Function of a Subunit Isoforms of the V-ATPase in pH Homeostasis and In Vitro Invasion of MDA-MB231 Human Breast Cancer Cells

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Running title: Function of V-ATPase a subunit isoforms in invasion

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It has previously been shown that highly invasive MDA-MB231 human breast cancer cells express V-ATPases at the cell surface whereas the poorly invasive MCF7 cell line does not. Bafilomycin, a specific V-ATPase inhibitor, reduces the in vitro invasion of MB231 cells but not MCF7 cells. Targeting of V-ATPases to different cellular membranes is controlled by isoforms of subunit a. mRNA levels for a subunit isoforms were measured in MB231 and MCF7 cells using quantitative RT-PCR. The results show that although all four isoforms are detectable in both cell types, levels of a3 and a4 are much higher in MB231 than in MCF7 cells. Isoform-specific siRNAs were employed to selectively reduce mRNA levels for each isoform in MB231 cells. V-ATPase function was assessed using the fluorescent indicators SNARF-1 and pyranine to monitor the pH of the cytosol and endosomal/lysosomal compartments, respectively. Cytosolic pH was decreased only on knock-down of a3, whereas endosome/lysosome pH was increased on knock-down of a1, a2 and a3. Treatment of cells with siRNA to a4 did not affect either cytosolic or endosome/lysosome pH. Measurement of invasion using an in vitro transwell assay revealed that siRNAs to both a3 and a4 significantly inhibited invasion of MB231 cells. Immunofluorescence staining of MB231 cells for V-ATPase distribution revealed extensive intracellular staining, with plasma membrane staining observed in approximately 18% of cells. Knock-down of a4 had the greatest effect on plasma membrane staining, leading to a 32% reduction. These results suggest that the a4 isoform may be responsible for targeting V-ATPases to the plasma membrane of MB231 cells and that cell surface V-ATPases play a significant role in invasion. However, other V-ATPases affecting the pH of the cytosol and intracellular compartments, particularly those containing a3, are also involved in invasion.

The leading cause of mortality from cancer is metastasis, making inhibition of metastasis an important strategy in controlling cancer progression. The metastatic cascade involves a series of steps that include escape of cells from the site of the primary tumor into the circulation or lymphatic system and the extravasation of cells from the circulation or lymphatic system into secondary sites (1,2). Both of these processes require the tumor cells to display an invasive phenotype in which they degrade extracellular matrix of the surrounding tissue. The vacuolar H⁺ ATPases (or V-ATPases) are a family of ATP-dependent proton pumps that have been implicated in tumor cell invasion (3). V-ATPases are upregulated in tissue samples from highly invasive pancreatic carcinomas (4) and treatment of a human cancer cell line with antisense oligonucleotides to the V-ATPase c subunit decreased invasion in vitro (5). Inhibition of V-ATPase expression in hepatocellular carcinoma cells using siRNAs reduces invasiveness of these cells in vitro and metastasis in vivo (6). In addition, treatment of highly invasive MDA-MB231 human breast cancer cells with the
specific V-ATPase inhibitors bafilomycin or concanamycin inhibits the invasion of these cells in vitro (7).

How might V-ATPases function in tumor cell invasion? Because V-ATPases are present in both intracellular compartments and the plasma membrane, there are a number of possibilities. Plasma membrane V-ATPases are poised to acidify the extracellular environment and have been shown to be active in transport of protons across the plasma membrane of a number of tumor cell types, including MB231 cells (7-9). Extracellular acidification is required for activation of secreted cathepsins, proteases that normally reside in lysosomes (10), but which have been shown to be secreted by a variety of cancer cells and to be important for tumor cell invasion (11). Once in the extracellular space, activated cathepsins can both degrade extracellular matrix proteins and activate other secreted proteases involved in invasion, such as matrix metalloproteases (MMPs) (11,12). Intracellular V-ATPases may also function in invasion by either aiding in the proteolytic activation of cathepsins or MMPs within lysosomes or secretory vesicles or by assisting in the trafficking of vesicles containing these proteases to the cell surface.

To begin to unravel the role of V-ATPases in tumor cell invasion, we have investigated the role of particular isoforms of the V-ATPase in the invasive phenotype of a human breast cancer cell line. V-ATPases are multisubunit complexes composed of a peripheral domain (V₁) responsible for ATP hydrolysis and an integral domain (V₀) that carries out proton transport (3). The core V₀ domain in mammals contains five subunits (a,d,c,c” and e), with the information necessary to target the V-ATPase to different cellular membranes located in the a subunit (13-19). Subunit a is a 100 kDa integral membrane protein containing a 50 kDa hydrophilic N-terminal domain located on the cytoplasmic side of the membrane and a 50 kDa C-terminal domain containing multiple membrane spanning segments (20). Targeting information is localized to the cytoplasmic N-terminal domain (21). In mammals subunit a exists in four isoforms (a1-a4) (15-19,22-24). The a1 and a2 isoforms appear to localize primarily to intracellular compartments, with a1 present in synaptic vesicles (25) and clathrin-coated vesicles in brain, while a2 is present in Golgi and endosomal compartments (18,26). The a3 isoform is responsible for targeting the V-ATPase to the plasma membrane of osteoclasts, where it plays an essential role in bone resorption (18,19,23,27), although a3 has also been localized to lysosomes and insulin-containing secretory vesicles in pre-osteoclasts (18) and pancreatic islet cells (17), respectively. The a4 isoform functions to target V-ATPases to the apical membrane of renal intercalated cells, where they are responsible for acid secretion into the urine (15,16,24). a4 has also been localized to the apical membrane of epididymal clear cells (28).

