

Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Surgical instrument handle

ARTG Identifier 221929

ARTG Start date 2/04/2014

Product Category Medical Device Included Class 1

GMDN 47829

GMDN Term Surgical instrument handle

Intended Purpose This is a hand-held manual surgical instrument designed to attach to the

proximal end of a surgical instrument to allow the surgeon to perform manipulations with the instrument, typically manual rotation of a bone

screw or tensioning of a nut during a surgical procedure.

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	

ARTG Standard Conditions

- The automatic conditions applicable to the inclusion of all kinds of medical devices in the Register are as specified in section 41FN of the Therapeutic Goods Act 1989.
- The standard conditions that are imposed under section 41FO of the Therapeutic Goods Act 1989 when kinds of medical devices are included in the Register are as set out in the following paragraphs.
- For a medical device included in the Register under Chapter 4 and imported into Australia, the Sponsor must ensure that information about the Sponsor is provided in such a way as to allow the sponsor to be identified.
- Each sponsor shall retain records of the distribution of all of the sponsor's medical devices included in the Register under Chapter 4. In the case of records relating to a Class AIMD medical device, Class III medical device, or Class IIb medical device that is an implantable medical device, the distribution records shall be retained for a minimum period of 10 years. In the case of records relating to any other device, the distribution records shall be retained for a minimum period of 5 years.
- The sponsor of a medical device included in the Register under Chapter 4 shall keep an up to date log
 of information of the kind specified in Regulation 5.8.
- It is a condition of inclusion in the ARTG that the sponsor of a medical device that is an AIMD, Class III or implantable Class IIb provides three consecutive annual reports to the Head of the Office of Product Review, Therapeutic Goods Administration following inclusion of the device in the ARTG (as specified in 5.8 of the regulations). Annual reports are due on 1 October each year. Reports should be for the period 1 July to 30 June. The first report following the date of inclusion in the ARTG must be for a period of at least six months but no longer than 18 months. Subsequent reports are to be provided on 1 October for a further 2 years. The annual report must include all complaints and adverse events received by the manufacturer relating to problems with the use of the device that have been received by them over the year. For orthopaedic implant prosthesis that have been re-classified from Class IIb to Class III medical devices, annual report information must be submitted if the device meets either of the following criteria: I.The device was subject to a TGA application audit based on revision rate when the device transitioned from Class IIb to Class III; and/or II.No devices were supplied to the Australian marketplace before 30 June 2012. As per the standard automatic condition, annual reports should be submitted each year for the first three years of inclusion as a Class III medical device on the ARTG.
- · Where a medical device included in the Register, contains a substance which is included in the Fourth

Schedule to the Customs (Prohibited Imports) Regulations or the Eighth Schedule to the Customs (Prohibited Exports) Regulations the Sponsor shall, at the time of importation or exportation of the medical device, be in possession of a licence and a permission for importation or exportation of each consignment of the goods as required by those regulations.

• A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Products Covered by This Entry

1. Surgical instrument handle

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 221929 ARTG Start Date: 2/04/2014



Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Drill bit, surgical

ARTG Identifier 221931

ARTG Start date 2/04/2014

Medical Device Included Class 1 **Product Category**

GMDN 32390

GMDN Term Drill bit, surgical

Intended Purpose A device that is opperated manualually that, when rotated will cut into

bone creating a hole of the same dimension as the diameter of the bit.

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	

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medical device, be in possession of a licence and a permission for importation or exportation of each consignment of the goods as required by those regulations.

• A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Products Covered by This Entry

1. Drill bit, surgical

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 221931 ARTG Start Date: 2/04/2014



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Department of HealthTherapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Punch, dental, soft-tissue, reusable

ARTG Identifier 221934
ARTG Start date 2/04/2014

Product Category Medical Device Included Class 1

GMDN 45329

GMDN Term Punch, dental, soft-tissue, reusable

Intended Purpose This is a hand held dental instrument that is intended to be used to

remove with precision a circular area of gum tissue during dental

surgery.

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	

ARTG Standard Conditions

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• A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Products Covered by This Entry

1. Punch, dental, soft-tissue, reusable

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 221934 ARTG Start Date: 2/04/2014



Australian Government

Department of Health

Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Trephine, <specify>

ARTG Identifier 221935

ARTG Start date 2/04/2014

Medical Device Included Class 1 **Product Category**

GMDN 14146

GMDN Term Trephine, <specify>

Intended Purpose This is a hand held cylindrical or crown saw used for the removal of a

disc of bone or other firm tissue during dental procedures in the mouth.

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	

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• A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Products Covered by This Entry

1. Trephine, <specify>

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 221935 ARTG Start Date: 2/04/2014



Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Driver, <specify>

 ARTG Identifier
 221930

 ARTG Start date
 2/04/2014

Product Category Medical Device Included Class 1

GMDN 16868

GMDN Term Driver, <specify>

Intended Purpose This is a hand held metal instrument designed to impart force on

another instrument. The distal end is shaped to mate with the instrument being driven into some form of tissue. The proximal end is designed to

absorb and transmit an impact force.

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	

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Products Covered by This Entry

1. Driver, <specify>

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 221930 ARTG Start Date: 2/04/2014



Australian Governmen

Department of HealthTherapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Wrench, surgical

ARTG Identifier 221932

ARTG Start date 2/04/2014

Product Category Medical Device Included Class 1

GMDN 32871

GMDN Term Wrench, surgical

Intended Purpose This is a hand held device that is a surgical instrument, manually

operated, with fixed jaws for gripping, turning, or twisting an object,

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	

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medical device, be in possession of a licence and a permission for importation or exportation of each consignment of the goods as required by those regulations.

• A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Products Covered by This Entry

1. Wrench, surgical

Product Specific Conditions

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Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

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Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Osteotome set, dental

ARTG Identifier 221936

ARTG Start date 2/04/2014

Product Category Medical Device Included Class 1

GMDN 44886

GMDN Term Osteotome set, dental

Intended Purpose This is a hand held dental surgical osteotomes inteded to be used sed

during dental osteotomy for shaping and condensing bone.

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	

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Products Covered by This Entry

1. Osteotome set, dental

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 221936 ARTG Start Date: 2/04/2014



Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Surgical mallet head

ARTG Identifier 221933

ARTG Start date 2/04/2014

Product Category Medical Device Included Class 1

GMDN 46976

GMDN Term Surgical mallet head

Intended Purpose This is a hand held device that is inteded to be used to replace the head

of some types of surgical mallets and hammers and used during dental

procedures in the mouth.

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Products Covered by This Entry

1. Surgical mallet head

Product Specific Conditions

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Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 221933 ARTG Start Date: 2/04/2014



Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Dental precision attachment, ball

ARTG Identifier 222929

ARTG Start date 30/04/2014

Product Category Medical Device Included Class IIa

GMDN 38576

GMDN Term Dental precision attachment, ball

Intended Purpose This is a device used during dental implant procedures that has an

attachment with ball-formed male part and ring-formed female part.

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	DV-2014-MC-05062-1

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- It is a condition of inclusion in the ARTG that the sponsor of a medical device that is an AIMD, Class III or implantable Class IIb provides three consecutive annual reports to the Head of the Office of Product Review, Therapeutic Goods Administration following inclusion of the device in the ARTG (as specified in 5.8 of the regulations). Annual reports are due on 1 October each year. Reports should be for the period 1 July to 30 June. The first report following the date of inclusion in the ARTG must be for a period of at least six months but no longer than 18 months. Subsequent reports are to be provided on 1 October for a further 2 years. The annual report must include all complaints and adverse events received by the manufacturer relating to problems with the use of the device that have been received by them over the year. For orthopaedic implant prosthesis that have been re-classified from Class IIb to Class III medical devices, annual report information must be submitted if the device meets either of the following criteria: I.The device was subject to a TGA application audit based on revision rate when the device transitioned from Class IIb to Class III; and/or II.No devices were supplied to the Australian marketplace before 30 June 2012. As per the standard automatic condition, annual reports should be submitted each year for the first three years of inclusion as a Class III medical device on the ARTG.
- Where a medical device included in the Register, contains a substance which is included in the Fourth Schedule to the Customs (Prohibited Imports) Regulations or the Eighth Schedule to the Customs (Prohibited Exports) Regulations the Sponsor shall, at the time of importation or exportation of the

