Accepted Manuscript

Structure-based design, synthesis and biological evaluation of a novel series of isoquinolone and pyrazolo[4,3-c]pyridine inhibitors of fascin 1 as potential antimetastatic agents

Stuart Francis, Daniel Croft, Alexander W. Schüttelkopf, Charles Parry, Angelo Pugliese, Ken Cameron, Sophie Claydon, Martin Drysdale, Claire Gardner, Andrea Gohlke, Gillian Goodwin, Christopher H. Gray, Jennifer, Konczal, Laura McDonald, Mokdad Mezna, Andrew Pannifer, Nikki Paul, Laura Machesky, Heather McKinnon, Justin Bower



PII:	S0960-894X(19)30056-3
DOI:	https://doi.org/10.1016/j.bmcl.2019.01.035
Reference:	BMCL 26272
To appear in:	Bioorganic & Medicinal Chemistry Letters
Received Date:	26 November 2018
Revised Date:	24 January 2019
Accepted Date:	29 January 2019

Please cite this article as: Francis, S., Croft, D., Schüttelkopf, A.W., Parry, C., Pugliese, A., Cameron, K., Claydon, S., Drysdale, M., Gardner, C., Gohlke, A., Goodwin, G., Gray, C.H., Jennifer, Konczal, McDonald, L., Mezna, M., Pannifer, A., Paul, N., Machesky, L., McKinnon, H., Bower, J., Structure-based design, synthesis and biological evaluation of a novel series of isoquinolone and pyrazolo[4,3-*c*]pyridine inhibitors of fascin 1 as potential antimetastatic agents, *Bioorganic & Medicinal Chemistry Letters* (2019), doi: https://doi.org/10.1016/j.bmcl. 2019.01.035

This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

Structure-based design, synthesis and biological evaluation of a novel series of isoquinolone and pyrazolo[4,3-*c*]pyridine inhibitors of fascin 1 as potential anti-metastatic agents

Stuart Francis^{a, *}, Daniel Croft^a, Alexander W. Schüttelkopf^a, Charles Parry^a, Angelo Pugliese^a, Ken Cameron^a, Sophie Claydon^a, Martin Drysdale^{a, 1}, Claire Gardner^a, Andrea Gohlke^a, Gillian Goodwin^a, Christopher H. Gray^a, Jennifer^a Konczal^a, Laura McDonald^a, Mokdad Mezna^a, Andrew Pannifer^{a, 2}, Nikki Paul^b, Laura Machesky^{b, c}, Heather McKinnon^a and Justin Bower^a

^a Drug Discovery Unit, CRUK Beatson Institute, Glasgow G61 1BD (UK)
^b Cell Migration Laboratory, CRUK Beatson Institute, Glasgow G61 1BD (UK)
^c Institute of Cancer Sciences, University of Glasgow, G61 1QH, Glasgow, UK
*Corresponding author at: Drug Discovery Unit, CRUK Beatson Institute,
Glasgow G61 1BD (UK). *E-mail address*: <u>s.francis@beatson.gla.ac.uk</u>
¹ Current address: Broad Institute, Cambridge, MA 02142, US
² Current address: Medicines Discovery Catapult, Mereside, Alderly Park,
Alderley Edge SK10 4TG, UK

Keywords: virtual screening, fragments, drug discovery, medicinal chemistry, cancer

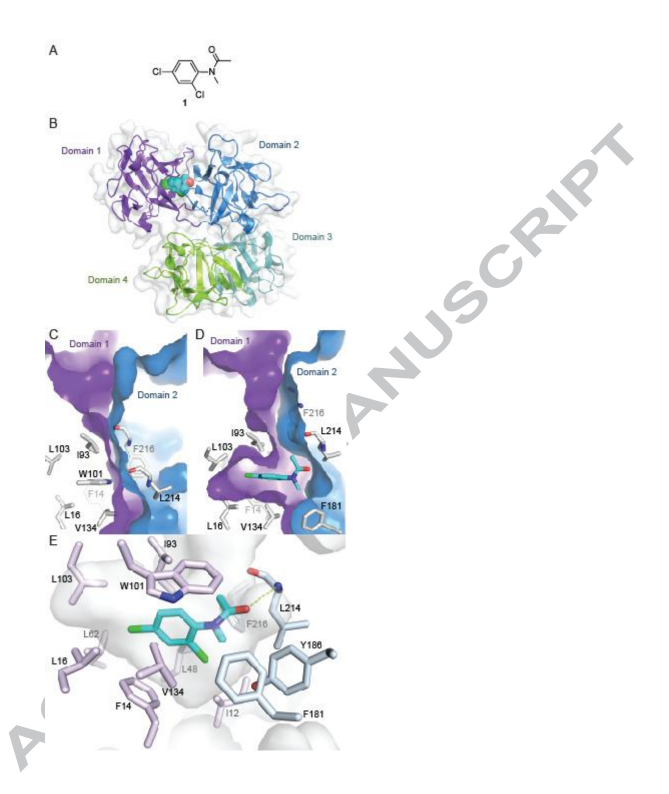
Abstract

Fascin is an actin binding and bundling protein that is not expressed in normal epithelial tissues but overexpressed in a variety of invasive epithelial tumors. It has a critical role in cancer cell metastasis by promoting cell migration and invasion. Here we report the crystal structures of fascin in complex with a series of novel and potent inhibitors. Subsequent structure-based elaboration of this and related compounds enabled the development of a series with nanomolar affinities for fascin, good physicochemical properties and the ability to inhibit fascin-mediated bundling of filamentous actin. These compounds provide promising starting points for fascin-targeted anti-metastatic therapies.

