



Outsource Smarter

Keep Your Pharmacy at the Ready

Critical medications from a partner you can trust.

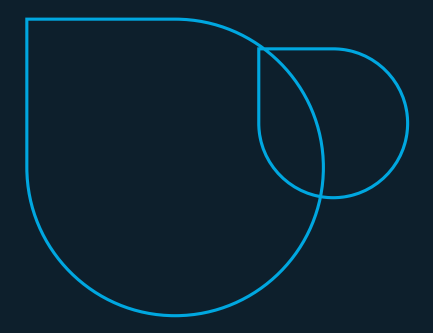
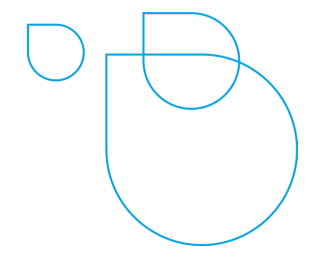


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Product Catalog

Why SCA Pharma?



As a reliable source of high-quality sterile admixtures and pre-filled syringes, we're proud to be your partner in delivering life-saving drugs to those who need them most.

Feel free to reach out should you have any questions about the products listed in this catalog or require assistance in placing your order. We're here to help you make smarter outsourcing decisions, including improved demand planning and optimizing inventory tracking and control.

Working together, we will make sure you have the medications you need, when you need them.







We understand that the most effective 503B solution is a partner you can trust. SCA Pharma collaborates with your management team to implement strategies that lower labor costs, reduce drug waste, and keep your pharmacy at the ready while freeing up manpower and resources to focus energy on other things.

A discerning commitment to quality ensures every product in this catalog meets or exceeds cGMP standards. Our lifecycle approach to quality management covers every phase of every process, and each batch SCA Pharma produces must meet stringent release specifications including the following:

- Sterility testing
- Visual Inspection and (Acceptable Quality Limit) AQL
- EM data incorporated into release criteria
- Endotoxin testing
- Potency testing
- Real time stability testing to support Beyond Use Dating
- UV overwrap protection
- Storage conditions

Understanding SCA Pharma's Compounding Processes

SCA Pharma compounds products using Finished Drug Product (FDP) and API (bulk). The API must appear on the FDA's drug shortage list at the time of compounding, distribution and dispensing. The starting pathway for each individual product is clearly defined in the product list under the "Product Source" heading.

In both processes, safety is achieved through proper application of current Good Manufacturing Practices (cGMP). As an additional measure for sterility assurance, SCA Pharma performs a sterile filtration on all products regardless of starting materials prior to filling into final containers.

Finished Drug Product (FDP)

FDP compounding uses an FDA-approved drug as the starting material which is aseptically processed into ready-to-use, single-dose products. This process may also be referred to as compounding from commercial, as the source NDC# can be traced back to a specific commercial drug product.

Key Characteristics of SCA Pharma FDP Products:

- Only FDA-approved commercially available sterile drugs are utilized.
- Source NDC#, traceable to the commercial starting material, is available for every product.
- Only products with a source NDC# traceable to a commercial drug product are delineated as being sourced from an FDP.
- No SCA products labeled as being sourced from FDP are produced from a bulk drug or feature an NDC# traceable to an API. We do not consider sterile filtered or terminally sterilized bulk or pooled products made from API and then filled under sterile aseptic conditions to be from FDP.

Active Pharmaceutical Ingredient (API)

A bulk drug substance purchased from an FDA-registered facility is compounded and sterilized to produce ready-to-use, single-dose products. This process may also be referred to as compounding from powder or non-sterile-to sterile compounding, as the source NDC# is traced back to the starting API.

Key Characteristics of SCA Pharma API Products:

- Each API must be accompanied by a Certificate of Analysis and is tested for potency and identity prior to use in compounding.
- The API must appear on the FDA's drug shortage list at the time of compounding, distribution and dispensing; or
- Appear on the FDA's list of bulk drug substances for which there is a clinical need (503B bulks list).

Products

All products offer both linear and 2D barcodes.



