Prevention of chemotherapy-induced nausea and vomiting: focus on fosaprepitant

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Abstract: Fosaprepitant is a prodrug of aprepitant, a neurokinin₁ (NK₁) receptor antagonist used in prophylactic antiemetic regimens used prior to cytotoxic chemotherapy. Fosaprepitant is being developed to provide a parenterally administered alternative to the orally administered aprepitant. Fosaprepitant is rapidly converted to aprepitant and an intravenous dose of 115 mg is bioequivalent to 125 mg orally, with similar plasma concentrations at 24 hours. In phase I and II trials fosaprepitant shows efficacy, but the large randomized efficacy studies have utilized aprepitant. When it is added to dexamethasone and a 5-HT₃ receptor antagonist on day 1 prior to chemotherapy aprepitant improves the control of acute post chemotherapy emesis and when continued on days 2 and 3 with dexamethasone it demonstrated even greater improvement in the control of delayed emesis. This has been shown with both cisplatin-containing regimens and those based upon cyclophosphamide and an anthracycline. Fosaprepitant is well tolerated with mild to moderate venous irritation being the only additional toxicity to those seen with oral aprepitant, and that is a function of dose, concentration, and infusion rate. Headaches are the other toxicity most commonly reported. Fosaprepitant can be used as a parenteral alternative to aprepitant in regimens to control chemotherapy-induced emesis.

Keywords: fosaprepitant, aprepitant, neurokinin₁ receptor, emesis, chemotherapy

Introduction
The introduction of cytotoxic chemotherapy, and particularly cisplatin, was associated with nausea and vomiting that did not respond to conventional doses of the then available antiemetcs, exemplified by metoclopramide and prochlorperazine. Of the factors that predicted nausea and vomiting the most significant was the cytotoxic drug, as each drug displayed a different emetic potential, and there were different mechanisms by which the cytotoxics could cause vomiting (Andrews et al 1998; Hesketh 1999). The most common pattern of nausea and vomiting is acute emesis which commences within hours of receiving chemotherapy and lasts over the first 24 hours. This can be followed by delayed emesis commencing near the beginning of the first day and often lasting for at least 5 days (Kris et al 1985). Those patients who experience severe post-chemotherapy emesis are then prone to developing anticipatory emesis as a conditioned response (Morrow 1982). Patients who have poor control of post chemotherapy emesis also demonstrate deterioration in their quality of life (Osoba et al 1997).

It was the discovery that two important mechanisms for post-chemotherapy emesis were mediated through 5 hydroxytryptamine, (5HT₁) and neurokinin₁ (NK₁) receptors and the development of antagonists that saw a great impact made upon the control of chemotherapy-induced vomiting and to a lesser extent nausea. The 5HT₁ receptors, predominantly in the small bowel, were major mediators of acute emesis, and the first of the antagonists, ondansetron, when given prior to chemotherapy, revolutionized the control of post chemotherapy acute emesis. Ondansetron and dexamethasone controlled acute emesis in over 80% patients (Gralla et al 1999). Patients, however, were still
listing nausea and vomiting in their top three side effects even after the great improvement in the control of acute emesis (Boer-Dennert et al 1997). This was due to the incidence of delayed emesis which can occur in 20%–25% patients in the absence of acute emesis, and which was being underestimated by clinicians by up to 30% (Grunberg et al 2004).

Unfortunately, in only 50% patients was the delayed phase of emesis, caused by drugs such as cisplatin, controlled by ondansetron and dexamethasone. It was the dexamethasone that was the most active drug, suggesting that a different mechanism was responsible for delayed compared with acute emesis (Olver et al 1996).

Substance P, a tachykinin, binds to \( \text{NK}_1 \) receptors in the brain stem which send messages to the vomiting center and induce vomiting. Blocking the \( \text{NK}_1 \) receptor lessens vomiting after cisplatin, and a variety of other emetic stimuli (Diemunsch and Grelot 2000).

With the development of the orally active \( \text{NK}_1 \) receptor antagonist, aprepitant, it was found that when it was added to ondansetron and dexamethasone it improved the control of cisplatin-induced acute emesis, but when continued for 2 further days had a major impact on the control of the delayed phase of the post-chemotherapy emesis (Hesketh et al 2003; Poli-Bigelli et al 2003).

Fosaprepitant \((L-758,298\text{ or } MK-0517)\) is a prodrug of aprepitant that can be administered intravenously and is converted into aprepitant within 30 minutes (Navari 2007).

