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Encapsulation into hyaluronic acid-based nanogels improves the selectivity index of the snake cathelicidin Ab-Cath

Miriam E. van Gent, PhDa,⁎, Sylvia N. Kłodzińska, PhDb, Maureen Severin, BSc a, Muhanad Ali, PhDa, Bjorn R. van Dooidewaard, BSc c, Erik Bos, MSc d, Roman I. Koning, Assistant professor d, Jan Wouter Drijfhout, Associate professor e, Hanne M. Nielsen, Professor b, Peter H. Nibbering, Associate professor a

aDepartment of Infectious Diseases, Leiden University Medical Center, 2300 RC Leiden, the Netherlands
bCenter for Biopharmaceuticals and Biobarriers in Drug Delivery (BioDelivery), Department of Pharmacy, Faculty of Health and Medical Sciences, University of Copenhagen, Universitetsparken 2, DK-2100 Copenhagen, Denmark
cDepartment of Cell and Chemical Biology, Leiden University Medical Center, 2300 RC Leiden, the Netherlands
dElectron Microscopy Facility, Department of Cell and Chemical Biology, Leiden University Medical Center, 2300 RC Leiden, the Netherlands
eDepartment of Immunology, Leiden University Medical Center, 2300 RC Leiden, the Netherlands

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Abstract

The antimicrobial peptide Ab-Cath, is a promising candidate for development as treatment for antimicrobial resistant (AMR) bacterial infections. Future clinical use is hampered by Ab-Cath's cationic peptidic nature and limited therapeutic window. Here, we evaluated hyaluronic acid-based nanogels for encapsulation of Ab-Cath to circumvent these limitations. Using microfluidics, monodispersed anionic nanogels of 156–232 nm encapsulating >99 % Ab-Cath were prepared. Unprecedented, lyophilization using polyvinyl alcohol and dextran-40 provided Ab-Cath nanogel protection and allowed easy dose adjustment. Lyophilized and redispersed Ab-Cath nanogels were as effective as Ab-Cath solution in killing AMR Staphylococcus aureus, Acinetobacter baumannii and Escherichia coli in biological fluids, and in reducing S. aureus and A. baumannii biofilms. Importantly, encapsulation of Ab-Cath in nanogels reduced Ab-Cath's cytotoxic effects on human fibroblasts by ≥10-fold. Moreover, cutaneous application of Ab-Cath nanogels eliminated bacteria colonizing 3D human skin. These findings affirm the use of nanogels to increase the selectivity index of antimicrobial peptides.

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Keywords: Antimicrobial peptide Ab-Cath; AMR infection; Nanogel; Lyophilization method; Drug delivery

Background

Chronic wound infections are responsible for considerable morbidity and significantly contribute to the high cost of health care. The majority of these infections are caused by Staphylococcus aureus, however most often chronic wound infections are polymicrobial, where two or more species of Gram-positive and/or Gram-negative bacteria occupy the infection site.1,2 Usually, these infections are accompanied with formation of biofilms, i.e. complex structures of bacteria and their self-produced molecules, such as extracellular polymeric substances.3 Biofilm-residing bacteria are 10–1000 times more resistant to antibiotics compared to their planktonic counterparts.3,4 This, in combination with the rising numbers of antimicrobial resistance worldwide emphasizes the urgent need for novel broad-spectrum antimicrobial agents as alternatives to current antibiotics. Antimicrobial peptides (AMPs) may fulfil this role.

A candidate AMP is Ab-Cath, that belongs to the family of cathelicidins. This peptide's sequence was mined from the genome of the snake Anilos bituberculatus. Ab-Cath is an amphipathic peptide spanning 32 amino acids and is considerably cationic. The peptide is highly effective against antimicrobial resistant (AMR) bacteria and, in particular, against Gram-negative strains (Voet and Nibbering, personal communication).
Importantly, Ab-Cath maintains bactericidal effects against bacteria in plasma and whole blood, though its accompanying moderate cytotoxicity to mammalian cells may challenge its development into a therapeutic dosage form.

One strategy to improve the therapeutic potential of Ab-Cath is encapsulation of the peptide into a drug delivery system composed of biodegradable and biocompatible polymers, such as hyaluronic acid (HA), which has antiadhesive and antibiofilm properties towards bacteria. HA may be modified with octenyl succinic anhydride (OSA) to produce a polymer with amphiphilic properties, allowing the polymer to self-assemble into soft flexible nanogels composed of hydrophobic and hydrophilic zones within the nanogel matrix. Nanogels composed of OSA-HA reduced the cytotoxicity of a range of molecules, including AMPs, peptidomimetics, and antibiofilm peptides. These nanogel formulations are produced as aqueous dispersions by using microfluidic-assisted self-assembly, yet the nanogels and the encapsulated peptides typically display limited shelf-lives under this condition. Lyophilization of nanogels is commonly applied to improve drug formulation stability, long-term storage, product shelf-life, and to ease distribution. Development of a lyophilization method is therefore essential for translation of Ab-Cath nanogels to the clinic.

