# MACHEREY-NAGEL

# TakaRa Available from and supported by Takara Bio USA, Inc.

# NucleoMag® Blood 200 μL

Automated purification of genomic DNA from 200 µL blood samples on the platform KingFisher® Flex



#### Introduction

The isolation of genomic DNA from whole blood samples is an initial and crucial step for various diagnostic workflows. Clinical applications such as genotyping, HLA typing, biomarker discovery, newborn screening, and pharmacogenetics are widely performed in laboratories worldwide. To enable a state of the art molecular diagnostic, the sensitivity and performance of biomolecular detection methods like aPCRs, next-generation sequencing, and microarray analysis, is constantly improved. A main aspect to face these workflow requirements, is to facilitate the extraction of highly pure DNA in substantial amounts. To provide a fast and consistent sample processing, MACHEREY-NAGEL designed the NucleoMag® Blood 200 µL kit, specialized for automated purification of genomic DNA from 200 µL whole blood samples (fresh or frozen, EDTA or citrate treated) in a 96-well format. The optimized protocol allows the processing of 96 samples within 45 minutes including a sample lysis on the KingFisher® platform.

## Product at a glance

NucleoMag <sup>®</sup> Blood 200 μL			
Technology	Magnetic bead technology		
Sample material	$\leq$ 200 $\mu L$ whole blood (fresh or frozen, EDTA or citrate treated)		
Preparation time	Approx. 45 min on KingFisher® Flex for 96 samples incl. sample lysis		
Typical yield	2–8 μg (200 μL blood; depending on sample quality)		
Elution volume	50–100 μL		
Binding capacity	0.4 μg/μL beads		
Elution volume	2–8 μg (200 μL blood; depending on sample quality) 50–100 μL		

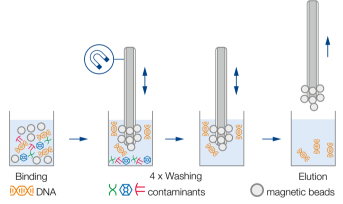
King Fisher® Flex	
Sample volume	20–5000 μL
Capacity	24/96 samples (8 plates per deck)
Heating/cooling	4–96 °C
Size/weight	60 x 38 x 68 cm/28 kg

# Material and Methods

Whole blood (fresh, frozen, treated with EDTA or citrate) is lysed at room temperature with Lysis Buffer MBL1 and Proteinase K. Following lysis incubation, binding of DNA to the NucleoMag® B-Beads is achieved by the provision of Binding Buffer MBL2.



Subsequent DNA isolation is performed on the automation platform KingFisher® Flex. The isolation principle is based on reversible adsorption of nucleic acids to paramagnetic NucleoMag® B-Beads under appropriate buffer conditions.

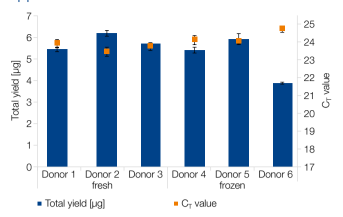


#### Workflow on automation platform

After magnetic separation, the NucleoMag® B-Beads are washed four times to remove contaminants and salts using three different wash buffers (MBL3, 80% ethanol and MBL4). Highly pure genomic DNA is finally eluted under low ionic strength conditions in a slightly alkaline elution buffer (MBL5).

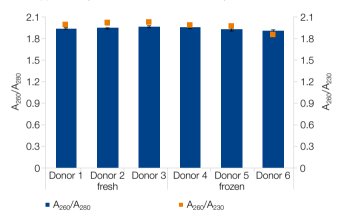


## Application data



#### Isolation of genomic DNA from fresh and frozen human blood samples

DNA was isolated from fresh and frozen 200  $\mu$ L human blood samples (n = 8) using the NucleoMag® Blood 200  $\mu$ L kit on a KingFisher® Flex platform. The total yield was determined by UV-spectrometry (dark blue bars). A subsequent qPCR analysis (orange squares) was performed with a Taqman® Probe for a 250 bp  $\beta$ -Actin amplicon using the SensiFast Probe Lo-ROX kit from Bioline on an Applied Biosystems® 7500 Real-Time PCR System.

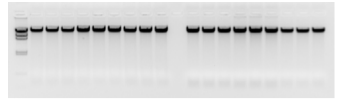


#### Purity of isolated nucleic acids from fresh and frozen human blood samples

DNA was isolated from fresh and frozen 200  $\mu$ L human blood samples (n = 8) using the NucleoMag® Blood 200  $\mu$ L kit on a KingFisher® Flex platform. The purity was determined by UV-spectrometry resulting into an average  $A_{260}/A_{280}$  value of 1.93  $\pm$  0.2 (dark blue bars) and into an average  $A_{260}/A_{230}$  value (orange squares) of 1.96  $\pm$  0.05.

#### M Donor 1 Donor 2 Donor 3

Donor 4 Donor 5 Donor 6



#### Integrity of isolated nucleic acids

The integrity of the isolated nucleic acids from fresh (donor 1-3) and frozen (donor 4-6) blood samples was analyzed by gel electrophoresis (10  $\mu$ l per eluate; 1 % TAE-gel; M: Lambda DNA/Hind III – Thermo Scientific).

# Automate your DNA extraction from blood samples

MACHEREY-NAGEL delivers a ready to go solution for your high throughput DNA extraction from various blood samples, such as fresh, frozen, treated with EDTA or citrate. We adapted the NucleoMag® Blood 200  $\mu$ L kit on instruments of the KingFisher® series to speed up your nucleic acid purification workflow.

- Reliable performance and excellent yields from various blood sample material
- Speed up your DNA extraction by processing of 96 samples in less than 45 minutes (including sample lysis)

### Ordering information

Product	Specifications	Preps	REF
NucleoMag <sup>®</sup> Blood 200 μL	Kit based on magnetic bead technology for the isolation of genomic DNA from 200 $\mu$ L blood samples including NucleoMag <sup>®</sup> B-Beads, buffers, and Proteinase K	1 x 96 4 x 96	744501.1 744501.4
KingFisher® Accessory Kit B*	Deep-well Plates, Deep-well Tip Combs, and Elution Plates for 4 x 96 NucleoMag $^{8}$ Blood 200 $\mu$ L preps using the KingFisher $^{8}$ Flex platform	4 x 96	744951

<sup>\*</sup>For use on KingFisher® Flex

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