

AHARIBO Protein System 360

Product	Catalog no	Rxns.
AHARIBO Protein System 360	#AHA360PS	12

Shipping: Blue Ice

<u>Storage Conditions</u>: store components according to the storage conditions reported on the labels, and on Page 6 of this manual.

Shelf Life: 12 months

<u>Description</u>: AHARIBO Protein System 360 is a complete solution for the study of *de novo* synthesized peptides combining the reagents to carry on the wet lab part and the associated LC-MS analysis and bioinformatics services. The kit includes reagents and components for 12 proteomic reactions (from *de novo* synthesized peptides pull down) and a USB pen drive providing access to the bundled LC-MS and Data Analysis services.

Suitable for: Eukaryotic cell lines

For Research Use Only. Not Intended for Diagnostic or Therapeutic Use.

		/		
Mod 080264	v1.1	Date Exp/rev 17/03/2025	Edited by MA PB LM	Pag 1 of pag 14

Table of Contents

Kit storage info	3
Additionally Required Materials	3
INTRODUCTION	4
WORKFLOW OVERVIEW	4
A. SAMPLE PREPARATION	5
A.1 Sample Recommendations	5
A.1.1. Cell Medium Recommendations	5
Optimal Workflow Recommendations	6
B. Pull-down of de-novo synthesized protein	7
Step B1. CELL LYSIS	8
A Before starting the experiment – Supplemented Lysis Buffer (SLB)	8
Adherent Cells lysis	8
Suspension Cells lysis	9
B.2 AU calculation - Input lysate quantification	11
B.2.1 Measure Lysate AU	11
B2.2 Calculate the volume of lysate and diluting buffer needed for the pulldown	11
Step B3. Click reaction (material for 1 reaction)	12
APPENDIX	13
App.1 Lysis buffer supplementation issues	13

Kit storage info

	Qty.	Storage
4°C components	1 box	4°C
-20°C components	1 box	-20°C
USB pen drive	1 box	RT

 Table 1. Kit composition (in boxes) and storage temperature.

Additionally Required Materials

- RiboLock RNase inhibitor (Thermo Scientific catalog no. EO0381)
- Nuclease free water / DEPC water
- Dnase I (Thermo Scientific catalog no. EN0521)
- Methionine-free medium (e.g., for DMEM Thermo Scientific catalog no. 30030)
- Sodium deoxycholate 10% in nuclease-free water
- Nanodrop ND-1000 UV-VIS Spectrophotometer
- o microcentrifuge and nonstick RNase-free microcentrifuge tubes (1.5 mL)
- Automatic wheel (rotator)
- Magnetic separation device for 1.5 mL tubes
- o Mixer
- o Vortex
- o lodoacetamide

Mod 080264	v1.1	Date Exp/rev 17/03/2025	Edited by MA PB LM	Pag 3 of pag 14

INTRODUCTION

AHARIBO (AHA-mediated RIBOsome isolation) is an innovative proteogenomic analysis solution that effectively explores the quantitative relationship between transcript and protein levels by capturing active translation. This reliable and accurate approach overcomes the limitations of traditional methods like polysome profiling and affinity purification, which are often labor-intensive and yield poor correlations between mRNA and protein levels.

Our streamlined 360 approach simplifies the entire *de novo* synthesized peptide detection workflow, from sample to results. Users retain full experimental control, performing the peptide pull-down using the kit's wet lab components. Once ready, simply contact Immagina to schedule a convenient pick-up for delivery to our local partner. Immagina then handles the mass spectrometry acquisition and data analysis. Users access raw data, results, and reports via a dedicated portal. (See the included USB drive for complete guidelines.)

Each kit includes two pick-ups, allowing for the simultaneous submission of multiple samples. Additional shipments can be arranged at the user's expense. This kit is compatible with eukaryotic cell lines, both freshly harvested and flash-frozen. Immagina's mission is to develop unique and smart enabling technologies to break down the walls in complex Translational studies. Please visit our website https://immaginabiotech.com/ for a complete overview of our products and services and our proprietary technologies.



