RESEARCH ARTICLE

Cell Based Drug Delivery: *Micrococcus luteus* Loaded Neutrophils as Chlorhexidine Delivery Vehicles in a Mouse Model of Liver Abscesses in Cattle

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Abstract

The recent WHO report on antibiotic resistances shows a dramatic increase of microbial resistance against antibiotics. With only a few new antibiotics in the pipeline, a different drug delivery approach is urgently needed. We have obtained evidence demonstrating the effectiveness of a cell based drug delivery system that utilizes the innate immune system as targeting carrier for antibacterial drugs. In this study we show the efficient loading of neutrophil granulocytes with chlorhexidine and the complete killing of *E. coli* as well as *Fusobacterium necrophorum* in *in-vitro* studies. *Fusobacterium necrophorum* causes hepatic abscesses in cattle fed high grain diets. We also show in a mouse model that this delivery system targets infections of *F. necrophorum* in the liver and reduces the bacterial burden by an order of magnitude from approximately $2 \times 10^6$ to $1 \times 10^5$.

Introduction

*Fusobacterium necrophorum* is a Gram-negative, rod shaped and obligate anaerobic bacterium, which is frequently associated with necrotic infections in animals [1]. *F. necrophorum* subspecies *necrophorum* is associated with liver abscess and foot rot in animals [2]. This organism is a major cause of infections including hepatic abscesses and necrotic laryngitis in feedlot cattle, both resulting from aggressive grain-feeding programs [3]. Foot rot and lameness in dairy and beef cattle and hepatic abscesses in feedlot cattle are of major economic concerns to the cattle industry [4], [5]. In feedlots, liver abscesses range from 12–32%, depending on various management and dietary factors [6]. In some feedlots, prevalence can be as high as 90–95% [7].
Cattle with liver abscesses have reduced feed intake, reduced weight gain, decreased feed efficiency, and decreased carcass dressing percentage. Liver abscesses and subsequently liver condemnations are of a major economic concern to beef producers and packers, as the liver accounts for roughly 2% of the weight of the carcass [3]. Economic loss incurred by the beef industry attributable to liver discounts alone has increased from $8.8 million in 1999 to $29.9 million in 2011 (NBQA audit 2011). Antibiotics such as tylosin used as feed additive causes significant reduction in the incidence of liver abscesses but does not prevent liver abscesses [8, 9].

The World Health Organization reports that there is a dramatic increase of microbial resistance against antibiotics [10]. As fewer antibiotics are discovered and made available for use, a different drug delivery approach is urgently needed. There is increased concern among Public Health personnel and consumers over the use of antibiotics for growth promotion in food animals, and several European countries have banned the use of antibiotics as growth promoters (European Commission, 1998). The restriction of antimicrobial use will lead to a much higher incidence of liver abscesses and considerable economic loss to the cattle industry. Therefore, technologies and practices minimizing antimicrobial use are highly appealing.

Recently, liposomal delivery systems have been successfully employed in targeting single bacteria, as well as communities of biofilms [12]. Liposomal delivery systems are of increasing interest, because they are often capable of enhancing the therapeutic index of a drug while minimizing its adverse effects [13]. However, it is evident from the literature that the targeting efficacy of liposomes is limited to roughly 10 percent, because of their clearance from the bloodstream and the natural limitations of the available passive targeting mechanism. [14] Here, we evaluated the use of neutrophils as transport cells with an active mechanism to target the infection site [15].

Neutrophils (neutrophil granulocytes) are the most abundant white blood cell in most species of mammals, are an important part of the innate immune system and are known to be “first responders” to infections and environmental exposure [15]. Neutrophils are “experts” in recognizing and phagocytizing bacteria [16]; hence, if deactivated nonpathogenic bacteria could be loaded with antimicrobial drugs they should be readily taken up by neutrophils. We asked whether bacteria filled with broad-band antibacterial drugs could be transported within defensive cells to a site of infection and whether the therapeutic payload could be effectively released once the defensive cells have reached their target. Our choice of bacterium was Micrococcus luteus, because this bacterium is part of the natural human skin flora and has one of the smallest known genomes. [17]. Chlorhexidine was selected as broad-band antimicrobial because it is a bactericidal and bacteriostatic as well as a topical disinfectant [18]. It is commonly used in mouthwash and contact lens cleaning solutions. Chlorhexidine’s mechanism of action is the attachment to the bacterial cell wall and membrane due to its positive charge [19]. It then disrupts the integrity of the semipermeable membrane, allowing an uncontrolled flow of ions that kills the cell [20, 21]. The uptake time for CHX has been reported as very rapid at about 20 seconds [22]. Although this study is focused on an animal pathogen using a mouse model, the concept established in the study will also be applicable to human diseases.

