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Harnessing cyclotides to design and develop novel peptide GPCR ligands

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Cyclotides are plant-derived cyclic, disulfide-rich peptides with a unique cyclic cystine knot topology that confers them with remarkable structural stability and resistance to proteolytic degradation. Recently, cyclotides have emerged as promising scaffold molecules for designing peptide-based therapeutics. Here, we provide examples of how engineering cyclotides using molecular grafting may lead to the development of novel peptide ligands of G protein-coupled receptors (GPCRs), today's most exploited drug targets. Integrating bioactive epitopes into stable cyclotide scaffolds can lead to improved pharmacokinetics and oral activity as well as selectivity and high enzymatic stability. We also discuss and highlight the importance of engineered cyclotides as novel tools to study GPCR signaling.

Introduction

Plants have historically been recognized as a rich source of therapeutic agents and they continue to profoundly inspire drug discovery. In fact, a myriad of today's drugs are plant-derived

natural products or their derivatives.¹ For instance, the sesquiterpene lactone artemisinin, isolated from *Artemisia annua*,² and the indole alkaloids vinblastine and vincristine, derived from *Catharanthus roseus*,³ constitute mainstays of modern malaria and cancer treatment, respectively.

In recent years, a considerable amount of literature has been accumulated, highlighting the potential of natural products as ligands of G protein-coupled receptors (GPCRs).^{4,5} These transmembrane receptors with more than 800 members encoded in the human genome are targeted by approximately 35% of US Food and Drug Administration-approved drugs, thus highlighting their

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druggability.⁶ Upon activation by a diverse array of ligands ranging from ions, and photons, small molecules and lipids to proteins and peptides, GPCRs interact with heterotrimeric G proteins composed of three subunits, G_{α} , G_{β} and G_{γ} , to generate second messengers and initiate cellular signaling cascades.⁷ While the G_{α} subunit modulates adenylyl cyclase, phospholipase C- β , RHO-guanine nucleotide exchange factors or cGMP phosphodiesterase, the $G_{\beta\gamma}$ complex is involved in the regulation of G protein-coupled inwardly rectifying potassium channels, voltage-gated Ca^{2+} channels, mitogen-activated protein kinases and many more downstream effector molecules.⁷ Binding of β -arrestins to activated GPCRs phosphorylated by GPCR kinases results in their uncoupling from heterotrimeric G proteins and targeting to clathrin-coated pits for endocytosis.⁷ Subsequently, GPCRs may encounter early endosomes, from where they either recycle to the cell membrane or undergo translocation for lysosomal degradation.⁸ Owing to their structural diversity and crucial role in a variety of physiological processes, GPCRs constitute the most attractive class of drug targets in modern pharmacology.

The vast majority of natural products acting as ligands of GPCRs are small molecules, but over the past few years nature-derived peptides have gained significant momentum in GPCR ligand discovery and development.^{9,10} Such peptide GPCR ligands have been mainly isolated from venomous animals, bacteria, fungi and plants and they provide a blueprint to produce novel peptidic GPCR ligands.⁹ Over 50 GPCR peptide drugs, most of which are chemically modified peptides, have been approved to date and more than 150 peptides have entered human clinical trials.^{10,11} Of approved drugs, five backbone-cyclic peptides targeting GPCRs have been approved as drugs.¹⁰ Bremelanotide is an example of a backbone-cyclic peptide selectively targeting the melanocortin 4 receptor (MC4R) whereas pasireotide, lanreotide, octreotide and vapreotide are backbone-cyclic peptides modulating the somatostatin GPCRs.¹⁰

In this article, we focus on a class of plant-derived cyclic, disulfide-rich peptides called cyclotides, and shed light on how they can be exploited to develop and engineer novel peptide GPCR ligands with improved pharmacological properties. We highlight the immense potential of plant-derived cyclotides as ligands of GPCRs.

Cyclotides

Structure, biosynthesis and function of native cyclotides

Cyclotides are the largest class of ribosomally synthesized and post-translationally modified plant-derived cyclic peptides.¹² They are characterized by their unique cyclic cystine knot motif comprising a head-to-tail cyclic backbone and a cystine knot formed by six conserved cysteine (Cys) residues.¹³ This highly rigid structure provides them with exceptional stability against thermal, chemical or enzymatic degradation.^{14,15} Biosynthesis of cyclotides from gene-encoded precursors that possess a N-terminal signal peptide, N- and C-terminal flanking leader propeptides with single or multiple core peptide domains involves a cascade of events.^{16,17} Entering the secretory pathway *via* the endoplasmic reticulum enables proper folding of the cyclotide precursors and disulfide bond formation.¹⁸ Post-translational modifications of cyclotides including proteolytic cleavage and subsequent head-to-tail cyclization is thought to take place in the vacuole, leading to production of mature cyclotides.¹⁹ The key enzymes enabling cyclotide processing at the C-terminus and backbone cyclization, are asparaginyl endopeptidases that specifically target an Asn or Asp residue at the C-terminus, and have been elucidated over the last decade.^{20,21} More recently, Rehm and colleagues identified members of the papain-like cysteine proteases as cyclotide N-terminal processing enzymes which play a crucial role in the biosynthetic production of cyclotides.²²



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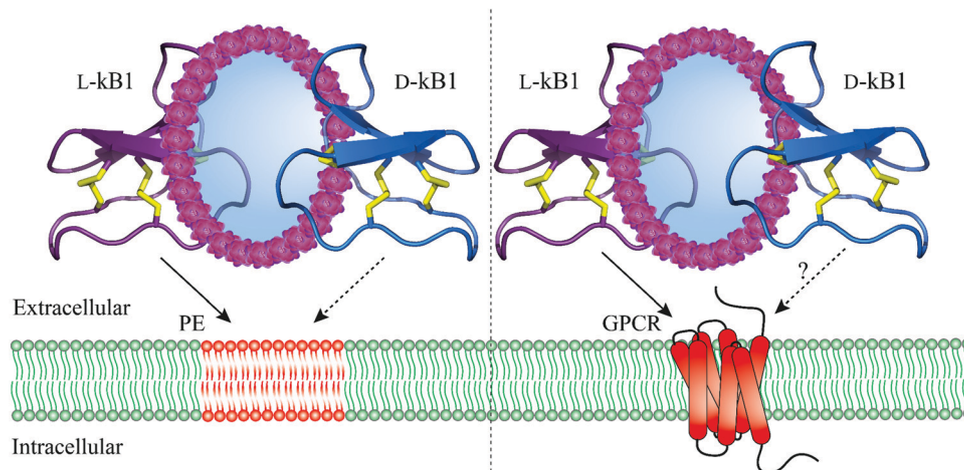


