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Impaired basolateral sorting of pro-EGF causes isolated recessive renal hypomagnesemia

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Primary hypomagnesemia constitutes a rare heterogeneous group of disorders characterized by renal or intestinal magnesium (Mg2+) wasting resulting in generally shared symptoms of Mg2+ depletion, such as tetany and generalized convulsions, and often including associated disturbances in calcium excretion. However, most of the genes involved in the physiology of Mg2+ handling are unknown. Through the discovery of a mutation in the EGF gene in isolated autosomal recessive renal hypomagnesemia, we have, for what we believe is the first time, identified a magnesiotropic hormone crucial for total body Mg2+ balance. The mutation leads to impaired basolateral sorting of pro-EGF. As a consequence, the renal EGFR is inadequately stimulated, resulting in insufficient activation of the epithelial Mg2+ channel TRPM6 (transient receptor potential cation channel, subfamily M, member 6) and thereby Mg2+ loss. Furthermore, we show that colorectal cancer patients treated with cetuximab, an antagonist of the EGFR, develop hypomagnesemia, emphasizing the significance of EGF in maintaining Mg2+ balance.

Introduction
Magnesium (Mg2+) is established as a central electrolyte in a large number of cellular metabolic reactions, including DNA synthesis, neurotransmission, and hormone receptor binding. It is a component of GTPase and a cofactor for Na,K-ATPase, adenylyl cyclase, phosphoinositide kinases, and phosphofructokinase (1). Mg2+ is also important for the regulation of parathyroid hormone release (2, 3). Accordingly, Mg2+ deficiency (plasma Mg2+ concentrations below 0.70 mM) has an effect on multiple body functions. Symptoms of Mg2+ deficiency are mostly related to muscle dysfunction, such as tetany, prolonged QT interval, and cardiac arrhythmias (4). Children with hypomagnesemia often present with tetany and/or convulsions. Hypomagnesemia is a problem frequently observed in more than 10% of hospitalized patients and occurrences can be as high as 65% in intensive care patients (5). A long-term complication seen in many adult patients with chronic hypomagnesemia is chondrocalcinosis, which can lead to impairment of joint function (4). Mg2+ deficiency can be secondary to systemic diseases (for instance, diabetes mellitus and Crohn disease) or to the use of osmotic agents, diuretics, and drugs such as cyclosporin and cisplatin (6). In addition, primary Mg2+ deficiency is observed in several monogenetic disorders. Failure of early diagnosis or noncompliance with treatment can be fatal or result in permanent neurological damage.

The plasma Mg2+ concentration is regulated within a narrow range by changes in urinary Mg2+ excretion in response to altered uptake by the intestine. Thus, the kidney plays a key role in Mg2+ homeostasis (4, 7). Most renal reabsorption of Mg2+ occurs in the proximal tubule and the thick ascending limb of the loop of Henle via a passive paracellular transport process, but the fine-tuning of the Mg2+ excretion takes place in the distal convoluted tubule (DCT), where Mg2+ is reabsorbed via an active transcellular transport process (6, 7). Apical entry into DCT cells is mediated by the Mg2+-permeable channel TRPM6 (transient receptor potential cation channel, subfamily M, member 6) driven by a favorable transmembrane voltage (8). The mechanism of basolateral Mg2+ transport into the interstitium is unknown. Mg2+ has to be extruded against an unfavorable electrochemical gradient, most likely by a Na+/Mg2+-dependent exchange mechanism and/or a Mg2+-ATPase. Finally, 3%–5% of the filtered Mg2+ is excreted in the urine.

Despite the critical role in Mg2+ handling, the exact mechanisms of transepithelial Mg2+ transport remain obscure. Studies of disorders with primary hypomagnesemia are very important to gaining more insight into the molecular and cellular mechanisms that underlie Mg2+ (re)absorption. Genetic studies in families with hereditary renal Mg2+ wasting syndromes have identified several genes that are either directly or indirectly involved in active Mg2+ handling. In the past few years, genetic studies of inborn errors of the Mg2+ balance revealed several new proteins unexpectedly involved in transepithelial Mg2+ transport in the DCT, e.g., thiazide-sensitive sodium chloride cotransporter (NCC), the γ subunit of the Na+,K-ATPase, and the previously mentioned epithelial Mg2+ channel, TRPM6 (9–12).

