One-Pot Diastereoselective Synthesis of Pyrrolopiperazine-2,6-diones by a Ugi/Nucleophilic Substitution/N-Acylation Sequence

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ABSTRACT: The diastereoselective synthesis of two families of pyrrolopiperazine-2,6-diones is presented. These compounds were prepared by one-pot Ugi/nucleophilic substitution/N-acylation/debenzoylation/(elimination) sequences. This novel route provides straightforward access to a wide variety of pyrrolopiperazine-2,6-diones with high chemical yields and complete diastereoselectivities. The proposed synthetic strategy poses a significant improvement compared to the syntheses of pyrrolopiperazine-2,6-diones previously described, as it allows introduction of different substituents to the C4 position and the diastereoselective generation of a new stereogenic center on the bridgehead carbon (C8a).

The development of new and efficient syntheses of N-heterocycles is of paramount importance because of their relevance in the pharmaceutical and fine chemicals industries. Such importance relies on the fact that the structures of many of these systems are found in molecules displaying biological activity. This is the case of fused bicyclic piperazines, found in various natural products presenting antifungal, antibacterial, anxiolytic, or antitumoral activities. Among the different methodologies described to synthesize fused heterocycles, as pyrrolodiketopiperazines, multicomponent reactions (MCR) represent an interesting strategy to address the access to these systems.

Regarding pyrrolo-2,5-diketopiperazines, different methodologies based on the Ugi reaction have been described, for instance the Ugi/deprotection/cyclization sequence (UDC), the Ugi four-center three-component reaction (U-4C-3CR), and the Joullié–Ugi/postcondensation sequence. On the other hand, pyrrolo-2,6-diketopiperazines can be prepared through a Ugi five-center four-component reaction (U-5C-4CR) followed by a postcondensation step. In this case, the proline is employed as a doubly functionalized reactant, with the alcohol used as solvent acting as the fourth component. This leads to a 1,1′-diiminodicarboxylic derivative which affords the corresponding piperazine after treatment with a strong base.

Another interesting family of pyrrolodiketopiperazines are those in which the pyrrolo nucleus is a γ-lactam. However, although the enantiopure form of their 2,5-diketopiperazine derivatives can be easily synthesized starting from glutamic acid and other α-amino acids, the synthesis of enantiopure 2,6-diketopiperazine–pyrrolidinone systems is challenging. Indeed, although the methodologies based on the Ugi reaction are chemically efficient, their characteristic low diastereoselectivity represents an important drawback of these approaches. Thus, Ciufolini et al. have reported the synthesis of these pyrrolopiperazine-2,6-diones by a two-step sequence, a Ugi five-center four-component reaction (U-5C-4CR), followed by the cyclization promoted by trifluoroacetic acid, yielding products with diastereoselectivities ranging from 10:1 to 1.5:1 (Scheme 1a). Analogous systems have been described by Ugi, starting from γ-ketoacids and α-aminoesters, although the stereochemical aspect is, again, a major issue (Scheme 1b). Moreover, the synthesis of the non-substituted C8a analogue following this strategy is an extremely expensive alternative. To the best of our knowledge, only one highly diastereoselective synthetic methodology furnishing pyrrolopiperazine-2,6-diones has been reported that fulfills these conditions, a five-step route with no multicomponent reaction being involved. However, this path presents an important shortcoming: the substituent

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introduced in the C4 position of the molecule must be an aromatic moiety, because they are introduced through an electrophilic aromatic substitution (Scheme 1c).

Prompted by the promising biological activity of 2,6-diketopiperazine−pyrrolidinone fused systems,15 herein we present a novel synthetic strategy for the synthesis of this class of compounds that improves those currently described and overcomes some of their limitations. This route allows high chemical yields and an almost quantitative diastereoselectivity of diketopiperazine−pyrrolidinone derivatives bearing an unsubstituted C8a position and different substituents in the C4 position (Scheme 1d).

