





In situ Hybridisation and Detection

FISH detection assays-v2

	FISH kit					
Label	Product number	# assays				
Green	CxxxK.2000.05	5 T				
Green	CxxxK.2000.10	10 T				
Orange	CxxxK.3000.05	5 T				
Orange	CxxxK.3000.10	10 T				
Green and orange	CxxxK.2030.05	5 T				
Green and orange	CxxxK.2030.10	10 T				
Green and orange	C801K.5206	20 T				

PANPATH REMBRANDT® FISH DETECTION KITS



PAGE 1 OF 20

Contents

			page
Chapter 1		Introduction	03
	1.1	Intended use	03
	1.2	The FISH principle	03
	1.3	Controls	03
	1.4	Contents of a REMBRANDT® FISH	03
	1.5	Materials required but not included	04
	1.6	Single product catalogue numbers, storage and shelf life	04
	1.7	Safety precautions	05
	1.8	Performance precautions	05
	1.9	Preparations of reagents in advance	06
	1.10	Preparations of reagents and materials needed but not supplied	06
	1.11	Microscope and accessories	07
Chapter 2		REMBRANDT® FISH protocol	09
	2.1	Specimen collection	09
	2.2	Pre-treatment of specimen	09
	2.3	Hybridisation procedure and post-hybridization washes	10
Chapter 3		Interpretation of the results	12
	3.1	Guidelines for interpretation	12
	3.2	Quality control	12
Chapter 4		Limitations of procedure	14
	4.1	Limitations	14
Chapter 5		References	15
Chapter 6		Trouble shooting guide	16
	6.1	Introduction	16
		Cytological specimen	16
	6.3	Paraffin embedded tissue sections	17
Immateria	l pro	pperty information	20

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Chapter 1 Introduction

1.1 Intended use

REMBRANDT has been designed for the detection of specific DNA or RNA sequences by using the fluorescence in situ Hybridisation (FISH) technique in cytological specimen or FFPE tissue sections. In extensive tests and field application the REMBRANDT FISH kits were proven to be very robust, and FISH results were highly reproducible. For the specific intended use of a REMBRANDT FISH detection assay, please see the product specific datasheet.

1.2 The FISH principle

FISH enables the detection of specific DNA or RNA sequences in histological and cytological specimens, without losing the often very essential morphological details. The principle of FISH is based on a "reaction" (= hybridisation) between a specifically labelled DNA or RNA sequence (= probe) and a DNA or RNA sequence present in the sample (= target). In case of matching sequences, a hybrid between the probe and target will be formed. Non-specific hybrids are washed out by the stringency wash procedure (PanWash 4). The formed hybrids can easily be visualised via fluorescent microscopy. The REMBRANDT® probes are labelled with fluorochromes; the fluorochromes used in the kits depend on specific probe(s) in the kit (Arsham et al., 2017).

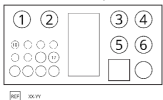
1.3 Controls

Use of controls is an essential part of the routine. To ensure that the FISH procedure is performed correctly and that observed positive and/or negative staining are specific, controls should be included in each experiment by the end-user.

1.4 Contents of a REMBRANDT® FISH detection assay

Application: cytological specimen

Contents REMBRANDT® FISH detection assays



Pos.	Contents
1	Pepsin diluent
2	Pepsin powder
3	PanWash 4
4	PanWash 4
5	PanWash 4
6	PanWash 4
10	Product specific probe
17	Mounting medium fluorescence

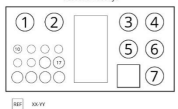
Product label		Vol.		REF
DIGEST	PEPSIN DIL	15	ml.	R018R.0000
DIGEST	PEPSIN POW	1	gr.	R011R.0000
PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
PROBE	SPECIFIC PROBE	75 150	μl μl	XXXXX.YYYY
SUPPORT	MOUNT FLU	1	ml.	Z000R.0050

PANPATH REMBRANDT® FISH DETECTION KITS



Application: FFPE tissue sections

Contents REMBRANDT® FISH detection assays



Pos.	Contents	Product label		Vol.		REF
1	Pepsin diluent	DIGEST	PEPSIN DIL	15	ml.	R018R.0000
2	Pepsin powder	DIGEST	PEPSIN POW	1	gr.	R011R.0000
3	PanWash 4	PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
4	PanWash 4	PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
5	PanWash 4	PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
6	PanWash 4	PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
7	Pre-treatment buffer	PRE-TREAT	Na-Citrate 100x	15	ml.	R026R.0000
10	Product specific probe	PROBE	SPECIFIC PROBE	75	μl	XXXXX.YYYY
	P. C. C.			150	μΙ	
17	Mounting medium fluorescence	SUPPORT	MOUNT FLU	1	ml.	Z000R.0050

1.5 Materials required but not included

- Xylene for dewaxing paraffin sections

- Fixative for cytological specimens

- Distilled or deionised water

- Ethanol series - 100% Ethanol

- 96% Ethanol

- 70% Ethanol

- 70%

- HCl 0.01 M

- Phosphate buffered saline (PBS); pH 7.2-7.4

- Pipettes and tips to deliver 10-1000 μL

- Immersion oil for fluorescence microscopy

- Timer(s)

- Fluorescence microscope and accessories

- Oven(s), heating block(s) and water baths

- Incubation chamber

- Mixing device i.e. vortex

- Magnetic stirrer

- (Surface) thermometer; calibrated

- pH meter and calibrators

- Coverslips

- Slides

- Microwave (optional)

1.6 Single product catalogue number, storage and shelf life

REMBRANDT	p	robe mix; 1 vial	Catalogue No	CXXXP.XXXX
Quantity	:	75 or 150 μl; 5 or 10 assays/kit	Storage	2 – 8°C
Composition : specific probe DNA fluorescently labelled, ready to use in hybridisation buffer.				

Pepsin digesti	ion	reagent; 1 vial	Catalogue No	R011R.0000
Quantity	:	1 gr; powder to be dissolved in 8 mL deionised water, to be aliquoted in portions of e.g. 1200 μ l or 1200 μ l depending on the amount of slides per test run	Storage powder dissolved solution	2 – 25°C -20°C
Composition	:	Pepsin powder		

Pepsin dilue	ent; 1 vial	Catalogue No	R018R.0000
Quantity	: 15 mL 1M to be diluted 100x with deionised water	Storage	
	~F 011		

PANPATH REMBRANDT® FISH DETECTION KITS

Path

PAGE 4 OF 20

		to 0.01 M HCI	1 M HCI 0.01 M HCI	2 – 25°C 2 – 25°C
Composition	:	HCI, 1M		

Concentrated PanWash 4, 25x SSC; 4 vials Catalogue No				R025R.0000
Quantity	:	15 mL concentrated solution to be diluted 12.5x (= 2x SSC) with deionised water and confirm (and adjust if needed) to pH 7.4	Storage conc. solution diluted solution	2 – 25°C ambient temp.
Composition	:	Sodium chloride, sodium citrate and surfactant		

Concentrated	Pre-treatment buffer, 100x Na-Citrate; 1 vial (optional)	Catalogue No	R025R.0000
Quantity	 15 mL concentrated solution to be diluted 100x (= 1x Na- Citrate) with deionised water and confirm (and adjust if needed) pH 6.0 	Storage conc. solution diluted solution	
Composition	: Natrium citrate		

Mounting med	liui	m; 1 vial	Catalogue No	Z000R.0050
Quantity	:	2 mL	Storage	2 – 8°C
Composition	:	DAPI (4,6-diamidino-2-phenylindole), DABCO (diazabicycloglycerol	-2.2.2- octane) in	TRIS buffer and

- Store kit and its contents at 2-8°C.
- Store the dissolved and aliquoted reagents at recommended temperatures (see above).
- When used and stored as indicated, the kit is stable until the expiry date printed on the box.

