Prevention of enteric bacterial infections and modulation of gut microbiota with conjugated linoleic acids producing *Lactobacillus* in mice

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**Abstract**

Probiotics are recognized for outcompeting pathogenic bacteria by competitive receptor-mediated colonization and secretion of functional metabolites which are antimicrobial against certain microbes as well as improving host’s gut health and immunity. Recently, we have constructed a bioactive *Lactobacillus casei* (LC) strain, LC<sup>mcra</sup>, by inserting *mcra* (myosin cross-reactive antigen) gene, which stimulates the conversion of conjugated linoleic acids. In this study, we evaluated the modulation of gut microbiome and protective roles of LC<sup>mcra</sup> against pathogenic *Salmonella enterica* serovar Typhimurium (ST) and enterohemorrhagic *E. coli* (EHEC) infections in BALB/cJ mice. We observed that LC<sup>mcra</sup> colonized efficiently in mice gut intestine and competitively reduced the infection with ST and EHEC in various locations of small and large intestine, specifically cecum, jejunum, and ileum (*p* < 0.05). Positive modulation of the cecal microbiota, for example, higher relative abundances of Firmicutes, lower relative abundances of Proteobacteria, and increased bacterial species diversity/richness, was detected in ST-challenged mice pretreated with LC<sup>mcra</sup> based on 16S metagenomic sequencing. Cytokine gene expression analysis indicated that mice pretreated with LC<sup>mcra</sup> associated with attenuated bacterial pathogen-induced gut inflammation. Furthermore, mice fed daily with LC<sup>mcra</sup> for one week could protect themselves from the impairments caused by enteric infections with ST or EHEC. These impairments include weight loss, negative hematological changes, intestinal histological alterations, and potential death. This in vivo study suggests that daily consumption of novel conjugated linoleic acids over-producing probiotic effectively improves intestinal microbiota composition and prevents/combats foodborne enteric bacterial infections with pathogenic *Salmonella* and diarrheagenic *E. coli*.

**Introduction**

The majority of human gut epithelial surfaces are colonized and safeguarded by a tremendous number of microorganisms including bacteria, viruses, fungi, and protozoans which are known as common gut microflora; each of them is crucial in forming and balancing a complex ecosystem with microbial diversity. These large number of microorganisms build up a microbial genetic repertoire approximately 100 times greater than that of the human host. Diversity of these microbes, specifically number of diverse bacterial species, is essential for good health and immunity of host. According to recent reports, human distal gastrointestinal (GI) tract can house more than 1000 distinct bacterial species, and the total number was estimated to be larger than 10<sup>14</sup> CFU/gm of fecal material. Bacteroidetes, Firmicutes, Proteobacteria, and Actinobacteria are the prevalent bacterial phyla in human gut microbiota and each of these phyla contains dozens of bacterial genus and hundreds of species.

In a gut ecosystem with homeostatic condition, most of the commensal bacteria colonize and survive symbiotically, whereas conditions such as immunodeficiency, malnutrition, and antibiotic-therapy cause dysbiosis and imbalance of commensal bacteria that induce pathogenesis and cause diseases. Furthermore, broad-spectrum antibiotic therapy or any other detrimental conditions may disturb the gut ecosystem balance long-term or lead to chronically irritated bowels.
reducing the number of beneficial bacteria and increasing the number of opportunistic pathogens and their toxic products that further weaken the host defense and/or induce inflammation and damage.\textsuperscript{11} As a consequence of imbalanced gut microflora, opportunistic pathogens, their produced metabolites, proteins, and/or toxins can take over the gut ecosystem and negatively impact host gut health.

Salmonella and diarrheagenic Escherichia coli generally infect human gut intestine through consumption of contaminated foods and/or drinks.\textsuperscript{12–14} Once these Gram-negative enteric pathogenic bacteria arrive in host gut, their complex type III secretion systems are activated, enabling them to introduce effector proteins directly into cell cytoplasm. Series of pathogenesis through type III secretion systems induce systematic infections causing acute or chronic inflammation and other serious disorders in the host.\textsuperscript{15} However, such enteric illness is usually facilitated by compromised gut immunity and dysbiotic gut microbiota which provide those enteric bacterial pathogens with weakened colonization resistance.\textsuperscript{16} On the other hand, traditional antibiotic therapy has been found to lyse enterohemorrhagic E. coli (EHEC) which further increases the risk for hemolytic-uremic syndrome (HUS), a post infectious sequelae in the patients.\textsuperscript{17,18} In these situations, pro- and prebiotics can be considered as priority in prevention and treatment of such foodborne bacterial pathogen-induced enteric illness.\textsuperscript{13,19,20}

With a promising scheme, it allows an establishment or recovery of the healthy enteric microbial ecosystem by introducing native, exogenous, or genetically engineered beneficial probiotics without inducing deleterious effects (like antibiotics) on human commensal gut bacteria.\textsuperscript{16,21}

Recently, we constructed and reported the role of a multi-functional Lactobacillus casei probiotic strain overexpressing myosin cross-reactive antigen gene (mcra), named as LC\textsuperscript{+mcra}.\textsuperscript{22} Several groups of researchers have demonstrated the health-beneficial effects of conjugated linoleic acids, such as anti-carcinogenesis, anti-oxidant, and anti-microbial effects.\textsuperscript{16,23,24} Similarly, we have also revealed the anti-pathogenic and anti-inflammatory properties of linoleic acids over-producing L. casei (LC\textsuperscript{+mcra}) based on in vitro examination. Here in this study, we aimed to evaluate the protective roles of LC\textsuperscript{+mcra} on modulating/recovering gut intestinal microflora composition and combating/alleviating foodborne enteric bacterial pathogenic infections \textit{in vivo} based on mice model.

