

Sep 25, 2023

Opentrons Pipeline: DNA Extraction with the Omega Biotek Stool Kit

DOI

dx.doi.org/10.17504/protocols.io.dm6gp39njvzp/v1



Kristina N Vsevolodova¹, Gideon Erkenwick², Mrinalini Watsa¹

¹San Diego Zoo Wildlife Alliance; ²Field Projects International

In Situ Laboratories
Tech. support email: info@insitulabs.org



Kristina N Vsevolodova

San Diego Zoo Wildlife Alliance

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Protocol Citation: Kristina N Vsevolodova, Gideon Erkenwick, Mrinalini Watsa 2023. Opentrons Pipeline: DNA Extraction with the Omega Biotek Stool Kit . **protocols.io** <https://dx.doi.org/10.17504/protocols.io.dm6gp39njvzp/v1>

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Protocol status: Working

Created: September 19, 2023

Last Modified: September 25, 2023

Protocol Integer ID: 88037

Keywords: Opentrons, OT2, DNA extraction, Extraction, Omega Biotek, Stool , Stool extraction, insitulabs, automated, dna extraction with the omega biotek stool kit, omega biotek stool kit, opentrons pipeline, dna extraction, opentron, automated pipetting robot, pipetting robot, pipeline, automated pipeline, extraction, well plate of dna, stool lysate, omega biotek, 200ul opentron, dna, stool, robot, filtered tip box, lysing of the stool

Funders Acknowledgements:

Gordon and Betty Moore Foundation

Grant ID: 9772

Revive and Restore Catalyst Science Fund

Grant ID: 2021-024

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Abstract

This protocol is an automated pipeline to extract a full 96-well plate of DNA from stool lysates prepared in two separate plates. Lysing of the stool can be performed as you wish, depending on species and their diet. The protocol itself begins after lysates have been created.

This protocol was developed and optimized for the following:

Platform: Opentrons OT-2 automated pipetting robot

Kit: Omega Biotek

Tips Used: 4 boxes (4 × 200uL Opentrons Filtered Tip boxes)

Recommended number of samples: 96



Guidelines

This protocol is recommended for a full 96-well plate. Do not use it for less than 12 columns. A version for variable sample numbers is forthcoming in the next version.

Things to consider:

Before beginning: Load your lysates into two plates, a 1 mL deep well with 300uL of lysate and a 200uL Plate with 200uL of lysate. This is because, during the COVID pandemic, locating 2mL deep well plates for small labs (i.e. to buy in numbers less than 10,000) was impossible to do. So, we made our protocol flexible to be able to use a larger amount of lysate (total of 500uL) without the 2mL deepwell plate.

Step 4: The 50mL Falcon tubes are recommended as means of sterile transport of accurate reagent amounts between the lab and the OT-2. They can be replaced with a different brand of tube or reservoir.

Step 6: Import the labware file BEFORE you import your protocol or it will give an error. This protocol has been validated against Opentrons software app version 6.2.1

Step 7.2: In the newer software versions for the OT-2, calibration is not required before every run. In some cases the user will not be asked to calibrate the machine, and they should not have to.

Materials

- 5x Nest 50mL Falcon Tubes
- 4x Opentrons 200µL Filter Tips
- 1x VWR 96 Deep Well Plates 1mL
- 2x NEST 1-Well Reservoirs, 195 mL
- 2x NEST 12-Well Reservoirs, 15 mL
- 1x 96-Well PCR Plate Non-skirt, 200µl
- 3x Nest skirted PCR Plate
- 2x Aluminium Seals
- 6× 2mL Tubes
- 100-1000µL pipette
- 1000µL pipette tips
- Incubator or water bath that can reach 70°C or more



Protocol materials

☒ 10% Bleach

☒ Distilled Water

☒ 70% Alcohol

☒ 70% Alcohol

☒ XP2 Binding Buffer **Omega Biotek Catalog #PDR040**

☒ Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50**

☒ XP2 Binding Buffer **Omega Biotek Catalog #PDR040**

☒ Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50**

☒ VHB Buffer **Omega Biotek Catalog #VHB-440**

☒ SPM Buffer **Omega Biotek Catalog #SPM-300**

☒ SPM Buffer **Omega Biotek Catalog #SPM-300**

☒ Elution Buffer **Omega Biotek Catalog #PDR048**

☒ SPM Buffer **Omega Biotek Catalog #SPM-300**

☒ VHB Buffer **Omega Biotek Catalog #VHB-440**

☒ SPM Buffer **Omega Biotek Catalog #SPM-300**

☒ Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50**

☒ Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50**

☒ Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50**

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☒ Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50**

