Staphylococcus Pasteuri (BCVME2) Resident In Buffalo Cervical Vaginal Mucus (CVM) - A Potential Source of Estrus-Specific Sex Pheromone(s)

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Research Article

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Abstract

There are microbes resident in the reproductive tract, some of which could be pathogenic while a few others would, perhaps, play important roles in protecting the reproductive tract from infections. Volatile compounds are known to play role as sex pheromones that attract the males for coitus during estrus or heat. It is likely that these compounds themselves are secondary metabolites of the bacterial flora resident in the vagina. In order to substantiate this hypothesis, bacteria were isolated from cervico-vaginal mucus (CVM) of buffalo during various phases of the oestrous cycle and identified, based on morphological, biochemical and molecular characteristics, as *Bacillus* during preestrus as well as diestrus, and *Staphylococcus* during all phases of the oestrous cycle. But, the populations of *Staphylococcus* differed between different phases of the oestrous cycle, the predominant forms being *S. warneri* (BCVMPE1_1) during preestrus, *S. pastueri* (BCVME2) during oestrus and *S. epidermis* (BCVMDE3) during diestrus. Mice, when used as sensors, efficiently differentiated the oestrus-specific *S. pastueri* (BCVME2) from the others. HS-GC-MS analysis showed that *S. pastueri* (BCVME2) produces key volatile compounds viz., acetic, propanoic, isobutyric, butyric, isovaleric and valeric acids. In addition, it is evidenced that *S. pasteuri* (BCVME2) volatiles influence the sexual behaviours such as flehmen and mounting of the bull. Thus, the paper reports that *S. pasteuri* (BCVME2) is the potential source of vaginal pheromone(s) during oestrus in buffalo.

Introduction

The cervico-vaginal mucus (CVM) microbial community is associated with reproductive health and fertility in mammals. For example, the cervical secretions are known to play crucial roles in the reproductive success of cows (Adnane et al. 2018). CVM of the healthy cow contains aerobic as well as anaerobic microbiota, especially those belonging to Enterobacteriaceae family, and also *Staphylococcus* sp., *Streptococcus* sp., *Enterococcus* sp. and *Lactobacillus* sp. (Otero et al. 1999; Wang et al. 2013). The CVM microbiota are influenced by hormonal, immunological and nutritional factors, and even the antibiotics and hormones that are used as drugs for treatment of the host (Otero et al. 2006). The dynamics of microbiota in the uterus and vagina have been well studied in the cows under various pathophysiological conditions (Ault et al. 2019; Bicalho et al. 2017a; Bicalho et al. 2017b; Bicalho et al. 2017c; Galvao et al. 2019). In buffalo, association of CVM microbiota with haematological parameters, such as counts of the white blood cells (WBC) lymphocytes, monocytes, granulocytes, etc., during the different phases of oestrous cycle has been reported (Joshi et al. 2020). The role of CVM bacteria in reproduction and pheromone releases in bovine has been reviewed in adequate detail (Srinivasan et al. 2021). However, elaborate investigation on estrus-specific bacteria in buffalo pheromone has not yet been taken up.

Research so far has revealed that the culture-independent method using next-generation sequencing (NGS) is appropriate to find the diversity of CVM microbiome with particular attention to the buffalo oestrous cycle (Srinivasan et al. 2019). Proteobacteria, Actinobacteria, Firmicutes, Bacteroidetes and Tenericutes are the most common CVM of buffalo. Among these five families, Firmicutes has been found
to be highly abundant during oestrus compared to the other phases (Srinivasan et al. 2019). The volatile chemical compounds of CVM can vary among the phases of oestrous cycle, and it is well known that bulls are able to differentiate oestrus and non-oestrus cows based on pheromone(s) present in the vaginal secretion (Karthikeyan and Archunan 2013; Rajanarayanan and Archunan 2004; Sankar and Archunan 2004).