Results

Probiotics preventing ST infection induced physiological abnormalities in mice

The weight of each mouse was monitored every day for the purpose of investigating if probiotics preventive administration could rescue mice from weight loss due to ST/EHEC infection (Figure 1). Within the entire 4-week rearing, a total of 12 mice in control group (no probiotic given), 7 mice in group given wild-type probiotic LC strain, and 1 mouse in group given linoleic acid over-expressed mutant LC\textsuperscript{+mcra} strain were sacrificed due to their health abnormality induced by ST infection. These sacrificed individuals included 8 mice from control and 5 mice from LC treatment found self-death due to ST challenge, but none from LC\textsuperscript{+mcra} treatment, which provided us the ST survival rates as 60% in control group, 75% in LC group, and 100% in LC\textsuperscript{+mcra} group (Figure 2). The death of the mice was generally accompanied with extreme (>20%) weight loss to approximately 8–10 g.

At the end of week 2, the average weight of mice in control group reached approximately 14–16 g, whereas both groups of mice which were given either LC or LC\textsuperscript{+mcra} gained weight at range of 1–2 g more compared to the control group of mice. Once mice were challenged with ST, the average weight gain trend of mice in control group which was not given probiotic was suspended and remained at 14.65 g during 1\textsuperscript{st} week of post-challenge. Then the weight of those mice decreased to 14.36 g and 13.47 g at 2\textsuperscript{nd} and 3\textsuperscript{rd} post-infection weeks, respectively (Figure 1A). However, the mice which were administered LC\textsuperscript{+mcra} kept continuing to gain average weight. In spite of the negative effect induced by ST infection, mice which were given LC\textsuperscript{+mcra} gained weight at 16.88 g, 17.02 g, and 19.12 g at the 1\textsuperscript{st}, 2\textsuperscript{nd}, and 3\textsuperscript{rd} week of post-infections, respectively (Figure 1C). The wild-type probiotic, LC fed mice exhibited mild effects in
maintaining the average body weight during the first two weeks of ST infection and gained approximately 1.5 g weight at the end of 3rd post-infection week (Figure 1B). On the other hand, we failed to observe any negative effects including average weight loss induced by EHEC infection. However, the oral administration of LC⁺mcra in mice was more effective in stimulating the weight gain by 1.1g and 1.4 g at the 2nd and 3rd post-infection weeks compared with mice administered LC (Figure 1D,E,F).

**Reduction on colonization of ST and EHEC in probiotics fed mice**

Either LC or LC⁺mcra was orally administered to mice in order to examine their colonization ability in mouse gut and evaluate their preventive role in altering enteric pathogenic bacterial colonization and infection in gastrointestinal tract of mice using BALB/cJ mice model. According to the colonization data collected from two individual mice trials, both LC and LC⁺mcra were able to colonize well in gut of BALB/cJ mice but the genetically modified probiotic strain, LC⁺mcra, could colonize in the mice gut more aggressively compared to the wild-type LC strain. Further, both LC and LC⁺mcra significantly reduced the colonization and infection of both enteric bacterial pathogens, ST and EHEC in BALB/cJ mice. We found that mice fed with LC⁺mcra could defend ST infection remarkably and recover fully within a week of challenge. Specifically, mice highly colonized with LC⁺mcra strain showed significantly reduced cecal colonization with ST (approximately 1 log CFU/g) compared to the group of mice which were given wild-type LC strain at all three time points (14, 21, and 28 d) (Figure 3A).

To compare the colonization of ST in jejunum, we observed that LC or LC⁺mcra pre-administered mice were colonized with lower number of ST at
range of 1.0 to 2.3 log CFU ST per gram jejunum fluids at 1\textsuperscript{st} week post-infection, 0.9 and 2.5 log CFU/g at 2\textsuperscript{nd} week post-infection, and 1.3 and 3.7 log CFU/g at 3\textsuperscript{rd} week post-infection (Figure 3B). Similarly, LC and LC\textsuperscript{+mcra} pre-administered mice were colonized with ST in lower rate of 1.7 and 2.2 log CFU per gram ileum fluids at 1\textsuperscript{st} week post-infection, 0.9 and 1.9 log CFU/g on 2\textsuperscript{nd} week post-infection, and 1.2 and 3.4 log CFU/g on 3\textsuperscript{rd} week post-infection to the control mice (Figure 3C).

The significant reduction on ST gut intestinal colonization was also observed in form of decreased ST fecal shedding (Figure 3D). On the 8\textsuperscript{th} day after mice were challenged with ST, both groups of mice administered with either wild-type probiotic LC or genetically modified probiotic LC\textsuperscript{+mcra} strain were colonized with reduced number (0.8 to 1.1 log CFU/mL) ST in feces but the differences became unsubstantial at the 9\textsuperscript{th} day. However, notable major effectiveness of LC\textsuperscript{+mcra} started to exhibit in mice after 1\textsuperscript{st} week post-infection, at which 1.3 log CFU/mL less ST was recovered from mice feces. In the subsequent two weeks, ST fecal shedding from LC\textsuperscript{+mcra} fed mice were observed to be continuously reduced by 1.1 and 2.1 log CFU/mL.

On the other hand, mice which were pretreated with LC barely reduced the EHEC colonization in jejunum and ileum, whereas mice pretreated with LC\textsuperscript{+mcra} showed significant influence in EHEC colonization resistance (Figure 4). Specifically, LC\textsuperscript{+mcra} fed mice were capable of significantly reducing the colonization of EHEC at 2.3, 1.6, and 0.9 log CFU/g in cecum, 1.6, 1.8, and 2.7 log CFU/g in jejunum, and 2.8, 1.8 and 2.1 log CFU/g in ileum at the 1\textsuperscript{st}, 2\textsuperscript{nd}, and 3\textsuperscript{rd} week post-challenge (Figure 4A,B,C). Meanwhile, consequential decreased EHEC fecal shedding was detected in LC\textsuperscript{+mcra} fed mice as well. However, only insignificant reductions (0.1 to 0.5 CFU EHEC less per mL feces) were found during the first two days after EHEC challenge on EHEC-free mice (the 8\textsuperscript{th} and 9\textsuperscript{th} day). The LC\textsuperscript{+mcra} administration substantially lowered 0.9, 1.9, and 2.2 CFU/mL

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\includegraphics[width=\textwidth]{figure3.png}
\caption{Effect of LC\textsuperscript{+mcra} on reducing colonization of ST in mice gut intestine. The bacterial numbers of ST at 14, 21, and 28 days in cecum (a), jejunum (b), ileum (c), and feces (d) from ST-infected mice with no probiotic treatment, LC, or LC\textsuperscript{+mcra} 1-week pretreatment were investigated in triplicate. Different letters (‘a’ through ‘c’) at individual time point (day 14, 21, or 28) are significantly different ($p < 0.05$) in the numbers of ST among control, LC pretreatment, and LC\textsuperscript{+mcra} pretreatment.}
\end{figure}
EHEC fecal shedding at the 1st, 2nd, and 3rd post-infection weeks in comparison with control (Figure 4D).

