

**GenoExplorer™ miRNA
Labeling Kit**
Catalog # 1301,1302

Version C
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User Manual

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Literature Citation

When describing a procedure for publication using these products, we would appreciate that you refer to them as the GenoExplorer™ miRNA Labeling Kit and the GenoExplorer™ miRNA Biochips Kit.

Patents and Trademarks

GenoExplorer is a trademark of GenoSensor. The GenoExplorer™ miRNA Labeling Kit and the GenoExplorer™ miRNA Biochips Kit are covered by patents pending.

Product Overview

The GenoExplorer™ miRNA Labeling Kit

- **Selective and Uniform Labeling**

GenoExplorer™'s labeling system employs a 5' end ligation approach. This offers greater selectivity in the labeling of RNA compared with more traditional 3' approaches since RNA molecules with capped 5' ends (*e.g.*, intact mRNA) will not be labeled. The ligation reaction also ensures that the labeling is uniform in that a single adaptor is attached to each RNA molecule with an uncapped 5' end. GenoExplorer™ currently offers a biotin label.

- **Simplified Sample Prep Protocol**

With GenoExplorer™'s more selective labeling system, RNA samples do not require the additional miRNA/small RNA isolation step as some labeling protocols recommend. Users can begin their labeling reaction with total RNA. It is important, however, to ensure that small RNA is retained during the initial RNA extraction procedure.

- **Streptavidin-Dye Staining**

The GenoExplorer™ full kit also includes a streptavidin-dye conjugate which is used for the post-hybridization dye staining step. This step is required for RNA labeled with biotin.

Kit Components and Storage Conditions

GenoExplorer™ miRNA Full Kit for 20 or 4 Reactions

| Components | 20 Reaction Amount | 4 Reaction Amount | Storage |
|---------------|------------------------|-------------------|---------|
| Buffer L | 140 µl | 28 µl | -20° C |
| Enzyme L | 10 µl | 2 µl | -20° C |
| SA-S Dye | 20 µl | 5 µl | -20° C |
| Buffer S | 2 vials of 1.4 ml each | 0.56 ml | -20° C |
| User's Manual | Web download | Web download | |

Shipping and Storage

GenoExplorer™ miRNA Labeling Kit reagents are shipped on dry ice. Components should be stored at temperatures shown in the above table. At proper storage conditions, components are stable for 1 year from the date received. Expiration dates are also noted on product labels.

Safety Warnings and Precautions

For research use only. Not recommended or intended for the diagnosis of disease in humans or animals. Do not use internally or externally in humans or animals. Consider all chemicals as potentially hazardous. Only persons trained in laboratory techniques and familiar with the principles of good laboratory practice should handle these products. Wear suitable protective clothing such as laboratory overalls, safety glasses, and gloves. Exercise caution to avoid contact with skin or eyes: if contact should occur, wash immediately with water (Material Safety Data Sheet for products is available upon request).

Additional Required Materials

Recommended: GenoExplorer™ miRNA Biochips Kit
Recommended: GenoExplorer™ miRNA Probe Set
Recommended: GenoExplorer™ Reagents for Hybridization Assay
Total RNA containing the small RNA
RNase-free water
Adjustable pipettors
RNase-free tips
RNase-free polypropylene microcentrifuge tubes (0.2, 0.5 or 1.5 ml)
Graduated cylinder
Microcentrifuge
Incubator set at 37° C
Incubator or heating block set at 75°
Heating block at 95° C
Hybridization Station (optional), for automated hybridization (*e.g.*, Genomic Solutions, Tecan) **or**
Coverslips, for manual hybridization (*e.g.*, Erie LifterSlips™, Product # 22x30I-2-4374)
Slide Chamber, for manual hybridization (*e.g.*, Corning® Microarray Hybridization Chamber, Product #2551)
Incubator or water bath set at 42° C, for manual hybridization
Bottles for storing diluted Wash Buffers
Centrifuge or air blower
Microarray scanner (*e.g.*, Axon, Agilent, Parkard) and image processing software
Several wash containers appropriately sized to the number of chips being used

