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# Subtype-Specific Sorting of The $ET_A$ Endothelin Receptor by A Novel Endocytic Recycling Signal for G Protein-Coupled Receptors

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**Abbreviations:** DiI-LDL, dioctadecyl-tetramethylindocarbocyanine-low density lipoprotein; GFP, green fluorescent protein; GPCR, G protein-coupled receptor; GRK, G protein-coupled receptor kinase; PDZ, PSD-95/Dlg/ZO-1; Tf-Rhod, tetramethylrhodamine-labeled transferrin.

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### **Abstract**

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We have previously reported that endocytic sorting of ET<sub>A</sub> endothelin receptors to the recycling pathway is dependent on a signal residing in the cytoplasmic carboxyl-terminal region. The aim of the present work was to characterize the carboxyl-terminal recycling motif of the ET<sub>A</sub> receptor. Assay of truncation mutants of the ET<sub>A</sub> receptor with increasing deletions of the carboxyl-terminal tail revealed that amino acids 390 – 406 contained information critical for the ability of the receptor to recycle. This peptide sequence displayed significant sequence similarity to several protein segments confirmed by X-ray crystallography to adopt antiparallel β-strand structures (β-finger). One of these segments was the β-finger motif of nNOS (neuronal nitric oxide synthase) reported to function as an internal PDZ (PSD-95/Dlg/ZO-1) domain-binding ligand (PDZ ligand). Based on these findings, the three-dimensional structure of the recycling motif of ET<sub>A</sub> receptor was predicted to attain a β-finger conformation acting as an internal PDZ ligand. Site-directed mutagenesis at residues that would be crucial to the structural integrity of the putative  $\beta$ -finger conformation or PDZ ligand function prevented recycling of the ET<sub>A</sub> receptor. Analysis of more than 300 G protein-coupled receptors (GPCRs) identified 35 different human GPCRs with carboxyl-terminal sequence patterns that fulfilled the structural criteria of an internal PDZ ligand. Among these are several receptors reported to follow a recycling pathway. In conclusion, recycling of ET<sub>A</sub> receptor is mediated by a motif with the structural characteristics of an internal PDZ ligand. This structural motif may represent a more general principle of endocytic sorting of GPCRs.

## Introduction

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The physiological effects of the vasoactive peptide endothelin-1 (ET-1)<sup>1</sup> are mediated by the ET<sub>A</sub> and ET<sub>B</sub> receptors, which belong to class A G protein-coupled receptors (GPCRs) (Yanagisawa, 1988). In the vasculature, ET<sub>A</sub> receptors residing on smooth muscle cells mediate prolonged vasoconstriction, whereas ET<sub>B</sub> receptors, which are located on the plasma membrane of endothelial cells, are primarily considered to cause NO-mediated vasodilatation (Yanagisawa, 1989). In addition, considerable evidence now also supports a role for the ET<sub>B</sub> receptor in clearance of plasma ET-1 from the circulation (Berthiaume, 2000). We have previously shown that agonist-induced internalization of the two receptor subtypes depends on a mechanism involving G protein-coupled receptor kinase, arrestin, clathrin, and dynamin (Bremnes, 2000). After internalization, however, the two receptor subtypes are targeted to different intracellular fates. The ET<sub>A</sub> receptor follows the recycling pathway through the pericentriolar recycling compartment and subsequently reappears on the plasma membrane, whereas the ET<sub>B</sub> receptor is directed to lysosomes for degradation (Bremnes, 2000). In terms of physiological effects, rapid recycling of the ET<sub>A</sub> receptor may provide basis for reestablishment of the signaling response, and thus, for the sustained vasoconstriction mediated by this receptor.

A key event of endocytic transport of signaling receptors is their sorting to either divergent recycling or degradative membrane pathways. The prevailing model proposes that internalized receptors are prevented from recycling by becoming sequestered and retained in multivesicular bodies (Sorkin, 2002). According to this model, recycling is considered the default destiny of membrane receptors not being targeted to lysosomes. The model is based mainly on studies of endocytic trafficking of the transferrin receptor and the epidermal growth factor

receptor (Mukherjee, 1997). Rapid recycling of these receptors has been reported even after all cytoplasmic residues have been removed (Mayor, 1993). This passive view of recycling has recently been challenged at least as far as GPCRs are concerned. Emerging evidence, including a previous report from our laboratory (Paasche, 2001), indicates that recycling of GPCRs is dependent on signals residing in the cytoplasmic carboxyl-terminal region (Cao, 1999; Hirakawa, 2003; Kishi, 2001; Tanowitz, 2003; Trejo, 1999). Analysis of carboxyl-terminally truncated ET receptors and ET<sub>A</sub>/ET<sub>B</sub> chimera revealed that lysosomal trafficking appears to be the "default" pathway without any requirement for cytoplasmic carboxyl-terminal sorting signals (Paasche, 2001). In contrast, recycling of the ET<sub>A</sub> receptor was dependent on a signal residing in the carboxyl-terminal tail of this receptor subtype. So far, however, the structural characteristics of a motif involved in sorting of a GPCR to the recycling pathway has been determined for the  $\beta_{\rm 2}$ adrenergic receptor ( $\beta_2$ -AR) only (Cao, 1999). The carboxyl-terminal end of the  $\beta_2$ -AR contains a structure or sequence pattern (DSSL) typically recognized by class1 PDZ (PSD-95/Dlg/ZO-1) domain proteins (Hall, 1998). The PDZ ligand of the β<sub>2</sub>-AR has been reported to bind EBP50/NHERF1 (Ezrin-radixin-moesin (ERM)-binding phosphoprotein-50/Na<sup>+</sup>/H<sup>+</sup> exchange regulatory factor 1), and to promote recycling of the  $\beta_2$ -AR by linking the receptor to the actin cytoskeleton through the ERM-binding domain of EBP50 (Hall, 1998; Cao, 1999). Furthermore, when grafted onto the carboxyl-terminal end of the  $\delta$ -opioid receptor ( $\delta$ -OR), this four-residue sequence (DSLL) was shown to re-route endocytosed  $\delta$ -OR from the degradative to the recycling pathway (Gage, 2001). On the other hand, the carboxyl-terminal PDZ ligand of the  $\beta_1$ -AR does not appear to be critical for recycling of this receptor subtype (Xiang, 2003). Furthermore, many receptors that are efficiently sorted to the recycling pathway, including the ET<sub>A</sub> receptor, lack classical carboxyl-terminal PDZ binding motifs (Bremnes, 2000; Hirakawa, 2003; Kishi, 2001;

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Tanowitz, 2003; Trejo, 1999). Yet, for several of these receptors the cytoplasmic carboxylterminal region has been shown to provide signals critical for sorting of the receptors to the recycling pathway (Bremnes, 2000; Hirakawa, 2003; Kishi, 2001; Tanowitz, 2003; Trejo, 1999). Thus, the aim of the present study was to identify and characterize the structural motif in the carboxyl-terminal tail of the  $ET_A$  receptor that dictates recycling of the receptor. Bioinformatics and mutational analysis of  $ET_A$  receptor sorting provided several lines of evidence that an internal PDZ ligand of the carboxyl-terminal region targets the receptor to the recycling pathway. Furthermore, additional evidence suggests that internal PDZ ligands may represent a more

general principle involved in sorting of many G protein-coupled receptors.

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## **Materials and Methods**

## **Plasmid Constructs**

Subcloning of human transferrin receptor cDNA into pcDNA1 and of the open reading frame of human ET<sub>A</sub> receptor into the mammalian expression vector pcDNA3 (pcDNA3-ET<sub>A</sub>) has been described previously (Bremnes, 2000). The hemagglutinin (HA) epitope-tagged dominant negative EBP50 truncation mutant EBP50ΔERM in pcDNA3 was generously provided by Dr. Mark von Zastrow, University of California, San Francisco (Cao, 1999).

## Construction of ET<sub>A</sub> Receptor Truncation Mutants

Truncation mutants of the ET<sub>A</sub> receptor with deletions of the cytoplasmic carboxyl-terminal tail were constructed in pcDNA3 by PCR directed mutagenesis. The truncation mutants ET<sub>A</sub>- $\Delta$ 417, ET<sub>A</sub>- $\Delta$ 406, ET<sub>A</sub>- $\Delta$ 401, ET<sub>A</sub>- $\Delta$ 398, ET<sub>A</sub>- $\Delta$ 394, and ET<sub>A</sub>- $\Delta$ 390 were made by replacement of D418, W407, G402, P399, M395, or S391, respectively, by stop codon and introduction of a *XbaI* restriction site downstream of the stop codon. The 3'-prime *Eco*RI-*XbaI* fragment of wild type ET<sub>A</sub> (pcDNA3-ET<sub>A</sub>) was replaced by the corresponding *Eco*RI/*XbaI*-digested PCR product to introduce the deletions indicated above. The cDNA constructs were verified by DNA sequence analysis using the dideoxy chain termination method. Construction of ET<sub>A</sub> receptor with in-frame carboxyl-terminal fusion of green fluorescent protein (ET<sub>A</sub>-GFP) in the mammalian expression vector pEGFP-N1 (BD Biosciences Clontech) has been described previously (Bremnes, 2000). ET<sub>A</sub>- $\Delta$ 406-GFP and ET<sub>A</sub>- $\Delta$ 401-GFP, i.e. ET<sub>A</sub> receptor truncation mutants with in-frame fusion of GFP at the carboxyl-terminal truncation site were made by introduction of a *Bam*H1 restriction site at the terminal amino acid in pcDNA3-ET<sub>A</sub>- $\Delta$ 406 and pcDNA3-ET<sub>A</sub>- $\Delta$ 401 using PCR-

directed mutagenesis. The EcoR1/BamH1 fragments of these modified vectors were subsequently transferred to EcoR1/BamH1-digested pEGFP-N1-ET<sub>A</sub>-GFP to generate pEGFP-N1-ET<sub>A</sub>- $\Delta$ 406-GFP and pEGFP-N1-ET<sub>A</sub>- $\Delta$ 401-GFP, respectively.