SCA Pharma works closely with medical facilities to ensure the products we produce match the up-to-date needs of our customers. We currently offer admixed pharmaceuticals in the following categories:

Anesthetic

Ketamine

Lidocaine

Antiarrhythmic

Diltiazem

Antibiotic

Vancomycin

Anticoagulant

Sodium Citrate

Beta-Adrenergic Blocking Agents

Labetalol

Electrolyte

Potassium Chloride

Potassium Phosphate

Neuromuscular Blocker

Rocuronium

Succinylcholine

Opioid Analgesic

Fentanyl

Fentanyl + Bupivacaine

Fentanyl + Ropivacaine

Hydromorphone

Methadone

Morphine

Oxytocic Agent

Oxvtocin

Vasopressor

Ephedrine

Epinephrine

Norepinephrine

Phenylephrine



Product List



NDC	Product Description	Product Source	BUD
	Diltiazem		
70004-0541-35	Diltiazem 1 mg/mL (125 mg) in 125 mL 0.9% Sodium Chloride 100 mL Bag (Total Volume 125 mL)	FDP	120 Days
70004-540-35	Diltiazem 1 mg/mL (125 mg) in 125 mL 5% Dextrose 100 mL Bag (Total Volume 125 mL)	FDP	105 Days
	Ephedrine		
70004-0604-09	Ephedrine 5 mg/mL in 0.9% Sodium Chloride 5 mL fill 6 mL Syringe	FDP	90 Days
70004-0604-09-K	Ephedrine 5 mg/mL in 0.9% Sodium Chloride 5 mL fill 6 mL Syringe (Kit Check)	FDP	90 Days
70004-0604-12	Ephedrine 5 mg/mL in 0.9% Sodium Chloride 10 mL fill 12 mL Syringe	FDP	90 Days
70004-0604-12-K	Ephedrine 5 mg/mL in 0.9% Sodium Chloride 10 mL fill 12 mL Syringe (Kit Check)	FDP	90 Days
70004-0605-09	Ephedrine 10 mg/mL in 0.9% Sodium Chloride 5 mL fill 6 mL Syringe	FDP	90 Days
70004-0605-09-K	Ephedrine 10 mg/mL in 0.9% Sodium Chloride 5 mL fill 6 mL Syringe (Kit Check)	FDP	90 Days
	Epinephrine		
70004-613-40	Epinephrine 4 mg (16 mcg/mL) in 0.9% Sodium Chloride 250 mL Bag	FDP	90 Days
70004-608-40	Epinephrine 5 mg (20 mcg/mL) in 0.9% Sodium Chloride 250 mL Bag	FDP	90 Days
70004-607-40	Epinephrine 8 mg (32mcg/mL) in 0.9% Sodium Chloride 250 mL Bag	FDP	90 Days
	Fentanyl (PF)		
70004-0202-22	Fentanyl 10 mcg/mL in 0.9% Sodium Chloride 50 mL fill 60 mL Syringe	FDP	105 Days
70004-0229-32	Fentanyl 10 mcg/mL in 0.9% Sodium Chloride 100 mL Bag	FDP	120 Days
70004-0229-40	Fentanyl 10 mcg/mL in 0.9% Sodium Chloride 250 mL Bag	FDP	120 Days
70004-203-32	Fentanyl 20 mcg/mL in 0.9% Sodium Chloride 100 mL Bag	FDP	120 Days
70004-0200-17	Fentanyl 50 mcg/mL in 25 mL fill 30 mL Syringe	FDP	180 Days
70004-0222-30	Fentanyl 50 mcg/mL in 50 mL DEHP/PVC-Free Bag	FDP	105 Days

