Complementary use of aesculin hydrolysis and inositol fermentation in the characterisation of Klebsielleae

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SUMMARY Rates of aesculin hydrolysis and inositol fermentation adequately differentiate Klebsielleae from other Enterobacteriaceae. In combination with tests for motility and growth in potassium cyanide medium, presumptive differentiation between Klebsielleae, and confirmation of Klebsiella pneumoniae, is obtained.

Aesculin hydrolysis and inositol fermentation have frequently been incorporated with success in biochemical testing schemes for differentiation between clinical isolates of Enterobacteriaceae.

Wasilauskus (1971) demonstrated that Klebsielleae could be readily differentiated from other Enterobacteriaceae by virtue of hydrolysis of aesculin to 6,7 dihydroxy coumarin, the reaction product combining with iron to form a black compound.

Inositol fermentation has also been widely used in differentiation of the Enterobacteriaceae, and inositol has been incorporated in specific screening media (Donovan, 1966; Barr and Mahood, 1976) and primary isolation medium (Davis and Matsen, 1974) for Klebsielleae. Barr and Mahood (1976), however, showed that some clinical isolates of Klebsiella spp, particularly Klebsiella pneumoniae and K. ozaenae, fermented inositol weakly and slowly, and demonstrated that fermentation by other Klebsiella spp could also appear to be weak or absent due to a rapid conversion of acid to neutral metabolites.

Since both aesculin hydrolysis and inositol fermentation have been utilised in parallel in some schemes for differentiation of Enterobacteriaceae (Chadwick et al., 1974), this investigation was carried out to elucidate and compare the ability of Klebsielleae and other Enterobacteriaceae to metabolise these two substrates. The results obtained were utilised to assess the value of the complementary use of aesculin hydrolysis and inositol fermentation, in an inositol-hydrogen sulphide-motility medium (Barr and Mahood, 1976), in the identification of Klebsielleae.

The value of these observations in the identification of Enterobacteriaceae by a replica agar plating method, modified from that described by Chadwick et al. (1974), is discussed.

Material and methods

ORGANISMS

Clinical isolates A total of 710 clinical isolates of Gram-negative bacilli examined comprised 434 cultures derived from urinary tract infections and 275 cultures isolated in the routine bacteriological laboratory from other sources. Of these, 234 cultures were identified as Klebsielleae by the methods of Chadwick et al. (1974) and Cowan (1974). An additional 20 clinical isolates of Klebsiella spp were obtained from Ruchill Hospital, Glasgow.

Stock cultures The following 26 cultures were also examined: Klebsiella aerogenes NCTC 8172; K. atlantae NCTC 9496; K. edwardsii NCTC 5054; K. pneumoniae (4) (NCTC 9633, 204, 8632, and 10246); K. ozaenae (4) (NCTC 5050, 9659, 10313, and 8883); K. rhinoscleromatis (2) (NCTC 5046 and 5047); Enterobacter aerogenes NCTC 10006; Enterobacter cloacae NCTC 10005; Serratia marcescens NCTC 9940; Escherichia coli NCTC 10418; Proteus mirabilis NCTC 5887; P. morganii NCTC 235; P. vulgaris NCTC 4175; P. rettgeri NCTC 7475; Providencia B, subgroup B, NCTC 10318; Providencia spp NCTC 10286; Citrobacter freundii NCTC 45

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SCREENING MEDIA PREPARATION

Seven basic screening media were employed. Aesculin bile agar (Difco), Simmon’s citrate agar (Difco), Phenylalanine agar (Difco), Xylose-lysine-deoxycholate agar (Difco), and Ornithine decarboxylase agar, derived from the fluid decarboxylase medium of Fay and Barry (1972), were utilised as described by Chadwick and Barry (1974). Two further screening media employed were inositol-hydrogen sulphide-motility agar medium (Barr and Mahood, 1976) and a potassium cyanide (KCN) agar medium based on that described by Braun (1938).

The inositol-hydrogen sulphide-motility medium contained (g/l): Tryptone (Oxoid), 10-0; NaCl, 5-0; Agar (Oxoid), 3-0; 2-3,5, triphenyl-tetrazolium chloride, 0-05; Fe₂NH₄PO₄, 0-3; Na₂SO₄.5H₂O, 0-2; K₃HPO₄, 0-3; acetoin, 8-5; and inositol, 20-0. Bromothymol blue in aqueous solution was added to give a final concentration of 0-003%. The medium was sterilised by autoclaving at 115°C for 10 minutes, acetoin and 2,3,5, triphenyltetrazolium chloride were added as filter sterilised aqueous solutions, and the pH value was adjusted to 6-6.

