Journal of Materials Chemistry B



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Cite this: J. Mater. Chem. B, 2024, **12**, 8966

Received 16th February 2024, Accepted 15th July 2024

DOI: 10.1039/d4tb00319e

rsc.li/materials-b

1. Introduction

Protein and peptide materials have become very popular over the last decade for a wide variety of biological uses and beyond, thanks to their inherent bioactivity, cytocompatibility, and biodegradability.^{1–3} In particular, those containing antimicrobial peptides (AMPs) are highly attractive in light of the emergence of antimicrobial resistance against traditional antibiotics, although their design entails a set of challenges to maintain their bioactivity.^{4,5} Recombinant fusion biotechnology has emerged as

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Materials derived from the human elastin-like polypeptide fusion with an antimicrobial peptide strongly promote cell adhesion[†]

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Protein and peptide materials have attracted great interest in recent years, especially for biological applications, in light of their possibility to easily encode bioactivity whilst maintaining cytocompatibility and biodegradability. Heterologous recombinant expression to produce antimicrobial peptides is increasingly considered a convenient alternative for the transition from conventional methods to more sustainable production systems. The human elastin-like polypeptide (HELP) has proven to be a valuable fusion carrier, and due to its cutting-edge properties, biomimetic materials with antimicrobial capacity have been successfully developed. In this work, we have taken advantage of this platform to produce a difficult-to-synthesise sequence as that of the human β -defensin 1 (hBD1), an amphipathic cationic peptide with structural folding constraints relevant to its bioactivity. In the design of the gene, highly specific endoproteinases recognition sites were introduced to release the active forms of hBD1. After the expression and purification of the new fusion construct, its biological activity was evaluated. It was found that both the fusion biopolymer and the released active forms can inhibit the growth of Escherichia coli in redox environments. Remarkably, 2D and 3D materials derived from the biopolymer showed a strong cell adhesion-promoting activity. These results suggest that HELP represents a multitasking platform that not only facilitates the production of bioactive domains and derived materials but could also pave the way for the development of new approaches to study biological interactions at the molecular level.

a promising platform to produce antimicrobial peptides. Standard chemical synthesis encounters several burdens that lead to increased production timelines and costs. These challenges include the hurdle of synthesising difficult sequences or lengthy peptides, inefficient purification processes due to the primary structure of the AMPs, and the need for the proper folding (*e.g.*, the occurrence of disulfide bridges, post-translational modifications, *etc.*) to maintain the antimicrobial efficacy.^{6,7} In contrast, the recombinant production in *Escherichia coli* offers cost-effectiveness, scalability, and efficient expression systems, resulting in high yields and establishing the basis for large-scale protein production.⁸ To date, many fusion carriers have been developed and used for AMP production. Nevertheless, the elastin-like polypeptides (ELPs) are still the least exploited.⁹

ELPs are a class of recombinant biopolymers inspired by elastin and therefore endowed with the inverse transition property, a thermo-responsive behaviour that allows these components to shift from the solvated form to the suspension state, depending on the temperature of the environment.¹⁰

The human elastin-like polypeptide (HELP) fusion carrier developed in our laboratory differs from most other ELPs

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[†] Electronic supplementary information (ESI) available: HhBD1 cloning and purification; stability of HELP and HhBD1; hBD1 accessible surface area calculation, radial diffusion assays; oscillatory rheology analysis; mass spectrometry; fluorescence analyses of cell cultured on coatings and matrices. See DOI: https:// doi.org/10.1039/d4tb00319e

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described in the literature as it possesses the cross-linking domains characteristic of tropoelastin.¹¹ These domains can be cross-linked by an enzymatic reaction, resulting in the formation of cytocompatible hydrogel matrices. This mild process does not affect cell viability, allowing its use for cell encapsulation and culture.¹² Notably, the cross-linking domains are also the main target for elastase, an enzyme secreted by activated polymorphonuclear leukocytes during inflammation.¹³ Consequently, the presence of elastolytic proteases of different origins can trigger the release of any domain fused to HELP, as well as any compound embedded in the gel matrix.¹⁴ Recently, a HELP-AMP fusion with the antimicrobial peptide indolicidin has been described. In addition to the successful high-yield expression, the HELP-AMP fusion product retained both the HELP thermo-responsive properties and the AMP activity, demonstrating the advantage of using this carrier for the production of AMP domains.15

The human β -defensin 1 (hBD1) belongs to the defensions family, a class of small, cationic antimicrobial peptides with a well-defined tertiary structure stabilised by three specific disulfide bridges that are conserved among all β-defensins.¹⁶ hBD1 is constitutively expressed and produced by several types of epithelial cells, such as those of the urinary tract, pancreatic duct, respiratory tract, and intestine, as well as by keratinocytes.¹⁷ Recent evidence suggests that the members of the β -defensin family are multifunctional peptides that not only have functions related to host defence,^{18,19} but are also involved in cell proliferation and migration,²⁰ wound healing, tissue regeneration,^{21,22} and tumour inhibition,²³ positioning them as attractive components for translational applications. hBD1 plays a critical role in the innate immune system's defence against microbial infections and constitutes one of the body's first lines of defence against pathogens. It is well-known for its broad-spectrum antimicrobial activity against various bacteria, fungi, and viruses.²⁴ However, the length of the sequence, the presence of six cysteine residues essential for the correct folding, as well as the need to increase hBD1 availability to produce multifunctional and biomimetic materials are the main challenges for its production by chemical synthesis. This has led us to employ the HELP carrier for the recombinant expression of hBD1 as an alternative and more sustainable route for the synthesis of this peptide.

