

PAPER

View Article Online
View Journal | View Issue

Cite this: *Biomater. Sci.*, 2025, **13**, 3915

Enhancing hand hygiene compliance through the long-lasting antimicrobial effects of nitric oxide-releasing hand sanitizer gel†

Manjyot Kaur Chug, Gabrielle Aluisio, Cole Bousquet, Mark Garren,  Yun Qian,  Joseph H. Campbell  and Elizabeth J. Brisbois *

Effective hand hygiene is crucial for reducing the transmission of disease-causing pathogens. While alcohol-based hand sanitizers have become popular, their increased usage during the COVID-19 pandemic raised concerns about their short-lived activity and potential side effects. The increased application of hand sanitizers and harmful side effects has necessitated an effective alternative with prolonged and enhanced antimicrobial properties which could result in a reduced number of sanitizer applications. To address these issues and improve antimicrobial efficacy, this study developed a nitric oxide (NO)-releasing hand sanitizer (NORel) gel enriched with other antimicrobial and moisturizing ingredients like ethanol, tea tree oil, and glycerin. The NORel gel underwent comprehensive analysis, including assessments of pH for 60 d, rheology, NO release, cytocompatibility, and *in vitro* and *ex vivo* antimicrobial effectiveness on rabbit skin proving its ability to eliminate over 97% of bacteria and fungi, including antibiotic-resistant strains. One NORel gel formulation, NORel2, demonstrated antimicrobial activity comparable to a commercial alcohol-based gel containing 62% ethyl alcohol, achieving a reduction of more than 5 logs in *S. aureus* bacteria on a rabbit skin model. Additionally, the NORel gel significantly outperformed the commercial alcohol gel by maintaining its antimicrobial efficacy on infected rabbit skin, showing a persistent activity with a 1.6-log reduction in viable *S. aureus* 2 h after application. This research introduces a biocompatible NO-releasing gel with superior antimicrobial properties compared to common alcohol-based sanitizers, making it an effective hand hygiene solution to reduce infections, especially in high-risk environments.

Received 6th March 2025,
Accepted 26th May 2025
DOI: 10.1039/d5bm00359h
rsc.li/biomaterials-science

1. Introduction

Infectious diseases have become an enduring challenge, affecting individuals and communities worldwide. The burden of infectious diseases encompasses strains on healthcare infrastructure, economic losses, and disruption of daily life. In hospitals, healthcare-associated infections (HCAIs) occur when the patient does not present with infection upon admittance. According to the US Centers for Disease Control and Prevention (CDC), approximately 1.7 million hospitalized patients each year contract HCAIs while receiving medical treatment. Alarming, over 98 000 of these patients, equating to one in 17 cases, succumb to their HCAIs.¹ These infections can lead to severe bodily infections including bloodstream, urinary tract, surgical site, and ventilator-associated pneumonia, which prolong the duration of hospital stays, increase

morbidity and mortality, and may necessitate additional diagnostic and therapeutic interventions.^{1,2}

Infectious microorganisms have multiple pathways by which an individual can encounter them. These pathways can be broadly categorized as direct or indirect (Fig. 1A). Direct sources involve close contact with infected individuals, and transmission can occur through airborne means, or through direct physical contact with



Fig. 1 (A) Various sources of hand contamination. (B) Structural composition of bacteria (C) a detailed, step-by-step guide to the methodology for creating NO-releasing hand sanitizer (NORel) gel.

play a pivotal role in preventing HCAs.^{3–5} When executed correctly and consistently, hand hygiene significantly reduces the risk of healthcare-associated infections, community-acquired infections, the spread of contagious diseases, and healthcare costs associated with treating HCAs. The World Health Organization (WHO) advocates that effective hand hygiene is the single most important practice to prevent and control HCAs.^{1,6,7}

Hand hygiene can be achieved through handwashing with soap and water or using hand sanitizers. While handwashing remains the gold standard, it is not always feasible in various settings, such as healthcare facilities, public transportation, or during emergencies. In such situations, hand sanitizers are a convenient and effective alternative due to their portability and ease of use. Conventional alcohol-based hand sanitizers primar-

ily rely on ethanol or isopropyl alcohol, typically in concentrations ranging from 60% to 95%, as their only active ingredient. These sanitizers have proven highly effective against a broad spectrum of microorganisms due to the ability of ethanol to infiltrate microbial membranes (Fig. 1B), denature proteins and impede microbial growth.^{8,9} However, their efficacy depends on several factors, including the type and concentration of active ingredients, application technique, and contact time. Additionally, their rapid evaporation upon application limits their residual activity, necessitating frequent reapplication to maintain protection. Frequent use of hand sanitizers with alcohol has been shown to cause skin dryness, irritation, acute toxicity, and other dermatological side effects.¹⁰

This rapid vaporization is facilitated by the heat from the user's skin. Once all of the alcohol has vaporized, the skin's

surface is largely devoid of pathogens. However, this sanitized surface is vulnerable to rapid recolonization by airborne or surface-dwelling microorganisms, which can breed in the nutrient-rich environment that results from cells killed by the sanitizer. As a result, there exists a brief sanitization window, typically lasting only 1 to 2 min after the initial application, during which the sanitization effect remains effective.¹¹ Beyond this window, the initial application loses its relevance in maintaining a pathogen-free surface.¹² In situations requiring prolonged protection, such as in hospital-based settings, the limited persistent activity of hand sanitizers poses challenges.