medical device, be in possession of a licence and a permission for importation or exportation of each consignment of the goods as required by those regulations.

• A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Products Covered by This Entry

1. Dental precision attachment, ball

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 222929 ARTG Start Date: 30/04/2014



Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Cement, dental, polymer based

ARTG Identifier 222930

ARTG Start date 30/04/2014

Product Category Medical Device Included Class IIa

GMDN 35870

GMDN Term Resin-composite dental cement

Intended Purpose This is a polymer-based material dispensed as a powder or liquid

containing polymethyl methacrylate, a filler, a plasticiser or a

polymerization initiator, intended for use in the cementation or fixation of restorations and appliance during dental procedures in the mouth.

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	DV-2014-MC-05062-1

ARTG Standard Conditions

- The automatic conditions applicable to the inclusion of all kinds of medical devices in the Register are as specified in section 41FN of the Therapeutic Goods Act 1989.
- The standard conditions that are imposed under section 41FO of the Therapeutic Goods Act 1989 when kinds of medical devices are included in the Register are as set out in the following paragraphs.
- For a medical device included in the Register under Chapter 4 and imported into Australia, the Sponsor must ensure that information about the Sponsor is provided in such a way as to allow the sponsor to be identified.
- Each sponsor shall retain records of the distribution of all of the sponsor's medical devices included in the Register under Chapter 4. In the case of records relating to a Class AIMD medical device, Class III medical device, or Class IIb medical device that is an implantable medical device, the distribution records shall be retained for a minimum period of 10 years. In the case of records relating to any other device, the distribution records shall be retained for a minimum period of 5 years.
- The sponsor of a medical device included in the Register under Chapter 4 shall keep an up to date log
 of information of the kind specified in Regulation 5.8.
- It is a condition of inclusion in the ARTG that the sponsor of a medical device that is an AIMD, Class III or implantable Class IIb provides three consecutive annual reports to the Head of the Office of Product Review, Therapeutic Goods Administration following inclusion of the device in the ARTG (as specified in 5.8 of the regulations). Annual reports are due on 1 October each year. Reports should be for the period 1 July to 30 June. The first report following the date of inclusion in the ARTG must be for a period of at least six months but no longer than 18 months. Subsequent reports are to be provided on 1 October for a further 2 years. The annual report must include all complaints and adverse events received by the manufacturer relating to problems with the use of the device that have been received by them over the year. For orthopaedic implant prosthesis that have been re-classified from Class IIb to Class III medical devices, annual report information must be submitted if the device meets either of the following criteria: I.The device was subject to a TGA application audit based on revision rate when the device transitioned from Class IIb to Class III; and/or II.No devices were supplied to the Australian marketplace before 30 June 2012. As per the standard automatic condition, annual reports should be submitted each year for the first three years of inclusion as a Class III medical device on the ARTG.
- · Where a medical device included in the Register, contains a substance which is included in the Fourth

Schedule to the Customs (Prohibited Imports) Regulations or the Eighth Schedule to the Customs (Prohibited Exports) Regulations the Sponsor shall, at the time of importation or exportation of the medical device, be in possession of a licence and a permission for importation or exportation of each consignment of the goods as required by those regulations.

