

## TO WHOM IT MAY CONCERN

To any governmental departments,  
registration and/or trade offices in MOLDOVA

### Distribution Authorisation Letter

This letter confirms that **Sanmedico**  
**Mun. Chisinau**  
**Str. Petricani 88/1 of. 10**  
**Republica MOLDOVA**

is the **legal, exclusive and sole** representative of **TECO Medical Instruments Production + Trading GmbH, Dieselstr. 1, 84088 Neufahrn NB, Germany**, for the territory of **MOLDOVA** only for all TECO products listed below. **Sanmedico** may participate in public and private tenders, providing sales to all TECO customers in the territory. We as manufacturer certify that our warranty is duly passed to the purchaser through **Sanmedico** for the price, delivery schedules and the specifications of the published literature, catalogues and fully covering the commodities offered.

**Sanmedico** will provide the following information to TECO GmbH when so required in relation to its market surveillance activities:

Reporting of incidents to TECO must take place within 3 working days  
Serial number of the device, exact location of the device and the user.

Validity: January 1<sup>st</sup>, 2023 to December 31<sup>st</sup>, 2024


Termination: Confirmation ends automatically on Dec. 31<sup>st</sup> of 2024  
and must be then renewed.

#### Products:

- Coatron M1 Semi-automated 1-channel Coagulometer (out of production)
  - Coatron M2 Semi-automated 2-channel Coagulometer (out of production)
  - Coatron X Eco Semi-automated 1-channel Coagulometer
  - Coatron X Pro Semi-automated 2-channel Coagulometer
  - Coatron X Top Semi-automated 4-channel Coagulometer
  - Coatron A4 Fully automated Coagulometer, 4 optic channels
  - Coatron A6 Fully automated Coagulometer, 6 optic channels
  - Coatron A6 plus Fully automated Coagulometer, 6 optic channels
  - Hemostasis Reagents Complete product line
- all instruments with complete accessory, consumables and spare parts

This document is signed in Neufahrn, Germany, on January 18<sup>th</sup>, 2023

TECO Medical Instruments Production+Trading GmbH

  
Christian Hoetzel



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# Certificate of Approval

This is to certify that the Management System of:

## **TECO Medical Instruments, Production + Trading GmbH**

Dieselstr. 1, 84088 Neufahrn, Germany

has been approved by LRQA to the following standards:

**ISO 13485:2016**

Approval number(s): ISO 13485 – 00038268

**The scope of this approval is applicable to:**

Design, development, manufacturing, storage and sales of coagulation instruments and in-vitro-diagnostic reagents used in the hemostaseology and coagulation.



**Paul Graaf**

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Area Operations Manager, Europe

Issued by: LRQA Limited



0001



# KONFORMITÄTSERKLÄRUNG DECLARATION OF CONFORMITY

Doc#200/08-2022

Hersteller / Manufacturer: **TECO Medical Instruments  
Production + Trading GmbH**  
 Adresse / Address: **Dieselstrasse 1, 84088 Neufahrn, Germany**  
 Marktakteur / Actor ID SRN: **DE-MF-000022642 <https://ec.europa.eu>**

Wir erklären hier für die im Anhang A ( Seite 2 – 23 IVD Produkte) spezifizierten Produkte dass sie gemäß der Richtlinie für In-vitro-Diagnostika Medizinprodukte 98/79/EC klassifiziert sind als allgemeine IVD.

Diese Konformitätserklärung wird unter der alleinigen Verantwortung des Herstellers i.V.m. Artikel 110 Abs.3 und Abs.4 der Verordnung (EU) 2017/746 und des § 8 Abs.1 des Medizinprodukte-Durchführungsgesetzes, in der jeweils geltenden Fassung, ausgestellt.

Im Falle eigenmächtiger Veränderungen am Produkt oder der nicht bestimmungsgemäßen Verwendung verliert diese Erklärung ihre Gültigkeit.

We declare herewith for the products specified in Annex A ( page 2 - 23 IVD products) that they are classified as general IVD according to the In Vitro Diagnostic Medical Devices Directive 98/79/EC.

This declaration of conformity is issued under the sole responsibility of the manufacturer in according to article 110 para.3 and para.4 of Regulation (EU) 217/746 and section 8 para.1 of the Medical Device Law Implementing Act.

In case of unauthorised modifications to the products or un-intended use, this declaration loses its validity.

|  |   |
|--|---|
| Sie entsprechen den anwendbaren Anforderungen der Richtlinie:  | They meet applicable requirements of:   |
| Richtlinie 98/79/EG über In-vitro-Diagnostika klassifiziert gemäß Artikel 9 als "alle anderen Produkte"  | Directive 98/79/EC on in-vitro-diagnostic medical devices classified according to article 9 as „all other products“                                   |
| Die Qualitätssicherung entspricht den Anforderungen der Richtlinie 98/79/EG über In-vitro-Diagnostika für diese Art von Produkten.                   | The Quality Assurance is in accordance with the requirements of Directive 98/79/EC on in-vitro-diagnostic medical devices for those kind of products. |
| Der implementierte QM-Prozess entspricht der EN ISO 13485:2021   | The implemented QM Process complies with EN ISO 13485:2021  |
| Die vorstehende Konformitätserklärung ist gültig für alle Chargen dieser Produkte, die nach dem Datum der Unterzeichnung in Verkehr gebracht wurden. | The above mentioned declaration of conformity is valid for all lots of this product, which are distributed after the date of signature.               |
| Das Konformitätsbewertungsverfahren entspricht Anhang III der Richtlinie 98/79/EG über In-vitro-Diagnostika für diese Art von Produkten.             | The conformity assessment procedure complies with Annex III of Directive 98/79/EC on in-vitro-diagnostic medical devices for those kind of products.  |

Ort und Datum der Unterzeichnung: **Neufahrn, 2022-08-31**  
Place and date of issue:



Christian Hötzi  
Verantwortliche Person / PRRC

Doc#200/08-2022

## KONFORMITÄTSERKLÄRUNG – DECLARATION OF CONFORMITY

Directive 98/79/EC Annex A  
 Übrige Produkte – Reagenzien für In-vitro-Diagnostika  
 Other products – Reagents for in vitro diagnostic – general IVD

| Pos. | Article No | Tradename           | Unit                                    | Generic Device Term                        | EMDN / GMDN Code<br>EUDAMED DI                |
|------|------------|---------------------|---|--|---|
| 1    | A0230-040  | TEClot PT-S (Quick) | 10x4ml PT-S                             | Prothrombin time ( quick test )            | W0103020101 / 30539<br>B-PTS-A0230-040X7      |
| 2    | A0230-100  | TEClot PT-S (Quick) | 10x10ml PT-S                            | Prothrombin time ( quick test )            | W0103020101 / 30539<br>B-PTS-A0230-100WY      |
| 3    | A0260-050  | TEClot PT-B (Owren) | 5x10ml PT-B                             | Prothrombin time ( quick test )            | W0103020199 / 55986<br>B-PTB-A0260-050G2      |
| 4    | A0320-050  | TEClot APTT-S       | 10x5ml APTT-S                           | Activated partial thromboplastin time      | W0103020102 / 55982<br>B-APTT-A0320-050AM     |
| 5    | A0401-020  | TEClot TT           | 10x2ml TT                               | Thrombin time / reptilase / batroxbin time | W0103020103 / 55988<br>B-TT-A0401-0207P       |
| 6    | A0511-020  | TEClot FIB          | 10x2ml FIB                              | Fibrinogen assays (factor i)               | W0103020201 / 55997<br>B-FIB-A0511-020N2      |
| 7    | A0511-050  | TEClot FIB          | 10x5ml FIB                              | Fibrinogen assays (factor i)               | W0103020201 / 55997<br>B-FIB-A0511-050NB      |
| 8    | C1010-020  | TEChrom AT          | 6x6ml reagent FXa<br>3x3 ml substrate   | Antithrombin                               | W0103020602 / 56156<br>B-AT-C1010-020HL       |
| 9    | D2010-012  | Red D-Dimer         | 3x4ml latex<br>3x7ml reaction buffer    | D-Dimer                                    | W0103020503 / 47349<br>B-DD-D2010-0126W       |
| 10   | D2020-005  | Blue D-Dimer LC     | 1x5ml latex LC<br>1x7ml reaction buffer | D-Dimer                                    | W0103020503 / 47349<br>B-DD-D2020-0057E       |
| 11   | P8001-010  | TECal N             | 10x1ml                                  | Calibration plasma for haemostasis         | W0103020701 / 45786<br>B-CAL-P8001-005X8      |
| 12   | P8200-005  | TECal DD            | 5x1ml                                   | Calibration plasma for haemostasis         | W0103020701 / 47348<br>B-CAL-P8200-005XX      |
| 13   | P6001-010  | TEControl N         | 10x1ml                                  | Control plasma for haemostasis             | W0103020702 / 30590<br>B-CTRL-P6001-010H7     |
| 14   | P6101-010  | TEControl A         | 10x1ml                                  | Control plasma for haemostasis             | W0103020702 / 30590<br>B-CTRL-P6101-010HQ     |
| 15   | P6201-010  | TEControl A Plus    | 10x1ml                                  | Control plasma for haemostasis             | W0103020702 / 30590<br>B-CTRL-P6201-010J9     |
| 16   | P5001-010  | TEClot Factor II    | 10x1ml                                  | Coagulation factor ii ( prothrombin )      | W0103020202 / 30542<br>B-FAC-II-P5001-010ML   |
| 17   | P5101-010  | TEClot Factor V     | 10x1ml                                  | Coagulation factor v                       | W0103020204 / 30544<br>B-FAC-V-P5101-010AN    |
| 18   | P5201-010  | TEClot Factor VII   | 10x1ml                                  | Coagulation factor vii                     | W0103020205 / 30545<br>B-FAC-VII-P5201-0107B  |
| 19   | P5301-010  | TEClot Factor VIII  | 10x1ml                                  | Coagulation factor viii                    | W0103020207 / 30547<br>B-FAC-VIII-P5301-01097 |
| 20   | P5401-010  | TEClot Factor IX    | 10x1ml                                  | Coagulation factor ix                      | W0103020208 / 30548<br>B-FAC-IX-P5401-0106C   |
| 21   | P5501-010  | TEClot Factor X     | 10x1ml                                  | Coagulation factor x                       | W0103020209 / 30549<br>B-FAC-X-P5501-010EQ    |
| 22   | P5601-010  | TEClot Factor XI    | 10x1ml                                  | Coagulation factor xi                      | W0103020210 / 30551<br>B-FAC-XI-P5601-010A8   |
| 23   | P5701-010  | TEClot Factor XII   | 10x1ml                                  | Coagulation factor xii                     | W0103020211 / 30552<br>B-FAC-XII-P5701-010CJ  |