☒ Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50**

☒ Elution Buffer **Omega Biotek Catalog #PDR048**

☒ SPM Buffer **Omega Biotek Catalog #SPM-300**

☒ VHB Buffer **Omega Biotek Catalog #VHB-440**

☒ XP2 Binding Buffer **Omega Biotek Catalog #PDR040**

☒ Elution Buffer **Omega Biotek Catalog #PDR048**

☒ Qubit® dsDNA HS Assay Kit **Thermo Fisher Scientific Catalog #Q32854**

☒ Qubit® 3.0 Fluorometer **Thermo Fisher Scientific Catalog #Q33216**

☒ Quantus(TM) NGS Starter Package **Promega Catalog #E5150**

☒ Quantus(TM) Fluorometer **Promega Catalog #E6150**



Quant-iT™ PicoGreen® dsDNA Assay Kit **Life Technologies Catalog #P11496**

Nuclease-Free Water, 150ml **Promega Catalog #P1195**

Mag-Bind Stool DNA 96 Kit **Omega Biotek Catalog #M4016-01**

Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50**

100% Ethanol

Elution Buffer **Omega Biotek Catalog #PDR048**

Elution Buffer **Omega Biotek Catalog #PDR048**

Elution Buffer **Omega Biotek Catalog #PDR048**

Elution Buffer **Omega Biotek Catalog #PDR048**

10% Bleach

Distilled Water

70% Alcohol

100% Ethanol

100% Ethanol

100% Ethanol

100% Ethanol

Troubleshooting

Before start

Clean the OT2 deck and walls with:

10% Bleach 1 rinse

Distilled Water 1 rinse

70% Alcohol 2 rinses

Note

Avoid wetting electronic parts.



Before starting

1 Ingredients

⊗ 100% Ethanol

⊗ Nuclease-Free Water, 150ml **Promega Catalog #P1195**

⊗ Mag-Bind Stool DNA 96 Kit **Omega Biotek Catalog #M4016-01** (Contains everything for extraction including lysis reagents)

OR separate reagents (not including lysis reagents):

- ⊗ SPM Buffer **Omega Biotek Catalog #SPM-300**
- ⊗ VHB Buffer **Omega Biotek Catalog #VHB-440**
- ⊗ XP2 Binding Buffer **Omega Biotek Catalog #PDR040**
- ⊗ Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50**
- ⊗ Elution Buffer **Omega Biotek Catalog #PDR048**

Note

For QC on individual level the

⊗ Qubit® dsDNA HS Assay Kit **Thermo Fisher Scientific Catalog #Q32854** on a

⊗ Qubit® 3.0 Fluorometer **Thermo Fisher Scientific Catalog #Q33216** can be used

OR the ⊗ Quantus(TM) NGS Starter Package **Promega Catalog #E5150** on a


⊗ Quantus(TM) Fluorometer **Promega Catalog #E6150**

For QC on plate level use the

⊗ Quant-iT™ PicoGreen® dsDNA Assay Kit **Life Technologies Catalog #P11496**

1.1 For the 1×96 Kit

Dilute SPM Buffer with  70 mL ⊗ 100% Ethanol per bottle and store at

 Room temperature .



Dilute VHB Buffer with  28 mL  100% Ethanol per bottle and store at  Room temperature .

1.2 For the 4×96 Kit

Dilute SPM Buffer with  70 mL  100% Ethanol per bottle and store at  Room temperature .

Dilute VHB Buffer with  112 mL  100% Ethanol per bottle and store at  Room temperature .

2 Materials

5x Nest 50mL Falcon Tubes for holding accurate reagent amounts before starting protocol.

4x Opentrons 200µL Filter Tips

1x VWR 96 Deep Well Plates 1mL for 300µL of lysate.

2x NEST 1-Well Reservoirs, 195 mL for extraction waste collection during protocol.

2x NEST 12-Well Reservoirs, 15 mL for holding reagents during protocol.

1x 96-Well PCR Plate Non-skirt, 200µL for 200µL of lysate.

