Model system for optimising mRNA non-isotopic in situ hybridisation: Riboprobe detection of lysozyme mRNA in archival gut biopsy specimens

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Abstract

The aim of this study was to optimise conditions for mRNA detection by non-isotopic in situ hybridisation (NISH) using biotinylated and digoxigenin labelled riboprobes. Because lysozyme gene transcripts are present at high concentrations in Paneth and other alimentary cells, archival gut biopsy specimens were chosen as a model system for these experiments. Most of the variables in NISH, from unmasking of mRNA, to its ultimate detection by peroxidase or alkaline phosphatase-based detection systems, were examined in detail. The most important findings were that simultaneous heating of tissue targets and riboprobes at 95°C for 15 minutes before hybridisation at 50°C for two hours gave the most intense signal for lysozyme mRNA in Paneth cells, Brunner’s glands, and lamina propria macrophages; digoxigenin labelled riboprobes gave a higher signal to noise ratio than their biotinylated counterparts, and probes 600 base pairs long were superior to shorter probes. It is concluded that the mRNA NISH method may be generally useful for detecting gene transcription in archival clinical biopsy specimens.

Non-isotopic in situ hybridisation (NISH) has been developed for the precise localisation of nucleic acids in intact cells. In diagnostic molecular pathology most of the effort has been directed towards DNA identification by NISH in archival tissue samples. For full pathobiological analysis of disease, equally sensitive methodology for mRNA detection in archival biopsy needs to be developed. The most commonly used probes for mRNA detection are radiolabelled riboprobes, the signal being detected by autoradiography. Single radioactive oligonucleotides have also been used but these have theoretical disadvantages which may be circumvented for some gene transcripts by non-isotopically labelled oligonucleotide cocktails. The development of reproducible NISH protocols for riboprobe mRNA detection would have several advantages: non-isotopic probes have a longer shelf-life; biohazard and disposal are non-problematic; time-consuming autoradiography is avoided; and labelled riboprobes are easily produced. NISH also provides more precise cellular resolution than autoradiography.

Biotinylated probes have been used extensively to label DNA probes but endogenous biotin produces background noise in some organs. Since the introduction of digoxigenin labelled probes for NISH, the general experience has been that background is less of a problem with this reporter: it is a plant alkaloid and, unlike biotin (vitamin H), it is not a physiological tissue constituent and is uncharged at neutral pH. The improved methodology for digoxigenin detection makes this the reporter of choice in most NISH DNA experiments.

Lysozyme mRNA in archival clinical gut biopsy specimens was chosen as a model system to optimise NISH procedures for identification of gene transcription in archival clinical biopsy specimens. This protein and mRNA are present in appreciable quantities in well defined cell populations, such as Paneth cells and lamina propria mononuclear cells. We analysed the following NISH variables for mRNA localisation by digoxigenin and biotin labelled riboprobes: (i) unmasking conditions for mRNA; (ii) prehybridisation requirements; (iii) hybridisation conditions; (iv) optimisation of detection systems; and (v) the efficacy of RNase inhibitors.

Methods

All reagents were obtained from Sigma (UK) or BDH (UK) unless otherwise stated. Sixteen routine formalin fixed, paraffin wax embedded endoscopic biopsy specimens of histopathologically normal small or large bowel were obtained from our diagnostic archival file: these included two rectal biopsy specimens of Crohn’s colitis. Four small bowel surgical resection specimens were also studied after various fixation times. Sections (4 μm thick) were mounted on multi-well slides with four wells (12 mm in diameter) per slide (Henley, Essex) that had been previously coated with aminopropyl-triethoxyxilane, and baked at 75°C for 60 minutes and overnight at 60°C. Slides were heated at 75°C (10 minutes) and immediately transferred while still hot, to xylene in which they were gently washed with shaking at 22°C twice for 10 minutes each wash to remove the wax. Slides were immersed in...
methanol for 10 minutes, twice, and quickly rinsed (three times) in double distilled water (ddH₂O). Sections were warmed in ddH₂O at 37°C while the proteases were prepared (see below).

**RIBOPROBES**

The lysozyme probe was a 642 base pair human lysozyme cDNA, subcloned in both orientations into the Hinc II site of pGEM-3 (Promega Biotech, Madison, Wisconsin, USA). The recombinant plasmids were purified, linearised with Hind III, and transcribed with T7 RNA polymerase to generate anti-sense RNA probes. For sense RNA probes, the plasmids were linearised with EcoRI and transcribed with SP6 polymerase. An unrelated control probe, the 800 base pair fragment of the 3' end of a p53 cDNA, was subcloned into the Smal/BamHI sites of pBluescribe (Stickland J, Evans M, McGee J O'D, unpublished observations). EcoRI and Hind III linearised plasmids were used to synthesise both sense and anti-sense riboprobes with T3 and T7 RNA polymerases, respectively.

