Escape from hsa-miR-519c enables drug-resistant cells to maintain high expression of ABCG2

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Abstract
Overexpression of ABCG2 has been reported in cell lines selected for drug resistance and it is widely believed to be important in the clinical pharmacology of anticancer drugs. We and others have previously identified and validated two microRNAs (miRNA; hsa-miR-519c and hsa-miR-520h) targeting ABCG2. In this study, the shortening of the ABCG2 3′ untranslated region (3′UTR) was found to be a common phenomenon in several ABCG2-overexpressing resistant cell lines, which as a result removes the hsa-miR-519c binding site and its repressive effects on mRNA stability and translation blockade, thereby contributing to drug resistance. On the other hand, reduced expression of hsa-miR-520h, previously thought to have allowed ABCG2 overexpression, was found to be caused by the sequestration of the miRNA by the highly expressed ABCG2. In drug-sensitive cells, inhibitors against hsa-miR-519c and hsa-miR-520h could augment the cytotoxic effect of mitoxantrone, suggesting a substantial role for both miRNAs in controlling ABCG2 level and thereby anticancer drug response. However, in drug-resistant cells, altering the levels of the two miRNAs did not have any effect on sensitivity to mitoxantrone. Taken together, these studies suggest that in ABCG2-overexpressing drug-resistant cells, hsa-miR-519c is unable to affect ABCG2 expression because the mRNA lacks its binding site, whereas hsa-miR-520h is sequestered and unable to limit ABCG2 expression. Given the recent observation that a truncated 3′UTR is also observed in ABCG2-overexpressing human embryonic stem cells, our results in drug-resistant cell lines suggest that 3′UTR truncation is a relatively common mechanism of ABCG2 regulation.

Introduction
ABCG2 is an efflux transporter important in clinical pharmacology and in normal tissue protection (1). ABCG2 was first identified and is often observed in drug-resistant cancer cells (2–6). However, relatively little is known about the mechanisms underlying its upregulation. Most studies of this question have focused on the ABCG2 promoter (7–14). Less is known about its regulation at the 3′ untranslated region (3′UTR) of the gene. MicroRNAs (miRNA) represent a large class of endogenous 19- to 25-nucleotide-long noncoding RNA molecules that are capable of inducing mRNA degradation and/or translational repression by forming imperfect hybrids with the 3′UTR sequences of their target miRNAs (15–17). We showed that ABCG2 mRNA adopts a longer 3′UTR in a parental S1 colon cancer cell line than in its drug-resistant S1M1 80 subline and that a miRNA (hsa-miR-519c; hereafter referred to as miR-519c) decreases endogenous ABCG2 mRNA and protein levels by acting through a putative miR-519c binding site located within the longer 3′UTR region found only in the parental cells (18). These findings suggest that escape from miRNA-mediated translational repression and mRNA degradation could contribute to overexpression of ABCG2 in drug-resistant cancer cells.

miRNAs play important roles in the regulation of basic cell functions, including proliferation, differentiation, and apoptosis (16, 19). Recently, attention has been paid to the involvement of miRNA in the acquisition of drug resistance by cancer cells (20–24). In this study, we sought to examine the ABCG2 3′UTR in various ABCG2-overexpressing resistant cell lines and to assess the functional role of the two previously validated miRNAs (hsa-miR-519c and hsa-miR-520h, hereafter referred to as miR-520h) targeting ABCG2.

Materials and Methods
Tissue Culture
Several pairs of parental and resistant cell lines were used. The human colon cancer cell line S1 and its resistant sublines (S1M1 0.4, S1M1 0.8, S1M1 1.6, S1M1 3.2, S1M1 80, and S1M1 200), obtained by selection in mitoxantrone in a stepwise manner, have been previously described (4, 25). A549, H460, MCF-7, SF295, and SW620 cells were studied along with their drug-resistant counterparts developed by stepwise selection of parental cells in increasing concentrations of anticancer agent. A549 Beca250 (26), H460 MX20...
(27), MCF-7 FLV1000 (5), SF295 MX2000, and SW620 Ad300 (28) were maintained in 250 nmol/l bacitracin, 20 nmol/l mitoxantrone, 1,000 nmol/l flavopiridol, 2,000 nmol/l mitoxantrone, and 300 ng/ml doxorubicin, respectively. ABCG2 is overexpressed in A549 Beca250, H460 MX20, MCF-7 FLV1000, and SF295 MX2000, contributing to drug resistance; no MRPl or P-glycoprotein was detected in these resistant cell lines (5, 27). The SW620 Ad300 subline does not express ABCG2 but has high levels of P-glycoprotein as the mechanism of drug resistance (28). The cell lines were maintained in improved MEM (S1, S1M1 series of resistant sublines, MCF-7, and MCF-7 FLV1000) or RPMI 1640 (A549, A549 Beca250, H460, H460 MX20, SF295, SF295 MX2000, SW620, and SW620 Ad300) supplemented with 10% fetal bovine serum, 100 units/ml streptomycin sulfate, and 100 units/ml penicillin G sulfate and incubated at 37°C in 5% CO2.

