Rab1 GTPase and Dimerization in the Cell Surface Expression of Angiotensin II Type 2 Receptor

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ABSTRACT

The physiological function of angiotensin II (Ang II) is mediated through the Ang II type 1 (AT1R) and type 2 (AT2R) receptors. Our previous studies have demonstrated that cell surface targeting of AT1R is regulated by Rab and Sar1 GTPases and the F(x)6LL motif in the membrane-proximal C terminus. However, the molecular mechanisms underlying the export of nascent AT2R remain poorly defined. In this report, we determined the role of Rab1 GTPase, which specifically controls protein transport from the endoplasmic reticulum (ER) to the Golgi, and receptor dimerization in the biosynthesis of AT2R. Cell surface expression of AT2R was augmented by transient expression of Rab1 and attenuated by dominant-negative Rab1 mutants and small interfering RNA-mediated knockdown of Rab1. Consistently, AT2R inhibition of epidermal growth factor–activated extracellular signal–regulated kinase 1/2 was significantly reduced by the Rab1 mutants, indicating that endogenous Rab1 modulates the cell surface targeting and signaling of AT2R. It is of interest to note that Rab1 augmented the overall expression of AT2R and its mRNA, whereas the Rab1 mutants attenuated the total AT2R expression and enhanced ubiquitin–dependent AT2R degradation. Furthermore, our previously characterized ER export-deficient AT1R mutant in which the F(x)6LL motif was mutated formed both homodimers and heterodimers with AT2R. Dimerization of the AT1R mutant with AT2R blocked AT2R trafficking to the cell surface, suggesting constitutive dimerization of both receptors in the ER and an important role of dimerization in ER export of the receptors. These data demonstrate for the first time that Rab1 GTPase and dimerization modulate export traffic from the ER to the cell surface of newly synthesized AT2R.

Angiotensin II (Ang II) plays an important role in modulation of the physiological function of virtually all organs and in the development of many diseases, such as diabetes, hypertension, myocardial infarction, congestive heart failure, and stroke (Murphy et al., 1991; Hein et al., 1995; Gallinat et al., 2000; Hunyady and Catt, 2006; Mehta and Griendling, 2007). The function of Ang II is mediated through activation of the cell surface Ang II type 1 receptor (AT1R) and Ang II type 2 receptor (AT2R), both of them belong to the seven transmembrane-spanning receptor superfamily coupled to heterotrimeric G proteins (GPCRs). It has been well demonstrated that AT1R mainly couples to the Gq family G proteins to activate several downstream effectors, whereas the function of AT2R is largely mediated through G\textsubscript{i}, to antagonize the AT1R action (Kang et al., 1994; Huang et al., 1996; Zhang and Pratt, 1996; Matsubara, 1998; Hunyady and Catt, 2006).

Similar to signaling mechanisms, the intracellular trafficking of AT1R and its role in regulating receptor function and in the pathogenesis of certain diseases have been extensively studied (Matsubara, 1998; Mehta and Griendling, 2007). These studies have strongly demonstrated that the function of AT1R is critically regulated by its intracellular trafficking, including endocytosis, recycling, degradation, and maturation. We have recently demonstrated that AT1R exit from the endoplasmic reticulum (ER) is mediated by a motif consisting of a phenylalanine and a dileucine spaced by six residues

ABBREVIATIONS: Ang II, angiotensin II; AT1R, angiotensin II type 1 receptor; AT2R, angiotensin II type 2 receptor; GPCR, G protein–coupled receptor; ER, endoplasmic reticulum; HEK, human embryonic kidney; AT2Rm, angiotensin II type 1 receptor mutant; HA, hemagglutinin; ERK, extracellular signal–regulated kinase; GFP, green fluorescent protein; EGF, epidermal growth factor; PCR, polymerase chain reaction; PBS, phosphate-buffered saline; FBS, fetal bovine serum; siRNA, small interfering RNA; PAGE, polyacrylamide gel electrophoresis; MG132, N-benzylxoycarbonyl (Z)-Leu-Leu-leucinal; AR, adrenergic receptor; ALK, activin-like kinase.
[F(x)]_{L,L} in the membrane-proximal C terminus and a single leucine residue in the first intracellular loop of the receptor (Duverney et al., 2004, 2009; Zhou et al., 2006). We have also demonstrated that the cell surface expression of AT1R and AT1R-mediated signaling in HEK293 cells and in cardiomyocytes can be manipulated through modifying the function of the Ras-like small GTPases, including Rab1, Rab2, Rab6, and Sar1 (Wu et al., 2003; Filipanau et al., 2004, 2006; Dong and Wu, 2007; Dong et al., 2008), which are involved in protein transport along the early secretory pathway. Rab1 specifically regulates anterograde protein transport from the ER to the Golgi, and Sar1 coordinates the formation of ER-derived COPII-coated vesicles on the ER membrane that exclusively mediate protein exit from the ER, whereas Rab2 and Rab6 are involved in the retrograde protein transport between the ER and Golgi (Plutner et al., 1991; Jin et al., 1996; Lupashin and Waters, 1997; Allan et al., 2000; Sato and Nakano, 2004; Stephens and Pepperkok, 2004; Altan-Bonnet et al., 2006). In contrast to AT1R, the molecular mechanisms underlying the intracellular trafficking, particularly export from the ER through the Golgi to the cell surface of AT2R, remain largely unknown.

