THE CATALYTIC MECHANISM OF THE MARINE-DERIVED MACROCYCLASE, PATGmac

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Abstract: Cyclic peptides are a class of compounds with high therapeutic potential, possessing bioactivities including anti-tumour and anti-viral (including anti-HIV). Despite their desirability, efficient design and production of these compounds has not yet been achieved. The catalytic mechanism of patellamide macrocyclization by PatG macracyclase domain has been investigated using computational methods. We applied a quantum mechanics/molecular mechanics (QM/MM) methodology, specifically ONIOM(M06-6-311++G(2d,2p):ff94/B3LYP/6-31G(d):ff94). The mechanism proposed here begins with a proton transfer from Ser783 to His 618 and from the latter to Asp548. Nucleophilic attack of Ser783 to the substrate leads on to the formation of an acyl enzyme covalent complex. The leaving group (AYDG) of the substrate is protonated by the substrate’s N-terminus leading to the breakage of the P1-P1’ bond. Finally, the substrate’s N-terminus attacks the P1 residue, decomposing the acyl-enzyme complex forming the macrocycle. We found that the formation and decomposition of the acyl-enzyme complex have the highest activation free energies (21.1 kcal.mol⁻¹ and 19.8 kcal.mol⁻¹ respectively), typical of serine proteases. Understanding the mechanism behind the macrocyclization of patellamides will be important to the application of the enzymes in the pharmaceutical and biotechnological industries.

Introduction

The pharmaceutical industry has become excellent at developing small-molecule drugs (below 600 MW) hitting classical compact binding sites. However, there remain a large number of non-classical targets with extended binding sites which cannot be modulated using small molecules, but instead need biologics (antibodies/native peptides) to obtain a therapeutic effect. Of the top selling drugs on the market 7 out of 10 are biologics aimed at complex diseases such as rheumatoid arthritis and cancer.[11] Macrocycles (500-2000 Da) are a third class of therapeutics which are able to modulate the same complex, extended, targets as biologics, but are easier to administer, hit intracellular targets, and may have a lower associated cost of goods.[9] Of the 68 approved macrocycle pharmaceuticals, 27 are cyclic peptides[8], 1 of which is orally available. Cyclic peptides have a number of advantages over linear ones including reduced susceptibility to metabolism, improved membrane permeability and an entropic advantage on binding to a target. The pharmaceutical industry is struggling with two aspects of macrocycles: their design and efficient production. Most processes to generate peptide macrocycles rely on the use of high dilution conditions to prevent oligomerisation, using controlled addition conditions making these approaches non-viable for large-scale synthesis.[6a,b] Alternative synthetic approaches have been developed including on-bead macrocyclization requiring attachment to the solid support via an amino acid side chain and a 3-dimensional orthogonal protecting group strategy.[5] Biological methods of macrocyclization include sortase-mediated ligation, but this results in the incorporation of the pentapeptide LPXTG in the cyclic peptide where X is variable.[6b]

Efficient formation of cyclic peptides without leaving a residual sequence in the final macrocycle is desirable. Natural cyclic peptides can be formed either via a non-ribosomal peptide synthetase route, or the more recently discovered superfamily of ribosomally produced and post-translationally modified peptides (RiPPs).[9] RiPPs are formed using a common biosynthesis, in which a precursor peptide, comprised of a leader sequence followed by a core peptide often flanked by signal sequences, is modified via the action of processing enzymes, which install post-translational modifications in the core peptide. The matured core peptide is then removed from the leader and signal sequences, liberating the mature, active peptide. In many cases, this last step results in the formation of a peptide macrocycle. Three RiPP macrocyclases have been defined; butelase-1 an asparagine/aspartate peptide ligase that is responsible for the formation the plant cyclic peptide cyclotides,[7] GmpOB, a polyol oligopeptidase involved in α-amanitin biosynthesis[8] and PatG macracyclase (PatGmac), involved in the biosynthesis of the cyanobactins which are cyanobacterially derived azole/azoline containing cyclic peptides.[9] The requirements of the PatGmac are a core peptide sequence of 6-11 residues which may include D-amino acids or unnatural amino acid residues, an AYD(GE) signal at the C-terminus which is not incorporated in the final cyclic peptide, and a cyclic residue (proline or thiazoline) at the C-terminal of the core peptide which is included in the macrocycle.[10]

