

# NucleoSpin<sup>®</sup> eDNA Water

# **User manual**

eDNA isolation



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## eDNA Isolation

## Protocol at a glance (Rev.01)

	NucleoSpin <sup>®</sup> eDNA Water					
1	Release eDNA from filter		5 mL tube	2 mL DISSOLVE 25 μL Liquid Proteinase K Mix Agitate (10 min, RT)		
2	Recover lysate		Ö	4,500 x <i>g</i> , 2 min		
3	Remove inhibitory substances		Ö	360 μL ACID 4,500 x <i>g</i> , 2 min		
4	Bind eDNA to the solid phase		Ö	50 μL PREC 25 μL NucleoTrap <sup>®</sup> suspension Mix Agitate (5 min, RT) 4,500 x <i>g</i> , 3 min		
		V	Ò	400 μL RESU, mix Transfer 10,000 x <i>g</i> , 1 min		
5	Wash and dry solid phase		Ö	400 μL WB, mix 10,000 x <i>g</i> , 1 min		
			Ö	300 μL WB, mix Transfer 10,000 x <i>g</i> , 2 min		
6	Elute eDNA		Ö	100 μL BE, mix 1min, RT 10,000 x <i>g</i> , 1 min		



## eDNA Isolation Simplified protocol at a glance (Rev.01)

NucleoSpin <sup>®</sup> eDNA Water					
1	Release eDNA from filter	-	► 5 mL tube	1.5 mL DISSOLVE 25 μL Liquid Proteinase K Mix Agitate (10 min, RT)	
2	Recover lysate			Use of syringe recommended for adsorbent filter (e.g. glass fiber) and not required for non-adsorbent filter (e.g. cellulose acetate).	
				10,000 x <i>g</i> , 1 min	
		<b>\$</b>	Ò	270 μL ACID	
3	Remove inhibitory				
	substances		Ò	10,000 x <i>g</i> , 2 min	
				Transfer	
	Bind eDNA to the solid phase			38 µL PREC	
				25 μL NucleoTrap <sup>®</sup> suspension	
4			Ò	Mix	
				Agitate (5 min, RT)	
				4,500 x <i>g</i> , 3 min	
	Wash and dry solid phase	÷			
				400 μL RESU, mix	
				Transfer	
			$\bigcirc$	10,000 x <i>g</i> , 1 min	
5				400 µL WB, mix	
			$\bigcirc$	10,000 x <i>g</i> , 1 min	
		S.		300 µL WB, mix	
		e	$\bigcirc$	Transfer	
		Real Provide American Science Provide American		10,000 x <i>g</i> , 2 min	
				100 ul BE mix	
6	Elute eDNA		Ċ	1min. BT	
				10.000 x <i>a</i> . 1 min	



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## 1 Components

## 1.1 Product content

	NucleoSpin <sup>®</sup> eDNA Water	
REF	740402.10	740402.50
Buffer DISSOLVE	30 mL	125 mL
Liquid Proteinase K	600 μL	1.5 mL
Buffer ACID	6 mL	30 mL
Precipitation Buffer PREC	1.5 mL	2 x 1.5 mL
NucleoTrap <sup>®</sup> Suspension	300 μL	1.5 mL
Resuspension Buffer RESU	5 mL	25 mL
Wash Buffer WB	10 mL	50 mL
NucleoSpin <sup>®</sup> eDNA XS column (light blue rings – plus Collection Tubes)	10	50
Elution Buffer BE	13 mL	13 mL
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# 1.2 Reagents, consumables, and equipment to be supplied by user

## Reagents and consumables:

- Filtration system Filter. Use of an ethylene oxide treated, round Glass Fiber Filter with 45 mm diameter (REF 740564; see section 6.2 Ordering information) or a similar round filter is recommended. Alternatively, membrane-type round filter (e.g. polyester, regenerated cellulose) or cartridge filters (e.g. Sterivex<sup>™</sup>) may be used.
- NucleoSpin Filters Midi (REF 740607; see section 6.2 Ordering information) are required for extraction protocol 5.1.1.
- Collection tubes 15 mL (15 mL High-Clarity Polypropylene Conical Tube, FALCON<sup>®</sup> REF 352096) are required for extraction protocol 5.1.1.
- 5 mL tubes (e.g. Eppendorf Biopur grade, recommended, or similar) are recommended for dissolution of eDNA from round filter according to protocol 5.1.2.
- Collection Tubes (1.5 mL) for elution
- Ethanol p.a. for filter drying in case of filter storage
- Disposable pipet tips (aerosol barrier pipet tips are recommended)

- DNA decontamination solution, e.g. bleach or Virkon Aquatic for decontamination of reusable materials and surfaces (e.g. water collection device, lab bench, pincers).
- Optional for turbid water samples: Filter flocs MN 2101 (REF 281120) with a support filter MN 750N (formats and REF upon request) to use as prefilter for turbid water (see section 6.2 Ordering information).