**Efficient colonization of LC<sup>+</sup>mcra in mice gut**

In order to examine the correlation between probiotic colonization and reduction on intestinal bacterial pathogens, we also compared the colonization level of both LC and LC<sup>+</sup>mcra in different portion of mice gut (Figure 5). The one-week daily oral administration led to high and stable cecal colonization level of LC<sup>+</sup>mcra above 10<sup>6</sup> CFU/g throughout 3 weeks afterwards, which were significantly higher than wild-type LC (Figure 5A). A similar trend was found in mice jejunum (Figure 5B), whereas, LC<sup>+</sup>mcra only exhibited numerical higher ileum colonization than wild-type LC (Figure 5C).

The fecal shedding number of administered LC were observed to raise after 1<sup>st</sup> day consumption (Figure 5D). Specifically, LC<sup>+</sup>mcra fecal shedding colonies gradually increased from 4.8 log CFU/mL, reached 5.8 log CFU/mL at the next day of final daily administration, and slightly decreased to around 5 log CFU/mL after 3 weeks. Whereas, fecal shedding colonies of wild-type LC were observed significantly lower (by 0.4-1.5 log CFU/mL) than LC<sup>+</sup>mcra. They reached 5.1 log CFU/mL as peak at the next day of final daily administration and ended up with lower than 3.5 log CFU/mL after 3 weeks.

**Mice hematology**

The hematological changes in mice with ST infection with or without pretreated with probiotic strains at various time points were summarized in Table 3. When compared with control group mice with placebo, ST challenge resulted in dramatic increase of red blood cells (RBC) but decrease of white blood cells (WBC) and platelets (PLT). Both pretreatment of LC or LC<sup>+</sup>mcra alleviated the increment of RBC and loss of WBC/PLT colonies.
in mice during Salmonellosis. LC$^{+mcra}$ treatment on mice could further help the mice maintain their normal levels of RBC, WBC, and PLT.

To further evaluate the WBC composition in blood collected from the mice challenged with ST with or without pretreated with probiotic strains, we investigated neutrophils, lymphocytes, monocytes, eosinophils, and basophils counts in different time points, which is summarized in Table 4. The numbers of neutrophils and lymphocytes in blood collected from mice challenged with ST were found to be notably reduced, whereas the monocytes, eosinophils, and basophils levels in mice with salmonellosis were detected to be significantly higher. The pretreatment with either probiotic strain, wild-type LC or mutant LC$^{+mcra}$, in ST-infected mice assisted in maintenance of their normal WBC composition (including all five cell types studied) at similar levels compared with control group.

**Histopathological changes in mouse cecum**

The histological examination of mouse cecal sections is shown in Figure 6. Tissue of cecum collected from the control group mice (Figure 6A) and ST-challenged mice with pre-administration of LC$^{+mcra}$ (Figure 6C) exhibited normal intestinal villi, microvilli, and goblet cells. In comparison, salmonellosis induced variable levels of histological alterations and abnormalities consisting of severe goblet cell depletion, villi/microvilli elimination, and inflammatory infiltrations between circular folds in cecum sections from ST infected mice which were not pretreated with probiotics (Figure 6D,E,F). However, the tissue of cecum collected from the mice administered with LC showed symptoms of salmonellosis, but the induced histopathological changes were mild, such as slight goblet cell reduction and slight changes of villi/microvilli (Figure 6B).
Regulation on expression of intestinal inflammatory cytokine genes

The regulation of cecal inflammatory cytokine gene expressions during 3-week ST infection as well as 1-week probiotic pre-administration is displayed in Figure 7. Specifically, ST infection induced up-expression of 4 pro-inflammatory cytokines IL-1β, IL-6, INF-γ, TNF-α genes and 1 anti-inflammatory cytokine IL-10 gene in mice cecal tissue cells. The up-regulation levels ranged from 2.2 to 7.8 log folds with the highest values for INF-γ gene and the highest expression at two weeks after ST challenge (Day 21). Another anti-inflammatory cytokine TGF-β gene was found down-regulated in ST infected mice cecum by...
1.5 to 2.9 log folds. The expression of intestinal inflammation-related cytokine genes in LC\textsuperscript{mcra} pretreated mice were manipulated at a positive manner. For example, all 4 pro-inflammatory cytokine genes provoked by ST were suppressed significantly by 1.3 to 5.3 log folds through three weeks after challenging compared to mice with no probiotic protection; expression of anti-inflammatory cytokines IL-10 and TGF-β genes were stimulated notably in comparison with either control or ST infection with no probiotic prevention.

**Modulation of murine gut microbiota due to probiotic pretreatment and salmonellosis**

To compare the gut microbial composition in various groups of mice including pretreated with probiotic strains and/or challenged with ST, we randomly selected their cecal contents (5 mice from each group) for 16S metagenomic sequencing and taxonomic classification. According to the taxonomic profile at the phylum level (Figure 8A), Firmicutes were the predominant phylum (63.51%), followed by Bacteroidetes (29.37%), in fecal samples collected.