The purpose of this study was to determine the a subunit isoform expression profile of the human breast cancer cell line MDA-MB231 and to determine the effect of reducing expression of each of the four a subunit isoforms on V-ATPase function and the invasiveness of these cells in vitro. The results suggest distinct functions of a subunit isoforms in control of cytosolic and endosome/lysosome pH and a role for both intracellular and plasma membrane V-ATPases in tumor cell invasion.

**EXPERIMENTAL PROCEDURES**

**Cell Culture**

Human breast cancer cell lines, MDA-MB231 and MCF7, were purchased from American Type Culture Collection (ATCC). Cells were grown in Falcon™ T-75 flasks in Dulbecco’s Modified Eagle Medium (DMEM) with phenol, 4.5 g/L D-glucose, 4 mM (584mg/L) L-glutamine, 10% FBS (Gibco), 60 mg/L sodium pyruvate (Gibco) supplemented with 10% FBS (Gibco), MEM non-essential amino acids (Gibco), 60 μg/ml penicillin and 125 μg/ml streptomycin (Gibco). Cells were grown in a 95% air/5% CO₂ humidified environment at 37°C.

**Real Time Reverse Transcription PCR**

Cells were harvested, lysed and RNA was isolated using Qiagen’s RNeasy® Mini Kit. After RNA isolation, mRNA was isolated with the MicroPoly(A) Purist™ Kit from Ambion®. Total RNA or mRNA concentration was quantified using Quant-iT RiboGreen® RNA reagent (Molecular Probes™). One step quantitative RT-
PCR was performed in a 96 well format on a Stratagene MX-3000P® QPCR system using Brilliant® SYBR® Green QRT-PCR Master Mix Kit (Stratagene). The PCR cycling sequence consisted of 30 minutes at 50°C to allow for reverse transcription, then a heat inactivation and denaturation step for 10 minutes at 95°C; this was followed by 40 cycles of 30s at 95°C, 1 min at 55°C, and 30s at 72°C to allow for denaturation, annealing and extension, respectively. To quantitate the results, cDNA clones of the a1, a2 (ATCC), a3, and a4 (Open Biosystems) isoforms were purchased. The cDNA sequences were verified by sequencing. Plasmid DNA for each isoform isolated from E. coli was quantitated by measuring the OD at 260 nm, serial dilutions were made and these DNA standards were used during the quantitative RT-PCR reactions to facilitate quantification of the initial mRNA levels for each experimental sample by use of a standard curve.

RNA Interference

siRNA pools specific for the a1, a2, a3 or a4 isoform were purchased from Dharmacon. Each pool contained four siRNAs specific for the appropriate a subunit isoform. MB231 cells were plated in 60mm dishes at 4X10⁵ cells/dish in the media described above without antibiotic and incubated overnight. Cells were transfected with 20 nM siRNA directed against a1 or a2, 100 nM siRNA directed against a3 or 10 nM siRNA directed against a4 according to the manufacturer’s directions. Briefly, the siRNA was mixed 1:1 with OPTI-MEM™ (Gibco), allowed to incubate for 5 minutes and then mixed with transfection reagent (Dharmafect2™, Dharmacon). The siRNA/transfection reagent mix was incubated for 20 minutes at room temperature and then added to the appropriate volume of DMEM + serum and 4 ml was added to each dish. After incubation of cells with siRNA for 24 hrs the media was changed to siRNA-free media and cells were incubated for additional time (depending upon the assay) prior to harvesting. For all experiments, data were collected 96 hrs post-transfection. To quantify reduction in the a subunit isoform mRNA levels, quantitative RT-PCR was performed as described above using RNA isolated from cells after siRNA treatment. To confirm the specificity of knockdown, we used primers specific for the isoforms that were not targeted by the siRNA treatment.

Loading cells with fluorescent pH indicators

To evaluate the localization of fluorescent pH indicators, cells were grown onto round coverslips (25 mm) in DMEM supplemented with FBS. After 48 hrs in culture, cells were incubated for 12 hrs with 1 mM 8-hydroxypyrene-1,3,6-trisulfonic acid (pyranine), a pH fluorescent indicator, in DMEM supplemented with FBS. Cells take-up pyranine by pinocytosis which effectively labels endosomal and lysosomal compartments (29). The cytosol was loaded with 7 µM SNARF-1, a pH fluoroprobe. This probe is internalized by incubating cells for 30 min with the acetoxymethyl (AM) ester form of SNARF-1(30). Cellular esterases cleave the AM groups leaving the charged form of SNARF-1 in the cytosol. Fluorescence images of cells were obtained using 20X objective and a digital camera (Orca, Hamamatsu). Cells loaded with SNARF-1 were excited at 534 nm and the fluorescence emission signal collected at 590 nm (long bandpass filter).

