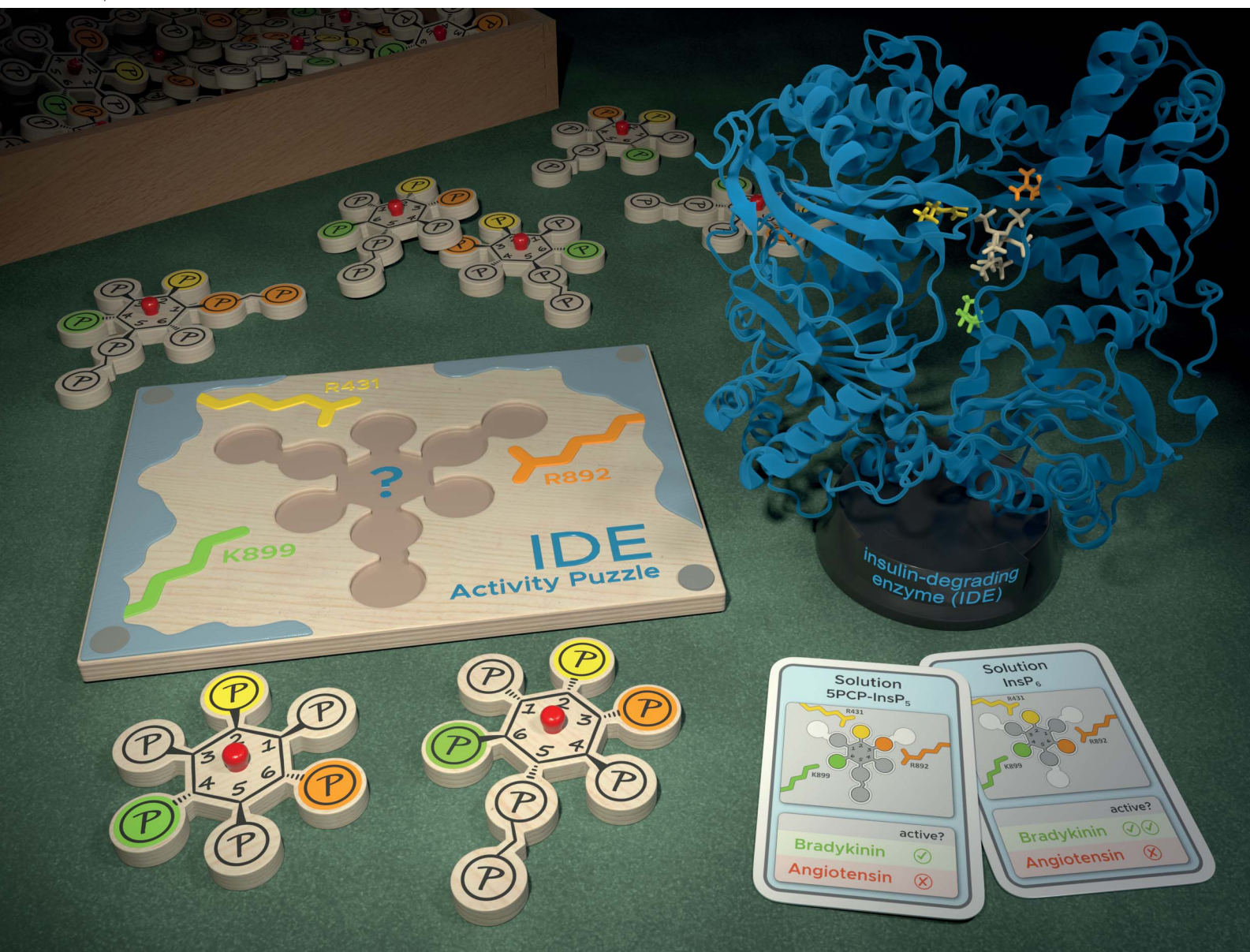


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Dissecting the activation of insulin degrading enzyme by inositol pyrophosphates and their bisphosphonate analogs†

