

A Sysmex Group Company

CytoCell

Instructions For Use

REF: LPH 006-S / LPH 006

13q14.3 Deletion Probe



PROFESSIONAL USE ONLY



Further information and other languages available at www.ogt.com

Limitations

This device is designed to detect genomic losses larger than the region covered by the red clone in this probe set, which includes the 13q14.3 region. Genomic losses outside this region or partial losses of this region may not be detected with this product.

The test is not intended for: use as a stand-alone diagnostic, prenatal testing, population-based screening, near-patient testing or self-testing. This product is intended for laboratory professional use only; all results should be interpreted by suitably-qualified staff, taking into account other relevant test results.

This product has not been validated for use on sample types or disease types other than those specified in the intended use.

Reporting and interpretation of FISH results should be consistent with professional standards of practice and should take into consideration other clinical and diagnostic information. This kit is intended as an adjunct to other diagnostic laboratory tests and therapeutic action should not be initiated on the basis of the FISH result alone.

Failure to adhere to the protocol may affect the performance and lead to false positive/negative results.

This kit has not been validated for purposes outside of the intended use stated.

Intended Use

The CytoCell 13q14.3 Deletion Probe is a qualitative, non-automated, fluorescence *in situ* hybridisation (FISH) test used to detect chromosomal deletions in the 13q14.2-q14.3 region on chromosome 13 in Carnoy's solution (3:1 methanol/acetic acid) fixed haematologically-derived cell suspensions from patients with confirmed or suspected chronic lymphocytic leukaemia (CLL) or multiple myeloma (MM).

Indications

This product is designed as an adjunct to other clinical and histopathological tests in recognised diagnostic and clinical care pathways, where knowledge of 13q14.3 deletion status would be important for clinical management.

Principles of the Test

Fluorescence *in situ* hybridisation (FISH) is a technique that allows DNA sequences to be detected on metaphase chromosomes or in interphase nuclei from fixed cytogenetic samples. The technique uses DNA probes that hybridise to entire chromosomes or single unique sequences, and serves as a powerful adjunct to G-banded cytogenetic analysis. This technique can now be applied as an essential investigative tool within prenatal, haematological and solid tumour chromosomal analysis. Target DNA, after fixation and denaturation, is available for annealing to a similarly denatured, fluorescently labelled DNA probe, which has a complementary sequence. Following hybridisation, unbound and non-specifically bound DNA probe is removed and the DNA is counterstained for visualisation. Fluorescence microscopy then allows the visualisation of the hybridised probe on the target material.

Probe Information

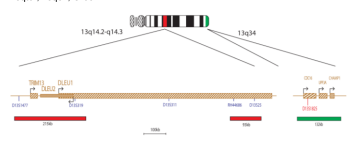
Rearrangements leading to the loss of all or part of the long arm of chromosome 13 are seen frequently in a wide range of haematological disorders.

Chromosome 13q aberrations occur in 16-40% of multiple myeloma cases (MM), most of them being complete monosomy 13 (85%), while the remaining 15% constitute deletion of 13q^{1,2,3}. A case study of multiple myeloma patients narrowed

down the critical deleted region to 13q14⁴. Historically, deletions of 13q have been associated with poor prognosis in MM but now it is believed that its prognostic relevance may be related to its association with other concurrent genetic lesions^{3.5}. Deletions affecting 13q14 are also the most frequent structural genetic aberrations in chronic lymphocytic leukaemia (CLL)^{6.7.8}. This region is found to be heterozygously deleted in 30-60% and homozygously deleted in 10-20% of CLL patients⁹. The survival rate has been shown to be similar for the two groups¹⁰. Patients with 13q14 deletions are classified as very low risk, in the absence of any other genetic lesions¹¹.

Two non-coding RNA genes, DLEU1 (*deleted in lymphocytic leukemia* 1) and DLEU2 (*deleted in lymphocytic leukemia* 2), plus the genetic marker D13S319, span the pathogenic critical region of 13q14¹². DLEU1 is considered to be the most likely CLL-associated candidate tumour suppressor gene within the 13q14 region¹³. Subsequently, D13S319, located between the RB1 gene and D13S25 and within the DLEU1 locus, was found to be deleted in 44% of CLL cases¹⁴. It has also been postulated that a gene telomeric to the D13S319 region, encompassing D13S25, may be important in cases with hemizygous deletions and that this gene is a putative tumour suppressor gene¹⁵.

