Gallic Acid Ameliorates Aspergillus Fumigatus Keratitis Through Reducing Fungal Load and Suppressing the Inflammatory Response

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PURPOSE. The purpose of this study was to explore the antifungal and anti-inflammatory effects of gallic acid (GA) on Aspergillus fumigatus (A. fumigatus) keratitis.

METHODS. CGK-8 assay and Draize eye test were used to determine the non-cytotoxic concentration of GA in RAW264.7 cells and an A. fumigatus keratitis mouse model. The antifungal effects of GA were analyzed using minimal inhibitory concentration (MIC), biofilm formation test, fungal adherence assay, calcofluor white staining, and propidium iodide staining. The therapeutic effects of GA were estimated by slit lamp photographs, clinical score, hematoxylin and eosin (H&E) staining, and Periodic acid-Schiff staining in vivo. Immunofluorescence staining and myeloperoxidase assay were conducted to identify neutrophil infiltration and activity. RT-PCR, ELISA, and Western blot were performed to detect the expression of pro-inflammatory cytokines and Nrf2/HO-1.

RESULTS. In HCECs and A. fumigatus keratitis mouse model, GA at 100 μg/mL did not affect cell viability, thus this concentration was applied to subsequent experiments. In vitro, GA significantly inhibited A. fumigatus growth, biofilm formation, and adhesion. In vivo, 100 μg/mL GA alleviated the severity of fungal keratitis (FK) by repressing fungal load, reducing neutrophil infiltration, and lowering MPO activity. Besides, the expression of IL-1β, TNF-α, LOX-1, and COX-2 was inhibited, whereas Nrf2 and HO-1 expression was enhanced at both mRNA and protein levels in the 100 μg/mL GA treated group in comparison to PBS control.

CONCLUSIONS. GA ameliorates FK severity through inhibiting A. fumigatus load, reducing neutrophils infiltration, downregulating the expression of pro-inflammatory cytokines, and enhancing the Nrf2/HO-1 pathway, which provides new insight into A. fumigatus keratitis treatment.

Keywords: gallic acid (GA), fungal keratitis (FK), antifungal, anti-inflammatory, Nrf2/HO-1

Fungal keratitis (FK) is an intractable ocular infection that can result in a corneal ulcer, perforation, and even blindness, which attribute to not only the presence of fungal hyphae, but also severe inflammatory response. The incidence of FK ranges from 6% to 56% in various regions of the world, suggesting the need to explore effective antifungal and anti-inflammatory agents.

FK is commonly caused by A. fumigatus, Aspergillus flavus (A. flavus), and Fusarium. Over the past 10 years, A. fumigatus has become the most prevalent airborne fungal pathogen, causing severe and usually fatal invasive infections in immunocompromised hosts in developed countries. Due to agricultural ocular trauma, postoperative corneal infection, long-term use of contact antibiotics and lenses, and excessive use of corticosteroids, the incidence of FK has been increasing, especially in developing countries and tropical regions. However, current antifungal drugs have significant limitations. For example, in a toxicity evaluation by Foster et al., amphotericin B was found to severely retard the rate of debrided corneal closure and manifested dramatic pathologic changes (evaluated on the bases of scores for quality of regenerating epithelium, stromal edema and haze, and iritis) that worsened each day with the continuation of therapy. Hence, from the abovementioned discussion, local toxicities present an important aspect of the challenge in the ocular application of amphotericin B. The other commonly used antifungal drug is natamycin. But natamycin has poor permeability, which barely passes an intact corneal epithelium due to its molecular mass (666 Da). In support, animal studies showed that the efficacy of natamycin was substantially lessened in intact versus debrided corneas. More importantly, amphotericin B and natamycin are costly, and repeated application in a relatively short period is required to prevent FK re-occurrence and drug resistance.
therapeutic strategy may reduce follow-up visits, especially those who cannot afford medical care. Therefore, it is urgent to find effective and affordable treatment for FK.

Gallic acid (GA; 3,4,5-trihydroxy benzoic acid) is a natural polyphenol ubiquitously present in fruits, vegetables, legumes, and herbal medicines.8 Previous studies have reported that GA and its derivatives possess a variety of biological properties, such as antioxidative, anti-viral, anticancer, and anti-inflammatory activities.9 For example, GA can bind and quench reactive oxygen species (ROS) with high affinity, thereby ameliorating oxidative stress-associated tissue damage in many diseases, including cancers, liver diseases, neuroinflammation, and inflammatory bowel disease.10–13 GA exhibits broad-spectrum antimicrobial activities, including antifungal, antibacterial, and antiviral in vitro.14 GA significantly inhibited Trichophyton rubrum growth by repressing the expression of two key enzymes involved in fungal ergosterol biosynthesis, squalene epoxidase, and sterol 14α-demethylase P450 (CYP51), to interfere with fungal membrane integrity,15 and the fungicidal efficiency was comparable to fluconazole.14 Recently, GA extracted from Cassia fistula has been shown to have a potent antifungal activity on fluconazole-resistant Candida species isolated from patients with HIV,16 whereas the antifungal mechanisms remain unclear.

On the other hand, corneal inflammation and its associated complications are the leading cause of vision impairment in FK. The excessive inflammatory response induced aggregation of immune cells, such as neutrophils, macrophages, and dendritic cells, further promotes the production of pro-inflammatory cytokines, such as IL-1β, TNF-α, and COX-2, resulting in corneal damage, nubecula, and even corneal perforation.17 Thus, controlling the excessive immune response in FK is beneficial in preventing inflammation-related corneal injury. In addition to its antifungal property, GA plays an anti-inflammatory role in different animal models and human subjects. For instance, GA attenuated lipopolysaccharides (LPS)-induced neuroinflammation and neurotoxicity in the rat brain and BV-2 microglial cells and significantly reduced IL-1β expression.18 GA also acts as the gastroprotective property in ethanol-induced gastric ulcer rats via downregulating the serum levels of pro-inflammatory cytokines, including IL-1β and tumor necrosis factor α (TNF-α) and activating nuclear factor-erythroid 2-related factor 2 (Nrf2)/heme oxygenase (HO-1) pathway.19 However, to date, there is no study investigating the role of GA in FK.

In this study, we demonstrate the antifungal and anti-inflammatory effects of GA in mouse macrophages and an A. fumigatus keratitis mouse model. GA treatment could alleviate the severity of A. fumigatus keratitis in mice not only by reducing fungal load but also by repressing the expression of pro-inflammatory cytokines and enhancing the Nrf2/HO-1 pathway. Our findings identified the antifungal and anti-inflammatory related mechanisms of GA in A. fumigatus keratitis, providing a new alternative for FK treatment.

**Materials and Methods**

**Gallic Acid and Natamycin Preparation**

GA powder (CAS 149-91-7; Macklin Biochemical Co. Ltd., Shanghai, China) was dissolved in phosphate-buffered saline (PBS; Solarbio, Beijing, China) at a storage concentration of 10 mg/mL, and further diluted to the required concentration with Dulbecco’s Modified Eagle Medium (DMEM; Solarbio) for RAW264.7 cells and human corneal epithelial cells (HCECs), and Sabouraud liquid medium which contains protein, agar, and glucose for fungal conidia treatment.

Natamycin (NAT) powder (CAS 7681-93-8; Macklin Biochemical Co. Ltd., Shanghai, China) was dissolved in PBS (Solarbio) at a storage concentration of 8 mg/mL, and further diluted to the required concentrations with DMEM (Solarbio) for RAW264.7 cells, and Sabouraud liquid medium for fungal conidia treatment.

**Aspergillus Fumigatus Culture**

*A. fumigatus* standard strain 3.0772 was obtained from China General Microbiological Culture Collection Center (Beijing, China), and stored in the refrigerator at −80°C. Its aerobic and culture temperature was 25 to 28°C. The second generation of *A. fumigatus* was inoculated in 150 mL Erlenmeyer flasks containing Sabouraud liquid medium. After 48 hours of shaking at 120 rpm, 37°C, the hyphae were washed using sterile PBS 3 times and collected by centrifugation at 5000 rpm for 15 minutes. The supernatant was discarded, and the hyphae were resuspended by 5 mL DMEM.

Foractivehyphae preparation, we pipetted 10 μL of diluted fungi into a hemocytometer and counted the number of hyphae under the microscope. According to the counting results, the hyphae solution was then diluted using DMEM to a final concentration of approximately 3 × 10⁸ hyphal fragments/mL. The active *A. fumigatus* hyphae were also inoculated onto Sabouraud agar plates and incubated for 36 to 48 hours at 37°C. Spores were scraped, diluted with PBS, and counted using the same approach as described above.

They were adjusted to 3 × 10⁹/mL for minimum inhibitory concentration (MIC), biofilm formation test, fungal adherence assay, calcofluor white staining, and Propidium iodide (PI) staining.

The inactive hyphae were prepared using active hyphae by overnight incubation with 75% alcohol. After being washed by DMEM, the inactive hyphae were diluted and counted using the same approach as described above. The final concentration was also adjusted to 10⁹ hyphal fragments/mL.

**Cell Culture and A. fumigatus Stimulation**

RAW264.7 cells (Chinese Academy of Sciences, Shanghai, China) were cultured as described previously.20 Briefly, RAW264.7 cells were seeded in 6- and 12-well plates for 24 hours until 80% to 90% confluency, then stimulated with inactive hyphae at 3 × 10⁶ hyphal fragments/mL for 1 hour, followed by GA incubation at a final concentration of 100 μg/mL or NAT at 8 μg/mL. RAW264.7 cells were collected at 8 hours for RT-PCR to detect the mRNA levels of IL-1β, TNF-α, LOX-1, and COX-2, and collected at 24 hours for Western blot to examine the protein levels of HO-1 and Nrf2. The supernatant at 24 hours was used for ELISA to estimate the protein levels of IL-1β and TNF-α.

