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Epidermal Growth Factor Cytoplasmic Domain Affects ErbB Protein Degradation by the Lysosomal and Ubiquitin-Proteasome Pathway in Human Cancer Cells^{1,2}

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Abstract

The cytoplasmic domains of EGF-like ligands, including EGF cytoplasmic domain (EGFcyt), have important biological functions. Using specific constructs and peptides of human EGF cytoplasmic domain, we demonstrate that EGFcyt facilitates lysosomal and proteasomal protein degradation, and this coincided with growth inhibition of human thyroid and glioma carcinoma cells. EGFcyt and exon 22-23–encoded peptide (EGF22.23) enhanced procathepsin B (procathB) expression and procathB-mediated lysosomal degradation of EGFR/ErbB1 as determined by inhibitors for procathB and the lysosomal ATPase inhibitor BafA1. Presence of mbEGFctF, EGFcyt, EGF22.23, and exon 23–encoded peptides suppressed the expression of the deubiqitinating enzyme ubiquitin C-terminal hydrolase-L1 (UCH-L1). This coincided with hyperubiquitination of total cellular proteins and ErbB1/2 and reduced proteasome activity. Upon small interfering RNA–mediated silencing of endogenously expressed UCH-L1, a similar hyperubiquitiny-lation phenotype, reduced ErbB1/2 content, and attenuated growth was observed. The exon 23–encoded peptide region of EGFcyt was important for these biologic actions. Structural homology modeling of human EGFcyt showed that this molecular region formed an exposed surface loop. Peptides derived from this EGFcyt loop structure may aid in the design of novel peptide therapeutics aimed at inhibiting growth of cancer cells.

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Introduction

Members of the family of epidermal growth factor (EGF)–like ligands are synthesized as membrane-anchored proforms and are enzymatically cleaved to release extracellular and intracellular domains. The interaction of the extracellular part of EGF-like ligands with the membranebound EGF receptors, ErbB1-4, is well established and regulates normal growth, differentiation, and tumorigenesis [1]. The important roles of the cytoplasmic domain (cyt) of EGF-like ligands are emerging. The cyt domains of transforming growth factor α (TGF α -cyt) and amphiregulin (ARcyt) are important for cell polarity, intracellular vesicular trafficking [2–5], and basolateral sorting [6–8]. Soluble neuregulin-1 cyt acts as a Address all correspondence to: Dr Thomas Klonisch, Department of Human Anatomy and Cell Science, Faculty of Medicine, University of Manitoba, 130-745 William Ave, Winnipeg, Manitoba, Canada R3E 0J9. E-mail: klonisch@cc.umanitoba.ca

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nuclear transcriptional suppressor for several regulators of apoptosis [9,10], and its interaction with cytosolic LIM-kinase 1 is implicated in visual-spatial cognition [9,11]. Heparin binding-epidermal growth factor (HB-EGF)cyt can affect cell survival [12] and induces S-phase entry by interacting with the cyclin A transcriptional repressor promyelocytic leukemia zinc finger protein and its heterodimerization partner B-cell lymphoma 6 [13-15]. Nuclear localization of HB-EGFcyt is linked to aggressive transitional cell carcinoma [16]. Membraneanchored betacellulin (BTC) undergoes intramembrane processing by presenilin 1 and/or presenilin 2-dependent gamma secretase to generate a free BTCcyt fragment that becomes palmitoylated as a prerequisite for gamma secretase-dependent processing and nuclear membrane localization of BTCcyt [17]. Impaired basolateral sorting of pro-EGF and inadequate renal EGFR activation caused by the P1070L mutation in human proEGFcyt results in rare autosomal recessive renal hypomagnesemia syndrome [18]. Transgenic (tg) mice overexpressing human proEGFcyt showed stunted growth [19]. In human thyroid carcinoma cells, EGFcyt alters the acetylation state of the microtubules and inhibits tumor cell invasion into elastin matrix by a reduced expression of (pro)cathepsin-L and impaired SNAP25-mediated process of vesicle-membrane fusion [20,21].

Ligand-dependent ErbB receptor degradation involves the binding of exogenous EGF to the EGFR (ErbB1), which triggers membraneanchored surface receptor phosphorylation and initiates EGFR internalization/degradation along the endolysosomal pathway [22,23]. Lysosomal hydrolases of the cathepsin family proteolytically process EGFR [24], with procathepsin B (procathB) playing a major role in EGFR degradation [25,26]. ErbB degradation also involves the ubiquitinproteasome system (UPS). UPS malfunction can increase the levels of activated ErbB1/EGFR and ErbB2/Neu antigen, and this may contribute to carcinogenesis [27,28]. Ubiquitination, the conjugation of a highly conserved 76-amino-acid ubiquitin (Ub) molecule to a substrate, is an energy-dependent multistep process [29]. Monoubiquitinin signal will target ErbB receptors for endosomal degradation, whereas a polyubiquitination tag is believed to mark the receptors for proteasomal degradation [30,31]. Before proteasomal degradation, deubiquitination enzymes (DUBs), including ubiquitin C-terminal hydrolase UCH-L1 and Ub-specific processing proteases (UBPs), cleave and recycle free Ub from its ubiquitinated substrate [32]. UCH-L1 maintains intracellular levels of free ubiquitin UCH-L1, and this preserves normal proteasome function [33]. UCH-L1 is expressed in mature neurons and testicular stem cells and is present during early embryogenesis [34]. Overexpression of UCH-L1 is found in numerous cancers and is correlated with advanced stages of disease and poor prognosis for patient survival [35].

In the present study, we demonstrate two novel roles of EGFcyt as an inducer of lysosomal procathB expression and a silencer of UCH-L1 expression in human thyroid and glioma cancer cells. Both events resulted in a decrease in cellular ErbB1/EGFR and ErbB2/Neu receptor levels and reduced cancer cell growth. This function was critically dependent on the presence of the exon 23–encoded peptide. Molecular modeling showed that this exon 23–encoded domain formed a loop structure in human EGFcyt.

