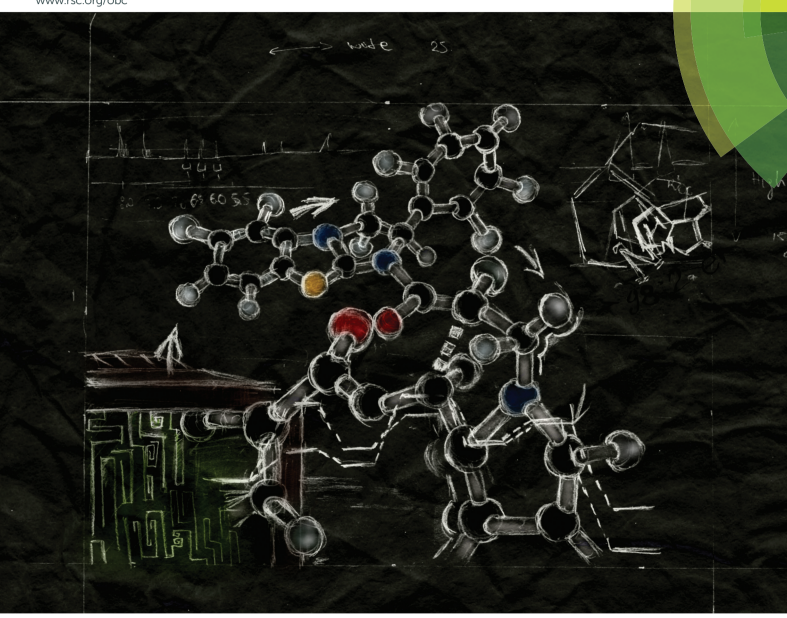
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Isothiourea-catalysed enantioselective pyrrolizine synthesis: synthetic and computational studies†

Daniel G. Stark,^a Patrick Williamson,^a Emma R. Gayner,^b Stefania F. Musolino,^a Ryan W. F. Kerr,^a James E. Taylor,^a Alexandra M. Z. Slawin,^a Timothy J. C. O'Riordan,^c Stuart A. Macgregor*^b and Andrew D. Smith*^a

The catalytic enantioselective synthesis of a range of *cis*-pyrrolizine carboxylate derivatives with outstanding stereocontrol (14 examples, >95:5 dr, >98:2 er) through an isothiourea-catalyzed intramolecular Michael addition-lactonisation and ring-opening approach from the corresponding enone acid is reported. An optimised and straightforward three-step synthetic route to the enone acid starting materials from readily available pyrrole-2-carboxaldehydes is delineated, with benzotetramisole (5 mol%) proving the optimal catalyst for the enantioselective process. Ring-opening of the pyrrolizine dihydropyranone products with either MeOH or a range of amines leads to the desired products in excellent yield and enantioselectivity. Computation has been used to probe the factors leading to high stereocontrol, with the formation of the observed *cis*-steroisomer predicted to be kinetically and thermodynamically favoured

Introduction

The 5,5-bicyclic pyrrolizine and pyrrolizidine structural motifs that contain a bridgehead nitrogen atom are present within the core of many biologically active natural products¹ such as that of dehydroretronecine 1 (Fig. 1).2 This natural product, along with many other derivatives, originates from metabolism of pyrrolizidine alkaloids (PAs), a natural alkaloid prevalent in plant life throughout nature.3 Such PA-derived molecules are known to be potent heptatoxins,4 carcinogens,5 teratogens and genotoxins.7 Compelled by such levels of biological activity, many of these natural products and their derivatives have become commercial therapeutic agents. For example, the partially reduced pyrrolizine mitomycin C 2, isolated from Streptomyces caespitosus or Streptomyces lavendulae, is a potent antitumour drug with a broad scope of applications.8 The nonsteroidal anti-inflammatory drug (NSAID), Licofelone 3 has shown great promise in osteoarthritis treatment through a dual inhibition of 5-LOX/COX.9 Additionally, another pyrrol-

Fig. 1 Representative biologically active pyrrolizines.

