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**Statement**

To whom it may concern,

We HTL-STREFA S.A., located at Adamówek 7, 95-035 Ozorków, Poland  
manufacturer of:

Haemolance Plus safety lancet type 420

hereby confirm that our products are in conformity with European Union Medical  
Device Directive 93/42/EEC amendment by 2007/47/EC as per EC Certificate reference  
number: 84587CE01 issued by DEKRA Certification B.V., Arnhem, The Netherlands,  
Notified Body Identification Number 0344.

Our current EC Certificate is valid until 24 May 2024 therefore, our products  
indicated on the EC Certificate may still be placed on the EU market in accordance with  
article 120 of Medical Device Regulation (MDR 2017/745).

In the same time HTL-Strefa S.A. would like to inform that we are in transition period  
to adopt the requirements of the MDR 2017/745 in EU.

The process of obtaining EC certificate complies with Medical Device Regulation (EU)  
2017/745 (MDR) is pending.

Yours sincerely,



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Justyna Żemigala

Regulatory Affairs Manager