Application of Whey Permeate Fermented with Kefir Grains for the Shelf-Life Improvement of Food and Feed

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Abstract

Fungal contamination by Aspergillus parasiticus and A. flavus causes negative effects on the production of food cereals. Kefir is an ancient fermented beverage obtained by the fermentation of different substrates with kefir grains. An important waste produced by the dairy cheese industry is the whey permeate, which nowadays is a strong ambient contaminant. The aim of this work was the standardization of the whey permeate fermenting conditions with kefir grains, the assessment of the antifungal activity of the cell-free-supernatants (CFS) from these fermentations and to compare it with that obtained with CFS of milk fermented with kefir grains. Finally, we studied if the addition of kefir fermented whey permeate to food (bread) and feed (poultry) could produce shelf life improvement. The optimal condition to obtain CFS with fungicidal effect was a fermentation with 10% w/v of kefir grains, at 30°C, for 24 hours until a pH 3.7. We found that CFS from whey permeate caused fungal inhibition, whereas CFS of kefir grains grown in milk showed lower antifungal activity. Additionally, the addition of kefir-fermented whey permeates in food (bread) and feed (poultry) improved their resistance to fungal contamination. This is the first report about the application of kefir-fermented whey permeate to improve the shelf life, suggesting its potential use as a biopreservative.

Keywords

Aspergillus parasiticus, Whey Permeate, Kefir Supernatants, Bread, Poultry Feed

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1. Introduction

Fungal contamination of food by *Aspergillus flavus* and *A. parasiticus* causes large food and feed losses especially in tropical areas, where temperature and humidity favor its development [1] [2]. Contamination problems are frequent in food processing plants, where fungal conidia from raw materials are difficult to eradicate from the environment. Even though molds are destroyed during the baking process, contamination arises from mold spores derived from atmosphere and from surfaces during the cooling, finishing or wrapping procedures [3]. High temperature and humidity in the processing environment cause the fungal prevalence and food contamination.

One alternative that has been widely explored is the application of lactic-acid bacteria, which produce organic acids, exhibit antibacterial and antifungal activities [4]-[9].

Kefir is a sour milk slightly carbonated with a low alcohol content that is obtained through the use of kefir grains. These grains are clusters of lactic- and acetic-acid bacteria along with yeasts in a structural matrix of polysaccharides and proteins. The microorganisms present are responsible for the lactic, acetic, and alcoholic fermentation of milk that yields a product with characteristic organoleptic properties [10].

Kefir-fermented milk supernatants at a concentration of 10% (v/v) in a broth completely inhibited the fungal growth of *A. flavus* and *Fusarium graminearum* [11]. Cell free supernatants obtained from whey permeate fermented with kefir grains inhibited the *A. parasiticus* and *F. graminearum* growth and the aflatoxin B1 and zearalenone production [5] [6].

Argentina is the fifth world cheese producer after the European Union, USA, Russia and Brazil. The estimated production is around 580,000 tons for 2016 [12]. Cheese whey produced after cheese manufacture has diverse applications, mainly the obtention of whey proteins for feed additive. However, after producing protein concentrates, cheese whey permeate (WP) remains as a waste. Due to its composition whey permeate is a strong ambient contaminant [13].

The consumers’ demand of biopreservatives in foods, instead of chemical preservatives has arisen since the XX century. The application of microorganisms classified as GRAS (Generally recognized as safe) such as lactic acid bacteria and their metabolites strongly promoted the research in this area [14]. Even though the obtention of these preservatives require further studies in scaling the volume process, the results obtained are very interesting to continue the assessment of natural preservatives offered to a public increasingly interested in natural foods. Kefir fermented products using WP as substrate could give a solution for the ambient contamination and additionally could be used as biopreservative for food and feed with special application against fungal contamination. Although production costs are not analyzed in this work, the decrease of industrial effluents combined with the development of biopreservatives could generate environmental benefits. Besides, the small companies could avoid unnecessary legal disputes for the effluents disposal.

