Improved Methods for Assessing Therapeutic Potential of Antifungal Agents Against Dermatophytes and their Application in the Development of NP213, a Novel Onychomycosis Therapy Candidate

Running Title: Novel Onychomycosis Testing Methods and New Peptide Therapy

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Abstract

Onychomycosis is a common, difficult to treat nail infection, mainly caused by dermatophytes. Current therapies are not wholly effective and are associated with manifold side effects. The development of treatments for onychomycosis is challenging because standard \textit{in vitro} tests are not predictive of antifungal efficacy within the nail. We have developed a new antifungal agent, NP213, for the treatment of onychomycosis. NP213 is based on endogenous host defence peptides (HDP) produced within the nail. We compared the \textit{in vitro} activity of NP213 and existing antifungal agents using conventional antimicrobial susceptibility test (AST) systems and more physiologically relevant models based on human nail. We observed that the standard \textit{in vitro} AST methodologies failed to predict the efficacy of antifungal agents within the nail. To address that, we present a more physiologically relevant modified AST method. This method, alongside other standard \textit{in vitro} assessments of activity (including mechanism of action and time of kill studies) better reflected the activity of NP213 and other antifungal agents within the nail than standard \textit{in vitro} AST methods. NP213 is a rapidly acting, fungicidal peptide, superior to existing antifungal agents \textit{in vitro}. It penetrated the nail more effectively than other antifungals, as confirmed using an optimised \textit{in vitro} nail infection model. The data presented supports the current clinical development status of NP213 as a novel agent for treating onychomycosis. We propose that the modified tests developed and applied for NP213 characterisation are the most relevant to use for screening any potential therapeutic candidates for onychomycosis.
Onychomycosis (fungal infection of the nail) is notoriously difficult to treat. The majority of patients receiving any of the limited number of currently available treatments (mainly azole- or allylamine-based) often fail to respond or relapse (1-3). The nail is a highly effective biological barrier and, hence, delivery of therapeutic agents to the nail and nail bed is challenging (4, 5). Keratin, the major constituent of the nail and the primary carbon and nitrogen source for dermatophytes binds to and inactivates many of the existing small molecule antifungal classes, further compromising therapy (6-8). Not surprisingly perhaps, the overall efficacy of current antifungal agents in onychomycosis is poor (4). Paradoxically, the potent in vitro antifungal data for the active agents of these therapies would indicate otherwise. On the one hand safer, more effective therapies are required for onychomycosis. On the other, more clinically predictive, physiologically relevant in vitro systems to test new compounds with therapeutic potential for this common condition are required.

To circumvent the limitations and challenges of drug delivery to the nail, we have taken a biological approach to combatting onychomycosis and designed a novel therapeutic agent derived from antimicrobial, or host defence, peptides (HDP). HDP are an essential component of the innate immune response to infection, and are expressed and produced in the skin and nail (9-12). We have developed a synthetic antimicrobial peptide, NP213 (Novexatin), which is now in late-stage clinical development as a topical therapy (in a water-based solution) for the treatment of onychomycosis. Preliminary data from a phase I and IIa clinical study in a total of 60 subjects confirmed that when dosed daily for 28 d, NP213 was very well tolerated in patients with onychomycosis and
brought about improvement of nail appearance (during and sustained after application; the NP213 effect was greater than placebo) and mycological improvement/resolution of infection when nails were assessed for dermatophyte burden 180 d post a 28 d application cycle of NP213 or placebo (NP213 effect was greater than placebo in patients with mild to moderate onychomycosis) (unpublished data). Here, we present the development and characterization of NP213 using standard in vitro tests, as well as a novel in vitro nail infection system and a modified in vitro MIC-determination protocol developed herein as more physiologically relevant for testing the activity of anti-dermatophyte agents. We show that NP213 is a promising, novel, first-in-class therapy candidate for nail fungus. We also demonstrate a reverse paradox to that detailed above for small molecule antifungals. Namely, standard in vitro AST systems [e.g. Clinical and Laboratory Standards Institute (CLSI) or European Committee on Antimicrobial Susceptibility Testing (EUCAST) methods employing nutrient-rich media (13-15)] do not accurately predict the activity of NP213 and other antifungals within the nail. We recommend that the optimised in vitro nail infection system and the modified in vitro AST protocol we present are used as a novel development strategy in conjunction with standard in vitro tests for more accurate assessment of the curative potential of new onychomycosis agents.

RESULTS AND DISCUSSION

Rational design of NP213. Most HDP are short (<50 aa), cationic, and amphipathic peptides produced by all complex life forms as first line defence against microbes (16-18). Using HDP as a blueprint, NP213 (Arg7) was designed as a highly
hydrophilic and positively charged (net charge +7) cyclic peptide. We anticipated that the charge would facilitate nail penetration by NP213, since the nail is a negatively charged aqueous hydrogel under physiological conditions (19). Further, NP213 is smaller than endogenous HDP (7 aa vs. ca. 30<100 aa) that are known to be present within the nail (9, 11, 12, 20, 21). One of the known drawbacks of peptide and protein therapeutic candidates is their susceptibility to hydrolysis, especially proteolysis (22). This is of particular concern with anti-dermatophyte agents, as dermatophytes produce multiple proteases and peptidases that enable them to hydrolyse keratin (23-25). NP213 was designed specifically to circumvent these issues. Firstly, as an all-arginine peptide, the limited sequence diversity and only R-R bonds within the peptide are susceptible to very limited classes of endoproteases (26). Secondly, the cyclic structure of NP213 minimises the probability of hydrolysis by exoproteases. NP213 was synthesized by standard solid phase synthesis processes as an acetate peptide salt in amorphous crystalline form, its purity determined by reverse phase-high performance liquid chromatography (data not shown), and used as an aqueous solution for the ensuing \textit{in vitro} testing.