Materials and Methods

Cell Culture

Human follicular thyroid carcinoma cell line FTC-133 and the human glioma cell line LN-18 were cultured in HAM-F12 medium (Fisher, Ottawa, Ontario, Canada) and 10% fetal calf serum (Sigma, Oakville, Ontario, Canada). Stable transfectants of the follicular thyroid carcinoma cell line FTC-133 harboring pcDNA4/HisMaxC empty plasmid control or pcDNA4/HisMaxC containing constructs encoding for the complete human EGF cytoplasmic domain (exons 22-24; EGFcyt), EGFcyt encoding for exon 22 and 23 (EGF22.23), a natural splice version of EGFcyt lacking exon 23 (EGFdel23) [20], and the human EGF, juxtamembrane, transmembrane, and cytoplasmic domain with C-terminal FLAG epitope (mbEGFctF; gift from Dr S. Wiley) [36] (Figure 1*A*) were described previously [19–21]. Transient transfectants of the human glioma cell line LN-18 were generated by using effectin (Qiagen, Toronto, Ontario, Canada) and using 0.4 µg of EGF constructs and empty vector (mock control). Protein lysates were harvested 48 hours after transient transfection for Western blot analysis.

RNA Isolations and Reverse Transcription–Polymerase Chain Reaction

Total RNA of all transfectants was isolated with TRIzol reagent (Life Technologies, Burlington, Ontario, Canada), and RNA amounts were determined by spectrophotometry at 260 and 280 nm. Total RNA (1 μ g) was used for first-strand complementary DNA (cDNA) synthesis using the SuperScript reverse transcriptase kit and 500 ng/ml of oligo d(T) primer (both Life Technologies). For reverse transcription–polymerase chain reaction (RT-PCR), 1 μ l of cDNA and 25 pM of specific forward and reverse primer were used (Table 1). The PCR cycle profile consisted of an initial denaturation at 95°C for 15 minutes followed by 30 cycles of denaturation at 95°C for 1 minute, annealing at 60°C for 1 minute, and elongation at 72°C for 2 minutes. A negative control devoid of cDNA template was included. Single-PCR products were separated in a 1% agarose gel in TAE buffer (EMD, San Diego, CA), visualized by ethidium bromide staining, and sequenced to confirm identity.

Immunohistochemistry

Five-micrometer sections of paraformaldehyde-fixed and paraffinembedded human thyroid tissues were treated with 3% hydrogen peroxidase to block endogenous peroxidase activity. Nonspecific binding were blocked with 3% bovine serum albumin and 10% goat normal serum (Sigma). The sections were incubated with a rabbit polyclonal antiserum against UCH-L1 (1:100; Enzo Biochem, New York, NY) overnight at 4°C. Sections were washed 3 × 10 in Tris (pH 7.4) and 0.1% Tween 20 buffer and incubated with a secondary goat antirabbit antibody (1:200; New England Bio [NEB], Ipswich, MA) for 1 hour at room temperature. Staining was developed using the DAB kit (Pierce, Nepean, Ontario, Canada).

RNA Silencing

Mock and EGF23del transfectants at 60% confluence were transfected with 50 nM siRNA-UCHL-1 (AM51331, s105082, ID7345) or the nonsilencing scrambled sequence (5'AATTCTCCGAACGTGTCA-CGT3') as a control (both Ambion, Austin, TX) using Lipofectamine 2000 (Life Technologies). After 48 and 96 hours, *UCH-L1* gene expression and UCH-L1 protein levels were assessed by RT-PCR and Western blot analysis, respectively.

BrdU Proliferation Assay

Stable and transient transfectants were seeded at 10^3 cells/well into 96-well plates and cultured overnight in serum-free medium. Cells were preincubated with exogenous EGF (10 ng/ml) for 8 hours and with



Figure 1. (A) Schematic representation of human EGF constructs used in this study: cyt cytoplasmic domain; e22-24, exons 22 to 24; JR, juxtamembrane region; TM, transmembrane domain. (B) Western blot analysis revealed an exclusive up-regulation of procathB in stable human thyroid carcinoma transfectants expressing EGFcyt and EGF22.23. Deletion of the exon 23–encoded peptide region in EGFdel23 and membrane-bound mbEGFctF failed to show an induction of procathB expression. (C) Treatment with specific inhibitors for procathB/L (ZFF-FMK, 10 μ M), cathB (CA-74-ME, 10 μ M), and Iysosomal ATPases (Baf-A1, 1 μ M) resulted in enhanced EGFR protein levels in the presence of EGFcyt and EGF22.23 (not shown) but failed to alter procathB and EGFR protein levels in mock. (D) Western blot detection revealed an up-regulation of EGFR in the presence of ZFF-FMK (10 μ M) in EGFcyt but not in mbEGFctF and mock transfectants. Addition of exogenous EGF did not oppose the ZFF-FMK effect nor did EGF alter cellular EGFR protein levels. (E) Exclusive expression of procathB protein in EGFcyt thyroid transfectants. Addition of exogenous EGF did not oppose the ZFF-FMK effect nor did EGF alter procathB protein levels. (F) BrdU proliferation assays revealed increased proliferation of proEGFcyt and mock cells on EGF treatment (10 ng/ml). The inhibitors ZFF-FMK (10 μ M) and Baf-A1 (1 μ M) did not alter cell growth nor did they interfere with the EGF-induced proliferation.

Table 1. Primer Sequences Used for the Amplification of Target Genes Listed.

Target	Primer	Primer Sequence (from 5' to 3')	
UCHL-1	Forward	CCC CGA GAT GCT GAA CAA AGT G GGG	
	Reverse	CTG CCT GTA TGG CCT CA	
UCHL-3	Forward	GCT GGA GGC CAA TCC CGA GG	
	Reverse	GGT ATC TGG CTC GTT CTT CAG G	
EGFR	Forward	TGT GAG GTG GTC CTT GGG AAT TTG G	
	Reverse	TGC TGA CTA TGT CCC GCC ACT GGA	
EGFcyt	Forward	ACTCAGAAGCTGCTATCGAAAAACCC	
	Reverse	CTCACTGAGTCAGCTCCATTTGGTG	
ErbB2	Forward	GAA CCC CCA GCT CTG CTA CCA	
	Reverse	CGT AGA AAG GTA GTT GTA GGG AC	
GAPDH	Forward	CATCACCATCTTCTAGGACCG	
	Reverse	TGACCTTGCCCACAGCCTTG	

specific inhibitors. BrdU assays were performed according to the manufacturer's protocol (Roche, Mississauga, Ontario, Canada). Reactions were stopped using 1 M H_2SO_4 , and the absorbance was measured in an ELISA reader at 450 nm.

