Gram Positive Bacterial Lipoteichoic Acid Role in a Root Canal Infection - A Literature Review

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Abstract

Bacteria and its by-products are found to be the main cause of pulpal and periapical infection of tooth. Infected root canals of tooth harbours a wide variation of microbial flora that includes both Gram-positive and Gram-negative microorganisms. Bacterial components such as Lipopolysaccharide (LPS) of gram negative bacteria and Lipoteichoic Acid (LTA) of gram positive bacteria have the potential to enter the peri-apical tissue of tooth and initiate the inflammatory process. After microbial death that occurs either due to body's defence cells or by antibiotic action, bacterial cell wall components such as LTA are released which can persist inside macrophages for prolonged periods causing chronic inflammation. Once these cell-wall components are recognized by the body immune surveillance cells, numerous inflammatory mediators are released leading to inflammation and subsequent pathological consequences. The purpose of this review is intend to summarize the role of gram positive bacterial component LTA in causing endodontic infection and use of potential therapeautic agents against LTA.

Keywords: Lipoteichoic acid (LTA), endodontic infection, Enterococcus faecalis, lipoteichoic acid

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INTRODUCTION

Evidence based treatment is the new paradigm in global health care. A wide variation of microbial flora is normally present in infected root canals of tooth. A thorough understanding of the microbiota associated with pulp pathology is the basis for the success of endodontic treatment. The microbiota involved in endodontic treatment failure are facultative anaerobes such as Enterococcus, Actinomyces species and Candida. Gram positive Enterococcus faecalis (E.faecalis) is commonly found in high percentage in sites exhibiting root canal failures and it survives as a single organism or as a major component of the flora in the root canals of tooth. Bacterial virulence factors which are primarily involved in various stages of endodontic as well as peri-apical infections promotes bacterial colonization and activation of host defence mechanisms. Among various virulence factors of gram positive bacteria, Lipoteichoic acid (LTA) which has pathogenic potential similar to that of gram-negative bacteria virulence factor Lipopolysaccharide, plays a key role in serious inflammatory responses1,2,3.

Teichoic acids (TAs) are specific polymers present on gram-positive bacterial cell surfaces. The word “teichoic” originates from the Greek word teikhos, meaning “wall”3. Wall-teichoic acid (WTA) are those Teichoic acids (TA’s) that are attached to the cell wall, where as those attached to the cell membrane are called as lipoteichoic acid (LTA). Armstrong et al was the first person to observe WTA in 1958 from cell wall fractions of Lactobacillus plantarum, Bacillus subtilis, and Staphylococcus aureus4. LTAs were identified by Kelemen et al. in 1961 in cell membrane fractions5.

LTA structure and function varies across different gram positive bacterial species. The backbone of TA’s are made of polymers consisting of repeating polyolphosphate units. The backbone of LTA’s are composed of poly-glycerol phosphate, and the WTA backbone are composed of poly-ribitol phosphate6. LTA are amphiphilic polysaccharides covalently linked to a terminal glycolipid moiety; and are released either during bacterial multiplication or after bacteriolysis caused by beta-lactam antibiotics or bactericidal cationic peptides, phospholipase A, cathepsins etc.7,8. LTA released after cell lysis binds specifically to target cells CD14 and to Toll-like receptors or binds non-specifically to membrane phospholipids. LTA bound to target cells causes activation of complement system and release of proinflammatory cytokines, reactive oxygen and nitrogen species from neutrophils and macrophages leading to indirect tissue injury7. The role of LTA in cell-to-cell interactions in biofilms, its role in causing cell envelope homeostasis were studied using the LTA deficient strain of Streptococcus gordonii9. LTA synthase(LtaS) enzyme inhibitor was found to prevent the growth of gram-positive antibiotic-resistant S. aureus in mice model9.

The present review attempts to summarize the immunostimulatory potential of gram-positive bacterial component LTA in causing endodontic infection and use of potential therapeutic agents against LTA inflammatory action.

Data source

A review of literature was performed by electronic search of Pubmed and Google Scholar for publications on LTA studies carried out mainly in relation to field of dentistry. Literature search was done using the key words - Lipoteichoic acid, endodontic infection, Enterococcus faecalis lipoteichoic acid, and endodontic medicaments. Articles were chosen using the following inclusion criteria: articles published in English language from 1953- January 2020, studies related to LTA role in endodontic infection were considered mainly. A total of 60 articles related to the field of Endodontics involved in studying the role of LTA in activating immune system in root canal and periapical infection were considered. Structure and role of LTA of those bacteria not exhibiting a potential role in endodontic infection were not considered.