Materials and Methods

Ethics Statement

This study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The protocol was approved by the Institutional Animal Care and Use Committee at Kansas State University (IACUC approved protocol #3298).
Loading of *M. luteus* with CHX

*M. luteus* (ML) was grown in an overnight culture in LB broth at 37°C and 200 rpm. The cells were separated from the medium and washed with PBS buffer. To load *M. luteus* with chlorhexidine (CHX), the cell pellet of 1 ml cell-suspension was resuspended in concentrated CHX (the stock solution is 2% w/v). The use of PBS buffer or sodium chloride solution to dilute the CHX solution led to unwanted precipitations.

The amount of CHX per 1×10⁸ cells of *M. luteus* was quantified using HPLC. CHX-loaded *M. luteus* cells were lysed with a 1% Triton X Tris lysis buffer (pH 7.2). The cell fragments were removed by centrifugation and the supernatant was analyzed via HPLC (Column: Phenomenex Kinetex 5u XB-C18, Detector: Waters 2998 Photodiode Array Detector). A 20 μl sample of the supernatant was injected for analysis and the UV-Vis spectrum was recorded from 400 to 700 nm. The absorption peak of CHX occurring at 630 nm was used for calibration and data analysis.

Thioglycollate Induced Peritonitis for isolation and *ex-vivo* use of neutrophils

Syngeneic mice were used to isolate circulating leukocytes for loading with appropriate antimicrobials. Mice were injected with 2.5 ml of autoclaved, aged thioglycollate medium (Fisher Scientific) intraperitonially. Twelve hours later, mice were placed under isoflurane anesthesia (2.0%-4.0%) and neutrophils were isolated by peritoneal lavage. The mice were then euthanized. All studies were approved by Institutional Animal Care and Use Committee at Kansas State University.

Flow Cytometry Analysis of Neutrophils

After the extraction of neutrophils from the host mouse, cells were counted and resuspended in RPMI medium at a concentration of 5×10⁵ cells/ml (cut-off for the flow cytometer). The cells were subsequently exposed to dead bacterial samples ranging from 10 to 100 bacterial cells per neutrophil granulocyte. The cell cultures were incubated for two hours at 37°C in a 25ml T-flask. Then the protocol for the PI/annexin 5 apoptosis assay was carried out [23].

Zeta Potential Measurements

Zeta potential measurements were performed using the ZetaPALS Zeta Potential Analyzer (Brookhaven Instruments Corporation), as described previously [24]. *M. luteus* was suspended in 1 X PBS buffer at 1×10⁹ cells/ml. The influence of CHX-hydrochloride on the zeta potential was studied over a range of concentrations from 0 μg/ml to 1000 μg/ml CHX.

In-vitro drug delivery against *E. coli* and *F. necrophorum*

The extracted neutrophils (PMN) were loaded with 20 bacteria per neutrophil (4–6×10⁶ neutrophils per sample). The samples were incubated at 37°C for two hours. Sample types were PMN + ML, PMN + ML modified with CHX, PMN and RPMI buffer.

A starter culture of *E. coli* C600N was grown overnight from a single colony in LB broth containing ampicillin at a final concentration of 50ng/ml and nalidixic acid at a final concentration of 12ng/ml. A 0.1ml volume of the overnight culture was added to 5ml of fresh LB broth containing no antibiotics, and grown to an O.D.₆₀₀ of 0.3. Following centrifugation the bacterial pellet was resuspended in 1X PBS to achieve 1×10⁷ CFU/ml. The bacterial culture was then incubated with 10% v/v complement inactivated mouse serum for 20 minutes at 37°C to opsonize the bacteria. The opsonized bacteria were then mixed with equal amount of PMNs
(1:1 MOI and 1:1 volume of each) and incubated at 37°C for 90 minutes, on a rocker platform. The tubes were then spun down at 5,000 RPM for 5 minutes, the supernatant was discarded and the pellet was resuspended in 100μl of 1X PBS. 100μl of 1% Triton/Tris was added to the resuspended pellet to lyse the eukaryotic cells, and was gently mixed to ensure that no bubbles were formed. This mixture was allowed to sit at room temperature for 5 minutes. 800μl of LB broth was added to the mixture to bring up the final volume to 1ml, which was then serially diluted and plated on LB agar plates containing a final concentration of 50ng/ml ampicillin and 12ng/ml nalidixic acid to enumerate the bacteria.