(Recital 23 of Directive 98/79/EC on In Vitro Diagnostics Medical Devices) - Annex A - general IVD



# KONFORMITÄTSERKLÄRUNG DECLARATION OF CONFORMITY

Doc#001-01/06-2022

Hersteller / Manufacturer: **TECO Medical Instruments  
Production and Trading GmbH**  
Adresse / Address: **Dieselstrasse 1, 84088 Neufahrn, Germany**  
Marktakteur / Actor ID SRN: **DE-MF-000022642 <https://ec.europa.eu>**

Die hier benannten Produkte der generischen Produktgruppe erfüllen die Anforderungen der aufgeführten Verordnungen, Richtlinien und Normen. Im Falle eigenmächtiger Veränderungen am Produkt oder der nicht bestimmungsgemäßen Verwendung verliert diese Erklärung ihre Gültigkeit.

Diese Konformitätserklärung wird unter der alleinigen Verantwortung des Herstellers ausgestellt.

## **BASIS UDI-DI 426018278CMX81152**

IVD - halb-automatische Blutgerinnungsmessgeräte - Handelsbezeichnung, Typ, Kat.-Nr.  
IVD - semi-automated Coagulation Systems - trade name, type, model, Cat.-No.

## **Coatron X Eco / Coatron X Pro / Coatron X Top**

**81 101 10**

**81 101 20**

**81 101 40**

The products of the generic product group named here fulfil the requirements of listed regulations, directives and standards. In the case of unauthorised modifications to the product or use not in accordance with the intended purpose, this declaration becomes invalid.

This declaration of conformity is issued under the sole responsibility of the manufacturer.

### **Verordnung (EU) 2017/746**

für in-vitro Diagnostika-IVDR  
und dem harmonisierten Standard am 2022-05-12:  
Risikoklassifizierung gemäß Artikel 47–Anhang VIII  
**Regel 5 b – „Klasse A“**

Konformitätsbewertungsverfahren gemäß:  
(EU) 2017/746 Artikel 17 (Anhang II+III)

Angewandte Normen zur Sicherstellung der  
grundlegenden Anforderungen an Leistung und  
Sicherheit:

EN ISO 18113-3:2011  
DIN EN 62304:2018  
DIN EN 62366-1  
DIN EN 62366-1:2017  
DIN EN 61326-1:2013  
DIN EN 55011:2009 + A1:2010  
IEC 61010-1:2010, AMD1:2016  
IEC 61010-2-101:2015  
IEC 61010-1:2010

Richtlinie 2011/65/EU RoHS III  
(incl. (EU) 2015/863) - DIN EN IEC 63000

QM-System gemäß (EU) 2017/746 Art.10(8)  
angewandter Standard: EN ISO 13485:2021

### **Regulation (EU) 2017/746**

for In-vitro diagnostic medical devices  
and it's harmonized standard at 2022-05-12:  
Risk classified according to article 47 annex VIII  
**Rule 5 b – "Class A"**

Conformity assessment procedure in accordance with:  
(EU) 2017/746 Article 17 (annex II+III)

Standards applied to ensure the essential requirements  
for performance and safety:

EN ISO 18113-3:2011  
DIN EN 62304:2018  
DIN EN 62366-1  
DIN EN 62366-1:2017  
DIN EN 61326-1:2013  
DIN EN 55011:2009 + A1:2010  
IEC 61010-1:2010, AMD1:2016  
IEC 61010-2-101:2015  
IEC 61010-1:2010

Directive 2011/65/EU RoHS III  
(incl. (EU) 2015/863 - DIN EN IEC 63000

QM-Systems in accordance with (EU) 2017/746 art.10(8)  
Applied standard procedure: EN ISO 13485:2021

Ort und Datum der Unterzeichnung: Neufahrn, 2022-06-21  
Place and date of issue:

Matthias Dieckmann  
General Manager



Christian Hötzel  
Verantwortliche Person / PRRC



# KONFORMITÄTSERKLÄRUNG

## DECLARATION OF CONFORMITY

Doc#100/07-2021

Wir / We

### TECO Medical Instruments Production and Trading GmbH

Name des Herstellers / Manufacturer's name

**Dieselstrasse 1, 84088 Neufahrn, Germany**

Anschrift / Address

erklären in alleiniger Verantwortung, dass die unten gelisteten IVD Zubehör Produkte:  
*declare under our own responsibility, that the IVD accessories products, listed below:*

|  |                           |
|--|---------------------------|
| Doppelküvette / <i>Double cuvette</i>                    | Ref. 19 000 02            |
| Einzelküvette / <i>Single cuvette</i>                    | Ref. 20 000 02, 24 100 00 |
| 4-fach Küvette / <i>Cuvette 4 pos/ea</i>                 | Ref. 80 521 10            |
| 6-fach Küvette / <i>Cuvette 6 pos/ea</i>                 | Ref. 80 560 00            |
| 6-fach Küvette (micro) / <i>Cuvette 6 pos/ea (micro)</i> | Ref. 80 570 00            |

allen anwendbaren Anforderungen folgender Richtlinien entsprechen: *meet all applicable requirements of:*

1. Richtlinie 98/79/EG über In-vitro Diagnostika und ihrem Zubehör, klassifiziert gemäß Artikel 9 als: "alle anderen Produkte"- im Sinne von Zubehör zu In vitro Diagnostika gemäß Artikel 1.

*1. Directive 98/79/EC on In-vitro diagnostic medical devices and their accessories, classified according to article 9 as: "all other products" – and in term of accessories for in vitro diagnostics according to article 1.*

2. Richtlinie 2011/65/EU (RoHS III)

*2. Directive 2011/65/EU (RoHS III)*

Das QM-System des Herstellers ist zertifiziert nach:

*The QM-system of the manufacturer is certified for:*

**EN ISO 13485:2016**

***EN ISO 13485:2016***

Konformitätsbewertungsverfahren gemäß:

*Conformity assessment procedure according to:*

Gemäß Anhang III der Richtlinie 98/79/EG

*According to Annex III of Directive 98/79/EC*

Ort und Datum der Unterzeichnung:  
Place and date of issue:

Neufahrn, 27.07.2021  
Neufahrn, July 27, 2021

Matthias Dieckmann  
General Manager



# TECO

MEDICAL INSTRUMENTS  
PRODUCTION+TRADING GMBH

Dieselstraße 1  
D-84088 Neufahrn N.B.  
fon: +49-8773/707 80-0  
fax: +49-8773/707 80-29

Neufahrn, 26/04/2018

## TO WHOM IT MAY CONCERN

We confirm that the instruments Coatron X Eco, Coatron X Pro and Coatron X Top have a closed cuvette system. Cuvettes have to be purchased with voucher identification code from TECO GmbH.