Given the interest in the α-alkylidene-γ-lactam scaffold in the development of compounds with biological properties as Michael acceptors toward bionucleophiles,16 α-methylidene-γ-lactams fused with 2,6-diketopiperazines, systems which have not been described so far, were also synthesized following a similar strategy.

The proposed synthetic path is based on a Ugi/nucleophilic substitution sequence, which leads to pyrrolidin-2-ones; a subsequent N-acylation followed by a debenzoylation step would lead to the desired pyrrolopiperazine-2,6-diones.

Initially, we employed the three α-aminoesters 1a−c to study the viability of the outlined route, together with 3-bromopropionic acid (2a), phenylglyoxal (3), and cyclohexyl isocyanide (4a) (Scheme 2). The reason behind the selection of phenylglyoxal as the carbonyl component is that the benzoyl group favors the enol tautomer,17 promoting cyclization through a C-alkylation reaction; besides, this group can be easily removed in a subsequent step by a retro-Claisen-like reaction.18 So, a solution of the commercial hydrochloride form of the corresponding α-aminoesters 1a−c was treated with potassium hydroxide in methanol for 10 min. Subsequently, 3-bromopropionic acid (2), phenylglyoxal (3), and cyclohexyl isocyanide (4a) were added, and the mixture was stirred for 24 h. The spontaneous cyclization of Ugi adducts 5 afforded the corresponding pyrrolidin-2-one 6 with a poor diastereoselectivity (Table 1). The glycine derivative required longer reaction times due to the lower solubility of the intermediate Ugi adduct 5a in methanol or treatment with a catalytic amount of cesium carbonate (Scheme 2).

Upon isolation of pyrrolidonones 6a−c, they were treated with cesium carbonate (2 equiv) in acetonitrile and heated to reflux for 1 h. Surprisingly, despite the low nucleophilicity of the carbonate anion, debenzoylated pyrrolopiperazine-2,6-


**Table 1. Results for the Synthesis of Pyrrolidin-2-ones 6**

| entry | 1 (R<sup>1</sup>) | 6 (%) | dr<sup>a</sup> |
|-------|------------------|-------|---------------|
| 1     | 1a (H)           | 6a (54)<sup>b</sup> | -             |
| 2     | 1b (Ph)          | 6b (41)<sup>b</sup>  | 51:49         |
| 3     | 1c (CH(CH<sub>3</sub>)<sub>3</sub>) | 6c (35)<sup>b</sup> | 50:50         |

<sup>a</sup>Determined by <sup>1</sup>H NMR spectroscopy in the reaction mixture. <sup>b</sup>Ugi adduct 5a was the only product observed after 24 h (78% yield).

Yield referred to the conversion of the Ugi adduct to the pyrrolidinone. Conversion is achieved by refluxing 5a in acetonitrile with a catalytic amount of cesium carbonate for 1 h. Chemical yield of the major diastereomer after purification by column chromatography. Chemical yield of the diastereomers mixture after purification by column chromatography.

**Scheme 3. Diastereoselective Synthesis of Pyrrolidinone-Fused Piperazine-2,6-diones 7 and X-ray Molecular Structure of Pyrrolopiperazine-2,6-dione 7c<sup>a</sup>**

![Scheme 3](image_url)

<sup>a</sup>The Olex2 plot is at the 30% probability level.

**Table 2. Results for the Synthesis of Pyrrolidinone-Fused Piperazine-2,6-diones 7 from Pyrrolidinones 6**

| entry | 6 (R<sup>1</sup>) | 7 (%) | dr<sup>a</sup> |
|-------|------------------|-------|---------------|
| 1     | 6a (H)           | 7a (84)<sup>b</sup> | -             |
| 2     | 6b (Ph)          | 7b (89)<sup>b</sup> | >95:5         |
| 3     | 6c (CH(CH<sub>3</sub>)<sub>3</sub>) | 7c (82)<sup>b</sup> | >95:5         |

<sup>a</sup>Determined by <sup>1</sup>H NMR spectroscopy in the reaction mixture. <sup>b</sup>Racemic mixture.

