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FOSAPREPITANT - fosaprepitant injection, powder, lyophilized, for solution MSN LABORATORIES PRIVATE LIMITED
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INCHRIGHTS OF PRECEDENG INFORMATION

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INCREDIBLE AND ADMINISTRATION OF THE PROPERTY OF THE

prevention of (1):

- scale and delayed nausea and vomiting associated with initial and repeat courses of highly emetogenic cancer chemotherapy (HEC) including high-dose clapitatis.

- calculated nausea and uniting associated with initial and repeat courses of moderately emetogenic cancer chemotherapy (HEC).

<u>Limitations of like (1)</u>
• Foragrepitant for injection has not been studied for treatment of established nausea and somiting.

**Recognitional Section 1. The control of the contr

cays 2 and 2.

Administer focuspreplant for injection through a central venous catheter as an intravenous infusion over 30 minutes (32 years to 17 years) or 60 minutes (6 months to less than 12 years).

Complete the infusion approximately 30 minutes prior to chemotheraps:

Complete the Indian September | 20 milutes plant to Chemotherapy
Commission Assessment

See And Presentable (Hornocolous the Confidence Information, D.1.2)

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COMMANDACTIONS

Notice Information Commission

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Most common adverse reaction in adults (a 2H) are: Mague, Gardhaa, neutropenia, authenia, anexis, penipheral neuropathy, inskapenia, dystepsia, unitary tract infection, pain in extremity. (6.1)
 Adverse reactions in pediatrics are similar to adults.

To report SUSPECTED ADVERSE REACTIONS, contact MSN Pharmaceuticals Inc. at 1-855-668-2569 or FDA at 1-808-0088 or

DRIS INTERACTIONS

See Full Prescribing Information for a list of clinically algorithms drug interaction. (4, 5.1,5.4, 7.1, 7.2)

Additional pediatric use information is approved for Merck Sharp & Dohme LLC's EMBLD (bicaprepliant) for injection vicesveer, due to Merck Sharp & Dohme LLC's marketing exclusivity rights, this drug product is not labeled with that information.

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See 17 for PATIENT COUNSELING INFORMATION.

Revised: 10/2022

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8.1 Pregnancy
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1 INDICATIONS AND USAGE Freapreplant for rigidize, its ordered and a state of the prevention of the prevention of the prevention of the prevention of a contract of the prevention of the prevention of a contract of the prevention of the prevent

emetogenic cancer chemotherapy (MEC). Limitations of Use.

• Fosapreptant for injection has not been studied for the treatment of established nausea and vomiting.

→ ✓ → APMINISTRATION

2.1 Prevention of Nausea and Vomiting Associated with HEC and MEC in Adult Patients

Adult Patients

The recommended dosage of fosagreplant for injection, decamethisone, and a 5HT, artisppint for the prevention of nauses and somiting associated with administrate of HTC or MEC in addition, and with a first properties of HTC or MEC in addition, and indicate fosagreplant for rejection as an intravenous infusion on Day 1 over 20 to 30 minutes completing the inflation appearatisately a first-innexts, parts or thermotherapy.

Workling Associated with HTC.

	Day 1	Day 2	Day 3	Day 4	
fosaprepitant for rjection	150 mg intravenously over 20 to 30 minutes	none	none	none	
Dexamethasone*	12 mg orally	8 mg orally	d mg orally twice daily	5 mg orally twice daily	
5-HT ₂ antagonist	See selected 5- HT ₂ antagonist prescribing information for the recommended	none	none	none	

*Administer decarmethations 30 minutes prior to chemisther apy treatment on Day 1 and in the morning on Days 2 through 4. Also administer decarmethations in the overlags on Days 3 and 4. A 50% dissage reduction of decarme

Table 2 Recommended Adult Dosing for the Prevention of Nausea and Vomiting Associated with MEC

	Day 1
rosaprepitant for niection	150 mg intravenously over 20 to 30 minutes
Dexamethasione*	12 mg orally

*Administer decamethisaces 30 minutes prior to chemotherapy treatment on Day 1. A 50% dosage reduction of decamethisaces is recommended to account for a drug interaction with inapropletant for lyacino, in sec China Pharmacology (22.3)].

2.2 Prevention of Manaea and Vomiting Associated with MEC and MEC in Preduct Publishers.

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1 moré 6.

