INFLAMMATORY BOWEL DISEASE

Possible role of REG Iα protein in ulcerative colitis and colitic cancer

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Background and aims: Although regenerating gene (REG) Iα protein may be involved in the inflammation and carcinogenesis in the gastrointestinal tract, its pathophysiological role in ulcerative colitis (UC) and the resulting colitic cancer remains unclear. We investigated expression of the REG Iα gene and its protein in UC and colitic cancer tissues. We examined whether cytokines are responsible for REG Iα gene expression and whether REG Iα protein has a trophic and/or an antiapoptotic effect on colon cancer cells.

Methods: Expression of REG Iα mRNA and its gene product in UC tissues was analysed by real time reverse transcription-polymerase chain reaction and immunohistochemistry, respectively. The effects of cytokines on REG Iα promoter activity were examined in LoVo cells by luciferase reporter assay. The effects of REG Iα protein on growth and H2O2-induced apoptosis were examined in LoVo cells by MIT and TUNEL assays, respectively.

Results: REG Iα protein was strongly expressed in inflamed epithelium and in dysplasias and cancerous lesions in UC tissues. The level of REG Iα mRNA expression in UC tissues correlated significantly with severity of inflammation and disease duration. REG Iα promoter activity was enhanced by stimulation with interferon γ or interleukin 6. REG Iα protein promoted cell growth and conferred resistance to H2O2-induced apoptosis in LoVo cells. REG Iα protein promoted Akt phosphorylation and enhanced Bcl-xL and Bcl-2 expression in LoVo cells.

Conclusions: The REG Iα gene is inducible by cytokines and its gene product may function as a mitogenic and/or an antiapoptotic factor in the UC-colitic cancer sequence.

MATERIALS AND METHODS

Tissue specimens and histological examination

Colon biopsy specimens were obtained by endoscopy from 24 patients with UC (15 men and nine women; mean age 45.6 years (range 19–79); mean disease duration 6.4 years (range 0–19)), four patients with Crohn’s disease (two men and two women; mean age 36.0 years (range 26–50)); mean disease duration 5.5 years (range 0–15)), eight patients with proctitis (six men and two women; age range 40–79 years), 10 patients with sporadic colon adenoma (seven men and three women; age range 55–85 years), and five normal controls (five men; age range 33–38 years) at Kyoto University Graduate School of Medicine. Tissue specimens were used for real time reverse transcription-polymerase chain reaction (RT-PCR) and histological analyses. This work was done with the approval of the Review Board of Kyoto University Hospital, and informed consent was obtained from all patients.

A total of seven colitic cancer lesions (location: five rectum, one sigmoid, one descending; histology: four well differentiated adenocarcinomas, three mucinous adenocarcinomas) were obtained from surgically resected specimens from four UC patients (two men and two women; age range 44–58 years; disease duration 11–25 years) and eight sporadic...
colon cancer lesions (location: three rectum, two sigmoid, one descending, one ascending, one caecum; two well differentiated adenocarcinomas, six moderately differentiated adenocarcinomas) were obtained from eight non-UC patients (five men and three women; age range 65–79 years) at Dokkyo University School of Medicine. The colitic cancer tissue specimens were fixed in 10% formalin solution, embedded in paraffin, and subjected to histological analyses. Sporadic colon cancer tissue specimens were used for real-time RT-PCR and histological analyses. This work was done with the approval of the Dokkyo University Surgical Pathology Committee, and informed consent was obtained from all patients.

The diagnosis of UC was based on established endoscopic and histological criteria, and the degree of inflammation was evaluated according to Matts' grade throughout the experiments.

**Immunohistochemical staining**

Immunohistochemical stainings for proliferating cell nuclear antigen (PCNA) and REG Iα protein were performed, as described previously, using antihuman PCNA antibody (PC10; Dako Japan, Kyoto, Japan; dilution 1:1000) and anti-human REG Iα antibody (dilution 1:2000). A cancerous specimen was considered positive for REG Iα when more than 20% of the tumour cells were positively stained.