This work describes the design, the production, the biochemical characterisation of the HELP-hBD1 fusion biopolymer (HhBD1) and the derived hBD1 peptides, as well as the biological assessment of their activity towards model microorganisms and cell lines. Derived materials such as coatings and matrices based on this new fusion biopolymer are evaluated for their interaction with biological systems.

2. Results and discussion

2.1 Production and characterisation of the HELP recombinant fusion with hBD1

hBD1 was employed to functionalise HELP to develop biomimetic components with antimicrobial properties. It has been

described that this AMP is physiologically synthesised as a 68residue prepropeptide (GenBank: CAA63405.1), the product of the *DEFB1* gene.²⁵ However, this product is further processed to originate functional domains ranging from 36 to 47 amino acids, which differ from each other by N-terminal truncation.²⁶ Two major fragments endowed with antimicrobial activity have been described; one of 47 amino acids, 27,28 and another one of 36 amino acids, which is described as the most active domain against bacterial cells.^{29,30} For this reason, the fragment of 47 amino acids was selected as the fusion partner at the Cterminus of the HELP carrier (Fig. 1). Remarkably, this last sequence does not contain any acidic residue.¹¹ Thus, the unique aspartic acid at the beginning of the 36-residue hBD1 domain represents a specific target for the Asp-N endoproteinase, allowing the release of this active peptide from the recombinant fusion biopolymer. Following the same strategy, the addition of a triplet encoding a glutamic acid just before the open reading frame of the 47-residue hBD1 domain resulted in the presence of a second specific cleavage site, in this case for the Glu-C endoproteinase, as schematically shown in Fig. 1.

After cloning, the fusion biopolymer, named HhBD1 (HELPhBD1, M_W 50694.45 Da), was expressed and produced in *E. coli* following a recombinant approach and purified by the inverse transition cycle, taking advantage of the thermo-responsive behaviour of the elastin domain, which is well described for the purification of ELP and ELP-based biopolymers (Fig. S1, ESI†).^{33,34} On average, more than 250 mg of pure HhBD1 were obtained per litre of bacterial culture using our lab-scale production system, which is in agreement with yields reported for other ELP-AMP fusions.³⁵ Electrophoretic analysis of the



EGNFLTGLGHRSDHYNCVSSGGQCLYSACPIFTKIQGTCYRGKAKCCK

Fig. 1 Schematic representation of the recombinant HhBD1 fusion biopolymer. The hBD1 sequence (grey-blue, 47 amino acids) is fused at the C-terminus of HELP (black, his-tag; green, cross-linking domains; red, hydrophobic domains of the human elastin). The hBD1 sequence is flanked by a glutamic acid residue, the specific proteolytic cleavage site for Glu-C (purple scissors). The 36 amino acids domain (blue) can be obtained by cleavage with the Asp-N specific protease (pink scissors). The image of the hBD1 fusion domain was generated with Mol*Viewer³¹ from the PDB entry 1E4S.³² pure HhBD1 (Fig. S1, lane 3, ESI⁺) showed the presence of additional bands that exhibited slower migration than that expected for the HhBD1 monomer, suggesting the presence of multimeric HhBD1 forms due to the formation of interchain disulfide bridges given that the primary structure of hBD1 contains six cysteine residues. The stability of the fusion protein at different pH values (see ESI,† Table S1) and the susceptibility to proteolysis by elastase were tested, and results are reported in Fig. S2 and S3 (ESI⁺). The HELP biopolymer was unaffected by both acidic and basic conditions. Intriguingly, prolonged incubation of HhBD1 in basic conditions resulted in the formation of a hydrogel-like layer on the bottom of the recipient with the consequent decrease in the concentration of HhBD1 in the supernatant.

As described above, the primary sequence of the HhBD1 biopolymer contains unique glutamic and aspartic acid residues, which are the recognition sites for highly specific endoproteinases. Therefore, the purified HhBD1 was treated with Glu-C and Asp-N to verify the release of the two hBD1 fragments (5069.42 Da and 3934.57 Da, respectively). The electrophoretic analysis (Fig. 2) showed that after the endoproteolytic reactions, most of the HhBD1 signal disappeared, accompanied by the formation of two new bands. The most prominent one migrated at the same level of the HELP band (about 50 kDa) and the other one ran along with the electrophoretic front (Fig. 2, boxed in purple), suggesting that the cleavage converted most of HhBD1 into HELP and hBD1. Both endoproteolytic reactions were performed on HELP as control and no effect on this biopolymer was observed, as expected (Fig. 2).

Turbidimetry analysis of the pure fusion biopolymer showed that, although no temperature-dependent transition

HIBD HED + ASY HED HIBD HED X CHUC

weth Protocol

■ 66 kDa

∎45 kDa

30 kDa

14 kDa

Fig. 2 Representative image of the 10% SDS-PAGE analysis of Glu-C and Asp-N specific cleavage of HhBD1 and HELP biopolymers. The electrophoretic signals corresponding to HhBD1 (blue arrow), HELP (red arrow), and the released hBD1 domains (boxed in purple) are indicated. Molecular mass markers (MM): bovine serum albumin, 66 kDa; ovalbumin, 45 kDa; carbonic anhydrase, 30 kDa and lysozyme, 14 kDa. Coomassie blue staining

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Fig. 3 Turbidimetric analysis of the HELP and HhBD1 biopolymers at 2 mg mL⁻¹ as a function of temperature. HELP (red symbols) and HhBD1 (blue symbols) were dissolved in 10 mM Tris (open symbols) and 10 mM Tris/0.15 mM NaCl (filled symbols) buffers, pH 8.

was observed when the biopolymer was dissolved in buffer alone (Fig. 3, blue open symbols), in the presence of a nearly physiological concentration of NaCl, HhBD1 showed a sharper transition at a lower temperature (T_t 28 °C) compared to HELP under the same conditions (T_t 37.3 °C) (Fig. 3, compare red filled with blue filled symbols). This effect was already observed when HELP was fused with indolicidin.¹⁵ It has been described that the fused domains, when folded, may act providing a surface in close proximity to the carrier and that the surface properties, rather than the overall hydrophobicity of the domain, play a dominant role in modulating the Tt of elastinlike carriers.³⁶ hBD1 exhibits an amphipathic conformation endowed with a predominantly apolar surface area (see ESI,† Table S2). This is consistent with the observed thermoresponsive behaviour of this HELP fusion biopolymer, which exhibits a lower Tt than the HELP carrier alone.