Administra fossar epitant for hipetion on Day 1 over 30 minutes (12 years to 17 years) or 50 minutes (12 years to 17 years) or 50 minutes (8 morths to less than 12 years), completing the inhalon approximately 50 minutes prior to Administra prior to Administration (17 minutes) or 50 minutes prior to Administration (17 minutes) or 50 minutes prior to Administration (17 minutes) or 50 minutes prior to Administration (18 minutes) or 50 minutes prior to Administration (18 minutes) or 50 minutes (18 minutes) or 50 minutes (18 minutes) or 50 minutes) or 50 minutes (18 minutes) or 50 minutes) or 50 minutes (18 minutes) or 50 minutes (18 minutes) or 50 minutes) or 50 minutes (18 minutes) or 50 minutes) or 50 minutes) or 50 minutes (18 minutes) or 50 minutes)

Drug	Age	Regimen
Posapreptant for injection	12 Years to 17 Years	150 mg intravenously over 30 minutes
	2 Years to less than 12 Years	4 mg/kg (maximum dose 150 mg) intravenously over 60 minutes
	6 Months to less than 2 Years	5 mg/kg (maximum dose 150 mg) intravenously over 60 minutes
Dexamethasonet	6 Months to 17 Years	If a corticosteroid, such as dexamethasone is co-administered, administer 50% of the recommended corticosteroid dose on Days 1 and 2.
5-HT ₂ antagonist	6 Months to 17 Years	See selected 5-HT3 antagonist prescribing information for the recommended dosage

* Dosing in pediatric patients less than 6 kg is not recommended

† Administer decumethasone 30 minutes prior to chemotherapy treatment on Day 1

Fosiacreptient for injection Dosage Regimen for Use with Multi-Day Chemotherapy,
Regimens.

Constitution

For politicity patients weighing at least 6 kg neceiving multi-day regimens of HEC or MEC, administed fosspreplature. For synction on Days 1.2, was 3. Administer fosspreplature for injection as an infrarenous institution frought a certain/avenus cultivater on Days 1 and apreplature capsular on for insuspreplature for principlature on Days 2 and 3, as shown in Administer capsular of the principlature of the principlature on Days 2 and 3, as shown in Administer for synchron or positive or principlature on Days 2 and 3, as shown in Administer for synchron or positive or principlature on Days 2 and 3, as shown in Administer for synchron or positive or for suspension on Days 2 and 3, as shown in Administer for synchron or for suspension on Days 2 and 3, as shown in Days 2 and 3, as shown in Days 3 and 3, and

MIRIZER (6 10 TOTALes was member app. The miritals prior to chemichler app. The miritals prior to chemichler app. The miritals prior to chemichler app. The miritals prior to chemicals and to chemical appears and to chemical associated with Single or Multi-day Regimens of HEC or MEC in Pediatric Patients 6 Months* to 17 Years

	Age of Pediatric Population	Day 1	Day 2	Day 3
Fosaprepitant for injection*	12 years to 17 years	115 mg intrivenously over 30 minutes	80 mg orally (apreptiant capsules) [†]	80 mg orally (apreptiant capsules) [†]
	6 months to less than 12 years		2 mg/kg orally (Fosaprepitant for oral suspension)F	2 mg/kg orally (Fosiaprepitant for oral suspension) ²

		22.71190	80 mg)	(maximum dose 80 mg)				
Dexamethasone	17 years	If a corticosteroid, such as dexamethasone, is co- administered, administer 50% of the recommended corticosteroid dose on Days 1 through 4.						
5-HT3 antagonist	6 months to 17 years		T3 antagonist presc e recommended do					

* Dosing in pediatric patients less than 6 kg is not recommended.
Ter patient 12 years to 17 years untable to washer or at capsules, fosspreighant for **Ter patient 12 years of age who weight a less 40 kg and who are able to washer or at capsules, apropher capsules can be used instead on Days 2 and 3 **Asternizate describeduses can be used instead on Days 2 and 3 **Asternizate describeduses 20 **Political port to characterizate preferred capsules can be used instead on Days 2 and 3 **Asternizate describeduses 20 **Political port to characterizate preferred can Day 1.

Additional pediatric use information is approved for Merck Sharp & Dohme LLC's EMENE (fosipreplant) for injection. However, due to Merck Sharp & Dohme LLC's marketing exclusivity rights, this drug product is not labeled with that information.

