PEOPLE HAVE PRIORITY

Instructions for use





PL-40 H

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WARNING! (risk of injury)



ATTENTION! (to prevent damage occurring)



General explanations, without risk to persons or objects



Not for re-use



Call customer service

on the handpiece drive / handpiece sleeve



Follow instructions for use



Date of manufacture



Do not dispose of with domestic waste



Data Matrix code for product information including UDI (Unique Device Identification)



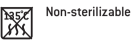
CE mark with identification number of the Notified Body



Non-ionizing electromagnetic radiation



The medical device with reference to electrical safety, mechanical safety and fire prevention conforms to UL 60601-1:2006, CAN/CSA-C22.2 No.601.1-M90:2005, CAN/CSA-C22.2 No. 60601-1:2008, ANSI/AAMI ES 60601-1:2005 25UX [Control No.]





Sterilizable up to the stated temperature

Thermo washer disinfectable



Ж

Not suitable for intracardiac application — Type BF appliance



Catalogue number

SN



DC – direct current

on the foot control

	CE mark with identification number of the Notified Body	(((₊))) ▲	Non-ionizing electromagnetic radiation	REF	Catalogue number
	Do not dispose of with domestic waste		DC – direct current	SN	Serial number
	Data Matrix code for product information including UDI (Unique Device Identification)	IPX1	Protection against dripping water	~~~	Date of manufacture
c SN u	s UL Component Recognition Mark indicates compliance with Canadian and		Foot control cordless C-NW		Reset

U.S. requirements

on the packaging



CE mark with identification number of the Notified Body



This way up



Fragile, handle with care



Keep dry



»Der Grüne Punkt« (The Green Dot) trademark of Duales System Deutschland GmbH



Trademark of RESY OfW GmbH for identification of recyclable transport and outer packaging of paper and cardboard



Data Matrix code for product information including UDI (Unique Device Identification)



Data structure in accordance with Health Industry Bar Code



Temperature limitation



Humidity limitation

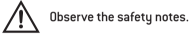


Caution! According to Federal law, this medical device may only be sold by or on the order of a dentist, physician or any other medical practitioner licensed by the law of the State in which he or she practices and intends to use or order the use of this medical device.

1. Introduction

For your safety and the safety of your patients

These Instructions for Use explain how to use your medical device. However, we must also warn against possible hazardous situations. Your safety, the safety of your team and, of course, the safety of your patients are of paramount importance to us.



Intended use

- PL-40 H: Battery driven electrical drive unit with wireless foot controller to perform cleaning and polishing of tooth surfaces and fillings.
- C-NW: Foot control for operation of medical electrical equipment.



Misuse may damage the medical device and hence cause risks and hazards for patient, user and third parties.

Qualifications of the user

We have based our development and design of the medical device on the target group »dentist, dental hygienist, dental employees (prophylaxis) and dental assistants«.

Introduction

C E Production according to EU Directive The handpiece drive meets the requirements of Directive 93/42/EEC. 0297 The foot control meets the requirements of Directive 93/42/EEC and RED Directive 2014/53/EU.

Responsibility of the manufacturer

The manufacturer can only accept responsibility for the safety, reliability and performance of the medical device when it is used in compliance with the following directions:

- The medical device must be used in accordance with these instructions for use. >
- Only the components approved by the manufacturer may be replaced (0-ring). >
- Modifications or repairs must only be undertaken by an authorized W&H service partner (see page 48). >
- The foot control has no components that can be repaired by the user. >
- The electrical installation at the premises must comply with the regulations laid out in IEC 60364-7-710 (»Installation of > electrical equipment in rooms used for medical purposes«) or with the regulations applicable in your country.
- Unauthorized opening of the medical device invalidates all claims under warranty and any other claims. >

Improper use, unauthorized assembly, modification or repair to the medical device, non-compliance with our instructions or the use of accessories and spare parts which are not approved by W&H, invalidates all claims under warranty and any other claims.

2. Electromagnetic compatibility (EMC)



Medical electrical equipment is subject to particular precautions with regard to EMC and must be installed and put into operation in accordance with the EMC notes included.

W&H guarantees the compliance of the device with the EMC requirements only when used with original W&H accessories and spare parts. The use of accessories and spare parts not approved by W&H can lead to an increased emission of electromagnetic interference or to a reduced resistance against electromagnetic interference.