izine based NSAID, Ketorolac 4, has found success as a commercial analgesic. 10

Given the value and potential of these bicyclic compounds, a variety of synthetic routes towards these motifs have been designed, with many syntheses involving classic total synthesis approaches towards specific target molecules. ¹¹ In recent years the state-of-the-art in catalytic pyrrolizine syntheses has involved diastereoselective multi-step reaction processes such

^aEaStCHEM, School of Chemistry, University of St Andrews, North Haugh, St Andrews, Fife, KY16 9ST, UK. E-mail: ads10@st-andrews.ac.uk

^bInstitute of Chemical Sciences, Heriot-Watt University, Edinburgh, EH14 4AS, UK. E-mail: s.a.macgregor@hw.ac.uk

 $[^]c$ Syngenta, Jealott's Hill International Research Centre, Bracknell, RG42 6EY, UK † Electronic supplementary information (ESI) available: NMR spectra, HPLC analysis and computational co-ordinates. Data available. 12 CCDC 1483759. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c60b01557c

as the phosphine-catalysed domino reaction developed by Tong and co-workers¹³ (Fig. 2a) or the gold-catalysed process by Matsuya and co-workers (Fig. 2b).14 Catalytic enantioselective methodologies that enable efficient access to this desirable structural motif are relatively limited. Within this area, Cho and co-workers showed that an enantioselective organocatalysed Michael addition-aldol approach could generate functionalised pyrrolizines with excellent diastereo- and enantiocontrol (18 examples, >95:5 dr and 95:5 to 99:1 er, Fig. 2c), although relatively high catalyst loadings were employed to promote this process. 15 In spite of these advances there is still a clear requirement for easily accessible and reliable catalytic methodologies that can produce stereodefined chiral pyrrolizine derivatives with high levels of efficiency and enantiocontrol.

Following seminal work from Romo and co-workers using ammonium enolates generated from carboxylic acids, 16 ourselves and others,¹⁷ have used isothioureas¹⁸ to catalyse a range of formal [2 + 2], $^{19} [3 + 2]^{20}$ and $[4 + 2]^{21}$ cycloaddition processes that employ an ammonium enolate intermediate.²² Related intramolecular Michael addition-lactonisation cascades from enone-acid substrates have been used to generate simple heterocyclic products such as THFs and pyrrolidines with excellent enantioselectivity. 23 Building upon this previous work, the application of this strategy to construct the highly desirable pyrrolizine core in a catalytic enantioselective fashion starting from pyrrole-derived enone acid substrates

a. Tong (2015): γ-Addition/[3+2] cycloaddition domino reaction PPh₃ (20 mol%) Na₂CO₃ (1.2 equiv) R^1 OAc PhMe, rt. 12 h 15 examples typically 72-95% yield CO₂Bn typically >95:5 dr b. Matsuya (2015): Au-catalysed [3+2]-cycloaddition/cyclisation $CO_2CH_2CF_3$ (Ph₃P)AuCl (cat.) .CO2Et AgOTf (cat.) PhCO₂H CICH2CH2CI, 65-80 °C 6 h 11 examples 48-68% yield, >95:5 dr c. Cho (2010): Organocatalytic Michael addition-Aldol reaction OTMS $Ar = 3,5-(CF_3)_2C_6H_3$ i) (20 mol%) PhCO₂H (40 mol%) PhMe, -10 °C, 18 h 18 examples typically 60-88% yield

Fig. 2 Current catalytic methods for pyrrolizine synthesis.

ii) NaBH₄ (1 equiv)

EtOH, -10 °C, 0.5 h

>95:5 dr, 95:5 to 99:1 er

such as 5 is investigated (Fig. 3). At the onset of these studies the main challenges were envisaged to arise from the incorporation of the electron-rich N-functionalised pyrrole core within the target enone-acid. A robust method to access this structural motif has not been reported previously, while the effect of incorporating this planar electron-rich aromatic structure upon stability and reactivity, as well as the conformational and steric effects upon diastereo- and enantioselectivity were unknown. Furthermore, the potential for competitive intramolecular Friedel-Crafts acylation of the pyrrole via a mixed anhydride or acyl ammonium ion intermediate, 24 or alternatively β-elimination from an ammonium enolate, needed to be assessed. To further enhance our understanding of this process, we also wanted to probe the course of the proposed cascade cyclisation process via computation in order to understand the factors leading to stereocontrol. Very limited computational analysis of the use of isothiourea-derived ammonium enolates in catalysis has been reported.²⁵ To date only the single report from Muck-Lichtenfeld and Studer concerning the intermolecular formal 1,3-dipolar cycloaddition of azomethine imines with mixed anhydrides under isothiourea catalysis has incorporated DFT analysis.20a

