

Multilocus Sequence Types in Czech Neonatal GBS Strains from 2004 to 2008

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Summary

The first multilocus sequence typing results of Czech neonatal *Streptococcus agalactiae* isolates are presented in this paper. The aims of the study were to prove if the Czech isolates belong to potential invasive clonal complex CC17 and to further analyse serotype – sequence type relationships.

Twelve sequence types detected among 77 isolates were assigned to 5 clonal complexes. Sequence type variability was found for most of our collection serotypes. As many as 83 % of the invasive isolates were covered by as few as 3 sequence types: S17, ST23 and ST19. ST17 was the only sequence type strictly tied to serotype III. “Hyperinvasive” ST17 was identified in 85 % of the cerebrospinal fluid isolates (6 out of 7), but in 32 % of the blood isolates only.

Key Words: *Streptococcus agalactiae* – MLST, invasive – neonatal – serotype.

Souhrn

Straková L., Musílek M., Motlová J.: MLST streptokoků skupiny B u novorozenců v České republice v letech 2004 – 2008

V článku jsou prezentovány první výsledky charakterizace 77 českých novorozeneckých kmenů *Streptococcus agalactiae* metodou multilokusové sekvenační typizace. Cílem této studie bylo ověřit, zda české izoláty patří k potenciálně invazivnímu klonálnímu komplexu CC17, a dále analyzovat vztahy mezi sérotypy a sekvenčními typy (ST). Dvanáct zjištěných ST bylo přiřazeno k 5 klonálním komplexům. U většiny sérotypů našeho souboru byla zjištěna variabilita sekvenčních typů. Potenciálně invazivní ST17 byl jediným sekvenčním typem striktně vázaným na sérotyp III. ST17, ST23 a ST19 dohromady tvořily 83 % invazivních kmenů. „Hyperinvazivní“ ST17 byl zachycen u 85 % (6 ze 7) izolátů z mozkomíšního moku, ale pouze u 32 % (7 z 22) kmenů *S. agalactiae* izolovaných z hemokultur.

Klíčová slova: *Streptococcus agalactiae* – MLST – invazivní – novorozenecký – serotyp.

Introduction

Streptococcus agalactiae (Group B Streptococcus, GBS) is a well known pathogen capable of causing both mild and severe infections. Neonatal sepsis and meningitis are among the most threatening ones. In the Czech Republic, a perinatal GBS disease preventive program, based on prenatal screening and antibiotic prophylaxis, is in place [11]. Approximately one quarter of Czech pregnant women are colonised by GBS. The level of colonisation varies depending, among others, on the cultivation method and swab site (rectal / vaginal) [12]. The percentage of women fulfilling the criteria for peripartum prophylaxis is relatively high. With

increasing antibiotic resistance in mind, there are attempts to specify more precisely the target groups for intervention.

One of the possibilities how to improve the prevention strategy is a better definition of „hypervirulent“ microbial clones. One of the internationally easily comparable methods capable of defining such clones is multilocus sequence typing (MLST). Isolates are assigned to sequence types (STs) based on partial sequencing of several housekeeping genes. STs are further grouped into clonal complexes (CC) based on the identity of some of the alleles.

The aims of the study were to prove if the Czech isolates belong to potential invasive clonal complex CC17 [3, 6] and to further analyse serotype – sequence type relationships.

Material and methods

In this study, 77 non-redundant GBS isolates were tested. Thirty-six isolates were obtained from neonates with mild disease and/or carriers; 12 from neonates with invasive disease likely to be caused by GBS (GBS was isolated from a primarily non-sterile site and other source of infection was not proved) and from 29 neonates with invasive disease. The invasive isolates were sent to the National Institute of Public Health, Prague, by clinical microbiology hospital laboratories throughout the Czech Republic from 2004 to 2008 and were obtained from primarily sterile sites, i.e. cerebrospinal fluid (CSF) and/or blood.

All isolates were serotyped using the Czech National Reference Laboratory for Streptococci and Enterococci in-house antisera and commercial latex agglutination set (Statensserum Institute, Denmark). MLST was carried out by the standard method [5] using a capillary sequencing machine (ABI Prism 3100). The sequences and allele combinations were compared with the MLST database (<http://pubmlst.org/sagalactiae> accessed on 03/08/2009). A clonal complex was defined as a maximum difference in 3 out of 7 alleles.

Discussion

The clonal variability of invasive GBS was relatively low in comparison with other pathogens (*Streptococcus pyogenes*, *Streptococcus pneumoniae*, *Neisseria meningitidis*). The Czech neonatal invasive GBS clones were similar to those studied so far in Western Europe and Northern America [4, 8, 9, 10]. Dominant representation of one serotype in one particular ST was seen (ST1-serotype V, ST19-serotype III, ST23-serotype Ia) similarly to other authors [2]. Our results confirm a „special position“ of ST17 among other STs [1]. ST17 is evolutionary distant from the other STs detected in our collection; we did not observe capsular switching (serotype change) in this particular ST. Additionally, ST17 is the most common sequence type among the Czech

Table 1: MLST clonal complexes in Czech GBS from neonates

CC (STs)	No. of invasive isolates (serotypes)	No. of susp. invasive isolates (serotypes)	No of non-invasive isolates (serotypes)	Total
CC 1 (ST1, ST7)	3 (V, IV)	5 (V, III, NT)	12 (V, Ia, II, III, VII, VIII, NT)	20
CC 10 (ST8, ST10, ST12, ST358)	0	0	6 (Ib, II, NT)	6
CC 17 (ST17)	13 (III)	4 (III)	8 (III)	25
CC 19 (ST19, ST28, ST182)	5 (III, NT)	2 (III)	6 (III, VIII, NT)	13
CC 23 (ST23, ST88)	8 (Ia, III, NT)	1 (Ia)	4 (Ia, NT)	13
Total No.	29	12	36	77

Notes: data in brackets-serotypes/sequence types detected; CC-clonal complex; NT-nontypeable; ST-sequence type

Results

Among 77 neonatal GBS isolates, 13 sequence types were assigned. Clonal complex CC1 involved ST1 (n=19) and ST7 (n=1). Parts of CC10 were ST10 (n=1), ST8 (n=2), ST12 (n=2) and ST358 (n=1); parts of CC19 were ST19 (n=10), ST28 (n=2), and ST182 (n=1). ST23 (n=12) and ST88 (n=1) were allocated to CC23. CC17 comprised isolates of a single only sequence type – ST17 (n=25). Seventeen (68 %) out of 25 ST17 neonatal isolates were invasive/potentially invasive. Overall distribution of CC among invasive and non-invasive isolates is shown in Table 1. Serotype – ST relationships are summarised in the same table (Table 1). ST17 was the only ST to be strictly tied to serotype III.

neonatal GBS isolates (45 %, 13 out of 29 invasive, vs. 22 %, 8 out of 36 non-invasive isolates). „Rapid detection“ methods based on PCR aimed at „hyperinvasive“ ST17 [7] would capture 85% (6 of 7) of our CSF isolates. Unfortunately, such a method would miss 68 % (15 of 22) of the Czech neonatal isolates from blood.

Conclusion

The Czech invasive neonatal GBS isolates have similar genomic characteristics as isolates from other countries studied so far. „Rapid detection“ methods aimed solely at “hyperinvasive” ST17 would miss 55 % (16 of 29) of our invasive isolates tested.

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