

# **User manual** NucleoMag 96 Tissue

June 2006/Rev. 01





# Table of contents

1	Kit c	ontents	4	
2	Proc	luct description	5	
	2.1	The basic principle	5	
	2.2	Kit specifications	5	
	2.3	Magnetic separation systems	6	
	2.4	Adjusting the shaker settings	6	
	2.5	Handling of beads	7	
	2.6	Elution procedures	8	
3 Storage conditions and preparation of working solutions				
4	4 Safety instructions – risk and safety phrases			
5	Gen	eral procedure	11	
	5.1	Protocol for the purification of genomic DNA from tissue samples,		
		cultured cells, or bacteria	13	
6	Арр	endix	16	
	6.1	Troubleshooting	16	
	6.2	Ordering information	18	
	6.3	Product use restriction / warranty	18	

# 1 Kit contents

	NucleoMag 96 Tissue		
Cat. No.	1 x 96 preps 744 300.1	4 x 96 preps 744 300.4	24 x 96 preps 744 300.24
NucleoMag-B-Beads	2.8 ml	11.5 ml	70 ml
Lysis Buffer T1	25 ml	100 ml	2 x 300 ml
Binding Buffer MB2	45 ml	2 x 90 ml	9 x 120 ml
Wash Buffer MB3	75 ml	300 ml	2 x 900 ml
Wash Buffer MB4	75 ml	300 ml	2 x 900 ml
Wash Buffer MB5	125 ml	500 ml	3 x 1000 ml
Elution Buffer MB6	25 ml	100 ml	2 x 300 ml
Proteinase K (lyophilized)	75 mg	4 x 75 mg	24 x 75 mg
Proteinase Buffer PB	3,6 ml	15 ml	3 x 35 ml
Protocol	1	1	1

# Material to be supplied by user:

Product	Cat. No.	Pack of
Separation Plate for magnetic beads separation, e.g. Square-well Block	740670	20
Lysis Tubes for incubation of samples and lysis, e.g. MN Tubes Strips	740637	5 sets
Cap Strips for Lysis Tubes	740638	30
Elution Plate for collecting purified DNA, e.g. 0,3 ml Microtiterplate, u-bottom e.g. 0,3 ml Microtiterplate, flat-bottom	740672 740673	20 20

# 2 **Product description**

# 2.1 The basic principle

The **NucleoMag 96 Tissue** procedure is based on reversible adsorption of nucleic acids to paramagnetic beads under appropriate buffer conditions. Tissue samples, cells or bacteria are lysed with SDS / proteinase K solution (buffer T1). For the adjustment of the binding conditions under which nucleic acids bind to the paramagnetic beads buffer MB2 and the NucleoMag-B-Beads Binding are added to the lysate. After magnetic separation the paramagnetic beads are washed twice to remove contaminants and salts using wash buffers MB3 and MB4. There is no need for a drying step as ethanol from previous wash steps is removed by a final incubation of the beads in buffer MB5. Finally, highly purified DNA is eluted with low-salt elution buffer (MB6) and can directly be used for downstream applications. The **NucleoMag 96 Tissue** kit can be used either manually or automated on standard liquid handling instruments or automated magnetic separators.

# 2.2 Kit specifications

**NucleoMag 96 Tissue** is designed for rapid manual and automated small-scale preparation of highly pure genomic DNA from tissue samples, cells or bacteria using the NucleoMag 96 SEP (see ordering information) or other magnetic separation systems (see section 2.3). Manual time for the preparation of 96 samples is about 120 minutes. The purified DNA can be used directly as template for PCR, blotting, or any kind of enzymatic reactions.

**NucleoMag 96 Tissue** allows easy automation on common liquid handling instruments. The actual processing time depends on the configuration of the instrument and the magnetic separation system used. Typically, 96 samples can be purified in less than 120 minutes using the NucleoMag SEP on the automation platform.

The kit provides reagents for the purification of up to 20  $\mu$ g of pure genomic DNA from suitable samples (up to 20 mg tissue, up to 1x 10<sup>7</sup> cells or up to 1 ml of an overnight culture of bacteria) with an A<sub>260/280</sub> ratio  $\geq$  1.6–1.9 and typical concentration of 20–50 ng/µl. Depending on the elution volume used concentrations of 10–150 ng/µl can be obtained.