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Inositol poly- and pyrophosphates (InsPs and PP-InsPs) are densely phosphorylated eukaryotic messengers, which are involved in numerous cellular processes. To elucidate their signaling functions at the molecular level, non-hydrolyzable bisphosphonate analogs of inositol pyrophosphates, PCP-InsPs, have been instrumental. Here, an efficient synthetic strategy to obtain these analogs in unprecedented quantities is described – relying on the use of combined phosphate ester-phosphoramidite reagents. The PCP-analogs, alongside their natural counterparts, were applied to investigate their regulatory effect on insulin-degrading enzyme (IDE), using a range of biochemical, biophysical and computational methods. A unique interplay between IDE, its substrates and the PP-InsPs was uncovered, in which the PP-InsPs differentially modulated the activity of the enzyme towards short peptide substrates. Aided by molecular docking and molecular dynamics simulations, a flexible binding mode for the InsPs/PP-InsPs was identified at the anion binding site of IDE. Targeting IDE for therapeutic purposes should thus take regulation by endogenous PP-InsP metabolites into account.

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Introduction

Inositol-based signaling molecules are central eukaryotic messengers and include the highly phosphorylated, diffusible inositol polyphosphates (InsPs) and inositol pyrophosphates (PP-InsPs). The PP-InsPs are most densely phosphorylated, and one or two high-energy pyrophosphate groups are attached to the *myo*-inositol scaffold. Genetic studies have linked PP-InsPs to a wide range of biological processes in mammals, including insulin signaling, spermatogenesis, and blood coagulation.^{1–3} To mediate such varied cellular responses, distinct molecular mechanisms of action have been proposed for PP-InsPs. These mechanisms comprise allosteric regulation of enzyme activity, competition with phosphatidyl inositols for lipid binding domains, and protein pyrophosphorylation.^{4–7} Although allosteric regulation of protein function by PP-InsPs is readily conceivable, it is currently not well understood how subtle structural changes of the ligand are translated to notable alterations in protein activity.

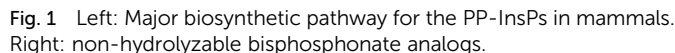
Insulin-degrading enzyme (IDE) is a conserved zinc metalloprotease, which is involved in the degradation of numerous bioactive peptides, including insulin, glucagon, amyloid beta peptides, bradykinin and angiotensin.^{8,9} Consequently, potent and selective modulation of IDE activity could provide a gateway to treat major diseases, such as type 2 diabetes and Alzheimer's disease. Despite the large therapeutic potential of targeting IDE, the regulatory mechanisms governing IDE substrate selectivity are not well characterized. Notably, IDE possesses an anion binding site remote from its active-site, and binding of poly-anions, such as ATP, can promote the proteolytic cleavage of some IDE substrates, in particular short peptide substrates.^{10–14} It was recently demonstrated that also phosphatidylinositols and inositol polyphosphates enhanced proteolytic activity of IDE, but only limited data was presented on the effect of PP-InsPs.¹⁵

The investigation of the properties and functions of PP-InsPs is often hindered by the lability of the pyrophosphate moieties, which can undergo hydrolysis in physiological contexts or during synthesis. Therefore, non-hydrolyzable methylene bisphosphonate analogs of PP-InsPs have been developed (PCP-InsPs, Fig. 1). These stable mimics have been applied as PP-InsP surrogates in biochemical assays,^{16–19} structural studies,²⁰ as mechanistic tools^{17,21,22} and as affinity reagents.^{23,24} Previous strategies to obtain the PCP-InsPs relied on phosphoryl chloride (PCP-Cl) reagents to append the PCP moiety in modest yield, while requiring high stoichiometry of the reagents and

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Encouraged by these results, we next applied the PCP-amidite reagents to the synthesis of 1,5-(PCP)₂-InsP₄ (Scheme S3†). Appending two PCP-moieties had been painstaking in the past using the phosphoryl chloride reagents, which required reagent recycling, and provided only modest yields (20–30%).^{18,20} Likewise, reacting 4 eq. of PCP-amidite **3a** with diol **6a** resulted in a mixture of the desired product **7a** with compounds bearing only one bisphosphonate moiety. The steric hindrance of the benzyl group presumably prevented the full conversion to the di-functionalized product. By contrast, when 4 eq. of PCP-amidite **3b** were reacted with diol **6b**, we obtained the doubly modified bisphosphonate **7b** in excellent yield (95%). Once again, this strategy proved scalable, providing over one gram of

