Controlling subcellular localization to alter function: Sending oncogenic Bcr–Abl to the nucleus causes apoptosis

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Altering the subcellular localization of signal transducing proteins is a novel approach for therapeutic intervention. Mislocalization of tumor suppressors, oncogenes, or factors involved in apoptosis results in aberrant functioning of these proteins, leading to disease. In the case of chronic myelogenous leukemia (CML), cytoplasmic Bcr-Abl causes oncogenesis/proliferation. On the other hand, nuclear entrapment of endogenous Bcr–Abl (in K562 human leukemia cells) causes apoptosis. The goal of this study was to determine whether ectopically expressed Bcr–Abl could cause apoptosis of K562 cells when specifically directed to the nucleus via strong nuclear localization signals (NLSs). A single NLS from SV40 large T-antigen or four NLSs were subcloned to Bcr–Abl (1NLS–Bcr–Abl or 4NLS–Bcr–Abl). When transfected into K562 cells, only 4NLS–Bcr–Abl translocated to the nucleus. Bcr–Abl alone was found to localize in the cell cytoplasm, colocalizing with actin due to its actin binding domain. 1NLS–Bcr–Abl also localized with actin. Apoptosis induced by 4NLS–Bcr–Abl was evaluated 24 h post-transfection by morphologic determination, DNA staining, and caspase-3 assay. This is the first demonstration that altering the location of ectopically expressed Bcr–Abl can kill leukemia cells. Multiple NLSs are required to overcome Bcr–Abl binding to actin, thus driving it into the nucleus and causing apoptosis.

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1. Introduction

The causative agent for 95% of all CML cases, Bcr–Abl, is derived from the fusion of the breakpoint cluster region (Bcr) gene on chromosome 22 and the Abelson leukemia oncogene (Abl) on chromosome 9. This reciprocal translocation results in an abnormal, shortened chromosome — “Philadelphia chromosome” or Ph (+) phenotype [1,2]. The resulting Bcr–Abl fusion protein acts as an oncoprotein, and the constitutive activation of tyrosine kinase activity of Abl leads to cell proliferation. Although Gleevec is currently the “gold standard” drug of choice for Bcr–Abl positive CML [3–5], resistance to treatment with Gleevec occurs. This is mostly due to mutations in the Bcr-Abl kinase domain that render it unable to bind to Gleevec [6–8]. Some mutations create a more potent Bcr–Abl oncogene and accelerate disease progression [9]. Other mechanisms for resistance include Bcr–Abl amplification or overexpression, clonal evolution, a decrease in Gleevec bioavailability or cell exposure, and upregulation of drug efflux pumps [6,10]. Many other tyrosine kinase inhibitors (TKIs) are being studied and developed, with a few already approved. Nonetheless, Bcr–Abl also has the potential to develop resistance to these molecules. In addition, since FDA approval, potentially fatal side effects of Gleevec have been uncovered. These include cardiotoxicity [11], the possibility of developing other cancers (due to blockade of tumor suppressor p63) [12], and acute renal failure [13]. Therefore, finding alternative strategies to Gleevec therapy are necessary.

In the cytoplasm Bcr–Abl acts as an oncogene by interacting with multiple signal transduction pathways that transmit anti-apoptotic and mitogenic signals [14]. The key pathways involve ras, MAP kinases, the STAT family, PI3 kinase, and mG, among others [15]. In the nucleus, Bcr–Abl may cause apoptosis due to nuclear Abl’s ability to stabilize p73 and activate its pro-apoptotic functions. Vigneri and Wang have previously shown that nuclear entrapment of Bcr–Abl in K562 cells results in apoptosis, and requires an active tyrosine kinase domain to do so [2]. They used Gleevec to stimulate Bcr–Abl to go to the nucleus (by unknown mechanism), followed by nuclear entrapment by leptomycin B (LMB), a general inhibitor of nuclear export. After washout of Gleevec, Bcr–Abl’s tyrosine kinase activity is re-activated, and the cells undergo spontaneous apoptosis. Unfortunately LMB causes neuronal toxicity and cannot be used therapeutically. This study attempts to address a fundamental question that remains: Does the apoptosis caused by nuclear entrapment of Bcr–Abl require depletion of Bcr–Abl from the cytoplasm, or is it sufficient to send ectopically expressed Bcr–Abl to the nucleus to cause apoptosis?

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2. Materials and methods

2.1. Subcloning and construction of plasmids

Full length Bcr–Abl was removed from pEYK3.1 retroviral vector (a kind gift from Dr. George Daley, Harvard Medical School, Boston) using EcoRI and cloned into pEGFP-C1 (Clontech, Mountainview, CA) at the EcoRI site to make EGFp–Bcr–Abl.

