Amoebicidal Activity of Poly-Epsilon-Lysine Functionalized Hydrogels

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Acanthamoeba keratitis (AK) is a rare infection but may result in severe visual impairment or blindness.1,2 Acanthamoeba are opportunistic protozoan, amoebic parasites that live naturally in soil, air, and aquatic habitats in one of two forms: trophozoites (mobile active form) or cysts (highly resilient, double-walled, dormant cysts).3 Trophozoites and cysts are both able to infect the cornea, skin, and central nervous system.4 The conversion of trophozoites into cysts results from adverse environmental conditions.5,6

The corneal epithelium normally provides a barrier to invading pathogens, but Acanthamoeba invade the cornea through a breach in the corneal epithelium. Contact lens (CL) wear is a significant risk factor for the development of AK, and up to 80% of cases of AK are associated with the use of CLs or lens solutions contaminated with waterborne Acanthamoeba. In particular, there has been an increase in AK associated with the increased use of soft CLs.7–9 In addition, a lack of compounds effective against Acanthamoeba cysts in CL solutions and tap water quality are also contributing risk factors of AK in CL wearers.10–13

To initiate infection, trophozoites adhere to the corneal epithelium via mannose-decorated glycoproteins and invade the cornea as enzymes such as matrix metalloproteases, serine, and cysteine proteases degrade the corneal stroma.14 Within the stroma, trophozoites may encyst, which may lead to a persistent, relapsing keratitis.15 Successful treatment of AK requires early diagnosis,16 but the diagnosis is challenging and may be mistaken for a bacterial, viral, or fungal keratitis. Treatment of established AK is difficult and currently reliant on intensive treatment with chlorhexidine, polyhexamethylene biguanide, brolene, or hexamide, with variable and often poor outcomes.17

Poly-epsilon-lysine (pεK) is a cationic peptide with intrinsic antimicrobial properties and that reportedly disrupts the...
cell membrane and cell wall of microbes showing broad-spectrum antimicrobial activity against Gram-positive and Gram-negative bacteria, yeasts, and fungi. Our previous studies have demonstrated pK hydrogels are effective against laboratory strains of *Escherichia coli*, *Staphylococcus aureus*, and *Pseudomonas aeruginosa* while having no toxicity toward corneal epithelial cells. Further functionalization of the free amine groups of the hydrogel with covalently bound pK (pK+) increased pK levels with a 10-fold increase in amine functionality due to additional pK molecules when compared with nonfunctionalized hydrogels. Further studies associated this increase in pK with increased antimicrobial activity against *P. aeruginosa*, reducing the number of viable bacteria in vitro and ex vivo corneal infection models.

The aim of this study was to evaluate whether pK offers amoebicidal activity against *Acanthamoeba* in both trophozoite and cyst forms. The effects of pK solution, pK hydrogel, and pK+ hydrogel treatment upon both trophozoite and cyst forms of *Acanthamoeba* were investigated at 24 hours and 7 days in vitro, and significant toxicity was demonstrated with both forms. Using an ex vivo porcine cornea model of AK, we showed that no *A. castellanii* were detected in the stroma after application of pK+ or pK hydrogels.

**Methods**

**pK Solution Preparation**

pK (Bainafo; Zhengzhou Bainafo Bioengineering Co., Ltd., Zhengzhou 450006, Henan Province, China) (20 mM) was prepared in sterile PBS (Oxoid, Hampshire, UK) to a final concentration of 2×10^4 amoeba/mL in PYG media. Trophozoites and Cysts

Trophozoites and cysts were used at 2×10^3 amoebae per well (100 μL) in a 96-well plate. Then, 100 μL of pK solution at 2× concentration was added to each well to a final working volume of 200 μL, with a final pK concentration range of 0 to 4.34 mM. Chlorohexidine (CHX) (Sigma-Aldrich) at 0.02% was used as a positive control for toxicity of trophozoites and cysts. Toxicity of pK against *A. castellanii* was assessed using propidium iodide (PI) (Thermo Fisher Scientific, Loughborough, UK) added to wells at a final concentration of 1 μg/mL. Dead trophozoites/cysts imaged using a fluorescent microscope and the number of dead red-labeled *A. castellanii* were compared with the number of nonstained trophozoites and cysts per field of view expressed as percentage dead (%) compared with total live and dead.

