| Justification: | Function: | Electronic signature justification: | Signed: Date (yyyy-mm-dd) - Time (hh:mm): |
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| Issued: | QA | Ulrika Svensson - SEHRBHNU | 2023-05-03 - 16:08 |
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| Approved: | OP | Martin Richardson - MARRIC | 2023-05-05 - 12:45 |
| Released: | | | |

This document has been electronically signed by the persons above.



DECLARATION OF CONFORMITY

FOR THE PRODUCT(S)

TheraBite® ActiveBand™ Kit Basic UDI: 7331791-JAW-A-000-0000-QQ

We, Atos Medical AB, hereby declare under our sole responsibility that the devices listed below comply with European Medical Devices Regulation (EU) 2017/745 and, any other applicable Union legislation and, clause 6.6 of Schedule 3 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002 and, UK Medical Devices Regulations 2002, including amendments in effect at the issuance date.

Intended use/purpose:

The TheraBite ActiveBand is an elastic silicone band used together with the TheraBite Jaw Mobilizer to increase and/or maintain muscle strength and endurance of the muscles of mastication (chewing muscles). The TheraBite ActiveBand is intended for single-patient use only.

Hörby, Sweden. Date as stated above

Martin Richardson, Senior Vice President Operations & Quality on behalf of the CEO of Atos Medical AB.

Manufacturer: Atos Medical AB

Kraftgatan 8, SE-242 35 Hörby

8, SE-242 33 HOIDY

Email: Web:

Telephone:

+46 (0)415 198 00 Info@atosmedical.com www.atosmedical.com

SRN number: SE-MF-000000725

Competent Authority Medical Products Agency

Sweden

FOR THE PRODUCT(S)

7331791-JAW-A-000-0000-QQ

| REF | Device name | Class | GMDN code |
|------|--------------------------|-------|-----------|
| 8260 | TheraBite ActiveBand Kit | 1 | 17802 |

In compliance with Therapeutic Goods (Medical Devices) Regulations 2002

Each kind of medical device to which the system has been applied complies with the applicable provisions of the essential principles, the classification rules, and the full quality assurance procedures, at each stage, from the design of the device until its final inspection before being supplied.

In compliance with UK Medical Devices Regulations 2002 as amended

Each kind of medical device to which the system has been applied fulfils the applicable obligations imposed by Annex VII and the device meets the provisions of the applicable section of the Medical Devices Regulations.

Authorized representative/UK Responsible Person:

Atos Medical UK Ltd Tottle Road Cartwright House Nottingham Nottinghamshire NG2 1RT England, United Kingdom

Common Specification(s) as per Article 9, and other Union Legislation(s)

- No relevant Common Specifications as defined in Article 9 to list.
- No relevant Union Legislations to list
- No European Representative



FOR THE PRODUCT(S)

TheraBite® Bite Pad

Basic UDI: 7331791-JAW-A-000-0001-QT

This Declaration of Conformity is issued under the sole responsibility of Atos Medical AB. Atos Medical AB declares that the devices covered by the present declaration are in conformity with the Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices and the requirements specified herein are fulfilled and, clause 6.6 of Schedule 3 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002 including amendments in effect at the issuance date.

Intended use/purpose:

The TheraBite Bite Pads are self-adhesive pads intended to protect the user's teeth.

Hörby, Sweden, date as stated on last page

Martin Richardson, Senior Vice President Operations & Quality on behalf of the CEO of Atos Medical AB.

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Manufacturer: Atos Medical AB

Kraftgatan 8, SE-242 35 Hörby

Sweden

Telephone: Email: Web: +46 (0)415 198 00 Info@atosmedical.com www.atosmedical.com

SRN number: SE-MF-000000725

Competent Authority Medical Products Agency

FOR THE PRODUCT(S)

7331791-JAW-A-000-0001-QT

| REF | Device name | Class* | GMDN code |
|-------|---------------------------------------|--------|-----------|
| PA001 | TheraBite Bite Pad, Regular (4 pcs) | 1 | 17802 |
| PA002 | TheraBite Bite Pad, Edentulous (4pcs) | 1 | 17802 |
| PA003 | TheraBite Bite Pad, Pediatric (4 pcs) | 1 | 17802 |

^{*}Product Classification according to Annex VIII EU Medical Device Regulation 2017/745

In compliance with Therapeutic Goods (Medical Devices) Regulations 2002

Each kind of medical device to which the system has been applied complies with the applicable provisions of the essential principles, the classification rules, and the full quality assurance procedures, at each stage, from the design of the device until its final inspection before being supplied.

In compliance with UK Medical Devices Regulations 2002 as amended

Each kind of medical device to which the system has been applied fulfils the applicable obligations imposed by Annex VII and the device meets the provisions of the applicable section of the Medical Devices Regulations.

