SYMBOL GLOSSARY

American Orthodontics utilizes symbols that are in conformance to EN 980 as listed in the European Harmonized Standards list; ISO 15223-1 and ISO 7010 as listed in the US FDA's Consensus Standards. Other symbols deemed necessary, but not on the harmonized/consensus list are also found below. Symbols will appear on packaging/labeling and instructions for use where applicable.



Manufacturer

FDA Consensus Standard ISO 15223-1 REF # 5.1.1 EU Harmonized Standard BS EN 980 REF # 5.12 Indicates the medical device manufacturer, as defined in EU Directives 90/385-EEC, 93/42/EEC



Authorized Representative

in the European Community FDA Consensus Standard ISO 15223-1 REF # 5.1.2 EU Harmonized Standard BS EN 980 REF # 5.13 Indicates the Authorized representative in the European Community



CE Marking

Complies with European directives



1434 EU Notified Body Number Complies with European directives



Catalogue Number

FDA Consensus Standard ISO 15223-1 REF # 5.1.6 EU Harmonized Standard BS EN 980 REF # 5.1 Indicates the manufacturer's catalogue number so that the medical device can be identified



Batch Code

FDA Consensus Standard ISO 15223-1 REF # 5.1.4 EU Harmonized Standard BS EN 980 REF # 5.4 Indicates the manufacturer's batch code so that the batch or lot can be identified



Serial Number

FDA Consensus Standard ISO 15223-1 REF # 5.1.7 EU Harmonized Standard BS EN 980 REF # 5.5 Indicates the manufacturer's serial number so that a specific medical device can be identified



Quantity

Indicates the amount of product included



Use By Date

FDA Consensus Standard ISO 15223-1 REF # 5.1.4 EU Harmonized Standard BS EN 980 REF # 5.3 Indicates the date after which the medical device



Do Not Re-use

FDA Consensus Standard ISO 15223-1 REF # 5.4.2 EU Harmonized Standard BS EN 980 REF # 5.2 Indicates a medical device that is intended for one use, or for use on a single-patient dureing a single



Caution

FDA Consensus Standard ISO 15223-1 REF # 5.4.4 EU Harmonized Standard BS EN 980 REF # 5.11 Indicates the need for the user to consult the instructions for use for important cautionary information such as warnings and precautions that cannot, for a variety of reasons, be presented on the medical device itself



Keep away from sunlight FDA Consensus Standard ISO 15223-1 REF # 5.3.2 EU Harmonized Standard BS EN 980 REF # 5.20 Indicates a medical device that needs protection from light sources



Temperature Limit

FDA Consensus Standard ISO 15223-1 REF # 5 3 7 EU Harmonized Standard BS EN 980 REF # 5.17.3 Indicates the temperature limits to which the medical device can be safely exposed



Humidity Limitation

FDA Consensus Standard ISO 15223-1 REF # 5.3.8 Indicates the range of humidity to which the medical device can be safely exposed



Consult Instructions for Use

FDA Consensus Standard ISO 15223-1 REF # 5.4.3 EU Harmonized Standard BS EN 980 REF # 5.18 Indicates the need for the user to consult the instructions for use



Non-Sterile

FDA Consensus Standard ISO 15223-1 REF # 5.2.7 EU Harmonized Standard BS EN 980 REF # 5.23 Indicates a medical device that has not been subjected to a sterilization process



Contains or Presence of Natural Rubber Latex

FDA Consensus Standard ISO 15223-1 REF # 5.4.5 EU Harmonized Standard BS EN 980 REF # 6.2 Indicates the presence of natural rubber latex or dry natural rubber latex as a material of construction within the medical deivce or the packaging of the medical device



Product is not made with Natural Rubber Latex

Product is not made with or contains natural

Statement "Caution: Federal Law restricts this device to sale to or on the order of a dentist/orthodontist" 21CFR801.109(b)



Danger or Warning (EC) No 1272/2008 [CLP] REF # GHS05

Corrosive cat 1 Indicates product may cause corrosive damage to metals, as well as skin, eyes



Danger

(EC) No 1272/2008 [CLP] REF # GHS06 Toxic cat. 1-3 Indicates product can cause death or toxicity with short exposure to small amounts



Danger or Warning

(EC) No 1272/2008 [CLP] REF # GHS02 Flammable Indicates fire hazard



(EC) No 1272/2008 [CLP] REF # GHS07 Toxic cat. 4 Irritant cat. 2 or 3

Lower systematic health hazards Indicates product may cause less serious health effects or damage ozone layer



Nickel-Chromium Warning

Indicates product contains Nickel and/or Chromium. Patients with an identified allergy to these metals should not use this product



Magnetic Field

Indicates interaction with metallic objects may product Pinch Hazards



FDA Consensus Standard ISO 7010 REF # P007 Indicates product can be harmful to pacemaker



Health Hazard GHS08 WHMIS 2015

Indicates product may cause or suspected of causing serious health effects



Not Sterilized

Indicates product is not sterilized by the