Probe Specification 13q14.2-q14.3, Red 13qter, 13q34, Green



The 13q14.2-q14.3 probes, labelled in red, cover the D13S319 and D13S25 markers. The 13qter subtelomere specific probe (clone 163C9), labelled in green, allows identification of chromosome 13 and acts as a control probe.

Materials Provided

Probe: 50µl per vial (5 tests) or 100µl per vial (10 tests) The probes are provided premixed in hybridisation solution (formamide; dextran sulphate; saline-sodium citrate (SSC)) and are ready to use.

Counterstain: 150µl per vial (15 tests)

The counterstain is DAPI antifade (ES: 0.125µg/ml DAPI (4,6-diamidino-2-phenylindole)).

Warnings and Precautions

- 1. For *in vitro* diagnostic use. For professional use only.
- 2. Wear gloves when handling DNA probes and DAPI counterstain.
- 3. Probe mixtures contain formamide, which is a teratogen; do not breathe fumes or allow skin contact. Handle with care; wear gloves and a lab coat.
- DAPI is a potential carcinogen. Handle with care; wear gloves and a lab coat.
 Dispose of all hazardous materials according to your institution's guidelines for bazardous waste disposal
- Operators must be capable of distinguishing the colours red, blue and green.
- 7. Failure to adhere to the outlined protocol and reagents may affect the
- performance and lead to false positive/negative results.
- 8. The probe should not be diluted or mixed with other probes.
- Failure to use 10µl of probe during the pre-denaturation stage of the protocol may affect the performance and lead to false positive/negative results.

Storage and Handling



The kit should be stored between -25°C to -15°C in a freezer until the expiry date indicated on the kit label. The probe and counterstain vials must be stored in the dark.



The probe remains stable throughout the freeze-thaw cycles experienced during normal use (where one cycle constitutes the probe's removal from and replacement into the freezer) and is photostable for up to 48 hours after being exposed to continuous lighting conditions. All efforts must be made to limit exposure to

Equipment and Materials Necessary but not Supplied

- Calibrated equipment must be used:
- 1. Hotplate (with a solid plate and accurate temperature control up to 80°C)
- 2. Calibrated variable volume micropipettes and tips range $1\mu l$ $200\mu l$ 3. Water bath with accurate temperature control at $37^{o}C$ and $72^{o}C$

light and temperature changes.

- Microcentrifuge tubes (0.5ml)
- 5. Fluorescence microscope (Please see Fluorescence Microscope Recommendation section)
- 6. Phase contrast microscope
- 7. Clean plastic, ceramic or heat-resistant glass Coplin jars
- 8. Forceps
- 9. Calibrated pH meter (or pH indicator strips capable of measuring pH 6.5-8.0)
- 10. Humidified container
- 11. Fluorescence grade microscope lens immersion oil
- 12. Bench top centrifuge

13. Microscope slides

- 14. 24x24mm coverslips
- 15. Timer
- 37°C incubator
 Rubber solution a
- 17. Rubber solution glue
- 18. Vortex mixer
- 19. Graduated cylinders
- Magnetic stirrer
 Calibrated thermometer
- 21. Calibrated thermometer

Optional Equipment not Supplied

1. Cytogenetic drying chamber

Reagents Needed but not Supplied

- 1. 20x saline-sodium citrate (SSC) Solution
- 2. 100% Ethanol
- Tween-20
 1M Sodium hydroxi
- 1M Sodium hydroxide (NaOH)
 1M Hydrochloric acid (HCl)
- 6. Purified water

Fluorescence Microscope Recommendation

Use a 100-watt mercury lamp or equivalent and oil immersion plan apochromat objectives 60/63x or 100x for optimal visualisation. The fluorophores used in this probe set will excite and emit at the following wavelengths:

Fluorophore	Excitation _{max} [nm]	Emission _{max} [nm]
Green	495	521
Red	596	615

Ensure appropriate excitation and emission filters that cover the wavelengths listed above are fitted to the microscope. Use a triple bandpass DAPI/green spectrum/red spectrum filter or a dual bandpass green spectrum/red spectrum filter for optimal simultaneous visualisation of the green and red fluorophores.

