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# Further optimization of the M<sub>1</sub> PAM VU0453595: Discovery of novel heterobicyclic core motifs with improved CNS penetration

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# Abstract

This letter describes the continued chemical optimization of the VU0453595 series of  $M_1$  positive allosteric modulators (PAMs). By surveying alternative 5,6- and 6,6-heterobicylic cores for the 6,7-dihydro-5*H*-pyrrolo[3,4-*b*]pyridine-5-one core of VU453595, we found new cores that engendered not only comparable or improved  $M_1$  PAM potency, but significantly improved CNS distribution ( $K_ps$  0.3 to 3.1). Moreover, this campaign provided fundamentally distinct  $M_1$  PAM chemotypes, greatly expanding the available structural diversity for this valuable CNS target, devoid of hydrogen-bond donors.

# **Graphical Abstract**



#### Keywords

M<sub>1</sub>; Muscarinic acetylcholine receptor; Positive allosteric modulator (PAM); CNS penetration; Structure-Activity Relationship (SAR)

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Positive allosteric modulators (PAMs) of the muscarinic acetylcholine receptor subtype 1 (M<sub>1</sub>) have garnered a great deal of attention as a novel therapeutic approach for the treatment of the cognitive and negative symptom domains of schizophrenia, especially targeting NMDA receptor hypofunction.<sup>1-8</sup> Moreover, M<sub>1</sub> PAMs are also of interest in general cognition enhancement and Alzheimer's disease.<sup>1-4,9-12</sup> Since we reported on the discovery of the first M1 PAM, BQCA,<sup>13,14</sup> numerous M1 PAMs have been reported in the primary and patent literature (most by scaffold hopping VU and Merck series) with many conserved moieties that consistently engender low CNS penetration ( $K_p s < 0.3$ ).<sup>15-25</sup> Our latest entry into M1 PAMs was the result of three distinct high-thoughput screening campaigns,<sup>26</sup> which resulted in novel indole-, azaindole- and isatin-based M<sub>1</sub> PAM scaffolds.<sup>25,27-29</sup> Of these, the isatin VU0119498 (1) was a unique PAM in that it potentiated all of the  $G_{q}$ -coupled mAChRs (M<sub>1</sub>, M<sub>3</sub> and M<sub>5</sub>) with equivalent potency and efficacy.<sup>27</sup> Subsequent optimization efforts identified 'molecular switches' that gave rise to a series of highly selective  $M_5$  PAMs,  $^{30-32}$  as well as ML137 (2), a highly selective  $M_1$  PAM by virtue of the fluorophenyl pyrazole moiety.<sup>28</sup> Carbonyl deletion provided lactam 3, and surveying regioisomeric lactams afforded VU0451725 (4), with improved DMPK properties over 2 and **3**.<sup>33</sup> Finally, an aza-scan identified the the 6,7-dihydro-5*H*-pyrrolo[3,4-*b*]pyridine-5-one core of VU0453595 (5), which proved a useful in vitro and in vivo tool, demonstrating efficacy in rodent models of pharmacologically-induced NMDA hypofunction.<sup>33</sup> In this Letter, we detail an optimization campaign surveying alternative 5,6- and 6,6-heterobicyclic cores, alternate moieties for the pyrazole, and walking additional fluorines around the central phenyl ring to ultimately provide multiple novel M<sub>1</sub> PAM scaffolds with comparable or improved rat M<sub>1</sub> PAM potencies and improved CNS distribution (K<sub>p</sub>s 0.4 to 3.1).

