Epoetin alfa in platinum-treated ovarian cancer patients: results of a multinational, multicentre, randomised trial

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This multicentre, open-label, controlled clinical trial assessed the effects of epoetin alfa treatment on haematologic and quality of life (QOL) parameters in 182 anaemic (Hb ≤ 12 g dl⁻¹) ovarian cancer patients receiving platinum chemotherapy. Patients were randomised 2:1 to receive epoetin alfa 10 000–20 000 IU three times weekly plus best standard treatment (BST) or BST only. Main study end points were changes from baseline in haemoglobin (Hb) level, transfusion requirements, and QOL. For the epoetin alfa group, mean Hb increased by 1.8 g dl⁻¹ by weeks 4–6 and was significantly increased from baseline through study end (P < 0.001). The mean change in Hb from baseline was significantly (P < 0.001) greater for epoetin alfa than BST patients at all postbaseline evaluations. Significantly fewer epoetin alfa than BST patients required transfusion(s) after the first 4 weeks of treatment (7.9 vs 30.5%; P < 0.001). Also, significant (P < 0.04) differences favouring the epoetin alfa group over the BST group were found for all three median CLAS scores (Energy Level, Ability to Do Daily Activities, Overall QOL) and the median average CLAS score during chemotherapy. These findings support use of epoetin alfa to increase Hb levels, reduce transfusion use, and improve QOL in anaemic ovarian cancer patients receiving platinum chemotherapy.

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Ovarian cancer is the fourth most common cancer in women and the second most common gynaecological cancer (Harries and Gore, 2002a, b). Although the 5-year survival rate for women with low-risk stage I epithelial ovarian cancer can be as high as 90% (Memarzadeh and Berek, 2001), this form of cancer is frequently not detected until it is in its advanced stages with corresponding 5-year survival rates of 20–40 and 10% for stage III and stage IV disease, respectively (Heintz et al, 2001). Standard therapy for ovarian cancer is cytoreductive surgery, followed in most cases by platinum/paclitaxel combination chemotherapy (Harries and Gore, 2002a). Although chemotherapy has been shown to increase survival for women with ovarian cancer, most eventually relapse and die (Harries and Gore, 2002b). Therefore, palliating symptoms and maintaining quality of life (QOL) have become primary goals in disease management.

Anaemia in ovarian cancer patients, while often related to the disease itself, also commonly results from myelosuppression induced by repeated cycles of platinum-based chemotherapy. In a large-scale audit of patients in the United Kingdom receiving chemotherapy, which included 856 patients with ovarian cancer, the proportion of ovarian cancer patients with anaemia (haemoglobin (Hb) < 11 g dl⁻¹) rose from about 25% after chemotherapy cycle 1 to 50% after cycle 6, despite 41% of these patients having received at least one blood transfusion during treatment (Barrett-Lee et al, 2000). The most commonly administered chemotherapeutic agents administered in this subgroup were carboplatin (63%), and a combination of doxorubicin + cisplatin + cyclophosphamide (15%).

