Regulation of Phosphorylation of the c-erbB-2/HER2 Gene Product by a Monoclonal Antibody and Serum Growth Factor(s) in Human Mammary Carcinoma Cells

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Monoclonal antibody (MAb) 4D5 was used to analyze the phosphorylation of $p185^{nEK2}$, the gene product of c-erbB-2/HER2, in SK-BR-3 cells. Culture in the continuous presence of 4D5 reduced the in vivo steady-state levels of p185HERZ phosphorylation by 80% in a dose-dependent manner, suggesting that MAb 4D5 may have interfered with the activation of phosphorylation of p185^{HER2}. The observed MAb-mediated reduction of p185" ^{encee} phosphorylation could not be completely accounted for by down-regulation. When cultures were grown under serum-free conditions, the steady-state levels of p185 new phosphorylation were reduced by 56%, and addition of 4D5 further inhibited phosphorylation to 20% of steady-state levels. With continuous exposure to increasing concentrations of newborn calf serum in these cultures, there was a linear increase in tyrosine-specific phosphorylation of p185^{HERZ}, reaching a 5.4-fold increase with 10% newborn calf serum. Phosphorylation of p185^{HER2} in the presence of newborn calf serum was not attributable to stimulation of the epidermal growth factor receptor by epidermal growth factor or by transforming growth factor- α . Extension strated a significant capacity of serum to induce p185^{HER2} phosphorylation. The demonstration of antibodyof these observations to two other mammary carcinoma cell lines, MDA-MB-453 and BT-474, also demonmediated partial inhibition of phosphorylation under serum-free conditions suggests that mammary carcinoma cells may also produce and secrete a factor or factors which may activate p185^{HER2}. Our observation that growtb-inhibitory MAb 4D5 is able to reduce the phosphorylation of p185^{72RZ} by newborn calf serum and by
a cellular-derived factor(s) suggests the existence of a growtb factor(s) which uses phosphorylation of p185^{HER2} as a signal transduction pathway to regulate cell proliferation.

Proto-oncogenes are a group of normal genes which play important roles in the regulation of cell proliferation and function (2, 5). Abnormalities in the expression, structure, or activity of proto-oncogene products contribute to the development and maintenance of the malignant phenotype in complex but important ways (36, 37, 46). Evidence that the gene products of several activated proto-oncogenes are either growth factors or growth factor receptors has suggested a possible link between proto-oncogenes and growth factors (20). For example, the receptor for macrophage colony-stimulating factor is identical to the product of c -fms (35), and c-erbB-1 encodes the receptor for epidermal growth factor (13) and transforming growth factor- α (TGF- α) (43). Growth factor receptors encoded by proto-oncogenes are transmembrane glycoproteins with intrinsic tyrosine-kinase activity (22). Receptor tyrosine kinases are activated by binding of their respective ligands, the growth factors (48). This activity is thought to be an integral part of signal transduction processes involved in the regulation of cell proliferation (21). Overexpression of some growth factor receptors has been shown to induce transformed properties in recipient cells (11, 32), possibly because of excessive activation of signal transduction mechanisms. Furthermore, a number of tumor cells with increased expression of growth factor receptors also produce ligands for these receptors (10).

HER2 (also known as c-erbB-2 or c-neu), the human

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homolog of the rat proto-oncogene neu (9), encodes a 185-kDa transmembrane glycoprotein with intrinsic tyrosine kinase activity which is presumed to be the receptor for an as-yet-unidentified ligand (3, 39). p185^{HER2} also has homology to, but is distinct from, the epidermal growth factor receptor (EGF-R), which is the product of c-erbB-1. Both proteins have a cysteine-rich extracellular domain, a transmembrane domain, and an intracellular tyrosine kinase (4, 31, 47). In spite of sequence homology between c-erbB-2 and c-erbB-1, EGF does not bind to $p185^{nE_{K2}}$ (33). $p185^{nE_{K2}}$ has been shown to be overexpressed or amplified or both in a number of human malignancies: breast (45), ovarian (38), thyroid (1), lung (7), salivary gland (34), and stomach (50). In addition, p185^{HER2} is a potent oncogene capable of inducing transformation and tumorigenesis when overexpressed in NIH 3T3 cells (12, 19). Overexpression of p185HER2 also induces tumor cell resistance to macrophage killing (15). Thus p185^{HER2} may have an important role in the development and maintenance of human tumors.