2.3 Preparation of Fosaprepitant for injection

Step 1	Assiptically ispect 5 mt. 0.9% Sodium Chloride injection, USP into the voil kissure that 0.9% Sodium Chloride injection, USP is added to the rial along the visit wall in order to prevent fearning. Swirt the visit quest kvoid shaking and jetting 0.9% Sodium Chloride Injection, USP into the Visit.
Step 2	Aseptically prepare an infusion bag filled with 145 mL of 1.9% Sodium Chibride Injection, USP.
Step 3	Asspitially withdraw the entire volume from the vial and transfer not the infusion bug centaining 145 mt. of 0.9% Sodium Chibrid nijection, USP to yield a total volume of 150 mtLand a fin concentration of 1 mg/mL.
Step 4	Gently invert the bag 2 to 3 times.
Step 5	Observation the solution to the advantage from the gregard offshoot 2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.
Step 6	If necessary, for volumes less than 150 ms, the calculated volume cape transferred to an appropriate size bag or syringe prior teatministration by infusion.
Step 7	Before administration, inspect the bag for particulate matter and discoloration. Discard the bag if particulate and/or discoloration are observed.

The recommended date of finage-replane for rigidition is based on the patient's age and sergice.

Coulomb Do not mice or reconstitute finage-replane for injection with seathers for which is formed to the contraction of the

3 DOSAGE FORMS AND STRENGTHS
Fosapreptiant for injection: 150 mg fosapreptiant, white to off white cake or powder in single-dose glass via for reconstitution.

CONTRAINDICATIONS
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See Table 7 and Table 8 for a listing of potentially significant drug interactions (see Drug interactions (7-1,7-2)).

3.2 hypersemblely feaction, including anaphylaxis and anaphylactic shock, during or soon after enhusion of loagerplant have occurred. Symptoms excluding lashing, expleme, appears, hypersen, hypersenshild year facilities and deminister perspective medical featings, to not reinstitute footspread and underside perspective medical featings, to not reinstitute footspreadured in palents with opportunities of the perspective medical featings, to not reinstitute footspreadured in palents with opportunities of the perspective medical featings, to not reinstitute footspreadured in palents with opportunities of the perspective medical featings, to not reinstitute footspreadured in palents with opportunities of the perspective medical featings, to not reinstitute footspreadured in palents and opportunities of the perspective medical featings, to not reinstitute footspreadured in palents and palents and

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5.4 Discrease in IMR with Concombant Warfarin
Caudinnistration of Tosapreplant
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Chicial Pharmacology (22.3)
Monte the IMR is planted on chronic warfare threapy in the 2week prints, particularly at 7 to 10 days, following initiation of fosapreplant
with each Chicial Chicago you'de law Cong foraction (7-12).

was neare, remnomarily cycle late Oray Information (7.1);

2.3. Bits of Induced Plattacy of Hormand Contraceptions
Upon condeministration with freesprepture, the efficacy of hormand contraception
may be returned unique administration of an ext. 2 days foliosing the last does of
attribute or facts by methods of contraception during treatment, with freesprepture
and for 1 month following administration of freesprepture (pure
Use in Specific Populations (6.31);

6 ADVERSE REACTIONS
The following clinically significant adverse reactions are described elsewhere in the ADVENSE REALTIONS
 The following clinically significant adverse reactions are described else labeling:
 Hypersensibity Reactions (see Warnings and Precautions (5.29)
 Infusion Site Reactions (see Warnings and Precautions (5.3))
 C.1 Clinical Trials Experience

Bocause clinical trials are conducted under reliefly varying conditions, adverse reaction rales observed in the clinical trials of a drug clientot be directly compared to raines in the clinical trials of a drug clientot be directly compared to of another drugs and may not relief bethe present on clinical practice. The overal safety of losspreptant for spectron was evaluated in approximately 1800 adult and potentine potents.

adult are present peaces. Associated with the Proportion of Managa and Victoritys Associated with degrees absociation in Adults for the Proportion of Managa and Victoritys Associated with In an active-centrelated chical trial in patients receiving MECs, safety was evaluated in 304 patients receiving a stage does of insepreptiant for repiction in combination with ordinated and an adult of the patients of the patients of the patients of the patients of patients receiving a language of the patients of the patients of the patients of patients of the patients of the patients of the patients of the patients of Table 6 Most Common Adverse Reactions in Patients Receiving MEC*