It is well known that severe anaemia is associated with an array of debilitating symptoms; however, even mild anaemia (Hb level 10–12 g dl⁻¹) can have serious negative consequences for patients. An Hb level < 12 g dl⁻¹ has been associated with increased risk of transfusion (Ray-Coquard et al, 1999), increased fatigue (Cella, 1997; Holzner et al, 2002), and a less-than-optimal QOL (Crawford et al, 2002). Recently, important data on anaemia and performance status were obtained by the European Cancer Anaemia Survey (ECAS). A prospective, multinational observational survey, ECAS evaluated the prevalence, incidence, and treatment of anaemia in 15 367 European cancer patients, 1741 of whom had gynaecologic malignancies (Ludwig et al, 2004). Analysis of the survey data showed that the prevalence and incidence of anaemia in the gynecologic subgroup were 81.4 and 74.8%; however, despite its high prevalence and incidence, anaemia remained untreated in 57.3% of patients with this symptom (Barrett-Lee et al, 2005). The analysis additionally showed a significant correlation between low Hb levels and poor performance score, as assessed by WHO criteria (P < 0.001, R = −0.18). From this it can be inferred that a substantial proportion of anemic gynaecologic cancer patients experience a decline in functional capacity, with a subsequent decline in QOL. Given the many negative consequences of anaemia for ovarian cancer patients, maintaining optimal Hb levels should be considered an essential aspect of supportive care.
Epoetin alfa therapy has been shown in both double-blind, placebo-controlled, and open-label studies to increase Hb levels in cancer patients receiving platinum- or nonplatinum-based chemotherapy, correcting anaemia, decreasing transfusion requirements, and subsequently improving patients’ QOL (Abels, 1992; Leitgeb et al., 1994; Glaspy et al., 1997; Demetri et al., 1998; Dammacco et al., 2001; Gabrilove et al., 2001; Littlewood et al., 2001; Thomas, 2002; Janinis et al., 2003; Shasha et al., 2003; Savonije et al., 2004; Chang et al., 2005; Witzig et al., 2005). However, few studies to date have examined the outcome of anaemia treatment in ovarian cancer patients, and none have specifically evaluated the impact of anaemia treatment on QOL in this population (ten Bakker Huinkink et al., 1998). Given the relative lack of data on anaemia treatment in ovarian cancer patients, we conducted a study to determine the possible benefits of epoetin alfa treatment with respect to transfusion reduction, QOL, and anaemia-related symptoms, including fatigue, in anemic ovarian cancer patients.

MATERIALS AND METHODS

Study patients and design

This was a phase IV, multinational, multicentre, randomised, open-label, comparative clinical trial conducted between March 1999 and June 2001. Female patients 18 years of age or older with a confirmed diagnosis of ovarian cancer, at least mild anaemia (Hb <12 g dL⁻¹), and an ECOG performance score of 0, 1, 2, or 3 were enrolled. Patients were to be receiving or scheduled to receive platinum-based chemotherapy and were to have a life expectancy of at least 6 months. Patients with untreated iron, folate, or vitamin B₁₂ deficiency or anaemia due to factors other than cancer or its treatment were excluded. Also excluded were those who had received a blood transfusion within 14 days prior to study entry or who had severe illness or surgery within 7 days of study entry. All patients provided written informed consent before study entry. The study was undertaken after approval of the protocol by the Independent Ethics Committee of each centre, and was conducted in accord with the Guidelines for Good Clinical Practice and the Declaration of Helsinki, South Africa amendment 1996.

Enrolled patients were randomised 2:1 to receive either epoetin alfa plus best standard treatment (BST; transfusion of red blood cells, as needed) or BST only. Outside of the United States, epoetin alfa is manufactured by Ortho Biologics, LLC, and distributed and marketed as EPREX® or ERYPO® by Ortho Biotech, a division of Janssen-Cilag. Epoetin alfa was administered initially at a dosage of 10 000 IU three times weekly (t.i.w.) subcutaneously (or 5000 IU t.i.w. for patients with body weight <45 kg). The initial dose was maintained throughout the first cycle of a 4-week cycle of chemotherapy or the first two cycles of a three-week cycle. If, at the end of the initial period, the reticulocyte count had not increased by >40 000 µL⁻¹, or Hb had not increased by >1 g dL⁻¹ above the baseline level, the dose of epoetin alfa was doubled (maximum allowed dosage, 20 000 IU t.i.w.). If the Hb level exceeded 14 g dL⁻¹ at any time, study drug was withheld until the Hb level had declined to <12 g dL⁻¹ and then was restarted at a dose 25–50% lower than the previous dose. If the Hb level increased by ≥2 g dL⁻¹/month⁻¹, the dose was reduced by 25–50% to maintain the Hb rate of increase at <2 g dL⁻¹/month⁻¹. Dosage reductions could be achieved by omitting one of the weekly doses of epoetin alfa. No adjustment to the dosage was made if Hb level increased in response to transfusion. The planned duration of study treatment was a maximum of 28 weeks, which included 18–24 weeks of chemotherapy (maximum, six cycles) plus up to 4 weeks after the last chemotherapy dose.