Following lysis of samples with proteinase K **NucleoMag 96 Tissue** can be processed completely at room temperature, however, elution at 55°C will increase the yield by about 15–20%.

NucleoMag B-Beads are highly reactive, superparamagnetic beads. The binding capacity is 0.4  $\mu$ g of gDNA per 1  $\mu$ l of NucleoMag-B-Bead Suspension, 1  $\mu$ l of suspension contains 130  $\mu$ g of beads.

# 2.3 Magnetic separation systems

For use of **NucleoMag 96 Tissue** the use of the magnetic separator NucleoMag SEP is recommended. Separation is carried out in a 2 ml Square-well Block (see ordering information). If the kit is used with other common separators, see suppliers ordering information for suitable separation plates.

Magnetic separator	Separation plate or tube		
NucleoMag SEP (Cat. No. 744 900)	Square-well Block (MN, Cat. No. 740 670)		
Tecan Te-MagS	1.5 ml tubes without lid (Sarstedt)		

## Static magnetic pins

Separators with static magnetic pins, e.g. NucleoMag SEP (for manual use and for use on liquid handling workstations): This type of separator is recommended in combination with a plate shaker, e.g. H+P Variomag<sup>®</sup> Teleshake (H+P Labortechnik AG, Bruckmannring 28, D-85764 Oberschleißheim, Germany, www.hp-lab.de), for optimal resuspension of the beads during the washing and elution steps. Alternatively, beads can be resuspended in the buffer by pipetting up and down several times. For fully-automated use on liquid handling workstations a gripper tool is required, that transfers the plate to the magnetic separator for separation of the beads or to the shaker module for resuspension of the beads.

## Movable magnetic systems

Separators with moving magnets, e.g. Te-MagS (for automated use only): Magnetic pins/rods are moved from one side of the well to the other and vice versa. Beads follow this movement and are thus pulled through the buffer during the wash and elution steps. Separation takes place when the system stops.

## Automated separators

Separators with moving magnets, e.g. ThermoElectron's King Fisher instruments: Magnetic beads are transferred into suitable plates or tubes. Beads are resuspended from the rod-covered magnets. Folowing binding, washing or elution beads are collected again with the rod-covered magnets and transferred to the next plate or tube.

# 2.4 Adjusting the shaker settings

When using a plate shaker for the washing and elution steps the speed settings have to be checked carefully for each specific separation plate and shaker to prevent cross-contamination from well to well. Proceed as follows:

## Checking shaker speed for wash steps:

- load 600 µl dyed water to the wells of the separation plate. Place the plate on the shaker and start shaking with a moderate speed setting for 30 seconds. Turn off the shaker and check plate surface for small droplets of dyed water.
- Increase speed setting, shake for an additional 30 seconds, and check plate surface for droplets again.

• Continue increasing the speed setting until you observe droplets on top of the separation plate. Reduce speed setting, check again and use this setting for the washing step.

## Checking shaker speed for elution step:

 apply 100-200 µl dyed water to the wells of the collection plate and proceed as described above.

# 2.5 Handling of beads

## Distribution of beads

A homogenous distribution of the magnetic beads to the individual wells of the separation plate is essential for a high well-to-well consistency. Therefore, before distributing the beads make sure that the beads are completely resuspended. Shake the storage bottle well or place it on a vortexer shortly. Premixing magnetic beads with the binding buffer allows easier homogenous distribution of the beads to the individual wells of the separation plate. During automation, a premix step before aspirating the beads/binding buffer mixture from the reservoir is recommended to keep the beads resuspended.

## Magnetic separation time

Attraction of the magnetic beads to the magnetic pins depends on the magnetic strength of the magnetic pins, the selected separation plate, distance of the separation plate from the magnetic pins, and the volume to be processed. The individual times for complete attraction of the beads to the magnetic pins should be checked and adjusted on each system. It is recommended to use the separation plates or tubes specified by the supplier of the magnetic separator.