The <sup>1</sup>H NMR spectra of pyrolopiperazine-2,6-diones 7a–c show a triplet of triplets at ca. 4.5 ppm, corresponding to the proton of the cyclohexane’s methine group, which confirms that cyclization took place through the nitrogen atom coming from the isocyanide, together with a signal around 4.3 ppm, due to the proton linked to C8a; in addition, no signals attributable to the benzyl group are observed, in agreement with the occurrence of a debenzoylation process (see SI). The stability of the former is in agreement with the stereochemistry of the chiral center coming from the corresponding α-aminoester and would generate the most stable diastereoisomer of pyrolopiperazine-2,6-diones 7b,c as observed (Scheme 3, Table 2).

**Scheme 4. Proposed Mechanism for the Diastereoselective Synthesis of Pyrrolopiperazine-2,6-diones 7**

![Scheme 4](image_url)

The chemical yield was remarkably improved. The scope of this diastereoselective synthesis was extended to other α-aminoesters and isocyanides and, as expected, all pyrrolopiperazine-2,6-diones 7b,c were observed (Scheme 3, Table 2).

**Table 3. Diastereomeric Purity of Pyrrolidinone Derivatives 7**

| entry | 7 (R<sup>1</sup>) | dr<sup>a</sup> |
|-------|------------------|---------------|
| 1     | 7a (54)<sup>b</sup> | -             |
| 2     | 7b (65)<sup>b</sup> | 51:49         |
| 3     | 7c (53)<sup>b</sup> | 50:50         |

<sup>a</sup>Determined by <sup>1</sup>H NMR spectroscopy in the reaction mixture.

Yield referred to the conversion of the Ugi adduct to the pyrrolidinone. Conversion is achieved by refluxing 5a in acetonitrile with a catalytic amount of cesium carbonate for 1 h. Chemical yield of the major diastereomer after purification by column chromatography.

by the presence of water traces in the solvent employed in the reaction. In this way, the intramolecular N-acylation would take place, resulting in the diketopiperazine—pyrrolidinone fused system B, with the concomitant removal of the methoxide group. At this point, it is difficult to prevent the retro-Claisen reaction because this methoxide group would attack the benzyl group, yielding methyl benzoate, as could be detected in the raw product by <sup>1</sup>H NMR spectroscopy; furthermore, this step would be favored by the thermodynamic stability of enolate D. This would be the key step in the stereoreactive outcome, because the stereocenter of the former pyrrolidin-2-one 6 would have been destroyed. Protonation of enolate D would be controlled by the stereochemistry of the chiral center coming from the former pyrrolidin-2-one 6 would have been destroyed. The stability of the former is in agreement with the stereochemistry determined for these compounds in solution by NOESY experiments and with that observed for 7c in the solid state. This result could be explained by the steric hindrance exerted by the R<sup>1</sup> substituent, which constrained the bicyclic system in the (4S,8aR)-7c epimer.

Bearing in mind that the configuration of C5 in pyrrolidin-2-ones 6 does not determine the stereochemistry of the final pyrolopiperazine-2,6-diones 7, the synthetic route leading to them was performed in one pot, without isolating the pyrrolidin-2-one intermediates 6. Thus, initially, the synthesis of pyrrolidin-2-ones 6 in methanol was conducted and, after removing the solvent, without any further purification, the residue was dissolved in acetonitrile and treated with cesium carbonate. Analysis of the raw product by <sup>1</sup>H NMR spectroscopy revealed the formation of only one diastereomer of pyrolopiperazine-2,6-diones 7b,c, but above all, the global chemical yield was remarkably improved. The scope of this one-pot two-step sequence was then assayed with other α-aminoesters and isocyanides and, as expected, all pyrolopiperazine-2,6-diones 7b,c were obtained in high yields and in their enantiopure form except for glycine derivative 7a, which was isolated as a racemic mixture. (Scheme 5, Table 3). Finally, an attempt was made to carry out this sequence using a single solvent to simplify the experimental procedure, adding cesium...
carbonate to the reaction mixture without removing the solvent. Thus, although the use of methanol in both stages afforded complex mixtures, the use of acetonitrile allowed the isolation of pyrrolopiperazines 7. However, chemical yields were significantly lower when only one solvent was employed (Table 3, entries 3, 4 vs 5, 6).