In this study, we aimed to formulate Ab-Cath in OSA-HA nanogels to reduce cytotoxicity while maintaining the antimicrobial activity of Ab-Cath, i.e. to increase the peptide’s selectivity index. Moreover, an effective lyophilization method was developed allowing for prolonged storage and concentration of the nanogels.

**Methods**

**Materials**

Hyaluronic acid (HyaCare, 50 kDa) was purchased from Evonik Nutrition and Care (Essen, Germany). Octenyl succinic anhydride (OSA, 97 %), lysozyme egg white, D-mannitol, trifluoroacetic acid (TFA), Triton X-100 and bovine serum albumin (BSA) were obtained from Sigma-Aldrich (St. Louis, MO, USA). Sodium bicarbonate (NaHCO₃), sodium hydroxide (NaOH), calcium chloride (CaCl₂), calcium chloride (CaCl₂) and human serum (male, type AB) were obtained from Merck (Darmstadt, Germany). Ab-Cath (acetyl-KRFKKFFRKVKKGVHRYFKKNKFYIAATIPYYG-amide; 4159.6 g/mol) was purchased from Avivia (Nijmegen, Netherlands). Tran-40 was a kind gift from Avivia (Nijmegen, Netherlands). Uranyl acetate (UA) was obtained from Fluka (Charlotte, NC, USA). Analytical grade solvents for UPLC analysis included ultrapure water (Veolia Purelab Chorus 1, High Wycombe, UK) and acetonitrile (100 %, VWR, PA, USA). Ultrapure water was obtained from a MilliPore system (Merck), and phosphate buffered saline (PBS) from Fresenius Kabi (Graz, Austria). Triptocoy broth (TSB), brain heart infusion (BHI) and Mueller-Hinton (MH) agar were purchased from Oxoid (Basingstoke, UK). Human plasma was obtained from Sanquin (Leiden, Netherlands). Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 1 % (v/v) GlutaMAX™, penicillin-streptomycin (pen/strep) and trypsin-EDTA were obtained from Gibco (Waltham, MA, USA). Microplates were purchased from Greiner BioOne (Alphen a/d Rijn, Netherlands) and culture plates and inactivated fetal bovine serum (FBSi) from Corning (NY, USA).

**Modification of hyaluronic acid with octenyl succinic anhydride**

Octenyl succinic anhydride-modified hyaluronic acid (OSA-HA, 17 %–32 % degree of substitution, see Figs. S2–S3) was synthesized as described by Eenschoten et al. Briefly, 1.25 g HA was dissolved into 50 mL ultrapure water and NaHCO₃ was added to yield a 2 M carbonate solution. Afterwards the pH was adjusted to pH 8.5 with 0.5 M NaOH and OSA was added dropwise to the HA solution to reach a 50:1 molar ratio of OSA:HA. The solution was left to react overnight at room temperature. Then, the reaction product was dialyzed against ultrapure water at 4 °C with water refreshment regularly until the conductivity reached a value lower than 5 μS/cm. Finally, the produced OSA-HA was lyophilized and characterized with 1H NMR.

**Preparation of Ab-Cath OSA-HA nanogels**

Ab-Cath OSA-HA nanogels were produced at room temperature using a microfluidic chip design previously described. Ab-Cath and OSA-HA were dissolved in ultrapure water to a concentration of 1500 μg/mL (10x final peptide concentration) and 500 μg/mL, respectively. The solutions were filled into three gastight fixed Luer lock tip glass syringes (Prosense, Oosterhout, Netherlands) mounted on 3 NE-300 syringe pumps (Prosense) to be able to control the flow rates. The OSA-HA solution was injected into the outer streams of the microfluidic chip at a flow rate of 0.99 mL/min and the peptide solution was injected into the middle stream at a flow rate of 0.22 mL/min, resulting a combined flow of 2.2 mL/min. The produced nanogels contained 150 μg/mL Ab-Cath and 500 μg/mL OSA-HA. For the experiments either freshly prepared nanogels or lyophilized nanogels were used; the concentration of peptide is given as the total peptide concentration present in the sample. Freshly prepared nanogels were stored at 4 °C and used within one week. For lyophilization of nanogels, cryoprotectant solutions containing 1–50 mg/mL D-mannitol, dextran-40 or PVA were added in a 1:1 (v/v) ratio to the nanogels, and the nanogel solution was lyophilized overnight using an Alpha 1–4 LSCbasic freeze dryer (Martin Christ Gefriertrocknungsanlagen, Osterode am Harz, Germany) with RV3 vacuum pump and EMF10 oil mist filter (Edwards, Burgers Hill, UK). Freshly prepared nanogels were diluted, and lyophilized nanogels resuspended, into the relevant media before use.

