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Multidrug Resistance Protein 1 (MRP1/ABCC1)-Mediated Cellular **Protection and Transport of Methylated Arsenic Metabolites** Differs between Human Cell Lines

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ABSTRACT

The ATP-binding cassette (ABC) transporter multidrug resistance protein 1 (MRP1/ABCC1) protects cells from arsenic (a proven human carcinogen) through the cellular efflux of arsenic triglutathione [As(GS)₃] and the diglutathione conjugate of monomethylarsonous acid [MMA (GS)₂]. Previously, differences in MRP1 phosphorylation (at Y920/S921) and N-glycosylation (at N19/N23) were associated with marked differences in As(GS)₃ transport kinetics between HEK293 and HeLa cell lines. In the current study, cell line differences in MRP1-mediated cellular protection and transport of other arsenic metabolites were explored. MRP1 expressed in HEK293 cells reduced the toxicity of the major urinary arsenic metabolite dimethylarsinic acid (DMAV), and HEK-WT-MRP1-enriched vesicles transported DMAV with high apparent affinity and capacity ($K_{\rm m}$ 0.19 μ M, $V_{\rm max}$ 342 pmol·mg⁻¹protein·min⁻¹). This is the first report that MRP1 is capable of exporting DMAV, critical

for preventing highly toxic dimethylarsinous acid formation. In contrast, DMAV transport was not detected using HeLa-WT-MRP1 membrane vesicles. MMA(GS)₂ transport by HeLa-WT-MRP1 vesicles had a greater than threefold higher V_{max} compared with HEK-WT-MRP1 vesicles. Cell line differences in DMA and MMA(GS)₂ transport were not explained by differences in phosphorylation at Y920/S921. DMAV did not inhibit, whereas MMA(GS)2 was an uncompetitive inhibitor of As(GS)₃ transport, suggesting that DMA^V and MMA(GS)₂ have nonidentical binding sites to As(GS)₃ on MRP1. Efflux of different arsenic metabolites by MRP1 is likely influenced by multiple factors, including cell and tissue type. This could have implications for the impact of MRP1 on both tissue-specific susceptibility to arsenic-induced disease and tumor sensitivity to arsenic-based therapeutics.

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Introduction

Arsenic is a proven human carcinogen, causing lung, skin, and bladder tumors (IARC, 2012). Chronic arsenic exposure is associated with increased incidences of kidney and liver tumors and a myriad of noncancerous adverse health effects (Platanias, 2009; IARC, 2012; Naujokas et al., 2013). Millions of people worldwide are exposed to levels of arsenic above the World Health Organization acceptable level of 10 μ g/l, predominantly through the consumption of groundwater naturally contaminated with inorganic arsenic [arsenite (As^{III}) and arsenate (As^V)] (Rahman et al., 2009). In addition to environmental

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exposures, arsenic trioxide is used clinically in the treatment of acute promyelocytic leukemia and is in clinical trials for the treatment of other hematologic and solid tumors (Kritharis et al., 2013; Ally et al., 2016; Cicconi and Lo-Coco, 2016; Falchi et al., 2016). Other arsenic compounds are also in clinical trials for the treatment of various cancers (Khairul et al., 2017). Thus, understanding the cellular handling of arsenic is critical for the development of therapeutics to treat chronic arsenic exposure and to maximize the clinical effectiveness of arsenic based drugs.

Cellular uptake of arsenic has been recently reviewed (Mukhopadhyay et al., 2014; Roggenbeck et al., 2016). Once inside most mammalian cells, arsenic undergoes extensive methylation (Vahter, 1999; Drobna et al., 2010). In humans, the four major methylation products are monomethylarsonic acid (MMAV), monomethylarsonous acid (MMAIII), dimethylarsinic acid (DMAV), and dimethylarsinous acid (DMAIII) (Thomas et al., 2007). Arsenic methylation has a significant impact on its toxicity, tissue distribution, and retention (Thomas et al., 2004, 2007; Wang et al., 2015). Although arsenic methylation results in an increased rate of arsenic whole body clearance (Drobna et al., 2009, 2010; Hughes et al., 2010) and reduces susceptibility to acute arsenic toxicity (Yokohira et al., 2010, 2011), trivalent methylated forms of arsenic (MMA^{III} and DMA^{III}) are considered bioactivation products because they are more reactive

ABBREVIATIONS: ABC, ATP-binding cassette; As^{III}, arsenite; As^V, arsenate; As(GS)₃, arsenic triglutathione; DMA^{III}, dimethylarsinous acid; DMA^V, dimethylarsinic acid; GSH, glutathione; HEK293, human embryonic kidney 293; IC50, half-maximal inhibitory concentration; MMAIII, monomethylarsonous acid; MMAV, monomethylarsonic acid; MMA(GS)2, monomethylarsenic diglutathione; MRP, multidrug resistance protein; MSD, membrane spanning domain; NBD, nucleotide binding domain; PIC, phosphatase inhibitor cocktail; WT, wild type.

metabolites than As^{III} (Petrick et al., 2000; Styblo et al., 2000; Mass et al., 2001; Kligerman et al., 2003; Moe et al., 2016).

In addition to methylation, arsenic can be conjugated with reduced glutathione (GSH/GS) (Leslie, 2012). Arsenic triglutathione [As(GS)₃] and the diglutathione conjugate of the highly toxic MMA^{III} [MMA(GS)₂] have been isolated from rat bile and mouse urine, thus these two As-GSH complexes are formed physiologically and account for a major fraction of arsenic in urine and bile (Kala et al., 2000, 2004; Suzuki et al., 2001; Cui et al., 2004; Bu et al., 2011). The ATP-binding cassette (ABC) transporter multidrug resistance protein 1 (MRP1, gene symbol *ABCC1*), along with the related MRP2 and MRP4 (*ABCC2* and *ABCC4*, respectively) mediate the cellular export of multiple methylated and/or glutathionylated metabolites of arsenic (Kala et al., 2000, 2004; Leslie et al., 2004; Carew and Leslie, 2010; Carew et al., 2011; Banerjee et al., 2014; Shukalek et al., 2016).

MRP1 is a 190-kDa phosphoglycoprotein with three polytopic membrane spanning domains (MSDs) and two nucleotide binding domains (NBDs) arranged as MSD0-MSD1-NBD1-MSD2-NBD2 (Cole, 2014). MRP1 confers resistance to a chemically diverse array of anti-cancer drugs and is involved in the cellular export of physiologic compounds including GSH, glutathione disulfide, 17β -estradiol 17-(β -D-glucuronide), and leukotriene C_4 (Cole, 2014). Furthermore, MRP1 transports a variety of xenobiotics often conjugated to GSH, glucuronate, or sulfate (Jedlitschky et al., 1996; Loe et al., 1996a,b; Leslie et al., 2005). Included in this list are the arsenic metabolites $As(GS)_3$ and $MMA(GS)_2$, the glutathionylated forms of inorganic arsenic (As^{III} and As^V), and MMA^{III} , respectively (Leslie et al., 2004; Carew et al., 2011).

