

New
miRNA Probes

miRNA Product Catalog

2020

(US)

Precision Medicine Research

- Cancer of Unknown Primary (CUP)
- Poorly Differentiated Tumors
- Undifferentiated Tumors
- Grading and Staging of Cancer



All-in-One

Precision Medicine

Dear Customer,

We are pleased to present the BioGenex miRNA Product Catalog for 2018. As a vertically integrated company, we develop, manufacture and market highly innovative and fully automated systems for cancer diagnosis, prognosis and therapy selection.

Xmatrix® systems redefine complete automation for the molecular pathology laboratory and standardize the protocol from baking through final cover-slipping in three simple steps - Load, Click and View. Compared to any other system on the market, Xmatrix® systems offer clean intense stain(s), automate more assay steps, and enable automation of technologies for the future molecular pathology laboratory.

- Xmatrix® ELITE integrates All-in-One staining of IHC, ISH, miRNA, ISH, special stains and beyond
- Xmatrix® Infinity is a high-performance staining platform for life sciences and translational research
- Xmatrix® ULTRA Dx is the next-generation system with new features such as Auto Drain, Auto DAB mixing and with new technologies
- Xmatrix® ULTRA Rx is the next-generation system with new features and technologies for life sciences and translation research
- NanoMtrx® 300 is a fully-automated, 30-slide benchtop compact system with micro-chamber® for IHC, ISH
- NanoMtrx® 100 is a fully-automated, 10-slide benchtop compact system with micro-chamber® for IHC and ISH
- Xmatrix® NANO VIP is a ten-slide automated system specifically designed for FISH
- Xmatrix® MINI enables *in situ* PCR and nucleic acid hybridization with tools for building micro-chamber

miRNA-guided diagnostics is a powerful molecular approach for evaluating clinical samples through miRNA detection and/or visualization. To date, this approach has been successfully used to diagnose, manage, and/or monitor a wide range of neoplastic and non-neoplastic diseases.

We offer a full selection of high quality fluorescent labeled human miRNA detection probes, ideal for sensitive and specific *in situ* hybridization of miRNA

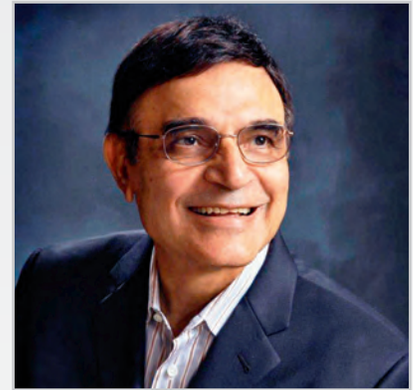
To maintain our tradition of offering superior solutions for the emerging needs of your laboratory, apart from miRNA, we offer a broad range of molecular pathology products for IHC, ISH, multiplex and special staining of tissues including 400+ primary antibodies, molecular probes, detection systems, and ancillaries. These are offered for standardized, reliable and consistent results to support the needs of molecular pathology laboratories of today, tomorrow and beyond.

BioGenex is committed to the core values of innovation, reliability, productivity, quality, superior after-sales support and service for complete customer satisfaction. These values are represented by our company's colors that stand for "energy and innovation" (orange) and "reliability" (blue). We unconditionally guarantee all of our products and services.

I invite you to learn more about our exciting products and future development through this catalog and our new website at www.biogenex.com. Should you have any suggestions for improving our products and services, I encourage you to write me directly at k.kalra@biogenex.com.

Give us an opportunity and experience the difference.

Warm Regards,
Krishan Kalra, Ph.D.
CEO



“ To become a global molecular medicine company providing affordable solutions for life science research and personalized medicine ”

Dr. Krishan Kalra

- Innovation
- Quality
- Service
- Reliability
- Productivity

Table of Contents

Overview.....	v
Ordering Information.....	vi
General Information.....	vii
Additional Information	vii

MicroRNA Probes

New microRNA Probes.....	02
microRNA Probes.....	05
Hybridization Detection System.....	44
Substrates and Chromogens.....	45

Automated Systems

Automated Platforms for Molecular Pathology.....	47
Xmatrix® ELITE.....	48
Xmatrix® ULTRA Dx.....	49
Xmatrix® Infinity.....	50
Xmatrix® ULTRA Rx.....	51
NanoMtrx® 300.....	52
NanoMtrx® 100.....	53
NanoVIP®.....	54
Xmatrix® MINI.....	55

Tissue Pre-treatment & Nucleic Acid Retrieval

EZ-DeWax™ Solution.....	57
Nucleic Acid Retrieval Method.....	57
Enzymes for Tissue Digestion.....	59
i500Plus™.....	58
EZ-Retriver®.....	58

Consumables and Ancillary Reagents

Microscope Slides & Coverslips.....	61
Pipette tips.....	62
Accessories	62
Buffers	63
Counterstain & Mounting Media.....	63

MicroRNA Tissue Control

Positive Control Slides and Barrier Slides.....	67
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General Terms and Conditions.....	73
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Overview

BioGenex celebrated its 36th anniversary serving the anatomic pathology market. We take great pride in providing premier service and support while bringing new and technologically advanced products to the market.

BioGenex provides a “Total Solution” for slide-based cell and tissue analysis. Our products include a wide variety of antibodies, highly sensitive detection kits, automated systems, probes and ancillary products. Our automated systems streamline operations in molecular and cellular pathology laboratories, providing effective tools for the detection and diagnosis of cancer and other diseases. BioGenex continues to innovate as evidenced by the launch of the Xmatrix® Staining System which provides complete automation “From Microtome to Microscope”.

We are committed to providing our customers and our distributors with flexible, innovative and cost-effective tools for clinical diagnostics, life science research and drug discovery.

Service

We value you and your business. We want our relationship to be one of total satisfaction. Our Technical Support Specialists provide fast troubleshooting advice and technical information and they are responsive to your individual needs. Just visit our website at www.biogenex.com, send an e-mail to support@biogenex.com or call toll free at 1-(800)-421-4149 from 7:00 AM to 4:00 PM (PST), Monday through Friday, with your request.

Quality

BioGenex is committed to excellence by providing high-quality products. We offer a broad range of products which are manufactured using state-of-the-art equipment in controlled environments. They are stringently tested to ensure that they meet or exceed functional, dimensional, and environmental requirements and are compliant with federal regulations. Our automated systems are designed for high-throughput at a low cost of ownership. They provide consistent quality results with ease-of-use and maximum flexibility for clinical diagnostics, life science research, and drug discovery markets.

Reliability

BioGenex products give consistent, reproducible and reliable results. Our automated systems are highly reliable and dependable, giving our customer peace of mind.

Innovation

BioGenex has a rich history of innovation in the field of Immunohistochemistry (IHC) and *In situ* Hybridization (ISH). BioGenex has a strong intellectual portfolio, consisting of several US and foreign-issued patents, in the areas of

- DNA labeling and amplification
- Antigen retrieval and deparaffinization
- Automation of tissue and cell sample preparation
- Automated IHC, and staining of nucleic acids
- Nucleic acid retrieval for tissues

Productivity

BioGenex has automated cell and tissue analysis to accelerate clinical diagnostics and drug discovery development. We have developed the total walk-away, industrial scale automated systems to streamline and standardize an array of processes for cell and tissue testing in IHC, ISH/CISH, FISH, and image analysis applications. We offer a “Total Solution” automating every aspect of the histology slide preparation “From Microtome to Microscope”. These technologies significantly increase laboratory operation productivity for clinical diagnostics, drug discovery and life sciences research applications by providing high-quality staining and imaging solutions.

Ordering Information

BioGenex Customer Service

Please call our Customer Service department from 07:00 A.M. to 04:00 P.M. (PST), Monday through Friday, to place an order or to inquire about an existing order.

Telephone (toll-free)	1-(800)-421-4149 (Option 1)
Fax	1-(510)-824-1490
Online Orders	www.biogenex.com
E-mail	customer.service@biogenex.com
Mail Orders	BioGenex Laboratories, Inc. Attention to: Customer Service 49026 Millmont Drive Fremont, CA 94538

Quote request can also be placed via our website.

To expedite the order process, please include the following information on your purchase order or correspondence:

- Purchase order number
- Customer number
- Name, phone and fax number of person ordering
- Shipping address (please do not use P.O. Box number)
- Billing address (if different from above)
- Name of product, catalog number, quantity, and price
- Special shipping instructions
- Credit card number and expiration date (for credit card payments)

International Orders

To place an order from outside the US, please contact your local BioGenex channel partner/distributor. For online orders please visit our website www.biogenex.com For countries where BioGenex does not have any channel partners/distributors, please e-mail us at internationalcs@biogenex.com

Opening a New BioGenex Account

First time orders paid by credit card (see under Payment) will be processed and shipped immediately. For other payment methods please accept a delivery time of up to five business days for credit verification purposes.

Credit Terms

Net 30 days in U.S. Dollars, upon approval. Overdue accounts are subject to a finance charge of 1.5% per month (18% per annum).

Confirming Orders

To avoid duplication of your shipment, please mark boldly "confirming order - please do not ship" on your order.

Pricing

All prices are quoted in U.S. dollars, exclusive of state and county sales tax, where applicable. Prices are valid only for shipments within U.S. and are subject to change without notice. Please inquire about our standing order and quantity discount policies.

Shipping

Shipping and handling charges are prepaid and added to the invoice. They vary with the destination, weight and content, and are available upon request at order entry and are indicated on the invoice. Reagent orders received by 2:00 P.M. (PST), Monday through Thursday, will generally be Expedited Shipping for Next Day Delivery. Early A.M. and Saturday delivery are available upon request.

Payment

All payments must be made in U.S. dollars. The following methods of payment are accepted:

- Bank transfer (see invoice for instructions)
- Check, drawn on a U.S. bank, made payable to: "BioGenex Laboratories, Inc."
- MasterCard®
- Visa®
- American Express®

Return Policy

Reagents are covered by the following Total Quality Assurance policy which states:

If you are not completely satisfied with the quality of our reagents, you may return them to us along with poor stained slides and filled RMA form for a refund or replacement, at our option.

BioGenex's liability is limited to a refund or replacement, at our option.

Please obtain a Return Material Authorization (RMA) number from Customer Service prior to the return of a product.

Returns, which are caused by unsatisfactory product performance, must be made within 30 days of delivery and will be subject to a 30% restocking fee.

Returns or replacements cannot be accommodated for expired products.

As BioGenex is an ISO13485 and USFDA compliant IVD manufacture, we can't accept returned products without return material authorization, RMA. All returned products without RMA will be trashed.

General Information

Web Site

For the latest information on new product releases listed pricing, special offers and for placing an online order, please visit our new website, www.biogenex.com

Customer Support

Our technical support and customer service specialists are ready to provide fast and detailed Information for your questions and needs. Please call our toll-free number to reach us.

Customer Service USA

Tel: 1-(800)-421-4149 (Option 1)

Fax: 1-(510)-824-1490

E-mail: customer.service@biogenex.com

Technical Support USA

Tel: 1-(800)-421-4149 (Option 2)

Fax: 1-(510)-824-1490

E-mail: support@biogenex.com

Website: www.biogenex.com

Corporate Office

BioGenex Laboratories, Inc.

49026 Milmont Drive

Fremont, CA 94538

Tel: 1-(800)-421-4149

Fax: 1-(510)-824-1490

Corporate Business

For general business matters not related to product orders or inquiries, please call us at 1-(800)-421-4149 or fax your correspondence to our main corporate business fax: 1-(510) 824-1490.

Trademarks

The following are trademarks of BioGenex Laboratories, Inc. USA

BioGenex®	EZ-AR™
EZ-Retriever®	MultiLink®
EZ-DeWax™	GenoMx®
i500 Plus™	Xmatrx®
Power Block™	XMOUNT™
AccuSlide®	XViz™
OptiPlus™	Super Mount®
InSite®	XISH™
XWash™	eXACT™

Additional Information

Nationwide Training Workshops

As a service to our customers, BioGenex has developed lectures and workshops on the full range of Immunohistochemistry and *in situ* Hybridization techniques. Please call our Technical Support Department or Regional Account Executive for more information on how you can participate in our educational workshops. Topics include the following:

- Basic Immunohistochemistry
- Cancer Panels
- Microwave-Based Antigen Retrieval
- ER/PR Immunostaining
- Troubleshooting
- Automation
- *in situ* Hybridization
- Double Staining
- Multiplexing and Co-detection of Protein and Nucleic Acid Biomarkers

We raise awareness of miRNA detection issues and recommend research directions to help pathologists integrate miRNA testing into clinical decision-making.

Free Technical Literature

In addition to the educational brochures produced by BioGenex, we offer other technically useful information to the histopathology specialists on our website, www.biogenex.com where you can download our data sheet, product catalog or relevant presentation that may accompany each product assay protocols, kit instruction manuals and conference posters. Please call our Technical support department to request specific items or to add your name to our mailing list.

Technology Partnering Opportunities

We are always interested in licensing innovative technology that will be useful to our customers. If you are a researcher and have new antibody clones or other new diagnostic technologies please think of BioGenex as a potential partner in marketing your inventions and discoveries. We have the scientific expertise and marketing experience necessary for the successful commercialization of your technical achievements. BioGenex has an active Research and Development program fully staffed with PhD and MD professionals who are experienced in immunopathology, protein chemistry, and molecular biology. For more information on technology transfer opportunities, please contact us at customer.service@biogenex.com



MicroRNA Probes



New

MicroRNA Probes

MicroRNAs (miRNAs) are endogenous, non-coding RNAs known to regulate gene expression by translational repression or RNA cleavage. Since miRNA has been observed to deregulate during progression of different cancer stages from normal to malignant and metastasis, the expression profile as a result of this deregulation can be exploited as a potential biomarker for cancer characterization.

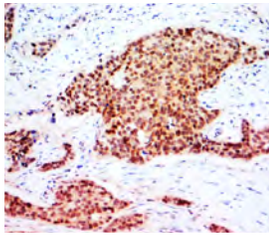
Automated and manual protocols

- Optimized for automated ISH staining by Xmatrx[®] ELITE
- Ready-to-use(RTU) reagents for FFPE tissues
- ISH Detection System and ancillaries

Highly Specific and Sensitive Probes

- Proprietary technology for clean intense stains
- *in situ* context of tissue morphology
- Positive control tissue slides

Hsa-miR-299-3p

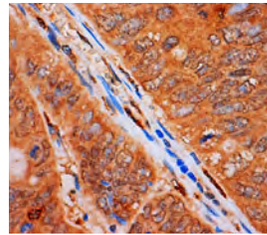


Hsa-miR-299-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM299-3P-100 (ASR*)
 Specificity: miR-299-3p
 Recommended Barrier: FB-HM299-3P
 Control:

miRNA-299-3p has been reported to modulate replicative senescence in endothelial cells and may be the target for potential clinical use to decrease invasiveness of breast cancer. The expression level of miRNA-299-3p identified statistically significant difference in melanoma samples. The fluorescinated hsa-miR-299-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-556

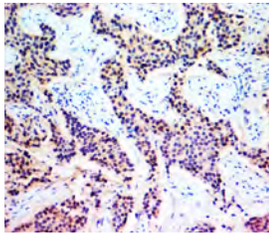


Hsa-miR-556 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM556-100 (ASR*)
 Specificity: miR-556
 Recommended Barrier: FB-HM556
 Control:

miRNA-556 is a novel marker of human colorectal cancer cells. The expression level of miRNA-556 is important for short disease free survival and overall survival in stage II colon cancer which may suggest its important role as a valuable aid to therapeutic decision marking in colorectal cancer (CRC) disease progress. The fluorescinated hsa-miR-556 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-301a-3p

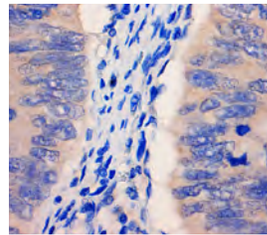


Hsa-miR-301a-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM301A-3P-100 (ASR*)
 Specificity: miR-301a
 Recommended Barrier: FB-HM301A-3P
 Control:

miRNA-301a-3p is down-regulated in pancreatic cancer cells and contributes to development of estrogen independence to lead to the invasion of breast cancer. The expression level of miRNA-301a-3p identified statistically significant difference in melanoma. The fluorescinated hsa-miR-301a-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-656-3p

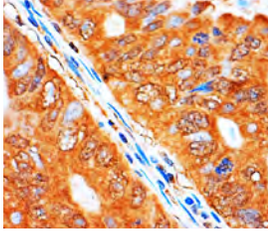


Hsa-miR-656-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM656-3P-100 (ASR*)
 Specificity: miR-656-3p
 Recommended Barrier: FB-HM656-3P
 Control:

miRNA-656-3p has been reported to express in colon tissues. The expression level of miRNA-656-3p identified high-risk patients of TNM-stage II colon cancer which may suggest its important role as a valuable aid to classify patients in TNM-stage II cancer with high risk of recurrence. The fluorescinated hsa-miR-656-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

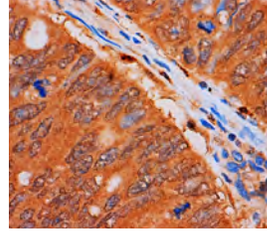
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-671-3p

Hsa-miR-671-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM671-3P-100 (ASR*)
 Specificity: miR-671-3p
 Recommended Barrier: FB-HM671-3P
 Control:

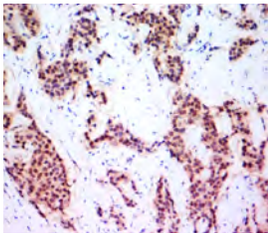
miRNA-671-3p is reported to express in colon tissues and functions as a tumor suppressor in breast cancer by influencing the Wnt signaling pathway. The expression level of miRNA-671-3p identified high-risk patients of TNM-stage II colon cancer. The fluorescinated hsa-miR-671-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-5010-3p

Hsa-miR-5010-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM5010-3P-100 (ASR*)
 Specificity: miR-5010-3p
 Recommended Barrier: FB-HM5010-3P
 Control:

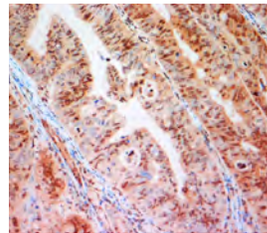
miRNA-5010-3p is reported to be dysregulated in colon adenomas. The expression level of miRNA-5010-3p identified high-risk patients of TNM-stage II colon cancer which may suggest its important role as a valuable aid to classify patients in TNM-stage II cancer with high risk of recurrence. The fluorescinated hsa-miR-5010-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-1537

Hsa-miR-1537 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM1537-100 (ASR*)
 Specificity: miR-1537
 Recommended Barrier: FB-HM1537
 Control:

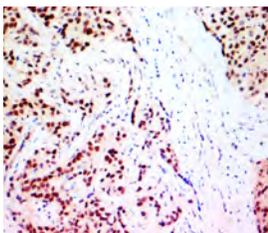
miRNA-1537 is reported to be up-regulated in melanoma and related with Her2 subtype breast cancer patients survival rate. The expression level of miRNA-1537 identified statistically significant difference in melanoma samples which may suggest its important role as a diagnostic biomarker and improve the precision and accuracy of melanoma detection and monitoring. The fluorescinated hsa-miR-1537 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-5100

Hsa-miR-5100 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM5100-100 (ASR*)
 Specificity: miR-5100
 Recommended Barrier: FB-HM5100
 Control:

The expression level of miRNA-5100 is increased in non-small-cell lung cancer and pancreatic cancer. miRNA-5100 identifies high-risk patients of TNM-stage II colon cancer which may suggest its important role as a valuable aid to classify patients in TNM-stage II cancer with high risk of recurrence. The fluorescinated hsa-miR-5100 probe is designed to localize this miRNA in FFPE tissue by *in situ*.

Hsa-miR-4787-3p

Hsa-miR-4787-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM4787-3P-100 (ASR*)
 Specificity: miR-4787-3p
 Recommended Barrier: FB-HM4787-3P
 Control:

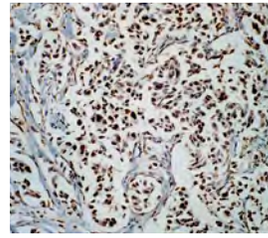
miRNA-4787-3p is a potential important marker for breast cancer. The expression level of miRNA-4787-3p identified statistically significant difference in melanoma samples which may suggest its important role as a diagnostic marker for melanoma. The fluorescinated hsa-miR-4787-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-1

Hsa-miR-1 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): PR026-100 (ASR*)
 Specificity: miR-1
 Recommended Barrier: FB-HM001
 Control:

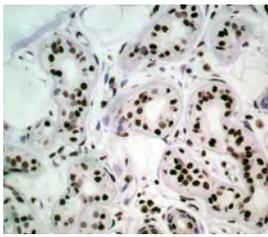
miR-1 plays a key role in the development and differentiation of smooth and skeletal muscles. The fluorescinated hsa-miR-1 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-let-7c

Hsa-miR-let-7c detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM007C-100 (ASR*)
 Specificity: let-7c
 Recommended Barrier: FB-HM007C
 Control:

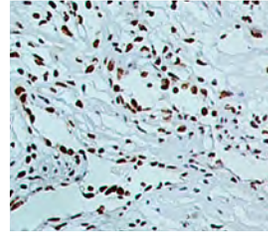
Data suggest that miR-let-7c suppresses androgen receptor expression and activity via regulation of myc expression. The fluorescinated hsa-miR-let-7c probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-let-7a

Hsa-miR-let-7a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM007A-100 (ASR*)
 Specificity: let-7a
 Recommended Barrier: FB-HM007A
 Control:

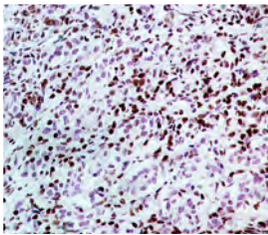
miR-let-7a has been shown to directly alter cell cycle progression and proinflammatory cytokine production. The fluorescinated hsa-miR-let-7a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-let-7d

Hsa-miR-let-7d detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM007D-100 (ASR*)
 Specificity: let-7d
 Recommended Barrier: FB-HM007D
 Control:

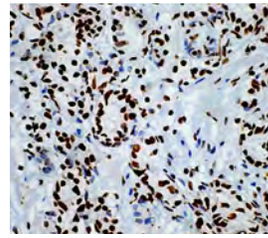
Let-7 family gene was first discovered in the nematode as a key developmental regulator. The expression of let-7 family has been reported to be lower in multiple tumor tissue than in normal tissue. The fluorescinated hsa-miR-let-7d probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-let-7b

Hsa-miR-let-7b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM007B-100 (ASR*)
 Specificity: let-7b
 Recommended Barrier: FB-HM007B
 Control:

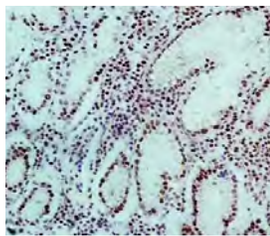
Let-7 family gene was first discovered in the nematode as a key developmental regulator. The expression of let-7 family has been reported to be lower in multiple tumor tissue than in normal tissue. The fluorescinated hsa-miR-let-7b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-let-7e

Hsa-miR-let-7e detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM007E-100 (ASR*)
 Specificity: let-7e
 Recommended Barrier: FB-HM007E
 Control:

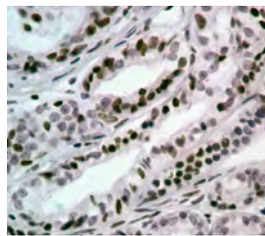
miR-let-7e plays a pivotal role in stem cell differentiation and its loss results in reversion of embryogenesis and dedifferentiation. The fluorescinated hsa-miR-let-7e probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-let-7g

Hsa-miR-let-7g detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM007G-100 (ASR*)
 Specificity: let-7g
 Recommended Barrier: FB-HM007G
 Control:

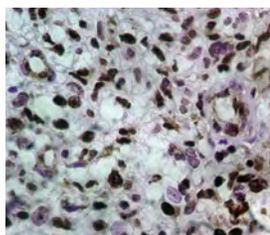
Let-7 family gene was first discovered in the nematode as a key developmental regulator. The expression of let-7 family has been reported to be lower in multiple tumor tissue than in normal tissue. The fluorescinated hsa-miR-let-7g probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-15a

Hsa-miR-15a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM015A-100 (ASR*)
 Specificity: miR-15a
 Recommended Barrier: FB-HM015A
 Control:

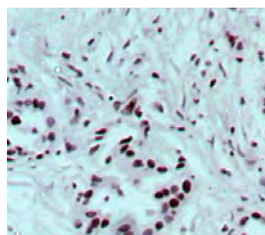
miR-15a might function as a tumor suppressor in the disease, and its expression has been reported to be lower in multiple tumor tissue than in normal tissue. The fluorescinated hsa-miR-15a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-9

Hsa-miR-9 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM009-100 (ASR*)
 Specificity: miR-9
 Recommended Barrier: FB-HM009
 Control:

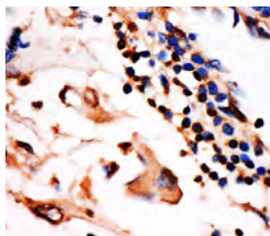
A series of miR-9 targets, such as nuclear factor (NF)- κ B1, caudal type homeobox 2 (CDX2), chromobox protein homolog 7 (CBX7), and methylenetetrahydrofolate cyclohydrolase (MTHFD2), were associated with tumorigenesis. The fluorescinated hsa-miR-9 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-15b

Hsa-miR-15b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM015B-100 (ASR*)
 Specificity: miR-15b
 Recommended Barrier: FB-HM015B
 Control:

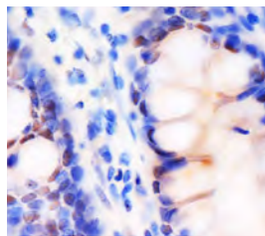
MiR-15b plays an important role in DNA damage response and repair mechanisms, thus protects cells from genotoxic stress. Recently, it has been reported that the expression of miR-15b may be altered following exposure to various genotoxic stressors including radiation, hydrogen peroxide and etoposide. The fluorescinated hsa-miR-15b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-10b

Hsa-miR-10b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM010B-100 (ASR*)
 Specificity: miR-10b
 Recommended Barrier: FB-HM010B
 Control:

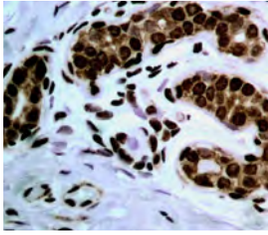
miR-10b has been identified as a target gene of transforming growth factor- β (TGF- β 1) which is a multifunctional cytokine that induces EMT in multiple cell types. The fluorescinated hsa-miR-10b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-16