	Fosaprepitant for injection, ondansetron, and dexamethasone! (N=504)	Ondansetron and dexamethasone ¹ (N=497)
atique	15%	13%
darrhea	13%	11%
neutropenia	8%	7%
astherna	4%	3%
anemia	3%	2%
peripheral neuropathy	3%	2%
eukopenia	2%	1%
Syspepsia	2%	1%
urinary tract infection	2%	1%
oan in extremity		

Gen in microsides 25 May 15 Ma

DEC. In an action controlled clinical study is patients receiving INCL safety was evaluated for the action controlled clinical study is patients receiving INCL safety was evaluated for patients receiving the 3-day regimen of oral proplator. Inter Chical Studies (2.4.1). The patients receiving INCL safety in Studies is the Studies of the INCL study with long patient and except point in the Studies of the INCL studies of

0.5%. 3.1%, reported in the Inscriptorate group companed to the surseplant group conceptual, proceedings, and process of the Inscriptorate group conceptual, process of the Inscriptorate group conceptual, process of the Inscriptorate group conceptual group conce

protein in usuli, patento receivorgi a single consi or trosuperplant the greater is consisted with appeals may be a consisted or core with foreign the single consistence of the process of the consistence of the consistence

6.2 Postmarketing Experience

The following deview mactions have been identified during post-approval use of fossprephent, Decause these reactions are reported volarizing from a population of fossprephent, Decause these reactions are reported volarizing from a population of the fossprephent, and a product of the population of th

7 DRUG INTERACTIONS 7.1 Effect of Fosaprepitan Drugs

When administered introvermously, foliagreplate, a producy of sereplate, it converted to aproplate within 30 minutes. Therefore, drug retractions following administration of learning plates of the roles and produced and drug plates and the roles plate street with one plane plates and the role aproplate, weak inhabition of CPSA4 continues for 2 days after simple does administration. Steply seen followed to the role of CPSA4 continues for 2 days after simple does administration. Steply are independent of CPSA4, Aproplate in a substance in ministration of CPSA4, Aproplate in substance of CPSA4, Aproplate in substance of CPSA4, are contracted used in followed planes are consistent of CPSA4 are contracted used in followed planes.

bistrates of CYP3A4 are contraindicated with fosaprepliant [see dications (4)]. Dosage adjustment of some CYP3A4 and CYP2C9 substrates may inted, as shown in Table 7.

