



March 16, 2017

Food and Drug Administration
10903 New Hampshire Avenue
Document Control Center - WO66-G609
Silver Spring, MD 20993-0002

NuVasive Incorporated
Mary Adams
Regulatory Affairs Consultant
7475 Lusk Blvd.
San Diego, California 92121

Re: K162313

Trade/Device Name: NuVasive Next Generation NVM5 System
Regulation Number: 21 CFR 874.1820
Regulation Name: Surgical Nerve Stimulator/Locator
Regulatory Class: Class II
Product Code: PDQ, ETN, GWF, HAW, IKN, OLO
Dated: February 15, 2017
Received: February 16, 2017

Dear Ms. Adams:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

<http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm> for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

<http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>.

Sincerely,

Michael J. Hoffmann -S

for Carlos L. Peña, PhD, MS
Director
Division of Neurological
and Physical Medicine Devices
Office of Device Evaluation
Center for Devices and Radiological Health

Enclosure

Indications for Use

510(k) Number (if known)
K162313

Device Name
NuVasive® Next Generation NVM5 System

Indications for Use (Describe)

The Next Generation NVM5® System is a medical device that is intended for intraoperative neurophysiologic monitoring during spinal surgery, neck dissections, thoracic surgeries, and upper and lower extremities. The device provides information directly to the surgeon, to help assess a patient's neurophysiologic status. NVM5 provides this information by electrically stimulating nerves via electrodes located on surgical accessories and monitoring electromyography (EMG), transcranial or lumbar motor evoked potential (MEP) or somatosensory evoked potential (SSEP) responses of nerves. The System also integrates Bendini™ software used to locate spinal implant instrumentation for the placement of spinal rods.

- **XLIF (Detection)** — The XLIF (Detection) function allows the surgeon to locate and evaluate spinal nerves, and is used as a nerve avoidance tool.

- **Basic & Dynamic Screw Test** — The Screw Test functions allow the surgeon to locate and evaluate spinal nerves by providing proximity information before, during or after bone preparation and placement of bone screws.

- **Free Run EMG** — The Free Run EMG function identifies spontaneous EMG activity of spinal nerves by continually displaying alive stream waveform of any mechanically induced myotome contractions.

- **Twitch Test (Train of Four)** — The Twitch Test function allows the surgeon to assess moderate degrees of neuromuscular block in effect by evaluating muscle contraction following a train of four stimulation pulses.

- **MEP** — Transcranial or lumbar (i.e., conus in region of L1-L2) stimulation techniques for motor evoked potentials are used to assess for acute dysfunction in axonal conduction of the corticospinal tract and peripheral nerves. The ME? function provides an adjunctive method to allow the surgeon to monitor spinal cord and motor pathway integrity during procedures with a risk of surgically induced motor injury.

- **SSEP** — The SSEP function allows the surgeon to assess sensory spinal cord function in surgical procedures during which the spinal cord is at risk.

- **Remote Access** - The remote monitoring and local wireless control provides real-time capabilities to the NG-NVM5 System for additional physicians.

- **Bendini** — The Bendini Spinal Rod Bending function is used to locate spinal implant system instrumentation (screws, hooks) to determine their relative location to one another to generate bend instructions to shape a spinal rod. A surgeon is able to use those instructions and bend a rod using the Bendini Bender, a mechanical rod bender.

Type of Use (Select one or both, as applicable)

Prescription Use (Part 21 CFR 801 Subpart D)

Over-The-Counter Use (21 CFR 801 Subpart C)

CONTINUE ON A SEPARATE PAGE IF NEEDED.

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510(k) Summary

In accordance with Title 21 of the Code of Federal Regulations, Part 807, and in particular 21 CFR §807.92, the following summary of information is provided:

A. Submitted by:

Mary Adams
Regulatory Affairs Consultant
NuVasive, Incorporated
7475 Lusk Blvd.
San Diego, California 92121
Telephone: (909) 991-5235
Date Prepared: March 13, 2017

B. Device Name

Trade or Proprietary Name: *NuVasive® Next Generation NVM5® System*
Common or Usual Name: Neurological surgical monitor;
Stereotaxic Instrument
Classification Name: Surgical Nerve Stimulator/Locator;
Evoked response electrical stimulator;
Neurological stereotaxic instrument;
Electromyography (EMG) monitor/stimulator
Device Class: Class II
Classification: §874.1820, §882.1870, §882.4560, §890.1375
Product Code: PDQ, ETN, GWF, HAW, IKN, OLO

