



### DECLARATION OF CONFORMITY

for CE – marking according to Annex III par. 3  
of Czech Government Act No. 56/2015 Coll. and EC Directive 98/79 on diagnostic medical devices  
in vitro

**Document number:** TF-V.01/9/2/11 ENG

**Manufacturer:** ITEST plus, s.r.o.

**Address:** ITEST plus, s.r.o.  
Kladská 1032/44c  
500 03 Hradec Králové  
Česká republika

**Product type:** **Diagnostic and Antimycotic Discs, Carbohydrates Diagnostic Tablets**

**Catalogue no.:** DD 501; DD 5011; DD 502; DD 5021; DD 503; DD 5031; DD 504;  
DD 505; DD 506; DD 5061; DD 507; DD 509; DD 511; DD 512; DD 5121;  
DD 513; DD 5131; DD 514; DD 5141; DD 515;

DD 601; DD 6011; DD 6012; DD 602; DD 6021; DD 6022; DD 603;  
DD 6031; DD 6032; DD 604; DD 6041; DD 605; DD 6051; DD 6052;  
DD 606; DD 6061; DD 6062; DD 607; DD 6071; DD 6072; DD 608;  
DD 6081; DD 6082; DD 609; DD 6091; DD 6092; DD 610; DD 6101;  
DD 6102; DD 611; DD 6111; DD 6112; DD 612; DD 6121; DD 6122;  
DD 613; DD 6131; DD 6132

DT 707, DT 711

SO 341, PP 61

#### Intended Use:

- The discs are used for fast and simple identification of microbial infectious diseases, based on biochemical characteristics of microbes.
- Antimycotic discs are determined for susceptible determination of microscopic fungi with antifungal preparates in diffusion method.

These products are for laboratory use only.



**Manufacturer hereby declares under his own responsibility that above listed products are in compliance with applicable legislature:**

- a) Czech Government Act No. 56/2015 Coll., on Technical requirements on diagnostic medical devices in vitro, as amended
- b) Directive of European Parliament no. 98/79/EC for diagnostic medical devices in vitro
- c) Czech Act no. 268/2014 Coll., on Diagnostic medical devices in vitro, and Amendment no. 90/2021 Coll., which changes the Act no. 268/2014 Coll., on medical devices and change of Act no. 634/2004, Coll., and Executive order no. 187/2021 Coll., which changes Executive order 62/2015 Coll., on implemetation of some regulations in Act on medical devices
- d) requirements of Annex I of Czech Government Act No. 56/2015 Coll., on Technical requirements on diagnostic medical device in vitro,

safe for intended use under standard conditions.

**References:**

ČSN EN ISO 13485 ed. 2 Změna A11:2022  
ČSN EN ISO 14971 ed. 2 Změna A11:2022  
ČSN EN ISO 15223-1: 2022  
ČSN EN 1041: 2009+A1: 2014  
ČSN EN 13612, opr.1: 2003  
ČSN EN ISO 20417: 2021  
ČSN EN ISO 14644-1:2019

**This Declaration was issued in compliance with Annex III of Czech Government Act No. 56/2015 Coll., on Technical requirements on diagnostic medical devices in vitro, as amended.**

**From 26. 05. 2022 the manufacturer will also meet the requirements of Regulation (EU) 2017/746 of the European Parliament and of the Council on in vitro diagnostic medical devices updated on 25. 01. 2022, according to art. 110, paragraphs (3) and (4) temporary provisions which define obligations for manufacturer to fulfill some specific requirements of IVDR 2017/746 as follows:**

- **post-market surveillance,**
- **vigilance,**
- **registration of economic operator**

**considering that the manufacturer is not allowed to implement any significant changes in design and/or intended use for above listed in vitro diagnostic medical devices after 26. 05. 2022.**

**Any unauthorised changes to this document, would make this declaration invalid.**

**Issue date:** 22. 06. 2022

**Name and Signature:** RNDr. Jiří Pospíšil, CSc.,  
Executive Chief Officer

**Place of issue:** Hradec Králové, Czech republic

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