These data showed the successful expression of the HhBD1 fusion biopolymer, which retained the thermo-responsive behaviour of HELP and displayed the expected cleavage pattern established in the design of the synthetic gene.

Biological activity evaluation of the HhBD1 biopolymer 2.2

After the production of purified HhBD1 exploiting the thermoresponsive behaviour of the HELP carrier, the first concern was to verify whether the fusion biopolymer retained the antimicrobial properties of the hBD1 domain. Although it was reported that the minimum inhibitory concentration of hBD1 (MIC) against Staphylococcus aureus and E. coli strains was 4-8 mg L^{-1} and 16-32 mg L^{-1} , respectively,³⁰ we could not detect any activity of HhBD1 following this method. Many reasons withstand behind this observation, especially the uncontrolled redox state of the fusion biopolymer in which the disulfide bridge formation is uncontrolled and unspecific and cannot be blocked due to the recombinant nature of HhBD1, and the impossibility of performing MIC assays adding reducing agents to the media, as these are toxic to the bacterial

HhBD1

HELP

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cells in solution. It has been described that the presence of cysteines is crucial for the antimicrobial activity of hBD1, especially the C-terminal cysteines, and that its antimicrobial capacity is closely related to the reduction of the disulfide bridges, and thus it can be evidenced in reducing environments.²⁹ Following a modification of the radial diffusion assay described by these authors, in the presence of 2 mM DTT, we were able to detect the antimicrobial activity for the HhBD1 fusion biopolymer towards E. coli ATCC 25922 strain when 100 µg of HhBD1 were deposited (containing approximately 10 µg of hBD1). In contrast, no inhibition halos were observed in the absence of the reducing agent (see ESI,† Fig. S4). This activity was also retained after the cleavage of HhBD1 with Glu-C and Asp-N, and no significant differences were observed when comparing the diameters of the inhibition zone of the digestion mixtures with those of the fusion biopolymer (Fig. 4). HELP alone did not show any antimicrobial activity in any of the tested conditions (see ESI,† Fig. S1B and C), confirming that the antimicrobial activity of the HhBD1 fusion biopolymer is due to the presence of the hBD1 domain.

The effect of HhBD1 on the viability of eukaryotic cells was tested using the WST-1 viability assay, in which the metabolic activity of osteoblastic (Fig. 5(A)) and fibroblastic (Fig. 5(B)) cells in the presence of different concentrations of HhBD1 and HELP was assessed. The viability assays did not evidence cytotoxicity on the tested cell lines. Rather, as already reported for HELP,³⁷ they revealed a moderate pro-proliferative effect of both biopolymers.

The evaluation of the biological activity of the HhBD1 biopolymer revealed that the fusion construct retained the antimicrobial activity, depending on the redox condition of the environment, as already described for hBD1. Furthermore, no cytotoxic effects on eukaryotic cells were detected at any of the concentrations tested (up to 500 μ g mL⁻¹).

Taken together, these data point to HhBD1 as a noncytotoxic candidate component to produce biomaterials and composites endowed with antimicrobial activity that can be specifically triggered in a reducing environment.



Fig. 4 Antimicrobial activity of HhBD1. (A) Radial diffusion assay performed in the presence of 2 mM DTT by incubating *E. coli* ATCC 25922 strain with the untreated HhBD1 biopolymer and with HhBD1 cleaved with Glu-C and Asp-N. Lysozyme was used as the positive control. (B) The diameter of the inhibition zones was measured and statistically analysed using a one-way ANOVA test. No significant differences (n.s.) were found with p < 0.05.



Fig. 5 Viability of (A) MG-63 osteoblast and (B) NIH-3T3 fibroblast cell lines treated with HELP (red bars) and HhBD1 (blue bars) biopolymers at different concentrations. Values were normalised to the untreated control culture (black dotted lines).

2.3 HhBD1-based 3D matrix production and characterisation

The presence of cross-linking domains in the HELP carrier has been demonstrated as a valuable prerequisite for preparing hydrogel matrices endowed with biological functionality.¹⁵ Following an already described enzymatic method,¹² the matrix-forming capacity of HhBD1 was explored and compared with that of HELP. 4% (w/v) aqueous solutions of HhBD1 in the presence of transglutaminase showed a capacity to form a porous, spongy, hydrogel-like matrix indistinguishable from that of HELP, as shown by scanning electron microscopy (SEM) analysis (Fig. 6(A)).

