REVIEW

Evolving role of neurokinin I-receptor antagonists for chemotherapy-induced nausea and vomiting

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Abstract: To examine pharmacologic and clinical characteristics of neurokinin 1 (NK₁)-receptor antagonists (RAs) for preventing chemotherapy-induced nausea and vomiting (CINV) following highly or moderately emetogenic chemotherapy, a literature search was performed for clinical studies in patients at risk of CINV with any approved NK₁ RAs in the title or abstract: aprepitant (capsules or oral suspension), HTX019 (intravenous [IV] aprepitant), fosaprepitant (IV aprepitant prodrug), rolapitant (tablets or IV), and fixed-dose tablets combining netupitant or fosnetupitant (IV netupitant prodrug) with the 5-hydroxytryptamine type 3 (5HT₃) RA palonosetron (oral or IV). All NK₁ RAs are effective, but exhibit important differences in efficacy against acute and delayed CINV. The magnitude of benefit of NK₁-RA-containing three-drug vs twodrug regimens is greater for delayed vs acute CINV. Oral rolapitant has the longest half-life of available NK₁ RAs, but as a consequence should not be administered more frequently than every 2 weeks. In general, NK₁ RAs are well tolerated; however, IV rolapitant was recently removed from US distribution, due to hypersensitivity and anaphylaxis, and IV fosaprepitant is associated with infusion-site reactions and hypersensitivity presumed related to its polysorbate 80 excipient. Also, available NK, RAs have potential drug–drug interactions. Adding an NK,

RA to $5HT_3$ RA and dexamethasone significantly improves CINV control vs the two-drug regimen. Newer NK₁ RAs offer more formulation options, higher acute-phase plasma levels, or improved tolerability, and increase clinicians' opportunities to maximize benefits of this important class of antiemetics.

Keywords: aprepitant, chemotherapy-induced nausea and vomiting, fosaprepitant, netupitant, neurokinin 1-receptor antagonists, rolapitant

Plain-language summary

This review aims to evaluate the unmet need for superior control of a common side effect of chemotherapy, known as chemotherapy-induced nausea and vomiting (CINV). Prevention of CINV maintains the patient's quality of life and minimizes CINV-related hospital visits. Several guidelines exist that recommend specific drug regimens for CINV treatment. One class of drugs recommended to prevent CINV, known as neurokinin 1-receptor antagonists (NK₁ RAs), is underused in clinical practice. Several NK₁ RAs are available, which have pharmacologic and clinical differences including formulation (intravenous vs oral), efficacy, and safety profiles. These differences should guide a physician's choice of treatment for each patient. An NK₁ RA can be added to an antiemetic regimen, a combination of drugs for preventing nausea and vomiting that includes a 5-hydroxytryptamine type 3 RA and corticosteroid. This regimen can significantly reduce episodes of vomiting and the need for additional medications. However, nausea control remains suboptimal, and further research is needed to find better antiemetic regimens to prevent vomiting and nausea successfully, specifically CINV. Some of the newer, improved NK₁ RAs can add maximum benefit to the antiemetic-drug regimen.

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Introduction

Nausea and vomiting (NV) are common, distressing adverse effects of chemotherapy.^{1,2} Chemotherapy-induced NV (CINV) significantly affects patients' daily functioning,²⁻⁴ quality of life,^{1,5-8} and ability to eat.^{2,6} Patients with uncontrolled CINV require more health care resources and incur greater health care costs.^{3,8-10} Poorly controlled or severe CINV can prompt a chemotherapy dose reduction or cycle delay,¹¹ ultimately affecting chemotherapy outcomes.

CINV incidence depends on several factors, including female sex,¹² young age (<50 years),^{13,14} and anxiety,¹⁵ but the key determinant is the chemotherapy regimen's emetogenicity.¹⁶ Antiemetic guidelines classify chemotherapeutic agents as having high, moderate, low, or minimal risk of inducing CINV.^{16–19} Without effective prophylaxis, highly emetogenic chemotherapy (HEC) induces vomiting in >90% of patients who receive it, and moderately emetogenic chemotherapy (MEC) induces vomiting in 30%–90% of recipients.¹⁶ CINV has a relapsing–remitting–relapsing time course. Patients usually experience intense CINV within 1–2 hours of initiating chemotherapy, lasting for about 24 hours (acute phase). Symptoms usually recede, but reemerge at 48–72 hours (delayed phase).²⁰

Guidelines for CINV prophylaxis have been developed by the National Comprehensive Cancer Network (NCCN),¹⁶ American Society of Clinical Oncology (ASCO),¹⁷ and Multinational Association of Supportive Care in Cancer (MASCC) and European Society of Medical Oncology.^{18,19} These include recommendations for preventing acute and delayed CINV tailored to the emetogenicity of the chemotherapy regimen.¹⁶⁻¹⁹ For most patients receiving HEC or MEC, a three- or four-drug regimen is recommended to prevent acute CINV.¹⁶⁻¹⁹ The standard three-drug regimen consists of a combination of a 5-hydroxytryptamine type 3 (5HT₂)-receptor antagonist (RA), a neurokinin 1 (NK₂) RA, and dexamethasone, 16-19 with olanzapine added for four-drug regimens recommended by ASCO and NCCN for patients receiving HEC.16,17 The MASCC guidelines recommend a three-drug regimen of a 5HT, RA and dexamethasone with either an NK, RA or olanzapine (if nausea is an issue).¹⁸ NCCN guidelines offer an alternative three-drug regimen for HEC or MEC: olanzapine, palonosetron, and dexamethasone.16 Patients receiving HEC or MEC should also receive antiemetics on chemotherapy days 2-4 to prevent delayed CINV, the choice of agent(s) depending on the antiemetic regimen received for acute CINV prophylaxis.¹⁶⁻¹⁹

Antiemetic prophylaxis aims for complete CINV prevention,²⁰ best achieved with multiple agents targeting

different emetogenic pathways.¹⁶ Unfortunately, many patients do not receive guideline-recommended antiemetic regimens,^{21–25} so are more likely to experience CINV.^{21,23–25} The reasons for poor adherence to CINV-guideline recommendations are unclear, but evidence suggests that physicians and patients perceive CINV differently.^{26,27} For example, physicians tend to underestimate the nausea that patients experience,²⁵ particularly during the delayed phase,²⁶ and prescribers, but not patients, often identify cost as a barrier to using effective antiemetic prophylaxis.²⁷

Despite comprehensive antiemetic guidelines, unmet medical needs remain in CINV management, especially for better nausea control (particularly delayed nausea). Moreover, use of certain drug classes, especially NK₁ RAs, is suboptimal,^{23,24} possibly reflecting a poor understanding of their appropriate use. This review aims to examine the pharmacologic, pharmacokinetic, and clinical features of NK₁ RAs and how they affect clinical efficacy and safety, enabling physicians to make informed, evidence-based, and rational therapeutic decisions about using these agents for CINV prophylaxis.

Overview of NK, RAs

CINV is mediated by a complex neural network in the gut and central nervous system, so combination antiemetic regimens are indicated to target multiple pathways. One pathway involves the action of substance P on NK₁ receptors in the gut and central nervous system. Chemotherapy induces substance P release in these regions during acute and delayed CINV, so blocking the NK₁ receptor may prevent acute and delayed emesis.²⁸ In addition, there is evidence of "cross talk" between the emetic pathways, such that a combination of a 5HT₃ RA and an NK₁ RA has synergistic antiemetic effects.²⁸

Several NK₁ RAs are available in the United States for use in combination with other antiemetics for CINV prevention. Aprepitant (Emend; Merck, Whitehouse Station, NJ), rolapitant (Varubi; Tesaro, Waltham, MA), and netupitant (Akynzeo; Helsinn Therapeutics, Iselin, NJ) are orally administered.^{29–31} Fosaprepitant (Emend IV; Merck) is a prodrug of aprepitant, permitting intravenous (IV) administration.³² In late 2017, IV formulations of aprepitant (HTX019, Cinvanti; Heron Therapeutics, San Diego, CA) free of polysorbate 80 and other synthetic surfactants³³ and rolapitant³¹ were approved in the United States. Most recently, in April 2018, fosnetupitant (Akynzeo), the prodrug of netupitant, was approved in the United States, allowing IV administration of netupitant.²⁹ Unlike the other NK₁ RAs, netupitant and fosnetupitant are available only in a fixed combination with the $5HT_3$ RA palonosetron (netupitant/palonosetron [NEPA] oral or IV).²⁹

Approved NK, **RAs** Formulations and indications

Orally administered NK₁ RAs are available as tablets, capsules, and oral suspension.²⁹⁻³¹ The aprepitant oral suspension can be used in almost any age-group, including in infants aged ≥ 6 months, while aprepitant capsules are only for patients aged ≥ 12 years.³⁰

Currently approved NK₁ RAs have similar but subtly different indications listed in their prescribing information. The approved formulations are summarized in Table 1 according to their brand name, route of administration, indication, and year of approval.^{29–33} IV aprepitant (Cinvanti) is a polysorbate 80- and synthetic surfactant-free formulation containing natural excipients.³³ Fosnetupitant (the prodrug of netupitant included in IV NEPA) was developed without the need for a surfactant emulsifier or solubility enhancer.^{29,34} The IV formulation of fosaprepitant contains polysorbate 80,³² and IV rolapitant (Varubi) contains the synthetic surfactant polyoxyl 15 hydroxystearate.³¹

