

USE AND MAINTENANCE HANDBOOK FOR
**DIN 13260-2 TERMINAL UNITS / PROBES AND
 ACCESSORIES**
 CONFORM TO EN ISO 9170-1 STANDARD



93/42/EEC

1. General information

1.1 Introduction

Pres Block DIN 13260-2 terminal units, probes and accessories are medical devices "CE" marked, built to comply with 93/42/EEC European directive, EN ISO 9170-1 and DIN 13260-2 standards.

Before using the device, please read this handbook and ensure it is read by authorized operators.

1.2 Destination of use

Pres Block terminal units, probes and accessories have been designed to connect in a safe way medical equipment to the medical gases distribution system.

1.3 Classification and information under 93/42/EEC directive

The medical device:

- is identified with class of risk IIb, according to rule 11 of Annex IX of directive 93/42/EEC;
- is neither for single use, nor sterile;
- does not intentionally contain latex;
- does not incorporate neither medicinal substances, nor biological or animal tissues, nor human blood;
- does not contain phthalates classified as carcinogenic, mutagenic or toxic to reproduction.

1.4 Markings

Markings on every device are:

"PB" or "Pres Block"	Identifies Pres Block as the manufacturer
"CE0476"	Identifies the Notified Body that approved the CE certification for the device
Symbol of the gas distributed	according to EN ISO 9170-1 standard

Batch Numbers and other markings:

Sockets	
Referenced standard	
Batch number: either as a data stamp [XYYY], followed by one or two letters identifying the gas [ZZ] or as "[LOT] XYYYZZ". In both cases "XX" stands for the year of manufacture	
Name of the Customer	(optional)
Base Blocks: Please refer to handbook FTGTD05Ze	
Probes and Adaptors	
"[LOT] XYYYZZ"	Batch number, where XX identifies the year of manufacture
Referenced standard	(optional)

2. Technical information

2.1 Components

Available components are:

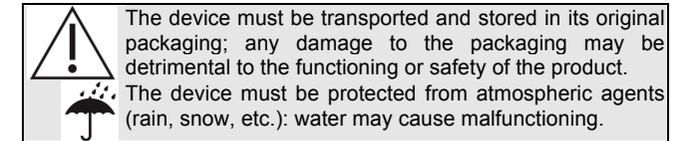
Name	Code (initial part)	Description
BASE BLOCK	ZARMOR	Part connected to the distribution system. Please refer to handbook FTGTD05Ze
SOCKET	VGTD VGTD	Part connected to the base block by a gas-specific interface. Contains the gas-specific connection point with the probe.
PROBE	IGTD / IGTGD	Male component, which is connected into and retained by the socket. Without check valve.
	IIGTD / IIGTGD	Male component which is connected into and retained by the socket. With check valve.
ADAPTOR	AGTD	Component for special applications.
SET	AGTD	Set made up of several components intended as a single device.

2.2 Technical and Operating characteristics

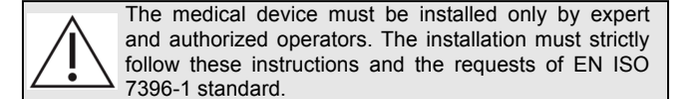
In compliance with the following standards:	DIN 13260-2 and EN ISO 9170-1	
Transport, storage and working temperatures range:		-20 ÷ +60° C
Transport, storage and working humidity range:		10 ÷ 90 %
Atmospheric pressure limits:		700 ÷ 1060 hPa
Materials used:	Brass (type CW614N) with and without nickel plating, Zamak (Zinc alloy), AISI (302/304/420), ABS, PA66, PE, POM, EPDM	
Lubricants:	Compatible with distributed gases (especially for O ₂)	
Distributed gases and relative working pressures:	O ₂ ; N ₂ O; CO ₂ ; Air, mixtures	from 320 kPa to 600 kPa
	Vacuum	from -10 kPa to -60 kPa
	N ₂ ; Air-800	from 560 kPa to 1200 kPa

2.3 Transport and storage

The original packaging is designed to guarantee the proper transport and storage of the product, in compliance with the requests of EN ISO 15001 standard.



3. Installation



The medical device must be installed at an easily accessible height, reducing as much as possible the risk of damages, due to furniture or other medical devices, and granting a distance ≥ 100 mm between 2 consecutive terminal units.

All the materials used during installation must be compatible with the gases supplied and the materials of the device. Every precaution must be taken to guarantee and maintain the cleanliness of the device according to EN ISO 15001 standard. Before installation, verify that the device is compatible with the devices it must be connected with, especially if supplied by different manufacturers.

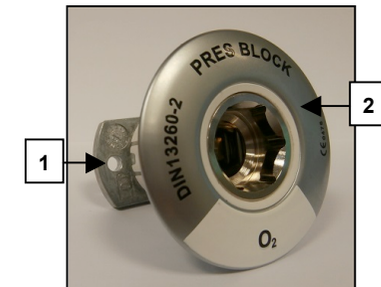
Before use, connect the device to equipotential earth.