It has been documented that male dogs and mice differentiate oestrus and non-oestrus cows by sniffing the odour of vaginal fluid, urine, saliva, faeces and milk of the latter (Fischer-Tenhagen et al. 2013; Rameshkumar et al. 2008; Sankar and Archunan 2005). It has been established that bovine bulls exhibit flehmen and mounting behaviours after perceiving unique volatiles from females during oestrus (Karthikeyan et al. 2013; Rajanarayanan and Archunan 2004; Sankar and Archunan 2004). The flehmen is demonstration of the most distinct and important aspect of the premating behaviour. The flehmen behaviour of bulls has been used to identify and quantify pheromone production in buffaloes and cows (Karthikeyan et al. 2013; Rajanarayanan and Archunan 2004; Sankar and Archunan 2004).

Volatile compounds such as acetic acid and propanoic acid have been identified in cow vaginal mucus (Sankar and Archunan 2011) and oleic acid from buffalo vaginal secretion (Karthikeyan and Archunan 2013) during oestrus, which act as female sex pheromones. Interestingly, acetic acid has been found in buffalo urine, with a substantial increase in concentration during estrus relative to other phases of the oestrous cycle (Muniasamy 2016). The bacterial populations present in the CVM are reported to potentially affect the sexual attraction of ewes (Ungerfeld and Silva 2005). The report suggests that fatty acids may act as pheromones as well as estrus indicators in buffaloes and that these microbes might be responsible for the specific odour released at oestrus.

On the basis of substantial evidences, buffalo CVM has been shown to produce estrus-specific volatiles that affect the reproductive behaviour of male buffalo during mating. However, the source of the buffalo CVM pheromone compounds is not well established. Evidence for the impact of CVM bacteria and their role in production of volatiles in the context of male attraction towards female buffalo is inadequate. The present study was therefore intended to i) demonstrate the bacterial community in buffalo CVM through a culture-based method during different phases of the oestrous cycle, and (ii) screen the oestrus-specific bacteria using mice as model sensor system, iii) find if bacterial secretory substance contains volatiles, and iv) evaluate bull behaviour in response to estrus-specific bacterial secretory substance of CVM in order to confirm that the very bacterial volatiles are the pheromones that elicit specific premating behaviours in the males.

### Materials And Methods

#### Animal and sample collection

The healthy female buffaloes (n = 12) and phases of oestrous cycle were monitored as per the protocol of Rajanarayanan and Archunan (2004). CVM collection was carried out based on the procedure and
Isolation and characterization of bacteria

For the isolation of bacteria, the CVM was collected from buffaloes representing the three phases of the oestrous cycle. CVM, 0.5 mL, was homogenized in 4.5 mL of sterile distilled water. An aliquot of 0.1 mL of the homogenate was serially diluted and plated in Nutrient-MacConkey-De Man, Rogosa and Sharpe- and Mannitol salt- agar by standard plate count method (Cassoli et al. 2016). Plates were incubated at 37°C for 24–48 hr. Following the incubation, the bacterial colonies were counted and expressed by log values (Log CFU/mL i.e. Log10 (CFU / (dilution factor x aliquot). Morphologically distinct colonies alone were picked and further processed. Primary characterization was carried out based on colony morphology which included colour, size and shape. Further, the isolates were subjected to Gram staining and biochemical tests for catalase, oxidase, sugar fermentation and haemolysis.

Extraction of genomic DNA and PCR amplification of 16S rRNA gene

Genomic DNA of bacterial isolates was extracted based on the method of Vingataramin and Frost (2015). The quality of gDNA was evaluated on 1.0% agarose gel. Fragment of 16S rRNA gene was amplified by PCR with the primers 27F AGAGTTTGATCMTGGCTCAG and 1492R CGGTTACCTTGTTACGACTT. PCR reactions were executed in a total volume of 30 µL containing 15 µL 2x PCR premix, 2 µL template DNA, 0.5 µL each of the primer and 12 µL sterile distilled water. The cycling parameters used were as follows: initial denaturation at 94°C for 3 min, followed by 35 cycles of denaturation at 94°C for 30 sec, annealing at 50°C for 30 sec and extension at 72°C for 90 sec, and then a final elongation at 72°C for 7 min (Coombs and Franco, 2003). Eurofins Genomics India Pvt. Ltd (Bangalore, India) conducted the 16S rRNA gene sequencing.