Hsa-miR-16 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM16-100
 Specificity: miR-16
 Recommended Barrier: FB-HM16
 Control:

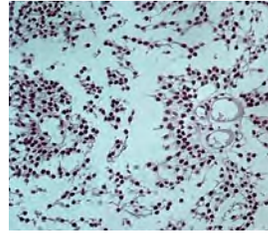
A recent meta-analysis showed that miR-16 family members have a relatively high value as promising biomarkers in diagnosing cancers. Another meta-analysis showed that the pooled sensitivity and specificity of miR-16 were 90% and 79.3% in diagnosing gastric cancer, which indicated that the measurement of elevated miR-16 levels in plasma could be a potential marker for gastric cancer. The fluorescinated hsa-miR-16 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-17

Hsa-miR-17 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM017-100 (ASR*)
 Specificity: miR-17
 Recommended Barrier: FB-HM017
 Control:

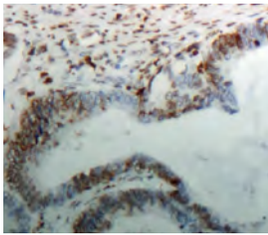
miR-17-92 is a polycistronic microRNA cluster that contains multiple microRNA components, each of which has a potential to regulate hundreds of target mRNAs. The fluorescinated hsa-miR-17 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-19a

Hsa-miR-19a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM019A-100 (ASR*)
 Specificity: miR-19a
 Recommended Barrier: FB-HM019A
 Control:

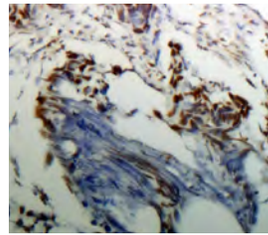
The suppressor of cytokine signaling 1 (SOCS1) is a novel target of miR-19a and miR-19a expression is inversely correlated with SOCS1 expression in tumor cells and tissues. The fluorescinated hsa-miR-19a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-17-3p

Hsa-miR-17-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM017-3P-100 (ASR*)
 Specificity: miR-17-3p
 Recommended Barrier: FB-HM017-3P
 Control:

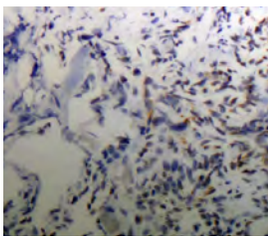
Tumor cell proliferation, colony formation, cell survival and invasion. Both miR-17-5p and miR-17-3p repressed TIMP metalloproteinase inhibitor 3 (TIMP3) expression. The fluorescinated hsa-miR-17-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-19b-3p

Hsa-miR-19b-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM019B-3P-100 (ASR*)
 Specificity: miR-19b-3p
 Recommended Barrier: FB-HM019B-3P
 Control:

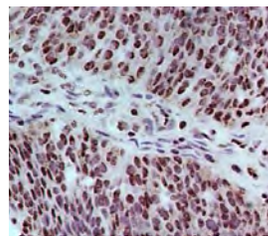
miR-19b-3p was identified to be the novel potential plasma miRNA candidate to detect some tumors. The fluorescinated hsa-miR-19a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-18a

Hsa-miR-18a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM018A-100 (ASR*)
 Specificity: miR-18a
 Recommended Barrier: FB-HM018A
 Control:

Hsa-miR-18a is located in the miR-17-92 cluster and reported to be highly expressed in multiple tumor tissues. The fluorescinated hsa-miR-18a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

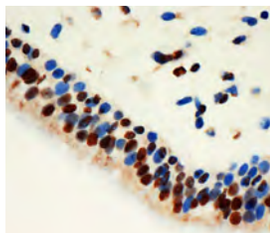
Hsa-miR-20a

Hsa-miR-20a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM020A-100 (ASR*)
 Specificity: miR-20a
 Recommended Barrier: FB-HM020A
 Control:

miR-20a was up-regulated in some tumor tissue and it contributed to tumor progression. The fluorescinated hsa-miR-20a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

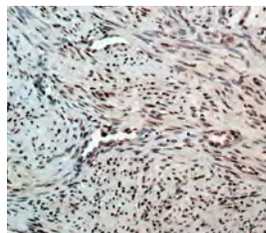
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-21

Hsa-miR-21 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM021-100 (ASR*)
 Specificity: miR-21
 Recommended Barrier: FB-HM021
 Control:

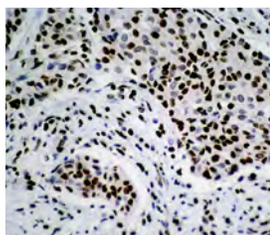
miR-21 is shown to involve in diverse biologic processes such as cell differentiation, proliferation, and apoptosis, presumably by modulating target proteins. The target genes of miR-21 include PTEN and programmed cell death 4 (PDCD4). The fluorescinated hsa-miR-21 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-23a

Hsa-miR-23a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM23A-100 (ASR*)
 Specificity: miR-23a
 Recommended Barrier: FB-HM23A
 Control:

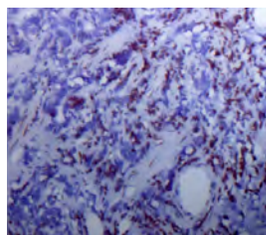
miR-23a is a miRNA cluster located in chromosome 19p13.12, which can function as an oncogene in several human malignancies. The fluorescinated hsa-miR-23a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-21-3p

Hsa-miR-21-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM021-3p-100 (ASR*)
 Specificity: miR-21-3p
 Recommended Barrier: FB-HM021-3P
 Control:

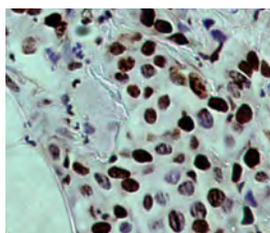
miR-21-3p has been shown to directly reduce the expression of two methionine adenosyltransferase genes by targeting their 3'-UTRs. The overexpression of miR-21-3p increases intracellular S-adenosylmethionine contents. The fluorescinated hsa-miR-21-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-23b

Hsa-miR-23b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM023B-100 (ASR*)
 Specificity: miR-23b
 Recommended Barrier: FB-HM023B
 Control:

miR-23b dysregulation may be associated with tumor progression. The fluorescinated hsa-miR-23b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-22

Hsa-miR-22 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM022-100 (ASR*)
 Specificity: miR-22
 Recommended Barrier: FB-HM022
 Control:

miR-22 sequence locates on the short arm of chromosome 17, in a minimal loss of heterozygosity region. miR-22 was found to be dysregulated in tumor tissue. The fluorescinated hsa-miR-22 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

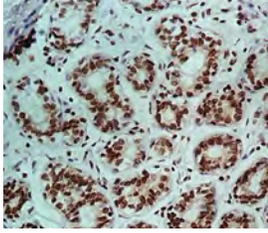
Hsa-miR-24-2

Hsa-miR-24-2 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM24-2-100 (ASR*)
 Specificity: miR-24-2
 Recommended Barrier: FB-HM24-2
 Control:

miR-24 governs cellular development and proliferation, acting as a tumor suppressor or oncogene in a cell type-specific manner. Multiple studies have demonstrated that miR-24 regulates the cell cycle both positively and negatively. The fluorescinated hsa-miR-24-2 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-24-3p

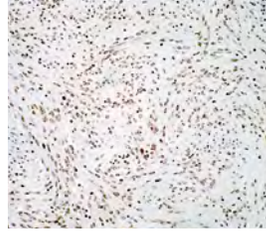


Hsa-miR-24-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM024-3P-100 (ASR*)
 Specificity: miR-24-3p
 Recommended Barrier Control: FB-HM024-3P

Recently, it has been shown that overexpression of miR-24-3p could alter T-cell proliferation and affect cellular gene expression through downregulation of mitogen activated protein kinase (MAPK) pathway. Thus imply the clinical relevance and prognostic value of tumor-derived exosomal miR-24-3p in T-cell dysfunction. The fluorescinated hsa-miR-24-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-27a

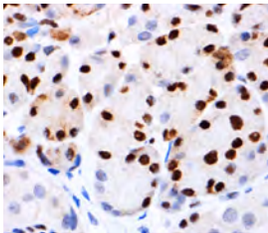


Hsa-miR-27a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM027A-100 (ASR*)
 Specificity: miR-27a
 Recommended Barrier Control: FB-HM027A

Data suggested that miR-27a suppresses ZBTB10/RINZF expression, and this novel zinc finger protein inhibits Sp1-dependent activation of the gastrin gene promoter. The fluorescinated hsa-miR-27a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-25

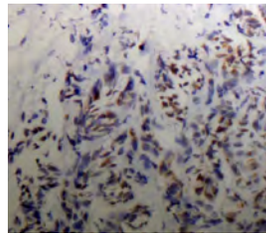


Hsa-miR-25 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM25-100 (ASR*)
 Specificity: miR-25
 Recommended Barrier Control: FB-HM25

miR-25 levels increase in human heart failure, and treatment with an anti-sense RNA molecule was recently reported to halt disease progression and improves cardiac function. The fluorescinated hsa-miR-25 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-27b

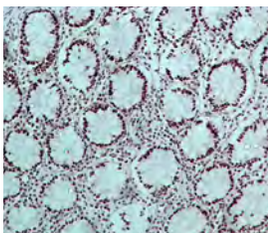


Hsa-miR-27b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): Hsa-miR-27b (ASR*)
 Specificity: miR-27b
 Recommended Barrier Control: FB-HM027B

miR-27b has been identified as an oncogenic microRNA and is highly expressed in tumor cells. Inhibition of miR-27 by antisense molecules decreases cell proliferation. The fluorescinated hsa-miR-27b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-26a

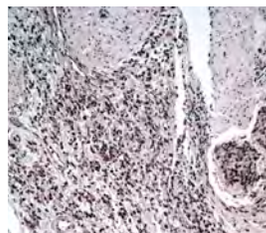


Hsa-miR-26a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM026A-100 (ASR*)
 Specificity: miR-26a
 Recommended Barrier Control: FB-HM026A

miR-26 expression is induced in response to hypoxia and upregulated during smooth muscle cell (SMC) differentiation and neurogenesis. Moreover, miR-26 is consistently down-regulated in a wide range of malignant tumors. The fluorescinated hsa-miR-26a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-28-3p

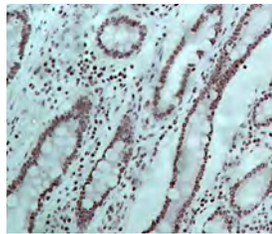


Hsa-miR-28-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM028-3P-100 (ASR*)
 Specificity: miR-28-3p
 Recommended Barrier Control: FB-HM028-3P

miR-28-3p is down-regulated in tumor samples compared with normal samples. miR-28-3p increase tumor cell migration and invasion in vitro. The fluorescinated hsa-miR-28-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

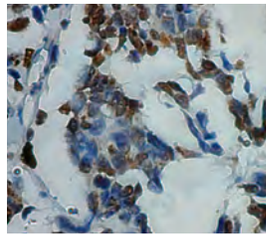
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-28-5p

Hsa-miR-28-5p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM028-5P-100 (ASR*)
 Specificity: miR-28-5p
 Recommended Barrier: FB-HM028-5P
 Control:

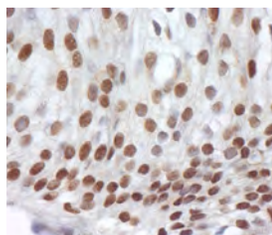
miR-28-5p is down-regulated in tumor samples compared with normal tissue. miR-28-5p increase tumor cell migration and invasion in vitro. The fluorescinated hsa-miR-28-5p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization

Hsa-miR-29c

Hsa-miR-29c detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM029C-100 (ASR*)
 Specificity: miR-29c
 Recommended Barrier: FB-HM29C
 Control:

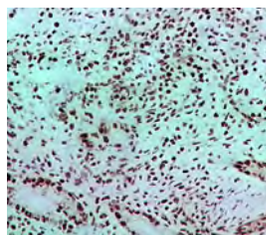
Mir-29 microRNA families are involved in regulation of various types of tumors. mir-29 was shown to play an inhibitory role in tumorigenesis. Many mammalian genomes encode four closely related miR-29 precursors that are transcribed in two transcriptional units. The fluorescinated hsa-miR-29c probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-29a

Hsa-miR-29a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM29A-100 (ASR*)
 Specificity: miR-29a
 Recommended Barrier: FB-HM29A
 Control:

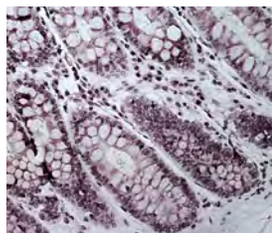
Ectopic expression of miR-29a in mouse hematopoietic stem cells (HSC) promoted self-renewal of myeloid progenitors, leading to a myeloproliferative disorder. The fluorescinated hsa-miR-29a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-30b

Hsa-miR-30b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM030B-100 (ASR*)
 Specificity: miR-30b
 Recommended Barrier: FB-HM030B
 Control:

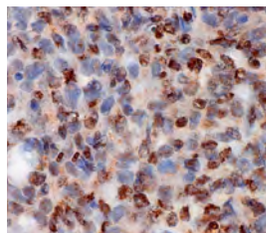
miR-30b promoted the metastatic behavior of tumor cells by directly targeting the GalNAc transferase GALNT7, which resulted in increased synthesis of the immunosuppressive cytokine IL-10, and reduced immune cell activation and recruitment. The fluorescinated hsa-miR-30b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-29b-3p

Hsa-miR-29b-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM29B-3P-100 (ASR*)
 Specificity: miR-29b-3p
 Recommended Barrier: FB-HM29B-3P
 Control:

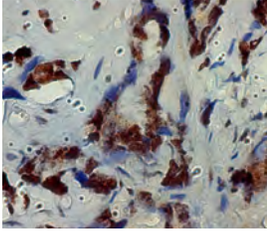
miR-29b-3p was found to be dysregulated in several tumor tissues. The fluorescinated hsa-miR-29b-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-30c

Hsa-miR-30c detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM030C-100 (ASR*)
 Specificity: miR-30c
 Recommended Barrier: FB-HM030C
 Control:

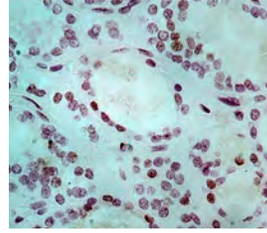
miR-30c involved in regulating a number of tumor associated genes. It has been shown that the integrin ITGB3 and the ubiquitin conjugating E2 enzyme (UBC9) are downregulated by miR-30. It has also been suggested that the TP53 protein may be a target of miR-30c and miR-30e. Members of the miR-30 family have been found to be highly expressed in heart cells. The fluorescinated hsa-miR-30c probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-30e

Hsa-miR-30e detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM030E-100 (ASR*)
 Specificity: miR-30e
 Recommended Barrier: FB-HM030E
 Control:

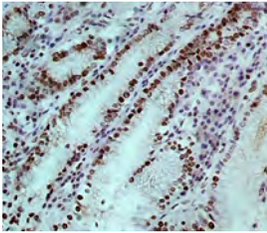
miR-30e involved in regulating a number of tumor associated genes. It has been shown that the integrin ITGB3 and the ubiquitin conjugating E2 enzyme (UBC9) are downregulated by miR-30. It has also been suggested that the TP53 protein may be a target of miR-30c and miR-30e. Members of the miR-30 family have been found to be highly expressed in heart cells. The fluorescinated hsa-miR-30e probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-34c

Hsa-miR-34c detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM34C-100 (ASR*)
 Specificity: miR-34c
 Recommended Barrier: FB-HM34C
 Control:

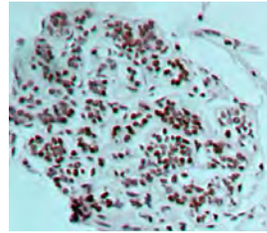
miR-34c has also been reported to be downregulated in several tumor types. Moreover, dysregulation of miR-34c has been proven to regulate tumor cell proliferation, apoptosis, senescence, migration and invasion. The fluorescinated hsa-miR-34c probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-31

Hsa-miR-31 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM031-100 (ASR*)
 Specificity: miR-31
 Recommended Barrier: FB-HM031
 Control:

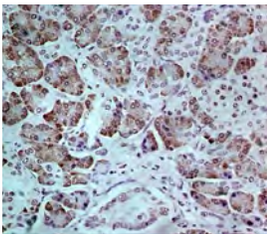
miR-31 is known as a tumor suppressor miRNA. miR-31 is frequently deleted and is the most underexpressed microRNA in certain tumors. It has been shown to affect the levels of tumor suppressor protein p53. The fluorescinated hsa-miR-31 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-92a

Hsa-miR-92a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM092A-100 (ASR*)
 Specificity: miR-92a
 Recommended Barrier: FB-HM092A
 Control:

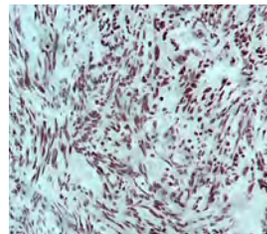
miR-92a is highly expressed in some tumors. The fluorescinated hsa-miR-92a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-34a

Hsa-miR-34a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM034A-100 (ASR*)
 Specificity: miR-34a
 Recommended Barrier: FB-HM034A
 Control:

The human miR-34a precursor is transcribed from chromosome 1. miR-34a itself is a transcriptional target of p53, suggesting a positive feedback loop between p53 and miR-34a. Thus, miR-34a functions as a tumor suppressor, in part, through a SIRT1-p53 pathway. miR-34 dysregulation is involved in the development of some tumors. The fluorescinated hsa-miR-34a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

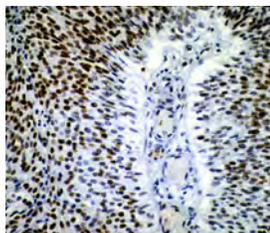
Hsa-miR-95

Hsa-miR-95 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM095-100 (ASR*)
 Specificity: miR-95
 Recommended Barrier: FB-HM095
 Control:

miR-95 expression was up-regulated in some tumors miR-95 increased proliferation by directly targeting SNX1. miR-95 expression levels correlated inversely with SNX1 protein levels. The fluorescinated hsa-miR-95 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

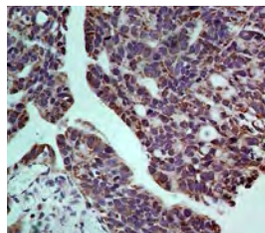
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-96

Hsa-miR-96 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM096-100 (ASR*)
 Specificity: miR-96
 Recommended Barrier: FB-HM096
 Control:

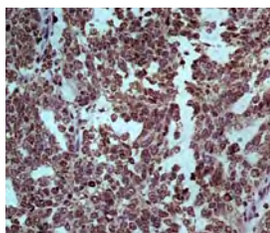
miR-96 expression decreases the transcript and protein levels of FOXO1 by binding to one of two predicted binding sites in the FOXO1 3'-UTR sequence. The fluorescinated hsa-miR-96 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-99b

Hsa-miR-99b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM099B-100 (ASR*)
 Specificity: miR-99b
 Recommended Barrier: FB-HM099B
 Control:

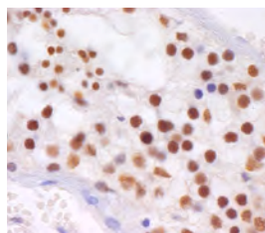
miR-99 family members miR-99a, -99b, and -100 were downregulated in tumor cell lines relative to the parental cell lines. miR-99 family members were also downregulated in human tumor tissue compared with normal tissue. The fluorescinated hsa-miR-99b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-98

Hsa-miR-98 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM098-100 (ASR*)
 Specificity: miR-98
 Recommended Barrier: FB-HM098
 Control:

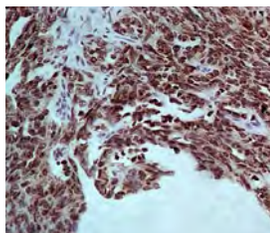
The ectopic expression of miR-98 inhibited tumor cell proliferation, invasion, and angiogenesis through repressing ALK4 and MMP11 expression. The fluorescinated hsa-miR-98 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-100

Hsa-miR-100 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM100-100 (ASR*)
 Specificity: miR-100
 Recommended Barrier: FB-HM100
 Control:

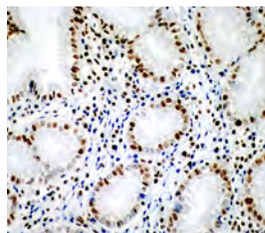
miR-100 is lost in many tumors and have potential function as a tumor suppressor. miR-100 inhibits the tumorigenicity, motility and invasiveness of tumor cells, and is commonly downregulated in human tumors due to hypermethylation. The fluorescinated hsa-miR-100 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-99a

Hsa-miR-99a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM099A-100 (ASR*)
 Specificity: miR-99a
 Recommended Barrier: FB-HM099A
 Control:

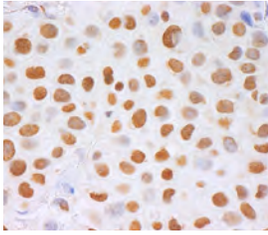
miR-99 family members miR-99a, -99b, and -100 were downregulated in tumor cell lines relative to the parental cell lines. miR-99 family members were also downregulated in human tumor tissue compared with normal tissue. The fluorescinated hsa-miR-99a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-101-3p

Hsa-miR-101-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM101-3P-100 (ASR*)
 Specificity: miR-101-3p
 Recommended Barrier: FB-HM101-3P
 Control:

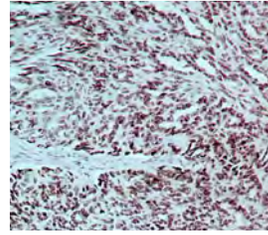
NDY1 up-regulation is shown to trigger the binding of EZH2 and NDY1 to the miR-101 locus. Activation of this pathway is essential for the epigenetic regulation of gene expression elicited by FGF-2. The fluorescinated hsa-miR-101-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-106a

Hsa-miR-106a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM106A-100 (ASR*)
 Specificity: miR-106a
 Recommended Barrier: FB-HM106A
 Control:

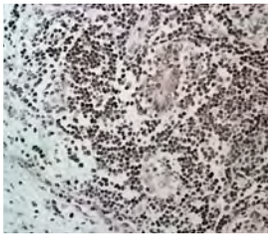
Sp1 and Egr1 are found to have an important role in miR-106a transcription and thus indirectly regulate the expression of IL-10 post-transcriptionally. The fluorescinated hsa-miR-106a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-124

Hsa-miR-124 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM124-100 (ASR*)
 Specificity: miR-124
 Recommended Barrier: FB-HM124
 Control:

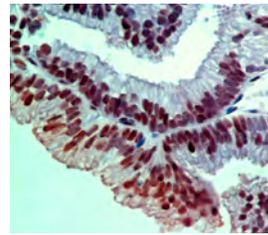
The mature miR-124 sequence is processed from 3 separate premature sequences, located at chromosomes 8p23.1 (miR-124-1), 8q12.3 (miR-124-2) and 20q13.33 (miR-124-3). The fluorescinated hsa-miR-124 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-107

Hsa-miR-107 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM107-100 (ASR*)
 Specificity: miR-107
 Recommended Barrier: FB-HM107
 Control:

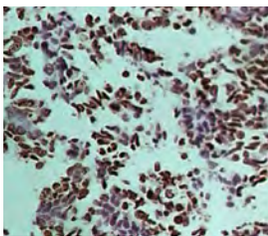
miR-107 is a microRNA expressed by human tumor specimens and regulated by p53. miR-107 decreases hypoxia signaling by suppressing expression of hypoxia inducible factor-1 β (HIF-1 β). miR-107 may have a tumor suppressor function by directly targeting CDK6 to inhibit the proliferation and invasion activities. The fluorescinated hsa-miR-107 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-125a

Hsa-miR-125a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM125A-100 (ASR*)
 Specificity: miR-125a
 Recommended Barrier: FB-HM125A
 Control:

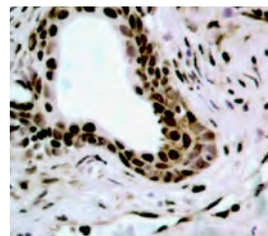
miR-125 family has been reported to be implicated in a variety of tumors and other diseases as either repressors or promoters. miR-125 family play crucial roles in many different cellular processes like cell differentiation, proliferation and apoptosis by targeting many different transcription factors, matrix-metalloprotease, and growth factors. The fluorescinated hsa-miR-125a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-122

Hsa-miR-122 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM122-100 (ASR*)
 Specificity: miR-122
 Recommended Barrier: FB-HM122
 Control:

miR-122 is specifically repressed in a subset of primary tumors that are characterized by poor prognosis. The loss of miR-122 results in an increase of cell migration and invasion and that restoration of miR-122 reverses this phenotype. miR-122 is a marker of hepatocyte-specific differentiation and an important determinant in the control of cell migration and invasion. The fluorescinated hsa-miR-122 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

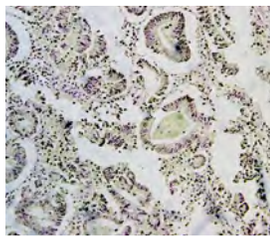
Hsa-miR-125b

Hsa-miR-125b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM125B-100 (ASR*)
 Specificity: miR-125b
 Recommended Barrier: FB-HM125B
 Control:

Enforced miR-125b expression in mammary cells is shown to decrease cell proliferation by inducing G2/M cell cycle arrest and reduced anchorage-independent cell growth of cells of mammary origin. The fluorescinated hsa-miR-125b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

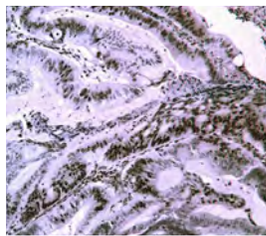
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-126

Hsa-miR-126 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM126-100 (ASR*)
 Specificity: miR-126
 Recommended Barrier: FB-HM126
 Control:

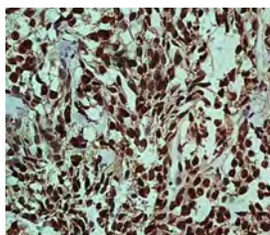
miR-126 is a microRNA expressed predominately by endothelial cells and controls angiogenesis. The fluorescinated hsa-miR-126 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-129

Hsa-miR-129 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM129-100 (ASR*)
 Specificity: miR-129
 Recommended Barrier: FB-HM129
 Control:

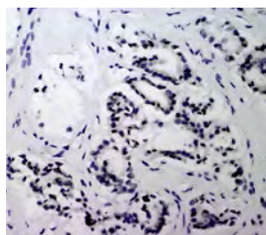
miR-129-5p expression is down-regulated in several tumor types. miR-129-5p promotes apoptosis and enhances chemosensitivity, while decreased miR-129-5p expression, as a result of hypermethylation of the miR-129 promoter, is associated with poor clinicopathological factors, such as clinical stage and progression in several tumors. The fluorescinated hsa-miR-129 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-127-3p

Hsa-miR-127-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM127-3P-100 (ASR*)
 Specificity: miR-127-3p
 Recommended Barrier: FB-HM127-3P
 Control:

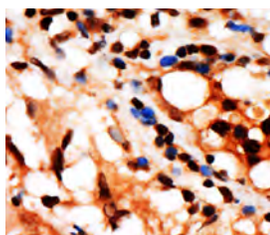
miR-127 is highly expressed in normal prostate and bladder tissues. miR-127 functions to regulate the expression levels of genes involved in lung development, placental formation and apoptosis. The fluorescinated hsa-miR-127-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-130b

Hsa-miR-130b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM130B-100 (ASR*)
 Specificity: miR-130b
 Recommended Barrier: FB-HM130B
 Control:

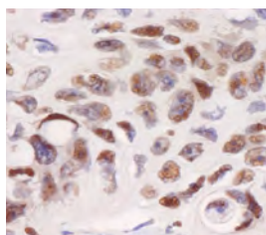
miR-130b, located at the 22q11 locus, plays an oncogenic or suppressive role in several tumors. The fluorescinated hsa-miR-130b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-128

Hsa-miR-128 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM128-100
 Specificity: miR-128
 Recommended Barrier: FB-HM128
 Control:

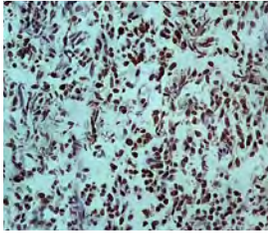
miRNA-128 is the most abundant brain-enriched microRNA that is induced during neuronal differentiation. Apart from brain, miRNA-128 has also been found in the skeletal muscle. Down regulation of miRNA-128 has been reported in several brain cancers such as glioblastoma, medulloblastoma and neuroblastoma. The fluorescinated hsa-miR-128 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-132

Hsa-miR-132 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM132-100 (ASR*)
 Specificity: miR-132
 Recommended Barrier: FB-HM132
 Control:

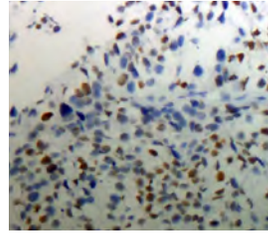
miR-132, transcribed from an intergenic region on human chromosome 17, is aberrantly expressed in many tumor types. The fluorescinated hsa-miR-132 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-133a

Hsa-miR-133a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM133A-100 (ASR*)
 Specificity: miR-133a
 Recommended Barrier: FB-HM133A
 Control:

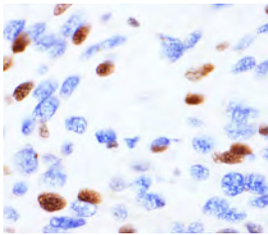
miR-133a is downregulated in some tumor types. The fluorescinated hsa-miR-133a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-135b

Hsa-miR-135b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM135B-100 (ASR*)
 Specificity: miR-135b
 Recommended Barrier: FB-HM135B
 Control:

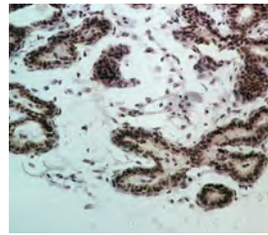
miR-135b is involved in the progression of several types of tumors and it is frequently dysregulated in tumor tissue. The fluorescinated hsa-miR-135b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-133b

Hsa-miR-133b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM133B-100 (ASR*)
 Specificity: miR-133b
 Recommended Barrier: FB-HM133B
 Control:

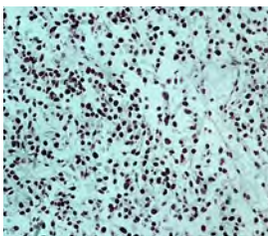
miR-133b is significantly downregulated in many tumor types. The fluorescinated hsa-miR-133b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-136

Hsa-miR-136 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM136-100 (ASR*)
 Specificity: miR-136
 Recommended Barrier: FB-HM136
 Control:

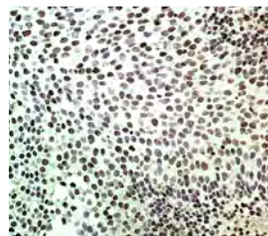
miR-136 was significantly downregulated in tumor specimens. The low-level expression of miR-136 is significantly associated with a more aggressive and/or poor prognostic phenotype. The fluorescinated hsa-miR-136 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-135a

Hsa-miR-135a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM135A-100 (ASR*)
 Specificity: miR-135a
 Recommended Barrier: FB-HM135A
 Control:

miR-135a is significantly downregulated in the tumor cell lines and plays a tumor-suppressive role. miR-135a expression is downregulated in the majority of human tumor tissues compared with pair-matched adjacent non-tumor tissues. The fluorescinated hsa-miR-135a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

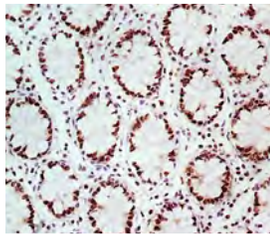
Hsa-miR-137

Hsa-miR-137 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM137-100 (ASR*)
 Specificity: miR-137
 Recommended Barrier: FB-HM137
 Control:

Recently studies revealed that miR-137 play essential roles in tumorigenesis. miR-137 modulates tumor cell growth, invasion and sensitivity. miR-137 was significantly down-regulated in tumors and inhibited proliferation of tumor cells by targeting PAK2. The fluorescinated hsa-miR-137 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

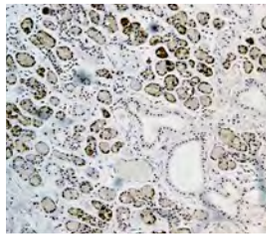
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-138

Hsa-miR-138 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM138-100 (ASR*)
 Specificity: miR-138
 Recommended Barrier: FB-HM138
 Control:

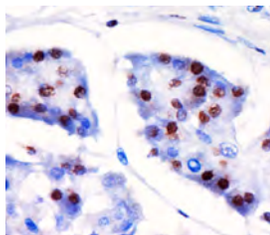
The down-regulation of microRNA-138 has been frequently observed in tumors with decreased levels of cell proliferation and colony formation. The fluorescinated hsa-miR-138 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-141

Hsa-miR-141 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM141-100(ASR*)
 Specificity: miR-141
 Recommended Barrier: FB-HM141
 Control:

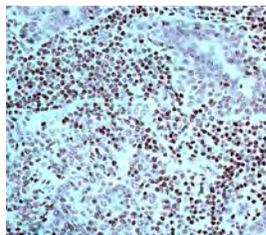
miR-141, along with miR-200c, is an important member of the miR-200 family for regulating the epithelial to mesenchymal transition. The fluorescinated hsa-miR-141 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-139

Hsa-miR-139 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM139-100
 Specificity: miR-139
 Recommended Barrier: FB-HM139
 Control:

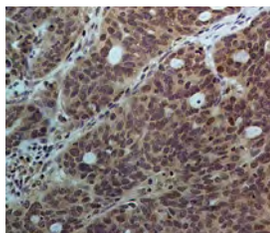
miRNA-139 is located in 11q13.4 and has anti-oncogenic and antimetastatic activity in humans. miR-139 may be the candidate to serve as promising biomarkers with sufficient sensitivity and specificity for the diagnosis of cancer in a clinical setting. The fluorescinated hsa-miR-139 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-142-3p

Hsa-miR-142-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM142-3P-100 (ASR*)
 Specificity: miR-142-3p
 Recommended Barrier: FB-HM142-3P
 Control:

miR-142-3p is involved in the progression of several tumor types. The fluorescinated hsa-miR-142-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-140

Hsa-miR-140 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM140-100 (ASR*)
 Specificity: miR-140
 Recommended Barrier: FB-HM140
 Control:

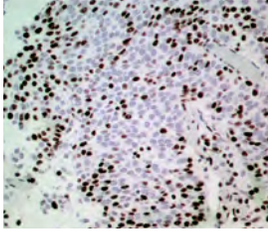
miR-140 functions as a tumor suppressor in many tumors and is significantly downregulated in human tumor tissues. Overexpression of miR-140 inhibited tumor growth, invasion, and metastasis. The fluorescinated hsa-miR-140 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-143

Hsa-miR-143 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM143-100 (ASR*)
 Specificity: miR-143
 Recommended Barrier: FB-HM143
 Control:

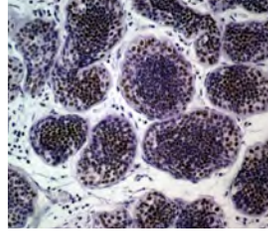
miR-143 specifically targets PKC ϵ and regulates its expression. Anti-miR-143 promotes cell proliferation, decreases apoptosis and up-regulates PKC ϵ expression. The fluorescinated hsa-miR-143 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-144

Hsa-miR-144 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM144-100 (ASR*)
 Specificity: miR-144
 Recommended Barrier: FB-HM144
 Control:

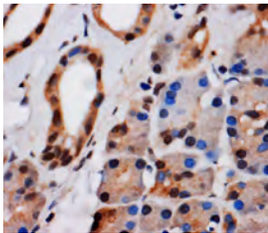
miR-144 is shown to promote cell proliferation, migration and invasion through repression of PTEN and targeted by zinc finger X-chromosomal protein. The fluorescinated hsa-miR-144 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization

Hsa-miR-146b

Hsa-miR-146b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM146B-100 (ASR*)
 Specificity: miR-146b
 Recommended Barrier: FB-HM146B
 Control:

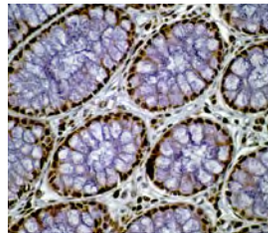
The expression of miR-146b-5p is known to be dysregulated in solid tumors and acts either as a tumor suppressor or promoter. The fluorescinated hsa-miR-146b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-145

Hsa-miR-145 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM145-100
 Specificity: miR-145
 Recommended Barrier: FB-HM145
 Control:

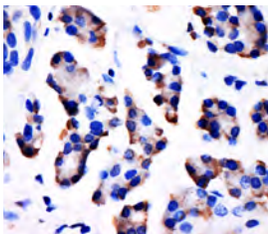
MiR-145 could serve as a tumor suppressor by targeting paxillin gene, it inhibited TGF- β -induced epithelial-mesenchymal transition and invasion through repression of SMAD3 in non-small cell lung cancer cells, it played pivotal roles in bladder cancer cells by regulating ubiquitin-like with PHD and ring finger domains 1. These findings provide novel insights into the potential mechanisms of cancer oncogenesis and suggest novel therapeutic strategies. The fluorescinated hsa-miR-145 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-147b

Hsa-miR-147b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM147B-100 (ASR*)
 Specificity: miR-147b
 Recommended Barrier: FB-HM147B
 Control:

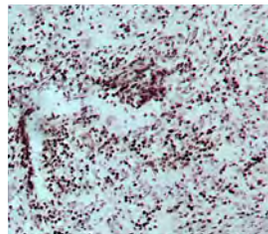
Studies demonstrated the participation of miR-147b in a negative feedback loop that is able to inhibit the pro-inflammatory response of macrophages to multiple TLR ligands. The fluorescinated hsa-miR-147b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-146a

Hsa-miR-146a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM146A-100 (ASR*)
 Specificity: miR-146a
 Recommended Barrier: FB-HM146A
 Control:

miR-146a plays a mechanistic role of in endotoxin-induced differential cross-regulation of TLR Signaling. The fluorescinated hsa-miR-146a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

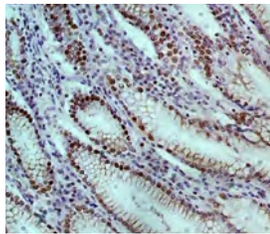
Hsa-miR-148a

Hsa-miR-148a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM148A-100 (ASR*)
 Specificity: miR-148a
 Recommended Barrier: FB-HM148A
 Control:

miR-148a expression is downregulated in several types of tumors and plays multiple roles as a tumor suppressor. The fluorescinated hsa-miR-148a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

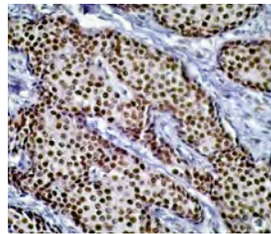
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-148b

Hsa-miR-148b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM148B-100 (ASR*)
 Specificity: miR-148b
 Recommended Barrier: FB-HM148B
 Control:

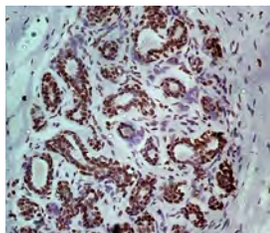
miR-148b was significantly downregulated in human tumors. Overexpression of miR-148b suppressed the growth of tumor cells, attributable to induction of apoptosis and cell-cycle arrest at S-phase. The fluorescinated hsa-miR-148b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-151a-3p

Hsa-miR-151a-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM151A-3p-100 (ASR*)
 Specificity: miR-151a-3p
 Recommended Barrier: FB-HM151A-3P
 Control:

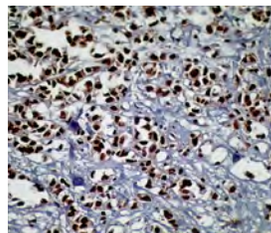
miR-151a has been demonstrated to be directly regulated by the p53-family of transcription factors and contributes to the tuning of p53-induced responses. The fluorescinated hsa-miR-151a-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-149

Hsa-miR-149 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM149-100 (ASR*)
 Specificity: miR-149
 Recommended Barrier: FB-HM149
 Control:

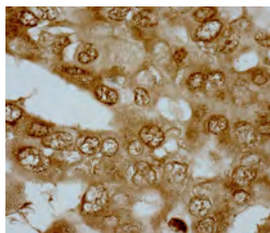
miR-149 has been identified to be a suppressor of tumor metastasis. Increased miR-149 levels block lung colonization in vivo. Low level of miR-149 and high level of GIT1 was significantly associated with advanced stages of tumor, as well as with lymph node metastasis. The fluorescinated hsa-miR-149 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-152-3p

Hsa-miR-152-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM152-3p-100 (ASR*)
 Specificity: miR-152-3p
 Recommended Barrier: FB-HM152-3P
 Control:

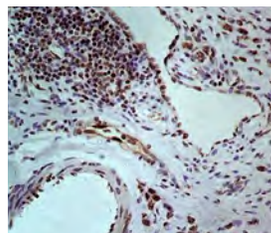
miR-152 is suggested to play a role in S-phase and G2/M-phase cell cycle progression of diploid fibroblasts. The fluorescinated hsa-miR-152-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-150

Hsa-miR-150 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM150-100 (ASR*)
 Specificity: miR-150
 Recommended Barrier: FB-HM150
 Control:

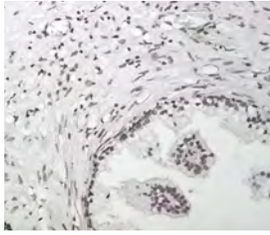
miR-150 is mainly expressed in the lymph nodes and spleen and is highly up-regulated during the development of mature T and B cells. The fluorescinated hsa-miR-150 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-153

Hsa-miR-153 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM153-100 (ASR*)
 Specificity: miR-153
 Recommended Barrier: FB-HM153
 Control:

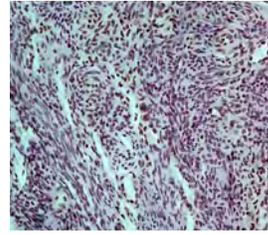
miR-153 upregulation promoted tumor invasiveness by indirectly initiating matrix metalloprotease enzyme 9 productions. Overexpression of miR-153 in tumor cells enhanced the G1/S transitional promoter, cyclin D1 expression, and decreased cyclin-dependent kinase (CDK) inhibitor, p21(Cip1) expression via downregulation of PTEN tumor suppressor gene and activated AKT signaling. The fluorescinated hsa-miR-153 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-154

Hsa-miR-154 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM154-100 (ASR*)
 Specificity: miR-154
 Recommended Barrier: FB-HM154
 Control:

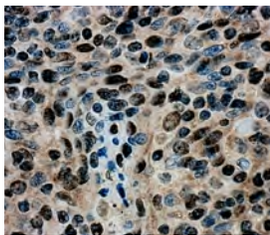
miR-154 is deregulated and functions as a candidate tumor suppressor in some tumors. miR-154 was decreased in tumor tissues and cell lines. Ectopic expression of miR-154 remarkably suppressed cell proliferation and colony formation, migration and invasion. The fluorescinated hsa-miR-154 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-181b

Hsa-miR-181b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM181B-100 (ASR*)
 Specificity: miR-181b
 Recommended Barrier: FB-HM181B
 Control:

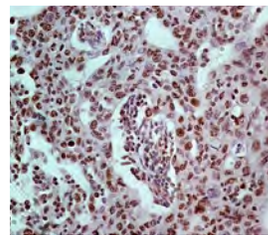
The downregulated miR-181b was involved in oncogenesis. miR-181b functioned as tumor suppressors which triggered growth inhibition, induced apoptosis and inhibited invasion in tumor cells. The fluorescinated hsa-miR-181b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-155

Hsa-miR-155 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM155-100 (ASR*)
 Specificity: miR-155
 Recommended Barrier: FB-HM155
 Control:

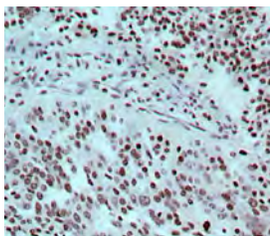
miR-155 is expressed in a variety of immune cell types and present at low levels in most of these cells until their activation by immune stimuli such as toll-like receptor ligands. The fluorescinated hsa-miR-155 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-181c

Hsa-miR-181c detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM181C-100 (ASR*)
 Specificity: miR-181c
 Recommended Barrier: FB-HM181C
 Control:

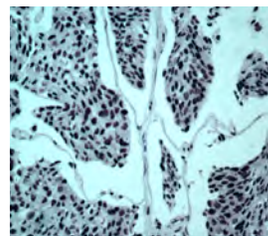
Aberrant miR-181c expression is related to many tumor types. The fluorescinated hsa-miR-181c probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-181a

Hsa-miR-181a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM181A-100 (ASR*)
 Specificity: miR-181a
 Recommended Barrier: FB-HM181A
 Control:

miR-181a expression was upregulated in metastatic tumors and serves as a predictive biomarker for metastasis and patient survival. miR-181a expression is highly associated with the development of metastatic disease. The fluorescinated hsa-miR-181a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

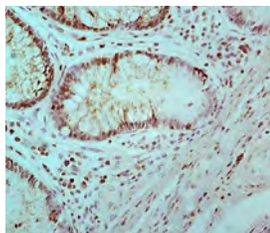
Hsa-miR-182

Hsa-miR-182 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM182-100 (ASR*)
 Specificity: miR-182
 Recommended Barrier: FB-HM182
 Control:

miR-182, member of a miRNA cluster is located at chromosomal locus 7q31-34, is commonly overexpressed in many tumor types. The fluorescinated hsa-miR-182 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

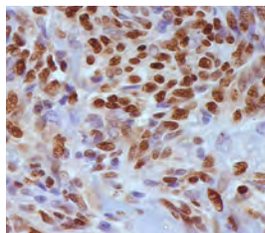
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-183

Hsa-miR-183 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM183-100 (ASR*)
 Specificity: miR-183
 Recommended Barrier: FB-HM183
 Control:

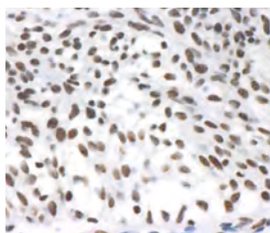
The level of miR-183 expression in tumor tissue has been reported to be higher than adjacent normal tissues, and miR-183 regulates diverse mediators of tumor survival and function, including targeting the tumor suppressor Bmi-1, EGR1, PTEN and SMAD4. The fluorescinated hsa-miR-183 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-185

Hsa-miR-185 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM185-100 (ASR*)
 Specificity: miR-185
 Recommended Barrier: FB-HM185
 Control:

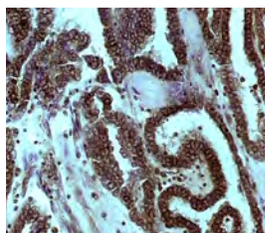
miR-185 has been identified as an important factor in several tumors. This relates to the fact that miR-185 is closely associated with tumor size, pTNM stage, lymph node, and perineural invasion. The fluorescinated hsa-miR-185 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-183-3p

Hsa-miR-183-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM183-3P-100 (ASR*)
 Specificity: miR-183-3p
 Recommended Barrier: FB-HM183-3P
 Control:

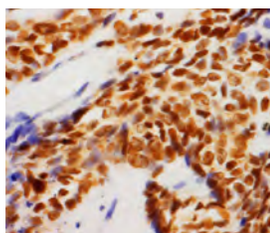
miR-183-3p was up-regulated in tumor tissues when compared with the corresponding normal tissues. Moreover, the expression of miR-183-3p in tumor tissue was found to be associated with lymph node metastasis, clinical stage, and EGFR mutation. The fluorescinated hsa-miR-183-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-186

Hsa-miR-186 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM186-100 (ASR*)
 Specificity: miR-186
 Recommended Barrier: FB-HM186
 Control:

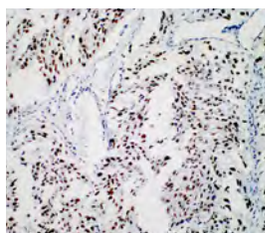
Overexpression of miR-186 in tumor cells inhibited proliferation by inducing G1-S checkpoint arrest. miR-186 expression promoted cell-cycle progression and accelerated the proliferation of tumor cells. The fluorescinated hsa-miR-186 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-184

Hsa-miR-184 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM184-100 (ASR*)
 Specificity: miR-184
 Recommended Barrier: FB-HM184
 Control:

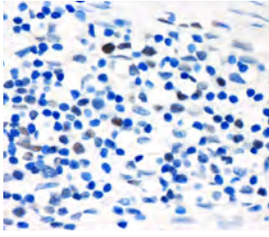
miR-184 may be oncogenic or tumor suppressive in different tumor types. The fluorescinated hsa-miR-184 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-187

Hsa-miR-187 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM187-100 (ASR*)
 Specificity: miR-187
 Recommended Barrier: FB-HM187
 Control:

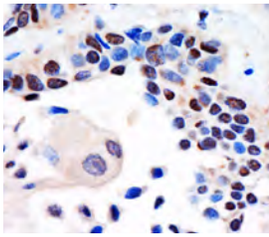
miR-187 is shown to overexpress in the subtype exhibiting loss of chromosome 11q but not in the MYCN amplified subtype. The fluorescinated hsa-miR-187 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-190a

Hsa-miR-190a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM190a-100
 Specificity: miR-190a
 Recommended Barrier: FB-HM190a
 Control:

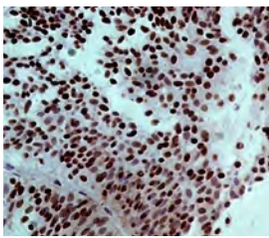
miR-190a belongs to the miRNA family and is located in the tail intron regions of two genes on 15q22.2. miR-190a is downregulated in aggressive neuroblastoma and prostate cancer. The miR-190a mediated effects rely on an extensive network of molecular changes in tumor cells and affects several transcriptional factors, tumor suppressor and interferon response pathways. The fluorescinated hsa-miR-190a probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-190b

Hsa-miR-190b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM190b-100
 Specificity: miR-190b
 Recommended Barrier: FB-HM190b
 Control:

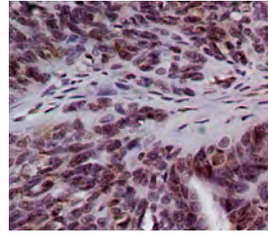
miR-190b negatively regulates tumor suppressor Bcl-2, possibly confers radio-sensitivity in gastric cancer cells. Also, miR-190b has been identified as a potential biomarker for ERα(+) breast cancer. The fluorescinated hsa-miR-190b probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-191

Hsa-miR-191 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM191-100 (ASR*)
 Specificity: miR-191
 Recommended Barrier: FB-HM191
 Control:

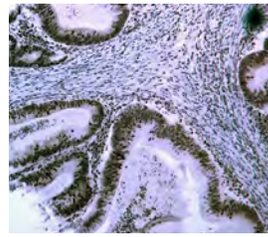
miR-191 has been found to be dysregulated in a large number of different types of human tumors. The fluorescinated hsa-miR-191 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-192

Hsa-miR-192 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM192-100 (ASR*)
 Specificity: miR-192
 Recommended Barrier: FB-HM192
 Control:

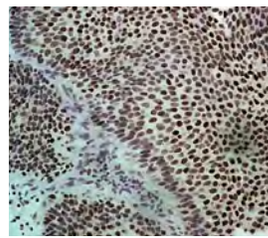
miR-192 is thought to be positive regulators of p53, a human tumor suppressor. It has also been suggested that miR-192 could be used as a biomarker for drug-induced liver damage. The fluorescinated hsa-miR-192 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-193a-3p

Hsa-miR-193a-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM193A-3P-100 (ASR*)
 Specificity: miR-193a-3p
 Recommended Barrier: FB-HM193A-3P
 Control:

miR-193a-3p induces the accumulation of intracellular reactive oxygen species, DNA damage in tumor cells. Furthermore, miR-193a-3p directly recognizes the 3'-UTR of the ERBB4 transcript and regulates ERBB4 expression, one of four ErbB receptor tyrosine kinase family members. The fluorescinated hsa-miR-193a-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

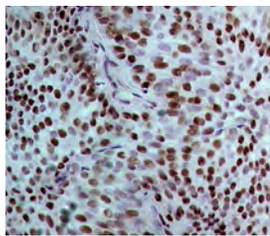
Hsa-miR-193b

Hsa-miR-193b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM193B-100 (ASR*)
 Specificity: miR-193b
 Recommended Barrier: FB-HM193B
 Control:

Aberrant expression of miR-193b is frequently observed in tumor tissues and it acts as a tumor suppressor in many types of tumors. miR-193b is down-regulated in tumor tissue and can promote tumorigenesis by inhibiting stathmin 1 and urokinase-type plasminogen activator (uPA). The fluorescinated hsa-miR-193b probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

