Investigation of Effect of Ripening and Processing on Prebiotic Potential of Banana

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ABSTRACT
Objective: Daily consumption of raw banana improves human health because of its resistant starch. Breakdown of starch during ripening and processing, makes it unavailable for fermentation by gut microbiota. The objective of this study was to evaluate the effect of ripening and processing on prebiotic potential of banana. Methods: Fermentation ability has been assessed in vitro and in vivo. In vitro prebiotic evaluation of banana was carried out for raw, ripe and processed banana as a sole carbon source in modified medium. In vivo evaluation was carried out after 4 weeks of administration of raw and ripe banana pulp flour to male Wistar rats. Results and Discussion: In vitro prebiotic evaluation of banana resulted increase in Lactobacillus acidophilus populations, pH reduction, increase in total titratable acidity than control. In a 4-week animal trial, daily administration of raw banana pulp flour increased CFU/gm of Lactobacillus species in caecum content and lowered the pH of the caecum significantly. Acidic pH of gut created by fermentation of prebiotic provides unfavorable condition for the growth of pathogens. Bacteriocin produced by Lactobacilli inhibited the growth of pathogens. Protection of large intestine from pathogens improves health of large intestine. Physicochemical changes during ripening and processing significantly reduced the prebiotic potential of banana.

Key words: Banana, Lactobacillus acidophilus, Prebiotic.

INTRODUCTION
Banana (Musa acuminata) is the popular nutritious food ingredient of South Indian dishes. Banana is mostly consumed either in raw, ripe and processed form in South India. Keralites (people from the Kerala) give raw banana powder to their children since long time. Starch and fibre imparts typical starchy taste to raw banana and decrease its palatability.1 every medium sized banana is rich source of carbohydrate with low fat content. Banana is also a good source of vitamin B9, potassium and vitamin C. Raw banana is rich in resistance starch (RS), which is not absorbed in the small intestine of healthy individual.2 RS has attracted interest due to its health effect on the colon.3 Fermentation of resistant starch in large intestine produces short chain fatty acids (SCFAs), carbon dioxide, methane and hydrogen. SCFAs are volatile fatty acids existing in straight- and branched-chain conformations. Common SCFAs include formic, acetic, propionic, butyric, isobutyric, valeric, isovaleric, and caproic acids. Acetic, propionic and butyric acids were studied extensively for beneficial effect on human health. It reduces pH of the colon, the formation of carcinogetic amines, improves fecal excretion, and mainly butyrate protects the bowels from colon cancer.4 The enzymes initiate chemical reactions as well as accelerate the reaction during ripening.1,6 Physical and chemical changes occur during ripening impart soft texture and sweet taste to banana and improve palatability of ripe banana. High percentage of sugar imparts sweet taste to ripe banana.7 Banana chips, processed form of banana is mostly accepted throughout the world. Frying is routinely used method to prepared banana chips. Frying affects the prebiotic potential of raw banana. Among various banana products on market, fried chips and candy hold major share of market.7 Most of the investigators explored the effect of roasting on prebiotic potential of other materials. very few data is available for effect of frying on prebiotic potential.

The effect of ripening and frying on fermentation of banana is less studied. So, the aim of present study was to investigate effect of raw, ripe and processed banana on the growth of Lactobacilli and to investigate in vivo prebiotic efficacy of raw and ripe banana pulp.

MATERIALS AND METHODS
Preparation of banana pulp flour (BPF)
Raw (banana with full green peel, stage 1 of ripening) and ripe (banana with full yellow, stage 6 of ripening) bananas were obtained from local market, Pune. Peeled fruits were cut into slices. Slices were then dipped in 0.5% (w/v) citric acid solution for 10 min, to avoid browning, drained and dried in oven at 60°C for 18 hr. The dried samples were ground in a household grinder. The powder was to pass through 60 mesh screens to obtain banana flour.8 Raw and ripe banana was sliced using slicer and then deep fried in ground nut oil. Fried chips were kept on tissue paper to absorb excess oil. Fried chips were converted into powder described above.