NDC	Product Description	Product Source	BUD
70004-0200-32	Fentanyl 50 mcg/mL in 100 mL Bag	FDP	105 Days
70004-0202-32	Fentanyl 10 mcg/mL in 0.9% Sodium Chloride 100 mL Bag	API	120 Days
70004-0202-40	Fentanyl 10 mcg/mL in 0.9% Sodium Chloride 250 mL Bag	API	120 Days
70004-213-32	Fentanyl 20 mcg/mL in 0.9% Sodium Chloride 100 mL Bag	API	120 Days
70004-0205-32	Fentanyl 20 mcg/mL in 0.9% Sodium Chloride 100 mL Bag	API	120 Days
70004-0200-22	Fentanyl 50 mcg/mL in 0.9% Sodium Chloride 50 mL fill 60 mL Syringe	API	105 Days
70004-0211-30	Fentanyl 50 mcg/mL in 0.9% Sodium Chloride 50 mL Bag	API	105 Days
	Fentanyl + Bupivacaine (PF)		
70004-0244-40	Fentanyl 2 mcg/mL + Bupivacaine 0.0625% in 0.9% Sodium Chloride 250 mL Bag	FDP	105 Days
70004-0253-40	Fentanyl 2 mcg/mL + Bupivacaine 0.1% in 0.9% Sodium Chloride 250 mL Bag	FDP	105 Days
70004-0231-32	Fentanyl 2 mcg/mL + Bupivacaine 0.125% in 0.9% Sodium Chloride 100 mL Bag	FDP	105 Days
70004-0254-40	Fentanyl 2 mcg/mL + Bupivacaine 0.125% in 0.9% Sodium Chloride 250 mL Bag	FDP	105 Days
70004-0231-22	Fentanyl 2 mcg/mL + Bupivacaine 0.125% in 0.9% Sodium Chloride 50 mL in 60 mL Syringe	FDP	90 Days
70004-0271-22	Fentanyl 2 mcg/mL + Bupivacaine HCl 0.125% in 0.9% Sodium Chloride 50 mL fill 60 mL Syringe	API	90 Days
70004-0231-40	Fentanyl 2 mcg/mL + Bupivacaine 0.125% in 0.9% Sodium Chloride 250 mL Bag	API	105 Days

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NDC	Product Description	Product Source	BUD
	Fentanyl + Ropivacaine (PF)		
70004-0264-22	Fentanyl 2 mcg/mL + Ropivacaine 0.1% in 0.9% Sodium Chloride 50 mL fill 60 mL Syringe	FDP	105 Days
70004-0260-32	Fentanyl 2 mcg/mL + Ropivacaine 0.2% in 0.9% Sodium Chloride 100 mL Bag	FDP	105 Days
70004-0280-32	Fentanyl 2 mcg/mL and Ropivacaine HCl 0.2% in 0.9% Sodium Chloride 100 mL Bag	API	105 Days
	Hydromorphone		
70004-0300-18	Hydromorphone 0.2 mg/mL in 0.9% Sodium Chloride 30 mL Syringe	FDP	105 Days
70004-0300-16	Hydromorphone 0.2 mg/mL in 0.9% Sodium Chloride 30 mL fill 35 mL Plungerless Syringe	FDP	105 Days
70004-0300-30	Hydromorphone 0.2 mg/mL in 0.9% Sodium Chloride 50 mL Bag	FDP	105 Days
70004-0300-55	Hydromorphone 0.2 mg/mL in 0.9% Sodium Chloride 100 mL Bag	FDP	105 Days
70004-0303-17	Hydromorphone 1 mg/mL in 0.9% Sodium Chloride 25 mL fill 30 mL Syringe	FDP	105 Days
70004-0303-16	Hydromorphone 1 mg/mL in 0.9% Sodium Chloride 30 mL fill 35 mL Plungerless Syringe	FDP	105 Days
70004-0303-21	Hydromorphone 1 mg/mL in 0.9% Sodium Chloride 30 mL fill 60 mL Syringe	FDP	105 Days
70004-0303-30	Hydromorphone 1 mg/mL in 0.9% Sodium Chloride 50 mL Bag	FDP	105 Days
70004-0300-22	Hydromorphone HCl 0.2 mg/mL in 0.9% Sodium Chloride 50 mL fill Syringe (10 mg/50 mL)	FDP	105 Days
	Ketamine		
70004-0435-09	Ketamine 10 mg/mL in 0.9% Sodium Chloride 5 mL fill 6 mL Syringe (50 mg/5 mL)	API	90 Days
70004-0435-12	Ketamine 10 mg/mL in 0.9% Sodium Chloride 10 mL fill 12 mL Syringe (100 mg/10 mL)	API	90 Days
	Labetalol		
70004-0700-28	Labetalol 5 mg/mL 4 mL fill 5 mL Syringe	FDP	190 Days
-			

