Minocycline modulates NFκB phosphorylation and enhances antimicrobial activity against *Staphylococcus aureus* in mesenchymal stromal/stem cells

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Abstract

**Background:** Mesenchymal stromal/stem cells (MSCs) have demonstrated pro-healing properties due to their anti-inflammatory, angiogenic, and even antibacterial properties. We have shown previously that minocycline enhances the wound healing phenotype of MSCs, and MSCs encapsulated in poly(ethylene glycol) and gelatin-based hydrogels with minocycline have antibacterial properties against *Staphylococcus aureus* (SA). Here, we investigated the signaling pathway that minocycline modulates in MSCs which results in their enhanced wound healing phenotype and determined whether preconditioning MSCs with minocycline has an effect on antimicrobial activity. We further investigated the in-vivo antimicrobial efficacy of MSC and antibiotic-loaded hydrogels in inoculated full-thickness cutaneous wounds.

**Methods:** Modulation of cell signaling pathways in MSCs with minocycline was analyzed via western blot, immunofluorescence, and ELISA. Antimicrobial efficacy of MSCs pretreated with minocycline was determined by direct and transwell coculture with SA. MSC viability after SA coculture was determined via a LIVE/DEAD® stain. Internalization of SA by MSCs pretreated with minocycline was determined via confocal imaging. All protein and cytokine analysis was done via ELISA. The in-vivo antimicrobial efficacy of MSC and antibiotic-loaded hydrogels was determined in Sprague-Dawley rats inoculated with SA. Two-way ANOVA for multiple comparisons was used with Bonferroni test assessment and an unpaired two-tailed Student’s t test was used to determine p values for all assays with multiple or two conditions, respectively.

**Results:** Minocycline leads to the phosphorylation of transcriptional nuclear factor-κB (NFκB), but not c-Jun NH₂-terminal kinase (JNK) or mitogen-activated protein kinase (ERK). Inhibition of NFκB activation prevented the minocycline-induced increase in VEGF secretion. Preconditioning of MSCs with minocycline led to a reduced production of the antimicrobial peptide LL-37, but enhanced antimicrobial activity against SA via an increased production of IL-6 and SA internalization. MSC and antibiotic-loaded hydrogels reduced SA bioburden in inoculated wounds over 3 days and accelerated reepithelialization.

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Conclusions: Minocycline modulates the NFκB pathway in MSCs that leads to an enhanced production of IL-6 and internalization of SA. This mechanism may have contributed to the in-vivo antibacterial efficacy of MSC and antibiotic-loaded hydrogels.

Keywords: Mesenchymal stromal cells, Mesenchymal stem cells, Biomaterials, Minocycline, NFκB, Antimicrobial, Hydrogel, Animal model

Background
Bone marrow-derived mesenchymal stromal/stem cells (MSCs) have demonstrated pro-healing properties in addition to their differentiation potential. Such pro-healing properties include an anti-inflammatory cytokine expression profile, decreased fibrotic granular tissue formation, enhanced keratinocyte proliferation, anti-myofibroblast differentiation, elevated secretion of growth factors and proliferation of microvascular endothelial cells with subsequent angiogenesis, and even antibacterial properties [1–10]. MSCs encapsulated in poly(ethylene glycol) and gelatin-based hydrogels for spatially and temporally controlled cellular presentation to wound sites have demonstrated the promotion of full-thickness cutaneous wound healing [11–14]. Extracellular matrix (ECM)-incorporated scaffolds for MSCs such as these hydrogels have been shown not only to enhance cellular delivery, but also to enhance the bioactivity of biomaterial scaffolds for MSCs through enhanced proliferation and migration [15]. Hydrogels loaded with minocycline alone or in combination with vancomycin and linezolid in addition to MSCs have been shown to prevent the bioburden of *Staphylococcus aureus* (SA) infection [16, 17]. Interestingly, minocycline has been shown to enhance the wound healing phenotype of MSCs in culture and in hydrogels that includes increased invasion capacity, proliferation, ECM attachment, adhesion molecule production, and growth factor production with subsequent angiogenesis [17]. However, the mechanism of MSC wound healing phenotype enhancement has not yet been investigated in detail.

MSC behavior and ultimate expression of wound healing properties is governed by the activation of signaling pathway proteins such as mitogen-activated protein kinase (MAPK/ERK), transcriptional nuclear factor-κB (NFκB), and c-Jun NH2-terminal kinase (JNK/SAPK) [18–20]. Previous studies have demonstrated that the NFκB and MAPK pathways can be activated due to the stimulation of injury or inflammatory factors, such as TNF-α, that leads to some of the pro-healing phenotype enhancements we have observed in MSCs after treatment with minocycline. These observed effects include an increase in growth factor secretion, adhesion molecule expression, and cytokine production [17, 18, 21–23]. Additionally, MSCs have demonstrated an increase in proliferation, migration, and growth factor secretion through the MAPK pathway activation from the antibacterial peptide LL-37 produced by MSCs that has also been shown to contribute to MSC antibacterial potential against *Escherichia coli* and SA [9, 10, 24, 25]. In this study, we investigate the signaling pathway that is modulated by minocycline in MSCs. We then determine whether minocycline has an effect on the MSC production of LL-37 that could have contributed to the MSC enhancement observed previously. We demonstrate that minocycline modulates the phosphorylation of the NFκB pathway in MSCs but inhibits LL-37 production, which led us to investigate the antimicrobial effects of MSCs pretreated with minocycline. We show that MSCs pretreated with minocycline have a significantly enhanced antibacterial capacity against SA due to an increase in IL-6 production and enhancement of SA internalization. Lastly, we demonstrate the in-vivo antibacterial efficacy of minocycline, vancomycin, linezolid, and MSC-loaded hydrogels in SA-inoculated full-thickness cutaneous wounds.

