Organic & Biomolecular Chemistry



View Article Online

PAPER

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Cite this: Org. Biomol. Chem., 2024, **22**, 3230

Stereodivergent synthesis of 2-oxooligopyrrolidines by an iterative coupling strategy[†]

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Natural linear polyamines play diverse roles in physiological processes by interacting with receptors at the cellular level. Herein, we describe the stereodivergent synthesis of oligopyrrolidines, which are conformationally constrained polyamines. We synthesized dimeric and trimeric 2-oxo-oligopyrrolidines using an iterative coupling strategy. The key to our success is an iridium-catalyzed *trans/cis*-selective nucleophilic addition and subsequent *threo/erythro*-stereoselective reduction. The synthesized pyrrolidines show varying cytotoxicities against a human cancer cell line depending on the number of rings and their stereochemistry.

Received 4th March 2024, Accepted 26th March 2024 DOI: 10.1039/d4ob00350k

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Introduction

Polyamines are organic cations containing a repeating structure of acyclic alkyl amines. Natural polyamines such as putrescine (1), spermidine (2) and spermine (3) exhibit a broad range of physiological functions such as cell replication and the modulation of gene expression (Scheme 1a).¹ Moreover, polyamines have attracted attention as "privileged" templates to induce versatile biological activities (Scheme 1b).² In 1988, the Melchiorre group documented tetraamine disulfide benextramine (4) as an irreversible α -adrenoreceptor antagonist.^{2a} Subsequently, 4 was also found to bind to other receptors such as nicotinic receptors and muscarinic receptors. These studies indicated that synthetic polyamines are attractive candidates in drug discovery. Woster and co-workers reported antibacterial polyamines targeting bacterial membranes.³ Compound 5 exhibits bactericidal activity against various Gram-positive bacteria, including methicillin-resistant Staphylococcus aureus (MRSA) due to electrostatic interaction between the positively charged polyamines and the negatively charged bacterial membrane. Bissati and co-workers reported thiourea-based polyamine 6 as being potent against the malaria parasite.⁴

Cyclic polyamines with stereogenic centers are promising structural motifs for controlling the three-dimensional

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arrangements of protonated amine groups, which is crucial for recognizing the target polyanion structures, as seen in DNA and mRNA (Scheme 1c).⁵ In 1997, Ganesh's group reported the first example using spermine analogues 7 and 8, which have a pyrrolidine backbone. Their synthetic analogues exhibited stronger binding to the DNA triplex than natural spermine.⁶ Müller, Koert and co-workers synthesized the trans-threo-trans oligopyrrolidines 9 and 10, which have helical conformations due to the rigid structure of pyrrolidines.⁷ The rate of ssRNA cleavage by oligopyrrolidine 10 was faster than that of natural spermine. Higuchi and colleagues reported pentamine-based inhibitors of lysine-specific demethylase 1 (LSD1) and LSD2 such as 11, which are promising potent anticancer agents.⁸ These pentamines exhibit better LSD1- and 2-inhibitory activities than simple linear polyamines, probably through conformational restriction with the three trans-cyclopentane units contained in the six stereogenic centers. They also found that the LSD-inhibitory activities depended on the stereochemistry of each artificial polyamine.

Results and discussion

To obtain three-dimensionally constrained polyamines, we took interest in 2-oxo-oligopyrrolidines **12** possessing diverse stereochemistry at C2'–C5 on the cyclic amines (Scheme 2). Casiraghi and coworkers documented pioneering reports on a building-block-based synthesis of 2-oxo-oligopyrrolidines by a coupling reaction using *N*-Boc-lactam **13** and 2-silyloxypyrrole **16** (Scheme 2a).⁹ The method began with LiEt₃BH reduction of *N*-Boc-lactam **13**, followed by acid-mediated formation of *N*,*O*-acetal **14**. Addition of TBSOTf to **14** at -90 °C formed the

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[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d4ob00350k



