

BioDerm, Inc. FDA Establishment Registration & Device Listing 2024

Establishment: BIODERM, INC.

12320 73rd 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

Proprietary Name: 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



<u>Proprietary Name:</u> FreeDerm Adhesive Remover; FreeDerm Bottles; FreeDerm

Wipes

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

<u>Proprietary Name:</u> 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)

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

<u>Proprietary Name:</u> BioDerm Penis Clamp; KindKlamp

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



Proprietary Name: BioPlus+ Barrier Film

Classification Name: TAPE AND BANDAGE, ADHESIVE

Product Code: 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: Faceplate Strip

Classification Name: BANDAGE, ELASTIC

Product Code: 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