**The pharmacology of fosaprepitant**

Fosaprepitant dimeglumine is a white powder which is freely water soluble and is a phosphoryl prodrug of aprepitant (Hale et al 2000). Its antiemetic properties are attributable to aprepitant, which is a selective neurokinin 1 (\( \text{NK}1 \)) receptor antagonist with low affinity for \( \text{NK}2 \) and \( \text{NK}3 \) receptors (Watson et al 1998). It inhibits chemotherapy emesis by penetrating the brain and occupying central \( \text{NK}1 \) receptors for a sufficient duration to inhibit both the acute and delayed phases of emesis (Tattersall et al 2000). Antiemetic efficacy with aprepitant increases with receptor occupancy up until a dose of 125 mg orally, but there is no greater benefit with higher doses (Hargreaves 2002).

Fosaprepitant 115 mg given intravenously is bioequivalent to aprepitant 125 mg given by mouth with similar plasma concentrations at 24 hours (Merck and Co Inc 2007). It has been trialed in single daily doses for up to 4 days. Fosaprepitant is converted to aprepitant within 30 minutes after the end of an infusion. Aprepitant is 95% bound to plasma proteins. In vitro studies show that aprepitant is metabolized in the liver primarily by CYP3A4, with minor metabolism by CYP1A2 and CYP2C19.

Preclinical toxicology studies of bolus fosaprepitant administered in seconds to dogs and rats showed that concentrations of \( \leq 1\text{ mg/mL} \) were well tolerated. Concentrations up to 25 mg/mL at low doses \((2–4\text{ mg/kg/day})\) were well tolerated in dogs but intermediate concentrations \((10\text{ mg/mL})\) given at higher doses \((32\text{ mg/kg/day})\) caused venous irritation (Lasseter et al 2007).

There have been seven metabolites identified in human plasma, which are only mildly active. Following a single intravenous dose of 14C-labeled fosaprepitant 57% of the radioactivity was recovered in the urine and 45% in the feces. The terminal half-life of aprepitant following the administration of aprepitant is around 14 hours (Merck and Co Inc 2007).

No dose adjustment is considered necessary based on gender, race, or age although there are no data available for fosaprepitant in patients less that 18 years old. Based on pharmacokinetics no dose adjustment is required for severe renal insufficiency including those patients on dialysis (Bergman et al 2005). Similarly there is no clinically meaningful difference in the pharmacokinetic parameters patients with mild to moderated hepatic insufficiency (Child-Pugh score 7–9) and no data on severe hepatic insufficiency.

**Drug interactions**

As a moderate inhibitor of CYP3A4, fosaprepitant/aprepitant should not be co-administered with drugs such as pimozide, terfenadine, astemizole, or cisapride where the inhibition may result in elevated concentrations of these drugs, with serious consequences. Similar care should be taken with drugs with a narrow therapeutic index that are metabolized by CYP3A4 such as cyclosporine, sirolimus, and tacrolimus. As fosaprepitant/aprepitant induces the metabolism of drugs metabolized by CYP2C9, the co-administration with drugs metabolized by this mechanism such as warfarin should be monitored carefully (Depre et al 2005). This is also the case for tolbutamide and phenytoin, which may result in them achieving lower plasma concentrations than desirable. The concentrations of oral contraceptives may be decreased with co-administration of aprepitant, so alternate contraception should be used.

Of the drugs to be given with fosaprepitant, there is no evidence of clinically meaningful interactions with \( \text{SHT}_3 \) antagonists, including palonosetron (Blum et al 2003; Shah et al 2005). There are a number of chemotherapy agents metabolized by CYP3A4 such as taxanes, etoposide,
irinotecan, ifosfamide, imatinib, and vinca alkaloids. Oral aprepitant did not influence the pharmacokinetics of docetaxel or vinorelbine and there has been no obvious interaction clinically when administered with etoposide, vinorelbine and paclitaxel (Nguyen et al 2005; Loos et al 2007). Aprepitant does inhibit cyclophosphamide bioactivation and thiopeta metabolism, but the effects are small (de Jonge et al 2005).

Aprepitant increases the AUC of dexamethasone, a substrate of CYP3A4 2.2 fold; therefore the co-administered doses of dexamethasone should be halved, as was done in the two phase III trials that established the efficacy of aprepitant in preventing high-dose cisplatin-induced emesis (McCrea et al 2003).