N-acyliminium ion 15, which underwent the vinylogous Mannich reaction^{10,11} with 16 to give trans-erythro and ciserythro 2-oxo-bispyrrolidines 17α and 17β . Pd/C-catalyzed hydrogenation of the resulting bispyrrolidines 17 provided 18, which could be converted to oligo-heterocyclic compounds by the same four-step sequence. Their pioneering work successfully demonstrated the utility of the iterative approach. However, some synthetic issues remained unsolved. First, reduction of amide carbonyls required a stoichiometric amount of a strong reductant. Second, transformation of the resulting hemiaminal to N,O-acetal 14 prior to the vinylogous Mannich reaction was essential to form the unstable N-acyliminium ion 15 under acidic conditions. In addition, only two out of the four possible stereoisomers were obtained by their method. To resolve these issues, we envisioned an iterative approach12 based on N-methoxylactams.13,14 Our method begins with an iridium-catalyzed reductive vinylogous Mannich reaction between N-methoxylactam 19 and 2-silyloxy-

pyrrole 22.^{15,16} Iridium-catalyzed hydrosilylation of **19** would give N,O-acetal 20. Subsequent addition of an acid to 20 forms the N-methoxyiminium ion 21, which could undergo a cis/ trans-selective vinylogous Mannich reaction with 22 to provide enamide 23. We employed N-methoxylactams instead of N-Boc-lactams due to the following reasons. N-Methoxylactams undergo Ir-catalyzed hydrosilylation under mild conditions, while *N*-Boc-lactams do not.¹⁷ Formation of the N-methoxyiminium ion 21 is feasible by direct addition of an acid without isolation of N,O-acetal 20. Thus, a reductive vinylogous Mannich reaction would give enamide 23 in a one-pot process. In addition, N-methoxyiminium ions are known to be more electrophilic than N-alkyliminium ions such as the N-benzyliminium ion, leading to a high yielding vinylogous Mannich reaction.¹⁸ The resulting enamide 23 could be converted to threo- or erythro-2-oxo-bispyrrolidines 24 by stereoselective reduction. These sequences would enable the stereodivergent synthesis of all four possible diastereomers of 24. The key to success is the development of appropriate reaction conditions to control the cis/trans configurations on the pyrrolidine ring, and the threo/erythro configurations on the rotational bond. Ultimately, we found that the stereocontrol of trans- or cis-selectivity could be achieved using kinetic or thermodynamic conditions, respectively. Subsequent threo/ erythro-selectivity was controlled by the choice of reducing agent. Iterative application of this process enabled the construction of 2-oxo-oligopyrrolidines 12. In this paper, we demonstrate the synthesis of all four possible diastereomers of 2-oxo-bispyrrolidines 24. The method is applicable to the synthesis of 2-oxo-trispyrrolidines 25 en route to 2-oxo-oligopyrrolidines 12.

Our stereodivergent nucleophilic addition was evaluated using N-methoxylactam 19 and 2-silyloxypyrrole 22 (Scheme 3). After extensive screening, trans-selective nucleophilic addition under kinetic control was realized as follows. Treatment of a solution of 13 in CHCl₃ with (Me₂HSi)₂O and a catalytic amount of IrCl(CO)(PPh₃)₂ initiated hydrosilylation to generate the N-methoxyiminium ion 21. Subsequently, the addition of Sc(OTf)₃ and 2-silyloxypyrrole 22 at -60 °C generated the N-methoxyiminium ion 26. Aqueous work-up afforded a mixture of four lactams and two enamides 27-30.19 Successful trans-selectivity is dependent on the nature of the solvent used, and $CHCl_3$ was found to be the best solvent [trans-(27 + 28: cis-(29 + 30) = 4:1]. Although the vinylogous Mannich reaction produced a mixture of six diastereomers, these diastereomers converged to two enamides 28 and 30 through deprotonation with NaHMDS, and α -selective protonation with AcOH at -78 °C. Thus, the reductive vinylogous Mannich reaction and subsequent α -selective protonation gave enamides 28 and 30 in a trans-selective fashion (73% for 2 steps, 28:30 = 3.7:1).20

With the optimized conditions for the *trans*-selective nucleophilic addition reaction in hand, we examined the *cis*-selective reaction as shown in Scheme 4. After iridium-catalyzed hydrosilylation of **19**, the vinylogous Mannich reaction using MeCN as a co-solvent at room temperature was found to



show *cis*-selectivity. The resulting mixture converged to two enamides **30** and **28** by a regioselective deprotonation/protonation sequence (51%, 2 steps, *cis/trans* = 5.7:1).²⁰

