Isolation and Characterization of a Colonic Autoantigen Specifically Recognized by Colon Tissue-bound Immunoglobulin G from Idiopathic Ulcerative Colitis

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Abstract

Patients with idiopathic ulcerative colitis (UC) have a colon-bound antibody (CCA-IgG) that reacts with colon tissue extracts. We have partially characterized a colon protein that is specifically recognized by CCA-IgG. CCA-IgG was eluted from operative colon specimens from 10 patients with UC. A colon tissue-bound IgG was similarly eluted from six patients with Crohn’s colitis, two with ischemic colitis, and one with diverticulitis. Purified serum IgG from patients with Crohn’s disease, from normal subjects and a patient with myeloma were also used as additional controls. For detection of antigen(s), tissue extracts were prepared from 26 specimens of colon (UC, 12; Crohn’s disease, 6; normal, 4; other controls, 4), 8 specimens of human normal stomach, duodenum, ileum, and liver (2 each). Tissue extracts were also prepared from rats and mice, including germ-free rat colon and rat’s fetal colon. Immunorecognition of CCA-IgG to the tissue extracts was examined by affinity-column chromatography and by transblot analysis. Tissue-extracted proteins were electrophoresed in SDS-polyacrylamide gel, transferred to nitrocellulose sheet, and probed with iodinated CCA-IgG, colonic IgG from other inflammatory bowel disease patients, UC serum IgG, and control serum IgG. Although many proteins were present in colon tissue extracts, 9 of 10 CCA-IgG consistently recognized a protein of 40 kD. None of the nine IgG preparations from colon specimens of patients with Crohn’s colitis and other colonic inflammatory diseases reacted with the 40-kD protein. Five of six symptomatic UC serum IgG and none of eight control serum IgG reacted with the 40-kD protein. The 40-kD protein was present in all colon specimens and it appeared to be organ specific. It was absent in mouse and rat tissues, including colon. The 40-kD protein is not actin and nor a part of the Ig molecule. These results suggest that the 40-kD protein is a colonic “autoantigen” that may initiate a specific IgG antibody response in UC.

Methods

Human specimens

Operative specimens of colon were obtained from 22 patients, including 12 patients with ulcerative colitis, 5 patients with Crohn’s colitis, 2 patients with ischemic colitis, and 2 with diverticulitis. Histologically normal segments of colon were obtained during colonic carcinoma resection from six additional patients with colon carcinoma. Specimens of histologically normal tissues, including ileum, duodenum, stomach, and liver (two each) were obtained from eight patients who underwent abdominal surgery for various reasons. Tissues were received within half an hour of surgery, separated from serosal fat, washed with phosphate-buffered saline (PBS) (10 mM sodium phosphate, 0.15 M sodium chloride, pH 7.5) and stored at −80°C in 10-g aliquots. The study was approved by the Human Experimentation Committee of the Albert Einstein College of Medicine.

10 of 12 patients with UC were operated on due to refractoriness to medical treatment, and in 2 patients, early toxic megacolon were suspected before surgery. All patients were treated with corticosteroid either orally or by intravenous administration and sulfasalazine immediately before surgery. All patients with UC had extensive colitis extending up to right colon. Three of six patients with Crohn’s colitis had involvement of transverse and descending colon, two had ileocolitis with involvement of ascending colon and one had segmental colitis involving the sigmoid colon only. All six patients were treated with sulfasalazine and four received corticosteroids as well. Of the two patients with ischemic colitis, one had involvement of the entire colon.
and the other patient had left hemicolectomy with involvement of splenic flexure area. Diverticulitis was localized in the sigmoid colon in both patients. Grossly diseased areas were used for these studies.

**Rat and mouse specimens**

Six healthy Wistar rats and 10 (nu/+) BALB/c mice were decapitated and specimens of stomach, liver, and the small and large intestine were dissected and stored at −80°C. Similar specimens were obtained from four nu/+ rats that were inbred and kept in germ-free isolators. 10 rat fetuses were obtained by sectioning the uterus of two full-term pregnant rats. The colon specimens from fetuses obtained from each mother were pooled and used for extraction of tissue proteins.