The aim of the present study was, therefore, to elucidate the gene defect and molecular mechanism underlying isolated recess-
sive renal hypomagnesemia (IRH), which is characterized by renal Mg\(^{2+}\) loss. To this end, a homozygosity-based mapping strategy and mutation detection was performed. In addition, the molecular mechanism of IRH was studied in vitro using patch clamp analysis and in vivo using clinical studies in humans.

**Results and Discussion**

IRH is characterized by low serum Mg\(^{2+}\) levels and mental retardation. Two affected sisters, V3 and V4 (Figure 1A), displayed low serum Mg\(^{2+}\) levels (0.53–0.66 mM) in combination with urinary fractional excretion (FE) values of Mg\(^{2+}\) of 4.8% and 3.6%, respectively. These values are well above an FE of 2%, indicating renal Mg\(^{2+}\) wasting as previously described (5, 13). Thus, the fact that the urinary excretion of Mg\(^{2+}\) was in the normal range while serum Mg\(^{2+}\) values were hypomagnesemic points to a diminished tubular Mg\(^{2+}\) reabsorption. Apart from hypomagnesemia, other biochemical abnormalities were not identified; in particular, there was no disturbance in urinary Ca\(^{2+}\) excretion (14). Both sisters showed psychomotor retardation during childhood, and presently they are moderately mentally retarded women who suffer from epileptic seizures. The previously identified genes involved in renal Mg\(^{2+}\) handling encoding the thiazide-sensitive sodium chloride cotransporter NCC (SLC12A3), paracellin-1 (CLDN16), the γ subunit of the NaK-ATPase (FXYD2), and the epithelial Mg\(^{2+}\) channel (TRPM6) have been excluded for these 2 affected sisters (8, 11, 12, 15). To determine the disease locus of this consanguineous family, a homozygosity-based mapping strategy with a set of highly polymorphic microsatellite markers spread over the genome was performed and followed by fine mapping. This resulted in the identification of a critical linkage interval with LOD score 2.66.

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**Figure 1**

Molecular analysis of EGF. (A) Pedigree of the family in which specific members suffer from IRH. Filled symbols represent affected individuals, half-filled symbols are heterozygous for the mutation C3209T, and open symbols represent clinically unaffected family members not screened for the mutation. Slash marks indicate deceased individuals, and double lines show consanguinity. (B) Schematic representation of the critical interval between polymorphic markers D4S2623 and D4S1575 on chromosome 4q and the intron-exon structure of the EGF gene, depicting the identified mutation. The position of the polymorphic markers is indicated by vertical bars. The horizontal arrow below the schematic boxed representation of EGF depicted between the polymorphic markers indicates the localization and orientation of the EGF gene. Cen, centromeric; tel, telomeric. (C) Genomic mutation sequence analysis of EGF in wild-type, homozygous, and affected individuals. The black bars under the sequences of the WT and heterozygous specify exon 22, resulting in the amino acid substitution P1070L. (D) Schematic model of pro-EGF, which resides as a type I membrane protein at the plasma membrane. The position of the P1070L mutation is depicted. (E) Sequence homology analysis of juxtamembrane domain and flanking residues. P1070 is strongly conserved among different species and forms the second proline of a basolateral PX domain and flanking residues.
(P < 0.004) on chromosome 4 of 18.4 cM between markers D4S2623 and D4S1575 (Figure 1B). A search for candidate genes within this region revealed the EGF gene, which was considered a highly relevant positional candidate since it had previously been linked to electrolyte homeostasis (16).

We sequenced EGF in the affected individuals (Figure 1A) and identified the homozygous mutation C3209T in exon 22, which cosegregated with the disorder and was absent in 126 ethnically matched control chromosomes (Figure 1C). The mutation caused the substitution of the highly conserved proline 1070 within the cytoplasmic tail of pro-EGF by leucine (P1070L) (Figure 1, D and E). We found that both parents and the 2 unaffected sisters (Figure 1A) exhibited the homozygous wild-type genotype (Figure 1, A and C).

