Evaluation of Substituted *N*,*N*'-Diarylsulfonamides as Activators of the Tumor Cell Specific M2 Isoform of Pyruvate Kinase

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The metabolism of cancer cells is altered to support rapid proliferation. Pharmacological activators of a tumor cell specific pyruvate kinase isozyme (PKM2) may be an approach for altering the classic Warburg effect characteristic of aberrant metabolism in cancer cells yielding a novel antiproliferation strategy. In this manuscript, we detail the discovery of a series of substituted N,N'-diarylsulfonamides as activators of PKM2. The synthesis of numerous analogues and the evaluation of structure–activity relationships are presented as well as assessments of mechanism and selectivity. Several agents are found that have good potencies and appropriate solubility for use as chemical probes of PKM2 including **55** (AC₅₀ = 43 nM, maximum response = 84%; solubility = 7.3 μ g/mL), **56** (AC₅₀ = 99 nM, maximum response = 84%; solubility = 5.7 μ g/mL), and **58** (AC₅₀ = 38 nM, maximum response = 82%; solubility = 51.2 μ g/mL). The small molecules described here represent first-in-class activators of PKM2

Introduction

Altered metabolism was among the earliest observed differences between cancer and healthy tissues. Normally, the cellular process of glycolysis converts glucose to pyruvate, which is subsequently converted to acetyl-CoA for entry into the Krebs cycle and ATP^{*a*} production. Under anaerobic conditions, pyruvate is converted to lactate and ATP generation is limited (Figure 1A). In seminal work, Otto Warburg reported that cancer cells have increased rates of lactate production even in the presence of oxygen (Figure 1B).^{1,2} This altered metabolism is thought to give tumor cells a selective growth advantage relative to normal cells.^{3,4} Overall, this change in metabolism in cancer cells is referred to as the Warburg effect and allows cells to meet the metabolic requirements of cell proliferation.

Pyruvate kinase operates at the final step of glycolysis, catalyzing the transfer of a phosphate group from phosphoenolpyruvate (PEP) to ADP, yielding one molecule of ATP and one molecule of pyruvate. Humans have two pyruvate kinase (PK) genes that each produce two different isozymes due to alternative splicing.^{5–7} The *Pyk* L/R gene produces two mRNAs with defined variations in the first exon, which are expressed in selective tissues including the liver (PKL form) and red blood cells (PKR form).⁸ The *Pyk* M gene also produces two mRNAs that leads to the expression of the M1 isozyme (PKM1), which is found in most normal adult tissues. However, in fetal tissue, alternative splicing results in the M2 isozyme (PKM2), with the splicing change occurring at a site required for the binding and allosteric activation of PKM2 by fructose-1,6-bis-phosphate (FBP).^{9–11} An early genetic observation associated with tumorigenesis recognized that normal adult PK isozymes are replaced by PKM2 in tumor cells.^{12–14} All tumors and cancer cell lines studied to date show exclusive expression of the PKM2 isoform.^{15,16} PKM2 has also been found in some normal adult tissues that require high rates of nucleic acid synthesis.

PKM2 exists in equilibrium between a low activity T-state and a high activity R-state. The active forms of PKM2, PKL, and PKR are tetrameric and are allosterically activated by FBP through binding at the aforementioned flexible loop region near the dimer-dimer interface of the tetramer. The allosteric regulation of PK isoforms by FBP provides an important feedback mechanism that activates PKM2 when this early stage glycolytic intermediate accumulates to sufficient levels. It has been proposed that the low activity T-state may represent a dimer form of PKM2, which shows weak affinity for PEP while formation of the R-state involves formation of the tetrameric form exhibiting increased affinity for this substrate. In tumor cells, PKM2 has been reported to exist primarily in its low activity, dimeric form.¹⁶ The binding of FBP to the flexible loop region of PKM2 is thought to

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^{*a*} Abbreviations: PK, pyruvate kinase; ATP, adenosine triphosphate; ADP, adenosine diphosphate; PEP, phosphoenolpyruvate; FBP, fructose-1,6-bis-phosphate; qHTS, quantitative high-throughput screen; LDH, lactate dehydrogenase.



Figure 1. (A) The fate of glucose metabolism to pyruvate in anaerobic and aerobic environments in differentiated, healthy tissue. (B) The fate of glucose metabolism to pyruvate in rapidly proliferating tissue and tumors. PKM2 is activated by FBP and inhibited by tyrosine-phosphorylated proteins derived from growth factor signaling.

stabilize the tetrameric form of the enzyme leading to activation. In contrast, PKM1 has a high affinity for PEP in its native state and is not activated by FBP.¹⁰ Functionally, the down-regulation of PKM2 activity in cancer cells is thought to aid in shunting key glycolytic intermediates toward pathways where they, in turn, are utilized as precursors for lipid, amino acid, and nucleic acid synthesis. Rapidly proliferating cells such as undifferentiated embryonic tissue and cancer cells require increased amounts of these fundamental cellular building blocks. Thus, the down-regulation of PKM2 activity provides a purposeful divergence from catabolic metabolism aimed at energy production toward an anabolic state aimed at providing the needed resources for rapid cellular construction.