Check the fluorescence microscope before use to ensure it is operating correctly. Use immersion oil that is suitable for fluorescence microscopy and formulated for low auto fluorescence. Avoid mixing DAPI antifade with microscope immersion oil as this will obscure signals. Follow manufacturers' recommendations in regards to the life of the lamp and the age of the filters.

Sample Preparation

The kit is designed for use on haematologically-derived cell suspensions fixed in Carnoy's solution (3:1 methanol/acetic acid) fixative, that are prepared according to the laboratory or institution guidelines. Prepare air dried samples on microscope slides according to standard cytogenetic procedures. The AGT *Cytogenetics Laboratory Manual* contains recommendations for specimen collection, culturing, harvesting and for slide making¹⁶.

Solution Preparation

Ethanol Solutions

Dilute 100% ethanol with purified water using the following ratios and mix thoroughly.

- 70% Ethanol 7 parts 100% ethanol to 3 parts purified water
- 85% Ethanol 8.5 parts 100% ethanol to 1.5 parts purified water

Store the solutions for up to 6 months at room temperature in an airtight container.

2xSSC Solution

Dilute 1 part 20xSSC Solution with 9 parts purified water and mix thoroughly. Check pH and adjust to pH 7.0 using NaOH or HCl as required. Store the solution for up to 4 weeks at room temperature in an airtight container.

0.4xSSC Solution

Dilute 1 part 20xSSC Solution with 49 parts purified water and mix thoroughly. Check pH and adjust to pH 7.0 using NaOH or HCl as required. Store the solution for up to 4 weeks at room temperature in an airtight container.

2xSSC, 0.05% Tween-20 Solution

Dilute 1 part 20xSSC Solution with 9 parts purified water. Add 5µl of Tween-20 per 10ml and mix thoroughly. Check pH and adjust to pH 7.0 using NaOH or HCl as required. Store the solution for up to 4 weeks at room temperature in an airtight container.

FISH Protocol

(Note: Ensure that exposure of the probe and counterstain to laboratory lights is limited at all times).

Slide Preparation

- Spot the cell sample onto a glass microscope slide. Allow to dry. (Optional, if using a cytogenetic drying chamber: slides should be spotted using a cytogenetic drying chamber. The chamber should be operated at approximately 25°C and 50% humidity for optimal cell sample spotting. If a cytogenetic drying chamber is not available, use a fume hood as an alternative).
- 2. Immerse the slide in 2xSSC for 2 minutes at room temperature (RT) without agitation.
- Dehydrate in an ethanol series (70%, 85% and 100%), each for 2 minutes at RT.
- Allow to dry.

Pre-Denaturation

- Remove the probe from the freezer and allow it to warm to RT. Briefly centrifuge tubes before use.
- 6. Ensure that the probe solution is uniformly mixed with a pipette.
- Remove 10µl of probe per test, and transfer it to a microcentrifuge tube. Quickly return the remaining probe to the freezer.
- Place the probe and the sample slide to prewarm on a 37°C (+/- 1°C) hotplate for 5 minutes.
- Spot 10µl of probe mixture onto the cell sample and carefully apply a coverslip. Seal with rubber solution glue and allow the glue to dry completely.

Denaturation

10. Denature the sample and probe simultaneously by heating the slide on a hotplate at 75°C (+/- 1°C) for 2 minutes.

Hybridisation

11. Place the slide in a humid, lightproof container at 37°C (+/- 1°C) overnight.

Post-Hybridisation Washes

- 12. Remove the DAPI from the freezer and allow it to warm to RT.
- 13. Remove the coverslip and all traces of glue carefully.
- Immerse the slide in 0.4xSSC (pH 7.0) at 72°C (+/- 1°C) for 2 minutes without agitation.
- Drain the slide and immerse it in 2xSSC, 0.05% Tween-20 at RT (pH 7.0) for 30 seconds without agitation.
- 16. Drain the slide and apply 10µl of DAPI antifade onto each sample.
- 17. Cover with a coverslip, remove any bubbles and allow the colour to develop in the dark for 10 minutes.
- 18. View with a fluorescence microscope (see Fluorescence Microscope Recommendation).

