DATA SHEET-V4 REMBRANDT® LSI 3QTER FISH DETECTION RESEARCH USE ONLY (RUO)

Ref C733K.3000.05 \(\overline{\sqrt{2}} \) 5 \\ C733K.3000.10 \(\overline{\sqrt{2}} \) 10 \\ T

Intended use

- The REMBRANDT® LSI 3qter FISH detection assay is for research use only and is intended for the detection of the 3q sub telomeric region by means of in situ hybridization.
- II. The REMBRANDT® LSI 3qter FISH detection assay is intended for the detection of the 3q sub telomeric region in fixed cells. A clinical diagnosis should not be established based on the performance of this test.
- III. The REMBRANDT® LSI 3qter FISH detection assay kit is a quantitative assay for the detection of the 3q sub telomeric region.
- IV. The intended users are qualified laboratory employees in cytology and/or pathology. The product is intended for professional use.

Clinical relevance

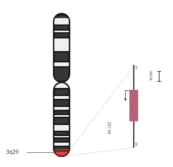
The 3g sub telomeric probe covers the 3g29 chromosomal location. This region harbors genes which are suspected to be tumor suppressor genes (TSGs), due to the association of 3g genetic aberrations with squamous cell lung cancer. non-small cell lung cancer, and other cancers (Chen et al., 2021; Zabarovsky et al., 2002). The loss of 3q is increasingly associated with (primary) breast cancer (Andersen et al., 1992; Yang et al., 2002) and with (early) invasive breast cancer (Martinez, 2001), and also with renal cell carcinoma (Favazza et al., 2017; Gatto et al., 2014; Hsieh et al., 2018). The most important TSGs associated with tumor-related pathways of the 3g-region are ALS2 C-Terminal Like (ALS2CL), EPH Receptor A3 (EPHA3) and xin actin binding repeat containing 1 (XIRP1) (Lee et al., 2010). The ALS2CL gene is associated with head and neck squamous cell carcinoma and breast cancer (Kim et al., 2021: Li et al., 2015), whereas EPHA3 gene is associated with (small cell) lung cancer (Peng et al., 2016; Zhuang et al., 2012), and XIRP1 gene is mainly associated with head and neck squamous cell carcinoma. The REMBRANDT LSI 3ater FISH detection assay is designed to detect the sub telomeric region, 3q29.





Probe specification

The REMBRANDT®LSI 3qter probe mix consists of a 231 kb probe and is available in an orange fluorescent detection (AF555). The REMBRANDT® LSI 3qter probes are premixed in a hybridization mixture (formamide, dextran sulphate and SSC) and are ready to use solutions.



Test principle

In a fluorescent in situ hybridization assay, a double stranded DNA probe labelled with a fluorochrome is used. The labelled DNA probe is diluted in a hybridization mixture. The hybridization mixture containing the DNA probe is added to the specimen. A co-denaturation will ensure that the genomic DNA and the probe DNA become single stranded. After denaturation, the probe DNA is able to hybridize to its complementary target sequences in the cells. In the REMBRANDT® LSI 3qter FISH detection assay, the fluorochrome is attached to the probe and the signals can be visualized directly by fluorescent microscopy after hybridization.

Reagents provided

Product name	Product number	Amount
Labelled LSI probe (depen REMBRANDT® LSI 3qter orange FISH probe mix	ding on size choice) C733P.3000.05 or C733P.3000.10	5 T 2 10 T
REMBRANDT® Pepsin powder	R011R.0000	1 g
REMBRANDT® Pepsin diluent	R018R.0000	15 ml
REMBRANDT® PanWash 4, 25X SSC	R025R.0000	4x 15 ml
REMBRANDT® Fluorescent Mounting	Z000R.0050	1 ml

Assay procedure

REMBRANDT® LSI 3qter FISH detection assay procedure for cytological specimen.

- Incubate slides in pre-heated proteolytic work solution (prepare according to section 1.9 of Manual-FISH) (R011R.000 + R018R.000) at 37 °C (100 µg/ml) for 15 minutes followed by a brief rinsing in 0.01M HCI (1x 2 minutes) and subsequent rinses in PBS (2x 1 minute)
- II. Flush wash slides in deionised water, followed by dehydration in graded ethanol series (ethanol 70%, 96%, 96%, 100%, 100%) 1 minute 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.

III. Homogenize probe solution (C733P.2030.YY) a spin briefly. Apply 15 μl of probe solution to each specimen. Cover all specimens with a cover slip (avoid air bubbles). Denature slides on a 80 °C hotplate or other heating device for 3 minutes

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, the temperature of the heating device may drop dramatically, thus causing incomplete denaturation.

