NOVEL MECHANISM FOR REGULATION OF EGF RECEPTOR
ENDOCYTOSIS REVEALED BY PROTEIN KINASE A INHIBITION

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Abbreviations: EGF, epidermal growth factor; EGFR, epidermal growth factor receptor (s); TxR, Texas-red; PKC, protein kinase C; PKA, protein kinase A; MβCD, methyl-β-cyclodextrin; Tf, transferrin; diFeTf, diferric transferrin; TfR, transferrin receptor; DAMGO, [D-Ala², MePhe⁴, Gly-ol³] enkephalin.

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ABSTRACT

Current models put forward that the epidermal growth factor receptor (EGFR) is efficiently internalized via clathrin-coated pits only in response to ligand-induced activation of its intrinsic tyrosine kinase and is subsequently directed into a lysosomal-proteasomal degradation pathway by mechanisms that include receptor tyrosine-phosphorylation and ubiquitylation. Here, we report a novel mechanism of EGFR internalization that does not require ligand binding, receptor kinase activity or ubiquitylation and does not direct the receptor into a degradative pathway. Inhibition of basal PKA activity by H89 and the cell-permeable substrate peptide Myr-PKI induced internalization of 40-60% unoccupied, inactive EGFR, and its accumulation into early endosomes without affecting endocytosis of transferrin and µ-opioid receptors. This effect was abrogated by interfering with clathrin function. Thus, the predominant distribution of inactive EGFR at the plasma membrane is not simply by default but involves a PKA-dependent restrictive condition resulting in receptor avoidance of endocytosis until it is stimulated by ligand. Furthermore, PKA inhibition may contribute to ligand-induced EGFR endocytosis since EGF inhibited 26% of its basal activity. On the other hand, H89 did not alter ligand-induced internalization of EGFR but doubled its half-time of down-regulation by retarding its segregation into degradative compartments, seemingly due to a delay in the receptor tyrosine-phosphorylation and ubiquitylation. Our results reveal that PKA basal activity controls EGFR function at two levels: 1) residence time of inactive EGFR at the cell surface by a process of “endocytic evasion”, modulating the accessibility of receptors to stimuli; 2) sorting events leading to the down-regulation pathway of ligand-activated EGFR, determining the length of its intracellular signaling. They add a new dimension to the fine tuning of EGFR function in response to cellular demands and cross-talk with other signaling receptors.
INTRODUCTION

The mechanisms that regulate the endocytic behavior of epidermal growth factor receptor (EGFR) have been a long-standing subject of intense research as a model system of regulated vesicular protein traffic associated to signaling, cellular demands and cancer (Carpenter, 2000; Ceresa and Schmid, 2000; Di Fiore and Gill, 1999; Schlessinger, 2000; Trowbridge et al., 1993; Wiley and Burke, 2001). Endocytosis provides a pathway for gradual attenuation or desensitization of receptor signaling and also allows a single receptor to transmit different signals from different locations in the cell before degradation, thus enhancing the range of modulation and response variability (Ceresa and Schmid, 2000; Di Fiore and Gill, 1999; Wiley and Burke, 2001). Defects in internalization and/or degradation pathways, as well as distinct endocytic routing displayed by different EGFR family members, have been associated with cell transformation and oncogenesis (Ceresa and Schmid, 2000; Di Fiore and Gill, 1999). The EGFR is also a downstream element of non-EGF like proliferation signals (Carpenter, 1999). All this prompts to search for mechanisms and factors able to control EGFR internalization, endosomal sorting and degradation, either in concert with ligand-induced events or independently of ligand binding. Serine-threonine kinases are interesting elements to explore since they play roles in both signaling and vesicular protein transport.

Ligand-mediated activation of the EGFR intrinsic tyrosine-kinase results in phosphorylation events involved in the signaling process, receptor endocytosis and final fate after internalization. Ligand binding induces structural changes in the EGFR that presumably expose cryptic codes in its cytosolic domain, favoring interaction with the clathrin-mediated endocytic apparatus (Boll et al., 1995; Nesterov et al., 1995a; Nesterov et al., 1995b), mainly through its adaptor AP-2 (Cadena et al., 1994; Nesterov et al., 1995a; Sorkin et al., 1996), although AP-2-independent processes have been also invoked (Nesterov et al., 1995b; Sorkin and Carpenter, 1993). In addition, EGFR activation stimulates the endocytic apparatus through tyrosine phosphorylation of a variety of downstream substrates needed for its efficient recruitment into coated pits (Lamaze and Schmid, 1995). The tyrosine kinase src is activated during ligand-induced EGFR endocytosis, phosphorylating and
redistributing clathrin to the plasma membrane (Wilde et al., 1999). In this scenario, only ligand-activated receptors would be efficiently recognized by the clathrin-endocytic machinery. Permanency of inactive EGFR at the cell surface is implicitly believed to be by default, grounded only on properties of receptor resting structure. On the other hand, sorting of EGF/EGFR from endosomes to a lysosomal degradation pathway depends on specific targeting information residing in its intracellular domain (Kil et al., 1999; Kurten et al., 1996) and has been recently shown to involve ubiquitylation of the receptor (Levkowitz et al., 1998) in a process influenced by protein kinase C (PKC) and mediated by the proto-oncogene product c-Cbl, that is tyrosine-phosphorylated by the EGFR (Bao et al., 2000).

Several studies have shown that the exocytic and endocytic pathways are tightly regulated by different combinations of serine-threonine kinases, including PKC and protein kinase A (PKA), presumably by controlling the generation of transport carriers emerging from the ER, Golgi complex or plasma membrane (Aridor and Balch, 2000; Goretzki and Mueller, 1997; Jamora et al., 1999; Lee and Linstedt, 2000; Muniz et al., 1996; Muniz et al., 1997; Pimplikar and Simons, 1994). PKC and PKA have been also implicated in transmodulation of ligand and/or kinase activity of the EGFR (Barbier et al., 1999; Carpenter, 1999; Ciardiello and Tortora, 1998; Schlessinger, 2000). However, only PKC has been additionally reported to modulate EGFR function by modifying its endocytic sorting (Bao et al., 2000). Instead, for several other proteins, including the low-density lipoprotein receptor-related protein (Li et al., 2001), urokinase-type plasminogen activator (Goretzki and Mueller, 1997) and Na+/H+ exchanger NHE3 (Hu et al., 2001), it has been recently described that PKA activity is required for their stimulated endocytosis. Inhibition of PKA activity or mutation of PKA phosphorylation sites inhibit the endocytosis of these proteins.