WORKFLOW OVERVIEW

Fig.1 Overview of the AHARIBO Protein System workflow. The cell manipulation is depicted in the workflow with the different components needed that are added during different time points (t0-t3) before pulldown of the newly synthesized peptide with the beads and subsequent peptide extraction. LC-MS is an example of a possible experiment that follow the pulldown.

N4 1 0000C 4		Data Fue / 17/02/2025		D
1000 080264	V1.1	Date Exp/rev 17/03/2025	Edited by IVIA PB LIVI	Pag 4 of pag 14

A. SAMPLE PREPARATION

A.1 Sample Recommendations

Reagents are optimized for experiments in 6-well plates, for different quantities, please scale up or down the amount of lysis buffer utilized in Step B1.

Typically, for experiments in 6-well plates, 150,000-250,000 adherent cells are seeded per well in 2 ml of cell growth medium. We recommend treating cells with AHA at 70 to 90% confluence when the cells are actively dividing. Please note that the amount of newly synthetize proteins with the translational machinery that can be isolated from a sample is strongly affected by its translational state and must be considered when programming experiments with the IMMAGINA AHARIBO Protein System 360. For instance, two lysates similarly concentrated (i.e., similar Abs260nm) but from different cell types or specimens (e.g. human vs mouse, or immortalized vs primary), or with different treatments (e.g. drugs and transfection reagents) could have completely different amounts of translating ribosomes, hence leading to different outcomes.

A.1.1. Cell Medium Recommendations

To prepare Methionine free complete medium, you should supplement the specific medium for your cell lines with all the components needed (e.g., Penicillin, L-glutamine, FBS, growth factors, etc.). If the Methionine-free medium that you purchase is also depleted from L-leucine, please add 10 μ L/mL of L-leucine (80 mM ready-to-use provided in the kit #IBT0441).

The complete Methionine-free medium must be prewarmed to 37°C or the right growing temperature before addition to the cells, to allow proper growth.

Mod 080264	v1.1	Date Exp/rev 17/03/2025	Edited by MA PB LM	Pag 5 of pag 14
			-	

Optimal Workflow Recommendations

- Please run up to 6 samples in parallel. Longer manipulation time may introduce an unwanted variability between the first and the last sample.
- Allocate at least 1 day for the completion of the entire workflow.
- This protocol has been optimized to perform all the reactions from cells growing at 37°C in a 6 well plate format. For other format and temperatures needed for your cell line, please adjust lysis buffer volume and temperature accordingly.
- The pulldown has been optimized for starting with 2 AU (Abs260nm) in 100 μL of diluted lysate.

Mod 080264	v1.1	Date Exp/rev 17/03/2025	Edited by MA PB LM	Pag 6 of pag 14
			•	

B. Pull-down of *de-novo* synthesized protein

AHARIBO Protein System 360 components and additional required materials needed in this section:

Step N	Kit component	Cat. nr.	Volume	Storage	Туре		Vial cap color
B1	Lysis buffer (LB)	# IBT0034	1 mL	-20°C	Vial	14 A 14 A 14 A	clear
B1	SDC 10%	Additionally Required Material					
B1	DNAse I	Additionally Required Material					
B1	RiboLock RNase Inhibitor	Additionally Required Material					
B1	sBlock	# IBT0451	50 µL	-20°C	Vial	14 A A A A A A A A A A A A A A A A A A A	clear
B1	Nuclease Free Water (NFW)	Additionally Required Material					
B1	L-azidohomoalanine (100 mM) (AHA)	# IBT0431	200 µL	-20°C	Vial	14 H H	clear
B1	L-Leucine (80mM) (LL)	# IBT0441	200 µL	-20°C	Vial		clear
B2	Washing Solution (WSS)	# IBT0461	15 mL	4°C	Bottle		
B2	Guanidinium Washing Solution (GWS)	# IBT0482	20 mL	4°C	Bottle		
B2	bBeads (bB)	# IBT0045	700 μL	4°C	Vial		Blue

Mod 080264	v1.1	Date Exp/rev 17/03/2025	Edited by MA PB LM	Pag 7 of pag 14

Step B1. CELL LYSIS

Before starting the experiment – Supplemented Lysis Buffer (SLB)

To ensure optimal reproducibility we recommend producing a fresh Supplemented Lysis Buffer (SLB) aliquot for each new experiment, right before proceeding with the Lysis Step. Combine the SLB by following Table 2 instructions and multiply the volumes according to the number of samples being processed (N). Please combine the different reagents following the left-to-right order. Note that 50 μ L of supplemented lysis buffer is the recommended volume for a 6 wells plate. For other sizes of dishes, please use proportional volumes.