Pharmacokinetic, receptor occupancy, and pharmacodynamic properties

Table 2 summarizes the pharmacokinetic characteristics of the NK₁ RAs currently approved in the United States and the agents' occupancy of NK₁ receptors in the brain. According to current US prescribing information, in healthy volunteers all of the oral formulations reach maximum plasma levels (C_{max}) in 3–5 hours.^{29–31,35} For IV formulations, C_{max} is reached within 30 minutes of the start of infusion.^{31–33} The elimination half-life ($t_{1/2}$) in healthy volunteers is 9–13 hours for aprepitant after oral or IV administration,^{30,32,33} but considerably longer for netupitant (oral 96 hours, IV 144 hours) and rolapitant (IV or oral 169–183 hours).^{29,31} The long $t_{1/2}$ of rolapitant explains why a single dose administered 1–2 hours before

Drug (brand name)	Administration route	Indication	Year of approval
Aprepitant (Emend) ³⁰	PO (capsules or suspension)	In adults (capsules or suspension) and pediatric patients (suspension, aged \geq 6 months; capsules, aged \geq 12 years), in combination with other antiemetics for: acute and delayed NV associated with initial and repeat courses of HEC, including high-dose cisplatin NV associated with initial and repeat courses of MEC (adults and pediatrics) in adults (capsules) for PONV	2003
Fosaprepitant (Emend) ³²	IV	In adults and pediatric patients aged ≥6 months, in combination with other antiemetics for: acute and delayed NV associated with initial and repeat courses of HEC, including high-dose cisplatin delayed NV associated with initial and repeat courses of MEC	2008
Netupitant/palonosetron capsule (Akynzeo) ²⁹	PO	In adults in combination with dexamethasone for: acute and delayed NV associated with initial and repeat courses of chemotherapy, including but not limited to HEC	US: 2014 EU: 2015
Rolapitant tablet (Varubi) ³¹	PO	In adults in combination with other antiemetic agents for: delayed NV associated with initial and repeat courses of emetogenic chemotherapy, including but not limited to HEC	US: 2015
Rolapitant IVª (Varubi) ³¹	IV	In adults in combination with other antiemetic agents for: delayed NV associated with initial and repeat courses of emetogenic chemotherapy, including but not limited to HEC	US: 2017
Aprepitant IV (Cinvanti) ³³	IV	In adults in combination with other antiemetics: acute and delayed NV associated with initial and repeat courses of HEC, including high-dose cisplatin NV associated with initial and repeat courses of MEC	US: 2017
Netupitant/palonosetron (Akynzeo) ²⁹	IV	In adults in combination with dexamethasone: acute and delayed NV associated with initial and repeat courses of HEC	US: 2018

 Table I Formulations of approved NK, RAs

Note: ^aManufacturer issued a press release on February 28, 2018 announcing the suspension of rolapitant IV distribution.⁸⁹

Abbreviations: HEC, highly emetogenic chemotherapy; IV, intravenous; MEC, moderately emetogenic chemotherapy; NK, RAs, neurokinin 1-receptor antagonists; PO, per os (oral).

Table 2 Comparative p	sharmacokinetic par	ameters and NK _I -re	Table 2 Comparative pharmacokinetic parameters and NK ₁ -receptor occupancy for approved NK ₁ -receptor antagonists in healthy volunteers	NK ₁ -receptor anta	gonists in healthy v	volunteers	
	Aprepitant PO (Emend) 125 mg ^{30,40,41}	Fosaprepitant IV (Emend) ^a I50 mg ^{32,40}	Netupitant/palonosetron PO (Akynzeo): ^b 300 mg netupitant ^{29,39}	Rolapitant PO (Varubi) 180 mg ^{31,35}	Rolapitant IV (Varubi) 166.5 mg ^{31,35}	Aprepitant IV (Cinvanti) I30 mg ^{33,85}	Fosnetupitant/palonosetron IV (Akynzeo): ^c 235 mg fosnetupitant ²⁹
Pharmacokinetic parameters	neters						
C _{max} (ng/mL)	1,600	4,200	434	968-992	1,986	6,100	841
t _{max} (hours)	4	<0.5	5	3-4	0.5	0.5	0.5
AUC (ng·h/mL)	19,600	37,400	14,402	118,252	124,016	45,460	12,012
$\mathbf{t}_{_{Y_{2}}}$ (hours)	9–13	9–13	96	169–183	169–183	9–13	144
Brain NK ₁ -receptor occupancy	upancy						
Tracer uptake (%)							
24 hours postdose	96	100	61-100	NA	NA	NA	NA
48 hours postdose	≥97 ^d	≥97	59–98	NA	NA	NA	NA
96 hours postdose	NA	NA	61–98	NA	NA	NA	NA
I 20 hours postdose	NA	40-75	NA	73	NA	NA	NA
Notes: Brain NK ₁ -receptor oc aprepitant after fosaprepitant a Abbreviations: AUC, area ur	ccupancy was estimated fi administration; ^b pharmacc ader the (concentration⊣	rom positron emission-tom okinetic parameters for net time) curve; C _{max} , maximal	Notes: Brain NK ₁ -receptor occupancy was estimated from positron emission-tomography scans after administration of the high-affinity NK ₁ -receptor ligand GR205171, radiolabeled with ¹¹ C, as a tracer. ¹⁹ Pharmacokinetic parameters for aprepitant after fossetupitant administration; ⁴ Preceptor occupancy achieved with aprepitant 165 mg. Abbreviations: ⁴ Preceptor occupancy achieved with aprepitant 165 mg. Abbreviations: AUC, area under the (concentration-time) curve: C _{mx} , maximal plasma concentration; IV, intravenous; NA, not available; NK ₁ , neurokinin 1; PO, per os (oral); t ₂₀ elimination half-life; t _{mx} , time to C _{mx} .	: high-affinity NK ₁ -recept ~ netupitant after fosnetu IA, not available; NK ₁ , ne	:or ligand GR205171, ra Jpitant administration; ⁴ eurokinin 1; PO, per os	diolabeled with ¹¹ C, as a receptor occupancy achi (oral); $t_{_{N}}$, elimination hal	racer. "Pharmacokinetic parameters for eved with aprepitant 165 mg. F-life; $t_{\rm max}$ time to $C_{\rm max}$.

chemotherapy is expected to prevent delayed CINV, as its protection against CINV is established for up to 5 days.^{36–38} However, rolapitant takes longer to achieve therapeutic concentrations, so may be less effective in the acute phase.³¹ In patients with cancer, systemic exposure to netupitant is lower than in healthy volunteers, but this has been reported to be clinically insignificant.²⁹

Plasma pharmacokinetic profiles of different NK₁ RAs suggest that all can rapidly bind to NK₁ receptors; however, there may be differences in the inherent ability of different NK₁ RAs to cross the blood–brain barrier.^{35,39–41} Receptor occupancy (RO) studies conducted in healthy volunteers have suggested that aprepitant reaches full RO within 24 hours.⁴¹ NK₁ RO by netupitant has been reported 24–96 hours postdose, and varies in different brain regions.³⁹ Brain NK₁ RO at 120 hours postdose has been reported at levels of 40%–75% for IV fosaprepitant⁴⁰ and 94% for oral rolapitant.⁴¹ Oral and IV rolapitant are highly plasma-bound (99.8%),³¹ which in conjunction with its longer t_{42} provides support for the high RO seen at 120 hours postdose, but RO data on earlier time points were not provided for either of the rolapitant formulations.³¹

Pharmacodynamics indicate that all NK, RAs undergo hepatic metabolism, with the potential to cause drug-drug interactions via the CYP enzyme system.²⁹⁻³³ Aprepitant and its prodrug fosaprepitant are substrates for CYP3A4. They can induce the enzyme and inhibit it weakly to moderately. Aprepitant and fosaprepitant also induce CYP2C9. As a result, concurrent use of the antipsychotic agent pimozide, a CYP3A4 substrate, is contraindicated with aprepitant or fosaprepitant, and there are warnings about using aprepitant (oral or IV) or fosaprepitant with other agents that are CYP3A4 substrates.^{30,32} Furthermore, using these agents with strong or moderate CYP3A4 inhibitors (eg, ketoconazole or diltiazem) may increase plasma concentrations of aprepitant, leading to an increased risk of drug-related adverse events. Conversely, the use of aprepitant or fosaprepitant formulations with strong CYP3A4 inducers (eg, rifampin) may reduce aprepitant plasma concentrations and decrease its efficacy.^{30,32,33} Because aprepitant and fosaprepitant induce CYP2C9, they can affect the clotting response to warfarin, so patients taking concomitant warfarin should have their international normalized ratio monitored.^{30,32,33} Aprepitant and fosaprepitant can also reduce the efficacy of oral contraceptives.