**RNA Degradation Analysis**

Parental and resistant cells in 10-cm tissue culture dishes were grown until subconfluent, and then actinomycin D (Sigma-Aldrich) was added to a final concentration of 5 μg/ml to arrest de novo RNA synthesis. At 0, 4, 8, and 16 h after actinomycin D treatment, the cells were harvested and ABCG2 mRNA was quantified by reverse transcription-PCR (RT-PCR) as previously described (18). c-myc and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) mRNA levels were also monitored as controls. The value recorded was the percentage of mRNA remaining compared with the amount before the addition of actinomycin D after normalization with GAPDH. All experiments were repeated thrice.

**Semiquantitative and Real-time RT-PCR**

Semiquantitative RT-PCR was done as previously described (18). Amplification of cDNA was done using the following primers: ABCA8, 5'-GTTGTTGTCACAAGCTTCCTTT-3' (forward) and 5'-ACGCTATGGCTTCCTTGGCTTC-3' (reverse); SMAD6, 5'-AGGAGAATGCTTCTAGGTCCATC-3' (forward) and 5'-TGAGGAAAGCCGCTTCCTTTC-3' (reverse); SMAD7, 5'-AGGAGAATGCTTCTAGGTCCATC-3' (forward) and 5'-TGAGGAAAGCCGCTTCCTTTC-3' (reverse); SMAD9, 5'-AGGAGAATGCTTCTAGGTCCATC-3' (forward) and 5'-TGAGGAAAGCCGCTTCCTTTC-3' (reverse); and SMAD10, 5'-AGGAGAATGCTTCTAGGTCCATC-3' (forward) and 5'-TGAGGAAAGCCGCTTCCTTTC-3' (reverse). GAPDH was used as the internal control.

ABCG2 mRNA expression in the S1M1 series of resistant cell lines was measured by real-time PCR using a LightCycler Fast Start DNA Master Taqman kit (Universal Probe Library; Roche Molecular Biochemicals) as previously described (18).

For the detection of miR-519c and miR-520h, 2 μg of total RNA were used in the RT reaction by using the QuantmiRNA RT kit (System Biosciences). Quantitative real-time PCR was done using the KAPA SYBR FAST qPCR kit (KapaBiosystems) in a LightCycler 480 Instrument I (Roche Applied Science). The mature miR-519c and miR-520h DNA sequence was used as the forward primer, and the 3' universal primer provided from the QuantmiRNA RT kit was used as the reverse primer. The human U6 RNA was amplified in parallel as the internal control. PCRs were done at 95°C for 10 min followed by 40 cycles of 95°C for 10 s and 60°C for 10 s. Fluorescence signal was acquired at the end of the elongation step of every PCR cycle (72°C for 10 s) to monitor the increasing amount of amplified DNA. ΔCt was calculated by subtracting the Ct of U6 from the Ct of miR-519c or miR-520h. ΔΔCt was then calculated by subtracting the ΔCt of the parental cells from the ΔCt of the corresponding resistant cells. Fold change of miRNAs was calculated by the equation 2^-ΔΔCt.

**3' Rapid Amplification of cDNA Ends Assay**

Total RNA was extracted from parental cells and resistant sublines using Trizol reagent (Invitrogen). 3' Rapid amplification of cDNA ends (RACE) assay was done as described previously (a detailed scheme is shown in Supplementary Fig. S1; ref. 18). DNA as a mixture from the PCR was subcloned into a pcR2.1 vector (Invitrogen). Multiple clones were then sequenced by the DNA sequencing core facility at the Laboratory of Experimental Carcinogenesis, National Cancer Institute. All sequences were aligned against reported mRNA sequence for ABCG2 (Genbank accession no. NM_0004872). For S1 and SW620 cells, 3' RACE analysis was done with total RNA harvested from cells after transfection with a miR-519c-specific inhibitor (Dharmacon).

**Cell Transfection with miRNA Inhibitors or Mimics**

miRNA (miRIDIAN) inhibitors or mimics against miR-519c or miR-520h were purchased from Dharmacon. Cells were transfected with up to 100 nmol/l of the specific inhibitors or mimics in a total volume of 6 ml by using DharmaFECT Duo Transfection Reagent (Dharmacon). To evaluate the role of the two miRNAs in the sensitivity of cells to ABCG2 substrate drugs, cells were seeded into 96-well plate for growth-inhibitory assays at 48 h after transfection with the miRIDIAN inhibitors or mimics.