Interestingly, As(GS)₃ is transported by MRP1 expressed in HEK293 cells with markedly different kinetics than by MRP1 expressed in HeLa cells (Shukalek et al., 2016). Further investigation revealed that MRP1 affinity and capacity for As(GS)3 was associated with the phosphorylation status of two residues in the linker region between NBD1 and MSD2 (Y920 and S921). Furthermore, the glycosylation status of two residues in the amino terminus (N19 and N23) influenced the stability of Y920 and/or S921 phosphorylation (Shukalek et al., 2016). Given this cell line difference, the first objective of the current study was to determine differences in arsenical cytotoxicity between the HEK-MRP1 and HeLa-MRP1 cell lines. The second objective was to use MRP1-enriched membrane vesicles isolated from HEK and HeLa cells to determine the cell line differences in MRP1 transport function. The third objective was to investigate the influence of Y920/S921 phosphorylation on the ability of HEK-MRP1-enriched membrane vesicles to transport arsenic metabolites in addition to $As(GS)_3$.

Materials and Methods

Materials. Carrier-free ⁷³As^V (158 Ci/mol) was purchased from Los Alamos Meson Production Facility (Los Alamos, NM). Tris base, GSH, ATP, AMP, sucrose, DMA^V (>99% purity), MgCl₂, creatine kinase, GSH reductase, creatine phosphate, NADPH, sodium (meta)-arsenite [Na₂AsO₂] (>99% pure), As (>98% purity), sodium metabisulfite [Na₂S₂O₅], and sodium thiosulfate [Na₂S₂O₃] were from Sigma-Aldrich (Oakville, Canada). Protease inhibitor cocktail tablets (Complete, Mini, EDTA-free) and PhosSTOP phosphatase inhibitor cocktail (PIC) tablets were purchased from Roche Applied Science (Laval, Canada). Phenylmethylsulfonyl fluoride was from Bioshop Canada Inc. (Burlington, Canada). Suprapur nitric acid was purchased from Merck (Darmstadt, Germany). MMAIII and DMAIII in the form diiodomethylarsine (CH₃AsI₂) and iododimethylarsine ([CH₃]₂AsI), respectively, were synthesized as previously described (Cullen et al., 2016) and were at least 99% pure as confirmed by NMR analysis. The rat monoclonal antibody (mAb) MRPr1 was from Novus Biologicals (Littleton, CO), while rabbit anti-Na⁺/K⁺ATPase (H-300) was purchased from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA).

Cell Lines and Stable Transfection. HEK293T cells were obtained from the American Type Culture Collection (Rockville, MD) and maintained in Dulbecco's modified Eagle's medium with 7.5% fetal bovine serum. HeLa and HEK293 cell lines stably expressing the empty pcDNA3.1(-) vector (HeLa-vector) and/or pcDNA3.1(-)-MRP1 (HeLa-MRP1 and HEK-MRP1) were gifts from Dr. Susan P.C. Cole (Queen's University, Kingston, Canada) and generated as described previously (Ito et al., 2001; Conseil and Cole, 2013). The HEK-vector expressing cell line (HEK-V4) was generated as described previously (Banerjee et al., 2014). HeLa stables were maintained in Roswell Park Memorial Institute (RPMI)-1640 medium with 5% calf serum and 600 µg/ml G418 (Geneticin). HEK293 stables were maintained in Dulbecco's modified Eagle's medium with 7.5% fetal bovine serum and 600 µg/ml G418. Stable cell lines were checked for the proportion of cells expressing MRP1 by flow cytometry (BD FACS Calibur, Cross Cancer Institute) using the MRP1-specific MAb MRPr1, as described previously (Hipfner et al., 1994; Leslie et al., 2003). Populations of less than 80% were not used in experiments.

Cytotoxicity Testing. The cytotoxicity of five arsenic species was measured using HEK-vector, HEK-MRP1, HeLa-vector, and HeLa-MRP1 as previously described (Carew et al., 2011). Briefly, cells were seeded in 96-well plates at 1×10^{-1} 10⁴ cells/well and grown for 24 hours. In quadruplicate, cells were treated with As^{III} (0.01–300 μ M), As^V (0.05–5000 μ M), MMA^{III} (0.03–30 μ M), DMA^{III} (0.01–300 μ M), or DMA^V (0.01–30 mM) for 72 hours. These doses were experimentally determined to range from nontoxic to causing complete loss of viability over 72 hours. The pH of DMAV was adjusted to pH 7.4 prior to treating cells. Cytotoxicity was determined using the tetrazolium-based MTS assay [CellTiter 96 Aqueous Non-Radioactive Cell Proliferation Assay (Promega, Madison, WI)], according to the manufacturer's instructions. The IC₅₀, defined as the concentration of arsenical that resulted in half the maximal toxic effect, for each arsenical was determined for HEK-MRP1 and HEK-vector cells using the sigmoidal dose-response equation in GraphPad Prism (GraphPad Software, La Jolla, CA). Relative resistance values (equivalent to relative protection), defined as the ratio of the IC50 value in HEK-MRP1 to that in HEK-vector, were determined for each arsenical tested.

Site-Directed Mutagenesis. Mutants of MRP1 (S905A-, S915A-, S916A-, S917A-, Y920F-, S921A-, Y920F/S921A-, Y920E/S921E-, and T931A-MRP1) were generated previously (Shukalek et al., 2016). In addition, several new potential phosphorylation mutants (S916E-, S918A-, S919A-, and S919E-MRP1) were generated using the QuikChange II XL site-directed mutagenesis kit (Stratagene; Agilent Technologies, Santa Clara, CA). pcDNA3.1(-)-MRP1 was used as the PCR template, and mutagenesis was carried out according to the manufacturer's instructions using mutagenic primers from Integrated DNA Technology (Coralville, IA); sequences are available upon request. The incorporation of desired mutations was confirmed by DNA sequencing (Molecular Biology Servicing Unit, University of Alberta, Edmonton, Canada).

Expression of Wild-Type and Mutant Forms of MRP1 in HEK293T Cells, Preparation of Membrane Vesicles, and Immunoblots. HEK293T cells were transfected using the calcium-phosphate method as described previously (Carew and Leslie, 2010) and incubated for 48- to 72-hours posttransfection. For membrane vesicle preparations, WT and MRP1 mutant-transfected cells were collected by centrifugation, layered with Tris sucrose buffer (50 mM Tris, pH 7.4, 250 mM sucrose) containing CaCl₂ (0.25 mM), EDTA-free protease inhibitors, and where indicated, phosSTOP PIC, and cell pellets stored at -80°C until plasma membrane-enriched vesicles were isolated as described previously (Carew and Leslie, 2010). Expression of WT and MRP1 mutants were confirmed by immunoblotting as described previously (Shukalek et al., 2016), using the MRPr1 antibody (1:10,000 dilution). Blots were also probed for Na⁺/K⁺-ATPase as a loading control using the Na⁺/K⁺-ATPase-specific antibody H-300 (1:10,000 dilution), except in the case of comparison of MRP1 levels between the two different cell lines (the Na+/K+-ATPase level was found to be different). Thus, blots were stained with Coomassie blue and normalized to total protein levels in each lane. Relative levels of MRP1 were quantified using ImageJ Software (National Institutes of Health, Bethesda, MD) and/or ImageLab Software (Bio-Rad, Hercules, CA).