Rolapitant is also slowly metabolized by CYP3A4, but does not induce or inhibit this enzyme.⁴² However, rolapitant metabolism also involves CYP2D6, so rolapitant should be used with caution in combination with other substrates for this enzyme if they have a narrow therapeutic index.³¹ For this reason, thioridazine is contraindicated in patients receiving rolapitant, and pimozide should be avoided.³¹ Rolapitant also inhibits the efflux transporters P-glycoprotein and BCRP, so increases systemic exposure to agents that are substrates of these transporters, including digoxin and sulfasalazine.⁴²

NEPA has no specific contraindications, but prescribing information includes warnings about the potential for hypersensitivity reactions and serotonin syndrome. As listed in the prescribing information, limitations of use include the fact that IV NEPA has not been studied in patients receiving anthracycline plus cyclophosphamide (AC)-based HEC and thus lacks data on potential hypersensitivity reactions in this group of patients. Because netupitant inhibits CYP3A4, drugs that are substrates or inducers of this enzyme should be avoided when NEPA is prescribed.²⁹

Clinical efficacy: complete response

A PubMed literature search was undertaken for clinical studies in patients at risk of CINV from 2003 to 2018, in which any approved NK₁ RA appeared in the title or abstract of the publication. A similar search was performed for published abstracts presented at major supportive-care congresses, ie, MASCC, ASCO, ASCO Palliative and Supportive Care Conference, and European Society of Medical Oncology, from 2016 to 2018. The results identified studies in which a threedrug regimen of an NK₁ RA, a 5HT₃ RA, and dexamethasone was compared with a two-drug combination of a 5HT₃ RA and dexamethasone.

Efficacy with highly emetogenic chemotherapy

Randomized controlled trials in patients receiving HEC are summarized in Table 3.36-38,43-56 The primary end point in most studies was complete response (CR), ie, no episodes of vomiting and no rescue antiemetic therapy. Most studies showed a significantly greater CR rate over the 5-day assessment period in groups receiving triple therapy compared with dual therapy.^{36–38,43,46–48,51,53–55} The exceptions were a study of fosaprepitant in women with gynecologic cancers undergoing combined radiotherapy and chemotherapy⁵² and two Japanese studies of oral aprepitant in patients with non-small-cell lung cancer who were receiving carboplatin-based regimens.49,50 In the Japanese studies, oral aprepitant-based triple therapy was significantly more effective than dual therapy in the subgroup of patients receiving carboplatin and pemetrexed \pm bevacizumab, but not in those receiving carboplatin and paclitaxel ± bevacizumab.49,50 The current NCCN guidelines classify carboplatin as HEC (where a three- or four-drug antiemetic regimen is recommended) if given at high doses, area under the concentration–time curve (AUC) \geq 4, and as MEC if AUC <4.¹⁶

Another consistent finding in comparative studies was a significantly higher rate of delayed CINV control with NK₁-RA-containing triple therapy vs steroid plus 5HT₃-RA dual therapy.^{36–38,43,46–48,51,53–55} Most studies also showed a higher CR rate during the acute phase.^{36,37,43,46–48,51,53–55} In addition, meta-analyses of randomized clinical trials have confirmed the significant and clinically relevant improvement in CR in patients receiving carboplatin-based chemotherapy when treated with the three-drug regimen containing an NK₁ RA compared with the dual-therapy combination.⁵⁷

Because female sex is a known risk factor for increased CINV,58 some NK₁-RA studies have analyzed CR rates in male and female participants receiving HEC. In a subgroup analysis of a trial in which patients received a 5HT, RA plus dexamethasone with or without oral aprepitant, CR rates in the oral aprepitant group were slightly lower in female (68.6%) than male (71.2%) participants, but still higher than with the two-drug regimen (36.8% and 55.0%, respectively).55 Similarly, in a post hoc multivariate analysis of two trials in which patients received a 5HT, RA plus dexamethasone with or without oral aprepitant, male sex was significantly associated with improved CR (P=0.023), but oral aprepitant improved CR regardless of patient sex.59 In a trial of oral palonosetron plus dexamethasone with or without oral netupitant, CR rates were higher in male than female participants, but all patients receiving oral NEPA had an incremental benefit in terms of CR.47 Therefore, although some differences in CR rates have been observed between male and female participants, both groups benefit from the addition of an NK, RA to a two-drug antiemetic regimen.

Across all randomized controlled trials, the magnitude of treatment difference in overall CR rate between two-drug and three-drug regimens for all patients ranged 3.6%–33% for oral aprepitant,^{43–46,48–51,54,55} 7%–17% for fosaprepitant,^{52,53} 13.1% for oral netupitant,⁴⁷ and 7.9%–15.8% for oral rolapitant.^{36–38} Aside from one study, the treatment difference in CR was consistently higher during the delayed vs acute phase.³⁶

It is difficult to compare treatment differences across studies, because of the variable patient populations and treatment regimens (ie, most HEC studies were cisplatin-based; Table 3). In some studies, patients in the three-drug arm received lower doses of dexamethasone than patients in the two-drug arm,^{44–47,51,53–55} whereas in other studies the dexamethasone dose was the same in both arms.^{36–38,43,49,50,52}

Aprepitant Cisplati Chawla et al Cisplati 2003 ⁴³ Cisplati Hesketh et al Cisplati 2003 ⁴⁶ Cisplati 2003 ⁵¹ With sol 2003 ⁵¹ Cisplati 2003 ⁵¹ Cisplati 2005 ⁴⁴ Cisplati 2005 ⁴⁴ Cisplati 2005 ⁴⁴ Cisplati 2005 ⁵⁴ Cisplati 2006 ⁵⁴ With sol Herrington et al Cisplati 2008 ⁵⁵ Mult p	Cisplatin-naïve adult patients		2000 Participante a companya da com		CR rate"			Treatment	Treatment difference, ^c P-value	alue
et al et al h et al elli et al et al l et al ston et al	latin-naïve adult patients				Acute CINV (day 1)	Delayed CINV (days 2–5)	Overall (days 1–5)	Acute CINV	Delayed CINV	Overall
et al h et al jelli et al t al st al ston et al	latin-naïve adult patients									
h et al felli et al et al et al ston et al		Cisplatin-based	Two-drug control	127	71.4	45.2	43.7	11.8, <0.05	27.5, <0.01	27.3, <0.01
h et al felli et al et al l et al ston et al	with solid tumors	regimens	en with	I 34₫	83.2	72.7	71.0			
h et al elli et al et al l et al ston et al			aprepitant							
elli et al et al l et al șton et al	Cisplatin-naïve adult patients	Cisplatin-based	Two-drug control	266	78.1	55.8	52.3	11.1, <0.001	19.6, <0.001	20.4, <0.00I
jelli et al et al l et al ston et al	with solid tumors	regimens	en with	264	89.2	75.4	72.7			
jelli et al et al l et al ston et al			aprepitant							
et al l et al gton et al	Cisplatin-naïve adult patients	Cisplatin-based	Two-drug control	286	68.4	46.8	43.3	14, <0.001	21, <0.001	19, <0.001
et al l et al ston et al	with solid tumors	regimens	en with	283	82.8	67.7	62.7			
st al l et al ston et al			aprepitant							
l et al ston et al	Cisplatin-naïve adult patients	Cisplatin-based	Two-drug control	72	49	32	26	22, <0.05	35, <0.05	33, <0.001
l et al ston et al	with solid tumors	regimens +	Three-drug regimen with	70	71	67	59			
l et al ston et al		concomitant AC ^e	aprepitant							
ston et al	Cisplatin-naïve adult patients	Cisplatin-based	Two-drug control	245	79.3	63.1	60.6	8.4, 0.005	11.0, 0.004	11.4, 0.003
ston et al	with solid tumors	regimens	en with	244	87.7	74.1	72.0			
ston et al			aprepitant							
	Adult patients with solid	Cisplatin-based	Two-drug control	16	56.2	31.2	31.2	14.2 (Iday),	28.I (I day,	20.7 (I day),
	ors	or AC regimens	Three-drug regimen with	30	70.4	59.3	51.9	10.5 (3 days),	31.2 (3 days),	24.4 (3 days),
		1	aprepitant for I day					NR		NR
			Three-drug regimen with	29	66.7	63.0	55.6			
			aprepitant for 3 days							
Takahashi et al Japan	Japanese cancer patients	Cisplatin-based	Two-drug control	150	83.3	51.7	50.3	3.7, NS	20.9, <0.001	20.2, <0.001
2010 ⁵⁵ aged	aged ≥20 years	regimens	en with	146ª	87.0	72.6	70.5			
			aprepitant							
Hu et al Chine	Chinese cisplatin-naïve adult	Cisplatin-based	Two-drug control	212	79.3	59.4	57.0	0.I, NS	14.6, 0.001	12.6, 0.007
2014 ⁴⁸ patie	patients with solid tumors	regimens	en with	209	79.4	74.0	69.6			
			aprepitant							
Ito et al CT-n	CT-naïve adult Japanese	Carboplatin-	Two-drug control	67	NR	NR	67.2	I	I	13.1, NS
2014 ⁴⁹ NSCI	NSCLC patients	based regimens	g regimen with	66	NR	NR	80.3			
			aprepitant							
ya et al	CT-naïve adult Japanese	Carboplatin-		39	100	76.9	76.9	0, NS	3.6, NS	3.6, NS
201550 NSC	NSCLC patients	based regimens	g regimen with	4	100	80.5	80.5			
			aprepitant							
Fosaprepitant										
Saito et al Japan	Japanese adult patients with	Cisplatin-	Two-drug control	173	81	49	47	13, 0.0006	16, 0.0025	17, 0.0015
2013 ⁵³ solid	solid tumors	containing	Three-drug regimen with	174	94	65	64			
		regimens	fosaprepitant							

