The Discovery, Isolation, and Classification of Various 
α-Haemolytic Micrococci which Resemble Aerococci

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SUMMARY

This work describes the identification and classification of a hitherto
unknown group of α-haemolytic micrococci isolated from bottles containing
dregs of fluid medicaments. The majority of the strains were both catalase-
positive and nitratase-positive. In view of several similarities between
these bacteria and the aerococci described by Shaw, Stitt & Cowan (1951),
and later by Williams, Hirsch & Cowan (1953), it is proposed that both
groups should be incorporated in a new bacterial family, Aerococcaceae,
despite the fact that the aerococci are catalase-negative and nitratase-
negative. The suggestion that a new family should be established is
prompted by the many dissimilarities between both the aforementioned
groups and the representatives of the families to which they are closest
according to the Bergey system. The new family ought to include one genus,
Aerococcus, and two species, one of which, A. viridans, should comprise
catalase-negative and nitratase-negative strains, and the other, A. cata-
lasicus, strains which are obligate catalase-positive and may also be
nitratase-positive.

INTRODUCTION

In the course of investigations made into the microbial flora found in the dregs
of fluid medicaments in bottles handed in at three Oslo pharmacies in exchange for
new medicines, I isolated α-haemolytic micrococcus strains which, in their haemo-
lytic, cultural and morphological properties, appear to be akin to the α-haemolytic
micrococci described by Shaw et al. (1951). It was proposed by Williams et al. (1953)
that these bacteria, which are generally present in air and dust, should form a new
genus, Aerococcus, and two species, one of which, A. viridans, should comprise
catalase-negative and nitratase-negative strains, and the other, A. cata-
lasicus, strains which are obligate catalase-positive and may also be
nitratase-positive.

Deibel & Niven (1960) found Aerococcus viridans to be identical, or very similar,
to Gaffkya homari and proposed that both these micro-organisms should be included
in a single species in the genus Pediococcus, since there are several characteristics
common to these two species and the pediococci.

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Tjeltveit & Clausen (1964) have recently shown *inter alia* that the strains of \( \alpha \)-haemolytic micrococci and a strain of *Aerococcus viridans* are homofermentative and produce lactic acid when a sufficient number of active bacterial cells are present during conversion of the glucose. The specific rotation of the lactic acid formed was not investigated.

*Cultural and morphological characteristics of the \( \alpha \)-haemolytic micrococcus strains*

Because the growth of these bacterial strains on solid media, such as ordinary agar and blood agar medium, resembles that of enterococci, they were at first considered to be \( \alpha \)-haemolytic species of *Streptococcus faecalis*. This assumption was borne out by their capacity for growth on 40% ox bile blood agar, in 6.5% sodium chloride beef-infusion peptone phosphate broth, and in the same broth with a normal content of sodium chloride (0.8%) but with a pH value of 9.6. However, microscopic examination of ordinary broth cultures failed to reveal any true chain formation. The cocci were comparatively large, being approximately 1-2 \( \mu \) in diameter, spherical or slightly irregular in shape, though not elongated; they occurred singly, in pairs, and in micrococcus-like formations. They were Gram-positive. Eleven culturally similar \( \alpha \)-haemolytic micrococcus strains were isolated from the medicine bottles and these strains were subjected to detailed study.

The \( \alpha \)-haemolytic strains develop at 22°, 30°, and 37°; growth is poorest at 22°, best at 37°. On 5% citrated horse blood agar the size of the colonies ranges from 0.5 to 2 mm. diam. after 18-24 hr of incubation at 37°. They are convex in shape and vary in colour from grey to white. On one occasion the colonies were clear, transparent, and formed 'craters' when they aged, as do pneumococcus colonies. The \( \alpha \)-haemolytic micrococci formed colonies with green zones on 'chocolate' agar. Aerobic cultivation for 24 hr at 37° in beef-infusion peptone phosphate broth and in the same broth with 1% glucose resulted in sparse growth and relatively meagre sediment. When grown in broth with a petroleum jelly seal, all strains showed slighter growth with the formation of a little sediment after 24 hr incubation at 37° than when cultured aerobically. All strains grew well in modified HS-medium with 10% normal horse serum (Clausen, 1956), but the best growth occurred in the upper third of the medium. When a comparison was carried out using the type strain *Aerococcus viridans* NCTC 8251, it was found that this—compared with the eleven \( \alpha \)-haemolytic micrococcus strains—produced relatively weaker growth and a granular sediment in the aforementioned media.