**Extraction and purification of CCA-IgG**

CCA-IgG was extracted and purified as described earlier (8) with slight modifications. A total of 22 colon specimens were used for extraction of tissue bound IgG. These included 10 patients with UC, 6 with Crohn’s colitis, 2 with ischemic colitis, 1 with diverticulitis, and 3 normal colons. Each colon specimen was minced, homogenized, and washed extensively with 0.01 M phosphate buffer with 0.15 M NaCl (PBS), pH 7.2, containing 2 mM phenylmethylsulfonyl fluoride (PMSF), 2 mM EDTA, and 2 mM sodium azide. The final wash was concentrated 1,000-fold and examined for immunoglobulin. Tissue-bound IgG was eluted in acid buffer using 0.02 M citrate, pH 3.2, and 0.1 M glycine-HCl, pH 2.8. The acid eluate was dialyzed against PBS until the pH was adjusted to 7.2 and concentrated by a dialysis membrane (pM 10; Amicon Corp., Scientific Sys. Div., Danvers, MA). IgG was purified by protein A coupled to Sepharose 4B. Purified IgG was examined by double diffusion in agar and sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) (10) and quantitated by radial immunodiffusion.

**Purification of serum IgG**

Serum immunoglobulin G (IgG) from six patients with symptomatic UC and five patients with active Crohn’s disease and three control subjects were purified by ammonium sulfate precipitation followed by DEAE-52 column chromatography (11) and examined by double immunodiffusion, immunoelectrophoresis, and by SDS-PAGE (10).

**Extraction of tissue proteins**

Each tissue specimen was thawed, minced into small pieces, washed five times in 5 vol of PBS, pH 7.2, containing 2 mM PMSF and 2 mM sodium azide. After centrifugation at 1,000 g for 5 min, the tissue pellet was homogenized in the same buffer including 1 M EDTA (5 vol [wt/vol] on ice in a Polytron (Kinematic type PT 1020350D; Switzerland) at 5°C speed for four 10-s intervals (9). Homogenates were centrifuged at 20,000 g for 20 min to remove large tissue particles and ultracentrifuged at 20,000 g for 60 min to obtain a clear supernatant. The protein concentration in the supernatants was adjusted to 5–10 mg/ml and were stored with 10% glycerol at −80°C in 2-ml aliquots. The same procedure was performed for organs obtained from rats and mice. Before each experiment, the extracts were thawed and centrifuged at 20,000 g for 60 min to remove aggregates.

**Affinity column chromatography to examine immune recognition of eluted colon tissue-bound IgG to tissue antigen(s)**

Using protein-A Sepharose CL-4B bound to IgG. 200 µl of washed, hydrated protein-A Sepharose CL-4B (Pharmacia Fine Chemicals, Inc., Piscataway, NJ) in PBS was incubated with 1–2 mg of CCA-IgG and control normal serum IgG. The gel was washed three times with PBS. Tissue extracts containing 5–10 mg of protein were passed slowly through the column. The gel was washed with 10 bed volumes of PBS. Proteins bound to the protein-A Sepharose CL-4B were eluted in 3 bed volumes of 0.02 M citrate buffer, pH 3.2. The resulting protein solution was neutralized with 0.1 M NaOH to pH 7.0 and lyophilized.

Using IgG coupled with cyanogen bromide–activated Sepharose 4B. Colon-eluted IgG and control serum IgG were coupled to Sepharose 4B in the ratio of 4 mg/ml of gel. Gels were equilibrated with 0.01 M PBS, pH 7.2. To remove nonspecific binding proteins, 0.5 to 1 ml of tissue extract in PBS/PMSF containing 5–10 mg of protein was incubated with myeloma IgG coupled Sepharose 4B (2 ml) overnight at 4°C in a side-to-side rotor. The gel was centrifuged at 1,000 g for 15 min and the supernatant was gently passed through small columns (100–200 µl) control serum IgG followed by CCA-IgG coupled Sepharose 4B. The gel was washed five times in 0.01 M PBS, pH 7.2. Bound proteins were eluted in three times gel volume of 2 M potassium thiocyanate in PBS (pH 7.4), diazylated against 0.05 M NH₄HCO₃, and concentrated.

Identification of affinity chromatography–prepared proteins. Proteins were electrophoresed in 7.5% and 10% SDS-PAGE. Gels were stained with 0.02% Coomassie Brilliant Blue. 50 µg of protein from each eluate was iodinated with 125I-Na using the iodogen method (12). Iodinated proteins were separated from free iodine by Sephadex G-25 column chromatography, reduced in 5% β-mercaptoethanol and analyzed by SDS-PAGE. The gels were subjected to autoradiography using a high-speed intensifying screen (Picer Corp., Highland Heights, OH).