## Construction of ET<sub>A</sub> Receptor Point Mutants

ET<sub>A</sub> receptors with point mutations in the cytoplasmic carboxyl-terminal tail were constructed by site-directed circular mutagenesis using pcDNA3-ET<sub>A</sub> as template and two complementary oligonucleotide primers harboring the desired mutation. The QuickChange site-directed mutagenesis kit (Stratagene) for circular mutagenesis was employed, and mutants were selected by *Dpn*1 and verified by DNA sequence analysis using the dideoxy chain termination method.

## **Cell Lines and Transfection**

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CHO-K1 cells (ATCC no. CRL-61) were maintained, propagated, and transfected as previously described (Bremnes, 2000; Paasche, 2001). For fluorescence microscopy experiments, cells were plated onto glass coverslips before transfection. For cytosolic Ca<sup>2+</sup> determination, CHO cells were grown within a 10 mm diameter polyethylene cylinder attached to the central part of a glass coverslip.

## Kinetics of [125I]-ET-1 Internalization and Recycling

Assay of internalization and externalization of wild type ET<sub>A</sub> receptor and the ET<sub>A</sub> receptor mutants were performed using receptor labeling with [125I]-ET-1 (Amersham Biosciences) as described previously (Bremnes, 2000; Paasche, 2001). [125I]-ET-1 binding to ET<sub>A</sub> receptor is essentially irreversible and the sum of surface-bound and internalized [125I]-ET-1 remained

constant during the assay. Thus, analysis of the fraction of receptors remaining intracellularly or on the surface of the plasma membrane as function of time enabled assay of receptor recycling as previously validated (Paasche, 2001).

## Radioligand Binding

Radioligand binding analysis of membrane preparations of transiently transfected CHO cells was performed according to Elshourbagy et al. (Elshourbagy, 1993) with minor modifications (Bremnes, 2000).

## Loading of CHO Cells with Fura-2 Acetoxymethyl Ester and Measurement of Cytosolic ${\rm Ca^{2+}}$ in Single Cells

Transiently transfected CHO cells on glass coverslips were washed once with Hank's balanced salt solution (HSS) and incubated in HSS solution containing 5 μM Fura-2 and 0.025% Pluronic F127 for 30 min at 37 °C. After incubation, the cells were washed once and then incubated at 37 °C for 30 min before assay of cytosolic Ca<sup>2+</sup> concentrations in response to stimulation with ET-1. The methodology and software for data acquisition and analysis of cytosolic Ca<sup>2+</sup> concentrations in single cells have been described previously (Røttingen, 1995). In brief, the cells were illuminated at the excitation wavelengths 345 nm and 385 nm 2.5 times per second. Fluorescent images were collected by a CCD video camera, stored on a videotape and simultaneously digitized by a framegrabber controlled by a computer. Cytosolic Ca<sup>2+</sup> concentrations were calculated from the ratio between fluorescence intensities at excitation wavelengths 345 nm and 385 nm at user-selected squares covering single cells. After recording of baseline fluorescence,

ET-1 was added (100 nM) and collection of fluorescent images were continued in order to measure agonist-stimulated Ca<sup>2+</sup> levels.

## **Phosphoinositide Hydrolysis**

ET-stimulated inositol phosphate generation was assayed in transfected CHO cells metabolically labeled with 3μCi myo-[<sup>3</sup>H]inositol as described previously (Paasche, 2001).

## Western Blot Analysis

To verify expression of the HA epitope-tagged carboxyl-terminal truncation mutant of EBP50, EBP50 $\Delta$ ERM (Cao, 1998), Western blot analysis of samples from untransfected and transiently transfected CHO cells was performed. The immunoblots were processed as described previously (Bremnes 2000), using anti-HA IgG<sub>2a</sub> and subsequently, horseradish peroxidase-conjugated antimouse IgG, and the ECL system (Amersham Biosciences) for detection of immunoreactivity.

## Analysis of Intracellular Trafficking of Receptor-GFP Fusion Proteins by Confocal Laser Scan Microscopy

For analysis of ET receptor trafficking, transfected CHO cells attached to glass coverslips were placed on ice, washed, and preincubated with cycloheximide (10 mg/ml) in HSS for 30 minutes to prevent accumulation of newly synthesized ET<sub>A</sub>-GFP in the Golgi apparatus. The cells were subsequently incubated with ET-1 (100 nM) for 1 h at 4 °C in binding buffer (HSS containing 20 mM HEPES pH 7.4, 0.2% bovine serum albumin, 0.1% glucose and 10 mg/ml cycloheximide). Endocytosis of agonist-stimulated ET<sub>A</sub>-GFP or GFP-tagged ET<sub>A</sub> receptor mutants was initiated by rapidly changing the cell medium to 37 °C. The cells were incubated for various periods of

time at 37 °C in order to investigate the intracellular trafficking pathways of the different receptor mutants. The cells were subsequently fixed in 4% paraformaldehyde before mounting onto object slides using Mowiol (Hoechst). Cells were examined with Leica TCS SP confocal microscope equipped with an Ar (488 nm) and two He/Ne (543 and 633 nm) lasers. A Plan apochromat  $100 \times 1.4$  oil immersion objective was used. Multi-labeled images were acquired sequentially with a CCD camera. Images were processed and overlaid using Photoshop 7.0.

## Uptake of Red Fluorescent Transferrin and LDL

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Loading of the pericentriolar recycling compartment with 25 µg/ml tetramethylrhodamine-conjugated transferrin (Tf-Rhod; Molecular Probes) was performed in CHO cells cotransfected with ET receptor-GFP cDNA and cDNA for the human transferrin receptor as described previously (Bremnes, 2000). For labeling of lysosomes, LDL-DiI (Molecular Probes) was bound to the cells simultaneously with ET-1 at 4 °C. Internalization of the agonist-bound receptor was subsequently performed at 37 °C for 60 min allowing receptor-bound LDL to enter lysosomal compartments. The cells were fixed in 4% paraformaldehyde before mounting of the samples and investigation by confocal laser scan microscopy.

Structure Predictions of The Cytoplasmic Carboxyl-Terminal Region of The ET<sub>A</sub> Receptor
The ICM Pro 3.0 program (Molsoft L.L.C., La Jolla, CA; available at <a href="www.molsoft.com">www.molsoft.com</a>) was
utilized for modeling and computer graphics visualizations (Abagyan, 1994). Using a crystal
structure of bovine rhodopsin as structural template (Palczewski, 2000), a three-dimensional
model of the human ET<sub>A</sub> receptor (Swiss-Prot # P25101) was constructed. In order to obtain
evidence of a structural motif within the cytoplasmic carboxyl-terminal region of the ET<sub>A</sub>

receptor, the PDB database of The Research Collaboratory for Structural Bioinformatics (<a href="www.rcsb.org">www.rcsb.org</a>) was searched for proteins with sequence similarity to the Q390 - N427 segment of the ETA receptor. The National Center for Biotechnology Information (NCBI) BLAST software (Altschul, 1997) maintained by the Swiss Institute of Bioinformatics (<a href="http://us.expasy.org/tools/blast/">http://us.expasy.org/tools/blast/</a>) was employed to identify proteins in the PDB database with amino acid sequence similarities to the Q390 - N427 segment of the ETA receptor. The structures of any identified proteins were retrieved and visualized computer graphically using the ICM software.

## Probing The Cytoplasmic Carboxyl-Terminal Region of Mammalian GPCRs for Internal PDZ Interaction Motifs

Amino acid sequences for ~300 different mammalian family A GPCRs excluding the olfactory receptors, 15 different family B GPCRs and 22 different family C GPCRs were obtained from the Swiss Prot and TrEMBL protein sequence databases (Abagyan, 1994). The ClustalX 1.82 protein analysis program (CSC - Scientific Computing Ltd) was first used to obtain crude alignments of these sequences. The resulting alignments were further edited and analyzed with the BioEdit computer program. Thereafter, the carboxyl-terminal regions were retrieved from these alignments, starting from the positions corresponding to V372 of the human ET<sub>A</sub> receptor (family A), E398 of the human secretin receptor (family B) and E865 of the human GABA<sub>B1</sub> subunit (family C). The ICM 3.0 program was employed to search for putative internal PDZ ligand motifs in the carboxyl-terminal tail sequences using the following amino acid sequence pattern {[?]\5}[ST][?][IVLMF]{[?]\3-6}[ST]{[?]\4}[RKH]{[?]\3}. {[?]\n} represents any n consecutive amino acids. [ST] represents either a serine or a threonine residue. The amino-terminal [ST] of

the sequence pattern corresponds to the  $P_{-2}$  position of the internal PDZ pseudopeptide. [IVLMF] represents a hydrophobic residue ( $P_0$  position), i.e. isoleucine, valine, leucine, methionine, or phenylalanine. [RKH] represents any positively charged residue, i.e. arginine, lysine or histidine.