Recent work has shown that replacement of PKM2 with PKM1 in cancer cells can relieve the Warburg effect and promote oxidative cellular metabolism characteristic of differentiated cells.¹⁷ Further, an assessment of growth in vivo of human lung cancer cells engineered to express only PKM1 showed delayed tumor formation in nude mouse xenografts. The tumors that did form in this model were shown to have re-expressed PKM2. In a related study, it was demonstrated that PKM2 has high affinity for binding phosphotyrosine peptides.¹⁸ Protein tyrosine phosphorylation is increased in tumor cells as a consequence of oncogenic intracellular signaling events that also drive uptake of excess nutrients. Binding of PKM2 to tyrosine phosphorylated proteins catalyzes the release of FBP from the enzyme and further down-regulates PKM2 activity to promote entry of glycolytic intermediates into biosynthetic pathways. Taken as a whole, these data suggest a therapeutic strategy whereby activation of PKM2 may restore normal cellular metabolism to a state characteristic of normal differentiated cells. We describe here the development of a PKM2 assay capable of identifying inhibitors and activators, the quantitative high-throughput screening (qHTS) of a large chemical library, and the identification and exploration of the structure activity relationships (SAR) of substituted N,N'-diarylsulfonamides as activators of PKM2.

Results

Pyruvate Kinase–Luciferase Coupled Assay. For the development of an assay for both activators and inhibitors of

PKM2, we took advantage of a well utilized luminescent assay detection system for protein kinases.^{19,20} Briefly, these assays use the ATP-dependent reaction catalyzed by firefly luciferase to measure ATP depletion and have been widely applied protein kinase assays. In the present assay, we applied this technology to measure ATP production catalyzed by PKM2. This provides for a robust increase in luminescent signal upon ATP product formation (see Supporting Information Table S1).²¹ The primary high-throughput screen was performed in 1536-well microtiter plates using $4 \mu L/well$ assay volume. Using this assay design, a quantitative high-throughput screen (qHTS)²² of nearly 300000 small molecules of the NIH Molecular Libraries²³ Small Molecule Repository was performed.

The results of this screening effort are detailed in Figure 2. Small molecule activators of PKM2 were more prevalent than inhibitors, occurring at 0.27% for high confidence 1a, 1b, and 2c concentration-response curves (CRCs; class 1 curves display two asymptotes, an inflection point, and $r^2 \ge 0.9$; subclasses 1a and 1b are differentiated by full (>80%) vs partial ($\leq 80\%$) response, respectively, while class 2a curves display a single left-hand asymptote and inflection point and full (>80%) response; see Supporting Information for complete curve class definitions).^{22,24} A cheminformatics analysis of the qHTS data included chemotype clustering, singleton identification, analysis of orthogonal activities, and structural considerations, which included physical properties and anticipated optimization potential. Potency range and maximum response was additionally considered (a visual representation is shown in Figure 2C). Ultimately, this analysis led us to focus on two lead structures represented by the substituted thieno-[3,2-b]pyrrole[3,2-d]pyridazinone 1 and substituted N,N'-diarylsulfonamide 2 shown in Figure 2D (a separate manuscript detailing the evaluation of **1** is in preparation).

Upon resynthesis, the PKM2 activity of 2 was confirmed in the aforementioned PK coupled luciferase assay that monitored ATP production and a fluorescent PK-lactate dehydrogenase coupled assay that monitored pyruvate production. Good agreement between the two assays was observed. For PKM2, the AC_{50} in the luciferase-coupled assay was approximately 4-fold more potent than the AC_{50} determined in the LDH coupled assay but the efficacy of the response was approximately 1.5- to 2-fold higher in the LDH coupled assay. The LDH assay was performed at high saturating ADP concentrations and low (0.1 mM) PEP concentrations, so differences that did occur between the two assays likely reflect preferences for the ADP-bound form of the enzyme. We discuss the data here using the luciferasecoupled assay but point to the LDH data for specific examples. The entire data set for both assays are available in PubChem (see Methods for assay identifiers (AIDs)). We confirmed that 2 and all reported analogues did not activate firefly luciferase (PubChem AID: 1379) nor showed fluorescence at the wavelength used for NADH detection (data not shown). As well, the activators identified here were inactive in a qHTS of the Leishmania mexicana pyruvate kinase (PubChem AIDs: 1721, 1722) that used the identical format as the PKM2 qHTS. Therefore, 2 appears to be a genuine activator of PKM2.