Our structural investigations have delineated the mode of action of PatGmac and have led to an understanding of the requirements described above.[9] PatGmac is a subtilisin-like serine protease produced by Prochloron sp. which is an obligate symbiont of the seasquirt Lissoclinum patella.[11] The normal extended conformation of the peptide substrate is prevented by bulky enzyme residues (Met660/Arg686/Phe684) protruding into the binding groove as a consequence of a disulphide bridge formation between Cys685 and Cys724. As a result of this only a bent substrate peptide can bind, and this is facilitated by the conformational properties of the proline or thiazoline residue at
the P1 position (P3, P2, P1 are the residues before the cleavage site and P1', P2' those after the cleavage site).
The unique structural feature of PatGmac is an insertion loop in the normal subtilisin sequence generating a helix-turn-helix motif forming a protective lid over the active site. The AYD signal at the C-terminus of the core peptide binds via the aspartate residue to basic residues in the helix-turn-helix motif (Arg589/Lys594/Lys598). After binding the substrate peptide, a normal serine protease mechanism ensues, but access by water is prevented by the strongly bound AYD signal, thus allowing the amino terminus of the core peptide to loop around and attack the acyl complex, resulting in cleavage of the recognition signal and cyclization of the core peptide.

In this work, we explored the mechanism of the reaction catalysed by PatGmac using computational methods, in order to describe, with atomistic detail, the most plausible catalytic mechanism of this enzyme. Considering the ability of the enzyme to macrocyclize an extensive range of nonactivated substrates, the data obtained in the present study has wide implications in a number of areas. Fully understanding the mechanism of PatGmac may lead to engineered analogues with improved properties which accept an even broader substrate range thus increasing its utility in a range of pharmaceutical and biotechnology applications.

Results and Discussion
In previous work, some of us proposed a catalytic mechanism for PatGmac\(^9\) (Figure 2) based on the structure of the enzyme macrocyclase domain and biochemical characteristics. It was suggested that the first step of the reaction is the nucleophilic attack of Ser783, aided by His618 which acts as a base, to the P1' carbonyl group of the substrate leading to the formation of an enzyme-substrate tetrahedral intermediate. After this first attack, the AYDG peptide is cleaved by protonation of the leaving group's N-terminal by His618. Lastly, the N-terminus of the substrate attacks the carbonyl group of the cysteine (P1) forming the macrocycle (Figure 2). In order to test the proposed mechanism and get a complete and detailed description of this catalytic mechanism, we have performed QM/MM calculations.

The first reaction step

In the reactants optimized structure, Ser783 is at 3.89 Å from the carbon atom of P1 and correctly oriented to begin the attack. Asp548 is hydrogen bonded to His618 (1.69 Å) and the latter is hydrogen bonded to Ser783 (1.54 Å) (Figure 3). The catalytic reaction begins with the typical proton transfer on the serine protease's catalytic triad, from Ser783 to His618. We defined the distance between \(N_\delta^1\) of His618 and the \(H_\gamma\) of Ser783 as the putative reaction coordinate. As the proton of Ser783 was being transferred to His618, the proton of the His618 imidazole ring spontaneously moved to Asp548 O-atom. We noticed that the structures of the reactants (Asp-COOH/His/OH-Ser) (Figure 3) and of the generated intermediate (Asp-COOH/His/H-O-Ser) (Figure 4) are very close in energy (2.4 kcal.mol\(^{-1}\)). The free energy barrier was calculated in 2.5 kcal.mol\(^{-1}\). At the transition state structure, both the proton transferred from Ser783 to His618 and the one that was consequently transferred from His618 to Asp548, assume an intermediate position between the respective residues (Figure 4).