Potassium cyanide agar medium was prepared from KCN broth medium (Braun, 1938; Möller, 1954), described by Cowan (1974), and supplemented with 0.5% agar (Oxoid). Potassium cyanide was added from a 0.5% aqueous solution to give a final concentration in agar of 0.125 g/l (1 part in 8000). Agar was cooled to 45°C before addition of KCN, dispensed immediately, and allowed to solidify at 4°C. Medium was freshly prepared and distributed on each day of use, and sensitivity was checked at frequent intervals with appropriate stock cultures.

Sterilin replidishes were used throughout, 3 ml of each medium except inositol-hydrogen sulphide-motility medium (4 ml) being dispensed in each compartment.

INOCULATION AND INCUBATION

All media were inoculated from a four-hour peptone water culture. In specific investigations sequential 10-fold dilutions of inocula were prepared in 9 ml peptone water. Motility medium was inoculated with a straight wire and other media with one drop from a Pasteur pipette. The peptone water culture used for inoculation of other media was reincubated for a further 16 hours at 37°C.

Ornithine decarboxylase agar was covered with liquid paraffin before incubation and all plates were incubated for 16 hours at 37°C.

INTERPRETATION OF BIOCHEMICAL REACTIONS

Utilisation of inositol, aesculin, and ornithine was detected by medium colour change, and citrate utilisation by a colour change and/or visible growth. Resistance to KCN was recorded by growth, and colony colour on xylose-lysine-deoxycholate agar was noted. Production of indole in peptone water was tested with Kovac’s reagent, and phenylalanine deaminase activity was detected by the addition of acidified 10% ferric chloride solution. Hydrogen sulphide production and motility were recorded after 16 hours, and again after a further 24-hour incubation period.

A semiquantitative determination of aesculin hydrolysis and inositol fermentation was recorded.

Aesculin hydrolysis was recorded after 4 hours and 16 hours as – (no hydrolysis), + + (distinct colour change usually limited to inoculation surface), and + + + (complete colour change within an individual inoculation compartment). Inositol fermentation was recorded as – (no fermentation), + (indicator colour change along line of stab inoculation), + + (clean zone of indicator colour change), and + + + (complete indicator change within individual inoculation compartment).

IDENTIFICATION OF CLINICAL ISOLATES

Clinical isolates were identified as far as possible using the basic biochemical differentiation scheme described by Chadwick et al. (1974). Where further tests were necessary these were carried out by conventional methods (Cowan, 1974).

Identification of Klebsiella isolates was confirmed by further biochemical tests by conventional methods, viz, methyl red and Voges-Proskauer fermentation of dulcitol, inositol, and glucose (gas); malonate and gluconate utilisation; and urease activity.

Results and discussion

The hydrolysis of aesculin and fermentation of inositol by 730 clinical isolates and 26 stock cultures of Gram-negative bacilli are summarised in Table I. National Collection of Type Culture strains examined gave reactions typical of the majority of clinical isolates of the same species.

HYDROLYSIS OF AESCULIN

Among the Gram-negative bacilli investigated, the genera Klebsiella, Enterobacter, and Serratia were readily distinguished from other genera by rapid hydrolysis of aesculin. Sixty-three per cent of Klebsiellaeae showed visible hydrolysis, by formation of the black reaction product after four hours.
Complementary use of aesculin hydrolysis and inositol fermentation in the characterisation of Klebsielleae

Table 1  Hydrolysis of aesculin and fermentation of inositol by 756 isolates of Gram-negative bacilli