Authorized representative/UK Responsible Person:

Atos Medical UK Ltd Tottle Road Cartwright House Nottingham

Nottinghamshire NG2 1RT England, United Kingdom

Common Specification(s) as per Article 9, and other Union Legislation(s)

- No relevant Common Specifications as defined in Article 9 to list.
- No relevant Union Legislations to list
- No European Representative

Document Number: VV-0544757 Status: Approved Version: 2.0 Name: DoC TheraBite Bite Pad 7331791-JAW-A-000-0001-QT

Document Approvals Approved Date: 2023-10-25

| Approval Task Verdict: Approve | SEHRBHNU Ulrika Svensson, Quality Assurance Specialist (ulrika.svensson-atosmedical@coloplast.com) Issuer 25-Oct-2023 12:26:33 GMT+0000 | |
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| Approval Task Verdict: Approve | MARRIC Martin Richardson, VP, International Manufacturing, Sweden, Germany & China (martin.richardson-atosmedical@coloplast.com) Management 25-Oct-2023 18:45:34 GMT+0000 | |

Document Number: VV-0544757 Status: Approved Version: 2.0 Name: DoC TheraBite Bite Pad 7331791-JAW-A-000-0001-QT



FOR THE PRODUCT(S)

TheraBite® Jaw Motion Rehabilitation System™ Basic UDI: 7331791-JAW-0-000-0000-98

This Declaration of Conformity is issued under the sole responsibility of Atos Medical AB. Atos Medical AB declares that the devices covered by the present declaration are in conformity with the Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices and the requirements specified herein are fulfilled and, clause 6.6 of Schedule 3 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002 including amendments in effect at the issuance date.

Intended use/purpose:

The TheraBite Jaw Motion Rehabilitation System is indicated for individuals who have, or are at risk of developing trismus (restrictions in their ability to open their jaw), and/or experience pain in the joints and/or muscles of the jaw. The device can also be used as a rehabilitation tool for postoperative physical therapy of the jaw, or to maintain the mouth open in a stable position, for example while performing dysphagia exercises. The TheraBite system is intended for single-patient use only.

Hörby, Sweden, date as stated on last page

Martin Richardson, Senior Vice President Operations & Quality on behalf of the CEO of Atos Medical AB.

Manufacturer: Atos Medical AB

Kraftgatan 8, SE-242 35 Hörby

Sweden

Telephone: Email: Web: +46 (0)415 198 00 Info@atosmedical.com www.atosmedical.com

SRN number: SE-MF-000000725

Competent Authority Medical Products Agency

FOR THE PRODUCT(S)

7331791-JAW-0-000-0000-98

| REF | Device name | Class* | GMDN code |
|-------|--|--------|-----------|
| TH001 | TheraBite Jaw Motion Rehabilitation System Adult | 1 | 17802 |
| TH002 | TheraBite Jaw Motion Rehabilitation System Pediatric | I | 17802 |

^{*}Product Classification according to Annex VIII EU Medical Device Regulation 2017/745

In compliance with Therapeutic Goods (Medical Devices) Regulations 2002

Each kind of medical device to which the system has been applied complies with the applicable provisions of the essential principles, the classification rules, and the full quality assurance procedures, at each stage, from the design of the device until its final inspection before being supplied.

In compliance with UK Medical Devices Regulations 2002 as amended

Each kind of medical device to which the system has been applied fulfils the applicable obligations imposed by Annex VII and the device meets the provisions of the applicable section of the Medical Devices Regulations.

Authorized representative/UK Responsible Person:

Atos Medical UK Ltd Tottle Road Cartwright House Nottingham Nottinghamshire NG2 1RT England, United Kingdom

Common Specification(s) as per Article 9, and other Union Legislation(s)

- No relevant Common Specifications as defined in Article 9 to list.
- No relevant Union Legislations to list
- No European Representative

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Document Number: VV-0544758 Status: Approved Version: 2.0 Name: DoC TheraBite Jaw Motion Rehab. System 7331791-JAW-0-000-0000-98

| Justification: | Function: | Electronic signature justification: | Signed: Date (yyyy-mm-dd) - Time (hh:mm): |
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| Released: | | | |

This document has been electronically signed by the persons above.



DECLARATION OF CONFORMITY

FOR THE PRODUCT(S)

TheraBite® Range of Motion Scale Basic UDI: 7331791-JAW-A-000-0002-QW

We, Atos Medical AB, hereby declare under our sole responsibility that the devices listed below comply with European Medical Devices Regulation (EU) 2017/745 and, any other applicable Union legislation and, clause 6.6 of Schedule 3 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002 and, UK Medical Devices Regulations 2002, including amendments in effect at the issuance date.

Intended use/purpose:

The Range of Motion Scale is used to monitor the progress of the rehabilitation program.

Hörby, Sweden. Date as stated above

Martin Richardson, Senior Vice President Operations & Quality on behalf of the CEO of Atos Medical AB.

Manufacturer: Atos Medical AB

Kraftgatan 8, SE-242 35 Hörby

Sweden

Telephone: Email: Web: +46 (0)415 198 00 Info@atosmedical.com www.atosmedical.com

SRN number: **SE-MF-000000725**

Competent Authority Medical Products Agency

FOR THE PRODUCT(S)

7331791-JAW-A-000-0002-QW

| REF | Device name | Class | GMDN code |
|-------|---------------------------------|-------|-----------|
| SC001 | TheraBite Range of Motion Scale | 1 | 17802 |

In compliance with Therapeutic Goods (Medical Devices) Regulations 2002

Each kind of medical device to which the system has been applied complies with the applicable provisions of the essential principles, the classification rules, and the full quality assurance procedures, at each stage, from the design of the device until its final inspection before being supplied.

In compliance with UK Medical Devices Regulations 2002 as amended

Each kind of medical device to which the system has been applied fulfils the applicable obligations imposed by Annex VII and the device meets the provisions of the applicable section of the Medical Devices Regulations.

Authorized representative/UK Responsible Person:

Atos Medical UK Ltd Tottle Road Cartwright House Nottingham Nottinghamshire NG2 1RT England, United Kingdom

Common Specification(s) as per Article 9, and other Union Legislation(s)

- No relevant Common Specifications as defined in Article 9 to list.
- No relevant Union Legislations to list
- No European Representative