



## **EU Declaration of Conformity** for product Anchor Tips, 300uL

REF

10036506

**Basic UDI-DI** 

91201239109889J4



**STRATEC Consumables GmbH** Sonystrasse 20, 5081 Anif, Austria

SRN: AT-MF-000023689

	of Anchor Tips, 300uL take sole res visions of the following regulation:	ponsibility for and hereby declare that Anchor Tips,
• REGULATION	I (EU) 2017/746 on in vitro medical	devices
RISK Class		
$\boxtimes$ A $\square$ B $\square$ C $\square$ D		
	uL are validated for their intended cessories, they are classified as class	d use in combination with the KleeYa System. In theirs A IVD medical devices.
CONFORMITY ROUTE		
☐ ANNEX IX Technica	l Documentation Examination	
$\square$ ANNEX IX Full Qua	lity System	
☐ ANNEX XI Producti	on Quality System	
⊠ ANNEX I & II + III		
Tips, 300uL in combin purposes requires a se	ation with the KleeYa System. The	mance Requirements is demonstrated for the Anchor application of the KleeYa System for in vitro diagnostic ording to REGULATION (EU) 2017/746 for the complete in combination with (e.g. assay).
was issued. Changes affecti		nd the regulatory requirements effectual at the date the Declaration ations trigger a review of the conformity assessment the Declaration
Date:	2022-05-25	stratec
Place:	Anif	consumables

Signature: Thomas Ehrenfeld

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