Researchers have explored various natural and synthetic antimicrobial ingredients in combination with alcohol to mitigate the limitations observed in commercial hand sanitizers. Emerging hand sanitizer technologies incorporate essential oils, aloe vera, benzalkonium chloride, *etc.* to prevent skin dehydration and help stabilize the product to increase its biocidal activity.^{13–15} One technology gaining traction is the use of hydrogels in hand sanitizers. Hydrogels possess many desirable qualities for hand sanitization, including pleasant hand-feel, ease of delivery, and reduced risk of spillage due to higher viscosity than liquid preparations.¹⁶ However, it is important to note that many of these studies primarily aim to improve the antimicrobial effects and reduce the toxicity of hand sanitizer gels. To the best of our knowledge, there is no existing report that not only strives to meet antimicrobial standards comparable to alcohol-based gels but also emphasizes the long-term antimicrobial action of these sanitizer gels. An ideal hand sanitizer should maintain its efficacy for extended periods, reducing the need for frequent reapplication and lowering the risk of infection from various pathogens.

Nitric oxide (NO) is a natural gasotransmitter in the human body with diverse biological functions. It plays a crucial role in the body's defense against pathogens and contributes to various physiological processes, making it a promising candidate for infection control, wound healing, vasodilation, and other therapeutic applications.^{17–19} The gaseous nature and short half-life of NO allows it to rapidly penetrate a wide range of microbial species, including bacteria, viruses, and fungi, regardless of their structural composition, without inducing resistance.²⁰ Given NO's highly reactive and unstable nature in physiological conditions, researchers have focused on designing stable NO donors like *S*-nitroso-*N*-acetylpenicillamine (SNAP) and *S*-nitrosoglutathione (GSNO) to emulate biological functions of NO externally for prolonged periods.^{21–23} These donors can be readily incorporated into various substrates (*e.g.*, polymers, gels, wound dressings) to achieve the therapeutic benefits of NO, which also include wound healing, blood flow regulation, and preventing blood clotting.^{24,25} While some NO-releasing technologies have made strides in the fight against infections, the opportunity to apply these principles to hand sanitizer products is an exciting frontier that remains largely unexplored.

Inspired by the benefits of NO, this study developed a no-rinse NO-releasing hand sanitizer gel (NORel) designed to

provide enduring antimicrobial advantages to combat HCAs. The NORel gel is comprised of NO donor SNAP in addition to ethanol, water, and carbomer, enriched with tea tree oil and glycerin for enhanced antimicrobial properties and skin protection (Fig. 1C). Following the preparation of NORel hand sanitizer gel, the comprehensive assessment of all prepared formulations was conducted in terms of pH measurement, rheological behavior, NO release kinetics, and gel stability. The killing efficiency of NORel gels was performed against different microorganisms *S. aureus*, *E. coli*, a clinically isolated antibiotic-resistant strain of Methicillin-resistant *S. aureus*, and *C. albicans* yeast. In a more challenging study, the persistence activity of NORel gel was evaluated using an explanted rabbit skin model and compared to a commercially available alcohol-based hand sanitizer gel. Finally, the biocompatibility of gels was conducted against 3T3 mouse fibroblast cells to assess the safety of the prepared NORel gel. This study is the first to report the combination of NO with conventional hand sanitizer ingredients designed for topical application. The NORel gel is anticipated to be a valuable addition to clinical settings and hospitals, where the risk of infection transmission is high. The development of NORel is expected to provide prolonged protection, surpassing the efficacy of conventional alcohol-based hand sanitizers in terms of antim

2.2. Synthesis of NO donor *S*-nitroso-*N*-acetylpenicillamine (SNAP)

The protocol for synthesizing the NO donor *S*-nitroso-*N*-acetylpenicillamine (SNAP) was adapted from a previous publication with slight modifications.²² Briefly, the precursor NAP was dissolved in a mixture of two parts water and three parts methanol. Next, 0.7 M of H₂SO₄ and 1.6 M of HCl were added to the solution, and sodium nitrite, pre-dissolved in water, was added drop by drop. The mixture was stirred for 10 min at room temperature (RT, ~23 °C). To facilitate the formation of SNAP, the mixture was then crystallized on ice under continuous nitrogen purging for 8 h, all while being shielded from ambient light. After the 8 h incubation period, a filtration setup was prepared using Whatman cellulose filter paper in a Buchner funnel connected to a vacuum suction. The SNAP crystals were collected on the filter paper through suction filtration. Following collection, the SNAP crystals were rinsed with ice-cold deionized water and left in a vacuum desiccator overnight for drying. Care was taken to protect the samples from exposure to light throughout the entire process. The purity of the synthesized SNAP crystals, >90%, was confirmed using chemiluminescence NOA and a UV-vis calibration curve, with SNAP exhibiting a characteristic absorption band at 340 and 590 nm.