**Transblot assay to examine immune recognition of eluted colon tissue bound IgG to tissue antigen(s)**

The assay was performed as described by Towbin et al. (13) with slight modifications. Briefly, equal amounts of tissue extracts were electrophoresed in SDS-PAGE (10%) and transferred to nitrocellulose sheet at 30 V over 16 h. To block nonspecific binding sites, the nitrocellulose sheet was incubated with 5% bovine serum albumin in 0.05 M Tris-HCl containing 0.15 M NaCl, pH 7.9, for 2 h at room temperature. In paired experiments, the nitrocellulose sheet containing transferred proteins was divided into two parts, each contains identical tissue extracts. The nitrocellulose strips were directly probed with iodinated CCA-IgG and control IgG. 50 µg of purified IgG (e.g., CCA-IgG and tissue eluted IgG from Crohn’s or ischemic colitis) was iodinated by the iodogen method (12) as described above. Each strip was incubated with 10 ml of 125I-IgG containing 3 × 10⁸ cpm/ml counts for 2 h at room temperature with constant gentle stirring. The strips were washed once with Tris-saline buffer, pH 7.9, containing 0.1% Triton X-100 and twice with Tris-saline buffer without Triton X-100, dried, and subjected to autoradiography. In the indirect method, strips were incubated with equal amounts of cold IgG for 2 h at room temperature followed by 10 ml of 125I-protein A (3 × 10¹⁰ cpm/ml) for 30 min. To ensure persistence of immunoreactivity of iodinated IgG, we electrophoresed anti-human IgG in parallel to the tissue extracts.

**Transblot assay to examine immune recognition of serum IgG to tissue antigen(s)**

For these experiments, PBS colon extracted proteins were partially purified by the use of DEAE cellulose ion exchange chromatography (11) with discontinuous KCl salt gradient. This semipurification step was necessary to reduce nonspecific bindings of serum IgG to several colon-extracted proteins. 10–15 mg of PBS extracted colonic tissue proteins were passed through a DEAE cellulose column (6-ml bed volume) equilibrated with potassium phosphate (pH 7.4) buffer. Proteins were eluted with 2 bed volumes of different concentrations of KCl such as 0.025, 0.05, 0.1, 0.2, 0.25, 0.3, 0.35, and 0.5 M. Eluted proteins were dialyzed, electrophoresed on SDS polyacrylamide gel, electroblotted to nitrocellulose sheets, and incubated with 125I-CCA-IgG and 125I-serum IgG from patients with UC, Crohn’s disease of colon with or without ileal involvement, and normal subjects. The strips were washed as mentioned above and subjected to autoradiography.

**Two-dimensional gel electrophoresis**

Human colon extract (100 µg/50 µl) was diluted with 2 vol of 8 M urea, 2% (wt/vol) Nonidet P-40, 2% Ampholines (1.6%, pH 5.7, and 0.4%, pH 3–10 [Sigma Chemical Co., St. Louis, MO]), 5% β-mercaptoethanol, 10% glycerol and incubated for 30 min on an isoelectric focusing (IEF) gel prepared as described by Leback and Rutter (14). The focused IEF gel was incubated for 1–2 h in an equilibration buffer.
Table I. Isolation of Colon Tissue-bound IgG from Patients with Inflammatory Bowel Diseases

<table>
<thead>
<tr>
<th>Patients</th>
<th>Number of specimens</th>
<th>Purified colon tissue-bound IgG (μg/g wet tissue (mean±SEM))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ulcerative colitis</td>
<td>10</td>
<td>74.2±19.6*</td>
</tr>
<tr>
<td>Crohn's disease</td>
<td>6</td>
<td>15.2±3.9</td>
</tr>
<tr>
<td>Ischemic colitis and diverticulitis</td>
<td>3</td>
<td>9.0±2.9</td>
</tr>
<tr>
<td>Normals</td>
<td>3</td>
<td>3.3±1.1</td>
</tr>
</tbody>
</table>

* P < 0.05.