Hence, it was confirmed that the purification and isolation of the pyrrolidin-2-one intermediate is not necessary, and that the reaction can be conducted in a one-pot two-step sequence, providing excellent stereochemical results. These results prompted us to apply this one-pot methodology to the synthesis of 2,6-diketopiperazines fused with α-methylidene-γ-lactams to introduce a methylene group as a convenient Michael acceptor toward bionucleophiles. Initially, for this purpose, 2-(bromomethyl)acrylic acid was selected, to introduce the methylidene group. So, the strategy previously introduced in the reactant and not generated during the reaction, the use of 3-bromo-2-(bromomethyl)propionic acid is undoubtedly a better choice as the overall yield is clearly higher. Therefore, the proposed one-pot two-step strategy proves itself a very powerful tool for the rapid and affordable diastereoselective synthesis of biologically relevant pyrrolopiperazine-2,6-diones.

In summary, a novel and completely diastereoselective one-pot methodology to synthesize pyrrolopiperazine-2,6-diones was presented, employing a simple Ugi/nucleophilic substitution/N-acylation/debenzylation/(elimination) sequence. This synthetic strategy represents a significant improvement over those described in the literature so far, as it allows introduction of a number of alkyl and aryl substituents to the C4 position, owing to the broad variety of commercially available α-aminoesters and the controlled generation of a new stereogenic center on the bridgehead carbon (C8a), starting from simple and inexpensive reagents, and reaching high chemical yields and quantitative diastereoselectivities, along with the possibility of introducing a methylene group as a Michael acceptor in the pyrrolidine ring.

| Table 3. Results for the Synthesis of Pyrrolidinone-Fused 2,6-Diketopiperazines 7 by a One-Pot Two-Step Sequence |
|---|
| entry | 1 (R1) | 4 (R2) | solvent | 7 (%) | dr |
| 1 | 1a (H) | 4a (CH2N2) | MeOH | 7a (78) | – |
| 2 | 1b (Ph) | 4a (CH2N2) | MeOH | 7b (86) | >95:5 |
| 3 | 1c (CH(CH3)2) | 4a (CH2N2) | MeOH | 7c (79) | >95:5 |
| 4 | 1c (CH(CH3)2) | 4a (CH2N2) | ACN | 7c (58) | >95:5 |
| 5 | 1d (CH2) | 4a (CH2N2) | MeOH | 7d (71) | >95:5 |
| 6 | 1d (CH2) | 4a (CH2N2) | ACN | 7d (56) | >95:5 |
| 7 | 1e (CH2Ph) | 4a (CH2N2) | MeOH | 7e (74) | >95:5 |
| 8 | 1b (Ph) | 4b (N(CH3)2) | MeOH | 7f (75) | >95:5 |
| 9 | 1c (CH(CH3)2) | 4b (N(CH3)2) | MeOH | 7g (81) | >95:5 |
| 10 | 1e (CH2Ph) | 4b (N(CH3)2) | MeOH | 7h (69) | >95:5 |

*aSolvent employed in the first stage. bChemical yield after purification. cDetermined by 1H NMR spectroscopy in the reaction mixture.