**Real time RT-PCR**

Total RNA was isolated from colonic biopsy samples and seven human colon cancer cell lines (Caco-2, Colo 205, DLD-1, HT-29, LoVo, SW403, and WiDr) with Trizol reagent (Gibco BRL, Rockville, Maryland, USA). Total RNA (5 μg) was reverse transcribed using oligo-dT primer (Applied Biosystems, Branchburg, New Jersey, USA) and RT product was amplified by PCR, as previously described. TaqMan quantitative real time PCR was performed with the ABI PRISM 7700 Sequence Detection System (Applied Biosystems, Foster City, California, USA). The following set of primers and probe for human REG Iα was prepared: human REG Iα 5′-CTA GAG GCA ACT GGA AAA TAC ATG TCT-3′ (sense), 5′-GTT GGA GAG ATG GTC CGG TTT-3′ (antisense), and 5′-FAM-AAC GGA GTC AAA AAT T (probe). In addition, a set of primers and probe for human glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was synthesised by Applied Biosystems.

Each amplification consisted of a 50 μl reaction mixture with 50 ng of cDNA, 250 nM of REG Iα probe (or 100 nM of GAPDH probe), 900 nM of REG Iα primer (or 200 nM of GAPDH primer), and 1×TaqMan universal PCR master mixture (Applied Biosystems). PCR cycling conditions were 50°C for two minutes, 95°C for 10 minutes, followed by 45 cycles at 95°C for 15 seconds and 60°C for 60 seconds. A template free negative control was included in all amplifications, and each assay was performed in duplicate. The intensity of the fluorescent dye was determined, and expression levels of REG Iα mRNA were normalised to GAPDH mRNA expression levels.

**Cell culture and treatment**

Colon cancer cell lines LoVo and SW403 were routinely maintained in RPMI1640 medium (Invitrogen, Grand Island, New York, USA) with 10% fetal bovine serum (Invitrogen) in a humidified incubator at 37°C with an atmosphere of 5% CO2. These cell lines were used for assessment of REG Iα promoter activity by interferon γ (IFN-γ; Roche, Mannheim, Germany), tumour necrosis factor α (TNF-α; Roche), interleukin (IL)-1β (Roche), IL-5 (Pepro Tech Inc, Rocky Hill, New Jersey, USA), IL-6 (Roche), IL-8 (Roche), and IL-13 (Pepro Tech Inc). In addition, LoVo cells were used for assays of cell growth and apoptosis.

**Luciferase activity assay and transfection**

The human REG Iα promoter from −1195 to +78 was generated from human stomach DNA by PCR using the following set of primers containing Mlu I and Bgl II restriction site, respectively: 5′-CTT AGG CGT GAA TTC CTG GGC TCA AGT GA-3′ and 5′-GCC GAA GAT TTT AGA TCT ACA GTG C-3′. The cloned nucleotides of the promoter were inserted into the position between Mlu I and Bgl II restriction site, upstream of the luciferase gene in the pGL3-Basic vector (Promega, Madison, Wisconsin, USA), and the construct was named hREG Iα-Luc.

LoVo and SW403 cells (2×104) were seeded 24 hours before transfection in 12 well plates (Iwaki, Funabashi, Japan). Cells were cotransfected with 700 ng of a hREG Iα-Luc construct and 7 ng of Renilla luciferase plasmid pRL-TK (as a control of transfection efficiency) in the Optimem medium (Gibco, Grand Island, New York, USA) using FuGENE 6 transfection reagent (Roche, Indianapolis, Indiana, USA) according to the manufacturer’s protocol. Forty eight hours later, cells were stimulated by IFN-γ, TNF-α, IL-1β, IL-5, IL-6, IL-8, and IL-13 for 3, 6, 12, and 24 hours. Luciferase assays were performed using the Dual Luciferase Reporter Assay system (Promega) according to manufacturer’s protocol. Firefly luciferase and Renilla luciferase activities were assayed by luminometer (Lumat LB 9506; Berthold, Germany). Results obtained were normalised for Renilla luciferase activity and expressed as fold of activity of the untreated cell group at the 0 hour time point.