## **Results**

Functional Characterization of Wild Type  $ET_A$  Receptor,  $ET_A$  Receptor Mutants and  $ET_A$ -GFP

Stimulation of CHO cells transfected with either wild type ET<sub>A</sub> receptor or the truncation mutants  $ET_A$ - $\Delta 406$  or  $ET_A$ - $\Delta 401$  with ET-1 (100 nM) generated rapid elevations in intracellular calcium concentrations (Fig. 1A). Untransfected cells, however, did not respond to stimulation with ET-1. Peak calcium concentrations appeared to be slightly higher in cells transfected with the ET<sub>A</sub>- $\Delta 406$  or ET<sub>A</sub>- $\Delta 401$  truncation mutants than in ET<sub>A</sub> wild type transfected cells, but these differences were not statistically significant. Thus, the ET<sub>A</sub> receptor mutants were found to have similar capacities for generation of second messenger responses as their wild type counterpart. In frame carboxyl-terminal fusion of green fluorescent protein (GFP) was employed to investigate the intracellular trafficking pathways of ET<sub>A</sub> receptor. As demonstrated in Fig. 1B, internalization of [1251]-ET-1 in CHO cells transfected with ET<sub>A</sub>-GFP showed that internalization of agonistbound ET<sub>A</sub>-GFP proceeded with similar kinetics as that of wild type ET<sub>A</sub> receptor. Furthermore, the capacity of ET<sub>A</sub>-GFP to mediate ET-1-stimulated inositol phosphate generation, as shown in Fig.1C, was nearly identical to that of wild type ET<sub>A</sub> receptor. Agonist-induced endocytosis of ET<sub>A</sub>-GFP was also examined with confocal laser scan microscopy. CHO cells transfected with ET<sub>A</sub>-GFP were incubated with ET-1 (100 nM) for 1h at 4 °C. Unbound ligand was subsequently removed and endocytosis of agonist-bound receptor was initiated by changing of the medium to 37 °C. As seen in Fig. 1D, ET<sub>A</sub>-GFP is initially localized predominantly at the plasma membrane. However, within minutes after agonist-induced endocytosis of ET<sub>A</sub>-GFP, the receptor was detected in a spot-like structure near the nucleus demonstrating agonist-dependent intracellular

transport. Fig. 1E demonstrates endocytosis and intracellular trafficking of Tf-Rhod in CHO cells transfected with human transferrin receptor. As shown, Tf-Rhod appears to be rapidly internalized and transported through endosomal vesicles to a perinuclear endosomal compartment. After 15 min Tf-Rhod is almost entirely located in the spot-like perinuclear structure previously demonstrated to represent the perinuclear recycling compartment (Presley, 1993). Accordingly, for optimal labeling of the recycling compartment in subsequent experiments, agonist-dependent endocytosis was allowed to proceed for 15 min.

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To identify the domain in the carboxyl-terminal tail of the ET<sub>A</sub> receptor that mediates recycling, a series of deletion mutants were constructed by introducing stop codons at positions D418 (ET<sub>A</sub>- $\Delta$ 417), W407 (ET<sub>A</sub>- $\Delta$ 406), G402 (ET<sub>A</sub>- $\Delta$ 401), P399 (ET<sub>A</sub>- $\Delta$ 398), M395 (ET<sub>A</sub>- $\Delta$ 394), or S391 (ET<sub>A</sub>- $\Delta$ 390). The ligand binding characteristics of these truncated receptors did not differ significantly from that of wild type ET<sub>A</sub> receptor (Table 1). As shown in Fig. 2, the ET<sub>A</sub>- $\Delta$ 406 truncation mutant showed a biphasic internalization curve similar to that of wild type ET<sub>A</sub> receptor. The declining part of this curve pattern (representing increasing fractions of receptor reappearing at the surface of the plasma membrane) has previously been shown to reflect recycling of the ET<sub>A</sub> receptor (Paasche, 2001). The ET<sub>A</sub>- $\Delta$ 417 mutant (not shown) also retained a biphasic internalization curve and hence the capability to undergo recycling. For the ET<sub>A</sub> receptor mutants with truncations amino-terminal to Q406 (ET<sub>A</sub>- $\Delta$ 401, ET<sub>A</sub>- $\Delta$ 398, ET<sub>A</sub>- $\Delta$ 394, ET<sub>A</sub>- $\Delta$ 390), however, the time course of internalization shifted from a biphasic to a monophasic event as shown in Fig. 2. We have previously shown that the signal for recycling resides distal to amino

acid O390 of ET<sub>A</sub> (Paasche, 2001). Thus, the current observations provide evidence that the amino acid sequence 390 – 406 contains the critical information conferring the ability of the receptor to undergo recycling. To further investigate the intracellular trafficking pathways of the truncation mutants  $ET_A$ - $\Delta 406$  or  $ET_A$ - $\Delta 401$ , green fluorescent protein (GFP) was fused to the carboxyl-terminal end of these receptor mutants for analysis of subcellular localization using fluorescence microscopy. ET<sub>A</sub>- $\Delta$ 406-GFP and ET<sub>A</sub>- $\Delta$ 401-GFP were transiently transfected into CHO cells and subjected to analysis of agonist-stimulated internalization. The GFP-tagged wild type ET<sub>A</sub> receptor (ET<sub>A</sub>-GFP) was analyzed in parallel for comparison. As shown in Fig. 3, after 15 min of ET-1 induced internalization, both ET<sub>A</sub>-GFP and ET<sub>A</sub>-Δ406-GFP accumulated in a perinuclear spot-like structure that was co-labeled with red fluorescent transferrin i.e. a prototypical marker of the perinuclear recycling compartment (Presley, 1993). ET-induced internalization of ET<sub>A</sub>-Δ401-GFP, however, was associated with spread endosome-like vesicular structures throughout the cytoplasm with only minor colocalization in the recycling compartment. This finding indicates that the transport route of  $ET_A$ - $\Delta 401$  differs from that of  $ET_A$ -GFP and ET<sub>A</sub>-Δ406-GFP. To investigate whether the spread vesicular distribution of internalized ET<sub>A</sub>- $\Delta 401$  was consistent with transport to lysosomes, the lysosomal compartments of ET<sub>A</sub>- $\Delta 406$ -GFP- or ET<sub>A</sub>-Δ401-GFP-transfected cells were labeled with red fluorescent LDL (DiI-LDL) (Handley, 1981). As shown in Fig. 4, neither ET<sub>A</sub>-GFP nor ET<sub>A</sub>-Δ406-GFP colocalized with red fluorescent LDL after 60 min of internalization. However, the spread endosome-like structures of internalized ET<sub>A</sub>-Δ401-GFP demonstrated extensive colocalization with red fluorescent LDL (Fig. 4).

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Searching of the PDB database revealed that specific domains of rat CD2 (PDB # 1hng), human interferon- $\gamma$  receptor  $\alpha$  chain (IFN- $\gamma$ R $\alpha$ ) (PDB # 1fyh), and human neuronal nitrogen oxide synthase (nNOS) (PDB # 1qau) displayed significant similarities to the Q390-D411 peptide segment of ET<sub>A</sub> (Fig. 5A). F103-Q124 of human nNOS, I131-T151 of human IFN-γRα, and Y129-N149 residues of rat CD2 displayed 36%, 40%, and 40% amino acid similarities. respectively, to the Q390-D411 peptide segment of the human ET<sub>A</sub> receptor. Furthermore, the identified segments of CD2, IFN-γRα, and nNOS all contained anti-parallel β-strands adopting βfinger structures. Indeed, the β-finger of human nNOS has been reported to function as an internal PDZ-ligand (Christopherson, 1999). In this motif, the proximal β-strand (also referred to as the pseudopeptide motif) structurally mimics distal carboxyl-terminal PDZ-ligands in its sequence-specific interaction with one of the PDZ domains of syntrophin. Analysis of the crystal structure of the PDZ-ligand of human nNOS in complex with the PDZ domain of syntrophin, revealed that the hydrophobic phenylalanine at the end of the proximal  $\beta$ -strand (G-strand) is exposed by a sharp  $\beta$ -turn and mimics the free carboxylate group normally required for the hydrophobic P<sub>0</sub> position of a carboxyl-terminal ligand (Harris, 2001). Based on sequence similarities between the ET<sub>A</sub> receptor and the proteins shown in Fig. 5A and B, and using the structure of nNOS as a template, a three-dimensional model of the K392-H410 peptide sequence of the ET<sub>A</sub> receptor was built. In the predicted structure, as shown in Fig. 5C, the proximal βstrand was found to contain a sequence conforming to Class 1 PDZ ligands with V398 and T396 of ET<sub>A</sub> representing the P<sub>0</sub> and the P<sub>-2</sub> positions, respectively, of the putative PDZ ligand.

