CARBOPLATIN- carboplatin injection, solution Ingenus Pharmaceuticals, LLC

Carboplatin Injection, USP Rx only

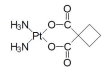
Carboplatin njection should be administered under the supervision of a qualified physician experienced in the use of cancer chemotherapeutic agents. Appropriate management of therapy and complications is possible only when adequate treatment facilities are readly available.

Bone marrow suppression is dose related and may be severe, resulting in infection and/or bleeding. Anemia may be cumulative and may require transfusion support. Vomiting is another frequent drug related side effect.

Anaphylactic-like reactions to carboplatin have been reported and may occur within mitutes of carboplatin hijection administration. Epinephrine, corticosteroids, and anthistamines have been employed to alleviate symptoms.

DESCRIPTION

Carboptath injection, USP is supplied as a sterile, pyrogen-free, 10 mg/mL aqueous solution of carboptath, USP. Carboptath, USP is a platinum coordination compound. The chemical name for carboptath, USP is platitum, disminime [1.1-cyclobutaneticarboxylato[2-0.07]-(SP-4-2), and carboptath, USP has the following structural formula:



C₆H₁₂N₂O₄ Pt M.W. 371.25

Carboplatin, USP is a crystalline powder. It is soluble in water at a rate of approximately 14 mg/mL, and the pH of a 1% solution is 5 to 7. It is virtually insoluble in ethanol, acetone, and dimethylacetamide.

CLINICAL PHARMACOLOGY

CLINICAL PHARMACOLOGY Carboptath, Rec Explain, produces predominantly interstrand DNA cross-links rather than DNA-protein cross-links. This effect is apparently cell-cycle nonspecific. The aquation of carboptim, which is thought to produce the active species, occurs at a siswer rate than in the case of csplatin. Despite this difference, It appears that both carboptian and csplatin nucleus equal numbers of drug -DNA cross-links, causing equivalent lesions and biological effects. The differences in potencies for carboptian and csplatin appear to be directly related to the difference in aquation rates.

cisplatin appear to be directly related to the difference in aquation rates. In patients with resatine clearances of about 60 mL/min or greater, plasma levels of instance arbightin decay in a biphasic manner after a 30-minute intravenous infusion of 300.110 to 500 mL/minute intravenous infusion of 300.110 to 500 mL/minute intravenous infusion of 800.110 to 500 mL/minute intravenous infusion of 800 mL/minute intravenous intervenous interv

The major route of elimination of carboptatin is renal excretion. Patients with creatinine clearances of approximately 60 mL/min or greater excrete 56% of the dose in the urine within 12 hours and 71% of the dose within 24 hours. All of the platimum in the 24-hour urine is present as carboptatin. Only 3% to 5% of the administered platimum is excreted in the urine between 24 and 96 hours. There are insufficient data to determine whether bilary excretion occurs.

In patients with creatinne clearances below 60 mL/min, the total body and renal clearances of carboplatin decrease as the creatinne clearance decreases. Carboplatin dosages should therefore be reduced in these patients (see DOSAGE AND ADMINISTRATION)

Ademication Long View The primary determinant of carboplath injection clearance is glomerular fitration rate (GFR) and this parameter of renal function is often decreased in elderly patients. Dosing formulas incorporating estimates of GFR (see DOSACE AND ADIMINSTRATION) to provide predictable carboplath injection plasma AUCs should be used in elderly patients to minimize the risk of toxicity.

CLINICAL STUDIES

Use with Cyclophosphamide for Initial Treatment of Ovarian Cancer In two prospectively randomized, controlled studies conducted by the National Cancer Institute of Canada, Clinical Trials Group (NCIC) and the Southwest Oncology Group (SWOG), 789 chemotheragy naive patients with advanced ovarian cancer were treated with carboplatin or cisplatin, both in combination with cyclophosphamide every 28 days for 6 courses before surgical reevaluation. The following results were obtained from both studies:

Comparative Efficacy

Overview of Pivotal Trials

	NCIC	SWOG
Number of patients randomized	447	342
Median age (years)	60	62
Dose of cisplatin	75 mg/m ²	100 mg/m ²
Dose of carboplatin	300 mg/m ²	300 mg/m ²
Dose of cyclophosphamide	600 mg/m ²	600 mg/m ²
Residual tumor < 2 cm (number of patients)	39% (174/447)	14% (49/342)

Clinical Response in Measurable Disease Patients		
	NCIC	SWOG
Carboplatin (number of patients)	60% (48/80)	58% (48/83)
Cisplatin (number of patients)	58% (49/85)	43% (33/76)
95% CI of difference (Carboplatin-Cisplatin)	(-13.9%, 18.6%)	(-2.3%, 31.1%)

Pathologic Complete Response*		
	NCIC	SWOG
Carboplatin (number of patients)	11% (24/224)	10% (17/171)
Cisplatin (number of patients)	15% (33/223)	10% (17/171)
95% CI of difference	(-10.7%, 2.5%)	(-6.9%, 6.9%)

 Image: Carboplatin-Csiplatin
 Image: Carboplatin
 Ima

Progression-Free Survival (PFS)