MMA(GS)₂, DMA^V, and As(GS)₃ Vesicular Transport Assays. Reduction of arsenate [⁷³As^V] into arsenite [⁷³As^{III}] and subsequent synthesis of ⁷³As(GS)₃ from ⁷³As^{III} and GSH were carried out as described previously (Reay and Asher, 1977; Shukalek et al., 2016). MMA(GS)₂ was synthesized from MMA^{III} and GSH as described previously (Carew et al., 2011; Banerjee et al.,

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2014). Vesicular transport was carried out in triplicate for each substrate, employing previously described methods (Carew et al., 2011; Banerjee et al., 2014; Shukalek et al., 2016). The amount of ⁷³As(GS)₃ transported was quantified using liquid scintillation counting, as described previously (Leslie et al., 2004; Shukalek et al., 2016). The amount of MMA(GS)₂ and DMA^V transported was quantified by inductively coupled plasma mass spectrometry using the standard addition method, as described previously (Banerjee et al., 2014). ATP-dependent transport was calculated by subtracting transport in the presence of AMP from transport in the presence of ATP, and data are expressed as picomoles or nanomoles As(GS)₃, MMA(GS)₂, or DMA^V transported per milligram protein per minute.

The linear range of DMA $^{\rm V}$ transport was determined by incubating the HEK-WT-MRP1 or HEK-vector membrane vesicles with DMA $^{\rm V}$ (1 μ M) in transport buffer at 37°C for the indicated time points. Kinetic parameters were determined by measuring the initial rate of DMA $^{\rm V}$ and MMA(GS)₂ transport at eight different substrate concentrations [0.1–2.5 μ M for DMA $^{\rm V}$ and 1–200 μ M for MMA(GS)₂] at 20 seconds and 1 minute, respectively. Curve fitting was done by nonlinear regression analysis using GraphPad Prism 6 software.

The ability of DMA^V and MMA(GS)₂ to inhibit ⁷³As(GS)₃ transport was first characterized by using a fixed concentration of ⁷³As(GS)₃ (1 μ M, 40 nCi) in the presence of increasing concentrations of DMA^V (0.04 and 1000 μ M) or MMA (GS)₂ (0.01–300 μ M). The conditions for the synthesis of As(GS)₃ resulted in the presence of 3 mM GSH in transport reactions (Leslie et al., 2004). Thus, As(GS)₃ inhibition experiments were completed under plus GSH conditions. This is important because the in vitro inhibition of MRP1 by certain compounds can be enhanced by physiologic concentrations of GSH (Cole and Deeley, 2006). IC₅₀ values were calculated for MMA(GS)₂ inhibition using GraphPad Prism 6 Software [nonlinear regression log(inhibitor) vs. response variable slope (4 parameters)]. To determine the mode of inhibition of ⁷³As(GS)₃ by MMA(GS)₂, the K_i of MMA(GS)₂ was determined by performing ⁷³As(GS)₃ (0.1–20 μ M, 40–100 nCi) transport in the presence of three different MMA (GS)₂ concentrations (5, 10, and 15 μ M), as described previously (Leslie et al., 2004).

All transport data were normalized as needed to correct for any difference in level of HeLa-WT-MRP1 or mutant MRP1 expressed in HEK cells relative to HEK-WT-MRP1 as determined by immunoblotting of each membrane vesicle preparation. Positive control transport experiments using MMA(GS)₂ or As(GS)₃ were run for each vesicle preparation, as described above and previously (Leslie et al., 2004; Carew et al., 2011).

Results

MRP1 Expressed in HEK293 Cells Decreases the Cytotoxicity of As^{III}, As^V, MMA^{III}, and DMA^V. Previously we showed that MRP1 stably expressed in HeLa cells reduced the toxicity of As^{III}, As^V, and MMA^{III}, but not MMA^V, DMA^{III}, or DMA^V, relative to HeLa cells expressing empty vector alone (Carew et al., 2011). Given that we recently reported substantial differences in the transport of As(GS)₃ between MRP1-enriched membrane vesicles isolated from HEK293 and HeLa cells (Shukalek et al., 2016), differences in arsenical cytotoxicity between the two cell lines were investigated. Thus, the cytotoxicity of five of these arsenicals (As^{III}, As^V, MMA^{III}, DMA^{III}, and DMA^V) in HEK-MRP1 and HEK-vector cell lines was determined in parallel with HeLa-MRP1 and HeLa-vector for comparison [Fig. 1; Table 1, and (Carew et al., 2011)].

The arsenic compounds that HEK-MRP1 cells conferred resistance to (relative to HEK-vector) were the same as the HeLa cell line pair (Fig. 1, A–C; Table 1), except that HEK-MRP1 conferred a significantly increased level of resistance against DMA^V (relative resistance value of 1.4, P < 0.05) [Fig. 1D; Table 1, and (Carew et al., 2011)]. In addition, HEK-MRP1 cells had a relative resistance of 9.2 against As^V (Fig. 1B; Table 1), whereas HeLa-MRP1 cells had a relative resistance of only 2.1 [Table 1 and (Carew et al., 2011)]. These results led us to characterize the cell line differences for DMA^V and As^V transport using MRP1-enriched membrane vesicles. Furthermore, although resistance to MMA^{III} was conferred by both HEK-MRP1 and HeLa-MRP1 cell lines [Fig. 1C; Table 1 and (Carew et al., 2011)], differences in transport characteristics between membrane vesicles prepared from these cell lines were also investigated for MMA(GS)₂.

Transport of DMA^V by MRP1-Enriched Membrane Vesicles. To test if differences in cytotoxicity were due at least in part to differences in ATP-dependent cellular efflux, DMA^V transport by HEK- and HeLa-WT-MRP1 membrane vesicles was measured at 0.05 and 1 μ M of DMA^V (Fig. 2A). MRP1-enriched vesicles prepared from transiently transfected HEK293T cells transported DMA^V (101 \pm 15 pmol· mg⁻¹ protein·min⁻¹ at 0.05 μ M and 264 \pm 42 pmol·mg⁻¹ protein·min⁻¹ at 1 μ M). In contrast, DMA^V transport was not detected for HeLa-WT-MRP1 vesicles (Fig. 2A), despite the fact that the same vesicle preparations were functional for MMA(GS)₂ transport (Fig. 4A), and

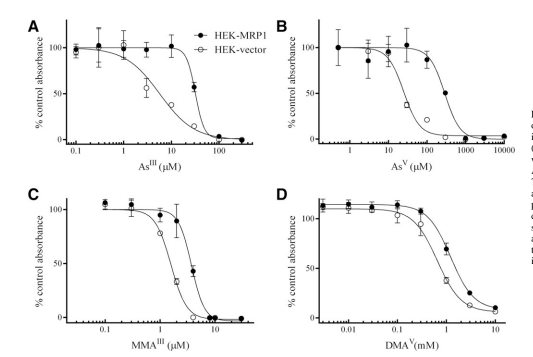


Fig. 1. Effect of selected arsenic compounds on the viability of HEK293 cells stably expressing human MRP1. Cells expressing empty vector (HEK-vector) (\bigcirc) and MRP1 (HEK-MRP1) (\bigcirc) were incubated in the presence of $\mathrm{As^{II}}$ (A), $\mathrm{As^{V}}$ (B), MMA^II (C), or DMA (D) for 72 hours. Cell viability was determined using a tetrazolium-based cytotoxicity assay. Data points are means (\pm S.D.) of quadruplicate determinations in a representative experiment; similar results were obtained in at least two additional experiments (mean IC₅₀ and relative resistance values from independent experiments are shown in Table 1).