| Table 4. Results for the Synthesis of α-Methylidene-γ-lactam-Fused 2,6-Diketopiperazines 8 by a One-Pot Two-Step Sequence |
|---|
| entry | 1 (R1) | 8 (%) | dr |
| 1 | 1a (H) | 8a (33) | – |
| 2 | 1b (Ph) | 8b (77) | >95:5 |
| 3 | 1c (CH(CH3)2) | 8c (38) | >95:5 |
| 4 | 1d (CH2) | 8d (61) | >95:5 |
| 5 | 1e (CH2Ph) | 8e (71) | >95:5 |

*aChemical yield after purification. bDetermined by 1H NMR spectroscopy in the reaction mixture. cStarting from 2-(bromomethyl)acrylic acid. dStarting from 3-bromo-2-(bromomethyl)propionic acid.

**EXPERIMENTAL SECTION**

**General Methods.** All reagents and solvents were purchased and used without any further purification. Melting points are not corrected. Optical rotations were measured on a Zeiss D-7082 polarimeter in a 1 dm cell, and concentrations are given in g/100 mL. 1H and 13C NMR spectra were recorded in CDCl3 at 300 and 75 on the basis of NOESY experiments. Again, computational studies for the epimers of 8c on C8a confirmed the higher stability of the (4S,8aS) diastereoisomer, with a difference of energy of 5.19 kcal mol−1 between epimers (see SI).

Thus, although atom economy was higher employing 2-(bromomethyl)acrylic acid, meaning that the double bond was introduced in the reactant and not generated during the reaction, the use of 3-bromo-2-(bromomethyl)propionic acid is undoubtedly a better choice as the overall yield is clearly higher. Therefore, the proposed one-pot two-step strategy proves itself a very powerful tool for the rapid and affordable diastereoselective synthesis of biologically relevant pyrrolopiperazine-2,6-diones.
M.Hz, respectively, on a Varian Mercury 300 system and a Bruker Avance III HD system; DEPT-135 experiments were conducted to assign carbon-13 signals. Chemical shifts are reported in parts per million with respect to residual solvent protons and coupling constants in hertz. High resolution mass spectra were recorded on a 6545 Q-TOF Agilent LC-MS mass spectrometer (positive electro-spray ionization mode, ESI (+)). X-ray diffraction studies were performed on a Bruker D8 VENTURE diffractometer.

**Procedure for the Synthesis of Ugi Adduct 5a.** Glycine methyl ester hydrochloride 1a (0.250 g, 2 mmol, 1.0 equiv) was treated with potassium hydroxide (0.101 g, 1.8 mmol, 0.9 equiv) in methanol (10 mL), and the mixture was sonicated for 10 min. Subsequently, 3-bromopropionic acid (2) (0.304 g, 2 mmol, 1 equiv) was added to the mixture, followed by phenylglyoxal hydrate (3) (0.304 g, 2 mmol, 1 equiv) and cyclohexyl isocyanide (4) (0.218 g, 2 mmol, 1 equiv). The reaction mixture was stirred at room temperature for 24 h and the obtained precipitate isolated by vacuum filtration, washed with cold methanol, and dried in vacuo.

(E)-Methyl 2-(3-Bromo-3-(cyclohexylamino)-1-hydroxoy-3-oxo-1-phenylprop-1-en-2-yl)propanamidocetate (5a). Pink solid. 78% yield, 727 mg. Mp 128 °C. 1H NMR (300 MHz, CDCl3) δ: 7.72 (d, J = 7.4 Hz, 2H), 7.53 (t, J = 7.4 Hz, 1H), 7.39 (t, J = 7.4 Hz, 2H), 7.30–7.22 (m, 5H), 6.28 (d, J = 7.6 Hz, 1H, NH), 5.40 (s, 1H), 3.70 (s, 3H), 3.48–3.34 (m, 1H), 2.77–2.52 (m, 4H), 1.66–0.68 (m, 10H). 13C{1H} NMR (75 MHz, CDCl3) δ: 196.6 (Cq), 176.6 (Cq), 169.8 (Cq), 168.3 (Cq), 135.2 (Cq), 134.6 (Cq), 133.2 (CH2), 129.4 (CH2), 129.1 (CH2), 128.5 (CH2), 128.3 (CH2), 76.1 (Cq), 61.3 (CH2), 52.8 (CH2), 49.0 (CH2), 32.0 (CH2), 31.8 (CH2), 30.9 (CH2), 29.7 (CH2), 28.9 (CH2), 25.2 (CH2), 24.6 (CH2), 24.3 (CH2). HRMS (ESI) m/z: [M + H]+ cated for C23H28BrN2O5 467.1186; found 467.1183.