Fig. 2 Molecular targets and mechanism of action of cyclotides. L-Kalata B1 (purple, PDB: 1nb1) strongly interacts with the phosphatidylethanolamine-rich plasma membrane (straight arrow, left) whereas D-kalata B1's (blue, PDB: 2jue) interaction with the phosphatidylethanolamine-rich plasma membrane is significantly reduced (dashed arrow, left). By contrast, L-kalata B1 exhibits affinity towards a GPCR (straight arrow, right) but whether D-kalata B1 shows decreased receptor interaction remains to be determined (dashed arrow, right). Disulfide bonds are shown in yellow and phosphatidylethanolamine-rich plasma membrane and a GPCR are colored in red. GPCR = G protein-coupled receptor; PE = phosphatidylethanolamine; PDB = protein data bank.

lower potency in hemolytic, insecticidal or anti-HIV assays than natural L-kB1.^{34,52} In a recent follow up study, Henriques and colleagues unveiled this chirality dependence by comparing peptide-lipid binding using mirror image model membranes.⁵³ Synthesis of non-natural chiral phospholipids deciphered that the native L-kB1 is able to bind with higher affinity to phospholipids with the same chirality that is observed in eukaryotic membranes, thereby suggesting that the chiral environment of lipid bilayers is capable of modulating the function of membrane-active peptides (Fig. 2).⁵³ In addition to Möbius cyclotides, the interaction of bracelet cyclotides with membranes has been also studied.⁵⁴ In contrast to Möbius cyclotides kB1 and kB2, bracelet cyclotide cycloviolacin O2 was shown to be more active and selective towards anionic membranes containing dioleoylphosphocholine and dioleoylphosphate.⁵⁴ Furthermore, interaction of this cyclotide with membrane lipids of *E. coli* resulted in a selective extraction of PE-containing lipids from the plasma membrane.⁵⁴

The unique interaction of cyclotides with PE confers them with the ability to enter into cells. Mutational studies on fluorescently labeled kB1 analogs revealed that in a MCF-7 breast cancer cell line the cytotoxic mutant [T20K]kB1 exhibits the ability to cross membranes more efficiently than the analog [V25K]kB1 with decreased cytotoxic activity.⁵⁵ Moreover, cellular uptake of labeled [T20K]kB1 resulted in no cell internalization at 4 °C, whereas treatment with a macropinocytosis inhibitor allowed for penetration, suggesting that kB1 enters cells *via* endocytosis.⁵⁵ In a follow up study, the underlying cell penetrating mechanism was elucidated using membrane-active or membrane-inactive kB1 variants labeled with a fluorophore *via* their Lys residue.⁵⁶ [T20K]kB1 was shown to internalize into a HeLa cervical cancer cell line at a nontoxic peptide concentration compared to the inactive mutants [E7K]kB1, [T16K]kB1, and [V25K]kB1 which were impaired in their capability to penetrate

the cells.⁵⁶ Reduced cellular uptake was observed at 4 °C and thus was demonstrated to be due to changes in membrane fluidity and the inhibition of endocytosis which was partially dependent on both dynamin and actin. Overall, these studies suggest that kB1 enters cells either *via* direct translocation or endocytosis by binding to the membrane through the interaction with PE lipids.⁵⁶

While cyclotides primarily interact with PE-rich membranes, GPCRs have been identified as another target (Fig. 2). In the area of cyclotide GPCR drug discovery an appropriate selection of controls used in the pharmacological assays ensures specific interaction of cyclotides with GPCRs, however, their interaction with membrane may represent a limitation for their use in GPCR drug development as this could lead to side effects. A seminal study on discovery of cyclotides as ligands of GPCRs was conducted by Koehbach and colleagues who unveiled the mechanisms underlying the oxytocic activity of cyclotides and their molecular targets.⁵⁷ Bioactivity-guided fractionation and in-depth pharmacological characterization of an *O. affinis* extract identified kB7 as being able to trigger contractions of uterine smooth muscle cells by partially activating human oxytocin and vasopressin V_{1a} receptors in the low μM concentration range (Table 1).⁵⁷ Sequence analysis of kB7 and oxytocin revealed significant homology and thus enabled design of an oxytocin-like peptide with nM affinity and selectivity towards the oxytocin receptor. In a more recent study, Keov *et al.* showed for the first time that kB7 directly interacts with an insect GPCR, the inotocin receptor, of *Tribolium castaneum* and *Lasius niger* by partially agonizing these receptors with an EC₅₀ of 7.2 and 2.2 μM, respectively (Table 1).⁵⁸ The capability of cyclotides to modulate GPCR signaling has recently led to the isolation of several peptides from *Carapichea ipeacacuanha*.⁵⁹ One of these peptides, caripe 8, elicited antagonistic activity at the corticotropin releasing factor type I receptor, a class B GPCR (Table 1).⁵⁹



