Justification:	Function:	Electronic signature justification:	Signed: Date (yyyy-mm-dd) - Time (hh:mm):
Issued:	QA	Ulrika Svensson - SEHRBHNU	2020-12-09 - 13:03
Reviewed:	QA	John Wennborg - JOHWEN	2020-12-09 - 13:23
Approved:	OP	Martin Richardson - MARRIC	2020-12-09 - 16:58
Released:	QA	Ulrika Svensson - SEHRBHNU	2020-12-09 - 17:41



# **DECLARATION OF CONFORMITY**

# Provox<sup>®</sup> Life<sup>™</sup> Adhesives

Basic UDI: 7331791-ADH-0-000-0001-CT

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 clause 6.6 of Schedule 3 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002.

#### Intended use/purpose:

Provox Life Adhesives are single use adhesives that provide attachment for Provox Life HMEs and accessories after total laryngectomy.

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 Tel: +46 (0)415 198 00

Email: info@atosmedical.com, Web: www.atosmedical.com

#### **Competent Authority:**

# 7331791-ADH-0-000-0001-CT

REF	Name	Class	GMDN code
7460	Provox Life Standard Adhesive Round	I	62175
7461	Provox Life Standard Adhesive Oval	I	62175
7462	Provox Life Standard Adhesive Plus	I	62175
7463	Provox Life Sensitive Adhesive Round	I	62175
7464	Provox Life Sensitive Adhesive Oval	I	62175
7466	Provox Life Sensitive Adhesive Plus	I	62175
8261	Provox Life Night Adhesive	I	62175
8263	Provox Life Stability Adhesive	I	62175

#### 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.

Justification:	Function:	Electronic signature justification:	Signed: Date (yyyy-mm-dd) - Time (hh:mm):
Issued:	QA	Ulrika Svensson - SEHRBHNU	2020-11-13 - 07:49
Reviewed:	QA	John Wennborg - JOHWEN	2020-11-13 - 12:51
Approved:	OP	Martin Richardson - MARRIC	2020-11-13 - 13:01
Released:	QA	Ulrika Svensson - SEHRBHNU	2020-11-17 - 07:23



# **DECLARATION OF CONFORMITY**

# Provox® Life™ BasePlate Adaptor Basic UDI: 7331791-HME-A-000-0005-FB

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 clause 6.6 of Schedule 3 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002.

#### Intended use/purpose:

Provox Life BasePlate Adaptor is an accessory that allows attaching medical device, e.g. an HME, with an ISO 15 mm standard connector to a Provox Life attachment.

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 Tel: +46 (0)415 198 00

Email: info@atosmedical.com, Web: www.atosmedical.com

#### **Competent Authority:**

# 7331791-HME-A-000-0005-FB

REF	Name	Class	GMDN code
8057	Provox Life BasePlate Adaptor	1	58705

#### 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.

Justification:	Function:	Electronic signature justification:	Signed: Date (yyyy-mm-dd) - Time (hh:mm):
Issued:	QA	Ulrika Svensson - SEHRBHNU	2021-02-04 - 16:47
Reviewed:	QA	John Wennborg - JOHWEN	2021-02-04 - 20:18
Approved:	OP	Martin Richardson - MARRIC	2021-02-05 - 08:52
Released:	QA	Ulrika Svensson - SEHRBHNU	2021-02-08 - 08:25



# **DECLARATION OF CONFORMITY**

# Provox® Life™ FreeHands HME Basic UDI: 7331791-HME-0-000-0008-XZ

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 clause 6.6 of Schedule 3 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002.

#### Intended use/purpose:

Provox Life FreeHands HME is a single use heat- and moisture exchanger intended for spontaneously breathing laryngectomized patients, utilizing a voice prosthesis and in combination with a Provox speaking valve or a DigiTop.

