Cultivation, Isolation and Characterization of Bacteriocin from Fresh Cow Milk and Meat Samples obtained from Lapai Market in Niger State Nigeria.

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ABSTRACT: This study focus on cultivation, isolation and characterization of Bacteriocin from fresh cow milk (FCM) and fresh cow meat (FMS) samples obtained from Lapai Market in Niger State, Nigeria. Potential bacteriocinogenic bacteria were screened with agar diffusion method on culture plates seeded with Staphylococcus and E.coli. Bacteriocinogenic isolates from FCM were Enterococcus faecium FCM5 and Bacillus megaterium FCM6, while isolates from FMS include Bacillus subtilis FMS1 and Micrococcus varians FMS2. Bacteriocins produced were assayed with pre-enriched MRS broth culture incubated at 30°C with adjusted pH (7.1) for 48 hrs and stored (4°C). The primed cultures were centrifuged (20,000rpm) for 30mins, supernatant were filtered (0.2µm membrane filters), precipitated with ammonium sulphate and steam sterilized (121°C at 15psi) for 15mins. Bacteriocin-like products (BLPs) were crystallized and tagged enterocin FCM5, bacillocin FCM6, bacillocin FMS1 and micrococin FMS2 according to bacteriocinogenic isolates' typing. After 48 hrs, Bacteriocinogenic load ranged between 5.1 - 6.13Log10 cfu/mL. Biopreservative indexes (BI) of BLPs (0.25 - 0.75 mg/mL) were effective against predominant food spoilers (Saccharomyces, Pseudomonas, Klebsiella and Micrococcus spp isolated from range of beverages, starchy and meaty meals). Quantified BLPs (20 - 1880 IU/mL) strengths had no direct correlation with the antibacterial spectral (6 - 19mm) expressed as zone of inhibition. Thus, efficacy of BLPs observed in this study, were not a function of BLP quantity but quality. Conclusively, Enterocin FCM5, Bacilliocins (FCM6 and FMS1) and Micrococin FMS2 are thermal and pressure stable BLPs that are effective against predominant food spoilers. ©JASEM

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Bacteriocins are proteinous metabolites of bacteria that have inhibitory activities against some related or non-related bacteria (Zhou et al., 2014). However, non-bacterial species such as fungi (Adebayo and Aderiye, 2010) are susceptible to bacteriocin. Popular among them are nisin, pediocin, colicin, aureocin, agrocin, alveicin, lactococin, divercin, subtilin, leucocin, lacticin, cuvaaticin, triflinoxin, halocin, thuricin etc.

Prominent lactic acid bacteria (LAB) that synthesizes bacteriocin-like products include pediococci, lactococci, lactobacilli, and enterococci. Bacteriocin-like products in combination with hydrogen peroxides, lactic acid and other organic acids are metabolites of LABs that naturally preserve foods from spoilage microbes. Most dominant LABs in fermented and non-fermented foods are good producers of bacteriocins. LABs that synthesize bacteriocins have been isolated from different food types and beverages such as beers, milk, fermented cucumber, cheese (Rekhef et al., 1995 Moham et al., 2012, Gonzales et al., 1994; Van Reenen et al., 1998) and fermented vegetables (Zhou et al.2014). The nutritive nature of meat products makes them a good medium that support broad spectrum of spoilage and benefious bacteria. LABs with bacteriocin-like metabolites have been isolated from meat products (Bromberg et al. 2004).

Global food safety agencies envisaged a world of healthy foods with natural food preservatives, which are generally regarded as safe (GRAS). Bacteriocins (Nisin) status as GRAS (Mohammed et al., 2016), have increased detailed studies on the potency and application of other bacteriocin as natural alternative substance for food preservations. Successful antimicrobial potencies of bacteriocins have been recorded (Mohammed et al., 2016; Gao et al. 2010). Most bacteriocins are progressively studied either as crude extract form, semi purified or purified form via culture sediments, precipitation (ammonium sulfate,
methanol extraction) and chromatographic activity. Considering the health risk associated chemical preservatives, there is a growing advocacy for natural or organic food preservatives. Thus, this study aimed to cultivate, isolate and assess bacteriocin from fresh cow milk and meat samples obtained from Lapai Market in Niger State, Nigeria.

**MATERIALS AND METHOD**

**Sample Collection:** A total of 1000 mL and 1000g of Fresh cow milk (FCM) and fresh meat samples (FMS) respectively were purchased from five different vendors at random locations in Lapai Market Niger state, Nigeria. Approximately from each vendor, 200 mL FCM and 200g of FMS were purchased and stored in sterile bottle and plastic bags respectively placed in ice bags <4°C while on transit to the laboratory (IBB university Lapai). Each of the five samples (FCM and FMS) were bulked together and labelled as FCM and FMS. Leftover foods from IBB University, cafeterias were aseptically collected in sterile plastic bags, and transferred to laboratory for isolation of test/indicators organisms.