**Size, polydispersity index and zeta potential of the nanogels**

Dynamic light scattering was used to determine the average nanogel size and polydispersity index (PDI), while the surface charge of the nanogels was estimated by the zeta potential (ZP). The size, PDI and ZP measurements were performed in ultrapure water at 25 °C using a Zetasizer Nano ZS (Malvern Instruments, Worcestershire, UK) equipped with a 633 nm laser and 173°
Transmission electron microscopy

Nanogels were visualized using negative stain transmission electron microscopy (TEM). Briefly, 200 mesh formvar and carbon coated copper EM grids (Agar Scientific, Stansted, UK) were glow-discharged by 0.2 mbar air for 1 min using the glow discharge unit of an EMITECH K950X (Quorum Technologies, Lewes, UK). Then, 3 μL of nanogel (containing 150 μg/mL Ab-Cath and 500 μg/mL OSA-HA) was applied per glow-discharged grid for 1 min and the grids were blotted to remove excess of sample. Subsequently, the grids were stained on droplets of 2 % (w/v) UA in water for 1 min, after which excess of the staining solution was removed with blotting paper. Imaging of the air-dried grids was performed at 120 kV on a Tecnai 12 electron microscope (ThermoFisher, Waltham, MA, USA). A 4k × 4k Eagle camera (ThermoFisher) was used to record images at 11,000× magnification.

Quantification of Ab-Cath

Quantification of Ab-Cath was performed using an ACQUITY H-class UPLC-MS system (Waters, Milford, MA, USA) with an LCT-premier mass detector (Waters). Chromatographic separation was carried out on an ACQUITY UPLC BEH C18 column (100 × 2.1 mm, 1.7 μm; Waters). The mobile phase consisted of eluent A (100 % water) and eluent B (100 % acetonitrile), both containing 0.05 % (v/v) TFA. Samples were run with a gradient of 5 to 75 % eluent B over 8 min at 0.5 mL/min at 50 °C. The data were analyzed using MassLynx Software V4.2 and a calibration curve of Ab-Cath (0.01–0.1 mg/mL).

Encapsulation of Ab-Cath into nanogels

The Ab-Cath amount encapsulated in nanogels was determined indirectly by measuring the residual amount of peptide present in the aqueous bulk phase after nanogel production. The aqueous bulk phase was obtained by centrifuging the nanogels at 100,000 × g; Waters). The mobile phase was 100% aqueous bulk phase was obtained by centrifuging the nanogels at 500,000 × g for 30 min. The encapsulation efficiency (EE) was quantified for 3 independent sample batch replicates. Calculations of the EE are based on the theoretical amount of Ab-Cath encapsulated, as only non-encapsulated Ab-Cath was measured:

\[
EE(\%) = \frac{\text{Total Ab-Cath} (\mu g) - \text{unencapsulated Ab-Cath} (\mu g)}{\text{Total Ab-Cath} (\mu g)} \times 100\%
\]

Similarly, the drug loading (DL) was calculated:

\[
DL(\%) = \frac{\text{Encapsulated Ab-Cath} (\mu g)}{\text{Total weight nanogels} (\mu g)} \times 100\%
\]

Release of Ab-Cath from nanogels

In vitro release studies with lyophilized Ab-Cath nanogels were performed in PBS using dialysis membranes (Spectra-Port® Float-a-Lyzer® G2, MWCO 100 kDa, Spectrum Labs, Breda, Netherlands). Prior to use, dialysis membranes were washed according to manufacturer's protocol. Next, dialysis membranes were coated with 1 mg/mL lysozyme egg white for 1 h at 37 °C and 200 rpm using an orbital shaker. Dialysis membranes were then washed with ultrapure water and loaded with 1 mL of Ab-Cath or Ab-Cath nanogels (at 300 μg/mL) and placed in 6 mL of release buffer while continuously shaking at 200 rpm. Temperature was maintained at 37 °C throughout the experiment and 1 mL samples were taken until 5 h, replacing sample volumes with equal volumes of PBS. From 5 h onwards, 6 mL samples were taken to maintain sink conditions. Samples were stored at -20 °C until analysis by UPLC. Results are expressed as percentage of Ab-Cath released from the nanogels compared to Ab-Cath solution.

Bacterial strains and culturing

Antimicrobial-resistant (AMR) strains used in this study include Acinetobacter baumannii strain (RUH875), methicillin-resistant Staphylococcus aureus (MRSA) strain LUH14616 (NCCB100829) and Escherichia coli strain LUH15174 (SPA012, invasive strain). Bacteria were stored in glycerol at −80 °C until use. In short, bacteria were cultured on blood agar plates overnight at 37 °C. Next, 3–5 colonies were cultured to mid-log phase in TSB for 2.5 h under continuous rotation. Mid-log phase bacteria were centrifuged at 1000 × g for 10 min, washed with PBS and resuspended in the preferred medium to the required concentrations based on the optical density at 600 nm.

In vitro killing assay

Mid-log phase bacteria were resuspended in PBS to a concentration of 5 × 10^6 colony forming units (CFU)/mL. Subsequently, 30 μL of PBS containing increasing concentrations of Ab-Cath, Ab-Cath nanogels or placebo nanogels, 50 μL of pooled human plasma (Sanquin) or human urine (healthy volunteers) and 20 μL of the bacterial suspension were pipetted into wells of a polypropylene V-shaped microplate. After incubation for 4 h at 37 °C under rotation at 200 rpm, the number of viable bacteria was assessed microbiologically. Results are expressed as lethal concentration (LC)_{99.9}, i.e. the lowest concentration of Ab-Cath killing 99.9 % of the inoculum.