Dimerization has been well described for a variety of GPCRs. Whereas some GPCRs constitutively dimerize in the ER and are then transported to the cell surface, other receptors may form dimers at the plasma membrane in response to agonist stimulation (Jones et al., 1998; Overton and Blumer, 2000, 2002; Szidonya et al., 2008). Although the precise function of receptor dimerization in vivo needs further investigation, many studies have demonstrated that GPCR dimerization may play an important role in regulating receptor folding, maturation, intracellular trafficking, ligand binding selectivity, and signaling. Similar to many other GPCRs, both AT1R and AT2R have been demonstrated to form homo- and heterodimers (AbdAlla et al., 2004; Hansen et al., 2004; Miura et al., 2006). In this report, as an initial approach to elucidating the molecular mechanism of export trafficking of newly synthesized AT2R, we determined the role of Rab1 GTPase and receptor dimerization in the assembly, export, and signal regulation of functional AT2R. Our data indicate that the ER-to-Golgi movement of nascent AT2R is modulated by Rab1 GTPase and that dimerization of AT2R occurs constitutively in the ER and controls ER export of the receptor. These data also suggest that AT2R signaling could be manipulated by modifying its export trafficking along the secretory pathway.

Materials and Methods

Materials. Angiotensin II type 1A receptor in vector pCDM8 and hemagglutinin (HA)-tagged ubiquitin (Ballabeni et al., 2004) were kindly provided by Dr. Kenneth E. Bernstein (Department of Pathology, Emory University, Atlanta, GA) and Dr. Ashok D. Pullikuth (Department of Pharmacology, Louisiana State University Health Sciences Center, New Orleans, LA). Antibodies against Rab1, Rab2, Rab4, Rab6, phospho-ERK1/2, and green fluorescent protein (GFP); rhodamine-conjugated anti-HA antibody; and protein G PLUS-agarose beads were purchased from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA). High-affinity fluorescein-conjugated anti-HA antibody and protein G PLUS-agarose beads were purchased from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA). Alexa Fluor 594-labeled secondary antibodi-
of PBS/1% FBS, the cells were resuspended and the fluorescence was analyzed as described above. Because the staining with the anti-HA antibodies was carried out in the unpermeabilized cells and only those receptors expressed at the cell surface were accessible to the anti-HA antibodies, the measured fluorescence reflected the amount of receptor expressed at the cell surface.

**Fluorescence Microscopy.** For fluorescence microscope analysis of receptor subcellular localization, HEK293 cells were grown on coverslips precoated with poly-L-lysine in six-well plates and transfected with 500 ng of GFP-tagged receptor cDNA. The cells were fixed with ice-cold methanol (−20°C) for 10 min and stained with 4,6-diamidino-2-phenylindole for 5 min. For subcellular localization of AT2R tagged with HA or Myc, HEK293 cells were permeabilized with PBS containing 0.2% Triton X-100 for 5 min and blocked with 5% normal donkey serum for 2 h. The cells were then incubated with antibodies against HA or Myc at a dilution of 1:50 for 1 h. After washing with PBS (3 × 5 min), the cells were incubated with Alexa Fluor 594-labeled secondary antibody (1:2000 dilution) for 1 h at room temperature. The coverslips were mounted, and fluorescence was detected with a DMRA2 epifluorescent microscope (Leica Microsystems, Inc., Deerfield, IL) as described previously (Wu et al., 2003). Images were deconvolved using SlideBook software and the nearest neighbor deconvolution algorithm (Intelligent Imaging Innovations, Denver, CO).

**Double-Stranded RNA-Mediated Interference of Endogenous Rab1.** A double-stranded siRNA with 19-nucleotide duplex RNA and 2-nucleotide 3′ dTdT overhangs (sense, 5′-GAUCCGAC-CAUCGGACUGdTT-3′ and antisense, 3′-dTdTCUAGGCCUGGU- GUACUGUGG-5′) targeting the sequence at the positions 136 to 156 relative to the start code (5′-AAGATCGGAAACATCGAGCTG-3′) of human Rab1 (accession NM030981) and a control nonsilencing siRNA duplex (sense, 5′-UUCUCGGACUGUCGACUdTdT-3′ and antisense, 5′-dTdTAAAGGCGCUUUCCAGGCU-3′) targeting a DNA sequence AATTCCGGAACGTGTGCTG were purchased from Qiagen (Valencia, CA). BLAST searches of the database reveals that this siRNA probe is specific for Rab1 and has no homology to other Rab proteins. HEK293 cells were plated on six-well dishes at a density of 2 × 10^5 cells/well for 12 to 16 h before transfection. The plasmid encoding selected receptor was mixed with siRNA and simultaneously transfected by using Lipofectamine 2000 reagent as described previously (Wu et al., 2003).