Cytosolic and Endosomal/Lysosomal pH Measurements

To verify the usefulness of pyranine and SNARF for in vivo pH measurements, 1 µM pyranine and 2 µM SNARF-1 (free acid) were dissolved in high
K+ buffers of increasing pH (from 5.5 to 8.5) and their spectral properties were evaluated using a fluorometer. The excitation spectra of pyranine were collected using an emission wavelength of 514 nm and the emission spectra of SNARF-1 were acquired using 534 nm as excitation wavelength. The significant properties of pyranine are the increase in the excitation peak at 465 nm and the decrease at 405 nm as pH is increased from 5.5 to 8.5. The significant properties of SNARF-1 are the increase in the emission signal at 644 nm and the decrease at 584 nm as pH is increased. Consequently, the fluorescence ratios at 465/405 nm and 644/584 nm can be used to monitor pH in endosomes/lysosomes and cytosol, respectively using a ratiometric approach that allows us to quantitate pH.

To evaluate the effect of siRNA treatment on steady state cytosolic and endosome/lysosome pH, cells growing on rectangular coverslips (8 x 22 mm) were treated with siRNA directed against the different α isoforms for 24 hrs as described above. 84 hrs post-transfection, cells were incubated with pyranine and/or SNARF-1 for 12 hrs as described above. Following incubation, for each experiment, 2 coverslips containing cells were washed, to remove external dye, and then transferred to a thermostated fluorescence chamber kept at 37°C and continuously perfused at 3 ml/min. The excitation spectra of pyranine in endosomes/lysosomes was acquired using an emission wavelength of 514nm and the emission spectra of SNARF-1 were acquired using 534 nm as excitation wavelength and the conversion of ratio values to pH\text{\textsuperscript{cyt}} and pH\text{\textsuperscript{endo/lys}} were performed.

**Invasion Assay**

Assays for in vitro invasion were performed using ChemoTx\textsuperscript{®} membranes (Neuroprobe) with an 8\textmu m pore size membrane and coated with Matrigel\textsuperscript{™} (BD Biosciences). A chemoattractant (FBS) was added to the trans-side of the membrane to induce invasive cells to digest the coating and migrate through the pores to the trans-side. Matrigel\textsuperscript{™} was diluted in PBS to a final concentration of 0.3 \mu g/\mu l and a total of 4\mu g was coated onto the membrane in each well. The membrane was allowed to dry overnight under vacuum at room temperature. Matrigel\textsuperscript{™}-coated membrane was re-hydrated with 50 \mu l DMEM + 4.5 g/L D-glucose, without phenol, L-glutamine or sodium pyruvate (Gibco) (termed Media) for at least two hours. 30 \mu l of the Media containing 5% FBS were added to the wells of the complementary ChemoTx\textsuperscript{®} plate to act as the chemoattractant. 72hrs post-transfection, cells were harvested to set-up the invasion assay. To ensure that only viable cells were loaded onto the membrane, cells were washed to remove unattached cells and then harvested via trypsinization. Viability of the harvested cells was assessed by trypan blue exclusion; less than 1% of the harvested cells stained with trypan blue. Cells were next brought to a concentration of 2X10\textsuperscript{5} cells/ml in Media that did not contain siRNA. For inhibitor-treated samples, cells were resuspended in Media containing either DMSO or 100 nM concanamycin A in DMSO and allowed to incubate for 15 minutes at 37°C. 5X10\textsuperscript{4} cells were seeded onto the rehydrated membrane which was then fitted onto the wells containing chemoattractant. The invading cells were incubated under normal growth conditions. After 24 hrs, the non-invaded cells and Matrigel\textsuperscript{®} were scraped from the top of the membrane using a cell scraper and the membrane with the invaded cells was fitted onto wells containing 30 \mu l 4% paraformaldehyde and incubated for 10 min at room temperature to fix the cells. The membrane was then fitted onto wells containing 30 \mu l of 4 \mu g/ml propidium iodide in warm PBS to stain the cells. The cells were incubated with propidium iodide for 30 minutes at 37°C and then moved to a Zeiss Axiovert 10 fluorescence microscope for analysis. Thirteen images per well were taken of the trans-side of the membrane. The number of invaded cells per well was counted from the images. Each condition was performed in triplicate and the number of invading cells averaged over the three wells. Percent invasion for treated cells was normalized to non-treated cells.