Fascin 1 (hereafter termed fascin) is an evolutionarily conserved actin binding protein that cross-links filamentous actin (F-actin) into tightly packed parallel bundles driving the formation of various cell surface protrusions, such as filopodia and invadopodia, that promote cell migration and invasion.¹ It's expression is highly restricted in adult human tissues.² Fascin is either absent or shows low expression in normal epithelial tissues but is overexpressed in a number of cancers, with expression levels correlating with overall cancer aggressiveness and predicting poor clinical outcome.^{3, 4} Conversely, genetic knockdown of fascin decreases tumor cell invasion both *in vitro* and *in vivo*,^{5, 6} suggesting that fascin may be a viable target for anticancer/antimetastatic drugs. Huang et al. have disclosed compounds which bind to fascin and inhibit actin bundling.⁷⁻⁹ However, in our hands a number of these compounds bound with modest (10–100 μ M) affinity and suffered from poor solubility.

In order to identify novel fascin compounds, with favourable properties, we undertook a surface plasmon resonance (SPR) screen using our collection of 1050 fragments and identified 53 hits (hit rate 5%). Follow-up protein crystallography yielded a range of fragments binding to multiple sites on the protein.

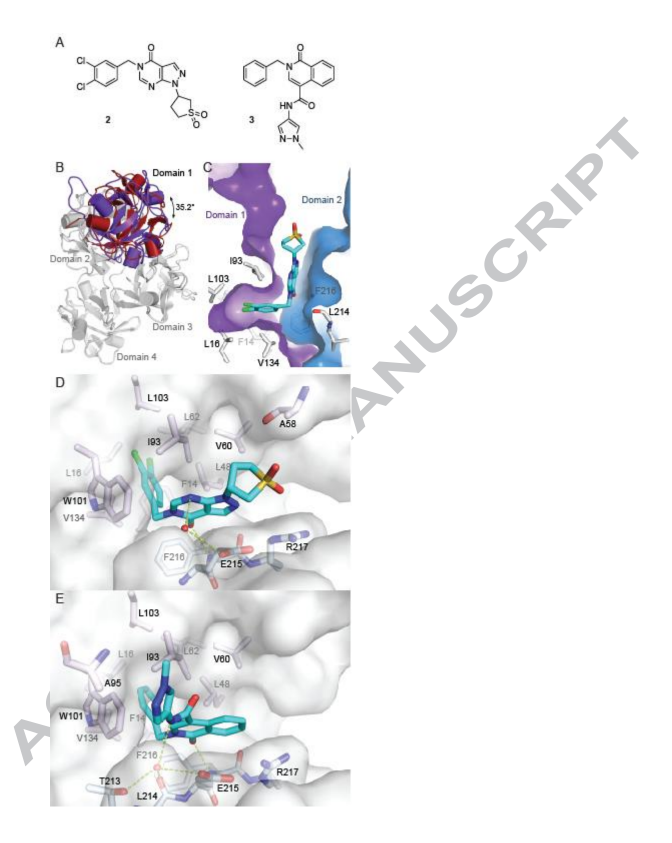
Figure 1. Fascin structure and fragment **1** binding. **A)** Chemical structure of compound **1. B)** Overall structure of fascin represented as a cartoon with the four β-trefoil domains colored differently and labelled. The bound compound **1** is shown as cyan spheres (PDB id 6I0Z). **C)** The interface between domains 1 and 2 in apo-fascin (PDB id 3P53)¹⁰. Domains are delineated by separate surfaces colored as in B) and labelled. Residues at the interface are shown as grey sticks and labelled with single-letter amino acid codes and residue numbers. **D)** Compound **1** bound to fascin, representation and view as in C) with the compound shown as cyan sticks. The structures were superimposed on domain 1 only. **E)** Detailed view of the binding site shown as a semi-transparent surface with surrounding protein residues drawn as sticks colored and labelled. Hydrogen bonds are shown as dashed pale green lines, other conventions as for D).



The *N*-phenylacetamide **1** (Figure 1A) binds efficiently (SPR $K_d = 92 \mu M$; ligand efficiency 0.43)¹¹ in a buried pocket between the first and second β trefoil domain (Figure 1B) and was therefore selected for further development.

