

# **Standard Pharmaceutical Product and Medical Device Information (Rx Product Only)**

Version 2021						Introduction Ty	pe: New I	tem		x Final Version			Date:	6/26/	6/2024
			PRODUCT INFORMA	TION						SPECIAL HAN	DLING AND STO	RAGE REQUI	REMENTS*		
Company Name: Camber Pharmaceuticals, Inc. Application: ANDA						a. Temperature – Indicate the USP temperature range for this product.									
Application Number for NDA/AN			ce):	21	2414				un romporata.	Temperature Range	Controlled Room		and 25 C (68	s° – 77° F)	
Medical Device Class, if applical			,.										`		
DUNS:	11-856-3719								1	Other Temperature Range	Requirement	Excursions r	permitted to 1	5°C to 30°C (	(59° – 86° F)
Proprietary Name (If Applicable) a		ame: Lenal	idomide Capsules 2.5 mg						T	(write in)					
Selling Unit NDC:	31722-257-28		Unit of Use NDC		31722-257-28	UPC: 3	331722257282		1	Notes					
UDI			CVX Code:			MVX Code:									
Description:	Lenalidomide Cap	oulos 2 E ma							i	Is this product to be shippe	to quotomoro on	202		No	1
Description.	Lenandonnide Cap	osules 2.5 mg								Is this product to be shipped				No	-
Active Ingredient(s):		Lenalidomide							1	is this product to be shipped	a to customers on	ary loc:		140	1
/ tours mg. salom(s).		Lonandonnao							b. Contact for	temperature excursion qu	estions:				
URL for Additional Product Inform	mation:	www.camberpharm	na.com						1	Name:		Soma Raju			
Address:	800 Centennial Av	ve, Suite 1				Address 2:				Number:		732-529-042	23		
City:	Piscataway				State:	NJ	Zip: 08854			Group E-mail:		somaraju@h	neterousa.cor	n	
Key Contact:	Customer Service	)			Email:	customerservice@d	camberpharma.com								
Phone Number:	1-866-827-3647				Fax:	732-562-8788			c. Special reg	ulations for product in any	states?			No	1
Product Therapeutic Classification	on:	Imide immunomodulator	y thalidomide analogue (cereblon m	odulator)						Special returns requirement	s for this product?			*Yes	
					_										
	ADDITI	ONAL PRODUCT IN	IFORMATION			PRODUCT DE	ESCRIPTION INFORM	MATION	d. Store produ	uct (unit of sale) upright?				No	1
The product is?			Is the Product	Direct And D	rop-Ship				] ]	Protect product (unit of sa	le) from liaht?			No	i
a legend device?		No	Is the Product	Unit of Use			28 ct		e. Shelf life:	r rotoot product (anii or ot	,g			24	Months
if yes, enter class #		110	Orphan Drug Status			Size:	20 01			Initial shelf life at launch (	if different):				Months
a product kit?		No	o.p.ia D. ag otatao				2.5 mg			indui onon mo at laanon (					1
if yes, list NDCs of		1.15	FDA Approval Status			Strength:	- 3				ORDER INFORI	MATION			
component parts						B F	Hard gelatin ca	apsule							
reverse numbered?		No				Dosage Form:				Unit of Sale		What is the	NDC selling	unit?	
co-licensed?		No	Allergens Present							x Bottle		1 Bottle of 2	8 Capsules		
latex-free?		Yes	Daime	nd Lactose		Product Shape	Capsule			Box/Carton		(Write-in, e.	g. 1 Box of 1	0 Vials)	
preservative-free?		Yes	Dairy a	id Lactose		Product Snape	e: '			Ampule					
correctional institution block?		No				Product Color	Pink opaque ca	ap and		Glass		Minimum or	der quantity	?	Yes
opioid?		No				Froduct Color	white opaque b	oody		Tube					
Cannabinoid?		No	Country of Origin	India		Product Imprir	Imprinted with '			Vial Liquid Sgl					
If Unit Dose, is item bar coded to u	unit dose for					r roudet imprii	and 'L1' on bod	dy		Vial Liquid Multi				ch package	type?
hospital scanning?			Is this product covered							Vial Powder Sgl		1	Each		
If Unit Dose, indicate NDC here:			Trade Agreements Act	TAA)?	No					Vial Powder Multi			Inner/Cartor	/Pack	
									]	Other: Write In			Case		
			FOR GENERIC DRUG PF	RODUCTS											
					Aut		'If Authorized Generic		PHARMACY ORDER / BILL UNIT						
I. Orange Book Rating:	AB						section fields are not a	applicable	Rec. sell unit to customer? Rx billing unit to pharm			acy:			
II. Generic Equivalent to What Bra	and?:	Revlimid										Each			
									(Write-in, e.g.	1 Vial)			Gram		
		DRUG SUPP	LY CHAIN SECURITY ACT	(DSCSA) INFO	RMATION								Milliliter		
Does supplier meet DSCSA defini	ition of monufacture	ror2	Yes	_	GLN:	0331722498975				ITEN	I AND PACKING I	NEOPMATIO	NI .		
Is product exempt from DSCSA?		err	No	-	GLN.	0331722490973				116	I AND I ACKING I	NI OKWATIOI	•		
									1		Dim	/IIC u	-4- \		
If yes, select exemption:					GCP:					Weight Lbs.		ions (US msn	•	Volume	Saleable #
Other exemption - Write in: Is product repackaged?			No		If was	ginal product purch	anna d		Item/Each:	-	Depth	Width	Height	(Cube)	Pieces
Is product repackaged?	e aveluciva dietribu	ttor?	Yes	-	direct from mi		laseu		item/Each:	0.08	1.53	1.53	2.38	5.57	1
Has FDA granted waiver/exception			No	$\dashv$			repackaged product		Box/Carton/B	undle/					
If yes, attach documentation from		ouuci:	140		r rovide sourc	e manuracturer for	repackageu product		Inner Pack:	under					
you, attaon accumentation no									Case:						
		GT	IN AND HIBCC PRODUCT	INFORMATION						2.4	9.84	6.5	4.13	264.15	24
									Pallet:						
Saleable Unit of Measure	S	Saleable Quantity	HIBCC		GTIN	<b>I-14</b>	Unit of Use G	TIN-14							
X Item/Each	_	1				1722257282	00331722257								
Box/Carton/Bundle/Inner Pack										COST INFORMATION			WHOLESAL	ER USE ONL	Y:
X Case		24			2033	1722257286									
Pallet									Regular Cost			Vendor #:			
									Invoice Cost (	(WAC) (\$)	\$20,157.36	Whsl. Code	#:		
									11						
									1 1			Fineline Co	de:		
									As of date:	5/12/2023		Fineline Co	de:		
									As of date:	5/12/2023		Fineline Co	de:		
												Fineline Co	de:		
			Attach copy of SAFETY D	ATA SHEET (SI	OS) or non hazar	d letter, PACKAGE I	NSERT, LABEL AND	PHOTO OF F				Fineline Co	de:		