## Washing the beads

Washing the beads can be achieved by shaking or mixing. In contrast to mixing by pipetting up and down mixing by shaker or magnetic mixing allows simultaneous mixing of all samples. This reduces the time and number of tips needed for the preparation. Resuspension by pipetting up and down, however, is in general more efficient than mixing by a shaker or magnetic mix.

Method	Resuspension Efficiency	Speed	Small elution volume possible	Number of tips needed
Magnetic mixing	+	++	+	Low
Shaker	++	++	+++	Low
Pipetting	+++	+*	++	High

\* 8-channel pipetting device

# 2.6 Elution procedures

Purified genomic DNA can be eluted directly with the supplied Elution Buffer MB6. Elution can be carried out in a volume of  $\geq$  50 µl. It is essential to cover the NucleoMag Beads completely with elution buffer during the elution step. The volume of dispensed elution buffer depends on the magnetic separation system (e.g. the position of the pellet inside the separation plate). For efficient elution the magnetic bead pellet should be resuspended completely in the elution buffer. For some separators high elution volumes might be necessary to cover the whole pellet.

Elution is possible at room temperature. However, the DNA yield can be increased by 15 - 20% if the elution step is performed at  $55^{\circ}$ C.

# 3 Storage conditions and preparation of working solutions

Attention:

Buffers MB2, MB3 and MB4 contain chaotropic salt! Wear gloves and goggles!

All components of the **NucleoMag 96 Tissue** kit should be stored at room temperature (20-25°C) and are stable for up to one year.

All buffers are delivered ready-to-use.

Before starting **NucleoMag 96 Tissue** protocol prepare the following:

 Before first use of the kit, add the indicated volume of Proteinase Buffer to lyophilized proteinase K. Proteinase K solution is stable at +4°C for up to 6 months. Storage at -20°C is recommended if the solution will not be used up during this period.

	NucleoMag 96 Tissue			
	1 x 96 preps	4 x 96 preps	24 x 96 preps	
Cat. No.	744 300.1	744 300.4	744 300.24	
	1 x 75 mg	4 x 75 mg	24 x 75 mg	
Proteinase K (lyophilized)	add 2.6 ml Proteinase Buffer	add 2.6 ml Proteinase Buffer to each vial	add 2.6 ml Proteinase Buffer to each vial	

#### Safety instructions – risk and safety phrases 4

Component	Hazard Contents	Hazard Symbo			Risk Phrases	Safety Phrases
MB2, MB3, MB4	Sodium perchlorate + ethanol < 50%	× >	Xn*	Harmful if swallowed. Flammable	R 10-22	S 7-13-16- 27
Proteinase K	Proteinase K, lyophilized		Xn, Xi <sup>**</sup>	Irritating to eyes, respiratory system and skin, may cause sensitization by inhalation	R 36/37/38- 42	S 22-24-26- 36/37

## **Risk Phrases**

R10	Flammable
R 22	Harmful if swallowed
R 36/37/38	Irritating to eyes, respiratory system and skin
R 42	May cause sensitization by inhalation

## **Safety Phrases**

S 13	Keep away from food, drink and animal feedstuffs
S 22	Do not breathe dust
S 24	Avoid contact with the skin
S 26	In case of contact with eyes, rinse immediately with plenty of water and seek medical advice
S 27	Take off immediately all contaminated clothing
S 36/37	Wear suitable protective clothing and gloves

<sup>\*</sup> Label not necessary, if quantity below 50 g or ml (according to 67/548/EEC Art. 25, 1999/45/EC Art. 12 and German GefStoffV § 42 and TRGS 200 7.1) \*\*\* Label not necessary, if quantity below 125 g or ml (according to 67/548/EEC Art. 25, 1999/45/EC Art. 12 and German

GefStoffV § 42 and TRGS 200 7.1)

