

Baxter

CARBON SPINE SURGICAL TABLETOP



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DIMENSIONS TABLETOP OVERALL (L X W X H)	CLASS I MEDICAL DEVICE WITH INTERNAL POWER SUPPLY, IPS
Dimensions Side Rails	
EU Side Rail	25 mm x 10 mm (1" x 0.4")
US Side Rail	28.6 mm x 9.5 mm (1.1" x 0.4")
Japan Side Rail	32 mm x 9 mm (1.3" x 0.4")
Net Weight	Ca. 95 kg (209.4 lbs)
Maximum Load	
Side Rail	60 Nm
Coupling	35 Nm
Patient Weight Capacity	
In Zero Position	225kg (496 lbs)
With Longitudinal Shift	180 kg (397 lbs)

OR TABLE COLUMN [Software Version 6.x.1.0 or higher]

The Carbon Spine Tabletop is compatible with the TS7500 column.

Height Adjustment	620 mm – 1170 mm (24.5" - 46")
Tilt	15°
Trendelenburg / Reverse Trendelenburg	± 20°

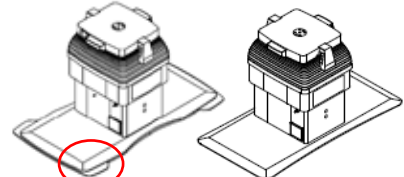
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Longitudinal Shift

With Stationary Column	
In Preferred Direction ¹	800 mm (31.5")
With Head Adapter	500 mm (19.6")
Not In Preferred Direction ¹	650 mm (23.6")
With Head Adapter	350 mm (13.7")
With Mobile Column ²	
Base Plate With Ears ³	500 mm (19.6")
With Head Adapter	200 mm (7.8")
Base Plate Without Ears ⁴	600 mm (23.0")
With Head Adapter	350 mm (13.7")
Adjustment Speed	80 mm/s (3.1"/s)

X-ray Area Head End

With Stationary Column	
In Preferred Direction ¹	1642 mm x 432 mm (64.6" x 17")
Not in Preferred Direction ¹	1492 mm x 432 mm (58.6" x 17")
With Mobile Column ²	
In Any Direction	1442 mm x 432 mm (56.7" x 17")



³ Base plate with ears

⁴ Base plate without ears

¹ In the preferred direction, the radiolucent area of the OR tabletop (head end) is on the right of the OR table column viewed from the column keypad. Please refer to the IFU for the Carbon Spine OR Tabletop for details.

² The preferred direction is not relevant when a mobile column is used. The head-end extension of the tabletop is restricted due to its construction type.

For more information please contact your local Baxter representative.

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The Carbon Spine is intended to be used by clinicians and medically qualified personnel. This medical device is a regulated healthcare product which, pursuant to such regulation bears a CE mark. Baxter recommends that you carefully read the detailed instructions for safe and proper use included in the documents accompanying the medical devices. The personnel of healthcare establishments are responsible for the proper use and maintenance of these medical devices. Baxter reserves the right to make changes without notice in design, specifications and models. The only warranty Baxter makes is the express written warranty extended on the sale or rental of its products. Baxter Medical Systems GmbH + Co.KG.