



© Automated Protein Normalization and Tryptic Digestion on a Biomek-NX Liquid Handler System V.2



Jan 12, 2022

Yan Chen¹, Tad Ogorzalek¹, Nurgul Kaplan Lease¹, Jennifer Gin¹, Christopher J Petzold¹

¹Lawrence Berkeley National Laboratory



dx.doi.org/10.17504/protocols.io.b3gtqjwn

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This protocol details steps to normalize the amount of protein for tryptic digestion in quantitative proteomic workflows by using a Biomek NX liquid handler system. It is optimized to normalize protein concentrations in a 96-well plate format and add TCEP, IAA, and trypsin.

This protocol works best as part of a semi-automated proteomic sample preparation workflow with:

<u>Automated Chloroform-Methanol Protein Extraction on the Biomek-FX Liquid Handler System</u>

and

<u>Automated Protein Quantitation with the Biomek-FX liquid handler system</u>

DOI

dx.doi.org/10.17504/protocols.io.b3gtqjwn

Yan Chen, Tad Ogorzalek, Nurgul Kaplan Lease, Jennifer Gin, Christopher J Petzold 2022. Automated Protein Normalization and Tryptic Digestion on a Biomek-NX Liquid Handler System. **protocols.io** https://dx.doi.org/10.17504/protocols.io.b3gtqjwn Yan Chen







Modular automated bottom-up proteomic sample preparation for highthroughput applications

Automation, Proteomics, Tryptic digestion, Biomek, Normalization, Sample preparation

protocol ,

Jan 04, 2022

Jan 12, 2022

Christopher Petzold Lawrence Berkeley National Laboratory

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Part of collection

Modular automated bottom-up proteomic sample preparation for high-throughput applications

- A Beckman-Coulter Biomek NX-S8 or NXP liquid handler system with an 8-pod head is used for this protocol. Alternative liquid handlers can be used with appropriate method development.
- Because different deck orientations and system components are possible, you will need to modify the method file (attached in the 'Before start' section) for your specific Biomek liquid handler system.

PCR Plate 96-well non-skirted Thermo Fisher Scientific Catalog #AB0600
PCR Tube Storage Rack Axygen Catalog #R96PCRFSP
Ammonium Bicarbonate LC-MS grade VWR Scientific Catalog #BJ40867-50G
20 uL pipet tips Molecular Bioproducts BioRobotix, Catalog #918-262
200 uL pipet tips Molecular Bioproducts BioRobotix, Catalog #919-262
Tris(2-carboxyethyl)phosphine hydrochloride (TCEP), SigmaAldrich, Catalog #C4706
Iodoacetamide, MilliporeSigma, Catalog #I1149
Trypsin, SigmaAldrich, Catalog #T6567-1MG

Wear proper PPE (gloves, safety goggles, and lab coat), and prepare solvents in a chemical fume hood.

Store organic solvents in a flammable storage cabinet when not in use. Discard used solvents and buffers in appropriate waste containers.

For this protocol you will need:



- A Beckman-Coulter Biomek NX-S8 or NXP liquid handler system with a 8-pod head
- Upload the attached method files and modify them to fit your deck and system configuration.
- **(i) Proteomics-Normalization Method.bmf**
- Proteomics-Plate to Plate Transfer.bmf
- Protein samples of known concentration

Chemicals to prepare:

- Prepare
 [M]100 Milimolar (mM) Tris(2-carboxyethyl)phosphine (TCEP) solution by dissolving □28.7 mg TCEP in □1 mL 100mM Ammonium Bicarbonate
- Prepare [M]200 Milimolar (mM) Iodoacetamide (IAA) solution by dissolving
 ■36.8 mg Iodoacetamide in ■1 mL 100mM Ammonium Bicarbonate
- Prepare [M]1 mg/ml Trypsin by adding □1 mL 1mM HCl to □1 mg Trypsin then vortex to mix

Store TCEP, IAA, and Trypsin in -20C. IAA is light sensitive. Store in amber tube (Fisher Scientific, Cat.#05-402-31).

Biomek NX-S8 input file preparation

- 1 After measuring protein concentration by the DC (Detergent Compatible) protein assay (Bio-Rad), export protein concentration report through MD Spectramax 250 software that controls the microplate reader. Copy the content in the exported text file and paste it into Excel, and then save as a UTF-8 format text file.
 - Example protein concentration file.txt
- 2 Use MS Excel or a Jupyter notebook to normalize the protein concentration to **□50 μg** and convert the spectrophotometer output file into the following two files in a format suitable for the Biomek NX-S8:

NX-AMBIC.csv

Α	В	С	D	E
scrpos	srcwell	destpos	destwell	vol
media	1	DestPlate	1	33.05
media	1	DestPlate	2	26.73
media	1	DestPlate	3	27.96
media	1	DestPlate	4	28.74
media	1	DestPlate	5	22.94
media	1	DestPlate	6	28.07
media	1	DestPlate	7	24.34
media	1	DestPlate	8	28.12
media	1	DestPlate	9	26.64
media	1	DestPlate	10	26.22