2.3. Preparation of NO-releasing (NORel) hand sanitizer gel

Hand sanitizer gels were prepared by initially dissolving carborer (105 mg) at room temperature (RT) into rapidly agitated ethanol (12 mL). Subsequently, DI water (7.12 mL) was added, and the mixture was stirred for 15 min at 200 rpm to achieve homogenous dispersion. After ensuring homogeneity, tea tree oil (40 µL) and glycerin (600 µL) were introduced into the blend. To include NO-releasing properties, *S*-nitroso-*N*-acetylpenicillamine (SNAP), a tertiary *S*-nitrosothiol (RSNO) NO donor, was incorporated into the formulations. To demonstrate the gel's adaptability, two distinct concentrations of SNAP were integrated into it. NORel1 and NORel2 hand sanitizer gels were formulated by adding 10.5 mg (10 wt%) and 31.5 mg (30 wt%) of SNAP, respectively, followed by additional stirring until SNAP dissolved fully and the liquid mixture became a uniform green color. Finally, triethylamine (TEA, 139.4 µL) was included for its coagulation and neutralizing properties, which transformed the liquid mixture into a viscous gel. A corresponding control without SNAP was prepared as a negative control for the study and labeled as the control gel. The resulting gels were transferred into 20 mL glass vials with caps for further characterization.

2.4. Characterization of NORel gel

2.4.1. pH measurement of gel. The initial pH assessment of NORel gels was conducted to assess the gel's safety. The pH of NORel1 and NORel2 gels was determined using a SevenCompact pH/Ion meter S220-std-kit (Mettler Toledo, Columbus, OH) with the probe directly immersed into each vial. Each sample was given a 10 min equilibration period before testing to reach room temperature and ensure precise

pH measurements. The pH probe was rinsed with DI water after each measurement. The final results are presented as the mean ± SD for each formulation ($n = 3$).

2.4.2. pH stability of gel. The pH stability of the gel was determined by tracking pH changes in hand sanitizer samples over 60 d. NORel gels were stored at various temperatures (−20 °C, 4 °C, room temperature (23 °C), and 37 °C) ($n = 3$) and pH was recorded on days 0, 5, 7, 14, 21, 28, 35, 42, 49, and 60 following the same method as 2.4.1. If the congealing agent were to

served as a reference to translate the absorbance readings from the study into compound concentrations within the samples. To ensure accuracy, the buffer solution's volume remained consistent across all samples throughout the experiment, preventing any variations in readings. The stability of the gels was assessed on days 0, 5, 7, 14, 21, and 28 d and compared to results obtained on day 0. Results from the study are reported as % NO remaining on each testing day relative to the initial amount of NO present in gels on day 0.

2.6. *In vitro* cytocompatibility screening

2.6.1. Preparation of 3T3 fibroblast cells. Mouse fibroblasts (NIH/3T3 cells, ATCC CRL-2522) were revived from cryopreserved stocks stored in the liquid nitrogen vapor phase. 3T3 cells were subcultured in minimal essential media (MEM) supplemented with 10% fetal bovine serum (FBS) and 1% streptomycin–penicillin (S–P) under a 5% CO₂-humidified atmosphere. Cells were grown for up to ten passages before discarding. Cells were grown up to 70% sub-confluency. Afterwards, cells were washed with 1× PBS (Ca²⁺ and Mg²⁺-free) and treated with 0.25% trypsin for 5 min. After detachment, cells were collected *via* centrifugation (200 rcf, 5 min) and resuspended in complete media. For cytotoxicity experiments, 12 mm hydrophilic polytetrafluoroethylene inserts (Millicell PICM01250) were pretreated with type I rat-tail collagen (~300 µg per insert) and then seeded with 3T3 cells in the intraluminal space to an initial density of 40 000 cells per cm² in 24-well plates. The extraluminal space was supplemented with 600 µL of complete media with 400 µL aliquoted in the intraluminal side. Cells were grown for 24 h prior to treatment.

2.6.2. Exposure of cells to NOrel gels. The aliquots of the gels (~100 mg) were added to the extraluminal side of the inserts with clean media added to both sides. Untreated wells (hereafter referred to as Blank) were also prepared with clean media. Cells were incubated for an additional 24 h. Afterward, the media was replaced with media supplemented with 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) to a final concentration of 0.5 mg mL⁻¹. Cells were incubated for an additional 3 h to facilitate the formation of the formazan precipitate. The supernatant was decanted taking caution to not perturb the crystalline layer. Afterward, the formazan precipitate was dissolved in dimethyl sulfoxide (DMSO, 350 µL per insert) and quantified for absorbance at 570 nm with a reference wavelength of 630 nm. Relative cellular viability was calculated according to eqn (1).

$$\text{Relative cell viability(\%)} = \frac{\text{Mean OD}_{\text{Treatment}}}{\text{Mean OD}_{\text{Blank}}} \times 100\% \quad (1)$$

Final data is reported as the mean percent viability ± SD for each formulation ($n = 4$ independent treatments per formulation).