## **Standard Pharmaceutical Product and Medical Device Information (Rx Product Only)**

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For Designated Drop Ship Only Products, Please Use Page 3

MATERIAL HA	ZARD CLASSIFICATION and TRANSPORTATION						
Is this product (check all that apply):							
a. Cytotoxic? Yes	SDS Hazard Classification						
b. CA Prop. 65 Carcinogen or Reproductive Toxicant?							
Is the product a CA Prop 65 carcinogen?	x Organic Corrosive						
Is the product a CA Prop 65 reproductive toxicant?	Inorganic Oxidizer						
Does the product label bear a CA Prop 65 warning?	Steroid/Androgen Contact Hazard						
2000 the product about a control of training.	- Constanting						
c. Contact Hazard?	Does the product have an Aerosol class? If yes, No						
d. Does this product require special clean-up instructions?	identify NFPA Storage Level:						
(If yes, attach SDS with special instructions.)	NFPA Storage Level:						
e. Does the product contain DEHP?							
Is this product regulated for shipment by DOT?	Is the product a NIOSH hazardous drug?						
(if yes, answer a-e below and provide SDS)	If yes, indicate which:  Group 2 items (non-antineoplastic that meets a hazard criterion)						
a. UN/Identification Number	il yes, indicate which.						
b. Proper Shipping Name	l l						
c. DOT Hazard Class	Hazardous Waste Identification						
d. Packing Group	THE STATE OF THE S						
e. Inhalation Hazard?	EPA Hazardous Waste Code: WT02, MN01, R006 Waste Characteristics D004						
Is this product regulated for shipment by IATA?  No	REMS or REGISTRY RESTRICTIONS						
(if yes, answer a-e below and provide SDS) a. UN/Identification Number	REMO OF REGISTRY RESTRICTIONS						
b. Proper Shipping Name	Is there a REMS on this product?						
c. DOT Hazard Class	If Yes, is it managed with a pharmacy registry?						
d. Packing Group	in res, is intantaged with a pharmacy registry?  Website URL:  www.lenalidomiderems.com						
e. Inhalation Hazard?	website offic.						
Is the product restricted for air shipment? If so, indicate restriction:	Med Guide Required Yes						
Passenger	Limited Distribution Requirement  Comments / Details: (For example, iPledge program?)  Must be a certified Lenalidomide REMS Program Location						
Cargo	Comments / Details: (For example, iPledge program?)  Must be a certified Lenalidomide REMS Program Location						
Passenger & Cargo							
Is this a reportable quantity? No	REMS: Yes						
RQ Threshold:	REMS Program Manager Name: Bristol Myers Squibb Phone: 1-888-423-5436						
Is this a marine pollutant? No	Supplier Manages REMS registry exclusively: No						
Is this product shipped utilizing an authorized DOT exception or Special Permit?	Wholesale distributor support: No						
No (if yes, identify method below)	Provider Name: DEA #:						
Limited Quantity	Site Enrollment Number assigned by Supplier:  NPI #:						
Consumer Commodity, ORM-D	by Supplier: NPI #:						
Small Quantity (49 CFR 173.4) Special Permit; DOT-SP	Comments						
Special Provision (listed in Column 7 of 49 CFR 172.101);	Continents						
SP#	Registry: Yes						
5r#							
ADD'L STORAGE INFORMATION	Registry Program Contact Name: REMS Call Center Phone: 1-888-423-5436						
	Comments Lenalidomide REMS is a shared REMS program						
Is the Product							
Controlled Substance? No Controlled Substance Code	RETURN INSTRUCTIONS						
Controlled by State(s)? No Listed Chemical (List I or II) No							
ARCOS Reportable? No If yes, indicate which:	Contact tel. # if product received damaged: 1-888-423-5436						