Role of Lipoteichoic Acid in Causing Tissue Injury

Literature search shows that LTA can activate classic and alternative pathways of the complement system, activaties macrophages to release inflammatory mediators and causes several tissue alterations including bone resorption10. Various in vitro studies have shown variations in immunostimulatory potential of LTA and this variations can be avoided by more gentle high purification process of LTA extraction which can maintain the immunostimulatory potential11. LTA isolated from E. faecalis stimulates leukocytes to release inflammatory mediators like tumor
necrosis factor alpha (TNF-α), nitric oxide (NO) and chemokines through the Toll-likereceptor2 (TLR2) signaling pathway in macrophages. It also has role in biofilm formation that enables bacteria to survive in adverse environments12, 13. Study carried out by Costa Junior et al has shown that *Streptococcus faecalis* LTA provoked a chronic inflammatory tissue reaction and can contribute to formation of a periapical lesion10.

LTA activates innate immune system which acts by detection of pathogens through specialized pattern recognition receptors like RIG-I-like receptor, the toll-like receptor family (TLR), and the nucleotide binding oligomerization domain (NOD)-like receptor (NLR) present in the cell membrane of immune and non-immune cells14. Among the pattern recognition receptors, TLR are key participants. Both LPS and LTA binds to CD14 receptor and activates signalling by TLR and induces release of proinflammatory cytokines such as Tumor Necrosis Factor alpha (TNF-α), Interleukin-1 (IL-1), Interleukin-8 (IL-8), Interleukin-6, and anti-inflammatory cytokines like interleukin-10 and interleukin-122,15. Studies have also shown that LTA causes release of prostaglandin E2 (PGE2), lysosomal enzymes and the generation of superoxide anion. These factors which have tissue-damaging (TNF-α, IL-1beta, IL-6, PGE2, lysosomal enzyme, superoxide anion) have been detected in periapical samples of patients with root canal infection12, 16, 17.

TNF-α is the most abundant cytokine and first bone resorptive mediator that has been detected in endodontic infection18. TNF-α induces osteoclast activation and also stimulates secretion of proteolytic enzymes and matrixmetalloproteinases (MMP), which are responsible for destroying extra cellular matrix of the bone tissue19, 20. In an experimental study conducted by Wang et al using clinical strain of *E. faecalis*, the authors concluded that *E. faecalis* LTA promoted pro-inflammatory cytokine TNF-α expression in macrophages by activating p38 Mitogen-activated protein kinases and nuclear factor kappa B (NF-κB) signalling pathways, but did not affect anti-inflammatory cytokines expression 21. Transcription factor NF-κB is involved in cytokine synthesis and TLRs activate the effector phase of the innate immune response mainly through NF-κ B pathway. In response to microbial infection, NLRP3 inflammasome that plays a key role in innate immunity, mediates caspase-1 activation which is essential for cleavage of pro-IL-1β and pro-IL-18 into their active mature form.

**Fig. 1.** LTA binds to CD14 receptor and are subsequently presented to Toll – like receptor (TLR). This stimulates the formation of a signalling complex that involves MyD88, IL-1 receptor associated kinase (IRAK), and tumour necrosis factor receptor-associated factor 6 (TRAF 6). Signalling of this complex leads to activation of pathways that regulate NFκB nuclear factor -κB; MAP: mitogen – activating protein leading to release of pro inflammatory cytokines. A primary Signalling pathway through NFκB causes expression of NLRP3 and pro-IL-1β. A second signal PAMPs (Pathogen-Associated Molecular Patterns) and DAMPs (Danger-Associated Molecular Pattern) activates NLRP3 inflammasome assembly that leads to activation of caspase-1 and the cleavage and secretion of active IL-1β . 14,22
A study carried out by Wang et al revealed that during periapical periodontitis, *E. faecalis* LTA directly induces NLRP3 inflammasome expression associated with the activation of the NF-κB signalling pathway, activating high expression of caspase-1 and release of IL-1β, consequently leading to periapical inflammation (Fig. 1).

Konrad Tietze et al. in their study found that gram-negative species produced high levels of TNF-α, while gram-positive species released higher quantities of the chemokine, IL-8. This difference might be due to the reason that gram-negative and gram-positive bacteria activates different TLRs. TLR-2 appears to be the primary mediator of the innate immune response to LTAs from various gram-positive bacteria.