Oscillatory rheology was performed on 4% biopolymer solutions containing 2 $\mu g m L^{-1}$ transglutaminase to follow the kinetics of the cross-linking reaction (Fig. 6). Although the gelation point is similar for the HELP and HhBD1 systems, the storage modulus of HhBD1 is anticipated compared to HELP, indicating faster enzyme kinetics (Fig. 6(B)). This could be due to the presence of additional glutamine and several lysine residues in the hBD1 domain which may be randomly involved in the cross-linking reaction, or higher solvent exposure of the cross-linking sites for HhBD1 relative to HELP. Both samples displayed elastic (G') and viscous (G'') moduli reaching a plateau after about 90 minutes, resulting in very similar storage moduli (1.0 \pm 0.1 kPa) for both matrices. This fact indicates that the HELP cross-linking domains in both biopolymers are the main target of the transglutaminase rather than the hBD1 domain. It is worth noting that the presence of the hBD1 domain lowered the Tt of the HhBD1 fusion biopolymer (Fig. 2(B)), likely favouring the proximity of the HhBD1 chains and resulting in a more efficient cross-linking process, which is in agreement with the faster gelation kinetics. Both HELP and HhBD1 samples formed strong gels, with elastic moduli that were over two orders of magnitude higher than the viscous moduli. The stress sweep analyses supported the main findings of the time sweep tests, with both gels showing a remarkably wide linear viscoelastic range (up to nearly 1 kPa) and with the fracture point of the HELP matrix being slightly anticipated at 1.2 ± 0.3 kPa (Fig. 6(C), red arrow) compared to

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Fig. 6 4% HELP and HhBD1 hydrogel matrices. (A) SEM analysis of the freeze-dried elastin-based hydrogel matrices. The bar is 40 μ m. (B) Oscillatory rheological time sweep and (C) frequency sweep analyses of HELP (red) and HhBD1 (blue) matrices. *G'*, elastic or storage modulus; *G''*, viscous or loss modulus.

that of the HhBD1 gel at 1.8 \pm 0.5 kPa (Fig. 6(C), blue arrow), suggesting more cross-linking points in the latter. The stability and nature of the hydrogels were further confirmed by frequency sweep analyses, which showed that the storage and loss moduli of both biopolymers were not affected by the applied frequency (see ESI,† Fig. S5). In summary, SEM and oscillatory rheology analyses evidenced that the HELP and HhBD1 matrices are comparable, although the addition of the antimicrobial domain slightly increased the cross-linking points and strength of the HhBD1 matrix.

The ability of HhBD1 to form a matrix was used as a convenient method to verify the specificity of the endoproteolytic cleavage by Glu-C and Asp-N. As described in the ESI,[†] each enzymatic reaction was performed on the matrix to cleave the fragments, which became soluble and were then recovered in the supernatant. The supernatant from each reaction was analysed by ESI-MS without further purification. The masses detected were consistent with those calculated for the two expected fragments, confirming the correctness of the amino acid sequence of the peptides and the high specificity of the reactions (see ESI,[†] Fig. S6A and B for Glu-C and Asp-N released fragments, respectively). Furthermore, these results confirmed the versatility of HELP fusion and the derived matrix to produce bioactive peptides.

2.4 Cell culture on HhBD1-derived substrates

2.4.1 Cell culture on HhBD1-based coatings. Components of the extracellular matrix send signals to cells and influence their ability to survive, divide, and exhibit certain developmental phenotypes. The cytocompatibility of the HELP and its derived materials has already been documented.^{12,38} HELP has never shown cytotoxicity towards the tested cells; however, despite supporting cell adhesion, it does not accelerate it.³⁹ Previous results demonstrated a high cell viability of osteoblast

and fibroblast cells in the presence of HhBD1 in the culture media (Fig. 5), thus, we decided to study its effect on cell adhesion and spreading. To evaluate the cell behaviour towards the HhBD1 fusion biopolymer, we prepared different surfaces based on this new construct to compare its effect with that of HELP. To analyse the cell response, HhBD1 and HELP were adsorbed on a substrate that does not support cell culture like the non-treated polystyrene, which here is referred to as NP (see also Experimental Section 4.4). Both MG-63 osteoblast and NIH-3T3 fibroblast cell lines were cultured on the coated and uncoated NP surfaces. Cells were also cultured on the standard tissue-plastic (TP) surface as the 100% adhesion control. Unexpectedly, the observation by optical microscopy showed that HhBD1 coating on NP significantly enhanced osteoblast and fibroblast cell adhesion, at the difference of the HELP coating on NP (Fig. 7(A) and 8(A)). This was confirmed by assessing both cell adhesion and viability of the two cell lines. The adhesion assay on the osteoblast cells showed that coating the NP surface with HELP only slightly improved cell adhesion (less than 20%), while the NP coating with HhBD1 increased adhesion to approximately 60%, demonstrating a remarkable ability to promote cell attachment (Fig. 7(B)). The viability assay showed that the cells that adhered to these surfaces were metabolically active (Fig. 7(C)). A similar effect was observed for the fibroblastic cell cultures. Approximately 90% of the cells remained attached to the HhBD1-coated NP surface (Fig. 8(B)), being metabolically active (Fig. 8(C)). For both cell lines, these assays showed that although the HELP coatings were able to slightly improve cell adhesion respect to the untreated NP surface, the HhBD1-coated surfaces demonstrated a higher capacity to enhance cell adhesion.

To further confirm these observations, thin-film coatings on NP were prepared with HELP and with HhBD1 to ensure a



Fig. 7 HhBD1 promoted the adhesion of osteoblastic cells. (A) Representative contrast phase microscopy images of MG-63 cultures on HELP- and HhBD1-coated NP surfaces. Cells cultured on untreated NP and TP (boxed) surfaces were the negative and positive controls for cell adhesion, respectively. (B) Adhesion assay using crystal violet staining and (C) viability assay of the attached cells. Values were normalised to the control cell culture on the TP surface and statistically evaluated using a one-way ANOVA test, ***p < 0.001. The bar is 200 nm.



