Major Parietal Cell Antigen in Autoimmune Gastritis with Pernicious Anemia is the Acid-producing H⁺,K⁺-Adenosine Triphosphatase of the Stomach

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Abstract

In autoimmune gastritis antibodies against a membrane-bound parietal cell antigen of previously unknown function are present in the sera of patients. In this study, a vesicular membrane preparation of porcine gastric mucosa cells was found to be a potent antigenic source. This preparation blocked >90% of antibody binding to a lysate of gastric mucosa cells. The membrane fraction contained H⁺,K⁺-ATPase (EC 3.6.1.36) as the major protein, which in sodium deoxycholate-polyacrylamide gel electrophoresis migrated with a weight of 92 kD. After reduction and alkylation, this component was resolved into two bands of similar staining intensity (92 and 88 kD). Immunoblotting analysis showed that sera of patients recognized antigen with pattern identical to the major protein of the vesicular membranes.

Protein A-Sepharose beads preincubated with immunoglobulins of five individual patients (but not control) sera were all found to reduce both the H⁺,K⁺-ATPase activity and the amount of parietal cell antigen of a preparation of vesicular membranes solubilized in n-octylglucoside. Taken together, the results of this study indicate that the major parietal cell antigen is identical to the acid-producing enzyme, H⁺,K⁺-ATPase, of the parietal cell.

Introduction

Autoimmune gastritis (chronic gastritis type A) is characterized by a disturbed parietal cell function resulting in a reduced gastric acid production (1). In advanced stages of the disease, achlorhydria and mucosal atrophy in the gastric body and fundus are found. Apart from producing acid, the parietal cells also secrete intrinsic factor (2), the 60-kD glycoprotein (3), which is essential for vitamin B₁₂ absorption. Deficiency of this vitamin results in pernicious anemia and neurological symptoms.

The pathogenesis of autoimmune gastritis seems to involve an immune-mediated destruction of the parietal cells. This is evidenced by a lymphocytic infiltrate in the mucosa, which consists predominantly of B cells (4), and by the production of autoantibodies (1, 5). Injection of patient IgG into rats reduces the gastric acid output and may induce atrophy of the mucosa (6). This suggests that the autoantibodies have a key role in the expression of the disease. The antibodies are directed against a protein localized in the canalicular areas of the cell (7). Since the function of this protein has remained elusive, we have made attempts to disclose its physiological role.

In the present study we have identified the parietal cell protein that seems to serve as the major antigen in autoimmune gastritis. This molecule is identical to the acid-producing enzyme, H⁺,K⁺-ATPase (EC 3.6.1.36), of parietal cells.

Methods

Patient sera and immunoglobulin preparations. Sera from ten patients, six women and four men, ages 35–82 yr (mean age 61 yr), with a diagnosis of pernicious anemia and low vitamin B₁₂ levels, were used. The subjects had impaired vitamin B₁₂ absorption by a Schilling test, which was normalized after addition of intrinsic factor. Eight patients had been further investigated and found to have achylia/hypochylia and/or atrophic fundic gastritis. All sera were positive in a routine immunofluorescence test for parietal cell antibodies. Control sera were obtained from healthy individuals lacking parietal cell antibodies. IgG were prepared by ammonium sulfate (1.6 M) precipitation of sera, followed by extensive dialysis against 20 mM Tris-HCl buffer, pH 7.4. The final concentration of the IgG preparations ranged between 10 and 15 mg/ml, as determined by the method of Lowry et al. (8).

Preparation of gastric mucosa cells, parietal cells, and vesicular membrane fractions. Porcine gastric mucosa cells were prepared by consecutive digestions with pronase and collagenase. Parietal cells were obtained by further purification of mucosa cells by centrifugation on Percoll gradients and elutriation (9, 10). The purity of parietal cell fractions was ~90%. Tubulovesicular membranes were prepared after homogenization of gastric mucosa and sucrose-Ficol gradient centrifugation (11). A porcine kidney membrane preparation containing Na⁺,K⁺-ATPase was obtained as described elsewhere (12).

