Gene silencing of human RAMP2 mediated by short-interfering RNA

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Abstract. Adrenomedullin (AM) is a regulatory peptide widely expressed, along its receptors, in cells and tissues, of which it controls many basic and specific functions acting in an autocrine-paracrine manner. However, the unequivocal demonstration of the physiological relevance of the regulatory role of AM would require the study of cells where the endogenous AM system has been suppressed. For this task we developed a technique to silence the AM gene in human umbilical vein endothelial cells (HUVECs) and the human embryonal kidney cell line (HEK-293). AM acts via two subtypes of receptor, named AM1 and AM2, which derive from the interaction of the calcitonin receptor-like receptors with two chaperones, called receptor activity modifying proteins (RAMP2 and RAMP3). Hence, we developed a protocol to suppress the human AM1 receptor by silencing the RAMP2 gene by transfection with short interfering RNAs (siRNAs). HUVECs were transfected using a new Ambion transfection reagent. RAMP2 gene silencing was determined in HUVECs by measuring RAMP2 mRNA levels in transfected and control cells by real-time polymerase chain reaction. The RAMP2 gene silencing was ~60% and was observed 48 h after transfection. Matrigel assay in vitro was carried out to evaluate the effects of siRNA sequences. HUVECs cells were plated on matrigel and the analysis of capillary-like tube formation showed that the cells were viable. The knockdown of the RAMP2 gene decreased the formation of tubes in response to 10^-8 M AM. The conclusion is drawn that siRNA technology can be successfully used in the investigations on AM and AM receptor functions.

Introduction

Adrenomedullin (AM) is a multi-functional regulatory peptide, originally isolated from human pheochromocytomas (1), that exerts potent vasorelaxant and hypotensive effects (reviewed in refs. 2,3). AM acts via a calcitonin receptor-like receptor (CRLR), whose selectivity for AM depends on its interaction with a family of receptor activity modifying proteins (RAMPs): RAMP1 generates CGRP1 receptors, while RAMP2 and RAMP3 give rise to two selective AM receptors, provisionally named AM1 and AM2 (reviewed in refs. 4,5).

AM and AM-receptor genes are widely expressed in cells and tissues, of which AM controls specific and basic functions. The best recognized regulatory actions of AM include the inhibition of Ca^{2+}-dependent agonist-stimulated aldosterone secretion from adrenocortical cells (reviewed in ref. 6) and the growth promoting effect on several normal and neoplastic cells (reviewed in refs. 3,7). Due to the rather low levels of circulating AM, it is commonly accepted that this peptide acts in an autocrine-paracrine manner (3,6). However, the physiological relevance of the AM system-mediated functional cell control is far from being satisfactorily demonstrated.

To address this issue, several investigators tried to ascertain whether the suppression of the endogenous AM system alters the cell functions which are known to be regulated by AM. The suppression of the endogenous AM system was obtained by treating in vitro the cells with either selective antagonists of AM receptors, such as AM22-52 and CGRP8-37 (4), or antisense oligonucleotides directed against the AM gene (8). Using this approach, evidence has been provided that the endogenous AM system plays a relevant role in the regulation of i) the secretion and growth of human and rat adrenocortical cells (9-13); ii) the growth of human fibroblasts and keratinocytes (14); and iii) the proliferation of cord blood hematopoietic stem cells (15).

The recent introduction of short interfering RNA (siRNA) technology (16) has led to the development of powerful sequence-specific tools able to induce a long-lasting gene-expression suppression in mammalian cells. We have recently developed a protocol to silence the AM gene in human umbilical vein endothelial cells (HUVECs) and human embryonal kidney cell line (HEK-293) (17), and here we describe the results of a study aimed to silence the RAMP2 gene in HUVECs by transfection with siRNAs. Gene silencing was evaluated by comparing the RAMP2 mRNA levels in siRNA-transfected and control cells, using real-time polymerase chain reaction (PCR) and by verifying whether transfected cells maintain their ability to arrange in capillary-like tubules in response to AM when cultured on Matrigel.

