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Bioorganic & Medicinal Chemistry xxx (2018) xxx-xxx

Contents lists available at ScienceDirect



Bioorganic & Medicinal Chemistry

journal homepage: www.elsevier.com/locate/bmc

Design and synthesis of a fragment set based on twisted bicyclic lactams

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ARTICLE INFO

Article history: Received 17 January 2018 Revised 15 February 2018 Accepted 16 February 2018 Available online xxxx

Keywords: Fragments Drug discovery Twisted amides Shape diversity

ABSTRACT

Current fragment sets tend to be dominated by flatter molecules, and their shape diversity does not reflect that of the fragments that are theoretically possible. The design and synthesis of a set of bridged fragments containing a bridgehead nitrogen is described. Many of these fragments contain twisted lactams whose modulated electronic properties may present unusual opportunities for interaction with target proteins. The demonstrated novelty, three-dimensionality and molecular properties of the set of 22 fragments may provide valuable, and highly distinctive, starting points for fragment-based drug discovery.

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1. Introduction

Over the last 15–20 years, fragment-based discovery has become a mainstream approach in medicinal chemistry.¹ Consequently, guidelines have been formulated to facilitate the assembly of fragment sets that target diverse relevant chemical space.² However, current fragment sets tend³ to be dominated by flatter (generally heteroaromatic) molecules whose shape diversity is not representative of the fragments that are theoretically possible.⁴ As a result, significant effort has been invested in the design of fragment sets with higher shape diversity.^{3a,d} More three dimensional (3D) fragments have inherently higher molecular complexity which has been argued⁵ to result in lower hit rates in fragment screens.⁶ Yet, such fragments are likely to offer distinctive opportunities for subsequent growth along specific vectors.

Here, we describe the design and synthesis of a fragment set that is based on a number of bicyclic ring systems containing a bridgehead nitrogen atom. Such ring systems are substructures within the frameworks of a diverse range of alkaloid natural products (see Fig. 1 for examples⁷) which can serve as an inspiration for drug discovery.⁸ The geometric constraints imposed by these ring systems can perturb functional group properties and characteristics by restricting or preventing the overlap of the bridgehead nitrogen lone pair with an adjacent π -system, and in the case of twisted amides, parameters have been developed to describe the

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https://doi.org/10.1016/j.bmc.2018.02.027

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extent of deformation.^{9,10} In extremely twisted amides, for example, the electronic properties of the functional group are more reminiscent of those of unconjugated amino ketones (for examples, see Fig. 2): this is reflected in the electrophilic reactivity of the carbonyl group^{10,11} and in the full¹² or partial¹³ *N*-protonation of the amide in contrast to the *O*-protonation observed with non-twisted amides. Despite the unusual and distinctive hydrogenbonding opportunities offered by such motifs, bicyclic lactams have barely been explored in a medicinal chemistry context, and may therefore provide new opportunities in drug discovery.¹⁴

2. Results and discussion

Our synthetic approach to bridged bicyclic fragments is shown in Scheme 1. Thus, $3-(\omega$ -carboxylate)-substituted piperidines **1** would be lactamised to yield a range of bridged lactams **2** (with variable n and R). It was envisaged that, with appropriate choice of substituent, addition of ring(s) (\rightarrow **3**) or functionalisation of the twisted amide (\rightarrow **4**) might be possible to yield related scaffolds. Finally, decoration would yield corresponding fragments for addition to a screening set.

2.1. Synthesis of bridged bicyclic lactams

Initially, a range of $3-(\omega-\text{carboxylate})$ -substituted piperidine substrates **7** was prepared (Scheme 2 and Table 1). The reductive aminations between the piperidin-3-one **5** and the amino esters **6a**–**d** gave the corresponding 3-amino piperidines **7a**–**d**. Alternatively, condensation of the piperidin-4-one **9** with pyrrolidine gave an enamine that was reacted directly with ethyl acrylate to give

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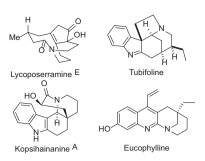
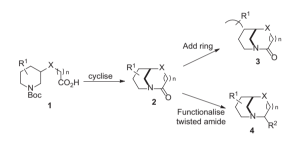


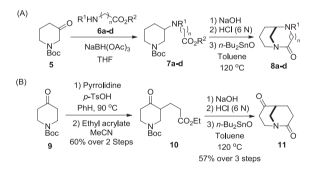
Fig. 1. Ring systems with bridgehead nitrogens embedded in alkaloids.