Methods
MSC isolation, characterization, and culture
MSCs were isolated from discarded filters of bone marrow harvests of healthy adult human donors based on a protocol approved by the University of Wisconsin Hospital and Clinics Regulatory Committee per our published protocols [12, 26]. Isolated MSCs at passage 4 were characterized for positive and negative markers via flow cytometry, and for multidifferentiation potential as described previously [12, 26, 27]. MSCs were cultured in 75-cm² tissue culture flasks (TPP, St. Louis, MO, USA) with Dulbecco’s Modified Eagle Medium (DMEM; Cellgro Mediatech, Inc., Corning, NY, USA), 10% fetal bovine serum (FBS), 2 mM L-glutamine, and 2 mM nonessential amino acids (NEAA) with medium changes every 3–5 days. Only MSC passages 4–8 were used in this study.

Minocycline treatment, protein extraction, and western blot
MSCs were cultured in six-well tissue culture plates (CellTreat, Shirley, MA, USA) at 300,000 wells/well in triplicate with 2 ml of MSC culture medium at 0, 50, or 100 μg/ml minocycline (Research Products International, Mt. Prospect, IL, USA) for 48 hours. MSCs were then
harvested and protein was extracted using a NE-PER® Nuclear and Cytoplasmic Extraction Kit with Halt™ protease and phosphatase inhibitor cocktail (Thermo Fischer Scientific). Cytoplasmic protein extract concentrations were determined using a DC™ Protein Assay (BioRad, Hercules, CA, USA) and the presence of specific proteins in the extracts was determined using a western blot analysis of 10 μg of protein as described previously [28]. Primary antibodies used were from Cell Signaling Technologies (Danvers, MA, USA): monoclonal rabbit α-phospho-NF-kappaB (catalog no. 3033), monoclonal rabbit α-NF-kappaB (catalog no. 8242), monoclonal rabbit α-phospho-JNK2 (catalog no. 9258), monoclonal rabbit α-phospho-MAPK (ERK 1/2) (catalog no. 4370), monoclonal rabbit α-MAPK (ERK 1/2) (catalog no. 4695), and monoclonal rabbit actin (catalog no. 4970) at 1:1000. The secondary antibody was used as alkaline phosphatase-conjugated goat α-rabbit IgG (1:30,000; GE Healthcare, Little Chalfont, UK). Membranes were developed for imaging with ECF substrate (GE Healthcare) and imaged on a Storm 840 Scanner (Amersham Bioscience, Amersham, UK) with ImageQuant TL software version 7.0 (GE Healthcare).

Immunofluorescence
MSCs were cultured overnight in six-well tissue culture plates (CellTreat) with glass coverslips added to the wells at 175,000 cells/well with 2 ml of MSC culture medium overnight. MSCs were then cultured with 0, 50, or 100 μg/ml minocycline for 24 hours. After treatment, media were removed and MSCs were washed three times with PBS. MSCs were then fixed for 10 minutes with 4% paraformaldehyde and washed once with PBS. The slides were blocked for 40 minutes using 4% BSA in TBS and then treated with the same primary rabbit α-phospho-NF-kappaB antibody stated earlier at 1:200 in 4% BSA in TBS at 4 °C overnight. The MSCs were washed four times for 10 minutes with TBS and then treated with goat anti-rabbit IgG Alexa Fluor 488 secondary antibody (ThermoFisher Scientific) at 1:200 in 4% BSA for 30 minutes at room temperature. MSCs were then washed twice for 10 minutes with TBS and stained with DAPI for 2 minutes, covered with a coverslip coated with VECTASHIELD® mounting medium, and imaged on an Olympus Fluoview FV1000.

NFκB phosphorylation and protein production analysis
Phosphorylation of NFκB residues was determined using a FACE™ NFκB p65 profiler (Active Motif, Carlsbad, CA, USA). Briefly, MSCs were plated at 50,000 cells/well overnight and then treated with or without minocycline for 24 hours and the protocol was continued using the manufacturer’s instructions. MSC proliferation was determined using the crystal violet stain included in the kit. VEGF, IL-6, TNF-α, and MCP-1 analysis was carried out using a DuoSet® ELISA kit system (R&D Systems, Minneapolis, MN, USA). LL-37 analysis was performed using an ELISA kit (Hyclut Biotech, Olsen, the Netherlands). Complement (C5, C5a, C3, C3b) analysis was performed using respective ELISA kits (Abcam, Cambridge, MA, USA). For VEGF analysis, MSCs were cultured in 24-well tissue culture plates (CellTreat) at 100,000 cells/well in triplicate with 1 ml of MSC culture medium at 0, 50, and 100 μg/ml minocycline with or without 10 μM of the NFκB inhibitor PDTC (BioVision, Milpitas, CA, USA) for 24 hours prior to analysis. For LL-37 production, a treatment condition was used where MSCs were stimulated with IFNγ (100 ng/ml; Sigma Aldrich) for 24 hours [10].

