Non-isotopic detection of in situ nucleic acid in cervix: an updated protocol

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SUMMARY  Recently, sensitive non-isotopic in situ hybridisation (NISH) methodology for the detection of human DNA and human papilloma virus (HPV) DNA in archival paraffin blocks of cervix was described. An amended protocol, now used in this laboratory for detection of these genes by NISH is presented. The amendments include the following: protease digestion at 37°C; tissue dehydration in air rather than ethanol; stringency washing in formamide solution; blocking non-specific binding of avidin alkaline phosphatase with a modified buffer; and increasing the concentration of avidin alkaline phosphatase for detecting low abundance DNA.

These changes simplify and increase the sensitivity of the protocol such that "Y" chromosome repeats are visualised in almost all female cells.

Visualisation of pHy2.1 autosomal repeat sequences has been used as a built-in sensitivity control for detection of human papillomavirus (HPV) nucleic acid in formalin fixed paraffin wax embedded cervical biopsy specimens. Using archival paraffin wax embedded material from the 1970s, we recently found that the pHY2.1 signal varies in consistency and intensity. In this paper we report modifications to the earlier protocol which increases the consistency and intensity of the pHy2.1 signal in this type of biopsy specimen; by extrapolation and experiment the HPV signal detected by in situ hybridisation is also increased.

Material and methods

1 Mount formalin fixed paraffin wax sections on glass multispot slides treated with aminopropyltriethoxysilane, dry at 37°C for 30 minutes, and bake sequentially at 75°C for 60 minutes and overnight at 60°C. In urgent cases biopsy specimens can be satisfactorily probed after the 75°C baking step. Sections can be stored indefinitely at 22°C.

2 Dewax sections by heating at 75°C (15 minutes) and transferring immediately to xylene at 22°C, changing xylene (10 minutes twice). Wash in 99% ethanol (10 minutes twice), wash in running tap water (five minutes), and preheat sections in distilled water in a coplin jar, kept at 37°C in a high water line water bath. Viral and human nucleic acids are unmasked by exposing sections to pepsin at 5 mg/ml (ca 2500 units/mg protein (Boehringer Mannheim, UK)) in 0.2 M hydrochloric acid for 15 minutes at 37°C. Pepsin, 500 mg, is dissolved by gentle agitation in 96 ml distilled water, preheated at 37°C, and 4 ml of 5 M hydrochloric acid added to a final concentration of 0.2 M hydrochloric acid. Refractory material is unmasked with proteinase K at 0.5 to 1 mg/ml (ca 20 units/mg protein (Boehringer Mannheim, UK)) in 0.01 M phosphate in 0.15 M sodium chloride, pH 7.4 (PBS), for 10 to 15 minutes at 37°C.

3 After digestion, wash sections thoroughly in PBS and dry at 37°C (five to 15 minutes).

4 Aliquots (9 μl) of hybridisation mix containing 10 ng of appropriate biotinylated probe are added to each well on multispot slides, covered with 14 mm diameter coverslips, and the slides placed in a moist, sealed Terasaki plate. Hybridisation mix consists of 50% formamide (Sigma, UK), 5% dextran sulphate (BDH, UK), 2 × SSC, 0.1 mM edetic acid, 0.5 μg sheared salmon sperm DNA, and 0.05 mM Tris-hydrochloric acid, pH 7.3; 1 × SSC = 0.15 M sodium chloride, 0.015 M sodium citrate.

5 Denature at 95°C for 15 minutes on a solid stainless steel plate in a hot air oven and hybridise at 42°C for two hours.

6 Wash slides at low stringency in three changes of 4 × SSC at 22°C (five minutes each change). If high stringency washing is required for specific HPV typing rather than human gene detection, the 4 × SSC washes are followed by two washes in 45% formamide in 1 × SSC at 37°C, pH 7.2 (10 minutes each wash), and 4 × SSC at 22°C for five minutes. After these

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washes, soak in blocking agent TBT (0-01 M Tris-hydrochloric acid, 0-15 M sodium chloride (pH 7-5)) containing 0-25% (w/v) bovine serum albumin (Sigma UK; No. A7906) and 0-05% Triton X 100 (v/v) at 22°C for 10 minutes.

7 Incubate sections at 22°C for 30 minutes with modified avidin alkaline phosphatase conjugate (Dako) diluted in TBT. Dilutions of 1/25 and 1/200 are satisfactory for low and high abundance nucleic acid detection, respectively.

8 Remove unbound conjugate by washing twice for five minutes in 0-01 M Tris-hydrochloric acid, 0-15 M sodium chloride (pH 9-5) and incubate for five to 30 minutes in substrate consisting of nitroblue tetrazolium (NBT) and 5-bromo-4-chloro-3-indolyl-phosphate (BCIP).3 In practice, the NBT-BCIP substrate/buffer is made up as follows.