In vitro biofilm breakdown assay

Mid-log phase bacteria were diluted in BHI to grow 24 h biofilms. Briefly, 100 μL of mid-log phase bacteria at 1 × 10^7 CFU/mL was added to each well of a polypropylene flat-bottom microplate and incubated for 24 h at 37 °C in humidified environment. The next day, planktonic bacteria were removed from the wells and biofilms were washed twice with 100 μL of PBS to remove remaining non-adherent bacteria. Biofilms were subsequently exposed to increasing concentrations of Ab-Cath, Ab-Cath nanogels, or placebo nanogels in PBS for 4 h at 37 °C. Plates were sealed with AmpliStar Adhesive Plate Seals (Westburg, Leusden, Netherlands). Medium controls were checked for possible contamination. Finally, biofilms were washed twice with PBS and biofilm-residing bacteria were harvested by detection optics. Malvern DTS v.6.20 software was used for data acquisition and analysis. Measurements were performed for 3 independent sample batch replicates.
sonication (Branson 1800 sonicator, 10 min) in 100 μL of PBS. The number of viable bacteria was assessed microbiologically. Results are expressed as the biofilm eradication concentration (BEC)_{99.9}, i.e. the lowest concentration of Ab-Cath killing 99.9 % of the bacteria in the biofilm.

**In vitro cytotoxicity assay using human primary skin fibroblasts**

Human primary skin fibroblasts (kindly provided by Dr. A. El Ghalbzouri, Department of Dermatology, LUMC) were grown in culture flasks using DMEM supplemented with 1 % (v/v) GlutaMAX™, 1 % (v/v) pen/strep and 5 % (v/v) FBSi. Fibroblasts were harvested using 0.05 % (v/v) trypsin-EDTA, washed and resuspended to 2 × 10⁵ cells/mL and finally 20,000 cells were seeded in 96-wells flat-bottom culture plates. Monolayers developed overnight at 37 °C and 5 % CO₂. After inoculation, the bacterial suspension was removed. HSEs were inoculated by the Cytotoxicity Detection Kit (Roche, Basel, Switzerland) according to manufacturer's instructions. Results are expressed as effective concentration (EC)_{50}, i.e. the highest concentration of Ab-Cath inducing 50 % reduction of cytotoxicity or metabolic activity. Non-linear regression curves with bottom and top restrictions at 0 and 100 % were fitted for each individual experiment to determine medians (and ranges) of the EC_{50} values.

**Calculation of the selectivity index**

The selectivity index, i.e. ratio between cytotoxicity and bactericidal activity, was calculated as follows:

\[
\text{Selectivity index} = \frac{\text{Cytotoxicity (μM)}}{\text{Bactericidal activity (μM)}}
\]

**3D human skin equivalents**

Human skin equivalents (HSEs) were cultured using Ker-CT cells (ATCC® CRL-4048™) as described previously. At least 2 days before infection, the culture medium was replaced by culture medium without antibiotics. The HSEs were inoculated with AMR _S. aureus_ LUH14616 or A. baumannii RUH875 at a concentration of 1 × 10⁵ CFU/model for 1 h at 37 °C and 5 % CO₂. After inoculation, the bacterial suspension was removed, the cells were washed with PBS and the colonized models were treated with Ab-Cath, lyophilized and dispersed Ab-Cath nanogels or placebo nanogels at the desired concentrations in PBS for 4 h after which the supernatants (non-adherent bacteria) were stored on ice, while the models (adherent bacteria) were homogenized (5000 rpm, 3 × 10 s, 10 s pause) using a bead-beater (Precellys 24 lysis and homogenization, Bertin Technologies). Bacterial counts in both fractions were assessed microbiologically. Results are expressed as individual values and medians of 3 individual measurements performed in duplicate.

**In vitro cytotoxicity towards 3D skin equivalents**

HSEs, cultured as described earlier, were exposed to Ab-Cath, lyophilized and dispersed Ab-Cath nanogels or placebo nanogels at the desired concentrations in PBS for 4 h including 1 % (v/v) Triton-X100 as positive control and PBS as negative control. Afterwards, LDH release from dead cells into the supernatants was detected by the Cytotoxicity Detection Kit (Roche, Basel, Switzerland) according to manufacturer’s instructions. Models were cut out, transferred to 24 well flat-bottom culture plates (Corning, NY, USA) and exposed to WST-1 reagent (Cell proliferation reagent WST-1; Roche) in Dulbecco’s Modified Eagle’s Medium (DMEM) medium (Gibco, Waltham, MA, USA) overnight to determine the metabolic activity of the cells in the models. Read-out of medium solutions without the models was performed according to manufacturer’s protocol. Results are expressed as percentage cytotoxicity or metabolic activity relative to the controls.