1.7 Safety precautions

- The hybridisation buffer in which the probes are mixed contains formamide, which is considered as a teratogen; contact with skin and mucous membranes should be avoided.
- —The Mounting medium contains DAPI (4,6-diamino-2-phenylindole) and 1,4-phenylenediamine which is a possible respiratory and dermal sensitizer; contact with skin and mucous membranes should be avoided.
- If reagents come into contact with skin or eyes, rinse with large volumes of clean water.
- Never pipette solutions by mouth.
- All hazardous materials should be disposed of according national guidelines.
- A material safety data sheet is available on request

1.8 Performance precautions

- Read all instructions before processing any assay.
- Store all reagents as recommended.
- **DO NOT** use reagents beyond their expiry date.

PANPATH REMBRANDT® FISH DETECTION KITS



PAGE 5 OF 20

- Allow all components to warm up to room temperature (20-25°C) before use.
- -Check all ready to use solutions for precipitates and other contaminations before use; if contaminations are present, the solution should be disposed of, and a new solution should be prepared.
- PanWash 4 concentrated solution may form crystals when stored at low temperatures. If heating
 does not dissolve the crystals, the solution should be disposed of and a new solution should be
 prepared.
- Homogenise all solutions before use.
- Homogenise probe solution and spin briefly before use.
- **DO NOT** substitute a reagent with one from another manufacturer.
- —Use treated glass slides, preferably REMBRANDT® Adhesive coated glass slides (Z000S.0002, PanPath B.V.).
- **DO NOT** re-use prepared, ready to use digestion reagent; dispose of residuals.
- With every new procedure, check the temperatures of pre-treatment solutions, denaturation device used, incubation device used and PanWash 4 solution by using a calibrated thermometer.
- Do not incubate more than 5 specimens at the same time in pre-set temperature baths/devices.
 Placing more than 5 specimens in such a device will cause a temperature drop, and thus will not provide the correct temperature.
- Pre-treatment, Denaturation and Hybridisation are the key steps in the procedure, if any of these steps in the procedure have not been carried out properly, the results may not be relied upon.
- Fluorochromes are light sensitive; therefore, all steps that do not require manipulation should be performed under exclusion of light.

1.9 Preparation of supplied reagents in advance

Pepsin stock solution:

Dissolve the pepsin digestion reagent in 8 mL distilled or deionised water (upon receipt of the kit). Aliquot in portions of i.e. 1200 or 600 μ L and store at- 20° C. This depends on the expectation of the amount of slides that are subject per test run.

Pepsin diluent:

Dilute the supplied pepsin diluent (1M HCl) 100x to 0.01 M HCl with distilled or deionised water.

Proteolytic work solution:

Prepare fresh work solution just before use and discard non-used solution:

- -Dilute aliquoted proteolytic reagent to 1.25 mg/mL for FFPE and 100 μg/ml for cytological specimen in 0.01 M HCl.
- -Pre-heat 0.01 M HCl solution at 37°C, add the required pepsin stock solution just before use and mix.

Do not re-use

PANPATH REMBRANDT® FISH DETECTION KITS



PAGE 6 OF 20

PanWash 4:

- Dilute a vial of the supplied 15 mL PanWash 4, 25x SSC to **2x SSC** with 172.5 mL deionised or distilled water to a total of 187.5 ml (concentration is 2x SSC).
- Dilute 0.8 mL of a vial of the supplied PanWash 4, 25x SSC to **0.1x** SSC with 199.2 mL deionised or distilled water to a total of 200 ml (concentration is 0.1x SSC).

Before use:

- a) Fill staining jars with SSC solution.
- b) Pre-heat 2x SSC at 42°C and 0.1x SSC at 61°C (if applicable)

Do not re-use

Pre-treatment buffer (included depending assay type; see product specific datasheet): Dilute the supplied 15 mL Pre-treatment buffer, 100x Na-Citrate to 1x Na-Citrate with 1485 mL deionised or distilled water.

Do not re-use

1.10 Preparation of reagents and materials needed but not supplied

Dewaxing reagents

i.e. xylene, ethanol 100% and methanol

Do not re-use

Dehydration reagents

i.e. graded ethanol series (ethanol 70%-96%-100%)

PBS (phosphate buffered saline)/ Tween® 20 (0.05%)

- Deionised water:

950 ml

- Sodium chloride (NaCL):

3

-Potassium chloride (KCI):

 $0.2\,$ g

- di-Sodium hydrogen phosphate di-hydrate (Na₂HPO₄.2H₂O):

1.78 8

- Potassium di-hydrogen phosphate (KH₂PO₄):

0.24 g

-Tween® 20

0.5 m

Adjust pH to 7.4 +/- 0.2 and adjust to 1000 mL with deionised water

Do not re-use

PANPATH REMBRANDT® FISH DETECTION KITS

Pan Path

PAGE 7 OF 20

1.11 Microscope and accessories

Microscope

A fluorescence microscope is needed to validate the *in situ* hybridisation results.

Oculars and objectives

10X oculars are sufficient, 20X or 40X objectives are usually used for scanning the section. It is recommended to use at least a 63X oil immersion objective for final analysis.

Light source

The light source of regular microscopes is often 50 or 100 W. However a 100 W light source is recommended to obtain optimal results.

Filters

Most, if not all microscope manufactures, can deliver the needed single and multi-band pass filters. The single and multi-band pass filters needed for the REMBRANDT® FISH detection assays are:

Filter set for DAPI

excitation G 365 nm

beam splitter FT 395 nm emission IP 420 nm

Filter set for green fluorochrome

Excitation BP 450 nm – 490 nm

beam splitter FT 510 nm

emission BP 515 nm – 565 nm

Filter set for orange fluorochrome

excitation BP 546 nm /12

beam splitter FT 580 emission LP 590 nm

Triple band filter set (DAPI / FITC / TRITC)

Immersion oil

The use of quality immersion oil is recommended; the immersion oil should have low auto-fluorescence and be suited for use in fluorescence microscopy.

PANPATH REMBRANDT® FISH DETECTION KITS



PAGE 8 OF 20

Chapter 2 REMBRANDT® FISH Protocol

- All incubation steps should be performed in a closed (dark) incubation chamber/staining jar which contains a liquid (water) creating a saturated moisturised environment. During the incubation steps, evaporation of reagents should be prevented.
- Once the hybridisation procedure has been started the specimen should not be allowed to dry except for those procedural steps that mention "air-dry".
- Allow all reagents to reach the temperatures required for the respective incubation
- Work in a fume hood, use forceps and wear protective laboratory clothing and powder free examination gloves.