Table 1 Native and grafted cyclotides as ligands of mammalian GPCRs

Cyclotide	Sequence	Target (MOA)	Affinity/potency (μM)	Ref.
Cyclopsychotride A	<i>cyclo</i> -CGESCVPFICTVTALLGCSCSKSKVCYKNSIP	NTR1 (antagonist)	IC ₅₀ 3	127
Kalata B7 ^a	<i>cyclo</i> -CGETCTLGTCTYQGTCTCSWPICKRNGLPV	OXR, V1AR (partial agonist)	K _i 50/EC ₅₀ 12 K _i 12/EC ₅₀ 4.8	57
Caripe 8	<i>cyclo</i> -CGESCVPFICITAAGCSCKKKVCYRNGVIP	CRFR1 (antagonist)	n.d.	59
ckb-kal	<i>cyclo</i> -CGETCVGGTCTNTPGCTCSWPVCKRPPGFSP	BKR1 (antagonist)	n.d.	76
[GHFRWG;23-28]kB1	<i>cyclo</i> -CGETCVGGTCTNTPGCTCSWPVCGHFRWGV	MC4R (agonist)	K _i 0.029/EC ₅₀ 0.580	89
MCo-CVX-5C ^b	<i>cyclo</i> -CPKILQRCRRSDCPGACICRNGYCGSYRXCRCpRRBCYXKV	CXCR4 (antagonist)	IC ₅₀ 0.019	95
MCo-CVX-6D ^c	<i>cyclo</i> -CPZILQRCRRSDCPGACICRNGYCGSYRXCRCpRRBCYXKV	CXCR4 (antagonist)	IC ₅₀ 0.00015	99
MCo-AT1-7 ^d	<i>cyclo</i> -CPKILQRCRRSDCPGACICRNGYCGSGEIVYRXGGV	MAS1R (agonist)	n.d.	105

GPCRs are named according to UniProt. More details on GPCR nomenclature are available in IUPHAR/BPS Guide to Pharmacology. n.d. refers to not determined ligand's mode of action (MOA). Grafted epitope sequences are highlighted in bold and italics. ^a Kalata B7 has been shown to interact with an insect GPCR, notocin receptor, with an EC₅₀ of 7.2 μM (*Tribolium castaneum*) and 2.2 μM (*Lasius niger*), respectively. ^b B, X, and p denote the amino acid residues 2-naphthylalanine, citrulline, and D-Pro, respectively. ^c B, X, and p represent the amino acid residues 2-naphthylalanine, citrulline, and D-Pro, respectively, while Z indicates 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA). ^d X is L-2,3-diaminopropionic acid residue.

Cutting-edge structural biology techniques and advanced *in silico* methods developed over the past decade have broadened our knowledge of GPCR structure and function.^{6,60} With more than 20 GPCR structures in complex with peptides available, there are ample opportunities for rational, structure-based design of peptides.^{10,61} Moreover, these new technologies are being implemented to perform homology modelling and peptide-receptor interaction studies. For instance, Koehbach *et al.* utilized homology modelling to shed more light on the interaction of cyclotides with the oxytocin receptor.⁵⁷ In line with the data of neurotensin (8–13) interacting with the extracellular surface of the neurotensin receptor,⁶² they revealed interactions of cyclotides with the extracellular loops of the oxytocin receptor and thus possible penetration into the ligand-binding pocket of peptide GPCRs despite their bulky size.^{57,63}

Cyclotides as grafting scaffolds for GPCR ligand design and development

The development of peptide-based GPCR drugs, and peptide drugs in general, has been limited by the typically poor pharmacokinetic profiles of peptides (*e.g.* enzymatic instability,

renal clearance) as well as low oral bioavailability and poor membrane permeability.⁹ Thus, novel strategies to tackle these challenges of peptides as therapeutics are vitally required. In this regard, the structural diversity, exceptional stability and multiple biological activities of cyclotides make them promising starting points for the discovery and development of novel GPCR ligands with unique pharmacological features. The concept of “molecular grafting” which delineates the insertion of a foreign epitope derived from a bioactive peptide or protein into the cyclotide scaffold,⁶⁴ thereby endowing cyclotides with new biological activities, has created much interest in the field of cyclotide research (Fig. 3 and 4). Grafting epitopes onto cyclotides, in particular kB1 (Fig. 3) and *Momordica cochinchinensis* trypsin inhibitor-I (MCoTI-I) (Fig. 4), has been successfully applied to a wide range of molecular targets, including receptors and enzymes;⁶⁵ however, herein we focus on its application and amenability in the context of designing and developing peptide GPCR ligands. The evidence based on the published knowledge of GPCR structures bound to endogenous peptides implies that α -, β -, or γ -turn motifs are the most common patterns of ligand recognition by GPCRs.⁶⁶ Since these stable structural elements govern peptides interactions with GPCRs⁶⁶ and as cyclotides have independently evolved

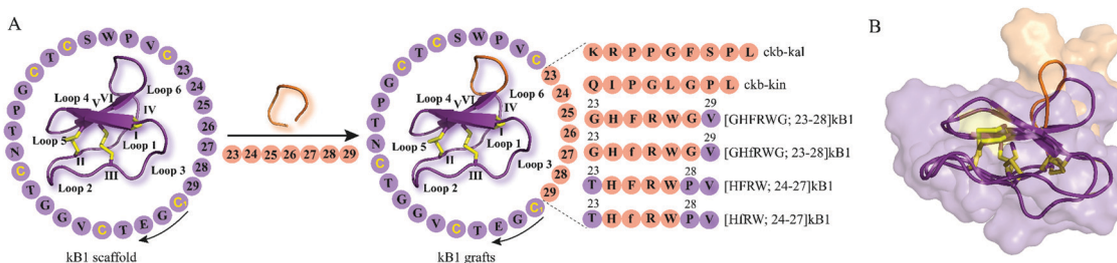


Fig. 3 Examples of kB1 grafts. (A) A bioactive peptide epitope (orange) is inserted into loop 6 of kB1 (PDB: 1nb1) leading to stable peptide GPCR ligands with desired biological activity. Amino acid residues of native kB1 are shown in purple while grafted sequences are indicated in orange. Start and direction of the sequence is highlighted by a cysteine residue and a black arrow, respectively. Sequences and positions of bioactive epitopes grafted in loop 6 of kB1 discussed in this review are shown. Disulfide bonds or cysteine residues are highlighted in yellow and are labeled I–VI. Loops of cyclotides are labeled 1–6. (f) Represents the amino acid residue D-Phe. The starting point of the sequence has been taken from published studies. (B) Cartoon and surface overlay of kB1 and [GHFRWG; 23–28]kB1 (PDB: 2lur) as melanocortin pharmacophore highlighting the only key difference in the grafted loop 6. PDB = protein data bank; kB1 = kalata B1.



