BACTERIAL CHARACTERISATION

Growth and cultural characteristics of Calymmatobacterium granulomatis – the aetiological agent of granuloma inguinale (Donovanosis)

A. B. M. KHARSANY, A. A. HOOLEN, P. KIEPIELA, T. NAICKER* and A. W. STURM

Department of Medical Microbiology and *Electron Microscope Unit, Faculty of Medicine, University of Natal, Durban, South Africa

Granuloma inguinale is a chronic destructive granulomatous disease of the genitalia. The clinical diagnosis is often unreliable and the definitive diagnosis is based on the visualisation of ‘Donovan bodies’ in tissue smears or biopsy specimens. The organism implicated in its aetiology, Calymmatobacterium granulomatis, was reported to have been cultured >30 years ago, but little is known about the organism because of its fastidious nature and the difficulty in culturing it. Twenty-two biopsy specimens from female patients with clinical and laboratory-confirmed granuloma inguinale were treated with amikacin 10 mg/L and inoculated in a monocyte co-culture system with peripheral blood mononuclear cells (PBMC) from a single donor and autologous sera. The method was subsequently modified by pretreatment of specimens with vancomycin 5 mg/L and metronidazole 10 mg/L in addition to amikacin 10 mg/L for the purpose of decontamination, pooled blood donor PBMC and by the use of heat-inactivated fetal calf serum instead of autologous serum for culture. This modified method was used to culture additional biopsy specimens and genital ulcer scrapings from female and male patients, respectively. All monocyte co-cultures were examined by a rapid Giemsa (RapiDiff) stain and by an indirect immunofluorescence test with immune sera. Representative cultures were examined by transmission electron microscopy. C. granulomatis was successfully isolated in pure culture by the monocyte co-culture system from four biopsy specimens and 14 genital ulcer scrapings. The cultured organisms were visible both intra- and extra-cellularly and were extremely pleomorphic, with characteristic single and bipolar condensation. The numbers of the organisms increased after each passage. All positive cultures showed bright fluorescence when tested with immune sera. Transmission electron microscopy of the cultured bacteria demonstrated a typical gram-negative cell wall consisting of an outer membrane, middle electron opaque layer and an inner plasma membrane. The capsule was thick and electron dense. Numerous electron dense granules were present within the cytoplasm.

Introduction

Granuloma inguinale, also known as Donovanosis, is a chronic granulomatous disease involving the genitalia and surrounding sites. It is found in specific geographical foci, namely New Guinea, north western Australia, south-east India, the Caribbean, parts of South America, parts of central Africa [1] and the KwaZulu/Natal region of South Africa [2].

In 1905, Donovan first described the characteristic intracytoplasmic inclusion bodies in mononuclear cells which were present in smears from oral lesions of a patient who also had ulcerated lesions of the genitalia [3]. These were recognised as the aetiological agent and named Calymmatobacterium granulomatis [4]. Currently the laboratory diagnosis of granuloma inguinale relies on the observation of ‘Donovan bodies’ in tissue smears or biopsy specimens examined by Giemsa and Wright stains [5]. To increase the sensitivity and specificity of the diagnosis, Dieterlies, Warthin-Starry [6] and the rapid Giemsa (RapiDiff) [7] stains have been used.

No real progress has been made with regard to culture of the aetiological agent. In 1943, Anderson [8] reported growth of a gram-negative capsulate bacillus in the yolk of embryonated eggs from a tissue specimen rich in Donovan bodies. Thereafter, between 1943 and 1951, several reports on the primary isolation.
of the micro-organism in the yolk sac of embryonated eggs [9–12] and chick brain [13] appeared and a few of these cultures were reported to have adapted to growth on cell-free media such as the fresh yolk medium of Dienst et al. [12] and Locke-yolk medium of Dulaney et al. [14]. The fresh yolk medium of Dienst et al. [12] was modified by replacing the egg yolk with lactalbumin hydrolysate and Anderson’s originally passed strain was said to have been cultured in 1959 [15]. Sodium azide and brilliant green were added to this medium to make it selective and in 1962 Goldberg reported the isolation of an organism resembling C. granulomatis from the faeces of a patient with cervical granuloma inguinale [16].