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## Intracellular Trafficking of $ET_A$ Receptors with Mutations that Perturb The PDZ Recognition Sequence or Structural Integrity of The $\beta$ -Finger

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To provide further evidence that the 390-406 peptide sequence of ET<sub>A</sub> possesses PDZ ligand properties, we used site-directed mutagenesis at residues that would be crucial to the structural integrity of the putative ligand. Fig. 6A and B show the kinetics of agonist-induced internalization of ET<sub>A</sub> wild type and ET<sub>A</sub> receptor mutants with substitutions perturbing the proximal strand of the  $\beta$ -finger, i.e. the strand containing the ligand directly interacting with a PDZ domain-containing protein. Agonist-stimulated internalization of the ET<sub>A</sub> T396A/S397A mutant was substantially impaired as compared to its wild type counterpart (Fig. 6A). Furthermore, the internalization curve was shifted from a biphasic to a monophasic event, which indicates that mutations of the P<sub>-1</sub> and P<sub>-2</sub> positions of the putative PDZ ligand prevent recycling. Mutations of V398 which represents the hydrophobic P<sub>0</sub> position into alanine or glycine (Fig. 6B) did not affect the initial rate of internalization, but abolished the biphasic curve pattern indicating severely perturbed recycling. In order to test our structural model further, mutations at residues that would perturb the distal strand of the  $\beta$ -finger were also made. As shown in Fig. 6C and D, agonist-induced internalization of the ET<sub>A</sub>-T403A/S404A mutant exhibited a monophasic internalization curve similar to that of the ET<sub>A</sub>-T396A/S397A mutant where both the initial rate of internalization and recycling capabilities were affected. The I405A and I405G mutations had less dramatic, albeit similar effects as V398A and V398G mutations. Inhibition of recycling was more pronounced upon mutation to alanine but was observed for both the ET<sub>A</sub>-I405A and ET<sub>A</sub>-I405G mutants (Fig. 6D). To study the intracellular trafficking pathways of these point-mutated receptors, GFP fusion proteins of ET<sub>A</sub>-T396A/S397A and ET<sub>A</sub>-T403A/S404A mutants were made. Agonist-stimulated internalization of receptor was investigated together with simultaneous

uptake of red fluorescent Tf-Rhod or DiI-LDL as described under *Materials and Methods*. As shown in Fig. 3, neither the ET<sub>A</sub>-T396A/S397A-GFP nor ET<sub>A</sub>-T403A/S404A-GFP mutants colocalized with transferrin receptor in the pericentriolar recycling compartment as opposed to ET<sub>A</sub>-GFP. However, in cells transfected with ET<sub>A</sub>-T396A/S397A-GFP or ET<sub>A</sub>-T403A/S404A-GFP, labeled with DiI-LDL, and stimulated with ET-1 for 60 min extensive colocalization of receptor and DiI-LDL was detected in lysosomes (Fig. 4). Taken together, these results support our hypothesis that recycling of ET<sub>A</sub> receptor is mediated via PDZ domain interactions.

Analysis of T396, S397, T403 and S404 of ET<sub>A</sub> as Potential Sites of Phosphorylation

As described above, substitutions of serine and threonine in the ET<sub>A</sub>-T396A/S397S or ET<sub>A</sub>-T403A/S404A double mutants affected not only the recycling capabilities but also the rate of internalization. Based on these observations we examined whether the indicated threonine and serine residues were sites of phosphorylation by introducing negative charge in the form of an aspartate residue that could mimic a phosphorylated state in these positions. We constructed mutants of ET<sub>A</sub>-Δ406 and "full length" ET<sub>A</sub> receptors where T396, S397, T403 and S404 were substituted with aspartate. Mutations into alanine were also made for parallel comparison. As shown in Fig. 7A and B, the aspartate and alanine substitutions of "full length" ET<sub>A</sub> and ET<sub>A</sub>-Δ406 abolished the biphasic internalization kinetics and significantly reduced the rate of internalization in a near identical manner. From these observations we conclude that receptor internalization and recycling are similarly affected upon mutation to alanine or aspartate.

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## Effect of A Dominant Negative EBP50 Mutant on Agonist-Stimulated Internalization and Recycling of The $\mathrm{ET}_A$ Receptor

To investigate whether or not recycling of the  $ET_A$  receptor is dependent on EBP50, we used a dominant negative truncation mutant of EBP50 (EBP50 $\Delta$ ERM) that lacks the ERM-binding domain but retains the PDZ domain reported to be required for interaction with the  $\beta_2$ -AR (Hall, 1998). CHO cells were transfected with the  $ET_A$  receptor alone or together with EBP50 $\Delta$ ERM. Western blot analysis, as shown in Fig. 8, confirmed the expression of EBP50 $\Delta$ ERM in the transiently transfected cells. The biphasic internalization kinetics of the  $ET_A$  receptor, which indicates recycling, was not altered upon expression of EBP50 $\Delta$ ERM (Fig. 8).

## Probing for Internal PDZ Ligand Motifs in GPCRs

A specific sequence pattern required to fulfill the structural criteria of an internal PDZ ligand (described under *Materials and Methods*) were defined and used to construct an algorithm that could be employed to screen GPCRs. Analysis of the carboxyl-terminal regions of more than 300 mammalian family A, B and C GPCRs, starting after predicted endpoints of transmembrane helix 7, revealed 35 different human GPCRs that fulfilled the sequence criteria of internal PDZ ligand motif. Fig. 9 shows alignment of the identified carboxyl-terminal sequences of 27 human GPCRs identified in the above analysis for which the cognate ligand is known and 8 orphan GPCRs.

## Discussion

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In the present study, the carboxyl-terminal motif of  $ET_A$  receptor that dictates receptor recycling has been identified and characterized. Structural modeling and mutagenesis of the cytoplasmic carboxyl-terminal region of  $ET_A$  receptor revealed that receptor recycling is mediated by a motif with the structural characteristics of an internal PDZ ligand.

Composite data from studies of ET<sub>A</sub> and ET<sub>B</sub> receptor chimera where the carboxylterminal tail of the receptors were interchanged (Paasche, 2001), and from studies of increasing truncations of the carboxyl-terminal region of ET<sub>A</sub>, revealed that the 17 amino acid polypeptide segment Q390-Q406 of ET<sub>A</sub> receptor constitutes the minimal peptide sequence required to provide a functional signal for recycling. Surprisingly, the identified polypeptide segment was part of a slightly larger segment (Q390-D411) of the carboxyl-terminal region of ET<sub>A</sub> that displayed striking similarities to peptide sequences of nNOS, IFN-γRα and CD2 for which a common secondary and tertiary structure had been determined. The shared structure of the latter proteins consisted of anti-parallel  $\beta$ -strands forming  $\beta$ -fingers that in nNOS have been reported to function as a PDZ ligand (Hillier, 1999). Although the crystal structure of the ET<sub>A</sub> receptor has not yet been determined, prediction of the secondary and tertiary structures of the Q390-D411 polypeptide segment of ET<sub>A</sub> receptor revealed that this segment may also consist of anti-parallel  $\beta$ -strands attaining  $\beta$ -finger conformation. The proximal and distal  $\beta$ -strands of the putative  $\beta$ finger could be supported by several hydrogen bonds between residues within the Q390-D411 polypeptide segment, one of which was a hydrogen bond typically found between the first and the fourth residue of the loop between the β-strands. The latter is considered crucial in stabilizing the β-turn (Coligan, 2004). Additional lines of evidence also reported in this study consistently supported involvement of the putative  $\beta$ -finger of  $ET_A$  in receptor recycling. Targeted mutations

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of residues within the proximal β-strand (ET<sub>A</sub>-T396A/S397A, ET<sub>A</sub>-V398A, ET<sub>A</sub>-V398G, ET<sub>A</sub>-T403A/S404A, ET<sub>A</sub>-I405A, ET<sub>A</sub>-I405G), or distal β-strand (ET<sub>A</sub>-T403A/S404A, ET<sub>A</sub>-I405A, ET<sub>A</sub>-I405G) all disrupted recycling of ET<sub>A</sub> receptor. These observations indicate that mutations that affect the structural integrity of the β-finger abolish receptor recycling. However, in order to provide unequivocal evidence of a PDZ-mediated mechanism of ET<sub>A</sub> receptor recycling, binding of the putative β-finger and a PDZ-containing protein would have to be demonstrated and implicated a role in receptor recycling. Strikingly, the proximal  $\beta$ -strand of the putative  $\beta$ -finger of ET<sub>A</sub> also contained a pseudopeptide sequence conforming to the canonical class I PDZ ligand similar to that of the proximal  $\beta$ -strand of the nNOS  $\beta$ -finger. According to the predicted  $\beta$ -finger of ET<sub>A</sub>, V398 represents the indispensable hydrophobic residue at the P<sub>0</sub> position of the putative PDZ ligand. As demonstrated in the present study, alanine substitution of the P-2, P-1, or P0 residues of the PDZ pseudopeptide ligand in the proximal β-strand essentially eliminated recycling of ET<sub>A</sub> receptor. These results correlate with the loss of function caused by alanine substitution of the corresponding PDZ pseudopeptide residues (T109A or F111A at the P<sub>-2</sub> and P<sub>0</sub> positions, respectively) of nNOS (Harris, 2001).