2.1 Specimen collection

Cytological specimen

Fixation in 70% ethanol is a commonly used for cytological specimen, it preserves the morphology and target DNA. Other fixation methods such as Carnoy's fixative are also applicable.

Paraffin embedded tissue sections

A standard procedure for tissue fixation and embedding usually involves the use of formalin and paraffin. The optimal tissue block size is 0.5 cm³. The formalin should be buffered and fixation times should (preferably) not exceed 12 hours. Excess and/or insufficient fixation may yield suboptimal morphology and target preservation. Embedding in paraffin should not exceed temperatures of 65°C.

Sample preparation: stretch 4 μ m paraffin sections on distilled water of 38-40°C without any additives and collect sections on bio-adhesive (i.e. organ silane) coated glass slides. Bake the slides at 65°C in a dry air oven for 1 hour. Slides can be used immediately or they can be stored at room temperature for up to 3 months.

Prior to FISH, slides need to be dewaxed in subsequent fresh xylene baths for 2×10 minutes. Incomplete removal of formalin and/or paraffin may affect the result of the procedure. Remove the xylene by placing the slides in subsequent 100% ethanol (2×5 minutes), hydrate in graded ethanol series, deionised water and continue with pre-treatment.

2.2 Pre-treatment of specimen

Cytological specimen

- Incubate slides in pre-heated proteolytic work solution at 37 °C (i.e. 100 μg/ml) for 15 minutes.
- Flush wash in deionised water, followed by dehydration in graded ethanol series (ethanol 70%, 96%, 96%, 100% and 100%), 1 minutes each and air-dry slides for 15 minutes.

PANPATH REMBRANDT® FISH DETECTION KITS



PAGE 9 OF 20

Paraffin embedded tissue sections

For making the DNA accessible for the probes and to obtain reliable and reproducible results, pretreatment procedures are necessary.

- Place slides in jar filled with pre-treatment buffer (R026R.0000), place the jar in a microwave set at i.e. 900W and incubate up until boiling, subsequently reset microwave at 180W and proceed with the incubation for 10 minutes followed by a 20 minute cool down period at room temperature; all in the same solution. Flush wash slides in deionised water at room temperature.
- Incubate slides in pre-heated proteolytic work solution at 37 °C (1.25 mg/ml) for 15 minutes.
- Flush wash in deionised water, followed by dehydration in graded ethanol series (ethanol 70%, 96%, 96%, 100% and 100%), 1 minutes each and air-dry slides for 15 minutes.

Do not treat more than 5 slides at the same time, because the temperature of the pre-heated solutions may drop dramatically, thus causing incomplete pre-treatment. Additionally, allow the slides to air-dry as recommended; otherwise sections will be lost.

2.3 Hybridisation procedure and post-hybridization washes

Denaturation and Hybridisation

- Homogenize probe solution and spin briefly. Apply 10-15 μl of probe solution to each specimen. Cover all specimens with a cover slip (avoid air bubbles).
- Place slides on an 80 °C hotplate or other heating device and incubate for 3 minutes in case of cytological specimen and 10 minutes in case of FFPE tissue sections (denaturation).

Work in a pre-set order to ensure that all slides have been incubated at 80 °C for the exact same time! Do not denature more than 5 slides at the same time, because the temperature of the heating device may drop dramatically, thus causing incomplete denaturation.

Transfer slides into a moist environment and incubate for 16 hours at 37 °C.

Differentiation (stringent wash) and rinsing

- Remove coverslips by submerging the slides in PBS at room temperature. Soak the slides until the coverslips fall off.
- Incubate slides in pre-heated PanWash 4, buffer concentration and incubation temperature differ per specific product (see product specific datasheet).
- Incubate slides in PBS at room temperature for 1 minute.
- Dehydrate in graded ethanol series. Air-dry slides for 15 minutes (in the dark).

Pan Path Do not incubate more than 5 slides at the same time in PanWash 4 (2x SSC and 0.1x SSC), because the temperature of PanWash 4, may drop dramatically, causing wrong stringency conditions.

Coverslipping

Mount slides by applying the supplied mounting medium (Z000R.0050) and coverslip.



Chapter 3 Interpretation of results

3.1 Guidelines for interpretation

- For specific interpretation guidelines and scoring criteria, see REMBRANDT® FISH detection product specific datasheet.
- For assessment it is important to select representative areas of the specimen.
- Over digestion can lead to DNA loss and affect the gene signal numbers in individual nuclei. Score
 only nuclei that have relatively intact nuclear borders.

3.2 Quality control

Positive control

It is advised to run a positive and negative control simultaneously with each FISH assay. As a positive control, one may also use validated in-house specimen. If the controls fail to demonstrate the expected staining, the result on the test specimen must be validated as invalid.

Assay requirements

- Noise-to-signal percentage
 - A FISH probe is detected via fluorescent microscopy. The probe will emit a fluorescent signal. However, a fluorescent signal can also be visualized in the cells. Therefore, the noise-to-signal cutoff needs to be determined for FISH probes. If the fluorescent signal in the background is too close to the actual probe signal, the probe is not useable in diagnostics. For good signal interpretation, the noise-to-signal percentage should be $\leq 40\%$.
- Hybridization efficiency
 Signals may only be interpreted if the probe signals are visualized in ≥98% of the cells.

For additional requirements, see the product specific datasheet.

Invalid results

- In case the control does not stain adequately, the results of the test slides cannot be accepted. The
 run should be repeated in order to rule out procedural errors. Please check the trouble shooting
 section below for further possibilities.
- Poor cell morphology may lead to unacceptable noise-to-signal percentage. In these cases we
 advise to assess the cell morphology quality in parallel brightfield stained specimen before
 performing a repeat test.
- If high background signals are present over the cytoplasm obscuring the true signals, the assay should be repeated.
- If nuclear borders are lost or persistent green or red auto-fluorescence masks true signals, the enzymatic digestion was not optimal and the test should be repeated.
- If hybridisation signals are not reliable/identifiable, the assay should be repeated.

For additional requirements, see the product specific datasheet

PANPATH REMBRANDT® FISH DETECTION KITS



PAGE 12 OF 20

Limitations of Procedure Chapter 4

4.1 Limitations

- The REMBRANDT® FISH detection assays are solely applicable for the detection of corresponding DNA as described in the product specific intended use, which may be present in the test specimen.
- Medical decisions may not be taken based on this test. This is a research use only product.
- Sample fixation methods should be performed according to the recommended sample preparation methods.
- Many factors can influence the performance of the FISH procedure. Failure in detection can be due to i.e. improper sampling, handling, the time lapse between tissue removal and fixation, the fixation time, processing fixed specimen, the bio-adhesive on the slide, incubation times, pretreatment procedures, incubation temperatures, freezing, thawing, washing, heating drying, reagent contaminations and interpretation of results.
- The performance of REMBRANDT® FISH detection assays was tested and validated when using the exact procedure as listed in the product specific data sheets, modifications to the procedure may alter the performance characteristics.
- The REMBRANDT® FISH detection assays test results are not to be relied on in case the sampling, sampling method, quality, sample preparation, reagents used, controls and procedure followed is not optimal.
- The medical profession should be aware of risks and factors influencing the intensity, the absence or presence of FISH signals which cannot be foreseen when applying this test.
- The user should carefully consider the risk and use of sample material for this test in case the sample material does not contain sufficient or representative test material.
- Laboratory personnel performing the test should be knowledgeable, professional and be able to interpret the test results.
- The specific FISH detection assay should only be used for the loci the probe targets.



