Method to generate highly stable D-amino acid analogs of bioactive helical peptides using a mirror image of the entire PDB

Michael Garton\(^a\), Satra Nim\(^b\), Tracy A. Stone\(^bc\), Kyle Ethan Wang\(^d\), Charles M. Deber\(^bce\), and Philip M. Kim\(^sed\,1\)

\(^a\)Donnelly Centre for Cellular and Biomolecular Research, University of Toronto, Toronto M5S 3E1, Canada; \(^b\)Department of Biochemistry, University of Toronto, Toronto M5S 1A8, Canada; \(^c\)Division of Molecular Medicine, Research Institute, Hospital for Sick Children, Toronto MSG 0A4, Canada; \(^d\)Department of Molecular Genetics, University of Toronto, Toronto M5S 1A8, Canada; and \(^e\)Department of Computer Science, University of Toronto, Toronto M5S 2E4, Canada

Biologics are a rapidly growing class of therapeutics with many advantages over traditional small molecule drugs. A major obstacle to their development is that proteins and peptides are easily destroyed by proteases and, thus, typically have prohibitively short half-lives in human gut, plasma, and cells. One of the most effective ways to prevent degradation is to engineer analogs from dextrorotary (D)-amino acids, with up to 10\(^5\)-fold improvements in potency reported. We here propose a general peptide-engineering platform that overcomes limitations of previous methods. By creating a mirror image of every structure in the Protein Data Bank (PDB), we generate a database of \(~2.8\) million D-peptides. To obtain a D-analogue of a given peptide, we search the (D)-PDB for similar configurations of its critical “hotspot” residues. As a proof of concept, we apply our method to two peptides that are Food and Drug Administration approved as therapeutics for diabetes and osteoporosis, respectively. We obtain D-analogs that activate the GLP1 and PTH1 receptors with the same efficacy as their natural counterparts and show greatly increased half-life.

Proteins and peptides have a number of properties that make them highly effective as therapeutic agents. These include very precise specificity, high binding affinity, low toxicity, and low risk of drug–drug interactions. Their diversity also provides very broad coverage of disease targets. Despite this, there are relatively few peptide drugs approved—around 60—with around 1,500 small molecule drugs. One major reason for this is their susceptibility to degradation by proteases and rapid renal clearance (1). Consequently, they often have prohibitively short gut, blood plasma, and intracellular half-lives. Peptides therefore tend to have low i.v. bioavailability and especially poor oral bioavailability, requiring frequent injections and severely limiting their use. Many peptide drug candidates struggle to progress beyond preclinical experiments due to bioavailability considerations. An array of techniques designed to stabilize peptides and increase their half-life has emerged and is currently driving a rapid expansion in drug candidates (2). These include PEGylation, backbone modifications, cyclization, stapling, and lipidation (3). One of the most effective approaches is the incorporation of dextrorotary (D)-amino acids (4, 5).

All amino acids except glycine exhibit chirality and, therefore, can exist in one of either (D) or levorotary (L) forms—so-called because of their influence on plane-polarized light. (D)-amino acids are occasionally found in nature (e.g., in some venomous, antibiotics, and peptidoglycan cell walls); however, this is extremely rare (6). Biology is peculiarly homochiral and constructed almost exclusively from the (L)-enantiomer. A useful consequence of this is that (D)-proteins are highly resistant to degradation and have low immunogenicity (7). The fundamental change in backbone–side-chain connectivity and geometry means they are not recognized as proteins by many (L)-proteins—including proteases. Consequently, (D)-proteins are reported to have greatly increased gut, blood plasma, and intracellular half-lives (8). Better cell penetration has also been reported in some cases (9, 10). This behavior can impart potency improvements of up to five orders of magnitude compared with their (L)-counterparts (11).

There are currently two approaches to engineering proteins with (D)-amino acids. Both approaches have significant limitations that preclude application to the majority of known or putative therapeutic peptides and drug targets (12–14). Simply replacing (L) for (D)-amino acids is generally ineffective as side-chain orientations with respect to the target are completely altered (15). Fig. 1A shows the consequence of simple (D) replacement in helical peptides. Change in side-chain orientation prevents correct binding geometry and destroys target binding.

