

EU Technical Documentation Assessment Certificate

mdc medical device certification GmbH

Kriegerstr. 6, 70191 Stuttgart, Germany Notified body (identification number 0483)

hereby certifies that the company (SRN: DE-MF-000005179)

PHS Medical GmbH

Ederweg 3 34277 Fuldabrück Germany

has submitted a technical documentation for the devices listed on the following pages in accordance with Annexes II and III of Regulation (EU) 2017/745, which fulfils the following requirements:

Annex IX - Chapter II (Assessment of the Technical Documentation)

of Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices.

The certificate consists of 3 pages. Details of the devices affected by this certificate as well as further information and conditions are included on the following pages.

 Valid from:
 2023-05-24
 Registration No.
 D1069500035

 Valid until:
 2028-05-23
 Evaluation Report No.
 P22-00991-240013

Stuttgart, 2023-05-24

Head of Notified Body





Devices:

Product:

NuPort-CT with and without Introducer

Intended purpose:

NuPort®/C-Port® port systems are subcutaneously implantable medical devices used when repeated access to the vascular system is indicated as part of therapy. These products provide the user with an easy-to-find site for cannula insertion to administer fluids, such as drugs, into the vascular system or to collect blood samples. The port is accessed by percutaneous puncture with a (non-coring) port cannula. The NuPort®/C-Port® port systems are available as standard vascular access ports, which are NOT suitable for high pressure injections, and as port systems for CECT high pressure injections. High pressure injections are performed using an appropriate infusion set with a non-coring port cannula. Port systems are intended for single use and are explanted and disposed of at the end of therapy.

Risk class: III

Basic-UDI-DI: 42503746PortsystemTitanNF

Product:

NuPort-LP-CT with and without Introducer

Intended purpose:

NuPort®/C-Port® port systems are subcutaneously implantable medical devices used when repeated access to the vascular system is indicated as part of therapy. These products provide the user with an easy-to-find site for cannula insertion to administer fluids, such as drugs, into the vascular system or to collect blood samples. The port is accessed by percutaneous puncture with a (non-coring) port cannula. The NuPort®/C-Port® port systems are available as standard vascular access ports, which are NOT suitable for high pressure injections, and as port systems for CECT high pressure injections. High pressure injections are performed using an appropriate infusion set with a non-coring port cannula. Port systems are intended for single use and are explanted and disposed of at the end of therapy.

Risk class: III

Basic-UDI-DI: 42503746PortsystemTitanNF

Product:

C-Port-CT with and without Introducer

Intended purpose:

NuPort®/C-Port® port systems are subcutaneously implantable medical devices used when repeated access to the vascular system is indicated as part of therapy. These products provide the user with an easy-to-find site for cannula insertion to administer fluids, such as drugs, into the vascular system or to collect blood samples. The port is accessed by percutaneous puncture with a (non-coring) port cannula. The NuPort®/C-Port® port systems are available as standard vascular access ports, which are NOT suitable for high pressure injections, and as port systems for CECT high pressure injections. High pressure injections are performed using an appropriate infusion set with a non-coring port cannula. Port systems are intended for single use and are explanted and disposed of at the end of therapy.

Risk class: III

Basic-UDI-DI: 42503746PortsystemPSUC6



Product:

C-Port with and without Introducer

Intended purpose:

NuPort®/C-Port® port systems are subcutaneously implantable medical devices used when repeated access to the vascular system is indicated as part of therapy. These products provide the user with an easy-to-find site for cannula insertion to administer fluids, such as drugs, into the vascular system or to collect blood samples. The port is accessed by percutaneous puncture with a (non-coring) port cannula. The NuPort®/C-Port® port systems are available as standard vascular access ports, which are NOT suitable for high pressure injections, and as port systems for CECT high pressure injections. High pressure injections are performed using an appropriate infusion set with a non-coring port cannula. Port systems are intended for single use and are explanted and disposed of at the end of therapy.

Risk class: III

Basic-UDI-DI: 42503746PortsystemTitanNF

Product:

C-Port II with and without Introducer

Intended purpose:

NuPort®/C-Port® port systems are subcutaneously implantable medical devices used when repeated access to the vascular system is indicated as part of therapy. These products provide the user with an easy-to-find site for cannula insertion to administer fluids, such as drugs, into the vascular system or to collect blood samples. The port is accessed by percutaneous puncture with a (non-coring) port cannula. The NuPort®/C-Port® port systems are available as standard vascular access ports, which are NOT suitable for high pressure injections, and as port systems for CECT high pressure injections. High pressure injections are performed using an appropriate infusion set with a non-coring port cannula. Port systems are intended for single use and are explanted and disposed of at the end of therapy.

Risk class: III

Basic-UDI-DI: 42503746PortsystemPSUC6

Product:

Percutaneous introducer set consisting of puncture cannula, split sheath, guide wire with dispenser and insertion aid, tunneler, syringe

Intended purpose:

The Introducer sets from PHS Medical GmbH are sterile accessories used exclusively for the insertion of the intravascular catheters included in the subcutaneously implantable port systems from PHS Medical GmbH. The product is used to assist the physician in implanting the PHS Medical port systems through a minimally invasive procedure known as the Seldinger technique.

Risk class: III

Basic-UDI-DI: 42503746IntroducerYF

Notes:

For the placing on the market of the devices an EU Quality Management System Certificate according to Annex IX, Chapter I of Regulation (EU) 2017/745 on medical devices is also required.