**Measurement of ERK1/2 Activation.** HEK293 cells were cultured on six-well plates and transfected with 1 μg of AT2R-GFP. Thirty-six to 48 h after transfection, the cells were serum-starved for at least 3 h and then stimulated with Ang II or with EGF. For measurement of ERK1/2 activation by EGF, HEK293 cells were transfected with the control pDNA vector or Rab1S25N and then stimulated with EGF at a concentration of 10 ng/ml for different times (2–20 min) at 37°C. The stimulation was terminated by addition of 1× SDS-gel loading buffer (Wu et al., 2003; Dong and Wu, 2007). After solubilization of the cells, 20 μl of sample was separated by 12% SDS-PAGE. Activation of ERK1/2 was determined by measuring the level of ERK1/2 phosphorylation with phosphospecific ERK1/2 antibodies. The blots were then stripped and reprobed with total ERK1/2 antibodies to confirm equal loading of the gels.

**Real-time Quantitative PCR.** Total RNA was isolated from HEK293 cells transfected with Rab1 using the standard RNA extraction procedure. Real-time quantitative PCR was performed in a total volume of 20 μl consisting of 2× IQ SYBR Green Super mix (Bio-Rad Laboratories, Hercules, CA; containing iTaq DNA polymerase, dNTPs with dUTP, SYBR Green I, 20 nM fluorescein, and stabilizers) and 200 nM primers. The forward and reverse primers for AT2R were 5′-CAGCAGCTTCTTCGGTTGAAT-3′ and 5′-CTTATCGCGTTTGGTTTGAC-3′, respectively, and the cycling conditions were 95°C for 30 s, then 50 cycles consisting of 20 s at 95°C and 30 s for annealing and extension at 60°C. The β-actin mRNA level was simultaneously measured as an internal control using the forward primer 5′-TGTGATGTTGGAATGGGTCAGA-3′ and the reverse primer 5′-TGTTGATGTTGGAATGGGTCAGAA-3′. The AT2 mRNA level was normalized to β-actin and expressed as -fold change.

**Measurement of AT2R Ubiquitination.** HEK293 cells were cultured on 100-mm dishes and transfected with 3 μg of AT2R-GFP and 3 μg of HA-ubiquitin together with 9 μg of the pCDNA vector, Rab1, or Rab1S25N. After transfection for 36 h, the cells were treated with MG132, a proteasome inhibitor, at a concentration of 10 μM for 12 h. The cells were then washed with PBS and incubated with 500 μl of buffer containing 50 mM Tris, pH 7.4, 150 mM NaCl, 1% Nonidet P-40, 0.5% sodium deoxycholate, 0.1% EDTA, and protease inhibitors at 4°C for 1 h with gentle rotation. After centrifugation for 15 min at 14,000g, 20 μl of protein G PLUS-agarose beads was added into the supernatant for 1 h to remove unspecified binding proteins. The precleared supernatant was then incubated with 2 μg of anti-GFP antibodies and rotated overnight at 4°C followed by incubation with 12.5 μl of protein G-agarose beads for 4 h at 4°C. The beads were then washed twice with buffer, and the bound proteins were eluted with 2× SDS-gel loading buffer. AT2R-GFP and HA-ubiquitin bound to AT2R in the immunoprecipitates were detected by immunoblotting using anti-GFP and HA antibodies, respectively.

**Protein Complementation Experiments.** Homom- and heterodimerization of AT1R and AT2R was measured by protein complementation experiments. GFP and its variants have a property that makes them useful for detecting bimolecular protein interactions. When these proteins are split into the N- and C-terminal fragments, neither fragment is fluorescent when expressed alone in cells nor does coexpression result in the reconstitution of a fluorescent protein. When these complementary N- and C-terminal fragments are fused to two proteins that do associate to form a complex, these two fragments can associate and correctly fold to produce a fluorescent protein (Hu et al., 2002). HEK293 cells were cultured on six-well plates and transfected with 1 μg of vectors expressing the Venus1 or Venus2 fusion proteins. Twenty-four hours after transfection, cells were harvested and washed once with PBS. The cells were then suspended in PBS containing 0.1% glucose and distributed into 96-well microplates (white OptiPlate; PerkinElmer Life and Analytical Sciences, Boston, MA). Fluorescent signals were read using a Fusion instrument (PerkinElmer Life and Analytical Sciences) with a time read of 1 s, a photomultiplier tube voltage of 500, and a gain of 488 nm and an emission filter of 535/25 nm as described previously (Dupré et al., 2007).