2.7. *In vitro* antimicrobial activity of NOrel gel

2.7.1. Growing microbial culture. Antimicrobial activity of the NO

buffered saline. Samples were immediately processed if not stored at $-20\text{ }^{\circ}\text{C}$ until future experiments.

To evaluate the efficacy of NO-releasing (NOREl) hand sanitizer gels, the rabbit skin samples were taken out of the freezer at $-20\text{ }^{\circ}\text{C}$ and soaked in Deionized water to thaw the specimen. Once thawed, the samples were cut into fifteen 8 mm diameter circles using a biopsy punch and placed into a 24-well plate. To disinfect the specimens, a 1000 μL of diluted chlorhexidine (CHXD) disinfectant was added to each well. After 15 min, the CHXD was washed out by two 1000 μL aliquots of sterilized PBS. The samples were soaked in fresh PBS for 2–3 h to fully remove the effect of disinfectant. The samples were then removed and briefly rinsed again in fresh PBS and transferred into a new 24-well plate for testing.

2.8.2. Preparation of bacterial culture. *S. aureus* bacteria is commonly found on the skin and mucous membranes of humans and animals. While *S. aureus* is typically harmless on the skin, it can lead to infections if it enters the body through medical devices, breaks, or cuts in the skin or mucous membranes. It is particularly opportunistic and can cause infections under various conditions. Therefore, *S. aureus* was chosen as a model organism to test the *ex vivo* efficacy of NOrel hand sanitizer in preventing the spread of bacteria. The overnight culture of *S. aureus* was prepared following the same protocol as 2.7.1.

2.8.3. Disinfection efficacy of NOrel gel. To investigate the disinfection efficacy of NOrel gel, 500 μL of *S. aureus* bacteria (10^6 – 10^8 CFU mL^{-1}) was exposed to an 8 mm skin punch out and incubated at $37\text{ }^{\circ}\text{C}$, for 12 h. After incubation, bacteria suspension from the skin specimen was removed and the samples were washed with fresh, sterile PBS to remove free-floating/loosely adhered bacteria. Gels were weighed and $\sim 120\text{ mg}$ of gel was exposed to skin samples adhered with bacteria and left to act for 6 h at $37\text{ }^{\circ}\text{C}$. To compare the efficacy of NOrel in eradicating viable bacteria, *S. aureus* control (remained untreated) and a commercial alcohol-based gel were used as corresponding controls. Samples were gently rinsed with sterile PBS to remove loosely adhered or dead cells and resuspended into 1 mL of sterile PBS. To extract viable adhered bacteria after gel exposure, samples were homogenized and vortexed for 60 s each and plated using bacteria spiral plater (Eddy Jet 2, IUL Instruments) and colonies were enumerated using an automated colony counter (Sphere Flash, IUL Instruments). The % reduction in viable bacterial adhesion was determined using eqn (3) where C represents the concentration of viable bacteria in CFU cm^{-2} for treatment groups (commercial alcohol-based gel and NOrel2 gel) with respect to bacteria control that received no treatment.

$$\% \text{ reduction} = \frac{C(\text{Untreated}) - C(\text{Treated})}{C(\text{Untreated})} \times 100\% \quad (3)$$

2.8.4. Investigation of prolonged effectiveness of NOrel gel. To assess the prolonged efficacy of NOrel gel, the NOrel and commercial alcohol-based hand sanitizer gels were exposed to skin samples first and then exposed to bacteria.

