

DRYPRO Σ II



DRYPRO III is a new design in the range of Konica Minolta's Dry Laser Imagers that fits the needs of medical professionals looking for a high resolution table top printer.

The DRYPRO III creates images of unrivalled sharpness, utilizing the latest precision optics from Konica Minolta to produce a 50-µm pixel pitch; the highest available resolution.

The new compact and highly efficient Laser Imager is easy to operate, offers an intuitive workflow and is available with a wide range of film sizes.



High-Quality images 50-μm

A semiconductor laser is used to produce a 50-µm pixel pitch (508 dpi), resulting in high image resolution.

Powerful image-processing algorithms are utilized to simultaneously optimize both the image smoothness as well as the text sharpness.

Diagnostic clarity is preserved and patient data is always readable, regardless of the print size.



Four film sizes and two trays

Variety of four film sizes: 14x17", 11x14", 10x12" and 8x10".

The DRYPRO ΣII is equipped with two film trays so that two different film sizes can be used simultaneously.

Consistent quality

The DRYPRO ΣII density control function maintains the output density via automatic measurement. The system also automatically calibrates whenever a tray of film with a new lot-number is loaded.

User-friendly design

The DRYPRO ΣII is easy to operate and offers an intuitive workflow. Film exchange requires a simple cartridge insertion. The innovative cartridges allow easy film size adjustment and support various modalities such as CR, CT, Ultrasound and MRI.

Space-saving & fast

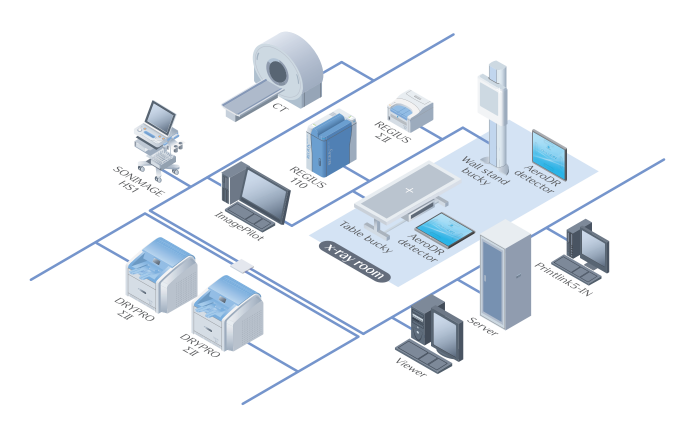
DRYPRO ΣII is a desktop printer and combines reliability and convenience with remarkable operating efficiency, all in a compact body. It features a



footprint as small as 65 x 63 cm² and is designed for use in small clinics and high throughput general hospitals with a speed up to 110 sheets/hour (for 8 x 10").

Network functions designed for open and flexible environments

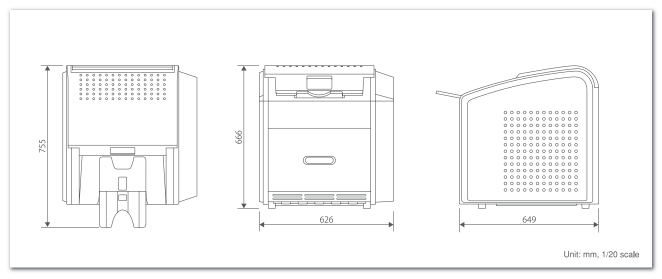
DRYPRO III is a networked imager, capable of connecting directly to DICOM print compliant devices. In addition to DICOM basic grayscale print functions, the DRYPRO ΣII supports Presentation LUT, which enables printed film to more accurately match diagnostic monitors.



PRODUCT SPECIFICATIONS

- Exposure source Semiconductor Laser
- Film size 14"x17"(35x43cm), 11"x14"(28x35cm), 10"x12"(25x30cm), and 8"x10"(20x25cm) selectable
- Film Dry Imaging Recording film SD-S / 125sheets per cartridge
- Pixel size 50µm (508 dpi)
- Processing capability 14x17-inch size sheets approximately 70 sheets/hour 8x10-inch size sheets approximately 110 sheets/hour
- Network connection Ethernet 10 BASE-T / 100BASE-T / 1000BASE-T
- Film supply Daylight setting
- Number of supply trays 2ch
- Output grades 14bit (16384 grades)
- Network connectivity DICOM Print Management SCP
- Operating condition 15-33°C 20-80% RH, non-condensing
- Power 90-130V(50/60Hz) 9A / 180-264V (50/60Hz) 4.5A
- Dimensions W:626mm x D:649mm x H:666mm
- Weight 79 kg
- Footprint 0.41m²

