**FDA cleared for treatment of Major Depressive Disorder - (Standard TMS)**MagPro® stimulators R20, R30, R30 with MagOption, X100, and X100 with MagOption are all FDA 510(k) cleared for the treatment of Major Depressive Disorder in adult patients who have failed to receive satisfactory improvement from prior antidepressant medication in the current episode (k170114, k150641 and k171481).

**FDA cleared for treatment of Major Depressive Disorder – (Theta Burst)**MagPro® stimulators R30 with theta burst option, X100 and X100 with MagOption are all FDA 510(k) cleared for the treatment of Major Depressive Disorder in adult patients who have failed to receive satisfactory improvement from prior antidepressant medication in the current episode (k173620)

**FDA cleared for stimulation of peripheral nerves for diagnostic purposes**

MagPro® stimulators R20, R30, R30 with MagOption, X100, and X100 with MagOption are all FDA 510(k) cleared for stimulation of peripheral nerves for diagnostic purposes (k160280, k061645, k091940)

The use of MagPro devices for other than FDA cleared intended uses is considered investigational. In accordance with US federal regulations, an IDE and/or IRB approval may be required for use of MagPro devices for uses distinct from the FDA cleared Indication(s) for Use. The sponsor is responsible for obtaining appropriate, required approvals. Please consult FDA's website ([www.fda.gov](https://www.fda.gov/medicaldevices/deviceregulationandguidance/howtomarketyourdevice/investigationaldeviceexemptionide/default.htm)) for more information.

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