	Laboratory Section	Form	BMU	-FM-65	Rev. 0	05 P.	AGE 1 OF 4
and the state of Medical Deli	Alle Selection of the s	ชื่อเอกสาร : Sample Record Form					
	San	m		Date Rec	(For Staff	`Only)	
Client Name:							
	ition / Institution				•••••	••••••	
	tuon / mstituton					•••••••••••	
	Fax:						
161	T dX		E-man	•••••	•••••	•••••	•••••
1. Desired Tes	sting for Medical Devices (🗹 C	heck the empty box)					
						Code	
	Test		Normal*	Fast Trac		r Staff Only)	
	1. Cytotoxicity Testing	. Cytotoxicity Testing					
	1.1 ISO certified (in-scope)	14,200 Bah	17,400) Baht			
	1.2 ISO uncertified (out-of-scope)	.2 ISO uncertified (out-of-scope)			00 Baht		
	2. Hemolysis Testing	10,500 Bah	31,50	00 Baht			
		5,00	0 Baht				
		Rabbit Blood	10,000 Baht				
	3. Composition Ananlysis	2,700 Bah	5,30	0 Baht			
	4. Microbiology Testing						
	4.1 Bioburden Testing		1,800 Baht	5,30	0 Baht		
	4.2 Bioburden Validation (Staff C	2,700 Baht	7,40	0 Baht			
	(sample <u>withou</u> t pores/threads/fiber)						
	4.3 Bioburden Validation (Staff C	5,300 Bah	12,60	00 Baht			
	(sample with pores/threads/fiber)						
_	4.4 Sterility Testing (14 days testing	3,700 Bah	10,50	0 Baht			
	4.5 Antibacterial Susceptibility to						
	Agar Diffusion Method	Agar Diffusion Method Broth Dilution Method					
	4.5.1 Staphylococcus aureus	4.5.1 Staphylococcus aureus			0 Baht		
	4.5.2 Staphylococcus epidermidis	4.5.2 Staphylococcus epidermidis			0 Baht		
	4.5.3 Pseudomonas aeruginosa		1,800 Bah	5,30	0 Baht		
	4.5.4 Escherichia coli		1,800 Bahi	_	0 Baht		
	4.5.5 Staphylococcus aureus MRSA (drug resistant)			=	0 Baht		
	4.5.6 Candida albican	3 700 Baht	I 1 10.50	0 Baht		(

2. Issuing Test Results (Customer will receive one Thai version of the Test Report as default)

4.5.7 Porphyromonass gingivalis

Requesting additional Test Results
English Version of the Test Report (4,000 Baht)
Thai Version of the Supporting Document (4,000 Baht)
English Version of the Supporting Document (4,000 Baht)

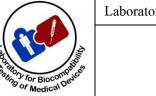
9,000 Baht

26,300 Baht

 $\textbf{\textit{Note}: The Supporting Document includes details about sample preparation, testing process, and qualitative test result}$

^{*}Regular Track means that the client will receive the service within 60 days from payment date.

^{**}Fast Track means that the client will receive the service within 14 days from payment date.



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3.	Details of Sample (for test report)							
	3.1 Name:							
	3.3 Type of material (e.g., Plastic, Steel,	Textile):						
	3.4 Model	Grade	Color	Size / Volume	cm / n			
	Feature	Lot Number		Amount	gram / sheet / roll / se			
	Manufacturer	Country		Dealer				
	Additional Notes:							
	3.5 Storage condition:							
4.	Supporting Documents for Samples	○ No ○ Y	/es	Others (specify)	 			
•	Warnings / Cautions / How-to-use		/es	Others (specify)				
 5.	Campany's Nama and Address specifi	ad in the Test Penert						
٥.		Company's Name and Address specified in the Test Report Name - Address						
	- 1							
6.	Company's Name and Address specific	ed in the Tax Invoice						
	According to the Test Report							
	Others (please specify)							
	Tax Identification Number		Bra	anch				
7.	Payment							
	□ Cash □	Transfer						
8.	Receiving Test Report	In person		By post office, to the address	s indicated above ***			
	Or another address specified here:							
9.	Receiving Samples In person, alon	ng with the Test Report						
	By mail, along with the Test Report *** By mail, according to the address in (No. 5.) ***							
	***Additional shipping cost will be a	pplied						
10.	Decision Criteria (Referenced according	g to ILAC G8:09:2019 G	uidelines on D	Decision Rules and Statement of Cor	nformity)			
	In the case that customers:							
	On't desire	to decide the test resi	ults criteria. Th	ne laboratory will report according to	o the Test Report			
		Criteria (No. 11)						
	Desire	to decide the test resi	ults criteria. Th	ne test results will be reported in acc	cordance with the			
		guidelines provided.	However, it m	oust be samples that are accredited for	or the laboratory.			