A starter culture of \textit{F. necrophorum} subsp. \textit{necrophorum} strain 8L1 was grown overnight from a single colony in pre reduced anaerobically sterilized brain heart infusion broth (PRAS-BHI). A 0.3 ml aliquot of the overnight culture was added to 10ml of fresh PRAS-BHI broth and grown to an O.D. \textsubscript{600} of 0.7. The bacteria were spun down to form pellets. Resuspension of bacterial pellet in PBS, opsonization, incubation with neutrophils, lysis of eukaryotic cells from the suspension, serial-dilution were all performed as described above for \textit{E. coli}, but under strict anaerobic conditions. Enumeration was performed by plating on blood agar followed by incubation at 39°C in an anaerobic chamber (Forma Anaerobic System; Thermo Scientific).

\textbf{Preliminary toxicity Study}

In order to determine whether chlorhexidine gluconate was toxic to mice, we conducted \textit{in-vivo} toxicity assays. Briefly, ten week old BALB/c mice were given tail vein intra-venous (IV) injections containing approximately one million neutrophils loaded with an average of 20 cells of \textit{M. luteus} modified with either 1% or 2% chlorhexidine gluconate (CHX). The animals were observed for 5 days post challenge to determine whether they demonstrated any clinical signs of toxicity. At the end of 5 days, the mice were euthanized and their kidneys, liver, brain, lung, spleen and heart were fixed in 10% formalin for histopathological evaluation.

\textbf{Mouse-treatment Study}

\textit{Fusobacterium necrophorum} subsp. \textit{necrophorum} strain 8L1 was grown overnight from a single colony in PRAS-BHI. 0.3ml of the starter culture was added to 10ml of fresh PRAS-BHI broth and grown to an O.D. \textsubscript{600} of 0.7. 1 ml of this culture was diluted 1:40 to achieve a final concentration of approximately 4×10\textsuperscript{6} CFU/ml [25]. 400μl of the diluted bacteria was injected intra-peritonially into 10 week old BALB/c mice. The mice were observed for two days before they were treated with neutrophils carrying antimicrobial cargo or controls.

10 week old BALB/c mice were randomly assigned into 6 groups with 9 mice per group. On day zero, all mice were infected intra-peritonially with an infectious dose of \textit{F. necrophorum}. On day 3 of the experiment, all mice were given intravenous tail vein injections containing 100μl of the following treatments or controls- mice in group 1 were treated with one million neutrophils containing an average of 20 cells of \textit{M. luteus} loaded with chlorhexidine. Mice in group 2 were treated with one million neutrophils containing unmodified, heat deactivated \textit{M. luteus}, mice in group 3 were treated with one million neutrophils and mice in group 4 were treated with 1X PBS. The animals were monitored for clinical signs for 5 days after treatment and were euthanized if any clinical signs developed. The mice were euthanized 5 days post treatment and all mice were examined post mortem for abscesses in the livers. Livers of the mice were weighed and homogenized in a tissue homogenizer for 1 min in modified lactate (ML) broth [26]. The \textit{F. necrophorum} bacterial load in the homogenate was enumerated using most probable number (MPN) analysis [27] using ML broth. Homogenized liver tissue samples were streaked on
blood agar plates for bacterial isolation and identification using Rapid ANAII system (Thermo Scientific). Liver, lung and spleen were fixed in formalin for histopathological evaluation.

**Results**

**Bacterial Loading and CHX retention**

CHX was identified by its absorption peak at 630 nm via HPLC. The concentration of CHX in the bacterial sample was determined by a calibration curve with the range 500–4000 μg CHX/ml in 500 μg increments. The calculated concentration of CHX is 204.6±19.6 μg CHX/10⁸ cells of *M. luteus*.