Product in combination with other devices

The REMBRANDT® FISH detection assays are intended for stand-alone usage. The assay is intended to be used in combination with standard formalin fixed or cytological specimen, hot plate(s), stove(s), incubation device(s), water bath(s), temperature and incubation time control(s), and other reagents (not supplied with this reagent) and a microscope. The combination has been tested and validated. Since the formalin fixed or cytological specimen, hot plate(s), stove(s), incubation device(s), water bath(s), temperature controls, incubation time control(s) and other not supplied reagents such as but not limited to fixation and other reagents and a microscope is not combined with the device as a product, conformity with the essential requirements is not applicable. Laboratory assay validation should always be established by the end-users.

Chapter 5 References

- Arsham, M. S., Barch, M. J., & Lawce, H. J. (2017). The AGT Cytogenetics Laboratory Manual The AGT Cytogenetics Laboratory Manual Edited by (Vol. 4).
- Duffy, L., Zhang, L., R., D., & M., A. (2012). Quality Control Considerations for Fluorescence In Situ Hybridisation of Paraffin-Embedded Pathology Specimens in a Diagnostic Laboratory Environment. Latest Research into Quality Control, Figure 1. https://doi.org/10.5772/51266
- Mascarello, J. T., Hirsch, B., Kearney, H. M., Ketterling, R. P., Olson, S. B., Quigley, D. I., Rao, K. W., Tepperberg, J. H., Tsuchiya, K. D., & Wiktor, A. E. (2011). Section E9 of the American College of Medical Genetics technical standards and guidelines: Fluorescence in situ hybridization. *Genetics in Medicine*, 13(7), 667–675. https://doi.org/10.1097/GIM.0b013e3182227295

Chapter 6 Trouble shooting guide

6.1 Introduction

This Trouble Shooting Guide is intended to support you in obtaining optimal results with PanPaths REMBRANDT* FISH detection assays (Duffy et al., 2012) (Arsham et al., 2017).

It is of course always possible that you encounter a problem which is not covered by this Trouble Shooting Guide, or that you still have doubts about your results. In such cases, please do not hesitate to contact your local supplier or PanPath B.V. directly. Since we consider your problem as our problem, we will do our utmost to find a solution.

6.2 Cytological specimen

Problem	Possible causes	Remedies			
Weak or no signals	Sample preparation	→ Make sure samples are prepared according to protocol			
	Proteolytical pre- treatment	→ Make sure correct concentration of pre-treatment solutions is used			
		→ Make sure pre-treatments are performed at correct temperatures			
		→ Make sure dehydration steps are performed as recommended			
		→ Make sure that all pre-treatment steps are performed			
	Hybridisation procedure	→ Homogenize probe solution prior to applying probe on the section			
	PanWash 4 temperature and incubation time	→ Make sure correct temperatures are maintained and that incubation time of washing step is followed			
	Coverslips removed with force	→ Make sure that slides are soaked and coverslips are not removed with force			
	■ Air-dry omitted	→ Make sure that when air-dry is recommended, this is done properly			
	Air bubbles under the coverslip	→ Coverslip on a dark surface to visualise if any			
	■ Insufficient dehydration	→ Prepare fresh dehydration solutions			
	 Photo-damage to the fluorescently labelled probe 	→ Try to prevent prolonged light exposure and work in a dark environment if applicable			

PANPATH REMBRANDT® FISH DETECTION KITS



PAGE 16 OF 20

	■ No target sequence present	→ Use appropriate controls				
	 Microscope and accessories 	→ Check filter sets				
		ightarrow Check lamps				
		ightarrow Check running hours of lamp				
		→ Check lamp alignment				
		ightarrow Check and clean lenses and mirror				
		ightarrow Call microscope technical service				
		→ Use recommended objectives and filters as described in the product specific datasheet				
Non-specific background staining and	Cellular debris in cell preparation	Perform additional wash steps with fresh fixative to remove debris				
signal variation	■ Denaturation temperature too high	→ Make sure temperature is 80 °C				
	■ Denaturation step too long	→ Denature no longer than 5 minutes				
	■ Washing temperature	→ Make sure correct temperatures are maintained				
		→ Make sure pH is adjusted to 7.4				
	■ Proteolytic pre- treatment too strong	Make sure that the concentration of pepsin is as described in the product specific datasheet and do not exceed the incubation time in proteolytic work solution				

6.3 Paraffin embedded tissue sections

Problem	Possible causes	Remedies
No section left on the slides or morphology detected	■ Sample preparation	→ Make sure samples are prepared according to protocol, the tissue is fixed in neutral buffered formalin and the slides are air dried well
detected	■ Tissue section too thin	$ ightarrow$ Optimal thickness of the tissue is 2-4 μm
	Wrong (side of) glass slide used	→ Use only plus coated glass slides
	Pepsin concentration too high	→ Make sure correct concentration of pepsin is used.
	Proteolytic pre-treatment step too long	→ Reduce time of proteolytic pre-treatment step

PANPATH REMBRANDT® FISH DETECTION KITS



PAGE 17 OF 20

r		
	■ Denaturation	 → Make sure temperature is 80°C → Denature no longer than 10 minutes
	■ Coverslips removed with force	Make sure that slides are soaked and coverslips are not removed with force
	■ Post-fix solution omitted	→ Make sure post-fix incubation is performed
	■ Air-dry omitted	→ Make sure that when air-dry is recommended, this is done properly
Weak or no signals	■ Tissue fixation	→ Only use buffered formalin fixative and check fixation time (prevent over-fixation)
	■ Deparaffinisation	→ Renew dewax series
	■ Pre-treatment	→ Make sure correct concentration of pre-treatment solutions is used
		→ Make sure pre-treatments are performed at correct temperatures
		→ Make sure dehydration steps are performed as recommended
		→ Make sure that all pre-treatment steps are performed
	Denaturation	ightarrow Make sure temperature is 80 °C
	 Hybridisation procedure 	→ Homogenize probe solution prior to applying probe on the section
	PanWash 4 temperature and incubation time	→ Make sure correct temperatures are maintained and that incubation time of washing step is followed
	Air bubbles during denaturation and hybridisation	→ Make sure that when covering section with cover slip, air bubbles are not present
	Insufficient amount of probe used	→ Make sure that sufficient amount of probe is used to cover the section
	No target sequence present	→ Use appropriate controls
	 Photo-damage to the fluorescently labelled probe 	→ Try to prevent prolonged light exposure and work in a dark environment if applicable
	Microscope and accessories	→ Check filter sets
		→ Check lamps
		→ Check running hours of lamp

PANPATH REMBRANDT® FISH DETECTION KITS



→ Check lamp alignment						
		ightarrow Check and clean lenses and mirror				
		→ Call microscope technical service				
Non-specific background	■ Tissue section too thick	$ ightarrow$ Optimal thickness of the tissue is 2-4 μm				
staining and signal variation	■ Tissue crumbled	→ Make sure tissue is stretched completely				
Signar variation	■ Deparaffinisation	→ Dewax series				
	■ Denaturation temperature too high	→ Make sure temperature is 80 °C				
	 Hybridisation: uneven distribution of probe due to air bubbles/uneven section 	→ Repeat procedure on new section				
	■ Denaturation step too long	→ Denature no longer than 10 minutes				
	■ Drying out of the section	Incubate all procedure steps in a moisturised environment; prevent evaporation				
	■ Washing temperature	→ Make sure correct temperatures are maintained				
		→ Make sure pH is adjusted to 7.4				

One should always bear in mind that the staining intensity and the level of background (or nonspecific) staining may depend on the type of tissue used.

