DRUG RESISTANCE AND *ACANTHAMOEBA* KERATITIS: THE QUEST FOR ALTERNATIVE ANTIPROTOZOAL CHEMOTHERAPY

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SUMMARY

Trophozoites and cysts of 20 isolates of Acanthamoeba from the cornea and five from related samples were tested in vitro for sensitivity to ten drugs (three aromatic diamidines, two aminoglycosides, two macrolides, a polyene macrolide antibiotic, an organoarsenical and an antimetabolite) and two cationic antiseptics (chlorhexidine and polyhexamethylene biguanide, PHMB). Only chlorhexidine and PHMB showed uniform amoebacidal activity. Aromatic diamidines (pentamidine isethionate, propamidine isethionate and diminazene aceturate) generally proved effective against both forms of the amoeba; only pentamidine gave synergy with the biguanide while propamidine gave an additive effect. Other drugs tested proved erratic or ineffective against different isolates. Chlorhexidine alone, or together with propamidine, was subsequently used in two patients with proven Acanthamoeba keratitis; the causative isolates were sensitive to the individual compounds and to the combination in vitro. The treatment provided resolution of the clinical disease; amoebae were shown to be nonviable by histology and culture. The combination of chlorhexidine and propamidine is recommended for treatment of proven Acanthamoeba keratitis.

Keratitis associated with *Acanthamoeba* infection is a relatively rare, sight-threatening condition occurring most often in contact lens wearers, where there has been inappropriate or inadequate disinfection of contact lens systems. The clinical presentation of the disease is often mistakenly diagnosed as herpes simplex or fungal infection. This results in inappropriate anti-microbial agents being administered. Early features such as photophobia and recurrent epithelial

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trate, dendritiform patterns and localised oedema, especially in a young person, as well as an association with contact lens wear, should suggest the possibility of *Acanthamoeba* keratitis. This can be confirmed by isolation and cultivation of the protozoan from corneal scrapes or biopsy material. Once a definitive diagnosis has been achieved, appropriate anti-acanthamoebal drug therapy will invariably be required.

The attests to a wide variety of drugs providing species or strains, both *in vivo* and *in vitro*. In clinical practice, however, it is important to note that some of the agents, for example some of the aromatic diamidines, may merely inhibit replication or induce encystment of the tropozoite form, rendering it quiescent as a cyst, and thus often resistant to conventional drug therapy. In such circumstances the cyst retains the potential to exacerbate the disease on discontinuation of the drugs or, if infected tissue is retained, following corneal transplantation.

In the United Kingdom, empirical combination therapy of propamidine, dibromopropamidine and neomycin has proved efficacious in some patients. ¹⁰ Drug resistance, ¹¹ as well as allergic or toxic reactions after prolonged therapy with propamidine, ¹² has limited the use of this combination, and has prevented its widespread acceptance in clinical practice.

Novel approaches to chemotherapy of *Acanthamoeba* keratitis continue to be forthcoming. For example, a formulation containing the cationic antiseptic polyhexamethylene biguanide at fairly low concentration, alone or in combination with propamidine and/or neomycin, has proved very effective against both trophozoites and cysts of *Acanthamoeba* derived from proven clinical cases of the infection.^{13,14} Drug therapy, however, may be complicated by a number of factors, mostly associated with failure to attend to the natural history and metabolism of the protozoan within the diseased cornea and also to the pharmacology of the selected drug(s) within this location.

The of this study was to suggest a more rational approach to chemotherapy of *Acanthamoeba* keratitis based on *in vitro* drug sensitivity studies on cultures of *Acanthamoeba* isolated from patients and contact lens accoutrements (details of which have been published pre-Representatives of selected drug classes and cationic antiseptics were used singly or, where considered appropriate, in combination with each other, in order to determine whether a potent acanthamoebacidal action could be identified *in vitro*, where the mechanism of action could be and the effect subsequently exploited *in vivo*. Illustrative case reports are used to highlight some important aspects.