Despite the reports of these early successes, there has been no subsequent publication on the growth or nature of the organism or its susceptibility to antimicrobial agents. Furthermore, no strains are available from any type culture collection worldwide. Recently, the successful growth of C. granulomatis in a monocyte co-culture system from biopsy specimens of three patients with ulcerative lesions was reported [17]. The present study reports on the modification of this co-culture technique and its application in the growth of the organism from tissue scrapings of genital lesions.

Materials and methods

Development of co-culture system

Over a 3-year period from Jan. 1993 to Dec. 1995, all female patients with ulcerative genital lesions presenting to the antenatal and gynaecology outpatient clinics of King Edward VIII Hospital, Durban, were investigated. Smears or tissue biopsy specimens, or both, were obtained from ulcerative lesions and investigated for chancroid, herpes genitalis, lymphogranuloma venerum, syphilis and granuloma inguinale by conventional laboratory tests. All ‘Donovan body’-positive biopsy specimens were processed for co-culture in monocytes.

To obtain peripheral blood mononuclear cells (PBMC), 30 ml of peripheral blood were obtained from a single normal healthy volunteer. This was mixed with an equal volume of Hank’s Balanced Salts Solution (HBSS; Whittaker M. A. Bioproducts, MD, USA) and passed through density gradient centrifugation with Histopaque 1077 (Sigma). The PBMC collected from the interface were rinsed in HBSS and suspended in RPMI 1640 (Whittaker M. A. Bioproducts) supplemented with 2 mM L-glutamine and unheated autologous serum 10%. The cells were transferred to tract vials containing glass coverslips (Bibby Sterilin) to allow monocyte adherence for 1 h at 37°C in CO₂ 5% in air. Each vial was then washed free of non-adherent cells with supplemented RPMI 1640 medium. The viability (99%) of PBMC was assessed with the trypan blue stain.

Biopsy specimens were suspended in supplemented RPMI 1640 and treated with amikacin (Bristol Laboratories) at a concentration of 10 mg/L for 2 h to eliminate contaminating bacteria, and then rinsed in PBS, pH 7.2, homogenised with a Dounce tissue grinder and resuspended in supplemented RPMI 1640. Each specimen (500 µl) was inoculated into three vials of monocytes. The vials were incubated for 1 h at 37°C in CO₂ 5% in air and 1 ml of fresh supplemented RPMI 1640 was added, followed by incubation for a further 48 h. Subcultures were made on to a fresh monocyte monolayer. In order to check for contaminating flora (Table 1) and to determine whether any cell-free media would support the growth of C. granulomatis (Table 2),

Table 1. Culture media and incubation conditions used to check monocyte cultures for the presence of contaminating flora

<table>
<thead>
<tr>
<th>Medium</th>
<th>Incubation temperature</th>
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<tbody>
<tr>
<td>Blood agar</td>
<td>25°C, 33°C, 37°C</td>
</tr>
<tr>
<td>MacConkey agar</td>
<td>37°C</td>
</tr>
<tr>
<td>Chocolate agar</td>
<td>25°C, 37°C</td>
</tr>
<tr>
<td>Sabouraud's agar</td>
<td>37°C</td>
</tr>
<tr>
<td>Brucella agar</td>
<td>25°C, 37°C</td>
</tr>
<tr>
<td>Blood agar (10%)</td>
<td>25°C, 37°C</td>
</tr>
<tr>
<td>Thioglycollate broth</td>
<td>25°C, 37°C</td>
</tr>
<tr>
<td>Mueller Hinton agar with isovitalex 1%</td>
<td>33°C, 37°C</td>
</tr>
<tr>
<td>Vancomycin 3 mg/ml, choloatised horse blood (5%)</td>
<td>33°C, 37°C</td>
</tr>
<tr>
<td>Wilkens Chalgren agar</td>
<td>37°C</td>
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</tbody>
</table>

