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# MagAttract + Metapolyzyme metagenomic gDNA extraction from urine V.2

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Dogstails



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Coming soon.

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Protocol status: Working We use this protocol and it's working

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## Abstract

A protocol for the metagenomic extraction of bacterial DNA from urine samples (optimised using dog urine), for use in a rapid diagnostics pipeline. At the end of the protocol, the DNA is cleaned up and ready for rapid barcoding (SQK-RBK004) library preparation for nanopore sequencing (or whatever other application you want to do).

Unless otherwise stated, all reagents should be included in the listed kits.

### Guidelines

This protocol, an adaptation of Qiagen's MagAttract HMW DNA kit, was developed by Natalie Ring and Alison Low for the Dogstails project, a collaboration between the Roslin Institute and the Royal (Dick) School of Veterinary Studies funded by the Dogs Trust. We are grateful to the dogs (and their owners) who donated samples to the R(D)SVS's Hospital for Small Animals, many of which were used in the development of this protocol.

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@NatalieAnneRing

## Materials

### Kits

- Urine sample from which to extract metagenomic gDNA
- X MagAttract HMW DNA kit Qiagen Catalog #67563
- X ProNex Size-Selective Purification System Promega Catalog #NG2001
- 🔀 Qubit® dsDNA HS Assay Kit Thermo Fisher Scientific Catalog #Q32854

#### **Other reagents**

- 50 mM Tris, 10 mM EDTA, ph8.0 ("buffer P1")
- MetaPolyzyme Sigma Aldrich Catalog #MAC4L-5MG
- X Nuclease-free Water
- X Distilled Water

### Equipment

Equipment	
DNA LoBind tubes, 1.5 mL	NAME
Tubes	TYPE
Eppendorf	BRAND
022431021	SKU
https://online-shop.eppendorf.us/US-en/Laboratory-Consumables Tubes-PF-56252.html	-44512/Tubes-44515/DNA-LoBind- K
1.5 mL	SPECIFICATIONS

OR

Equipment	
SafeSeal reaction tube, 1.5 ml, PP, PCR Performance Tested, Low DNA-binding	NAME
Tubes	TYPE
Sarstedt	BRAND
72.706.700	SKU
https://www.sarstedt.com/en/products/laboratory/screw-cap-micro-tubes-reaction-tubes/reaction-tubes/reaction-tubes/product/72.706.700/	LIN K
1.5 mL SPECIF	ICATIONS

Equipment		
Magnetic Stand	NAME	
Magnetic Stand	TYPE	
Thermo Scientific	BRAND	
MR02	SKU	
https://www.thermofisher.com/order/catalog/product/MR02 <sup>LINK</sup>		
Any magnetic rack that fits your tubes will suffice. SPECIFICATIONS		

Equipment	
Centrifuge	NAME
Benchtop Centrifuge	TYPE
Eppendorf	BRAND
5405000441	SKU
https://online-shop.eppendorf.us/US-en/Centrifugation-44533/Centrifuges-44534/Centrifuge PF-243560.html	-5425- <sup>LIN</sup> K
Any benchtop centrifuge will suffice	SPECIFICATIONS

Equipment	
ThermoMixer	NAME
Benchtop Incubator	TYPE
Eppendorf	BRAND
5382000023	SKU
https://online-shop.eppendorf.us/US-en/Temperature-Control-and-Mixing-44518/Instruments 44519/Eppendorf-ThermoMixerC-PF-19703.html	S- LIN K
Any heat block will suffice	SPECIFICATIONS

Equipment	
Mini-centrifuge	NAME
Centrifuge	TYPE
Fisher	BRAND
S67601B	SKU
https://www.fishersci.com/shop/products/fisherbrand-standard-mini-centrifuge-standard-micentrifuge/s67601b	ini- <sup>LIN</sup> K
Any standard mini centrifuge with adapters for different tube sizes will suffice	SPECIFICATIONS