These observations suggest that receptor-associated tyrosine kinase activity of overexpressed proto-oncogene protein products is important for the regulation of cell growth. We have developed a panel of monoclonal antibodies (MAbs) reactive with domains of the human EGF-R (23) and $p185$ ^{HER2} (17, 18) in intact cells and have demonstrated antiproliferative effects of these antibodies in vitro (16, 18, 23) and in vivo (29). Antibody 4D5, which is specifically directed against p185^{HER2}, exhibits strong antiproliferative activity on cultured human breast tumor cell lines which overexpress $p185^{nE}$ (18). Since $p185^{nE}$ is a receptor

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with intrinsic tyrosine kinase activity, we investigated the modulation of p185^{HER2} phosphorylation by MAb 4D5. We report here that activation of phosphorylation of p185^{HER2} by serum was reduced in the presence of an excess of MAb 4D5 and that MAb-sensitive phosphorylation was mediated by a growth factor or factors other than TGF- α or EGF. Furthermore, SK-BR-3 cell-conditioned medium contained a factor(s) that could activate $p185^{HER2}$ phosphorylation and was partially inhibited by MAb 405.

MATERIALS AND METHODS

Materials. MAbs 405 (18) and 9G6 (44) were raised against human pl 85^{HER2} . MAbs 528 and 225 bind to the human EGF-R (23). Antiphosphotyrosine MAb PY-69 was obtained from ICN Biochemicals, Inc. Rabbit immunoglobulin to mouse immunoglobulins G (RAM) was supplied by Accurate Chemicals, Westbury, N.Y. 32P; (carrier free; 28.5 Ci/nmol) and $35S$ -labeled L-cysteine (1,030 Ci/mmol) were purchased from New England Nuclear, Boston, Mass.

Cell lines and cell culture. Human breast tumor cell lines SK-BR-3, BT-474, and MOA-MB-453 were obtained from the American Type Culture Collection. The A431 human epidermal carcinoma cell line was originally supplied by Gordon Sato. All cell lines except MOA-MB-453 (which was grown in L-15 medium) were maintained in Ham F-12- Dulbecco modified Eagle medium (1:1, vol/vol) (F-12/ OMEM) supplemented with 10% fetal bovine serum.

Labeling of p185 HER2 with ³²P₁ and [³⁵S]cysteine. Cells (3 \times $10⁵$) were plated in F-12/DMEM in each well of a six-well dish. Twenty-four hours later, cultures were washed with phosphate-free medium and incubated for up to 15 h in phosphate-free F-12/DMEM containing 0.4 mCi of $^{32}P_1$ per ml in the presence or absence of MAb and newborn calf serum. At desired times, cells were harvested in 400 μ l of lysis buffer (20 mM HEPES [N-2-hydroxyethylpiperazine- N' -2-ethanesulfonic acid; pH 7.5], 1% Triton X-100, 10% glycerol, 1.5 mM magnesium chloride, 1 mM ethyleneglycol bis- N, N, N', N' -tetraacetic acid, 0.1 mM phenylmethylsulfonyl fluoride, $10 \mu g$ of leupeptine per ml, 2 mM sodium orthovanadate) at 4°C for 20 min. The lysate was centrifuged at 10,000 rpm in an Eppendorf microfuge for 10 min, and then $60 \mu l$ of Pansorbin was added as described elsewhere (42). For labeling with $[35S]$ cysteine, the cells were washed with cysteine-free medium and refed with cysteine-free F-12/DMEM containing 0.15 mCi of [35S]cysteine per ml with or without 5% newborn calf serum.