	NCIC	SWOG
Median		
Carboplatin	59 weeks	49 weeks
Cisplatin	61 weeks	47 weeks
2-year PFS*		
Carboplatin	31%	21%
Cisplatin	31%	21%
95% CI of difference (Carboplatin-Cisplatin)	(-9.3, 8.7)	(-9, 9.4)
3-year PFS*		
Carboplatin	19%	8%
Cisplatin	23%	14%
95% CI of difference	(-11.5, 4.5)	(-14.1, 0.3)
(Carboplatin-Cisplatin)		
Hazard Ratio [†]	1.10	1.02
95% (Carboplatin-Cisplatin)	(0.89, 1.35)	(0.81, 1.29)

In a compartin-separation Kaplan-Meier Estimates Unrelated deaths accurring in the absence of progression were counted as events (progression) in the analysis. Analysis adjusted for factors found to be of prognostic significance were consiste with unadjusted analysis.

Survival

	NCIC	SWOG
Median		
Carboplatin	110 weeks	86 weeks
Cisplatin	99 weeks	79 weeks
2-year Survival°		
Carboplatin	51.9%	40.2%
Cisplatin	48.4%	39%
95% CI of difference (Carboplatin-Cisplatin)	(-6.2, 13.2)	(-9.8, 12.2)

3-year Survival"		I
Carboplatin	34.6%	18.3%
Cisplatin	33.1%	24.9%
95% CI of difference (Carboplatin-Cisplatin)	(-7.7, 10.7)	(-15.9, 2.7)
Hazard Ratio [†]	0.98	1.01
95% (Carboplatin-Cisplatin)	(0.78, 1.23)	(0.78, 1.30)
*Kaplan-Meier Estimates [†] Analysis adjusted for factors fo with unadjusted analysis.	und to be of prognostic	significance were consistent

Comparative Toxicity

The pattern of toxicity exerted by the carboplatin-containing regimen was significantly different from that of the cisplatin-containing combinations. Differences between the two studies may be explained by different cisplatin dosages and by different supportive care.

care. The carboplatin-containing regimen induced significantly more thrombocytopenia and, in one study, significantly more leukopenia and more need for transfusional support. The capitatri-containing regimen produced significantly more anemia in one study. However, no significant differences occurred in incidences of infections and hemorrhagic episodes Non-hematologic toxicites (enses, neurotoxicity, ototoxicity, renatio toxicity, hypomagnesemia, and alopecia) were significantly more frequent in the cleplathcontaining arms.

ADVERSE EXPERIENCES IN PATIENTS WITH OVARIAN CANCER NCIC STUDY

		STUDY		
		Carboplatin Arm		P-Values [†]
		Percent*	Percent*	
Bone Marrow				
Thrombocytopenia	<	70	29	< 0.001
	100,000/mm ³			
	<	41	6	< 0.001
	50,000/mm ³			
Neutropenia	< 2,000	97	96	ns
	cells /mm ³			
	< 1,000	81	79	ns
	cells /mm ³			
Leukopenia	< 4,000	98	97	ns
	cells /mm ³			
	< 2.000	68	52	0.001
	cells /mm ³			
Anemia	< 11 g/dL	91	91	ns
	< 8 g/dL	18	12	ns
Infections	4 0 g/uc	14	12	ns
Bleeding		10	4	ns
Transfusions		42	4 31	0.018
Gastrointestinal		42	71	0.010
				-
Nausea and		93	98	0.010
vomiting				
Vomiting		84	97	< 0.001
Other GI side		50	62	0.013
effects				
Neurologic				
Peripheral		16	42	< 0.001
neuropathies				
Ototoxicity		13	33	< 0.001
Other sensory		6	10	ns
side effects				
Central		28	40	0.009
neurotoxicity				
Renal				
Serum creatinine		5	13	0.006
elevations				
Blood urea		17	31	< 0.001
elevations				
Hepatic				
Bilirubin elevations		5	3	ns
SGOT elevations		17	13	ns
Alkaline	1	E		1
phosphatase		Γ	Γ	Γ
elevations				
Electrolyte loss				
Sodium		10	20	0.005
Potassium		16	22	ns
Calcium		16	19	ns
		16 63	19 88	
Magnesium		co	00	< 0.001
Other side		1	1	1
effects				_
Pain		36	37	ns
Asthenia		40	33	ns
Cardiovascular		15	19	ns
Respiratory		8	9	ns
Allergic		12	9	ns
Genitourinary		10	10	ns
Alopecia‡		50	62	0.017
Mucositis		10	9	ns
		14 V	r	1.2

 Mucositis
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 Values are in percent of evaluable patients
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ADVERSE EXPERIENCES IN PATIENTS WITH OVARIAN CANCER SWOG STUDY