\textbf{\textit{In vitro} activity of NP213 against dermatophytes using conventional test methods.}

\textbf{Time-of-kill, sporicidal and anti-hyphal activity.}

Dermatophytes grow vegetatively as hyphae but also produce spores (micro/macroconidia and occasionally chlamydospores), which are dormant, asexual, and non-motile cells, and are normally less susceptible to the antifungal activity of
existing drug classes, including terbinafine (TRB), ciclopirox, amorolfine, and bifonazole, than actively metabolising cells (27-30). Spore production in the nail (31) is thought to contribute to treatment failure and recurrence of onychomycosis (32). The time of kill of a nominal dose of 1000 mg/L NP213 was therefore determined against both spores and hyphae of *T. rubrum* NCPF0118. NP213 was equally efficacious against spores and hyphae of *T. rubrum* NCPF0118; a >3-log (99.9%) kill of spores and hyphae of *T. rubrum* NCPF0118 took 3–4 h (6.47–9.47 × 10^4 cfu/ml killed). By contrast, 0.01 mg/L (2 × MIC) of TRB failed to kill spores or hyphae of *T. rubrum* NCPF0118 even after 24 h (Fig. 1). This demonstrates that NP213 is fungicidal and killed more rapidly than TRB (33) under these test conditions.

NP213 targets the fungal cytoplasmic membrane. We hypothesized that the antifungal activity of NP213 occurs via membrane perturbation and disruption/permeabilization, as that is the most common mechanism of action of the cationic HDP used as a blueprint for the design of NP213 (18, 34). To establish whether that was indeed the case, propidium iodide (PI) staining of a spore suspension of *T. rubrum* NCPF0118 exposed to 500 or 1000 mg/L NP213 for 18 h was performed. PI is a membrane-impermeant fluorescent dye that fluoresces when it intercalates double-stranded DNA (35). Therefore, it is a useful indicator of membrane permeabilization and cell death (Fig. 2A). The number of PI-stained *T. rubrum* NCPF0118 cells in samples incubated with NP213 was significantly higher than that in controls in the absence of NP213 (p<0.005; one-way ANOVA with Tukey’s multiple comparisons test following log transformation of data) and was dose-dependent as the difference in the number of PI...
stained cells was significantly higher when *T. rubrum* NCPF0118 was exposed to 1000 mg/L NP213 compared to exposure to 500 mg/L NP213 (*p*<0.01; one-way ANOVA with Tukey's multiple comparisons test following log transformation of data). This suggested that membrane permeabilization occurred upon exposure to a range of concentrations of NP213. This further confirmed that NP213 was fungicidal and that at least one of its mechanisms of action involved membrane permeabilization.

Scanning electron microscopy (SEM) and transmission electron microscopy (TEM) analyses were then performed to investigate the effect of NP213 on *T. rubrum* cell morphology directly. For SEM analysis, *T. rubrum* NCPF0118 cells were exposed to NP213 (100 x MIC, 100 g/L), and two marketed antifungal agents, TRB (0.5 mg/L, 100 x MIC) or ciclopirox olamine (100 mg/L, 100 x MIC), for 48 h before SEM analysis (Fig. 2B and Fig. S1). The appearance of cells exposed to ciclopirox and TRB was similar to that of untreated cells. In contrast, the hyphae of cells exposed to NP213 appeared flatter, which was indicative of a potential loss of internal turgor pressure following membrane lysis (Fig. 2Biv). The presence of cellular debris on the surface of hyphae supported the occurrence of cell lysis. Since the SEM analysis cannot be used for a direct evaluation of membrane lysis, the effect of NP213 on *T. rubrum* NCPF0118 was also analysed by TEM (Fig. 2C). For TEM analysis, cross-sections of hyphae exposed to 10 g/L NP213 (10 x MIC) for 0, 6, 18 or 48 h revealed clear loss of cellular contents (Fig. 2Cii – iv) when compared to control cells, in which distinct cellular organelles were visible (Fig. 2Ci). This indicated that unlike currently available onychomycosis agents, NP213 is novel in directly targeting the fungal cytoplasmic membrane. Further, NP213
significantly reduced the number of germ tubes formed by *T. rubrum* NCPF0118 (Fig. S2), although it did not affect the length of the formed germ tubes (data not shown).

**Antimicrobial activity of NP213 is cationic charge-dependent.** Each molecule of NP213 has a net positive charge of +7 to facilitate penetration of the nail (19-21) and to enable initial binding of NP213 to negatively charged microbial membranes via electrostatic interactions, a characteristic of most positively charged HDP (18, 36, 37). The importance of the positive charge on the antifungal activity of NP213 was investigated by incubating *T. rubrum* NCPF0118 with 1 mM NP213 and different molar equivalents of polyanetholesulfonic acid (PASA; liquoid) (Fig. 3). PASA is an anionic derivative of polystyrene that binds to and neutralises antimicrobial factors in the blood, including complement (38). As the concentration of PASA increased relative to that of NP213, exceeding the equimolar ratio, the inhibition of growth of *T. rubrum* NCPF0118 by NP213 became less evident. The data was analysed by 2-way ANOVA and Dunnett’s multiple comparisons test to compare the effect of PASA with no exposure to PASA, which indicated that once the molar ratio of PASA:NP213 exceeded 2:1 then the lack of growth inhibition due to neutralisation of the positive charge of NP213 was statistically significant (*p*<0.0001). Thus, the positive charge of NP213 was indeed essential for its antifungal activity.

**NP213 MIC values determined using standard broth microdilution AST.** Considering its effect on fungal growth *in vitro*, the minimal inhibitory concentrations of NP213 required to achieve these actions were determined using the current standard broth microdilution procedure for filamentous fungi (13) against a broad range of dermatophytes and non-dermatophyte fungi that cause onychomycosis. Dermatophyte
MIC values were from <100 – 4000 mg/L and MFC values from 500 – 4000 mg/L (complete MIC data set for dermatophytes is provided in Table S1). NP213 also demonstrated activity against yeasts and non-dermatophyte fungi known to cause onychomycosis (data not shown). The dermatophyte MIC values, whilst high compared to values obtained for other antifungals including TRB, ciclopirox, and azoles used to treat onychomycosis, nevertheless demonstrated that NP213 can inhibit growth of a range of dermatophytes causative of onychomycosis. The possibility of adaptive resistance to NP213 by selected target dermatophytes was assessed in vitro with no evidence of resistance development against NP213 over 20 passages, but with evidence of resistance to TRB (data not shown).