An elegant solution to this problem in unstructured peptides is retro-inversion (RI). RI involves reversing the (D)-peptide sequence—flipping the termini and, thus, restoring the (L)-amino side chain angles. It has been used with some success on unstructured peptides (16, 17). The extended (D)-peptides assume side chain topology similar to their parent molecule but with inverted amide peptide bonds. However, RI usually fails if the peptide has secondary structure, owing largely to the topological properties of helices. (D)-peptides always adopt left-handed helices (18, 19), while (L)-peptide helices are always right-handed.

Significance

Using D-amino acids as the building blocks for bioactive peptides can dramatically increase their potency. However, simply swapping regular levorotary amino acids for dextrorotary (D)-amino acids alters the peptide surface topology and function is lost. Current methods to overcome this are not generally applicable and exclude the majority of therapeutic targets. By creating a mirror image of all \(111,867\) protein structures in the Protein Data Bank (PDB), we convert this repository into a D-peptide database with \(2.8\) million D-peptide structures. This D-PDB can be searched to find therapeutically active topologies, demonstrated here by the discovery of D-peptide GLP1R and PTH1R agonists. Evaluation of D-PDB coverage suggests that it holds candidates for most therapeutic targets and, thus, potentially contains hundreds of potent drug leads.

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RI and MIPD limitations mean that the majority of known and putative therapeutic targets are inaccessible to current (D)-peptide engineering techniques. Here, we propose an approach that overcomes these limitations and enables the design of helical (D)-peptides to a much broader range of targets. A schematic of our approach is shown in Fig. 1C.

The PDB contains over 110,000 naturally occurring and engineered structures. It is therefore a very rich source of information for the rational design of proteins. Our approach exploits this resource by creating a mirror image version of the entire repository—thereby rendering every structure in (D)-amino acids. We further compartmentalize the structures into ∼2.8 million helices and call this database the “(D)-PDB.” The (D)-PDB is then scanned—using structural alignment—for residue configurations that match the hotspot residue configurations of their partially proteolyzed (L)-peptides (Fig. 1C). Hotspot residues are those identified as contributing significantly to target recognition, binding, and receptor activation. They are a small subset of the full peptide—typically no more than three or four residues. Finding a structurally equivalent set in the (D)-PDB is therefore highly probable.

Using the glycogen-like-peptide (GLP1) and parathyroid hormone (PTH) as proof-of-concept test cases, we successfully generated (D)-helix agonists of the GLP1 and PTH1 receptors using matches discovered in the (D)-PDB.

**Results**

**(D)-PDB Construction.** Internal interactions of a protein are identical in its mirror image. This allowed the creation of a parallel protein database composed of (D)-proteins simply by flipping structure files with Cartesian coordinates along the x axis. Each flipped structure is composed entirely of (D)-amino acids and should fold exactly as the in silico structure shows when synthesized. A schematic showing (D)-PDB construction is shown in Fig. 2.

After removing any nonprotein molecules such as DNA, solvent, and ions, each file in the PDB is flipped along the x axis to create the mirror image. Nonhelical secondary structure was then removed and each helix was put into a separate file, totaling more than 2.8 million helix files. This separation ensured that hotspot alignments would only occur on relatively short, contiguous peptide regions. Redundancy was allowed, as even small differences—such as different side chain rotamers—increased the method power. Since protein regions without secondary structure can effectively be converted to (D)—experimentally using RI, we removed such regions from the (D)-PDB. Beta-sheet/strand structures were also removed for simplicity, and because therapeutic peptides tend to be either helical or unstructured.

**Query Preparation.** It is first necessary to identify or make a crystal structure—or NMR solution structure—of the functional (L)-peptide. A homology model could also be used. It should be noted that homology model effectiveness will likely be highly dependent on the degree of conservation with known structures. Residues critical to target binding and activity can then be identified—often from the literature—by alanine scanning mutagenesis. Ideally this is done experimentally, but with a target bound structure, it can also be carried out computationally using techniques such as thermodynamic integration or free energy perturbation.