## Equipment:

- Water collection device (e.g. bottle, bucket, can, canister, beaker)
- Filter holder, e.g.
  - o Thermo Scientific<sup>™</sup> Nalgene<sup>™</sup> Analytical Test Filter Funnels 250 mL, in which the 0.45 µm filter membrane can be replaced by the recommended glass fiber filter or a filter of preference.
  - o Filter holder for use in the field: ADVANTEC Polypropylene Filter Holder for 47 mm Membranes (Cat. No. 43303020)
- Manual pipettors
- Vacuum device (e.g. vacuum pump) to move the water sample through the filter
- Tweezers for filter handling
- Personal protection equipment (lab coat, gloves, goggles)
- Vortex mixer
- Centrifuge for 15 mL tubes with a swing-out rotor capable of reaching 4,500 x g for use of extraction protocol 5.1.1
- Centrifuge for microcentrifuge tubes (1.5 mL or 2 mL)
- Tube roller for 15 mL tubes. Alternatively, 5 mL tubes and MN Bead Tube Holder 5 mL (see section 6.2 Ordering information)

## 1.3 About this user manual

It is strongly recommended to read the detailed protocol sections of this user manual before using the **NucleoSpin<sup>®</sup> eDNA Water** for the first time. Experienced users, however, may refer to the Protocol at a glance instead. The Protocol at a glance is designed to be used only as a supplemental tool for quick referencing while performing the purification procedure.

All technical literature is available on the internet at www.mn-net.com.

## 2 **Product description**

# 2.1 The basic principle: NucleoSpin<sup>®</sup> eDNA Water – filtration method

NucleoSpin<sup>®</sup> eDNA Water is a kit intended for the isolation of eDNA from milliliter to liter water samples collected from natural freshwater and marine environments. The focus of the kit is the isolation of DNA released from diverse organisms into the water. Organisms targeted are for example populations of fish, amphibians, reptiles, crustaceans, mammals, birds, plants and other organisms, that came in contact with the sampled water.

eDNA can be described as any DNA released from organisms into the environment. With regard to water samples, eDNA can be derived from e.g. epithelial cells, excretions, decaying organisms or pieces of tissue, reproductive cells (e.g. sperms, spawn, pollen). Most eDNA is associated with particles, such as cells, organelles, or bound to grains of inorganic or organic material. For this reason, eDNA can be obtained from water samples by filtration.

After filtration of an adequate water volume, DNA from the water is retained on the filter. The filter can either be directly processed as well as being stored for later processing. DNA is released from the filter and with a special dissolving buffer (Buffer DISSOLVE) and proteinase. Subsequently, DNA is absorbed to a particular matrix (NucleoTrap<sup>®</sup>), which is captured with a NucleoSpin<sup>®</sup> XS column, washed and finally eluted with a low salt elution buffer.

## 2.1.1 NucleoSpin® eDNA Water – precipitation method

Alternatively, eDNA can be isolated from a smaller water volume (e.g. up to 40 mL) without the need for filtration. In this procedure an active agent and a solid matrix are added to a 40 mL water sample. eDNA it is bound to the NucleoTrap<sup>®</sup>matrix, which is then captured with a with a NucleoSpin<sup>®</sup> XS column, washed and finally eluted with low salt buffer. For this procedure, the active agent Buffer (PREC) has to ordered separately (see section 6.2 Ordering information).

## 2.2 Product specification

- NucleoSpin<sup>®</sup> eDNA Water is designed for the isolation of eDNA from milliliter to liters of environmental water samples such as river, stream, creek, lake, pond, lagoon, bay, sea or ocean.
- NucleoSpin<sup>®</sup> eDNA Water recovers small to large eDNA fragments.
- A filtration protocol as well as a precipitation protocol is supported by the kit. For the precipitation method, extra buffer PREC has to be ordered.
- DNA yield strongly depends on the individual sample type and processed water volume. Total DNA yields of a few ng up to 3.5 µg have been observed from 1 L creek water
- Isolated DNA is suitable for diverse downstream applications such as PCR, qPCR and metabarcoding.

Table 1: Kit specifications at a glance				
Parameter	NucleoSpin <sup>®</sup> eDNA Water – filtration method	NucleoSpin <sup>®</sup> eDNA Water – precipitation method		
Technology	Silica beads/membrane technology			
Format	Filtration + beads + XS column	Precipitation + beads + XS column		
Sample material	environmental	water samples		
Sample amount	approximately 100 – 4000 mL*	up to 40 mL		
Fragment size	small to large			
Typical yield	Sample dependent (ng - µg)			
Ratio A <sub>260/280</sub> ; A <sub>260/230</sub>	typically not applicable due to low DNA content in the sample			
Elution volume	100 μL			
Filtration time	sample dependent	0 min		
Preparation time	approx. 60 min per 6 samples + filtration	approx. 30 min per 6 samples		
Binding capacity	not evaluated	not evaluated		

\*: The volume of filterable water depends very much on the filter used as well on the water quality (turbidity, suspended matter).