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Immaterial property information

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In situ Hybridisation and Detection

ISH detection assays-v1

ISH kit					
Label	Product number	# assays			
Biotin	CxxxK.0100.05	5 T			
Biotin	CxxxK.0100.10	10 T			
Digoxigenin	CxxxK.9900.05	5 T			
Digoxigenin	CxxxK.9900.10	10 T			
Biotin and Digoxigenin	CxxxK.0199.05	5 T			
Biotin and	CxxxK.0199.10	10 T			
Digoxigenin	CAXXII.0133.10	10 1			

Contents

			<u>page</u>
Chapter 1		Introduction	03
	1.1	Intended use	03
	1.2	The ISH principle	03
	1.3	Controls	03
	1.4	Contents of a REMBRANDT® ISH	03
	1.5	Materials required but not included	04
	1.6	Single product catalogue numbers, storage and shelf life	04
	1.7	Safety precautions	05
	1.8	Performance precautions	05
	1.9	Preparations of reagents in advance	06
	1.10	Preparations of reagents and materials needed but not supplied	06
	1.11	. Microscope and accessories	07
Chapter 2		REMBRANDT [®] ISH protocol	09
	2.1	Specimen collection	09
	2.2	Pre-treatment of specimen	09
	2.3	Hybridisation procedure and post-hybridization washes	10
Chapter 3		Interpretation of the results	12
	3.1	Guidelines for interpretation	12
	3.2	Quality control	
Chapter 4		Limitations of procedure	13
	4.1	Limitations	13
Chapter 5		References	15
Chapter 6		Trouble shooting guide	16
	6.1	Introduction	16
	6.2	-,	16
	6.3	Paraffin embedded tissue sections	17
Immateria	Ipro	operty information	20

Chapter 1 Introduction

1.1 Intended use

REMBRANDT has been designed for the detection of specific DNA or RNA sequences by using the *in situ* Hybridisation (ISH) technique in cytological specimen or FFPE tissue sections. In extensive tests and field application the REMBRANDT ISH kits were proven to be very robust, and ISH results were highly reproducible. For the specific intended use of a REMBRANDT ISH detection assay, please see the product specific datasheet.

1.2 The ISH principle

ISH enables the detection of specific DNA or RNA sequences in histological and cytological specimens, without losing the often very essential morphological details. The principle of ISH is based on a "reaction" (= hybridisation) between a specifically conjugated DNA or RNA sequence (= probe) and a DNA or RNA sequence present in the sample (= target). In case of matching sequences, a hybrid between the probe and target will be formed. Non-specific hybrids are washed out by the stringency wash procedure (PanWash 4). The formed hybrids can easily be visualized after detection with corresponding antibodies by fluorescent or brightfield microscopy. The REMBRANDT* probes are conjugated to biotin or digoxigenin; the conjugate used in the kits depend on specific probe(s) in the kit (Arsham et al., 2017).

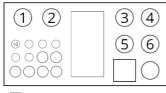
1.3 Controls

Use of controls is an essential part of the routine. To ensure that the ISH procedure is performed correctly and that observed positive and/or negative staining are specific, controls should be included in each experiment by the end-user.

1.4 Contents of a REMBRANDT® ISH detection assay

Application: cytological specimen

Contents REMBRANDT® ISH detection assays



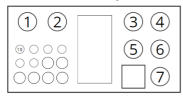
Pos.	Contents
1	Pepsin diluent
2	Pepsin powder
3	PanWash 4
4	PanWash 4
5	PanWash 4
6	PanWash 4
10	Product specific probe

Product label		Vol.		REF
DIGEST	PEPSIN DIL	15	ml.	R018R.0000
DIGEST	PEPSIN POW	1	gr.	R011R.0000
PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
PAN-WASH 4	SSC 25X	15	ml.	R025R.0000
PROBE	SPECIFIC PROBE	75 150	μl ul	XXXXXX.YYYY



Application: FFPE tissue sections

Contents REMBRANDT® ISH detection assays



Pos.	Contents
1	Pepsin diluent
2	Pepsin powder
3	PanWash 4
4	PanWash 4
5	PanWash 4
6	PanWash 4
7	Pre-treatment buffer

10 Product specific probe

Product label			Vol.		REF		
DIGEST	PEPSIN DIL		15	ml.	R018R.0000		
DIGEST	PEPSIN POW		1	gr.	R011R.0000		
PAN-WASH 4	SSC 25X		15	ml.	R025R.0000		
PAN-WASH 4	SSC 25X		15	ml.	R025R.0000		
PAN-WASH 4	SSC 25X		15	ml.	R025R.0000		
PAN-WASH 4	SSC 25X		15	ml.	R025R.0000		
PRE-TREAT	Na-Citrate 100x		15	ml.	R026R.0000		
PROBE	SPECIFIC PROBE		75	μl	XXXXXX.YYYY		
			150	μl			

1.5 Materials required but not included

- Xylene for dewaxing paraffin sections
- Fixative for cytological specimens
- Distilled or deionised water
- Ethanol series
- 100% Ethanol
- 96% Ethanol
- 70% Ethanol
- HCI 0.01 M

REF XX-YY

- Phosphate buffered saline (PBS); pH 7.2-7.4
- Pipettes and tips to deliver 10-1000 μL
- Immersion oil for fluorescence or brightfield microscopy
- Timer(s)

- Fluorescence microscope and accessories
- Oven(s), heating block(s) and water baths
- Incubation chamber
- Mixing device i.e. vortex
- Magnetic stirrer
- (Surface) thermometer; calibrated
- pH meter and calibrators
- Coverslips
- Slides
- Microwave (optional)

1.6 Single product catalogue number, storage and shelf life

REMBRANDT® probe mix; 1 vial			Catalogue No	CXXXP.XXXX	
Quantity	:	75 or 150 μl; 5 or 10 assays/kit	Storage	2 – 8°C	
Composition: specific conjugated probe DNA, ready to use in hybridisation buffer.					