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Materials and methods

Cells and reagents. HUVECs were obtained from Clonetics (Cambrex Bio Science, Milan, Italy). Endothelial cell growth medium (ECGM) was provided by PromoCell (PBI International, Milan, Italy), DNA oligonucleotide templates by MWG Biotech (Florence, Italy), Matrigel by Becton Dickinson Labware (Bedford, MA), and human AM peptide by Phoenix Pharmaceutical (Belmont, CA). Fetal calf serum (FCS), 4',6-diamino-2-phenylindole dilactate (DAPI), collagen, and all other chemicals and reagents were purchased from Sigma-Aldrich Corp. (St. Louis, MO).

siRNA preparation. Four siRNA for human RAMP2 were designed, chemically synthesized and purchased from Dharmacon (Lafayette, CO) in deprotected and desalted form. Two scramble sequences for RAMP2 were selected from two RAMP2-siRNA sequences. Sfold Web Server (http://sfold.wadsworth.org) was used to design specific scramble-siRNA (17). They were generated using the Silencer siRNA construction kit (Ambion, Austin, TX) (18). All sequences are shown in Table I. As a positive control Ambion’s Silencer GAPDH-siRNA and scrambled sequences were used. Cy2-labeled siRNA was synthesized using the Silencer siRNA labeling kit (Ambion), purified, resuspended and hybridized according to the manufacturer’s protocol. The fluorescent siRNA was used to follow the transfection with fluorescence microscopy.

Cell culture. HUVECs were cultured in 0.1% collagen-coated flasks in ECGM, supplemented with 10% FCS, 0.4% ECGS/heparin, 1 μg/ml hydrocortisone, 50 μg/ml gentamycin and 50 μg/ml amphotericin-B. For all experiments HUVECs were used at passages 4-6. Before transfection, the cells were trypsinnized, and after trypsine inactivation, resuspended in normal growth medium and maintained at 37°C while the transfection complexes were prepared.

Transfection reagent and siRNA concentration. The duplexes, added in a single pool at a concentration of 10 or 50 nM, were used to transfect cells using the lipid-based agent for Reverse Transfection siPORT™ NeoFX™ (Ambion). SiPORT NeoFX was diluted into serum-free medium. The siRNA sequences were diluted into serum-free medium to a final concentration of 10 or 50 nM, and then the diluted siPORT NeoFX and RNAs were combined. The formed transfection complexes were incubated for 10 min at room temperature and then dispensed into the empty wells of a culture plate. The cells were pipetted into the culture plate wells containing the transfection complexes. They were plated into 24- or 6-well plates at 2x10⁴ or 15x10⁴ cells/well, respectively.

Visualization of transfection. RAMP2-siRNA was chemically labeled using the Silencer Cy2 siRNA labeling kit (Ambion). Briefly, 5 μg of RAMP2-siRNAs was labeled using 7 μl of labeling reagent, and then purified and resuspended, as indicated by the manufacturer’s protocol. The cells were transfected with 10 nM Cy2-RAMP2-siRNA. After 20 h cells were fixed using methanol/acetic acid (3:1 v/v) and examined under a fluorescence and a reverted phase-contrast light microscope.

HUVEC transfection with siRNAs. The cells were transfected into 6-well culture plates with 10 or 50 nM of RAMP2-siRNA, GAPDH-siRNA and scrambled sequences. All quantities of siRNA tested were incubated in duplicate, and the experiments were replicated 3 times. The cells were incubated under their normal growth conditions and were monitored for gene silencing 48 h after transfection by real-time RT-PCR.

Total RNA extraction and RT. HUVECs were lysed directly in the culture wells, and total RNA extraction was performed with the SV total RNA isolation system (Promega Corporation, Madison, WI), RNA was eluted into RNase-free water, measured by spectrophotometry, and stored at -80°C until use

Table I. siRNA sequence.