# 2.3 Handling of sample material - Stability of eDNA in the water sample

With collection / removal of a water sample from its natural environment, changes of eDNA amount and distribution in the sample will start. Sample storage can significantly influence the stability of the eDNA. For best results, the time from sample collection to onset of eDNA isolation should not exceed several hours, if the filter is not stabilized e.g. by drying or freezing. Ideally, water samples should be filtrated in the field at the collection site. Filters can be dried with ethanol and stored/transported for later eDNA isolation. If direct filtration is not an option, water samples can be cooled to 0 °C – 4 °C for several hours before eDNA extraction. For the precipitation method (section 5.2), Buffer PREC can be added to a 40 mL water sample at the collection site / in the field in order to stabilize the eDNA within the water sample.

## 2.4 Stability of DNA on the filter

eDNA can be stabilized on a filter by briefly passing ethanol (5 mL) over the filter in order to dry the filter. The ethanol-wet filter can be stored for several days in a closed tube at ambient temperature avoiding a spoilage of the sample/filter.

## 2.5 Elution procedures

A volume of 100  $\mu L$  is recommended for eDNA elution from the NucleoSpin<sup>®</sup> eDNA XS column. A smaller volume for elution is not recommended.

# 2.6 Size, yield and quality of eDNA obtained from water samples

DNA fragments from approximately 200 bp to >10,000 bp have been observed. Yield of total DNA can be in the range of few ng up to 3.5  $\mu$ g per 1 L water sample. Average fragment length and concentration of the isolated eDNA depend on the water sample being processed.

Some types of water sample tend to cause a carryover of brownish substances, such as humic acids. For such samples a clean up with the NucleoSpin<sup>®</sup> Inhibitor Removal kit (section 6.2 Ordering information) is recommended. Typically, eDNA eluates obtained are suitable for diverse downstream analysis.

## 2.7 Stability of isolated DNA

Due to the low DNA content of typical water samples, the resulting low total amount of isolated DNA, fragmentation, and the absence of DNase inhibitors (the elution buffer does not contain EDTA), the eluates should be kept on ice for short term storage and frozen at -20°C or below for long term storage for optimal results.

# 3 Storage conditions and preparation of working solutions

Attention: Resuspension Buffer RESU contains chaotropic salt! Wear gloves and goggles!

Caution: Resuspension Buffer RESU contains chaotropic salt which can form highly reactive compounds when combined with bleach (sodium hypochlorite). DO NOT add bleach or acidic solutions directly to the sample-preparation waste.

- Liquid Proteinase K can be stored at room temperature (18–25 °C) and is stable for at least one year. However, store Liquid Proteinase K at 4 °C or -20 °C after first use.
- All other kit components should be stored at room temperature (18–25 °C) and are stable for at least one year.
- Storage / transport at lower temperatures may cause precipitation of salt in buffer. If any
  precipitate is visible, heat the solution to 50°C for 30 min while mixing it, let the solution
  cool down to room temperature.

## 4 Safety instructions

The following components of the NucleoSpin® eDNA kits contain hazardous contents.

Wear gloves and goggles and follow the safety instructions given in this section.

Only harmful features do not need to be labeled with H and P phrases up to 125 mL or 125 g.

Mindergefahrliche Eigenschaften mussen bis 125 mL oder 125 g nicht mit H- und P-Satzen gekennzeichnet werden.

Component	Hazard contents	GHS symbol	Hazard phrases	Precaution phrases
Inhalt	Gefahrstoff	GHS-Symbol	H-Sätze	P-Sätze
Resuspension Buffer RESU	guanidinium thiocyanate 45–60 % <i>Guanidinthiocyanat 45–60</i> % CAS 593-84-0	WARNING ACHTUNG	302, 412	264W, 273, 301+312, 330
Wash Buffer WB	ethanol 55–75 % Ethanol 55–75 % CAS 64-17-5	DANGER GEFAHR	225	210, 233
Precipitation Buffer PREC	1-dodecylpyridiniumchloride 5–16.5 % 1-Dodecylpyridiniumchlorid 5–16.5 % CAS 104-74-5	DANGER GEFAHR	302, 314, 317, 411	260sh, 264W, 273, 280sh, 303+361+353, 305+351+338, 310, 330

The symbol shown on labels refers to further safety information in this section. Das auf Etiketten dargestellte Symbol weist auf weitere Sicherheitsinformationen dieses Kapitels hin.

## Hazard phrases

H 225	Highly flammable liquid and vapor. Flüssigkeit und Dampf leicht entzündbar.
H 302	Harmful if swallowed. Gesundheitsschädlich bei Verschlucken.
H314	Causes severe skin burns and eye damage. Verursacht schwere Verätzungen der Haut und schwere Augenschäden.
H 317	May cause an allergic skin reaction. Kann allergische Hautreaktionen verursachen.
H 411	Toxic to aquatic life with long lasting effects. Giftig für Wasserorganismen, mit langfristiger Wirkung.
H 412	Harmful to aquatic life with long lasting effects. Schädlich für Wasserorganismen, mit langfristiger Wirkung.

