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Secretory TAT-peptide-mediated protein transduction of LIF receptor α-chain distal cytoplasmic motifs into human myeloid HL-60 cells

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Abstract

The distal cytoplasmic motifs of leukemia inhibitory factor receptor α-chain (LIFRα-CT3) can independently induce intracellular myeloid differentiation in acute myeloid leukemia (AML) cells by gene transfection; however, there are significant limitations in the potential clinical use of these motifs due to liposome-derived genetic modifications. To produce a potentially therapeutic LIFRα-CT3 with cell-permeable activity, we constructed a eukaryotic expression pcDNA3.0-TAT-CT3-cMyc plasmid with a signal peptide (ss) inserted into the N-terminal that codes for an ss-TAT-CT3-cMyc fusion protein. The stable transfection of Chinese hamster ovary (CHO) cells via this vector and subsequent selection by Geneticin resulted in cell lines that express and secrete TAT-CT3-cMyc. The spent medium of pcDNA3.0-TAT-CT3-cMyc-transfected CHO cells could be purified using a cMyc-epitope-tag agarose affinity chromatography column and could be detected via SDS-PAGE, with antibodies against cMyc-tag. The direct administration of TAT-CT3-cMyc to HL-60 cell culture media caused the enrichment of CT3-cMyc in the cytoplasm and nucleus within 30 min and led to a significant reduction of viable cells (P < 0.05) 8 h after exposure. The advantages of using this mammalian expression system include the ease of generating TAT fusion proteins that are adequately transcripted and the potential for a sustained production of such proteins in vitro for future AML therapy.

Key words: Leukemia inhibitory factor; TAT-HIV1; Protein transduction domain; Acute myeloid leukemia; LIF receptor

Introduction

Leukemia inhibitory factor (LIF), as a member of the IL-6 family of cytokines, produces biological effects via binding to its receptor, which consists of a low-affinity LIF receptor α-chain (referred to as LIFRα) and a high-affinity subunit (referred to as gp130), both of which are shared with other IL-6 family cytokines (1). It has been reported that the 136-145-amino acid (aa) region of the distal C-terminal in cytoplasm, which contains 5-tyrosine residues (Y5) and several YXXQ motifs, is genetically conserved among gp130, LIFRα, and G-CSFR in numerous known cell lines (2,3). Thus, these motifs could initiate intracellular signaling by triggering LIFRα-gp130 heterodimerization or gp130-gp130/LIFRα-LIFRα homodimerization via Janus kinase-signal transducer and activator of transcription (Jak/STAT) and subsequently influence the growth or differentiation of leukemia cells (4). Intriguingly, because previous findings have confirmed that the enrichment of triple YXXQ motifs of the LIFRα cytoplasmic domain (LIFRα-CT3) via liposome transfection can inhibit proliferation and induce differentiation in human myeloid leukemia HL-60 cells (5,6),
we hypothesize that the LIFRα-CT3 polypeptide could also achieve a therapeutic effect intracellularly, thus representing a potential future peptide-targeting leukemia therapy.

Advances in protein delivery and the identification of several protein transduction domains have facilitated the delivery of proteins/peptides to cells or organs (7–10). The HIV-trans-activating transduction domain (TAT-PTD) has 11 aa (TAT-PTD <sup>49</sup>–<sup>57</sup>: YGRKKRRQRRR) and can perform the intracellular delivery of proteins across the plasma membrane (11,12). Although the mechanism of TAT-facilitated cellular uptake remains controversial and uncertain, TAT has been widely applied as a tool for protein transduction in vivo and in vitro after fusion with various full-length or truncated peptides (10–13). The technology of generating TAT fusion proteins requires the synthesis of a fusion protein in which TAT is linked to the molecule of interest via the use of a bacterial expression vector. In general, the TAT fusion protein is also linked to some sort of tag so as to facilitate its subsequent purification. The purified recombinant fusion protein could be directly added to mammalian cells in culture or injected in vivo into an animal (14). The above technique is generally highly applicable but laborious; in addition, a protein that is derived from a prokaryotic expression system is potentially more limited by its lack of splicing and the associated post-transcription processing systems or post-translation modifying systems in comparison to eukaryotic expression systems.

