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Second-generation tricyclic pyrimido-pyrrolo-oxazine mTOR inhibitor with predicted blood-brain barrier permeability†

Chiara Borsari, [‡]a Erhan Keles, [‡]a Andrea Treyer, ^b Martina De Pascale, ^a Paul Hebeisen, ^{ac} Matthias Hamburger ^b and Matthias P. Wymann ^{*a}

Highly selective mTOR inhibitors have been discovered through the exploration of the heteroaromatic ring engaging the binding affinity region in mTOR kinase. Compound **11** showed predicted BBB permeability in a MDCK-MDR1 permeability *in vitro* assay, being the first pyrimido-pyrrolo-oxazine with potential application in the treatment of neurological disorders.

Introduction

The mechanistic target of rapamycin (mTOR) operates downstream of phosphoinositide 3-kinase (PI3K). The mTOR protein kinase is part of two functionally distinct multiprotein mTOR complexes, mTORC1 and mTORC2.^{1,2} mTOR promotes cell growth and proliferation and is dysregulated in cancer and central nervous system (CNS) disorders,³ such as Parkinson's, Alzheimer's, Huntington's disease, and epilepsy.⁴ Tuberous sclerosis complex (TSC) is a negative regulator of TORC1. TSC loss-of-function causes hyperactivation of TORC1 signalling leading to the development of tuberous sclerosis, which causes epilepsy, behavioural changes and cognitive impairment in >90% of patients. Rapamycin is an allosteric inhibitor of TORC1, and is the prototype of first generation mTOR inhibitors. Everolimus, a rapalog, showed efficacy in decreasing the frequency and severity of epileptic seizures and has been approved for the treatment of TSC patients in 2018.⁵ Rapalogs have, however, significant immunosuppressive effects, which are pronounced due to their limited ability to cross the blood brain barrier (BBB).⁶ Therefore, the identification of highly selective mTOR inhibitors with optimized brain permeability is an ongoing challenge for the treatment of CNS disorders. Lead optimization strategies on different scaffolds led to the discovery of ATP-competitive mTOR kinase inhibitors (TORKi).

targeting both mTORC1 and mTORC2, including CC-223,⁷ INK128,⁸ AZD2014,⁹ PQR620,¹⁰ PQR626 (ref. 11) and a thiazolopyrimidine derivative.¹² Recently, we have disclosed a conformational restriction strategy to identify highly selective TORKi.¹³ A systematic structure-activity relationship (SAR) study pinpointed the pyrazine derivative **12b** as potent inhibitor of mTOR kinase ($K_i = 8.0$ nM), showing a 212-fold selectivity over the structurally related PI3K α (see Table 1 for chemical structure).¹⁴ However, **12b** displayed a limited ability to cross the BBB, preventing its application in the treatment of CNS disorders such as epilepsy. Given the unexplored activity of the tricyclic pyrimido-pyrrolo-oxazine compounds in CNS indications, we selected this privileged scaffold as starting point for the development of novel lead compounds with potential brain permeability.

Study design and synthesis

Herein, we investigated the effect of the heteroaryl moiety, engaging the binding affinity region in the mTOR kinase, on ligand potency and brain penetration. In several compound classes, isosterism between 6-membered diazaheterocycles and thiadiazole derivatives has been exploited to improve potency, enhance selectivity, alter physical properties, modify and reduce metabolism.¹⁵ The introduction of fluorine into molecular scaffolds¹⁶ is known to modulate lipophilicity, and plays an important role in BBB permeability.

We explored different bioisosters, and difluoro/trifluoro-substituted heteroaromatic rings on the tricyclic scaffold. The synthetic procedures for library preparation are reported in Schemes 1 and 2. Sulfamidate **22** was prepared starting from a chiral amino alcohol – (S)-serine – in several steps (ESI† for details).^{17,18} The tricyclic pyrimido-pyrrolo-oxazine scaffold was generated by sequential alkylation and a nucleophilic aromatic substitution reaction of dichloropyrimidine **12** with

^a Department of Biomedicine, University of Basel, Mattenstrasse 28, 4058 Basel, Switzerland. E-mail: Matthias.Wymann@Unibas.ch

^b Pharmaceutical Biology, Pharmacenter, University of Basel, Klingelbergstrasse 50, 4056 Basel, Switzerland