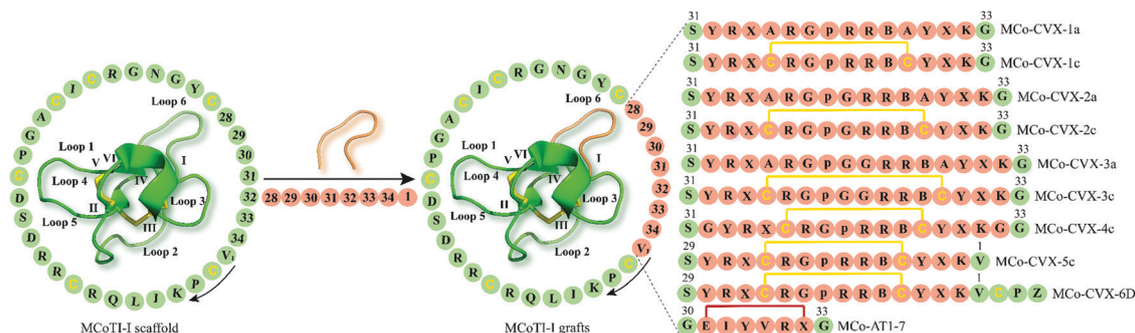


Fig. 4 Examples of cyclotide GPCR ligands grafted onto loop 6 of MCoTI-I scaffold (MCoTI-II NMR structure was used as template, PDB: 1ib9). Amino acid residues of native MCoTI-I are shown in green while grafted sequences are highlighted in orange. Start and direction of the sequence is indicated by a valine residue and a black arrow, respectively. The starting point of the sequence has been taken from published studies. Disulfide bonds or cysteine residues are shown in yellow and are labeled I–VI. B and p indicate the amino acid residues 2-naphthylalanine and D-Pro, respectively, while Z represents 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA). X denotes citrulline residue except for MCo-AT1-7 where X is an L-2,3-diaminopropionic acid residue. The isopeptide bonds formed between the glycine residues and epitope are shown in red. MCoTI-I = *Momordica cochinchinensis* trypsin inhibitor-I; PDB = protein data bank.

these structural features in their framework,⁶⁷ then it is likely cyclotides can not only act as defense molecules in plants but also serve mankind as a valuable source for developing peptide GPCR therapeutics. We now provide examples of how cyclotides, used as drug design scaffolds, can help to increase enzymatic stability, oral activity and selectivity of bioactive epitopes towards their target GPCRs.

Targeting bradykinin receptors by a cyclotide scaffold to treat chronic pain

Chronic pain represents one of today's most challenging issues faced by the health-care sector, with more than 1.9 billion people worldwide suffering from this medical condition.⁶⁸ Peptides derived from venomous animals such as cone-snails, snakes, spiders or scorpions have been shown to be a valuable source for the development of peptide therapeutics to treat chronic pain.^{69,70} Ziconotide, a synthetic version of ω -conotoxin MVIIA (ω -MVIIA) isolated from cone-snail *Conus magus*, is an example of a peptide drug approved for the treatment of chronic pain.⁷¹ However, ziconotide, – a blocker of N-type voltage-sensitive calcium-channels, Ca_v2.2 – develops centrally mediated side effects and thus needs to be intrathecally administered, restricting its use to only a limited number of patients with severe chronic pain.⁷¹ In this context, antagonists of bradykinin receptors, the prototypical GPCRs, have attracted increasing interest as therapeutic agents for the treatment of chronic and inflammatory pain.^{72,73} Bradykinin, kallidin and their des-Arg metabolites (des-Arg⁹-bradykinin and Lys-des-Arg¹⁰-bradykinin), collectively referred to as kinins, are short-lived peptide mediators with a half-life in blood of less than 30 seconds which are generated by virtue of the serine protease kallikrein that proteolytically cleaves the precursor protein kininogen.⁷³ Kinins mediate their pharmacological effects by targeting bradykinin B1 and B2 GPCRs.⁷³ Although the bradykinin B2 receptor is ubiquitously expressed in healthy tissues, the expression of bradykinin B1 receptor is predominantly associated with pathological tissue damage, suggesting that

targeting bradykinin B1 receptors might be a promising approach to managing pain.⁷⁴ Icatibant (HOE 140) is a potent and long-acting peptidomimetic bradykinin B2 receptor antagonist that has been approved to treat hereditary angioedema.⁷⁵ However, as treatment with icatibant requires injection by the parenteral route, developing selective long-lived and orally active bradykinin B1 receptor peptide therapeutics represents an attractive strategy yet with significant challenges.

Wong and colleagues have explored grafting onto cyclotides as a strategy to design and develop novel orally active and selective bradykinin B1 receptor peptide antagonists.⁷⁶ Prompted by previous studies reporting des-Arg¹⁰-[Leu⁹]-kallidin (DALK) as a bradykinin B1 receptor antagonist,⁷⁷ they opted to graft this epitope as well as des-Arg⁹-kinestatin (DAK) onto loop 6 of the kB1 framework which has been previously shown to be amenable to molecular grafting (Fig. 3A).⁷⁸ Accordingly, loop 6 of kB1 (ckb) was substituted with either DALK (kal) or DAK (kin) resulting in two cyclic bradykinin antagonists: ckb-kal and ckb-kin (Fig. 3A).⁷⁶ Additionally, a linear analog of ckb-kin, kb-kin, was designed to corroborate the necessity of the cyclic backbone to engineer stable, orally active bradykinin B1 receptor antagonists. Compared to the control peptide DALK with a half-life of less than five minutes, ckb-kal, ckb-kin and the linear variant kb-kin showed enhanced *in vitro* stability in human serum and lasted longer than six hours.⁷⁶ These data provide evidence that the cystine knot architecture of kB1 may be effective in preventing enzymatic degradation of the designed peptide analogs. The toxicity of engineered cyclic and linear peptides against NIH3T3 cells was assessed, whereby the treatment with linear kb-kin led to about 10% cell death after 48 hours when using concentrations higher than 1 μ M.⁷⁶ Subsequent fluorescence-based functional assays were performed to elucidate the mode of action of the engineered peptides. All three peptides were capable of inhibiting intracellular calcium levels upon receptor activation with a bradykinin B1 receptor agonist des-Arg⁹-bradykinin (Table 1), but no antagonistic effect was observed following cell treatment with the bradykinin B2 receptor agonist bradykinin.⁷⁶ The receptor