To elucidate the mechanism leading to the *cis*-selectivity, we investigated the effect of reaction temperature (Table 1). After iridium-catalyzed hydrosilylation of **19** in toluene at room temperature, addition of **22** in the presence of 7 mol% of $Sc(OTf)_3$ at -78 °C promoted the vinylogous Mannich reaction. Interestingly, a mixture of four lactams and two enamides **27–30** was obtained with slight *trans*-selectivity [99%, *cis*-(**29** + **30**): *trans*-(**27** + **28**) = 1:1.7] (entry 1). The vinylogous Mannich reaction at room temperature was found to be *cis*-selective [31%, *cis*-(**29** + **30**): *trans*-(**27** + **28**) = 3.1:1], although some decomposition was observed (entry 2). To increase both yield and selectivity, the addition of MeCN was proved to be effective [62%, *cis*-(**29** + **30**): *trans*-(**27** + **28**) = 5.5:1]. These results indicated that the *cis*-selectivity was achieved under thermodynamic conditions.

To clarify the thermodynamic nature, the reversibility of the vinylogous Mannich reaction was further confirmed by a crossover experiment (Scheme 5). A solution of **27–30** in toluene/ MeCN was treated with 1-benzyloxy-2-silyloxypyrrole **31** in the presence of a catalytic amount of Sc(OTf)₃. Subsequent isomerization to enamides through the deprotonation/protonation protocol gave enamide **33** [44% (2 steps), *trans* : *cis* = 1 : 2.0], along with a mixture of **28** and **30** [44% (2 steps), **28** : **30** = 1 : 7.0].These results clearly indicated that enamide **33** was formed by a retro-Mannich/Mannich reaction under equilibrium conditions.

Having established the *cis/trans*-selective vinylogous Mannich reaction, we investigated the stereoselective reduction of trans-enamide 28 and cis-enamide 30 (Scheme 6). Hydrogenation of trans-enamide 28 with Rh/Al₂O₃ formed the 2-oxo-bispyrrolidines, favoring erythro lactam 34 over threo lactam 35 (Scheme 6a, 76%, 34:35 = 3.2:1).²⁰ In contrast, hydride reduction under acidic conditions reversed this stereoselectivity. Thus, treatment of trans-enamide 28 with NaBH3CN and HCl in the presence of 15-crown-5 gave a mixture of 2-oxobispyrrolidines 34 and 35 with threo selectivity (78%, 34:35 = 1:1.8).^{20,21} cis-Enamide 30 showed similar stereoselectivity (Scheme 6b). In contrast, Rh/Al₂O₃-catalyzed hydrogenation of cis-enamide 30 predominantly provided threo-lactam 36 (70%, 36:37 = 2.9:1²⁰ and hydride reduction with NaBH₃CN and TFA produced erythro lactam 37 as a major product (66%, 36:37 = 1:2.0). Reduction of *trans-erythro* lactam 34 was achieved by hydrosilylation with 1 mol% of IrCl(CO)(PPh₃)₂ and $(Me_2HSi)_2O$, followed by the addition of Sc(OTf)₃ to give bispyrrolidine 38.

We demonstrated the feasibility of iterative assembly of a pyrrolidine unit by investigating the stereoselective synthesis of 2-oxo-trispyrrolidines **41** and **42** (Scheme 7). Under the optimized kinetic conditions for *trans*-selective addition, *trans*-erythro lactam **34** was converted to *trans* enamide **39**, along with the formation of minor *cis* enamide **40** (66% (2 steps), **39**: **40** = 4.5:1).²⁰ Rh/Al₂O₃-catalyzed hydrogenation promoted erythro-selective reduction of enamide **39**, providing erythro lactam **41** and *threo* lactam **42** in 75% yield (**41**: **42** = 4.9:1).²⁰ The lactam carbonyl of 2-oxo-trispyrrolidine **41** was success-



Scheme 3 trans-Selective vinylogous Mannich reaction under kinetic conditions.



Table 1 Vinylogous Mannich reaction under thermodynamic conditions^a



Entry	Temp.	Solv.	cis-(29 + 30) : trans-(27 + 28)	Combined yield (%)
1	−78 °C	None	1:1.7	99
2	rt	None	3.1:1	31
3	rt	MeCN	5.5:1	62

^a Optimal reaction conditions: **19** (1 equiv.), IrCl(CO)(PPh₃)₂ (1 mol%), (Me₂HSi)₂O (1.4 equiv.), toluene, rt, 1 h; Sc(OTf)₃ (7 mol%), 22 (3.0 equiv.), co-solvent, 5 h.