The uptake of CHX loading also was observed over a 4 hour time frame in one hour increments showing that with an increased loading time more CHX is taken up. The results, however, are not significant (p>0.05) and all samples for *in-vitro* and *in-vivo* experiments have been loaded for 24h to ensure a steady state. Genuit et al., reported that uptake of chlorhexidine by bacteria occurs within 30 seconds [28]. This explains why the uptake of chlorhexidine did not increase significantly between the first and the fourth hour.

Another method to assess CHX loading and, more importantly, where in the bacterial cell CHX is located, is a correlation of zeta potential measurement and UV-Vis analysis of loaded CHX. Since the mechanism of action for CHX is governed by the attraction of the positively charged CHX molecule to the negative surface charge of the bacterial cell an increase of this surface charge can be measured. Fig 1 (left panel) shows the increased zeta potential for a CHX-hydrochloride concentration from 0 μg/ml to 1000 μg/ml loading stock solution in 100μg/ml increments. The data has been normalized to allow a nonlinear regression of the underlying logarithmic shape of this dataset. The zeta potential reaches from -32.86 ± 1.88 mV (0%) at 0μg/ml CHX to -9.63 ± 2.22 mV (100%) at 1mg/ml CHX.

We subsequently correlated the zeta potential data to the actual amount of CHX loaded into the bacterial cell as we detect it by lysing the cells and measuring the CHX concentration in the lysate. The detected linear correlation in Fig 1 (right panel) obtained by the least squares

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**Fig 1.** Zeta potential data (surface charge) for *M. luteus* loaded with CHX-hydrochloride over a range of concentrations from 0 μg/ml to 1000 μg/ml CHX. The data has been normalized (0%-100% equals an increase of the zeta potential from -32.86 ± 1.88 mV to -9.63 ± 2.22 mV) in order to fit a logarithmic function (%ζ = 33.48•log(CHX-concentration)). The panel on the right shows correlation of the measured zeta potential (x-axis) with the measured absorption of CHX in the lysate (y-axis). A linear trend with an R² value of 0.9045 is also shown.

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regression model, suggests that most of the CHX is bound on the surface of the cells or integrated into the cell wall rather than taken up into the cell’s cytoplasm. If CHX would enter the cytoplasm a further increase of CHX concentration in the cell lysate would be expected, while the zeta potential is maintained at its upper limit of roughly -10mV. However this is not the case. A possible explanation is that the negative surface charge has reached a level too weak to further attract the positively charged CHX molecules.

The ability of \textit{M. luteus} to retain CHX it has taken up is vital to ensure that the neutrophil’s quality as a drug carrier is not impaired by cytotoxic effects of released CHX on the way to the site of the infection.

Data (not shown) indicates that after 5 days, 5% of the loaded CHX are released by the cells into the buffer. Given the average lifespan of neutrophils of 3–5 [29] days this presents a satisfying result in terms of protecting the neutrophil carrier cell from cytotoxic CHX effects.

**Neutrophil loading and survival \textit{ex vivo}**

To determine the \textit{ex-vivo} neutrophil loading efficiency for modified bacteria, \textit{M. luteus} was loaded with a fluorescent rhodamine-B dye. The neutrophil sample was analyzed via flow cytometry and compared to an unloaded neutrophil sample. Fig 2 (left panel) shows that approximately 70% of the neutrophil population had taken up the rhodamine-b modified \textit{M. luteus} after a 2h loading period.

\textit{Ex-vivo} neutrophil survival during the loading phase is critical for this drug delivery approach. Neutrophils do not survive for more than a day \textit{ex-vivo} and additional stress by exposure to a bacterial sample can further reduce the survival time dramatically. It has been shown

![Graph showing neutrophil survival](image)

**Fig 2.** Comparison of the fluorescence of an unloaded neutrophil population with a neutrophil population loaded with rhodamine-modified \textit{M. luteus}. Panel on the right shows neutrophil survival after 3 and 6 hours for ratios of 10, 20 and 100 bacteria/neutrophil determined by PI/annexin 5 apoptosis assay and flow cytometry.

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that phagocytic activity increases the oxygen consumption of neutrophils by the factor of two [30]. Fig 2 (right panel) shows that ratios of 10–20 bacteria per neutrophil give reasonable survivability of the neutrophil when compared to the control group. If exposed to a ratio of 100 bacteria per neutrophil the survival drops below 10% for both time points.