**Transfection and expression of the human REG Iα cDNA**

The full length of human REG Iα cDNA was inserted into the pIRE2-EGFP vector containing cytomegalovirus promoter driving the enhanced green fluorescent protein gene (EGFP) (Clontech, Palo Alto, California, USA). After cloning and verifying the nucleotides of the human REG Iα cDNA by sequencing, the construct was named pIRE2-hREG Iα, and pIRE2-EGFP vector without insert was used as a control.

Plasmids were stably transfected into LoVo cells using FuGENE 6 transfection reagent, as described above. To select cells with stable expression of the pIRE2-hREG Iα and pIRE2-EGFP, cells were cultured for over 3–4 weeks in medium containing G418 (Gibco; 400 μg/ml) and surviving colonies were pooled.

**Preparation of conditioned medium**

To prepare conditioned medium, we cultured human embryonic kidney (HEK) 293T cells in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum. According to the manufacturer’s protocol, cells were transfected with 10 μg of pIRE2-hREG Iα or control plasmid using FuGENE 6 transfection reagent. The medium was replaced by serum free RPMI1640 medium after a 48 hour incubation period. The conditioned medium was then collected and stored frozen as a source of recombinant REG Iα protein.

**Cell growth assay**

Cell growth assay was assessed by Cell Counting Kit-8 (Dojindo, Kumamoto, Japan). LoVo cells (5×103), stably transfected with pIRE2-hREG Iα (LoVo-REG Iα cells) or pIRE2-EGFP (LoVo-EGFP) vector, were plated in 96 well microplates. Cells were incubated in serum free RPMI1640 medium for 24, 48, and 72 hours. After addition of 10 μl of the Cell Counting Kit-8 reagent followed by a three hour incubation, the plates were read at 450 nm in a spectrophotometer (Molecular...
To assess the effect of REG Iα in the medium, we added anti-REG Iα antibody (50 μg/ml) to serum free RPMI1640 medium, and cells incubated for 72 hours were also evaluated in the same procedure.

Cell growth assay was also performed to examine the trophic effect of REG Iα protein on colon cancer cells. Briefly, LoVo cells (5×10³) were cultured in 96 well microplates for 24 hours. Then, the medium was changed to conditioned medium containing human recombinant REG Iα or control medium, and cells were incubated for 24, 48, and 72 hours. To assess the effect of REG Iα in the medium, we added anti-REG Iα antibody to control or REG Iα containing medium, and cells incubated for 72 hours were also evaluated in the same procedure.

**TUNEL assay**

LoVo-REG Iα cells (1×10⁶) and control LoVo-EGFP cells were cultured in four well culture slides (Falcon, Bedford, Massachusetts, USA). Twenty four hours later, cells were incubated for two hours with different concentrations (0–10 mmol/l) of H₂O₂ in serum free medium. Thereafter, cells were incubated in routine medium for 24 hours. After washing with phosphate buffered saline, slides were fixed with 10% buffered formalin for 15 minutes and then stained by In Situ Cell Death Detection Kit (Roche) according to the supplied protocol. TdT mediated dUTP nick end labelling (TUNEL) index was calculated as the percentage of positive cells. To assess the effect of REG Iα in the medium, we added anti-REG Iα antibody (50 μg/ml) to routine medium immediately after treatment with H₂O₂ (5 mmol/l).