Probiotic culture maintenance
The starter culture of Lactobacillus acidophilus was procured from National Collection of Industrial Microorganism (NCIM-5426), Pune (MS, India). The culture was maintained on deMan Rogosa Sharpe (MRS) agar (Himedia, India) at 37°C for 24 h. Sub-culturing was done in MRS media every 15 days.

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Growth of *L. acidophilus* on BPF

MRS broth is extensively studied selective medium for growth of Lactobacillus. Modified MRS broth was prepared as per the method described by Mandalari *et al.* (2007), using BPF as sole carbon source medium. BPF was added to the broth at a final concentration of 5%. The experimental media was sterilized at 121°C for 15 min before inoculation. Colonies isolated from MRS agar plates were pre-cultured twice in MRS liquid broth at 37°C, first for 24 h and then for 18 h, to ensure that all the cells were harvested from the early stationary phase. The bacterial suspensions 2% (v/v) was then used to inoculate the BPF medium. In all cases, the initial microbial concentration was approximately at 10^6 CFU/mL. Modified media was incubated in micro-aerophillic conditions for 48 h. Samples were taken periodically to determine optical density, % titratable acidity, pH and antimicrobial activity against *E. coli*.

Monitoring of growth

Growth of *L. acidophilus* was monitored by measuring optical density using UV-Visible spectrophotometer (Shimadzu Analytical Pvt. Ltd, Japan) at 600 nm. Measured values of optical density were plotted on growth curves. The maximum specific speed for the growth was calculated during the exponential growth phase through the following equation (Kask *et al.* 2003):\(^{10}\)

\[
\ln \left( \frac{\text{optical density at the end of the exponential growth phase}}{\text{optical density at the beginning of the exponential growth phase}} \right) = \frac{\ln \left( \mu_{\text{max}} \right)}{t_d} \times \text{Time interval between observations}
\]

The doubling time was determined through equation

\[
t_d = \ln 2 / \mu_{\text{max}}
\]

Where, \(\mu_{\text{max}}\) - maximum specific growth rate, \(t_d\) - doubling time

Changes in pH of modified media containing banana pulp flour

The pH was measured by an electronic digital type pH meter (Equip-Tronic, Mumbai, India). pH 4.0 and pH 7.0 buffer solutions were used to standardize the pH meter. Measured pH values were plotted on graphs. From graphs the rates of decrease of pH values were calculated.

Titratable acidity

0.01 M NaOH with phenolphthalein as an indicator was used to measure the titratable acidity (as % lactic acid). The % titratable acidity was calculated by using formula:11

\[
% \text{Titratable Acidity} = \frac{9 \times \text{titer value} \times \text{normality of NaOH} \times \text{Dilution}}{\text{Weight of the sample taken}} \times 100
\]

Antimicrobial activity against *E. coli*

Cell-free supernatant was collected by centrifugation at 4,000 rpm for 15 min (C-24, Cooling centrifuge, Remi Instrument LTD, India). Antimicrobial assay was performed by agar well diffusion method. MacConkey agar was used to grow *E. coli*. Sterilized agar was allowed to solidify. Over night culture of *E. coli* was spread on solidified agar and 100 µL of cell free supernatant was added in 6 mm wells. Plates were kept at 4°C for 4 h to facilitate diffusion. Then plates were incubated aerobically at 37°C for 24 h. Zone of inhibition was recorded in duplicate.

In vivo prebiotic potential of raw and ripe BPF

Amongst raw, ripe and processed BPF, raw and ripe BPF showed comparatively good *in vitro* prebiotic efficacy to that of fried BPF. Hence only two samples were assessed *in-vivo*.