lmmunoprecipitation and SDS-polyacrylamide gel electrophoresis. Aliquots $(350 \mu l)$ of the cell lysates (or equal amounts of trichloroacetic acid-precipitable counts per minute) containing $32P$ -labeled or $[35S]$ cysteine-labeled $p185^{HER2}$ were subjected to immunoprecipitation with 10 µg of MAb 9G6, 528, or PY-69 at 4°C for 2 h. Immune complexes were collected by absorption to RAM-protein A-Sepharose beads at 4°C for 1 h. Beads were washed three times with 1 ml of buffer (20 mM HEPES [pH 7.5], 150 mM NaCl, 0.1% Triton X-100, 10% glycerol, 2 mM sodium orthovanadate). Washed pellets were mixed with 40μ I of sample loading buffer (10 mM Tris HCI [pH 6.8], 1% sodium dodecyl sulfate [SOS], 0.2% 2-�-mercaptoethanol, 10% glycerol, 0.001% bromophenol blue), heated at 95°C for 5 min, and resolved on a 7% SOS-polyacrylamide slab gel (26). The efficiency of precipitating labeled receptor with MAb 9G6 is 80 to 90% when this procedure is used. Low-molecular-mass colored markers (Amersham Corp.) were used as standards. Phosphoamino acid analysis. The band corresponding to

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FIG. 1. Effect of MAb 4D5 on steady-state levels of p185HER2 phosphorylation in SK-BR-3 cells. Subconfluent cultures were labeled with ${}^{32}P_1$ (400 µ.Ci in 1 ml of phosphate free F-12/DMEM supplemented with 5% newborn calf serum) in the continuous presence of different amounts of antibody for 15 h. Detergent extracts were made, and p185HER2 was immunoprecipitated by using MAb 9G6 and then resolved by 7% SDS-polyacrylamide gel electrophoresis (Materials and Methods). An autoradiogram resulting from 16 h of exposure of the dried gel is shown here. The arrow indicates the position of ³²P-labeled pl85^{*HER*2}. Lane 1, Control cells; lanes 2 to 4, cells treated with MAb 405 at 30, 150, and 300 nM, respectively; lane 5. cells treated with 400 nM F(ab) fragment of MAb 4D5. The amounts (in counts per minute) of $p185^{HER^2}$ in each lane were 4,453 (lane 1), 1,967 (lane 2), 1.785 (lane 3), 1,040 (lane 4), and 335 (lane 5). Counts were corrected by subtracting the background of 60 cpm. The results shown are representative of results in six different experiments.

the 185-kOa HER2 protein, resolved as described above, was excised out of the gel. ^{32}P -labeled p185 HER2 in a gel slice was partially hydrolyzed with 200 μ l of 6 N HCl at 110°C for 1 h. Two portions (10 μ l each) of the hydrolysate were taken for measurement of radioactivity in a liquid scintillation counter to determine the total incorporation of $32P$ into the p185HER2 receptor. The rest of the hydrolysate was dried, suspended in distilled water, and applied to a Oowex AG1-X8 column. The column was washed with distilled water, and the absorbed $32P$ -labeled materials were eluted with 0.5 N HCI and lyophilized. The recovery of radioactivity by this procedure was 78 to 85%. 32P-phosphoamino acids mixed with unlabeled carrier phosphoamino acids (phosphoserine, phosphothreonine, and phosphotyrosine [1:1:1)) were analyzed by thin-layer electrophoresis as described elsewhere (8).

RESULTS

MAb 4D5 reduces amount of $32P$ -labeled p185 $HER2$. MAb 405 was used to investigate the regulation of phosphorylation. SK-BR-3 cells, which have an amplified c-erbB-2 gene (45), were cultured for 15 h in medium containing $^{32}P_i$ in the continuous presence of various concentrations of MAb 405. The p185^{HER2} from these cells was immunoprecipitated with another anti-p185 HEK2 MAb, 9G6, which recognizes a dis-</sup> tinct epitope of $p185^{HER2}$, and resolved by SDS-polyacrylamide gel electrophoresis. Results of such an experiment are shown in Fig. 1. Treatment of cells with 405 reduced in vivo steady-state levels of ^{32}P -labeled pl85^{HER2} up to 80% in a dose-dependent manner (lanes 2 through 4). There was 49% \pm 8% reduction in phosphorylation by 150 nM MAb 4D5 in eight different experiments. When the F(ab) fragment of 405 was used instead of intact antibody, comparable or greater reduction of ^{32}P -labeled p185 HER2 was observed (lane 5). As a control, SK-BR-3 cells were incubated with another MAb, 225 lgGI, specifically directed against the EGF-R. and there was no effect on the amount of $32P$ -labeled p185 $HER2$ (unpublished data). The reduction in steady-state levels of 32 P-labeled pl85^{HER2} was not due to interference by 4D5 with MAb 9G6 during the immunoprecipitation reaction, as immunoprecipitation performed with another polyclonal an-