The vibrational frequency analysis revealed an imaginary frequency corresponding to the reaction coordinate (an antisymmetric stretch involving both proton transfers) confirming the nature of the transition state (996.52 \(i\).cm\(^{-1}\)). However, another small negative frequency (31.30 \(i\).cm\(^{-1}\)) was found, as a consequence of the more relaxed geometry optimization. Thermal corrections and zero point energies were included in the free energy calculation, despite the existence of a very small second imaginary constant. These corrections have a small contribution to the corresponding free energy values in agreement to what has been reported in the literature regarding proton transfers between catalytic residues (ca. 1.2 kcal/mol).\(^{[26]}\) As the barrier was extremely shallow and not rate-limiting it was not optimized further.

The small free energy of reaction suggests that both structures may exist simultaneously being the latter the most favourable to initiate the catalytic reaction. The possibility of the first step to begin with a direct nucleophilic attack of Ser783 to the substrate, with the proton transfer occurring during the attack was also considered. Defining the distance between Ser783 and P1 as reaction coordinate, it was found that the energy barrier associated with this alternative is very high (>30 kcal.mol\(^{-1}\)). Our proposal for the first step has a much smaller energy barrier.
(2.5 kcal.mol$^{-1}$) indicating that the proton transfer leading to an intermediate with enhanced nucleophilicity for attack at the substrate amide.

**The second reaction step**

The deprotonated Ser783 attacks the P1 carbonyl group forming the enzyme linked tetrahedral intermediate. A study of this step was made with a linear transit scan using the optimized structure after the proton transfer described above as reactants, and defining the distance between the O$_\gamma$ of Ser783 and the carbonyl carbon of P1, which is initially 3.90 Å (Figure 4), as the reaction coordinate to be scanned.

A conformational change in His618 took place before the formation of the tetrahedral intermediate, with the approximation of Ser783 to the P1 carbonyl group. His618 changed its rotamer by rotating around the C$_\beta$-C$_\gamma$ bond, breaking the hydrogen bond with Ser783 hydroxyl and forming another with the Ser783 carbonyl, while keeping the hydrogen bond with Asp548. Additionally, a hydrogen bond between the attacking oxygen of Ser783 and Asn717 amine was formed (Figure 5). This conformational change led to a stable intermediate (INT2) before Ser783 has completed the nucleophilic attack on the substrate. A free energy barrier of 4.8 kcal.mol$^{-1}$ in relation to INT1 was found, characterized by one imaginary frequency (744.16 i.cm$^{-1}$) that corresponds to the coordinated stretching of the Ser783 oxygen-P1 carbonyl carbon and of the N-H bond of the terminal amino group of the substrate.

This proton, together with the Asn717 amine, constitutes the oxyanion hole which stabilizes the oxyanion of the acyl-enzyme complex. The interactions between them may consist of low-barrier hydrogen bonds (LBHB) considering, particularly, their short length (< 2.0 Å). The existence of LBHB between the oxyanion and the oxyanion hole in serine proteases has already been proposed$^{[31]}$ although others have considered them to be simple electrostatic interactions.$^{[32]}$ In this case we can clearly see the transition from an electrostatic hydrogen bond in the reactants to a hydrogen bond where the proton is shared between both acceptors, commonly known as a single-well hydrogen bond, which corresponds to an extreme case of a low barrier hydrogen bond where the barrier vanishes. However, we note that this structure, achieved at the transition state, is not a stable, long-lived interaction. Whether or not this kind of interactions are catalytic or anticatalytic is a matter of debate.$^{[33]}$

**The third reaction step**

Starting from the INT2 optimized structure, the attack of Ser783 to the substrate was conducted until the formation of the tetrahedral intermediate, using the distance between the Ser783 deprotonated oxygen and the P1 carbonyl carbon (3.62 Å) as reaction coordinate (Figure 5). This attack had a free energy barrier of 21.1 kcal.mol$^{-1}$ in relation to INT2. This rate-limiting transition state (TS3) was further freely optimized, having one imaginary frequency (31.83 i.cm$^{-1}$) that corresponds to the coordinated stretching of the Ser783 oxygen-P1 carbonyl carbon and of the N-H bond of the terminal amino group of the substrate.