In both arms, red blood cell transfusion was permitted during the study if judged necessary, but physicians were asked to refrain from transfusing patients unless the Hb level was <9 g dL⁻¹.

Administration of white cell growth factor was permitted, and a daily dose of 200-mg elemental iron as oral iron supplementation was recommended to prevent restriction of erthropoesis. Transferrin saturation ≤20% was considered indicative of inadequate iron stores and iron deficiency.

Efficacy and safety evaluations

The primary efficacy end point was the difference between the treatment groups in change in Hb level from baseline to study end. Secondary efficacy end points included within-group change in Hb level from baseline to study end and between-group differences in proportions of patients considered complete responders, partial responders, or nonresponders. Complete responders were defined as patients who demonstrated an Hb increase ≥1 g dL⁻¹ above baseline without transfusion within the preceding 4 weeks. Partial responders were defined as patients who achieved an Hb increase of ≥0.5 g dL⁻¹ but <1 g dL⁻¹, and nonresponders, as those who either were transfused or demonstrated an Hb increase of <0.5 g dL⁻¹ above baseline.

Other efficacy end points included change in proportion of patients transfused and change in QOL scores from baseline to study completion. Quality of life was assessed using the patient-rated Cancer Linear Analog Scale (CLAS, also known as the Linear Analog Scale Assessment (LASA)), which measures Energy Level, Ability to Do Daily Activities, and Overall QOL, and the Functional Assessment of Cancer Therapy-Anaemia (FACT-An), for which the FACT-General (FACT-G Total) scale, FACT-An Fatigue subscale, and Nonfatigue subscale were assessed. Both the FACT-An and the CLAS scales are cancer specific and sensitive to Hb levels (Cella, 1997; Glaspy et al., 1997; Demetri et al., 1998; Gabrilove et al., 2001). Haemoglobin levels, transfusion data, and QOL scores were obtained within 7 days prior to the first dose of study medication and at study completion or early termination. During the study, Hb levels were measured and transfusion data collected on completion of study weeks 4 or 6, 8 or 9, 12, and 16 or 18; QOL was assessed after 4 or 6 weeks (CLAS only), 8 or 9 weeks (CLAS and FACT-An), and 12 weeks (CLAS only). Additionally, tumour response to chemotherapy and/or radiotherapy was assessed at study end or the final visit.

Safety and tolerance of epoetin alfa were evaluated by the usual methods, including monitoring adverse events. Adverse events were reported by patients throughout the study either spontaneously or in response to general, nondirect questioning by the investigator.

Statistical analyses

The primary analysis was based on the intent to treat (ITT) population. For efficacy evaluations, changes between baseline and each monthly value for Hb level were analysed using both analysis of variance (ANOVA) and the Wilcoxon rank-sum test. The proportion of patients transfused during the treatment period was analysed by Fisher’s exact test, changes from baseline in QOL scores were analysed using the Wilcoxon signed-rank-sum test, and tumour stage was compared using the Wilcoxon rank-sum test. All P-values were unadjusted and were derived from two-sided tests. A P-value of ≤0.05 was considered to indicate statistical significance.

A total of 145 evaluable patients were required to complete the study to have a 90% power to detect a difference (2.0 g dL⁻¹) between the epoetin alfa and BST groups in change in Hb from baseline to last evaluation (primary variable), with randomisation assignment to one of two treatment arms in a 2:1 ratio. Tests of significance were one- or two-sided, with α set at 0.05 or 0.025. The study was not powered for secondary efficacy variables, including QOL.
RESULTS

In total, 182 patients were enrolled in the study, 173 (114 epoetin alfa; 59 BST) of whom were eligible for efficacy evaluation (ITT population). The nine ineligible patients were excluded because of data recording on differently designed case report forms (4), misdiagnosis (3), or withdrawal of patient consent (2). The majority (91) of evaluable patients were seen at centers in the United Kingdom; the remaining patients were treated at centers in Austria (27), Greece (26), Sweden (22), The Netherlands (5), and Denmark (2). Of the 173 patients, 145 completed the study. The other 28 were discontinued prematurely for the following reasons: adverse event, insufficient response, noncompliance, asymptomatic/cure, disease progression, chemotherapy discontinued, and cancer not ovarian. Three of the 28 patients, all in the epoetin alfa group, died during the study as a result of their malignancy. All three patients were included in the study despite having a life expectancy of <6 months at entry, and their deaths were not unexpected.