(i) NBT (10 mg) (Sigma; N6876) is weighed into an Eppendorf tube. Dissolve in 0-2 ml dimethylformamide and add 1 ml of substrate buffer (0-1 M Tris-hydrochloric acid, 0-1 M sodium chloride, and 0-005 M magnesium chloride, pH 9-5). Transfer to a larger vessel containing 30 ml of substrate buffer at 37°C.

(ii) Weigh out 5 mg of BCIP into a glass tube and dissolve in 0-2 ml of dimethylformamide.

(iii) The BCIP solution is added in drops to the NBT solution and stirred. This final substrate mix is stored in 5 ml aliquots at −20°C until required.

9 A signal is evident in one to two minutes when HPV copy number is high. Terminate reaction by washing for five minutes in tap water and immersing for five minutes in 10% formalin containing 0-15 M sodium chloride. The final preparations are rinsed in

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Fig 1a Detection of autosomal "Y sequences" with old protocol in archival cervical biopsy specimen. (Not counterstained). Nuclei are just visible at this low magnification.

Fig 1b Parallel section to 1a but "Y sequences" detected in situ by new protocol. (Not counterstained). "Y sequences" are clearly evident in nuclei, at identical magnification.

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distilled water, counterstained with pyronin, light haematoxylin or light green if required, and mounted in glycerine jelly or crystal/Mount (Biomeda, USA). This last may be postmounted in De Pex (RA Lamb, UK).

Results

Results with the old \(^1\) and present procedure are illustrated in figs 1a and b.

Discussion

Five principal modifications to the original method have been described. Firstly, nucleic acids are unmasked by pepsin/hydrochloric acid or proteinase K/PBS at 37°C. Secondly, the ethanol dehydration step before in situ probing has been amended. Thirdly, the stringency washing stage has been improved. Fourthly, a different blocking and washing buffer has been introduced which increases the sensitivity and simplifies the procedure. Fifthly, avidin-alkaline phosphatase conjugates have been used at different dilutions. These modifications give the most consistent and sensitive results for human papilloma virus (HPV) detection by non-isotopic in situ hybridisation (NISH).

With pepsin/hydrochloric acid, blue nucleic acid signals are visualised against an essentially clear background. Proteinase K not only obtains viral nucleic acid signals but also enhances greatly the autosomal Y signal (fig 1b). This may be due to its greater effectiveness in digesting native proteins. It also tends to induce a pale pink cellular background with viral probes, seldom seen after pepsin/hydrochloric acid treatment.

Interestingly, the autosomal Y signal is lost after high stringency washings. The Y signal from Y bodies in sections of male tonsil withstands such treatment (unpublished observation). This finding suggests that autosomal Y repeat sequences in female cells are only weakly homologous to repeat sequences in the Y body of male cells, or that the stringency conditions in NISH are not entirely similar to standard nucleic acid analytical procedures. Further studies on this phenomenon are clearly necessary.

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References


Requests for reprints to: Professor J O'D McGee, Nuffield Department of Pathology, John Radcliffe Hospital, Level 1, Headington, Oxford OX3 9DU, England.
Mucins in Histopathology

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Four symposia will cover the topics: transplantation; image analysis; quantitation of immunofluorescence and intracellular probing; and leukemia/lymphoma immunophenotyping.
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For detailed information regarding the programme, advance registration, and submission of abstracts write to: Mariano La Via, Clinical Applications of Cytometry, P.O. Box 39778, Charleston, South Carolina 29407, USA.

External Quality Assessment in Histopathology and Cytology
External Quality Assessment is of increasing importance in histopathology and cytology. With the advent of the cytology proficiency testing scheme and the need to control quality of the newly implemented breast cancer screening, the Royal College of Pathologists and the Association of Clinical Pathologists are presenting a special one day symposium at St Bartholomew's Hospital, London EC1, on Monday April 24, 1989. Fee £40.00 including buffet lunch.
Further details from: Professor G Slavin, Department of Histopathology, St Bartholomew's Hospital, West Smithfield, London EC1.

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The Association of Clinical Pathologists runs a locum bureau for consultant pathologists.
Applicants with the MRC Path who would like to do locums and anyone requiring a locum should contact Dr DH Orrell, Department of Pathology, Royal Lancaster Infirmary, Ashton Road, Lancaster LA1 4RP.

Corrections
The references to the paper by Burns, Graham, and McGee were incorrect at time of press (J Clin Pathol 1988;41:897–99) and should be as follows:

References

An error was inadvertently made to the title of the paper by Dr Savitzipori (J Clin Pathol 1988;41:1099–103). It should have read: Cerebral involvement associated with Escherichia coli 0157: H7 in humans and gnotobiotic piglets.

Broadsheet 120 January 1989: Folio 7 of the above broadsheet: (Guide to Diagnosis of inborn errors of metabolism in district general hospitals, has been reprinted owing to a series of printing errors (enclosed with this issue).