The chemistry to access new analogs, if not commercially available, was straightforward (**Scheme 1**).<sup>34</sup> The fluorinated heterobiaryl tail moities were readily prepared in two steps as either a benzyl chloride **7** or a benzyl mesylate **8** from commercial benzyl alcohols **6**. Various 5,6- and 6,6-heterobiaryl systems were then alkylated with either **7** or **8** to provide analogs **10**. A subsequent Suzuki coupling installed the heterobiaryl motif, delivering analogs **11**. Quinolinone and naphthyridinone analogs **11** of **9**, were made in a single step from **12**, and based on our previous work, cores such as **15** were also accessed in a simple three step procedure.<sup>34</sup>

SAR was steep for the diverse analogs **11**, with many compounds devoid of  $M_1$  PAM activity on both human or rat  $M_1$ , or displaying species bias towards rat  $M_1$  PAM activity. In general, the 2,6-difluoro analogs were active whereas mono-fluoro and *des*-fluoro phenyl congeners were inactive as  $M_1$  PAMs. Representative SAR is shown in Table 1 for a subset of analogs **16**, possessing an *N*-Me-indazole attached at the 4-position to the 2,6-difluorophenyl ring. While only rat  $M_1$  data is shown, analogs **16** were uniformly 2- to 3-fold less potent on human  $M_1$  (with many > 10 µM). Here, various 6,6-hetrobicyclic ring systems were comparably active (r $M_1$  EC<sub>50</sub>s 4.3-4.9 µM) across quinazolin-4(3*H*)-ones (**16a**), pyrido[3,4-*d*]pyrimidin-4(3*H*)-ones (**16b**), quinoxalin-2(1*H*)-ones (**16c**) and naphthyridin-5(6*H*)-ones (**16d**). These analogs possessed favorable *in vitro* DMPK profiles (rat and human  $f_u$ s of 0.01 to 0.04) and moderate predicted hepatic clearance (CL<sub>hep</sub>s of 40-44 mL/min/kg). However, they were superior to the lead **5** in terms of brain distribution

(K<sub>p</sub>), wherein **16a-d** displayed K<sub>p</sub>s (rat brain:plasma ratios) of 0.35 to 2.16, and when corrected for fraction unbound in plasma and brain homogenate binding, the K<sub>puu</sub>s ranged from 0.3 to 0.77 – a major advance in the context of M<sub>1</sub> PAMs. Notably, **16c** (VU0478436) afforded a >6-fold increase in CNS penetration over **5**. The 5,6-congener **16e** (VU0486691),based on a dihydroimidazol[1,2-*c*]pyrimidin-5(3*H*)-one core, showed enhanced M<sub>1</sub> PAM potency (rM<sub>1</sub> EC<sub>50</sub> = 1.7  $\mu$ M, 50% ACh Max), improved *in vitro* DMPK profile (rat and human *f*<sub>u</sub>s of 0.08 and 0.04, respectively and moderate rat predicted hepatic clearance (CL<sub>hep</sub> = 40 mL/min/kg)). Moreover, **16e** demonstrated a rat K<sub>p</sub> of 0.35 and a K<sub>puu</sub> of 0.3. This finding led us to explore additional 5,6-heterobicyclic cores.

SAR proved steep as additional 5,6-hetrobicyclic cores were prepared and evaluated, with the vast majority devoid of  $M_1$  PAM activity. During this effort, it was also shown that regioisomeric *N*-Me indazoles had a profound effect on  $M_1$  PAM activity (**Fig. 2**). Interestingly, the 4-positional isomer **17** was devoid of  $M_1$  PAM activity, while in contrast, the 5-positional isomer **18** was a potent  $M_1$  PAM (EC<sub>50</sub> = 1.7 µM, 50% ACh Max, pEC<sub>50</sub> = 5.76+0.02) with very attractive *in vitro* DMPK properties (rat and human  $f_u$ s of 0.06 and 0.05, respectively, low rat hepatic clearance (CL<sub>hep</sub> = 29 mL/min/kg) and a large free fraction in rat brain homogenate binding,  $f_u = 0.098$ ). Moreover, **18** (VU0484061) possessed a rat brain:plasma ratio (K<sub>p</sub>) of 0.40 and a K<sub>puu</sub> of 0.70. Once again, mL/min/kg). However, they were superior to the lead **5** in terms and in comparison to the known  $M_1$  PAMs with low K<sub>p</sub>s and K<sub>puu</sub>s, both **16c** and **18** truly stand out. It is important to point out that the majority of M<sub>1</sub> PAMs possess two or more hydrogen bond donors (typically a *trans*-2-hydroxy cylohexyl amide moiety) that likely engenders the poor CNS penentration due to P-gp efflux or low permeability.<sup>13-25</sup>