Of 119 patients with available drug exposure information, 117 commenced dosing with 30 000 IU week \(^{-1}\), whereas two patients commenced dosing with 20 000 and 27 000 IU week \(^{-1}\), respectively. (The two latter patients violated the protocol requirement for an initial starting dose of 30 000 IU week \(^{-1}\), but this was considered a minor violation and the patients were therefore included for analysis.) In total, 10 patients (8%) required dose increases during the study.

Table I  Demographic and clinical characteristics at baseline (intent-to-treat population, \(N = 173\))

| Characteristic           | Epoetin alfa (\(n = 114\)) | Best standard treatment (\(n = 59\)) |
|--------------------------|----------------------------|-------------------------------------|
| Mean age, year (± s.d.)  | 59.1 (± 10.6)              | 60.3 (± 11.2)                       |
| Range                    | 35.0–87.0                  | 30.0–79.0                           |
| Mean Hb level (g dl\(^{-1}\)) | 10.75 ± 0.94              | 10.66 ± 0.83                        |
| ECOG performance score (n, %)* | |                                     |
| 0                        | 56 (49.1)                  | 27 (45.8)                           |
| 1                        | 47 (41.2)                  | 29 (49.2)                           |
| 2                        | 11 (9.6)                   | 3 (5.1)                             |
| 3                        | 0 (0.0)                    | 0 (0.0)                             |
| 4                        | 0 (0.0)                    | 0 (0.0)                             |
| Tumour stage, n (%)      |                            |                                     |
| I                        | 12 (10.5)                  | 10 (16.9)                           |
| II                       | 13 (11.4)                  | 3 (5.1)                             |
| III                      | 58 (50.9)                  | 32 (54.2)                           |
| IV                       | 28 (24.6)                  | 13 (22.0)                           |
| Unknown                  | 3 (2.6)                    | 1 (1.7)                             |
| Metastatic disease, n (%)|                            |                                     |
| Unknown                  | 1 (0.9)                    | 0 (0.0)                             |
| None                     | 27 (23.7)                  | 13 (22.0)                           |
| Abdominal                | 66 (57.9)                  | 32 (54.2)                           |
| Liver                    | 12 (10.5)                  | 6 (10.2)                            |
| Lymphatic                | 12 (10.5)                  | 6 (10.2)                            |
| Lung                     | 7 (6.1)                    | 5 (8.5)                             |
| Other type               | 34 (29.8)                  | 17 (28.8)                           |
| Previous surgery, n (%)  |                            |                                     |
| n                        | 94 (82.5)                  | 49 (83.1)                           |

*0 = able to carry out normal activities, 1 = restricted physical activity/ambulatory/light work, 2 = ambulatory/capable of all self-care/unable to work, 3 = capable of only limited self-care, 4 = completely disabled.

Baseline demographic and clinical characteristics were generally comparable between the two treatment groups (Table I). Mean Hb levels at baseline were 10.8 g dl\(^{-1}\) for patients given epoetin alfa and 10.7 g dl\(^{-1}\) for those given BST. More than 90% of the patients in each group were receiving carboplatin or carboplatin plus paclitaxel, and the remainder were receiving cisplatin. The frequencies of 3- and 4-week chemotherapy cycles also were comparable between the two groups (3-week cycles: epoetin alfa, 73.7% ; BST, 72.9%) (4-week cycles: epoetin alfa, 26.3%; BST, 27.1%).