MATERIALS AND METHODS

Acanthamoeba and Cultivation

Eighteen Acanthamoeba corneal isolates, including three from the same patient (TB) taken at different time intervals, were initially investigated and its storage case were also included. From another (MT) and from the contact lens storage case and one from the water supply at (the home was negative) were included.

On completion of testing these another patient presented with *Acanthamoeba* keratitis. Isolates from a corneal biopsy and scrape as well as a soft contact lens were examined. This afforded the opportunity of assessing the *in vitro* findings from the 18 samples in a clinical setting.

amoebae were maintained by routine passage on to 1.5% high clarity agar No. 1 (LAB M) made up in was spread with heat-killed *Klebsiella aerogenes* and moistened intermittently with amoebal saline prior to incubation in air at either 25 °C, 32 °C or 35 °C.

In order to obtain sufficient numbers of each of the 25 isolates for *in vitro* drug screening, the surface of each plate was flooded with amoebal saline and agitated in order to permit transfer of amoebae to sterile plastic 75 cm² flasks (Sterlin, CelCult) containing approximately 50–100 cm³ of a defined growth medium.¹⁶

After several transfers, the amoebae were incubated in this medium for approximately 72 at either 25 °C or 32 °C.

In order to determine the purity of Acanthameoba cultures, Giemsa staining was performed. Viability of each Acanthamoeba culture was assessed using a 0.2% trypan blue. Cyst populations were obtained by incubating trophozoite cultures for about 7 days at 25 °C or 32 °C. Purity and viability were determined retrospectively, as described contact and viability were used and viability were used drug sensitivity studies.

Drugs

Aqueous (100 μg/ml) of drug or disinfectant were prepared immediately prior to use and filter-sterilised using Gelman filters with 0.22 μm pore size. Compounds used for assessment of amoebacidal action are shown in Table I. These agents were selected action are consequence of literature reports of their efficacy, or that of related drugs or against Acanthamoeba, either in wivo.

Drug and Antiseptic Screening

Drug and antiseptic screening was performed using a series of sterile 96-well microtitre plates containing a standardised concentration of 2×10^4 organisms per $100~\mu l$ of medium per well. One hundred with of doubling dilutions of each compound ($100-0.8~\mu g/ml$) were produced vertically for each of the 12 compounds tested. Lids were secured, then the contents of plates mixed gently for 10 minutes on a plate rotator prior to incubation at either 25 °C or 32 °C.

Sensitivity of isolates was assessed after 48 hours of incubation, by recording either the lowest concentration of drug or antiseptic which resulted in complete lysis or degeneration of trophozoites and non-viability of resulting cysts (minimum trophozoite amoebacidal concentration, MTAC) or, for cysts, the lowest concentration of test compound that resulted in no excystment and trophozoite replication (minimum cysticidal concentration, MCC),¹³

Table I. Drugs and antiseptics used in this study, their classification and proposed antimicrobial mechanism of action

Agent	Class	Same of the same o
Chlorhexidine digluconate ¹ (chlor)*	Cationic antiseptic	Membrane function
Polyhexamethylene biguanide ^a (phmb)	Cationic antiseptic	Membrane function
Propamidine isethionate ³	Aromatic diamidine	DNA synthesis
Pentamidine isethionate ⁴ (penta)	Aromatic diamidine	DNA synthesis
Diminazine aceturate ^{b.5} (dim)	Aromatic diamidine	DNA synthesis
Neomycin sulphate ⁶ (neo)	Aminoglycoside antibiotic	Protein synthesis
Paromomycin sulphate ⁷ (paro)	Aminoglycoside antibiotic	Protein synthesis
Amphotericin B ⁸	Polyene macrolide antibiotic	Membrane (ergosterol) biosynthesis
(dir.)	Macrolide antibiotic	Protein synthesis
Spiramycin ^{d,10} (spir)	Macrolide antibiotic	Protein synthesis
Cymelarsan ^{b,11} (cymel)	Organoarsenical	Energy metabolism
α -Difluoromethylornithine ^{b 12} (dfmo)	Antimetabolite	Substrate-enzyme reaction (inhibitor of ornithine decarboxylase)

Gift from: aMoorfields Eye Hospital; bProf. F. W. Jennings; Lilly Research Centre Ltd; dRhone-Poulenc Ltd.