The EGF gene consists of 24 exons encoding a large type I membrane–anchored precursor protein of 1,207 amino acid residues that exists as a membrane-bound molecule, which is proteolytically cleaved into pro-EGF to finally generate the 53–amino acid peptide hormone EGF (17). EGF has a profound effect on cell differentiation in vivo and is a potent mitogenic factor for a variety of cultured cells of both ectodermal and mesodermal origin (18). EGF belongs to the EGF-like family of growth factors, which bind with high affinity to the EGFR. Other members of this family are TGF-α, amphiregulin, heparin-binding EGF-like growth factor, betacellulin, and epiregulin (19–21). These membrane-anchored growth factor precursors are characterized by the fact that they are biologically active at the cell surface although they can be proteolytically cleaved to release soluble, diffusible factors (19–21).

To study the tissue distribution of pro-EGF and EGFR, we performed RT-PCR on various human tissues (Figure 2A). For EGF, PCR amplification products were detected in, e.g., kidney, salivary gland, cerebrum, and prostate, whereas no expression was detected in the adrenal gland, liver, and placenta. The EGFR showed a ubiquitous expression pattern since all tissues tested were positive by RT-PCR analysis (Figure 2A). Immunohistochemical analysis on rat kidney sections showed that EGF is consistently coexpressed with NCC, a marker of the DCT (22) (Figure 2B). Interestingly, the immunopositive staining is mainly localized along the apical domain in DCT. Coffey and coworkers demonstrated that human pro-EGF overexpressed in Madin-Darby canine kidney (MDCK) cells is delivered equally to the apical and basolateral membrane but is found predominantly at the apical membrane domain (23). They proposed that preferential ectodomain cleavage at the basolateral surface explains the apparent apical localization of pro-EGF. Sack and Talor provided evidence for the existence of specific binding sites in tubular basolateral membranes, suggesting a physiologic role of EGF in the kidney (24). In the kidney, the EGFR was detected in glomerular endothelial cells, peritubular capillaries, and arteriolar walls as well as along the thick ascending limb of Henle loop and DCT (25, 26). Immunohistochemical analysis revealed that the EGFR was located particularly along the basolateral membrane of the tubular cells. The DCT reabsorbs approximately 10% of the filtered Mg²⁺ load, and the reabsorption rate in this segment defines the final urinary Mg²⁺ excretion, as virtually no reabsorption takes place beyond this section (1). In kidney, the epithelial Mg²⁺ channel TRPM6 is specifically expressed in DCT, where it forms the rate-limiting step of epithelial Mg²⁺ transport (8). Thus, pro-EGF and TRPM6 are both predominantly

![Figure 2](image_url)

Expression profiling of EGF and EGFR.

(A) The expression of EGF and EGFR mRNA was determined by RT-PCR on various human tissues. Signals for EGF were detected in, e.g., kidney, salivary gland, prostate, and cerebrum whereas no signals were detected in the adrenal gland, cerebellum, liver, lung, and placenta. The EGFR showed a ubiquitous expression pattern since PCR amplification products were obtained in all tissues tested. hEGF, human EGF; hEGFR, human EGFR.

(B) Immunohistochemical analysis of EGF (green) and thiazide-sensitive sodium chloride cotransporter (NCC, red) in rat kidney sections (upper panel, overview of a cortical region; lower panel, magnified image of an immunopositive tubule). EGF colocalized with NCC, a marker for the DCT. Original magnification, ×180 (B, top panels); ×360 (B, bottom panels).
expressed in DCT, the main site of active renal Mg\(^{2+}\) reabsorption (1, 16, 17, 27). To determine whether EGF can regulate the activity of TRPM6, human embryonic kidney 293 (HEK293) cells were transiently transfected with TRPM6 and treated for 30 minutes with EGF (10 nM), resulting in increased channel activity (Figure 3A). Western blot analysis demonstrated endogenous expression of EGFR in these HEK293 cells (data not shown). EGF dose dependently stimulated TRPM6 activity with half maximal effective concentration of 1.7 nM, an apparent affinity that is in the physiological range as reported previously (Figure 3B) (24).

