ULS™

ULS™ aRNA Bioti<mark>n</mark> Labeling Kit

For 25 labeling reactions

Product code FA-010

For laboratory use only Research purposes only



KREATECH'S ULS™ aRNA Biotin Labeling Kit

This kit is intended for RESEARCH USE ONLY. IT IS NOT INTENDED FOR DIAGNOSTIC APPLICATIONS and/or COMMERCIAL PURPOSES.

Important

Open the kit immediately and store all components as instructed on page 6

- Read the entire Instruction manual before starting your experiment.
- Do not mix reagents from different kits.
- During the preparation of reagents and throughout the entire procedure please observe the safety regulations issued for laboratories concerning handling of samples.
- Dispose of reagents according to relevant local regulations.
 Take appropriate safety precautions such as wearing a lab coat, gloves and eye protection.

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A. Assay Materials

I. Components and Storage

Component	Amount	Storage
BIO-ULS™	125 µL	4°C
10 x Labeling Buffer	100 µL	4°C
KREA <i>pure</i> ™ columns	25 pcs	4°C
KREA <i>block</i> ™	1.5 mL	-20°C*

^{*} KREAblock is shipped at 4°C, store at -20°C upon delivery.

II. Reagents and Buffers not Included in Kit Fragmentation reagents (Ambion Cat # 8740)

B. General Information

I. Background

Gene expression profiling using DNA-microarrays has been growing rapidly over the last decade. When working with material from fine needle biopsy, laser capture micro-dissection material and in fact with patient material in general, the amount of target material is limited. This has necessitated the need to carry out target amplification, generally by linear amplification based on a protocol first described by van Gelder and Eberwine¹. This procedure is based on the reverse transcription from mRNA into double stranded cDNA using an Oligo(dT) primer containing a T7 RNA polymerase promoter sequence. Linear amplification is achieved during the subsequent IVT reaction using T7 polymerase with cDNA as a functional template.

The ULS aRNA Biotin Labeling Kit has been designed to enable the generation of unmodified amplified aRNA using natural nucleotides, which can then be labeled with BIO-ULS. This kit is intended for Affymetrix GeneChip® users.

II. Principle of ULS Labeling

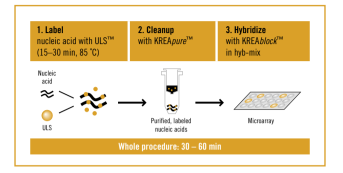
The proprietary ULS technology is based on the stable binding properties of a platinum complex to biomolecules. The ULS molecule consists of a platinum complex, a detectable molecule and a leaving group which is displaced upon reaction with the target. This ULS molecule forms a co-ordinative bond, firmly coupling the ULS to the target. ULS labels DNA, and RNA by binding to the N7 position of guanine. In proteins, ULS binds to nitrogen and sulphur containing side chains of methionine, cysteine and histidine (see figure inside back cover). ULS is available coupled to a variety of labels and haptens, including fluorochromes and biotin. ULS thereby enables one-step non-enzymatic labeling of nucleic acids to be achieved within 30 minutes. ULS labeling can be performed with or without enzymatic amplification, prior to labeling.

III. ULS aRNA Biotin Labeling Process

The procedure of expression analysis with the ULS aRNA Biotin Labeling Kit is as follows:

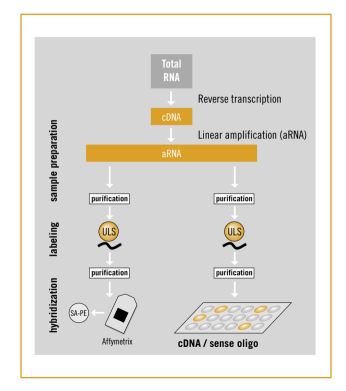
- aRNA is generated from isolated total RNA via linear amplification using natural unmodified nucleotides
- 2. aRNA is non-enzymatically labeled with ULS reagent (30 min)
- 3. The labeled aRNA is purified with the KREApure column
- 4. The labeled aRNA is fragmented
- 5. The fragmented and labeled aRNA is hybridized to a microarray in the presence of KREA*block*

Figure 2: A 30-60 minute protocol for DNA microarray applications



IV. Schematic Overview of the ULS aRNA Biotin Labeling Process

Figure 3. Schematic Overview of ULS aRNA Biotin Labeling Process.



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C. Protocol

I. Total RNA Isolation

A wide variety of RNA isolation techniques are available, e.g using Trizol (Invitrogen) extraction followed by RNeasy column purification (QIAGEN). Irrespective of which isolation procedure is used the RNA material should be free from DNA and other contaminants. Assessment of the purity and yield of your RNA should be carried out by

- Running your RNA on a 1% agarose gel. The integrity of the total RNA is determined by observing the ribosomal bands
- Determining the OD₂₆₀. For all RNAs OD_{260/280} should be >1.9 and OD_{260/230} should be >2.1

II. Linear Amplification

ULS technology allows labeling of aRNA generated from a variety of commercially available kits, e.g. Message amp aRNA amplification kit (Ambion), Mega script T7 kit (Ambion).

