

ExpressArt Mag FFPE Clear DNAREady kit

FFPE DNA isolation kit

Catalogue No. 9020-A100

for 100 DNA isolations from archival paraffin samples

with Magnetic Beads for DNA isolation

No separate paraffin removal step !

This protocol provides the required laboratory procedures

**For additional information about ExpressArt technology and products
see**

**<http://www.amp-tec.com/products>
» more information**

Kit Contents

Box I	Volume per DNA isolation	Kit contents for 100 isolations
Lysis Buffer (LB)	870 µl	87 ml
FFPE Clear a formulation of <i>non-toxic, non-odorous</i> organic solvents	500 µl	55 ml
Demodification Solution (DeS)	130 µl	15 ml
Nuclease-free water	50 µl	5 ml

Box II	Volume per DNA isolation	Kit contents for 100 isolations
NucleoGuard	5 µl	0.6 ml
Proteinase K, PCR Grade*	30 µl	powder
DNase-Free RNase**	45 µl (working solution)	750 µl concentrate
DiB, Dilution buffer for DNase	39.13 µl	2x 2 ml
Magnetic Beads Stock	100 µl	10 ml

* Dissolve proteinase K (powder) in 4 ml water (25 mg/ml); store at -20°C.

**Prepare “RNase working solution”

Prepare appropriate volumes as stock solution, required volumes for one sample:

7.5 µl RNase concentrate + 37.5 µl water.

Unused Aliquots are stored at -20°C

Please note, these reagents are not included in the kit

Isopropanol

Required instruments

Thermomixer (Eppendorf #5350 000.013 or “Thriller” from PEQLAB #91-7010) or equivalent

Magnetic separator

Tissue samples

Please note:

1) to avoid negative surface effects (contact to air), do not use the first 2 slides for preparation of nucleic acids.

2) no need for immediate processing of slides,

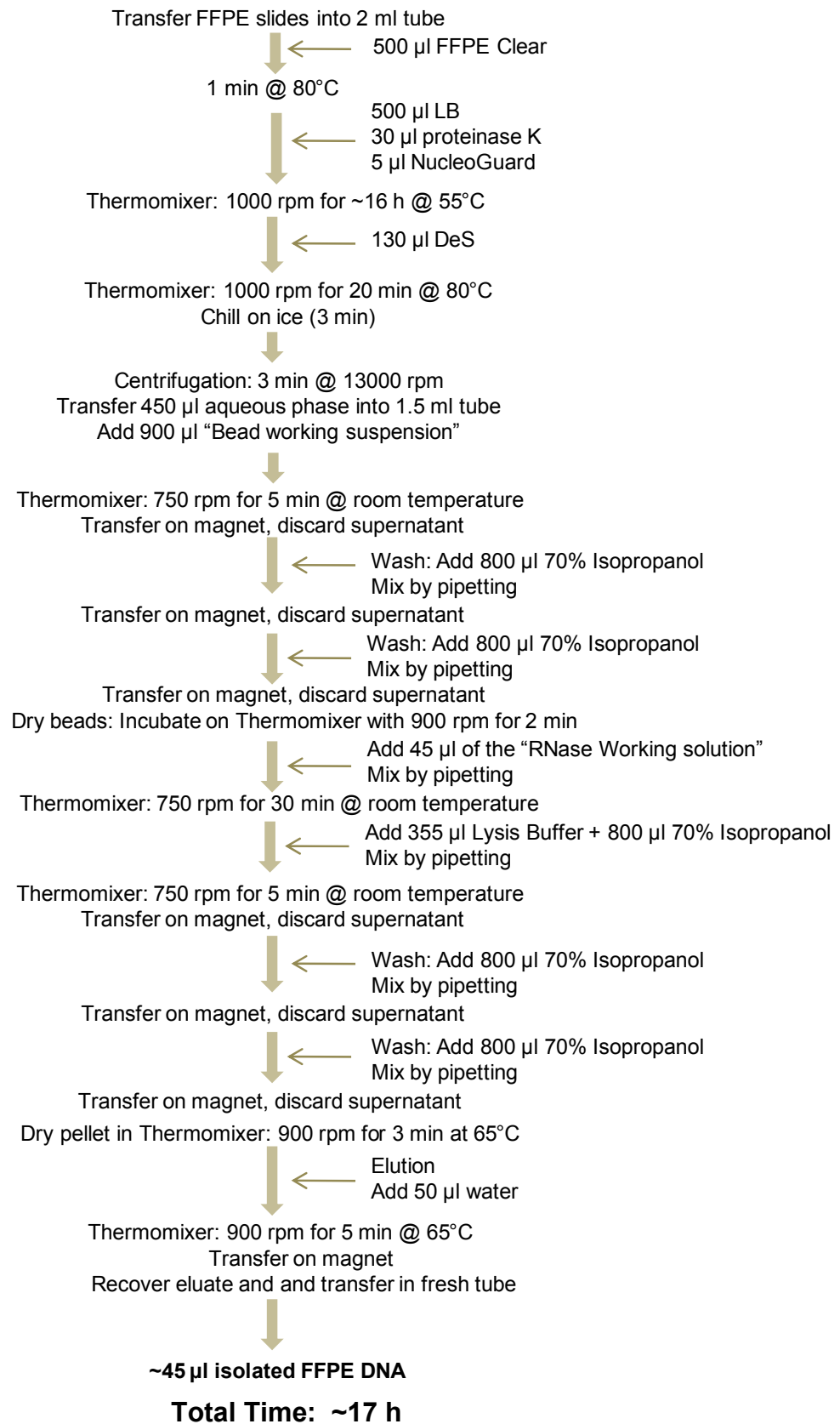
freezing and short-term storage (up to 3 days) at -80°C is even advantageous:

