, Barron R, Kartashov A, Xu H, Lyman GH (2015) Risk and consequences of chemotherapy-induced febrile neutropenia in patients with metastatic solid tumors. J Oncol Pract 11:47–54. https://doi.org/10.1200/JOP.2014.001492
5. Lyman GH, Dale DC, Culakova E, Poniewierski MS, Wolff DA, Kuderer NM, Huang M, Crawford J (2013) The impact of the granulocyte colony-stimulating factor on chemotherapy dose intensity and cancer survival: a systematic review and meta-analysis of randomized controlled trials. Ann Oncol 24:2475–2484. https://doi.org/10.1093/annonc/mdt226
6. Smith TJ, Bohlke K, Lyman GH, Carson KR, Crawford J, Cross SJ, Goldberg JM, Khatcheressian JL, Leigh NB, Perkins CL, Somlo G, Wade JL, Wozniak AJ, Armitage JO (2015) Recommendations for the use of WBC growth factors: American Society of Clinical Oncology clinical practice guideline update. J Clin Oncol 33:3199–3212. https://doi.org/10.1200/JCO.2015.62.3488
7. Denduluri N, Lyman GH, Wang Y, Morrow PK, Barron R, Patt D, Bhomwik D, Li X, Bhor M, Fox P, Dhanda R, Saravanan S, Jiao X, Garcia J, Crawford J (2018) Chemotherapy dose intensity and overall survival among patients with advanced breast or ovarian cancer. Clin Breast Cancer 18:380–386. https://doi.org/10.1016/j.clbc.2018.02.003
8. Denduluri N, Patt DA, Wang Y, Bhor M, Li X, Favret AM, Morrow PK, Barron RL, Asmar L, Saravanan S, Li Y, Garcia J, Lyman GH (2015) Dose delays, dose reductions, and relative dose intensity in patients with cancer who received adjuvant or neoadjuvant chemotherapy in community oncology practices. J Natl Compr Canc Netw 13:1383–1393. https://doi.org/10.6004/jnccn.2015.0166
9. Becker PS, Griffiths EA, Alwan LM, Bachiashvili K, Brown A, Cool R, Curtin P, Dinner S, Gojo I, Hicks A, Kallam A, Kidwai WZ, Kloth DD, Kraut EH, Landsburg D, Lyman GH, Miller R, Mukherjee S, Patel S et al (2020) NCCN Guidelines insights: hematopoietic growth factors, Version 1.2020. J Natl Compr Canc Netw 18:12–22. https://doi.org/10.6004/jnccn.2020.0002

10. National Comprehensive Cancer Network (NCCN) (2021) NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines): hematopoietic growth factors (Version 2.2021). Available at: https://www.nccn.org/professionals/physician_gls/default.aspx. Accessed 26 Mar 2021

11. Vogel CL, Wojtkiewicz MZ, Carroll RR, Tjulandin SA, Barajas-Figueroa LJ, Wiens BL, Neumann TA, Schwartzberg LS (2005) First and subsequent cycle use of pegfilgrastim prevents febrile neutropenia in patients with breast cancer: a multicenter, double-blind, placebo-controlled phase III study. J Clin Oncol 23:1178–1184. https://doi.org/10.1200/JCO.2005.09.102

12. Neulasta® (pegfilgrastim) (2021) Full prescribing information. Amgen Inc. Thousand Oaks, CA

13. Patel J, Rainess RA, Benfield MJ, Rogers KML, Moore DC, Larc C, Arnall JR (2021) Retrospective analysis of clinical outcomes associated with the use of pegfilgrastim on-body injector in patients receiving chemotherapy requiring granulocyte colony-stimulating factor support. Hosp Pharm 56:77–80. https://doi.org/10.1177/0018578719867659

14. Collins JM, Fleming GF, Christ TN (2019) Comparison of the incidence of febrile neutropenia in obese and normal weight breast cancer patients receiving myelosuppressive chemotherapy and prophylactic pegfilgrastim. J Oncol Pharm Pract 25:1112–1118. https://doi.org/10.1177/1078155218776471

15. Mahtani RL, Crawford J, Rifkin R, Dale D, Brookhard A, Gawade PL, Lweis S, Lawrence T, Belani R, Lyman GH (2020) A multicenter, prospective, observational study to determine the incidence of febrile neutropenia (FN), persistence and G-CSF utilization among cancer patients at high risk for FN receiving pegfilgrastim by an on-body injector (OBI) versus other FN prophylaxis strategies. Presented at: San Antonio Breast Cancer Symposium; December 8–11, 2020; Virtual

16. Richardson DB, Kinlaw AC, MacLehose RF, Cole SR (2015) Standardized binomial models for risk or prevalence ratios and differences. Int J Epidemiol 44:1660–1672. https://doi.org/10.1093/ije/dyv137

17. Robins JM, Hernan MA, Brumback B (2000) Marginal structural models and causal inference in epidemiology. Epidemiology 11:550–560. https://doi.org/10.1097/00001648-200009000-00011

18. Weycker D, Doroff R, Hanau A, Bowers C, Belani R, Chandler D, Lonshteyn A, Bensink M, Lyman GH (2019) Use and effectiveness of pegfilgrastim prophylaxis in US clinical practice: a retrospective observational study. BMC Cancer 19:792. https://doi.org/10.1186/s12885-019-6010-9

19. Weycker D, Bensink M, Lonshteyn A, Doroff R, Chandler D (2019) Use of colony-stimulating factor primary prophylaxis and incidence of febrile neutropenia from 2010 to 2016: a longitudinal assessment. Curr Med Res Opin 35:1073–1080. https://doi.org/10.1080/03007995.2018.1358851

20. Shah N, Hatfield M, Lawrence T, Manjelievskaia J, Moynihan M, Bonafe M (2020) Incidence of febrile neutropenia in chemotherapy cycles with pegfilgrastim receipt via on-body injector versus pre-filled syringe. J Manag Care Spec Pharm 26:S15–S16

21. Lyman GH, Allcott K, Garcia J, Stryker S, Li Y, Reiner MT, Weycker D (2017) The effectiveness and safety of same-day versus next-day administration of long-acting granulocyte colony-stimulating factors for the prophylaxis of chemotherapy-induced neutropenia: a systematic review. Support Care Cancer 25:2619–2629. https://doi.org/10.1007/s00520-017-3703-y

22. Gawade PL, Li S, Henry D, Smith N, Belani R, Kelsh MA, Bradbury BD (2020) Patterns of granulocyte colony-stimulating factor prophylaxis in patients with cancer receiving myelosuppressive chemotherapy. Support Care Cancer 28:4413–4424. https://doi.org/10.1007/s00520-020-05295-2

23. Hawkins A, Murphy A, McNamara M, Gawade PL, Belani R, Kelsh MA (2020) A survey of oncologists’ perceptions and opinions regarding the use of granulocyte colony-stimulating factors. J Cancer Educ 35:178–186. https://doi.org/10.1007/s1097-019-01638-8

24. Stephens JM, Bensink M, Bowers C, Hollenbeak CS (2018) Travel burden associated with granulocyte colony-stimulating factor administration in a Medicare aged population: a geospatial analysis. Curr Med Res Opin 34:1351–1360. https://doi.org/10.1080/03007995.2017.1358158

25. Villafuerte-Gutierrez P, Villalon L, Losa JE, Henriquez-Camacho C (2014) Treatment of febrile neutropenia and prophylaxis in hematologic malignancies: a critical review and update. Adv Hematol 2014:986938. https://doi.org/10.1155/2014/986938

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