3.1 Base blocks

Please refer to handbook FTGTD05Ze

3.2 Sockets

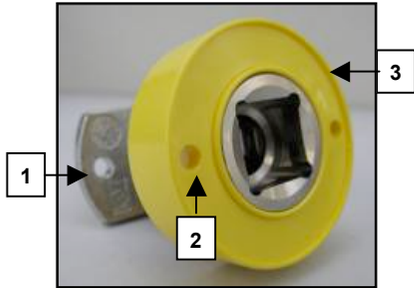
VGTD-OR (release sleeve supplied not assembled)



Perform the following operations in succession:

- align the 2 fixing holes I/D 4,5 mm (1) with the 2 females M4 threaded of the base block;
- fix the socket onto the base block tightening at the same time the 2 M4 screws (supplied with the base block) with an Allen wrench;
- fix the release sleeve (2) onto the socket;
- Turn the release sleeve (2) in the right position, it is possible turn 90° in 90°;
- apply the red label "NOT OPERATIVE", taking it away only when all final tests of the system have been performed.

VGTD-OR (release sleeve supplied assembled)



Perform the following operations in succession:

- align the 2 fixing holes I/D 4,5 mm (1) with the 2 females M4 threaded of the base block;
- fix the socket onto the base block tightening at the same time the 2 M4 screws (supplied with the base block) with an Allen wrench, through the 2 holes I/D 4,5 mm (2) of the release sleeve;
- apply the gas identification disc / label onto the release sleeve (3);
- apply the red label "NOT OPERATIVE", taking it away only when all final tests of the system have been performed.

3.3 Probes

Probe with barb end terminal IGT...PG... and IIGT...PG...

Perform the following operations in succession:

- fix the barb end to the flexible hose (I/D 6 or 8 mm) with a suitable single-use clip or crimped ferrule, according to EN ISO 5359 standard.

Probe with threaded terminal IGT...F... and IIGT...F...

Perform the following operations in succession:

- fix the threaded terminal granting the tightness with a suitable O-ring or PTFE tape.

3.4 Adaptors

Perform the following operations in succession:

- fix the adaptor to the panel using suitable screws;
- fix the barb end to the flexible hose (I/D 6 mm) with a suitable single-use clip or crimped ferrule, according to EN ISO 5359 standard.



Once installation is completed, verify the tightness: no leakages are allowed.
Afterwards testing required by EN ISO 7396-1 standard must be run.

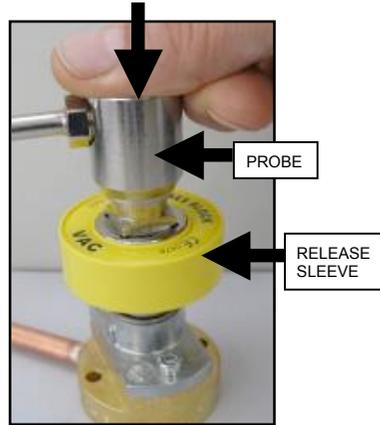
4. Use

4.1 Functioning

SOCKET / PROBE CONNECTION

1. Verify that the socket and the probe are designed for the same gas and that they refer to the same standard;
2. Grasp the probe and turn it aligning its profile with the corresponding profile of the socket;
3. Insert the probe into the socket with enough force to oppose the upstream pressure, until the release sleeve of the socket clicks into place;
4. Release the probe verifying that it remains anchored into the

socket.



SOCKET / PROBE DISCONNECTION

1. Keep the probe, in order to prevent its quick projection;
2. Push the release sleeve of the socket, towards the base block.



4.2 Cleaning

For cleaning use only distilled water or diluted ethyl alcohol.

4.3 Safety

Before use, personnel must familiarize with the control devices and their operations.

If the medical device is out of order or in maintenance, disconnect the gas supply and place a visible sign stating "NOT IN SERVICE. DO NOT USE".



Do not use the medical device for gases or pressures different from those for which it is designed.
Do not crash the probe, in order not to spoil its functionality.

Do not try to insert probes designed for a different gas and/or different standard into the sockets. This may damage both devices.

5. Maintenance

The maintenance program must envisage a minimum of biannual inspections with particular reference to:

- easy coupling and disconnection;

- wear or damage;
 - contamination;
 - marking – labelling;
 - tightness;
 - flow rate according to EN ISO 7396-1.
- These operations must be recorded.

The following maintenance kits are available for the sockets:

- VGTD-OR-RIC (for all gases except VAC);
- VGTD-OR-V-RIC (only for VAC).



Every precaution must be taken to maintain the cleanliness according to the requests of EN ISO 15001 standard.

During maintenance and repair only original parts must be used.

Do not use lubricants that aren't compatible with the distributed gases at their operating pressures. Especially in the case of contact with oxygen there might be safety issues relating to fire or explosion, if an incompatible lubricant is used.

Once maintenance is completed, testing required by EN ISO 7396-1 standard must be run again



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