Supplied as solutions: 10.05% Sterets, Unisept; 20.02% contains hypromellose eye drops, 0.3%; % Brolene, May & Baker (contains benzalkonfum chloride); 6Minims, Smith & Nephew (contains phenylmercuric nitrate).

Supplied as solids: ⁴Pentacarinat, May & Baker; ⁷Sigma; ⁸Fungizone i/v, Squibb: ¹² donated compounds.

*Abbreviations in parentheses are used in Figs. 1 and 2 and Table II.

after thoroughly cystaspec of residual drug, and in the medium described above. 16 Observations were performed in duplicate using an inverted microscope.

For *in vitro* combination testing, a chequerboard method was used.¹⁷ With this procedure, four possible outcomes of drug–drug or drug–antiseptic combinations were possible:¹⁸

- 1. Additivity, where the result with the two compounds was equivalent to their sum when used separately.
- 2. Autonomy (or indifference), where the result with the two compounds was not different from the result with the more effective compound used alone.
- 3. Antagonism, where the with the two compounds was less than the additive response.
- 4. Synergism, where the result with the two compounds was greater than the additive response.

Five replicates per determination were performed¹⁹ using the combinations shown in Table II; findings were based on results obtained from 7 of the 18 corneal isolates.

CASE REPORTS

Patient TB

Drug resistance occurred after treatment of *Acantha-moeba* keratitis with propamidine alone; topical neomycin had been withdrawn because of allergy. The isolate collected in early treatment was retested for drug and antiseptic sensitivities, as were two later isolates. Both the latter have been recorded before as temperature-sensitive and propamidine-resistant.

Patient AT

Treatment was given with topical betamethasone—neomycin q.i.d. for 4 weeks, followed by propamidine q.i.d. alone for 3 weeks, prior to laboratory confirmation of *Acanthamoeba* keratitis by corneal scrape and biopsy. The isolated protozoan grew poorly at 35 °C, compared with those from contact lens and storage case. AT had worn Acuvue (Johnson & Johnson) disposable soft contact lenses (FDA Group 4) and had used Softab (Alcon; a chlorine-based system) following the manufacturer's instructions for contact lens disinfection.²⁰

AT responded satisfactorily over a 12 month period to a combination of propamidine and neomycin, before undergoing a penetrating keratoplasty due to scarring in the visual axis. Histology revealed a few degenerate cysts. There was no recurrence of infection after 1 year.

Patient MT

Treatment of this patient's *Acanthamoeba* keratitis was initially with a topical propamidine–neomycin combination, which induced a toxic reaction in the cornea within 2 months. At this stage trophozoites were still present in a corneal biopsy.

Guttae chlorhexidine (0.02% w/v in 0.9% saline) alone

was administered for 9 months a further adverse reaction. With this course of treatment relapse of *Acantha-moeba* infection was not apparent. Isolates from a corneal biopsy, contact lens and its storage case grew confluently at 35 °C. MT had worn Acuvue disposable soft contact lenses (FDA Group 4) and used Softab for contact lens disinfection.²⁰

AB

A 48-year-old man with mild atopy and early keratoconus had recently changed contact lenses from daily-wear soft to rigid, gas-permeable lenses. Total one sterilising solution containing dividingen was used for lens hygiene. The patient attended an eye casualty department with a dendritiform corneal ulcer, stromal oedema and a mild anterior chamber reaction. This was considered to be herpetic kerato-uveitis and treatment was commenced with acyclovir ointment and prednisolone 0.5% drops. He was then referred to one of us (C.M.K.). Intensive antiviral and steroid therapy did not bring about improvement. The eye became increasingly painful, injected, and a ring abscess developed. A corneal biopsy was performed 2 months after initial presentation. Acanthamoeba was observed histologically in the corneal stroma and cultured from it. The protozoan was seen on the hydrogel contact lens and cultured from washings.