subtype selectivity was further corroborated in a competition binding study: all three engineered bradykinin antagonists competitively displaced a fluorescein-des-Arg⁹-bradykinin agonist in HEK293 cells transiently expressing the bradykinin B1 receptor. The analgesic activity of the kB1-based peptides was then evaluated in an *in vivo* model of visceral pain induced by acetic acid thus leading to abdominal constriction (writhing) in mice.⁷⁶ Both intraperitoneal (1 mg kg⁻¹) and oral administration (10 mg kg⁻¹) of grafted cyclotides demonstrated that ckb-kal acts as the most potent antagonist in the abdominal constriction assay followed by ckb-kin and then the linear control kb-kin.⁷⁶ Intriguingly, intraperitoneal treatment with the DALK control resulted in analgesic activity similar to the cyclotide grafts, but no inhibition of writhing frequency was observed after oral application. Two-way analysis of variance on oral activity revealed no significant differences between intraperitoneal and oral administration for ckb-kal and ckb-kin, while there was a significant difference for the linear analogue kb-kin and the control peptide DALK. This study thus underpins the importance of the cyclic backbone of kB1 to retain and augment oral activity.

Exploiting the cyclotide framework to generate stable and selective melanocortin 4 receptor agonists

Obesity is a medical condition imposing a staggering burden on society and public health. Over the past few decades, obesity has reached pandemic proportions, with more than 100 million children and 600 million adults suffering from obesity (body mass index > 30 kg m⁻²) with about 4 million deaths and 120 million disability-adjusted life-years as a consequence of excess body weight.^{79,80} Furthermore, obesity is a high-risk factor for comorbidities, including cardiovascular disease, diabetes mellitus, kidney dysfunction, cancer and musculoskeletal disorders.⁷⁹ Liraglutide, a glucagon-like peptide 1 (GLP-1) analogue, has been recently approved to treat obesity,⁸¹ presumably acting by modulating appetite and body weight *via* activation of the GLP-1 receptor (GLP-1R) within glutamatergic neurons.⁸² In addition to targeting GLP-1R, the melanocortin system emerged as an alternative target to treat obesity⁸³ and comprises five GPCRs, the melanocortin receptors (MCR 1–5). The endogenous peptide agonists include the adrenocorticotrophic hormone and α , β , and γ -melanocyte-stimulating hormones (MSH), which are all produced by proteolytic cleavage of the precursor protein proopiomelanocortin.⁸³ Interestingly, the melanocortin system is unique in that it is the only GPCR system having two endogenous inverse agonists, *i.e.* agouti protein and agouti-related protein.⁸⁴ Whereas MC1R, MC2R and MC5R are mainly expressed in melanocytes, the adrenal cortex and various peripheral tissues, respectively, high expression levels of MC3R and MC4R are detected in the central nervous system.⁸⁵ MC4R is of particular interest as a target for the pharmacological treatment of obesity since activation of this GPCR leads to reduced food intake.⁸⁶ In fact, setmelanotide is a potent MC4R peptide agonist that has completed phase II clinical trials to treat obesity and hyperphagia in patients with genetic defects in the leptin-melanocortin signaling pathways.⁸⁷

Given an unmet clinical need for MCR agonists, Eliassen *et al.* inserted a tetrapeptide sequence (His-Phe-Arg-Trp) of MSH and adrenocorticotrophic hormones that is required to activate the MCRs⁸⁸ into loop 6 of the kB1 scaffold (Fig. 3A).⁸⁹ The tetrapeptide was flanked either by the remaining native amino acids in loop 6, *i.e.* T at the N-terminus and PV at the C-terminus or between G residues which were used instead of the native kB1 residues to prevent possible nonactive conformations of the tetrapeptide owing to direct replacements (Fig. 3A).⁸⁹ In total, four cyclotide grafts were synthesized: [GHFRWG; 23–28]kB1, [GHRfWG; 23–28]kB1, [HFRW; 24–27]kB1 and [HfRW; 24–27]kB1 (Fig. 3A) and were subjected to an in-depth pharmacological characterization. All four peptides were tested in a radioligand binding assay and [GHFRWG; 23–28]kB1 was shown to displace a radiolabeled ligand from the MC4R in a concentration-dependent manner with a K_i of 29 nM (Table 1).⁸⁹ Importantly, [GHFRWG; 23–28]kB1 exhibited 107- and 314-fold increased affinity towards MC4R over the MC1R and MC5R, respectively, while no binding to MC3R was detected at ligand concentrations up to 10 μ M. To assess the ability of [GHFRWG; 23–28]kB1 to activate the MC4R, a functional cAMP assay was carried out. Compared to the endogenous ligand α -MSH with a potency of 3.7 nM, [GHFRWG; 23–28]kB1 was demonstrated to induce the activation of MC4R with an EC₅₀ of 580 nM (Table 1).⁸⁹ These data suggest that incorporation of the tetrapeptide into the kB1 scaffold results in a possible steric hindrance with the receptor and may account for the lower potency of the grafted cyclotide. Structural analysis by nuclear magnetic resonance spectroscopy revealed similar structural elements of [GHFRWG; 23–28]kB1 and kB1 indicating that [GHFRWG; 23–28]kB1 retained a native cyclotide fold (Fig. 3B). Additionally, surface-exposed hydrophobic residues of [GHFRWG; 23–28]kB1 such as His, Phe or Trp were surrounded by polar or charged residues while the Arg residue in loop 6 was shown to be highly solvent-exposed.⁸⁹ Compared to α -MSH, [GHFRWG; 23–28]kB1 exhibited a significantly improved *in vitro* proteolytic stability with more than 80% of the peptide being intact after five hours. This study underscores the great potential of the kB1 scaffold to develop stable and selective agonists targeting MC4R for the treatment of therapeutic indications such as obesity.