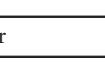
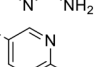
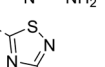
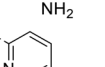
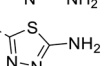
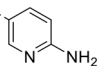
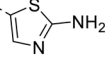
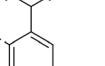
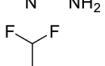
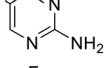
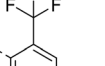
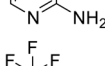
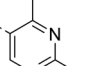
^c PIQUR Therapeutics AG, Hochbergerstrasse 60, 4057 Basel, Switzerland

† Electronic supplementary information (ESI) available: Experimental procedures and characterisation for all new compounds. Experimental details and raw data for biological assays, ¹H, ¹³C NMR, HRMS spectra and HPLC chromatograms for all novel compounds. See DOI: [10.1039/d0md00408a](https://doi.org/10.1039/d0md00408a)

‡ These authors have contributed equally.



Table 1 Heteroaromatic rings investigation

		Cellular assays IC_{50}^a [nM]		<i>In vitro</i> binding assays K_i^b [nM]					
Name	Ar	pPKB S473	pS6 S235/236	PI3K α	mTOR	SI c	clog P^d	PSA d	
12b e		94 \pm 2.5	60 \pm 13	1695 \pm 277	8.0 \pm 0.92	212	1.74	102.5	
1 e		77 \pm 6.8	55 \pm 5.5	536 \pm 22	5.1 \pm 0.4	105	1.85	102.5	
2		512 \pm 59	261 \pm 7.1	806 \pm 20	58 \pm 5.8	14	2.38	102.5	
3		162 \pm 3.1	84 \pm 20	847 \pm 216	11 \pm 2.4	77	1.84	102.5	
4		135 \pm 14	80 \pm 7.8	1547 \pm 278	8.2 \pm 0.6	189	1.80	102.5	
5 e		133 \pm 16	61 \pm 12	2147 \pm 400	10 \pm 1.6	215	2.59	89.6	
6		157 \pm 5.4	102 \pm 4.2	2485 \pm 251	15 \pm 2.8	166	2.37	89.6	
7		253 \pm 10	67 \pm 4.8	232 \pm 33	12 \pm 0.7	19	2.82	89.6	
8		116 \pm 7.5	119 \pm 9.9	17 \pm 0.30	34 \pm 9.9	0.5	2.33	102.5	
9		1089 \pm 34	776 \pm 138	272 \pm 10	118 \pm 9.6	2.3	3.46	89.6	
10		446 \pm 45	694 \pm 131	55 \pm 0.93	249 \pm 39	0.22	3.11	102.5	
11		549 \pm 56	348 \pm 44	>20 000	33 \pm 1.3	>600	3.46	89.6	

^a IC₅₀s were measured using a 7-point 1:2 serial dilution and each concentration was measured in triplicate. ^b Compounds were tested for the *in vitro* binding to the ATP-binding site of PI3K α and mTOR using a commercially available LanthaScreen time-resolved FRET (TR-FRET) displacement assay. IC₅₀s were measured using a 10-point 1:4 serial dilution. Each concentration was performed in duplicate, IC₅₀ to K_i conversion factor = 9.84 (p110 α), 2.05 (mTOR). Data are reported as mean \pm SD. ^c Selectivity index calculated as $K_i(\text{mTOR})/K_i(\text{p110}\alpha)$. ^d Calculation of log P and PSA was performed with ChemAxon (<https://www.chemaxon.com>). ^e Previously reported in ref. 14.

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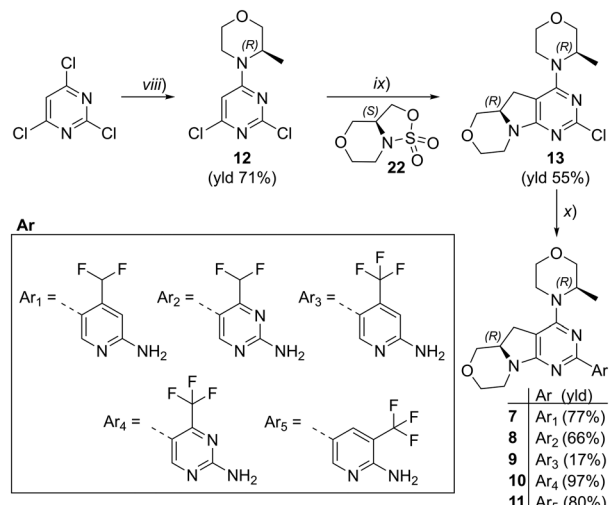
(S)-sulfamide **22** using *n*-BuLi and copper(I) iodide (cat.) followed by acidic hydrolysis of the intermediary sulfaminic acid and base-mediated ring closure.¹⁴ To introduce a 1,2,4-thiadiazole and a pyrazine, a tributyltin group was appended on **13**, followed by Stille coupling between **14** and halides (Scheme 1a). The Stille reaction of 3-bromo-5-chloro-1,2,4-