Riboprobes were labelled with digoxigenin-11-UTP (Boehringer, Germany) or biotin-11-UTP at concentrations of 350 μM/l and 1-375 mM/l, respectively, in the transcription reaction using a protocol based on that of Höltké and Kessler. The labelled probes were digested with DNase and extracted with phenol/chloroform and chloroform. Probe size and qualitative assessment of label incorporation was by gel electrophoresis of glyoxal denatured probe, followed by transfer to nitrocellulose filters, and biotin probes were detected as described: for digoxigenin detection, anti-digoxigenin antibody was used. Full length probes (median length 600 base pairs) were hydrolysed with alkali to an average size of 150 nucleotides (100–200 base pairs), as estimated by non-isotopic Northern blotting.

**NON-ISOTOPIC IN SITU HYBRIDISATION**

**Proteolysis**

Nucleic acids were “unmasked” by exposing sections to pepsin/HCl or proteinase K for 20 and 15 minutes respectively at 37°C. Pepsin/HCl (specific activity = 3-200 IU/mg protein) was prepared as described previously. Proteinase K was dissolved to a concentration of 500 μg/ml (specific activity = 20 IU/mg protein, Boehringer Mannheims, UK), in phosphate buffered saline (PBS) (10 mM NaH₂PO₄ 1-5 mM KH₂PO₄, 150 mM NaCl, pH 7.4). After proteolysis sections were quickly rinsed in PBS, ddH₂O, and post-fixed in 2% paraformaldehyde (w/v) in PBS for five minutes, followed by quick rinses in PBS, ddH₂O, methanol, and dried at 37°C.

Prehybridisation/hybridisation

To assess the effect of prehybridisation sections were covered with 140 μl of hybridisation mix without probe; PBS alone was used as a control. Sections were incubated in moist Terasaki plates for 30 minutes at 37°C. Slides were drained and 8 μl of hybridisation mix containing digoxigenin labelled (0-4 ng/ml), or biotinylated antisense riboprobe (0-06 ng/ml), was pipetted on to each well and individual wells covered with a circular glass coverslip (14 mm in diameter); these were not sealed. Hybridisation mix contained 50% (v/v) formamide, 5% (v/v) dextran sulphate, 2 x SSC, 0-1 mM EDTA, 1 mM TRIS-HCI (pH 7.3), and *Escherichia coli* tRNA to a final concentration of 100 mg/ml; 1 x SSC = 0-15 M sodium chloride, 0-015 M sodium citrate. All solutions used in the hybridisation mix were prepared using ddH₂O that had been previously treated with 0-1% (v/v) diethyl pyrocarbonate (DEPC).

The multi-well slides in moist Terasaki plates were subsequently treated as follows. One set was heated at 95°C in a hot air oven for 15 minutes and hybridised at 42°C or 50°C for two, four, six, or 16 hours. The second set were hybridised (without preheating at 95°C) at 42°C or 50°C for the same times. Each set was also hybridised with hydrolysed and non-hydrolysed riboprobes under both sets of conditions.

Negative controls consisted of sections on the same multi-well slide incubated with the same quantity of labelled lysozyme sense probe, or with hybridisation mix only; unrelated digoxigenin labelled p53 sense and anti-sense riboprobes were also hybridised to consecutive sections. Competitive studies were carried out by incubating sections with hybridisation mix containing both labelled anti-sense and sense lysozyme riboprobes.

**Post-hybridisation treatment**

Coverslips were removed by washing for 10 minutes in 2 x SSC at 22°C (twice, for six minutes each) and sections incubated with 150 μg/ml RNase A (Sigma UK) in 2 x SSC for 30 minutes at 37°C (140 μl per well). To investigate the effect of post-hybridisation washes on background noise, slides were washed at higher stringency in 0-2 x SSC/50% formamide at 50°C for 30 minutes. Slides were then washed in TBT blocking solution (0-05 M TRIS/HCl, 0-1 sodium chloride, pH 7-5, containing 0-25% (w/v) bovine serum albumin and 0-05% Triton X100 (w/v) at 22°C for 10 minutes.