Parietal cell antibody assays. Parietal cell antibodies were measured in patient sera by routine indirect immunofluorescence analysis using fixed sections of rat stomach. An ELISA was developed to determine antibody binding to gastric mucosa cells and vesicular membranes. Flat-bottomed microtiter plates (Nunc, Copenhagen, Denmark) were coated overnight at +4°C with 100-μl aliquots of porcine gastric mucosa cells (2 × 10⁶ cells; 0.5 mg protein, lysed in 100 μl 0.2% Triton X-100) diluted 50-fold with PBS or with 100-μl aliquots of vesicular membranes diluted in PBS. Subsequently, the wells were washed four times with 250 μl PBS with 0.5% vol/vol Tween 20 (PBS-T). Then, 100-μl aliquots of patient sera diluted 1:200 in PBS-T were added and incubated for 1 h at room temperature followed by four washes. Subsequently, 100 μl of β-galactosidase conjugated rabbit anti-human IgG (Pharmacia Fine Chemicals, Uppsala, Sweden) was added. After 1 h at room temperature the wells were washed four times. A substrate solution containing 100 μl of ortho-nitrophenyl-β-galactoside (Pharmacia Fine Chemicals) was added for 30 min at room temperature. The reaction was stopped with 100 μl 0.66 M Na₂CO₃ and the absorbance measured at 405 nm.

PAGE experiments. Antigen preparations were separated by electrophoresis in polyacrylamide gels, 4% stacking and 8% separating gels.

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in Tris-glycine buffer with SDS (SDS-PAGE) as described by Laemmli (13). The antigen preparations were analyzed nonreduced, as well as after reduction (0.2 M β-mercaptoethanol) and alkylation (0.2 M iodoacetamide). Protein staining of the gels was performed with Coomassie Blue R-250.

In immunoblotting experiments the separated proteins were transferred after electrophoresis onto nitrocellulose membranes (0.45 μm pore size; Bio-Rad Laboratories, Richmond, CA) for 16 h at 200 mA. The membranes were blocked by incubation in 1% BSA-PBS for 30 min, washed twice in PBS-t for 30 min, incubated with dilutions of patient and control sera in PBS-t for 60 min, and washed twice in PBS-t followed by incubation with a 1:200 dilution of horseradish peroxidase rabbit anti-human IgG conjugate (Dakopatts AB, Hagersten, Sweden) in PBS-t buffer for 60 min. The blots were washed twice in PBS-t and then incubated with a solution containing 4-chloro-1-naphtol (color development reagent, Bio-Rad Laboratories) to detect antibody binding to the blots (14). All incubations and washes were carried out at room temperature.

**Solubilization procedure.** A vesicular membrane preparation (7.2 mg protein/ml) was solubilized with 1.3% n-octylglucoside (Boehringer Mannheim Biochemicals, Mannheim, FRG) at 4°C for 30 min followed by centrifugation at 70,000 g for 1 h. The solubilize (6.8 mg protein/ml), when tested for ATPase, showed reduced enzyme activity (15) estimated to be 5% of the native preparation.

**Adsorption experiments.** Protein A (1 mg) conjugated to Sepharose beads (Pharmacia Fine Chemicals) were preincubated at 37°C for 60 min with IgG preparations (1.5 mg protein) of sera from patients with autoimmune gastritis and from healthy controls, respectively. The amounts of IgG bound to the protein A-Sepharose beads were the same with patient or control preparations. After washing in fresh buffer (20 mM Tris-HCl, pH 7.4), 35 μg of solubilized membranes in 0.16% n-octylglucoside was added and incubated for 1 h at 37°C. The mixture was centrifuged (2,000 g) and 150 μl of the supernatant (5 μg protein) was analyzed for ATPase activity in incubation with a buffer containing 5 μM 32P-labeled ATP (~ 25,000 CPM), 2 mM MgSO4, 10 mM KCl, and 20 mM Hepes, pH 7.4, for 30 min at 30°C (11, 16). In the absence of K+ in the incubation buffer < 5% of the total ATPase activity remains (11). The amount of 32Pi formed was quantitated as an ammonium molybdate complex extracted into an isobutanol/toluene (1:1) phase. In the absence and presence of added enzyme, 0.8 and 8.2% of the ATP substrate was hydrolyzed, respectively. IgG-coated beads caused a nonspecific, ~ 40%, reduction of the enzyme activity, possibly in part due to residual ammonium ions in the IgG solutions.

The supernatants were also analyzed for the presence of parietal cell antigen. This was accomplished by measuring inhibition in an antibody binding assay. Microtiter plates precoated with a preparation of porcine vesicular membranes (0.4 μg/well) were incubated with 25 μl of supernatants, together with a 1:10,000 dilution of a patient serum with a high titer (> 1:400 as determined by routine immunofluorescence testing) of parietal cell antibodies. Subsequently, the inhibitory effect of the different supernatants on antibody binding was determined by the ELISA described above.