**Minimum Inhibitory Concentration**

The MIC assay method refers to the standardized microdilution method.21 GA was diluted using Sabouraud liquid medium by a 2-fold dilution method to 12.5, 25, 50, 100, 200, 400, and 800 μg/mL in a sterile Eppendorf tube, and *A. fumigatus* spores at 3 × 10⁶/mL were added to each
tube. After being mixed adequately, each solution was transferred into a 96-well plate (100 μL/well) with 6 repetitions for each concentration. After 48 hours of incubation at 37°C, the optical density (OD) was measured at 540 nm with a microplate reader. Two control conditions (GA-free medium with or without conidium) were simultaneously set up for each plate.

**Cell Counting Kit-8 Assay**

Human corneal epithelial cells (HCECs; PCS-700-010; kindly provided by the Ocular Surface Laboratory Zhongshan Ophthalmic Center, China) were cultured using the previous method. HCECs (3 × 10^4/mL) were suspended and transferred into 96-well plates with sterile DMEM and incubated for 24 hours until 90% confluence. After being rinsed with sterile PBS, each well was incubated with 100 μL different concentrations of GA diluted with DMEM for 48 hours, followed by a 2-hour incubation with 10 μL CCK-8 (MedChemExpress) at 37°C. Then, OD at 450 nm was measured by a microplate reader.

**Draize Eye Test**

To evaluate the ocular toxicity of GA in mouse models, a Draize eye test was performed as described previously. Briefly, the left eyes of the mice were treated with 5 μL of GA (10^4 and 100 μg/mL), and the right eyes were treated with PBS 3 times a day as a control. At 1, 3, and 5 days post-treatment, 1 μL of 1% luciferin sodium (Meilunbio, Dalian, China) was dropped into the conjunctival sac of each mouse, and corneal fluorescein staining was examined by slit-lamp microscopy under cobalt blue light (n = 5/group/time). Corneal fluorescein staining (CFS) scores were evaluated according to the Organization for Economic and Co-Operative Development (OECD) grading scale, which involved weighting and summing six components of directly observable changes in the eye's anterior segment, including the density and area of corneal opacification, the severity of iritis, conjunctival redness, edema, and discharge.

**Biofilm Formation**

A colorimetric microtiter plate assay was performed to determine the biofilm-forming capacity of *A. fumigatus* exposed to different concentrations of GA. Conidia suspension (3 × 10^7/mL) was transferred into a 24-well plate with 1000 μL per well and incubated for 48 hours at 37°C. After biofilm formed, wells were gently decanted and rinsed with sterilized PBS. Then, 1000 μL Sabouraud medium containing GA (0, 12.5, 25, 50, 100, and 200 μg/mL) was added to each well in triplicate and incubated for 48 hours at 37°C. After air drying, biofilms were fixed with glutaraldehyde and stained with 100 μL per well of 1% crystal violet (Solarbio) for 15 minutes. Unbound dye was washed off with PBS. After air drying, dye bound to biofilm was released with 100 μL of 95% ethanol per well without shaking. The OD of each well was measured at 620 nm using a microplate reader.

**Calcofluor White Staining**

Calcofluor white staining was conducted to determine the cell wall integrity of *A. fumigatus*. GA was diluted in different concentrations (0, 50, 100, and 200 μg/mL) with a Sabouraud medium and was incubated with *A. fumiga-

**Propidium Iodide Staining**

To examine the cell membrane integrity of *A. fumigatus*, PI staining was performed as previously described. The conidia suspension of *A. fumigatus* (2 × 10^6 hyphal fragments/mL) was seeded on a 6-well plate and incubated at 37°C for 24 hours. Then, the hyphae were washed, centrifuged (12,000 rpm for 10 minutes), transferred to a new 6-well plate, and then incubated with 0, 50, 100, and 200 μg/mL GA at 37°C for 24 hours. The 8 μg/mL NAT was used as a positive control. After harvesting and washing with sterile PBS, the fungal hyphae were incubated with PI (Leagene Biotechnology, Beijing, China) solution for 15 minutes at RT in the dark. Images were photographed with a fluorescence microscope (Zeiss Axiovert, Jena, Germany; at 20× magnification).

**Fungal Adhesion Assay**

HCECs were inoculated to the chambered slides (four/slide) and incubated at 37°C until 30% confluency. After being rinsed with sterile PBS, cell chambers were incubated with conidia suspension (3 × 10^4/mL) for 1 hour and treated with GA at 100 μg/mL for 3 hours at 37°C. Then, the slides were washed with sterile PBS, air-dried, and stained through hematoxylin and eosin (H&E). The number of spores that adhered to the surface of HCECs was counted, averaged, and expressed as the number of adherent spores/cells. All specimens were observed and photographed by optical microscopy (Axio Vert; Zeiss, Jena, Germany; at 400× magnification).