Sequencing and phylogenetic analysis of 16S rRNA gene

The amplicons were purified using pre-sequencing kit and sequenced on ABI 3730xl automated sequencer (Perkin Elmer Applied Biosystems, Foster City, CA, USA) and ABI Prism BigDye Terminator Cycle Sequencing Ready Reaction Kit when AmpliTaq® DNA polymerase was used as per protocols recommended by the manufacturer.

The dataset of 16S rRNA gene sequences was prepared for BLAST search. All individual representative sequences were performed at NCBI (Altschul et al. 1990). Sequences were obtained from BLAST searches based on E-value where identity scores differed depending on the size of the gene family. Nucleotide compositional parameters were predicted from BioEdit 7.0.4 (Hall 2011). Besides the BLAST results, 16S rRNA gene sequences of the most closely related taxa were retrieved from the GenBank database and
aligned using the ClustalW tool implemented in MEGA X (Kumar et al. 2018). MEGA X was used to reconstruct the phylogenetic tree adopting neighbour-joining (NJ) method using bootstrap values based on 1000 replications (Kim et al. 2019). 16S rRNA gene sequences were deposited in NCBI database using BankIt (http://www.ncbi.nlm.nih.gov).

**Isolation of volatiles from bacterial culture**

The selected bacterial colonies from CVM collected during various phases of oestrous cycle (i.e. preestrus, estrus and diestrus) were inoculated in 250 mL of LB (Luria Bertani) broth and incubated in rotary shaker at 150 rpm for 48 hr. Further, the culture was centrifuged at 5000 x g for 10 min (Tredwell et al. 2011). The supernatant was subjected to further studies.

**Discrimination of bacterial volatiles using mice**

Screening of bacteria odour was performed using mice as the odour detector (Sankar and Archunan 2005). Twenty four male mice (12 week old) were used for the odour preference and the study was performed by Y-maze apparatus made of tin sheets and glass plates (Sankar and Archunan 2005). The behaviour study was conducted according to the standard protocol (Achiraman and Archunan, 2006). The bacterial supernatants (100 µL) were presented in 2 mL vials having minute pores and mice were introduced into the central chamber to begin in a common place, and the duration and number of visits were observed for 15 min. Three mice were used as test animals for each sample. Use of mice in this experiment was approved by the Institutional Ethics Committee (IAEC) of Bharathidasan University (BDU/IAEC/2017/NE/13Dt.21.03.2017). Ultimately, the potential odour-producing bacterial isolates were selected and taken to further HS-GC-MS analysis.

**Identification of bacterial volatiles adopting headspace gas chromatography-mass spectrometry (HS-GC/MS)**

The supernatant (1 mL) collected from *S. pasteuri* culture was subjected to HS analysis. HS-GC/MS analysis was performed on an Agilent mass spectrometer (Agilent Technologies, DE, USA) coupled with GC system (5977 MSD, 7679A Headspace Sampler, Agilent Technologies). Bacterial volatiles were separated by capillary GC column DB-VRX (20 m x 180 µm x 1 µm). Helium was used as the carrier gas at a flow rate of 1 mL/min. The chromatographic conditions for the analysis was as follows: a) initial temperature 40°C, for 3 min, b) 150°C for 1 min at a rate of 7°C/min, and c) 240°C at a rate of 60°C. Samples (1µL) were injected in split mode (30 sec) and the split ratio was 10:1. The separated compounds were ionized by electron energy at 70 eV in positive ion mode. Peak identification was done based on mass spectral interpretation and comparisons using NIST National Institute of Standard and Technology (NIST) library (Boots et al. 2014).
Bull behavioural observation towards CVM bacterial volatile samples

Non-estrus female buffaloes (n = 6) and bulls (n = 6) were selected for the behaviour studies. The supernatant of bacterial culture, estrus CVM, non-estrus CVM and LB broth were subjected to bull behaviour analysis. Each sample (5 mL) was individually soaked in cotton and rubbed on the vulvar region of non-estrus buffalo (i.e. dummy cow). The bulls were allowed to sniff the test animals, and behaviours were observed for 30 min (Karthikeyan and Archunan 2013). The flehmen and mounting behaviours exhibited by the bull in response to the test samples were recorded.