Prior to leaving this unique sub-series of M<sub>1</sub> PAMs, one last library of analogs was prepared with more diverse tail pieces within the 16b and 16c 6,6-heterobicyclic cores. Again, SAR was steep, and few active M1 PAM resulted. However, this last campaign afforded three M1 PAMs 19-21 with diverse profiles (Fig. 3). Here, the pyrido[2,3-b]pyrazin-2(1H)-one 19 (VU0486384) was a potent and efficacious rat M<sub>1</sub> PAM (EC<sub>50</sub> =  $2.8 \mu$ M,  $80\pm1\%$  ACh Max, pEC<sub>50</sub> = 5.56 $\pm$ 0.12), with a favorable fraction unbound in plasma (rat and human  $f_{us}$  of 0.04), high predicted hepatic clearance ( $CL_{hep} = 60 \text{ mL/min/kg}$ ), yet excellent CNS pentration ( $K_p = 1.64$  and  $K_{puu} = 1.1$ ). The nature of the heterobiaryl moiety played a key role in analogous pyrido [3,4-d] pyrimidin-4-(3H)-one congeners 20 and 21. The isoquinoline analog **20** was a weak rat M<sub>1</sub> PAM (EC<sub>50</sub> = 6.1  $\mu$ M, 42±3% ACh Max, pEC<sub>50</sub> = 5.21±0.11) with equivalent plasma fraction unbound ( $f_{us}$  of 0.03) for rat and human, and the best K<sub>p</sub> to date for an M1 PAM of 3.1 (and a Kpuu of 2.7). In sharp contrast, the more basic N-Me benzimidazole congener **21** was of comparable potency (EC<sub>50</sub> =  $5.3 \mu$ M,  $65\pm4\%$  ACh Max, pEC<sub>50</sub> = 5.28±0.14), good plasma fraction unbound (rat and human  $f_{us}$  of 0.04 and 0.06, respectively), but no detectable CNS penetration (brain levels below the level of quantitation, BLQ). These data show that subtle pKa modulation can dramtically impact Kn.

Finally, the concept of divergent signal bias, mediated by stabilization of unique conformers of the GPCR by the allosteric ligand, has emerged, and in many instances is critical for avoiding adverse effect liabilites.<sup>1-4,35-38</sup> Thus, our lab surveys the propensity of new  $M_1$  PAM ligands to display signal bias.<sup>38-41</sup> For VU0453595 (**5**), DiscoverRX assessed activities

of the M<sub>1</sub> PAM against human M<sub>1</sub> in a calcium flux assay, as well on β-arrestin recruitment and internalization.<sup>39</sup> PAM **5** was shown to be a modest human M<sub>1</sub> PAM (EC<sub>50</sub> = 1.9 µM, 79% ACh Max) with no effect on receptor internalization (EC<sub>50</sub> >10 µM) and modest effect on β-arrestin recruitment (EC<sub>50</sub> = 2.6 µM, 57% Max). New M<sub>1</sub> PAM **16b** was evaluated similarly, and was found to be a modest human M<sub>1</sub> PAM (EC<sub>50</sub> = 5.3 µM, 65% ACh Max) with no effect on receptor internalization (EC<sub>50</sub> >10 µM), yet a submicromolar effect on βarrestin recruitment (EC<sub>50</sub> = 980 nM, 33% Max). At this point, the *in vivo* ramification of these profiles across signal transduction pathways are unclear, but we are tracking and noting differences between M<sub>1</sub> PAMs and plan to investigate more thoroughly once a

In summary, we report on the further optimization of the *in vivo* tool  $M_1$  PAM, VU0453595 (5). A diverse array of 5,6- and 6,6-heterobicyclic cores were developed as novel  $M_1$  PAMs with unprecedented levels of CNS penetration ( $K_ps$  0.3 to 3.1 and  $K_{puus}$  of 0.3 to 2.7) and lacking the prototypical hydrogen-bond donor motifs. While these  $M_1$  PAMs are too weak to advance as clinical candidates, the improved disposition of these new chemoptypes represent fundamentally new starting points for further chemical optimization. Additional refinements are in progress and will be reported in due course.

collection of M<sub>1</sub> PAM ligands with diverse profiles (and comparable PK) are accumulated.