Inhibitors

For BrdU assays and Western blots, cells were incubated with proteasome inhibitors MG132 (10 μ M) or lactacystine (1 μ M) for 2 hours and with cath-B inhibitor ZFF-FMK (10 μ M), cath-B/-L inhibitor CA-74-ME (10 μ M), and the inhibitor of lysosomal ATPases BafA1 (1 μ M) for 3 hours (all Calbiochem, San Diego, CA) and ErBb1 inhibitor AG1478 (10 μ M) for 1 hour (Sigma-Aldrich, Oakville, Ontario, Canada). Batimastat (Sigma-Aldrich) was used as described earlier [36]. Cellular proteins were extracted with 1× Laemmli buffer (4% SDS, Tris-HCl pH 7.6, mercaptoethanol, bromophenol blue).

MTT Cell Viability Assay

Cells were seeded at 10³ cells/well into 96-well plates and cultured overnight in serum-free medium. Cells were incubated with specific inhibitors to determine their cytotoxicity effect on cells. MTT assays were performed according to the manufacturer's protocol (Life Technologies).

Transient Transfection with EGFcyt Peptides

Parental FTC-133 cells were transfected with N-terminally fluorescein isothiocyanate (FITC)–labeled 10- to 12-mer synthetic peptides (EZBioLab, Carmel, IN) derived from the exon 22– and exon 23– encoding peptide sequences of human EGFcyt using the Chariot kit according to manufacturer's instructions (Ambion; Table 2). Transfection efficiency was consistently more than 70% as determined by counting FITC-positive cells in a Z1 upright fluorescent microscope (Zeiss, Jena, Germany).

Subcellular Fractionation and Western Blot Analysis

Cells were seeded in 25-cm² flasks for 3 days under normal serum conditions. At 80% confluence, cell protein lysates were taken using a nuclear, cytoplasmic, and membrane extraction kit according to the company protocol (Pierce). For Western blot, transfectants were seeded at 8.5×10^4 cells/6 wells for 24 hours in serum-free medium with and without treatment of MG132 and lactacystine (both at 1 μ M). For Western blot analysis, 20- μ g/ml proteins were transferred onto a Hybond nitrocellulose membrane (Thermo Scientific, Nepean, Ontario, Canada) and nonspecific binding sites were saturated for 1 hour at room temperature in blocking buffer containing Tris-buffered saline/0.01% Tween 20 plus 5% skimmed milk. Membranes were incubated overnight at 4°C with rabbit polyclonal antibodies to UCHL-1 (1:1000;

Enzo Biochem), histone 1 (H1), ErbB1, ErbB1^{Tyr1045}, and ErB2 (all 1:1000; NEB), and a monoclonal antibody (mAb) to ubiquitin (1:2000; Enzo Biochem), respectively. All antibodies were diluted in blocking buffer. A mouse mAb against β -actin (1:20,000; Sigma) was used to determine equal protein loading per lane. On incubation with a horseradish peroxidase–conjugated goat anti-rabbit (1:2000; NEB) or goat antimouse secondary antibody (1:20,000; Sigma) for 2 hours at room temperature, specific immunoreactive bands were detected with an ECL kit (Thermo Scientific, Nepean, Ontario, Canada).

Immunoprecipitation

FTC-133-EGFcyt, FTC-133-mbEGFctF, and corresponding mock transfectants (1×10^6 cells) were harvested at 80% confluence, lysed in 400 µl of lysis buffer (50 mM HEPES [pH 7.4], 1% Triton X-100, 100 mM NaCl, 10 mM NaF, 1 mM EDTA, 1 mM sodium vanadate) for 1 hour on ice and centrifuged at 13,000g for 15 minutes at 4°C. Anti-EGFR antibody (10 µg/ml; Santa Cruz, Santa Cruz, CA) was added in lysis buffer and incubated overnight at 4°C. Immunocomplexes were purified by protein A/G agarose beads (30 µl/200 µl; Sigma) with gentle rocking for 4 hours at 4°C, pelleted for 60 seconds, and washed 3× in 500 µl of ice-cold lysis buffer. Pellets were resuspended twice in 100 µl of 3× SDS sample buffer (1.25 M Tris, pH 6.8, 10% SDS, 20% glycerol, 10% mercapthoethanol, 2% bromophenol blue), heated to 90°C for 5 minutes followed by Western blot detection of immunoprecipitated EGFR (1:1000, NEB). Membranes were stripped with buffer containing 0.2 M glycine and 0.05 Tween 20 for 15 minutes at room temperature, washed, blocked with 5% milk, and probed for c-Cbl (1:1000; Santa Cruz) and ubiquitin (Ub; 1:2000) (Cedarlane, Burlington, Ontario, Canada).

Proteasome Activity Assay

Transfectants expressing cytosolic EGFcyt, mbEGFctF, and mock cells were seeded in 96-well plates at 1×10^3 cells/well. Cells were pretreated with 10 μ M of MG132 for 1 hour and soluble EGF for 8 hours (10 ng/ml). Proteasomal activity assays were performed according to the manufacturer's protocol (Promega, Madison, WI).

Homology Modeling of the EGFcyt Domain

The homology modeling approach was used to generate 3D structures of EGFcyt composed of 149 amino acids [37]. The Modeller Web-based server was used in model building. On the basis of sequence identity values ranging between 35% and 38%, the structure of the receptor binding protein P2 of PRD1 (PDB code: 1n7v) was used as one of the templates. The strain energy of the model was generated

Table 2. Peptide Sequences Used in This Study.