Mode of Action. We examined the mode of action **2** through analysis of the steady-state kinetics of PEP and ADP by the PKM2.¹¹ As discussed, FBP is known to allosterically activate PKM2 through induction of an enzyme state with a high affinity for PEP. In the absence of

A	Activity	Distribution			Curve	e Classific	ation		
	Activity	Distribution	1.1	1.2	2.1	2.2	3	4	5
	Activation	No. Cmpds.	134	318	325	1541	2094		
	Adivation	% library	0.05	0.11	0.11	0.55	0.74	276309	1 [
	Inhibition	No. Cmpds.	4	25	31	582	983	97.73%	0.00%
	minibition	% lib ra ry	0.00	0.01	0.01	0.21	0.35		[
2000 1500 1001 5- 2- 901- -10-	Log(Cmpd), M	300,000 150,000 4 5ample #	750 Samole #	0.8	-6 Log(Cmpd), M	300 200 Activity 100 (%) 4	eo-{		

Figure 2. (A) Summary of actives from PKM2 qHTS of 276309 small molecules. (B) Concentration response curve (CRC) representation of the full qHTS data set. (C) CRC's for actives from the primary screen associated with curve classes 1a, 1b, and 2a showing potency and maximum response. (D) Chemical structures of the two lead activators of PKM2 substituted thieno[3,2-*b*]pyrrole[3,2-*d*]pyridazinone NCGC00031955 (1) and substituted *N*,*N*'-diarylsulfonamide NCGC00030335 (2).



Figure 3. (A) Initial velocity as a function of PEP and ADP concentration in the presence (\triangle) or absence (\bigcirc) of **2** (10 μ M). (B) Initial velocity as a function PEP and ADP concentration in the presence (\bigcirc) or absence (\bigcirc) of FBP (10 μ M). V_{o} , initial rate in pmol/min as determined in the PK-LDH coupled assay (kinetic assays were carried out at approximately 22 °C with 10 nM PKM2 using [KCl] = 200 mM, [MgCl₂] = 15 mM, and with either [ADP] or [PEP] = 4.0 mM; see Supporting Information).

activator, hPK shows low affinity for PEP ($K_{\rm M} \sim 1.5$ mM). In the presence of **2** or FBP, the $K_{\rm M}$ for PEP decreased nearly 10-fold to 0.26 ± 0.08 mM or 0.1 ± 0.02 mM, respectively, with lesser effects on $V_{\rm max}$ (values of 245 pmol/min with or without FBP and 265 pmol/min with **2**). The addition of excess PEP abolishes the activation of PKM2 by **2** or FBP. However, varying the concentration of ADP in the presence and absence of **2** shows that the steady-state kinetics for this substrate are not significantly affected ($K_{\rm M}$ for ADP = 0.1 mM in either condition). Thus, our primary lead **2** lowered the $K_{\rm M}$ for PEP but had no affect on the $K_{\rm M}$ for ADP (Figure 3A) similar to what we observed for FBP (Figure 3B).

Synthesis of Substituted N,N'-Diarylsulfonamides and Selected Analogues. The synthetic strategy utilized to build various N,N'-diarylsulfonamide analogues was a straightforward sequence of coupling reaction, deprotection, and a second coupling reaction as detailed in Scheme 1.25 Specifically, monoboc protected piperazine was coupled to numerous aryl sulfonyl chlorides to provide the corresponding boc-protected N-arylsulfonamides. These intermediates were deprotected and subsequently coupled to a second aryl sulfonyl chloride to provide numerous N, N'-diarylsulfonamide analogues. All final compounds were purified by preparative scale HPLC, and the yields for these procedures were typically high. The same procedure was utilized to explore alternate diamines between each aryl-sulfonamide moiety including cyclic diamines of different ring size (analogue 34), linear diamines (analogues 35-39), ring systems with an internal secondary amine and an exocyclic amine (analogues 40-47), and analogues with variously substituted piperazines (analogues 48-51) (scheme not shown). A similar procedure was utilized for the synthesis of several amide versions of these reagents (represented by compounds 22 and 33) in order to explore a comparative SAR of sulfonamide versus amide functionality (scheme not shown). We were also interested in exploring several of the related sulfone derivatives akin to the lead structure (Scheme 2). To synthesize these derivatives, we treated N-boc-4-bromopiperidine with various aryl sulfides to afford the appropriately substituted thiol ethers. Oxidation to the sulfone followed by boc deprotection and a second coupling to various aryl sulfonyl chlorides to provided 4-(arylsulfonyl)-1-(arylsulfonyl)piperidine analogues (represented by analogues 21 and 32). Finally, it was of interest to explore N, N'-diarylsulfonamide analogues on the related piperazin-2-one core (Scheme 3). These derivatives were accessed through treatment of piperazin-2-one with 1 equiv of various aryl sulfonyl chlorides which preferentially coupled to the free amine moiety. The resulting intermediate was converted to the 1,4-bis(arylsulfonyl)piperazin-2-ones by deprotonation of the amide with LHMDS, followed by addition of various aryl sulfonyl chlorides to generate the desired products in good yields (represented by analogues 52 and 53).