C. Predicate Devices

The subject *NuVasive Next Generation (NG) NVM5 System* is substantially equivalent to the following predicate devices:

NuVasive NVM5 System K152942 (Primary Predicate)
Medtronic NIM 3.0 Nerve Integrity Monitor K083124 (Additional Predicate)
NuVasive NVM5 System K132694 (Reference Predicate for current density of the cutaneous electrodes)

D. Device Description

The *NG-NVM5 System* is a medical device that is intended for intraoperative neurophysiologic monitoring during spinal surgery, neck dissections, thoracic surgeries, and upper and lower extremities. The device provides information directly to the surgeon, to help assess a patient's neurophysiologic status. *NG-NVM5* provides this information by electrically stimulating nerves via electrodes located on surgical accessories and monitoring electromyography (EMG), motor evoked potential (MEP) or somatosensory evoked potential (SSEP) responses of nerves. Moreover, a Twitch Test function is utilized to test the ability of the nerve to respond, or contract, following four stimulation pulses to determine the presence of neuromuscular block.

The System also integrates Bendini® software used to locate spinal implant instrumentation for the placement of spinal rods as well as providing guidance to support the delivery of pedicle

screws during EMG monitoring. Lastly, the system offers real-time control capabilities (local wireless control) to the NG-NVM5 System by additional monitoring physicians, in addition to remote monitoring capabilities.

In summary, the *NG-NVM5 System* includes the following five (5) software functionalities / modalities:

1. Electromyography (EMG)
2. Motor Evoked Potential (MEP)
3. Somatosensory Evoked Potential (SSEP)
4. Remote Access
5. Bendini

The *NG-NVM5 System* hardware consists of a Patient Module (PM) and computer, as well as accompanying accessory components which consist of an assortment of disposable conductive probes, electrodes, and electrode leads.

E. Intended Use

The Next Generation NVM5® System is a medical device that is intended for intraoperative neurophysiologic monitoring during spinal surgery, neck dissections, thoracic surgeries, and upper and lower extremities. The device provides information directly to the surgeon, to help assess a patient's neurophysiologic status. NVM5 provides this information by electrically stimulating nerves via electrodes located on surgical accessories and monitoring electromyography (EMG), transcranial or lumbar motor evoked potential (MEP) or somatosensory evoked potential (SSEP) responses of nerves. The System also integrates Bendini™ software used to locate spinal implant instrumentation for the placement of spinal rods.

- **XLIF (Detection)** – The XLIF (Detection) function allows the surgeon to locate and evaluate spinal nerves, and is used as a nerve avoidance tool.
- **Basic & Dynamic Screw Test** – The Screw Test functions allow the surgeon to locate and evaluate spinal nerves by providing proximity information before, during or after bone preparation and placement of bone screws.
- **Free Run EMG** – The Free Run EMG function identifies spontaneous EMG activity of spinal nerves by continually displaying a live stream waveform of any mechanically induced myotome contractions.
- **Twitch Test (Train of Four)** – The Twitch Test Function allows the surgeon to assess moderate degrees of neuromuscular block in effect by evaluating muscle contraction following a train of four stimulation pulses.
- **MEP** – Transcranial or lumbar (i.e., conus in region of L1-L2) stimulation techniques for motor evoked potentials are used to assess for acute dysfunction in axonal conduction of the corticospinal tract and peripheral nerves. The MEP function provides an adjunctive method to allow the surgeon to monitor spinal cord and motor pathway integrity during procedures with a risk of surgically induced motor injury.
- **SSEP** – The SSEP function allows the surgeon to assess sensory spinal cord function in surgical procedures during which the spinal cord is at risk.

- Remote Access - The remote monitoring and local wireless control provides real-time capabilities to the NG-NVM5 System for additional physicians.
- Bendini – The Bendini Spinal Rod Bending function is used to locate spinal implant system instrumentation (screws, hooks) to determine their relative location to one another to generate bend instructions to shape a spinal rod. A surgeon is able to use those instructions and bend a rod using the Bendini Bender, a mechanical rod bender.

F. Technological Characteristics

As was established in this submission, the subject *NG-NVM5 System* is substantially equivalent to other predicate devices cleared by the FDA for commercial distribution in the United States. The subject device was shown to be substantially equivalent and have equivalent technological characteristics to its predicate device through comparison in areas including design, intended use, material composition, and functions. The technological differences within this 510(k) that were shown to be substantially equivalent to the predicates include additional angular and offset spinal parameter assessment.