Once the hotspot residues are identified, various atom sets are designated within each residue. Usually these are pairs of atoms, but in the case of ring-containing amino acids such as Phe, Tyr, and Trp, a set may include three. Each set is ranked according to its importance to target interaction, with level 1 being the highest. Level 1 usually means the atom pair or triplet furthest from the backbone and, thus, closest to the target. We assume that if level 1 can be matched, the remaining side chain atoms need not match to be effective. This greatly increases the chance of a match in the (D)-PDB. The other levels are used if level 1 atom pairs do not produce any suitable matches. Lower atom level matches can be used because one of the residue’s rotamers—above this level—will usually correctly position the level 1 atoms.
Intramolecular clash can occur between these rotamers and nonhotspot residues in the match. This is identified by full reconstruction of the match, with rotamers that allow correct level 1 positioning. Matches thereby considered nonviable are discarded. Any three atoms of a ring (or ripples) can be used to ensure that the correct planarity is represented. Default atom levels for each standard amino acid are shown in Fig. S1.

Another way to increase the likelihood of a (D)-PDB match is to group residues by similarity. For example, if a query hotspot is Arg, then matches with both Arg and Lys could be allowed. A (D)-peptide Lys match may be effectively used in the final design, or mutated to (D)-Arg with little effect on helix integrity. Similarity residue groupings are shown in Fig. S1 and, with atom renaming, can be used in combination with atom levels to maximize (D)-match likelihood.

In many cases, (L)-peptides of interest have both helical and unstructured elements. Only hotspot in the helical region are used, on the basis that unstructured peptide cannot be generated by RI, and added in postprocessing. To facilitate RI linkage, the last helical residue immediately adjacent to the unstructured region is designated a “junction” residue and included in the alignment (D)-PDB scan. Only backbone atoms (N, O, C, and CA) are used for junctions unless the junction is also a hotspot. Using backbone atoms ensures the postmatch-added RI unstructured peptide will be oriented in the same direction as the (L)-equivalent was. This ensures that correct arrangement of unstructured hotspots in relation to structured hotspots is possible when the unstructured RI region is attached.

For the RI peptide sections to be attached, D-matches need to have opposite sequence direction to the L-query. For example, if a query hotspot is Arg, then matches with both Arg and Lys could be allowed. A (D)-peptide Lys match may be effectively used in the final design, or mutated to (D)-Arg with little effect on helix integrity. Similarity residue groupings are shown in Fig. S1 and, with atom renaming, can be used in combination with atom levels to maximize (D)-match likelihood.

Test Case: GLP-1. GLP-1 is currently of interest as a diabetes mellitus and obesity treatment (25) and was chosen as a good proof-of-concept test case. It is a particularly challenging case that tests the limits—and demonstrates the full utility—of this method. It involves multiple helices, multiple unstructured regions, negatively charged hotspots, positively charged hotspots, hydrophobic hotspots, ringed hotspots, and a junction—hotspot residue. GLP-1 is a helical GPCR agonist, and this makes engineering a (D)-analog very difficult using conventional methods. There is good availability of structures and hotspot residue information (25) together with a structure for the ligand bound to the extracellular domain of the B-class GPCR (26).