Scheme 4 *cis*-Selective vinylogous Mannich reaction and regioselective deprotonation/protonation.

fully reduced by iridium-catalyzed hydrosilylation followed by

and 2-oxo-trispyrrolidines against Jurkat cells (human T-cell

The antiproliferative activity of the series of mono-, bis-,

leukemia cells) is summarized in Table 2. Comparison of the activity of 2-oxo-pyrrolidine 19, 2-oxo-bispyrrolidines 34-37, and 2-oxo-trispyrrolidines 41 and 42 revealed clear relationships between the number of rings and the compound's antiproliferative activity. 2-Oxo-bispyrrolidines 34-37 were found to be more potent than 2-oxo-pyrrolidine 19, with 2-oxo-trispyrrolidines 42 displaying the strongest activity, approximately 10-fold when compared to monomeric 2-oxo-pyrrolidine 19.

0N-78

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the addition of $Sc(OTf)_3$ to give trispyrrolidine 43.



Scheme 6 Synthesis of four stereoisomers of 2-oxo-bispyrrolidines.

Table 2 IC₅₀ values of synthetic compounds^a

IC_{50} [μM]
15.5
2.00
44.1

^{*a*} The antiproliferative activity against Jurkat cells was evaluated by the standard MTT assay. Cells were treated for 72 h with increasing concentrations of the evaluated compounds.

Stereochemistry played an important role in the antiproliferative activity of the compounds. Of the 2-oxo-bispyrrolidines **34–37**, *trans-threo* lactam **35** exhibited the highest activity. A similar tendency was observed for the trimeric compounds, although only two diastereomers were tested. Thus, the activity of 2-oxo-trispyrrolidine **42** with a *trans-threo* configuration at C2–C5' was 8-fold higher than that of **34** with a *trans-erythro* configuration. Interestingly, 2-oxo-oligopyrrolidines **34** and **41** exhibited stronger activity than the corresponding **38** and **43**, their fully reduced counterparts.

Conclusion

In conclusion, the results reported here provide an iterative and stereodivergent method to construct 2-oxo-oligopyrrolidines. *trans/cis*-Selectivity in the nucleophilic addition to amide carbonyls was controlled by choosing either kinetic or thermodynamic conditions. Subsequent *erythro/threo*-selective reduction of the enamides was realized by hydrogenation with Rh/Al₂O₃ or hydride reduction with NaBH₃CN. The developed conditions enabled the stereodivergent synthesis of all four possible diastereomers of the 2-oxo-bispyrrolidines. Moreover, the established method was successfully applied to the synthesis of the trimeric derivatives. The antiproliferative activity



Scheme 7 Stereoselective synthesis of 2-oxo-trispyrrolidines 41 and 42.

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of the synthetic lactams against human cancer cells revealed the role of the number of rings and their stereochemistry in the cytotoxicity of the compounds. Detailed investigations of the antiproliferative activities and RNA binding assays of compounds are underway.

Author contributions

Conceptualization: Y.S., N.C., T.O., and T.S.; data curation: Y. S., K.T., M.F., and T.M.; formal analysis: M.F., T.M.; funding acquisition: N.C., T.S.; investigation: Y.S., K.T., M.F., Su.S., K. S., and T.M.; methodology: Y.S., M.F., T.M.; project administration: T.O., T.S.; resource: Si.S., N.C., T.S.; software: N/A; supervision: Si.S., N.C., T.O., T.S.; validation: N/A; visualization: Y.S., T.O., T.S.; writing – original draft: Y.S., T.O.; writing – review and editing: Si.S., N.C., T.O., T.S.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research was supported by a Grant-in-Aid for Scientific Research (B) from MEXT (22H02084), the Tobe Maki Foundation, the JGC-S Scholarship Foundation, the Kato Memorial Bioscience Foundation, the Sasakawa Scientific Research Grant from The Japan Science Society (2022-0613), and the Amano Institute of Technology Foundation. The financial support of the Yoshida Scholarship Foundation to Y. Soda is gratefully acknowledged.

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- 19 α -Protonation of the oxypyrrole moiety in compound 19 provides *trans* or *cis*-enamide (28 and 30), and γ -protonation gave two *trans* or *cis*- α , β -unsaturated compounds each (27 α , 29 α , 29 β and 29 β). The details are described in the ESI.[†]
- 20 These diastereomers were separated by HPLC, see the ESI for details.†
- 21 Their stereochemistry was determined by NOESY measurement of the converted cyclic urea (see the ESI).†