**Detection of phosphorylated, non-phosphorylated Akt, and Bcl family proteins**

LoVo cells were cultured in 10 cm dishes for 24 hours. After washing with phosphate buffered saline, the medium was changed to conditioned medium containing human recombinant REG Iα or control medium. LoVo cells (5×10³) were cultured in four well culture slides for 24 hours, followed by a 24 hour incubation with conditioned medium containing human recombinant REG Iα or control medium. Then, cells were incubated for another two hours with different concentrations (0–10 mmol/l) of H₂O₂, and subjected to TUNEL assay 24 hours later. To assess the effect of REG Iα in the medium, we added anti-REG Iα antibody to control or REG Iα containing medium immediately after treatment with H₂O₂ (5 mmol/l).
recombinant REG Iα or control medium, and cells were incubated for another 12 hours. Cells were then mixed with lysis buffer containing 20 mM Tris-HCl (pH 7.4), 150 mM NaCl, 2 mM EDTA, 1% Nonidet P-40 (NP-40), 50 mM NaF, and 10 mM EDTA. After incubation, the cells were lysed with lysis buffer containing 20 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1% Nonidet P-40 (NP-40), 50 mM NaF, and 10 mM EDTA. Protein extract (10 μg) was fractionated by sodium dodecyl sulphate polyacrylamide gel electrophoresis. After transfer to a polyvinylidene difluoride membrane, western blots were performed using anti-REG Iα antibodies (Sigma Chemical Co., St Louis, Missouri, USA), anti-Mcl-1 (BD Sciences, San Jose, California, USA), anti-Bcl-2, anti-Bcl-xL, anti-Akt, antiphospho specific Akt (Ser473) (New England Biolabs, Beverly, Massachusetts, USA), and anti-Bax (Santa Cruz Biotechnology, Santa Cruz, California, USA). The relationship between REG Iα mRNA expression and several clinicopathological factors, and found that REG Iα mRNA expression was significantly enhanced as the severity of endoscopic or histological features increased (fig 1B, 1C). Moreover, REG Iα mRNA expression in UC patients of long duration (>10 years) was significantly greater than those of short duration (<10 years) (fig 1D). We further subdivided patients by both disease duration and histological findings and analysed REG Iα mRNA expression in each group. As shown in fig 1E, in high-Matts score group REG Iα gene expression level in patients with long duration UC was significantly greater than that in patients with short duration UC. In addition, in low-Matts score group REG Iα gene expression level in patients with long duration was also greater than in patients with short duration although this was not statistically significant (p = 0.16). REG Iα mRNA expression in UC tissues was significantly correlated with disease duration (fig 1F) but not with age or sex (data not shown).

Expression of REG Iα protein and PCNA in normal colonic and ulcerative colitis mucosa, and in dysplasia and colitic cancer

In normal colonic mucosa, REG Iα immunoreactivity was detected only in a few epithelial cells in the basal portion of crypts (fig 2A). Colonic epithelial cells positive for PCNA were also observed in the basal portion of crypts (fig 2B).

In UC mucosa, both the number of REG Iα positive cells in crypts and the intensity of REG Iα immunoreactivity were increased. Strongly REG Iα positive cells were mainly observed in the lower part of the colonic mucosa (fig 2C). The number of PCNA positive cells in crypts was also increased, and the distribution of such cells was similar to that of REG Iα positive cells (fig 2D).

REG Iα immunoreactivity was detected in the cytoplasm of not only dysplastic but also colitic cancer cells (fig 2E, 2G). REG Iα expression was detected in all seven of the colonic cancer lesions, originating from four patients examined. PCNA was also strongly expressed in both dysplastic and carcinomatous cells in all colitic cancer tissues examined (fig 2F, 2H).

Expression of REG Iα mRNA in colon cancer cell lines, sporadic colon adenomas, and cancers

Expression of REG Iα mRNA was detected in all seven colon cancer cell lines examined using the RT-PCR method (fig 3). Then, we examined REG Iα mRNA expression in colon adenomas, cancers, and their neighbouring normal colon mucosa by real time RT-PCR. REG Iα mRNA expression level in colon adenomas (n = 10; 159 (51)) was significantly higher than that in normal colon mucosa (n = 10; 1.0 (0.2))

Figure 2  Immunostaining for REG Iα protein and proliferating cell nuclear antigen (PCNA) in normal colon (A, B), ulcerative colitis (C, D), dysplasia (E, F), and colitic cancer (G, H) tissues. REG Iα: A, C, E, G; PCNA: B, D, F, H. Bars = 100 μm.