	Lysis buffer (LB)	Sodium deoxycholate (SDC) 10% (W/V)	DNase I 1 U/µL	RiboLock RNase Inhibitor 40 U/µL	sBlock	Final Volume
N=1	43.5 µL	5 µL	0.25 μL	0.25 µL	0.5 µL	50 µL
N=						

 Table 2. Recipe for the supplementation of the provided lysis buffer.

The final concentration of reagents in the SLB is as follows: Sodium deoxycholate (1%), DNase I (5U/mL), and RiboLock RNase Inhibitor (200 U/mL).

Please if the SLB appears as a whiteish and cloudy solution, do not proceed and check Appendix 1.

Adherent Cells lysis

- □ B1.1a Remove the medium from the 6 well plate.
- B1.2a Wash cells (80% confluency) by adding 1 mL of PBS and removing it after the wash.
 Repeat the wash.
- □ B1.3a Add 1 mL/well of methionine-free medium (supplemented as needed) to the cells and incubate at 37°C for 40 minutes.
- $\Box~$ B1.4a Add 10 μL of AHA reagent to the cells, and incubate for 60 min at 37°C.
- □ B1.5a Place the plate on ice and wash the cells with 1mL/well of cold PBS.
- □ B1.6a Remove all residual PBS with a pipette. All the PBS must be removed before proceeding with the lysis to avoid diluting the lysis buffer.
- B1.7a Perform the lysis directly adding 45 µL of the complete Supplemented Lysis Buffer (Table 2) to each cell well and scrape vigorously. Mechanical scraping helps the downstream processing by disrupting the cell membrane and releasing the cellular contents, including ribosomes. To ensure good lysis, follow these guidelines for mechanical scraping:

Mod 080264	v1.1	Date Exp/rev 17/03/2025	Edited by MA PB LM	Pag 8 of pag 14
		• • • •		

- Before scraping, make sure you are working in a sterile environment using appropriate aseptic techniques.
- Prepare your sample by adding the necessary lysis buffer or solution as per the protocol.
- Using a suitable tool such as a cell scraper, spatula, or pipette tip, gently scrape the surface of the cell culture dish or tissue to dislodge the cells.
- Apply consistent but gentle pressure to ensure thorough scraping while avoiding excessive force that may introduce debris.
- Scrape in a systematic manner, covering the entire surface area to ensure an even distribution of lysed cells.
- Continue scraping until you observe the desired level of cell detachment and release of cellular material.
- Transfer the lysate to a suitable collection vessel, such as a microcentrifuge tube, for further processing or analysis.
- □ **B1.8a** Collect the cell lysate in a 1.5 mL microcentrifuge tube and pellet the cell debris and nuclei by centrifugation at 20,000 g for 5 min at 4°C.
- □ B1.9a Transfer the supernatant to a new tube and keep it on ice for 20 min.
- B1.10a Check the absorbance of the cell lysate at 260 nm, we suggest using a Nanodrop setting the "nucleic acid" function and using 1.5 μL of the supplemented lysis buffer as blank (for troubleshooting check A.3 AU calculation - Input lysate quantification).
- □ B1.11a incubate the cell lysate with 10mM final lodoacetamide for 45 min in the dark
- □ **B1.12a** If the sample is not processed the same day, please store the sample at -80°C or in a cryogenic storage system to maintain its stability until further processing.

