Uncertainties in screening and prevention of Group B Streptococcus disease

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Short summary: Our article presents current research gaps related to GBS screening strategy, disease prognostic factors, serocorrelates of protection and women’s attitudes to screening and prevention strategies. It provides clearly identified research priorities to stimulate researchers and funders to address these gaps.

**Abstract**

In autumn 2016, the UK Department of Health (now Department of Health and Social Care), convened two meetings to discuss how to address research evidence gaps in order to minimise the impact of infant Group B Streptococcal (GBS) disease in the United Kingdom. At this meeting, a number of research priorities were highlighted including: improvements in screening for GBS colonization in pregnant women and offering intrapartum antibiotic prophylaxis, point of care testing and understanding the effect of widespread intrapartum antibiotic use on long term infant health. Further discussions involved investigating the feasibility of a large prospective study of pregnant women and their infants to understand the role of antibody in the protection against GBS disease in infancy following maternal exposure to GBS colonization. This viewpoint summarises the research uncertainties identified by this meeting.

**Introduction**

Group B Streptococcus (GBS) is a bacterium carried in the urogenital and gastrointestinal or urinary tract in the general population. However, approximately 20% of pregnant women carry GBS at any one time and there is a risk to the infant associated with passage of the bacterium from mother to infant around birth1. GBS can cause septicaemia, pneumonia, meningitis and death in up to 2% of infants born to colonized women in the absence of intrapartum antibiotic prophylaxis (IAP).2 GBS disease primarily occurs in babies in two forms: early onset (EOGBS), occurring between birth and day six of life, and late onset (LOGBS), occurring from day seven to day 90 of life, with the disease practically disappearing thereafter. The risk factors associated with LOGBS are poorly understood. Survivors of GBS disease have a higher risk of long-term neurodevelopmental impairment (particularly following meningitis) that can severely impact quality of life.3 In the UK, a recent national surveillance study has shown that the incidence of culture-confirmed EOGBS disease appears to be rising, from 0.48/1000 (95% confidence interval (CI) 0.43-0.53) livebirths in 2000 to 0.57/1000 (95% CI 0.52-0.62) in 2014, despite a clinical risk factor-based IAP policy introduced in 2003.4 Over the same period of time, LOGBS incidence also appears to have risen from 0.24/1000 (95% CI 0.21-0.28) to 0.37/1000 (95% CI 0.33-0.41) livebirths. GBS is now the leading cause of severe bacterial infection in neonates5 and bacterial meningitis4 in UK infants. Several unanswered questions exist concerning how to prevent all forms of GBS infection.

Policies for GBS screening vary between countries, with some offering universal screening (e.g. through microbiological testing) to all pregnant women whilst others undertake this selectively.6 In the UK, the Royal College of Obstetrics and Gynaecology (RCOG)7 and National Institute for Health and Care Excellence (NICE)8 recommendations have been to offer IAP to women identified as having risk factors for GBS: previous baby with invasive GBS infection, GBS bacteriuria in the current pregnancy, vaginal or rectal swab positive for GBS in the current pregnancy or maternal intrapartum pyrexia above 380C. However, the recent national surveillance study has shown that only 35% of a cohort of 429 UK and Irish cases with early-onset GBS disease had one or more of these risk factors compared to 65% in the 2000–2001 study.9 Since this surveillance was undertaken, the RCOG guidelines have been updated (in 2017)7. These revised guidelines now include preterm labour as an additional risk factor for the offer of IAP and a recommendation for women who have previously had a positive GBS result to either be treated as a carrier in this pregnancy or to offer repeat testing and IAP if appropriate.This could potentially add up to 12% more women to the proportion who might be offered IAP 7. Nevertheless, a risk factor based approach provides only limited protection against invasive EOGBS disease in the infant.

In around half of high-income countries, women not already identified as at increased risk of their baby developing EOGBS are offered a test for GBS colonization at 35-37 weeks’ gestation (‘universal screening’) with subsequent IAP for those whose swab is positive for GBS (around 18-20%).6 Both the risk-based and universal screening approaches entail giving IAP to a substantial proportion of pregnant women, the great majority of whom would not have had an affected baby even without the IAP.

Further uncertainties exist in how best to detect those infants at risk of both early and late onset GBS infection. Current test-based screening is developed to identify maternal GBS colonization rather than neonatal or infant invasive disease. The risk factors used to identify women to be offered IAP are generic clinical risk factors and are not sensitive or specific for EOGBS.

. A recent systematic review concluded that the evidence base addressing the potential impact of widespread use of IAP for the mother (anaphylaxis, antimicrobial resistance, medicalisation of labour) and baby (short- and long-term effects on the gut microbiome, lengthier stay in hospital, antimicrobial resistance) was limited and that further large, high-quality, and longitudinal observational studies across countries would improve our understanding in this area.10 Given that in the UK approximately 20% of women are colonized yet only one in a hundred infants born to women carrying GBS develop invasive GBS disease4, there is a need to understand other factors that contribute to natural protection from infection, such as antibody concentrations. Such data would also facilitate vaccine development and licensure.