**Statistics.** Experimental data were analyzed for statistical significance by the unpaired t test.

**Results.**

In initial experiments the binding of antibody to microtiter wells, which were coated with lysates of gastric mucosa cells, was determined. Marked antibody binding was observed with sera of patients with autoimmune gastritis (Fig. 1, left). The mean±SD absorbance at 405 nm of 10 patient sera was 0.39±0.12, compared with 0.19±0.06 for sera of 10 healthy controls (P < 0.0005). In subsequent experiments, fractions of mucosa cells were examined for antigenicity. The antigen is known to be membrane bound (7) and a preparation of vesicular membranes isolated from gastric mucosa cells was found to be ~ 100-fold more potent than mucosa cell lysates (Fig. 1, right). In a separate experiment the capacity of vesicular membranes to inhibit parietal cell antibody binding to gastric mucosa cell lysates was studied. More than 90% of antibody binding was blocked by high concentrations of gastric vesicular membranes, whereas porcine kidney membrane preparations, rich in Na+/K+-ATPase, did not affect antibody binding (Fig. 2). These data thus indicated that gastric vesicular membranes contain the major autoantibody binding material.

Further experiments were conducted to characterize the parietal cell antigen. When subjected to SDS-PAGE under
In nonreducing conditions, multiple protein bands of gastric parietal cell preparations were stained (Fig. 3). A specific immunoblot signal of 92 kD was always obtained in a position with only a minor amount of protein staining. In some experiments, weak nonspecific bands, corresponding to the major protein components of mucosa cell lysates, were observed with both patient and control sera. In the vesicular membrane preparation, one major protein band with an approximate molecular weight of 92 kD was present. Upon immunoblotting analysis of nonreduced material, the antigen was localized in most experiments mainly to a 92-kD band (Figs. 3 and 4). In some instances, however, a blurred, broad band of 65–75 kD was seen in addition to the 92-kD band (Fig. 4). The basis of this variable behavior of nonreduced material upon immunoblotting analysis was not further explored.

After reduction (0.2 M β-mercaptoethanol) and alkylation of the vesicular membranes, the immunoblotting results were reproducible. The major antigen appeared in two sharp bands of similar intensity with weights ~ 92 and 88 kD (Fig. 4). These bands were also seen in the protein staining, indicating that the parietal cell antigen corresponded to the major protein present in the vesicular membrane fraction. After reduction with high concentration (1M) of β-mercaptoethanol, the 92-kD material appeared as a 88-kD band, and in addition as a broad band of lower mobility ~ 100–105 kD. With low concentration of the agent (0.002 M) a minor amount of the 88-kD band was found, the 92-kD band being the major component. Again, in these experiments the immunoblotting patterns corresponded to the major protein stainings (data not shown).

When vesicular membranes solubilized in n-octylglucoside were incubated with protein A–Sepharose beads precoated with five patient or five control IgG, all patient immunosor-
crosomal fraction of homogenates of human gastric mucosa (18). At the ultrastructural level, the antigen is localized to the cell membrane forming the microvilli of the parietal cells, a part of the cell suggested to play a role in the production of hydrochloric acid (7). Many patients (40–60%) with pernicious anemia, due to autoimmune gastritis, also have antibodies against intrinsic factor (17), a glycoprotein secreted by the parietal cells. The experiments of the present investigation demonstrated the ability of porcine vesicular membranes to inhibit the majority of antibody binding to gastric mucosa cell lysates. This indicates that, from a quantitative point of view, the major antigen of parietal cells is localized to vesicular membranes. These preparations are rich in tubulovesicles of parietal cells (11) and contain as the major component a protein that can be phosphorylated by [γ-32P]ATP (19). This protein represents H⁺,K⁺-ATPase (19, 20), the hydrogen- ion pump of the stomach, which has been reported to have a weight ~ 94 kD in its subunit form (11, 15). Recently, a complete nucleotide sequence of a cDNA clone of the rat stomach H⁺,K⁺-ATPase, corresponding to a 114-kD protein, was described (21). In the present study a weight of 92 kD was determined for the major porcine parietal cell antigen in the nonreduced state. We have obtained similar data with a preparation of tubulovesicles of human origin (unpublished observations). Upon reduction and alkylation, two separate bands of 92 and 88 kD emerged. These bands had similar intensity in protein staining and were equally antigenic in immunoblotting experiments, suggesting that the bands may represent two different populations of the enzyme subunit. Apparently, these two proteins are not disulfide linked, but reduction and alkylation affects intrachain reactive group/groups. The parallelism between major protein staining and immunoblotting patterns in experiments with membrane material was considered of prime importance as the results indicated identity between the major protein (H⁺,K⁺-ATPase) of the gastric vesicles and the parietal cell antigen. The details of the reactions taking place after reduction and alkylation, as well as the basis for the molecular weight differences between the different bands, will be the subject of further studies.