Treatment was commenced with 0.02% chlorhexidine in isotonic saline drops every hour for a week then 2-hourly, combined with 0.1% propamidine at the same frequency, and 3-hourly prednisolone 1%. The eye became and comfortable within 7 days though the patient remained. The abscess began regressing and the epithelium slowly healed. At 2 months there was a central stromal opacity and 3 mm overlying epithelial defect.

The patient next presented as an emergency with a 2-day history of sudden discomfort and further loss of vision. Cornea was found to be perforated centrally with an intumescent lens. An emergency keratoplasty, extracapsular lens extraction and posterior chamber lens

Table II. In vitro combination testing of selected drugs against seven Acanthamoeba corneal isolates

Combination	Effect on MTAC	Effect on MCC		
Isolates (TB 1-2; 1, 2, 10, 13; AB)				
phmb + pentamidine	Synergy (slight)	Synergy (slight)		
phmb + neomycin	Additivity	Additivity		
phmb + dirithromycin	Autonomy	Autonomy		
propamidine + neomycin	Additivity	Additivity		
pentamidine + neomycin	Additivity	Additivity		
*diminazine + neomycin	Additivity	Additivity		
*diminiazine + dirithromycin	Autonomy	Autonomy		
*pentamidine + dfmo	Autonomy	Autonomy		
*cymelarsan + dfmo	Autonomy	Autonomy		
*neomycin + dirithromycin	Autonomy	Antagonism		
Isolate AB				
chlorhexidine + propamidine	Additivity	Additivity		
chlorhexidine + pentamidine	Additivity	Synergy (slight)		
chlorhexidine + neomycin	Additivity	Additivity		

MTAC, minimum trophozoite amoebacidal concentration; MCC, minimum cysticidal_concentration. *Not including

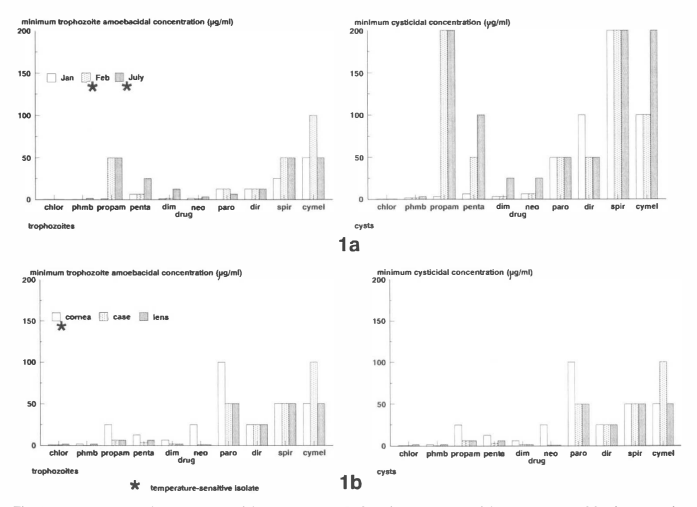


Fig. 1. (a) Minimum trophozoite amoebacidal concentration (MTAC) and minimum cysticidal concentration (MCC) of ten agents for three corneal isolates from patient TB. (b) MTAC and MCC of ten agents for corneal, storage case and contact lens isolates from patient AT. For abbreviations see Table 1.

implant was performed. Half the excised cornea was sent for culture and half for histopathology. *Streptococcus acidominimus* (weakly α-haemolytic) was cultured from the corneal tissue but it remained persistently culturenegative for *Acanthamoeba*. Degenerate cysts were seen in the corneal stroma but no trophozoites. The corneal epithelium appeared healthy.

Post-operatively treatment was continued with Brolene (May & Baker) and chlorhexidine with prednisolone (1% 2-hourly) and Polytrim (Burroughs Wellcome) drops 2-hourly. Three months after keratoplasty the graft remained clear with no signs of further infection. The corrected vision was 6/12.