1. Synthesis of the PCP-amidite reagents



2. Synthesis of the PCP-InsPs analogs



Scheme 1 PCP-amidite reagents for the synthesis of PCP-InsPs.

compound **7b**. The bisphosphonate analog 1,5-(PCP)₂-InsP₄ (**2**) was obtained from intermediate **7b** following a synthetic route similar to the one used for **1**, and batches of 50–100 mg of **2** were isolated at a time.

Differential modulation of IDE activity by PP-InsPs

We next wanted to investigate the influence of PCP-InsPs/PP-InsPs on the activity of insulin degrading enzyme (IDE) (Fig. 2A). Membrane-bound and diffusible inositol phosphates were recently shown to stimulate cleavage of IDE substrates, in particular the cleavage of small peptide substrates.¹⁵ However, only little data was provided for PP-InsPs. Given the wide substrate scope of IDE, and the increasing interest in PP-InsP signaling and metabolism, elucidating whether PP-InsPs govern IDE activation and substrate selectivity is important before further exploring IDE as a pharmacological target.

The activation of IDE by PP-InsPs was assayed using a well-established model fluorogenic peptide (Fig. 2B).^{31,32} Cleavage of the model peptide by IDE results in a fluorescence recovery that can be followed over time. Initial rates were measured for various peptide concentrations in the absence or presence of InsP₆, 5PP-InsP₅, 1,5-(PP)₂-InsP₄, or their non-hydrolyzable counterparts. All ligands activated IDE, in particular at low peptide concentrations (<20 μM) (Fig. 2C and S1†). At higher peptide concentrations, all ligands displayed a substrate

inhibition profile. Such behavior had already been observed in the case of the lower inositol phosphates, and could be due to competition with the peptide for a secondary substrate binding site.¹⁵ Interestingly, InsP₆ was the least efficient activator, whereas 5PP-InsP₅ and 1,5-(PP)₂-InsP₄ activated the cleavage of the peptide most strongly (Fig. 2C and S1†). The additional pyrophosphate on 1,5-(PP)₂-InsP₄ did not seem to promote further activation of the enzyme as compared to 5PP-InsP₅. The PCP analogs mirrored this trend and induced the proteolytic cleavage by IDE more strongly than InsP₆, although they exhibited lower potency than the natural molecules (Fig. 2C, D and S1†). Because the pK_a values for the methylene bisphosphonate group are slightly elevated compared to the pyrophosphate group,³³ the overall protonation states of the non-hydrolyzable analogs differ at physiological pH, potentially altering their interaction with the polyanion binding site of the protein.

We next determined the effect of PP-InsPs on IDE activation with respect to the natural substrates bradykinin and angiotensin. Using an HPLC assay, the initial rates of the enzyme were measured in the absence and presence of InsP₆ and PP-InsP/PCP-InsP ligands. Cleavage of bradykinin was strongly activated by InsP₆ and PP-InsPs (Fig. 2E). Contrary to the model peptide substrate, the strength of activation was very similar for InsP₆, 5PP-InsP₅ and 1,5-(PP)₂-InsP₄, and not more pronounced for the inositol pyrophosphates. The PCP-InsPs again displayed