TABLE 1
Resistance of MRP1-transfected HEK293 and HeLa cells to inorganic and methylated arsenic species

	$IC_{50} (\mu M) (\pm S.D.)$		Relative Resistance ^a	$IC_{50} (\mu M) (\pm S.D.)$		Relative Resistance
	HEK-Vector	HEK-MRP1	Relative Resistance	HeLa-Vector	HeLa-MRP1	Relative Resistance
As ^{III} As ^V MMA ^{III} DMA ^{III} DMA ^V	4.9 ± 1.0 26 ± 6.7 1.7 ± 0.12 1.2 ± 1 1090 ± 290	$25.5 \pm 4.2 221 \pm 62 3.9 \pm 0.5 1.6 \pm 1.6 1500 \pm 250$	5.3 ± 0.9**** 9.2 ± 3.6*** 2.3 ± 0.2*** 1.2 ± 0.2 1.4 ± 0.2*	7.0 ± 1.2 156 ± 55 5.0 ± 1.2 6 ± 0.5^{b} 690 ± 430	$ \begin{array}{c} 17.5 \pm 2.4 \\ 303 \pm 51 \\ 7.6 \pm 1.3 \\ 7 \pm 2.3^{b} \\ 680 \pm 450 \end{array} $	$2.5 \pm 0.3***$ $2.1 \pm 0.5**$ $1.5 \pm 0.2*$ 1.2 ± 0.4^{b} 1.0 ± 0.13

"Relative resistance factors were calculated by dividing the IC_{50} values obtained for the MRP1-expressing cell line by the IC_{50} values obtained for empty vector expressing cell line. Values shown are the mean (\pm S.D.) obtained from at least three independent experiments. IC_{50} values for the MRP1 expressing cell lines were compared with the IC_{50} values for the empty vector control cell lines using a Student's t test (*P< 0.05, **P< 0.001, ***P< 0.001, ****P< 0.0001).

comparable levels of MRP1 were present (Supplemental Fig. 1). For certain compounds transported by MRP1, GSH can either be required for or enhance transport (Cole, 2014). To determine whether this was the case for DMA^V, transport assays were completed in the presence of 3 mM GSH and this had no effect on HEK- or HeLa-WT-MRP1 DMA^V transport activity (data not shown). Thus, we report for the first time that MRP1 is capable of transporting this important methylated arsenic metabolite, at least under certain conditions.

Kinetic Analysis of MRP1-Mediated DMA^V Transport. Transport of DMA^V by HEK-WT-MRP1 membrane vesicles was then further characterized. The linear range of DMA^V (1 μ M) transport versus time was measured for HEK-WT-MRP1 and HEK-vector control membrane vesicles (Fig. 2B). Transport by HEK-WT-MRP1 was linear for up to 30 seconds and reached a maximal activity of 138 ± 24 pmol·mg⁻¹ protein at 30 seconds. ATP-dependent transport of DMA^V by HEK-vector was very low and similar to transport observed in the presence of AMP. HEK-WT-MRP1 membrane vesicle transport of DMA^V was characterized kinetically by determining the initial rates of transport over several concentrations of DMA^V (Fig. 2C; Table 2). HEK-MRP1 was found to transport DMA^V with high apparent affinity and capacity ($K_{\rm m}$ of 0.19 ± 0.06 μ M and $V_{\rm max}$ of 342 ± 37 pmol· mg⁻¹ protein·min⁻¹).

Analysis of DMA^V Transport by HEK-MRP1 Phosphorylation Mutants. Due to the complete lack of DMA^V transport by HeLa-WT-MRP1 and substantial transport by HEK-WT-MRP1 membrane vesicles, this cell line difference was further explored. We previously showed that HEK293 and HeLa cell line differences in As(GS)₃ transport by MRP1 were associated with differences in phosphorylation at Y920/S921 (Shukalek et al., 2016). To determine if phosphorylation differences at these sites were responsible for the difference in DMA^V transport by MRP1 between the two cell lines, DMA^V transport by MRP1-Y920/S921 dephosphorylation- and phosphorylation-mimicking mutants was investigated.

Dephosphorylation-mimicking HEK-Y920F/S921A-MRP1 membrane vesicles exhibited a complete loss of DMA^V transport (Fig. 3A), suggesting these sites are critical for the cell line difference. However, the phosphorylation-mimicking HEK-Y920E/S921E-MRP1, which we expected would restore DMA^V transport, also completely lacked DMA^V transport (Fig. 3A). Furthermore, individual dephosphorylation mimicking mutants Y920F-MRP1 and S921A-MRP1 also completely lost function (data not shown). Mutant membrane vesicle preparations had MRP1 levels similar to HEK-WT-MRP1 (Supplemental Fig. 1) and were functional for MMA(GS)₂ (Fig. 4B; Table 2) and/or As(GS)₃ (Supplemental Fig. 2A) transport.

Analysis of DMA^V Transport by HeLa-WT-MRP1 in the Presence of Phosphatase Inhibitors. The possible modulation of DMA^V transport by MRP1 through phosphorylation was also investigated by studying the transport in MRP1-enriched vesicles

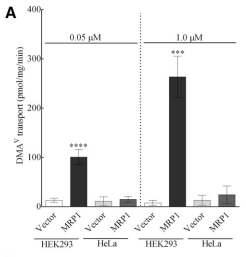
prepared from HeLa cells in the presence or absence of a phosphatase inhibitor cocktail (PIC) at two different substrate concentrations $(0.05 \text{ and } 1 \mu\text{M})$ (Fig. 3B). We previously showed for As(GS)₃ that HeLa-WT-MRP1 membrane vesicles prepared in the presence of a PIC have a 19- and 12-fold increase in $K_{\rm m}$ and $V_{\rm max}$, respectively, compared with HeLa-WT-MRP1 prepared in the absence of a PIC (Shukalek et al., 2016). In contrast, the inclusion of a PIC did not influence DMAV transport by HeLa-WT-MRP1 at either concentration (Fig. 3B). To ensure that the HeLa-WT-MRP1 ± PIC membrane vesicles used in the DMAV transport assays were functional, and the PIC active, As(GS)3 transport experiments were completed on the same vesicle preparations (Fig. 3C). Consistent with our previous report (Shukalek et al., 2016), As(GS)₃ transport by HeLa-WT-MRP1 vesicles, prepared in the presence of a PIC, was significantly higher than for HeLa-WT-MRP1 vesicles prepared in the absence of a PIC. These results could suggest that mechanisms other than phosphorylation are likely responsible for the cell line difference observed in DMA^V transport by HEK-WT-MRP1 compared with HeLa-WT-MRP1. Alternatively, a differential phosphorylation site that remains stable in the absence of PIC could be involved.