Results

Table I summarizes the yield of tissue-bound purified IgG from colon specimens of patients with UC, Crohn’s colitis, ischemic colitis, diverticulitis, and normal colon. The amount of tissue bound IgG was highest in UC specimens with a mean of 74.2 μg/g wet tissue when compared with Crohn’s disease colon of 15.2 μg/g of wet tissue and ischemic colitis and diverticulitis of 9.0 μg/g of wet tissue. The yield of CCA-IgG was significantly (P < 0.05) higher in UC compared with the tissue-eluted IgG from Crohn’s disease and other control tissue specimens (Table I). Tissue-bound IgG was barely detectable in the normal colon. Crude CCA contained mostly IgG, and minute amounts of IgA, albumin, and traces of other unidentified proteins. Protein A–Sepharose 4B removed all IgG present in the crude acid eluate (Fig. 1, and Fig. 2a, lanes 1 and 2).

The immune recognition of CCA-IgG was initially examined by affinity chromatography. After elution of CCA-IgG, the residual tissue pellet was rehomogenized in PBS with 0.1% Triton X-100 and the extracted proteins were passed through the CCA-IgG bound protein-A column. The bound proteins were eluted by citrate buffer at pH 3.2 and were examined by SDS-PAGE as shown in Fig. 2a. Crude CCA (Fig. 2a, lane 1) contained several proteins, including albumin, heavy and light chains of IgG which was separated by protein-A column (Fig. 2a, lane 2). Fig. 2a, lane 4 was the acid eluate and it contained CCA-IgG and an additional 40-kD protein. This suggests that the residual tissue pellet from which CCA-IgG was eluted contained an extractable 40-kD protein(s) which bound to CCA-IgG. Control serum IgG bound to the protein A column did not recognize the 40-kD protein.

The recognition of the 40-kD protein by CCA-IgG was also demonstrated in additional immunoabsorbant column experiments using cyanogen bromide-activated Sepharose 4B coupled to CCA-IgG and control IgG (Fig. 2b). The acid eluates following passage of colon extracts were iodinated, electrophoresed in SDS polyacrylamide gel and autoradiographed and they contained Ig, albumin, and the 40-kD protein (Fig. 2b). The 40-kD protein was present in normal colon extract (Fig. 2b, lane 1) and in the autologous extract of the UC colon specimen (Fig. 2b, lane 2), a part of which was used for extraction of CCA-IgG.

Figure 1. Double immunodiffusion in agar. Protein-A purified CCA-IgG is at the central well. Peripheral wells contain antiserum to γ (designated by Y), α (designated by a), K, and μ chains, and Fab and Fc fragments of IgG.
To examine the specific binding protein(s) with CCA-IgG, we subsequently used more sensitive and direct method of transblot analysis, in which very small amounts of colon eluted IgG were needed. In Fig. 3, the extreme right hand lane demonstrates transferred colon-extracted total proteins from SDS-PAGE to the nitrocellulose sheet that was stained with Coomassie Brilliant Blue. There are many proteins in the colon extract with a wide range of molecular weights. When probed with \( ^{125}\text{I}-\text{CCA-IgG} \), only one band at 40-kD was seen; no bands were seen with control normal serum IgG (Fig. 3). Both UC and normal colon extracts contained the 40-kD protein(s) specifically recognized by CCA-IgG.

In separate experiments, equal amounts of both CCA-IgG and colon tissue-bound IgG eluted from patients with Crohn's disease or ischemic colitis were used in transblot assays against the same extracts of colon from patients with UC and normal controls. Because of the small amount of elutable tissue bound IgG in patients with ischemic colitis and diverticulitis,

Figure 3. An autoradiogram from transblot experiment in which a normal and a UC colon extracts were probed with \( ^{125}\text{I}-\text{CCA-IgG} \) (UC-colon-IgG) and \( ^{125}\text{I} \) control human serum IgG. The extreme right lane shows the Coomassie Blue-stained nitrocellulose sheet strip containing transferred proteins from a normal colon extract. Of all the transferred proteins only one protein of 40-kD reacted with the CCA-IgG. This protein was present in both colon extracts. No recognition was found with control serum IgG. MP, marker proteins. Also note that \( ^{125}\text{I} \)-control serum IgG was recognized by anti-human IgG (α-IgG). K, \( \times 1,000. \)