**Figure 8.** Mice cecal microbial community phylum-level structure. Bacterial distributions at phylum level in cecal contents from individual pooled dataset were depicted in terms of (a) control mice providing placebo and without ST infection, (b) mice infected with ST, (c) mice daily administered with LC for one week followed by ST challenge, and (d) mice daily administered with LC\textsuperscript{mcra} for one week followed by ST challenge.
from the control group mice which were given placebo (primary control). The relative abundance of Proteobacteria was 0.92% with individual variation from 0.54% to 1.35%. Significant difference in gut microbial community at phylum level was observed in ST infected mice (Figure 8B), specifically the group of mice which were not pretreated with probiotic strains; though the predominant phylum was still Firmicutes (61.26%), the relative abundance of Bacteroidetes was notably decreased to 17.81% and the relative abundance of Proteobacteria boosted to 14.86% (Figure 8B). One-week daily administration of probiotic (LC or LC+mcra) positively shaped the phylum level gut microbiota in the group of mice which were challenged with ST (Figure 8C,D). More specifically, in comparison with ST infected mice which were not pretreated with probiotic (secondary control), the microbial composition of probiotic pretreated mice after challenged with ST were dominated with Firmicutes and they were raised by 5.67 and 13.34% in the groups of mice which were pretreated with LC and LC+mcra, respectively. Whereas in the same pre-treated groups of mice, the relative abundances of cecal Proteobacteria were reduced by 13.17 and 14.17%, respectively (Figure 8C,D).

At genus level, Bacteroides (18.50%) was identified being the highest abundant in cecal contents of primary control group of mice, followed by Ruminococcus (7.17%), Blautia (7.02%), Johnsonella (4.39%), and Lactobacillus (1.80%) (Figure 9). The relative abundances of Salmonella and Enterobacter were observed less than 0.01% of the total gut bacterial composition in the cecal content of the group of mice which were not challenged ST or challenged with ST but pretreated with probiotic strains (Figure 9). Whereas, the gut microbiota genus in ST-infected mice (without pretreatment with probiotic strains) exhibited significantly higher abundances of Salmonella (5.27%) and Enterobacter (3.72%), but lower abundances of Bacteroides (9.98%), Blautia (5.16%), Johnsonella (3.34%), and Lactobacillus (0.17%) (Figure 9). Other gut microbial genus-level noticeable differences between ST-infected mice and control included reduced abundance of Anaerobranca, Anaeroplasm, and Butyrvirbio but increased abundance of Akkermansia, Desulfobacter, Enteroccus, Klebsiella, Leptolyngbya, Natronincola, Staphylococcus, and Tolumonas, Trabulsiella (Figure 9). Compared with secondary control, probiotic pretreatments notably increased the relative abundances of Bacteroides, Blautia, Escherichia, Johnsonella, and Lactobacillus as well as lowered Salmonella, Enterobacter, Klebsiella, Tolumonas, and Trabulsiella. Particularly, the mice which were pre-treated with LC+mcra were partly protected from Salmonella colonization and the relative abundance of Enterobacter was as low as to the control group. Significant escalation of their relative abundances of Bifidobacterium (0.12%), Blautia (8.43%), and Lactobacillus (9.18%) was also detected (Figure 9).

The overall cecal bacterial species diversity was observed minimal in the cecum content of the group of mice which were challenged with ST but not pretreated with probiotic strains but the cecal bacterial species diversity was promoted by LC+mcra pretreatment and protection (Figure 10). Specifically, compared with primary control, the mice group infected with ST exhibited significantly reduced gut intestinal microbial diversity at species level which was indicated by various alpha-diversity indexes including Chao-1, Fisher-alpha, Margalef’s richness, and Simpson (numerically higher), and Shannon (Figure 10). However, the one-week daily pre-administration with LC+mcra, but not with wild-type LC, before ST challenging caused a notably increased bacterial species diversity in cecum compared with secondary control group and even higher in comparison with primary control group.

Discussion

The probiotic strain LC+mcra with 7-fold upregulation in its expression level of mcra gene coding linoleate isomerase has been found with prominently significant 21-fold higher rate in total lino-leic acids production per bacterial cell.22 In a previous study, we revealed in vitro that LC+mcra could competitively exclude the growth and adhesive activity of both ST and EHEC and at meanwhile, suppress their vital virulence gene factors. Moreover, though effectiveness of probiotics in combating enteric bacterial pathogens is still controversial, several researchers have suggested that secondary metabolites of probiotics such as CLA might enhance their overall in vivo health-beneficial functions.16,25–27 Here in the current study, we systematically and in-depth
investigated the double effects of both *Lactobacillus* and CLA on murine gut health. According to our results, 1-week consecutive consumption of LC+mca through oral administration efficiently prevented/eased the following *Salmonella* infection. Although probiotic administration through water might generate variance of bio-availability in mice gut, it is worth mentioning that early-staged oral probiotic gavage possesses high risk in potential induced injury in 3-week-old mouse esophagus. As the balance of bacterial fecal shedding with beneficial and detrimental ratio serves as a key indicator of the gut intestinal health.\(^2^8\) correspondingly we observed in our study the reduced colonization of ST/EHEC in both fecal content and intestinal fluids. Though in this study, we as well as some previous researchers, observed that EHEC were not able to cause enough infection and death in mice, but the *Salmonella* colonization in mice was claimed to be restricted by bioactive probiotics or functional fatty acids oral supplements *in vivo*,\(^2^7,2^9–3^1\) in which the virulence gene factors of *Salmonella* were suggested to be manipulated.\(^3^2,3^3\)

On the other hand, probiotic itself was addressed to be capable of reducing intestinal...
pathogens through physical repelence and colonization resistance.\textsuperscript{34–38} Interestingly, our \textit{in vivo} findings in which we observed that either wild-type or genetically engineered \textit{L. casei} remarkably diminished ST/EHEC colonization in cecum, jejunum, and ileum also agreed with previous reports based on \textit{in vitro} test.\textsuperscript{22} Further, LC\textsuperscript{+mcra} displayed more intensive reductions of colonization of ST/EHEC and/or their severity of infection considering the extraneous strengthening effects implemented by its over-promoted CLA production.\textsuperscript{16} In fact, CLA has been documented and linked with antimicrobial active against several enteric bacterial pathogens including \textit{Salmonella} though the specific mechanism are still under study.\textsuperscript{25,39} Most importantly, the \textit{in vivo} examination based on BALB/cJ mice justified the protective roles of LC\textsuperscript{+mcra} on combating enteric bacterial pathogens, following and matching with previous \textit{in vitro} outcomes relied on various pathogenic bacterial strains.\textsuperscript{22,40,41}