To visualize the subcellular localization of the complemented fluorescent proteins, HEK293 cells were harvested 24 h after transfection and seeded on laminin-coated coverslips for 4 h at 37°C. The cells were fixed for 20 min in PBS, pH 7.4, containing 3% paraformaldehyde, washed three times in PBS, and incubated for 1 h at room temperature in PBS containing 0.2% normal goat serum plus 0.2% Triton X-100. Confocal microscopy was performed with an unmodified LSM-510 system (with photomultiplier tube detection; Carl Zeiss Inc., Thornwood, NY) using a 63/1.4 oil Plan-Apochromat objective. GFP (green) was excited at 488 nm with an argon laser, emitting fluorescence at 510 nm (Dupré et al., 2006, 2007).

**Immunoblotting.** Western blot analysis of protein expression was carried out as described previously (Wu et al., 2003). Samples were separated by SDS-PAGE and transferred onto polyvinylidene difluoride membranes. The signal was detected using ECL Plus (PerkinElmer Life and Analytical Sciences) and a luminescent image analyzer (LAS-1000 Plus; Fuji Film, Tokyo, Japan) and quantitated using the Image Gauge, version 3.4 (Fuji Film).

**Statistical Analysis.** Differences were evaluated using Student’s t test, and p < 0.05 was considered as statistically significant. Data are expressed as the mean ± S.E.
Results

Role of Rab1 GTPase in the Cell Surface and Total Expression of AT2R. To determine whether AT2R targeting to the cell surface is regulated by Rab1 GTPase, we determined the effect of expressing Rab1 and its dominant-negative mutants or Rab1 siRNA on the cell surface expression, total expression, subcellular distribution, and signaling of AT2R. In the first series of experiments, HA-AT2R was transiently expressed together with Rab1 or its dominant-negative mutants Rab1S25N and Rab1N124I, and the cell surface expression of AT2R was quantified by flow cytometry after staining with anti-HA antibodies in unpermeabilized cells. The expression of wild-type Rab1 increased the cell surface expression of AT2R by 2.3-fold, whereas Rab1S25N and Rab1N124I significantly inhibited the cell surface expression of AT2R by 82 and 65%, respectively, compared with cells transfected with the empty pcDNA3 vector (Fig. 1A).

To determine whether Rab1 expression altered total AT2R expression, GFP-tagged AT2R was expressed with Rab1, Rab1S25N, or Rab1N124I and the overall expression of AT2R was determined by flow cytometry measuring the GFP signal. The expression of Rab1 significantly enhanced total expression of AT2R by 38%, whereas Rab1S25N and Rab1N124I inhibited total AT2R expression by 69 and 44%, respectively, compared with cells transfected with the empty pcDNA3 vector (Fig. 1B). The ratio of cell surface to total expression of AT2R was markedly increased by expression of Rab1 and decreased by expression of dominant-negative Rab1 mutants (Fig. 1C). These data indicate that AT2R transport to the cell surface uses a Rab1-dependent pathway and the expression level of endogenous Rab1 is a rate-limiting factor for AT2R targeting to the plasma membrane.

In the second series of experiments, we determined the effect of Rab1S25N and Rab1N124I on the subcellular distribution of AT2R. To exclude the possibility that the epitope tagging may produce nonspecific effects on AT2R trafficking, AT2R tagged with three different epitopes (HA or Myc at the N terminus and GFP at the C terminus) was used. AT2R tagged with these different tags robustly expressed at the cell surface (Fig. 2A), suggesting that epitope tagging did not influence the subcellular localization of the receptor. Consistent with its inhibitory effect on the cell surface expression of AT2R as quantified by flow cytometry, transient expression of Rab1S25N and Rab1N124I resulted in a marked perinuclear distribution of AT2R (Fig. 2A), consistent with well-established Rab1 function in regulating ER-to-Golgi protein transport. To further prove that intracellular accumulation of AT2R was indeed caused by expression of the Rab1 mutants, we determined the colocalization of HA-tagged AT2R and GFP-tagged Rab1 mutants. HA-AT2R was arrested in the perinuclear region of the cells that were transfected with GFP-Rab1S25N or GFP-Rab1N124I (Fig. 2B). AT2R did not colocalize with either Rab1 mutant (Fig. 2B) as AT2R and the dominant-negative Rab1 mutants are presumably localized to the ER/Golgi and the cytoplasm, respectively.