Figure 4. An autoradiogram from transblot experiment where two UC colon extracts (UC1 and UC2) were probed with \( ^{125}\text{I}-\text{CCA-IgG} \) (UC-colon IgG) and Crohn's disease colon tissue eluted IgG (CD-colon IgG). CCA-IgG reacted only with the 40-kD protein that was present in both UC colon extracts. CD-colon IgG did not react with any of the transferred proteins. Anti-human IgG (α-IgG) reacted with both CCA-IgG and CD-colon IgG, indicating the persistence of immunoreactivity following radiiodination. MP, marker protein; K, \( \times 1,000. \)

Figure 2. Affinity chromatography to examine immune recognition of CCA-IgG to colonic extracts. (a) 0.02% Coomassie Brilliant Blue staining of the SDS-PAGE (10%). All samples were reduced with 5% \( \beta \)-mercaptoethanol. Crude CCA (lane 1) contained albumin and heavy and light chains of IgG. IgG was almost completely separated by Protein-A column (lane 2 and Fig. 1). PBS wash of the Protein-A column (lane 2) after passage of crude CCA contained mostly albumin and traces of heavy and light chains. To examine whether CCA-IgG binds with any protein(s) in the residual UC colon tissue pellet following extraction of CCA-IgG, we further homogenized the pellet in PBS containing 0.1% Triton X-100 and passed extracted proteins through the CCA-IgG bound Protein-A column. Lane 4 contained the acid (pH 3.2) eluate that demonstrates heavy and light chains of CCA-IgG as expected and an additional protein of 40 kD. Lane 3 contained concentrated final PBS wash of CCA-IgG bound Protein-A column after passage of the pellet extract showing no residual unbound protein. Lane 5 contained marker proteins that are included from above downwards 92-, 66-, 45-, 31-, and 14-kD proteins. K, \( \times 1000. \) (b) Autoradiogram of SDS-PAGE (7.5%) of proteins eluted by 2 M potassium thiocyanate from CCA-IgG coupled to cyanogen bromide activated Sepharose 4B columns following incubation of normal colon (lane 1) and UC colon (lane 2) extracts. The bound proteins were iodinated, electrophoresed and autoradiographed. The eluted proteins contained from above downwards large amount of Ig, albumin (66K), and also the distinct band of 40-kD protein. The 40-kD protein was present in both normal colon extract (lane 1) and the autologous colon extract (lane 2).
similar experiments, tissue-eluted IgG from Crohn’s colitis did not react with the 40-kD protein in colon extracts from Crohn’s colitis and normal colon. The recognition of this 40-kD protein(s) by CCA-IgG was demonstrated with 9 of the 10 CCA-IgG preparations and none of the 9 control tissue eluted IgG nor with serum IgG or myeloma IgG.

To determine whether the 40-kD protein is present in other human tissues, we probed extracts of normal ileum, duodenum, stomach, and liver with 125I-CCA-IgG and control serum IgG. The 40-kD protein was not detected by CCA-IgG in any of these tissues, except colon (Fig. 5). Control IgG did not react with any extract, including colon. One of the colon extracts was from the autologous colon (Fig. 5) from which the CCA-IgG was extracted. The 40-kD protein was present in this autologous colon extract. Similar results were obtained with three additional patients with UC.

To examine species specificity, we performed experiments using extracts of tissue from conventional rats and rats inbred and kept in germ-free conditions and mice that included colon, small intestine, stomach, and liver. Extracts of colon specimens from rat fetus were also used. There were many protein bands in the 40-kD area, as seen by Coomassie Blue staining of the nitrocellulose sheet. However, CCA-IgG did not react with any of the tissue extracts (Fig. 6). Fig. 6 demonstrates a paired experiment in which rat tissue extracts and human tissue extracts were probed with the same 125I-CCA-IgG. Although the 40-kD proteins were detected by 125I-CCA-IgG in the human colon extracts, it was not seen in all rat tissue extracts, including the colon. One human tissue extract was obtained from the ileocecal area, which was grossly normal. This patient had a right hemicolectomy for carcinoma of the ascending colon. Although ileal tissue extract did not react with CCA-IgG, extract from the ileocecal area reacted (Fig. 5).