RESULTS

Patient TB (Fig. 1A)

Trophozoites and cysts of all three corneal isolates were highly sensitive to chlorhexidine and polyhexamethylene biguanide (PHMB). While trophozoites and cysts from the first isolate were sensitive to propamidine, the two subsequent isolates showed resistance as previously tested, a phenomenon apparently related to temperature¹¹ but not to the action of the antiseptics. Both forms of the amoeba for all isolates were sensitive to neomycin. Amphotericin B,

cymelarsan and α -difluoromethylornithine (α -DFMO) were ineffective against trophozoites and cysts for all three isolates. This patient, in whom medical treatment failed with propamidine and arsenicals, ¹¹ would probably have benefited from therapy with cationic antiseptics.

Patient AT (Fig. 1B)

Trophozoites and cysts of all three isolates (cornea, contact lens and storage case) were highly sensitive to chlorhexidine and PHMB. Trophozoites and cysts from the corneal isolate, but not the others, were less sensitive to propamidine and were temperature-sensitive (confluent growth at 25 °C, poor at 35 °C); this trend was also evident for the other two diamidines. Neomycin was more effective than paromomycin but the corneal isolate was more resistant than the others as regards both trophozoites and cysts. Amphotericin B, cymelarsan and $\alpha\text{-DFMO}$ were ineffective.

Patient MT (Fig. 2A)

Trophozoites and cysts of all three isolates (cornea, storage case, workplace water sample) were highly sensitive to chlorhexidine and PHMB and grew well at 32 °C. Propamidine was more effective than either pentamidine or

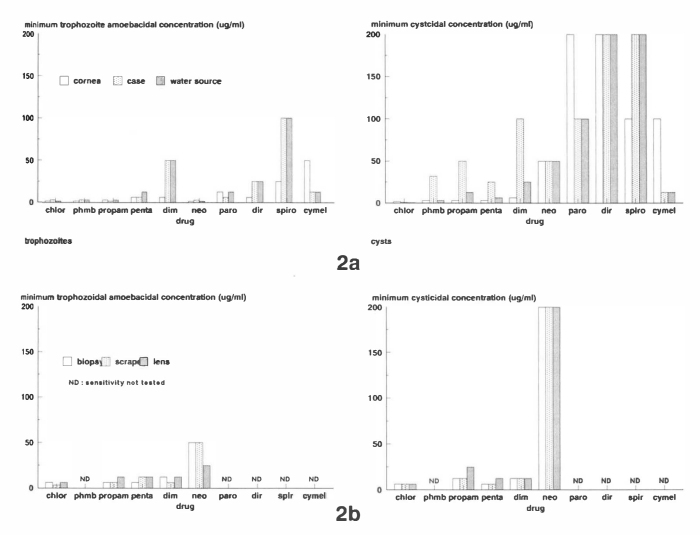


Fig. 2. (a) MTAC and MCC of ten agents for corneal, contact lens storage case and water-source isolates from patient MT. (b) MTAC and MCC of ten agents for corneal biopsy, corneal scrape and contact lens washing isolates from patient AB. For abbreviations see Table I.

diminazine against trophozoites of all three isolates, but pentamidine was most effective against cysts. For all three isolates, the storage case isolate was less sensitive to diamidines than either the corneal or water sample isolates. Trophozoites of all three isolates were sensitive to neomycin but cysts were resistant. Amphotericin B, cymelarsen and α -DFMO were ineffective.

Patient AB (Fig. 2B)

Trophozoites and cysts of all three isolates (corneal biopsy, scrape and contact lens washings) were sensitive at the upper limit to chlorhexidine; the biopsy sample only was tested against PHMB and gave similar values. All three isolates were sensitive to diamidines but again at the upper limit (MTAC 6.3–25 $\mu g/ml$, MCC 6.3–25 $\mu g/ml$). Trophozoites and cysts of the three isolates were resistant to acyclovir – which has been tested since it has been suggested that other antiviral agents maybe ineffective against Acanthamoeba.

