A simple and reliable pretreatment protocol facilitates fluorescent in situ hybridisation on tissue microarrays of paraffin wax embedded tumour samples

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MATERIALS AND METHODS

Breast tissue microarray

The breast tumour tissue microarray was constructed as described previously.5 Sections (5 µm thick) were prepared from a tissue microarray block containing 100 cores and placed on to either positively charged slides (BDH, Poole, Dorset, UK) or silanised slides (BDH; Dako, Glostrup, Denmark).

Multi-tissue microarray

A multi-tissue type array was purchased from Ambion (Landmark Tissue MicroArray Test Array, Lot number 022P21A; Ambion, Huntingdon, Cambridgeshire, UK). The Ambion TMA contained approximately 50 specimens/slide of normal or diseased tissues of the following types: liver, thyroid, lung, prostate, muscle, breast cancer, lymph node, lung cancer, testis, prostate cancer, kidney, and colon cancer.

Fluorescence in situ hybridisation

Probes

In total, 30 probes were tested in hybridisations to TMAs, of which 23 were in house labelled and seven were commercial probes. In house probes (DNA isolated from BAC clones) encompassing genes and loci of interest were labelled with either digoxigenin–11-dUTP, estradiol–15-dUTP, biotin–16-dUTP (Roche Biochemicals, Lewes, East Sussex, UK), or Spectrum Orange–dUTP by nick translation (Vysis, Abott Diagnostics, Maidenhead, Berkshire, UK). These in house probes were robust and reproducible for FISH using paraffin wax embedded cancer tissues in general and in particular TMAs.

Abbreviations: FISH, fluorescent in situ hybridisation; SSC, saline sodium citrate; SST, saline sodium citrate containing 0.05% Tween 20; TMA, tissue microarray
included 14 BACs on chromosome 8p12 (obtained from Research Genetics, InVitrogen, Paisley, UK and the Sanger Centre, Hinxton, Cambridgeshire, UK), four BACs on chromosome 11q13 (obtained from BACPAC Resource Center, Oakland Children’s Hospital, California, USA), three BACs on 12q24 (obtained from the Sanger Centre), BAC containing GARP (kind gift from Vysis, USA), and a plasmid clone containing centromere chromosome 7 sequences (provided by Dr M Rocchi, Resources for Molecular Cytogenetics, University of Bari, Italy). The labelled probes were precipitated in the presence of human COT1 DNA (Roche Biochemicals) and resuspended in hybridisation buffer (Hybrisol VI; Qbiogene, Nottingham, UK). Commercial directly labelled probes purchased from Vysis were: LAVysion with EFGR–Spectrum Red, C-MYC–Spectrum Gold, loci 5p15.2–Spectrum Green, centromere of chromosome 6–Spectrum Aqua, cyclin D1–Spectrum Orange, centromere of chromosome 8–Spectrum Orange, and centromere of chromosome 11–Spectrum Orange.

**Pretreatment protocols**

Figure 1 is a flow chart comparing the steps involved in the pretreatment of paraffin wax embedded tissue sections either with the 1M sodium thiocyanate method or the 10mM citric acid buffer.

**Sodium thiocyanate method**
- Xylene/RT/10mins/3x
- 100% EtOH/RT/10mins/2x
- Air dry
- 0.2N HCl/37°C/30mins
- Wash dH2O/RT/5mins
- Wash 2× SSC/RT/5 mins
- 1M NaSCN/80°C/30 mins
- Wash 2× SSC/RT/5mins/2x
- Wash in dH2O
- 1–20 mg/ml pepsin in 0.2N HCl pH 2.0 for 10 mins/37°C
- Wash 2× SSC/RT/5mins/2x
- Ethanol dehydration
- Co-denature and hybridise

**Citric acid buffer method**
- Xylene/RT/10mins/3x
- 100% EtOH/RT/10mins/2x
- Air dry
- 10 mM citric acid buffer pH 6.0/80°C/30 mins to 2 hours
- Wash 2× SSC/RT/5mins/2x
- Wash in dH2O
- 1–20 mg/ml pepsin in 0.2N HCl pH 2.0 for 10 mins/37°C
- Wash 2× SSC/RT/5mins/2x
- Ethanol dehydration
- Co-denature and hybridise

**In situ hybridisation**

Aliquots containing 100 ng of home labelled probes or 5 µl of the LAVysion probes (Vysis) were co-denatured with the tissues at 80°C for 10 minutes and hybridised at 37°C for 48–72 hours. After hybridisation, slides were washed in 0.4× saline sodium citrate (SSC)/0.3% Nonidet P-40 at 73°C for three minutes, followed by 2× SSC at room temperature. Slides hybridised with directly labelled probes were dehydrated in an ethanol series, air dried and counterstained with 4’6-diamidino-2-phenylindole in antifade solution (Vectashield; Vector Laboratories, Peterborough, UK).

