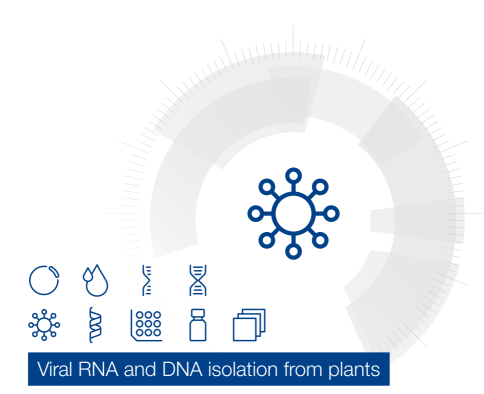
MACHEREY-NAGEL

User manual



■ NucleoSpin® Plant Pathogen

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Table of contents

1 C	1 Components		4
	1.1	Kit contents	4
	1.2	Reagents, consumables, and equipment to be supplied by user	5
	1.3	About this user manual	5
2 P	rod	uct description	6
4	2.1	The basic principle	6
4	2.2	Kit specifications	6
2	2.3	Handling, preparation, and storage of starting materials	7
4	2.4	Lysis and disruption of sample material	7
4	2.5	Elution procedures	8
3 S	Stora	age conditions and preparation of working solutions	9
4 Safety instructions		10	
4	4.1	Disposal	10
5 Protocols		11	
Į.	5.1	Nucleic acid isolation from plant material	11
6 A	ppe	endix	15
(6.1	Troubleshooting	15
(6.2	Ordering Information	18
6	6.3	Product use restriction / warranty	19

1 Components

1.1 Kit contents

	NucleoSpin [®] Plant Pathogen	
REF	10 preps 740170.10	50 preps 740170.50
Lysis Buffer PFL	8 mL	30 mL
Reduction Buffer PFR	5 mL	5 mL
Binding Buffer PFB	10 mL	30 mL
Wash Buffer PFW1	8 mL	30 mL
Wash Buffer PFW2 (concentrate)*	6 mL	25 mL
RNase-free H ₂ O	13 mL	13 mL
NucleoSpin® Plant Pathogen Columns (light blue rings)	10	50
Collection Tubes (2 mL)	30	150
Collection Tubes (1.5 mL)	10	50
Leaflet	1	1

 $^{^{\}star}$ For preparation of working solutions and storage, see section 3.

1.2 Reagents, consumables, and equipment to be supplied by user

Reagents

• 96 – 100 % ethanol (for preparation of Buffer PFW2)

Consumables

- Disposable pipette tips
- NucleoSpin® Bead Tubes Type G (optional, see section 6.2 for ordering information)
- NucleoSpin[®] Filter Columns (optional, see section 6.2 for ordering information)

Equipment

- Manual pipettes
- Centrifuge for microcentrifuge tubes
- Equipment for sample disruption and homogenization (see section 2.4)
- Thermoshaker or water bath
- Personal protection equipment (lab coat, gloves, goggles)

1.3 About this user manual

It is strongly recommended for first time users to read the detailed protocol sections of the **NucleoSpin® Plant Pathogen** kit before using this product. 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 online at www.mn-net.com.

Please contact Technical Service regarding information about any changes to the current user manual compared with previous revisions.

2 Product description

2.1 The basic principle

The **NucleoSpin® Plant Pathogen** kit is designed for the isolation of viral nucleic acid (RNA, DNA) from diverse plant material, e.g. tobacco, pepper, cauliflower, broccoli, radish, hemp, and hop.

First, plant material is mechanically disrupted (e.g., with a pipette tip or with NucleoSpin® Bead Tubes, grinding in liquid nitrogen, or any other suitable disruption method) in lysis buffer containing large amounts of chaotropic ions. This lysis buffer immediately inactivates RNases, which are present in virtually all biological materials. After transfer of the lysate supernatant, a binding solution is added which creates appropriate binding conditions which favor adsorption of RNA/DNA to the silica membrane. Washing steps with two different buffers remove salts, metabolites and macromolecular cellular components. Pure RNA/DNA is finally eluted under low ionic strength conditions with RNase-free water. The eluate contains viral nucleic acids in case of infected plant material was used as sample as well as plant nucleic acids.

The RNA preparation using <code>NucleoSpin® Plant Pathogen</code> can be performed at room temperature. The eluate, however, should be treated with care because RNA is very sensitive to trace contaminations of RNase, often found on general lab ware, fingerprints, and dust. To ensure RNA stability, keep RNA frozen at -20 °C for short-term or at -70 °C for long-term storage.