Haematopoietic response

The evaluation of haematopoietic response was based on the 171 patients who received uninterrupted treatment. In the epoetin alfa group, the Hb level increased by a mean of 1.8 g dl\(^{-1}\) after the first 4–6 weeks of treatment, and was significantly (\(P<0.001\)) increased above baseline at all time points (Figure 1). In contrast, Hb levels in the BST group changed little over the course of treatment. Differences between the epoetin alfa and BST groups were significant (\(P<0.001\), ANOVA or t-test) at all post-baseline evaluations. The highest Hb levels in the epoetin alfa group were observed after weeks 8–9 and 12. Mean ± s.d. increases in Hb level from baseline after 8–9 weeks were 2.0 ± 1.5 g dl\(^{-1}\) for the epoetin alfa group vs 0.3 ± 1.0 g dl\(^{-1}\) for the BST group; corresponding values after 12 weeks were 1.8 ± 1.3 vs 0.0 ± 1.1 g dl\(^{-1}\). At study completion or early termination, mean changes in Hb level from baseline were 1.6 ± 1.5 and 0.3 ± 1.3 g dl\(^{-1}\), respectively, for the epoetin alfa and BST groups.

The distribution of patients in the two treatment groups by Hb level during treatment is illustrated in Figure 2. As shown, patients treated with epoetin alfa above the 25th percentile had Hb values higher than those for BST patients in the 75th percentile at weeks 8–9 and week 12, and in the 95th percentile after weeks 16–18. Median (interquartile range) Hb levels for the epoetin alfa and BST groups were 13.0 (2.5) vs 11.0 (1.4) g dl\(^{-1}\), respectively, after 8–9 weeks, and 12.9 (1.5) g dl\(^{-1}\) vs 10.8 (1.5) g dl\(^{-1}\), respectively, after 12 weeks. Overall, more patients in the epoetin alfa group than in the BST group had Hb increased by ≥ 1 g dl\(^{-1}\) (responders, 78 vs 32% for epoetin alfa and BST, respectively; Table 2). Conversely, fewer patients in the epoetin alfa group did not respond to treatment. The difference in the proportions of both responders...
and nonresponders between the treatment groups was statistically significant ($P < 0.001$).

**Transfusion use**

A significantly smaller proportion of patients in the epoetin alfa group (nine out of 114, or 7.9%) than in the BST group (18 out of 59, or 30.5%) were transfused at least once after the first 4 weeks of treatment ($P < 0.001$, Fisher’s exact test). Also, significant differences in transfusion rate favouring epoetin alfa were noted at all evaluations except week 12, at which time the difference favoured epoetin alfa, but not significantly: week 4 or 6, 5.9% vs 16.1%, $P = 0.048$; week 8 or 9, 0.0% vs 14.0%, $P < 0.001$; week 12: 1.6% vs 5.3%, not significant; weeks 16–18, 0.0% vs 19.2%, $P = 0.007$; and up to 28 weeks: 1.8% vs 13.8%, $P = 0.004$.

**Quality of life**

Of the 173 evaluable patients, 102 (64 epoetin alfa, 38 BST) had paired CLAS data for baseline and after 12 weeks, and 141 (91 epoetin alfa, 50 BST) had such data for baseline and end of study. Analysis of these data (Figure 3) showed significant differences from baseline favouring epoetin alfa over BST for all three CLAS change scores (Energy Level, Ability to Do Daily Activities, Overall QOL) and the average median CLAS change score during chemotherapy ($P \leq 0.04$; after 12 weeks: $P \leq 0.003$; Wilcoxon signed rank-sum test). At the final visit, the median increase was 10.0 mm for each parameter in the epoetin alfa group compared with median increases of 1.25–2.85 mm for these parameters in the BST group; differences between the groups did not achieve statistical significance ($P = 0.054–0.118$). Results of within-treatment analysis showed that median scores for each of the three CLAS scales and the average median CLAS score were significantly increased from baseline at all four evaluation points in the epoetin alfa group ($P < 0.001$, Wilcoxon signed rank test), whereas no significant change from baseline was detected at any evaluation point in the BST group. In the epoetin alfa group, the average median CLAS score increased by 36% from baseline (56.85 mm) to last observation (77.30 mm), with increases of up to 45% (to 82.70 mm) noted after 12 weeks. In contrast, the BST group showed little change in average median score from baseline (62.00 mm) to last observation (70.00 mm) or to any evaluation point during chemotherapy (maximum, 63.30 mm after 8 or 9 weeks).