SAR of Substituted *N*,*N'*-Diarylsulfonamides and Selected Analogues. The lead *N*,*N'*-diarylsulfonamide **2** identified from the primary screen was found to possess an AC₅₀ value versus PKM2 of 0.111 μ M and maximum responses versus PKM2 (relative to activation by FBP) of 92%, respectively (Table 1). In the LDH assay, which used high saturating ADP levels and low (0.1 mM) levels of PEP, we found greater average efficacy but lower potency for **2** (AC₅₀ values of 0.75 μ M and a maximum responses of 150%). We focused

Scheme 1^a



^{*a*} Conditions and reagents: (i) TEA, CH₂Cl₂, 0 °C; (ii) TFA, CH₂Cl₂, 0 °C; (iii) TEA, CH₂Cl₂, 0 °C.

Scheme 2^{*a*}



^{*a*} Conditions and reagents: (i) K₂CO₃, DMF; (ii) mCPBA, CH₂Cl₂, 0 °C; (iii) TFA, CH₂Cl₂, 0 °C; (iv) TEA, CH₂Cl₂, 0 °C.

Scheme 3^a

our early efforts on both improving these values and further understanding of each chemotypes SAR potential and limitation. One initial focus involved symmetric versions of the N,N'-diarylsulfonamides. As such, symmetry was examined utilizing the 6-(2,3-dihydrobenzo[b][1,4]dioxine) heterocycle (analogue 3) and the 4-methoxybenzene ring (analogue 4). Each analogue had slightly diminished AC₅₀ values (270 and 171 nM, respectively). From here, one aryl sulfonamide unit was held constant while exploring the SAR of the other aryl sulfonamide. Compounds 5-20 are representative examples from this strategy whereby the 6-(2,3-dihydrobenzo[b]-[1,4]dioxine) heterocycle remained constant. The most effective substitutions involved electron withdrawing groups in the 2- and 6-positions of the phenyl ring [for instance 2,6difluorobenzene (analogue 10, $AC_{50} = 65$ nM, maximum response = 94%), 2,6-difluoro-4-methoxybenzene (analogue 12, $AC_{50} = 28$ nM, maximum response = 92%), and 2,6difluoro-3-phenol (analogue 14, $AC_{50} = 52$ nM, maximum response = 95%]. Given the improved activity observed with the difluoro analogues, we next chose to hold the 2,6-difluorobenzene ring constant and vary the opposite side with both substituted phenyl rings and various heterocycles (compounds 23-31). Several heterocycles were tolerated, including the 7-(3,4-dihydro-2H-benzo[b][1,4]dioxepine) moiety (analogue 25, $AC_{50} = 103$ nM, maximum response = 100%) and the 6-(2-methylbenzo[d]thiazole) moiety (analogue **31**, $AC_{50} = 86$ nM, maximum response = 104%). The 2-napthyl and 6-(2,2-dimethylchroman) derivatives provided significant enhancement in terms of maximum response (analogues 28 and 29, $AC_{50} = 66$ and 93 nM, maximum response = 138% and 119%, respectively). All amide versions of these compounds were found to be inactive in both assays (represented by derivatives 22 and 33). The sulfone derivatives that were explored showed loss of potency. Interestingly, this loss was more severe when the sulfone moiety bridged the piperidine ring system to the 6-(2,3-dihydrobenzo[b]-[1,4]dioxine) system relative to substituted phenyl rings (i.e., 2,6-difluorophenyl) as illustrated by analogues 21 and $32(AC_{50} = 254 \text{ and } 863 \text{ nM}, \text{maximum response} = 104\% \text{ and}$ 110%, respectively).

Several linear diamines and several alternate diamine core ring systems were examined. For these studies, we retained the 2,6-difluorophenyl and 6-(2,3-dihydrobenzo[*b*][1,4]dioxine) heterocycle as the two aryl substituents to afford comparative uniformity. The results are shown in Table 2 and demonstrate that the piperazine and the related 1,4-diazepane (analogue **34**, AC₅₀ = 895 nM, maximum response = 120%) had clear advantages over other diamine moieties. Ligations with linear diamines ranging from 2 to 6 carbons in length (analogues **35–39**) were found to have diminished potencies. Cis and trans versions of the cyclohexane-1,4-diamines conferred similar loss in activation potency. Interestingly, the trans version of this analogue performed significantly better than the cis version. Numerous analogues with one secondary amine contained in 4-, 5- and 6-membered rings and one



^a Conditions and reagents: (i) TEA, CH₂Cl₂, 0 °C; (ii) LHMDS, THF, -78 °C - then Ar₂SO₂Cl.