Fig. 2 Inositol pyrophosphates modulate IDE activity distinctly, depending on the small peptide substrate. (A) Structure of rat IDE (pdb: 3tuv; identity to human IDE: 95%), highlighting relevant binding regions. IDE is composed of four domains, represented here in pale cyan (domain 1), teal (domain 2), light grey (domain 3) and dark grey (domain 4). Amino acids of relevant sites are highlighted in different colors: (1) anion binding site, orange; (2) active site, red; (3) peptide exosite, blue. (B) Principle of the fluorogenic activity assay. When intact, the fluorescence of the peptide is quenched. Upon cleavage of the peptide by IDE, fluorescence is recovered and can be measured as a read-out for enzyme activity. (C) Activity of IDE on a fluorogenic peptide in presence or absence of InsPs (5 μM). Experiments were performed at 37 °C in 50 mM Tris buffer, pH 7.4, 50 mM NaCl. All experiments were performed at least in triplicate, and standard error of the mean of these replicates are shown as intervals. (D–F) Activity of IDE for various peptides (peptide concentration: 10 μM, InsP concentration if present: 5 μM; IDE concentration: 0.05 μg mL⁻¹). Error bars show standard error of the mean of at least three independent experiments. Statistical significance was assessed using one-way ANOVA. All samples containing InsP₆/PP-InsPs were compared to the "No InsP" sample. The result of Dunnett's multiple comparison test is shown above each bar. (ns: non significant, $p > 0.1$; no marker: $p < 0.1$; *: $p < 0.05$; **: $p < 0.01$; ***: $p < 0.001$, ****: $p < 0.0001$). (F) is reproduced on a magnified scale in the ESI (Fig. S3†).

less potency as activators, compared to the natural molecules. Unexpectedly, InsP₆ and PP-InsP/PCP-InsPs had no activating effect on the proteolytic cleavage of angiotensin. Neither InsP₆, nor the PP-InsPs/PCP-InsPs accelerated the rate of angiotensin cleavage. In fact, angiotensin cleavage was even slightly inhibited by 1,5-(PP)₂-InsP₄ (Fig. 2F).

Overall, the InsPs/PP-InsPs influenced IDE activity in markedly different ways, depending on the peptide substrate. While cleavage of the model peptide substrate was most strongly promoted by the pyrophosphate-containing metabolites 5PP-InsP₅ and 1,5-(PP)₂-InsP₄, IDE catalyzed cleavage of bradykinin was activated similarly by InsP₆, 5PP-InsP₅ and 1,5-(PP)₂-InsP₄. By contrast, no activating effect of InsPs/PP-InsPs was observed with angiotensin as the substrate, indicating a high degree of flexibility and adaptability in the system.

PP-InsPs target the anion binding site of IDE

Deciphering the interplay between IDE, InsPs/PP-InsPs and distinct peptide substrates at the molecular level could advance our understanding of the regulation of this central

enzyme. Unfortunately, attempts to obtain co-crystal structures of IDE and the PCP-InsP analogs remained unsuccessful. Also, HDX-MS data could not pinpoint distinct conformational changes upon binding of PP-InsP ligands. While a decrease in deuterium exchange rate was evident in the peptide binding region (peptide 351–359) in samples containing both the bradykinin substrate and 5PCP-InsP₅, a significant effect on the PCP-InsP–IDE-substrate complex could not be described (Fig. S4†). These results may indicate a more dynamic nature of the PP-InsP–IDE interactions, making it challenging to capture them experimentally.

As an alternative approach, we employed computational methods to explore possible interactions between PP-InsPs and IDE. Potential binding modes of InsP₆ and 5PCP-InsP₅ to IDE were first investigated by molecular docking using GLIDE module implemented in the program Schrödinger.^{34,35} These two ligands were chosen, since DFT-optimized structures of InsP₆ and 5PCP-InsP₅ based on density functional theory (DFT) (Table S2†) had been described previously.^{20,36} Both ligands were docked to the ATP binding site of rat IDE, which exhibits