- IV. Transfer the slides into a moist environment and incubate for 16 hours at 37 °C.
- V. Remove coverslips by soaking the slides in PBS at room temperature
- VI. Incubate the slides in diluted, pre-heated PanWash 4 (R025R.000) (prepare according to section 1.9 of Manual-FISH)

Do not incubate more than 5 slides at the same time in PanWash 4, the temperature of the PanWash 4 may drop dramatically, causing wrong stringency conditions.

- VII. Incubate the slides in PBS at room temperature for 1 minute
- VIII. Dehydrate the slides in graded ethanol series (70%, 96%, 96%, 100%, 100%) 1 minute each and air-dry the slides for 15 minutes (in the dark)

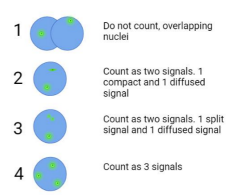
Mount the slides by applying mounting medium (Z000R.0050) and coverslip

Interpretation of results

Hybridization of the REMBRANDT® LSI 3gter probe is viewed using a fluorescence microscope equipped with appropriate excitation and emission filters for orange detection: λ_{exc} 555 nm, λ_{em} 572 nm. Allowing visualization of orange fluorescent signal concentrated at the 3q sub telomeric region and the blue counterstained chromosomes and nuclei. The enumeration of the 3g sub telomeric region is conducted by microscopic examination of interphase nuclei. The fluorescently-stained 3q sub telomeric regions stand out brightly against the general fluorescence of the nucleus. The LSI 3qter procedure enables visual enumeration of copy numbers of the 3g sub telomeric region within the nuclei. The assay results are reported as the percentage of nuclei with 0, 1, 2, 3, 4, and >4 fluorescent signals. Each fluorescent signal corresponds to a copy of the 3q sub telomeric region.

Enumerate the fluorescent signals in the interphase nucleus using a 40X or 63X magnification. Objectives with higher magnification (eg, 63X or 100X) should be used to verify or resolve questions about split or diffused signals. Enumerate at least 100 nuclei per slide for accurate analysis.

- •Two signals in close proximity and approximately the same sizes but not connected by a visible link are counted as 2 signals.
- •Count a diffuse signal as 1 signal if diffusion of the signal is contiguous and within an acceptable boundary.
- •Two small signals connected by a visible link are counted as 1 signal.
- •Enumerate the number of nuclei with 0, 1, 2, 3, 4, or >4 signals. Count nuclei with 0 signals only if there are other nuclei with at least 1 signal present in the field of view. If the accuracy of the enumeration is in doubt, repeat the enumeration in another area of the slide.
- •Do not enumerate nuclei with uncertain signals (Arsham et al., 2017)



Performance characteristics Analytical Sensitivity and Specificity

The analytical sensitivity and specificity were investigated within PanPaths analytical performance assessment. Precision was investigated for the REMBRANDT®LSI 3qter FISH detection assay and results are available upon request.

Analytical sensitivity

The analytical sensitivity was determined in three levels. The normal cut-off percentage was determined based on assessment of 200 individual nuclei in two samples of healthy interphase lymphocytes from peripheral blood. The beta inversion was used to determine the percentage of normal-cut off. For the noise-to-signal percentage, the noise and signal values were determined for 100 signals in interphase lymphocytes from peripheral blood of two independent samples. The differences between noise and signal were evaluated using a 95% confidence interval. For the hybridization efficiency, 200 individual nuclei were assessed for the presence of FISH signals.

Performance characteristic Normal cut-off percentage	Outcome 10%
Noise-to-signal cut-off	13%
percentage Hybridization efficiency	99%

Analytical specificity

The analytical specificity was determined in two levels. The theoretical specificity was determined by sequencing analysis of the probe DNA and mapping on Hg38. The practical specificity was determined by assessing the hybridization pattern in metaphase chromosomes of interphase lymphocytes from peripheral blood.

Performance characteristic	Outcome
Theoretical specificity	Mapped on chromosome 3,
	qter
Practical specificity	100%

Limitations of Procedure

- i) The REMBRANDT® LSI 3qter FISH detection assay is solely applicable for the detection of the 3q sub telomeric region, which may be present in cell preparations (cytological specimen i.e. interphase lymphocytes from peripheral blood samples).
- ii) Either human tissue sections or human cytological preparations may be used. Samples must be fixed in buffered formalin or alcohol. In tissue sections are required, the sections should be prepared in a 4 µm thickness. Furthermore, the tissues should be glued to the glass slides with a bio-adhesive (e.g. organ silane), dried at room temperature, subsequently dried at 37 °C overnight and lastly completely deparaffinized in xylene and alcohol series and air dried.