there is no negative effect on DNA quality; frequently, DNA yields will increase significantly

Up to 5 FFPE slides (10 µm thick with appr. 1-2 cm² of tissue) per isolation can be used in the lysis procedure

Work Flow

RNA isolation from FFPE samples using the ExpressArt Mag FFPE Clear RNAREady kit



1. Sample preparation

Insert paraffin sections in reaction tube

Immediately (no prior paraffin removal required) add 500 µl **FFPE Clear**

(**Note:** For immediate processing, it can be more convenient to prepare tubes with FFPE Clear, and transfer paraffin sections directly into the solvent, avoiding "electrostatic problems")

Heat for 1 min @ 80°C.

Add 500 µl **Lysis Buffer (LB)** + 30 µl Proteinase K (25 mg/ml) + 5 µl NucleoGuard

2. Lysis: Incubation in Thermomixer (1000 rpm) **overnight (ca. 16 h) @ 55°C**

3. Demodification: Add 130 µl **Demodification Solution (DeS)** (to 535 µl lysate)

Vortex, then incubation in Thermomixer (1000 rpm) for 20 min @ 80°C

Chill on ice (3 min)

4. DNA Purification

Before you start, prepare "bead suspension"

For 1 sample: Transfer 100 µl of Magnetic Beads Stock into 1.5 ml tube, place on magnet, discard liquid and resuspend pellet in 900 µl 70% isopropanol.

For multiple samples, use two suspension steps

Example for **10 samples:** **Step 1:** Transfer 1 ml of Magnetic Beads Stock into 1.5 ml tube, place on magnet, discard liquid and resuspend pellet in 1 ml of 70% isopropanol by pipetting.

Step 2: Transfer into 15 ml tube, add 8 ml 70% isopropanol and mix thoroughly.

4.1. Recovery of the aqueous lysate

Centrifuge lysate (from step 3): 3 min @ 13'000 rpm

This results in three zones: pellet at the bottom (debris), **aqueous phase** with the lysate in the middle and organic supernatant at the top.

4.2. DNA binding to beads

Transfer 450 µl of the **aqueous phase** (avoid transfer of the organic supernatant) in 1.5 ml reaction tube

Mix 450 µl of the aqueous lysate with 900 µl bead suspension by pipetting (= 1350 µl)

Incubate on Thermomixer (750 rpm) for 5 min @ room temperature

Transfer on magnet, discard supernatant

4.3. First washing steps

Wash 2x with 800 µl 70% Isopropanol: Mix by pipetting, place on magnet, remove liquid

4.4. RNA-Digestion

Incubate on Thermomixer (900 rpm) for 2 min @ room temperature to dry the beads.

Add 45 µl RNase working solution to beads, mix by pipetting.

Incubate on Thermomixer (750 rpm) for 30 min @ room temperature

4.5. RNase inactivation and DNA recovery

Add 355 µl Lysis Buffer + 800 µl 70% Isopropanol and mix by pipetting

Incubate on Thermomixer (750 rpm) for 5 min @ room temperature

Transfer on magnet, discard supernatant

4.6. Second washing steps

Wash 2x with 800 µl 70% Isopropanol: Mix by pipetting, place on magnet, remove liquid

4.7. Elution

Incubate on Thermomixer (900 rpm) for 3 min at 65°C to dry the beads completely

Add 50 µl water, mix by pipetting and make sure to soak the beads completely

Incubate on Thermomixer (900 rpm) for 5 min @ 65°C

Transfer on magnet and transfer supernatant in fresh tube

DNA is recovered in ~45 µl.

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