Human *Toxoplasma gondii*–specific Secretory Immunoglobulin A Reduces *T. gondii* Infection of Enterocytes In Vitro

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Abstract

Whey from 17 women (four acutely infected with *Toxoplasma gondii*, eight chronically infected, and five uninfected) was studied. *T. gondii*–specific secretory IgA antibodies were demonstrated by ELISA in whey from acutely infected and one of eight chronically infected women. Such antibodies to tachyzoite proteins of ≤14, 22, 26–28, 30, 46, 60, 70–80, and > 100 kD (eliminated by protease but not periodate or neuraminidase treatment) were demonstrated in whey from acutely infected subjects when Western blots were probed with their whey and antibodies to human secretory IgA or IgA or secretory piece. Secretory IgA from four of eight chronically infected women recognized the 46- and 69-kD epitopes. Other whey samples were negative. Incubation of *T. gondii* tachyzoites with whey or purified secretory IgA from acutely infected (but not seronegative) women caused 50–75% reduction in infection of enterocytes in vitro. Whey reactive with the 46-kD epitope from three of six chronically infected women caused less (≥40%) inhibition. Whey and purified secretory IgA from two of three acutely infected women agglutinated tachyzoites. Whey did not result in complement-dependent lysis of *T. gondii*. These results indicate that it may be possible to produce human secretory IgA to *T. gondii* capable of reducing initial infection of enterocytes, as such IgA is present during natural infection. They also demonstrate candidate epitopes for such protection. (*J. Clin. Invest. 1992; 90:2585–2592.*) Key words: *Toxoplasma gondii* infection • whey • secretory antibody • epitopes • agglutination • fab

Introduction

*Toxoplasma gondii* can cause death, systemic illness, and loss of sight, hearing, cognitive, or other neurological function when it is acquired congenitally or in immunocompromised individuals (1, 2). One strategy for development of protective preparations against organisms that initially infect via the intestine has been to stimulate intestinal, organism-specific secretory IgA (3–19). As *T. gondii* is initially acquired perorally (1), we studied whether humans produced such *T. gondii*–specific secretory IgA. Milk was used as the source of the secretory IgA. Epitopes recognized by such IgA then were characterized and the effect of this secretory IgA on infection of enterocytes in vitro was determined.

Methods

Collection of milk samples. Milk samples were obtained from 17 women. Serologic tests were performed by Dr. Jack Remington’s laboratory (Stanford University and Palo Alto Medical Research Institute, Palo Alto, CA) as described (1, 2). Serologic test results for these women are in Table I. Five were uninfected (i.e., *T. gondii* seronegative in the Sabin Feldman Dye test [negative]); seven were chronically infected (i.e., *T. gondii* IgG seropositive, but IgM seronegative in the double sandwich IgM ELISA [chronic]; one had been infected periconceptually although serologic tests retained an acute pattern at the time her milk was obtained (subacute); and four were acutely infected (i.e., *T. gondii* IgM seropositive and AC/HS (a differential agglutination test) acute and three of these were mothers of a congenitally infected infant [acute]). Milk samples were obtained according to National Institutes of Health guidelines concerning human subjects. Milk samples were collected by the women through manual expression and were immediately frozen at −20°C. Before analyses, samples were centrifuged at 10,000 g for 1 h to remove lipids. The clear middle layer was collected. Samples were divided into separate aliquots and stored at −135°C until analyzed. The studies described herein were performed within 1 yr of collection of the milk samples.

Preparation of *T. gondii* antigen for ELISA. Swiss Webster (SW) female mice (Harlan Sprague Dawley, Inc., Indianapolis, IN) that weighed 18–20 g were used. 2 × 10⁶ RH strain tachyzoites in 0.2 ml isotonic saline were injected intraperitoneally into Swiss Webster mice. 2 d later, 5 ml of isotonic PBS, pH 7.4, was injected into the peritoneal cavity and the ascites—PBS suspension containing the parasite was aspirated. This suspension was forced through a 27-gauge needle to disrupt infected peritoneal exudate cells and filtered through a funnel lined with glass wool. It was then passed through a polycarbonate membrane filter with a 3-μm pore diameter (Nuclepore Corp., Pleasanton, CA) to remove host cells. The filtered parasites were washed three times in PBS and centrifuged at 400 g for 15 min at 4°C and were resuspended in distilled water at a concentration of 5 × 10⁶/ml; they were then sonicated three times for 30 s each time, and centrifuged at 10,000 g for 15 min at 4°C to remove cellular debris. The concentration of the soluble *T. gondii* lysate antigen was determined by the Lowry method (20).