In this manuscript we report the realisation of this strategy to facilitate the catalytic enantioselective synthesis of these valuable heterocyclic products 6 and 7. A straightforward three-step synthetic route to the enone acid starting materials from readily available pyrrole-2-carboxaldehydes is delineated, with commercially available benzotetramisole (BTM) proving the optimal catalyst for the enantioselective process. Furthermore, the use of computational analysis allows insight into the origin of stereocontrol in this intramolecular cascade process.

This work: isothiourea-catalysed enantioselective pyrrolizine synthesis

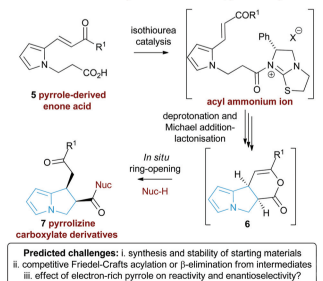


Fig. 3 Proposed isothiourea-catalysed Michael addition-lactonisation methodology for pyrrolizine synthesis.

Results and discussion

Substrate synthesis

Initial studies set out to devise a practical synthetic route towards the target pyrrole-derived enone-acid substrates. Substrate 11 was identified as a model system, and was initially prepared from pyrrole 2-carboxaldehyde by N-alkylation, ester hydrolysis and Wittig olefination. However, attempted synthesis of a range of pyrrole-derived enone-acid substrates or scale-up of this synthetic route proved low yielding and irreproducible when directed towards alternative substrates. After a thorough investigation into alternative synthetic approaches, a reliable three-step sequence starting from the corresponding pyrrole 2-carboxaldehyde was established (Scheme 1). Aldol-condensation with the requisite ketone provides a general and chromatography-free preparation of pyrrole-enones 8. Treatment with sub-stoichiometric t-BuOK and methyl acrylate gives the N-alkylation product 9, with subsequent ester hydrolysis providing the desired enone-acid substrates 10 in good overall yield with a wide scope available. This reliable synthetic sequence allowed the preparation of a range of substrates 11-18 in up to 58% yield over three steps and on multigram scale, and allows for substituent variation within both the pyrrole and enone components.²⁶ Direct chlorination of **11** with *N*-chlorosuccinimide (NCS) led to substrate 19, while an analogous procedure using acetone for the aldol reaction procedure, followed by alkylation and ester hydrolysis gave methyl enone 20.

Reaction optimisation

With a reliable synthetic route to pyrrole-derived enone acid substrates in hand, the feasibility and subsequent optimi-

R1 NH Me Ar (2.0 equiv)

EtOH, rt, 24 h

R1 NH 8

$$CO_2Me$$

(1-10 mol%)

 CO_2H

Ar EtOH: CO_2H
 CO_2H

Ar = Ph 11

 CO_2H

Ar = 4-BrC₆H₄ 12

Ar = 3-BrC₆H₄ 13

Ar = 2-thienyl 14

Ar = 2-naphthyl 16

Ar (2.0 equiv)

EtOH, rt, 24 h

R1 NH 8

 CO_2Me
 CO_2Me
 CO_2Me

Ar = A-BrC₆H₄ 19

 CO_2H
 CO_2H

Scheme 1 Optimised synthesis of pyrrole-derived enone-acid substrates.