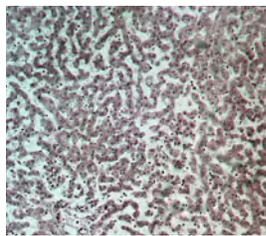
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-194

Hsa-miR-194 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM194-100 (ASR*)
 Specificity: miR-194
 Recommended Barrier: FB-HM194
 Control:

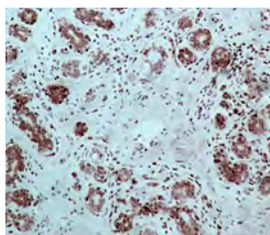
miR-194 is expressed in liver parenchymal cells, and in human gastrointestinal tract. miR-194 plays a role in the activation of stellate cells during liver fibrogenesis. miR-194 expression varies in human organs and in different status of hepatocyte differentiation. miR-194 is an epithelial cell-specific marker in the liver. The fluorescinated hsa-miR-194 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-197

Hsa-miR-197 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM197-100 (ASR*)
 Specificity: miR-197
 Recommended Barrier: FB-HM197
 Control:

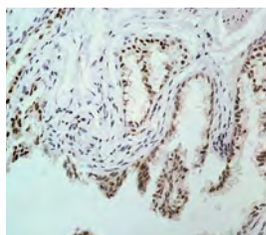
miR-197 is an onco-miR which functions as a key repressor of p53-dependent apoptotic cascade in tumor cells. The fluorescinated hsa-miR-197 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-195

Hsa-miR-195 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM195-100 (ASR*)
 Specificity: miR-195
 Recommended Barrier: FB-HM195
 Control:

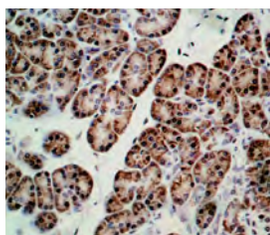
miR-195 is aberrantly expressed in multiple types of disease. miR-195 was significantly downregulated in tumors. The fluorescinated hsa-miR-195 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-198

Hsa-miR-198 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM198-100 (ASR*)
 Specificity: miR-198
 Recommended Barrier: FB-HM198
 Control:

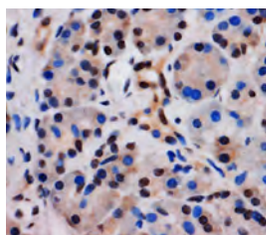
It has been reported that several genes can be targeted by miR-198 in different type of tumors and miR-198 has different functions during tumor progression. miR-198 has been shown to be a tumor suppressor by inhibition of tumor cell growth, migration and invasion. The fluorescinated hsa-miR-198 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-196a

Hsa-miR-196a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM196A-100 (ASR*)
 Specificity: miR-196a
 Recommended Barrier: FB-HM196A
 Control:

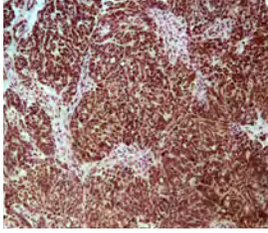
miR-196a is a microRNA that suppresses the expression of specific homeobox genes that are of high relevance for the development of human embryos. The fluorescinated hsa-miR-196a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-199a

Hsa-miR-199a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM199A-100 (ASR*)
 Specificity: miR-199a
 Recommended Barrier: FB-HM199A
 Control:

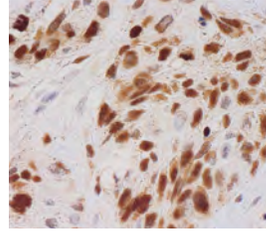
miR-199a, which is encoded from the opposite strand of DN2 (Dynamin 2 is a key component of endocytic machinery that is transcriptionally suppressed by HIF-1), is shown to exert reciprocal negative regulation upon HIF-1 α and HIF-2 α . The fluorescinated hsa-miR-199a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-200a

Hsa-miR-200a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM200A-100 (ASR*)
 Specificity: miR-200a
 Recommended Barrier: FB-HM200A
 Control:

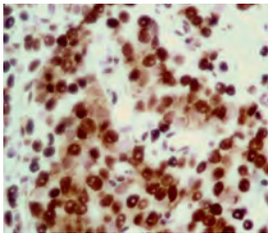
Gain and loss of function assays for miR-200a is shown to lead to a significant differential and converse expression of epithelial mesenchymal transition (EMT)-related genes. The fluorescinated hsa-miR-200a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-203a-3p

Hsa-miR-203a-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM203A-3p-100 (ASR*)
 Specificity: miR-203a-3p
 Recommended Barrier: FB-HM203A-3P
 Control:

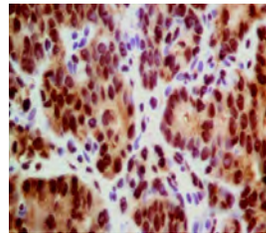
miR-203 is an antiproliferative microRNA involved in skin differentiation that targets the 3'-UTR of the "stemness-maintaining" transcription factor $\Delta Np63\alpha$. The fluorescinated hsa-miR-203a-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-200b

Hsa-miR-200b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM200B-100 (ASR*)
 Specificity: miR-200b
 Recommended Barrier: FB-HM200B
 Control:

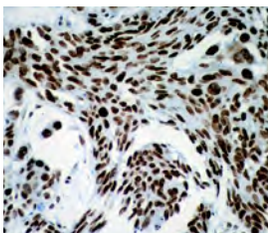
miR-200b targets v-ets erythroblastosis virus E26 oncogene homolog 1 (Ets-1) and is down-regulated by hypoxia to induce angiogenic response of endothelial cells. The fluorescinated hsa-miR-200b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-204

Hsa-miR-204 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM204-100 (ASR*)
 Specificity: miR-204
 Recommended Barrier: FB-HM204
 Control:

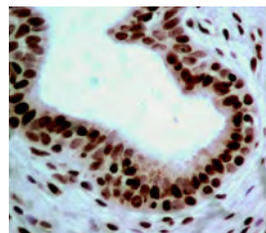
miR-204 targeting of the Ankrd13A gene is found to control both nesenchymal neural crest and lens cell migration. The fluorescinated hsa-miR-204 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-200c

Hsa-miR-200c detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM200C-100 (ASR*)
 Specificity: miR-200c
 Recommended Barrier: FB-HM200C
 Control:

Overexpression of the miR-200c is reported to lead to reduced expression of transcription factor 8 and increased expression of E-Cadherin. The fluorescinated hsa-miR-200c probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

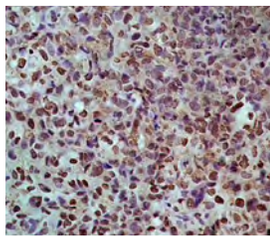
Hsa-miR-205

Hsa-miR-205 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM205-100 (ASR*)
 Specificity: miR-205
 Recommended Barrier: FB-HM205
 Control:

miR-205 is capable of suppressing epithelial to mesenchymal transition by targeting the transcriptional factors ZEB1 and SIP1. miR-205 has also been shown to regulate E-Cadherin and possibly target PTEN. The fluorescinated hsa-miR-205 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

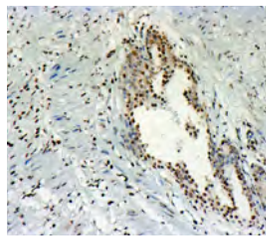
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-206

Hsa-miR-206 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM206-100 (ASR*)
 Specificity: miR-206
 Recommended Barrier: FB-HM206
 Control:

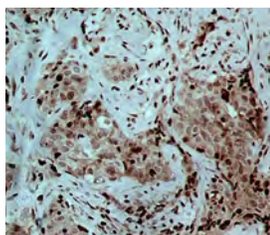
miR-206 targets HSP60 leading to accelerated glucose-mediated apoptosis in cardiomyocytes. miR-206 is reported to decrease endogenous ER α mRNA and protein levels. The fluorescinated hsa-miR-206 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-212

Hsa-miR-212 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM212-100 (ASR*)
 Specificity: miR-212
 Recommended Barrier: FB-HM212
 Control:

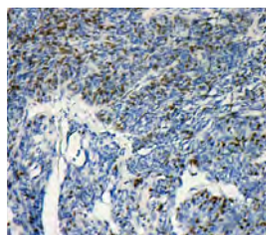
miR-212 expression is essential for the proper development, maturation and function of neurons. miR-212 deregulation is associated with several neurological disorders, such as Alzheimer's disease. The fluorescinated hsa-miR-212 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-210

Hsa-miR-210 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM210-100 (ASR*)
 Specificity: miR-210
 Recommended Barrier: FB-HM210
 Control:

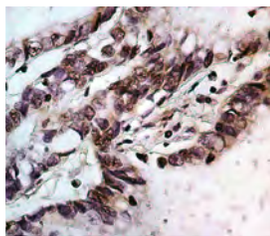
miR-210 has been strongly linked with the hypoxia pathway, and is upregulated in response to hypoxia-inducible factors. It is also overexpressed in cells affected by cardiac disease and tumors. miR-210 has been studied for its effects in rescuing cardiac function after myocardial infarcts via the up-regulation of angiogenesis and inhibition of cardiomyocyte apoptosis. The fluorescinated hsa-miR-210 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-214

Hsa-miR-214 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM214-100 (ASR*)
 Specificity: miR-214
 Recommended Barrier: FB-HM214
 Control:

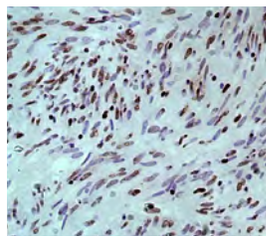
miR-214 expression level is associated with metastasis and invasion of tumors. miR-214 could inhibit the proliferation capacity, migration and invasion ability of HeLa cells. Plexin-B1, a target of miR-214, may function as an oncogene in human tumor HeLa cells. The fluorescinated hsa-miR-214 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-211

Hsa-miR-211 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM211-100 (ASR*)
 Specificity: miR-211
 Recommended Barrier: FB-HM210
 Control:

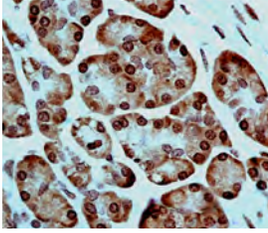
miR-211 is localized on intron 6 of the Trpm1 gene at 15q13-q14, a locus that is frequently lost in neoplasms. miR-211 functions and the effect of loss-of-function have been described in normal and tumor tissues. The fluorescinated hsa-miR-211 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-215

Hsa-miR-215 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM215-100 (ASR*)
 Specificity: miR-215
 Recommended Barrier: FB-HM215
 Control:

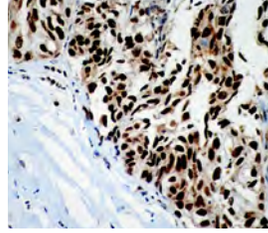
miR-215 identified from the microRNA cluster site at chromosome 1q41, has been reported to function as a tumor suppressor in a variety of human tumors by positive regulate p53. miR-215 suppressed the expression of key targets such as thymidylate synthase (TS), dihydrofolate reductase, and denticleless protein homolog (DTL). The fluorescinated hsa-miR-215 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-216a

Hsa-miR-216a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM216A-100 (ASR*)
 Specificity: miR-216a
 Recommended Barrier: FB-HM216A
 Control:

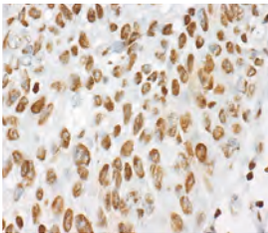
It was shown that TGF- β activates Akt in glomerular mesangial cells by inducing the miR-216a and miR-217, both of which target PTEN, an inhibitor of Akt activation. The fluorescinated hsa-miR-216a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-218

Hsa-miR-218 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM218-100 (ASR*)
 Specificity: miR-218
 Recommended Barrier: FB-HM218
 Control:

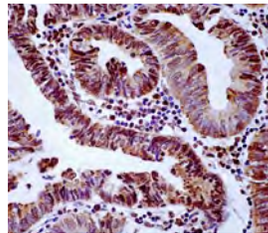
miR-218 is reported to be part of a regulatory circuit involving the Slit-Robo1 pathway. Decreased miR-218 levels eliminate Robo1 repression which activates the pathway through the interaction between Robo1 and Slit2. The fluorescinated hsa-miR-218 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-216b

Hsa-miR-216b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM216B-100 (ASR*)
 Specificity: miR-216b
 Recommended Barrier: FB-HM216B
 Control:

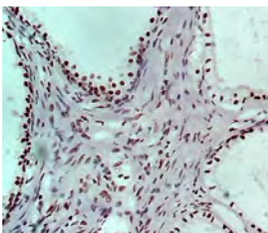
miR-216b was identified as a tumor suppressor in many tumors. Forced expression of miR-216b in Rink-1 cells inhibits cell proliferation and colony formation, which is correlated with reduced expression levels of epidermal growth factor receptor and matrix metalloproteinase-14 (MT1-MMP). Furthermore, miR-216b is dysregulated in bone marrow mesenchymal stem cells, and is associated with nonalcoholic fatty liver disease. The fluorescinated hsa-miR-216b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-221-3p

Hsa-miR-221-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM221-3p-100 (ASR*)
 Specificity: miR-221-3p
 Recommended Barrier: FB-HM221-3P
 Control:

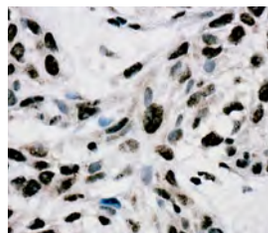
miR-221, together with miR-222, is encoded in tandem from a gene cluster located on chromosome X. Both miRNAs have been shown to directly target p27kip1, Bmf, PTEN, Mdm2, PUMA, and TRPS1. The fluorescinated hsa-miR-221-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-217

Hsa-miR-217 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM217-100 (ASR*)
 Specificity: miR-217
 Recommended Barrier: FB-HM217
 Control:

miR-217 targets oncogenes or tumor suppressor genes such as KRAS/WASF3 in different cell types by inhibiting tumor cell growth and anchorage-independent colony formation. The fluorescinated hsa-miR-217 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

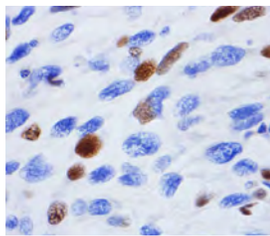
Hsa-miR-222

Hsa-miR-222 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM222-100 (ASR*)
 Specificity: miR-222
 Recommended Barrier: FB-HM222
 Control:

miR-222, together with miR-221, is encoded in tandem from a gene cluster located on chromosome X. Both miRNAs have been shown to directly target p27kip1, Bmf, PTEN, Mdm2, PUMA, and TRPS1. The fluorescinated hsa-miR-222 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

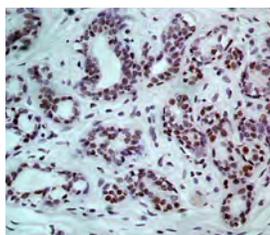
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-223

Hsa-miR-223 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM223-100 (ASR*)
 Specificity: miR-223
 Recommended Barrier: FB-HM223
 Control:

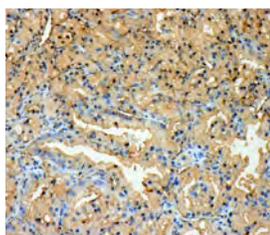
miR-223 is a hematopoietic specific microRNA with crucial functions in myeloid lineage development. It plays an essential role in promoting granulocytic differentiation. In some tumors the miR-223 downregulation is correlated with higher tumor burden, disease aggressiveness, and poor prognostic factors. The fluorescinated hsa-miR-223 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-224

Hsa-miR-224 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM224-100 (ASR*)
 Specificity: miR-224
 Recommended Barrier: FB-HM224
 Control:

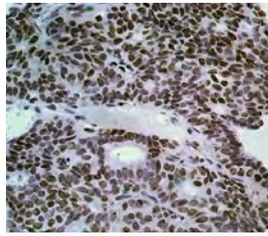
miR-224 could play an oncogenic role in the cellular processes of tumors. miR-224 has been shown to be involved in the tumorigenesis and development. The fluorescinated hsa-miR-224 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-296

Hsa-miR-296 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM296-100 (ASR*)
 Specificity: miR-296
 Recommended Barrier: FB-HM296
 Control:

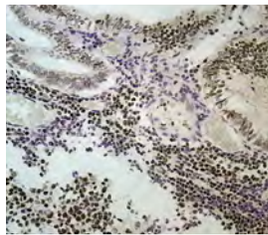
miR-296 was found to be located on chromosome 20q13.32, and it was reported that the 20q13.32–13.33 chromosome region is deleted in 20% of tumor tissues. In a recent study, it was demonstrated that miR-296 modulates tumor invasiveness by modulating HMGA1 expression. The fluorescinated hsa-miR-296 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-297

Hsa-miR-297 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM297-100 (ASR*)
 Specificity: miR-297
 Recommended Barrier: FB-HM297
 Control:

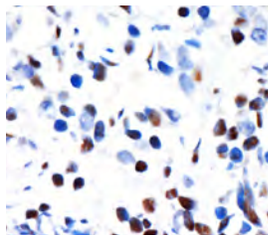
miR-297 was downregulated in human tumor tissues and negatively correlated with expression levels of MRP-2. The fluorescinated hsa-miR-297 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-300

Hsa-miR-300 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM300-100 (ASR*)
 Specificity: miR-300
 Recommended Barrier: FB-HM300
 Control:

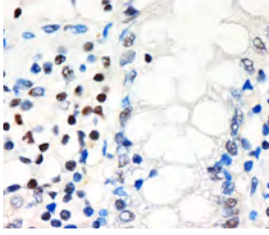
miR-300 was upregulated in several tumor types. miR-300 inhibits epithelial to mesenchymal transition and metastasis by targeting Twist. The fluorescinated hsa-miR-300 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-302b

Hsa-miR-302b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM302b-100
 Specificity: miR-302b
 Recommended Barrier: FB-HM302b
 Control:

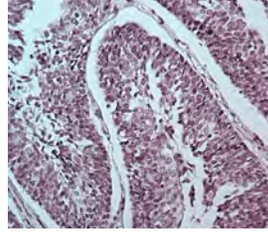
miRNA-302b is located in 11q13.4 and has anti-oncogenic and antimetastatic activity in humans. miR-302b may be the candidate to serve as promising biomarkers with sufficient sensitivity and specificity for the diagnosis of cancer in a clinical setting. The fluorescinated hsa-miR-302b probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-326

Hsa-miR-326 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM326-100
 Specificity: miR-326
 Recommended Barrier: FB-HM326
 Control:

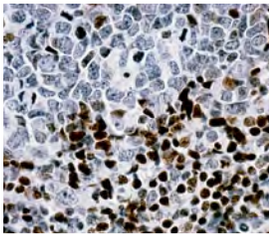
miR-326 is localized in the intron 1 of *Arb1* gene, and a well-known downstream component of Hedgehog signaling in cerebellar neuronal progenitor and tumor cells. miR-326 is also involved in Th-17 cells differentiation and progress of multiple sclerosis disease. The fluorescinated *hsa-miR-326* probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-330

Hsa-miR-330 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM330-100 (ASR*)
 Specificity: miR-330
 Recommended Barrier: FB-HM330
 Control:

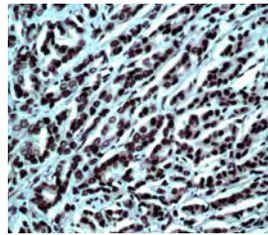
The expression of miR-330 in tumor cells enhanced cellular proliferation, promoted cell migration and invasion, and dampened cell apoptosis. The fluorescinated *hsa-miR-330* probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-328

Hsa-miR-328 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM328-100 (ASR*)
 Specificity: miR-328
 Recommended Barrier: FB-HM328
 Control:

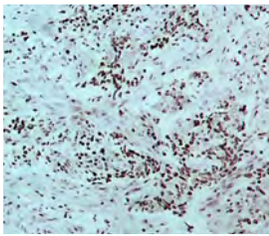
A study shows that miR-328 regulates zonation morphogenesis by targeting expression of hyaluronan receptor CD44. The fluorescinated *hsa-miR-328* probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-331-3p

Hsa-miR-331-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM331-3P-100 (ASR*)
 Specificity: miR-331-3p
 Recommended Barrier: FB-HM331-3P
 Control:

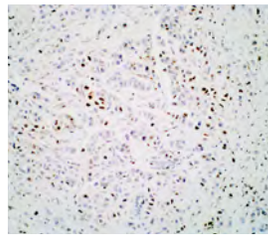
miR-331-3p expression is decreased in tumor tissue comparing to normal tissue. The fluorescinated *hsa-miR-331-3p* probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-329

Hsa-miR-329 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM329-100 (ASR*)
 Specificity: miR-329
 Recommended Barrier: FB-HM329
 Control:

miR-329 functions as a tumor suppressor in some malignancies. miR-329 was decreased in metastatic tumor tissues compared with primary tumor tissues. Overexpression of miR-329 substantially suppressed cell proliferation, colony formation, migration and invasion of tumor cells. The fluorescinated *hsa-miR-329* probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

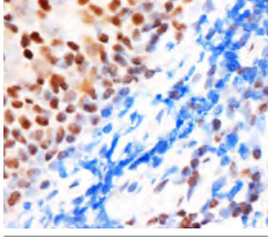
Hsa-miR-335

Hsa-miR-335 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM335-100 (ASR*)
 Specificity: miR-335
 Recommended Barrier: FB-HM335
 Control:

Differential microRNA expression analyses reveal that miR-335 is significantly down-regulated upon differentiation of human mesenchymal stem cells. The fluorescinated *hsa-miR-335* probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

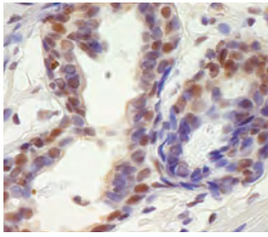
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-337

Hsa-miR-337 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM337-100
 Specificity: miR-337
 Recommended Barrier: FB-HM337
 Control:

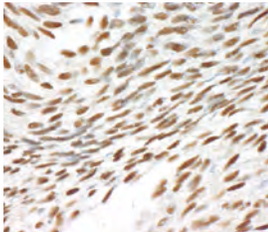
Many studies have shown miR-337 to be involved in tumor cell proliferation, migration, and invasion⁵. Its expression was found to be related to the tumor prognosis in some patients. One recent study showed miR-337 was minimally expressed in pancreatic ductal adenocarcinoma (PDAC) tissues, and its level was related to TNM stage, lymph node status, and survival in PDAC patients, which suggested that miR-337 could be used as determinants of PDAC patient prognosis. The fluorescinated hsa-miR-337 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-338-3p

Hsa-miR-338-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM338-3P-100 (ASR*)
 Specificity: miR-338-3p
 Recommended Barrier: FB-HM338-3P
 Control:

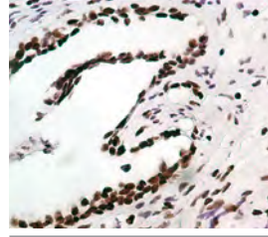
miR-338-3p was transcribed from the intron 8 of apoptosis-associated tyrosine kinase (AATK) gene, located on chromosome 17q25, playing a critical role in promoting cell death, neuronal differentiation and neurite extension. miR-338-3p could act as a tumor suppressor in several tumor types. The fluorescinated hsa-miR-338-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-339-5p

Hsa-miR-339-5p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM339-5P-100 (ASR*)
 Specificity: miR-339-5p
 Recommended Barrier: FB-HM339-5P
 Control:

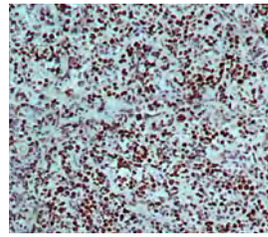
miR-339-5p targets BCL-6 and dramatically inhibited tumor cell migration and invasion *in vitro*. In addition, it has been reported that Dicer-regulated miR-339-5p promotes resistance of tumor cells to cytotoxic T-lymphocytes by down-regulation of ICAM-1. The fluorescinated hsa-miR-339-5p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-342-3p

Hsa-miR-342-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM342-3P-100 (ASR*)
 Specificity: miR-342-3p
 Recommended Barrier: FB-HM342-3P
 Control:

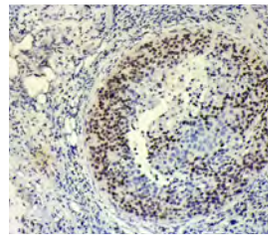
The level of miR-342-3p was significantly increased in tumor, and was inversely associated with the prognosis of patients. The fluorescinated hsa-miR-342-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-361

Hsa-miR-361 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM361-100 (ASR*)
 Specificity: miR-361
 Recommended Barrier: FB-HM361
 Control:

miR-361 was significantly downregulated in serum of tumor patients. The level of miR-361 was lower in tumor than in benign disease and healthy individuals. The fluorescinated hsa-miR-361 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

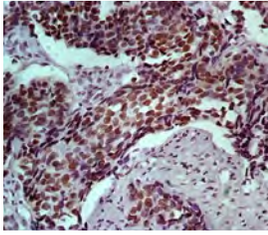
Hsa-miR-362

Hsa-miR-362 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM362-100 (ASR*)
 Specificity: miR-362
 Recommended Barrier: FB-HM362
 Control:

miR-362 is significantly up-regulated in tumor and associated with tumor progression. Inhibition of miR-362 in tumor cells dramatically decrease the cell proliferation, clonogenicity, migration and invasion *in vitro* as well as tumor growth and metastasis *in vivo*. The fluorescinated hsa-miR-362 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

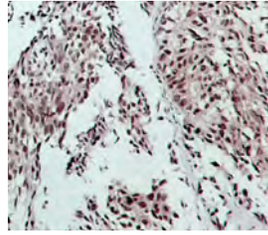
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-365a-3p

Hsa-miR-365a-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM365A-3P-100 (ASR*)
 Specificity: miR-365a-3p
 Recommended Barrier: FB-HM365A-3P
 Control:

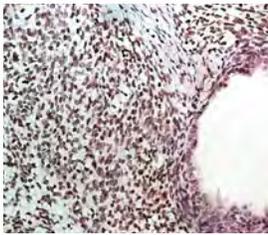
miR-365 is a direct negative regulator of IL-6. Ectopic expression of a miR-365 inhibitor elevated IL-6 expression. The negative regulation of miR-365 was strictly dependent on a microRNA binding element in the 3'-UTR of IL-6 mRNA. The fluorescinated hsa-miR-365a-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-374a

