




## Declaration of Conformity

Manufacturer	<b>ArjoHuntleigh AB</b> Hans Michelsensgatan 10 211 20 Malmö, Sweden
Single Registration Number	SE-MF-000000696
Declaration	ArjoHuntleigh AB as the manufacturer of the following medical devices, takes sole responsibility and declares conformity with the applicable provisions of Medical Device Regulation (EU) 2017/745 concerning medical devices.
Device Family Name	<b>Maxi Twin Compact, Non-AC-powered patient lift</b> Models: KTCBxxxxx-xx
Basic UDI-DI	<b>5060693520112VK</b>
GMDN Number and Term	<b>GMDN Number: 12330</b> Mobile patient lifting system, battery-powered
Additional Information	Also complies with the following EU Legislation: Machinery Directive 2006/42/EC RoHS Directive 2011/65/EU
Risk Class and Rule	 Class I, Rule 13

APPROVED BY		
Title: Quality Director	Signature:	Electronically signed by: Anna Nowotna Reason: I am signing as approver of this document Date: Jul 23, 2024 13:58 GMT+2
Name: Anna Nowotna	Date: 	
Title: Senior Regulatory Compliance Manager	Signature:	Electronically signed by: david moynham Reason: I am signing as approver of this document Date: Jul 23, 2024 12:04 GMT+1
Name: David Moynham	Date: 	

On behalf of ArjoHuntleigh AB: Place: Poznan