Comparing the putative  $\beta$ -finger of ET<sub>A</sub> receptor with that of nNOS also revealed conservation of a positive residue in the distal  $\beta$ -strand. R121 in the distal  $\beta$ -strand of nNOS forms a salt bridge with D62 and plays important role in binding of nNOS to the PDZ domain of syntrophin, presumably by stabilizing the  $\beta$ -finger formation (Harris, 2001). The specific function of K408 in the putative  $\beta$ -finger motif of ET<sub>A</sub> receptor, i.e. the positive residue aligning with R121 of the distal  $\beta$ -strand nNOS, is unclear since this residue is removed in the ET<sub>A</sub>- $\Delta$ 406 truncation mutant which is still capable of recycling. However, it has been reported that the  $\beta$ -

finger of nNOS is a fairly rigid structure, which even by itself, separate from nNOS, may retain some residual hairpin structure (Wang, 2000).

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Previous reports have shown that agonist-stimulated desensitization and internalization of ET<sub>A</sub> and ET<sub>B</sub> receptors depend on GRK-catalyzed phosphorylation of serine and threonine residues in the cytoplasmic carboxyl-terminal region (Bremnes, 2000; Freedman, 1997). However, the specific residues of agonist-occupied ET<sub>A</sub> receptors phosphorylated by GRK have yet to be determined. Conceivably, introduction of negative charge by phosphorylation of any serine or threonine residue of the carboxyl-terminal region may to some extent relieve intramolecular constraints sufficient to promote binding of β-arrestin with subsequent desensitization and internalization of the receptor. However, mimicking of ET<sub>A</sub> receptor phosphorylation by substituting the indicated threonine and serine residues with aspartic acid (ET<sub>A</sub>-T396D/S397D; ET<sub>A</sub>-T403D/S404D) did not lead to increased rates of agonist-induced internalization. Rather, the aspartic acid substitutions impaired agonist-induced internalization to similar extent as alanine substitution of the same residues. (ET<sub>A</sub>-T396A/S397A; ET<sub>A</sub>-T403A/S404A). Thus, the reduced initial rates of agonist-induced internalization of the ET<sub>A</sub> receptor mutants do not appear to be due to loss of GRK-catalyzed phosphorylation. A more plausible interpretation of the observed findings is that substitution of the serine and threonine residues of the proximal and distal β-strands disrupt the β-finger and thereby abolish ET<sub>A</sub> receptor recycling. As serially connected mechanisms, impaired endosomal trafficking may also impact on the initial step of endocytosis, i.e. agonist-induced internalization of the ET<sub>A</sub> receptor.

To our knowledge, this is the first demonstration implicating a putative internal PDZ ligand structure in endocytic sorting of a GPCR. An imminent issue then, was to what extent the putative internal PDZ ligand of ET<sub>A</sub> receptor might represent a more widely employed principal

in endocytic sorting of GPCRs. Interestingly, as many as 35 different GPCRs fulfilled the principal criteria of an internal PDZ ligand motif. However, our search parameters only entailed class I PDZ domains. In this respect, inclusion of parameters covering class II and III PDZ domain proteins could potentially identify even more GPCR with internal PDZ ligand motifs. As shown in Fig. 9, the loop connecting the antiparallel  $\beta$ -strands of the putative  $\beta$ -finger structures of the aligned GPCRs consists of varying number of residues. Indeed, the variation in loop length allowed in the search parameters for  $\beta$ -finger-like PDZ ligand motifs in GPCRs was supported by evidence of maintained PDZ binding affinity upon insertion of additional residues in the  $\beta$ -turn region of the PDZ ligand of nNOS (Harris, 2001).

Among the human GPCRs shown in Fig. 9 are several receptors that have been reported to follow intracellular recycling pathways, i.e. the  $\alpha_{1D}$ -adrenergic,  $AT_1$  angiotensin,  $NK_1$  neurokinin,  $Y_1$  neuropeptide, LH luteinizing hormone, 5-HT $_{2A}$  serotonin, SST $_3$  somatostatin, and  $V_{1A}$  vasopressin receptors (Kishi, 2001; Benya, 1994; Bhattacharyya, 2002; Chauvin, 2002; Garland, 1996; Gicquiaux, 2002; Hein, 1997; Innamorati, 1999; Kreuzer, 2001; McCune, 2000; Tseng, 1995; Marchese, 2001). Conversely, peptide sequences with similarity to the search parameters could not be identified in GPCRs sorting to lysosomes, e.g. the ET $_B$  receptor, PAR-1, CXCR4 chemokine receptor,  $\delta$ -opioid receptor and rat LH receptor (Bremnes, 2000; Kishi, 2001; Trejo, 1999; Gage, 2001; Marchese, 2001). Thus, the proposed model of internal PDZ ligand recognition appears to be a consistent mechanism for sorting GPCRs to the recycling pathway. The rat and human LH receptor homologs deserve special attention because of divergent sorting to lysosomes and endocytic recycling, respectively (Kishi, 2001). Interestingly, the primary structure of rat LH receptor differs from its human receptor homolog at residues critical to maintain the putative PDZ ligand-containing  $\beta$ -finger. Thus, rat LH receptor may be regarded as

a naturally occurring variant of human LH receptor that does not contain an operative PDZ ligand. This contention is supported by several reports of point mutations or deletion of amino acids in the carboxyl-terminal region of human LH receptor that divert the receptor from the recycling pathway to the lysosomal trafficking pathway (Kishi, 2001; Galet, 2004; Hirakawa, 2003). Indeed, these targeted amino acids were apparently all critical to maintain the PDZ pseudopeptide ligand or  $\beta$ -finger conformation according to our proposed model. Recently, a report by Ascoli and colleagues demonstrated that efficient recycling was also shown to be dependent on the upstream hydrophobic amino acid L683. According to the alignment in Fig. 9, L683 may represent the "0-position" of a putative PDZ ligand-containing  $\beta$ -finger formation.

Although sorting of GPCRs to the recycling pathway may depend on uniform mechanisms, different PDZ domain proteins may be involved. For example involvement of EBP50 does not appear to be a consistent finding (Cao, 1999; Kishi, 2001). In the present study a dominant negative deletion mutant of EBP50 (EBP50 $\Delta$ ), lacking the ERM binding domain, did not affect recycling of the ET<sub>A</sub> receptor despite massive expression of EBP50 $\Delta$ .

In the present study, bioinformatic and mutational analysis provide strong evidence that recycling of the ET<sub>A</sub> receptor is mediated by a motif with the structural characteristics of an internal PDZ ligand. This motif which is lacking in the carboxyl-terminal region of ET<sub>B</sub> endothelin receptors, provides a mechanism of the divergent sorting of the ET<sub>A</sub> and ET<sub>B</sub> receptor subtypes. Furthermore, we provide evidence that internal PDZ ligand motifs may represent a more general principle of endocytic sorting of GPCRs. The important challenge of the future will be to identify the protein that recognizes and binds the putative PDZ ligand motif of ET<sub>A</sub> receptor and directs the receptor to the recycling pathway.

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## **Footnotes**

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## **Legends for Figures**

Fig. 1 Functional characteristics of wild type ET<sub>A</sub> receptor, carboxyl-terminally truncated ET<sub>A</sub> receptors and ET<sub>A</sub>-GFP fusion protein. (A) Time course of cytosolic Ca<sup>2+</sup> concentrations in ET-1 stimulated CHO cells transiently transfected with wild type ET<sub>A</sub> (····) receptor or the ET<sub>A</sub> truncation mutants ET<sub>A</sub>- $\Delta$ 406 (—) and ET<sub>A</sub>- $\Delta$ 401 (—). The cells were loaded with Fura-2 as described under Materials and Methods. ET-1 (100 nM) was added 30 seconds after recording of light emission was initiated (arrow). Untransfected cells (–) stimulated with ET-1 were used as control of background levels. The data are mean of at least 80 cells from three independent experiments. (B) Internalization kinetics of [125I]-ET-1 in CHO cells transiently transfected with wild type  $ET_A$  receptor or  $ET_A$  receptor with carboxyl-terminal fusion of GFP. Data are mean  $\pm$ S.D. of three parallel wells and representative of at least three independent experiments. (C) Inositol phosphate accumulation in CHO cells transiently transfected with wild type ETA receptor or ET<sub>A</sub>-GFP stimulated in the absence or presence of ET-1 (100 nM). Data are mean  $\pm$ S.D. of three parallel wells and representative of at least three independent experiments. (D) Photomicrographs demonstrating ET-1-induced endocytosis of ET<sub>A</sub>-GFP in CHO cells. CHO cells on cover glass were transiently transfected with ETA-GFP and analysed by confocal laser scan microscopy at 0, 15 and 60 min after initiation of ET-1 induced internalization. Cells were pretreated with 10 mg/ml cycloheximide for 30 min before addition of ET-1 (100 nM) and during the entire uptake period to clear newly synthesized receptor-GFP fusion protein from Golgi. The cells were subsequently fixed in paraformaldehyde at 4 °C and investigated by confocal laser scan microscopy. (E) Internalization of Tf-Rhod in CHO cells transfected with human transferrin

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receptor. The cells were fixed in paraformaldehyde at the indicated time points after addition of Tf-Rhod and analyzed by confocal laser scan microscopy. Note that Tf-Rhod becomes highly

enriched in a perinuclear compartment after 15 minutes of internalization.