Fig. 3A–G illustrates the findings for trophozoites and cysts obtained for the remaining 13 corneal isolates. As

described above, chlorhexidine and PHMB were most efficacious. All isolates with the exception of nos. 2 and 10 for trophozoites and nos. 1 and 13 for cysts were sensitive to propamidine; there were no obvious trends with the other two diamidines. Aminoglycosides were relatively ineffective against cysts while the trophozoites of four isolates (nos. 1, 4, 7, 10) were insensitive also. Trophozoites and cysts from all isolates were insensitive to macrolides, amphotericin B, cymelarsan and α -DFMO.

Fig. 3H shows the average values for MTAC and MCC for 10 of the 12 drugs tested. Chlorhexidine and PHMB were the most active compounds against both trophozoites and cysts. The diamidines were the next drug class in order of acanthamoebacidal activity. Neither the aminoglycosides, macrolides nor the arsenical were effective against cysts and showed an increasing inability to destroy trophozoites.

Table II gives findings obtained from *in vitro* combination of selected drugs and PHMB against six corneal isolates (i.e. TB 1, 2; 1, 2, 10, 13) and selected drugs and chlorhexidine against isolate AB. The only combinations that gave a slight synergistic response were the cationic antiseptics (chlorhexidine or PHMB) and pentamidine;

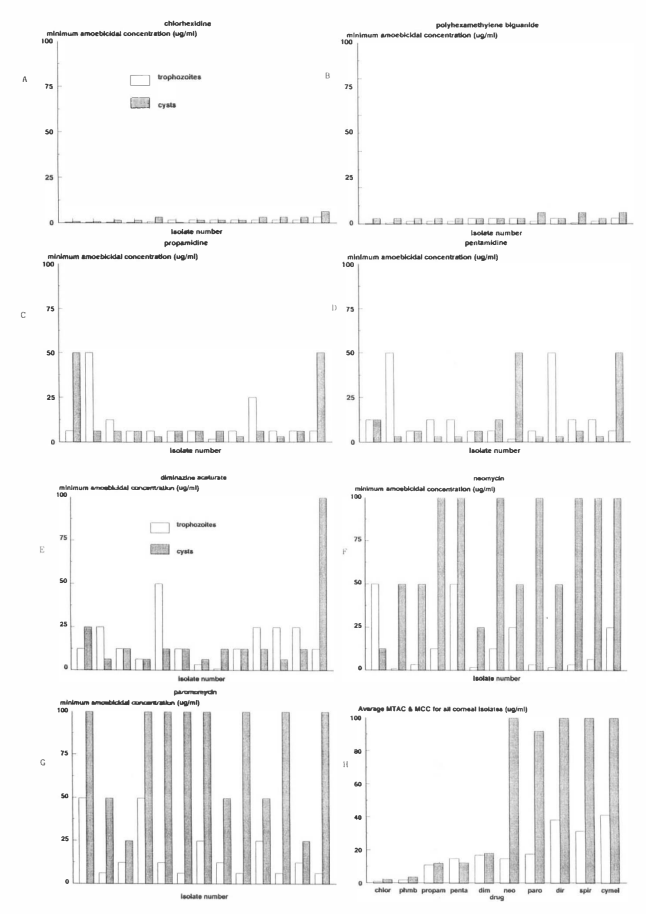


Fig. 3. MTAC and MCC for individual drugs: A, chlorhexidine; B, PHMB; C, propamidine; D, pentamidine; E, diminazene, F, neomycin; G, paramomycin; H, average MTAC and MCC for all corneal isolates.

additivity was found between chlorhexidine and propamidine for the AB isolate, a successful combination in practice. All three diamidines showed additivity with neomycin as did the combination of PHMB and neomycin. Other combinations showed autonomy, except that of neomycin and dirithromycin, where there was antagonism.

