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 Zentralstelle der Länder  
 für Gesundheitsschutz  
 bei Arzneimitteln und  
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 BS-MDR-099



Product Service

## EU Quality Management System Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapters I and III  
 (Class IIa and Class IIb Devices)

**No. G10 044751 0176 Rev. 00**

<b>Device(s):</b> Diagnostic Ultrasound System	<b>Risk Classification:</b> IIa	<b>CND code:</b> Z110401
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**The validity of this certificate depends on conditions and/or is limited to the following:** ./.

<b>Revision History including Changes:</b>	Revision / Issue Date / Report Rev. 00 / 2019-11-21 / SH1905502
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