**A. fumigatus Keratitis Mouse Models**

Healthy female 8-week-old C57BL/6 mice (SiPeiFu Biotechnology Co. Ltd., Beijing, China) weighed between 20 and 30 grams (g) were treated according to the ARVO Statement regarding the Use of Animals in Ophthalmology and Vision Research. The establishment of the *A. fumigatus* keratitis model was in accordance with Zhan et al. Mice were abdominally anesthetized with 4 mL of 8% chloral hydrate, and each left eye was a blank control. The right corneal stroma was injected with 2.5 μL *A. fumigatus* conidia suspension (3 × 10^7/mL) with a microsyringe (10 μL; HAMILTON, type 1701) under a slit-lamp microscope (DS126431; Cannon, Japan; at 4× magnification). After 4 hours, 5 μL of GA (100 μg/mL) was dripped on the right eye 3 times a day in the GA-treated groups, and 5 μL of PBS was dripped on the right eye 3 times a day as a control. The 8 μg/mL NAT was used as comparator treatment. Based on the observation under a slit lamp at 1, 3, and 5 days post-infection (p.i.), the severity of keratitis (n = 8/group) was evaluated by clinical score referred to the 12-point scoring system. Briefly, each aspect of opacity density, opacity area,
and surface regularity scores were 0 to 4 points. A total score of 0 to 5 is a mild infection, 6 to 9 is a moderate infection, and 10 to 12 is a severe infection. The corneas at 3 and 5 days p.i. were harvested for RT-PCR. The corneas at 3 days p.i. were removed for myeloperoxidase (MPO). The mouse eyeballs at 3 days p.i. were removed for H&E staining, PAS staining, and immunofluorescence staining (IFS).

### Myeloperoxidase Assay

Mice corneas at 3 days p.i. (n = 5/group) were harvested and homogenized, then followed with the instruction of the MPO test kit (Nanjing Jiancheng Bioengineering Institute, Nanjing, China). MPO was measured spectrophotometrically at 460 nm at 37°C. The line slope was related to the MPO activity (unit/g cornea).

### Hematoxylin and Eosin Staining

Eyeballs of C57BL/6 mice (n = 3/group) were harvested at 3 days p.i. were cut in optimum cutting temperature (O.C.T.) compound (Sakura Tissue-Tek, Torrance, CA, USA). After liquid nitrogen froze, eyeballs were sectioned into 9 μm under cryostat and fixed in acetone. Slices were treated by conventional H&E staining and photographed under an optical microscope (Axio Vert; Zeiss, Jena, Germany; at 200× magnification).

### Periodic Acid Schiff Staining

Mice eyeballs at 3 days p.i. (n = 3/group) were harvested, embedded, and frozen in O.C.T., and then sectioned by 9 μm under cryostat. Sections were stained with a glycogen PAS staining kit (Leagene, Beijing, China) following the manufacturer's instructions. Each slice was photographed under an optical microscope (Axio Vert; Zeiss, Jena, Germany; at 400× magnification).

### Immunofluorescence Staining

The immunofluorescence step refers to the previous study. The mouse eyeballs (n = 3/group) at 3 days p.i. were removed, embedded, and frozen at O.C.T., then sliced into 9 μm-thick tissue sections. After being blocked with goat serum (1:100, Solarbio) for 30 minutes at 37°C, the sections were incubated with anti-NIMP-R14 antibody (1:100, Santa Cruz Biotechnology, Santa, Cruz, CA, USA) at 4°C overnight. Then, slices were stained with FITC-conjugated goat anti-mouse antibodies (1:50; Elabscience, Wuhan, China) at 4°C overnight. The slides were washed with PBS, and then incubated with anti-NIMP-R14 antibody (1:100; Santa Cruz Biotechnology, Santa, Cruz, CA, USA) at 4°C overnight. After washing, the slides were stained with 4′,6-diamidino-2-phenylindole dihydrochloride (DAPI; 1:5; Solarbio). The slides were observed under the fluorescence microscope (Zeiss Axio Vert, Jena, Germany; at 200× magnification).

### Real-time PCR

Total RNA of RAW264.7 cells and mouse corneas (n = 5/group) were extracted using the RNAiso Plus reagent (TaKaRa, Dalian, China) and quantified by NanoDrop ND-1000 Spectrophotometry (Thermo Fisher Scientific). RNA samples were reverse transcribed using HiScript III-RT SuperMix-cjj (Vazyme, Nanjing, China) to produce a cDNA template. The procedure of RT-PCR was referred to in previous experimental research. The corresponding primer pair sequences used for RT-PCR are shown in the Table.