Binding of compound **1** induces a hydrophobic pocket in domain 1, mainly by displacing Trp101. In addition, domain 1 pivots by $\approx 15^{\circ}$, reorienting residues on either side of the domain interface and creating a narrow channel to the protein surface (Figures 1C and 1D). The dichlorophenyl moiety of **1** fills the hydrophobic pocket, while the acetamide engages in hydrophilic interactions with domain 2, with its carbonyl accepting a hydrogen bond from the backbone of Leu214 (Figure 1E). While the completely enclosed binding site makes compound **1** a highly efficient ligand, it hampered attempts to develop this series.

Figure 2. Fascin undergoes a significant conformational change upon binding of compounds **2** and **3**. **A)** Chemical structures of **2** and **3**. **B)** Apo-fascin (PDB id 3P53, red) and the fascin·**2** complex (PDB id 6l10, purple) superimposed on the relatively rigid domains 2–4 (grey; RMSD = 1.1 Å for 348 C α atoms), viewed along the domain 1 rotation axis. **C)** Compound **2** bound to fascin oriented as Figures 1C/D (proteins superimposed on domain 1). **D)** Detail view of the fascin·**2** complex. Conventions as in Figure 1E, the red sphere represents a bound water molecule. **E)** Detailed view of the fascin·**3** complex (PDB id 6l11).



The importance of this fragment binding site was highlighted when we determined the crystal structure of compound **2** (Figure 2A) previously identified by Huang et al. to inhibit fascin-mediated actin bundling.¹² To

accommodate the larger compound 2, domain 1 undergoes a more dramatic conformational change, rotating 35° around an axis almost parallel to a line connecting the first two β -trefoil domains (Figure 2B). For the compound binding site (Figure 2C) this almost pure rotation results in a 6 Å 'drop' of domain 2 relative to domain 1, so that the Phe216 side chain, which in the fascin compound 1 complex lines the channel, now extends the hydrophobic pocket (cf. Figures 1D and 2C) where the compound's dichlorobenzyl moiety binds. More importantly, the conformational change opens up the channel so that it can accommodate the rest of the compound. The pyrazolopyrimidinone core rests on a relatively flat surface created by the salt-bridged side chains of Glu215 and Arg217. Additionally, compound 2 accepts a hydrogen bond from Phe216 and makes a relatively weak water-bridged interaction with the side chain of Glu215 (Figure 2D). The induced conformational changes provide 2 with access to the protein surface creating opportunities for structure-driven compound elaboration. Notably, the recently disclosed structure of fascin in complex with an unrelated small molecule bound in the same site⁹ shows fascin adopting a similar conformation (RMSD < 1.3 Å for 483 C α atoms).

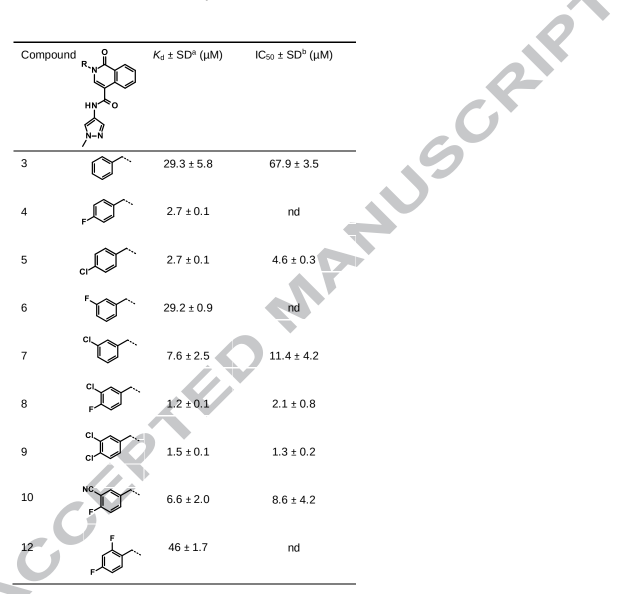
To identify compounds which could potentially fill these additional areas a hybrid virtual screen was carried out. The first method was structure-based: 809 commercial analogues of compound **2** and \approx 2.7 million lead-like molecules from our in-house virtual library were docked¹³ in the enclosed site. To prioritize the docked compounds, a multi-parameter optimization was performed using Pareto efficiency¹⁴ as the ranking technique with calculated free energy of binding (from docking), QED¹⁵ and interaction pattern similarity