FIG. 2. Specificity of the reduction of $32P$ -labeled p185 H^{ER2} by MAb in SK-BR-3 cells in the presence or absence of MAb 405. Subconfluent cells were labeled with ³²P_i for 15 h. The cells were lysed in 600 µI of extraction buffer and divided into two equal parts of 250 µI each. Immunoprecipitation was performed with anti-pl85 MAb (lanes l to 3) or with anti-EGF-R MAb 528 (lanes I' to 3'). An autoradiogram of a dried gel is shown here. Lane 1 and 1'. Control; lanes 2 and 2', 30 nM MAb 405; lanes 3 and 3', 150 nM MAb 405. Counts per minute: lane l, 5,985; lane 2, 3,798; lane 3, 3,120; lane l'. 853; lane 2', 779; lane 3'. 932. Abs, Antibodies.

tibody (18) recognizing the carboxy-terminal 17 amino acids of pl85^{HER2} gave similar results (unpublished data).

Next, we examined the possibility of general inhibitory effects of MAb 405 on the steady-state levels of other ³²P-labeled receptor proteins by analyzing the amount of ³²P-labeled p185^{HERZ} and ³²P-labeled EGF-R in the same experiment (Fig. 2). These results indicated that there was no reduction of 32P-labeled EGF-R during 15 h of treatment of SK-BR-3 cells with 150 nM MAb 405. which had reduced the amount of $32P$ -labeled pl85 HER by 48%.

Analysis of reduction of $p185^{MER2}$ phosphorylation. The reduction of steady-state levels of $32P$ -labeled p185 H^{ER2} by MAh 4D5, shown in Fig. 1 and 2. could result from downregulation of p185^{HER2} and/or interference in the activation of $p185^{HER2}$ phosphorylation by a direct or indirect mechanism(s). In initial studies to explore these possibilities, parallel cultures of cells were metabolically labeled with $[35S]$ cysteine or $32P_i$. During 11 h of concurrent incubation with MAb 4D5, there was a 45% reduction in ³²P-labeled p185^{HERZ} (Fig. 3A) and only a 14% reduction in ³⁵S-labeled 1255-120 p1365-120
p185^{HER2} (Fig. 3B). This suggests that the reduced ³²P label in p185^{HERZ} in the presence of MAb 4D5 can only partially be attributed to reduced p185^{HER2} content. Next, we performed a similar experiment comparing the capacities of the monovalent F(ab) fragment of MAb 405 and an intact MAb 4D5 to affect the reduction of ³⁵S-labeled p185^{HER2}. There was no change in 35 S-labeled pl85 $^{\prime\prime\prime\prime\prime\prime}$ in the presence of F(ab), but there was a 26% reduction caused by MAb 4D5 (Fig. 3C, lanes 3 and 2, respectively). The results obtained in the immunoprecipitation experiments documented in Fig. 3A through C were confirmed by immunoblotting (D) . Immunoblotting of the 32P-labeled SK-BR-3 cell extracts used in Fig. 2 demonstrated only a marginal reduction in the content of pl85^{11ER2} protein when cells were cultured in the presence of MAb 405 but a substantial reduction in the amount of $32P$ -labeled p185^{HER2} (Fig. 2). The expression of EGF-R was not affected. Immunoblotting of similar unlabeled SK-BR-3 extracts also demonstrated very little reduction in the content of $p185^{HER2}$ by MAb 4D5 (Fig. 3D, experiment 2). These findings indicate that increased receptor catabolism induced by a MAb cannot fully account for the observed reduction in $32P$ labeling and show [with $F(ab)$] that reduced labeling is dissociated from reduced content of $p185^{HER2}$.

