

Application Inform	nation / I	Information de soumission		
*Licence Name/Nom de l'homologation:		Application Number/Numéro de soumission		
ANOSMIC COVID-19 SMELL TESTER		319520		
Licence # / # de l'homologation:		Application Type/Type de soumission:		
Manufacturer/Fabricant:		Interim Order  Company ID/Identificateur d'entreprise:		
VIROCULE INC.		160784		
Risk Class:		Rationale:		
1		Rule 7(1)		
*Licence Type/Type d'homologation:		Rationale:		
Single Device				
$\square$ Contains Controlled Substance(s)	Contains	s Drug(s) Contains Biological Material(s)		
*Intended Use and/or Indications for Use/ Utilisation Prévu	e et/ou Ind	lications		
identify anosmia " the loss of sme	ell " wh daily if	nables the user to smell the vapor to nich is a major symptom of COVID-19. Ithey have the coronavirus and take Center.		
OEN	1 Licence	Information		
OEM Licence Name :		OEM Manufacturer :		
OEM Intended Use and/or Indications for Use				
Reason for Change		Comment(s)		
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		Comment(s)		
Change in name and/or address of the				
Private Label Manufacturer				
Private Label License name change				
Private Label Device name change				
Addition of device(s)				
Deletion of device(s)				
Not sure what kind of review documents are needed for this of they provided a manual and brochure. They did not provide a labelling)  Certificate Screening Checklist:		ficate when asked but gave their MDEL (only for packaging and		
Cert # (new):		Cert Revisions / Comments (If Applicable):		
Cert. # (old):				



Replacing Existing Cert on File (Y/N):				
				_
Criteria	conforms	Comme	nts/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"		$\square$ new	v □ 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			_	
Kevin Nguyen ▼		Date:2	020/08/14	_
Device Licensing Services Division  Medical Devices Directorate				
Review Division - D	LSD Commu	nication		
Review Division Screener Action:				
Review Division Screener Response:				
TOTION DIVISION CONCENSION RESPONSE.				
			Date:	
Project Philips Communication				
Review Division Screener Medical Devices Directorate				

*
7

Santé Canada

Health Canada Check GMDN

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