In the present study, we have developed an alternative technology that offers advantages in terms of the application of TAT-mediated transduction techniques. We fused TAT-PTD <sup>49</sup>–<sup>57</sup> with LIFRα-CT3 in the recombinant plasmid pcDNA3.0-ss-TAT-CT3-cMyc with a signal peptide (ss) inserted into the N-terminal. Next, the ss-TAT-CT3-cMyc fusion protein was expressed in Chinese hamster ovary (CHO) cells before their culture supernatants were purified through an anti-cMyc agarose affinity column. When we compared the ss-TAT-CT3-cMyc fusion protein to its ss-CT3-cMyc counterpart, the ss-TAT-CT3-cMyc fusion protein was found to be capable of being secreted from CHO cells and subsequently demonstrated a distinct capacity to be delivered into human myeloid leukemia HL-60 cells. Furthermore, we foresee that such transformed cells could be a sustained source of protein transduction domain (PTD) fusion peptides and other macromolecules in vivo.

**Material and Methods**

**Reagents**

Sall, Nhel, Xhol, and BamHI restriction enzymes were purchased from Invitrogen™ (USA). DNA polymerase, PCR purification kit, gel extraction kit, and plasmid mini-prep kit were obtained from Dalian Takara™ (China). The pcDNA3.0 eukaryotic expression vector was purchased from Invitrogen. The rabbit anti-human LIFRα C-terminal antibody was obtained from Sigma™ (USA). All of the primers that were used in this study were synthesized by Sangon™ (China).

**RT-PCR and vector construction**

Total RNA was extracted from 1 x 10<sup>7</sup> HL-60 cells (15). For pcDNA3.0-ss-CT3-cMyc, the ss-CT3-cMyc cDNA was amplified from the pcDNA3.0-gp190CT3 plasmid, which was kindly provided by Yang et al. (5), via a standard polymerase chain reaction with the forward primer F1 (5'-CCGCGATCCGCGCCACCATGGGTTTCAAGGTGCGC-3') and reverse primer R (5'-CGGCTCGAGCTACAGATTTTCAGGTTCTTCTCTTCTGAGATGAGTTTTTGTTCCTCCAGCG-3'). The forward primer F1 contained BamHI, Nhel, SalI (in bold above), and a 21-amino acid peptide, the sequence of which was originated from a human antibody heavy chain gene at the N-terminals as a signal peptide (16) (as underlined above). The reverse primer R contained Xhol (in bold above) and a 10-aa cMyc epitope tag (17) (EQKLISEEDL) at the C-terminal for further protein purification (as underlined above). An approximately 471-bp amplified fragment was cut with BamHI and Xhol and later ligated into a pcDNA3.0 vector, which had been previously digested with BamHI and Xhol, to create the corresponding pcDNA3.0-ss-TAT-CT3-cMyc expression vector.

For pcDNA3.0-ss-TAT-CT3-cMyc, two oligonucleotides were first synthesized and annealed so as to generate a double-stranded oligonucleotide with restriction sites for the Nhel and SalI restriction enzymes and to encode 11 aa (YGRKKRRQRRR) from the basic domain of HIV1-TAT. The sequences were: 5'-CTAGCCTATGCGAGAAGAAGCGGAAGACACAGCGACGAAGAAGAG (the TAT sequence is underlined) and 5'-CTCGAGCTTTCCTGCTGGATCGCTTGTTTTGGCTTTGTTGCATAG-3'. The forward primer F1 contained BamHI, Nhel, SalI (in bold above), and a 21-amino acid peptide, the sequence of which was originated from a human antibody heavy chain gene at the N-terminals as a signal peptide (16) (as underlined above). The reverse primer R contained Xhol (in bold above) and a 10-aa cMyc epitope tag (17) (EQKLISEEDL) at the C-terminal for further protein purification (as underlined above). An approximately 471-bp amplified fragment was cut with BamHI and Xhol and later ligated into a pcDNA3.0 vector, which had been previously digested with BamHI and Xhol, to create the corresponding pcDNA3.0-ss-TAT-CT3-cMyc expression vector.

