New phenotypic typing scheme for group B streptococci

S R Heard, J A Mawn

Abstract

Aims: To develop a new typing system for group B streptococci based on \(^{35}\)S-methionine-labelled protein profiles of bacterial proteins.

Methods: 377 clinical isolates of group B streptococci were examined by incorporation of \(^{35}\)S-methionine into bacterial proteins under strict anaerobic conditions. After sodium dodecyl sulphate polyacrylamide gel electrophoresis, autoradiography was performed. The patterns produced were visually analysed and categorised into clusters of organisms based on the pattern of band production between 32–46 kilodaltons.

Results: 294 of the typed strains classified into seven different groups designated A–G. 32 strains failed to incorporate \(^{35}\)S-methionine sufficiently to be grouped and 11 strains did not fall into one of the seven identified groups. Typability, reproducibility, and discrimination of the system was evident.

Conclusions: This typing system may help to distinguish between colonising and invasive strains of the organism.

(J Clin Pathol 1993;46:145–148)

Group B streptococci are commensals of the gastrointestinal and female genital tracts. Up to 28% of pregnant women may carry this organism in the vagina.\(^1\) It is an important neonatal pathogen, responsible for both early and late onset neonatal sepsis which is characterised by a potentially life-threatening pneumonia, septicemia, or meningitis. The incidence of neonatal disease varies, with two to three cases per 1000 live births being reported from the United States\(^2\) and 0-3 per 1000 live births in the United Kingdom.\(^4\) Prematurity and low birthweight, prolonged rupture of membranes, and maternal pyrexia are recognised as important risk factors in susceptible mother-infant pairs.\(^5\) Maternal sepsis accounts for 10–20% of septicemas after delivery in the United States: group B streptococci is the second most common blood culture isolate in this setting.\(^6\) The pathogenesis of early onset neonatal disease is related to the degree of genital colonisation in the mother\(^7\) which may be reduced by prophylaxis with chlorhexidine during delivery,\(^8\) and to additional risk factors which contribute to the potential for the organism to cause invasive disease.\(^9\) The epidemiology of late onset disease is less well defined but both maternal and nosocomial sources have been implicated.\(^10\) Cross-infection resulting in nosocomial outbreaks in nurseries has also been well documented.\(^11\)

Current typing methods for the organism are based on a reference laboratory typing scheme using serotyping followed, if necessary, by phage typing. This combined approach may be required because although most clinical strains can be serotyped, this system lacks sufficient discrimination, and less than 80% of strains can be typed using bacteriophages alone.\(^12\) Serological classification is based on Lancefield's original description of two cell wall carbohydrate antigens, the group specific common antigen carried by all strains, and the type specific capsular antigen. Clinical strains are currently divided serologically according to the presence of specific antigen into seven groups (Ia, Ib, Ia/c, II, III, IV, VI).\(^13\) Groups I, II, and III are associated with early onset neonatal disease, while type III causes 90% of all cases of both early and late onset meningitis and 90% of all cases of late onset disease.\(^14\) In addition to the polysaccharide antigens, C proteins on the surface of group B streptococci form a complex composed of at least four different components (\(\alpha, \beta, \gamma\), and \(\delta\)) which are found on all Ib strains and in up to two thirds of type II and III strains.\(^15\) Additional protein antigens, termed R and X, have also been described. These are primarily associated with type II and III strains.\(^16\) An epidemiological study using restriction endonuclease analysis has recently demonstrated the usefulness of this technique in examining recurrent invasive disease\(^17\) and in outbreaks,\(^18\) but it has not been formulated into a typing system.

Neither serological nor phage typing systems can distinguish between colonising strains and those which have the potential to cause invasive disease in neonates. It is clear that host immunological factors, including the amount of type specific protective antibody that is present, are of critical importance in the development of invasive disease,\(^19\) but pathogenic factors related to particular strains of invasive organisms are still not well defined.

We were specifically interested in investigating the role of bacterial proteins in strain speciation as there is evidence that at least the C protein is a likely virulence factor and may, in fact, be a potentially good candidate for a vaccine.\(^20 21\) The immunological study of the disease has certainly identified one aspect of
invasive disease, but it may be that characteristics of the organism itself can account, in part, for its clinical behaviour. To date, serological, phage, and molecular methods at classification systems have failed to be of benefit in identifying potentially "high-risk" invasive strains. The typing system presented here attempts to classify the organism at the phenotypic level and is based on the development of a typing scheme using 35S-methionine incorporation into bacterial proteins.

Methods
Sequential group B streptococci isolates grown under standard laboratory conditions from high vaginal swabs received in the clinical laboratory were purified and stored in glycerol broth at −70°C. After detailed experimental conditions for the optimal incorporation of 35S-methionine into group B streptococci had been ascertained, the most important of which is the requirement for strict anaerobic conditions, the stored strains were inoculated on to pre-reduced blood agar and Islam’s media (Oxoid, Basingstoke) and were incubated in an anaerobic chamber (10% H2, 10% CO2, 80% N2; Don Whitley Instrumentation) for 24 hours. 35S-methionine (Amersham International) at a final concentration of 10 μCi was added to 100 μl aliquots of sterile pre-reduced methionine free modified Eagle’s medium (Flow Laboratories) which was inoculated with a standard loop of group B streptococci calculated to 20 μg of protein. The mixture was then incubated for two to eight hours at 37°C under strict anaerobic conditions. An equal volume of double strength cracking buffer (2% 2-mercaptoethanol, 20% sodium dodecylsulphate (SDS), and 40% glycerol) was then added to stop the incorporation. After boiling for 5 minutes and cooling, 35 μl samples were loaded into sample lanes of a 4% stacking gel and electrophoresed into a 10% SDS-acrylamide (BDH Laboratories) resolving gel. 3C-methylated molecular weight markers (Range: 14.3–200 kilodaltons) to a final activity of 0.1 μCi were run with each gel, and as the system became clarified, appropriate specific controls were also used. Eight standard NCTC strains were also analysed using this typing methodology. Twenty five per cent of the strains were typed between two and four times to assess the reproducibility of the method.