- Cytological specimen should be prepared as required by the user, fixed with cytological fixation agent, rinsed in distilled water prior to the ISH procedure and air dried.
- iv) 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 size of the tissue specimen in the fixation medium, the fixation time, processing fixed tissue, the thickness of the section, the bio-adhesive on the slide, deparaffinisation procedure, incubation times, proteolytic pre-treatment, detection reagents, incubation temperatures and interpretation of results.
- v) The performance of the ISH procedure is also affected by the sensitivity of the method and the presence of the 3q sub telomeric region. In case the limit of the sensitivity is reached a false negative reaction may be the result.
- vi) The REMBRANDT®LSI 3qter FISH detection assay results should not be relied on in case the sampling, sampling method, sample quality, sample preparation, reagents used, controls and procedure followed are not optimal or as described the working protocols.
- ix) The medical profession should be aware of risks and factors influencing the fluorescent signal intensity while interpretating the test result. Microscopy settings might influence the signal intensity and/or interpretation.
- x) Laboratory personnel performing the test should be trained and knowledgeable to be able to interpret the test results

Storage and handling

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

Product	Product number	Storage conditions	
REMBRANDT® LSI 3qter orange FISH probe mix	C733P.XXXX	2-8 °C	
REMBRANDT® Pepsin powder	R011R.0000	Powder: 2- 25°C, ambient temperature	
		Dissolved: - 20°C	
REMBRANDT® Pepsin diluent	R018R.0000	Concentrated solution and diluted: 2-25°C, ambient temperature	
REMBRANDT® PanWash 4, 25X SSC	R025R.0000	Concentrated solution and diluted: 2-25°C, ambient temperature	

REMBRANDT® Z000R.0050 2-8 °C Fluorescent mounting medium



Hazard statements

H315 - Causes skin irritation

H319 - Causes serious eye irritation

H351 - Suspected of causing cancer

H360D - May damage the unborn child

H373 - May cause damage to organs through prolonged or

repeated exposure

Precautionary Statements

P202 - Do not handle until all safety precautions have been read and understood

P280 - Wear protective gloves/protective clothing/eye protection/face protection

P302 + P352 - IF ON SKIN: Wash with plenty of water and soap P305 + P351 + P338 - IF IN EYES: Rinse cautiously with water for

several minutes. Remove contact lenses, if present and easy to do. Continue rinsing

P308 + P313 - IF exposed or concerned: Get medical advice/attention

P362 + P364 - Take off contaminated clothing and wash it before reuse

P405 - Store locked up

Additional information

Product in combination with other devices

The REMBRANDT® DNA probes are intended for standalone usage. The assay is intended to be used in combination with standard cytological preparation methods, hot plate(s), stove(s), incubation device(s), water bath(s), temperature and incubation time control(s), proteolytic-, detection- and other reagents (not supplied with this reagent) and a microscope. The combination has been tested and validated. Since the standard cytological preparation methods, 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 proteolytic reagents, detection reagents and a microscope is not combined with the device as a product, conformity with the essential requirements is not applicable. Assay validation criteria are mentioned in 'Interpretation of the Results' and are also depending on clinical state of the sample, which may influence the validation criteria.

For additional information regarding the REMBRANDT® assays, a manual is included which specifies the following subjects:

- Controls
- Materials required but not included
- Storage and shelf-life
- Performance precautions
- Preparations of reagents
- Specimen collection
- Quality control
- Trouble shooting guide

Technical assistance

For technical assistance regarding the products performance, please contact info@panpath.nl or call +31 495499090. Visit our website for reprints of datasheets or additional documentation. www.panpath.nl

Literature list

Arsham, M. S., Barch, M. J., & Lawce, H. J. (2017). The AGT Cytogenetics Laboratory Manual The AGT Cytogenetics Laboratory Manual Edited by (Vol. 4).

Andersen, T. I., Gaustad, A., Børresen, A.-L., Farrants, G. W., Nesland, J. M., Ottestad, L., & Tveit, K. M. (1992). Genetic alterations of the tumour suppressor gene regions 3p, 11p, 13q, 17p, and 17q in human breast carcinomas. *Genes, Chromosomes and Cancer, 4*(2), 113–121. https://doi.org/10.1002/gcc.2870040203

Arsham, M. S., Barch, M. J., & Lawce, H. J. (2017). The AGT Cytogenetics Laboratory Manual The AGT Cytogenetics Laboratory Manual Edited by (Vol. 4).