Table 2. Cell-free culture media used for the attempted recovery of C. granulomatis

<table>
<thead>
<tr>
<th>Medium</th>
<th>Incubation temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dulaney's slants</td>
<td></td>
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<tr>
<td>-modified: 50% egg yolk in thioglycollate medium and thioglycollate overlay</td>
<td>25°C, 37°C</td>
</tr>
<tr>
<td>-modified: 50% egg yolk in Locke's solution and thioglycollate overlay</td>
<td>25°C, 37°C</td>
</tr>
<tr>
<td>-modified: 50% egg yolk in Locke's solution and normal human sera (1 in 10) and Locke's solution overlay</td>
<td>25°C, 37°C</td>
</tr>
<tr>
<td>Lactalbumin hydrolysate (LH)</td>
<td></td>
</tr>
<tr>
<td>-with sodium azide and with brilliant green</td>
<td>25°C, 37°C</td>
</tr>
<tr>
<td>-with activated charcoal</td>
<td>25°C, 37°C</td>
</tr>
<tr>
<td>-LH replaced with phytone peptone</td>
<td>25°C, 37°C</td>
</tr>
<tr>
<td>-LH replaced with yolk from 5–7-day embryo</td>
<td>25°C, 37°C</td>
</tr>
<tr>
<td>Embryonic yolk</td>
<td></td>
</tr>
<tr>
<td>-with macerated chick heart</td>
<td>25°C, 33°C, 37°C</td>
</tr>
<tr>
<td>Cystine agar slants</td>
<td></td>
</tr>
<tr>
<td></td>
<td>25°C, 33°C, 37°C</td>
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</tbody>
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various culture media were inoculated and incubated at appropriate temperatures and atmospheric requirements for a period of 7 days. The coverslip from each vial was air-dried, stained by Gram's and rapid Giemsa (Rapi-Diff) [7] stains and examined by light microscopy at ×1000 magnification. During the development of the culture system, a total of 22 biopsy specimens was subjected to this procedure for co-culture. Pure cultures of C. granulomatis were obtained from only three of these specimens and the remaining 19 cultures were overgrown with various micro-organisms, despite amikacin treatment. Twelve of the 19 cultures contained bacteria morphologically resembling C. granulomatis among the mixed cultures.

Transmission electron microscopy
Cells from the three positive pure co-cultures were scraped off the coverslips. These suspensions were centrifuged at 12 000 rpm for 15 min and the deposits were fixed in glutaraldehyde 1% in Eagle's minimal essential media with Earle's balanced salts solution (EMEM) (Whittaker M. A. Bioproducts) for 2 h. The specimens were thereafter washed in EMEM and post-fixed in osmium tetroxide 1% in 0.1 M cacodylate buffer for 1 h. The specimens were then dehydrated through ascending grades of alcohol and embedded in araldite epoxy resin. Sections (50 nm) were cut on a Reichert ultracut microtome and stained with Reynold's lead citrate and uranyl acetate before examination with a Joel 100C transmission electron microscope.

Modification of the co-culture method
The co-culture method was modified in three ways: the decontamination procedure for specimens, the source of PBMC and the use of fetal calf serum instead of autologous serum for culture. To establish which antibiotics could be used for specimen decontamination, all contaminating organisms were cultured and tested for their susceptibility to a variety of antimicrobial agents. Thereafter, subsequent biopsy specimens were subjected to a combination of vancomycin (Eli Lilly, USA) 5 mg/L, metronidazole (Rhone-Poulenc) 10 mg/L, and amikacin (Bristol Laboratories) 10 mg/L for 2 h and this combination was found to be most appropriate in eliminating bacterial contamination. As a regular supply of blood from laboratory volunteers is difficult to obtain, PBMC isolated from HIV-negative blood donors attending the Natal Blood Transfusion Services were used. Buffy coat fractions were mixed with pre-warmed unsupplemented RPMI 1640 and transported to the laboratory, and immediately subjected to density gradient centrifugation as described above. The autologous serum was replaced with heat-inactivated fetal calf serum (Delta Bioproducts, South Africa). This modified method was used for co-culturing additional biopsy specimens from female patients and tissue scrapings obtained from patients presenting with genital ulcer disease to the Sexually Transmitted Diseases (STD) Clinic, City Health, Durban.