MSC cocultures with *Staphylococcus aureus*
MSCs were cultured overnight in 48-well tissue culture plates (CellTreat) at 50,000 cells/well with 0.5 ml of MSC culture medium. After overnight culture, media were removed and replaced with either fresh MSC media alone, fresh MSC media at 100 μg/ml minocycline, or fresh MSC media at 80 μg/ml vancomycin and cells were cultured for 24 hours. SA (strain #29213; ATCC, Manassas, VA, USA) culture vials stored at −80 °C were thawed, plated on tryptic soy agar (TSA; Becton Dickinson) and incubated on a rotator at 37 °C until the optical density at 600 nm reached 0.1 (1 × 10⁸ CFU/ml) as measured by the spectrophotometer (Nanodrop 2000x UV-vis Spectrophotometer; Thermo Scientific). After MSC treatments, the media were removed and each well was washed three times with 0.5 ml of PBS and each well was replaced with fresh MSC media without minocycline. Immediately following, 20 μl of the SA suspension (200 CFU) was added to the MSC cultures and incubated at 37 °C for 6 hours. Cultures of MSC media alone and TSB alone with 200 CFU SA were included as controls. After 6 hours of culture, 50 μl of the coculture supernatant was removed and diluted serially (1:10) for colony forming ability enumeration as described previously [16]. Immediately following cocultures, supernatants were collected and spun at 12,000 × g for 10 minutes. Supernatants were removed from bacterial cell pellets and placed in a clean Eppendorf tube and stored at −20 °C until cytokine analysis. LIVE/DEAD® stain was applied to the MSCs alone, fresh MSC media at 100 μg/ml vancomycin and cells were cultured for 24 hours. SA (strain #29213; ATCC, Manassas, VA, USA) culture vials stored at −80 °C were thawed, plated on tryptic soy agar (TSA; Becton Dickinson) and incubated on a rotator at 37 °C until the optical density at 600 nm reached 0.1 (1 × 10⁸ CFU/ml) as measured by the spectrophotometer (Nanodrop 2000x UV-vis Spectrophotometer; Thermo Scientific). After MSC treatments, the media were removed and each well was washed three times with 0.5 ml of PBS and each well was replaced with fresh MSC media without minocycline. Immediately following, 20 μl of the SA suspension (200 CFU) was added to the MSC cultures and incubated at 37 °C for 6 hours. Cultures of MSC media alone and TSB alone with 200 CFU SA were included as controls. After 6 hours of culture, 50 μl of the coculture supernatant was removed and diluted serially (1:10) for colony forming ability enumeration as described previously [16]. Immediately following cocultures, supernatants were collected and spun at 12,000 × g for 10 minutes. Supernatants were removed from bacterial cell pellets and placed in a clean Eppendorf tube and stored at −20 °C until cytokine analysis. LIVE/DEAD® stain was applied to the MSCs alone, fresh MSC media at 100 μg/ml vancomycin and cells were cultured for 24 hours. SA (strain #29213; ATCC, Manassas, VA, USA) culture vials stored at −80 °C were thawed, plated on tryptic soy agar (TSA; Becton Dickinson) and incubated on a rotator at 37 °C until the optical density at 600 nm reached 0.1 (1 × 10⁸ CFU/ml) as measured by the spectrophotometer (Nanodrop 2000x UV-vis Spectrophotometer; Thermo Scientific). After MSC treatments, the media were removed and each well was washed three times with 0.5 ml of PBS and each well was replaced with fresh MSC media without minocycline. Immediately following, 20 μl of the SA suspension (200 CFU) was added to the MSC cultures and incubated at 37 °C for 6 hours. Cultures of MSC media alone and TSB alone with 200 CFU SA were included as controls. After 6 hours of culture, 50 μl of the coculture supernatant was removed and diluted serially (1:10) for colony forming ability enumeration as described previously [16]. Immediately following cocultures, supernatants were collected and spun at 12,000 × g for 10 minutes. Supernatants were removed from bacterial cell pellets and placed in a clean Eppendorf tube and stored at −20 °C until cytokine analysis. LIVE/DEAD® stain was applied to the MSCs alone, fresh MSC media at 100 μg/ml vancomycin and cells were cultured for 24 hours. SA (strain #29213; ATCC, Manassas, VA, USA) culture vials stored at −80 °C were thawed, plated on tryptic soy agar (TSA; Becton Dickinson) and incubated on a rotator at 37 °C until the optical density at 600 nm reached 0.1 (1 × 10⁸ CFU/ml) as measured by the spectrophotometer (Nanodrop 2000x UV-vis Spectrophotometer; Thermo Scientific). After MSC treatments, the media were removed and each well was washed three times with 0.5 ml of PBS and each well was replaced with fresh MSC media without minocycline. Immediately following, 20 μl of the SA suspension (200 CFU) was added to the MSC cultures and incubated at 37 °C for 6 hours. Cultures of MSC media alone and TSB alone with 200 CFU SA were included as controls. After 6 hours of culture, 50 μl of the coculture supernatant was removed and diluted serially (1:10) for colony forming ability enumeration as described previously [16].
(PeproTech, Rocky Hill, NJ, USA) during SA cocultures and were analyzed similarly [29]. MSCs were also cultured against SA in a transwell system to prevent cell-to-cell contact. MSCs were plated at 175,000 cells/well in the bottom of a six-well 0.4-μm transwell plate (Corning) with 2 ml of culture medium and cultured overnight. MSCs were then treated with or without minocycline for 24 hours and washed three times with PBS. SA was grown as already stated and 600 CFU (60 μl) of SA was added to 1 ml of culture medium in the top transwell and cocultured for 6 hours. After coculture, SA colony forming analysis was done as already stated and SA attachment to the top transwell surface was carried out using a gentian violet stain protocol as described previously [30].

