Instructions for Use (IFU)

Protective Plug



Manufacturer:



Smart Denture Conversions, LLC 56 Hunter St Suite 320 Apex NC, 27511 855-550-8787

www.SmartDentureConversions.com

Training:

The following descriptions are insufficient to allow immediate use of the Smart Denture Conversions' system. Knowledge of implant-prosthetic treatment and instruction in the handling of the Smart Denture Conversions' system provided by an operator with relevant experience is necessary. It is strongly recommended that new and experienced users of Smart Denture Conversions' products complete special training before using a new product for the first time. Smart Denture Conversions offers a variety of training options. Please go the www.SmartDentureConversions.com for more information.

Product Description:

Protective Plugs are used to mask the mating surface of the TiBase while adding acrylic to the intaglio surface of the denture. The tip of the Protective Plug engages the PEEK Cap of the Separable Fastener that remains in the TiBase after the pickup. The table below summarizes the items:

Name	Part Number	Compatible Separable Fastener	Material	# of Uses
Protective Plug	SF-008	All	6061 T6/7075 T6 Aluminum	Single

Indications for Use:

The Protective Plugs are intended to be used in the lab, on a sterile denture with TiBases that have been picked up. The supplied Protective Plugs are intended for single use and a single patient. Re-use of single use devices creates a potential risk of patient or user infection and misfitting components. For more specific information on process steps, please refer to the Technique Manual located on the website www.SmartDentureConversions.com.

Contraindications:

It is contraindicated to using Smart Denture Conversions' Protective Plugs in:

- Patients who are medically unfit for an oral surgical procedure.
- Patients in whom adequate sizes, numbers or desirable positions of implants are not reachable to achieve safe support of functional or eventually parafunctional loads.
- Patients who show signs of allergy or hypersensitivity to the chemical components of the materials listed in the chart above.

Warning:

- Components are to be used by dental health care professionals and are to be used in patients subject to dental implant treatment.
- If the indication or the nature of use is not clear, do not use until all points have been clarified.
- Do not use if package is damaged.
- Always inspect components before use. Do not use damaged, deformed, corroded, or discolored components.
- Ensure products are secured against aspiration when handled intraorally. Aspiration of products may lead to infection or unplanned physical injury.

IFU-008 Rev-C (Draft) Page 1 of 4

Protective Plug



- Failure to follow the procedures outlined in these instructions may lead to any or all of the following complications: Aspiration or swallowing of a component, follow-up treatment, incorrect impression resulting in incompatible restorations.
- As the clinical outcome of dental treatment is influenced by multiple variables, even if the product is used
 according to the instructions for use the residual risks described below can. anaphylaxis (severe allergic
 reaction); aspiration or swallowing of components; pain; local infection; inflammation; local irritation; loss of
 product function; follow-up treatment.
- Smart Denture Conversions is not liable for damage resulting from use outside the intended use of the product.

Cautions/Precautions:

The following precautions are required before or during treatment:

- Do not use Smart Denture Conversion components after the expiration date indicated on the packaging.
- All products intended for single use must not be reused. Re-use of single use devices creates a potential risk of patient or user infection and misfitting components.
- Before every procedure make sure that all required components, instruments, and auxiliary equipment are complete, in operating order and available in the required quantity.
- If, due to unfavorable anatomical conditions, instruments do not fit or cannot be used for other reasons, the course of treatment planned with them must not be continued and alternatives must be sought.
- Always wear suitable personal protective equipment for your own safety.
- Position the patient such that the danger of aspiration of components is minimized.
- All components used in the patient's mouth must be secured to prevent aspiration and swallowing.

Magnetic Resonance Imaging (MRI) Safety Information:

All Smart Denture Conversion LLC products which remain in the patient's body have not been evaluated for safety and compatibility in the MR environment. They have not been tested for heating, migration, or image artifact in the MR environment. The safety of the Smart Denture Conversion LLC products in the MR environment is unknown. Scanning a patient who has such a product may result in patient injury.

Cleaning and Sterilization Instructions:

Components are delivered non-sterile by Smart Denture Conversions and are intended for single use. Prior to use, the devices must be cleaned and sterilized by the user.

