CONVERSION OF LINOLEIC ACID TO DIFFERENT FATTY ACID METABOLITES BY *Lactobacillus Plantarum* 13-3 AND IN SILICO CHARACTERIZATION OF THE PROMINENT REACTIONS

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ABSTRACT

*Lactobacillus plantarum* strains have been extensively used in food processing and preservation. *L. plantarum* also has the potential to convert polyunsaturated fatty acids, e.g. linoleic acid (LA) into bioactive and less toxic fatty acid metabolites. The objective of this study was to assess the capability of probiotic *L. plantarum* 13-3 to convert Linoleic Acid (LA) to different fatty acid metabolites in the medium supplemented with differential concentrations of LA, and the relevant reactions were characterized by *in silico* calculation. *L. plantarum* 13-3 was grown in MRS medium containing LA from 1% to 10%, and the fatty acid metabolites formed in the medium were identified and quantitated by GC-MS and *in silico* studies were done to confirm the enzymatic reactions involved in its conversion. The results showed that *L. plantarum* 13-3 could convert LA at different concentrations to 5 different fatty acid metabolites i.e., (Z)-Ethyl heptadec-9-enoate, 9,12-Octadecadienoic acid (Z, Z), methyl ester, Octadec-9-enoic acid, cis-11,14-Eicosadienoic acid, methyl ester and (Z)-18-Octadec-9-enolide. Among these metabolites, the formation of an long chain fatty acid Octadeca-9-enoic Acid, also known as 18:1, N-9 or Delta(9)-Octadecenoic acid, is classified as a member of the Long-chain fatty acids in media supplemented with 4% to 10% LA, is being reported for the first time. Putative candidate enzymes involved in biotransformation of LA into fatty acid metabolites were identified in whole genome of *L. plantarum* 13-3, sequenced previously. *In silico* studies confirmed that several enzymes including the linoleate isomerase, acetoacetate decarboxylase and oxidoreductase may be involved in biotransformation of LA into fatty acid metabolites. These enzymes could effectively bind the LA molecule mainly by forming hydrogen bonding between the acidic groups of LA and the proline residues at the active sites of the enzymes validating the putative reacting partners.

Keywords: *Lactobacillus plantarum*, fatty acid metabolites, Linoleate isomerase, Acetoacetate decarboxylase, Dehydrogenase, *in silico* study.

INTRODUCTION

Linoleic acid (c18:2 n-6) (LA) is the most abundant polyunsaturated fatty acid (PUFA), also known as an essential fatty acid and it is a significant constituent of low-density lipoproteins [1]. LA cannot be produced endogenously in animals and hence its only source is via dietary intake [2]. Algae and plants contain adequate amounts of the Δ12- and Δ15-desaturase enzymes and as a result LA and α-linolenic acid are the most widespread fatty acid found in plant tissues and oils.

As a PUFA, LA can be oxidized through endogenous enzymes and reactive oxygen species in the circulation. Metabolites of LA are formed as a result of oxidation by the action of endogenous enzymes. LA and its byproducts are known to exhibit various biological effects and they are involved in metabolic disorders and cancer [3-7]. The irony of an essential fatty acid contributing in both beneficial and pathological processes may be explained by its conversion to biologically active metabolites [8-9]. Several metabolic processes in the host are regulated by gut microbiota, some of these processes include lipid metabolism, glucose metabolism and homeostasis [10]. The gut microbiota is also capable of performing various processes that cannot be carried out by the host and these processes can give rise to microbially produced or modulated metabolites which works as metabolic substrates and signaling molecules in the host, with key implications for host metabolism and health [11]. Many bacteria have been noticed that their growth has been inhibited by polyunsaturated fatty acids (PUFA) such as Linoleic acid (LA; 18:2Δ9Z,12Z). PUFA in high concentration is toxic and may block native fatty acid biosynthesis by the inhibition of enoyl-ACP reductase [12-15]. Bacteria have developed a detoxification mechanism and are capable of enzymatically hydrogenate PUFA by completely reducing the double bonds on the carbon chain, i.e., bio hydrogenation, producing non-toxic saturated fatty acids as the end product [16]. This process includes various steps and has been greatly described for LA and oleic acid (OA; 18:1 Δ9Z) that are converted to non-toxic saturated stearic acid (SA; 18:0) by rumen microbiota [16-19].