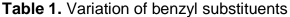
(based on compound 2 interaction fingerprints)¹⁶ as the parameters to optimize. The second method was ligand-based using shape-similarity¹⁷ to find analogues of compound **2** in our in-house library. Combining the results, 110 commercial compounds were tested by SPR at a screening concentration of 100 µM. The resulting binders were validated by co-crystallization and SPR dose-response experiments. Eighteen compounds were confirmed to bind in the same site as **2**, and three yielded K_d values <100 μ M. Of these three, we decided to progress the isoquinolinone **3** (SPR $K_d = 29.3 \pm 5.8 \mu$ M; Figures 2A and 2E), which maintains affinity similar to 2 despite substituent truncations from both the benzyl and the core ring system. The pyrazolamide moiety forms an edge-to-face aromatic interaction with the Trp101 side chain. It also moves the bridging water seen in the fascin compound 2 complex structure deeper into the binding site. This water now hydrogen-bonds to the Thr213 side chain, the Leu214 backbone carbonyl and the ligand amide in addition to the Glu215 side chain, which 'follows' it by adopting a different rotamer (Figures 2D and 2E). Analysis of this water with MOE 3D-RISM¹⁸ suggests it is relatively tightly bound ($\Delta G = -3.9$ kJ/mol) and that displacing it would likely be energetically unfavorable.

In order to measure the effect of these compounds on fascin activity we developed a robust and highly reproducible *in vitro* F-actin bundling assay, based on a previously published method.¹⁹ Unlike compound **1**, both **2** and **3** inhibit fascin in this bundling assay. Compound **3** inhibits actin bundling with an IC₅₀ of 67.9 \pm 3.5µM, giving an approximately two-fold fall-off from SPR binding affinity. A similar fall-off is seen for other compounds in this series.

Considering that both 1 and 2 place a substituted phenyl into the hydrophobic pocket, it seemed plausible that the plain benzyl of **3** is not optimal for filling the available space. Additionally, given that this induced pocket is likely to show some plasticity, it may be possible to open it up further by extending the compound. To investigate this, we synthesized a series of benzyl derivatives of **3** (Table 1). Introduction of a *para*-halo substituent (compounds **4** and **5**) increased binding affinity \approx 14-fold compared to 3. The crystal structure of 5 bound to fascin shows a binding mode without substantial changes in either ligand or protein conformation compared to the fascin-3 complex (Figure S1A). A single *meta*-fluoro substituent (6) is equipotent while *m*-chloro substitution (7) improved affinity, though less so than 4 or 5. This may be partly explained by the slight differences in binding mode for meta substituted compounds compared to 3, in which the benzyl moiety is rotated to line up the m-substituent with a pocket indentation (Figure S1B). In light of this conformational divergence, it is not surprising that m,p-dihalo compounds exhibit sub-additive affinity gains, though both 8 and 9 showed improved binding over mono- or unsubstituted analogues with SPR K_d values of 1.2 and 1.5 µM respectively. The structure of the fascin 9 complex shows the compound adopts a conformation with the phenyl rotated to an intermediate position compared to the *meta* and *para* analogues (Figure S1C). Larger *meta* or *para* substituents (*e.g.* **10** and **11**; Figure S1D) are tolerated by expanding the hydrophobic pocket. However, the associated conformational penalty consistently makes these compounds less potent. Introduction of an ortho substituent decreases potency both in combination with a para substituent

(12) or as a single substituent (data not shown). Taken together the SAR suggests that m,p-dichloro substitution (9) is close to optimal in this series.





^a K_d : SPR binding affinity, values are mean ± standard deviation from n=2-10 separate experiments; ^b IC₅₀: compound concentration that reduces actin bundling by 50%, values are mean ± standard deviation

We next investigated altering the bicyclic scaffold of **3** (Table 2), taking care to retain the pyridinone moiety, which we found to be essential for binding. Complete deletion of the outer ring (**13**) led to a significant loss of affinity, which can be partly recovered with the addition of a 3-amino substituent (**14**),

which donates an additional hydrogen bond to the Phe216 backbone carbonyl. The naphthyridinones **16** and **17** maintain binding affinities similar to the parent isoquinolinone, while placement of the naphthyridinone nitrogen next to the amide (**15**) results in the formation of an intramolecular hydrogen bond stabilising a conformation that is not optimal for fascin binding (Figure S2A), and thus a decrease in binding affinity. For comparison, a matched pyrazolopyridinone **19** shows 7-fold weaker binding.

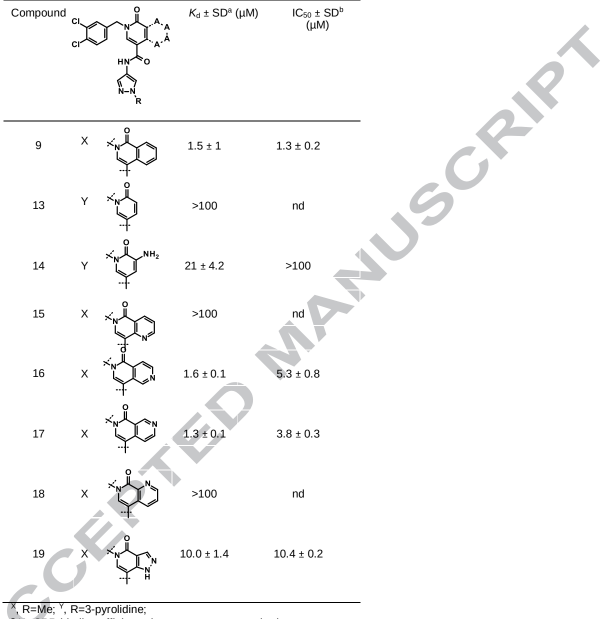
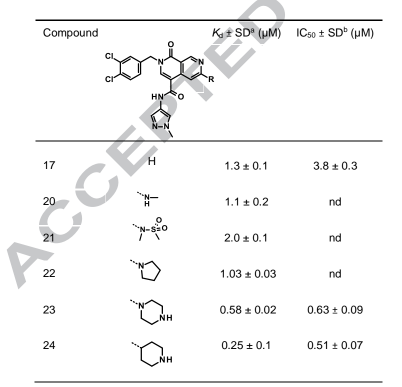
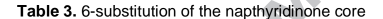