### Enzyme-Linked Immunosorbent Assay

RAW264.7 cells cultured in 6-well plates were stimulated with inactive *A. fumigatus* hyphae for 1 hour, then treated with 100 μg/mL GA for 24 hours. The supernatant was collected and centrifuged. The ELISA kit (R&D Systems, Minneapolis, MN, USA) was used to detect IL-1β and TNF-α protein levels according to the manufacturer’s instructions. The absorbance was measured at 450 nm with a microplate reader.

### Western Blot

RAW264.7 cells were lysed in RIPA buffer (Solarbio) containing PMSF (Solarbio) and phosphatase inhibitor (MCE), sonicated, and centrifuged at 12,000 rpm for 5 minutes. BCA kit (Elabscience) was used for protein concentration determination. Total proteins were separated on SDS-PAGE and transferred onto polyvinylidene difluoride (PVDF) membrane (Solarbio). The membranes were blocked in blocking buffer (Solarbio) and incubated with primary antibodies against GADPH (1:2000; Elabscience), HO-1 (1:1000; Abcam), and
Nrf2 (1:1000; Abcam) at 4°C overnight, followed by incubation with goat anti-rabbit secondary antibodies at 37°C for 1.5 hours. Blots were visualized using enhanced chemiluminescence (ECL; Thermo Fisher Scientific, Waltham, MA, USA).

### Statistical Analysis

The experimental data were analyzed by GraphPad Prism8.0.2 (GraphPad, San Diego, CA, USA) and expressed by mean ± SEM. The unpaired, two-tailed Student’s t-test was used to analyze the statistical significance between the two groups. One-way analysis of variance (ANOVA) with Bonferroni’s post hoc test was performed to examine the difference among three or more groups. The clinical score in different groups was determined by the Mann-Whitney U test. Image J (National Institutes of Health, Bethesda, MD, USA) was used to analyze the images of IFS and Western blot. Any P < 0.05 was considered as significant (ns = no significance). All experiments were repeated at least three times to ensure accuracy.

### RESULTS

#### Evaluation of GA Toxicity In Vitro and In Vivo

CCK-8 assay and Draize eye test were performed to determine the toxicity of GA in HCECs and an A. fumigatus mouse model, respectively. In HCECs, GA did not affect the cell viability at 100 μg/mL for 48 hours (Fig. 1A). However, GA started inhibiting cell viability at 200 μg/mL (see Fig. 1A), suggesting that GA was cytotoxic to HCECs at 200 μg/mL. In healthy mouse cornea, no pathological or morphological damage was observed in the corneal epithelium of mice treated with 100 μg/mL GA at days 1, 3, and 5 (Fig. 1B). Thus, GA with a concentration of no more than 100 μg/mL was considered safe and applied to the subsequent in vitro and in vivo experiments.

#### Antifungal Effects of GA In Vitro

We demonstrate that GA has antifungal effects on \emph{A. fumigatus} \emph{in vitro}. GA started to inhibit \emph{A. fumigatus} growth at 100 μg/mL (Fig. 2A) and markedly inhibited the biofilm formation of \emph{A. fumigatus} starting from 25 μg/mL in a dose-dependent manner (Fig. 2B). H&E staining revealed that the number of \emph{A. fumigatus} conidia adherent to HCECs was significantly reduced in the 100 μg/mL GA-treated group compared with the PBS group (Figs. 2C, 2D). This indicated that GA suppressed \emph{A. fumigatus} adhesion at 100 μg/mL. Moreover, calcofluor white staining and PI staining were performed to evaluate the integrity of the fungal cell wall and membrane. Less fungal cell wall components labeled by calcofluor white were detected in GA-treated \emph{A. fumigatus} than in PBS control (Fig. 2E), suggesting GA disrupted the cell wall integrity. Upon GA treatment, the fluorescent intensity of nuclear and chromosomes exposed from dead fungal cells with impaired membrane was enhanced (Fig. 2F), which also supported that GA may break \emph{A. fumigatus} cell membranes. These findings reveal the antifungal property of GA, which is associated with inhibiting...
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FIGURE 2. Antifungal effects of GA in vitro. MIC of GA at concentrations of 0, 12.5, 50, 100, 200, 400, and 800 μg/mL for 48 hours in HCECs. GA started to inhibit A. fumigatus growth at 100 μg/mL (A). Biofilm formation of A. fumigatus exposed to different concentrations of GA (0, 12.5, 25, 50, 100, and 200 μg/mL) for 48 hours. GA markedly inhibited biofilm formation of A. fumigatus starting from 25 μg/mL in a dose-dependent manner (B). H&E staining based fungal adhesion assay showed the number of A. fumigatus conidia adherent to HCECs was significantly reduced in the 100 μg/mL GA-treated group compared with PBS control (×400 magnification) (C). Quantitative results of H&E staining (D). Calcofluor white staining of A. fumigatus treated with PBS, Caspofungin acetate (positive control), or different concentrations of GA (50, 100, and 200 μg/mL) for 48 hours in vitro (×20 magnification). GA disrupted the cell wall integrity in a dose-dependent manner (E). PI staining of A. fumigatus treated with PBS, natamycin (positive control), or different concentrations of GA (50, 100, and 200 μg/mL) for 24 hours in vitro (×20 magnification) (F). All data were presented as mean ± SEM. MIC and biofilm formation results were analyzed by one-way ANOVA with Bonferroni's post hoc test, and an unpaired, two-tailed Student's t-test was used to compare fungal adhesion results.

biofilm formation and adhesion and disrupting the fungal cell membrane integrity.