EGFcyt Coding Region of Peptides	Peptide No.	Peptide Sequence (N'- to C'-terminus)
Exon 22	P1	TQKLLSKNPKNP
	P2	KNPYEESSRDVR
	P3	DVRSRRPADTED
	P4	TEDGMSSCPQP
Exon 23	P1	PQPWFVVIKE
	P2	VIKEHQDLKN
	P3	DLKNGGQPVA
	P4	QPVAGEDGQAAD
Scrambled peptide	P5	QPVIHDGVAAE

Peptides (10-12 mer) were designed to cover the complete exons 22 and 23 and have a three amino acid overlap at the N-terminus with each consecutive C-terminal peptide. All peptides were FITC conjugated at the N-terminus to monitor cellular uptake.



Figure 2. (A, C) In thyroid cancer cells, EGFcyt and EGF22.23 caused a 50% decrease in proliferation when compared to EGFdel23 and mock. Treatment with exogenous EGF (10 ng/ml), MG132 (10 μ M), and lactacystine (1 μ M) either completely or partially antagonized the growth inhibitory action of EGFcyt and EGF22.23, respectively. Dual treatment of cells with exogenous EGF plus MG132 or lactacystine (1 μ M) showed that the proteasome inhibitor did not antagonize the growth-promoting action of EGF. (B and D) MbEGFctF had no growth-inhibitory effect and treatments with EGF (10 ng/ml) MG132 (10 μ M) and lactacystine (1 μ M) failed to affect cell proliferation in mbEGFctF and mock cells.

separately in the GROMOS force field using the 53A6 parameter set optimized for molecular dynamic simulations. After building the 3D model of human EGFcyt, all atomic positions were locked, and required hydrogen atoms were added to the backbone structure of the molecule. Molecular dynamics simulation was performed at 1×10^6 for 50 picoseconds for further optimization in GROMACS (version 3.3). This involved a brief, steepest-descent run that used a maximum step protocol of 1.5 Å, and a maximum tolerance of 2000 kJ/mol per nanometer followed by a more extensive conjugate gradient minimization with tolerances of 200 kJ/mol per nanometer. After building and refinement of the 3D EGFcyt model, the Protein Structure & Model Assessment Tools were used to verify the quality of the model. This tool is capable of verifying a number of aspects of model qualities such as (i) local model quality estimation (anolea atomic mean force potential, em-

pirical force field, composite scoring function for model quality estimation), (ii) global model quality estimation (all atom distance–dependent statistical potential), (iii) stereochemistry check (protein structure verification, stereochemical quality check; minimum resolution, 2.5 Å), and (iv) structural features (secondary structure, geometrical features, and solvent exposure assignment, analysis of protein structure motifs). On the basis of these assessment results, model quality was evaluated according to the Ramachandran plot and the amino acid residues in the allowed, disallowed region and overall G factor.

Statistical Analysis

All experiments were repeated at least three times for two different clones. For multiple experiment comparisons, analysis of variance table and Tukey test were used with P < .05 considered significant.

Results

EGFcyt Affects Cathepsin-B–Mediated EGFR Degradation in Cancer Cells

Cathepsin-B (cathB) plays an important role in the lysosomal degradation of EGFR [26]. Here we demonstrate that of all EGF constructs tested (Figure 1A), soluble EGFcyt and EGF22.23 upregulated procathB in thyroid cancer (Figure 1B) and glioma cells (Figure W1A). This effect was not observed with membraneanchored mbEGFctF, the EGFdel23 splice variant with a deletion of exon 23, or mock cells (Figures 1B and W1A), suggesting cellular compartment-specific EGFcyt effects and a functional role for the EGFcyt exon 23 peptide region. Both immunoreactive cathB and CD63 were colocalized in mock and EGFcyt clones, indicating lysosomal localization of procathB (data not shown). The presence of EGFcyt coincided with diminished EGFR and ErbB2 expression in stably transfected thyroid cancer cells (Figures 1 C and 6, A and B) and transiently transfected LN-18 glioma cells (Figure W1B). To determine an involvement of lysosomal procathB in this EGFR downregulation, EGFcyt thyroid transfectants and mock were incubated

> 12,000 10,000 8,000

> > 6,000 4,000 2,000 0

> > > mock

RLU/s

A.

with specific inhibitors for cathB (ZFF-FMK, 10 μ M), cathL/B (CA-74-ME, 10 μ M), or the inhibitor of lysosomal ATPases BafA1 (1 μ M). Treatment with these three inhibitors resulted in a marked up-regulation of EGFR in the EGFcyt transfectants but not in mock (Figure 1*C*). ZFF-FMK also caused the up-regulation of EGFR in EGFcyt glioma transfectants (Figure W1*C*). However, in mbEGFctF and mock transfectants cellular levels of EGFR (Figures 1*D* and W1*C*) and procathB (Figures 1*E* and W1*D*) remained unchanged. The EGFR present in the thyroid (Figure W4, *B-D*) and EGFcyt glioma (Figures W1*E* and W3) transfectants was bioactive in response to exogenous EGF and caused a maximal proliferative response similar to mock, whereas BafA1 and ZFF-FMK alone had no effect on proliferation (Figures 1*F* and W1*E*).

EGFcyt Affects Proteasome Activity in Cancer Cells

mbEGFctF

We determined the ability of structurally distinct EGFcyt domains to affect the proliferation of human thyroid and brain cancer cells. Cancer cells expressing soluble EGFcyt and EGF22.23 showed significantly reduced BrdU incorporation when compared with mock and/or EGFdel23 (Figure 2*A*) and mbEGFctF (Figures 2*B* and



cyt

mock

also emphasized the importance of the exon 23-encoded peptide region in this process because EGFdel23 and mock failed to affect proteasome activity.



Figure 4. (A) *UCH-L1* gene activity and (B) UCH-L1 protein were exclusively downregulated in the presence of EGFcyt, EGF22.23, and mbEGFctF. (C) Specific 10- to 12-mer peptides covering the complete EGFcyt peptide region encoded by exons 22 and 23 were used to determine the specificity of exon-derived peptides for the silencing of UCH-L1 by EGFcyt. Exclusively peptides encoded by the exon 23– derived peptide sequences of EGFcyt, but not scrambled peptides or peptides derived from pEGFcyt exon 22, were capable of downregulating UCH-L1 expression at the gene and protein levels. Representative immunodetection of UCH-L1 in (D) papillary and (E) undifferentiated human thyroid cancer tissues. (F and G) No specific immunostaining was detected in normal human thyroid tissues (data not shown) and when the primary antiserum was replaced by a species-specific nonimmune serum at the same concentration. Magnification, ×20.