Table 2. Variation of the bicyclic scaffold

^X, R=Me; ^Y, R=3-pyrolidine; ^a K_d : SPR binding affinity, values are mean ± standard deviation from n=2-10 separate experiments; ^b IC₅₀: compound concentration that reduces actin bundling by 50%, values are mean ± standard deviation from n=2-4 separate experiments; nd: not determined.

Considering their improved physicochemical properties and chemical tractability, we decided to progress a naphthyridinone, initially by adding substituents to replicate the space filling of the cyclic sulphone of **2**. A binding mode comparison (Figure S2B) suggested that substitution from C5 would provide the best-matched vector, though 6-substituents should be able to access the same space. Due to concerns about the potential clash between a 5-substituent and the pyrazoloamide preventing the compound from adopting the preferred fascin-binding conformation, we opted to elaborate **17** through substitution from the 6-chloro derivative (Table 3). The introduction of simple





^a K_d : SPR binding affinity, values are mean ± standard deviation from n=2-10 separate experiments; ^b IC₅₀: compound concentration that reduces actin bundling by 50%, values are mean ± standard deviation from n=2-4 separate experiments; nd: not determined.

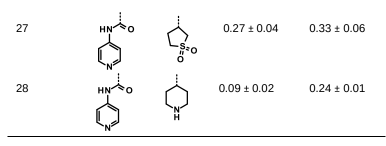
alkylamine substituents (20, 22) had no significant effect on fascin binding. The sulphonamide (21) aiming to more closely replicate the sulphone of compound 2 also failed to improve affinity. The piperazine 23 gave slightly improved fascin binding, despite ostensibly placing a partial positive charge close to the guanidine group of Arg217 (cf. Figure 2E). The piperidine analogue (24) shows further improved fascin binding and inhibition. The crystal structure of the fascin-compound 24 complex shows the compound binding broadly as expected (Figures S3A and S3B). The naphthyridinone makes an additional water-bridged interaction with the Phe216 backbone carbonyl. The piperidine orients to fill the space between Ala58 and Arg217 (Figure S3C), but does not make any specific interactions, as its amine points towards bulk solvent. Comparing the binding modes of 24 and 9 reveals that the core ring system of 24 is rotated slightly (Figure S3D), possibly to optimize the fit of the piperidine. This small rotation has an amplified effect on the position of the pyrazoloamide, bringing the pyrazole closer to Trp101 and Arg225 while at the same time improving the hydrogen bond from the amide to the bridging water. The displacement observed for 24 suggests that naphthyridinones with bulky 6-substituents may be sterically restricted from adopting an optimal core binding mode. In light of this we chose to revisit the pyrazolo series, where we had previously observed that the equally bulky cyclic sulphone substituent of 2 was compatible with the 'standard' core orientation (Figure S2B). The truncated compound 25 showed diminished binding (Table 4), revealing that addition of either the pyrazoloamide in the 7position (to obtain 19) or the cyclic sulphone at N1 (to obtain 2) to this compound improves fascin affinity more than 50-fold and 23-fold, respectively.

Combining these key features into one compound gave **26**, with an SPR *K*d of 0.6 μ M, suggesting near-additive gains in binding affinity compared to **24** and almost closing the potency gap to our best naphthyridinones.

We next investigated changes to the heteroaromatic amide. In line with the predicted importance of the bridging water discussed above, any changes to the amide (N-alkylation, carbonyl deletion) abolish fascin binding (not shown). On the other hand, the methylpyrazole interacts with the protein through stacking with the indole of Trp101. A number of compounds with alternative heterocycles were made to probe this interaction, focusing on five- and six-membered nitrogen-containing heterocycles. Of these the pyridin-4-yl **27** yielded the largest, albeit still modest, improvement over **26** in terms of both affinity and inhibition. QM calculations (Figure S4) suggest that at least for (hetero)aryl derivatives improved stacking with Trp101 may be a major factor in this increase in potency.