3x Nest skirted PCR Plate for holding non-skirt plate and for final elutions.

2x Aluminium Seals

6× 2mL Tubes

100-1000µL pipette

1000µL pipette tips

Incubator or water bath that can reach 70°C or more

- 2.1 Autoclave the NEST 1-Well Reservoirs, 195 mL and NEST 12-Well Reservoirs, 15 mL before use. These can be rinsed and autoclaved and reused and need not be purchased new for each extraction. Slight yellowing of product can occur, but we do not see that it affects final outcomes in any discernible way.

3 Opentrons Equipment List

Equipment

OT-2	NAME
Liquid handler	TYPE
Opentrons	BRAND
OT-2	SKU

On the right pipette mount use the P300M

Equipment

OT-2 8 Channel Electronic Pipette	NAME
Pipette	TYPE
Opentrons	BRAND
P300M	SKU
https://shop.opentrons.com/8-channel-electronic-pipette/	LINK

Magnetic Module to place in Slot 7

Equipment

OT-2 Magnetic Module GEN2

NAME

Module

TYPE

Opentrons

BRAND

999-00098

SKU

<https://shop.opentrons.com/magnetic-module-gen2/> LINK

Prepare reagents

- 4 After the reagents are properly diluted and materials are ready, prepare the following amounts:

A	B	C	D
Item Name	Amount per sample [uL]	Amount for 96 samples [uL]	Amount for 96 samples * 1.1 overage[uL]
Mag-bind Bead Particles CH Round 1	10	960	1056
XP2 Binding Buffer Round 1	300	28800	31680
Mag-bind Bead Particles CH Round 2	10	960	1056
XP2 Binding Buffer Round 2	300	28800	31680
Wash 1: VHB Buffer	400	38400	42240
Wash 2: SPM Buffer	400	38400	42240
Wash 3: SPM Buffer	400	38400	42240
Elution Buffer	100	9600	10560



- 4.1 Fill one **Nest 50mL Falcon Tube** with the amount of
 XP2 Binding Buffer **Omega Biotek Catalog #PDR040** Round 1 and
 Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50** Round 1 required in
the table, and mix well. Keep at Room temperature . Split between two tubes if
needed.
- 4.2 Fill one **Nest 50mL Falcon Tube** with the amount of
 XP2 Binding Buffer **Omega Biotek Catalog #PDR040** Round 2 and
 Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50** Round 2 required in
the table, and mix well. Keep at Room temperature . Split between two tubes if
needed.
- 4.3 Fill one **Nest 50mL Falcon Tube** with the volume indicated in the table with wash 1:
 VHB Buffer **Omega Biotek Catalog #VHB-440** . Keep at Room temperature .
- 4.4 Fill one **Nest 50mL Falcon Tube** with the volume indicated in the table with wash 2:
 SPM Buffer **Omega Biotek Catalog #SPM-300** . Keep at Room temperature .
- 4.5 Fill one **Nest 50mL Falcon Tube** with the volume indicated in the table with wash 3:
 SPM Buffer **Omega Biotek Catalog #SPM-300** . Keep at Room temperature .
- 4.6 Distribute Elution Buffer **Omega Biotek Catalog #PDR048** between each 2mL tube
evenly.
- 4.7 Set an incubator or water bath to 70 °C and heat
 Elution Buffer **Omega Biotek Catalog #PDR048** in the incubator to 70 °C .

OT-2 script definitions

5 Definition of samples and labware:

5.1 Lysed Sample Plate 2 200uL

Remainder of lysed samples that will be added to Slot 7 in second round of bead incubation.



Position: Slot 2, **96-Well PCR Plate Non-skirt, 200µl** with 200uL of sample lysis on top of an empty **Nest skirted PCR Plate** (used as a base)

Name in the Deck: Lysis plate 2

Labware name in the protocol: denvillewithaxxygenbase_96_wellplate_200ul

Sample name in the script: lysate

5.2 **Lysed Sample Plate 1 300uL**

Lysed samples that will undergo the first round of bead incubation.

Position: Slot 7, **Opentrons Magnetic Module** with **VWR 96-Well Deep Well Plate** full of 300uL of sample lysis on top

Name in the Deck: Lysis plate 1

Labware name in the protocol: vwr_96_wellplate_1000ul

Sample name in the script: magsamps

5.3 **Sample Elution 1**

Samples that have been eluted from the beads for the first round elution 1.