The \( \alpha \)-haemolytic micrococci were examined according to the methods described below.

**METHODS**

Formulae are not given for media the composition of which must be assumed to be generally known. Unless otherwise stated, incubation was carried out at 37°. All cultures were incubated aerobically.

*Optimum temperature for growth.* Blood agar plates were prepared by addition of 5% citrated horse blood to a medium consisting of 1% peptone (Danish, Orthana Bacteriological Brand), 0.8% NaCl, 0.2% Na\textsubscript{2}HPO\textsubscript{4}, and 1.8% agar (Japanese, quality Kobe I) in aqueous beef-infusion (pH = 7.4). Plates were poured to a
constant depth, about 4 mm. Each individual strain was subcultured to three blood agar plates; one was incubated at 22°, the others at 30°, and 37°. Growth was recorded after 24 and 48 hr.

**Morphology.** Unstained wet preparations of 24 hr cultures of beef-infusion peptone phosphate broth with and without 1% glucose were examined. Gram-preparations of the broth and blood agar cultures were examined.

**Haemolysis** was tested on 5% citrated horse blood agar after 24 and 48 hr incubation.

**Ox bile resistance** was studied by inoculation on to 40% ox bile blood agar (40% autoclaved ox bile, 8% defibrinated horse blood, and 52% agar medium with 2-6% agar, quality Kobe 1) and incubation for 20 hr.

**Tellurite resistance** was investigated by inoculating the strains on to McLeod’s blood agar containing 0-04% potassium tellurite. Growth or lack of growth was recorded after 20 hr incubation.

**Thermoresistance** was tested by heating a 20 hr beef-infusion peptone phosphate broth culture at 60° (in a water bath) for 30 min., cooling, and subculturing to new broth and blood agar. The media were incubated for 48 hr before the results were recorded. Each broth culture to be tested was drawn sufficiently far up a Pasteur pipette to enable the ends of the pipette to be sealed without heating the cultures. The pipette was then plunged so deeply into a water bath at 60° that the entire section moistened with culture was submerged.

**Growth tests** were carried out in beef-infusion peptone phosphate broth with 6-5% NaCl or with a pH of 9-6; incubation period, 72 hr.

**Methylene-blue reductase** was tested in milk culture to which had been added 0-1% methylene blue. The reduction or lack of reduction was recorded after 20 and 48 hr and finally after 5 days’ incubation.

**Reduction prior to coagulation of litmus milk** was observed after 12 and 24 hr, and daily for 5 days.

**Utilization and acid formation of certain sugars, alcohols and glycosides.** 0-5% (aesculin 0-2%) of the following sugars, etc., were added to peptone water +5% normal horse serum +1% Andrade’s indicator: lactose, glucose, maltose, sucrose, raffinose, galactose, xylose, mannitol, dulcitol, glycerol, starch, dextrin, salicin, aesculin. Incubation period: 14 days.

**Proteolytic activity:** ability to liquefy gelatin was tested in stab cultures incubated at 22° for 30 days.

The ability to liquefy inspissated ox serum was determined by incubation at 22° for up to 14 days.

**Formation of catalase.** 10% H₂O₂ was added to 24 hr agar slant cultures and observed for up to 5 min. to determine whether gas was formed. Control tests were performed on non-inoculated agar slant media after 24 hr incubation.

**Formation of nitratase.** Four-day cultures in nitrate broth were tested by adding Griess–Ilosvay reagent. Negative tests were checked by adding powdered zinc to prove that there was nitrate left in the culture medium. A surplus of nitrite in a culture can ‘conceal’ a positive reaction by decolorizing the red stain; for this reason the test was performed daily for up to 4 days (Shaw et al. 1951).