GA Improved A. Fumigatus Keratitis Outcomes in Mice

To explore the therapeutic effect of GA, 100 μg/mL GA was applied to mice with A. fumigatus keratitis, and clinical scores were used to evaluate the severity of keratitis at 1, 3, and 5 days p.i. Slit-lamp photographs illustrated less ulcer area in GA-treated corneas at 3 and 5 days p.i. compared to PBS control (Fig. 3A). Consistently, GA treatment versus PBS significantly decreased the clinical scores at 3 and 5 days p.i., whereas there was no difference observed at 1-day p.i. (Fig. 3B), indicating that GA could ameliorate the severity of A. fumigatus keratitis at 3 and 5 days p.i. However, there was no significant difference in clinical scores between GA and the natamycin group, suggesting that GA treatment had the same therapeutic effects as natamycin. We then examined the fungal load using PAS staining. Fewer hyphae were shown in GA and natamycin-treated corneas compared with the PBS group (Fig. 3D). Therefore, we hypothesize that GA may improve A. fumigatus keratitis prognosis in mice by inhibiting fungal growth and alleviating the inflammatory response, and the therapeutic effects of GA might be comparable to natamycin.

GA Repressed Neutrophil Infiltration and MPO Activity In Vivo

In order to determine the anti-inflammatory effect of GA, neutrophil infiltration and MPO activity were analyzed in GA-, natamycin-, and PBS-treated mice with A. fumigatus keratitis. IFS revealed that 100 μg/mL GA significantly inhibited the infiltration and accumulation of neutrophils in the corneal stroma at 3 days p.i. compared with PBS. This inhibitory impact on neutrophil recruitment was also found in the natamycin-treated group (Figs. 4A, 4B). Similarly, GA- and natamycin- versus PBS-treated mice corneas showed markedly reduced MPO activity at 3 days p.i. (Fig. 4C). These results suggested that GA and natamycin treatment could inhibit neutrophil vitality in A. fumigatus keratitis mice.

GA suppressed pro-inflammatory cytokine production and enhanced the expression of Nrf2 and HO-1 in vitro and in vivo. We demonstrate that GA downregulated the expression of pro-inflammatory cytokines in RAW264.7 cells and A. fumigatus keratitis mouse models. The inactivated hyphae were used to stimulate RAW264.7 cells for 1 hour, followed
FIGURE 3. GA improved *A. fumigatus* keratitis outcomes in mice. Representative slit-lamp photographs (A) and corresponding clinical scores (B) of *A. fumigatus* keratitis mice corneas with PBS, 100 μg/mL GA, and 8 μg/mL natamycin treatment at 1, 3, and 5 days p.i. (*n* = 8/group/time). PAS staining (for fungal hyphae) (C) and H&E staining (D) of *A. fumigatus* keratitis mice eyeballs with PBS, 100 μg/mL GA, and 8 μg/mL natamycin treatment at 3 days p.i. (*n* = 3/group; ×100). The clinical score in different groups was analyzed using the Mann-Whitney *U* test (ns: no significance).

FIGURE 4. GA repressed neutrophil infiltration and MPO activity in vivo. Representative IFS photographs of *A. fumigatus* keratitis mice corneas treated with PBS, 100 μg/mL GA, and 8 μg/mL natamycin at 3 days p.i. (*n* = 3/group). Green: NIMP-R14 labeled neutrophils; Blue: DAPI stained nucleus (×200 magnification) (A). IFS quantitative analysis of area fractions of neutrophils (B). Quantitative analysis of MPO activity levels of *A. fumigatus* keratitis mice corneas treated with PBS, 100 μg/mL GA, and 8 μg/mL natamycin at 3 days p.i. (*n* = 5/group) (C). All data were mean ± SEM and analyzed by an unpaired, two-tailed Student’s *t*-test.
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Figure 5. GA suppressed pro-inflammatory cytokine production in RAW264.7 cells and *A. fumigatus* keratitis mouse models. RAW264.7 cells were stimulated with inactivated *A. fumigatus* hyphae for 1 hour and treated with 100 μg/mL GA, 8 μg/mL natamycin, and PBS for 8 hours for RT-PCR and 24 hours for ELISA. RT-PCR results of the mRNA levels of IL-1β (A), TNF-α (B), COX-2 (C), and LOX-1 (D), and ELISA results of IL-1β (E), TNF-α (F), IL-6 (G), COX-2 (H), and LOX-1 (I) were detected at 3 and 5 days post-treatment (n = 6/group/time). All data were mean ± SEM. RT-PCR and ELISA result from RAW264.7 cells were analyzed by one-way ANOVA with Bonferroni’s post hoc test. An unpaired, two-tailed Student’s t-test was used for RT-PCR results from *A. fumigatus* mouse models (N: normal; Af: *A. fumigatus*; ns: no significance).