Ruhlmann et al 2016 ⁵²	Women undergoing RT + CT for gynecologic cancers	Cisplatin + RT for 5 weeks	Two-drug control Three-drug regimen with fosaprepitant	116 118	88 92	NR NR	65 72	4, 0.255	1	7, 0.194
Netupitant							_			
Hesketh et al	CT-naïve adult patients with	Cisplatin-	Two-drug control	136	89.7	80.1	76.5	8.8, ≤0.01	10.3, ≤0.05	13.1, 0.004
20 I 4 ⁴⁷	solid tumors	containing	Three-drug regimen with	135^{a}	98.5	90.4	89.6			
		regimens	netupitant							
Zhang et al	CT-naïve adult patients with	Cisplatin-	Three-drug control	416	87.0	74.3	72.4	–2.5, NR	3.6, NR	I.4, NR
201856	solid tumors	containing	Three-drug regimen with	412	84.5	9.77	73.8			
		regimens	netupitant							
Rolapitant										
Rapoport et al	Adult patients with cancer	Cisplatin-	Two-drug control	16	66.7	48.9	46.7	20.9, 0.001	14.7, 0.045	15.8, 0.032
201536		containing	Three-drug regimen with	90ª	87.6	63.6	62.5			
		regimens	rolapitant							
Rapoport et al 2015 ³⁷ –	Cisplatin-naïve adult patients	Cisplatin-	Two-drug control	535	76.6	58.5	60.2	7.0, 0.0001	10.3, 0.0045	11.2, 0.0005
pooled analysis of two	with solid tumors	containing	Three-drug regimen with	535	83.6	68.8	71.4			
studies		regimens	rolapitant							
Schwartzberg et al	HEC- or MEC-naïve adult	AC regimens	Two-drug control	359	76.7	59.6	54.9	0.2, 0.9659	7.2, 0.0465	7.9, 0.0332
201538	patients with solid tumors		Three-drug regimen with	344	76.9	66.8	62.8			
			rolapitant							
Votes: ^a Antiemetic regimens (between three-drug regimen gr	comprised a 5HT ₃ -receptor antagon oup vs control: ^d additional group(s)	iist and dexamethasone received a lower-than-	, with or without an NK ₁ -recep approved dose of NK ₁ -receptor	otor anta inhibitor	gonist; ^b CR was (, and are not sho	defined as no episod(wn in the table; °dox(es of vomiting and orubicin + cycloph	no rescue antiem osphamide; ^f statist	etic therapy; ^c diffel ical comparison vs	ence in CR rate placebo arm not
Votes: ^a Antiemetic regimens (between three-drug regimen gr	Notes: *Antiemetic regimens comprised a 5HT ₃ -receptor antagonist and dexamethasone, with or without an NK ₁ -receptor a between three-drug regimen group vs control, "additional group(s) received a lower-than-approved dose of NK ₁ -receptor inhib undershows there are no similinear difference in the CP are between originar stations constrant and ver Los or dove Lo	ist and dexamethasone received a lower-than-	approved dose of NK ₁ -receptor i	otor anta inhibitor	gonist; ^b CR was (, and are not sho	defini vwn ir	ed as no episode the table; °dox	ed as no episodes of vomiting and 1 the table; ^e doxorubicin + cycloph	ed as no episodes of vomiting and no rescue antiem 1 the table: °doxorubicin + cyclophosphamide: 'statist	Notes: "Antiemetic regimens comprised a 5HT ₃ -receptor antagonist and dexamethasone, with or without an NK ₁ -receptor antagonist; ^b CR was defined as no episodes of vomiting and no rescue antiemetic therapy; "difference in CR rate between three-drug regimen group vs control: "additional group(s) received a lower-than-approved dose of NK ₁ -receptor inhibitor, and are not shown in the table: "doxorubicin + cyclophosphamide: 'statistical comparison vs placebo arm not indershor: there are constroned in the CR rate hervoen parianter abive a not dow 1 or on dow 1.0. or dow 1.3.

undertaken; there was no significant difference in the CR rate between patients taking aprepitant only on day 1 –3. Abbreviations: AC, anthracycline + cyclophosphamide; CINV, chemotherapy-induced nausea and vomiting: CR, complete response; CT, chemotherapy; HEC, highly emetogenic chemotherapy; 5HT₃, 5-hydroxytryptamine type 3; MEC, moderately emetogenic chemotherapy; NK, neurokinin 1; NR, not reported; NS, not significant; NSCLC, non-small-cell lung cancer; RT, radiotherapy.

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Several meta-analyses have confirmed that three-drug regimens containing an NK₁ RA are significantly more effective than two-drug regimens for achieving CR in patients with CINV.^{60–62} The estimated risk difference for overall CR between the two types of antiemetic regimens is 14% (95% CI 12%–17%) in patients receiving any type of HEC, 16% (95% CI 14%–19%) in patients receiving cisplatin-based HEC, and 11% (95% CI 7%–15%) in patients receiving AC-based HEC,⁶¹ all of which exceed the level considered to be clinically meaningful (\geq 10%).^{63,64}

Few studies have directly compared the efficacy of three-drug antiemetic regimens using different NK, RAs. Studies directly comparing different three-drug regimens in patients receiving HEC have found similar CR rates among regimens. For example, comparable rates of overall, acute, and delayed CR were reported in patients receiving a fosaprepitant- or oral aprepitant-based three-drug regimen.⁶⁵ In a comparison of CR rates in patients receiving oral NEPA plus dexamethasone vs those receiving oral aprepitant plus ondansetron plus dexamethasone as an exploratory end point, there was no statistical significance,⁴⁷ and the threshold of clinically meaningful difference was not met,63,64 as shown in Table 3. In a recent study, CR rates showed noninferiority of oral NEPA plus dexamethasone vs oral aprepitant plus granisetron plus dexamethasone.56 Another study showed no significant difference between oral NEPA-based and oral aprepitant-based triple therapy in patients receiving HEC,66 including in the subgroup of patients receiving carboplatinbased chemotherapy.67

Recently, researchers have investigated the effect of adding olanzapine to a three-drug antiemetic combination in randomized, double-blind, placebo-controlled trials with aprepitant or fosaprepitant.^{68,69} The olanzapine plus three-drug combination elicited a significantly higher CR rate in the acute, delayed, and overall phases compared with placebo plus a three-drug combination in patients receiving HEC.⁶⁹ A study in patients receiving HEC or MEC also found significantly higher CR rates with olanzapine treatment in the delayed and overall phases.⁶⁸

Efficacy with moderately emetogenic chemotherapy

Studies in which an NK₁ RA was added to a standard twodrug regimen of a 5HT₃ RA and dexamethasone in patients receiving MEC are summarized in Table 4.^{38,70–82} However, in some studies, patients received chemotherapy regimens that have since been recategorized as HEC,^{70,71,80–82} such as AC-based regimens. Carboplatin is now classified as HEC if given at high doses (AUC \geq 4) and as MEC at lower doses (AUC <4).¹⁶ Most studies have shown a significantly greater improvement in overall CR rates with the three-drug vs the two-drug regimen, with greater differences observed in HEC-treated patients.^{70–72,75–80} Like the HEC studies, the triple-antiemetic combination tended to have a more marked effect on CR rates in the delayed than the acute phase in patients receiving MEC. This difference between acute and delayed antiemetic effect was most marked in two studies in which patients were receiving carboplatin-based chemotherapy for solid tumors^{47,80} and another in which patients with multiple myeloma were receiving high-dose melphalan prior to autologous stem-cell transplant.⁷⁷

Female sex is one of several risk factors for increased CINV,⁵⁸ and a difference in CR rates in the overall phase between male (83.0%) and female (77.9%) patients was reported in a trial of a $5HT_3$ RA plus dexamethasone with or without oral rolapitant in patients receiving carboplatin-based chemotherapy. However, both groups had significantly higher CR rates than the sex-matched patients in the control group (67.7% and 62.1%, respectively).⁷²

A recent meta-analysis supported the incremental benefit of an NK, RA, and suggested that the magnitude of effect of NK, RA-based triple therapy on CR varied depending on the MEC regimen administered.⁸³ The effect on overall CR was greatest in patients who were receiving carboplatin-based chemotherapy (Figure 1), with a risk difference of 15% between two-drug regimens and NK, RA-based three-drug regimens. A significant effect in favor of the three-drug regimen was also seen in patients who received MEC that did not contain oxaliplatin or carboplatin, but not in patients receiving oxaliplatin-based regimens.83 The odds of achieving CR in the acute and delayed phases were significantly better with the three-drug than with the two-drug regimens in patients taking carboplatin. Also, delayed CR rates were significantly higher following a three-drug regimen in MEC patients not receiving oxaliplatin or carboplatin (Figure 1).83

For patients receiving MEC, current guidelines recommend NK₁ RAs in those at high risk of CINV.¹⁶ These include female patients, those aged <55 years, people without a history of habitual alcohol use, and nonsmokers.⁸⁴

Clinical efficacy: nausea control

Because CR and other measures of antiemetic efficacy generally focus on control of emesis, some trials have also included nausea end points, although nausea control is typically a secondary or exploratory end point. This patient-reported outcome is often measured using a 100 mm VAS (0, no nausea; 100, worst possible nausea)^{36,46,56} or a 4-point nausea-severity score (0, none; 1, mild; 2, moderate; 3, severe).^{53,55,75} End points reported include "no nausea"