For probes labelled indirectly with digoxigenin, estradiol, or biotin, the slides were blocked with 3% bovine serum albumin made up in 2× SSC containing 0.05% Tween 20 (2× SST) for 20 minutes and washed three times with 2× SST at room temperature. This was followed by incubation at 37°C for 30 minutes with either antidigoxigenin–fluorescein isothiocyanate, anti-estradiol–rhodamine (Roche Biochemicals), or avidin–CY5 (Amersham, Little Chalfont, Buckinghamshire, UK). The slides were washed three times in 2× SST at room temperature. The slides were mounted as described above.

**Image analysis**

Slides were examined under a Zeiss Axioplan 2 epifluorescence microscope (Zeiss, Welwyn Garden City, UK) equipped with single band pass emission filters. Digital images were taken using a Hamamatsu ORCA II camera (Hamamatsu, Welwyn Garden City, UK) operated via the SmartCapture software (Digital Scientific, Cambridge, UK).

**RESULTS**

Comparing the effect of citric acid buffer and sodium thiocyanate pretreatment protocol on tissue attachment

First we evaluated the effect of the different pretreatments on tissue morphology and attachment to the slides. Pretreating paraffin wax embedded sections with 1M sodium thiocyanate
at 80°C for 30 minutes was most detrimental to the condition of the sections, especially for TMAs, where the cores were “floating” loosely from the slide. The loss of tissue from the slide was more apparent after protein digestion, independent of the concentration and type of enzyme used (pepsin, proteinase K, or protease). When the TMAs were pretreated with NaSCN, approximately 44.5% of the cores were lost (fig 2) when compared with citric acid buffer treatment. The loss of cores after sodium thiocyanate pretreatment was consistent in several TMA sections tested. The attachment of the tissues to the slide was not improved by the use of aminopropyltriethoxysilane coated slides or Dako silanised slides (46% loss).

Reducing the length of incubation to 10 minutes or reducing the temperature to 45°C improved the attachment of the tissue sections, but the quality of the FISH signals was compromised (data not shown). Less than 1% of the cores were lost using our pretreatment method (fig 2) on more than 30 TMA sections used.

We found that the alternative heating methods with citric acid buffer used daily in immunohistochemistry laboratories were also gentler on the tissue sections and less cores were lost when compared with the sodium thiocyanate protocol. Only 5% of tissue cores were lost when TMAs were heated in a pressure cooker, whereas a loss of only 2% was seen after microwaving (data not shown).

Comparing the effect of citric acid buffer and sodium thiocyanate pretreatment on FISH signals

We assessed the quality of the FISH signals by comparing the relative signal intensities of each fluorophore as demonstrated by the exposure time and the amount of autofluorescence background. Signal intensity was comparable using both treatment protocols, citric acid buffer and sodium thiocyanate. The main difference was in the amount of autofluorescence background, which was much higher using the sodium thiocyanate treatment (data not shown). This made the scoring of signals by eye difficult and automated digital capture of the images impossible. No difference in signal quality and autofluorescence background was found between the three citric acid buffer based methods, which were significantly better than the sodium thiocyanate method, and allowed for scoring by eye.

The citric acid buffer protocol worked robustly with both centromeric probes and locus/gene specific probes (fig 3). The quality of the FISH signals generated from in house labelled probes was comparable to that obtained using commercial probes, for example GARP versus cyclin D1 (fig 3F). In addition to breast cancer tissues, we found that our pretreatment protocol worked without further optimisation for other types of tissues, such as colon, lung, lymph node, prostate, muscle, kidney, and Ewing's sarcoma, suggesting that the protocol is robust (fig 3A–D). Neither the citric acid buffer nor the sodium thiocyanate protocol worked well with liver, spleen, or thyroid tissues because of the high autofluorescence emitted by the tissue and by blood cells.