Christian Hoetzl  
General Manager  
TECO Germany



Quality Management

We are certified

Voluntary participation in regular monitoring according to ISO 9001:2008



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Dieselstraße 1

D-84088 Neufahrn N.B.

fon: +49-8773/707 80-0

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# CERTIFICATE

for: **Mr. Vitalie Goreacii**

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Company: **Sanmedico SRL**  
Str. Petricani 88/1, oficiul 10  
Chisinau - Rep. Moldava MD-2059  
MOLDOVA

have participated with success at the intensive training session:

**Application and technical training for following instruments:**

- **Coatron X series**
  - **Installation**
  - **Application**
  - **General use, also in combination with TECAM Software**
  - **Technical and After Sales Service**

Supervisors: **Mr. Chr. Hoetzi and Mrs. Wendy Guo**

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Place of Training: **TECO – Germany**

Date: **November 18<sup>th</sup>, 2019**



Christian Hoetzi  
General Manager





TOP  
INNOVATION  
2017 - 2018

Clotting  
Chromogenic  
Immunturbidimetric

# Coatron

Semi-automated  
Coagulation Analyzer Series

With 1, 2 or 4 optical channels



# TECO

Innovation in Coagulation

# A new area of manual and semi-automated Coagulation Analyser rise up

The Coatron X instrument line is a consequent continuation in the development of the Coatron product line. Over 25 years in experience and innovation is the reference for our new Coatron X instrumentation line.

The unique detection principle in combination with the high-level analytical algorithm calculates exact, precise and reproducible results.

Easy in operation – self instructing user dialogue - reliable

## **Highest optical resolution, enlarged optic range, smallest sample and reagent volume**

0.1 mOD, 0 - 3800 mOD, just with 75 µL sample and reagent volume

## **Complete optical analysis**

No further parts required, like balls, stirrers etc.

## **Adaptation of the light level**

Automatic light level adjustment of the optic channels to each sample

## **Exclusion of disturbance**

Stray light reduction, exact temperature control, all parameter are preset

“Complete range of Coagulation Analysis with the highest standard and reliability. The new generation of Coagulation instruments with optical detection are here.”

Coatron X - product family



With 1, 2 or 4 optical channels.

[www.teco-medical.com](http://www.teco-medical.com)

## Prepared for the daily routine and the upcoming requirements

### One instrument – many possibilities

The Coatron X family is prepared to work with one, two or four channels. The built-up and functionality is specifically designed to each instrument version and requirements. The operation with the intuitive user dialogue and handling of the detection results are easy and effective. The possibility to connect the instrument to the **TECO Cloud** offers new perspective of instrument, reagent and consumables verification and handling. The precise and correct patient result is what we want to secure.



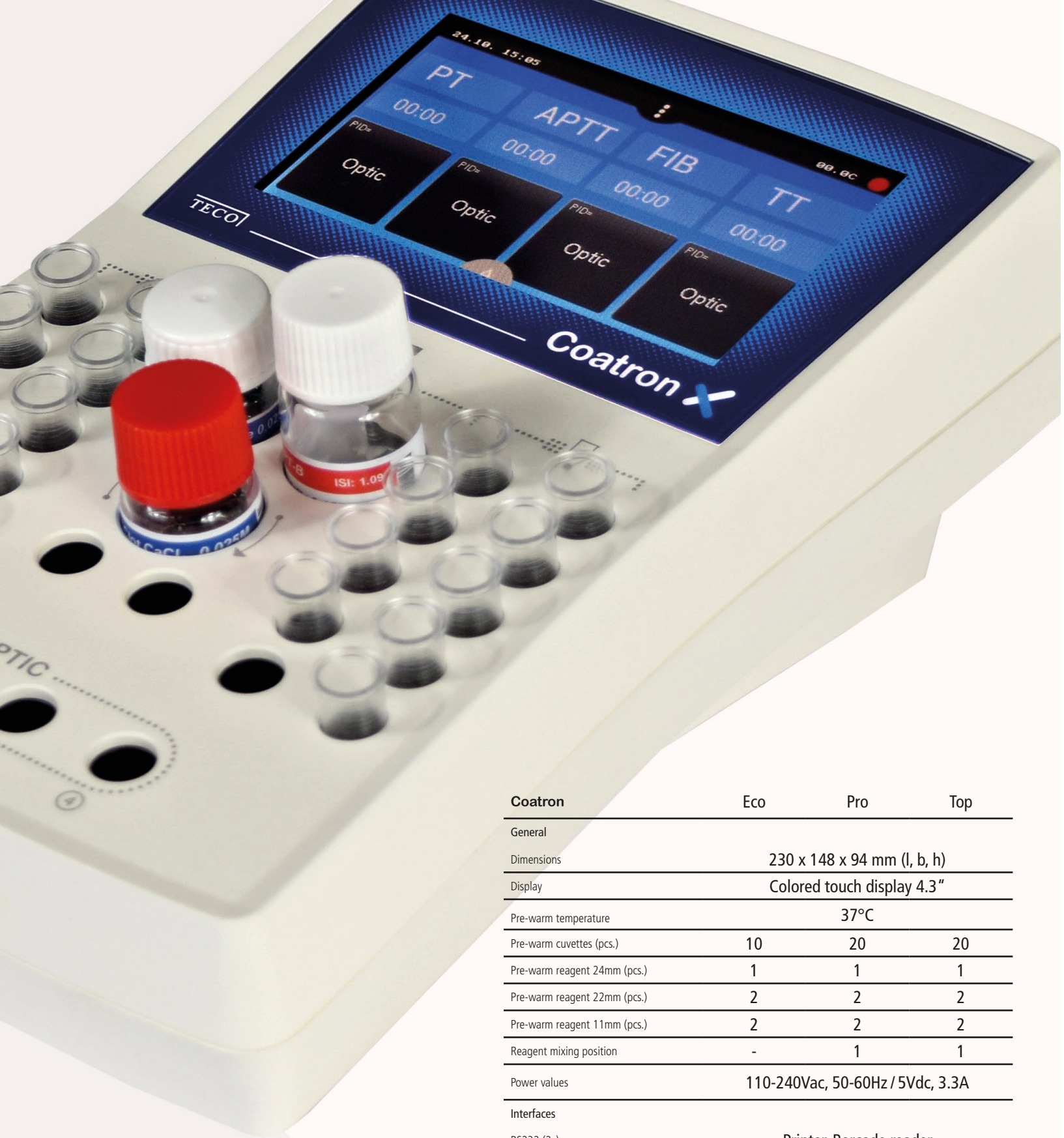
### Quality is our basic demand

TECO develop and produce with qualified and specialized companies, located in Germany. High reliability, nearly maintenance free instruments are our benefit. Our reference is 25 years, in worldwide laboratories, with satisfied users.



### TECO Cloud Services – A strong data bank and application service behind

All instrument versions of the Coatron X family are connectable via Bluetooth to Smart-devices, like mobile devices, tablets, etc. with a specific APP or direct access to the TECO Cloud Services.



| Coatron                      | Eco                              | Pro                | Top      |
|------------------------------|----------------------------------|--------------------|----------|
| General                      |                                  |                    |          |
| Dimensions                   | 230 x 148 x 94 mm (l, b, h)      |                    |          |
| Display                      | Colored touch display 4.3"       |                    |          |
| Pre-warm temperature         | 37°C                             |                    |          |
| Pre-warm cuvettes (pcs.)     | 10                               | 20                 | 20       |
| Pre-warm reagent 24mm (pcs.) | 1                                | 1                  | 1        |
| Pre-warm reagent 22mm (pcs.) | 2                                | 2                  | 2        |
| Pre-warm reagent 11mm (pcs.) | 2                                | 2                  | 2        |
| Reagent mixing position      | -                                | 1                  | 1        |
| Power values                 | 110-240Vac, 50-60Hz / 5Vdc, 3.3A |                    |          |
| Interfaces                   |                                  |                    |          |
| RS232 (2x)                   | Printer, Barcode reader          |                    |          |
| USB (2x)                     | Network, Firmware update         |                    |          |
| Bluetooth                    | TECO Cloud, App                  |                    |          |
| Optic / tests                |                                  |                    |          |
| Optic channels               | 1                                | 2                  | 4        |
| Wavelength (nm)              | 620 (red)                        | 405 (UV)           | 405 (UV) |
| Global Coag. tests           | PT, APTT, TT, FIB                |                    |          |
| Specific Coag. tests         | -                                | individual factors |          |
| Chromogenic Coag. tests      | -                                | AT, PC             |          |
| Latex based tests            | D-Dimer                          |                    |          |
| Whole blood tests            | PT-B                             | -                  | -        |



