



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

July 1, 2015

Orthomerica Products, Inc.
David Hooper
Manufacturing Engineer
6333 North Orange Blossom Trail
Orlando, Florida 32810

Re: K151147
Trade/Device Name: Starband® and Starlight®
Regulation Number: 21 CFR 882.5970
Regulation Name: Cranial Orthosis
Regulatory Class: Class II
Product Code: OAN, MVA
Dated: April 27, 2015
Received: May 5, 2015

Dear David Hooper:

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 yours,

Carlos L. Pena -S 

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)

K151147

Device Name

STARband® and STARlight®

Indications for Use (Describe)

The STARband and STARlight are intended for medical purposes for use on infants from 3 to 18 months of age, with moderate-to-severe non-synostotic positional plagiocephaly, including infants with plagiocephalic-, brachycephalic- and scaphocephalic-shaped heads by applying mild pressure to prominent regions of the infant's cranium in order to improve cranial symmetry and/or shape. These devices are also indicated for adjunctive use for infants from 3 to 18 months of age whose synostosis has been surgically corrected, but who still have moderate-to-severe cranial deformities including plagiocephalic-, brachycephalic- and scaphocephalic-shaped heads.

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.

This section applies only to requirements of the Paperwork Reduction Act of 1995.

DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.

The burden time for this collection of information is estimated to average 79 hours per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to:

Department of Health and Human Services
Food and Drug Administration
Office of Chief Information Officer
Paperwork Reduction Act (PRA) Staff
PRASStaff@fda.hhs.gov

"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number."

510(k) Summary

I. Applicant Information

Name: Orthomerica Products, Inc.
Address: 6333 North Orange Blossom Trail
Orlando, FL 32810
Telephone: (407) 290-6592
Facsimile: (407) 290-2419

FDA Establishment Registration Number

1058152

Contact Information

Contact Person: David Hooper, Manufacturing Engineer
Address: 6333 North Orange Blossom Trail
Orlando, FL 32810
Telephone: (407) 290-6592
Facsimile: (407) 290-2419
Email: dhooper@orthomerica.com
Date Prepared: April 27th, 2015

II. Submission Information

Type: Traditional 510(k) Submission
Proprietary Name: STARband® and STARlight®
Common Name: Cranial Orthosis
Classification: Class II (special controls); OAN; MVA; 21 CFR 882.5970
Classification Name: Cranial Orthosis

III. Manufacturer Site

Name: Orthomerica Products, Inc.
Address: 6333 North Orange Blossom Trail
Orlando, FL 32810
Telephone: (407) 290-6592
Facsimile: (407) 290-2419
FDA Establishment Registration Number: 1058152

IV. Description of Device/Modification

The STARband and STARlight redirects the head growth to improve proportion and symmetry. The practitioner takes a plaster impression or 3-dimensional captured image of the infant's head to acquire the existing shape. The mold is sealed and filled with plaster or the 3-dimensional image is carved from a rigid polyurethane foam blank to create a positive model of the head shape. The positive model is modified to obtain greater symmetry and space in the areas of flattening. The STARband and STARlight provide total contact over the prominent or bossed areas of the baby's head to discourage growth there. Over the course of treatment, the inside of the band is further modified by the practitioner to provide space for growth to occur in the flat or depressed areas. The shape of the STARband and STARlight directs growth into the areas of least resistance and creates a precise pathway for the head shape to improve in symmetry and proportion.

The STARband and STARlight product families as it was released in K141842 are essentially still the same devices. The STARband Side Opening design and STARband Bi-Valve design is made with an outer shell of 5/32" polyethylene-polypropylene copolymer plastic with an inner liner made of 1/2" pelite polyethylene foam or 1/2" Aliplast foam (closed cell polyethylene). The STARlight Side Opening design and the STARlight Bi-Valve design are made of a plastic shell of 5/32" – 1/4" clear Surlyn or 1/8" - 7/32" Clear Co-Polyester. The STARlight PRO (Post-operative Remolding Orthosis) design is made of 1/4" to 3/8" clear Surlyn. Optional Aliplast (closed cell polyethylene) padding is available for the clear plastic bands and in addition, optional Reston (polyurethane – 3M Medical Product) foam is available for the STARlight PRO design.

The STARband Side Opening design and the STARlight Side Opening design has a top opening and a side opening. The band is held in place by a Velcro® strap (1½" for STARband Side Opening and 1" for STARlight Side Opening) across the side opening. The STARlight PRO has two side openings, no top opening, and is held in place by a Velcro strap across each side opening. The STARlight Bi-Valve design and the STARband Bi-Valve design consist of two plastic shells that overlap with a superior sliding mechanism. The right and left overlap tabs are connected via a Velcro strap with chafe and loop.

The proposed device modification is the addition of a new 3-dimensional shape capture method, specifically, the Rodin4D M4DScan System and the BodyScan System. The BodyScan System is the same as the M4DScan System (Hardware and Software included) except it is distributed in the United States by TechMed 3D, Inc. on behalf of Rodin4D. This system uses a structured white light projector and a single camera to detect the shape of the surface captured. This system utilizes LED white light which is a

non-coherent (i.e. non-laser) light source. Because this system utilizes a non-coherent light source, it is safe to use on infant patients under all circumstances.

V. Statement of Indications and Intended Use

Statement of Indications:

The STARband and STARlight are intended for medical purposes for use on infants from 3 to 18 months of age, with moderate-to-severe non-synostotic positional plagiocephaly, including infants with plagiocephalic-, brachycephalic- and scaphocephalic-shaped heads by applying mild pressure to prominent regions of the infant's cranium in order to improve cranial symmetry and/or shape. The devices are also indicated for adjunctive use for infants from 3 to 18 months of age whose synostosis has been surgically corrected, but who still have moderate-to-severe cranial deformities including plagiocephalic-, brachycephalic- and scaphocephalic-shaped heads.

Intended Use:

The STARband and STARlight are designed to treat infants with abnormal head shapes from age 3 to 18 months and is available by prescription only. Since growth is the driving factor in head shape correction, the infants wear the STARband or STARlight for approximately 23 hours per day. The most common head deformities are positional plagiocephaly, brachycephaly, and scaphocephaly. The STARband and STARlight have also been cleared to treat unresolved head deformities in patients who have undergone surgery to correct craniosynostosis. The same principles of cranial remolding apply to positional deformities and post-operative patients.

VI. Predicate Devices

- STARband and STARlight, Cranial Orthosis, K141842

VII. Summary of Technological Characteristics

The M4DScan/BodyScan System proposed in this 510(k) is an additional method to capture the infant’s head shape for the fabrication of the STARband and STARlight Cranial Orthosis. The technological characteristics and the underlying principles of operation of the STARband and STARlight Cranial Orthosis shall remain exactly the same. The inclusion of the M4DScan /BodyScan System is the focus of this submission and that change is indicated in **Table 1** under the Approved 3-Dimensional Imaging Devices section.

Table 1 – Comparison of Predicate Device cleared in K141842 to the Proposed Device

Feature	From K141842	Proposed Device
Intended Use	Maintains total contact over areas of bossing or protrusion and creates voids over areas of depression or flattening to redirect cranial growth toward greater symmetry.	Maintains total contact over areas of bossing or protrusion and creates voids over areas of depression or flattening to redirect cranial growth toward greater symmetry.
Materials	<p>Material for STARband Side Opening design and STARband Bi-Valve design</p> <ul style="list-style-type: none"> - Outer shell of 5/32” copolymer plastic - An inner liner of 1/2” Pelite polyethylene foam or 1/2” Aliplast foam <p>Material for STARlight Side Opening design and STARlight Bi-Valve design</p> <ul style="list-style-type: none"> - 5/32” - 1/4” clear Surlyn or 1/8” – 7/32” Clear Co-Polyester plastic shell <p>Material for STARlight PRO design</p> <ul style="list-style-type: none"> - 1/4” – 3/8” clear Surlyn <p>Closure for Bivalve design</p> <ul style="list-style-type: none"> - Sliding/Overlap closure system - Chicago screw (or similar) for top sliding mechanism - 1” Velcro strap - 1” chafe buckle - Speedy rivets <p>Closure for STARband Side Opening design</p> <ul style="list-style-type: none"> - 1 ½” Velcro Strap - 1 ½” chafe buckle - A Gap Block made from ½” firm Pelite polyethylene foam 	<p>Material for STARband Side Opening design and STARband Bi-Valve design</p> <ul style="list-style-type: none"> - Outer shell of 5/32” copolymer plastic - An inner liner of 1/2” Pelite polyethylene foam or 1/2” Aliplast foam <p>Material for STARlight Side Opening design and STARlight Bi-Valve design</p> <ul style="list-style-type: none"> - 5/32” - 1/4” clear Surlyn or 1/8” – 7/32” Clear Co-Polyester plastic shell <p>Material for STARlight PRO design</p> <ul style="list-style-type: none"> - 1/4” – 3/8” clear Surlyn <p>Closure for Bivalve design</p> <ul style="list-style-type: none"> - Sliding/Overlap closure system - Chicago screw (or similar) for top sliding mechanism - 1” Velcro strap - 1” chafe buckle - Speedy rivets <p>Closure for STARband Side Opening design</p> <ul style="list-style-type: none"> - 1 ½” Velcro Strap - 1 ½” chafe buckle - A Gap Block made from ½” firm Pelite polyethylene foam