Stability of Finished Slides

Finished slides remain analysable for up to 1 month if stored in the dark at/or below RT.

Procedural Recommendations

- 1. Baking or ageing of slides may reduce signal fluorescence
- Hybridisation conditions may be adversely affected by the use of reagents other than those provided or recommended by Cytocell Ltd
- Use a calibrated thermometer for measuring temperatures of solutions, waterbaths and incubators as these temperatures are critical for optimum product performance
- The wash concentrations, pH and temperatures are important as low stringency can result in non-specific binding of the probe and too high stringency can result in a lack of signal
- 5. Incomplete denaturation can result in lack of signal and over denaturation can also result in non-specific binding
- 6. Over hybridisation can result in additional or unexpected signals
- Users should optimise the protocol for their own samples prior to using the test for diagnostic purposes
- 8. Suboptimal conditions may result in non-specific binding that may be misinterpreted as a probe signal

Interpretation of Results

Assessing Slide Quality

The slide should not be analysed if:

- Signals are too weak to analyse in single filters in order to proceed with analysis, signals should appear bright, distinct and easily evaluable
- There are high numbers of clumped/overlapping cells obstructing the analysis
 >50% of the cells are not hybridised
- There is excess of fluorescent particles between cells and/or a fluorescent
- There is excess of nuclescent particles between cens and/or a nuclescent haze that interferes with the signals - in optimal slides the background should appear dark or black and clean
- Cell nucleus borders cannot be distinguished and are not intact

Analysis Guidelines

- Two analysts should analyse and interpret each sample. Any discrepancies should be resolved by assessment by a third analyst
- Each analyst should be suitably qualified according to recognised national standards
- Each analyst should score independently 100 nuclei for each sample. The first
 analyst should start the analysis from the left side of the slide and the second
 analyst from the right one
- · Each analyst should document their results in separate sheets
- Analyse only intact nuclei, not overlapped or crowded nuclei or nuclei covered by cytoplasmic debris or high degree of autofluorescence
- Avoid areas where there is excess of cytoplasmic debris or non-specific hybridisation
- Signal intensity may vary, even with a single nucleus. In such cases, use single filters and/or adjust the focal plane
- In suboptimal conditions signals may appear diffuse. If two signals of the same colour touch each other, or the distance between them is no greater than two signal widths, or when there is a faint strand connecting the two signals, count as one signal
- If in doubt about whether a cell is analysable or not, then do not analyse it

Analysis Guidelines		
	Do not count – nuclei are too close together to determine boundaries	
	Do not count overlapping nuclei – all areas of both nuclei are not visible	
••••	Count as two red signals and two green signals – one of the two red signals is diffuse	
• • •	Count as two red signals and two green signals – the gap in one red signal is less than two signal widths	

Expected Results Expected Normal Signal Pattern



In a normal cell, two red and two green signals (2R, 2G) are expected.

Expected Abnormal Signal Patterns



In a cell with a hemizygous deletion of the 13q14.3, the expected signal pattern will be one red and two green signals (1R, 2G).



In a cell with a homozygous deletion, the expected signal pattern will be no red and two green signals (0R, 2G).

13q deletions in CLL are recognised as being heterogenous; small deletion within the 13q region may result in a small residual signal with this probeset.

Other signal patterns are possible in aneuploid/unbalanced specimens.

Known Cross-Reactivity

The green 13qter probe may show cross-hybridisation to the centromere of chromosome 19 and the p-arms of other chromosomes.

Adverse Event Reporting

If you believe this device has malfunctioned or suffered a deterioration in its performance characteristics which may have contributed to an adverse event (e.g. delayed or misdiagnosis, delayed or inappropriate treatment), this must be reported immediately to the manufacturer (**email:** vigilance@ogt.com).

If applicable, the event should also be reported to your national competent authority. A list of vigilance contact points can be found at: http://ec.europa.eu/growth/sectors/medical-devices/contacts/.