Fig. 3 shows the process of preparing GLP1 structures to query the (D)-PDB. Unbound NMR solution structures (PDB ID code: 4G7ZM) and the receptor bound crystal structure (PDB ID code: 5IOL) were used as starting points. GLP-1 is composed of two helices joined by a four-residue flexible linker. Each helix was set up separately with a view to relinking two matches using the retro inverted linker sequence. Helix 1 runs from T7 to Y13. Helix 2 runs from A18 to K28. In helix 1, T7 and D9 are identified as hotspot residues, while T7 and Y13 act as junction residues. F17, I18, and L20 are the hotspot for helix 2, while A18 and K28 are the junction residues. Fig. 3B shows how hotspot and junction residues are prepared following extraction from their structure. First, the junction residue backbone atoms are rotated about the CA-CB axis. Then and by 108.5° about the CA along a defined plane. This ensures that a (D)-match can accept correctly oriented RI linker and terminal tail sequence in postmatch processing. Following this, six query structures are generated for helix 1, and 27 for helix 2, one for each combination of atom levels (Fig. 3C). In the event of no good match, each of the 33 query structures can be rerun using chemically similar residues. Fig. 3D delineates the order in which each of these was prepared, together with the combinations of atom levels used in each case. K34, while not a definitive hotspot, has been shown to contribute slightly. For this reason, both Lys and Arg were queried before “any,” as a positive charge was slightly preferred.

(D)-Match Output Processing. After running each query variant sequentially as outlined in Fig. 3, a number of matches were located in the (D)-PDB. Match quality was measured using the root-mean-square deviation (rmsd) of every atom level combination with corresponding level combinations—if they exist—in every (D)-PDB file. The rmsd cutoff was set for <1.5 Å, although <1.0 Å is ideal if possible. The best match for helix 1 was found in 3s6d, pdb at 0.5 Å, and in 4rzf.pdb at 0.9 Å for helix 2. Fig. 4A shows query—match structural alignments and—together with colored dots—indicates the successful query variants. Match sequences are reverse ordered due to the junction backbone 180° rotation, allowing RI peptide extension as planned. Both sequences are substantially different to their (L)-query. These matches were then combined with RI unstructured regions to construct the full (D)-analog of GLP-1 (Fig. 4B). A full (D)-analog structure was constructed and docked to the GLP1 receptor (GLP1R) ECD structure (Fig. 4C). Residues R12 and Q13 were found to clash with the receptor and were therefore mutated to alanine. W3 and H23 were also provisionally mutated to original query residue types—subject to checks on helix integrity.

The full (D)-analog sequence was checked for helix integrity using PSI-pred (27) (Fig. 3D). In addition to the mutations, this was to check that secondary structure is preserved when matched helices are removed from their full protein context. It also provides assurance that the full (D)-analog can fold in the same way as its components. This is necessary in order that the (D)-peptide configuration of hotspot residues closely resembles that of the (L)-peptide, and is presented as such to the target. It also
such as unwanted structure induced shows total loss of (L)-GLP-1 Best (D)-match results and full (D)-peptide construction. shows that (D)-peptide sequentially test the (D)-PDB until close matches are μ a requirement and showed no activity. ∼ Garton et al. value of 59.6 nM To demonstrate the general maximum stimulation by For- Preparation of GLP-1 queries for scanning the (D)-PDB. (is thereby established. value of 2.2 www.pnas.org/cgi/doi/10.1073/pnas.1711837115 C de decreasing quality (atom pairs or triplets according to estimated import to target binding. ensures (D)-peptide matches have reversed sequence order junction residues that have their backbone atoms rotated 180°. Rotation charge evaluation of (L)-GLP1. An isoelectric point prediction of pH 4.66, and net charge evaluation of −1.9 at pH 7 for (D)-GLP1 using pepcalc (28), indicates that it has good solubility and, thus, is suitable for experimental validation.

Experimental Validation of (D)-GLP1. The best candidate was then synthesized from (D)-amino acids and tested for its capacity to activate GLP1R. Binding of GLP1 to the GLP1R has previously been shown to activate adenyl cyclase (AC) with consequent production of cAMP, which, in turn, activates protein kinase A (PKA) to phosphorylate and activate cAMP response element-binding protein (CREB). In the present study, we investigated the ability of (D)-GLP1 peptide to induce activation of GLP1R and compared the response with native (L)-GLP1 peptide. We generated a stable GLP1 receptor/CRE-luciferase expressing HEK293 cell line and observed a cAMP-inducible luciferase expression following treatment with Forskolin (Fig. 5A). (D)-GLP1 peptide increased luciferase expression in GLP1 receptor expressing HEK293 cells, but was inactive in pCDNA3.1 HEK293 cells. (L)-GLP1 peptide displayed an EC₅₀ value of 59.6 nM with 67.2% efficacy relative to maximum stimulation by Forskolin (Fig. 5A). (D)-GLP1 peptide also increased luciferase expression in GLP1 receptor expressing HEK293 cells (Fig. 5A).