H89 is a potent, highly specific and reversible inhibitor of PKA (Chijiwa et al., 1990), which, in combination with a panel of other kinase inhibitors, provides a convenient way to analyze the function of PKA in diverse vesicular transporting pathways (Aridor and Balch, 2000; Goretzki and Mueller, 1997; Jamora et al., 1999; Lee and Linstedt, 2000; Muniz et al., 1996; Muniz et al., 1997). Here we report that inhibition of PKA activity by H89 or the PKA inhibitory peptide Myr-PKI
induced endocytosis and selective redistribution of empty inactive EGFR into early endocytic compartments. Thus, basal PKA activity participates in a previously unsuspected mechanism that keeps inactive EGFRs predominantly at the cell surface by abrogating their interaction with the endocytic apparatus. Additionally, we found that H89 delayed the endocytic sorting of ligand-activated EGFR to a degradative lysosomal-proteosomnal pathway, very likely as a consequence of retarding the receptor autophosphorylation and ubiquitylation, suggesting that the time course of intracellular signaling before degradation is also under the influence of basal PKA activity.
MATERIALS AND METHODS

Reagents and antibodies

Human recombinant EGF and cDNA encoding the human EGFR were provided by Drs. Pablo Valenzuela and Carlos George-Nascimento, Chiron Co., Emeryville, CA. cDNA encoding the N-terminal FLAG-tagged murine µ-opioid receptor was provided by Drs. Paulette Zaki and Christopher Evans, University of California, Los Angeles. Dulbecco’s modified Eagle’s medium containing high glucose, Protein A-Sepharose, butyric acid and all other reagents were purchased from Sigma. Fetal bovine serum was purchased from HyClone. H89, forskolin, H8, H7, Myr-PKI (14-22), KT5720, staurosporine, chelerythrine, Calphostin C, 5,6-dichloro-1-β-D-ribofuranosylbenzimidazole (DRB) and genistein were from Calbiochem. CKI-7 was obtained from Seikagaku Corp and wortmannin and DAMGO was from Sigma.

For immunoblot detection, the ECL system from Life Sciences, Amersham Corp. was utilized. Cell surface labeling reagents, EZLink™ sulfo-NHS-biotin and EZLink™ sulfo-NHS-SS-biotin, were from Pierce. Cell culture reagents were purchased from GIBCO and Sigma. Tissue culture plastics were from Nunc. Polyclonal antibody EGFR984 has been previously characterized (Faúndez et al., 1992) and was obtained against the peptide described (Kriss et al., 1985), corresponding to residues 984-996 (DDVVDADEYLIPQ) of the EGFR cytosolic tail. Hybridomas producing monoclonal anti-EGFR antibodies HB8506 reacting with the extracellular domain of the EGFR and antiphosphotyrosine HB8190 were purchased from ATCC. HB8190 can detect phosphorylated tyrosines both in immunoblot and immunoprecipitation (Faúndez et al., 1992). Anti-ubiquitin antibody was from Santa Cruz Biotechnology, Inc. M2 antibody against FLAG was from Sigma.

Cell culture, transfection, and drug treatments

N2a cells, cultured in Dulbecco’s modified Eagle’s medium, 4.5 g/liter glucose, supplemented with 7.5% fetal bovine serum and antibiotics (50 mU/ml penicillin, 100 µg/ml streptomycin, and 50 µg/ml fungizone) were transfected with a vector pBK-CMV into which the cDNA of the human EGFR was sub-cloned, or with pcDNA3 vector into which the cDNA of
FLAG-µ-opioid receptor was subcloned. Transfection was done with lipofectAMINE (Gibco BRL) according to the manufacturer’s instructions. Permanently transfected cells were selected in 0.8 mg/ml G418 and maintained in the mentioned culture medium supplemented with 0.2 mg/ml G418 (Gibco-BRL). Expression was induced by 10 mM sodium butyrate for 12-24 h, following our previously described protocols (Marzolo et al., 1997). K721A cells expressing a point mutant kinase-negative EGFR and Her14 expressing wt EGFR (Felder et al., 1990), were a kind gift from Dr. J. Schlessinger (New York University Medical Center, New York). In all the experiments the cells were cultured to about 80% confluency and then were serum-starved by incubation during 24 h in media supplemented with only 0.3 % FBS (Hyclone GiBCO). In the case of H89, KT5720, CKI-7, staurosporin and DRB, stock solutions were prepared in DMSO and, therefore, control conditions included concentrations of 0.2-0.4% DMSO.

**Indirect immunofluorescence and colocalization with transferrin-Texas Red**

Cells were grown on glass coverslips, subjected to the different experimental conditions and then washed with PBS and fixed for 30 min at room temperature with 4% paraformaldehyde in PBS supplemented with 0.1 mM CaCl₂ and 1 mM MgCl₂ (PBS-CM). After washing three times with PBS plus 0.2% gelatin (300 Bloom, Sigma Chemical Co.) for 5 min each, the cells were permeabilized with 0.2 % Triton-X100 for 10 min at room temperature and incubated for 1 h at 25°C with anti-EGFR monoclonal antibody HB8506 (1/100 in PBS). Incubation with the secondary antibody, anti mouse IgG FITC (1/100 in PBS, 1 hr at 25°C), was made after six washes in PBS-gelatin. Trafficking of the internalized EGFR through the early endocytic compartments was analyzed together with transferrin-TxR. The cells were washed three times with cold D-MEM and then incubated with 50 µg/ml of transferrin-TxR (Molecular Probes) for different time periods at 37°C, washed and processed for indirect immunofluorescence of the EGFR. Fluorescence images were collected on a Zeiss Axioplan microscope and photographed using a 63x immersion objective and the Axiocam camera. Images were transferred to a computer workstation running Axiovision imaging software.
Receptor Binding and Internalization Assays

Human $^{125}$I-EGF was prepared by the chloramine T method as described (Faúndez et al., 1992). The specific activities of labeled ligand were typically 50,000-70,000 cpm/ng. Binding assays were done in Hanks solution with 20 mM HEPES and 0.1% BSA during 2 h at 4°C. Free ligand was removed by washing 4 times with ice-cold PBS 0.1% BSA. Cell bound ligand was released in 1 M NaOH for 1 h at room temperature. Nonspecific binding was assessed in the presence of 1 µg/ml of unlabeled human EGF and subtracted from the total. Samples were quantified using a gamma counter (COBRA™II, Packard, Canberra Co.). Scatchard analysis was performed with the LIGAND program (Munson and Rodbard, 1980). To monitor $^{125}$I-EGF internalization, cells in 12-well dishes were incubated with 50 ng/ml of $^{125}$I-EGF in binding medium for 2 h at 4°C. After washing the unbound radioligand, the cells were transferred to 37°C for different time periods. Assays were ended by rapidly cooling the monolayers in ice-water followed by 4 washes with ice-cold PBS 0.1% BSA to remove unbound ligand and then surface and internalization assessment. The established acid-wash method (Haigler et al., 1980) was made incubating the cells for 6 min with 0.2 M acetic acid (pH 2.5) containing 0.5 M NaCl at 4°C. The acid wash was combined with another short rinse with the same acidic solution to determine the amount of surface-bound (SUR) $^{125}$I-EGF. The cells were then lysed in 1 N NaOH to quantitate internalized (IN) radioactivity. $^{125}$I-EGF degradation was assessed by TCA precipitation of incubation media. All data represents specific $^{125}$I-EGF binding assessed by adding 100-fold molar excess of cold EGF with each point in duplicate.

DiFeTf was prepared from apoTf as described elsewhere (Bali and Harris, 1990), and iodinated with the chloramine T method in similar conditions as for EGF, achieving an specific activity of 13500 cpm/ng.