Feature	From K141842	Proposed Device
	<ul style="list-style-type: none"> - Large Flange, Blind Rivet <p>Closure for STARlight Side Opening design and the STARlight PRO design:</p> <ul style="list-style-type: none"> - 1” Velcro Strap - 1” chafe buckle Optional tamper resistant strap (qty 2 for the STARlight PRO design) 	<ul style="list-style-type: none"> - Large Flange, Blind Rivet <p>Closure for STARlight Side Opening design and the STARlight PRO design:</p> <ul style="list-style-type: none"> - 1” Velcro Strap - 1” chafe buckle Optional tamper resistant strap (qty 2 for the STARlight PRO design)
Product Design	Custom made cranial orthosis, approximately 6 to 10oz in weight. STARlight PRO weighs 12.5 to 18.5 oz.	Custom made cranial orthosis, approximately 6 to 10oz in weight. STARlight PRO weighs 12.5 to 18.5 oz.
Production	<ul style="list-style-type: none"> - Form orthosis from a positive mold of infant’s head - Positive mold is formed based upon measurements of the infant’s head taken by an approved 3-dimensional imaging device from which a 3-dimensional image is made or from a traditional plaster cast - The 3-dimensional image is used to produce a positive mold using a 5-axis routing machine 	<ul style="list-style-type: none"> - Form orthosis from a positive mold of infant’s head - Positive mold is formed based upon measurements of the infant’s head taken by an approved 3-dimensional imaging device from which a 3-dimensional image is made or from a traditional plaster cast - The 3-dimensional image is used to produce a positive mold using a 5-axis routing machine
Approved 3-Dimensional Imaging Devices	<ul style="list-style-type: none"> - STARscanner I - STARscanner II - Omega Scanner - scanGogh-II - 3dMDhead System - 3dMDcranial System - 3dMDflex System - SmartSoc System 	<ul style="list-style-type: none"> - STARscanner I - STARscanner II - Omega Scanner - scanGogh-II - 3dMDhead System - 3dMDcranial System - 3dMDflex System - SmartSoc System - M4DScan/BodyScan System
Testing	<p>Repeatability and Reproducibility (R&R) Analysis</p> <ul style="list-style-type: none"> - Utilized uniform shapes with known dimensions that represent various sizes of pediatric patients between ages 3 to 18 months of age - Compared proposed device to cast and predicate device - Associated parameters includes A-P and M-L - Proposed device is substantially equivalent to predicate device <p>Cranial Shape Capture Accuracy Study</p> <ul style="list-style-type: none"> - Utilized a representative cranial 	<p>Repeatability and Reproducibility (R&R) Analysis</p> <ul style="list-style-type: none"> - Utilized uniform shapes with known dimensions that represent various sizes of pediatric patients between ages 3 to 18 months of age - Compared proposed device to cast and predicate device - Associated parameters includes A-P and M-L - Proposed device is substantially equivalent to predicate device <p>Cranial Shape Capture Accuracy Study</p> <ul style="list-style-type: none"> - Utilized a representative cranial

Feature	From K141842	Proposed Device
	<p>shape that possesses a predefined shape with known dimensions</p> <ul style="list-style-type: none"> - Compared proposed device to cast and predicate device - Associated Coordinate Planes (A-P; M-L; P-D and various Radius Parameters; Squareness; Flatness) - Proposed device is substantially equivalent to predicate device <p>Material Biocompatibility Testing</p> <ul style="list-style-type: none"> - Cytotoxicity –Agar Diffusion - Closed Patch Sensitization - Primary Dermal Irritation 	<p>shape that possesses a predefined shape with known dimensions</p> <ul style="list-style-type: none"> - Compared proposed device to cast and predicate device - Associated Coordinate Planes (A-P; M-L; P-D and various Radius Parameters; Squareness; Flatness) - Proposed device is substantially equivalent to predicate device <p>Material Biocompatibility Testing</p> <ul style="list-style-type: none"> - Cytotoxicity –Agar Diffusion - Closed Patch Sensitization - Primary Dermal Irritation

The STARband and STARlight Cranial Orthosis have already received FDA 510(k) clearance under K141842 for being manufactured from a 3-dimensional imaging device that utilizes a non-coherent light source and takes 2-dimensional (2D) images for shape capture (3dMD Systems and SmartSoc System). The M4DScan/BodyScan System utilizes the same technology as the FDA 510(k) cleared 3dMD and SmartSoc Systems of a non-coherent light source and takes continuous 2D images from triangulated positions for shape capture. Considering that STARband and STARlight are still the same device as it was in the predicate device and that the shape capture devices have the same technological characteristics; the STARband and STARlight Cranial Orthosis are substantially equivalent to the predicate device.

