03- Application Form

APPLICATION FOR INVESTIGATIONAL TESTING AUTHORIZATION

(disponible en français)

1.	DEV	ICE CLASSIFICA	ATION			
	\checkmark	Class II	□ Class	III		Class IV
1.	[Not	e: this is the devic	it appears on label) e name for which the At	athorization will be iss	ued]	
	Pulse o	ximeter				
2.	_	OTOCOL IDENT	IFICATION: agnosis or treatment fo	or which the device w	vill b	e sold.
	The pur present pressure	rpose of this devicing clinicians and	e is to non-invasively musers with an oxygen sa	easure oxyhemoglobituration (SpO2), which	n and h is a	l deoxyhemoglobin,
3.			SS OF MANUFACTU and address to which to	`		,
	Compa	ny Name	Glia Inc.			
	Street A	Address/P.O. Box:	54 Craig St.			
	City: Lo	ondon				
	Provinc	ce/State: Ontario				
	Postal/Z	Zip Code: N6C 1E	8			
	Country	y: Canada				
	Contact	t Name and Title:	Dr. Tarek Loubani,	Primary investigator a	nd Eı	mergency Physician
	Telepho	one No.: +1	519 488 6475	Fax No.: NA		
	E-Mail	Address:	tarek@tarek.org			

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4. MAILING ADDRESS FOR REGULATORY CORRESPONDENCE (if different from 4)

Note: (i) The Authorization will be **issued** to Company named in Item 4 but will be **sent** to the Company shown below if different. (ii) The Company named below must be authorized by the manufacturer named in Item 4 to submit an authorization application on their behalf.

Company Name				
Street Address/P.O. Box				
City Province/State Postal/Zip Code Country				
			Contact Name and Title:	
			Telephone No.:	Fax No.:
			E-Mail Address:	
DEVICE TYPE (check one o				
Single Device	. ✓			
Medical Device Group				
Medical Device Family				
Medical Device Group Family				
Test Kit				
System				
PREFERRED NAME CODE	a) optional			
	Province/State Postal/Zip Code Country Contact Name and Title: Celephone No.: E-Mail Address: DEVICE TYPE (check one only) Single Device Medical Device Group Medical Device Family Medical Device Group Family Cest Kit System			

(disponible en français)

8.	DEVICE	HICACE	CATE	CODV
δ.	DEVICE	USAUL	CAIL	してひだY

(73) Anaesthesiology	✓
(74) Cardiovascular	
(76) Dental	
(77) Ear, Nose & Throat	
(78) Gastroenterology & Urology	
(79) General & Plastic Surgery	
(80) General Hospital	✓

(84) Neurology	
(85) Obstetrics & Gynaecology	
(86) Ophthalmology	
(87) Orthopaedics	
(89) Physical Medicine	
(90) Radiology/Imaging	

FOR IVDDs ONLY

(75) Chemistry	
(81) Haematology	
(82) Immunology	

(83) Microbiology	
(88) Pathology	
(91) Clinical Toxicology	

9. DOES THIS DEVICE CONTAIN A DRUG?

(Note: this question does not apply to IVDDs)

Yes□ No**x**

If yes

Brand /Trade Name of Drug:
Active Ingredient:
Drug Manufacturer:
Applicable Drug Identification Number (if any):

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10. **DEVICE DETAIL**

Please provide the following information, where applicable for each component device, part or accessory.

Name of Device, Components, Parts and Accessories as per product label	or Device Identificati Number if previou assigned	on Model or Catalogue number sly
Pulse Oximeter – control unit	None assigned	POX-C001
Pulse Oximeter – finger probe	None assigned	POX-F001

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11. ATTACHMENTS

In addition to items 1 to 11, of the Application for investigational testing, please indicate (49) which of the relevant information requirements listed below, are included as attachments to this application, or will be provided at a later date. For details regarding content and format, please refer to the guidance documents "Preparation of an Application for Investigational Testing - Medical Devices" and "Preparation of an Application for Investigational Testing - in Vitro Diagnostic Devices"

	Attached	To Come
Background Information	x	
Risk Assessment	NA	
Ethics Committee or IRB Approval(s)	X	
Protocol	X	
Device Label	X	
Investigator Agreements	x	

12.	If this Device contains a drug and it does not have a Drug Identification Number, I the			
	Manufacturer of this device attest that the (\square drug meets) meet) acceptable standards of safety, efficacy and quality.	$(\Box$ drug does not		
	I hereby certify that the information provided on this application and in any attached documentation is correct, complete and in accordance with all relevant sections of the <i>Medical Devices Regulations</i> .			

Name of Signing Official: Tarek Loubani

Signed: Date: 2018 August 8