The oligonucleotides 5′-CCGAAAGCAAAAAGAAGAGAGAGATA-GAAT-3′ and 5′-CCGATCTACTCTTCTCTCTTGTGTT-3′ were ligated to pEGFP–Bcr–Abl at the BspEI site. The oligonucleotide insert encodes for the nuclear localization signal (NLS) from SV40 large T-antigen (amino acids P5KRRKV) and is flanked with the BspEI-digested sequence. The ligation resulted in the formation of pEGFP–1NLS–Bcr–Abl and pEGFP–4NLS–Bcr–Abl (a concatemer consisting of four nuclear localization signals).

The bases encoding key residues (T65A and Y66A) in the EGFP sequence were then ligated into the pEGFP-C1 vector using EcoRI and cloned into pEGFP–C1 (Clontech, Mountainview, CA) to eliminate EGFP fluorescence (for use in some co-transfection experiments). The primers used for the mutagenesis were 5′-CTCGT-GACCCTCTGCGCCGCGTGCAGCCTC-3′ and its reverse complement. This plasmid encodes 4NLS–Bcr–Abl with non-fluorescent EGFP.

2.2. Cell line and culture conditions

Bcr–Abl positive K562 cells (human chronic myelogenous leukemia cell line), from our collaborator Dr. K. Elenitoba-Johnson (Univ. of Michigan), were cultured as suspension cells in RPMI 1640 supplemented with 10% FBS (HyClone Laboratories, Logan, UT), 1% penicillin–streptomycin (100U/ml, Gibco BRL, Grand Island, NY), 0.1% gentamycin (Hyclone), and 1%-glutamine (Hyclone). Cells were maintained in a 5% CO2 incubator at 37 °C. Cells were split at a density of 0.5×105/ml two days before transfection.

2.3. Transfection

Transient transfections were performed using Amaza Nucleofector II according to the Amaza protocol for K562 cells. Briefly, 2×106 cells were pelleted from a cell density of 1–5×105 cells/ml, and then resuspended in 100 µL Amaza Solution V. This solution was then added to 10 µg DNA and transfected in an Amaza cuvette using program T-013. Following, 500 µL RPMI was added to the cuvette and cells were transferred to 15 mL RPMI and plated in a 75 cm2 flask for caspase-3 assays. Small aliquots (200 µL) of cells were plated into 4-well live-cell chambers for fluorescence microscopy (Lab-tek chamber slide system, 2 mL, Nalge NUNC International, Naperville, IL) for determination of transfection efficiency. For co-localization experiments where 2 plasmids were transfected simultaneously, 5 µg of each plasmid was used; 5 µg of a single plasmid was used for comparison studies. Cells were incubated for 20–24 h before any other assays were performed.

To calculate the transfection efficiency, four or more fields of cells were counted under the 40X objective. The number of transfected cells (as indicated by the EGFP expression; see methods below for microscope settings) was divided by the total number of cells to obtain transfection efficiency.

2.4. Caspase-3 activity assay

The induction of apoptosis was monitored through the enzymatic activity of caspase-3 using the EnzChek Caspase-3 Assay Kit #1 (Molecular Probes, Eugene, OR) following the manufacturer's protocol. Briefly, 1.5×106 cells were pelleted and resuspended in 50 µL 1× cell lysis buffer followed by a freeze–thaw cycle. The lysed cells were centrifuged for 5 min at 2100 × g. As a control, 1 µL of an mM caspase-3 inhibitor was added to one of the lysates of untransfected K562 cells. 2× substrate (50 µL) solution was then added to 50 µL of lysate and incubated at room temperature for 30 min. A standard curve was made using known amounts of 7-amino-4-methylcoumarin (AMC). Fluorescence was then measured at an excitation wavelength of 355 nm and an emission wavelength of 460 nm. Caspase assays were performed three or more separate times (n = 3). Data was represented as relative fluorescence units per cell, taking transfection efficiencies into account (Fig. 5).

2.5. Actin staining and microscopy

K562 cells transfected with EGFP–Bcr–Abl were fixed with formaldehyde and stained for actin using BODIPY 558/568 Phalloidin from Molecular Probes (Eugene, OR) according to manufacturer’s protocol. Briefly, cells were washed with PBS, pelleted, and resuspended in 4% formaldehyde (in PBS). Cells were incubated in formaldehyde solution for 10 min at room temperature and subsequently washed in PBS. Cells were then placed in 1 mL of 0.1% Triton X-100 (in PBS) for 5 min followed again by washing with PBS. The cells were then incubated with staining solution for 20 min at room temperature and then washed with PBS. Following staining the cells were viewed at 40× with an Olympus IX701F inverted fluorescence microscope (Scientific Instrument Company, Aurora, CO) using the HQ:TRITC filter to detect actin, and a high-quantity narrow band GFP filter (excitation HQ480/20 nm, emission HQ510/20 nm, with beam splitter Q4951p) to detect EGFP. Cells were photographed using a F-View Monochrome CCD camera.