![Figure 5](http://www.jci.org)  
**Figure 5.** An autoradiogram from transblot experiment in which extracts of four colon specimens, one from Crohn’s disease (CD), one normal colon (N. Col.), and two UC colons (UC1 and UC2) were electrophoresed and transferred to nitrocellulose sheet. In addition, extracts of histologically normal human stomach (ST), duodenum (DUO), ileum, and liver were electrophoresed. All extracts were probed with 125I-CCA-IgG. The 40-kD protein(s) were detected in all colon extracts. Extracts of stomach, duodenum, ileum, and liver did not react with the CCA-IgG. 125I-CCA-IgG reacted with the autologous colon extract as indicated in the figure. K, × 1,000.

![Figure 6](http://www.jci.org)  
**Figure 6.** An autoradiogram from transblot experiment in which human tissue extracts and rat tissue extracts prepared from liver, stomach (ST), small intestine (S.I.), and large intestine (L.I.) were probed with 125I-CCA-IgG (UC-colon IgG). All human colon extracts, including UC, normal (Nor.) colon, and normal ileocecal (Nor. IL-CeC) area, contained the 40-kD protein as detected by CCA-IgG. Neither the human liver nor any of the rat tissue extracts reacted with the CCA-IgG. MP, marker protein; K, × 1,000.

To determine the immunoreactivity of the 40-kD protein with serum IgG, we purified serum IgG from six patients with symptomatic UC, five patients with active Crohn’s disease involving colon with or without ileal involvement, and three normal subjects. Initial immunoblot experiments using serum IgG and colon tissue extracts proteins caused significant high background due to nonspecific bindings of several proteins with serum IgG from all subjects. To reduce this nonspecific binding, we developed a method for purification and enrichment of the 40-kD protein by ion exchange chromatography with discontinuous KCl salt gradients. Proteins eluted with 0.35 M KCl contained enriched 40-kD protein as detected by CCA-IgG (Fig. 7). Subsequently, 0.35 M KCl eluted proteins were used to examine the immune recognition by each of 14 purified serum IgG (Fig. 7). Five of six patients with symptomatic UC reacted with the 40-kD protein by immunoblot analysis; no immune recognition was observed by serum IgG from five patients with symptomatic Crohn’s disease and three normal subjects.

To examine the homogeneity and determine the isoelectric point (pl) value of the 40-kD protein, we performed two-dimensional SDS-PAGE followed by immunoblot studies, as shown in Fig. 8. The sample resolved by IEF demonstrated a major band of 40-kD size immunoreactive to CCA-IgG at pH 5.8 and two minor components of the same molecular weight with more acidic pl.

To examine whether the 40-kD protein is a part of the immunoglobulin molecule, we performed experiments using both affinity column chromatography and transblot system. An UC colon extract was sequentially passed through affinity columns prepared with purified serum IgG, anti-human IgG, IgA, and IgM (IgG fraction; Cappel Laboratories, Cochranville, PA) coupled with cyanogen bromide–activated Sepharose 4B. The PBS elutes containing unbound proteins and the 2-M potassium thiocyanate eluates from these columns containing the bound proteins were examined by transblot assays using
electrophoresed in lanes 1-4. All lanes contained 80 μg of total proteins isolated from the same colon tissue. MP stands for marker proteins and α-IgG contained anti-human IgG. The 40-kD protein(s) is enriched and a limited number of proteins, compared with PBS colon extract (Fig. 3, extreme right lane), can be seen in the 0.35 M KCl eluates. (b) Autoradiogram from transblot experiment after the same nitrocellulose strips as in a were incubated with radioiodinated CCA-IgG, serum IgG from a symptomatic patient with UC (UC-IgG), an active Crohn’s disease patient (CD-IgG), and serum IgG from a normal subject. The 40-kD protein reacted with the CCA-IgG and also with UC serum IgG. Neither CD-IgG nor normal serum IgG reacted with the 40-kD protein. There were some other nonspecific faint bands that were common to UC and control serum IgG.

125I-CCA-IgG. The 40-kD protein was present in all PBS eluates (Fig. 9) and did not bind to any of the column(s). In separate transblot experiments, colon-extracted proteins were probed with radioiodinated anti–human IgG, IgA, and IgM. Although there was recognition of IgG, IgA, and IgM, respectively, in the colon extracts, no reaction was noted with the 40-kD protein. These results suggest that the 40-kD protein is not derived from an Ig molecule.