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Fig. 2 Intracellular trafficking of ET<sub>A</sub> receptor with increasing truncation of the carboxylterminal region. Internalization kinetics of [ $^{125}$ I]-ET-1 in CHO cells transiently transfected with wild type ET<sub>A</sub> ( $\bullet$ ) or the ET<sub>A</sub> receptor truncation mutants ET<sub>A</sub>- $\Delta$ 406 ( $\nabla$ ), ET<sub>A</sub>- $\Delta$ 401 ( $\nabla$ ) and ET<sub>A</sub>- $\Delta$ 390 ( $\circ$ ). 24 hours after transfection CHO cells were incubated with [ $^{125}$ I]-ET-1 (25 pM in 1 ml) for 3 h at 4 °C to label surface ET<sub>A</sub> receptors. The internalization assay was initiated by rapid transfer of the cells to 37 °C for the time indicated. The fraction of intracellular receptor at each time point was measured as described under *Materials and Methods*. Inlet: Agonist-bound receptor (fmol/well) at plasma membrane surface at t = 0. Data are mean  $\pm$  S.D. of three parallel wells and representative of at least three independent experiments.

Fig. 3 Subcellular localization of agonist-stimulated GFP-tagged ET<sub>A</sub> wild type receptor and ET<sub>A</sub> receptor with mutations affecting the putative carboxyl-terminal β-finger and internal PDZ ligand. Probing for receptor colocalization with transferrin in the recycling compartment. Photomicrographs of confocal laser scan microscopic analysis of CHO cells cotransfected with ET<sub>A</sub>-GFP, ET<sub>A</sub>-Δ406-GFP, ET<sub>A</sub>-Δ401-GFP, ET<sub>A</sub>-T396A/S397A-GFP, or ET<sub>A</sub> T403A/S404A-GFP and transferrin receptor. After binding of ET-1 (0.1 μM) at 4 °C, agonist-stimulated internalization of the ET receptors was analyzed after 15 min at 37 °C. Tf-Rhod was added during the 15 min period for simultaneous labeling of the perinuclear recycling

compartment. As shown in the overlays to the right, ET<sub>A</sub>-GFP and ET<sub>A</sub>- $\Delta$ 406-GFP display prominent colocalization with transferrin receptor. ET<sub>A</sub>- $\Delta$ 401-GFP, ET<sub>A</sub>-T396A/S397A-GFP or ET<sub>A</sub>-T403A/S404A-GFP on the other hand, exhibit spread vesicular distribution lacking colocalization with transferrin receptor. Bar is 10  $\mu$ M.

Fig. 4. Subcellular localization of agonist-stimulated GFP-tagged ET<sub>A</sub> wild type receptor and ET<sub>A</sub> receptor with mutations affecting the putative carboxyl-terminal β-finger and internal PDZ ligand. Probing for receptor colocalization with LDL in lysosomes.

Photomicrographs of confocal laser-scan microscopic analysis of CHO cells transiently transfected with ET<sub>A</sub>-GFP, ET<sub>A</sub>-Δ406-GFP, ET<sub>A</sub>-Δ401-GFP, ET<sub>A</sub>-T396A/S397A-GFP, or ET<sub>A</sub>-T403A/S404A-GFP. Agonist-induced receptor internalization was performed for 60 min together with DiI-LDL to probe for lysosomal sorting. As shown in the overlays to the right, the spread vesicular distribution of ET<sub>A</sub>-Δ401-GFP, ET<sub>A</sub>-T396A/S397A-GFP and ET<sub>A</sub>-T403A/S404A-GFP coclocalized with LDL in lysosomes. ET<sub>A</sub>-GFP and ET<sub>A</sub>-Δ406-GFP could not be detected in DiI-LDL-labeled lysosomes. Bar is 10 μM.

Fig. 5 Analysis of the primary and secondary structures of the Q390-D411 peptide sequence of the ET<sub>A</sub> receptor. The Q390-D411 sequence of the ET<sub>A</sub> receptor aligned with rat CD2 (A), human INF- $\gamma$ R $\alpha$  (A), and human nNOS (B). The secondary structures of the aligning sequences of CD2, INF- $\gamma$ R $\alpha$  and nNOS as determined by X-ray crystallographic studies are indicated with arrows ( $\beta$ -strand structure) and lines (coil structure). The indicated sequence of nNOS is reported

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to form a  $\beta$ -finger motif operating as an internal PDZ ligand. C. Predicted structure of the K392-H410 peptide sequence of the ET<sub>A</sub> receptor. The predicted sequence consists of two antiparallel  $\beta$ -strands forming a  $\beta$ -finger. The proximal  $\beta$ -strand contains the sequence pattern recognized by class 1 PDZ domains, where the V398 represents the "P<sub>0</sub> position" of the pseudopeptide ligand.

Fig. 6 Intracellular trafficking of ET<sub>A</sub> receptors with mutations at selected residues that would perturb the β-finger or internal PDZ ligand function. A-D. Internalization kinetics of [ $^{125}$ I]-ET-1 in CHO cells transfected with the indicated ET<sub>A</sub> receptor mutants. All receptor mutants are compared with internalization kinetics of [ $^{125}$ I]-ET-1 in parallel cells transfected with wild type ET<sub>A</sub> receptor ( $\bullet$ ). A. ET<sub>A</sub>-T396A/S397A (O) (mutations at the P<sub>-1</sub> and P<sub>-2</sub> positions of the putative PDZ ligand). B. ET<sub>A</sub>-V398A (O) and ET<sub>A</sub>-V398G ( $\blacktriangledown$ ) (mutations at the P<sub>0</sub> position of putative PDZ ligand) ET<sub>A</sub>. C. ET<sub>A</sub>-T403A/S404A (O) (residues of the distal β-strand). D. ET<sub>A</sub>-I405A (O) and ET<sub>A</sub>-I405G ( $\blacktriangledown$ ). I405 is located in the distal β-strand region. Inlets: Agonist bound receptor (fmol/well) at plasma membrane surface at t = 0. The data are mean  $\pm$  S.D. of three parallel wells and representative of at least three independent experiments.

Fig. 7 Intracellular trafficking of ET<sub>A</sub> receptor mutants with substitution of the putative phosphorylation sites T396, S397, T403 and S404 with alanine or negatively charged aspartic acid residues. A. Internalization kinetics of [ $^{125}$ I]-ET-1 in CHO cells transiently transfected with ET<sub>A</sub>-T396D/S397D/T403D/S404D, ET<sub>A</sub>-T396A/S397A/T403A/S404, or wild type ET<sub>A</sub>. B Internalization kinetics of [ $^{125}$ I]-ET-1 in CHO cells transiently transfected with ET<sub>A</sub>  $\Delta$ 406/T396D/S397D/T403D/S404D, ET<sub>A</sub>- $\Delta$ 406/T396A/S397A/T403A/S404A, or ET<sub>A</sub>- $\Delta$ 406.

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The data are mean  $\pm$  S.D. of three parallel wells and representative of at least three independent experiments.

Fig. 8 Lack of effect of a dominant negative EBP50 deletion mutant (EBP50 $\Delta$ ERM) on ET<sub>A</sub> receptor internalization and recycling. Kinetics of [ $^{125}$ I]-ET-1 internalization in CHO cells transiently transfected with wild type ET<sub>A</sub> receptor alone ( $\bullet$ ) or together with EBP50 $\Delta$ ERM (O). The data are mean  $\pm$  S.D. of three parallel wells and representative of at least three independent experiments. Western blot analysis of CHO cells cotransfected with HA epitope-tagged EBP50 $\Delta$ ERM confirms expression of EBP50 $\Delta$ ERM in the transfected CHO cells. A single immunoreactive band migrated at the expected molecular mass of EBP50 $\Delta$ ERM as indicated.

Fig. 9 GPCRs containing putative internal PDZ ligand motifs. Alignment of putative PDZ ligand-containing  $\beta$ -finger structures in the carboxyl-terminal region of GPCRs identified by an algorithm searching for conserved sequence patterns similar to that of the  $\beta$ -finger of nNOS. The figure shows 35 different human GPCRs identified in the analysis as described in *Materials and Methods*. Orphan receptors are indicated by asterix. The receptors are grouped according to family A, B and C GPCRs and aligned with the  $\beta$ -finger of nNOS. Variation in allowed  $\beta$ -loop length is 3-6 amino acids. Gray shading indicates conserved amino acids. All receptors contain a putative PDZ ligand with conserved amino acids at positions conforming to the P<sub>-2</sub> and P<sub>-1</sub> positions of class 1 PDZ ligands. In addition, all GPCRs contain a conserved positively charged residue involved in stabilization of the  $\beta$ -finger structure. As indicated in the figure, a few receptors contain more than one putative  $\beta$ -finger in the carboxyl-terminal region.

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**Table 1.**Radioligand binding characteristics of wild type  $ET_A$  receptor and  $ET_A$  receptor mutants

pertubing the putative recycling motif. The properties of ET-1 binding, including the number of

binding sites, were determined in CHO cells transiently transfected with  $ET_A$  wild-type,  $ET_A$ - $\Delta 406$ ,  $ET_A$ - $\Delta 401$ ,  $ET_A$ - $\Delta 390$ ,  $ET_A$ -T396A S397A,  $ET_A$ -T403A S404A,  $ET_A$ - $\Delta 406$ -GFP and  $ET_A$ - $\Delta 401$ -GFP. Binding of <sup>125</sup>I-ET-1 was performed on membranes as described under *Materials*and *Methods* to determine the equilibrium dissociation constants ( $K_d$ ) and the maximal binding

( $B_{max}$ ) for the different receptors. The data are mean  $\pm$  S.D. of three parallels and representative

of at least three independent experiments.