Previous in vivo studies showed that the predominant form of EGF released from epithelial cells is the high-molecular mass 160–170 kDa EGF, as found at high concentrations in serum, urine, and milk (28, 29). To investigate the functional effect of the identified mutation, HEK293 cells were transfected with wild-type or mutant pro-EGF and after 2 days their supernatant was collected. TRPM6-expressing HEK293 cells were subsequently incubated for 30 minutes with these preconditioned media and analyzed for channel activity using the patch clamp technique. Interestingly, the supernatant obtained from mutant pro-EGF–expressing HEK293 cells failed to stimulate TRPM6 activity whereas a significant channel activation was observed with the wild-type EGF–containing supernatant (Figure 3C). Quantification of EGF in both supernatants using ELISA indicated that the secretion of mutant pro-EGF is impaired (Figure 3D).

To demonstrate that the observed stimulatory effects are due to EGFR activation, a preincubation with cetuximab was performed. This IgG human/mouse chimeric monoclonal antibody competes with the EGF receptor (EGFR) to block binding of the ligand and subsequent signal transduction mediated via the receptor-associated tyrosine kinase and prevents phosphorylation of the EGFR and other downstream proteins.

**Figure 3**

Electrophysiological analysis of pro-EGF and pro-EGF–P1070L on TRPM6 channel activity. (A) Average time course of outward (at +80 mV) current densities from HEK293 cells transfected with TRPM6 in control conditions (open circles) and after EGF treatment (30 minutes, 10 nM, filled circles) in comparison with mock-transfected cells without EGF treatment (triangles) and after EGF stimulation (filled triangles). n = 17–20. (B) Dose-response curve of EGF-induced current in TRPM6-transfected HEK293 cells indicating half maximal effective concentration of 1.7 nM. n = 11–14. (C) Histogram presenting the current densities at +80 mV of TRPM6-transfected cells (200 seconds after establishment of the whole-cell configuration) treated with mock (mock), pro-EGF (EGF WT) or pro-EGF–P1070L (EGF-P1070L) supernatant. All treatments were performed for 30 minutes at 37°C. White bars indicate experimental conditions in which HEK293 cells were transfected with mock DNA whereas black bars indicate HEK293 cells transfected with TRPM6. Asterisk indicates significance in comparison with nontreated TRPM6-transfected cells. P = 0.024; n = 6. (D) Supernatants of mock-, pro-EGF–, or pro-EGF–P1070L–transfected HEK293 cells were analyzed by ELISA. Asterisk indicates mock and pro-EGF–P1070L supernatant were significantly different from EGF supernatant. P = 0.006; n = 4. (E) Histogram summarizing the current density (pA/pF) at +80 mV (200 seconds after break-in) of HEK293 cells treated with the apical or basolateral supernatant of MDCK cells stably transfected with either wild-type pro-EGF, pro-EGF–P1070L, or mock DNA (30 minutes, 37°C). White bars indicate experimental conditions in which HEK293 cells were transfected with mock DNA whereas black bars indicate HEK293 cells transfected with TRPM6. Crosses indicate significance in comparison with the mock treatment at the apical side (wild-type pro-EGF apical, P = 0.0001, n = 6; pro-EGF–P1070L apical, P = 0.029, n = 6), and pound symbol represents significance in comparison with the mock treatment at the basolateral side (wild-type pro-EGF basolateral P = 0.031, n = 6). (F) Schematic model illustrating how pro-EGF–P1070L mutation leads to impaired basolateral sorting of pro-EGF, resulting in abrogated stimulation of the EGFR localized at the basolateral membrane. Activation of the EGFR in DCT by EGF is necessary to prevent renal Mg\(^{2+}\) wasting by stimulation of the epithelial Mg\(^{2+}\) channel TRPM6.
proteins in the signal transduction cascade (30). The stimulatory effect of the supernatant collected from pro-EGF–expressing HEK293 cells on TRPM6 was significantly inhibited by preincubation for 1 hour with cetuximab (191 ± 22 pA/pF, 491 ± 62 pA/pF, and 286 ± 48 pA/pF for TRPM6-expressing HEK293 cells measured at +80 mV and treated with mock supernatant, EGF supernatant, or EGF supernatant plus cetuximab, respectively; P < 0.05, n = 6). Thus, the stimulatory effect of the supernatant on TRPM6 activity is due to direct activation of the EGFR.

