Systematic Review and Meta-analysis of Short-versus Long-Acting Granulocyte Colony-Stimulating Factors for Reduction of Chemotherapy-Induced Febrile Neutropenia

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ABSTRACT

Introduction: Short- and long-acting granulocyte-colony stimulating factors (G-CSFs) are approved for the reduction of febrile neutropenia. A systematic literature review was performed to identify randomized controlled trials (RCTs) and non-RCTs reporting the use of G-CSFs following chemotherapy treatment.

Methods: Medline®/Medline in-process, Embase®, and the Cochrane Library were searched for studies published between January 2003 and June 2016. A hand-search of relevant conference proceedings was conducted for meetings held between 2012 and 2016. Eligible studies were restricted to those reporting a direct, head-to-head comparison of short- versus long-acting G-CSFs for reduction of chemotherapy-induced febrile neutropenia. Risk-of-bias assessments were performed for full publications only.

Results: The search strategy yielded 4044 articles for electronic screening. Thirty-six publications were evaluated for the meta-analysis: 11 of 12 RCTs and 2 of 24 non-RCTs administered doses of the short-acting G-CSF filgrastim for ≥ 7 days. In RCT studies, there was no statistically significant difference in outcomes of interest between short- and long-acting G-CSFs. In non-RCTs, the overall risk was lower with...

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long-acting G-CSF than with short-acting G-CSF for incidence of febrile neutropenia [overall relative risk (RR) = 0.67, *P* = 0.023], hospitalizations (overall RR = 0.68, *P* < 0.05), and chemotherapy dose delays (overall RR = 0.68, *P* = 0.020).

**Conclusions:** Overall, the weight of evidence from RCTs indicates little difference in efficacy between the short- and long-acting G-CSFs if dosed according to recommended guidelines. There is some evidence for greater efficacy for long-acting G-CSFs in non-RCTs, which may be a result of under-dosing of short-acting G-CSFs in general practice in real-world usage.

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**INTRODUCTION**

Chemotherapy-induced neutropenia is one of the most frequent adverse events reported during cytotoxic chemotherapy treatment [1]. Neutropenia increases the risk of infection, often leading to febrile neutropenia (FN) [1]. In the United States, the annual incidence of FN was estimated at 91,560 cases in 2012 [2]. In the United Kingdom, the estimated annual incidence of FN in 2012 was 19.4 per 1000 oncology admissions [3]. Development of FN during chemotherapy may necessitate an interruption or cessation of treatment with chemotherapy or result in a dose reduction in subsequent treatment cycles, which can jeopardize treatment response and, ultimately, patient survival [1]. Furthermore, FN may lead to life-threatening adverse events such as neutropenic sepsis, a significant cause of mortality in patients with cancer and neutropenia [4]. The potential economic impact of FN is also significant. In the United States, mean costs of cancer-related neutropenia or FN hospitalizations were estimated at $18,880 per first FN-related hospitalization across 2007–2010 and $24,770 per stay in 2012 [2, 4].

Prophylaxis using granulocyte colony-stimulating factors (G-CSFs) decreases the risk of FN, the severity and duration of neutropenia, and the number of chemotherapy dose reductions or delays [5]. Furthermore, G-CSF prophylaxis supports delivery of optimal chemotherapy and improves survival outcomes [6, 7]. Filgrastim was the first G-CSF to be approved for the reduction of FN, and the effect of this short-acting G-CSF is dependent on the duration of treatment [8]. A 2014 study by Weycker et al. demonstrated that the risk for chemotherapy-induced neutropenic complications was substantially higher with 1–3 days [odds ratio 2.4 (95% CI 1.6–3.4)] and 4–6 days [odds ratio 1.9 (95% CI 1.3–2.8)] versus ≥ 7 days of treatment with filgrastim prophylaxis [8]. Product labeling recommends filgrastim administration once daily for up to 2 weeks or until the absolute neutrophil count (ANC) has reached 10,000/mm³ following its chemotherapy-induced nadir [9]. However, most patients treated in clinical practice receive daily G-CSF prophylaxis for ≤ 7 days and such treatment is associated with worse neutropenia-related clinical outcomes [10, 11]. A pegylated form of filgrastim (pegfilgrastim) is also available; it is administered once per 2- to 3-week chemotherapy cycle because of its longer half-life [12].