sation of the isothiourea-catalysed Michael addition-lactonisation protocol was conducted (Table 1). Utilising achiral isothiourea DHPB 22, i-Pr2NEt and t-BuCOCl provided the desired pyrrolizine dihydropyranone 21 from enone-acid 11 in a moderate 41% isolated yield but with excellent >95:5 dr (entry 1). Alternative methods for in situ generation of a reactive carboxylate derivative, such as Mukaiyama's reagent 26, did not improve conversion or isolated yield (entry 2). A significant improvement in yield was observed when the equivalents of both t-BuCOCl and i-Pr2NEt were increased from 1.5 to 3.0 equiv., with 21 obtained in 84% yield. Subsequent studies assessed the feasibility of an enantioselective process, with chiral isothiourea catalysts 23-25 examined. Tetramisole·HCl 23 and benzotetramisole (BTM) 24-mediated reactions (entries 4 and 5) provided 21 in excellent yield, >95:5 dr and with >99:1 er. Application of HyperBTM 25 proved marginally less effective, with the product 21 obtained in high yield and dr but with reduced 96.5:3.5 er (entry 6). Notably in all of these catalytic processes no competitive products arising from either Friedel-Crafts acylation or β-elimination were observed, despite the proposed intermediacy of an acyl ammonium ion. The effect of reduced catalyst loading using BTM 24 and tetramisole·HCl 23 was next evaluated. Using 5 mol% BTM 24 maintained excellent levels of diastereo- and enantioselectivity, giving product 21 in 82% yield, while using 5 mol% tetra-

Table 1 Enantioselective Michael addition-lactonisation optimisation

O Ph 11 CO ₂ H	Reagent i-Pr ₂ NEt solvent, 0 °C then catalyst i-Pr ₂ NEt rt, t (h)	Ph H, O 21
N S P	h N S	Ph:NS
22 DHPB 2	?3 Tetramisole∙HCl	24 BTM
i-Pr _{//} , N	Tfo	⊕ N Br O Et
25 HyperBTM	26 Muk	aiyama's reagent

Entry	Reagent ^a (equiv.)	Catalyst (mol%)	Yield ^b (%)	dr ^c	er^d
1	t-BuCOCl (1.5)	22 (10)	41	>95:5	_
2	26 (1.5)	22 (10)	45	>95:5	_
3	t-BuCOCl (3.0)	22 (10)	84	>95:5	_
4	t-BuCOCl (3.0)	23 (10)	84	>95:5	>99:1(ent)
5	t-BuCOCl (3.0)	24 (10)	81	>95:5	>99:1
6	t-BuCOCl (3.0)	25 (10)	81	>95:5	96.5:3.5
7	t-BuCOCl (3.0)	24 (5)	82	>95:5	>99:1
8	t-BuCOCl (3.0)	23 (5)	67	>95:5	>99:1(ent)
9	t-BuCOCl (3.0)	24 (1)	42	>95:5	>99:1
10	t-BuCOCl (3.0)	23 (1)	34	>95:5	99:1(ent)

^a Applied in 1:1 ratio with *i*-Pr₂NEt. ^b Isolated yield. ^c Determined by ¹H NMR of the crude reaction product. ^d Determined by chiral HPLC.

misole·HCl 23 led to reduced but still acceptable 67% vield (entries 7 and 8). Further reduction of the catalyst loading to 1 mol% showed the same trend in reactivity, with BTM-24 giving 21 in 42% isolated yield, and tetramisole·HCl 23 a lower 34% yield, yet still in high dr and er (entries 9 and 10). All further studies used BTM 24 (5 mol%) as the optimal reaction conditions for this enantioselective process.

As an alternative strategy to the isolation of dihydropyranone 21 in situ ring-opening with a suitable nucleophile was investigated to provide access to pyrrolizine carboxylate derivatives (Table 2). Taking pyrrole-derived enone-acid 11 under the optimum catalysis conditions the resulting dihydropyranone 21 can be readily ring-opened in situ with MeOH giving pyrrolizine methyl ester 27 in excellent 86% yield, >95:5 dr and >99:1 er. Ring-opening using both primary and secondary amines to give pyrrolizine amides also proved successful. For example, use of allylamine provided 28 in 81% yield, >95:5 dr and 98.5: 1.5 er, while ring-opening with pyrrolidine gave pyrrolizine amide 29 in quantitative yield, >95:5 dr and >99:1 er. Morpholine, N-Boc piperazine and tetrahydrothieno[3,2-c]pyridine could also be utilised, giving the corresponding pyrrolizine amides 30-32 in good to excellent yield (60-75%), >95:5 dr and >99:1 er. Upon standing the pyrrolizine carboxylate products proved considerably more stable to storage than the

Table 2 Michael addition-lactonisation/ring-opening protocol^{a,b,c}

>95:5 dr, >99:1 er

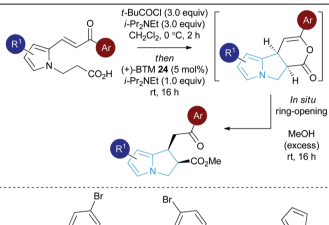
95:5 dr, >99:1 er

dihydropyranone 21, therefore ring-opening with MeOH was used as the general procedure when exploring further substrate scope.