The IN/SUR method (Wiley and Cunningham, 1982) was used to estimated the endocytic rate constant $k_e$, as the slope calculated by linear regression of the biphasic curve resulting from the plot of the ratio of internalized (IN) to surface (SUR) ligand (IN/SUR) versus time.
**Cell surface biotinylation and immunodetection of EGFR**

The EGFR present at the cell surface was assessed by biotinylation assays and by surface immunoprecipitation with the antibody HB8506 followed by Protein-A-Sepharose (Bravo-Zehnder et al., 2000). Briefly, for biotinylation, cells were washed three times in PBS-CM at 4°C, and then sulfo-NHS-Biotin (Pierce, Rockford, IL) was added to a final concentration of 0.5 mg/ml for 30 min. The monolayers were washed six times in ice-cold PBS and incubated with NH₄Cl 50 mM/PBS-CM for 10 min to block free biotin. Biotin assay for endocytosis experiments with reducible sulfo-NHS-SS-biotin was made as described (Le Bivic et al., 1989), reducing surface sulfo-NHS-SS-Biotin with 50 mM glutathion for 30 min, in 90 mM NaCl, 1 mM MgCl₂, 0.1 mM CaCl₂, 60 mM NaOH, 10% FBS. The cells were then lysed and immunoprecipitated with anti-EGFR antibody HB8506 previously linked to Protein A Sepharose beads for 2 hrs. The beads were washed six times in ice-cold buffer for 5 min each and incubated in sample buffer for 5 minutes at 65°C. The eluates were run in a 7.5% acrylamide gel. Protein detection was made by blot analysis with streptavidin-HRP conjugate and the ECL detection kit (Amersham) in nitrocellulose filters, after electrotransferring the biotinylated surface proteins.

**Immunoprecipitation**

Immunoprecipitations were performed following established procedures with minimal modifications (Faúndez et al., 1992; Marzolo et al., 1997). Cells were lysed with 1% Triton X-100 in 50 mM Hepes, 150 mM NaCl, 1 mM EGTA, 2 mM MgCl₂, glycerol 10% supplemented with an antiprotease mixture (4 mM PMSF, 0.28 TIU/ml aprotinin; 4 µg/ml of pepstatin, leupeptin and antipain). Total cell protein was determined by Bio-Rad protein assay (Bio-Rad Richmond, CA) using bovine serum albumin as standard. Immunoprecipitations of equivalent total protein amounts were performed at 4°C for 2 h using the primary antibody previously bound to 30 µl of Protein A-Sepharose CL-4B beads per sample. The beads were washed six times with ice-cold buffer (20 mM Hepes, 150 mM NaCl, 10% glycerol and 0.1% TritonX-100). The beads in each tube were heated to 65°C for 5 min in 20 µl of sample buffer (62.5 mM Tris, pH 6.8, 2% SDS, 100 mM dithiothreitol, 10% glycerol, 0.005% bromophenol blue), and then clarified by centrifugation. Proteins were
separated by SDS-PAGE (Laemmli, 1970) on 7.5% acrylamide gels and transferred to nitrocellulose membranes (Towbin et al., 1979). Membranes were blotted for proteins as indicated and visualized using horseradish peroxidase-conjugated secondary antibody and ECL detection system. The tyrosine phosphorylation experiments were made in the lysis buffer supplemented with 1.5 mM MnCl₂, 1 mM MgCl₂ and 1 mM sodium orthovanadate in which case intact cells were incubated at 37°C either in the absence or presence of 100 ng/ml EGF. For immunoprecipitation of EGFR present on the cell surface, the cells were first metabolically labeled and then incubated with HB8506 antibody at 4°C for 2 h. Cells were starved in methionine/cysteine free D-MEM medium (Gibco) with 0.3% FBS for 30 min, and then incubated with 0.1 mCi (EasyTaq, Express Labeling Mix) of ³⁵S-labeled methionine and cysteine (ICN Radiochemicals, Cleveland, OH) in 1 ml of D-MEM for 12 h at 37°C. Fluorograms were developed on pre-flashed Kodak AR X-Omat films (Eastman). Fluorograms or immunoblots were digitalized in a VISTA-T630 UMax scanner driven by Adobe Photoshop 3.1 and quantitative analysis was done with the NIH Image 1.55 (fpu) software.

**Inhibition of clathrin-mediated endocytosis by cytosol acidification, potassium or cholesterol depletion**

Cells were incubated in DMEM 0.3% FBS overnight before the experiments. Cytosol acidification (Sandvig et al., 1987) was carried out by washing the cells twice in DMEM pH 7.4 and then incubating them at 37°C for 10 min in DMEM plus 10 mM acetic acid, pH 5.0. For the intracellular potassium depletion (Larkin et al., 1983), the cells were rinsed twice with buffer A (20 mM Hepes, 140 mM NaCl, 1 mM CaCl₂, 1 mM MgCl₂, pH 7.4), incubated in hypotonic buffer A (buffer A diluted 1:1 with water) for 5 min at 37°C, and then rinsed and incubated again in buffer A for additional 30 min at 37°C. Cholesterol depletion was done with methyl-β-cyclodextrin (MβCD) (Rodal et al., 1999; Subtil et al., 1999) in Her14 cells. The cells were washed twice in PBS, incubated in DMEM 0.3% FBS with 10 mM MβCD for 30 min at 37°C and then incubated with or without H89 in the same medium for additional 10 min. After these procedures, the cells were rinsed twice with PBS before adding either EGF (100 ng/ml) for 10 min or 20 μM H89 for 1 h at 37°C.
RESULTS

**H89 provokes a delay in the EGF/EGFR complex degradation**

To analyze the effects of H89 on EGF/EGFR endocytosis and degradation we first used transfected neuroblastoma N2a cells as a model system that provided cells expressing different levels of the receptor. We selected colonies of transfected cells expressing about 200,000 receptors/cell, hereafter called N2a-15 cells. These transfected cells displayed high (Kd: 0.08 nM) and low affinity (Kd: 0.56 nM) forms of the EGFR (our unpublished results), as described for most cellular systems. Instead, in non-transfected N2a cells, we could not detect endogenous EGF-R by $^{125}$I-EGF binding, immunoblot, and immunofluorescence.

Cells treated for 1 h with 20 $\mu$M H89 and exposed to $^{125}$I-EGF at 4ºC for 2 h were incubated at 37ºC for various time periods to allow internalization of the ligand-receptor complex. As previously shown in human melanoma cells (Goretzki and Mueller, 1997), H89 did not affect $^{125}$I-EGF endocytosis (Fig. 1A). In fact, the internalization rate showed no significant differences between control ($k_e = 0.149$ min$^{-1}$) and H89 treated cells ($k_e = 0.143$ min$^{-1}$) (Fig. 2C). However, H89 diminished the degradation of the internalized $^{125}$I-EGF (Fig. 1B and C) and provoked a delay in the degradation of ligand-activated EGFR, increasing its half-time of down regulation (Fig. 2A). N2a-15 cells incubated with saturating concentrations of EGF (100 ng/ml) showed a progressive disappearance of the EGFR with a half-time of about 34 min, that increased to 77 min in cells pre-treated for 1 h with H89 (Fig. 2B). The level of receptor degradation after the delay was finally similar (80%) to that achieved by ligand alone.

