A randomised, placebo-controlled, double-blind study of aprepitant in nondrinking women younger than 70 years receiving moderately emetogenic chemotherapy

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Background: We evaluated the efficacy of aprepitant plus granisetron and an increased dose of dexamethasone in selected patients undergoing moderately emetogenic chemotherapy (MEC).

Methods: Nondrinking women <70 years undergoing MEC were randomly assigned to aprepitant (day 1, 125 mg; days 2 and 3, 80 mg) or placebo. Dexamethasone on days 1–3 was 12, 4, and 4 mg with aprepitant and 20, 8, and 8 mg with placebo. The primary end point was complete response (CR; no emesis or rescue therapy) during 120 h of the first cycle. Logistic regression analysis was performed to identify predictors of overall CR.

Results: Of the 94 patients enrolled, 91 were assessable. Most received carboplatin-based chemotherapy. In the aprepitant (n = 45) and placebo (n = 46) groups, the overall, acute (day 1), and delayed (days 2–5) CR rates were 62% and 52%, 98% and 96%, and 62% and 52%, respectively. Although not statistically significant, the overall CR rate was 10% higher in the aprepitant group. Both regimens were well tolerated. On multivariate analysis, advanced ovarian cancer (OR, 0.26 (0.10–0.72)) was independently associated with a lower CR.

Conclusion: Even with an increased dose of dexamethasone, aprepitant seemed more effective than placebo in these selected patients undergoing MEC; however, delayed phase management remains a significant problem.

Despite considerable progress in prevention, chemotherapy-induced nausea and vomiting (CINV) remain the most feared adverse effects among patients with cancer. Uncontrolled CINV can limit the dose intensity of chemotherapy and seriously compromise a patient’s quality of life (Oo and Hesketh, 2005). The incidence of CINV depends primarily on the dose and type of chemotherapeutic agents administered. Published guidelines consistently classify carboplatin, irinotecan, and oxaliplatin as moderately emetogenic chemotherapy (MEC; Kris et al, 2006; Roila and Fatigoni, 2006). The risk of CINV also depends on gender and age; female and younger patients are at

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greater risk. In contrast, patients with a history of heavy alcohol consumption have a lower risk of CINV (Pollera and Giannarelli, 1989; Oseba et al., 1997; Schwartzberg, 2007; Hesketh, 2008; Hesketh et al., 2010).

For MEC regimens not based on anthracycline/cyclophosphamide (AC), recent guidelines, such as the National Comprehensive Cancer Network (NCCN) Antiemesis Guidelines version 1.2012 (Ettinger et al., 2012) and the American Society of Clinical Oncology (ASCO) guidelines for antiemetics (Basch et al., 2012), generally recommend the use of a 5-hydroxytryptamine 3 (5-HT3) receptor antagonist and dexamethasone ‘with or without’ the neurokinin-1 receptor antagonist, aprepitant. Aprepitant can enhance the prevention of CINV in patients who receive highly emetogenic chemotherapy (HEC), such as cisplatin (Hesketh et al., 2003; Poli-Bigelli et al., 2003; Schmoll et al., 2006) or AC (Warr et al., 2005). According to the NCCN guidelines, aprepitant use is recommended only for select patients receiving MEC, such as carboplatin or irinotecan. However, the characteristics of these ‘select’ patients are unclear, and no randomised trials support this strategy for non-AC MEC. The Multinational Association of Supportive Care in Cancer (MASCC) does not recommend aprepitant for non-AC MEC regimens (Roila et al., 2010). Rapoport et al. (2010) reported a phase III, gender-stratified trial of aprepitant in 848 patients, showing that aprepitant significantly improved the primary end point of no vomiting as well as the secondary end point of complete response (CR) rate after MEC, including AC or non-AC regimens. Dexamethasone was administered only on day 1. In the subgroup analysis of 428 patients undergoing non-AC MEC, although the primary end point of no vomiting was significantly different (aprepitant group, 83.2%; placebo group, 71.3%), the secondary end point of overall CR rate did not differ significantly between the groups (aprepitant group, 74.2%; placebo group, 65.5%). Addition of dexamethasone on days 2 and 3 may have increased the overall CR rate and reduced the difference in efficacy between aprepitant and placebo. This trial was not considered sufficiently compelling to recommend the standard use of aprepitant in non-AC chemotherapy. However, Waqar et al. (2008) reported that among lung cancer patients, vomiting occurred in a higher proportion of women (31%) compared with men (8%) within 72 h after carboplatin administration (area under the curve (AUC), 5), suggesting that aprepitant might be effective in ‘select’ patients, such as women.