(25)-Methyl 2-(2-Benzoyl-2-(cyclohexylcarbamoyl)-5-oxopyrrolidin-1-yl)-2-phenylacetate (6b). Yellow oil (5:1 Hex/ EtOAc). 65% yield, 600 mg (as diastereomers mixture). 1H NMR (300 MHz, CDCl3) δ: 7.72 (d, J = 7.4 Hz, 2H), 7.53 (t, J = 7.4 Hz, 1H), 7.39 (t, J = 7.4 Hz, 2H), 7.30–7.22 (m, 5H), 6.28 (d, J = 7.6 Hz, 1H, NH), 5.40 (s, 1H), 3.70 (s, 3H), 3.48–3.34 (m, 1H), 2.77–2.52 (m, 4H), 1.66–0.68 (m, 10H). 13C{1H} NMR (75 MHz, CDCl3) δ: 196.6 (Cq), 176.6 (Cq), 169.8 (Cq), 168.3 (Cq), 135.2 (Cq), 134.6 (Cq), 133.2 (CH2), 129.4 (CH2), 129.1 (CH2), 128.5 (CH2), 128.3 (CH2), 76.1 (Cq), 61.3 (CH2), 52.8 (CH2), 49.0 (CH2), 32.0 (CH2), 31.8 (CH2), 30.9 (CH2), 29.7 (CH2), 28.9 (CH2), 25.2 (CH2), 24.6 (CH2), 24.3 (CH2). HRMS (ESI) m/z: [M + H]+ cated for C23H28BrN2O5 467.1186; found 467.1183.

**General Procedure for the Synthesis of Pyrrolidinone-Fused 2,6-Diketopiperazines.** Method A. A mixture of pyrrolidin-2-ones 6a–e (0.2 mmol, 1 equiv) and cesium carbonate (0.4 mmol, 2 equiv) in acetonitrile (3 mL) was stirred and heated to reflux with a heating block for 1 h. Subsequently, the solvent was removed under reduced pressure and the crude product dissolved in chloroform. This solution was washed with water and the organic layer dried over sodium sulfate, filtered, and concentrated to dryness. The residue was purified by column chromatography, employing SiO2 as stationary phase and a hexane/ethyl acetate mixture as eluent. Method B. The corresponding α-aminoester hydrochloride 1a–e (2 mmol, 1 equiv) was treated with potassium hydroxide (1.8 mmol, 0.9 equiv) in methanol (10 mL) and the resulting suspension sonicated for 10 min. Subsequently, 3-bromopropionic acid (2) (2 mmol, 1 equiv) was added to the reaction mixture, followed by phenylglyoxal (3 mmol, 1 equiv) and isocyanide 4a,b (2 mmol, 1 equiv). The reaction mixture was stirred at room temperature for 24 h, and the solvent was removed under reduced pressure. The raw product was dissolved in acetonitrile (10 mL), treated with cesium carbonate (4 mmol, 2 equiv), heated to reflux with a heating block for 1 h, and henceforth treated as indicated in method A. Method C. The corresponding α-aminoester hydrochloride 1a–e (2 mmol, 1 equiv) was treated with potassium hydroxide (1.8 mmol, 0.9 equiv) in acetonitrile (10 mL) and the resulting suspension sonicated for 10 min. Subsequently, 3-bromopropionic acid (2) (2 mmol, 1 equiv) was added to the reaction mixture, followed by phenylglyoxal (3 mmol, 1 equiv) and isocyanide 4a,b (2 mmol, 1 equiv). The reaction mixture was stirred at room temperature for 24 h, and then cesium carbonate (4 mmol) was added. The reaction mixture was stirred and heated to reflux with a heating block for 1 h. From there, the reaction was conducted according to the procedure described for method A.