You can find the current EMC manufacturer's declaration on our website at http://wh.com or request a copy directly from the manufacturer.



HF communication equipment

Do not use any portable and mobile HF communication equipment (e.g., mobile telephones) during operation. These may affect medical electrical equipment.



The medical device may be interfered by other equipment, even if these other devices comply with CISPR (International special committee on radio interference) emission requirements.

3. Scope of delivery

REF	Description
30317000	Handpiece drive
30316000	Foot control C-NW with Stick
07969610	Charger with adaptor
05882600	Handpiece holder

4. Safety notes



- > Before using the medical device for the first time, store it at room temperature for 24 hours.
- > Check the medical device for damage and loose parts each time before using.
- > Do not operate the medical device if it is damaged.
- > Always ensure that the correct operating conditions are provided.
- > Perform a test run each time before using.
- > Never touch the patient and the electrical contacts on the medical device simultaneously.
- > Only put the medical device into operation when the handpiece sleeve is attached.
- > Keep the foot control away from magnetic fields.
- > Replace the foot control as soon as the resistance is noticeably reduced.



> Do not expose the medical device to any violent mechanical impacts.

Battery



- > Do not charge the battery unattended.
- > As soon as the charging cycles start to deteriorate send the medical device to an authorized W&H service partner.
- > Defective or worn-out batteries must only be replaced by an authorized W&H service partner.



- > Charge the battery of the medical device as soon as the status LED flashes.
- > Incorrect use of the rechargeable battery can cause fire or corrosion.

Safety notes



P The medical device is classed as »conventional equipment« (closed equipment without protection against the ingress of water).



The medical device is not approved for operation in potentially explosive atmospheres.



ChargerOnly use the charger supplied.



Hygiene and maintenance prior to initial use
> Clean and disinfect the medical device.
> Sterilize the handpiece sleeve.

System failure

A total system failure does not constitute a critical fault. Simply switch the unit off and then on again.

Safety notes

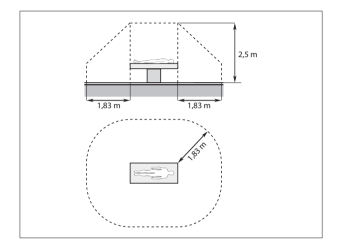


Risks due to electromagnetic fields

The functionality of implantable systems, such as cardiac pacemakers and implantable cardioverter defibrillators (ICD) can be affected by electric, magnetic and electromagnetic fields. This medical device complies with the reference values defined in EN 50527-2-1/2016 for unipolar and bipolar pacemakers and is therefore suitable for patients with pacemakers.

- > Find out if patient and user have implanted systems before using the medical device and consider the application.
- > Keep a safe distance of at least 7 cm between the medical device and the cardiac pacemakers.
- > Make appropriate emergency precautions and take immediate action on any signs of ill-health.
- > Symptoms such as raised heartbeat, irregular pulse and dizziness can be signs of a problem with a cardiac pacemaker or ICD.

Safety notes



The patient environment (see diagram) encompasses the area up to 2.50 m above the patient and 1.83 m in all horizontal directions.



The charger must not be used within the patient environment.



The Prophy Angles are disposable articles.



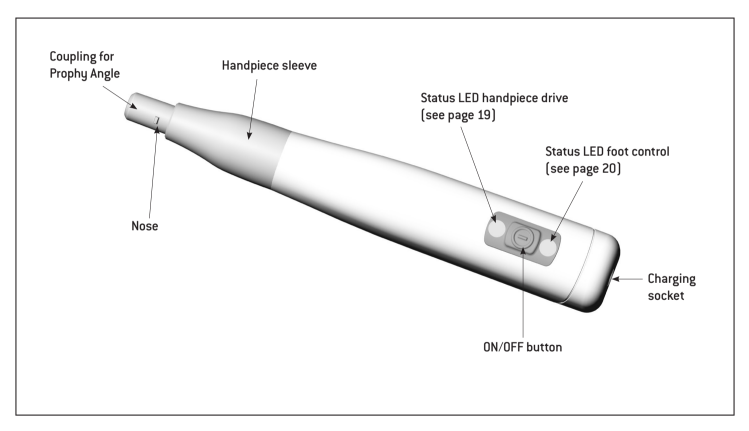
> Use only Prophy Angles which are in perfect condition. Follow the operating instructions of the manufacturer.