NX-AMBIC.csv output table

NX-protein.csv

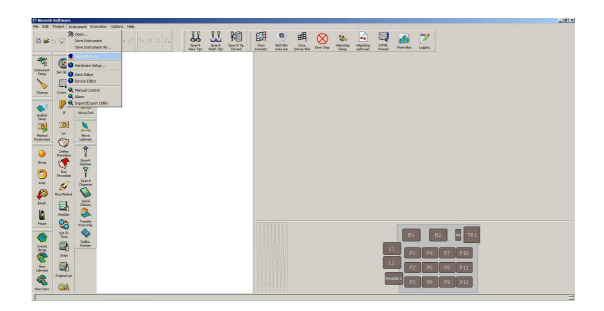
Α	В	С	D	E
scrpos	srcwell	destpos	destwell	vol
SrcPlate	1	DestPlate	1	10.95
SrcPlate	2	DestPlate	2	17.27
SrcPlate	3	DestPlate	3	16.04
SrcPlate	4	DestPlate	4	15.26
SrcPlate	5	DestPlate	5	21.06
SrcPlate	6	DestPlate	6	15.93
SrcPlate	7	DestPlate	7	19.66
SrcPlate	8	DestPlate	8	15.88
SrcPlate	9	DestPlate	9	17.36
SrcPlate	10	DestPlate	10	17.78

NX-protein.csv output table

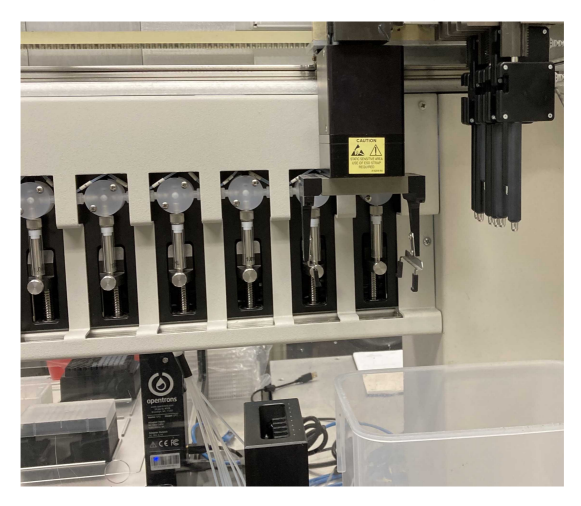
Biomek NX-S8 preparation

3 Open Biomek software program from Biomek NX-S8 control computer. In the "Instrument" drop-down menu, select "Home all Axes" to prepare the instrument for use and purge air from the tubing and syringes.

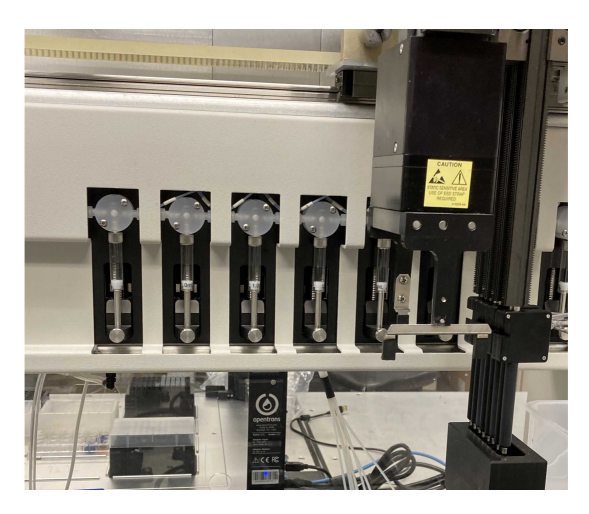




Biomek software that controls the operation of Biomek NX-S8 liquid handler system



Homing all axes



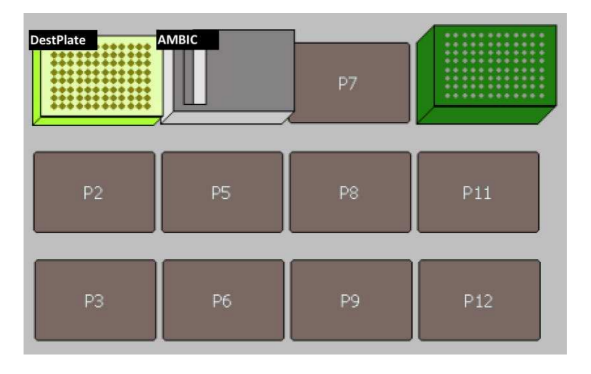
Purging air

Buffer Transfer

4 Go to "Open Method." Select the "Proteomics" folder and open the method "Proteomics-Normalization Method."

Because different deck orientations and system components are possible, you will need to modify the method file (attached in the 'Before start' section) for your specific Biomek liquid handler system.