Observational studies evaluating neutropenia-related outcomes in patients receiving myelosuppressive chemotherapy suggest that pegfilgrastim is more effective than filgrastim [10, 13–23]. Literature reviews that evaluated the efficacy of short- and long-acting G-CSFs have mainly focused on clinical trial data [24, 25]. However, a recent review of real-world comparative effectiveness studies found that risks of FN and FN-related complications were generally lower for prophylaxis with pegfilgrastim than with short-acting G-CSFs [26]. Therefore, a systematic literature review (SLR) was performed to identify both randomized controlled trial (RCT) and non-RCT studies that reported the use of G-CSF for the reduction of FN following myelosuppression due to chemotherapy.
METHODS

Systematic Literature Review (SLR)

Medline®, Medline in-process, Embase®, and the Cochrane Library were searched using the OVIDSP interface to identify studies published between January 1, 2003, and August 11, 2015. Search strings consisted of text words and Medical Subject Headings terms for G-CSFs, drug names, neutropenia, and study design (Supplementary Appendix Tables S1 and S2). The search strategy was executed on August 11, 2015, and refreshed on June 15, 2016, to capture recent full-text publications. Searches were designed to overlap by 3 months with the original search date of August 2015 to allow for indexing lag within the databases.

To capture the latest studies not yet published as full-text articles and/or supplemental results of previously published studies, a hand-search of relevant conference proceedings held between 2012 and 2015 (or the most recent 3 years available) was conducted on August 24, 2015 (Supplementary Appendix Table S3), and refreshed on June 23, 2016, to include conferences held between 2015 and 2016. Bibliographic reference lists of eligible SLRs and meta-analyses were also examined for relevant publications. The final search result from each database was limited to references published in the English language.

Eligibility Criteria

Publications involving studies of adults (aged > 18 years) with non-myeloid malignancies receiving myelosuppressive anticancer drugs or adults with acute myeloid leukemia receiving induction or consolidation chemotherapy were included in the review. Publications of interest included primary/secondary G-CSF prophylaxis or treatment with lenograstim (Granocyte®), filgrastim (Neupogen®, Zarzio®, Nivestim®, Ratiogranastim®), pegylated filgrastim (pegfilgrastim, Neulasta®), lipegfilgrastim (Longquex®), Ro 25-8315, empegfilgrastim, maxy-G34, PEG-rHuG-CSF, and BK0026. Eligible studies were further restricted to those reporting a direct, head-to-head comparison of short- versus long-acting G-CSFs. Full details of the inclusion and exclusion criteria are provided in Supplementary Appendix Table S4. Outcomes of interest included the incidence of FN, neutropenia-related or all-cause hospitalizations, and chemotherapy dose reductions or delays due to neutropenia.

Risk-of-Bias Assessment

Two analysts screened the references in parallel, based on title and abstract. Risk of bias for full publications only was assessed by two independent reviewers; disputes were resolved by a third researcher. All publications that met the predefined inclusion/exclusion criteria for the review were obtained as full articles and reassessed against the review criteria. This process was fully compliant with the 2009 Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines.

Meta-analysis

Results were analyzed by pair-wise meta-analysis. Results from relevant studies were entered into a 2 × 2 table and risk ratios were calculated. The Chi squared test of heterogeneity was used to test the assumption that the true effect did not differ between studies. If the Chi squared test gave a low P value or a large Chi squared (I²) statistic relative to its degree of freedom and I² was high, there was evidence of heterogeneity and a random-effect meta-analysis was used. When heterogeneity was low, fixed-effect meta-analysis was used with Mantel–Haenszel methods.

It was originally agreed to exclude publications where the duration of administration of short-acting G-CSF was < 7 days or those where the duration was not reported; however, among the non-RCTs identified, only two studies [6, 27–31] stated that treatment was given ≥ 7 days, and therefore there were insufficient data to perform a sensitivity analysis that included only these studies. One non-RCT reported that long-acting G-CSF was given on the same day as chemotherapy [32]; this study...
was omitted from the meta-analysis. Among the RCTs, one study reported administration of the short-acting G-CSF for < 7 days [33]. A sensitivity analysis for incidence of FN was performed by removing this study. All statistical analyses were performed using Stata (v.12).

Compliance with Ethics Guidelines

This article is based on previously conducted studies and does not contain any studies with human participants or animals performed by any of the authors.