CYP3A4 Substi	
Pimozide	mes .
Clinical Impact	ncreased pimozide exposure
intervention	Posapreobant is contraindicated (see Contraindications (4)).
Servanton Servadiazeoines	POSABTRIDEARS IN CONTRARIDEARRO ISSNE CONTRARIDEARRORS (47).
Clinical Impact	increased exposure to midazolam or other benzodiazepines
Larrical Impact	metabolized via CYP3A4 (alprazolam triazolam) may increase the risi
	of adverse reactions (see Clinical Pharmacology (12,31).
	Monitor for benzodiazepine-related adverse reactions.
intervention Decamethasione	Monitor for benzodazepine-related adverse reactions.
Clinical Impact	increased decamethasone exposure/see Clinical Pharmacolog
ofenuention	(12.3)].
Intervention	Reduce the dose of oral dexamethasone by approximately 50% (see
	Dosage and Administration (2.1)].
Methybrednisolo	ne
Cârsical Impact	increased methylprednisolone exposure (see Clinical Pharmacolog
	(12.3)J.
ntervention	Reduce the dose of oral methylprednisolone by approximately 50%
	on Days 1 and 2 for patients receiving HEC and on Day 1 for patients
	receiving MEC.
	Reduce the dose of intravenous methylpredrisolone by 25% on Days
	1 and 2 for patients receiving HEC and on Day 1 for patients receiving
	MEC.
Chemotherapeut	ic agents that are metabolized by CYP3A4
Clinical Impact	Increased exposure of the chemotherapeutic agent may increase the
	risk of adverse reactions (see Clinical Pharmacology (12.3)).
ntervention	Architecture, vincristine, or ifosfamide or other chemotherapeut
	 Monitor for chemotherapeutic-related adverse reactions.
	Stoposide vinorebine pacitized and docetaxel
	 No dosage adjustment needed.
Hormonal Contra	centiasi
Clinical Impact	Decreased hormonal exposure during administration of and for 28
	days after administration of the last dose of fosapreplant (see
	Warnings and Precautions (5.5), Use in Specific Populations (8.3), an
	Clinical Pharmacology (12.3)).
ntervention	Effective alternative or back-up methods of contraception (such as
THE PERILED	condoms and spermicides) should be used during treatment with
	osapreptant and for 1 month following administration of
	osapreptant.
Corporation:	
Examples	orth control pills, skin patches, implants, and certain IUDs
Examples	
	birth control pills, skin patches, implants, and certain IUDs
CYP2C9 Subst	birth control pills, skin patches, implants, and certain IUDs
CYP2C9 Substi	birth centrel pills, skin patches, implants, and certain IUDs
CYP2C9 Subst	birth control pils, skin patches, implants, and certain IUDs autes Decreased warfarin exposure and decreased prothrombin time (INR)
CYP2C9 Substi Warfarin Clinical Impact	inth control pile, skin patches, implents, and certain IUDs units. Decreased werfarin exposure and decreased prothrombin time UNIX See Warnings and Precasions (5.4). Clinical Pharmacology (12.3).
CYP2C9 Substi	birth control pills, skin patches, implants, and certain IUDs wites Decreased warfarin exposure and decreased prothromain time (INR) Sae Warnings and Procusions (5.4). Clinical Pharmacology (22.3)]. n patients on Promis warfar Interior year new the prothromain time n patients on Promis warfar Interior year memory the prothromain.
CYP2C9 Substi Warfarin Clinical Impact	arth control pills, skin patches, implients, and certain IUDs sites Decreased warfarin appoint and decreased proteoministins (RIM) Decreased warfarin appoint and decreased proteoministins (RIM) In palliants on chronic warfain their apy, montrol the proteoministins in palliants on chronic warfain their apy, montrol the proteoministic research.
CYP2C9 Substi Warfarin Clinical Impact	birth control pills, skin patches, implants, and certain IUDs wites Decreased warfarin exposure and decreased prothromain time (INR) Sae Warnings and Procusions (5.4). Clinical Pharmacology (22.3)]. n patients on Promis warfar Interior year new the prothromain time n patients on Promis warfar Interior year memory the prothromain.
CYP2C9 Substi Warfarin Clinical Impact	arth control pills, skin patches, implients, and certain IUDs sites Decreased warfarin appoint and decreased proteoministins (RIM) Decreased warfarin appoint and decreased proteoministins (RIM) In palliants on chronic warfain their apy, montrol the proteoministins in palliants on chronic warfain their apy, montrol the proteoministic research.
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EYP2C9 Substi Warfarin Clinical Impact Intervention	bith control pile, skin publisher, replaces, and certain 1925s 2688. Discussed warfare outpours and discussed profesorible time 1986 saw Warming and Pre-easities (2.4), Chical Pharmocology (22.3); n patient, on thorn warfar therapy, recent the profesorible and patient and therapy and pre-easities (2.4), Chical Pharmocology (22.3); n patient on thorn warfar therapy, recent the profesorible safernitar atom of fosagraphant with each chemotherapy cycle.
CYP2C9 Substi Warfarin Clinical Impact Intervention	Diffs control pile, skin publisher, implients, and certain ISUSs MEAN TOTAL STATE AND ADDRESS AND AD
EYP2C9 Substi Warfarin Clinical Impact Intervention	print contrargibs, sten patients, requires, and center IDSs. 284 Consequent and feet mappingues and decreased protriormals since IDSs. Consequent and feet mappingues and decreased protriormals since IDSs. Consequent and the contract of the contract o
CYP2C9 Substi Warfarin Clinical Impact Intervention	port contrargibs, his packness, registers, and certain follows: **Commission and pack registers and an article programmes from the commission and
CYP2C9 Substi Warfarin Clinical Impact Intervention	print contrargibs, sten patients, requires, and center IDSs. 284 Consequent and feet mappingues and decreased protriormals since IDSs. Consequent and feet mappingues and decreased protriormals since IDSs. Consequent and the contract of the contract o

7.2 Effect of Other Drugs on the Pharmacokinetics of Fosaprepitant/Aprepitant

Aprephant is a CYP344 substrate (see Clinical Pharmacology (12.3)). Co-administration of foogreeplant with drops that are inhibitors or induces of CYP344 may result in concentrations of aprephant, respectively, as shren in Table 8.