The aim of this work was to study the kinetic parameters and antifungal effect of kefir grains grown in whey permeates and the application of kefir-fermented products on bread and poultry feed in order to improve their shelf-life. Finally, we studied the sensory properties of bread added with kefir-fermented products.

2. Materials and Methods

2.1. Fungal Cultures and Preparation of Conidia Inoculum

*A. flavus* AFUNL5 was isolated from cereals at Laboratorio de Micología, Universidad Nacional del Litoral (Argentina). *F. graminearum* Fg44 strains were provided kindly by Dr. Teresa Alconada of the Centro de Investigacion y Desarrollo en Fermentaciones Industriales (CINDEFI) of the Universidad Nacional de La Plata (Argentina). *A. terreus* CMUNLP1, *A. fumigatus* CMUNLP2, *Penicillium sumatrense* CMUNLP3, *P. crustosum* CMUNLP4, *Trichoderma longibrachiatum* CMUNLP5 and *Rhizopus microspores* var. *rhizopodiformis* CMUNLP6 were isolated from the commercial poultry feed Nutrisur® BB (Nutrisur, La Plata, Argentina). *A. parasiticus* CMUNLP7 was isolated from corn (Dra. León at Cathedra of Microbiology-National University of La Plata). *A. parasiticus* NRRL 2999 was obtained from the Agricultural Research Service Culture Collection (Washington, DC, USA). Fungi were maintained at 4°C in soft agar (2 g/L agar in water) until used. The inoculum was prepared by growing the fungi on PDA (Potato Dextrose Agar) slants (Britania, Argentina) for 7 days at 30°C. After incubation, 10 ml of 0.01% (w/v) Sodium Lauryl Sulfate (SLS) in 1% (w/w) sodium chloride solution were added to the tubes and spores were loosened by gently scraping with a spatula, and serial dilutions
were made. The number of spores, about $5 \times 10^5$/ml was assessed by means of account in a Neubauer chamber [15] [16].

### 2.2. Preparation of Cell-Free Supernatants and Organic Acid Solutions

The kefir grains used -CIDCA AGK1 were characterized at the Centro de Investigación y Desarrollo en Criotecnología de Alimentos (CIDCA), UNLP [17] [18] and stored in whole milk at $-20^\circ$C. The kefir grains were activated through two consecutive passages of fermentation in commercial ultra-high-temperature-processed (UHT) milk (Sancor, Santa Fe, Argentina). The grains were inoculated into UHT milk or sterile WP at a concentration of 10% (w/v) and incubated at $30^\circ$C for 24 or 48 h. The fermentation products were then separated from the grains by passage of the fermentation mixture through a sieve of 1 mm$^2$ mesh size and the microorganisms present precipitated by centrifuging for 15 min at 14,000 g in an Eppendorf 5415D centrifuge (Eppendorf, Hamburg, Germany).

The standardization of the kefir grains CIDCA AGK1 was carried out as follows: Kefir grains CIDCA AGK1 grown in milk were adapted to WP by consecutive passages of fermentation (24 h at $30^\circ$C) during four weeks (kefir grains WP-4) or twelve weeks (kefir grains WP-12). The fermentation products obtained from these kefir grains were used on different assays (acidification kinetic, biomass production and addition to food and feed).

The resulting kefir-fermented-whey-permeate cell-free supernatants (KFWP CFS) or kefir-fermented-milk cell-free supernatants (KFM CFS) were obtained by filtration through a microcellulose filter of 0.22-μm pore size (Sigma-Aldrich, St. Louis, USA) and stored at $-20^\circ$C.

### 2.3. Fungal Inhibition Assays with KFWP CFS and KFM CFS

The assays were performed on Petri dishes with the basal medium (BM) containing malt extract (1% w/v) (Biorbiker, Beauvais, France), yeast extract (2% w/v) (Biorbiker, Beauvais, France) and agar (2% w/v) (Merck, Darmstadt, Germany). The media were autoclaved at 121°C for 15 min. The agar medium at 45°C was mixed with the KFWP CFS or with lactic and acetic acid mixtures in order to obtain different final concentrations. Twenty milliliters of supplemented medium were plated in Petri dishes (80 mm diameter) and the final pH of the medium was determined with a Hannah pH211 Microprocessor pH meter.

The assays with KFWP CFS in the media included two different controls to investigate the effect of filter-sterilized WP on fungal growth. In the first, the BM was used alone, while in the second the BM was supplemented with 70% (v/v) unfermented WP.

Three plates each of controls or with additions of KFWP CFS or organic acids were inoculated with 10 μL of the conidia suspensions at $5 \times 10^5$ conidia/mL dispensed by micropipetting in the center of the solidified growth medium.

The diameter of the circular inoculums obtained was assumed as the colony initial diameter. Inoculated plates were incubated at 30°C inside plastic boxes containing dishes of water to prevent dehydration. Growth was allowed until the maximum 80 mm diameter, correspondent to the total Petri dish invasion by mycelium; all cultures were kept in the boxes during 30 days in order to determine the total fungal inhibition [19]. Given that filamentous molds grow on solid media forming a circular colony around the initial inoculation zone, colony diameters were measured each daily by placing the Petri dishes on a millimeter scale illuminated from beneath by a light box. Four diameter measures were taken from the center of each colony and the results were calculated from the mean diameter of the replicate colonies [20].

### 2.4. Determination of Growth Rate ($K_D$) and Lag Phase (Lag)

Growth rate $KD$ (mm/h), was calculated from the regression slope of colony diameter versus time during the linear growth phase, using the Sigma Plot 9.0™ software. The lag phase (Lag) is the time in hours required for the colony to grow beyond the inoculation zone (typically 5 - 7 mm). This value corresponded to the point on the abscissa where the regression line intersected the horizontal line representing the initial inoculation-zone diameter. Thus,
where: $D_0 = \text{diameter of the inoculation zone}$, $Y_0 = \text{intersection of the regression line}$ with the ordinate and $KD$ = slope of the regression line (i.e., growth rate; [15] [21]).

2.5. Evaluation of the Fungistatic and/or Fungicidal Effect of KFWP CFS

From the plates of the growth-inhibition experiment where no fungal development occurred, the agar in the central inoculation zone was cut out and placed in 80-mm plates containing unsupplemented BM in order to assess whether the inhibitory effect on fungal development had been fungicidal or simply fungistatic. The plates were then incubated for 30 days at 30°C to investigate the fungal-growth capability upon removal from the source of inhibition. This determination was carried out after 7, 15 and 30 days of incubation.

2.6. Effect of KFWP CFS Preparations on the Fungal Germination

We conducted assays on the inhibition of conidial germination in the presence of KFWP CFS preparations and unfermented cell-free whey supernatants after the dilution and pH adjustment of those fractions. A volume of 10 μL of a suspension $5 \times 10^5 \text{ conidia/mL}$ of each fungal isolate was added to a volume of 190 μL of each treatment in a 96-well enzyme-linked-immunosorbent-assay (ELISA) plate (Cellstar, Frickenhausen, Germany). Treatments were performed as follows: KFWP CFS (24 h of fermentation and pH 3.7), unfermented WP acidified with lactic and acetic acid mixture (at the same concentrations measured in the KFWP CFS) and adjusted pH at 3.7 or unfermented WP acidified with HCl and adjusted pH at 3.7. Controls were performed with cell-free supernatant of WP added with the same concentration of conidia as used with the experimental samples. The plates were incubated for 48 h at 30°C and the optical density (OD) at 580 nm in the wells was monitored with a Synergy HT™ ELISA reader from Biotek Instruments (Winooski, USA). The percent inhibition was calculated by:

\[
\% \text{Inhibition} = 100(\text{OD control} - \text{OD treated})/\text{OD control}
\]

(2)

The extent of inhibition obtained was scored semiquantitatively according to Gerez et al. [7], where inhibitions of 20% or greater were considered positive, <40% low, ≥40% but <70% moderate, and ≥70% high. Each determination was made in quintuplicate. Three independent assays were performed. Differences were statistically tested using one-way analysis of variance (ANOVA) and the Tukey test to determine significant differences (α = 0.05).

2.7. High-Performance-Liquid-Chromatography Analysis of Lactic and Acetic Acids in KFWP and KFM CFS

The concentrations of lactic and acetic acids in the KFWP and KFM CFS were measured by high-performance liquid chromatography (HPLC) in a chromatograph (Agilent Technologies, series 1200, Santa Clara, CA, USA) with an Aminex HPX-87H ion-exchange column (Bio-Rad, Hercules, CA, USA), a mobile phase of 45 mM H₂SO₄ (Merck, Darmstadt, Germany), a flow rate of 0.6 mL/min, and an ultraviolet detector measuring at 214 nm [16].

Pure acetic acid (Merck, Darmstadt, Germany) at concentrations of 0.42, 0.83, 1.67, 3.33, 11.70, 25.00, and 66.10 mM and lactic acid (Carlo Erba®, Milan, Italy) at 5.55, 11.10, 44.40, 88.80, 111.00, 155.00 and 209.00 mM were used for the control curves.

2.8. Calculation of the Concentrations of Undissociated Organic Acids

The concentrations of the undissociated acids were calculated by the following equation:

\[
HA = \frac{[TA] \times [H^+]}{([H^+] + Ka)}
\]

(3)

where $[HA]$ is the undissociated acid concentration (mM). $Ka$ is the equilibrium constant of lactic (pKa 3.79) or acetic acid (pKa 4.75) and $[TA]$ is the total acid concentration (mM).
2.9. Preparation of Supplemented Poultry Feed

A 200 mL aliquot of fermented KFWP was added to 200 g of poultry feed Nutrisur® BB. Then the supplemented feed was dried in a convection oven at 50°C until a water activity \((a_w)\) of 0.95 ± 0.05 was achieved. By the same methodology, unfermented WP to give the same concentration as the fermented product were added separately to poultry feed. Feed supplemented with water and dried by the same procedure was used as control.

The water activity \((a_w)\) of supplemented poultry feed was measured at 20°C with an AquaLab Series 3 TE meter (AquaLab, Pullman, WA, USA). The pH was measured using a pH211 pH meter equipped with an HI 1330B microelectrode (Hanna Instruments, Ann Arbor, MI, USA).

2.10. Preparation of Supplemented Bread

Bread was prepared as described below: wholemeal flour 410 g; commercial compressed yeast (Casla®, Argentina) 35 g; brown sugar 40 g; butter (La Paulina®, Santa Fe, Argentina) 80 g; salt (Dos Anclas®, Argentina) 10 g; water 200 mL. To produce bread supplemented with KFWP or unfermented WP, water (200 mL) was totally replaced by KFWP or unfermented WP (200 mL).

An amount of 35 g commercial yeast was mixed with 20 g brown sugar and water (or KFWP or unfermented WP). After 15 min, the mixture was placed into an electric mixer Kenwood KM 005 (Kenwood®, Hampshire, UK), which was used at medium speed. Simultaneously, remaining flour, brown sugar and water or KFWP was added together with the salt and butter. Mixing process took place during 20 min. The resulting mass was stretched, rolled and placed in fermenter Pauna Beta 107 (Pauna®, Buenos Aires, Argentina) at 30°C with saturated water vapor for 45 minutes for fermentation. Mass was baked in an oven (Ariston®, Argentina) preheated to 210°C for 27 minutes. The resulting bread was cooled at 25°C for 2 h. Bread was cut and slices were used to perform fungal contamination assays and sensory analysis.

2.11. Determination of the Resistance to Artificial Fungal Contamination of Bread Loaf Prepared with Kefir-Fermented Milk

A fungal suspension (of \(A.\) flavus AFUNL5, \(A.\) parasiticus NRRL 2999 or \(P.\) sumatrense CMUNLP3) of \(10^4\) conidia/mL was prepared. Then, 1 mL of this suspension was pulverized per 100 g bread, whereas the uncontaminated bread control was treated in the same way but with sterile distilled water. The samples were then left to dry before storage at 14°C in sealed polyethylene bags. Thereafter the bags were inspected daily for determination of visible signs of the fungal presence [7]. The intervening time was considered the bread resistance to the fungal contamination.

2.12. Sensory Analysis

In order to study the effect of the KFWP on the sensory characteristics of bread, two different techniques were performed [22].

2.12.1. Similarity Test

Each triangle test set consisted of 3 samples (2 identical samples and 1 different sample) placed on a tray, and the panelists were asked to identify the different sample. The panelists were asked to taste the samples in the order indicated on their ballot sheets [23]. The values obtained were analyzed using the following equation.

\[
LS\% = \left(1.5x\left(x/n\right) - 0.5\right) + 1.5xZ_{\beta}x \sqrt{n \left(x - x^2 \right)/n^3} \times 100
\]

where \(LS\%\) = upper limit of the panelist’s percentage; \(x\) = number of correct answers; \(n\) = number of panelists; \(Z_{\beta} = 1.64\) (value corresponding to one tail of the normal curve), \(\beta = 5\%\). Statistical parameters for similarity were \(\alpha = 0.10; \beta = 0.05\) with an estimated proportion of discriminators (Pd) of 30% [24].

2.12.2. Acceptability Test

This analysis was made by using the 9-point hedonic scale, where the panelists qualified the product with a score between 1 (“dislike very much”) to 9 (“I like very much”) [25].
2.13. Statistical Analysis

All the growth parameters were analyzed by the SIGMAPLOT 10.0® software. The results of three independent assays are presented as the mean values ± standard deviation (SD). Differences in growth kinetics were tested for significance by the analysis of variance (ANOVA) to determine significant effects at p < 0.05 by means of the STATGRAPHICS Plus 5.1® software. All the experiments were performed at least in triplicate.

3. Results

Characterization of Kefir Grains Grown in Milk and WP

The standardization of kefir grain fermentation in WP was carried out with three types of kefir grains CIDCA AGK1: kefir grains grown in milk (KFM), kefir grains grown in WP during four weeks (WP-4) and twelve weeks (WP-12). Kefir grains grown in milk and those adapted to WP were able to ferment WP (Figure 1). When kefir grains WP-12 were grown in WP the acidification ability decreased in comparison with grains WP-4 (Figure 1(a)). However, kefir grains grown in milk and grains WP-4 reached a similar pH after 24 h fermentation (around pH 3.7), in contrast to WP-12 (pH 4.51) (Figure 1(a)). On the other hand, kefir grains adapted to WP decreased their biomass. Kefir grains WP-12 showed a higher decrease than WP-4, whereas kefir grains grown in milk increased their biomass (Figure 1(b)). However, the counts of LAB and yeasts remained unchanged in all three conditions.

Fermentation products of kefir grains grown in milk (KFM) and kefir grains WP-4 and WP-12 grown in WP were obtained, centrifuged, sterilized by filtration and used in the fungal-inhibition assays. KFWP CFS-4 (cell-free supernatants from WP fermented with WP-4) showed fungicidal activity, whereas KFWP CFS-12 (cell-free supernatants from WP fermented with WP-12) showed poor antifungal activity (Table 1).

Moreover, the KFWP CFS presented a change in the concentration of undissociated acids (lactic plus acetic acids). KFWP CFS-12 showed lower concentration of undissociated acids than KFWP CFS-4. KFWP CFS-4 with fungicidal activity (70% v/v CFS in the medium) showed a concentration of undissociated acids similar to that observed in kefir fermented milk cell-free supernatants (KFM CFS) added at the same concentration (70% v/v CFS) in the medium. KFM CFS showed no-fungicidal activity. KFM CFS showed a lower antifungal activity, which could be due to the presence of milk compounds that produce a protective effect on the fungus. Also, the presence of WP or the decrease of the pH (for the addition of strong acid) per se caused no-antifungal activity (Table 1). These results suggest that kefir grains lose their fermentative and antifungal activity when they were grown in WP during long-term periods, even though the counts of LAB and yeasts were similar to both kefir grains grown in WP during four and twelve weeks.

To improve the knowledge about antifungal activity of kefir, KFWP CFS-4 was used for germination inhibition assays. CFS of WP acidified with lactic and acetic acids (at the same concentration found in KFWP CFS-4) and CFS of WP acidified with HCl were also assayed. Two behaviors were observed. P. sumatrense CMUNLP3, P. crustosum CMUNLP4 and F. graminearum Fg44 showed a higher sensitivity due that these fungi presented a similar inhibition for KFWP CFS, WP acidified with organic acids and WP acidified with HCl. In contrast, A. parasiticus NRRL2999, A. parasiticus CMUNLP7, A. flavus AFUNL5, A. terreus CMUNLP1, A. fumigatus CMUNLP2, T. longibrachiatum CMUNLP5 and Rh. microspores var. rhizopodiformis CMUNLP6 showed a lower sensitivity due to they were inhibited by KFWP CFS, whereas inhibition decreased with WP acidified with organic acids or WP acidified with HCl (Figure 2). These findings suggest that the antifungal activity of KFWP is produced, partially, for the presence of lactic and acetic acids plus the presence of other metabolites produced by kefir microorganisms.

Poultry feed was added with KFWP (containing total kefir microorganisms) and then artificially contaminated with several fungi. Table 2 shows that poultry feed added with KFWP showed a significant increase of the time lapsed until appearing visible fungal contamination (>30 days) for all fungi analyzed, whereas poultry feed control and poultry feed added with unfermented WP showed lower time for the visible fungal contamination (between 12 and 14 days).

Due to the promising results found, KFWP (containing total kefir microorganisms) was used as a component to produce whole-meal bread, which was artificially contaminated with several fungi. Both controls (Bread Control and Bread added with unfermented WP) showed not significant differences in the time lapsed until appearing visible fungal contamination. Bread loaves added with KFWP showed a significant increase of the time
Figure 1. Acidification kinetic (a) and biomass change (b) of kefir grains CIDCA AGK1 adapted to WP during different periods of time (○ 4 weeks and (●) 12 weeks) or grown in (□) milk. After adaptation, 10 g of grains were inoculated in 100 ml of WP and assayed for acidification kinetic. Counts of LAB and yeasts after 48 h in milk: 2.84 ± 0.35 × 10^8 CFU/mL and 1.15 ± 0.51 × 10^7 CFU/mL, respectively. Counts of LAB and yeasts after 48 h in WP with grains WP-4: 1.08 ± 0.14 × 10^8 CFU/mL and 2.24 ± 0.38 × 10^7 CFU/mL, respectively. Counts of LAB and yeasts after 48 h in WP with grains WP-12: 1.66 ± 0.45 × 10^8 CFU/mL and 3.52 ± 0.20 × 10^7 CFU/mL, respectively.
Table 1. $K_D$ and Lag phase of *A. parasiticus* NRRL2999 challenged against KFWP CFS obtained from kefir grains in WP for 4 weeks and 12 weeks and KFM CFS.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Assays with KFWP CFS-4</th>
<th>Assays with KFWP CFS-12</th>
<th>Assays with KFM CFS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$K_D$ (mm/h)</td>
<td>Lag phase (h)</td>
<td>Undissociated acids (mM)</td>
</tr>
<tr>
<td>BM* plus unfermented WP or milk</td>
<td>0.82 ± 0.02</td>
<td>18.3 ± 1.2</td>
<td>0.40</td>
</tr>
<tr>
<td>BM plus unfermented WP or milk acidified with HCl</td>
<td>0.81 ± 0.01</td>
<td>19.5 ± 2.5</td>
<td>0.71</td>
</tr>
<tr>
<td>BM plus KFWP CFS 25% v/v</td>
<td>0.68 ± 0.02</td>
<td>17.5 ± 1.1</td>
<td>6.05</td>
</tr>
<tr>
<td>BM plus KFWP CFS 50% v/v</td>
<td>0.69 ± 0.01</td>
<td>33.2 ± 2.1</td>
<td>16.47</td>
</tr>
<tr>
<td>BM plus KFWP CFS 70% v/v</td>
<td>0</td>
<td>&gt;720.0</td>
<td>34.09</td>
</tr>
</tbody>
</table>

*Bm*: basal medium.