**H89 inhibits ligand-induced tyrosine-phosphorylation and ubiquitylation of the EGFR**

Ligand binding induces EGFR tyrosine phosphorylation and subsequently ubiquitylation, which has been functionally related to sorting into a degradation pathway (Bao et al., 2000; Levkowitz et al., 1998). Under the effect of H89, the levels of EGFR tyrosine-phosphorylation and ubiquitylation remained undetectable within the first 5 min of EGF stimulation (Fig. 3A), appearing in the next time period of 10 min (lane 7), when almost 90% of the ligand activated receptors have been already internalized, as shown by cell surface biotinylation assays (Fig. 3B, lane 4). The slower
kinetics of these ligand-induced modifications could account for the delayed down-regulation of the receptor seen under H89 treatment.

**H89 induces ligand-independent internalization of EGFR**

During the acid wash experiments we noticed that pre-treating cells with 20 µM H89 decreased the levels of \(^{125}\text{I}\)-EGF binding. Scatchard analysis showed that H89 induces complete loss of the high affinity forms of the EGF-R and a dramatic decrease in the low affinity binding sites (Fig. 4A). Overall, more than 50% of the binding sites disappeared in 1 h of H89 treatment. To study whether this was due to endocytic removal of empty inactive EGFR from the cell surface, we performed cell surface biotinylation assays (Fig. 4B; lanes 1 and 2) and cell-surface immuno-detection with an antibody against the external domain of the EGF-R (Fig. 4B; lanes 3 and 4). Both experiments showed decreased levels of EGFR at the cell surface after 1 h of 20 µM H89 treatment, the total cellular content of the receptor remaining unchanged (Fig. 4B; lanes 5 and 6). In contrast with its fate after ligand-induced endocytosis, the H89-internalized receptor was not degraded for at least 4 h (Fig. 4C). Immunofluorescence made in N2a-15 and in Her14 cells (Honegger et al., 1987), that are more flattened, showed the receptor accumulated in a juxtanuclear compartment (Fig. 4D), probably corresponding to recycling endosomes that in many cells appear as a bright perinuclear spot (Mukherjee et al., 1997; Trowbridge et al., 1993).

Experiments of cell surface biotinylation with the cleavable reagent sulfo-NHS-SS-biotin (Le Bivic et al., 1989) directly demonstrated H89-induced endocytosis of unoccupied EGFR. Glutathion added to the medium eliminates the biotin only from cell surface proteins since internalized proteins become inaccessible to it. N2a-15 cells were first incubated with sulfo-NHS-SS-biotin at 4ºC and then warmed to 37ºC in the presence or absence of 20 µM H89 for 5, 10 and 15 min. Internalization was readily detectable within the first 5 min of H89 treatment (Fig. 5A; lane 2) while it was not observed under control conditions (Fig. 5A; lanes 1, 3 and 5). The EGFR lost biotin label during the next time periods of 10 and 15 min (lane 6), indicating that a proportion of the internalized receptors recycled back to the cell surface.
The endocytic effect of H89 was also seen by immunofluorescence (Fig. 5B). After 1 h of incubating N2a-15 or NIH-3T3 Her14 cells with H89, the EGFR ends up accumulated in a juxtanuclear endocytic compartment reminiscent of recycling endosomes.

**EGFR endocytosis induced by H89 does not involve the intrinsic protein tyrosine-kinase activity, tyrosine-phosphorylation and ubiquitylation of the receptor**

In contrast with the effects induced by EGF, we could not detect any significant change in the phosphotyrosine content of the EGFR when cells were treated with 20 µM H89 (Fig. 6A). Furthermore, H89 was able to induce internalization of the mutant receptor K721A which lacks tyrosine kinase activity (Honegger et al., 1987), as seen by immunofluorescence (Fig. 6B), ligand-binding and cell surface biotinylation experiments (our unpublished results). In addition, cells incubated with 1 mCi/ml $^{32}$P for 4 h showed no changes in the phosphorylation status of the EGFR upon H89 treatment (our unpublished results), making unlikely modifications of serine-threonine phosphorylation of the receptor.

It has been recently described that EGFR ubiquitylation can occur at the cell surface and therefore might provide additional endocytic information (Stang et al., 2000). However, in contrast with the effects of ligand stimulation (Fig. 6C, lanes 8-12), we could not find evidence of EGFR ubiquitylation during H89-induced endocytosis (Fig. 6C, lanes 2-6), congruently with the fact that EGFR ubiquitylation requires its tyrosine-phosphorylation (Levkowitz et al., 1998).

**H89-induced internalization of EGFR occurs via clathrin-mediated endocytosis**

The endocytic pathway induced by H89 seems to be saturable. The average levels of EGFR endocytosis induced by 20 µM H89 for 1 h and measured as remanent ligand binding were 57% ± 5% (n=10) for N2a-15 cells, that express 200,000 receptors/cell and 44% ± 4% (n=6) for Her14 cells expressing 270,000 receptors/cell. Instead, it was only 4% ± 1% (n=6) in A431 cells expressing around 4,000,000 receptors/cell and 3% ± 2% (n=4) in another colony (N2a-3) of transfected N2a cells which express about 1,000,000 receptors/cell. Similar saturability has been demonstrated for ligand-induced internalization of EGFR via clathrin-coated pits (Wiley, 1988; Wiley et al., 1991).
Cytosol acidification (Sandvig et al., 1987) and potassium depletion (Larkin et al., 1983), that block endocytosis by clathrin-coated pits, inhibited H89-induced as well as EGF-induced disappearance of EGFR from the cell surface (Fig. 7A). Furthermore, treatment with 10 mM MβCD, that causes acute cholesterol depletion from the plasma membrane and has been recently shown to decrease clathrin-coated pit function (Rodal et al., 1999; Subtil et al., 1999), was also able to inhibit receptor endocytosis induced by H89 (Fig. 7B). Clathrin-mediated EGFR endocytosis induced by ligand is also inhibited by the tyrosine kinase inhibitor genistein that would interfere with phosphorylation events mediated not only by the intrinsic receptor kinase but also by src (Wilde et al., 1999). Genistein also abolished EGFR endocytosis induced by H89 (Fig. 7B). All these observations indicate that H89 promotes an interaction of EGFR with the clathrin-mediated endocytic machinery.

**H89 effects on EGFR endocytosis correlated with inhibition of PKA activity.**

H89 has been currently used as a potent, highly selective and reversible inhibitor of PKA (Chijiwa et al., 1990; Muniz et al., 1996; Muniz et al., 1997), since it is approximately a 1000-fold more efficient inhibitor for PKA than for PKC or other known kinases (Chijiwa et al., 1990). At 30 μM, H89 maintains selectivity for PKA inhibition in intact cells without affecting other kinases measured in extracts of treated cells (Chijiwa et al., 1990) while at 90 μM, H89 inhibits Golgi vesiculation induced by the drug illimaquinone involving PKD instead of PKA (Jamora et al., 1999).