Next we addressed the possibility that the reduction in $32P$ -labeled p185 HER2 associated with exposure to MAb 4D5 could be related to a change in the level of expression of pl85HER? on the plasma membrane or to the extent of

FIG. 3. Analysis of the reduction of $p185^{HER2}$ phosphorylation in SK-BR-3 cells treated with MAb 4D5. Cells were labeled with ³²P; (A) or $[³⁵S]$ cysteine (B) in the presence or absence of MAb 4D5 (150) nM) for 11 h. Samples were prepared and separated as described in Materials and Methods. The autoradiogram shown here was obtained by 6 h of exposure. (C) Cells were labeled with $[35S]$ cysteine for 11 h in the presence of MAb 405 (150 nM, lane 2) or F(ab) <400 nM. lane 3) or with culture medium (lane 1). Samples were prepared and immunoprecipitation was carried out as described in Materials and Methods. An autoradiogram of a dried gel is shown here. Quantitation of the $p185^{HER2}$ bands was obtained by densitometric scanning (A through C) or by determining radioactivity associated with bands (A and B). Quantitation by determining the radioactivity
associated with p185^{HER2} bands in panels A and B gave results similar to those with densitometric scanning, and there was a 27% \pm
3% additional reduction in ³²P-labeled p185^{H ER2} compared with
³⁵S-labeled p185^{HER2}. (D) Immunoblotting of p185^{HER2} and EGF-R proteins. In experiment 1 (Exp.1), ³²P-labeled SK-BR-3 cell extracts (50 μ g of protein) used in Fig. 2. lanes 1 and 2, were resolved on a 7% SDS-polyacrylamide gel and then immunoblotted with anti-P185 M:\b 9G6 or anti-EGF-R polyclonal antibody RK-11. Experiment 2 shows the immunoblotting of unlabeled SK-BR-3 cell extracts µrepared following culture for 15 h with or without 30 nM MAb 405. Since some of the extracts used here were radiolabeled, immunoblotted membranes were visualized by using a protein A-gold etmancement kit (30). Abs, Antibodies.

down-regulation of receptor protein. First, we determined what fraction of the 35 S-labeled p185 $11ER^2$ is present on the cell surface at 37°C (Fig. 4A). In these experiments.
p185^{*HER2*} expressed on the plasma membrane was identified by its capacity to bind MAb 405 prior to cell lysis. The results indicate that $19\% \pm 4\%$ (average from three different experiments) of total 35 S-labeled p185 $11ER^2$ is expressed on the cell surface under these experimental conditions; thus pl85^{HER2} is available for down-regulation by MAb 4D5. Down-regulation of EGF-R has been shown to be dependent on temperature (41). To confirm that down-regulation of surface pi ^{185</sub>HER2} also is reduced at 4° C, experiments were performed to analyze the effect of temperature on the abundance of ³⁵S-labeled p185^{*HER2*} on the cell surface. Results indicated that at $4^{\circ}C$ the amount of total ³⁵S-labeled p185 $^{\prime\prime\prime\prime\prime\prime\prime}$ expressed on the surface increased to 35% \pm 3% (data not shown) compared with $19\% \pm 4\%$ of total ³⁵Slabeled $p185^{\prime\prime E\bar{\kappa}2}$ at 37°C.

In order to define the contribution of down-regulation to MAb-induced reduction in p185^{11ER2} phosphorylation, we

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FIG. 4. (A) Quantitation of surface expression of ³⁵S-labeled p185HER₂. Cells were labeled with [³⁵S]cysteine for 11 h. At the end of incubation, some cultures were lysed in 500 µI of lysis buffer for the determination of total 35 S-labeled p185 meck by immunoprecipitation with 10 µg of 405 (Materials and Methods). For measuring the surface expression of 35 S-labeled pl85^{HER2}, cultures were washed with phosphate-buffered saline and further incubated with F-12/ DMEM-20 mM HEPES (pH 7.5) containing 20 µg of high-affinity MAb 405 per ml for 1 h at 4°C. The cultures were washed, lysed in 500 µI of extraction buffer, and processed for immunoprecipitation by adding RAM-protein A-Sepharose beads but no more MAb 405 during the immunoprecipitation procedure. The results of two representative experiments are shown here. Lanes 1 and 3, Total 35 S-labeled p185^{HER2}; lanes 2 and 4, 35 S-labeled p185HERZ on the cell surface. (B) Effect of MAb 4D5 on surface expression of $p185^{HER2}$ in a temperature shift experiment. SK-BR-3 cells were first equilibrated with 32P; for 4 h at 37°C (lane 1) and then further incubated with or without MAb 405 for an additional 11 h (in the continuous presence of 32P;) either at 37°C (lanes 2 and 3) or at 4°C (lanes 4 and 5). The autoradiogram shown here was obtained by exposing lanes 1 to 3 for 24 h and lanes 4 and 5 for 96 h. Quantitation of the amount of ³²P associated with pl85^{HERZ} bands was obtained by densitometric scanning of the autoradiogram and by determining the radioactivity associated with $p185^{HEN2}$ bands (A and B).

