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¹ C-Glucopyranosyl-1,2,4-triazoles As New Potent Inhibitors of 2 Glycogen Phosphorylase

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Supporting Information

ABSTRACT: Glycogen phosphorylase inhibitors are considered as potential antidiabetic agents. $3-(\beta-D-Glucopyranosyl)$ -5-substituted-1,2,4-triazoles were prepared by acylation of O-10

perbenzoylated N^1 -tosyl-C- β -D-glucopyranosyl formamidrazone and subsequent removal of the protecting groups. The best inhibitor was $3-(\beta-D-glucopyranosyl)-5-(2-naphthyl)-1,2,4-triazole (<math>K_i = 0.41 \mu M$ against rabbit muscle glycogen phosphorylase 12 13

KEYWORDS: 1,2,4-Triazole, C-glucopyranosyl derivative, bioisoster, glycogen phosphorylase, inhibitor

nhibitors of enzymes are among classics of medicinal 16 Lchemistry, and many drug molecules' activity is due to 17 decreasing the efficiency of these catalytic proteins. In a 18 chemical biological approach, finding an enzyme inhibitor is the 19 result of a good match of the biological and chemical spaces 20 represented by a binding site of an enzyme and a small 21 molecule, respectively, fitting to each other with considerable 22 strength. Among several methods to design inhibitors, 23 bioisosteric replacement of structural elements of existing 24 molecules is widely applied and in many cases results in higher 25 activity or other advantageous property of the new compound.² Glycogen phosphorylase (GP) is the main regulatory enzyme 27 of glycogen metabolism. GP, catalyzing the rate determining 28 step of glycogen degradation in the liver by phosphorolysis, is 29 directly responsible for the regulation of blood glucose levels. 30 Therefore, GP has been a validated target in combating 31 noninsulin-dependent or type 2 diabetes mellitus (T2DM), and 32 its inhibitors are considered as potential antidiabetic agents. The biochemical and pharmacological background of this research has been thoroughly summarized in several reviews of 35 the past decade; therefore, the reader is kindly referred to those ³⁶ papers. ^{3–5} Furthermore, possible application of GP inhibitors 37 in intervention of other diseased states associated with GP 38 activity (e.g., cardiovascular disorders, 6 ischemic lesions, 7,8 and 39 tumorous growth⁷) has also been under investigations.

Several classes of compounds^{9,10} were shown to be inhibitors 41 of GP. The most widely studied group of molecules is that of 42 glucose derivatives, 11,12 which bind primarily to the active site 43 of GP. 13 The best glucose derivatives are submicromolar 44 inhibitors of rabbit muscle GPb, the prototype of GPs. 14 45 Glucopyranosylidene-spiro-thiohydantoin ($K_i = 29.8 \mu M$ 46 against rat liver GP) was shown to exert considerable in vivo 47 blood sugar diminishing activity. 15

N-Acyl- β -D-glucopyranosylamines (compounds 1 in Chart 1) 49 were among the first GP inhibitors, ¹⁶ and many analogous 50 derivatives were investigated. ^{17–20} In this series, *N*-(2naphthoyl)- β -D-glucopyranosylamine (1 R = 2-naphthyl) was 51 the best inhibitor, 18 which also served as a lead structure for 52 bioisosteric replacements. As illustrated in Chart 1, enzymatic 53 tests²¹ as well as crystallographic studies¹⁹ revealed high 54 similarity of amide (1) and 1,2,3-triazole (2) type inhibitors 55 both in binding strength and structural features of the enzyme- 56 inhibitor complexes. Kinetic tests of bioisosteric oxadiazoles 22,23 57 3-5 demonstrated that the constitution of the heterocycle had 58 a strong bearing on the inhibition: the most efficient inhibitor 59 in these series was $5-(\beta-D-glucopyranosyl)-3-(2-naphthyl)-60$ 1,2,4-oxadiazole (5), which had a similar efficiency to that of 1. 61

Other investigations on C-glucopyranosyl heterocycles with 62 condensed rings showed that benzothiazole 7 was much less 63 efficient than benzimidazole 8.²⁴ An X-ray crystallographic 64 study of the RMGPb-8 complex revealed a specific H-bond 65 between NH of the heterocycle and the main chain C=O of 66 His 377, 25 and the stronger binding of 8 was attributed to this 67 interaction, which cannot exist in the case of 7.

On the basis of these preliminaries, synthesis and study of 69 1,2,4-triazoles of type 6 were envisaged anticipating that the H- 70 bond donor capacity of this heterocycle would result in 71 stronger inhibitors of GP.