**DETECTION OF NISH SIGNAL**

**Biotinylated probes**

Slides were incubated in a moist chamber at 22°C for 30 minutes with mouse monoclonal anti-biotin antibody (Dako UK) diluted 1 in 50 in TBT; biotinylated (Fab')2 rabbit anti-mouse immunoglobulins (Dako UK) diluted 1 in 200 for 30 minutes; washed in TBS (50 mM TRIS/HCl, pH 7-2, 100 mM NaCl, 1 mM MgCl₂), and incubated for 30 minutes with alkaline phosphatase or peroxidase conjugated streptavidin (Dako UK) diluted 1 in 50 or 1 in 75, respectively, in TBT containing 5% non-fat dried milk (Cadbury UK). After two five minute washes in TBS alkaline phosphatase was detected by incubation in nitroblue tetrazolium containing 5-bromo-4-chloro-3-indolyl-phosphate (NBT/BCIP) substrate in the dark—that is, in a drawer or regular cupboard, for one to 16 hours at 22°C. Peroxidase was detected by incubating in 3-amoine-9-ethylcarbazole (AEC), hydrogen peroxide.
mRNA NISH in archival biopsy specimens

**Background**

mRNA detection in archival biopsy specimens can be achieved using mRNA in situ hybridisation (NISH) techniques. This method involves the hybridisation of labelled riboprobes, which are then detected using a range of histochemical methods.

**Methods**

1. **Preparation of Tissue Specimens**
   - The specimens are treated with RNAse inhibitors and DNAse to prevent degradation of RNA or DNA.
   - The tissue fragments are fixed in paraformaldehyde and post-fixed in TBT.
   - The slides are then washed in TBS and dried in a stream of heated air.

2. **Hybridisation**
   - The riboprobes are hybridised to the tissue sections at optimal conditions.
   - The hybridisation is carried out at 50°C, as this temperature is optimal for many riboprobes.

3. **Detection**
   - The hybridised riboprobes are detected using a variety of methods, such as digoxigenin-labelling and alkaline phosphatase conjugation.
   - The labelled signals are visualised using alkaline phosphatase substrates, such as Fast Red or BCIP.

4. **Counterstaining**
   - The sections are counterstained with methyl green or haematoxylin and eosin, and mounted in glycerol jelly.

**Results**

The most widely used protocols for mRNA detection with riboprobes require hybridisation at 42°C–50°C for 12 to 16 hours. This is usually preceded by prehybridisation treatment of slides and followed by post-hybridisation digestion with RNAses. We used this protocol as the “comparator system” against which we judged the effect of modifying the variables tested (see below). The protocol which has been evolved detected lysozyme mRNA in Paneth cells, Brunner’s glands, and in some cases, mononuclear cells in the lamina propria (fig 1, fig 3A).

**Hybridisation Temperature**

The intensity of the signal increased slightly when the hybridisation temperature was increased from 42°C to 50°C (fig 2A). The use of higher temperatures (55°C, 60°C, and 65°C) did not increase signal intensity over that at 50°C (table). Drying of the sections was a problem using sealed slides above 55°C.

**Preheating Target and Probe**

When tissue target and probe were heated for 15 minutes at 95°C simultaneously on the slide before hybridising, the signal intensity was

**Figure 1** Lysozyme mRNA in Paneth cells and Brunner’s glands in an archival small bowel biopsy specimen. This section was hybridised with (A) biotin and (B) digoxigenin labelled lysozyme anti-sense riboprobe using optimal conditions for mRNA detection. Background is higher with the biotinylated probe.

**Figure 2** (A) Small bowel biopsy specimens hybridised for two hours at 50°C, without a preheating step, with digoxigenin labelled lysozyme anti-sense riboprobe. Paneth cells are just visible (arrow) using the three-step alkaline phosphatase procedure. (B) Small bowel biopsy specimens, heated at 95°C for 15 minutes with digoxigenin labelled lysozyme anti-sense riboprobe, and hybridised for two hours at 50°C. Paneth cells are intensely labelled and lamina propria macrophages are visible.
greatly increased and the number of cells in which lysozyme mRNA was detected also increased. With this initial heating step the optimum signal was achieved by hybridising subsequently at 50°C for two hours (fig 2B); prolonged hybridisation did not significantly increase the signal (table). In practice, therefore, we heat at 95°C for 15 minutes and hybridise at 50°C for two hours.

ENZYME DIGESTION AND PREHYBRIDISATION OF TISSUE SECTIONS

Using the conditions outlined in the preceding paragraph, pepsin-HCl digestion of target was superior to that of proteinase K. Although proteinase K gave a slightly more intense signal, the background was higher and tissue morphology poorer. Prehybridisation with hybridisation mix without probe did not improve the signal to noise ratio obtained with riboprobes in these tissues.

HYDROLYSED AND NON-HYDROLYSED PROBES

Hydrolysis of lysozyme riboprobes (to a median size of 100–200 base pairs) before hybridisation did not increase signal intensity. In fact, unhydrolysed probes, median size 600 base pairs, gave a more intense signal (figs 3A, 3B, table).