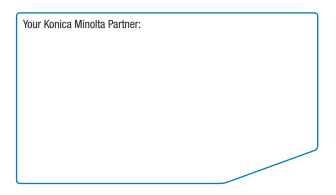
DIMENSIONS



Brand Name: DRYPRO Σ II : LASER IMAGER Туре : DRYPRO SIGMA 2 Model

Storing and handling unused film

Unused film should be stored in a cool, dark place (25° or below), where it will not be affected by radiation.



www.konicaminolta.eu





Certificate

KONICA MINOLTA Medical & Graphic Imaging Europe B.V.

We hereby confirm that

Sergiu Sorocovici

has successfully participated in a technical training from December 17thth to 18th, 2014 for

Drypro Σ, Drypro 832, Drypro 873, Printlink 5 (TTDP&TTDS)

Training content:

- Installation
 Configuration
- Adjustments Troubleshooting
 - Basic Application QC

Munich, December 20th, 2014

Patrick Winkler Service Manager EMEA Hans-Joachim Kock Trainer



REGISTRUL DE STAT AL DISPOZITIVELOR MEDICALE

Введите текс	т для п	оиска												
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DM000349456	5	FILME PENTRU REPRODUCERE DIN TEHNOLOG LASER	II			SD-Q 11X14" MEDICAL FILM 50 SH)	A3R7	Jap	ponia	KONICA MINOLTA, INC.	INTERMED S.R.L.	Rg04-000097	15-04-2022
DM000349465	5	FILME PENTRU REPRODUCERE DIN TEHNOLOG LASER	II			SD-P 10X12" MEDICAL FILM 125 SH		A5FJ	Jap	ponia	KONICA MINOLTA, INC.	INTERMED S.R.L.	Rg04-000097	15-04-2022
DM000349470	0	FILME PENTRU REPRODUCERE DIN TEHNOLOG LASER	II			SD-P 14X17" MEDICAL FILM 125 SH		A5FR	Jap	ponia	KONICA MINOLTA, INC.	INTERMED S.R.L.	Rg04-000097	15-04-2022
DM000349460	0	FILME PENTRU REPRODUCERE DIN TEHNOLOG LASER	II			SD-Q 14X17" MEDICAL FILM 125 SH		A3RD	Jap	ponia	KONICA MINOLTA, INC.	INTERMED S.R.L.	Rg04-000097	15-04-2022
DM000349455	5	FILME PENTRU REPRODUCERE DIN TEHNOLOG LASER	II			SD-Q 10X12" MEDICAL FILM 125 SH		A3R6	Jap	ponia	KONICA MINOLTA, INC.	INTERMED S.R.L.	Rg04-000097	15-04-2022
DM000349454	4	FILME PENTRU REPRODUCERE DIN TEHNOLOG LASER	II			SD-Q 8X10" MEDICAL FILM 125 SH		A3R5	Jap	ponia	KONICA MINOLTA, INC.	INTERMED S.R.L.	Rg04-000097	15-04-2022
DM000349476	5	SCANER DE IMAGINI CU LASER				LASER IMAGER DRYPRO SIGMA2			Jap	ponia	KONICA MINOLTA, INC.	INTERMED S.R.L.	Rg04-000097	15-04-2022



EU DECLARATION OF CONFORMITY

Manufacturer

Name

KONICA MINOLTA, INC.

Address

1 Sakura-machi, Hino-shi, Tokyo, 191-8511, Japan

Single Registration Number

Pending

declares, sole responsibility, that the following product

Generic Device Group:

Laser Imagers

Type:

LASER IMAGER

Model (Product Name):

DRYPRO SIGMA 2

Basic UDI-DI: Intended Purpose:

4560141920000688T
The device is intended for use in the acquisition and process of

radiographic images of human anatomy. It is intended to replace radiographic film/screen system in general-purpose diagnostic

procedures.