W2A) transfectants. This reduction in cancer cell growth was not the result of increased cell death as determined by cytotoxicity assays (data not shown). EGF-induced EGFR activation antagonized the growth inhibitory effect of EGFcyt in thyroid cancer (Figures 2, A and C, and W4, A and B) and glioma (Figure W3) cells. Treatment of mock, EGFdel23, and mbEGFctF clones with soluble EGF caused a small increase in cell proliferation (Figures 2, A-D, W3, and W4, A and B). We determined if cleavage of the extracellular EGF part of mbEGFctF may account for the increased proliferation of mbEGFctF clones compared with EGFcyt transfectants. BrdU proliferation assays in the presence of the matrix metalloproteinase (MMP) inhibitor batimastat failed to alter the proliferation of mbEGFctF in thyroid transfectants, indicating that the increased proliferation observed in mbEGFctF clones was not a result of liberation of EGF from the mbEGFctF protein with subsequent EGFR activation (Figure W4A). However, batimastat was able to decrease the proliferation of LN-18, suggesting EGFR activation as a result of proteolytic release of endogenous EGF-like ligand(s) in these glioma cells. We detected the expression of endogenous proEGF by PCR in LN-18 cells (data not shown). The decrease in proliferation with batimastat was significant in EGFcyt and mbEGFctF LN-18 transfectant but not in the mock controls (Figure W3).

The proteasome inhibitors MG132 and lactacystine increased EGFR levels in thyroid cancer cells (Figure 6*B*) and glioma (Figure W2*B*) and enhance growth rates of EGFcyt and EGF22.23 transfectants (Figure 2, *A* and *C*). We detected a significant reduction in proteasomal activity in the presence of EGFcyt and EGF22.23 (Figure 3, *A* and *B*) and, to a lesser extent, in mbEGFctF (Figure 3, *A* and *C*). Proteasomal activity assays in the presence of increasing concentrations of MG132 confirmed the inhibitory effect of EGFcyt, EGF22.23 (Figure 3*B*) and mbEGFctF (Figure 3*C*) and indicated similar inhibition kinetics. Almost complete proteasomal inhibition was observed at 1 μ M MG132 in all transfectants (Figure 3, *B* and *C*). Treatment with soluble EGF and/or the EGFR inhibitor AG1478 failed to increase proteasomal activity in EGFcyt clones (Figure 3*A*). Cytotoxicity assays done in the presence of MG132, lactacystine, and EGF confirmed that the





Figure 5. (A) Representative Western blots showing the level of ubiquitinated proteins in EGFcyt and mock cells. In the presence of EGFcyt and mbEGFctF, the level of ubiquitinated total protein and protein fractions derived from subcellular compartments was increased. This enhanced protein ubiquitination was not observed with EGFdel23 (not shown) and mock. Ponceau S–stained blots served as loading control. EGFR served as marker for the membrane fraction, β -actin for the total and cytoplasmic fraction and histone 1 (H1) as marker for the nuclear fraction. (B) Transfection with a specific siUCH-L1 (50 nM) construct for 72 and 96 hours successfully silenced UCH-L1 expression in UCH-L1⁺ mock cells. (C) Western blot analysis showed that silencing of UCH-L1 protein coincided with increased levels of total ubiquitinated (Ub–) proteins in siUCH-L1 treated mock compared with UCH-L1⁺ mock and UCH-L1⁻ EGFcyt cells treated with scrambled siRNA.



Figure 6. (A) Although RT-PCR did not show a difference in ErbB1 and ErbB2 gene expression (data not shown), Western blot analysis revealed a decrease of ErbB1 and ErbB2 protein in the presence of EGFcyt and EGF22.23. No change in ErbB status was detected in the presence of mbEGFctF, EGFdel23, and mock. (B) Treatment with the proteasome inhibitors MG132 and lactacystine increased the amount of ErbB1 and ErbB2 protein in EGFcyt and mbEGFctF transfectants as shown in representative Western blots. Proteasome inhibitors had no effect on ErbB1/2 protein levels in EGFdel23 (data not shown) and mock transfectants. Equal protein loading was assessed by detecting β -actin.

reduction of proteasomal activity in the EGFcyt clones was not the result of increased cell death (data not shown).

EGFcyt Is a Novel Regulator of UCH-L1 in Cancer Cells

Microarray analysis identified and subsequent RT-PCR and Western blot data identified ubiquitin C-terminal hydrolase (UCH-L1) as a target gene in thyroid cancer cells. Thyroid cancer transfectants expressing EGFcyt and EGF22.23 showed the silencing of UCH-L1 at the mRNA (Figure 4A) and protein levels (Figure 4B). In the presence of mbEGFctF, UCH-L1 gene expression was attenuated at the mRNA (Figure 4A) and protein levels (Figure 4B). Treatment with soluble EGF for up to 4 hours did not change UCH-L1 protein levels in the EGFcyt and mbEGFctF clones (Figure 4B). LN-18 glioma cells showed strong UCH-L1 expression, but transient transfection with EGFcyt or mbEGFctF caused markedly reduced UCHL-1 protein levels (Figure W2C). UCH-L1 down-regulation by EGFcyt, EGF22.23, and mbEGfctF was specific for UCH-L1 and not observed for UCH-L3 in glioma (data not shown) and thyroid cancer cells (Figure 4A). UCH-L1 is a member of the DUB protein family and plays an important role in the processing of Ub as part of protein degradation.