**In-vitro Results**

The delivery system was tested in-vitro on *E. coli* and *F. necrophorum* cultures. In the literature, the MIC-value of CHX is 0.25–8 μg/ml [28]. In our assays, the MIC50 value of CHX for *E. coli* C600N was 19.5 μg/ml, and the MIC50 value of CHX for *F. necrophorum* strain 8L1 was 9 μg/ml. These values were exceeded with a CHX concentration of 60.4 ± 4.7 μg/ml in the in-vitro tests. In both cases, the targeted bacterium was killed completely in the treatment group. Control groups containing neutrophils or neutrophils loaded with unmodified *M. luteus* reduced the bacterial count when compared to the no-neutrophil-control group, but failed to completely eliminate the bacteria, leaving viable CFUs in the range of 10^6–10^8. This drop in the CFU count can be attributed to the increased release of bactericidal reactive oxygen species from neutrophils as shown by [31]. Fig 3 shows the results of the in-vitro tests using the neutrophil

![Graph showing CFU/mL vs Treatment](image-url)
based CHX delivery system on *E. coli* C600N. The CHX treatment group was able to kill all *E. coli* C600N so that no colonies were found on LB agar plates. Similarly, Fig 4 shows the complete killing of *F. necrophorum* using the neutrophil based CHX delivery system. Again the control groups containing live neutrophils show a reduced CFU count that can likely be attributed to the bactericidal properties of the phagocyte.

**In-vivo** Results: Toxicity and safety assessment

A preliminary toxicity study showed no toxic effects on the mice, and all of the mice appeared normal and healthy after treatment with neutrophils loaded with 1% or 2% chlorhexidine at any time during the five day observation period after transplant. Moreover, microscopic examination of tissue sections by a pathologist (S.N.) revealed no lesions.

In the final in-vivo mouse experiment the CHX treatment group showed significant results (p<0.05, Pairwise Two-Sided Multiple Comparison Analysis; Dwass, Steel, Critchlow-Fligner Method [32, 33, 34]) against all control groups. Each mouse in the CHX treatment group
received a dose of 0.056 mg CHX carried in the loaded neutrophils. The CFU count was reduced by an order of magnitude from an average of approximately $2 \times 10^6$ in the control groups to $1 \times 10^5$. Fig 5 shows the CFU count results of homogenized and plated liver samples.

A subsequent spectroscopic analysis of the mouse liver samples showed no traces of residual CHX five days after the treatment injection. The damage caused to an untreated liver by *F. necrophorum* can be seen in Fig 6, while livers of the treatment group showed no lesions.

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**Fig 5. Results of the in-vivo experiment.** MP + CHX is the treatment group containing neutrophils loaded with CHX modified *M. luteus*. The PBS group is the control group containing pure RPMI medium while PMN and PMN + MP represent the neutrophil control groups without and with unmodified *M. luteus*. DOI: 10.1371/journal.pone.0128144.g005

PBS: Control group
PMN + MP + CH: CHX Treatment Group
PMN + MP: Neutrophil + unmodified *M. luteus* control
PMNs only: Neutrophil control group
Discussion

Here, we present the following findings: (1) A non-pathogenic bacterium (*Micrococcus luteus*) can be used as a vector for efficiently loading neutrophils with drugs *ex-vivo*, (2) Chlorhexidine (a broad spectrum disinfectant) can be efficiently taken up and trapped by the bacteria, (3) Mouse neutrophils containing the therapeutic cargo can kill *F. necrophorum in-vitro*, and (4) Mouse neutrophils loaded *ex-vivo* with the *M. luteus* containing antimicrobial drugs can be used as vehicles for targeted therapy of liver abscesses in mice caused by *F. necrophorum*.