A free energ}$^{[31]}$y barrier of 4.8 kcal.mol$^{-1}$, in relation to INT1 was found, clearly corresponding to the reaction coordinate. We will refer to this point as TS2 (Figure 5). In that structure His618 is placed between Ser783 side chain and carbonyl oxygen atoms at 2.22 Å and 2.28 Å respectively. Ser783 which initially was at 3.90 Å from the P1 residue stays at 3.57 Å on TS2 and at 3.61 Å on INT2.
The fourth reaction step

The conformational rearrangement that His618 makes it unlikely that it protonates the leaving group in the third step of the catalytic reaction, as initially proposed and which is typical for other serine proteases. After the rearrangement His618 stays at 5.97 Å from P1' and not adequately oriented (Figure 6), the substrate’s N-terminal (charged) amine group would be the most suitable candidate to protonate P1' amine.

Therefore, we conducted a linear transit scan using the distance between the NH3+ terminal proton and the P1' amine group, which is 2.01 Å in the INT3 geometry, as reaction coordinate (Figure 6). A transition state (TS4) was then freely optimized and an imaginary frequency (1184.98 i cm⁻¹) was found which corresponds to an asymmetric stretch between the transferred proton and the two nitrogen atoms (Figure 7). We found an energy barrier of 3.1 kcal.mol⁻¹ relatively to INT3. With the transfer of the proton, a very stable intermediate (INT4) is obtained (ΔGstep4 = -23.8 kcal.mol⁻¹), the bond P1-P1' is cleaved but the AYDG tetrapeptide is retained on the active site of the enzyme, at 2.95 Å from P1, on the INT4 optimized structure (Figure 7).

The fifth reaction step – Macrocyclization of the substrate

To complete the catalytic mechanism of PatGmac, the substrate’s N-terminus attacks the carbon atom of P1 carbonyl group closing the macrocycle. We have used the distance between the N-terminus and the P1 carbonyl carbon (4.99 Å) as reaction coordinate. We found that during this step the N-terminus donates a proton to Ser783, consequently decomposing the enzyme-substrate tetrahedral structure since the bond between P1 and Ser783 is cleaved. The macrocycle is formed and the AYDG peptide which was at 2.95 Å from P1 at the beginning of this step is further displaced. This step had a free energy barrier of 19.8 kcal.mol⁻¹, in relation to INT4, and originates very stable products (ΔGstep5 = -42.6 kcal.mol⁻¹) (Figure 8), being the displacement of the AYDG peptide an important factor contributing to the great stability of the products. The freely optimized TS5 geometry shows one imaginary frequency (136.59 i cm⁻¹), corresponding to the reaction coordinate. The present description of this step differs from the
first proposed mechanism[9], in that, the substrate’s N-terminus
donates a proton to Ser783 rather than His618.

The reaction rates reported for PatGmac are approximately 1
per day.[10, 36] Thus, the Gibbs free energy barrier for the
macrocyclization reaction may be estimated from the transition
state theory, resulting in an observed experimental free energy
of $\approx$ 24 kcal.mol$^{-1}$, which is a comparable value to the obtained
in the present work (21.1 kcal.mol$^{-1}$).