**Immunocytochemistry**

A polyclonal antibody against the V-ATPase subunit E (provided by Dr. Reuveni) was used to determine the in situ localization of V-ATPases (31). MB231 cells were plated onto 60mm dishes containing poly-D-lysine coated 12mm round
coverslips. Approximately 24 hrs later the cells were treated with siRNA for 24 hrs. The media was then changed to siRNA-free media and the cells were allowed to incubate for an additional 72 hours. 96hrs post siRNA transfection, the cells on coverslips were retrieved. Using a 200 µl pipette tip, the confluent monolayer was scratched to create a wound to induce migration. Wounding was performed because it has previously been shown that cells utilizing V-ATPases for pH regulation and migration target them to the plasma membrane at the leading edge (31). Cells were then incubated for an additional 4 hrs, washed, fixed with 4% paraformaldehyde, and permeabilized with 0.1% Triton X-100. The cells were blocked with 1% BSA in PBS for 1 hr and then incubated with anti-E antibody at a 1:500 dilution overnight. The cells were then rinsed with PBS and then treated with Alexa Fluor® 488 conjugated goat anti-rabbit secondary (1:500) and Alexa Fluor® 594 phalloidin (to stain F-actin, 1:250) (Invitrogen) in 1% BSA-PBS were added to the cells. After 1 hr incubation at room temperature, the cells were rinsed with PBS. The cells were prepared for viewing using ProLong® Gold (Invitrogen) mounting medium which was allowed to cure at room temperature for 24 hrs. The fluorescent cells were imaged on either the Zeiss Axiovert 10 fluorescence microscope or a Leica TCS SP2 confocal microscope. To quantitate the appearance of plasma membrane staining, 30-40 consecutive images were taken along the wound. For each image, the total number of cells was counted and the number showing a distinct line of staining at the plasma membrane was counted. The ratio of cells with plasma membrane staining versus the total number of cells was calculated.

**Cathepsin L Secretion**

Cells plated in 60 mm dishes were treated with siRNA and allowed to incubate for 24 hrs at 37°C. Media was replaced with siRNA-free media and cells were incubated for an additional 48 hrs The media was then replaced with serum-free media and cells were incubated for an additional 24 hrs. 96 hours post-transfection the conditioned media from each dish was recovered and concentrated 80-fold using an Amicon® Ultra (4ml, 10K cut-off, Millipore). Samples of the concentrated media for each condition were resolved by SDS PAGE and blotted for Cathepsin L using a monoclonal antibody from Zymed Laboratories.

**Statistical analysis**

All results are expressed as means +/- standard deviation. The significant differences were determined by ANOVA for each experiment, followed by pairwise t-test to compare individual treatments to non-treated. All statistical tests were considered significant at P< 0.05.

**RESULTS**

**mRNA levels of a subunit isoforms in MDA-MB231 and MCF7 cells as measured using quantitative RT-PCR.** Because of the role V-ATPases have been reported to play in invasiveness of breast cancer cells, it was of interest to compare the mRNA expression profiles for a subunit isoforms in the highly invasive breast cancer cell line MDA-MB231 with those of the poorly invasive cell line MCF7. Quantitative RT-PCR was performed with isoform-specific primers on mRNA isolated from each cell line, using plasmid-borne cDNAs encoding each isoform as standards. As can be seen in Fig.1, although all four a subunit isoforms can be detected in both cell lines, their relative abundance differs dramatically between MB231 and MCF7 cells. Thus the predominant a subunit isoform in MB231 cells is a3 whereas the predominant isoform in MCF7 cells is a1, followed closely by a2. Comparing the ratio of expression of each isoform between the two cell lines (Fig.1C) reveals that both the a3 and a4 isoforms are expressed at much higher levels in MB231 than MCF7 cells.

**Selective knockdown of each subunit a isoform mRNA using siRNA.** In order to evaluate the role of a subunit isoforms in V-ATPase function and invasiveness, we employed isoform-specific siRNAs to reduce the level of each a subunit isoform mRNA in MB231 cells. The effect of siRNA treatment on mRNA levels for each isoform was assessed using quantitative RT-PCR. mRNA was isolated from cells treated with siRNA for 24 hrs followed by 72 hrs of post-treatment incubation. As shown in Fig.2, greater than 50% reduction in mRNA levels was observed for each a subunit isoform. Moreover, reductions in mRNA
levels were isoform-specific, although it should be noted that there does appear to be some compensatory up-regulation of the a1, a2 and a3 mRNA levels upon treatment with siRNA to a4. These results indicate that the siRNA treatment is effective at selectively knocking down mRNA levels for each a subunit isoform.