by 100 μg/mL GA, 8 μg/mL natamycin, and PBS treatment. Both mRNA and protein production of pro-inflammatory factors in RAW264.7 cells was enhanced by *A. fumigatus* treatment (Figs. 5A–F). In comparison with PBS treated group, the mRNA levels of IL-1β (see Fig. 5A), TNF-α (see Fig. 5B), cyclo-oxygenase-2 (COX-2; see Fig. 5C), and lectin-like oxidized low-density lipoprotein receptor-1 (LOX-1; see Fig. 5D) were decreased in GA treated cells at 8 hours post-treatment. Consistently, protein levels of IL-1β (see Fig. 5E), and TNF-α (see Fig. 5F) were reduced in the GA-treated group at 24 hours post-treatment. Whereas in natamycin-treated cells, less reduction was found in the expression of these cytokines but with no statistical significance. In *A. fumigatus* keratitis mouse models, the mRNA expression of IL-1β (Fig. 5G), TNF-α (Fig. 5H), COX-2 (Fig. 5I), and LOX-1 (Fig. 5J) was significantly elevated by *A. fumigatus* infection and markedly repressed by 100 μg/mL GA and 8 μg/mL natamycin versus PBS eye drop treatment at 3 and 5 days p.i. No statistical significance was observed between GA- and natamycin-treated groups. These results indicated that the anti-inflammatory role of GA could be associated with the Nrf2/HO-1 signaling pathway.
**Figure 6.** GA enhanced Nrf2 and HO-1 expression in RAW264.7 cells and *A. fumigatus* keratitis mouse models. RAW264.7 cells were stimulated with inactivated *A. fumigatus* hyphae for 1 hour and treated with 100 μg/mL GA or PBS for 8 hours for RT-PCR and 24 hours for Western blot. RT-PCR results of the mRNA levels of Nrf2 (A) and HO-1 (B) in RAW264.7 cells. Western blot results of HO-1 (E) in RAW264.7 cells and the corresponding quantitative analysis (D). Western blot results of HO-1 (H) in RAW264.7 cells and the corresponding quantitative analysis. In an *A. fumigatus* mouse model, mice corneas were treated with 100 μg/mL GA or PBS eye drop 3 times/day, and the mRNA levels of Nrf2 (G), HO-1 (H) were detected at 3 and 5 days p.i. (*n* = 6/group/time). All data were mean ± SEM. RT-PCR and Western blot results from RAW264.7 cells were analyzed by one-way ANOVA with Bonferroni’s post hoc test. An unpaired, two-tailed Student’s *t*-test was used for RT-PCR results from *A. fumigatus* mouse models (N: normal; Af: *A. fumigatus*; ns: no significance).

**DISCUSSION**

Our study investigates GA’s therapeutic effect on FK in vitro and in the *A. fumigatus* keratitis mouse model. We identified that GA at 100 μg/mL had no cytotoxic effects within 48 hours in HCECs and no corneal toxicity for 5 days of treatment. Thus, we considered that 100 μg/mL GA could be applied as a safe concentration for local corneal therapy.

GA is an abundant phenolic compound in various natural foods. It has a tri-hydroxylated phenolic structure, a benzene ring, a carboxylic grouping, and three hydroxyl groups that endow GA antioxidant properties. Studies showed that GA had an antifungal ability to inhibit fungi sporulation, impede germination of saprobe fungi, and destroy biofilm formation. Consistently, we demonstrated that 100 μg/mL GA could repress conidia growth, germination, and biofilm formation of *A. fumigatus*. More importantly, GA reduced the average percentage of adhered conidia adhering to HCECs compared with PBS control, indicating that GA has anti-adhesion capacity. It has been reported that GA could degrade chitin, the major structural component of the phytopathogenic fungi cell walls, by producing the enzyme chitinase. This finding coincides with our staining results, in which 100 μg/mL GA impaired *A. fumigatus* membrane integrity. Besides, GA reduced the activity of CYP51, a key enzyme in the process of fungi ergosterol synthesis. Ergosterol is one of the fungal plasma membrane components, playing a pivotal role in membrane fluidity, permeability, and signal transduction pathways. Thus, we suspect that GA destroys *A. fumigatus* cell membranes and increases membrane permeability by inhibiting the activity of CYP51 enzymes.