Figure 3  Detection of REG Iα mRNA in various human colon cancer cell lines by reverse transcription-polymerase chain reaction. GAPDH, glyceraldehyde-3-phosphate dehydrogenase.
REG Ix in ulcerative colitis and colitic cancer

The effect of several cytokines on human REG Ix promoter activity was analysed by transient expression assays in two human colon cancer cell lines (fig 4). In both LoVo and SW403 cells transfected with the hREG Ix-Luc construct, luciferase activity was significantly elevated at three hours after stimulation with IFN-γ (100 and 500 U/ml) and this elevation was sustained for 24 hours. Similarly, human REG Ix promoter activity was enhanced in both cell lines at three hours after treatment with IL-6 (100 and 1000 U/ml) and again the elevation was sustained for 24 hours. In contrast, treatment with TNF-α, IL-1β, IL-5, IL-8, or IL-13 did not affect luciferase activity in the two cell lines.

**Effects of cytokines on REG Ix promoter activity in colon cancer cell lines**

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**Effect of REG Ix on cell growth in LoVo cells**

LoVo cells transfected with pRES2-hREG Ix (LoVo-REG Ix) showed significantly higher WST-8 cleavage levels than LoVo cells transfected with pRES2-EGFP (LoVo-EGFP; control) at each time point (24–72 hours) of incubation (fig 5A). Increased WST-8 cleavage in the LoVo-REG Ix cell group was suppressed almost to control levels by addition of anti-REG Ix antibody (fig 5B), suggesting that enhanced cell growth in LoVo-REG Ix cells is caused by secreted REG Ix protein.

REG Ix-conditioned medium significantly increased WST-8 cleavage in LoVo cells after 24–72 hours of incubation (fig 5D). As shown in fig 5E, the increased WST-8 cleavage in REG Ix-treated LoVo cells was abolished by concomitant administration of anti-REG Ix antibody.

**Antia apoptotic effect of REG Ix in LoVo cells**

The LoVo-REG Ix cell group showed significantly lower TUNEL positivity than the LoVo-EGFP group when they were...
treated with H2O2 at concentrations of 2.5–10 mM, suggesting that REG Ix overexpressing cells are more resistant to apoptosis induced by H2O2 (fig 6A). The decreased TUNEL positivity in the LoVo-REG Ix cell group was significantly reversed by treatment with anti-REG Ix antibody (fig 6B), suggesting that secreted REG Ix protein confers an anti-apoptotic effect on colon cancer cells.

LoVo cells treated with REG Ix conditioned medium showed a significantly lower TUNEL positivity than cells treated with control medium when they were exposed to H2O2 at concentrations of 2.5–10 mM (fig 6C). The decreased TUNEL positivity in REG Ix treated LoVo cells was significantly reversed by concomitant administration of anti-REG Ix antibody (fig 6D).