This step was done to test the efficiency of

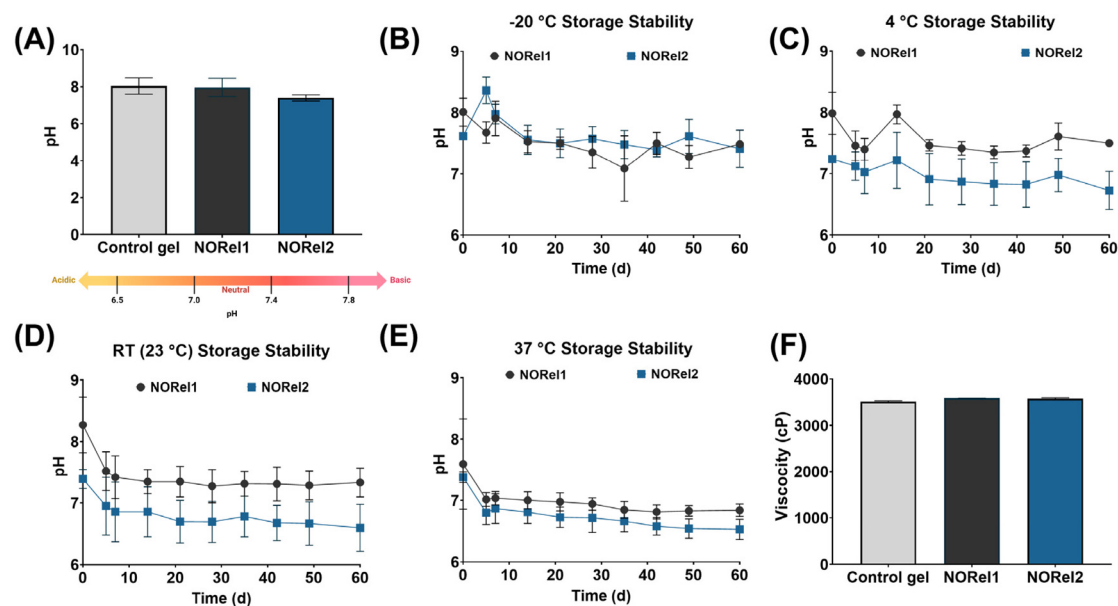


Fig. 2 (A) pH analysis of NOReL hand sanitizer gel. pH stability of NOReL gels at (B) $-20\text{ }^{\circ}\text{C}$ freezer (C) $4\text{ }^{\circ}\text{C}$ fridge (D) room temperature ($23\text{ }^{\circ}\text{C}$) and (E) $37\text{ }^{\circ}\text{C}$ (physiological temperature), up to 60 d of storage. (F) Average viscosity measurements of NOReL and control gels. Data represents mean $\pm$ SD, $n = 3$.

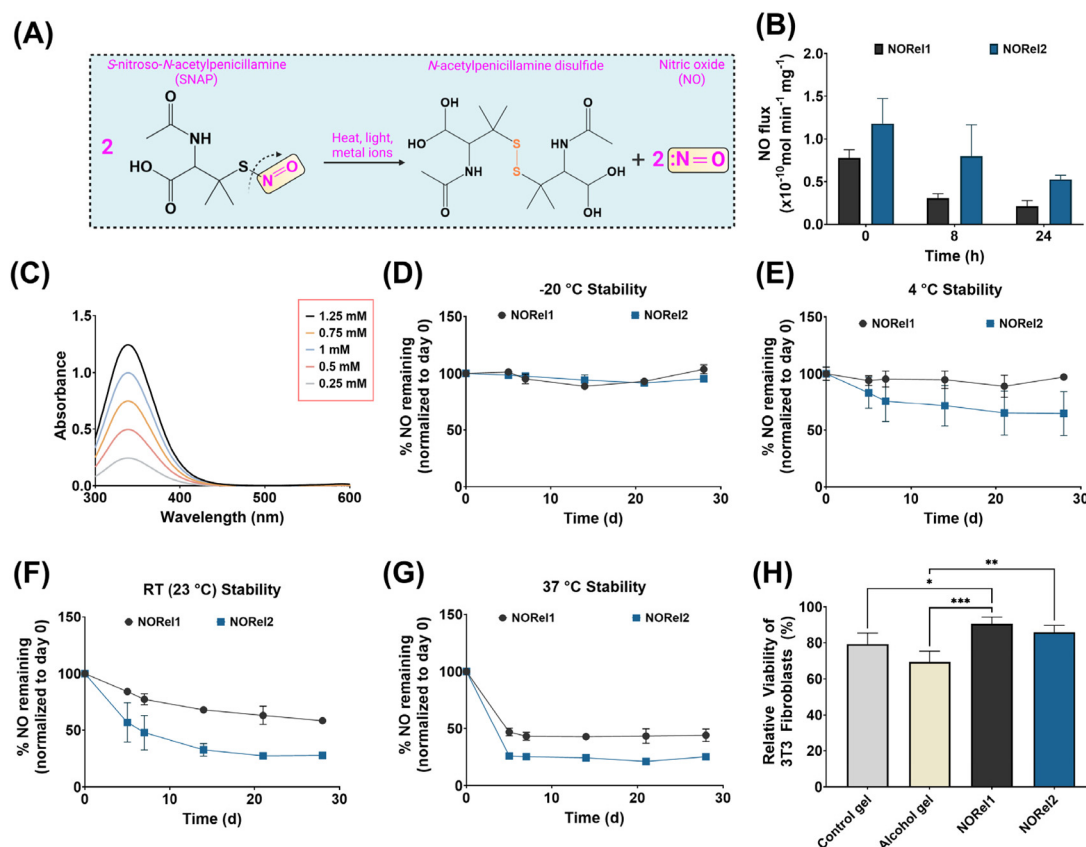
SNAP. The pH values for the 0%, NOReL1, and NOReL2 gels were 8.04 ± 0.42 , 7.96 ± 0.48 , and 7.40 ± 0.16 , respectively (Fig. 2A). Notably, the incorporation of SNAP led to a reduction in the pH level of the NOReL gel, attributable to the acidic nature of the SNAP moiety with the carboxylic acid group.²² However, the pH of the gel remained within the safe range for skin application, posing no risk of skin imbalances.²⁸