5 Click on "Instrument Setup."



Deck setup

6 Set up the deck (refer to the deck setup picture above):

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Α	В	С
Deck Label	Labware	Reagent
DestPlate	PCR plate 96-well non-skirted (Thermo	
	Fisher, Cat.#AB0600) on a yellow PCR rack	
AMBIC	Biomek Reservoir (discontinued)	Ammonium Bicarbonate buffer
tips	200 uL pipet tips (Molecular Bioproducts	
	BioRobotix, Cat.#919-262)	

Deck materials

- 7 Click on "Transfer From File."
- 8 Copy the NX-AMBIC.csv file generated in Excel or via a Jupyter Notebook into the directory that the method is designated to read. For example C:\Users\jbei\Desktop\Proteomics Methods\CSV files\

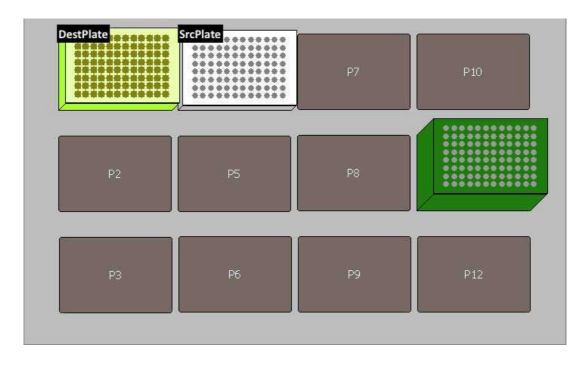
- 9 Click on the 2nd "View Datasets" to check that you have copy and pasted the correct volumes in the 96-well format.
- 10 Click "Finish" to make sure there are no error messages.

11 Click the "Run" button (green arrow) to start.

5m

Protein Transfer

- 12 Go to "Open Method." Select the "Proteomics" folder and open the method "Proteomics-Plate to Plate Transfer."
- 13 Click on "Instrument Setup."



Deck setup

14 Set up the deck (refer to the deck setup picture above):

30s

Α	В	С
Deck Label	Labware	Reagent
DestPlate	PCR plate 96-well non-skirted (Thermo Fisher, Cat.#AB0600) on a yellow PCR rack	Ammonium bicarbonate buffer
SrcPlate	PCR plate 96-well non-skirted (Thermo Fisher, Cat.#AB0600) on a yellow PCR rack	protein
tips	20 uL pipet tips (Molecular Bioproducts BioRobotix, Cat.#918-262) or	
	200 uL pipet tips (Molecular Bioproducts BioRobotix, Cat.#919- 262)	

Deck materials

- 15 Click on "Transfer From File."
- Copy the **NX-protein.csv** file generated by Excel or via a Jupyter Notebook into the directory that the method is designated to read. For example: **C:\Users\jbei\Desktop\Proteomics**Methods\CSV files\
- 17 Click on "View Datasets" to check that you have copy and pasted the correct volumes in the 96well format.
- 18 Click "Finish" to make sure there are no error messages.
- MANUAL STEP: Use a multichannel pipette to mix protein samples completely right before starting.
- 20 Click the Run button (green arrow) to start.

8m

Trypsin Digestion

21 Chemicals to prepare:

protocols.io

- Prepare [M]100 Milimolar (mM) Tris(2-carboxyethyl)phosphine (TCEP) solution by dissolving ■28.7 mg TCEP in ■1 mL 100mM Ammonium Bicarbonate
- Prepare [M]200 Milimolar (mM) Iodoacetamide (IAA) solution by dissolving

 □36.8 mg Iodoacetamide in □1 mL 100mM Ammonium Bicarbonate
- Prepare [M]1 mg/ml Trypsin by adding □1 mL 1mM HCl to □1 mg Trypsin then vortex to mix

Store TCEP, IAA, and Trypsin in -20C. IAA is light sensitive. Store in amber tube (Fisher Scientific, Cat.#05-402-31).

- 22 Add the following reagents to the normalized protein plate, in this order:
 - 1. **2.5** μl 100 mM TCEP
 - 2. **2.5 μl 200 mM IAA**
 - 3. **1 µl 1 mg/mL Trypsin**

The final concentrations will be 1 μ g/ μ l protein (in 50 μ l total volume), 5 milimolar (mM) TCEP, 10 milimolar (mM) IAA , and 1 μ l Trypsin (1 mg/ml) (1:50 trypsin:protein ratio). Adjust as needed for your data acquisition protocols.

23 Incubate at § 37 °C for © 04:00:00 to © 16:00:00 .

20h

Clarifying Digestion Reaction

15m

24 After digestion, spin at **(3)14000 rpm, 00:15:00** .

15m

Pipet supernatant into new PCR plate. Seal and store at 8 -20 °C until ready for LC-MS/MS analysis (e.g., <u>Discovery Proteomics protocol</u>, <u>Targeted Proteomics protocol</u>).

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Citation: Yan Chen, Tad Ogorzalek, Nurgul Kaplan Lease, Jennifer Gin, Christopher J Petzold Automated Protein Normalization and Tryptic Digestion on a Biomek-NX Liquid Handler System https://dx.doi.org/10.17504/protocols.io.b3gtgjwn