<table>
<thead>
<tr>
<th>Sequence range</th>
<th>siRNA sequence</th>
</tr>
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<tbody>
<tr>
<td>RAMP2-siRNA(1) 545-563</td>
<td>Sense 5'-CCUCAUCACUCUUUGUAGUA-3'</td>
</tr>
<tr>
<td></td>
<td>Antisense 3'-GGAGUAGUGAGAACAUCU-5'</td>
</tr>
<tr>
<td>RAMP2-siRNA(2) 290-308</td>
<td>Sense 5'-UCAAAUUGGACCCUAAUCGAA-3'</td>
</tr>
<tr>
<td></td>
<td>Antisense 3'-AGUUAACCUGGAAGCUU-5'</td>
</tr>
<tr>
<td>RAMP2-siRNA(3) 358-376</td>
<td>Sense 5'-GAGAAUGCCUGGACGACUU-3'</td>
</tr>
<tr>
<td></td>
<td>Antisense 3'-CUCUAACGGACCUCGUGA-5'</td>
</tr>
<tr>
<td>RAMP2-siRNA(4) 417-435</td>
<td>Sense 5'-GAGGAGGAUCUUUGAGA-3'</td>
</tr>
<tr>
<td></td>
<td>Antisense 3'-CUCUCCUAGUAGAAGCUU-5'</td>
</tr>
<tr>
<td>RAMP2-siRNA(1) SCRAMBLE</td>
<td>Sense 5'-ACCAUGCUUUAUUCGGAU-3'</td>
</tr>
<tr>
<td></td>
<td>Antisense 3'-UAGCGGAUAAAGGCAGU-5'</td>
</tr>
<tr>
<td>RAMP2-siRNA(2) SCRAMBLE</td>
<td>Sense 5'-UAAGAGUCGUUCUGUGA-3'</td>
</tr>
<tr>
<td></td>
<td>Antisense 3'-UACAGGAGAAGCUUUA-5'</td>
</tr>
</tbody>
</table>
The RT of the total RNA was carried out as follows: oligo dT (Promega) and 1 μg of target RNA were incubated at 70°C for 15 min and at 4°C for 5 min, and then added to 20 μl of a reaction mixture containing 0.5 mM dNTPs (Sigma-Aldrich), 3 mM MgCl₂, 1 U ribonuclease inhibitor, ImProm-II reaction buffer and ImProm-II reverse transcriptase (Promega). The mixture was incubated at 25°C for 15 min, at 42°C for 60 min, and at 70°C for 15 min to inactivate reverse transcriptase.

Real-time PCR. Real-time PCR was carried out as previously described (20-22). Briefly, 1 μl of the RT-reaction solution (diluted to 12 ng/μl) and 12.5 μl of IQ SYBR-Green Supermix (Bio-Rad Laboratories, Milan, Italy), containing 50 mM KCl, 20 mM Tris-HCl (pH 8.4), 3 mM MgCl₂, 0.8 mM dNTPs, SYBR-Green and 6.25 U iTaq DNA polymerase, were mixed to a final volume of 25 μl. PCR was performed in a Bio-Rad I-Cycler iQ detection system, using the following protocol: denaturation program (95°C for 3 min), 35 cycles of two steps of amplification (95°C for 15 sec and annealing for 30 sec), and melting curve (60-90°C with a heating rate of 0.5°C/10 sec). Primer sequences, annealing temperature and accession numbers are shown in Table II. During the exponential phase, the fluorescence signal threshold was calculated, and the fraction number of PCR cycles required to reach the threshold (cycle threshold, Ct) was determined. Ct values decreased linearly with increasing input target quantity, and were used to calculate the relative mRNA expression, according to a mathematical quantification model (23). The specificity of amplification was tested at the end of each run by real-time PCR melting analysis, using the I-Cycler iQ software 3.0. All samples were amplified in duplicate and compared with the respective control. GAPDH or HMBS was used as a reference to normalize data.

Morphogenesis assay on matrigel. Unpolymerized Matrigel was placed in the well (40 μl/well) of a 24-well plate (1.28 cm²/well) and allowed to polymerize for 1 h at 37°C. Twenty-four hours from transfection with siRNA-RAMP2, HUVECs were trypsinized and 2x10⁴ cells were plated in 500 μl medium. Some wells were supplemented with 10⁻⁸ M AM. After 20 h of incubation in a 95% air-5% CO₂ humidified atmosphere at 37°C, the cells were fixed and their ability to give rise to a mesh of capillary-like tubules was assessed under a reverted phase-contrast light microscope, as detailed previously (24-26).