## The details make the difference

### Coatron X

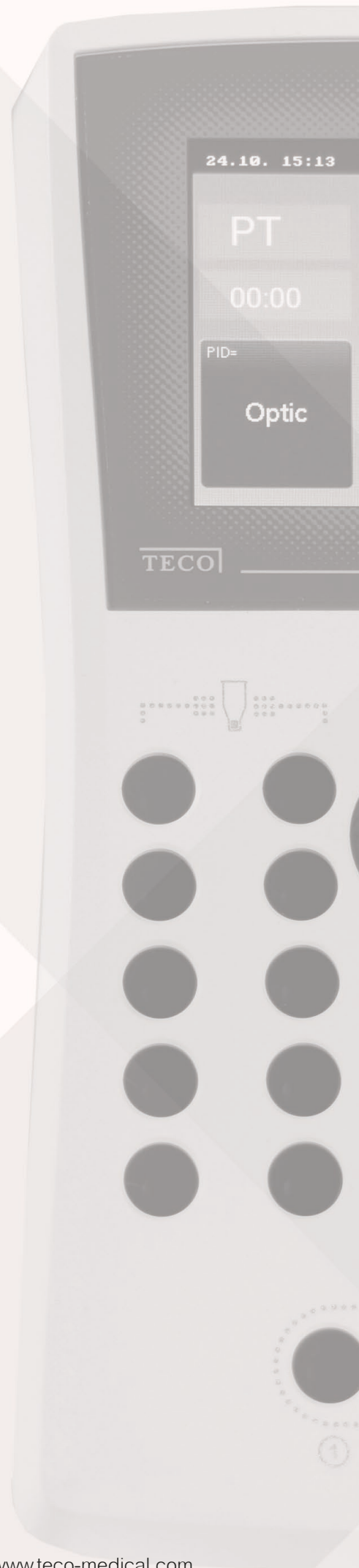
The remarkable details in every single component is achieved by selecting of premium suppliers.

The performance of a high level instrument is strongly depending on the concept in general and the perfect usability to reach the requirements of a modern laboratory analyser.

Priority No. 1 was to get a daily routine reliability and easy-to-use operation.

### Software and connection possibilities

With the Coatron X product family starts a new time line in analysis management and service maintenance. Operation via intuitive, colored touchscreen, as well patient result management are perfectly optimized.



## Operation details

| Coatron   | Eco | Pro      | Top |
|---|-----|----------|-----|
| <b>Operation</b>  |     |          |     |
| Touchscreen 4.3"  | ✓   | ✓        | ✓   |
| Real time clock   | ✓   | ✓        | ✓   |
| Stopwatch   | ✓   | ✓        | ✓   |
| Language selection                                      | ✓   | ✓        | ✓   |
| <b>Interfaces</b>                                       |     |          |     |
| USB to LIS  | ✓   | ✓        | ✓   |
| Network to LIS<br>(TECAM software required)             | ✓   | ✓        | ✓   |
| <b>Management</b>                                       |     |          |     |
| Test calibration  | ✓   | ✓        | ✓   |
| Tracking to Pat.ID, Patient ID,<br>Sample ID or Auto ID | ✓   | ✓        | ✓   |
| Automatic optic start<br>(no Starterpipette required)   | ✓   | ✓        | ✓   |
| Double determination                                    | ✗   | ✓        | ✓   |
| Sample management (ID)                                  | ✗   | ✓        | ✓   |
| Reagent management (ID)<br>(lot und expiry)             | ✗   | ✓        | ✓   |
| Internal result databank                                | ✗   | ✓        | ✓   |
| Patient identification with barcode                     |     | optional |     |



### Intuitive operation and control

Clear and easy to operate user dialogue with a high quality colored touchscreen

- Direct usable
- Short learning phase
- Logic, intuitive operation
- Reliable touchscreen surface
- Quick touch response



### For small and mediate laboratory requirements

Concept is suitable for daily routine work in Coagulation laboratories and hospitals

- Three different versions available, depending on number of samples per day
- In maximum up to 4 optic channels available

### Interfaces

#### RS232 (2x)

- For external serial printer and external barcodereader

#### LIS/USB

#### Bluetooth



Integrated barcode scan for reagents.





### **TECO Cloud Services**

#### **A strong data bank and application service behind**

All instrument versions of the Coatron X family are connectable via Bluetooth to Smart-devices, like mobile devices, tablets, etc. with a specific APP or direct access to the TECO Cloud Services.



For trading partners worldwide, please visit our web-page

**TECO Medical Instruments Production + Trading GmbH**  
Dieselstr. 1, 84088 Neufahrn, Germany  
Tel.: +49 (0) 8773 70780-0, Fax +49 (0) 8773 70780-29  
info@teco-gmbh.com, www.teco-medical.com

**TECO**  
Innovation in Coagulation



**Utilizare**

Acest produs este utilizat pentru determinarea timpului de protrombină (PT) în plasmă, conform Quick<sup>1,2</sup>. Testul este sensibil la activitatea factorilor de coagulare implicați pe calea “extrinsecă”, și anume: II, V, VII, X și fibrinogen și prin urmare este utilizat pentru terapie anticoagulantă orală cu inhibitori ai Vitaminei-K precum Warfarin sau Marcumar. De asemenea, se utilizează și pentru determinarea cantitativă a factorilor de coagulare implicați pe calea “extrinsecă”. PT-ul măsoară timpul de coagulare extrinsecă (activarea factorului VII) a plazmei de testare după adăugarea reagentului PT.

**Continut și Determinări**

| Product       | TEClot PT-S | TEClot PT-S | TEClot PT-S |
|---------------|-------------|-------------|-------------|
| Cat.No.       | A0230-010   | A0230-040   | A0230-100   |
| PT-S Reagent* | 5x2 mL      | 10x4 mL     | 10x10 mL    |

**Determinări**

|             |          |          |           |
|-------------|----------|----------|-----------|
| Coatron M** | 200 Det. | 800 Det. | 2000 Det. |
| Coatron A4  | 100 Det. | 400 Det. | 1000 Det. |
| Coatron A6  | 200 Det. | 800 Det. | 2000 Det. |

\*conține un extract din creier de iepure cu Buffer, stabilizatori și Clorură de Calciu.

\*\*Metoda micro (75μl în total)

**Preparare**

Reconstituiți cu apă de înaltă puritate cu volumul indicat pe eticheta flaconului

|           |           |           |
|-----------|-----------|-----------|
| A0230-010 | A0230-040 | A0230-100 |
| 2 mL      | 4 mL      | 10 mL     |

Lăsați la temperatura camerei pentru cel puțin 15 minute cu învârtiri/rotiri ocazionale. Plasați reagentul în aparat și lăsați să se incubeze pentru încă 15 minute. Reactivul sedimentează și trebuie învârtit/rotit înainte de fiecare testare. La aparatele Coatron puteți utiliza o bară de amestecare pentru asta.

**Păstrare și Stabilitate**

Reagenții nedesfăcuți și păstrați între 2<sup>o</sup>-8<sup>o</sup>C, sunt stabili pînă la data expirării indicate pe ambalaj. Reagenții deschiși deja, sunt stabili :

|            |                                  |                      |                   |
|------------|----------------------------------|----------------------|-------------------|
|            | 2 <sup>o</sup> -8 <sup>o</sup> C | 20-25 <sup>o</sup> C | 37 <sup>o</sup> C |
| PT reagent | 5 zile                           | 36 ore               | 8 ore             |

### Măsuri de precauție

Evitați contactul cu pielea și ochii. Purtați haine de protecție adecvate. Eliminați componentele în conformitate cu reglementările locale pentru materialele infecțioase. Toate componentele sunt verificate pentru HIV, VHB, VHC. Cu toate acestea, produsele din sângele uman ar trebui considerate potențial infecțioase.

### Colectarea și depozitarea probei <sup>4</sup>

1. Obțineți sângele venos prin puncție curată a venei.
2. Amestecați imediat 9 părți de sânge cu 1 parte de 3,2% de Citrat de Na (0.105M) și amestecați bine.
3. Centrifugați specimenul la 1500g pentru 10 min (trombocite < 10000/ $\mu$ L).
4. Separați plazma după centrifugare și depozitați în tuburi de sticlă siliconizată.
5. Utilizați plazma în timp de 4 ore sau înghețați și dezghețați doar înainte de utilizare.