To identify particular peptide regions in the EGFcyt domain responsible for this down-regulation of UCH-L1, we used FITC-labeled 10- to 12-mer peptides that covered the entire exon 22 and exon 23 region of human EGFcyt (Table 2). Thyroid cells were transfected with the different FITC-peptide conjugates, and transfection efficiency was determined in each case to be at 60% to 70% (counted fraction of FITC-stained to unstained cells). All peptides derived from the exon 23 peptide region caused a complete loss of UCH-L1 protein, whereas all exon 22–encoded peptides and the scrambled control peptide failed to affect UCH-L1 expression (Figure 4C). Thus, the exon 23–encoded peptide region of the human EGF cytoplasmic domain, as a membrane-bound



Figure 7. (A) UCH-L1 siRNA treatment (50 nM) successfully silenced endogenous UCH-L1 expression in mock. MG132 (10 μ M) treatment did not influence silencing of UCH-L1 in mock cells. (B) On UCH-L1 knockdown, Western blot analysis showed reduced ErbB1 and ErbB2 protein levels. However, in mock treated simultaneously with MG132 (10 μ M) and siUCH-L1 (50 nM), this receptor down-regulation was abolished, suggesting that proteasome activity was essential for the degradation of ubiquitinated ErbB1/2. β -Actin was used to assess equal protein loading. (C) Silencing of UCH-L1 gene activity in mock resulted in an approximately 50% reduction in proliferation, which was similar to the growth rates determined for UCH-L1–negative EGFcyt transfectants.



Figure 8. (A) Western blot detection of total fractions of EGFR and c-Cbl plus/minus treatment with EGF before IP; β -actin was used to assess equal protein loading. IP experiments and subsequent Western blot analysis confirmed the attenuated EGFR levels in the presence of EGFcyt. Unstimulated EGFcyt cells revealed enhanced levels of ubiquitinated EGFR similar to mock treated with EGF. In the presence of EGFcyt, coimmunoprecipitation of c-Cbl with EGFR revealed reduced c-Cbl–EGFR interaction compared with mock, which increased markedly on EGF treatment. An IgG1a isotype control antibody was used as a negative control in the IP experiments. (B) Western blot detection of total fractions of EGFR and c-Cbl plus/minus treatment with EGF before IP; β -actin was used to assess equal protein loading. IP results for EGFR, c-Cbl and ubiquitinated EGFR in mbEGFctF and mock cells plus/minus EGF treatment showed comparable results. Treatment with exogenous EGF caused phosphorylation of EGFR at residue Tyr¹⁰⁴⁵ (see Figure W4, *C* and *D*) and enhanced the interaction between c-Cbl and activated EGFR in all transfectants studied. An IgG1a isotype control antibody was used as a negative control antibody was used as a negative control antibody was used as a negative control in the IP experiments. (C) A structural model of human EGFcyt was generated based on the structure of the receptor binding protein P2 of PRD1 (PDB code: 1n7v) used as template. This molecule has up to 38% amino acids identity with human EGFcyt. The exon 23–encoded peptide region marked in red was found to form an exposed loop.

or soluble form, was identified to have an essential role in suppressing UCH-L1 expression in cancer cells.

The expression of UCH-L1 in human thyroid tissues has not been reported. We performed immunohistochemistry and demonstrated, for the first time, the presence of immunoreactive UCH-L1 in human papillary (Figure 4D) and undifferentiated (Figure 4E) thyroid carcinoma tissues. Normal human thyroid tissue (not shown) and control sections with primary antiserum replaced by nonimmune serum were

devoid of UCH-L1 (Figure 4, F and G). Our results suggest the exclusive expression of UCH-L1 in thyroid cancer tissues.

EGFcyt Decreases UCH-L1, Causes Hyperubiquitination of Cellular Proteins, and Reduces ErbB1/2 Receptor Levels

Cells with impaired UCH-L1-mediated DUB function are known to show high levels of protein ubiquitination [38]. Western blot analysis of total, cytoplasmic, nuclear, and membrane-bound protein

fractions showed increased levels of ubiquitinated cellular proteins in the presence of either EGFcyt or mbEGFctF (Figure 5A). RT-PCR and Western blot analysis confirmed the successful specific knockdown of UCH-L1 for more than 96 hours in UCH-L1-positive EGFdel23 and mock clones (Figure 5B). Silencing of UCH-L1 resulted in significantly elevated levels of ubiquitinated proteins at 72 and 96 hours in mock transfectants (Figure 5C). Next, we addressed if a reduced level of UCH-L1 can affect the level of ErbB receptors, particularly ErbB1/ EGFR and ErbB2/Neu antigen, and whether this may contribute to the attenuated cell growth in the presence of EGFcyt. We failed to detect changes in ErbB1 gene expression in any of the transfectants investigated (data not shown) but observed a significant decrease in ErbB1 and ErbB2 protein content exclusively in the presence of EGFcyt (and EGF22.23) in thyroid cancer cells (Figure 6, A and B) and glioma transfectants (Figure W1B). No change in ErbB1/2 protein content was detectable in mbEGFctF, EGFdel23, or mock cells (Figures 6, A and B, and W1B). Treatment with proteasome inhibitors MG132 and lactacystine resulted in increased ErbB1/2 protein levels exclusively in proEGFcyt and mbEGFctF transfectants (Figures 6B and W2B). Upon si-UCH-L1 treatment, we detected decreased levels of UCH-L1 mRNA (data not shown), UCH-L1 protein (Figure 7, A and B) and ErbB1/2 protein (Figure 7B). Treatment with MG132 failed to affect UCH-L1 levels in siUCH-L1-treated cells (Figure 7A) but abolished the siUCH-L1-induced decrease in ErbB1/2 (Figure 7B), resulting in protein levels similar to untreated and si-control-treated mock (Figure 7, A and B). The siUCH-L1-treated mock cells with a decrease in ErbB1/2 levels showed significantly reduced cell proliferation (Figure 7C). We performed immunoprecipitation (IP) assays with a specific antibody to EGFR on mock and UCH-L1-negative EGFcyt thyroid transfectants plus/minus EGF treatment to determine whether the presence of EGFcyt coincided with the increased level of ubiquitinated EGFR. Irrespective of EGF treatment, the IP revealed attenuated total EGFR protein levels in EGFcyt clones compared with mock (Figure 8A). Ubiquitinated EGFR was clearly observed in untreated EGFcyt transfectants but only weakly detectable in mock (Figure 8A). In mock, but not in EGFcyt transfectants, EGF treatment resulted in an increased presence of ubiquitinated EGFR (Figure 8A). IP results for mbEGFcyt and mock transfectants show similar EGFR levels, and in both cases, we observed an increase in ubiquitinated EGFR on EGF treatment (Figure 8B). When the anti-EGFR was replaced by a nonimmune IgG1a control serum, no immunoprecipitated proteins were detected (Figure 8, A and B). In all transfectants investigated, EGF treatment caused a marked increase in the phosphorylation of EGFR tyrosine residue 1045 (EGFR-Tyr¹⁰⁴⁵) (Figure W4C), which was inhibited in the presence of the EGFR inhibitor AG1478 (Figure W4D). We performed co-IP experiments for c-Cbl, known to ubiquitinate [22,39,40] EGFR-Tyr¹⁰⁴⁵ [41] and assist in the sorting process of the EGFR [42]. EGF treatment resulted in significantly increased c-Cbl coimmunoprecipitation with EGFR in mock, EGFcyt (Figure 8A), and mbEGFctF clones (Figure 8B).