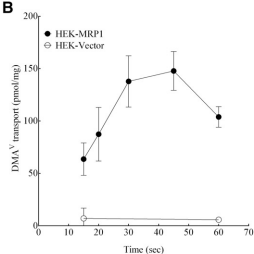
The lack of DMA^V transport by all mutants of Y920/S921 could also be due to the disruption of phosphorylation/dephosphorylation of neighboring sites that are critical for DMA^V transport. We screened the influence of seven additional putative phosphorylation sites in the linker region (S905-, S915-, S916-, S917-, S918-, S919-, T931-MRP1) by mutating them to phospho-mimicking and/or dephospho-mimicking residues in full-length MRP1, expressing them in HEK293 cells, preparing membrane vesicles, and measuring DMA^V transport. Surprisingly, all mutants lacked DMA^V transport activity, despite the fact that they were comparable to HEK-WT-MRP1 for As(GS)₃ transport and MRP1 level (Supplemental Fig. 2).

Inhibition of As(GS)₃ Transport by DMA^V. To further characterize the differences in interaction of As(GS)₃ and DMA^V with MRP1, transport of As(GS)₃ (1 μ M) by HEK-WT-MRP1 vesicles was measured in the presence of DMA^V. DMA^V did not inhibit As(GS)₃ transport even at a concentration of 1 mM (data not shown), four orders of magnitude greater than the apparent K_m of DMA^V for MRP1 (Table 2).

MRP1 Does Not Transport As^V. Given the striking difference in As^V relative resistance levels between the HeLa and HEK293 cell line pairs [Fig. 1B; Table 1 and (Carew et al., 2011)], the ability of HEK-MRP1 and HeLa-MRP1 membrane vesicles to transport As^V was investigated. As^V (1 and 10 μ M) was not transported by MRP1-enriched membrane vesicles prepared from HEK293 cells in the presence or absence of 3 mM GSH (data not shown). Similar results were obtained for membrane vesicles prepared from HeLa-MRP1 cells (data not shown). These results are consistent with a lack of detectable As^V transport (in the presence and absence of GSH) by MRP1-enriched

^bPreviously published in Carew et al. (2011) but done in parallel with HEK experiments.





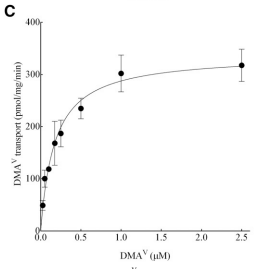


Fig. 2. ATP-dependent transport of DMA^V by MRP1-enriched membrane vesicles. Transport experiments were done with membrane vesicles (20 μ g of protein) prepared from HEK293T cells transiently transfected with WT-MRP1 (black bars or symbols) or empty pcDNA3.1(-) (vector) (white bars or symbols) or HeLa cells stably transfected with WT-MRP1 (dark gray bars) or empty vector (light gray bars). For individual experiments, transport was done in triplicate, and then reactions were pooled for analysis by inductively coupled plasma mass spectrometry (ICP-MS). Bars and symbols represent the means (\pm S.D.) of three independent experiments. (A) Vesicles were incubated for 20 seconds at 37°C in transport buffer with DMA^V (0.05 or 1 μ M). Statistically significant differences in DMA^V transport were determined using a one-way ANOVA followed by a Dunnett's multiple comparisons

vesicles prepared from the H69AR small cell lung cancer cell line and our previous conclusion that inorganic arsenic (As^{III} and As^V) is metabolized to As(GS)₃ before transport by MRP1 (Leslie et al., 2004).

Transport of MMA(GS)₂ by MRP1-Enriched Membrane Vesicles. We previously showed that MMA^{III} is transported by HEK-MRP1 in the form MMA(GS)₂ and that HeLa-MRP1 cells confer protection against MMA^{III} relative to HeLa-vector cells (Carew et al., 2011). In the current study, HEK-MRP1 cells were also found to confer protection against MMA^{III} relative to HEK-vector (Table 1). To determine if differences in transport characteristics existed between cell lines, MMA (GS)₂ transport by HEK- and HeLa-WT-MRP1-enriched vesicles was compared. Under the conditions tested, MMA(GS)₂ (1 μ M) transport by HeLa-WT-MRP1 vesicles was 1.6-fold higher (P < 0.01) than HEK-WT-MRP1 vesicles (Fig. 4A).

Kinetic Analysis of MRP1-Mediated MMA(GS)₂ Transport. To determine if the increased transport of MMA(GS)₂ was due to changes in $K_{\rm m}$ and/or $V_{\rm max}$, MRP1-mediated transport of MMA(GS)₂ was measured at an initial rate over eight different concentrations of MMA(GS)₂ (Fig. 4B; Table 2). HEK- and HeLa-WT-MRP1 membrane vesicles were found to have similar apparent affinity for MMA(GS)₂ $(K_{\rm m} {\rm of } 23 \pm 2.2 {\rm and } 33 \pm 24 ~\mu {\rm M}, {\rm respectively}), {\rm similar to what we}$ previously reported for HEK-WT-MRP1 (Carew et al., 2011). In contrast, the V_{max} was 3.4-fold higher for MMA(GS)₂ transport by HeLa-WT-MRP1 (V_{max} of 17 \pm 5.3 nmol·mg⁻¹·min⁻¹) than HEK-WT-MRP1 (V_{max} of 4.9 \pm 0.5 nmol·mg⁻¹·min⁻¹) membrane vesicles (Fig. 4B; Table 2). In addition, kinetic characterization of MMA(GS)₂ transport by the double dephosphorylation-mimicking mutant HEK-Y920F/S921A-MRP1 previously demonstrated to have substantially reduced apparent $K_{\rm m}$ and $V_{\rm max}$ values for As(GS)₃ relative to HEK-WT-MRP1(Shukalek et al., 2016) was completed. Interestingly, HEK-Y920F/S921A-MRP1 had very similar apparent affinity ($K_{\rm m}$ of 24 \pm 5.5 μ M) and capacity ($V_{\rm max}$ of 5.3 \pm 2.2 nmol·mg⁻¹·min⁻¹) to that of HEK-WT-MRP1 (Fig. 4B; Table 2). Consistent with this, the phosphorylation-mimicking mutant HEK-Y920E/S921E-MRP1 also had similar apparent affinity and capacity to HEK-WT-MRP1 (mean $K_{\rm m}$ of 24 μ M, $V_{\rm max}$ of 5.5 nmol·mg⁻ protein·min⁻¹, n = 2) (Table 2). These data suggest that, in contrast with As(GS)₃ transport, phosphorylation of Y920/S921 has little influence on the transport of MMA(GS)₂ by MRP1.