**pK Cytotoxicity Against Human Corneal Epithelial Cells**

Human corneal epithelial (hCE-T) cells (donated by Kaoru Araki-Sasaki, Kansai Medical University, Moriguchi, Japan) were cultured at 37°C and 5% CO2. hCE-T cells were cultured in Dulbecco's modified Eagle's medium/Ham's F12 media (DMEM/F12) (Thermo Fisher Scientific) containing 5% (v/v) fetal calf serum (Biosera; Labtech, Heathfield, UK) with 1% (v/v) fungizone and penicillin/streptomycin (Sigma-Aldrich) supplementation. hCE-T cells were seeded in 96-well plates at 1×10^4 hCE-T cells and incubated for 3 days until 80% confluence was achieved. Media were replaced with fresh media containing pK (0–17.36 mM) and incubated at 37°C and 5% CO2 for 24 hours and 7 days. At each time point, viability/toxicity assays were performed using the Live/Dead viability/cytotoxicity kit for mammalian cells (Thermo Fisher Scientific) following the manufacturer's instructions. hCE-T cells were incubated with Calcein-AM and ethidium homodimer 1 for 30 minutes and fluorescent live (green)/dead (red) cells imaged using an imaged Zeiss Apotome (Germany) live-cell microscope.

**pK Hydrogel Toxicity Assay Against Trophozoites and Cysts**

Under sterile conditions, 10-mm discs of pK and pK+ hydrogels or commercial hydrophilic cast-molded CL (Hydrogel CL; Filcon II 2, 77% water content; Ultradisc, Leighton Buzzard, UK) were added to 48-well plates using sterile forceps. Trophozoites or cysts were used at 1×10^4 amoebae per well in 500 μL of PBS (nutrient deplete) or PYG media (nutrient replete) and incubated at 28°C for either 24 hours or 7 days. Toxicity of pK hydrogels was measured as described above following the addition of 1 μg/mL propidium iodide to label dead trophozoites and cysts.
Ex Vivo Porcine Cornea Keratitis Model

Porcine eyes were obtained from a local abattoir within 6 hours of slaughter, and corneas were processed with agarose supports to establish an air-liquid interface ex vivo organ culture model as previously described by Kennedy et al. and Supplementary 1. Trophozoites were suspended in 10 μL PYG media containing $1 \times 10^5$ amoebae and pipetted onto an air-dried region of the central cornea and allowed to absorb for 30 minutes at room temperature (assessed by visual analysis). Once the droplet was no longer visible, 10-mm discs of commercial CL, pεK, or pεK+ hydrogels were applied to corneas. Control infected and noninfected corneas were run in parallel. DMEM (+10% FBS) was added up to the scleral boundary of the cornea ($\sim$3 mL) and incubated for 7 days at 37°C, 5% CO₂.

Following infection, pεK+ and pεK hydrogel lenses and commercial hydrogel CLs were removed from corneas, resuspended in PYG media, and vortexed for 30 seconds to remove *A. castellanii* from lenses, and the solution was transferred to a 6-well plate to monitor regrowth. Quantification of *A. castellanii* infection of corneas was achieved by trephining (Blink Medical Ltd, Solihull, UK) 10-mm discs from the central cornea. To isolate trophozoites, corneal discs were cut into quadrants and further quartered, resuspended in 3 mL PBS, and briefly vortexed, followed by homogenization using a Qiagen TissueRuptor (Qiagen, Manchester, UK) for 15 seconds. The homogenate was centrifuged at 1000 $\times$ g for 10 minutes, resuspended in PYG media (+ glucose 1.5% (w/v)), transferred to a well of a 6-well plate, and cultured for 7 days to monitor regrowth. *A. castellanii* from each experimental condition were imaged at 7 days using a Nikon (Nikon Europe BV, Netherlands) Ti-E microscope and quantified following resuspension of *A. castellanii* in PYG media using a hemocytometer.