Using cyclotide scaffolds to generate antagonists of CXC-motif-chemokine receptor 4

The CXC-motif-chemokine receptor 4 (CXCR4) was initially identified in 1996 as a co-receptor required for the entry of the human immunodeficiency virus type 1 (HIV-1) into host cells.⁹⁰ Along with its endogenous chemokine CXCL12, CXCR4 participates in the control of cellular migration, organogenesis, hematopoiesis, and immune responses.^{91,92} Malfunctioning of CXCR4 is associated with a variety of medical conditions such as cancer, autoimmune diseases and atherosclerosis.^{91–93} Consequently, multiple synthetic CXCR4 ligands have been developed as antagonists of the CXCR4 (*e.g.* plerixafor/AMD3100 is an approved CX4R antagonist).⁹⁴

A decade ago, Aboye *et al.* reported the design and synthesis of a cyclotide-based antagonist of the HIV-1 viral replication



capable of selectively targeting CXCR4.⁹⁵ Using a MCoTI-I scaffold they grafted a circularly permuted version of CVX15 that was designed by fusing the N- and C-termini directly or *via* a flexible Gly_n linker (*n* = 1, 2) and eliminating D-Pro⁸ and Pro⁹ residues, thereby resulting in the new N- and C-termini with Tyr¹⁰ and Lys⁷, respectively (Fig. 4).⁹⁵ CVX15 is a cyclic peptide analog of the horseshoe crab peptide, polyphemusin, that has been previously shown to inhibit HIV by antagonizing CXCR4.⁹⁶ Furthermore, in addition to substitution of Gln at position 6 by citrulline (CVX15 Gln6Cit), native Cys residues of CVX15 forming the disulfide bridge were replaced with Ala residues (Fig. 4). In a CXCR4- β -lactamase U2OS cell-based fluorescence assay, all cyclotide grafts concentration-dependently inhibited CXCR4 activated by endogenous CXCL12, yet with disparate potency ranging from $23.8 \pm 0.3 \mu\text{M}$ (MCo-CVX-3a) to $19 \pm 3 \text{ nM}$ (MCo-CVX-5c) (Table 1).⁹⁵ Compared to the most potent inhibitor MCo-CVX-5c, the peptide CVX15 Gln6Cit showed an EC₅₀ value of $71 \pm 13 \text{ nM}$. Furthermore, cyclotides MCo-CVX-1c and MCo-CVX-3c were approximately 10 times more potent than the cyclotide mutant versions substituted with an Ala residue instead of Cys (MCo-CVX-1a and MCo-CVX-3a), highlighting the importance of Cys residues for biological activity.⁹⁵ Additionally, the length of the Gly linker was critical for the biological activity of the cyclotide grafts. For instance, the MCo-CVX-1c was the most potent antagonist with an EC₅₀ of $0.10 \pm 0.01 \mu\text{M}$, which was designed by linking directly the N- and C-termini of the CVX15 peptide.⁹⁵ Incorporation of additional residues on MCo-CVX-2c and MCo-CVX-3c generated moderately potent molecules with EC₅₀ values of 2 and 3 μM , respectively, suggesting that Gly residues enhance flexibility of the peptide thus affecting the potency by presumably leading to a greater entropic penalty.⁹⁵ Moreover, the position of the epitope in loop 6 was important for the biological activity of the grafted cyclotides. The most active cyclotide was MCo-CVX-5c (EC₅₀ = $19 \pm 3 \text{ nM}$) (Table 1), grafted between Gly residues 30 and 34, while MCo-CVX-1c (grafted at Asn³²) and MCo-CVX-4c (grafted at Asn³² and with additional Gly linker) showed EC₅₀ values of $102 \pm 12 \text{ nM}$ and $39 \pm 1 \text{ nM}$, respectively. Also, MCo-CVX-5c inhibited not only CXCL12-induced Erk phosphorylation and internalization of CXCR4 but also the entry and replication of CXCR4-tropic HIV-1 in human lymphocyte MT4 cells (EC₅₀ = $2.0 \pm 0.3 \text{ nM}$).⁹⁵ Importantly, MCo-CVX-5c exhibited about three times higher potency than raltegravir, an approved drug to treat HIV infection.⁹⁷ Determination of peptide *in vitro* stability in human serum revealed that MCo-CVX-5c is more stable than the native MCoTI-I, the grafted peptide CVX15 Gln6Cit or the linear, reduced, and alkylated version of MCo-CVX-5c. Biolayer interferometry analysis also indicated that the cyclotide MCo-CVX-5c is capable of binding to serum proteins with association and dissociation rate constants of $(3.6 \pm 0.7) \times 10^3 \text{ M}^{-1} \text{ s}^{-1}$ and $(1.4 \pm 0.2) \times 10^{-2} \text{ s}^{-1}$, respectively.⁹⁵ Serum binding has been reported to be a valuable strategy to prolong serum half-life of peptides.⁹⁸

In a follow-up study, the grafted cyclotide MCo-CVX-5c was used as a template to synthesize MCo-CVX-6 in which a Lys residue in loop 6 was replaced by an Arg residue.⁹⁹ MCo-CVX-6 was then

conjugated to the metal-chelating agent 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA) *via* its ϵ -amine group of the Lys residue in loop 1, resulting in MCo-CVX-6D (Fig. 4). MCo-CVX-6D showed increased *in vitro* binding affinity for CXCR4 ($K_i = 0.15 \pm 0.01 \text{ nM}$) (Table 1) and was exploited in a radiolabeled cyclotide with ⁶⁴Cu²⁺ [⁶⁴Cu]MCo-CVX-6D suitable for positron emission tomography-computed tomography (PET-CT) imaging studies in mice carrying CXCR4 glioblastoma xenografts.⁹⁹ Subsequent *in vivo* studies demonstrated significant uptake of radiolabeled MCo-CVX-6D in tumors expressing CXCR4, whereas the MCoTI-I control peptide labeled with [⁶⁴Cu-DOTA] mainly accumulated in the kidneys.⁹⁹ These data are in line with a report by Wang *et al.* who determined renal clearance of MCoTI-II as the major elimination route.¹⁰⁰ These studies highlight the ability of MCoTI-I framework to accommodate sequence variations to develop stable and potent antagonists of the CXCR4.

