

## **Quality Assessment Certificate**Notified Body 2843

This is to certify that ABS ITALY S.R.L. did undertake the relevant type approval procedures for the type of equipment identified below which was found to be in compliance with the requirements of Marine Equipment Directive (MED) 2014/90/EU, subject to any conditions in the schedule attached hereto.

CERTIFICATE NUMBER: 07-280652-9-MED

MANUFACTURER: Ocenco Incorporated

MANUFACTURER PLANT

LOCATION: Pleasant Prairie, Wisconsin, USA

AUTHORISED REPRESENTATIVE: Interspiro AB

EC Type Examination Certificate,

Number: see page 3 DATED: see page 3

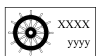
THIS CERTIFICATE IS ISSUED IN COMPLIANCE WITH CONFORMITY ASSESSMENT MODULE D OF THE REGULATIONS AND DIRECTIVES LISTED ABOVE.

ISSUE DATE: 24-oct-2025 EXPIRATION DATE: 05-feb-2030

ELECTRONICALLY SIGNED BY: E. Brina

This certificate authorizes the manufacturer or his authorised representatives, in conjunction with the EC Type Examination (Module B) Certificates listed, to affix the "Mark of Conformity" in accordance with articles 9 & 10 of the Directive.

Example for the application of the "Mark of Conformity":



2843 Number of the Notified Body responsible for the quality surveillance module.

XXXX The year in which the mark is affixed.

The manufacturer shall issue a Declaration of Conformity for each product with reference to the EC Type Examination Certificate and the Quality Assessment Certificate.

Entry Date:

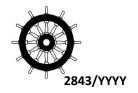
Name of Equipment Manufacturer:

	10225 82nd Avenue Pleasant Prairie Wisconsin, WI 53158 USA	
	Tel: E-mail: Website:	+1-262 947-9000 daniel.brtis@ocenco.com www.ocenco.com
Authorized Representative:	Interspiro AB, Kemistvagen 21 18379 Taby Box 2853 18728 Sweden Tel. +46 8 636 5100	
Equipment/Component:	Emergency Escape Breathing Device (EEBD)	
Model:	M-20.2	
Scope:	European Union Marine Equipment Directive 2014/90/EU, Commission implementing Regulation (EU) 2025/1533: Item MED/3.41 - Emergency Escape Breathing Device (EEBD)	
Comments:	Quality system approved in accordance with the requirements of the European Union Marine Equipment Directive 2014/90/EU for Item MED/3.41c.	
Limitations:	This certificate authorizes the manufacturer or his authorized representative, in conjunction with valid EC Type Examination (Module B) Certificates detailed on page 3 to affix the "Mark of Conformity" in accordance with Articles 9 and 10 of the Directive.	
77		loses its validity if the manufacturer makes any anges to the approved quality system.
	each product Certificate and	rer shall issue a Declaration of Conformity for with reference to the EC Type Examination d this Quality Assessment Certificate in h Article 16 of the Directive.
Revisions:		07-280652-9-MED, replaces Certificate N°. IED Dated: 06-feb-2025

24-oct-2025

Ocenco Incorporated

## Markings & Declaration of Conformity:



## Example for the application of the "Mark of Conformity":

2843 Number of the Notified Body responsible for the quality surveillance module.

YYYY The year in which the mark is affixed.

## The following products are covered by this Quality Assessment Certificate:

MED item No.	MED Certificate No.	Expiry date
MED/3.41c	07-280652-6-EC	23-oct-2030