Fig. 3. Preparation of GLP-1 queries for scanning the (D)-PDB. (A) GLP-1 structures and sequence, including the free peptide in solution (Left) and receptor ECD bound structure (Right). Free GLP-1 was solved using NMR and reveals a central unstructured linker region in contrast to GLP-1R bound. Hotspot and junction residues are annotated in green and blue, respectively. (B) Hotspot are extracted separately for helix one and two, together with junction residues that have their backbone atoms rotated 180°. Rotation ensures (D)-peptide matches have reversed sequence order—a requirement for RI linker and tail attachment. (C) Levels (1–3) are assigned to hotspot atom pairs or triplets according to estimated import to target binding. (D) Atom levels are combined with similar residues. A combination order of decreasing quality—to sequentially test the (D)-PDB until close matches are identified—is thereby established.

(d)-GLP1 peptide displayed an EC₅₀ value of 2.2 μM with a similar efficacy as the (L)-GLP1 peptide. A scrambled version of (D)-GLP1 was simultaneously tested as a negative control—to account for any nonspecific effects—and showed no activity.

To investigate the mechanisms underlying the effects of (D)-GLP1 peptide on GLP1R, we studied downstream effects of activating GLP1R with (D)-GLP1 peptide. We looked to find if activation of GLP1R with (D)-GLP1 peptide would induce phosphorylation of ERK1/2 and Akt. In HEK293 cells expressing GLP1R, 10 μM (L)-GLP1 peptide evoked a robust increase in ERK activation as assessed by the increase in phospho-ERK1/2 (Fig. S4). The maximum level of phospho-ERK1/2 was achieved around 60 min after stimulation. (D)-GLP1 peptide at a concentration of 10 μM also activated ERK1/2, evoking a maximum increase of phospho-ERK1/2 around 60 min after stimulation. The level of phospho-ERK1/2 was sustained after 120 min following (D)-GLP1 treatment while the signal decreased after 60 min with (L)-GLP1.

Resistance to protease degradation is one of the most useful properties of D-peptides generally. We carried out quantitative analysis of the (D)-GLP1 Proteinase K (ProtK) resistance and compared with (L)-GLP1. Fig. 5C shows total loss of (L)-GLP1 in <1 h, while 80% of (D)-GLP1 can still be detected after 6 h exposure to ProtK.

Test Case 2: Parathyroid Hormone. To demonstrate the general applicability of this technique, another test case was selected. Parathyroid hormone (PTH) is an FDA-approved treatment for osteoporosis delivered by daily s.c. injection. Osteoporosis affects ~200 million people worldwide but only a fraction receive highlights any influence that unstructured RI regions may have on the helix or vice versa—such as unwanted structure induced into an RI region by adjoining helix. Fig. 4D shows that (D)-GLP1 has approximately the same secondary structure profile as (L)-GLP1. An isoelectric point prediction of pH 4.66, and net charge evaluation of −1.9 at pH 7 for (D)-GLP1 using pepcalc (28), indicates that it has good solubility and, thus, is suitable for experimental validation.