Vesicular membranes containing H⁺,K⁺-ATPase can be solubilized by detergents such as n-octylglucoside and cholate (22, 23). After solubilization enzymatic activity is associated with higher molecular weight forms thought to reflect tetrameric, trimeric, and dimeric forms of the protein in addition to monomeric material (23). Solubilization is accompanied by a gradual loss of enzymatic activity. In the present study, ~ 5% of the original enzyme remained in 0.16% n-octylglucoside. Solubilizes containing monomeric enzyme regain enzymatic activity when added to liposome preparations (22). In this study, solubilized vesicular membranes were used to investigate direct antibody binding. Solubilized enzymatic activity was found to bind to antibody-coated protein A–Sepharose beads, which similarly bound parietal cell antigen. The data thus further demonstrate the identity between H⁺,K⁺-ATPase and the major parietal cell antigen, as suggested by the electrophoresis and immunoblotting experiments.

Autoimmune gastritis is accompanied by a reduced gastric acid production and a gradual loss of the gastric mucosa, which is infiltrated by a large number of lymphocytes, mainly B cells (4). Injection of IgG from patients into experimental animals results in reduced acid production and mucosal atrophy (6). A direct inhibitory effect of patient IgG on acid secretion of bull frog gastric mucosa in vitro has been reported, as well as an inhibition of gastrin-stimulated carbonic anhydrase activity as analyzed by a cytochemical assay (24). Further, gastrin receptor blocking antibodies have been described by the use of a radioreceptor assay with isolated parietal cells and by inhibition of gastrin stimulation of aminopyrine accumulation in parietal cells (25), an indirect index of HCl production (26). In this latter study, gastrin receptor antibodies were detected only in a fraction of patients with pernicious anemia, whereas all patients studied had parietal cell antibodies. These investigators reported a 65–70-kD protein, albeit not detectable in protein staining, as the presumptive microsomal antigen in pernicious anemia. The antigen was identified by immunoblotting analysis using patient sera and parietal cell homogenates from rat and dog or gastric microsomes from mouse, dog, and rat (27). The protein appeared antigenic only in sulfhydryl nonreduced state. Further, a second, distinct protein of ~ 88 kD was also reported to react, but less well than the 65–70-kD band, with parietal cell microsomal antibodies. This protein was detected only with dog mucosa and appeared antigenic only in sulfhydryl reduced form.

Our results and interpretation of electrophoretic and immunoblotting experiments are at variance with those of Dow et al. (27). In our opinion, the major parietal cell antigen is the protein band of 92 kD typically found with nonreduced material and the two sharp bands of 92 and 88 kD present after reduction and alkylation of vesicular membranes. The antigenicity of this material, in the nonreduced as well as reduced and alkylated forms, was demonstrated with the use of pooled patient sera. Identical results were obtained when several individual sera were tested (not shown). However, we have noted that certain patient sera, despite being positive for parietal cell antibody by immunofluorescence and reactive against native vesicular membranes in ELISA, do not give an immunoblot signal with reduced and alkylated vesicular membrane material after electrophoresis in SDS-PAGE. This suggests that such sera contain antibodies directed against epitopes sensitive to denaturation (unpublished observations). As exemplified in Fig. 4, a blurred, broad band of 65–75 kD was sometimes found upon immunoblotting of nonreduced vesicular membranes. In the 65–75 kD position no corresponding protein staining was detected. These observations are in accordance with those described by Dow et al. (27) regarding a 65–70-kD band. The basis of this variable antigenic behavior is the subject of further investigation.

Thus, the present study has identified the acid-producing enzyme, H⁺,K⁺-ATPase, of parietal cells as the major antigen in autoimmune gastritis. Preliminary results from our laboratory indicate that some autoantibodies against this antigen can directly block the enzyme activity (unpublished observations). Parietal cell antibodies possibly interfere with parietal cell function via complement-mediated mechanisms, and via antibody-dependent cell-mediated cytotoxic reactions, by altering cellular function directly or after the internalization of membrane-bound antibodies. Patients with autoimmune gastritis may perhaps have, as discussed above, antibodies against the gastrin receptor, in addition to antibodies against the major parietal cell antigen. The extent to which H⁺,K⁺-ATPase antibodies and gastrin receptor antibodies occur simultaneously, and the separate roles of such antibodies in provoking inhibition of gastric acid production and cellular atrophy in patients with autoimmune gastritis, remains to be clarified.
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