**Formation of plasminocoagulase.** Tests for this enzyme were performed by mixing equal parts of 24 hr broth culture and a 1/10 dilution of citrated human plasma in
sterile saline solution, followed by incubation in a water bath at 37° for 4 hr and storage overnight at room temperature.

*Formation of H₂S.* Stab cultures in lead acetate agar were incubated for 14 days.

*Hydrolysis of arginin* in broth accompanied by formation of NH₃ was demonstrated by mixing equal parts of Nessler's reagent and 48 hr arginine-containing broth culture.

*Hydrolysis of sodium hippurate (1%)* in broth into sodium benzoate after 5 days' cultivation was determined by means of 12% FeCl₃ solution with 0·2% concentrated HCl (Roemer, 1948).

*Hydrolysis of urea.* Christensen (1946) urea medium with indicator was used to demonstrate formation of NH₃ by urease activity. Incubation period: 14 days.

*Methyl red (MR) and Voges-Proskauer (VP) tests* were carried out with the following incubation periods and temperatures: MR test: 4, 5 and 7 days at 30°. VP test: 3 and 4 days at 30°.

*Determination of final pH* in 1% glucose beef-infusion peptone phosphate broth (pH = 7·4) after 14 days. The pH was measured with a Beckman pH meter.

The following bacteria were used in certain tests as reference species: *Aerococcus viridans* NCTC 8251, *Pediococcus cerevisiae* NCTC 8066 and *P. acidilactici* NCIB 6990.

**RESULTS**

Most of the results of these investigations are given in Table 1. Comments on the results not tabulated are given below.

The test for tellurite resistance gave varying results, as one or two of the strains were tellurite-resistant, but the majority were not, or showed only traces of growth on tellurite medium. This test would appear to be of no significance in classifying these bacteria.

Growth in broth with 6·5% NaCl was investigated with a positive result on all micrococcus strains as well as on *Aerococcus viridans*.

The test for hydrolysis of sodium hippurate gave positive results in the case of eight of the eleven strains examined, and two showed traces of hydrolysis. The test was not performed on aerococci.

In tests not recorded in the tables, the α-haemolytic micrococcus strains did not form soluble haemolysin, they were non-pathogenic when injected intraperitoneally into white mice, and did not precipitate with streptococcus antisera of group D (Clausen, 1961).

The α-haemolytic micrococci proved biochemically different from *Aerococcus viridans* in the following ways. (The biochemical properties of the aerococcus strains were determined by Williams et al. 1953, and Shaw et al. 1951).

1. Reduction prior to coagulation of litmus milk. Five of eleven strains of α-haemolytic micrococcus reduced litmus in milk prior to coagulation, whereas none of 29 aerococcus strains did so.

2. Acid production from raffinose. All α-haemolytic micrococcus utilized this carbohydrate, whereas only four of twelve aerococci did so.

3. Acid production from mannitol. Only one of eleven α-haemolytic micrococcus strains utilized this alcohol, whereas five out of twelve aerococci are reported to have done so.
a-Haemolytic micrococci

(4) Liquefaction of inspissated ox serum. Nine out of eleven a-haemolytic micrococcus strains liquefied inspissated ox serum, whereas none of 29 aerococcus strains did so.

(5) Formation of catalase. Eight out of eleven a-haemolytic micrococcus strains were catalase-positive, the majority being relatively strong formers of catalase.

Table 1. Cultural and biochemical properties of a group of a-haemolytic micrococci compared with various strains of Aerococcus viridans

<table>
<thead>
<tr>
<th>Property</th>
<th>a-Haemolytic micrococci</th>
<th>No. positive out of 11 strains tested</th>
<th>A. viridans</th>
<th>No. positive out of 12 strains tested*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optimum temperature for growth 37°C</td>
<td>+</td>
<td>11</td>
<td>+</td>
<td>66%†</td>
</tr>
<tr>
<td>a-Haemolytic on blood agar</td>
<td>+</td>
<td>11†</td>
<td>+</td>
<td>12</td>
</tr>
<tr>
<td>Growth on 40% ox bile agar</td>
<td>+</td>
<td>11</td>
<td>+</td>
<td>12</td>
</tr>
<tr>
<td>Resist 60° for 30 min.</td>
<td>+</td>
<td>11</td>
<td>+</td>
<td>12</td>
</tr>
<tr>
<td>Growth in broth at pH 9.6</td>
<td>+</td>
<td>11</td>
<td>+</td>
<td>11</td>
</tr>
<tr>
<td>Reduction of 0.1% methylene blue in milk</td>
<td>−</td>
<td>1 (2)§</td>
<td>−</td>
<td>0</td>
</tr>
<tr>
<td>Reduction prior to coagulation of litmus milk</td>
<td>+</td>
<td>5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Utilization of</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lactose</td>
<td>+</td>
<td>9</td>
<td>+</td>
<td>10</td>
</tr>
<tr>
<td>Glucose</td>
<td>+</td>
<td>11</td>
<td>+</td>
<td>12</td>
</tr>
<tr>
<td>Maltose</td>
<td>+</td>
<td>11</td>
<td>+</td>
<td>12</td>
</tr>
<tr>
<td>Saccharose</td>
<td>+</td>
<td>11</td>
<td>+</td>
<td>12</td>
</tr>
<tr>
<td>Raffinose</td>
<td>+</td>
<td>11</td>
<td>−</td>
<td>4</td>
</tr>
<tr>
<td>Galactose</td>
<td>+</td>
<td>11</td>
<td>+</td>
<td>97%†</td>
</tr>
<tr>
<td>Xylose</td>
<td>−</td>
<td>0</td>
<td>+</td>
<td>41%†</td>
</tr>
<tr>
<td>Mannitol</td>
<td>−</td>
<td>1</td>
<td>+</td>
<td>5</td>
</tr>
<tr>
<td>Dulcitol</td>
<td>−</td>
<td>0</td>
<td>−</td>
<td>0†</td>
</tr>
<tr>
<td>Glycerol</td>
<td>+</td>
<td>9</td>
<td>+</td>
<td>97%†</td>
</tr>
<tr>
<td>Starch</td>
<td>−</td>
<td>4</td>
<td>−</td>
<td>0†</td>
</tr>
<tr>
<td>Dextrin</td>
<td>−</td>
<td>4</td>
<td>−</td>
<td>38%†</td>
</tr>
<tr>
<td>Salicin</td>
<td>+</td>
<td>8</td>
<td>+</td>
<td>83%†</td>
</tr>
<tr>
<td>Aesculin</td>
<td>+</td>
<td>8</td>
<td>+</td>
<td>12†</td>
</tr>
<tr>
<td>Gelatin liquefaction</td>
<td>−</td>
<td>0</td>
<td>−</td>
<td>0</td>
</tr>
<tr>
<td>Liquefaction of inspissated ox serum</td>
<td>+</td>
<td>9</td>
<td>−</td>
<td>0†</td>
</tr>
<tr>
<td>Catalase formation</td>
<td>+</td>
<td>8</td>
<td>−</td>
<td>0†</td>
</tr>
<tr>
<td>Nitratase formation</td>
<td>+</td>
<td>6 (7)§</td>
<td>−</td>
<td>0†</td>
</tr>
<tr>
<td>Plasma coagulase formation</td>
<td>−</td>
<td>0</td>
<td>−</td>
<td>0</td>
</tr>
<tr>
<td>H₂S formation</td>
<td>−</td>
<td>0</td>
<td>−</td>
<td>0</td>
</tr>
<tr>
<td>Hydrolysis of arginine</td>
<td>−</td>
<td>0</td>
<td>−</td>
<td>0</td>
</tr>
<tr>
<td>Hydrolysis of urea</td>
<td>−</td>
<td>0</td>
<td>−</td>
<td>0</td>
</tr>
<tr>
<td>MR positive</td>
<td>−</td>
<td>0‡</td>
<td>+</td>
<td>93%†</td>
</tr>
<tr>
<td>VP positive</td>
<td>−</td>
<td>0</td>
<td>−</td>
<td>0</td>
</tr>
<tr>
<td>Final pH</td>
<td>5-2-5-6</td>
<td>11</td>
<td>5-5-5-8</td>
<td>12</td>
</tr>
</tbody>
</table>