*T. gondii*–specific sIgA ELISA. Wells of polystyrene microtiter plates (Dynatech Laboratories, Inc., Chantilly, VA) were coated with 100 μl *T. gondii* lysate antigen (50 μg/ml) in 0.1 M carbonate buffer, pH 9.6. After an overnight incubation at 4°C, the plates were washed three times with 0.1 M PBS containing 0.5% Tween 20 (Sigma Chemical Co., St. Louis, MO), pH 7.2 (PBS-T), and postcoated with 4% bovine serum albumin, fraction V (Calbiochem Corp., La Jolla, CA) in PBS (PBS-B) for 1 h at 37°C. 100 μl of twofold serial dilutions of the whey samples in PBS-TB was added to each well. The plates were incubated for 1 h at 37°C followed by three washes in PBS-T. 100 μl of alkaline phosphatase–conjugated goat anti–human secretory IgA (Cappel Laboratories, West Chester, PA) diluted in PBS-TB was added to each well. The plates were incubated for 1 h at 37°C followed by three washes in PBS-T. 100 μl of nitrophenyl phosphate (1 mg/ml; Sigma Chemical Co.) diluted in 0.05 M carbonate buffer, pH 9.6, was added to each well followed by an incubation for 1 h at 37°C, and the
Table 1. Serologic Test Results for Women from Whom Whey was Obtained

<table>
<thead>
<tr>
<th>Status of infection</th>
<th>Subject number</th>
<th>Infant DOB</th>
<th>Estimate of trimester of acquisition</th>
<th>Data sera obtained</th>
<th>Date milk obtained</th>
<th>Reciprocal of serum dye test*</th>
<th>Serum IgM ELISA*</th>
<th>Serum IgA ELISA*</th>
<th>Serum AC/HS*</th>
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<td>11/90</td>
<td>Second</td>
<td>9/90</td>
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<td>8,000</td>
<td>7.7</td>
<td>9.4</td>
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<td>2</td>
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<td>Third</td>
<td>11/90</td>
<td>11/90</td>
<td>1,024</td>
<td>3.9</td>
<td>5.6</td>
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<tr>
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<td>3/89</td>
<td>Third</td>
<td>7/89</td>
<td>5/89</td>
<td>4,096</td>
<td>4.8</td>
<td>ND</td>
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<td>&gt;1,600/3,200</td>
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<td>8/90</td>
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<td>9/89</td>
<td>9/89</td>
<td>Negative</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>

* At the time milk sample was obtained; AC/HS refers to the "Differential agglutination" test of Desmonts, Thulliez, and Remington. This test uses acetone (AC) and formalin (HS)-fixed tachyzoites. It is useful in determining whether infection was acquired in the 6 mo before the time when the sample was taken. The results for the acutely and subacutely infected women demonstrate that their infections were recently acquired. § Treated with spiramycin during gestation. IgG by EIA, IgM by IFA.

absorbance was determined at 405 nm using an ELISA reader. Specific antibody titer was determined as the last dilution that yielded an absorbance two times that of the negative control.

**Total IgA ELISA.** The quantification of total IgA in whey using an ELISA was performed as described under **T. gondii**–specific IgA ELISA with modifications. Briefly, microtiter plates were coated with 100 µl anti-human secretory component (1 µg/ml; Sigma Chemical Co.) rather than **T. gondii** lyse antigen and fivefold serial dilutions of milk samples were used. Standard curves were constructed using known amounts of IgA (0.145–710 µg/ml). Correlations between log_{10} OD readings and log_{10} concentration of IgA were linear in this range and highly significant (R = 0.97).

**Isolation of IgA.** Milk samples were diluted 1:1 with PBS and applied to a 5.0 ml column of immobilized jacalin crosslinked to 6% beaded agarose (Pierce Chemical Co., Rockford, IL) (21). The column was washed with 25 ml of PBS. The IgA was eluted with PBS containing 0.1 M methylsulfoxide (Sigma Chemical Co.) and 1-mL fractions were collected. The peak fractions were pooled and the buffer was exchanged with PBS using a microconcentration system, molecular weight cutoff, 30,000 (Centricon 30; Amicon Corp., Danvers, MA). The concentrate was filtered using a 0.45-µm filter (Gelman Sciences Inc., Ann Arbor, MI) and then stored at −135°C.

**Maintenance of enterocytes.** Enterocytes (ATCC CRL 1592, IEC-6 [22]) were grown in 75-cm² flasks containing Dulbecco's minimal essential medium (DMEM) (Gibco Laboratories, Grand Island, NY), 10% FCS (which had been heated for 1 h at 56°C to inactivate complement) (HyClone Laboratories Inc., Logan, UT), containing 1.0% glutamine, 100 U/ml penicillin, 100 µg/ml streptomycin, and 2.5 µg/ml amphotericin B (DMEM, 10% FCS). They were split 1:5 for propagation or harvested for a challenge when they were confluent (~3–4 d after initiation of cultures).

**Preparation of Me49 tachyzoites for in vitro challenge.** Me49 tachyzoites were propagated in an L929 macrophage cell line (kindly provided by Dr. Frank Fitch, The University of Chicago, Chicago, IL) in 75-cm² flasks (Nunc, Roskilde, Denmark) containing DMEM, 10% FCS. L929 cells were split 1:10 (10^6) every 3–4 d. Free tachyzoites along with tachyzoite-infected cells from a 3–4-d infected monolayer were split 1:10 and 10^7 were added to 10^6 uninfected L929 cells to propagate the tachyzoites.

To prepare tachyzoites for in vitro challenge, infected monolayers were scraped with a rubber policeman to dislodge free tachyzoites and infected cells. The suspension was passed through a 27-gauge needle and a funnel lined with glass wool and was filtered free of host cells as described (23). The tachyzoites were centrifuged at 400 g at 20°C for 15 min. The pellet then was resuspended in 1 ml of Medium 199 (Gibco Laboratories) that contained 1.0% glutamine, 100 U/ml penicillin, and 100 µg/ml streptomycin (M199). The tachyzoites were counted in trypan blue, and the concentration was adjusted to 1.6 × 10^7/ml. 100-µl aliquots were dispensed into individual tubes containing 100 µl milk or IgA from either seropositive or seronegative control subjects and incubated at 37°C while rocking on a platform rocker (Belco Biotechnology, Vineland, NJ) at 10 rpm. Sera from uninfected and infected mice (23) were also incubated with tachyzoites as described above, as internal controls in each assay. After 30 min, the volumes were adjusted to 400 µl with M199 containing 10% FCS.

**In vitro challenge.** 100 µl of enterocytes (2 × 10^5/ml) were plated in each well of a microtiter plate (Flow Laboratories, Inc., McLean, VA) before challenge. After incubation for 1 to 3 h at 37°C, 5% CO₂, the monolayers were challenged with 100 µl of treated Me49 strain tachyzoites (4 × 10^8/ml). After 1 h, 25 µl [3H]uracil (50 Ci/mmol, 100 µCi/ml; Amersham Corp., Arlington Heights, IL) was added to each well followed by incubation for 20 h. Tachyzoites were harvested by alternating vigorous washing and aspirating with isotonic saline. They were collected on fiberglass filters with a multiple automated cell harvester (M.A. Bioproducts, Walkersville, MD [24]). Filters were dried and the tachyzoites were retained on each filter counted by liquid scintillation spectrophotometry to quantitate tachyzoite multiplication (23). Reduction in infection was expressed as the percent reduction of [3H]uracil incorporation compared with the seronegative whey control. To compare individual experiments, counts per minute of the

1. Abbreviation used in this paper: DMEM, Dulbecco's minimal essential medium.
negative control in separate cultures were normalized to 10,000 cpm. This was done by assigning a value of 10,000 to control wells in which enterocytes were challenged with T. gondii pretreated with media and FCS only (i.e., untreated controls). Thus, normalized counts per minute were calculated as follows: 10,000 × (cpm treated [e.g., with whey]/cpm untreated control).

In addition, enterocytes were cultured and challenged in four-chamber slides (LabTek; Nunc, Naperville, IL) as previously described for macrophages (23), and numbers of enterocytes per field (×400), percent infected cells, and mean number of tachyzoites per vacuole were assessed as previously described (23).

Tachyzoite uptake. To determine whether whey was toxic to the enterocytes and could result in fewer cells per culture well, 100 μl of enterocytes that contained 2 × 10^5 enterocytes was plated in each well. Also, seven serial twofold dilutions of enterocytes were plated in triplicate to produce a standard curve for comparison. This was done because DNA and protein assays could not be used to quantitate enterocytes as the cultures were not washed before processing and, thus, measurement of DNA or protein would also reflect detached cells in the culture supernatant. After incubation for 1 to 3 h at 37°C, 5% CO₂, and 100% humidity, 100 μl of diluted whey samples was added to the monolayers. After 1 h, 25 μl [3H]thymidine (50 Ci/mmol, 50 μCi/ml; Amersham Corp.) was added to each well followed by incubation for 20 h. Enterocytes were harvested by alternating vigorous washing and aspirating with isotonic saline. They were collected on fiberglass filters with a multiple automated cell harvester. Filters were dried and the DNA labeled with [3H]thymidine that was retained on each filter was counted by liquid scintillation spectrophotometry to estimate the number of enterocytes that were present and thus incorporating [3H]-thymidine.

Sabin Feldman dye test. This was performed as described (24).

Agglutination. This was performed using Formalin-fixed tachyzoites with the standardized direct agglutination test kit (Biomerieux, Marcy-L'Etoile, France) according to the manufacturer's instructions. Dilutions of whey and sera (dye test negative and positive for internal controls) were 1:40 and 1:4,000.

SDS-PAGE and electrophoretic transfer. RH tachyzoites were obtained from tissue culture. They were dissolved in SDS gel sample buffer (1% 2-mercaptoethanol [Sigma Chemical Co.], 500 mM Tris [Sigma Chemical Co.], 1% SDS, and 10% glycerol, pH 6.8). Approximately 2.5 × 10^7 tachyzoites (200 μg protein) per lane were loaded onto a 12% Laemmli gel and subjected to electrophoresis at 30 mA for 4 to 5 h (25). The T. gondii antigens resolved by SDS-PAGE were transferred to a polyvinylidene difluoride membrane (Millipore Corp., Bedford, MA) using a semidyry electroblotting apparatus (American Bionetics, Hayward, CA). The transfer buffer consisted of 25 mM Tris, 40 mM 6-amino-hexanoic acid, and 20% methanol, pH 9.4. A constant current of 2.5 mA/cm² was applied for 1 h at room temperature.

To determine whether antigens were polypeptides, lipids, or carbohydrates, T. gondii antigens were also treated with proteinase K (1 U/ml; Sigma Chemical Co.) for 1 h, neuraminidase (1 U/ml; Sigma Chemical Co.) for 4 h, or periodate (10 nM, Sigma Chemical Co.) for 30 min and blotted as described (25).

Western blot. Polyvinylidene difluoride membranes were blocked overnight at 4°C with 5% non-fat powdered milk in buffer (140 mM NaCl, 0.05% Tween 20, in 15 mM Tris, pH 8.0) (M-TNT). Strips were cut and incubated with whey samples diluted 1:1 with M-TNT for 1 h. After three washes with 5% M-TNT, strips were incubated for 1 h with horseradish peroxidase–conjugated anti–human secretory component (Sigma Chemical Co.) diluted 1:1,000 in M-TNT. Blots were also probed with anti–human secretory IgA (Cappel Laboratories) and anti–human IgA (Sigma Chemical Co.). Sera from uninfected and infected mice (23) probed with goat antibody to mouse immunoglobulins (Cappel Laboratories) were also included in each assay as internal controls. Strips were washed three times in PBS and developed using a PBS substrate solution that contained 0.5 mg/ml diaminobenzidine (Sigma Chemical Co.), 0.2 mg/ml CoCl₂ (Sigma Chemical Co.), and 0.03% H₂O₂ (Fisher Scientific, Fairlawn, NY) for 15 min. An estimation of the molecular weight of the reactive polypeptides was determined using pre-stained molecular weight standards (Amersham Corp.) as a reference.

Statistics. Within each experiment there were triplicate cultures for each treatment group. Each experiment was repeated two or more times. Results are expressed as mean±SD. To permit comparisons between experiments, counts per minute in individual experiments were normalized to 10,000 cpm (i.e., [cpm treated/cpm untreated] × 10,000). Effects of treatment of tachyzoites in the enterocyte challenge were analyzed using the Neuman-Keuls multiple-range test. Correlation coefficients were determined by regression analysis. Differences were considered significant when P < 0.05.

Results

Secretory IgA ELISA. Whey from three of three acutely infected mothers contained T. gondii–specific secretory IgA when tested in ELISAs (Fig. 1). ELISA values with whey from the chronically infected and seronegative women were substantially less (Fig. 1). There was no definitive association between concentrations of total IgA and presence of T. gondii–specific secretory IgA demonstrated by ELISA (Fig. 1). Specifically, data in Fig. 1 indicate that two women had whey with higher amounts of total IgA and specific antibody to T. gondii and two women had whey with higher amounts of total IgA and no specific antibody to T. gondii. Two of four samples with substantial total IgA had substantial specific IgA antibody to T.
gondii, and 1 of 12 samples without substantial total IgA had substantial specific IgA antibody to *T. gondii*.

*Western blots.* Epitopes recognized by this secretory IgA were defined by Western blot (Figs. 2 and 3 and summarized in Table II). Whey was demonstrated to contain some complete secretory IgA for the first patients identified, as it was detected when antibody to secretory piece and IgA were used as well as antibody to human secretory IgA (Fig. 2). Treatment of nitrocellulose strips of electrophoresed *Toxoplasma* antigens with periodate (10 nM) for 30 min, neuraminidase (1 U/ml) for 4 h, or protease (protease K) (1 U/ml) for 1 h demonstrated that all these antibodies were directed against proteins.

*Enterocyte challenge.* An in vitro enterocyte challenge assay was developed to determine whether these antibodies had potential biological significance in protection against *T. gondii*. Me49 strain tachyzoites were incubated with whey, and in subsequent experiments with jacalin-purified secretory IgA subclass 1. Whey from the initially studied acutely infected mother, and to a lesser extent whey from the initially studied chronically infected mother, inhibited replication after in vitro *T. gondii* challenge in a dose-dependent manner (Fig. 4). Replication was measured as tritiated uracil uptake (Fig. 4). These results were confirmed when whey from all 16 women were studied simultaneously (Fig. 5 A and B). Results were the same whether data are expressed as normalized data from replicate experiments (Fig. 5 A) or counts per minute for a single representative experiment (Fig. 5 B).

IgA1 was purified using a jacalin affinity column. This IgA1 from the acutely infected mother, but not the seronegative mother, also inhibited *T. gondii*. Data from a representative experiment are in Fig. 6.

Whey was not toxic to the enterocytes. There was no difference in thymidine uptake of enterocytes treated with media,
Table II. Correlation of Presence of T. gondii–specific Secretory IgA in ELISA, Epitopes Recognized in Western Blots, and Inhibitory Effect of Whey on Subsequent Replication of T. gondii in Enterocytes

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<thead>
<tr>
<th>Molecular mass of epitope</th>
<th>Acute</th>
<th>Subacute/chronic</th>
<th>Uninfected</th>
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<tr>
<td></td>
<td>1*</td>
<td>2</td>
<td>3*</td>
</tr>
<tr>
<td>&gt; 100</td>
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<td>+</td>
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<td>≤ 14</td>
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T. gondii–specific ELISA results (OD)  917  642  266  ND  25  64  78  69  134  115  61  1  111  42  -9  65  58

Percent inhibition* > 75  50  50  ND  0  50  40  40  15  15  0  0  0  0  0  0  0

++, strongly positive; +, present; ±, equivocal. * Subject number. † This whey sample also contained secretory IgA that recognized epitopes between 14 and 22 kD. ‡ Treated with Spiramycin during gestation. ‡* Whey from this seronegative mother had nonspecific reactivity in some assays. ‡† Percent inhibition, percent inhibition of T. gondii infection of enterocytes due to whey. Inhibition appeared to correlate with presence of T. gondii secretory IgA to epitopes ≤ 46 kD demonstrated in Western blots.

Whey from the seronegative mother, or whey from the acutely infected mother. This assay detected less than twofold differences in enterocyte numbers accurately. Numbers of cells per high-power field evaluated by light microscopy were not altered by incubation with the whey sample.