**Statistics**

Differences between 2 groups (Ab-Cath, Ab-Cath nanogels or placebo nanogels) were evaluated by a Kruskal-Wallis test, followed by a Mann-Whitney rank sum test using Graphpad Prism software version 6.0 (Graph Pad Software). Differences were considered statistically significant when p < 0.05.

**Informed consent statement**

All human primary skin fibroblasts used in this study were isolated from surplus skin collected according to article 467 of the Dutch Law on Medical Treatment Agreement and the Code for Use of Human Tissue of the Dutch Federation of Biomedical Scientific Societies. The Declaration of Helsinki principles were followed when working with human primary cells.

**Results**

**Physicochemical properties of freshly produced Ab-Cath nanogels**

Ab-Cath OSA-HA nanogels were prepared using increasing concentrations of Ab-Cath at a peptide to polymer ratio of 0.3:1 (w/w). Produced nanogels ranged in size from 195 nm to 465 nm; all being monodisperse, negatively charged, and with high encapsulation efficiency and drug loading (Table 1). Incorporation of increasing amounts of Ab-Cath resulted in less negatively charged nanogels and increased nanogel sizes. OSA-HA nanogels containing 150 μg/mL Ab-Cath were chosen for further evaluation, considering a preferred particle size between 150 and 200 nm.⁶,¹⁷

**Stability of freshly produced Ab-Cath nanogels during storage**

The physicochemical properties of OSA-HA nanogels were monitored over a 15-day period to determine the colloidal stability of the nanogels in ultrapure water stored at 4 °C. Results...
showed no significant (p > 0.05) differences in size, PDI and ZP for both Ab-Cath and placebo nanogels during storage (Fig. 1). Therefore, OSA-HA nanogels were considered physically stable over this period.

**Antimicrobial activity of freshly produced Ab-Cath nanogels**

Next, we compared the antimicrobial activity of Ab-Cath nanogels to Ab-Cath and, as control, placebo nanogels on AMR *S. aureus* and *A. baumannii* in 50 % (v/v) plasma and *E. coli* in urine. After 4 h, 24 h and 48 h of exposure, the bactericidal activity of Ab-Cath nanogels was similar to Ab-Cath (Table 2).

Placebo nanogels did not show any bactericidal activity at the tested concentrations.

**Physicochemical properties of lyophilized Ab-Cath nanogels**

Lyophilization was applied to allow further testing of more concentrated Ab-Cath nanogels, i.e. antibiofilm and cytotoxic activities. As this process could potentially destabilize the nanogels, the ability of various cryoprotectants to protect the nanogels during lyophilization was investigated. TEM microscopical examination of fresh Ab-Cath nanogels revealed individual spherical nanogels with sizes below 300 nm.

### Table 1

<table>
<thead>
<tr>
<th>Ab-Cath (μg/mL)</th>
<th>OSA-HA (μg/mL)</th>
<th>Size (nm)</th>
<th>PDI</th>
<th>ZP (mV)</th>
<th>EE (%)</th>
<th>DL (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>500</td>
<td>238 ± 25</td>
<td>0.15 ± 0.06</td>
<td>−32.6 ± 2.7</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>150</td>
<td>500</td>
<td>194 ± 38</td>
<td>0.06 ± 0.04</td>
<td>−16.5 ± 3.1</td>
<td>99.7 ± 0.1</td>
<td>23</td>
</tr>
<tr>
<td>200</td>
<td>666.7</td>
<td>367 ± 160</td>
<td>0.06 ± 0.04</td>
<td>−14.3 ± 6.4</td>
<td>99.7 ± 0.1</td>
<td>23</td>
</tr>
<tr>
<td>250</td>
<td>833.3</td>
<td>465 ± 163</td>
<td>0.06 ± 0.05</td>
<td>−11.4 ± 1.6</td>
<td>99.6 ± 0.1</td>
<td>23</td>
</tr>
</tbody>
</table>

Data are presented as mean ± SD of three to 24 independent sample batch replicates. OSA-HA = octenyl succinic anhydride-modified hyaluronic acid; PDI = polydispersity index; ZP = zeta potential; EE = encapsulation efficiency; DL = drug loading.

