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**TESTED[®]
DEVICE**

DENSO WAVE Inc.
Resin (U-100 C-N)

Report No. DE 2006-1161

DUPLICATE

Statement of
Qualification

Single product
Biological Resistance

Customer	DENSO WAVE Inc. 1, Yoshiike, Kusaki, Agui-cho, Chita-gun 470-2297 Aichi Japan
Component tested	
Category:	Materials
Subcategory:	Plastics
Product name:	Resin (U-100 C-N) (manufacturing date: 4/2020; color: transparent; serial number: PLATE_2020-08)
Biological resistance test	
Standards/Guidelines:	ISO 846 The norms stated generally refer to the version valid at the time of the tests.
Test environment parameters:	Microbiological laboratory:.....S2
Test procedure parameters:	<ul style="list-style-type: none">• Procedure A (resistance to fungi) using spore suspension of spores containing the following test strains:<ul style="list-style-type: none">– <i>Aspergillus niger</i> ASM 1957– <i>Chaetomium globosum</i> ASM 1962– <i>Paecilomyces variotii</i> ASM 1961– <i>Penicillium pinophilum</i> ASM 1944– <i>Trichoderma virens</i> ASM 1963• Procedure C (resistance to bacteria) using bacteria suspension containing the following test strain: <i>Pseudomonas aeruginosa</i> DSM 1253• Incubation at 29±1 °C with a relative humidity of ≥95 %; visually inspection after four (4) weeks

Test result / Classification

The biological resistance of Resin (U-100 C-N) regarding to growth intensity was investigated in accordance with ISO 846 and classified with the following result:

Biological resistance	Growth intensity	Classification
Procedure A (resistance to fungi)	3	weak
Procedure C (resistance to bacteria)	2	weak
Overall result	weak	


The classification is based on a worst-case consideration of the Procedures A and C. In the process, growth intensity was assessed according to the classification system used in ISO 846:

Classification: fungi (Procedure A)
0 = excellent 2, 3 = weak
1a, 1b, 1c = good 4, 5 = none

Classification: bacteria (Procedure C)
0 = excellent 2 = weak
1 = good 3 = none

The measuring devices used for the qualification tests are calibrated at regular intervals; their results can be traced back to national and international standards. In cases where no national standards exist, the test procedure implemented complies with the technical regulations and norms applicable at the time of the test. The relevant documentation can be viewed on request at any time.

Detailed information and parameters of the test environment can be found in the Fraunhofer IPA test report.

Fraunhofer Institute for Manufacturing Engineering and Automation IPA	DE 1409-725 Report No. first document	Stuttgart, July 15, 2015 Place, date of first document issued
Department of Ultraclean Technology and Micromanufacturing	DE 2006-1161 Report No. current document	Stuttgart, November 4, 2020 Place, current date
Nobelstrasse 12 70569 Stuttgart Germany	on behalf of  Dr.-Ing. Frank Bürger, Project Manager Fraunhofer IPA	