Table 8 Effects of Other Drugs on Pharmacokinetics of

 -	u Lines	Fos				·	

Clinical Impact	Significantly increased exposure of apreptiant may increase the risk of adverse reactions associated with fosapreptiant [see Adverse Reactions (6.1), Clinical Pharmacology (12.3)].
intervention	Avoid concomitant use of fosapreplant
Examples	Moderate inhibitor: Sklarem Sk
Strong CYP3/	4 Inducers
Clinical Impact	Substantially decreased exposure of apreptiant in patients chronically taking a strong CYP3A4 inducer may decrease the efficacy of fosapreptiant (see Clinical Pharmacology (12.3)).
Intervention	Avoid concomitant use of fosapreplant.

a USE IN SPECIFIC POPULATIONS

2.1 Prepanency
3.8 Marriages
3.4 Marriages
4.5 Marriage

risk of major birth defects and miscarriage in clinically recognized pregnancies is 2 to 4% and 15 to 20%, respectively

Data Animal Data

communities in enthinguised development studies in rats and rabbits, aproplant was administered in enthinguised development at crad does up to 1000 might bette daily rats) and up to the maximum toler lated days are 23 mightights of publishes. To enthinguishes also might great daily represent and up to the communities and days are 23 mightights of publishes. To enthinguishes are might great and are publishes to enthinguishes and are publishes of the publishes are daily and in the daily are da

property of solidor at 2 mg/splay ware several and a solidor and a solidor at 2 mg/splay ware several and a solidor at 2 mg/splay and a solido

The safety and effectiveness of a single dose and a 3-day regimen of fosaprepitant have been established in pediatric patients 6 months to 17 years for the prevention of acute and delayed nausea and vomiting associated with initial and repeat courses of MEC and MEC.

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10 OVERODAGE
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Insoluble in r-bezarie.

Each vila of losspreplant for irjection for administration as an intravenous infusion contains 245.3 mg of fosspreplant direquismine equivalent to 50 mg of fosspreplant direquismine equivalent to 50 mg of fosspreplant fire act and the following native in preferents: debate disodium 15.4 mg), lactose arrhydrous (375 mg), polysorbate 80 (75 mg), sodium hydrochekie action flygrenches action (for pri adjalantem).

12.1 Mechanism of Action
Fosapreptiant is a prodrug of aprepliant and accordingly, its antiemetic effects are attributable to apreptiant.

Aprephant is a selective high-affinity antagenist of human substance P/heurokinin 1(NK₂) receptors. Aprephant has little or no affinity for serotonin (5-HT₂), dopamine, and corticosteroid

reciptors, the targets of existing therapies for chemotherapy-induced massins and vomiting CRVID, Aprengation that been shown in a mirral model or in this control of the c

12.2 Pharmacodynamics
Cacilia: Electrophysiolog:
In a randomied, double-blind, positive-controlled, thorough QTc study, a single 200mg dose of Fosspertal (approximately -13 times the recommended dose) had no effect on the QTc interval. 12.3 Pharmacokinetics

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Jack Pharmacetische (Administration)

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Table 9 Systemic Exposures of Aprepitant for Single Dose Fosaprepitant for Injection Regimen in Pediatric Patients

Population	Single Dose of	n				
	fos aprepitant for Injection Regimen	AUC 0-26hr- (mcg*hr/mL)	C _{max} (mcg/mL)	C ₂₄ (mcg/mL)	C ₆₂ (mcg/mL)	C ₇₂ (mcg/mL)
12 Years to 17 Years	150 mg	29.4	3.4	0.7	ND*	ND*
6 Years to less than 12 Years	4 mg/kg	35.2	3.6	0.7	0.2	0.05
2 Years to less than 6 Years		28.2	3.1	0.4	0.1	0.02
6 Months to less than 2 Years	5 mg/kg	32.7	3.3	0.4	NET	ND*

No — Not Determined. Pharmacolisatic samples seem on collected to support the parameter value of infraset.

Ne — Not Extinsized, The generatic mann could not be estimated due to values being believe the limitation of quantification.

3-day NOT/OneONOT for superspictual Regimes: Structural and greeplast systemic exposures in partners 6 months to set than 1.2 years and observed systemic exposures in partners. Sometimes to contain to set than 1.2 years and observed systemic exposures in partners (Compl. on Day 1 and concentrations at the end of Day 1 (Cyst.), Day 2 (Cya) and Day 3 (Cyst.).