To determine the optimal storage conditions for preserving the stability of carbomer and SNAP, NOReL1, and NOReL2 gels were subjected to storage at various temperatures ($-20\text{ }^{\circ}\text{C}$, $4\text{ }^{\circ}\text{C}$, room temperature ($\sim 23\text{ }^{\circ}\$

stabilize carbomers in NOrel gels. Hydrogen bonds, which are attractive forces between molecules, have a profound impact on the gel's properties. In this case, glycerin played a crucial role by facilitating the formation of hydrogen bonds among the gel's molecules, thereby enhancing its structural integrity and viscosity. Additionally, the SNAP moiety, with its carboxylic acid and secondary amine groups, played a dual role in the gelation process. Firstly, it contributed to the neutralization of carboxylic acids, akin to TEA. Secondly, SNAP participated in the formation of hydrogen bonds, similar to glycerin.³⁷ These combined effects resulted in an increased quantity of hydrogen bonds within the gel. Consequently, when higher concentrations of SNAP were introduced into the gel, the number of hydrogen bonds formed increased, leading to augmented gel strength and viscosity. This intricate interplay of carbomer, TEA, glycerin, and SNAP collectively contributed to the stability, consistency, and overall performance of the gel. These findings align with previous research on Pluronic F127 where the viscosity of gel increased due to the addition of RSNO donor.³⁸ With these adjustable physical properties, further investigations into NO release were pursued.

3.3. Nitric oxide release kinetics

In the pursuit of robust infection control measures, the role of NO has gained considerable attention for its potential. Nonetheless, the intrinsic instability of NO donors in solution has presented a substantial hurdle, limiting the ability to achieve sustained and efficacious levels of NO. To overcome this, inventive strategies have emerged, centered on the use of polymeric matrices, that can stabilize and prolong the release of NO from materials.³⁹ These strategies offer a protective shield to NO donors, preserving NO and prolonging its release characteristics that enable consistent, regulated, and highly efficient release of NO. For this reason, a robust NO donor SNAP was used to formulate the hand sanitizer gel. In the presence of heat, light, and metal ions, the thiol bond is cleaved and can readily release NO from the donor (Fig. 3A). The NO release from NOrel hand sanitizer gel was characterized using a chemiluminescence method (Fig. 3B). The study aimed to assess the adaptability of NOrel gel by varying the SNAP content and its impact on NO release under physiologically relevant conditions. Specifically, NOrel1 and NOrel2 gels were



subjected to NO release testing within an amber sample cell maintained at a temperature of 37 °C. The results demonstrated that, on average, NORe12 gel exhibited the highest NO release rates during the entire 24 h study period. The concentration of SNAP within the gel significantly influenced the general trends in NO release. Notably, at 0, 8, and 24 h, NORe1 gel released 0.78 ± 0.10 , 0.30 ± 0.05 , and $0.21 \pm 0.07 \times 10^{-10}$ mol min⁻¹ mg⁻¹ of NO, respectively, while NORe12 released 1.18 ± 0.29 , 0.80 ± 0.37 , and $0.53 \pm 0.05 \times 10^{-10}$ mol min⁻¹ mg⁻¹ of NO, respectively (Fig. 3B). Over time, the levels of NO declined as SNAP decomposed in the gel. Prior studies have examined NO release from hydrogels composed of Pluronic F127, alginate, gelatin, and similar materials for antibacterial purposes that showed comparable levels of NO from hydrogels over 24 h.^{38,40} However, the combination of traditional hand sanitizer components with NO represents a novel approach that has not been explored before. In summary, both NORe1 and NORe12 gels have demonstrated their ability to release NO under physiological conditions, indicating the increase in NO release by altering the NO donor amount within the gel.

These outcomes validate the feasibility and effectiveness of NORe1 hand sanitizer gel as a proof-of-concept. Importantly, it should be noted that the duration, quantity, and extent of NO release can be further tailored to specific situational needs by changing the NO donor concentration, varying the type of NO donor, or even changing the gel composition. The enhanced NO release exhibited by NORe1 gels, in combination with other antimicrobial agents like ethanol and tea tree oil, holds therapeutic potential, particularly in combatting the proliferation of opportunistic pathogens. This innovative approach could have significant implications in the field of infection control and prevention.

3.4. SNAP stability in NORe1 gel

The stability of SNAP in alcohol-based formulations, in conjunction with carbomer, under various thermal conditions is not well understood. This lack of understanding could potentially affect the shelf-life of NORe1 gels. From a commercial perspective, it is crucial to determine the optimal storage conditions for these gels, considering potential clinical applications. To investigate the stability, NORe1 and NORe12 gels were subjected to different temperatures (−20 °C, 4 °C, room temperature (~23 °C), and 37 °C) to simulate real-world storage conditions and the percent of NO (%NO) remaining in the gels was monitored over 28 d, relative to NO donor SNAP present in fresh gels on day 0. The absorption spectra of SNAP at 340 nm wavelength were used to analyze SNAP concentrations in gel formulations by dissolving 100 mg of gel in 1 mL of 10 mM PBS (pH 7.4) with 100 μM EDTA. A standard curve of SNAP in PBS-EDTA was plotted to quantify the amount of SNAP in the gel (Fig

and the commercial alcohol-based gel alternative, suggesting acceptable lack of cytotoxicity of the gel in absence of NO release. In contrast, both the NOrel1 and NOrel2 formulations demonstrated an upward trend in cellular proliferation with respect to the alcohol alternative ($p < 0.05$), while the NOrel1 gel formulation also demonstrated improved viability compared to the gel control. The increase in viability with the NOrel1 and NOrel2 gel formulations is likely attributable to presence of NO donor SNAP, which has previously shown a proliferative effect for fibroblast cells in several previous NO-releasing materials.^{45–47} By promoting fibroblast proliferation, the NO-releasing gels offer a key benefit over the control and commercial gels in performing better against bacteria while concurrently developing cytoprotective effects ideal for topical application.