In a 4 weeks animal experiment, the male Wistar rats had free access to diet and water and were housed in ventilated room with a 12h:12h light/ dark photoperiod at 23±2°C. All procedures had approval of the Institute Animal Ethical Committee. After a week of acclimatization to experimental conditions, the rats were randomly divided into three groups (n=6). Control (group-I) received distilled water; Group-II and Group III received raw and ripe BPF (2g/day/kg of rat) suspended in distilled water. After 1 week of treatment stool samples were collected for every week and immediately stored at 4°C. Freshly voided fecal material was collected to study faecal moisture, pH and bacterial concentrations.\(^{12}\) The water content of the luminal stools was calculated by weight difference between fresh and dried (kept during 24 h at 65°C) samples. For pH measurement, faecal content was suspended in water, homogenized by vortexing and pH values was measured using a pH-meter. At the end of 4th week animals were scarified, transferred to laminar flow cabinet and caecum content was removed anaerobically. pH and bacterial count of caecum content was determined. Ten-fold serial dilutions were made in the medium and aliquots of 0.1 ml of the appropriate dilution was spread onto the following agar media to determine bacterial count; MRS agar for *lactobacilli* and Rogosa agar (Himedia, India) supplemented with 0.5 g/L cysteine hydrochloride for *Bifidobacterium* and culture plates were incubated microaerophically for *Lactobacillus* and anaerobically for *Bifidobacterium* at 37°C for 24–48 h.\(^{13}\)

Microbiological analysis of caecum content

For the isolation and counting of probiotic bacteria each one of the caecum samples were aseptically diluted by 0.1% (w/v) sterile peptone water. Aliquots of these suspensions were transferred to test tubes containing 9 mL with the same peptone water, so as to obtain serial decimal dilutions (10^{-1} to 10^{-6}).

Later, an aliquot of each diluted sample was transferred to plate dishes containing specific agar as described by Vinderola and Reinheimer *et al.* (1999).\(^{14}\) the counting was performed by a Colony Counter apparatus.

Statistical analysis

All the experiments were conducted at least twice and triplicate samples were used for each test. Data was collected and analyzed by using one way analysis of variance and Bonferroni’s test. All statistical analysis was performed using Graph Pad InStat.

RESULTS

Growth of *L. acidophilus* in presence of BPF

Lag phase of *L. acidophilus* growth curve was decreased, and log phase was increased significantly after addition of raw BPF, as sole carbon source to modified MRS medium (Figure 1). Then, bacterial growth curve reached to stationary phase after 32 h and turned to be relatively constant for 48 h. High optical density of *L. acidophilus* was observed in presence of raw BPF (Figure 2).

Growth kinetics

The maximum specific growth rates (\(\mu_{\text{max}}\)) and the doubling time (\(t_d\)) of the media containing different carbon source after 48 h of anaerobic fermentation are shown in Table 1. *L. acidophilus* showed the fastest growth (\(\mu_{\text{max}}=0.173\) h^{-1}) on raw BPF compared to ripe (\(\mu_{\text{max}}=0.028\) h^{-1}) and processed one. Doubling time of *L. acidophilus* was very less in presence of raw BPF (\(t_d=4.006\) h)
Results and Discussion

Growth kinetics of L. acidophilus

Table 1: Growth kinetics of L. acidophilus in presence of raw, ripe and processed BPF.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Raw BPF</th>
<th>Processed Raw BPF</th>
<th>Ripe BPF</th>
<th>Processed ripe BPF</th>
</tr>
</thead>
<tbody>
<tr>
<td>$h_{max}$ (h⁻¹)</td>
<td>0.173</td>
<td>0.028</td>
<td>0.048</td>
<td>0.030</td>
</tr>
<tr>
<td>Doubling Time (t₆₃)</td>
<td>4.006</td>
<td>24.75526</td>
<td>14.44057</td>
<td>23.10491</td>
</tr>
</tbody>
</table>

Figure 1: Growth of L. acidophilus in reconstituted MRS broth either with raw, ripe and processed BPF.

Figure 2: Maximum OD shown by L. acidophilus in presence of raw, ripe and processed BPF.

The inhibitory activity was distinct towards gram negative E. coli (Figure 4), common causative agent of large intestine. Highest zone of inhibition (16.5±2.12 mm) against E. coli was shown by supernatant obtained from incubation mixture of raw BPF compared to ripe (11 ±2.82 mm) and processed BPF.

Table 2: The titratable acidity of the medium supplemented with raw, ripe and processed banana inoculated with L. acidophilus.

<table>
<thead>
<tr>
<th>Incubation time</th>
<th>Titratable acidity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>Raw BPF</td>
</tr>
<tr>
<td>24 h</td>
<td>0.07±0.03</td>
</tr>
<tr>
<td>48 h</td>
<td>0.09±0.04</td>
</tr>
</tbody>
</table>

Values presented as means ±SD.

which proves the ability of raw BPF content to support the growth of L. acidophilus.

Changes in pH of fermentation media in presence of raw, ripe and processed BPF

Addition of BPF reduces pH of modified MRS medium. Significant reduction in pH from 6.9 (t=0) to 5-5.5 (t=48 h) showed by raw BPF (Figure 3). Rate of decrease in pH was highest in presence of raw BPF. Reduction in pH showed strong relationship to production of lactic acid which is main metabolic product produced by the Lactobacilli. Acidic pH generates unfavorable environment for the growth of pathogens.

Titratable acidity

Increase in percent titratable acidity (Table 2) was observed with raw BPF; 2.16±0.04 to that of ripe; 1.15±0.09, processed raw BPF; 0.57±0.08, processed ripe BPF; 0.79±0.05 and control; 0.09±0.04, after 48 h of fermentation. Produced lactic acid and short chain fatty acids might be contributing increased percentage of titratable acidity.

Antimicrobial activity against E. coli

The inhibitory activity was distinct towards gram negative E. coli (Figure 4), common causative agent of large intestine. Highest zone of inhibition (16.5±2.12 mm) against E. coli was shown by supernatant obtained from incubation mixture of raw BPF compared to ripe (11 ±2.82 mm) and processed BPF.

In vivo prebiotic investigation using Wistar male rats

Four-week administration of raw BPF increased CFU/gm of Lactobacilli significantly in caecum content compared to ripe BPF (Table 3). Increased beneficial microflora in caecum indicates improved beneficial micro flora of gut and vice versa. Table 3 showed significant increase in Lactobacilli spp. after administration of raw BPF (8.57±0.07⁶ log CFU/g) to male wistar rats compared to control (8.04±0.03log CFU/g). E. coli species of caecum content also decreased significantly when rats were fed with raw BPF (5.03±0.15Log CFU/ g of wet caecum content) to that of ripe BPF and control.

DISCUSSION

Disturbed microbial flora can be restored by probiotics, prebiotics and combination of two, referred as symbiotic. Probiotics were extensively studied and having safety records to its credit to restore disturbed gut microbiota. Sensitivity of probiotics to external environmental conditions such as temperature, pH, humidity and internal environment like pH of GI tract renders it less effective for modulation of gut microbiota. Raw BPF significantly increased optical density, lowered the pH and showed less doubling time was observed for L. acidophilus to that of ripe, processed BPF and control. High dietary fibre content of raw banana makes it suitable for the growth of L. acidophilus. Reduced pH of the incubation mixture is an indirect measurement of acid produced during fermentation by L. acidophilus. As undigested BPF was subjected to in vitro fermentation; easily available substrate of BPF such as sucrose, fructose and glucose might be contributing to increase OD along with dietary fibre. All these substrate utilized by L. acidophilus to produce lactic acid and pyruvic acids using Embden-Meyerhof-Parnas pathway using NADH as the cofactor and the enzyme lactate dehydrogenase. Lactic acid and other fatty acids (short chain fatty acids) produced by Lactobacilli contributes to the maintenance of a low pH which is thought to be an important control mechanism preventing colonization by pathogens. Titratable acidity was also found to be increased by L. acidophilus, L. fermentum and L. casei when grown in presence of fructooligosaccharides as a prebiotic. Pyruvate is immediately converted to end products such as linear and branched SCFAs such as acetate, propionate, butyrate and carbon dioxide, hydrogen, methane, and water. These produced SCFAs also might be responsible for increase in percent titratable acidity. Lactic acid bacteria produce inhibitory compounds such as organic acids, hydrogen peroxide, diacetyl and bacteriocin. Bacteriocins are the ribosomally produced cationic proteins inhibit pathogenic bacteria living in the same ecological environment. Lactobacilli use bacteriocin as a
digested in upper gastrointestinal tract. Undigested resistant starch in upper gastrointestinal tract gets fermented in large intestine to produce SCFAs. Linear SCFAs such as acetate, propionate and butyrate have various health benefits to human health\(^{20}\) the ability of raw banana starch to reach at the large intestine increased the CFU/g of Lactobacilli spp in caecum content of male wistar rats compared to ripe. Significant decrease in E. coli, common pathogen of large intestine was also observed in caecum content (Table 3). Increase in lactobacilli competitively inhibits the growth of pathogen such as E. coli. No significant increase in Bifidobacteria was observed after administration of raw BPF. In addition, the pH of the intestinal caecum content was decreased in in treated group. Significant decrease in pH during in vitro fermentation of raw BPF (Figure 3) and increase in titratable acidity (Table 2) showed correlation with lowered caecum pH.

No significant increase in count of Lactobacilli in ripe BPF group compared to raw BPF could be due to breakdown of starch, hemicellulose and pectin substances resulting in softness of the tissue. Synthesis of fruit characteristics like volatiles, pigments and organic acid requires energy which is produced from the breakdown of starch. The enhancement in the activity of enzymes of glycolytic and Kreb’s cycle help the fruit to assimilate energy as ATP produced from the breakdown.\(^{21}\) Therefore ripe banana showed less optical density, reduced lowered pH during in vitro fermentation study and less stimulation of Lactobacilli and Bifidobacteria in vivo study.

Modern food manufacturing methods destroy most forms of RS and make it unavailable to large intestine. Frying of banana to prepare chips may decrease availability of RS in colon drastically, which in turn not significantly increased the Lactobacilli and Bifidobacteria spp in caecum content. (Isolated colonies of Lactobacilli, Bifidobacterium and E. coli on specific agar were confirmed by respective biochemical tests and 16S rDNA performed at Genombio Technologies Pvt. Ltd., data not shown) Lactic acid bacteria exclude adhesion of pathogens and enhances secretion of simple or complex molecules which regulates gut health.\(^{22}\) The formation of harmful amines and carcinogenic substances are reduced by lactic acid bacteria (Lactobacilli and Bifidobacteria) in the colon.\(^{23}\)

Ingestion of unripe banana pulp not only beneficial because of starch but natural antioxidants polyphenols could confer additional health benefits to human health.\(^{24}\)

**CONCLUSION**

Raw BPF significantly promoted the growth of L. acidophilus in vitro, because of presence of resistant starch. Daily administration of raw BPF to male wistar rats modulated intestinal environment via stimulating the growth of beneficial species and inhibiting the growth of pathogens. Other important constituents of banana such as potassium, iron, and vitamin B\(_6\), sugars like glucose, sucrose and fructose

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**Table 3:** Effect of the intake of raw and ripe BPF (2 g/day/rat), for 4 weeks, on the percentage of water, pH values, Lactobacilli and Bifidobacteria spp. in the caecum content of rats.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Lactobacilli spp. (Log CFU/g of wet caecum content) *</th>
<th>Bifidobacteria spp. (Log CFU/g of wet caecum content) *</th>
<th>E. coli (Log CFU/g of wet caecum content) *</th>
<th>pH Caecum#</th>
<th>% water Caecum#</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>8.04±0.03</td>
<td>7.14±0.08</td>
<td>5.83±0.20</td>
<td>6.37±0.13</td>
<td>72.4±1.5</td>
</tr>
<tr>
<td>Raw BPF</td>
<td>8.57±0.07</td>
<td>7.44±0.05</td>
<td>5.03±0.15</td>
<td>6.19±0.16</td>
<td>76.7±1.2</td>
</tr>
<tr>
<td>Ripe BPF</td>
<td>8.17±0.07</td>
<td>7.30±0.05</td>
<td>5.79±0.15</td>
<td>6.41±1.0</td>
<td>73.8±1.0</td>
</tr>
</tbody>
</table>

*Values presented as means and standard errors

Mean values with different superscripts are significantly different according to the Bonferroni’s test

(*P<0.01 vs. Control group, *P<0.05 vs. Ripe BPF).
can exhibit more health benefits. Polyphenols compounds of unripe banana flour would add up health benefits as natural antioxidants. In vitro and in vivo studies showed promising prebiotic potential of raw BPF.

**CONFLICT OF INTEREST**

The authors declare no conflict of interest.

**REFERENCES**