VALCYTE (valganciclovir) tablets, for oral use VALCYTE (valganciclovir) for oral solution Initial U.S. Approval: 2001

WARNING: HEMATOLOGIC TOXICITY, IMPAIRMENT OF FERTILITY, FETAL TOXICITY. MUTAGENESIS AND CARCINOGENESIS

See full prescribing information for complete boxed warning.

Hematologic Toxicity: Severe leukopenia, neutropenia, anemia, thrombocytopenia, pancytopenia, and bone marrow failure including aplastic anemia have been reported in patients treated with VALCYTE. rment of Fertility: Based on animal data and limited human data, VALCYTE may cause temporary o

permanent inhibition of spermatogenesis in males and suppression of fertility in females. (5.3)
Fetal Toxicity: Based on animal data, VALCYTE has the potential to cause birth defects in humans. (5.4) s and Carcinogenesis: Based on animal data, VALCYTE has the potential to cause cancers in humans. (5.5)

-- RECENT MAJOR CHANGES -----Dosage and Administration Recommended Dosage in Pediatric Patients (2.3)

----- INDICATIONS AND USAGE -----VALCYTE is a deoxynucleoside analogue cytomegalovirus (CMV) DNA polymerase inhibitor indicated for:

Treatment of CMV retinitis in patients with acquired immunodeficiency syndrome (AIDS). Prevention of CMV disease in kidney, heart, and kidney-pancreas transplant patients at high risk.

Pediatric Patients (1.2 Prevention of CMV disease in kidney and heart transplant patients at high risk.

DOSAGE AND ADMINISTRATION			
	Adult Dosage (2.2)		
Treatment of CMV retinitis	Induction: 900 mg (two 450 mg tablets) twice a day for 21 days		
	Maintenance: 900 mg (two 450 mg tablets) once a day		
Prevention of CMV disease in heart or kidney-pancreas transplant patients	900 mg (two 450 mg tablets) once a day within 10 days of transplantation until 100 days post-transplantation		
Prevention of CMV disease in kidney transplant patients	900 mg (two 450 mg tablets) once a day within 10 days of transplantation until 200 days post-transplantation		
Pediatric Dosage (2.3)			
Prevention of CMV disease in kidney transplant patients 4 months to 16 years of age	Dose once a day within 10 days of transplantation until 200 days post- transplantation according to dosage algorithm (note the calculation of creatinine clearance using a modified Schwartz formula in children)		
Prevention of CMV disease in heart transplant patients 1 month to 16 years of age	Dose once a day within 10 days of transplantation until 100 days post- transplantation according to dosage algorithm (note the calculation of creatinine clearance using a modified Schwartz formula in children)		

- VALCYTE for oral solution and tablets should be taken with food. (2.1, 12.3) VALCYTE tablets should not be broken or crushed. (2.6)
- Adult patients should use VALCYTE tablets, not VALCYTE for oral solution. (2.1)
- Adults with renal impairment: Adjust dose based on creatinine clearance. For adult patients receiving hemodialysis
- a dose recommendation cannot be given. (2.5, 8.6, 12.3) - DOSAGE FORMS AND STRENGTHS

Tablets: 450 mg. (3)

Oral Solution: 50 mg per mL. (3)

--- CONTRAINDICATIONS --Hypersensitivity to valganciclovir or ganciclovir. (4

--- WARNINGS AND PRECAUTIONS ----• Acute renal failure: Acute renal failure may occur in elderly patients (with or without reduced renal function), patients who receive concomitant nephrotoxic drugs, or inadequately hydrated patients. Use with caution in elderly patients or those taking nephrotoxic drugs, reduce dosage in patients with renal impairment, and monitor renal

-- ADVERSE REACTIONS ---Adult patients: Most common adverse reactions and laboratory abnormalities (reported in at least one indication by greater than or equal to 20% of patients) are diarrhea, pyrexia, fatigue, nausea, tremor, neutropenia, anemia,

leukopenia, thrombocytopenia, headache, insomnia, urinary tract infection, and vomiting. (6.1)

Pediatric patients: Most common adverse reactions and laboratory abnormalities (reported in greater than or equal to 20% of pediatric solid organ transplant recipients) are diarrhea, pyrexia, upper respiratory tract infection, urinary tract infection, vomiting, neutropenia, leukopenia, and headache. (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact Genentech at 1-888-835-2555 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

---- DRUG INTERACTIONS

Imipenem-cilastatin: Seizures were reported in patients receiving ganciclovir and imipenem-cilastatin. Concomitant use is not recommended unless the potential benefits outweigh the risks. (7)
 Cyclosporine or amphotericin B: When coadministered with valganciclovir, the risk of nephrotoxicity may be

increased, Monitor renal function, (5.2, 7) enolate mofetil (MMF): When coadministered with valganciclovir, the risk of hematological and renal toxicity may be increased. Monitor for ganciclovir and MMF toxicity. (7)

Other drugs associated with myelosuppression or nephrotoxicity: Due to potential for increased toxicity, consider for concomitant use with valganciclovir only if the potential benefits are judged to outweigh the risks. (7)
Didanosine: Ganciclovir coadministered with didanosine may increase didanosine levels. Monitor for didanosine toxicity (e.g., pancreatitis), (7)

 Probenecid: May increase ganciclovir levels. Monitor for evidence of ganciclovir toxicity. (7) ----- USE IN SPECIFIC POPULATIONS

Lactation: Breastfeeding is not recommended with use of VALCYTE. (8.2

See 17 for PATIENT COUNSELING INFORMATION and FDA-approved patient labeling

FULL PRESCRIBING INFORMATION: CONTENTS*

WARNING: HEMATOLOGIC TOXICITY, IMPAIRMENT OF FERTILITY, FETAL TOXICITY, MUTAGENESIS AND CARCINOGENESIS

INDICATIONS AND USAGE Adult Patients

DOSAGE AND ADMINISTRATION

General Dosing Information Recommended Dosage in Adult Patients with Normal Renal Function

Preparation of VALCYTE for Oral Solution

2.5 Dosage Recommendation for Adult Patients with Renal Impairment
2.6 Handling and Disposal

DOSAGE FORMS AND STRENGTHS

CONTRAINDICATIONS WARNINGS AND PRECAUTIONS Hematologic Toxicity

Acute Renal Failure Impairment of Fertility Fetal Toxicity Mutagenesis and Carcinogenesis

ADVERSE REACTIONS Clinical Trials Experience

DRUG INTERACTIONS USE IN SPECIFIC POPULATIONS

Females and Males of Reproductive Potential Pediatric Use

Renal Impairment

8.7 Hepatic Impairment OVERDOSAGE DESCRIPTION 12 CLINICAL PHARMACOLOGY

Mechanism of Action 12.3 Pharmacokinetics

13 NONCLINICAL TOXICOLOGY 3.1 Carcinogenesis, Mutagenesis, Impairment of Fertility 14 CLINICAL STUDIES

14.1 Adult Patients14.2 Pediatric Patients

15 REFERENCES

HOW SUPPLIED/STORAGE AND HANDLING PATIENT COUNSELING INFORMATION

*Sections or subsections omitted from the full prescribing information are not listed.

FULL PRESCRIBING INFORMATION

WARNING: HEMATOLOGIC TOXICITY, IMPAIRMENT OF FERTILITY, FETAL TOXICITY. MUTAGENESIS AND CARCINOGENESIS

tologic Toxicity: Severe leukopenia, neutropenia, anemia, thrombocytopenia, pand bone marrow failure including aplastic anemia have been reported in patients treated with VALCYTE (see Impairment of Fertility: Based on animal data and limited human data, VALCYTE may cause temporary or

permanent inhibition of spermatogenesis in males and suppression of fertility in females *[see Warnings* Fetal Toxicity: Based on animal data, VALCYTE has the potential to cause birth defects in humans [see

Warnings and Precautions (5.4)].
Mutagenesis and Carcinogenesis: Based on animal data, VALCYTE has the potential to cause cancers in humans [see Warnings and Precautions (5.5)].

INDICATIONS AND USAGE

1.1 Adult Patients

<u>Treatment of Cytomegalovirus (CMV) Retinitis:</u> VALCYTE is indicated for the treatment of CMV retinitis in patients with acquired immunodeficiency syndrome (AIDS) [see Clinical Studies (14.1)].

Prevention of CMV Disease: VALCYTE is indicated for the prevention of CMV disease in kidney, heart, and kidney-pancreas transplant patients at high risk (Donor CMV seropositive/Recipient CMV seronegative [D+/R-]) [see Clinical

1.2 Pediatric Patients

<u>Prevention of CMV Disease:</u> VALCYTE is indicated for the prevention of CMV disease in kidney transplant patients (4 months to 16 years of age) and heart transplant patients (1 month to 16 years of age) at high risk *[see Clinical* Studies (14.2)1.

2 DOSAGE AND ADMINISTRATION 2.1 General Dosing Information

Adult nations should use VAI CYTE tablets not VAI CYTE for oral solution.

VALCYTE for oral solution and tablets should be taken with food [see Clinical Pharmacology (12.3)]. VALCYTE for oral solution (50 mg/mL) must be prepared by the pharmacist prior to dispensing to the patient [see Dosage and Administration (2.4)]

2.2 Recommended Dosage in Adult Patients with Normal Renal Function

For dosage recommendations in adult patients with renal impairment [see Dosage and Administration (2.5)].

<u>Treatment of CMV Retinitis:</u>
• Induction: The recommended dosage is 900 mg (two 450 mg tablets) taken orally twice a day for 21 days.
• Maintenance: Following induction treatment, or in adult patients with inactive CMV retinitis, the recommended

Prevention of CMV Disease:

For adult patients who have received a heart or kidney-pancreas transplant, the recommended dosage is 900 mg (two 450 mg tablets) taken orally once a day starting within 10 days of transplantation until 100 days

For adult patients who have received a kidney transplant, the recommended dosage is 900 mg (two 450 mg

2.3 Recommended Dosage in Pediatric Patients

dosage is 900 mg (two 450 mg tablets) taken orally once a day.

<u>Prevention of CMV Disease in Pediatric Kidney Transplant Patients:</u> For pediatric kidney transplant patients 4 months to 16 years of age, the recommended once daily mg dose (7 x BSA x CrCl) should start within 10 days of post-

 $\begin{array}{l} \underline{Prevention \ of \ CMV \ Disease \ in \ Pediatric \ Heart \ Transplant \ Patients:} \ For \ pediatric \ heart \ transplant \ patients \ 1 \ month \ to \ 16 \ years \ of \ age, the \ recommended \ once \ daily \ mg \ dose \ (7 \times BSA \times CrCI) \ should \ start \ within \ 10 \ days \ of \ transplant \ attitude \ days \ of \ transplant \ days \ of \ \ of \ transplant \ days \ of \ transplan$ until 100 days post-transplantation.

