

Partially Saturated Bicyclic Heteroaromatics as an sp^3 -enriched Fragment Collection

David G. Twigg, Noriyasu Kondo, Sophie L. Mitchell, Warren R. J. D. Galloway, Hannah F. Sore, Andrew Madin and David R. Spring*

Dedicated to Professor Stuart L. Schreiber on the occasion of his 60th birthday

Abstract: Fragment-based lead generation has proven to be an effective means of identifying high-quality lead compounds for drug discovery programmes. However, the fragment screening sets often used are principally comprised of sp^2 -rich aromatic compounds, limiting the structural (and hence biological) diversity of the library. Herein, we describe strategies for the synthesis of a series of partially saturated bicyclic heteroaromatic scaffolds with enhanced sp^3 character. Subsequent derivatisation led to a fragment collection featuring the regio- and stereo-controlled introduction of substituents on the saturated ring system, often featuring the formation of new stereocentres.

Fragment-based drug discovery (FBDD) is a well-established method for generating high-quality hits and leads.^[1] The approval of B-Raf kinase inhibitor vemurafenib (Zelboraf) in 2011^[2] and Bcl-2 inhibitor venetoclax (Venclexta) in 2016,^[3] coupled with the ongoing evaluation of over 20 candidates in clinical trials,^[4] validates this approach as a complementary strategy to other hit discovery techniques such as high-throughput screening.^[5] While the growing prevalence of fragment-based approaches is encouraging, evaluation of many existing fragment libraries shows a predominance of (hetero)aromatic, ‘flat’ compounds, with a deficiency of chiral, sp^3 -rich examples.^[6,7]

Studies by Ritchie et al.^[8] and Lovering et al.^[9] demonstrate improvements in project progression by, for example, increasing the fraction of sp^3 centres within molecules or restricting the number of aromatic rings. Furthermore, computational analysis demonstrates that greater 3D conformational character is observed in compounds that have been clinically evaluated in humans, compared to those found in commercial libraries.^[10] This indicates the importance of sp^3 -richness in both the design of screening collections and the subsequent development of hits to leads.

Examples of previous studies aiming to synthesise collections of sp^3 -enriched fragments have been limited. Diversity-oriented synthesis^[7,11] and natural product derivatives^[12]

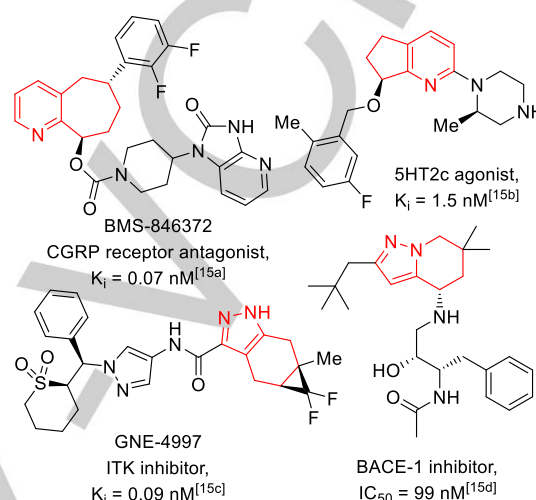
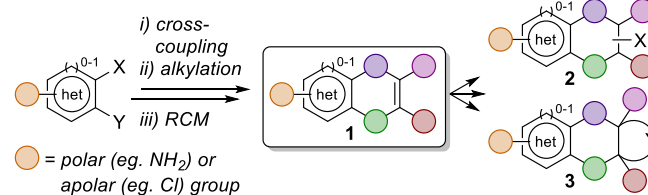


Figure 1: Selected examples of bioactive compounds containing functionalised partially saturated bicyclic heteroaromatics (highlighted in red).

have been used to generate 3D fragment collections, but there remains an unmet need to access new scaffold types. Recently there have been calls^[13] for new approaches and methodologies to design fragments with multiple synthetically accessible growth vectors in three dimensions, to allow rapid and efficient elaboration of hits to leads after initial screening, with some early success.^[14]

With these points in mind, the study described herein sought to develop efficient synthetic routes to a series of partially saturated bicyclic heteroaromatic (PSBH) fragments with enhanced sp^3 content relative to existing fragment libraries. Compounds featuring PSBHs have been shown to display bioactivity against a range of targets (Figure 1),^[15] and so a series of related fragments might be expected to serve effectively as a screening collection for FBDD applications.