**Internalization of *Staphylococcus aureus* by MSCs**

MSCs were cultured overnight in six-well tissue culture plates (CellTreat) with glass coverslips added to the wells at 175,000 cells/well with 2 ml of MSC culture medium. MSC treatment with or without minocycline and SA culture preparation was performed as already stated, and then 60 μl of the SA suspension (600 CFU) was added to the MSC cultures and incubated at 37 °C for 4 hours. After coculture, media were removed and the wells were washed three times with PBS. Cells were fixed with 4% paraformaldehyde for 10 minutes, and then washed with PBS. Cells were then stained with Vacnomycin BOPIDY® FL Conjugate (ThermoFisher Scientific) at 1 μg/ml with ActinGreen® 488 ReadyProbes™ (ThermoFisher Scientific) in PBS for 10 minutes at room temperature and washed with PBS. Coverslips were removed from the wells and stained with DAPI for 2 minutes, covered with a coverslip coated with VECTASHIELD® mounting medium, and imaged on an Olympus Fluoview FV1000. Total present SA was characterized accordingly [26, 27]. MSC passages 4–6 were used for this in-vivo study. A day before the experiment, silicone O-rings (McMaster-Carr, Atlanta, GA, USA) were placed under ultraviolet light in a biosafety cabinet for at least 20 minutes and planktonic SA was grown in TSB to OD600 = 0.1 as stated earlier, aliquoted in 1 ml volumes in Eppendorf tubes, and frozen at −80 °C for overnight storage. On the day of the experiment, rat eyes were lubricated with Lubrifresh® (Amazon, Seattle, WA, USA) and treated with buprenophine (0.05 mg/kg). The rat dorsum was shaved and scrubbed with Betasept (4%; Amazon) and sterile saline. Two silicone O-rings were glued to the shaved dorsum of the rats using CrazyGlue Advanced Gel (Elmer’s Products Inc., Westerville, OH, USA) to inhibit healing by contraction, which is the accepted method for examining the healing of full-thickness wounds in rats [32]. Full-thickness wounds were created (two per rat) on the dorsum of the shaved rat using 8-mm biopsy punches (Miltex GmbH, York, PA, USA) to inhibit healing by contraction, which is the accepted method for examining the healing of full-thickness wounds in rats [32]. Full-thickness wounds were created (two per rat) on the dorsum of the shaved rat using 8-mm biopsy punches (Miltex GmbH, York, PA, USA) to inhibit healing by contraction, which is the accepted method for examining the healing of full-thickness wounds in rats [32]. Full-thickness wounds were created (two per rat) on the dorsum of the shaved rat using 8-mm biopsy punches (Miltex GmbH, York, PA, USA) to inhibit healing by contraction, which is the accepted method for examining the healing of full-thickness wounds in rats [32].

**Genetic expression analysis**

MSCs were washed with PBS three times after MSC/SA cocultures in 48-well plates. MSC RNA was isolated by lysing MSCs in culture with 0.5 ml of TRIzol reagent (ThermoFisher Scientific) per well and the protocol was continued as described previously [17]. RT-qPCR was performed on synthesized cDNA using a StepOnePlus Real-Time PCR System (ThermoFisher Scientific) using TaqMan Expression Assay Probes for *IL-6* and *GAPDH*. Data pertaining to mRNAs were collected quantitatively and the CT number was corrected by CT readings of the corresponding internal control of *GAPDH*.

**In-vivo application of hydrogels in *Staphylococcus aureus*-inoculated full-thickness wounds**

All animal experimental protocols were approved by the Institutional Animal Care and Use Committee (IACUC) of the University of Wisconsin-Madison. Female Sprague–Dawley rats (250–300 g, 12 weeks old) were grouped into three different cohorts (n = 3 rats per treatment type for each time point). Sprague–Dawley bone marrow-derived MSCs were similarly isolated, cultured, and characterized accordingly [26, 27]. MSC passages 4–6 were used for this in-vivo study. A day before the experiment, silicone O-rings (McMaster-Carr, Atlanta, GA, USA) were placed under ultraviolet light in a biosafety cabinet for at least 20 minutes and planktonic SA was grown in TSB to OD600 = 0.1 as stated earlier, aliquoted in 1 ml volumes in Eppendorf tubes, and frozen at −80 °C for overnight storage. On the day of the experiment, rat eyes were lubricated with Lubrifresh® (Amazon, Seattle, WA, USA) and treated with buprenorphine (0.05 mg/kg). The rat dorsum was shaved and scrubbed with Betasept (4%; Amazon) and sterile saline. Two silicone O-rings were glued to the shaved dorsum of the rats using CrazyGlue Advanced Gel (Elmer’s Products Inc., Westerville, OH, USA) to inhibit healing by contraction, which is the accepted method for examining the healing of full-thickness wounds in rats [32]. Full-thickness wounds were created (two per rat) on the dorsum of the shaved rat using 8-mm biopsy punches (Miltex GmbH, York, PA, USA) to inhibit healing by contraction, which is the accepted method for examining the healing of full-thickness wounds in rats [32]. Full-thickness wounds were created (two per rat) on the dorsum of the shaved rat using 8-mm biopsy punches (Miltex GmbH, York, PA, USA) to inhibit healing by contraction, which is the accepted method for examining the healing of full-thickness wounds in rats [32]. Full-thickness wounds were created (two per rat) on the dorsum of the shaved rat using 8-mm biopsy punches (Miltex GmbH, York, PA, USA) to inhibit healing by contraction, which is the accepted method for examining the healing of full-thickness wounds in rats [32].