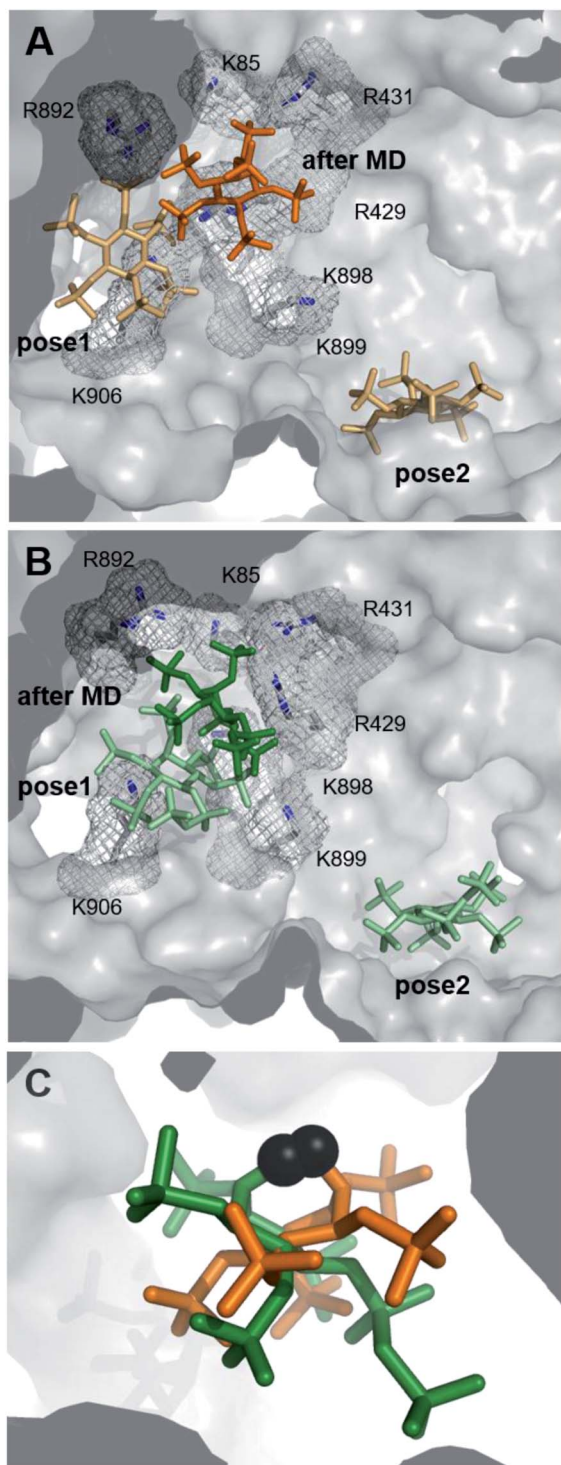


Fig. 3 Computational investigation of the interactions of InsP_6 and 5PCP-InsP_5 with IDE. Two binding poses for InsP_6 (A) and 5PCP-InsP_5 (B) were obtained by docking experiments, which converged to one binding position after MD simulations. One exemplary pose after 100 ns MD simulation (pose after MD) is shown for both InsP . Basic amino acids (Lys and Arg) in pose after MD are highlighted as sticks and in the mesh representation. Protons are omitted for clarity. (C) Superimposition of MD binding sites of InsP_6 (orange) and 5PCP-InsP_5 (green) illustrate the variable orientations of InsPs when binding to IDE. Protons are omitted for clarity. Phosphate groups at the 2-position of the inositol ring are represented as dark grey spheres to facilitate comparison.

a sequence identity of 95% with respect to human IDE (pdb: 3tuv).¹⁰ For both ligands, two binding poses were identified close to the ATP binding site, independent of their ionization state (Fig. 3A and B). In pose 1, similar side-chain interactions (R429, R892, K896, and K898) to those reported for ATP¹⁰ were involved in the recognition of InsP -phosphoryl groups. As InsP_6 and 5PCP-InsP_5 do not, or only slightly, overlap with ATP, further contacts of the ligands in pose 1 were established with R893, K906, and R920. In contrast, in pose 2, which was more remote from the ATP binding site, the ligands interacted mostly with K425, K571, and K632. Both poses were populated in the molecular docking analysis by various energetically comparable sub-poses with minor structural variations, including rotation and translation (Fig. S5†). These observations indicate a high degree of flexibility of InsP_6 and 5PCP-InsP_5 when binding to IDE. As a result, it was impossible to assign interactions of specific phosphoryl groups with individual amino acids.