Effect of Rab1 on the mRNA Level and Ubiquitination of AT2R. Our preceding data demonstrated that expression of Rab1 and its dominant-negative mutants altered the overall expression of AT2R, which is in contrast to their effect on the total expression of many other exogenously expressed GPCRs, such as AT1R, α2β-AR, and β2-AR in HEK293 cells (Wu et al., 2003). Therefore, it is interesting to further study how Rab1 influences total AT2R expression. We first determined whether Rab1 expression could alter total mRNA level of AT2R. HEK293 cells were transiently transfected with AT2R-GFP together with pcDNA3.1 (Control), wild-type (WT) Rab1 or the dominant-negative Rab1 mutants Rab1S25N and Rab1N124I. AT2R expression at the cell surface was quantitated by flow cytometry after incubation with anti-HA antibodies as described under Materials and Methods. The data shown are percentage of the mean value obtained from the cells transfected with HA-AT2R and pcDNA3.1 and are presented as the mean ± S.E. of four experiments. B, effect of Rab1 on the overall expression of AT2R. HEK293 cells were transfected with AT2R-GFP together with pcDNA3.1 (Control), WT Rab1, Rab1S25N, or Rab1N124I. Total AT2R expression was quantitated by flow cytometry measuring GFP fluorescence. The data shown are percentage of the mean value obtained from the cells transfected with AT2R-GFP and pcDNA3.1 and presented as the mean ± S.E. of four experiments. C, ratio of cell surface AT2R to total AT2R expression generated from the data shown in A and B. The data presented as the mean ± S.E. of four experiments. * P < 0.05 versus control.
GPCR degradation may be mediated through ubiquitination-dependent pathway. Therefore, we determined whether expression of Rab1 could promote ubiquitination of AT2R. GFP-tagged AT2R and HA-tagged ubiquitin were coexpressed in HEK293 cells, and the ubiquitination state of AT2R was determined by immunoblotting for the HA-tagged ubiquitin after immunoprecipitation of the receptor using anti-GFP antibodies. In the absence of MG132, a proteasome inhibitor, AT2R in the immunoprecipitates was not detectable. In the presence of MG132, a proteasome inhibitor, AT2R in the immunoprecipitates was detected. Therefore, we determined whether expression of Rab1 could increase AT2R ubiquitination.

**Inhibition of the Cell Surface Expression and Intracellular Accumulation of AT2R by siRNA-Mediated Knockdown of Rab1.** To further define the function of Rab1 in regulating AT2R trafficking, we determined the effect of siRNA-mediated knockdown of endogenous Rab1 on the cell surface expression and subcellular distribution of AT2R. Introduction of Rab1 siRNA into HEK293 cells dramatically reduced the Rab1 expression level compared with control cells without transfection or transfected with control siRNA (Fig. 4A). In contrast, Rab1 siRNA transfection had no effect on the expression of the closely related small GTPases Rab2, Rab4, and Rab6 (Fig. 4A), demonstrating that the Rab1 siRNA selectively altered Rab1 expression. Similar results were obtained in two separate experiments. Molecular mass is indicated in the left.

**Effect of Rab1 Function on AT2R Signaling.** It has been demonstrated that AT2R inhibits EGF-
mediated ERK1/2 activation (De Paolis et al., 2002). To determine whether Rab1 is capable of regulating AT2R signaling through modifying receptor export to the cell surface, we determined whether transient expression of Rab1S25N could alter the inhibitory effect of AT2R on EGF-mediated ERK1/2 activation. AT2R was transiently expressed with or without expression of Rab1S25N in HEK293 cells and stimulated with EGF alone or EGF plus Ang II. EGF stimulation potently activated ERK1/2, which can be inhibited by Ang II stimulation in cells expressing AT2R. Attenuation of EGF-mediated ERK1/2 activation by AT2R was markedly inhibited in cells expressing Rab1S25N (Fig. 5A). These data suggest that Rab1S25N blocks AT2R transport to the cell surface and attenuates its function.

To determine whether the effect of Rab1S25N on ERK1/2 activation by EGF and Ang II/AT2R was due to direct influence of Rab1S25N on the EGF-mediated ERK1/2 activation, we measured the effect of Rab1S25N on EGF-mediated ERK1/2 activation. EGF stimulation robustly activated ERK1/2 in a time- and dose-dependent manner (Fig. 5, B and C), and ERK1/2 activation by EGF was almost identical in cells transfected with the control empty vector compared with cells transfected with Rab1S25N. These data suggest that inhibition of Rab1 function does not significantly influence EGF-mediated ERK1/2 activation and imply that Rab1 modulation of AT2R effect on EGF-mediated ERK1/2 activation is not due to the effect of Rab1 on the EGF/EGFR-mediated ERK1/2 activation per se.