In the in-vivo application of hydrogels in *Staphylococcus aureus*-inoculated full-thickness wounds, rats were randomly assigned to a treatment, with both wounds on the same animal receiving the same treatment. The treated wounds were allowed to dry for 2 minutes, then covered with Tegaderm™ film (3 M Healthcare, Neuss, Germany) and wrapped with cling gauze followed by Vetrap™ (3 M) tape. At 1 and 3 days after treatment application, rats were euthanized with Beuthanasia (0.2 ml/kg; Schering-Plough Animal Health Corp., Union, NJ, USA). The wounds were then imaged, before one wound bed per
rat was harvested and put into 1 ml of sterile PBS with sterile beads for SA CFU enumeration, and one wound bed per rat was harvested, fixed in 10% neutral buffered formalin for 48 hours, paraffin embedded, and H&E stained [14, 33]. Wound beds collected for SA CFU enumeration were homogenized using a bullet blender (three 5-minute cycles; Next Advanced, Averill Park, NY, USA) and then diluted serially and cultured on agar plates for colony counting as already stated. Photomicrographs were taken of histological sections from H&E staining and the epidermal thickness of the healing wound tissue with clearly demarcated epidermal and dermal tissue boundaries was measured by averaging five epithelialized keratinocyte widths from randomized locations including the central region of the wound and areas adjacent to the wound margin.

Statistical methods
All assays were conducted with at least three replicates per condition. Two-way ANOVA for multiple comparisons was used with a Bonferroni test assessment to determine \( p \) values for all assays with three or more conditions, and an unpaired two-tailed Student’s \( t \) test was used to determine \( p \) values for all assays with only two conditions. \( p < 0.05 \) was considered statistically significant. All analyses were performed with GraphPad Prism 5.0. Error bars represent the standard deviation of the specific group mean.

Results
Modulation of NFkB by minocycline in MSCs
MSCs cultured with 100 \( \mu \)g/ml minocycline for 48 hours led to the presence of a phosphorylated NFkB western blot band from the protein extract that was not present in protein extracts derived from MSCs treated with 0 or 50 \( \mu \)g/ml minocycline for 48 hours. There were no differences in phosphorylation of JNK or ERK between all of the MSC culture conditions (Fig. 1a). Immunofluorescence shows that NFkB is phosphorylated by minocycline in a dose-dependent manner (Fig. 2a). NFkB is phosphorylated at the serine 468 residue when MSCs are treated with 50 \( \mu \)g/ml minocycline (\( p < 0.001 \) vs 0 \( \mu \)g/ml) and with 100 \( \mu \)g/ml minocycline (\( p < 0.0001 \) vs 0 \( \mu \)g/ml). NFkB is phosphorylated at the 536 residue when MSCs are treated with 50 \( \mu \)g/ml minocycline (\( p < 0.05 \) vs 0 \( \mu \)g/ml) and with 100 \( \mu \)g/ml minocycline (\( p < 0.0001 \) vs 0 \( \mu \)g/ml). Total levels of NFkB remained unchanged between treatment conditions (Fig. 1c). There was an increase in MSC proliferation after 24 hours of culture when MSCs were treated with 50 \( \mu \)g/ml minocycline (\( p < 0.0001 \) vs 0 \( \mu \)g/ml) and with 100 \( \mu \)g/ml minocycline (\( p < 0.0001 \) vs 0 \( \mu \)g/ml, \( p < 0.0001 \) vs 50 \( \mu \)g/ml) (Additional file 1: Figure S1).

Effects of minocycline on VEGF and antimicrobial peptide LL-37 production
There was an increase in VEGF production by MSCs when treated with 50 \( \mu \)g/ml minocycline (\( p < 0.05 \) vs 0 \( \mu \)g/ml) and with 100 \( \mu \)g/ml minocycline (\( p < 0.05 \) vs 0 \( \mu \)g/ml). The increased production of VEGF by minocycline was not present when MSCs were cultured with the NFkB inhibitor PDTC (control, \( p < 0.01 \); 50 \( \mu \)g/ml, \( p < 0.01 \); 100 \( \mu \)g/ml, \( p < 0.05 \)) (Fig. 2). There was a decrease in LL-37 production in MSCs treated with 100 \( \mu \)g/ml minocycline for 24 hours (\( p < 0.001 \) vs MSC control), and there was an increase in LL-37 production in MSCs treated with 100 ng/ml IFN-\( \gamma \) for 24 hours (\( p < 0.05 \) vs MSC control) (Fig. 3).