A recently published collection of articles has highlighted the potential global burden and the scale of the problem2. Independently (though with overlap with some of the experts), the UK Department of Health (now Department of Health and Social Care), convened a working group to examine some of the research uncertainties that could be tackled in the UK with appropriately commissioned funding. In this article we present a consensus view on current research gaps in evidence on screening, diagnostic tests and vaccination with the aim of stimulating research in this area.

**1: Screening approaches for GBS**

Two main approaches to the screening of pregnant women are widely practised in high- and middle-income settings: universal screening or risk-factor based screening. The Centers for Disease Control and Prevention, American College of Obstetrics and Gynecologists, American Academy of Pediatrics, American Society for Microbiology and American College of Nurse-Midwives have recommended IAP to prevent EOGBS since the early 1990s.11 Following a large multicentre cohort study in 200212 that suggested universal screening to be superior to risk-based screening, guidance in the USA recommended universal screening (using a vaginal and a rectal swab) for GBS colonization at 35-37 weeks’ gestation and among women with threatened preterm delivery and unknown colonization status, and administration of high dose intravenous benzylpenicillin or ampicillin in labor in those with a positive GBS swab.13,14 Additional risk factors for which IAP may be offered include: women with GBS bacteriuria in current pregnancy; a previous infant with GBS disease; vaginal or rectal swab positive for GBS in the current or a previous pregnancy, women with unknown colonization status and intrapartum risk factors such as prolonged rupture of membranes or maternal intrapartum pyrexia above 380C. In the UK, clinical risk factor screening (as outlined above, with the addition of preterm labour) is undertaken. Since the introduction of IAP policies,13,14 culture-confirmed EOGBS disease in the USA declined from 1.7 per 1000 live births in the 1990s to 0.21 per 1000 live births in 2015.7,15,16

The use of clinical risk factor based IAP strategies rather than universal screening is based on the assessment that the introduction of routine microbiological screening may not reduce EOGBS-related mortality and morbidity sufficiently to be cost-effective. The recent National Screening Committee report assessing GBS screening concluded serious concern that it was not clear whether benefits associated with screening outweighed the harms for the majority of the population and that large numbers of women would be offered and take antibiotics when they do not need to, increasing potential risks associated with widespread antibiotic use in both woman and infant.17 A UK test accuracy study found that only 28.9% (89/307) of women with clinical risk factors actually carried GBS (whether tested by rapid intrapartum PCR-based methods or following enriched culture medium of swabs in late pregnancy) and that 19% (205/1080) of women with no clinical risk factors carried GBS.18

A recent Cochrane review evaluating IAP for known maternal GBS colonization identified four randomized trials involving 852 GBS positive women, three of which (more than 20 years old), compared ampicillin or penicillin to no treatment; no clear differences in newborn deaths were found although the occurrence of early GBS infection in the newborn was reduced with antibiotics (risk ratio (RR) 0.17, 95% CI 0.04 to 0.74).19 The review was critical of the quality of the studies, including their small size, and considered there to be a high risk of bias in their methodology and execution, with further adequately sized, double-blind controlled trials needed.

Whilst there is broad agreement that IAP should be offered in the presence of risk factors, there is currently no formal international consensus as to whether IAP is best directed for women through universal screening or based on the presence of clinical risk factors. A European consortium has previously called for a universal intrapartum GBS screening strategy using rapid real time PCR testing.20 A review of universal antenatal culture-based screening for maternal GBS colonization to prevent early-onset GBS disease against programme appraisal criteria for the UK National Screening Committee recommended that the criteria for recommending a universal screening programme were not currently met. The review concluded that further research would require randomized controlled trial evidence, with economic modelling to evaluate the associated costs.21

Several studies from the USA indicate that at least 40% of all infants are exposed to IAP for clinical reasons or because their mothers had a GBS-positive swab in pregnancy22,23. Intrapartum antibiotics are delivered at a key time in the development of the infant’s intestinal microbiome. These microbiota may be important for metabolic, nutritional, physiological and immunological development and rapidly evolve following birth. If the initial exposure to maternal microbiota is altered, such as following a Caesarean birth, studies demonstrate differences in the type, variation and distribution of organisms. The same may be true following IAP and these effects could therefore have long-term health implications. The microbiome has been implicated in adult disease such as obesity, allergy and atopy, inflammatory bowel disease, and the development of colon cancer24. However, causal links have yet to be established and there is a vital need to understand infant gut microbial community succession and to study the impact of early life exposure to IAP on the processes involved in establishing a healthy microbiome.