Important! With the ULS protocol, amplification is carried out prior to labeling with the detection moieties. Enzymatic reactions should be carried out using only unmodified nucleotides. This results in better yields, longer fragments and a more stable amplified RNA sample.

Furthermore, aRNA samples need to be clean of divalent cations (e.g. Mg²⁺) salt and other (wash) buffer components which could disturb the labeling efficiency.

Be aware that some components in silica based purification systems can inhibit the ULS reaction. A final wash step using 80% ethanol (PA) before elution should prevent this. Alternatives: alcohol precipitation of eluted material or cleanup using an EDTA, sephadex combination.

III. aRNA Labeling

a. ULS-Labeling Procedure

Briefly spin all required reagents to collect contents of tubes

- Take 20 μg of aRNA ensure final concentration in labeling reaction is above 50 ng/μL
 - In general suboptimal modification degrees of the labeled material are achieved if final concentration of the aRNA in the labeling mixtures is below 50 ng/µL)
- 2. Add 5 μL of BIO-ULS per 20 μg aRNA
- 3. Add 1/10 volume of 10x Labeling solution
- 4. Adjust with RNase-free water to final volume and mix by pipetting (see example set-up below)
- 5. Label sample by incubation for 30 minutes at 85°C
- Place samples on ice, spin down to collect contents of tube before proceeding with purification using the KREApure columns

Example of BIO-ULS labeling of 20 μg aRNA

aRNA (20 μg) + RNAse free water	22.0 µL
BIO-ULS	5.0 μL
10 x labeling solution	3.0 µL
Total volume	30 μL

b. Dye Removal Using KREA*pure* Columns

Removal of free ULS label using KREApure columns

(20800 g is equivalent to 14,000 rpm on eppendorf 5417C)

- 1. Resuspend column material by vortexing
- 2. Loosen cap 1/4 turn and snap off the bottom closure
- 3. Place the column in a 2 mL collection tube
- 4. Pre-spin the column for 1 minute at 20800 g
- 5. Discard flow-through and re-use collection tube
- 6. Wash the column with 300 µL RNase free water

- 7. Discard collection tube and flow-through
- 8. Put column in a new (RNase free) 1.5 mL micro centrifuge tube
- 9. Add ULS-labeled aRNA on to column bed
- 10. Spin column for 1 minute at 20800 g
- 11. Flow through is purified labeled aRNA

c. aRNA Fragmentation

- 1. Transfer labeled sample to a microfuge tube and add 1/10 volume of 10 x fragmentation buffer (Ambion) to decrease the fragment size to 60-200 bases. (e.g. 4 μ L in final volume of 40 μ L)
- 2. Incubate at 70°C for 15 minutes
- 3. Spin the vial briefly and add $1\mu L$ of stop solution Ambion), mix by pipetting (the labeled aRNA can form aggregates which dissolve by pipetting) and place on ice until further use

IV. Preparation of Labeled Material for Hybridization

- 1. KREA*block* should be added to 1/4 final volume of the hybridization mixture (e.g. 25 μ L of KREA*block* in a 100 μ L hybridization volume)
- 2. Hybridize and wash slides according to own protocol

D. Trouble Shooting

I. Total RNA and aRNA Preparation and Analysis

Problem	Possible Reasons and Suggestions
OD ₂₆₀ not within	Cause: Impure RNA
parameters	Remedy: Repeat RNA clean-up kit using
	commercial kit or precipitate RNA and
	dissolve again

II. ULS Labeling

Problem	Possible Reasons and Suggestions
Degree of	Cause: There may be salt present which
labeling too low	disturbs labeling
	Remedy: Clean up aRNA and ensure final
	80% ethanol wash step is used with
	silica based columns (see C II)
	Cause: Incorrect ratio of labeling reagent
	to aRNA
	Remedy: Ensure use of instructed amount
	of ULS per μg of aRNA
	Remedy: Ensure concentration of the
	labeling reaction is above 50 ng/µL
High labeling	Cause: Incorrect ratio of labeling reagent
density	to aRNA
	Remedy: Ensure use of instructed amount
	of ULS per μg of aRNA

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III. Array Hybridization and Detection

Problem	Possible Reasons and Suggestions
Background on	Cause: Too much sample added to
the slide	microarray
	Remedy: Reduce sample amount
	Cause: Insufficient blocking
	Remedy: Add more or alternative blockers
	to microarray

E. Appendix

I. Determination of RNA Quality

Measure A_{260} and A_{280} nm using a spectrophotometer and calculate $OD_{260/280}$. For good quality RNA this value should be between 1.9 and 2.1.

F. References

Van Gelder RN et al. (1990), Proc Natl Acad Sci USA 87: 1663–1667

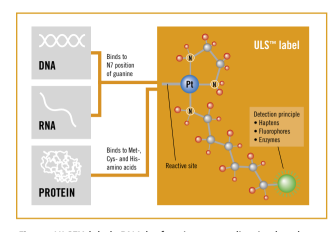


Figure. ULS^{TM} labels RNA by forming a coordinative bond on the N7 position of guanine.



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