Table 1 – Comparison of Technical Characteristics

Specification/ Property	Predicate Device	Predicate Device	Subject Device
	Medtronic NIM 3.0 Nerve Monitoring System (K083124)	NuVasive NVM5 System (K152942 and K132694)	NuVasive NG-NVM5 System
Intended Use / Indications for Use	<p>Intended Use: The NIM 3.0 is intended for locating and identifying cranial and peripheral motor and mixed motor-sensory nerves during surgery, including spinal cord and spinal cord nerve roots. The APS electrode is an accessory intended for providing automatic periodic stimulation to nerves when used with the Medtronic Nerve Monitoring Systems.</p> <p>Indications for NIM 3.0 EMG Monitoring Procedures include:</p> <p>Intracranial, Extracranial, Intratemporal, Extratemporal, Neck Dissections, Thoracic Surgeries, and Upper and Lower Extremities</p> <p>Indications for Spinal procedures which may use NIM 3.0 EMG monitoring include:</p> <p>Degenerative Treatments, Pedicle Screw Procedures, Fusion Cages, Rhizotomy, Orthopedic Surgery, Open and Percutaneous Lumbar and Cervical Surgical Procedures, and Thoracic Surgical Procedures.</p>	<p>The NVM5® System is a medical device that is intended for intraoperative neurophysiologic monitoring during spinal surgery. The device provides information directly to the surgeon, to help assess a patient’s neurophysiologic status. NVM5 provides this information by electrically stimulating nerves via electrodes located on surgical accessories and monitoring electromyography (EMG), transcranial or lumbar motor evoked potential (MEP), or somatosensory evoked potential (SSEP) responses of nerves. The System also integrates Bendini® software used to locate spinal implant instrumentation for the placement of spinal rods.</p> <ul style="list-style-type: none"> • XLIF (Detection) – The XLIF (Detection) function allows the surgeon to locate and evaluate spinal nerves, and is used as a nerve avoidance tool. • Basic & Dynamic Screw Test – The Screw Test functions allow the surgeon to locate and evaluate spinal nerves by providing proximity information before, during or after bone preparation and placement of bone screws. • Free Run EMG – The Free Run EMG function identifies spontaneous EMG activity of spinal nerves by continually displaying a live stream waveform of any mechanically induced myotome contractions. • Twitch Test (Train of Four) – The Twitch Test Function allows the surgeon to assess moderate degrees of neuromuscular block in effect by evaluating muscle contraction following a train of four stimulation pulses. • MEP – Transcranial or lumbar (i.e., conus in region of L1-L2) stimulation techniques for motor evoked potentials are used to assess for acute dysfunction in axonal conduction of the corticospinal tract and peripheral nerves. The MEP function provides an adjunctive method to allow the surgeon to monitor spinal cord and motor pathway integrity during procedures with a risk of surgically induced motor injury. • SSEP – The SSEP function allows the surgeon to assess sensory spinal cord function in surgical procedures during which the spinal cord is at risk. • Remote Reader – The Remote reader function provides real time remote access to the NVM5 System for a monitoring physician outside of the operating room • Guidance – The Guidance function is intended as an aid for use in either open or percutaneous pedicle cannulation procedures in the lumbar and sacral spine (L1-S1) of adult patients, and when used in conjunction with radiographic imaging and EMG, allows the surgeon to assess the angulation of system accessories relative to patient spinal anatomy for the creation of a cannulation trajectory for bone screw placement. • Bendini – The Bendini Spinal Rod Bending function is used to locate spinal implant system instrumentation (screws, hooks) to determine their relative location to one another to generate bend instructions to shape a spinal rod. A surgeon is able to use those instructions and bend a rod using the Bendini Bender, a mechanical rod bender. 	<p>The <i>Next Generation NVM5® System</i> is a medical device that is intended for intraoperative neurophysiologic monitoring during spinal surgery, neck dissections, thoracic surgeries, and upper and lower extremities. The device provides information directly to the surgeon, to help assess a patient’s neurophysiologic status. <i>NG-NVM5</i> provides this information by electrically stimulating nerves via electrodes located on surgical accessories and monitoring electromyography (EMG), transcranial or lumbar motor evoked potential (MEP) or somatosensory evoked potential (SSEP) responses of nerves. The System also integrates Bendini™ software used to locate spinal implant instrumentation for the placement of spinal rods.</p> <ul style="list-style-type: none"> • XLIF (Detection) – The XLIF (Detection) function allows the surgeon to locate and evaluate spinal nerves, and is used as a nerve avoidance tool. • Basic & Dynamic Screw Test – The Screw Test functions allow the surgeon to locate and evaluate spinal nerves by providing proximity information before, during or after bone preparation and placement of bone screws. • Free Run EMG – The Free Run EMG function identifies spontaneous EMG activity of spinal nerves by continually displaying a live stream waveform of any mechanically induced myotome contractions. • Twitch Test (Train of Four) – The Twitch Test Function allows the surgeon to assess moderate degrees of neuromuscular block in effect by evaluating muscle contraction following a train of four stimulation pulses. • MEP – Transcranial or lumbar (i.e., conus in region of L1-L2) stimulation techniques for motor evoked potentials are used to assess for acute dysfunction in axonal conduction of the corticospinal tract and peripheral nerves. The MEP function provides an adjunctive method to allow the surgeon to monitor spinal cord and motor pathway integrity during procedures with a risk of surgically induced motor injury. • SSEP – The SSEP function allows the surgeon to assess sensory spinal cord function in surgical procedures during which the spinal cord is at risk. • Remote Access - The remote monitoring and local wireless control provides real-time capabilities to the NG-NVM5 System for additional physicians. • Bendini – The Bendini Spinal Rod Bending function is used to locate spinal implant system instrumentation (screws, hooks) to determine their relative location to one another to generate bend instructions to shape a spinal rod. A surgeon is able to use those instructions and bend a rod using the Bendini Bender, a mechanical rod bender.