Lactic acid bacteria (LAB) are the most important and common starter cultures used in fermented dairy products, and they may originate from the microflora of raw milks (e.g., bovine, ovine, caprine) but most commonly are inoculated intentionally during product manufacture.

LAB are generally regarded as safe (GRAS) microorganism which have been traditionally used in food fermentation for a long history. Traditionally fermented dairy products are considered as a key source of functional microorganism, e.g. LAB and ingredients [20]. Many LAB strains are able to produce various bacteriocins, exopolysaccharides, fatty acids, etc. engaging their useful health effects. Many LAB strains have shown different promising bioactivities on human health, including antimicrobial activity, prevention and treatment of diarrhea, relief of symptoms caused by lactose intolerance, anti-mutagenic and anti-carcinogenic activities and stimulation of the immune system [21]. However, the uncertainties of the influence from these LAB strains on the quality of functional foods and their bioactivity keeping in food matrix frequently prove their application in modern food industry [22].

*Lactobacillus plantarum* is a facultative homofermentative lactic acid bacteria (LAB). It may get energy from different sugars, and it inhabits an adaptive niche such as gastrointestinal, vaginal, and urogenital tracts, vegetables, dairy products, and fermented foods [23-28]. This bacterium also carries significant properties for not only the manufacture of a diversity of food and wine but also vitamin production, bacteriocin, probiotic, antifungal, and potential antitumors agents [29-33].

*L. plantarum* is homofermentative having the competency to transform growth-inhibiting free polyunsaturated fatty acids, e.g. linoleic acid (LA) at a comparatively higher concentration, into bioactive conjugated LA (CLA) and other less toxic fatty acids metabolites e.g. hydroxy fatty acids, o xo fatty acids, conjugated fatty acids and saturated trans fatty acids, as the intermediates to eventually produce saturated monoenes (OA and trans-vaccenic acid) [34-36].

*In silico* analysis of the whole genome sequences of *L. plantarum* 13-3, exposed the presence of many enzymes including acetoacetate decarboxylase, Oxidoreductase and linoleate isomerase.

We validated our findings by employing in silico analysis to characterize the relevant reactions involving the enzymes that were identified using the whole genome sequence of *L. plantarum* 13-3. Our findings would have great prospect for human nutrition, nutraceuticals and food industry and will pave way into identifying the relevant reactions and doses for biotransformation of LA into desirable fatty acid metabolites.
**MATERIALS AND METHODS**

**Microorganism and growth conditions**

*L. plantarum* 13-3 isolated from Tibetan Kefir preserved in the culture bank of Dairy Laboratory in Beijing Technology and Business University of China was repeatedly revived for three times at 37°C in MRS medium (Beijing Asboxing Co Ltd) containing 2.0% glucose, 1.0% meat extract, 1.0% tryptone, 0.5% yeast extract, 0.1% Tween 80, 0.2% K2HPO4, 0.5% sodium acetate, 0.2% diaminonium citrate, 0.02%, MgSO4 x 7H2O and 0.005%, MnSO4 x H2O was used in this study. Distilled water was used as solvent for dissolving medium components and medium was adjusted to pH 5.5 and sterilized at 121°C for 15 minutes. The fresh MRS medium added with different concentrations of LA was inoculated with 1% of the activated culture of *L. plantarum* 13-3 for growth and production of metabolites at 37°C.

**L. plantarum 13-3 Screening for the production of fatty acid metabolites**

MRS media was supplemented with linoleic acid (LA). During growth of *L. plantarum* 13-3 in the MRS medium containing 1%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, 9% and 10% (w/v) of LA, culture samples were taken at 0, 8, 16, 24, 36 and 48 h for the determination of viable counts by plate count method as expressed by colony forming units (CFUs) per mL, and for determination of fatty acid metabolites as explained below.

**Spectrophotometric determination of fatty acid metabolites**

The culture sample were centrifuged at the speed of 13,000 xg for 5 min at 4°C and 1 mL of the supernatant was mixed with 2 mL of isopropanol. After addition of 1.5 mL of hexane, the mixture was extensively vortexed in order to extract the lipids and then allowed to stand for 5 min. The hexane layer was collected and the absorbance was measured at 233 nm. The fatty acid metabolites were extracted by using hexane/isopropanol (2:1, v/v) solution at room temperature, and the extracts were washed with distilled water and then dehydrated with anhydrous sodium sulfate [37].

