DePuy Synthes Cranio-maxillofacial Distraction System - May Reverse Directions After Surgery

Recall Class: Class I

Date Recall Initiated: April 16, 2014


Manufacturing Dates: April 20, 2009 through April 15, 2011

Distribution Dates: November 3, 2009 to April 14, 2014


Use: The DePuy Synthes Cranio-maxillofacial (CMF) Distraction System is an implant used to lengthen and/or stabilize the lower jawbone (mandibular body) and the side of the lower jaw (ramus). This device is used in pediatric and adult patients to correct birth (congenital) or post-traumatic defects of the jaw by gradually lengthening the bone (distraction).

Recalling Firm:
Synthes (USA)
1302 Wrights Lane East
West Chester, PA 19380

Reason for Recall: DePuy Synthes is recalling certain lots of the Cranio-maxillofacial Distraction System because the device may reverse direction and lose the desired distraction distance after surgery.

- Infants are at the highest risk for injury if the device fails because sudden obstruction of the trachea can occur. This could lead to respiratory arrest, and result in death.
- Children or adults with the ability to maintain an open airway are at less risk for serious injury because failure of the device would not result in tracheal obstruction and could be medically reversible.
- In all patient populations, failure of the device may result in the need for surgical intervention to replace the failed device.

There have been 15 reports of injury associated with the use of this device.

http://www.fda.gov/MedicalDevices/Safety/ListofRecalls/ucm411589.htm?source=govd... 9/1/2014
Public Contact: Customers who have questions about this recall may contact DePuy Synthes Customer Support at 1-800-479-6328, Monday – Friday, 9:00 a.m. to 8:00 p.m. Eastern Time or the sales consultant.

FDA District: Philadelphia District Office

More Information about this Recall:

On April 16, 2014, DePuy Synthes sent an Urgent Notice (http://www.depuysynthes.com/binary/org/DPY_SYN/Products/Images/CMF/Recall%202013%20CNL%20CMF%20Distraction%20System%20(AB%20BC%20Distractor%20Bodies)%20Customer%20Notification%20Final%204-15-2014.pdf) to their customers. The notice identified the problem, affected products and tells customer to:

- Review their inventory and remove affected lots from stock.
- Call DePuy Synthes at 1-800-479-6329 for a return authorization number
- Complete and return the verification in the letter included with the notice.

About Class I Recalls

Class I recalls are the most serious type of recall and involve situations in which there is a reasonable probability that use of these products will cause serious adverse health consequences or death.

Health care professionals and consumers may report adverse reactions or quality problems they experienced using these products to MedWatch: The FDA Safety Information and Adverse Event Reporting Program (https://www.accessdata.fda.gov/scripts/medwatch/) either online, by regular mail or by FAX.

Additional Resources

### Class 1 Device Recall DePuy Synthes Craniofacial Distraction System

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#### Class 1 Device Recall DePuy Synthes Craniofacial Distraction System