Inhibition of As(GS)₃ **Transport by MMA**(GS)₂. Although the $K_{\rm m}$ of As(GS)₃ for MRP1 was >10-fold higher for HEK-WT-MRP1 compared with HeLa-WT-MRP1 and HEK-Y920F/S921A-MRP1 (Shukalek et al., 2016), no difference between the apparent $K_{\rm m}$ value for MMA(GS)₂ transport was observed for these membrane vesicles (Fig. 4; Table 2). This result suggested that MMA(GS)₂ and As(GS)₃ interact at nonidentical binding sites. To begin to characterize the differences in interaction of As(GS)₃ and MMA(GS)₂ with MRP1, transport of As(GS)₃ (1 μ M) by HEK-WT-MRP1 vesicles was measured in the presence of increasing concentrations of MMA(GS)₂ (Fig. 5A). MMA(GS)₂ was found to potently inhibit As(GS)₃ transport with an IC₅₀ value of 11 \pm 1.5 μ M.

The inhibition of $As(GS)_3$ transport by $MMA(GS)_2$ was further characterized by measuring the effect of $MMA(GS)_2$ (5, 10, and 15 μM)

post hoc test with HEK-Vector as the control group (***P < 0.001; ****P < 0.0001). (B) Time course of ATP-dependent DMA $^{\rm V}$ transport was determined by incubating membrane vesicles with DMA $^{\rm V}$ (1 μ M) in transport buffer at 37°C for the indicated time points. (C) HEK-WT-MRP1 membrane vesicles were incubated for 20 seconds at 37°C with increasing concentrations of DMA $^{\rm V}$ (0.025–2.5 μ M). Data were fitted using a one-site Michaelis-Menten kinetic model with GraphPad Prism6.

TABLE 2

Kinetic parameters of MMA(GS)₂ and DMA^V transport by MRP1 (and phosphorylation mutants)

Kinetic parameter								
Compound	Cell Line	Variant	n	$K_{\rm m}~(\mu{\rm M})$	$V_{ m max}{}^a$			
DMA ^V	НЕК293Т	WT	3	0.19 ± 0.06	342 ± 37^{b}			
$MMA(GS)_2$	HEK293T	WT	3	23 ± 2.2	4.9 ± 0.5			
		Y920F/S921A	4	24 ± 5.5	5.3 ± 2.2			
		Y920E/S921E	2	29, 18	5.8, 5.0			
	HeLa	WT	3	33 ± 24	16.8 ± 5.3**			

 $[^]aV_{\rm max}$ values are in nmol mg $^{-1}$ protein min $^{-1}$ unless otherwise indicated and were corrected for MRP1-level relative to HEK-WT-MRP1 as described in *Materials and Methods*. b pmol mg $^{-1}$ protein min $^{-1}$.

on As(GS)₃ (0.1–20 μ M) transport (15 μ M shown in Fig. 5B). Michaelis-Menten analysis showed that MMA(GS)₂ at each concentration tested reduced both the apparent $K_{\rm m}$ and $V_{\rm max}$ values, suggesting an uncompetitive mode of inhibition with an average $K_{\rm i}$ of 7.3 \pm 5.1 μ M (\pm S.D., n = 3). These data are consistent with MMA(GS)₂ and As(GS)₃ interacting at nonidentical binding sites.

Discussion

The proven human carcinogen arsenic naturally contaminates the drinking water of hundreds of millions of people worldwide. One of the most affected countries is Bangladesh, where the arsenic contamination has been referred to as "the largest mass poisoning of a population in

history" (Smith et al., 2000). Understanding the cellular handling of arsenic, including efflux pathways, is critical for the prevention and treatment of arsenic-induced disease.

We investigated the ability of MRP1 to confer resistance to and/or transport important methylated arsenic metabolites when expressed in HEK293 cells compared with HeLa cells. The cellular resistance conferred by MRP1 against different arsenic species is useful information; however, resistance levels can be influenced by cellular metabolism and uptake efficiency. To draw conclusions about MRP1-mediated transport of specific arsenic compounds, it was critical to measure their transport directly using MRP1-enriched membrane vesicles. The population of membrane vesicles that is accumulating MRP1 substrates are inside-out, allowing the measurement of ATP-dependent transport with minimal influence of metabolism and cellular uptake. This allows the MRP1 contribution to cellular export of specific arsenic compounds to be evaluated and allows the accurate determination of kinetic parameters.

The most pronounced difference between cell lines was for DMA^V, which HEK-MRP1 cells conferred resistance to and HEK-WT-MRP1 membrane vesicles transported with high apparent affinity and capacity (Fig. 2; Table 1). This is the first report that MRP1 is capable of transporting this important arsenic metabolite. In contrast, HeLa-MRP1 cells did not confer resistance to DMA^V relative to HeLa-vector cells (Carew et al., 2011), and HeLa-WT-MRP1 membrane vesicles did not have detectable DMA^V transport activity (Fig. 2A).

The relative resistance conferred by MRP1 expressed in HEK293 cells was small, but significant (1.4-fold, P < 0.05, Fig. 1C; Table 1). This marginal resistance was in contrast with the high-affinity and high-capacity transport of DMA^V observed with HEK-WT-MRP1-enriched membrane vesicles (Fig. 2C; Table 2). A likely explanation for the

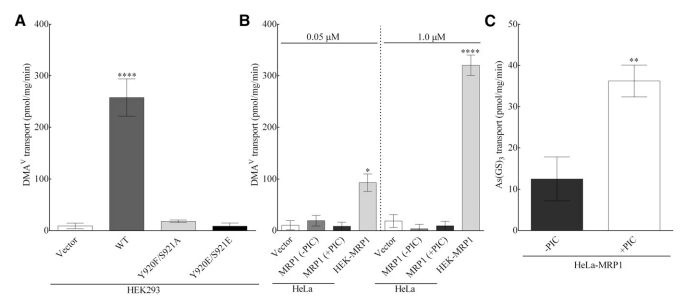


Fig. 3. Effect of Y920/S921-MRP1 mutation and/or phosphatase inhibitors on ATP-dependent transport of DMA^V or As(GS)₃ by MRP1-enriched membrane vesicles. Transport experiments were done with membrane vesicles (20 μ g of protein) prepared from HEK293T cells transiently transfected with WT-MRP1, Y920F/S921A-MRP1, Y920F/S921E-MRP1, or empty pcDNA3.1(-) (vector) or from HeLa cells stably transfected with WT-MRP1 or empty vector. Bars and symbols represent the means (\pm S. D.) of three independent experiments. (A) Membrane vesicles from HEK293T cells transiently transfected with WT-MRP1 (dark gray bars), Y920F/S921A-MRP1 (light gray bars), Y920F/S921E-MRP1 (black bars), or vector (white bars) were incubated for 20 seconds at 37°C with DMA^V (1 μ M). For individual experiments, transport was done in triplicate, and then reactions were pooled for analysis by ICP-MS. Statistically significant differences in DMA^V transport were determined using a one-way ANOVA followed by a Dunnett's multiple comparisons post hoc test using HEK-Vector as the control group (****P < 0.0001). (B) Membrane vesicles from HeLa cells stably transfected with vector (light gray bars) or WT-MRP1 prepared in the presence (white bars) or absence (black bars) of a PIC were incubated for 20 seconds at 37°C with DMA^V (0.05 or 1 μ M). HEK-WT-MRP1 membrane vesicles were used as a positive control. For individual experiments, transport was done in triplicate, and then reactions were pooled for analysis by ICP-MS. Statistically significant differences in DMA^V transport were determined using a one-way ANOVA followed by a Dunnett's multiple comparisons post hoc test using HeLa-Vector as the control group (**P < 0.05; *****P < 0.0001). (C) ATP-dependent transport of As(GS)₃ by MRP1-enriched membrane vesicles from stably transfected HeLa cells prepared with (white bar) or without (black bar) PIC. The membrane vesicles were incubated for 3 minutes at 37°C with As(GS)₃ (1 μ M). Statistically significant differences in As(GS)₃ transport wer

^{**}Significantly different from HEK293-MRP1 (P < 0.01; one-way ANOVA with Tukey's multiple comparisons post hoc test).