thiadiazole was regioselective for C-5 coupling to give bromo derivative **15**. In contrast, no site-selective cross-coupling between the tin derivative **14** and 3-chloro-6-iodopyridazine was observed, and a mixture of chloro and iodo derivatives was obtained (**16**, 50% chloro, 44% iodo; see ESI†). The heteroaryl halide **16** was aminated by treatment with aqueous

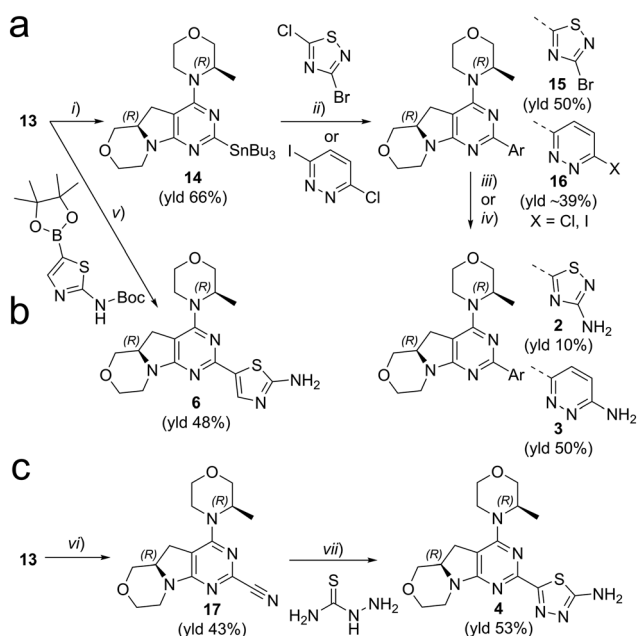
ammonia at 100 °C to yield the desired pyridazine-substituted compound **3** in moderate yield. The conversion of the bromide **15** into an amine by using ammonia was not successful. *tert*-Butyl carbamate in presence of caesium carbonate was employed for the Pd-catalysed synthesis of the *N*-Boc-protected derivative of **15**. In a one-pot procedure, treatment with HCl in dioxane generated the final 1,2,4-thiadiazole derivative **2**. To prepare the 1,3,4-thiadiazole-substituted compound **4**, the chloropyrimidine **13** was converted into the nitrile **17** using tetraethylammonium cyanide in presence of DABCO. Coupling of nitrile **17** and thiosemicarbazide followed by cyclization at 65 °C in presence of TFA yielded compound **4** (Scheme 1c). Subsequently, thiazole and difluoromethyl- and trifluoromethyl-substituted pyridines/pyrimidines were installed on intermediate **13** using a Suzuki cross-coupling reaction with boronic acid pinacol esters to generate the final compounds **6–11** (Schemes 1b and 2).

Results and discussion

The library was tested for *in vitro* binding to recombinant mTOR and PI3K α (K_i), and for PI3K/mTOR signalling in A2058 melanoma cells (IC_{50} for phosphorylated S6 ribosomal protein [pS6, Ser235/236] and phosphorylated protein kinase B [PKB/Akt, Ser473] to detect mTORC1 and mTORC2 activity). The *in vitro* and cellular profiling of the novel set of



Scheme 2 Synthesis of compounds **7–11**. Reagents and conditions: (viii) morpholine derivative, DIPEA, DCM, 0 °C → rt, o/n; (ix) 1) *n*-BuLi, CuI, -78 °C → r.t., o/n; 2) HCl conc., MeOH, 45 °C, 4–6 h; 3) NaOH, H₂O, r.t., 1–16 h; (x) 1) boronic ester (for **7, 9–11**) or boronic ester generated *in situ* (for **8**), XPhosPdG2 (cat.), K₃PO₄, dioxane/H₂O, 95 °C, 2–16 h; 2) HCl, dioxane/H₂O, 60 °C, 3–16 h (for **7–9, 11**). See ESI† for details.