		Carboplatin Arm	Cisplatin Arm	P-Values [†]
		Percent*	Percent*	
Bone Marrow				
Thrombocytopenia		59	35	< 0.001
	100,000/mm ³			
	<	22	11	0.006
	50,000/mm ³			
Neutropenia	< 2,000	95	97	ns
	cells /mm ³			
	< 1,000	84	78	ns
	cells /mm ³			
Leukopenia	< 4,000	97	97	ns
	cells /mm ³			
	< 2,000	76	67	ns
	cells /mm ³			
Anemia	< 11 g/dL	88	87	ns
	< 8 g/dL	8	24	< 0.001
Infections		18	21	ns
Bleeding		6	4	ns
Transfusions		25	33	ns
Gastrointestinal				
Nausea and	1	94	96	ns
vomiting	1	-	-	[
Vomiting		82	91	0.007
Other GI side		40	48	ns
effects				
Neurologic				
Peripheral		13	28	0.001
neuropathies		15	20	0.001
Ototoxicity		12	30	< 0.001
Other sensory		4	6	ns
side effects		7	0	113
Central		23	29	ns
neurotoxicity		25	29	115
Renal				-
Serum creatinine		7	38	< 0.001
elevations		′	20	<0.001
Blood urea				
elevations		ī	-	-
Hepatic				-
Bilirubin elevations		5	3	ns
SGOT elevations		23	16	
			20	ns
Alkaline phosphatase		29	20	ns
prospratase elevations				
Electrolyte loss				
				+
Sodium Potassium		-	-	F.
		-	1	1
Calcium		-	-	1
Magnesium		58	77	< 0.001
Other side				1
effects		l		-
Pain		54	52	ns
Asthenia		43	46	ns
Cardiovascular		23	30	ns
Respiratory		12	11	ns
Allergic		10	11	ns
Genitourinary		11	13	ns
Alopecia [‡]		43	57	0.009
Mucositis		6	11	ns
*Values are in percen	t of evaluable na			

Values are in percent of evaluable patients
 [†]ns = not significant, p>0.05
 [‡]May have been affected by cyclophosphamide dosage delivered

Use as a Single Agent for Secondary Treatment of Advanced Ovarian Cancer

In two prospective, randomized controlled studies in patients with advanced ovaria cancer previously treated with chemotherapy, carboplatin achieved 6 clinical compl responses in 47 patients. The duration of these responses ranged from 45 to 71+ weeks.

INDICATIONS

Initial Treatment of Advanced Ovarian Carcinoma

Initial Treatment of Advanced Ovarian Carcinoma Carboptin higherino is indicated for the high treatment of advanced ovarian carcinoma in established combination with other approved chemotherapeutic agents. One established combination regime consists of carboptian and cyclophosphamide. Two randomized controlled studies conducted by the NCC and SWOG with carboptain versus cisplath, both in combination with cyclophosphamide. Two equivalent overall survival between the two groups (see CLINICAL STUDIES). There is limited statistical power to demonstrate equivalence in overal pathobgic complete response rates and bing-term survival (> 3 years) because of the small number of palents with these outcomes: the small number of palents with resid tumor < 2 cm after hitld surgery also limits the statistical power to demonstrate equivalence in this subgroup.

Secondary Treatment of Advanced Ovarian Carcinoma

Carboplatin njection is indicated for the pallative treatment of patients with ovariar carcinoma recurrent after prior chemotherapy, including patients who have been previously treated with cisplatin.

Within the group of patients previously treated with cisplatin, those who have developed progressive disease while receiving cisplatin therapy may have a decreased response rate.

CONTRAINDICATIONS

Carboplatin injection is contraindicated in patients with a history of severe allergic reactions to cisplatin or other platinum-containing compounds.

Carboplatin injection should not be employed in patients with severe bone marrow depression or significant bleeding.

WARNINGS

Bone marrow suppression (leukopenia, neutropenia, and thrombocytopenia) is dose-dependent and is also the dose-limiting toxicity, Peripheral blood counts should be frequently monicored during carboplain injecton treatment and, when appropriate, unit recovery is achieved. Median nadr occurs at day 21 in patients receiving single agent carbopatin. In general, single intermittent courses of carbopatin should not be repeated until leukocyte, neutrophi, and platelet counts have recovered.

Since anemia is cumulative, transfusions may be needed during treatment with carboplatin, particularly in patients receiving prolonged therapy.

Catalogian, park dar yr i paeris telewnig probliget i magy: Bone marrow suppression is increased in patients who have received prior therapy, especially regimens including cisplatin. Marrow suppression is also increased in patients with imparied kiloney function. Initial carbopikin injection dosages in these patients should be appropriately reduced (see DOSAEC AND ADMINISTRATION) and blood counts should be carefully monthered between courses. The use of carbopikin in combination with other bone marrow suppressing therapies must be carefully manage with respect to dosage and timing in order to minime additive effects.

Carbopiath has imited nephrotoxic potential, but concomitant treatment with aminoglycosides has resulted in increased renal and/or audiologic tox/city, and cauton must be exercised when a patient receives both drugs. Clinically significant hearing bis has been reported to occur in pediatric patients when carbopiath was administered at higher than recommended doses in combination with other obtoxics agents.

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Although peripheral neurotoxicity is infrequent, its incidence is increased in patients older than 65 years and in patients previously treated with cisplatin. Pre-existing carbophatin-duce neurotoxicity does not worsen in about 70% of the patients receiving carbophatin as secondary treatment.

Loss of vision, which can be complete for light and colors, has been reported after the use of carboptian with doses higher than those recommended in the package insert. Vision appears to recover totally or to a significant extent within weeks of stopping these high doses.