Fig. 4. Effect of siRNA-mediated knockdown of Rab1 on the cell surface expression and subcellular distribution of AT2R. A, HEK293 cells cultured in six-well dishes were transiently transfected with control siRNA or Rab1 siRNA as described under Materials and Methods. At 48 h after transfection, total homogenate protein was separated by 12% SDS-PAGE, and the expression levels of Rab1, Rab2, Rab4, and Rab6 were determined by immunoblotting using Rab isoform-specific antibodies (left). Quantitative data are expressed as percentages of Rab protein expression in cells transfected with control siRNA and are presented as the means ± S.E. of three individual experiments (right). *P < 0.05 versus control siRNA. B, effect of Rab1 siRNA on AT2R expression at the cell surface. HEK293 cells were transfected with HA-AT2R together with control or Rab1 siRNA as described under Materials and Methods. AT2R expression at the cell surface was measured by flow cytometry after staining by anti-HA antibodies. The data shown are percentage of the mean value obtained from the cells transfected with HA-AT2R and control siRNA and are presented as the mean ± S.E. of three experiments. *P < 0.05 versus control siRNA. C, effect of Rab1 siRNA on AT2R subcellular localization. HEK293 cells were transfected with GFP-conjugated AT2R together with control or Rab1 siRNA as described above. The subcellular distribution of AT2R was revealed by detecting GFP fluorescence. The data are representative images of at least four individual experiments. Scale bar, 10 μm.

Fig. 5. Effect of Rab1 on AT2R function in regulating EGF-mediated ERK1/2 activation. A, attenuation of AT2R inhibition on EGF-activated ERK1/2 by Rab1S25N. HEK293 cells cultured on six-well dishes were transfected with HA-AT2R together with control or Rab1 siRNA as described under Materials and Methods. AT2R expression at the cell surface was measured by flow cytometry after staining by anti-HA antibodies. The data shown are percentage of the mean value obtained from the cells transfected with HA-AT2R and control siRNA and are presented as the mean ± S.E. of three experiments. *P < 0.05 versus control siRNA. B, effect of Rab1 siRNA on AT2R expression at the cell surface. HEK293 cells were transfected with HA-AT2R together with control or Rab1 siRNA as described under Materials and Methods. AT2R expression at the cell surface was measured by flow cytometry after staining by anti-HA antibodies. The data shown are percentage of the mean value obtained from the cells transfected with HA-AT2R and control siRNA and are presented as the mean ± S.E. of three experiments. *P < 0.05 versus control siRNA. C, effect of Rab1 siRNA on AT2R subcellular localization. HEK293 cells were transfected with GFP-conjugated AT2R together with control or Rab1 siRNA as described above. The subcellular distribution of AT2R was revealed by detecting GFP fluorescence. The data are representative images of at least four individual experiments. Scale bar, 10 μm.

To determine whether Rab1 is capable of regulating AT2R signaling through modifying receptor export to the cell surface, we determined whether transient expression of Rab1S25N could alter the inhibitory effect of AT2R on EGF-mediated ERK1/2 activation. AT2R was transiently expressed with or without expression of Rab1S25N in HEK293 cells and stimulated with EGF alone or EGF plus Ang II.
Dimerization of the ER-Retained AT1R Mutant with Wild-Type AT1R and AT2R. AT1R and AT2R have been well demonstrated to form homodimers and heterodimers (AbdAlla et al., 2004; Hansen et al., 2004; Miura et al., 2006). We have recently demonstrated that the AT1Rm, in which the F(X)₆LL motif in the membrane-proximal C terminus was mutated, was unable to transport to the cell surface and was arrested in the ER (Duvernay et al., 2004). We first asked whether the export-deficient AT1Rm retained its ability to form homodimers as measured using a protein complementation assay based on the reconstitution of the GFP variant Venus. As shown in Fig. 6A, both wild-type AT1R and AT2R formed homodimers, reconstituting Venus fluorescence compared with negative controls consistent with many reports in the literatures (AbdAlla et al., 2004; Hansen et al., 2004; Miura et al., 2006). Similar to wild-type AT1R and AT2R, the mutant AT1Rm also formed homodimers (Fig. 6A). Under the same experimental conditions, the activin-like kinase (ALK) 5 (a transforming growth factor-β receptor) formed homodimers but did not form heterodimers with either AT1R or AT2R (Fig. 6A). We then used confocal microscopy to visualize the subcellular localization of reconstituted dimers that would indicate where AT1R, AT2R, and AT1Rm formed homodimers. As shown in Fig. 6B, the GFP fluorescence was observed both at the cell surface and intracellular compartments in cells expressing either AT1R or AT2R, suggesting that AT1R and AT2R could form dimers at the plasma membrane and intracellular compartments. In contrast, the GFP fluorescence was seen only in intracellular compartments, presumably ER, but not at the plasma membrane, in cells expressing AT1Rm (Fig. 6B).

We next determined whether AT1Rm could undergo heterodimerization with wild-type AT1R and AT2R. Similar to heterodimerization between wild-type AT1R and AT2R, protein complementation experiments again showed that AT1Rm heterodimerized with both AT1R and AT2R (Fig. 6A). These data demonstrate that mutation of the F(X)₆LL motif in the C termini does not prevent receptor dimerization. It is important to note that these data suggest that AT1R and AT2R dimerization (homo- and heterodimerization) occurs constitutively during their biosynthesis and maturation in the ER.