Suspension Cells lysis

- □ B1.1a Collect the suspension cells (80% confluency) by pelleting them using a centrifuge (e.g., centrifugate at 300 g for 5 min at RT). Remove the supernatant.
- □ **B1.2a** Wash pelleted cells with 1 mL of PBS. Pellet them again by centrifugation and remove the supernatant. Repeat the wash and remove completely the supernatant after centrifugation.
- □ B1.3a Add 1 mL/well of methionine-free medium (supplemented as needed) to the cells and incubate at 37°C for 40 minutes.
- \square B1.4a Add 10 µL of AHA reagent to the cells, and incubate for 60 min at 37°C.
- B1.5a Collect the treated suspension cells by pelleting. Wash the cells with 1mL of cold PBS.
 Pellet them again by centrifugation.
- □ B1.6a Remove all residual PBS with a pipette. All the PBS must be removed before proceeding with the lysis to avoid diluting the lysis buffer.
- B1.7a Perform the lysis directly adding 45 µL of the complete Supplemented Lysis Buffer (Table 2) to the cell pellet. Lysate the cells pipetting up and down at least 30 times with a 200 µL pipette without generating bubbles.

March 00000C4		Data Frankan 17/02/2025		D 0 f 14
1000 080264	VI.I	Date Exp/rev 17/03/2025	Edited by IVIA PB LIVI	Pag 9 of pag 14

- □ **B1.8a** Collect the cell lysate in a 1.5 mL microcentrifuge tube and pellet the cell debris and nuclei by centrifugation at 20,000 g for 5 min at 4°C.
- □ B1.9a Transfer the supernatant to a new tube and keep it on ice for 20 min.
- □ B1.10a Check the absorbance of the cell lysate at 260 nm, we suggest using a Nanodrop setting the "nucleic acid" function and using 1.5 µL of the supplemented lysis buffer as blank (for troubleshooting check A.3 AU calculation Input lysate quantification). If the sample is not processed the same day, please store the sample at -80°C or in a cryogenic storage system to maintain its stability until further processing.
- □ B1.12a incubate the cell lysate with 10mM final lodoacetamide for 45 min in the dark
- □ B1.13a If the sample is not processed the same day, please store the sample at -80°C or in a cryogenic storage system to maintain its stability until further processing.

IMPORTANT!!

Negative control test.

To perform the negative control test, please use cell lysate of the same cell line not incubated with AHA. This **negative control lysate** will be used later as control for the pulldown experiment. Please remember that this will decrease the number of pulldowns for each single negative control experiment.

To avoid nonspecific binding, incubate the negative control lysate with 10mM final lodoacetamide for 45 min in the dark.

Mod 080264	v1.1	Date Exp/rev 17/03/2025	Edited by MA PB LM	Pag 10 of pag 14
				.0

B.2 AU calculation - Input lysate quantification

B.2.1 Measure Lysate AU

Cells should be lysed following Step B1 a, or b instructions depending on your specimen type. The AU of your sample is measured using a spectrophotometer, most commonly a Nanodrop. Set the instrument so to measure the Abs at 260 nm (usually Nucleic Acid function) and measure the absorbance of your lysate using the Supplemented Lysis Buffer (SLB) as blank (see Before starting the experiment – Lysis Buffer Supplementing & Table 3). The use of different lysis buffers is strongly discouraged because it may interfere with the efficiency of ribosome pull-down and with the AU calculation (some components may absorb at 260 nm).

<u>If the instrument does not allow the use of the SLB as blank</u>, please use water instead, then record the absorbance of both the SLB and the lysate and subtract the absorbance of the SLB to the lysate.

Example:

- \Box Supplemented Lysis buffer SLB Abs260nm = 7 AU
- \Box Specimen Abs260nm = 17 AU
- □ Absorbance value of lysate = 17 7 = 10 AU

B2.2 Calculate the volume of lysate and diluting buffer needed for the pulldown

The absorbance of your sample depends on your sample characteristics (type of cell and amount) if resuspended in the same volume of lysis buffer.

In Step X, to calculate the volume of lysate to dilute in WSS buffer to reach 2 AU in 100 μ L, follow the examples below.

Example 1: Nanodrop absorbance value of lysate at 260 nm = 10 AU.