Minocycline effects on MSC antibacterial activity
MSCs pretreated with 100 \( \mu \)g/ml minocycline for 24 hours demonstrated complete inhibition of SA colony forming abilities (\( p < 0.0001 \) vs media control), while MSCs not pretreated with minocycline or MSCs pretreated with 80 \( \mu \)g/ml vancomycin demonstrated a decrease in SA colony forming abilities (\( p < 0.01 \) vs MSC control). MSCs pretreated with minocycline demonstrated a significantly lower colony forming ability inhibition of SA than MSCs not pretreated with minocycline (\( p < 0.001 \)) and MSCs pretreated with vancomycin (\( p < 0.0001 \)) (Fig. 4a). MSCs not pretreated with minocycline demonstrated an increase in LL-37 production when cocultured with SA (\( p < 0.05 \) vs MSCs), while MSCs pretreated with 100 \( \mu \)g/ml minocycline demonstrated an inhibition of LL-37 production with and without SA coculture (\( p < 0.05 \) vs MSCs). MSCs not pretreated with minocycline and cocultured with SA produced higher levels of LL-37 compared to MSCs pretreated with minocycline without SA coculture (\( p < 0.01 \)) and compared to MSCs pretreated with minocycline with SA coculture (\( p < 0.01 \)) (Fig. 4b). Both MSCs not pretreated with minocycline and MSCs pretreated with minocycline demonstrated an increase in IL-6 production when cocultured with SA (\( p < 0.0001 \) vs MSCs), with minocycline-preconditioned MSCs demonstrating a higher production of IL-6 compared to MSCs not preconditioned with minocycline when both were cocultured with SA (\( p < 0.01 \)) (Fig. 4c). MSCs not pretreated with minocycline and cocultured with SA demonstrated an increase in IL-6 gene expression (\( p < 0.05 \) vs MSCs) and MSCs pretreated with minocycline and cocultured with SA demonstrated an increase in IL-6 gene expression (\( p < 0.0001 \) vs MSCs), with minocycline-preconditioned MSCs demonstrating higher levels of IL-6 gene expression compared to MSCs not preconditioned with minocycline when both were cocultured with SA (\( p < 0.001 \)) (Additional file 2: Figure S2). SA colony forming abilities were higher when cocultured with MSCs pretreated with minocycline with anti-IL-6 present in cocultures compared to when cocultured with MSCs
pretreated with minocycline without anti-IL-6 present in cocultures ($p < 0.01$) (Additional file 3: Figure S3). There were no differences in TNF-α or MCP-1 production levels between all MSC culture conditions (Fig. 4d, e).

There was a higher level of viable MSCs after coculture with SA when pretreated with minocycline ($p < 0.01$ vs untreated) and there was a lower level of necrotic MSCs after coculture with SA when pretreated with minocycline ($p < 0.0001$ vs untreated) (Fig. 5a). Live/dead images with only cell borders visible suggest a higher presence of necrotic MSCs in the untreated condition (Fig. 5b). Attachment of SA to the transwell surface in transwell cocultures was decreased when cocultured with MSCs pretreated with minocycline ($p < 0.0001$ vs media control) and SA colony forming abilities were lowered, but not absent, in transwell cocultures when cocultured with MSCs pretreated with 100 μg/ml minocycline ($p < 0.05$ vs media control) (Fig. 6a, b).

MSCs demonstrated a higher level of SA internalization when pretreated with 100 μg/ml minocycline ($p < 0.0001$ vs MSCs) (Fig. 7a, b). SA colony forming abilities were higher in cocultures with MSCs pretreated with minocycline when internalization inhibitor chloroquine ($p < 0.01$ vs MSCs pretreated with minocycline) or...
3-methyladenine ($p < 0.001$ vs MSCs pretreated with minocycline) was included in the minocycline pretreatment. SA colony forming abilities were still lower in cocultures with MSCs pretreated with minocycline and chloroquine ($p < 0.01$ vs media control) and in cocultures with MSCs pretreated with minocycline and 3-methyladenine ($p < 0.01$ vs media control) (Fig. 7c). Production of complement protein C5 was higher in MSCs cocultured with SA when not pretreated with minocycline ($p < 0.0001$ vs MSCs, $p < 0.0001$ vs MSCs pretreated with minocycline + SA). Production of the cleaved complement protein C5a was higher in MSCs cocultured with SA when not pretreated with minocycline ($p < 0.0001$ vs MSCs, $p < 0.0001$ vs MSCs pretreated with minocycline + SA). Production of the complement protein C3 was higher in MSCs cocultured with SA when not pretreated with minocycline ($p < 0.05$ vs MSCs, $p < 0.0001$ vs MSCs pretreated with minocycline + SA). Production of the cleaved complement protein C3b was higher in MSCs cocultured with SA when not pretreated with minocycline ($p < 0.0001$ vs MSCs, $p < 0.0001$ vs MSCs pretreated with minocycline + SA) (Fig. 7d–g).