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The proper orientations and sequences of pcDNA3.0-ss-TAT-CT3-cMyc were confirmed by automated sequencing.

**Transfection**

We transfected CHO cells using the FuGENE® 6 Transfection Reagent (Roche Applied Science, USA) in a serum-containing medium according to manufacturer guidelines. The transfection efficiencies using FuGENe Reagent and DNA ratios of 3:1 and 6:1 (µL and µg for the amount of reagent and DNA, respectively) were similar and approximately twice as high as that obtained using a DNA ratio of 3:2. We carried out transfections with two plasmids: pcDNA3.0-ss-CT3-cMyc and pcDNA3.0-ss-TAT-CT3-cMyc. We selected stable transformed cell lines, which were designated as CHO-CT3-cMyc and CHO-TAT-CT3-cMyc, using 2600 µg/mL Geneticin (Invitrogen) 24 h after transfection. The Geneticin sensitivities of the parent CHO cells were determined in pre-
liminary experiments. The transformed cells were maintained in the same antibiotic concentrations.

Cell culture
CHO cells were maintained in RPMI-1640 medium (Gibco-BRL, Germany) containing 10% fetal bovine serum (FBS, Gibco, Scotland), 100 U/mL penicillin, and 100 μg/mL streptomycin (Gibco-BRL) at 37°C and 5% CO2 in a humidified incubator (18).

The human myeloid leukemia HL-60 cell line was purchased from Cell Bank, Chinese Academy of Sciences (China), and cultured as described elsewhere (19).

Protein expression and purification
Stably transfected CHO-CT3-cMyc and CHO-TAT-CT3-cMyc cells (1.0 x 10^5 cells/mL) were cultured in 20 mL RPMI-1640 medium supplemented with 10% FBS, penicillin/streptomycin, and 2600 μg/mL Geneticin in a 75-cm² culture flask. When the cultures were near confluence, the medium was replaced with 15 mL serum-free RPMI-1640 without Geneticin. The medium was collected after 48-72 h of incubation at 37°C and centrifuged at approximately 2400 g for 15 min. The supernatant was used immediately or kept at -20°C. The process was repeated until 1 L supernatant was collected. After collection, the culture supernatants were combined and filtered through a 0.45-μm filter so as to remove cells and cell debris. Next, the supernatants were loaded onto a gel filtration chromatography column that had been conjugated with anti-cMyc agarose according to manufacturer protocols (A7470, Sigma-Aldrich, Germany). The purified protein was eluted with 0.1 M ammonium hydroxide at pH 11 to 12 and subsequently neutralized with 1 N acetic acid. The concentration of the fusion protein was determined using a BCA protein assay kit (Pierce, USA) and bovine serum albumin as standard.

Western blots
For the Western blots, we cultured CHO-CT3-cMyc or CHO-TAT-CT3-cMyc cells in serum-free medium. We harvested the medium at near confluence, concentrated it using a 3000 MWCO Microcon Centrifugal Filter Device™ (Millipore, USA) at 4°C (20), separated it by 15% SDS-PAGE, and electroblotted it onto polyvinylidene difluoride (PVDF) membranes (15). For immunoblotting, we used a 9E10 anti-Myc epitope tag monoclonal antibody (SC-40, Santa Cruz Biotechnology, USA) at a dilution of 1:2000 and a secondary peroxidase-labeled anti-mouse IgG antibody at a dilution of 1:5000. For the blocking and dilution of the antibodies, we used 1X TBS/Casein Blocker (Bio-Rad, USA). Protein molecular weight markers were purchased from Beyotime (P0062, China).