<table>
<thead>
<tr>
<th>Total number of isolates</th>
<th>Hydrolysis of aesculin&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Fermentation of inositol&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>4 Hours</td>
<td>16 Hours</td>
</tr>
<tr>
<td></td>
<td>--- + + + +</td>
<td>--- + + + +</td>
</tr>
<tr>
<td>Klebsiella aerogenes/oxytoca</td>
<td>202 60 142 0</td>
<td>0 0 202 17 18 176</td>
</tr>
<tr>
<td>atlantae/edwardsii</td>
<td>K. pneumoniae</td>
<td></td>
</tr>
<tr>
<td></td>
<td>15 11 4 0</td>
<td>0 0 15 6 8 1</td>
</tr>
<tr>
<td>K. ozoaena</td>
<td>12 7 5 0</td>
<td>0 1 11 2 4 6</td>
</tr>
<tr>
<td>K. rhinoscleromatis</td>
<td>4 0 4 0</td>
<td>0 0 4 0 2 2</td>
</tr>
<tr>
<td>Enterobacter aerogenes</td>
<td>16 10 6 0</td>
<td>0 3 13 0 2 13</td>
</tr>
<tr>
<td>Enterobacter/Serratia&lt;sup&gt;a&lt;/sup&gt;</td>
<td>33 20 13 0</td>
<td>0 4 29 17 8 7</td>
</tr>
<tr>
<td>Proteus mirabilis</td>
<td>56 56 0 0</td>
<td>56 0 56 0 0 0</td>
</tr>
<tr>
<td>P. vulgaris</td>
<td>5 5 0 0</td>
<td>2 3 0 5 0 0</td>
</tr>
<tr>
<td>P. morganii</td>
<td>11 11 0 0</td>
<td>10 1 0 8 3 0</td>
</tr>
<tr>
<td>P. rettgeri</td>
<td>20 20 0 0</td>
<td>15 3 2 4 2 3</td>
</tr>
<tr>
<td>Escherichia coli</td>
<td>316 316 0 0</td>
<td>302 10 4 300 15 1</td>
</tr>
<tr>
<td>Citrobacter spp.</td>
<td>44 44 0 0</td>
<td>41 3 0 41 2 1</td>
</tr>
<tr>
<td>Providencie spp.</td>
<td>3 3 0 0</td>
<td>3 0 3 2 0 1</td>
</tr>
<tr>
<td>Pseudomonas aeruginosa</td>
<td>22 22 0 0</td>
<td>19 3 22 0 0</td>
</tr>
</tbody>
</table>

<sup>a</sup>Hydrolysis of aesculin was recorded as (-) no hydrolysis, (+) clear zone of colour change due to hydrolysis, (+++) complete change within inoculation compartment.

<sup>b</sup>Fermentation of inositol was recorded as (-) no fermentation, (+) indicator change along line of stab inoculation, (++) clear zone of indicator colour change, (+++) complete indicator change within individual inoculation compartment.

<sup>a</sup>Isolates include Enterobacter spp. and Serratia spp. other than Enterobacter aerogenes.

NB Apart from Klebsielleae only five isolates, all Proteus rettgeri, gave a positive reaction on both substrates.

incubation, and aesculin hydrolysis of all Klebsielleae, excepting one, resulted in a complete colour change within the inoculation compartment after 16 hours' incubation.

No isolates of non-Klebsielleae exhibited any hydrolysis of the substrate after four hours' incubation. Although 27 (5-6%) of non-Klebsielleae isolates screened showed detectable hydrolysis of aesculin after 16 hours, hydrolysis by only six isolates resulted in a complete colour change within the inoculation compartment. Indeed, with the exception of four isolates of E. coli and two isolates of P. rettgeri, aesculin hydrolysis by non-Klebsielleae was restricted to the vicinity of bacterial growth on the surface of inoculation.

Inoculation with dilutions, prepared from the original peptone water, demonstrated that reduction of inocula of Klebsielleae, from 10<sup>7</sup> viable organisms per ml to 10<sup>5</sup> viable organisms per ml, did not result in diminution in aesculin hydrolysis within 16-hour incubations. Similar dilutions of inocula of non-Klebsielleae resulted only in a change in aesculin hydrolysis of Pseudomonas spp where a distinct reduction in activity was noted.

**FERMENTATION OF ISOINETOL**

A semiquantitative determination of inositol fermentation by Gram-negative bacilli in an inositol-hydrogen sulphide-motility medium was undertaken. As previously described (Barr and Mahood, 1976), and as indicated in Table 1, Klebsielleae could be divided into two groups on the basis of rate of inositol fermentation. K. aerogenes/oxytoca/atlantae/edwardsii and Enterobacter aerogenes were normally rapid fermenters while K. pneumoniae, K. ozoaena, K. rhinoscleromatis, and Enterobacter/Serratia, other than Enterobacter aerogenes, were frequently slow or non-fermenters of inositol (Table 1).

