Iron Contents of Different Colonial Types of Neisseria gonorrhoeae

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INTRODUCTION

Four characteristic colonial types of Neisseria gonorrhoeae were described by Kellogg et al. (1963, 1968), of which types 1 and 2 were virulent in human volunteers whereas types 3 and 4 appeared to be laboratory variants and avirulent. In the course of studies on Kellogg's types, Jephcott & Reyn (1971) reported a fifth type. This colonial type 5 was found in about 1% of cultures of human cervical and urethral specimens but its pathogenicity has not been established (Brown & Kraus, 1974). Differences between gonococcal colonial types include possession of pili by types 1 and 2 but not by the other types (Jephcott, Reyn & Birch-Andersen, 1971; Swanson, Kraus & Gotschlich, 1971). Recently, Walstad, Guymon & Sparling (1977) reported that type 3 and dark-coloured variants of types 1 and 2 had larger amounts of outer membrane protein than light-coloured variants of types 1 and 2, and light-coloured colonial type 4.

Bullen, Rogers & Griffiths (1974) drew attention to the importance of iron metabolism in the pathogenicity of some micro-organisms. Although iron is essential for the growth of gonococci, its possible role in pathogenicity has not been studied. Edwards & Seamer (1960) determined the iron content of Corynebacterium diphtheriae and showed that there was a critical optimum iron content for maximum toxin production. We have undertaken the present study to determine the iron content of the various Kellogg colonial types of gonococci believing there may be a similar correlation between iron levels and pathogenicity.

METHODS

Strains. The strains of gonococci studied were: r62 (originally isolated in Kellogg's laboratory) from which types 1, 2, 3 and 4 were derived [these were used freshly subcultured from storage in liquid nitrogen (−120 °C)]; types 1 and 4 of r62 (ANM) [these were the same types except that they had already been passaged several times in the basic liquid medium (ANM) of Hafiz & McEntegart (1976)]; freshly isolated strains from patients attending the Special Clinic at the Royal Infirmary, Sheffield, designated gc338 (types 1, 2, 3 and 4), gc41 (types 1, 2, 3 and 4) and gc113 (types 4 and 5).

Growth and preparation of dried organisms. All the strains were grown on GC agar (Difco) plus 2% defined supplement (Kellogg et al., 1963) for 18 to 20 h at 35 °C in an atmosphere of air plus 10% (v/v) CO₂ and enhanced humidity. The colonies were confirmed as gonococci by their oxidase and fermentation reactions and their typical bright fluorescence when treated with a specific antigonococcal conjugate (Difco). The gonococcal colonial types were selected using the special lighting system of Jephcott & Reyn (1971). Each of the selected types of all strains were grown in bulk on batches of 70 standard (90 mm) GC agar plates (Difco) and, after confirming that the gonococci were of the desired colonial type, the bacteria were harvested into 10 ml deionized water in iron-free Universal containers. The organisms were washed once
RESULTS AND DISCUSSION

The assay over seven separate experiments using FeSO\textsubscript{4} standards was linear up to at least 5.59 µg Fe and was sensitive to 0.22 µg Fe which corresponded to a ΔE of 0.05 above a zero control. The mean value of $E_{538}$ and standard deviation (S.D.) at 5.59 µg Fe was 1.29 ± 0.082 with a standard error of the mean (S.E.M.) of ± 0.031; these S.D. and S.E.M. values were typical of those for other levels of iron.

The values obtained for the iron contents of gonococci (Table 1) fell within the range of those quoted by Neildands (1974) for other species of bacteria. However, the iron contents of the pathogenic colonial types 1 and 2 were lower than those of the relatively avirulent types 3 and 4. Also the iron contents of types 1, 2, 3 and 4 of stored f62 were lower but in
similar proportion to one another as those of the colonial types of the freshly isolated strain GC338. The long storage in liquid nitrogen could have been responsible for this because after several passages in ANM the iron contents of types 1 and 2 of f62 (ANM) equalled those of the equivalent types of GC338.

Waring et al. (1953) demonstrated an increased pathogenicity for animals of Brucella suis rendered relatively iron-deficient in culture media compared with 'iron-normal' cultures of the same organism. They found iron-deficient Brucella suis was more invasive and produced a poorer inflammatory response in the host and suggested that the increased virulence was due to the host defence mechanism being more adversely affected by 'iron-deficient' Brucella suis. Payne & Finkelstein (1975) found that when relatively avirulent gonococci of colonial types 3 and 4 were inoculated along with iron into chick embryos their virulence increased, though iron had little or no effect on the virulence of types 1 and 2 in the same host. They suggested that iron might compensate for the absence of a virulence factor in types 3 and 4 by interfering with the chick embryo defence system. Recent observations in our laboratory suggest that the death of chick embryos injected intravenously with gonococci was not due to infection but to some form of endotoxic shock. Furthermore, there is ample evidence that in the course of experimental infection with Gram-negative organisms there is a release of endotoxins which bring about significant changes in the metabolism of iron by the host (Sussman, 1974). On the basis of our present observations, an increase in the critical environmental iron in the host might provide the necessary level of iron to express gonococcal virulence.

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REFERENCES