# 5 General procedure

1	Lyse samples (up to 20 mg tissue, up to 1x10 <sup>7</sup> cells or bacteria pellet from up to 1 ml overnight culture).	Add 25 µl proteinase K solution and 200 µl T1 buffer, mix and incubate at 56°C, 1-3 h or overnight	
2	Clear lysate by centrifugation, transfer 225 µl of cleared lysate to a suitable separation plate (e.g. Square-well Block) for further processing.	5,600 x g	
3	Bind DNA to NucleoMag Beads.	225 µl cleared lysate 24 µl B-beads 360 µl MB2 shake 5 min at RT optional: mix by pipetting up and down	
4	Remove supernatant.	after 2 min separation remove supernatant	
5	MB3 wash step.	600 μl MB3 shake 5 min, RT optional: mix by pipetting up and down after 2 min separation remove supernatant	

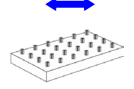
6 MB4 wash step.

600 µl MB4 shake 5 min, RT

optional: mix by pipetting up and down

after 2 min separation remove supernatant





8 MB5 wash step.

900 µl MB5 60 sec incubation aspirate and discard supernatant

Note: Do not resuspend beads in MB5 buffer

9 Elute genomic DNA and transfer to Elution Plate.

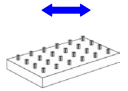
50 – 200 µl MB6 shake 5-10 min, RT

optional: mix by pipetting up and down

after 2 min separation transfer eluted DNA

> Optional: Elution at 55°C





# 5.1 Protocol for the purification of genomic DNA from tissue samples, cultured cells, or bacteria

This protocol is designed for magnetic separators with static pins (e.g. NucleoMag SEP) and suitable plate shakers (e.g. H+P Variomag<sup>®</sup> Teleshake). This protocol is for manual use and serves as a guideline for adapting the kit to robotic instruments. Contact MACHEREY-NAGEL for available scripts or support protocols.

## 1 Lyse samples

Calculate the amount of lysis stock required: for each sample 25  $\mu$ l of proteinase K solution + 200  $\mu$ l buffer T1 are required. Prepare lysis stock solution accordingly and vortex.

Never prepare the lysis stock solution more than 15 min before addition to the samples. Proteinase K tends to self digestion when incubated in T1 without substrate.

Transfer 225 µl of the resulting stock solution to each lysis tube containing up to 20 mg of tissue sample (e.g. mouse tail section), or up to  $1 \times 10^7$  cultured cells or up to 1 ml of an overnight culture of bacteria. Close the individual tubes. Mix by vigorous shaking for 10 – 15 s. Spin briefly (15 s; 1,500 × g) to collect any sample at the bottom of the tube.

The sample must be submerged in the solution.

Incubate the tubes containing the samples at 56°C until complete lysis is obtained (at least 1-3 h or overnight). For cultured cells incubation can be carried out at 70°C for 10-15 min. For optimal lysis, mix occasionally during incubation. Make sure that the lysis tubes are securely closed.

If RNA-free DNA is crucial for downstream applications, an RNase digest may be performed: Add 20 µl RNase A (20 mg/ml) solution (not included, see ordering information)and incubate for additional 5 min at room temperature.

2 Remove residual debris, e.g. hair and/or bones or cell debris, in the lysate by centrifugation (5 min;  $5,600 - 6,000 \times g$ ) and transfer 225 µl of the lysate into the wells of a suitable separation plate (e.g. Square-well Block).

**3** Add 24 μl of NucleoMag-B-Beads and 360 μl buffer MB2 to 225 μl of cleared lysate obtained from previous step. Mix immediately by shaking (5 min) at room temperature.

Alternatively, pipette up and down 10 times and incubate 5 min at room temperature

Note: NucleoMag-B-Beads and Binding Buffer MB2 can be premixed. Per well to be processed mix 24  $\mu$ l of NucleoMag-B-Beads with 360  $\mu$ l Binding Buffer MB2. Vortex briefly. Depending on the dead volume of the reservoir, additional amounts of bead suspension and binding buffer are necessary.

Be sure to resuspend the NucleoMag-B-Beads before removing them from the storage bottle. Vortex storage bottle briefly until a homogenous suspension has formed.

4 Separate the magnetic beads against the side of the wells by placing the separation plate on the magnetic separator. Wait at least 2 min until all the beads have been attracted to the magnet. Remove and discard supernatant by pipetting.