## **Precaution phrases**

P 210	Keep away from heat/sparks/open flames/hot surfaces. No smoking. Von Hitze, heißen Oberflächen, Funken, offenen Flammen sowie anderen Zündquellenarten fernhalten. Nicht rauchen.
P 233	Keep container tightly closed. Behälter dicht verschlossen halten.
P260sh	Do not breathe dust/vapors. Staub/Dampf nicht einatmen.
P 264W	Wash with water thoroughly after handling. Nach Gebrauch mit Wasser gründlich waschen.
P 273	Avoid release to the environment. Freisetzung in die Umwelt vermeiden.
P 280sh	Wear protective gloves/eye protection. Schutzhandschuhe/Augenschutz tragen.
P 301+312	IF SWALLOWED: Call a POISON CENTER/doctor if you feel unwell. BEI VERSCHLUCKEN: Bei Unwohlsein GIFTINFORMATIONSZENTRUM/Arzt anrufen.
P 303+361+353	IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water [or shower]. BEI BERÜHRUNG MIT DER HAUT (oder dem Haar): Alle kontaminierten Kleidungsstücke sofort ausziehen. Haut mit Wasser abwaschen [oder duschen].
P 305+351+338	IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing. BEI KONTAKT MIT DEN AUGEN: Einige Minuten lang behutsam mit Wasser ausspülen. Eventuell vorhandene Kontaktlinsen nach Möglichkeit entfernen. Weiter ausspülen.
P 310	Immediately call a POISON CENTER/doctor. Sofort GIFTINFORMATIONSZENTRUM/Arzt anrufen.
P330	Rinse mouth. Mund ausspülen.

For further information please see Material Safety Data Sheets (www.mn-net.com). Weiterfuhrende Informationen finden Sie in den Sicherheitsdatenblattern (www.mn-net.com).

## 5 Protocols

Before starting the procedure:

- Make sure to work in an environment which minimizes risk of DNA contamination: close doors and windows; prevent uncontrolled air draft; wear clean personal protection equipment; change gloves frequently. Ideally, work in a controlled positive pressure lab.
- Decontaminate your working environment (e.g. lab bench surfaces) before starting the procedure. Bleach or Virkon Aquatic can be used for decontamination (see: Quality assurance project plan – eDNA monitoring of bighead and silver carps; prepared for U.S. Fish and Wildlife Service; USFWS Midwest Region, Bloomington, MN, 2019)
- Make sure to decontaminate any reusable material (e.g. water collection container, filter holder, pincers/tweezers).
- When working with hypochlorite solutions, bleach, or Virkon Aquatic please take care to read the instructions in sections 3 and 4.
- Set up your filtration system before starting the procedure.
- Make sure to have all materials available, depending on your experimental set up (see section 1.2 for required and optional materials).
- Make sure to mix the NucleoTrap<sup>®</sup> suspension well before withdrawing an aliquot. It is
  important that the NucleoTrap<sup>®</sup> particles are released from the bottom of the tube and
  dispersed well within the liquid.

## 5.1 Protocol for eDNA isolation by filtration method

## Water collection

Collect water in a suitable container. For good eDNA coverage, collect water subsamples from different places of the water system to be analyzed and pool them. The optimal volume of water depends on the quality of the water and the abundance of the target DNA. To start with, a target volume of 1 L is recommended. The volume of filterable water depends very much on the filter used as well on the water quality (turbidity, suspended matter).

Note: Make sure to use a DNA-free water collection container and filtering equipment.

## Water filtration

As a filter we recommend to use the MN Glass Fiber Filter (45 mm, EO-treated, see section 6.2 Ordering information) to minimize the risk of DNA contamination. Other filter types (mixed cellulose ester, cellulose acetate, nylon, polyvinylidene fluoride, cellulose nitrate, polycarbonate) can be used but may show a lower filtration capacity or lower eDNA yield.

Filtration can be performed by either inserting the recommended Glass Fiber Filter (grid pattern facing down, side of water outlet) into an e.g. Nalgene<sup>™</sup> Analytical Test Filter Funnel, 250 mL (Thermo Scientific<sup>™</sup>) in which the 0.45 µm filter membrane is replaced by the recommended filter or a similar device (see Figure 1 below).

Alternatively, the round filter can be inserted into an ADVANTEC Polypropylene Filter Holder for 47 mm Membranes (Cat. No. 43303020).



## Figure 1 Usage of the recommended round Glass Fiber Filter (45 mm, EO-treated).

Alternatively, cartridge filters like Sterivex<sup>™</sup> Filter Units may be used, but might show a lower filtration capacity. Cartridge filters used should not require more than approximatel 2 mL solution for eDNA detachment from the filter (i.e. inner volumen of the cartridge should not exceed approximately 5 mL).

Let the water flow through the filter, e.g. by either negative pressure (vacuum) for the Nalgene<sup>™</sup> Analytical Test Filter Funnel or by negative or positive pressure for the ADVANTEC filter holder.