Receptor	$K_d$	$B_{ m max}$
	pM	pmol/mg
ET <sub>A</sub> wt	$71 \pm 3$	$3.2 \pm 0.15$
$ET_A \Delta 406$	$100 \pm 7$	$1.6 \pm 0.11$
$ET_A \Delta 401$	$96 \pm 6$	$1.8 \pm 0.11$
$ET_A \Delta 390$	$87 \pm 8$	$1.7 \pm 0.55$
ET <sub>A</sub> T396A S397A	$81 \pm 2$	$3.3 \pm 0.08$
ET <sub>A</sub> T403A S404A	$89 \pm 6$	$3.4 \pm 0.24$
$ET_A \Delta 406$ -GFP	$63 \pm 4$	$6.0 \pm 0.41$
ET <sub>A</sub> Δ401-GFP	$139 \pm 6$	$2.3 \pm 0.10$

Fig.1

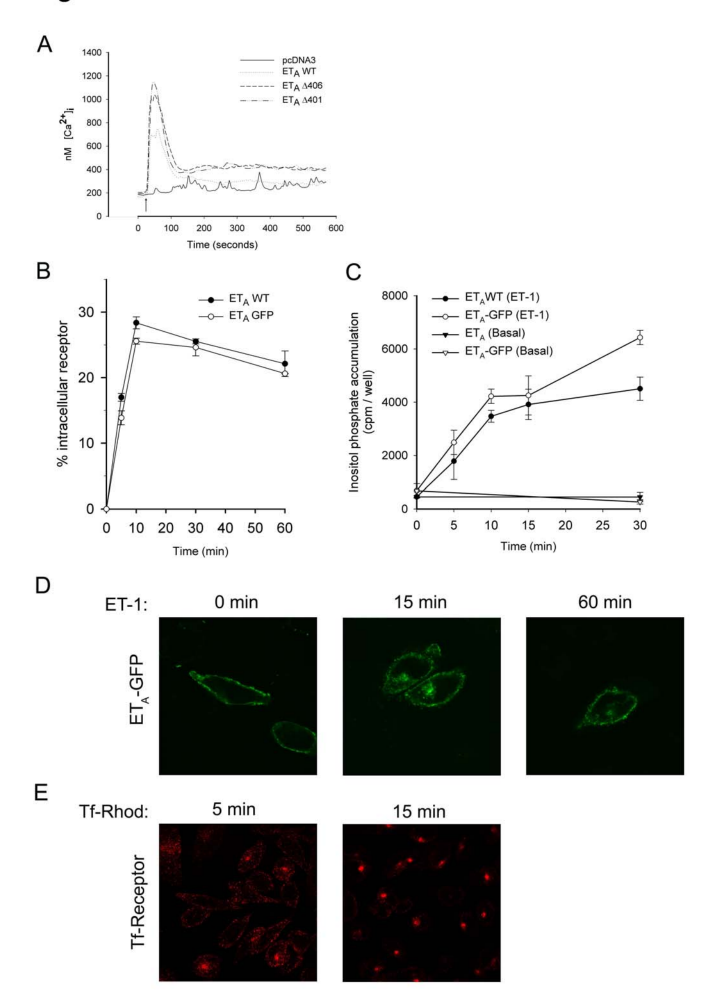
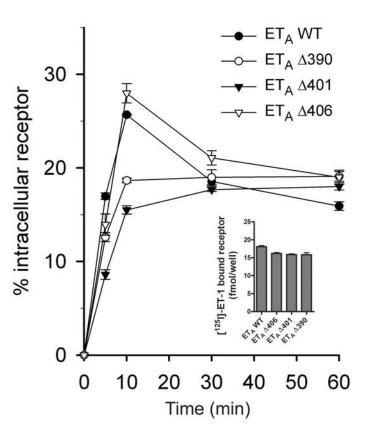
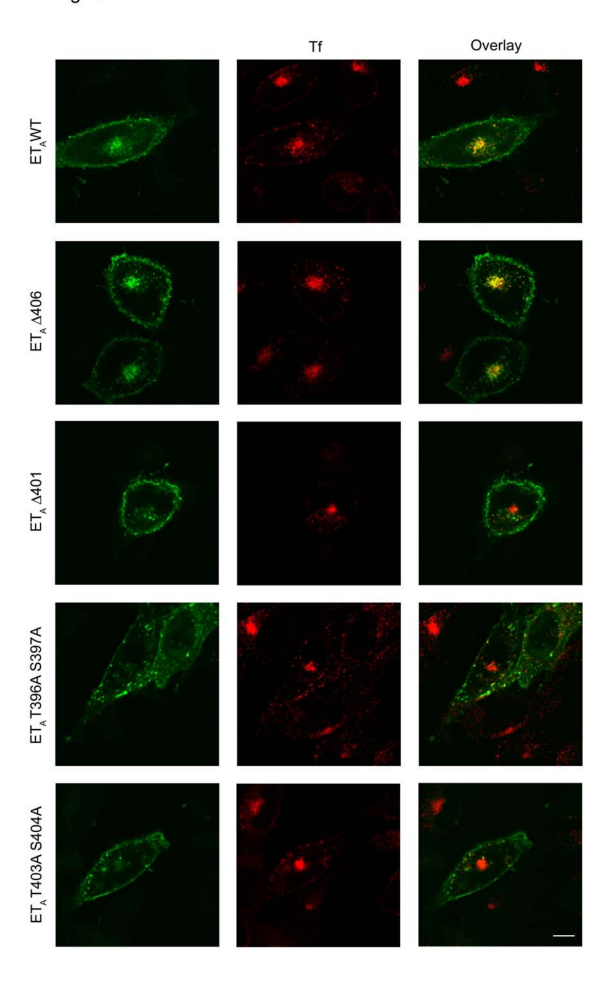


Fig.2





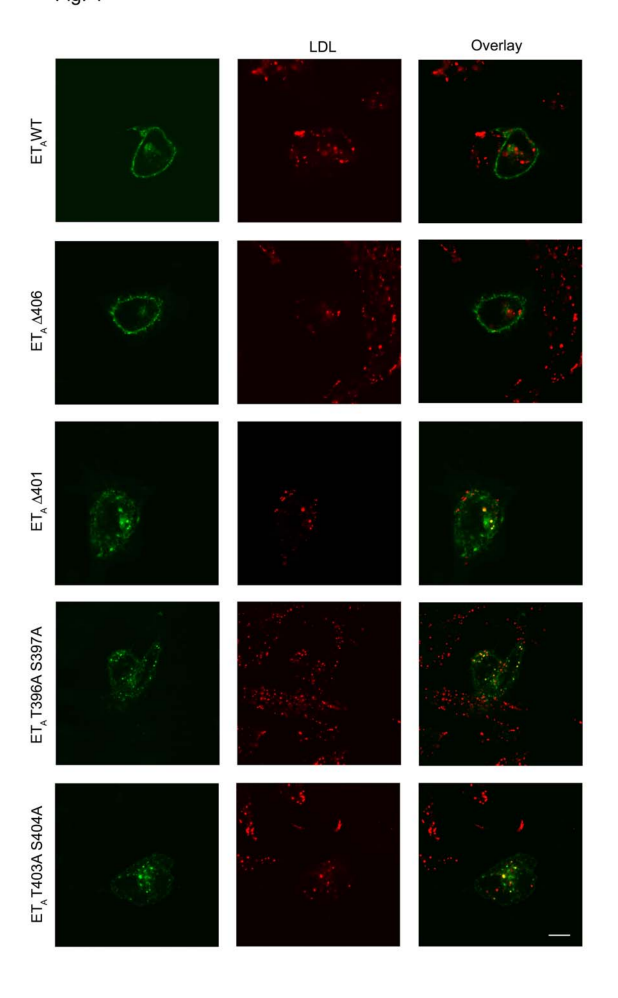
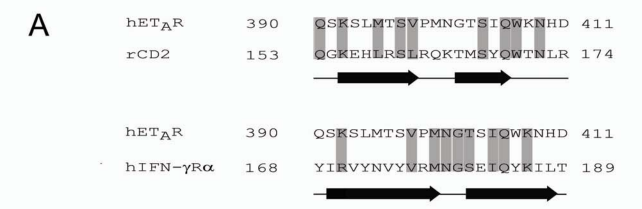
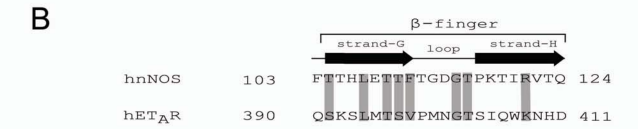