Design of cyclotide-based peptides with angiotensin-like activity

The renin-angiotensin system orchestrates essential physiological processes in the human body including blood pressure and fluid balance.¹⁰¹ These physiological events are executed and maintained by a number of enzymes and endogenous peptides. Angiotensin I is produced by the protease renin which cleaves the precursor protein angiotensinogen, whereas angiotensin II is generated by angiotensin-converting enzyme which cleaves angiotensin I.¹⁰¹ Angiotensin-converting enzyme in turn is capable of cleaving angiotensin I and angiotensin II to form angiotensin 1-9 and angiotensin 1-7, respectively.¹⁰¹ Accumulating evidence underpins the implication of the renin-angiotensin system in carcinogenesis and its modulation by endogenous angiotensin (AT) 1-7.¹⁰² Studies have shown that the activation of the proto-oncogene MAS 1 receptor (MAS1R) induced by AT 1-7 significantly inhibits human lung cancer growth both *in vitro* and *in vivo*.¹⁰³ Hence, targeting MAS1R provides an appealing strategy to develop therapies against lung cancer.

Driven by these studies and owing to the poor pharmacodynamic and pharmacokinetic profiles of AT 1-7,¹⁰⁴ Aboye and colleagues grafted an AT 1-7-like peptide into loop 6 of the MCoTI-I cyclotide, with Asp¹ and Pro⁷ being replaced by diaminopropionic acid and Glu, respectively (Fig. 4).¹⁰⁵ The AT 1-7 analogue was placed between Gly³⁰ and Gly³³ residues of MCoTI-I using the β -amino and γ -carboxylic groups, thereby generating two isopeptide bonds in the new MCo-AT 1-7 cyclotide (Fig. 4). This enabled the AT 1-7 analog to retain its positively and negatively charged groups at the N- and C-termini.¹⁰⁵ The synthesized MCo-AT 1-7 showed striking *in vitro* stability in human serum with a half-life of 39 hours compared to its linearized, reduced and alkylated version, which degraded within 57 minutes.¹⁰⁵ These results further stress the significance of the CCK motif for the enzymatic stability of cyclotides.

The ability of the grafted cyclotide to modulate the MAS1R signaling was assessed in a cell-based fluorescence assay in a CHO cell line stably expressing the receptor and measuring



intracellular NO levels. The MCo-AT 1–7 cyclotide was shown to act as an agonist of the MAS1R by increasing intracellular NO in a concentration-dependent manner (Table 1).¹⁰⁵ Co-incubation of MCo-ATI 1–7 with the MAS1R peptide antagonist A779 reinforced the specificity of the cyclotide graft as no increase in the intracellular NO concentration was observed. Overall, the findings of this study support the idea of using the MCoTI-I framework as a springboard for the development of peptide GPCR ligands with angiotensin-like activity.

Conclusions and perspective

Cyclotides have the potential to address some of the major challenges of peptide therapeutics including serum as well as gastrointestinal stability, oral activity and membrane permeability. In this review, we have elaborated on several examples in which grafting of bioactive epitopes onto cyclotide templates has proven to be a powerful strategy to facilitate the development of peptide GPCR ligands with enhanced pharmacological properties such as oral activity, receptor selectivity and/or resistance to proteolysis (Fig. 3, 4 and Table 1). Besides the potential of grafted cyclotides as therapeutics, we have referred to a study in which cyclotides have been shown to be a promising diagnostic tool by targeting GPCRs to enable the early detection of cancer and to monitor its progression.

However, several limitations and challenges associated with molecular grafting need to be addressed in future research.

Correct folding of grafted cyclotides is one of the key challenges which may explain why no engineered cyclotide targeting a GPCR of interest has yet entered human clinical trials. This being the case, a broader understanding of the processes involved in oxidative folding of cyclotides and how grafting influences folding would be valuable. Exploiting *de novo* design of peptides,¹⁰⁶ recently discovered cyclization enzymes^{107,108} or hybrid methods relying on nature-inspired biosynthetic reactions¹⁰⁹ are promising strategies to facilitate the folding of grafted cyclotides. In another relevant development, Yin *et al.* recently established a method based on the structure of anchor residues of the epitope and scaffold to guide form and function in grafted peptides.¹¹⁰ Another challenge is that the development of cyclotides bioengineered to unleash immune responses suggests that there is a possibility of inadvertent immunogenicity occurring for some cyclotide constructs.^{111,112} Compared to protein therapeutics,¹¹³ grafted cyclotides are less prone to be immunogenic due to their small size and resistance to proteolysis; however, careful assessment of preclinical immunogenicity studies prior to clinical evaluation is regarded as imperative.

Despite these challenges, grafted cyclotides represent powerful ligands and tools to illuminate GPCR pharmacology. For instance, it is known that native cyclotides do not cross the blood–brain barrier (BBB),¹⁰⁰ however, this feature may be advantageous to target GPCRs specifically expressed in the peripheral nervous system. Difelikefalin is an excellent example of a peptide drug with poor penetration of the BBB, that was introduced into the clinic in 2019 to treat postoperative pain by selectively targeting