**Semiloop RT-PCR for miRNAs**

Semiloop RT-PCR for evaluating expression of mature miR-519c and miR-520h was done as previously described (18).

**Growth Inhibition Assays**

Mitoxantrone was obtained from Sigma-Aldrich. The growth-inhibitory effect of mitoxantrone under the various miRIDIAN inhibitor/mimic transfection conditions as indicated was tested using the sulforhodamine B proliferation assay (29). At 48 h after transfection with either the miRNA inhibitors or mimics, cells were inoculated into 96-well microtiter plates in 100 μl at plating densities ranging from 5,000 to 20,000 per well depending on the doubling time of individual cell lines. The cells were then treated with mitoxantrone at a range of concentrations and allowed to incubate at 37°C in 5% CO2 for 96 h. Each drug concentration was tested in quadruplicate and controls were tested in replicates of eight. Dose-modifying factors were calculated for the miRNA inhibitor or mimic with mitoxantrone by dividing the IC50 for mitoxantrone without the miRNA inhibitor/
mimic by the IC$_{50}$ for mitoxantrone in the presence of the miRNA inhibitor/mimic. Each experiment was carried out independently at least thrice. To determine whether differences between IC$_{50}$ values were significant, a Student's t test was done with $P < 0.05$ being considered significant.

**Western Blot Analysis**

Whole-cell lysates prepared from S1 and its resistant sublines were separated by SDS-PAGE and subjected to immunoblot analysis with the respective antibodies (ABCG2, Kamiya Biomedical Co.; GAPDH, American Research Products). The blot was analyzed in the Odyssey (LI-COR) IR imaging system after incubation with a 1:10,000 dilution of the goat anti-mouse (IRDye800CW) or anti-rabbit (IRDye680) secondary antibody (LI-COR).

**Flow Cytometry**

Detection of ABCG2-mediated efflux was done by flow cytometry assays as previously described (30). Briefly, cells were trypsinized and incubated in 10 μmol/L phenylboric acid A (PhA) with or without 10 μmol/L of the ABCG2-specific inhibitor fumitremorgin C (FTC) in complete medium (phenol red-free RPMI 1640 with 10% FCS) at 37°C in 5% CO$_2$ for 30 min. Subsequently, the cells were washed once with cold complete medium and then incubated for 1 h at 37°C in PhA-free complete medium continuing with 10 μmol/L FTC to generate the FTC/Efflux histogram, or without FTC to generate the Efflux histogram. Cells were then washed with cold Dulbecco’s PBS and placed on ice in the dark until analyzed.

To determine cell surface expression of ABCG2, cells were trypsinized and resuspended in 2% bovine serum albumin/Dulbecco’s PBS to which was added the phycoerythrin-conjugated anti-ABCG2 antibody 5D3 (eBioscience) or phycoerythrin-conjugated mouse IgG2b negative control antibody (eBioscience). The cells were incubated with antibody for 30 min at room temperature and washed with Dulbecco’s PBS. Surface expression of ABCG2 was calculated as the difference in mean channel number between the 5D3 antibody histogram and the negative control antibody histograms.

Samples were analyzed on a FACSort flow cytometer (Becton Dickinson). Phycoerythrin fluorescence was detected with a 488-nm argon laser and a 585-nm bandpass filter, whereas PhA was detected using a 635-nm red diode laser and a 561-nm filter. At least 10,000 events were collected for all flow cytometry studies. By gating on forward versus side scatter, debris was eliminated, and dead cells were excluded based on propidium iodide staining.

**Fluorescence In situ Hybridization Analysis**

Fluorescence in situ hybridization (FISH) analysis was done using whole chromosome paint kits for chromosome 4 in Spectrum Orange and chromosome 17 in Spectrum Green (Vysis). Earlier studies had been done using an ABCG2 bacterial artificial chromosome probe (>50 kb and containing the 3' end of the gene; National Center for Biotechnology Information Clone Registry ID RP11-368G2; Invitrogen) to localize the ABCG2 gene to normal and rearranged chromosomes. The ABCG2 bacterial artificial chromosome probe localizes to chromosome position 4q22 and has previously been used to study ABCG2 overexpression (31). Standard FISH hybridization procedure was used. All slides were counterstained with 4',6-diamidino-2-phenylindole.