Drugs such as rifampicin and St John’s Wort, strong CYP3A4 inducers, reduce the concentration of aprepitant/fosaprepitant whereas a strong CYP3A4 inhibitor like ketoconazole can increase the AUC of aprepitant five-fold. Food effects on the absorption of aprepitant are minimized because of its nanoparticle formulation (Olver et al 2007).

**Early phase clinical trials of fosaprepitant**

In the initial development of aprepitant, two of the phase II trials used the intravenous prodrug L-758298, now called fosaprepitant. These trials were conducted in patients who were receiving their first ever dose of at least 50 mg/m² cis-fosaprepitant. These trials used the intravenous prodrug L-758298, now called fosaprepitant whereas a strong CYP3A4 inhibitor like ketoconazole can increase the AUC of aprepitant five-fold. Food effects on the absorption of aprepitant are minimized because of its nanoparticle formulation (Olver et al 2007).

In the first of these fosaprepitant at a dose of either 60 mg or 100 mg was compared with ondansetron 32 mg given 1 hour before the chemotherapy. There was no significant difference in acute postchemotherapy emesis, with 37% on fosaprepitant and 52% on ondansetron having no emesis. In the delayed phase, 72% receiving fosaprepitant had no emesis compared with 30% receiving ondansetron (McCrea et al 2001).

In the next study, fosaprepitant was studied in combinations. One group of patients received fosaprepitant 100 mg with intravenous dexamethasone 20 mg on day 1 and then oral aprepitant (MK-869) 300 mg on days 2 to 5. The second group had the same drugs on day 1 and placebo on days 2 to 5 and the third group received intravenous ondansetron 32 mg added to dexamethasone on day 1 and placebo on days 2 to 5. Acute emesis was best controlled by the ondansetron and dexamethasone group (83%) compared with only 40% when the fosaprepitant groups were combined. However, the fosaprepitant groups did better in the delayed phase of the emesis with complete responses recorded for group 1 in 59%, for group 2 in 46%, and for group 3 in 38% (p < 0.05 group 1 vs group 3) (van Belle et al 2002). These trials suggested that NK₁ receptor antagonists were going to be best in combination antiemetic regimens.

Later,fosaprepitant was developed as an alternative intravenous formulation to oral aprepitant. A study was performed to find the bioequivalent dose of fosaprepitant, in a polysorbate 80 vehicle, in terms of aprepitant AUC (area under the concentration time curve) to 125 mg aprepitant (Lasseter et al 2007). The study was in 3 parts. In parts 1 and 2 fosaprepitant doses from 90 to 150 mg were investigated, and based on those results 2 doses of fosaprepitant, 100 and 115 mg, were selected for the randomized open label crossover test of bioequivalence in part 3. Blood samples were collected over 72 hours following the drug administration for aprepitant assays. Patients ranged in age from 18 to 45 years. Across all three parts of the study 106 subjects received fosaprepitant doses ranging from 90 to 150. There were no subjects discontinued because of adverse events. In parts 1 and 2 neither the 150 mg dose nor the 90 mg dose met the AUC bioequivalence criteria relative to 125 mg aprepitant. The 115 mg dose proved bioequivalent to aprepitant. It is rapidly converted to aprepitant with a half-life in the plasma of 2.3 minutes and complete conversion occurs within 30 minutes. It is not extensively distributed to the tissues. Although fosaprepitant resulted in a higher maximum concentration than aprepitant, the trough concentrations at 24 hours were equivalent, suggesting similar NK₁ receptor occupancy. This suggests that fosaprepitant 115 mg can be used interchangeably with 125 mg aprepitant.

**Efficacy**

The efficacy data come from the trials of aprepitant. Initially there were 3 other phase II studies with aprepitant. A 351-patient placebo-controlled study in cisplatin-induced emesis studied 4 groups: the first used granisetron and dexamethasone on day 1 with placebo days 2 to 5; the second added in aprepitant on day 1 and then 2 to 5; the third arm used aprepitant the day before chemotherapy and then with dexamethasone on day 1 but without the granisetron, and then continued aprepitant days 1 to 5; and the final arm was similar but with the aprepitant and dexamethasone starting on day 1 (Campos et al 2001). This trial confirmed the need for a 5HT₃ antagonist as part of triple therapy on day 1 for controlling the acute phase of emesis and confirmed the benefit of aprepitant in delayed emesis.