Scheme 1 Synthesis of compounds **2–4, 6**. Reagents and conditions: (a) (i) Pd(amphos)Cl₂, bis(tributyltin), dioxane, reflux, 3 h; (ii) halide derivative, Pd(PPh₃)₄, dioxane, 90 °C, o/n; (iii) 1) Pd₂(bda)₃/xantphos, *tert*-butyl carbamate, Cs₂CO₃, dioxane, 90 °C, o/n, 2) HCl in dioxane (for **2**); (iv) NH₃ aq. DMSO, 100 °C, o/n (for **3**). (b) (v) 1) boronic ester, XPhosPdG2 (cat.), K₃PO₄, dioxane/H₂O, 95 °C, 2–16 h; 2) HCl, dioxane/H₂O, 60 °C, 3–16 h; (c) (vi) tetraethylammonium cyanide, DABCO, DMSO, 140 °C, 1.5 h. (vii) TFA, 65 °C, o/n. See ESI† for details.

conformationally restricted compounds is reported in Table 1. Pyrimidine (**1**), pyrazine (**3**) and pyridine (**5**) have been compared with their bioisosteres involving the replacement of $-\text{CH}=\text{CH}-$ by sulphur. Replacement of the pyrimidine with a 1,2,4-thiadiazole led to a >10-fold decrease in mTOR affinity, reducing compound selectivity [K_i (mTOR) **1** = 5.1 nM, **2** = 58 nM; K_i (PI3K α)/ K_i (mTOR) **1** = 105, **2** = 14]. On the contrary, comparable activity and selectivity were observed for the bioisosteric pairs **3–4** and **5–6** (Table 1). The presence of a difluoro or trifluoromethyl group in *para*-position of 2-amino pyridines (**7** and **9**) caused a selectivity drop compared to **12b** [K_i (PI3K α)/ K_i (mTOR) < 20]. The pyrimidine derivatives **8** and **10** were more potent on PI3K α than on mTOR kinase [K_i (PI3K α)/ K_i (mTOR) < 1]. A gain in mTOR potency and selectivity over PI3K was achieved moving the CF₃-group from *para* (compound **9**) to *meta* position (**11**) on the pyridine scaffold. **11** showed the highest selectivity [K_i (PI3K α)/ K_i (mTOR) > 600] among these tricyclic derivatives, and outperformed the selectivity of **12b**. Compound **11** displayed a moderate cellular activity (IC_{50} for phosphorylated PKB/Akt and for phosphorylated S6 < 550 nM).

While difluoromethyl- and trifluoromethyl-substituted pyridines/pyrimidines have been explored in pre-clinical and clinical candidates targeting the PI3K-mTOR pathway,^{10,19–21} to the best of our knowledge no mTOR inhibitor bearing thiazole or 1,3,4-thiadiazole moieties has been disclosed yet. With no prior hints to stability information, we characterised compounds **4** and **6** with respect to their CYP related metabolism, after incubation with individual human recombinant CYP isoenzymes CYP1A1 and CYP1A2. These isoenzymes have been selected since they are known to be implicated in the metabolism of structurally similar molecules.¹⁹ **4** was highly stable with both CYP1A1 and 1A2



(>70% remaining compound after 60 minutes of incubation, Fig. 1). **6** displayed moderate to high stability with CYP1A1 and CYP1A2 (48% and 87% remaining compound, respectively), and comparable to that of the corresponding bioisosteric compound **5** (Fig. 1). Moreover, the (*R*)-3-methylmorpholine derivative **5** presented a higher stability with 1A1 isoenzyme with respect to the corresponding unsubstituted-morpholine **27** (see ESI†). Due to the highly variable expression of CYP1A1 in liver and extrahepatic tissues, the high stability towards degradation by CYP1A1 isoenzyme is an asset in drug development and facilitates the establishment of a therapeutic dose.