Exposure of HL-60 cells to purified fusion proteins
To test the transduction of the CT3-cMyc or TAT-CT3-cMyc fusion protein, we first cultured HL-60 cells as described in Ref. 19. Before the direct administration of purified fusion proteins, the HL-60 cells were centrifuged and rinsed three times with PBS to eliminate any possible FBS-induced effects. Next, the cells were re-seeded on six-well plates at a concentration of 1 x 10^4 cells per well with serum-free medium and received fusion proteins at different final concentrations. The same volume of PBS was also added as an internal control. In general, we fixed the corresponding HL-60 cultures after 30 min, 4 h, and 8 h of exposure and examined them under a fluorescence microscope.

Fluorescence microscopy
The pre-staining treatment of non-adherent HL-60 cells (~1 x 10^5/mL) was carried out as described in Ref. 21. The smears were incubated with a mouse-anti-human cMyc primary antibody that had been diluted in blocking serum (1:50), incubated for 30 min at 37°C, and then rinsed for 5 min in 4X saline-sodium citrate/0.1% Tween 20. This process was repeated two additional times. The blocking step was repeated before the smears were incubated with a fluorescein isothiocyanate-conjugated secondary antibody that had been diluted in blocking solution (1:200) and subsequently rinsed as above. The smears were eventually mounted in an anti-fade solution for analysis. The nuclei of HL-60 cells were counterstained with DAPI, visualized with a fluorescence microscope (Leica, Germany) at 630X magnification, and photographed using a charge-coupled camera.

Cell viability and statistical analysis
The viability of HL-60 cells was assessed with the cell counting kit-8 (CCK-8; Beyotime) assay as described by Ting et al. (22) and Lang et al. (23). The HL-60 cells were centrifuged and rinsed three times with PBS so as to eliminate any possible FBS-induced effect. Next, the cells were re-seeded on a 96-well plate at a concentration of 1 x 10^4 cells per well and either exposed to the CT3-cMyc protein or to the TAT-CT3-cMyc protein (30 μM) for 8 h with serum-free medium. The same volume of PBS was also added as an internal control. The culture medium was then changed to RMPI 1640 with 10% FBS. The CCK-8 reagents were added at 24, 48, 72, and 96 h, respectively, after adding the CT3-cMyc protein or the TAT-CT3-cMyc protein. Plates were incubated at 37°C for 1 h and absorbance at 450 nm was recorded.

The cell viability tests were performed at least three times, and the data are reported as means ± SEM. The results were considered to be statistically significant if P < 0.05 as determined by the chi-square test. The analysis was conducted using SPSS 17.0.

Results
Transfection and establishment of cell lines that express and secrete TAT-CT3-cMyc and CT3-cMyc
The full-length LIFRα-CT3 consists of 118 aa residues
To produce a cell-permeable LIFRα-CT3 with an epitope tag (-cMyc), a eukaryotic expression vector bearing the TAT-CT3-cMyc gene was constructed. A recombinant plasmid without a TAT domain was also constructed as the control (Figure 1). Both pcDNA3.0-ss-TAT-CT3-cMyc and pcDNA3.0-ss-CT3-cMyc plasmids are designed for high-level stable expression in mammalian hosts and for the secretion of proteins that are fused at the N-terminal to the human antibody heavy chain gene.

We established stably transfected cell lines that expressed the two fusion proteins (the ~18.6-kDa ss-TAT-CT3-cMyc and ~17.0-kDa ss-CT3-cMyc). CHO-CT3-cMyc and CHO-TAT-CT3-cMyc cells were observed to secrete CT3-cMyc and TAT-CT3-cMyc, respectively, into the culture medium. Western blots of the concentrated, serum-free spent medium of the CHO-CT3-cMyc and CHO-TAT-CT3-cMyc cells, which were prepared using a monoclonal antibody to cMyc, revealed a single band of protein with the same migration as that observed in SDS-PAGE. The serum-free spent medium of wild-type CHO cells was also concentrated and labeled by the same procedure as an internal control. The molecular weight of CT3-cMyc or TAT-cMyc-CT3-cMyc was approximately 18 kDa, whose specificity was confirmed to one protein band extracted from the spent medium of wild-type CHO cells as the negative control (Figure 2).