Rendered Structural Modeling Reveals an Exposed Exon 23–Encoded Peptide Loop in EGFcyt

The molecular structure of human EGFcyt has not been reported. We used homology modeling based on the receptor binding protein P2 of PRD1 (PDB code: 1n7v), a modeling template displaying up to 38% amino acid identity with human EGFcyt. The rendered model of the cytoplasmic domain of human EGF revealed the exon 23–encoded peptide region we identified as a biologically important molecular module of EGFcyt to form an exposed loop structure potentially capable of participating in protein interactions involving EGFcyt (Figure 8C).

Discussion

In the present study, we identified soluble EGFcyt as a novel factor affecting lysosomal (pro)cathB-mediated degradation and proteasomal activity in cancer cells. We had previously demonstrated the presence of soluble human EGFcyt in the cytoplasm of cancer cells [19,20]. The increased procathB expression and down-regulation in EGFR protein were dependent on the exon 23-encoded peptide region present in EGFcyt and EGF22.23, confirming our previous findings of the important role of this peptide region as an essential molecular determinant for the in vitro and in vivo biologic activity of the EGF cytoplasmic domain [19-21]. CathB has been recognized as an important and specific degradation mechanism for EGFR in the endolysosomal compartment [26]. EGF ligand-mediated activation results in monoubiquitination at multiple sites within the tyrosine activation domain of EGFR [30,43], which serves as a signal for receptor internalization [30,44,45] and regulates sorting of endosomal EGFR to lysosomes [46,47]. Continuous EGFR ubiquitination by the E3 ubiquitin ligase c-Cbl is required for lysosomal targeting and degradation of the EGFR [42,48]. The specific inhibition of cathB and the elevation of the acidic pH in the endosomallysosomal compartment by the vacuolar-type ATPase (V-ATPase) inhibitor bafilomycin A1 (BafA1), known to disrupt the vesicular proton gradient and raise the pH of acidic vesicles [49], effectively blocked EGFR degradation in the presence of EGFcyt in cancer cells.

The presence of soluble and membrane-anchored EGFcyt and EGF22.23, all containing the exon 23-encoded peptide region, was also associated with reduced proteasome activity which could be further suppressed with increasing concentrations of MG132, a nonselective proteasome inhibitor known to also inhibit procathB [50]. To discriminate between these two inhibitory effects, we replaced MG132 with the selective proteasome inhibitor lactacystine [51] but obtained similar results. We identified the deubiquitin esterase UCH-L1, a major cellular ubiquitin hydrolase [33], as a novel transcriptional target that was specifically downregulated or completely silenced in the presence of mbEGFctF and EGFcyt, respectively. Intriguingly, the silencing of UCH-L1 was also achieved using small peptides designed according to the exon 23-encoded peptide sequence of EGFcyt. UCH-L1 is involved in diverse aspects of tumor cell adhesion and motility [52,53], but its role in tumor cell growth is still unclear and may depend on the specific cell context [54-57]. Uch-L1 transgenic mice recently provided conclusive evidence for a role of UCH-L1 as an oncogene. These mice displayed a high incidence of sporadic tumors, primarily consisting of lymphomas and lung adenoma [52]. In humans, prostate cancer is among numerous cancers shown to express UCH-L1 [58]. Here we provide evidence for the presence of UCH-L1 in human thyroid carcinoma. UCH-L1 was absent in normal thyroid tissues, suggesting that UCH-L1 may qualify as a novel marker for thyroid cancer cells.

Proper control of cellular ubiquitin levels is crucial for normal cell functions and requires the balanced actions of ubiquitin ligases and DUB enzymes [47,59–61]. Malfunction of the ubiquitin pathway due to UCH-L1 depletion in the presence of proEGFcyt, proEGF22.23, and exon23 peptides or UCH-L1 decrease in mbEGFctF clones coincided with increased ubiquitination of total, cytoplasmic, membrane, and nuclear protein fractions, including EGFR. This likely triggered a depletion of the cellular pool of free ubiquitin that caused the decrease

in proteasome activity observed in thyroid cancer cells [33,62,63]. Furthermore, the UCH-L1^{negative}, procathB^{high} EGFcyt phenotype resulted in lysosomal degradation of ErbB1/2 receptors as determined by our cathB/lysosomal inhibitor studies and significantly reduced cell growth in our cancer cell models. Importantly and contrary to human fibroblasts from patients with eight different lysosomal storage diseases, the attenuated UCH-L1 expression in our cancer cell models was not associated with increased cell death [62].