**Microbial Isolation:** Standard methods of culture media preparation were done according to methods of Cowan and steel (2004). From the bulked sample, 100 mL of FCM was diluted in 900 mL of distilled water and vigorously agitated for 1 minute. With sterile pipette, 1 mL of the solution was diluted serially with 0.1% peptone water up to the 3rd fold and 1mL of each sample were pour plated on lactic acid Bacteria medium (LABM), nutrient agar (oxoid) and De Man Rogosa sharpe (MRS) broth (Oxoid) incubated at 30°C for 24 hours. Similar protocols were replicated using 100g of chopped (~2cm³) FMS steeped in distilled water overnight. Colonies were enumerated and expressed as Log10 CFU/mL. Pure colonies were obtained by subculturing isolates on fresh media and stored in slant agars prior to characterization using relevant biochemical test (Cowan and Steel, 2004).

**Test/Indicator Organism Protocol:** *Staphylococcus aureus* and *E. coli* were selected as indicator organisms to screen prospective bacteriocin producing isolates. Prominent strains from food samples were selected as the test organisms. Among the selected species, include *Saccharomyces cerevisiae* from beverages, *Klebsiella* sp (meat sample), *Micrococcus varians* (meat) and *Pseudomonas* sp (variety of carbohydrate solid foods).

**Bacteriocin Assay:** Screening and cultivation of Bacteriocin: Prospective bacteriocin synthesizing pure isolates of FCM and FMS were initially screened using agar well diffusion methods. Aliquots (50uL) of 48hr old pure cultures of FCM and FMS isolates were discharged into wells (7-mm diameter) of fresh nutrient agar culture plates seeded with either *Staphylococcus aureus* or *E. coli* as indicator organisms. Isolates with zone of inhibitions were designated as positive or otherwise negative. Only positive designated isolates were cultivated in 250mL of pre-enriched MRS broth culture for 48 hours at 30°C with adjusted pH (7.1) and stored at 4°C prior crude extract purification and quantification assays.

**Crude Extraction, semi purification and quantification of Bacteriocin:** Protocols for crude extract of bacteriocin were sequential by centrifuging primed culture (48 hr old) at 20,000 rpm (≤4°C) for 30 mins. Supernatant of solution were adjusted to pH of 7.1, before filter sterilization using whiteman® membrane nylon filter (0.2µm) and 5mg/ml catalase (C-100 bovine liver, Sigma) was added to eliminated peroxides and lactic acids effect (Kacem et al. 2005). The solution was tagged as bacteriocin crude extract (BCE) and used for bacteriocin antibacterial assays. BCEs were further precipitated with ammonium sulphate, steam sterilized (121°C at 15 psi) for 15minutes and tagged as bacteriocin-like product (BLP). Quantification of BLPs were determined at spectrophotometric wavelength of 450nm and expressed as IU/mL equivalent of Nisin(Sigma, USA) standard solution.

**Nisin Standardization Protocol:** Nisin Standard was prepared according the methods of Papagianni et al. (2006) by adding 0.1g of nisin to solution (10 ml 0.02 N HCl and 0.75% NaCl) and steam sterilized at 121°C for 15minutes at 15psi and tagged standard nisin solution(10²UI/ml). From this standard solution, 0.2, 0.4, 0.6, 0.8 and 1.0 concentrations with index factor of 10³ IU/mL were prepared and used to plot standard nisin concentration-absorbance(@450nm) chart with $R^2 = 0.959$.

**Antibacterial assay:** The antibacterial activity of the BCEs were determined using agar well diffusion method described by Mohammed et al., (2016) with slight modifications. From each BCEs, 10mL was collected and labelled as 1.0 Au/mL, and three sequential dilutions (0.75 AU/mL, 0.5 AU/mL, 0.25 AU/mL) were made from it. From each concentration, 500 µL of each were drawn and discharged into 5-mm diameter wells in bioassay agar plates (with 5 wells per plate: 4 sample and 1control) seeded with specific spectrum of test organisms.
Distilled water was used as control on each plate. All plates were incubated at 37°C for 24 h and zone of inhibitions diameters were measured. Minimum inhibitory concentration (MIC) was determined using the least concentration of BCEs that induced Zone of inhibition and expressed as arbitrary units (AU) per ml(AU/mL). Where 1 AU/mL = dilution factor of standardized solution of BLPs.

![Graph showing bacterial load of BLPs Isolates from FCM and FMS](image1)

**KEY:** FCM = Fresh Cow Milk, FMS = Fresh Meat Sample, BLPs = Bacteriocin-like-products

**RESULTS AND DISCUSSION**

Bacteriocinogenic load cultivated for 48hrs at 30°C with adjusted pH (7.1) were relatively stable at the range of 5.1 Log cfu/mL - 6.13Log cfu/mL (Fig. 1). Irrespective of the source of Bacteriocinogenic isolates, there were no significantly different (P<0.05) among bacterial loads. Perhaps, similarity of the culture conditions was responsible for the stable bacterial kinetics. The ranges of bacterial loads were within the findings of Malheiros et al., (2015), which documented a range of 5.62 - 9.26 log cfu/mL. However, it seems that bacterial load have neither direct correlation with antibacterial spectral nor BLPs quantities within 48 hrs cultivation period.