**Effect of a subunit isoform reduction on cytoplasmic and endosomal/lysosomal pH in MB231 cells.** To investigate the effect of reducing each subunit a isoform on V-ATPase function in MB231 cells, the fluorescence indicators SNARF-1 and pyranine were employed to monitor cytosolic and endosome/lysosome pH, respectively. Cells loaded with SNARF-1 show a homogeneous distribution of fluorescence consistent with a cytoplasmic distribution (Fig.3A). By contrast, cells allowed to endocytose the membrane impermeant pH indicator pyranine show punctate staining (Fig.3B). Cells loaded with both pyranine and the lysosomal marker LysoTracker® Red (Figs.3C-E) show extensive co-localization of these two probes (Fig.3E), indicating that pyranine localizes to the endosome/lysosome compartment. The pH dependence of the spectral properties of pyranine and SNARF-1 are shown in Fig.4. For pyranine, the excitation spectra at an emission wavelength of 514nm show an increase in the excitation peak at 465nm and a decrease at 405nm with increasing pH. The ratio of fluorescence at 465nm and 405nm can therefore be used to quantitate endosome/lysosome pH. For SNARF-1 the emission spectra at an excitation wavelength of 534nm shows an increase at 644nm and a decrease at 584nm with increasing pH. Cytosolic pH can therefore be determined from the ratio of fluorescence at 644 and 584.

Using this ratiometric approach, cytosolic and endosome/lysosome pH were quantitated following treatment of MB231 cells with isoform-specific siRNAs. As shown in Fig.5A, treatment with siRNA specific for a3 significantly reduced the cytosolic pH by nearly 0.2 units whereas siRNA specific for a1, a2 or a4 had little effect. By contrast, treatment of cells with siRNA for a1, a2 and a3 all significantly increased the pH of endosomes/lysosomes by 0.2-0.25 units whereas a4 siRNA had no significant effect (Fig.5B). These results suggest that under steady-state conditions the a3 isoform plays a role in controlling both cytosolic and endosome/lysosome pH whereas the a1 and a2 isoforms function in regulation of only endosome/lysosome pH.

**Effect of siRNA treatment on cellular distribution of V-ATPases as assessed using immunofluorescence.** To determine the effect of treatment of cells with siRNAs specific for a subunit isoforms on the cellular distribution of V-ATPases, immunofluorescence studies were performed using an antibody directed against the E subunit of the V-ATPase (part of the peripheral V1 domain). This antibody was employed because of the lack of suitable antibodies capable of recognizing the a subunits in an isoform-specific manner in these cells. Cells were induced to migrate by creation of a “wound” in the cell monolayer. Representative immunofluorescence images of cells at the border of such “wounds” are shown in Fig.7A. As can be seen, extensive intracellular staining was observed, indicative of V-ATPase localization to intracellular compartments (endosomes, lysosomes and Golgi).
In addition, plasma membrane staining was observed in approximately 18% of cells, with staining primarily localized to the leading edge, consistent with previous reports (7,31). Next, the effect of treatment with isoform-specific siRNAs on cellular distribution of V-ATPase was assessed. No detectable effect on the intracellular distribution of V-ATPases was observed following treatment with any of the siRNAs (data not shown). Treatment of cells with siRNAs specific for a3 or a4 but not a1 or a2 resulted in a reduction in the fraction of cells showing plasma membrane staining (Fig.7B), with a4 resulting in the largest decrease (32%). These results suggest that the a4 isoform may be responsible for targeting of the V-ATPase to the plasma membrane in these cells.

**Effect of siRNA treatment on secretion of pro-Cathepsin L** To determine whether treatment with any of the isoform-specific siRNAs had effects on protein trafficking and secretion, the levels of a secreted protein (pro-Cathepsin L) in the media were compared for cells treated with siRNAs and untreated cells. As shown by the representative blot in Fig.8, none of the treatments with the isoform-specific siRNAs had a significant effect on secretion of pro-Cathepsin L.

**DISCUSSION**

Although V-ATPases have been implicated in tumor cell invasion, nothing is known concerning which populations of V-ATPase are functioning in this process. Thus, inhibitors like bafilomycin and concanamycin, while specific for V-ATPases, do not distinguish between V-ATPases containing different subunit isoforms or localized to intracellular versus plasma membranes. This latter problem is due to the membrane permeability of bafilomycin and concanamycin. Similarly, treatment of cells using siRNAs to subunits common to all V-ATPases (such as subunit c) (6), results in inhibition of all V-ATPases in the cell.

Targeting of V-ATPases to different cellular compartments is controlled by isoforms of subunit a, part of the integral V₀ domain (15-19,21-24,28). In mammals, there are four isoforms of subunit a, with a3 and a4 responsible for plasma membrane targeting in osteoclasts and renal intercalated cells, respectively (15-19,22-24). In order to begin to understand the role of different populations of V-ATPases in tumor cell invasiveness, we wished to compare the expression profile of subunit a isoforms in two human breast cancer cell lines with different metastatic potential. MDA-MB231 are highly invasive cells that display plasma membrane V-ATPases and show invasion that is inhibited by bafilomycin and concanamycin (7,30,31). By contrast, MCF7 cells are poorly invasive, show little evidence of plasma membrane V-ATPases and display invasion that is not sensitive to V-ATPase inhibitors at the concentrations that affect MB231 cells (7). The results using quantitative RT-PCR demonstrate that while mRNA encoding all four a subunit isoforms can be detected in both cell types, both a3 and a4 are expressed at much higher levels in MB231 cells as compared to MCF7 cells. It is interesting to note that while the a3 isoform is expressed in a variety of cell types and tissues in addition to osteoclasts (19,22), the a4 isoform has thus far been detected almost exclusively in renal cells (15,24). This result suggests that MB231 cells have up-regulated an atypical isoform of subunit a.