Since actin is 43 kD and colon tissue contains actin, we examined whether the 40-kD protein could be actin and if CCA-IgG is directed to actin. Fig. 10 demonstrates an autoradiogram from a transblot experiment after incubation of a

125I-CCA-IgG with transferred proteins of a UC colon extract. Passed sequentially through affinity columns prepared with purified serum IgG and anti–human IgG, IgA, and IgM coupled to cyanogen bromide–activated Sepharose 4B. The PBS elutes containing unbound proteins were probed with 125I-CCA-IgG. The 40-kD protein was present in all the eluates including the original UC extract. These results suggest that the 40-kD protein is not a part of Ig, UC1, original UC colon extract; UC1a, PBS eluate from serum IgG coupled Sepharose 4B column; UC1b, PBS eluate from anti–human IgM coupled Sepharose 4B column; UC1c, PBS eluate from anti–human IgA coupled Sepharose 4B column; UC1d, PBS eluate from anti–human IgM coupled Sepharose 4B column; MP, marker proteins; α-H-γ, antisera against human γ-chain. K, × 1,000.

UC colon extract, pure chicken actin (Sigma Chemical Co.) and human fibroblast extract with 125I-CCA-IgG and antiactin antibody. CCA-IgG, although recognizing the 40-kD protein,

125I-CCA-IgG. The 40-kD protein was present in all PBS eluates (Fig. 9) and did not bind to any of the column(s). In separate transblot experiments, colon-extracted proteins were probed with radioiodinated anti–human IgG, IgA, and IgM. Although there was recognition of IgG, IgA, and IgM, respectively, in the colon extracts, no reaction was noted with the 40-kD protein. These results suggest that the 40-kD protein is not derived from an Ig molecule.

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UC colon extract, pure chicken actin (Sigma Chemical Co.) and human fibroblast extract with 125I-CCA-IgG and antiactin antibody. CCA-IgG, although recognizing the 40-kD protein,
did not react with actin that was recognized by antiactin. Antiactin reacted with actin present in the fibroblast extract and the UC extract.

Discussion

These studies confirm demonstration (8, 9) of a colon tissue-bound IgG antibody in patients with UC, which we termed CCA-IgG. Using affinity-column chromatography and transblot, we demonstrated that CCA-IgG recognizes a colon tissue-specific 40-kD protein(s). This reactivity was specific for CCA-IgG and was absent with colon tissue-eluted IgG from patients with Crohn’s disease and ischemic colitis and purified serum IgG from healthy subjects and a myeloma patient. Furthermore, anti-human IgG, IgA, and IgM also did not react with the 40-kD protein.

In previous experiments (8, 9), we did not detect tissue-bound IgG in colon from patients with Crohn’s colitis and normal patients (colon cancer patients). With improved methods and using several protease inhibitors, we increased the yield of elutable colon tissue-bound IgG in UC, and isolated small amounts of tissue-bound IgG from colon specimens of patients with Crohn’s colitis, ischemic colitis, and diverticulitis. However, tissue-bound IgG is significantly (P < 0.05) higher in patients with UC than in patients with Crohn’s or ischemic colitis (Table I). Tissue-bound IgG in normal controls was nil or a trace amount that was not sufficient to perform further studies.

In previous studies using an indirect immunofluorescence assay (8) and radioimmunoprecipitation method (9), we demonstrated preferential recognition of UC colon tissue extract by CCA-IgG as compared with control IgG and control colon tissue. Using direct and more sensitive transblot experiments, we confirmed the disease specificity of CCA-IgG compared with several control IgG preparations, including colon tissue-eluted IgG. A striking finding is the unique recognition of a possibly single (or multiple with close molecular weights) 40-kD protein among many proteins present in colon extracts. In contrast to our previous reports, CCA-IgG recognized the 40-kD protein in diseased and normal colon extracts probably due to increased sensitivity of the current assay. The other possibility is that in the indirect immunofluorescence assay used earlier, the “antigen(s)” in normal colon tissue is not “exposed” to be detected by CCA-IgG, whereas in patients with UC it is exposed. Whether there is any difference in the quantity of the 40-kD protein in different colon tissue is unknown. Further studies of purification of the 40-kD protein and its quantitation may answer this issue. If it is not exposed in normal tissue, it will be important to study what factors are involved in triggering antigenicity.