**Extraction of fatty acids from the medium.**

For the analysis by gas chromatography (GC), the culture samples were centrifuged at 1900 rpm for 5 minutes at 4°C to remove the cells. An internal standard (C17:0, heptadecanoic acid, 98% pure; Macklin) was added to 5 mL of the supernatant fluid to give a final concentration of 0.15 g/ml. Then 5 mL of isopropanol was added and vortexed for 30 s. Subsequently 2 mL of isopropanol was added and vortexed for 30 s. Lastly, 5 mL of n-hexane was added to this mixture, vortexed for 3 min, incubated for 30 min, and centrifuged at 1900 rpm for 5 min. The upper hexane layer containing fatty acid methyl esters (FAME) was collected and was dried under a stream of liquid nitrogen [38].

**Gas chromatography and mass spectrometry (GC-MS)**

The GC-MS analysis was done by using Shimadzu GC-2010 instrument coupled with a Dual Stage TSP (Ultra) mass spectrometer. The FAME sample of 2 μL was injected in a split mode, set at 10:1 split ratio at 250°C. The carrier gas was helium at a constant flow rate of 1 mL/min. The separation was conducted on a highly polar (TR-Wax MS, 30 m length x 0.25 mm i.d. x 0.25μm thickness) and fused silica capillary column (Thermo Fisher Scientific). The initial oven temperature was held at 170°C for 1 min, then increased at 0.8 °C/min to 200°C. The temperature of line transfer was at 250°C, and the ion source was controlled at 200°C. The MS detector was operated in an electron ionization (EI) voltage of 70 eV under a mass scan range of 33–450 amu (m/z).

**Identification of fatty acid metabolites**

Chemical identification was conducted by comparison of the mass spectra (MS) of the peaks with those found in the National Institute of Standard and Technology library (NIST, 2014).

**In silico characterization of conversion of LA to various fatty acid metabolites by L. plantarum 13-3**

Enzymes responsible for catalyzing the relevant reactions for conversion of LA to different fatty acid metabolites in *L. Plantarum* 13-3 were identified by analysis of the whole genome sequence (GCA_004028315.1) sequenced earlier using SWISS-MODEL that is a fully automated protein structure homology modelling server [39] The protein module was visualized by protein visualizing program Discovery Studio 3.5 [40]. The LA molecule was drawn in ChemSketch v15.0.9 (Chemaxon) assigned with proper 2D orientation, and the structure of each compound was analyzed for connection error in bond order. Energy of the molecules was minimized using Avogadro [41] with MMFF94 force field. Docking studies calculations were performed by Autodock Tool [42]. Protein (LAI, DH and DC) and ligands (LA) structures were converted to pdbqt file by MGL Tools 1.5.6 rc1. The interactions of complex protein-ligand conformations were analyzed using Autodock Tools 4.2 [42], and Discovery Studio 4.1 [40].

**Statistical Analysis.**

Statistical analysis of the data was done using ANOVA SAS version 9. Evaluation of the significance of differences between groups was performed with one-way ANOVA as mentioned in figure legends.