It is now widely recognised that standard antimicrobial susceptibility testing protocols, including CLSI (13) and EUCAST (15) protocols, fail to adequately account for the environmental factors that are present during host-pathogen interactions in vivo and can appreciably impact antibiotic efficacy in vitro, and that the media on which these tests are based do not mimic the host environment (39-43). Furthermore, standard antimicrobial susceptibility tests were devised as long ago as 1961 and were not designed to interrogate the activity of non-antibiotic antimicrobials, including antimicrobial peptides (AMP) such as NP213. We therefore developed and tested novel in vitro methods for antimicrobial susceptibility testing of anti-dermatophyte agents under more physiologically appropriate conditions.

In vitro activity and ex vivo efficacy of NP213 testing using novel, physiologically relevant methods.
Development of a modified MIC determination method for dermatophytes. The broth microdilution methods used to determine the MIC and MFC values utilise the relatively nutrient-rich RPMI-1640 liquid medium as a test matrix (13). This medium contains 2 g/l glucose as the main carbon source, no protein, and only small amounts of individual amino acids (0.01–0.2 g/l). By contrast, the human nail is predominantly made up of different types of keratin that are highly cross-linked by disulphide bonds (44, 45) (45). Consequently, the nutrient content of RPMI-1640 liquid medium bears little resemblance to the nutrients available to fungi growing in/on the nail.

We sought to develop an antifungal susceptibility testing protocol that would resemble clinical/in vivo conditions more closely than conventional MIC testing methods, based on keratin; the major carbon and nitrogen source for dermatophytes within human nail. The medium used for broth microdilution MIC testing (13) was therefore modified to a protein-based test medium containing almost no carbohydrate (44), from the carbohydrate-rich (2.0% w/v as glucose), protein-free RPMI-1640 liquid medium. Different sources of keratin, representative of keratin-rich epithelia and known to support dermatophyte growth were selected (46-49). Powdered human nail material, human skin keratin, or sheep’s wool keratin, all as suspensions in 10 mM sodium phosphate buffer, pH 7.0 were investigated as a nutrient source in vitro for dermatophyte growth. Suspensions of test media precluded monitoring fungal growth by turbidity and so antifungal susceptibility was assessed by fluorescence-based determination of fungal metabolism using the metabolic activity indicator, AlamarBlue™ (50). In preliminary antimicrobial susceptibility testing in modified media containing a
nominal concentration of the different keratins at 0.5% (w/v), the MIC values of NP213 against *T. rubrum* NCPF0118 were 4–62.5-fold lower than those obtained when the fungi were grown in RPMI-1640 liquid medium (Table 1). Interestingly, MICs varied, depending on the source of the keratin, with MICs for NP213 lowest in media containing powdered human nail. To determine whether these reductions in MIC, particularly that seen in human nail powder-based media, was in part due to reduced background metabolism of the fungi versus when in glucose-rich RPMI 1640, the metabolic growth of *T. rubrum* NCPF0118 was assessed for up to 168 h in 10 mM sodium phosphate buffer, pH 7.0, containing 0.5% (w/v) of the different keratins was assessed without the presence of NP213. This revealed (Fig. 4) that in fact only fungi cultured in the human nail powder-derived keratin had a metabolic growth profile in line with that achieved in RPMI-1640. Growth was somewhat reduced versus either of these culture conditions in the skin and sheep’s wool keratin-based media. Human nail material was therefore selected for subsequent studies as not only is this the most relevant keratin source when investigating onychomycosis and potential therapeutic agents but seemed to best support the growth of *T. rubrum*. We next determined whether 0.5% (w/v) nail powder was optimal as the concentration of nutrient source to employ in these assays. Interestingly, concentrations as low as 0.1% (w/v) human keratin supported the metabolic activity of *T. rubrum* NCPF0118, but not to the extent where metabolic activity was in line with that seen in RPMI-1640 liquid medium as a benchmark, whereas concentrations in the range of 0.25% to 0.75% (w/v) nail powder (plateauing thereafter) in 10mM sodium phosphate did achieve the same metabolic activity as RPMI-1640 liquid medium and were employed in subsequent experiments.
Antifungal activity of NP213 assessed using the modified MIC determination method. The MIC values of NP213 were further investigated in the modified antimicrobial test system (containing only human nail material) alongside those for TRB and ciclopirox olamine against *T. rubrum* NCPF0118. Activity in this system was compared with MICs obtained using the CLSI broth microdilution protocol (Table 2). The MIC (determined by the CLSI broth microdilution procedure) of NP213 was reduced by up to 128-fold when determined using the modified protocol. The apparent MIC of TRB increased 2–16-fold and that of ciclopirox increased as much as 16-fold, but was not completely inhibited despite it being described that these antifungal agents are known to bind to nail keratin with a concomitant inhibition of antifungal efficacy (6, 8, 51). We believe that this nail keratin-based medium (containing 0.5% (w/v) powdered human nail) is optimal for determining the antifungal activity of a range of onychomycosis treatment candidates as it supports high levels of dermatophyte metabolic activity (Fig. 4) whilst at the same time, being a more accurate demonstration of the physiological conditions encountered by onychomycosis-causing fungi in the nail.

Overall, our data support the application of the modified MIC method we developed in the assessment of activity of antifungal agents against dermatophytes. The levels of keratin in this medium are not inhibitory to antifungal agents known to bind to and be inactivated by keratin. This method facilitates testing against fungi in a metabolic state more closely resembling their status within the nail and as detailed below, activity of these agents within the nail.

Optimisation of an *in vitro* model of nail infection. The *in vitro* activity and mechanism of action determined for NP213, the MIC data obtained using the modified
MIC method and preliminary clinical data obtained thus far together suggested that NP213 might be highly efficacious in the nail environment. *In vitro* nail models have previously been described for testing candidate onychomycosis therapies, including those based on sections of human nail, animal hoof slices and films of hair keratin (52-55). However, these are not entirely physiologically relevant because animal hoofs do not closely resemble the human nail in terms of keratin content. Further, the human nail models generally do not use full-thickness human nails or only use very small fragments thereof. We therefore optimised the *in vitro* full thickness model of dermatophyte nail infection (as described in the Materials and Methods; Fig. 5). A higher fungal inoculum in our own model versus other published *in vitro* models (2 x 10^7 spores/nail fragment) facilitated successful establishment of deep-seated infections with *T. rubrum* NCPF0118 (Fig. 5B, Fig. 6B). Similar infections were also established with *T. rubrum* ATCC-MYA-4438 and *T. rubrum* S52d0 (data not shown).