Hsa-miR-374a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM374A-100 (ASR*)
 Specificity: miR-374a
 Recommended Barrier: FB-HM374A
 Control:

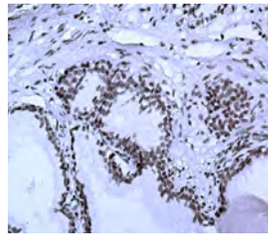
miR-374a was overexpressed in the tumors. Besides, miR-374a was involved in the tumorigenesis and metastasis by regulating the Wnt/catenin pathway. The fluorescinated hsa-miR-374a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-372

Hsa-miR-372 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM372-100 (ASR*)
 Specificity: miR-372
 Recommended Barrier: FB-HM372
 Control:

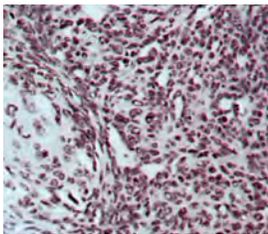
miR-372 belongs to the miR-371-372 gene cluster, which is located on chromosome 19q13.42. Recent studies demonstrated that miR-372 regulates the cell cycle, apoptosis, invasion, and proliferation in many types of human tumors. The fluorescinated hsa-miR-372 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-374b

Hsa-miR-374b detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM374B-100 (ASR*)
 Specificity: miR-374b
 Recommended Barrier: FB-HM374B
 Control:

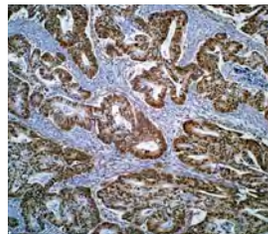
miR-374b is downregulated in tumor tissue and is an independent predictor of biochemical recurrence-free survival. The fluorescinated hsa-miR-374b probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-373

Hsa-miR-373 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM373-100 (ASR*)
 Specificity: miR-373
 Recommended Barrier: FB-HM373
 Control:

miR-373 stimulated tumor cell migration and invasion in vitro and in vivo. Certain tumor cell lines depend on endogenous miR-373 activity to migrate efficiently. The fluorescinated hsa-miR-373 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

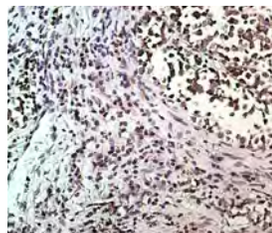
Hsa-miR-375

Hsa-miR-375 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM375-100 (ASR*)
 Specificity: miR-375
 Recommended Barrier: FB-HM375
 Control:

It has been shown that overexpression of miR-375 down-regulates while knockdown of miR-375 up-regulates CLDN1 mRNA and protein, respectively. The fluorescinated hsa-miR-375 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-376c

Hsa-miR-376c detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM376C-100 (ASR*)
 Specificity: miR-376c
 Recommended Barrier: FB-HM376C
 Control:

miR-376c was found to have potential complementary binding sites on the 3'UTR of ALK7 mRNA. miR-376c belongs to an evolutionary conserved miRNA family which also includes miR-376a, miR-376a* and miR-376b, and these genes are found in a syntenic cluster on human chromosome 14. The fluorescinated hsa-miR-376c probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-378a

Hsa-miR-378a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM378A-100 (ASR*)
 Specificity: miR-378a
 Recommended Barrier: FB-HM378A
 Control:

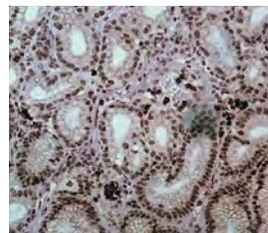
miRNA-378 promotes cell survival and angiogenesis by targeting SuFu and Fus-1 expression. The fluorescinated hsa-miR-378a probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-379

Hsa-miR-379 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM379-100 (ASR*)
 Specificity: miR-379
 Recommended Barrier: FB-HM379
 Control:

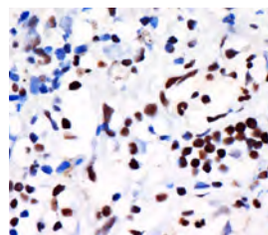
miR-379, is located on chromosome 14q32, 31. In the context of tumor, miR-379 regulates interleukin-11 (IL-11) production and decreased in tumor. The fluorescinated hsa-miR-379 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-381

Hsa-miR-381 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM381-100 (ASR*)
 Specificity: miR-381
 Recommended Barrier: FB-HM381
 Control:

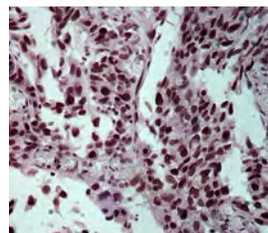
Recent functional studies have demonstrated that miR-381 serves as a tumor suppressor and is associated with radio-sensitivity in tumor cells. Overexpression of miRNA-381 confers increased radio-sensitivity of tumor cells, promotes nonaggressive phenotype, and growth inhibition. miRNA-381 exerts its biological functions through the regulation of various target genes, such as MITF, LRRC4, ID1, MDR1, BRD7, and WEE1. The fluorescinated hsa-miR-381 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-382

Hsa-miR-382 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM382-100
 Specificity: miR-382
 Recommended Barrier: FB-HM382
 Control:

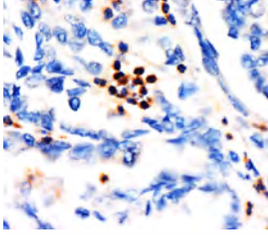
miR-382 has been found to have a decreased expression and the ability to suppress tumorigenesis in colorectal cancer and lung cancer. Moreover, the expression levels of miR-382 is purported to be associated with last-stage and tumor metastasis in NSCLC patients. The fluorescinated hsa-miR-382 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-383

Hsa-miR-383 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM383-100 (ASR*)
 Specificity: miR-383
 Recommended Barrier: FB-HM383
 Control:

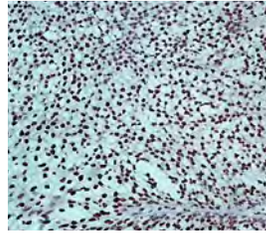
Downregulation of miR-383 promotes tumor cell invasion by targeting IGF1R. miR-383 promoted the expression of miR-320 and enhanced miR-320-mediated suppression of granulosa cell (GC) proliferation. miR-383 was up-regulated in the follicular fluid of polycystic ovarian syndrome (PCOS) patients, while the expression of E2F1 and SF-1 was down-regulated in GCs. The fluorescinated hsa-miR-383 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-384

Hsa-miR-384 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM384-100
 Specificity: miR-384
 Recommended Barrier: FB-HM384
 Control:

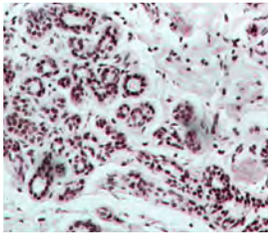
miR-384 is a brain-enriched miRNA, highly expressed in hippocampus and downregulated in glioma tissues and glioma cell lines. The fluorescinated hsa-miR-384 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-412

Hsa-miR-412 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM412-100 (ASR*)
 Specificity: miR-412
 Recommended Barrier: FB-HM412
 Control:

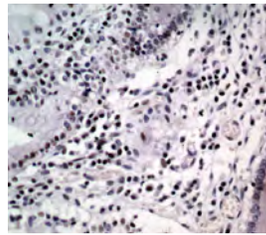
miR-412 was observed to be upregulated in the tumor tissues compared with normal tissues. mRNA bound to the AGO2 complex (RIP-Chip) identified a set of miR-412 target genes that are involved in neuronal cell death processes. The fluorescinated hsa-miR-412 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-409-3p

Hsa-miR-409-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM409-3P-100 (ASR*)
 Specificity: miR-409-3p
 Recommended Barrier: FB-HM409-3P
 Control:

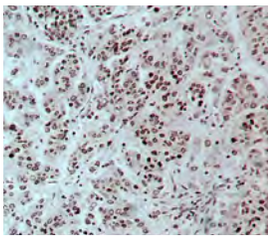
miR-409-3p was significantly downregulated in tumor cell lines and tissues. Overexpression of miR-409-3p in SGC-7901 tumor cells dramatically suppressed cell proliferation and induced cell apoptosis both *in vitro* and *in vivo*. The transcriptional regulator PHF10 was a target of miR-409-3p. The fluorescinated hsa-miR-409-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-422a

Hsa-miR-422a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM422A-100 (ASR*)
 Specificity: miR-422a
 Recommended Barrier: FB-HM422A
 Control:

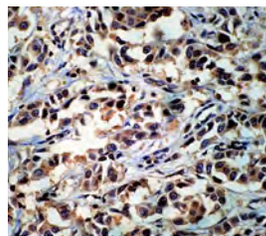
miR-422a plays a protective role in tumors where significantly reduced expression has been observed in tumors when compared to the normal tissue counterparts. miR-422a also inhibits pathways that stimulate tumor cell proliferation. The fluorescinated hsa-miR-422a probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-410

Hsa-miR-410 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM410-100 (ASR*)
 Specificity: miR-410
 Recommended Barrier: FB-HM410
 Control:

miR-410 was significantly downregulated in the tumor tissue. The expression of miR-410 was inversely associated with MET in human tumor tissues. Restoring expression of miR-410 led to proliferation inhibition and reduced invasive capability. miR-410 plays an important role in regulating MET-induced AKT signal transduction. The fluorescinated hsa-miR-410 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

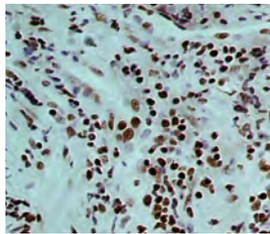
Hsa-miR-423-3p

Hsa-miR-423-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM423-3P-100 (ASR*)
 Specificity: miR-423-3p
 Recommended Barrier: FB-HM423-3P
 Control:

miR-423 is located on chromosome 17 and lies within the first intron of the gene nuclear speckle splicing regulatory protein (NSRP1) which is involved in alternate splicing of mRNAs. The fluorescinated hsa-miR-423-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

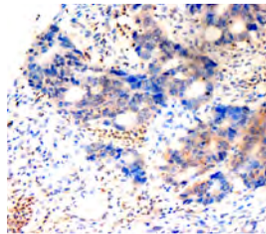
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-424

Hsa-miR-424 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM424-100 (ASR*)
 Specificity: miR-424
 Recommended Barrier: FB-HM424
 Control:

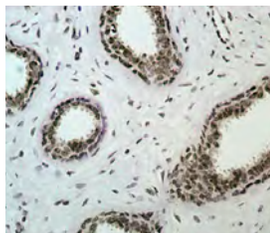
Hypoxia induces miR-424 expression and that miR-424 in turn suppresses the level of PDCD4 protein, a tumor suppressor. The inhibition of miR-424 enhanced apoptosis and increased the sensitivity of tumor cells. miR-424 levels are inversely correlated with PDCD4 expression in clinical tumor samples. The fluorescinated hsa-miR-424 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-433

Hsa-miR-433 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM433-100
 Specificity: miR-433
 Recommended Barrier: FB-HM433
 Control:

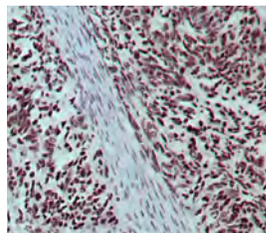
miR-433 has been reported to be dysregulated in several malignancies, including ovarian cancer, liver cancer and colorectal cancer. miR-433 is also highly expressed in brain, variation in the miRNA-433 binding site of FGF20 confers risk for Parkinson disease. The fluorescinated hsa-miR-433 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-425

Hsa-miR-425 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM425-100 (ASR*)
 Specificity: miR-425
 Recommended Barrier: FB-HM425
 Control:

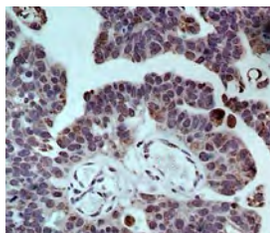
miR-425 has been identified as a potential biomarker in many types of tumors. miR-425 has been reported to promote tumorigenicity and aggressiveness in tumors. The fluorescinated hsa-miR-425 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-449a

Hsa-miR-449a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM449A-100 (ASR*)
 Specificity: miR-449a
 Recommended Barrier: FB-HM449A
 Control:

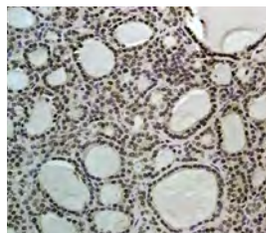
miR-449a is downregulated in human tumor tissue and possesses potential tumor suppressor function. miR-449a-mediated growth arrest in tumor cells is dependent on the Rb protein. The fluorescinated hsa-miR-449a probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-429

Hsa-miR-429 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM429-100 (ASR*)
 Specificity: miR-429
 Recommended Barrier: FB-HM429
 Control:

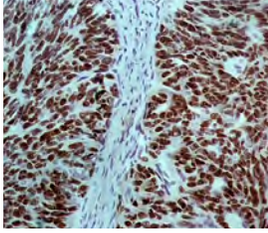
miR-429, a member of the miR-200 family of microRNAs, was significantly downregulated in tumor tissues and cell lines. miR-429 inhibited the proliferation and growth of tumor cells *in vitro* and *in vivo*. Downregulation of miR-429 may contribute to carcinogenesis and the initiation of epithelial-mesenchymal transition (EMT) by targeting Onecut2. The fluorescinated hsa-miR-429 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-450b-3p

Hsa-miR-450b-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM450B-3P-100 (ASR*)
 Specificity: miR-450b-3p
 Recommended Barrier: FB-HM450B-3P
 Control:

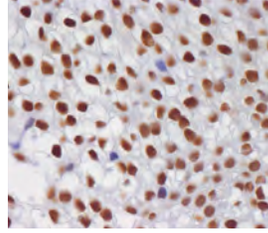
miR-450b-3p inhibits HER3 expression and represses the downstream signal transductions of HER family. Overexpression of miR-450b-3p inhibits tumor cells clonogenic potential and enhances their sensitivity to trastuzumab, a monoclonal antibody that binds to the HER2 receptor, or doxorubicin through repressing proliferative signal pathways mediated by HER3/HER2/PI3K/AKT. The fluorescinated hsa-miR-450b-3p probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-451

Hsa-miR-451 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM451-100 (ASR*)
 Specificity: miR-451
 Recommended Barrier: FB-HM451
 Control:

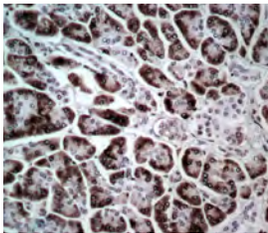
miR-451 gene is located on chromosome 17 at 17q11.2. miR-451 regulates the drug-transporter protein P-glycoprotein, potentially promoting resistance to the chemotherapy drug Paclitaxel. miRNA-451 is widely dysregulated in human tumors and plays a critical role in tumorigenesis and tumor progression. The fluoescinated hsa-miR-451 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-486-3p

Hsa-miR-486-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM486-3P-100 (ASR*)
 Specificity: miR-486-3p
 Recommended Barrier: FB-HM486-3P
 Control:

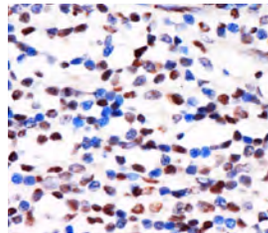
miR-486-3p dysregulation was observed in several tumors. Overexpression of miR-486-3p resulted in a moderate decrease of mature erythroid cells, indicating a possible inhibitory effect on erythropoiesis. The fluoescinated hsa-miR-486-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-483

Hsa-miR-483 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM483-100 (ASR*)
 Specificity: miR-483
 Recommended Barrier: FB-HM483
 Control:

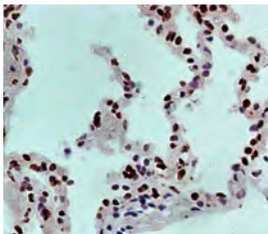
miR-483 is located within intron 2 of the IGF2 locus. The expression level of miR-483 alone can accurately diagnose a tumor as benign or malignant. miR-483 also highly expressed in colon, breast and liver tumor. The fluoescinated hsa-miR-483 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-489

Hsa-miR-489 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM489-100
 Specificity: miR-489
 Recommended Barrier: FB-HM489
 Control:

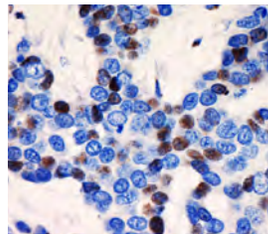
miR-489 has been reported to mediate chemoresistance in ovarian cancer and breast cancer. Akt3 and Smad3 could be the downstream target of miR-489. The fluoescinated hsa-miR-489 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-486

Hsa-miR-486 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM486-100 (ASR*)
 Specificity: miR-486
 Recommended Barrier: FB-HM486
 Control:

miR-486 plays a tumor-suppressor role. miR-486 is located at Chromosome 8p11, a region of frequent genomic loss in multiple tumors. miR-486 is significantly downregulated in tumor. miR-486 inactivation is required for the expression of several pro-oncogenic traits, and that this is likely mediated through miR-486 targeting the OLFM4 antiapoptotic factor. The fluoescinated hsa-miR-486 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

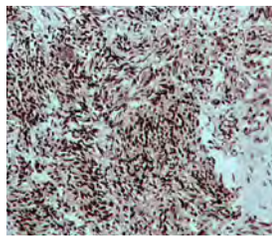
Hsa-miR-491

Hsa-miR-491 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM491-100
 Specificity: miR-491
 Recommended Barrier: FB-HM491
 Control:

miR-491-5p is located in the fourth intron of FOCAD, it has been reported to be involved in several cancer types. miR-491-5p can act as a tumor suppressor by targeting JMJD2B in breast cancer, or targeting TRIM28 in glioma. The fluoescinated hsa-miR-491 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

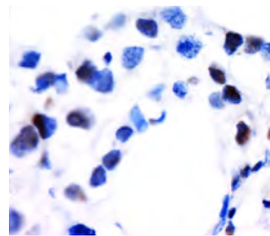
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-494

Hsa-miR-494 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM494-100 (ASR*)
 Specificity: miR-494
 Recommended Barrier: FB-HM494
 Control:

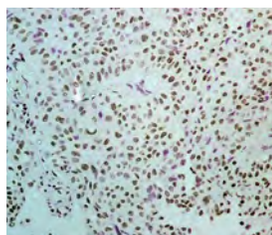
miR-494 regulates the expression of phosphatase and tensin homolog (PTEN) post-transcriptionally and functions as a micro-oncogene in carcinogenesis induced by anti-BPDE. The fluorescinated hsa-miR-494 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-498

Hsa-miR-498 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM498-100
 Specificity: miR-498
 Recommended Barrier: FB-HM498
 Control:

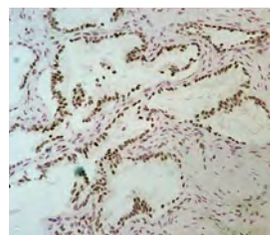
miR-498 is downregulated and correlated with non-small cell lung cancer progression, which might be a putative prognostic biomarker or therapeutic target in NSCLC treatment. The fluorescinated hsa-miR-498 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-495

Hsa-miR-495 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM495-100 (ASR*)
 Specificity: miR-495
 Recommended Barrier: FB-HM495
 Control:

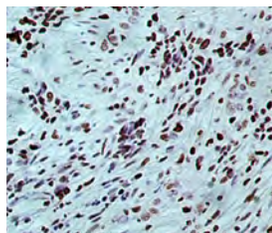
miR-495 was dramatically decreased in tumor cell lines and ectopic expression of miR-495 drastically retarded the proliferation and tumorigenicity in *in vitro* and *in vivo* assays, suggesting that downregulation of miR-495 may associate with features of tumor and that it functions as an antimir. The fluorescinated hsa-miR-495 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-502

Hsa-miR-502 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM502-100 (ASR*)
 Specificity: miR-502
 Recommended Barrier: FB-HM502
 Control:

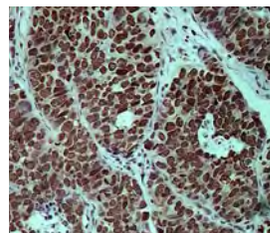
The expression of miR-502 was downregulated in tumor patient specimens compared with the paired normal control samples. The fluorescinated hsa-miR-502 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-497

Hsa-miR-497 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM497-100 (ASR*)
 Specificity: miR-497
 Recommended Barrier: FB-HM497
 Control:

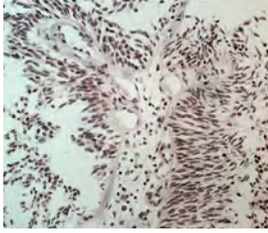
miR-497 locates at 17p13.1, and is frequently deleted in human tumors. miR-497 showed significant growth-suppressive activity with induction of G1 arrest. miR-497 overexpression led to the aberrant cell proliferation in hepatocarcinogenesis. The fluorescinated hsa-miR-497 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-505

Hsa-miR-505 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM505-100 (ASR*)
 Specificity: miR-505
 Recommended Barrier: FB-HM505
 Control:

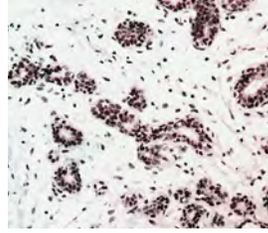
miR-505 functions as a tumor suppressive microRNA. FGF18, a proangiogenic factor, is directly regulated by miR-505. miR-505 inhibits cell proliferation by inducing apoptosis. The fluorescinated hsa-miR-505 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-508-3p

Hsa-miR-508-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM508-3P-100 (ASR*)
 Specificity: miR-508-3p
 Recommended Barrier: FB-HM508-3p
 Control:

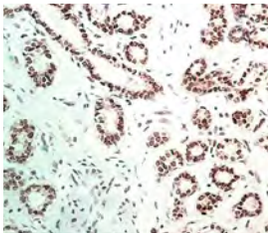
miR-508-3p (member of the miR-506 family) is located on Xq27.3, which is a fragile site of the human X chromosome. The very limited reports about miR-508-3p are controversial according to different tumor types. The fluorescinated hsa-miR-508-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-511

Hsa-miR-511 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM511-100 (ASR*)
 Specificity: miR-511
 Recommended Barrier: FB-HM511
 Control:

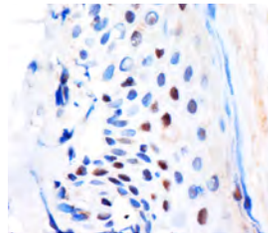
3'-UTRs of TLR4 I and TLR4 II were miR-511 target sites and that miR-511 knockdown enhanced TLR4 protein levels in differentiating dendritic cells. Downregulation of miR-511 expression was found in ovarian tumor tissues. The fluorescinated hsa-miR-511 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-509-3p

Hsa-miR-509-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM509-3P-100 (ASR*)
 Specificity: miR-509-3p
 Recommended Barrier: FB-HM509-3P
 Control:

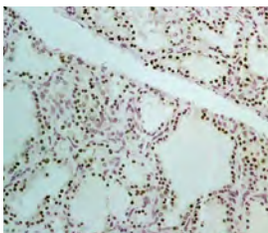
It was reported that miR-509-3p may function as a tumor suppressor. The expression level of miR-509-3p is lower in tumor than in the adjacent normal tissues and ectopic expression of miR-509-3p inhibits renal cell growth and migration. The fluorescinated hsa-miR-509-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-514a

Hsa-miR-514a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM514a-100
 Specificity: miR-514a
 Recommended Barrier: FB-HM514a
 Control:

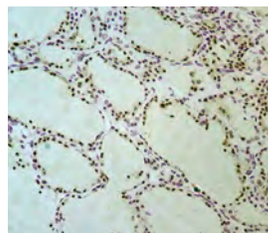
miR-514a is a member of a cluster of miRNAs on chrXq27.3 that has been implicated in the malignant transformation of melanocytes and tumor progression. However, in ovarian carcinoma, this miRNA cluster has been demonstrated as a tumor suppressor. The fluorescinated hsa-miR-514a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-510

Hsa-miR-510 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM510-100 (ASR*)
 Specificity: miR-510
 Recommended Barrier: FB-HM510
 Control:

miR-510, is elevated in tumor samples while absent in the matched non-tumor tissue samples. The fluorescinated hsa-miR-510 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-517a-3p

Hsa-miR-517a-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM517A-3P-100 (ASR*)
 Specificity: miR-517a-3p
 Recommended Barrier: FB-HM517A-3P
 Control:

miR-517a-3p was differentially expressed in tumor 95D and 95C cell lines that have different metastatic potential. Manipulation of miR-517a-3p expression changed tumor cell proliferation, migration and invasion capacity. MiR-517a-3p directly regulated FOXJ3 expression by binding to FOXJ3 promoter. The fluorescinated hsa-miR-517a-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

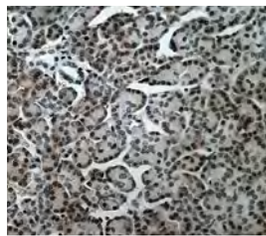
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-520C

Hsa-miR-520c detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM520C-100 (ASR*)
 Specificity: miR-520c
 Recommended Barrier: FB-HM520C
 Control:

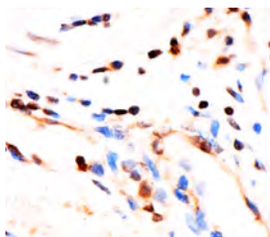
miR-520c is an important miRNA and has been characterized as oncogenes. In tumor cells, miR-520c stimulated tumor cell migration and invasion by suppressing the expression of CD44. The fluorescinated hsa-miR-520c probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-541

Hsa-miR-541 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM541-100 (ASR*)
 Specificity: miR-541
 Recommended Barrier: FB-HM541
 Control:

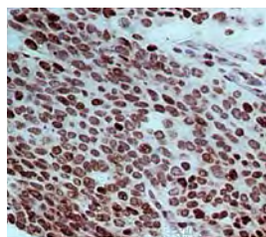
miR-541 directly regulates HER2 expression in tumor. The fluorescinated hsa-miR-541 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-524

Hsa-miR-524 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM524-100
 Specificity: miR-524
 Recommended Barrier: FB-HM524
 Control:

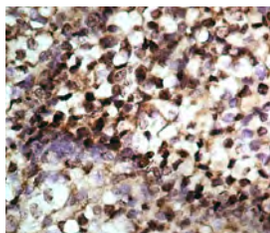
miR-524 targets both BRAF and ERK2 genes, the key regulators of the MAPK pathway, and affect melanoma cell migration and proliferation. miR-524 is also a brain-enriched miRNA, which is associated with the pathological grade and overall survival of gliomas. The fluorescinated hsa-miR-524 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-544

