Functional Reconstitualion of the Human Epidermal Growth Factor Receptor System in *Xenopus* Oocytes

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Abstract. We have expressed the human EGF receptor (hEGF-R) in *Xenopus* oocytes by injecting mRNA synthesized in vitro using SP6 vectors containing receptor cDNAs. Each oocyte could express over $1 \times 10^6$ receptors of a single affinity class and these were able to bind and rapidly internalize EGF. Occupancy resulted in receptor tyrosine autophosphorylation, downregulation, and release of intracellular calcium. Occupied receptors also rapidly induced meiotic maturation in stage VI oocytes. Receptors lacking tyrosine kinase activity bound EGF normally, but did not downregulate or induce any biological responses. The rate of oocyte maturation was proportional to hEGF-R occupancy and was significantly faster than progesterone-induced maturation at nanomolar EGF concentrations. Mutant hEGF-R truncated at residue 973 displayed identical phenotypes in both mammalian cells and oocytes in that they were defective in their ability to release intracellular calcium, undergo ligand induced internalization and receptor downregulation. However, these receptors were fully capable of inducing oocyte maturation. The remarkable retention of specific biological activities of different hEGF-R in the context of oocytes suggests that this receptor system interacts with generally available cellular components that have been conserved during evolution. In addition, it suggests that cell surface tyrosine kinase activity may play an important role in regulating resumption of the cell cycle.

A central question in biology is the mechanism by which growth factors regulate entry into the cell cycle. Polypeptide mitogens such as EGF bind to specific surface receptors which possess intrinsic tyrosine kinase activity (Carpenter, 1985; Cohen, 1983). Phosphorylation of cellular substrates on tyrosine residues leads to a multitude of responses such as ion transport (Chen et al., 1987), induction of receptor internalization (Glenney et al., 1988), gene induction, and eventually cell division (Carpenter, 1985; Chen et al., 1987). Although receptor tyrosine kinase activity is necessary for the mitogenic action of EGF, it has been difficult to identify those events and cellular substrates directly involved in regulating the cell cycle. Partly this is due to the plethora of responses to growth factors and the asynchronous movement of mammalian cells through the cell cycle. It is thus difficult to identify a particular biochemical event as being crucial to either a specific response or to a responsive subpopulation of cells. Indeed, experimental systems which do not suffer from these drawbacks, such as yeast and *Xenopus* oocytes, have proven to be more amenable to studies on the biochemical regulation of the cell cycle (Maller, 1987; Simanis et al., 1987).

Recent studies in *Xenopus* oocytes have identified the homologue of the yeast *cdc2* gene product as a central component in the progesterone-mediated resumption of meiosis (Dunphy et al., 1988; Gautier et al., 1988). This protein (p34<sup>cdc2</sup>) is a component of the maturation promoting factor (MPF) that appears during meiosis and has serine kinase activity (Labbe et al., 1989; Simanis et al., 1987). Numerous studies have focused on the regulation of p34<sup>cdc2</sup> by other proteins (Draetta et al., 1989; Dunphy and Newport, 1989) and p34<sup>cdc2</sup> has been identified as a major substrate of tyrosine kinases in mammalian cells (Draetta et al., 1988). Despite a growing body of evidence on how p34<sup>cdc2</sup> is regulated in cycling cells, little is known regarding the mechanisms of its initial activation (Maller, 1988). Although most studies have concluded that phosphorylation of p34<sup>cdc2</sup> on tyrosine residues inhibits its activity (Dunphy and Newport, 1989; Morla et al., 1989), tyrosine kinase activity has been shown to facilitate oocyte maturation (Maller, 1987; Morgan et al., 1986). Unfortunately, oocytes lack sufficient numbers of receptors possessing tyrosine kinase activity for biochemical analyses (Maller and Koontz, 1981). In addition, direct microinjection of tyrosine kinases such as the insulin receptor (Maller, 1987) and src kinase (Spivack et al., 1984) into oocytes does not trigger maturation. Thus the relationship between activation of growth factor receptor tyrosine kinase activity and the p34<sup>cdc2</sup> pathway remains obscure.

Since oocytes express foreign proteins when injected with a suitable mRNA (Dawid and Sargent, 1988), we sought to...
develop a method for the expression of high levels of human EGF receptors (hEGF-R) in oocytes. Since a number of cDNA vectors containing site-directed mutants of the hEGF-R are available (Chen et al., 1987), this approach allows manipulation of the receptor system at the molecular level. By using SP6 vectors to direct the in vitro synthesis of large amounts of mRNA for oocyte injection (Krieg and Melton, 1987), we obtained the expression of over 1 × 10^10 receptors per oocyte. Significantly, we found that the hEGF-R can function normally in oocytes and its tyrosine kinase activity can trigger meiosis.

Materials and Methods

Oocyte Isolation and Culture

Xenopus laevis were purchased from the South African Snake Farm (Fish Hoek, Cape Province) and maintained as previously described (Opresko and Karpf, 1987). Animals were injected with 500 IU HCG at least 2 wk before oocyte isolation by manual dissection as previously described (Opresko and Karpf, 1987). Oocytes were maintained in 50% L-15 medium containing 5% calf serum and 1% BSA (Wallace et al., 1980) for 18 h before injection with mRNA. The oocytes were injected with ~20 ng mRNA in 10–20 nl of water and maintained in the above medium for 40 h before evaluation. All injections were performed with a custom pneumatic injector calibrated by video microscopy.