Equipment		
Vortex Mixer	NAME	
Vortex Mixer	ТҮРЕ	
VWR	BRAND	
97043-562	SKU	
https://us.vwr.com/store/catalog/product.jsp?catalog_number=97043-562 <sup>LINK</sup>		

### **Protocol materials**

- 🔀 Nuclease-free Water
- X MagAttract HMW DNA kit Qiagen Catalog #67563
- 🔀 Qubit® dsDNA HS Assay Kit Thermo Fisher Scientific Catalog #Q32854
- X MetaPolyzyme Merck MilliporeSigma (Sigma-Aldrich) Catalog #MAC4L-5MG
- X ProNex Size-Selective Purification System Promega Catalog #NG2001
- 🔀 Distilled Water
- 🔀 Qubit® dsDNA HS Assay Kit Thermo Fisher Scientific Catalog #Q32854

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

### Before start

- "Buffer P1" is required for the metapolyzyme lysis incubation: 50 mM Tris, 10 mM EDTA, pH 8.0
- Metapolyzyme is used here at a concentration of 3.3 mg/ml (resuspend 5 mg lyophilized powder in 1.5 ml PBS pH 7.5)
- We recommend using low DNA-binding tubes throughout, but definitely for the elution/storage of DNA

Exte	ended pre-lysis spin down	
1	Pellet 2× 1.5 ml aliquots of urine in 1.5 ml tubes by centrifuging at maximum speed (~13,000 RPM/16,000 xg) for 20 minutes, then discard supernatant 3 mL urine	20m
	16,000 x g, Room temperature, 00:20:00	
	Note	
	We have found that this extended spin at the beginning of the protocol results in much better yield of bacterial gDNA, especially in samples with low bacterial abundance	
Met	apolyzyme & Proteinase K Lysis	
2	Resuspend cell pellets (which might be invisible) and combine in 160 $\mu$ l buffer P1 (50 mM Tris, 10 mM EDTA, pH 8.0)	
	📕 160 μL buffer P1	
3	Add 20 $\mu l$ metapolyzyme (3.3 mg/ml, 5 mg resuspended in 1500 $\mu l$ PBS) and mix by flicking the tube	
	Δ 20 µL metapolyzyme (3.3 mg/ml)	
4	Incubate on a thermomixer for 60 minutes at 37°C with 900 RPM shaking	1h
	<b>\$</b> 900 rpm, 37°C, 01:00:00	
5	Add 20 $\mu I$ MagAttract proteinase K and mix by flicking the tube	

👗 20 μL proteinase K

6 Incubate on a thermomixer for 30 minutes at 56°C with 900 RPM shaking

**()** 900 rpm, 56°C, 00:30:00

### MagAttract DNA isolation and washing

7 Add 150 µl MagAttract buffer AL and mix by pulse vortexing

👗 150 μL buffer AL

Note

Our standard "pulse vortex" is 10 short (<1 second) pulses per tube

8 Add 15 μl MagAttract Suspension G and 280 μl MagAttract buffer MB and mix by pulse vortexing

 $\stackrel{\text{L}}{=}$  15 µL Suspension G

🕹 280 μL Buffer MB

Note

Make sure the magnetic beads (Suspension G) are really well mixed before adding them! The whole suspension should be black, not separated into a bead layer and a clear layer. We usually resuspended by vortexing for 10 or more seconds.