**Results**

**Stability and 3'UTR Length of ABCG2 mRNA in Parental S1 Cells and Its ABCG2-Overexpressing Mitoxantrone-Resistant Sublines**

We previously observed that ABCG2 mRNA in the parental S1 colon cancer cell line was found to adopt different...
lengths at the 3′UTR, with a long (∼2 kb) and multiple shorter (200–400 bp) 3′UTRs, whereas only the shorter ones exist in the resistant S1M1 80 subline (18). As the parental S1 cells underwent stepwise selection with increasing concentrations of mitoxantrone, individual selection steps were maintained at 0.4, 0.8, 1.6, 3.2, 80, and 200 μmol/L of mitoxantrone and named accordingly (4, 25). Overexpression of ABCG2 was observed in the whole series with detection of upregulated mRNA and protein expression in as early as the first resistant step (i.e., S1M1 0.4; Fig. 1A). We asked whether an increase in mRNA stability and shortening of 3′UTR could contribute to the overexpression of ABCG2 in the early and low-resistance steps. Degradation of mRNA was followed by RT-PCR (18) after the cells were treated with actinomycin D to inhibit RNA synthesis. Whereas ABCG2 mRNA in S1M1 0.4, S1M1 1.6, and S1M1 80 was fairly stable up to 16 hours after actinomycin D treatment, ABCG2 in the parental cells had a half-life of ∼6 hours.

Figure 2. A, 3′RACE analysis of ABCG2 mRNA in the S1M1 series of resistant cell lines. The schematics for the 3′RACE assay can be found in Supplementary Fig. S1. The first PCR and the nested PCR 1 gave the same pattern of PCR products. Only the nested PCR 1 is shown. In miR-519c inhibitor–transfected S1 parental cells, a long PCR product (∼2 kb, long arrow) and three much smaller fragments (200, 300, and 400 bp, shorter arrows) were obtained. In all resistant S1M1 sublines, only the smaller fragments were seen. Nested PCR 2 was done to further confirm the existence of the long ABCG2 3′UTR fragment in the parental S1 cells only. B, FISH analysis of chromosome metaphase spreads with whole chromosome paints. Chromosome 4 is painted with WCP 4 Spectrum Orange and chromosome 17 is painted with WCP 17 Spectrum Green; 4′,6-diamidino-2-phenylindole counterstain. S1 parent line: two normal chromosomes 4; S1M1 0.4, S1M1 0.8, S1M1 1.6, S1M1 3.2, and S1M1 80: one normal chromosome 4, one normal 17, and t(4;17)(q21-q22; p13). Earlier studies had shown that the ABCG2 signal is split at the breakpoint (31).
(Fig. 1B). GAPDH and c-myc, controls with long (>24 hours) and short (15–40 minutes) mRNA half-lives, respectively, were found to yield opposite degradation profiles (data not shown), confirming that actinomycin D efficiently stopped transcription in our experiments. These results suggested a reduced degradation rate for ABCG2 mRNA even in the early selection steps.

3′RACE reactions were done to examine the 3′ ends of the ABCG2 mRNA and to define the location of polyadenylated sites in the S1M1 series of resistant sublines (18). The nested PCR 1 gave rise to a long fragment and multiple shorter fragments in the parental S1 cells but only to short fragments in the resistant sublines (Fig. 2A). Nested PCR 2 further confirmed the existence of the long ABCG2 3′UTR in the S1 cells but not in the resistant sublines. There was no significant reaction with samples primed without RT (data not shown).

Chromosomal Rearrangement in the Early Steps of the S1M1 Series and Its Coincidence with the Shortening of ABCG2 3′UTR

The chromosomal rearrangements of the S1 parental and S1M1 80-resistant cells have been characterized (31). A balanced translocation t(4;17)(q21-q22; p13), presumably at the 3′end of the ABCG2 gene, was found in the resistant S1M1 80 subline using FISH and spectral karyotyping (SKY) analysis. It was then hypothesized that the sequences juxtaposed to ABCG2 after the translocation conferred the increased ABCG2 expression in S1M1 80 cells, although the breakpoint has never been characterized. In the present study, the early selection steps (S1M1 0.4, S1M1 0.8, S1M1 1.6, and S1M1 3.2) were assessed for chromosomal translocations as a possible initial event for induction of ABCG2 and the subsequent multidrug resistance phenotype. FISH analysis using whole chromosome paints was done to identify normal and rearranged chromosomes in cytogenetic preparations from the parent S1 and the S1M1-resistant sublines. Interestingly, the same translocation reported in S1M1 80, involving chromosome 4 downstream of the ABCG2 gene, was also observed in all selection steps (Fig. 2B). The parental S1 cells had no chromosome 4 rearrangement and no copy number gain or amplification of chromosome 4. It is notable that the S1M1 80 cells have this gene rearrangement, and despite intense selection pressure in very high concentrations of mitoxantrone, ABCG2 has never undergone gene amplification.