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#### Figure 1.

Evolution of the development of the VU0119498 series of  $M_1$  PAMs, culminating in VU0453595 (**5**), a moderately potent PAM (r $M_1$  EC<sub>50</sub> = 3.2  $\mu$ M, 75% ACh Max and 3x less potent on h $M_1$ ) with modest CNS penetration ( $K_p$  = 0.3). In this work, we survey alternative 5,6- and 6,6-heterobicyclic cores and pyrrole replacements in an attempt to increase CNS penetration.



#### Figure 2.

The impact of positional isomers of the *N*-Me indazole in the context of dihydropyrazolo[1,5-*a*]pyrazin-4-(5*H*)-ones **17** and **18**.



## Figure 3.

Additional M<sub>1</sub> PAMs **19-21** based on 6,6-heterobicyclic cores with a diverse range of pharmacological and DMPK properties.



#### Scheme 1.

*Reagents and conditions*: (a) Ghosez's reagent or SOCl<sub>2</sub>, DCM, rt, 65-78%; (b) MsCl, Et N, DCM, 0 °C, 75-88%; (c) **7** or **8**, Cs<sub>2</sub>CO<sub>3</sub>, MeCN, 70 °C, 52-80%; (d) Het-B(OH), Pd(dppf)Cl, Cs<sub>2</sub>CO<sub>3</sub>, THF:H<sub>2</sub>0 (10:1),  $\mu$ w 140 °C, 22-69%; (e) ethyl glyoxalate, 51-68%; (f) Br(CH<sub>2</sub>)<sub>2</sub>NHBoc, Cs<sub>2</sub>CO<sub>2</sub>, DMF, rt, 16h, 98%; (g) HCl, 1,4-dioxane, rt, 1.5 h, 90%; (h) Na<sub>2</sub>CO<sub>3</sub>, 1,4-dioxane/H<sub>2</sub>0, rt 3 hr, 90%.

#### Table 1

Structures and activities of analogs 16.

Het F F 16				
Cpd	Het	rM <sub>1</sub> EC <sub>50</sub> (μM) <sup>a</sup> [% ACh Max ±SEM]	rM <sub>1</sub> pEC <sub>50</sub> (±SEM)	Rat K <sub>p</sub> (K <sub>p,uu</sub> ) <sup>b</sup>
16a		4.7 [45 <u>+</u> 4%]	5.32 <u>+</u> 0.10	0.79 (0.32)
16b	O N N	4.7 [73 <u>+</u> 5%]	5.32 <u>+</u> 0.08	0.77 (0.60)
16c	N N N N N N N N N N N N N N N N N N N	4.9 [67 <u>+</u> 3%]	5.31 <u>+</u> 0.02	2.16 (0.77)
16d		4.3 [52 <u>+</u> 4%]	5.37 <u>+</u> 0.07	0.52 (0.49)
16e	N N N	1.7 [50 <u>+</u> 5%]	5.77 <u>+</u> 0.04	0.35 (0.30)

<sup>a</sup>Calcium mobilization assays with rM<sub>1</sub>-CHO cells performed in the presence of an EC<sub>20</sub> fixed concentration of acetylcholine; values represent means from three (n=3) independent experiments performed in triplicate.

<sup>b</sup>Total and calculated unbound braimplasma partition coefficients determinec at 0.25 hr post-administration of an IV cassette dose (0.20-0.25 mg/kg) to male, SD rat (*n*=1), in conjunction with *in vitro* rat plasma protein and brain homogenate binding assay data.