The targets of this study featured a variable aromatic heterocycle fused to a partially saturated carbocycle. The heterocycle could bear either a polar (eg. amino) group, which should greatly enhance aqueous solubility, necessary for fragment screening at higher concentrations,^[7,14] or alternatively a hydrophobic (eg. chloro) group, able to forge key interactions



Scheme 1: General synthetic strategy toward PSBH scaffolds **1** and subsequent incorporation of new functionalities (**2**) or rings (**3**).

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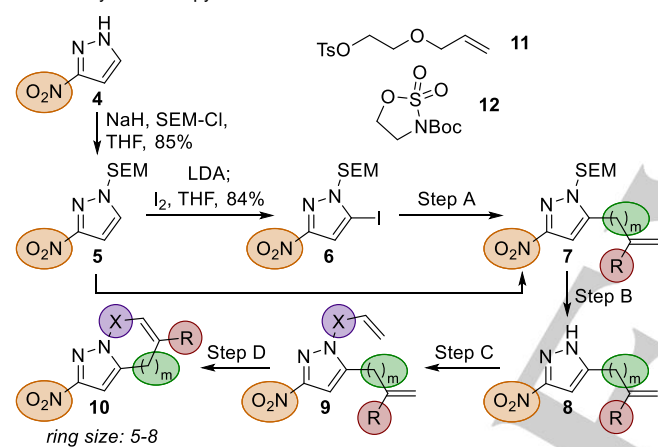
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with protein targets.^[16] The synthetic route (Scheme 1) employed a modular and divergent approach, using simple cross-coupling and alkylation reactions to install a pair of terminal olefins which could be reliably cyclised via ring-closing metathesis (RCM).^[17] This allowed for excellent control of the carbocycle ring size and the position and orientation of the resultant endocyclic olefin growth vector, which could undergo subsequent functionalisation to produce a range of fragments suitable for screening and/or further elaboration.

We selected pyrazole and pyridine as representative aromatic heterocycles. Whilst previous studies have synthesised related structures,^[18] they have incorporated less control over the position of the olefin and do not feature the amino group found in many of our compounds. Furthermore there are only very few examples where the olefin is used as a branch-point and further functionalised beyond simple reduction.^[15a,19]

Starting from readily available 3-nitro-1H-pyrazole **4**, SEM protection, selective iodination and subsequent Suzuki coupling with potassium vinyltrifluoroborate gave vinyl derivative **7a** (Table

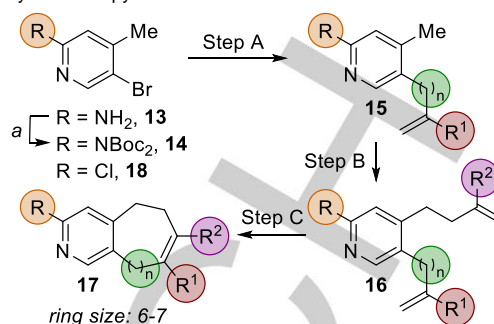
Table 1. Synthesis of pyrazole-based PSBH scaffolds.



m	R	X	Step			
			A (7)	B (8) ^[c]	C (9) ^[d]	D (10) ^[g]
a	H		85% ^[a]	80%	55%	45% ^[h]
b	H		-	-	86%	84%
c	H		-	-	55%	89%
d	H		-	-	59% ^[e]	65%
e	H		-	-	29% ^[f]	83%
f	Me		76% ^[a]	69%	44%	90%
g	H		61% ^[b]	82%	70%	93%
h	H		-	-	36%	59% ^[h]

Reaction conditions: [a] R'-BF₃K (1.5 equiv), Pd(dppf)Cl₂.CH₂Cl₂ (5 mol%), K₂CO₃ (3.0 equiv), THF/H₂O, 70 °C. [b] **5** (1.0 equiv), LDA (1.2 equiv), CuBr (20 mol%), allyl bromide (1.2 equiv), THF, -78 °C to RT. [c] TFA, CH₂Cl₂, RT. [d] NaH (1.5 equiv), alkyl bromide (1.5 equiv), THF or DMF, 70 °C. [e] NaH (1.5 equiv), **11** (1.5 equiv), THF, 70 °C. [f] **12** (2.0 equiv), K₂CO₃ (3.0 equiv), DMF, RT, 43%; then NaH (1.5 equiv), allyl iodide (1.5 equiv), DMF, RT, 68%. [g] Grubbs II (10 mol%), CH₂Cl₂, 40 °C. [h] Hoveyda-Grubbs II (10 mol%), toluene, 110 °C or CH₂Cl₂, RT.