Pepsin digestion reagent; 1 vial Catalogue I			Catalogue No	R011R.0000
Quantity	:	1 gr; powder to be dissolved in 8 mL deionised water, to be aliquoted in portions of e.g. 1200 μ l or 1200 μ l depending on the amount of slides per test run	Storage powder dissolved solution	2 – 25°C -20°C
Composition	:	Pepsin powder		

PANPATH REMBRANDT® ISH DETECTION KITS



PAGE 4 OF 20

Pepsin diluent; 1 vial		Catalogue No	R018R.0000
Quantity	: 15 mL 1M to be diluted 100x with deionised water to 0.01 M HCl	Storage 1 M HCI 0.01 M HCI	2 – 25°C 2 – 25°C
Composition	: HCI, 1M		

Concentrated PanWash 4, 25x SSC; 4 vials Catalogue No R025R.000			R025R.0000	
Quantity: 15 mL concentrated solution to be diluted 12.5x (= 2x SSC) with deionised water and confirm (and adjust if needed) to pH 7.4		Storage conc. solution diluted solution	2 – 25°C ambient temp.	
Composition	:	Sodium chloride, sodium citrate and surfactant		

Concentrated	Concentrated Pre-treatment buffer, 100x Na-Citrate; 1 vial (optional) Catalogue No R025R.0000			
Quantity	 15 mL concentrated solution to be diluted 100x (= 1x Na- Citrate) with deionised water and confirm (and adjust if needed) pH 6.0 	Storage conc. solution diluted solution	2 – 25°C ambient temp.	
Composition	: Natrium citrate			

- Store kit and its contents at 2-8°C.
- Store the dissolved and aliquoted reagents at recommended temperatures (see above).
- When used and stored as indicated, the kit is stable until the expiry date printed on the box.

1.7 Safety precautions

- The hybridisation buffer in which the probes are mixed contains formamide, which is considered as a teratogen; contact with skin and mucous membranes should be avoided.
- If reagents come into contact with skin or eyes, rinse with large volumes of clean water.
- Never pipette solutions by mouth.
- All hazardous materials should be disposed of according national guidelines.
- A material safety data sheet is available on request

1.8 Performance precautions

- Read all instructions before processing any assay.
- Store all reagents as recommended.
- **DO NOT** use reagents beyond their expiry date.
- Allow all components to warm up to room temperature (20-25°C) before use.
- -Check all ready to use solutions for precipitates and other contaminations before use; if contaminations are present, the solution should be disposed of, and a new solution should be prepared.
- PanWash 4 concentrated solution may form crystals when stored at low temperatures. If heating does not dissolve the crystals, the solution should be disposed of and a new solution should be prepared. Pan

PANPATH REMBRANDT® ISH DETECTION KITS

PAGE 5 OF 20

Path

- Homogenise all solutions before use.
- Homogenise probe solution and spin briefly before use.
- − DO NOT substitute a reagent with one from another manufacturer.
- Use treated glass slides, preferably REMBRANDT® Adhesive coated glass slides (Z000S.0002, PanPath B.V.).
- − **DO NOT** re-use prepared, ready to use digestion reagent; dispose of residuals.
- With every new procedure, check the temperatures of pre-treatment solutions, denaturation device used, incubation device used and PanWash 4 solution by using a calibrated thermometer.
- Do not incubate more than 5 specimens at the same time in pre-set temperature baths/devices.
 Placing more than 5 specimens in such a device will cause a temperature drop, and thus will not provide the correct temperature.
- Pre-treatment, Denaturation and Hybridisation are the key steps in the procedure, if any of these steps in the procedure have not been carried out properly, the results may not be relied upon.

1.9 Preparation of supplied reagents in advance

Pepsin stock solution:

Dissolve the pepsin digestion reagent in 8 mL distilled or deionised water (upon receipt of the kit). Aliquot in portions of i.e. 1200 or 600 μ L and store at- 20° C. This depends on the expectation of the amount of slides that are subject per test run.

Pepsin diluent:

Dilute the supplied pepsin diluent (1M HCl) 100x to 0.01 M HCl with distilled or deionised water.

Proteolytic work solution:

Prepare fresh work solution just before use and discard non-used solution:

- -Dilute aliquoted proteolytic reagent to 1.25 mg/mL for FFPE and 100 μ g/ml for cytological specimen in 0.01 M HCl.
- -Pre-heat 0.01 M HCl solution at 37°C, add the required pepsin stock solution just before use and mix.

Do not re-use

PanWash 4:

- Recommended for non-stringent washes:

Dilute a vial of the supplied 15 mL PanWash 4, 25x SSC to **2x SSC** with 172.5 mL deionised or distilled water to a total of 187.5 ml (concentration is 2x SSC).

Recommended for stringent washes:

Dilute 0.8 mL of a vial of the supplied PanWash 4, 25x SSC to **0.1x** SSC with 199.2 mL deionised or distilled water to a total of 200 ml (concentration is 0.1x SSC).

Before use:

a) Fill staining jars with SSC solution and pre-heat to the desired temperature.

Do not re-use

PANPATH REMBRANDT® ISH DETECTION KITS



PAGE 6 OF 20

Pre-treatment buffer (included depending assay type; see product specific datasheet): Dilute the supplied 15 mL Pre-treatment buffer, 100x Na-Citrate to 1x Na-Citrate with 1485 mL deionised or distilled water.

Do not re-use

1.10 Preparation of reagents and materials needed but not supplied

Dewaxing reagents

i.e. xylene, ethanol 100% and methanol

Do not re-use

Dehydration reagents

i.e. graded ethanol series (ethanol 70%-96%-100%)

PBS (phosphate buffered saline)/Tween® 20 (0.05%)

- Deionised water:

950 ml

- Sodium chloride (NaCL):

8 8

-Potassium chloride (KCI):

 $0.2\,$ g

- di-Sodium hydrogen phosphate di-hydrate (Na₂HPO₄.2H₂O):

1 78 8

- Potassium di-hydrogen phosphate (KH₂PO₄):

 $0.24 \, \mathrm{g}$

-Tween® 20

 $0.5 \, \text{ml}$

Adjust pH to 7.4 +/- 0.2 and adjust to 1000 mL with deionised water

Do not re-use

1.11 Microscope and accessories

Microscope

A fluorescence or brightfield microscope is needed to validate the *in situ* hybridisation results, depending on the application of the end-user.

Oculars and objectives

10X oculars are sufficient, 20X or 40X objectives are usually used for scanning the section. It is recommended to use at least a 63X oil immersion objective for final analysis.

Light source

The light source of regular microscopes is often 50 or 100 W. However a 100 W light source is recommended to obtain optimal respits

PANPATH REMBRANDT® ISH DETECTION KITS

PAGE 7 OF 20



Path

Immersion oil

The use of quality immersion oil is recommended; the immersion oil should be suited for use in fluorescence or brightfield microscopy, depending on the application of the end-user

Chapter 2 REMBRANDT® ISH Protocol

- <u>In case of detection with fluorescent antibodies:</u> All incubation steps should be performed in a closed (dark) incubation chamber/staining jar which contains a liquid (water) creating a saturated moisturised environment. During the incubation steps, evaporation of reagents should be prevented.
- Once the hybridisation procedure has been started the specimen should not be allowed to dry except for those procedural steps that mention "air-dry".
- Allow all reagents to reach the temperatures required for the respective incubation
- Work in a fume hood, use forceps and wear protective laboratory clothing and powder free examination gloves.

2.1 Specimen collection

Cytological specimen

Fixation in 70% ethanol is a commonly used for cytological specimen, it preserves the morphology and target DNA. Other fixation methods such as Carnoy's fixative are also applicable.

Paraffin embedded tissue sections

A standard procedure for tissue fixation and embedding usually involves the use of formalin and paraffin. The optimal tissue block size is 0.5 cm³. The formalin should be buffered and fixation times should (preferably) not exceed 12 hours. Excess and/or insufficient fixation may yield suboptimal morphology and target preservation. Embedding in paraffin should not exceed temperatures of 65°C.

