





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Choose stronger fixation.^{†,‡,1}

Consult this chart to transition from your current device to the stronger absorbable fixation^{†,‡} of the MaxTack™ motorized fixation device.

Former product	Former product description	Number of tacks	Former product code
 Securestrap™ device [§]	Securestrap™ absorbable strap fixation device 7.2 mm tack length	25	STRAP25
 OptiFix™ device [∅]	OptiFix™ absorbable fixation system 6.7 mm tack length	15 30	0113127 0113126
 SorbaFix™ device [¶]	SorbaFix™ absorbable fixation system 6.0 mm tack length	15 30	0113115 0113116

Medtronic product	Medtronic product description	Number of tacks	Medtronic product code
 MaxTack™ motorized fixation device	Motorized fixation device with 30 absorbable tacks	30	MAXTACK30

Make the switch today – contact your Medtronic representative or visit us at [medtronic.eu](https://www.medtronic.eu)

† Compared to competitive SorbaFix™, OptiFix™, Securestrap™, CapSure™[®] and Medtronic absorbable ReliaTack™ devices.
 ‡ Based on benchtop data, not necessarily indicative of human clinical outcomes.
 § Licensed as Ethicon Securestrap Absorbable Strap Fixation Device.
 ∅ Licensed as OptiFix Absorbable Fixation System.
 ¶ Licensed as SorbaFix Absorbable Fixation System, Open, with Absorbable Fasteners.
 1. Based on internal report #RE00437048, MaxTack™ device fixation strength claims. January 2023.

Please note that the intended use of a product may vary depending on geographical approvals. See the device manual(s) for detailed information regarding the intended use, the (implant) procedure, indications, contraindications, warnings, precautions, and potential adverse events. If a device is eligible for eIFU usage, instructions for use can be found at Medtronic's website manuals.medtronic.com. Medtronic products placed on European markets bear the CE mark and the UKCA mark (if applicable).
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