To determine whether the ER-retained AT1Rm could function as a dominant-negative mutant inhibiting AT2R transport to the cell surface, AT1Rm was expressed together with HA-AT2R in HEK293 cells, and the cell surface expression and subcellular distribution of AT2R were analyzed. The subcellular localization of the receptors showed that AT2R was trapped in the perinuclear region, presumably in the ER (Fig. 6B). These data indicate that AT1Rm may exert a dominant-negative effect on AT2R cell surface targeting.

**Discussion**

The physiological function of Ang II is mediated through activating AT1R and AT2R at the plasma membrane, which couple to different G proteins to produce largely opposite effects. We have characterized previously the regulation of exit from the ER and subsequent transport from the ER through the Golgi to the cell surface of AT1R (Wu et al., 2003; Duvernay et al., 2004; Filipeanu et al., 2004; Dong and Wu, 2007; Dong et al., 2008). However, the regulators involved in the AT2R export trafficking and the role of export trafficking in AT2R signaling remain poorly defined. Here, we have demonstrated that the small GTPase Rab1 is an important determinant of surface trafficking AT2R. We have also shown that both homo- and heterodimerization of AT1R and AT2R occur in the ER and that a single defective trafficking signal in one monomer equivalent in a receptor dimer is sufficient to block cell surface transport, suggestive of an important quality control step in the function of dimeric GPCRs.

Rab proteins represent the largest branch of the Ras-like small GTPase superfamily. Rab1 GTPase specifically modulates ER-to-Golgi protein transport (Plutner et al., 1991; Jin et al., 1996; Lupashin and Waters, 1997; Allan et al., 2000).
As an initial approach to elucidating the molecular mechanism of GPCR export trafficking, we have determined the role of Rab1 in the cell surface targeting of several GPCRs, including AT1R, α1-AR, α2B-AR, β1-AR, and β2-AR (Wu et al., 2003; Filipeanu et al., 2004, 2006). Our previous studies have shown that the ER-to-cell surface transport of different GPCRs is mediated through distinct pathways. Whereas α2B-AR transport is independent of Rab1, AT1R export is dependent on Rab1 (Wu et al., 2003; Filipeanu et al., 2004, 2006). Interestingly, augmentation of Rab1 function by overexpressing wild-type Rab1 facilitates the cell surface transport of endogenous AT1R in cardiomyocytes but not exogenously expressed AT1R in HEK293 cells (Wu et al., 2003; Filipeanu et al., 2004). Based on these sensitivities, ER-to-cell surface transport of GPCRs can be divided into three different pathways (Dong et al., 2007). The first pathway is inhibited by attenuating Rab1 function and facilitated by increasing Rab1 function and is used by AT1R and α1-AR in cardiac myocytes. The second pathway is inhibited by attenuating Rab1 function, but not sensitive to increasing Rab1 function (such as β1-AR and β2-AR). The third pathway is not influenced by manipulating Rab1 function and is used by α2B-AR. Our current data indicate that AT2R cell surface transport is inhibited by the dominant-negative Rab1 mutants and Rab1 siRNA and facilitated by overexpressing Rab1, indicating that AT2R export is mediated through a Rab1-dependent pathway and the endogenous Rab1 level is a rate-limiting factor for its transport.

It is apparent that AT1R and AT2R have distinct sensitivities to the manipulation of Rab1 function. First, attenuating Rab1 function inhibits more potently the cell surface expression of AT2R than AT1R. Our previous and current data demonstrate that Rab1S25N inhibits the cell surface expression of AT1R and AT2R by 65 and 82%, respectively, in HEK293 cells (Wu et al., 2003). Likewise, Rab1N124I inhibits the cell surface expression of AT1R and AT2R by 45 and 65%, respectively (Wu et al., 2003). Such a moderate inhibition of Rab1N124I on endogenous AT1R was also observed in neonatal cardiomyocytes by 36% and in HL-1 myocytes by 44% (Filipeanu et al., 2004). Second, responses of AT1R and AT2R to the augmentation of Rab1 function are also different. Our previous data demonstrated that the cell surface transport of exogenously expressed AT1R was not altered by overexpressing Rab1 in HEK293 cells (Wu et al., 2003). In contrast, Rab1 overexpression increased AT2R at the cell surface by 130% in HEK293 cells. These data indicate that functional manipulation of components of the transport machinery such as Rab1 to alter protein ER-to-Golgi transport along the secretory pathway may provide a novel means for differential modulation of the export trafficking of AT1R and AT2R.