In our conditions, assessment of PKA activity in extracts of intact cells treated with H89 for 1 h showed an IC$_{50}$ of about 12 μM (Fig. 8A). This correlated with the IC$_{50}$ of 13 μM estimated for the effect of H89 on $^{125}$I-EGF binding (Fig. 8C). Ligand-binding activity decreased within the range of 5-20 μM H89, whereas at higher H89 concentrations this effect was gradually lost until it became completely abolished at 300 μM. Thus, other serine-threonine kinases inhibited at these high H89 concentrations (Chijiwa et al., 1990; Cotlin et al., 1999) are likely required for the endocytosis of EGFR acting downstream of PKA. The time course of PKA inhibition was also congruent with an involvement of PKA in the endocytic effect of H89. Substantial inhibition of PKA activity was
achieved as fast as 5 min of H89 coincident with the time course of H89-induced endocytosis of EGFR (Fig 8B and D) (see also Fig. 5). Strikingly, EGF by itself causes a 26% decrease in the levels of PKA activity (Fig. 8B), suggesting that this effect might play a role in EGFR ligand-induced endocytosis.

Myr-PKI, a specific peptide substrate for PKA considered to be one of its most highly selective inhibitors (Harris et al., 1997), was also able to induce EGFR endocytosis, as it was most clearly seen in HeLa cells (Fig. 9). Either 20 µM H89 or 100 µM Myr-PKI led to redistribution of the receptor from the cell surface to intracellular compartments, progressively concentrated in perinuclear regions. Within the first 15 min of H89 treatment, linear arrays of vesicles resembling tubulo-vesicular structures, were frequently seen converging toward these perinuclear regions.

Instead, inhibitors of PKC (H-7, calphostin-C and chelerythrine), casein kinase I and II (CKI-7 and DRB) and tyrosine kinases (genistein) were all ineffective as inductors of EGFR endocytosis (our unpublished results). Wortmannin, an inhibitor of PI-3 kinase that participates in the endocytosis of several proteins (Chen and Wang, 2001), has no effect either (our unpublished results).

**Changes in the intracellular pathway of the EGFR and redistribution of transferrin endocytic compartments upon H89 treatment**

The endosomal compartment is a system of interconnected tubulo-vesicular elements currently and broadly divided by functional and structural criteria into early (sorting and recycling) and late (pre-lysosomal) endosomes (Mukherjee et al., 1997; Trowbridge et al., 1993). The endocytic pathway followed by EGFR under the effect of H89 was analyzed by double fluorescence using Tf-TxR as marker for early endosomal compartments (Mukherjee et al., 1997; Trowbridge et al., 1993). The EGFR internalized by EGF stimulation extensively colocalized with Tf-TxR during the first 10-15 minutes and then was segregated from Tf-TxR into distinct vesicular structures, as expected for proteins following recycling and degradation pathways (Fig. 10, 30 min). After one hour, the staining of the EGFR diminished in most cells. Instead, the EGFR internalized during H89 treatment in the absence of ligand showed a maintained colocalization with Tf-TxR and both were
gradually concentrated in a juxtanuclear compartment very likely corresponding to recycling endosomes (Mukherjee et al., 1997). Such distribution was not achieved by the EGFR in cells treated with EGF alone. Finally, when Her14 cells were incubated with EGF and H89, the pattern of EGFR endocytosis frequently showed tubulo-vesicular endocytic compartments and the colocalization of EGFR and Tf-TxR lasted longer than in cells treated with EGF alone, since it was still observed in many cells after 60 min of H89 treatment (Fig. 10). This means that EGF/EGFR complexes were more slowly segregated from early endosomes to lysosomes under the effects of H89, as suggested also by the delayed receptor down-regulation and ligand degradation seen before.

**H89 does not induce endocytosis of transferrin and µ-opioid receptors, but provoked changes in the early endocytic compartments**

Indirect immunofluorescence of the human TfR made in HeLa cells showed that H89 causes changes in the distribution of early endocytic compartments. During the first 5-10 min of H89 incubation, the TfR was seen in linear arrays of vesicles, possibly corresponding to tubulo-vesicular structures, converging toward the perinuclear region (Fig. 11A), a pattern similar to that showed by the EGFR under the effects of H89 alone (see Figure 9). After 60 min of H89 treatment, these linear arrays were rarely observed while instead perinuclear vesicles became predominant. In spite of these changes, the internalization rate of Tf was not affected by 20 µM H89, neither in N2a (control, \( k_e = 0.013 \text{ min}^{-1} \); H89, \( k_e = 0.017 \text{ min}^{-1} \), n=6) nor in HeLa cells (control, \( k_e = 0.107 \text{ min}^{-1} \); H89, \( k_e = 0.109 \text{ min}^{-1} \), n=6) (Fig. 11B). This is consistent with previous reports in HeLa cells and A431 cells in which the internalization rate of Tf was not affected by low concentrations of H89, being only decreased at higher concentrations (300 µM) due to casein kinase II inhibition (Cotlin et al., 1999). Treatment with either H89 (20 µM) or Myr-PKI peptide (100 µM) for 60 min did not modify the cell surface \(^{125}\text{I}-\text{diFeTf} \) binding activity (our unpublished results), indicating that recycling of continuously internalized TfR was also not altered.

In addition, to test other kind of signaling receptors whose clathrin-mediated endocytosis is induced by ligand we assessed the effects of H89 on µ-opioid receptors, pertaining to the class of G protein-coupled receptors (Keith et al., 1996). H89 did not induce endocytosis of a flag-tagged µ-
opioid receptor expressed in permanently transfected N2a cells, contrasting with the clear endocytic effect exerted by its ligand DAMGO (Fig. 11C). Therefore, the mechanism involving basal PKA activity by which the EGFR avoids the endocytic apparatus seems to be rather selective.
DISCUSSION

Our results revealed new regulatory systems controlling the permanency of inactive EGFR at the cell surface and the efficiency of intracellular segregation of ligand-activated EGFR to a degradation pathway, both at the expense of basal PKA activity.

The most striking finding was that inhibition of PKA basal activity, by either low concentrations of H89 (IC$_{50}$=12 µM) or the PKA substrate peptide Myr-PKI, in the absence of ligand, induced rapid internalization of inactive EGFR through the clathrin-mediated pathway. Such an effect was not generalized. We could not detect H89 induction of endocytosis either in transferrin receptors which are constitutively internalized (Mukherjee et al., 1997; Trowbridge et al., 1993) or in µ-opioid receptors that undergo endocytosis upon ligand-binding (Keith et al., 1996). The magnitude of the endocytosis depended on the PKA inhibitor used, the cell line, and the levels of receptor expressed. Close to 60% of EGFR was removed from the cell surface in N2a-15 cells and accumulated in early endosomes without degradation for at least 4 hours. All this reveals the existence of mechanisms controlling the relative distribution of EGFR between the cell surface and endocytic compartments, indicating also that the predominant distribution of inactive EGFR at the plasma membrane is not simply by default but through a regulated process of “endocytic evasion”, amenable to modulation according to cellular demands. Basal PKA activity exerts some kind of restrictive condition resulting in EGFR avoidance of endocytosis until it is stimulated by ligand, either maintaining the receptor endocytic codes cryptic or an EGFR internalization-promoting element inactive.