Note: Do not disturb the attracted beads while aspirating the supernatant. The magnetic pellet might not be visible in this step. Remove supernatant from the opposite side of the well.

Remove the separation plate from the magnetic separator.

**5** Add 600 μl Wash Buffer MB3 to each well and wash the bead/DNA complex by shaking (5 min) at room temperature.

Alternatively, pipette up and down 15 times.

Separate all of the magnetic beads against the side of the well by placing the separation plate on the magnetic separator. Aspirate and discard the supernatant.

Note: Supernatant may have a pale brownish color, magnetic bead pellet is visible now.

Remove the separation plate from the magnetic separator.

**6** Add 600 μl Wash Buffer MB4 to each well and wash the bead/DNA complex by shaking (5 min) at room temperature.

Alternatively, pipette up and down 15 times.

Separate all of the magnetic beads against the side of the well by placing the separation plate on the magnetic separator. Aspirate and discard the supernatant.

Note: Supernatant is colorless, magnetic bead pellet is clearly visible

Leave the separation plate on the magnetic separator for step 7!

**7** Add 900 μl Wash Buffer MB5 to each well and incubate for 60 s while the beads are still attracted on the magnet. Then aspirate and discard the supernatant.

Note: Do not resuspend the beads in Wash Buffer MB 5. This step is required to remove traces of ethanol and eliminates a drying step. Resuspension of beads in buffer MB5 will decrease DNA yield.

8 Add desired volume of Elution Buffer MB6 ( $50 - 200 \mu$ l) to each well, remove the separation plate from the magnetic separator, and resuspend the bead/DNA complex by shaking ( $5 - 10 \mu$ ).

Alternatively, pipette up and down 10 times and incubate 5 – 10 min.

Separate the magnetic beads against the side of the wells by placing the separation plate on the magnetic separator. Wait 2 min until all the beads have been attracted to the magnet. Transfer the supernatant containing the purified genomic DNA to the Elution Plate.

Note: The yield can be increased by 15 - 20% by using prewarmed elution buffer ( $55^{\circ}$ C) or by incubating the bead/elution buffer suspension at  $55^{\circ}$ C for 10 min.

# 6 Appendix

# 6.1 Troubleshooting

## Problem Possible cause and suggestions

Elution buffer volume insufficient

 Beads pellet must be covered completely with elution buffer

## Insufficient performance of elution buffer during elution step

• Remove residual buffers during the separation steps completely. Remaining buffers decrease efficiency of following wash steps and elution step.

## Beads dried out

• Do not let the beads dry as this might result in lower elution efficiencies.

## Partial elution in Wash Buffer MB5 already

Poor DNA yield

• Keep the beads on the magnet while dispensing Wash Buffer MB5. Do not resuspend beads in this buffer, and do not incubate beads in this buffer for more than 2 min, as this buffer is water-based and might elute the DNA already.

## Aspiration of attracted bead pellet

• Do not disturb the attracted beads while aspirating the supernatant, especially when the magnetic pellet is not visible in the lysate.

## Incubation after dispensing beads to lysate

• Mix immediately after dispensing NucleoMag Beads/Binding Buffer MB2 to the lysate.

## Insufficient washing procedure

• Use only the appropriate combinations of separator and plate, e.g. MN Square-well Block in combination with NucleoMag SEP.

Low purity

• Make sure that beads are resuspended completely during the washing procedure. If shaking is not sufficient to resuspend the beads completely mix by repeated pipetting up and down.