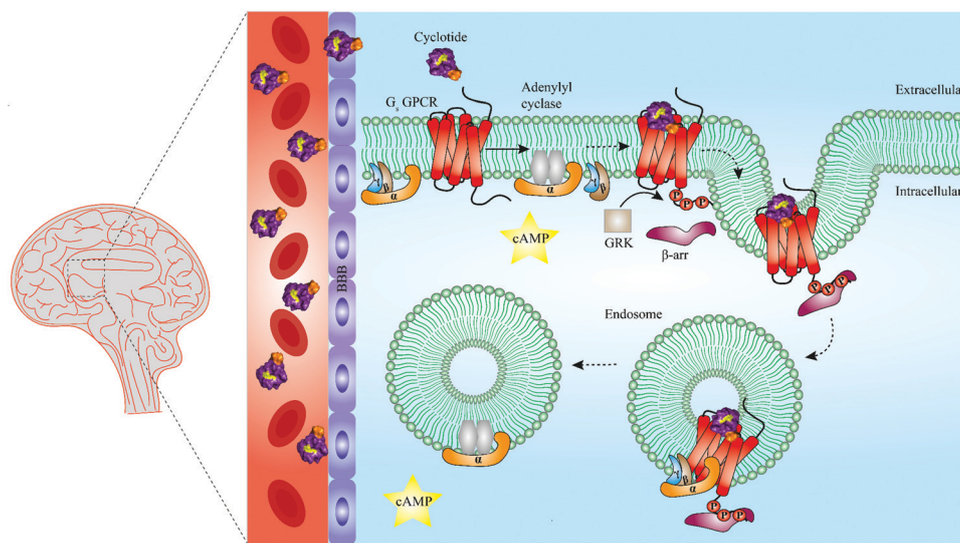


Fig. 5 Summary of unresolved functions of engineered cyclotides as GPCR ligands. Native cyclotides have restricted capabilities to cross the blood–brain barrier (BBB), yet they are suitable to develop cyclotide grafts to target GPCRs in the central nervous system by penetrating the BBB. They may be harnessed as a tool to study biased signaling of GPCRs by either activating G protein-dependent or β -arrestin-dependent signaling pathways. Additionally, cyclotides may be able to initiate endosomal GPCR signaling of G_s protein-coupled receptors (G_s GPCR). For instance, following G_s GPCR activation by a grafted cyclotide, conformational change of a G_s GPCR leads to dissociation of the heterotrimeric G protein complex and production of cAMP (second messenger) *via* activation of adenylyl cyclase. While modulation of G protein-dependent pathways by cyclotides is well-studied (straight arrow), β -arrestin recruitment and thus β -arrestin-dependent pathways are poorly investigated (dashed arrows). Phosphorylation of a G_s GPCR by GRKs results in β -arrestin recruitment and initiation of clathrin-mediated endocytosis. According to a recent study published by Nguyen *et al.*, once internalized, G_s GPCR and β -arrestin form a megaplex that activates G_s protein to enable a sustained endosomal G_s -mediated signaling.¹²¹ GRK = G protein-coupled receptor kinases, cAMP = cyclic adenosine monophosphate, G_s = stimulatory G protein, GPCR = G protein-coupled receptor.



κ -opioid receptors.¹⁰ Targeting κ -opioid receptors in the periphery has emerged as an appealing strategy to develop novel analgesics without centrally mediated side effects including sedation, dysphoria and hallucinations.¹¹⁴ Given the remarkable stability of cyclotides and their suitability for molecular grafting as well as their limited capability to penetrate the BBB, there is great potential to develop novel peripherally restricted cyclotide-based peptide GPCR therapeutics.

In recent years, cell penetrating peptides have gained impetus as a class of peptides capable of carrying and translocating large proteins or hydrophilic molecules across cell membranes.¹¹⁵ Intriguingly, cyclotides have been demonstrated to possess cell penetrating properties. For example, Ji *et al.* chemically engineered a cyclotide that antagonized an intracellular interaction of the tumor suppressor p53 and the oncoprotein Hdm2 both *in vitro* and *in vivo*.¹¹⁶ However, it has been known for a long time that cell penetrating peptides can remain trapped in endosomes during endocytosis, leading to their lysosomal degradation and limited ability to reach their target sites and to convey biological activity.¹¹⁵ Whether and how cyclotides escape endosomes remains to be determined, but the work of Ji *et al.* demonstrates that grafted cyclotides are active in the cytoplasm.¹¹⁶ Furthermore, Huang and colleagues have optimized the cell penetrating properties of cyclotides in order to foster their intracellular uptake.¹¹⁷ It is thus of interest to explore if the capacity of cyclotides to penetrate cell membranes can be leveraged in the future to engineer and develop cyclotide-based peptide ligands able to target GPCRs beyond barriers and reach areas such as the central nervous system (Fig. 5). Recent studies offer evidence that canonical GPCR signaling is not confined to the plasma membrane but may also occur in endosomes, suggesting that targeting endosomal GPCRs is a feasible strategy with therapeutic potential.^{118–121} This raises the further question of whether cyclotides equipped with a given bioactive epitope may modulate endosomal GPCR signaling (Fig. 5).

In the context of developing GPCR drugs devoid of centrally mediated deleterious side effects, the past five years have witnessed a renaissance in studies of the biased signaling of GPCRs. The ability of GPCR ligands to selectively activate one pathway over another¹²² has led to development of several peptide biased ligands preferably activating either G protein-dependent or β -arrestin-dependent signaling pathways linked to either beneficial or detrimental effects. For example, Zhang *et al.* identified a G protein-biased peptide agonist of the glucagon-like peptide-1 receptor by screening large combinatorial libraries suggesting that G protein-biased peptide agonists targeting GLP-1R are promising molecules to treat type II diabetes.¹²³ On the other hand, β -arrestin biased peptide agonists targeting angiotensin II receptor type I have been proposed to have therapeutic potential for the treatment of cardiomyopathy.¹²⁴ The ability of engineered cyclotides to engage in β -arrestin-biased signaling remains an unexplored area that warrants further investigation (Fig. 5). All cyclotide grafts discussed in this review have been chemically synthesized based on either Möbius or trypsin inhibitor cyclotide

frameworks due to the fact that bracelet cyclotides are more difficult to fold *in vitro*,¹²⁵ albeit they are more prevalent in nature than Möbius or trypsin inhibitor cyclotides.¹²⁶ Hence, it remains an elusive question for now to know if grafting bioactive epitopes onto bracelet cyclotide scaffolds would lead to the development of cyclotide GPCR ligands with distinct pharmacological properties, *e.g.* improved potency or target selectivity.

Given the importance of GPCRs as drug targets,⁶ the unique diversity of naturally occurring cyclotides as well as advances in molecular grafting methodologies,¹¹⁰ we are confident that plant-derived cyclotides will provide new opportunities for GPCR drug development.

Conflicts of interest

There are no conflicts to declare.

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