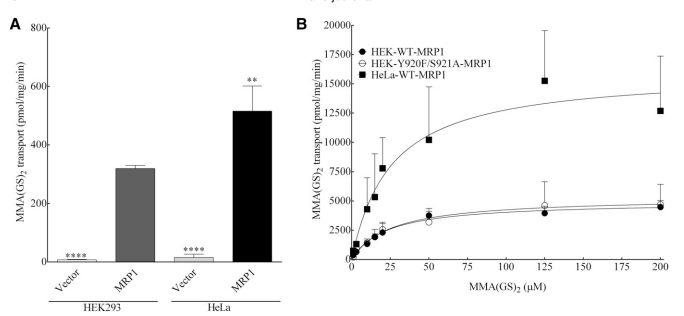


Fig. 4. ATP-dependent transport of MMA(GS)₂ by MRP1-enriched membrane vesicles. Transport experiments were done with membrane vesicles (20 μ g of protein) prepared from HEK293T cells transiently transfected with WT-MRP1, Y920F/S921A-MRP1, or empty pcDNA3.1(-) (vector) or from HeLa cells stably transfected with WT-MRP1 or vector. For individual experiments, transport was done in triplicate, and then reactions were pooled for analysis by ICP-MS. Bars and symbols represent the means (+S.D.) of at least three independent experiments. (A) Vesicles prepared from HEK293T cells transiently transfected with WT-MRP1 (dark gray bars) or vector (white bars) or HeLa cells stably transfected with WT-MRP1 (black bars) or vector (light gray bars) were incubated for 1 minute at 37°C in transport buffer with MMA(GS)₂ (1 μ M). Statistically significant differences in MMA(GS)₂ transport were determined using a one-way ANOVA followed by a Dunnett's multiple comparisons post hoc test using HEK-WT-MRP1 as the control group (**P < 0.01; ****P < 0.0001). (B) Vesicles prepared from HEK293T cells transiently transfected with WT-MRP1 were incubated for 1 minute at 37°C with increasing concentrations of MMA(GS)₂ (1–200 μ M). Data were fitted using a one-site Michaelis-Menten kinetic model with GraphPad Prism6.

difference in results between the two assays is that DMA^V is poorly taken up by cells (Delnomdedieu et al., 1995; Dopp et al., 2004, 2005; Naranmandura et al., 2007, 2011), including HEK293 cells (Banerjee et al., 2014). Thus, it is likely that the data generated with the inside-out MRP1-enriched membrane vesicles, with no requirement for cell entry, more accurately reflects what is occurring after formation of DMA^V within the cell. Humans are predominantly exposed to As^{III} and As^V in drinking water, which are taken up by cells efficiently (Roggenbeck et al., 2016) and then converted to methylated products (e.g., DMA^V).

As a starting point for determining a mechanism for cell line differences in MRP1-mediated DMAV transport, we investigated the potential contribution of differential phosphorylation. Mutation of two phosphorylation sites (Y920/S921-MRP1), previously reported as responsible for cell line differences in the transport of As(GS)3, to both phospho- or dephospho-mimicking amino acids, surprisingly resulted in a complete loss of DMA^V transport. Individual mutant HEK-Y920F-MRP1 and HEK-S921A-MRP1 membrane vesicles, shown previously to transport As(GS)3 to a similar extent as HEK-WT-MRP1, also completely lacked DMAV transport. The inclusion of a PIC during the preparation of HeLa-WT-MRP1 membrane vesicles did not result in a gain of DMAV transport activity (although MRP1-mediated As(GS)₃ transport was increased), suggesting that either phosphorylation was not important or a stable phosphorylation site (not influenced by the PIC) was involved. Mutation of multiple other putative phosphorylation sites in the linker region also resulted in a complete loss of HEK-MRP1 DMAV transport. Our data suggest that DMAV transport by HEK-WT-MRP1 membrane vesicles is extremely sensitive to alterations in the linker region between NBD1 and MSD2. The reasons for this are currently not understood and require further investigation. Differences in posttranslational modifications and/or protein:protein interactions that alter the structure of this region could potentially explain the cell line differences in MRP1-mediated DMAV transport. Indeed, there is some

suggestion in the literature that this linker region is important for protein: protein interactions and that such interactions may be modulated by phosphorylation (Yang et al., 2012; Ambadipudi and Georges, 2017).

MRP1 transport of DMA^V is the second DMA^V efflux pathway to be identified. Previously, we reported that the related MRP4 transports DMA^V with similar affinity ($K_{0.5}$ 0.22 \pm 0.15 μ M for MRP4 vs. $K_{\rm m}$ 0.19 \pm 0.06 μ M for MRP1), but through a cooperative mechanism (Hill coefficient 2.9 \pm 1.2) and assuming equal protein levels, lower capacity ($V_{\rm max}$ 32 \pm 3 pmol·mg⁻¹ protein·min⁻¹ for MRP4 vs. $V_{\rm max}$ 342 \pm 37 pmol·mg⁻¹ protein·min⁻¹ for MRP1) [(Banerjee et al., 2014, 2016) and Table 2]. The tissue expression and cellular localization of MRP4 likely make it critical for urinary elimination of hepatic metabolites (Banerjee et al., 2014). The localization of MRP1 to the basolateral surface of epithelial cells and expression in specific cell types of most tissues (undetectable protein levels in human hepatocytes), likely makes MRP1 important for cellular/tissue protection rather than playing a role in arsenic elimination.

DMA^V is an arsenic compound with low toxicity relative to trivalent arsenic species (Moe et al., 2016). The efflux of DMA^V from the cell is critical to prevent the reduction of DMA^V to the highly reactive DMA^{III} (Nemeti and Gregus, 2013). Furthermore, export of DMA^V would likely prevent product inhibition of arsenic (+3 oxidation state) methyltransferase, allowing the formation and cellular export of more DMA^V. The reducing intracellular environment might suggest that DMA^{III} is the predominant form of dimethylated arsenic within the cell; however, this has been difficult to prove and DMA^V has been detected in human cell lines and mouse liver homogenate (Currier et al., 2011). The highly reactive DMA^{III} is highly protein bound and unlikely to be available for cellular export (Hippler et al., 2011; Shen et al., 2013). An equilibrium between DMA^{III} and DMA^V will exist within the cell, and the high-affinity high-capacity export of DMA^V by MRP1 would provide a good mechanism for cellular detoxification.