2.2 Kit specifications

- NucleoSpin® Plant Pathogen is recommended for the isolation of RNA from diverse
 plant tissues like leaves, stems, roots. The kit is not suitable for the isolation of small RNA
 (< 200 nt).
- Typically, 10-100 mg sample input is recommended per preparation.
- The kit allows the isolation of up to 20 µg RNA/DNA, suitable for downstream applications such as qRT-PCR, cDNA synthesis, Northern blotting and others.

Kit specifications at a glance		lance
	Parameter	NucleoSpin® Plant Pathogen
	Format	Mini spin column
	Use	For research use only
	Target	RNA/DNA (viral and plant)
	Handling	Centrifugation
	Sample material	≤ 100 mg plant material
	Fragment size	> 200 nt
	Typical yield	up to 20 µg total nucleic acid
	A ₂₆₀ /A ₂₈₀	1.9-2.1 (might vary for strongly infected plants)

Kit speci	fications	at a gl	ance
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 A_{260}/A_{230} ~ 2 (might vary for strongly infected plants)

Typical RIN (RNA 7-9 (for sample with strong virus symptoms, RIN might vary)

Integrity Number)

Elution volume 50 µL

Preparation time 25 min/6 preps

Binding capacity 200 µg

2.3 Handling, preparation, and storage of starting materials

RNA is not protected against digestion by plant RNase until the sample material is flash frozen or disrupted in the presence of RNase inhibiting or denaturing agents. Therefore it is important that samples are processed as fresh as possible or flash frozen in liquid N_2 immediately and stored at -70 °C. If frozen samples are used as sample material, it is very important that the sample will only thaw during the mechanical disruption in the presence of lysis buffer. Otherwise the RNA quality will be immediately impaired.

Plant material lysed in Lysis buffer PFL can be stored at -20 °C for at least 2 weeks.

Wear gloves at all times during the preparation. Change gloves frequently.

2.4 Lysis and disruption of sample material

For most plant sample material a mechanical disruption is a necessity. Several disruption options are possible.

Plant material crushing with a pipet tip

To release sufficient virus particle from infected plant material it is typically sufficient to crush plant material (without lysis buffer) within a 1.5 mL tube with a disposable pipet tip by grinding and crushing until some microliter of leaf juice appear. After addition of lysis buffer and mixing the supernatant is transfered into a fresh tube before addition of binding buffer.

Mortar, pestle and liquid nitrogen

This common sample disruption method can be used for most sample types. It typically gives excellent RNA quality; however, this method is not recommended due to the risk of cross contamination from sample to sample.

Bead tubes

NucleoSpin® Bead Tubes Type G (see section 6.3 for ordering information) are recommended in combination with a swing-mill (e.g., MM200, MM300, MM400 (Retsch®) for most plant materials. Bead Tubes typically give highest yield, avoid any cross-contamination, and enable time efficient sample disruption.

The MN Bead Tube Holder should not be used for disruption of plant material with NucleoSpin® Bead Tubes Type G because it is usually insufficient.

When using bead tubes for sample disruption the subsequent filtration with a NucleoSpin® Filter (see ordering information) is recommended in order to remove debris from the lysate.

2.5 Elution procedures

It is possible to adapt the elution method and elution volume in order to achieve optimal RNA concentrations for the respective downstream application. In addition to the standard method described in the individual protocols (recovery rate about 70 – 90 %), modifications are possible.

- High yield: Perform two elution steps with the volume indicated in the individual protocol.
 About 90 100 % of bound nucleic acid will be eluted.
- High yield and high concentration: Elute with the standard elution volume and apply the eluate once more onto the column for re-elution.

Eluted RNA should immediately be kept on ice for optimal stability. For short-term storage freeze at -20 $^{\circ}$ C, for long-term storage freeze at -70 $^{\circ}$ C.

3 Storage conditions and preparation of working solutions

Attention: Buffers PFL and PFW1 contain chaotropic salt. Wear gloves and goggles!

CAUTION: Lysis Buffer contains guanidine hydrochloride 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.

All kit components should be stored at $15-25\,^{\circ}\mathrm{C}$ and are stable until: see package label. Storage at lower temperatures may cause precipitation of salts.

Such precipitates can be easily dissolved by incubating the bottle at 40 °C before use.

