



Type Approval Application Checklist		
Details:	Document checklist (to be completed by the Applicant):	
1. Applicant: (Company Name and Address of each location, Phone Number, Email address, Contact Person. Please provide information relating to your human and technical resources (including laboratories and/or inspection facilities), and its functions and relationship in a larger corporation/group, if any) MED証明書に記載の御社住所、お電話番号、email,ご担当者名をお書きください。 Request for Marine Services (Form2502TA)と同じ記載をお願いします。	Existing ISO 9001 certificate	
2. Manufacturer: Please note that this is name will be listed on the certificate (if different from 1: Company Name and Address, Phone Number, Email address, Contact Person. Please explain your relationship with the Applicant, any relevant legal obligations)	Existing ISO 9001 certificate	
製造所の場所が設問1と違う場合は,ご住所、お電話番号、email ,ご担当者名を記入してくた	さい。	
3. Authorized Representative: (Name and Address, Phone Number, Email address, Contact Person. Please note : This is required for the Manufacturers not located in the territory of at least one European Union member State applying to the MED Certification). Details of Authorised Representative will not be listed on the certificate	Written mandate	
認定代理人の会社名とご住所を記入してください。		
4. Place(s) of Production: (if different from 1 or 2: Company Name and Address, Phone Number, Email address, Contact Person.) 製造場所が設問1又は2と違う場合は、ご住所、お電話番号、email,ご担当者名を記入してく	□ Existing ISO 9001 certificate ださい。	
5. Product: Name: 例:Lifejacket and Immersion Suit Description: Type:	Documentation to be submitted (for each application): General/functional description of the product Technical documentation including test report(s) Copies of accreditation certificates and	
Application: Marine/Offshore/Industrial (delete as appropriate) Ratings: Standards and/and other normative documents for which certification is sought: Other conditions:	 schedules (for the test house(s)) Analysis and assessment of risk(s) Product Specification/Literature/ data sheets Design Drawings, sufficient to fully define the product Software Quality Plan 	
6. Type Approval Certificate: (Multiple options may be applicable) New Renew Amend LR Type Approval 該当する項目にチェックしてください MED Module B Module D Module B Module D Module F Module Recognition MCA Transport Canada	 Copies of existing Module B EC Type Examination Certificates Copies of EC Declarations of Conformity Relevant Existing Certificates 	
Draft Type Approval Certificate required (will be issued prior to issue of final Certificate in order to allow a review)		
7. For Renewal or Amendments to an existing Certificate please state previous Certificate Number(s):	該当する項目にチェックしてください	
更新又は変更がある場合は、該当する証明書 No.を記載してください。 In addition, if you have a Module D/E Certificate to be amended please list the Certificate number: Module D/Eの修正がある場合は、該当する証明書 No.を記載して下さい。		

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8. Have any changes/amendments been made to the following since p 変更または改正がある場合は、該当する項目にチェッ Product Documentation Technical		If yes, to any changes please provide: Detailed description of changes Relevant documentation
9. Do you outsource any processes, production, or activities relating to U Yes ロNo 外部委託がある場合は詳細を記載して下さ		☐ If yes, please provide details, including information concerning all outsourced processes used that will/may affect conformity to requirements; if another legal entity is used for producing the certified product(s) that is different from your entity, then appropriate contractual arrangements shall be established with that entity.
10. Testing: Specified standards: (Including (Inter)National standards, International Conventions, Rules) Proposed Test Programme, Test Report/Drawings Environmental Testing in accordance with LR Test Specification No. 1: ENV1 – controlled environments only, to producer's specification ENV2 – enclosed spaces subject to temperature, humidity and vibration: 5°C to 55°C ENV3 – enclosed spaces subject to generated heat from other equipment: 5°C to 70°C Existing Test Reports ENV4 – mounted on reciprocating machinery: 5°C to 55°C ENV5 – open decks: -25°C to +70°C 環境試験を行う場合は該当項目にチェック Additional tests e.g. IP65: please state して下さい. Lot下さい.		
11. Please provide all other information such as information for initial evaluation and surveillance activities, e. the locations where the certified product(s) are produced and contact personnel at these locations. 上記の項目以外で何かあれば記載して下さい。		
12. Comments: 上記の項目以外で何かあれば記載して下さい。		
13. Declaration: I declare that information provided is true and complete and that the same application has not been lodged with any other notified body	14. Client 's Name (block capitals please): ご署名と日付、 ご署名された方のお名前をアルファベット大文字でご記入下さい。 Signature:	
15. Application review conducted by (Name, date and signature): (LRV use only)	LRVが記入します。	

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