Sterilization is to be performed corresponding to the following scheme:

1. **Preparation for sterilization:** Place components in a sterilization pouch which is legally marketed (for the US market: FDA-cleared) for use with the recommended sterilization parameters. Every sterilization package must have a sterilization indicator and sterilization date.

2. Sterilization:

Method	Cycle	Temperature	Exposure Time*	Dry Time	Cooling Time	Reference
Steam	Dynamic Air Removal (Prevacuum)	132°C (270°F)	4min	20min	30min	ANSI/AAMI
Steam	Gravity Displacement	121°C (250°F)	30min	30min	30min	TIR12:2010
Steam	Gravity Displacement	132°C (270°F)	15min	30min	30min	

^{*}Minimum exposure times, the operating times are longer and may vary depending on the device.

Storage, Handling and Transportation:

The devices must be stored in a dry place in their original packaging at room temperature and protected from direct sunlight. Improper storage may compromise essential material and design characteristics, leading to device failure.

IFU-008 Rev-C (Draft) Page **2** of **4**

Protective Plug



Disposal:

Safely discard potentially contaminated or no longer usable medical devices as healthcare (clinical) waste in accordance with local healthcare guidelines, country and government legislation or policy. Separation, recycling, or disposal of packaging material shall follow local country and government legislation on packaging and packaging waste, where applicable. If there is no current legislation, pack them in a perforating waste/sharps disposal container and dispose of them in hospital waste.

Basic UDI-DI Information:

The following table lists the Basic UDI-DI information for the devices described in this IFU.

Product	Catalog Number	Included Parts	Basic UDI-DI Number
Protective Plug 6PK, SDC	PP6PK	SF-008	+D990PP6PK0
Protective Plug 10PK, SDC	PP10PK	SF-008	+D990PP10PK0
Premium Starter Kit, SDC	PSK	SF-008	+D990PSK0
Premium Starter Kit, Straumann	PSK-ST	SF-008	+D990PSK-ST0
Premium Starter Kit, Tilobe	KDTL-PSK	SF-008	+D990KDTL-PSK0
Premium Starter Kit, Paltop	KDIH-PSK	SF-008	+D990KDIH-PSK0
Premium Starter Kit, Biohorizons	BHHD-PSK	SF-008	+D990BHHD-PSK0
Premium Starter Kit, TSV	ZVTS-PSK	SF-008	+D990ZVTS-PSK0
Premium Starter Kit, Low Profile	ZVLP-PSK	SF-008	+D990ZVLP-PSK0
Press-On Cap 10PK, ZimVie	ZV-POC10PK	SF-008	+D990ZV-POC10PK0
Press-On Cap 6PK, ZimVie	ZV-POC6PK	SF-008	+D990ZV-POC6PK0

Validity:

Upon publication of these instructions for use, all previous versions are superseded.

Availability:

Some items of Smart Denture Conversions are not available in all countries.

Warranty:

Please visit <u>www.SmartDentureConversions.com</u> for the most up to date warranty information.

IFU-008 Rev-C (Draft) Page **3** of **4**

Protective Plug



Symbols Glossary:

The following Symbols may be present on the device labeling or in information accompanying the device. Refer to the device labeling or accompanying information for the applicable symbols.

•••	Manufacturer	~~ <u> </u>	Date of Manufacture
	Use By Date	SN	Serial Number
LOT	Batch Code	REF	Catalog Number
UDI	Unique Device Identifier	MD	Medical Device
CE	CE Mark	UK RP	UK Representative
UK	United Kingdom Conformity Assessment Mark	UK 0086	United Kingdom Conformity Assessment Mark with Approved Body Number
EC REP	European Representative	NON	Non-Sterile
STERILE	Comes Sterilized	STERILE A	Comes Sterilized using Aseptic Processing
STERILEEO	Comes Sterilized using Ethylene Oxide Processing	STERILE R	Comes Sterilized using Irradiation Processing
STERILE	Comes Sterilized using Dry Heat Processing	STERMIZE	Do Not Resterilize
2	Do Not Reuse	i	Consult Instructions for Use
#	Keep Dry	*	Keep Away from Sunlight
RX Only	For Prescription Use Only	<u> </u>	Caution, Consult Accompanying Documents

IFU-008 Rev-C (Draft) Page 4 of 4