3-Glycosyl-5-substituted-1,2,4-triazoles were described in the 73 literature mainly with furanoid rings in reactions of C- 74 glycofuranosyl (thio)formimidates with hydrazide or amidra-75 zone reagents^{26–28} or transforming a 2,5-anhydro-D,L-allono- 76 lactone derivative with aminoguanidine.²⁹ 3-Glycopyranosyl-5- 77 substituted-1,2,4-triazoles could not be located in the literature; 78 the only C-glycopyranosyl-1,2,4-triazoles were 1,3,5-trisubsti- 79 tuted derivatives obtained from glycosyl cyanides with 1-aza-2-80 azoniaallene salts³⁰ or with hydrazonoyl chlorides in the 81 presence of Yb(OTf)3.31

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Chart 1. Selected Inhibitors of Glycogen Phosphorylase and Their ${\rm Efficiency}^a$

 aK_i [μ M] against RMGPb for R=2-naphthyl. b A K_i value of 2.4 μ M was measured independently by Oikonomakos and co-workers. 22

Synthesis of the desired 3-glucopyranosyl-5-substituted-1,2,4-84 triazoles of type $\bf 6$ was planned by adaptation of a literature protocol³² in which acylation of N^1 -tosylamidrazones gave 3,5-86 disubstituted-1-tosyl-1,2,4-triazoles. Removal of the N-tosyl group was foreseen under conditions usually applied for N-88 desulfonylation of nitrogen heterocycles.³³

To start the syntheses, O-perbenzoylated β -D-glucopyranosyl 90 formimidate³⁴ 9 was reacted with tosylhydrazide to give the 91 necessary tosylamidrazone 10 in good yield (Scheme 1). 92 Reaction of 10 with acetyl chloride furnished tosyl-triazole 11a, 93 which was N-detosylated by tetrabutylammonium fluoride 94 (TBAF) to 12a. With acetoxyacetyl chloride 10 gave a mixture 95 of 11b and 12b indicating that the N-tosyl group is prone to 96 splitting off under the acylation conditions. The crude mixture 97 of 11b and 12b was treated with TBAF to produce 12b in 61% 98 yield for the two steps. Acylations of 10 with aromatic acid 99 chlorides were accompanied by complete N-detosylation 100 thereby simplifying the preparation of 12d-f, which were 101 obtained in good yields. Removal of the O-acyl protecting 102 groups was effected under Zemplén conditions to give test 103 compounds 6a and 6c-f in good to excellent yields.

Scheme 1. Synthesis of $3-(\beta-D-Glucopyranosyl)-5-substituted-1,2,4-triazoles (6)$

i) 1.5 equiv TsNHNH₂, dry CH₂Cl₂, rt; ii) 1.5 equiv RCOCl, 1.8 equiv pyridine, dry CHCl₃, 0 °C to rt; iii) TBAF, dry THF, reflux; iiv) ~1M NaOMe in MeOH. rt.

		Conditions and yields (%)					
	R		11		12ª		6
a	-CH ₃	ii	69	iii	88 ^b	iv	73
b	-CH ₂ OCOCH ₃	-	-	ii, iii	61°	-	-
c	-CH₂OH	-	-	-	-	iv	93^d
d	-C ₆ H ₅	-	-	ii	69	iv	62
e	-C ₆ H ₄ -4- <i>t</i> Bu	-	-	ii	58	iν	71
f	2-naphthyl	-	-	ii	56	iv	81

3-(β -D-Glucopyranosyl)-5-substituted-1,2,4-triazoles **6** were 104 assayed against RMGPb as described earlier, ³⁵ and the kinetic 105 results, showing the compounds to be competitive inhibitors, 106 are summarized in Table 1. Methyl (**6a**) and hydroxymethyl 107 t1 (**6c**) derivatives proved weak inhibitors in the micromolar 108 range and were significantly less efficient than the parent 109 amides **1a** and **1c**, respectively. Appending unsubstituted 110 aromatic groups to the 1,2,4-triazole ring as in **6d** and **6f** led 111

Table 1. Inhibition^a of RMGPb by Compounds 6 and Comparison to Other Nonclassical Bioisosteres

 ${}^{a}K_{i}$ [μ M] b Calculated from the IC₅₀ value by using a web-based tool. ³⁶

112 to a remarkable strengthening of the inhibition. While 1,2,4-113 oxadiazoles $\mathbf{5d}$ and $\mathbf{5f}$ were practically equipotent with the 114 corresponding amides $\mathbf{1d}$ and $\mathbf{1f}$, triazoles $\mathbf{6d}$ and $\mathbf{6f}$ inhibited 115 the enzyme by ~ 1 order of magnitude stronger, respectively. 116 This indicated that the possibility for the formation of a H-117 bond was advantageous for the binding, rendering compound 118 $\mathbf{6f}$ to one of the most efficient glucose analogue inhibitors of 119 GP known to date. Introduction of a t-butyl substituent in the 120 4-position of the phenyl group as in $\mathbf{6e}$ resulted in a much 121 weaker inhibitor. This observation may reveal that the active 122 site of GP, where these compounds may bind to, can not 123 accommodate a bulky aliphatic moiety.

Further studies to establish the binding peculiarities of these inhibitors by X-ray crystallographic investigation of the enzyme—inhibitor complexes as well as molecular dockings to predict other efficient derivatives based on this skeleton are in progress.

In conclusion, a new method was elaborated for the synthesis of hitherto unknown 3-(β -D-glucopyranosyl)-5-substituted-131 1,2,4-triazoles. These compounds inhibited rabbit muscle GPb, and the 5-(2-naphthyl) derivative with its submicromolar inhibition proved one of the best inhibitors of the enzyme.

134 ASSOCIATED CONTENT

135 Supporting Information

136 Representative synthetic procedures, enzyme kinetic measure-137 ments, and compound characterization. This material is 138 available free of charge via the Internet at http://pubs.acs.org.

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151 **Notes**

152 The authors declare no competing financial interest.

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