Specification/ Property	Predicate Device	Predicate Device	Subject Device
	Medtronic NIM 3.0 Nerve Monitoring System (K083124)	NuVasive NVM5 System (K152942 and K132694)	NuVasive NG-NVM5 System
Software Modalities / Functionalities	<ul style="list-style-type: none"> Stimulating EMG (manual and APS) Free run EMG 	<ul style="list-style-type: none"> XLIF Detection Basic & Dynamic Screw Test Free Run EMG Twitch Test (Train of Four) MEP SSEP Remote Reader Guidance Bendini <ul style="list-style-type: none"> NuvaMap O.R. NuvaLine spinal parameter assessment tools (optional) 	<ul style="list-style-type: none"> XLIF (Detection) Basic & Dynamic Screw Test Free Run EMG Twitch Test MEP SSEP Remote Access (<i>Remote Monitoring and Local Wireless Control</i>) Bendini <ul style="list-style-type: none"> NuvaMap O.R.
Algorithms	<ul style="list-style-type: none"> APS 	<ul style="list-style-type: none"> XLIF Detection Basic & Dynamic Screw Test Free Run EMG Twitch Test (Train of Four) MEP SSEP Guidance Bendini – Rod bending and spinal parameter assessment 	<ul style="list-style-type: none"> XLIF Detection Basic & Dynamic Screw Test Free Run EMG Twitch Test (Train of Four) MEP SSEP Remote Access Bendini – Rod bending and spinal parameter assessment
Total Available Channels	Up to 8	32	32
Headbox/ Patient Module	Unknown	Yes	Yes
IEC 60601-1 Compliant	Yes	Yes	Yes
Full Scale View Range	±2.5 µV to ± 50 mV	± 0.5µV to ± 8mV	± 0.5µV to ± 5mV
Frequency Response	15 Hz to 1.85 kHz	3 Hz to 4.8 kHz	3 Hz to 1.5 kHz
User Interface	Medtronic supplied computer with keyboard and mouse	NuVasive-supplied computer with optional touch screen and/or keyboard/mouse Mobile device (for NuvaLine only)	NuVasive-supplied computer with separate touch screen and/or keyboard/mouse Mobile device
Remote Access	No	Yes	Yes
Train of Four Testing	N/A	Yes	Yes