Fig. 8 HhBD1 promotes the adhesion of fibroblastic cells. (A) Representative contrast phase microscopy images of NIH-3T3 cultures on HELPand HhBD1-coated NP surfaces. Cells cultured on untreated NP and TP (boxed) surfaces were the negative and positive controls for cell adhesion, respectively. (B) Adhesion assay using crystal violet staining and (C) viability assay of the attached cells. Values were normalised to the control cell culture on the TP surface and statistically evaluated using a one-way ANOVA test, ***p < 0.001. The bar is 200 nm.

controlled coverage of the cell seeding surfaces with the two biopolymers (see Experimental Section 4.4). Fluorescence analysis was performed on the two cell lines cultured on the thinfilms (Fig. 9(A) and (B)). To better highlight the pro-adhesive effect, cells were analysed 5 hours after seeding to prevent them from producing their extracellular matrix and starting to adhere to the substrate. Actin filaments were stained with Alexa Fluor 594 phalloidin, and nuclei were counterstained with DAPI. Consistently with previous results, only a few cells showed adhesion to the NP surface and to the HELP thin-film, whereas the HhBD1 thin-film coating strongly promoted cell adhesion to a level comparable to the culture on the TP control surface (Fig. 9(A) and (B)). DAPI staining further evidenced the difference between cultures on HELP and HhBD1 (see ESI,† Fig. S4). These results showed that the presence of the hBD1 domain on the surface has a robust pro-adhesive effect. In both cell lines, the morphology of the cells cultured on HhBD1 thinfilms resembled those cultured on the control TP surface (Fig. 9). The cells cultured on the HhBD1 substrate showed highly organised actin meshwork. A slightly lower degree of spreading with respect to the cells cultured on control TP was observed. However, this could be related to irregularities on the surface of the thin-film as well as an uneven distribution of hBD1 on the surface. The cells cultured on HELP showed a different morphology, remaining rounded and showing a tendency to aggregate rather than spread, resembling the cultures on the untreated NP substrate. The DAPI staining emphasised the difference between the cultures on HELP and HhBD1 (see ESI,† Fig. S7).

2.4.2 Cell culture on HhBD1-based 3D matrix. The matrix-forming capacity of HELP could provide further evidence for the observed adhesion capacity conferred by the presence of the



Fig. 9 Fluorescence microscopy analysis of (A) MG-63 osteoblast and (B) NIH-3T3 fibroblast cell lines grown on HhBD1 and HELP thin-film NP coated surfaces. F-actin was labelled with Alexa Fluor 594 phalloidin, and nuclei were counterstained with DAPI. The bar is 50 µm.

hBD1 domain. It was observed that although the HELP matrix is not cytotoxic and can be successfully used to encapsulate cells, proliferation was delayed, and cultures resulted in the formation of islets rather than monolayers.¹² However, it has also been reported that the fusion of adhesion signals to the HELP promotes cell adhesion on the derived hydrogel matrix.⁴⁰ To verify whether the hydrogel network derived from HhBD1 could support cell adhesion and growth, 4% HhBD1 matrices were prepared. Before seeding, extensive washing with water was performed to prevent the presence of the enzyme and of the non-crosslinked biopolymer. 24 hours after seeding, the cells cultured on these matrices were stained with toluidine blue (Fig. 10), and the effect of the presence of the hBD1 domain in the matrix became even more evident, especially when the two cultures on HELP and HhBD1 matrices were compared (Fig. 10(A) and Fig. S8A, ESI⁺). As previously observed,¹² cells seeded on HELP matrices remained round and aggregated, whereas cells cultured on the HhBD1 matrices adopted the morphology already observed for the cultures on TP and on the



Fig. 10 HhBD1 hydrogel matrices supported cell adhesion of osteoblast and fibroblast. (A) Representative contrast phase images of cells cultured on the hydrogel matrices and stained with toluidine blue. (B) Viability assay of cells cultured on the hydrogel matrices. Values were normalised to the TP cell growth control and statistically evaluated using a one-way ANOVA test, ***p < 0.001. The bar is 200 nm.

HhBD1 NP coated surfaces. To evaluate the cell adhesion capacity of the matrices, a WST-1 viability assay was employed as an indirect measurement of the number of cells able to attach to the matrices. Thus, before the assay, the samples were washed to remove the cells that were not attached to the matrices. The results further confirmed the remarkable difference between the two culture substrates (Fig. 10(B)).

Fluorescence microscopy analysis was carried out to examine the morphology and cytoarchitecture of the cells cultured on these matrices (Fig. 11 and Fig. S8, ESI[†]). Only few cell nuclei were visualised on the HELP matrix (Fig. S8B, ESI[†]), whereas on the HhBD1 matrix, the presence of a high number of stained nuclei was evident, indicating a robust cell adhesion of both osteoblast and fibroblast cells (Fig. S8B, ESI[†]). Both cell lines cultured on the HhBD1 matrices exhibited cells with a well-organised actin meshwork (Fig. 11), albeit cell dimensions appeared reduced to some extent compared to cells cultured on the standard TP surface (Fig. 9). This slight reduction in spreading may be due to the heterogeneous and irregular surface of the matrix as well as to the different surface stiffness respect that of the TP surface. However, cells were firmly attached and displayed a properly shaped cytoskeleton (Fig. 11).