Construction of the hEGF-R Vector

The pLOLB vector was constructed from the pSP64 (polyA) vector (Promega Biotech, Madison, WI) by replacing the Eco RI linearization site with a Not I site. The pXER vector containing the full-length cDNA for the hEGF-R as well as the M^121 and c^973 truncation mutations was obtained from Drs. Gordon Gill and Michael Rosenfeld (University of California, San Diego, CA). The hEGF-R cDNAs were excised by cutting with Hind III, end-filling with Klenow fragment, and then cutting with Xba I. The cDNAs were cloned into the pLOLB vector using the Xba I and Sma I sites, yielding the pOBER vector. The SP6 RNA polymerase transcription initiation site was 95 nucleotides upstream of the hEGF-R coding region which terminated 310 nucleotides from the polyA tail. This plasmid was linearized with Not I and mRNA was transcribed using SP6 RNA polymerase (Krieg and Melton, 1987). To obtain capped mRNA, diguanosine triphosphate was present in a fivefold molar excess to rGTP. The newly synthesized mRNA was treated with RNAase-free DNase, extracted with chloroform/phenol, precipitated and resuspended in DEPC-treated water to a concentration of 0.5–1.5 µg/µl and frozen in small aliquots. The concentration of mRNA was determined spectrophotometrically after 0.50 column chromatography.

Receptor Labeling

The hEGF-R was biosynthetically labeled by incubating mRNA-injected oocytes for 48 h in bicarbonate-free, DME with 10% normal methionine and cysteine, 10% dialyzed calf serum, and 20 mM Hepes buffer, pH 7.6 (Wallace and Misulovin, 1978). This medium was diluted 1:1 with 20 mM Hepes buffer and contained 1 mCi/ml [35S]Translabel (ICN Radiochemicals, Irvine, CA). Groups of five oocytes were homogenized in 1 ml of 12.5 mM CHAPS, 10 mM NaCl, 1 mM EDTA 10 mM Tris-Cl, pH 8.0 and 100 µg/ml each aprotinin, leupeptin, chymostatin, and pepstatin, and 3 mM PMSF. The insoluble residue was removed by centrifugation and the labeled hEGF-R was immunoprecipitated with the 528 mouse monoclonal antibody (Gill et al., 1984). Mouse B22 cells expressing high levels of the tyrosine kinase active (Kin') hEGF-R (Glenney et al., 1988) were labeled overnight as described above using normal strength DME and 50 µCi/ml of [35S]Translabel. Solubilized immunoprecipitates were subjected to SDS gel electrophoresis using a 7.5% gel. Gels were treated with EN3HANCE before fluorography. Molecular weights of the labeled bands were determined using prestained molecular weight markers (Bio-Rad Laboratories, Richmond, CA).

The presence of phosphotyrosine in the hEGF-R was determined by Western blot analysis. Groups of oocytes injected 40 h previously with the indicated mRNA were incubated either with or without EGF. After a 30-min extraction at 0°C with the CHAPS solution described above containing 10 mM NaF and 0.1 mM Na_2VO_4, SDS and DTT were added to 1% and 12 mM respectively. Samples were boiled and subjected to SDS gel electrophoresis using 5–15% gradient gels. The proteins were transferred to nitrocellulose using a PolyBlot apparatus (Fisher Scientific Co., Pittsburgh, PA) and probed with [125I]-labeled antiphosphotyrosine monoclonal antibody PY20 (ICN Biochemicals Inc., Cleveland, OH) as previously described (Glenney et al., 1988).

Histone Kinase Assay

The assay for the activation of oocyte histone H1 kinase was based on the protocol previously described (Labbe et al., 1988). Groups of three oocytes were rinsed once and homogenized in 20 µl of 50 mM β-glycerophosphate, 10 mM MgCl_2, 7.5 mM EGTA, 1 mM DTT; 20 mM Hepes, pH 7.5 and 100 µg/ml each aprotinin, leupeptin, chymostatin, and pepstatin. Insoluble material was removed by centrifugation for 1 min at 8,000 rpm in a tabletop swinging bucket microfuge (Savant Instruments, Inc., Hicksville, NY) maintained at 4°C. Homogenates (15 µl) were mixed with 5 µl of histone reaction mixture (5 mg/ml lysine-rich histones (HuS from Sigma Chemical Co., St. Louis, MO), 15 mM MgCl_2, 1 mM ATP, and 50 µCi/ml [γ-32P]ATP from Amersham Corp., Arlington Heights, IL). After 10 min, the reaction was terminated by adding 40 µl of 2% SDS, 2% β-mercaptoethanol, and boiling for 2 min. The samples were subjected to SDS gel electrophoresis using 12% gels followed by staining, drying, and autoradiography.
ph. Quantitation of the amount of label in the bands was determined using a densitometer (model 620; Bio-Rad Laboratories). Exposure times of the autoradiographs were adjusted to remain in the linear range of the densitometer as determined by parallel samples of known specific activities.