**TAT-mediated delivery of CT3-cMyc protein to human myeloid HL-60 cells**

To evaluate the ability of TAT-related CT3 delivery to human myeloid leukemia HL-60 cells, we compared the transmembrane delivery of CT3-cMyc and TAT-CT3-cMyc after their direct administration to the serum-free medium of the HL-60 cells at various concentrations or times. Purified CT3-cMyc and TAT-CT3-cMyc fusion proteins were added to a total of 1 x 10^5 HL-60 cells each so as to achieve final concentrations of 10, 30, and 50 µg/mL at 30 min, 4 h, and 8 h. The identical PBS volumes were added to HL-60 cells to act as controls. The intracellular distributions of these fusion proteins were examined by immunofluorescence.

On a time basis, when the administered doses of both fusion proteins were 30 µg/mL, which were similar to the PBS controls (Figure 3), no cMyc-positive cells were found in the HL-60 cells that were obtained from the CT3-cMyc-treated group after 4 h (Figure 3); however, cMyc-positive cells were readily detected in the HL-60 cells that were obtained from the TAT-CT3-cMyc-treated group 30 min after their administration (Figure 3). TAT-CT3-cMyc proteins were primarily found in the nuclei of HL-60 cells. The TAT-CT3-cMyc protein levels, which were calculated based on the number and fluorescence intensity of cMyc-positive cells, peaked at around 30 min to 1 h after their administration and gradually decreased thereafter, although they were still detectable 8 h after administration. These data indicate that TAT-CT3-cMyc but not CT3-cMyc possesses the ability to deliver LIFRα-CT3.

On a dose-dependent basis, because we knew the approximate timing of the transmembrane delivery of TAT-CT3-cMyc, we further explored the dose-response relationships between the fusion protein dosages and fluorescence intensities of cMyc-positive cells after 1 h of TAT-CT3-cMyc fusion protein administration. cMyc-positive cells were readily detected in the HL-60 cells obtained from the TAT-CT3-cMyc-treated group in the 10-µg/mL sample (Figure 4). TAT-CT3-cMyc proteins were primarily found in the nuclei of HL-60 cells. As the protein concentrations were increased, the TAT-CT3-cMyc protein concentrations, which were calculated on the basis of the fluorescence intensity...
of cMyc-positive cells, peaked at a concentration of 30 μg/mL and did not significantly change when compared to the increased concentration up to 50 μg/mL. These data indicate that approximate 30 μg/mL TAT-CT3-cMyc fusion protein was the ideal and saturated dosage close enough in this experimental design for the transmembrane delivery of $1 \times 10^5$ HL-60 cells \textit{in vitro}.

**TAT-CT3-cMyc-induced change in HL-60 cell viability**

To assess the effect of delivered LIFRα-CT3 fusion protein...
protein in HL-60 cells, we compared the cell viability changes after the direct administration of fusion proteins to the serum-free medium of the HL-60 cells at 30 μg/mL for 8 h. No significant difference in cell viability was observed after 24 h of incubation with PBS, CT3-cMyc protein or TAT-CT3-cMyc protein (P > 0.05). A significant decrease in cell viability was observed for HL-60 cells incubated with TAT-CT3-cMyc for 48, 72, and 96 h compared to cells incubated with PBS or CT3-cMyc protein (P < 0.05), based on the CCK-8 assay (Figure 5). A significant decrease in cell viability was not observed in the presence of CT3-cMyc or of an equal amount of PBS as the internal control of wild-type HL-60 cells.

Discussion

In the present study, we have shown that an LIFRα-CT3 fusion protein that bears the protein transduction domain of the HIV1-TAT protein, a signal peptide from the human antibody heavy chain gene, and a cMyc-epitope-tag can be derived from stably transformed eukaryotic expression CHO cells and subsequently delivered into human myeloid leukemia HL-60 cells.