DISCUSSION

Failure of drug therapy in Acanthamoeba keratitis has been recognised for some time but the reasons are not always understood in the absence of drug sensitivity testing. This study suggests that some of the commonly used drugs such as neomycin and paromomycin are not effective amoebacides – a finding in accordance with that of other workers. 13 source case reports (TB, AT) suggest that resistance may as a result of low-dose, single-drug anti-amoebic therapy. Sensitivity testing can therefore be useful on all occasions. It is not sufficient to rely on of the organism from the contact lens or its storage case since amoebae from these sources may have different sensitivities from that isolated from the cornea, particularly when the diagnosis has been late and the has been pretreatment with a variety of drugs. Since an animal model is not yet available for the sensitivity testing of Acanthamoeba isolates, in vitro assessment is a necessary but relatively unsatisfactory

Agents tested in vitro and found to have an effect on different species and strains of Acanthamoeba include: clotrimazole²¹ and ketoconazole,²² although like other azoles the effect is to be amoebastatic rather than amoebacidal,²³ and with some drugs in this group the organisms may be highly resistant;²⁴ 5-fluorocytosine (a nucleotide although this drug has been found ineffective by other workers;^{27,28} the diamidines pentamidine isethionate, ^{10,29} although others have reported insensitivity to this drug, ²² hydroxystilbamidine isethionate, ^{25,30} diminizene aceturate¹⁰ and propamidine isethionate, 10,29 although again this drug has been identified as insensitive in other studies, except if combined with dimethylsulphoxide (DMSO);31 membrane-active peptides, the magainins,³² The effect being enhanced when in combination with silver nitrate or propamidine; pimaricin;²⁸ amphotericin B or AB methyl ester,²² although found to be ineffective in other studies;²⁷ certain inhibitors of biosynthesis;³³ trifluoroperazine;³⁴ the aminoglycosides paramomycin, ^{24,30} which was ineffective in other studies, ²⁵ and neomycin, 30 also without effect in other studies except if propamidine; ³¹ polymyxin E (colistin); ²² acriflavine³⁰ (although some workers have reported resistance,²⁸) and other acridines.³⁵

There is considerable disparity regarding *in vitro* efficacy of drugs which are active against *Acanthamoeba*; none demonstrate uniform activity against all isolates, and there is differential sensitivity between trophozoites and cysts, the former being more sensitive than the latter. Furthermore, some reports are based on mixed trophozoite

and cyst drug-sensitivity studies which may cause confusion with drugs that merely induce encystment and are not acanthamoebacidal in action.

Several compounds have been used with varying effect in the clinical setting. These include: itraconazole plus miconazole;³⁶ clotrimazole;³⁷ ketoconazole;³⁸ dibromopropamidine plus propamidine and neomycin; 10 propamidine isethionate as Brolene;³⁹ propamidine combination with neomycin-polymyxin B-gramidicin as Neosporin (Calmic);⁴⁰ Neosporin with or without miconazole or ketoconazole;41 pimaricin plus Neodecadron (dexphosphate, neomycin amethasone sulphate) hydroxyuracil, rifampicin and atropine;²⁸ PHMB solution contained 0.3% hypromellose, 0.45% NaCl, 0.37% KCl, 0.19% borax and 0.19% boric acid) alone or in combination with propamidine, 13 or PHMB in assessment tion of teams tears combined with propamidine and neomycin.¹⁴

As with in vitro testing, clinical reports suggest that drug selection has been relatively haphazard. As a result, it was decided to investigate a cohort of a corneal isolates from cases of three from contact-lens-associated materials and one from a water supply used to clean the storage case. Most of the drugs and antiseptics used in the present study have been previously assessed for potential anti-acanthamoebal activity. Two macrolides, however, were included (dirithromycin and spiramycin), since erythromycin is known to induce encystment of Acantha*moeba*. 42 The organoarsenical cymelarsen was included as a novel representative of this group with less inherent toxicity than earlier arsenical compounds which, in exhibit relatively poor activity against Acanthamoeba. 11.25 a cationic antiseptic, was selected for inclusion since it has been shown to have good antiacanthamoebal activity in vitro⁴³ and had previously been suggested anecodotally for therapy one of us (D.V.S.). PHMB, a related but as vet unlicenced for topical use in shown to have considerable activity against Acanthamoeba, in vitro and in vivo.