The study was underpowered for FACT-An analysis. However, univariate analysis demonstrated trends favouring epoetin alfa over BST for the FACT-An Nonfatigue score ($P = 0.087$) after 8–9 weeks and for the FACT-G Total ($P = 0.081$), FACT-An Fatigue ($P = 0.173$), and FACT-An Nonfatigue ($P = 0.082$) subscale scores at study end.

**Tumour response**

At the end of treatment, the two study groups were similar with respect to the proportions of patients in each group with complete response, partial response, or no response to cancer treatment (Table 3). However, proportionally more patients in the epoetin alfa group than in the BST group had progressive disease, although this difference was not significant ($11$ vs $2\%$, $P = 0.425$). Examination of the patients’ medical histories showed that the
The results of our study show that epoetin alfa can effectively increase Hb levels and reduce transfusion use for ovarian cancer patients receiving platinum-based chemotherapy. The mean Hb levels for the epoetin alfa group were significantly increased from baseline at all evaluations (P<0.001) and 78% of patients who received epoetin alfa demonstrated a complete response to anaemia treatment (Hb increase ≥1 g dl⁻¹ without transfusion within the previous 4 weeks). Response was rapid, with a 1.8 g dl⁻¹ increase in mean Hb level at 4–6 weeks, and a maximum increase of 2.0 g dl⁻¹ at 8–9 weeks. This rate of Hb increase compares favourably to rates seen in previous studies in patients with gynaecologic or other tumour types. In two large studies with patients receiving platinum or nonplatinum chemotherapy, mean Hb increases were about 1 g dl⁻¹ at week 4 and 2 g dl⁻¹ at week 8 (Gabrilove et al, 2001; Littlewood et al, 2001). In a study of epoetin alfa in patients with gynaecologic malignancies receiving poly-

**Table 3** Tumour response to cancer treatment

| Tumour response | Epoetin alfa (n = 114) | Best standard treatment (n = 59) |
|-----------------|-----------------------|---------------------------------|
| Complete, n (%) | 55 (48)               | 33 (56)                         |
| Partial, n (%)  | 25 (22)               | 19 (32)                         |
| None, n (%)     | 7 (6)                 | 5 (9)                           |
| Progressive disease, n (%) | 13 (11) | 1 (2) |
| Unknown, n (%)  | 14 (12)               | 1 (2)                           |

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23 trials showed that epoetin alfa therapy significantly improved CLAS (20–25%), FACT-Fatigue (17%), and FACT-An (12%) scores from baseline \( (P = 0.05) \), whereas scores for control groups were relatively unchanged or worsened \( (Jones et al., 2004) \). Of interest, one of these trials, a randomised, double-blind controlled study \( (Littlewood et al., 2001) \), showed that although a significantly greater proportion of placebo-treated patients than epoetin alfa-treated patients were transfused, the mean Hb level of the placebo patients was unchanged from baseline over the course of the study and their QOL did not improve, but rather, worsened. In contrast, epoetin alfa-treated patients demonstrated a significant increase in Hb level from baseline, and showed significant improvement in QOL domains.