Hsa-miR-544 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM544-100 (ASR*)
 Specificity: miR-544
 Recommended Barrier: FB-HM544
 Control:

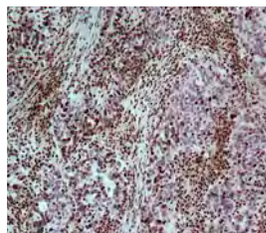
miR-544 exhibited a progression-associated downregulation in tumors. The fluorescinated hsa-miR-544 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-532-5p

Hsa-miR-532-5p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM532-5P-100 (ASR*)
 Specificity: miR-532-5p
 Recommended Barrier: FB-HM532-5P
 Control:

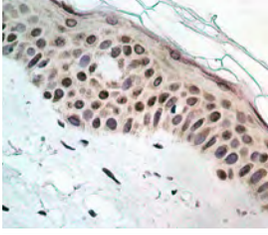
miR-532-5p was differentially expressed in tumor 95D and 95C cell lines that have different metastatic potential. Manipulation of miR-532-5p expression changed tumor cell proliferation, migration and invasion capacity. MiR-532-5p directly regulated FOXJ3 expression by binding to FOXJ3 promoter. The fluorescinated hsa-miR-532-5p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-545-5p

Hsa-miR-545-5p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM545-5p-100 (ASR*)
 Specificity: miR-545-5p
 Recommended Barrier: FB-HM545-5P
 Control:

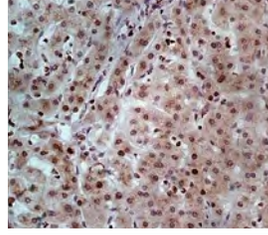
Low miR-545 levels in tumors promote tumor cells growth, and are associated with reduced survival in patients. miR-545 inhibits the proliferation of tumor cells both in vitro and in vivo. The fluorescinated hsa-miR-545-5p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-573

Hsa-miR-573 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM573-100 (ASR*)
 Specificity: miR-573
 Recommended Barrier: FB-HM573
 Control:

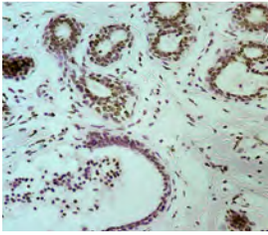
miR-573 has been reported to act as a tumor suppressor gene. The fluorescinated hsa-miR-573 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-610

Hsa-miR-610 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM610-100(ASR*)
 Specificity: miR-610
 Recommended Barrier: FB-HM610
 Control:

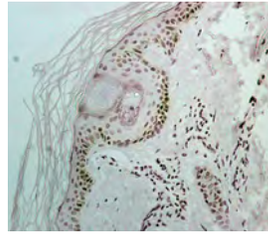
miR-610 which were downregulated in tumor and may be exploited for therapeutic intervention to inhibit tumor progression and metastasis. miR-610 suppresses tumor cell proliferation. The fluorescinated hsa-miR-610 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-574-3p

Hsa-miR-574-3p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM574-3P-100 (ASR*)
 Specificity: miR-574-3p
 Recommended Barrier: FB-HM574-3p
 Control:

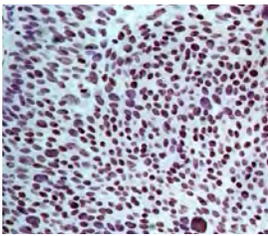
miR-574-3p was downregulated in clinical tumor tissues, and knockdown of endogenous miR-574-3p abrogated the tamoxifen-mediated growth suppression of MCF-7 cells. The fluorescinated hsa-miR-574-3p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-614

Hsa-miR-614 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM614-100 (ASR*)
 Specificity: miR-614
 Recommended Barrier: FB-HM614
 Control:

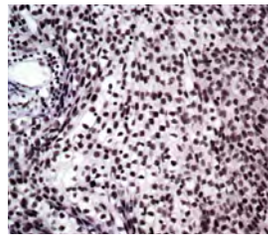
miR-614 inhibited tumor cells invasion and proliferation. The fluorescinated hsa-miR-614 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-590

Hsa-miR-590 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM590-100 (ASR*)
 Specificity: miR-590
 Recommended Barrier: FB-HM590
 Control:

Downregulation of miR-590 by nicotine has been found to play a key part in the generation of atrial fibrosis by atrial structural remodeling. Expression of miR-590 was downregulated in a number of tumor cell lines. The down-regulation of miR-590-5P may result in the dysregulation of its target genes. The fluorescinated hsa-miR-590 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

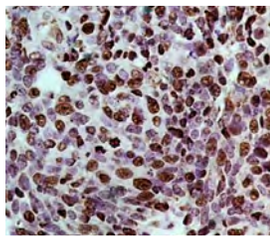
Hsa-miR-615

Hsa-miR-615 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM615-100 (ASR*)
 Specificity: miR-615
 Recommended Barrier: FB-HM615
 Control:

Expression of microRNA miR-615 is reported in various tumors. The ectopic expression of miR-615 reduced the cell growth and migration. Expression of miR-615 is epigenetically activated by DNA methylation in tumor cells. The fluorescinated hsa-miR-615 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

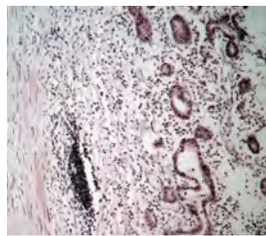
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-622

Hsa-miR-622 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM622-100 (ASR*)
 Specificity: miR-622
 Recommended Barrier: FB-HM622
 Control:

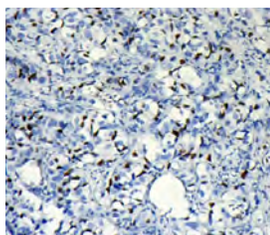
Expression of miR-622 is downregulated in tumors. Ectopic expression of miR-622 promotes invasion, tumorigenesis and metastasis of tumor cells both in vitro and in vivo. The fluorescinated hsa-miR-622 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-628

Hsa-miR-628 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM628-100 (ASR*)
 Specificity: miR-628
 Recommended Barrier: FB-HM628
 Control:

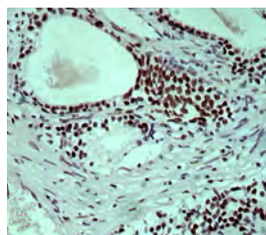
miR-628 was significantly downregulated in tumors when compared with normal ones. The fluorescinated hsa-miR-628 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-625

Hsa-miR-625 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM625-100 (ASR*)
 Specificity: miR-625
 Recommended Barrier: FB-HM625
 Control:

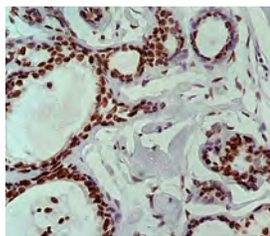
miR-625 has been shown to be downregulated in tumors. miR-625 is responsible for the regulation of metastasis in tumor cells, and therefore downregulation of miR-625 results in increased metastasis. The fluorescinated hsa-miR-625 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-629

Hsa-miR-629 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM629-100 (ASR*)
 Specificity: miR-629
 Recommended Barrier: FB-HM629
 Control:

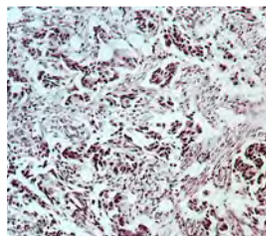
miR-629 is upregulated in many tumor tissues. miR-629 activates IL-6–JAK–STAT3 signaling in tumor cells, which in turn upregulates miR-629 expression. The fluorescinated hsa-miR-629 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-627

Hsa-miR-627 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM627-100 (ASR*)
 Specificity: miR-627
 Recommended Barrier: FB-HM627
 Control:

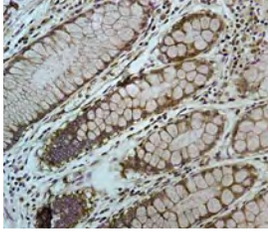
miR-627 is a major epigenetic regulator in vitamin D induced growth inhibition of tumorous cells upon stimulation by calcitriol. miR-627 acts on target gene JMJD1A (jumonji domain containing 1A), the gene encoding a histone demethylase which is upregulated under hypoxia and promotes tumor growth in tumor cells. Overexpression of miR-627 decreased JMJD1A and suppressed the expression of growth-promoting and differentiating genes, GDF15. The fluorescinated hsa-miR-627 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-630

Hsa-miR-630 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM630-100 (ASR*)
 Specificity: miR-630
 Recommended Barrier: FB-HM630
 Control:

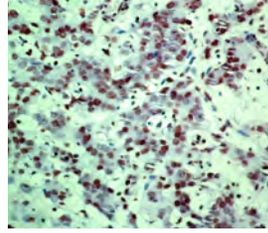
miR-630 has recently been identified to be implicated in many critical processes in human malignancies. miR-630 expression was significantly increased in tumor specimens compared with that in adjacent normal specimens. The fluorescinated hsa-miR-630 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-638

Hsa-miR-638 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM638-100 (ASR*)
 Specificity: miR-638
 Recommended Barrier: FB-HM638
 Control:

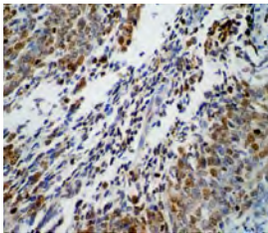
miR-638 has been reported to be downregulated in several types of tumor, and may therefore function as a tumor suppressor gene. The fluorescinated hsa-miR-638 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-648

Hsa-miR-648 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM648-100 (ASR*)
 Specificity: miR-648
 Recommended Barrier: FB-HM648
 Control:

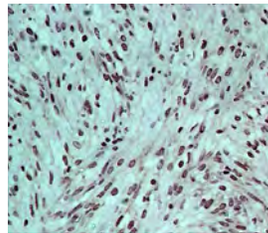
The miR-648 gene is present in the first intron of MICAL3, encoding a member of the microtubule associated monooxygenase, calponin, and LIM domain-containing (MICAL) family of flavoprotein monooxygenases, which participate in axon guidance, actin remodeling, and redox activity in promoting vesicle-docking complexes in the process of exocytosis. The fluorescinated hsa-miR-648 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-641

Hsa-miR-641 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM641-100 (ASR*)
 Specificity: miR-641
 Recommended Barrier: FB-HM641
 Control:

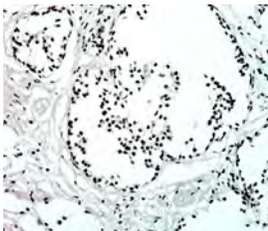
miR-641 is an uncharacterized microRNA located at intron-1 of the AKT2 gene and is reported to co-regulate and cooperate with AKT. The fluorescinated hsa-miR-641 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-650

Hsa-miR-650 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HHM0650-100 (ASR*)
 Specificity: miR-650
 Recommended Barrier: FB-HM650
 Control:

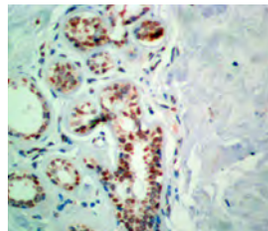
miR-650 is involved in lymphatic and distant metastasis in human tumors. The ectopic expression of miR-650 promotes tumorigenesis and proliferation of tumor cells. The fluorescinated hsa-miR-650 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-642a-5p

Hsa-miR-642a-5p detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM642A-5P-100 (ASR*)
 Specificity: miR-642a-5p
 Recommended Barrier: FB-HM642A-5P
 Control:

miR-642a-5p targets Toll-like Receptor 4 in monocytes. The fluorescinated hsa-miR-642a-5p probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

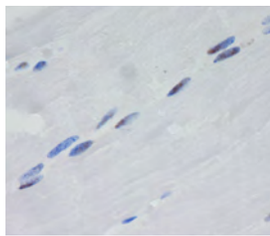
Hsa-miR-663a

Hsa-miR-663a detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM663A-100 (ASR*)
 Specificity: miR-663a
 Recommended Barrier: FB-HM663A
 Control:

miR-663 may be a potential tumor suppressor. The fluorescinated hsa-miR-663a probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

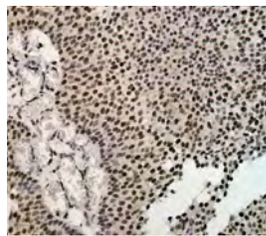
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-675

Hsa-miR-675 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM675-100
 Specificity: miR-675
 Recommended Barrier: FB-HM675
 Control:

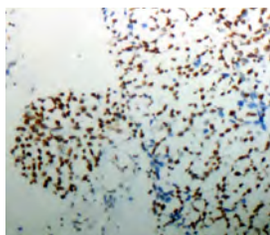
miR-675 is embedded in H19's first exon and expressed in the placenta from the gestational time point when placental growth normally ceases. miR-675 has an essential function in skeletal muscle differentiation and regeneration by targeting BMP pathway and Cdc6, a DNA replication initiation factor. The fluorescinated hsa-miR-675 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-765

Hsa-miR-765 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM765-100 (ASR*)
 Specificity: miR-765
 Recommended Barrier: FB-HM765
 Control:

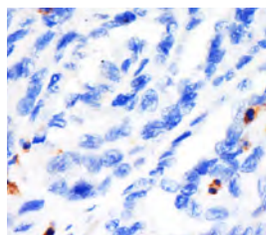
miR-765 is a fulvestrant-induced and ER β -associated miRNA, and it targets an oncogenic protein HMGA1. The fluorescinated hsa-miR-765 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-708

Hsa-miR-708 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM708-100 (ASR*)
 Specificity: miR-708
 Recommended Barrier: FB-HM708
 Control:

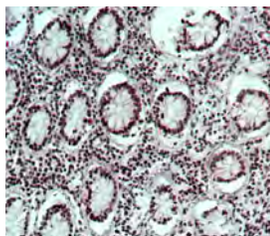
miR-708 is located on chromosome 11q14.1 and is encoded in intron 1 of the ODZ4 gene. It is highly expressed in the brain and eyes. High miR-708 expression levels are observed in tumors due to their oncogenic role in tumor growth and progression. miR-708 overexpression results in increased cell proliferation, migration, and invasion. The fluorescinated hsa-miR-708 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-766

Hsa-miR-766 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM766-100
 Specificity: miR-766
 Recommended Barrier: FB-HM766
 Control:

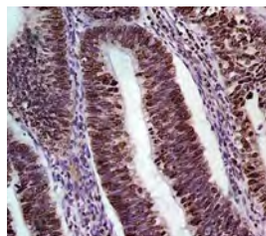
Growing evidence indicates that miR-766 acts as a tumor promoter or suppressor in multiple cancers, including cutaneous carcinoma, lung adenocarcinoma, colorectal cancer and renal cell carcinoma. The fluorescinated hsa-miR-766 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-718

Hsa-miR-718 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM718-100 (ASR*)
 Specificity: miR-718
 Recommended Barrier: FB-HM718
 Control:

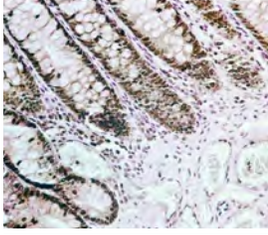
Decreased expression of miR-718 was associated with tumor aggressiveness. The fluorescinated hsa-miR-718 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-802

Hsa-miR-802 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM802-100 (ASR*)
 Specificity: miR-802
 Recommended Barrier: FB-HM802
 Control:

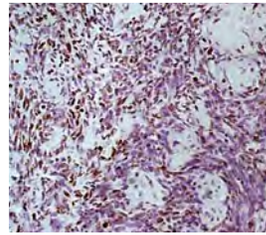
Enriched expression of miR-802 promoted cell proliferation in tumor cells by negatively targeting cell cycle inhibitor p27 protein as against the normal tissues. The fluorescinated hsa-miR-802 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-874

Hsa-miR-874 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM874-100 (ASR*)
 Specificity: miR-874
 Recommended Barrier: FB-HM874
 Control:

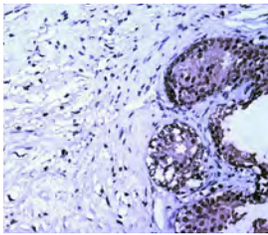
miR-874 has been identified as a tumor-suppressor and is reportedly down-regulated in some types of tumor. The fluorescinated hsa-miR-874 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization

Hsa-miR-1181

Hsa-miR-1181 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM1181-100 (ASR*)
 Specificity: miR-1181
 Recommended Barrier: FB-HM1181
 Control:

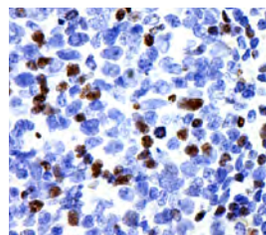
Recently, it has been shown that overexpression of miR-1181 inhibited, whereas down-regulation of miR-1181 promoted, tumor stem cells (CSCs)-like phenotypes in vitro and tumorigenicity in vivo. This indicated that downregulated or low expression of miR-1181 is associated with poor overall survival and disease-free survival of the tumor patients. The fluorescinated hsa-miR-1181 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-940

Hsa-miR-940 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM940-100 (ASR*)
 Specificity: miR-940
 Recommended Barrier: FB-HM940
 Control:

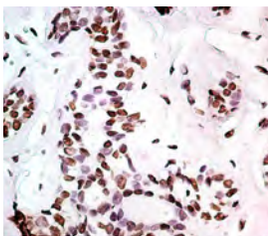
The dysregulation of miR-940 has been found in various tumors. miR-940 was highly expressed in normal tissues compared with tumors, and miR-940 inhibited migratory and invasive potential of tumor cells. miR-940 promotes tumor cell invasion and metastasis by downregulating ZNF24. The fluorescinated hsa-miR-940 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-1244-1

Hsa-miR-1244-1 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM1244-1-100
 Specificity: miR-1244-1
 Recommended Barrier: FB-HM1244-1
 Control:

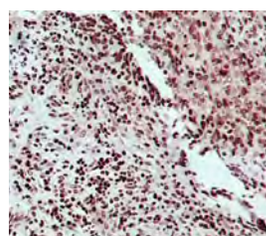
miR-1244 acts as a tumor suppressor in lung cancer by reducing its proliferation, survival and invasion, and its under-expression is highly associated with patients' survival. Recent studies also suggest that miR-1244 is associated with progression of prostate cancer cells to antiandrogen therapy resistance. The fluorescinated hsa-miR-1244-1 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-944

Hsa-miR-944 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM944-100 (ASR*)
 Specificity: miR-944
 Recommended Barrier: FB-HM944
 Control:

miR-944 expression has been detected in several tumor types, and is more abundant in tumor samples than in their normal counterparts. High expression of miR-944 is also associated with tumor recurrence. The fluorescinated hsa-miR-944 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

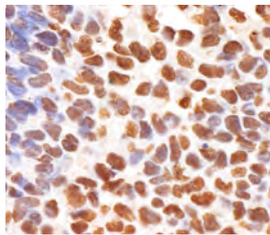
Hsa-miR-1247

Hsa-miR-1247 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM1247-100 (ASR*)
 Specificity: miR-1247
 Recommended Barrier: FB-HM1247
 Control:

Aberrant expression of miR-1247 has been found in several tumors and is predicted to play an important role in the pathological processes of tumor by miRNA-regulated network analysis. The fluorescinated hsa-miR-1247 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

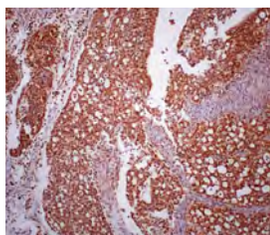
*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hsa-miR-1258

Hsa-miR-1258 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM1258-100 (ASR*)
 Specificity: miR-1258
 Recommended Barrier Control: FB-HM1258

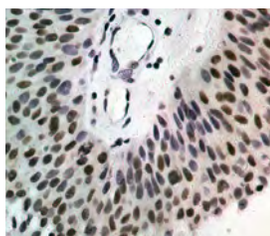
miR-1258 may play an important role in tumor development and progression by regulating the expression of HPSE. The fluorescinated hsa-miR-1258 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-1285

Hsa-miR-1285 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM1285-100 (ASR*)
 Specificity: miR-1285
 Recommended Barrier Control: FB-HM1285

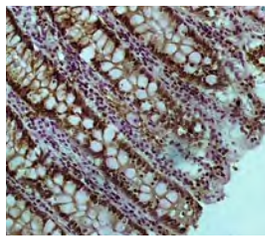
Genome-wide gene expression analysis data show that transglutaminase 2 (TGM2) is directly regulated by miR-1285. The fluorescinated hsa-miR-1285 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-1296

Hsa-miR-1296 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM1296-100 (ASR*)
 Specificity: miR-1296
 Recommended Barrier Control: FB-HM1296

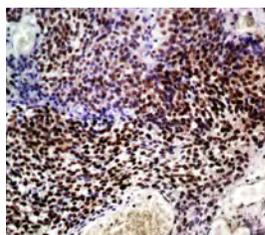
miR-1296 is downregulated in tumor and that MCM2 is one of its targets. The fluorescinated hsa-miR-1296 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-1297

Hsa-miR-1297 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM1297-100 (ASR*)
 Specificity: miR-1297
 Recommended Barrier Control: FB-HM1297

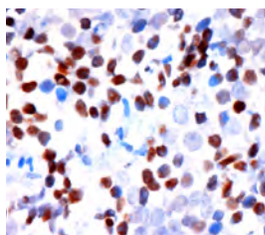
It has been reported that miR-1297 acts as a tumor suppressor by suppressing in vitro and in vivo expression of TRIB2/PTEN and further increasing C/EBP α expression thereby inhibits cell proliferation, migration, and tumorigenesis. miR-1297 inhibits the Cox-2/PGE-2 signaling pathway causing higher levels of miR-1297 in normal tissues than corresponding tumor tissues. The fluorescinated hsa-miR-1297 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-1826

Hsa-miR-1826 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM1826-100 (ASR*)
 Specificity: miR-1826
 Recommended Barrier Control: FB-HM1826

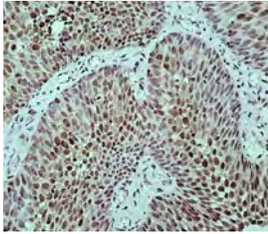
miR-1826 expression was significantly lower in tumor tissues and lower expression was significantly associated with overall shorter survival. miR-1826 also inhibited tumor cell proliferation, invasion and migration. miR-1826 plays an important role as a tumor suppressor by down-regulating beta-catenin and MEK1 in VHL inactivated tumors. The fluorescinated hsa-miR-1826 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Hsa-miR-3978

Hsa-miR-3978 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM3978-100
 Specificity: miR-3978
 Recommended Barrier Control: FB-HM3978

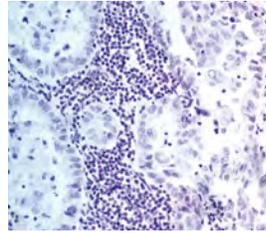
Differential expression of miR-3978 in lung cancer patients is observed⁵. Putative targets of miR-3978 have not been well defined. However, miR-3978 may target LGMN during metastatic progression of peritoneal gastric cancer patients. The fluorescinated hsa-miR-3978 probe is designed to localize this miRNA in FFPE tissue by *in situ* hybridization.

Hsa-miR-4723

Hsa-miR-4723 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM4723-100 (ASR*)
 Specificity: miR-4723
 Recommended Barrier: FB-HM4723
 Control:

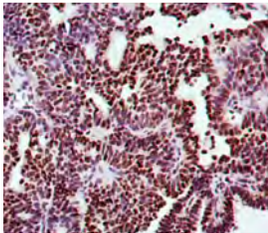
miR-4723 expression is attenuated in tumor and is significantly correlated with poor survival outcome and tumor progression. Functional studies using tumor cell lines showed that reconstitution of miR-4723 expression led to significant decreases in cell growth, invasion and migration. The fluorescinated hsa-miR-4723 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

Scramble

Negative staining of scramble probe in FFPE tissue

Ready-to-use (Manual): PR032-100 (ASR*)
 Specificity: Scramble
 Recommended Barrier: FB-PR032
 Control:

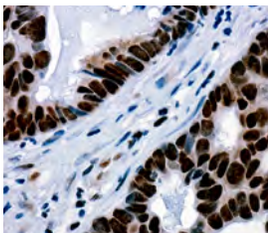
The scramble probe does not identify any miRNA sequences in human FFPE and freshly prepared frozen tissues. The probe sequence does not share homology with miRNA sequences available in the miRBase database.

Hsa-miR-9500

Hsa-miR-9500 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): HM9500-100 (ASR*)
 Specificity: miR-9500
 Recommended Barrier: FB-HM9500
 Control:

The expression levels of miR-9500 were reduced in tumor cells and tumor tissues compared with normal tissues. Overexpression of miR-9500 impeded cell migration in human tumor cells. The fluorescinated hsa-miR-9500 probe is designed to localize this miRNA in FFPE tissue by in situ hybridization.

U6

U6 detected in FFPE tissue stained with DAB

Ready-to-use (Manual): PR031-100 (ASR*)
 Specificity: U6
 Recommended Barrier: FB-PR031
 Control:

The U6 probe identifies a small nuclear RNA U6 sequence in human FFPE and freshly prepared frozen tissues. The probe sequence does not share homology with miRNA sequences available in the miRBase database.

*Analyte Specific Reagent. Analytical and performance characteristics are not established.

Hybridization Detection System

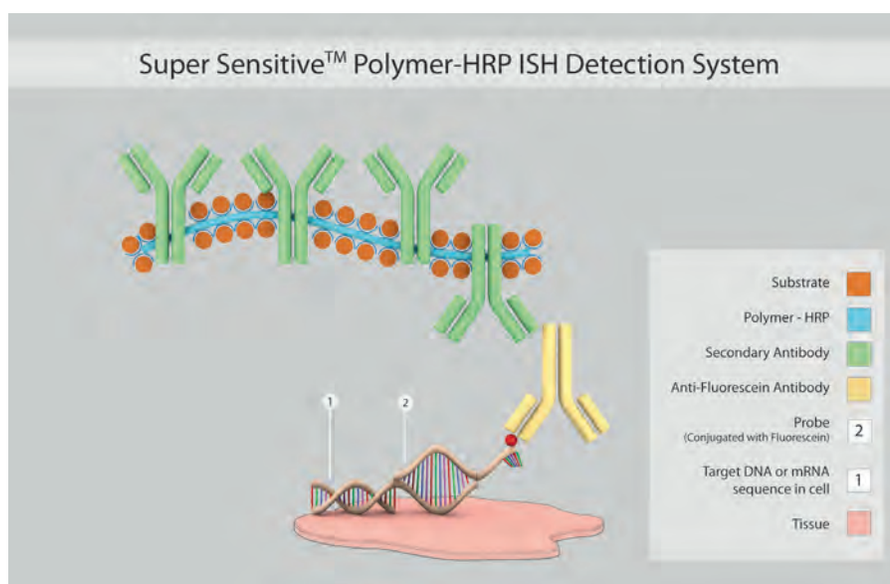
in situ Hybridization (ISH) is a powerful technique for detecting and localizing specific nucleic acid sequences within cells or tissues. This is achieved by the hybridization of a labeled probe to the specific RNA/DNA sequence within the cell and subsequent detection of the bound probe. ISH technique enables the semi-quantification of mRNA expression and helps determine the temporal and spatial patterns of gene expression in cells, tissue and whole animals. ISH technique can also be used for detection of intracellular pathogens with a very high degree of sensitivity.