Histology

A separate set of corneas from each experimental condition was fixed in 10% (v/v) neutral buffered formalin (Sigma-Aldrich) for 18 hours. Corneas were processed using a Leica ASP300 (Nussloch, Germany) tissue processor. Paraffin-embedded tissue was sectioned at a thickness of 5 μm and stained with periodic acid–Schiff (PAS) (Abcam, Cambridge, UK) following the manufacturer’s protocol for positive detection of chitin in the cell wall of *A. castellanii*. Tissue sections were imaged using a Nikon CI upright microscope using a 20× and 40× objective.
FIGURE 2. Dose response of pEK solutions against human corneal epithelial cell line hCE-T at 24 hours and 7 days. pEK solutions (0–17.36 mM) added to hCE-T cells and dead cells were labelled with PI (red) and live cells labeled with calcein AM (green). (A) (i) Quantification of the percentage dead hCE-T cells in the total population at 24 hours. The red data point indicates the toxic dose of pEK toward trophozoites and cysts. (ii) Corresponding fluorescent images of hCE-T cells; dead cells were labeled with PI (red) and live cells labeled with calcein AM (green). (B) (i) Quantification of the percentage dead hCE-T cells in the total population at 7 days. The red data point indicates the toxic dose of pEK toward trophozoites and cysts. (ii) Corresponding fluorescent images of hCE-T cells; dead cells were labeled with PI (red) and live cells labeled with calcein AM (green). Scale bars: 50 μm.

Statistical Analysis

Experiments were performed in triplicate (n = 3), with three wells per experiment and five fields of view in each well. One-way ANOVA was performed with a post hoc Tukey’s analysis, and P < 0.05 was considered significant. Statistical analysis was performed using Prism software version 8.02.263 (GraphPad Software, La Jolla, CA, USA). Error bars are shown as standard deviations of biological replicates.

RESULTS

Amoebicidal, Both Trophozoicidal and Cysticidal, Activity of pEK Solution Against *A. castellanii*

Trophozoicidal and cysticidal effects of pEK solution (0–4.34 mM) on *A. castellanii* were assessed following incubation for 24 hours and 7 days. Toxicity was assessed by the number of dead, red PI-stained *A. castellanii* compared to nonstained live *A. castellanii* and expressed as percentage death at each dose.

After 24-hour pEK treatment of trophozoites, 80% (SD ± 6.14%) toxicity was observed at 0.54 mM pEK (∗P < 0.0001) compared to untreated trophozoites, and no further increase in toxicity above 80% was achieved at any higher concentrations (Fig. 1Ai). After 24 hours, trophozoites remained attached to the tissue culture plastic surface (TCPS) surface showing characteristic morphology, while treatment with CHX showed 100% toxicity. An increased dose of pEK increased the number of dead *A. castellanii*, while live *A. castellanii* detached from the surface and did not show characteristic trophozoite morphology. At 7 days posttreatment, amoebicidal effects of pEK upon trophozoites were observed at lower pEK doses (Fig. 1Aii). After 7-day treatment with lower concentrations of pEK (0.016 mM), the number of dead trophozoites increased to 99% (SD ± 1.04%) (∗P < 0.0001).

Effects of pEK solution upon cysts after 24-hour treatment showed 76% (SD ± 1.50%) cyst death at 0.54 mM and above (∗P < 0.0001) (Fig. 1Bi). At 7 days (Fig. 1Bii), treatment with pEK solution led to increased number of dead cysts at lower pEK concentrations. Toxicity against cysts reached 77% (SD ± 2.14%) at 0.004 mM (∗P < 0.0001), and cyst death did not significantly increase (∗P < 0.05) further with increased pEK concentrations. At 24 hours and 7 days, only live cysts were visible in control images, while in comparison, CHX showed red PI-stained cysts in the field of view.