Related Products from GenoSensor

GenoExplorer™ microRNA Full Kit (Cat# 1101C – 1199C)
GenoExplorer™ microRNA Biochips Kit (Cat# 1201C – 1299C)
GenoExplorer™ microRNA Probe Set (Cat# 1401C – 1499C)
GenoExplorer™ Reagents for Hybridization Assay (Cat# 1500's)

GenoExplorer™ Labeling Protocol

General Description

The GenoExplorer™ miRNA Labeling Kit (patent pending) provides a direct end-labeling method. Biotin labels are ligated to the 5' ends of RNA molecules which do not contain a 5'-capped structure. These RNA molecules include rRNA, tRNAs, regulatory small RNAs such as microRNA, siRNA, snRNAs, and other RNA transcripts of yet unknown function. Due to labeling selectivity, this method has resulted in low false positive hybridization signals that are usually caused by mRNA, which are the most highly complex sequences of genome transcripts. This protocol uses directly isolated RNA without RNA target amplification, and ultimately reflects the cellular microRNA molar ratio, thus providing for the reliability of hybridization signals. The streptavidin-conjugated dye stain (SA-S Dye) and the staining buffer (Buffer S) reagents are also included in this kit. Their preparation and use in the post-hybridization staining procedure is described below in the hybridization assay section.

Handling RNA Samples

When working with RNA, always use proper microbiological aseptic techniques. Use RNase- and DNase-free reagents, water, glassware and plasticware. Use non-powdered gloves during all steps of sample labeling, chip hybridization, washing, detection, and scanning.

RNA Preparation

The GenoExplorer™ miRNA Labeling Kit (patent pending) provides an easy and quick way to label microRNAs and other small RNAs. Total RNA isolation (not provided) using traditional methods such as Trizol is recommended. Some commercial kits can be used. Users should be aware of the harvest efficiency for small RNAs when choosing them. Checking with manufacturers is highly recommended. A total RNA starting amount of 5 to 10 micrograms is recommended.

High quality and sufficient amounts of RNA samples is crucial for experiments with microarrays. RNA quality can be evaluated by visualizing the RNA on a gel, as well as by calculating the A_{260}/A_{280} ratio. On a denaturing gel (or on an ordinary agarose gel in denaturing buffer) the RNA should appear as two bright distinct bands that represent the 28S and 18S ribosomal species. The 28S band should be brighter than the 18S band. Tailing of these major bands down the gel, or a background smear behind these bands that gets heavier at lower molecular weights can indicate degradation of the RNA. Degraded RNA will produce high background and low signal intensity microarray results.

miRNA Labeling Procedure

This procedure is used to attach a biotin label to the 5' ends of RNA molecules.

1. Place Buffer L on ice and thaw for 15-20 minutes. Check for any precipitate. If necessary, warm the solution to 37°C and agitate to dissolve the precipitate completely.
Note: Aliquot is recommended to minimize thaw/freeze cycles
2. Mix Buffer L by vortexing followed by brief centrifugation.
3. Combine reagents according to Table 1 below for a single reaction. Reagents should be combined in an RNAase-free microcentrifuge tube and all reagents should be kept on ice during set up of the reaction. For high accuracy, pipet the viscous Buffer L slowly.

Table 1: miRNA Labeling Reaction Mix

| Reagents | Volume (µl) |
|-------------------|-------------|
| Buffer L | 7.0 |
| RNA (2.5-10µg) | Adjustable |
| RNAase-free water | Adjustable |
| Enzyme L | 0.5 |
| Total | 20 |

4. Mix reaction thoroughly by pipetting up and down several times.
5. Incubate at 20° C for 3 hours.
6. Incubate at 75° C for 20 min to inactivate the enzyme.
7. Store on ice until ready for the hybridization step. The labeled sample can also be stored at -70° C and used later.