Corticosteroids are recommended for the prevention of acute and delayed emesis following HEC and MEC (Ioannidis et al., 2000). The recommended dose of dexamethasone on the first day of MEC is 8 mg (Roila et al., 2010; Basch et al., 2012) or 12 mg (Ettinger et al., 2012). On the other hand, a 20-mg dose of dexamethasone may prevent CINV more effectively in patients who receive HEC (IGAR, 1998). Although carboplatin-based MEC is less emetogenic than HEC, the optimal dose of dexamethasone in selected patients who receive carboplatin may be more closely resemble to that in patients who receive cisplatin because carboplatin is a platinum agent. However, it remains unclear which patients who receive carboplatin-based MEC would benefit from an increased dose of dexamethasone equivalent to that used for HEC, given the potentially greater risk of adverse effects.

We hypothesised that women <70 years who did not drink alcohol were at a high risk for CINV, even after non-AC MEC. However, it is unknown whether 5-HT3 receptor antagonist and an increased dose of dexamethasone equivalent to that used for HEC adequately prevent CINV or whether additional treatment with aprepitant prevents CINV more effectively in these selected patients. We conducted a multicenter, placebo-controlled, double-blind, randomised phase II study to evaluate the effectiveness of aprepitant for preventing CINV after carboplatin- or irinotecan-based MEC in nondrinking women <70 years.

**Patients and methods**

### Study design

This trial was conducted between January 2011 and September 2012 after the approval from each site’s institutional review board. Written informed consent was obtained from all the patients who were enrolled with the use of an online registration system. The patients were stratified according to performance status (PS; 0 or 1–2), institution, and chemotherapy regimens (carboplatin or irinotecan), then randomly assigned to the aprepitant group or placebo group according to a computer-generated, blinded allocation schedule. The investigator, study nurses, and participants remained blinded to the treatment assignments. To ensure in-house blinding, the assigned treatment and intravenous dexamethasone were dispensed by a pharmacist who was not otherwise involved in the study. Matched placebos for oral aprepitant were used to maintain double blinding. Patients completed a diary to report vomiting episodes, rescue therapy use, and daily nausea assessments from the initiation of chemotherapy infusion (0 h) until the morning of day 6 (120 h) after chemotherapy. This study has been registered in the University Medical Information Network Clinical Trials Registry as No. 000004998.

### Patient population

Women aged 20–69 years with histologically confirmed malignancies who were naïve to aprepitant and scheduled to receive carboplatin- or irinotecan-based regimens were included in this study. Patients with a history of alcohol consumption, defined as ≥1 alcoholic drinks per week, were excluded. MEC regimens authorised for use in this study were as follows: (i) carboplatin (AUC 6 mg/min m−1) plus intravenous cytotoxic antitumor drugs, such as paclitaxel and pemetrexed; (ii) carboplatin (AUC 5 mg/min m−1) plus paclitaxel (200 mg m−2); (iii) carboplatin (AUC 5 mg/min m−1) plus liposomal doxorubicin (30 mg m−2); and (iv) irinotecan (≥150 mg m−2) plus fluorouracil, bevacizumab, or cetuximab. Eligible patients had to have an Eastern Cooperative Oncology Group PS of 0–2 and an estimated life expectancy of at least 3 months and had to meet the following laboratory criteria: neutrophil count ≥1500 mm−3; platelet count ≥100 000 mm−3; aspartate aminotransferase and alanine aminotransferase ≤2.5 times the upper limit of the normal range at the facility; total bilirubin ≤1.5 times the upper limit of the normal range at the facility; and creatinine ≤1.5 times the upper limit of the normal range at the facility. We also excluded patients at risk of vomiting for other reasons (symptomatic brain metastasis, meningeal infiltration, epilepsy, active peptic ulcers, gastrointestinal obstruction, concomitant abdominal, or pelvic radiotherapy), pregnant, nursing, or possibly pregnant women.

