



## EU DECLARATION OF CONFORMITY



### Identification of the declarant and manufacturer:

**Name** Andromedical S.L.  
**Address** Gran via 6, 28013 Madrid, Spain  
**VAT** ESB82545096  
**Email** [legal@andromedical.com](mailto:legal@andromedical.com)

### Product identification:

**Product description** Penile traction device (PTD)  
**Product name** Andropeyronie  
**Product models** N/A  
**Classification** Medical Device Class I  
**Type:** Non-sterile, Non-measurement, Non-reusable surgical instrument, non-invasive, non-incorporated medical substances, non-active, device.  
**Test-Standards** ISO 10993; EN71-3:2002  
**GMDN Code** 46340, Penile traction splint.  
**UMDNS Code** 23850, Penile tissue expanders, traction.

The Product to which this declaration on refers to, is manufactured according to the Medical Device Regulation (EU) 2017/745 on the approximation of the laws of Member States and is registered in Class I according to Annex VIII. Conformity is assured according to the guidelines set out in Annex X. The Product is produced in accordance with harmonized standards. The product carries the CE Mark. Notified body intervention is not required.

The manufacturer, hereby, bears full responsibility for the production in accordance with the requirements stated in this declaration.

Place and Date: Madrid, 10/10/2020

DocuSigned by:

*Eduardo Gómez de Diego*

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Dr. Eduardo Gómez de Diego

CEO & CRO

[www.andromedical.com](http://www.andromedical.com)

Phone-Fax: +34-911-981-740

Gran via 6  
28013 Madrid  
SPAIN