To determine the role that different isoforms of subunit a play in V-ATPase function and tumor cell invasion, isoform-specific siRNAs were employed to selectively reduce mRNA levels for each of these isoforms in MB231 cells. These siRNAs were shown to reduce mRNA levels for the appropriate a subunit species in an isoform-specific manner. To assess the effects of siRNA treatment on V-ATPase function, the pH of the cytosolic and endosome/lysosome compartments was quantitated using the fluorescent pH indicators SNARF and pyranine, respectively. SNARF is taken up by cells as the acetoxymethyl ester and becomes trapped in the cytosol following cleavage by cytosolic esterases (30). As a result, cells labeled with SNARF show a diffuse, cytoplasmic staining pattern (Fig.3A). Pyranine is a membrane impermeant probe which is internalized by cells via fluid phase endocytosis (29). The localization of pyranine to the endosome/lysosome compartment was confirmed by co-localization with the lysosomal marker Lysotracker® Red (Fig.3E). The results show that treatment of MB231 cells with siRNA specific for a3 both acidifies the cytosol and alkalinizes the...
endosome/lysosome compartment. This result is consistent with the presence of a3 in the endosome/lysosome compartment, where inhibition of V-ATPase activity prevents transport of protons out of the cytosol as well as into the lumen of endosomes/lysosomes. Similar conclusions can be drawn for the a1 and a2 isoforms with regards to their effect on endosome/lysosome pH, however no significant effect on the steady state pH of the cytosol is observed (Fig.5A). This may be related to the much lower levels of expression of a1 and a2 relative to a3 in MB231 cells (Fig1A). It is also possible that a1 or a2 are involved in the targeting of V-ATPases to the endosome/lysosome compartment such that their reduction secondarily results in effects on endosome/lysosome acidification. The fact that no significant effect on the intracellular staining pattern was observed by immunofluorescence following treatment with any of the isoform-specific siRNAs, however, argues against this possibility.

To evaluate the role of a subunit isoforms in the invasiveness of MB231 cells, an in vitro invasion assay was employed. The results show that only siRNAs specific for a3 and a4 significantly inhibit invasion. This result is somewhat surprising in light of the absence of any effect of a4-specific siRNA treatment on either cytosolic or endosome/lysosome pH. To assess the effect of siRNA treatment on V-ATPase localization, immunofluorescence was performed using an antibody that recognizes the E subunit (part of the V_1 domain). The results demonstrate the presence of V-ATPase at the plasma membrane in a significant fraction of MB231 cells induced to migrate by creation of a “wound” in the cell monolayer. In addition, V-ATPase tends to localize to the leading edge of the cell, as indicated by co-localization with F-actin. Treatment of cells with siRNA to a4 reduces the fraction of cells showing plasma membrane staining by 32% relative to untreated controls. Inhibition of a3 expression also reduced plasma membrane staining, although to a lesser degree (18%). These results suggest that the a4 isoform (and perhaps to a lesser degree a3) are involved in targeting the V-ATPase to the plasma membrane of MB231 cells and that this cell surface V-ATPase plays a role in the invasiveness of these cells. It should be noted that although the decrease in plasma membrane staining with a4-siRNA treatment is consistent with the a4 isoform being part of the plasma membrane V-ATPase, it does not eliminate the possibility that a4 is involved in the trafficking to the plasma membrane of V-ATPase complexes containing a different isoform. The fact that all of the other siRNA treatments lead to smaller effects on plasma membrane staining, however, argues against this possibility. The absence of any effect of a4 knock-down on cytosolic pH suggests that the cell surface V-ATPases containing a4 do not contribute significantly to cytosolic pH homeostasis in these cells, or that their function in this context can to some degree be compensated by a partial up-regulation of other a subunit isoforms (Fig.2). It is interesting to note in this regard that a1, a3 and a4 have all been reported to be present in the plasma membrane of renal proximal tubule cells, so that a subunit isoforms may have partially overlapping functions (26).

We speculate that a4-containing V-ATPases at the cell surface may promote the invasiveness of MB231 cells by locally acidifying the extracellular environment (i.e., the space between the cell surface and the extracellular matrix). This extracellular acidification may promote invasion by activation of secreted proteases, such as cathepsins, which require an acidic environment for optimal activity. Because the a4 isoform is typically expressed only in renal cells (15,16,24), we suggest that MB231 cells have up-regulated an atypical form of the a subunit for the purpose of enhancing their invasive potential. Additional experiments measuring the effect of siRNA treatment on proton secretion into the extracellular space and looking at whether a4 expression levels in other tumor cell types correlates with their invasive properties will be required to test these ideas.