Several recent studies indicate alterations in infant microbiota in IAP-exposed and IAP-unexposed infants up to 12 months of life. Two Canadian studies, Baby and Mi and CHILD have recently published evidence suggesting early differences in the microbiota of low risk term infants exposed to IAP compared to unexposed infants and those born by caesarean section.25,26 These studies suggest cumulative dysbiosis with IAP and caesarean section and modifying effects of breastfeeding. An Italian study of 84 infants followed for 30 days indicated reduced numbers of beneficial *Bifidobacterium* in IAP-exposed infants, which was further exacerbated in those who were formula rather than breastfed.27 It is therefore possible that IAP may also alter the initial colonising microbiota and this could have long-term health implications. Several larger cohort studies are now underway to investigate the long-term consequences of peri-partum antibiotic exposure following caesarean section on the infant intestinal microbiome and future disease risk.

Box 1 summarises research questions associated with GBS natural course and screening.

2: Testing strategies for bacterial load/virulence – colonization to invasive disease

It is recognized that one major barrier to screening and prophylaxis for GBS disease is that current tests detect GBS colonization in pregnant women rather than predicting infant invasive disease. A better understanding of the factors that determine bacterial virulence and host susceptibility is needed in order to enable development of a microbiological test that better targets the women (and babies) that require antibiotic prophylaxis and avoids giving unnecessary antibiotics to the women and babies who would otherwise remain well. This research should entail underpinning biological studies first, before addressing technological issues (e.g. development of cheaper PCR-based tests that incorporate antibiotic sensitivities) and subsequent clinical evaluation of any new microbiological test.

There is a paucity of data regarding bacterial factors that influence transmission of GBS from mother to fetus and neonate, maintaining homeostasis in the infant intestine or causing invasive disease. Recently, a number of virulence factors and GBS lineages such as hypervirulence Clonal Complex (CC) 1728 and surface proteins (e.g. Rib, Alp and Pilus proteins)29 have been implicated in increased disease risk and colonization persistence. Additionally, the initial inoculum (woman’s bacterial load at point of transmission) has also been associated with an increased risk of EOGBS.30 Further such insights could allow more targeted implementation of IAP to only those women who carry such variants of GBS most likely to cause EOGBS, thus reducing the IAP currently offered to all women with GBS colonization. For LOGBS, even less is understood. It is unclear whether LOGBS is predominantly derived from environmental sources (horizontal transmission) or whether the infant gut harbours pathogens that cause episodes of LOGBS because of genetic alterations after transmission, as has been described with other pathogens such as pneumococcus9,31. A recent study of LOGBS on a single neonatal unit using whole genome sequencing, suggested that the majority of cases in that setting were likely to reflect nosocomial transmission32. Information about the characteristics of GBS that increase the risk of neonatal colonization and persistence may have important implications for more targeted IAP and for second generation vaccine development. Such knowledge would also enable screening tests to be adapted towards these specific markers.

Development of tests that could reliably predict which women will transmit GBS to their child, and which infants will go on to develop disease would be a substantial addition to the current screening programmes. Developing our understanding of the role of immunological factors and GBS strains on birth outcomes and combining this information with the population structure of GBS in colonization and disease will strengthen our knowledge of potential vaccine coverage and molecular diagnostic targets.

Several molecular diagnostic tools are emerging as potential candidates for more rapid identification of invasive GBS disease or to rapidly identify women who are GBS colonized. Rapid diagnostics for invasive disease, including the MinIon®, loop-mediated isothermal amplification,33 and optical immunoassays whilst those for rapid intrapartum colonization screening include PCR-based methods.34 However, little data are publicly available about their sensitivity and specificity for GBS detection in the clinical setting. Recently, Whole genome sequencing (WGS) has been used to investigate GBS colonization factors, providing an opportunity to investigate beyond serotyping and overcomes some of the sensitivity issues with the current latex agglutination tests.35 WGS could be used to identify antimicrobial resistance genes that might be useful targets for any future screening test in the context of penicillin allergy, in order to make better choices around IAP, but as yet the methods are cumbersome and could not be developed in real time.

. An alternative could be a PCR-based method that could target those genes identified by WGS as conferring clindamycin resistance, such as those developed for pneumococcal disease.36

However, it is important to consider that looking at the GBS genomics alone will not provide sufficient evidence for a more targeted IAP. Host factors will also play an important role in identifying those who do and do not progress from colonisation to disease.

Box 2 outlines the research questions surrounding GBS testing.

**3: Group B Streptococcus vaccines**

Clinical evaluation of GBS vaccines using a reduction of invasive neonatal disease as a primary outcome requires large studies, which are therefore best carried out in settings with relatively high prevalence. It is estimated that an efficacy study of approximately 60,000 pregnant women in countries with a disease incidence of more than one in a thousand live births would be required to detect a 75% reduction in early onset and late onset GBS disease.37 This figure assumes that the vaccine would cover approximately 90% of circulating serotypes (ST).37 An alternative approach could be to establish immune correlates of protection based on vaccine or natural antibody studies such has been used for meningococcal and higher valency formulations of pneumococcal polysaccharide-conjugate vaccines.