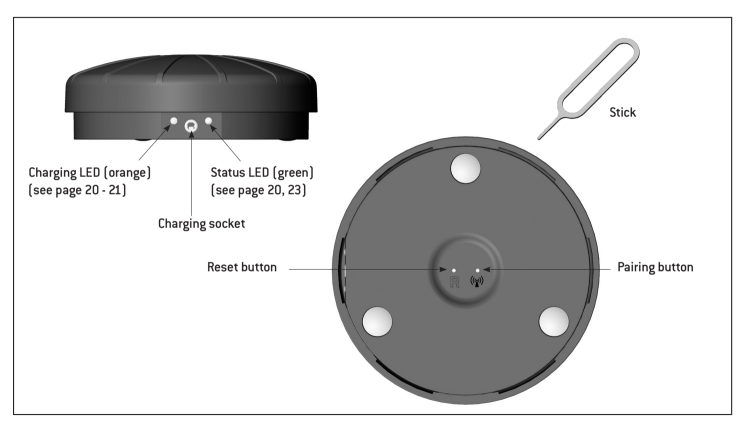
> Insert the Prophy Angle only when the medical device is stationary.
 > Never touch the Prophy Angle while it is still rotating.



> Only use Prophy Angles with plastic shanks for the Doriot system. Prophy Angles with metal shanks damage the clamping chuck system.

5. Description





Description



Standby mode

- > The handpiece can be activated with the ON/OFF button.
- > If the handpiece drive is not used for longer than 4 minutes, it returns to standby mode automatically.

LED	steady	flashes	flashes intermittently
GREEN	 → Battery is 25–100 % charged → Pairing successful > Handpiece drive is ready for operaion 	→ Pairing active	
ORANGE	→ Battery is charging	 → Battery is 2–25 % charged > Complete the treatment > Do not start any further treatment > Charge the battery 	 → Battery is 1 % charged > Charge the battery
RED		 → Error message > Contact an authorised W&H service partner 	

LED	flashes	flashes alternately	
ORANGE	 → Battery of foot control is flat > Complete the treatment > Charge the battery of the foot control 	 → Pairing unsuccessful > Troubleshooting with pairing problems (see page 23) 	



Standby mode

> The foot control can be activated by pressing.

LED	steady	steady	flashes	flashes intermittently*
	• 🕞 •	• _G •		• • •
GREEN	→ Connection to paired medical device established		→ Foot control is attempting to establish a connection to the paired medical device	 → Battery is flat > Charge the battery
ORANGE		→ Battery is charging		

* The LED flashes for 40 milliseconds every 4 seconds

6. Start-up



^o Charge the medical device fully before you use them for the first time.



• Attach the adaptor to the power supply unit.



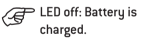
• Connect the charging cable into the foot control charging socket.

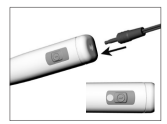


Plug the charger into an power socket.



LED orange: Battery is charging.





- Connect the charging cable up to the charging socket on the handpiece drive.
- ELD orange: Battery is charging.
 - LED green: Battery is charged.
- The handpiece drive will not switch to standby mode as long as it is plugged into the charging cable.

You can query the battery status when the handpiece drive is switched on and during the charging process.



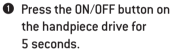
> Briefly press the ON/OFF button:

LED	flashes	Battery status			
	3 x green	75–100 %			
	2 x green	50–75 %			
	1 x green	25–50 %			
	orange	2–25 %			

Start-up

The foot control and handpiece drive are already paired when delivered! If pairing is active: Status LED (green) flashes on the foot control.

- > Both devices need to be in pairing mode for it to be possible to pair the handpiece drive and foot control.
- > To activate the handpiece drive's pairing mode, place it in the vicinity of the foot control.

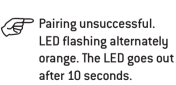


Æ LED flashing green: The handpiece drive is in pairing mode for 30 seconds.

After 3 seconds the status I FD switches from flickering to flashing. The foot control's pairing mode is now activated

Pairing

Use the stick to press the pairing button on the foot control for 3 seconds.



Pairing successful.

LED green.









