## Viewpoint

# **The Problem of Funding Off-label Deep Brain Stimulation** Bait-and-Switch Tactics and the Need for Policy Reform

P. Justin Rossi, BA<sup>1</sup>; James Giordano, PhD<sup>2</sup>; Michael S. Okun, MD<sup>3</sup>

#### $\gg$ Author Affiliations

" (C) (~

**JAMA Neurol Published Online: January 2017** 2017;74;(1):9-10. doi:10.1001/jamaneurol.2016.2530

#### 

Deep brain stimulation (DBS) is currently approved by the US Food and Drug Administration to treat Parkinson disease, essential tremor, and dystonia. However, so-called off-label use of DBS may be permissible under research-based or compassionate use guidelines to treat severe, medication-refractory cases of other neurological and psychiatric disorders such as Tourette syndrome and obsessive-compulsive disorder.

While affording promising outcomes, DBS surgery and its associated postoperative care is expensive. Mean initial surgical costs are US \$65000 per patient, and battery replacements can add an additional \$10000 to \$20000 in costs during the first 36 months postimplantation (depending on brain target and amount of electricity required).<sup>1</sup> These costs can be daunting because governmental and commercial insurance providers are reluctant to subsidize off-label therapies. Coverage depends on preauthorization requests that require exhaustive documentation of a patient's medical history and peer-to-peer review with an insurance provider's medical director.

### Get Access

View Full Text | Download PDF