The recommended once daily dosage of VALCYTE is based on body surface area (BSA) and creatinine clearance (CrCl) derived from a modified Schwartz formula, and is calculated using the equation below:

Pediatric Dose (mg) = 7 x BSA x CrCl (calculated using a modified Schwartz formula). If the calculated Schwartz creatinine clearance exceeds 150 mL/min/1.73m², then a maximum value of 150 mL/min/1.73m² should be used in the equation. The k values used in the modified Schwartz formula are based on pediatric patient age, as shown Mosteller BSA $(m^2) = \sqrt{\frac{\text{Height}(cm) \times \text{Weight}(kg)}{3600}}$

Schwartz Creatinine Clearance (mL / min / 1.73 m^2) = $\frac{k \ x \ \text{Height}(cm)}{\text{Serum Creatinine (mg / dL)}}$

Table 1 k Values According to Pediatric Patient Age* k value Pediatric Patient Age Infants less than 1 year of age with low birth weight for gestational age Infants less than 1 year of age with birth weight appropriate for gestational age Children aged 1 to less than 2 years

Boys aged 2 to less than 13 years Girls aged 2 to less than 16 years Boys aged 13 to 16 years The k values provided are based on the Jaffe method of measuring serum creatinine, and may require correction when enzymati

Monitor serum creatinine levels regularly and consider changes in height and body weight and adapt the dose as

All calculated doses should be rounded to the nearest 25 mg increment for the actual deliverable dose. The oral dispenser is graduated in 0.5 mL increments. A 50 mg dose is equivalent to 1 mL. if the calculated dose exceeds 900 mg, a maximum dose of 900 mg should be administered. VALCYTE for oral solution is the preferred formulation since it provides the ability to administer a dose calculated according to the formula above; however, VALCYTI tablets may be used if the calculated doses are within 10% of available tablet strength (450 mg). For example, in the calculated dose is between 405 mg and 495 mg, one 450 mg tablet may be taken. Before prescribing VALCYTE tablets, pediatric patients should be assessed for the ability to swallow tablets.

2.4 Preparation of VALCYTE for Oral Solution

Wearing disposable gloves is recommended during reconstitution and when wiping the outer surface of the bottle/cap and the table after reconstitution. Prior to dispensing to the patient, VALCYTE for oral solution must be prepared by the pharmacist as follows [see How Supplied/Storage and Handling (16)]:

 Measure 91 mL of purified water in a graduated cylinder.
 Measure 91 mL of purified water in a graduated cylinder.
 Shake the VALCYTE bottle to loosen the powder. Remove the child resistant bottle cap and add approximately half the total amount of water for constitution to the bottle and shake the closed bottle well for about 1 minute. Add the emainder of water and shake the closed bottle well for about 1 minute. This prepared solution contains 50 mg of

valganciclovir free base per 1 mL. Remove the child resistant bottle cap and push the bottle adapter into the neck of the bottle

 Close bottle with child resistant bottle cap tightly. This will assure the proper seating of the bottle adapter in the bottle and child resistant status of the cap. • Store constituted oral solution under refrigeration at 2°C to 8°C (36°F to 46°F) for no longer than 49 days. Write the discard date of the constituted oral solution on the bottle label.

The patient package insert, which includes the dosing instructions for patients, and 2 oral dispensers should be dispensed to the patient [see Patient Counseling Information (17)].

2.5 Dosage Recommendation for Adult Patients with Renal Impairment

Serum creatinine levels or estimated creatinine clearance should be monitored regularly during treatment. Dosage recommendations for adult patients with reduced renal function are provided in Table 2. For adult patients on hemodialysis (CrCl less than 10 mL/min), a dose recommendation for VALCYTE cannot be given [see Use in Specific Populations (8.5, 8.6), Clinical Pharmacology (12.3)].

Table 2 Decore Decommondations for Adult Deticate with Impaired Devel Constitute

	lable 2 Dosage Recommendations for Adult Patients with Impaired Renai Function			
	VALCYTE 450 mg Tablets			
CrCl* (mL/min) Induction Dose Maintenance/Prevention Dos				
	≥ 60	900 mg twice daily	900 mg once daily	
	40 – 59	450 mg twice daily	450 mg once daily	
	25 – 39	450 mg once daily	450 mg every 2 days	
	10 – 24	450 mg every 2 days	450 mg twice weekly	
	< 10 (on hemodialysis)	not recommended	not recommended	

n estimated creatinine clearance in adults is calculated from serum creatinine by the following formulas:

(140 - age [years]) x (body weight [kg]) For males = (72) x (serum creatinine [mg/dL])

For females = 0.85 x male value

Dosing in pediatric patients with renal impairment can be done using the recommended equations because CrCl is a

2.6 Handling and Disposal

Caution should be exercised in the handling of VALCYTE tablets and VALCYTE for oral solution. Tablets should not be broken or crushed. Because valganciclovir is considered a potential teratogen and carcinogen in humans, caution should be observed in handling broken tablets, the powder for oral solution, and the constituted oral solution [see Warnings and Precautions (5.4, 5.5)]. Avoid direct contact with broken or crushed tablets, the powder for oral solution, and the constituted oral solution with skin or mucous membranes. If such contact occurs, wash thoroughly with soap and water, and rinse eyes thoroughly with plain water.

Handle and dispose VALCYTE according to guidelines for antineoplastic drugs because ganciclovir shares some of the

DOSAGE FORMS AND STRENGTHS

 VALCYTE tablets: 450 mg, pink, film-coated convex oval tablets with "VGC" on one side and "450" on the other VALCYTE for oral solution: 50 mg per mL, supplied as a white to slightly yellow powder for constitution, forming

a colorless to brownish yellow tutti-frutti flavored solution. Available in glass bottles containing approximately

CONTRAINDICATIONS

VALCYTE is contraindicated in patients who have had a demonstrated clinically significant hypersensitivity reaction (e.g., anaphylaxis) to valganciclovir, ganciclovir, or any component of the formulation [see Adverse Reactions (6.1)].

Hematologic Toxicity

Severe leukopenia neutropenia anemia thrombocytopenia pancytopenia and hope marrow failure including aplastic anemia have been reported in patients treated with VALCYTE or ganciclovir. VALCYTE should be avoided if the absolute neutrophil count is less than 500 cells/µL, the platelet count is less than 25,000/µL, or the hemoglobin is less than 8 g/dL, VALCYTE should also be used with caution in patients with pre-existing cytopenias and in patients suppressive drugs or irradiation. Cytopenia may occur at any time during treatment and may worse with continued dosing. Cell counts usually begin to recover within 3 to 7 days after discontinuing drug. In patients with severe leukopenia, neutropenia, anemia and/or thrombocytopenia, treatment with hematopoietic growth factors

Due to the frequency of neutropenia, anemia, and thrombocytopenia in patients receiving VALCYTE *Isee Adverse* Reactions (6.1)], complete blood counts with differential and platelet counts should be performed frequently, especially in infants, in patients with renal impairment, and in patients in whom ganciclovir or other nucleoside analogues have previously resulted in leukopenia, or in whom neutrophil counts are less than 1000 cells/µL at the beginning of treatment. Increased monitoring for cytopenias may be warranted if therapy with oral ganciclovir is changed to VALCYTE because of increased plasma concentrations of ganciclovir after VALCYTE administration [see Clinical Pharmacology (12.3)].

5.2 Acute Renal Failure

Acute renal failure may occur in:

• Elderly patients with or without reduced renal function. Caution should be exercised when administering VALCYTE to geriatric patients, and dosage reduction is recommended for those with impaired renal function [see Dosage and Administration (2.5), Use in Specific Populations (8.5, 8.6)].

Patients receiving potential nephrotoxic drugs. Caution should be exercised when administering VALCYTE to

patients receiving potential nephrotoxic drugs.

• Patients without adequate hydration. Adequate hydration should be maintained for all patients.

5.3 Impairment of Fertility

or permanent inhibition of spermatogenesis in males, and may cause suppression of fertility in females. Advise patients that fertility may be impaired with use of VALCYTE [see Use in Specific Populations (8.1, 8.3), Nonclinical oxicoloav (13.1)1. 5.4 Fetal Toxicity

Based on animal data and limited human data, VALCYTE at the recommended human doses may cause temporary

Ganciclovir may cause fetal toxicity when administered to pregnant women based on findings in animal studies. When given to pregnant rabbits at dosages resulting in 2 times the human exposure (based on AUC), ganciclovir caused malformations in multiple organs of the fetuses. Maternal and fetal toxicity were also observed in pregnant mice and rabbits. Therefore, VALCYTE has the potential to cause birth defects. Pregnancy should be avoided in femal patients taking VALCYTE and in females with male partners taking VALCYTE. Females of reproductive potential should be avoided in females with male partners taking VALCYTE. be advised to use effective contraception during treatment and for at least 30 days following treatment with VALCYTI because of the potential risk to the fetus. Similarly, males should be advised to use condoms during and for at least ment with VALCYTE [see Dosage and Administration (2.6), Use in Specific Pop 8.3), Nonclinical Toxicology (13.1)].

5.5 Mutagenesis and Carcinogenesis

Animal data indicate that ganciclovir is mutagenic and carcinogenic. VALCYTE should therefore be considered a potential carcinogen in humans [see Dosage and Administration (2.6), Nonclinical Toxicology (13.1)].

ADVERSE REACTIONS The following serious adverse reactions are discussed in greater detail in other sections of the labeling:

 Hematologic Toxicity [see Warnings and Precautions (5.1)].
 Acute Renal Failure [see Warnings and Precautions (5.2)]. Impairment of Fertility [see Warnings and Precautions (5.3)]

Fetal Toxicity [see Warnings and Precautions (5.4)].
Mutagenesis and Carcinogenesis [see Warnings and Precautions (5.5)].

The most common adverse reactions and laboratory abnormalities reported in at least one indication by greater than or equal to 20% of adult patients treated with VALCYTE tablets are diarrhea, pyrexia, fatigue, nausea, tremor, neutropenia, anemia, leukopenia, thrombocytopenia, headache, insomnia, urinary tract infection, and vomiting. The most common reported adverse reactions and laboratory abnormalities reported in greater than or equal to 20% of pediatric solid organ transplant recipients treated with VALCYTE for oral solution or tablets are diarrhea, pyrexia, upper respiratory tract infection, urinary tract infection, vomiting, neutropenia, leukopenia, and headache.