### Table 2

<table>
<thead>
<tr>
<th>Exposure</th>
<th>LC&lt;sub&gt;99.9&lt;/sub&gt; (μM)</th>
<th>Ab-Cath solution</th>
<th>Ab-Cath nanogel</th>
<th>Placebo nanogel</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>S. aureus</em> (LUH14616)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 h</td>
<td>1.6 (1.6–3.2)</td>
<td>1.6 (1.6–3.2)</td>
<td>&gt;12.8</td>
<td></td>
</tr>
<tr>
<td>24 h</td>
<td>0.8 (0.8–3.2)</td>
<td>0.8 (0.8–3.2)</td>
<td>&gt;12.8</td>
<td></td>
</tr>
<tr>
<td>48 h</td>
<td>1.6</td>
<td>3.2 (0.8–6.4)</td>
<td>&gt;12.8</td>
<td></td>
</tr>
<tr>
<td><em>A. baumannii</em> (RUH875)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 h</td>
<td>0.2 (0.1–0.4)</td>
<td>0.4 (0.2–0.4)</td>
<td>&gt;6.4</td>
<td></td>
</tr>
<tr>
<td>24 h</td>
<td>0.1</td>
<td>0.2 (0.05–0.2)</td>
<td>&gt;6.4</td>
<td></td>
</tr>
<tr>
<td>48 h</td>
<td>0.2</td>
<td>0.4 (0.2–0.4)</td>
<td>&gt;6.4</td>
<td></td>
</tr>
<tr>
<td><em>E. coli</em> (LUH15174)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 h</td>
<td>0.8 (0.4–1.6)</td>
<td>0.8 (0.4–1.6)</td>
<td>&gt;3.2</td>
<td></td>
</tr>
</tbody>
</table>

Bactericidal activity on AMR bacteria upon 4 h, 24 h and 48 h exposure to increasing concentrations of Ab-Cath, Ab-Cath nanogels and placebo nanogels in 50 % (v/v) plasma (*S. aureus* and *A. baumannii*) or urine (*E. coli*) was assessed. Results are expressed as median and ranges of 2–4 independent experiments performed in duplicate and is shown as LC<sub>99.9</sub>, i.e. the lowest concentration of peptide to kill 99.9 % of the bacteria.
(Fig. 2A), whereas collapsed nanogels and aggregates were seen after lyophilization without cryoprotectants (Fig. 2B). Lyophilization of Ab-Cath nanogels in presence of 10 mg/mL PVA (Fig. 2C) and 10 mg/mL dextran-40 (Fig. 2D), but not 50 mg/mL mannitol (Fig. 2E), maintained the morphology of the nanogels. Further investigations revealed no major differences in size, PDI and ZP between lyophilization without cryoprotectants and lyophilization in presence of PVA, dextran-40 or mannitol (Table 3). Use of mannitol was restricted due to its low solubility. Observed differences between the two methods are a direct result of different sample preparation, with TEM measurements performed on air dried samples and DLS on nanogel solutions. The use of both TEM and DLS thus provides complimentary data, allowing for a comprehensive understanding of the behavior of nanogels upon lyophilization with and without cryoprotectants. Together, these data indicate that both PVA and dextran-40 protect Ab-Cath nanogels during lyophilization.

Table 3
Physicochemical properties of lyophilized and redispersed Ab-Cath nanogels.

<table>
<thead>
<tr>
<th>Lyophilized</th>
<th>Cryoprotectant</th>
<th>Cryoprotectant (mg/mL)</th>
<th>Size (nm)</th>
<th>PDI</th>
<th>ZP (mV)</th>
<th>EE (%)</th>
<th>DL (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No</td>
<td>–</td>
<td>–</td>
<td>160 ± 7</td>
<td>0.08 ± 0.03</td>
<td>−19.7 ± 2.3</td>
<td>100.0 ± 0.0</td>
<td>23.0 ± 0.0</td>
</tr>
<tr>
<td>Yes</td>
<td>–</td>
<td>–</td>
<td>280 ± 99</td>
<td>0.17 ± 0.08</td>
<td>−24.8 ± 2.3</td>
<td>98.9 ± 1.1</td>
<td>22.9 ± 0.2</td>
</tr>
<tr>
<td>Yes</td>
<td>Mannitol</td>
<td>50</td>
<td>205 ± 31</td>
<td>0.17 ± 0.02</td>
<td>−20.3 ± 3.1</td>
<td>96.8 ± 4.1</td>
<td>22.5 ± 0.7</td>
</tr>
<tr>
<td>Yes</td>
<td>PVA</td>
<td>10</td>
<td>261 ± 28</td>
<td>0.17 ± 0.03</td>
<td>−25.5 ± 0.8</td>
<td>84.4 ± 5.9</td>
<td>20.2 ± 1.1</td>
</tr>
<tr>
<td>Yes</td>
<td>Dextran-40</td>
<td>10</td>
<td>308 ± 71</td>
<td>0.24 ± 0.04</td>
<td>−23.0 ± 3.3</td>
<td>98.2 ± 1.6</td>
<td>22.8 ± 0.3</td>
</tr>
</tbody>
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Nanogels were produced using 150 μg/mL Ab-Cath and 500 μg/mL OSA-HA. Data are presented as mean ± SD of three to four independent sample batch replicates. Abbreviations: OSA-HA = octenyl succinic anhydride-modified hyaluronic acid; PDI = polydispersity index; ZP = zeta potential; EE = encapsulation efficiency; DL = drug loading; PVA = polyvinyl alcohol.
Release of Ab-Cath from lyophilized and redispersed nanogels

The levels of Ab-Cath released from OSA-HA nanogels redispersed in PBS after lyophilization in presence of PVA or dextran-40 gradually increased until maximum values amounted to 48 % and 36 % at 25 h, respectively (Fig. 3). Ab-Cath solution rapidly diffused through the dialysis membrane, reaching its maximum at around 25 h (Fig. S4).

