Addiction Therapy Services Product Catalog



Addiction Therapy Services Product Catalog

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AddictionTherapyServices.com





8 mg of naloxone HCI in each nasal spray device



NDC: 59467-679-01

Packaging:

1 Carton containing two ready-to-use nasal spray devices

To learn more, contact your Hikma Account Manager or email us at usaddictiontherapy.com

Kloxxado® is a registered trademark of Hikma Pharmaceuticals USA Inc.



BUPRENORPHINE Sublingual Tablets C-III

PRODUCT DESCRIPTION

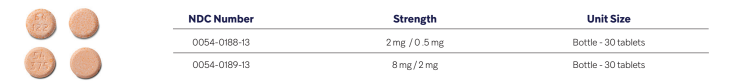
Round/White Sublingual Tablets

54	NDC Number	Strength	Unit Size	
	0054-0176-13	2 mg	Bottle - 30 tablets	
	0054-0177-13	8 mg	Bottle - 30 tablets	

BUPRENORPHINE AND NALOXONE Sublingual Tablets USP C-III

PRODUCT DESCRIPTION

Round/Speckled Peach to Peach Sublingual Tablets



BUPRENORPHINE AND NALOXONE Sublingual Film C-III

PRODUCT DESCRIPTION Orange/Rectangular Film	NDC Number	Strength	Unit Size
	43598-579-30	2 mg / 0 .5 mg	30 Pouches each containing 1 Sublingual Film
Buprenorphine and Nakozone θ Sublingual Film 8 mg/2 mg δενιή δενιή δενιή	43598-580-30	4 mg/1 mg	30 Pouches each containing 1 Sublingual Film
Bit can at the final and a data. The source is a data are seen to an address of the source of the so	43598-582-30	8 mg / 2 mg	30 Pouches each containing 1 Sublingual Film
	43598-581-30	12 mg / 3 mg	30 Pouches each containing 1 Sublingual Film

BUPRENORPHINE AND NALOXONE Sublingual Film C-III is manufactured by Lohmann Therapy Systems, (LTS), West Caldwell, NJ 07006 for Dr. Reddy's Laboratories Inc., Princeton, NJ 08540, USA. Please visit https://dailymed.nlm.nih.gov/ for additional information specific to this product, including the Full Prescribing Information with complete indications for use, warnings, precautions and adverse reactions.

DISKETS[®] Dispersible Tablets C-II (Methadone Hydrochloride Tablets for Oral Suspension USP)

PRODUCT DESCRIPTION

Light Pinkish-Orange Cross-Scored Tablets. Orange-Pineapple Flavored

54	NDC Number	Strength	Unit Size
	0054-4538-25	40 mg	Bottle - 100 tablets

METHADONE HCI Tablets USP C-II

PRODUCT DESCRIPTION

Round/White Tablets

NDC Number	Strength	Unit Size
0054-4570-25	5 mg	Bottle - 100 tablets
0054-4571-25	10 mg	Bottle - 100 tablets

METHADONE HCI Oral Concentrate USP C-II

PRODUCT DESCRIPTION	NDC Number	Strength	Unit Size
Clear/Colorless Solution - Flavorless	0054-0391-68	10 mg / mL	1000 mL Bottle
Clear/Red Colored Solution - Cherry flavored	0054-0392-68	10 mg / mL	1000 mL Bottle

Please visit www.hikma.com/us for additional product information, including the Full Prescribing Information with complete indications for use, warnings, precautions and adverse reactions for each product.



Ordering Instructions

Mandatory items for order processing:

The following licenses must be on file with Hikma:

- Current DEA Registration Certificate
- State License
- Completed Due Diligence Questionnaire

C-II Ordering:

- Follow directions printed on the back of the DEA-222 Form.
- Shipment can only be made to the address shown on the form.
- If your name and address have changed, contact your regional DEA office for a new supply of forms.
- 222 Form MUST be mailed. 222 Forms that are emailed or faxed will NOT be accepted.

222 Forms must be mailed to the following address: Hikma Pharmaceuticals USA Inc. 1809 Wilson Rd., 08-118 Columbus, OH 43228

• Federal Regulations do not allow us to accept forms that have corrections, alterations or write overs.

C-III Ordering:

- Follow directions printed on the ATS Product Order Form.
- The clinic name, address, and DEA number MUST match the information printed on DEA Certificate of Registration.
- Order form can be emailed or faxed.

Shipping:

- Our distribution center ships orders Monday through Friday.
- Next business day delivery by 10:30 to most U.S. addresses via FedEx Priority Overnight.