Before starting any NucleoSpin® Plant Pathogen protocol prepare the following:

Wash Buffer PFW2: Add the indicated volume of 96-100% ethanol (see table below) to Wash Buffer PFW2. Mark the label of the bottle to indicate that ethanol was added. Wash Buffer PFW2 can be stored at 15-25% for at least one year.

	A Plant and Fungi	
REF	10 preps 740120.10	50 preps 740120.50
Wash Buffer PFW2 (concentrate)	6 mL Add 24 mL ethanol	25 mL Add 100 mL ethanol

4 Safety instructions

When working with the **NucleoSpin® Plant Pathogen** kit, wear suitable protective clothing (e.g., lab coat, disposable gloves, and protective goggles). For more information consult the appropriate Material Safety Data Sheets (MSDS available online at **http://www.mn-net.com/msds**).



Caution: Guanidine hydrochloride in Buffer PFL can form highly reactive compounds when combined with bleach! Thus, do not add bleach or acidic solutions directly to the sample preparation waste.

The waste generated with the **NucleoSpin® Plant Pathogen** kit has not been tested for residual infectious material. A contamination of the liquid waste with residual infectious material is highly unlikely due to strong denaturing lysis buffer treatment, but it cannot be excluded completely. Therefore, liquid waste must be considered infectious and should be handled and discarded according to local safety regulations.

4.1 Disposal

Dispose hazardous, infectious or biologically contaminated materials in a safe and acceptable manner and in accordance with all local and regulatory requirements.

5 Protocols

5.1 Nucleic acid isolation from plant material

Before starting the preparation:

- Check if Wash Buffer PFW2 was prepared according to section 3.
- Set thermoshaker or water bath to 56 °C.
- During storage, especially at low temperatures, a precipitate may form in some buffers.
 Such precipitates can be easily dissolved by incubating the bottle at 40 °C before use.

1 Homogenize sample - release of viruses

Option A: Sample crushing with pipet tip

Place approximately 10 – 100 mg sample material (typically plant leaf) into a 1.5 mL tube (not provided).

Using a disposable pipet tip, grind the sample by circular and stinging movements until several microliter plant juice appears.

Note: Appearance of plant juice might not be obvious for partially dried samples. For dry sample materials the homogenization option B (bead tubes) is recommended.

Add 300 µL Buffer PFL to the sample.

Add 12 µL Buffer PFR to the tube.

Note: A Premix of PFL and PFR can be prepared for several samples on a daily basis.

300 μL PFL 12 μL PFR

Grind sample

Mix

Vortex the tube for approximately 10 seconds.

Incubate lysis tube for ${\bf 5}$ min at ${\bf 56}$ °C. Mix occasionally or use the thermoshaker device.



56 °C, 5 min

Centrifuge for 1 \min at 14,000 \times g in order to sediment cell debris.



14,000 x g,

<u>Note:</u> If the cell debris pellet is not sufficiently solid, centrifuge for a longer time (e.g., 3 min) and / or at 20,000 x g.



Transfer sample

Transfer approximately 300 μL supernatant into a fresh tube (not provided).

Option B: Bead Tubes (not provided)

Add 300 µL Buffer PFL into NucleoSpin® Bead Tubes Type G.



300 µL PFL 12 µL PFR

Transfer

Add 12 μL Buffer PFR to the tube.

Transfer sample to the NucleoSpin® Bead Tube Type G.

sample

Place the Bead Tube into a swing-mill and **agitate twice** for **30 s** at 30 Hz with intermediate position change (please refer to the manufacturers' instructions for proper use of the instrument)



Incubate NucleoSpin® Bead Tube Type G for 5 min at 56 °C.

56 °C, 5 min

Remove steel balls from the Bead Tube.

Į

Attention: Removal of steel balls is necessary in order to avoid tube damage during subsequent centrifugation.

Centrifuge for 1 min at 14,000 x α in order to sediment cell

14,000 x g,

debris.

Note: If the cell debris pellet is not sufficiently solid, centrifuge

for a longer time (e.g. 3 min) and / or at 20,000 x g.

Continue with the clear supernatant.

Filtrate Lysate

Insert a NucleoSpin® RNA Plant and Fungi Filter Column (not provided, please inquire) into a Collection Tube (see ordering information).



Load lysate

<u>Note:</u> Alternatively use a 2 mL microcentrifuge tube with lid (not provided). This facilitates mixing by vortexing in step 2.

Load the clear lysate from the bead tube onto the column.

Centrifuge for 1 min at 14,000 x g.