Chen, C., Pan, Y., Bai, L., Chen, H., Duan, Z., Si, Q., Zhu, R., Chuang, T.-H., & Luo, Y. (2021). MicroRNA-3613-3p functions as a tumor suppressor and represents a novel therapeutic target in breast cancer. *Breast Cancer Research*, 23(1), 12. https://doi.org/10.1186/s13058-021-01389-9

Favazza, L., Chitale, D. A., Barod, R., Rogers, C. G., Kalyana-Sundaram, S., Palanisamy, N., Gupta, N. S., & Williamson, S. R. (2017). Renal cell tumors with clear cell histology and intact VHL and chromosome 3p: a histological review of tumors from the Cancer Genome Atlas database. *Modern Pathology*, 30(11), 1603–1612. https://doi.org/10.1038/modpathol.2017.72

Gatto, F., Nookaew, I., & Nielsen, J. (2014). Chromosome 3p loss of heterozygosity is associated with a unique metabolic network in clear cell renal carcinoma. *Proceedings of the National Academy of Sciences*, 111(9). https://doi.org/10.1073/pnas.1319196111

Hsieh, J. J., Le, V. H., Oyama, T., Ricketts, C. J., Ho, T. H., & Cheng, E. H. (2018). Chromosome 3p

- Loss–Orchestrated VHL, HIF, and Epigenetic Deregulation in Clear Cell Renal Cell Carcinoma. *Journal of Clinical Oncology*, 36(36), 3533–3539. https://doi.org/10.1200/JCO.2018.79.2549
- Kim, H. A. J., Shaikh, M. H., Lee, M., Zeng, P. Y. F., Sorgini, A., Akintola, T., Deng, X., Jarycki, L., Khan, H., MacNeil, D., Khan, M. I., Mendez, A., Yoo, J., Fung, K., Lang, P., Palma, D. A., Patel, K., Mymryk, J. S., Barrett, J. W., ... Nichols, A. C. (2021). 3p Arm Loss and Survival in Head and Neck Cancer: An Analysis of TCGA Dataset. Cancers, 13(21), 5313. https://doi.org/10.3390/cancers13215313
- Lee, D. J., Schönleben, F., Banuchi, V. E., Qiu, W., Close, L. G., Assaad, A. M., & Su, G. H. (2010). Multiple tumor-suppressor genes on chromosome 3p contribute to head and neck squamous cell carcinoma tumorigenesis. *Cancer Biology & Therapy*, 10(7), 689–693. https://doi.org/10.4161/cbt.10.7.12886
- Li, H., Yang, B., Xing, K., Yuan, N., Wang, B., Chen, Z., He, W., & Zhou, J. (2015). A preliminary study of the relationship between breast cancer metastasis and loss of heterozygosity by using exome sequencing. Scientific Reports, 4(1), 5460. https://doi.org/10.1038/srep05460
- Martinez, A. (2001). Chromosome 3p allele loss in early invasive breast cancer: detailed mapping and association with clinicopathological features. *Molecular Pathology*, 54(5), 300–306. https://doi.org/10.1136/mp.54.5.300
- Peng, J., Wang, Q., Liu, H., Ye, M., Wu, X., & Guo, L. (2016). EPHA3 regulates the multidrug resistance of small cell lung cancer via the PI3K/BMX/STAT3 signaling pathway. *Tumor Biology*, 37(9), 11959– 11971. https://doi.org/10.1007/s13277-016-5048-4
- Yang, Q., Yoshimura, G., Mori, I., Sakurai, T., & Kakudo, K. (2002). Chromosome 3p and breast cancer. Journal of Human Genetics, 47(9), 453–459. https://doi.org/10.1007/s100380200064
- Zabarovsky, E. R., Lerman, M. I., & Minna, J. D. (2002). Tumor suppressor genes on chromosome 3p involved in the pathogenesis of lung and other cancers. *Oncogene*, 21(45), 6915–6935. https://doi.org/10.1038/sj.onc.1205835
- Zhuang, G., Song, W., Amato, K., Hwang, Y., Lee, K., Boothby, M., Ye, F., Guo, Y., Shyr, Y., Lin, L., Carbone, D. P., Brantley-Sieders, D. M., & Chen, J. (2012). Effects of Cancer-Associated EPHA3 Mutations on Lung Cancer. JNCI: Journal of the National Cancer Institute, 104(15), 1183–1198. https://doi.org/10.1093/jnci/djs297

Disclaimer: This document is valid until the product expiry on the kit label

DATA SHEET-V3 REMBRANDT® LSI 3QTER-ISH DETECTION RESEARCH USE ONLY (RUO)

Intended use

- I. The REMBRANDT® LSI 3qter-ISH detection assay is for research use only and is intended for the detection of the human 3q sub telomeric region by means of in situ hybridization.
- II. The REMBRANDT® LSI 3qter-ISH detection assay is intended for the detection of the 3q sub telomeric region in fixed cells. A clinical diagnosis should not be established based on the performance of this test.
- III. The REMBRANDT® LSI 3qter-ISH detection assay kit is a quantitative assay for the detection of the 3q sub telomeric region.
- IV. The intended users are qualified laboratory employees in cytology and/or pathology. The product is intended for professional use.