As outlined above, the localization of EGFR is restricted to the basolateral membrane whereas pro-EGF is present at both the apical and basolateral membranes (23). This prompted us to investigate in detail the cellular metabolism of pro-EGF. Recently, several basolateral sorting motifs were identified, including the sequence motif PXXP (with X being an arbitrary amino acid), determining the targeting of proteins to the basolateral membrane in MDCK cells (31). Our mutation substitutes the second highly conserved proline in the cytoplasmic 1067PKNP1070 motif into a leucine (Figure 1D). We hypothesized that this mutation results in improper trafficking of pro-EGF, which prevents an adequate secretion of the hormone into the circulation. Since EGFR is only localized basolaterally, the mutation should prevent the ligand from reaching its receptor. Interestingly, He and coworkers have shown previously that a 22–amino acid sequence in the EGFR juxtamembrane domain contains autonomous sorting information necessary for basolateral localization of the receptor in MDCK cells (31). This sorting motif contains a polyproline core comprising residues P667 and P670 (667PXXP670). Site-directed mutations at P667 and P670 were associated with impaired basolateral delivery of the EGFR.

To determine the effect of the mutation on the sorting of pro-EGF in polarized epithelial cells, MDCK cells stably expressing either wild-type or mutant pro-EGF were grown on semipermeable filter supports until they reached confluency. After 2 days, media of the basolateral and apical compartment were collected individually. Next, HEK293 cells expressing TRPM6 were treated with the collected preconditioned media of wild-type pro-EGF, mutant pro-EGF, or mock-expressing cells. Equal stimulation of TRPM6 channel activity was observed when the cells were incubated with the apically collected media of wild-type or mutant pro-EGF–expressing MDCK cells (Figure 3E).

However, stimulation of TRPM6 channel activity was only observed with basolaterally collected wild-type pro-EGF medium, not with the basolateral mutant pro-EGF media (Figure 3E). Of note, the apically and basolaterally collected mock medium did not affect TRPM6 activity. These observations suggest that the basolateral release of mutant pro-EGF is diminished, seriously hampering the EGF-dependent activation of the basolaterally localized EGFR (Figure 3F).

Clinical trials directed to the treatment of patients with colorectal cancer have demonstrated that cetuximab is synergistic with chemotherapy for these patients (32). We observed that serum Mg²⁺ levels gradually decreased in all studied colorectal cancer patients on cetuximab treatment (Figure 4A). This is in line with a recent report showing that a significant number of similarly treated patients developed hypomagnesemia (30). Our patients treated with cetuximab displayed an FE of Mg²⁺ of 5.1%, which is inappropriately high given their hypomagnesemia (Figure 4B). As previously established by Agus (5), an FE of Mg²⁺ above 2% in the presence of hypomagnesemia indicates renal Mg²⁺ loss.

The 2 affected sisters with IRH exhibited a similar relatively high Mg²⁺ excretion profile (Figure 4B). The urinary Mg²⁺ patterns of the patients treated with EGFR antibodies, the patients described in our Dutch family, and patients with hypomagnesemia with secondary hyperparathyroidism (HSP; OMIM 230200) were identical and suggest mutual defects in renal TRPM6 activity. Indeed, preincubation with cetuximab abolished the stimulatory effect of EGF on TRPM6 activity as shown by patch clamp analysis of TRPM6-expressing HEK293 cells (Figure 4C). Taken together, these data strongly suggest a pivotal role for EGFR signaling in the maintenance of normal Mg²⁺ balance and indicate TRPM6 as a critical link between EGFR inhibition and IRH.
Mutation analysis of the EGF gene was performed by forward sequencing of the coding regions of the gene of the youngest affected family member. Forward and reverse sequencing of exon 22 was subsequently performed on all family members.