**EGF Binding to Oocytes**

1 h before binding, the cells were placed into fresh 50% L-15 solution containing 125 μg/ml sodium ipodate which inhibits the deiodination activity of oocytes (Opresko et al., 1980). 125I-EGF was prepared as previously described (Wiley and Cunningham, 1982). Scatchard analyses (Scatchard, 1949) were performed at 0°C to prevent internalization. Oocytes were brought to equilibrium (4-6 h) with 125I-EGF concentrations ranging from 5 × 10^{-10} to 1.6 × 10^{-7} M. The labeled EGF was not diluted with unlabeled EGF to avoid potential problems associated with differential binding between labeled and unlabeled ligand (Wiley, 1985). A minimum of 10 oocytes were used to evaluate binding at each 125I-EGF concentration. Non-specific binding was determined in parallel using water-injected oocytes and was always <10% of total binding. To allow internalization, oocytes were incubated at 20°C for the indicated times. After the incubation, cells were washed in three changes of solution O-R2 and placed in the upper chamber of an in vitro synthesis of hEGF-R mRNA containing a 30-residue polyA tail. Injection of this polyA mRNA resulted in very efficient synthesis of hEGF-R by recipient oocytes. Shown in Fig. 1 B is the net binding of 125I-EGF to individual oocytes at both 0 and 22°C. Binding of 125I-EGF to mRNA-injected oocytes at 0°C, a temperature that prevents receptor internalization, was usually 10-fold greater than water-injected controls. However, a significant increase in 125I-EGF binding was observed at 22°C, probably as a consequence of ligand internalization. To separate the contribution of surface receptor expression from receptor internalization, we examined 125I-EGF binding at 0°C and then analyzed the ability of the expressed receptors to internalize ligand.

Groups of oocytes were injected with either water or mRNA and cultured for different lengths of time. The ability of oocytes to bind 125I-EGF at 0°C in both stage IV/V (1.0 mm) and VI (1.3 mm) oocytes was then evaluated. As shown in Fig. 2 A, significant 125I-EGF binding could be observed by 24 h and this continued to increase for the entire period of the experiment. The 1.0-mm-diam oocytes accumulated surface hEGF-R at a rate of ∼7 × 10^6 h^{-1} between 24- and 72-h postinjection while the accumulation rate of oocytes 1.3-mm diameter was ∼2.3 × 10^6 h^{-1}. Within this period, there was no sign of an upper limit in the number of receptors observed on the oocyte surface. Even after correcting for differences in oocyte volume and surface area, stage VI oocytes were still more efficient in expressing surface hEGF-R, consistent with previous observations that protein synthesis rates in these cells are significantly higher than in smaller oocytes (Wasserman et al., 1984).

The hEGF-R expressed by stage VI oocytes were of a single affinity class. Shown in Fig. 2 B is a Scatchard plot of 125I-EGF binding to mRNA-injected oocytes at 0°C. Non-specific binding to water-injected oocytes at 0°C usually ranged between 5 and 10% and equilibrium binding was did not result in any observable 125I-EGF binding by oocytes. We therefore obtained the pXER expression vector containing the cdNA encoding the full-length hEGF-R (Lin et al., 1986). Direct microinjection of this vector into the oocyte nucleus resulted in a significant, but highly variable binding of 125I-EGF to recipient cells (data not shown). To improve the reproducibility of hEGF-R expression, we constructed the vector shown in Fig. 1 A. This vector supports the in vitro synthesis of hEGF-R mRNA containing a 30-residue polyA tail. Injection of this polyA mRNA resulted in very efficient synthesis of hEGF-R by recipient oocytes. Shown in Fig. 1 B is the net binding of 125I-EGF to individual oocytes at both 0 and 22°C. Binding of 125I-EGF to mRNA-injected oocytes at 0°C, a temperature that prevents receptor internalization, was usually 10-fold greater than water-injected controls. However, a significant increase in 125I-EGF binding was observed at 22°C, probably as a consequence of ligand internalization. To separate the contribution of surface receptor expression from receptor internalization, we examined 125I-EGF binding at 0°C and then analyzed the ability of the expressed receptors to internalize ligand.

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Figure 3. The human EGF receptor expressed by oocytes is full length and displays EGF-induced autophosphorylation. (A) Immunoprecipitation of metabolically labeled hEGF receptor. Oocytes were injected with mRNA encoding either the normal hEGF-R (Kin+) or the hEGF-R in which lysine at position 721 was altered to a methionine by site-directed mutagenesis (Kin-). Metabolically labeled hEGF-R from mouse B82 cells were used as a control. After 40-h labeling, the receptor was immunoprecipitated using the 528 monoclonal antibody. (B) Western blot analysis of EGF-induced tyrosine phosphorylation in oocytes. Stage V oocytes injected 40 h previously with the indicated mRNA were incubated either with (+) or without (−) 1.7 × 10⁻⁸ M EGF for 5 min followed by extraction and Western blot analysis using ¹²⁵I-labeled antiphosphotyrosine monoclonal PY20 (Glenney et al., 1988).

achieved by 4 h (data not shown). Although normal human fibroblasts also express receptors of a single affinity class (Carpenter et al., 1975; Knauer et al., 1984), the calculated equilibrium affinity for the hEGF-R in oocytes was 3.6 × 10⁻⁸ M, which is ~10-fold lower than that observed in human fibroblasts (Wiley et al., 1989). In the experiments shown in Fig. 2 B, each oocyte expressed ~1.5 × 10⁹ hEGF-R, or ~100-fold the number of endogenous insulin/IGF-1 receptors (Mailer and Koontz, 1981).