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						hPK, M2	hPK, M2
	#	Х	Y	Ar ₁	Ar ₂	AC ₅₀ (μM) ^a	Max. Res. ^b
	2	NA	NA	6-(2,3-dihydrobenzo[b][1,4]dioxine)	4-methoxybenzene	0.111	92
Ar₁−S−N N−S−Ar₂	3	NA	NA	6-(2,3-dihydrobenzo[b][1,4]dioxine) 6-	-(2,3-dihydrobenzo[b][1,4]dioxine) 0.270	90
	4	NA	NA	4-methoxybenzene	4-methoxybenzene	0.171	88
2.4	5	SO_2	Ν	4-cyanobenzene	NA	0.029	44
2-4	6	SO_2	Ν	4-chlorobenzene	NA	0.154	100
	7	SO_2	Ν	4-fluorobenzene	NA	0.094	100
Ar ₁ -X-Y_N-SO	8	SO_2	Ν	3-fluorobenzene	NA	0.316	107
0 \	9	SO_2	Ν	2-fluorobenzene	NA	0.089	114
5-22	10	SO_2	Ν	2,6-difluorobenzene	NA	0.065	94
Ę,	11	SO_2	Ν	2,4,5-trifluorobenzene	NA	0.090	105
	12	SO_2	Ν	2,6-difluoro-4-methoxybenzene	NA	0.028	92
	13	SO_2	Ν	2,5-difluoro-4-propylbenzene	NA	0.757	69
5 F	14	SO_2	Ν	2,6-difluoro-3-phenol	NA	0.052	95
20-00	15	SO_{2}	Ν	2,4-difluorobenzene	NA	0.124	113
	16	SO_2	Ν	phenyl	NA	0.202	108
	17	SO_2	Ν	3-(trifluoromethyl)benzene	NA	0.209	39
	18	SO_2	Ν	3-methoxybenzene	NA	0.113	90
	19	SO_2	Ν	2-pyridine	NA	0.542	103
	20	SO_2	Ν	2-pyridine 1-oxide	NA	> 10	82
	21	$\overline{SO_2}$	CH	2,6-difluorobenzene	NA	0.254	104
	22	CO	Ν	2,6-difluorobenzene	NA	inactive	NA
	23	SO_2	Ν	4-methoxybenzene	NA	0.090	102
	24	SO_2	Ν	2,6-difluorobenzene	NA	0.066	74
	25	SO_2	Ν	7-(3,4-dihydro-2H-benzo[b][1,4]dioxepine) NA	0.103	100
	26	SO_2	Ν	5-(benzo[d][1,3]dioxole)	NA	0.191	61
	27	SO_2	Ν	7-(4-methyl-3,4-dihydro-2 <i>H</i> -pyrido[3,2- <i>b</i>][1,4]02	xazine) NA	2.71	94
	28	SO_2	Ν	2-naphthalene	NA	0.066	138
	29	SO_2	Ν	6-(2,2-dimethylchroman)	NA	0.093	119
	30	SO_2	Ν	5-(1-methyl-1 <i>H</i> -indole)	NA	0.387	91
	31	SO_2	Ν	6-(2-methylbenzo[d]thiazole)	NA	0.086	104
	32	SO_2	СН	6-(2,3-dihydrobenzo[b][1,4]dioxine)	NA	0.863	110
	33	CO	Ν	6-(2,3-dihydrobenzo[b][1,4]dioxine)	NA	inactive	NA

 a AC₅₀ values were determined utilizing the luminescent pyruvate kinase-luciferase coupled assay (ref 22) and the data represents the results from three separate experiments. See the Supporting Information section for comparative values from the fluorescent pyruvate kinase–lactate dehydrogenase coupled secondary assay (ref 27). Maximum response (Max Res) is % activity at 57 μ M of compound. See Methods for normalization. See Supporting Information section for standard deviations.

Table 2. SAK of Sciected <i>N</i> . <i>N</i> - Dai visuitonamides with Divergent Diamine wor	Table 2.	SAR of Selec	ted N.N'-Dar	vlsulfonamides	with Divergent	Diamine Moietie
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	#	COPE	hPK, M2	hPK, M2 Mar. Paulo
	#	CORE	$4C_{50} (\mu M)^{\circ}$	Max. Res.
C C	10	piperazine	0.065	94
	34	1,4-diazepane	0.866	120
	35	N,N'-(ethane-1,2-diyl)	> 15	60
∽o´ F	36	N,N'-(propane-1,3-diyl)	3.85	106
10, 34-47	37	N,N'-(butane-1,4-diyl)	7.97	113
	38	N,N'-(pentane-1,5-diyl)	2.33	114
	39	N,N'-(hexane-1,6-diyl)	4.83	110
	40	N,N'-((trans)-cyclohexane-1,4-diy	d) 2.11	91
	41	N,N'-((cis)-cyclohexane-1,4-diyl) > 35	61
	42	HN-_N	9.00	100
	43	N	3.69	101
	44	HN	> 10	84
	45	N	> 15	82
	46		3.05	108
	47	NJNH	4.47	93

 a AC₅₀ values were determined utilizing the luminescent pyruvate kinase–luciferase coupled assay (ref 22), and the data represents the results from three separate experiments. See the Supporting Information for comparative values from the fluorescent pyruvate kinase–lactate dehydrogenase coupled secondary assay (ref 27). Maximum response (Max Res) is % activity at 57 μ M of compound. See Methods for normalization. See Supporting Information for standard deviations.