2.6. DNA staining and EGFP microscopy

K562 cell nuclei were stained by the addition of 0.8 µL Hoechst 33342 (10 mg/ml) (Invitrogen, Carlsbad, CA) to 1 mL of cells in a Lab-Tek®II 4-well live-cell chamber (Nalge NUNC) and incubated for 30 min at 37 °C. Pictures were taken approximately 24 h after transfection using a fluorescence microscope with high-quantity narrow band GFP filter (to detect EGFP) and Cyan GFP v2 filter (excitation D436/20 nm, emission D480/40 nm, with beam splitter 455dcpl) to detect H33342. To minimize photobleaching of EGFP chromophore, cells were imaged using neutral density filters in combination with short exposure times. An air stream incubator (Neutek ASI 400, Burnsville, VA) with a variable temperature control was used to maintain the microscope stage and the live-cell chambers at 37 °C. All filters were purchased from Chroma Technology (Brattleboro, VT).

2.7. Cell tracing

Quantitation of Bcr–Abl in the nucleus and cytoplasm was carried out by measuring the fluorescence intensity of EGFP, tagged to the Bcr–Abl, as previously described [18]. All the images were analyzed using analySIS® software (Soft Imaging System, Lakewood, CO).

2.8. Statistical analysis

All experiments were done at least in triplicate (n ≥ 3). The difference between the percent nuclear intensity values (for Fig. 4)
3. Results

When transfected into K562 cells, Bcr–Abl (pEGFP–Bcr–Abl) localizes in the cytoplasm and forms a distinctive ring around the cell (Fig. 1). This localization is indicative of binding to actin, and is expected due to previous reports of Bcr–Abl localization with actin [19]. Co-localization of EGFP–Bcr–Abl and actin is shown in Fig. 1.

Additional nuclear localization signal(s) were added to EGFP–Bcr–Abl to attempt to overcome Bcr–Abl binding to actin. Two plasmids were created, one with 1 NLS subcloned to EGFP–Bcr–Abl, and one with 4 NLSs subcloned to EGFP–Bcr–Abl. When transfected into K562 cells, only EGFP–4NLS–Bcr–Abl localized to the nucleus (Fig. 2G) with the distinct absence of the actin ring surrounding the cell (compared to 2A, wt–Bcr–Abl). The 1NLS construct, while somewhat dispersed throughout the cell, still formed a peripheral actin ring (Fig. 2D), just like wt–Bcr–Abl (Figs. 2A and 1).

Fig. 2. A–C. EGFP–Bcr–Abl transfected into K562 cells; fluorescent (EGFP), nuclear staining with H33342, and phase contrast images, respectively. D–F. EGFP–1NLS–Bcr–Abl transfected into K562 cells; fluorescent (EGFP), nuclear staining with H33342, and phase contrast images, respectively. G–H. EGFP–4NLS–Bcr–Abl transfected into K562 cells; fluorescent (EGFP), nuclear staining with H33342, and phase contrast images, respectively. White arrows (H) indicate apoptotic cells undergoing massive DNA segmentation. Black arrows (I) indicate that these same cells have undergone cell shrinkage, a morphological hallmark of apoptosis [20–24]. Black arrowheads (I) show cells undergoing necrotic/cytoplasmic blebbing. Scale bar (5 \( \mu \)m) representative for all panels (A–I).

was analyzed using an unpaired t-test with Welch’s correction. One-way ANOVA with Tukey’s multiple comparisons post-test was used to assess the differences between relative fluorescence intensity values from the caspase-3 assay (Fig. 5). All statistics were calculated using GraphPad Prism (San Diego, CA).

Table 1

<table>
<thead>
<tr>
<th>Construct transfected</th>
<th>No. of cells transfected</th>
<th>No. transfected cells with segmented DNA</th>
<th>Percentage of cells with segmented DNA</th>
</tr>
</thead>
<tbody>
<tr>
<td>EGFP–Bcr–Abl</td>
<td>178</td>
<td>7</td>
<td>3.9</td>
</tr>
<tr>
<td>EGFP–1NLS–Bcr–Abl</td>
<td>154</td>
<td>5</td>
<td>3.2</td>
</tr>
<tr>
<td>EGFP–4NLS–Bcr–Abl</td>
<td>375</td>
<td>46</td>
<td>12.3</td>
</tr>
</tbody>
</table>

Morphological changes indicating apoptosis include cytoplasmic blebbing, cell shrinkage, and fragmentation of cells into smaller bodies [20–24]. These morphological changes are seen obviously in cells transfected with 4NLS–Bcr–Abl, comparing Fig. 2I (apoptosis) to Fig. 2C and F (no apoptosis). Black arrows indicate shrunken cells (with cell shrinkage indicating apoptosis). Black arrowheads (I) show cells undergoing necrotic/cytoplasmic blebbing.