Specific Performance Characteristics Analytical Specificity

Analytical specificity is the percentage of signals that hybridise to the correct locus and no other location. The analytical specificity was established by analysing a total of 200 target loci. The analytical specificity was calculated as the number of FISH signals that hybridised to the correct locus divided by the total number of FISH signals hybridised.

Table	 Analytical 	Specificity	for the	13q14.3 Deletion Probe

Probe	Target Locus	No. of Signals Hybridised to the Correct Locus	Total No. of Signals Hybridised	Specificity (%)
Red 13q14.3	13q14.3	200	200	100
Green 13qter	13qter, 13q34	200	200	100

Analytical Sensitivity

Analytical sensitivity is the percentage of scoreable interphase cells with the expected normal signal pattern. The analytical sensitivity was established by analysing interphase cells across different normal samples. The sensitivity was calculated as the percentage of scoreable cells with the expected signal pattern (with a 95% confidence interval).

Table 2. Analytical Sensitivity for the 13q14.3 Deletion Probe

No. of Cells with Expected Signal Patterns	No. of Cells with Scoreable Signals	Sensitivity (%)	95% Confidence Interval
481	500	96.2	1.6

Characterisation of Normal Cut-off Values

The normal cut-off value, in association with FISH probes, is the maximum percentage of scoreable interphase cells with a specific abnormal signal pattern at which a sample is considered normal for that signal pattern.

The normal cut-off value was established using samples from normal and positive patients. For each sample, the signal patterns of 100 cells were recorded. The Youden index was calculated to find the threshold value for which Sensitivity + Specificity-1 is maximised.

Table 3. Characterisation of Normal Cut-off Values for the 13q14.3 Deletion Probe

Abnormal signal pattern	Youden Index	Normal Cut-off (%)
1R, 2G or 0R, 2G	0.95	7

Laboratories must verify cut-off values using their own data^{17, 18}.

Precision and Reproducibility

Precision is a measure of the natural variation of a test when repeated several times under the same conditions. This was assessed by analysing repeats of the same lot number of probe tested on the same sample, in the same conditions on the same day.

Reproducibility is a measure of the variability of a test and has been established in terms of sample-to-sample, day-to-day and batch-to-batch variability. Day-to-day reproducibility was assessed by analysing the same samples on three different days. Batch-to-batch reproducibility was assessed by analysing the same samples using three different to numbers of probe on one day. Sample-to-sample reproducibility was assessed by analysing three replicates of a sample on one day. For each sample, signal patterns of 100 interphase cells were recorded and the percentage of cells with the expected signal pattern was calculated.

The reproducibility and precision were calculated as the Standard Deviation (STDEV) between replicates for each variable and overall mean STDEV.

Table 4. Reproducibility and Precision for the 13q14.3 Deletion Probe

Variable	Standard Deviation (STDEV)
Precision	0.72
Sample-to-sample	0.58
Day-to-day	0.96
Batch-to-batch	1.40
Overall deviation	1.03

Clinical Performance

The clinical performance was established on a representative sample of the intended population for the product. For each sample, the signal patterns of \geq 100 interphase cells were recorded. A normal/abnormal determination was made by comparing the percentage of cells with the specific abnormal signal pattern to the normal cut-off value. The results were then compared to the known status of the sample.

The results of the clinical data were analysed in order to produce sensitivity, specificity and cut off values using a one-dimensional approach.

Table 5. Clinical Performance for the 13q14.3 Deletion Probe

Variable	Result
Clinical Sensitivity (true positive rate, TPR)	96.3%
Clinical Specificity (true negative rate, TNR)	99.1%
False Positive rate (FPR) = 1 – Specificity	0.9%

Additional Information

For additional product information please contact the CytoCell Technical Support Department.

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Guid	Guide to Symbols				
	REF	en: Catalogue number			
	IVD	en: In vitro diagnostic medical device			
	LOT	en: Batch code			
	i	en: Consult instructions for use			
		en: Manufacturer			
	\Box	en: Use-by date			

 _25°C
 en: Temperature limit

 Image: Constant sufficient for <n> tests

 Image: Constant sufficient for <n> tests

 Image: Constant sufficient for <n> tests

Patents and Trademarks

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