



Lloyd's  
Register

Notified Body authorised by the MCA



Maritime &  
Coastguard  
Agency

## EC (MODULE D) CERTIFICATE OF CONFORMITY

LLOYD'S REGISTER VERIFICATION LIMITED (LRV), designated as a "notified body" under the terms of the Merchant Shipping (Marine Equipment) Regulations 2016 (S.I. 2016 No. 1025) did undertake an assessment of the subject manufacturer's Quality System as per requirements of Annex II Conformity to type based on quality assurance of production process (Module D) of Marine Equipment Directive (MED) 2014/90/EU and Commission Implementing Regulation (EU) 2017/306 indicating design, construction and performance requirements and testing standards for marine equipment and was found to conform with the requirements for the Product Types below.


<b>Manufacturer (Applicant)</b>	Daniamant Limited
<b>Address</b>	Unit 3, The Admiral Park Airport Service Road Hilsea Portsmouth Hampshire, PO3 5RQ United Kingdom (UK)
<b>Reference Regulation Item (No &amp; designation)</b>	<b>Regulation 2017/306 MED/1.2</b> Position-Indicating Lights for life saving appliances (a) For survival craft and rescue boats (b) For lifebuoys (c) For lifejackets

Approval is subject to continued maintenance of the requirements of the above mentioned Directives and to all products continuing to comply with the standards and conditions of EC Type Examination Certificates issued by Lloyd's Register Verification or other Notified Body when they are of the above Designation(s).

Approval is further subject to continued maintenance of the certified quality management system in accordance with the requirements of LRQA certificate number LRQ10007676 or an equivalent replacement thereof.

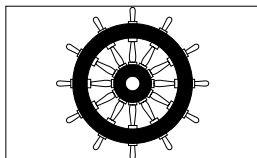
Authorisation is hereby given to the manufacturer to use the LRV Notified Body Registration Number 0038 in accordance with the requirements of the specified Directives in relation to the described products.

**This certificate remains valid unless cancelled or revoked, provided that products manufactured under this Certificate remain satisfactory in service and the above quality management system continues to be approved.**

First issue	3 November 2016		
Date of issue	26 October 2017	Expiry date	25 January 2018
Certificate No.	MED 1650265/M5	Signed	
Sheet No	1 of 2	Name	J. Deboer For and on behalf of Lloyd's Register Verification LRV EC Distinguishing No. 0038

**Note:**

This certificate is issued under the authority of the MCA. No product shall be manufactured under this Certificate unless a valid EC Type Examination Certificate (Module B) is held on that product's Technical File. The manufacturer shall advise the Notified Body of all proposed modifications or changes to a product for which an EC Type Examination Certificate (Module B) has been issued, and of proposed changes of manufacturing location or process, and shall retain copy of their written authorisation or Certification of such changes.



0038/yyyy

Subject to the Manufacturer's compliance with the foregoing, and those conditions of the Directive, the Manufacturer or his Authorised Representative is allowed to affix the 'Mark of Conformity' to products of the types shown above.

yyyy = The year in which the mark is affixed.

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Issue number 1

Date 26 October 2017	Quote this reference on all future communications SOUTSO/TA/QA/MED/JD
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### EC QUALITY SYSTEM (MODULE D) CERTIFICATE No. MED 1650265/M5

#### Places of Production

Daniamant Limited  
Unit 3 Admiral Park  
Airport Services Road  
Portsmouth  
PO3 5RQ

Daniamant ApS  
Industrivej 24C  
3550 Slangerup  
Denmark

Module B No's	Regulation Item number	Date of issue / revision	Notified body
MED 1650264	A.1/1.2(c)	03/11/2016	0038
MED 1650280	A.1/1.2(c)	21/11/2016	0038
MED 1650281	A.1/1.2(a)	22/11/2016	0038
MED 1650282	A.1/1.2(a)	22/11/2016	0038
MED 1750055	A.1/1.2(a)	24/03/2017	0038
MED 1750058	MED/1.2(a)	27/03/2017	0038
MED 1750059	MED/1.2(a)	27/03/2017	0038
MED 1750107	MED/1.2(b)	29/05/2017	0038
MED 1750137	MED/1.2(a)	02/08/2017	0038
MED 1750186	MED/1.2(b)	23/10/2017	0038
MED 1750187	MED/1.2(b)	23/10/2017	0038
MED 1750188	MED/1.2(b)	23/10/2017	0038
MED 1750189	MED/1.2(b)	23/10/2017	0038
MED 1750190	MED/1.2(b)	23/10/2017	0038
BABT-MED001093 issue 02	A.1/1.2(a)	11/11/2013	0168
BABT-MED001094 issue 02	A.1/1.2(a)	20/03/2013	0168
BABT-MED000040 issue 02	A.1/1.2(a)	29/04/2014	0168
BABT-MED000041 issue 01	A.1/1.2(a)	17/12/2012	0168
BABT-MED000042 issue 02	A.1/1.2(a)	19/08/2015	0168
BABT-MED001105 issue 02	A.1/1.2(c)	04/08/2016	0168
BABT-MED001106 issue 02	A.1/1.2(c)	19/08/2015	0168
BABT-MED001111 issue 02	A.1/1.2(c)	21/11/2013	0168
BABT-MED000095 issue 01	A.1/1.2(a)	19/12/2016	0168
BABT-MED000104 issue 01	A.1/1.2(b)	01/08/2017	0168