



BioDerm, Inc. FDA Establishment Registration & Device Listing 2024

Establishment:	BIODERM, INC. 12320 73 rd Court North Largo, FL 33773
Registration Number:	1063299
FEI Number*:	3001949129
Status:	Active
Initial Distributor/Importer:	Yes *Note Firm May Have Additional Establishment Types. Please Review Listings For Further Information.
Date Of Registration Status:	2023
Owner/Operator:	BIODERM, INC. 12320 73rd Court North Largo, FL 33773
Owner/Operator Number:	9032576
<u>Proprietary Name:</u>	CathGrip Double Strap; CathGrip Oval; CathGrip PEG Protect; CathGrip Single Strap; Tube Securement Device; Low Profile CathGrip Single Strap; Low Profile CathGrip Double Strap; Large LVAD Securement Device; Medium LVAD Securement Device; CathGrip Extra Small
Classification Name:	DEVICE, INTRAVASCULAR CATHETER SECUREMENT
Product Code:	KMK
Device Class:	1
Regulation Number:	880.5210
Medical Specialty:	General Hospital
Registered Establishment Name:	BIODERM, INC.
Registered Establishment Number:	1063299
Owner/Operator:	BIODERM, INC.
Owner/Operator Number:	9032576
Establishment Operations:	Contract Manufacturer; Manufacturer



Proprietary Name:

Classification Name: SOLVENT, ADHESIVE TAPE
Product Code: KOX
Device Class: 1
Regulation Number: 878.4730
Medical Specialty: General & Plastic Surgery
Registered Establishment Name: BIODERM, INC.
Registered Establishment Number: 1063299
Owner/Operator: BIODERM, INC.
Owner/Operator Number: 9032576
Establishment Operations: Specification Developer

FreeDerm Adhesive Remover; FreeDerm Bottles; FreeDerm Wipes

Proprietary Name:

Classification Name: DEVICE, PASTE-ON FOR INCONTINENCE, NON-STERILE
Product Code: NOA
Device Class: 1
Regulation Number: 876.5250
Medical Specialty: Gastroenterology
Registered Establishment Name: BIODERM, INC.
Registered Establishment Number: 1063299
Owner/Operator: BIODERM, INC.
Owner/Operator Number: 9032576
Establishment Operations: Contract Manufacturer; Manufacturer

Faceplate Seals; Men's Liberty Acute with CathGrip; Men's Liberty Acute with CathGrip and Towel; Men's Liberty Acute without CathGrip; Men's Liberty; Men's Liberty Acute; BioDerm XLS (Oval)

Proprietary Name:

Classification Name: CLAMP, PENILE
Product Code: FHA
Device Class: 1
Regulation Number: 876.5160
Medical Specialty: Gastroenterology
Registered Establishment Name: BIODERM, INC.
Registered Establishment Number: 1063299
Owner/Operator: BIODERM, INC.
Owner/Operator Number: 9032576
Establishment Operations: Manufacturer

BioDerm Penis Clamp; KindKlamp



Proprietary Name:

Classification Name:	BioPlus+ Barrier Film
Product Code:	TAPE AND BANDAGE, ADHESIVE
Device Class:	KGX
Device Class:	1
Regulation Number:	880.5240
Medical Specialty:	General Hospital
Registered Establishment Name:	BIODERM, INC.
Registered Establishment Number:	1063299
Owner/Operator:	BIODERM, INC.
Owner/Operator Number:	9032576
Establishment Operations:	Specification Developer

Proprietary Name:

Classification Name:	Faceplate Strip
Product Code:	BANDAGE, ELASTIC
Device Class:	FQM
Device Class:	1
Regulation Number:	880.5075
Medical Specialty:	General Hospital
Registered Establishment Name:	BIODERM, INC.
Registered Establishment Number:	1063299
Owner/Operator:	BIODERM, INC.
Owner/Operator Number:	9032576
Establishment Operations:	Manufacturer