

LABORATORY SERVICE REQUEST (LSR) - BIOCOMPATIBILITY (ISO)

CLIENT INFO

(**Instructions**: Use ONE form for each group of similar samples requiring the same storage, handling, analyses, and compliance. Fill in the form as completely as possible. Submit the form along with sample(s). Initiation of sample analysis will be delayed if form is not complete.)

Sponsor: (Send Report To)	Invoice To:
Contact Name:	AP Contact Name:
Company Name:	Company Name:
Address:	Address:
City/State/Zip:	City/State/Zip:
Country:	Country:
Phone:	Invoice To Email:
Email:	AP Email:
PBL Quote Number:	AP Phone:
PO Number:	Credit Card to be used for payment

TEST ARTICLE INFO Test Article ID (Please use the exact wording you want to appear in the final report.) Lot Number: Part Number: Other Identifier: (ex. Sample Code) **Expiration Date:** Quantity of Test Articles Submitted: (Please indicate the number of units, volume, and/or weight of test article.) Physical Description: Other: Solid Powder Device Liquid Gel Hazardous: Sample Disposition: (Check one box) Storage Condition: Controlled Substance: (include MSDS if samples are hazardous, client (Samples will be discarded 30 days after report will incur charges for disposal of hazards.) issuance unless otherwise indicated.) Room Temperature ∏ No Not Hazardous Discard 2 to 8°C Reactive Hazardous Waste Municipal Waste -10 to -25°C 🗌 Yes Biohazard Return Unused 🗌 Return All -60 to -90°C Schedule: Toxic Other: Carrier: Account:



TEST ARTICLE INFO (CONT.)

List part(s) of the Test Article that should be tested:		
List part(s) of the Test Article that should be excluded:		
Final intended use/application of Test Article?:		
Can Test Article be cut?		
Sterility Status: Non-Sterile Sterile (Please indicate method)		
Is Test Article absorbent? No Yes (Additional test article(s) required for absorbing capacity determination)		
Unknown (Requirement for absorbing capacity determination will be assessed by Pacific BioLabs)		
Extraction Conditions: (for tests other than the Cytotoxicity tests)		
□ 121°C for 1 hr □ 70°C for 24 hrs □ 50°C for 72 hrs □ 37°C for 72 hrs □ Other:		
Surface Area of one device (cm ²): Thickness (mm):		
ISO 10993-12, The standard surface area can be used to determine the volume of extraction vehicle needed. This area includes the combined area of both sides of the sample and excludes indeterminate surface irregularities. When surface area cannot be determined due to the configuration of the sample, a mass/volume of extracting fluid shall be used. Unusually complex surface area calculations that are to be performed by PBL may incur additional charges. Extraction of large samples may incur additional media charges.		
Note: Non-patient contacting portions of the medical device should, if possible, be excluded either physically from test sample extracts or by exclusion of the surface area in the calculation of the extraction ratio. When this is not possible, the extraction ratio shall be justified, Ensure that all contacting portions are covered by the selected extraction vehicle volume.		
The extraction ratios are selected based on the thickness of the sample to be tested. Please use the following criteria to select the proper extraction ratio (ISO 10993-12):		
□ $3 \text{ cm}^2 / 1 \text{ mL}$ for samples $\ge 0.5 \text{ mm}$ thick		
6 cm ² / 1 mL for samples < 0.5mm thick		
Regulatory Treatment: Non-regulatory GGMP GLP		
Complete the following section if GLP:		
GLP Stability Testing and Test Article Characterization (Certificate of Analysis):		
Will be provided during study Will not be provided during study		
 Note: GLP testing requires, by Federal regulation, accurate identification of the test and control articles (e.g., a lot number, batch number or sample code) that allows unambiguous traceability of the tested material. GLP testing also requires that the testing laboratory document characterization of the test and control articles for stability, strength, purity, and composition, or other characteristics which will appropriately define the test or control article. This information is typically provided by the Sponsor and is required for each batch of test or control article tested; this documentation is often in the form of a Certificate of Analysis (C of A). For testing of biomedical devices that contain no drugs, Pacific BioLabs requires documentation of the identity, composition, and stability of the material tested. Failure to provide this information will be noted as noncompliance with GLP regulations in the Final Report. 		
Rush Services: No Yes (will incur a 50% surcharge)		
Report Format: PDF PDF and Paper Paper (Note: A fee will apply for paper copy of report.)		