**Antifungal activity of NP213 in the optimised *in vitro* model of nail infection.** We next used the *in vitro* nail model to determine whether NP213 was active against *Trichophyton* spp. infecting the human nail. Excised human nails were prepared and infected as described in Materials and Methods. NP213 (in a water-based vehicle) or comparator currently marketed antifungals were applied to infected nails for 28 d on a daily basis. Viable fungi were extracted and cultured on potato dextrose (PDA) agar to assess the infection status of the nails as detailed in the Materials and Methods. NP213 application to nails successfully eradicated different strains of *T. rubrum* from infected nails after only 28 d of daily application compared to vehicle alone or water (Table 3), unlike the comparator topical onychomycosis agents ciclopirox and amorolfine (Fig. 6A).
Importantly, there was no evidence of a placebo effect for NP213, as treatment with the vehicle alone caused no significant reduction in the number of *T. rubrum* cfu recovered at the end of the experiments (Kruskal-Wallis test of log transformed data (*p*<0.0001)) (Table 3 and Fig. 6A). Placebo formulations are not available for the ciclopirox or amorolfine comparator products meaning that a placebo effect of these organic-solvent containing products can not be ruled out as a contributing factor of any antifungal effects observed for these products. In addition to culturing viable dermatophytes, fungal elements within the nail were analysed by TEM (Fig. 6B) revealing only the presence of non-viable hyphal ‘ghosts’ (loss of intracellular contents from fungi, but fungal cell walls remaining relatively intact) in NP213-treated nails and more evidence of cellular damage than the intact, viable cells within infected nails treated with ciclopirox or amorolfine (or other agents such as itraconazole, data not shown). Importantly, these data also confirm NP213’s ability to penetrate full nail thickness nail and remain bioactive therein, as the sections visualised by TEM were those were taken from the ventral aspect of nail samples.

The activity/biostability of NP213 within full-thickness human nail was further assessed for up to 11 months following cessation of application of NP213 for 28 d to eradicate an *in vitro* nail infection (Fig. 7). Infected and NP213-treated nails, cleared of fungal infection, were re-exposed to *T. rubrum* NCPF0118 after 3, 5, 8 and 11 months following cessation of NP213 treatment. Infection could not be re-established in NP213 treated nails, but did successfully establish in control, untreated nails inoculated alongside as controls (Fig. 7). Log transformed data for each re-infection time-point was analysed by one-way ANOVA with Holm-Sidak’s multiple comparisons test to compare...
vehicle and NP213 treatment and the reduction in counts following NP213 treatment was statistically significant ($p<0.0001$) at every re-infection time-point.

To further validate our *in vitro* nail model and importantly, to confirm that NP213 was also able to eradicate infection from full-thickness, onychomycotic nails from patients with established, chronic clinical infection, we assessed NP213 efficacy *ex vivo* in infected nails from patients with clinically confirmed onychomycosis caused by dermatophytes (Fig. 8). Only dermatophytes were isolated and enumerated from untreated nails, facilitated by the use of selective dermatophyte test medium (DTM) agar (Becton, Dickinson U.K. Ltd), but it is possible that other infectious agents were present in the naturally infected nails. Moreover, nails may have been infected with more than one dermatophyte strain. Nonetheless, NP213 successfully eradicated all dermatophyte infection from these onychomycotic nails with dermatophyte burdens of $4.3 \times 10^3$ CFU/ml to $>2 \times 10^7$ cfu/ml, a ‘natural state’ inoculum range replicated in our *in vitro* nail model described above.

The predicted value of the novel, combined testing strategy developed herein, combining the standard and modified *in vitro* assays and *in vitro* nail model assays is, we believe, significant for future testing of novel anti-dermatophyte agents. Alongside standard *in vitro* tests, the approach we now recommend mitigates, in part, the paradox of *in vitro* testing results and activity within the nail for anti-dermatophyte agents. It also provides conditions which are more representative of the behaviour of dermatophytes within the nail and of the behaviour of antifungal agents in a nutrient-poor biological matrix such as the nail.
CONCLUSIONS

The need for new, safe and more effective antifungal agents as onychomycosis therapies is obvious and significant. Further, a clear need exists for more predictive in vitro testing methods that can be applied to all antifungal agents. We addressed the first point, the need for new antifungals, by developing NP213. We addressed the second point, the need for better testing methods, by developing a novel, expanded testing system that goes beyond standard, non-physiologically relevant MIC experiments, and employs nail infection models and human nail-based media (the latter as the basis for modified AST assays). Importantly, the tests described herein are predictive of the activity of antifungal agents within the nail and allow them to be tested against pathogenic dermatophytes in a (metabolic) state in which they would exist within the nail. Without this full package of data, the high MIC/MFC values obtained using standard methods (Table 1) would likely have been considered too high to pursue further development of NP213, a now late clinical stage therapeutic candidate for onychomycosis.

MATERIALS AND METHODS

Peptides and chemicals. NP213 was synthesized as an acetate salt (>95% purity) by solid phase synthesis (PolyPeptide Group, France; Almac Group, UK; Ambiopharm Inc, USA). NP213 was prepared as a lyophilized powder and dissolved in sterile-deionized water (sdH2O) to a concentration of 400 mg/ml, unless otherwise stated. All other chemical and growth media were obtained from Sigma-Aldrich (UK), except where otherwise stated.
Fungal isolates. A complete list of fungal isolates used in these experiments can be found in Supplementary Table S1. Dermatophytes were purchased from the National Collection of Pathogenic Fungi (Public Health England, UK), American Type Culture Collection (LGC Standards, UK) and the Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH (DSMZ, Germany). Clinical isolates of dermatophytes were obtained from Prof Michel Monod (Centre Hospitalier Universitaire Vaudois, Switzerland) and Prof Boni Elewski (University of Alabama at Birmingham, USA). Dermatophyte cultures were grown on Sabouraud dextrose (SAB) agar (Oxoid Ltd., UK), for up to 14 d at 30°C. Fungi were stored at -85°C with 0.5% (v/v) dimethylsulfoxide as cryoprotectant. For experimental purposes, dermatophytes were cultured at 30°C for up to 14 d on PDA (Oxoid Ltd).