(Cop):
Table 10
Systemic Exposures of Aprepitant for 3-Day IV/Oral/Oral Regimen in Pediatric

Population	3-Day Dose	Geometric Mean						
	Fosaprepitant (IV)Oral/Oral*)	AUC 0-24hr- (mcg*hr/mL	C _{max} (mcg/mL)	C ₂₄ (mcg/mL)	(mcg/mL)	C ₇₂ (mcg/mL		
12 Years to 17 Years	115/80/80 mg	18.0	3.0	0.4	0.2	NET		
6 Years to less than 12 Years	3/2/2 mg/kg	25.7	2.7	0.5	0.3	0.3		
2 Years to less than 6 Years		20.2	2.3	0.3	0.2	0.2		
6 Months to less than 2 Years		16.6	1.9	0.2	0.1	0.1		

*W on Day 1, Oral on Day 2, and Oral on Day 3

*NE — Not Estimated. The permetric mean could not be estimated due to values being below the limitation of quantification.

Plasma concentrations of losspreptiant are negligible within 15 – 30 minutes after the completion of the intuison in polistic polisions.

Comparison of the relation to partners; patients.

See

1. See

Control and Contro

Additional pediatric use information is approved for Merck Sharp & Dohme LLC's EMEND (fosiapreptiant) for injection. However, due to Merck Sharp & Dohme LLC's marketing exclusivity rights, this drug product is not labeled with that information.

an Indicational Conference on Assessment of Particles

13.1 Contribution Conference on Assessment of Particles

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Mutagenesis. Aprepitant and fosiaprepitant were not genotoxic in the Ames test, the human lymphobiastoid cell (TX6)

mutagenesis test, the rat hepatocyte DNA strand break test, the Chinese hamster ovary (CHO) cell chromosome aberration test and the mouse micronucleus test. Impairment of fertility for the control of t

recognition, with animomorphic and expenses of a popular content of a perspect, in the condition of a condition

CLINICAL STORES

14.1 Proceedings of Received Services and Varieting Associated with REC in Adults

14.2 Proceedings of Received Services and Varieting Associated with REC in Adults

14.2 Proceedings of Received Services and Varieting Associated Services (Fig. 1971) and compared to a 3
day of services (Fig. 1971) and received Services (Fig. 1971) and Compared to a 3
day of services (Fig. 1971) and received Services (Fig. 1971) and Compared to a 3
day of services (Fig. 1971) and received Services (Fig. 1971) and Compared to a 3
day of services (Fig. 1971) and Services (Fig. 1971)

Table 12 Treatment Regimens in Adult HEC Trial*				
	Day 1	Day 2	Day 3	Day 4
Fosaprephart Regi	men			
Fosapreplant for injection	150 mg intravenously over 20 to 30 minutes approximately 30 minutes prior to chemother apy	none	none	none
Oral dexamethasone ²	12 mg	8 mg	8 mg twice daily	8 mg twice daily
Ondansetron	Ondansetron ²	none	none	none
Oral aprepitant Re	gimen			
Aprepitant capsules	125 mg	80 mg	80 mg	none
Or al dexameth as on	d 12 mg	8 mg	8 mg	8 mg
Ondarsetron	Ondansetroni	none	none	none

*Temperapation for implicition placeline, appropriate cognision placeline and data semichimistree placeline for in eventure, and Supp. 3 of a five several to make the final semiconal production of the country of the

Table 13 Percent of Adult Patients Receiving HEC Responding by Treatment Group and Phase — Cycle 1

ENDPOINTS	Fosaprepitant for Injection Regimen (N = 1106)*	Oral Aprepitant Regimen (N = 1134)*	(95% CI)
PRIMARY ENDPO	NT		
Complete Respon	nse ^p		
Overall ⁶	71.9	72.3	-0.4 (- 4.1, 3.3)
ECONDARY END	POINTS		
Complete Respon	se ²		
Delayed phase!	74.3	74.2	3.5, 3.7)
No Vomiting			
Overaff	72.9	74.6	-1.7 (- 5.3, 2.0)

"N: Namber of patients included in the primary analysis of complete response. TDRFerence and Confidence interval (CI) were calculated using the method proposed by "Complete Response — no vomething and no use of rescue therapy. "Coveral — to to 120 hours post-initiation of cisplain chemotherapy. "Delayed phase — 25 to 120 hours post-initiation of cisplain chemotherapy.