Statistical analysis

Analysis of variance (ANOVA) was carried out for behaviour data analysis by using SPSS (V.2.3) package, and images were plotted in GraphPad Prism (V.8.0.2).

Results

Isolation of bacteria from buffalo CVM

The bacteria isolated from CVM during preestrus, estrus and diestrus were present as 38 CFU/mL, 51 CFU/mL and 36 CFU/mL, respectively as determined adopting serial dilution method. Thus, cumulatively as many as 125 bacterial colonies were observed covering all three phases. Amongst them, 6 isolates were chosen, based on morphological characters such as smoothness and colour (white or creamy yellow), for higher level analyses. Biochemical characterization such as Gram staining, sugar fermentation tests (glucose, fructose, maltose and sucrose) and haemolytic assay showed (Table S1). The bacteria found in oestrus CVM are Gram positive, non-haemolytic, and fermentable sugars, including glucose, fructose, maltose, and sucrose, with acid production.

Molecular characterization of bacterial isolates showed that expression of Staphylococcus community differed between the three phases of the oestrous cycle, as follows: *Staphylococcus warneri* (BCVMPE1_1) during preestrus, *S. pasteuri* (BCVME2) during estrus and *S. epidermidis* (BCVMDE3) during diestrus. With regard to Bacillus resident in CVM, *B. subtilis* (BCVMPE1) and *B. toyonensis* (BCVMPE2) were limited to preestrus, whereas *B. proteolyticus* (BCVMDE5) showed up only during diestrus. The sequences have been deposited at NCBI under the accession number: MT789231.1 (BCVMPE1_1), MT705012.1 (BCVMPE2), MT598012.1 (BCVMPE1), MT598013.1 (BCVMDE3), MT598014.1 (BCVMDE3), and MT705014.1 (BCVMDE5).

Construction of the phylogenetic tree
Pairwise 16S rRNA gene sequence similarities were determined using BLASTn analysis. Based on the identity (98% and above from the query coverage) the species was selected. The variations in nucleotide composition were identified for CVM bacterial isolates (in percentage as well as numbers). G + C base pair was found to be high, above 52% (Table S2). NCBI-BLAST was practiced to identify isolates having > 98% similar sequence at inter- and intra-specific levels. The result showed that the tree could be grouped into two major clusters. The first cluster consists of various species of *Staphylococcus* genus; the second cluster consists of *Bacillus* species as represented in phylogenetic tree analysis (Fig. 1A). The molecular taxonomic characterization showing the differential expression in respect to three phases of oestrous cycle is presented in Fig. 1B.

**Discrimination of the bacterial volatiles from CVM adopting mice behavioural assay**

There were significant differences among the bacterial volatiles across different phases of the oestrous cycle. The behavioural assessment of the mice revealed greater attraction towards estrus-specific bacteria than the non-estrus bacterial sample (Fig. 2). The estrus-specific bacterial volatiles was found to be most attractive as mice visited them frequently and spent more time than the non-estrus bacterial volatiles ($p < 0.001$). The behavioural study further expounded that the estrus *S. pasteuri* (BCVME2) most attracted the mice.

**Identification of volatiles produced by bacteria as revealed in headspace GC-MS**

The GC-MS analysis of cell free supernatant of estrus-specific bacteria isolated from CVM revealed the nature of volatiles (Fig. 3). Six volatiles *viz.*, acetic acid, propanoic acid, isobutyric acid, butyric acid, isovaleric acid and valeric acid are produced by *S. pasteuri* (Table 1). Acetic acid appeared to be at a higher peak intensity among bacterial volatiles, followed by isovaleric acid, valeric acid, isobutyric acid, butyric acid, and propanoic acid.