In most cases, \textit{Salmonella} infections are associated with diarrhea, weight loss, dramatic alterations in composition of blood cells, as well as death.\textsuperscript{14,42–44} Accordantly we detected $10^5$–$10^7$ CFU/g intestinal colonization of ST induced salmonellosis and caused around 8% weight loss, 52% higher level of RBC, 19% and 71% lower levels of WBC (especially neutrophils and lymphocytes) and PLT, and severe cecal inflammation in the survival mice. The physical, hematological, and gut intestinal abnormalities mentioned above in our \textit{in vivo} examination contributed in the 40% death rate of mice challenged with enteric bacterial pathogen ST. However, probiotics secreting different types of functional fatty acids initiate attenuation in over-reactive gut inflammation through anti-inflammatory activities,\textsuperscript{16,22,24} which correlates with the LC\textsuperscript{+mcra} (CLA) mediated relative up-regulation of murine intestinal anti-inflammatory cytokine genes from mice under salmonellosis found in our study. Therefore, apart from the

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\caption{Bacterial diversity at species level in murine cecum. The assessment of alpha-diversity including Observed number of taxa species (A), Chao-1 (B), Fisher’s alpha (C), Margalef’s richness (D), Simpson index (E), and Shannon index (F) was determined and analyzed among control mice providing placebo and without ST infection, mice infected with ST, mice daily administered with LC for one week followed by ST challenge, and mice daily administered with LC\textsuperscript{+mcra} for one week followed by ST challenge. Standard deviations among individual group members were provided. Different letters (‘a’ through ‘c’) are significantly different ($p < 0.05$) among groups of control, infection, and infection following probiotic pretreatments.}
\end{figure}
direct colonization competition and repellence, daily administration of probiotics, especially \( \text{LC}^{+\text{mcr}} \), also prevented regular salmonellosis symptoms and maintained the overall physical and gut health condition of mice through mediating immuno-modulation. In future study, several other tissues including kidney, liver, and lung could be examined to further investigate whether \( \text{LC}^{+\text{mcr}} \) pretreatment prevented ST systemic infection.

To address concerns from the host’s point of view, the maintenance of intact and operative gut intestine physiological condition is crucial in both metabolism and symbiotic intestinal microbiota composition. In our study, \( \text{LC}^{+\text{mcr}} \) and its byproduct CLA prevented ST-induced elimination of goblet cells, villi, and microvilli as well as the inflammatory infiltrations between circular folds in cecum, which maintained the overall functions in terms of intestinal nutrients absorption and profoundly raised the survival rate (0 death) in mice. As a matter of fact, CLA has also been previously tested in colitis and inflammatory bovine disease recovery and showed significant gut health benefits, but the specific mechanism was not explained. Here, our findings based on CLA are in support of these researches and suggest a protective mechanism from both bacterial colonization and host histology sides.

A balanced gut microbial ecosystem serves as the crucial defense against colonization and infection with enteric pathogens. \( \text{Salmonella} \) infection could have negatively impacted the gut intestinal microbial composition by diminishing the abundances of Firmicutes including \( \text{Lactobacillus} \) and \( \text{Bifidobacterium} \), and simultaneously favoring the dominance of Proteobacteria inducing follow-up opportunistic infections. In our study, we observed the increasing abundances of \( \text{Salmonella} \) and \( \text{Enterobacter} \) with overall reduced bacterial species diversity following ST challenge in mice, whereas \( \text{LC}^{+\text{mcr}} \) pre-administration successfully prevented the negative shifting of gut microbiota which was induced by ST infection. As a matter of fact, several researchers also observed that CLA-containing diets were able to alter the fatty acids metabolism and maintaining homeostatic gut microflora. The healthier gut microbial distribution as well as their diversity in specific genus and species are influenced by the consumption of CLA-producing probiotic. Particularly at genus level, the higher abundances of \( \text{Lactobacillus} \) and \( \text{Bifidobacterium} \) positively influence the specific gut immune homeostatic programming and signaling pathways which contributes in defending and excluding endogenous or exogenous pathogens. Furthermore, \( \text{Blautia} \) usually participates in nutrient absorption and digestion, which contributes in maintaining host’s gut intestinal nutritional status even the balance of gut is disrupted by pathogenic bacterial infection. On the other hand, reduced abundance of opportunistic pathogenic bacterial colonization like \( \text{Enterobacter} \) and \( \text{Klebsiella} \) minimized the risk of gut microbiota dysbiosis and further systematic infection. A balanced intestinal microbiota strengthen the first-line gut intestinal defense system against multiple pathogenic bacterial infections, which may possess a strong correlation and be the explanation of reduced bacterial pathogen colonization and inflammation in mice gut. A follow-up research on accumulative quantity of CLA in murine gut intestine may help in revealing dynamic interactions among the complex microbial ecosystem including established probiotic strains, their metabolic CLA, endogenous gut commensal flora, and exogenous bacterial pathogens.

Based on previous research, EHEC oral challenge on mice with distinct genetic configuration and gut microbial composition can result in various levels of colonization, morbidity, and mortality. Specifically, EHEC dose as low as \( 10^2 \) CFU led to intense cecal colonization and death in germ-free mice whereas for conventional mice model like BALB/c, considerably higher dose of EHEC was requisite to cause diseases. In some cases, infectious dose of EHEC less than \( 10^{10} \)–\( 10^{11} \) CFU/mouse failed to even introduce cecal colonization and infection, which parallel with our findings. Based on the current study, BALB/cJ mice orogastrically challenged with \( 10^7 \) CFU EHEC were colonized at a range from \( 10^2 \) to \( 10^7 \) CFU/g of EHEC in intestinal fluid collected from cecum, jejunum, and ileum but failed to produce any visible physiological abnormalities or mortality in mice. This finding could be explained by the relative resistance in BALB/c mice toward EHEC through shorter
shedding duration and producing higher serum/fecal levels of O157-specific IgA. On the other hand, LC\(^{+}\)mcra, as we observed in vitro\(^{22}\) and predicted for in vivo, stood out in reducing the colonization level of EHEC as well as preventing from kidney histological abnormalities and weight loss in BALB/cJ mice. Further research dependent on germ-free or compromised commensal flora mouse model might be substantial in revealing how LC\(^{+}\)mcra involved in defending host from EHEC pathogenesis and post-infectious complications.