LIF is so named because it can induce a differentiation of the M1 murine myeloid leukemia cell line (24,25). LIFRα-TAT, as reported in our previous research, was observed to have the capacity to activate signal transducer and activator of transcription3 (STAT3) in HL-60 cells by transfection, to initiate LIF-related intracellular signaling, and to facilitate both increasing differentiation and decreasing proliferation (5,26-28); however, transfection or virus-mediated gene delivery may cause an irreversible genetic modification, which makes these deliveries significantly unacceptable for possible clinical practice (29,30). Peptide-based cell delivery systems are greatly expanded by the recognition of PTDs and synthetic peptides with translocation properties (14,31,32). In the present study, we have shown that the TAT-CT3-cMyc fusion protein can be expressed in eukaryotic system CHO cells, secreted into the medium, and efficiently purified. The purified protein was shown to be able to penetrate not only cells but also the nuclear membranes of HL-60 cells (33).

The technique for the production of TAT-related fusion proteins in general requires the synthesis and purification of such proteins using prokaryotic expression systems (34-36). As an alternative, we have developed a method based on the application of a mammalian expression vector, pcDNA3.0. This vector is designed for high-level expression and can be inserted into other domains so as to regenerate novel properties. In our particular case, the resultant novel property was the ability of the fusion proteins to be secreted out of the CHO cells into serum-free medium supernatants, to be effectively concentrated and directly administered to target leukemia cell. As a prototype, we developed stable transformed cell lines (37) with easily detected sensitivities in fixed cells to illustrate the advantages of this technique.

The purity of the recombinant TAT-CT3-cMyc was greater than 90%, as determined by SDS-PAGE. Previous studies have shown that secreted TAT fusion proteins are able to transduce target cells but with very low efficiency (37). According to our study, this is not always the case. The original structure of ss-TAT-CT3-cMyc without mutations was observed to be qualified enough to ensure the secretion of TAT-CT3-cMyc, which is vital in exploring the structure on which TAT-PTD is based. More specifically, the 30 μg/mL TAT-CT3-cMyc fusion protein concentration used in our study was highly effective in transmembrane location within an hour and was primarily located in the nucleus. We speculate that this phenomenon may have been due first to the location of TAT-PTD, which was located downstream of the signal peptide on the N-terminal, which, in turn, may have somehow transformed itself and may have protected the specific sites from being cleaved by endogenous furins (38,39); second, the signal peptide of our study was never applied to the TAT-based eukaryotic expressing system, which may also contribute to the change in protein conformation leading to potential damage.

There are remaining unsolved problems in CHO cell-based TAT-fusion-protein production. The level of protein expression in CHO-transfected cells constantly varies from cell to cell. Some researchers attribute this variation to the inherently stochastic nature of gene expression. The stochastic mechanism in gene expression operates in both prokaryotes and eukaryotes and may explain phenotypic
variations in isogenic cell populations (40). In our observation, TAT-CT3-cMyc and CT3-cMyc were poorly expressed in the transformed CHO cells when determined by immunofluorescence using antibodies against cMyc-epitope-tag but highly detectable in a spent medium of corresponding CHO cells (data not shown). Hence, we further speculate that the gene involved in this liposome-based methodology may induce epigenetic alterations of host cells that make it hard to monitor the expression levels in CHO cells but may somehow not affect the efficiency when fusion proteins are capable of being secreted outside the host cells. Such evidence and reasoning make CHO cells, which are a eukaryotic expression system, an ideal tool for LIFRα-CT3-based acute myeloid leukemia therapy.

The use of a mammalian secretory system to generate TAT-related fusion proteins facilitates their preparation because of its apparent soluble form and more reliable transcripational structure so that they can be directly added to cultured cells. The positive use of LIF in vitro, which is defined as promoting myeloid differentiation and inhibiting cell proliferation, can be achieved by TAT-CT3-based extracellular administration of TAT-CT3-cMyc to HL-60 cells. We have also presumed that the therapeutic application of this technique, whereby parent cells have been transfected with constructs that code for the PTD fusion proteins, could benefit leukemia patients (14).

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