Coagulase-negative *Staphylococcus hominis* is a part of the commensal bacterial microflora of healthy people. Simultaneously, these bacteria are recognized as important causes of nosocomial infections, especially in patients with indwelling prosthetic devices and in immunocompromised neonates (Chaves et al., 2005; Götz 2006; Kaufman and Fairchild, 2004; Rodhe et al., 2006). This bacterium can be responsible for bloodstream infections, sepsis, endocarditis, peritonitis, bone and joint infections (Kloos and Bannerman, 1999). Similar to other staphylococci, the formation of stable biofilm on medical devices or on host tissues is thought to be the major pathogenic factor of *S. hominis* (Chokr et al., 2006; Fredheim et al., 2009). Many studies revealed that distinct staphylococci clones can become endemic and that these clones can persist in the hospital environment (Kelly et al., 2008; Muldrew et al., 2008; Rodríguez-Aranda et al., 2009; Treviño et al., 2009). On the other hand, it has been observed that staphylococcal isolates obtained in hospitals display a wide genetic diversity (Fredheim et al., 2009; Nunes et al., 2005; Szczuka et al., 2010, Szczuka et al., 2012). This could be due to the fact that the pathogens do not represent a single evolutionary line but rather they are composed of several lines (Melless et al., 2004; Miragaia et al., 2008; Rijnders et al., 2009; Pupo et al., 1997; Szczuka and Kaznowski 2004). Although *S. hominis* is the third most commonly isolated species among CoNS, there is limited information available on the epidemiology of *S. hominis*. There is also a lack of easy-to-use and rapid methods for molecular typing of *S. hominis*. Pulsed-field gel electrophoresis (PFGE) is considered the “gold standard” for molecular typing of *Staphylococcus epidermidis* (Wang et al., 2003). In view of the above, alternative methods have been proposed. Repetitive-element PCR (rep-PCR) analysis based on multi-copy elements of the staphylococcal genome has good reproducibility and discriminatory power (Krzymińska et al., 2012) and recently rep-PCR has been commercially adapted to automated format known as the DiversiLab System (bioMérieux, France) (Treviño et al., 2009). Recently, Johansson et al. (2006) proposed multiple-locus variable-number tandem repeat analysis (MLVA) for the discrimination of clinical isolates of

**Clonal Analysis of *Staphylococcus hominis* Strains Isolated from Hospitalized Patients**

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**Abstract**

*Staphylococcus hominis* is a part of normal skin flora, but it is also a cause of nosocomial infections. The aim of this study was to investigate the genetic relatedness of 62 strains of *S. hominis* obtained from hospitalised patients during an 11-year period. For the discrimination of these clinical strains we used repetitive sequence-based PCR method (BOX-PCR) and multiple-locus variable-number tandem repeat analysis (MLVA). BOX-PCR analysis revealed a large genetic diversity among clinical strains and we did not find a predominant clone with the ability to persist in a hospital environment. MLVA is not as discriminatory as BOX fingerprinting and would not be a useful method for epidemiological studies.

**Keywords:** *Staphylococcus hominis*, epidemiological studies, BOX-PCR fingerprinting, MLVA analysis

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S. epidermidis. This method relies on the detection of different copy numbers of short repeated sequence that are arranged in tandem arrays.

The purpose of this study was to determine the genetic relationship among S. hominis strains isolated from clinical specimens from patients treated in a hospital in 2001–2012. Clones have been differentiated by repetitive sequence based PCR method (BOX-PCR) and multiple-locus variable-number tandem repeat analysis (MLVA).

Clinical isolates of S. hominis were obtained in the period 2001–2012 from clinical specimens from patients treated in different departments of a 700-bed Regional Hospital in Poznań, Poland. Strains were recovered from well-defined infections in patients. Identification of bacterial cultures was performed using the Vitek 2 system (bioMérieux, France). The reference strain of S. hominis ATCC 2122 was used in this study.

Bacteria were grown on blood agar at 37°C overnight, and DNA was extracted with the Genomic DNA Plus kit (A&A Biotechnology, Poland). The PCR reaction was performed using primer with the sequence (5’CTACGGCAAGGCGACGCTGACG3’) complementary to BOX elements of bacterial genomic DNA (Versalovic et al., 1991). The PCR was performed by methods described by Wieser and Busse (2000). The resulting fragments were electrophoresed in 1.5% agarose gel. The DNA in gels was stained with ethidium bromide, visualized on a UV light transilluminator, and documented with V.99 Bio-Print system (Vilber Lourmat, Torcy, France).