6.1 Clinical Trials Experience

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect rates observed in practice. Valganciclovir, a prodrug of ganciclovir, is rapidly converted to ganciclovir after oral administration. Adverse reactions

known to be associated with ganciclovir usage can therefore be expected to occur with VALCYTE eatment of CMV Retinitis in AIDS Patients: In a clinical study for the treatment of CMV retinitis in HIV-infected patients, the adverse reactions reported by patients receiving VALCYTE tablets (n=79) or intravenous ganciclovir (n=79) for 28 days of randomized therapy (21 days induction dose and 7 days maintenance dose), respectively, included diarrhea (16%, 10%), nausea (8%, 14%), and headache (9%, 5%). The incidence of adverse reactions was

similar between the group who received VALCYTE tablets and the group who received intravenous ganciclovir. The frequencies of neutropenia (ANC less than 500/µL) were 11% for patients receiving VALCYTE tablets compared with 13% for patients receiving intravenous ganciclovir. Anemia (Hgb less than 8 g/dL) occurred in 8% of patients in each group. Other laboratory abnormalities occurred with similar frequencies in the two groups. Adverse reactions and laboratory abnormalities are available for 370 patients who received maintenance therapy with VALCYTE tablets 900 mg once daily in two open-label clinical trials. Approximately 252 (68%) of these patients received VALCYTE tablets for more than nine months (maximum duration was 36 months). Table 3 and Table 4 show

pooled selected adverse reactions and abnormal laboratory values from these patients Table 3 Pooled Selected Adverse Reactions Reported in greater than or equal to 5% of Patients who Received VALCYTE Tablets Maintenance Therapy for CMV Retinitis

	Patients with CMV Retinitis
Adverse Reactions according to Body System	VALCYTE Tablets (N=370) %
Gastrointestinal system	
Diarrhea	41
Nausea	30
Vomiting	21
Abdominal pain	15
General disorders and administrative site conditions	
Pyrexia	31
Nervous system disorders	
Headache	22
Insomnia	16
Neuropathy peripheral	9
Paresthesia	8
Eye disorders	
Retinal detachment	15

Table 4 Pooled Selected Laboratory Abnormalities Reported in Patients Who Received VALCYTE Tablets

	Patients with CMV Retinitis
Laboratory Abnormalities	VALCYTE Tablets (N=370) %
Neutropenia: ANC/µL < 500 500 - < 750 750 - < 1000	19 17 17
Anemia: Hemoglobin g/dL < 6.5 6.5 - < 8.0 8.0 - < 9.5	7 13 16
Thrombocytopenia: Platelets/μL < 25000 25000 - < 50000 50000 - < 100000	4 6 22
Serum Creatinine: mg/dL > 2.5 > 1.5 – 2.5	3 12

Prevention of CMV Disease in Solid Organ Transplant Patients: Table 5 shows selected adverse reactions regardless of severity with an incidence of greater than or equal to 5% from a clinical trial (up to 28 days after study treatment) where heart, kidney, kidney-pancreas and liver transplant patients received VALCYTE tablets (N=244) or oral panciclovir (N=126) until Day 100 post-transplant. The majority of the adverse reactions were of mild or moderate

Table 5 Percentage of Selected Grades 1–4 Adverse Reactions Reported in greater than or equal to 5% of Adult Patients From a Study of Solid Organ Transplant Patients

Adverse Reactions	VALCYTE Tablets (N=244) %	Oral Ganciclovir (N=126) %
ointestinal disorders		
nea	30	29
ea	23	23
ing	16	14
ous system disorders		
ors	28	25
ache	22	27
nnia	20	16
ral disorders and nistration site conditions		
ia	13	14

Table 6 shows selected adverse reactions regardless of severity with an incidence of greater than or equal to 5% from another clinical trial where kidney transplant patients received either valganciclovir once daily starting within 10 days post-transplant until Day 100 post-transplant followed by 100 days of placebo or valganciclovir once daily until ay 200 post-transplant. The overall safety profile of VALCYTE did not change with the extension of prophylaxis until Table 6 Percentage of Selected Grades 1–4 Adverse Reactions Reported in greater than or equal to

5% of Adult Patients from a Study of Kidney Transplant Patien

Adverse Reactions	VALCYTE Tablets Day 100 Post-transplant (N=164) %	VALCYTE Tablets Day 200 Post-transplant (N=156) %
Gastrointestinal disorders		
Diarrhea	26	31
Nausea	11	11
Vomiting	3	6
Nervous system disorders		
Tremors	12	17
Headache	10	6
Insomnia	7	6
General disorders and administration site conditions		
Pyrexia	12	9

Table 7 and Table 8 show selected laboratory abnormalities reported with VALCYTE tablets in two trials in solid organ

Laboratory Abnormalities	VALCYTE Tablets (N=244) %	Ganciclovir Capsules (N=126) %
Neutropenia: ANC/μL		
< 500	5	3
500 - < 750	3	2
750 – < 1000	5	2
Anemia: Hemoglobin g/dL		
< 6.5	1	2
6.5 - < 8.0	5	7
8.0 - < 9.5	31	25
hrombocytopenia: Platelets/µL		
< 25000	0	2
25000 - < 50000	1	3
50000 - < 100000	18	21
Serum Creatinine: mg/dL		
> 2.5	14	21
> 1.5 – 2.5	45	47

Table 8 Selected Laboratory Abno	ormalities Reported in a Study of Adult Kidney Transplant Patients*			
Laboratory Abnormalities	VALCYTE Tablets Day 100 Post-transplant (N=164) %	VALCYTE Tablets Day 200 Post-transplant (N=156) %		
Neutropenia: ANC/μL < 500 500 – < 750 750 – < 1000	9 6 7	10 6 5		
Anemia: Hemoglobin g/dL < 6.5 6.5 - < 8.0 8.0 - < 9.5	0 5 17	1 1 15		
Thrombocytopenia: Platelets/μL < 25000 25000 - < 50000 50000 - < 100000	0 1 7	0 0 3		
Serum Creatinine: mg/dL > 2.5 > 1.5 – 2.5	17 50	14 48		

Laboratory abnormalities are those reported by investigators Other adverse drug reactions from VALCYTE in clinical trials in CMV retinitis and solid organ transplant patients
Other adverse drug reactions with VALCYTE in clinical trials in either patients with CMV retinitis or solid organ transplant patients that occurred in at least 5% of patients are listed below.

Eve disorders: retinal detachment, eve pain

Gastrointestinal disorders: dyspepsia, constipation, abdominal distention, mouth ulceration General disorders and administration site conditions: fatique, pain, malaise, asthenia, chills, peripheral edema Hepatobiliary disorders: hepatic function abnormal

Infections and infestations: candida infections including oral candidiasis, upper respiratory tract infection, influenza, urinary tract infection, pharyngitis/nasopharyngitis, postoperative wound infection

Injury, poisoning, and procedural complications: postoperative complications, wound secretion Metabolic and nutrition disorders: decreased appetite, hyperkalemia, hypophosphatemia, weight decreased

Musculoskeletal and connective tissue disorders: back pain, myalgia, arthralgia, muscle spasms

Psychiatric disorders: depression, anxiety

Nervous system disorders: insomnia, neuropathy peripheral, dizziness

Respiratory, thoracic and mediastinal disorders: cough, dyspnea

Renal and urinary disorders: renal impairment, creatinine clearance renal decreased, blood creatinine increased,

Skin and subcutaneous tissues disorders: dermatitis, night sweats, pruritus Vascular disorders: hypotension

Other adverse reactions with VALCYTE in clinical trials in either patients with CMV retinitis or solid organ transplant patients that occurred in less than 5% of patients are listed below.

Blood and lymphatic disorders: febrile neutropenia, pancytopenia, bone marrow failure (including aplastic anemia) Cardiovascular disorders: arrhythmia

Eve disorders: macular edema Gastrointestinal disorders: pancreatitis

Ear and labyrinth disorders: deafness

Hemorrhage: potentially life-threatening bleeding associated with thrombocytopenia

Immune system disorders: hypersensitivity Infections and infestations: cellulitis, sepsis Injury, poisoning, and procedural complications; postoperative pain, wound dehiscence

Investigations: aspartate aminotransferase increased, alanine aminotransferase increased Musculoskeletal and connective tissue disorders: limb pain

Nervous system disorders: seizure, dysquesia (taste disturbance)

Psychiatric disorders: confusional state, agitation, psychotic disorder, hallucinations Renal and urinary disorders: renal failure

Adverse Reactions in Pediatric Patients: VALCYTE for oral solution and tablets have been studied in 179 pediatric solid organ transplant patients who were at risk for developing CMV disease (aged 3 weeks to 16 years) and in 24 neonates with symptomatic congenital CMV disease (aged 8 to 34 days), with duration of ganciclovir exposure ranging from 2 to 200 days [see Use in Specific

Prevention of CMV Disease in Pediatric Solid Organ Transplant Patients: The most frequently reported adverse reactions (greater than 10% of patients), regardless of seriousness, in pediatric solid organ transplant patients taking VALCYTE until Day 100 post-transplant were diarrhea, pyrexia, upper respiratory tract infection, vomiting, anemia, neutropenia, constipation and nausea. The most frequently reported adverse reactions (greater than 10%) of patients) in pediatric kidney transplant patients treated with valganciclovir until Day 200 post-transplant were upper respiratory tract infection, urinary tract infection, diarrhea, leukopenia, neutropenia, headache, abdominal pain, tremor, pyrexia, anemia, blood creatinine increased, vomiting, and hematuria.

In general, the safety profile was similar in pediatric patients compared to that observed in adult patients. However the rates of certain adverse reactions, and laboratory abnormalities, such as upper respiratory tract infection, pyrexia, nasopharyngitis, anemia, and abdominal pain were reported more frequently in pediatric patients than in adults [see Use in Specific Populations (8.4), Clinical Studies (14.2). Neutropenia was reported at a higher incidence in the two in the pediatric population.

The overall safety profile of VALCYTE was similar with the extension of prophylaxis until Day 200 post-transplant in high risk pediatric kidney transplant patients. However, the incidence of severe neutropenia (ANC $< 500/\mu$ L) was nigher in pediatric kidney transplant patients treated with VALCYTE until Day 200 (17/57, 30%) compared to pediatric kidney transplant patients treated until Day 100 (3/63, 5%). There were no differences in the incidence of severe (Grade 4) anemia or thrombocytopenia in patients treated 100 or 200 days with VALCYTE.

6.2 Postmarketing Experience

The following adverse reactions have been identified during post-approval use of VALCYTE. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure. As VALCYTE is rapidly and extensively converted to ganciclovir, any adverse reactions associated with ganciclovir might also occur with valganciclovir

Agranulocytosis Granulocytopenia

In general, the adverse reactions reported during the postmarketing use of VALCYTE were similar to those identified during the clinical trials.

7 DRUG INTERACTIONS

In vivo drug-drug interaction studies were not conducted with valganciclovir. However, because valganciclovir is rapidly and extensively converted to ganciclovir, drug-drug interactions associated with ganciclovir will be expected for VALCYTE. Drug-drug interaction studies with ganciclovir were conducted in patients with normal renal function Following concomitant administration of VALCYTE and other renally excreted drugs, patients with impaired renal function may have increased concentrations of ganciclovir and the coadministered drug. Therefore, these patients should be closely monitored for toxicity of ganciclovir and the coadministered drug. Established and other potentially significant drug interactions conducted with ganciclovir are listed in Table 9.