If the water sample clogs the filter, the uses of Filter flocs MN 2101 and a support filter MN 750N are recommended as a prefilter. In order to do so, insert a Nalgene<sup>™</sup> Analytical Test Filter Funnel containing the support Filter MN750N and approximately 1.6 g of filter flakes into a second Funnel which contains the GF-5 filter.

#### Storage/transport of filters (optional)

If storage / transport of the filter is required after filtration of water, add 5 mL of ethanol (*p.a.* quality) and let it pass through the filter. The ethanol displaces residual water from the filter and thus conserving the eDNA on the filter. Place the ethanol-wet filter into a suitable tube, e.g. a 5 mL reaction tube (e.g. Eppendorf Biopur grade).

After storage/transport of the filter, remove the filter from the tube and briefly let residual ethanol evaporate, before starting the eDNA release step.

## 5.1.1 Extraction protocol

Please note: this protocol requires the use of NucleoSpin<sup>®</sup> Filter Midi, 15 mL tubes and corresponding centrifuges. If you do not have such equipment available in your workspace, please use protocol 5.1.2. instead.

## 1 eDNA release from filter

Place the round filter as a whole into a suitable tube, e.g. a 5 mL reaction tube (e.g. Eppendorf Biopur grade, recommended, see figure above) or a 15 mL collection tube (not provided).

<u>Note:</u> Do not cut the filter. Make sure that top side (the former water facing side) of the filter is now facing inward, so that the subsequently added DISSSOLVE buffer can easily contact the filtration residue.

<u>Note:</u> If a cartridge filter is used, add DISSOLVE and Liquid Proteinase K into the cartridge.

Add 2 mL of buffer DISSOLVE into the tube.

Add 25 µL of Liquid Proteinase K into the buffer DISSOLVE and close the tube.

Vortex for 10 seconds. Make sure that the entire filter is wetted by the DISSOLVE buffer

Incubate the tube on a roller in a 5 mL tube (recommended) or 15 mL tube or on the MN Bead Tube Holder (for 5 mL tubes) for 10 min with moderate agitation. The filter should be surrounded by/submerged in the DISSOLVE buffer.

<u>Note:</u> Strong agitation/vortexing should be avoided. Some filter types (e.g. nitrocellulose, glass fiber) might disintegrate upon strong vortexing, which can subsequently interfere with the purification process.

## 2 Lysate recovery

Transfer the filter, the solution, and any sediment visible in the lysis tube into a NucleoSpin<sup>®</sup> Filter Midi resting in a 15 mL tube (not provided).

Centrifuge for 2 min at 4,500 xg. The lysate is recovered in the 15 ml tube.

Remove and discard the NucleoSpin® Filter Midi containing the round filter.

<u>Note:</u> The recommended filter type is a glass fiber filter absorbing most of buffer DISSOLVE. Other non-membrane filter likely will absorb most of the buffer, too. Therefore, the use of the NucleoSpin<sup>®</sup> Filter Midi is recommended in order to recover most of the solution. If other type of filters are used (e.g. membrane filter) the use of NucleoSpin<sup>®</sup> Filter Midi might be dispensable due to the low absorption of liquid by membrane filters. However, applying the NucleoSpin<sup>®</sup> Filter Midi is still recommended for optimum yield and ease of use.

<u>Note:</u> If a cartridge filter is used, remove the lysate from the cartridge and put the lysate into a 15 mL Collection Tube (not provided).

### 3 Removal of inhibitory substances

Add **360 µL buffer ACID** to the recovered lysate.

Invert the tube once to mix the buffer ACID into the solution.

<u>Note:</u> Do not mix thoroughly – pellets should not be resuspended. This step contributes to the removal of inhibitory substances, especially for humic acid rich (brownish) water samples.

Centrifuge the tube for 2 min at 4,500 x g.

Transfer the supernatant without any sediment into a fresh Collection Tube (15 mL, not provided).

#### 4 eDNA binding to the solid phase

Add **50 µL buffer PREC** to the supernatant.

Add 25 µL NucleoTrap® suspension and close the tube.

Note: Make sure to mix the suspension well before withdrawing an aliquot.

Incubate the tube at room temperature  $(18 - 25^{\circ}C)$  for 5 min on a roller.

<u>Note:</u> Alternatively, the tube can be incubated on shaker with moderate shaking. eDNA will bind to the NucleoTrap<sup>®</sup> suspension in this step.

Centrifuge the tube for 3 min at 4,500 x g.

Note: The NucleoTrap<sup>®</sup> with eDNA bound to it, is sedimented in this step.

Remove and discard the supernatant.

<u>Note:</u> Remove the supernatant with a pipet or turn the tube and place it upside down on a paper tissue to remove residual supernatant. Some droplets may stay in the tube – this does not compromise the purification process.

### 5 Wash and dry solid phase

Add **400 µL Resuspension Buffer RESU** to the NucleoTrap<sup>®</sup> pellet and resuspend it by vortexing.

Transfer the suspension into a 1.5 mL tube (not provided).

Centrifuge for 1 min at 10,000 xg.

