

Certificate



Quality Management System EN ISO 13485:2016

Registration No.: SX 1307131-1


Organization: Viant Aura GmbH
Staatsstr. 5
97773 Aura
Germany

The scope of certification also covers the following:

No.	Facility	Scope
/01	c/o Viant Aura GmbH Staatsstr. 5 97773 Aura Germany	Contract manufacturer of stainless steel components for medical devices based on seamless and welded precision tubes, as well as section tubes, turned tube parts, formed tube parts and tube assemblies.
/02	c/o Viant Aura GmbH Struthberg 19 97773 Aura Germany	Contract manufacturer of stainless steel components for medical devices based on section tubes and turned tube parts, as well as cleaning and packaging under controlled environmental conditions.

Report No.: 3336040-90
Effective date: 2021-03-16
Expiry date: 2024-03-15
Issue date: 2021-03-16




Roland Gruber
TÜV Rheinland LGA Products GmbH
Tillystraße 2 · 90431 Nürnberg · Germany

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Organization: Viant Aura GmbH
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Scope: Contract manufacturer of stainless steel components for medical devices based on seamless and welded precision tubes, as well as section tubes, turned tube parts, formed tube parts and tube assemblies.

The Certification Body of TÜV Rheinland LGA Products GmbH certifies that the organization has established and applies a quality management system for medical devices. Proof has been furnished that the requirements specified in the abovementioned standard are fulfilled. The quality management system is subject to yearly surveillance.

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A blue ink handwritten signature, appearing to read 'Roland Gruber', written over a horizontal line.

Roland Gruber
TÜV Rheinland LGA Products GmbH
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