PROBE DETECTION

The most sensitive detection system was the three-step procedure using NBT/BCIP as substrate for alkaline phosphatase. The signal appeared at almost the same time in Brunner’s glands and Paneth cells, and was optimal in about 20–30 minutes. To obtain a signal in lamina propria macrophages it was necessary to incubate in substrate for 45 to 60 minutes. While the signal obtained with digoxigenin probes stood out very clearly from a clean background, biotinylated probes produced higher background with these substrate incubation times. With biotinylated probes the background level was high after 45 minutes’ incubation in substrate and unacceptable after incubations longer than one hour. NBT/BCIP gave a much more intense signal than Fast red; it was also superior to peroxidase detection of probes by AEC/H2O2.

RNASE INHIBITORS

The use of general measures to protect against free RNAses (gloves, washes with DEPC treated water, and glassware treated with RNase inhibitors) did not improve the intensity of the signal obtained without using gloves, washing the slides with ultrapure water only, and using only clean glassware. All of our reagents, however, were made up in DEPC treated water.

To examine the effects of fixation on mRNA retention four small bowel biopsy specimens were routinely fixed for two to 48 hours in 10% buffered formalin and paraffin wax embedded. Sections were hybridised under optimal conditions with digoxigenin labelled riboprobes (table). The intensity of signal in all cell types was similar, irrespective of fixation time.

<table>
<thead>
<tr>
<th>Probe*</th>
<th>Heating (95°C/15 minutes)</th>
<th>Temperature</th>
<th>Time (hours)</th>
<th>Signal intensity†</th>
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*All probes were digoxigenin labelled.
†Signal intensity was scored independently as 1 + to 5 + by two observers; ± indicates that a signal was just visible.
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CONTROLS
All of the controls with labelled sense probes and hybridisation mix only were negative. The p53 anti-sense probe did not react with Paneth cells or any of the other cell types that reacted with the lysozyme riboprobe. Sections hybridised with both labelled lysozyme anti-sense and sense probes showed a reduction in signal intensity in this competitive reaction. The incubation with RNase A before hybridisation abolished the signal. Digestion with DNase before hybridisation either did not affect or increase signal intensity. RNase digestion after hybridisation was effective in reducing background but did not affect the specific signal in Paneth cells, etc.

Discussion
We investigated the transcription of the lysozyme gene and its product in routinely processed tissue sections of human bowel. The heating step at 95°C for 15 minutes is the most relevant innovation to improve the sensitivity of NISH on archival tissue sections. Heating before hybridisation produces a strong signal without background after short hybridisations (two hours), which is not attainable without heating. Results, therefore, can be obtained in one day. The heating step probably removes secondary structure from the mRNA, and perhaps riboprobe.

The model used here shows the advantages of digoxigenin as a reporter molecule. The pattern and intensity of the signal obtained with riboprobes labelled with biotin or digoxigenin was the same, but biotinylated probes gave a much higher background that was not abolished even with post-hybridisation RNase treatment. This was particularly so with long enzyme substrate incubations. A higher signal to noise ratio, therefore, was obtained with digoxigenin labelled probes, particularly with the three-step detection system. Prehybridisation treatments and high stringency post-hybridisation washing did not reduce non-specific background, irrespective of the reporter molecule, suggesting that background is not due to specific hybridisation of probe to tissue components.

Most of the protocols for mRNA detection with riboprobes use hydrolysed probes of 150–200 base pairs, the reasoning being that short fragments penetrate more easily into cells. In our experiments, however, hydrolysis of RNA probes 600 to 100–200 base pairs did not give a signal as intense as non-hydrolysed probes. This would argue that probe size is not the only element involved in the production of the final signal.

The identification of lysozyme protein has been used as a marker of macrophages in a variety of organs. In our experiments lysozyme mRNA in lamina propria cells in normal bowel was almost undetectable. A positive signal for lysozyme mRNA in lamina propria cells, however, was obtained in the rectal biopsy specimens of Crohn’s disease. Taken together, these observations suggest that lysozyme transcription and translation are upregulated in those macrophages present in inflammatory disease.

Other protocols for mRNA detection by in situ hybridisation emphasise that, as RNAses are ubiquitous, special measures are necessary, such as gloves, washes with DEPC treated H₂O₂, and treatment of glassware with RNase inhibitors, to avoid degradation of mRNA. In our experiments all the reagents were made up with DEPC treated water. Incorporating the other measures was found to be unnecessary. Formalin fixation of tissue may protect the mRNA in the nucleolus by cross-linking them to tissue proteins.

In conclusion, this methodology gave consistent results in 16 endoscopic small bowel and two rectal biopsy specimens as well as four small bowel surgical samples that were morphologically normal or chronically inflamed. Heating at 95°C for 15 minutes before hybridisation allows hybridisation time to be reduced to two hours, compared with the standard overnight incubation procedures using riboprobes. Thus the technique can be easily completed in one working day and may be applicable for use in routine laboratories.

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