The cationic antiseptics showed outstanding efficacy against all isolates, with chlorhexidine giving the lowest MTAC and MCC 3H). The aromatic diamidines as a group were second in order of efficacy, although in keeping with previous literature reports the effect varied considerably between the isolates. A diminazine (an encystment-enhancing agent),8 showed satisfactory activity against some isolates, a finding in keeping with that of et al. 10 but at odds with the results of other workers. 30 The aminoglycosides again showed variability amongst the strains, and had no effect against cysts, both findings confirming previous literature reports. 13 Macrolides showed similar behaviour to aminoglycosides but were less effective. A note of caution should be introduced, however, since the macrolides used were prodrugs, the metabolites being more effective, at least against bacteria.44 The seasons showed poor activity against both forms of the protozoan. Others have reported similar resistance to arsenicals. 11.25 In keeping with the findings of

Table III. In vitro combination testing of drugs against Acanthamoeba corneal biopsy isolate from patient AB

Combination	Effect on mean trophozoite amoebacidal concentration	Effect on mean concentration
phmb + pentamidine	Synergy (slight)	Synergy (slight)
phmb + neomycin	Additivity	Additivity
phmb + propamidine	Additivity	Additivity
propamidine + neomycin	Additivity	Additivity
pentamidine + neomycin	Additivity	Additivity
+ propamidine	Additivity	Additivity
chlorhexidine + pentamidine	Additivity	Synergy (slight)
chlorhexidine + neomycin	Additivity	Additivity

other workers. 45 the inhibitor of ornithine

α-DFMO, had no is, however, reason to believe that other components of polyamine metabolism in *Acanthamoeba* may yet be found suitable as the basis for development of more active chemotherapy against the protozoan. 46,47

Single drug therapy of *Acanthamoeba* keratitis with currently used compounds appears inadequate, and may lead to emergence of drug resistance. Combination therapy must be considered. Notable in this context *in vivo* is the combination of and propamidine; and propamidine, or neomycin and propamidine; and propamidine, or neomycin, dibromopropamidine and propamidine; and, *in vitro*, DMSO and propamidine isethionate. In the present *in vitro* study, additive effects were observed with cationic antiseptics propamidine or neomycin, and slight synergy between the antiseptics and pentamidine (Tables II, III).

Following the demonstration of in vitro efficacy of chlorhexidine two patients have been treated with this drug. One patient (MT), who had developed an idiosyncratic reaction to both neomycin and propamidine, was treated satisfactorily with monotherapy chlorhexidine. The other patient (AB) received combination therapy of chlorhexidine with propamidine with rapid control of the Acanthamoeba infection. sis evident, however, that useful anti-acanthamoebic drugs may not have universal activity against all amoebae. We believe that, in general, combination therapy should always be employed, firstly because of the possibility of an additive effect and secondly to prevent the emergence of resistance. On the basis of these two patients, plus one other now treated successfully for 3 months with a similar combination to AB, and on anecdotal evidence from several personal communications, chlorhexidine seems to be well tolerated in the eve.

The findings from the present study are suggestive of membrane effects, which permit easier access of drug into the amoebae. Cationic antiseptics such as chlorhexidine⁴⁸ and, to a lesser extent, neomycin perturb the plasmalemma; this may facilitate entry of an effective drug as an aromatic diamidine. Diamidines are either inhibitors of *S*-adenosylmethionine decarboxylase in *Acanthamoeba*,⁴⁹ or drugs which interact with the nucleic acid of the organism. In addition they may act

in a way such as occurs in human neutrophilic granulocytes by for co-factors⁵² or cytoplasmic enzymes,⁵³ or may on their own exert an inhibitory effect on multiplication of *Acanthamoeba*. Analogues of the diamidine series⁵⁴ may considerably enhance this effect.

The combination of chlorhexidine and propamidine seems to have had effective amoebacidal action within the cornea. This could shorten the time during which anti-acanthamoebic drugs are required and their frequency of This, in turn, may reduce the likelihood of toxic reaction, 12 and obviate effects of inherent 29 or acquired 11 resistance to the diamidine.

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Key words: Acanthamoeba keratitis, Antiprotozoal Exercises Chlorhexidine, Diamidines, Drug resistance.

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