Circulating heterogeneous antibodies against various alimentary tract antigens, components of intestinal bacteria, e.g., Escherichia coli 0:14 polysaccharide, antigen(s) from germ-free rat feces, and rat colon epithelial glycoproteins, have been demonstrated in UC and in Crohn’s disease (3, 4, 16–19). These antibodies belonged to IgG, IgA, and IgM classes (19). The colon antigen from rats is a heat-stable polysaccharide, located in the mucus-producing cells of the colonic mucosa and in the mucus (20). Various epithelial cell-associated components from murine small intestine, but not from kidneys, have been found to cross-react with sera from both UC and Crohn’s disease patients (21). Using fluorescence-activated cell-sorter, circulating antibodies reactive against colon epithelial cells from Wistar rats have been reported in patients with UC (22). In contrast to these studies, using CCA-IgG, the 40-kD protein was not detected in the intestinal extracts (both small and large intestines) of two strains of rats, including germ-free rats and mice, suggesting the species specificity of the 40-kD protein. We recently identified intestinal tissue antigens specific for Crohn’s disease tissue that reacted with Crohn’s disease serum IgG and not with UC serum IgG (23). The molecular weights of these proteins were 160,000, 120,000, and 110,000 and CCA-IgG did not react to any of these proteins present in Crohn’s disease tissue (Fig. 5). Whether the specific recognition of the 40-kD protein in human colon extracts by CCA-IgG is related to a bacterial antigens such as E. coli and enterobacterial common antigen is unknown. Immunofluorescent studies using E. coli and bacteroids grown from feces of normal subjects did not react to CCA-IgG (9). Further studies using purified colonic bacterial proteins are needed to examine the cross-reactivity of the 40-kD protein.

Serum and purified serum IgG from patients with symptomatic UC demonstrated antibody-dependent cell-mediated cytotoxicity against a colon cancer cell line RPMI-4788 and not against HeLa cells (24, 25). Whether CCA-IgG recognizes the surface protein(s) in the colon cancer cell line and whether it is cytotoxic is unknown. The 40-kD protein detected by CCA-IgG appears to be colon-specific, since extracts of human tissues from ileum, duodenum, stomach, and liver did not react with CCA-IgG. The 40-kD protein is not actin, as it does not react with antiactin. Moreover, though CCA-IgG reacts with the 40-kD protein, it does not react with actin. Serum IgG from patients with symptomatic UC reacted with the 40-kD protein as demonstrated by immunotransblot assay. Serum IgG from patients with active Crohn’s disease and control subjects did not react with the 40-kD protein. Using the purified 40-kD protein, further studies are needed to examine whether CCA-IgG and circulating antibodies from different patients with UC are similar, particularly in relation to their antigen specificity.

The fact that 9 of the 10 CCA-IgG and none of the 9 control diseased colon tissue eluted IgG demonstrated immunoreactivity suggests this phenomenon to be unique to UC. Whether this is primary or secondary cannot be completely resolved until the 40-kD protein is further characterized. The presence of the 40-kD protein in UC tissue and in normal colon tissue suggests its possibility to be an “autoantigen” that indeed reacted with the CCA-IgG purified from the autologous colon as well (Fig. 5). However, the role of the 40-kD protein in initiating the disease process, if any, is unknown. Specific autoantibodies against the double-stranded DNA in systemic lupus erythematosus, against thyroglobulin in autoimmune thyroiditis, and against acetylcholine receptor in myasthenia gravis play important roles in the pathogenesis of these diseases (24–26). Many patients who are severely ill have high titers of the specific autoantibodies and monitoring these circulating autoantibodies was extremely helpful in assessing the clinical course (27–30). Future studies using purified 40-kD protein as antigen may enable identification and accurate measurement of CCA-IgG in the circulation, which may help in diagnosis and assessing the clinical activity of the disease.

Acknowledgments

The authors gratefully acknowledge help from Dr. Richard J. Stockert for the two-dimensional gel experiments; valuable discussions with
Anatol G. Morell, Srilata Bagchi, Jayanta Roy Chowdhury, Betty Diamond, and Barbara Birshtein; and the excellent secretarial assistance of Anna Caponigro in preparing the manuscript. This study was supported by research grant NIADDK-AM-21832 from the National Institutes of Health, Bethesda, MD. Dr. Das is the recipient of the Irma T. Hirschl Career Development Award.

References