Toxicity of pEK Solution Against hCE-T Cells

Maximum toxicity of pEK treatment toward trophozoites and cysts was observed at 0.54 mM at 24 hours and at 0.016 mM and 0.004 mM at 7 days for trophozoites and cysts, respectively. Toxicity of pEK solution at different concentrations was determined upon confluent monolayers of hCE-T cells.

After 24-hour treatment with pEK solution, confluent monolayers of live hCE-T cells (stained green with calcein-
AM dye) were observed after application up to 0.54 mM peK, with comparable dead hCE-T cells (stained red with PI) to untreated controls ($P > 0.05$) (Figs. 2Ai, Aii). At 1.09 mM peK and above, live cells were reduced, with increased dead cells per image compared to untreated controls ($P < 0.05$). At 7 days, treatment with peK 0 to 0.067 mM was not toxic to hCE-T cells ($P > 0.05$) and, at 0.135 mM and above, proved toxic to hCE-T cells with a reduction in live hCE-T cells ($P < 0.05$) (Figs. 2Bi, Bii).

**Trophozoicidal Activity of peK Hydrogels Against *A. castellanii***

Toxicity of peK$^+$ hydrogels toward trophozoites at 24 hours and 7 days was assessed. At 24 hours, trophozoites grown on TCPS, commercial hydrogel CL, and nonfunctionalized peK hydrogels adhered to the surface, with few cysts forming and minimal trophozoite death, identified by PI staining in $<1\%$ of the population (Figs. 3Ai, Aii) with no significant difference compared to untreated controls ($P < 0.05$). Trophozoites cultured on peK$^+$ hydrogels, however, showed 98.52% (SD $\pm$ 1.82%) ($P < 0.0001$) death, similar to CHX (97.93% [SD $\pm$ 1.38%]; $P < 0.0001$) (Fig. 3Aii), compared to untreated controls, commercial CLs, and peK hydrogels. There was no significant difference in death between peK$^+$ hydrogels and CHX ($P = 0.094$).

At 7 days, growth on peK$^+$ hydrogels showed 83.31% (SD $\pm$ 7.96%) toxicity against trophozoites compared to untreated controls ($P < 0.0001$). The total number of live and dead trophozoites decreased by 7 days compared to 24 hours, with dead trophozoites being degraded within the media, accounting for the lower percentage reduction. In comparison, trophozoites cultured on nonfunctionalized peK hydrogels, commercial hydrogel CL, and TCPS all showed growth of trophozoites with negligible toxicity ($<1\%$ in the total population (Figs. 3Bi, Bii), with no significant difference ($P > 0.05$) compared to untreated controls. Treatment with CHX showed 96.01% (SD $\pm$ 4.12%) dead trophozoites and was slightly more effective compared to peK$^+$ hydrogels ($P = 0.024$).

**Cysticidal Activity of peK Hydrogels Against *A. castellanii***

Cysts were cultured on TCPS, peK and peK$^+$ hydrogels, and commercial hydrogel CLs in both nutrient-rich PYG medium or nonnutrient PBS to mimic dormant or proliferating conditions for 24 hours or 7 days. Cysts cultured in PBS remained in cyst form on TCPS, peK or peK$^+$ hydrogels, or commercial hydrogel CLs. At 24 hours under control untreated conditions, the percentage of dead cysts in the overall population was 13.18% (SD $\pm$ 5.29%). There was an increase in dead cysts when cultured on peK$^+$ hydrogels of 70.59% (SD $\pm$ 10.93%) and CHX of 69.37% (SD $\pm$ 6.68%), both significantly different from untreated controls, peK hydrogels, or commercial CL ($P < 0.0001$) (Fig. 4A). After 7 days,
Amoebicidal Poly-Epsilon-Lysine+ Hydrogels

