In routine bacteriology the swarming of *Proteus* on plate cultures often masks the identity or presence of other bacteria which may be present. Mixed cultures including *Proteus* are often obtained from open wounds or sinuses following abdominal or chest operations, so that unless some special selective medium is used it is usually impossible to identify or to isolate other bacteria from a culture containing *Proteus*. Frequently \( \beta \)-haemolytic streptococci are, missed in such cultures, and thus one often fails to identify an organism causing some infection which may have become secondarily infected with *Proteus*.

Numerous methods have been published for the inhibition of swarming of *Proteus* on culture plates, the simplest being the use of 6 per cent agar (Hayward and Miles, 1943). This high concentration of agar often leads to technical difficulties in preparation, and if the plates are not sufficiently dried, or if a very moist specimen is inoculated, *Proteus* is not necessarily prevented from swarming. Many chemicals have been incorporated in agar to make selective media, the most popular method being that of adding 1/5,000 sodium azide (Snyder and Lichstein, 1940). Unfortunately, when sodium azide blood-agar plates are used the blood tends to haemolyse and the plates do not keep satisfactorily before use. Jones and Handley (1945) described a satisfactory highly selective medium for the isolation of *Salmonellae* from material contaminated with *Proteus*.

In 1931 Krämer and Koch reported the results of their experiments using 14 different types of media for the inhibition of swarming, and showed how the addition of chloral hydrate prevented the swarming of *Proteus*. In this country the practical use of chloral hydrate plates appears to be little known except in certain laboratories; this paper has therefore been written in order to bring it to the knowledge of everyone as being a practical method.

After certain preliminary experiments it has been found that chloral hydrate in a final concentration of 1/1,000 in nutrient agar or blood agar inhibits the swarming of *Proteus* but allows good growth of *Proteus* and of most other bacteria. Experience with the method coincides with that of Krämer and Koch, who found that practically all bacteria produced typical colonies. One great advantage is that it allows good growth of \( \beta \)-haemolytic streptococci, and also permits the production of \( \beta \) haemolysis, which thus enables one to detect and isolate single colonies from cultures heavily infected with *Proteus*. It is an advantage to incubate the plates for a full 24 hours, although after 18 hours the \( \beta \) haemolysis of haemolytic streptococci can usually be detected with ease.

On 1/1,000 chloral hydrate plates colonies of *Proteus* appear as discrete colonies from 1 to 2 mm. in diameter, so that if necessary one can pick off single colonies of *Proteus* and also single pure colonies of other bacteria from mixed cultures. This is an advantage over media which inhibit the growth of *Proteus*, as on them colonies of other bacteria need not necessarily be pure, as the *Proteus* bacilli just fail to grow but lie dormant and live on the plate. Weaker concentrations, such as 1/1,500, allow the colonies of *Proteus* to become large and semi-confluent, making it difficult to isolate other types of colonies present. At 1/2,000 slight spreading occurs.

Stronger concentrations inhibit the growths of *Proteus* and of other bacteria, but also inhibit the production of \( \beta \) haemolysis on blood-agar plates by haemolytic streptococci.
Preparation of Plates

The method of preparation of chloral hydrate plates is simple. A stock 10 per cent solution of chloral hydrate is made in distilled water and an appropriate amount added to the medium to give a final strength of 1/1,000. It is an advantage to make several plates at one time, adding the chloral hydrate to the bulk base at the time of pouring the plates, otherwise a very small volume must be added to the medium for each plate. It is not necessary to sterilize the stock solution, as it is apparently self-sterilizing at this strength, but it is an obvious advantage to exclude bacteria, especially spore-bearers, by using sterile distilled water. If preferred, the stock solution may be autoclaved without interfering with its inhibitory properties. Once the plates are poured they should be dried in the usual way, but not overdried.

The chloral hydrate blood-agar plates may be stored in the refrigerator for a week before use without the occurrence of haemolysis. Any stored plate should be dried in the incubator for about half an hour before inoculation.

Krämer and Koch report having found occasional batches of blood-agar plates giving atypical growths, but over the last three years no very unusual growth has been found as long as the correct percentage is accurately added to the media. Slight increase in the concentration due to the inaccurate addition of too small a volume of stock solution will lead to irregular results.

Conclusions

1. Chloral hydrate in a final concentration of 1/1,000 in nutrient agar or blood-agar plates inhibits the swarming of Proteus.

2. The chloral hydrate in the strength recommended permits the growth of Proteus as single colonies 1 to 2 mm. in diameter.

3. Chloral hydrate in this strength allows the production of β haemolysis on blood-agar plates by haemolytic streptococci.

4. Chloral hydrate plates permit the identification and isolation in pure culture of bacteria from mixed cultures contaminated with Proteus.

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Chloral Hydrate Plates for the Inhibition of Swarming of Proteus
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