These data, consistent with previous observations, confirmed that the presence of the hBD1 domain conferred to



Fig. 11 Fluorescence microscopy analysis of MG-63 and NIH-3T3 cells cultured on HhBD1 hydrogel matrices. F-actin was labelled with Alexa Fluor 594 phalloidin, and nuclei were counterstained with DAPI. The bar is 50 μ m.

the HELP matrix a strong signalling effect for cell attachment towards both the tested lines. Interestingly, in contrast to the antimicrobial activity, the pro-adhesive capacity toward eukarvotic cells did not require reducing environmental conditions, suggesting that the peptide conformation may be a crucial factor in modulating its bioactivity. Coatings, thin-films, and hydrogel matrices derived from HhBD1 supported the osteoblasts and fibroblasts adhesion. Overall, these data revealed a pro-adhesive activity of the hBD1 domain, which, to our knowledge, is described for the first time. Although it is known that hBD1, like other β -defensins, is a multifunctional factor,^{24,41} a pro-adhesive activity has so far only been demonstrated for hBD5.⁴² Notably, it has recently been described that hBD1 can specifically bind to the pore region of the Kv1.3 potassium channel, leading to conformational changes while retaining the activation properties of the channel itself.⁴³ There is now plenty of evidence that different classes of K⁺ channels play a role in integrin-dependent adhesion.⁴⁴ It has been demonstrated that these channels can be physically associated with integrins, such that the conformational change of the channel in turn affects the structure of the adjacent integrin subunit, leading to activation of the latter and cell adhesion.45 Interestingly, these channels are reported to be involved in the regulation of cell size in non-excitable tissues.46 Our results are consistent with these findings and may even point to a possible unexplored mechanism related to cell adhesion.

3. Conclusions

Due to its length and conformation, the hBD1 is considered a difficult-to-synthesise sequence. The HELP carrier has been successfully employed as an alternative and more sustainable system to produce large amounts of this antimicrobial peptide with respect to chemical synthesis. The HhBD1 fusion biopolymer was produced and characterised, verifying the antimicrobial domain(s) release capacity achieved by the appropriate design of the synthetic gene. The analyses confirmed the potential of the new biopolymer, which retained both the thermo-responsive properties and the antimicrobial capacity, holding great promise to produce biocompatible and multifunctional materials. Furthermore, enzymatic crosslinking yielded remarkably strong and stable hydrogels from HhBD1. Unexpectedly, a strong cell adhesion-promoting activity was detected for the surfaces treated with the HhBD1 biopolymer as well as for the derived hydrogel matrix. This finding is consistent with recent data that indicate that hBD1 is a multifaceted factor with still unveiled biological functions and thus harbouring promise for a variety of clinical applications. Despite the development of countless strategies and materials for surface engineering, achieving efficient cell adhesion still represents a challenge. Our data indicates that the novel HhBD1 biopolymer is a versatile component for the fabrication of coatings and scaffolds that promote cell adhesion and that can be safely integrated into biological systems due to their biotic origin. In addition to being an alternative route to produce bioactive

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peptides and an effective way to obtain materials that directly integrate the functional domain, the HELP platform turned out to be a valuable system to study, by a reductionist approach, the biological interactions that are still unknown or not fully elucidated at the molecular level. The opportunity to embed a functional peptide in a biomimetic moiety, the availability of a suitable inactive control, as well as the possibility to choose between different configurations, such as the liquid solution and 2D or 3D material settings, are the main features of a powerful tool to decipher the complexity of biological mechanisms.

4. Experimental

4.1 Production of the recombinant HhBD1 fusion biopolymer

Cloning, expression, and purification. The DNA sequence coding for hBD1 (GenBank: AAB21494.1) was cloned at the C-terminal region of the synthetic HELP gene, exploiting the unique restriction sites. The expression and purification of the recombinant HELP and HhBD1 biopolymers were carried out according to established protocols (see ESI†).¹⁵ The purified biopolymer was analysed by 9% acrylamide SDS-PAGE stained with Coomassie blue as previously described.⁴⁷ The purified product was freeze-dried and stored at -20 °C for further use.

Physico-chemical characterization. Purified biopolymers were dissolved in 10 mM Tris or 10 mM Tris/0.15 mM NaCl buffer pH 8 to a final concentration of 2 mg mL⁻¹. The solutions were equilibrated at 4 °C for 16 hours before analysis. Turbidimetric analysis was performed by measuring absorbance at $\lambda = 350$ nm in a temperature range from 20 to 50 °C at a heating scan rate of 0.2 °C min⁻¹ using a Jenway 6300 spectrophotometer (Hong Kong, China). The inverse transition temperature (T_t) was determined as the temperature at which the absorbance value reached 50% of its peak. For each biopolymer, three replicates were performed, and representative data sets were plotted using GraphPad Prism 10.1.0 (270) software (Boston, USA).

Specific release of hBD1 domains from the HhBD1 fusion biopolymer. The reactions with Glu-C (#P8100S, New England Biolabs, Ipswich, USA) and Asp-N (#55576-49-3, Roche, Basel, Switzerland) enzymes were set up for both HhBD1 and HELP biopolymers at a final concentration of 6 mg mL⁻¹ in 15 μ L of 100 mM ammonium bicarbonate buffer at pH 8. Glu-C reactions were performed at an enzyme concentration of 8.3 ng μ L⁻¹ for 5 hours at room temperature, while Asp-N reactions were carried out at a final concentration of 2.6 ng μ L⁻¹ for 18 hours at 37 °C. After the incubation with the specific endoproteinase, 15 μ L of Laemmli loading buffer were added to each sample to stop the reaction and 3 μ L of this mixture were analysed on 9% SDS-PAGE.