The absolute number of surface hEGF-R expressed by oocytes depended both on the amount of injected mRNA, the particular batch of oocytes, the oocyte stage, and the time allowed for synthesis. Typically, this number ranged between 3 × 10⁹ and 3 × 10¹⁰ receptors/oocyte. However, the surface area of a 1.0-mm oocyte is approximately that of 50,000 fibroblasts (Opresko and Wiley, 1987a), translating to a surface receptor density equivalent to between 0.6 and 6 × 10⁵ for a typical mammalian cell. Although this receptor density is high, it falls within the range reported for transfected mammalian cells and is substantially less than that observed for A431 cells (Chen et al., 1987; Haigler et al., 1979). In addition, the number of endogenous vitellogenin receptors expressed on the oocyte surface is between 0.3 and 3 × 10¹⁰ (Opresko and Wiley, 1987a). Thus the surface densities of hEGF-R expressed by oocytes is high, but well within the capacity of the oocyte plasma membrane.

We also synthesized mRNA from a hEGF-R cDNA that contains methionine rather than lysine at residue 721 (Chen et al., 1987). Receptors made from this mutant cDNA lack intrinsic tyrosine kinase activity, but are unaltered in their ability to bind EGF (Chen et al., 1987). Oocytes injected with mRNA derived from the “wild-type” (Kin+) and M¹²¹ (tyrosine kinase inactive [Kin-]) constructions synthesized equivalent numbers of receptors as a function of injected mRNA and both bound EGF with the same affinity (data not shown).

To determine whether these receptors were normally processed, oocytes were incubated with [³⁵S]methionine/cysteine followed by immunoprecipitation with a monoclonal antibody to the hEGF-R (Gill et al., 1984). As shown in Fig. 3 A, a protein was immunoprecipitated with the same mobility as the hEGF-R synthesized by transfected mammalian cells. The additional bands found in the oocyte immunoprecipitate are nonspecifically adsorbed, sulfated yolk proteins
The EGF Receptor Is Fully Functional in Oocytes and Induces Meiotic Maturation

Our ability to synthesize hEGF-R in oocytes allowed us to test which functional aspects of the receptor are intrinsically dependent on tyrosine kinase activity. For example, the Kin- hEGF-R in mammalian cells is unable to transmit a biological signal (Chen et al., 1987) or to trigger receptor downregulation (Glenney et al., 1988). Kin- receptors do not undergo downregulation because of their lack of ligand-induced internalization. We therefore examined the relative ability of Kin+ and Kin- hEGF-R to induce these events in the oocytes. To determine whether both Kin+ and Kin- hEGF-R undergo endocytosis and downregulation, we incubated oocytes with a high concentration of 125I-EGF. The extent of internalization and surface binding was then quantitated by acid stripping (Wiley and Cunningham, 1982). As shown in Fig. 4, both Kin+ and Kin- receptors internalized EGF. However, the specific internalization rate of the Kin+ receptor was about twice that of its Kin- counterpart (Fig. 4).

Figure 4. Internalization and down-regulation of the kinase positive and kinase negative hEGF-R by oocytes. (A) Binding and internalization of Kin+ (•, ○) receptors. Oocytes were incubated with 3.3 × 10^-8 M 125I-EGF (5.7 × 10^8 cpm/mmol) and at the indicated times groups of eight oocytes were rinsed and the relative amount of surface-associated and internalized 125I-EGF determined by acid stripping. (B) Binding and internalization of Kin- (○, □) receptors. Same experiment described in A, but the oocytes were injected with mRNA derived from the Kin- hEGF-R. (C) Internalization plots of the Kin+ (•) and Kin- (○) receptors. The kinetics of internalization was determined as described above using 1.7 × 10^-9 M 125I-EGF and the data were transformed into internalization plots. The slope of these plots is proportional to the specific internalization rate of the receptor, yielding a value of 0.21 min^-1 for the Kin+ receptor, and 0.11 min^-1 for the Kin- receptor.

...are also found in sham-injected oocytes (data not shown). The molecular weight of the labeled band was 170,000, indicating that the hEGF-R is fully glycosylated by oocytes (Soderquist et al., 1988).

