

Application Informa	ation / Informat	tion de soumission
*Licence Name/Nom de l'homologation:		Application Number/Numéro de soumission
ANTI-SARS-COV-2 ELISA (IGG) Licence # / # de l'homologation:		313727 Application Type/Type de soumission:
o		Interim Order
Manufacturer/Fabricant:		Company ID/Identificateur d'entreprise:
EUROIMMUN MEDIZINISCHE LABORDIAGNOSTII	KA AG	116604
Risk Class:		Rationale: Class III by IVDD Rule 2(b)(i)
*Licence Type/Type d'homologation:		Rationale:
Test Kit ▼		
☐ Contains Controlled Substance(s) ☐ Co	ontains Drug(s)	Contains Biological Material(s)
*Intended Use and/or Indications for Use/ Utilisation Prévue	et/ou Indications	
The enzyme immunoassay (ELISA) determination of human antibodies of the immunoglobulin class IgG agor citrate plasma to support the diagnosis of SARS-Cotthe direct pathogen detection Moreover serology can be	gainst SAR: V-2 infectio	S-CoV-2 in serum, EDTA, heparin
product is designed for use as IVD. The test can be process	sed fully au	tomatically.
product is designed for use as IVD. The test can be process		•
product is designed for use as IVD. The test can be process	Licence Informa	ation
product is designed for use as IVD. The test can be process	Licence Informa	•
product is designed for use as IVD. The test can be process	Licence Informa	ation
product is designed for use as IVD. The test can be process OEM DEM Licence Name:	Licence Informa	ation
product is designed for use as IVD. The test can be process OEM DEM Licence Name:	Licence Informa	ation
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use	Licence Informa	ation anufacturer :
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change	Licence Informa	ation anufacturer :
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device	Licence Informa	ation anufacturer :
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change	Licence Informa	ation anufacturer :
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change License name change	Licence Informa OEM M	ation anufacturer :
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change License name change Device name change	Licence Informa OEM M	ation anufacturer :
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change License name change Device name change Change to the purpose/indication of	Licence Informa OEM M	ation anufacturer :
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change License name change Device name change Change to the purpose/indication of license	Licence Informa OEM M	ation anufacturer :
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change License name change Device name change Change to the purpose/indication of license Addition of device(s) Deletion of device(s)	Licence Informa OEM M	ation anufacturer: Comment(s)
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change License name change Device name change Change to the purpose/indication of license Addition of device(s) Deletion of device(s) Reason for Change	Licence Informa OEM M	ation anufacturer :
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change License name change Device name change Change to the purpose/indication of license Addition of device(s) Deletion of device(s)	Licence Informa OEM M	ation anufacturer: Comment(s)
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change License name change Device name change Change to the purpose/indication of license Addition of device(s) Deletion of device(s) Reason for Change Change in name and/or address of the	Licence Information OEM M	ation anufacturer: Comment(s)
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change License name change Device name change Change to the purpose/indication of license Addition of device(s) Deletion of device(s) Reason for Change Change in name and/or address of the Private Label Manufacturer	Licence Information of the control o	ation anufacturer: Comment(s)
product is designed for use as IVD. The test can be process OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change License name change Device name change Change to the purpose/indication of license Addition of device(s) Deletion of device(s) Reason for Change Change in name and/or address of the Private Label Manufacturer Private Label License name change	Licence Informa OEM M	ation anufacturer: Comment(s)

Certificate Screening Checklist:



Cert # (new):	Cert Revisions /	Comments (If Applicable):
Cert. # (old):		
Replacing Existing Cert on File (Y/N):		
Criteria	conforms	Comments/info for MDS
Issued to full name of manufacturer as it appears on application/licence and label.		
Issued to complete civic address matching application/licence and label.		
Criteria are ISO 13485:2016 and Medical Devices Regulations – Part 1 – SOR 98/282		
Scope activities limited to design, development, manufacture, production, servicing, installation, or distribution.		
Activities include "manufacture" or "production" Activities include "design" or "development and development" for class III/IV devices.		
Scope is unambiguous and covers app./lic. devices. Does not contain product names/models/licence numbers.		
Auditing Organisation is Authorized or Recognized		
Statement of Authorization or Recognition.		
Field labelled "Effective Date"		
Field labelled "Expiry Date"		
Validity period ≤ 3 years		
Unique identification code labelled "certificate number" or "certification document number"		□ new □ revised
Name, title, and signature of certification authority		
Pagination (page x or y) included on all pages . All pages present.		
Method to verify validity		
	.	
Screening	g Decision	
Accept		▼
Liem Whelan		Date : April 14 th , 2020
Device Licensing Services Division Medical Devices Bureau	_ _	

Santé Canada

Check GMDN

Code	GMDN Name	GMDN	GMDN Description	lssue	Action for Screener	Regulatory Action	Reference Material
		Code					