Calibration curve for BLPs quantification was obtained using plot of absorbance value against concentration (UI/mL) of Nisin standard solution. The curve was resolved as linear model (Y = 0.182X) with coefficient (R²) of 0.959 (Fig. 2). The Chart was validated based on more than 95% confidence (R² = 0.959). Malheiros et al., (2015) used fitted models and coefficients (R² values) for bacteriocin assays in their attempt to optimize the growth of bacteriocin production by *Lactobacillus* subsp. *sakei* 2a.
Similarly, Venigalla et al., (2017) validated the precision of quadratic models for bacteriocin assay based on the R² values (0.911), which was significantly lower than the R² (0.959) observed in this study. Perhaps, the purity level of the standardized Nisin (sigma USA) used in this study, accounted for the disparity compared to the crude bacteriocin of Venigalla et al., (2017).

Minimum inhibitory concentrations (MIC) were assessed and expressed as biopreservative index (BI) of bacteriocins (FCM and FMS). BLPs of FCM6 and FMS1 with BI of 0.25mg/mL each, were adjudged as better biopreservatives than FCM5 (0.75mg/mL) and FMS1 (0.50mg/mL). Test organisms to assay the BI factor, bacteriocins of bacillocins (FCM6 and FMS1) would be efficient to preserve foods against bacterial spoilers of non-beverage than micrococin FMS 2, and enterococs FCM5 would be best to preserve beverages.

| BLPs          | MIC/BI (mg/mL) | Test organism (predominate isolate from source) | Source of indicator organism |
|---------------|----------------|-----------------------------------------------|------------------------------|
| Enterocin FCM5| 0.75            | *Saccharomyces* sp                        | Beverages                   |
| Bacillocin FCM6| 0.25            | *Pseudomonas* sp                             | Starchy food                |
| Bacillocin FMS1| 0.25            | *Klebsiella* sp                              | Meaty foods                |
| Micrococin FMS2| 0.50            | *Micrococcus* sp                             | Meaty foods                |

KEY: FCM = Fresh Cow Milk, FMS = Fresh Meat Sample, BI = Biopreservative Index, BLP = Bacteriocin-Like Products, MIC = Minimum Inhibitory Concentrate.

Assessing the quantity of BLPs after 48 hrs at 30°C cultivation, showed that Micrococin FMS2 (1.88 IU/mL) was significantly high compared to decreasing order of Enterocin FCM5 (0.23 IU/mL), BacillocinFCM6 (0.13 IU/mL), and Bacillocin FMS1 with 0.02 IU/mL (Fig. 3). The cultivation conditions (pH 7.1 and 30°C) seem to be optimal for Micrococin FMS2 synthesis.

However, Papagianni et al., (2006), correlated the bacteriocin concentration and antibacterial assay (zone of inhibition). Their models seem to be species sensitive, as only one among three species of *Lactobacillus* plot was correletively strong. Delgado et al. (2005) also demonstrated that quotients of linear parallel models of zone of inhibition and indicator organisms' sensitivity are capable of quantifying bacteriocin activity. Both references above utilized relativity principles to quantity...
bacteriocins, which are similar to the scope of this study that used absorbance relativity values expressed as equivalent of Nisin.  Bacillocins (FCM6 and FMS1) have the highest zone of inhibitions of 18 mm and 19 mm respectively when 1ml (~10^2 UI/mL equivalent of Nisin) was tested against predominant bacterial food spoilers (Pseudomonas and Klebsiella Spp). In declining trend, microcin FMS2 (11mm) was effective against related species (Micrococcus spp) and the least was enterocin FCM5 with 6mm zone of inhibition against Saccharomyces spp (Fig. 4). Yang et al., (2012) observed that BLPs of lactobacillus species inhibited related species.

Similarly in this study, microcin FMS2 inhibited Micrococcus sp. Previous studies of Mohammed et al. (2016); Yang et al., (2012) Papagianni et al. (2006), Sobrino-Lopez and Martín-Belloso (2008) and as well this study, observed inhibition of non-related species by BLPs. Obvious stability and antimicrobial activities of BLPs after treatments (121°C at 15psi), confers heat and pressure stable status to BLPs of FCMs and FMSs. Some of the gram negative pressure-resistant bacteria such Pseudomonas and E. coli (Black et al., 2005), were effectively inhibited by BLPs of this study. Invariably, they could be possibly used in hurdle technology for preservation of pressure treated foods. Conclusively, BLPs of Enterocin FCM5, Bacilliocins (FCM6 and FMS1) and Micrococin FMS2 are thermal and pressure stable BLPs that are effective against predominant food spoilers.

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