ANALYSIS REQUIRED - Please check all required tests		
CYTOTOXICITY TESTS ISO 10993-5:		
Agar Diffusion		
Direct Contact		
MEM Elution (extract test)		
Please select extraction condition: 🔄 37°C for 24 hrs 🔄 37°C for 72 hrs (extended extraction)		
MTT - Quantitative Evaluation (extract test)		
Please select extraction condition: 37°C for 24 hrs 37°C for 72 hrs (extended extraction)		
Please select if images of the 24-hour and 48-hour scoring are to be included in final report (additional fee will apply). (Final Reports will not include images unless otherwise indicated.)		
Yes, include cell images No, cell images are not required		
SENSITIZATION TESTS ISO 10993-10:		
Closed Patch Test for Delayed-Type Hypersensitivity		
Maximization Test for Delayed-Type Hypersensitivity		
Please select extraction medium: Cottonseed Oil Saline No Extract (Direct Application) Other:		
IRRITATION TESTS ISO 10993-23:		
Intracutaneous (intradermal) Reactivity Test		
Dermal Irritation Test		
Ocular Irritation Test		
Vaginal Irritation Test		
Rectal Irritation Test		
Penile Irritation Test		
Oral Irritation Test (Hamster Cheek Pouch)		
Please select extraction medium: Cottonseed Oil Saline No Extract (Direct Application) Other:		
SYSTEMIC TESTS <i>ISO 10993-11</i> :		
Acute Systemic Toxicity Test		
Please select extraction medium: Cottonseed Oil Saline No Extract (Direct Application) Other:		
Material Mediated Pyrogenicity (Saline Extract Only)		
SUBACUTE TOXICITY TESTS ISO 10993-11:		
$\square Repeated Dose Toxicity Study \rightarrow Custom designed, as per protocol.$		
SUBCHRONIC/CHRONIC TOXICITY TESTS ISO 10993-11:		
$\square Repeated Dose Toxicity Study \rightarrow Custom designed, as per protocol.$		
IMPLANTATION TESTS* ISO 10993-6:		
Please select route:		
Please select duration: 7 Days 14 Days 30 Days 60 Days 90 Days 180 Days 365 Days		
*Histopathology is included in all implantation studies.		
HEMOCOMPATIBILITY ISO 10993-4:		
Hemolysis: ASTM Method		
Please select route: Direct and Indirect Direct Indirect		



ANALYSIS REQUIRED - Please check all required tests (continued)				
SUBCONTRACTED TESTS				
GENOTOXICITY <i>ISO 10993-3</i> :				
🗌 Ames Test 🔄 Mouse Lymphoma 📄 Chromosomal Aberration	Mouse Micronucleus			
Please select extracts: Saline DMSO PEG 400	Other:			
HEMOCOMPATIBILITY ISO 10993-4:				
Prothrombin Time (PT) Partial Thromboplastin Time (PTT)				
Platelet Aggregation Platelet Count				
Compliment Activation				
In Vivo Thrombogenicity				
☐ Other:				
Additional Service: Autoclaving (121°C for ~30 minutes) will ind				
OTHER TESTS/SPECIAL INSTRUCTIONS:				
The signature of the Sponsor (or Sponsor's representative) below is assurance	that 1) the study is appropriate to the Sponsor's project			
goals and that no alternative in vitro or decreased in vivo animal use procedure	es are available to meet the stated purpose of the study, 2)			
the species chosen is appropriate to the stated purpose of the study and that use of alternative species has been considered, 3) the study				
is not an unnecessary duplication of previous work, and 4) the number of animals used is appropriate to establish biological or statistical significance as required by the study. The Sponsor also specifies that documentation for the above assurances may be obtained from the				
Sponsor.				
	DATE			
TESTING AUTHORIZED BY (Please Sign):	DATE:			

(Signature and date, or electronic signature is required for testing to begin; unsigned LSR forms will not be processed)