Shorter Hybridisation Times Using CytoCell Probes



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What binds us, makes us.

S Chatters, F Partheniou, A Hobbs, G Fonseka, A Castro-Justo, A Gocza-Blasko, K Mak-Hannah, R Frodsham, M Lawrie Cytocell Ltd., Oxford Gene Technology (OGT).

Introduction

Fluorescence *in situ* hybridisation (FISH) analysis is the 'gold-standard' method for the detection of balanced and unbalanced chromosomal rearrangements plus gains and deletions in neoplastic specimens. Standard FISH protocols incorporate an overnight hybridisation step; however, shorter hybridisations are sometimes desired as a result of laboratory operational requirements. The CytoCell[®] FISH probe range from Oxford Gene Technology (OGT) delivers bright, clear and precise signals when hybridised overnight. The aim of this project was to determine whether the hybridisation times for standard CytoCell 'off-the-shelf' CE-marked IVD probes could be reduced without detrimental effect on probe performance.

Materials and Methods

Ten CytoCell Haematology CE-marked IVD probe kits were selected for analysis, as shown in Table 1, below.

Probe Name	Cat. No.
BCR/ABL Translocation, Dual Fusion	LPH 007
MLL Breakapart	LPH 013
IGH Breakapart	LPH 014
P53 Deletion	LPH 017
Del(20q) Deletion	LPH 020
Del(5q) Deletion	LPH 024
Del(7q) Deletion	LPH 025
BCR/ABL Plus Translocation, Dual Fusion	LPH 038
CKS1B/CDKN2C (P18) Amplification/Deletion	LPH 039
P53/ATM Probe Combination	LPH 052

Table 1: Probe selection.

Following the standard CytoCell protocol, (Table 2), four replicates of each probe set were hybridised to Carnoy's-fixed peripheral blood lymphocyte and bone marrow samples and each replicate hybridised for five differing times: one hour, two hours, three hours, four hours and overnight (sixteen hours), making a total of 160 separate hybridisations.

Step 1	Spot slide with 10µl of cell sample and dehydrate.	Step 5	Hybridise the slide in a humid, lightproof container at 37°C (+/-1°C).
Step 2	Apply 10µl of probe onto dehydrated cell sample.	Step 6	Wash the slide in 0.4xSSC (pH 7.0) at 72°C (+/- 1°C) for 2 minutes.
Step 3	Place coverslip onto slide and seal.	Step 7	Wash the slide in 2xSSC+0.05% Tween-20 (pH 7.0) at RT for 30 seconds.
Step 4	Denature on a hotplate at 75°C (+/-1°C) for 2 minutes.	Step 8	Apply DAPI counterstain provided and view under a fluorescent microscope.

Table 2: Outline of Standard CytoCell FISH Procedure.

Each of the hybridisation replicates were analysed and assessed for individual probe component intensity and sensitivity by two independent analysts, scoring a target number of one hundred interphase nuclei each, according to CytoCell standard QC procedure, and validated against individual probe intensity measured using MetaSystems® Isis and Metafer image analysis software.

Results and Discussion

The independent scoring and intensity data were normalised against the data obtained for standard overnight hybridisation. This enabled relative sensitivity and intensity relative values to be calculated.

Absolute intensity, by pixel, and eye intensity values were combined to give a signal intensity index for each probe set. These indices were analysed by fluorophore colour and sample type, using a standard overnight hybridisation as baseline to give a measure of any reduction in signal intensity. Sensitivity data were normalised in the same manner, giving an indication of the probe sensitivity, alongside standard deviation values. These results are shown in Tables 3 and 4, and Figures 2 and 3.

At a hybridisation of four hours, the intensity and sensitivity indices for all sets of data are above 0.8 – the cut-off for CytoCell QC analysis. This indicates the presence of bright easily-scored signals with little dropout for all fluorophore types in all cell types analysed. Furthermore, many of the indices for hybridisation times below four hours are above or approaching the 0.8 value, which still indicates presence of strong bright signals, confirming the robustness of CytoCell probes when used with shorter hybridisation times. It is of note that even with a reduction of signal intensity below that of the CytoCell QC cut-off, signals remained bright and scoreable.

Hyb Time	All Probe Sets	Green Signals Only	Red Signals Only	Aqua Signals Only	PB Lymphs Only	Bone Marrow Only
ON	1.00	1.00	1.00	1.00	1.00	1.00
4	0.88	0.92	0.85	0.80	0.85	0.90
3	0.76	0.79	0.74	0.80	0.71	0.81
2	0.61	0.61	0.61	0.70	0.56	0.66
1	0.52	0.51	0.50	0.70	0.49	0.55

Table 3: Mean Intensity Index.

Hyb Time	All Pro	obe Sets	PB Lymphocytes Only		Bone Marrow Only	
	Mean	SD	Mean	SD	Mean	SD
ON	1.00	-	1.00	-	1.00	-
4	0.92	0.15	0.94	0.07	0.91	0.19
3	0.83	0.12	0.83	0.10	0.83	0.14
2	0.82	0.12	0.84	0.10	0.79	0.14
1	0.76	0.19	0.71	0.20	0.80	0.18

Table 4: Sensitivity Index

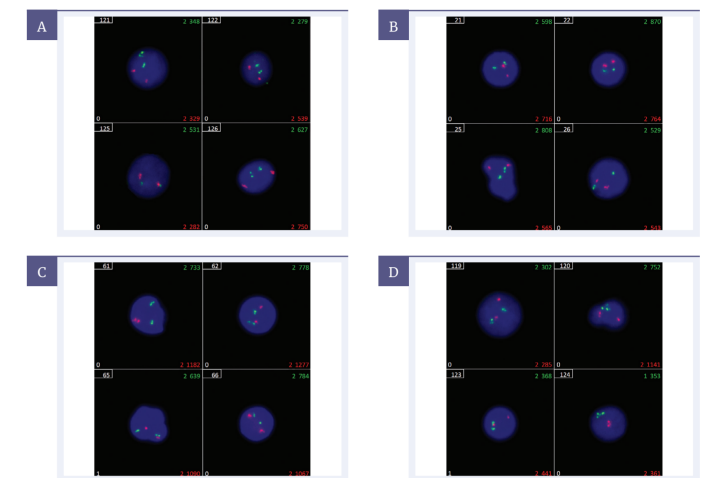


Figure 1: Comparison of hybridisation results. LPH007 probe set at A overnight, B four hours, C three hours, D one hour. Output from Metafer at 60x.



Figure 2: Mean Intensity Index.



Figure 3: Mean Sensitivity Index.

Conclusions

These data show that, for a range of probe and specimen types, the hybridisation time for standard CE IVD-marked CytoCell probes may be shortened from overnight to four hours or fewer, whilst retaining excellent signal sensitivity and intensity. This gives laboratories the flexibility to validate and use off-the-shelf CytoCell probes with reduced hybridisation times, facilitating same-day reporting, should this be required.