Table 2. Synthesis of pyridine-based PSBH scaffolds.

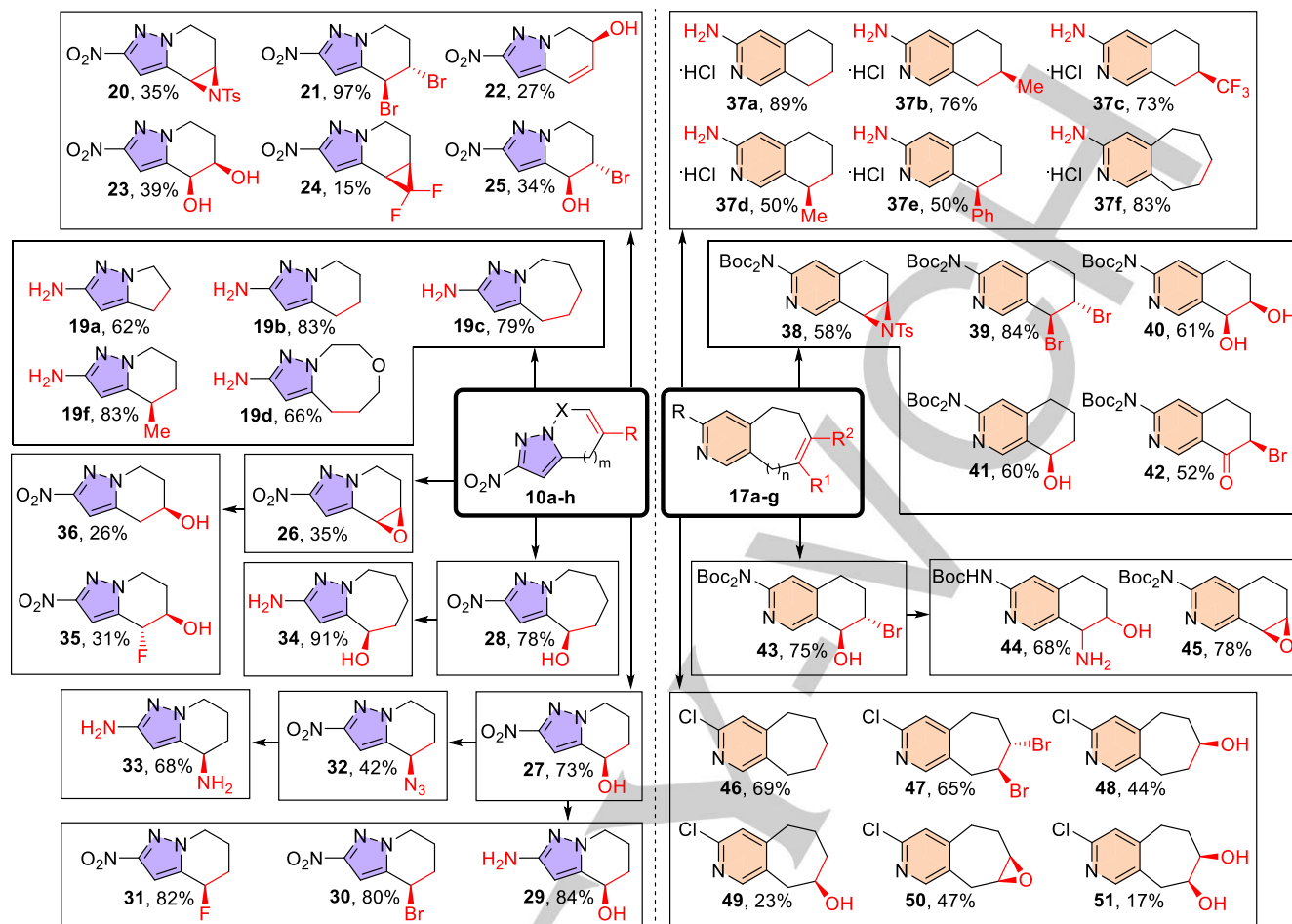


R	n	R ¹	R ²	Step		
				A (15) ^[b]	B (16) ^[e]	C (17) ^[f]
a	Boc ₂ N	0	H	85%	60%	95%
b	Boc ₂ N	0	Me	-	69%	94%
c	Boc ₂ N	0	CF ₃	-	77%	54%
d	Boc ₂ N	0	Me	87%	60%	94%
e	Boc ₂ N	0	Ph	92%	68%	83%
f	Boc ₂ N	1	H	78% ^[c]	41%	65%
g	Cl	1	H	73% ^[d]	79%	91%