Among non-Klebsielleae, 39 isolates (8-1%) fermented inositol under the experimental conditions described. Eleven of 20 isolates of P. rettgeri gave a strong fermentation reaction comparable to that shown by the majority of the K. aerogenes/oxytoca/atlantae/edwardsii and Enterobacter aerogenes group. Inositol fermentation by other non-Klebsielleae, including 15 isolates of E. coli, was recorded as weak.

**COMPLEMENTARY VALUE OF AESCULIN HYDROLYSIS AND INOSITOL-HYDROGEN SULPHIDE-MOTILITY MEDIA**

It has been shown here (Table 1) that during routine screening of Gram-negative bacilli, inositol fermentation by a considerable number of Klebsielleae was weak or absent. Difficulties in differentiation of Klebsielleae on this basis from other Enterobacteriaceae was compounded by the ability of many strains of P. rettgeri to ferment inositol strongly and the demonstration of weak inositol fermentation, particularly by other Proteus isolates and by E. coli. The evidence presented demonstrates that Klebsiella spp and Enterobacter aerogenes could not be distinguished as a group on the basis of a semiquantitation of inositol fermentation alone.

However, all strains of Klebsielleae examined, excepting one, showed a similar rapid hydrolysis of
sulphide-motility
and potassium
of
rettgeri,
48
are
quoted
number of
"Fermentation of
Motility
was
aerogenes,
Table 2 Typical test reaction
cation
in parallel could
media
from
non-Klebsielleae investigated, only five isolates,
was
hydrolysis, inositol
included
isolates showed
comparable activity.

The results suggested that the use of these two
media in parallel could be utilised in the
differentiation of Klebsielleae and in the presumptive
identification of particular species within that group. Among
non-Klebsielleae investigated, only five isolates, all
P. rettgeri, gave a positive reaction in both aesculin
and inositol media.

An analysis of the results obtained for aesculin
hydrolysis, inositol fermentation, and motility is
summarised in Table 2. Growth in KCN medium
was subsequently added to this biochemical test
sequence to enable the differentiation of K. pneu-
moniae (Friedlander's bacillus) from other Klebsiellea
spp.

Since all excepting one Klebsielleae isolate in-
vestigated demonstrated evidence of rapid aesculin
hydrolysis during 16-hour incubations, only strains
showing strong aesculin hydrolysis have been in-
cluded in Table 2. No isolate giving rapid aesculin
hydrolysis was shown to produce hydrogen sul-
phide in the inositol-hydrogen sulphide-motility
medium. The results clearly demonstrate that the
complementary use of these two test media would be
of value in the presumptive identification and
differentiation of Klebsielleae.

With the use of aesculin alone, six isolates only
were encountered which gave rapid hydrolysis
typical of Klebsielleae. With the use of an inositol-
screening medium alone a much larger number of
non-Klebsielleae (39) was encountered, the majority
being differentiated from Klebsiellea spp on the basis
of motility. Only two isolates of P. morganii could be
differentiated from motile Klebsielleae on the basis
of hydrogen sulphide production, and no inositol-
fermenting isolate of P. rettgeri, 16 in number,
produced hydrogen sulphide in the inositol-hydrogen
sulphide-motility medium.

The use of the inositol medium in parallel did,
however, allow a presumptive differentiation among
the Klebsielleae to be made. The non-motile rapid
inositol-fermenting species K. aerogenes/oxytoca/a-
atlantae/edwardsii are differentiated particularly from
K. pneumoniae and K. ozanae, and among the motile
Klebsielleae, Enterobacter aerogenes is differentiated
from other Enterobacter spp and Serratia spp.

The utilisation of these two test media, and KCN
agar medium, in parallel may enable the diagnostic
bacteriologist to detect significant differences among
isolates of Klebsielleae before embarking on exten-
sive biochemical testing.

**Table 2** Typical test reaction patterns, given by rapid aesculin hydrolysing (+ + +) isolates, on inositol-hydrogen
sulphide-motility and potassium cyanide medium: percentage isolates of each species giving particular reactions
are quoted