3.6. Antibacterial efficacy of NOrel hand sanitizer

The broad-spectrum antimicrobial efficacy of NOrel gel was assessed against four microbes frequently linked to hospital-acquired infections: *S. aureus*, *E. coli*, *C. albicans*, and a clinical

isolate of methicillin-resistant *S. aureus* (MRSA) using microbial-killing assay. The antimicrobial effectiveness test demonstrated that the prepared formulations of NOrel gels had significant antimicrobial activity against Gram-positive, and Gram-negative bacteria and *C. albicans* yeast (Fig. 4A–D). The killing efficiencies of each sanitizer gel are reported in Table S3.† The control gel, without SNAP, individually led to >70% reduction in pathogen viability compared to untreated control, resulting in log reductions of 0.81 ± 0.04 , 1.05 ± 0.12 , 1.25 ± 0.10 , and 0.56 ± 0.12 against *S. aureus*, *E. coli*, MRSA, and *C. albicans*, respectively. This can be attributed to the antimicrobial efficacy of the ethanol and tea tree oil in the gel. The addition of SNAP to gel significantly enhanced its antimicrobial potential, with NOrel1 gel achieving log reductions of 1.97 ± 0.04 , 1.98 ± 0.09 , 1.71 ± 0.04 , and 1.04 ± 0.35 against all microbial strains under evaluation.

Notably, NOrel2 gel, containing higher amounts of SNAP, exhibited the highest efficacy among all formulations, achieving log reductions of 2.30 ± 0.27 , 2.34 ± 0.17 , 2.32 ± 0.15 , and

1.59 ± 0.28 against *S. aureus*, *E. coli*, *MRSA*, and *C. albicans*, respectively, compared to untreated control (Fig. 4E–H). These findings are consistent with the NO release levels observed in Fig. 3B, where higher levels of NO release correlated positively with the amount of SNAP incorporated into the gel. *N*-Acetylpenicillamine, the synthetic precursor to SNAP, does not produce any antibacterial effect, which has led to the conclusion in literature that the bactericidal effect of SNAP is solely a result of the NO released from the gel.⁴⁸ While numerous antibiotics focus on specific pathways within microorganisms, NO utilizes a multifaceted approach to deactivate microorganisms. It disrupts essential proteins, damages DNA, and affects cell membranes, among other actions.^{49–52} This comprehensive strategy enables NO to exhibit broad-spectrum antimicrobial properties, effective against various bacteria, fungi, and viruses. This characteristic of NO has been consistently proven in prior research, and the outcomes observed with this innovative material are in line with findings from previously published studies.^{21,53}

3.7. Disinfection efficacy of NOrel gel using *ex vivo* infected rabbit skin model

The antimicrobial effectiveness of NOrel gel was further assessed through an *ex vivo* animal study using rabbit skin specimens. The explanted skin specimens from euthanized rabbits were exposed to *S. aureus* bacteria for 12 h to replicate real-world conditions, allowing bacteria to adhere to the skin. Subsequently, NOrel2 gel, chosen due to its superior NO release and *in vitro* antimicrobial performance compared to NOrel1, was applied and left to incubate on the skin for 6 h to enable NO action against the bacteria (Fig. 4I). In this *ex vivo* study, NOrel2 gel's performance was compared to that of a commercial alcohol-based gel, which served as a secondary test group. Both NOrel2 and the alcohol-based gel were assessed against a bacterial control group, which consisted of infected skin samples with no active antimicrobial treatment. Results from this investigation revealed that NOrel2 gel exhibited similar antimicrobial effects to the commercial alcohol-based gel containing 62% ethyl alcohol as its active component. Notably, both the commercial alcohol and NOrel2 gel demonstrated exceptional bacterial reduction, with >99.99% bacterial eradication (Fig. 4J) and reductions of 5.28 ± 0.07 and 5.23 ± 0.33 log in bacterial viability compared to the untreated control, respectively (Fig. 4K and L). Similar animal models have been employed in previous studies to demonstrate the antimicrobial potential of gels on skin.⁵⁴ Various NO-releasing gels have been developed, serving as versatile platforms for angiogenesis, antimicrobial purposes, wound healing therapies, and formulations in the form of topical lotions.^{55,56} As an endogenous regulatory molecule, NO serves multiple advantageous roles, including providing localized immunity against infectious pathogens.