**RESULT AND DISCUSSION**

*L. plantarum* 13-3 exhibited similar growth pattern at the concentrations of LA from 1% (w/v) to 10% (w/v) as shown in Fig. 1A. For all the concentrations of LA, it was observed that *L. plantarum* 13-3 increased its growth up to 24 h, subsequently entered stationary phase till 36 h, and then the growth decreased till 48 h. There was slightly increase in the growth of the strain with the increase of the LA concentration. On the contrary, many other LAB strains were reported to be inhibited to different extent by LA and their tolerance to LA varied [37,39-40]. While earlier studies showed that even lower LA levels (25 µg/ml) could inhibit bacterial growth [37], *L. plantarum* 13-3 was able to grow well at the concentration of LA up to 10 % (w/v), indicating its relatively high tolerance to LA. (Fig1B), showed that production of total fatty acid metabolites gradually increased with the increase of LA concentration from 1% (w/v) to 10% (w/v) as indicated by the increased absorption at the wavelength of 233 nm. This indicated that *L. plantarum* 13-3 exhibited high tolerance to LA by converting LA to less toxic fatty acid metabolites. Previous researcher also advocated that conversion of free LA to LA analogues might function as a detoxification mechanism in bacteria and a stronger LA tolerance indicated a higher productivity of LA analogues/metabolites [37,39,45-46]. Furthermore, more fatty acid metabolites were produced during the stationary phase of growth of *L. plantarum* 13-3 and the production reduced when the death phase started. Similar findings were also reported with other microbial producers of LA fatty acid metabolites/analogues [47-48]. Significantly, it should be highlighted that the spectrophotometric method does not differentiate between isomers of CLA since it is based on measurement of the conjugated double bond in the fatty acid [37].

Fig 1C shows different fatty acid metabolites produced by *L. plantarum* 13-3 with different concentrations of LA in the medium with different time interval of 8, 16, 24, 36 and 48 hours.
The fatty acid metabolites produced by *L. plantarum* 13-3 at different concentrations of LA were identified and quantitated by GC (Table 1). Total of 5 fatty acid metabolites were identified including one long chain fatty acid, octadec-9-enoic acid C₁₈H₃₃O₂, one ethyl ester, (Z)-ethyl heptadec-9-enoate C₁₇H₃₁O₂, two methyl esters 9,12-octadecadienoic acid (Z,Z)- methyl ester C₁₈H₃₂O₂ also known as methyl linoleate which is found in clove and is mainly used as a flavoring agent and cis-11,14-eicosadienoic acid, methyl ester, and (Z)-18-octadec-9-enolide as shown in Table 1. The following compounds, (Z)-ethyl heptadec-9-enoate, 9,12-octadecadienoic acid (Z, Z)- methyl ester, and cis-11,14-eicosadienoic acid methyl ester were identified under all concentrations of added LA as shown in Table 1. These findings proved that the ability of *L. plantarum* 13-3 to produce the above mentioned fatty acids. While, the octadec-9-enoic acid was produced by *L. plantarum* 13-3 under the concentration of added LA ranged from 4% to 10%. Finally, (Z)-18-octadec-9-enolide was produced only at 10% of added LA as listed in Table 1.

| Name                                         | RT   | 1% | 2%  | 3%  | 4%  | 5%  | 6%  | 7% | 8%  | 9%  | 10% |
|-----------------------------------------------|------|----|-----|-----|-----|-----|-----|----|-----|-----|-----|
| (Z)-Ethyl heptadec-9-enoate                    | 15.112 | + | +  | +   | +   | +   | +   | +  | +   | +   | +   |
| 9,12-Octadecadienoic acid (Z,Z)- methyl ester | 15.729 | + | +  | +   | +   | +   | +   | +  | +   | +   | +   |
| Octadec-9-enolic acid                         | 29.670 | - | -  | +   | +   | +   | +   | +  | +   | +   | +   |
| cis-11,14-Eicosadienoic acid, methyl ester    | 31.935 | + | +  | +   | +   | +   | +   | +  | +   | +   | +   |
| (Z)-18-Octadec-9-enolide                      | 32.002 | - | -  | -   | -   | -   | -   | -  | -   | -   | +   |

+ detected, - not detected

**In silico studies for relevant reactions**

**Linoleate Isomerase**

Linoleate isomerase is the enzyme which has the responsibility for the conversion of one isomeric form of ligand to another form via enzymatic mechanism. Isomers are different form of the same ligands having same molecular formula but different structural formula having different 3D orientation in space. The genome of *L. plantarum* 13-3 was blasted to discourse different linoleate isomerase giving many hits, and best hit was selected for further study.

