


<b>DECLARATION OF CONFORMITY</b>			
<b>Product Name:</b> Safehip Active		<b>Product no.:</b> 8190	
<b>APPROVED BY</b> TYTEX		<b>APPROVED BY</b> CUSTOMER	<b>VALID FROM</b> 30. April 2021
		<b>VERSION NO.</b> 1.1	

**DECLARATION OF CONFORMITY**

According to Regulation 2017/745 on Medical Devices (MDR), Tytex A/S hereby declares to fulfil all relevant requirements for the below mention products as set out in Annex I General safety and performance requirements, Annex II Technical documentation, Annex III Technical documentation on post-market surveillance and Annex XIV Clinical evaluation and post-market follow-up.

The Safehip Active are class I, rule I, non-sterile CE marked products intended as hip protection for elderly people (protection).

SRN: DK-MF-000003620

Product number	Product name	Size	Colour	Basic UDI-DI
8190 50 – 01	Safehip Active	S	Black	57039361285150AAJN
8190 50 – 03	Safehip Active	M	Black	57039361285150AAJN
8190 50 – 05	Safehip Active	L	Black	57039361285150AAJN
8190 50 – 07	Safehip Active	XL	Black	57039361285150AAJN
8190 50 – 09	Safehip Active	2XL	Black	57039361285150AAJN

This declaration is issued under the sole responsibility of Tytex A/S



Kim Remin Ankjaer  
 VP of Quality & Environment  
 Legal Representative

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