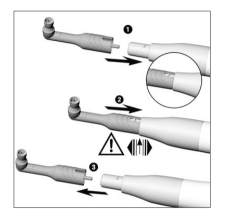
Troubleshooting with pairing problems

- > Remove any metallic objects located between the foot control and handpiece drive.
- > Change the position of the foot control.
- > Eliminate any sources of interference (e.g. brush motors, mobile telephones, radios, WLAN, ...)
- > Use the stick to press the reset button on the foot control and try pairing again.



If the pairing problem cannot be remedied using the steps described above, the unit will need to be inspected by an authorized W&H service partner.

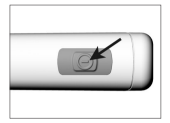
Start-up



Prophy Angle Cup or Brush

- Position the groove on the Prophy Angle with the nose of the handpiece drive.
 Push the Prophy Angle onto the handpiece drive until the limit stop.
 - Verify full engagement.
- Hold the handpiece sleeve firmly. Remove the Prophy Angle.

7. Handpiece drive



Switch on Press the ON/OFF button.



Press the foot control to variably control the speed of the disposable contra-angle handpiece.

 \mathcal{F} Press the foot control as far as it will go to attain the maximum speed of 3,000 rpm.

The following light signals are shown on the foot control:

Foot control pressed	
Status LED (green) flashes	Foot control is attempting to establish a connection to the paired medical device
Status LED (green) steady	Connection to paired medical device established



Switch off

• Keep the ON/OFF button depressed for 2 seconds.

Handpiece drive



Do not hold the handpiece drive at eye level!

- > Attach the Prophy Angle to the handpiece drive.
- > Operate the handpiece drive with the foot control.



In the event of operating malfunctions (e.g., vibrations, unusual noise, overheating, coolant failure or leakage) **stop the medical device immediately** and contact an authorized W&H service partner.

8. Hygiene and maintenance

Follow your local and national laws, directives, standards and guidelines for cleaning, disinfection and sterilization.



> Wear protective clothing, safety glasses, face mask and gloves.



Cleaning agents and disinfectants

- > Read the notes, follow the instructions and heed the warnings provided by the manufacturers of cleaning agents and/or disinfectants.
- > Use only detergents which are intended for cleaning and/or disinfecting medical devices made of metal and plastic.
- > It is imperative to comply with the concentrations and exposure times specified by the manufacturer of the disinfectant.
- > Use disinfectants which have been tested and found effective by the Verbund für Angewandte Hygiene e.V. (VAH = Association for Applied Hygiene), the Österreichischen Gesellschaft für Hygiene, Mikrobiologie und Präventivmedizin (ÖGHMP = Austrian Society for Hygiene, Microbiology and Preventive Medicine), the Food and Drug Administration (FDA) and the U.S. Environmental Protection Agency (EPA).
- > The user is responsible for validating its process if the specified cleaning agents and disinfectants are not available.

Hygiene and maintenance



The product lifetime and the medical device's ability to operate correctly are mainly determined by mechanical stress during use and chemical influences due to processing.

> Send worn or damaged medical devices and/or medical devices with material changes to an authorized W&H service partner.



Processing cycles

> We recommend to replace the handpiece sleeve after 600 processing cycles.

Hygiene and maintenance



> Clean the medical device immediately after every treatment.

> Wipe the entire handpiece drive, the handpiece sleeve, the handpiece holder and the foot control with disinfectant.



> Ensure that no fluids enter the medical device.



> Switch the handpiece drive off.> The charger must not be connected.



Note that the disinfectant used during pre-treatment is only for personal protection and cannot replace the disinfectant step after cleaning.

> Do not place the handpiece drive, the handpiece sleeve, the handpiece holder and the foot control in liquid disinfectant or in an ultrasonic bath.



> Do not immerse the handpiece drive and the foot control in water or clean them under running water.

Handpiece sleeve / Handpiece holder

- > Clean the handpiece sleeve and the handpiece holder under running tap water (< 35°C / < 95°F).
- > Rinse and brush off all internal and external surfaces.
- > Remove any liquid residues using compressed air.



> W&H recommends wiping down with disinfectant.



Evidence of the handpiece drive, the handpiece sleeve, the handpiece holder and the foot control basic suitability for effective manual disinfection was provided by an independent test laboratory using the »mikrozid® AF wipes« disinfectant (Schülke & Mayr GmbH, Norderstedt).