Time-of-kill method. Spore suspensions of ~1 × 10^5 spores/ml were prepared (13) and used directly to determine the time-of-kill for spores. To test the germlings (germinated spores/short hyphae), spores were allowed to germinate for 12 h at 30°C and examined microscopically (Axiovert 40 CF microscope, Carl Zeiss Ltd, UK) to ensure that germination had occurred, and used in the time of kill experiment as ‘germlings’. For the time-of-kill experiment, 0.5–1 × 10^5 cells/ml were prepared in 10 ml of RPMI-1640 liquid medium and incubated aerobically for 24 h at 30°C. Aliquots (20 μl) were removed at the indicated time-points, serially diluted in RPMI-1640 liquid medium [in the absence of NP213 and in the presence of 3% (w/v) PASA to neutralise residual NP213], and 50 μl (in triplicate) were plated on PDA agar and incubated for up to 7 d at 30°C or until clear fungal colonies were evident in controls. The colonies were then counted to establish fungal killing.
**PI staining.** A spore suspension of *T. rubrum* NCPF0118 (1 × 10^4 spores/ml) was exposed to 0 mg/L, 500 mg/L, and 1000 mg/L NP213 for 18 h at 30°C. Following this, the fungi were exposed to PI, as modified from the manufacturer’s instructions in FungaLight™ CFDA, AM/Propidium Iodide Yeast Vitality Kit (FisherScientific, UK), where fungi were exposed to 16 μM PI for 15 min at 30°C. The cells were viewed by fluorescence microscopy (x 400 magnification) (Zeiss Axiovert 40) and the percentage of cells stained with PI was determined in six independent fields of view in duplicate experiments.

**SEM and TEM.** *T. rubrum* NCPF0118 was prepared for electron microscopy by growing in 12-well polystyrene tissue culture plates (Greiner Bio-one, UK) on sterile 0.2-μm polycarbonate membrane filters (Thermo-Fisher Scientific, UK) submerged in RPMI-1640 medium for 7 d at 30°C. After 7 d, fungi were exposed to either NP213 (2000 mg/L), TRB (0.01 mg/L), or ciclopirox olamine (2 mg/L) for 48 h at 30°C. Filters were then aseptically transferred to sterile Bijou containers (7 ml), submerged in 2.5% (v/v) glutaraldehyde in 100 mM sodium phosphate buffer (pH 7.3), and stored at 2–8°C for 48 h prior to processing for electron microscopy analysis. For SEM analysis, the cells were processed by ethanol and hexamethyldisilazane dehydration series before gold mounting. For TEM analysis, the cells were dehydrated by passing through ethanol and acetone series before being embedded in wax resin, stained with uranyl acetate/lead citrate stains to improve contrast, sectioned at 90 nm, and mounted onto copper grids for TEM analysis. TEM and SEM micrographs were acquired using a JEM-1400 Plus JEOL TEM and a Zeiss EVO MA10 SEM, respectively. Microscopy was performed in the Microscopy and Histology Core Facility at the University of Aberdeen (UK).
Conventional determination of MIC and MFC values. Antifungal susceptibility testing to determine the MIC was conducted by the broth microdilution procedure for filamentous fungi (M38-A2) (13). The MFC was determined according to the method described by Ghannoum, et al. (2011) (56). Briefly, once MIC testing was completed, the contents of the wells were mixed thoroughly; 100 μl culture from each well was spread plated onto PDA agar and incubated at 30°C for up to 7 d. Absence of any growth was deemed fungicidal, whereas growth from wells that contained concentration of antifungals at or above the MIC was deemed fungistatic. Experiments were carried out in triplicate with triplicate samples in each experiment. Selected HDPs have been shown to bind to polystyrene, resulting in reduced activity in antimicrobial susceptibility tests employing such plasticware (57). However, parallel MIC determinations using polypropylene and polystyrene microwell plates with T. rubrum NCPF0118 demonstrated no evidence of such an effect with NP213, with identical MICs in both polypropylene and polystyrene 96-well plates (data not shown). Since polystyrene 96-well plates are optically clear and give lower background optical densities at 530 nm than polypropylene plates, these were used in all relevant experiments.

Modified antifungal susceptibility testing procedure. In selected experiments, antifungal susceptibility testing was carried out as described above, but with the following modifications. RPMI-1640 liquid medium was substituted for 10 mM sodium phosphate buffer (pH 7.0) containing 0.1, 0.25, 0.5, 0.75, and 1.0% (w/v) powdered toenail suspensions and 0.0025% (w/v) AlamarBlue™ (Thermo-Fisher Scientific, UK). AlamarBlue™ is a cell viability reagent that uses the reducing power of living cells to quantitatively measure viability that can be detected using a fluorescence-based plate
reader. Human nails were obtained from an NHS podiatrist (NHS Grampian, UK), following appropriate ethical approval (NHS Grampian Research Ethics Committee REC reference: 05/S0801/115) and informed written consent, from donors following nail avulsion performed because of ingrown toenails. All nails were disease-free, based on the podiatrist’s diagnosis. Nails were cut into small fragments (<5 mm) and ground to a fine powder in liquid nitrogen using a mortar and pestle. Following this, the powdered nails were passed through a fine-meshed sieve and the sieved nail powder was used to prepare suspensions in 20 mM sodium phosphate buffer, pH 7.0, which was subsequently diluted 1:1 following addition of the fungal inoculum. Nail powder suspensions were sterilised by autoclaving for 20 min at 121°C.