In a further phase II study of 159 patients, granisetron and dexamethasone was the standard treatment with other groups
adding aprepitant to this on day 1, then either continuing aprepitant until day 5 or using placebo form days 2 to 5 (Navari et al 1999). There was a significantly superior result for acute emesis in the two groups who received triple therapy on day 1 (93% and 94% for complete control of emesis vs 67% p < 0.001) and in the delayed phase those groups receiving aprepitant were significantly better (82% and 78% vs 33%, p < 0.001) although the difference between just day 1 aprepitant and daily dosing for 5 days did not reach statistical significance.

A further phase II was designed to refine the dosing of aprepitant, adding to ondansetron plus dexamethasone 125 mg aprepitant on day 1 and 80 mg days 2 to 5 compared with 40 mg on day 1 then 25 mg on days 2 to 5 or placebo (Chawala et al 2003). Again the patients receiving aprepitant had superior control of acute and delayed emesis, but nausea was only better controlled in the delayed phase. The efficacy of aprepitant was proven definitively in 2 phase II trials in patients receiving their first ever dose of cisplatin ≥70 mg/m² over ≤3 hours. The standard therapy was considered as ondansetron and dexamethasone on day 1 followed by dexamethasone on days 2 to 4. The aprepitant arms added 125 mg of oral aprepitant on day 1 and then 80 mg days 2 and 3 giving just the dexamethasone on day 4 (Hesketh et al 2003; Poli-Bigelli et al 2003). Recording the Poli-Bigelli study first then the Hesketh study, the aprepitant arms were statistically significantly superior in overall complete response rates (62.7% vs 43.3%, p < 0.001 and 72.1% vs 52.3%, p < 0.001) and for acute emesis (82.8% vs 68.4%, p < 0.001 and 89.2% vs 78.1%, p < 0.001). However, the most impressive response differences were in delayed emesis (67.7% vs 46.8%, p < 0.001 and 74.4% vs 55.8%, p < 0.001). Again, nausea was better controlled only in the delayed phase. The efficacy of aprepitant was maintained over 6 cycles.

A specific study of multiple cycles of high-dose cisplatin used ondansetron and dexamethasone for the acute phase and compared a placebo for the delayed phase with two different dosing schedules of aprepitant on days 1 to 5 (de Wit et al 2003). The complete response rate was maintained over 6 cycles in the aprepitant groups but not in the placebo group.

In randomized studies, the selection of the control arm is vital. Given that it had been common practice to continue the 5HT₁ antagonist with the dexamethasone for 4 days, a study in 489 patients receiving high-dose cisplatin randomized them to either ondansetron and dexamethasone for on each day or triple therapy with aprepitant on day 1 followed by aprepitant and dexamethasone as given in the large randomized trials (Schmoll et al 2006). The aprepitant regimen yielded a significantly improved 5-day overall response rate (72% vs 61%, p = 0.003) with a 9% improvement in protection from nausea and vomiting on day 1 and 11% on days 2 to 5.

Following the studies with cisplatin-containing regimens, the efficacy of adding aprepitant to other chemotherapy regimens was trialed. In a large study 866 patients who were treated with cyclophosphamide and anthracyclines, which as single agents have moderated emetic potential but as a combination arguably are of high emetic potential, were studied (Warr et al 2005). Patients were given oral ondansetron and dexamethasone with either aprepitant on days 1, 2, and 3 or just followed by 2 days of ondansetron. The overall 5-day complete response rate was (50.8% vs 42.5%, p = 0.015) in favor of the aprepitant group. Nausea was not as well controlled as vomiting particularly in the delayed phase. The improved efficacy was maintained over 4 cycles of chemotherapy (Herrstedt et al 2005).

In a small trial, the role of aprepitant as a salvage therapy to add into the antiemetic regimen in subsequent cycles if patients had failed to respond to a 5HT₁ receptor antagonist and dexamethasone was tested for both cisplatin and other chemotherapy (Oechsle et al 2006). The addition of aprepitant significantly improved activity.

There are no large studies of the efficacy of aprepitant in children or adolescents but case reports in adolescents are promising (Smith et al 2005).