Shortening of the ABCG2 mRNA 3′UTR in Other ABCG2-Overexpressing Resistant Cell Lines

To determine whether 3′UTR truncation was unique to the S1M1 80 cell line, the length of the ABCG2 mRNA 3′UTR in other ABCG2-overexpressing resistant cell lines was examined by 3′RACE assay. Parental S1 and SW620 cells were transfected with specific miR-519c inhibitor to provide higher ABCG2 expression levels for the 3′RACE assay (Fig. 3A). For all parental cell lines tested (S1, MCF-7, SF295, H460, A549, and SW620), 3′RACE assay gave a long 3′UTR fragment and multiple shorter fragments (Fig. 3B, top). However, in the ABCG2-overexpressing resistant sublines (S1M1 80, MCF-7 FLV1000, SF295 MX2000, H460 MX20, and A549 Beca250), only the shorter fragments were obtained. Interestingly, in P glycoprotein–overexpressing SW620 Ad300 cells, where there is no ABCG2 overexpression, the long 3′UTR fragments persist together with the other shorter fragments observed in other resistant cell lines (Fig. 3B). The existence of the long ABCG2 3′UTR in the
parental cells and in the SW620-resistant subline without ABCG2 overexpression was further confirmed by nested PCR (Fig. 3B, bottom). Because the 3′ RACE assay may have missed the long 3′ UTR PCR products because of their lower PCR efficiency, we also conducted RT-PCR experiments by using total RNA from the pairs of parental and resistant cell lines, and primers complementary to different regions in the ABCG2 3′ UTR cDNA. Six separate and overlapping PCR fragments (A–F) that covered the reported ABCG2 3′ UTR sequence (NM_004872) were amplified (Supplementary Fig. S2A; ref. 18). All six PCR fragments were obtained with the predicted size in all RT samples tested, eliminating the possibility of alternate splicing at the ABCG2 3′ UTR. Calibration curves were constructed using a range of concentrations of input genomic DNA from S1 cells. The relative abundance of different 3′ UTR fragments in each cell line, compared with the most upstream region A, was then estimated by reading from the calibration curves. For all parental cell lines tested, as well as the SW620 Ad300 subline without ABCG2 overexpression, the relative abundance of the six 3′ UTR regions was found to be similar (Supplementary Fig. S2B). However, for the ABCG2-overexpressing resistant cell lines, the relative abundance of the more downstream 3′ UTR regions decreased from region A to F, suggesting a higher proportion of truncated 3′ UTR in the resistant cells (Supplementary Fig. S2B).

**Stability of ABCG2 mRNA in Various Parental and Resistant Cell Lines**

Because the various resistant sublines (except the P glycoprotein–expressing SW620 Ad300 subline) lack the long ABCG2 3′ UTR fragment, as reported for S1M1 80 cells (18), the ABCG2 mRNA in these resistant cells may also be more stable than that in the parental counterparts. The half-life of ABCG2 mRNA in this panel of parental and drug-resistant cells was measured after treatment with actinomycin D to inhibit RNA synthesis. Whereas ABCG2 mRNA in the resistant sublines is fairly stable up to 24 hours after actinomycin D treatment, ABCG2 in the parental cells has a half-life of ~12 hours (except in S1 where the half-life is ~8 hours), which is comparable with the number reported by Nakanishi et al. (12) for MCF-7 cells. Thus, the reduced degradation rate of ABCG2 mRNA may have contributed to the upregulation of ABCG2 in these resistant cells. In resistant P glycoprotein–overexpressing SW620 Ad300, ABCG2 mRNA stability was found to be similar to that in the parental cells (Fig. 4).

**Sequence Analysis of ABCG2 3′ UTR with Different Lengths**

We cloned the 3′RACE PCR mixture as a whole (first PCR) into the TOPO vector and subjected the resulting clones to DNA sequencing. ABCG2 3′ UTR transcripts of five different lengths were found. Consistent with the 3′ RACE assay, whereas we found a long (~2 kb) and a few much shorter (100–400 bp) 3′ UTR transcripts in the parental cells, only the shorter ones were found in the resistant cells. A summary of the 3′ UTR ends found for the various pairs of parental and resistant cell lines is tabulated in Supplementary Fig. S3. The numbering of nucleotides at the 3′ UTR is relative to the first nucleotide in the ABCG2 transcript designated in NM_004827. The sequence of the 3′ UTR was identical to the full-length sequence except for the early termination, indicating that multiple site polyadenylation rather than alternate splicing was involved.