Change in the Concentration of Clinical Comment

Table 9 Established and Other Potentially Significant Drug Interactions with Ganciclovir

Concomitant Drug	Ganciclovir or Concomitant Drug	
Imipenem-cilastatin	Unknown	Coadministration with imipenem- cilastatin is not recommended because generalized seizures have been reported in patients who received ganciclovir and imipenem- cilastatin.
Cyclosporine or amphotericin B	Unknown	Monitor renal function when VALCYTE is coadministered with cyclosporine or amphotericin B because of potential increase in serum creatinine [see Warnings and Precautions (5.2)].
Mycophenolate mofetil (MMF)	⇔ Ganciclovir (in patients with normal renal function) ⇔ MMF (in patients with normal renal function)	Based on increased risk, patients should be monitored for hematological and renal toxicity.
Other drugs associated with myelosuppression or nephrotoxicity (e.g., adriamycin, dapsone, doxorubicin, flucytosine, hydroxyurea, pentamidine, tacrolimus, trimethoprim/ sulfamethoxazole, vinblastine, vincristine, and zidovudine)	Unknown	Because of potential for higher toxicity, coadministration with VALCYTE should be considered only if the potential benefits are judged to outweigh the risks.
Didanosine	↔ Ganciclovir ↑ Didanosine	Patients should be closely monitored for didanosine toxicity (e.g., pancreatitis)
Probenecid	↑ Ganciclovir	VALCYTE dose may need to be reduced. Monitor for evidence of ganciclovir toxicity.

8 USE IN SPECIFIC POPULATIONS 8.1 Pregnancy

After oral administration, valganciclovir (prodrug) is converted to ganciclovir (active drug) and, therefore, VALCYTE is

expected to have reproductive toxicity effects similar to ganciclovir. In animal studies, ganciclovir caused materna and fetal toxicity and embryo-fetal mortality in pregnant mice and rabbits as well as teratogenizity in rabbits a exposures two-times the human exposure. There are no available human data on use of VALCYTE or ganciclov in pregnant women to establish the presence or absence of drug-associated risk. The background risk of major birth defects and miscarriage for the indicated populations is unknown. However, the background risk in the U.S. general population of major birth defects is 2–4% and the risk of miscarriage is 15–20% of clinically recognized regnancies. Advise pregnant women of the potential risk to the fetus [see Warnings and Precautions (5.3), Use in Specific Populations (8.3)]. Clinical Considerations
Disease-associated maternal and/or embryo/fetal risk

Most maternal CMV infections are asymptomatic or they may be associated with a self-limited mononucleosis-like

syndrome. However, in immunocompromised patients (i.e., transplant patients or patients with AIDS) CMV infections may be symptomatic and may result in significant maternal morbidity and mortality. The transmission of CMV to the fetus is a result of maternal viremia and transplacental infection. Perinatal infection can also occur from exposure of

the neonate to CMV shedding in the genital tract. Approximately 10% of children with congenital CMV infection are

symptomatic at birth. Mortality in these infants is about 10% and approximately 50-90% of symptomatic survivin newborns experience significant morbidity, including mental retardation, sensorineural hearing loss, microcephal seizures, and other medical problems. The risk of congenital CMV infection resulting from primary maternal CMV infection may be higher and of greater severity than that resulting from maternal reactivation of CMV infection.

Doses resulting in two-times the human exposure of ganciclovir (based on the human AUC following a single intravenous infusion of 5 mg per kg of ganciclovir) resulted in maternal and embryo-fetal toxicity in pregnant mice and rabbits as well as teratogenicity in the rabbits. Fetal resorptions were present in at least 85% of rabbits and mice. Rabbits showed increased embryo-fetal mortality, growth retardation of the fetuses and structural abnormalities of multiple organs of the fetuses including the palate (cleft palate), eyes (anophthalmia/microphthalmia), brain (hydrocephalus), jaw (brachygnathia), kidneys and pancreas (aplastic organs). Increased embryo-fetal mortality was also seen in mice. Daily intravenous doses of approximately 1.7 times the human exposure (based on AUC) administered to female mice prior to mating, during gestation, and during lactation caused hypoplasia of the testes and seminal vesicles in the male offspring, as well as pathologic changes in the nonglandular region o

Data from an ex-vivo human placental model showed that ganciclovir crosses the human placenta. The transfer occurred by passive diffusion and was not saturable over a concentration range of 1 to 10 mg/mL.

8.2 Lactation

Risk Summary

No data are available regarding the presence of valganciclovir (prodrug) or ganciclovir (active drug) in human milk, the effects on the breastfed infant, or the effects on milk production. Animal data indicate that ganciclovir is excreted in the milk of lactating rats. The Centers for Disease Control and Prevention recommend that HIV-infected mothers not breastfeed their infants to avoid risking postnatal transmission of HIV. Advise nursing mothers that breastfeeding is not recommended during treatment with VALCYTE because of the potential for serious adverse events in nursing infants and because of the potential for transmission of HIV [see Boxed Warning, Warnings and Precautions (5.1, 5.3,

8.3 Females and Males of Reproductive Potential

nales of reproductive potential should undergo pregnancy testing before initiation of VALCYTE [see Use in Specific Contraception

Because of the mutagenic and teratogenic potential of VALCYTE, females of reproductive potential should be advised

o use effective contraception during treatment and for at least 30 days following treatment with VALCYTE [see

Oosage and Administration (2.6), Warnings and Precautions (5.4, 5.5), Nonclinical Toxicology (13.1)].

Because of its mutagenic potential, males should be advised to use condoms during and for at least 90 days following, treatment with VALCYTE [see Dosage and Administration (2.6), Warnings and Precautions (5.3, 5.5),

Intertility
VALCYTE at the recommended doses may cause temporary or permanent female and male infertility [see Warnings and Precautions (5.3), Nonclinical Toxicology (13.1)].

Pregnancy Testing

In a small, open-label, non-randomized clinical study, adult male renal transplant patients receiving VALCYTE for CMV prophylaxis for up to 200 days post-transplantation were compared to an untreated control group. Patients were followed-up for six months after VALCYTE discontinuation. Among 24 evaluable patients in the VALCYTE group, the mean sperm density at the end of treatment visit decreased by 11 million/mL from baseline; whereas, among 14 evaluable patients in the control group the mean sperm density increased by 33 million/mL. However, at the follow-up visit among 20 evaluable patients in the VALCYTE group the mean sperm density was comparable to that observed among 10 evaluable patients in the untreated control group (the mean sperm density at the end of follow-up visit increased by 41 million/mL from baseline in the VALCYTE group and by 43 million/mL in the untreated group)

VALCYTE for oral solution and tablets are indicated for the prevention of CMV disease in pediatric kidney transplant valor for the first sound and a dates at a limited of the prevention of the seaso in pediatric Adrey at a risk for developing CMV disease [see Indications and Usage (1.2), Dosage and Administration (2.3)].

The use of VALCYTE for oral solution and tablets for the prevention of CMV disease in pediatric kidney transplant patients 4 months to 16 years of age is based on two single-arm, open-label, non-comparative studies in patients 4 nonths to 16 years of age. Study 1 was a safety and pharmacokinetic study in pediatric solid organ transplant patients idney, liver, heart, and kidney/pancreas). VALCYTE was administered once daily within 10 days of trans for a maximum of 100 days post-transplantation. Study 2 was a safety and tolerability study where VALCYTE

The use of VALCYTE for oral solution and tablets for the prevention of CMV disease in pediatric heart transplant patients 1 month to 16 years of age is based on two studies (Study 1 described above and Study 3) and was supported by previous demonstration of efficacy in adult patients [see Clinical Pharmacology (12.3), Clinical Studies (14.2)]. Study 3 was a pharmacokinetic and safety study of VALCYTE in pediatric heart transplant patients less than 4 months of age who received a single dose of VALCYTE oral solution on each of two consecutive days. A

The safety and efficacy of VALCYTE for oral solution and tablets have not been established in children for prevention of CMV disease in pediatric liver transplant patients, in kidney transplant patients less than 4 months of age, in heart transplant patients less than 1 month of age, in pediatric AIDS patients with CMV retinitis, and in infants with

A pharmacokinetic and pharmacodynamic evaluation of VALCYTE for oral solution was performed in 24 neonates with congenital CMV infection involving the central nervous system. All patients were treated for 6 weeks with a combination of intravenous ganciclovir 6 mg per kg twice daily or VALCYTE for oral solution at doses ranging from 14 mg per kg to 20 mg per kg twice daily. The pharmacokinetic results showed that in infants greater than 7 days to 3 months of age, a dose of 16 mg per kg twice daily of VALCYTE for oral solution provided ganciclovir systemic exposures (median AUC_{b-12} =23.6 [range 16.8–35.5] mcg·h/mL; n=6) comparable to those obtained in infants up to 3 months of age from a 6 mg per kg dose of intravenous ganciclovir twice daily (AUC_{0-12i}=25.3 [range 2.4–89.7] mcg·h/mL; n=18) or to the ganciclovir systemic exposures obtained in adults from a 900 mg dose of VALCYTE tablets twice daily. However, the efficacy and safety of intravenous ganciclovir and of VALCYTE have not been established for the treatment of congenital CMV infection in infants and no similar disease occurs in adults; therefore, efficacy cannot

8.5 Geriatric Use

Studies of VALCYTE for oral solution or tablets have not been conducted in adults older than 65 years of age. Clinical studies of VALCYTE did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects. In general, dose selection for an elderly patient should be cautious, usually starting at the low end of the dosing range, reflecting the greater frequency of decreased hepatic, renal, or cardiac function, and of concomitant disease or other drug therapy. VALCYTE is known to be substantially excreted by the kidneys, and the risk of toxic reactions to this drug may be greater in patients with impaired renal function. Because renal clearance decreases with age, VALCYTE should be administered with consideration of their renal status. Renal function should be monitored and dosage adjustments should be made accordingly [see Dosage and Administration

Dose reduction is recommended when administering VALCYTE to patients with renal impairment [see Dosage and

Administration (2.5), Warnings and Precautions (5.2), Clinical Pharmacology (12.3)]. For adult patients on hemodialysis (CrCl less than 10 mL/min), VALCYTE tablets should not be used. Adult hemodialysis patients should use ganciclovir in accordance with the dose-reduction algorithm cited in the CYTOVENE®-IV complete product information section on DOSAGE AND ADMINISTRATION: Renal Impairment [see Dosage and Administration]

8.7 Hepatic Impairment The safety and efficacy of VALCYTE have not been studied in patients with hepatic impairment

be useful in reducing serum concentrations in patients who have received an overdose of VÁLCYTE [see Clinical Pharmacology (12.3)]. Adequate hydration should be maintained. The use of hematopoietic growth factors should be considered [see Warnings and Precautions (5.1) and Clinical Pharmacology (12.3)].

following adverse events: Hematological toxicity: myelosuppression including pancytopenia, bone marrow failure, leukopenia, neutropenia,

granulocytopenia

VALCYTE contains valganciclovir hydrochloride (valganciclovir HCl), a hydrochloride salt of the L-valyl ester of ganciclovir that exists as a mixture of two diastereomers. Ganciclovir is a synthetic guanine derivative active against

9-yl) methoxy]-3-hydroxypropyl ester, monohydrochloride. Valganciclovir HCl is a polar hydrophilic compound with a solubility of 70 mg/mL in water at 25°C at a pH of 7.0 and an n-octanol/water partition coefficient of 0.0095 at pH 7.0. The pKa for valganciclovir HCl is 7.6.