Experimental study carried out by Baik et al indicates that highly pure and structurally intact *E. faecalis* LTA stimulates TLR2 preferentially rather than TLR4. Predominant stimulation of TLR2 by gram-positive bacteria can also be influenced by the method of bacterial killing, such as heat killing or ethanol killing, since heat killing process can denature and inactivate proteinaceous antigens capable of stimulating TLR4. Studies have also stated that heat-killed *E. faecalis* has a unique feature that it was able to stimulate both TLR2 and TLR4, possibly because *E. faecalis* may possess heat-stable constituents which has the potential to stimulate TLR4. Despite the stimulation of both TLR2 and TLR4 by *E. faecalis*, TLR2 is more important than TLR4 in the dentine inflammation because numerous TLR2-positive macrophages were detected, and the expression of TLR2 rather than TLR4 was greatly induced in inflamed dental pulp tissues.

*E. faecalis* LTA increased macrophages autophagy which can have potential effect on process of periapical infection. *E. faecalis* LTA inhibits osteoclast differentiation from bone marrow derived macrophages through TLR2 signalling pathway, and also causes the macrophages to maintain their phagocytic potential and their inflammatory capacities, possibly contributing to refractory apical periodontitis. Similar results were obtained in another study where *E. faecalis* LTA effectively inhibited osteoclastogenesis of bone marrow macrophages in the presence of RANK via the transcription factor RBP-Jκ. The receptor activator of nuclear factor-KB ligand (RANKL)/osteoprotegerin (OPG) are the prime regulators in bone resorption and have main functional role in the differentiation, activation and survival of osteoclasts and osteoclast precursors. RANKL triggers osteoclastogenesis thereby increasing bone loss, whilst OPG has the opposing effect. The results of study carried out Zhao et al shows that *E. faecalis* LTA upregulated the ratio of RANKL/OPG in a dose-dependent way and that only a lower concentrations was required to stimulate RANK.

Yaguang Tian et al assessed the role of *E. faecalis* LTA on the apoptosis of osteoblasts and stated that the apoptotic effect of *E. faecalis* LTA on osteoblast was in a dose-dependent manner and LTA can inhibit the proliferation of osteoblast-like cells. Periapical lesion may have an epithelial lining which acts as a barrier preventing the host cells from getting affected by slowing down the invasion of bacteria and their products into the host tissues. *E. faecalis* LTA has the ability to disrupt this epithelial barrier and affect osteoblasts in the periapical region.

LTA up-regulates the expression of TLR2 and it was found that TLR2 activated odontoblast exhibits a down regulation in its special function of dentin matrix synthesis and mineralization and in turn these TLR2 activated odontoblast orient its activity on initiating innate immune response by secreting chemokines. An increased amount of inflammatory chemokine CCL2 and CXCL10 was found in supernatants from LTA-stimulated odontoblasts.

In the work done by Wang PL et al, the authors stated that *Streptococcus mutans* LTA causes apoptosis of human dental pulp cells which is important for initiation and progression of pulpitis. LTA causes rapid rise of macrophage Vascular Endothelial Growth Factor (VEGF) expression that may lead to pulpal necrosis or even to the prolonged maintenance of the inflammatory reaction after root canal therapy. In the experimental study carried out using *Streptococcus mutans* LTA, it was found that very low concentrations of LTA itself can stimulate macrophages in the dental pulp to secrete VEGF. VEGF is a strong inducer of angiogenesis and also increases vascular permeability. VEGF expression in dental pulp can cause increased intrapulpal pressure and pain.

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In a study carried out to evaluate time dependent expression of complement 5a and its receptor C5aR, it was found that LTA at a concentration of 1 µg/ml stimulates human dental pulp cells to activate the expression of C5a which was highest at 48 hours. Following initiation of inflammation, complementary protein fragments such as C3a, C4a and C5a are formed which causes degranulation of basophils/mast cells and release high amount of histamine, serotonin and vascular active mediators. The study also states that with in the timeline considered, the proinflammatory effect of LTA is stronger which is indicated by higher expression of IL-6 by LTA than does LPS. Streptococcus mutans LTA were found to have high affinity for hydroxypatite, suggesting LTA’s significant role in colonization of gram positive bacteria on dentinal surfaces.

The viable but nonculturable state is a survival mechanism adopted by bacteria when exposed to adverse environmental conditions and LTA has its role during this period as it was found that E.faecalis LTA quantity was doubled during this state. During this state bacteria loses its growth potential but retains its viability and pathogenicity and start its regular division upon restoration of normal growth conditions.