Accepted models of EGFR endocytosis state that activation of its intrinsic tyrosine-kinase domain is required for efficient receptor recruitment into clathrin-coated pits by exposing cryptic endocytic codes that interact with AP-2 adaptors (Cadena et al., 1994; Nesterov et al., 1995a; Sorkin et al., 1996). EGFR could also become ubiquitylated at the cell surface and this modification might eventually serve as endocytic information (Stang et al., 2000). However, we could detect neither an increased tyrosine phosphorylation nor ubiquitylation of the EGFR receptor during H89 treatment. Furthermore, H89 provoked internalization of the mutant receptor K721A which lacks tyrosine
kinase activity (Felder et al., 1990) and is not a substrate of c-Cbl ubiquitylation (Levkowitz et al., 1998). In spite of lacking these modifications several observations involved a clathrin-mediated pathway in the EGFR endocytic effects of H89. Analysis of cells expressing high levels of EGFR showed that the endocytic process activated by H89 was saturable, as previously described for receptors internalized via clathrin-coated pits (Wiley, 1988; Wiley et al., 1991). Three procedures that inhibit clathrin-coated pit function, cytosol acidification (Sandvig et al., 1987), potassium depletion (Larkin et al., 1983), and acute cholesterol depletion (Rodal et al., 1999), effectively blocked internalization of EGFR induced by either EGF or H89. Finally, internalization through caveolae, where EGFR has been localized in fibroblasts (Mineo et al., 1996; Smart et al., 1995), could be discarded because N2a cells do not express caveolin-1 and therefore do not form caveolae (Gorodinsky and Harris, 1995; Shyng et al., 1994). Whether or not H89 induces interaction of EGFR with AP-2 is an interesting possibility to explore in future experiments. In certain conditions, this interaction can occur independently of receptor tyrosine phosphorylation levels and ligand-binding (Boll et al., 1995; Sorkin and Carpenter, 1993), or might not even be necessary for EGFR endocytosis, as suggested by studies made with mutant receptors lacking AP-2 binding sites (Nesterov et al., 1995b) and with an AP-2 dominant negative µ2 subunit responsible for recognizing endocytic codes (Nesterov et al., 1999).

The fate of internalized EGFR under the effect of H89 was different from that seen in cells stimulated with EGF. It was not substantially delivered to a proteasomal/lysosomal pathway as indicated by a lack of degradation during 4 h of H89 treatment and an extensive colocalization with Tf-TxR seen first in peripheral endocytic compartments and then in a punctate juxtanuclear region with the characteristic pattern of recycling endosomes. This supports the notion that receptor tyrosine-phosphorylation and ubiquitylation, which are not induced by H89, are indeed required for its sorting to a degradation pathway (Bao et al., 2000; Levkowitz et al., 1998). The effects of H89 are also different from those described for PKC activation, which results in transient internalization of unoccupied EGFR from the cell surface but the receptors recycled to the cell surface in just 1 h (Beguinot et al., 1985; Lin et al., 1986). PKA inhibition also changed the intracellular distribution of
TfR but seemingly without functional consequences. Whereas after one hour of PKA inhibition almost 50% of the EGF binding activity disappeared from the cell surface due to internalization no changes were detected in Tf binding, indicating that its recycling remained unaffected, as occurred also with its internalization rate.

EGFR endocytosis induced by PKA inhibition is the first example of protein segregation into a vesicular pathway induced by kinase inhibition and highlights the importance of basal kinase activity in selective protein trafficking. All previous reports showed that kinase activation or inhibition resulted in the corresponding induction or abrogation of protein vesicular transport, both in the exocytic and endocytic routes (Aridor and Balch, 2000; Goretzki and Mueller, 1997; Jamora et al., 1999; Jilling and Kirk, 1996; Lee and Linstedt, 2000; Muniz et al., 1996; Muniz et al., 1997; Pimplikar and Simons, 1994). For instance, H89 inhibits trans-Golgi network-to-cell surface transport by decreasing PKA dependent formation of transport vesicles (Muniz et al., 1996; Muniz et al., 1997). Inhibition of PKA and abrogation of PKA phosphorylation sites also decreased the stimulated endocytosis of several proteins (Goretzki and Mueller, 1997; Hu et al., 2001; Li et al., 2001).

Ligand-induced EGFR endocytosis involves several downstream substrates of the receptor and activation of the protein tyrosine kinase src (Benmerah et al., 1998; Confalonieri et al., 2000; Lamaze and Schmid, 1995; Wilde et al., 1999). The EGFR endocytic pathway primarily triggered by PKA inhibition seems also to require several kind of kinases, including other serine-threonine kinases besides PKA as well as tyrosine-kinases. In fact, the effect of H89 on EGFR endocytosis diminished gradually at concentrations higher than 50 µM H89 and disappeared at 300 µM, most likely due to inhibition of additional serine-threonine kinases. A tyrosine-kinase distinct from that of the EGFR, that might correspond to src, is also involved since the endocytic effect of H89 was abolished by genistein and we showed that neither the tyrosine-kinase activity of the receptor nor its tyrosine-phosphorylation were required.

There is increasing interest in disclosing control systems able to modulate intracellular trafficking of ligand-activated EGFR since it continues signaling from endocytic compartments
before degradation (Carpenter, 2000; Ceresa and Schmid, 2000; Di Fiore and Gill, 1999; Schlessinger, 2000; Trowbridge et al., 1993; Wiley and Burke, 2001). Recent studies have shown that PKC activation prevents EGFR ubiquitylation by c-Cbl ligase and causes the receptor to recycle to the cell surface instead of being sorted to lysosomes (Bao et al., 2000). Our results indicate that PKA could be an additional regulatory element acting in an opposite way to PKC, that is, fostering segregation of EGF/EGFR complexes from early endosomes to degradation. In fact, H89 caused a delay in the trafficking of ligand-activated EGFR out of early endosomes, as indicated by an expanded time of colocalization with Tf-TxR, determining a two-fold increased half-time of the receptor down-regulation, whereas its internalization rate remained unchanged. Both ligand-induced autophosphorylation as well as ubiquitylation of the EGFR were retarded and reduced by H89, and this probably interfered with the ubiquitylation-dependent process that drives the EGFR into its degradation pathway (Bao et al., 2000; Levkowitz et al., 1998). Interestingly, both modifications occurred at a time when most receptors were already internalized under EGF stimuli, suggesting that when PKA is inhibited the EGF/EGFR complexes start to signal from intracellular compartments. This might provide a model system to compare cell surface versus intracellular signaling, an issue of great interest (Carpenter, 2000; Ceresa and Schmid, 2000; Di Fiore and Gill, 1999; Schlessinger, 2000; Trowbridge et al., 1993; Wiley and Burke, 2001).

There are two isoforms of PKA, defined PKA-I and PKA-II, which share identical catalytic subunits but differ in their regulatory subunits, RI and RII respectively (Taylor et al., 1990). It is tempting to speculate that PKA-II is the one involved in controlling the EGFR endocytic behavior since it has been localized in membranes and organelles, including Golgi complex, plasma membrane and early endosomes, whereas PKA-I is cytosolic (Griffiths et al., 1990). In principle, changes in the EGFR endocytic behaviour induced by decreasing PKA activity could result from variations in the expression of these enzyme isoforms, known to occur during differentiation, cell growth and neoplastic transformation (Ciardiello and Tortora, 1998).