Substrate scope: Michael addition-lactonisation/methanolysis

With a reliable synthetic route to pyrrole-derived enone acid substrates available, the generality of the isothiourea catalysis using BTM 24 as the catalyst and MeOH as the nucleophile for in situ ring-opening was evaluated (Table 3). Variation of the enone-substituent of the substrate was first investigated with a range of aryl groups accommodated. Brominated aryl units

Table 3 Michael addition-lactonisation/methanolysis scope: variation of Michael acceptor^{a,b,c}



>95:5 dr, >99:1 er

^a Isolated yield. ^b dr determined by ¹H NMR of the crude reaction product. ^c er determined by chiral HPLC.

^a Isolated yield. ^b dr determined by ¹H NMR of the crude reaction product. ^c er determined by chiral HPLC.

42

79% yield

>95:5 dr. >99:1 er

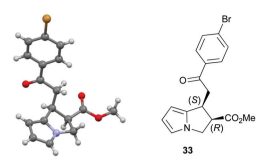


Fig. 4 Molecular representation of the X-ray structure of 33.

can be included to give 33 and 34 in 78% and 74% yield, respectively, with both generated as single diastereoisomers in >99:1 er. Incorporation of the heterocyclic 2-thienyl substituent was readily tolerated giving 35 in 77% yield, >95:5 dr and 99:1 er. Electron-rich groups (4-MeOC₆H₄) can be installed, giving 36 in excellent 93% yield, >95:5 dr and >99:1 er.

Both 1-naphthyl and 2-naphthyl units can also be incorporated to access the corresponding products 37 and 38 in good yields and excellent enantioselectivity, although attempted catalysis using Me-enone 20 did not result in any conversion to product. Variation within the pyrrole core of the pyrrolizine carboxylate product was also assessed. Chlorinated product 39 was produced in 66% yield with good 94:6 dr and as a single enantiomer (>99:1 er). The core motif can be expanded to the hexahydro-1H-pyrroloindole structure with the corresponding product 40 achieved in 53% yield and excellent stereoselectivity (>95:5 dr and >99:1 er).²⁷ The reaction to form 33 was readily carried out on a 1 gram scale, giving the desired product in 67% yield in >95:5 dr and >99:1 er. The relative and absolute configuration within 33 was assigned by X-ray crystallography analysis, with the configuration within all other products assigned by analogy (Fig. 4).²⁸

Product derivatisation

To exemplify that this methodology can potentially be used as a basis for further synthetic elaboration, product derivatisation within both the pyrrole core and aromatic ketone substituent was explored (Scheme 2). For example, the synthesis of product 33 using this cascade methodology gives a product with the bromine functional handle. Through the use of a Suzuki–Miyaura coupling 33 was readily elaborated to access the polyheteroaromatic pyrrolizine 41 in 60% isolated yield and with no loss of stereointegrity. Alternatively, a simple chlorination of the pyrrole core within 27 was conducted using NCS to access chloropyrrolizine 42 in 79% yield.

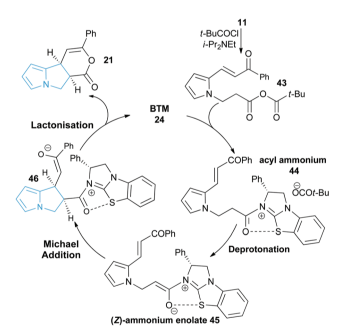
Proposed mechanism and computational studies

A proposed catalytic cycle for the synthesis of pyrrolizine dihydropyranone **21** from enone-acid **11** is shown in Scheme 3. Firstly, pivaloyl chloride reacts with the enone-acid to form mixed anhydride **43**, with subsequent nucleophilic attack from the Lewis base catalyst BTM **24** generating acyl ammonium ion **44**. Deprotonation to form (*Z*)-ammonium enolate **45**, followed

a. Suzuki-Miyaura coupling: ArB(OH)₂ (1.2 equiv) Pd(PPh₃)₄ (10 mol%) Na₂CO₃ (3.0 equiv) CO₂Me DME, 85 °C, 5 h 33 Ar = OMe 60% yield >95:5 dr, >99:1 er >95:5 dr, >99:1 er Boo b. Pyrrolizine chlorination: o' NCS (1.1 equiv) CH2Cl2, 0 °C, 1 h

Scheme 2 Product derivatisation.