Experimental Validation of (D)-GLP1. The best candidate was then synthesized from (D)-amino acids and tested for its capacity to activate GLP1R. Binding of GLP1 to the GLP1R has previously been shown to activate adenyl cyclase (AC) with consequent production of cAMP, which, in turn, activates protein kinase A (PKA) to phosphorylate and activate cAMP response element-binding protein (CREB). In the present study, we investigated the ability of (D)-GLP1 peptide to induce activation of GLP1R and compared the response with native (L)-GLP1 peptide. We generated a stable GLP1 receptor/CRE-luciferase expressing HEK293 cell line and observed a cAMP-inducible luciferase expression following treatment with Forskolin (Fig. 5A). (D)-GLP1 peptide increased luciferase expression in GLP1 receptor expressing HEK293 cells, but was inactive in pCDNA3.1 HEK293 cells. (L)-GLP1 peptide displayed an EC₅₀ value of 59.6 nM with 67.2% efficacy relative to maximum stimulation by Forskolin (Fig. 5A). (D)-GLP1 peptide also increased luciferase expression in GLP1 receptor expressing HEK293 cells (Fig. 5A).

Fig. 4. GLP-1 Best (D) match results and full (D)-peptide construction. (A) (L)-query sequences (Upper) showing hotspots (green), junction residues (blue), and remaining original sequence (gray). Closest matching (D) structures are shown with atom levels annotated with dots corresponding to colors from Fig. 3. Match sequences are significantly different to query sequences. Helix 1 is highlighted light gray and helix 2 is dark gray (Lower). (B) Full (D)-analog construction from best (D) match helix sequences juxtaposed with RI linker and terminal tail sequences. (C) Construction of D-analog structure from match helices and modeled linker. Docking to GLP-1 R ECD identifies potential steric clashes, circumvented by mutation to alanine. Reintroduction of native peptide side chains at two junction positions is also judged prudent. (D) PSI-PRED predicts that correct secondary structure is maintained in the (D)-analog, with medium to high confidence (blue bars). (E) Solubility check results predict good solubility.
PTH, partly due to the lack of an oral delivery option. (D)-peptides have shown some oral bioavailability in human trial (29, 30) and, thus, a (D)-analog of PTH could be of interest. PTH is also of interest for treating hyperparathyroidism (31) and to promote bone growth following fracture (32).

The same process as described for GLP-1 was repeated: Crystal structures and hotspot residues were identified from the literature (33, 34). Fig. S5A shows PTH (1–34) with hotspot residues colored green and junctions in blue, again split into two helices. Helix one hotspots + junctions found a closest (D)-PDB match of 0.95 Å, while helix two was 0.82 Å (Fig. S5B). Reconstruction of the full (D)-peptide using RI for the linker and terminal tails is shown in Fig. S5C. A structural model of the (D)-PTH was constructed and positioned on the receptor to align with hotspot residues. Several mutations were introduced to remove clash and enhance similarity to (L)-PTH (M1–M5).

As with GLP-1 and GLP1R, binding of PTH (residues 1–34) to the parathyroid receptor (PTH1R) has also been shown to activate AC, triggering cAMP production. This activates PKA to phosphorylate and activate CREB. Fig. 6A shows that the (D)-PTH designed here activates PTH1R with a potency and efficacy comparable to (L)-PTH and Forskolin. Protease stability was also calculated and again showed a dramatic difference in degradation rate between the (L) and (D) versions (Fig. 6B and Fig. S5D). All of the (L)-PTH is degraded in under 1 h, while more than 85% of the (D)-analog is still detectable at 6 h.

**Discussion**

We have developed a method for converting (L)-peptides to highly stable (D)-analogs by exploiting a mirror image version of the PDB. We used it to engineer (D)-peptide analogs of the agonists GLP1 and PTH that successfully activate GLP1R and PTH1R, respectively. It is a simple and inexpensive method to implement, especially if a starting structure is already available—or can be obtained with reasonable confidence by homology modeling. Otherwise, helical (L)-peptide structures are mostly straightforward to obtain using X-ray crystallography or NMR. Information on hotspot residues can often be sourced from the literature, or otherwise obtained by straightforward alanine scanning mutagenesis experiments.

An interesting point emerging from this study is a powerful confirmation of the notion of hotspot residues. We have shown that a peptide can be totally changed—except for the position a small subset of residues—and function is still retained.