2-Cyclohexylidihydropyrrolo[1,2-a]pyrazine-1,3,6-(2H,4H,7H)-trione (7a). Brown oil (2:1 H2O/EtOAc). 78% yield, 390 mg. 1H NMR (300 MHz, CDCl3) δ: 4.84 (d, J = 18.6 Hz, 1H, 4.47 (dt, J = 12.3, 8.8 Hz, 1H), 4.30–4.21 (m, 1H, 3.78 (d, J = 16.8 Hz, 1H), 2.59–2.52 (m, 2H, 2.19–2.14 (m, 2H), 1.91–1.83 (m, 10H). 13C{1H} NMR (75 MHz, CDCl3) δ: 196.6 (Cq), 175.7 (Cq), 170.3 (Cq), 167.8 (Cq), 134.7 (Cq), 133.3 (Cq), 129.3 (CH3), 128.5 (CH3), 76.7 (CH3), 52.5 (CH2), 49.2 (CH2), 44.9 (CH2), 32.4 (CH2), 32.2 (CH2), 29.4 (CH2), 28.5 (CH2), 25.3 (CH2), 24.9 (CH2), 24.8 (CH2). HRMS (ESI) m/z: [M + Na]+ cated for C15H13N2O4Na 409.1739; found 409.1733.

(45,8aS)-2-Cyclohexyl-4-pentylidihydropyrrolo[1,2-a]pyrazine-1,3,6(2H,4H,7H)-trione (7b). Pale yellow oil (2:1 Hex/EtOAc). 86% yield, 561 mg. [α]D = −149.8° (c = 0.26, MeOH). 1H NMR (300 MHz, CDCl3) δ: 7.45–7.21 (m, 1H), 6.03 (s, 1H, 4.61 (tt, J = 12.2, 3.7 Hz, 1H), 4.23–4.13 (m, 1H, 2.64–2.19 (m, 4H, 1.97–1.03 (m, 10H). 13C{1H} NMR (75 MHz, CDCl3) δ: 173.0.
(4S,8aS)-2-Cyclohexyl-4-isopropylidihydropyrrolo[1,2-a]pyrazine-3,6(2H,4H)-trione (7c). White solid (2:1 Hex/EtOAc). 79% yield, 461 mg. Mp 73–75 °C. [α]D = +2.65° (c = 2, MeOH). 1H NMR (300 MHz, CDCl3) δ: 4.87–4.82 (m, 3H), 2.60–1.97 (m, 5H), 1.74–0.94 (m, 10H), 0.99 (d, J = 6.8 Hz, 3H). 13C(H) NMR (75 MHz, CDCl3) δ: 173.8 (Cq), 171.2 (Cq), 169.5 (Cq), 156.7 (Cq), 145.7 (Cq), 141.3 (Cq), 133.8 (Cq), 126.8 (Cq), 125.8 (Cq), 123.2 (CH), 121.7 (CH), 121.4 (CH), 53.4 (CH), 29.0 (CH2), 28.7 (CH2), 26.3 (CH2), 26.2 (CH2), 25.1 (CH2), 23.0 (CH3), 19.4 (CH3), 19.3 (CH3). HRMS (ESI) m/z: [M + H]+ calc for C21H22N2O3 341.1540; found 341.1549.