The opposing effects of Rab1 and Rab1S25N on the total AT2R synthesis are likely mediated through distinct molecular mechanisms. Similar to cell surface expression, total AT2R is enhanced by Rab1 and attenuated by Rab1 mutants. These data are in contrast to our previous data showing that Rab1 did not significantly alter the overall expression of several exogenously expressed GPCRs, including AT1R (Wu et al., 2003). We have further demonstrated that Rab1 expression markedly increased the AT2R mRNA level, suggesting that Rab1-enhanced cell surface expression of AT2R is at least in part due to increased AT2R synthesis. These data also suggest that Rab1, as a well characterized trafficking regulator, is able to modulate protein synthesis at the transcriptional level. However, the molecular mechanism underlying Rab1 function in modulating AT2R synthesis needs further investigation. It is possible that Rab1 overexpression may alter mRNA stability, which would eventually influence protein synthesis. Consistent with this possibility, a recent study has demonstrated that Rab1 depletion significantly decreases the RNA levels of hepatitis C virus (Sklan et al., 2007). However, Rab1S25N had no effect on the mRNA levels of AT2R. Rather, it dramatically enhanced AT2R ubiquitination. These data indicate that similar to many other GPCRs, AT2R undergoes ubiquitination and ubiquitination-dependent proteolysis seems to be main determinant for the reduction in the total AT2R expression caused by Rab1S25N. Interestingly, although expression of Rab1 and Rab1S25N results in opposing effects on the total and cell surface expression of AT2R, they similarly enhance AT2R ubiquitination. It is possible that the Rab1-promoted AT2R ubiquitination is caused by the increase in the total AT2R synthesis induced by Rab1 expression. Rab1 overexpression markedly facilitates total AT2R synthesis, which will contribute to the increased cell surface expression of the receptor. Meanwhile, not all the newly synthesized AT2R are able to export from the ER and transport to the cell surface, some receptors will remain in the ER and then target from the degradation pathway via ubiquitination.

It is important to note that our previous results and the current study strongly indicate that, in addition to modulating export trafficking of AT1R and AT2R, Rab1 is also capable of regulating their function. It has been well demonstrated that AT1R and AT2R couple to different G proteins and activate opposite signal transduction pathways (Murphy et al., 1991; Mehta and Griendling, 2007). We have shown previously that increase Rab1 expression facilitates AT1R function in regulating ERK1/2 activation and growth in cardiac myocytes (Wu et al., 2001; Filipeanu et al., 2004). Here, we demonstrate that expression of dominant-negative Rab1 mutants also inhibited AT2R function. Rab1 function in regulating AT1R and AT2R signaling is probably mediated through modulating their ER-to-Golgi trafficking, which would eventually alter the number of the receptors at the cell surface. Therefore, our studies provide a possibility to manipulate signaling of very closely related GPCRs (e.g., AT1R versus AT2R) by manipulating their trafficking along the early secretory pathway.

AT1R and AT2R form homodimers and heterodimers (AbdAlla et al., 2004; Hansen et al., 2004; Miura et al., 2006). However, where they form dimers and the role of the dimerization in the ER export remain elusive. Our current data have demonstrated that AT1R and AT2R dimers could be detected in both intracellular compartments and at the plasma membrane. Furthermore, similar to AT1R, the AT1R mutant defective in the ER export homodimerized and heterodimerized with AT1R and AT2R. These data indicate that homo- and heterodimerization of AT1R and AT2R occur constitutively and as early as in the ER. More interesting to note is that the AT1R mutant markedly blocked the cell surface transport of wild-type AT2R revealed by subcellular distribution. These data suggest that AT1R mutant may function as a dominant-negative mutant to block the transport of wild-type AT2R. The inhibition of AT2R cell surface expression by the AT1R mutant is probably mediated through heterodimerization between two receptors. The AT1R mutant defective in the ER export forms heterodimers with wild-type AT2R and traps AT2R inside the cell. This pos-
sibility is strongly supported by the inhibition of cell surface expression of GPCRs by heterodimerization of wild-type receptors with their intracellularly retained receptor mutants. These data provide strong evidence implicating that homo- and heterodimerization of AT1R and AT2R play an essential role in their export from the ER and subsequent transport to the cell surface. These data also provide a useful means for manipulating the function of Ang II receptors by using their export-defective mutants.

In summary, our previous and current studies have demonstrated that the cell surface and total expression of AT1R and AT2R have different sensitivities to the manipulation of Rab1 function, although export trafficking of both receptors is mediated through a Rab1-dependent pathway. The ER export-defective AT1R mutant functions as a dominant-negative mutant inhibiting AT2R transport to the cell surface, which is mediated through heterodimerization, suggesting that AT1R and AT2R constitutively form dimers in the ER and that dimerization plays an important role in their export from the ER. Taken together, these data provide strong evidence indicating that the export trafficking of AT1R and AT2R is mediated through distinct, highly regulated processes and that the function of Ang II may be manipulated through modulating its receptor antiporter trafficking along the early secretory pathway.

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