RESULTS

Literature and Conference Search

The original search yielded 3970 articles, 3770 of which were excluded, resulting in 200 articles assessed for eligibility by full review (Supplementary Appendix Fig. 1i). Of these, 162 were excluded on full review and 6 additional publications were identified through supplemental searches. Therefore, a total of 44 references met criteria for inclusion [10, 13–23, 27, 29–59]. The refresher search yielded 530 articles, 514 of which were excluded, resulting in 16 articles assessed for eligibility by full review (Supplementary Appendix Fig. 1ii). Of these, 12 were excluded on full review, and no further relevant abstracts were identified through supplemental searches. Therefore, a total of four additional references met criteria for inclusion: two full publications from abstracts identified in the original screen [6, 28], one abstract containing further data from a previously identified abstract [60], and one poster containing data from a new study [61].

A detailed description of the studies can be found in Supplementary Appendix Tables S5 and S6. In total, 45 publications (17 RCTs; 28 non-RCTs) were included in the SLR [6, 10, 13–23, 27–61]. Of these, 36 (12 RCTs; 24 non-RCTs) were evaluated for the subsequent meta-analysis (Supplementary Appendix Table S7).

RCTs

Seventeen publications reported RCT data: five congress abstracts [34, 42, 47–49, 60], four full foreign language publications with an English abstract [52, 56, 58, 59], and eight full English language publications [33, 35, 36, 45, 51, 53, 55, 57]. Publications reported data from RCTs conducted in China (n = 6) [51, 52, 56–59], multiple countries worldwide (n = 3) [35, 36, 53], Russia (n = 2) [34, 48, 60], India (n = 2) [47, 49], Europe (n = 1) [42], Korea (n = 1) [45], Italy (n = 1) [33], and the United States (n = 1) [55]. Studies compared pegfilgrastim with filgrastim (n = 10) [33, 35, 36, 42, 45, 49, 51, 53, 55, 57], the long-acting G-CSF emepgfilgrastim with filgrastim (n = 2) [34, 48, 60], and a pegylated G-CSF with an unpegylated daily dose G-CSF (drug name not provided; n = 5) [47, 52, 56, 58, 59].

Non-RCTs

Twenty-eight publications reported non-RCT data: 11 congress abstracts [14–17, 29, 31, 39–41, 43, 44], one congress poster presentation [61], and 16 full publications [6, 10, 13, 18–23, 27, 28, 30, 32, 37, 38, 46, 50, 54]. Publications reported non-RCT data from the United States (n = 10) [10, 15, 16, 18–20, 22, 30, 38, 46], Spain (n = 5) [13, 17, 21, 43, 44], Canada (n = 3) [14, 40, 41], Germany (n = 3) [37, 54, 61], Greece (n = 2) [23, 32], multiple countries worldwide (n = 1) [39], Portugal (n = 1) [27, 28], Singapore (n = 1) [29], Austria (n = 1) [50], and the United Kingdom (n = 1) [6, 31].

Twenty-two studies compared filgrastim with pegfilgrastim [6, 10, 14–20, 22, 23, 29–32, 38–41, 43, 44, 46, 54]: in one of these studies, the granulocyte-macrophage colony-stimulating factor sargramostim was administered as a concomitant therapy to both treatment arms [30]. Four studies compared filgrastim or lenograstim with pegfilgrastim [13, 21, 50, 61], and two studies compared a filgrastim biosimilar with filgrastim and pegfilgrastim [27, 28, 37].
Incidence of FN

**Evidence from RCTs**

Eleven RCTs assessed the incidence of FN with short- and long-acting G-CSFs (Supplementary Appendix Table S8) [33–36, 45, 48, 49, 51–53, 57, 60]. Of these, nine studies included patients by specific cancer type [33–36, 45, 48, 49, 53, 57, 60] and two included patients across cancer types [51, 52]. None of the RCTs reported a statistically significant difference in the incidence of FN between short- and long-acting G-CSFs [33–36, 45, 48, 49, 51–53, 57, 60].

**Evidence from Non-RCTs**

Thirteen non-RCTs investigated the incidence of FN with short- and long-acting G-CSFs (Supplementary Appendix Table S8) [10, 13, 14, 18, 21, 23, 27–29, 32, 38, 41, 50, 54]. Of these, 11 were retrospective cohort studies using data from an administrative claims database [13, 18], a pharmacy prescription database [29], patient medical records [10, 14, 21, 38, 50], or records of patients who previously participated in prospective trials [23, 32], one reported a subanalysis of data from an RCT [54, 62], and one reported data from a single-center retrospective study that did not specify the data source [27, 28]. One study was prospective and observational in design [41]. Studies assessed short-versus long-acting G-CSF prophylaxis in patients with breast cancer \( n = 7 \) [14, 23, 27, 28, 32, 41, 50, 54], across cancer types \( n = 5 \) [10, 13, 18, 21, 38], and in patients with lymphoma \( n = 1 \) [29].