14.2 Prevention of Nausea and Vomiting Associated with MEC in Adults

In a randomized, gurdak daduk-bödi, achte comparator controlled study, in a randomized, gurdak daduk-bödi, achte comparator controlled study, in consideration with collecturation and scannidations to liceage particle direquipment (consideration with collecturation) and controlled study of the controlled study of the

(51%), oxalplatin (24%), and cyclophosphamide (Table 14 Treatment Regimens in Adult MEC Trial*

	Day 1	Day 2	Day 3
osaprepitant Regimen			
osapreptant for Injection	150 mg intravenously over 20 to 30 minutes approximately 30 minutes prior to chemotherapy	none	none
Draf Dexamethasone ⁷	12 mg	none	none
Oral Ondansetron [‡] Standard Therapy	8 mg for 2 doses	none	none
Oral Dexamethasone	20 mg	none	none
Oral Ondansetron [‡]	8 mg for 2 doses	8 mg twice	8 mg twice daily

* Fossproylant for rigition placebo and decamethaction for the production of the pro

The primary endpoint was complete response (defined as no vomiting and no rescue therapy) in the delayed phase (25 to 120 hours) of chemotherapy-induced nausea and vomitino. The results by the restiment or occup are shown in Table 13.

Table 15 Percent of Adult Patients Receiving MEC Responding by Treatment Group

ENDPOINTS	Fosaprepitant for Injection Regimen (N = 502)*	P-Value	Treatment Difference (95% CI)
PRIMARY ENDPOINT			
Complete tesponse ²			
			10.4 (5.1, 15.9)

*N: Number of patients included in the intention to treat population.
*Complete Response = no vomiting and no use of rescue therapy.

*Delayed phase = 25 to 120 hours post-initiation of chemotherapy.

16 HOW SUPPLIED/STORAGE AND HANDLING
No. 151 — Single-dose glass vial containing 150 mg of fesspreptiant as a while to off
white hoppiblace due to prowder for reconstitution. Sopplied as feditives:
Single-State of the State of S

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2 PATIST COURSELING POTENTIAL SHADOWS AND ASSESSED AS

	Fosaprepitant
١	for injection

Road this Patient information before you start receiving fosapreptant for injection and each time you are scheduled to re-coagreptant for injection. There may be new information. This information does not take the place of taking with your healthcare provider about your medical condition or treatment.

What is fosaperuptant for hijection? osaperuptant for hijection is a prescription made the used with other medicines that treat neuroes and control of the properties of the properties of the prevent neuroes and vomiting cased by certain arti-curact cit-termetherapy imedicines. Fosaperuptant for injection is not used to treat neuroes and vomiting that you aready how. It is not brown if the properties for ejection is a size and effection in critisms have form of morethin of age.

Who should not recain financepitant for high tion?

Do not notice foragraphism for injection?

Do not notice foragraphism for injection if your

are always to foragraphism, represent, or uny of the injections in foragraphism for injection. See the

end of this sulfet for a complete six of the injections in foragraphism for injection.

are usually parsions (GMPA*).

It should I tell my healthcare provider before receiving fosaprepitant for injection? re receiving fosaprepitant for injection, tell your healthcare provider if you: we low problems preparate or plan to become pregnant. It is not known if fosaprepitant for injection can harm

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Allow the instict this you see, samp as no treem our strong your neutricare provises or prammaca, want 
god a new middler.

How we'll bracebee footaprepitant for injection?

Modella 19 years of age and older:

rosuprepitant for injection will be given on Day 1 of chemotherapy treatment. It will be given to you by

rosuprepitant for injection will be given on Day 1 of chemotherapy treatment. It will be given to you by

rosuprepitant provinces in your value about 50 to 60 minutes before you start your chemotherapy
                                                 terent.

dren 6 months to 17 years of age:
apreparatifor rigiction will be given to your child by intravenous (IV) infusion into a large vein through a
of IV line called a central venous catheter, about 1 hour to 1 % hours before the start of their
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el your healthcare provider if you have any side effect that bothers you or that does not go away. These re not all of the possible side effects of foispreptant for injection. For more information ask your subthcare provider or pharmacist. All your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-DDB.

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ional pediatric use information is approved for Merck Sharp & Dohme LLC's EMEND (fossprephant) for on. However, due to Merck Sharp & Dohme LLC's marketing exclusively rights, this drup product is no id with this information. Market Information has been approved by the U.S. Food and Drug Administration.

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