Fig. 2. Unstrained amide 12 and Kirby's most twisted amide 13.



Scheme 1. Envisaged synthetic approach to bridged bicyclic fragments.



Scheme 2. Synthesis of bridged bicylic lactams (see Table 1).

the 3-substituted piperidin-4-one **10**. Boc deprotection of **7a–d** and **10**, and ester hydrolysis, was followed by Bu_2SnO -mediated cyclisation to give the corresponding lactams **8a–d** and **11**.¹⁵ The approach enabled the synthesis of both bicyclo[3.3.1]nonane and

Table 1	
Synthesis of bridged bicyclic lactams (see Scheme 2)	

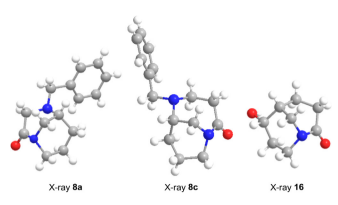


Fig. 3. X-ray crystal structures of 8a, 8c and 16.

bicyclo[4.3.1]decane bicyclic ring systems in moderate yield. Unfortunately, the approach did not enable the synthesis of the alternate bicyclo[3.2.2]nonane, bicyclo[4.2.2]decane and bicyclo [4.1.1]octane ring systems, presumably owing to increased ring strain in these species. Furthermore, deprotection of benzyl-protected **8a** and **8c** and Cbz-protected **8d** was not successful under a range of hydrogenation conditions (see Fig 3).

The distortion¹⁰ of the bridged bicyclic lactams **8a–d** and **11**, and their derivatives (see below), was assessed. ¹³C NMR spectroscopy is a valuable technique for investigating amide distortion, with the chemical shift of the carbonyl carbon serving as a sensitive probe of strain (see Fig. 2; unstrained δ -lactam **12**, δ_C : 165 ppm; Kirby's most twisted amide **13**, δ_C : 200 ppm).^{11,16} The lactams based on bicyclo[3.3.1]nonane ring systems (e.g. **8a**, **8b** and **11**; amide carbonyl δ_C : 181–185 ppm) were markedly more strained than those (e.g. **8c** and **8d**; amide carbonyl δ_C : 174–177 ppm) based on bicyclo[4.3.1]decane ring systems.

Strain was also assessed by analysis of the X-ray crystal structures of **8a** and **8c** and a derivative of **11** (lactam **16**, see below), and determination of standard amide distortion parameters (Supporting Information). For example the sum of the bond angles around the amide nitrogen (Θ) deviated further from 360° in the bicyclo[3.3.1]nonanes (**8a**: Θ = 337°; **16**: Θ = 341°) than in the bicyclo[4.3.1]decane ring system **8c** (Θ = 349°).

2.2. Synthesis of hetaryl-annulated scaffolds

The potential to access alternative scaffolds by annulation of heteroaromatic rings by exploiting the ketone in bicyclic lactam **11** was then explored (Scheme 3). Thus, reaction of the ketone **11** and propargylamine, catalyzed by 2.5 mol% NaAuCl₄,¹⁷ gave the related pyrido-fused lactam **14** in 43% yield. In a similar vein, reaction of **11** with 2-iodoaniline, catalyzed by 20 mol% Pd(OAc)₂, gave the indolo-fused lactam **15** in 53% yield.

Substrates	Intermediate synthesis		Lactam synthesis		
	Product	Yield,%	Product	Yield ^b %	δ_{C}^{c}
5 , 6a (n = 1; R^1 = Bn; R^2 = Et)	7a (n = 1; R^1 = Bn; R^2 = Et)	68	8a $(n = 1; R^1 = Bn)$	46	182.9
5 , 6b (n = 1; R^1 = Me; R^2 = Me)	7b (n = 1; R^1 = Me; R^2 = Me)	57	8b $(n = 1; R^1 = Me)$	45	182.0
5 , 6c (n = 2; $R^1 = Bn$; $R^2 = Et$)	7c (n = 2; R^1 = Bn; R^2 = Et)	72	8c $(n = 2; R^1 = Bn)$	59	176.
5 , 6d (n = 2; $R^1 = H$; $R^2 = Et$)	7d $(n = 2; R^1 = Cbz; R^2 = Et)$	74 ^a	8d $(n = 2; R^1 = Cbz)$	52	176.
9, ethyl acrylate	10	60	11	57	182.

^a Isolated after Cbz protection: CbzCl (1.1 eq.), NaHCO₃ (6.0 eq.) in CH₂Cl₂.

^b Yield over 3 steps.

^c Lactam carbonyl.

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