Sample preparation: stretch 4 μ m paraffin sections on distilled water of 38-40°C without any additives and collect sections on bio-adhesive (i.e. organ silane) coated glass slides. Bake the slides at 65°C in a dry air oven for 1 hour. Slides can be used immediately or they can be stored at room temperature for up to 3 months.

Prior to ISH, slides need to be dewaxed in subsequent fresh xylene baths for 2 x 10 minutes. Incomplete removal of formalin and/or paraffin may affect the result of the procedure. Remove the xylene by placing the slides in subsequent 100% ethanol (2 x 5 minutes), hydrate in graded ethanol series, deionised water and continue with pre-treatment.

2.2 Pre-treatment of specimen

Cytological specimen

- Incubate slides in pre-heated proteolytic work solution at 37 °C (100 μg/ml) for 15 minutes.
- Flush wash in deionised water, followed by dehydration in graded ethanol series (ethanol 70%, 96%, 96%, 100% and 100%), 1 minutes each and air-dry slides for 15 minutes.

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Paraffin embedded tissue sections

For making the DNA accessible for the probes and to obtain reliable and reproducible results, pretreatment procedures are necessary.

- Place slides in jar filled with pre-treatment buffer (R026R.0000), place the jar in a microwave set at i.e. 900W and incubate up until boiling, subsequently reset microwave at 180W and proceed with the incubation for 10 minutes followed by a 20 minute cool down period at room temperature; all in the same solution. Flush wash slides in deionised water at room temperature.
- Incubate slides in pre-heated proteolytic work solution at 37 °C (1.25 mg/ml) for 15 minutes.
- Flush wash in deionised water, followed by dehydration in graded ethanol series (ethanol 70%, 96%, 96%, 100% and 100%), 1 minutes each and air-dry slides for 15 minutes.

Do not treat more than 5 slides at the same time, because the temperature of the pre-heated solutions may drop dramatically, thus causing incomplete pre-treatment. Additionally, allow the slides to air-dry as recommended; otherwise sections will be lost.

2.3 Hybridisation procedure and post-hybridization washes

Denaturation and Hybridisation

- Homogenize probe solution and spin briefly. Apply 10-15 μl of probe solution to each specimen. Cover all specimens with a cover slip (avoid air bubbles).
- Place slides on an 80 °C hotplate or other heating device and incubate for 3 minutes in case of cytological specimen and 10 minutes in case of FFPE tissue sections (denaturation).

Work in a pre-set order to ensure that all slides have been incubated at 80 °C for the exact same time! Do not denature more than 5 slides at the same time, because the temperature of the heating device may drop dramatically, thus causing incomplete denaturation.

Transfer slides into a moist environment and incubate for 16 hours at 37 °C.

Differentiation (stringent wash) and rinsing

- Remove coverslips by submerging the slides in PBS at room temperature. Soak the slides until the coverslips fall off.
- Incubate slides in pre-heated PanWash 4.

The concentration of PanWash 4 and the temperature may strongly depend on the detection system used. Recommend washes for non-stringent conditions are: 2x 5 minutes in 2x SSC at 42°C. For a stringent wash: 2x 5 minutes in 2x SSC at 42°C followed by 2x 5 minutes in 0.1x SSC at 61°C. Do not incubate more than 5 slides at the same time in PanWash 4 (2x SSC and 0.1x SSC), because the temperature of PanWash 4, may drop dramatically, causing wrong stringency conditions.

PANPATH REMBRANDT® ISH DETECTION KITS



PAGE 10 OF 20

- Incubate slides in PBS at room temperature for 1 minute.
- Dehydrate in graded ethanol series. Air-dry slides for 15 minutes (in the dark).

Recommended antibody detection systems

The appropriate detection system should be evaluated by the end-user. Recommended detection systems are listed below:

Digoxigenin detection	Biotin detection
R003R.0000	R041R.0000
REMBRANDT®	REMBRANDT®
Sheep aDig-AP conjugate	Goat aBio-AP Fab
	conjugate
R004R.0000	R042R.0000
REMBRANDT®	REMBRANDT®
Sheep aDig-HRP	Goat aBio-HRP Fab
conjugate	conjugate

After incubation with conjugated antibodies, use an appropriate detection system. Recommended chromogenic detection systems are listed below:

AP detection	HRP detection
R008R.0000	R007R.0000
REMBRANDT® NBT/BCIP	REMBRANDT® AEC
substrate	substrate
	+
	R010R.0000
	REMBRANDT® AEC
	buffer



Chapter 3 Interpretation of results

3.1 Guidelines for interpretation

- For specific interpretation guidelines and scoring criteria, see REMBRANDT® ISH detection product specific datasheet.
- For assessment it is important to select representative areas of the specimen.
- Over digestion can lead to DNA loss and affect the gene signal numbers in individual nuclei. Score
 only nuclei that have relatively intact nuclear borders.

3.2 Quality control

Positive control

It is advised to run a positive and negative control simultaneously with each ISH assay. As a positive control, one may also use validated in-house specimen. If the controls fail to demonstrate the expected staining, the result on the test specimen must be validated as invalid.

Invalid results

- In case the control does not stain adequately, the results of the test slides cannot be accepted. The
 run should be repeated in order to rule out procedural errors. Please check the trouble shooting
 section below for further possibilities.
- Poor cell morphology may lead to unacceptable results. In these cases we advise to assess the cell morphology quality in parallel brightfield stained specimen before performing a repeat test.
- If high background signals are present over the cytoplasm obscuring the true signals, the assay should be repeated.
- If hybridisation signals are not reliable/identifiable, the assay should be repeated.

For additional requirements, see the product specific datasheet.



Chapter 4 Limitations of Procedure

4.1 Limitations

- The REMBRANDT® ISH detection assays are solely applicable for the detection of corresponding DNA as described in the product specific intended use, which may be present in the test specimen.
- Medical decisions may not be taken based on this test. This is a research use only product.
- Sample fixation methods should be performed according to the recommended sample preparation methods.
- Many factors can influence the performance of the ISH procedure. Failure in detection can be
 due to i.e. improper sampling, handling, the time lapse between tissue removal and fixation, the
 fixation time, processing fixed specimen, the bio-adhesive on the slide, incubation times, pretreatment procedures, incubation temperatures, freezing, thawing, washing, heating drying,
 reagent contaminations and interpretation of results.
- The performance of REMBRANDT® ISH detection assays was tested and validated when using
 the exact procedure as listed in the product specific data sheets, modifications to the procedure
 may alter the performance characteristics.
- The REMBRANDT® ISH detection assays test results are not to be relied on in case the sampling, sampling method, quality, sample preparation, reagents used, controls and procedure followed is not optimal.
- The medical profession should be aware of risks and factors influencing the intensity, the absence or presence of ISH signals which cannot be foreseen when applying this test.
- The user should carefully consider the risk and use of sample material for this test in case the sample material does not contain sufficient or representative test material.
- Laboratory personnel performing the test should be knowledgeable, professional and be able to interpret the test results.
- The specific ISH detection assay should only be used for the loci the probe targets.