C-II Ordering Instructions

Sample Single-Sheet 222 Form with instructions:

PURCHASER INFORMATION REGISTRATION INFORMATION REGISTRATION #: REGISTERED AS: SCHEDULES: ORDER FORM NUMBER: DATE ISSUED:		me	20	SUPPLIER DEA NUMBER:# R W 0 4 9 8 4 1 9 PART 2: TO BE FILLED IN BY PURCHASER <u>Hikma Pharmaceuticals USA Inc.</u> BUSINESS NAME <u>1809 Wilson Road, 08-18</u> STREET ADDRESS <u>Columbus, OH 43228</u> CITY, STATE, ZIP CODE														
Van int or Sigr	ne and Tit Type Name and and indi	tle Title cate Sigi	PURCHASER	TO FILLEI PURCI	ET 5: BE D IN BY HASER	(nar AL ⁻ Sig	ne in p TERN natur	art 2) i ATE I e- by 1	if order DEA f	r is end # upplier	orsed to	anoth	ier su			be filled in by	irst supplie	
EM	NO. OF PACKAGES	PACKAGE SIZE	NAME OF ITEM	NUMBER REC'D	DATE REC'D	PAR	T 4: 1	О ВЕ			BY S					NUMBER DATE SHIPPED SHIPPED		
1		1000mL	Methadone HCI Oral Conc- Cherry			-			INA	HUNA		3000	9			SHIFFED	SHIFFE	
2		1000mL	Methadone HCI Oral Conc- Clear															
3	50	100	Methadone HCI 40mg Diskets											-				
4	20	100	Methadone HCI Tablets, 5mg				5						7					
5	10	100	Methadone HCI Tablets, 10mg											7			-	
6																		
7												T						
8													\square					
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Purchaser Information:

- Print/Type name and title.
- Sign and indicate signature authority.
 For example, "attorney-in-fact", "power of attorney", "designated agent", or "secretary" may be used.
- Fill in current date.
- Customer must complete the following columns:
 - No. of packages
 - Package size
 - Name of item(s): Please distinguish between Cherry and Clear Methadone HCI Oral Conc.
 Fill in "Last Line Completed"

2 Supplier Identification

 Enter DEA number, name, and address of supplier. DEA#: RW0498419
 Hikma Pharmaceuticals USA Inc.
 1809 Wilson Road, 08-118
 Columbus, OH 43228

3 Alternate Supplier Identification

- Leave this portion of the form blank.
- **4** Controlled Substance Shipment
 - Leave this portion of the form blank.

6 Controlled Substance Receipt

• Purchaser will enter the number of packages received and date received for each line item on its copy of the original order form.

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Purchaser must make a copy of the original order form for its records before mailing the original to the supplier.

Mail the original copy to the address below. Hikma Pharmaceuticals USA Inc. 1809 Wilson Road, 08-118 Columbus, OH 43228

Controlled Substance Ordering System

Benefits of CSOS

- Improves accuracy by eliminating the paper 222 form
- Quicker turn-around times
- Lower costs per transaction
- Introduces greater security and helps ensure DEA compliance

Overview

- Customer applies for a digital signing certificate issued by the DEA
- Upon approval by the DEA, customer downloads the digital certificate
- Contact Hikma to gain access to the csos.hikma.com website
- Customer can begin ordering their products electronically

How to apply for a digital certificate?

To obtain a digital certificate, a DEA registrant must enroll in the CSOS program through the DEA website (deaecom.gov). Every individual who wants to sign electronic orders for controlled substances must enroll with the DEA to acquire his or her own personal digital certificate.

Choose your applicant type:

- Registrant Individual who is authorized to sign DEA applications. The Registrant applicant must name him/herself as Coordinator or delegate the role to another Principle Coordinator or existing subscriber.
- Coordinator Any individual employed by the DEA Registrants organization. The Principle Coordinator
 is a required role for each DEA registration number in the CSOS program and will be the primary
 CSOS contact. If the Registrant is requesting to be the Coordinator, he or she should apply only
 as a registrant and indicate him or herself as the Coordinator.
- Power of Attorney Individual with the authority to sign controlled substance orders for a DEA Registrant.

What happens after I enroll?

Once the enrollment process is complete, within about six weeks the DEA will send an activation notice via email and postal mail containing your access code and password. These two items (access code and password) must be married up in order for you to retrieve your digital certificate from the DEA website.