**Subjects with colorectal cancer treatment.** We prospectively studied consecutive patients with metastatic colorectal cancer for which treatment with cetuximab was initiated at the University Hospital Gasthuisberg (Leuven, Belgium). Most patients were included in phase II or III trials. Cetuximab was kindly donated by Merck KGaA. Informed consent was obtained from all subjects investigated in this study.

**Calculation of FE<sub>Na</sub>.** The FE<sub>Na</sub> was calculated using the following formula:

\[
FE_{Na} = \frac{U_{na} \times P_{cr}}{(0.7 \times P_{creat}) \times U_{Cr}} \times 100
\]

where \(U_{na}\) and \(P_{cr}\) refer to the urine and serum concentrations of Mg<sup>2+</sup> (Mg) and creatinine (Cr). Serum Mg<sup>2+</sup> concentration was multiplied by 0.7 since only approximately 70% of the circulating Mg<sup>2+</sup> was unbound by albumin and therefore able to be filtered across the glomerulus.

**Expression profiling.** PCRs were performed in 50 µl reaction volumes containing 1 µl of cDNA, 10 pmol of each primer (Supplemental Table 2), 2.5 mM MgCl<sub>2</sub>, 200 µM of each deoxyribonucleotide triphosphate (dATP, dTTP, dCTP, and dGTP), and 0.5 U of Taq polymerase. PCRs had an initial denaturation stage of 1 minute at 95°C, followed by 39 cycles of 30 seconds at 95°C, 1 minute at 55°C, 1 minute at 72°C, and a final extension step at 72°C for 10 minutes.

**Immunohistochemistry.** Staining of rat kidney sections for NCC and EGF was performed on 7-µm sections of paraffin-embedded kidneys. Sections were stained with affinity-purified anti-NCC (1:200) and affinity-purified rabbit anti-EGF (1:100) (Calbiochem; EMD Biosciences). Images were made with a Bio-Rad MRC 1000 laser scanning confocal imaging system using a x60 oil-immersion objective.

**Linkage analysis.** Samples of peripheral EDTA blood were collected from each person, and genomic DNA was isolated using standard procedures. Homozygosity mapping was performed with a genome-wide set of 400 evenly distributed microsatellite markers (LMS-MD10 2.5; Applied Biosystems), with an average intermarker distance of 10 cM, on the basis of Marshfield genetic maps (http://research.marshfieldclinic.org/genetics/GeneticResearch/compMaps.asp). The average heterozygosity of these markers was 0.76. Five additional markers, D4S1570, D4S2623, D4S2392, D4S1615, and D4S422, were selected for fine mapping purposes from the public databases on chromosome 4 (http://research.marshfieldclinic.org/genetics/GeneticResearch/data/Maps/Map4.txt and http://genome.ucsc.edu/cgi-bin/hgGateway). The analysis of the markers was according to the protocol provided for the LMS-MD10 version 2.5 with some small modifications. PCRs were performed in 10 µl reaction volumes containing 40 ng of genomic DNA, 5 pmol of each primer, 2.5 mM MgCl<sub>2</sub>, 250 µM dATP, dTTP, dCTP, and dGTP, each 10 mM Tris/HCl (pH 8.3), 50 mM KCl, and 0.4 U of Taq polymerase (AmpliTaq TM Gold; Applied Biosystems). PCRs had an initial denaturation stage of 12 minutes at 95°C, followed by 10 cycles of 15 seconds at 94°C, 15 seconds at 55°C, and 30 seconds at 72°C, then 22 cycles of 15 seconds at 89°C, 15 seconds at 55°C, and 30 seconds at 72°C, and a final extension step at 72°C for 10 minutes. Reactions for each marker were performed separately, with products being pooled into size-specific sets before typing. Markers were typed on an ABI 3100 sequencer (Applied Biosystems) using GeneMapper 4.0 software (Applied Biosystems). Allele binning was performed with the Excel 2000 (Microsoft) macro linkage designer developed by van Camp and coworkers (37), and...
we checked Mendelian inheritance of alleles with PedCheck 1.0 software (http://watson.hgen.pitt.edu) (38). Multipoint LOD scores were calculated with the GeneHunter, version 2.1 release 5, program in the easyLINKAGE software package (39).