Classification:

Class I, Rule 1, according to Annex VIII of REGULATION (EU)

2017/745

Serial Number:

from A9R4-201363 to A9R4-999999 (A9R4) from A9R5-201363 to A9R5-999999 (A9R5)

referred to in this declaration conforms with the following EU law(s):

REGULATION (EU) 2017/745, confirmed by the procedure of its Annex IX,

DIRECTIVE 2014/53/EU and Directive 2011/65/EU

and conforms with the following standard(s):

EN ISO 13485:2016, EN ISO 14971:2012, EN 1041:2008,

EN ISO 15223-1:2016, EN 60601-1:2006+A1:2013, EN 60601-1-2:2015,

EN 60601-1-6:2010+A1:2015, EN 60825-1:2007, EN 62366:2008+A1:2015,

EN 62304:2006 for REGULATION (EU) 2017/745,

EN 300 330 V2.1.1 for DIRCTIVE 2014/53/EU,

EN IEC 63000:2018 for Directive 2011/65/EU

and that this declaration is valid upon approval for release of each product.

EU Representative

Name Konica Minolta Business Solutions Europe GmbH

Address Capellalaan 65, 2132 JL, Hoofddorp, The Netherlands

Single Registration Number NL-AR-000002026

Signed for and on behalf of manufacturer:

Tokyo Japan, 2021-04-22

(Place and date of issue)

HAJIME NOZAWA

General Manager.

Quality Assurance Operations

Healthcare Business Unit

Healthcare Business Headquarters

(Name, function)

(Signature of equivalent authorized by the manufacturer)



Quality Management System EN ISO 13485:2016

Registration No.:

SX 2003379-1

Organization:

KONICA MINOLTA, INC.

1 Sakura-machi, Hino-shi, Tokyo, 191-8511 Japan

Scope:

Design, Development and Manufacture of Direct Digitizers (Computed Radiography Systems and Digital Radiography Systems), Radiographic Film Processors, Diagnostic Ultrasound Transducers, Ultrasound Needle Guides, Picture Archiving and Communication Systems (PACS), Software for Picture Archiving and Communication Systems (PACS), Diagnostic X-ray Digital Imaging System Workstation, Software for Diagnostic X-ray Digital Imaging System Workstation, Print Management Systems, Pulse Oximeters, Software for Pulse Oximeters, Physiological patient Monitor and Jaundice Meters

Design, Development and Manufacture of Fluoroscopic Image Storage Devices (Plate, Cassette and Cassette Plate)

Design, Development, Manufacture and Distribution of Laser Imagers, Diagnostic X-ray Systems, Diagnostic Ultrasound Systems and Pulse Oximeter Probes

Manufacture and Distribution of Medical Imaging Films and X-ray Films (X-ray Films for General Radiography, X-ray Films for Photofluorography and Dental X-ray Films for General Use)

The Certification Body of TÜV Rheinland LGA Products GmbH certifies that the organization has established and applies a quality management system for medical devices.

Proof has been furnished that the requirements specified in the abovementioned standard are fulfilled. The quality management system is subject to yearly surveillance.

Report No.:

150247687-301

Effective date:

2022-04-19

Expiry date:

2025-04-18

Issue date:

2022-04-11

DAKKS

Deutsche
Akkreditierungsstelle
D-ZM-14169-01-02

Michiaki Aihara
TÜV Rheinland LGA Products GmbH
Tillystraße 2 · 90431 Nürnberg · Germany

Maihara



Quality Management System EN ISO 13485:2016

Registration No.:

SX 2003379-1

Organization:

KONICA MINOLTA, INC.

1 Sakura-machi Hino-shi, Tokyo, 191-8511 Japan

The scope of certification also covers the following:

No.

Facility

/01-1

c/o KONICA MINOLTA, INC.

Tokyo site 1 Sakura-machi, Hino-shi, Tokyo, 191-8511 Japan

Design and Development of Radiographic Film Processors. Picture Archiving and Communication Systems (PACS), Software for Picture Archiving and Communication Systems (PACS), Diagnostic X-ray Digital Imaging System Workstation, Software for Diagnostic X-ray Digital Imaging System Workstation, Print Management Systems, Pulse Oximeters, Software for Pulse Oximeters, Physiological patient Monitor and Jaundice Meters

Design and Development of Fluoroscopic Image Storage Devices (Cassette)

Design and Development and Distribution of Laser Imagers, Diagnostic X-ray Systems, Diagnostic Ultrasound Systems and Pulse Oximeter Probes

Distribution of Medical Imaging Films, X-ray Films (X-ray Films for General Radiography, X-ray Films for Photofluorography and Dental X-ray Films for General Use)

Design, Development and Manufacture of Direct Digitizers (Computed Radiography Systems and Digital Radiography Systems), Diagnostic Ultrasound Transducers and Ultrasound Needle Guides

Design, Development and Manufacture of Fluoroscopic Image Storage Devices (Plate and Cassette Plate)

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2022-04-11

Deutsche Akkreditierungsstelle D-ZM-14169-01-02

TÜVRheinland Machano Michiaki Aihara

TÜV Rheinland LGA Products GmbH

Tillystraße 2 · 90431 Nürnberg · Germany



Quality Management System EN ISO 13485:2016

Registration No.:

SX 2003379-1

Organization:

KONICA MINOLTA, INC.