Clinical relevance

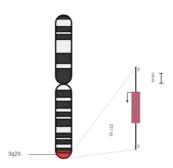
The 3g sub telomeric probe covers the 3g29 chromosomal location. This region harbors genes which are suspected to be tumor suppressor genes (TSGs), due to the association of 3g genetic aberrations with squamous cell lung cancer, non-small cell lung cancer, and other cancers (Chen et al., 2021; Zabarovsky et al., 2002). The loss of 3q is increasingly associated with (primary) breast cancer (Andersen et al., 1992; Yang et al., 2002) and with (early) invasive breast cancer (Martinez, 2001), and also with renal cell carcinoma (Favazza et al., 2017; Gatto et al., 2014; Hsieh et al., 2018). The most important TSGs associated with tumor-related pathways of the 3g-region are ALS2 C-Terminal Like (ALS2CL), EPH Receptor A3 (EPHA3) and xin actin binding repeat containing 1 (XIRP1) (Lee et al., 2010). The ALS2CL gene is associated with head and neck squamous cell carcinoma and breast cancer (Kim et al., 2021; Li et al., 2015), whereas EPHA3 gene is associated with (small cell) lung cancer (Peng et al., 2016; Zhuang et al., 2012), and XIRP1 gene is mainly associated with head and neck squamous cell carcinoma. The REMBRANDT LSI 3gter-ISH detection assay is designed to detect the sub telomeric region, 3g29.





Probe specification

The REMBRANDT® LSI 3qter probe mix consists of a 231 kb probe and is available in a digoxigenin or biotin conjugation. The REMBRANDT® LSI 3qter probes are pre-mixed in a hybridization mixture (formamide, dextran sulphate and SSC) and are ready to use solutions.



Test principle

In an *in situ* hybridization assay, a double stranded DNA probe labelled with a hapten is used. The labelled DNA probe is diluted in a hybridization mixture. The hybridization mixture containing the DNA probe is added to the specimen. A co-denaturation will ensure that the genomic DNA and the probe DNA become single stranded. After denaturation, the probe DNA is able to hybridize to its complementary target sequences in the cells. In the REMBRANDT® LSI 3qter-ISH detection assay, the haptens are attached to the probe and the signals can be visualized after detection by corresponding antibodies by fluorescent or brightfield microscopy.

Reagents provided

iteageille provid	c u	
Product name	Product	Amount
	number	
Labelled LSI probe (depe		ize choice)
REMBRANDT® LSI	C733P.0100.05	Σ/ _{5.T}
3qter-ISH biotin probe	or	V 5 1
mix	C733P.0100.10	∑ 10 T
REMBRANDT® LSI	C733P.9900.05	$\sqrt{\Sigma}$
3qter-ISH digoxigenin	or	_ <u>∨</u> 5 T
probe mix	C733P.9900.10	Σ/10.Τ
REMBRANDT® Pepsin	R011R.0000	1 g
powder		
REMBRANDT® Pepsin	R018R.0000	15 ml
diluent		
REMBRANDT®	R025R.0000	4x 15 ml
PanWash 4, 25X SSC		
REMBRANDT® Pre-	R026R.0000	15 ml
treatment buffer		

Assay procedure

REMBRANDT® LSI 3qter-ISH detection assay procedure for cytological specimen and FFPE tissue sections.

- Specimen collection: for a detailed description of the specimen collection for cytological specimen or FFPE tissue sections see section 2.1 Specimen collection of the Manual ISH
- II. For FFPE tissue sections, after dewaxing, place slides in jar with pre-treatment solution (R026R.0000) in microwave set at i.e. 900W and incubate up until boiling. Subsequently, reset microwave at 180W and incubate for 10 minutes, followed by cooling down for 20 minutes at room temperature. Flush wash slides in deionised water.
- III. Incubate slides in pre-heated proteolytic work solution (prepare according to section 1.9 and 1.10 of Manual ISH (R011R.000 + R018R.000) at 37 °C. Paraffin-embedded sections (1.25 mg/ml) or cytological specimen (100 µg/ml) for 15 minutes.
- IV. Flush wash slides in deionised water, followed by dehydration in graded ethanol series (ethanol 70%, 96%, 96%, 100%, 100%) 1 minute 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.