Study	Patients	Chemotherapy	Study Patients Chemotherapy Antiemetic regimens ^a	z	CR rate ^b			Treatm	Treatment difference.	nce, ^c
		2	D					P-value		
				1	Acute CINV (day 1)	Delayed CINV (days 2-5)	Overall (days I–5)	Acute CINV	Delayed CINV	Overall
Aprepitant	-	-	-				-			
Warr et al	CT-naïve adult patients with	Cyclophosphamide-	Two-drug control	428	69	49	42	ŵ	6,	9,
200578	breast cancer	based MEC	Three-drug regimen with aprepitant	438	76	55	51	0.034	0.064	0.015
Yeo et al	Chinese women with breast	AC-based	Two-drug control	62	72.6	57.8	41.9	-0.5,	6.6,	4.9,
200981	cancer	regimens ^d	Three-drug regimen with aprepitant	62	72.1	64.4	46.8	0.95	0.51	0.58
Rapoport et al	HEC- or MEC-naïve adult	Any MEC regimen	Two-drug control	418	80.3	60.9	56.3	8.9,	9.9,	13.4,
20 I 0 ⁷⁶	patients with solid tumors		Three-drug regimen with aprepitant	430	89.2	70.8	68.7	<0.001	<0.001	<0.001
Tanioka et al	Aprepitant-naïve, nondrinking	Irinotecan- or	Two-drug control	46	95.7	52.1	52.1	2.1,	10.1,	10.1,
2013 ⁸²	Japanese women aged 20–69 years	carboplatin-based MEC	Three-drug regimen with aprepitant	45	97.8	62.2	62.2	SN	0.33	0.33
Schmitt et al	Adult patients with multiple	High-dose	Two-drug control	181	90	46	41	7,	26,	17,
2014 ⁷⁷	myeloma undergoing	melphalan regimen	Three-drug regimen with aprepitant	181	97	60	58	0.022	0.011	0.0042
	conditioning prior to autologous SCT									
Nishimura et al	lapanese patients	Oxaliplatin-based	Two-drug control	206	92.4	75.4	74.3	2.3,	9.6,	10.7,
2015 ⁷⁵	aged ≥20 years with	MEC	Three-drug regimen with	207	94.7	85.0	85.0	0.37	0.02	0.01
	colorectal cancer		fosaprepitant or aprepitant ^e							
Yahata et al	Japanese women aged 20–80	TC regimen ^d	Two-drug control	152	90.4	49.3	47.3	3.6,	14.3,	14.3,
2016 ⁸⁰	years with gynecologic cancers		Three-drug regimen with aprepitant	155	94.0	63.6	61.6	NS	0.0072	0.0073
Kim et al	Korean patients aged \ge 20	CT containing one	Two-drug control	243	97.9	71.2	70.4	-2.1,	3.1,	3.0,
201773	years with solid tumors	or more MEC agents	Three-drug regimen with aprepitant	237	95.8	74.3	73.4	ΣŢ	ЪŢ	0.458
Fosaprepitant			•							
Kitayama et al	CT-naïve Japanese adult	MEC (mostly	Two-drug control	35	94	74	74	7,	-5,	- 5,
201574	patients with solid tumors	oxaliplatin- or	Three-drug regimen with	35	100	69	69	SN	NS	NS
		irinotecan-based)	fosaprepitant							
Nishimura et al	Japanese patients	Oxaliplatin-based	Two-drug control	206	92.4	75.4	74.3	2.3,	9.6,	10.7,
2015 ⁷⁵	aged ≥ 20 years with	MEC	Three-drug regimen with	207	94.7	85.0	85.0	0.37	0.02	0.01
	colorectal cancer		fosaprepitant or aprepitant ^e							
Weinstein et al	HEC- or MEC-naïve adult	Non-AC MEC	Two-drug control	507	91.0	68.5	6.9	2.2,	10.4,	10.2,
201679	patients with solid tumors		Three-drug regimen with	508	93.2	78.9	77.1	0.0184	<0.001	<0.001
			fosaprepitant							
Netupitant										
Aapro et al	CT-naïve adult patients with	AC-containing	Two-drug control	725	85.0	69.5	66.6	3.4,	7.4,	7.7,
20147	solid tumors	regimens ^a	Three-drug regimen with netupitant	724	88.4	76.9	74.3	0.047	0.001	0.00 I
									(Co	(Continued)

 Table 4 CRs in three-drug vs two-drug regimen trials in MEC (approved agents at recommended doses)

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Study	Patients	Chemotherapy	Antiemetic regimens ^a	z	CR rate ^b			Treatme P-value	Treatment difference, ^c P-value	ice, ^c
				I	Acute CINV (day 1)	Acute CINV Delayed CINV Overall (day 1) (days 2–5) (days 1–	Overall Acute (days I–5) CINV	Acute CINV	Delayed CINV	Overall
Aapro et al	CT-naïve adult patients with	AC-containing	Two-drug control	651	NR	NR	66.6–74.6	1	1	7.7–13.6,
201770	solid tumors	regimens ^d	Three-drug regimen with netupitant 635 NR	635	NR	NR	74.3-83.8 ^g			≤0.001
Rolapitant			1							
Schwartzberg et al	HEC- or MEC-naïve adult	Non-AC regimens	Two-drug control	307 84.4	84.4	63.8	61.2	6.3,	12.3,	13.6,
2015 ³⁸	patients with solid tumors		Three-drug regimen with rolapitant	322	90.7	76.1	74.8	0.0163	0.0008	0.0003
Hesketh et al 2016 ⁷² –	HEC- or MEC-naïve adult	Carboplatin-	Two-drug control	209	88.0	65.6	64.6	3.7,	16.7,	15.6,
subgroup analysis of Schwartzberg et al 2015 ³⁸	patients with solid tumors	containing regimens	Three-drug regimen with rolapitant	192 91.7	91.7	82.3	80.2	0.231	<0.001	<0.001
Notes: ^a Antiemetic regimens between three-drug regimen gi categorized as HEC. The AC r	comprised a 5HT ₃ -receptor antagonis oup vs control. ^d At the time the stud egimen for Yeo et al ⁶¹ was doxorubic	st and dexamethasone, with by was performed, AC and ⁻ in + cyclophosphamide. °TI	Notes: "Antiemetic regimens comprised a 5HT ₃ -receptor antagonist and dexamethasone, with or without an NK ₁ -receptor antagonist. ^b CR was defined as no episodes of vomiting and no rescue antiemetic therapy. ^c Difference in CR rate between three-drug regimen group vs control. ^d At the time the study was performed, AC and TC regimens were classified as MEC. AC has since been categorized as HEC, and carboplatin at the dose used in the study by Yahata et al ⁸⁰ is also categorized as HEC. The AC regimen for Yeo et al ⁸¹ was doxorubicin + cyclophosphamide. ^a These patients received either aprepitant or fosaprepitant as part of the three-drug regimen. Not tested, because significance in the key end point	CR was s since b osaprep	defined as no epis een categorized a: itant as part of the	odes of vomiting and r HEC, and carboplatin three-drug regimen. ¹	no rescue antierr at the dose usec Not tested, beca	netic therap I in the stuc use signific:	y. 'Difference Jy by Yahata ei ince in the key	in CR rate t al ⁸⁰ is also r end point

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Abbreviations: AC, anthracycline + cyclophosphamide; CINV, chemotherapy-induced nausea and vomiting; CR, complete response; CT, chemotherapy; HEC, highly emetogenic chemotherapy; 5HT, 5-hydroxytryptamine type 3; MEC, (overall CR rate) was not significant. ⁸These data pertain to multiple cycles of treatment.

emetogenic chemotherapy: NK,, neurokinin 1; NR, not reported: NS, not significant; NT, not tested for significance; SCT, stem-cell transplant; TC, paclitaxel + carboplatin moderately

 $(<5 \text{ mm on } 100 \text{ mm VAS or } 0 \text{ on a } 4\text{-point scale})^{36,46,56}$ and "no significant nausea" (<25 mm on 100 mm VAS or 0 and 1 on a 4-point scale),^{53,55,75} and assessments may be made during acute, delayed, and overall phases, but are more often assessed during the overall phase.

Efficacy with highly emetogenic chemotherapy

Randomized controlled trials in patients receiving HEC are summarized in Table 5.^{36,37,43,46,47,49,51–56} Most of these trials reported no significant improvements in nausea control (percentage of patients with no nausea or no significant nausea) in any CINV phase with the addition of an NK, RA to a two-drug antiemetic regimen. Of those trials that did report a significant improvement in nausea control during any phase, 36,37,43,47,51,55 only two trials and one pooled analysis of two trials reported significant improvements in nausea control across the acute, delayed, and overall phases.^{36,37,47} In trials where addition of an NK₁ RA significantly improved nausea control in the overall phase (which was assessed most frequently), nausea-control rates ranged 49%-52.7% for "no nausea"37,43,51 and 52%-89.6% for "no significant nausea"37,47 in patients who received the three-drug regimen. However, comparisons across trials must be made with caution, because of differences in study design, patient populations, nausea assessments, chemotherapy regimens, and antiemetic regimens.

Efficacy with moderately emetogenic chemotherapy

Randomized controlled trials in patients receiving MEC are summarized in Table 6.38,70-72,74-82 About half the MEC trials tabulated reported no significant improvements in nausea control in any CINV phase with the addition of an NK, RA to a two-drug antiemetic regimen. Of those that did report a significant improvement in nausea control in any phase, 70-72,75,76,79,80 none showed significant improvements in nausea control across the acute, delayed, and overall phases. In trials where addition of an NK₁ RA significantly improved nausea control in the overall phase, nausea-control rates ranged 62.5%-74.6% for no nausea^{71,72} and 73.6%-88.8% for no significant nausea^{75,76,79,80} in patients who received the three-drug regimen. However, again, differences in study design, patient populations, nausea assessments, chemotherapy, and antiemetic regimens administered limit comparisons across trials.