Stabilitatea plazmei: 4 ore la 18-26°C, 8 ore la 2-8°, 30 zile la -20°C, 6 luni la -70°C

### Procedura

#### A. Metoda automată: Coatron A

| Prothrombin Time |            | A4          |     | A6         |     |            | A4     | A6 |        | A4       | A6 |
|------------------|------------|-------------|-----|------------|-----|------------|--------|----|--------|----------|----|
| PAT              | Patient    | 50 $\mu$ l  | CP1 | 25 $\mu$ l | CP1 | Incubation | 0s     |    | SENS   | 2        |    |
| BUF              | IBS Buffer | 0 $\mu$ l   | P39 | 0 $\mu$ l  | P79 | Maxtime    | 120s   |    | POINTS | 4        |    |
| CLR              | -          | 0 $\mu$ l   | -   | 0 $\mu$ l  | -   | Unit       | 251    |    | MIX    | No       |    |
| DP               | -          | 0 $\mu$ l   | P00 | 0 $\mu$ l  | P00 | Method     | Coag   |    | Clean  | 0        | 0  |
| R0               | -          | 0 $\mu$ l   | P00 | 0 $\mu$ l  | P00 | Math       | log XY |    | Multi  | 1        | 3  |
| R1               | -          | 0 $\mu$ l   | P00 | 0 $\mu$ l  | P00 | CT-Mech    | No     |    | S-Corr | 0%       |    |
| R2               | PT Reagent | 100 $\mu$ l | P25 | 50 $\mu$ l | P46 | Deadtime   | 7s     |    | T-Corr | 30% - 4s |    |

#### B. Metoda manuală:

1. Incubați reagentul PT la 37°C pentru cel puțin 10 minute
2. Pipetați **25  $\mu$ l** din specimen în cuveta pentru testare. Incubați la 37°C pentru 1-2 min
3. Adăugați **50  $\mu$ l** de reagent PT (37°C) și începeți simultan testul.
4. Înregistrați/măsurați în secunde timpul de coagulare.

### Rezultate așteptate

Secunde tipice : 11-18 sec

Intervalul normal: 70 - 130% 0.85 – 1.15 INR

Oricum, rezultatele sunt influențate de aparate, tehnică, calibrare etc. Se recomandă ca fiecare laborator să-și stabilească propriul interval, specific aparatelor utilizate.

## Standardizare și Calibrare

Rezultatul PT este exprimat în secunde sau activitate (% Quick) sau INR (International Normalised Ratio).

Rezultatele INR:

-au fost calculate din timpul normal și valoarea ISI (international sensitivity index/ indicele internațional de sensibilitate). Primul este obținut prin testarea plazmei proaspete a unui grup de indivizi sănătoși. Valoarea ISI este stabilită în certificatul de analiză specific fiecărui lot.

$$INR = \left( \frac{\text{Patient PT}}{\text{Normal PT}} \right)^{ISI}$$

Rezultatele % activității (Quick):

-au fost calculate dintr-o curbă de calibrare, care a fost pregătită după plasma de referință (e.g. **TECAL N**) și diluări în soluții saline ca 0.9% de NaCl2 sau TECLOT IBS buffer. Sunt recomandate cel puțin 3 sau mai multe puncte de calibrare . Curba de calibrare trebuie confirmată cu plasma de control cu interval normal și patologic.

| <b>% din normal diluat în soluție salină</b> | <b>100%*</b> | <b>50%</b> | <b>25%</b> | <b>12,5%**</b> |
|--|--------------|------------|------------|----------------|
|  | Fără diluție | 1+1        | 1+3        | 1+7            |

\*media de la cel puțin 21 de indivizi sănătoși este definită ca 100%<sup>5</sup>

\*\*diluția de 12.5% poate cauza rezultate “+++” în unele cazuri, pentru că nivelul de diluție al fibrinogenului este prea mare pentru detecție optică.

## Controlul Calității

TEControl sau altă plasmă de control comercială trebuie utilizată pentru controlul calității performanței la o frecvență în conformitate cu practicile bune de laborator (GLP). TEControl poate fi înghețat o singură dată după reconstituire. 120-150 μl păstrat închis în tuburi de polipropilen la -20°C , este stabil timp de 30 zile.

## Restricții/ Limitări

O mare grijă și atenție trebuie de avut la factorii care pot părea ne semnificativi.

### A. Colectarea probei. Evitați:

1. Folosiți doar tuburi de plastic sau sticlă siliconizată.
2. Amestecarea întârziată a sîngelui cu anticoagulant.
3. Contaminarea cu tromboplastină tisulară.
4. Raportul greșit de sînge cu anticoagulant.
5. Probele hemolizate, icterice sau lipemice pot interfera sistemele optice.

### B. Tehnici de Laborator:

1. Efectuați testul la 37°C.
2. Utilizați doar apă cu puritate înaltă.
3. pH-ul optim este 7.0-7.5.
4. Valoarea ISI nu e constantă în primile 30 min după reconstituire.
5. Reagentul face sedimente , de aceea trebuie de agitat înainte de fiecare testare.

## Caracteristici de performanță

Performanțe tipice pe aparatul Coatron M4:

| Precision:       | CV% (în timpul testării) | CV% (între testări) |
|------------------|--------------------------|---------------------|
| Normal control   | < 3,0                    | < 5,0               |
| Abnormal control | < 3,0                    | < 5,0               |













## Garantie

Acest produs este garantat doar dacă se utilizează în conformitate cu informațiile de pe eticheta proprie și din instrucțiune. TECO nu acordă garanție dacă produsul este utilizat în alt scop și în niciun caz TECO nu va fi răspunzător pentru daunele care rezultă din garanția expresă menționată mai sus.

## Referințe:

1. Quick, A.J., The Hemorrhagic Diseases and the Physiology of Hemostasis. Charles C. Thomas: Springfield, IL. 1942.
2. Quick, A.J., Hemorrhagic Diseases. Lea and Febiger: Philadelphia. 1957.
3. Miale, J.B., Laboratory Medicine-Hematology, 4th Edition. C.V. Mosby: St. Louis. 1972.
4. National Committee for Clinical Laboratory Standards: Guidelines for the Standardized Collection, Transport and Preparation of Blood Specimens for Coagulation Testing and Performance of Coagulation Assays.
5. Besselaar A M H P van den, Lewis SM, Mannucci P n Poller L. 1993. Status of present and candidate International Reference Preparations (IRP) of thromboplastin for prothrombin time. Thromb Hemostas 69; 85
6. Besselaar A M H P van den. 1991. The significance of the International Normalized Ratio (INR) for oral anticoagulant therapy. H17CC 3; 146153.

Symbol keys

|  |  |   |  |   |  |
|--|--|---|--|---|--|
|  Expiry date    |  In Vitro Diagnostica |  Biological hazard |  Catalogue Number |  Reconstitute with dest. water |  Consult accompanying documents |
|  Store at 2-8°C |  EU conformity        |  Manufacturer      |  Lot. Number      |  Ready to use                  |  Authorized Representative      |



IVD

REF

A0501-010, A0501-025, A0511-020, A0511-050

**Utilizare**

TECLOT FIB este utilizat pentru determinarea cantitativă a fibrinogenului în plasma umană, conform tehnicii dezvoltată de Clauss<sup>1</sup>. Nivelul de fibrinogen poate crește ca rezultat al inflamației, sarcinii sau utilizarea contraceptivelor orale<sup>2</sup>. Nivelul scăzut de fibrinogen poate fi identificat în situații concrete precum boli de ficat sau DIC (Coagulare intravasculară diseminată). Deficiențele congenitale includ afibrinogenaemia (nici un fibrinogen detectabil), hypofibrinogenaemia (<1 mg/ml) și dysfibrinogenaemia (molecula anormală de fibrinogen).

**Continut și preparare**

| Product               | TECLOT FIB Kit-10 | TECLOT FIB Kit-25 | TECLOT FIB | TECLOT FIB |
|-----------------------|-------------------|-------------------|------------|------------|
| Cat.No.               | A0501-010         | A0501-025         | A0511-020  | A0511-050  |
| Reactivul de trombină | 5x2 mL            | 5x5 mL            | 10x2 mL    | 10x5 mL    |
| IBS Buffer            | 1x125 mL          | 1x125 mL          | -          | -          |
| TECal Normal          | 1x1 mL            | 1x1 mL            | -          | -          |
| TEControl A           | 1x1 mL            | 1x1 mL            | -          | -          |

**Determinări**

|            |          |           |          |           |
|------------|----------|-----------|----------|-----------|
| Coatron M* | 400 Det. | 1000 Det. | 800 Det. | 2000 Det. |
| Coatron A4 | 200 Det. | 500 Det.  | 400 Det. | 1000 Det. |
| Coatron A6 | 200 Det. | 500 Det.  | 400 Det. | 1000      |

\*Metoda micro (75μL în total)

1. Reactivul de trombină:  
 Conține trombină de bovină (~80NIH) cu stabilizatori.  
 REF: A0501-010/A0511-020: reconstituiți cu 2 ml de apă purificată.  
 REF: A0501-025/A0511-050: reconstituiți cu 5 ml de apă purificată.
2. IBS Buffer: Gata de utilizare. Conține soluție salină tamponată Imidazol.
3. TECal Normal: reconstituiți cu 1 ml de apă purificată. Conține plasma umană citrată.
4. TEControl A: reconstituiți cu 1 ml de apă purificată. Conține plasma umană citrată.



Rotiți ușor după reconstituire și lăsați timp de 15 minute la temperatura camerei. Amestecați bine înainte de utilizare. Nu agitați/ scuturați.

**Păstrare și stabilitate**

Reagenții care nu au fost deschiși sunt stabili pînă la data expirării indicate pe ambalaj, păstrați la temperatura de 2°-8°C. Reagenții deschiși deja se păstrează:

|                        |         |          |         |
|------------------------|---------|----------|---------|
| Reactivul de trombină* | 2-8 °C  | 15-25 °C | 37 °C   |
|                        | 12 zile | 5 zile   | 24 ore  |
| TEControl sau Plasma   | 2-8 °C  | 15-25 °C | -20 °C  |
|                        | 8 ore   | 4 ore    | 30 zile |

\*Reactivul trebuie de protejat razele ultraviolete și evaporare.

### **Măsuri de precauție**

Evitați contactul cu pielea și ochii. Purtați haine de protecție adecvate. Eliminați componentele în conformitate cu reglementările locale pentru materialele infecțioase. Toate componentele sunt verificate pentru HIV, HBV, HCV. Cu toate acestea, produsele din sângele uman ar trebui considerate potențial infecțioase.

### **Colectarea și păstrarea probelor**<sup>3</sup>

1. Obțineți sânge venos printr-o puncție curată a venei.
2. Amestecați imediat 9 părți de sânge cu 1 parte de 3.2% sodium citrate (0.105M) și amestecați bine.
3. Centrifugați proba la 1500g timp de 10 minute (trombocite < 10000/μL).
4. Separați plasma după centrifugare și păstrați în plastic sau tuburi de sticlă siliconată.
5. Folosiți plasma în timp de 4 ore sau depozitați/păstrați înghețat și dezghețați numai înainte de folosire.

### **Procedura**

#### A. Metoda automată Coatron A

| Fibrinogen |            | A4   |     | A6   |     |            | A4     | A6 |        | A4 | A6 |
|------------|------------|------|-----|------|-----|------------|--------|----|--------|----|----|
| PAT        | Patient    | 10μl | CP1 | 10μl | CP1 | Incubation | 0s     |    | SENS   | 0  |    |
| BUF        | IBS Buffer | 90μl | P39 | 90μl | P79 | Maxtime    | 120s   |    | POINTS | 4  |    |
| CLR        | -          | 0μl  | -   | 0μl  | -   | Unit       | 769    |    | MIX    | No |    |
| DP         | -          | 0μl  | P00 | 0μl  | P00 | Method     | Coag   |    | Clean  | 1  | 3  |
| R0         | -          | 0μl  | P00 | 0μl  | P00 | Math       | log XY |    | Multi  | 1  | 1  |
| R1         | -          | 0μl  | P00 | 0μl  | P00 | CT-Mech    | Yes    |    | S-Corr | 0% |    |
| R2         | Fibrinogen | 50μl | P29 | 50μl | P49 | Deadtime   | 3s     |    | T-Corr | 0% |    |

#### B. Metoda manuală Coatron M

1. Prepararea diluțiilor pentru Standard, Control și Pacient.

| Diluția Standard    | Plazma         | IBS Buffer |
|---------------------|----------------|------------|
| 1:5                 | 200μL Standard | 800μL      |
| 1:10                | 500μL 1:5 STD  | 500μL      |
| 1:20                | 500μL 1:10 STD | 500μL      |
| 1:40                | 500μL 1:20 STD | 500μL      |
| Pacient sau Control | 100μL Plasma   | 900μL      |

2. Pipetați 50 μl de standard diluat sau plasma pacientului (1:10) într-o cuvetă pentru testare. Preîncălziți la 37°C timp de 1-2 minute.
3. Adăugați 25 μl de reactiv de trombină și simultan începeți testul.  
Pentru alte aparate, consultați manualul cu instrucțiuni specifice mai detaliate.

### **Calibrare**

Ca referință trebuie de utilizat TECal Normal sau altă Plasma standard comercială preparată în care fibrinogenul a fost determinat. (200-300mg/dL). Reprezintă grafic timpul de coagulare obținut cu fiecare din diluțiile standard ale fibrinogenului pe axa-y , opus concentrației de fibrinogen (mg/dL) pe axa-x utilizând hîrtia grafică log-log. Linia cea mai potrivită trebuie de determinat prin analiza regresiei liniare. Fibrinogenul din probele de plasmă poate fi determinat prin interpolare din curba de calibrare.

### **Rezultate aseptate**

Rezultatele normale tipice sunt 180-450 mg/dL<sup>4,5</sup>. Oricum , rezultatele sunt influențate de metoda de detecție a coagulării și poate varia de la laborator la laborator. Este recomandat ca fiecare laborator să-și stabilească propriul interval normal specific aparatului utilizat.

### **Controlul Calității**

Pentru un control de încredere al calității performanței , trebuie utilizată TEControl sau altă plasmă de control comercială la o frecvență în conformitate cu practicile bune de laborator (GLP). TEControl poate fi înghețat o singură dată după reconstituire. 120-150 μl păstrat închis în tuburi de polipropilen la -20°C , este stabil timp de 30 zile.

### **Limitări / Restricții**

A. Colectarea probei.EVITAȚI:

1. Folosiți doar tuburi de plastic sau sticlă siliconizată.
2. Întîrzierea amestecării sîngelui cu anticoagulant.
3. Contaminarea cu tromboplastină tisulară.
4. Raportul greșit dintre sînge și anticoagulant.
5. Probele hemolizate, icterice sau lipemice pot interfera sistemele optice.

B . Tehnici de Laborator:

1. Efectuați testul la 37°C.
2. Utilizați doar apă cu puritate înaltă.
3. pH-ul optim este 7.0-7.5.

### **Caracteristici de performanță:**

| Precizia         | CV%(în timpul testării) | CV%(între testări) |
|------------------|-------------------------|--------------------|
| Normal control   | < 5.0                   | < 5.0              |
| Abnormal control | < 5.0                   | < 10.0             |

(Performanțe tipice pentru aparatul Coatron M4)


### **Garantie**

Acest produs este garantat doar dacă se utilizează în conformitate cu informațiile de pe eticheta proprie și din instrucțiune. TECO nu acordă garanție dacă produsul este utilizat în alt scop și în niciun caz TECO nu va fi răspunzător pentru daunele care rezultă din garanția expresă menționată mai sus.

## Referinte

1. Clauss, A., Gerinnungsphysiologische Schnellmethode zur bestimmung des Fibrinogens. Acta Haematol., 1957, 17: 237-246.
2. Shaw, T.S., Assays for Fibrinogen and its Derivatives, CRC Crit. Rev. Clin. Lab. Sci., 1977, 8: 145-192.
3. National Committee for the National Laboratory (NCCLS) Standards: Collection transport and preparation of blood specimens for coagulation testing and performance of coagulation assays. Document H21-A2, vol. 11, No. 23, 1991.
4. Scully, R.E. et al., Normal Reference Laboratory Values, N. Eng. J. Med., 1980, 302(37) : 37-48.
5. Okuno, T. and Selenko, V., Amer. J. Med. Tech., 1972, 38(6) : 196-201.

Symbols key:

|  |  |   |   |  |
|--|--|---|---|--|
|  Expiry date    |  In Vitro Diagnostica |  Biological hazard |  Catalogue Number |  Consult accompanying documents |
|  Store at 2-8°C |  EU conformity        |  Manufacturer      |  Lot. Number      |  Authorized Representative      |





IVD

REF

A0590-125

### Intended Use

The IBS Buffer solution is optimally formulated for use on Coagulation Analyzers. Use in accordance with the recommended Operators Manuals for installing and replacing Owrens Veronal Buffer (OVB). The IBS can be used as the diluent for preparing plasma dilutions in the performance of Fibrinogen determinations and Coagulation Factor Assays with all manual, mechanical, or photo-optical means of clot detection. Follow Reagent manufacturer's recommended procedures for preparation of plasma dilutions using Imidazole Buffered Saline.

### Contents & Determinations

|            |            |
|------------|------------|
| Product    | IBS Buffer |
| Cat.No.    | A0590-125  |
| IBS Buffer | 1x125 mL   |

### Preparation

IBS: pH 7.3 - 7.4, liquid  
Ready to use.

### Storage and Stability

Unopened reagents are stable until the expiration date shown on the label stored at 2-8°C.

### Precautions

Avoid contact with skin and eyes. Wear suitable protective clothing. Dispose components in compliance with local regulations for infectious material.

