

1-5 mL E. coli

culture

11,000 x g,

30 s

+ 250 µL A2

Mix gently

RT, 5 min

+ 350 µL A3

Mix

## NucleoSpin<sup>®</sup> Plasmid / Plasmid (NoLid) – isolation of high-copy plasmid DNA using a

vacuum manifold (Rev. 01, June 2016)

This protocol is only a supplement to the kit's general user manual. Please refer to the kit manual for more detailed information regarding safety instructions, product-specific disclaimers, and especially preparations needed before starting the procedure. The latest version of the user manual is available at **www.mn-net.com/usermanuals** or can be requested from our technical service (tech-bio@mn-net.com). Safety data sheets (SDS) can be downloaded from **www.mn-net.com/MSDS**.

## 1 Cultivate and harvest bacterial cells

Use **1–5 mL** of a saturated *E. coli* culture and pellet cells in a standard benchtop microcentrifuge for **30 s** at **11,000 x** *g*. Discard supernatant and remove as much of the liquid as possible.

<u>Note</u>: For isolation of low-copy plasmids volumes of culture and buffers need to be adapted. Refer to section 5.2 in the NucleoSpin<sup>®</sup> Plasmid / Plasmid (NoLid) user manual.

## 2 Lyse cells

Add 250 μL Buffer A1. Resuspend the cell pellet<br/>completely by vortexing or pipetting up and down. Make<br/>sure no cell clumps remain before addition of Buffer A2!+ 250 μL A1<br/>ResuspendAttention: Check Buffer A2 for precipitated SDS prior

Attention: Check Buffer A2 for precipitated SDS prior to use. If a white precipitate is visible, warm the buffer for several minutes at 30–40 °C until any precipitate is dissolved. Mix thoroughly and cool buffer down to room temperature (18–25 °C).

Add **250 μL Buffer A2**. **Mix gently** by inverting the tube **6–8 times**. Do not vortex to avoid shearing of genomic DNA. Incubate at **room temperature** for a maximum of **5 min** or until the lysate appears clear.

Add **350 µL Buffer A3**. **Mix thoroughly** by inverting the tube **until the blue samples turn colorless completely**! Do not vortex to avoid shearing of genomic DNA!



\* Reduction of atmospheric pressure



Place the NucleoSpin<sup>®</sup> Plasmid / Plasmid (NoLid) Column into a Collection Tube (2 mL). Centrifuge for **1 min** at **11,000 x** g to remove Buffer A4 completely. Make sure the spin column does not come in contact with the flowthrough while removing it from the centrifuge and the collection tube.

<u>Note</u>: Residual ethanol from Buffer A4 might inhibit enzymatic reactions. Total removal of ethanol can be achieved by incubating the columns for 2–5 min at 70 °C prior to elution.

## 7 Elute DNA

Place the NucleoSpin<sup>®</sup> Plasmid / Plasmid (NoLid) Column into a new 1.5 mL microcentrifuge tube (not provided).

Add **50 µL Buffer AE** and incubate at **room temperature** (18–25 °C) for **1 min**. Centrifuge for **1 min** at **11,000 x** *g*.

-0.2 to -0.4 bar\*, 5 min

> 11,000 x *g*, 1 min

+ 50 µL AE

RT, 1 min

11,000 x *g*, 1 min

\* Reduction of atmospheric pressure