In conclusion, the current study has demonstrated a substantial influence of CLA over-producing probiotic strain, LC\(^{+}\)mcra exerted on Salmonella and pathogenic E. coli infections in conventional mice. Specifically, mice orally given LC\(^{+}\)mcra daily for one week minimized EHEC colonization and protected themselves from ST-facilitated serious salmonellosis which was observed by notably reduced fecal shedding and intestinal colonization of ST, amelioration on acute inflammation, and prevention on hematological and histological abnormalities. In depth metagenomic analysis revealed that LC\(^{+}\)mcra pretreatment modulated mice cecal bacterial community with increased diversity which are predominated with comparative higher Firmicutes and lower Proteobacteria. The outstanding protective roles of LC\(^{+}\)mcra against ST and EHEC infection plus its profound effectiveness over wild-type LC may provide a promising option for prophylaxis on pathogenic Salmonella and diarrheagenic E. coli infections and reduce enteric bacterial infections.

Methods

Bacterial strain and growth conditions

*Lactobacillus casei* (LC, ATCC 334) and our laboratory generated linoleic acid over-expressed *L. casei, LC\(^{+}\)mcra\(^{22,40}\)* were used as probiotics while *Salmonella enterica* serovar Typhimurium (ST, ATCC 14028) and enterohemorrhagic *Escherichia coli* EDL933 (EHEC, ATCC700927) were chosen as enteric bacterial pathogens in this study. *Lactobacillus* strains LC and LC\(^{+}\)mcra were grown on regular MRS agar and MRS agar containing 100 mg/mL 5-fluorouracil separately at 37°C for 24 h in the presence of 5% CO\(_2\) (Forma\(^{\text{TM}}\) Scientific CO\(_2\) water jacketed incubator, Thermo Fisher Scientific Inc., Waltham, MA, USA).\(^{22}\) ST and EHEC were grown on LB agar (EMD Chemicals Inc., Gibbstown, NJ, USA) for 18 h at 37°C under aerobic conditions (Thermo Scientific MAXQ 4450, Thermo Fisher Scientific Inc., Waltham, MA, USA).

Mouse model and animal experiments

The 3-week-old BALB/cJ Mice (approximately 8–10 g) were purchased from The Jackson Laboratory (Bar Harbor, ME USA) and reared in static micro-isolating cages with cellulose Bio-Performance bedding and huts as environmental enrichment. Teklad standard rodent diet and regular tap water were provided for mice feeding and drinking, respectively. A total of 90 mice (45 males and 45 females) were used for each trial. Following a completely randomized method, 90 mice were randomly assigned to 9 groups (designated A1 to C3) resulting in 10 mice per group; two cages were assigned to each group with a total of 5 mice per cage. Mice cages were changed weekly, and each individual mouse was weighed and monitored with health examinations daily. At the end of the second, third, and fourth week, 3, 3, and 4 mice from each group respectively, were randomly selected and euthanized with CO\(_2\) inhalation in euthanasia chamber for organ samples collection.

Feeding probiotic to BALB/cJ mice and challenging with ST and EHEC

Overnight culture of LC or LC\(^{+}\)mcra in MRS broth were diluted in fresh 5 mL MRS broth at 1:50 and allowed for 3 h further growth. The bacterial cells in exponential phase were harvested following centrifugation at 3,000 × g for 15 min, PBS washing, and resuspension in 1.0 mL PBS. A final concentration of 10\(^{11}\) CFU/mL was adjusted with PBS and used to feed mice. The design of in vivo mouse trial was summarized in Table 1. Probiotic (either 10\(^9\) CFU/mL LC or LC\(^{+}\)mcra) cells were maintained in water bottle filled with regular tap water for group B and C and fed to mice from Day 1 to Day 7. Control mice, group A, was fed with regular tap water only.

Overnight culture of ST and EHEC bacterial cells in LB broth were diluted in fresh 5 mL LB
broth at 1:50 and allowed for 3–4 h further growth at 37°C. The exponential phase bacterial cells were harvested and washed by centrifugation at 3,000 × g for 15 min and resuspended in 1.0 mL of PBS. A final concentration of bacterial cells was adjusted to 10^8 CFU/mL in PBS. On Day 7, an aliquot of 100 µL ST or EHEC suspension containing approximately 10^7 CFU was fed to mice in groups 2 or 3 respectively, with oral gavage, and the mice were reared thereafter for another 3 weeks. Mice in group 1 was orogastrically fed with 100 µL PBS and served as control.

### Sample collection and processing

In order to estimate the bacterial fecal shedding, fecal samples were collected from each mouse in sterile Whirl-Pak bags using sterile spoons at Day 1, 2, 3, 6, 7, 8, 9, 14, 21, and 28 for PBS serial dilution and plating on specific agar plates. In order to investigate the bacterial colonization in mice gut, intestine, ileum, jejunum, and cecum from each euthanized mouse were separated and harvested. Then the ileum, jejunum, and cecal fluids were serial diluted with PBS, followed by plating on specific agar plates. Specifically, MRS agar for LC, MRS agar containing 100 mg/mL 5-fluorouracil for LC+ mcra, XLT-4 agar for ST, and MacConkey agar for EHEC were used, respectively.

Mice cecum was kept in RNA Later for further RNA extraction, cDNA reverse transcription, and inflammation-related gene expression level analysis. For hematological analysis, the blood samples from each mouse was collected from heart in VACUETTER® Heparin tubes (Greiner Bio-One, Monroe, NC, USA) and further analyzed with a ProCyte Dx® Hematology Analyzer (IDEXX, Westbrook, ME, USA) according to the manufacturer’s instructions.