Until recently, the progress of topical NO treatments was impeded by difficulties in securely storing and effectively delivering NO to infection or inflammation sites. Fortunately, with the advancement of stable NO donors, it has become



Fig. 5 (A) Schematic illustration of difference between standard alcohol containing sanitizer vs. NO-releasing hand sanitizer gel (NORel). Alcohol containing gels with no secondary antimicrobial action evaporate quickly and lack in continuous activity overtime. NO releasing gel with other

NO at physiologically relevant levels for at least 24 h and all gel variants exhibited biocompatibility, confirming their safe nature. NOrel gels effectively eliminated bacteria and fungi, including antibiotic-resistant MRSA, with over 97% efficiency compared to untreated controls. Their effectiveness was also confirmed on an infected rabbit skin model that showed more than 99.99% efficacy against *S. aureus* bacteria. Furthermore, NOrel gels demonstrated persistent antimicrobial effects even 2 h after application, in contrast to commercial alcohol-based gels, which showed no significant impact due to their short-lived action and rapid evaporation. This research presents a biocompatible NO-releasing gel with superior antimicrobial properties compared to alcohol-based sanitizers, offering an effective hand hygiene solution, especially in high-risk environments. The NOrel gel is expected to provide extended protection, eliminate antibiotic-resistant pathogens, and mitigate skin-related side effects.

Data availability

The data supporting this article have been included as part of the ESI.† Data will be made available upon a reasonable request.

Conflicts of interest

Dr Elizabeth J. Brisbois is a co-founder of Nytrix, Inc., a startup company which is involved in exploring possibilities of using nitric oxide-releasing materials for medical applications.

Acknowledgements

This work was supported by the National Institute of Health through the funds received under NIH R01HL151473. Graphics were created by the authors using the BioRender.com software.

References

- 1 M. Haque, M. Sartelli, J. McKimm and M. Abu Bakar, *Infect. Drug Resist.*, 2018, **11**, 2321–2333.
- 2 *Preventing Health Care-Associated Infections*, ed. A. S. Collins, Agency for Healthcare Research and Quality, Rockville (MD), 2008.
- 3 H. Saito, K. Inoue, J. Ditai, B. Wanume, J. Abeso, J. Balyejussa and A. Weeks, *Antimicrob. Resist. Infect. Control*, 2017, **6**, 1–12.
- 4 F. Tusabe, J. Nanyondo, M. J. Lozier, M. Kesande, O. Tumuhairwe, M. Watsisi, F. Twinomugisha, A. Medley, J. Mutoro and M. Lamorde, *Am. J. Trop. Med. Hyg.*, 2023, **109**, 191.
- 5 E. L. Larson, J. Cimi

- 33 S. Nurman, R. Yulia, Irmayanti, E. Noor and T. Candra Sunarti, *Sci. Pharm.*, 2019, **87**, 32.
- 34 R. Hamed, T. Al Baraghthi, A. Z. Alkilani and R. Abu-Huwaij, *J. Pharm. Innovation*, 2016, **11**, 339–351.
- 35 S. Sommatitis, M. C. Capillo, C. Maccario, R. Rauso, E. D'Este, M. Herrera, M. Castiglioni, R. Mocchi and N. Zerbinati, *Gels*, 2023, **9**, 108.
- 36 M. T. Islam, N. Rodríguez-Hornedo, S. Ciotti and C. Ackermann, *Pharm. Res.*, 2004, **21**, 1192–1199.
- 37 A. N. Lyapunov, E. P. Bezuglaya, N. A. Lyapunov and I. A. Kirilyuk, *Pharm. Chem. J.*, 2015, **49**, 639–644.
- 38 L. M. Estes Bright, M. R. S. Garren, M. Ashcraft, A. Kumar, H. Husain, E. J. Brisbois and H. Handa, *ACS Appl. Mater. Interfaces*, 2022, **14**, 21916–21930.
- 39 M. K. Chug, N. Crutchfield, M. Garren, H. Handa and E. J. Brisbois, *ACS Appl. Bio Mater.*, 2024, **7**, 2993–3004.
- 40 S. Ghalei, M. Douglass and H. Handa, *ACS Biomater. Sci. Eng.*, 2021, **8**, 273–283.
- 41 W. R. Mathews and S. W. Kerr, *J. Pharmacol. Exp. Ther.*, 1993, **267**, 1529–1537.
- 42 R. Kumar, H. Massoumi, M. K. Chug and E. J. Brisbois, *ACS Appl. Mater. Interfaces*, 2021, **13**, 25813–25824.
- 43 L. Grossi, P. C. Montecvecchi and S. Strazzari, *J. Am. Chem. Soc.*, 2001, **123**, 4853–4854.
- 44 International Organization for Standardization (ISO), Biological evaluation of medical devices – Part 5: Tests for *in vitro* cytotoxicity, 3rd edn, ISO 10993-5:2009, 2009, <https://www.iso.org/standard/36406.html>.
- 45 M. Garren, P. Maffe, A. Melvin, L. Griffin, S. Wilson, M. Douglass, M. Reynolds and H. Handa, *ACS Appl. Mater. Interfaces*, 2021, **13**, 56931–56943.
- 46 M.