Constructs. The full-length EGF precursor in vector plasmid SAD (pSAD) was kindly provided by G.I. Bell (17) (University of Chicago, Chicago, Illinois, USA). The full-length EGF precursor P1070L mutant was obtained by in vitro mutagenesis of the pSAD-EGF vector according to the protocol described by the manufacturer (Invitrogen). The construct was verified by DNA sequence analysis. To generate plasmid CB7 constructs for the stable expression of the full-length EGF precursor and the P1070L variant in MDCK type I cells (41) were grown in DMEM containing 5%.

Two days after

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The bicistronic expression vector pCINeo-IRES-GFP with EGF or preconditioned medium. Subsequently, cells were analyzed by DMEM (modified as described above) was performed prior to treatment conditioned medium. Cetuximab pretreatment for 1 hour with 2

TRPM6 was measured after a 30-minute preincubation with 10 nM mouse of EGF on TRPM6 activity in HEK293 cells transiently transfected with according to the manufacturer’s protocol (Stratagene). Primers used were 5′-CGGGGTACCGCCACATTGCCACTCTATATC-3′ and 5′-CCGCTCTAGAGTCACTGAGTCAGCTCCATTTG-3′. PCR products were cloned into the plasmid CB7 vector using the restriction enzymes KpnI and XbaI and verified by DNA sequence analysis.

Electrophysiology. The bicistronic expression vector pCINeo-IRES-GFP containing the full-length open reading frames of N terminally HA-tagged human TRPm6 (GenBank accession number NM_017662) was used to coexpress human TRPm6 and enhanced GFP in HEK293 cells. HEK293 cells were grown in DMEM (BioWhittaker) containing 10% (v/v) fetal calf serum, 2 mM l-glutamine, and 10 μg/ml Ciproxin at 37°C in a humidity-controlled incubator with 5% (v/v) CO2. Cells were transiently transfected with the respective constructs using Lipofectamine 2000 (Invitrogen), as described previously (40), and electrophysiological recordings were performed 48 hours after transfection. Transfected cells were identified by their green fluorescence when illuminated at 480 nm. Nontransfected (GFP-negative) cells from the same batch were used as controls. Patch clamp experiments were performed in the tight seal whole-cell configuration at room temperature (20–25°C) using an EPC-10 patch clamp amplifier computer controlled by PatchMaster Classic 1.20 software (HEKA Elektronik). Currents were digitized at 10 kHz and digitally filtered at 2.9 kHz. Patch pipettes had resistances between 2 and 5 MW after filling with the standard intracellular solution. Cells were held at 0 mV, and voltage ramps of 200 ms duration ranging from –100 to +100 mV were applied every 2 seconds. Extracting the current amplitudes at +80 mV from individual ramp current records provided an assessment of the temporal development of membrane currents. Current densities were obtained by normalizing the current amplitude to the cell membrane capacitance. The time course of current development was determined by measuring the current at +80 mV. Steady state protocols were applied from holding potentials of 0 mV and consisted of 400-ms steps to –100 to +100 mV (increment of 20 mV). The standard pipette solution contained 150 mM NaCl, 10 mM EDTA, and 10 mM HEPES/NaOH (pH 7.2). Extracellular solutions contained 150 mM NaCl and 10 mM HEPES/NaOH (pH 7.4) supplemented with 1 mM CaCl2. The osmolarity of the solutions was adjusted to 300–310 mOsm/kg with mannitol.

Pretreatment with EGF, preconditioned media, and cetuximab. The effect of EGF on TRPm6 activity in HEK293 cells transiently transfected with TRPm6 was measured after a 30-minute preincubation with 10 nM mouse EGF (Sigma-Aldrich) in DMEM (modified as described above) or the preconditioned medium. Cetuximab pretreatment for 1 hour with 2 μg/ml in DMEM (modified as described above) was performed prior to treatment with EGF or preconditioned medium. Subsequently, cells were analyzed by the patch clamp technique as described above.