Reaction conditions: [a] Boc₂O (2.5 equiv), DMAP (0.1 equiv), THF, 70 °C. [b] R'-BF₃K or R'-B(MIDA) (1.5 equiv), Pd(dppf)Cl₂.CH₂Cl₂ (10 mol%), K₂CO₃ (3.0 equiv), THF/H₂O, 70 °C. [c] Allyltributyltin (1.1 equiv), Pd(PPh₃)₄ (10 mol%), KF (2 equiv), toluene, 110 °C. [d] *i*-PrMgCl.LiCl (1.5 equiv), allyl bromide (1.2 equiv), THF, -15 °C to RT. [e] LDA (1.2 equiv), alkyl bromide (1.5 equiv), THF, -78 °C to RT. [f] Grubbs II (5 mol%), CH₂Cl₂, 40 °C.

1). Deprotection followed by *N*-alkylation with an alkyl bromide of varying C-chain length provided metathesis precursors **9a-c**, which upon treatment with either Grubbs' or Hoveyda-Grubbs' 2nd generation catalysts yielded the desired scaffolds **10a-c**. Inclusion of further heteroatoms in the formation of medium-sized partially saturated rings was achieved by treatment of vinyl intermediate **8a** with either tosylate **11** (leading, after RCM, to *O*-containing fragment **10d**), or 3-Boc-1,2,3-oxathiazolidine 2,2-dioxide **12** which gave access to the *N*-containing scaffold **10e** after allylation and metathesis. Use of a different Suzuki coupling partner gave methyl-substituted product **7f** which could be elaborated to PSBH fragment **10f**. Alternatively, direct allylation at the C-5 position of SEM-protected intermediate **5** could be achieved upon treatment with lithium diisopropylamide (LDA) and allyl bromide. This led, in an analogous way, to scaffolds **10g-h** with non-conjugated olefins.

A similar approach was used to generate PSBH scaffolds from pyridine **13** (Table 2). Attempts to mask the 2-amino group as a nitro group proved ineffective as, despite successful cross-coupling reactions, the 2-nitropyridines were unstable to strong base and did not undergo the desired alkylations at the 4-methyl position. Mono-Boc protection was also unsuitable due to poor yields in the cross-coupling step, possibly due to catalyst chelation. The 2-amino group could be rendered synthetically tractable, however, either with bis-Boc protection or via substitution for a 2-chloro group, which itself can serve as a synthetic handle.^[20]



Scheme 2: Functionalisation of PSBH scaffolds. For reaction conditions, see SI.

Bis-Boc substrate **14** (prepared in one step from **13**) was functionalised at the 5-position using either Suzuki coupling (for vinyl substituents) or Stille coupling (for allyl substituents) to produce intermediates **15a,d-f**. Treatment with LDA and trapping of the resultant anion with a variable alkyl bromide electrophile gave a range of metathesis substrates **16a-f**, which under standard ring-closing metathesis conditions yielded PSBH scaffolds **17a-f**. The 2-chloro substrate **18** could be allylated in the 5-position using an excess of *i*-PrMgCl.LiCl and trapping the resultant organometallic intermediate with allyl bromide. Allylation at the 4-methyl position and RCM thus yielded scaffold **17g** in superior yields.

Following PSBH synthesis, a series of simple one-, two- or three-step functionalisations were performed on selected pyrazole and pyridine scaffolds to demonstrate the synthetic utility of the olefin π -bond as a growth vector and to generate a variety of new stereocentres (Scheme 2).

Catalytic hydrogenation of nitropyrazoles **10a-d,f** served to reduce both the olefin π -bond and the nitro group in moderate to good yields, revealing the latent amino functionality and, in the case of **19f**, creating a new stereocentre. Other one-step reactions include aziridination, dibromination, allylic oxidation, dihydroxylation, difluorocyclopropanation, hydroxybromination, epoxidation and hydroboration (**20-28**), introducing functionalities at the 4-, 5- and 6-positions of the fused pyrazole systems.

Demonstrating that these initial products can serve as intermediates to other fragments, the products of hydroboration can react further to incorporate Br, F and N substituents (**30-33**), whilst epoxide **26** can be opened by nucleophiles such as fluoride and hydride to form fluorohydrin **35** and alcohol **36**. Whilst the yields of some reactions were modest, sufficient material was obtained for full characterisation and future screening campaigns.