exocyclic primary amine were examined (analogues 42-47) and found to be less active than the original lead compounds

in both assays. In addition to the derivatives shown in Table 2, numerous bicyclic and spirocyclic diamines were examined

Table 3. SAR of Selected N,N'-Diarylsulfonamides with Substitutions on the Piperazine Ring

	#		р	C	D	v	v	hPK, M2	hPK, M2
	#	A	В	U	D	Λ	Y	$AC_{50}(\mu M)^{*}$	Max. Kes.
	10	Н	Н	Н	Н	NA	NA	0.065	94
P→N-S→	48	Me	Η	Н	Η	NA	NA	4.34	109
	49	Н	Me	Н	Н	NA	NA	3.10	99
10, 48-51 F	50	Н	Η	Me	Η	NA	NA	9.18	108
	51	Н	Н	Η	Me	NA	NA	2.93	108
	52	NA	NA	NA	NA	0	NA	0.114	105
∽o x y f	53	NA	NA	NA	NA	NA	0	2.42	97
52, 53									

 a AC₅₀ values were determined utilizing the luminescent pyruvate kinase–luciferase coupled assay (ref 22), and the data represents the results from three separate experiments. See the Supporting Information for comparative values from the fluorescent pyruvate kinase–lactate dehydrogenase coupled secondary assay (ref 27). Maximum response (Max Res) is % activity at 57 μ M of compound. See Methods for normalization. See Supporting Information for standard deviations.

Table 4. S	SAR of Se	elected N,N	V'-Diarylsulf	onamides [Including	Solubility	Assessment
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	#	n	Ar ₁	Ar ₂	hPK, M2 AC ₅₀ (µM) ^a	hPK, M2 Max. Res. ^b	Solubility ^c (µM)	Solubility (µg/mL)
	10	1	2,6-difluorobenzene	6-(2,3-dihydrobenzo[b][1,4]dioxine)	0.065	94	< 1.1	< 0.5
$Ar_1 = \ddot{S} = \dot{N}$ $N = \ddot{S} = Ar_2$	34	2	2,6-difluorobenzene	6-(2,3-dihydrobenzo[b][1,4]dioxine)	0.866	120	5.6	2.7
ö (7 ö	54	1	3-aniline	2,6-difluoro-4-methoxybenzene	0.026	90	< 0.7	< 0.4
10, 34, 54-59	55	1	3-aniline	6-(2,3-dihydrobenzo[b][1,4]dioxine)	0.043	84	7.3	4.1
	56	1	3-aniline	2,6-difluorobenzene	0.099	84	5.7	3.0
	57	2	3-aniline	2,6-difluoro-4-methoxybenzene	0.223	92	26.3	15.1
	58	2	3-aniline	6-(2,3-dihydrobenzo[b][1,4]dioxine)	0.038	82	51.2	29.0

 a AC₅₀ values were determined utilizing the luminescent pyruvate kinase–luciferase coupled assay (ref 22), and the data represents the results from three separate experiments. See the Supporting Information for comparative values from the fluorescent pyruvate kinase–lactate dehydrogenase coupled secondary assay (ref 27). b Max Res value represents the % activation at 57 μ M of compound. c Kinetic solubility analysis was performed by Analiza Inc. and are based upon quantitative nitrogen detection as described (www.analiza.com). The data represents results from three separate experiments with an average intraassay %CV of 4.5%. d LogD analysis was performed by Analiza Inc. and are based upon octanol/buffer partitioning and quantitative nitrogen detection of sample content as described (www.analiza.com). NT = not tested.

(for instance 2,6-diazabicyclo[3.2.2]nonane and 2,7-diazaspiro[4.4]nonane) and found to be less active than the corresponding piperazine and 1,4-diazepane analogues (data not shown).

Several piperazine rings were synthesized and evaluated with a single methyl addition proximal to either the 2,6difluorophenyl or 6-(2,3-dihydrobenzo[b][1,4]dioxine) heterocycle. Additional consideration was given to the absolute stereochemistry of the methyl group. The results are detailed in Table 3 and show that these analogues were less potent than the unmodified ring systems. Another piperazine ring modification was the incorporation of a carbonyl moiety α to the ring nitrogens. Here, the amine to lactam conversion proximal to the 6-(2,3-dihydrobenzo[b][1,4]dioxine) heterocycle resulted in an active derivative (analogue 52, $AC_{50} =$ 114 nM, maximum response = 105%). The same amine to lactam conversion adjacent to the 2,6-difluorobenzene resulted in a loss of potency (analogue 53, $AC_{50} = 2.42 \,\mu M$, maximum response = 97%). The activities of these agents again demonstrate the lack of symmetric SAR for this chemotype.