### Treatment

The doses of each drug according to the study group are shown in Table 1. On day 1, administration of the first MEC

| Table 1. Study of the drug schedule |
| Regimen | Day 1 | Day 2 | Day 3 |
|---|---|---|---|
| **Aprepitant group** | | | |
| Aprepitant | 125 mg orally | 80 mg orally | 80 mg orally |
| Granisetron | 1 mg IV | 4 mg IV | 4 mg IV |
| Dexamethasone | 12 mg IV | | |
| **Placebo group** | | | |
| Aprepitant | 0 mg orally | 0 mg orally | 0 mg orally |
| Granisetron | 1 mg IV | 8 mg IV | |
| Dexamethasone | 20 mg IV | | |

Abbreviation: IV = intravenous.
Aprepitant in selected patients receiving MEC

Statistical analysis. The primary end point was the rate of CR, defined as no vomiting or retching episodes with no rescue medication for 120 h from the start of the first cycle of MEC. The criteria were applied to evaluate CR in the acute (0–24 h), delayed (24–120 h), and overall (0–120 h) phases. The following key secondary end points were also analysed: (i) no emesis; (ii) no rescue therapy; (iii) no significant nausea (nausea score: none and mild); (iv) no nausea (nausea score: none and mild); and (v) total control (no emesis, no rescue therapy, and no nausea (nausea score: 0)).

This study was a randomised phase II trial comparing the aprepitant group with the placebo group. In the phase III trial of aprepitant in patients who received non-AC MEC (Rapoport et al, 2010), the CR rate in the placebo group was 65.5%. This rate was anticipated to be lower in nondrinking women <70 years receiving only carboplatin- or irinotecan-based regimens; however, the difference was expected to be offset by the additional doses of dexamethasone on days 2 and 3. Therefore, we estimated the CR rate in the placebo group to be 65% in our study. Treatment with aprepitant was expected to increase the CR rate by 20% on the basis of the results of phase III trials of aprepitant in patients who received HEC (Hesketh et al, 2003; Poli-Bigelli et al, 2003), given that CINV induced by MEC in these selected patients would be similar to CINV by HEC. Assuming a one-sided significance level of 10% for testing the primary hypothesis, the sample size was calculated based on the hypothesis that CR rate of the placebo group, estimated at 65%, would improve by 20% in the aprepitant group. Given a total sample size of 90 patients (45 per group), the statistical power was estimated to be 82%. Assuming that approximately 5% of the subjects would be withdrawn or drop out, the target sample size was set at 94 in total and 47 per group.

Exploratory subgroup analysis of predictive factors of CR in the overall phase was performed by logistic regression analysis. Variables with P values of ≤0.10 on univariate analysis and clinically important variables (age, PS, allocation) were included in the multivariate analysis. All statistical analyses were performed using the IBM SPSS Statistics 20 (IBM, Armonk, NY, USA).

RESULTS

Patients. A total of 94 patients were enrolled in this study and randomly assigned to one of the two treatment arms (Figure 1). Of these, 91 patients were included in the full analysis set. Both treatment groups had similar baseline demographics (Table 2). Most patients (98%) underwent carboplatin-based chemotherapy. Common malignancies were ovarian/peritoneal cancer (55%) and uterine endometrial cancer (38%). Thirty-nine (43%) patients were 60–69 years old.