As in the case of other platinum-coordination compounds, allergic reactions to carbopatin have been reported. These may occur within minutes of administration should be managed with appropriate supportive therapy. There is increased risk of allergic reactions including anaphylaxis in patients previously exposed to platinum therapy (see CONTRIAINDICATIONS and ADVERSE REACTIONS). Allergic Reactions) High dosages of carboplatin (more than 4 times the recommended dose) have resulted in severe abnormalities of liver function tests.

In server examininates on when function tests. Carboptish injection may cause feal harm when administered to a pregnant woman. Carboptish has been shown to be embryotoxic and treatogenic in rats. There are no adequate and well-controlled studies in pregnant ownen. If this drug is used during pregnancy, or if the patient becomes pregnant while receiving this drug, the patient should be apprived of the potential hazard to the fetus. Women of childbearing potential should be advised to avoid becoming pregnant.

PRECAUTIONS

General

Needles or intravenous administration sets containing aluminum parts that may come in contact with carboplatin injection should not be used for the preparation or administration of the drug. Aluminum can react with carboplatin causing precipitate formation and loss of potency. Drug Interactions

The renal effects of nephrotoxic compounds may be potentiated by carboplatin

Carcinogenesis, Mutagenesis, Impairment of Fertility

carcnogeness, Mutagenesis, Impairment of Fertility
The carcinogenic potential of carchopain has not been studied, but compounds with similar mechanisms of action and matagenetic profiles have been reported to be carcinogenic. Cost ain has been used on to the mutaget both in yours and it wice. It has also been shown to be embryotaxic and treatogenic in rais receiving the drug during organogenesis. Secondary malignancies have been reported in association with multi-drug therapy.

Pregnancy

Pregnancy category D See WARNINGS

Nursing mothers

It is not known whether carboplatin is excreted in human milk. Because there is possibility of toxicity in nursing infants secondary to carboplatin treatment of ti mother, it is recommended that breastfeeding be discontinued if the mother is with carboplatin injection.

Pediatric Use

Safety and effectiveness in pediatric patients have not been established (see WARNINGS, "audiologic toxicity").

Geriatric Use

Geniatric Use Of the 789 patients in hitial treatment combination therapy studies (NCIC and SWOG), 395 patients were treated with carboplatin in combination with cyclophosphamide. Of these, 141 were over 65 years of data or 1 these trials, age was not a prognostic factor for survival. In terms of safety, elderly patients treated with carboptist were more likely to develop severe thromoscytopenis than younger patients in a combined database of 1,942 patients (141 were \geq 55 years or doiler. In these trials, age was not a prognostic factor for survival. In terms of safety, elderly patients treated with single agent carboptist for different tumor types, a similar incidence of adverse events was seen in patients 65 years and older and in patients less than 65. Other reported clinical experience is and itedimided differences in reportes between ethely and yoursus patients, but preater sant-bidy of in the older individuals can be used by ADMINISTRATION).

ADVERSE REACTIONS

For a comparison of toxicities when carboplatin or cisplatin was given in combination with cyclophospharmide, see CLINICAL STUDIES, Use with Cyclophospharmide for Initial Treatment of Ovarian Cancer, Comparative Toxicty.

ADVERSE EXPERIENCES IN PATIENTS WITH OVARIAN CANCER

		First Line Combination Therapy [*] Percent	Second Line Single Agent Therapy [†] Percent
Bone Marrow			
Thrombocytopenia	< 100,000/mm ³	66	62
	< 50,000/mm ³	33	35
Neutropenia	< 2,000 cells /mm ³	96	67
	< 1,000 cells /mm ³	82	21
Leukopenia	< 4,000 cells /mm ³	97	85
	< 2,000 cells /mm ³	71	26
Anemia	< 11 g/dL	90	90
	< 8 g/dL	14	21
Infections		16	5
Bleeding		8	5
Transfusions		35	44
Gastrointestinal			
Nausea and vomiting		93	92
Vomiting		83	81

Other GI side effects	46	21
Neurologic		
Peripheral neuropathies	15	6
Ototoxicity	12	1
Other sensory side effects	5	1
Central neurotoxicity	26	5
tenal		
erum creatinine levations	6	10
llood urea elevations	17	22
lepatic		
Bilirubin elevations	5	5
GOT elevations	20	19
Ikaline phosphatase	29	37
levations		
lectrolyte loss		
odium	10	47
otassium	16	28
alcium	16	31
lagnesium	61	43
Other side effects		
'ain	44	23
sthenia	41	11
ardiovascular	19	6
tespiratory	10	6
llergic	11	2
Senitourinary	10	2
lopecia‡	49	2
Aucositis Use with Cyclophosphamide for I	8	1

Use with Cyclophosphamide for Initial Treatment of Ovarian Cancer: Data are based on the experience of 33 patients with ovarian carer (regardless of baseline study, who received initial combination threngy with carboplath and Cyclophosphamide in two randomized combiled studies conducted by SWOG and MCC (see CLINICAL STODES). Combination with cyclophosphamide as well as duration of treatment may be combined on two recombined studies and the duration of treatment may be combined on two recombined studies and the duration of treatment may be based on the experience of 535 patients with prevuous treated ovarian carcinoma (regardless of baseline status) who received single agent carboplatin.

