



2020

CERTIFICATE OF REGISTRATION

This certifies that:

VEPRO AG
MAX-PLANCK STR. 1-3
PFUNGSTADT Hessen, GERMANY 64319

is registered with the U.S. Food and Drug Administration for FY 2020 pursuant to Title 21, 807 et seq. of the United States Code of Federal Regulations:

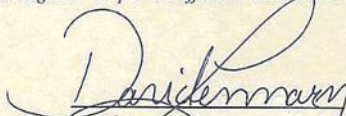
Establishment Registration:	9614981
DUNS No.:	32-166-8584
Device Classification Name:	SYSTEM, IMAGE PROCESSING, RADIOLOGICAL
Product Code:	LLZ
Regulation Number:	892.2050
Official Correspondent and U.S. Agent:	Registrar Corp 144 Research Drive, Hampton, Virginia, 23666, USA Telephone: +1-757-224-0177 • Fax: +1-757-224-0179

Registrar Corp will confirm that such registration remains effective upon request and presentation of this certificate until the end of the year stated above, unless said registration is terminated after issuance of this certificate. Registrar Corp makes no other representations or warranties, nor does this certificate make any representations or warranties to any person or entity other than the named certificate holder, for whose sole benefit it is issued. This certificate does not denote endorsement or approval of the certificate-holder's device or establishment by the U.S. Food and Drug Administration. Registrar Corp assumes no liability to any person or entity in connection with the foregoing.

Pursuant to 21 CFR 807.39, "Registration of a device establishment or assignment of a registration number does not in any way denote approval of the establishment or its products. Any representation that creates an impression of official approval because of registration or possession of a registration number is misleading and constitutes misbranding."

The U.S. Food and Drug Administration does not issue a certificate of registration, nor does the U.S. Food and Drug Administration recognize a certificate of registration. Registrar Corp is not affiliated with the U.S. Food and Drug Administration.


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 David Lennarz
 Executive Director
 Registrar Corp
 Dated: November 18, 2019

Establishment Registration & Device Listing

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Establishment:
 VEPRO AG
 MAX-PLANCK STR. 1-3
 PFUNGSTADT Hesse, DE 64319
Registration Number: 9614981
FEI Number*: 3004562407
Status: Active
Date Of Registration Status: 2024

Owner/Operator:
 VEPRO AG
 MAX-PLANCK STR. 1-3
 PFUNGSTADT, DE 64319
Owner/Operator Number: [9030770](#)

Official Correspondent:
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 Registrar Corp
 144 Research Drive
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Email: David.Lennarz@Registrarcorp.Com

* Firm Establishment Identifier (FEI) should be used for identification of entities within the imports message set

FDA | U.S. Food and Drug Administration



Annual Registration Successful

Facility: VEPRO AG, PFUNGSTADT, Hesse, GERMANY

You have successfully updated your registration and listing information for 2025.

Your registration will be valid through Dec 31, 2025.

Be sure to print this page for your records.

The next registration renewal period is October 1 - December 31, 2025.

Registering your facility and listing devices does not, in any way, constitute FDA approval of your facility or devices.

You may contact the FDA with any questions at reglist@cdrh.fda.gov.

The Owner/Operator Number for this Registration is: 9030770.

Facility Information

Registration Number: 9614981
 Initial Importer: N
 Facility Name: VEPRO AG
 Address: MAX-PLANCK STR. 1-3, --
 PFUNGSTADT, Hesse, 64319,
 GERMANY
 Foreign Trade Zone: N

Owner/Operator Information

Owner/Operator Number: 9030770
 Contact Name: GERLINDE - GABRIEL
 Company: VEPRO AG
 Address: MAX-PLANCK STR. 1-3, --
 PFUNGSTADT, 64319, GERMANY
 Telephone: 49 - 615 - 7800600
 Fax: -
 E-mail: Accounting@vepro.com

Official Correspondent Information

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 DUNS Number: 139242874

United States Agent Information

Contact First Name: David
 Contact Last Name: Lennarz
 Contact Title: Mr
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 Phone: 757 - 2240177
 Fax: 757 -2240179
 DUNS Number: 139242874
 E-mail: david.lennarz@registrarcorp.com

Device Listings

Listing Number	Premarket Submission Number	Premarket Submission Type	Product Code(s)	Device Name(s)	Activities	Importers
D061761	K972215		LLZ	System, image processing, radiological	Manufacturer	VEPRO North America, Inc.