4.2 Antimicrobial and biological evaluation of HhBD1

Radial diffusion assay. Antimicrobial activity was tested for HhBD1 and the products from the reactions with the specific endoproteinase Glu-C and Asp-N described in the previous

section. In parallel, control reactions were performed under the same conditions with the HELP biopolymer. All reagents and biopolymers were sterilised by 0.22 µm filtration. The final reaction volume was 500 µL containing 3 mg of biopolymer. After incubation, the reaction mixture was frozen at -80 °C and then freeze-dried. 60 µL of water were added to each freezedried reaction sample to concentrate the biopolymer to approximately 50 μ g μ L⁻¹ to perform the killing assay. The antimicrobial activity against E. coli ATCC 25922 strain was assessed using a modification of the radial diffusion assay described by Schroeder et al.29 Briefly, a single colony of E. coli from a fresh agar plate was used to inoculate 3 mL of 2.1% (w/v) Mueller-Hinton Broth pH 7.3 (Merck Millipore, Massachusetts, USA). 300 µL of the overnight bacterial culture were diluted in 10 mL of 2.1% (w/v) Mueller-Hinton broth and incubated at 37 °C with continuous shaking (150 rpm) for approximately 2-2.5 hours until an optical density (OD) of approximately 0.5 units was reached. At this point, the bacterial cells were harvested, washed three times with ice-cold NaPi buffer (10 mM sodium phosphate pH 7.3) and then diluted in buffer to 0.1 OD units. For the killing assay, 450 µL of this bacterial solution was mixed with 25 mL of NaPi containing 0.21% (w/v) Mueller-Hinton powder, 1% (w/v) low EEO-agarose (Sigma-Aldrich, Missouri, USA) with or without 2 mM dithiothreitol (DTT, Sigma-Aldrich, Missouri, USA). These mixtures were poured onto 10×10 cm plates (# 82.9923.422, Sarstedt, Numbrecht, Germany) and then cooled to RT to solidify before holes of approximately 2 mm diameter were punched using a glass Pasteur pipette connected to a vacuum pump. The holes were filled with 2 µL of the solutions prepared as described above containing approximately 100 µg of biopolymer. Lysozyme (2 μ g per well) was used as the positive control. The plates were incubated at room temperature for 1 hour to allow the biopolymers to diffuse, and then they were transferred to 37 °C for up to 48 hours. Images were captured, and the diameter of the inhibition zones was measured using ImageJ software.48 The data were statistically analysed using a one-way analysis of variance with n = 6 and p < 0.05.

Cell culture and viability assay. MG-63 and NIH-3T3 cell lines were routinely grown in Dulbecco's modified Eagle's medium (DMEM, Sigma-Aldrich, Missouri, USA) supplemented with 2 mM L-glutamine, 100 µg mL⁻¹ streptomycin, and 100 units mL⁻¹ penicillin and containing 10% (v/v) heatinactivated fetal bovine serum. The cells were maintained in 25 cm² flasks at 37 °C in an atmosphere with saturated humidity and 5% CO2. Cells were seeded in a 96-well tissueculture polystyrene (TP) flat-shaped bottom microplate (Sarstedt, Numbrecht, Germany) at a density of 10^4 cells per cm² in 100 μ L of supplemented DMEM and cultured under standard conditions for 24 hours. Subsequently, the supernatant was replaced with 100 μ L of fresh medium containing 2× serial dilutions starting from 500 μ g mL⁻¹ of HELP and HhBD1, and the cells were further cultured for 24 hours. 5 µL of WST-1 reagent (Roche, Basel, Switzerland) were added per well and incubated at 37 °C for 60 minutes. Absorbance was measured at 450 nm using a microplate reader (Synergy H1, BioTek, Winooski, USA).

4.3 HhBD1 3D matrix production and characterisation

3D matrix production. Hydrogel matrices were prepared following the enzymatic cross-linking method described previously.¹² 4% (w/v) sterile aqueous solutions of HELP and HhBD1 were mixed with microbial transglutaminase (N-Zyme Biotec GmbH, Germany) to a final concentration of 2 μ g μ L⁻¹ and the cross-linking reaction was carried out at room temperature for 1 hour. After the reaction, the matrices were washed with excess water to remove unreacted components and immediately used or stored at 4 °C.

Scanning electron microscopy. 4% HELP and HhBD1 were prepared as described above. After washing with water, they were frozen at -80 °C and freeze-dried. Slices were carefully cut, mounted on aluminium stubs covered with double-sided carbon tape, and sputter-coated with chromium using a Q150T ES plus coater (Quorum Technologies Ltd, UK). Analysis was performed using a scanning electron microscope (Gemini 300, Zeiss, Oberkochen, Germany) operating in secondary electron detection mode. The working distance was 9.3 mm, and the acceleration voltage was 5 kV.

Oscillatory rheological analysis. 225 μ L of 4% (w/v) aqueous solutions of HhBD1 and HELP were mixed, by pipetting upsidedown, with 7.5 μ L of 60 μ g μ L⁻¹ of transglutaminase and transferred to a flat rheometer plate of 20 mm diameter (Malvern Kinexus Ultra Plus Rheometer, Alfatest, Milan, Italy). The rheometer was then lowered to a gap of 0.6 mm, and a time sweep analysis was performed for 90 minutes (1 Hz, 1 Pa, 25 °C with Peltier temperature controller), followed by a frequency sweep (4 Pa) and a stress sweep (1 Hz). The storage (*G'*) and loss (*G''*) moduli of the hydrogels were recorded from 0.1 to 10 Hz (stress 4 Pa, within the linear regime). Each test was performed in at least 2 replicates. For the graphical representation of the time and frequency sweeps, the mean value of two representative data sets was plotted, while a representative data set was used for the stress sweep analysis.

4.4 Cell culture on HhBD1-based substrates

Preparation of the coatings. Coatings were prepared either by adsorption or by deposition of the biopolymers to obtain a thin-film. Coatings by adsorption were done in a sterile 96-well nontreated polystyrene (NP) flat-shaped bottom microplate (Vetrotecnica, Padova, Italy). 100 μ L of 4 mg mL⁻¹ sterile (by 0.2 μ m filtration) aqueous solution of each biopolymer were added per well and incubated overnight at 5 °C. Subsequently, the solution was removed, and the wells were washed two times with 200 μ L sterile water. After the washes, the microplate was air-dried under a sterile hood.