The tyrosine kinase activity of the hEGF-R in oocytes was determined by Western blot analysis using an anti-phosphotyrosine (anti-pTyr) monoclonal antibody (Glenney et al., 1988). As seen in Fig. 3 B, only oocytes injected with mRNA encoding the Kin+ hEGF-R showed an EGF-mediated appearance of a new tyrosine-phosphorylated protein. The molecular weight of the EGF-induced band was 170,000, indicating that it is the autophosphorylated hEGF-R (Downward et al., 1984). These results demonstrate that oocytes can synthesize a fully glycosylated and enzymatically functional EGF receptor.

Figure 5. Occupied EGF receptors can induce meiotic maturation in oocytes. (A) Kinetics of EGF and progesterone-induced maturation. Oocytes injected with mRNA encoding the Kin+ (●) or Kin- (○) hEGF-R were exposed to 1.7 × 10^-8 M EGF. Alternatively, water-injected oocytes were exposed to 10 μg/ml progesterone (□). At the indicated times, they were scored for the appearance of the characteristic white spot at the apex of the animal hemisphere, indicating GVBD. (B) Effect of EGF concentration on the rate of maturation. Groups of oocytes (n = 7) expressing the Kin+ hEGF-R were treated with the indicated concentrations of EGF. The time of GVBD was then scored as indicated in A. The results are shown as the mean time of GVBD ± the standard deviation. Control oocytes were treated with 10 μg/ml progesterone.
4 C). Significantly, surface binding dropped to less than one-third the initial values by 80 min in cells expressing Kin+ receptors while surface binding for the Kin- receptor remained constant. This indicates that Kin+ receptors are undergoing ligand-induced internalization and downregulation, consistent with the observations of higher internalization rates for the Kin+ receptor. To confirm downregulation of Kin+ receptors, we conducted a "steady state" analysis of surface binding (Wiley and Cunningham, 1981) at 22°C to allow for endocytosis (Wallace et al., 1973). Oocytes were incubated with increasing concentrations of [125I]-EGF and allowed to approach a steady state of surface binding. The maximum extent of surface binding was then determined by extrapolation and compared to equilibrium binding at 0°C. The results of this experiment showed that oocytes that displayed an average of 2.4 × 10^6 receptors at 0°C (before EGF exposure) had a maximum of 8.5 × 10^6 surface receptors at steady state binding. This confirms that oocytes lose approximately two-thirds of their surface receptors by downregulation. The absence of downregulation of Kin- receptors does not seem to be due to differences in their ability to recycle relative to Kin+ receptors. EGF internalized by either receptor is transferred to a light endosomal compartment and eventually to yolk platelets (data not shown). This indicates that intrinsic kinase activity induces hEGF-R downregulation at the level of internalization in oocytes as it does in mammalian cells (Chen et al., 1987; Glenney et al., 1988).

To determine whether the hEGF-R could induce a biological response in oocytes, we examined the effect of EGF on meiotic maturation. It has already been established that incubating oocytes with high concentrations of insulin and IGF-I can facilitate and sometimes induce oocyte maturation (LeGoascogne et al., 1984; Mailer and Koontz, 1981). However, since microinjected tyrosine kinases such as the src protein or insulin receptor cannot induce maturation (Mailer, 1987; Spivack et al., 1984), a direct relationship between tyrosine kinase activity and maturation is uncertain. We therefore compared the relative ability of EGF to induce maturation in oocytes expressing either the Kin+ or Kin- hEGF-R. As shown in Fig. 5 A, EGF was very effective in inducing maturation in oocytes that expressed the Kin+ receptor. However, treatment of oocytes expressing Kin+ receptors with EGF was without effect, even after 24-h treatment. In addition, neither water-injected nor uninjected oocytes responded to EGF. Germinal vesicle breakdown (GVBD) in EGF-treated oocytes was confirmed by sectioning of fixed oocytes. Cytoplasm from EGF-matured oocytes could also induce maturation in recipient oocytes with the same kinetics as cytoplasm from progesterone-stimulated oocytes (data not shown), confirming the presence of MPF (Wasserman and Masui, 1975a).

We were quite surprised to find that EGF could induce maturation significantly faster than progesterone (Fig. 5 B). However, the rate of EGF-induced maturation was highly dependent on both the concentration of EGF and the level of hEGF-R expression at the oocyte surface. As shown in Fig. 5 B, reducing the concentration of EGF resulted in a decrease in the rate of maturation to below that observed for progesterone. When the level of surface hEGF-R expression was decreased to below 10^9 per oocyte, the maximum rate of EGF-stimulated maturation also fell below that typically induced by progesterone, but was still much faster than insulin-induced maturation (data not shown). However, we occasionally found batches of oocytes in which progesterone induced maturation nearly as fast as that induced by EGF. Therefore, the mechanism(s) which regulates the sensitivity of oocytes to progesterone does not necessarily affect their sensitivity to EGF. Nevertheless, the intrinsic tyrosine kinase activity of the EGF receptor is essential for the induction of maturation and EGF can stimulate oocyte responses at physiological ligand concentrations.