When all documentation has been received, contact Hikma Addiction Therapy Services (Toll Free 833.449.3475) to gain access to the csos.hikma.com website. Hikma will assist you in setting up your account and provide training for use of the system.

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Return Goods Policy Effective April 1, 2020

Hikma Pharmaceuticals USA Inc. ("Hikma") Return Goods Policy (this "Policy") applies to the return and/or credit of product(s) purchased by a direct customer of Hikma. This Policy also applies on returns from indirect customers that are returning through the wholesaler from the original purchase. Unless otherwise required by regulation or law or expressly agreed by the parties, the following policy applies to Hikma product(s).

Product Returns

 Upon receipt of a Return Authorization or box label(s), actual returns are to be forwarded to the processing facility at the following location: Inmar RX Solutions, Inc.
 3845 Grand Lakes Way, Suite 125 Grand Prairie, Texas 75050

Return Authorization ("RA") Procedures for Expired Products

- Request for an RA (box labels) may be made by any of the below methods through Hikma's third party reverse logistics processor, Inmar RX Solutions, Inc. ("Inmar"):
 - Accessing Inmar's website at <u>https://returns.</u> <u>healthcare.inmar.com</u> (a PDF copy of your debit memo will need to be uploaded); or
 - E-mail the debit memo to <u>rarequest@inmar.com.</u> Include: NDC#; lot#; and the expiration date(s) assigned to each item; or
 - Fax your debit memo to Inmar at 817-868-5343.
 - PLEASE NOTE: All third-party return processors must contact Inmar for a RA.

Returnable Product - For Reimbursement

- Authorized expired product, which shall be defined as: Product returned within six (6)months prior to the expiry date, or within twelve (12) months thereafter, in full and unopened containers with a Hikma label, purchased directly from Hikma and returned directly to Inmar.
- Recalled product, as stated on a recall notice issued by Hikma, which is returned directly to Inmar after requesting and receiving a RA from Inmar.
- A DEA Form 222 is required in order to return C-II controlled substances. Please send DEA Form 222 requests to: Fax # (817) 868-5342 or E-mail: 222@inmar.com.
- Products meeting the "Conditions for Credit" and not included on the "Non-Returnable Product" listing as set forth below.

Non-Returnable Product - No Credit

• Partial units, except where mandated by federal, state or local laws.

- An inner pack included within a saleable package with a different NDC.
- Private Labelled, Repackaged, Reconstituted, and/or Contract Manufactured product.
- Product(s) sold at no cost including, but not limited to donations and samples.
- Product(s) with more than six (6) months expiration dating or product greater than twelve (12) month from the expiration date assigned to such product.
- Product(s) not purchased directly from Hikma or the customer's authorized distributor/wholesaler.
- Product(s) sold as short-date, close-out, special promotion, and/or sold as non-returnable.
- Customer overstocked product, unless prior approval from Hikma is received.
- Product(s) damaged due to negligence or insurable causes, such as fire, floods, and/or natural disasters.
- Product(s) purchased through a bankruptcy sale.
- Product(s) damaged/deteriorated due to negligence, including, but not limited to improper handling or storage by the customer.
- Product(s) received by Inmar thirty (30) days or more after the date assigned on the RA.
- Product(s) purchased or distributed contrary to federal, state or local laws.
- Product(s) sold to any city/municipality, county, state and/or federal entity for the purpose of stockpiling directly by Hikma or through an authorized wholesaler or distributor of record.
- Product(s) with defaced or missing Hikma labels which do not clearly display the product's expiration date, NDC and lot number including, but not limited to products with a prescription label.
- Product(s) purchased outside of the United States and its territories, commonwealths and possessions, including, but not limited to the District of Columbia and the commonwealth of Puerto Rico.
- Product(s) purchased for future events including speculative purposes.
- Expired returns with a returnable value of \$25.00 or less in value.

Return Goods Policy Effective April 1, 2020

Shipping Errors

- Hikma must be notified of any shipping disputes within three (3) business days of receipt of product(s). Product(s) shipped in error by Hikma must be returned within thirty (30) business days of shipment to receive credit. Product(s) returned after thirty (30) business days of shipment shall be considered excess stock and will not be eligible for credit.
- If the error involves products which are controlled substances, Hikma must be notified within 24 hours of receipt of the order of any overages or mistakes in such controlled substances order.
- For clarity, customers will limit approved damage returns to packages/cases that are damaged and unsaleable to qualify for credit from Hikma.