Thus, the bacterium *M. luteus* is used as a liposome-like protective shell for the active ingredient, chlorhexidine. The additional stability provided by the bacterial cell wall is key to the successful transport of CHX. In our experience liposomes are disrupted by CHX (data not shown). While liposomes rely on their size (passive targeting) and possible targeting ligands on their outer surface to deliver the drug to its target, the cell based approach described here is characterized by a much more specific targeting mechanism as well as an active form of movement. Neutrophils, the targeting component of this drug delivery system, are recruited via chemotaxis into infected tissue [35]. Bacterial antigens naturally enhance the uptake of modified, drug carrying *M. luteus* by neutrophils while the structure of cell wall and membrane represent a natural version of a caged liposome with superior stability against shear- and osmotic stresses. This is a major difference to the bacterial ghost system where inner and outer membranes of Gram-negative bacteria fuse to create an empty bacterial shell. While this has the advantage of a minimal introduction of bacterial DNA into the host organism the bacterial ghost shell releases 40% of the drug it carries within one day. Using intact *M. luteus* as drug carrier has the advantage that approximately 95% of the chlorhexidine is retained within the dead bacterial cell during transport after uptake by the neutrophil. On the other hand, apoptosis of the neutrophil, which naturally occurs after reaching the target, is a dramatic event, which accomplishes the release of a therapeutically effective amount of chlorhexidine. Importantly, the *M. luteus* protects the neutrophil granulocyte, our targeting carrier, from cytotoxic effects of the drug and enhances uptake via phagocytosis. If liposomes were used, they would fuse to the neutrophil and release the free drug into the interior, where it could be cytotoxic.

Fig 6. Image A shows a mouse liver from the control group with lesions (marked by the arrows) caused by *Fusobacterium necrophorum*. Image B shows a liver from the CHX treatment group with no visible lesions.

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Chlorhexidine is a potentially suitable bactericidal drug showing activity even against MRSA strains [36, 37]. Using neutrophils as delivery vehicle has the advantage of a rapid targeted delivery to an infection site due to a fast inflammation response. This allows localized treatment with low drug dosages. The short lifespan of neutrophils [28] guarantees a quick release of the drug.

This cell-delivered antibacterial therapy was efficacious in the mouse model even though only a small amount of chlorhexidine was administered (60 micrograms in 1.3–10^6 neutrophils), and only one dose was administered. Multiple daily injections of the cell therapy may have completely eliminated all bacteria. It is also possible that the cell-based therapy discussed here could be further enhanced by combining two or more agents in the bacteria. For example, silver has been shown to greatly enhance the efficacy of antibiotics, even in cases of drug resistance [38].

Effective antibacterial drugs exist, even for multidrug resistant bacteria, but they are limited by severe toxicity issues. For example, colistin, a polymyxin antibiotic discovered more than fifty years ago but discontinued due to its toxicity is now again in the clinic to treat cystic fibrosis patients, for which there is no other recourse [39]. The approach presented here could allow such drugs to be used more effectively because (1) cells are used as a Trojan horse to shield normal tissues and cells from the drug, and (2) only very small amounts of the drug are required.

Bacterial resistance to antimicrobials is now a global threat to humanity. The findings presented here may offer the prospect of a new armament in the perpetual arms race against multidrug resistant bacteria, since it utilizes a cell-based targeted approach for single or combinatorial delivery of antiseptics or antibiotics that would otherwise be toxic. It is intriguing that a very small amount of drug only administered once can be leveraged into a therapeutic benefit when actively targeted to the site of infection by neutrophils.

Supporting Information

S1 File. HPLC raw data of CHX-loading quantification.
(XLSX)

S2 File. Raw data of anti E. coli and anti F. necrophorum in vitro experiments.
(XLSX)

S3 File. In vivo data of mice MPN liver scores.
(XLSX)

S4 File. Loaded neutrophil survival data.
(XLSX)

S5 File. M. luteus zeta potential and correlation to amount loaded.
(XLSX)

S6 File. CHX retention by M. luteus day 1–5.
(XLSX)

S7 File. Direct observation of CHX in M. luteus.
(XLSX)

S8 File. Quantification of the loaded neutrophil population.
(XLSX)
Author Contributions
Conceived and designed the experiments: DLT SHB SN. Performed the experiments: SOW SM HA LC TBS. Analyzed the data: WH SM SOW. Contributed reagents/materials/analysis tools: DLT SHB SN. Wrote the paper: SOW SM HA.