1 Sakura-machi Hino-shi, Tokyo, 191-8511 Japan

/01-2

c/o KONICA MINOLTA, INC.

Tokyo site

2970 Ishikawa-machi, Hachioji-shi, Tokyo 192-8505 Japan

Design and Development of Radiographic Film Processors. Picture Archiving and Communication Systems (PACS), Software for Picture Archiving and Communication Systems (PACS), Diagnostic X-ray Digital Imaging System Workstation, Software for Diagnostic X-ray Digital Imaging System Workstation, Print Management Systems, Pulse Oximeters, Software for Pulse Oximeters, Physiological patient Monitor and Jaundice Meters

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Distribution of Medical Imaging Films, X-ray Films (X-ray Films for General Radiography, X-ray Films for Photofluorography and Dental X-ray Films for General Use)

Design, Development and Manufacture of Direct Digitizers (Computed Radiography Systems and Digital Radiography Systems), Diagnostic Ultrasound Transducers and Ultrasound Needle Guides

Design, Development and Manufacture of Fluoroscopic Image Storage Devices (Plate and Cassette Plate)

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SX 2003379-1

Organization:

KONICA MINOLTA, INC.

1 Sakura-machi Hino-shi, Tokyo, 191-8511 Japan

The scope of certification also covers the following:

/01-3

c/o KONICA MINOLTA TECHNOPRODUCTS CO..

LTD. Tokyo site 1 Sakura-machi. Hino-shi, Tokyo, 191-8511 Japan Design and Development of Radiographic Film Processors. Picture Archiving and Communication Systems (PACS), Software for Picture Archiving and Communication Systems (PACS). Diagnostic X-ray Digital Imaging System Workstation, Software for Diagnostic X-ray Digital Imaging System Workstation, Print Management Systems, Pulse Oximeters, Software for Pulse Oximeters, Physiological patient Monitor and Jaundice Meters

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Michiaki Aihara

TÜV Rheinland LGA Products GmbH Tillystraße 2 · 90431 Nürnberg · Germany



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Registration No.:

SX 2003379-1

Organization:

KONICA MINOLTA, INC.

1 Sakura-machi Hino-shi, Tokyo, 191-8511 Japan

The scope of certification also covers the following:

/02

c/o KONICA MINOLTA TECHNOPRODUCTS CO.,

LTD.

Sayama site 2-2-1 Hirosedai, Sayama-shi, Saitama 350-1328 Japan

Manufacture of Direct Digitizers (Computed Radiography Systems and Digital Radiography Systems), Laser Imagers, Radiographic Film Processors, Diagnostic X-ray Systems, Diagnostic Ultrasound Systems, Diagnostic Ultrasound Transducers, Ultrasound Needle Guides, Picture Archiving and Communication Systems (PACS), Software for Picture Archiving and Communication Systems (PACS), Diagnostic X-ray Digital Imaging System Workstation, Software for Diagnostic X-ray Digital Imaging System Workstation, Print Management Systems. Pulse Oximeters, Pulse Oximeter Probes, Software for Pulse Oximeters, Physiological patient Monitor and Jaundice Meters

Manufacture of Fluoroscopic Image Storage Devices (Plate, Cassette and Cassette Plate)

/03

c/o KONICA MINOLTA **TECHNOPRODUCTS**

CO., LTD. Kuki site 6201-6 Sanga, Shobu-cho, Kuki-shi. Saitama 346-0104 Japan

Manufacture of Direct Digitizers (Computed Radiography Systems and Digital Radiography Systems) and Laser Imagers

Manufacture of Medical Imaging Films, X-ray Films (X-ray Films for General Radiography, X-ray Films for Photofluorography and Dental X-ray Films for General Use)

TÜVRheinland

Report No.: 150247687-301 Effective date: 2022-04-19 Expiry date: 2025-04-18 Issue date: 2022-04-11

> TÜV Rheinland LGA Products GmbH Tillystraße 2 · 90431 Nürnberg · Germany



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