All doses in this insert are specified in terms of valgancicloving

12 CLINICAL PHARMACOLOGY 12.1 Mechanism of Action

Valganciclovir is an antiviral drug with activity against CMV [see Microbiology (12.4)].

transplant patients (Table 10).

Valganciclovir is a prodrug of ganciclovir. Valganciclovir $C_{\text{\tiny max}}$ and AUC are approximately 1% and 3% of those of harmacokinetics in Adults: The pharmacokinetics of ganciclovir after administration of valganciclovir tablets have

Table 10 Ganciclovir Pharmacokinetics* in Healthy Volunteers and HIV-positive/CMV-positive Adults Administered VALCYTE Tablets 900 mg Once Daily with Food

i k parameter	14	Value (IVICALI ± 3D)	
AUC _{0-24h} (mcg·h/mL)	57	29.1 ± 9.7	
C _{max} (mcg/mL)	58	5.61 ± 1.52	
Absolute oral bioavailability (%)	32	59.4 ± 6.1	
Elimination half-life (hr)	73	4.08 ± 0.76	
Renal clearance (mL/min/kg)	20	3.21 ± 0.75 (1 study, n=20)	
*Data were obtained from single and multiple dose studies in healthy volunteers, HIV-positive patients, and HIV-positive/CMV-positive patients with and without retinitis. Patients with CMV retinitis tended to have higher ganciclovir plasma concentrations than patients			

evaluated in HIV- and CMV-seropositive patients, patients with AIDS and CMV retinitis, and in solid organ

The systemic ganciclovir exposures attained following administration of 900 mg VALCYTE tablets once daily were comparable across kidney, heart and liver transplant recipients (Table 11).

Table 11 Ganciclovir Pharmacokinetics in Solid Organ Transplant Recipients Administered VALCYTE Tablets 900 mg Once Daily with Food

Parameter	Heart Transplant Recipients (N=17)	Liver Transplant Recipients (N=75)	Kidney Transplant Recipients* (N=68)
AUC _{0-24h} (mcg·h/mL)	40.2 ± 11.8	46.0 ± 16.1	48.2 ± 14.6
C _{max} (mcg/mL)	4.9 ± 1.1	5.4 ± 1.5	5.3 ± 1.5
Elimination half-life (hr)	6.58 ± 1.50	6.18 ± 1.42	6.77 ± 1.25

was administered once daily within 10 days of transplantation for a maximum of 200 days post-transplantation in pediatric kidney transplant patients. The results of these studies were supported by previous demonstration of efficacy in adult patients [see Adverse Reactions (6.1), Clinical Pharmacology (12.3), Clinical Studies (14.2)].

physiologically based pharmacokinetic (PBPK) model was developed based on the available pharmacokinetic data from pediatric and adult patients to support dosing in heart transplant patients less than 1 month of age. However, due to uncertainty in model predictions for neonates, VALCYTE is not indicated for prophylaxis in this age group.

e extrapolated from intravenous ganciclovir use in adults.

(2.5), Warnings and Precautions (5.2), Use in Specific Populations (8.6), Clinical Pharmacology (12.3)]. 8.6 Renal Impairment

(2.5) and Clinical Pharmacology (12.3)1.

Experience with VALCYTE Tablets: An overdose of VALCYTE could possibly result in increased renal toxicity [see Dosage and Administration (2.5), Use in Specific Populations (8.6)]. Because ganciclovir is dialyzable, dialysis may

Reports of adverse reactions after overdoses with valganciclovir, some with fatal outcomes, have been received from clinical trials and during postmarketing experience. The majority of patients experienced one or more of the

Hepatotoxicity: hepatitis, liver function disorder Renal toxicity: worsening of hematuria in a patient with pre-existing renal impairment, acute kidney injury, elevated

Gastrointestinal toxicity: abdominal pain, diarrhea, vomiting Neurotoxicity: generalized tremor, seizure

VALCYTE is available as a 450 mg tablet for oral administration. Each tablet contains 496.3 mg of valganciclovir HCl (corresponding to 450 mg of valganciclovir), and the inactive ingredients microcrystalline cellulose, povidone K-30, ridone and stearic acid. The film-coat applied to the tablets contains Opadry Pink®.

VALCYTE is also available as a powder for oral solution, which when constituted with water as directed contains 50 mg/mL valganciclovir free base. The inactive ingredients of VALCYTE for oral solution are sodium benzoate, fumaric acid, povidone K-30, sodium saccharin, mannitol and tutti-frutti flavoring.

Absorption, Distribution, Metabolism, and Excretion
The pharmacokinetic (PK) properties of VALCYTE are provided in Table 12.

Table 12	Pharmacokinetic Propert	ies of Ganciclovir and Valganciclovi	Associated with VALCYTE
		Volgonojolovin	Consistavia

	Valganciclovir	Ganciclovir		
Absorption		•		
T _{max} (h) median (min-max) (fed conditions)		2.18 1.7h to 3.0h		
Food effect (high fat meal/fasting): PK parameter ratio and 90% confidence interval ^a		C _{max} : 1.14 (0.95, 1.36) AUC: 1.30 (1.07, 1.51) ^a		
		T_{max} : \longleftrightarrow		
Distribution				
% Bound to human plasma proteins (ex vivo)	Unknown	1-2% over 0.5-51 mcg/mL		
Cerebrospinal fluid penetration	Unknown	Yes		
Metabolism		•		
	Hydrolyzed by intestinal and liver esterases	No significant metabolism		
Elimination				
Dose proportionality		AUC was dose proportional under fed conditions across a valganciclovi dose range of 450 to 2625 mg		
Major route of elimination	Metabolism to ganciclovir	Glomerular filtration and active tubular secretion		
t _{1/2} (h)		See Tables 10 and 11		
% Of dose excreted in urine	Ur	known		
% Of dose excreted in feces	Unknown			

ately 600 total calories (31.1 g fat, 51.6 g carbohydrates and 22.2 g protein) to 16 HIV-positive subject

Table 13 Pharmacokinetics of Ganciclovir from a Single Oral Dose of 900 mg VALCYTE Tablet

Renal Impairment: The pharmacokinetics of ganciclovir from a single oral dose of 900 mg VALCYTE tablets were evaluated in 24 otherwise healthy individuals with renal impairment. Decreased renal function results in decreased clearance of ganciclovir and increased terminal half-life (Table 13).

Estimated Creatinine Clearance* (mL/min)	N	Apparent Clearance (mL/min) Mean ± SD	AUC _{last} (mcg·h/mL) Mean ± SD	Half-life (hours) Mean ± SD
51-70	6	249 ± 99	49.5 ± 22.4	4.85 ± 1.4
21-50	6	136 ± 64	91.9 ± 43.9	10.2 ± 4.4
11-20	6	45 ± 11	223 ± 46	21.8 ± 5.2
≤ 10	6	12.8 ± 8	366 ± 66	67.5 ± 34

*Creatinine clearance calculated from 24-hour urine collection.

Hemodialysis reduces plasma concentrations of ganciclovir by about 50% following VALCYTE administration. Adult patients receiving hemodialysis (CrCl less than 10 mL/min) cannot use VALCYTE tablets because the daily dose of VALCYTE tablets required for these patients is less than 450 mg [see Dosage and Administration (2.5) and Use in

Pharmacokinetics in Pediatric Patients: The pharmacokinetics of ganciclovir were evaluated following the admi istration of valganciclovir in 63 pediatric solid organ transplant patients aged 4 months to 16 years, and in 16 pediatric heart transplant patients less than 4 months of age. In these studies, patients received oral doses of valganciclovi (either VALCYTE for oral solution or tablets) to produce exposure equivalent to an adult 900 mg dose [see Dosage and istration (2.3), Adverse Reactions (6.1), Use in Specific Populations (8.4), Clinical Studies (14.2)].

In studies using the pediatric valganciclovir dosing algorithm, the pharmacokinetics of ganciclovir were similar across organ types and age ranges (Table 14). Relative to adult transplant patients (Table 11), AUC values in pediatric

patients were somewhat increased, but were within the range considered safe and effective in adults. Table 14 Ganciclovir Pharmacokinetics by Age in Pediatric Solid Organ Transplant Patients

		Age Group						
Organ	PK Parameter mean (SD)	< 4 months	4 months to ≤ 2 years	> 2 to < 12 years	≥ 12 years			
	N	14ª	6	2	4			
Heart (N=26)	AUC _{0-24h} (mcg·h/mL)	66.3 (20.5)	55.4 (22.8)	59.6 (21.0)	60.6 (25.0)			
	C _{max} (mcg/mL)	10.8 (3.30)	8.2 (2.5)	12.5 (1.2)	9.5 (3.3)			
	t _{1/2} (h)	3.5 (0.87)	3.8 (1.7)	2.8 (0.9)	4.9 (0.8)			
	N		2	10	19			
Kidney (N=31)	AUC _{0-24h} (mcg·h/mL)	NA	67.6 (13.0)	55.9 (12.1)	47.8 (12.4)			
	C _{max} (mcg/mL)		10.4 (0.4)	8.7 (2.1)	7.7 (2.1)			
	t _{1/2} (h)		4.5 (1.5)	4.8 (1.0)	6.0 (1.3)			
	N		9	6	2			
Liver (N=17)	AUC _{0-24h} (mcg·h/mL)	NA	69.9 (37.0)	59.4 (8.1)	35.4 (2.8)			
	C _{max} (mcg/mL)		11.9 (3.7)	9.5 (2.3)	5.5 (1.1)			
	t _{1/2} (h)		2.8 (1.5)	3.8 (0.7)	4.4 (0.2)			

Ages ranged from 26 to 124 days

Pharmacokinetics in Geriatric Patients: The pharmacokinetic characteristics of VALCYTE in elderly patients have not

<u>Drug Interactions:</u> In vivo drug-drug interaction studies were not conducted with valganciclovir. However, because valganciclovir is rapidly and extensively converted to ganciclovir, interactions associated with ganciclovir will be expected for VALCYTE [see Drug Interactions (7)].

Table 15 and Table 16 provide a listing of established drug interaction studies with ganciclovir. Table 15 provides the effects of coadministered drug on ganciclovir plasma pharmacokinetic parameters, whereas Table 16 provides the effects of ganciclovir on plasma pharmacokinetic parameters of coadministered drug.