References
1. Tadepalli S, Stewart G C, Nagaraja T G, Narayanan S K. (2008) Human Fusobacterium necrophorum Strains Have a Leukotoxin Gene and Exhibit Leukotoxic Activity. Journal of Medical Microbiology 57.2: 225–31. Web.
2. Narayanan S K, Stewart G C, Chengappa M M, Willard L, Shuman W, Wilkerson M et al. (2002) Fusobacterium necrophorum Leukotoxin Induces Activation and Apoptosis of Bovine Leukocytes. Infection and Immunity 70.8: 4609–620. Web. PMID: 12117974
3. Nagaraja T G, Chengappa M M. (1998) Liver abscesses in feedlot cattle: a review. J Anim Sci January 76:287–98 PMID: 9464910
4. Checkley S L, Janzen E D, Campbell J R, McKinnon J J. (2005) Efficacy of vaccination against Fusobacterium necrophorum infection for control of liver abscesses and footrot in feedlot cattle in western Canada. Can Vet J. 46(11): 1002–1007 PMID: 16363327
5. Van Donkersgoed J, Jewison G, Mann M, Cherry B, Altwasser B, Lower R et al. (1997) Canadian beef quality audit. Can Vet J.; 38:217–225 PMID: 9105719
6. Brink D R, Lowry S R, Stock R A, Parrott J C. (1990) Severity of liver abscesses and efficiency of feed utilization of feedlot cattle. J. Anim. Sci. 68:1201–1207 PMID: 2385638
7. Nagaraja T G, Lechtenberg K F. (2007) Liver Abscesses in Feedlot Cattle. Veterinary Clinics of North America: Food Animal Practice 23.2: 351–69. Web.
8. Nagaraja T G, Beharka A B, Chengappa M M, Carroll L H, Raun A P, Laudert S B et al. (1999) Bacterial flora of liver abscesses in feedlot cattle fed tylosin or no tylosin. J Anim Sci, 77, 973–978 PMID: 10328365
9. Tan Z L, Lechtenberg K F, Nagaraja T G, Chengappa M M, Brandt R T. (1994) Serum neutralizing antibodies against Fusobacterium necrophorum leukotoxin in cattle with experimentally induced or naturally developed hepatic abscesses. J Anim Sci, 72, 502–508 PMID: 8157357
10. Antimicrobial Resistance Global Report on Surveillance. Genève: World Health Organization, (2014). Print.
11. Bolhuis A, Janice R. (2014) DNA as a Target for Antimicrobials. Bioorganic Chemistry: n. pag. Web.
12. Dhandapani N V, Thapa A, Sandip G, Shrestha A, Shrestha N, Bhattachar R S. (2013) Liposomes as novel drug delivery system: a comprehensive review. Int. J. Res. Pharm. Sci. (Madurai, India), 4, 187
13. Alhariri M, Aghani M, Amiri A. (2013) Liposomal antibiotics for the treatment of infectious diseases. Expert Opin. Drug Delivery, 10, 1515 doi: 10.1517/17425247.2013.822860 PMID: 23886421
14. Maruyama K. (2011) Intracellular targeting delivery of liposomal drugs to solid tumors based on EPR effects. Adv. Drug Delivery Rev, 63, 161 doi: 10.1016/j.addr.2010.09.003 PMID: 20869415
15. Timar C I, Lorincz A M, Ligeti E. (2013) Changing world of neutrophils. Pfluegers Arch., 465, 1521. doi:10.1007/s00424-013-1285-1 PMID: 23636774
16. Hurst J K. (2012) What really happens in the neutrophil phagosome? Free Radical Biol. Med., 53, 508 doi: 10.1016/j.freeradbiomed.2012.05.008 PMID: 22609248
17. Young M, Artsabanov V, Beller H R, Chandra G, Chater K F, Dover L G et al. (2010) Genome Sequence of the Fleming Strain of Micrococcus Luteus, a Simple Free-Living Actinobacterium. Journal of Bacteriology 192.3: 841–60. Web doi:10.1128/JB.01254-09 PMID: 19948807
18. Leikin J B, Paloucek F P. (2008) Chlorhexidine Gluconate. Poisoning and Toxicology Handbook (4th ed.), Informa, pp. 183–184
19. Puig S M, Montiel C J M, Almerich S J M. (2008) Use of chlorhexidine varnishes in preventing and treating periodontal disease: a review of the literature. Med Oral Patol Oral Cir Bucal; 13:E257–60. PMID: 18379452