Handpiece sleeve



W&H recommends automated cleaning and disinfection using a washer-disinfector (WD). Read the notes, follow the instructions and heed the warnings provided by the manufacturers of washer-disinfectors, cleaning agents and/or disinfectants.



> The handpiece drive, the handpiece holder and the foot control are not approved for automated processing in a washer-disinfector and for sterilization.



^a Evidence of the medical device's basic suitability for effective automated disinfection was provided by an independent test laboratory using the »Miele PG 8582 CD« washer-disinfector (Miele & Cie. KG, Gütersloh) and the »Dr. Weigert neodisher[®] MediClean forte« cleaning agent (Dr. Weigert GmbH & Co. KG, Hamburg).

> Cleaning at 55°C (131°F) – 5 minutes

> Disinfection at 93°C (200°F) - 5 minutes

> Ensure that the medical device is completely dry internally and externally after cleaning and disinfection.

> Remove any liquid residues using compressed air.



- > Check the medical device after cleaning and disinfection for damage, visible residual soiling and surface changes.
- > Reprocess any medical devices that are still soiled.
- > Sterilize the handpiece sleeve following cleaning and disinfection.

Handpiece sleeve



Pack the handpiece sleeve in sterilization packages that meet the following requirements:

- > The sterilization package must meet the applicable standards in respect of quality and use and must be suitable for the sterilization method.
- > The sterilization package must be large enough for the sterilization goods.
- > The filled sterilization package must not be under tension.

Handpiece sleeve



W&H recommends sterilization according to EN 13060, EN 285.



Read the notes, follow the instructions and heed the warnings provided by the manufacturers of steam sterilizers.
 The program selected must be suitable for the handpiece sleeve.

Recommended sterilization procedures

- > Steam sterilization (type B, S)
- > Sterilization time at least 3 minutes at 134°C (273°F)
- > Maximum sterilization temperature 135°C (275°F)



 Evidence of the handpiece sleeve basic suitability for effective sterilization was provided by an independent test laboratory using the LISA 517 B17L steam sterilizer (W&H Sterilization S.r.I., Brusaporto (BG)) and the Systec VE-150 steam sterilizer (Systec).

> »Dynamic-air-removal prevacuum cycle« (type B): temperature 134°C (273°F) – 3 minutes*

> »Steam-flush pressure-pulse cycle« (type S): temperature 134°C (273°F) – 3 minutes*

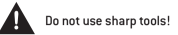
*EN 13060, EN 285, ISO 17665

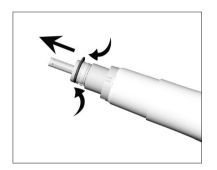
Handpiece sleeve



> Store sterile goods dust-free and dry.
> The shelf life of the sterile goods depends on the storage conditions and type of packaging.

9. Replacing the O-ring





- Pull the handpiece sleeve off the handpiece drive.
- Squeeze the 0-ring between your thumb and index finger firmly so that it forms a loop.

• Pull the old O-ring off.

• Push the new O-ring on in its place.

10. Servicing



Regular checks

Regular servicing of function and safety including the accessories is necessary and should be carried out at least once every three years, unless shorter intervals are prescribed by law. The inspection must be undertaken by a qualified organization and must include the following procedures:

Handpiece drive

- > External visual inspection
- > Function test

Foot control

- > External visual inspection
- > Function test with check to see if the maximum speed can be reached



The regular checking must only be performed by an authorized W&H service partner.

Servicing

Repairs and returns

In the event of operating malfunctions immediately contact an authorized W&H service partner. Repairs and maintenance work must only be undertaken by an authorized W&H service partner.



> Ensure that the medical device has been completely processed before returning it.

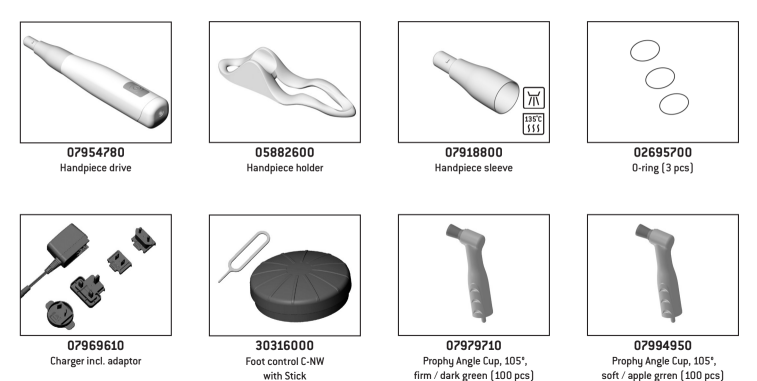


> Always return equipment in the original packaging!