**Activation of the MPF/p34^cdc2 Complex**

Progesterone-stimulated maturation involves the activation of the MPF/p34^cdc2 complex which contains a serine/threonine histone H1 kinase activity (Arion et al., 1988; Labbe et al., 1989). The activation of the MPF/p34^cdc2 complex is thought to be essential for subsequent events in meiosis. However, the rapid maturation stimulated by EGF and its de-
Expression of an Internalization-defective hEGF-R in Oocytes

When we first examined the effect of EGF on oocyte maturation, we noted that oocytes expressing the Kin+ hEGF-R appeared morphologically normal until ~6-8 h after EGF addition. After this time, oocytes became mottled and then visibly degenerated. This is similar to what has been observed for maturation induced by both H-ras injection (Birchmeier et al., 1985), high concentrations of divalent cations and ionophores (Wasserman and Masui, 1975b), and other nonphysiological agents that can affect intracellular calcium distribution (Smith, 1989). Although the hEGF-R can release intracellular calcium in mammalian cells (Sawyer and Cohen, 1981), progesterone treatment does not induce calcium release in oocytes (Cork et al., 1987). Therefore, we felt that this additional response could be responsible for the eventual oocyte degeneration. To test this hypothesis, we made use of a recently described mutant of the hEGF-R that lacks the carboxy-terminal domain distal to residue 973 (c'973 hEGF-R). This receptor has full kinase activity, but is defective in its ability to release intracellular calcium and to undergo ligand-induced internalization (Chen et al., 1989). Significantly, the c'973 receptors are fully capable of inducing mitosis in mammalian cells, suggesting that calcium release and induced internalization are not required for induction of cell division. We were thus interested to determine whether these receptors would induce meiotic maturation in oocytes without causing oocyte degeneration.

We inserted the c'973 hEGF-R into our SP6 vector and synthesized the appropriate mRNA. When injected into oocytes, this mRNA directed the synthesis of a 150,000-kD hEGF-R protein, as determined by [35S]methionine/cysteine labeling followed by immunoprecipitation (data not shown). This is the expected size of the c'973 hEGF-R. When injected with the same amount of mRNA, 125I-EGF binding at 0°C to oocytes expressing the c'973 receptor was indistinguishable from those expressing the wild-type receptor (data not shown). We also observed that the addition of EGF did not result in autophosphorylation of the receptor molecule, confirming that all of the autophosphorylation sites had been removed by the c'973 truncation.

Since the c'973 receptor is internalization defective and...
does not undergo downregulation in mammalian cells, we were interested in confirming this receptor attribute in oocytes. As shown in Fig. 7 B, internalization plot analysis demonstrated that the specific internalization rate of the c'973 receptors was significantly lower than the K+ hEGF-R. On the average, the internalization rate of the c'973 was 0.03 min⁻¹, or between 35 and 40% of the rate observed in parallel sets of oocytes expressing the wild-type K+ hEGF-R. As expected, surface binding of 125I-EGF to oocytes expressing the wild-type hEGF-R displayed a time-dependent decrease at 20°C as a consequence of downregulation while surface binding to oocytes expressing the c'973 hEGF-R reached a constant steady state binding (Fig. 7 A). Binding at 0°C confirmed that treatment of oocytes expressing c'973 hEGF-R with high concentrations of EGF for 2 h at 20°C did not decrease the initial number of surface receptors (data not shown). We conclude that the internalization/downregulation defective phenotype of the c'973 receptor in mammalian cells is preserved in Xenopus oocytes.

To determine the relative ability of the different hEGF-R to release intracellular calcium stores in oocytes, cells expressing the appropriate receptor were loaded overnight with 45Ca, rinsed, and placed in a continuous flow chamber. The release of 45Ca from the oocytes was then evaluated after adding EGF to the perfusate. As shown in Fig. 8, the addition of EGF to oocytes expressing the wild-type hEGF-R resulted in a rapid release of preload 45Ca. Water-injected oocytes or those expressing Kin receptors displayed no response to the addition of EGF (data not shown). In contrast to the situation with the wild-type receptor, EGF addition to oocytes expressing the c'973 receptor resulted in very little release of 45Ca. However, oocytes could release calcium in response to ionomycin (Fig. 8). The relative ability of EGF to induce calcium release in oocytes expressing either wild-type or c'973 hEGF-R is essentially identical to what has been observed in mammalian cells (Chen et al., 1989).

Once we had confirmed that the c'973 hEGF-R preserved their phenotype in the Xenopus oocyte system, we examined the question of whether they could induce meiotic maturation. We found that the addition of EGF to oocytes expressing c'973 hEGF-R indeed resulted in rapid meiotic maturation. However, the rate of maturation we observed was not significantly different than what we had observed after stimulation of oocytes expressing the wild-type receptor. In addition, the number of occupied receptors required for induction of maturation was similar between the wild-type and c'973 receptors (data not shown). This indicates that the signaling mechanism required for the induction of maturation is not significantly different between the two receptor constructions. However, as shown in Fig. 9, the morphology of oocytes matured in response to the c'973 hEGF-R was indistinguishable from that displayed by oocytes that matured in response to progesterone. Prolonged incubations with EGF did not result in any visible abnormalities or morphological changes, other than those normally associated with maturation. Thus the removal of the internalization/calcium domain from the hEGF-R results in a receptor which is fully capable of inducing meiotic maturation, but which does not trigger the negative side effects observed with the wild-type receptor.