Conclusions

We have explored by computational approaches the
macrocyclization reaction catalyzed by PatGmac. Our results
showed that the mechanism followed by this enzyme is different
to those typical of serine proteases albeit with some similarities
(Scheme 1). The typical proton transfer on the Ser-His-Asp
catalytic triad occurs, as we describe in the first reaction step, as
well as the formation of the acyl-enzyme intermediate (second
mechanistic step). We found, however, that the protonation of
the AYDG leaving group of the substrate is most probably made
by the substrate’s NH$_3^+$ terminus and not by His618 (Scheme 1),
due to steric impediments (Figure 2). This enzyme differs from
typical serine proteases where deacylation of the enzyme-
substrate complex occurs by an attack of a water molecule[30]
regenerating the enzyme. In PatGmac, the active site is shielded
from water[9] and because of that, the deacylation is achieved by
the attack of the substrate’s N-terminus which also protonates
the acyl complex. The formation of the peptide bond is common
to macrocyclization of other peptide substrates, namely other
cyanobactins.[9, 26]

This new mechanism differs from that proposed previously
(Figure 2, Scheme 1)[9] the central difference is His618 undergoes
a conformational rearrangement and does not
protonate the leaving group. Rather it is the incoming substrate
amino terminus that protonates the leaving group.

This study contributes to further understanding the mechanism
of macrocyclization of PatG substrate. As cyclic peptides have
been seen as having great interest for industry, particularly
pharmaceutical, more knowledge about their natural synthesis
also contributes to improve the efficiency of large scale
production of such compounds. The findings in this work
suggest that adapting the enzyme process to utilize different
substrates would need careful consideration of the pKa of the
incoming nucleophile. Regarding the active site residues, the
Asp His Ser triad is mandatory for the function of the enzyme
and have to be maintained in any engineered analogue while
Asn717 stabilizes the tetrahedral-intermediate.

Viable options to increase the reaction rate have to focus on the
rate-limiting step. For so, the most promising way to achieve
catalysis would be to optimize the oxyanion hole.

Possibly the presence of a positively charged residue at position
717 (i.e. Lys) would further favour the nucleophilic attack of
Ser783 to the substrate which we found that it is a rate limiting
step of the reaction. However, the size of Lys and the proximity
with the positive N-terminus are obstacles for the correct
placement and protonation of the lysine.

Methods

Energetic profile of the PatGmac catalytic mechanism

Figure 9 shows the energetic pathway of the macrocyclization
reaction catalyzed by PatGmac at the ONIOM(M06/6-311++G(2d,2p):Amber//B3LYP/6-31G(d):Amber) level.