Fungal metabolic activity was monitored by fluorescence (λex 530 nm, λem 590 nm) following the reduction of AlamarBlue™ every 24 h for up to 168 h, or until high metabolic activity (≥100,000 U) was observed in the inoculated controls in the absence of any antifungal agent using a BioTek Synergy HTX Multi-Mode Reader (BioTek U. S., Winooski, USA). As a control, antimicrobial susceptibility testing was conducted in RPMI-1640 liquid medium and MICs were determined by measuring both the optical density and metabolic activity. Sodium phosphate buffer (pH 7.0) alone did not support dermatophyte growth (data not shown). The experiments were performed in triplicate with three replicates per each experiment.

**In vitro** human nail model of *Trichophyton* spp. infection and **ex vivo** treatment of onychomycotic nails. An experimental model of *Trichophyton* spp. nail infection was optimised to simulate onychomycosis under laboratory conditions, and to test the antifungal efficacy of NP213 and antifungal comparators (Fig. 5). As an inert
support and to provide a moist atmosphere, sterile water agar plates were prepared in 90-mm Petri dishes [1.5% (w/v) agar in sterile deionised H₂O]. Petri dishes were divided into three sections and individual pieces of sterile silicone rubber tubing (1-cm diameter; 3-mm height) were placed in each section allowing the investigation of three nail fragments per plate. This prevented direct contact between the nails and the agar surface or the lid of the Petri dish. Such set-up was necessary because commercial agar preparation components may bind to cationic peptides, including defensins and polymyxins (58, 59), inhibiting their diffusion and reducing activity. Silicon tubing is permeable to oxygen and CO₂ (60), and its use ensured O₂ access to support fungal growth. Its diameter was sufficiently small to stably support the nail fragments.

Uninfected human nails (sourced under ethical approval as described in the section: modified antifungal susceptibility testing procedure) were cleaned of all residual skin and debris before cutting to size (ca. four fragments were obtained from an adult male great toe nail), and their surface area and thickness were determined using a micrometer (Fig. 5Bi). Nail fragments were submerged in 10 ml of dH₂O and sterilized by autoclaving at 134°C for 20 min. They were then mounted with the ventral (nail bed side, concave) surface facing upwards on the silicon tubing rings. Nail samples were allocated to provide nails of similar average size and thickness for each treatment. Nails were left to dry and incubated to ensure sterility for 7 d in a humid atmosphere at 30°C. Each nail fragment was inoculated on its ventral surface with a spore suspension containing ca. 2 × 10⁷ spores of *Trichophyton* spp. in 0.15 M NaCl (0.01 ml/cm² nail surface area). Petri dishes were incubated at 30°C inside a sealed plastic box containing an sdH₂O reservoir to avoid dehydration, for 14 d, at which point the hyphal
growth was clearly visible on the entire surface of the nail fragment (Fig. 5Bii). Nails within the same Petri dish were always exposed to the same fungal inoculum and treatment, mitigating the risk of cross-contamination. Randomly sampled nails were subjected to TEM analysis (described above) to confirm infection had been established with fungal penetration of the nail fragments (Fig. 5Biii, uninfected, and 5Biv, infected).

For ex vivo assessment of the antifungal activity of NP213, nails from patients diagnosed with onychomycosis (obtained under ethical approval as described in the section: modified antifungal susceptibility testing procedure) were surface-cleaned with 70% isopropyl alcohol swabs (Robinson Healthcare Ltd, UK), but not sterilised.

All nail fragments, whether naturally infected or infected in vitro, were treated by application of NP213, comparator antifungal agents or controls to the dorsal side of the nail. The dorsal (convex) aspect of the nail is that normally exposed to the environment. Antifungal agents were applied (0.01 ml/cm² nail surface area) daily and distributed over the surface of the nail area with a sterile nail lacquer brush applicator (VWR International Ltd., UK), as specified for each experiment.

Following 28 d of daily treatment with NP213, nail fragments were transferred to sterile 2.0-ml microcentrifuge tubes containing 1 ml of SAB broth containing 6% (w/v) PASA, to neutralise residual NP213 (Fig. 3) (61), and with 100 mg of sterile 0.5 mm glass beads. For nail fragments treated with ciclopirox, SAB broth was supplemented with 800 μM iron (III) chloride, to inactivate ciclopirox (62), a concentration that did not affect T. rubrum growth (data not shown). All other nails were processed in SAB broth with 100 mg of sterile 0.5 mm glass beads. The contents of the tubes were mixed thoroughly for 1 min using a bead-beater to release fungal cells from the nail matrix for
enumeration, and stepwise 10-fold dilutions were prepared in SAB broth containing 3% (w/v) PASA. Samples were plated on PDA, incubated at 30°C for 14 d, and colonies counted. Recovery of microbes from nails from onychomycosis patients, NP213-treated and untreated, was carried out using DTM agar (Becton, Dickinson U.K. Ltd) rather than PDA.

Statistical analysis. All statistical analyses were carried out using GraphPad Prism (GraphPad Software, La Jolla, USA). Statistical significance was determined as described in the text.

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DKM, JR, CSS, LM, and VD conducted the experiments. DKM, CSS, and DO were responsible for experimental design and data analysis. DKM, CSS, and DO wrote the manuscript.

DKM, JR and VD are employees of NovaBiotics Ltd. DO is a director of NovaBiotics Ltd. CSS and LM are former employees of NovaBiotics Ltd.

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Niewerth M, Kunze D, Seibold M, Schaller M, Korting HC, Hube B. 2003. Ciclopirox olamine treatment affects the expression pattern of Candida albicans genes encoding virulence factors, iron metabolism proteins, and drug resistance factors. Antimicrob Agents Chemother 47:1805-17.
TABLE 1  Effect of keratin source on the antifungal activity of NP213 against *T. rubrum* NCPF0118. In these experiments, MIC was defined as the concentration of NP213 required for a 100% inhibition of fungal metabolic activity, determined by the modified version of the broth microdilution procedure (see Material and Method for complete details). Median values are shown for three technical replicates from three replicate experiments.

| Keratin source [0.5% (w/v)] in Phosphate buffer<sup>a</sup> | Median MIC (mg/L) |
|---------------------------------------------------------------|-------------------|
| Human nail                                                   | 16–32             |
| Human skin                                                   | 125               |
| Lamb’s wool                                                  | 250               |
| RPMI-1640                                                    | 1000              |