Super Sensitive™ (Manual) & XISH (Xmatrx®) One-Step Polymer-HRP Detection System

This is a novel detection system using a non-biotin polymeric technology that makes use of Poly-HRP reagent. As the system is not based on the Biotin-Avidin System, problems associated with endogenous biotin are completely eliminated. The technology allows excellent cell penetration ability for intense staining, compared with other polymer HRP.

Features & Benefits:

- Clean Stain without endogenous biotin background
- High signal to noise ratio for intense stain
- Universal system for all fluorescein labeled probes
- Available in barcode tagged (XISH kit) for Automation or in dropper bottles (Super Sensitive™ kit) for manual staining



ISH Detection Systems Composition

SKU	Size	α Fluor.	Polymer HRP	DAB buffer	DAB Chromo.	Peroxide block	Power block	Hematox	Prot. K	Hybrid. buffer	NAR-1	Washes A,B,E,F
DF400-25K	25 test	2 mL	2 mL	5 mL	2 mL	3 mL	3 mL	3 mL	3 mL	6 mL	2 mL	10 mL
DF400-50KE	50 test	3 mL	3 mL	10 mL	2 mL	5 mL	5 mL	5 mL	5 mL	6 mL	3 mL	20 mL
DF400-YADE Xmatrx® Elite	100 test	5 mL	5 mL	4x5 mL + 5 barcoded vials	7 mL	10 mL	10 mL	10mL	5 mL	NA	5 mL	2x10 mL

Product	Size	Cat. No.	Description
NAR1	250 mL	HK873-5K	Microwave based nucleic acid retrieval for manual use only

Substrates and Chromogens

BioGenex offers complete Substrate Packs for immunohistochemical staining with alkaline phosphatase and peroxidase labels. The kits are designed to reduce substrate preparation time and minimize exposure to chemical hazards. The chart below summarizes the substrates offered, indicating enzyme and standard mounting media compatibility.

Features & Benefits:

- High Resolution AEC and Liquid DAB
- Rapid Development Time
- Ready-to-use(RTU) Solutions
- Long-Term Stability

The chart below summarizes the compatibility of mounting medium, chromogens and counterstains

Chromogen	Stain Color	Enzyme used	Solubility in Alcohol/Xylene	Compatible with Hematoxylin	Compatible Mounting Media
AEC	Brick Red	HRP	Yes	Yes	Aqueous or Super Mount
DAB	Brown	HRP	No	Yes	Aqueous, Super Mount or Xmount
Elegance Red	Red	AP	No	Yes	Aqueous, Super Mount or Xmount
Fast Red	Red	AP	Yes	Yes	Aqueous or Super Mount
New Fuchsin	Red	AP	Yes	Yes	Aqueous or Super Mount

ISH - Substrates and Chromogens Packs – Manual & Open system **

Product Name	60 Tests*	250 Tests*	500 Tests*/Large
Fast Red	NA	NA	HK182-5KE
Elegance Red	NA	NA	HK144-5KE
New Fuchsin (400 slides)	NA	NA	HK183-5KE
Two Component DAB (BUFFER+CHROMOGEN) (1000 slides)	NA	NA	HK542-XAKE
AEC (BUFFER+CHROMOGEN)	NA	HK092-5KE	HK092-YAKE
AEC (Concentrated BUFFER+CHROMOGEN)	NA	NA	HK129-YAKE
AEC One Step Sol.	HK139-06K	NA	HK139-50K

* 100 µL/test of prepared reagent

** Reagent vials for Xmatrix® need to be purchased separately



Automation



Automated Platforms for Molecular Pathology

BioGenex pioneers in the design, development and manufacturing of advanced systems for automation of cell- and tissue-based staining. To accommodate diverse laboratory needs we offer an array of clinical and research automation platforms that meet globally accepted quality standards (ISO13485:2003 & ISO9001:2008), are approved by the FDA and are especially designed to improve laboratory workflow, productivity and reproducibility.

Xmatrx® systems (NANO VIP, MINI, INFINITY, ELITE and ULTRA) are the direct result of our platform technology innovation. They offer a variety of automation, throughput and assay applications. Our key technology differentiators include the eXACT™ temperature control and reaction micro-chamber- improving IHC results and enabling Nucleic Acid-based Diagnostics (NADx).

1. Clinical platforms, support LIMS connectivity for data tracking and management, contain barcode enabled technologies and include over 400+ optimized protocols with ready to use reagents in barcode labeled vials (Xmatrx® vials). These systems are FDA approved for In Vitro Diagnostic (IVD) applications including: immuno-histochemistry (IHC), *in situ* hybridization (ISH), codetection and special staining.

Clinical Platforms /Application	IHC	ISH/CISH	Double Staining	Special Stains	IF
Xmatrx® ELITE	√	√	√	√	√
Xmatrx® ULTRA Dx	√	√	√	√	√
i6000™ Diagnostics	√	NA	√	√	√

2. Research platforms, offer infinity possibilities for translational and clinical research. They include flexible open system software for easily creating, editing and saving protocols and enable automation of any slide-based assay including immuno-histochemistry (IHC), *in situ* hybridization (ISH), fluorescence *in situ* hybridization (FISH), immuno-fluorescence (IF), co-detection and multiplex applications (double and triple stains; IHC/ISH), *in situ* PCR, micro-RNA and special staining.

Research Platforms /Application	IHC	ISH/CISH	Double Staining	Special Stains	FISH	IF	miRNA ISH	Multiplexing (ISH + IHC)	In Situ PCR
Xmatrx® Infinity	√	√	√	√	√	√	√	√	√
Xmatrx® ULTRA Rx	√	√	√	√	√	√	√	√	√
i6000™ Infinity	√	NA	√	√	NA	√	NA	NA	NA
NanoMtrx® 300	√	√	NA	NA	NA	NA	√	NA	NA
NanoMtrx® 100	√	√	NA	NA	NA	NA	√	NA	NA

3. Nucleic Acid Diagnostics (NAD) dedicated Platforms: Xmatrx® NANO VIP and MINI, are the most economical and flexible automation platforms for FISH, ISH and In-Situ Hybridization. These systems are small in size, contain 10 independent eXACT™ thermal cyclers that can run 10 different protocols simultaneously. These instruments contain on-board wash and waste drainage systems, audio-visual alerts and a user-friendly software with ability to add or delete cycles, store protocols for future use and perform, deparaffinization, antigen retrieval, hybridization, washing and up to 45 PCR cycles.

NAD Platforms /Application	ISH/CISH	FISH	miRNA ISH	In Situ PCR
NanoVIP®	√	√	√	√
Xmatrx® MINI	√	√	√	√

4. Other Systems: The i500 Plus is a LIMS enabled Barcode label printer for integrated digitized data tracking.

Other Systems	Description
EZ-Retriever™	Pre-treatment and antigen retrieval system using a programmable microwave oven with built-in temperature controls
i500™ Plus	LIMS enabled barcode label primer compatible with Xmatrx®

Clinical Platforms

Xmatrix[®]ELITE

Three Simple Steps



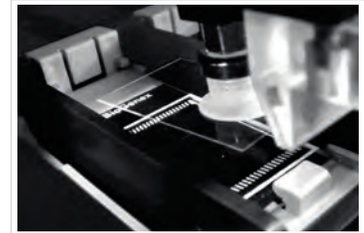
The most advanced fully automated system for IHC, ISH, SS Co-detection, and multiplexing

- 40 independent protocols simultaneously
- Fully automated, including baking, dewaxing & antigen retrieval
- eXACT™ Temperature Control on every slide (RT-105 °C)
- Bar-Coded reagent vials and slides to eliminates human errors
- Wide reagent dispense volumes: 10 µL to 850 µL
- BioGenex's proprietary coverslip mechanism
- Over 400+ optimized protocols with ready to use (RTU) reagents
- LIMS - enabled data tracking and management
- Liquid level sensor for accurate reagent dispensing
- System allows use of 3rd party antibodies

* optional software

Xmatrix® ULTRADx

Next Generation Fully Automated Staining System



All-in-One - IHC, ISH, SS and Co-detection

Fully Automated System from Microtome to Microscope... For the Molecular Pathology Laboratory of Present, Future and Beyond

- Next generation fully-automated slide staining system with Baking, Dewaxing & Antigen Retrieval
- Auto-DAB enabled – On-board automated mixing of chromogen and buffer
- 40 independent protocols simultaneously
- Bar-Coded reagent vials and slides to eliminates human errors
- eXACT™ temperature control on every slide (RT-105 °C)
- Wide reagent dispense volumes: 10 µL to 850 µL
- Auto drain disposal system
- Liquid level sensor for accurate reagent dispensing
- BioGenex's proprietary coverslip mechanism
- LIMS - enabled data tracking and management
- High throughput - 100 slides per day, 60 slides in eight-hour shift, and 40 slides in delayed overnight run
- Over 400+ optimized protocols with ready-to-use(RTU) reagents in barcoded vials
- Intuitive software designed for ease of use and flexibility
- System allows use of 3rd party antibodies
- Multiple slide processing options – Random, Continuous and STAT
- Work Flow status indicator

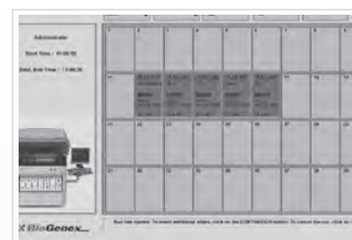
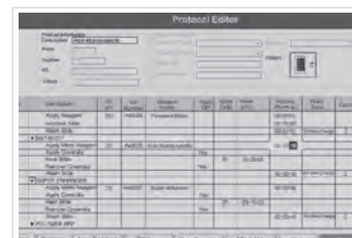
*Expected release: 2020

Research Platforms

Xmatrix[®] Infinity

Infinite Possibilities...

...For Translational and Clinical Research



All-in-One - IHC, IF, ISH, CISH, FISH, SS, *in situ* PCR and miRNA...

- Intelligent and flexible system offering infinite possibilities – IHC, ISH, FISH, SS, CISH, IF, Multiplexing and Co-detection
- Simultaneous optimization of up to 40 parameters in single run
- Reaction micro-chamber reduces micro-reagent consumption by up to 90%
- 40 independent thermocyclable (PCR) workstations
- Intuitive software designed for ease of use and flexibility
- Reports for inventory management and regulatory compliance
- Multiple slide processing options – Random, Continuous and STAT

NanoMtrx[®]300



State-of-the-art • Fully Automated All-In-One IHC, ISH, and Special Stains

- Compact 30 slide benchtop system
- 30 slides under 2.5 hours
- Generates 70% less waste
- Separates hazardous waste
- Uses standard slides
- Easy set-up and low maintenance

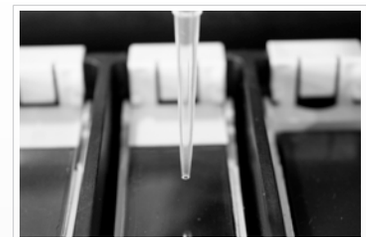
NanoMtrx[®]100



State-of-the-art • Fully Automated IHC and ISH System

- Multi-format specimen processing: FFPE, frozen, cell preparations, smears, and FNAs
- Micro-chamber[®] for uniform staining throughout the slide
- Temperature controlled micro-chambers[®] for minimal reagent consumption
- Gentle wash and blow-dry to eliminate tissue lift-off
- On-board auto-DAB mixing
- Generates 70% less waste
- Fast turnaround time of 2 hours with simultaneous 10 slide processing
- Intuitive user-friendly GUI

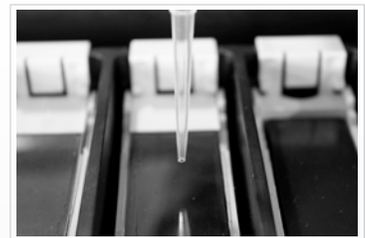
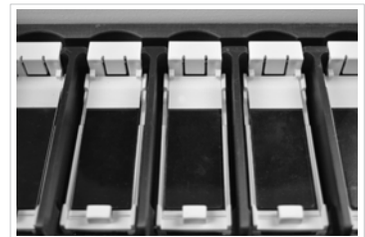
Nucleic Acid Diagnostic (NAD) Platforms

NanoVIP[®]*eFISHiency System for FISH Automation***All-in-One - ISH, FISH, miRNA ISH and IHC**

- Next generation fully-automated slide staining system
- Economical and affordable
- Flexible Open System Software - create, edit and save protocols for future use
- Simultaneous Optimization of 10 different protocols at the same time
- eXACT™ Temperature Control on every slide (RT-105 °C)
- Wide reagent dispense volumes: 10 µL to 850 µL
- Liquid level sensor for accurate reagent dispensing
- BioGenex's proprietary coverslip mechanism
- Intuitive software designed for ease of use and flexibility

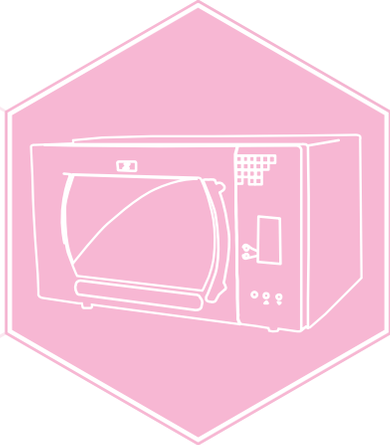
Xmatrix[®] MINI

eFISHiency Workstation



All-in-One - FISH, *in situ* PCR and ISH

- High performance *in situ* PCR and FISH
- Hybridizer with eXACT™ temperature controls
- 10 independent thermal cyclers
- Built-in touch screen display for easy operations
- Facility of on-board wash with effective waste drainage system
- Audio-visual alerts and on screen color-coded error alerts
- User-friendly software with ability to add/delete cycles, store protocols for future use and perform up to 45 PCR cycles



Tissue Pre-treatment & Nucleic Acid Retrieval 

De-Waxing Solutions

One-Step DeWaxing and Rehydration Reagent

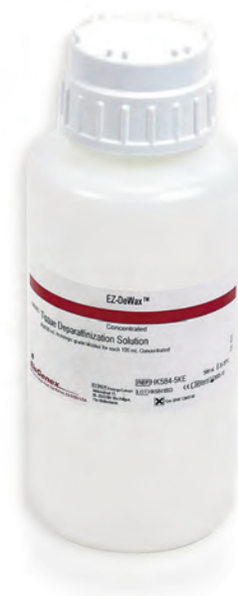
BioGenex deparaffinization solutions are “one-step” products that simultaneously enables the removal of paraffin and allows rehydration of the tissue with a single reagent. In the past, formalin-fixed, paraffin-embedded tissue sections were traditionally deparaffinized with highly toxic, noxious chemicals (i.e.xylene, xylene equivalents). BioGenex, a pioneer in the Immunohistochemistry technology, offers xylene-free products that removes the paraffin from mounted tissue slides easily and rapidly.

1. EZ-DeWax™ Sol. – For all BioGenex manual methods.
2. X-DeWax™ Sol. – Optimized for Xmatrix® automation.

Features & Benefits

- Effectively removes paraffin and allow rehydration of the tissue in one step.
- Reduces deparaffinization time from 45 minutes to 10 minutes.
- Eliminates use of toxic solvents (xylene) and minimizes hazardous waste.
- Ready-to-use(RTU) or 2x solutions (to be diluted 1:1 with ethanol) are available.

Product	1000 mL ^(RTU)	500 mL ^(2x)	1 Gallon ^(2x)
X-DeWax (Xmatrix®)	HX015-XAK	HX016-XAK	HX016-XEK
EZ-DeWax (Manual)	HK585-5k	HK584-5k	NA



Nucleic Acid Retrieval Method

BioGenex is the inventor of Nucleic Acid Retrieval enabling technology. This technology is an effective way of unmasking DNA in formalin-fixed, paraffin-embedded tissue sections using microwave heating. The Nucleic Acid Retrieval technique breaks the formalin induced cross-linking bonds between DNA and proteins, as well as protein-protein cross-linking thereby allowing better penetration of probes and accessibility of DNA for binding. Nucleic Acid Retrieval (NAR-1) is recommended instead of proteinase K when DNA targeting probes are used.

Advantages of the method:

- Reduces time for probe incubation
- Consistent and reliable staining quality
- Eliminates false-negative staining results
- Easy to use - Can be used in both microwave or Xmatrix® Automation protocols
- Non-hazardous, non-flammable and odorless – Safe and Eco-friendly

Product	Method	Features & Recommended Use
NAR-1	Microwave, 95-100 °c	Excellent for DNA targeting probes

Enzymes for Tissue Digestion

Some tissues require the use of enzymatic pre-treatment before staining to achieve standardized results depending on the antibodies and their different incubation and pre-treatment requirements. Each kit contains three or four vials of lyophilized enzyme powder and 15 mL of reconstitution buffer, enabling you to make fresh enzyme solutions as needed.

1. Proteinase K in a ready-to-use(RTU), RNase-free solution and is recommended for use with RNA targeting probes.
2. The Trypsin and Pepsin kits contain well-established enzymes suitable for routine pre-treatment at 37 °C. Pepsin is recommended as pretreatment for FISH applications.
3. Protease XXIV kits contain a universal digestive agent that allows for fast and effective pre-treatment at room temperature.

i500 Plus™

LIS Enabled Barcode Label Printer

Integrated Digitized Data Tracking System

- For printing chemical resistant barcode labels
- Compatible with Xmatrix®
- User-friendly software
- Synchronization of protocol information
- Efficient system
 - Eliminates human error
 - Helps reduce operating cost
 - Fast turn-around



EZ-Retriever® System

Pre-treatment and Antigen Retrieval System

- DeWax, re-hydration and antigen retrieval in one step
- Optimized factory protocols
- User-defined protocols
- High throughput - 96 slides in 20 minutes
- Microwavable containers
- Programmable time and temperature controls
- Built-in probe measures solution temperature in real time
- Time saving and uniform heating
- Eco-friendly solutions





Consumables & Ancillary Reagents



Microscope Slides & Coverslips

OptiPlus™ Positive-Charged Microscope Slides provide a strong adhesive surface for tissues and cells to prevent tissue displacement during harsh pre-treatments such as enzymatic digestion and the microwave Antigen Retrieval method. These slides are ideal for automated systems. Additionally, each slide has a frosted end for easy labeling. The OptiPlus™ Positive-Charged Barrier Slides have all the advantages of our regular OptiPlus™ slides, but also contain hydrophobic barriers that allow the quantity of reagents per slide to be tailored to the size of the specimen. These slides eliminate reagent waste without the need to use a PAP pen, thereby reducing set-up time in manual assays as well as in automated systems. The permanent hydrophobic barriers are compatible with dewaxing solutions and other reagents. The slides are suitable for use with frozen tissue sections, formalin-fixed paraffin sections, and cytology preparations.

Xmatrx® Automated Staining Systems

OptiPlus™ Barrier Slides for Xmatrx® (U.S. & Foreign Equivalent Patents Pending) contain a double hydrophobic barriers that allows formation of an oil seal to prevent evaporation of microreagents during high temperature steps and prolonged incubations. Four different configurations are available:

1. A single test area of 25 x 40 mm (>80 µL of reagent recommended)
2. A single test area of 25 x 25 mm (>40 µL of reagent recommended)
3. A single test area of 18 x 18 mm (>10 µL of reagent recommended)
4. Two test area per slide, each measuring 18 x 18 mm

Coverslips are optimized for use on Xmatrx® staining systems and come in three configurations to accommodate the different barrier slides.



Microscope Barrier Slides & Coverslips for Xmatrx®

Product	1 Box	1 Case
Barrier Slides, 18 x 18 mm (72/box, 1440/case)	XT128-SL	XT128-CL
Barrier Slides, 18 x 18 mm, 2-Zone (72/box, 1440/case)	XT114-SL	XT114-CL
Barrier Slides, 25 x 25 mm (72/box, 1440/case)	XT108-SL	XT108-CL
Barrier Slides, 25 x 40 mm (72/box, 1440/case)	XT134-SL	XT134-CL
Coverslips, 18 x 18 mm (175/box, 1750/case)	XT121-YBX	XT121-XBK
Coverslips, 25 x 25 mm (90/box, 900/case)	XT122-90X	XT122-YQK
Coverslips, 25 x 40 mm (50/box, 500/case)	XT118-50X	XT118-YRK

Microscope Slides & Accesories for Manual

Product	1 Box	1 Case
Barrier Slide, 3 x 1/3 Test Areas	XT014-SL	XT014-CL
Barrier Slides, 2/3 Test Area	XT013-SL	XT013-CL
Microscopic Slides	XT002-SL	XT002-CL
PAP pen (For 500 to 1000 Slides)-1 unit	XT001-PP	N/A

Pipette tips

BioGenex pipette tips are made of high-quality polypropylene and are RNase and heavy metals-free when untampered. Inner surface is extremely smooth and requires minimum wetting. 1 mL pipette tips are optimized for use on BioGenex Xmatrx® Staining Systems, while 200 µL tips are optimized for Xmatrx® staining systems.

Pipette tips for Xmatrx®

Product	1 Box	1 Case
Pipette Tips, 1 mL (192/box, 960/case)	XT105-01X	XT104-05X
Pipette Tips, 200 µL (960/box, 4800/case)	XT146-01X	XT145-05X

Consumables kits for Xmatrx®

Item	SKU	Size	Barrier Slides 25 x40 mm	Barrier Slides 25 x40 mm	Coverslips 25 x 40 mm	Coverslips 25 x 40 mm	1 mL Pipette Tips	200 µl Pipette Tips
IHC kit	XT148-YCDE	200 test	216	NA	1000	NA	384	960
ISH kit	XT144-YAD	100 test	NA	104	NA	900	384	960

Accessories

1. Antigen Retrieval Accessories Kits

The Antigen Retrieval Accessory Kit consists of slide holders and slide baths that make it convenient and compatible with any of the several Antigen Retrieval solutions. To accommodate microwave heating, the slide baths and slide holders are made of heat-stable thermoplastic polyolefin and hydrocarbon polymers of acetal resins. These accessories may be used in a microwave or a pressure cooker.

Item	SKU	Slide Bath + Lid	Slide Holder
24- Slide Accessory kit	MW001-SU	1	1 (24- slide capacity)
72- Slide Accessory kit	MW001-HB	3	3 (72- slide capacity)

2. NordicWare® Microwave Pressure Cooker

Placing the NordicWare® Microwave Pressure Cooker within a microwave is an effective method for enhancing staining with the Antigen Retrieval technique. The heat produced under enhanced pressure can reduce the build up of gas bubbles on the surface of tissues. This improves the intensity of staining, accompanied by preservation of tissue and cell morphology. This pressure cooker is also optimized for use with various BioGenex Antigen Retrieval solutions. BioGenex Catalog number: NW001-PC.



3. PAP Pen for Tissue Staining

The PAP pen is a useful pen-like tool for immunohistochemical staining methods. It is designed to prevent the waste of valuable reagents by forming a water-repellent barrier around the specimen. This barrier creates the proper surface tension to hold an antibody solution or detection reagents within the target area on the slide. The surface tension provided by the PAP pen circle ensures that only the amount of antibody solution needed for sufficient reaction will be applied. Since over-flooding of the slide is eliminated, wiping of excess fluid around the specimen can be avoided. The PAP pen can be used for immunostaining of paraffin sections, frozen sections, and for fluorescent antibody methods. The PAP pen contains a special formulation, which is water repellent. It can be removed, if desired, with xylene or xylene substitutes after the staining procedure is completed. BioGenex Catalog Number: XT001-PP, sufficient for use on 500-1000 slides.

Buffers

Buffers and diluents are available for immunohistochemistry, *in situ* hybridization Special Stains and most other applications.

- General buffers, such as PBS (pH 7.6) and TBS (pH 7.6, 0.1M) can be used for washing/rinsing of slides.
- Super Sensitive™ Wash Buffer is phosphate buffered saline (pH 7.4) with surfactant and is used to ensure optimal staining with even spreading of antibodies and other reagents to avoid inconsistent results.

Buffers - Manual & Automation

Product Name	500 mL ^(20x)
Phosphate Buffered saline	HK091-9K
Super Sensitive Wash Buffer	HK583-5K
Tris Buffer (Wash Buffer) 3/Pack (dry powder to make 3L)	HK098-5K

Counterstains and Mounting Media

BioGenex offers the following counterstains for use in Immunohistochemistry, *in situ* Hybridization and other applications with either manual or automated staining systems.

- Mayer's hematoxylin is a blue stain that does not contain alcohol and therefore is compatible with both alcohol soluble non-permanent chromogens (AEC, Fast Red & New Fuchsin) and alcohol-insoluble chromogens (DAB & Elegance Red). It is alcohol and xylene insoluble and therefore compatible with most clearing agents and mounting media.

Product Name	1 mL ^(RTU)	6 mL ^(RTU)	250 mL ^(RTU)
Hematoxylin, Mayer's (IHC, ISH)	NA	HK100-5K	HK100-9K

Mounting of all stained biological specimens is an essential step before their microscopic evaluation. Mounting also enables the slides to be archived for long periods of time. The mounting medium may be used to attach a coverslip or may itself serve as a coverslip substitute. The choice of mounting medium depends on whether long-term or short-term preservation is desired, and whether the mounting procedure is chemically compatible with the chromogen and the counterstain.

- SuperMount® Permanent Mounting Medium is a polymer based aqueous mounting media that does not require the use of a coverslip. This innovative, patented mounting medium (BioGenex's U.S. Patent No. 5,492,837) is designed to preserve biological specimens for long-term storage. SuperMount® medium is compatible with most aqueous and organic-soluble dyes and chromogens including AEC, DAB, Elegance Red, Fast Red, New Fuchsin, BCIP/NBT, Rhodamine, Fluorescein, Texas Red, Phycoerythrin, Phycocyanin, and Fat Stain (Oil Red O). The refractive index of SuperMount® yields greater transparency and clarity of specimens to be examined under the microscope. SuperMount® can be used for the mounting of all biological specimens, including stained tissue sections, cytospin preparations, and blood smears.