Culture from tissue scrapings
In a study to determine the aetiology of genital ulcer disease amongst adult males, 200 consecutive patients with genital ulceration were investigated. Tissue scrapings were obtained with a 100 μl plastic bacteriological loop (Bibby Sterilin) and collected in 500 μl of PBS, pH 7.2. One hundred μl were transferred to 500 μl of decontamination medium containing RPMI 1640 supplemented with fetal calf serum 10% and amikacin 10 mg/L, vancomycin 5 mg/L and metronidazole 10 mg/L. After 2 h the tubes were centrifuged at low speed, excess supernate was discarded and the deposit was resuspended in 1 ml of RPMI 1640 supplemented with fetal calf serum 10% without antibiotics, before inoculation on to mononuclear cells.

Immunofluorescence
Slides of all positive co-cultures were incubated for 30 min at 37°C in a humidified chamber with sera from two patients with granuloma inguinale who presented with lesions of 3 months and 1 year duration. Serum from a volunteer with no past or present history of genital ulcer disease was also tested. All sera were tested at dilutions of 1 in 80 and 1 in 160. The slides were washed and re-incubated with fluorescein isothiocyanate-labelled mouse anti-human IgG (Wellcome Diagnostics) for 30 min. Smears obtained from the ulcerative lesions of patients with granuloma inguinale which showed characteristic ‘Donovan bodies’ were used as positive controls. Cultures of laboratory strains of Klebsiella pneumoniae, K. oxytoca, K. aerogenes, Enterobacter cloacae, Citrobacter freundii and Escherichia coli were used as negative controls. Both positive and negative control slides were tested with the patients’ sera and serum from the volunteer.

Results
During development of the co-culture system, pure cultures of C. granulomatis were obtained from three of 22 biopsy specimens from women with granuloma inguinale (Donovanosis). An additional biopsy specimen yielded pure growth after use of the modified method. Fourteen of the 200 tissue scrapings from men presenting with genital ulcers to the STD Clinic yielded growth in the monocyte co-culture after incubation for 48 h. None of these grew on any of the cell-free culture media (Table 1) that were incubated for 7 days.

All co-cultures, i.e., the four from biopsy specimens and 14 from tissue scrapings, tested positive in the indirect immunofluorescence test with the patients’ sera (Fig. 1A) and negative with the serum from the volunteer at both dilutions. Direct smears from lesions
with 'Donovan bodies' were also positive (Fig. 1B) with patients' sera and negative with serum from the volunteer, whilst smears of the control bacterial cultures were negative (Fig. 1C) with both sets of sera.

The co-cultures remained viable on subculture for at least eight passages. Micro-organisms with the typical morphology of the aetiological bacteria were visualised both intra- and extra-cellularly in co-cultures.
The extracellular bacteria were in close proximity to the monocytes. The staining reaction revealed characteristic single and bipolar condensation as described for 'Donovan bodies' (Fig. 2A). The bacteria were pleomorphic with either bulging or tapered ends. Dividing bacilli were evident. The size range was 0.5–2.0 \( \mu \text{m} \) in width and 1.0–2.5 \( \mu \text{m} \) in length. After subsequent passages an increase in the number of bacterial cells was observed with a number of coccobaccillary forms present (Fig. 2B). Around all the cells a translucent area was present, suggestive of capsule formation. The single and bipolar condensation was still evident. Gram's stain of the cultures showed that the bacteria were gram-negative (Fig. 2C), although they stained poorly.