An important consideration was the aqueous solubility of the analogues presented in this study. To assess the solubility of selected analogues, including **10** and **34**, we utilized a commercial provider of aqueous solubility profiles.²⁶ The majority of analogues examined had solubility levels (measured in both μ g/mL and μ M) below the detectable limit. The solubility of the seven-member **34** registered at the low values of 5.6 μ g/mL (Table 4). The lack of aqueous solubility is, therefore, a major liability for these agents as molecular probes of PKM2. To rectify this, we designed and synthesized numerous analogues incorporating solubilizing functionalities on one or both aryl groups of selected compounds (including pyridines, anilines, and *N*-acetyl anilines). Table 4 presents selected results. In general, aniline substitutions placing the amine meta relative to the sulfonamide functionality resulted in highly active agents including **55** (AC₅₀ = 43 nM, maximum response = 84%), **56** (AC₅₀ = 99 nM, maximum response = 84%), and **58** (AC₅₀ = 38 nM, maximum response = 82%). Gratifyingly, selected analogues incorporating this moiety also had amplified water solubilities including **57** (26.3 μ g/mL) and **58** (51.2 μ g/mL) (Table 4). Overall, **58** possesses the best combination of activity and solubility of all the analogues presented in this study.

Selectivity of Chosen N,N'-Diarylsulfonamide Analogues. With a better understanding of the SAR for this chemotype, we next concerned ourselves with the selective activation of PKM2 versus PKM1, PKR, and PKL. An appropriate tool compound aimed at further delineating the role of PKM2 as a critical contributor in the Warburg effect requires a high degree of selective activation of PKM2 relative to other PK targets with particular consideration for PKM1. Members of each chemotype were assayed versus PKM1, PKR, and PKL (the assay design was similar as that described for PKM2, and details are located in the Methods and Supporting Information). All analogues in the N, N'-diarylsulfonamides class were found to be inactive versus PKM1. This is consistent with the lack of allosteric regulation for the PKM1 isoform. The activity versus PKR and PKL varied from compound to compound but generally showed maximum responses below 30% for both isozymes in both assay formats. The selectivity for 2 is shown in Figure 4.



Figure 4. Selectivity assessment for 2 versus PKM2 (○), PKM1 (■), PKL (□), and PKR (●).

Conclusion

Warburg's finding that cancer cells show altered cellular respiration and metabolism ranks as one of the earliest observations in cancer biology. A key realization associated with the Warburg effect is the re-expression of PKM2 in all cancer cells leading to increased availability of glycolyctic intermediates for biosynthesis of amino acids, nucleic acids, and lipid building blocks for cellular construction. The expression of PKM2 in tumor cells makes this enzyme an attractive target for therapeutic intervention in cancer. We have described here the discovery and optimization of a series of substituted N,N'-diarylsulfonamide activators of PKM2, which should be useful pharmacological tools to study the antiproliferative effects of PKM2 activation. This chemotype activates the PKM2 isoform by increasing the affinity for phosphoenolpyruvate (PEP) with little effect on kinetics of ADP binding similar to the natural activator FBP. The synthesis of the lead structure and numerous analogues allowed the identification of compounds with low nanomolar potencies and activation responses similar to full activation afforded by the native allosteric activator FBP. In addition, unlike FBP, these agents were found to be selective for PKM2 versus PKM1, PKL, and PKR isoforms. SAR explorations included both evaluations of PKM2 activation and aqueous solubility. From these studies, analogues 10 (AC₅₀ = 65 nM), **12** (AC₅₀ = 28 nM), **14** (AC₅₀ = 52 nM), **28** (AC₅₀ = 66 nM), **54** (AC₅₀ = 26 nM), **55** (AC₅₀ = 43 nM), **56** (AC₅₀ = 99 nM), and 58 (AC₅₀ = 38 nM) were identified as highly potent activators of PKM2. NCGC00185916-01 (58) was identified as a small molecule PKM2 activator that combined good potency as a PKM2 activator with appropriate aqueous solubility (51.2 μ g/mL). The PKM2 activators described here represent first-in-class small molecule activators of PKM2 and studies to examine these compounds' antiproliferative capacity and their ability to alleviate the Warburg effect are currently underway.