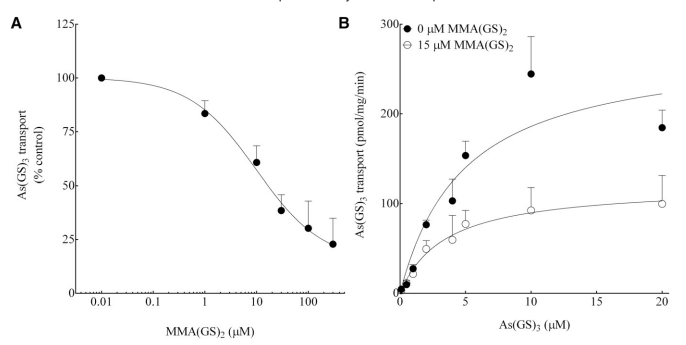


Fig. 5. MMA(GS)₂ inhibits MRP1 mediated ATP-dependent transport of As(GS)₃. Transport experiments were done with membrane vesicles (20 μ g of protein) prepared from HEK293T cells transiently transfected with WT-MRP1. Individual experiments were done in triplicate and ⁷³As quantified using liquid scintillation counting. (A) Vesicles were incubated with ⁷³As(GS)₃ (1 μ M, 40 nCi) for 3 minutes at 37°C in transport buffer with increasing concentrations of MMA(GS)₂ (0.01–300 μ M). Data points represent the mean (+S.D.) of three independent experiments. (B) Vesicles were incubated with increasing concentrations of ⁷³As(GS)₃ (0.1–20 μ M, 40–100 nCi) for 3 minutes at 37°C in transport buffer in the presence of MMA(GS)₂ (15 μ M). Data points represent means (+S.D.) of triplicate determinations in a single experiment. Two independent additional experiments with MMA(GS)₂ at 5 and 10 μ M were done to calculate the K_1 (7.3 ± 5.1 μ M, ±S.D., n = 3).

Out of the five arsenic compounds tested HEK-MRP1 conferred the highest level of resistance against As^V (9-fold) followed by As^{III} (5-fold). Consistent with previous studies using MRP1-enriched membrane vesicles isolated from H69AR cells (Leslie et al., 2004), HEK- and HeLa MRP1-enriched vesicles did not transport As^V in the presence or absence of GSH. Once inside the cell, As^V is reduced to As^{III} and then enters the methylation pathway (Cullen, 2014). Thus, our results are consistent with As^V and As^{III} being converted to As(GS)₃ before efflux by MRP1, as we showed previously (Leslie et al., 2004). The reason why HEK-MRP1 cells confer higher levels of resistance to As^V than As^{III} is not understood. As^V enters cells more slowly (through Na⁺-dependent phosphate transporters) than As^{III} (through aquaglyceroporins) (Mukhopadhyay et al., 2014; Roggenbeck et al., 2016), and this could influence the methylation and glutathionylation of arsenic and alter the metabolites available for MRP1 export.

HEK-MRP1 and HeLa-MRP1 cell lines both conferred significantly higher levels of resistance to MMA^{III} than their respective vector controls [(Carew et al., 2011) and Table 1]. MMA(GS)₂ was transported with comparable apparent affinity by HeLa-WT-MRP1 and HEK-WT-MRP1 membrane vesicles; however, the $V_{\rm max}$ was 3.4-fold higher for HeLa-MRP1 membrane vesicles. Kinetic parameters for MMA(GS)₂ transport were not significantly different between HEK-WT-MRP1 and HEK-Y920F/S921A-MRP1 or HEK-Y920E/S921E-MRP1, suggesting that differential phosphorylation at these sites is not responsible for the cell line differences in $V_{\rm max}$. Consistent with these phosphorylation sites being important for the interaction between MRP1 and As(GS)₃, but not MRP1 and MMA(GS)₂, we found that MMA(GS)₂ was an uncompetitive inhibitor of As(GS)₃ transport. Thus, increasing concentrations of As(GS)₃ did not overcome MMA(GS)₂ inhibition (Fig. 5B), providing support for the idea that As(GS)₃ and MMA(GS)₂ do not share identical binding sites.

Arsenic was previously reported to activate kinase and inhibit phosphatase pathways (Rehman et al., 2012; Beauchamp et al., 2015),

and we had postulated cellular exposure to arsenic would result in a shift to a prophosphorylation state of Y920/S921-MRP1 (Shukalek et al., 2016). This in turn would result in the switch of MRP1 from a high-affinity, low-capacity transporter of As(GS)₃ to a more efficient low-affinity, high-capacity As(GS)₃ transporter (Shukalek et al., 2016). Phosphorylation of these residues appear to be important specifically for As(GS)₃, but not for MMA(GS)₂ or DMA^V (this study) or as previously reported for methotrexate, leukotriene C4, or 17β -estradiol 17-(β -Dglucuronide) (Loe et al., 1996b; Stride et al., 1997; Shukalek et al., 2016). Why phosphorylation of MRP1 at Y920/S921 has an impact on As(GS)3, but not other arsenic metabolites is unknown. Potentially, MRP1 exports As(GS)₃ over a broad concentration range to reduce As III availability for the formation of more toxic trivalent methylated forms. The $K_{\rm m}$ values for As(GS)₃ ($K_{\rm m}$ range ~0.3–4 μ M) (Leslie et al., 2004; Shukalek et al., 2016) and DMA^V ($K_{\rm m}$ 0.19 μ M) (Table 2) are much lower than MMA(GS)₂ ($K_{\rm m}$ range 11-33 μ M) [Table 2 and (Carew et al., 2011)]. At low levels of arsenic exposure, MRP1 is potentially important for the export of As(GS)₃, and any DMA^V that is formed (preventing the formation of the highly toxic DMAIII). During higher cellular arsenic exposure, MRP1 phosphorylation allows it to still export As(GS)3 efficiently; MMA(GS)2 accumulation might start to occur and MRP1 would be able to export this, and potentially DMAV. It is worth noting that the transport of DMAV by MRP1 is remarkably more efficient than reported for any other transporter and arsenical combination (Roggenbeck et al., 2016), providing support for MRP1 being an important transport pathway for DMA^V under environmentally relevant exposure conditions.

Differences in MRP1 transport of DMA^V (this study) and As(GS)₃ (Shukalek et al., 2016) by membrane vesicles isolated from different cells raises the possibility that MRP1 could have a distinct role in arsenic efflux, depending in which tissue and/or cell type it is expressed. MRP1 has been indirectly implicated in the protection of specific tissues from

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arsenic toxicity, including kidney and brain (Kimura et al., 2005, 2006; Dringen et al., 2016; Wang et al., 2016). Furthermore, MRP1 could play a role in resistance to arsenic-based therapies and this could be modified depending upon the tumor type. This study lays the groundwork for further investigation into how the cellular environment influences the function of MRP1, particularly for the cellular detoxification of important arsenic metabolites.

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Authorship Contributions

Participated in research design: Banerjee, Kaur, Whitlock, Carew, Le, Leslie

Conducted experiments: Banerjee, Kaur, Whitlock, Carew.

Contributed new reagents or analytic tools: Le.

Performed data analysis: Banerjee, Kaur, Whitlock, Carew, Leslie.

Wrote or contributed to the writing of the manuscript: Banerjee, Kaur, Whitlock, Leslie.

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