### RNA extraction, cDNA synthesis, and quantitative RT-PCR

Extraction of mice intestinal RNA was carried out using TRIzol® Reagent (Life Technologies Co., Carlsbad, CA, USA) following previous methods. The cDNA synthesis was performed according to the manufacture’s instruction of qScript cDNA SuperMix. The PCR reaction mixture containing 10 µL PerfeCTa SYBR Green Fast Mix (Quanta Biosciences, Beverly, MA, USA), 2 µL of each 100 nM primer (Table 2), 2 µL of cDNA (10 ng), and 4 µL of RNase-free water was amplified using an Eco Real-Time PCR system.

| Gene   | Primer | Sequence (5’-3’) | Function                  |
|--------|--------|------------------|--------------------------|
| 18S rRNA | Sense | TTAGAGTTGTTCAAGACGGCCCGA | Housekeeping gene |
| IL-1β  | Sense | TCTTGGCAAATGCCTGCTTG | Inflammatory cytokine gene |
| IL-6   | Sense | ACACATGTTTCTCGGAGAAAATCCTGG | Pro-/Anti-inflammatory cytokine gene |
| IFN-γ  | Sense | TCTGCAAGTGCAACATGTGCTTG | Anti-inflammatory cytokine gene |
| TGF-β  | Sense | ATGCATGACGTGGCTGGCTGG | Pro-inflammatory cytokine gene |
| TNF-α  | Sense | GGCAACAGCTTCGGGATAATG | Anti-inflammatory cytokine gene |

### Table 1. Mice groups, numbers per group, and their treatment/infection.

| Group (#) | Mice (n) | Probiotic Treatment (daily during 1st week) | Pathogen challenge (beginning of 2nd week) |
|-----------|----------|-------------------------------------------|------------------------------------------|
| A1        | 10       | PBS LC1                                   | PBS ST3                                   |
| B1        | 10       | PBS LC1                                   | EHEC                                      |
| C1        | 10       | PBS LC1                                   | EHEC                                      |
| A2        | 10       | PBS LC1                                   | EHEC                                      |
| B2        | 10       | PBS LC1                                   | EHEC                                      |
| C2        | 10       | PBS LC1                                   | EHEC                                      |
| A3        | 10       | PBS LC1                                   | EHEC                                      |
| B3        | 10       | PBS LC1                                   | EHEC                                      |
| C3        | 10       | PBS LC1                                   | EHEC                                      |

1 Wild-type L. casei; 2 Mutant L. casei; 3 S. Typhimurium; 4 Enterohemorrhagic E. coli
Table 3. Hematological changes and comparison of mice with and without *Salmonella* infection.

| Day | Control | Infection | LC | LC+mcra | Control | Infection | LC | LC+mcra | Control | Infection | LC | LC+mcra |
|-----|---------|-----------|----|---------|---------|-----------|----|---------|---------|-----------|----|---------|
| 14  | 10.15 ± 0.14<sup>a</sup> | 13.03 ± 4.20<sup>b</sup> | 10.84 ± 0.55<sup>b</sup> | 10.60 ± 0.46<sup>b</sup> | 3.48 ± 0.35<sup>a</sup> | 2.95 ± 0.80<sup>b</sup> | 3.12 ± 0.61<sup>a</sup> | 3.55 ± 0.42<sup>a</sup> | 7 ± 3<sup>a</sup> | 2 ± 2<sup>c</sup> | 5 ± 2<sup>b</sup> | 7 ± 3<sup>a</sup> |
| 21  | 10.34 ± 0.21<sup>b</sup> | 14.89 ± 3.87<sup>a</sup> | 10.72 ± 0.65<sup>b</sup> | 10.51 ± 0.33<sup>b</sup> | 3.65 ± 0.25<sup>a</sup> | 2.96 ± 0.40<sup>b</sup> | 3.13 ± 0.69<sup>a</sup> | 3.54 ± 0.38<sup>a</sup> | 8 ± 1<sup>a</sup> | 2 ± 1<sup>c</sup> | 5 ± 2<sup>b</sup> | 8 ± 3<sup>a</sup> |
| 28  | 10.29 ± 0.13<sup>b</sup> | 15.64 ± 4.37<sup>a</sup> | 11.84 ± 2.14<sup>b</sup> | 10.71 ± 0.53<sup>b</sup> | 3.59 ± 0.22<sup>a</sup> | 2.90 ± 0.49<sup>b</sup> | 3.26 ± 0.30<sup>a</sup> | 3.51 ± 0.29<sup>a</sup> | 7 ± 2<sup>a</sup> | 2 ± 1<sup>c</sup> | 4 ± 2<sup>b</sup> | 8 ± 2<sup>a</sup> |

<sup>1</sup>Mice with no probiotic protection infected with *Salmonella* Typhimurium
<sup>2</sup>Mice pre-administered with wild-type *Lactobacillus casei* followed by *S.* Typhimurium infection
<sup>3</sup>Mice pre-administered with mutant *L. casei* followed by S. Typhimurium infection

*Means with different letters (a-c) for each type of blood cells (red blood cell, white blood cell, and platelets) in different groups (control, infection, infection following LC pretreatment, and infection following LC<sup>mcra</sup> pretreatment) at individual time point (day 14, 21, and 28) are significantly different at p < 0.05
observations were performed under a light microscope (BA210E, Motic Asia, Hong Kong, China).

**Metagenomic sequencing and analysis**

Mice cecal contents were harvested during the 4th week (Day 22–28) and 5 samples from each group of control (PBS only), ST infection, LC pretreatment followed by ST infection, or LC$^{+mcra}$ pretreatment followed by ST infection were randomly selected for metagenomics analysis. Microbial genomic DNA extraction was carried out using QIAamp Fast DNA Stool Kit (QIAGEN, Valencia, CA, USA) following instructions from the manufacturer. The variable V3 and V4 regions of microbial 16S rRNA gene were targeted for phylogenetic classifications. DNA libraries were prepared for equimolar-pooling using Nextera DNA Library Preparation Kit and Nextera Index Kit (Illumina, San Diego, CA, USA) according to the manufacturer’s instructions. Paired-end sequencing (2 × 300 bp) was conducted on Illumina MiSeq using MiSeq v3 600-cycle kit (Illumina, San Diego, CA, USA). Sequence data was processed through MiSeq Reporter – BaseSpace for FASTQ Workflow generation followed by taxonomic

