

Specialty Distributors

Veloxis has worked with the specialty distribution teams from major wholesalers to ensure availability and ease of ordering ENVARSUS XR.

	Customer Service Number	Customer Service Hours	Emergency Orders Number
ASD Specialty Healthcare* <small>A division of AmerisourceBergen</small>	1(800) 746-6273	Monday - Thursday 7:00am - 6:30pm CST Friday 7:00am - 6:00pm CST	1(800) 746-6273
McKesson Plasma and Biologics	1(877) 625-2566	Monday - Friday 8:00am - 6:30pm CST	1(877) 625-2566
Cardinal Specialty Pharmaceutical Distribution <small>A division of Cardinal Health</small>	1(866) 476-1340	Monday 8:00am - 5:00pm CST Tuesday - Friday 8:00am - 6:00pm CST	1(866) 476-1340 <small>After hours, select "Emergency Order" option</small>
Bioridge	1(973) 564-8004 bob.anderson@basembia.com	Monday - Friday 8:30am - 5:00pm EST	1(973) 564-8004
Morris and Dickson	1(800) 388-3833	Monday - Thursday 8:00am - 6:00pm CST	N/A

* You can reach ASD Specialty Healthcare account set-up at 1 (877) 654-7808, Monday-Thursday 8:30am - 6:30pm CST and Friday 8:30am - 6:00pm CST

INDICATIONS AND USAGE

ENVARSUS XR is indicated for the prophylaxis of organ rejection in de novo kidney transplant patients in combination with other immunosuppressants.

ENVARSUS XR is also indicated for the prophylaxis of organ rejection in kidney transplant patients converted from tacrolimus immediate-release formulations in combination with other immunosuppressants.

IMPORTANT SAFETY INFORMATION




WARNING: MALIGNANCIES AND SERIOUS INFECTIONS

Increased risk for developing serious infections and malignancies with ENVARSUS XR or other immunosuppressants that may lead to hospitalization or death

Please see additional Important Safety Information on pages 3-4 and, full Prescribing Information, including Boxed Warning.

Available Strengths, Package Sizes and NDCs

ENVARUSUS XR is available in a variety of tablet strengths and package sizes to meet the needs of patients and customers.

Strength						
	0.75 mg Tablet		1 mg Tablet		4 mg Tablet	
Tablet per Bottle	30	100	30	100	30	100
NDC #	68992-3075-3	68992-3075-1	68992-3010-3	68992-3010-1	68992-3040-3	68992-3040-1
ASD Specialty Healthcare						
	45526	45525	45524	45523	45522	45521
Cardinal Specialty Pharmaceutical Distribution						
	5151956	5151998	5152020	5152046	5152046	5169974
McKesson Plasma Biologics						
	3490026	3490018	3489994	3489986	3489978	3489945

Note: Some wholesaler systems may request an 11-digit NDC number. If that is the case, please add a zero “0” just before the last 1 or 3 in the 10-digit NDC number.

\$ Co-Pay Support* and Patient Assistance Program

Veloxis is committed to ensuring access to ENVARUSUS XR for all patients, regardless of their situation. \$0 Co-Pay support is available for eligible commercially insured patients*.

For uninsured patients or patients without coverage for ENVARUSUS XR, our patient assistance program offers ENVARUSUS XR at no cost.

To access these programs, please contact Veloxis Transplant Support
1-844 VELOXIS (835-6947)

* Please contact Veloxis Transplant Support for full program details.

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CONTRAINDICATIONS

ENVARUSUS XR is contraindicated in patients with known hypersensitivity to tacrolimus.

WARNINGS AND PRECAUTIONS

Lymphoma and Other Malignancies: Immunosuppressants, including ENVARUSUS XR, increase the risk of developing lymphomas and other malignancies, particularly of the skin. Post-transplant lymphoproliferative disorder (PTLD), associated with Epstein-Barr Virus (EBV), has been reported in immunosuppressed organ transplant patients.

Serious Infections: Immunosuppressants, including ENVARUSUS XR, increase the risk of developing bacterial, viral, fungal, and protozoal infections, including opportunistic infections. These infections may lead to serious, including fatal, outcomes.

Not Interchangeable with Other Tacrolimus Products - Medication Errors: Medication errors, including substitution and dispensing errors, between tacrolimus capsules and tacrolimus extended-release capsules were reported outside the U.S. This led to serious adverse reactions, including graft rejection, or other adverse reactions due to under- or over-exposure to tacrolimus. ENVARUSUS XR is not interchangeable or substitutable with tacrolimus extended-release capsules, tacrolimus capsules or tacrolimus for oral suspension.

New Onset Diabetes after Transplant: ENVARUSUS XR caused new onset diabetes after transplant (NODAT) in kidney transplant patients, which may be reversible in some patients. African-American and Hispanic kidney transplant patients are at an increased risk.