(4S,8aS)-5-Cyclohexyl-5-methyl-4-phenylidihydropyrrolo[1,2-a]pyrazine-3,6(2H,4H)-trione (7a). Yellow oil (3:1 Hex/EtOAc). 79% yield, 521 mg. [α]D = –63.0° (c = 0.11, acetone). 1H NMR (300 MHz, CDCl3) δ: 7.42–7.27 (m, 7H), 6.16 (t, J = 2.8 Hz, 1H), 6.13 (s, 1H), 5.53 (t, J = 2.4 Hz, 1H), 4.62 (t, J = 12.2, 3.7 Hz, 1H), 4.19 (dd, J = 8.8, 5.2 Hz, 1H), 3.25–3.06 (m, 2H), 1.88–1.18 (m, 10H). 13C(H) NMR (75 MHz, CDCl3) δ: 170.4 (Cq), 167.7 (Cq), 166.1 (Cq), 136.2 (Cq), 118.6 (CH), 54.5 (CH), 54.3 (CH), 44.6 (CH), 29.3 (CH), 28.7 (CH), 27.7 (CH2), 26.4 (CH2), 26.3 (CH2), 25.2 (CH3). HRMS (ESI) m/z: [M + H]+ calc for C22H23N2O3 363.1709, found 363.1710.

(4S,8aS)-2-Cyclohexyl-4-isopropylidihydropyrrolo[1,2-a]pyrazine-3,6(2H,4H)-trione (7b). Orange oil (3:1 Hex/EtOAc). 85% yield, 446 mg. 1H NMR (300 MHz, CDCl3) δ: 6.11 (t, J = 2.7 Hz, 1H), 5.50 (t, J = 2.6 Hz, 1H), 4.96 (d, J = 18.7 Hz, 1H), 4.49 (t, J = 11.8, 3.7 Hz, 1H), 4.28 (t, J = 6.7 Hz, 1H), 3.90 (d, J = 18.8 Hz, 1H), 3.19 (dt, J = 5.4, 2.7 Hz, 2H), 2.26–0.82 (m, 10H). 13C(H) NMR (75 MHz, CDCl3) δ: 170.4 (Cq), 167.0 (Cq), 166.1 (Cq), 136.2 (Cq), 118.6 (CH), 54.5 (CH), 54.3 (CH), 44.6 (CH), 29.3 (CH), 28.7 (CH), 27.7 (CH2), 26.4 (CH2), 26.3 (CH2), 25.2 (CH3). HRMS (ESI) m/z: [M + H]+ calc for C22H23N2O3 363.1709, found 363.1710.

(4S,8aS)-2-Cyclohexyl-4-isopropylidihydropyrrolo[1,2-a]pyrazine-3,6(2H,4H)-trione (7c). White solid (3:1 Hex/EtOAc). 75% yield, 455 mg. Mp 143–145 °C. [α]D = –103.3° (c = 0.16, MeOH). 1H NMR (300 MHz, CDCl3) δ: 7.30–7.23 (m, 3H), 7.14–7.11 (m, 2H), 5.04 (t, J = 5.3 Hz, 1H), 4.49 (t, J = 12.2, 3.8 Hz, 1H), 3.41–3.33 (m, 2H), 3.20 (dd, J = 13.8, 5.5 Hz, 1H), 2.41–1.08 (m, 14H). 13C(H) NMR (75 MHz, CDCl3) δ: 173.3 (Cq), 170.8 (Cq), 170.4 (Cq), 135.3 (Cq), 129.2 (Cq), 129.0 (Cq), 129.7 (Cq), 129.6 (Cq), 127.6 (CH), 127.5 (CH), 55.4 (CH), 55.1 (CH), 54.1 (CH), 37.4 (CH), 29.4 (CH), 29.1 (CH), 28.5 (CH2), 26.3 (CH2), 26.2 (CH2), 25.2 (CH2), 22.0 (CH3). HRMS (ESI) m/z: [M + H]+ calc for C21H22N2O3 341.1540; found 341.1549.
Light yellow oil (3:1 Hex/EtOAc). 71% yield, 500 mg. 

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