V. Homogenize probe solution (C801P.0199.YY) and spin briefly. Apply 10-15 µl of probe solution to each specimen. Cover all specimens with a cover slip (avoid air bubbles). Denature slides on a 80 °C hotplate or other heating device, 3 minutes for cytological specimen and 10 minutes for FFPE tissue sections.

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, the temperature of the heating device may drop dramatically, thus causing incomplete denaturation.

- VI. Transfer the slides into a moist and dark environment and incubate for 16 hours at 37
- VII. Remove coverslips by soaking the slides in PBS at room temperature.
- VIII. Incubate the slides in diluted, pre-heated PanWash 4 (R025R.0000) (prepare according

to section 1.9 of Manual-ISH). For cytological specimen and FFPE tissue sections, 2x 5 minutes in 2x SSC at 42 °C. For cytological specimen, subsequently incubate 2x 5 minutes in 0.1x SSC at 61 °C.

Do not incubate more than 5 slides at the same time in PanWash 4, the temperature of the PanWash 4 may drop dramatically, causing wrong stringency conditions.

IX. Appropriate detection system should be evaluated by the end-user. Recommended detection systems are listed below

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

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

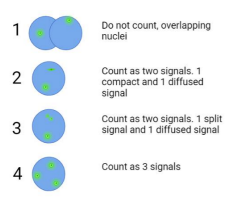
Interpretation of results

Hybridization of the LSI 3qter probe is conducted by microscopic examination of interphase nuclei (fluorescence or brightfield, depending on antibodies used for detection). The fluorescently or chromogenic-stained 3q sub telomeric regions stand out brightly against the nucleus. The enumeration of the 3q sub telomeric region is conducted by microscopic examination of interphase nuclei. The LSI 3qter procedure enables visual enumeration of copy numbers of the 3q sub telomeric region within the nuclei. The assay results are reported as the percentage of nuclei with 0, 1, 2, 3, 4, and >4 signals. Each signal corresponds to a copy of the 3q sub telomeric region.

Enumerate the signals in the interphase nucleus using a 40X or 63X magnification. Objectives with higher magnification (eg, 63X or 100X) should be used to verify or resolve questions about split or diffused signals. Enumerate at least 100 nuclei per slide for accurate analysis.

- •Two signals in close proximity and approximately the same sizes but not connected by a visible link are counted as 2 signals.
- •Count a diffuse signal as 1 signal if diffusion of the signal is contiguous and within an acceptable boundary.

- •Two small signals connected by a visible link are counted as 1 signal.
- •Enumerate the number of nuclei with 0, 1, 2, 3, 4, or >4 signals. Count nuclei with 0 signals only if there are other nuclei with at least 1 signal present in the field of view. If the accuracy of the enumeration is in doubt, repeat the enumeration in another area of the slide.
- •Do not enumerate nuclei with uncertain signals



Performance characteristics

The REMBRANDT® LSI 3qter-ISH detection assay was analytically validated for REMBRANDT® LSI 3qter orange detection. The results of the direct fluorescent assay are shown. However, for the REMBRANDT® LSI 3qter-ISH assay, the detection system may influence the performance characteristics and the REMBRANDT® LSI 3qter in combination with different detection systems should be evaluated carefully by the end-user.

Analytical sensitivity

The analytical sensitivity was determined in three levels. The normal cut-off percentage was determined based on assessment of 200 individual nuclei in two samples of healthy interphase lymphocytes from peripheral blood. The beta inversion was used to determined the percentage of normal-cut off. For the noise-to-signal percentage, the noise and signal values were determined for 100 signals in interphase lymphocytes from peripheral blood of two independent samples. The differences between noise and signal were evaluated using a 95% confidence interval. For the hybridization efficiency, 200 individual nuclei were assessed for the presence of ISH signals.

Performance characteristic	Outcome
Normal cut-off percentage	10%
Noise-to-signal cut-off percentage	13%
Hybridization efficiency	99%

Analytical specificity

The analytical specificity was determined in two levels. The theoretical specificity was determined by sequencing analysis of the probe DNA and mapping on Hg38. The practical specificity was determined by assessing the hybridization pattern in metaphase chromosomes of interphase lymphocytes from peripheral blood.