According to these results, the TS2 has the highest free energy.
However, that is not the rate-limiting step since both TS3 and
TS5 have higher energy barriers relatively to the preceding
intermediate which are very stable. These TS represent the
formation of the enzyme-substrate complex (TS3) and the
formation of the macrocycle and accompanying decomposition
of the acyl-enzyme complex (TS5). The TS3 has an energy
barrier slightly higher than TS5 (21.1 kcal.mol$^{-1}$ and 19.8
kcal.mol$^{-1}$ respectively), but given their proximity, both will be
relevant for the observed rate of this reaction (note that the
difference between the two is very narrow and probably close to
the accuracy of the methodology for relative energies between
similar molecular structures). Both the formation and
decomposition of the tetrahedral complex have been identified
as slow steps previously[30, 34] consistent with our results here
and with our previous mass spectrometric observation of the
acyl PatGmac intermediate. The PES of the reaction shows that
it is strongly exothermic with the products 76.0 kcal.mol$^{-1}$ more
stable than the reactants. All mechanistic steps are exothermic
with step 3 (formation of the acyl-enzyme intermediate) being
the sole exception.
We started the modelling of the system using an X-ray structure of the subtilisin-like domain of the PatGmac enzyme containing an analogue of the substrate in the active site (PDB: 4AKT, 2.63 Å resolution). This structure had a mutation (His168Ala) on the catalytic triad of the active site (Asp548, His618 and Ser783) which was reverted by superposition with a structure of the free enzyme (PDB: 4AKS, 2.19 Å resolution) that contained the original residue. The coordinates of Ala168 on the 4AKT structure were then replaced by those of His618 of the free enzyme structure. The hydrogen bonding network between the three residues of the catalytic triad and the proximity between the deprotonated nitrogen of the imidazole ring of His618 and the hydroxyl group of Ser783, observed on the free enzyme structure, is mandatory to generate a productive conformation and to initiate the catalytic mechanism. Thus, we naturally assumed that the rotameric state and position of His618 is similar in the free enzyme and in the enzyme complexed with the substrate. Additionally, in the X-ray structure, a loop composed by residues 651-657 was missing and was modelled using the program MODELLER. This loop is located away from the active site (>|10 Å), thus, the modelling performed should not have a significant effect on the study of the catalytic mechanism. One natural substrate has the sequence lle-MeOxH-Ala-ThH-Ile-OxH-Phe-ThH-Ala-Tyr-Asp-Gly, the (Ala-Tyr-Asp-Gly) are the recognition residues cleaved (at the ThH-Ala bond) during the reaction (ThH = thiazoline, OxH = oxazoline, MeOxH = methyl oxazoline). The mimic peptide on the 4AKT structure had a different sequence (Val-Pro-Ala-Pro-Ile-Pro-Phe-Pro-Ala-Tyr-Asp-Gly) in which the azoline heterocycles had been replaced by Pro. Hence, the heterocycles that on the X-Ray structure were mimicked by prolines, were corrected and missing parts were modelled in GaussView. We have preserved the original coordinates for most of the residues of the substrate. Figure SI-1 shows the original precursor peptide and the modelled peptide. We used the X- Leap software to protonate the complex, assuming that all residues were in their physiological protonation states, and 23 Na+ counter-ions were added to neutralize the charge of the system. Additionally, we surrounded the system with 15830 water molecules using a truncated rectangular box of TIP3P water molecules with a minimum distance of 12 Å between any atom of the protein and the faces of the box.

We performed a two-step minimization using the Amber 12 simulation package (parm 98SB force field) in order to relax the system by removing eventual tensions and clashes. First, the water molecules and counter-ions were minimized with the remainder of the system fixed (steepest descent algorithm for the first 500 cycles, and conjugate gradient algorithm for the last 1500 steps); and, second, the position of all atoms (steepest descent algorithm for the first 5000 cycles, and conjugate gradient algorithm for the last 10000 steps) of the model was minimized.

We ran molecular dynamics (MD) simulations, starting from the structure obtained after the minimization procedure, to see if the modelled structure was stable and preserved. First we warmed the system from 0 to 300 K in a 200 ps long simulation maintaining a constant volume and using periodic boundary conditions.

Then, a 15 ns production run was conducted using periodic boundary conditions with the isobaric-isothermal ensemble defining a pressure of 1 atm and a temperature of 300 K using the Langevin thermostat and the Berendsen barostat for that purpose. The cutoff for the Lennard-Jones interactions was set to 10 Å. The Coulomb interactions were treated using the Particle Mesh-Ewald (PME) method, with a cutoff of 10 Å for the real part of the sum. The time step of the simulation was 2 fs. The potential energy of the system was treated with the LeapFrog integration algorithm.