11. W&H Accessories and spare parts



Use only original W&H accessories and spare parts or accessories approved by W&H. Supplier: W&H partners



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12. Technical data

-20°C bis +60°C (-4°F to +140°F)
8% to 80% (relative), non-condensing
+10°C to +35°C (+50°F to +95°F)
15% to 80% (relative), non-condensing

Handpiece drive	PL-40 H		
Battery type:	Li-lon		
Runtime:	8 treatments with a polishing duration of 6 min.		
Standby:	automatically after 4 min.		
Charging time:	approx. 2 h		
Rated voltage:	3,7 V		
Rated capacity:	680 mAh		
Max. speed	3.000 rpm		
Maximum torque:	2 Ncm		
Dimensions (WxDxH):	160 x 25 x 28 mm		
Weight:	118 g		

Technical data

Foot control	C-NW		
Battery type:	Li-ion		
Runtime:	approx. 2 months		
Standby:	automatically if not actuated		
Charging time:	approx. 3 h		
Rated voltage:	3.7 V		
Rated capacity:	680 mAh		
Dimensions (WxDxH):	117 x 117 x 38 mm		
Weight:	190 g		

Charger	
Rated voltage:	100 - 240 V
Permissible voltage fluctuation:	± 10 %
Frequency:	50 – 60 Hz
Power:	7 VA

Technical data

Classification according to Paragraph 6 of the General Specifications for the Safety of Medical Electrical Equipment according to IEC 60601-1/ANSI/AAMI ES 60601-1



Charger: Class II medical electrical equipment

Handpiece drive: Internally powered



Type BF applied part (not suitable for intracardiac application)



The C-NW foot control is protected against vertically falling drops of water (IPX1 as per IEC 60529)

Pollution level:	2
Overvoltage category:	II
Altitude:	up to 3,000 m above sea level



Temperature information

Temperature of the hand	piece at the operat	or side: n	nax. 55°C (max. :	L31ºF)
Temperature of the hand	piece at the patient	tside: m	nax. 50°C (max. :	L22°F)

13. Disposal



Ensure that the parts are not contaminated on disposal.

Follow your local and national laws, directives, standards and guidelines for disposal.



- > Medical device> Waste electrical equipment
- > Packaging

Explanation of warranty terms

This W&H medical device has been manufactured with great care by highly qualified specialists. A wide variety of tests and controls guarantees faultless operation. Please note that claims under warranty can only be validated when all the directions in the Instructions for Use have been followed.

As the manufacturer, W&H is liable for material or manufacturing defects within a warranty period of 12 months from the date of purchase. Accessories and consumables (handpiece holder, stick) are excluded from the warranty.

We accept no responsibility for damage caused by incorrect handling or by repairs carried out by third parties not authorized to do so by W&H!

Claims under warranty accompanied by proof of purchase must be sent to the vendor or to an authorized W&H service partner. The provision of service under warranty extends neither the warranty period nor any other guarantee period.

12 month warranty

Authorized W&H service partners

Find your nearest W&H service partner at http://wh.com Simply go to the menu option »Service« for full details. Alternatively please contact:

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W&H Impex Inc., 6490 Hawthorne Drive, Windsor, Ontario, N8T 1J9, Canada t + 1 800 2656277, 1 519 9446739, f + 1 519 9746121, E-Mail: service.ca@wh.com

W&H Impex Inc., 33091 W Jefferson Ave. Brownstown, MI-48173, USA t + 1 800 2656277, 1 519 9446739, f + 1 519 9746121, E-Mail: service.us@wh.com

A-DEC AUSTRALIA CO. INC., Unit 8, 5-9 Ricketty Street, Mascot NWS 2020, Australia t + 61 2 83324000, f + 61 2 83324099, E-Mail: a-dec@a-dec.com.au

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