CINVANTI® (aprepitant) injectable emulsion Coding Reference Guide

ICD-10 Diagnosis Code		
Code	Description	
R11.0	Nausea	
R11.10	Vomiting, unspecified	
R11.11	Vomiting without nausea	
R11.12	Projectile vomiting	
R11.13	Vomiting of fecal matter	
R11.14	Bilious vomiting	
R11.2	Nausea with vomiting, unspecified	
T45.1X5*	Adverse effects of antineoplastic and immunosuppressive drugs	
Z41.9*	Encounter for other procedures for purposes other than remedying health state	
Z51.11* [†]	Encounter for antineoplastic chemotherapy	

*Supplementary Classification Code TRequired when given within 48 hours of moderately or highly emetogenic chemotherapy

National Drug Code (NDC)	
NDC	Description
47426-0201-01	Single-dose vial 130 mg IV

Coding for CINVANTI	
HCPCS Code	Description
J0185 [‡]	Injection, aprepitant, 1 mg

[‡]This code may be used for CINVANTI administered on or after January 1, 2019.

Modifier	Description
JW Modifier	Drug amount discarded/not administered to any patient (Indicate quantity discarded)
JZ Modifier§	Zero drug amount discarded/not administered to any patient
JG Modifier	Modifier for drug or biological acquired with 340B drug pricing program discount
TB Modifier	Modifier for drug or biological acquired with 340B drug pricing program discount; reported for informational purposes

Professional Services and CPT Codes	
CPT Code	Description
96367	Intravenous infusion, for therapy, prophylaxis, or diagnosis; additional sequential infusion of a new drug/substance, up to 1 hour
96375	Therapeutic, prophylactic or diagnostic IV push, new substance/drug

Hospital Service and Supplies		
Revenue Codes	Description	
0631	Single source drug	
0636	Drugs requiring detailed coding	

The coding information contained herein is for informative purposes only, and is not a guarantee of coverage or reimbursement for any product or service. This information is not intended to substitute for the physician's independent diagnosis or treatment of each patient. Coding requirements may vary by payer; please consult the payer to determine which codes are required.

For questions regarding CINVANTI billing and coding please call Heron Connect at 1-844-HERON11 (1-844-437-6611) from 8 AM to 5 PM ET, Monday through Friday.

Please see Indications and Important Safety Information on reverse.

www.cinvanti.com



[§] Effective July 1, 2023, providers are **required** to report the JZ modifier on all claims that bill for drugs from single-dose containers that are separately payable **when there are no discarded amounts**. Providers may start using the modifier as of January 1, 2023, however, after July 1,2023 use of the modifier is required.

CINVANTI® (aprepitant) injectable emulsion

Indication

CINVANTI is a substance P/neurokinin-1 (NK1) receptor antagonist, indicated in adults, in combination with other antiemetic agents, for the prevention of:

- acute and delayed nausea and vomiting associated with initial and repeat courses of highly emetogenic cancer chemotherapy (HEC) including high-dose cisplatin as a single-dose regimen.
- delayed nausea and vomiting associated with initial and repeat courses of moderately emetogenic cancer chemotherapy (MEC) as a single-dose regimen.
- nausea and vomiting associated with initial and repeat courses of MEC as a 3-day regimen.

<u>Limitations of Use:</u> CINVANTI has not been studied for treatment of established nausea and vomiting.

Important Safety Information

Contraindications

CINVANTI is contraindicated in patients with hypersensitivity to any of the components of CINVANTI. Concurrent use of pimozide with CINVANTI is contraindicated.

Warnings and Precautions

Clinically Significant CYP3A4 Drug Interactions

Aprepitant is a substrate, weak-to-moderate (dose-dependent) inhibitor, and an inducer of CYP3A4.

- Use of CINVANTI with other drugs that are CYP3A4 substrates may result in increased plasma concentration of the concomitant drug.
 - Use of pimozide with CINVANTI is contraindicated due to the risk of significantly increased plasma concentrations of pimozide, potentially resulting in prolongation of the QT interval, a known adverse reaction of pimozide.
- Use of CINVANTI with strong or moderate CYP3A4 inhibitors (e.g., ketoconazole, diltiazem) may increase plasma concentrations of aprepitant and result in an increased risk of adverse reactions related to CINVANTI.
- Use of CINVANTI with strong CYP3A4 inducers (e.g., rifampin) may result in a reduction in aprepitant plasma concentrations and decreased efficacy of CINVANTI.

Hypersensitivity Reactions

Serious hypersensitivity reactions, including anaphylaxis, during or soon after administration of CINVANTI have occurred. Symptoms including dyspnea, eye swelling, flushing, pruritus, and wheezing have been reported. If hypersensitivity reactions occur, discontinue CINVANTI. Do not reinitiate CINVANTI in patients who experience these symptoms with previous use.

Decrease in INR with Concomitant Warfarin

Co-administration of CINVANTI with warfarin, a CYP2C9 substrate, may result in a clinically significant decrease in the International Normalized Ratio (INR) of prothrombin time. Monitor the INR in patients on chronic warfarin therapy in the 2-week period, particularly at 7 to 10 days, following initiation of CINVANTI with each chemotherapy cycle.

Risk of Reduced Efficacy of Hormonal Contraceptives

The efficacy of hormonal contraceptives may be reduced during administration of and for 28 days following the last dose of CINVANTI. Advise patients to use effective alternative or back-up methods of non-hormonal contraception during treatment with CINVANTI and for 1 month following administration of CINVANTI or oral aprepitant, whichever is administered last.

Use in Specific Populations

Avoid use of CINVANTI in pregnant women as alcohol is an inactive ingredient for CINVANTI. There is no safe level of alcohol exposure in pregnancy.

Adverse Reactions

The most common adverse reactions are:

- Single-dose fosaprepitant with MEC (≥2%): fatigue, diarrhea, neutropenia, asthenia, anemia, peripheral neuropathy, leukopenia, dyspepsia, urinary tract infection, pain in extremity.
- 3-day oral aprepitant with MEC (≥1% and greater than standard therapy): fatigue and eructation.
- Single-dose fosaprepitant with HEC: generally similar to 3-day oral aprepitant. In addition, infusion site reactions (3%) occurred.
- Single-dose CINVANTI (≥2%): headache and fatigue. The safety profile of CINVANTI in healthy subjects who received a single 2-minute injection was similar to that seen with a 30-minute infusion.

Report side effects to the FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