NDC	Product Description	Product Source	BUD
	Lidocaine		
70004-0720-12	Lidocaine 1% 10 mL fill 12 mL Syringe (PF)	FDP	90 Days
70004-0723-09	Lidocaine 2% 5 mL fill 6 mL Syringe (PF)	FDP	90 Days
70004-0723-09-K	Lidocaine 2% 5 mL fill 6 mL Syringe (PF) (Kit Check)	FDP	90 Days
	Methadone		
70004-0380-05	Methadone 10 mg/mL 1 mL fill 3 mL Syringe	FDP	105 Days
	Morphine		
70004-0100-16	Morphine 1 mg/mL in 0.9% Sodium Chloride 30 mL fill 35 mL Plungerless Syringe	FDP	105 Days
70004-0100-59	Morphine 1 mg/mL in 0.9% Sodium Chloride 100 mL Bag	FDP	105 Days
70004-0103-16	Morphine 5 mg/mL in 0.9% Sodium Chloride 30 mL fill 35 mL Plungerless Syringe	FDP	105 Days
	Norepinephrine		
70004-0781-40	Norepinephrine 4 mg in 0.9% Sodium Chloride 250 mL Bag	FDP	90 Days
70004-0784-40	Norepinephrine 8 mg in 0.9% Sodium Chloride 250 mL Bag	FDP	90 Days
70004-0785-40	Norepinephrine 16 mg in 0.9% Sodium Chloride 250 mL Bag	FDP	190 Days
	Oxytocin		
70004-085-44	Oxytocin 30 units added to 0.9% Sodium Chloride 500 mL Bag	FDP	75 Days

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NDC	Product Description	Product Source	BUD
	Phenylephrine		
70004-0811-12	Phenylephrine 40 mcg/mL in 0.9% Sodium Chloride 10 mL fill 12 mL Syringe (400 mcg/10mL)	FDP	240 Days
70004-0811-12-K	Phenylephrine 40 mcg/mL in 0.9% Sodium Chloride 10 mL fill 12 mL Syringe (400 mcg/10mL) (Kit Check)	FDP	240 Days
70004-0816-12	Phenylephrine 80 mcg/mL in 0.9% Sodium Chloride 10 mL fill 12 mL Syringe	FDP	240 Days
70004-0816-12-K	Phenylephrine 80 mcg/mL in 0.9% Sodium Chloride 10 mL fill 12 mL Syringe (Kit Check)	FDP	240 Days
70004-0810-11	Phenylephrine 100 mcg/mL in 0.9% Sodium Chloride 5 mL fill 12 mL Syringe	FDP	240 Days
70004-0810-11-K	Phenylephrine 100 mcg/mL in 0.9% Sodium Chloride 5 mL fill 12 mL Syringe (Kit Check)	FDP	240 Days
70004-0810-12	Phenylephrine 100 mcg/mL in 0.9% Sodium Chloride 10 mL fill 12 mL Syringe	FDP	240 Days
70004-0810-12-K	Phenylephrine 100 mcg/mL in 0.9% Sodium Chloride 10 mL fill 12 mL Syringe (Kit Check)	FDP	240 Days
70004-0810-22	Phenylephrine 100 mcg/mL in 0.9% Sodium Chloride 50 mL fill 60 mL Syringe	FDP	240 Days
70004-0811-40	Phenylephrine 10 mg in 0.9% Sodium Chloride 250 mL Bag	FDP	95 Days
70004-0808-40	Phenylephrine 20 mg in 0.9% Sodium Chloride 250 mL Bag	FDP	95 Days
70004-0807-40	Phenylephrine 25 mg in 0.9% Sodium Chloride 250 mL Bag	FDP	95 Days
70004-0825-40	Phenylephrine 40 mg in 0.9% Sodium Chloride 250 mL Bag	FDP	95 Days
70004-0806-40	Phenylephrine 50 mg in 0.9% Sodium Chloride 250 mL Bag	FDP	95 Days
70004-0805-40	Phenylephrine 100 mg in 0.9% Sodium Chloride 250 mL Bag	FDP	95 Days
	Potassium Chloride		
70004-0832-40	Potassium Chloride 20 mEq added to 0.9% Sodium Chloride 250 mL Bag	FDP	90 Days
70004-0833-44	Potassium Chloride 40 mEq added to 0.9% Sodium Chloride 500 mL Bag	FDP	90 Days
	Potassium Phosphate		
70004-0841-40	Potassium Phosphate 15 mMol added to 0.9% Sodium Chloride 250 mL Bag	FDP	105 Days