**Discussion**

Small molecules are known to influence the immunomodulatory and antibacterial properties of MSCs. Adipose tissue-derived MSCs can be treated with rapamycin which leads to the production of anti-inflammatory molecules that could help prevent acute graft-versus-host disease (aGVHD) [34]. Menstrual-derived MSCs have shown a synergistic antimicrobial effect with antibiotics in improving survival rates and enhancing bacterial clearance with reduced organ injury in an animal sepsis model [35]. Minocycline has an immunomodulatory effect on bone marrow-derived MSCs that leads to a synergistic effect with minocycline itself in treating autoimmune encephalomyelitis [36]. Bone marrow-derived MSCs have therapeutic benefits in other multiple applications where the effects of antibiotic treatment are unknown, and it would be beneficial to investigate the effects of minocycline treatment on these applications. For example, MSCs increase the thickness of the ventricular wall in a heart failure model, and they improve cardiac function with enhanced ventricular remodeling and reduced cardiac fibrosis in a heart transplantation model [37, 38]. This effect could be influenced by a minocycline-induced production of VEGF [17]. MSCs can also influence macrophage polarization, and MSCs have been shown to increase the frequency of alternatively activated macrophages in the heart after acute myocardial infarction [39]. The effect of minocycline-treated MSCs on macrophage polarization would be a logical future study to investigate further.
immunomodulatory effects of minocycline-treated MSCs [40, 41].

We have shown previously that minocycline can enhance the wound healing properties in human bone marrow MSCs, but the mechanism of enhancement was still unknown [17]. Here, we demonstrate that this enhancement of the wound healing phenotype may be due to the phosphorylation and potential activation of NFκB in MSCs. NFκB is a transcription factor that affects gene expression changes in responses to stimuli including cytokines and bacterial antigens that can ultimately regulate immune responses to infection [42, 43]. NFκB controls the expression of inducible chemokines, cell adhesion molecules, and vasoactive and anti-apoptotic proteins [44–46]. Activation of the NFκB pathway in MSCs due to stimulation by TNF-α, LPS, or hypoxia has led to the increased production of multiple growth factors by MSCs including VEGF, FGF2, IGF-1, and HGF that could play a crucial role in the wound healing attributes of MSCs [21]. Activation of the NFκB pathway though the TNF receptor family can lead to the cytoplasmic phosphorylation of NFκB at serine 536, as
observed in this study, that leads to IκB kinase ε (IKKe) translocation to the nucleus, where it can associate with the promoter regions of several NFκB target genes and phosphorylate other transcription factors, ultimately leading to the transcription of a subset of NFκB target genes [47]. Activation of the NFκB pathway in MSCs due to stimulation by TNF-α has also shown to enhance IL-6, vascular cell adhesion molecule-1 expression (VCAM-1), and intercellular adhesion molecule 1 (ICAM-1) production [18, 47, 48]. These studies suggest that the NFκB pathway may be activated through the interaction of minocycline with the TNF receptor family which leads to the recruitment of an IκB kinase (IKK) that phosphorylates inhibitor of kappa B (IκB) on NFκB and ultimately leads to IκB polyubiquitination and degradation by the proteasome which allows for cytoplasmic p65 phosphorylation and downstream effects [49]. This mechanism of NFκB cytoplasmic phosphorylation and IKKe nuclear translocation may be responsible for the minocycline-induced enhancement of VEGF and IL-6 production, as well as enhanced adhesion molecule production in MSCs as we observed previously. Indeed, this is suggested by inhibiting the NFκB pathway in combination with minocycline treatment that ultimately prevented the minocycline-induced increase in MSC VEGF production [17].

In addition to NFκB phosphorylation, treatment of MSCs with the antibacterial peptide LL-37 increases MSC migration, proliferation, growth factor expression, and tissue regeneration capacity similar to what we
observed in our previous study [17, 24]. Interestingly, LL-37 activates the NFκB pathway in embryonic endothelial progenitor cells that enhances recruitment to muscle tissue and in periodontal ligament cells that led to enhanced VEGF production [50, 51]. MSCs produce LL-37 in response to infection that leads to antibacterial activity against *E. coli*, *Pseudomonas aeruginosa*, and SA [9]. LL-37 produced by MSCs has been shown to be synergistic with the aminoglycoside antibiotic genetin in a cystic fibrosis infection treatment model and decreased the severity of lung injury in *E. coli*-infected lungs [10, 52]. However, our results show that minocycline actually inhibits the production of LL-37 in MSCs both alone and when stimulated by SA, suggesting that the NFκB pathway is activated by minocycline itself and not from an induction of LL-37 production. Because LL-37 has been proven to be a vital peptide in MSC antibacterial activity, we determined whether minocycline treatment had an effect on MSC antibacterial activity against SA.

Interestingly, pretreatment of MSCs with minocycline leads to greatly enhanced antibacterial activity against SA in which no colony formation was observed after coculture. MSC IL-6 production is increased in MSCs cocultured with SA, but both production and gene expression of IL-6 was higher in MSCs after SA coculture when pretreated with minocycline. NFκB activation through lipopolysaccharide (LPS) stimulation has been shown to control IL-6 production in macrophages, and an increase in IL-6 through NFκB activation was not accompanied by an increase in TNF-α [53] as we also observed in this study. IL-6 stimulates antibacterial responses of immune regulating cells and has been shown to initiate antibacterial activity of keratinocyte grafts against SA through an increase in the synthesis of cytokines that could be cytotoxic to SA [54]. Although we demonstrate that IL-6 is a major factor in the minocycline-enhanced antibacterial activity of MSCs, it is not the only factor, because reduction of IL-6 with an antibody does not lead to recovery of the full colony forming ability of SA. Previous studies suggest that phagocytosis of SA may be a part of the MSC antibacterial profile [55, 56].