Efficacy. The percentages of patients with CR in the overall, acute, and delayed phases for each treatment are shown in Figure 2. The CR rate in the overall phase was superior but not significantly higher in the aprepitant group than in the placebo group (aprepitant group, 62.2% (28 out of 45); placebo group, 52.1% (24 out of 46); P = 0.33). The difference was 10.1 percentage points (90% confidence interval, −7% to 27%). The acute phase efficacies were similarly high in both the groups (aprepitant group, 97.8% (44 out of 45); placebo group, 95.7% (24 out of 46)) and the delayed phase efficacy was the same as the overall efficacy. For each predefined, secondary end point and treatment, the overall phase efficacies are shown in Table 3. The median day of the first episode of vomiting or of rescue use was day 4 in both the groups. Eighteen (20%) patients used rescue therapy in the absence of clinically significant nausea or vomiting. The CR rates in the 14 patients with...
a previous history of chemotherapy were similar in the treatment groups (aprepitant group, 78% (7 out of 9); placebo group, 80% (4 out of 5)). The results of univariate logistic regression analysis of factors related to CR are shown in Table 4. Pretreatment plasma levels of estradiol or progesterone did not significantly correlate with the overall CR. Variables tested in the multivariate analysis were age, PS, history of chemotherapy, treatment allocation (aprepitant or placebo), ascites/peritoneal dissemination, and advanced (stage III/IV, recurrence) ovarian/primary peritoneal cancer. After controlling for these factors, the only independent predictor of a lower CR was advanced ovarian/primary peritoneal cancer (OR, 0.26 (0.10–0.72); P = 0.010; Table 5). Among the 39 patients with advanced ovarian/primary peritoneal cancer, the overall CR was 45% (10 out of 22) in the aprepitant group and 29% (5 out of 17) in the placebo group, respectively.

Tolerability. Safety was evaluated in all the 92 subjects who were assigned to treatment, including the patient who discontinued chemotherapy due to a hypersensitivity reaction. Generally, the adverse event profile did not markedly differ between the groups, although myalgia/arthralgia was more common in the placebo group (Table 5). One patient in the placebo group experienced grade 2 upper gastrointestinal bleeding on day 2, which might have been caused by dexamethasone; however, most events were mild and self-limiting. No grade 3–4 adverse events were observed except for a grade 4 hypersensitivity reaction to paclitaxel (n = 1) in the aprepitant group. A slight decrease in the incidence of anorexia was noted in patients receiving aprepitant (74%) compared with patients receiving the placebo (85%).

To our knowledge, this study is the first randomised trial to evaluate aprepitant in ’select’ patients, that is, nondrinking female patients <70 years who mainly received carboplatin-based MEC. Nearly half (47.9%) of the patients experienced vomiting or used rescue medication after MEC despite treatment with granisetron and an increased dose of dexamethasone. This was much lower than our expected CR rate of 65% and partly due to the low CR rate of 29% in 17 patients with advanced ovarian cancer. Addition of aprepitant seemed to be effective even with an increased dose of dexamethasone; however, improvement is required in the delayed phase CR rate of 62.2%. The addition of aprepitant resulted in a 10.1% non-significant improvement in the overall CR rate, which was also lower than our initial expectation of 20%. We had thus overestimated the benefit of aprepitant for the prevention of non-AC MEC. This non-significant result might be attributed to the small sample size in our trial, and a 10.1% improvement might be statistically significant in a larger study group. In a study of 857 patients with breast cancer who received AC, however, the efficacy of aprepitant was only an 8.3% improvement of CR rate (aprepitant group, 50.8%; placebo group, 42.5%; Warr et al, 2005), and all guidelines recommend the standard use of aprepitant in patients who receive AC chemotherapy. We still believe that a 10.1% improvement is promising and that further confirmatory phase III trials of aprepitant in this population are warranted. The overall CR rates in the 14 patients with a previous history of chemotherapy were similar between the groups, suggesting previous chemotherapy did not influence the overall results of our study.

Among the 91 subjects included in the full analysis set, the proportion of patients without significant nausea was 80%, which was higher than the 59% of patients who did not use rescue therapy (Table 3). Indeed, 20% of the patients used rescue therapy in the absence of subjective significant nausea or vomiting. This finding
suggests that one of the reasons for rescue use may be anxiety, which is the major cause of anticipatory nausea in cancer chemotherapy (Nerenz et al., 1986). Prophylactic benzodiazepines such as lorazepam or alprazolam may improve the delayed phase CR for such patients.

The use of palonosetron instead of granisetron might improve delayed CINV in this setting because the MASCC (Roila et al., 2010) and ASCO guidelines (Basch et al., 2012) recommend palonosetron as the preferred 5-HT3 receptor antagonist for non-AC MEC regimens. However, no clinically relevant differences between palonosetron and other 5-HT3 receptor antagonists have been demonstrated by randomised phase III trials for non-AC MEC regimens. Moreover, a recent study from the Rochester Cancer Center, New York demonstrated that delayed nausea was significantly improved by additionally giving dexamethasone on days 2 and 3; however, no difference was evident between palonosetron and granisetron after HEC or MEC (Roscoe et al., 2011). In patients who receive aprepitant, the difference between palonosetron and granisetron would be expected to be small. Other methods with the potential to improve delayed CINV are increased doses of dexamethasone on days 2 and 3, addition of dexamethasone, aprepitant, or both on days 4 and 5, addition of prochlorperazine or an 5-HT3 receptor antagonist on days 2–5, or addition of olanzapine on days 1–4 (Navari et al., 2011).