Our present results together with recent observations on the regulation of phosphodiesterase (Grange et al., 2000) allow to propose an attractive mechanism of transmodulation of EGFR
endocytic behavior by other growth factors and hormones. PKA activity decreased when cAMP levels diminished as the result of stimulation of c-AMP-phosphodiesterase PDE4D3 by endogenously produced phosphatidic acid (PA) (Grange et al., 2000). The ubiquitous enzyme phospholipase D (PLD), that produces PA and plays an important role in vesicular trafficking (Roth, 1999), is transiently activated by multiple external stimuli, including hormones and growth factors, acting through G-protein-coupled or tyrosine-kinase cell surface receptors (Jones et al., 1999). Therefore, it is expected that stimuli leading to activation of PLD should result in decreased PKA activity conducive to EGFR endocytosis. Furthermore, this mechanism could be also operative in the ligand-induced EGFR endocytosis. EGF is able to activate PLD (Yeo and Exton, 1995) and PLD inhibition decreases ligand-induced EGFR endocytosis (Shen et al., 2001). Interestingly, we showed a 26% inhibition of PKA activity in N2a-15 cells stimulated by EGF.

The identification of PKA regulated elements responsible for maintaining the EGFR at the cell surface would be necessary to further understand the receptor endocytic mechanisms and to disclose potential new targets for the interference with its function in cancer. It would be also of interest to assess whether other members of the EGFR family respond to H89 in a similar manner.
ACKNOWLEDGEMENTS

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LEGENDS TO FIGURES

Figure 1: Effect of H89 on the internalization and degradation kinetics of previously bound 125I-EGF

Transfected N2a-15 cells were incubated in the absence or presence of 20 µM H89 for 1 h at 37°C and then with 125I-EGF (50 ng/ml) for 2 h at 4°C to saturate the EGFR at the cell surface. After washing off unbound ligand, the cells were warmed to 37°C and incubated for different periods of time. Cells previously treated with H89 were kept in the presence of the drug. 125I-EGF disappearance from the surface (A) and internalization (B) were assessed by acid wash. Degradation (C) was estimated from TCA soluble radioactivity in the media. H89 did not affect ligand-induced internalization but clearly inhibited ligand degradation.

Figure 2: H89 provokes a delay in EGFR down-regulation without affecting internalization kinetics

Permanently transfected N2a-15 cells expressing EGFRs were treated with EGF (100 ng/ml) at 37°C in the absence or presence of 20 µM H89 for the indicated time periods. (A) Immunoblot of EGFR detected with polyclonal antibody (EGFR984) and revealed by ECL. (B) The relative amount of EGFR assessed in the immunoblot is expressed as percentage of control. EGF induced almost 70% receptor degradation during the first hour versus 40% in the presence of H89. After two hours, receptor degradation became similar in cells treated with or without H89. In (C), the cells were incubated during different time periods at 37°C with 125I-EGF (50 ng/ml) alone or together with 20 µM H89. The amount of internalized (Internal) and surface-associated (Surface) radiolabel was determined as described under the “Experimental Procedures”. Data were plotted using the IN/SUR method showing a $k_e$ of 0.149 min$^{-1}$ and 0.143 min$^{-1}$ for the control and H89 treated cells, respectively.

Figure 3: H89 delayed both tyrosine phosphorylation and ubiquitylation of the EGFR

(A) N2a-15 cells were treated with EGF (200 ng/ml) alone or together with 20 µM H89 for 5, 10 and 15 min at 37°C. The cells were lysed in the presence of 200 µM sodium orthovanadate and the EGFR was immunoprecipitated with HB8506 antibody and analyzed by western blot with
either anti-phosphotyrosine (pTyr) or anti-ubiquitin (Ub) antibodies. The blots were then stripped and incubated with the polyclonal anti-EGFR antibody (EGFR984) to assess total EGFR mass. (B) Cells treated with EGF for the indicated periods of time were placed at 4°C and biotinylated with sulfo-NHS-Biotin to determine the percentage of EGFR remaining on the cell surface (upper panel). In another experiment (lower panel), the cells were first incubated with reducible sulfo-NHS-SS-Biotin at 4°C and then placed at 37°C in the presence or absence of EGF. At the indicated time points, the sulfo-NHS-SS-Biotin in the cell surface was reduced with 50 mM glutathion at 4°C. Only internalized EGFRs remain biotinylated. In both experiments the EGFRs were immunoprecipitated and analyzed by western blot with streptavidin-HRP and ECL. Note that about 90% of the receptor mass was already internalized after 10 min of EGF incubation.

**Figure 4: H89 diminishes both high and low affinity binding sites and the amount of empty EGFR at the surface of transfected N2a cells.**

(A) Scatchard analysis of 125I-EGF (327100 cpm/pmol) shows that 20 µM H89 provoked 50% reduction of total number of binding sites and disappearance of the high affinity component. (B) the amount of EGFR at the plasma membrane was assessed by cell-surface biotinylation, followed by immunoprecipitation and streptavidin-peroxidase detection (lanes 1 and 2), and by immune-detection of the receptor at the cell surface of intact 35S-methionine-cysteine labeled cells, using the anti-EGFR monoclonal antibody HB8506, followed by precipitation of immune complexes with Protein A-Sepharose (lanes 3 and 4). In each case the cells were incubated for 1 h in the absence (lanes 1 and 3) or presence of 20 µM H89 (lanes 2 and 4). Both methods show a decreased mass of empty EGFR at the cell surface of about 50% when treated with H89. Total EGFR immunoprecipitated from N2a-15 cells did not change under the effects of H89 (lanes 5 and 6). (C) Immunoblot of 300,000 cells treated for up to 4 h with H89 20 µM shows no degradation of the EGFR. (D) N2a-15 cells were incubated with or without H89 20 µM for 1 h at 37°C and then analyzed by immunofluorescence with HB8506 anti-EGFR monoclonal antibody and FITC-anti-mouse. An intense intracellular EGFR staining appears upon H89 treatment. Bars, 10 µm.
Figure 5: H89 induces the internalization and accumulation of EGFR in intracellular compartments

(A) Internalization of the EGFR was monitored by incubation with EZ-Link™ sulfo-NHSS-Biotin at 4°C and incubation at 37°C in the absence or presence of H89 for 5, 10 and 15 min. After reduction with glutathion, the EGFR was immunoprecipitated with the HB8506 antibody and Protein A Sepharose, and then analyzed by western blot with streptavidin-HRP and ECL. Endocytosis occurred already after the first 5 min of H89, as detected by resistance of biotinylated-EGFR to treatment with glutathion at 4°C. (B) N2a-15 and Her14 cells incubated for the indicated times at 37°C with 20 μM H89, were fixed with 4% paraformaldehyde and permeabilized with 0.2% Triton X-100. EGFR was detected by immunofluorescence with HB8506 anti-EGFR monoclonal antibody and FITC-anti-mouse. Bars, 10 μm.

