Early detection of *Chlamydia trachomatis* using fluorescent, DNA binding dyes

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**SUMMARY** HeLa 229 cells were infected with genital tract strains of *Chlamydia trachomatis*. After incubation for varying times the infected cells were fixed and stained with the fluorescent DNA binding dyes Hoechst 33258 or DAPI for comparison with conventional Giemsa stain. Fluorochrome-treated preparations were examined by incident ultraviolet fluorescence microscopy and the Giemsa-stained preparations by dark-ground light microscopy. Chlamydial inclusion bodies could be identified unambiguously as early as 18 hours after infection of HeLa 229 cells using either Hoechst 33258 or DAPI but not until some 48 hours in Giemsa-stained preparations. The DNA rich chlamydial elementary bodies in infected egg yolk suspension were readily detected using Hoechst 33258. The fluorescent dye technique was simpler and more rapid than Giemsa staining. Using Hoechst 33258 it is possible to speed up the identification of chlamydial isolates growing in tissue culture.

Chlamydiae are prokaryotic microorganisms structurally related to bacteria but lacking the energy systems necessary for independent existence. Unlike bacteria they undergo a complex growth cycle initiated by infection of the host cell with DNA rich, 0.3 μm diameter, elementary bodies. The elementary bodies enlarge to form the metabolically active, 1 μm diameter, RNA rich, reticulate bodies. These reticulate bodies divide and multiply within host cell vacuoles, giving rise to a characteristic inclusion body. Ultimately, elementary bodies differentiate from the reticulate bodies and are released from the cell some 48-72 hours after infection.

*Chlamydia trachomatis* infections are diagnosed by growth of the agent in specially prepared tissue culture cells for some three days. The culture is then examined for the characteristic, glycogen-containing inclusion bodies using iodine, fluorescent antibody, or Giemsa stain. Giemsa-stained inclusions are more readily visualised using dark-ground microscopy (Darougar *et al.*, 1971). This paper describes a novel method of staining chlamydial inclusions, which permits the unambiguous detection of developing chlamydial reticulate bodies in tissue culture for the common genital tract strains after only 18 hours’ growth. The method exploits the ability of Hoechst 33258, a benzimidazole compound, specifically to form a fluorescent complex with DNA but not RNA by selective binding to the adenine-thymine rich regions of double-stranded DNA (Arndt-Jovin and Jovin, 1977).

**Methods**

**Material**

*C. trachomatis* strains UW4/GCX (serotype H), IOL-238/R (serotype G), and LGV.440/LN (serotype L1) were kindly provided by Dr S. Darougar, Institute of Ophthalmology, London. The strains were passaged once through the yolk sac of embryonated hens’ eggs and then frozen at −70°C.

Hoechst 332581 was obtained through the kind auspices of Mr J. W. Neaves, Hoechst UK Limited, Salisbury Road, Hounslow, Middx, UK. DAPI (4′-6-diamidino-2-phenylindole hydrochloride) was originally obtained from Professor O. Dann, Institut für angewandte Chemie, 852 Erlangen, West Germany and is now available commercially from Serva Feinbiochemica, PO Box 10 52 60, Heidelberg D-69, West Germany. Diethylamino ethyl dextran

1Bisbenzimide trihydrochloride. Now available from Aldrich Chemical Co Ltd, Gillingham, Dorset, UK

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(DEAE dextran) of molecular weight 500000 was obtained from Sigma Chemicals, Poole, Dorset, UK.