analyzed the effect of incubation with MAb 4D5 at 4°C. In these studies, cells were first equilibrated with $3^{2}P_1$ for 4 h at 37°C (Fig. 48, lane 1) and then maintained at 37°C (lanes 2 and 3) or shifted to 4°C (lanes 4 and 5) for an additional 11 h, with or without MAb 405. A comparison of the labeled material in Janes 2 and 4 in Fig. 48 (ftuorographs exposed for 24 and 96 h, respectively) showed a significant 85% reduction in ³²P labeling of $p185^{n2}$ during 11 h at 4^{n}C compared with labeling at 37°C. However, incubation of cells at 4°C did not prevent a further substantial MAb-mediated reduction in steady-state levels of $p185^{HER2}$ phosphorylation: there was a 34% decrease at 4°C (compare lanes 4 and 5) and a 51% decrease at 37°C (compare lanes 2 and 3). Taken together, these observations indicate that MAb-induced reduction in
p185^{HER} phosphorylation cannot be completely accounted for by down-regulation.

Experiments were performed to determine whether the F(ab) fragment might have the capacity to act as an agonist by activating tyrosine phosphorylation. The results in Fig. 5 indicate that the addition of F(ab) for 15 min slightly stimulated in vivo tyrosine phosphorylation of $p185$ ^{HER2} in cultures labeled with 32P (Fig. 5, lane 2'). However, there was no activation in cultures exposed to F(ab) for a longer treatment of 60 min (Fig. 5, lane 3 '). The observation that the F(ab) fragment of 4D5 does not down-regulate the $35S$ -

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FIG. 5. Partial agonist nature of F(ab). Subconfluent SK-BR-3 cells were labeled with ${}^{32}P_i$ for 15 h. Some cultures were treated with 400 nM F(ab) for the indicated times. The cells were lysed in 800 μ I of extraction buffer. The lysates were divided into two equal parts of 350 µI each and then immunoprecipitated with MAb 906 (lanes 1 to 3) or with antiphosphotyrosine MAb PY-69 (lanes 1' to 3'). An autoradiogram resulting from a 1-h exposure of dried gel is shown here. Lane 1, Control; lane 2, F(ab) incubation for 15 min; lane 3, F(ab) incubation for 60 min.

labeled $p185^{HER2}$ but can act for a short time as a partial agonist is interesting; however, we have not attempted to further characterize these properties in the present study.

Activation of phosphorylation of p185^{BER2} in presence or absence of newborn calf serum. Next, we investigated the possible source of the factor(s) that might stimulate
p185^{HER2} phosphorylation. As shown in Fig. 6A, culturing the cells in serum-free medium resulted in a steady-state level of phosphorylation of pl85^{HER2} reduced 56% (lane 1) compared with that observed in the continuous presence of newborn calf serum (lane 3). The addition of MAb 4D5 in serum-free culture conditions further reduced p185^{HER2}