Position: Slot 3, Empty **Nest skirted PCR Plate** (to receive elution 1)

Name in the Deck: Sample Elution Plate 1

Labware name in the protocol: nest_96_wellplate_100ul_pcr_full_skirt

Sample name in the script: eluates

5.4 **Sample Elution 2**

Samples that have been eluted from the beads for the second round elution 2.

Position: Slot 2, **96-Well PCR Plate Non-skirt, 200µl** with 200uL of sample lysis on top of an empty **Nest skirted PCR Plate** is replaced with new **Nest skirted PCR Plate** (to receive elution 2)

Name in the Deck: Sample Elution Plate 2

Labware name in the protocol: nest_96_wellplate_100ul_pcr_full_skirt


Sample name in the script: eluates2

Prepare the OT-2

- 6 Before loading your protocol, load the following labware files into your Opentrons app:



This labware definition allows us to use a non-skirted plate in the Opentrons app by inserting it into a skirted plate, and also allows us to use a 200uL plate (where our skirted plates that clip in are only 100uL. Feel free to replace with your own labware here).

 vwr_96_wellplate_1000ul.json 11KB

This labware definition is for the **1mL deepwell plate** from VWR. Note the rounded wells work well with the magnet.

Load this python file to the Opentrons app:  OT2_Omegabiotekfecal_v4.0.py 13KB

6.1 Definition of Protocol Variables:

This protocol is written per column, best working for multiples of 8. Therefore, if you want to modify the sample number just open the script in a text editor program, and modify the following value in line 3 of the script:

"numSamps": 96 → Indicates the number of samples that you will process.

Note

This protocol is recommended for a full plate of 96 samples (12 columns). Any less columns are not optimal.

7 Arrange the OT-2 deck

7.1 Slot 1: **NEST 12-Well Reservoirs, 15 mL** with reagents preloaded in the following order:

	A	B	C	D	E	F	G	H	I	J	K	L
	Well 1	Well 2	Well 3	Well 4	Well 5	Well 6	Well 7	Well 8	Well 9	Well 10	Well 11	Well 12
	Wash 3: SP M Buffer	Wash 3: SP M Buffer	Wash 3: SP M Buffer	Wash 3: SP M Buffer	EMPTY	EMPTY	EMPTY	EMPTY	EMPTY	EMPTY	EMPTY	Elution Buffer (when prompted)

Slot 2: **96-Well PCR Plate Non-skirt, 200µl** with 200uL of sample lysis on top of an empty **Nest skirted PCR Plate**

Slot 3: Empty **Nest skirted PCR Plate** (to receive elution 1)

Slot 4: **NEST 12-Well Reservoirs, 15 mL** with reagents preloaded in the following order:



	A	B	C	D	E	F	G	H	I	J	K	L
	Well 1	Well 2	Well 3	Well 4	Well 5	Well 6	Well 7	Well 8	Well 9	Well 10	Well 11	Well 12
	XP2 Binding Buffer and Magbind Particles Round 1 then 2	XP2 Binding Buffer and Magbind Particles Round 1 then 2	XP2 Binding Buffer and Magbind Particles Round 1 then 2	EMPTY	Wash 1: VH Buffer	Wash 1: VH Buffer	Wash 1: VH Buffer	Wash 1: VH Buffer	Wash 2: SPM Buffer	Wash 2: SPM Buffer	Wash 2: SPM Buffer	Wash 2: SPM Buffer

Slot 5: Opentrons 200µL Filter Tips**Note**

It is possible to use Opentrons 200µL Filter Tips or Opentrons 300 Tips (as in the image below). We usually use Opentrons 200µL Filter Tips to avoid cross contamination. The tips are in fact exactly the same dimensions, except that the P200F has a filter, while the P300 does not, and is therefore able to hold more liquid.

