Lab Use



# **REQUEST FORM 005A**

# Labelled Relative Quantitation Analysis

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### **SECTION A**

Contact details	
Name	Billing Address
Organisation / Institution	Email Telephone
Purchase Order Number	Fax

Sample Information - Please complete details on all pages & sign page 4

## Proteomics Analysis price guide as of March 2022. Consult our website for latest price information

Serv ☑	rice 005A – Labelled Relative Quantitation Analysis & Proteome Mapping  1D-LC map (Proteome Mapping) – Mandatory sample viability check for all iTRAQ/TMT experiments		Price (USD) \$330 per experiment
	<b>4-plex iTRAQ</b> - Sample labelling and analysis by 2D LC-MS/MS with automatic database analysis	Up to four samples	\$6,600 per experiment
	4-plex iTRAQ; duplicate experiment (2 x 4-plex)		\$9,900
	<b>4-plex iTRAQ</b> ; triplicate experiment (3 x 4-plex)		\$13,200
	<b>8-plex iTRAQ</b> - Sample labelling and analysis by 2D LC-MS/MS with automatic database analysis	Up to eight samples	\$8,250 per experiment
	8-plex iTRAQ; duplicate experiment (2 x 8-plex)		\$12,375
	8-plex iTRAQ; triplicate experiment (3 x 8-plex)		\$16,500
	<b>6-plex TMT;</b> Sample labelling and analysis by 2D LC-MS/MS with automatic database analysis	Up to six samples	\$7,500 per experiment
	<b>6-plex TMT</b> ; duplicate experiment (2 x 6-plex)		\$11,250
	6-plex TMT; triplicate experiment (3 x 6-plex)		\$15,000
	<b>10-plex TMT</b> ; Sample labelling and analysis by 2D LC-MS/MS with automatic database analysis	Up to ten samples	\$11,000 per experiment
	<b>10-plex TMT</b> ; duplicate experiment (2 x 10-plex)		\$16,500
	10-plex TMT; triplicate experiment (3 x 10-plex)		\$22,000
	Data and result files for publication purposes		\$275

### Lab Use Only:

	1D experiment	Replicate 1	Replicate 2	Replicate 3
Sample Received:				
Processed/Operator:				
Spot set:				
MS analysis/Operator:				
Report Reference:				
Checked Workflow:				
Checked Report:				
Storage:				

### **SECTION B**

# NOTE:

- 40  $\mu$ L plasma is required for a single experiment and 80  $\mu$ L plasma is required for duplicate experiments
- Liquid samples should be in a volume of 100  $\mu L$  200  $\mu L$  and have a concentration of 2 mg/mL 5 mg/mL
- Acetone precipitated sample pellets should contain a minimum of 100  $\mu g$  of protein but not exceed 200  $\mu g$  per tube.
  - Excess of 200 µg will not be processed.
- Samples should contain equivalent amounts of protein per tube.

<b>Description of sample:</b> (Liquid, Pellet, Lyophilised)	
Amount of starting material: (eg. 100 mg Leaf, 1 mg muscle, no. of cells, etc.)	
Method used for protein estimation:	
Buffer composition: Note; please state buffer composition before drying, extraction buffer.	
Any other treatments or chemicals present including volume and final concentration: (eg. Acetone precipitation, sucrose, reduction reagents, etc.)	

# Comments:

### **SECTION C**

# **Experimental Design**

Please i. Circle appropriate response

Is this a single experiment? Yes No

Is this experiment? iTRAQ 4-plex iTRAQ 8-plex TMT 6-plex TMT 10-plex

No. of replicates 1 2 3

If more than 1 replicate Repeat experimental design for each replicate (Please print another copy of this page and fill in the details)

# iTRAQ 4-plex

Label	114	115	116	117
Sample name				
Amount of protein				
PI Ref				

Control Sample (please circle) 114 115 116 117

# iTRAQ 8-plex

Label	113	114	115	116
Sample name				
Amount of protein				
PI Ref				
Label	117	118	119	121
Sample name				
Amount of protein				
PI Ref				

Control Sample (please circle) 113 114 115 116 117 118 119 121

# **TMT 6-plex**

Label	126	127	128
Sample name			
Amount of protein			
PI Ref			
Label	129	130	131
Label Sample name	129	130	131
	129	130	131

Control Sample (please circle) 126 127 128 129 130 131

TMT 10-plex

<u> </u>					
Label	126	127N	127C	128N	128C
Sample name					
Amount of protein					
PI Ref					
Label	129N	129C	130N	130C	131
Sample name					
Amount of protein					
PI Ref					

Control Sample (please circle) 126 127N 127C 128N 128C 129N 129C 130N 130C 131

### **SECTION D**

Further details on database for protein identification

Effective protein identification by mass spectrometry is highly dependent on access to an appropriate database. Answers to the following questions will guide the data analysis pipeline.

1.	What is the target organism?
2.	What other contaminating organisms are likely to be present in the sample provided?
3.	What are the most taxonomically related species of the target organism?
4.	Is the database for the target organism or its related species available in the NCBI or Swiss-Prot
	databases, otherwise where can they be downloaded? Please provide details.
	t <u>e</u> ase consider <i>De novo</i> peptide sequencing (Service 002) if the target species is not available or not Il represented in the NCBI or Swiss-Prot databases.
Please	sign here below:
1.	I have read and understood the Proteomics Analysis Price List and agree to the charges and to Proteomics International's standard Terms and Conditions (available at: <a href="http://www.proteomics.com.au/analytical-services/terms-and-conditions/">http://www.proteomics.com.au/analytical-services/terms-and-conditions/</a> ).
2.	Hazards: I declare that the sample(s) are non-harmful, non-infectious and non-radioactive.
3.	I have completed both pages of this submission form with details for each sample submitted for analysis.
4.	For students, please ensure supervisor signs this form.
Not	e: Please be aware that samples are destroyed by analysis and cannot be returned.
uthor	ised Signature Date: