Proteolytic Degradation of Exocrine and Serum Immunoglobulins

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Abstract The susceptibility of exocrine and serum immunoglobulins and antibodies to proteolytic degradation was assessed. Colostral and duodenal fluid exocrine 11S IgA, monomeric serum IgA, and IgG were digested with trypsin, chymotrypsin, or duodenal fluid. Exocrine IgA was more resistant to digestion than were the serum immunoglobulins. Under conditions of the experiments, most of colostral IgA retained its 11S quaternary structure, including the secretory piece; the portion degraded was reduced almost entirely to peptides.

The superior resistance of exocrine IgA was also demonstrated by digestion of serum IgG and nasal exocrine IgA diphtheria antitoxins with trypsin or duodenal fluid. Selective precipitation of trypsin-digested antibody molecules to heavy chains, light chains, or secretory piece revealed that the differences in susceptibility to digestion were due to differences in lability of the Fc portions of the IgA and IgG antibody molecules. The Fc portions of IgG antibody molecules were degraded or cleaved from the Fab units of the molecules, whereas the Fab-like portions of IgG antibody molecules remained associated with their Fab-like units and the secretory piece. On the other hand, trypsin treatment did not affect the antigen binding ability of the Fab parts of either the exocrine IgA or IgG antibodies.

The Fab-like portions of exocrine IgA may be protected from trypsin degradation by the quaternary structure of the 11S molecules, which includes a dimer of 7S IgA subunits and the secretory piece.

Introduction

Exocrine 11S IgA is the predominant immunoglobulin of most human external secretions, including intestinal fluid (2–4), colostrum, and milk (4, 5). Antiviral and antibacterial antibodies of the IgA class (6–9) have been identified in these fluids, suggesting that exocrine IgA plays a role in the immune defenses of the gut. Because immunoglobulins in gastrointestinal fluids are exposed to potent proteolytic enzymes, their susceptibility to proteolytic degradation may influence their biological activities.

Evidence concerning the stability of antibodies and immunoglobulins exposed to proteolytic enzymes is meager and inconclusive. The biological activity of certain milk antibodies resisted peptic digestion more effectively than did the activity of corresponding serum antibodies (9). The activities of other serum and exocrine antibodies were equally susceptible to trypsin (9, 10) or peptic (10) digestion. Tomasi and coworkers (11, 12) have stated that exocrine IgA resists digestion by several proteolytic enzymes which readily degrade serum immunoglobulins, but experimental evidence has not been published.

In this study, human exocrine and serum immunoglobulins and antibodies were digested with trypsin, chymotrypsin, or duodenal fluid. Susceptibility to digestion was assessed principally by the following two approaches: (a) digests of the immunoglobulins were fractionated by gel filtration, and the per cent of each immunoglobulin remaining intact was estimated from the spectrophotometric absorption curves or by radial immunodiffusion; and (b) diphtheria antitoxins prepared from nasal secretions or serum were digested, and the amounts of radio-labeled diphtheria toxoid bound by intact antibodies or by their fragments were estimated. The results indicated that exocrine IgA was more resistant to digestion than were IgG or serum IgA, and...
that IgA antitoxin was more resistant than serum IgG antitoxin.

METHODS

Purification of immunoglobulins and diphtheria antitoxins. Colostral 11S IgA, the secretory piece (SP) free in colostrum, and serum IgG were purified as described (13). Colostral IgA and SP were purified by diethyl aminomethyl (DEAE)-cellulose (Mannex-DEAE, Mann Research Labs, New York) and Sephadex (Pharmacia Fine Chemicals, New Market, N. J.) gel filtration from pooled, defatted colostrum which had been stored at −20°C. Monomeric serum IgA was prepared by a modification of the zinc sulfate method (14). The supernatant of the zinc sulfate at pH 7.8. The precipitate was twice filtered through a column of Sephadex G-200 (2.5 × 100 cm). Protein was eluted by upward-flowing borate-buffered saline (0.15 M NaCl in 0.01 M borate buffer, pH 8.0, containing 0.001 M EDTA). The final preparation contained less than 5% IgG and polymerized IgA.

Monomeric IgA was prepared from 11S colostral IgA by reduction with 50 mM dithioerythritol and alkylation with iodoacetamide as described (13). The 7S peak was isolated by Sephadex G-150 gel filtration.

Diphtheria antitoxins were prepared from serum or secretions of adults who had been immunized either by nasal instillation or subcutaneous injection of diphtheria toxoid (15). Exocrine IgA antitoxin was purified from nasal secretions by DEAE-cellulose column chromatography and Sephadex G-200 gel filtration. IgG antitoxin was isolated from serum by DEAE-cellulose column chromatography.

Enzymes and duodenal fluid. Twice-crystallized Type II1 bovine pancreatic trypsin (12,000 BAAE U/mg) was purchased from the Sigma Chemical Company, St. Louis, Mo. Bovine pancreatic alpha chymotrypsin (crystallized, salt-free, A grade, 1240 chymotrypsin neutral fraction [NF] U/mg) and soybean trypsin inhibitor (three-times crystallized, B grade) were purchased from Calbiochem, Los Angeles, Calif. Stock solutions of the enzymes (10 mg/ml in 0.001 N HCl) were stored at −20°C. Duodenal fluid was aspirated from fasting adults, snap frozen, and stored for a few days to a few weeks at −20°C before use. The subjects had various gastrointestinal disorders, but did not have pancreatic insufficiency; some were hypogammaglobulinemics.

Proteolytic activity of trypsin and duodenal fluid was estimated by casein digestion according to Kunitz (16). High protein casein was purchased from Calbiochem. Specific activity of the trypsin was 3.9 × 10^4 Kunitz units.

Tryptic hydrolysis of p-toluene sulphonyl-l-arginine methyl ester (TAME) was estimated by the method of Hummel (17) in Determinates TAME (Worthington Biochemical Corp., Freehold, N. J.).

Digestion of antigen and estimation of toxoid binding. Nasal fluid or serum antitoxin specimens were incubated with trypsin or duodenal fluid at a concentration of immunoglobulin (20 µg of N per ml or 0.13 mg of protein per ml) similar to that of IgA in normal duodenal fluid (2–4). Digestion was conducted with trypsin:substrate ratios of 1:23 or 1:10 (w/w) for 3 or 17 hr at 38°C in 0.05 M Tris-HCl buffer, pH 8.0. Trypsin digestion was stopped by a 4-fold excess of soybean trypsin inhibitor. The amount of 125I-labeled diphtheria toxoid bound by IgA or IgG antitoxin was then estimated as described (22). Soluble antigen-antibody complexes were formed by incubation (1 hr) of 0.2 ml of digested antitoxin preparation with excess radio-labeled toxoid. (Duodenal fluid digests were incubated at 4°C, trypsin digests at 38°C.) Excess rabbit antibody specific for immunoglobulin heavy chains, light chains, or SP then was added to the appropriate antitoxin mixture in order to coprecipitate toxoid bound only to intact antibodies or to Fab fragments as well as to intact antibodies. Anti-γ or anti-Fab antibodies were added to the IgG antitoxin mixtures; anti-α, anti-SP, or anti-Fab antibodies were added to the IgA antitoxin mixtures. Control (undigested) antitoxins were handled identically, except that duodenal fluid or a trypsin-trypsin inhibitor mixture was added to them just before incubation with toxoid. After further incubation of the antitoxin-toxoid mixtures (24 hr, 4°C), the precipitates were washed with cold borate-buffered saline, dissolved in 0.6 ml of 0.5 N NaOH, and the radioactivity was measured. The amount of toxoid bound by the precipitated immunoglobulin was estimated by comparing the sample's radioactivity with that of standard tubes containing a known amount of toxoid-125I. The amounts of labeled toxoid bound by enzyme-treated and untreated antitoxins were compared.

RESULTS

Digestion of exocrine and serum immunoglobulins. Colostral 11S IgA, serum IgA, or IgG (0.3–1.5 g/100 ml solutions) were incubated with trypsin, chymotrypsin, or duodenal fluid at 38°C for 8 hr in 0.05 M Tris-HCl buffer, pH 8.0. The ratio of trypsin or chymotrypsin to immunoglobulin was 1:23 (w/w). Duodenal fluid was diluted 2- to 5-fold in Tris-HCl buffer so that its proteolytic activity, measured by casein digestion, was similar to that of trypsin. Thus the proteolytic activity of the fluid was less and the immunoglobulin

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concentrations many times greater than those usually found in duodenal fluid (2-4). (Trypsin digestion was stopped by the addition of a 2-fold excess (w/w) of soybean trypsin inhibitor; chymotrypsin and duodenal fluid digestion was stopped by freezing.) Digestion mixtures were fractionated by Sephadex G-150 gel filtration (Fig. 1).

After digestion of colostrum IgA with trypsin or chymotrypsin, nearly all protein eluted as a single peak in the volume corresponding to that of untreated 11S IgA. Antigenic analysis demonstrated that the protein contained a-chain, light-chain, and SP antigenic determinants on a single molecule. A very small amount of small peptides also was eluted. Similar results were obtained when IgA preparations from two other colostrum pools were digested with trypsin. After digestion with duodenal fluid, colostral IgA yielded two additional small peaks. The first peak, a shoulder on the 11S
peak, eluted slightly earlier than IgG, and contained α-chain and light-chain determinants, but no SP determinants. Less than 3% of the recovered material was in this peak. The second small peak eluted slightly later than albumin and had only light-chain determinants. About 90% of the immunoglobulin which was degraded was degraded to peptides.

Trypsin, chymotrypsin, or duodenal fluid digests of serum IgA contained a major peak which eluted in the volume corresponding to that of the untreated protein and which contained α-chain and light-chain determinants. A much smaller peak, which eluted slightly later than albumin and contained only light-chain determinants, and considerable small peptide material also were produced.

All three enzyme preparations degraded most of IgG to fragments antigenically identical to Fab and Fc fragments of papain-digested IgG as well as to small peptides. The per cent of each immunoglobulin remaining intact after digestion was calculated from the gel filtration elution curves. The total area under each curve and the area attributable to undegraded protein were estimated separately; the area attributable to enzyme or duodenal fluid protein was excluded. 85–90% of colostral IgA was intact after digestion with trypsin or chymotrypsin, whereas only about 45–60% of serum IgA or IgG were intact (Fig. 2). Duodenal fluid more effectively degraded all three immunoglobulins, but colostral IgA was still most resistant. Serum IgA was slightly more resistant than IgG.

The stability of immunoglobulins in duodenal fluid was assessed further under conditions more closely simulating those within the duodenal lumen. Undiluted duodenal fluids containing physiologic concentrations (2, 3) of immunoglobulins were incubated at 38°C for 8 hr, and concentrations of the immunoglobulins before and after incubation were estimated by radial immunodiffusion. When duodenal fluids containing 0.30 and 0.44 mg/ml of IgA from two immunologically normal adults were incubated, the IgA concentrations decreased to 70 and 75%, respectively, of preincubation concentrations (Table I). (IgA in these unincubated fluids was proven to be the IIS exocrine variety by antigenic analysis and gel filtration separation.) When serum IgG was added to these fluids to achieve concentrations of 0.3 mg/ml and the mixtures incubated, IgG concentrations decreased to 18 and 27% of the preincubation values. Colostral IgA, serum IgA, or IgG (0.3 mg/ml) were incubated in duodenal fluids from two hypogammaglobulinemic patients. Digestion caused much less reduction in the concentration of colostral IgA than in the concentrations of serum IgA or IgG (Table I). Proteolytic activities of the fluids, measured by casein digestion, were equivalent to 0.5–2.5 mg of trypsin per ml.

The possibility was considered that IgA concentrations in the digests may have been overestimated because of the presence of large amounts of immunoglobulin fragments with α-chain determinants. An aliquot of the duodenal fluid-colostral IgA digest in which the IgA...
concentration was estimated by immunodiffusion to have decreased to 57% of the preincubation concentration (Table I) was separated on a column of Sephadex G-150. IgA in the eluted fractions was then estimated by the radial immunodiffusion system augmented by radiolabeled antibody. After digestion 51% of the immunoglobulin was recovered in the volume corresponding to that of undigested colostral IgA, and only about 10% of material with heavy-chain determinants was found in later-eluting fractions.

**Table I**

<table>
<thead>
<tr>
<th>Duodenal fluid*</th>
<th>Secretory IgA</th>
<th>Serum IgA</th>
<th>IgG</th>
</tr>
</thead>
<tbody>
<tr>
<td>% of preincubation concentration</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal</td>
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<td>—</td>
<td>18</td>
</tr>
<tr>
<td>—</td>
<td>70</td>
<td>27</td>
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<tr>
<td>γ-Globulin-deficient</td>
<td>57</td>
<td>14</td>
<td>11</td>
</tr>
<tr>
<td>—</td>
<td>79</td>
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<td>24</td>
</tr>
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</table>

* Colostral IgA, serum IgA, and IgG were added to the γ-globulin-deficient fluids; serum IgG was added to the normal fluids.

Digestion of reduced-alkylated colostral IgA and assessment of trypsin inhibitory activity of IgA. Monomeric 7S IgA isolated from reduced and alkylated 11S IgA was digested with trypsin at an enzyme:immunoglobulin ratio of 1:25 (w/w) in 0.05 M Tris-HCl buffer, pH 8.0. After 8 hr at 38°C, virtually all immunoglobulin had been degraded to small peptides or to fragments with light-chain determinants only.

In order to exclude the possibility that exocrine IgA or SP possesses trypsin inhibitory activity, trypsin hydrolysis of TAME was measured in the presence or absence of colostral IgA or free SP. Addition of a 25-fold excess (w/w) of either protein to the incubation mixture containing 0.7 μg of trypsin did not significantly decrease the rate of hydrolysis. A 4-fold excess of soybean trypsin inhibitor caused nearly complete inhibition.

**Diphtheria antitoxin digestion.** When either the serum IgG or nasal IgA antitoxin preparation was treated with trypsin and precipitated with anti-Fab antibodies, the amount of toxoid bound by precipitated antitoxin was approximately 100% of that bound in the untreated preparations (Table II). This finding indicated that the antigen binding sites in the antitoxin molecules had not been degraded by digestion. When anti-γ chain antibodies were added to the toxoid–IgG antitoxin mixture (17 hr incubation), only 19% of the bound toxoid was precipitated, whereas anti-α chain antibodies added to the IgA antitoxin digest coprecipitated 93% of IgA-bound toxoid, and 85% was precipitated by anti-SP antibodies.

The antitoxins were also digested for 3 hr in duodenal fluid having proteolytic activity equivalent to 1.3 mg of trypsin per ml (about 200 times greater activity than that of trypsin in the preceding experiment). After digestion, only 16% of toxoid bound by IgG antitoxin was coprecipitated by anti-γ chain antibodies, whereas 84% of toxoid bound to exocrine IgA antitoxin was coprecipitated by anti-α chain antibodies. The greater susceptibility of IgG antitoxin to digestion was also demonstrated in a 3 hr incubation with trypsin (Table II).

DISCUSSION

The studies consistently demonstrated that exocrine IgA was more resistant to proteolytic digestion than were IgG or serum IgA immunoglobulins or antibodies. The 11S quaternary structure of exocrine IgA evidently remained largely intact after digestion, as indicated by chromatographic fractionation and antigenic analysis of digested colostral IgA, and by the finding that the antigen combining ability of trypsin-digested nasal IgA antitoxin had not been dissociated from its α chains or SP. The superior stability of exocrine IgA even at physiologic concentrations in undiluted duodenal fluid was impressive. After a 3 hr incubation in fluids with potent proteolytic activity, more than 80% of IgA antitoxin was precipitated with anti-α chain antibodies, whereas only 16% of IgG antitoxin was precipitated by antibodies of γ chains. In immunodiffusion experiments, the concentration of exocrine IgA incubated in duodenal fluid decreased much less than did the concentrations of IgG and serum IgA. Because digested serum and exocrine IgA both yielded no or only small amounts of heavy chain-containing fragments, the immunodiffusion estimates of immunoglobulin concentrations probably accurately reflect the difference in stability between exocrine and serum IgA. On the other hand, estimates of IgG concentrations in digests probably were elevated modestly by the presence of Fe fragments. This artifact would have caused the amount of IgG degraded to be underestimated, and thus would not have influenced the conclusion that exocrine IgA is more resistant to duodenal fluid digestion than is IgG.

Studies using diphtheria antitoxins revealed that differences in resistance of exocrine IgA and IgG antibodies to trypsin were due to differences in lability of the Fe portions of the molecules. After a 17 hr digestion, exocrine IgA antitoxin was almost entirely precipitated by anti-α chain antibodies, whereas only a small fraction of IgG antitoxin was precipitated by anti-γ chain antibodies. These findings indicated that the Fe portions of most IgG antitoxin molecules had been degraded or cleaved, while the corresponding portions of most IgA...
antitoxin molecules were antigenically intact and attached to their Fab units. On the other hand, digested IgG and exocrine IgA antitoxins both completely retained toxoid binding ability that could be precipitated by antibodies to light chains, indicating that the Fab portions of both immunoglobulins remained intact and were equally resistant to digestion.

Why exocrine IgA resists tryptic hydrolysis is unknown, but the explanation may lie in the complex quaternary structure of the immunoglobulin, which consists of dimers of 7S IgA bound to SP by noncovalent and disulfide linkages acting in concert (5, 13). We did not find that γA monomers derived from reduced and alkylated IgA are unusually stable; however, because reduction and alkylation may have affected the lability of the 7S molecules, this evidence is not conclusive. The possibility that IgA proteins, which constitute a higher proportion of IgA in secretions than in serum (23, 24), are more resistant to proteolysis than are IgA proteins was not investigated. It seems unlikely, however, that the modest differences reported in IgA subclass concentrations could account for the large difference in resistance of serum and exocrine IgA to whole duodenal fluid digestion. The polymer fraction of serum IgA, held together exclusively by disulfide bonds, is reportedly susceptible to enzymatic hydrolysis (11), suggesting that SP may stabilize the exocrine IgA molecule. Although this possibility is attractive, the essential role of SP in maintaining the quaternary structure of exocrine IgA has not been established (nor was free SP or exocrine IgA effective in inhibiting trypsin). The finding that α-chain proteins from patients with intestinal lymphoproliferative diseases form polymers in vivo that are held together by disulfide bonds and noncovalent forces (25) suggests that the similar linkages of exocrine IgA may be sufficient for stabilization of the immunoglobulin. Since the α-chain disease protein polymers are more resistant to digestion with papain (25a) and pepsin than γA myeloma globulins are, the resistance of exocrine IgA to tryptic digestion may also be independent of the SP component.

Our findings are consistent with previous studies of tryptic degradation of immunoglobulins. Trypsin cleaves IgG principally into biologically-active Fab and Fc fragments (26, 27). Fab fragments of trypsin-digested IgG antibodies were found to retain antigen-binding ability (27). Papain degrades normal serum and myeloma IgA to 3.5S fragments corresponding to the Fab portion of IgG, but no fragments corresponding to Fc fragments are produced (28). Cederblad, Johansson, and Rymo (29) found colostral IgA difficult to degrade with trypsin until they added 0.01 μM cysteine. Considerable amounts of Fab-like fragments, but no Fc-like fragments, then were produced. The presence of cysteine, which also enhances tryptic digestion of γG globulins (26), may explain the greater degradation of colostral IgA in their studies than in ours. In contrast to our conclusions regarding tryptic digestion of immunoglobulins, Wilson and Williams (30) concluded that exocrine IgA is not especially resistant to peptic digestion, because pepsin-digested colostral IgA yielded an amount of peptides intermediate between amounts produced by two groups of IgA myeloma proteins. The relative susceptibility of serum and exocrine immunoglobulins to peptic digestion therefore may differ from their susceptibility to tryptic digestion, although the gel filtration data presented by Wilson and Williams actually suggest that less colostral IgA than either group of myeloma proteins was degraded to fragments smaller than the parent molecules, despite similarities in the amounts of peptides produced.

Whether secretory IgA enjoys biological advantages because of its resistance to digestion is uncertain, since in these studies the antigen-binding ability of both IgG and exocrine IgA antitoxin molecules were unaffected by tryptic digestion. On the other hand, the biological role of the Fe portion of IgA, which is protected in exocrine IgA, is unknown. Neither serum nor exocrine IgA fix complement after nonspecific aggregation (31-33) or in antigen-antibody complexes (34). An important role for the Fe portion of IgA probably will be established, nevertheless, because the corresponding portions of other immunoglobulins (IgG, IgM, and IgE) are biologically important. The biological significance of the resistance of secretory immunoglobulins to tryptic digestion deserves investigation.

**ACKNOWLEDGMENTS**

We are grateful to Mrs. Bonita L. DeVald for technical assistance.

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**Table II**

<table>
<thead>
<tr>
<th>Digestion</th>
<th>Antitoxin</th>
<th>Heavy chain</th>
<th>Fab</th>
<th>SP</th>
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<tbody>
<tr>
<td>Trypsin</td>
<td>17</td>
<td>IgG</td>
<td>19</td>
<td>102</td>
</tr>
<tr>
<td>E:S,* 1:23</td>
<td>1:10</td>
<td>IgA</td>
<td>93</td>
<td>103</td>
</tr>
<tr>
<td>Duodenal fluid</td>
<td>3</td>
<td>IgG</td>
<td>16</td>
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<tr>
<td></td>
<td></td>
<td>IgA</td>
<td>84</td>
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<tr>
<td>Trypsin</td>
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<td>IgG</td>
<td>52</td>
<td>103</td>
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<tr>
<td>E:S, 1:10</td>
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<td>IgA</td>
<td>89</td>
<td>97</td>
</tr>
</tbody>
</table>

*Enzyme:substrate (immunoglobulin) w/w.

1 Seligmann, M. Personal communication.
REFERENCES