Figure 2. Percentage of inhibition of conidial germination of different fungi challenged against KFWP CFS (white bars), CFS of WP acidified with organic acids (black bars) and CFS of WP acidified with HCl (gray bars). KFWP CFS were obtained as described in Materials and Methods. WP was artificially acidified with lactic plus acetic acids at the same concentrations detected in KFWP CFS.
lapsed until appearing visible fungal contamination related to the controls for the three fungi analyzed (Table 3).

In order to determine whether bread added with KFWP could be accepted by the consumers, sensory tests were carried out. Triangle tests showed that panelists could differentiate between bread loaves elaborated with WP (unfermented WP or KFWP added to the recipe) and bread loaves control elaborated without WP (Table 4). Additionally, acceptability tests were carried out. Bread loaves added with KFWP showed a good acceptability (7.3 out of 10 points). Moreover, a value of 71.3% panelists answered that they would buy the product (Table 4). These results suggest that the application of KFWP in bread could be used to improve the resistance to fungal contamination and that a good percentage of the consumers would buy bread added with KFWP.

4. Discussion

Kefir grains are a complex community constituted by LAB, yeasts and acetic-acid bacteria. Milk fermented by kefir grains has been reported to contain, in general, lactic and acetic acids, acetaldehyde, ethanol, carbon dioxide (in equilibrium with carbonic acid), B vitamins, and hydrocarbons such as diacetyl [17] [26]. In our experiments, we reported that kefir grains fermented WP, as reported previously [5] [6]. However, we reported for the first time the effect of the WP on the kinetic parameters of kefir grains. We demonstrated that WP as a culture medium during long-term periods caused a decrease on the fermentative ability and the biomass production (Figure 1). When kefir grains CIDCA AGK1 were grown in milk, they increased their biomass. Similar results were obtained by Guzel-Seydim et al. [27], with kefir grains grown in nonfat reconstituted milk.

### Table 2. Poultry feed added with KFWP or unfermented WP and contaminated artificially with different fungus strains.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>aw of feed</th>
<th>Time lapsed until appearing visible fungal contamination (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>A. flavus AFUNL5</td>
</tr>
<tr>
<td>Control</td>
<td>0.948</td>
<td>13.0 ± 1.3</td>
</tr>
<tr>
<td>Poultry feed added with unfermented WP</td>
<td>0.953</td>
<td>13.0 ± 0.8</td>
</tr>
<tr>
<td>Poultry feed added with KFWP</td>
<td>0.957</td>
<td>&gt;30.0*</td>
</tr>
</tbody>
</table>

1 aw was measured by an AquaLab Series 3 TE meter (AquaLab, Pullman, WA, USA). 2 Poultry feed was inoculated with 10⁴ conidia/mL per each 100 g of feed. * Statistical differences between the values of each column. Two independent assays were carried out.

### Table 3. Bread loaves added with KFWP or unfermented WP and contaminated artificially with different fungus strains.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>aw of bread</th>
<th>Time lapsed until appearing visible fungal contamination (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>A. flavus AFUNL5</td>
</tr>
<tr>
<td>Bread Control</td>
<td>0.943</td>
<td>5.0 ± 0</td>
</tr>
<tr>
<td>Bread added with unfermented WP</td>
<td>0.941</td>
<td>6.7 ± 2.1</td>
</tr>
<tr>
<td>Bread added with KFWP</td>
<td>0.947</td>
<td>12.7 ± 1.5*</td>
</tr>
</tbody>
</table>

1 aw was measured by an AquaLab Series 3 TE meter (AquaLab, Pullman, WA, USA). 2 All bread loaves were inoculated to 10⁴ conidia/mL per each 100 g of bread. Two independent assays were carried out. * Statistical differences between the values of each column.