The observed SAR was rationalized using computational modelling studies. We investigated the putative interactions of **1–6** and **11** with ATP-binding sites of mTOR and PI3K α . After energy minimization, the mTOR-ligand complexes maintained pivotal interactions including H-bonds (i) between Asp2195 and the NH₂-group of the aminopyridine, and (ii) between the oxygen atom of the (*R*)-3-methylmorpholine and the hinge region backbone amide of Val2240. The (*R*)-3-methylmorpholine has been previously reported as a crucial feature to achieve selectivity for mTOR kinase over PI3Ks.^{12,14} The orientation of the amino group on the 1,3,4-thiadiazole in **4** and on thiazole in **6** resembles that of the 2-amino substituted pyrimidine (**1**), pyrazine (**3**) and pyridine (**5**, Fig. 2b for **3–4**), leading to the establishment of an H-bond network with Asp2195 and Glu2190. On the contrary, the NH₂-group on the 1,2,4-thiadiazole (**2**) is 2.8 Å away from the 2-amino group of **1**. The different orientation prevents an H-bond interaction with Glu2190 (6 Å distance), which plays a pivotal role in the stabilization of inhibitor binding (Fig. 2a). This explains the reduced affinity for mTOR kinase of **2** with respect to compounds **1, 3–6**. Docking studies of compound **11** into PI3K α and mTOR revealed that the 3-trifluoromethyl group is well accommodated in mTOR (Fig. 2c), while it induces steric clashes with Lys802 and Asp933 in PI3K α (Fig. 2d). This is due to the different orientation of Lys2187 (mTOR) and Lys802 (PI3K α), which leads to a broader binding affinity pocket in mTOR compared to that of PI3K α . The orientation of Lys2187 (mTOR)/Lys802

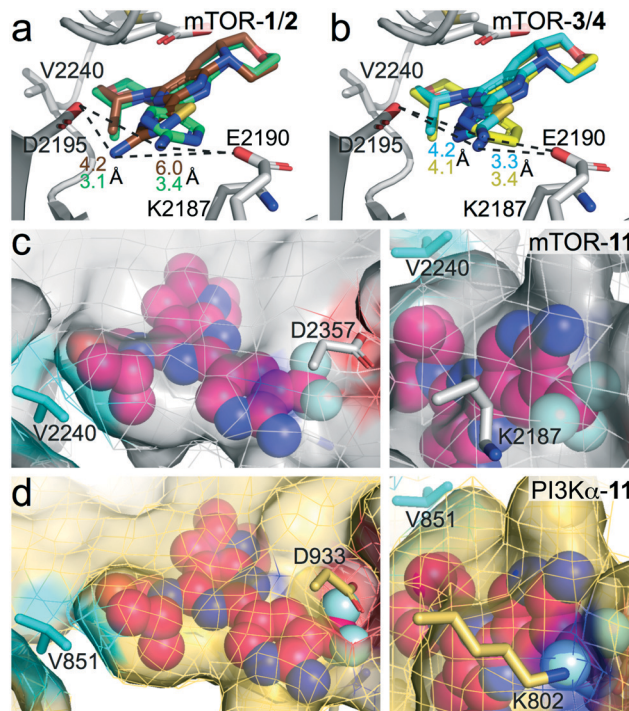


Fig. 2 (a and b) Superimposition of (a) mTOR-1 and mTOR-2 complexes and (b) mTOR-3 and mTOR-4 complexes. mTOR, grey; 1, green; 2, brown; 3, yellow; 4, cyan. The distances between the 2-amino group and Glu2190/Asp2195 are depicted as dashed black lines. Distances values are coloured as the inhibitors. (c and d) Docking of compound **11** (magenta) to (a) mTOR (grey) starting from PDB 4JT6,²² and (b) PI3K α (wheat) starting from PDB 6OAC.²¹ No steric clashes are present with Asp2357 and Lys2187 in mTOR (c). Steric clashes with Asp933 and Lys802 in PI3K α are shown (d).

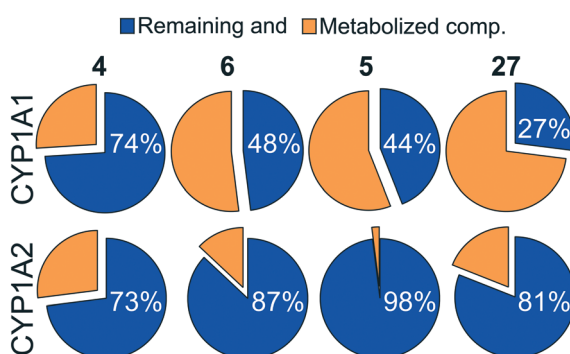


Fig. 1 Pie chart showing the percentage of remaining compounds **4–6** and **27** (blue) after 60 min of incubation with the indicated human recombinant CYP isoenzymes.