14,000 x g, 30 s

<u>Note:</u> In some cases a small pellet will form. This pellet does not have to be removed and can be processed together with the supernatant.

Note: If the sample does not pass the column completely, centrifuge at 20,000 x g for additional 3 min.

Transfer approximately 300 μ L supernatant into a fresh tube (not provided).

2 Adjust nucleic acid binding conditions

Add 300 μ L Buffer PFB to the lysate and mix by pipetting.

Incubate for 5 min at room temperature.



300 µL PFB RT, 5 min

3 Bind nucleic acid

For each preparation take one NucleoSpin® Plant Pathogen Column (light blue ring) preassembled with a Collection Tube.

Load the sample (typically approx. 600 µL) but not more than 650 µL onto the NucleoSpin Plant Pathogen column.

Centrifuge for 30 s at 14,000 x g.

Discard collection tube with flowthrough and insert the column into a fresh Collection Tube (2 mL, provided).



Load 650 µL sample

 $14,000 \times q$ 30 s



14,000 x q, 30 s

Wash and dry silica membrane

1st wash

Add 500 µL Buffer PFW1 onto the column.

Centrifuge for 1 min at 14.000 x a.

Discard collection tube with flowthrough and insert column into a fresh Collection Tube (2 mL, provided).



500 uL PFW1

14,000 x g, 1 min

2nd wash

Add 500 µL Buffer PFW2 onto the column.

Centrifuge for 1 min at 14,000 x g.

Discard flowthrough and reuse collection tube.



500 μL PFW2

14,000 x q, 1 min

3rd wash

Add 500 µL Buffer PFW2 onto the column.

Centrifuge for 1 min at 14,000 x g.

Discard flowthrough and discard collection tube unless the following additional wash step is included.



500 μL PFW2

 $14,000 \times q$ 1 min

Optional:. For some plant samples comprising less than 10 mg plant material the third wash step might be omitted.

Add 500 µL Wash Buffer PFW2 onto the column.

Centrifuge for 1 min at 14,000 x g.

Discard collection tube with flowthrough.

5 Elute nucleic acid

Insert column into a fresh Collection Tube (1.5 mL, provided).

Add 50 µL RNase-free H₂O onto the column.

Incubate for approximately 1 min at room temperature.

Centrifuge for 1 min at 14,000 x g.

If higher nucleic acid concentrations are desired, elution can be done with 40 μ L. Overall yield, however, will decrease when using smaller volumes.



For further alternative elution procedures see section 2.5

6 Appendix

6.1 Troubleshooting

Problem	Possible cause and suggestion		
	Too much sample material		
Clogged	Use less sample material		
NucleoSpin® Plant Pathogen Column	 and / or centrifuge for 3 min at 20,000 x g prior to addition of ethanol to the lysate 		
	• and / or centrifuge the column for 3 min at 20,000 xg after loading the ethanolic lysate onto the column .		
	Highly symptomatic, partially dried or coloured leaves		
	 Samples in advanced stages of viral infection might show reduced nucleic acid yield. If possible, use material showing less viral infection symptoms. 		
Poor RNA/DNA	Sample with high starch content was heat incubated		
quality or yield	 Samples such as potato tubers, maize kernels, wheat kernels and similar should not be incubated at elevated temperatures during the RNA purification procedure 		
	However, banana fruit tissue of ripe fruits should be heat incubated in order to obtain high RNA yield.		
Poor RNA/DNA	Washing steps not sufficient		
purity and or colored silica membrane / eluate	Perform an additional wash step with Buffer PFW1.		

Problem

Possible cause and suggestion

RNase contamination

 Create an RNase-free working environment. Wear gloves during all steps of the procedure. Change gloves frequently. Use of sterile, disposable polypropylene tubes is recommended. Keep tubes closed whenever possible during the preparation. Glassware should be oven-baked for at least 2 hours at 250 °C before use.

Insufficient sample quality

 Control sample harvest, storage, and lysis. Make sure that samples are harvested, stored and lysed adequately in order to preserve RNA integrity. Whenever possible, use fresh material. If this is not possible, flash freeze the samples in liquid nitrogen. Samples should always be kept at -70 °C. Never allow tissues to thaw before addition of Lysis Buffer. Perform disruption of samples in liquid nitrogen.

Poor RNA quality or yield

Insufficient sample disruption

 Choose a different disruption method. If one disruption method gives unsatisfactory results, try an alternative disruption method.