• A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Products Covered by This Entry

1. Cement, dental, polymer based

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 222930 ARTG Start Date: 30/04/2014



Australian Government

Department of Health

Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Gauge, depth, dental implant

ARTG Identifier 222931

ARTG Start date 30/04/2014

Product Category Medical Device Included Class Im

GMDN 31846

GMDN Term Gauge, depth, dental implant

Intended Purpose This is a hand-held dental instrument used in surgical implant

procedures to accurately measure various depths. This device has

graduations on the measuring tip denoting specific depths

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	DV-2014-MC-05062-1

ARTG Standard Conditions

- The automatic conditions applicable to the inclusion of all kinds of medical devices in the Register are as specified in section 41FN of the Therapeutic Goods Act 1989.
- The standard conditions that are imposed under section 41FO of the Therapeutic Goods Act 1989 when kinds of medical devices are included in the Register are as set out in the following paragraphs.
- For a medical device included in the Register under Chapter 4 and imported into Australia, the Sponsor must ensure that information about the Sponsor is provided in such a way as to allow the sponsor to be identified.
- Each sponsor shall retain records of the distribution of all of the sponsor's medical devices included in the Register under Chapter 4. In the case of records relating to a Class AIMD medical device, Class III medical device, or Class IIb medical device that is an implantable medical device, the distribution records shall be retained for a minimum period of 10 years. In the case of records relating to any other device, the distribution records shall be retained for a minimum period of 5 years.
- The sponsor of a medical device included in the Register under Chapter 4 shall keep an up to date log of information of the kind specified in Regulation 5.8.
- It is a condition of inclusion in the ARTG that the sponsor of a medical device that is an AIMD, Class III or implantable Class IIb provides three consecutive annual reports to the Head of the Office of Product Review, Therapeutic Goods Administration following inclusion of the device in the ARTG (as specified in 5.8 of the regulations). Annual reports are due on 1 October each year. Reports should be for the period 1 July to 30 June. The first report following the date of inclusion in the ARTG must be for a period of at least six months but no longer than 18 months. Subsequent reports are to be provided on 1 October for a further 2 years. The annual report must include all complaints and adverse events received by the manufacturer relating to problems with the use of the device that have been received by them over the year. For orthopaedic implant prosthesis that have been re-classified from Class IIb to Class III medical devices, annual report information must be submitted if the device meets either of the following criteria: I.The device was subject to a TGA application audit based on revision rate when the device transitioned from Class IIb to Class III; and/or II.No devices were supplied to the Australian marketplace before 30 June 2012. As per the standard automatic condition, annual reports should be submitted each year for the first three years of inclusion as a Class III medical device on the ARTG.
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(Prohibited Exports) Regulations the Sponsor shall, at the time of importation or exportation of the medical device, be in possession of a licence and a permission for importation or exportation of each consignment of the goods as required by those regulations.

• A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Products Covered by This Entry

1. Gauge, depth, dental implant

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 222931 ARTG Start Date: 30/04/2014



Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Dental implant suprastructure device, temporary, reusable

ARTG Identifier 222928

ARTG Start date 30/04/2014

Product Category Medical Device Included Class IIa

GMDN 46122

GMDN Term Dental implant suprastructure device, temporary, reusable

Intended Purpose This is a prefabricated device used to create a suprastructure on a

dental implant to mimic preparations of natural teeth. It is used during dental implant restorative and laboratory procedures to provide a base (the intermediate fixture level) for a temporary restoration during the healing and sculpturing period of the soft-tissue prior to the fabrication and installation of the final restoration This is a reusable device.