**Reduced Levels of miR-520h Found in ABCG2-Overexpressing Resistant Cells**

As discussed previously, miR-520h binds to both long and short ABCG2 3′ UTR in parental S1 and resistant S1M1 80 cells, whereas miR-519c binds to only the long ABCG2 3′ UTR in the parental S1 cells (18). We therefore examined the expression level of these two miRNAs by quantitative real-time PCR in the various parental and resistant cells to
try to elucidate their role in regulating ABCG2. Interestingly, the levels of miR-520h in the resistant cells (except SW620 Ad300) were reduced 2- to 3-fold relative to those in the respective parental cells (Fig. 5). In contrast, miR-519c levels in the resistant cells were found to be slightly higher in the resistant cells than in the parental cells.

High Levels of ABCG2 in Resistant Cells Serve as a Sink for miR-520h in the Resistant Cells

Although the reduction in miR-520h expression in the resistant cells may mediate the overexpression of ABCG2, we also considered an alternate possibility—that the high levels of ABCG2 could serve as a sink for miR-520h in the resistant cells, yielding the same observation. ABCG2 was thus knocked down by transfecting parental MCF-7 and resistant MCF-7 FLV1000 cells with a pool of four small interfering RNA duplexes specifically designed against ABCG2 (ON-TARGETplus small interfering RNA, Dharmacon). ABCG2 was knocked down significantly by 90% and 60% in MCF-7 and MCF-7 FLV1000, respectively (Supplementary Fig. S4). Whereas the miR-519c level was not affected by ABCG2 small interfering RNA transfection, the miR-520h level increased >2-fold in the ABCG2-silenced MCF-7 FLV1000 cells. The effect of lower endogenous miR-520h levels in the resistant cells on four of its putative target genes (BRCA2, SMAD6, ABCA8, and ID1a), predicted by the miRDB miRNA target prediction database (32), was also evaluated by RT-PCR (Supplementary Fig. S5). Compared with the parental MCF-7 cells, no consistent increase in these putative miR-520h target genes was observed in the resistant MCF-7 FLV1000 cells despite the low levels of miR-520h in these cells.

Effects of miRNA Silencing and Forced Expression on Sensitivity of Cells to the ABCG2 Substrate Mitoxantrone

We previously showed that resistant cells could have overexpressed ABCG2 by escaping repression mediated by miR-519c. To further assess the intrinsic effects of miR-519c and miR-520h on sensitivity toward ABCG2 substrate drugs, we did cytotoxicity assays using sulforhodamine B to compare the growth of S1 and MCF-7 cells in which the endogenous pool of miR-519c or miR-520h was altered by transfection with synthetic miRIDIAN inhibitor or mimic (Dharmacon). Cells were seeded into 96-well plates for cytotoxicity assay, cell surface ABCG2 expression and PhA toxicity assay (Fig. 6; Supplementary Fig. S7). On the other hand, miRNA mimics against either miR-519c or miR-520h did not have any appreciable effect on the expression or function of ABCG2 in either S1 or MCF-7 cells (Fig. 6; Supplementary Fig. S7).

ABCG2 mRNA expression and efflux function were also induced, to varying extent, in other parental cell lines on transfection with miR-519c or miR-520h inhibitors (Supplementary Fig. S8A and B).

Discussion

High levels of ABCG2 expression are found in normal tissues involved in xenobiotic protection, in stem cells, and in drug-resistant cancer cell lines (2–6). However, little is known about the mechanisms underlying its upregulation, especially at the 3′UTR. Previously, we reported that the ABCG2 mRNA is more stable in the mitoxantrone-selected and ABCG2-overexpressing resistant subline SIM1 80 compared with the ABCG2 mRNA in S1 parental cancer cell line (18). Our results suggested that miR-519c binds to the 3′-untranslated region of the ABCG2 transcript and reduces its stability, leading to the overexpression of ABCG2. The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6). The data from quadruplicate samples were analyzed for significant differences by the paired Student’s t test (Supplementary Fig. S6).

Figure 5. Differential miR-519c and miR-520h expression levels between the parental and ABCG2-overexpressing resistant cells. Expression of the two miRNAs was measured by quantitative real-time PCR. U6 small RNA was used as the internal control. Columns, mean from three independent and reproducible experiments; bars, SD.
Figure 6. A, RT-PCR analysis of ABCG2 and MDR1 mRNA expression in S1 and S1M1 80 cells transfected with specific inhibitor/mimic against miR-519c or miR-520h. GFP from cotransfection was also measured to assure sufficient transfection efficiency. ABCG2 expression was increased by miR-519c and miR-520h inhibitor in the parental S1 cells, but no change was recorded in the resistant S1M1 80 cells. B, ABCG2 surface expression and FTC-inhibitable PhA efflux in miRNA inhibitor or mimic-transfected S1 cells. Left column, cells were incubated for 30 min with either phycoerythrin-labeled negative control antibody (solid lines) or the phycoerythrin-labeled anti-ABCG2 antibody 5D3 (dashed lines) according to the manufacturer’s instructions. Right column, transfected cells were incubated in PhA with or without 10 μmol/L FTC for 30 min at 37°C, washed, and then allowed to efflux for 1 h at 37°C in substrate-free medium continuing with (dashed lines, FTC/Efflux histogram) or without (solid lines, Efflux histogram) FTC. Representative histograms are shown. No change in ABCG2 expression or PhA efflux was observed in S1M1 80 cells on the miRNA inhibitor or mimic transfection (data not shown).