Performance characteristic	Outcom	е		
Theoretical specificity	Mapped gter	on	chromosome	3,
Practical specificity	100%			

Limitations of Procedure

- i) The REMBRANDT® LSI 3qter-ISH detection assay is solely applicable for the detection of the 3q sub telomeric region, which may be present in cell preparations (cytological specimen i.e. interphase lymphocytes from peripheral blood samples).
- ii) Either human tissue sections or human cytological preparations may be used. Samples must be fixed in buffered formalin or alcohol. In tissue sections are required, the sections should be prepared in a 4 µm thickness. Furthermore, the tissues should be glued to the glass slides with a bio-adhesive (e.g. organ silane), dried at room temperature, subsequently dried at 37 °C overnight and lastly completely deparaffinized in xylene and alcohol series and air dried.
- iii) Cytological specimen should be prepared as required by the user, fixed with cytological fixation agent, rinsed in distilled water prior to the ISH procedure and air dried
- iv) 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 size of the tissue specimen in the fixation medium, the fixation time, processing fixed tissue, the thickness of the section, the bio-adhesive on the slide, deparaffinisation procedure, incubation times, proteolytic pre-treatment, detection reagents, incubation temperatures and interpretation of results.
- v) The performance of the ISH procedure is also affected by the sensitivity of the method and the presence of the 3q sub telomeric region. In case the limit of the sensitivity is reached a false negative reaction may be the result.
- vi) The REMBRANDT® LSI 3qter-ISH detection assay results should not be relied on in case the sampling, sampling method, sample quality, sample preparation, reagents used, controls and procedure followed are not optimal or as described the working protocols.
- vii) The medical profession should be aware of risks and factors influencing the fluorescent signal intensity while interpretating the test result. Microscopy settings might influence the signal intensity and/or interpretation.

Laboratory personnel performing the test should be trained and knowledgeable to be able to interpret the test results.

Storage and handling

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

Product REMBRANDT® LSI 3qter-ISH probe mix	Product number C733P.XXXX	Storage conditions 2-8 °C
REMBRANDT® PanWash 4, 25X SSC	R025R.0000	Concentrated solution and diluted: 2-25°C, ambient temperature
REMBRANDT® Pepsin powder	R011R.0000	Powder: 2-25°C, ambient temperature
REMBRANDT® Pepsin diluent	R018R.0000	Dissolved: -20°C Concentrated solution and diluted: 2-25°C, ambient temperature
REMBRANDT® Pre-treatment buffer	R026R.0000	Concentrated solution and diluted: 2-25°C, ambient temperature



Hazard statements

H315 - Causes skin irritation

H319 - Causes serious eye irritation

H351 - Suspected of causing cancer

H360D - May damage the unborn child

H373 - May cause damage to organs through prolonged or

repeated exposure

Precautionary Statements

P202 - Do not handle until all safety precautions have been read and understood

P280 - Wear protective aloves/protective clothing/eve protection/face protection

P302 + P352 - IF ON SKIN: Wash with plenty of water and soap P305 + P351 + P338 - IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsina

P308 + P313 - IF exposed or concerned: Get medical

P362 + P364 - Take off contaminated clothing and wash it before reuse

P405 - Store locked up

Additional information

Product in combination with other devices

The REMBRANDT® DNA probes are intended for standalone usage. The assay is intended to be used in combination with standard cytological preparation methods, hot plate(s), stove(s), incubation device(s), water bath(s), temperature and incubation time control(s), proteolytic-, detection- and other reagents (not supplied with this reagent) and a microscope. The combination has been tested and validated. Since the standard formalin fixed, paraffin embedded tissue blocks, standard tissue freezing, tissue sectioning (microtome), standard cytological preparation methods, 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 proteolytic reagents, detection reagents and a microscope is not combined with the device as a product, conformity with the essential requirements is not applicable. Assay validation criteria are mentioned in 'Interpretation of the Results' and are also depending on the target load, which may influence the validation criteria

For additional information regarding the REMBRANDT® assays, a manual is included which specifies the following subjects:

- Controls
- Materials required but not included
- Storage and shelf-life
- Performance precautions
- Preparations of reagents
- Specimen collection
- Quality control
- Trouble shooting guide

Technical assistance

For technical assistance regarding the products performance, please contact info@panpath.nl or call +31 495499090. Visit our website for reprints of datasheets or additional documentation. www.panpath.nl

Literature list

Arsham, M. S., Barch, M. J., & Lawce, H. J. (2017). The AGT Cytogenetics Laboratory Manual The AGT Cytogenetics Laboratory Manual Edited by (Vol. 4).