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	DV-2014-MC-05062-1

ARTG Standard Conditions

- The automatic conditions applicable to the inclusion of all kinds of medical devices in the Register are as specified in section 41FN of the Therapeutic Goods Act 1989.
- The standard conditions that are imposed under section 41FO of the Therapeutic Goods Act 1989 when kinds of medical devices are included in the Register are as set out in the following paragraphs.
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- Each sponsor shall retain records of the distribution of all of the sponsor's medical devices included in the Register under Chapter 4. In the case of records relating to a Class AIMD medical device, Class III medical device, or Class IIb medical device that is an implantable medical device, the distribution records shall be retained for a minimum period of 10 years. In the case of records relating to any other device, the distribution records shall be retained for a minimum period of 5 years.
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 of information of the kind specified in Regulation 5.8.
- It is a condition of inclusion in the ARTG that the sponsor of a medical device that is an AIMD, Class III or implantable Class IIb provides three consecutive annual reports to the Head of the Office of Product Review, Therapeutic Goods Administration following inclusion of the device in the ARTG (as specified in 5.8 of the regulations). Annual reports are due on 1 October each year. Reports should be for the period 1 July to 30 June. The first report following the date of inclusion in the ARTG must be for a period of at least six months but no longer than 18 months. Subsequent reports are to be provided on 1 October for a further 2 years. The annual report must include all complaints and adverse events received by the manufacturer relating to problems with the use of the device that have been received by them over the year. For orthopaedic implant prosthesis that have been re-classified from Class IIb to Class III medical devices, annual report information must be submitted if the device meets either of the following criteria: I.The device was subject to a TGA application audit based on revision rate when the device transitioned from Class IIb to Class III; and/or II.No devices were supplied to the Australian

marketplace before 30 June 2012. As per the standard automatic condition, annual reports should be submitted each year for the first three years of inclusion as a Class III medical device on the ARTG.

- Where a medical device included in the Register, contains a substance which is included in the Fourth Schedule to the Customs (Prohibited Imports) Regulations or the Eighth Schedule to the Customs (Prohibited Exports) Regulations the Sponsor shall, at the time of importation or exportation of the medical device, be in possession of a licence and a permission for importation or exportation of each consignment of the goods as required by those regulations.
- A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Products Covered by This Entry

1. Dental implant suprastructure device, temporary, reusable

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 222928 ARTG Start Date: 30/04/2014



Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Dental implant suprastructure device, permanent

ARTG Identifier 222927

ARTG Start date 30/04/2014

Product Category Medical Device Included Class IIa

GMDN 44879

GMDN Term Dental implant suprastructure device, permanent

Intended Purpose This is a prefabricated device that is incorporated into, or creates, the

necessary suprastructure mimicking preparations of natural teeth. It is used during dental implant restorative procedures and will provide the permanent intermediate fixture level between the dental implant and the final restoration. These are abutments, healing caps, fixation scres and

sleeves.

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	DV-2014-MC-05062-1

ARTG Standard Conditions

- The automatic conditions applicable to the inclusion of all kinds of medical devices in the Register are as specified in section 41FN of the Therapeutic Goods Act 1989.
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 - It is a condition of inclusion in the ARTG that the sponsor of a medical device that is an AIMD, Class III or implantable Class IIb provides three consecutive annual reports to the Head of the Office of Product Review, Therapeutic Goods Administration following inclusion of the device in the ARTG (as specified in 5.8 of the regulations). Annual reports are due on 1 October each year. Reports should be for the period 1 July to 30 June. The first report following the date of inclusion in the ARTG must be for a period of at least six months but no longer than 18 months. Subsequent reports are to be provided on 1 October for a further 2 years. The annual report must include all complaints and adverse events received by the manufacturer relating to problems with the use of the device that have been received by them over the year. For orthopaedic implant prosthesis that have been re-classified from Class IIb to Class III medical devices, annual report information must be submitted if the device meets either of the following criteria: I.The device was subject to a TGA application audit based on revision rate when the device transitioned from Class IIb to Class III; and/or II.No devices were supplied to the Australian

marketplace before 30 June 2012. As per the standard automatic condition, annual reports should be submitted each year for the first three years of inclusion as a Class III medical device on the ARTG.