Table 1: Volatile compounds of *S. pasteuri* (BCVME2) isolated from buffalo CVM.
Bull behaviour conditioned by the volatiles of *S. pasteuri*

The bulls were attracted to the maximum towards intact estrus CVM followed by supernatant of *S. pasteuri*, LB broth, and non-estrus CVM. The frequency of flehmen response exhibited by bull was highest to the intact estrus CVM (5.83 ± 1.16) as well as *S. pasteuri* secretory substance (4.66 ± 1.21). By contrast, bull showed significantly lesser attraction to non-estrus CVM (0.67 ± 1.03) and no response at all to LB
broth. Likewise, the mounting behaviour exhibited by bull was remarkably higher towards intact estrus CVM \( (4.83 \pm 1.94) \) and \( S. \text{pasteuri} \) secretory substance \( (4.5 \pm 1.64) \). However, bull showed less response towards non-estrus CVM \( (0.5 \pm 1.22) \) and no response at all towards LB broth. However, the flehmen and mounting behaviours observed during exposure to estrus CVM sample and \( S. \text{pasteuri} \) secretory substance are almost similar in response. Details of flehmen and mounting behaviours of bull are given in Fig. 4.

**Discussion**

Our previous study showed differences in the CVM bacterial community with respect to various phases of the oestrous cycle (Srinivasan et al. 2019), in which the Firmicutes phylum was more abundant during the oestrus phase. Probably, expression of certain specific bacterial genes is higher during estrus phase than the other phases. The present investigation revealed expression of specific gene(s) in \( S. \text{pasteuri} \) during the estrus phase. The available information indicates that the CVM bacterial strains get influenced during estrus in buffalo. It has also been reported that the vaginal bacterial population is affected by the cyclical variation during the oestrous cycle (Otero et al. 1999; Otero et al. 2000).

In the present study, \( S. \text{pasteuri} \) were incidentally observed in buffalo CVM during estrus. Similar studies have been reported in various animals. Association of vaginal \( \text{Simonsiella} \) spp in reference to estrus phase has been documented in lions and leopards (Callealta et al. 2018). It is interesting to note that \( \text{Streptococcus} \) and \( \text{Staphylococcus} \) are the most dominant bacteria in healthy cows during the reproductive cycle (Amin et al. 1996). \( \text{Staphylococcus} \) (70%) and \( \text{Escherichia} \) (15%) were observed more frequently during natural and induced estrus of ewes (Orihuela et al. 2019) and all the isolates had a high sensitivity to ciprofloxacin antimicrobials that evaluated vaginitis regulation (Mohammed et al. 2017). The present finding supports the previous report of the presence of \( \text{Staphylococcus pasteuri} \), which comes under the phylum Firmicutes, and is abundant during estrus (Srinivasan et al. 2019).

In addition, bacteria such as \( \text{Bacillus} \) spp., \( \text{Staphylococcus} \) spp., and \( \text{Streptococcus} \) spp., have been identified in vagina of ewe (Manes et al. 2010) as well as cows (Otero et al. 1999; Otero et al. 2000; Zambrano-Nava et al. 2011). These reports match with the present finding of bacterial community identified in CVM of buffalo. On the other hand, major bacterial genera in the vagina of Gyr and Nellore breed cows include \( \text{Aeribacillus}, \text{Bacillus}, \text{Clostridium}, \text{Bacteroides} \) and \( \text{Ruminococcus} \) (Giannattasio-Ferraz et al. 2019). The present analysis also depicted the presence of \( \text{Bacillus} \) spp. in buffalo exclusively during preestrus and diestrus phases. Notably, \( \text{Lactobacilli} \) were found to be very low in bovine vaginal microbiota which was confirmed by culture-dependent and culture-independent methods (Otero et al. 2000; Quereda et al. 2020; Srinivasan et al. 2019). The present finding reveals the absence of \( \text{Lactobacillus} \) sp. in buffalo CVM. Overall, the results obtained in the present study are consistent with previous reports.

The mice system was used to screen the volatile compound samples. Among 7 samples tested with mice, \( S. \text{pasteuri} \) secretory substance exhibited to be more attractive than the other cultures. The results are
consistent with an earlier report that mice are capable of discriminating the estrus and non-estrus samples. A recent study provides supportive evidence that mice are highly sensitive to volatiles produced by the microbes (Peixoto et al. 2018). Truly, the mice study helped to pick out the bacteria potentially responsible for volatiles production.