<sup>a</sup>Phosphate buffer, 10 mM sodium phosphate buffer, pH 7.0.
TABLE 2 Antifungal activity of NP213, terbinafine hydrochloride (TRB), and ciclopirox olamine was determined against *T. rubrum* NCPF0118 in RPMI-1640 liquid medium or 10 mM sodium phosphate buffer, pH 7.0, supplemented with powdered human nail material. MIC\textsubscript{100} values were determined using a modified broth microdilution procedure for incubation with human nail material, and a standard broth microdilution in RPMI-1640 (1, 2), and compared. The MICs reported for RPMI-1640 medium were determined based on optical density changes at 530 nm and metabolic activity, with identical results. All experiments were conducted at least three times, with three technical replicates for each experiment.

| MIC\textsubscript{100} in mg/L (fold MIC change\textsuperscript{a}) | NP213  | Terbinafine | Ciclopirox |
|-------------------------------------------------|--------|-------------|------------|
| RPMI-1640                                       | 1000   | 0.005       | 0.5        |
| Nail content                                    |        |             |            |
| (%) (w/v)                                       |        |             |            |
| 0.1                                             | 8–32 (↓32–128) | 0.01–0.02 (↑ 2–4) | 2–4 (↑ 4–8) |
| 0.25                                            | 8–32 (↓32–128) | 0.01–0.04 (↑ 2–8) | 2–4 (↑ 4–8) |
| 0.5                                             | 16–32 (↓32–64) | 0.01–0.08 (↑ 2–16) | 2–4 (↑ 4–8) |
| 0.75                                            | 8–32 (↓32–128) | 0.02–0.04 (↑ 4–8) | 8 (↑ 16)    |

\textsuperscript{a}MIC fold-change in human nail compared to MIC obtained in RPMI-1640 liquid medium is given in parentheses.
TABLE 3 Survival of *T. rubrum* in the *in vitro* infected human nail model (14 d at 30°C) following exposure to 10% (w/v) NP213 in a water-based vehicle or control vehicle without NP213 following daily application for 28 d. The data are presented as the log transformed mean and 95% confidence intervals and analysed by the Kruskal-Wallis test. Data is derived from 3 nail fragments per treatment and triplicate counts per nail fragment. CI = confidence interval.

| Dermatophytes recovered (log (mean cfu/ml & 95% CI)) | Prior to treatment | Vehicle | 10% (w/v) NP213 | p value treatment: vehicle |
|------------------------------------------------------|--------------------|---------|----------------|---------------------------|
| *T. rubrum* NCPF0118                                 | 6.6 (6.5, 6.7)     | 5.9 (5.5, 6.3) | 0.0 (0.0, 0.0) | 0.0082                    |
| *T. rubrum* ATCC-MYA-4438                            | 4.3 (4.0, 4.7)     | 6.0 (5.7, 6.5) | 0.0 (0.0, 0.0) | <0.0001                   |
| *T. rubrum* S29d0                                    | 6.5 (6.2, 6.8)     | 7.0 (6.7, 7.2) | 0.0 (0.0, 0.0) | <0.0001                   |
FIGURE LEGENDS

FIGURE 1 Incubation time required for fungicidal activity of NP213 or terbinafine (TRB) against spores or germlings (germinated hyphae) of T. rubrum NCPF0118 and T. rubrum ATCC-MYA-4438. A. Antimicrobial efficacy of NP213 against T. rubrum NCPF0118. B. Antimicrobial efficacy of NP213 against T. rubrum ATCC-MYA-4438. C. Antimicrobial efficacy of TRB against T. rubrum NCPF0118. Experiments were conducted three times with three technical replicates in each experiment. Data shown is from one representative experiment. Dotted lines on the graphs represent 3 log (99.9%) kill.

FIGURE 2 Mode of action of NP213. A. Propidium iodide (PI) staining of T. rubrum NCPF0118 incubated with NP213. T. rubrum NCPF0118 spores were exposed to NP213 for 18 h at 30°C, followed by a 15-min exposure to PI. Cells were examined by fluorescence microscopy (× 400 magnification). The total number of cells and the number of PI-stained cells were counted in six independent fields of view for each sample in duplicate experiments. The log transformed mean number of PI-stained cells is presented, and error bars represent 95% confidence intervals. Statistical analysis was conducted using one-way ANOVA followed by Tukey’s multiple comparisons test to determine significance (** p<0.001; * p<0.01) between NP213-treated and untreated cell populations. B. SEM analysis of the effect of a 48-h exposure to antifungals on the morphology of T. rubrum NCPF0118. (i) untreated cells; (ii) 100 mg/L ciclopirox; (iii) 0.5 mg/L TRB; and (iv) 100 mg/ml NP213. Single arrows, flattening of the hyphae; double arrows, bulges within the hyphae; arrowheads, damage to the surface of the hyphae;
asterisks, possible blebbing on the hyphal surface. All sample treatments were conducted in triplicate and at least 30 images were acquired for each treatment. Figures are representative images from a single replicate. C. TEM analysis of the effect of duration of incubation with NP213 (10 mg/ml) on the morphology of *T. rubrum* NCPF0118. CW, cell wall; PM, plasma membrane; N, nucleus; V, vacuole. Arrowheads, separation of the plasma membrane from the cells wall; double arrows, plasma membrane destruction. The images are representative of three replicates. Scale bars are 500 nm for control, 6 h, and 48 h treatments; and 100 nm for 18-h treatment.

**FIGURE 3** Antifungal activity of NP213 is dependent on its positive charge. *T. rubrum* NCPF0118 (ca. $1 \times 10^3$ CFU/ml) was incubated aerobically for 96 h at 30°C in the presence of 1 mM NP213 (1093.3 mg/L) and increasing molar equivalents of PASA in RPMI-1640 liquid medium. Data are presented as the mean and error bars represent the 95% confidence intervals of three biological replicates, each containing three technical replicates. Data was analysed for significance by 2-way ANOVA with Dunnett’s multiple comparisons test. PASA alone, at the concentrations used, had no effect on the growth of *T. rubrum* NCPF0118 (data not shown).