With respect to inhibition of invasion by knock-down of a3, it seems most likely that this is due, at least in part, to inhibition of intracellular V-ATPases, since the effect on invasion is more significant than the effect observed for plasma membrane staining. Moreover, the effects of a3 knock-down on cytoplasmic and endosome/lysosome pH are most consistent with a3 having an intracellular role. There are a number of possible functions that intracellular V-ATPases may play in promoting invasion, including intracellular activation of secreted proteases.
involved in degradation of extracellular matrix and trafficking of these proteases to the cell surface. Preliminary experiments monitoring protein secretion in MB231 cells have not detected any significant changes following siRNA treatment (Fig.8), although this does not rule out more selective effects on particular trafficking pathways. Additional experiments will be required to elucidate the role of a subunit isoforms in protein trafficking in mammalian cells. It should be noted that up-regulation of intracellular and plasma membrane V-ATPases may play additional roles in cancer progression beyond promotion of tumor cell invasion. For example, increased V-ATPase activity in intracellular compartments may aid in the drug resistance of tumor cells by driving uptake into intracellular vesicles via proton-coupled drug transporters (32-35). Intracellular and plasma membrane V-ATPases may also function in cytoplasmic pH homeostasis. This is particularly important in tumor cells which, because of their high glycolytic rates (32), often require additional aid in acid secretion from the cytoplasm. In summary, this study represents the first examination of the role of V-ATPase a subunit isoforms in pH regulation and invasiveness of breast cancer cells, and suggests the possibility that V-ATPases may represent an effective and potentially selective therapeutic target in controlling tumor cell metastasis.

REFERENCES


**FOOTNOTES**

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1. Abbreviations:  V-ATPase, vacuolar proton-translocating ATPase; MB231, MDA-MB231; RT-PCR, reverse transcription PCR; ConA, concanamycin A; DMSO, dimethyl sulfoxide; NT, non-treated; QRT-PCR, quantitative RT-PCR. Cytosolic pH, pH\text{cyt}; Endosomes/Lysosomes pH, pH\text{E/L}

**FIGURE LEGENDS**

**Figure 1. mRNA levels of subunit a isoforms in MB231 and MCF7 cells.** Using mRNA isolated from cells, quantitative RT-PCR was used to determine the mRNA levels of the different subunit a isoforms, as described under Experimental Procedures. Plasmids expressing the cDNA for each isoform were used to construct a standard curve. All values are normalized to total mRNA loaded as determined by Quant-iT RiboGreen® reagent. The values reported are the ratio of ng of isoform-specific mRNA to the total ng of mRNA. A) a subunit isoform-specific mRNA levels in MB231 cells. B) a subunit isoform-specific mRNA levels in MCF7 cells. C) Ratio of a subunit isoform-specific mRNAs in MB231 versus MCF7 cells. Inset shows data for a1 and a2 on an expanded scale. n=7, error bars = standard deviation.

**Figure 2. Isoform specificity of mRNA knockdown using a subunit isoform-specific siRNAs.** siRNA pools from Dharmacon were used to reduce the mRNA levels of each isoform in MB231 cells as described under Experimental Procedures. The level of knockdown was quantitated by performing QRT-PCR on RNA from cells harvested 96 hrs post siRNA treatment. Knockdown is reported as the ratio of the ng of mRNA in treated cells versus the ng of mRNA in non-treated (NT) cells as determined using the standard curve after QRT-PCR. Values are means, error bars = standard deviation, n=5. *P<0.01 vs. non-treated (NT).

**Figure 3. Fluorescent pH indicators in cytosol and endosomes/lysosomes.** MB231 cells were loaded with 1 mM 8-hydroxypyrene-1,3,6-trisulfonic acid (pyranine), to label endosomes/lysosomes or 7 µM of the acetoxymethyl (AM) ester form of SNARF-1 to label the cytosol as described under Experimental Procedures. (A) Fluorescence image of cells loaded with SNARF-1, excited at 534 nm and the fluorescence emission signal collected at 590 nm (long bandpass filter). (B) Same field as for A, except that pyranine was excited at 465 nm and its emission collected at 514 nm (20 nm bandpass). (C-E) Cells incubated with pyranine as in (A) were washed, to remove extracellular pyranine, and further incubated with 2 µM LysoTracker® Red DND-99 for 3 min to label endosomes/lysosomes. (C) Cells were transferred to the microscope chamber and the fluorescence of pyranine was excited at 458 nm and the emission collected from 514-540 nm. (D) The fluorescence of LysoTracker® Red was excited at 543 nm and the emission collected from 580-650 nm. (E) Shows the merge of C and D and indicates that these fluoroprobes exhibit significant co-localization.