### Table 4. White blood cell counts in mice with and without *Salmonella* infection.

| White Blood Cells | Group                          | 14  | 21  | 28  |
|-------------------|-------------------------------|-----|-----|-----|
| Neutrophils       | Control                       | 1.31 ± 0.40$^a$ | 1.34 ± 0.40$^a$ | 1.50 ± 0.21$^a$ |
|                   | Infection1                    | 0.78 ± 0.36$^b$ | 0.41 ± 0.22$^b$ | 0.27 ± 0.14$^b$ |
|                   | LC1                           | 1.02 ± 0.30$^{ab}$ | 1.07 ± 0.37$^{ab}$ | 1.08 ± 0.48$^{ab}$ |
|                   | LC$^{+mcra}$ 3                | 1.27 ± 0.56$^a$ | 1.16 ± 0.11$^{ab}$ | 1.20 ± 0.23$^{ab}$ |
| Lymphocytes       | Control                       | 2.19 ± 0.22$^a$ | 2.50 ± 0.13$^a$ | 2.44 ± 0.41$^a$ |
|                   | Infection                     | 1.18 ± 0.25$^b$ | 0.92 ± 0.15$^b$ | 0.73 ± 0.18$^b$ |
|                   | LC1                           | 2.05 ± 0.78$^a$ | 2.03 ± 0.13$^a$ | 1.82 ± 0.26$^a$ |
|                   | LC$^{+mcra}$ 3                | 2.37 ± 0.37$^a$ | 2.53 ± 0.15$^a$ | 2.46 ± 0.52$^a$ |
| Monocytes         | Control                       | 0.04 ± 0.01$^a$ | 0.04 ± 0.02$^a$ | 0.04 ± 0.02$^a$ |
|                   | Infection                     | 0.16 ± 0.04$^a$ | 0.14 ± 0.06$^a$ | 0.19 ± 0.04$^a$ |
|                   | LC1                           | 0.05 ± 0.02$^a$ | 0.06 ± 0.03$^a$ | 0.08 ± 0.03$^a$ |
|                   | LC$^{+mcra}$ 3                | 0.05 ± 0.02$^a$ | 0.05 ± 0.02$^a$ | 0.05 ± 0.01$^a$ |
| Eosinophils       | Control                       | 0.06 ± 0.04$^a$ | 0.07 ± 0.03$^a$ | 0.07 ± 0.03$^a$ |
|                   | Infection                     | 0.16 ± 0.03$^a$ | 0.16 ± 0.04$^a$ | 0.17 ± 0.05$^a$ |
|                   | LC1                           | 0.09 ± 0.04$^a$ | 0.11 ± 0.01$^a$ | 0.09 ± 0.03$^a$ |
|                   | LC$^{+mcra}$ 3                | 0.07 ± 0.03$^a$ | 0.06 ± 0.03$^a$ | 0.06 ± 0.04$^a$ |
| Basophils         | Control                       | 0.01 ± 0.01$^c$ | 0.01 ± 0.01$^c$ | 0.01 ± 0.01$^c$ |
|                   | Infection                     | 0.03 ± 0.01$^a$ | 0.03 ± 0.02$^a$ | 0.04 ± 0.02$^a$ |
|                   | LC1                           | 0.02 ± 0.01$^b$ | 0.02 ± 0.01$^b$ | 0.02 ± 0.01$^b$ |
|                   | LC$^{+mcra}$ 3                | 0.01 ± 0.01$^c$ | 0.01 ± 0.01$^c$ | 0.01 ± 0.01$^c$ |

*Means with different letters (a–c) for each type of WBC (neutrophils, lymphocytes, monocytes, eosinophils, and basophils) in different groups (control, infection, infection following LC pretreatment, and infection following LC$^{+mcra}$ pretreatment) at individual time point (day 14, 21, and 28) are significantly different at $p < 0.05$.

**Histopathology analysis**

Intestinal tissue samples were taken from mice after euthanization and were stored in neutral buffered formalin (4% formaldehyde; pH 7.4) at 4°C for further processing. Once the samples were removed from fixative, they were dehydrated with increasing concentrations of ethanol, cleared in xylene, and embedded in paraffin. Microtome (LEICA RM2065, Leica Biosystems, Buffalo Grove, IL, USA) was used to harvest 5 μm thick paraffin sections followed by heat fixing at 37°C overnight. Then the slices were stained with hematoxylin and eosin and mounted with DPX mounting medium 13512 (Electron Microscopy Sciences, Hatfield, PA, USA). Histological
classification based on Greengenes database (http://greengenes.lbl.gov/). Demultiplexing was performed using only perfect index recognition (mismatch = 0) followed by removing PhiX reads. 16S sequence length below 1250 bp or with more than 50 wobble bases was filtered, and all entries classified with no genus or species were also filtered. The relative abundances and alpha/beta-diversity indices were calculated using ‘vegan’, ‘phyloseq’, and ‘metagenomeseq’ R package and plotted in Excel.

**Statistical analysis**

All data were analyzed by the SPSS software. Comparison among multiple mice groups were performed with the one-way analysis of variance followed by Tukey’s and Bonferroni’s tests. Survival rate among multiple mice groups were analyzed with Log-Rank test. For all tests, significant differences were considered on the basis of p values below a significant level of 0.05.

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**Contributors**

MP conceived the study, designed and carried out the experiments, analyzed the data, and wrote the manuscript. ZT contributed to in vivo experiments and manuscript draft. PP and CBe assisted in experimental preparation, sample processing, and microbiological experiments. CBi participated in reviewing and editing the manuscript. JM provided sequencing resources and helped in metagenomics. DB supervised the research and finalized the manuscript. All authors approved the final manuscript.

**Ethics approval and consent to participate**

Mice in vivo experiments were performed in ABSL2 facilities in Department of Animal and Avian Sciences, University of Maryland in accordant with protocol #R-NOV-17-55 approved by the Institutional Animal Care and Use Committee (IACUC). The best effort was made for minimizing the suffer of animals. To ensure animal welfare, mice were monitored and recorded for physical appearance and body weight on a daily basis during experimental period.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

**Competing interests**

The authors declare that they have no competing interests.

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