Table 2 – Performance Testing Summary for STARband and STARlight Cranial Orthosis

Test	Test Method Summary	Results
Repeatability and Reproducibility (R&R) Analysis	<p>The purpose of this test is to analyze the repeatability and reproducibility (components of precision) of the proposed shape capture device. The data obtained is compared to the casting process and the predicate device.</p> <p>Utilized uniform shapes with known dimensions that represent various sizes of pediatric patients between ages 3 to 18 months of age. Associated test parameters include A-P and M-L.</p>	<p>The shape capture device passed the acceptance criteria. The relevancy of the test determined substantial equivalence of the proposed shape capture device.</p>
Cranial Shape Capture Accuracy Study	<p>The purpose of this test is to analyze the accuracy of the proposed shape capture device. The data obtained is compared to the casting process and the predicate device.</p> <p>Utilized a representative cranial shape that possesses a predefined shape with known dimensions. Associated parameters analyzed include A-P, M-L, P-D, various radius parameters, squareness, and flatness.</p>	<p>The shape capture device passed the acceptance criteria for all parameters. The relevancy of the test determined substantial equivalence of the proposed shape capture device.</p>

The STARband and STARlight are essentially the same Cranial Orthosis. The main difference between the STARband and STARlight are the materials used to produce them. The STARband and STARlight materials have been biocompatibility tested, and the results of the tests are listed below in **Table 3**.

Table 3 – Biocompatibility Testing Summary for STARband and STARlight Cranial Orthosis

Material	Test	Results	Conclusion
Surlyn	Closed Patch Sensitization	A score of 0.00/0.00 (Test/Control) was given for both Incidence and Severity in the 24 hour and 48 hour scoring interval.	Not a Sensitizer No Erythema or Edema Formation
Surlyn	Primary Dermal Irritation	Primary Irritation Index: 0.00	Negligible Dermal Response
Surlyn	Cytotoxicity – Agar Diffusion	Cell culture treated with test sample exhibited no reactivity (Grade 0).	Non-cytotoxic
Copolymer with Pelite Foam	Closed Patch Sensitization	A score of 0.00/0.00 (Test/Control) was given for both Incidence and Severity in the 24 hour and 48 hour scoring interval.	Not a Sensitizer No Erythema or Edema Formation
Copolymer with Pelite Foam	Primary Dermal Irritation	Primary Irritation Index: 0.06	Negligible Dermal Response
Copolymer with Pelite Foam	Cytotoxicity – Agar Diffusion	Cell culture treated with test sample exhibited no reactivity (Grade 0).	Non-cytotoxic
Copolymer with Aliplast Foam	Closed Patch Sensitization	A score of 0.00/0.00 (Test/Control) was given for both Incidence and Severity in the 24 hour and 48 hour scoring interval.	Not a Sensitizer No Erythema or Edema Formation
Copolymer with Aliplast Foam	Primary Dermal Irritation	Primary Irritation Index: 0.00	Negligible Dermal Response
Copolymer with Aliplast Foam	Cytotoxicity – Agar Diffusion	Cell culture treated with test sample exhibited slight reactivity (Grade 1).	Non-cytotoxic

VIII. Summary and Conclusions of Non-Clinical Performance Data

The M4DScan/BodyScan System was evaluated for substantial equivalence. The system uses flashes of structured light which is equivalent to the flash from a consumer grade camera and is safe to use on infants without eye protection. The shape capture repeatability and reproducibility was evaluated and determined to be acceptable. An additional, Cranial Shape Capture Accuracy Study was performed concluding that the M4DScan/BodyScan System yields a product that is substantially equivalent to the predicate device. With sufficient accuracy and no concerns with the safety of the system, the M4DScan/BodyScan System was determined to have a safety and effectiveness profile similar to the predicate device for capturing infant head shape data to manufacture the STARband and STARlight Cranial Orthosis.