Remove and discard the supernatant.

Note: The pellet might be translucent and thus be hardly visible in this step.

Add **400 \muL Wash Buffer WB** to the NucleoTrap<sup>®</sup> pellet and resuspend pellet by vortexing.

Centrifuge for 1 min at 10,000 xg.

Remove and discard the supernatant.

Note: The pellet is typically easily visible in this step.

Add **300 \muL Wash Buffer WB** to the NucleoTrap<sup>®</sup> pellet and resuspend pellet by vortexing

Transfer the NucleoTrap<sup>®</sup> suspension onto a **NucleoSpin<sup>®</sup> eDNA XS column**, placed in a Collection Tube (2 mL; provided).

Drying step: Centrifuge for 2 min at 10,000 xg.

Discard the flowthrough and place the column into an elution tube (1.5 mL; not provided).

#### 6 Elute eDNA

Add **100 µL Elution Buffer BE** onto the NucleoSpin® eDNA XS column.

Vortex for approximately 5–10 seconds until the NucleoTrap<sup>®</sup> material is completely resuspended.

Incubate approximately 1 min at room temperature.

Centrifuge for 1 min at 10,000 x g.

Discard the column and use the eDNA eluate for analysis.

## 5.1.2 Simplified extraction protocol

Please note: this protocol is intended for users who do not have the option of using 15 mL tubes and corresponding centrifuges in their workspace. If you have such equipment available in your workspace, we recommend to follow the protocol 5.1.1. instead. Also note that this protocol recommends the use of the MN Bead Tube Holder for 5 mL Tubes. Please check 6.2. for ordering information. Depending on the type of filter used (high or low liquid absorbend), a 5 mL syringe may be required for solution recovery from the filter (see ordering information).

## 1 eDNA release from filter

Place the round filter as a whole into a suitable tube, e.g. a 5 mL reaction tube (e.g. Eppendorf Biopur grade, recommended, see figure above).

<u>Note:</u> Do not cut the filter. Make sure that top side (the former water facing side) of the filter is now facing inward, so that the subsequently added DISSSOLVE buffer can easily contact the filtration residue.

Note: If a cartridge filter is used, add DISSOLVE and Proteinase into the cartridge.

Add 1.5 mL of buffer DISSOLVE into the tube.

Add 25 µL of Liquid Proteinase K into the buffer DISSOLVE and close the tube.

Vortex for 10 seconds. Make sure that the total filter is wetted by the DISSOLVE buffer.

Incubate the tube on the MN Bead Tube Holder (for 5 mL tubes) for 10 min with moderate agitation. The filter should be surrounded by/submerged in the DISSOLVE buffer.

<u>Note:</u> Strong agitation/vortexing should be avoided. Some filter types (e.g. nitrocellulose, glass fiber) might disintegrate upon strong vortexing, which can subsequently interfere with the purification process.

## 2 Lysate recovery

**For absorbent filters** (e.g. glass fiber filters), transfer the filter and the solution into a 5 mL syringe. Squeeze the liquid out of the filter by compressing it with the syringe and collect the liquid in a 2 mL tube (not provided). Discard the syringe containing the round filter.

<u>Note:</u> If filter material has disinterated due to excessive agitation/vortexing, the use of Syringe filter, PES, 25 mm, 5  $\mu$ m (see ordering information) is recommended: pass the solution from the syringe through the syringe filter.

**For non-absorbent filters** (e.g. membrane filters) it is sufficient to remove the filter from the DISSOLVE buffer, let it drip-off briefly and discard the filter. Transfer the lysate into a fresh 2 mL tube (not provided). Alternatively, the filter can be pushed aside within the tube with a pipet tip in a way that the solution can be recovered directly from the tube without prior filter removal.

<u>Note:</u> The recommended filter type is a glass fiber filter absorbing most of buffer DISSOLVE. Other non-membrane filters are highly likely to absorb most of the buffer as well. In such cases, the use a 5 mL syringe is recommended in order to recover most of the solution. If other types of filters are used (e.g. membrane filter) the use of a syringe is dispensable due to the low absorption of liquid by membrane filters.

<u>Note:</u> If a cartridge filter is used, remove the lysate from the cartridge and put the lysate into a 2 mL Collection Tube (provided).

#### Centrifuge the recovered lysate for 1 min at 10,000 x g.

Note: A loose sediment might form, which can stay in the tube.

#### 3 Removal of inhibitory substances

Add 270 µL buffer ACID to the tube.

Invert the tube once to mix the buffer ACID into the solution.

<u>Note:</u> Do not mix thoroughly – pellets should not be resuspended.

Centrifuge for 2 min at 10,000 x g and transfer the supernatant without any sediment into a fresh Collection Tube (2 mL, not provided).

Note: Sediment might be very loose.

### 4 eDNA binding to the solid phase

Add 38 µL buffer PREC to the recovered supernatant.

Add 25 µL NucleoTrap<sup>®</sup> suspension and close the tube.

Note: Make sure to mix the suspension well before withdrawing an aliquot.

