

Quality Assessment Certificate Notified Body 0729

This is to certify that ABS Europe Ltd., as a Notified Body under the authorization of UK MCA by Merchant Shipping Equipment Regulation, 2016, S.I. 2016 Number 1025 and MSN 1874 (M+F), as amended, did undertake the relevant conformity assessment of the manufacturing plant listed and same was found to be in compliance with the provisions of these Regulations and EU Marine Equipment Directive 2014/90/EU of 18th September 2016

CERTIFICATE NUMBER: 07-HS280652-5-MED

MANUFACTURER: Ocenco, Inc.

MANUFACTURER PLANT

LOCATION: Pleasant Prairie, Wisconsin, USA

PORT OFFICE: Chicago Port

AUTHORISED REPRESENTATIVE: Interspiro AB, Kemistvagen 12 18379 Taby, Sweden

EC Type Examination Certificate,

Number: **07-HS280652-3-EC** DATED: 25 November 2015

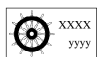
This Certificate is issued in compliance with Conformity Assessment Module f D of the Regulations and Directives listed above.

ISSUE DATE: 24th March 2020 Expiration Date: 5th February 2025

SIGNATURE: N. Tyagi

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":



0729 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.

This certificate evidences compliance with one or more of the Rules, Guides, standards or other criteria of ABS or a statutory, industrial or manufacturer's standards. ABS Europe Ltd. is a Notified Body under the authorization of UK MCA by Merchant Shipping Equipment Regulation, S1 2016, No. 1025 and MSN 1874 (M+F), as amended. Compliance with the provise laid down in EU Directive 2014/90/EU, will be determined as governed by the terms and conditions as contained in EU MED application form – CLS-JBA-00131.



Certificate No.:	07-HS280652-5-MED

Entry Date: 24 March 2020

Name of Equipment/ OCENCO, Inc.

Component manufacturer 10225 82nd Avenue, Pleasant Prairie,

WI 53158, USA

Tel: +1-262 947-9000 Fax: +1-262 947-9020

E-mail: mikekay@ocenco.com Website: www.ocenco.com

Authorized Representative: Interspiro AB,

Kemistvagen 12 18379 Taby, 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) 2019/1397: 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/3.41.

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.

This certificate loses its validity if the manufacturer makes any unapproved changes to the approved quality system.

The manufacturer shall issue a Declaration of Conformity for each product with reference to the EC Type Examination Certificate and this Quality Assessment Certificate in accordance with Article 16 of the Directive.

Markings & Declaration of Conformity:



0729/YYYY

0729 Number of the Notified Body responsible for the quality surveillance module.

YYYY The year in which the mark is affixed.

Revisions: Certificate No. 07-HS280652-5-MED dated 24 March 2020 replaces

Certificate No. 07-HS280652-4-MED