The issue for GBS, however, is that although there are data to support the concept of an immune correlate of protection,38 it is difficult to link this evidence to the current vaccines in development. There have been several case-control studies that provide evidence that serocorrelates of protection against infant GBS disease are achievable.39-41 These studies ranged in size from 25,000 to 140,000 pregnant women and captured between 33 and 109 cases of neonatal GBS disease. Each study provided evidence that higher antibody concentrations were linked to a reduced probability of contracting disease. However, no study was sufficiently powered to provide a definitive answer and, as the studies were all assessed using different assays and data analysis, it is not possible to compare and pool these results. There is an urgent need to develop a consensus around a validated serocorrelate of protection.

To facilitate more rapid licensure and availability of a GBS vaccine for prevention of early onset and late onset infant GBS disease, a serocorrelate of protection against neonatal invasive GBS disease will undoubtedly prove useful. However, when applying a serocorrelate of protection against a neonatal disease with a defined risk period, where the prevention strategy is vaccinating the pregnant woman, additional factors including placental IgG antibody transfer and antibody decay must be considered. These factors are important as a serocorrelate will likely need to demonstrate that vaccines can generate antibody titers in the mother that can be effectively transferred and persist within the infant so that they are protected not only against early onset, but also late onset disease.

Determining a consensus for a protective antibody concentration is not easily achieved, as protective antibody concentrations may vary by serotype 40,42, and the assessment of immunogenicity varies by the assay methods employed.38,42 It is clear that a well characterized assay is required that has been demonstrated to be robust, reproducible, specific and precise.

Several initiatives are currently underway to facilitate the determination of a serocorrelate of protection against invasive disease including standardisation of assays.43 Capsular polysaccharide CRM197 conjugate vaccine,44 tetanus toxoid protein conjugate vaccine and an Alp/Rib protein adjuvanted vaccine are all in development.45

Box 3 outlines research questions surrounding the development and testing of a GBS vaccine for use in the UK.

**Conclusions**

Many questions remain unanswered and require more evidence if we are to truly reduce the burden and impact of GBS disease in the United Kingdom and other countries with high disease burden. The focus of the research community should be to come together to answer these complex questions as a priority.

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**Box 1: Research Questions around GBS natural course and screening**

1. What is the clinical and cost-effectiveness of universal screening (and treatment) for Group B Streptococcus using the best-available microbiological tests in reducing early onset GBS-related sepsis, mortality and morbidity, compared to current risk factor-based screening?
2. What are the medium and long-term clinical sequelae and costs of infants with early-onset and late-onset GBS disease, stratified by clinical presentation?
3. How can existing datasets and/or routine data be used to collect this information?
4. Does intrapartum antibiotic prophylaxis have an impact on the infant’s microbiome and if so what is the clinical significance of this change on short and long-term outcomes?
5. What factors affect the adoption or uptake of a) different screening and testing approaches and b) intrapartum antibiotic prophylaxis after risk-based screening?

**Box 2: Research Questions around GBS testing**

1. What factors in the mother, infant and bacterium influence the development of invasive GBS disease and how do these relate to the identification of maternal (or neonatal) colonization?
2. Can a microbiological test for GBS be developed for clinical practice to detect GBS isolates at high risk of causing invasive disease rather than colonization alone that has sufficient accuracy, and convenience (for women and the health service), and could be implemented into current healthcare?
3. Can rapid, sensitive diagnostics be developed that improve the identification of infants with invasive GBS disease?
4. What is the appropriate reference standard against which to measure any new rapid diagnostic test?

5. Could incorporation of antibiotic resistance genes into a PCR primer set enable development of a rapid test with additional information on antibiotic susceptibility in the context of penicillin allergy?

**Box 3:** **Development and testing of a vaccine for Group B Streptococcus**

1. What are the serological correlates of protective immunity against invasive GBS infection in UK women and infants?
2. What is the safety, immunogenicity and effect on colonization, in pregnant women and their infants, of a multivalent GBS vaccine given from 28 weeks of pregnancy, and what is the tolerability, acceptability and safety profile of the vaccine in the UK population?
3. What are the attitudes and knowledge of a) parents-to-be and b) healthcare professionals on antenatal vaccination for GBS and how can we learn from recent implementation of other vaccines into pregnancy (e.g. pertussis) to inform this?
4. How do these attitudes and knowledge vary across groups of different ethnicity and socio-economic status and how would we work with these groups to maximise engagement, including attendance and take-up?
5. What is the role of midwives, family physicians, obstetricians and patient support groups in ensuring good take-up?