### Table 4. Acceptability of bread loaves added with KFWP or unfermented WP into recipe or sprayed with KFWP.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Triangle Test</th>
<th>Acceptability punctuation</th>
<th>Percentage of consumers that would buy the product</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control Bread</td>
<td>A</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Bread added with unfermented WP</td>
<td>B</td>
<td>7.7 ± 1.6</td>
<td>82.0</td>
</tr>
<tr>
<td>Bread added with KFWP</td>
<td>B</td>
<td>7.3 ± 1.7</td>
<td>71.3</td>
</tr>
</tbody>
</table>

*Treatments with the same letter were not differentiated by the panelists. NA: not analyzed.
Previously, Rimada and Abraham [28] found that kefir grains CIDCA AGK1 are able to ferment deproteinized whey (obtained from skimmed milk) which composition was 5.34% lactose and 0.32% protein. Other authors have demonstrated that kefir grains CIDCA AGK10 are able to ferment cheese whey, which presented a concentration 1% and 6.5% of protein and lactose, respectively [29]. In our assays, kefir grains fermented WP, which contained 0.02% and 3.32% of protein and lactose, respectively (data not shown).

We found that the counts of LAB and yeasts of KFWP-4, KFWP-12 and KFM presented no-changes. However, with this methodology we did not detect changes in the quality of the microbial community that could indicate alterations in the balance of lactobacilli species. These results suggest that there are microbial population changes which cause a decrease in acidification ability and a possible loss of the microbial population responsible for producing the kefiran (matrix which contains the kefir microorganisms) (Figure 1).

On the other hand, we showed that KFWP CFS exerts fungicidal activity against *A. parasiticus* NRRL2999. Our results agreed with those obtained with kefir grains grown in cheese whey and WP that also showed antifungal activity against diverse fungi [5] [6] [9]. KFWP CFS showed fungicidal activity against *A. parasiticus* [5] and *F. graminearum* [6]. However, we found that kefir grains decreased their antifungal activity when they were in WP during long-term periods (twelve weeks). These results suggest that it is recommended to use KFWP CFS obtained with kefir grains grown in WP during short-term periods (not more than four weeks) to obtain satisfactory antifungal effect. Additionally, KFM CFS showed lower antifungal effect and no-antifungal activity due to, possibly, the presence of milk compounds which protect the fungus (Table 1). Similar results were obtained with kefir-fermented milk supernatants against *A. flavus* AFUNL5 [30]. Additionally, KFWP CFS showed higher antifungal activity than WP acidified with organic acids or WP acidified with HCl (Figure 2). These findings suggest that the antifungal activity of KFWP is owed to the presence of lactic and acetic acids plus the presence of other metabolites produced by kefir microorganisms such as other organic acids or bacteriocins. The latter ones are a wide group of compounds with potential antimicrobial activity. In this sense, bacteriocin F1 produced by *Lactobacillus paracasei* subsp. tolerans isolated from Tibetan kefir grains exhibited antimicrobial activity, with a strong heat and pH stability [31].

On the other hand, the application of KFWP to poultry feed or bread caused the improvement of resistance to fungal contamination in both substrates (Table 2 & Table 3). Our results are in agreement with those reported by Londero *et al.* [9], who demonstrated that kefir-fermented whey improve the poultry feed resistance to fungal contamination. In addition, Gerez *et al.* [7] found that the application of a strain of commercial yeast plus four strains of *Lactobacillus* to bread improved the resistance to contamination from two days (control bread) to 5 days (treated bread). Additionally, the results presented here are in concordance with those reported by Gamba *et al.* [6], who found that kefir-fermented milk added at a concentration of 10% w/v to corn arepas improved their shelf life and their resistance to fungal contamination.

### 5. Conclusions

The presence of KFWP CFS in high concentrations exhibited a fungicidal capability against *A. parasiticus* NRRL2999. This capability was lost when kefir grains were grown in WP during long-term periods.

Unfermented WP acidified with lactic plus acetic acid CFS showed lower antifungal activity than that caused by KFWP CFS, suggesting the presence of other metabolites produced by kefir microorganisms that improve the antifungal activity. The application of KFWP in poultry feed and bread caused an improvement of resistance to fungal contamination. Additionally, bread added to KFWP showed a good acceptance.

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### Conflict of Interest

There is no conflict of interest with other co-authors for the publication of this manuscript in this journal. All the co-authors have contributed in the preparation of the manuscript up to the submission stage.
References