Table 15 Results of Drug Interaction Studies with Ganciclovir: Effects of Coadministered Drug on

Ganciclovir Pharmacokinetic Parameters						
Coadministered Drug	Ganciclovir Dosage	N	Ganciclovir Pharmacokinetic (PK) Parameter			
Mycophenolate mofetil (MMF) 1.5 g single dose	5 mg/kg IV single dose	12	No effect on ganciclovir PK parameters observed (patients with normal renal function)			
Trimethoprim 200 mg once daily	1000 mg every 8 hours	12	No effect on ganciclovir PK parameters observed			
Didanosine 200 mg every 12 hours simultaneously administered with ganciclovir	5 mg/kg IV twice daily	11	No effect on ganciclovir PK parameters observed			
	5 mg/kg IV once daily	11	No effect on ganciclovir PK parameters observed			
Probenecid 500 mg every 6 hours	1000 mg every 8 hours	10	AUC ↑ 53 ± 91% (range: -14% to 299%) Ganciclovir renal clearance ↓ 22 ± 20% (range: -54% to -4%)			

Coadministered Drug	Ganciclovir Dosage	N	Coadministered Drug Pharmacokinetic (PK) Parameter
Oral cyclosporine at therapeutic doses	5 mg/kg infused over 1 hour every 12 hours	93	In a retrospective analysis of liver allograft recipients, there was no evidence of an effect on cyclosporine whole blood concentrations.
Mycophenolate mofetil (MMF) 1.5 g single dose	5 mg/kg IV single dose	12	No PK interaction observed (patients with normal renal function)
Trimethoprim 200 mg once daily	1000 mg every 8 hours	12	No effect on trimethoprim PK parameters observed
Didanosine 200 mg every 12 hours	5 mg/kg IV twice daily	11	AUC ₀₋₁₂ ↑70 ± 40% (range: 3% to 121%) C _{max} ↑49 ± 48% (range: -28% to 125%)
Didanosine 200 mg every 12 hours	5 mg/kg IV once daily	11	AUC ₀₋₁₂ \uparrow 50 ± 26% (range: 22% to 110%) C _{max} \uparrow 36 ± 36% (range: -27% to 94%)

Mechanism of Action: Valganciclovir is an L-valyl ester (prodrug) of ganciclovir that exists as a mixture of two diastereomers. After oral administration, both diastereomers are rapidly converted to ganciclovir by intestinal and hepatic scharges. Ganciclovir is a synthetic analogue of 2'-deoxyguanosine, which inhibits replication of human CMV in cell culture and in vivo.

 $In CMV-infected \ cells, ganciclovir\ is\ initially\ phosphorylated\ to\ ganciclovir\ monophosphate\ by\ the\ viral\ protein\ kinase,$ pUL97. Further phosphorylation occurs by cellular kinases to produce ganciclovir triphosphate, which is then slowly metabolized intracellularly (half-life 18 hours). As the phosphorylation is largely dependent on the viral kinase, phosphorylation of ganciclovir occurs preferentially in virus-infected cells. The virustatic activity of ganciclovir is due to inhibition of the viral DNA polymerase, pUL54 by ganciclovir triphosphate.

Antiviral Activity: The quantitative relationship between the cell culture susceptibility of human herpes viruses to intivirals and clinical response to antiviral therapy has not been established, and virus sensitivity testing has not been attivities and emindred response to attiviting the standardized. Sensitivity test results, expressed as the concentration of drug required to inhibit the growth of virus in cell culture by 50% (EC_{so}), vary greatly depending upon a number of factors including the assay used. Thus, the eported EC, values of ganciclovir that inhibit human CMV replication in cell culture (laboratory and clinical isolates have ranged from 0.08 to 22.94 μ M (0.02 to 5.75 mcg/mL). The distribution and range in susceptibility observed in one assay evaluating 130 clinical isolates was 0 to 1 μ M (35%), 1.1 to 2 μ M (20%), 2.1 to 3 μ M (27%), 3.1 to 4 μ M (13%), 4.1 to 5 μ M (5%), less than 5 μ M (less than 1%). Ganciclovir inhibits mammalian cell proliferation (CC_{s0}) in cell culture at higher concentrations ranging from 40 to greater than 1,000 μ M (10.21 to greater than 250 mcg/mL). Bone marrow-derived colony-forming cells are more sensitive [CC_{so} value = 2.7 to 12 μM (0.69 to 3.06 mcg/mL)].

Viral Resistance:
Cell culture: CMV isolates with reduced susceptibility to ganciclovir have been selected in cell culture. Growth of CMV strains in the presence of ganciclovir resulted in the selection of amino acid substitutions in the viral protein kinase pUL97 (M460IV, L595S, G598D, and K599T) and the viral DNA polymerase pUL54 (D301N, N410K, F412V, P488R,

L516R, C539R, L545S, F595I, V812L, P829S, L862F, D879G, and V946L). In vivo: Viruses resistant to ganciclovir can arise after prolonged treatment or prophylaxis with valganciclovir by selection of substitutions in pUL97 and/or pUL54. Limited clinical data are available on the development of clinical resistance to ganciclovir and many pathways to resistance likely exist. In clinical isolates, seven canonical pUL97 substitutions, (M460V/I, H520Q, C592G, A594V, L595S, and C603W) are the most frequently reported ganciclovir resistance-associated substitutions. These and other substitutions less frequently reported in the literature, or

Table 17 Summary of Resistance-associated Amino Acid Substitutions Observed in the CMV of Patients Failing Ganciclovir Treatment or Prophylaxis

F342Y, K359E/Q, L405P, A440V, M460I/V/T/L, V466G/M, C518Y, H520Q, P521L, del 590-593, A591D/V, C592F/G, A594E/G/T/V/P, L595F/S/T/W, del 595, del 595-603, E596D/G/Y, K599E/M, del 600-601, del 597-600, del 601-603, C603W/R/S/Y, C607F/S/Y, I610T, A613V

E315D, N408D/K/S, F412C/L/S, D413A/E/N, L501F/I, T503I, K513E/N/R, D515E, L516W, I521 P522A/L/S, V526L, C539G, L545S/W, Q578H/L, D588E/N, G629S, S695T, I726T/V, E756K, L773V, V781I, V787E/L, L802M, A809V, T813S, T821I, A834P, G841A/S, D879G, A972V, del 981-982, A987G

The presence of known ganciclovir resistance-associated amino acid substitutions was evaluated in a study that extended valganciclovir CMV prophylaxis from 100 days to 200 days post-transplant in adult kidney transplant patients at high risk for CMV disease (D+/R-) [see Clinical Studies (14.1)]. Five subjects from the 100 day group and four subjects from the 200 day group meeting the resistance analysis criteria had known ganciclovir resistance-associated amino acid substitutions detected. In six subjects, the following resistance-associated amino acid substitutions were detected within pUL97: 100 day group: A440V, M460V, C592G; 200 day group: M460V, C603W.

In three subjects, the following resistance-associated amino acid substitutions were detected within pUL54: 100 day group: E315D; 200 day group: E315D, P522S. Overall, the detection of known ganciclovir resistance-associated amino acid substitutions was observed more frequently in patients during prophylaxis therapy than after the ompletion of prophylaxis therapy (during therapy: 5/12 [42%] versus after therapy: 4/58 [7%]). The possibility of Il resistance should be considered in patients who show poor clinical response or experience persistent viral excretion during therapy. Cross-Resistance: Cross-resistance has been reported for amino acid substitutions selected in cell culture by

ganciclovir, cidofovir or foscarnet. In general, amino acid substitutions in pUL54 conferring cross-resistance to ganciclovir and cidofovir are located within the exonuclease domains and region V of the viral DNA polymerase. Whereas, amino acid substitutions conferring cross-resistance to foscarnet are diverse, but concentrate at and between regions II (codon 696–742) and III (codon 805–845). The amino acid substitutions that resulted in reduced susceptibility to ganciclovir and either cidofovir and/or foscarnet are summarized in Table 18.

Substitutions at amino acid positions pUL97 340-400 have been found to confer resistance to ganciclovir. Resistance data based on assays that do not include this region should be interpreted cautiously.

Table 18 Summary of pUL54 Amino Acid Substitutions with Cross-Resistance between Ganciclovir,

oluolovii, alia/ol i oscarilet					
Cross-resistant to cidofovir	D301N, N408D/K, N410K, F412C/L/S/V, D413E/N, P488R, L5011, T503I, K513E/N, L516R/W, I521T, P522S/A, V526L, C539G/R, L545S/W, Q578H, D588N, I726T/V, E756K, L733V, V787E, V812L, T813S, A834P, G841A, del 981-982, A987G				
Cross-resistant to foscarnet	F412C, Q578H/L, D588N, V715A/M, E756K, L733V, V776M, V781I, V787E/L, L802M, A809V, V812L, T813S, T821I, A834P, G841A/S, del 981-982				
	Cross-resistant to cidofovir				

13 NONCLINICAL TOXICOLOGY

observed in clinical trials, are listed in Table 17.

ote: Many additional pathways to ganciclovir resistance likely exist

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Long-term carcinogenicity studies have not been conducted with VALCYTE. However, upon oral administration alganciclovir is rapidly and extensively converted to ganciclovir. Therefore, like ganciclovir, valganciclovir is a

Ganciclovir was carcinogenic in the mouse at oral doses that produced exposures approximately 0.1x and 1.4x, respectively, the mean drug exposure in humans following the recommended intravenous dose of 5 mg/kg, based on area under the plasma concentration curve (AUC) comparisons. At the higher dose, there was a significant increase in the incidence of tumors of the preputial gland in males, forestomach (nonglandular mucosa) in males and females and reproductive tissues (ovaries, uterus, mammary gland, clitoral gland and vagina) and liver in females. At the lower dose, a slightly increased incidence of tumors was noted in the preputial and harderian glands in males, forestomach in males and females, and liver in females. Ganciclovir should be considered a potential carcinogen in humans.

Valganciclovir increases mutations in mouse lymphoma cells. In the mouse micronucleus assay, valganciclovir was clastogenic. Valganciclovir was not mutagenic in the Ames Salmonella assay. Ganciclovir increased mutations in mouse lymphoma cells and DNA damage in human lymphocytes in vitro. In the mouse micronucleus assay, ganciclovir was clastogenic. Ganciclovir was not mutagenic in the Ames Salmonella assay.

alganciclovir is converted to ganciclovir and therefore is expected to have similar reproductive toxicity effects as ganciclovir [see Warnings and Precautions (5.3)]. Ganciclovir caused decreased mating behavior, decreased fertility, and an increased incidence of embryolethality in female mice following intravenous doses that produced an exposure approximately 1.7x the mean drug exposure in humans following the dose of 5 mg per kg, based on AUC comparisons. Ganciclovir caused decreased fertility in male mice and hypospermatogenesis in mice and dogs following daily oral or intravenous administration. Systemic drug exposure (AUC) at the lowest dose showing toxicity n each species ranged from 0.03 to 0.1x the AUC of the recommended human intravenous dose. Valganciclovir caused similar effects on spermatogenesis in mice, rats, and dogs. These effects were reversible at lower doses but irreversible at higher doses. It is considered likely that ganciclovir (and valganciclovir) could cause temporary or permanent inhibition of human spermatogenesis.