Conditions for Credit

- Product eligible for return which is received and verified by Inmar (or destroyed by customer's agent with prior written approval by Hikma) within thirty (30) days of receipt of a RA with a valid RA number are eligible for credit.
- Excluding Non-Returnable Products as defined above.
- Customer agrees to a one (1%) percent current year return limitation based on customer's prior calendar year's purchase value of all return eligible products. Return value based on return credit dollars issued for both direct and indirect/third party customer returns.
- See below for how returnable product will be valued by Hikma, transportation and Hikma disclaimers.

Valuation of Returns and Credit Memos

- For direct customers, a credit will be issued based upon the lower of the current net invoice price at the time the returned product(s) is received by Inmar -OR- the lowest net invoice price paid in the prevailing 24 months.
- For indirect customers, a credit will be issued based upon the lower of the current net indirect price at the time the returned merchandise is received by Inmar -OR- the lowest net indirect contract price paid ("Lowest Indirect Price") in the prevailing 24 months from the wholesaler. If Hikma cannot identify the Lowest Indirect Price for a customer, then Hikma will use a predetermined indirect return price.
- Indirect returns will be credited through the wholesaler or distributor of purchase.
- The amount of credit issued or authorized by Hikma is directly correlated to what is validated by Inmar. In the

event of any conflict between the customer's claimed quantity and the quantity validated by Inmar, the quantity validated by Inmar shall control.

- For recalled product, current net sale price will be credited to ensure replacement costs are covered.
- Credit will be issued by Hikma in the form of a credit memo only.

Third Party Processors

- Third party processors must comply with all requirements of this Policy. Hikma will not pay or reimburse any service fees to the purchaser or third-party return processor (e.g., handling, processing, or freight charges incurred, etc.).
- Hikma will not process returns using pricing from the third party's internally generated price list.

Transportation

- Transportation charges, including prepaid freight and insurance, are the responsibility of the customer except when due to a Hikma error, as solely determined by Hikma.
- Hikma is not responsible for lost or damaged shipments of returned product(s). Insuring and tracking shipments are the responsibility of the customer.

Company Disclaimers

- Submission of the returned product does not constitute Hikma's acceptance for credit.
- Sales representatives are not permitted to authorize and/or pick-up returned products.
- Package size, lot number and lot expiration date will be obtained and verified after receipt of product by Inmar.
- Hikma reserves the right to refuse credit when product is returned through parties other than Inmar.
- All returns are subject to review by Hikma, and issuance of a RA number does not guarantee credit.
- Hikma reserves the sole right to determine whether items qualify under this Policy for return, credit or refund.
- Inmar's determination of the physical count of the returned products will be final. By returning products you authorize Hikma and its designee, as your agent, to destroy, without payment or other recourse, any returned product.
- Any and all credits provided pursuant to this Policy are only valid if redeemed within one (1) year of issuance. Any and all credits that are not redeemed within one (1) year of issuance shall be null and void.

Return Goods Policy Effective April 1, 2020

- Unauthorized deductions for returned product(s) will not be accepted.
- Hikma reserves the right to require proof of purchase source on all merchandise returned for credit or refund.
- Non-Hikma product(s) returned with Hikma product(s) will not be the responsibility of Hikma. Hikma reserves the right to charge customers for any costs incurred to process and destroy such non-Hikma product. Any such non-Hikma product will not be returned to the customer.

This Return Goods Policy supersedes all previous policies and may be modified by Hikma, from time to time, in its discretion. Hikma values the relationship it shares with its customers and will make a commercially reasonable attempt to provide thirty (30) days advance notification of policy changes. Customers will be expected to adhere to the most current policy which can be found on the Hikma website.



Corporate Headquarters

Hikma Pharmaceuticals USA Inc. 200 Connell Drive, Suite 4000 Berkeley Heights, NJ 07922 Tel: 732.542.1191 | Fax: 732.542.0940 hikma.com/us



Addiction Therapy Services Customer Service

Business Hours: 8am ET – 5pm ET, Monday – Friday Tel: 1.833.449.3475 | Fax: 866.464.1562 Email: usaddictiontherapy@hikma.com

Please visit www.hikma.com/us for additional product information, including the Full Prescribing Information with complete indications for use, warnings, precautions and adverse reactions for each product.

Product images may not reflect actual sizes and/or exact colors.

All other trademarks listed herein are the property of their respective owners and are used for illustrative purposes only. These trademark owners are not associated or affiliated with Hikma Pharmaceuticals USA, Inc.





Addiction Therapy Services Customer Service

Toll Free 1.833.449.3475 Fax: 1.866.464.1562 Email: usaddictiontherapy@hikma.com addictiontherapyservices.com

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