Figure 6: EGFR internalization induced by H89 does not involved tyrosine-phosphorylation, tyrosine-kinase activity or ubiquitylation of the receptor

(A) Transfected N2a-15 cells expressing EGFR were incubated for the indicated times with either 20 μM H89 (lanes 2-4) or 100 ng/ml EGF (lanes 5-7) at 37°C. The EGFR was immunoprecipitated with the monoclonal antibody-HB8506 and its phosphotyrosine content was assessed by immunoblot with an antiphosphotyrosine monoclonal antibody (HB8190). H89 did not provoke a detectable increase in tyrosine phosphorylation of the EGFR, as it can be clearly seen after EGF treatment. (B) K721A cells expressing a kinase-minus EGFR were incubated with or without 20 μM H89 and then fixed, permeabilized and the cellular distribution of the EGFR analyzed by indirect immunofluorescence. H89 induced internalization and intracellular accumulation of empty EGFR. (C) Her14 cells were incubated with 20 μM H89 (lanes 2-6) or 200 ng/ml of EGF (lanes 8-12) for different times at 37°C. The EGFR was then immunoprecipitated with anti-EGFR antibodies (HB8506) and analyzed by immunoblot with anti-ubiquitin antibody. H89 did not induce ubiquitylation of the receptor. The membranes were stripped and incubated with polyclonal anti EGFR antibody (EGFR984).
**Figure 7: H89 induces internalization of empty EGFR through a clathrin-mediated pathway involving a tyrosine-kinase activity distinct from the receptor.**

(A) Endocytosis of the EGFR was stimulated with either 20 µM H89 for 1 h or 100 ng/ml EGF for 10 min in control conditions or after cytosol acidification or potassium depletion in Her14 cells. Biotinylation of cell surface proteins was then done at 4ºC and the EGFR was immunoprecipitated with HB8506 and its biotinylation analyzed by western blot with streptavidin-HRP and ECL. Disappearance of the EGFR from the cell surface was lowered by these two procedures that inhibit clathrin-mediated endocytosis. The result is representative of three independent experiments. (B) Her14 cells were treated with or without 10 mM MβCD for 30 min or incubated for 60 min with or without 50 µM genistein, a tyrosine-kinase inhibitor, prior to the addition of 20 µM H89 for an additional 10 min at 37ºC. The cells were fixed with 4% paraformaldehyde, permeabilized with 0.2% Triton X-100 and analyzed by indirect immunofluorescence with HB8506 antibody for EGFR distribution. Both MβCD and genistein abrogate H89-induced endocytosis of EGFR. Since we previously showed that under H89 the intrinsic tyrosine-kinase of the receptor is not activated, other tyrosine-kinase activities should be involved.

**Figure 8: Correlation between H89 effects on PKA inhibition and decrease of ^125^I-EGF binding in transfected N2a cells.**

Cells were treated with different concentrations of H89 for 1 h at 37ºC (A and C), or for different times with 20 µM H89 (B and D), or 100 ng/ml of EGF alone (B) and then the PKA activity was assessed in cell extracts with the Signa TECT™ kit of Promega (A and B), and ^125^I-EGF binding was determined using saturation conditions (C and D). Concentration dependency of PKA inhibition *in vivo* by H89 showed an IC\textsubscript{50} of 12.2 µM (insert), in correlation with the IC\textsubscript{50} 13.3 µM observed for the decrease in ligand binding within a range of 5-20 µM H89, while at higher concentrations the effect became gradually lost. Almost maximal effect of 20 µM H89 was achieved between 5-15 min for PKA inhibition and at about 30 min for the decrease in ^125^I-EGF binding. Strikingly, EGF provoked 26% inhibition of PKA in N2a-15 cells.
Figure 9: H89 and the cell permeable peptide Myr-PKI induce internalization of EGFR in HeLa cells

HeLa cells were treated with or without 20 µM H89 or 100 µM Myr-PKI (14-22) peptide for the indicated time periods at 37ºC. Cells were fixed with methanol for 2 min at room temperature and then incubated with HB8506 (1/25) antibody and anti-mouse FITC. Both PKA inhibitors showed a similar endocytic effect upon EGFR. Linear arrays of vesicles or vesicle-tubular structures were seen, more frequently under H89 treatment. Bars, 10 µM

Figure 10: Different endocytic behavior of EGFR induced by H89 versus EGF

Her14 cells were incubated in the presence of EGF (100 ng/ml) or H89 (20 µM) or both, together with Tf-TxR (50 µg/ml) for the indicated periods of time at 37ºC. The cells were then washed in PBS, fixed with 4% paraformaldehyde and permeabilized with 0.2% Triton X-100 for immunofluorescence with HB8506 anti-EGFR monoclonal antibody and anti-mouse FITC. Merged pictures of the EGFR (green) and Tf (red) are presented. The EGFR stimulated by EGF colocalizes (yellow) with Tf in peripheral vesicles mostly during the first 10 min whereas segregation to different compartments is clearly apparent at 30-60 min. In contrast, during H89 treatment the EGFR co-localizes all the time with Tf and ends up accumulated in a juxtanuclear compartment seen as a bright spot. Incubation with EGF together with H89 shows frequent tubular endocytic compartments and colocalization of EGFR with Tf-TxR as a predominant pattern, even after 30-60 min of incubation, suggesting a delay in receptor segregation out of early endosomes. Bars, 10 µm.

Figure 11: H89 provoked changes in the intracellular distribution of transferrin receptors but did not affect its internalization rate and did not cause internalization of the G-protein-coupled µ-opioid receptor.

(A) HeLa cells were incubated during the indicated time periods with 20 µM H89 and the distribution of the TfR was analized by indirect immunofluorescence. The pattern of the its intracellular distribution changed first to linear arrays of vesicles converging toward perinuclear and then perinuclear vesicles were predominantly seen after 60 min of treatment. (B) For the internalization assays of 125IIdiTf, the cells were incubated in serum-free media during 1 h at 37ºC
before the experiments and then incubated with 250 ng/ml of \(^{125}\text{I}\text{dI Tf}\) for 2 h at 4\(^\circ\)C. After extensive washes with PBS-BSA 0.1\% to eliminate the unbound \(^{125}\text{I}\text{dI Tf}\), internalization was allowed by incubating the cells for different time periods at 37\(^\circ\)C, in the presence or absence of 20 \(\mu\text{M}\) H89. H89 did not affect the internalization rate determined by the IN/SUR method. (C) N2a cells permanently transfected with flag-tagged \(\mu\)-opioid receptors were incubated with either 20 \(\mu\text{M}\) H89 or their ligand DAMGO (10 \(\mu\text{M}\)) for 15 min. No evidence of internalization was observed with H89 in contrast with the endocytic pattern induced by the ligand. Bars, 10 \(\mu\text{M}\).
REFERENCES


Fig. 1
Salazar and González, 2001
Fig. 2
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Fig. 3
Salazar and González, 2001
Fig. 4
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Fig. 6
Salazar and González, 2001
A

- H89  EGF

Control

Cytosol acidification

K⁺ Depletion

B

Control  H89

MβCD + H89  Genistein + H89

Fig. 7
Salazar and González, 2001
Fig. 8
Salazar and González, 2001
Fig. 9
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EGFR
Transferrin-TxR

Fig. 10
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Fig. 11
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