* Most of these data are from Williams et al. (1953).
† According to Shaw et al. (1951): the percentage positive out of 29 strains tested.
‡ From distinctly α-haemolytic to faintly α'-haemolytic.
§ The figures in parentheses include also strains which show weak positive reactions.
|| Slow reduction.
** Examined in solid media.
** The negative MR results all showed weak to very weak pink, but they were distinctly different from a positive reaction.
Only one of the catalase-forming strains may be described as weakly positive, and one as a moderate to weak catalase former. These two strains did not reduce nitrate to nitrite. The aerococci are reported not to form catalase.

(6) Formation of nitrite from nitrate. Six (+1 uncertain) out of the eleven \(\alpha\)-haemolytic micrococcus strains formed nitrite from nitrate, whereas none of 29 aerococcus strains did so.

**Differences between the \(\alpha\)-haemolytic micrococi, Aerococcus viridans and the pediococci**

The \(\alpha\)-haemolytic micrococi and *Aerococcus viridans* have been compared with *Pediococcus cerevisiae* and *P. acidilactici* with the aid of various growth and resistance tests, which gave the following results: the \(\alpha\)-haemolytic micrococi and *A. viridans* did not grow at 37° (14 days) or 25° (30 days) in the pediococcal media, unhopped wort, yeast-water with autolysed yeast, or Bettge & Heller’s beer with starch (Jørgensen, 1956a), the last two media containing 2.5% (v/v) alcohol, whereas both pediococci grew well in all three media. The \(\alpha\)-haemolytic micrococi and the aerococcus were also MR and VP negative, whereas the pediococci were highly MR positive. On the other hand, only *P. cerevisiae* (not *P. acidilactici*) produced acetyl methyl carbinol. Moreover, the \(\alpha\)-haemolytic micrococi and the aerococcus grew in 6.5% NaCl broth and in broth with a pH of 9.6. Neither of the pediococci developed in these two media; this applied both when incubated at 37° for 5 days and when incubated at 25° for 15 days. Furthermore, the pediococci were not \(\alpha\)-haemolytic when grown on blood agar (*P. cerevisiae* may at the most produce a very weak greening reaction), they have a much lower final pH value (cf. the strong MR positive reaction) than the \(\alpha\)-haemolytic micrococi and the aerococci (Williams et al. 1953), and they form more typical microaerophilic colonies when grown under aerobic conditions on ordinary agar medium and blood agar. One species, *P. cerevisiae*, had a growth-optimum at a comparatively low temperature between 18 and 25° (Jørgensen, 1956b).

**DISCUSSION**

While it has been possible to demonstrate appreciable differences in the biochemical properties of the \(\alpha\)-haemolytic micrococi investigated and the aerococci described by Shaw et al. (1951) and Williams et al. (1953), it has also been shown that there is a great similarity between them, culturally and morphologically, and in regard to many biochemical criteria, especially to the tolerance tests: growth in broth with a pH of 9.6, growth on 40% ox bile blood agar, and resistance to 60° for 30 min. On the other hand the aerococci, unlike the \(\alpha\)-haemolytic micrococi, are reported to be negative with regard to several other properties.

The formation of catalase is probably the most important difference between the \(\alpha\)-haemolytic micrococi and the aerococci, and eight out of eleven of the former strains are catalase formers, whereas the aerococci are reported to be catalase negative. *Aerococcus viridans* (NCRL 8251) showed a negative or at the most a slight trace of a positive reaction. As the formation of catalase is a property characteristic of micrococi, but invariably absent from streptococci, a distinction should be made between catalase-positive and catalase-negative strains of the \(\alpha\)-haemolytic micro-
α-Haemolytic micrococci

cocci. The three catalase-negative strains investigated differed very little in other biochemical properties from the rest of the strains. However, they were nitrite-negative, as is *A. viridans*. A minor difference between them and the one aerococcus strain with which they have been compared (*A. viridans* NCTC 8251) was their formation of denser white colonies on blood agar; the α-haemolysis was weaker in the case of two of the strains, and they also appeared by microscopic examination to be somewhat larger in fluid medium. Moreover, *A. viridans*, unlike the α-haemolytic micrococcus strains, formed granular sediment in broth cultures.