>95:5 dr. >99:1 er



Scheme 3 Proposed mechanism of isothiourea-catalysed Michael addition-lactonisation.

by intramolecular Michael-addition forms the new C–C bond and two stereocentres in the initial cyclisation step. Intermediate 45 is believed to be stabilised by a favourable non-bonding or electrostatic O to S interaction and this has been investigated computationally (vide infra). Subsequent lactonisation releases catalyst 24 and the polycyclic dihydropyranone product 21, which can be ring-opened upon addition of a nucleophilic amine or alcohol to give the isolated pyrrolizine carboxylate derivative.

On the basis of this mechanistic hypothesis, the origin of the diastereo- and enantioselective formation of the pyrrolizine dihydropyranone products in this BTM 24-catalysed cascade process was further probed through density functional theory (DFT) calculations. Calculations employed the M06-2X functional and investigated the cyclisation of the parent enone-acid 21 reacting in the presence of 24 in dichloromethane solvent from either the (E)- or (Z)-form of the ammonium enolate. Full computational details, including results obtained with the B3LYP functional, are supplied in the ESI.†29

Fig. 5 shows computed profiles for the alternative reactions of the (Z)-ammonium enolate, (Z)-45, to form the cis- and trans-isomers of pyrrolizine dihydropyranone 21. From (Z)-45 two low energy transition states were located for the Michael cyclisation step: $TS1_{cis}$ at +5.2 kcal mol^{-1} and $TS1_{trans}$ at +10.0 kcal mol⁻¹. The onward reaction via $TS1_{cis}$ (highlighted in red) leads to 46_{cis} at -0.6 kcal mol⁻¹ in which the newly formed C4-C9 bond is rather long at 1.62 Å. Attack of the enolate oxygen then permits formation of a zwitterionic tetrahedral dihydropyran intermediate 47cis via TS2cis at +6.3 kcal mol^{-1} . Facile dissociation of BTM 24 from 47_{cis} via TS3_{cis} forms 21_{cis} at -18.2 kcal mol⁻¹. An analogous series of steps (highlighted in blue) was also characterized for the formation of 21_{trans} at -13.0 kcal mol⁻¹. Formation of 21_{cis} is therefore both thermodynamically and kinetically favoured, as both TS1_{cis} and TS2_{cis} are significantly lower than TS1_{trans} en route to 21_{trans}. A third pathway starting from the (E)-ammonium enolate precursor, (E)-45 (+3.7 kcal mol⁻¹) and leading to the enantiomeric-trans product was also characterised, but has a

significantly higher barrier of 16.7 kcal mol⁻¹ and can therefore be discounted (see ESI†).

It is notable that all the computed structures for both reaction pathways in Fig. 5 from enolate (Z)-45 up to the final BTM dissociation steps feature a co-planar [1,5]-S...O motif with [1,5]-S...O interatom distances ranging from 2.60 Å to 2.82 Å. The importance of non-bonding S···O interactions has been widely recognized in structural and medicinal chemistry in the solid state (often described as a stabilising $n_{\rm O}$ to $\sigma_{\rm C-S}^*$ interaction),30 and has been previously recognised as a controlling element in enantioselective isothiourea-catalyzed reaction processes.31 Although the origin of this interaction is still under debate, 32 in all calculated structures the [1,5]-S...O distance is considerably below the sum of the van der Waals radii. Interestingly, the shortest distances calculated are found within (Z)-45 (2.64 Å) and zwitterionic intermediates (47_{cis} 2.60 Å and 47_{trans} 2.64 Å), presumably reflecting the formal negative charge on the oxygen atom within these structures. However, as this feature appears within all the structures in the computed reaction profiles, and varies in a similar way along those profiles, it does not appear to be a discriminating factor between the two pathways in Fig. 5.