In the narrative section that follows, the incidences of adverse events are based on data from 1,893 patients with various types of tumors who received carboplatin as single agent therapy.

Hematologic Toxicity

Hematologic Toxicity Bone marcrow suppression is the dose-imiting toxicity of carbopiatin. Thrombocytopenia with platelet counts below 50,000/mm³ occurs in 25% of the patients (35% of pretreated ovarian cancer patients); neutropenia with granubcyte counts below 1,000/mm³ occurs in 15% of the patients (21% of pretreated ovarian cancer patients); laukopenia with WBC counts below 2,000/mm³ occurs in 15% of the patients (26% of pretreated ovarian cancer patients). The nadir usually occurs about day 21 in patients receiving single agent theraps, by 60x 28, 05% of patients have patients counts about 100,000mm³. 74% have neutrophil counts above 2,000mm³, 67% have leukocyte counts above 4,000mm³.

Marrow suppression is usually more severe in patients with impaired kidney function Patients with poor performance status have also experienced a higher incidence of severe leukopenia and thrombocytopenia.

The hematologic effects, although usually reversible, have resulted in infectious or hemorrhagic complications in 5% of the patients treated with carbopatin, with drug related death occurring in less than 1% of the patients. Fever has also been reported in patients with neutropenia.

patents win neutropena. Anemia with menoglobin less than 11 g/dL has been observed in 71% of the patients who started therapy with a baseline above that value. The incidence of anemia increases with increasing exposure to carboplatin. Transfusions have been administered to 26% of the patients treated with carboplatin (44% of previously treated ovarian cancer patients). Bone marrow depression may be more severe when carboplatin is combined with other bone marrow suppressing drugs or with radiotherapy.

Gastrointestinal Toxicity

Gastrointestinal Toxicity Vomiting occurs in 65% of the patients (01% of previously treated ovarian cancer patients) and in about one- third of these patients. It is severe. Carboplatin, as a single agent or in combination, is significantly less enterlognic than cabpatin, however, patients previously treated with emetogenic agents, especially capatin, appear to be more prone to vomiting. Nusuea above occurs in an additional 10% to 15% of patients. Both masses and vomiting usually cease within 24 hours of treatment and are often responsive to antemetic measures. Although no conclusive efficacy data exist with the following schedules, prolonged administration of carboplatin, either by continuous 24-hour infusion or by daip puble doses given for 5 consecutive days, was associated with less severe womiting than the single-dose intermittent schedule. Emesis was increased when a directs observed requently were pain, in 17% of the patients, diarrhea, in 6%; and constipation, also in 6%.

Neurologic Toxicity

Neurologic Toxichty Periyhera neuropathisn have been observed in 4% of the patients receiving carboplatin (6% of pretroated owains cancer patients) with mild paresthesiles occurring most requerity. Carboplatin therapy produces significantly forem and less severe neurologic side effects than does therapy with cisplatin. However, patients older than 65 years and/or previously treated with cisplatin. However, patients older than 65 years and/or previously treated with cisplatin. However, patients older than 65 years and/or previously treated with cisplatin appear to have an increased risk (10%) (or perpheral neuropathies. In 70% of the patients with pre-exiting cisplatin-induced perpheral neuropathies. In 70% of the patients with pre-exiting cisplatin-induced perpheral neuropathies. In 70% of the patients with pre-exiting cisplatin-induced perpheral neuropathies. In 70% of the patients with pre-exiting cisplatin-induced perpheral neuropathies. The perpheral neuropathies and appear to be most oftem related to the use of antiemetics.

Although the overall incidence of peripheral neurologic side effects induced by carboplath is low, prolonged treatment, particularly in cisplatin pretreated patients, may result in cumulative neurotoxicity.

Nephrotoxicity

Nephrotoxicity Development of abnormal renal function test results is uncommon, despite the fact that carbophith, unlike csplath, has usually been administered without high-volume fluid hydration and/or forced duriess. The incidences of abnormal renal function tests reported are 6% for serum creatinne and 14% for blood urea ntrogen (10% and 22%, respective), in pretrated ovarian cancer patients). Most of these reported abnormalities have been mild and about one-half of them were reversible.

Creatinine clearance has proven to be the most sensitive measure of kidney function in patients receiving carboplath, and It appears to be the most useful test for correlating drug clearance and bone marrow suppression. Twenty-seven precent of the patients who had a baseline value of 60 mL/min or more demonstrated a reduction below this value during carboplath thre app.

Hepatic Toxicity

nepate loakery The incidences of abnormal liver function tests in patients with normal baseline values were reported as follows: total bifruibn; 5%, 5007, 15%, and akaline phosphates, 24%, (5%, 15%, and 5%, respectively, in partenside outrain cancer patients). These (5%, 15%, and 5%, respectively, in partenside outrain cancer patients). These abhough the role of metastatic tumor in the liver may complicate the assessment in many patients. In a limited series of patients receiving very high dosages of carboplatin and autobagous bone marrow transplantation, severe abnormalities of liver function tests were reported.