**Discussion**

Our results show that the human EGF receptor system can be functionally reconstituted in Xenopus oocytes. Other investigators have previously reported the synthesis of hEGF-R by oocytes after injection of mRNA derived from A431 cells (Simmen et al., 1984), but insufficient numbers were obtained for biological or biochemical studies. Our approach of synthesizing mRNAs in vitro from cDNA vectors allowed us to obtain sufficient amounts of mRNA to cause high levels of receptor expression. Indeed, the levels of surface receptor expression we observe (over 10¹⁰ per oocyte) approach that of many transfected mammalian cells (Chen et al., 1989).

The Xenopus oocyte is particularly useful for investigating the EGF receptor because they constitute a large, synchronized cell population with well-defined, stage-specific responses to growth factors (Maller, 1987; Taylor and Smith, 1987; Wallace and Misulovin, 1978). They also lack endogenous EGF receptors. The lack of 125I-EGF binding to oocytes that we and other investigators have observed is not simply due to their inability to bind mammalian EGF since the ligand can readily bind to Xenopus hepatocytes at nanomolar concentrations as well as induce DNA synthesis and receptor autophosphorylation (Wolffe et al., 1985). The

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**Figure 9.** EGF-induced maturation of oocytes expressing the c'973 hEGF-R. (A) Uninjected oocytes treated for 8 h with 10 μg/ml progesterone. (B) Oocytes were injected with 20 ng of mRNA encoding the c'973 hEGF-R and 40 h later were treated with 1.7 × 10⁻⁹ M EGF for 8 h. The EGF-treated oocytes matured at 3 h while the progesterone-treated cells matured at 4 h.
lack of endogenous EGF receptors allows one to determine which functional aspects of the receptor system are intrinsic to the receptor molecule itself, and those which require cellular components that are EGF receptor specific. The results of our study indicate that virtually all important aspects of this receptor system are intrinsic to the receptor molecule itself.

Analysis of ¹²⁵I-EGF binding to oocytes revealed a single affinity class of receptors at both 0°C (Fig. 1 B) and at 22°C (data not shown). However, this affinity (∼36 nM at both temperatures) was quite low relative to mammalian cells (Wiley et al., 1989). It has been reported that some cell types display multiple classes of EGF receptors, with the “high affinity” type being the ones active in signal transduction (Defize et al., 1989). However, many cells display only a single, low affinity class of EGF receptors and yet still retain full responsiveness to EGF (Knauer et al., 1984). We have previously proposed that the apparent affinity of cells for EGF can be dependent on factors extrinsic to the receptor itself, such as cell geometry, receptor density, and distribution (Wiley, 1985, 1988). The low affinity that we observe for the hEGF-R is most likely due to geometric and physical aspects of oocytes (such as the thick vitelline envelope) since the endogenous oocyte vitellogenin and insulin receptors are both primarily low affinity (Mailer and Koontz, 1981; Opresko and Wiley, 1987a). A significant connection between receptor affinity and signal transduction is unlikely, since despite the low affinity of hEGF-R expressed in oocytes, we were able to generate significant biological responses at subnanomolar EGF concentrations.

The hEGF-R binds and rapidly internalizes EGF (Fig. 4 C) at rates comparable to mammalian cells (Wiley, 1985). EGF also induced endocytic downregulation (Fig. 4 A). The role of receptor tyrosine kinase activity in these processes is the same in oocytes as in mammalian cells (Chen et al., 1987; Glenney et al., 1988) since it was required both for maximal rates of internalization and receptor downregulation. Removal of the carboxy region distal to residue 973 also eliminated downregulation as observed in mammalian cells (Chen et al., 1989). However, absolute differences between the specific internalization rates of the Kin+ and Kin- receptors were not as pronounced in oocytes as in mammalian cells. Typically, we have observed a 6–10-fold greater internalization rate of the Kin+ versus Kin- receptor in transfected mammalian cells (Chen et al., 1989), but we only observed a two- to three-fold difference in oocytes. However, since downregulation of the hEGF-R is due to ligand-induced (kinase-dependent) internalization (Chen et al., 1989), we expected to see correspondingly less downregulation. This was indeed the case since the extent of hEGF-R downregulation was only ∼66% in oocytes as compared to the 80–95% observed in mammalian cells (Chen et al., 1989; Knauer et al., 1984). The source of this quantitative difference between the oocyte and mammalian systems is currently unknown, but most likely is a result of the extremely high endocytic capacity of Xenopus oocytes (Opresko and Wiley, 1987b) since this would narrow differences between constitutive and ligand-induced internalization rates. Nevertheless, the dependency of downregulation on both kinase activity and specific domains in the carboxy region of the receptor argues that the mechanisms of EGF-induced internalization are probably the same in all cells.