Six non-RCT publications reported no significant difference between short- and long-acting G-CSFs [23, 27–29, 38, 41, 50], five reported that pegfilgrastim was significantly superior to filgrastim [10, 13, 14, 18, 54], one reported a numerical trend toward this result but did not perform a statistical analysis [21], and one reported that filgrastim was significantly more effective than pegfilgrastim [32].

**Meta-analysis: Short- versus Long-Acting G-CSFs on the Incidence of FN**

In the analysis of RCTs using a fixed-effect model (Fig. 1i), the overall risk for FN with long-acting G-CSFs was generally lower than with short-acting G-CSFs [overall relative risk (RR) = 0.86]; however, this difference was not statistically significant \( (P = 0.226) \). A sensitivity analysis, excluding one RCT wherein the duration of short-acting G-CSF administration was \(< 7 \text{ days} \) [33], confirmed the original analysis (overall RR = 0.87, \( P = 0.261 \)). In the analysis of non-RCTs using a fixed-effect model (Fig. 1ii), the overall effect of long-acting G-CSF on FN was almost the same as that of short-acting G-CSF (overall RR = 0.98; \( P = 0.681 \)). Using a random- rather than fixed-effect model (Supplementary Appendix Fig. 2), the overall risk for FN with long-acting G-CSF was generally lower than with short-acting G-CSF (overall RR = 0.67); this difference was statistically significant \( (P = 0.023) \).
analysis to be performed. In the meta-analysis of non-RCTs using a fixed-effect model (Fig. 1iii), the overall risk of FN-related hospitalizations associated with long-acting G-CSF was generally lower than with short-acting G-CSF (overall RR = 0.68; \( P < 0.05 \)).

Chemotherapy Dose Reductions or Delays due to Neutropenia

Evidence from RCTs

Four RCTs reported chemotherapy dose reductions and/or delays due to neutropenia; all studies reported similar levels of dose reduction or delay between short- and long-acting G-CSF (Supplementary Appendix Table S10) [33, 35, 36, 42].

Evidence from Non-RCTs

Ten non-RCTs reported chemotherapy dose reductions and/or delays due to neutropenia (Supplementary Appendix Table S10) [6, 13, 17, 21, 23, 27–29, 31, 32, 41, 54]. Of these, two reported similar incidences of dose reduction between daily G-CSF and pegfilgrastim [27, 28, 54], three reported no difference between treatments for this outcome or chemotherapy delays [29, 32, 41], two reported a statistically significant result for higher incidence of dose reduction or delay with daily G-CSF versus pegfilgrastim [13, 23], and two reported a numerical trend toward this result [17, 21]. The remaining non-RCT reported that 84.9% (pegfilgrastim) versus 69.5% (filgrastim) of patients achieved ≥ 85% relative dose intensity; authors did not state whether this was a significant result [31].

Meta-analysis: Short- versus Long-Acting G-CSFs on the Incidence of Chemotherapy Dose Reductions and Delays

There were not enough RCTs reporting chemotherapy dose reductions and delays to enable a meta-analysis. In the meta-analysis of non-RCTs using a fixed-effect model (Fig. 2), the overall risks of chemotherapy dose reductions (overall RR = 0.69, \( P < 0.05 \)) and delays (overall RR = 0.70, \( P < 0.05 \)) with long-acting G-CSF were generally lower than with short-acting G-CSF. In the meta-analysis of non-RCTs using a random-effect model (Supplementary Appendix Fig. 3), the overall risk of chemotherapy delays with long-acting G-CSF was generally lower than with short-acting G-CSF (overall RR = 0.68, \( P = 0.020 \)).

DISCUSSION

This SLR identified RCTs and non-RCTs that reported head-to-head comparisons of short-versus long-acting G-CSFs for the reduction of chemotherapy-induced FN. When short-acting G-CSF was dosed according to treatment guidelines, there was no statistically significant difference in the incidence of FN, hospitalizations, and chemotherapy dose reductions and/or delays between short- and long-acting G-CSFs among RCT results. This is important because FN-related hospitalizations are common, distressing, and costly [2, 4]. This analysis confirms the recommendations of the American Society of Clinical Oncology and the European Society of Clinical Oncology that indicate that either filgrastim or pegfilgrastim can be used for the prevention of treatment-related FN, with the choice of agent dependent on convenience, cost, and clinical situation [63, 64].