The Conversion of linoleic acid (LA) into several metabolites were studied by using computational approach. LA was optimized by using Avogadro and saved into PDB format. Linoleate isomerase gene was found in *L. plantarum* 13-3 which was blasted by using NCBI platform and then module by using Swiss model. Linoleate isomerase is a monomer protein having tertiary structure which consists of its backbone, side chain, hydrophobic moiety, hydrophilic moiety, acidic group and basic group shown in figure 2. The collective interaction of the linoleate isomerase with LA was carried by using Auto dock Vina Program. The optimized structure of linoleate isomerase and linoleic acid was used for Auto-dock study.
Table 2. Interaction of LA with Linoleate isomerase, and their result calculated by autodock.

| Binding Energy | -1.89 |
|----------------|-------|
| Ligand efficiency | -0.09 |
| Inhibition Constant | 41.27 |
| Intermol Energy | -5.61 |
| Vdw_hb_desolv energy | -4.94 |
| Electrostatic energy | -0.67 |
| Total internal | -0.4 |
| Calculated RMS | 0.0 |

After docking it was observed that Carboxylic Acidic moiety of LA formed Hydrogen bonding with Lysine (LYS A:549) and also showed electrostatic and VDW wall interactions. As Linoleate isomerase is NADPH dependent enzyme so Hydrogen from NADPH attack on the double bond of the LA ultimately the reaction of LA with Linoleate Isomerase and NADPH leading to the formation of different metabolites.

Figure 3: 2D Interaction of LA with Linoleate Isomerase, shows the 2D interaction of Linoleic Acid with the enzyme Linoleate Isomerase.

The interaction of the LA in the active site of the Linoleate Isomerase showing the Perfect fit because it occupies the active site of the Linoleate Isomerase which provides the pathway for enzymatic reaction.

The Acidic group of LA formed hydrogen bond with Lysine 549 (LYS:549). The acidic moiety holds by enzyme by hydrogen and electrostatic interaction, which provide medium for the double bond transfer reaction.

Figure 4: 3D interaction of the LA in the active site of Linoleate Isomerase.

Acetoacetate Decarboxylase

Acetoacetate decarboxylase belongs to the class of enzymes which plays a vital role in solvent production by catalyzing the decarboxylation of acetoacetate moieties, producing acetone and carbon dioxide. The production of solvent acetone by enzyme acetoacetate decarboxylase containing bacteria was utilized in large-scale industrial syntheses in the first half of the twentieth century. Recently we have explored in this study that beyond it, it also plays important role in fatty acid chemistry by converting LA into different fatty acid metabolites. The genome of L. plantarum 13-3 was blasted for the search of different Acetoacetate Decarboxylase giving many hit, and best hit was selected for further study. The Protein Structure of Acetoacetate Decarboxylase of L. plantarum 13-3 module by Swiss model is shown in figure below. The structure of enzyme Acetoacetate Decarboxylase having ligand in the active site is given below in figure 5.

Figure 6: 2D and 3D Interaction of LA with Acetoacetate Decarboxylase.

Figure shows 2D and 3D interaction of Linoleic Acid with the enzyme Acetoacetate Decarboxylase in L. plantarum 13-3.

The interaction of the LA in the active site of the Acetoacetate Decarboxylase showing the Perfect fit because it occupies the active site of the Acetoacetate Decarboxylase which provides the pathway for enzymatic reaction.

The Acidic group of LA formed two hydrogen bonds with Lysine (LYS A:114) and Arginine (AGR:106) and also showed electrostatic and VDW wall interactions.

The enzyme Acetoacetate Decarboxylase of L. plantarum 13-3 is an oligomer which consists of twelve monomer having active site amino acid include lysine 114, Arginine 106, Isoleucine 179, Tyrosine 177, Tyrosine 87, Leucine 115, Aspartate 219 and Aspartate 220. The Hydrogen bond between Arginine 106 and LA have a distance of 2.233 nm while the angle DHA (Donor Hydrogen Acceptor) is 159.62 and angle HAY (Hydrogen Acceptor on Y Axis) is 145.70 and LA having the distance of 2.97 nm has the DHA angle 98.73 and the HAY angle 165.99. The second hydrogen bond between Lysine 114 and LA have a distance of 2.314 nm while the angle DHA (Donor Hydrogen Atom) is 151.04 and angle HAY (Hydrogen Acceptor) is 106.001.

The following table 3 is showing the docking result of LA with Acetoacetate Decarboxylase L. plantarum 13-3.