Andersen, T. I., Gaustad, A., Børresen, A.-L., Farrants, G. W., Nesland, J. M., Ottestad, L., & Tveit, K. M. (1992). Genetic alterations of the tumour suppressor gene regions 3p, 11p, 13q, 17p, and 17q in human breast carcinomas. Genes, Chromosomes and Cancer, 4(2), 113-121. https://doi.org/10.1002/gcc.2870040203

- Arsham, M. S., Barch, M. J., & Lawce, H. J. (2017). The AGT Cytogenetics Laboratory Manual The AGT Cytogenetics Laboratory Manual Edited by (Vol. 4).
- Chen, C., Pan, Y., Bai, L., Chen, H., Duan, Z., Si, Q., Zhu, R., Chuang, T.-H., & Luo, Y. (2021). MicroRNA-3613-3p functions as a tumor suppressor and represents a novel therapeutic target in breast cancer. *Breast Cancer Research*, 23(1), 12. https://doi.org/10.1186/s13058-021-01389-9
- Favazza, L., Chitale, D. A., Barod, R., Rogers, C. G., Kalyana-Sundaram, S., Palanisamy, N., Gupta, N. S., & Williamson, S. R. (2017). Renal cell tumors with clear cell histology and intact VHL and chromosome 3p: a histological review of tumors from the Cancer Genome Atlas database. *Modern Pathology*, 30(11), 1603–1612. https://doi.org/10.1038/modpathol.2017.72
- Gatto, F., Nookaew, I., & Nielsen, J. (2014). Chromosome 3p loss of heterozygosity is associated with a unique metabolic network in clear cell renal carcinoma. *Proceedings of the National Academy of Sciences*, 111(9). https://doi.org/10.1073/pnas.1319196111
- Hsieh, J. J., Le, V. H., Oyama, T., Ricketts, C. J., Ho, T. H., & Cheng, E. H. (2018). Chromosome 3p Loss–Orchestrated VHL, HIF, and Epigenetic Deregulation in Clear Cell Renal Cell Carcinoma. *Journal of Clinical Oncology*, 36(36), 3533–3539. https://doi.org/10.1200/JCO.2018.79.2549
- Kim, H. A. J., Shaikh, M. H., Lee, M., Zeng, P. Y. F., Sorgini, A., Akintola, T., Deng, X., Jarycki, L., Khan, H., MacNeil, D., Khan, M. I., Mendez, A., Yoo, J., Fung, K., Lang, P., Palma, D. A., Patel, K., Mymryk, J. S., Barrett, J. W., ... Nichols, A. C. (2021). 3p Arm Loss and Survival in Head and Neck Cancer: An Analysis of TCGA Dataset. Cancers, 13(21), 5313. https://doi.org/10.3390/cancers13215313
- Lee, D. J., Schönleben, F., Banuchi, V. E., Qiu, W., Close, L. G., Assaad, A. M., & Su, G. H. (2010). Multiple tumor-suppressor genes on chromosome 3p contribute to head and neck squamous cell carcinoma tumorigenesis. *Cancer Biology & Therapy*, 10(7), 689–693. https://doi.org/10.4161/cbt.10.7.12886
- Li, H., Yang, B., Xing, K., Yuan, N., Wang, B., Chen, Z., He, W., & Zhou, J. (2015). A preliminary study of the relationship between breast cancer metastasis and loss of heterozygosity by using exome sequencing. Scientific Reports, 4(1), 5460. https://doi.org/10.1038/srep05460
- Martinez, A. (2001). Chromosome 3p allele loss in early invasive breast cancer: detailed mapping and association with clinicopathological features. *Molecular Pathology*, 54(5), 300–306. https://doi.org/10.1136/mp.54.5.300

- Peng, J., Wang, Q., Liu, H., Ye, M., Wu, X., & Guo, L. (2016). EPHA3 regulates the multidrug resistance of small cell lung cancer via the PI3K/BMX/STAT3 signaling pathway. *Tumor Biology*, 37(9), 11959– 11971. https://doi.org/10.1007/s13277-016-5048-4
- Yang, Q., Yoshimura, G., Mori, I., Sakurai, T., & Kakudo, K. (2002). Chromosome 3p and breast cancer. *Journal of Human Genetics*, 47(9), 453–459. https://doi.org/10.1007/s100380200064
- Zabarovsky, E. R., Lerman, M. I., & Minna, J. D. (2002). Tumor suppressor genes on chromosome 3p involved in the pathogenesis of lung and other cancers. *Oncogene*, 21(45), 6915–6935. https://doi.org/10.1038/sj.onc.1205835
- Zhuang, G., Song, W., Amato, K., Hwang, Y., Lee, K., Boothby, M., Ye, F., Guo, Y., Shyr, Y., Lin, L., Carbone, D. P., Brantley-Sieders, D. M., & Chen, J. (2012). Effects of Cancer-Associated EPHA3 Mutations on Lung Cancer. *JNCI: Journal of the National Cancer Institute*, 104(15), 1183–1198. https://doi.org/10.1093/jnci/djs297

Disclaimer: This document is valid until the product expiry on the kit label