Computed structures for the selectivity-determining transition states TS1cis and TS1trans are compared in Fig. 6. Both structures have similar distances (2.09 and 2.10 Å) for the forming C4···C9 bond, with Michael addition occurring anti-to the stereodirecting phenyl substituent and thus accounting for the observed enantioselectivity. In the favoured TS1cis arrangement the prostereogenic centres adopt an approximately eclipsed conformation with a H-C4···C9-H dihedral of 3°,

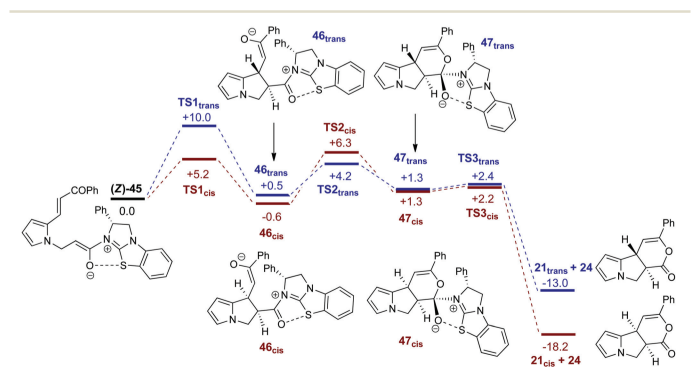


Fig. 5 Computed free energy profiles (M06-2X(CH₂Cl₂), kcal mol⁻¹) for the formation of cis- and trans-pyrrolizine dihydropyranones 21 from (Z)-ammonium enolate (Z)-45

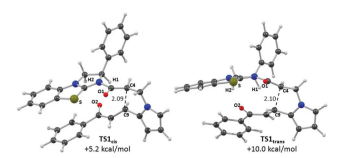


Fig. 6 Computed selectivity determining transition states TS1cis and TS1_{trans} with key atom labels and the forming C4...C9 distance highlighted in Å.

while in TS1_{trans} the corresponding H-C4···C9-H dihedral is 149.5°. Both transition states also exhibit short and approximately co-planar [1,5]-S...O1 contacts of ca. 2.7 Å between the BTM sulfur atom and the enolate oxygen. The near-parallel arrangement of the BTM moiety and the enone aryl substituent in TS1cis (inter-plane angle = 5.7°) appears set up for stabilising π -stacking interactions that should be captured by the M06-2X functional.²⁹ Indeed with the B3LYP functional (where such dispersion effects are not treated) the equivalent interplane angle = 28.1° and the preference for $TS1_{cis}$ is reduced to only 1.4 kcal mol⁻¹ (see ESI†). Also noticeable in **TS1**_{cis} are two short contacts between the forming oxy-anion on the enone (O2) and two C-H hydrogens upon the positively charged isothiouronium core (O1···H1 2.19 Å and O1···H2 2.37 Å). The enhanced acidity of these hydrogen atoms may facilitate some non-classical H-bonding and so confer greater stability on TS1cis over TS1trans where the analogous contacts are distinctly longer (O1···H1 2.72 Å and O1···H2 2.65 Å). While not particularly strong individually, these various effects likely combine to stabilise TS1cis over TS1trans and so account for the observed selectivity for the cis product.

Conclusions

In conclusion, an optimised and straightforward three-step synthetic route to a range of pyrrole-derived enone acid starting materials from readily available pyrrole-2-carboxaldehydes is delineated. The catalytic enantioselective synthesis of a range of cis-pyrrolizine carboxylate derivatives proceeds with outstanding stereocontrol (14 examples, >95:5 dr, >98:2 er) with benzotetramisole proving the optimal catalyst for this process. In situ ring-opening of the pyrrolizine dihydropyranone products with either MeOH or a range of amines leads to the desired products in excellent yield and enantioselectivity. Computation has been used to probe the factors leading to high stereocontrol in this reaction process, with the formation of the observed cis-stereoisomer predicted to be both kinetically and thermodynamically favoured. Further work from this laboratory will utilise this methodology for the synthesis of target molecules and probe the utility of isothioureas and other Lewis bases in enantioselective catalysis.

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