**FIGURE 4** Effect of keratin source on the metabolic activity of *T. rubrum* NCPF0118. *T. rubrum* NCPF0118 (ca. $1 \times 10^3$ spores/ml) was inoculated into 10 mM sodium phosphate buffer, pH 7.0, containing 0.5% (w/v) of each keratin source. Cultures were incubated for up to 216 h at 30°C and metabolic activity (AlamarBlue™ reduction) was
monitored every 24 h. Experiments were conducted three times with three technical replicates per experiment and data is presented as the mean ± SD.

**FIGURE 5** The optimized *in vitro* human nail model for evaluating the efficiency of antidermatophyte agents. A. Diagrammatic representation of the *in vitro* human nail model of *Trichophyton* spp. infection. Sterile nails were inoculated with *Trichophyton* spp. (ca. 2 × 10^7 spores/ml) on the ventral (lower) side and incubated at 30°C for 14 d, at which point the nail surface was entirely covered by hyphal growth and fungi had penetrated into the nail matrix. The nail fragments were then inverted so that the dorsal (upper) surface was facing upwards. Investigative substances and controls were applied on the dorsal side only. (Bi–iv). Images of an uninfected and infected nail from the *in vitro* nail infection model. i, Sterile nail fragment prior to inoculation with *T. rubrum* NCPF0118; ii, nail fragment following infection with *T. rubrum* NCPF0118 for 14 d at 30°C; iii, TEM of medial section of a sterile human nail fragment; iv, TEM of medial section of a human nail fragment following infection with *T. rubrum* NCPF0118 for 14 d at 30°C. Arrows indicate cross-sectional views of fungal hyphae within the nail matrix. The images are representative of three replicates. Scale bars, 5 μm.

**FIGURE 6** The efficacy of NP213 and other antifungal agents in the *in vitro* model of infection. A. Antimicrobial activity of an NP213 (in a water-based vehicle), 8% (w/v) ciclopirox nail lacquer, and 5% (w/v) amorolfine nail lacquer against *T. rubrum* NCPF0118. Data are representative of triplicate samples from three nail fragments and data is presented as the log transformed mean and error bars represent the 95%
confidence intervals. Statistical analysis was conducted using Kruskal-Wallis test of log transformed cfu/ml followed by Dunn’s multiple comparisons test to determine significance (** ** p<0.0001; * p<0.05). B. TEM analysis of the effect of NP213, ciclopirox, and amorolfine on fungal morphology within the nail following treatment for 28 d with 10% (w/v) NP213, 8% (w/v) ciclopirox nail lacquer, or 5% (w/v) amorolfine nail lacquer against *T. rubrum* NCPF0118. Nails were treated on a daily basis with the described compounds (and incubated at 30°C) and processed after 28 d to count the live fungi (colony counting) and for TEM. The effect of antifungal treatment was assessed for statistical significance by one-way ANOVA and Tukey’s multiple comparisons test. When compared to the water control, all results were statistically significant (p<0.05), with the exception of the vehicle control, which was not significant. The effect of NP213 when compared to the NP213 vehicle control was also statistically significant (p<0.05). For the TEM analysis, arrows indicate probable dead fungi with cell walls remaining but no evidence of intracellular content. Arrowheads indicate probable live fungi with evidence of intracellular content. The images are representative of three replicate nail fragments per treatment. Scale bars, 1 μm (NP213); 2 μm (amorolfine); 5 μm (ciclopirox and vehicle).

**FIGURE 7** NP213 remains active within the human nail for at least 335 d post-treatment termination, as evidenced by a lack of *T. rubrum* NCPF0118 growth following attempted re-infection of previously treated nail fragments. Fungal recovery from the control nails infected 28, 120, 180, 270, and 365 d after the initial infection, and from nails originally infected with *T. rubrum* NCPF0118 and treated with 10% (w/v) NP213 daily for 28 d at
30°C was determined as described in the Materials and Methods. After the treatment phase, nails were incubated at 30°C for the time points specified, in a humid atmosphere until processing to determine the numbers of surviving fungi. Log transformed colony counts represent the mean and 95% confidence intervals of triplicate samples from each of the three nails per time point.

**FIGURE 8 Ex vivo efficacy of NP213 against onychomycotic nails.** Eight great (large) toe nails from patients with clinically confirmed onychomycosis were treated daily with 10% (w/v) NP213 in an aqueous water-based vehicle or vehicle only for 28 d before recovery of dermatophytes (triplicate counts per nail fragment) on DTM agar. A. Combination data representing the activity of NP213 on all samples compared to vehicle (Unpaired t-test with Welch’s correction following log transformation of data; p<0.0001). B. Table showing the effect on individual nails. Treatment of nails with NP213 caused a statistically significant reduction in the number of cfu recovered for all nail samples treated with NP213 compared to vehicle (Mann-Whitney test following log transformation of data).
A

Dermatophytes recovered (log(CFU/ml))

Vehicle (placebo) 10% (w/v) NP213

****

B

Dermatophytes recovered (log (mean CFU/ml & 95% CI))

| Infected nail ID | Vehicle       | 10% (w/v) NP213 | p value compared to vehicle |
|------------------|---------------|-----------------|----------------------------|
| 6:169            | 3.98 (3.37, 4.55) | 0.0 (0.0, 0.0) | 0.0022                     |
| 5:152            | 4.36 (3.67, 5.04) | 0.0 (0.0, 0.0) | 0.0022                     |
| 9:226            | 3.63 (3.52, 3.73) | 0.0 (0.0, 0.0) | 0.0022                     |
| 8:212            | 7.30 (7.30, 7.30) | 0.0 (0.0, 0.0) | 0.0022                     |
| 12:281           | 4.30 (3.84, 4.76) | 0.0 (0.0, 0.0) | 0.0022                     |
| 3:89             | 7.02 (6.89, 7.15) | 0.0 (0.0, 0.0) | 0.0022                     |
| 4:101            | 5.01 (4.74, 5.27) | 0.0 (0.0, 0.0) | 0.0119                     |
| 4:111            | 6.62 (6.52, 6.71) | 0.0 (0.0, 0.0) | 0.0022                     |