Werner Fischer defined polymeric chains associated with bacterial membranes using the generic term “amphiphile” and stated five structural types of LTA (I-V). In the structure of E. faecalis LTA the hydrophilic chains are repetitious 1,3-polyglycerol phosphates and the C-2 position of the glycerol residues are nonstoichiometrically substituted with D-alanine and kojibiose residues. E. faecalis LTA belongs to classical Type 1 LTA whose structure shows D-alanine esters substitute and this confer positive charges onto an otherwise negatively-charged polymer. Study carried out by Fabretti et al shows that LTA has a potential functional role in biofilm formation of E. faecalis, and an anti-LTA antibody affects biofilm formation. The effect of D-alanine esters of teichoic acids on biofilm formation and adhesions have also been evaluated. The dlt operon, which causes expression of the protein that incorporates D-alanine into LTA, consists of 4 genes: dltA, dltB, dltC and dltD and inactivation of genes within dlt operon leads to total absence or decrease in D-alanine esters. The presence of D-alanine seem to be closely related to the inflammatory potential of LTA and absence of D-alanine residue affects the virulence of E. faecalis. D-alanylated LTA has the potential to restrict the action of autolysins and cell wall active antibiotics, and this helps bacterial envelop integrity to be maintained. Similar results were obtained by Poyart et al where a mutant strain of Streptococcus agalactiae which was lacking D-Alanine moiety on the LTA, exhibited a decrease in its virulence and was easily killed by macro phages and neutrophils. Cetyltrimethylammonium bromide and chlorhexidine showed improved antimicrobial action on lactobacillus casei in which the gene dlt D was inactivated.

Bovine mammary epithelial cells when stimulated with LTA resulted in upregulation of differentially expressed genes IL-8, CXCL6, CCL2 and IL-1α, proving that LTA play a role in initial inflammatory response. LTA from pathogenic and non-pathogenic bacteria differ in their immunostimulatory abilities. LTA from pathogenic bacteria such as S. aureus induces production of NO and TNFα with effective TLR2 activation, whereas LTA from Lactobacillus plantarum suppresses TLR2 activation by attenuating Pam2CSK4 ligand induced IL-8 production in epithelial cells. Considering these findings, it can be considered that LTA has a potential role in resistance against adverse conditions and also against root canal medicaments.

**Therapeutic Medicaments in Endodontics to Neutralise Pathological Lipoteichoic Acid Activity**

Antimicrobial agents are usually preferred for bacterial infection, but they are often not enough to obtain good treatment outcome, since there are only few effective endotoxin-neutralizing agents. The overproduction of inflammatory mediators by pathogenic bacteria can be decreased by using agents that target LPS and LTA. Studies are being carried out with various endodontic medicaments that can bind to and sequester LTA and prevent excessive inflammatory response. Table 1 gives an insight about various medicaments.

Stannous fluoride has clinical gingivitis benefits as it decreases the pathogenicity of plaque biofilms by blocking reactivity of LPS and LTA ligands with tissue receptors that are involved


Table 1. Summary of studies carried out to evaluate LTA neutralizing action by various therapeutic agents used

| Authors & Year | Study Purpose | Conclusion |
|----------------|---------------|------------|
| MatejaZorko et al 2008 [46] | Study evaluated the effect of alexidine and chlorhexidine on LTA and LPS | Study result showed that chlorhexidine had slightly better LPS neutralization action, while both chlorhexidine and alexidine showed similar LTA-neutralizing activity. Both chlorhexidine and alexidine had the potential to neutralize immunostimulatory cell-wall components released after bactericidal activity of antibiotics. |
| Jung EunBaik et al 2008b [47] | Study was carried out to evaluate the role of calcium hydroxide in inactivating E. faecalis LTA | The result of the study shows that inflammatory activity of LTA is inactivated by calcium hydroxide by affecting its TLR2-stimulating activity resulting in reduced TNF-α expression. |
| Jin-Kyung Lee et al 2009 [48] | Study evaluated if Chlorhexidine decreases the inflammatory potential of Enterococcus faecalis and its virulence factor, Lipoteichoic acid | Chlorhexidine can inactivate LTA of E. faecalis leading to a decrease in inflammatory responses. To totally neutralize LTA’s inflammatory potential, chlorhexidine greater than 0.2% concentration was needed for 24 hours of treatment time and with 2% chlorhexidine a treatment time more than 6 hours was needed. |
| Jung EunBaik et al 2011 [49] | Study was carried out to evaluate if calcium hydroxide deacylates E. faecalis LTA resulting in the loss of immunostimulating activity | Results of study states that calcium hydroxide could deacylate the glycolipid moiety of E. faecalis LTA, leading to impairment of immunostimulating activity. |
| Sung-Hoon Lee et al 2012 [50] | Study was carried out to investigate the antimicrobial and neutralizing effects of 4 Human β-Defensins (HβD 1,2,3,4) (HβD - cationic antimicrobial Peptides that plays a role in protecting the host against infectious microbes) on induction of proinflammatory cytokines by E. faecalis and E. faecalis LTA. | It was found that HβDs had an antibacterial effect against E. faecalis. All tested HβDs inhibited E. faecalis LTA induced expression of proinflammatory cytokine and intercellular adhesion molecule 1. HβD-3 showed strongest bactericidal and neutralizing effect against E. faecalis and E. faecalis LTA. |
| Reference | Description |
|-----------|-------------|
| Sun Woong Hong et al. 2016 [51] | Study was carried out to investigate whether NaOCl inactivates *E. faecalis* LTA. Treatment of *E. faecalis* LTA with NaOCl causes a reduced production of inflammatory mediators by LTA, as NaOCl deacylates the glycolipid moiety of *E. faecalis* LTA structure. |
| Wei Yin et al. 2020 [52] | Study evaluated effect of dioscin, an anti-oxidative stress drug, on the *E. faecalis* originated LTA. Results showed that LTA induced activation of NLRP3 inflammasome in mouse macrophages was inhibited by dioscin induced activation of NLRP3 inflammasome in mouse Macrophages. |

