

Infected breast milk associated with late-onset and recurrent group B streptococcal infection in neonatal twins: a genetic analysis

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▶ To cite this version:

Arnaud Gagneur, Geneviève Héry-Arnaud, Séverine Croly-Labourdette, Gisèle Gremmo-Feger, Sophie Vallet, et al.. Infected breast milk associated with late-onset and recurrent group B streptococcal infection in neonatal twins: a genetic analysis. European Journal of Pediatrics, 2009, 168 (9), pp.1155-1158. 10.1007/s00431-008-0903-y. hal-00557543

HAL Id: hal-00557543 https://hal.univ-brest.fr/hal-00557543v1

Submitted on 21 Jan 2011

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- 1 Infected breast milk associated with late-onset and recurrent group B
- 2 streptococcal infection in neonatal twins: a genetic analysis
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- 24 Acknowledgements
- 25 We wish to thank Tracey Montagnon for reviewing the manuscript.
- 26

27 Summary

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Asymptomatic excretion of group B streptococcus (GBS) in breast milk may be an under-recognised cause of neonatal and recurrent infection. We report the case of late-onset and recurrent infection in newborn twins resulting from ingestion of maternal breast milk infected with GBS. Genetic analysis of isolates is equally presented.

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35 Key-words: group B streptococcus; breast milk; recurrent infection; genetic36 analysis; late-onset disease

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39 Introduction

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41 Group B streptococcus (GBS) is the most frequent cause of neonatal sepsis 42 and meningitis. Most cases occur in the first week of life and are related to 43 vaginal carriage in the mother (early-onset disease). Conversely, late-onset 44 disease (between 1 week and 3 months of age) is less common and is hand-45 transmitted by nursery personnel or via other nosocomial or community 46 pathways [6]. Late- onset and recurrent disease have also been reported with 47 the ingestion of infected mother's milk, with some cases confirmed using 48 molecular techniques [2-4, 7, 9, 13]. We report in this article the case of late-49 onset and recurrent infection in newborn twins resulting from ingestion of 50 maternal breast milk infected with GBS. In this case genetic analysis 51 demonstrated that all GBS isolates (maternal breast milk and vaginal isolates; 52 twin CSF and blood isolates) were identical, but additional genetic analysis 53 also revealed that the GBS isolates were of a particularly virulent clone 54 belonging to serotype III, as are 90% of the strains responsible for late- onset 55 disease [3].

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57 Case report

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59 Premature twins were delivered by spontaneous vaginal delivery at 31 weeks 60 gestation, 48 hours following membrane rupture. The 26 year- old mother, 61 gravida 1, para 2, received intrapartum antibiotic prophylaxis (single dose of 62 penicillin G) due to positive vaginal GBS culture at 30 weeks gestation. Twin 63 1, female, weighed 1600g with an Apgar score of 10/10 at 1 and 5 minutes. 64 No respiratory failure was noted and early enteral feeding was started with raw 65 breast milk at day 1. Total enteral alimentation with breast milk was obtained 66 at day 6. Twin 2, male, weighed 1720g with an Apgar score of 4/9 at 1 and 5

67 minutes. He was intubated shortly after birth due to respiratory failure and received one dose of surfactant. By the 3rd hour of life, he was extubated and 68 69 nasal CPAP was initiated. Enteral alimentation with raw breast milk was 70 introduced at day 4. Antibiotic therapy with cefotaxime and amoxicillin was 71 prescribed for both infants due to incomplete prepartum antibiotic prophylaxis 72 and stopped at day 2 due to negative C- reactive protein as well as negative 73 gastric and blood cultures. Cerebral ultrasound examination was normal for 74 both infants at day 4.

75 On day 13, Twin 2 developed cardio-respiratory instability and blood culture 76 tested positive for group B streptococcus. Meningitis was suspected due to 77 elevated CSF protein concentration. Until day 16 Twin 1 was asymptomatic 78 with negative C- reactive protein control. On day 16, she developed 79 respiratory distress and subsequent blood and cerebrospinal fluid cultures 80 tested positive for GBS. Antibiotic treatment with amoxicillin at 200mg/day 81 was prescribed for 14 days, in association with an aminoglycoside during the 82 first 48 hours of treatment. Control blood cultures were negative after day 1, 3 83 and 5 of treatment. Cerebral ultrasound examination controls were normal for 84 both infants. At day 41 of life Twin 1 developed septic syndrome with parotitis 85 and was transferred to the NICU. Blood culture was positive for GBS. Cardiac 86 ultrasound examination was normal. Antibiotic therapy with cefotaxime 87 (200mg/kg/day) and tobramycin (5mg/kg/day) was initiated. At day 7 of 88 infection tomodensitometry examination identified cerebral microabcess and 89 modification of the antibiotherapy ensued, with the administration of 90 ciprofloxacin (30 mg/kg/day) associated with cefotaxime (250 mg/kg/day) for 3 91 weeks. Then oral amoxicillin was initiated for 3 additional weeks. Mastitis was 92 diagnosed in the infants' mother 24 hours following discovery of GBS infection 93 in Twin 1 (day 42). Milk culture tested positive for GBS and the maternal 94 infection was treated with amoxicillin for 10 days. Breastfeeding was

95 suspended and a 10- day preventive oral amoxicillin treatment given to the 96 non- infected twin (confirmed via negative blood and CSF cultures as well as 97 negative CRP controls). Following this infection, the infants remained on 98 pasteurized breast milk. Follow- up at one year showed no cerebral anomalies 99 upon ultrasound examination in association with normal neurological 100 examinations at 1 year of life.