Incubate the tube at room temperature (18 - 25°C) for 5 min on a roller.

<u>Note:</u> Alternatively, the tube can be incubated on shaker with moderate shaking. eDNA will bind to the NucleoTrap<sup>®</sup> suspension in this step.

Centrifuge the tube for 3 min at 4,500 xg.

Note: The NucleoTrap<sup>®</sup> with eDNA bound to it is sedimented in this step.

Remove and discard the supernatant.

<u>Note:</u> Remove the supernatand with a pipet or turn the tube and place it upside down on a tissue to remove residual supernatant. Some droplets may stay in the tube.

#### 5 Wash and dry solid phase

Add **400 µL Resuspension Buffer RESU** to the NucleoTrap<sup>®</sup> pellet and resuspend it by vortexing.

Centrifuge for 1 min at 10,000 x g.

Remove and discard the supernatant.

Note: The pellet might be translucent and thus be hardly visible in this step.

Add  $400~\mu L$  Wash Buffer WB to the NucleoTrap  $^{\odot}$  pellet and resuspend pellet by vortexing.

Centrifuge for 1 min at 10,000 x g.

Remove and discard the supernatant.

Note: The pellet is typically easily visible in this step

Add **300 \mu L Wash Buffer WB** to the NucleoTrap® pellet and resuspend pellet by vortexing

Transfer the NucleoTrap<sup>®</sup> suspension onto a **NucleoSpin<sup>®</sup> eDNA XS column**, placed in a Collection Tube (2 mL; provided).

Drying step: Centrifuge for 2 min at 10,000 xg.

Discard the flowthrough and place the column into an elution tube (1.5 mL; not provided).

## 6 Elute eDNA

Add 100 µL Elution Buffer BE onto the NucleoSpin® eDNA XS column.

Vortex for approximately 5–10 seconds until the NucleoTrap  $^{\ensuremath{\mathbb{R}}}$  material is completely resuspended.

Incubate approximately 1 min at room temperature.

Centrifuge for 1 min at 10,000 x g.

Discard the column and use the eDNA eluate for analysis.

# 5.2 Protocol for eDNA isolation from 40 mL water by direct precipitation (without filtration)

Before starting the procedure:

- Additional precipitation buffer PREC is required for this procedure (see ordering information).
- Make sure to work in an environment which minimizes risk of DNA contamination.
- Decontaminate your working environment (e.g. lab bench surfaces) before starting the procedure.
- Make sure to decontaminate any reusable material (e.g. pincers/tweezers).
- Make sure to have all materials available, depending on your experimental set up (see section 1.2 for required and optional materials)
- Make sure to mix the NucleoTrap<sup>®</sup> suspension well before withdrawing an aliquot. It is
  important that the NucleoTrap<sup>®</sup> particles are released from the bottom of the tube and
  dispersed well within the liquid.

## 1 Water collection

Collect **40 mL water** in a 50 mL tube (not provided). For good eDNA coverage, collect water subsamples from different places of the water system to be analyzed and pool them. The optimal volume of water depends on the quality of the water and the abundance of the target DNA.

Note: Make sure to use a DNA-free water collection container.

Add **800 µL PREC** to the 40 mL water sample and mix.

<u>Note:</u> The sample can be stored/transported for several days at room temperature  $(18-25^{\circ}C)$  preventing sample spoilage.

<u>Note:</u> If water volumes smaller than 40 mL are processed, adjust the volume of PREC proportionally.

## 2 eDNA binding to the solid phase

Add 25 µL NucleoTrap<sup>®</sup> suspension and close the tube.

Note: Make sure to mix the suspension well before withdrawing an aliquot.

Incubate the tube at room temperature (18-25°C) for 15-20 min on a roller.

<u>Note:</u> eDNA will bind to the NucleoTrap<sup>®</sup> suspension in this step.

Centrifuge the tube for 3 min at 4,500 x g.

<u>Note:</u> The NucleoTrap<sup>®</sup> is sedimented in this step.

Remove and discard the supernatant.

Note: Turn tube upside down to remove residual free supernatant.

## 3 Wash and dry solid phase

Continue the procedure as described in section 5.1 with Wash and dry solid phase.