The hEGF-R could also trigger a rapid release of intracellular calcium from oocytes. As was the case with mammalian cells (Chen et al., 1989), the ability of the hEGF-R to mediate this response was severely attenuated by removal of the receptor domain distal to residue 973 (Fig. 8). Removal of this sequence also decreased receptor internalization and prevented ligand-induced receptor downregulation (Fig. 7). To date, all phenotypic differences that have been observed between different mutant hEGF-R in mammalian cells have also been observed in Xenopus oocytes. The remarkable retention of specific biological activities of the hEGF-R in the context of oocytes suggests that this receptor system interacts with generally available cellular components that have been conserved during evolution.

We were quite surprised at the ability of the hEGF-R to rapidly induce meiotic maturation. A wide variety of pharmacological agents, receptors, oncogenes, kinases, and enzymes have been injected into oocytes to determine their ability/role in inducing meiotic maturation (Birchmeier et al., 1985; Huchon et al., 1981; Maller, 1987; Sehgal et al., 1988; Smith, 1989). This usually results in a “facilitation” of progesterone-induced maturation, but rarely maturation itself. Tyrosine kinases in particular have been poor inducers of maturation. Although directly injected src kinase (Spivack et al., 1984) and insulin receptors (Maller, 1987) will increase ribosomal protein S6 phosphorylation, they do not trigger maturation. Insulin and IGF-1 will facilitate progesterone-induced maturation in isolated oocytes (Hirai et al., 1983; LeGoascegne et al., 1984), but trigger maturation only rarely (Wallace and Misulovin, 1980) unless nonphysiological conditions are used, such as removal of potassium from the medium (Cicirelli et al., 1988). We have never observed insulin-induced maturation in Xenopus oocytes using standard conditions (i.e., physiological salines), although we have shown that insulin is a potent stimulator of vitellogenin uptake and oocyte growth (Opresko and Wiley, 1987b). Therefore, the role of tyrosine kinases in the induction of oocyte maturation has been uncertain. Our result with the reconstituted hEGF-R is the first direct evidence that tyrosine kinases can trigger the resumption of meiosis in oocytes. Other than MPF itself, the hEGF-R seems to be the most effective inducer of meiotic maturation yet described. This may be partially due to the high levels of receptors that are expressed, since lower receptor occupancies resulted in progressively slower response times (Fig. 5 B). However, the levels of hEGF-R we achieve in our system are less than those used by other investigators who directly microinjected maturation-inducing proteins such as the Harvey ras oncogene (Birchmeier et al., 1985). In addition, occupancy of only a small fraction of surface receptors was required to induce maturation (<5%). As a function of surface area, these numbers are comparable to the receptor levels available on Xenopus hepatocytes (Wolfe et al., 1985) and normal human fibroblasts (Carpenter et al., 1975) and are similar to the number of endogenous IGF-1/insulin receptors (Maller and Koontz, 1981). As a function of cell mass, oocyte expression of the hEGF-R is only a small fraction of the levels displayed by other cells. Thus the ability of the hEGF-R to induce maturation does not appear to be a nonspecific result of receptor overexpression, but instead to be an intrinsic property of the receptor tyrosine kinase activity.

Even though the hEGF receptor is capable of inducing
capable of inducing maturation. Indeed, a requirement for maturation faster than progesterone, it does not appear to in which to investigate these types of problems.

The Journal of Cell Biology, Volume 111, 1990 1670

It is not yet certain whether internalization of the hEGF-R is necessary to induce maturation, but our data support the hypothesis that it acts to phosphorylate a substrate at the cell surface. Although the c973 hEGF-R was only internalized at one-third the rate of its wild-type counterpart, it was fully capable of inducing maturation. Indeed, a requirement for activated receptors to reside at the oocyte surface would explain why previous efforts to demonstrate a role for tyrosine kinases in maturation have failed, since those studies directly microinjected kinases into the oocyte cytoplasm (Maller, 1987; Spinavc et al., 1984). Direct microinjection of progesterone also fails to induce maturation (Smith and Ecker, 1971), again suggesting that maturation triggered by hormone receptors may require action at the cell surface.

The ability of the hEGF-R to induce meiotic maturation in oocytes suggests that the fundamental processes regulating the eucaryotic cell cycle are highly conserved during evolution. Our results also point to a central role for tyrosine kinases in both the activation and regulation of this process. However, it is noteworthy that maturation induced by mutant hEGF-R that are deficient in triggering some biological responses, such as calcium release, more closely resembles that produced by physiological agents. Since progesterone treatment does not release intracellular calcium in oocytes (Cork et al., 1987), this is an "inappropriate" hormonal response. The ability of wild-type hEGF-R to trigger responses typical of its normal context suggests that the response machinery is similar in all cells and that different receptors function to orchestrate different responses. Correct induction of a final cellular response may require selective stimulation of the machinery. Indeed, it has been suggested that normal oocyte maturation occurs through both an inhibition and stimulation of selective signal transduction pathways (Smith, 1989). Thus qualitative as well as quantitative changes in growth factor receptors may be necessary to induce normal cell division in an otherwise inappropriate context. Because of the availability of a series of hEGF-R mutations with defined alterations in autophosphorylation, substrate specificity, ability to stimulate ion transport and receptor internalization (Chen et al., 1989), the oocyte is a good system in which to investigate these types of problems.

References

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Reconstitution of EGF Receptor System in Oocytes

1671

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