<table>
<thead>
<tr>
<th>Fermentation of Inositol</th>
<th>Motility</th>
<th>Growth on KCN agar</th>
<th>Organisms</th>
<th>% Isolates with reaction pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td>-</td>
<td>+</td>
<td>+</td>
<td>Klebsiella aerogenes, etc. (202)</td>
<td>82.1</td>
</tr>
<tr>
<td>+</td>
<td>-</td>
<td>+</td>
<td>K. aerogenes, etc. (202)</td>
<td>8.9</td>
</tr>
<tr>
<td>-</td>
<td>+</td>
<td>+</td>
<td>K. ozanae (11)</td>
<td>54.9</td>
</tr>
<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>K. rhinoscleromatis (4)</td>
<td>50.0</td>
</tr>
<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>K. aerogenes, etc. (202)</td>
<td>3.4</td>
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<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>K. ozanae (11)</td>
<td>27.2</td>
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<tr>
<td>+</td>
<td>+</td>
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<td>K. rhinoscleromatis (4)</td>
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<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>K. aerogenes, etc. (202)</td>
<td>0.5</td>
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<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Enterobacter aerogenes (13)</td>
<td>92.3</td>
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<td>+</td>
<td>+</td>
<td>Enterobacter/Serratia (29)</td>
<td>3.4</td>
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<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Proteus rettgeri (2)</td>
<td>100.0</td>
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<td>+</td>
<td>Enterobacter aerogenes (13)</td>
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<tr>
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<td>+</td>
<td>+</td>
<td>Enterobacter/Serratia (29)</td>
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<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Enterobacter/Serratia (29)</td>
<td>27.2</td>
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<td>+</td>
<td>+</td>
<td>+</td>
<td>K. pneumoniae (15)</td>
<td>6.6</td>
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<td>+</td>
<td>+</td>
<td>K. pneumoniae (15)</td>
<td>53.3</td>
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<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>K. pneumoniae (15)</td>
<td>40.1</td>
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<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Escherichia coli (4)</td>
<td>100.0</td>
</tr>
</tbody>
</table>

1. Fermentation of inositol was recorded as (−) no fermentation, (+) indicator colour change along line of stab inoculation, (+ + +) clear zone of indicator colour change. (+ + +) complete indicator change within individual inoculation compartment.
2. Motility was recorded after 16 hours and after 40 hours' incubation. Motility recorded as (±) indicates motility in majority of isolates.
3. Total number of isolates of each species examined (bracketed).
4. *Klebsiella aerogenes, etc.* includes *K. aerogenes/oxytoca/aatlantae/edwardsii.*
success, particularly in the identification of Gram-negative bacterial urinary tract infections. The scheme incorporates indole production, phenylalanine deaminase activity, ornithine decarboxylase activity, aesculin hydrolysis, inositol fermentation, and growth on xylose-l-lysine-deoxycholate agar. Within this scheme, however, inositol has been maintained in a motility agar medium, and a KCN agar medium has been introduced. The latter medium enabled differentiation of *K. pneumoniae* from other Klebsielleae and contributed to the differentiation between other genera.

In this scheme the evaluation of ornithine decarboxylase activity enables the differentiation of *Klebsiella* spp from other Klebsielleae and overcomes the need for dependence on demonstration of organism motility. This is clearly an advantage, because of the intrinsic difficulty in predictably demonstrating motility in agar in some 16-hour incubations and in view of the prevalence among hospital Enterobacteriaceae of non-motile isolates of *Serratia* spp and *Enterobacter* spp (Oberhofer and Hajkowsky, 1970).

By adopting this scheme, *E. coli* and *P. rettgeri*, which in a simple aesculin-inositol-motility screening regimen could be mistaken for Klebsielleae, are adequately differentiated. *P. rettgeri* is differentiated on the basis of phenylalanine deaminase activity. Rapid aesculin hydrolysing isolates of *E. coli* which, in the simple scheme, could not be differentiated from *K. pneumoniae*, except in the presence of an adequate demonstration of motility, are clearly separated on the basis of indole production. As far as we are aware, no reports of indole-positive *K. pneumoniae* have been noted, although indole-positive isolates of other Klebsielleae are frequently encountered (Rosner, 1970; Davis and Matsen, 1974). The rapid recognition of *K. pneumoniae* may be of special value, particularly if the widely held view that this species is potentially the most pathogenic *Klebsiella* spp was upheld.

In this laboratory, a compromise has been reached between the use of the simple biochemical testing scheme described and the more extensive biochemical identification screen. When staff are available during weekdays the complete range of tests are carried out and isolates are reported at genus or species level at the same time as antibiotic sensitivity reports are made.

At weekends, or other periods of staff shortage, only aesculin and inositol media are utilised. Although the limitations of this practice are noted, the system allows, in the majority of cases, the accurate reporting of isolates as *Klebsiella* spp, *Proteus* spp, or coliform species. For epidemiological studies, isolates examined and reported by the limited range of tests can be stored and identified at a later date.

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References