- Aqueous Mounting Medium is glycerol-based mounting medium that require the use of a coverslip. It is intended for short-term specimen storage and is compatible with most chromogens and counterstains.

- XMount™ Mounting Medium is a permanent mounting medium that has been optimized for use with BioGenex™ instrument for all BioGenex detection systems for immunohistochemistry (IHC), *In Situ* Hybridization (ISH) and special stains. XMount™ is intended for use with alcohol and xylene insoluble chromogens, such as DAB (for peroxidase systems) and Elegance Red (for alkaline phosphatase systems). XMount™ dries clear with an ideal refractive index similar to high quality glass and tissue elements. Mounted slides can be viewed with high magnification oil immersion lenses. Also, when mounting preparations stained with the BCIP/NBT substrate, crystal formation that may occur when using other media is minimized.

Mounting Medium

Product Name	15 mL ^(RTU)	50 mL ^(RTU)
Aqueous Mounting Medium - Manual	HK099-5K	NA
SuperMount™ Permanent Mounting Medium - Manual	HK079-5K	HK079-7K
Xmount™ Mounting Media (200 tests) – Barcoded	HX035-YCD	NA
Xmount™ Mounting Media (200 tests) – Xmatrix® Infinity	HX035-10X	NA



MicroRNA Tissue Control



Positive Control Slides and Barrier Slides

Positive control slides are made with tissue which has undergone processing identical to that of the test tissue. BioGenex provides positive control slides that enable one to confirm miRNA detection.

Barrier slides are positive control tissue slides with barriers to prevent loss of reagent.

Pack size: Positive Control slides (5 slides per pack)

Barrier slides (5 slides per pack)

Cat.No	Product	Recommended Positive Control	BarrierSlidesCat.No
HM001-100	Hsa-miR-1 Probe	Heart	FB-HM001
HM007A-100	Has-miR-7a Probe	Prostate, Intestine, Pancrease	FB-HM007A
HM007B-100	Hsa-miR-let-7b Probe	Prostate Ca	FB-HM007B
HM007C-100	Hsa-miR-Let-7c	Breast	FB-HM007C
HM007D-100	Hsa-miR-let-7d Probe	Prostate Ca, Prostate	FB-HM007D
HM007-100	Hsa-miR-7e	Breast, Lung	FB-HM007E
HM007G-100	Hsa-miR-let-7g Probe	Intestine	FB-HM007G
HM009-100	Hsa-miR-9	Stomach Ca, Colon Ca	FB-HM009
HM010B-100	Has-miR-10b Probe	Prostate Ca, Small Cell Lung Ca	FB-HM010B
HM015A-100	Hsa-miR-15a Probe	Thyroid	FB-HM015A
HM015B-100	Hsa-miR-15B Probe	TCC, Bladder Ca	FB-HM015B
HM016-100	Hsa-miR-16 Probe	colon	FB-HM016
HM017-100	Has-miR-17 Probe	Prostate Ca, Colon Ca, Colon Ca	FB-HM017
HM017-3P-100	Hsa-miR-17-3p	Prostate Ca, Colon Ca, Colon Ca	FB-HM017-3P
HM018A-100	Hsa-miR-18a	TCC	FB-HM018A
HM019A-100	Hsa-miR-19a	TCC	FB-HM019A
HM019B-3P-100	Hsa-miR-19b-3p	Prostate Ca	FB-HM019B-3P
HM020A-100	Hsa-miR-20A Probe	Ovary Ca, Stomach Ca	FB-HM020A
HM021-100	Hsa-miR-21 Probe	Breast Ca	FB-HM021
HM021-3P-100	Hsa-miR-21-3p	Breast Ca	FB-HM021-3P
HM022-100	Hsa-miR-22 Probe	Breast	FB-HM022
HM023A-100*	Hsa-miR-023A	-	FB-HM023A
HM023B-100	Hsa-miR-23b	Prostate Ca	FB-HM023B
HM024-3P-100	Hsa-miR-24-3P	T Cell Lymphoma	FB-HM024-3P
HM025-100	Hsa-miR-25	N. Breast/ N. Pancreas	FB-HM025
HM026A-100	Hsa-miR-26A Probe	Ca.Liver / N.intestine	FB-HM026A
HM026B-100	Hsa-miR-26B Probe	Ovary Ca	FB-HM026B
HM027A-100	Hsa-miR-27A	Breast, Breast Ca	FB-HM027A
HM027B-100	Hsa-miR-27b	Breast, Prostate Ca	FB-HM027B
HM028-3P-100	Hsa-miR-28-3P Probe	Colon, Hemangioma	FB-HM028-3P
HM028-5P-100	Hsa-miR-28-5P Probe	Non-Hodgkin's lymphoma	FB-HM028-5P
HM29A-100	Hsa-miR-029A	TCC	FB-HM29A
HM29b-3p-100	Hsa-miR-029b-3p	Colon	FB-HM29b-3p
HM029C-100	Hsa-miR-29C	Lung Ca	FB-HM029C
HM030B-100	Hsa-miR-30B Probe	Stomach Ca	FB-HM030B
HM030C-100	Hsa-miR-30C	Breast Ca	FB-HM030C
HM030-100	Hsa-miR-30E	Breast	FB-HM030E
HM031-100	Hsa-miR-31 Probe	Lymphonode testis	FB-HM031

*Please inquire

NOTE: The list for positive control slides is constantly being updated, depending upon tissue availability. Please call 1(800) 421-4149 for availability or visit our website at www.biogenex.com

Cat.No	Product	Recommended Positive Control	BarrierSlidesCat.No
HM034A-100	Hsa-miR-34A Probe	Breast, Prostate, Colon	FB-HM034A
HM34C-100*	Hsa-miR-034C	-	FB-HM34C
HM0650-100	Hsa-miR-650 Probe	GIST	FB-HM0650
HM092A-100	Hsa-miR-92A Probe	Lymphnode testis	FB-HM092A
HM095-100	Hsa-miR-95 Probe	Small cell lung Ca	FB-HM095
HM096-100	Hsa-miR-96	TCC, Colon Ca, Breast Ca	FB-HM096
HM098-100	Hsa-miR-98	Ovary Ca	FB-HM098
HM099A-100	Hsa-miR-99A Probe	GIST	FB-HM099A
HM099B-100	Hsa-miR-99B Probe	Breast, Colon	FB-HM099B
HM100-100	Hsa-miR-100 Probe	Testis	FB-HM100
HM101-100	Hsa-miR-101	LN	FB-HM101
HM101-3P-100	Hsa-miR-101-3p	LN	FB-HM101-3P
HM106A-100	Has-miR-106a Probe	Liver Ca, TCC, Colon Ca	FB-HM106A
HM107-100	Hsa-miR-107 Probe	Small cell lung Ca	FB-HM107
HM1181-100	Hsa-miR-1181	N. Ovary/N.Pancreas	FB-HM1181
HM122-100	Hsa-miR-122 Probe	Bone, Pancrease	FB-HM122
HM124-100	Hsa-miR-124 Probe	Ca.Ovary	FB-HM124
HM1247-100	Hsa-miR-1247 Probe	TCC, Bladder Ca, Lung Ca	FB-HM1247
HM1258-100	Hsa-miR-1258	TCC, Thyroid, Breast	FB-HM1258
HM125A-100	Hsa-miR-125A Probe	Prostate, Pancrease, Ovary Ca	FB-HM125A
HM125B-100	Has-miR-125b Probe	Ovary	FB-HM125B
HM126-100	Has-miR-126 Probe	Cervix, Ovary, Prostate, Breast, Intestine	FB-HM126
HM127-3P-100	Hsa-miR-127-3P Probe	TCC	FB-HM127-3P
HM1285-100	Has-miR-1285 Probe	Cervix, Ovary, NC, Prostate, Intestine, Breast	FB-HM1285
HM129-100	Hsa-miR-129	Stomach Ca	FB-HM129
HM1296-100	Hsa-miR-1296	Testis	FB-HM1296
HM1297-100	Hsa-miR-1297	Colon	FB-HM1297
HM130B-100	Hsa-miR-130B	Oesophagus Ca	FB-HM130B
HM132-100	Hsa-miR-132	TCC	FB-HM132
HM133A-100	Hsa-miR-133A Probe	Prostate Ca	FB-HM133A
HM133B-100	Hsa-miR-133B Probe	TCC	FB-HM133B
HM135A-100	Hsa-miR-135A Probe	Prostate Ca	FB-HM135A
HM135B-100	Hsa-miR-135B Probe	TCC	FB-HM135B
HM136-100	Hsa-miR-136	Small Cell Lung Ca, Stomach Ca	FB-HM136
HM137-100	Hsa-miR-137	TCC	FB-HM137
HM138-100	Hsa-miR-138	Colon Ca	FB-HM138
HM140-100	Hsa-miR-140	Ovary Ca	FB-HM140
HM141-100	Has-miR-141 Probe	TCC, Prostate	FB-HM141
HM142-100	Hsa-miR-142	Ca.Lung/Ca.Breast	FB-HM142
HM142-3P-100	Hsa-miR-142-3P Probe	Ca.Lung/Ca.Breast	FB-HM142-3P
HM143-100	Hsa-miR-143	Pancrease, Prostate Ca, Colon Ca	FB-HM143
HM144-100	Has-miR-144 Probe	Urinary bladder/Prostate	FB-HM144
HM145-100	Has-miR-144 Probe	Human prostate tissues	FB-HM145
HM146A-100	Hsa-miR-146a Probe	Breast, Intestine, Ovary	FB-HM146A
HM146B-100	Hsa-miR-146B	Prostate, TCC, Breast Ca	FB-HM146B
HM147B-100	Has-miR-147b Probe	Breast, Prostate	FB-HM147B
HM148A-100	Hsa-miR-148A Probe	Prostate, Colon, Breast, Testis	FB-HM148A

*Please inquire

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Cat.No	Product	Recommended Positive Control	BarrierSlidesCat.No
HM148B-100	Hsa-miR-148B Probe	Intestine, Breast, Lung	FB-HM148B
HM149-100	Hsa-miR-149	N.Breast/N.Colon	FB-HM149
HM150-100	Hsa-miR-150 Probe	Lymphonode testis	FB-HM150
HM151A-3p-100	Has-miR-151a-3p Probe	Breast, Thyroid, Esophagus, GB	FB-HM151A-3p
HM152-100	Has-miR-152 Probe	Thyroid, Ovary, Breast, Skin	FB-HM152
HM153-100	Hsa-miR-153	Colon Ca, TCC	FB-HM153
HM154-100	Hsa-miR-154	Lung	FB-HM154
HM155-100	Hsa-miR-155 Probe	Hodgkins Lymphoma	FB-HM155
HM181A-100	Hsa-miR-181A Probe	Sqc. Ca, TCC, Colon Ca	FB-HM181A
HM181B-100	Hsa-miR-181B Probe	TCC	FB-HM181B
HM181C-100	Hsa-miR-181C Probe	Breast Ca	FB-HM181C
HM182-100	Hsa-miR-182	Bladder Ca, Colon Ca, Lung Ca	FB-HM182
HM1826-100	Hsa-miR-1826 Probe	TCC, Bladder Ca	FB-HM1826
HM183-100	Hsa-miR-183	Breast Ca, TCC, Colon Ca, Ad. Ca. Intestine	FB-HM183
HM183-3p-100	Hsa-miR-183-3p	Breast Ca, TCC, Colon Ca, Ad. Ca. Intestine	FB-HM183-3p
HM184-100	Hsa-miR-184	BCC	FB-HM184
HM185-100	Hsa-miR-185	Kidney Ca, GIST	FB-HM185
HM186-100	Hsa-miR-186	Thyroid, Breast, TCC, Colon	FB-HM186
HM187-100	Hsa-miR-187 Probe	Prostate	FB-HM187
HM191-100	Hsa-miR-191 Probe	Lymphonode testis	FB-HM191
HM192-100	Hsa-miR-192 Probe	Colon	FB-HM192
HM193A-3P-100	Hsa-miR-193A-3P	Breast	FB-HM193A-3P
HM193B-100	Hsa-miR-193B	TCC	FB-HM193B
HM194-100	Hsa-miR-194 Probe	TCC	FB-HM194
HM195-100	Hsa-miR-195 Probe	Lymphonode testis	FB-HM195
HM196A-100	Has-miR-196a Probe	Lymphonode testis	FB-HM196A
HM197-100	Hsa-miR-197	N. Liver	FB-HM197
HM198-100*	Hsa-miR-198	-	FB-HM198
HM199A-100	Hsa-miR-199a	Liver Ca	FB-HM199A
HM200A-100	Has-miR-200a Probe	Breast, Prostate, Intestine	FB-HM200A
HM200B-100	Has-miR-200b Probe	TCC, Prostate	FB-HM200B
HM200C-100	Hsa-miR-200C	TCC, Prostate	FB-HM200C
HM203A-3P-100	Hsa-miR-203A	Ad. Ca, Esophagus Ca, TCC, RCC	FB-HM203A-3P
HM204-100	Has-miR-204 Probe	Breast	FB-HM204
HM205-100	Has-miR-205 Probe	Lymphonode testis	FB-HM205
HM206-100	Hsa-miR-206 Probe	Intestine, Breast	FB-HM206
HM210-100	Hsa-miR-210 Probe	Breast Ca, RCC	FB-HM210
HM211-100	Hsa-miR-211	Kidney	FB-HM211
HM212-100	Hsa-miR-212 Probe	Lung, Prostate, Liver Ca, Prostate Ca, GIST	FB-HM212
HM214-100	Hsa-miR-214 Probe	Ovary Ca	FB-HM214
HM215-100	Hsa-miR-215 Probe	Colon Ca, Prostate Ca	FB-HM215
HM216A-100	Has-miR-216a Probe	Lymphonode testis	FB-HM216A
HM216B-100	Hsa-miR-216B	Stomach Ca, Esophagus	FB-HM216B
HM217-100	Hsa-miR-217	N. Prostrate/ Ca. Liver	FB-HM217
HM218-100	Hsa-miR-218	Normal cervix/Ca. breast	FB-HM218
HM221-3P-100	Hsa-miR-221-3p	Kidney, Colon	FB-HM221-3P
HM222-100	Hsa-miR-222 Probe	Ca. Breast/ Ca. Lung	FB-HM222

*Please inquire

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Cat.No	Product	Recommended Positive Control	BarrierSlidesCat.No
HM223-100	Hsa-miR-223	N. Breast	FB-HM223
HM224-100	Hsa-miR-224 Probe	Breast Ca	FB-HM224
HM24-2-100	Hsa-miR-24-2	Sqc. Ca	FB-HM24-2
HM296-100	Hsa-miR-296	TCC, Prostate	FB-HM296
HM297-100	Hsa-miR-297	TCC	FB-HM297
HM300-100	Hsa-miR-300	Gall bladder, Ad. Ca, TCC	FB-HM300
HM328-100	Hsa-miR-328 Probe	Lymphonode testis, Tonsil	FB-HM328
HM329-100	Hsa-miR-329 Probe	Breast, Prostate	FB-HM329
HM330-100	Hsa-miR-330	Prostate, LN, TCC	FB-HM330
HM331-3P-100	Hsa-miR-331-3p	Prostrate Ca	FB-HM331-3P
HM335-100	Hsa-miR-335	Breast, Intestine, Ovary, Colon Ca	FB-HM335
HM337-100	Hsa-miR-337	Lymph Node	FB-HM337
HM338-3p-100	Hsa-miR-338-3p	Breast	FB-HM338-3p
HM339-5p-100	Hsa-miR-339-5p	Kidney, TCC	FB-HM339-5p
HM342-3p-100	Hsa-miR-342-3p	Testis	FB-HM342-3p
HM361-100	Hsa-miR-361 Probe	Prostate	FB-HM361
HM362-100	Hsa-miR-362 Probe	Prostate Ca, Lung, Lymphonode testis	FB-HM362
HM365A-3P-100	Hsa-miR-365A-3P	Ca. Prostate/Ca.Ovary	FB-HM365A-3P
HM372-100	Hsa-miR-372	Cervix	FB-HM372
HM373-100	Hsa-miR-373 Probe	Lymphonode testis	FB-HM373
HM374A-100	Hsa-miR-374A	Colon Ca, Colon, Breast Ca	FB-HM374A
HM374B-100	Hsa-miR-374B	Lymph Node	FB-HM374B
HM375-100	Has-miR-375 Probe	Colon, Hemangioma. Kidney	FB-HM375
HM376C-100	Hsa-miR-376C	Bone	FB-HM376C
HM378A-100	Hsa-miR-378A	Bladder Ca, Liver Ca, GIST	FB-HM378A
HM379-100	Hsa-miR-379	Prostate, TCC	FB-HM379
HM381-100	Hsa-miR-381	TCC, Breast	FB-HM381
HM383-100	Hsa-miR-383	Prostate Ca, Melanoma	FB-HM383
HM409-3P-100	Hsa-miR-409-3P Probe	Breast, Prostate	FB-HM409-3P
HM410-100	Hsa-miR-410 Probe	TCC, GIST	FB-HM410
HM412-100	Hsa-miR-412 Probe	GIST	FB-HM412
HM422A-100	Hsa-miR-422A	Stomach	FB-HM422A
HM423-3P-100	Hsa-miR-423-3p	TCC, Breast Ca	FB-HM423-3P
HM424-100	Hsa-miR-424 Probe	Breast Ca	FB-HM424
HM425-100	Hsa-miR-425	Breast	FB-HM425
HM429-100	Hsa-miR-429 Probe	Prostate, Ovary, Colon	FB-HM429
HM449A-100	Hsa-miR-449A Probe	Colon, Breast	FB-HM449A
HM450B-3P-100	Hsa-miR-450B-3P	Thyroid, Ovary	FB-HM450B-3P
HM451-100	Hsa-miR-451 Probe	Thyroid, Lung, Ovary	FB-HM451
HM4723-100	Hsa-miR-4723-5p	TCC	FB-HM4723
HM483-100	Hsa-miR-483	Lymphonode testis	FB-HM483
HM486-100	Hsa-miR-486 Probe	Lung	FB-HM486
HM486-3P-100	Hsa-miR-486-3P	Lung	FB-HM486-3P
HM494-100	Hsa-miR-494 Probe	Breast Ca	FB-HM494
HM495-100	Hsa-miR-495	TCC, Ovary, Breast Ca	FB-HM495
HM497-100	Hsa-miR-497 Probe	BCC, TCC	FB-HM497
HM502-100	Hsa-miR-502	Gall bladder	FB-HM502

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Cat.No	Product	Recommended Positive Control	BarrierSlidesCat.No
HM505-100	Hsa-miR-505	Breast, Intestine, Ovary, Prostate Ca	FB-HM505
HM508-3p-100	Hsa-miR-508-3p	Breast	FB-HM508-3p
HM509-3p-100	Hsa-miR-509-3p	TCC	FB-HM509-3p
HM510-100	Hsa-miR-510	Thyroid	FB-HM510
HM511-100	Hsa-miR-511	Thyroid, Breast	FB-HM511
HM517A-3p-100	Hsa-miR-517A-3p	Thyroid	FB-HM517A-3p
HM520C-100	Hsa-miR-520C	Breast	FB-HM520C
HM532-5p-100	Hsa-miR-532-5p	Ovary	FB-HM532-5p
HM541-100	Hsa-miR-541	Pancrease	FB-HM541
HM544-100	Hsa-miR-544 Probe	Intestine, Breast	FB-HM544
HM545-5P-100	Hsa-miR-545-5P	Breast	FB-HM545-5P
HM573-100	Hsa-miR-573	Skin	FB-HM573
HM574-3p-100	Hsa-miR-574-3p	Breast, TCC	FB-HM574-3p
HM590-100	Hsa-miR-590 Probe	Stomach Ca	FB-HM590
HM610-100	Hsa-miR-610	Breast	FB-HM610
HM614-100	Hsa-miR-614	BCC, Skin	FB-HM614
HM615-100	Hsa-miR-615	Breast, Intestine, Ovary, TCC	FB-HM615
HM622-100	Hsa-miR-622 Probe	Breast, Colon	FB-HM622
HM625-100	Hsa-miR-625 Probe	Intestine, Breast	FB-HM625
HM627-100	Hsa-miR-627	Breast	FB-HM627
HM628-100	Hsa-miR-628 Probe	Prostate	FB-HM628
HM629-100	Hsa-miR-629	Non-Hodgkin's lymphoma, Prostate Ca	FB-HM629
HM630-100	Hsa-miR-630	Breast Ca	FB-HM630
HM638-100	Hsa-miR-638	Colon Ca, TCC	FB-HM638
HM641-100	Hsa-miR-641	Breast, GB, Thyroid, Ovary	FB-HM641
HM642A-5p-100	Hsa-miR-642A-5p	Breast, Prostate, Lung	FB-HM642A-5p
HM648-100	Hsa-miR-648 Probe	RCC	FB-HM648
HM663A-100	Hsa-miR-663A Probe	Prostate	FB-HM663A
HM708-100	Hsa-miR-708	Bladder Ca	FB-HM708
HM718-100	Hsa-miR-718 Probe	Ovary, Intestine, LN	FB-HM718
HM765-100	Hsa-miR-765	Lung	FB-HM765
HM802-100	Hsa-miR-802	Intestine	FB-HM802
HM874-100	Hsa-miR-874	Intestine	FB-HM874
HM940-100	Hsa-miR-940	GIST	FB-HM940
HM944-100	Hsa-miR-944	Breast	FB-HM944
HM9500-100	Hsa-miR-9500	TCC	FB-HM9500
HM128-100	Hsa-miR-128	Brain Tumor	FB-HM128
HM139-100	Hsa-miR-139	Bladder	FB-HM139
HM190a-100	Hsa-miR-190a	Breast Cancer	FB-HM190a
HM190b-100	Hsa-miR-190b	Lung Ca.	FB-HM190b
HM193b-100	Hsa-miR-193b	Colorectal Ca	FB-HM193b
HM302b-100	Hsa-miR-302b	Gastric Ca.	FB-HM302b
HM326-100	Hsa-miR-326	Colorectal Ca.	FB-HM326
HM378a-100	Hsa-miR-378a	Colorectal Ca.	FB-HM378a
HM382-100	Hsa-miR-382	Lung Ca.	FB-HM382
HM384-100	Hsa-miR-384	RCC	FB-HM384
HM433-100	Hsa-miR-433	Colorectal Ca.	FB-HM433

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Cat.No	Product	Recommended Positive Control	BarrierSlidesCat.No
HM489-100	Hsa-miR-489	Breast Ca.	FB-HM489
HM491-100	Hsa-miR-491	Breast Ca.	FB-HM491
HM498-100	Hsa-miR-498	Lung Ca.	FB-HM498
HM514a-100	Hsa-miR-514a	Melanoma	FB-HM514a
HM524-100	Hsa-miR-524	Melanoma	FB-HM524
HM675-100	Hsa-miR-675	Skin	FB-HM675
HM766-100	Hsa-miR-766	Kidney	FB-HM766
HM1244-1-100	Hsa-miR-1244-1	Tonsil	FB-HM1244-1
HM3978-100	Hsa-miR-3978	Prostate ca.	FB-HM3978

NOTE: The list for positive control slides is constantly being updated, depending upon tissue availability. Please call 1(800) 421-4149 for availability or visit our website at www.biogenex.com

General Terms and Conditions

1. Order Information

- Credit Terms: BioGenex will review the customer credit application and finalize the terms (Credit Limit and Net Days) based on inputs provided and credit rating.
- Order Confirmation: To avoid shipment duplication, please indicate in bold "**CONFIRMING ORDER - PLEASE DO NOT SHIP**" on your order.

2. Conditions of Sale

- All prices are quoted in U.S. dollars, exclusive of Sales Tax (State and County), as applicable.
- If an order is not taxable, a tax exemption certificate must be provided.
- Products and prices are subject to change without any prior notice.
- Discounts: Please inquire about BioGenex quantity discount policies at 1-800-421-4149.
- Payment: All payments must be made in U.S. dollars. You may choose any mode of payment (Note: Online payment systems are implemented).

3. Return and Refund Policy

BioGenex reagents are covered by Quality Assurance (QA) policy:

- Returns will only be accepted with BioGenex Return Material Authorization (RMA). Please contact customer service for further assistance.
- BioGenex has a limited liability for a refund or replacement. The same is solely under the discretion of BioGenex management.
- A full refund will be provided when a product cannot perform according to data specifications.
- If client makes an error in ordering a product, a refund may be provided along with a 30% restocking fee.
- Express Delivery: Express delivery options are also available on request at an extra cost.
- BioGenex customer service for assistance:
Tel: 1-800-421-4149, Monday through Friday
7 AM – 4 PM PST or
E-mail at: customer.service@biogenex.com

4. Other Terms and Conditions

- BioGenex is committed to quality, innovation, service, and support. We believe that the high degree of quality control performed on all our products will help you with consistent and reproducible results.
- All orders are subject to acceptance by BioGenex and product availability.
- Delivery dates are estimates and BioGenex shall have no liability for any delays.
- There are no expressed, implied or statutory warranties,

including without limitation, the implied warranties of merchantability, fitness for a particular purpose and non-infringement of third party rights.

- Freight charges are prepaid and added to the invoice.
- BioGenex shall not be liable for any incidental, indirect, special or consequential damages, even if it is aware of the possibility of such damages. BioGenex's total liability for any order shall not exceed the amount paid by customer under such order.
- These terms and conditions constitute the entire agreement between the parties with respect to the products purchased hereunder.
- Any additional, different or inconsistent terms and conditions in a purchase order form or like forms used by customer to purchase, change, accept or otherwise process the orders are objected to and not binding on BioGenex.
- This agreement between the parties shall be governed by the laws of the State of California without regard to its conflicts of laws.
- Any dispute arising out of or related to this Agreement shall be resolved solely in the U.S. District Court for the Northern District of California or in San Francisco County, and in no other courts, and Customer hereby consents to the jurisdiction of, venue in and service of process from the aforementioned courts.

Super Sensitive Nucleic Acid System miRNA *In Situ* hybridization

Probe Design

- High specificity and sensitivity stemming from high-melting temperature probe
- Labeled with high-density reporter molecules to enable single copy gene visualization

miRNA *In Situ* hybridization

- Designed to provide intense super clean stains
- Localization of target cells in the spatial context
- Multiplexing miRNA, IHC & FISH targets

System Provides

- Optimized protocols
- Automation from microtome to microscope
- Ready to use probes and visualization system
- Ready to use reagents including stringency washes



In the U.S., call +1 (800) 421-4149
Outside the U.S., call +91-40-27185500



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