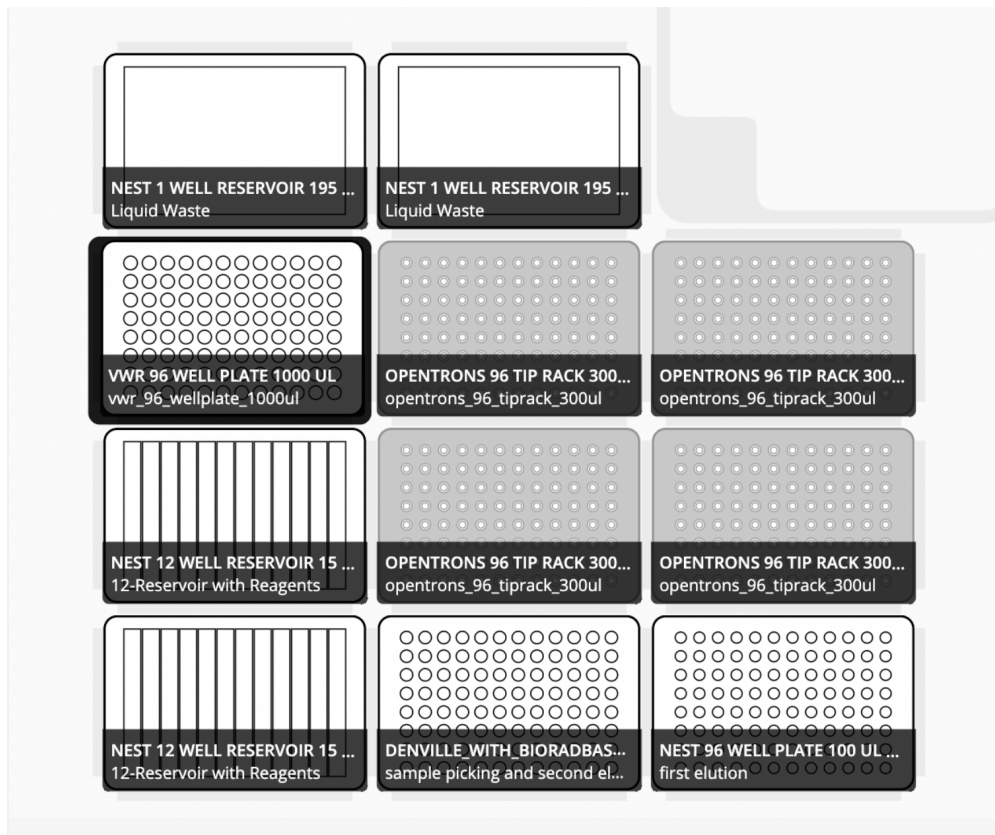
Slot 6: Opentrons 200µL Filter Tips

Slot 7: Opentrons Magnetic Module with VWR 96-Well Deep Well Plate full of 300uL of sample lysis on top

Slot 8: Opentrons 200µL Filter Tips**Slot 9: Opentrons 200µL Filter Tips**

Slot 10: NEST 1-Well Reservoirs, 195 mL (for waste)

Slot 11: NEST 1-Well Reservoirs, 195 mL (for waste)




Placement of LABWARE and TIPS in the OT2 Deck used for the Omega Biotek Stool extraction protocol. These materials are for purifying 96 samples.

7.2 Calibrate the deck if needed. Follow the on screen instructions.

Run the OT-2 protocol

1h 9m

7.3 Mixing Buffer and Particles with Lysed Sample Plate 1

Column 2 of tips in **Slot 8** will align with column 1 in **Slot 7** to mix the sample by aspirating and dispensing  10 μ L . The tips will then be returned to their original starting point. Each subsequent column of tips will continue on the same pattern until all of the samples are mixed.

7.4 Allowing beads to settle on the magnet

10m

The Opentrons Magnetic Module is engaged and incubates the mixed samples for


 00:10:00 .


7.5 Removing the supernatant

The supernatant is removed in two steps very gently to avoid removing settled beads. Supernatant is discarded in the Liquid waste NEST 1-Well Reservoir, 195 mL in Slot 10. Column 2 of tips in **Slot 8** will align with column 1 in **Slot 7** to remove the supernatant and then will return the tips back to their original starting point. Each subsequent column of tips will continue on the same pattern until all of the samples have their supernatant removed.


7.6 Adding VHB Buffer to Lysed Sample Plate 1 for wash 1

Column 2 of tips in **Slot 5** will transfer

 VHB Buffer **Omega Biotek Catalog #VHB-440** from wells 5-8 in **Slot 4** in two

 133 µL transfer steps to each sample in Lysed Sample Plate 1 in **Slot 7** without touching the lysates. The tips will then be dropped into the waste container.