Methods

Reagents. Kinase-Glo was obtained from Promega (Madison, Wi). ATP, PEP, LDH, and NADH were from Sigma. Reagents and solvents were purchased from Sigma, Alfa Aesar, Acros, Enamine, Oakwood Products, Matrix Scientific, or Chem-Impex International. Pyruvate kinase (EC 2.7.1.40), human M2 isoform, was expressed as a N-terminal His₆-tagged fusion protein in BL21(DE3)pLys cells. Preparation and purification was as described in Christofk et al.¹⁸

Luminescent Pyruvate Kinase-Luciferase Coupled Assay. Production of a luminescent signal based on the generation of ATP by pyruvate kinase was determined by using the ATPdependent enzyme firefly luciferase.²¹ Three μ L of substrate mix (at rt) in assay buffer (50 mM imidazole pH 7.2, 50 mM KCl, 7 mM MgCl₂, 0.01% Tween 20, 0.05% BSA) was dispensed into Kalypsys white solid bottom 1536-well microtiter plates using a bottle-valve solenoid-based dispenser (Kalypsys). The final concentrations of substrates in the assay were 0.1 mM ADP and 0.5 mM PEP. Then 23 nL of compound were delivered with a 1536-pin array tool²⁸ and 1 μ L of enzyme mix in assay buffer (final concentration, 0.1 nM pyruvate kinase, 50 mM imidazole pH 7.2, 0.05% BSA, 4 °C) was added. Microtiter plates were incubated at rt for 1 h and 2 μ L of luciferase detection mix (Kinase-Glo, Promega²¹ at 4 °C protected from light) was added and luminescence was read with a ViewLux (Perkin-Elmer) using a 5 s exposure/plate. Data was normalized for AC₅₀ values to control columns containing uninhibited enzyme (n), and AC_{100} inhibition (i) according the following equation: activation $(\%) = [(c - n)/(n - i)] \times 100$ where c = compound, n = DMSO neutral, i = no enzyme control. A % activity of 100% is approximately a 2-fold increase over basal assay signal (% activation by FBP was variable but averaged 100%). Monitoring of activation was accomplished using enzyme at $3 \times$ the final concentration. The primary qHTS data and confirmatory data are available in PubChem (AIDs: 1631, 1634, and 1751). Follow-up of synthesized analogues was determined using the same protocol with the exception that the enzyme concentrations for isoforms PKM1, L and R were 1, 0.1, and 0.1 nM, respectively (PubChem AIDs for M1, L and R bioluminescent assays are 1780, 1781, and 1782, respectively).

Fluorescent Pyruvate Kinase-Lactate Dehydrogenase Coupled Secondary Assay. All compounds were also tested in a kinetic mode by coupling the generation of pyruvate by pyruvate kinase to the depletion of NADH through lactate dehydrogenase.²⁷ For PKM2, 3 µL of substrate mix (final concentration, 50 mM Tris-Cl pH 8.0, 200 mM KCl, 15 mM MgCl₂, 0.1 mM PEP, 4.0 mM ADP, and 0.2 mM NADH) was dispensed into Kalypsys blacksolid 1536-well plates using the Aurora Discovery BioRAPTR flying reagent dispenser (FRD; Beckton-Dickenson, Franklin Lakes, NJ)²⁸ and 23 nL of compounds were delivered using a Kalypsys pin tool²⁹ and then 1 μ L of enzyme mix (final concentrations, 10 nM hPK-M2 and 1 µM of LDH) was added. Plates were immediately placed in ViewLux (Perkin-Elmer), and NADH fluorescence was determined at 30 s exposure intervals for between 3 and 6 min. Data were normalized to the uninhibited and EC_{100} activation using known activators such as FBP. The data has been deposited in PubChem (AID: 1540). Follow-up of synthesized analogues was determined using the same protocol (PubChem AIDs for L, M1 and R LDH-coupled assays are 1541, 1542, and 1543, respectively). This assay was also used to determine the $K_{\rm M}$'s for PEP and ADP in the presence and absence

of activator. Data was fit in GraphPad prism. Conversion of fluorescent units to pmols of NADH was performed using a standard curve of known NADH concentrations. Data was collected on the Perkin-Elmer Viewlux.

Synthesis of 2 and Analogues. Complete procedures for the synthesis of 2 and all analogues are presented in the Supporting Information. Proton NMR, HRMS, and purity (as judged by peak integration on two separate HPLC gradients) for all compounds are listed in the Supporting Information. 2: ¹H NMR (400 MHz, DMSO- d_6) δ : 2.94 (s, 8H), 3.86 (s, 3H), 4.23–4.45 (m, 4H), 7.01–7.08 (m, 1H), 7.08–7.19 (m, 4H), 7.57–7.67 (m, 2H). HRMS; calculated for C₁₉H₂₂N₂O₇S₂ (M+), 454.0868; found, 454.08718.

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Note Added after ASAP Publication. This paper was published ASAP on December 17, 2009 with incorrect data in Scheme 3. The revised version was published on December 23, 2009. Figure 2 was modified on January 6, 2010.

Supporting Information Available: Assay protocols, experimental procedures, and spectroscopic data (¹H, LC/MS and HRMS) are listed for all compounds. This material is available free of charge via the Internet at http://pubs.acs.org.

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