



## Schedule of Devices

The above Manufacturer's Declaration is valid for the following devices:

Identification of the device Device name / Basic UDI-DI (under MDR application)	MDR Device classification	If the MDR device is a substitute device, identification of the corresponding MDD device	MDD Certificate Reference(s) of the devices under MDR application, and the NB Identification (Name and number)	Original expiry date as indicated on the Directive Certificate prior to the extension of the validity (if applicable)	End date of extended validity/transition period
LifePort Kidney Transporter 1.1 (LKT101P)  081504502LKT101PZN	Ila	LifePort Kidney Transporter LKT101P  (Only Name Change)	Certificate: 10000320758-PA-NA-NOR  NB: DNV Product Assurance AS NB #: 2460	27 May 2024	31 December 2028
LifePort Kidney Transporter Disposable Perfusion Pack (LKT200)  LifePort Kidney Transporter Perfusion Pack with Optional Oxygenation Tubing (LKT200X)  081504502LKT200VQ	Ila	LifePort Kidney Transport Disposable Perfusion Pack LKT200  LifePort Kidney Transporter Disposable Perfusion Pack with Optional Oxygenation Tubing LKT200X  (Only Name Change)	Certificate: 10000320758-PA-NA-NOR  NB: DNV Product Assurance AS NB #: 2460	27 May 2024	31 December 2028



<p>LifePort Kidney Transporter Disposable SealRing Cannula (7x20) (CAN0720)</p> <p>LifePort Kidney Transporter Disposable SealRing Cannula (10x35) (CAN1035)</p> <p>LifePort Kidney Transporter Disposable Straight Cannula (3mm) (CAN0003)</p> <p>LifePort Kidney Transporter Disposable Straight Cannula (5mm) (CAN0005)</p> <p>LifePort Kidney Transporter Disposable Straight Cannula (8mm) (CAN0008)</p> <p>LifePort Kidney Transporter Disposable Coupler (CAN1000)</p> <p>LifePort Kidney Transporter Disposable Universal SealRing Cannula (3mm) (UCAN0003)</p> <p>LifePort Kidney Transporter Disposable</p>	Ila	<p>LifePort Kidney Transporter Disposable Cannula</p> <p>CAN0720 CAN1035 CAN0003 CAN0005 CAN0008 UCAN0003 UCAN0005 UCAN0007 UCAN0009</p> <p>LifePort Kidney Transporter Disposable Flexible Couplers CAN1000</p>	<p>Certificate: 10000320758-PA-NA-NOR</p> <p>NB: DNV Product Assurance AS NB #: 2460</p>	27 May 2024	31 December 2028
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<p>Universal SealRing Cannula (5mm) (UCAN0005)</p> <p>LifePort Kidney Transporter Disposable Universal SealRing Cannula (7mm) (UCAN0007)</p> <p>LifePort Kidney Transporter Disposable Universal SealRing Cannula (9mm) (UCAN0009)</p> <p>081504502Cannula6S</p>		<p>(Only Name Change)</p>			
<p>LifePort Kidney Transporter Disposable Sterile Drape (LKT300)</p> <p>081504502LKT300VV</p>	<p>Is</p>	<p>LifePort Kidney Transporter Disposable Sterile Drape LKT300</p>	<p>Certificate: 10000320758-PA-NA-NOR</p> <p>NB: DNV Product Assurance AS NB #: 2460</p>	<p>27 May 2024</p>	<p>31 December 2028</p>



# EC CERTIFICATE

## Full Quality Assurance System

Certificate No.: 10000320758-PA-NA-NOR

Project No.: PRJC-509108-2014-MSL-NOR

Valid Until: 27 May 2024

This is to certify that the quality system of:

### **Organ Recovery Systems, Inc.**

One Pierce Place, Suite 475W. Itasca, IL 60143 USA

For design, production and final product inspection/testing of:

### **LIFEPORTR KIDNEY TRANSPORTER, INCLUDING ACCESSORIES**

Has been assessed with respect to:

### **The conformity assessment procedure described in Annex II excluding section 4 of Council Directive 93/42/EEC on Medical Devices, as amended**

and found to comply

Further details of the product(s) and conditions for certification are given overleaf.

Place and date:  
**Høvik, 12 April 2021**

For the issuing office:  
**Notified Body 2460  
DNV Product Assurance AS**



**Palani Damodharan**  
Principal Assessor

Notice: The Certificate is subject to terms and conditions as set out in the Certification Agreement. Failure to comply may render this Certificate invalid.

NOTIFIED BODY 2460: DNV Product Assurance AS, Veritasveien 3, 1363 Høvik, Norway, Tel +47 67 57 88 00, [www.dnv.com](http://www.dnv.com)

ICP-4-5-i1-MDD-f2, rev.0

## Jurisdiction

Application of Council Directive 93/42/EEC of 14 June 1993, adopted as “Forskrift om Medisinsk Utstyr” by the Norwegian Ministry of Health and Care Services.

## Certificate history:

Revision	Description	Issue Date
0.0	Initial Certification	07 May 2020
1.0	Addition of Kidney Transporter	28 May 2020
2.0	Extension in scope – <b>LKT200X</b> new products and editorial corrections ( <b>changes in bold</b> )	12 April 2021

## Products covered by this Certificate:

Product Description	Product Name	Class
<b>LifePort</b> Kidney Transporter GMDN 16461	LKT100P LKT101P	Ila
<b>LifePort Kidney Transport Disposable</b> Perfusion Pack GMDN 48039	LKT200	Ila
<b>LifePort Kidney Transporter Disposable Perfusion Pack with Optional Oxygenation Tubing</b> GMDN 48039	<b>LKT200X</b>	<b>Ila</b>
<b>LifePort Kidney Transporter Disposable Cannula</b> GMDN 48039	CAN0720 CAN1035 CAN0003 CAN0005 CAN0008 UCAN0003 UCAN0005 UCAN0007	Ila

Certificate No.: 10000320758-PA-NA-NOR  
 Place and date: Høvik, 12 April 2021

	UCAN0009	
<b>LifePort Kidney Transporter</b> Disposable Flexible Couplers GMDN 48039	CAN1000	Ila
<b>LifePort Kidney Transporter</b> Disposable Sterile Drape GMDN 48039	LKT300	Is

The complete list of devices is filed with the Notified Body

**Sites covered by this certificate**

Site Name	Address
Organ Recovery System, Inc.	One Pierce Place, Suite 475W, Itasca, IL 60143, USA

**EU Representative**

Emergo Europe, Prinsessegracht 20, 2514 AP The Hague, The Netherlands

## Terms and conditions

The certificate is subject to the following terms and conditions:

- Any producer (see 2001/95/EC for a precise definition) is liable for damage caused by a defect in his product(s), in accordance with directive 85/374/EEC, as amended, concerning liability of defective products.
- The certificate is only valid for the products and/or manufacturing premises listed above.
- The Manufacturer shall fulfil the obligations arising out of the quality system as approved and uphold it so that it remains adequate and efficient.
- The Manufacturer shall inform the Notified Body of any intended updating of the quality system and the Notified Body will assess the changes and decide if the certificate remains valid.
- Periodical audits will be held, in order to verify that the Manufacturer maintains and applies the quality system. the Notified Body reserves the right, on a spot basis or based on suspicion, to pay unannounced visits.

The following may render this Certificate invalid:

- Changes in the quality system affecting production.
- Periodical audits not held within the allowed time window.

## Conformity declaration and marking of product

When meeting with the terms and conditions above, the producer may draw up an EC declaration of conformity and legally affix the CE mark followed by the Notified Body identification number.

End of Certificate