Schedule No. Is it a scheduled listed chemical product?: No	Is product returnable for credit: No						
CLASS OF TRADE RESTRICTION:	URL/Link to returns policy:						
No restriction: Select YES if sold to retail pharmacy, hospitals, clinics and physician offices  Yes	contact - www.lenalidomiderems.com						
Restricted to retail pharmacy only:  No	Special regulations or returns requirements for this						
Restricted to hospital, clinics, and physician offices only:  Restricted from US territories? (explain in comments)  No	product in certain states?  If so, which states? Other requirements? Comments?						
	Dispensed Product Returns: Return to prescriber, pharmacy or call 1-888-423-5436 to return directly to Lenalidomide REMS program. (All States)						
Comments: Distribution to validated Lenalidomide REMS Certified Dispensing Locations only.	Dispensed Product Returns: Return to prescriber, pnarmacy or call 1-888-423-9436 to return directly to Lenalidomide KEIMS program. (All States)  Non-Dispensed Product Returns: Return directly to Cambler's third party return goods processor. (All States)						
	Damaced in Transit Returns (by carrier): Return to Camber Distribution Center. (All States)						
MISCELLANE	OUS NOTES and/or Image of Product Barcode:						



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#### FOR DESIGNATED DROP SHIP PRODUCT ONLY - if not a designated drop ship, do not complete.

Order Method for De	esignated Drop Ship Product	Standard Order Receipt and Processing					
b. Autofax c. Fax d. Phone only e. Supplier Web Site only  Minimum Order Quantity: 1 Bottle  Supplier's Customer Service Number: 732-1  Contracted 3PL company / contact #: Nam Phor		Purchase order daily receipt cut off time by supplier Cut off time:  11:00 AM Monday - Thursday					
Expedited freight fees billed with each order:	No	Overnight receipt available: No					
Drop Ship service fee billed with each order:	No	PO Receipt cut off time:					
Drop Ship miscellaneous fees billed:  Comments:	No	Days of week overnight is available:  Monday Tuesday Wednesday Thursday Friday  Priority Overnight receipt available:					
Class of	Trade Restriction:	PO Receipt Cut off time:					
No restriction: Select YES if sold to retail pharma Restricted to retail pharmacy only: Restricted to hospital, clinics, and physician office Restricted from US territories? (explain in comme	acy, hospitals, clinics and physician offices  No es only:  No	Saturday Overnight receipt available:  PO Receipt Cut off time:  Phone: Fax: EDI:  Overnight Fees apply: Other fees apply:					
Other Data Informat	tion Required to Process PO:	Return Instructions					
Patient Procedure Date: Physician Name: Physician/Clinic Phone # Physician State License # Physician/Clinic DEA #: Physician/Clinic Specialty:  Miscel	e	Contact # if product is received damaged:  Is product returnable for credit:  URL/Link to returns policy:  https://www.camberpharma.com/partner-resources/#returned-goods-policy  Special regulations or returns requirements for this product in certain states?  Yes  If so, which states? Other requirements? Comments?  Dispensed Product Returns: Return to prescriber, pharmacy or call 1-888-423-5436 to return directly to Lenalidomide REMS program. (All States)	x467				
		Non-Dispensed Product Returns: Return directly to Camber's third party return goods processor. (All States)  Damaged in Transit Returns (by carrier): Return to Camber Distribution Center. (All States)					
		ADDITIONAL INFORMATION					
		Is product order for scheduled patient procedure?  Is product order for restocking purposes?  Yes					