- \Box 2 AU (final quantity) * 100 μ L (final volume) = 200 AU* μ L
- \Box 200 AU*µL /10 AU (lysate absorbance) = 20 µL of lysate to pipette
- \Box Put 20 µL of lysate in 80 µL of WSS buffer.

Example 2: Nanodrop absorbance value of lysate at 260 nm = 4 AU.

- \Box 2 AU (final quantity) * 100 μ L (final volume) = 200 AU* μ L
- \Box 200 AU*µL /4 AU (lysate absorbance) = 50 µL of lysate to pipette
- $\hfill\square$ Put 50 μL of lysate in 50 μL of WSS buffer.

Mod 080264	v1.1	Date Exp/rev 17/03/2025	Edited by MA PB LM	Pag 11 of pag 14

Step B3. Click reaction (material for 1 reaction)

- □ **B3.1** Remove the **bBeads** from 4°C and place the tube at RT for at least 15 min.
- B3.2 Dilute the lysate obtained at the end of STEP B1, with WSS to obtain a final value of 2 AU in 100 μL (see A2.2 Calculate the volume of lysate and diluting buffer needed for the pulldown section for more information).
- \square **B3.3** Add 50 µL of bBeads.
- □ **B3.4** Incubate for 180 min on a thermomixer 1000 rpm at room temperature.
- B3.5 Remove the tubes from the mixer. DO NOT CENTRIFUGATE but allow the entire solution with the beads to settle at the bottom of the tube. If residual solution is present on the lid, pull down the beads by gently flicking down the tube by hand 2 or 3 times.
- B3.6 Place the tube onto the magnet and let the beads collect for 2 3 minutes or until the supernatant is completely clear. If residual solution is present on the lid, pull down the beads by gently flicking down the magnet 2 or 3 times Remove and discard the supernatant with a pipette while the tube remains in contact with the magnet.
- $\hfill\square$ **B3.7** Add 700 μL of **GWS** solution to the beads.
- □ **B3.8** Incubate for 15 minutes on a thermomixer at 1,000 rpm at RT.
- □ **B3.9** Place the tube onto the magnet and remove and discard the supernatant with a pipette while the tube remains in contact with the magnet.
- □ B3.10 Repeat the steps from B3.7 to B3.9
- \square **B3.11** Wash the beads 2 times with 500 µL of NFW, 2 min each.
- \square **B3.12** Resuspend the beads in 200 µL of NFW.
- B3.13 Polypeptides are bound to the beads, transfer the suspension (beads in water) to a new
 1.5 mL tube. Store the Eppendorf containing the beads in water at -80°C until shipment.
- B3.14 Now you are ready to ship your samples to Immagina for proteomic analysis. Please refer to the guideline file found on the USB key included in your kit for detailed instructions on sample preparation and shipping.

Mod 080264	v1.1	Date Exp/rev 17/03/2025	Edited by MA PB LM	Pag 12 of pag 14
		2 ate 1, p) et 1, j ee, 1010		. «B == 0. p«B =

APPENDIX

App.1 Lysis buffer supplementation issues

Please check if, after adding Sodium deoxycholate a whiteish and cloudy solution appears. If so, please do not proceed with the lysis of the sample and toss the supplemented LB. Subsequently, warm up the SDC at RT and add it to a new aliquot of the not-supplemented LB. If the whiteish and cloudy solution persists, please contact our tech support (<u>techsupport@immaginabiotech.com</u>).

	Mod 080264	v1.1	Date Exp/rev 17/03/2025	Edited by MA PB LM		Pag 13 of pag 14
--	------------	------	-------------------------	--------------------	--	------------------

Contacts



Info

info@immaginabiotech.com

Sale support (quoting, ordering, and order status update) orders@immaginabiotech.com

Technical service (technical inquiries and quality complaints) techsupport@immaginabiotech.com



Viale dell'Industria, 47, 38057, Pergine Valsugana (TN), ITALY



https://www.immaginabiotech.com



Notes:

Mod 080264	v1.1	Date Exp/rev 17/03/2025	Edited by MA PB LM	Pag 14 of pag 14