Fig.5





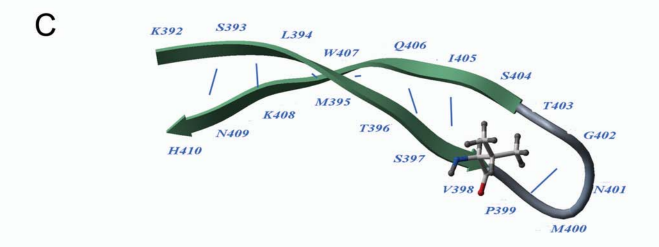
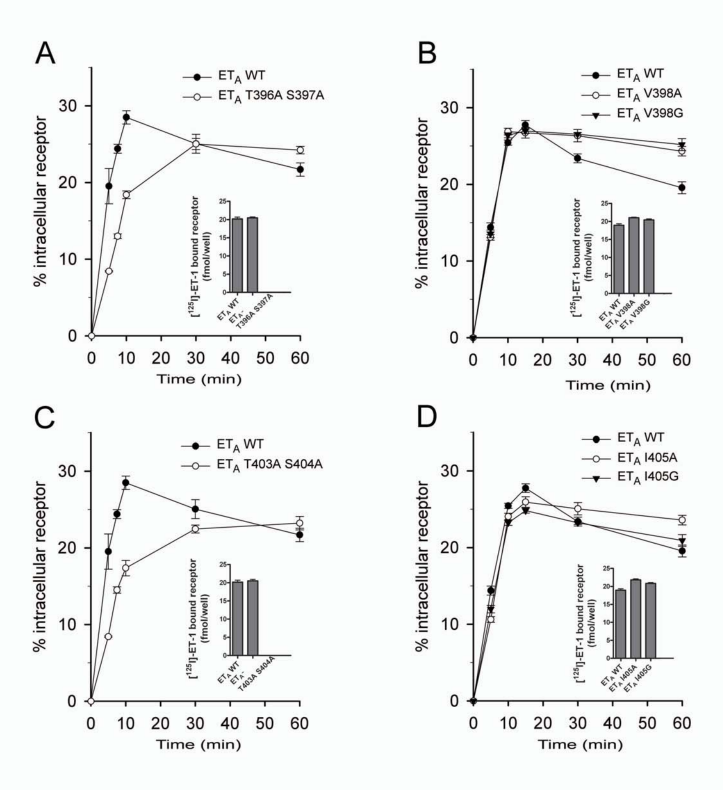


Fig.6





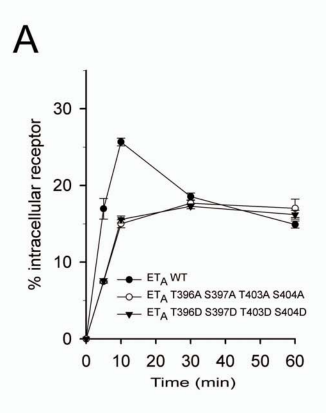
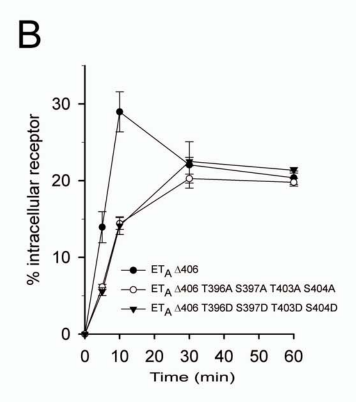
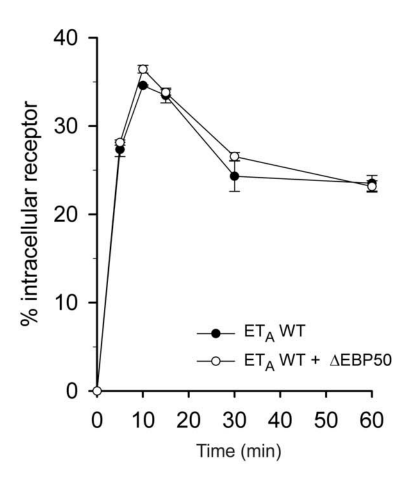
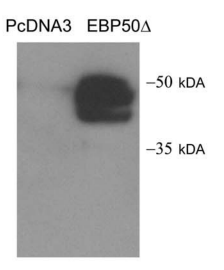


Fig.7







		eta-finger		
	20	$\beta$ -strand loop $\beta$ -strand	l	
nNOS	101	EGF <b>T</b> TH <b>L</b> E <b>T</b> T <b>F</b> TGD <b>GT</b> PKTI <b>R</b> VTQ	124	
A ET <sub>A</sub> R	388	CYQSKSLMTSVPMNGTSIQWKNHD	411	
AT <sub>1</sub> R	320	IPPKAKSHSNLSTKMSTLSYRPSD	343	
$AT_2R(1)$	323	NRFQQK <b>l</b> r <b>s</b> v <b>f</b> rvpi <b>t</b> wlqg <b>k</b> res	346	
$AT_2R(2)$	330	RSVFRVPI <b>TWL</b> -QGKRE <b>S</b> MSCR <b>K</b> SSS	354	
$V_{1A}R$	390	NNRSPTNSTGMWKDSPKSSKSIK	412	
BRS <sub>3</sub>	372	PGTGSIQMSEI-SVTSFTGCSVKQAE	396	
CCR <sub>6</sub>	336	CVRRKYKSSGFSCAGRYSENISRQTS	361	
Gal <sub>2</sub> R	325	AARGTHSGSVLERESSDLLHMSE	347	
$NK_1R$ (1)	344	TQG <b>S</b> VYKV <b>S</b> R <b>L</b> -ETTIS <b>T</b> VVGA <b>H</b> EEE	368	
$NK_1R$ (2)	370	EDGPKATPSSLDLTSNCSSRSDS	392	
$NMU_2R$	383	PCQSSMHNSHL-PTALSSEQMSRTNY	407	
$Y_1R$ (1)	345	DDYETIAMSTMHTDVSKTSLKQAS	368	
$Y_1R$ (2)	354	TMH <b>T</b> DVSK <b>T</b> S <b>L</b> KQA <b>S</b> PVAF <b>K</b> KIN	376	
$Y_4R$	347	EESEHLPLSTVHTEVSKGSLRLSG	370	
NPFF <sub>1</sub> R	373	FVVVRPSD <b>S</b> G <b>L</b> PSESGP <b>S</b> SGAP <b>R</b> PGR	398	
NPFF <sub>2</sub> R	477	TSNQLVQE <b>S</b> T <b>F</b> QNPHGE <b>T</b> LLYR <b>K</b> SAE	502	
SST <sub>3</sub> R		GKEMNGRVSQITQPGTSGQERPPS	390	
LHR		PSQSTLKLSTLHCQGTALLDKTRY	696	
$lpha_{ exttt{1D}}$ AR (1)		LGPFRRPTTQLRAKVSSLSHKIRA	523	
$lpha_{\text{1D}}$ AR (2)		AEAACAQR <b>S</b> E <b>V</b> EAV <b>S</b> LGVP <b>H</b> EVA	551	
$oldsymbol{eta}_3$ ar		AARPALFP <b>S</b> G <b>V</b> -PAARS <b>S</b> PAQP <b>R</b> LCQ	398	
5HT <sub>2A</sub> R		AKT <b>T</b> DNDC <b>S</b> M <b>V</b> ALGKQH <b>S</b> EEAS <b>K</b> DNS	461	
DP <sub>2</sub> R		LESVLVDDSELGGAGSSRRRRTSS	346	
LPA <sub>4</sub> R		INAHIRMESLFKTETPLTTKPSL	346	
S1P <sub>4</sub> R		HSGASTTDSSLRPRDSFRGSRSLS	368	
GPR75*		CGPSHSKESMVSPKISAGHQHCGQ	469	
Н963*			310	
HM74*		NSGEPWSPSYLGPTSNNHSKKGH	370	
CMKLR1* (1)		KKFKVA <b>L</b> F <b>S</b> R <b>L</b> VNAL <b>S</b> EDTG <b>H</b> SSY	347	
CMKLR1* (2)		SYP <b>S</b> HRSF <b>T</b> K <b>M</b> SSMNER <b>T</b> SMNE <b>R</b> ETG	371	
CMKLR1* (3)		SHRSFTKMSSMNERTSMNERETG	371	
GPR25*		LARRISSASSL-SRDDSSVFRCRAQA	352	
SALPR* (1)		REFRKA <b>L</b> K <b>S</b> L <b>L</b> WRIASP <b>S</b> ITSM <b>R</b> PFT	419	
SALPR* (2)		WRIASPSI <b>T</b> SM-RPFTA <b>T</b> TKPE <b>H</b> EDQ	427	
LGR <sub>5</sub> *		VWTRSKHPSLMSINSDDVEKQSC	824	
B PAC <sub>1</sub> R		ASSGVNGGTQL-SILSKSSSQIRMSG	460	
$VPAC_1R$ (1)		GSNGATCS <b>T</b> Q <b>V</b> SMLTRV <b>S</b> PGAR <b>R</b> SSS	449	
$VPAC_1R$ (2)		GATCSTQVSMLTRVSPGARRSSS	449	
PTH <sub>2</sub> R		SQSQVAASTRMVLISGKAAKIAS	483	
C mGlu <sub>5</sub> R		ARSSSSGSLMEQISSVVTRFTA	1070	
GABA <sub>B1</sub> R		IAEKEERVSEL-RHQLQSRQQLRSRR	926	
GABA <sub>B2</sub> R		PSIGGVDASCVSPCVSPTASPRHRHV	930	
$GABA_{BL}R^*$ (1)		RRAAQRARSHF-PGSAPSSVGHRANR	626	
$GABA_{BL}R*$ (2)	010	SHFPGSAP <b>S</b> S <b>V</b> GHRANR <b>T</b> VPGA <b>H</b> SRL	635	