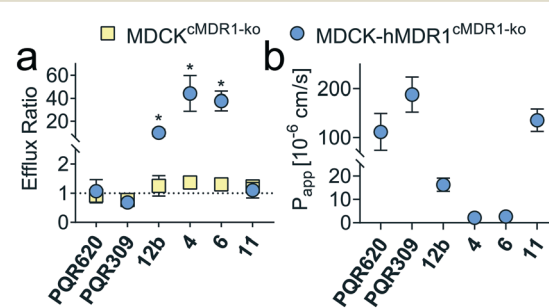


Fig. 3 (a) Efflux ratio (ER) in MDCK cells with and without hMDR1 grown on TransWell filters. An ER of 1 is indicative of the compound not being a hMDR1 substrate. (b) Apparent permeability (P_{app}) from apical to basolateral site in MDCK-hMDR1^{cMDR1-ko} cells. Numerical values are reported in Tables S1 and S2.† * $p < 0.05$, unpaired *t*-test, GraphPad Prism 8.0.2.



employed,²³ and the results compared with a control cell line expressing neither MDR1 ortholog (MDCK^{MDR1-ko}). Compounds **4** and **6**, as well as **12b** included as negative control, were identified as hMDR1 substrates (ER > 20) with low permeability ($P_{app} < 60 \times 10^{-6}$ cm s⁻¹), predicting poor BBB permeability *in vivo*. On the contrary, compound **11** was no hMDR1 substrate (ER ~ 1) and showed a P_{app} comparable to PQR309¹⁹ and PQR620,¹⁰ which are known brain penetrant PI3K and/or mTOR inhibitors (Fig. 3).

Conclusion

In summary, we have identified 3-CF₃-substituted pyridine, thiazole and 1,3,4-thiadiazole as unexplored heteroaromatic rings for mTOR inhibitors providing an exquisite mTOR selectivity. We have reported the first tricyclic pyrimido-pyrrolo-oxazine derivative showing MDCK-based BBB permeability. Altogether, the good *in vitro*/cellular potency, and the predicted brain penetration qualify **11** as second-generation tricyclic pyrimido-pyrrolo-oxazine with potential application in the treatment of CNS disorders.

Conflicts of interest

PH is a past employee of PIQUR Therapeutics AG, Basel; and PH and MPW are shareholders of PIQUR Therapeutics AG.