### Warranty

This product is warranted to perform in accordance with its labelling and literature. TECO disclaims any implied warranty of merchantability or fitness for any other purpose, and in no event will TECO be liable for any consequential damages arising out of aforesaid express warranty.

Symbols key:

|                |                      |                   |                  |                                |
|----------------|----------------------|-------------------|------------------|--------------------------------|
| Expiry date    | In Vitro Diagnostica | Biological hazard | Catalogue Number | Consult accompanying documents |
| Store at 2-8°C | EU conformity        | Manufacturer      | Lot. Number      | Authorized Representative      |



A0590-125

### Verwendungszweck

Die IBS Pufferlösung (Imidazole Buffered Saline) wird für die Verdünnung von Plasma verwendet werden, wie es z.B. bei der koagulometrischen Bestimmung von Fibrinogen, Einzelfaktoren oder auch Verdünnungsreihen für die Methoden Kalibrierung notwendig ist.

### Inhalte und Bestimmungen

|            |            |
|------------|------------|
| Produkt    | IBS Puffer |
| Kat.Nr.    | A0590-125  |
| IBS Buffer | 1x125 mL   |

### Vorbereitung

IBS: pH 7.3 - 7.4, flüssig  
Gebrauchsfertig

### Lagerung und Stabilität

Ungeöffnete Reagenzien sind bei Lagerung zwischen 2-8°C bis zum auf dem Etikett angegebenen Verfallsdatum haltbar.

### Vorsichtsmaßnahmen

Haut- und Augenkontakt vermeiden. Angemessene Schutzkleidung tragen. Bestandteile gemäß lokaler Vorschriften für infektiöse Materialien entsorgen.

### Garantie

Es wird garantiert, dass die Wirkungsweise dieses Produktes den Angaben auf der Packung und in der Produktliteratur entspricht. TECO haftet weder für die Verkäuflichkeit oder Eignung dieses Produktes für irgendwelche andere Zwecke noch für irgendwelche Folgeschäden, die sich aus der vorstehenden, expliziten Garantie ergeben.

Erklärung der Symbole:

|                  |                     |                    |                |                         |
|------------------|---------------------|--------------------|----------------|-------------------------|
| Verfallsdatum    | In-Vitro Diagnostik | Biologische Gefahr | Katalog-Nummer | Begleitpapiere beachten |
| Bei 2-8°C lagern | EU Konformität      | Hersteller         | Lot. - Nummer  | Bevollmächtigter        |



IVD

REF

P6001-010

**Intended Use**

Use as a normal control for following coagulation tests:

**PT, APTT, Thrombin time, Fibrinogen,  
Anti-thrombin and D-Dimer**

**Contents**

10 x 1mL freeze dried citrate-anticoagulated human plasma

**Preparation**

Reconstitute individual vials with **1,0 ml** distilled water. Allow to stand at room temperature, with occasional swirling, for 15 min before use. Be certain all particulate matter is well dissolved.

PT whole blood (TEClot PT-B): Reconstitute individual vials with **1,7 ml** distilled water.

**Storage & Stability**

Unopened vials are stable until the expiration date shown on the label stored at 2°-8°C.

Dissolved plasma change analytic levels below 10% if stored as following:

|         |         |          |
|---------|---------|----------|
| -20 °C  | 2-8 °C  | 20-25 °C |
| 1 month | 8 hours | 4 hours  |

Dissolved plasma can be refrozen only one time in aliquots (120-150µL). Stored at -20°C in closed polypropylene tubes, the aliquots must be used within 30 days.

**Precautions**

This product contains substance from human origin!  
Avoid contact with skin and eyes. Wear suitable protective clothing. Dispose components in compliance with local regulations for infectious material. All components are checked for HIV, HBV and HCV. However products from human blood should be considered as potentially infectious.

**Expected Results**

Refer to "Certificate of Analysis".

**Warranty**

This product is warranted to perform in accordance with its labelling and literature. TECO disclaims any implied warranty of merchantability or fitness for any other purpose, and in no event will TECO be liable for any consequential damages arising out of aforesaid express warranty.

Symbols key:

|                |                      |                   |                  |                                |
|----------------|----------------------|-------------------|------------------|--------------------------------|
| Expiry date    | In Vitro Diagnostica | Biological hazard | Catalogue Number | Consult accompanying documents |
| Store at 2-8°C | EU conformity        | Manufacturer      | Lot. Number      | Authorized Representative      |





## Verwendungszweck

Als normale Kontrolle für folgende Gerinnungstests verwenden:

**PT, APTT, Thrombinzeit, Fibrinogen,  
Antiithrombin und D-Dimer**

## Inhalt

10 x 1mL gefriergetrocknetes mit Zitrat versetztes gerinnungshemmendes Humanplasma

## Vorbereitung

Die einzelnen Fläschchen mit 1,0ml destilliertem Wasser anlösen. Fläschchen bei Raumtemperatur bis zur Anwendung unter gelegentlichen Verwirbeln 15 Minuten lang stehen lassen. Stellen Sie sicher, dass alle Partikel gut aufgelöst sind.

Vollblut PT (TEClot PT-B): einzelne Fläschchen mit 1,7ml destilliertem Wasser anlösen.

## Lagerung und Stabilität

Ungeöffnete Fläschchen sind bei Lagerung zwischen 2-8°C zum bis auf dem Etikett angegebenen Verfallsdatum haltbar.

Gelöstes Plasma verändern die analytischen Levels unter 10% wenn wie folgt gelagert:

|         |           |           |
|---------|-----------|-----------|
| -20 °C  | 2-8 °C    | 20-25 °C  |
| 1 Monat | 8 Stunden | 4 Stunden |

Gelöstes Plasma kann einmalig wiedereingefroren werden. Die Aliquots (120-150µL) sind 30 Tage haltbar, wenn sie in polypropylen Gefäßen bei -20°C aufbewahrt werden.

## Vorsichtsmaßnahmen

Dieses Produkt enthält Substanzen humanen Ursprungs! Haut- und Augenkontakt vermeiden. Angemessene Schutzkleidung tragen. Abfälle laut lokaler Regelungen für infektiöse Materialien entsorgen. Alle Bestandteile wurden auf HIV, HBV und HCV getestet. Trotzdem müssen Produkte aus menschlichem Blut immer als potentiell infektiös angesehen werden.

## Erwartete Ergebnisse

Lesen Sie das Analysenzertifikat

## Garantie

Es wird garantiert, dass die Wirkungsweise dieses Produkts den Angaben auf der Packung und in der Produktliteratur entspricht. TECO haftet weder für die Verkäuflichkeit oder Eignung dieses Produktes für irgendwelche andere Zwecke noch für irgendwelche Folgeschäden, die sich aus der vorstehenden, expliziten Garantie ergeben.

Erklärung der Symbole:

|                  |                     |                    |                |                         |
|------------------|---------------------|--------------------|----------------|-------------------------|
| Verfallsdatum    | In-Vitro Diagnostik | Biologische Gefahr | Katalog-Nummer | Begleitpapiere beachten |
| Bei 2-8°C lagern | EU Konformität      | Hersteller         | Lot. - Nummer  | Bevollmächtigter        |





IVD

REF

P6101-010

**Intended Use**

Use as an abnormal control for following coagulation tests:

**PT, APTT, Thrombin time, Fibrinogen,  
Antithrombin and D-Dimer**

**Contents**

10 x 1mL freeze dried citrate-anticoagulated human plasma

**Preparation**

Reconstitute individual vials with **1,0 ml** distilled water. Allow to stand at room temperature, with occasional swirling, for 15 min before use. Be certain all particulate matter is well dissolved.

PT whole blood (TEClot PT-B): Reconstitute individual vials with **1,7 ml** distilled water.

**Storage & Stability**

Unopened vials are stable until the expiration date shown on the label stored at 2°-8°C.

Dissolved plasma change analytic levels below 10% if stored as following:

| -20 °C  | 2-8 °C  | 20-25 °C |
|---------|---------|----------|
| 1 month | 8 hours | 4 hours  |

Dissolved plasma can be refrozen only one time in aliquots (120-150µL). Stored at -20°C in closed polypropylene tubes, the aliquots must be used within 30 days.

**Precautions**

This product contains substance from human origin!  
Avoid contact with skin and eyes. Wear suitable protective clothing. Dispose components in compliance with local regulations for infectious material. All components are checked for HIV, HBV and HCV. However products from human blood should be considered as potentially infectious.

**Expected Results**

Refer to "Certificate of Analysis".

**Warranty**

This product is warranted to perform in accordance with its labelling and literature. TECO disclaims any implied warranty of merchantability or fitness for any other purpose, and in no event will TECO be liable for any consequential damages arising out of aforesaid express warranty.