Bioequivalence studies

Approvals of IV rolapitant, HTX019 (IV formulation of aprepitant), and IV NEPA were based on demonstration of bioequivalence with the corresponding approved oral agents.^{31,33} A study in healthy volunteers showed that a single



Figure I CR with NK₁-RA-containing triple therapy vs dual therapy with a 5HT₃ RA and dexamethasone.

Notes: ORs from a meta-analysis of randomized studies in patients receiving different types of MEC. Data from Jordan et al.⁸³

Abbreviations: CR, complete response; 5HT₃, 5-hydroxytryptamine type 3; MEC, moderately emetogenic chemotherapy; NK₁, neurokinin 1; OR, odds ratio; RA, receptor antagonist.

IV dose of rolapitant 166.5 mg was bioequivalent to a single oral dose of 180 mg.³⁵ As expected, C_{max} was higher with IV than with oral rolapitant, and occurred at an earlier time point, but the elimination $t_{1/2}$ was similar. Both rolapitant formulations were well tolerated, with a similar overall incidence of adverse events.³⁵

A single dose of HTX019 130 mg was bioequivalent to the approved formulation of fosaprepitant 150 mg IV in healthy volunteers. Plasma concentrations of aprepitant from both infusions were essentially superimposable at 0.75 hours after administration. Both agents were well tolerated, although HTX019 was associated with a lower rate of infusion reactions.^{85,86}

Safety

In general, the NK₁ RAs are well tolerated and not associated with specific adverse events,⁸⁷ although it can be difficult to distinguish adverse events related to antiemetics from those associated with chemotherapy. In randomized comparisons, the incidence of associated adverse events for three-drug regimens containing oral NK₁ RAs was similar to that for two-drug regimens in patients receiving HEC or MEC.^{37,38,43,46,47,51,71,82} The most common adverse events with the oral agents are fatigue/asthenia, headache, hiccups, and constipation.⁸⁷

In a bioequivalence study of healthy volunteers, oral and IV formulations of rolapitant had a similar overall incidence

of adverse events. IV rolapitant contains polyoxyl 15 hydroxystearate, a synthetic surfactant with a limited safety profile.^{31,88,89} Two patients in the IV rolapitant group (2.8%)developed a mild infusion-site reaction, and the incidence of headache was higher with IV than with oral rolapitant (8.5% vs 3.0%, respectively).³⁵ In Phase I studies, IV rolapitant was less likely than oral rolapitant to exhibit drug interactions associated with P-glycoprotein or BCRP.42 Soon after the formulation's approval, a US Food and Drug Administration MedWatch safety alert was issued to health-care providers on January 16, 2018 warning against hypersensitivity reactions, including anaphylaxis and anaphylactic shock, which may occur during or following administration of IV rolapitant. Moreover, the alert recommended avoiding administration of the drug if the patient was hypersensitive to any ingredient of the drug formulation.88 Following that warning, a press release issued by the manufacturer on February 27, 2018 announced the suspension of IV rolapitant distribution.89

Fosaprepitant, the IV prodrug of aprepitant, is associated with a high incidence of infusion-site reactions and hypersensitivity, including anaphylaxis,^{32,90} and the prescribing information includes a warning about the risk of these events.³² Patients should be monitored during and after IV infusion of fosaprepitant, and discontinued if hypersensitivity reactions occur.³² In a Phase III trial of a two-drug regimen of a 5HT₃ RA plus dexamethasone with or without fosaprepitant in patients scheduled to receive non-AC MEC,⁷⁹

study	Patients	Chemotherapy	Antiemetic regimens ⁴	z	Nausea	Nausea (% of n	Nausea control (% of natients)		Betwee study o	Between group P-value for study group vs control	alue for trol
						Acute	Delayed	Overall	Acute	Delayed	Overall
Aprepitant	_						_	-			
Chawla et al	Cisplatin-naïve adult	Cisplatin-based	Two-drug control	127	No nausea [∈]	66.7	36.5	34.1	SN	<0.01	<0.01
200343	patients with solid tumors	regimens	Three-drug regimen with aprepitant	134 ^b		71.8	58.3	52.7			
			Two-drug control	127	No significant	87.3	62.7	58.7	NS	<0.01	<0.01
			Three-drug regimen with aprepitant	134	nausea ^d	90.8	83.3	81.7			
Hesketh et al	Cisplatin-naïve adult	Cisplatin-based	Two-drug control	266	No nausea [€]	69.1	47.7	44.2	٩N	SN	NS
2003*	patients with solid tumors	regimens	Three-drug regimen with aprepitant	264		72.3	51.0	47.5			
			Two-drug control	266	No significant	86.5	68.5	66.0	SS	SN	SN
			Three-drug regimen with aprepitant	264	nausea ^d	90.6	75.3	73.2			
Poli-Bigelli et al	Cisplatin-naïve adult	Cisplatin-based	Two-drug control	286	No nausea [€]	ЧP	40	39	I	<0.01	<0.05
200351	patients with solid tumors	regimens	Three-drug regimen with aprepitant	283		٩N	53	49			
			Two-drug control	286	No significant	٩N	65	64	I	NS	NS
			Three-drug regimen with aprepitant	283	nausea ^d	٩N	73	71			
Schmoll et al	Cisplatin-naïve adult	Cisplatin-based	Two-drug control	245	No significant	89.5	69.7	72.1	NS	NS	NS
2006 ⁵⁴	patients with solid tumors	regimens	Three-drug regimen with aprepitant	244	nausea ^d	92.1	73.1	75.9			
Takahashi et al	Japanese cancer patients	Cisplatin-based	Two-drug control	150	No nausea [€]	66.0	26.2	24.2	NS	NS	NS
20 I 0 ⁵⁵	aged ≥ 20 years	regimens	Three-drug regimen with aprepitant	I46 ^d		67.I	34.9	34.2			
			Two-drug control	150	No significant	88.0	56.4	55.7	NS	<0.01	SN
			Three-drug regimen with aprepitant	146	nausea ^f	90.4	72.6	69.2			
lto et al	CT-naïve adult Japanese	Carboplatin-based	Two-drug control	67	Nausea	20.0	56.7	58.3	SN	NS	SN
2014 ⁴⁹	NSCLC patients	regimens	Three-drug regimen with aprepitant	67	frequency ^g	25.0	46.7	46.7			
Fosaprepitant											
Saito et al	Japanese adult patients	Cisplatin-containing	Two-drug control	173	No nausea [€]	67.5	24.7	24.I	NS	NS	NS
2013 ⁵³	with solid tumors	regimens	Three-drug regimen with fosaprepitant	174		67.6	30.6	30.1			
			Two-drug control	173	No significant	84.9	58.4	58.4	NS	NS	NS
			Three-drug regimen with fosaprepitant	174	nausea ^f	90.2	66.5	65.3			
Ruhlmann et al	Women undergoing RT	Cisplatin + RT for	Two-drug control	116	No nausea ^e	62	NR	32	NS	I	NS
2016 ⁵²	+ CT for gynecologic	5 weeks	Three-drug regimen with fosaprepitant	118		71	NR	47			
	cancers		Two-drug control	116	No significant	90	NR	67	NS	I	NS
			Three-drug regimen with fosaprepitant	18	nausea ^f	95	NR	81			
Netupitant											
Hesketh et al	CT-naïve adult patients	Cisplatin-containing	Two-drug control	136	No significant	93.4	80.9	79.4	≤0.01	≤0.05	≤0.01
201447	with solid tumors	regimens	Three-drug regimen with netunitant	135	prostice	00	001	7 00			

