

RA 133/2024  
2025-08-20

**EU DECLARATION OF CONFORMITY**  
medical devices rev. 2

**Manufacturer's name:** HTL-Strefa S.A,  
ul. Adamówek 7,  
95-035 Ozorków,  
Poland

**SRN:** PL-MF-000002198

**Device's name:** safety lancets, personal lancets  
(as per the Product list)

**Classification:** IIa

**Classification rule:** 6

We, herewith declare under the sole responsibility of the manufacturer that the stated medical devices meet the provisions of Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices, amending Directive 2001/83/EC, Regulation (EC) No 178/2002 and Regulation (EC) No 1223/2009 and repealing Council Directives 90/385/EEC and 93/42/EEC.

This Declaration of Conformity is supported by the Quality Management System based on the standard for medical devices, confirmed by following Certificates:

- EN ISO 13485:2016, Certificate No.: SX 1023950-1 issued by TÜV Rheinland LGA Products GmbH
- ISO 13485:2016 (MDSAP), Certificate No.: MD 1023950-1-1 issued by TUV Rheinland of North America, Inc.

This Declaration of Conformity is valid for all products concerned bearing the CE Marking of Conformity specified in the annexed product list.

CE marked devices are covered by the EU Certificate Quality Management System, REGULATION (EU) 2017/745 on Medical Devices in accordance with Annex IX, Chapter I, Section 2 and 3, reference number: HZ 1023950-1 issued for first time on 2024-08-02. The certification was delivered by TÜV Rheinland LGA Products GmbH, Tillystraße 2, D-90431 Nürnberg, German, Notified Body Identification Number 0197 on the products concerned conforming to the required Technical Documentation and meeting the relevant provisions of Regulation (EU) 2017/745.

On behalf of HTL-Strefa S.A.  
Justyna Żemigala  
Regulatory Affairs Manager /  
Person Responsible for Regulatory Compliance

Ozorków,

Place, date 2025-08-20



Authorized signature, Name & Position

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**PRODUCT LIST**

This product list belongs to the EU Declaration of Conformity identified by document no RA 133/2024 and specifies the CE marked products concerned that HTL-Strefa S.A. intends to distribute in conformity with the provisions of the Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices, amending Directive 2001/83/EC, Regulation (EC) No 178/2002 and Regulation (EC) No 1223/2009 and repealing Council Directives 90/385/EEC and 93/42/EEC.  
The following list identifies the products by name and device type.

Device name and type / intended purpose	Commercial Name	Dimension of the needle	GMDN / EMDN code	Basic UDI-DI
<b>Safety lancet type 420</b> sterile, single-use medical devices intended for capillary blood sampling	<ul style="list-style-type: none"> <li>Haemolance Plus</li> <li>Safe-T-Lance Plus</li> </ul>	18G / 21G / 25G / 28G / 1.5 mm blade	61579 / V0104	590799609420M8, 590799609420XFY
	<ul style="list-style-type: none"> <li>McKesson Safety Lancets PUSH BUTTON</li> </ul>	28G		
<b>Safety lancet type 430</b> sterile, single-use medical devices intended for capillary blood sampling	<ul style="list-style-type: none"> <li>Prolance</li> </ul>	18G / 21G / 25G / 28G 1,5 mm blade	61579 / V0104	590799609430MB, 590799609430XG5
	<ul style="list-style-type: none"> <li>Single-Let</li> </ul>	28G		
<b>Safety lancet type 450</b> sterile, single-use medical devices intended for capillary blood sampling	<ul style="list-style-type: none"> <li>ergo Lance</li> </ul>	21G / 25G / 30G	61579 / V0104	590799609450MH, 590799609450XGF
<b>Safety lancet type 520</b> sterile, single-use medical devices intended for capillary blood sampling	<ul style="list-style-type: none"> <li>MediSafe Solo</li> </ul>	23G / 28G / 29G	61579 / V0104	590799609520MD, 590799609520XG7
	<ul style="list-style-type: none"> <li>MenaLancet Pro</li> <li>Securlancets unik</li> </ul>	23G / 29G		
	<ul style="list-style-type: none"> <li>diamet mySafety</li> <li>MedicoFine</li> </ul>	29G		
	<ul style="list-style-type: none"> <li>Single-Let Next</li> </ul>	28G		
<b>Safety lancet type 532</b> sterile, single-use medical devices intended for capillary blood sampling	<ul style="list-style-type: none"> <li>myLance</li> </ul>	23G / 28G	61579 / V0104	590799609532ML, 590799609532XGJ