NDC	Product Description	Product Source	BUD
	Rocuronium		
70004-850-09	Rocuronium 10 mg/mL 5 mL fill 6 mL Syringe	FDP	90 Days
70004-850-09-K	Rocuronium 10 mg/mL 5 mL fill 6 mL Syringe (Kit Check)	FDP	90 Days
70004-0850-12	Rocuronium 10 mg/mL 10 mL fill 12 mL Syringe	FDP	90 Days
70004-0850-12-K	Rocuronium 10 mg/mL 10 mL fill 12 mL Syringe (Kit Check)	FDP	90 Days
	Sodium Citrate		
70004-0900-25	Sodium Citrate 4% 3 mL fill 6 mL Syringe	FDP	105 Days
	Succinylcholine		
70004-0908-09	Succinylcholine 20 mg/mL 5 mL fill 6 mL Syringe	FDP	90 Days
70004-0908-09-K	Succinylcholine 20 mg/mL 5 mL fill 6 mL Syringe (Kit Check)	FDP	90 Days
70004-0908-12	Succinylcholine 20 mg/mL 10 mL fill 12 mL Syringe	FDP	90 Days
70004-0908-12-K	Succinylcholine 20 mg/mL 10 mL fill 12 mL Syringe (Kit Check)	FDP	90 Days
	Vancomycin (Refrigerated)		
70004-0920-59	Vancomycin 1 gram added to 0.9% Sodium Chloride 250 mL Bag	FDP	120 Days
70004-0923-59	Vancomycin 1.25 grams added to 0.9% Sodium Chloride 250 mL Bag	FDP	120 Days
70004-0924-59	Vancomycin 1.5 grams added to 0.9% Sodium Chloride 250 mL Bag	FDP	120 Days
70004-0924-44	Vancomycin 1.5 grams added to 0.9% Sodium Chloride 500 mL Bag	FDP	120 Days
70004-0926-44	Vancomycin 1.75 grams added to 0.9% Sodium Chloride 500 mL Bag	FDP	120 Days
70004-0928-44	Vancomycin 2 grams added to 0.9% Sodium Chloride 500 mL Bag	FDP	120 Days

PF = Preservative Free BUD is from the date of product compounding

End Product Testing

To guarantee product quality and safety, each batch is tested against rigorous specification criteria prior to release. SCA Pharma's End Product Testing protocol covers five major components and includes the generation and approval of a Certificate of Conformance (C of C) by Quality Assurance.

C of C acceptance criteria and test results become part of the completed batch record.

End Product Testing components:

Defect-Free

All sterile products must be free of critical/major defects prior to release. Skilled operators inspect 100% of units and an AQL level is independently verified.

Environmental Monitoring

Comprehensive environmental monitoring is performed during compounding per USP <1116> and cGMP.

Endotoxin Limits

All sterile products must pass applicable standards for endotoxin (pyrogen) limits. Test methods have been validated per USP <85> and cGMP guidelines.

Sterility Testing

Scan RDI (Rapid Microbiological Technology) enables quick turnaround time for sterility testing of sterile products which maximizes usable shelf life. Scan RDI usage has been qualified to be equivalent to or better than USP <71> traditional sterility methods.

Potency/Identity Specifications

Scientifically sound and appropriate specifications have been established for all products. Every lot is tested for both strength (potency) and identity by methods which are highly specific.

SCA Pharma's Commitment to Quality

Delivering the highest quality medications to our partners and their patients is SCA Pharma's top priority. Our comprehensive training and personnel qualification criteria ensure full compliance with regulatory requirements on every batch we produce.

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