Zhang et al 2018 ⁵⁶	CT-naïve adult patients with solid tumors	Cisplatin-containing regimens	Three-drug control Three-drug regimen with netupitant	416 412	No nausea ^c	67.8 68.9	54.3 53.2	51.4 49.3	SN	SN	NS
			Three-drug control Three-drug regimen with netupitant	416 412	No significant nausea ^d	87.3 89.8	72.8 78.2	70.4 75.7	NS	SN	NS
Rolapitant											
Rapoport et al	Adult patients with cancer	Cisplatin-containing	Two-drug control	16	No nausea [€]	NR	NR	R	NS	NS	NS
2015 ³⁶		regimens	Three-drug regimen with rolapitant	90		NR	NR	R			
			Two-drug control	16	No significant	73.3	47.8	42.2	<0.05	<0.05	<0.01
			Three-drug regimen with rolapitant	90	nausea ^d	86.5	64.4	63.2			
Rapoport et al	Cisplatin-naïve adult	Cisplatin-containing	Two-drug control	535	No nausea [€]	64	44	42	NS	0.0002	0.0004
2015 ³⁷ – pooled	patients with solid tumors	regimens	Three-drug regimen with rolapitant	535		70	56	52			
analysis of two			Two-drug control	535	No significant	83	67	65	0.0090	0.0108	0.0174
studies			Three-drug regimen with rolapitant	535	nausea ^d	88	74	72			
Notes: ^a Comprised nausea = <5 mm on	Notes: ^a Comprised a 5HT ₃ -receptor antagonist and dexamethasone, with or without nausea = <5 mm on 100 mm VAS; ⁴ no significant nausea = <25 mm on 100 mm VAS; ⁴	examethasone, with or wit a = <25 mm on 100 mm V	hout an NK,-receptor antagonist; ^b additional group(s) received a lower-than-approved dose of NK,-receptor inhibitor; and are not shown in the table; 'no AS; "no nausea = nausea score 0 on a 4-point scale (0, none; 1, mild; 2, moderate; 3, severe); 'no significant nausea = nausea scores 0 and 1 on a 4-point scale	oup(s) rec ale (0, nor	:eived a lower-than- ne; I, mild; 2, moder	approved d ate; 3, sever	ose of NK ₁ -ree re); ^f no significa	ceptor inhibito ant nausea = n	or, and are no ausea scores	ot shown in th 0 and 1 on a 4	e table; °no -point scale
(0, none; 1, mild; 2, n Abbreviations: CT	(0, none; 1, mild; 2, moderate; 3, severe): ⁸ nausea frequency evaluated daily by patient Abbreviations: CT, chemotherapy: HEC, highly emetogenic chemotherapy: 5HT ₃ ,	ency evaluated daily by pati togenic chemotherapy; 5H	ent questionnaire (no details reported). T ₃ , 5-hydroxytryptamine type 3; NK ₁ , neurokinin 1; NP, not performed; NR, not reported; NS, not significant; NSCLC, non-small-cell lung cancer; RT,	nin I; NP	, not performed; N	R, not repo	orted; NS, not	significant; N	SCLC, non-s	mall-cell lung	cancer; RT,

polysorbate 80, a surfactant used to solubilize fosaprepitant, and associated with infusion reactions and hypersensitivity when used in formulations of other pharmaceutical agents and vaccines.91,92 HTX019, the IV formulation of aprepitant, is free of polysorbate 80 and other synthetic surfactants.³³ In healthy volunteers, HTX019 was bioequivalent to fosaprepitant, but associated with a lower rate of infusion reactions.85 Within an hour of IV infusion, adverse events were reported in 20 participants receiving fosaprepitant and one participant receiving HTX019.85 However, as HTX019 contains aprepitant, the same active agent as fosaprepitant, the prescribing information for both IV agents includes the same warnings and precautions about hypersensitivity reactions.32,33 Oral and IV formulations of NEPA are now approved in the United States. In a randomized, double-blind Phase III study comparing IV and oral NEPA (each with dexamethasone) prior to initial and repeated cycles of non-AC HEC in 404 patients, both NEPA formulations were similarly well tolerated.93,94 No serious adverse events related to IV or oral NEPA were recorded, the most common adverse event was constipation in both treatment groups, and the incidence of

infusion-site reactions were reported in 2.2% of patients

who received fosaprepitant compared with 0.6% of patients who did not.32 These reactions may be associated with

adverse events did not increase over repeated cycles.93,94 No patients receiving IV NEPA developed an infusion-site reaction. There were no clinically relevant electrocardiographic abnormalities or cardiac safety concerns with either formulation.93,94

All NK, RAs have the potential for drug-drug interactions, 29-33 so careful assessment of concomitant medications is required when deciding which agent to use. For example, dexamethasone is a CYP3A4 substrate, so a lower dose of dexamethasone (12 mg) is recommended on day 1 of antiemetic treatment with regimens containing oral or injectable emulsion aprepitant, fosaprepitant, or oral or IV NEPA than with regimens including oral rolapitant (dexamethasone 20 mg).¹⁶ Because some NK, RAs are substrates, weakmoderate (dose-dependent) inhibitors, and inducers of CYP3A4, they may increase the plasma concentrations of chemotherapeutic agents that are metabolized by CYP3A4, including taxanes, irinotecan, vinca alkaloids, and tyrosinekinase inhibitors.^{87,95} Consequently, physicians should be vigilant for the possibility of an increased risk of adverse events when using NEPA, aprepitant, or fosaprepitant in patients receiving chemotherapy regimens containing these agents. Care should be taken when administering rolapitant with CYP2D6 substrates, including metoprolol and venlafaxine.87,92

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Study	Patients	Chemotherapy	Antiemetic regimens ^a	z	Nausea	Nausea	Nausea control		Betwee	Between group P-value	value
					measure	(% of p;	(% of patients)		for stud	for study group vs control	control
						Acute	Delayed	Overall	Acute	Delayed	Overall
Aprepitant											
Warr et al	CT-naïve adult patients	Cyclophosphamide-	Two-drug control	428	No nausea ^b	NR	NR	33	NR	NR	NS
200578	with breast cancer	based MEC	Three-drug regimen with aprepitant	438		NR	NR	33			
			Two-drug control	428	No significant	RR	NR	56	RR	NR	NS
			Three-drug regimen with aprepitant	438	nausea ^c	NR	NR	61			
Yeo et al	Chinese women with	AC-based regimens ^c	Two-drug control	62	No nausea ^b	NR	NR	35.5	NR	NR	NS
2009 ⁸¹	breast cancer		Three-drug regimen with aprepitant	62		NR	NR	30.6			
			Two-drug control	62	No significant	NR	NR	62.9	RR	NR	NS
			Three-drug regimen with aprepitant	62	nausea ^c	NR	NR	66.1			
Rapoport et al	HEC- or MEC-naïve adult	Any MEC regimen	Two-drug control	418	No significant	RR	NR	66.4	RR	NR	<0.05
201076	patients with solid tumors		Three-drug regimen with aprepitant	430	nausea ^c	NR	NR	73.6			
Tanioka et al	Aprepitant-naïve, non-	Irinotecan- or	Two-drug control	46	No nausea ^d	NR	NR	39	NR	NR	NS
20 I 3 ⁸²	drinking Japanese women	carboplatin-based MEC	Three-drug regimen with aprepitant	45		NR	NR	53			
	aged 20–69 years		Two-drug control	46	No significant	NR	NR	76	NR	NR	NS
			Three-drug regimen with aprepitant	45	nausea ^e	NR	NR	83			
Schmitt et al	Adult patients with	High-dose melphalan	Two-drug control	181	No nausea ^b	R	NR	78	NR	NR	NS
201477	multiple myeloma	regimen	Three-drug regimen with aprepitant	181		NR	NR	85			
	undergoing conditioning		Two-drug control	181	No significant	N R	NR	88	RR	NR	NS
	prior to autologous SCT		Three-drug regimen with aprepitant	181	nausea ^c	NR	NR	94			
Nishimura et al	Japanese patients aged	Oxaliplatin-based MEC	Two-drug control	206	No nausea ^d	90.2	61.8	59.6	NS	SN	NS
201575	≥20 years with colorectal cancer		Three-drug regimen with fosaprepitant or aprepitant ⁶	207		93.6	66.3	65.2			
			Two-drug control	206	No significant	96.2	81.4	80.9	NS	0.47	0.034
			Three-drug regimen with fosaprepitant or	207	nausea ^e	98.9	88.8	88.8			
			aprepitant								
Yahata et al	Japanese women aged	TC regimen ^g	Two-drug control	152	No nausea ^d	89.7	33.6	33.6	NS	NS	NS
2016 ⁸⁰	20–80 years with		Three-drug regimen with aprepitant	155		89.4	40.4	39.7			
	gynecologic cancers		Two-drug control	152	No significant	98.0	76.0	74.4	NS	0.027	0.014
			Three-drug regimen with aprepitant	155	nausea ^e	98.7	85.4	85.4			
Fosaprepitant											
Kitayama et al	CT-naïve Japanese adult	MEC (mostly oxaliplatin-	Two-drug control	35	No nausea ^h	77	46	46	NS	NS	NS
201574	patients with solid tumors	or irinotecan-based)	Three-drug regimen with fosaprepitant	35		86	60	60			