Antimicrobial activities of lyophilized and redispersed Ab-Cath nanogels

The bactericidal activity of lyophilized and redispersed Ab-Cath nanogels against planktonic AMR S. aureus and A. baumannii in 50 % human plasma and against E. coli in 50 % urine was similar to that of Ab-Cath (Fig. 4A and Table S2). In addition, the antibiofilm activities of lyophilized Ab-Cath nanogels and Ab-Cath on AMR S. aureus and A. baumannii biofilms were also comparable (Fig. 4B and Table S2). Of note, placebo nanogels and cryoprotectant solutions did not show any antibacterial activity at the tested concentrations (data not shown). Together, Ab-Cath nanogels were similarly effective as Ab-Cath solution against AMR bacteria.

Decreased cytotoxicity to human primary skin fibroblasts of lyophilized and redispersed Ab-Cath nanogels

Next, the cytotoxic activities of lyophilized Ab-Cath nanogels redispersed in culture medium supplemented with 0.5 % (v/v) human serum were compared to that of Ab-Cath by exposing monolayers of human primary skin fibroblasts for 4 h to these solutions. The EC50 of Ab-Cath for human primary skin fibroblasts amounted to 49 μM, while the EC50 values for Ab-Cath nanogels lyophilized without cryoprotectants or in presence of PVA or dextran-40 were 175 μM, 540 μM and 630 μM, respectively (Fig. 4C). In agreement, Ab-Cath reduced the metabolic activity of these cells at considerably lower concentrations than Ab-Cath nanogels (Fig. 4D). Placebo nanogels did not induce cytotoxicity in skin fibroblasts (Fig. S5).

Improved selectivity index of lyophilized and redispersed Ab-Cath nanogels

Comparison of the selectivity indexes, i.e. ratio between cytotoxicity and bactericidal or antibiofilm activity, revealed that loading in nanogels improved the overall selectivity index of Ab-Cath (Fig. 4E, F and Table S3). In addition, the selectivity indexes of Ab-Cath nanogels lyophilized in the presence of cryoprotectants were >1.5-fold higher than those of Ab-Cath nanogels lyophilized without cryoprotection. These data indicate that nanogel encapsulation of Ab-Cath allows the use of higher Ab-Cath concentrations without inducing cytotoxicity compared to Ab-Cath in solution. Overall, Ab-Cath nanogels lyophilized in presence of dextran-40 showed the most promising selectivity index for both planktonic and biofilm-residing AMR bacterial strains and was selected for further testing.

Lyophilized and redispersed Ab-Cath nanogels maintained effectiveness against AMR infections in a 3D human epidermal model, while they reduced cytotoxicity

Lastly, the efficacy of redispersed Ab-Cath nanogels lyophilized in the presence of dextran-40 and Ab-Cath were assessed against AMR S. aureus and A. baumannii colonizing a 3D human epidermal model. Results revealed a dose-dependent reduction in S. aureus and A. baumannii counts on the epidermal model upon exposure to Ab-Cath nanogels as well as to Ab-Cath solution (Fig. 5A, B). Exposure of the bacteria on the epidermal model to placebo nanogels was without effect on bacterial counts. Furthermore, the highest dose of Ab-Cath induced minor cytotoxicity towards the epidermal model, which was significantly higher (p = 0.0288) than Ab-Cath nanogels in the supernatant (Fig. 5C), but this difference was not significant in the subnatant (Fig. 5D).