Electrolyte Changes

The incidences of abnormally decreased serum electrolyte values reported were as follows: sodium, 29%; potassium, 20%; calckium, 22%; and magnesium, 29%; (47%, 25%, 33% and 43%, respectively, in pretrated ovarian cancer patients). Electrolyte supplementation was not routinely administered concomitantly with carboptain, and these electrolyte abnormalities were rarely associated with symptoms.

Allergic Reactions

Allergic Reactions Hypersensitivity to carboplatin has been reported in 2% of the patients. These allergic reactions have been similar in nature and severity to those reported with other platinum: containing compounds, i.e., rach, uritaria, erythema purutus, and rarely bronchospasm and hypotension. Anaphylactic reactions have been reported as part of postmarketing surveillance (ear ARNINGS). These reactions have been successfully managed with standard epinephrine, corticosteroid, and antihistamine therapy.

Injection Site Reactions

Injection site reactions, including redness, swelling, and pain, have been reported during postmarketing surveillance. Necrosis associated with extravasation has also been reported.

Other Events

Uner tvents Pain and asthenia were the most frequently reported miscelaneous adverse effects; their relationship to the tumor and to anemia was likely. Abpecia was reported (3%). Cardivascular, respiratory, genthourinary, and muccals side effects have occurred in 6% or less of the patients. Cardivascular events (cardiac falure, embolsm, ceretrovascular accidents) were fatal in less than 1% of the patients and did not appear to be related to chemotherapy. Cancer-associated hemolytic uremic syndrome has been reported rarely.

Malaise, anorexia, hypertension, dehydration, and stomatitis have been reported as part of postmarketing surveilance.

To report SUSPECTED ADVERSE REACTIONS, contact Ingenus Pharmaceuticals, LLC at (877) 748-1970 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch

OVERDOSAGE

There is no known antidote for carboplatin injection overdosage. The anticipated

DOSAGE AND ADMINISTRATION

NOTE: Aluminum reacts with carboplatin causing precipitate formation and loss of potency,

therefore, needles or intravenous sets containing aluminum parts that may come in contact with the drug must not be used for the preparation or administration of carboplatin injection. Single Agent Therapy

Carboplatin njection, as a single agent, has been shown to be effective in patients with recurrent ovarian carchona at a dosage of 360 mg/m² IV on day 1 every 4 weeks (aternatively see Formula Dosing). In general, however, single intermittent courses of carboplath should not be repeated until the neutrophil count is at kast 20,000 and the plateter count is nates 1100,000.

Combination Therapy with Cyclophosphamide

In the chemotherapy of advanced ovarian cancer, an effective combination for previously untreated patients consists of:

Carboplatin - 300 mg/m² IV on day 1 every 4 weeks for 6 cycles (alternatively see Formula Dosing).

Cyclophosphamide - 600 mg/m² IV on day 1 every 4 weeks for 6 cycles. For directions regarding the use and administration of cyclophosphamide please refer to its package insert (see CLINICAL STUDIES).

Intermittent courses of carboplatin in combination with cyclophosphamide should not be repeated until the neutrophil count is at least 2,000 and the platelet count is at least 100,000.

Dose Adjustment Recommendations

Pretreatment platelet count and performance status are important prognostic factors for severity of myelosuppression in previously treated patients.

The suggested dose adjustments for single agent or combination therapy shown in the table below are modified from controlled trials in previously treated and untreated patients with ovarian carcinoma. Bodo counts were done weekly, and the recommendations are based on the lowest post-treatment platelet or neutrophl value.

Platelets	Neutrophils	Adjusted Dose*
		(From Prior Course)
> 100,000	> 2,000	125%
50 to 100,000	500 to 2,000	No Adjustment
< 50,000	< 500	75%
cyclophosphamide in co adjusted at a lower leve	ombination. In the contro	ngle agent or to both carboplatin and lled studies, dosages were also e myelosuppression. Escalations studies.

Carboplatin injection is usually administered by an infusion lasting 15 minutes or longer. No pre- or post-treatment hydration or forced diuresis is required.

Patients with Impaired Kidney Function

Patients with creatine character values below 60 mL/min are at increased risk of severe bone marrow suppression. In renaly-impaired patients who received single agent carobaltist herapy, the incidence of severe leukopenia, neutropenia, or thrombocytopenia has been about 25% when the dosage modifications in the table below have been used.

Baseline Creatinine Clearance	Recommended Dose on Day 1
41 to 59 mL/min	250 mg/m ²
16 to 40 mL/min	200 mg/m ²

The data available for patients with severely impaired kidney function (creatinine clearance below 15 mL/min) are too limited to permit a recommendation for treatment. These dosing recommendations apply to the initial course of treatment. Subsequent dosages should be adjusted according to the patient's tolerance based on the degree of bone marrow suppression.

Formula Dosing

Formula Dosing Another approach for determining the initial dose of carboplatin hjection is the use of mathematical formulae, which are based on a patient's pre-existing renal function or renal function and desired plateter dark. Renal excremion is the major runce of elimination for carboplatin (see CLINICAL PHARMACDLOGY). The use of dosing formulae, as compared to empirical dose cacluation based on body surface area, adows compensation for patient variations in pretreatment renal function that might otherwise result in either underdosing (in patients with above average renal function) or overdosing (in patients with impaired renal function).