Problem	Possible cause and suggestions			
Suboptimal performance of DNA in downstream applications	<ul> <li><i>Carry-over of ethanol from wash buffers</i></li> <li>Be sure to remove all of the ethanolic wash solution, as residual ethanol interferes with downstream applications.</li> <li><i>Low purity</i></li> <li>see above</li> </ul>			
Carry-over of beads	<ul> <li><i>Time for magnetic separation too short</i></li> <li>Increase separation time to allow the beads to be completely attracted to the magnetic pins before aspirating any liquid from the well.</li> <li><i>Aspiration speed too high (elution step)</i></li> <li>High aspiration speed during the elution step may cause bead carry over. Reduce aspiration speed for elution step.</li> </ul>			
Cross contamination	<ul> <li>Contamination of the rims</li> <li>Do not moisten the rims of the separation plate when transferring the tissue lysate. If the rim of the wells is contaminated, seal the separation plate with self-adhering PE foil (see ordering information) before starting the shaker.</li> </ul>			

# 6.2 Ordering information

Product	Cat. No.	Pack of
NucleoMag 96 Tissue	744 300.1	1 x 96
NucleoMag 96 Tissue	744 300.4	4 x 96
NucleoMag 96 Tissue	744 300.24	24 x 96
Lysis buffer T1	740 940.25	25 ml
RNase A	740 505.50	50 mg
NucleoMag SEP	744 900	1
Square-well Blocks	740 670	20
Elution Plates, u-bottom	740 672	20
Elution Plates, flat-bottom	740 673	20
Self-adhering PE foil	740 676	50 sheets
MN Tube Strips	740 637	5 sets
Cap Strips	740 638	30 strips

# 6.3 **Product use restriction / warranty**

**NucleoMag 96 Tissue** kit components were developed, designed and sold **for research purposes only**. They are suitable **for in vitro uses only**. No claim or representation is intended for its use to identify any specific organism or for clinical use (diagnostic, prognostic, therapeutic, or blood banking).

It is rather the responsibility of the user to verify the use of the **NucleoMag 96 Tissue** kit for a specific application range as the performance characteristic of this kit has not been verified to a specific organism.

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. Please contact us if you wish an extra copy.

MACHEREY-NAGEL does not warrant against damages or defects arising in shipping and handling (transport insurance for customers excluded), or out of

accident or improper or abnormal use of this product; against defects in products or components not manufactured by MACHEREY-NAGEL, or against damages resulting from such non-MACHEREY-NAGEL components or products.

MACHEREY-NAGEL makes no other warranty of any kind whatsoever, and SPECIFICALLY DISCLAIMS AND EXCLUDES ALL OTHER WARRANTIES OF ANY KIND OR NATURE WHATSOEVER, DIRECTLY OR INDIRECTLY, EXPRESS OR IMPLIED, INCLUDING, WITHOUT LIMITATION, AS TO THE SUITABILITY, REPRODUCTIVITY, DURABILITY, FITNESS FOR A PARTICULAR PURPOSE OR USE, MERCHANTABILITY, CONDITION, OR ANY OTHER MATTER WITH RESPECT TO MACHEREY-NAGEL PRODUCTS.

In no event shall MACHEREY-NAGEL be liable for claims for any other damages, whether direct, indirect, incidental, compensatory, foreseeable, consequential, or special (including but not limited to loss of use, revenue or profit), whether based upon warranty, contract, tort (including negligence) or strict liability arising in connection with the sale or the failure of MACHEREY-NAGEL products to perform in accordance with the stated specifications. This warranty is exclusive and MACHEREY-NAGEL makes no other warranty expressed or implied.

The warranty provided herein and the data, specifications and descriptions of this MACHEREY-NAGEL product appearing in MACHEREY-NAGEL published catalogues and product literature are MACHEREY-NAGEL's sole representations concerning the product and warranty. No other statements or representations, written or oral, by MACHEREY-NAGEL's employees, agent or representatives, except written statements signed by a duly authorized officer of MACHEREY-NAGEL are authorized; they should not be relied upon by the customer and are not a part of the contract of sale or of this warranty.

Product claims are subject to change. Therefore please contact our Technical Service Team for the most up-to-date information on MACHEREY-NAGEL products. You may also contact your local distributor for general scientific information. Applications mentioned in MACHEREY-NAGEL literature are provided for informational purposes only. MACHEREY-NAGEL does not warrant that all applications have been tested in MACHEREY-NAGEL laboratories using MACHEREY-NAGEL products. MACHEREY-NAGEL does not warrant the correctness of any of those applications.

Please contact:

MACHEREY-NAGEL Germany Tel.: +49 (0) 24 21 969-270 e-mail: tech-bio@mn-net.com