**Optimising incubation time and pepsin digestion using the citric acid buffer protocol**

Incubating the slides for a minimum of 30 minutes was adequate for probes with highly repetitive sequences, such as centromeric probes, but when co-hybridising these probes with single copy probes, only the centromeric signals were consistently present in every cell. Incubating for 60 minutes resulted in strong signals for both types of probe. Longer incubation times resulted in no improvement in signal intensity or autofluorescence background (data not shown).
Citric acid buffer pretreated slides required no pepsin digestion if the probes were made of highly repetitive sequences, but a minimum of 1 mg/ml pepsin digestion for 15 minutes at 37°C was required for single copy probes. Increasing the concentration of pepsin to 5 mg/ml did reduce the autofluorescence background, but no difference in signal intensity was obtained. Using pepsin at concentrations greater than 10 mg/ml (data not shown) increased tissue digestion, significantly altered tissue morphology, and resulted in increased autofluorescence. In addition, we compared the efficacy of different types of proteolytic enzymes (pepsin, protease, and proteinase K) and found that proteinase K was the least suitable because it left highly fluorescent speckles within the cells (data not shown).

DISCUSSION

FISH, a technique that allows the visualisation of specific nucleic acid sequences in situ,\textsuperscript{12–14} has become a valuable technique for the analysis of chromosome/locus/gene copy number changes in paraffin wax embedded cancer samples, especially with the recent development of tissue microarrays.\textsuperscript{8–10}

In general, most paraffin wax embedded tissues are fixed with formalin, which results in the formation of methylene bridges between amino groups in the DNA and/or proteins; this helps to preserve the cellular composition and morphology of the tissue, but reduces the penetration of both antibodies and/or nucleic acid probes. Therefore, an important factor that determines the success of FISH on formalin fixed tissue is adequate unmasking of the target nucleic acids before hybridisation. These unmasking procedures or pretreatments used in FISH have been adapted from antigen retrieval techniques used in immunohistochemistry, which generally involve the application of high heat,\textsuperscript{15} chaotropic agents such as sodium thiocyanate,\textsuperscript{16} and protein digestion.\textsuperscript{17} The use of an acid, usually 0.2N HCl, has also been recommended because...
simple and reproducible paraffin FISH protocol

We have developed and validated an efficient pretreatment protocol for fluorescent in situ hybridization (FISH) that is easy, reproducible, and facilitates FISH on tissue microarrays (TMAs) of paraffin wax embedded tumour samples.

- Pretreatment with citrate acid buffer produces FISH signals with superior signal to noise ratios compared with sodium thiocyanate pretreatment.
- The best tissue attachment was achieved using our newly developed pretreatment protocol: less than 1% of TMA cores were lost.

Citrated acid buffer may act as deproteinising agents and increase probe penetration.19

We have found that sodium thiocyanate pretreatment is too harsh when applied to TMAs, because tissue sections were found to detach from the slides and were susceptible to mechanical disaggregation, caused by changing of the microscope focal plane. Loss of tissue cores presents further problems in sample identification and analysis on a TMA, where the identity of each section is based on the positional grid. The optimal concentration, incubation time, and temperature of these agents must be titrated for different tissue sections and tissue types. Therefore, finding the optimal conditions for all cores on a TMA is often difficult.

Leers et al reported recently that heating dewaxed tissue sections in 10mM citric acid buffer at 80°C for two hours followed by a short proteolytic step greatly improved the resolution of DNA flow cytometry profiles of paraffin wax embedded tumour tissue. The authors suggested that heating cells in an acidic environment led to the extraction of chromatin proteins and consequently allowed better binding of propidium iodide to DNA, thereby increasing fluorescence intensity. We decided to adapt this pretreatment protocol for FISH and the results presented here show that our revised protocol produced FISH signals with a significantly higher signal to noise ratio than the commonly used sodium thiocyanate pretreatment. Leers et al used tissue sections of 50 µm thickness, whereas TMA sections are only 4–5 µm thick. We found that the length of incubation could be reduced to one hour with no compromise in FISH signal intensity. FISH using the citric acid buffer protocol was successful for 22 locus specific probes, four centromeric probes, and four commercial locus specific probes in more than 90% of the cores on a TMA containing 100 cores of breast tissues (normal and tumour tissues). The cores where FISH was unsuccessful contained over 90% stromal material.

"Sodium thiocyanate pretreatment is too harsh when applied to tissue microarrays, because tissue sections were found to detach from the slides and were susceptible to mechanical disaggregation."19

“NATIVE DNA FLOW CYTOMETRY OF PARAFFIN-EMBEDDED TUMOR TISSUE. A SIMPLE AND REPRODUCIBLE PROTOCOL FOR FISH ON TMAS OF PARAFFIN WAX EMBOSS SAMPLES.”


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