Symbols key:

|                |                      |                   |                  |                                |
|----------------|----------------------|-------------------|------------------|--------------------------------|
| Expiry date    | In Vitro Diagnostica | Biological hazard | Catalogue Number | Consult accompanying documents |
| Store at 2-8°C | EU conformity        | Manufacturer      | Lot. Number      | Authorized Representative      |



**Verwendungszweck**

Als abnormale Kontrolle für folgende Gerinnungstests verwenden:

**PT, APTT, Thrombinzeit, Fibrinogen,  
Antithrombin und D-Dimer**

**Inhalt**

10 x 1mL gefriergetrocknetes mit Zitrat versetztes gerinnungshemmendes Humanplasma

**Vorbereitung**

Die einzelnen Fläschchen mit 1,0ml destilliertem Wasser anlösen. Fläschchen bei Raumtemperatur bis zur Anwendung unter gelegentlichen Verwirbeln 15 Minuten lang stehen lassen. Stellen Sie sicher, dass alle Partikel gut aufgelöst sind.

Vollblut PT (TEClot PT-B): einzelne Fläschchen mit 1,7ml destilliertem Wasser anlösen.

**Lagerung und Stabilität**

Ungeöffnete Fläschchen sind bei Lagerung zwischen 2-8°C zum bis auf dem Etikett angegebenen Verfallsdatum haltbar.

Gelöstes Plasma verändern die analytischen Levels unter 10% wenn wie folgt gelagert:

|         |           |           |
|---------|-----------|-----------|
| -20 °C  | 2-8 °C    | 20-25 °C  |
| 1 Monat | 8 Stunden | 4 Stunden |

Gelöstes Plasma kann einmalig wiedereingefroren werden. Die Aliquots (120-150µL) sind 30 Tage haltbar, wenn sie in polypropylen Gefäßen bei -20°C aufbewahrt werden.

**Vorsichtsmaßnahmen**

Dieses Produkt enthält Substanzen humanen Ursprungs! Haut- und Augenkontakt vermeiden. Angemessene Schutzkleidung tragen. Abfälle laut lokaler Regelungen für infektiöse Materialien entsorgen. Alle Bestandteile wurden auf HIV, HBV und HCV getestet. Trotzdem müssen Produkte aus menschlichem Blut immer als potentiell infektiös angesehen werden.











**Erwartete Ergebnisse**

Lesen Sie das Analysenzertifikat

**Garantie**

Es wird garantiert, dass die Wirkungsweise dieses Produkts den Angaben auf der Packung und in der Produktliteratur entspricht. TECO haftet weder für die Verkäuflichkeit oder Eignung dieses Produktes für irgendwelche andere Zwecke noch für irgendwelche Folgeschäden, die sich aus der vorstehenden, expliziten Garantie ergeben.

Erklärung der Symbole:

|  |   |  |  |   |
|--|---|--|--|---|
|  Verfallsdatum    |  In-Vitro Diagnostik |  Biologische Gefahr |  Katalog-Nummer |  Begleitpapiere beachten |
|  Bei 2-8°C lagern |  EU Konformität      |  Hersteller         |  Lot. - Nummer  |  Bevollmächtigter        |



IVD

REF

P8001-005

**Intended Use**

Use as a calibrator or normal control for following coagulation tests:

**PT, APTT, Thrombin time, Fibrinogen,  
Factors: II, V, VII, VIII, IX, X, XI, XII,  
Antithrombin, Protein-C, free Protein-S,  
D-Dimer**

**Contents**

5 x 1 mL freeze dried citrate-anticoagulated human plasma

**Preparation**

Reconstitute individual vials with **1,0 ml** distilled water. Allow to stand at room temperature, with occasional swirling, for 15 min before use. Be certain all particulate matter is well dissolved.

PT whole blood (TEClot PT-B CAT=A0260 xxx): Reconstitute individual vials with **1,7 ml** distilled water.

**Storage & Stability**

Unopened vials are stable until the expiration date shown on the label stored at 2°-8°C.

Dissolved plasma change analytic levels below 10% if stored as following:

|         |        |          |         |
|---------|--------|----------|---------|
| -20 °C  | 2-8 °C | 20-25 °C | 37°C    |
| 30 days | 24h    | 8h       | 2 hours |

Dissolved plasma can be refrozen only one time in aliquots (120-150µL). Stored at -20°C in closed polypropylene tubes, the aliquots must be used within 30 days.

**Precautions: Potential Biohazardous material**

This product contains substance from human origin! Avoid contact with skin and eyes. Wear suitable protective clothing. Dispose components in compliance with local regulations for infectious material. All components are checked for HIV, HBV and HCV. However products from human blood should be considered as potentially infectious.

**Performance Characteristics:**

Refer to "Certificate of Analysis".

**Limitations:**

The control plasma is subject to the limitations of the assay system (reagent + instrument). Results out of expected range may indicate deterioration, false test calibration or problems with one or more components of the test system

**Warranty**

This product is warranted to perform in accordance with its labelling and literature. TECO disclaims any implied warranty of merchantability or fitness for any other purpose, and in no event will TECO be liable for any consequential damages arising out of aforesaid express warranty.

Symbols key:

|                |                      |                   |                  |                                |
|----------------|----------------------|-------------------|------------------|--------------------------------|
| Expiry date    | In Vitro Diagnostica | Biological hazard | Catalogue Number | Consult accompanying documents |
| Store at 2-8°C | EU conformity        | Manufacturer      | Lot. Number      | Authorized Representative      |



### Verwendungszweck

Als Kalibrator oder Normalkontrolle für folgende Gerinnungstests verwenden:

**PT, APTT, Thrombinzeit, Fibrinogen,  
Faktoren: II, V, VII, VIII, IX, X, XI, XII,  
Antithrombin, Protein-C, freies Protein-S,  
D-Dimer**

### Inhalt

5 x 1mL gefriergetrocknetes mit Zitrat versetztes gerinnungshemmendes Humanplasma

### Vorbereitung

Die einzelnen Fläschchen mit 1,0ml destilliertem Wasser anlösen. Fläschchen bei Raumtemperatur bis zur Anwendung unter gelegentlichen Verwirbeln 15 Minuten lang stehen lassen. Stellen Sie sicher, dass alle Partikel gut aufgelöst sind.

Vollblut PT (TEClot PT-B CAT=A0260 xxx): einzelne Fläschchen mit 1,7ml destilliertem Wasser anlösen.

### Lagerung und Stabilität

Ungeöffnete Fläschchen sind bei Lagerung zwischen 2-8°C zum bis auf dem Etikett angegebenen Verfallsdatum haltbar.

Gelöstes Plasma verändern die analytischen Levels unter 10% wenn wie folgt gelagert:

|         |            |           |           |
|---------|------------|-----------|-----------|
| -20 °C  | 2-8 °C     | 20-25 °C  | 37°C      |
| 30 Tage | 24 Stunden | 8 Stunden | 2 Stunden |

Gelöstes Plasma kann einmalig wiedereingefroren werden. Die Aliquots (120-150µL) sind 30 Tage haltbar, wenn sie in polypropylen Gefäßen bei -20°C aufbewahrt werden.

### Vorsichtsmaßnahmen: Potentiell infektiöses Material

Dieses Produkt enthält Substanzen humanen Ursprungs! Haut- und Augenkontakt vermeiden. Angemessene Schutzkleidung tragen. Abfälle laut lokaler Regelungen für infektiöse Materialien entsorgen. Alle Bestandteile wurden auf HIV, HBV und HCV getestet. Trotzdem müssen Produkte aus menschlichem Blut immer als potentiell infektiös angesehen werden.

### Erwartete Ergebnisse

Lesen Sie das Analysenzertifikat

### Einschränkungen:

Das Kontrollplasma unterliegt den Einschränkungen der verwendeten Reagenzien und Geräte. Ergebnisse außerhalb des Sollbereichs können verursacht werden durch abgelaufene Materiale, ungültige Methodenkalibration oder Problemen an Reagenz, Gerät oder Zubehör.

### Garantie

Es wird garantiert, dass die Wirkungsweise dieses Produkts den Angaben auf der Packung und in der Produktliteratur entspricht. TECO haftet weder für die Veräußerlichkeit oder Eignung dieses Produktes für irgendwelche andere Zwecke noch für irgendwelche Folgeschäden, die sich aus der vorstehenden, expliziten Garantie ergeben.

Erklärung der Symbole:

|                  |                     |                    |                |                         |
|------------------|---------------------|--------------------|----------------|-------------------------|
| Verfallsdatum    | In-Vitro Diagnostik | Biologische Gefahr | Katalog-Nummer | Begleitpapiere beachten |
| Bei 2-8°C lagern | EU Konformität      | Hersteller         | Lot. - Nummer  | Bevollmächtigter        |