14 CLINICAL STUDIES

Induction Therapy of CMV Retinitis: In one randomized open-label controlled study, 160 patients with AIDS and newly diagnosed CMV retinitis were randomized to receive treatment with either VALCYTE tablets (900 mg twice daily for 21 days, then 900 mg once daily for 7 days) or with intravenous ganciclovir solution (5 mg per kg twice daily for 21 days, then 5 mg per kg once daily for 7 days). Study participants were: male (91%), White (53%), Hispanic (31%), and Black (11%). The median age was 39 years, the median baseline HIV-1 RNA was 4.9 log₁₀, and the median CD4 cell count was 23 cells/mm. A determination of GMV retinitis progression by the masked review of retinal photographs taken at baseline and Week 4 was the primary outcome measurement of the 3-week induction therapy. Table 19 provides the outcomes at 4 weeks.

Table 19 Week 4 Masked Review of Retinal Photographs in CMV Retinitis Study

	Intravenous Ganciclovir	VALCYTE Tablets
Determination of CMV retinitis progression at Week 4	N=80	N=80
Progressor Non-progressor	7 63	7 64
Death Discontinuations due to Adverse Events Failed to return	2 1 1	1 2 1
CMV not confirmed at baseline or no interpretable baseline photos	6	5

Maintenance Therapy of CMV Retinitis: No comparative clinical data are available on the efficacy of VALCYTE tablets for the maintenance therapy of CMV retinitis because all patients in the CMV retinitis study received open-label VALCYTE tablets after Week 4. However, the AUC for ganciclovir is similar following administration of 900 mg VALCYTE tablets once daily and 5 mg per kg intravenous ganciclovir once daily. Although the ganciclovir C_{\max} is lower following VALCYTE tablets administration compared to intravenous ganciclovir, it is higher than the C_{\max} obtained following oral ganciclovir administration. Therefore, use of VALCYTE tablets as maintenance therapy is supported by a plasma concentration-time profile similar to that of two approved products for maintenance therapy of CMV retinitis.

Prevention of CMV Disease in Heart, Kidney, Kidney-Pancreas, or Liver Transplantation: A double blind, double-dummy active comparator study was conducted in 372 heart, liver, kidney, or kidney-pancreas transplant patients at high risk for CMV disease (D+/R-). Patients were randomized (2 VALCYTE: 1 oral ganciclovir) to receive either VALCYTE tablets (900 mg once daily) or oral ganciclovir (1000 mg three times a day) starting within 10 days of transplantation until Day 100 post-transplant. The proportion of patients who developed CMV disease, including CMV syndrome and/or tissue-invasive disease during the first 6 months post-transplant was similar between the VALCYTE tablets arm (12.1%, N=239) and the oral ganciclovir arm (15.2%, N=125). However, in liver transplant patients, the incidence of tissue-invasive CMV disease was significantly higher in the VALCYTE group compared with the ganciclovir group. These results are summarized in Table 20.

Mortality at six months was 3.7% (9/244) in the VALCYTE group and 1.6% (2/126) in the oral ganciclovir group.

Table 20 Percentage of Patients with CMV Disease, Tissue-Invasive CMV Disease or CMV Syndrome by Organ Type: Endpoint Committee, 6 Month ITT Population

	CMV D	isease¹		Invasive isease	CMV Syndrome ²		
rgan	VGCV	GCV	VGCV	GCV	VGCV	GCV	
	(N=239)	(N=125)	(N=239)	(N=125)	(N=239)	(N=125)	
ver	19%	12%	14%	3%	5%	8%	
=177)	(22/118)	(7/59)	(16/118)	(2/59)	(6/118)	(5/59)	
dney	6%	23%	1%	5%	5%	18%	
=120)	(5/81)	(9/39)	(1/81)	(2/39)	(4/81)	(7/39)	
eart	6%	10%	0%	5%	6%	5%	
=56)	(2/35)	(2/21)	(0/35)	(1/21)	(2/35)	(1/21)	
dney/Pancreas	0%	17%	0%	17%	0%	0%	
=11)	(0/5)	(1/6)	(0/5)	(1/6)	(0/5)	(0/6)	
/ = oral ganciclovir VGCV = valganciclovir							

umber of patients with CMV disease = Number of patients with tissue-invasive CMV disease or CMV syndrom CMV syndrome was defined as evidence of CMV viremia accompanied with fever greater than or equal to 38°C on two or more occasions separated by at least 24 hours within a 7-day period and one or more of the following: malaise, leukopenia, atypical lymphocytosis, thrombocytopenia, and elevation of hepatic transaminases

revention of CMV Disease in Kidney Transplantation: A double-blind, placebo-controlled study was conducted in 326 kidney transplant patients at high risk for CMV disease (D+/R-) to assess the efficacy and safety of extending VALCYTE CMV prophylaxis from 100 to 200 days post-transplant. Patients were randomized (1:1) to receive VALCYTE tablets (900 mg once daily) within 10 days of transplantation either until Day 200 post-transplant or until Day 100 post-transplant followed by 100 days of placebo. Extending CMV prophylaxis with VALCYTE until Day 200 post-transplant demonstrated superiority in preventing CMV disease within the first 12 months post-transplant in high risk kidney transplant patients compared to the 100 day dosing regimen (primary endpoint). These results are summarized in Table 21

Table 21 Percentage of Kidney Transplant Patients with CMV Disease, Tissue-Invasive CMV Disease or CMV Syndrome, 12 Month ITT Population

	CMV D	CMV Disease ¹		Invasive Jisease	CMV Syndrome ²		
	100 Days	200 Days	100 Days	200 Days	100 Days	200 Days	
	VGCV	VGCV	VGCV	VGCV	VGCV	VGCV	
	(N=163)	(N=155)	(N=163)	(N=155)	(N=163)	(N=155)	
Cases	36.8%	16.8%	1.8%	0.6%	35.0%	16.1%	
	(60/163)	(26/155)	(3/163) ³	(1/155)	(57/163)	(25/155)	
VGCV = valganciclovir.							

Visure of variganizations.

Number of patients with CMV disease = Number of patients with tissue-invasive CMV disease or CMV syndrome

CMV syndrome was defined as evidence of CMV viremia accompanied with at least one of the following; fever (greater than or equal to 38°C), severe malaise, leukopenia, atypical lymphocytosis, thrombocytopenia, and elevation of hepatic transaminases

Two patients in the 100 day group had both tissue-invasive CMV disease and CMV syndrome; however, these patients are counted as having only tissue-invasive CMV disease.

The percentage of kidney transplant patients with CMV disease at 24 months post-transplant was 38.7% (63/163) for the 100 day dosing regimen and 21.3% (33/155) for the 200 day dosing regimen.

<u>Prevention of CMV in Pediatric Heart, Kidney, or Liver Transplantation:</u> Sixty-three children, 4 months to 16 years of age, who had a solid organ transplant (kidney 33, liver 17, heart 12, and kidney/liver 1) and were at risk for developing CMV disease, were enrolled in an open-label, safety, and pharmacokinetic study of oral VALCYTE (VALCYTE for oral solution or tablets). Patients received VALCYTE once daily within 10 days after transplant until a maximum of 100 days post-transplant. The daily doses of VALCYTE were calculated at each study visit based on body surface area and a modified creatinine clearance [see Dosage and Administration (2.3)].

The pharmacokinetics of ganciclovir were similar across organ transplant types and age ranges. The mean daily anciclovir exposures in pediatric patients were somewhat increased relative to those observed in adult solid organ ansplant patients receiving VALCYTE 900 mg once daily, but were within the range considered safe and effective in adults [see Clinical Pharmacology (12.3)]. No case of CMV syndrome or tissue-invasive CMV disease was reported within the first six months post-transplantation.

Prevention of CMV in Pediatric Kidney Transplantation: Fifty-seven children, 1 to 16 years of age, who had a renal transplant and were at risk for developing CMV disease, were enrolled in an open-label tolerability study of oral VALCYTE (VALCYTE for oral solution or tablets). Patients received VALCYTE once daily within 10 days after transplant until a maximum of 200 days post-transplant. The daily doses of VALCYTE were calculated at each study visit based on body surface area and a modified creatinine clearance [see Dosage and Administration (2.3)]. No case of CMV In body during a first and a mounted disease was reported within the first 12 months post-transplantat

15 REFERENCES

- 1. Brion LP, Fleischman AR, McCarton C, Schwartz GJ. A simple estimate of glomerular filtration rate in low birth weight infants during the first year of life: noninvasive assessment of body composition and growth. J of Ped
- 2. NIOSH [2014]. NIOSH list of antineoplastic and other hazardous drugs in healthcare settings. By Connor TH, MacKenzie BA, DeBord DG, Trout DB, O'Callaghan JP, Cincinnati, OH: U.S. Department of Health and Human Services, Centers for Disease Control and Prevention, National Institute for Occupational Safety and Health, DHHS (NIOSH) Publication No. 2014-138 (Supersedes 2012-150).

16 HOW SUPPLIED/STORAGE AND HANDLING

VALCYTE tablets: Supplied as 450 mg, pink, convex oval tablets with "VGC" on one side and "450" on the other t contains 450 mg valganciclovir. VALCYTE is supplied in bottles of 60 tablets (NDC 0004-0038-22). Store VALCYTE tablets at 20°C to 25°C (68°F to 77°F); excursions are permitted to 15°C to 30°C (59°F to 86°F)

VALCYTE for oral solution; Supplied as a white to slightly yellow powder blend for constitution, forming a colorless to brownish-yellow tutti-frutti flavored solution. Available in glass bottles containing approximately 100 mL of solution after constitution. Each bottle can deliver up to a total of 88 mL of solution. Each bottle is supplied with a bottle adapter and 2 oral dispensers (NDC 0004-0039-09).

Prior to dispensing to the patient, VALCYTE for oral solution must be prepared by the pharmacist [see Dosage and

Store dry powder at 20°C to 25°C (68°F to 77°F); excursions are permitted to 15°C to 30°C (59°F to 86°F) [see USP Store constituted solution under refrigeration at 2°C to 8°C (36°F to 46°F) for no longer than 49 days. Do not freeze.

17 PATIENT COUNSELING INFORMATION

Advise the patient to read the FDA-approved patient labeling (Patient Information and Instructions for Use)

Serious Adverse Reactions Inform patients that VALCYTE may cause granulocytopenia (neutropenia), anemia, thrombocytopenia and elevated creatinine levels and that dose modification or discontinuation of dosing may be required. Complete blood counts, platelet counts, and creatinine levels should be monitored frequently during treatment [see Warnings and Precautions

Pregnancy and Contraception Inform females of reproductive potential that VALCYTE causes birth defects in animals. Advise them to use effective

contraception during and for at least 30 days following treatment with VALCYTE. Similarly, advise males to use condoms during and for at least 90 days following treatment with VALCYTE [see Use in Specific Populations (8.1,

Advise patients that VALCYTE is considered a potential carcinogen [see Nonclinical Toxicity (13.1)].

Advise mothers not to breast-feed if they are receiving VALCYTE because of the potential for hematologic toxicity and cancer in nursing infants, and because HIV can be passed to the baby in breast milk [see Use in Specific Populations

Advise patients that VALCYTE may cause temporary or permanent female and male infertility [see Warnings and Precautions (5.3), Use in Specific Populations (8.3)].