Table 3. Interaction of LA with acetoacetate decarboxylase, and their result calculated by autodock.

| Binding Energy     | -2.31 |
|--------------------|-------|
| Ligand efficiency  | -0.12 |
| Inhibition Constant| 20.28 mM |
| Intermol Energy    | -5.41 |
| Vdw_hb_desolv energy | -4.54 |
| Electrostatic energy | -0.87 |
| Total internal     | -1.02 |
| Calculated RMS     | 0.0   |
Oxidoreductase

Oxidoreductase is a class of enzyme that are involved in catalyzing the reaction which involve the transfer of electrons from one molecule, the reductant, also called the electron donor, to another, the oxidant, also called the electron acceptor. So these enzymes usually involve the transfer of charge from catalyzing molecule leading to oxidation or reduction. This group of enzymes usually NADP or NAD+ dependent enzyme which use it as cofactors. Dehydrogenase is tetramer, composed of monomer A, B, C and D. The active site residue and substrate in it is reported in figure 7 given below.

The active site residue and substrate in it is reported in the following figures. The amino acids in the active site are Glycine 17, Tyrosine 206, Asparagine 90, Methionine 186, Glycine 189, Tyrosine 156, Glycine 92, Isoleucine 93, Asparagine 114 and Lysine 160. The amino acid GLY A:17 in the active site of oxidoreductase form hydrogen bonding with substrate (LA), Which provide the plate form for the enzymatic reaction by holding acidic moiety of substrate for reduction reaction. As these enzymes are cofactor dependent enzyme, so hydrogen from the FADH or NADPH attack on the unsaturated moiety and resulting the formation of reduction product. Moreover, the docking result is reported in table 4.

The following table 4 is showing the docking result of LA with Oxidoreductase in L. plantarum 13-3.

| Binding Energy | -1.64 |
| Ligand efficiency | -0.08 |
| Inhibition Constant | 62.68 |
| Intermol Energy | -4.98 |
| Vdw_hb_desolv energy | -5.29 |
| Electrostatic energy | -0.31 |
| Total_internal | -0.78 |
| Calculated RMS | 0.0 |

The Hydrogen bond between Lysine 17 and LA have a distance of 2.38 nm while the angle DHA (Donor Hydrogen Atom) is 130.161 and HAY (Hydrogen Acceptor) angle is 91.442.

**Figure 8.** 2D and 3D interaction of LA with the enzyme Oxidoreductase in L. plantarum 13-3, is showing the 2D and 3D interaction of Linoleic acid with the enzyme Dehydrogenase Oxidoreductase in L. plantarum 13-3.

CONCLUSION

*L. plantarum* 13-3 was shown to be a competent candidate for converting LA from 1% to 10% (w/v) to various fatty acid metabolites such as Z-Ethyl heptadec-9-enoate, 9,12-Octadecadienoic acid (Z,Z) , methyl ester, Octadec-9-enoic acid, cis-11,14-Eicosadienoic acid, methyl ester and (Z)-18-Octadec-9-enolide. Main reactions involved in this conversion include isomerization, dehydrogenation and reduction as confirmed by the *in silico* analysis. The putative linoleate isomerase and dehydrogenase catalyzing the relevant reactions were identified using the whole genome sequence of *L. plantarum* 13-3. These two enzymes were shown to bind the LA molecule at their active sites mainly by formation of hydrogen bonding between the acidic group of LA and the proline residues of the enzymes. Further study is required to investigate the role of the conversion from LA to its various fatty acid metabolites by *L. plantarum* 13-3 physiologically and the implications for exploring the potentiality in functional foods.

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The image contains a page from a document with a text body that includes scientific references and research findings. The text is about Lactobacillus plantarum and its properties, including its role in biofilm formation inhibition, bioinformatics studies, and its application in food science and technology. The text is dense with scientific terms and data from various studies and journals.

For example, a reference cites a study on the inhibition of biofilm formation of Streptococcus mutans by L. plantarum lipoteichoic acid. Another reference discusses the production of volatile phenols by Lactobacillus plantarum in wine conditions. The text also mentions the use of genomic diversity of Lactobacillus plantarum strains isolated from various origins.

The document appears to be a comprehensive review or an article that integrates multiple studies to present a detailed understanding of Lactobacillus plantarum's role and its applications.

To understand the context and the implications of these findings, one would need to refer to the original research papers cited within the text.