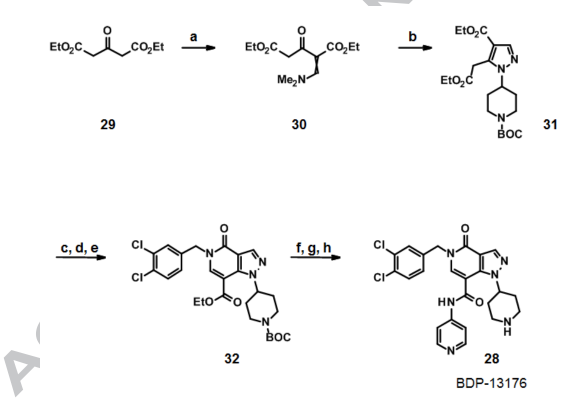
Compound	R ¹ -aroup	R ² -aroup	$K_{d} \pm SD^{a}$ (µM)	$IC_{50} \pm SD^{b}$ (µM)
25	Ν	Н	>500	nd
2	Ν		29.5 ± 8.3	67.6 ± 2.4
26			0.60 ± 0.05	0.63 ± 0.15

Table 4. Further substitutions of the pyrazolopyridinone scaffold



^a K_d : SPR binding affinity, values are mean ± standard deviation from n=2-10 separate experiments; ^b IC₅₀: compound concentration that reduces actin bundling by 50%, values are mean ± standard deviation from n=2-4 separate experiments; nd: not determined.

Given the success of **24** and to alleviate concerns about chemical stability of the cyclic sulphone substituent, we opted to replace it with a 1-piperidin-4-yl.



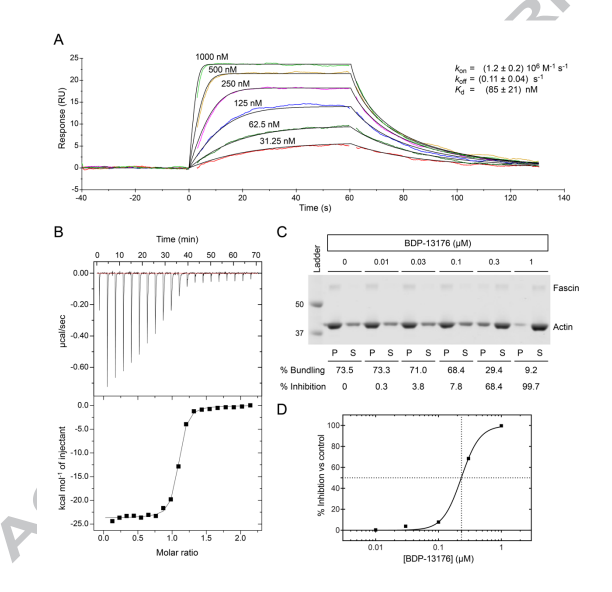
Scheme 1. Reagents and conditions: (a) *N, N*-dimethylformamide dimethyl acetal, ethanol r.t, (75%); (b) 4 *tert*-butyl 4-hydrazinopiperidine-1-carboxylate, ethanol, reflux., (27%); (c-e) Bredereck's, toluene reflux; 3,4-dichlorophenyl)methanamine, acetic acid, toluene; sodium ethoxide, ethanol

(3 steps 85%) (f) Sodium hydroxide, ethanol r.t. (84%), (g) <u>HATU</u>, *N,N*diisopropylethylamine, DMF, r.t., (83%), (h) TFA, DCM r.t. (52%).

Synthesis of this was achieved starting from the commercially available diethyl beta-ketoglutarate (29) and reacting with N,N-dimethylformamide dimethyl acetal (Scheme 1).²⁰ Addition of *tert*-butyl 4-hydrazinopiperidine-1carboxylate led to the pyrazole (31), which was further functionalised by with Bredereck's the refluxing reagent followed bv addition of dichlorobenzylamine. Partial cyclisation was observed at this stage, however, the crude product was fully cyclised with sodium ethoxide. This gave intermediate pyrazlopyridinone (32) in good yield over the three steps. Saponifaction of the ester, coupling of the 4-amino pyridyl and removal of the Boc protecting group furnished BDP-13176 (28). BDP-13176 further improved fascin affinity, giving us the best fascin binder (SPR K_d = 85 ± 0.02 nM, LE = 0.29, ITC K_d = 50 nM; Figures 3A and 3B) and actin bundling inhibitor (IC₅₀ = 240 ± 0.01 nM; Figures 3C and 3D).