A simple formula for calculating dosage, based upon a patient's glomerular filtration rate (GFR in mL/min) and carboplatin injection target area under the concentration versus time curve (AUC in mg/mL-min), has been proposed by Calvert. In these studies, GFR was measured by ^{S1}Cr-EDTA clearance.

CALVERT FORMULA FOR CARBOPLATIN DOSING Total Dose (mg) = (target AUC) × (GFR + 25) Note: With the Calvert formula , the total dose of carboplatin is calculated in mg , <u>not</u> mg/m².

The target AUC of 4 mg/mL+min to 6 mg/mL+min using single agent carboplatin appears to provide the most appropriate dose range in previously treated patients. This study also showed a trend between the AUC of single agent carboplatin administered to previously treated patients and the likelihood of developing toxicity.

	% Actual Toxicity in Previous	ly Treated Patients
	Gr 3 or Gr 4	
AUC (mg/mL·min)	Thrombocytopenia	Gr 3 or Gr 4 Leukopenia
4 to 5	16%	13%
6 to 7	33%	34%

Geriatric De

Because renal function is often decreased in elderly patients, formula dosing of carboplatin injection based on estimates of GFR should be used in elderly patients to provide predictable plasma carboplatin injection AUCs and thereby minimize the risk of to view. provide toxicity

PREPARATION OF INTRAVENOUS SOLUTIONS

Carboplatin injection is a premixed aqueous solution of 10 mg/mL carboplatin.

Carboplatin aqueous solution can be further diluted to concentrations as low as 0.5 mg/mL with 5% Dextrose in Water (D₅W) or 0.9% Sodium Chloride Injection, USP. Multin properties of the control of

HOW SUPPLIED

Each mL of carboplatin injection, USP contains 10 mg of carboplatin, USP in water for injection and is available as follows:

NDC Number	Contents	Size
50742-445-05	50 mg	5 mL Multidose vial
50742-446-15	150 mg	15 mL Multidose vial
50742-447-45	450 mg	45 mL Multidose vial
50742-448-60	600 mg	60 mL Multidose vial

STORAGE

Unopened vials of carboplatin injection, USP are stable to the date indicated on the package when stored at 20° to $25^{\circ}C$ (68° to $77^{\circ}F$) [See USP Controlled Room Temperature].

PROTECT FROM LIGHT.

Carboplatin injection, USP multidose vials maintain microbial, chemical, and physical stability for up to 14 days at 25°C following multiple needle entries.

Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration. Solutions for infusion should be discarded 8 hours after preparation.

HANDLING AND DISPOSAL

Caution should be exercised in handling and preparing carboplatin injection. Several guidelines on this subject have been published.¹⁻⁴ guidence on this subject name deel published.² The published of the subject of

NIOSH Alert: Preventing occupational exposures to antineoplastic and other hazardous drugs in healthcare settings. 2004. U.S. Department of Health and Human Bervices, Public Health Service, Centers for Disease Control and Prevention, National Institute for Occupational Safety and Health, DHHS (NIOSH) Publication No. 2004-165.

OSHA Technical Manual, TED 1-0.15A, Section VI: Chapter 2. Controlling occupational exposure to hazardous drugs. OSHA, 1999. http://www.osha.gov/dts/osta/otm/otm_vi/otm_vi2.html

American Society of Health-System Pharmacists. ASHP guidelines on handling hazardous drugs.

Am | Health-Syst Pharm. 2006;63:1172-1193.

Polovich M, White JM, Kelleher LO, eds. 2005. Chemotherapy and biotherapy guidelines and recommendations for practice. 2nd ed. Pittsburgh, PA: Oncology Nursing Society.

Manufactured for:

Ingenus Pharmaceuticals, LLC Orlando El 32811-7913 Made in Switzerland

Revised: 06/2023 ingenus 😵

Patient Insert Patient Insert CARBOplatin (kar boe pla tin) Injection

Rx Only

Read this entrice leaflet carefully. Keep it for future reference. This information will help you learn more about carboplain. It cannot, however, cover all the possible warnings or side effects relating to carboplain, and it does not list all of the benefits and risks of carboplath. Your doctor should always be your first choice for detailed information about your medical condition and your treatment. Be sure to ask your doctor about any questions you may have.

What is cancer?

Under normal conditions, the cells in your body divide and grow in an orderly, controlled fashon. Cell division and growth are necessary for the human body to perform its functions and to repair itself. Cancer cells are different from normal cells because they are not able to control ther growth. The reasons for this abnormal growth are not yet fully understood.

A tumor is a mass of unhealthy cells that are dividing and growing fast and in an uncontrolled way. When a tumor invades surrounding healthy body tissue it is known as a malignant tumor. A malignant tumor can spread (metastasize) from its original location to other parts of the body.

What is carboplatin?

Carboplatin is a medicine that is used to treat cancer of the ovaries. It acts by interfering with the division of rapidly multiplying cells, particularly cancer cells.

Who should not take carboplatin?

no should not take can oppating eatment with carboplatin is not recommended if you: are allergic to carboplatin or other platinum-containing products; have a weakened blood-forming system (bone marrow depression) or significant have a weakened blood-forming system (bone marrow depression) or bleeding;
are pregnant, intend to become pregnant, or are breastfeeding a baby.