Discussion

The present study demonstrates that the snake cathelicidin Ab-Cath can be efficiently encapsulated in OSA-HA nanogels through lyophilization and subsequent redispersion. This approach improved the selectivity index of Ab-Cath, allowing higher concentrations to be used without inducing cytotoxicity in comparison to Ab-Cath in solution. Furthermore, Ab-Cath nanogels showed promise in reducing bacterial counts in a 3D human epidermal model, and their cytotoxicity was significantly lower than that of Ab-Cath solution. This work paves the way for further development of Ab-Cath nanogels as potential therapeutic agents for the treatment of AMR infections.
with favorable physicochemical properties. This conclusion is based on the following findings. First, we produced nanogels encapsulating 150 μg/mL Ab-Cath ranging from 156 to 232 nm in size with very homogenous distribution, as evidenced by the low polydispersity index (PDI = 0.06) and by TEM. These results support previous observations that nanogel preparation by means of microfluidic-assisted self-assembly allows for a precise control over the self-assembly process of nanogels. Moreover, these nanogel sizes met our intended size range of 150–200 nm, which was based on previous observations that transport across biofilms is negatively correlated to nanogels with increasing size and nanogels in this size range are less likely to be removed by immune cells and the complement system. Second, Ab-Cath nanogels showed an overall reduction in the surface electro-negative net charge (−16.5 mV) compared to non-loaded nanogels (−32.6 mV), suggesting that in addition to being...
encapsulated in the nanogel this cationic peptide also binds to the nanogel's surface. It is hypothesized that both hydrophobic and electrostatic interactions between polymer and peptide are important for the nanogel surface binding by Ab-Cath. Third, the encapsulation efficiency of Ab-Cath in OSA-HA nanogels was very high (≥99 %) in comparison to that of the antibiofilm peptide DJK-5-loaded and peptoid-loaded OSA-HA nanogels.10,11 This is likely related to intrinsic properties of the AMP, such as length,19 hydrophobicity20 and charge.21 In addition, loading of Ab-Cath into nanogels amounted to 23 %, which is substantially higher than the <10 % drug loading achieved for most existing nanomedicines.22 This finding further emphasizes the potential of OSA-HA nanogels as high peptide drug-loading nanomedicine. Fourth, both Ab-Cath and non-loaded OSA-HA nanogels were physically stable over 15 days when stored at 4 °C in ultrapure water. This is in agreement with previous observations that nanogel dispersions with a ZP of 10–30 mV above or below zero are considered relatively to highly stable.23 Finally, we found that 26–33 % of Ab-Cath was released from the lyophilized nanogels upon redispersion in the first 5 h followed by a sustained release phase over at least 72 h. This release is slower than the release of DJK-5 from OSA-HA nanogels in 24 h,11 but similar to the release of novicidin from OSA-HA nanogels in 14 days,9 indicating that the release profile is peptide-specific. Sustained release is favorable in treatment of complex wound and burn infections, as it minimizes the frequency of painful wound dressing changes.24 The release of Ab-Cath from OSA-HA nanogels stabilizes after 72 h, indicating a slow sustained release phase and/or strong electrostatic interactions between cationic Ab-Cath and anionic OSA-HA polymer. The release rate is influenced by i) pH, ii) presence of degrading enzymes and iii) salt concentration.7 Therefore, these release percentages should not be directly translated to concentrations used in the in vitro studies, where nanogels are used at varying conditions.

Lyophilization was used to concentrate Ab-Cath nanogels and for long-term storage, which are both essential for further development of these nanogels as therapeutics. However, this study showed that lyophilization without a cryoprotectant damaged the nanogels, affected their size and ZP, and resulted in poor antibacterial activities of Ab-Cath nanogels. Potential factors contributing to destabilization of the nanogels during lyophilization are related to the stress of freezing, dehydration and rehydration processes. We found that addition of cryoprotectants PVA or dextran-40 prior to lyophilization proved to fully abolish these drawbacks when lyophilizing the nanogels. Effective cryoprotectant concentrations of 10 mg/mL (=5 % w/v) used in this study corroborate with concentrations used in previous studies: ≥5 % (w/v) of mannitol,25 1 % (w/v) dextran-70,26 ≥5 % (w/v) dextran-9025 and ≥5 % (w/v) of PVA.27 To the authors' knowledge, this is the first report on successful lyophilization of AMP-loaded OSA-HA nanogels.

Importantly, this study confirms that encapsulation of Ab-Cath in OSA-HA nanogels is a successful strategy to reduce cytotoxicity and maintain antimicrobial activity of Ab-Cath, thus improving its selectivity index. This conclusion is based on the following findings. Encapsulation of Ab-Cath in OSA-HA nanogels lyophilized in presence of cryoprotectants PVA or dextran-40 maintained antimicrobial activity against planktonic AMR S. aureus, A. baumannii and E. coli and biofilm-residing AMR S. aureus and A. baumannii, while cytotoxicity for human primary skin fibroblasts was reduced 11- to 13-fold. The maintained activities combined with reduced cytotoxicity of Ab-Cath nanogels could potentially be attributed to i) a reduction in Ab-Cath's hydrophobicity upon encapsulation in nanogels, further enhancing its selectivity towards bacterial membranes and/or ii) desensitization of the human fibroblasts to increasing Ab-Cath concentrations due to the slow release of Ab-Cath from nanogels. Moreover, lyophilized Ab-Cath nanogels were as effective upon redispersion as Ab-Cath in eliminating AMR S. aureus and A. baumannii colonizing a 3D human epidermal model, while minor cytotoxicity of Ab-Cath in this epidermal model was not found for the formulated peptide. Future efficacy testing of Ab-Cath nanogels and peptide solution should include ex vivo skin and/or animal models. Nevertheless, based on this study it can be concluded that Ab-Cath is an excellent antimicrobial to treat AMR bacteria, either planktonic, biofilm-residing or hiding in the wounded skin.

In summary, Ab-Cath was efficiently encapsulated in OSA-HA nanogels with excellent physicochemical and functional properties. In addition, a successful method to lyophilize Ab-Cath nanogels was developed, allowing for long-term storage and concentration of the nanogels. Furthermore, the Ab-Cath nanogels were successful in eliminating bacteria when cutaneously applied on bacterial colonized 3D human skin. Together, OSA-HA nanogels are promising as delivery system for the snake cathelicidin Ab-Cath to combat AMR bacteria.

CRediT authorship contribution statement


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Declaration of competing interest

The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision to publish the results.
Appendix A. Supplementary data

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References


