

PROTECH LEADED EYEWEAR INC, DBA PROTECH MEDICAL

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FDA REGISTRATION

1. GENERAL

ESTABLISHMENT

Protech Leaded Eyewear, Inc.

Business Trade Name: Protech Medical 1360 North Killian Drive, STE 2 Lake Park, FL 33403

2. PRODUCT LISTINGS

APRONS

Classification Name: Product Code: Device Class: Regulation Number: Medical Specialty: Apron, Protective IWO 1 892.6500 Radiology

BARRIER

Classification Name: Product Code: Device Class: Regulation Number: Medical Specialty: Barrier, Control Panel, X-Ray, Movable IWX 1 892.6500 Radiology

ISOLATION GOWNS

Classification Name:Non-surgical Isolation GownProduct Code:OEADevice Class:1Regulation Number:878.4040Medical Specialty:General & Plastic Surgery

Owner/Operator Number: 10025412 Owner/Operator: Protech Leaded Eyewear, Inc. 1360 North Killian Drive, STE 2 Lake Park, FL 33403

EYEWEAR

Classification Name: Product Code: Device Class: Regulation Number: Medical Specialty: Shield, Eye, Radiological IWS 1 892.6500 Radiology

GLOVES

Classification Name: Product Code: Device Class: Regulation Number: Medical Specialty: Radiographic Protection Glove IWP 1 892.6500 Radiology

Kindly note that all information contained herein is publicly available information which can be found on the FDA's website:

Registration Number: 1063300 FEI Number*: 1000488948

Date of Registration Status: 2021

Status: Active

* Access: https://www.fda.gov/medical-devices/device-registration-and-listing/search-registration-and-listing

* Select: "Search the Registration & Listing Database"

* Input: "Protech Leaded Eyewear" into the Establishment or Tradename Section and click on the Search Button.

NOTICE: The FDA does not market nor endorse company marketing or claims. Be cautious about those including the FDA logo on their literature.