- Where a medical device included in the Register, contains a substance which is included in the Fourth Schedule to the Customs (Prohibited Imports) Regulations or the Eighth Schedule to the Customs (Prohibited Exports) Regulations the Sponsor shall, at the time of importation or exportation of the medical device, be in possession of a licence and a permission for importation or exportation of each consignment of the goods as required by those regulations.
- A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Products Covered by This Entry

1. Dental implant suprastructure device, permanent

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 222927 ARTG Start Date: 30/04/2014



Therapeutic Goods Administration

Australian Register of Therapeutic Goods Certificate

Issued to

AB Dental Australia Pty Ltd

for approval to supply

AB Dental Australia Pty Ltd - Screw endosteal dental implant, two-piece

ARTG Identifier 222926

ARTG Start date 30/04/2014

Product Category Medical Device Included Class IIb

GMDN 55849

GMDN Term Screw endosteal dental implant, two-piece

Intended Purpose This is a sterile device made of alloplastic materials (that is intended to

be surgically implanted into alveolar and/or basal bone of the mandible or maxilla to provide support and a means of retention for a dental

prosthesis. This is a two-piece device.

Manufacturer Details	Address	Certificate number(s)
AB Dental Devices Ltd	3 Habosem Street Minrav Buildings Northern Industrial Area, Ashdod, 77101 Israel	DV-2014-MC-05062-1

ARTG Standard Conditions

- The automatic conditions applicable to the inclusion of all kinds of medical devices in the Register are as specified in section 41FN of the Therapeutic Goods Act 1989.
- The standard conditions that are imposed under section 41FO of the Therapeutic Goods Act 1989 when kinds of medical devices are included in the Register are as set out in the following paragraphs.
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- Each sponsor shall retain records of the distribution of all of the sponsor's medical devices included in the Register under Chapter 4. In the case of records relating to a Class AIMD medical device, Class III medical device, or Class IIb medical device that is an implantable medical device, the distribution records shall be retained for a minimum period of 10 years. In the case of records relating to any other device, the distribution records shall be retained for a minimum period of 5 years.
- The sponsor of a medical device included in the Register under Chapter 4 shall keep an up to date log
 of information of the kind specified in Regulation 5.8.
- It is a condition of inclusion in the ARTG that the sponsor of a medical device that is an AIMD, Class III or implantable Class IIb provides three consecutive annual reports to the Head of the Office of Product Review, Therapeutic Goods Administration following inclusion of the device in the ARTG (as specified in 5.8 of the regulations). Annual reports are due on 1 October each year. Reports should be for the period 1 July to 30 June. The first report following the date of inclusion in the ARTG must be for a period of at least six months but no longer than 18 months. Subsequent reports are to be provided on 1 October for a further 2 years. The annual report must include all complaints and adverse events received by the manufacturer relating to problems with the use of the device that have been received by them over the year. For orthopaedic implant prosthesis that have been re-classified from Class IIb to Class III medical devices, annual report information must be submitted if the device meets either of the following criteria: I.The device was subject to a TGA application audit based on revision rate when the device transitioned from Class IIb to Class III; and/or II.No devices were supplied to the Australian marketplace before 30 June 2012. As per the standard automatic condition, annual reports should be submitted each year for the first three years of inclusion as a Class III medical device on the ARTG.

- Where a medical device included in the Register, contains a substance which is included in the Fourth Schedule to the Customs (Prohibited Imports) Regulations or the Eighth Schedule to the Customs (Prohibited Exports) Regulations the Sponsor shall, at the time of importation or exportation of the medical device, be in possession of a licence and a permission for importation or exportation of each consignment of the goods as required by those regulations.
- A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Products Covered by This Entry

1. Screw endosteal dental implant, two-piece

Product Specific Conditions

No specific conditions have been recorded against this entry.

Therapeutic Goods Administration PO Box 100, Woden ACT 2606 Australia

Phone: 1800 020 653 Email: info@tga.gov.au ARTG Identifier: 222926 ARTG Start Date: 30/04/2014