The incidence of adverse events was similar between the epoetin alfa and BST groups in this study, although patients in the epoetin alfa group had a higher incidence of TVEs \( (12 \text{ events vs } 1 \text{ event}; \text{incidence } 8.3 \% \text{ vs } 1.7 \%) \). These findings are consistent with those of other randomised studies in which patients received epoetin alfa administered as either a t.i.w. \( (Dammacco et al., 2001; \text{Littlewood et al., } 2001) \) or once-weekly \( (q.w.) \) \( (Chang et al., 2005; Witzig et al., 2005) \) regimen. The overall incidence of adverse events in these studies ranged from 73 to 88% for patients in the epoetin alfa groups and from 75 to 86% for those in the control groups. Also, the incidence of TVEs was comparable to that reported in our study, ranging from 5 to 10.8% for the epoetin alfa groups and 3 to 7.9% for the control groups. The Hb level required for study enrollment/randomisation and baseline Hb in these studies are shown in Table 4. That all erythropoiesis stimulating agents may increase the risk for TVE development is well established and this information is in the agents’ product labelling. It must be mentioned in this regard, however, that the results of two recently reported studies have suggested an adverse impact on survival conferred by erythropoiesis stimulating agents \( (Henke et al., 2003; Leyland-Jones, 2003) \), and that in one of the studies, increased mortality was considered partly attributable to TVEs \( (Leyland-Jones, 2003) \). Both studies included cancer patients who would not normally receive erythropoiesis stimulating agents, namely, nonanemic patients, and patients with Hb levels higher than those recommended in the approved labelling. Interpretation of the results of these studies is complicated due to the study designs and imbalance of risk factors in the populations. In contrast, a meta-analysis of 27 randomised, controlled studies \( (N = 3287) \) of recombinant human erythropoietin (RHuEPO) showed that the relative risk for thromboembolic complications after RHuEPO treatment was not significantly increased compared with that of untreated patients \( (RR = 1.58, 95\% \text{ CI } = 0.94–2.66; 12 \text{ trials, } N = 1738) \). The absolute risk difference was 0.02 \( (95\% \text{ CI } 0.00–0.04) \), although the number of patients with no thrombotic event may have been underreported \( (Bohlius et al., 2005) \). Further, there was evidence of a trend toward improved overall survival with RHuEPO treatment. From the preceding, it can be concluded that epoetin alfa is safe and well tolerated when administered according
to labelling, and that the risk of TVE development in cancer patients receiving epoetin therapy may be substantially limited by targeting the Hb concentration to around 12 g dl⁻¹.

The natural history of ovarian cancer poses a unique challenge to anaemia management. In the majority of patients, this cancer is not diagnosed until it has reached more advanced stages. Although patients with advanced ovarian cancer typically respond well to first-line chemotherapy, most relapse and become candidates for further chemotherapy (Harries and Gore, 2002b). Prolonged disease, surgery, and repeated cycles of platinum-based or other chemotherapy all contribute to the development of anaemia in this population. Anaemia has been demonstrated to negatively affect cancer patients by increasing risk of transfusion and reducing QOL. The results of the present study support the use of epoetin alfa in anemic ovarian cancer patients to achieve a reliable haematologic response that is maintained throughout treatment. Evidence from this study further supports increasing Hb levels to ameliorate anaemia as a means of improving QOL – a primary goal of treatment for ovarian cancer patients with advanced disease and treatment-related anaemia.

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Appendix

The following investigators participated in this study: Dr K Joerg, Villach, Austria; Dr H Hausmaninger, Salzburg, Austria; Dr M Lahousen, Graz, Austria; Dr J Maier, Wels, Austria; Dr MR Mirza, Odense, Denmark; Dr M Antonopoulos, Athens, Greece; Dr P Kosmidis, Athens, Greece; Dr P Paraskevopoulos, Thessaloniki, Greece; Dr Hdankbaar, Hengelo, The Netherlands; Dr HP Sleeboom, Den Haag, The Netherlands; Dr P Willemse, Groningen, The Netherlands; Dr H Andersson, Göteborg, Sweden; Dr H Malmstrom, Linkoping, Sweden; Dr M Ridderheim, Lund, Sweden; Dr J Davis, Glasgow, UK; Dr A Hong, Wonford, UK; Dr CJ Irwin, Coventry, UK; Professor M Lind, Hull, UK; Dr P Murray, Colchester, UK; Dr SD Pledge, Sheffield, UK; Professor N Reed, Glasgow, UK; Dr H Thomas, Guildford, UK; Dr PM Wilkinson, Manchester, UK.

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