Separating physical contact of MSCs and SA through a transwell system led to the formation of SA colonies when cocultured with MSCs pretreated with minocycline, suggesting that cell-to-cell contact is necessary for optimal antibacterial activity of MSCs against SA. There are multiple ways in which MSCs can demonstrate antibacterial activity through cell-to-cell contact. MSC immunoregulation includes activation of the complement system proteins, which is generally suppressed by MSC production of factor H but can be induced through inflammatory factors that may be secreted by SA [57]. The activation of the NFκB pathway is correlated with the cleavage of complement protein C5 to C5a and C5b in MSCs that leads to a cascade which forms a Membrane Attack Complex (MAC) that induces cell death in microbes [58, 59]. However, in this study we see that
minocycline inhibits both production of C5 and the cleavage of C5 into C5a. The complement protein C3 is produced and cleaved into the active C3b protein by MSCs as well, which leads to opsonization that ultimately leads to bacterial phagocytosis [60, 61]. Again, minocycline inhibits C3 production and cleavage into C3b in MSCs, suggesting the cell-to-cell contact antibacterial activity is accomplished by different means. Because the complement system seems to be suppressed in MSCs by pretreatment with minocycline, it is possible that minocycline induces the production of factor H even in the presence of inflammatory factors produced by SA [57]. Bacterial internalization and an increase in IL-6 production are strongly correlated [55, 62]. The increased internalization of SA by MSCs may have been due to NFκB activation of minocycline, which increased IL-6 production in the presence of SA that ultimately induced the increased SA internalization, or the potential preservation of MSC viability by minocycline pretreatment may have allowed MSCs to carry on normal internalization activity.

Vancomycin and linezolid have different mechanisms of antibacterial activity against SA than minocycline, and
are not known to modulate NFκB in mammalian cells [63–65]. Additionally, minocycline has been shown to modulate neuroinflammation independently of its antimicrobial activity in brain abscesses caused by SA infection, suggesting minocycline’s antimicrobial and immunomodulatory enhancement effects on other cell types [66]. The specific minocycline-induced antimicrobial enhancement of MSCs may have contributed to the in-vivo antimicrobial efficacy of MSC and triple antibiotic-loaded hydrogels that reduced SA bioburden and increased reepithelialization in SA-inoculated wounds.

**Conclusions**

Minocycline modulates the NFκB pathway and enhances antibacterial activity against SA via an increase in IL-6 production and SA internalization by MSCs. MSC-loaded hydrogels containing minocycline reduce bacterial bioburden in SA-inoculated full-thickness cutaneous wounds.

**Additional files**

- **Additional file 1: Figure S1.** Showing proliferation of MSCs after 24 hours of minocycline treatment in a dose-dependent manner. (JPG 103 kb)
- **Additional file 2: Figure S2.** Showing expression of IL6 in MSCs when cultured alone or with SA with and without minocycline pretreatment. (JPG 171 kb)
- **Additional file 3: Figure S3.** Showing colony forming abilities of SA after 6 hours of coculture with media controls, MSCs with or without anti-IL-6, and MSCs pretreated with minocycline with or without antiIL-6. (JPG 146 kb)

**Abbreviations**

aGVHD: Acute graft-versus-host disease; ANOVA: Analysis of variance; BSA: Bovine serum albumin; CDNA: Complementary deoxyribonucleic acid; CFU: Colony forming unit; CT: Cycle threshold; ECM: Extracellular matrix; ELISA: Enzyme-linked immunosorbent assay; FG2: Fibroblast growth factor 2; GAPDH: Glyceraldehyde 3-phosphate dehydrogenase; H&E: Hematoxylin and eosin; HGF: Hepatocyte growth factor; ICAM-1: Intercellular adhesion molecule 1; IFN: Interferon gamma; IGF-1: Insulin-like growth factor 1; IgG: Immunoglobulin G; IKK: Ib kinase; IL-16: Interleukin-6; IkB: Inhibitor of kappa B; JNK/SAPK: c-Jun N-terminal kinase; LL-37: Cathelicidin-related antimicrobial peptide; LPS: Lipopolysaccharide; MAPK/ERK: Mitogen-activated protein kinase; MCP-1: Monocyte
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Availability of data and materials
The datasets used and analyzed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions
ADG conceived and designed the study, conducted the experiments, collected and assembled the data, analyzed and interpreted the data, wrote the main manuscript text, and prepared Figs. 1, 2, 3, 4, 5, 6, 7 and 8. PH and WJK provided laboratory facilities, administrative support, and provision of study materials and donors. ADG, PH, and WJK provided financial support. All authors reviewed and edited the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate
MSCs were isolated from discarded filters of bone marrow harvests of healthy adult human donors based on a protocol approved by the University of Wisconsin Hospital and Clinics Regulatory Committee per our published protocols (Protocol Number 2016-0298). Informed consent was not required for this study. All animal experimental protocols were approved by the Institutional Animal Care and Use Committee (IACUC) of the University of Wisconsin-Madison (Protocol Number V0 1060).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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