While (D)-PTH was comparable to (L)-PTH potency and efficacy, (D)-GLP1 potency was ~40-fold lower than the native peptide, albeit with similar efficacy. Further optimization could involve testing multiple candidates—as only one of seven (D)-GLP1 candidates produced by the method was tested—and refining with mutagenesis experiments. However, as a proof-of-concept study, we were primarily concerned with the less trivial problem of finding functional (D)-scaffolds. Affinity is a common victim of methods to engineer stability. However, increased stability and longer plasma half-life means that even a large affinity loss can still yield a net improvement in potency. Phospho-ERK experiments indicated that half-life was increased by ~fivefold. It is well established that GLP1R has rapid internalization and desensitization (35, 36); it is thus likely that this process is responsible for the relatively modest improvement in activity duration, rather than degradation. This was confirmed by protease K degradation experiments, which showed dramatic improvement in stability for both GLP-1 and PTH. It should be noted that our study compared the activity of (D)-analogs to the native hormones, and not to the many available analogs that may have higher potency (particularly in the case of GLP1). Therefore, depending on the application, much additional work may be required to optimize (D)-analogs and fully assess their therapeutic potential compared with currently approved solutions. This may involve introducing noncanonical amino acids or chemical modifications.

Nonhotspot residues can vary greatly between the original (L)-peptide and (D)-analogs engineered this way. While not contributing...
significant to the interaction, these differences may still adversely affect binding. For instance, bulky or charged (D)-peptide residues may interact with the target in a disruptive manner, especially if that space in the (L)-version is occupied by small or uncharged residues. Mutagenesis could be used to resolve this. GLP-1 was one of the more challenging cases: an agonist consisting of two helices connected by a flexible linker. Reconstructing the full peptide from two different (D) matches, such that both helices retain correct relative orientations to each other, is difficult. Cases with a single helix, fewer hotspots, or simpler antagonist action, will likely be more tractable. There is one RI limitation that our method does not overcome. Target interactions with the (L)-peptide backbone may be lost upon conversion to a (D)-peptide. Binding is likely to be adversely affected if it requires hydrogen bonding with the peptide backbone.

(D)-analogs generally avoid some of the limitations of stabilizing methods such as stapling, lipidation, PEPylation (2). These approaches can lead to significant conformational change that can adversely affect their activity. Reduced solubility is another common drawback associated with such approaches. In certain cases, where these limitations are not catastrophic, (D)-analogs could potentially be enhanced using these techniques. Combining approaches is likely to be additive or synergistic in terms of increasing half-life. As such, (D)-PDB matching can be seen as complementary rather than competing with them.

Peptide therapeutics are currently undergoing a huge expansion, and the market size is predicted to continue its dramatic increase over the next few years (37). The most recent published estimate for the number of peptides in clinical and preclinical development is 140 and 500, respectively (3). With ~80% of these likely to be helical (14), this means that over 500 of these are potentially immediately applicable for use with the (D)-PDB method. The majority of these are at present prohibited by the limitations of current methodologies. It should be noted that this estimate was published in January 2015 and, therefore, the current number of peptides in development is likely to now be significantly higher. Several (D)-amino acid containing peptide therapeutics have been approved for use, thus far indicating no inherent toxicity to humans (37).

Our work demonstrates proof of concept and leaves scope for further development into, for example, β-strand peptides, and the engineering of larger (D)-proteins. We anticipate that (D)-PDB matching will become another key tool for finding stable lead molecules in early stage drug discovery.

Materials and Methods

The latest protein database was downloaded and converted to the (D)-PDB by inverting the x-coordinate sign in each structure. Hotspot information for GLP-1 and PTH was readily available in the literature. Structural alignments were carried out with Click (38), while PSI-PRED (27) was used to predict the likely secondary structure, and PepCalc (28) was used to predict peptide solubility. Both (L)- and (D)-peptides were synthesized by Lifetech LLC. The HEK293 cell line was obtained from the American Type Culture Collection (ATCC), and cells stably expressing GLUT1 and reporter CRE-Gal4 Luciferase construct were read using a luminometer with a 480-nm filter. ProTek (Bioshop) digestions were repeated three times, and densitometry of bands was determined using ImageJ software (39). For a more detailed description, please refer to the Supporting Information.

