

EU Declaration of Conformity

with the Medical Device Regulation (MDR) 2017/745

We,	Wolturnu	s A/S, Skalhuse 31, 9240	Nibe, Denmark - DK-MF-0000	25274
	Name and Ad	ddress of Manufacturer – Sir	ngle Registration Number (SRN)	_
hereby, unde	er our sole respo	onsibility declare, that the	product	
		Manual Whee	elchair	
		Category Na	те	_
with model na	mes and referen	ce		
	Model	Basic UDI-DI	UDI-DI	
	Tukan A	57138250017G	5713825570485	
	Tukan B	57138250017G	5713825570492	
	Tukan C	57138250017G	5713825570508	1
	Tukan D	57138250017G	5713825570515	
	Tukan SL	57138250017G	5713825613106	
	Tukan Tilt	57138250017G	5713825570522	
	•	dical Device Regulation (N following standards DS/EN ISO 14	MDR) 2017/745 as class I medic 971:2019	al devices
		Standard(s)	
unable to wa	lk or who have a	a mobility problem. It is d	nded to provide mobility to per esigned for individual use, and The manual wheelchair can be	it can be
		rmity was written accordi t the premises of the man	ng to Annex 4 in MDR, and all s ufacturer.	supporting
Manufact	urer			2 /
February 28, 2	2024 - Nibe	Peter Libak, CEO	1,2/1	

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Signature of

authorized person

Name and position of

authorized person

Date and place of issue