UTR of ABCG2 in the S1 cells, thereby suppressing ABCG2 expression by either mRNA degradation or protein translation repression, but that the binding site for this miRNA is lost in the shorter 3’UTR present in the resistant S1M1 80 cells. Similar findings have recently been reported in an undifferentiated human embryonic stem cell line where its high ABCG2 expression was associated with only the short 3’UTR variant forms (34). In contrast, another differentiated human embryonic stem cell line with lower ABCG2 levels possesses the longer 3’UTR variant (34). Sandberg et al. (35) also found that rapidly proliferating cells express ABCG2 mRNAs with shorter 3’UTRs, presumably to escape miRNA regulation. In the present study, we examined this phenomenon in additional cancer cell lines to gain further insight into this mechanism of ABCG2 gene overexpression. The importance of ABCG2 and other drug transporters in oncology has not yet been clearly established. Certainly overexpression of ABCG2 and P-glycoprotein in clinical samples has been linked with poor prognosis in multiple settings (36, 37). However, the addition of inhibitors to therapeutic regimens did not improve outcome in multiple studies. This left transporters as therapeutic targets an unanswered question that has been renewed in recent years with the discovery that most of the tyrosine kinase inhibitors currently in clinical use are substrates or inhibitors of the transporters (1). Because the tyrosine kinase inhibitors are being tested in combination with chemotherapy, a method to confirm whether ABCG2 confers resistance to them, or is inhibited by them, or both is exceedingly relevant.

Earlier steps of selection in the S1M1 series were examined, and stabilization of ABCG2 mRNA and shortening of the 3’UTR were observed in all the earlier steps (Figs. 1B and 2A). We previously reported that the effect of miR-519c on inhibiting ABCG2 protein translation is more pronounced than that promoting mRNA degradation probably because of imperfect binding of miR-519c to its binding site on ABCG2 3’UTR (18). When we review the S1M1 series of resistant sublines, the increase in ABCG2 protein expression is more dramatic than that in ABCG2 mRNA levels (Fig. 1A). This lack of proportionality is consistent with a differential effect of the miRNA on mRNA stability and protein translation. Taken together, our results suggest that loss of miR-519c binding may be involved in the initial step of selection when these sublines acquired multidrug resistance, prompting a marked overexpression of ABCG2.

miR-520h has been reported to target ABCG2 in hematopoietic stem cells during their differentiation into progenitor cells (38). Unlike miR-519c, miR-520h binds to ABCG2 3’UTR in both parental and resistant cells. We found that ABCG2-overexpressing resistant cells have lower levels of miR-520h than their parental counterparts (Fig. 5). Interestingly, this is likely caused by the sequestration of miR-520h by the highly expressed ABCG2 in the resistant cells (Supplementary Fig. S4).

A balanced t(4;7) translocation downstream of the ABCG2 gene has been reported for S1M1 80 cells (31). Coincident with the shortening of the ABCG2 3’UTR in the early S1M1-resistant sublines, the same translocation was also found in these sublines starting from S1M1 0.4. Because the breakpoint for this translocation is believed to be 3’ to the ABCG2 gene, it remains to be determined if there is any correlation between the translocation and the differential 3’UTR length adopted in resistant cells. Although this chromosomal rearrangement was not identified in several other drug-resistant cell lines examined and was considered to be
a unique event (31), the possibility that this rearrangement could be found in other tumors cannot be excluded. Indeed, there is increasing evidence that fusion genes are important even in nonhematologic tumor types such as the EML4-ALK fusion gene recently identified as an important target in a subset of lung cancer (39, 40). Further investigation is warranted in other ABCG2-overexpressing resistant cancer cell lines and in primary clinical samples.

The concurrence of the 3′ translocation, distinct histone modifications at the promoter (14), and the shortening of 3′ UTR of ABCG2 in the resistant sublines, although possibly coincidental, suggests an interaction of multiple mechanisms of gene regulation. Intriguingly, besides the S1M1 series of resistant cells, all the ABCG2-overexpressing resistant human cancer cell lines tested in this study exhibited the absence of the long ABCG2 3′ UTR, effectively removing miR-519c-mediated repression on ABCG2 (Fig. 3B) and stabilizing the ABCG2 mRNA (Fig. 4). In contrast, the long ABCG2 3′ UTR is retained in the doxorubicin-resistant cell line derived from the SW620 colon cancer cell line (Ad300) without the overexpression of ABCG2 (Fig. 3B).