Effect of LTA against Bacterial biofilm

It is stated that LTA is involved in biofilm formation. However, LTA of few bacterial species like *Lactobacillus plantarum* which is a known probiotic agent in oral health has shown to have inhibitory effect on *Staphylococcus aureus* biofilm formation[66]. Biofilm formation of multispecies bacteria was decreased in dose dependent way by *Lactobacillus plantarum* LTA (Lp.LTA). Intracanal medicaments efficacy in removal of preformed multispecies biofilm was aided by *Lactobacillus plantarum* LTA[57]. Disruption of *E. faecalis* biofilms on dentin blocks were enhanced when dentin blocks were pretreated with Lp. LTA and then followed by endodontic medicaments use, than when using Lp. LTA or medicament alone, or when using both at the same time[58]. Among various Lactobacillus species studied, Lp. LTA has a potential biofilm blocking effect and this inhibition is seen not only on *E. faecalis* biofilm but also on other bacterial biofilms[59]. Compound 1771 that modulates LTA metabolism inhibited *Streptococcus mutants* biofilm formation and this effect was more pronounced when it was used along with myrcitin and fluoride.[60]

**CONCLUSION**

LPS is the commonly investigated bacterial virulence factor but taking into account that endodontic infections are polymicrobial in nature studies can also be focused on influence of other bacterial virulence factors. Little is known in the endodontic literature regarding the LTA of gram-positive bacteria in teeth with regard to its role to clinical finding. With this consideration further research related to LTA structure of common gram positive endodontic pathogen which may aid in development of methods to neutralize LTA pathological potential. Considering factors like LTA extraction method, contamination of LTA used in studies by endotoxins and structural deterioration during in vitro studies, further studies correlating role of LTA in clinical symptoms are to be carried out. The immunostimulatory effect of LTA in causing infection merit further investigation by studies to evaluate precise role of LTA in causing periapical infection, ways for elimination of LTA by developing novel therapeutic agents targeting LTA, and relationship of LTA contents and clinical symptoms for a better treatment outcome.

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**CONFLICT OF INTEREST**

The authors declare that there is no conflict of interest.
AUTHORS' CONTRIBUTION
VSR Conceptualized the study. VSR, MR, AV, and KV did the data curation. BRM visualized the study. MR and VS provided the necessary resources and did the supervision. VSR administered the project. VSAR wrote the original draft. VSR, BRM and AV did the supervision. VSR administered the project. MR and VS provided the necessary resources and AV did the data curation. BRM visualized the study.

VSR Conceptualized the study. VSR, MR, AV, and KV reviewed and edited the final manuscript. KV revised the manuscript. VSR wrote the manuscript. VSAR wrote the original draft. VSR, BRM and AV did the supervision. VSR administered the project. MR and VS provided the necessary resources and AV did the data curation. BRM visualized the study.

VSR Conceptualized the study. VSR, MR, AV, and KV reviewed and edited the final manuscript. KV revised the manuscript. VSR wrote the manuscript. VSAR wrote the original draft. VSR, BRM and AV did the supervision. VSR administered the project. MR and VS provided the necessary resources and AV did the data curation. BRM visualized the study.

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DATA AVAILABILITY
The datasets generated and analysed during the current study are available from the corresponding author on reasonable request.

ETHICS STATEMENT
This article does not contain any studies with human participants or animals performed by any of the authors.

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