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102 Analysis revealed all strains as belonging to serotype III. Epidemiological 103 relationships between maternal and neonatal GBS isolates were investigated 104 by pulsed-field gel electrophoresis (PFGE) of DNA restricted with Smal [11]. 105 Analysis was conducted on maternal vaginal and raw breast milk isolates (2) 106 isolates), a single Twin 2 blood culture isolate (1 isolate), and on Twin 1 CSF 107 and first- and -second blood culture isolates (3 isolates). All six isolates 108 displayed identical PFGE patterns, revealing their genetic relationship (figure 109 1A).

110 Characterization of isolate virulence was conducted by multiplex PCR 111 according to primers and method previously described [11, 12]. First, 112 amplification of the tRNA gene clusters at the 3' end of rRNA operons 113 produced a unique fragment of 1.2 Kb (figure 1B); second, hy/B amplification 114 produced a 0.3 Kb fragment, showing no IS1548 insertion within the gene 115 (figure 1B). This pattern was correlated to the invasive phylogenetic division I 116 defined by Musser et al. in multilocus enzyme electrophoresis (MLEE) 117 analysis [8, 10].

118

119 **Discussion**

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121 Cases reporting neonatal late and recurrent group B streptococcal disease 122 associated with raw maternal milk are rare, and few are the studies in which

genetic evidence is proposed for this scenario [2, 7, 13]. In our case, not only was total DNA macrorestriction analysis conducted, showing indistinguishable patterns for the six isolates, but additional genetic analysis also revealed that the GBS isolates belonged to a particularly virulent clone shown to produce more extracellular neuraminidase [8]. These isolates, as 90% of the strains responsible for late onset disease [3], belonged to serotype III.

129

130 If the physiopathology of early onset GBS disease is well-documented, little is
131 known about late or recurrent GBS infections. This case offers a novel
132 hypothesis explaining how GBS can cause neonatal infection from 7 days to 3
133 months following delivery.

134 In most of the rare cases described [7, 9], there were no signs of maternal 135 mastitis, indicating a silent maternal duct colonization. Moreover, National 136 Committee of Hygiene guidelines do not systematically screen for GBS in the 137 raw maternal milk supply [1]. These may be two reasons for the 138 underestimation of maternal milk as a source of GBS infection. In the present 139 case, genetic analysis affords evidence for maternal milk as the source of 140 neonatal GBS infection. A circular process was hypothesized by Kotiw et al 141 [7]. GBS initially colonizes the neonate's oropharynx mucosa from perinatal or 142 other sources, infecting maternal ducts during breastfeeding. The organism 143 multiplies in the milk ducts. As the microbial concentration increases in the 144 milk, the infant is re-infected during breastfeeding. Mastitis may or may not be 145 present [7]. However Olver et al described cases of GBS infection in preterm 146 infants fed with maternal milk via nasogastric tube alone [9]. Prematurity is a 147 recognized predisposing factor to GBS infection although breast milk 148 transmission was also described in term infants [4].

In our unit, expressed mother's milk is systematically pasteurized and frozenfor conservation in our lactarium, and each specimen of milk is screened for

151 bacteria before administration to preterm infants. However, preterm infants 152 might also receive raw, freshly-expressed breast milk from their mothers 153 present in the unit. When a preterm infant falls clinically ill while the mother is 154 breastfeeding, the mother's milk should be cultured to rule out or to document 155 possible breast milk transmission. Mother's milk feedings should be 156 suspended while providing banked milk pending culture result. If breast milk is 157 positive for GBS, adequate antibiotherapy should also be prescribed for the 158 mother. Byrne et al reported that it is possible to give the mother the 159 opportunity to continue breastfeeding as desired; she can be encouraged to 160 maintain her milk supply by pumping and discarding milk until appropriate 161 treatment is completed and negative breast milk cultures are obtained [4].

162

Asymptomatic excretion of GBS in breast milk may be an under-recognised cause of neonatal and recurrent infection. Recommendations should be established to prevent these infections, notably in the case of multiple births: 1/ treatment of both twins; 2 / recognition of the possibility of GBS breast milk infection in late onset or recurrent infection; 3/ suspension of breastfeeding upon suspicion of GBS breast milk infection in both of the children; 4/ search for GBS colonization in both mother and children.

However, it should also be noted that use of human milk in the intensive care
nursery decreases the incidence of nosocomial sepsis [5] and breastfeeding
should still be considered as the most appropriate nutrition for babies and
preterm infants [1].

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175 Conflict of interest statement

176 All authors, no conflict of interest.

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Figure 1

A. Pulsed-field gel electrophoresis (PFGE) of DNA restricted with *Sma*l of maternal and neonatal GBS isolates. All six isolates displayed identical PFGE patterns, showing their genetic relationship. Molecular weight (MW) ; maternal vaginal isolate (lane 1) ; Twin 1 CSF (lane 2) ; maternal raw breast milk isolate (lane 3) ; first (lane 4) and second (lane 5) Twin 1 blood culture isolates, and Twin 2 blood culture isolate (lane 6).

228 B. Ethidium bromide stain of 2% agarose gel showing multiplex PCR products 229 for the GBS clinical isolates. Amplification of the tRNA gene clusters at the 3' 230 end of rRNA operons produced a unique fragment of 1.2 Kb; amplification 231 hy/B gene produced a 0.3 Kb fragment, showing no IS 1548 insertion within the 232 gene. This pattern was correlated to the invasive phylogenetic division I 233 defined by Musser et al. in multilocus enzyme electrophoresis (MLEE) 234 analysis [8, 12]. MW, molecular weight ; lane 1, maternal vaginal isolate ; lane 235 2, twin 1 CSF isolate ; lane 3, maternal raw breast milk isolate ; lanes 4 and 5, 236 first and second twin 1 blood culture isolates.