## 6 Appendix

## 6.1 Troubleshooting

Problem	Possible cause and suggestions		
	<ul> <li>Too much water used. Use less water or use MN Filter Flocs MN 2101 and Filter MN750N in a prefilter.</li> </ul>		
	<ul> <li>Water contains too much insoluble substances. Use MN Filter Flocs MN 2101 and Filter MN750N in a prefilter in order to prevent/reduce risk of filter clogging or use less water. See figure below.</li> </ul>		
	Filter funnel 1		
Filter clogging	Filter funnel 2		
Filler clogging	1.6 g Filter flocs MN 2101		
	Support Filter MN 750 N		
	Glass Fiber Filter GF5		
	Figure 2 Example of the assembly of a prefilter with MN2101 filter flocs and support filter MN 750N.		
	<ul> <li>Membrane filter used. Membrane filter tend to clog much faster than the recommended one. Use the recommended filter.</li> </ul>		
	Filter bypass. Make sure that the filter is placed correctly in the filter holder ensuring that all water passes through the filter and that there is no filter bypass. Some filter types (e.g. membrane filter) tend to have higher risks of bypasses or filter breakages.		
NO DNA yiela	Inappropriate DNA quantification method. Some water types contain little eDNA. Make sure your DNA quantification is sensitive enough (e.g. quantification with PicoGreen dye or by PCR).		
	<ul> <li>Water quality: Some water samples contain predominantly highly degraded DNA while other water samples may contain larger amounts of high molecular weight DNA.</li> </ul>		
degraded	<ul> <li>Inappropriate sample collection, storage, transport: Make sure to process the sample as soon as possible after collection. If there is a delay, it is possible to cool the water for several hours. Alternatively, filters may be conserved by ethanol treatment.</li> </ul>		

Problem	Possible cause and suggestions		
	<ul> <li>DNA contamination during sample collection or processing: Make sure to work in an environment which minimizes risk of DNA contamination: close doors and windows; prevent uncontrolled air draft; wear clean personal protection equipment; change gloves frequently; ideally work in a controlled positive pressure lab.</li> </ul>		
Contamination of the eluate with DNA from	<ul> <li>Contaminated reusable materials (e.g. water collection container): Make sure to only use reusable materials which has been decontaminated.</li> </ul>		
unexpected species	<ul> <li>Contaminated working environment: Decontaminate your working environment (e.g. lab bench surfaces) before starting the procedure.</li> </ul>		
	<ul> <li>Run a no-sample extraction control in order to obtain information, which DNA does not originate from your sample.</li> </ul>		
	• Wrong or unspecific species "identification": Make sure that your detection method is sufficient specific for organisms or your interest.		

The NucleoSpin<sup>®</sup> eDNA Water kit is produced under controlled and monitored conditions with high hygienic standards. Nonetheless, it cannot be guaranteed that there is no contaminating DNA at all times of whatever source due to the omnipresence of DNA and the increasing sensitivity of DNA detection and assignment. Therefore, a no-sample extraction control (mock preparation) is recommended.

## 6.2 Ordering information

Product	REF	Pack of
Glass Fiber Filter (45 mm, EO-treated)	740564	50 pieces
NucleoSpin <sup>®</sup> Filter Midi	740607	50 pieces
MN Bead Tube Holder 5 mL	740459	1 piece
Filter flocs MN 2101, ashless, quantitative	281120	500 g
Filter MN 750 N (45 mm diameter)	please inquire for formats a	Ind REF
Disposable 5 mL syringe wiht Luer tip	729101	100 pieces
Syringe filter, PES, 25 mm, 5 $\mu$ m	729242	100 pieces
Buffer PREC	740568	50 mL
NucleoSpin <sup>®</sup> Inhibitor Removal	740408.10/.50	10/50 preps

## 6.3 Product use restriction / warranty

NucleoSpin® eDNA Water products are intended, developed, designed, and sold FOR RESEARCH PURPOSES ONLY, except, however, any other function of the product being expressly described in original MACHEREY-NAGEL product leaflets. MACHEREY-NAGEL products are intended for GENERAL LABORATORY USE ONLY! MACHEREY-NAGEL products are suited for QUALIFIED PERSONNEL ONLY! MACHEREY-NAGEL products shall in any event only be used wearing adequate PROTECTIVE CLOTHING. For detailed information please refer to the respective Material Safety Data Sheet of the product! MACHEREY-NAGEL products shall exclusively be used in an ADEQUATE TEST ENVIRONMENT. MACHEREY-NAGEL does not assume any responsibility for damages due to improper application of our products in other fields of application. Application on the human body is STRICTLY FORBIDDEN. The respective user is liable for any and all damages resulting from such application. 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It is rather in the responsibility of the user or - in any case of resale of the products - in the responsibility of the reseller to inspect and assure the use of the DNA/RNA/protein purification products of MACHEREY-NAGEL for a well-defined and specific application. MACHEREY-NAGEL shall only be responsible for the product specifications and the performance range of MN products according to the specifications of in-house quality control, product documentation and marketing material. This MACHEREY-NAGEL product is shipped with documentation stating specifications and other technical information. MACHEREY-NAGEL warrants to meet the stated specifications. MACHEREY-NAGEL's sole obligation and the customer's sole remedy is limited to replacement of products free of charge in the event products fail to perform as warranted. Supplementary reference is made to the general business terms and conditions of MACHEREY-NAGEL, which are printed on the price list. 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It cannot be guaranteed, that the NucleoSpin<sup>®</sup> eDNA Water kits are free of any detectable DNA at all times. However, the products are produced under controlled and monitored conditions with high hygienic standards to keep the risk of DNA contamination of whatever source as low as reasonably possible.



Plasmid DNA Clean up RNA DNA Viral RNA and DNA Protein High throughput Accessories Auxiliary tools



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