Treatment with REG Ix conditioned medium enhanced the phosphorylation of Akt in LoVo cells (fig 7A). Moreover, REG Ix gene induction or REG Ix conditioned medium on the growth of human colon cancer cells. LoVo cells transfected with pIRE2-hREG Ix (LoVo-REG Ix) or pIRE2-EGFP (LoVo-EGFP; control) plasmids were cultured for 24, 48, and 72 hours. WST-8 cleavage in each group was measured by ELSA (absorbance OD450), as described in materials and methods. (B) Effect of anti-REG Ix antibody on cell growth of LoVo-REG Ix and LoVo-EGFP. LoVo-REG Ix and LoVo-EGFP cells were incubated with or without anti-REG Ix antibody (Ab, 50 μg/ml) for 72 hours. (C) Source of REG Ix protein. Human embryonic kidney HEK293T cells were transfected with a human REG Ix cDNA expression plasmid or a control plasmid, and the conditioned medium conditioned by these cells was collected. Release of REG Ix protein into the conditioned medium of human REG Ix /cDNA transfected or control plasmid, and the medium conditioned by these cells was collected. Release of REG Ix protein into the conditioned medium of human REG Ix /cDNA transfected or control plasmid, and the medium conditioned by these cells was collected. Release of REG Ix protein into the conditioned medium of human REG Ix /cDNA transfected or control plasmid, and the medium conditioned by these cells was collected. Release of REG Ix protein into the conditioned medium of human REG Ix /cDNA transfected or control plasmid, and the medium conditioned by these cells was collected. Release of REG Ix protein into the conditioned medium of human REG Ix /cDNA transfected or control plasmid, and the medium conditioned by these cells was collected. Release of REG Ix protein into the conditioned medium of human REG Ix /cDNA transfected or control plasmid, and the medium conditioned by these cells was collected. Release of REG Ix protein into the conditioned medium of human REG Ix /cDNA transfected or control plasmid, and the medium conditioned by these cells was collected.
Iz conditioned medium clearly increased Bcl-2 and Bcl-xL expression in LoVo cells whereas REG Iz did not affect Mcl-1 expression (fig 7B).

DISCUSSION

In the present study, we have shown that REG Iz is expressed in only a few small cells in the mid to basal portion of a colonic crypt where putative stem cells reside and continuous cell renewal is occurring.\(^\text{21,22}\) Interestingly, in the gastric epithelium, REG Iz is expressed not only in enterochromaffin-like and chief cells but also in immature small cells in the proliferating neck zone of gastric glands.\(^\text{23,24}\) As REG Iz protein is indeed mitogenic to gastrointestinal epithelial cells,\(^\text{25}\) REG Iz produced in the proliferative zone may play a role in the self renewal of gastrointestinal epithelium under certain physiological conditions.

It is noteworthy that several investigators have identified REG Iz as one of the most abundantly expressed genes in the colonic mucosa of patients with UC by gene chip analyses.\(^\text{26,27}\) even though only a few REG Iz positive cells are present in normal colonic mucosa. Confirming this data, we demonstrated in this study that levels of both REG Iz gene and its protein were significantly increased in UC mucosa. Furthermore, we also showed that enhancement of REG Iz gene expression in the colonic mucosa was correlated with severity of inflammation in UC, as assessed both endoscopically and histologically. These data strongly suggest that REG Iz is important in the inflammatory process of UC.

What factors could be responsible for the increase in REG Iz expression in UC mucosa? As various cytokines play important roles in the inflammation of UC, we examined in this study whether those cytokines enhance transcription of the REG Iz gene in vitro, and found that both IFN-\(\gamma\) and IL-6 significantly stimulated REG Iz promoter activity in colon cancer cells. In contrast, IL-8, IL-1\(\beta\), TNF-\(\alpha\), IL-5, or IL-13 could not enhance REG Iz promoter activity in colon cancer cells. Previous studies have shown that, in addition to IFN-\(\gamma\) and IL-6, TNF-\(\alpha\) and CINC-2\(\beta\), a mouse homologue of IL-8, can enhance REG Iz gene expression in rat pancreatic acinar cells\(^\text{28,29}\) and in rat gastric mucosa in vivo.\(^\text{30}\) The discrepancies between these results and our present data may be due to the use of different species, cell types, or experimental conditions. It is well known that expression of both IL-6 and IFN-\(\gamma\) is prominently increased in UC mucosa.\(^\text{31-33}\) Thus although we did not measure levels of those cytokines in this study, it appears reasonable to suppose that increased levels of IL-6 and IFN-\(\gamma\) in the mucosa of UC are at least in part responsible for the enhanced expression of the REG Iz gene and its product. It may be noted that although UC is known as a Th2 dominant disease,\(^\text{34,35}\) none of Th2 cytokines tested in this study had any effect on REG Iz promoter activity. These data may suggest that the enhanced REG Iz expression in UC mucosa is not a Th2 specific phenomenon but rather a reflection of a general inflammatory condition where IL-6 and IFN-\(\gamma\) expression are elevated.