EGFR-activating EGF-like ligands can antagonize the function of EGFcyt [21]. We used the general MMP inhibitor batimastat to account for the possibility of a proteolytic release of (a) the bioactive extracellular EGF part of the membrane-anchored mbEGFctF that may stimulate EGFR (Figure 1A) [36] and/or (b) the MMP-mediated cleavage of endogenous membrane-bound EGF-like ligands. Such effects would explain differences observed between the EGFcyt and mbEGFctF transfectants because a liberated EGF/endogenous EGF-like ligand is expected to counteract the actions of the EGFcyt domain. Although batimastat had no effect in thyroid transfectants, it caused a small but significant decrease in proliferation with the glioma EGFcyt and mbEGFctF clones. The fact that we detected endogenous EGF expression in LN-18 may explain that both EGFcyt and mbEGFctF clones showed similar responses to batimastat and made proteolytic release of the EGF domain of the mbEGFctF construct unlikely. In addition, the batimastat experiments clearly showed that the release of endogenous EGF-like ligand had no significant effect on the responses measured, suggesting that the cellular effects observed for EGFcyt and mbEGFctF in thyroid and glioma cancer cells were genuine and, in the case of LN-18, only marginally affected by endogenously induced EGFR activity. Further support for a lack of enzymatic processing of mbEGFctF or its involvement in juxtacrine activation of EGFR on neighboring cells [36] came from the finding that mbEGFctF and mock clones showed similar levels of c-Cbl-EGFR interaction. The E3 ubiquitin ligase c-Cbl is known to ubiquitinate EGFR [22,39,40] and assist in the sorting process of this tyrosine kinase receptor [42]. As expected, the proto-oncogene and RING finger member c-Cbl coimmunoprecipitated with ubiquitinylated EGFR [42,64] and EGF-mediated EGFR phosphorylation caused the enhanced c-Cbl interactions with activated EGFR, suggesting that neither membrane-anchored nor soluble EGFcyt interfered with the ligandinduced c-Cbl-EGFR protein interaction.

This study demonstrated for the first time that EGFcyt can downregulate ErbB2/Neu. The ErbB2 receptor is commonly considered an endocytosis-impaired receptor and ligand-activated EGFR-ErbB2 dimers largely recycle back to the cell membrane for reactivation [65–67]. Despite its known association with the two E3 ubiquitin ligases CHIP and c-Cbl [67–70], the impaired ability of ErbB2 to undergo ubiquitination has been suggested as a potential cause [71]. Recently, the deubiquitinating activity of POH1, a component of the 19S proteasomal lid, was shown to have a critical role in the regulation of ubiquitinated ErbB2 levels [72]. We identified a novel role of UCH-L1 in facilitating the EGFcyt-mediated down-regulation of ErbB2, implicating UCH-L1 as an important novel DUB in EGFcyt-mediated regulation of both ErbB1 and ErbB2 [73].

Our data provide further evidence for a potential tumor suppressor function of the small EGFcyt peptide region encoded by exon 23 in cancer cells [19–21], and this involves UCH-L1. Our structural homology modeling predicted this exon 23–encoded peptide region to be an exposed loop structure accessible for protein-protein interactions. The combination of existing ErbB inhibitors and the UCH- L1 gene silencing effect of specific exon 23–encoded peptides may be a novel attractive therapeutic strategy for the treatment of patients with aggressive ErbB1/2–dependent cancers.

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Figure W1. (A) Western blot analysis revealed the up-regulation of procathB in human glioma cell line LN-18 transiently transfected with EGFcyt. This effect was not observed with mbEGFctF construct. (B) Decreased EGFR and ErbB2 protein levels in LN-18 transiently transfected with EGFcyt. This effect was not observed with mbEGFctF construct. (C) Western blot analysis revealed an up-regulation of EGFR in the presence of ZFF-FMK (10 μ M) in EGFcyt glioma transfectants. No change was observed in mbEGFctF transfectants and mock. (D) The same ZFF-FMK treatment did not alter cellular procathB expression in any of the LN-18 clones. (E) BrdU proliferation assays revealed decreased proliferation in LN-18 EGFcyt transfectants and treatment with the general MMP inhibitor batimastat caused a significant decrease in proliferation in EGFcyt and mbEGFctF clones suggestive of a possible weak EGFR activation through cleaved EGF-like ligand(s). In all cases, EGF treatment (10 ng/ml) caused a significant increase in proliferation in LN-18 EGFcyt clones. The procathB inhibitor ZFF-FMK (10 μ M) did not affect cell growth in the presence or absence of exogenous EGF.



Figure W2. (A) BrdU proliferation assays revealed a 12% increase in proliferation in LN-18 EGFcyt transfectants on MG132 treatment, which was enhanced further in the presence of EGF. The general MMP inhibitor batimastat caused a small but significant decrease in proliferation in LN-18 EGFcyt and mbEGFctF clones suggestive of a possible weak EGFR activation through cleaved EGF-like ligand(s). (B) Western blot analysis revealed an exclusive up-regulation of EGFR in the presence of MG132 (10 μ M) in LN-18 EGFcyt transfectants. No change was observed with mbEGFctF transfectants and mock. (C) Representative Western blot demonstrating that both EGFcyt and mbEGFctF glioma transfectants showed reduced levels of cellular UCH-L1. β -Actin served as a loading control.



Figure W3. BrdU proliferation assays showing the increased proliferation in EGFcyt glioma transfectants and mock on EGF treatment (10 ng/ml). The EGF-induced proliferation was blocked by the EGFR inhibitor AG1478 (10 μ M). AG1478 alone or in combination with the MMP inhibitor batimastat failed to significantly affect glioma cell growth.



Figure W4. (A) Treatment of EGF-responsive EGFcyt and mbEGFctF thyroid transfectants with the general MMP inhibitor batimastat and subsequent BrdU assays. The increased proliferation of the mbEGFctF compared with EGFcyt transfectants was not the result of EGFR activation due to the proteolytic release of the extracellular bioactive EGF domain of the mbEGFctF construct. Batimastat failed to cause a change in proliferation in any of the transfectants tested. (B) The EGF-induced (10 ng/ml) increase in proliferation of EGFcyt, mbEGFctF, and corresponding mock thyroid transfectants was blocked by the EGFR inhibitor AG1478 (10 μ M) as determined by BrdU proliferation assays. AG1478 itself did not affect cell growth. EGF treatment caused EGFcyt transfectants to significantly increase their proliferation rate to reach growth rates similar to mock. (C) Western blot detection of total EGFR and EGFR phosphorylated at residue Tyr¹⁰⁴⁵ (EGFR-Tyr¹⁰⁴⁵) in thyroid EGFcyt, mbEGFctF, and mock transfectants. EGF (10 ng/ml) caused the specific Tyr1045 phosphorylation of EGFR in all transfectants investigated. (D) EGFR phosphorylation at residue Tyr¹⁰⁴⁵ was completely blocked in the presence of the specific EGFR inhibitor AG1478.