GenoExplorer™ Labeled miRNA Hybridization Assay Guidelines

The GenoExplorer™ miRNA Labeling Kit has been optimized for microarray experiments using GenoExplorer™ microRNA chips and assay reagents. If microarrays other than GenoExplorer™ microRNA chips are used, consult the optimized protocols established by the microarray/probe set provider. However, it is critical to also include the post-hybridization staining step. Below are general guidelines that can be used for running the hybridization assay using GenoExplorer™ miRNA labeled samples. For a detailed protocol on running GenoExplorer™ labeled miRNA on GenoExplorer™ BioChips, please consult the user manual for the GenoExplorer™ full kit.

1. Prepare labeled RNA in hybridization buffer following the optimized protocol established by microarray/probe set provider.
2. Incubate the sample at 90-95° C for 5 min to denature.
3. Cool sample by spinning down briefly.
4. Hybridize samples on microarrays following the optimized protocols established by the microarray and/or probe set provider.
5. After completion of the hybridization, slides should be washed and dried according to optimized protocols established by the microarray and/or probe set provider. If hybridization is performed in an automated hybridization station, the drying step is probably not required.
6. Within 15 minutes of beginning the post-hybridization staining, the SA-S dye should be diluted 250-fold into Buffer S. For a single sample this dilution should be performed by adding 1 µl of the SA-S to 249 µl of buffer S followed by gentle mixing and brief centrifugation. Keep protected from light to minimize photobleaching of dye.
7. Load the 1:250 diluted SA-S dye stain solution on to the array and incubate at 25°C for 30 min. Shield the slide from light to minimize effects of photobleaching of the dye.
8. After completion of the post-hybridization staining step, slides should be washed and dried according to the optimized protocols established by the microarray and/or probe set provider.

Scanning, Detection and Image Analysis

Consult the scanner manufacturer instructions for laser and PMT settings. The SA-S dye stain signal can be detected using the scanner's 635 nm wavelength (Cy5) channel. Consult the .gal file provided by the microarray manufacturer for further data analysis.

Appendix

Troubleshooting Guide

Poor Hybridization Signal

| | |
|--|---|
| Poor quality RNA samples | <ul style="list-style-type: none"> • Use higher quality RNA samples • Use proper laboratory techniques when handling RNA samples • Label and hybridize more sample to microarray |
| Suboptimal amount of sample applied to microarray | <ul style="list-style-type: none"> • Label and hybridize more sample to microarray |
| Improper detection strategy | <ul style="list-style-type: none"> • Verify that detection instrumentation is compatible with detection reagents • Adjust detection settings |
| Signal lost by exposure to light, environmental conditions | <ul style="list-style-type: none"> • Minimize exposure to light |
| Poor biotin/SA-S dye detection | <ul style="list-style-type: none"> • Ensure that the samples have been purified and quantified properly before labeling • Ensure that the kit components have been stored properly • Ensure that the detection solution was prepared and stored properly |
| Hybridization signal “stripped” from microarray | <ul style="list-style-type: none"> • Ensure correct temperatures are used for each of the process steps. |
| Suboptimal hybridization time | <ul style="list-style-type: none"> • Extend hybridization time |

High Background

| | |
|---|--|
| Excess detection reagents remaining on microarray | <ul style="list-style-type: none">• Do not allow slides to dry out during hybridization protocol• Do not touch microarray directly or forcibly remove coverslip at any time |
| Excess sample applied to microarray | <ul style="list-style-type: none">• Quantify the amount of labeled sample and use less in hybridization |
| Ink or marker used to identify microarray | <ul style="list-style-type: none">• Avoid using markers or ink to identify slide; use a diamond scribe pen |

Dust

| | |
|--------------------|--|
| Dust on microarray | <ul style="list-style-type: none">• Avoid dust from environment• Minimize exposure of the microarray to the air• Use filtered distilled water for final wash |
|--------------------|--|

Technical Service

For more information or technical assistance, please call, write, fax, or email.

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Limited Warranty

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