Tissue Culture
HeLa 229 cells (Flow Labs, Irvine, Scotland) were grown to confluent monolayers on 13 mm diameter coverslips in disposable bijoux. Each bijou contained 1 ml of Eagle's Minimal Essential Medium, 10% v/v Foetal Calf Serum, 1% v/v non-essential amino acids, L-glutamine and sodium bicarbonate (Flow Labs) without antibiotics. The bijoux were inoculated with 100 μl volumes of a 1 in 100 dilution of 50% egg lethal dose chlamydiae and incubated for 3 hours at 35°C for UW4 and IOL 238 or 37°C for LGV to permit chlamydial adsorption. Before adsorption the infectivity of strains UW4 and IOL 238 was potentiated by washing the cells twice with 45 μg/ml of DEAE-dextran in Hanks salts followed by centrifugation of the chlamydiae on to the cells at 900 g for 1 hour (Kuo et al., 1971; 1972).

Staining
Chlamydial infected monolayers on 13 mm diameter No. 1 coverslips were washed twice with complete Dulbecco's phosphate buffered saline (PBS, Oxoid Limited) and fixed for 10 minutes with absolute methanol. The fixed cells were washed twice with PBS and then incubated for 10 minutes at 37°C with a freshly prepared solution of 10 μg/ml of Hoechst 33258 or DAPI in PBS. Stained cells were washed twice with PBS, dried, and mounted in 50% v/v glycerol in PBS cell surface down on a microscope slide for fluorescence microscopy. For comparative purposes chlamydial infected cells were stained in parallel with Giemsa stain (Johnson, 1975). As both Hoechst 33258 and DAPI bind to DNA they are possible carcinogens and should be handled accordingly.

Microscopy
Hoechst 33258 or DAPI stained material was examined with an Ortholux II microscope (E. Leitz, Wetzlar, West Germany) fitted with an HBO 200 ultraviolet source and Ploem 1 incident light illumination. The filters used were: a 4 mm BG38 in the lamhouse, 2 × 2 mm UG 1 filters for narrow band excitation at 365 nm, a TK 400/K400 dichroic mirror and suppression filter, and a K460 (460 nm) barrier filter.

Giemsa-stained material was examined using an Ortholux II microscope equipped with an oil immersion NA 1·2 dark-ground condenser.

Results
Figure 1 is a fluorescence micrograph showing the result of staining uninfected HeLa 229 cells with Hoechst 33258. Figure 2 shows the results of a parallel experiment, in which the cells were infected with LGV 440 for 48 hours. Chlamydial inclusions can readily be detected between the brightly fluorescent host cell nuclei. Lymphogranuloma venereum (LGV)

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**Fig. 1** Uninfected HeLa 229 cells monolayer stained with Hoechst 33258 for 10 minutes and examined by incident ultraviolet fluorescence microscopy. Only the host cell nuclei are stained. × 400.
chlamydiae differ from United Kingdom isolates of *C. trachomatis* in that they have a shorter growth cycle, do not require centrifugation to establish *in vitro* infection, and form larger inclusions. The experiment was repeated using two typical genital tract *C. trachomatis* strains, UW4 and IOL 238, taking samples after 1, 12, 18, 24, 36, 42, 48, 60, and 72 hours' incubation and comparing in parallel Hoechst 33258 and Giemsa-stained material.

In the Hoechst 33258 stained preparations, no significant differences could be seen between uninfected and chlamydial infected cells after 12 hours' incubation. However, after 18 hours' incubation definite chlamydial inclusions could be seen in the infected cells (Fig. 3). These inclusions were characteristically granular, consisting of approximately 1 μm reticulate bodies, and, being less rich in DNA, they were less fluorescent than the host cell nucleus. By 24 hours the inclusions were noticeably enlarged and at 36 hours were much more brightly fluorescent due to the development of DNA rich elementary bodies. The inclusions continued to enlarge up to 60 hours' incubation (Fig. 4), and by 72 hours large numbers of chlamydiae had been released from the cells. After 72 hours' incubation brightly fluorescent (DNA rich) particles of a similar size to chlamydial elementary bodies could be seen attached to the surface of the tissue culture cells (Fig. 5). Small, immature inclusions were also noted in 72-hour preparations as well as the large primary inclusions (Fig. 5). In view of the experimental conditions used, in which the concentration and molecular weight of DEAE-dextran was optimally titrated for the chlamydial strain so as to achieve a high rate of infectivity, it is believed that these small inclusions represent a limited cycle of re-infection occurring in the absence of further centrifugation. Mature chlamydial inclusions of strains UW4 or IOL 238 developing after 60 or 72 hours' incubation were readily detected by dark-ground microscopy in Giemsa-stained preparations (Fig. 6). However, in our culture system immature chlamydial inclusions in HeLa 229 cells infected for less than 48 hours could not be unambiguously detected by the dark-ground Giemsa stain technique. Chlamydial elementary bodies with their high ratio of DNA:RNA were readily identified as small, brightly fluorescent particles against an unstained background in Hoechst 33258 stained smears of chlamydial infected egg yolk sac (Fig. 7).