How is carboplatin used?

reve is carecoplatin used?
Only a professional experienced in the use of cancer drugs should give you this medication. Carecoplatin is given by dripping the medicine slowly and directly into a vein medication. Carecoplatin for you based on your weight, height, and kidney function. Carecoplatin may be given above or with other drugs. Treatment is usually repeated every four weeks for a number of cycles.

Before and after carboplatin treatment, your doctor may give you medication to lessen the nausea and vomiting associated with this cancer treatment.

What should you tell your doctor before starting treatment with carboplatin? Discuss the benefits and risks of carboplatin with your doctor before beginning

- Be sure to inform your doctor: If you are allergic to carboplatin or other platinum-containing products; If you are on intend to become pregnant, since carboplatin may harm the developing fetus. It is important to use effective birth control while you are being treated with carboplatin.
- If you are breastfeeding, since nursing infants may be exposed to carboplatin in this
- If you are breastreading, since nursing intrants may be exposed to carbopatin in this way;
 If you are taking other medicines, including all prescription and non-prescription (over-the-counter) drugs, since carboplatin may affect the action of other medicines;
 If you have any other medical problems, especially chicken pox (including recent exposure to add so or hidner with chicken pox), singles, hearing problems, intection, or kidney dasses, since treatment with carboplatin increases the risk and severity of these conditions.
- What should I avoid while taking carboplatin?

If you are pregnant or think you might be pregnant, or if you are breastfeeding, let your doctor know right away. Carboplatin may harm your developing fetus or breastfeeding baby. If you are a woman of childbearing age, you should use birth control to avoid getting pregnant while you are taking carboplatin.

yexus y preguant: wine you are taxing carboplatin. You should avoid contact with adults and children who have infections, and tell your doctor right away T you show signs of infection such as cough, fever, and/or chils. Also, while you are being treated with carboplatin or after you stop treatment, first check with you doctor before getting any minumizations (vaccinations). Avoid contact with adults or children who have received oral polio vaccine since they can pass the polio virus to you.

What are the possible side effects of carboplatin?

The most service side effects of carboplatin are: • bleeding and reduced blood cells, including reduced red blood cells (anemia) and platelets (needed for proper blood clotting), which may be severe enough to require blood transfusion. You should tell your doctor right away if you notice any unusual bruising or bleeding, hcluding black tary stoods or blood in

- the urine. infection-carboplain can temporarily lower the number of white blood cells in your blood, increasing the risk of infection: if e-threatening aleroic reaction-during and after treatment the doctor or nurse will observe you cardfully for signs of alergic reaction;

kidney and liver problems;
loss of hearing or ringing in the ears;

Contact your doctor right away if you experience any of these effects, or notice effects that worry you or are troublesome.

Of the less serious side effects associated with carboplatin treatment, the most common are nausea, vomking, diarrhea, loss of appette, hair loss and numbness, tingling, burning, or pain in the hands and feet.

This medicine was prescribed for your particular condition. It must be given under close medical supervision by a doctor trained in the use of drugs for the treatment of cancer.

This summary does not include everything there is to know about carboplatin. Medicines are sometimes prescribed for purposes other than those listed in patient leaflets. If you are sometimes prescribed for purposes other than those listed in patient leafter. If yo have questions or concerns, or want more information about carobplatin, your physician and pharmacist have the complete prescribing information upon which this information is based. You may want to read it and discuss it with your doctor. Remember, no written summary can replace careful discussion with your doctor. Manufactured for:

Ingenus Pharmaceuticals, LLC

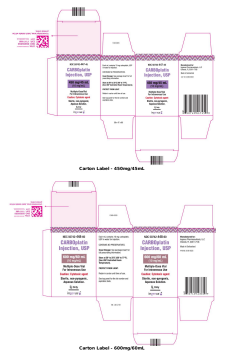
Orlando, FL 32811-7193. Made in Switzerland

Revised: 06/2023 ingenus 😵

PACKAGE LABEL.PRINCIPAL DISPLAY PANEL







CARBOPLATIN

	ormation				
		HUMAN PRESCRIPTION DRUG	Item Code (Source)		NDC:50742-447
Active Ingre					
		dient Name 05) (CARBOPLATIN - UNII:BG3F62ONI			th Strength 450 mg in 45 ml
nactive Ing					
ATER (UNII: 05		gredient Name		S	trength
Packaging					
tem Code	P	Package Description		ting Start Date	Marketing End Date
NDC:50742- 447-45	1 in 1 CARTON		07/01/20	23	
	45 mL in 1 VIA Combination P	L, MULTI-DOSE; Type 0: Not a roduct			
Marketing					
Marketing Category		ation Number or Monograph Citation			Marketing End Date
NDA	ANDA2084	87	07/01/2023	8	
ARBOPLA arboplatin inje	ction, solution	n			
Product Info		HUMAN PRESCRIPTION DRUG			
Product Type Route of Admi			item Code	(Source)	NDC:50742-448
	dient/Active	Moiety			
Active Ingre			De ele	of Strength	1 Strength
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