Thin-film coatings (100 µg of biopolymer per cm²) were prepared depositing 10 µl (1.25 µg µL⁻¹) of HhBD1 or HELP sterile aqueous solution (0.22 µm filtration) on the bottom (0.125 cm²) of the microwells of a µ-slide (15-Wells uncoated µ-Slides 3D, #81506, IBIDI, Grafelfing, Germany) and then air-dried under a sterile hood at room temperature.

Adhesion and viability assays of cells cultured on coatings. For cell adhesion assays on HELP and HhBD1 surfaces coated by adsorption, 5000 cells per well were seeded in a final volume of 100 µL of supplemented DMEM. Uncoated tissue-culturetreated (TP) and uncoated NP wells were used as controls. 24 hours after seeding, the cultures were inspected by phase contrast microscopy (Zeiss Primovert contrast phase microscope), and images were acquired using a Zeiss AxioCam 202 mono camera, coupled to Zeiss Zen 3.3 acquisition software (Zeiss, Germany). Before the assays, each well was washed with PBS to remove the unattached cells. Consequently, crystal violet staining and WST-1 cell viability assays were performed on the cells that remained on the surfaces. For the crystal violet assay, the cells were fixed with 50 µL of 100% methanol for 10 minutes on ice. After removing the supernatant, 50 µL of 0.5% crystal violet in 20% methanol were added and incubated for 10 minutes. After extensive washing with water, 50 µL of a 10% acetic acid solution was added to lyse the cells and absorbance was measured at 600 nm. To assess cell viability, 24 hours after seeding the medium was changed and replaced with 100 µL of fresh medium containing 5 µL of WST-1 reagent per well and incubated at 37 °C for 90 minutes. Then, absorbance was measured at 450 nm using a microplate reader.

Cell culture on HELP and HhBD1 matrices. 4% HhBD1 and HELP matrices (w/v) were prepared as described above depositing 10 μ L in the bottom of a well of a μ -Slide (15-Wells IbiTreat μ-Slides 3D, #81506, Ibidi, Grafelfing, Germany). The matrices were cross-linked for 1 hour and then washed extensively with sterile water. 5000 cells per well were seeded in a final volume of 50 µL of supplemented DMEM and cultured for 24 hours. The cultures were then inspected by phase contrast microscopy. The wells were washed with PBS. For toluidine blue staining, cells were fixed by adding 30 µL of 2% (v/v) paraformaldehyde in PBS per well and left at room temperature for 15 minutes. After washing twice with PBS, cells were stained with 10 µL of 0.5% (w/v) toluidine blue in 20% ethanol for 10 minutes. After extensive washing with water, the samples were analysed by phase contrast microscopy, and images were acquired. As an alternative to the crystal violet staining, cell adhesion on matrices was indirectly assessed by the WST-1 assay to evaluate the number of attached cells. After washing with PBS to remove the unattached cells, 50 µL of fresh supplemented DMEM containing 5 µL of WST-1 reagent were added per well. After 120 minutes at 37 °C the supernatant was transferred to a microwell plate, and absorbance was measured at 450 nm using a microplate reader.

Fluorescence analysis. Cell morphology was analysed on the cultures on HELP and HhBD1 thin-film coated surfaces as well as on 4% (w/v) matrices. The thin-film coatings were prepared in the μ -slide as described above, and 5000 cells per well were seeded on these coatings in 50 μ L of supplemented DMEM. These cultures were carried on for 5 hours. For cell morphology analysis on the HELP and HhBD1 matrices, 10 μ L of each 4% (w/v) biopolymer solution were deposited on glass coverslips (#01.4305.19, Vetrotecnica, Padova, Italy), and the cross-linking reaction was carried out as described above. The coverslips were placed in a sterile Petri dish, and 5000 cells in a volume of 20 μ L were seeded on each matrix and incubated for

15 minutes. Then, 15 mL of supplemented DMEM were added to the Petri dish and cultured for 24 hours.

All samples were washed three times with an excess of PBS, and cells were fixed by incubation with 4% (v/v) paraformaldehyde in PBS for 15 minutes at room temperature. Samples were then washed with PBS and blocked for 10 minutes with a solution containing 5% normal goat serum and 0.1% Triton X-100 in PBS. Then, 15 µL of blocking solution containing 4',6-diamidino-2-phenylindole (DAPI; Sigma-Aldrich, Missouri, USA) and Alexa Fluor 594 phalloidin (Molecular Probes, Eugene, USA) at a dilution of 100 ng mL⁻¹ and 0.53 U mL⁻¹, respectively were added to each sample and incubated at 4 °C for 2 hours. After washing three times for 10 minutes with 0.1% Triton X-100 in PBS, the µ-slide lid was sealed, and the coverslips were mounted on glass slides. Fluorescently labelled cells were visualised using a fluorescence microscope (Leica DMLS), and images were acquired with a Leica DFC450 C camera coupled to a Leica LAS v4.13 acquisition software (Leica Microsystems, Wetzlar, Germany). Image cropping, superimposition, and analysis were performed using ImageJ software.48

Statistical analysis. For graphical representation, the values were represented as the mean \pm SD. Unless otherwise indicated, at least three replicas were analysed. One-way analysis of variance (ANOVA) was carried out to compare the means of the different data sets within each experiment.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We thank Prof. Sabina Passamonti for her invaluable support and Dr Marco Stebel, Dr Davide Porrelli, and Dr Fabio Hollan for technical assistance. This work was supported by the H2020 Marie Sklodowska-Curie Action (AIMed project, grant no. 861138), the Horizon Europe STOP project, grant no. 101057961, and the iNEST (Interconnected North-Est Innovation Ecosystem) consortium funded by the European Union NextGenerationEU (PNRR). S.M. acknowledges funding from the Italian Ministry of University and Research through the PRIN program (SHAZAM grant no. 2022XEZK7K) funded by the European Union – Next Generation EU.

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