7.7 Mixing VHB with Lysed Sample Plate 1

Column 2 of tips in **Slot 8** will align with column 1 in **Slot 7** to mix the sample by aspirating and dispensing  10 µL . The tips will then be returned to their original starting point. Each subsequent column of tips will continue on the same pattern until all of the samples are mixed.

7.8 Incubating VHB Buffer and lysate

4m

The Opentrons Magnetic Module is engaged and incubates the mixed samples for


 00:04:00 .


7.9 Removing the supernatant from the wash

The supernatant is removed in two steps very gently to avoid removing settled beads. Supernatant is discarded in the Liquid waste NEST 1-Well Reservoir, 195 mL in Slot 10. Column 2 of tips in **Slot 8** will align with column 1 in **Slot 7** to remove the supernatant and then will return the tips back to their original starting point. Each subsequent column of tips will continue on the same pattern until all of the samples have their supernatant removed.

7.10 Adding SPM Buffer to Lysed Sample Plate 1 for wash 2


Column 3 of tips in **Slot 5** will transfer

 SPM Buffer **Omega Biotek Catalog #SPM-300** from wells 9-12 in **Slot 4** two

 133 µL transfer steps to each sample in Lysed Sample Plate 1 in **Slot 7** without touching the lysates. The tips will then be dropped into the waste container.




7.11 Mixing SPM with Lysed Sample Plate 1

Column 2 of tips in **Slot 8** will align with column 1 in **Slot 7** to mix the sample by aspirating and dispensing  10 μL . The tips will then be returned to their original starting point. Each subsequent column of tips will continue on the same pattern until all of the samples are mixed.

7.12 Incubating VHB Buffer and lysate

4m

The Opentrons Magnetic Module is engaged and incubates the mixed samples for


 00:04:00 .


7.13 Removing the supernatant from the wash

The supernatant is removed in two steps very gently to avoid removing settled beads. Supernatant is discarded in the Liquid waste NEST 1-Well Reservoir, 195 mL in Slot 10. Column 2 of tips in **Slot 8** will align with column 1 in **Slot 7** to remove the supernatant and then will return the tips back to their original starting point. Each subsequent column of tips will continue on the same pattern until all of the samples have their supernatant removed.


7.14 Adding SPM Buffer to Lysed Sample Plate 1 for wash 3

Column 4 of tips in **Slot 5** will transfer

 SPM Buffer Omega Biotek Catalog #SPM-300 from wells 1-3 in **Slot 1** in two

 133 μL transfer steps to each sample in Lysed Sample Plate 1 in **Slot 7** without touching the lysates. The tips will then be dropped into the waste container.


7.15 Mixing SPM with Lysed Sample Plate 1

Column 2 of tips in **Slot 8** will align with column 1 in **Slot 7** to mix the sample by aspirating and dispensing  10 μL . The tips will then be returned to their original starting point. Each subsequent column of tips will continue on the same pattern until all of the samples are mixed.

7.16 Incubating SPM Buffer and lysate

4m

The Opentrons Magnetic Module is engaged and incubates the mixed samples for

 00:04:00 .


7.17 Removing the supernatant from the wash



The supernatant is removed in two steps very gently to avoid removing settled beads. Supernatant is discarded in the Liquid waste **NEST 1-Well Reservoir, 195 mL** in Slot 11. Column 2 of tips in **Slot 8** will align with column 1 in **Slot 7** to remove the supernatant and then will return the tips back to their original starting point. Each subsequent column of tips will continue on the same pattern until all of the samples have their supernatant removed.


7.18 Allowing beads to air dry

1m

The **Opentrons Magnetic Module** is engaged for  00:01:00 to allow the

 Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50** to air dry.

7.19 Removing excess wash buffer

Column 2 of tips in **Slot 8** will align with column 1 in **Slot 7** to remove excess wash by aspirating  10 µL and dispensing into the Liquid waste **NEST 1-Well Reservoir, 195 mL** in Slot 11. The tips will then be returned to their original starting point. Each subsequent column of tips will continue on the same pattern until all of the wash buffer is removed.

Note

It is important to remove any residual wash buffer before allowing beads to dry as it contains alcohol. Alcohol could prevent a good elution in the next step and inhibit further processes.

7.20 Allowing beads to air dry

2m

The **Opentrons Magnetic Module** remains engaged for  00:02:00 to allow the

 Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50** to have a final air dry and then disengages.

