

Application Informa	ition / Inform	ation de soumission	
*Licence Name/Nom de l'homologation: FTD SARS-COV-2		Application Number/Numéro de soumission 315376	
Licence # / # de l'homologation:		Application Type/Type de soumission: Interim Order	
Manufacturer/Fabricant: FAST TRACK DIAGNOSTICS LUXEMBOURG S.A.R.L.		Company ID/Identificateur d'entreprise: 139974	
Risk Class:		Rationale: Classification Rule IVDD Rule 2(b)(i)	
*Licence Type/Type d'homologation:		Rationale:	
Test Kit ▼			
☐ Contains Controlled Substance(s) ☐ C	ontains Drug((s) Contains Biological Material(s)	
*Intended Use and/or Indications for Use/ Utilisation Prévue	et/ou Indications		
FTD SARS-CoV-2 is a qualitative in detection of severe respiratory syn coronavirus 2 (SARS-CoV-2) nuclei oropharyngeal swabs of patients w symptoms of SARS-CoV-2 infection epidemiological risk factors, who a Coronavirus Disease 2019 (COVID-The test is intended as an aid in the	drome c acids in ith signs a in conjui re suspec 19).	nasopharyngeal and and nction with clinical and eted of	
human coronavirus SARS-CoV-2. For in vitro diagnostic use.			
For in vitro diagnostic use.	Licence Inforn	mation	
For in vitro diagnostic use.		nation Manufacturer :	
For in vitro diagnostic use.			
OEM Intended Use and/or Indications for Use		Manufacturer :	
OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change			
OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device		Manufacturer :	
OEM Licence Name: OEM Intended Use and/or Indications for Use Reason for Change Change to classification of a device Manufacturer name change		Manufacturer :	
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		Manufacturer :	
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		Manufacturer :	
Por in vitro diagnostic use. 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		Manufacturer :	
Por in vitro diagnostic use. 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		Manufacturer :	
Por in vitro diagnostic use. 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		Manufacturer :	
Por in vitro diagnostic use. 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)		Manufacturer :	
Por in vitro diagnostic use. 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		Manufacturer :	
Por in vitro diagnostic use. 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)		Comment(s)	
Por in vitro diagnostic use. 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	OEM	Comment(s)	
Por in vitro diagnostic use. 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	OEM	Comment(s)	
Por in vitro diagnostic use. 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	OEM	Comment(s)	



Cert # (new):	Cert Revisions /	Comments (If Applicable):
Cert. # (old):		
cette ii (ola).		
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		▼
Jiazhen Minnie Dai		Date:2020-05-08
Device Licensing Services Division Medical Devices Bureau		

*
7

Santé Canada

Health Canada Check GMDN

Reference Material Regulatory Action **Action for Screener** Issue **GMDN Description** GMDN Code **GMDN Name**