

Proof of Authority

To use the CE mark on pressure equipment on the basis of internal production control plus supervised pressure equipment checks at random intervals according to module A2 of Decree 44/2016. (XI.28.) NGM and Directive 2014/68/EU of the European Parliament and Council

Certificate no.: **H/Ü 23 0274**

Name and address of the manufacturer: **Celitron Medical Technologies Kft.
H-2600 Vác
Nádas utca 2.
Hungary**

The manufacturer is authorized on the basis of the results of the verification of the preconditions to use the CE mark along with the identification number of the notified body within the scope of the validity on the pressure equipment manufactured by the manufacturer as follows:

CE 1008

Scope of validity **ISS AC575 Chamber**

Report No.: **E 122/2067/2024**

Valid until: **12.11.2025.**

(First issue of the Certificate: 02.02.2023)

Budapest, 03.12.2024



Ferenc Székely
Certifier

TÜV Rheinland InterCert Kft.
Industrial Services Business Unit
NoBo Number: 1008

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MS-0047845 Melléklet-4

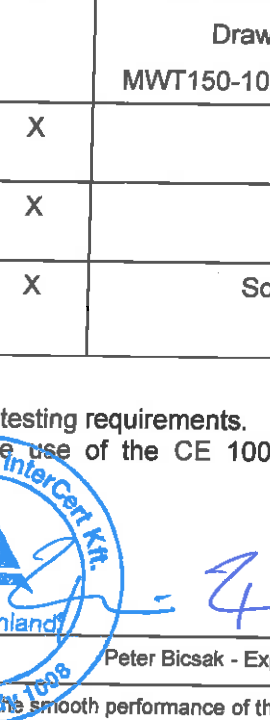
Rev.0

Supervision report

Supervised pressure equipment checks at random intervals



Identification number of the notified body:
1008

Report number		E122/2067/2024	
MODUL: A2	Internal production control plus supervised pressure equipment checks at random intervals	<input checked="" type="checkbox"/>	
C2	Conformity to type based on internal production control plus supervised pressure equipment checks at random intervals	<input type="checkbox"/>	
H1	Conformity based on full quality assurance plus design examination	<input type="checkbox"/>	
E	Acceptance of pressure equipment of the III and IV categories according to the diagrams 1, 2, 3, 4, 5, acc.to 38. § and article 4. item (1) and (2)	<input type="checkbox"/>	
D	Acceptance of pressure equipment of the III and IV categories according to the diagrams 1, 2, 3, 4, 5, acc.to 38. § and article 4. item (1) and (2).	<input type="checkbox"/>	
Manufacturer:/ Supplier: Celitron Medical Technologies Kft. Nádas utca 2. H-2600 Vác		Manufacturer's factory: Auth Impex Kft. Budapesti országút, hrsz.: 4032/32 H-7700 Mohács	
BASIS OF THE TEST:		Decree 44/2016 (XI.28.) NGM, Directive 2014/68/EU	
DOCUMENTS PRESENTED:		Technical documentation (Modul A2, C2, H1) <input checked="" type="checkbox"/> EU-type examination certificate (Module C2) <input type="checkbox"/> EC design examination certificate (Module H1) <input type="checkbox"/> Certificate of the QA certificate (Module E, D, H1) <input type="checkbox"/>	
TESTS CARRIED OUT:		Fulfilled	Comments
1.	The aforementioned documents, reports and certificates are available	X	ISS AC575 Chamber Drawing Nr.: MWT150-100-00-21-AD-13
2.	The technical documentation fulfills the requirements of the Directive	X	-
3.	The tests have been carried out in the necessary scope according to art. 3.2 of Annex 3. / Annex I.	X	-
4.	Test of one or more pressure equipment Charge-No.: 00087.13.23044	X	Scope
RESULTS:			
<p>The aforementioned tests have been carried out according to the specified testing requirements. The test documents are complete. In case of the modules A2, C2 the use of the CE 1008 mark on the aforementioned pressure equipment is approved until the next inspection.</p>			
Place: Budapest		Date: 03.12.2024.	
Attachment: -		 Peter Bicsak - Expert	

The most important data of the pressure equipment and the company data are stored for the smooth performance of the order.
Your data are safe.

The test results apply exclusively to the aforementioned subject of tests. The written approval of the testing laboratory is required to copy excerpts from the testing report.

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