Weinstein et alHEC- or MEC-naïve adult201679HEC- or MEC-naïve adult201679patients with solid tumorsAapro et alCT-naïve adult patients201770with solid tumors201770with solid tumors		I hree-drug regimen with tosaprepitant or aprepitant ⁶	207		93.6	66.3	65.2			
ein et al itant et al et al		Two-drug control Three-drug regimen with fosaprepitant or aprepitant ^f	206 207	No significant nausea ^e	96.2 98.9	81.4 88.8	80.9 88.8	NS	0.47	0.034
itant et al et al	Non-AC MEC	Two-drug control Three-drug regimen with fosaprepitant	507 508	No nausea ^b	R R	AR AR	61.6 65.3	R	NR	NS
et al et al		Two-drug control Three-drug regimen with fosaprepitant	507 508	No significant nausea ^c	R R	R R	78.3 83.1	R	NR	0.026
et al										
et al	AC-containing regimens ^g	Two-drug control	725	No significant	87.9	71.3	69.1	SN	0.014	0.020
et al		Three-drug regimen with netupitant	724	nausea ^c	87.3	76.9	74.6			
	AC-containing regimens ^g	Two-drug control	651	No significant	NR	R	69.1	NR	NR	0.020
		Three-drug regimen with netupitant	635	nausea ^{c,i}	NR	R	74.6			
Rolapitant										
Schwartzberg et al HEC- or MEC-naïve adult	MEC or AC-containing	Two-drug control	307	No nausea ^b	66	45	42	NS	NS	NS
2015 ³⁸ patients with solid tumors	regimens ^g	Three-drug regimen with rolapitant	322		65	48	45			
		Two-drug control	307	No significant	85	69	67	NS	NS	NS
		Three-drug regimen with rolapitant	322	nausea ^c	82	73	71			
Hesketh et al HEC- or MEC-naïve adult	Carboplatin-containing	Two-drug control	209	No nausea ^b	77.0	53.6	51.2	NS	0.034	0.023
2016 ⁷² – subgroup patients with solid tumors	regimens	Three-drug regimen with rolapitant	192		80.7	64.1	62.5			
analysis of		Two-drug control	209	No significant	91.4	74.2	72.7	NS	0.050	NS
Schwartzberg et al 2015 ³⁸		Three-drug regimen with rolapitant	192	nausea ^c	90.6	82.3	80.7			
Notes: ³ Antiemetic regimens comprised a 5HT ₃ -receptor antagonist and dexamethasone, with or without an NK ₁ -receptor antagonist. ⁸ No mausea = <5 mm on 100 mm VAS. ⁴ No significant nausea = <25 mm on 100 mm VAS. ⁴ No nausea = <10 mones: 1, mild: 2, moderate; 3, severe). ¹ No significant nausea = nausea score 0 and 1 on a 4-point scale (0, none: 1, mild: 2, moderate; 3, severe). ¹ Included in aprepirant and fosaprepirant sections, as patients received either agent. ⁴ At the time these studies were performed, AC and TC regimens were classified as MEC. AC has since been categorized as HEC, as has carboplatin at the dose used in the study by Yahata et al. ^{80 n} T oral control, defined as no nausea. ¹ Data for cycle 1 of this multicycle study. MK , neurotherapy; HEC, highly emetogenic chemotherapy; 5HT ₃ , 5-hydroxytryptamine type 3; MEC, moderately emetogenic chemotherapy; NK ₁ , neurokinin 1; NR, not reported; NS.	or antagonist and dexamethaso ild: 2. moderate: 3. severe). No es were performed, AC and TC study. CT, chemotherapy; HEC, highly	ie, with or without an NK ₋ receptor antagonist. ¹ significant nausea = nausea score 0 and 1 on a 4-pc egimens were classified as MEC. AC has since been emetogenic chemotherapy; 5HT ₃ , 5-hydroxytrypta	 ^bNo nat ^bNo nat ^boint scal ^bn catego ^amine type 	usea = <5 mm oi e (0, none; 1, mild rized as HEC, as h ve 3; MEC, moder	n 100 mr ; 2, mode as carbop ately eme	VAS. «No si ate; 3, sever latin at the dc togenic chem	ignificant naus e). ^f Included ir sse used in the otherapy; NK	sea = <25 a aprepitant e study by Y. c, neurokini	mm on 100 m and fosaprepi ahata et al. ⁸⁰ ^h T in 1; NR, not r	im VAS. ⁴ No tant sections, otal control, eported; NS,

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Discussion

Adding an NK, RA to an antiemetic regimen of a 5HT, RA and dexamethasone significantly reduces the incidence of emesis and rescue medication (as measured by CR) in patients at risk of CINV relative to dual therapy.^{60,62,83} For patients receiving HEC or AC, NK, -RA-containing therapy (as triple therapy or with the addition of olanzapine) is now recommended for CINV prevention in all major antiemetic guidelines.16-19 NCCN guidelines also include this three-drug regimen containing an NK, RA as a recommended option for patients receiving MEC,16 and ASCO guidelines recommend this three-drug regimen for patients receiving MEC that contains high-dose carboplatin.17 Some studies noted a lower CR in female patients than male patients receiving a three-drug antiemetic regimen containing an NK, RA, consistent with female sex being a known risk factor for CINV; however, in all cases CR rates were higher in both male and female patients receiving an NK1-RA-containing three-drug antiemetic regimen compared with a two-drug regimen.

The efficacy of NK₁ RAs in the control of nausea is less clear. Many studies did not measure nausea incidence or severity, and if they did, these were secondary or exploratory end points. Few studies to date have demonstrated a significant improvement in nausea control by adding an NK₁ RA to a two-drug antiemetic regimen, and those that did reported that only about half the patients treated with HEC experienced no nausea in the overall CINV phase. Therefore, there is still a need for more studies evaluating nausea end points and for better antiemetic regimens that improve nausea control.

Currently, NK, RAs are available as both oral and IV formulations. The oral route is convenient, but nonadherence to treatment may negatively affect efficacy. Some patients with cancer cannot tolerate oral treatments, some patients may have difficulty swallowing because of mucositis, and oral drug bioavailability may be compromised by diarrhea or gastrointestinal ulceration.⁹⁶ The IV formulations may be less convenient for patients and hospital staff, as they require patients to attend the clinic,96 but IV administration ensures treatment adherence and is suitable for patients with swallowing difficulties. Following the suspension of distribution of IV rolapitant, there are now three IV formulations of NK, RAs: fosaprepitant, which contains polysorbate 80 and is associated with a high incidence of infusion-site and hypersensitivity reactions;³² an IV aprepitant formulation (HTX019) that is free of polysorbate 80 and other synthetic surfactants, and appears to have an improved tolerability profile;^{33,85} and IV NEPA, free of surfactant emulsifiers and solubility enhancers.29

Among the oral agents, rolapitant has the longest $t_{1/2}$ and requires only a single dose to be administered prior to chemotherapy.³¹ The oral and IV NEPA fixed combinations are also administered only once before chemotherapy, whereas additional doses of oral aprepitant are required on days 2-3 to prevent delayed CINV.^{29,30} However, the long $t_{1/2}$ of oral rolapitant appears to offer no clinical advantage: an indirect meta-analysis of NK, RAs suggested that oral rolapitant was the least effective available agent in this class.⁶⁰ All NK₁ RAs may be associated with potential drug interactions.²⁹⁻³³ Those that are CYP3A4 substrates (aprepitant, fosaprepitant, and NEPA) should be given with a lower dose of dexamethasone on treatment day 1 (12 mg) than the dose used with oral rolapitant (20 mg).¹⁶ When deciding which NK, RA to use, physicians should consider the formulation, indication, pharmacology, efficacy, and safety of these agents, as well as any concomitant medications. It has been suggested that individualized antiemetic therapy, taking into account both treatment-related and patient-related risk factors, may be preferable to consensus guidelines, and patient-level CINV-predictive models have been proposed.97 In addition, NCCN guidelines recognize that the ultimate clinical decision on an appropriate antiemetic regimen may depend on the individual patient's situation and risk factors.16

Despite their clear benefits and recommendations in antiemetic guidelines, NK, RAs are underutilized in clinical practice. Some institutions may limit the use of more expensive branded antiemetics by asking physicians to use a 5HT, RA and dexamethasone in the first cycle, then add an NK, RA in later cycles if the patient experiences CINV in cycle 1. This practice is inconsistent with antiemetic guidelines for patients receiving HEC (and many receiving MEC), and ignores the fact that the patient's first experience with chemotherapy is most crucial for CINV prevention.²⁰ Patients whose CINV is controlled in the first cycle are more likely to do well in subsequent cycles, whereas patients who experience CINV during cycle 1 are more likely to develop refractory or anticipatory CINV.73,98 Patients who do not achieve complete CINV control have poor quality of life, incur greater costs, and use more health-care resources.^{8,90,99} One option that has been considered is the use of olanzapine instead of an NK, RA in combination with a 5HT₃ RA and dexamethasone. A randomized Phase III trial of olanzapine compared with oral aprepitant, both in combination with IV palonosetron and dexamethasone, in patients receiving cisplatin-based or AC-based HEC found no significant difference in CR rates between the two regimens,¹⁰⁰ but there was a significant

improvement in the control of nausea with olanzapine. In a meta-analysis of ten randomized controlled trials, olanzapine was more effective than oral aprepitant in the acute phase of CINV, but comparable in the delayed phase.¹⁰¹ A meta-analysis of 43 trials reported that an olanzapine-based triplet regimen improved nausea control, but was similar in CR to an NK₁-triplet regimen.¹⁰² There are strong economic and clinical arguments for the use of guideline-recommended antiemetic protocols that include an NK₁ RA in addition to a 5HT₃ RA and dexamethasone (with or without olanzapine) during the first and subsequent chemotherapy cycles. This is especially important as hospitals transition to reimbursement for quality care rather than fees for service, and oncologists will be encouraged to keep patients out of the emergency department and hospital.

In conclusion, this review of published data with NK, RAs highlights the efficacy of these agents in controlling emesis and rescue-medication use as part of three-drug or four-drug regimens, and the importance of patients receiving prophylactic regimens that comply with antiemetic guideline recommendations. For nausea control, the incremental benefit of using an NK, RA is less clear, so this remains an area for future research. While caution is needed in making cross-study comparisons, the available data suggest that the pharmacological differences between the NK₁-RA inhibitors, specifically the longer $t_{i_{\lambda}}$ of oral rolapitant, do not translate into enhanced clinical benefit, particularly within the HEC setting. Newer agents may offer key advantages in terms of better nausea control, tolerability, formulation options, and therapeutic plasma levels in the acute phase of CINV than the existing agents, and offer clinicians more opportunities to maximize the benefits of this important class of antiemetics.

Author contribution

LSS and RMN designed the systematic review, were responsible for the writing and critical revisions of the manuscript, read and approved the final manuscript, and agree to be accountable for all aspects of the work.

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