Table II. Primer sequence.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Accession number</th>
<th>Temperature</th>
<th>Oligo sequence</th>
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<tr>
<td>RAMP2</td>
<td>NM_005854</td>
<td>60°C</td>
<td>Sense 5’-CTGCTGGGCGCTGTCTGAA-3’</td>
<td>400</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Antisense 5’-TTCTGACCCCTGTTGCTG-3’</td>
<td>400</td>
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<tr>
<td>GAPDH</td>
<td>NM_002046</td>
<td>60°C</td>
<td>Sense 5’-CTCTCTGGCTCTCTGTTGAC-3’</td>
<td>400</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Antisense 5’-TGAGCGATGTTGCTGCGT-3’</td>
<td>400</td>
</tr>
<tr>
<td>HMBS</td>
<td>NM_000190</td>
<td>60°C</td>
<td>Sense 5’-GGCAATGCGGCTGCAA-3’</td>
<td>400</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Antisense 5’-GGGTACCCACCGCAATC-3’</td>
<td>400</td>
</tr>
</tbody>
</table>

Results

Using fluorescent RAMP2-siRNA, HUVEC transfection was monitored. Twenty hours after transfection, HUVECs exhibited red fluorescent spots in the cytoplasm near the nucleus (Fig. 1). Transfected cells were counted, and the efficiency of transfection was 80.7±4.1% (SD). HUVEC viability was assayed 24 h after transfection. Cells were trypsinized and counted in a Burker's camera: viability was 90% for control (cells incubated only with transfection reagent) and 85% for cells transfected with RAMP2-siRNA.

GAPDH and RAMP2-gene silencing in HUVECs was evaluated by real-time PCR 48 h after transfection with GAPDH-siRNA and RAMP2-siRNA. All siRNAs were tested at 10 and 50 nM (Fig. 2). GAPDH-siRNA was used as a control to verify the capacity of cells to receive siRNA, and to analyze down-regulation of the target gene. The GAPDH gene was silenced by ~40% with 10 nM GAPDH-siRNA and there was no silencing with 50 nM GAPDH-siRNA. RAMP2-siRNA induced a significant decrease in expression of RAMP2 mRNA compared to the controls. The gene was silenced by ~60-65% with 10 nM RAMP2-siRNA, and by only 30% with 50 nM of RAMP2-siRNA. Scrambled GAPDH- and RAMP2-siRNAs did not silence the respective genes.
After seeding on Matrigel, control HUVECs spread and aligned with each other to form branching anastomosing tubes with multi-centric junctions that within 20 h gave rise to a meshwork of capillary-like structures. AM (10⁻⁸ M) increased the length of the meshwork of the control, but not RAMP2-siRNA-transfected HUVECs (Fig. 3).

Discussion

Gene silencing by siRNAs, although being a rather novel and rapidly evolving technology, is already regarded as a very promising technique, not only for mammalian functional genomics, but also for therapeutical purposes (16). To achieve good gene silencing, it is necessary to design specific RNA sequences with appropriate characteristics (27). Following these directions, we developed a protocol to silence the RAMP2 gene in HUVEC cells.

In the present study, we used a new transfection reagent, the Ambion reverse transfection siPORT NeoFX. Reverse transfection is an alternative method of transfection, where cells are transfected while still in suspension (i.e. after trypsinization and prior to plating). This method displays a minimal toxicity and has been reported to improve transfection efficiency over the standard pre-plated method for many cell types tested and saves an entire day in the process. Presumably, the amount of exposed cell surface, and not the number of transfection complexes, is the limiting factor in the traditional adherent transfection. Reverse transfection is believed to increase cell exposure to transfection complexes, thereby leading to a greater transfection efficiency.

Using this technique, we showed an optimal transfection efficiency, and real-time PCR demonstrated a knockdown of RAMP2 gene expression by ~60-65%. Of interest, we observed that transfection with too much siRNA did not evoke more knockdown: 10 nM siRNA appears to be the best concentration. This observation is very important because transfection with high doses of siRNA has been reported to cause selected apoptosis and induction of stress-response genes in cultured cells (28). Moreover, evidence is available that low doses of siRNAs are not able to evoke a cytokine (interferon) response in mammalian cells (29), a finding of great relevance in light of the reported ability of inflammatory cytokines to induce AM-gene expression and AM release from different cell systems (30-34).

The AM receptor-mediated proangiogenic action of AM is well known and the sensitivity of in vitro Matrigel assay to test this effect of AM is well recognized (35-36). The good efficiency of our protocol for RAMP2 gene silencing and the ensuing impairment of AM1 receptor activity has been confirmed by the demonstration that control but not transfected HUVECs display a normal in vitro angiogenic response to AM.

Collectively, our present findings allow us to conclude that siRNA technology is a very promising tool for investigating the role of AM receptors in cell functional regulation.

References