**Figure 4. Spectral properties and pH dependence of fluorescence of SNARF-1 and pyranine.** 1 µM pyranine and 2 µM SNARF-1 (free acid) were dissolved in high K+ buffer and spectral properties evaluated as follows. The excitation spectra of pyranine were collected using an emission wavelength of 514 nm, whereas the emission spectra of SNARF-1 were acquired using 534 nm as the excitation wavelength. Note that these fluoroprobes do not exhibit spectral overlap and do not exhibit fluorescence resonance energy transfer (FRET) that could hamper the interpretation of the data. For pyranine, an increase in the excitation peak at 465 nm and a decrease at 405 nm as pH is increased from 5.5 to 8.5 is observed. 415 nm represents the isoeexcitation wavelength and is used to evaluate dye concentration and/or quenching artifacts. For SNARF-1, the emission signal at 644 nm decreases and at 584 nm
increases as pH is decreased. 600 nm represents the isoemissive wavelength. Consequently, the fluorescence ratios at 465/405 nm and 644/584 nm can be used to monitor pH in endosomes/lysosomes and cytosol, respectively, using a ratiometric approach that allows quantitation of pH.

Figure 5. Steady state pH of cytosol and endosomes/lysosomes in MB231 cells treated with a subunit isoform-specific siRNAs. Cells which had been treated with siRNA directed against a1, a2, a3, or a4 or no siRNA (Control) were incubated as described under Experimental Procedures followed by loading of endosomes/lysosomes with pyranine or cytosol with SNARF-1, as described. 96 hrs post-transfection, simultaneous measurements of pH\textsuperscript{cyt} and pH\textsuperscript{E/L} were performed in cells co-loaded with pyranine and SNARF-1 as described under Experimental Procedures. The conversion of ratio values to pH\textsuperscript{cyt} and pH\textsuperscript{E/L} were performed as described previously. Values are means, error bars = standard deviation, n=6. *P <0.05 vs. control

Figure 6. In vitro invasion of MB231 cells after siRNA treatment. In vitro invasion was assayed using Matrigel™ coated ChemoTx® membranes as described under Experimental Procedures. Cells treated with either concanamycin or siRNA were allowed to invade and then stained with propidium iodide. Fourteen images were taken of the trans-side of the membrane in each well and the number of cells per well were counted and the average over three wells was calculated. Invasion is reported as the ratio of the amount of invasion observed for treated cells divided by the amount of invasion observed for non-treated cells. For concanamycin experiments, non-treated (NT) samples include an equivalent volume of the solvent (DMSO). Values are means, error bars = standard deviation, n=5, *P<0.01 vs NT.

Figure 7. Immunostaining of MB231 cells with antibody against the V-ATPase and quantitation of plasma membrane staining. MB231 cells were grown as a monolayer on coverslips inside 60mm dishes. A wound was made through the monolayer to polarize the cells after which they were immunostained with an antibody directed against the E subunit of V-ATPase as described under Experimental Procedures. A) The three panels show fluorescence using anti-E (left), phalloidin to stain actin filaments (middle) and the merge (right), respectively. White arrows indicate the leading edge. B) Cells grown as described above were treated with siRNA against the four a subunit isoforms, allowed to reach confluence, wounded and immunostained as described under Experimental Procedures. Thirty to forty consecutive images were captured along the wound. For each image, the total number of cells was counted and the number showing a distinct line of staining at the plasma membrane (as indicated by the white arrow) was counted. The data is presented as fraction of cells with plasma membrane staining relative to non-treated. Values are means, error bars = standard deviation, n=3. *P < 0.01 vs. NT.

Figure 8. Secretion of Cathepsin L by MB231 cells. MB231 cells were treated with siRNA against the four a subunit isoforms or no siRNA (NT) for 24 hrs and the media was then replaced with media without siRNA. The cells were incubated for an additional 48 hrs in serum-containing media, 24 hrs in serum-free media and then conditioned media from these cells was collected and analyzed for the presence of pro-Cathepsin L by Western blotting using a commercially available antibody as described under Experimental Procedures. Shown is a representative result.
Figure 3
Figure 4

Excitation Spectra
pyranine

Emission Spectra
SNARF-1

Pyranine Emission= 514 nm

SNARF-1 Excitation= 534 nm

Wavelength (nm)

350 400 450 500 550 600 650 700

8.5

7.5

6.5

5.5
**Figure 5**

**A**
- a4 siRNA
- a3 siRNA
- a2 siRNA
- a1 siRNA
- CONTROL

Steady state pH\textsubscript{Cyt}

**B**
- a4 siRNA
- a3 siRNA
- a2 siRNA
- a1 siRNA
- CONTROL

Steady state pH\textsubscript{CEL}
Figure 6

Relative invasion (treated/nontreated)

Treatment: NT, ConA, a1, a2, a3, a4

* indicates significant difference.
Figure 7

A

anti - V-ATPase  phalloidin  merge

B

Fraction of cells with PM staining treated/nontreated

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* indicates a significant difference
Figure 8

pro-CatL

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Function of a subunit isoforms of the V-ATPase in pH homeostasis and In Vitro invasion of MDA-MB231 human breast cancer cells
Ayana Hinton, Souad R. Sennoune, Sarah Bond, Min Fang, Moshe Reuveni, G. Gary Sahagian, Daniel Jay, Raul Martinez-Zaguilan and Michael Forgac

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