Device name and type / intended purpose	Commercial Name	Dimension of the needle	GMDN / EMDN code	Basic UDI-DI
<b>Safety lancet type 545-549</b> sterile, single-use medical devices intended for capillary blood sampling	<ul style="list-style-type: none"> <li>Medlance</li> </ul>	21G / 23G / 28G 1,5 mm blade	61579 / V0104	5907996095455493K
<b>Safety lancet type 553-556</b> sterile, single use medical devices intended for capillary blood sampling	<ul style="list-style-type: none"> <li>Medlance Plus</li> </ul>	21G / 25G / 30G 0,8 mm blade	61579 / V0104	5907996095535563D, 590799609553556X4F
	<ul style="list-style-type: none"> <li>GlucoSmart</li> </ul>	21G / 30G		
	<ul style="list-style-type: none"> <li>Safe Digitest</li> </ul>	21G / 25G / 30G		
	<ul style="list-style-type: none"> <li>Solofix Safety</li> </ul>	21G/25G/ 0,8 mm blade		
<b>Safety lancet type 610</b> sterile, single-use medical devices intended for capillary blood sampling	<ul style="list-style-type: none"> <li>Acti-Lance</li> </ul>	17G / 23G / 28G	61579 / V0104	590799609610MF, 590799609610XG9
	<ul style="list-style-type: none"> <li>Safe Digitest PLUS</li> </ul>	23G / 28G		
<b>Personal lancet type 560</b> sterile, single-use medical devices intended to be used with a lancing	<ul style="list-style-type: none"> <li>droplet</li> </ul>	28G / 30G / 33G	61579 / V0104	590799609560MR, 590799609560XGT
	<ul style="list-style-type: none"> <li>Penlancet</li> </ul>	28G / 33G		
	<ul style="list-style-type: none"> <li>MyStar SylkFeel</li> </ul>	30G		
	<ul style="list-style-type: none"> <li>MenaLancet</li> </ul>			
	<ul style="list-style-type: none"> <li>Microdot</li> </ul>			
	<ul style="list-style-type: none"> <li>GlucoSmart fine</li> </ul>	33G		
	<ul style="list-style-type: none"> <li>Glucoject Lancets PLUS</li> </ul>			

On behalf of HTL-Strefa S.A.  
Justyna Żemigala  
Regulatory Affairs Manager /  
Person Responsible for Regulatory Compliance

Ozorków,

Place, date 2025-08-20



Authorized signature, Name & Position

The statement to the EC Declaration of Conformity (DoC No.: RA 133/2024; 2025-08-20)

We, HTL-STREFA.S.A. with its registered office at Adamówek no. 7, 95-035 Ozorków, Poland hereby declare that, Solofix Safety Lancet Type 553-556 are covered by the EU Declaration of Conformity (Doc No.: RA 133/2024; 2025-08-20) EU DoC of Solofix Safety Lancet which we issued and can be placed in the market as a medical device according to the Medical Device Regulation (EU)2017/745

B.Braun items	B. Braun Product name	HTL items	HTL Product name	EMDN Code
6183010DE	Solofix Safety Fine	7132	Safety lancet type 553 25G	61579/V0104
6183020DE	Solofix Safety Neonat	7133	Safety lancet type 556 0,8mm blade	61579/V0104
6183000DE	Solofix Safety Universal	7131	Safety lancet type 554 21G	61579/V0104

Yours faithfully



Anna Linke-Szewczyk  
Regulatory Affairs Senior Specialist  
Regulatory Affairs Department of HTL-Strefa S.A.