The pyridine-based scaffolds **17a-g** can undergo a similar range of transformations. Catalytic hydrogenation and subsequent acid-mediated deprotection of bis-Boc compounds **17a-f** generated novel fragments, many of which include new stereocentres (**37a-f**). Aziridination, dibromination, dihydroxylation, hydroboration, α -bromoketone formation and hydroxybromination were also carried out (**38-43**). Further reactions included amino-alcohol (**44**) and epoxide (**45**) formation, both from bromohydrin **43**. 3-Chloro fragments **46-51** could also be readily accessed using similar conditions.

Calculation of a range of physicochemical properties was carried out on all PSBH products. Almost all fragments were shown to conform to the so-called "Rule of Three", a set of criteria commonly associated with greater hit rates in fragment screening collections.^[21] Of particular note are the low mean values of molecular weight (190), SlogP (1.45) and 'fraction aromatic' (0.43) and the high mean number of chiral centres (0.88), especially when compared to existing commercial libraries (Table 3).

Table 3. Mean physicochemical properties of fragment collections.

Property ^[a]	Ideal Range ^[b]	This work	Chembridge	Maybridge
SlogP	0-2	1.45	1.31	2.55
MW	140-230	190	222	265
PSA	≤60	58.0	53.9	57.5
HBA	≤3	1.35	1.81	2.12
HBD	≤3	0.55	1.04	0.81
HAC	10-16	12.8	15.5	18.0
RBC	≤3	0.6	3.2	2.8
Chiral centres	0-1	0.88	0.27	0.18
Fraction Aromatic	-	0.43	0.42	0.52

[a] MW = molecular weight, PSA = polar surface area, HBA = number of hydrogen-bond acceptors, HBD = number of hydrogen-bond donors, HAC = heavy atom count, RBC = rotatable bond count. [b] Based on the guidelines used by Astex Pharmaceuticals.^[14,21] Green = within ideal range, orange = at extreme of ideal range, red = outside ideal range. See SI for further details.

In conclusion, we have developed simple, scalable routes to a series of partially saturated pyrazole- and pyridine-based scaffolds which can readily undergo a range of synthetic transformations to generate a collection of sp³-rich fragments, suitable either as screening members in a library or as intermediates to 'higher content fragments'. The compounds adhere to recognised guidelines for fragment physicochemical properties whilst displaying enhanced sp³ character, greater chirality and providing a range of three-dimensional growth vectors for synthetic development. It is envisioned that the strategy could be applied to a vast range of analogous scaffolds with varied heterocycles and substituents and that several of the functionalisation reactions detailed in Scheme 2 could be rendered asymmetric based on related precedent.^[19,22]

Acknowledgements

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Keywords: Synthetic methods • Drug discovery • Fused-ring systems • Nitrogen Heterocycles • Drug design

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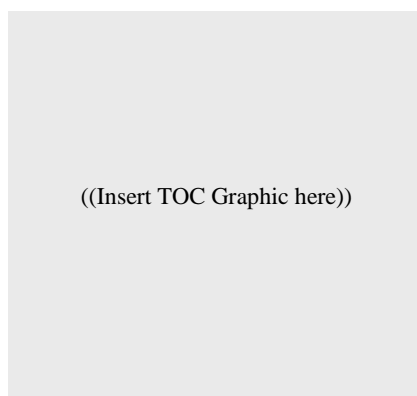
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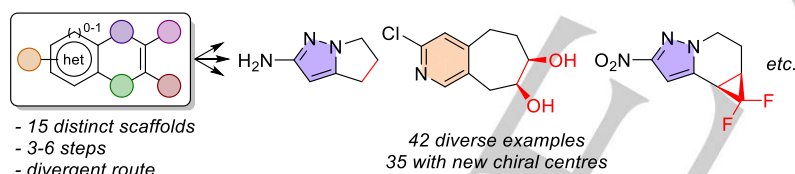
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**Partially Saturated Bicyclic
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 Fragment Collection**

2D or not 2D: A collection of partially saturated bicyclic pyridine- and pyrazole-based fragments, derived from a diverse set of readily accessible branch-point scaffolds, is presented. Their enhanced sp^3 content, compared to typical fragment libraries, allows for excellent control of 3D growth vectors in drug development applications.