Electrophoresis was performed using a Protean II dual vertical slab gel with 16 cm plates (Biorad Laboratories, Richmond, California) at 4°C. A constant current of 35 mA was applied to run the samples through the stacking gel, after which the current was increased to 55 mA for 3–4 hours. The gels were fixed (10% acetic acid, 10% isopropanol), dried, and autoradiographed. Analysis of the autoradiographs was performed independently by two researchers (SRH and JM) by visual analysis of the bands which were within the molecular weight range 30–46 kilodaltons.

Results
Three hundred and thirty seven strains were studied. A comparative visual analysis of the bands in the molecular weight range 30–46 kilodaltons showed that 294 (87%) of strains could be classified into seven groups designated a–g. Consideration of bands above or below this region complicated the classification, and did not contribute to reliable discrimination among strains. Thirty two strains (9%) failed to incorporate the 35S-methionine sufficiently to be grouped and 11 strains could not be classified into one of the seven groups. The 73 strains typed on more than one occasion gave the same result each time.

The groups distinguished by a distinctive pattern on 35S-methionine protein analysis are designated as a, b, c, d, e, f and g. Autoradiographs showing these seven groups are shown in fig 1. The distribution of the 294 typed strains within the seven strains is shown in the table. Figure 2 comprises autoradiographs of eight standard NCTC serotyped strains of group B streptococci. These show that different serogroups may fall within the same 35S-methionine group (strains typing as serogroup VI (lane 3) and III (lane 10) both type as f, and conversely, that the same serogroup may have different 35S-
methionine types (strains 8182 (lane 2) and 8100 (lane 9), which are both Ic strains, fall into groups f and b, respectively). Figure 3 shows previously untyped strains in relation to the seven standard strains in order to demonstrate the typing system "in use". Most strains were readily classifiable, but 3% of strains could not be classified within the seven existing standard groups, as shown in lanes 7 and 9, a mother and baby pair.

Discussion
The development of a new typing scheme for an organism is time consuming and laborious. For epidemiological purposes, many workers now use a variety of comparative methods to assess whether outbreak strains are the same or different, without placing them within the context of a well defined typing system. Another possible approach is to take a small number of strains classified by existing systems and apply the new system to these as a means of assessment. Our view, however, is that a systematic approach of investigating a large number of clinical isolates should be undertaken to define a comprehensive typing system with the potential to use it for both epidemiological and analytical purposes.

In the case of group B streptococci, existing classifications are based on serotyping systems which identify type specific cell wall carbohydrate groups with some protein antigens, and on a defined phage typing system. Restriction endonuclease analysis (REA) has recently been performed on 54 isolates and seems to help in discriminate between serologically similar strains, but produced 28 different REA patterns. A second study noted two particular restriction digest patterns from nine of 10 strains of neonates with early onset disease. This methodology has not, however, been developed into a coherent typing system and its very complexity suggests that it would have limited use for this purpose.

As there is evidence that bacterial proteins are involved in the pathogenesis of invasive group B streptococci disease, a typing scheme which is based on protein analysis is appropriate. 35S-methionine analysis has been used for epidemiological purposes to investigate outbreaks, but only in the case of Clostridium difficile has the technique been used to develop a comprehensive typing scheme which has been applied clinically.

Less than 10% of strains we tested could not be typed due to failure of incorporation of the 35S-methionine. Group B streptococci grows better under anaerobic conditions and also produces both increased haemolysis and more of its characteristic tan pigment on starch medium if it is grown anaerobically. Initial difficulties with 35S-methionine incorporation into the bacterial proteins were overcome by performing the incorporation under strict anaerobic conditions.

There was good discrimination among the groups with only 3% of strains not allocated to one of the seven identified groups. These are likely to form the basis for further extension and refinement of the system especially as it is applied over a wider geographical area. The system also allows for good reproducibility: the 73 strains which were typed on more than one occasion produced the same result. Reproducibility is more readily achieved by using appropriate controls on the typing gel. With the current complement of seven standard strains, seven standard strains, a molecular weight marker, and seven test strains can all be run on a single gel (fig 3). The strains analysed in this report have not been subject to serotyping because the basis of the serotyping system is the polysaccharide component of the organism. As anticipated, there was no relation between the serogroup and the 35S-methionine group among the serotyped NCTC strains examined. In 35 additional strains which have been both serotyped and typed by the re-
ported system, again no relation was apparent (personal observations).

The development of a new typing scheme based on protein analysis is needed to further our understanding of the epidemiology of group B streptococci and to attempt to identify strains which could be potentially pathogenic. Existing typing schemes are neither sufficiently discriminatory nor comprehensive to offer a solution to this problem. If the fundamental clinical issue is how can we determine which of the 30% of pregnancies colonised with group B streptococci are at risk from invasive disease in the neonate or mother, then several questions, including those related to the immunology of the disease, pathogenicity related to structure, adhesion, and possibly toxin production, and the identification of strains which can both colonise and invale, need to be considered. The \( ^{35} \)S-methionine system reported here is currently being used to study invasive strains of the organism to consider whether this phenotypic classification can be of clinical value in determining which pregnancies require early intervention for carriage of invasive group B streptococci strains.

We gratefully acknowledge the support of the Joint Research Board and the Special Trustees of St Bartholomew's Hospital.

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doi: 10.1136/jcp.46.2.145

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