Generation of an MDCK cell line stably expressing pro-EGF and pro-EGF–P1070L. MDCK type I cells (41) were grown in DMEM containing 5% (v/v) FCS (HyClone), 13 mM NaHCO3, 2 mM l-glutamine, and 10 μg/ml ciproxin (Bayer) at 37°C in a humidity-controlled incubator with 5% (v/v) CO2. For stable transfection of MDCK cells, 25 μg circular DNA was transfected using the CaCl2-phosphate precipitation technique (42). Cells were trypanosed 24 hours after transfection, seeded on Petri dishes, and cultured in DMEM containing 75 μg/ml hygromycin B (Invitrogen). Between 10 and 14 days after transfection, individual colonies were selected by means of cloning rings and expanded. Total RNA was extracted from each colony that had been grown until confluent in a 24-well culture plate using TRIzol. Total RNA Isolation Reagent (Invitrogen) according to the manufacturer’s protocol. The obtained RNA was subjected to DNase treatment (Promega) to prevent genomic DNA contamination. Thereafter, 1.5 μg of RNA was reverse transcribed by Moloney Murine Leukemia Virus Reverse Transcriptase (Invitrogen) as described previously (43). The cDNA was used to determine pro-EGF expression with the primers described in Supplemental Table 2. MDCK cells stably expressing plasmid CB7–EGF, plasmid CB7–EGF–P1070L, or empty vector (5 clones of each) were used to seed 10 24-mm Transwell filter wells of each condition (Corning Inc.). We seeded 1.5 × 105 cells/cm2 on permeable filter supports; these were grown for 4 days until they reached confluence and an electrical resistance of greater than 2,400 Ω·cm2. Next, the media were replaced by DMEM medium as described below but without FCS and harvested after 2 days.

Transfection of HEK293 cells. HEK293 cells were transiently transfected with pSAD-EGF, pSAD-EGF–P1070L, or empty-vector pCINeo-IRES-GFP cDNA using polyethyleneimine (Polysciences Inc.). Six million cells were seeded on a petri dish (57 cm2) (10 plates per vector) in a total volume of 5 ml DMEM. After 3 hours, the cells were transfected. The transfection mix for each plate consisted of 15 μg cDNA and 90 μl polyethyleneimine (1 μg/ml) in a total volume of 500 μl Opti-MEM (Invitrogen). The transfection mix was incubated for 20 minutes at room temperature. Subsequently, the transfection mix was added to the cells and incubated overnight. After 1 day of transfection, the culture media was replaced by 8 ml DMEM medium as described above but without FCS.

Collection and concentration of conditioned culture media. Two days after replacement of the medium of HEK293 and MDCK cells, the FCS-free culture media was collected and proteinase inhibitors were added (1 mM PMSF, 1 μg/ml pepstatin A, and 10 ng/ml leupeptin). Subsequently, the collected FCS-free culture media of HEK293 cells with added proteinase inhibitors was concentrated 400 times using Centriprep YM-3 followed by Microcon YM-3 (Millipore). Prior to concentration of the collected HEK293 medium, 2 ml of the collected FCS-free culture medium was kept separately and used for patch clamp analysis.

Human EGF ELISA. EGF concentrations were determined in 400 times concentrated culture supernatant by using Centriprep columns and a human EGF ELISA kit according to the manufacturer’s protocol (RayBiotech Inc.).

Acknowledgments

The authors thank H. van Bokhoven, F. Sweep, and G. Bell for providing genomic control DNA, blood samples, and the cDNA encoding human pro-EGF, respectively. We acknowledge J. van Zoelen for generously donating the antibody against pro-EGF and helpful discussions. The authors are grateful to A. van der Kemp for immunohistochemical analysis, K. Lee for electrophysiological measurements, and A. Heister for linkage studies. This study was supported financially by the Dutch Kidney Foundation (C02.2030 and C03.6017). The authors were further supported by grants from the European Science Foundation (EURYI 2006), the Netherlands Organization for Scientific Research (ZonMW 9120.6110), and the Human Frontier Science Program (RGP32/2004). S. Thébault is supported by an EMBO long-term fellowship (ALTF 727-2005),
and S. Tejpar and E. van Cutsem are senior clinical investigators of the Fund for Scientific Research — Flanders (FWO).

Received for publication January 31, 2007, and accepted in revised form May 8, 2007.