Twenty-four hours after transfection, cytochemical analyses of apoptosis, including overall cell morphology (Fig. 2C,F, and I), DNA morphology/segmentation (Fig. 2B,E, and H), were examined.

The characteristic change in nuclear morphology (DNA segmentation) is “the most accurate indicator of the involvement of apoptosis in the death of a cell” [22], where the nucleus changes shape from the normal round/oval shape into smaller, non-homogeneous segments. As shown in Fig. 2H, cells transfected with 4NLS–Bcr–Abl show this characteristic DNA segmentation into smaller pieces (white arrows), not seen in other cells (compare to Fig. 2B and E), including healthy cells (Fig. 3, right panel).

Transfected cells were also counted for DNA segmentation or nuclear morphology changes, an accurate indicator of apoptosis [22]. As shown in the Table 1, cells transfected with EGFP–4NLS–Bcr–Abl had a higher percentage of cells with apoptotic nuclei than EGFP–1NLS–Bcr–Abl and EGFP–Bcr–Abl. Only cells with obvious DNA segmentation (more than two nuclear fragments formed) were counted as segmented.

Bcr–Abl is known to form homodimers (tetramers) [25,26]. To determine if the 4NLS–Bcr–Abl was capable of multimerizing with Bcr–Abl, and altering its localization, a co-transfection (co-localization) experiment was performed. EGFP–4NLS–Bcr–Abl with mutated EGFP chromophore (not fluorescent) was transfected with fluorescent EGFP–Bcr–Abl. If the 4NLS–Bcr–Abl protein is able to drag Bcr–Abl into

Fig. 3. Healthy K562 cells. Left panel, phase contrast image. Right panel, nuclear staining with H33342. Nuclei of healthy cells are either round/oval or “kidney” shaped.

Fig. 4. Percent nuclear intensity for cells transfected with EGFP–Bcr–Abl alone (right column) or EGFP–Bcr–Abl and 4NLS–Bcr–Abl (left column). n = 5 cells for each group.
Nuclear entrapment and cytoplasmic depletion of wt-Bcr–Abl may have a synergistic effect on cell death. The nuclear entrapment results in apoptosis via the Abl domain which can cause cell death by stabilizing p73 and activating its pro-apoptotic functions [22,27,28]. Cytoplasmic depletion of wt–Bcr–Abl will remove its ability to interact with the signal transduction proteins involved in gene transcription, mitochondrial processing of apoptotic responses, cytoskeletal organization, and degradation of inhibitory proteins [29]. Here we report that ectopically expressed Bcr–Abl directed to the nucleus is sufficient to induce apoptosis without cytoplasmic depletion. However, it may be speculated that coupling cytoplasmic depletion with nuclear entrapment will synergistically enhance the apoptotic signal. To this end, our ultimate goal is to deplete wt–Bcr–Abl from the cytoplasm, and direct it to the nucleus by our previously described “protein switch” technology [18,30]. We exploit known dimerization domains (DD) of proteins to capture them and move them to a different subcellular compartment. An ectopically expressed “protein switch” can be designed to target any protein with a DD. The protein switch also contains inducible signal sequences to localize to a starting compartment in the cell (e.g., cytoplasm). Upon addition of ligand, the protein switch can move to a different subcellular compartment (e.g., nucleus). Our ultimate goal is to use the protein switch [18,30] to control the subcellular location of Bcr–Abl, and convert Bcr–Abl from an oncogene to an apoptotic factor.

Alternative therapies for CML are needed, despite the “blockbuster” results with Gleevec. Gleevec is not a cure for CML, and patients need to take it for the rest of their lives. Gleevec is also known to have severe side effects including cardiotoxicity [11] and the possibility of developing other cancers (due to blockade of tumor suppressor p53) [12]. Importantly, Gleevec resistance mostly occurs due to different point mutations in Bcr–Abl that make traditional small molecule inhibitors (like Gleevec) unable to bind/inhibit Bcr–Abl. In some cases, a decrease in Gleevec bioavailability/cell exposure and upregulation of drug efflux pumps [6,10] have led to resistance.

The protein switch concept is also being tested in our laboratory with the anti-apoptotic factor survivin, and the tumor suppressor p53 (mislocalized in cancers). Changing localization to alter function may prove to be a new therapeutic strategy for many types of cancer, including CML [31].

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