Although our study has focused on ABCG2 overexpression due to drug selection, it is possible that the same regulatory mechanism identified (i.e., stabilization of mRNA due to truncation at the 3′ UTR) could apply to other cells with high endogenous ABCG2 levels, as observed in the undifferentiated human embryonic stem cell line (34). To date, however, we have not confirmed ABCG2 3′ UTR and mRNA stabilization as a generalized mechanism contributing to intrinsic drug resistance due to high endogenous ABCG2 expression. None of the unselected cell lines with relatively high ABCG2 expression that we have examined to date (MCF-7, SF295, A549, H460, RPMI-8226, or KM12; 41) has had the truncated phenotype. Rather, ABCG2 adopts both a long and several short 3′ UTR fragments (Supplementary Fig. S1B). However, 3′ UTR length has not been examined in tumor samples with ABCG2 overexpression nor in putative cancer stem cells.

During the review of this manuscript, Pan et al. (42) identified another miRNA (miR-328) targeting ABCG2 3′ UTR. They also observed an inverse relation between the levels of miR-328 and ABCG2 in the parental MCF-7 and the resistant MCF-7/MX100 cells. Interestingly, the binding site for miR-328 on ABCG2 3′ UTR is also located in the long 3′ UTR species identified only in parental cells in our study. Although Pan et al. did not evaluate the ABCG2 3′ UTR length in the parental and resistant cells, we speculate that ABCG2 3′ UTR also adopts a long form in their resistant cell line model. Therefore, the interplay between differential miRNA expression and ABCG2 3′ UTR length could determine the precise mechanism by which ABCG2 is overexpressed in a particular resistant cell line.

Evidence pointing to the role of miRNAs in determining drug sensitivity and resistance is steadily emerging. Blower et al. (43) reported significant correlations in the NCI-60 tumor drug screen panel between miRNA expression patterns and potency patterns for a large set of compounds. It has been shown that a genetic polymorphism of the miR-24 miRNA binding site in dihydrofolate reductase gene could result in resistance to methotrexate (44). The upregulation of miR-214 has been shown to promote the survival of ovarian cancer cells and to induce cisplatin resistance through targeting the 3′ UTR of PTEN, which leads to downregulation of PTEN protein and activation of Akt pathway (45). More recently, it has been shown that the increased expression of miR-27a and miR-451 contributes to the multidrug resistance phenotype in A2780 human ovarian cancer cells (21), although the precise mechanism is not clear. In another study, miR-451 was validated as a true miRNA targeting MDR1 in MCF-7 cells (22).

We evaluated the role of miR-519c and miR-520h in affecting ABCG2 expression, function, and resistance to mitoxantrone. Unselected cells transfected with inhibitors against both miR-519c or miR-520h showed increased ABCG2, PhA efflux, and mitoxantrone resistance on removal of the repressive effect by the two miRNAs (Fig. 6; Supplementary Figs. S6 and S8). Although the studies raise the possibility of miRNA manipulation as a novel strategy to alter drug sensitivity, the experiments do show the limitations of such a strategy. The miRNA mimics do not reduce ABCG2 expression or function where levels are already low. Furthermore, neither miRNA inhibitor nor mimic affects expression in resistant SIM1 80 and MCF-7 FLV1000 cells. This may indicate that although the miRNAs are able to control low levels of ABCG2 mRNA, they cannot have a measurable effect on the very high levels found in drug-resistant cell lines.

Although reducing drug resistance through increasing miRNA expression is not achievable at this writing, the possibility that it could open new avenues of investigation. For example, we need to know whether the 3′ UTR truncation is observed in any drug resistance setting. Drug-selected cell lines, with exceedingly high expression of an ABC transporter, do not faithfully reproduce clinical drug resistance. We need to evaluate the 3′ UTR in clinical samples. Given the observation that both undifferentiated stem cells and proliferating cells have truncated 3′ UTR as part of their phenotype, it seems rational that tumors with ABCG2 overexpression may have truncated 3′ UTR as well. Identifying truncation would be convincing evidence of activation of the transporter in that tumor type. In the event that ABCG2 3′ UTR truncation can be confirmed in specific tumor types, methods of reversing the truncation will need to be developed. Because a genetic change is very unlikely as a cause, the shortening must be epigenetic in origin, and this leads to the possibility that an epigenetic therapy could be identified to modulate the length of the 3′ UTR. With the increasing numbers of epigenetic therapies on the horizon, the possibility is an intriguing one (46).

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

References


Escape from hsa-miR-519c enables drug-resistant cells to maintain high expression of ABCG2