9 Incubate on a thermomixer for 3 minutes at room temperature with 1,400 RPM shaking

30m

	<b>40</b> rpm, Room temperature , 00:03:00	
10	Spin down briefly, then pellet beads on magnet and remove supernatant	
11	Add 700 $\mu$ l MagAttract buffer MW1 and incubate on a thermomixer for 1 minute at room temperature with 1,400 RPM shaking	1m
	Δ 700 μL buffer MW1	
	<b>(5</b> 1400 rpm, Room temperature , 00:01:00	
12	Repeat steps 10 and 11	1m
13	Spin down briefly, then pellet beads on magnet and remove supernatant	
14	Add 700 μl MagAttract buffer PE and incubate on a thermomixer for 1 minute at room temperature with 1,400 RPM shaking	1m
	Ä 700 μL buffer PE	
	<b>(5</b> 1400 rpm, Room temperature , 00:01:00	
15	Repeat steps 13 and 14	1m
16	Spin down briefly, then pellet beads on magnet and remove supernatant	
17	Rinse the pelleted beads on the magnetic rack with 700 $\mu$ l distilled water by pipetting down the opposite wall of the tube, then incubate for 1 minute on the magnetic rack	
	a source opposite wait of the tabe, then incubate for riminute of the magnetic rack	
	L distilled water	

- 18 Remove distilled water
- 19 Repeat steps 17 and 18
- 20 Spin down briefly, then pellet beads on magnet and remove any remaining supernatant
- Add 50 μl nuclease-free water off the magnet, to resuspend the bead pellet
  - $\stackrel{\scriptstyle }{=}$  50 µL nuclease-free water
- 22 Incubate on a thermomixer for 3 minutes at room temperature with 1,400 RPM shaking
  - () 1400 rpm, Room temperature , 00:03:00
- 23 Spin down briefly, then pellet beads on magnetic rack and **keep supernatant** in a low-DNA binding 1.5 mL tube (e.g. <u>Eppendorf</u> or <u>Sarstedt</u>)

### **Qubit Pre-clean-up quantification**

- 24 Quantify DNA using Qubit dsDNA HS kit. If DNA concentration is an appropriate concentration for your experiment (for us, this means at least 0.2 ng/µl), continue to clean-up steps.
  - X Qubit<sup>®</sup> dsDNA HS Assay Kit VWR International Catalog #Q32854
  - 👗 1 μL DNA
  - $\stackrel{\scriptstyle }{=}$  199  $\mu$ L Qubit dsDNA HS working solution

### ProNex DNA clean-up

Add 150 μl room temperature ProNex beads to your entire tube of DNA (49 μl)

3m

	Δ 200 µL ProNex beads	
	Note	
	Like the magnetic beads in Suspension G, make sure the ProNex beads are really well mixed (10+ seconds of vortexing) immediately before you use them.	
26	Mix well by slowly pipetting up and down 10 times	
27	Incubate at room temperature for 10 minutes (no shaking needed)	10m
	00:10:00	
	Room temperature	
28	Spin down briefly, then pellet beads on magnet and remove supernatant	
29	Rinse the pelleted beads on the magnetic rack by pipetting 200 µl ProNex Wash Buffer down <b>the opposite wall of the tube</b> , then incubate at room temperature for 60 seconds (no shaking), then remove Wash Buffer	1m
	Δ 200 μL Wash Buffer	
	Com temperature	
	00:01:00	
30	Repeat step 26	
31	<b>Air-dry</b> (lid open) the sample on the magnetic rack for 5 minutes (longer is OK, no more than 60 minutes)	5m
	Room temperature	

00:05:00

32 Add 20 μl nuclease-free water off the magnet. Resuspend the pellet by **flicking the tube**, then incubate at room temperature for 5 minutes (no shaking needed)

Д	20 µL	nuclease-free	e water
e.	Room	temperature	

00:05:00

33 Spin down briefly, then pellet the beads on magnet and **keep supernatant** in a low DNAbinding tube

### Qubit post-clean-up quantification

34 Quantify DNA using Qubit dsDNA HS kit. If DNA concentration is an appropriate concentration for your experiment (for us, this means at least 0.2 ng/µl), continue to library preparation.

X Qubit<sup>®</sup> dsDNA HS Assay Kit VWR International Catalog #Q32854

👗 1 μL DNA

 $\Delta$  199 µL Qubit dsDNA HS working solution

5m