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The approved dosing schedule for short-acting G-CSF treatment is 5 μg/kg once daily, commencing 24 h after chemotherapy and continuing for up to 2 weeks or until the ANC has recovered (10,000/mm³) following its chemotherapy-induced nadir [9]. However, an analysis of administrative claims data demonstrated that almost half of filgrastim users received treatment for 6 days [46]. Furthermore, an observational study reported lower adherence to treatment guidelines and recommendations for timing of G-CSF initiation with use of filgrastim/lenograstim versus pegfilgrastim [65]. Among the studies identified in this SLR, few non-RCTs (2 of 24) and nearly all RCTs (11 of 12) indicated that short-acting G-CSF was administered for ≥ 7 days [6, 27–31, 34–36, 42, 45, 48, 49, 51–53, 57, 60]. This observation supports other evidence of under-dosing of short-acting G-CSFs in real-world clinical practice and that compliance is likely to be lower in routine clinical practice compared with RCTs. Furthermore, this may explain, in part, the difference between the results from the RCT and non-RCT studies in this analysis.

Most (11 of 17) RCTs included in this meta-analysis were described as open-label [36, 42, 45, 47, 48, 51, 52, 55, 57–59]. The absence of blinding in open-label studies may increase the risk of performance and detection bias. However, study design appeared to have minimal impact on the relative efficacy of short- versus long-acting G-CSFs; only 1 of 11 open-label studies reported a significant result across all outcomes of interest, with longer duration of grade 3+ neutropenia reported for
patients treated with short- versus long-acting G-CSF [57]. Furthermore, a risk of bias assessment (results not shown) indicated the majority of studies were of high quality, and most were published from 2010 onward. Substantial changes in clinical practice over this period were not likely, which may have impacted treatment response in patients.

Some inherent limitations of this type of review are acknowledged: some FN events may not have been captured accurately, differences in the algorithms used for FN make comparisons between studies difficult, and some studies may have been underpowered to detect any differences in FN and FN-related endpoints. It is also important to note that new studies may have been published since completion of the systematic review (June 2016). Additionally, for the evidence presented from retrospective studies (from sources such as claims databases and pharmacy prescription databases), the accuracy and completeness of the information is uncertain.

Overall, the weight of evidence indicates little difference in efficacy between the short- and long-acting drugs if the short-acting G-CSF is dosed according to recommended guidelines (RCT data) [33–36, 42, 45, 48, 49, 51–53, 57, 60]. There is some evidence for greater efficacy with long-acting G-CSFs in non-RCT studies [10, 13–23, 54, 61]. However, this may be a result of under-dosing of short-acting G-CSFs in real-world general practice. If there is no clinical difference between short- and long-acting G-CSFs, then treatment choice could be based on differences in patient quality of life or economic factors. Patients may prefer long-acting G-CSF treatment because fewer injections are needed [66]. Currently, the cost of the long-acting drug may outweigh any benefit of convenience. Filgrastim is less expensive than pegfilgrastim [30, 37, 67]. Nevertheless, use of fewer injections in routine practice than is recommended may be an attempt to limit treatment costs [24].

Biosimilars, biologic products that are highly similar to a licensed (i.e., reference or originator) biologic drug [68, 69], may provide a lower-cost alternative to originator biologic therapies. Filgrastim biosimilars are available worldwide [70]; the availability of short-acting G-CSF biosimilars might be a way to regulate cost while maintaining the recommended dosing schedule [24] to ensure improved health outcomes. A 2014 annual review issued by the Pharmaceutical Management Agency of New Zealand reported a significant reduction of costs in the total G-CSF market and a nearly 25% expansion in G-CSF usage following introduction of biosimilar filgrastim in 2012 [71]. Better access to G-CSFs also had a significant public health impact; the incidence of FN in women receiving chemotherapy for breast cancer decreased from one-third to less than 7% [71]. Biosimilars of pegfilgrastim are only just becoming available; the availability of long-acting G-CSF biosimilars may improve access to treatments that are more convenient for patients while remaining cost-effective.

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Compliance with Ethics Guidelines. This article is based on previously conducted studies and does not contain any studies with human participants or animals performed by any of the authors.

Data Availability. All data generated or analyzed during this study are included in this published article/as supplementary information files.

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