#### Date Posted
August 20, 2014

#### Recall Status
Open

#### Recall Number
Z-2148-2014

#### Recall Event ID
6807123

#### Premarket Notification 510(K) Number
K05013824

#### Product Classification
External Mandibular Fixator And/OR Distractor - Product Code MQN26

#### Product
DePuy Synthes Craniofacial Distraction System (AB Distractor Bodies and BC Distractor Bodies)

#### Code Information
Part nos. 04.315.003, 04.315.004, 04.315.005, 04.315.006 04.315.023 04.315.024 04.315.025 04.315.026 04.315.027 04.315.028 04.315.029 04.315.053 04.315.054 04.315.055 04.315.056 04.315.063 04.315.064 04.315.065 04.315.066 04.315.067 04.315.068, with lot nos. 6245205, 7129328, IS10400 6246203, 6694298, 6914674, 6977377, 6921359, 6981567, 7008430, 7081952, 7129329, 7460419, 7556184 6246984, 6342006, 6964974, 7129330, 7460420, 6246983, 7389334, 7422735, 7515818 6245460, 6342005, 6454138, 6945399, 6651393, 6816023, 6898959, 6921360, 7306561, 7515824 6370904, 6356443, 6410544, 6438375, 6625359, 6651392, 6894928, 7241620, 7351855, 7394070, 7408179, 7458272, 7458273, 7458274, IS10394 6450702, 6454139, 6553925, 6625840, 6883404, 6921586, 7031344, 7185897, 7399329, 7408181, 7422707, 7422708, 7422706, 7458299, 7515832, IS10365 6393118, 6393119, 6410545, 6454140, 6512859, 6625641, 6651390, 6684925, 7185895, 7408182, 7422703, 7422704, 7515829, 7515830, 6393120, 7408182, 7515833 7229138, 7399333, 7422700, 7545381, IS10366 6252759, 7129331, 7155681, 7418771, 7448753 6245463, 6438376, 6981568, 6983876, 7003685, 7129332, 7309797, 7185772, 7418848, 7448754 6251769, 7081953, 7310030, 7365542, 7418849, 7448752, 7448758, 7556149 6245458, 6342007, 7129333, 7422709, IS10395 625462, 6342003, 624546, 6245451, 6883290, 6983301, 7082907, 7269672, 7477692, 7475810, 7557623 6280719, 6342004, 6394297, 6651930, 673168, 6757617, 682454, 6824550, 6883304, 6883305, 6961025, 6984927, 7011382, 7056501, 7111867, 7118593, 7269687, 7351857, 7351858, 7351859, 7515861, 7422701, 7545379, 7579520, IS10395, 6651389, 6702172, 6883306, 6883307, 6942444, 6984926, 7111868, 7185692, 7306562, 7306563, 7306564, 7498975, 7515834, 7515826, IS10369 6651388, 6741835, 6824553, 6883309, 6913821, 6942445, 6961024, 7082908, 7111869, 7129252, 7158888, 7515858, 7515863, 7657633 6245185, 6942446, 7185887, 7306565, 7306566, 7408183, 7477691, 7515866, 7579433, 7565635, IS10390, IS10391, 6275129, 7082005, 7185890, 7269699, 7422697, 7509200, IS10386, IS10387

#### Recalling Firm/Manufacturer
Synthes, Inc.
1302 Wrights Ln E
West Chester, Pennsylvania 19380-3417

#### For Additional Information Contact
Customer Support
610-719-5000

#### Manufacturer Reason for Recall
DePuy Synthes is initiating a recall of certain lots of the Craniofacial Distraction System (AB Distractor Bodies and BC Distractor Bodies) because they may reverse post-operatively.

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http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfres/cfres.cfm?id=126898

9/1/2014
Class 2 Device Recall Synthes Cranio maxillofacial (CMF) Distraction System

Date Posted: July 12, 2013
Recall Status: Open
Recall Number: Z-1726-2013
Recall Event ID: 6567023
Premarket Notification 510(K) Number: K06013824
Product Classification: External Mandibular Fixator And/Or Distractor - Product Code MGN26

Product:
Synthes CMF Distraction System The product is intended for use as a bone stabilizer and lengthening (and/or transport) device for correction of congenital deficiencies or post-traumatic defects of the mandibulat body and ramus, where gradual bone distraction is required. It is intended for single use only.

Code Information:
Part # 04.315.067 with lot #s 6184270 and/or 22612-04.

Recalling Firm/Manufacturer:
Synthes USA HQ, Inc.
1302 Wight Ln E
West Chester, Pennsylvania 19380-3417

Consumer Instructions:
Contact the recalling firm for information.

For Additional Information Contact:
Synthes Recall Information Center
610-719-5450

Manufacturer Reason for Recall:
The firm initiated a voluntary recall of the BC Distractor Body which is part of the Cranio maxillofacial (CMF) Distractor System, due to a mis-alignment issue. There have been instances reported within the impacted lots in which the slot in the barrel of the BC distractor was rotated such that it was not possible to attach the B-type and C-type foot plates.

FDA Determined Cause:
COMPONENT CONTROLS (GMP - GOOD MANUFACTURING PRACTICE): Nonconforming Material/Component

Action:
SYNTHESES sent an Urgent Notice: Medical Device Recall letter dated June 12, 2013, to all affected customers. The letter identified the product, the problem, and the action to be taken by the customer. Customers were requested to examine their inventory for affected product, remove it from use, and return it to Synthes. If they have affected product they were instructed to call Synthes at 1-800-479-6329 to obtain a Return Authorization Number. Complete the Verification Section at the end of the letter and indicate the number of devices found and note the Return Authorization Number and return to Credit/Returns, Syntheses, 1101 Syntheses Avenue, Monument, CO 80132. If customers did not have the affected product they were asked to complete the Verification Section and fax it to 610-251-9005. Customers with questions were instructed to call 610-719-5450 or email FieldAction@synthes.com. For questions regarding this recall call 610-719-5450.

Quantity in Commerce: 10
Distribution: Nationwide Distribution including PA, CT, DE, and NY.
Total Product Life Cycle: TPLC Device Report27

9/1/2014