It is important to clarify whether REG Iz is involved in the carcinogenesis of UC associated colorectal cancer. Interestingly, we found in this study that REG Iz was expressed not only in the epithelial cells of UC mucosa but also in epithelial cells of dysplastic lesions and in colitic cancer cells. Moreover, we have clearly shown that REG Iz expression is significantly increased in patients with longstanding colitis that presents a high risk of colitic cancer.\(^\text{36,37}\) These findings strongly suggest involvement of REG Iz in the UC-colitic cancer sequence. During UC, colonic epithelial cells are continuously injured by inflammation that may induce sustained regeneration of epithelial cells. In the present study, we have shown that REG Iz protein has not only growth promoting but also antiapoptotic actions on colonic cells, and moreover that REG Iz protein exerts its antiapoptotic effect at least in part by activating Akt signalling and enhancing Bcl-xl and Bcl-2 expression. Thus REG Iz protein induced by UC may play a role in protecting colonic epithelial cells from apoptosis. Conversely, however, as REG Iz expression is sustained at high levels in chronic UC, its antiapoptotic action as well as its growth promoting effects may contribute to the development of colitic cancer from UC tissues. Of note, we also found that the distribution of REG Iz positive cells was similar to that of PCNA positive cells in colitic cancer tissues as well as in dysplastic and colitis mucosa. Therefore, REG Iz produced in epithelial cells as well as in cancer cells may exert a direct action on epithelial cells in colitis, and also on cancer cells themselves. In support of this view, addition of anti-REG Iz antibody to the incubation medium abolished not only the growth promoting effect of REG Iz conditioned medium but also the enhanced growth of REG Iz transfected cells. Similarly, anti-REG Iz antibody blocked the antiapoptotic effect of REG Iz conditioned medium and abrogated the reduction in apoptosis in REG Iz transfected cells. These data strongly suggest that REG Iz secreted from epithelial cells or cancer cells exerts growth promoting and antiapoptotic actions in a paracrine or autocrine fashion. On the other hand, it may be noted in this study that colitic cancers develop in only a limited number of patients with UC, although REG Iz is ubiquitously expressed in the inflamed UC mucosa. However, it may be emphasised that carcinogenesis is a complicated event in which many factors are involved. Thus REG Iz may play a role in the UC-colitic cancer sequence as one of the growth promoting as well as antiapoptotic factors, and function in colitic cancer development not as a tumour initiator but as a tumour promoter.

In the present study, we have shown that REG Iz gene expression is increased not only in UC mucosa and colitic cancers but also in sporadic colon adenomas or cancers, using real time RT-PCR analysis, although its expression is not always detected in sporadic colon cancers by northern blot analysis.\(^\text{38,39}\) In addition, REG Iz gene expression was detected in all seven colon cancer cell lines examined. These findings may suggest that REG Iz is involved not only in the UC-colitic cancer sequence but also in the adenoma-carcinoma sequence, although the mechanism of carcinogenesis in UC associated colorectal cancer is believed to be distinct from that in sporadic colorectal cancer.\(^\text{40,41}\) The role of REG Iz in the adenoma-carcinoma sequence should also be clarified in future studies.

In summary, we have shown that REG Iz is expressed not only in epithelial cells of UC mucosa but also in precancerous dysplastic epithelial cells and colitic cancer cells. The promoter activity of the REG Iz gene is stimulated by IFN-\(\gamma\) and IL-6. Moreover, REG Iz protein has both mitogenic and antiapoptotic effects on human colon cancer cells in vitro. Together, these results suggest that cytokine induced REG Iz in the colonic mucosa plays an important role in the development of UC associated colorectal cancer.

ACKNOWLEDGEMENTS

This work was supported in part by grants-in-aid from the Ministry of Education, Culture, Sports, Science and Technology, Japan. We are grateful to Dr Hiroshi Okamoto in Tohoku University Graduate School of Medicine, Sendai, Japan, for providing anti-REG Iz antibody.

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