GC-MS analysis indicated six volatiles, acetic acid, propanoic acid, isobutyric acid, butyric acid, isovaleric acid and valeric acid, as produced by *S. pasteuri*. It is known that short-chain fatty acids such as acetic-, propanoic- and butyric acid, are relatively abundant due to bacterial fermentation in colonic lumen which regulate entero-pathogenic colonization by host immunomodulation in human (Smith et al. 2013; Wrigley 2004). A recent study shows that acetic, valeric, caproic and myristoleic acids are at significantly higher levels in the milk of estrus cow (Zebari et al. 2019). The fatty acids, acetic and propanoic, have been identified in the faeces of cow, and varied in relation to the time of ovulation (Mozūraitis et al. 2017). The present investigation supports the previous reports of presence of acetic, propanoic and valeric acids as produced by CVM bacteria in estrus buffalo.

Since estrus-specific *S. pasteuri* produces volatiles, we focused on testing the role of *S. pasteuri* secretory substance in bull behaviours which confirmed the estrus-specificity and its use as a reliable estrus indicator. Interestingly, bull exhibited reproductive behaviours such as flehmen and mounting, which are important in assessing sexual desire of bull as response to *S. pasteuri* secretory substance. The present study also corroborates previous findings that the estrus-specific faeces pheromone has a significant effect on flehmen and mounting behaviour in buffalo (Karthikeyan et al. 2013). It has been reported that bulls exhibit this reproductive behavioural patterns in response to the presence of a mixture of volatiles in the CVM of estrus cow (Karthikeyan and Archunan 2013; Rajanarayanan and Archunan 2004; Sankar and Archunan 2004). Vaginal pheromones have been reported to be acetic acid, propanoic acid, and 1-iodoundane, in cows (Sankar and Archunan 2004) and oleic acid as an estrus indicator in buffalo (Karthikeyan and Archunan 2013) to stimulate flehmen and mounting behaviour response in bulls. The present findings suggest that the volatile compounds of bacterial origin may act as attractants of the bull and induce the olfactory system to elicit flehmen and mounting behaviours in order to maintain the health and management during reproduction. Since *S. pasteuri* is capable of secreting the volatiles viz., acetic, propanoic, isobutyric, butyric, isovaleric, and valeric acids, they can be considered as major bacterial volatiles that induce the buffalo bull’s sexual behaviour. For instance, acetic acid, 2-butanone and oleic acid have been shown to improve Zebu bull sexual behaviour and total sperm production (Mondal et al., 2019). Thus, this study reports the presence in the buffalo vaginal fluid of volatile compounds secreted by CVM bacteria which facilitate the expression of male reproductive behaviours, which in turn suggests that these volatiles are putative pheromone compounds produced during oestrus in buffalo.

Further studies are needed to find the mechanisms associated with the specific effects of *S. pasteuri* volatiles. Characterization of protein profile of *S. pasteuri* from buffalo CVM during estrus is required to understand the host-bacterial interaction and their role in chemo-signal communication that would enlighten the reproduction management in buffaloes.
Declarations

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The authors declare no conflicts of interest.

Availability of data and materials

Not applicable

Code availability

Not applicable

Authors' contributions

MS: Conceptualization, Investigation, Writing original manuscript. RLR: Data interpretation. DD: Conceptualization, Contributed to isolation of bacteria and culture methods. MAA: review & editing, GA: Conceptualization, Supervision, review & editing.

Ethics approval

Not applicable

Consent to participate

Not applicable
Consent for publication

All authors are agreed for the article communication and publication.

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**Figures**

**Figure 1**

The phylogenetic tree. A. The inter- and intra-specific evolutionary relationship of the strains isolated from CVM. B. Highlight of the specific expression of bacterial isolates in CVM during the estrus phase.
Figure 2

Ability of mice to differentiate the CVM bacteria by sniffing bacterial volatile samples. Bars show the number of visits by mice towards bacterial volatile from buffalo CVM.
Figure 3

Chromatogram of bacterial secretory substance. A. S. pasteuri and B. LB broth.
Figure 4

Behavioural assay. Bull behaviour towards the volatile sources. n=6; Values are represented as Mean±SEM. Bars having same letters indicate no significance.

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