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11. Test Report's Criteria

1. Reporting of In Vitro Cytotoxicity Test

1 Qualitative evaluation: The achievement of a numeric grade greater than 2, based on below table, is considered a cytotoxic effect.

Table - Qualitative morphological grading of cytotoxicity of extracts

Grade	Reactivity	Conditions of all cultures
0	None	Discrete intracytoplasmatic granules, no cell lysis, no reduction of cell growth
1	Slight	Not more than 20 % of the cells are round, loosely attached and without intracytoplasmatic granules, or show
		changes in morphology; occasional lysed cells are present; only slight growth inhibition observable.
2	Mild	Not more than 50 % of the cells are round, devoid of intracytoplasmatic granules, no extensive cell lysis; not more
		than 50 % growth inhibition observable.
3	Moderate	Not more than 70 % of the cell layers contain rounded cells or are lysed; cell layers not completely destroyed, but
		more than 50 % growth inhibition observable.
4	Severe	Nearly complete or complete destruction of the cell layers.

^{2.} Quantitative evaluation: Reduction of cell viability by more than 30 % is considered as a cytotoxic effect.

Reference: ISO 10993-5:2009 Biological evaluation of medical devices — Part 5: Tests for in vitro cytotoxicity

2. Reporting of In Vitro Hemolysis Test			3. Reporting of Bioburden Testing			
Hemolytic Index above Negative Control		Reaction Level	Grade	Reaction Level	Characteristic / Conditions	
0-2		No reaction	-	No microorganism observed	No microorganism is observed on the medium after culturing with the test sample.	
2-5		Slight reaction	+	Microorganisms observed	Microorganisms are observed on the medium after culturing with the test sample.	
>5		Severe Reaction				
Reference : ASTM F75	Reference : ASTM F756-13:2019 Standard Practice for Assessment of			Reference : ASTM D3516 Standard Test Methods for Ashing Cellulose		
Hemolytic properties of	Hemolytic properties of Materials					
4. Reporting of Bioburden Testing with Identification			5. Reporting of Sterility Testing			
Test Result	ult Characteristic / Conditions		Grade	Reaction Level		
Types of microorganisms	medium after culturing with the test sample are		-	N	Io microorganism observed	
Number of The number of microorganisms that grew on the medium microorganisms after culturing with the test sample.		+		Microorganisms observed		
Reference: ISO 11737-1: 2018 Sterilization of medical devices - Microbiological			Reference: ISO 11737 – 2:2019 Sterilization of medical devices -			
methods - Part 1: Determination of a population of microorganisms on products			Microbiological methods - Part 2: Tests of sterility performed in the validation of			
AMENDMENT I			a sterilization process			



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6. 1	Reporting of Susc	ceptibility Test by means of Agar diffusion	7. Reporting of Susceptibility Test by means of Broth dilution		
Grade	Reaction Level	Characteristic / Conditions	Test Result	Characteristic / Conditions of Cultured Cell	
-	No inhibition zone	No inhibition zone is presented around the test sample that was placed on the medium.	MIC	The lowest concentration of the sample that can inhibit the growth of microorganisms.	
+ Slight Inhibition zone is presented around the test inhibition zone sample that was placed on the medium.			*MIC: Minimum inhibitory concentration Reference: CLSI M07 10 th edition: 2015 Methods for dilution antimicrobial susceptibility tests for bacteria that Grow aerobically		
Reference : CLSI M02 13 th edition: 2018 Performance standards for antimicrobial disk susceptibility tests					

Notes						
	(If it is requested to test on samples with abnormalities or deviation from specified conditions, there will be a statement or a disclaimer – for example,					
	"This report is outside the scope of ISO/IEC 17025:2017", in the test report.					
	Sender	Receiver				
	()	()				
	Date	Date				
	Laboratory Mana	ager				
	()				
	Date					