Toxicity of pɛK+ hydrogels against *A. castellanii* cysts cultured in PBS buffer (nutrient deplete) at 24 hours and 7 days. (A) (i) Graph shows the percentage of dead cysts and (ii) fluorescent images of cysts when cultured on TCPS, commercial hydrogel CL, and pɛK and pɛK+ hydrogels for 24 hours. (B) (i) Graph shows the percentage of dead cysts to total live and dead cysts and (ii) fluorescent images of cysts when cultured on TCPS, commercial hydrogel CL, and pɛK and pɛK+ hydrogels for 7 days. CHX was run as a positive control for cyst death. Scale bars: 50 μm.

Dead cysts increased to 82.32% (SD ± 2.74%) on the pɛK+ hydrogel, and CHX was slightly more effective with 90.83% (SD ± 5.45%) (Fig. 4B) ($P = 0.03$), with both resulting in significantly higher cysticidal activity compared to untreated controls, pɛK hydrogels, and commercial CLs ($P < 0.0001$).

Cysts cultured in PYG medium on TCPS, commercial hydrogel CL, and pɛK hydrogels differentiated into trophozoites at 24 hours (Fig. 5A). In contrast, cysts cultured in PYG medium on pɛK+ hydrogels did not differentiate into trophozoites and showed 75.37% (SD ± 2.84%) reduction in viable cysts with no significant difference compared to CHX (70.27% [SD ± 4.84%]) ($P = 0.68$). Both pɛK+ hydrogel and CHX were significantly different from untreated controls, pɛK+ hydrogel, and commercial hydrogel CL ($P < 0.0001$).

By 7 days, dead cysts on pɛK+ hydrogel had increased to 87.14% (SD ± 5.79%), comparable to CHX with 82.36% (SD ± 6.24%) ($P = 0.69$), and were both significantly different from untreated controls, pɛK+ hydrogel, and commercial hydrogel CL ($P < 0.0001$).

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**Toxicity of pɛK Hydrogels Against *A. castellanii* in Ex Vivo Porcine Corneas**

Porcine corneas were infected with trophozoites for 7 days in the presence of commercial hydrogel CL, pɛK hydrogel, or pɛK+ hydrogel (Fig. 6). Histologic analysis of fixed corneas using PAS to label *A. castellanii* showed there were no *A. castellanii* within the corneal stroma in uninfected corneas. Infected corneas and infected in the presence of commercial hydrogel CL both showed cysts below the corneal surface. Infected corneas incubated with pɛK or pɛK+ hydrogels showed an absence of *A. castellanii* within tissue sections.

Numeration of *A. castellanii* present on lenses after incubation with corneas showed trophozoite regrowth at 7 days from commercial hydrogel CL ($2.68 \times 10^5$ amoebae) and pɛK hydrogel ($1.51 \times 10^5$ amoebae) ($P = 0.0004$) (Figs. 7Ai, Aii). No detectable viable *A. castellanii* retrieved were from the pɛK+ hydrogel, which was significantly different from other hydrogel lenses ($P < 0.0001$).

After 7 days, regrowth of *A. castellanii* from homogenized corneal buttons from infected corneas and corneas from under commercial hydrogel CL showed $4.89 \times 10^5$ and $3.12 \times 10^5$ amoebae per cornea, respectively ($P = 0.0018$) (Figs. 7Bi, Bii). *A. castellanii* regrowth from corneas under pɛK hydrogel was $5.88 \times 10^4$ amoebae per cornea, compared to infected and commercial hydrogel CL ($P < 0.0001$). No detectable *A. castellanii* were retrieved from corneas incubated with pɛK+ hydrogel, which was significantly different compared to infected and commercial hydrogel CL ($P < 0.0001$) corneas but not from pɛK hydrogel corneas ($P = 0.47$).