Reagents not applied or restored properly

- Prepare Buffer PFW2 by adding ethanol according to the description.
- Sample and reagents have not been mixed completely. Always vortex vigorously after each reagent has been added.

Kit storage

- Store kit components at room temperature. Storage at low temperature my cause salt precipitation.
- Keep bottles tightly closed in order to prevent evaporation or contamination

lonic strength and pH influence A_{260} absorption as well as ratio A_{260}/A_{280}

For adsorption measurement, use 5 mM Tris pH 8.5 as diluent.
 Please see also:

Poor RNA quality or yield (continued)

- Manchester, K L. 1995. Value of A_{260} / A_{280} ratios for measurement of purity of nucleic acids. Biotechniques 19, 208-209.
- Wilfinger, W W, Mackey, K and Chomczyski, P. 1997.
 Effect of pH and ionic strength on the spectrophotometric assessment of nucleic acid purity. Biotechniques 22, 474–481.

Problem	Possible cause and suggestion
	Carry-over of contaminants
	 Carefully load the lysate to the NucleoSpin® RNA Plant Pathogen Column and try to avoid a contamination of the upper part of the column and the column lid.
Low A_{260}/A_{230} ratio	 Make sure that a sufficient amount / concentration of RNA is used for quantification so that the A₂₃₀ value is significantly higher than the background level.
	\bullet Measurement of low amount / concentration of RNA will cause unstable $\rm A_{260}/A_{230}$ ratio values.
Centrifuge not capable to reach 14,000 xg	Use 11,000 x g or max speed of your centrifuge. Increase centrifugation time if necessary.
	Carry-over of ethanol or salt
	 Do not let the flowthrough touch the column outlet after the wash steps. Be sure to centrifuge at the corresponding speed for the respective time in order to remove last wash buffer completely.
Suboptimal performance of RNA in downstream	 Check if wash buffer has been equilibrated to room temperature before use. Washing at lower temperatures lowers efficiency of salt removal by wash buffer.
experiments	Store isolated RNA properly
	 Eluted RNA should always be kept on ice for optimal stability since trace contaminations of omnipresent RNases (general lab ware, fingerprints, dust) will degrade the isolated RNA. For short term storage freeze at -20 °C, for long term storage freeze at 70 °C.
	Beads not removed from Bead Tube
Damaged Bead Tubes Type G	 Remove steel balls from the Bead Tube by placing a magnet on top of the lid. Invert the tube once. Open the tube and remove steel balls attached to the lid.

6.2 Ordering Information

Product	REF	Preps/Pack of
NucleoSpin® Plant Pathogen	740170.10/.50	10/50
Lysis Buffer PFL	740122.30	30 mL
Reduction Buffer PFR	740123.5	5 mL
Wash Buffer PFW1	740119.30	30 mL
Wash Buffer PFW2 (concentrate)	740939/.1	50 mL/200 mL
NucleoSpin® Bead Tubes Type G	740817.50	50
MN 96 Bead Plate Type G	740855.1/.4	1 × 96/4 × 96
Collection Tubes (2 mL)	740600	1000

6.3 Product use restriction/warranty

All MACHEREY-NAGEL products are designed for their intended use only. They are not intended to be used for any other purpose. The description of the intended use of the products can be found in the original MACHEREY-NAGEL product leaflets. Before using our products, please observe the instructions for use and the safety instructions from the respective Material Safety Data Sheet of the product.

This MACHEREY-NAGEL product is carrying documentation stating specifications and other technical information. MACHEREY-NAGEL warrants to meet the stated specifications. The provided warranty is limited to the data specifications and descriptions as given in the original MACHEREY-NAGEL literature. No other statements or representations, written or oral, by MACHEREY-NAGELS employees, agents or representatives, except written statements signed by a duly authorized officer of MACHEREY-NAGEL are authorized. They should not be relied upon by the costumer and are not a part of a contract of sale or of this warranty.

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Please contact:

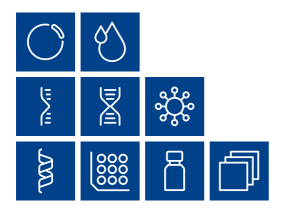
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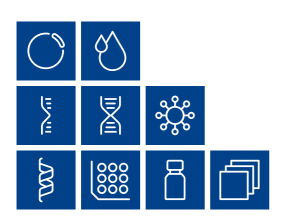
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Plasmid DNA
Clean up
RNA
DNA
Viral RNA and DNA
Protein
High throughput
Accessories
Auxiliary tools



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