Note

Do not let beads dry for too long to prevent cracking of the pellet.





Expected result



The color of beads will change from shining dark brown to light brown when dried.

7.21 Adding elution buffer to Lysed Sample Plate 1 for Elution 1

The user must remove the  2.0 mL tubes with

 Elution Buffer **Omega Biotek Catalog #PDR048** from the  70 °C incubator and pour 4 of these tubes into well 12 in **Slot 1**.

Column 5 of tips in **Slot 5** will transfer  60 µL of

 Elution Buffer **Omega Biotek Catalog #PDR048** from well 12 in **Slot 1** to column 1 in **Slot 7** and mix by aspirating and dispensing  40 µL . The tips will then be dispensed into the waste container. Each subsequent column of tips will continue on the same pattern until all of the samples are mixed with warmed elution buffer.



Note

If you own a [Temperature module from Opentrons](#) you can also use it to keep the elution buffer continually warm.

7.22 Incubating the beads with DNA in elution buffer


15m


The protocol is paused for  00:15:00 to allow for the

 Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50** to incubate the DNA in the elution buffer at  Room temperature .

7.23 Allowing beads to settle on the magnet


2m

The [Opentrons Magnetic Module](#) is engaged for  00:02:00 to give the

 Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50** time to settle on the magnet away from the elution.

7.24 Transferring each sample elution to Sample Elution Plate 1





Column 6 of tips in **Slot 5** will transfer  60 μL of each eluate to a new, clean **Nest skirted PCR Plate** in **Slot 3**. Each subsequent column of tips will continue on the same pattern until all of the sample eluates are transferred, extending into the tips in **Slot 9**. The tips will be returned to the tip boxes to be reused for Elution 2.



The **Opentrons Magnetic Module** is disengaged.

7.25 Adding elution buffer to Lysed Sample Plate 1 for Elution 2

The user must remove the  2.0 mL tubes with

 Elution Buffer **Omega Biotek Catalog #PDR048** from the  70 °C incubator and pour the remaining 2 tubes into well 12 in **Slot 1**. They must also replace the **96-Well PCR Plate Non-skirt, 200 μL** now empty of sample lysis on top of an empty **Nest skirted PCR Plate** in **Slot 2** with a new, clean **Nest skirted PCR Plate** to receive Elution 2



Column 5 of tips in **Slot 9** will transfer  40 μL of

 Elution Buffer **Omega Biotek Catalog #PDR048** from well 12 in **Slot 1** to column 1 in **Slot 7** and mix by aspirating and dispensing  20 μL . The tips will then be dispensed into the waste container. Each subsequent column of tips will continue on the same pattern until all of the samples are mixed with warmed buffer, extending into the tips in **Slot 6**.

7.26 Incubating the beads with DNA in elution buffer


15m


The protocol is paused for  00:15:00 to allow for the

 Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50** to incubate the DNA in the buffer at  Room temperature .

7.27 Allowing beads to settle on the magnet


2m

The **Opentrons Magnetic Module** is engaged for  00:02:00 to give the

 Mag-Bind Particles CH **Omega Biotek Catalog #MBPCH- 50** time to settle on the magnet away from the elution.



7.28 Transferring each sample elution to Sample Elution Plate 2



The same tips that were used for Elution 1 will transfer  40 µL of each eluate to a new, clean **Nest skirted PCR Plate** in **Slot 2**. Each subsequent column of tips will continue on the same pattern until all of the sample eluates are transferred. The tips will be dispensed into the waste container.

The **Opentrons Magnetic Module** is disengaged.

7.29 Storage of Sample Elution Plates 1 and 2

Cover the plates with an aluminium plate seal and store at  4 °C for use or  -20 °C for long term storage.

7.30 QC of Sample Elution Plates 1 and 2


See QC Note in Step 1 for options.


After finishing the protocol

1h 9m

8 Clean the OT2 deck and walls with:

 10% Bleach 1 rinse


 Distilled Water 1 rinse

 70% Alcohol 2 rinses

Note

Avoid wetting any electronic parts.

9 Clean OT2 module with:

 70% Alcohol 2 rinses

Note

Avoid wetting electronic parts.

10 Air dry OT2 robot and modules.



Protocol references

This protocol is based on the [manufacturer's protocol](#) from Omega Biotek.