Acknowledgements

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References

- 1 M. P. Wymann and R. Schneiter, *Nat. Rev. Mol. Cell Biol.*, 2008, **9**, 162–176.
- 2 D. D. Sarbassov, D. A. Guertin, S. M. Ali and D. M. Sabatini, *Science*, 2005, **307**, 1098–1101.
- 3 M. P. Wymann and R. Marone, *Curr. Opin. Cell Biol.*, 2005, **17**, 141–149.
- 4 Z. Z. Chong, Y. C. Shang, S. Wang and K. Maiese, *Future Neurol.*, 2012, **7**, 733–748.
- 5 L. Lechuga and D. N. Franz, *Expert Rev. Neurother.*, 2019, **19**, 913–925.
- 6 C. Brandt, P. Hillmann, A. Noack, K. Römermann, L. A. Öhler, D. Rageot, F. Beaufils, A. Melone, A. M. Sele, M. P. Wymann, D. Fabbro and W. Löscher, *Neuropharmacology*, 2018, **140**, 107–120.
- 7 Z. Jin, H. Niu, X. Wang, L. Zhang, Q. Wang and A. Yang, *Oncotarget*, 2017, **8**, 58469–58479.
- 8 E. K. Slotkin, P. P. Patwardhan, S. D. Vasudeva, E. de Stanchina, W. D. Tap and G. K. Schwartz, *Mol. Cancer Ther.*, 2015, **14**, 395–406.
- 9 K. G. Pike, K. Malagu, M. G. Hummersone, K. A. Menear, H. M. Duggan, S. Gomez, N. M. Martin, L. Ruston, S. L. Pass and M. Pass, *Bioorg. Med. Chem. Lett.*, 2013, **23**, 1212–1216.
- 10 D. Rageot, T. Bohnacker, A. Melone, J. B. Langlois, C. Borsari, P. Hillmann, A. M. Sele, F. Beaufils, M. Zvelebil, P. Hebeisen, W. Löscher, J. Burke, D. Fabbro and M. P. Wymann, *J. Med. Chem.*, 2018, **61**, 10084–10105.
- 11 C. Borsari, E. Keles, D. Rageot, A. Treyer, T. Bohnacker, L. Bissegger, M. De Pascale, A. Melone, R. Sriramaratnam, F. Beaufils, M. Hamburger, P. Hebeisen, W. Löscher, D. Fabbro, P. Hillmann and M. P. Wymann, *J. Med. Chem.*, 2020, **63**(22), 13595–13617.
- 12 S. Bonazzi, C. P. Goold, A. Gray, N. M. Thomsen, J. Nunez, R. G. Karki, A. Gorde, J. D. Biag, H. A. Malik, Y. Sun, G. Liang, D. Lubicka, S. Salas, N. Labbe-Giguere, E. P. Keaney, S. McTighe, S. Liu, L. Deng, G. Piuzzi, F. Lombardo, D. Burdette, J.-C. Dodart, C. J. Wilson, S. Peukert, D. Curtis, L. G. Hamann and L. O. Murphy, *J. Med. Chem.*, 2020, **63**, 1068–1083.
- 13 V. Cmiljanovic, P. Hebeisen, E. Jackson, F. Beaufils, T. Bohnacker and M. P. Wymann, WO2015049369, 2015.
- 14 C. Borsari, D. Rageot, A. Dall'Asen, T. Bohnacker, A. Melone, A. M. Sele, E. Jackson, J.-B. Langlois, F. Beaufils, P. Hebeisen, D. Fabbro, P. Hillmann and M. P. Wymann, *J. Med. Chem.*, 2019, **62**, 8609–8630.
- 15 N. A. Meanwell, *J. Med. Chem.*, 2011, **54**, 2529–2591.
- 16 J. J. Danon, T. A. Reekie and M. Kassiou, *Trends Chem.*, 2019, **1**, 612–624.
- 17 J. F. Rousseau, I. Chekroun, V. Ferey and J. R. Labrosse, *Org. Process Res. Dev.*, 2015, **19**, 506–513.
- 18 P. Hebeisen, A. Alker and M. Buerkler, *Heterocycles*, 2012, **85**, 65–72.
- 19 F. Beaufils, N. Cmiljanovic, V. Cmiljanovic, T. Bohnacker, A. Melone, R. Marone, E. Jackson, X. Zhang, A. Sele, C. Borsari, J. Mestan, P. Hebeisen, P. Hillmann, B. Giese, M. Zvelebil, D. Fabbro, R. L. Williams, D. Rageot and M. P. Wymann, *J. Med. Chem.*, 2017, **60**, 7524–7538.
- 20 C. Borsari, D. Rageot, F. Beaufils, T. Bohnacker, E. Keles, I. Buslov, A. Melone, A. M. Sele, P. Hebeisen, D. Fabbro, P. Hillmann and M. P. Wymann, *ACS Med. Chem. Lett.*, 2019, **10**, 1473–1479.
- 21 D. Rageot, T. Bohnacker, E. Keles, J. A. McPhail, R. M. Hoffmann, A. Melone, C. Borsari, R. Sriramaratnam, A. M. Sele, F. Beaufils, P. Hebeisen, D. Fabbro, P. Hillmann, J. E. Burke and M. P. Wymann, *J. Med. Chem.*, 2019, **62**, 6241–6261.
- 22 H. Yang, D. G. Rudge, J. D. Koos, B. Vaidalingam, H. J. Yang and N. P. Pavletich, *Nature*, 2013, **497**, 217–223.
- 23 M. Karlgren, I. Simoff, M. Backlund, C. Wegler, M. Keiser, N. Handin, J. Müller, P. Lundquist, A. C. Jareborg, S. Oswald and P. Artursson, *J. Pharm. Sci.*, 2017, **106**, 2909–2913.