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1. Bruno BJ, Miller GD, Lim CS (2013) Basics and recent advances in peptide and protein drug delivery. Ther Deliv 4:1443–1467.
2. Carib-Sil-Vergez M, Simon A, Sim M, Kim PM (2017) Strategies to alter the properties of motif-mediated protein-protein interactions as drug leads. Annu Rev Pharmacol Toxicol 57:39–60.
3. Fosgerau K, Hoffmann T (2015) Peptide therapeutics: Current status and future directions. Drug Discov Today 20:122–128.
4. Welch BD, VanDemark AP, Heroux A, Hill CP, Kay MS (2007) Potent D-peptide inhibitors of HIV-1 entry. Proc Natl Acad Sci USA 104:16828–16833.
5. Liu M, et al. (2010) D-peptide inhibitors of the p53-MDM2 interaction for targeted molecular therapy of malignant neoplasms. Proc Natl Acad Sci USA 107:14321–14326.
6. Kreil G (1997) D-amino acids in animal peptides. Annu Rev Biochem 66:337–345.
7. Uppalapati M, et al. (2016) A potent D-protein antagonist of VEGF-A is nonimmunogenic, metabolically stable, and longer-living in vivo. ACS Chem Biol 11:1058–1065.
8. Knezevic AE, Pentelute BL (2015) A D-amino acid at the N-terminus of a protein abrogates its degradation by the N-end rule pathway. ACS Cent Sci 1:423–430.
9. Nick CL, et al. (2010) (D)-Amino acid analogues of DT-2 as highly selective and superior inhibitors of GMP-dependent protein kinase lambda. Biochim Biophys Acta 1802:645–653.
10. Brugidou J, Legrand C, Mety J, Ribbie A (1995) The retro-inverso form of a homeobox-derived peptide is rapidly internalized by cultured neurons: A new basis for an efficient intracellular delivery system. Biochim Biophys Acta 1244:685–693.
11. Veine DM, Yao H, Stafford DR, Fas KS, Livant DL (2014) A D-amino acid containing peptide as a potent, noncovalent inhibitor of uS1 integrin in human prostate cancer invasion and lung colonization. Clin Exp Metastasis 31:379–393.
12. Li C, et al. (2010) Limitations of peptide retro-inversion isomerization in molecular mimicry. J Biol Chem 285:19572–19581.
13. Rabaut L, Olivier N, Melfy O (2012) Sequential native peptide ligation strategies for total chemical protein synthesis. Chem Soc Rev 41:7001–7015.
14. Law V, et al. (2014) DrugBank 4.0: Shedding new light on drug metabolism. Nucleic Acids Res 42:D1991–D1097.
15. Li C, et al. (2013) Functional consequences of retro-inversion isomerization of a mini-protein inhibitor of the p53-MDM2 interaction. Bioorg Med Chem 21:4045–4050.
16. Li H, et al. (2015) Novel retro-inversion peptide inhibitor reverses angiotensin receptor antagonist autobody-induced hypertension in the rabbit. Hypertension 65:793–799.
17. Ben-Yedidia T, Beignon AS, Partidos CD, Muller S, Arnon R (2002) A retro-inverso analogue of influenza virus hemagglutinin B-cell epitope 91-108 induces a strong mucosal and systemic immune response and confers protection in mice after intranasal immunization. Mol Immunol 39:523–531.
18. Hung LW, Kohnmura M, Ariyoshi Y, Kim SH (1998) Structure of an enantiomeric protein, D-mollin on 1.8 A resolution. Acta Crystallogr D Biol Crystallogr 104:16828–16833.
19. Novotny M, Kleywegt GI (2005) A survey of left-handed helices in protein structures. J Mol Biol 347:231–241.
20. Jochim AL, Arora PS (2010) Systematic analysis of helical protein interfaces reveals targets for synthetic inhibitors. ACS Chem Biol 5:919–923.