Figure 3. Comparison of SPR binding affinity, ITC binding parameters and functional bundling assay potency for BDP-13176. **A)** Multi-cycle kinetic curve for BDP-13176 binding to immobilised (His₈)₂-fascin. Data were fitted using a 1:1 kinetic binding model. **B)** ITC titration thermogram and referenced isotherm (first point removed) of BDP-13176 (150 µM in syringe) binding to fascin (15 µM in cell). Fitting of the isotherm to a 1:1 binding model resulted in the following binding parameters: $N = 1.05 \pm 0.01$, $K_d = (45 \pm 7)$ nM, $\Delta H = (-2400 \pm 200)$ cal mol⁻¹, $\Delta S = -45.9$ cal mol⁻¹ K⁻¹. **C)** Inhibition of fascin bundling

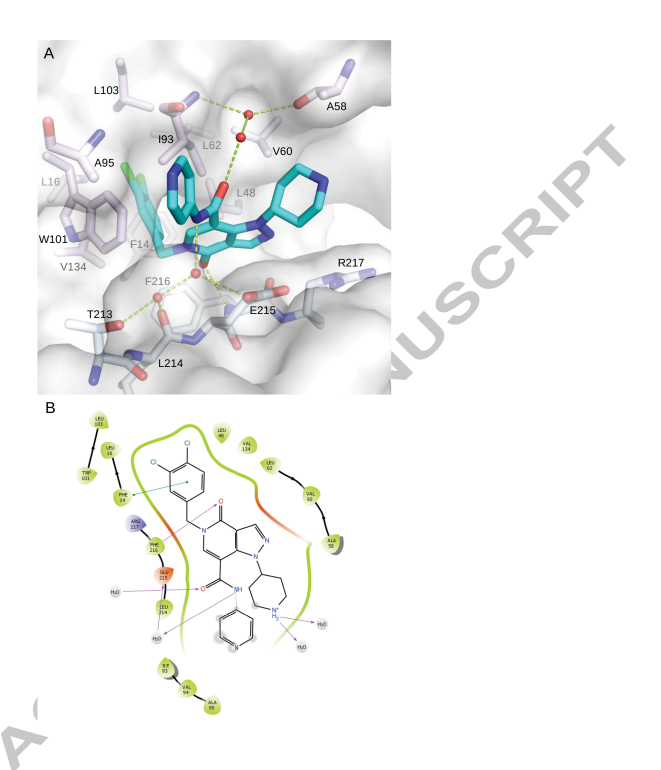
activity by BDP-13176. Inhibition of bundling activity can be seen as fascin and F-actin move from the pellet (P) to the supernatant (S) with increasing concentrations of BDP-13176. **D)** Concentration-response curve for bundling assay data shown in C).



The crystal structure of fascin in complex with BDP-13176 shows the compound adopting the expected overall binding mode (Figure 4), but with a slight core rotation in the opposite direction from that observed for **24** (Figure S5). The compound's *in vitro* physicochemical and DMPK properties (Table

S1) are within acceptable limits with low lipophilicity (LogD_{7.4}=1.8) and reasonable kinetic solubility (midpoint = 65μ M). The compound was also moderately stable in both human and mouse liver microsomes (Human Cl_{int}=6.4, Mouse Cl_{int}=11.8, μ L/min/mg protein). However, the compound has low Caco-2 permeability (A-B P_{app}=0.15x10⁻⁶ cm/s) and high efflux (efflux ratio=37.8).

Figure 4. Binding of BDP-13176 to fascin. **A)** Crystal structure of the fascin-BDP-13176 complex (PDB id 6I18). The bridging water seen in earlier complexes is replaced by two ordered waters that together make the same contacts as the single bound water. **B)** Interaction plot for BDP-13176 bound to fascin.



In conclusion, fragment screening coupled with crystallography and computational approaches allowed identification of a cryptic pocket within fascin. Using compound **2** ($K_d = 29.5 \pm 8.3 \mu M$)¹² as a starting point combined with virtual screening and structure-based design allowed us to develop potent and functionally active fascin binders. Crystal structures reveal the

compounds bind with a hydrophobic 'hook' in an induced pocket between the first two β -trefoil domains of fascin and from there extend towards the protein surface, effecting a substantial conformational change in domain 1. While the mechanism by which our compounds inhibit bundling remains unclear, it is worth noting that fascin's two major proposed actin binding regions involve domain 1 and cross a domain boundary. The ligand-induced conformational change would deform both these regions and thereby disrupt actin binding (Figure S6). We anticipate these compounds to be a useful foundation to further probe fascin's potential role in tumor invasion and metastasis.

Acknowledgements

This research was supported by funding from CRUK (grants A17196 and A17096) and the Pancreatic Cancer Research Fund. We would like to thank Diamond Light Source for beamtime (proposals mx6683, mx8659 and mx11651) and the staff of beamlines i02, i03 and i04 for their assistance. We also thank David Sumpton and Gillian McKay for mass spectrometric analyses, and Daniel James for informatics support.

Experimental Section

Experimental details can be found within the supplementary data file provided.

References

1. Li A, Dawson JC, Forero-Vargas M, et al. The actin-bundling protein fascin stabilizes actin in invadopodia and potentiates protrusive invasion. *Curr Biol.* 2010;20(4): 339-345. https://doi.org/10.1016/j.cub.2009.12.035.