**DISCUSSION**

This study demonstrated the antimicrobial peptide pɛK is effective at reducing viable *A. castellanii* trophozoites and
cysts, both as a solution and when covalently bound to pεK hydrogels using in vitro assays. pεK was nontoxic to hCE-T cells at doses up to 0.54 mM at 24 hours and up to 0.067 mM at 7 days, incorporating doses that were toxic to A. castellanii. pεK+ hydrogels demonstrated toxicity toward both cysts and trophozoites compared to nonfunctionalized pεK hydrogels and commercial hydrogel CLs. In our ex vivo porcine corneal infection model, pεK and pεK+ hydrogels prevented infection of A. castellanii within the cornea stroma, and reduced numbers of A. castellanii adhered to hydrogels.

Current preventative treatments against AK are targeted at maintaining good hygiene regimes and use of disinfectants within CL solutions. Therapeutic treatments for established AK involve the use of active reagents such as polyhexamethylene biguanide (0.02%) (or CHX 0.02%) and propanidide (0.1%) and hexamidine (0.1%), often in combination.15,20,27 Increased resistance to many of these reagents has been reported in different strains of Acanthamoeba.28,29 We have investigated only one particular strain of Acanthamoeba, but we have examined the effectiveness of pεK+ hydrogels at killing both trophozoites and cysts, the latter of which are particularly difficult to kill. MeniCare Pure CL solution (Menicon Ltd., Nagoya, Japan) contains pεK as an active ingredient, which is effective against Acanthamoeba. It is used as a disinfectant instead of polyhexamethylene biguanide, but this CL solution is only suitable for use with rigid gas-permeable CLs.

The effects of pεK solution against trophozoites and cysts were investigated in both nutrient-rich and nutrient-deplete environments. Toxic effects of pεK against A. castellanii occurred in a dose- and time-dependent manner against both cysts and trophozoites in nutrient-replete and nutrient-deplete environments. The effects of pεK were more effective against the trophozoites, which are easier to treat compared to the cysts.30–32

Current treatments for prevention or treatment of an AK are typically harsh and damaging to the ocular surface, in particular the corneal epithelium.33 We demonstrated that effective doses of pεK were toxic against trophozoites or cysts, respectively, but not to hCE-T cells. pεK is generally regarded as safe, is used in many applications,18 and offers potential use as a treatment for AK.

There are currently no commercial CLs available that offer antimicrobial activity, and CLs themselves provide a potential route of infection into the cornea. Having established pεK solution was toxic to A. castellanii, we demonstrated pεK+ hydrogels showed toxicity at 24 hours and 7 days toward A. castellanii, compared to the commercial CLs tested. pεK+ hydrogels offer inherent antiamoebicidal (both trophozoicial and cysticidal) activity without the potential need for topical application. CLs themselves provide a potential surface for Acanthamoeba to prolifer-
FIGURE 6. Ex vivo corneal infection of *A. castellanii* trophozoites in the presence of pεK⁺ hydrogels. Histologic sections of PAS staining of porcine corneas infected with *A. castellanii* trophozoites for 7 days. Top row left shows no infection in cornea, middle top shows infected cornea (insert highlighting *A. castellanii* cyst), and top right shows *A. castellanii* grown in the presence of a commercial hydrogel CL. Bottom row shows corneas infected with *A. castellanii* in the presence of pεK hydrogel and pεK⁺ hydrogels. Scale bars: 50 μm.

FIGURE 7. Growth of *A. castellanii* from pεK⁺ hydrogels and corneas from ex vivo infection model. (A) Lenses were removed from corneas and cultured in PYG media for 7 days. (i) Phase contrast images of *A. castellanii* regrowth from lenses and (ii) quantification at 7 days. (B) Regrowth of *A. castellanii* from cornea (i) phase contrast images and (ii) quantification at 7 days. Scale bars: 50 μm.
Amoebicidal Poly-Epsilon-Lysine<sup>+</sup> Hydrogels

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Amoebicidal Poly-Epsilon-Lysine\*$^+\,$ Hydrogels