On transmission electron microscopy, bacteria appeared singly or in clusters (Fig. 3A) with a number of extracellular dividing organisms (Fig. 3B). They exhibited a characteristic gram-negative cell wall (Fig. 3C), consisting of an outer membrane (OM), a middle electron opaque layer (MW) and an inner plasma membrane (IM). The periplasmic space (PS) between the outer and inner membrane was electron lucent. Both plasma and outer membranes were approximately 7.5 nm thick, each displaying the typical trilaminar nature. The capsule (G) was thick, dense and fuzzy (Fig. 3D). A membrane-like structure (TM) was observed on the periphery of the capsule. The cytoplasm was rich in ribosomes, many of them occurring in aggregates. Electron-dense granules (DG) were prominent (Fig. 3D). Pili (fimbriae) or flagella were not identifiable.

**Discussion**

The resurgence of granuloma inguinale in the KwaZulu/Natal region of South Africa prompted this attempt to culture the aetiological agent, *C. granulomatis*. Initial efforts to emulate the work of the early researchers (1940–1960) met with no success. This is also in keeping with the unavailability of the organism from any international type culture collection. However, when the yolk sacs of 5–7-day-old fertile chick eggs were used, the aetiological organism was maintained but could not be subcultured for growth.

The choice of PBMC for culture was based on the visualisation of the causative agent within mononuclear cells of specimens (smears and biopsy) of infected tissue. Such a system allows human monocytes and macrophages to control the growth of microbial pathogens and provides an opportunity to study host–parasite interactions [18]. During the initial phase of the study, PBMC were obtained from a

![Fig. 3. Electron microscopic characteristics of *C. granulomatis* from monocyte co-cultures. A, Large number of extracellular organisms. B, Dividing extracellular organisms. C, Gram-negative cell wall structure of the organism consisting of an outer membrane (OM), middle electron opaque layer (MW), inner plasma membrane (IM) with periplasmic space (PS) and evidence of a trilaminar membrane (TM) around the capsule. D, Characteristic thick, fuzzy and electron dense capsule (G) with numerous electron dense granules (DG) within the cytoplasm.](image-url)
single donor, a method which proved to be fairly restrictive. The availability of HIV-negative blood donor PBMC from the Natal Blood Transfusion services facilitated the regular supply of these cells. A natural extension of this would be the use of an immortalised monocyte cell line.

A predictable problem of growing organisms from genital site lesions is that of contamination with other less fastidious organisms which overgrow C. granulomatis and kill the PBMC as well. Initial attempts aimed to decontaminate these specimens by the use of an aminoglycoside only. These antibiotics do not enter eukaryotic cells and are, therefore, expected to kill all susceptible extracellular bacteria. This would include all extracellular C. granulomatis, because aminoglycosides have been used for the successful treatment of granuloma inguinale [1, 19]. However, it was hoped that there would be sufficient numbers of intracellular bacteria to maintain growth of the organism. This approach was successful and the growth of C. granulomatis from biopsy specimens of three women with granuloma inguinale was reported [17]. However, in view of the persisting contamination of biopsy specimens with aminoglycoside-resistant gram-positive cocci and anaerobic bacteria, metronidazole and vancomycin were added on the presumption that C. granulomatis is not an anaerobic organism. The combination of vancomycin and metronidazole together with amikacin was used for decontamination on additional biopsy specimens and on scrapings of genital ulcers. Successful culture of C. granulomatis was obtained in 14 of 200 men with genital ulcer disease from whom ulcer scrapings were collected.

Confirmation that organisms obtained in co-culture were C. granulomatis is based on three observations: the isolates did not grow on any of the extensive cell-free culture media listed in Table 2; they demonstrated a positive reaction in the immunofluorescence test with immune sera; and they showed striking morphological similarities with C. granulomatis by both light and transmission electron microscopy.