Inform patients that tasks requiring alertness may be affected including the patient's ability to drive and operate machinery as seizures, dizziness, and/or confusion have been reported with the use of VALCYTE [see Adverse]

Use in Patients with CMV Retinitis Inform patients that VALCYTE is not a cure for CMV retinitis, and they may continue to experience progression of

retinitis during or following treatment. Advise patients to have ophthalmologic follow-up examinations at a minimum of every 4 to 6 weeks while being treated with VALCYTE. Some patients will require more frequent follow-up.

form adult patients that they should use VALCYTE tablets, not VALCYTE for oral solution [see Dosage and

Inform patients to take VALCYTE with food to maximize bioavailability.

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PATIENT INFORMATION

VALCYTE (Val-site) for oral solution

What is the most important information I should know about VALCYTE?

VALCYTE can cause serious side effects, including

VALCYTE (Val-site)

- Blood and bone marrow problems. VALCYTE can affect the bone marrow lowering the amount
 of your white blood cells, red blood cells, and platelets and may cause serious and life-threatening
- Kidney failure. Kidney failure may happen in people who are elderly, people who take VALCYTE with certain other medicines, or people who are not adequately hydrated.
- Fertility problems. VALCYTE may lower sperm count in males and cause fertility problems. VALCYTE may also cause fertility problems in women. Talk to your healthcare provider if this is a
- Birth defects. VALCYTE causes birth defects in animals. It is not known if VALCYTE causes birth defects in people. If you are a female who can become pregnant, you should use effective birth itrol during treatment with VALCYTE and for at least 30 days after treatment. If you are pregnant, talk to your healthcare provider before starting treatment with VALCYTE. If you are a femal who can become pregnant, you should have a pregnancy test done before starting VALCYTE
- Tell your healthcare provider right away if you become pregnant during treatment with VALCYT Males should use condoms during treatment with VALCYTE, and for at least 90 days after treatment, if their female sexual partner can become pregnant. Talk to your healthcare provider if you have questions about birth control
- Cancer. VALCYTE causes cancer in animals and may potentially cause cancer in people.

Your healthcare provider will do regular blood tests during treatment with VALCYTE to check you for side effects. Your healthcare provider may change your dose or stop treatment with ALCYTE if you have serious side effects.

What is VALCYTE?

VALCYTE is a prescription antiviral medicine.

In adults, VALCYTE tablets are used: • to treat cytomegalovirus (CMV) retinitis in people who have acquired immunodeficiency syndrome

- (AIDS). When CMV virus infects the eyes, it is called CMV retinitis. If CMV retinitis is not treated, it can cause blindness
- to prevent CMV disease in people who have received a kidney, heart, or kidney-pancreas transplant and who have a high risk for getting CMV disease.

VALCYTE does not cure CMV retinitis. You may still get retinitis or worsening of retinitis during or after treatment with VALCYTE. It is important to stay under a healthcare provider's care and have your eyes checked at least every 4 to 6 weeks during treatment with VALCYTE.

In children, VALCYTE tablets or oral solution are used:

• to prevent CMV disease in children 4 months to 16 years of age who have received a kidney transplant and have a high risk for getting CMV disease.

• to prevent CMV disease in children 1 month to 16 years of age who have received a heart transplant

and have a high risk for getting CMV disease. It is not known if VALCTYE is safe and effective in children for prevention of CMV disease in liver transplant, in kidney transplant in infants less than 4 months of age, in heart transplant in infants less than 1 month of age, in children with AIDS who have CMV retinitis, and in infants with congenital CMV

Do not take VALCYTE if you have had a serious allergic reaction to valganciclovir, ganciclovir or any of the ingredients of VALCYTE. See the end of this leaflet for a list of the ingredients in VALCYTE. Before you take VALCYTE, tell your healthcare provider about all of your medical conditions,

including if you: have low blood cell counts

- have kidney problems are receiving hemodialysis
- are receiving radiation treatment
- are pregnant or plan to become pregnant. See "What is the most important information I should know about VALCYTE?"
- are breastfeeding or plan to breastfeed. It is not known if VALCYTE passes into your breast milk. You
- should not breastfeed if you take VALCYTE.

 o You should not breastfeed if you have Human Immunodeficiency Virus (HIV-1) because of the risk of passing HIV-1 to your baby.
- Talk to your healthcare provider about the best way to feed your baby.

Tell your healthcare provider about all the medicines you take, including prescription and overthe-counter medicines, vitamins and herbal supplements, VALCYTE and other medicines may affect each other and cause serious side effects. Keep a list of your medicines to show your healthcare provider and pharmacist.

You can ask your healthcare provider or pharmacist for a list of medicines that interact with VALCYTE. • Do not start taking a new medicine without telling your healthcare provider. Your healthcare provider can tell you if it is safe to take VALCYTE with other medicines

How should I take VAI CYTE? Take VALCYTE exactly as your healthcare provider tells you. Your dose of VALCYTE will depend on

- Adults should only take VALCYTE tablets. Children may take either VALCYTE tablets or oral solution. Take VALCYTE with food.
- Do not break or crush VALCYTE tablets. Avoid contact with your skin or eyes. If you come in contact with the contents of the tablet or oral solution, wash your skin well with soap and water or rinse vour eves well with plain water.
- If your child is prescribed VALCYTE for oral solution, your pharmacist will give you oral dosing dispensers to measure your child's dose of VALCYTE for oral solution. To be sure you receive the prescribed dose, it is important to use the dispenser provided to you. See the detailed Instructions or Use below for information about how to take VALCYTE for oral solution. Ask your pharmacist if you have any questions. If you lose or damage your oral dispensers and cannot use them, contact vour pharmacist.
- If you take too much VALCYTE, call your healthcare provider or go to the nearest hospital emergency room right away.

What should I avoid during treatment with VALCYTE? VALCYTE can cause seizures, dizziness, and confusion. You should not drive a car or operate machinery until you know how VALCYTE affects you

What are the possible side effects of VALCYTE VALCYTE may cause serious side effects, includin See "What is the most important information I should know about VALCYTE?"

The most common side effects of VALCYTE in adults include: · low white cell, red cell and platelet cell counts in blood tests

diarrhea fever headache fatique sleeplessness · urinary tract infection

- shaky movements (tremors) vomiting The most common side effects of VALCYTE in children include diarrhea vomiting
- low white blood cell counts in blood tests upper respiratory tract infection urinary tract infection

These are not all the possible side effects of VALCYTE.

Call your doctor for medical advice about side effects. You may report side effects to FDA

How should I store VALCYTE?

- Store VALCYTE tablets at room temperature between 68°F to 77°F (20°C to 25°C). • Store VALCYTE for oral solution in the refrigerator between 36°F to 46°F (2°C to 8°C), for no longer
- Do not freeze Do not keep VALCYTE that is out of date or that you no longer need.

Keep VALCYTE and all medicines out of the reach of children. General information about the safe and effective use of VALCYTE.

Medicines are sometimes prescribed for purposes other than those listed in a Patient Information leaflet. Do not use VALCYTE for a condition for which it was not prescribed. Do not give VALCYTE to other people, even if they have the same symptoms you have. It may harm them, You can ask your pharmacist or healthcare provider for information about VALCYTE that is written for health

What are the ingredients in VALCYTE?

Active ingredient: valganciclovir hydrochloride Inactive ingredients for tablets: microcrystalline cellulose, povidone K-30, crospovidone, and stearic acid. The film-coating applied to the tablets contains Opadry Pink®

Inactive ingredients for oral solution: sodium benzoate, fumaric acid, povidone K-30, sodium saccharin, mannitol and tutti-frutti flavoring.

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nstructions for Use **VALCYTE (Val-site)** for oral solutio

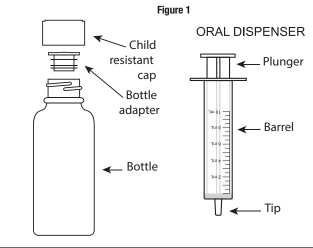
Be sure that you read, and that you understand and follow these instructions carefully to ensure proper dosing of the oral solution.

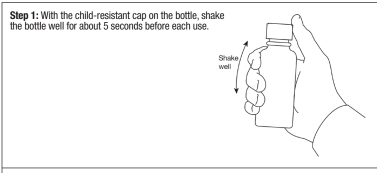
- Avoid contact with your skin or eyes. If you come in contact with the contents of the oral solution. wash your skin well with soap and water or rinse your eyes well with plain water.
- Always use the oral dispenser provided to give or take a dose of VALCYTE for oral solution. • Call your pharmacist if your oral dispenser is lost or damaged, and they will tell you how to continue to give or take a dose of VALCYTE for oral solution.

Do not use VALCYTE for oral solution after the discard date on the bottle.

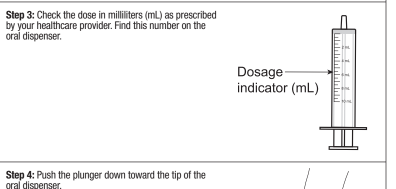
Ask your healthcare provider or pharmacist to show you how to measure your prescribed dose.

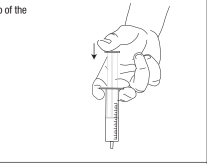
To take a dose of VALCYTE for oral solution, you will need the bottle of medicine and an oral dispenser provided with the medicine (see Figure 1). Your pharmacist inserts the bottle adapter in the VALCYTE for oral solution bottle.













Step 6: Carefully turn the bottle upside down with the oral dispenser in place. Pull the plunger to withdraw the prescribed dose If you see air bubbles in the oral dispenser, fully push in the plunger so that the oral solution Then withdraw your prescribed dose of VALCYTE

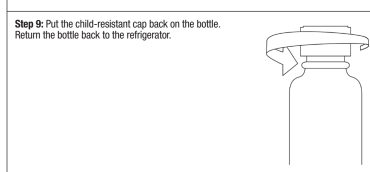
Step 7: I eave the oral dispenser in the bottle adapter and turn the bottle to an upright position. Slowly remove the oral dispenser from the bottle adapter.

Step 8: Give or take the dose of VAI CYTE for oral solution. · Place the tip of the oral dispenser

flows back into the bottle.

for oral solution.

 Slowly push down the oral dispenser plunger until the oral dispenser is empty.



Step 10: Rinse the oral dispenser with tap water after each use.

 Remove the plunger from the oral dispenser barrel by pulling the plunger all the way out of the barrel. Rinse the oral dispenser barrel and plunger with water and let them air dry.

When the oral dispenser barrel and plunger

are dry, put the plunger back into the oral

dispenser barrel for the next use.

Do not throw away the oral dispenser

How should I store VALCYTE for oral solution? • Store solution in the refrigerator at 36°F to 46°F (2°C to 8°C) for no longer than 49 days.

This Patient Information and Instructions for Use have been approved by the U.S. Food and Drug

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