**Discussion**

Non-gonococcal urethritis (NGU) is now more common than gonorrhoea and *C. trachomatis* can be isolated from approximately half these patients. Such infections cannot be regarded as trivial as *C. trachomatis* has been isolated direct from the
Fig. 3  18-hour inclusions (arrowed) of C. trachomatis UW4 in HeLa 229 cells after staining with Hoechst 33258. The characteristically granular nature of the inclusions can be seen. × 1000.

Fig. 4  DAPI stained HeLa 229 cells after infection with C. trachomatis IOL 238 for 60 hours. The chlamydial inclusions are now larger and more brightly fluorescent due to the presence of large numbers of the DNA rich elementary bodies. × 400.
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Fig. 5  HeLa 229 cells after infection for 72 hours with C. trachomatis UW4, stained with Hoechst 33258. Brightly stained, small, DNA rich particles (solid arrows) can be seen adherent to the surfaces of the tissue culture cells; these particles are comparable to those seen in chlamydial infected egg yolk sac (Fig. 7) and represent chlamydial elementary bodies. A mature primary inclusion (*) is present as well as two small inclusions (hollow arrows) comparable to those seen after 18 hours' infection (Fig. 3). These may be late developing primary inclusions or, more likely, a new cycle of infection in the previously uninfected cells. × 550.

diseased fallopian tubes in 30% of patients with acute salpingitis (Mårdh et al., 1977) while maternal infections have been associated with prematurity, pneumonia, and eye infections in the newborn (Rees et al., 1977; Beem and Saxon, 1977). Chlamydial infections are resistant to penicillins, cephalosporins, chloramphenicol, and aminoglycosides. Therefore, only rapid diagnosis would permit early treatment with erythromycin or tetracyclines.

Chlamydiae can be rapidly detected using fluorescent antibody techniques (Thomas et al., 1977). However, appropriate fluorescent antisera are not readily available, and, like other fluorescent antibody techniques, the method is time-consuming and requires considerable expertise. For this reason, relatively insensitive and non-specific conventional staining techniques are most commonly used for detecting chlamydiae, the morphology of mature chlamydial inclusions being sufficiently characteristic to permit definite identification. However, diagnosis of chlamydial infection might be made earlier by combining the specificity of modern DNA binding dyes with the sensitivity of fluorescence techniques.

Acridine orange is the classic fluorescent dye for staining virus-infected cells. Unfortunately, acridine orange reacts with RNA as well as DNA and was found to stain both the host cell cytoplasm and the developing reticulate bodies, making it difficult to detect early chlamydial maturation. Recently, two fluorescent dyes, DAPI and Hoechst 33258, have become available which act by non-intercalative binding to the adenine-thymine rich areas of double-stranded nucleic acid and which are highly specific for DNA (Arndt-Jovin and Jovin, 1977).
Fig. 6 Monolayer of HeLa 229 cells heavily infected for 72 hours with C. trachomatis UW4, stained with the conventional Giemsa method, and examined by dark-ground light microscopy. 72-hour inclusions such as these (arrows) are readily visualised by both techniques. However, unlike the Hoechst stain, the Giemsa method does not permit ready identification of early inclusions such as those of Fig. 3. × 350.