20. Martin A R. (1959) A technique for studying the action of antisepsics on bacteria in subcutaneous tissues, with special reference to chlorhexidine. Journal of Clinical Pathology, 12, 48–51 PMID: 13631081
21. Jones C G. (1997) Chlorhexidine: is it still the gold standard? Periodontology 2000, 15, 55–62 PMID: 9643233
22. McDonnell G, Denver A R. (1999) Antiseptics and Disinfectants: Activity, Action and Resistance. Clinical Microbiology Reviews 12.1: 147–79. Print. PMID: 9880479
23. Support Protocols. General Annexin V Staining Procedure. N.p., n.d. Web. 29 July 2014.
24. Perera A S, Wang H, Basel MT, Polkhref M R, Gamage P S, Kalita M et al. (2013) Channel Blocking of MspA Revisited. Langmuir 29: 308–315. doi: 10.1021/la3037296 PMID: 23214433
25. Narayanan S K, Chengappa M M, Stewart G C, Nagaraja T G. (2003) Immunogenicity and Protective Effects of Truncated Recombinant Leukotxin Proteins of Fusobacterium necrophorum in Mice. Veterinary Microbiology 93.4: 335–47. Web. PMID: 12713895
26. Tan Z L, Nagaraja T G, Chengappa M M. (1994) Selective enumeration of Fusobacterium necrophorum from the bovine rumen. Appl Environ Microbiol.; 60(4):1387–9. PMID: 8017925
27. Rowe R, Todd R, Waide J. (1977) Microtechnique for most-probable-number analysis. Appl Environ Microbiol.; 33(3): 675–80. PMID: 16345226
28. Genuit T, Bobichchio G, Napolitano L M, McCarter R J, Roghman M C. (2001) Prophylactic Chlorhexidine Oral Rinse Decreases Ventilator-Associated Pneumonia in Surgical ICU Patients. Surgical Infections, 2(1): 5–18. PMID: 12594876
29. Pillay J, Braber I D, Vrisekoop N, Kwast L M, De Boer R J, Borghans JA et al. (2010) In vivo Labeling with 2H2O Reveals a Human Neutrophil Lifespan of 5.4 Days. Blood 116.4: 625–27. Web. doi: 10.1182/blood-2010-01-259028 PMID: 20410504
30. Baldridge C W, Gerard R. W. (1933) The extra respiration of phagocytosis. Am. J. Physiol. 103: 235–236.
31. Reiss M, Roos D. (1978) Differences in Oxygen Metabolism of Phagocytosing Monocytes and Neutrophils. Journal of Clinical Investigation 61.2: 480–88. Web. PMID: 202614
32. Critchlow D E, Fligner M A. (1991) On Distribution-Free Multiple Comparisons in the One-Way Analysis of Variance. Communications in Statistics—Theory and Methods, 20, 127–139.
33. Dwass M, (1960) Contributions to Probability and Statistics, chapter Some k-Sample Rank-Order Tests. 198–202. Stanford, CA: Stanford University Press.
34. Steel R G, (1960) A Rank Sum Test for Comparing All Pairs of Treatments. Technometrics, 2, 197–207.
35. Nuzzi P A, Lokuta M A, Huttenlocher A. (2007) Analysis of Neutrophil Chemotaxis. Methods in Molecular Biology. Volume 370: pp 23–35 PMID: 17416985
36. Brookes S E, Walczak M A, Hameed R, Coonan P. (2002) Chlorhexidine Resistance in Antibiotic-Resistant Bacteria Isolated From the Surfaces of Dispensers of Soap Containing Chlorhexidine. Infection Control and Hospital Epidemiology. 23.11: 692–95. Web.
37. Cookson B. D., Bolton M. C., and Platt J. H.. “Chlorhexidine Resistance in Methicillin-resistant Staphylococcus Aureus or Just an Elevated MIC? An in-vitro and in-vivo Assessment.” Antimicrobial Agents and Chemotherapy 35.10 (1991): 1997–2002. Web.
38. Morones-Ramirez J R, Winkler J A, Spina C S, Collins J J. (2013) Silver Enhances Antibiotic Activity Against Gram-Negative Bacteria. Science Translational Medicine 5.190: 190ra81. Web.
39. Li J, Nation R L, Tumbridge J D, Milne R W, Coulthard K, Rayner C R et al. (2006) Colistin: The Re-emerging Antibiotic for Multidrug-resistant Gram-negative Bacterial Infections. The Lancet Infectious Diseases 6: 589–601. Web. PMID: 16931410