Product in combination with other devices

The REMBRANDT® ISH detection assays are intended for stand-alone usage. The assay is intended to be used in combination with standard formalin fixed or cytological specimen, hot plate(s), stove(s), incubation device(s), water bath(s), temperature and incubation time control(s), and other reagents (not supplied with this reagent) and a microscope. The combination has been tested and validated. Since the formalin fixed or cytological specimen, hot plate(s), stove(s), incubation device(s), water bath(s), temperature controls, incubation time control(s) and other not supplied reagents such as but not limited to fixation and other reagents and a microscope is not combined with the device as a product, conformity with the essential requirements is not applicable. Laboratory assay validation should always be established by the end-users.

Chapter 5 References

- Arsham, M. S., Barch, M. J., & Lawce, H. J. (2017). The AGT Cytogenetics Laboratory Manual The AGT Cytogenetics Laboratory Manual Edited by (Vol. 4).
- Duffy, L., Zhang, L., R., D., & M., A. (2012). Quality Control Considerations for Fluorescence In Situ Hybridisation of Paraffin-Embedded Pathology Specimens in a Diagnostic Laboratory Environment. Latest Research into Quality Control, Figure 1. https://doi.org/10.5772/51266
- Mascarello, J. T., Hirsch, B., Kearney, H. M., Ketterling, R. P., Olson, S. B., Quigley, D. I., Rao, K. W., Tepperberg, J. H., Tsuchiya, K. D., & Wiktor, A. E. (2011). Section E9 of the American College of Medical Genetics technical standards and guidelines: Fluorescence in situ hybridization. *Genetics in Medicine*, *13*(7), 667–675. https://doi.org/10.1097/GIM.0b013e3182227295

Chapter 6 Trouble shooting guide

6.1 Introduction

This Trouble Shooting Guide is intended to support you in obtaining optimal results with PanPaths REMBRANDT® ISH detection assays (Duffy et al., 2012) (Arsham et al., 2017).

It is of course always possible that you encounter a problem which is not covered by this Trouble Shooting Guide, or that you still have doubts about your results. In such cases, please do not hesitate to contact your local supplier or PanPath B.V. directly. Since we consider your problem as our problem, we will do our utmost to find a solution.

6.2 Cytological specimen

Problem	Possible causes	Remedies
Weak or no signals	■ Sample preparation	→ Make sure samples are prepared according to protocol
	Proteolytical pre- treatment	→ Make sure correct concentration of pre-treatment solutions is used
		→ Make sure pre-treatments are performed at correct temperatures
		→ Make sure dehydration steps are performed as recommended
		→ Make sure that all pre-treatment steps are performed
	■ Hybridisation procedure	→ Homogenize probe solution prior to applying probe on the section
	PanWash 4 temperature and incubation time	→ Make sure correct temperatures are maintained and that incubation time of washing step is followed
	Coverslips removed with force	→ Make sure that slides are soaked and coverslips are not removed with force
	■ Air-dry omitted	→ Make sure that when air-dry is recommended, this is done properly
	Air bubbles under the coverslip	→ Coverslip on a dark surface to visualise if any
	Insufficient dehydration	→ Prepare fresh dehydration solutions
	 Photo-damage to the fluorescently labelled probe 	→ Try to prevent prolonged light exposure and work in a dark environment if applicable
	■ No target sequence present	→ Use appropriate controls

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	■ Microscope and accessories	 → Check filter sets → Check lamps → Check running hours of lamp → Check lamp alignment → Check and clean lenses and mirror → Call microscope technical service → Use recommended objectives and filters as described in the product specific datasheet
Non-specific background staining and	Cellular debris in cell preparation	Perform additional wash steps with fresh fixative to remove debris
signal variation	■ Denaturation temperature too high	→ Make sure temperature is 80 °C
	■ Denaturation step too long	→ Denature no longer than 5 minutes
	■ Washing temperature	 → Make sure correct temperatures are maintained → Make sure pH is adjusted to 7.4
	■ Proteolytic pre- treatment too strong	Make sure that the concentration of pepsin is as described in the product specific datasheet and do not exceed the incubation time in proteolytic work solution

6.3 Paraffin embedded tissue sections

Problem	Possible causes	Remedies
No section left on the slides or morphology detected	■ Sample preparation	→ Make sure samples are prepared according to protocol, the tissue is fixed in neutral buffered formalin and the slides are air dried well
detected	■ Tissue section too thin	$ ightarrow$ Optimal thickness of the tissue is 2-4 μm
	Wrong (side of) glass slide used	→ Use only plus coated glass slides
	Pepsin concentration too high	→ Make sure correct concentration of pepsin is used.
	Proteolytic pre-treatment step too long	Reduce time of proteolytic pre-treatment step
	■ Denaturation	→ Make sure temperature is 80°C
		ightarrow Denature no longer than 10 minutes

PANPATH REMBRANDT® ISH DETECTION KITS



PAGE 17 OF 20

	T = 2	
	Coverslips removed with force	→ Make sure that slides are soaked and coverslips are not removed with force
	■ Post-fix solution omitted	→ Make sure post-fix incubation is performed
	■ Air-dry omitted	→ Make sure that when air-dry is recommended, this is done properly
Weak or no signals	■ Tissue fixation	→ Only use buffered formalin fixative and check fixation time (prevent over-fixation)
	■ Deparaffinisation	→ Renew dewax series
	■ Pre-treatment	→ Make sure correct concentration of pre-treatment solutions is used
		→ Make sure pre-treatments are performed at correct temperatures
		→ Make sure dehydration steps are performed as recommended
		→ Make sure that all pre-treatment steps are performed
	Denaturation	ightarrow Make sure temperature is 80 °C
	Hybridisation procedure	→ Homogenize probe solution prior to applying probe on the section
	PanWash 4 temperature and incubation time	→ Make sure correct temperatures are maintained and that incubation time of washing step is followed
	Air bubbles during denaturation and hybridisation	→ Make sure that when covering section with cover slip, air bubbles are not present
	Insufficient amount of probe used	→ Make sure that sufficient amount of probe is used to cover the section
	No target sequence present	→ Use appropriate controls
	Photo-damage to the fluorescently labelled probe	→ Try to prevent prolonged light exposure and work in a dark environment if applicable
	■ Microscope and	→ Check filter sets
	accessories	→ Check lamps
		→ Check running hours of lamp
		→ Check lamp alignment
		ightarrow Check and clean lenses and mirror
		→ Call microscope technical service



Non-specific background	■ Tissue section too thick	→ Optimal thickness of the tissue is 2-4 µm
staining and signal variation	■ Tissue crumbled	→ Make sure tissue is stretched completely
Signal variation	■ Deparaffinisation	→ Dewax series
	■ Denaturation temperature too high	→ Make sure temperature is 80 °C
	 Hybridisation: uneven distribution of probe due to air bubbles/uneven section 	→ Repeat procedure on new section
	■ Denaturation step too long	→ Denature no longer than 10 minutes
	■ Drying out of the section	Incubate all procedure steps in a moisturised environment; prevent evaporation
	■ Washing temperature	→ Make sure correct temperatures are maintained
		→ Make sure pH is adjusted to 7.4

One should always bear in mind that the staining intensity and the level of background (or nonspecific) staining may depend on the type of tissue used.

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