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 Tel: +46 (0)415 198 00

Email: info@atosmedical.com, Web: www.atosmedical.com

#### **Competent Authority:**

7331791-HME-0-000-0008-XZ

REF		Name	Class	GMDN code
7440	)	Provox Life FreeHands HME	I	58705
7477	7	Provox Life Sample Pack FreeHands HME	I	58705

#### 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.

Justification:	Function:	Electronic signature justification:	Signed: Date (yyyy-mm-dd) - Time (hh:mm):
Issued:	QA	Ulrika Svensson - SEHRBHNU	2021-02-12 - 12:10
Reviewed:	QA	John Wennborg - JOHWEN	2021-02-12 - 13:11
Approved:	OP	Martin Richardson - MARRIC	2021-02-12 - 13:14
Released:	QA	Ulrika Svensson - SEHRBHNU	2021-02-12 - 13:22



# **DECLARATION OF CONFORMITY**

# Provox<sup>®</sup> Life<sup>™</sup> HMEs

Basic UDI: 7331791-HME-0-000-0001-XC

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 clause 6.6 of Schedule 3 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002.

#### Intended use/purpose:

Provox Life HMEs are single use heat- and moisture exchangers for patients breathing through a tracheostoma.

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 Tel: +46 (0)415 198 00

Email: info@atosmedical.com, Web: www.atosmedical.com

#### **Competent Authority:**

# 7331791-HME-0-000-0001-XC

REF	Name	Class	GMDN code
7475	Provox Life Sample Pack Protect HME	1	58705
7476	Provox Life Sample Pack Energy HME	1	58705
8310	Provox Life Go HME	1	58705
8311	Provox Life Home HME	1	58705
8312	Provox Life Energy HME	1	58705
8313	Provox Life Protect HME	1	58705
8262	Provox Life Night HME	1	58705

#### 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.

Justification:	Function:	Electronic signature justification:	Signed: Date (yyyy-mm-dd) - Time (hh:mm):
Issued:	QA	Ulrika Svensson - SEHRBHNU	2021-02-04 - 16:46
Reviewed:	QA	John Wennborg - JOHWEN	2021-02-04 - 20:13
Approved:	OP	Martin Richardson - MARRIC	2021-02-05 - 08:53
Released:	QA	Ulrika Svensson - SEHRBHNU	2021-02-08 - 08:25



# **DECLARATION OF CONFORMITY**

# Provox® Life™ Sample Packs

Basic UDI: 7331791-KIT-0-000-0003-HV

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 clause 6.6 of Schedule 3 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002.

#### Intended use/purpose:

Provox Life HMEs are single use heat- and moisture exchangers for patients breathing through a tracheostoma.

Provox Life Adhesives are single use adhesives that provide attachment for Provox Life HMEs and accessories after total laryngectomy.

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 Tel: +46 (0)415 198 00

Email: info@atosmedical.com, Web: www.atosmedical.com

#### **Competent Authority:**

# 7331791-KIT-0-000-0003-HV

REF	Name	Class	GMDN code
7467	Provox Life Sample Pack Standard Round	I	58705
7468	Provox Life Sample Pack Standard Oval	1	58705
7469	Provox Life Sample Pack Standard Plus	I	58705
7470	Provox Life Sample Pack Sensitive Round	I	58705
7471	Provox Life Sample Pack Sensitive Oval	I	58705
7472	Provox Life Sample Pack Sensitive Plus	I	58705
7473	Provox Life Sample Pack Stability	I	58705
7474	Provox Life Sample Pack Night	I	58705

#### 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.

Justification:	Function:	Electronic signature justification:	Signed: Date (yyyy-mm-dd) - Time (hh:mm):
Issued:	QA	Ulrika Svensson - SEHRBHNU	2020-12-03 - 15:03
Reviewed:	QA	John Wennborg - JOHWEN	2020-12-07 - 11:48
Approved:	OP	Martin Richardson - MARRIC	2020-12-07 - 12:18
Released:	QA	Ulrika Svensson - SEHRBHNU	2020-12-07 - 12:59



# **DECLARATION OF CONFORMITY**

# **Provox® Life™ Shower**

Basic UDI: 7331791-ADH-A-000-0001-UB

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 clause 6.6 of Schedule 3 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002.

#### Intended use/purpose:

Provox Life Shower is a single patient use device intended to be placed into a Provox Life attachment to avoid water from entering the tracheostoma during showering.

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 Tel: +46 (0)415 198 00

Email: info@atosmedical.com, Web: www.atosmedical.com

#### **Competent Authority:**

# Document No: 10000043987 Edition: 03 Release date: 2020-12-07

# **DECLARATION OF CONFORMITY**

# 7331791-ADH-A-000-0001-UB

REF	Name	Class	GMDN code
8308	Provox Life Shower	1	62047

#### 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.