FIG. 6. (A) Detection of newborn calf serum-mediated phosphorylation of p185^{HER2}. Subconfluent SK-BR-3 cells were labeled with ${}^{32}P_1$ in the culture medium without (lanes 1 and 2) or with (lanes 3 and 4) 5% newborn calf serum for 15 h. Cultures analyzed in lanes 2 and 4 also were continuously exposed to 150 nM MAb 405. Samples were prepared and immunoprecipitated for assaying the amount of pl85^{H ER2} as described in the Materials and Methods. Quantitation of the pl85 bands was obtained by densitometric scanning of the autoradiogram. (B) Control experiment showing effect of serum on the ${}^{32}P_1$ labeling of EGF-Rs for 15 h in SK-BR-3 cell cultures. Cell extracts were immunoprecipitated with anti-EGF-R MAb 528, which recognizes one distinct band with an approximate molecular mass of 170 kDa (arrow). (C) Two-dimensional thin-layer electrophoresis pattern of 32P-phosphoamino acids in a hydrolysate of the $p185^{nE}$ immunoprecipitated in panel A. S, Phosphoserine; T, phosphothreonine; Y, phosphotyrosine. Number at lower left of each autoradiogram indicates the following culture conditions: l, with no serum; 2, with serum; 3. with serum and MAb 405. Tyrosine phosphorylation in control cells was visualized faintly on the autoradiogram but reproduces poorly.

phosphorylation (Fig. 6A, lane 2) to 20% of the steady-state levels achieved in the absence of newborn calf serum (Fig. 6A, lane l). Experiments were done to examine the capacity of newborn calf serum to stimulate tyrosine phosphorylation of p185^{HER2} in short-term treatment. There was no increased activation of phosphorylation when serum-free cultures were supplemented with newborn calf serum for 30 min at 37 or 4°C (data not shown).

To determine the specificity of the capacity of newborn
 $\frac{16}{5}$ steps to stimulate astivation of $\frac{6}{5}$ also also also also calf serum to stimulate activation of $p185^{HER2}$ phosphorylation in SK-BR-3 cells, we investigated the potential for serum activation of another closely related molecule, the EGF-R. There was no potentiating effect of newborn calf serum on phosphorylation of the EGF-R in SK-BR-3 cells (Fig. 6B).

Having shown an increase in the steady-state levels of p185^{HER2} phosphorylation induced by newborn calf serum and its reduction by MAb 405, we determined the phosphoamino acid content of p185^{HER2} under these conditions by two-dimensional thin-layer electrophoresis (Fig. 6C). p185^{HER2} from cells cultured in the absence of newborn calf serum contained predominantly phosphoserine and phosphotlineonine with little phosphotyrosine (Fig. 6C, blot 1). The presence of some phosphorylation on tyrosine can be demonstrated by longer exposure of the autoradiogram but is not visualized well in the figure shown. Quantitation of the relative amount of label in each amino acid was obtained by scraping the ninhydrin-ideiltified spots from the thin-layer plate for liquid scintillation counting. Activation by 5% n�wborn calf serum increased the total phosphoamino acid content 2.2-fold, while for phosphotyrosine, the increase was 3.9-fold (Fig. 6C, blot 2). Inhibition of newborn calf serum-mediated stimulation of p185^{HER2} phosphorylation by MAb 405 resulted in a parallel reduction in the content of all three phosphoamino acids (Fig. 6C, blot 3).

Tyrosine phospborylation of p185^{HER2}. To further quantitate tlie relative increase in the steady-state phosphotyrosine content of p185^{HER2} induced by newborn calf serum, cells were metabolically labeled with [35S]cysteine and assayed for the steady-state phosphotyrosine content of pl85^{HER2} by using antiphosphotyrosine MAb PY-69. MAb PY-69 was specific for phosphotyrosine in the immunoprecipitation reaction; i.e., we were able to show competion for binding with cold phosphotyrosine and not with phosphoserine in experiments with 35S-labeled EGF-R (data not shown). As illustrated in Fig. 7A, lanes l' to 4', the amount of phosphorylation of p185^{HER2} on tyrosine increased with the concentration of newborn calf serum in the culture medium. The level of activation of tyrosine phosphorylation in serum-free medium was 18% of that observed in 10% newborn calf serum, and in medium containing 2.5% serum, tyrosine phosphorylation was 35% of that observed with a serum concentration of 10%. As a control, equal amounts of labeled cell extracts were immunoprecipitated with anti-p185HER2 MAb 9G6 (Fig. 7A, lanes 1 through 4). There was no significant effect of serum on the levels of ³⁵S-labeled p185HER2 in the cells. To quantitate these results, the ratios of phosphotyroslne-associated counts to total counts associated with $p185^{HER2}$ are presented in Fig. 7B, which shows a dose-dependent increase of up to 5.4-fold with 10% newborn calf serum in the culture medium.