In direct smears the ‘Donovan bodies’ are seen intracytoplasmically within macrophages and are pathognomonic for granuloma inguinale [6]. The bacteria within these cells appear as either definite short coccobacilli or very pleomorphic capsulated forms. It is suggested that the pleomorphic forms are undergoing a process of bacterial elongation and active replication. The pleomorphic nature of the organism in culture has been described by several workers [9, 16, 20]. The bacteria in the monocyte co-culture after incubation for 48 h at 37°C in CO₂ 5% in air were extremely pleomorphic, varying in size and shape in comparison to those usually seen in direct smears, suggesting that these organisms were in a process of cell division and multiplication. The morphology of the bacteria varied after subsequent passages and was either pleomorphic or coccobacillary. During these subsequent passages the capsules and the single or bipolar staining reaction of the organisms were still evident. This morphological variation may result from the demonstration of different stages of growth by viable cultures in an optimal growth environment. Some authors have reported the growth of what they believed to be C. granulomatis on cell-free growth media [9, 14–16]. Although this study used all the media reported, no growth was obtained, despite following the published methodology for the preparation of the media. It is possible that unpublished details or unidentifiable differences in the ingredients may be the reason for these negative results. Another explanation could be that previously reported isolates of C. granulomatis which grew on cell-free media belonged to a different species.

Anderson et al. reported the presence of C. granulomatis specific antibodies in convalescent sera of patients with granuloma inguinale [21], whilst others have found similar responses when using Klebsiella antigen instead of Calymmatobacterium antigen [22, 23]. Although there are controversies regarding cross-reactivity with Klebsiella species in such a test, no reaction was observed with the control organisms, including the three species of Klebsiella. These results confirm the lack of cross-reactivity with other bacterial antigens. All the monocyte co-culture isolates and the direct smears containing ‘Donovan bodies’ used as positive controls showed bright fluorescence.

Previous reports of the electron microscopic features of C. granulomatis have been inconsistent. Davis et al. [24] described the presence of ‘bacteriophage’ attached to the cell wall and empty phage heads within the organism and it was suggested C. granulomatis is a phage modified bacterium. This phenomenon was confirmed by Davis in a later study [25], but was strongly refuted by Kuberski et al. [26]. In the present study there was no evidence of particles either within or on the micro-organism to suggest the presence of bacteriophage. Distinct electron dense granules were evident within the bacterial body and these would give the organism the characteristic single or bipolar ‘safety pin’ appearance.

The presence of the capsule has also been described and confirmed [24–26]. However, the description appears to be discrepant in relation to the present study. Both authors [24, 26] described the capsule as an electron lucent area around the organism, whereas the feature of the capsule in the present study was electron dense. The capsule characteristics have been described extensively for other bacterial species and invariably have an electron dense appearance [27]. One explanation could be that previous electron microscopic studies have been performed directly on clinical biopsy specimens, it is possible that these
electron lucent areas are in reality intracellular vacuoles in which the bacteria are situated and not bacterial capsules. In the present study, electron microscopic features are described from cultured specimens and not from biopsy specimens. Davis et al. [24] suggested that the capsule is unusual as it is large in size and has a capsular membrane. Whether the membrane-like structure observed on the periphery of the capsule in this study represents the capsular membrane seen in Davis' study is unclear. Clearly there was no evidence of pili (fimbriae) or flagella as suggested by Chandra et al. [28]. The bacterial cell wall was characteristically that of a gram-negative bacterium as described by others [24, 26].

This study developed a culture method for C. granulomatis which can be applied to routine clinical specimens such as ulcer scrapings, as well as biopsy samples of large lesions. The successful culture of the aetiological agent of granuloma inguinale will enable the collection of a sufficient number of bacterial strains to study the biological characteristics and virulence attributes of this micro-organism, to determine its antimicrobial susceptibility profile and to develop a definitive diagnostic test. This will lead to a better understanding of the epidemiology and pathogenesis of this disease.

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References