DAPI has been used for the spectrofluorimetric assay of DNA and, in the presence of RNA, can detect as little as $5 \times 10^{-10}$ g/ml of DNA (Kapuściński and Skoczylas, 1977). DAPI can be used for the detection of DNA viruses and mycoplasma in vitro (Russell et al., 1975). Most of the experiments described here were performed using Hoechst 33258 because of its ready availability; however, comparable results were obtained using DAPI, and this is now commercially available.

The method described has significant advantages. The staining technique is rapid, taking some 15 minutes as opposed to the 1 hour plus of Giemsa or fluorescent antibody staining. Only chlamydiae and host cell nuclei are stained; there is thus no possibility of confusing chlamydiae with host cell cytoplasmic components. Moreover, with incident or transmitted dark-ground illumination the chlamydiae are viewed under ideal contrast conditions: bright light blue fluorescence against a black background. The technique is far more sensitive than Giemsa staining; with both DAPI and Hoechst 33258 chlamydial inclusions in HeLa 229 cells can readily be identified after 18 hours’ growth. The presence of 1 μg/ml of emetine hydrochloride in the tissue culture medium during incubation of the chlamydial infected cells markedly increases the size of early (18 and 24 hour) chlamydial inclusions and prevents overgrowth of the host cells, making identification even easier. Emetine selectively blocks protein synthesis in eukaryotic host cells but leaves chlamydial (prokaryotic) protein synthesis unimpaired, favouring chlamydiae in their competition with the host cell for intracellular amino acid pools (Becker and Asher, 1972).

In our laboratory, clinical specimens for chlamydial culture are centrifuged onto monolayers of DEAE-dextran treated HeLa 229 cells followed by incubation of the infected cells in emetine-containing tissue culture medium before Hoechst staining. Isotope studies on chlamydial amino acid utilisation by strain UW4 growing in emetine-treated cells (data not shown) shows there is little point in incubating infected cells beyond some 40 hours. Beyond this point, loss of cells from the monolayer due to chlamydial release and the effects of emetine...
continuing

exceeds the rate of isotope incorporation due to
continuing chlamydial multiplication. In practice,
no difficulties have been experienced in identifying
Hoechst stained inclusions from clinical specimens
in HeLa cell monolayers. Most of the DNA-
containing debris likely to be encountered—
bacteria, yeasts, white cells, epithelia—is washed
from the preparations before and after staining, and
it is, in any case, quite unlike the characteristic
chlamydial inclusions. Inclusions of DNA-contain-
ing viruses (eg, herpes, adenovirus) could theoret-
cally be detected by Hoechst staining; indeed,
is this might be useful for the diagnosis of non-
bacterial eye infections. However, Hoechst-stained
virus inclusions are unlikely to be confused with
chlamydial inclusions and would not be expected to
develop in metabolically inhibited emetine or
cycloheximide treated host cells. Hoechst stain has
been successfully used in our laboratory for the
direct microscopic detection of chlamydiae in
clinical specimens. However, insufficient experience
has yet been gained to know how useful it will be in
this respect when compared with Giemsa stain.

The disadvantage of the Hoechst stain is its
requirement for a fluorescence microscope with
suitable filtration for the 365 nm excitation and
450-500 nm emission spectrum of the dyes. How-
ever, most diagnostic microbiology laboratories
have facilities for fluorescence microscopy. Incident
light illumination of the specimen is preferable, but
dark-ground transmitted illumination can be used.
Thus, for most laboratories the only extra capital
cost will be a suitable excitation and barrier filter.

By comparison with fluorescein conjugates used for
immunofluorescence, Hoechst 33258 stained material
usually fluoresces much brighter, is stable on storage,
and is more resistant to ultraviolet induced fading.

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