To optimize the predicted docking poses in a flexible protein environment, and to validate our hypothesis of heterogeneous InsP -IDE complexes, both poses of InsP_6 and 5PCP-InsP_5 were subjected to all-atom molecular dynamics (MD). During the MD simulations, both InsP molecules underwent noticeable re-orientation with respect to their starting poses, which reveals the limitations of the rigid docking approach. Interestingly, all models converged to only one principal binding mode, as illustrated in Fig. 3A and B. In this binding mode, however, the $\text{InsP}/\text{PP-InsP}$ molecules adopted a wide range of orientations (Fig. 3C and S6†), varying in rotation and slightly in translation. Comparison of the binding poses of InsP_6 and 5PCP-InsP_5 revealed that both ligands share interactions with the same key amino acids (R429, R431, R892, R899) (Fig. S6†). The interaction with K85, K898, and K906 occurred sporadically but not specifically for either of the two InsP molecules.

Taken together, the computational approach identified one main binding mode for InsP_6 and 5PCP-InsP_5 at the anion binding site of IDE. The binding to this basic cavity, however, was highly flexible, with a heterogeneous population of ligand conformations, which might explain the experimental difficulty for characterizing these protein-ligand complexes by common structural biology techniques. The observed heterogeneity also provides a first clue as to how these structurally very similar molecules may be able to dictate substrate specificity.

Conclusion

In sum, we utilized combined phosphoester-phosphoramidite reagents to efficiently synthesize and isolate non-hydrolyzable analogs of PP-InsPs . Two PCP-amidite reagents, compatible with distinct protecting group schemes, were synthesized in good yields. These new building blocks consolidate existing PCP-amidite reagents and will enable alternative synthetic strategies to install methylene bisphosphonate groups on chemically and/or pharmaceutically relevant target structures. The PCP-amidite reagents were then applied to the synthesis of PCP-InsPs , yielding unprecedented amounts of these analogs, due to the efficient coupling of the PCP-amidites to sterically encumbered hydroxyl groups. Beyond the high coupling



efficiency, this strategy also holds promise for further derivatization on the bisphosphonate moiety, as exemplified here by the introduction of a ^{13}C -label into the PCP group. In light of our current computational and biochemical data, it appears that the protonation states of InsPs/PP-InsPs may be an important factor governing the interactions with IDE. In the future, incorporating a fluorinated bisphosphonate group could be of interest, since it better mimics the pyrophosphate group with regard to the pK_a values.³³

We then utilized the PCP-InsP analogs, alongside their natural counterparts, to interrogate the mechanism of activation of IDE by PP-InsPs. Biochemical characterization revealed a strong dependence of protein activity on both the InsP/PP-InsP ligands and the peptide substrates. Subsequent MD simulations could pinpoint the interaction between IDE and InsPs/PP-InsPs to the anion binding site of IDE, and highlighted the high degree of flexibility in these interactions, as illustrated by the wide range of orientations of the ligands. To capture these dynamic interactions experimentally with high resolution will require the development of new molecular tools, such as PP-InsP-protein cross-linking reagents.

Because several IDE substrates have opposing functions (bradykinin *versus* angiotensin, insulin *versus* glucagon), it is interesting to speculate whether a fine-tuned balance of PP-InsP metabolites can influence complex organismal responses, such as vasodilation and carbohydrate metabolism, *via* IDE. As potent and selective pharmacological tools targeting PP-InsP metabolism are now becoming available,^{37,38} we should be able to address these questions in a more complex environment in the future. In return, understanding the modulation of IDE substrate multiplicity by PP-InsPs would also inform current drug development programs, targeting the inositol kinases.

Data availability

The data supporting this article have been included as part of the supplementary material.

Author contributions

S. H. and Do. F. designed the experiments and wrote the manuscript. S. H., Da. F. and K. F. synthesized and characterized the compounds described in this paper. S. H. performed all biochemical assays and analyzed the data. G. L. D. performed the HDX experiments and analyzed the data, supported by V. H., H. S. and T. U. designed the computational studies, T. U. and B. T. performed them and T. U. and H. S. analyzed the data. All authors reviewed the manuscript.

Conflicts of interest

The authors declare no competing financial interest.

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