Viewed in the light of these properties, the micrococcus strains investigated may be said to be closely akin to the aerococci, but the catalase-positive and nitratase-positive strains were not identical with the aerococcus species *Aerococcus viridans* proposed by Williams *et al.* (1953).

All the α-haemolytic micrococcus investigated and also the *Aerococcus viridans* strain examined were homofermentative lactic-acid formers (Tjeltveit & Clausen, 1964), but a reference staphylococcus strain, *Staphylococcus epidermidis*, was also found to have the same property. It may be mentioned that Orla-Jensen (1919) found that *S. aureus* formed optically inactive or laevorotatory, and *Micrococcus caseolyticus* dextrorotatory lactic acid from D-glucose, so that the fermentative powers of the strains cannot be regarded as decisive for their classification, either. Morphologically the bacterial strains were not streptococci, and it is obvious that the aerococci and the other catalase-negative microccci lack the conclusive criteria for classification within the enterococcus group, the only group they really can be compared with in the family Lactobacillaceae. The catalase-positive strains cannot be classified in any genus under Tribe I, Streptococceae, *Bergey's Manual* (1957) as all the genera within the Tribe are catalase-negative. From these comparative studies of the α-haemolytic microccci and of *A. viridans* on the one hand and of the pediococci on the other, it was found that the difference between the two groups is so great that there is no justification for including the first group in the genus *Pediococcus*. Nor, without further study, can it be decided whether *Gaffkya homari*, which is of marine origin, is identical with the air and dust bacterium *A. viridans*, as proposed by Deibel & Niven (1960).

The α-haemolytic microccci can be classified neither in the genus *Leuconostoc*, which includes only heterofermentative species, nor in the genus *Peptostreptococcus*, which includes only anaerobes. These α-haemolytic microccci differ, however, to such an extent from the representatives of the genera in the family Micrococceae, that they do not appear to belong here either. The genera of this family do not include α-haemolytic species. Deibel & Niven (1960) found that *Gaffkya homari* produced a strong greening reaction on sheep blood agar, whereas *Bergey's Manual* (1957) describes this bacterium as β-haemolytic on human blood agar. The only genus which might be considered, viz. *Micrococcus*, comprises only species which differ markedly from the bacteria being considered.

It is proposed to establish a new family in the Order Eubacteriales, because of the great specificity of the bacterial group in question. It is now suggested that the species *Aerococcus viridans*, which is catalase-negative, should be supplemented by a new, catalase-positive species. As the generic name of *Aerococcus* has formerly been used, this name should be retained, and the same designation should be incorporated in the new family name.
In the light of the foregoing data it is proposed that the following family, genus, and species names should be given to the bacteria:

- **Family** Aerococcaceae fam.nov.
- **Genus** 1 *Aerococcus* Williams et al.
- **Species** 1 *Aerococcus viridans* Williams et al. (Catalase-negative and nitratase-negative)
- **Species** 2 *Aerococcus catalasicus* sp.nov. (Catalase-positive and possibly also nitratase-positive)

Further details of the family, generic, and specific properties will be evident from the foregoing.

I wish to express my sincerest thanks to S. Dick Henriksen, M.D., Professor at the Medical Institute of Bacteriology at the University of Oslo, who was my chief at the time most of this work was performed, and who helped me to solve many of the problems I encountered in carrying out my investigations. I should also like to record my gratitude to R. Saxholm, M.D., Senior Medical Officer of the Bacteriological Department at the National Institute of Public Health, Oslo, for his generous support in connexion with the later investigations that form part of this work. Finally, particular thanks are due to Mrs Grethe Barbo, laboratory assistant, who rendered me valuable aid in performing many important experiments. I am indebted to S. Rasch, M.Sc., Ringnes Brewery, Oslo, for the pediococcal media he so kindly procured for me.

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