The sequence of S. hominis SK 119 was downloaded from the National Center for Biotechnology Information (NCBI) and analyzed for the presence of tandem repeat sequences with the software program Tandem Repeats Finder. The analysis identified 142 genomic regions with tandem repeat sequences and eleven regions were selected for further analysis on the basis of genomic location, repeat length, and copy number.

These tandem repeat sequences showed a minimum size of 18 bp for an individual repeat and a tandem copy number greater than two. Flanking primers were designed for the repeat loci based on the genome sequence of the SK 119 S. hominis strain. Using the Basic Local Alignment Search Tool (BLAST) we searched for the repeated sequence in other C80 S. hominis strain. The three repeated sequences were not found and for this reason these sequences were excluded from the studies. PCR was performed by the method described by Johansson et al. (2006). When performing MLVA, we noted PCR amplification failure at six loci (designated as Sh 3, Sh 4, Sh 5, Sh 7, Sh 8 and Sh 11) for many S. hominis isolates. The easiest explanation for this is that the targeted sequence is missing in these strains. PCR amplification at loci Sh 1 and Sh 2 were successful. The multiplex PCR was performed for the repeat motif TTGGAGTATTGATTATTGATT (locus Sh 1) and repeat motif TTGCTGTATTGATTATTGATT (locus Sh 2) using primers F-, GTTGCTTTGGCGTTTTTGT and R-GCTTGTAAGATGCGCTAAA for locus Sh 1 and F-TGCCTTCCTTGAGTTGAC and R-GCTTGTAAGATGCGCTAAA for locus Sh 2.

The results of the BOX and MLVA were analyzed using GelCompar II (version 3.0; Applied Maths, Kortrijk, Belgium) software. The percentages of fingerprint similarities were analyzed using the Dice coefficient. The UPGMA clustering method was used to create a dendrogram. The results of the MLVA and BOX analysis are presented in Fig. 1 and Fig. 2.

On the basis of BOX-PCR, sixty three genotypes were identified. The fingerprints of the isolates consisted of 8 to 14 bands, indicating the presence of several copies of the BOX sequence in the genomes of S. hominis strains. All strains showed a high degree of genetic diversity and had unique BOX-PCR patterns. The patterns of all S. hominis strains showed very low similarity (S = 26%). MLVA analysis allocated thirty nine isolates to 10 clusters, when a similarity level of 95% was applied. Strains grouped in these clusters were isolated from people treated in different wards of the hospital. We did not observe distribution of these strains into the clusters according to their source of isolation. Twenty four isolates showed unique MLVA patterns. The largest cluster was identified on the basis of one DNA fragment. Other clusters were designed on the basis of two or three DNA fragments. The results indicated that there was no sufficient variation in repeated regions to make them suitable candidates for sequence-based typing. It appears that strains showing the same MLVA patterns had completely different BOX banding patterns. Thus, we can conclude that MLVA is not as discriminatory as BOX analysis and can not be recommended as an epidemiological tool.

Despite the growing importance of S. hominis as a cause of nosocomial infections, there is limited information available on the epidemiology of S. hominis in the hospital setting. Our results revealed a large genetic diversity of S. hominis strains obtained from patients treated in one hospital over 11 years. We did not identify an endemic clone of S. hominis in the hospital setting with the ability to spread epidemically. Thus, patients were infected with clonally-unrelated strains.
and none of these strains persisted in the hospital environment. Previously, Chaieb et al. (2006) found that a single clone of S. hominis has been responsible for sepsis of thirteen neonates treated at a neonatology department in a Madrid hospital. In other reports, only a few S. hominis were investigated and these strains were grouped with other CoNS species using PFGE (de Silva et al., 2002; Chaieb et al., 2006; Spare et al., 2003). In our opinion, an accurate, fast and reproducible typing system would improve the epidemiological investigation of S. hominis. BOX-PCR appeared to be the more suitable method for epidemiological studies than MLVA. Strain differentiation by means of MLVA is very often based on single amplicon or two amplicons in the MLVA profiles. The result of this study demonstrated that the repeated regions chosen for MLVA analysis did not display sufficient variation for the discrimination of S. hominis isolates. BOX profiles as well as MLVA profiles did not contain a species-specific band and therefore these analyses cannot be used for identification of the S. hominis strains.

In conclusion, we observed a wide genetic diversity among S. hominis strains isolated from hospitalised patients treated in one hospital. BOX typing system is efficient in the analysis of large numbers of isolates and therefore suitable for epidemiological studies as well as for the detections of relatedness of isolates.

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Literature