As other new antiemetics are developed, combinations with aprepitant are tested. Palonosetron is a 5HT₁ receptor antagonist with a longer duration of activity which has been reported as having better activity against the delayed emesis associated with chemotherapy of moderate emetic potential. After determining that there were no pharmacokinetic problems with the combination, aprepitant with palonosetron and dexamethasone was found to be highly effective in preventing chemotherapy-induced nausea and vomiting in the days following the administration of cyclophosphamide chemotherapy of moderate emetic potential (Shah et al 2005; Grote et al 2006).

**Safety and tolerability**

Fosaprepitant and aprepitant are very well tolerated. In the initial phase 1 studies with aprepitant or its prodrug the adverse events that were reported more commonly for aprepitant groups over placebo were fatigue, somnolence, dizziness,
flushing, nausea, hiccups, headache, and menstrual problems. In the trials of L-758298 compared with ondansetron the only significant difference in toxicities was more diarrhoea with the prodrug than ondansetron, which itself would tend toward producing constipation (Cocquyt et al 2001; van Belle et al 2002). Most studies administered aprepitant as part of an antiemetic regimen which often included 5HT3 receptor antagonists, so it has been difficult to isolate the side effects specifically due to aprepitant. In the two large phase III trials the incidence of drug-related side adverse events for the aprepitant arm was 19.5% vs 14.4% in the control arm in the Poli-Bigelli study and 14.6% vs 11.0% in the Hesketh study (Hesketh et al 2003; Poli-Bigelli et al 2003). The most reported side effects were asthenia, anorexia, diarrhea, headaches, and hiccups. In the Warr study there was more constipation in the control arm and more dyspepsia in the aprepitant arm (Warr et al 2005).

In the study to establish the bioequivalence of fosaprepitant there were no serious adverse events or laboratory toxicities (Lasseter et al 2007). Headache and infusion site symptoms were the most reported events, but these were only of mild to moderate intensity. Fosaprepitant is tolerable at 1 mg/mL infused over 15 to 30 minutes but was found to cause irritation at 25 mg/mL at doses of 50 mg or 100 mg infused over 30 seconds.

**Patient quality of life and satisfaction**

The impact on patients of chemotherapy-induced nausea and vomiting goes beyond those side effects to be associated with a measurable deterioration in their global quality of life (Osoba et al 1997). More specifically the patients experiencing emesis also have more fatigue, anorexia, and insomnia.

In the large phase III trials of aprepitant in combination regimens for cisplatin and non-cisplatin induced emesis, a quality of life scale, the Functional Living Index Emesis (FLIE), was used (Martin et al 2003). The arms of the studies that contained aprepitant reported minimal or no impact of the emesis on daily life compared with those just receiving a 5HT3 antagonist and dexamethasone (74.4% vs 63.5% in the Poli-Bigelli study and 70.4% vs 55.6% in the Hesketh study) (Hesketh et al 2003; Poli-Bigelli et al 2003). In subgroup analyses this improvement in maintaining daily life activities was independent of sex and independent of age (Ma et al 2003; Martin et al 2003; Hesketh et al 2006). A similar outcome was reported in the Warr study where cyclophosphamide and an anthracycline was the chemotherapy, and those reporting minimal or no impact of emesis and quality of life were 63.5% on aprepitant vs 55.6% without, p = 0.019).

Cost effectiveness is also an important consideration for patients. It has been calculated that the routine use of aprepitant in antiemetic prophylaxis is cost effective in those situations where the likelihood of delayed emesis is high or the need will exist later for costly rescue medications (Moore et al 2007).

**Conclusions**

Fosaprepitant, a prodrug of aprepitant, is being developed as an intravenous alternative to oral aprepitant. It is rapidly converted to aprepitant after administration. The efficacy is the same as that of aprepitant. It is very well tolerated, with venous irritation being specific to this formulation and headache being the most frequent of the other toxicities. It seems that it is able to be used interchangeably with aprepitant and has been safely administered in single- and multiple-day schedules.

For example, in triple antiemetic therapy, fosaprepitant could be administered on day 1 with other antiemetics before intravenous chemotherapy and then on days 2 and 3 the NK1 therapy could be continued with oral aprepitant. An intravenous formulation may be more convenient in some circumstances and in some jurisdictions there are differing reimbursements for intravenous and oral drugs which may make an intravenous formulation more desirable, particularly if the income was being used to fund additional patient services.

Medical indications would include patients with severe mucositis or difficulty swallowing or any gastrointestinal disturbance that would make oral therapy problematic.

Fosaprepitant provides an intravenous alternative to oral formulations of NK1 receptor antagonists that is safe and effective.

**References**