Specification/ Property	Predicate Device	Predicate Device	Subject Device
	Medtronic NIM 3.0 Nerve Monitoring System (K083124)	NuVasive NVM5 System (K152942 and K132694)	NuVasive NG-NVM5 System
Needle Electrodes	Various	Various	Various
Surface Electrodes	Various	Various	Various, including new cutaneous electrodes for lumbar MEP
Electrode Leads	Various	Various	Various
Stimulating Probes	Various	Various	Various
Recording Channels	EMG	EMG, MEP, and SSEP	EMG, MEP, and SSEP
EMG			
EMG Modalities	<ul style="list-style-type: none"> Free Run EMG 	<ul style="list-style-type: none"> XLIF (Detection) Basic & Dynamic Screw Test Free Run EMG Twitch Test 	<ul style="list-style-type: none"> XLIF (Detection) Basic & Dynamic Screw Test Free Run EMG Twitch Test
XLIF (Detection)			
Types of Modes	Triggered Stimulation	Automatic Stimulation	Automatic Stimulation
Threshold Values for Color Alerts	No	Yes	Yes (Identical to predicate)
Audio feedback	No	Yes	Yes
Basic & Dynamic Screw Test			
Types of Modes	N/A	Automatic Stimulation	Automatic Stimulation
Threshold Values for Color Alerts	N/A	Yes	Yes (Identical to predicate)
Audio feedback	N/A	Yes	Yes
Free Run EMG			
Types of Modes	Manual Stimulation	Manual Stimulation	Manual Stimulation
Threshold Values for Color Alert	Yes	Yes	Yes (Identical to K152942)
Audio feedback	Yes	Yes	Yes

Specification/ Property	Predicate Device	Predicate Device	Subject Device
	Medtronic NIM 3.0 Nerve Monitoring System (K083124)	NuVasive NVM5 System (K152942 and K132694)	NuVasive NG-NVM5 System
Twitch Test			
Types of Modes	N/A	Manual and Automatic Stimulation	Manual and Automatic Stimulation
Threshold Values for Color Alerts	N/A	Yes	Yes
Audio feedback	N/A	Yes	Yes
MEP			
Types of Modes	N/A	Manual and Automatic Stimulation	Manual and Automatic Stimulation
Threshold Values for Color Alerts	N/A	Yes	Yes (Identical to predicate)
Audio feedback	N/A	Yes	Yes
SSEP			
Types of Modes	N/A	Manual Stimulation	Manual Stimulation
Threshold Values for Color Alerts	N/A	Yes	Yes (Identical to predicate)
Audio feedback	N/A	Yes	Yes
Remote Access			
Screen-sharing accessibility	Remote Monitoring	Remote Monitoring	Remote Monitoring and Local Wireless Control
Guidance			
Clinical Use	N/A	<ul style="list-style-type: none"> • Requires input derived from CT, MRI, or radiographic images • Intended to assist the surgeon in cannulating the pedicle based on user predefined trajectory • Integrated with EMG stimulation 	Not applicable
Performance Requirements	N/A	<ul style="list-style-type: none"> • Angular tolerance of $\pm 2^\circ$ • Confirmation of alignment to pre-planned trajectory • Seamlessly integrated with an insulated Jamshidi Needle 	Not applicable
IEC 60601 Compliant	N/A	YES	YES
User Interface	N/A	Touch screen, graphical user interface and audio	Not applicable

Specification/ Property	Predicate Device	Predicate Device	Subject Device
	Medtronic NIM 3.0 Nerve Monitoring System (K083124)	NuVasive NVM5 System (K152942 and K132694)	NuVasive NG-NVM5 System
Bendini			
Components	N/A	Optical (IR) tracking technology system, IR tracking instruments, computer.	Optical (IR) tracking technology system, IR tracking instruments, computer.
User Interface	N/A	Touch screen, graphical user interface and audio.	Touch screen, graphical user interface and audio.
IEC 60601 Compliant	N/A	YES	YES
Instrumentation	N/A	<ul style="list-style-type: none"> • IR Digitizer (with integrated passive spheres) • Rod Bender • Optional Mobile application 	<ul style="list-style-type: none"> • IR Digitizer (with integrated passive spheres) • Rod Bender • Optional Mobile application

G. Performance Data

Nonclinical testing was performed to demonstrate that the subject *NG-NVM5 System* is substantially equivalent to other predicate devices and to verify that the *NG-NVM5 System* meets design specifications and performance characteristics, based upon the intended use. The *NG-NVM5 System* was subjected to Verification and Validation Testing according to the Software Requirements Specifications defined for the system, to include the modifications made as part of the subject device. Laboratory bench top testing was performed to verify accuracy of angle and offset measurement and validate that the graphical user interface (GUI) and system components function as intended. Additionally, the current and power densities were calculated to support substantial equivalence of cutaneous electrodes.

The results of these studies showed that the subject *NG-NVM5® System* meets the same acceptance criteria as the performance of the predicate device, and the device was therefore found to be substantially equivalent.

H. Conclusions

Based on the indications for use, technological characteristics, performance testing, and comparison to predicate devices, the subject *NG-NVM5 System* has been shown to be substantially equivalent to legally marketed predicate devices.