Partial depletion of activating factor from newborn calf serum. Since our results indicated the presence of some activating factor(s) for $p185$ ^{HER2} phosphorylation in newborn calf serum, we sought confirmation of this observation by determining whether newborn calf serum could be de-

FIG. 7. (A) Detection of tyrosine-specific phosphorylation of pl85^{HER2} by newborn calf serum. Subconfluent SK-BR-3 cells were metabolically labeled with $[35S]$ cysteine in the presence of different concentrations of serum for 15 h. Cells were lysed in 650 μ l of extraction buffer. The lysates were divided into two equal parts of 300 µ each and then immunoprecipitated with anti-p185HER2 MAb 9G6 (lanes 1 to 4) or antiphosphotyrosine MAb PY-69 (lanes 1' to 4'). Other details of the assay were as described in the legend to Fig. 5. To detect the phosphotyrosine (P-Tyr) signal in cells cultured in the absence of serum (lane l'), it was necessary to expose the autoradiogram for 20 h, which resulted in overexposure of lanes l to 4. (8) To quantitate the data in panel A, the radioactivity associated with p185^{HER2} was determined by counting the excised bands in a liquid scintillation counter. The ratios of counts in p185HER2 phosphotyrosine over total counts in p185^{HER2} were plotted as a percentage of control with 0% newborn calf serum against the concentration of newborn calf serum used in the culture medium. (C) Depletion of the activator(s) of p185HER2 phosphorylation in serum. SK-BR-3 cells were labeled with ${}^{32}P_1$ in the absence or presence of serum for 15 h. Lane 1, Control without scrum; lane 2, medium with 5% newborn calf serum; lane 3, medium with 5% newborn calf serum depleted of factor(s) by three repetitive adsorptions of 4 h each on SK-BR-3 cells at 4"C; lane 4, control medium adsorbed on A431 cells. Cell extracts were prepared and pl85^{HER2} was assayed as described in Materials and Methods.

pleted of such a factor(s). In these experiments, phosphatefree medium containing 5% newborn calf serum was treated by repetitive absorption with SK-BR-3 cells (three times, for 4 h each time, at 4°C). Culture medium treated in an identical manner by adsorption with A431 cells, which do not express high levels of p185^{HER2}, was used as control. Figure 7C shows that there was a 53% reduction in p185HER2 phosphorylation in SK-BR-3 cells cultured in the presence of medium preadsorbed with SK-BR-3 cells (lane 3) compared with untreated medium (lane 2), and there was only a 16% reduction in p185^{HER2} phosphorylation when SK-BR-3 cell cultures were supplemented with medium preadsorbed with A431 cells (Fig. 7C, lane 4).

Activation factor(s) In newborn calf serum was not TGF-a or EGF. p185^{HER2} has been shown to be phosphorylated when EGF-Rs are activated by exposure to EGF or TGF- α , which are not ligands for p185^{HER2} (24, 40). To determine whether the newborn calf serum-mediated 2.2-fold enhancement of $p185^{HER2}$ phosphorylation resulted from the activity of TGF- α or EGF, we used anti-EGF-R MAb 528, which has

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Keep your litigation team up-to-date with **real-time alerts** and advanced team management tools built for the enterprise, all while greatly reducing PACER spend.

Our comprehensive service means we can handle Federal, State, and Administrative courts across the country.

Advanced Docket Research

With over 230 million records, Docket Alarm's cloud-native docket research platform finds what other services can't. Coverage includes Federal, State, plus PTAB, TTAB, ITC and NLRB decisions, all in one place.

Identify arguments that have been successful in the past with full text, pinpoint searching. Link to case law cited within any court document via Fastcase.

Analytics At Your Fingertips

Learn what happened the last time a particular judge, opposing counsel or company faced cases similar to yours.

Advanced out-of-the-box PTAB and TTAB analytics are always at your fingertips.

API

Docket Alarm offers a powerful API (application programming interface) to developers that want to integrate case filings into their apps.

LAW FIRMS

Build custom dashboards for your attorneys and clients with live data direct from the court.

Automate many repetitive legal tasks like conflict checks, document management, and marketing.

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Sync your system to PACER to automate legal marketing.