Table 3. Results of key end points in the overall phase

| End points                  | Aprepitant (n = 45) | Placebo (n = 46) | Diffence (90% CI) |
|-----------------------------|---------------------|-----------------|------------------|
| Complete response           | 28 (62%)            | 24 (52%)        | 0.33 10% (–7%, 27%) |
| No vomiting                 | 38 (83%)            | 36 (78%)        | 0.45 6% (–7%, 20%)  |
| No significant nausea a     | 38 (83%)            | 35 (76%)        | 0.32 8% (–5%, 22%)  |
| No rescue therapy           | 30 (67%)            | 24 (52%)        | 0.16 14% (–2%, 31%) |
| No nausea                   | 24 (53%)            | 18 (39%)        | 0.17 14% (–3%, 31%) |
| Total control b             | 21 (47%)            | 17 (37%)        | 0.35 10% (–7%, 27%) |

Abbreviation: CI = confidence interval. a No significant nausea; nausea score, none and mild. b Total control: no emesis, no rescue therapy, and no nausea; nausea score, 0.

Table 4. Logistic regression analysis of predictors of CR in the overall phase

| Age, years | n = 91 | Univariate P-value | Multivariate P-value | Odds ratio (95% CI) |
|------------|--------|--------------------|----------------------|--------------------|
| 60–69      | 39     | 0.90               | 1.00                 | 1.00 (0.39–2.55)   |
| 30–59      | 52     | 0.68               | 0.87                 | 0.86 (0.32–2.28)   |

Table 5. Patients with specific clinical adverse events of incidence ≥5% in at least one treatment group

| Aprepitant (n = 46) | Placebo (n = 46) |
|---------------------|------------------|
| Adverse events      | Grade 1 | Grade 2 | Grade 1 | Grade 2 |
| Anorexia            | 21      | 13     | 25      | 14      |
| Myalgia/arthritis   | 2       | 1      | 7       | 2       |
| Fatigue             | 4       | 3      | 4       | 2       |
| Constipation        | 2       | 1      | 1       | 1       |
| Diarrhea            | 3       | 1      | 2       | 1       |

No grade 3–4 adverse events were observed except grade 4 hypersensitivity (n = 1) in the aprepitant group.
and nearly half had undergone bilateral oophorectomy. Exploratory analysis showed that only advanced ovarian/peritoneal cancer was a significantly poor predictor of CINV, although the results of analysis might have been different or other variables might also have been found to be significant predictors of CR if the sample size had been larger. Among the 39 patients with advanced ovarian/peritoneal cancer, 86% (19 out of 22) in the aprepitant group and 88% (15 out of 17) in the placebo group had ascites or peritoneal dissemination. Therefore, the presence of ascites or peritoneal dissemination might have influenced CINV in patients with advanced ovarian/peritoneal cancer. Study groups including advanced ovarian/peritoneal cancer patients should be stratified according to cancer type in future antiemetic trials. Furthermore, aprepitant was associated with a 16% improvement in the overall CR in patients with advanced ovarian/peritoneal cancer (45% (10 out of 22) vs 29% (5 out of 17)); therefore, aprepitant might be more effective than placebo even in such patients, and additional studies are required to optimise treatment for this important subset of patients.

In conclusion, aprepitant in combination with granisetron and an increased dose of dexamethasone equivalent to that used for HEC was well tolerated and seemed more effective than placebo for the prevention of CINV in nondrinking women <70 years who received MEC. However, delayed-phase CINV remains a significant problem. Further confirmatory trials of aprepitant in this population are warranted.

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CONFLICT OF INTEREST

HM reported having accepted an unrestricted research grant and received honoraria from Ono Pharmaceutical Co., Ltd. The other authors declare no conflict of interest.

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