A structure of PatGmac complexed with substrate was taken from the MD simulation and was used as the starting point for the study of its reaction mechanism. We have chosen a structure of the system (after the equilibration) whose conformation was productive for the beginning of the catalytic cycle, which means that the distance between Ser783 and the substrate, and between the substrate’s N-terminal and P1 had to be small and appropriate for chemical reaction. The analysis of the trajectories has shown that, after the equilibration, these criteria are met in most of the structures, hence, the energetic cost for reaching a productive conformation should not be significant. To investigate the potential energy surface (PES) along the mechanism of the macrocyclization reaction we performed QM/MM calculations, widely used in enzymatic studies, applying an ONIOM scheme as implemented in the Gaussian 09 software package. The system, containing a total of 5200 atoms, was divided into a ‘QM layer’, containing 91 atoms (Figure 1), and an ‘MM layer’ which were treated at DFT and classical MM levels respectively. The high layer includes the catalytic triad (Ser783, His618 (side chain until Co) and Asp548 (side chain until Cβ)), Asn717 (side chain until Cγ) and P1 (Cys), P1′ (Ala), P2′ (Tyr) residues and the terminal ile of the substrate. A list of the atoms in the QM layer is given in table SI-1. For the QM layer we employed the B3LYP functional using 6-31G(d) basis set for geometry optimizations, which was shown to provide accurate results in previous studies, while in the low layer we used the AMBER parm 98SB force field. We used hydrogen atoms as site atoms where covalent bonds were in between the two layers. The interaction between the two layers was treated using an electrostatic embedding scheme. The study of each mechanism step began by conducting linear scans along the reaction coordinates. These corresponded to specific interatomic distances that connected the reactants to the products of each hypothesized reaction step. The precise reaction coordinates that were assumed are described in the main text, when each reaction step is discussed. The structures of the reactants, intermediates, transition states (TS) and products were then fully optimized, starting with the guesses taken from the linear transit scans, for the rate-limiting steps. In the case of chemical steps with very shallow barriers we have used the highest energy structure of the linear transit scan as very good approximations of the transition state. This procedure was motivated by the great complexity of performing free transition state geometry optimization in this very large and heterogeneous system. The differences in free energy that we got when we compared the two procedures (i.e. taking the structure from the adiabatic mapping and making a free geometry optimization, both alternatives calculated for the rate-limiting steps) were 0.4 and 2.6 kcal/mol. We performed vibrational frequency calculations for every stationary point, with those of the reactant, intermediates and product having no imaginary frequencies and those of the TS having just one, which in all cases was clearly related to the reaction coordinate. Even though the identity of the minima connected to each transition state was clear from the liner transit scans, and was further supported by the observation of the relevant normal mode, we ran further IRC calculations (albeit not to the full extent of the explored PESs), starting from the obtained TSs, to confirm that the minima that were connected to the TS were the ones that we were expecting. The final energies were obtained conducting single-point (SP) energy calculations on the optimized geometries using different density functionals (M06, B19BS and mPW1K in addition to B3LYP), known to have a good performance for thermodynamics and kinetics, and, a larger basis set, 6-311++G(d,2p), in the QM layer to improve the accuracy of the results. Table SI-2 shows the activation and reaction energies obtained with the different functionals. They provide a PES that translates into the same mechanism and are qualitatively equivalent. The values obtained with B3LYP seem a bit elevated comparatively to those obtained with the other functionals. The energy barriers for each step are similar in all cases. However, with B3LYP and M06 the energy barrier to TS3 is slightly higher than the barrier to TS5 whereas with mPW1K and B19BS the opposite is
observed. In the discussion we considered the results of the M06 functional because it has been shown to be the most appropriate for the description of the thermodynamics and kinetics of the chemistry of main-group elements.\textsuperscript{25-27} We have calculated also the zero-point energy, the entropy and the thermal corrections to obtain Gibbs free energies at 298.15 K, which is a comparable temperature to that of the water where the Indo-Pacific seastarip Lissoclinum patella naturally occurs.\textsuperscript{29} To calculate the entropy and free energy we have employed the particle in a box/rigid rotor/harmonic oscillator formalism. This is a physically clear and rigorous formalism to calculate the entropy and free energy of a system within a single-conformation model.

GD3 dispersion\textsuperscript{29} was included in the single point calculations as implemented in Gaussian 09 D.01.

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Fully understanding the mechanism of PatGmac may lead to engineered analogues with improved properties which accept an even broader substrate range, increasing its utility in a range of pharmaceutical and biotechnology applications. In the TOC graphic is presented the structures of the reactants and products of the macrocyclization catalysed by PatGmac.