In *A. fumigatus* keratitis mice, GA significantly alleviated the severity of FK. PAS staining showed GA obviously decreased the hyphae load in the infected sites, indicating that GA could improve infectious corneal ulcers by limiting the growth of *A. fumigatus*. Similar results were observed in natamycin-treated groups, which suggests that the antifungal effect of GA is comparable to natamycin. On the other hand, the less fungal load is complementary to alleviating inflammatory response and decreasing neutrophil infiltration. Neutrophils are vital inflammatory cells for eliminating intracellular microorganisms. Corneas infected with fungi attract neutrophils released from the limbal vessels to the site of the infection. But the aggregation of neutrophils may result in severe inflammation, which inevitably causes sustained infiltration of inflammatory cells, leading to undesired tissue damage, such as ulceration, cornea perforation, and even vision loss. Earlier studies demonstrated that GA could attenuate neutrophils infiltration and subsequent injury in isoproterenol-induced myocardial infarction. It is in agreement with our results, which showed that 100 μg/mL GA inhibited neutrophils infiltration, alleviated the severity of corneal edema, and made it with more ordered pathological structure in the stroma at 3 days p.i. Clinical scores in GA-treated groups were significantly lower than in PBS-treated groups, and GA-treated corneas were more transparent, with less ulcer area. Therefore, GA markedly contributes to ameliorating corneal damage in *A. fumigatus* keratitis by limiting fungal load and reducing neutrophil recruitment. Interestingly, the natamycin-treated group had a similar therapeutic effect in vivo experiments as GA. Although this therapeutic effect of natamycin could be attributed to its antifungal property, it is still unclear whether natamycin has
a similar anti-inflammatory potential as GA. Thus, in vitro experiments in RAW264.7 cells were conducted to explore the anti-inflammatory activity of GA and natamycin.

During the inflammation process, neutrophils also promote the release of pro-inflammatory chemokines, facilitating the inflammatory response.50,51 These released chemokines and cytokines recruit more neutrophils and other immune cells, activating an inflammatory cascade.45 IL-1β and TNF-α are important pro-inflammatory cytokines involved in the corneal anti-fungal immune response; mainly produced by activated monocytes, mediating acute inflammatory responses.17 LOX-1 belongs to the C-type lectin family, playing an essential role in the corneal antifungal immune response.18 COX-2, induced by proinflammatory cytokines, generates prostaglandins, which are supposed to cause inflammation.48 In our study, the inactivated A. fumigatus hyphae were used to stimulate cells. The deactivated fungi have no invasiveness and only stimulate the cells’ immune system through residual antigens. We showed that GA treatment decreased A. fumigatus stimulated overexpression of IL-1β, TNF-α, LOX-1, and COX-2 in vitro and in vivo, whereas natamycin only decreased them in vivo. This is because natamycin played an antifungal role in the process of A. fumigatus keratitis in mice. Less fungal load resulted in less extent of inflammation, thereby reducing the expression of inflammatory cytokines. However, in RAW264.7 cells, natamycin could not reduce the enhanced expression of pro-inflammatory cytokines stimulated by inactivated A. fumigatus hyphae, suggesting natamycin has no anti-inflammatory property. Instead, GA showed markedly inflammation-inhibiting activity in vitro.

We have recently described the involvement of the Nrf2/HO-1 signaling pathway in A. fumigatus keratitis.17,20 and GA treatment significantly enhanced A. fumigatus induced overexpression of NF-2 and HO-1 in FK mice models and RAW264.7 cells. These results inspired us that the anti-inflammatory mechanism of GA may be associated with the Nrf2/HO-1-1 pathways. In the LPS-induced chronic obstructive pulmonary disease (COPD) model, GA treatment resulted in increased protein levels of Nrf2 accompanied by an increment in the transcription of HO-1.39 Consistently, we observed that the expression of Nrf2 and HO-1 was elevated under the treatment of GA in A. fumigatus infected RAW264.7 cells. Although GA induced the enhancement of the Nrf2/HO-1 signaling pathway is validated in infected RAW264.7 cells, natamycin could not reduce the enhanced expression of pro-inflammatory cytokines stimulated by inactivated A. fumigatus hyphae, suggesting natamycin has no anti-inflammatory property. Instead, GA showed markedly inflammation-inhibiting activity in vitro.

In summary, our study investigates the antifungal and anti-inflammatory effects of GA on A. fumigatus keratitis in vitro and in vivo. We showed that GA could ameliorate A. fumigatus keratitis via inhibiting fungal growth and spore germination, decreasing neutrophils recruitment and infiltration, and increasing the expression of Nrf2 and HO-1, as well as reducing the expression of inflammatory cytokines, such as IL-1β, TNF-α, LOX-1, and COX-2, which suggests that GA could be a potentially effective drug for FK treatment.

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