2. Hashimoto Y, Kim DJ, Adams JC. The roles of fascins in health and disease. *J Pathol.* 2011;224(3): 289-300. https://doi.org/10.1002/path.2894.

3. Tan VY, Lewis SJ, Adams JC, Martin RM. Association of fascin-1 with mortality, disease progression and metastasis in carcinomas: a systematic review and meta-analysis. *BMC Med.* 2013;11: 52. https://doi.org/10.1186/1741-7015-11-52.

4. Ma Y, Machesky LM. Fascin1 in carcinomas: Its regulation and prognostic value. *Int J Cancer.* 2015;137(11): 2534-2544. https://doi.org/10.1002/ijc.29260.

5. Li A, Morton JP, Ma Y, et al. Fascin is regulated by slug, promotes progression of pancreatic cancer in mice, and is associated with patient outcomes. *Gastroenterology.* 2014;146(5): 1386-1396 e1381-1317. https://doi.org/10.1053/j.gastro.2014.01.046.

6. Minn AJ, Gupta GP, Siegel PM, et al. Genes that mediate breast cancer metastasis to lung. *Nature.* 2005;436(7050): 518-524. https://doi.org/10.1038/nature03799.

 Huang FK, Han S, Xing B, et al. Targeted inhibition of fascin function blocks tumour invasion and metastatic colonization. *Nat Commun.* 2015;6: 7465. https://doi.org/10.1038/ncomms8465.

8. Han S, Huang J, Liu B, et al. Improving fascin inhibitors to block tumor cell migration and metastasis. *Mol Oncol.* 2016;10(7): 966-980.

https://doi.org/10.1016/j.molonc.2016.03.006.

9. Huang J, Dey R, Wang Y, Jakoncic J, Kurinov I, Huang XY. Structural Insights into the Induced-fit Inhibition of Fascin by a Small-Molecule Inhibitor. *J Mol Biol.* 2018;430(9): 1324-1335.

https://doi.org/10.1016/j.jmb.2018.03.009.

10. Jansen S, Collins A, Yang C, Rebowski G, Svitkina T, Dominguez R. Mechanism of actin filament bundling by fascin. *J Biol Chem.* 2011;286(34): 30087-30096. https://doi.org/10.1074/jbc.M111.251439.

11. Hopkins AL, Groom CR, Alex A. Ligand efficiency: a useful metric for lead selection. *Drug Discovery Today.* 2004;9(10): 430-431. https://doi.org/https://doi.org/10.1016/S1359-6446(04)03069-7.

12. Huang X-Y, Shue, CY. Methods for Inhibiting Fascin. WO/2014/031732; 2014.

13. Friesner RA, Murphy RB, Repasky MP, et al. Extra precision glide: docking and scoring incorporating a model of hydrophobic enclosure for protein-ligand complexes. *J Med Chem.* 2006;49(21): 6177-6196. https://doi.org/10.1021/jm051256o.

14. Deb K, Agrawal S, Pratap A, Meyarivan T. A Fast Elitist Nondominated Sorting Genetic Algorithm for Multi-objective Optimization: NSGA-II. Berlin, Heidelberg: Springer Berlin Heidelberg; 2000:849-858.

15. Bickerton GR, Paolini GV, Besnard J, Muresan S, Hopkins AL. Quantifying the chemical beauty of drugs. *Nat Chem.* 2012;4(2): 90-98. https://doi.org/10.1038/nchem.1243.

16. Deng Z, Chuaqui C, Singh J. Structural interaction fingerprint (SIFt): a novel method for analyzing three-dimensional protein-ligand binding interactions. *J Med Chem.* 2004;47(2): 337-344.

https://doi.org/10.1021/jm030331x.

 Sastry GM, Dixon SL, Sherman W. Rapid shape-based ligand alignment and virtual screening method based on atom/feature-pair similarities and volume overlap scoring. *J Chem Inf Model.* 2011;51(10): 2455-2466. https://doi.org/10.1021/ci2002704.

Truchon JF, Pettitt BM, Labute P. A Cavity Corrected 3D-RISM
 Functional for Accurate Solvation Free Energies. *J Chem Theory Comput.* 2014;10(3): 934-941. https://doi.org/10.1021/ct4009359.

19. Yamashiro-Matsumura S, Matsumura F. Purification and characterization of an F-actin-bundling 55-kilodalton protein from HeLa cells. *J Biol Chem.* 1985;260(8): 5087-5097.

20. Grošelj U, Pušavec E, Golobič A, Dahmann G, Stanovnik B, Svete J. Synthesis of 1,5-disubstituted-4-oxo-4,5-dihydro-1H-pyrazolo[4,3-c]pyridine-7-carboxamides. *Tetrahedron.* 2015;71(1): 109-123. https://doi.org/10.1016/j.tet.2014.11.034.

Structure-based design, synthesis and biological evaluation of a novel series of isoquinolone and pyrazolo[4,3-*c*]pyridine inhibitors of fascin 1 as potential anti-metastatic agents