Nephrotoxicity: ENVARUSUS XR, like other calcineurin-inhibitors, can cause acute or chronic nephrotoxicity. Consider dosage reduction in patients with elevated serum creatinine and tacrolimus whole blood trough concentrations greater than the recommended range. The risk for nephrotoxicity may increase when ENVARUSUS XR is concomitantly administered with CYP3A inhibitors (by increasing tacrolimus whole blood concentrations) or drugs associated with nephrotoxicity.

Neurotoxicity: ENVARUSUS XR may cause a spectrum of neurotoxicities. The most severe neurotoxicities include posterior reversible encephalopathy syndrome (PRES), delirium, seizure, and coma; others include tremors, paresthesias, headache, mental status changes, and changes in motor and sensory functions.

Hyperkalemia: Mild to severe hyperkalemia, which may require treatment, has been reported with tacrolimus including ENVARUSUS XR. Concomitant use of agents associated with hyperkalemia may increase the risk for hyperkalemia.

Hypertension: Hypertension is a common adverse reaction of ENVARUSUS XR therapy and may require antihypertensive therapy.

Risk of Rejection with Strong CYP3A Inducers and Risk of Serious Adverse Reactions with Strong CYP3A Inhibitors:

The concomitant use of strong CYP3A inducers may increase the metabolism of tacrolimus, leading to lower whole blood trough concentrations and greater risk of rejection. In contrast, the concomitant use of strong CYP3A inhibitors may decrease the metabolism of tacrolimus, leading to higher whole blood trough concentrations and greater risk of serious adverse reactions. Therefore, adjust ENVARUSUS XR dose and monitor tacrolimus whole blood trough concentrations when coadministering ENVARUSUS XR with strong CYP3A inhibitors or strong CYP3A inducers.

WARNINGS AND PRECAUTIONS (continued)

QT Prolongation: ENVARSUS XR may prolong the QT/QTc interval and cause Torsade de Pointes. Avoid ENVARSUS XR in patients with congenital long QT syndrome. Consider obtaining electrocardiograms and monitoring electrolytes periodically during treatment in patients with congestive heart failure, bradyarrhythmias, those taking certain antiarrhythmic medications or other products that lead to QT prolongation, and those with electrolyte disturbances. When coadministering ENVARSUS XR with other substrates and/or inhibitors of CYP3A, a reduction in ENVARSUS XR dosage, monitoring of tacrolimus whole blood concentrations, and monitoring for QT prolongation is recommended.

Immunizations: Whenever possible, administer the complete complement of vaccines before transplantation and treatment with ENVARSUS XR. Avoid the use of live attenuated vaccines during treatment with ENVARSUS XR. Inactivated vaccines noted to be safe for administration after transplantation may not be sufficiently immunogenic during treatment with ENVARSUS XR.

Pure Red Cell Aplasia: Cases of pure red cell aplasia (PRCA) have been reported in patients treated with tacrolimus. If PRCA is diagnosed, consider discontinuation of ENVARSUS XR.

ADVERSE REACTIONS

De Novo kidney transplant patients: Most common adverse reactions (incidence $\geq 15\%$) reported with ENVARSUS XR are diarrhea, anemia, urinary tract infection, hypertension, tremor, constipation, diabetes mellitus, peripheral edema, hyperkalemia and headache.

Conversion of kidney transplant patients from immediate-release tacrolimus: Most common adverse reactions (incidence $\geq 10\%$) reported with ENVARSUS XR include: diarrhea and blood creatinine increased.

USE IN SPECIFIC POPULATIONS

Pregnancy: Based on postmarketing surveillance, registry and animal data may cause fetal harm. Advise pregnant women of the potential risk to the fetus.

Nursing Mothers: Tacrolimus is present in human milk. Discontinue drug or nursing, taking into account the importance of drug to the mother.

Females and Males of Reproductive Potential: Advise female and male patients of reproductive potential to speak with their healthcare provider on family planning options including appropriate contraception prior to starting treatment with ENVARSUS XR. Based on animal studies, ENVARSUS XR may affect fertility in males and females.

Pediatric Use: The safety and efficacy of ENVARSUS XR in pediatric patients have not been established.

Geriatric Use: Clinical studies of ENVARSUS XR did not include sufficient numbers of patients aged 65 and over to determine whether they respond differently from younger patients.

Renal Impairment: Frequent monitoring of renal function is recommended. Lower doses may be required.

Hepatic Impairment: Frequent monitoring of tacrolimus trough concentrations is recommended. With greater tacrolimus whole blood trough concentrations in patients with severe hepatic impairment, there is a greater risk of adverse reactions and dosage reduction is recommended.

Race: African-American patients may require higher doses to attain comparable trough concentrations compared to Caucasian patients. African-American and Hispanic kidney transplant patients are at an increased risk for new onset diabetes after transplant. Monitor blood glucose concentrations and treat appropriately.

To report SUSPECTED ADVERSE REACTIONS, contact Veloxis Pharmaceuticals, Inc. at 1-844-VELOXIS (835-6947) or FDA at 1-800-FDA-1088 or visit www.fda.gov/medwatch.