Microscopic analysis revealed that whey diminished infection of enterocytes by T. gondii but not subsequent replication of T. gondii. Specifically, in a representative experiment, in cultures challenged with T. gondii treated with whey from an acutely infected mother versus those treated with whey from an uninfected mother, there were 21 versus 40% infected enterocytes at 1 h after challenge and 26 versus 62% infected enterocytes at 18 h after challenge. Whey did not alter the mean number of tachyzoites per vacuole (≥ 4 vs. ≥ 4).

Sabin Feldman dye test. Human whey did not result in complement-dependent lysis of T. gondii.

Direct agglutination. Whey and jacalin affinity–purified secretory IgA from the two most recently infected mothers (subjects 1 and 2) agglutinated T. gondii tachyzoites. Whey from the other acutely infected mother (subject 3), who also was untreated, did not agglutinate T. gondii tachyzoites. This suggests that one mechanism of action of secretory IgA, but not the only one, could be agglutination of T. gondii.

Discussion

In earlier studies, we demonstrated that mice have intestinal IgA specific for T. gondii (26). Whey from uninfected, chronically, and acutely infected mothers was studied to determine whether humans also have T. gondii–specific secretory IgA and to obtain larger quantities of antibodies to facilitate characterization of such secretory IgA. The present studies demonstrate that T. gondii–infected humans produce secretory IgA specific to T. gondii. These antibodies recognize T. gondii proteins of ~ 14, 22, 26–28, 30–35, 46, 60, 70–80 and > 100 kD. Although SAG1 (p30) appeared to be the major epitope identified by whey from acutely infected versus chronically infected women, there was not an absolute correlation between inhibitory activity on T. gondii replication within enterocytes and presence of secretory antibody to SAG1 (Table II).

To simulate the natural route of infection, i.e., oral ingestion of T. gondii followed by subsequent invasion of the intestinal mucosa, an enterocyte challenge assay was developed.

Figure 4. Effect of incubation of Me49 strain T. gondii tachyzoites with whey on their subsequent replication in enterocytes in vitro. No Ab, refers to effect of media alone; no, effect of sera from seronegative mice or whey from a seronegative human mother; no, effect of sera from chronically infected mice or milk from the chronically infected mother; and no, effect of whey from the acutely infected mother. Differences between milk from uninfected and infected individuals were significant. ENT refers to counts per minute from enterocytes in culture alone; Me49 refers to counts per minute from the Me49 strain of T. gondii in culture alone.
Figure 5. Effect of treatment (incubation) of tachyzoites with whey from 16 women on Me49 strain T. gondii infection of enterocytes in vitro. (A) Normalized data pooled from at least two representative experiments. (B) Representative experiments (cpm). Numbers are the same as used in other Tables and Figures to indicate whey from individual subjects. Differences between whey from uninfected and infected individuals were significant. o, Untreated, refers to enterocytes challenged with Toxoplasma treated and cultured with media alone. • M+, treatment with sera from infected mice; □ M−, sera from uninfected mice; ●, treatment with whey from acutely infected woman; ■, treatment with whey from subacutely infected woman; †, treatment with whey from chronically infected woman; ◊, treatment with whey from uninfected woman. ENT refers to counts per minute from enterocytes in culture alone; Me49 refers to counts per minute from the Me49 strain of T. gondii in culture alone.

Figure 6. Effect of milk, secretory IgA isolated from whey using a jacalin-affinity column, and mouse sera on the subsequent replication of Me49 strain T. gondii tachyzoites in enterocytes challenged in vitro. •, Effect of media alone; ○, effect of sera from the seronegative mice or human whey or secretory IgA isolated from this whey (whey, jacalin) from the seronegative mother; ◊, effect of serum from the chronically infected mouse, and ●, effect of whey, or secretory IgA isolated from this milk (whey, jacalin), from the acutely infected woman. Dilutions of mouse sera were 1:10 and whey were 1:2. The sera from the infected mouse and milk and IgA from the infected individual significantly reduced counts per minute.

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The Me49 strain of *T. gondii* was adapted from its murine host to a tissue culture propagation system. The now rapidly dividing tachyzoites were used to infect rat enterocytes (22) in vitro. Quantitation of infection was assessed by incorporation of $[^3H]$uracil into multiplying tachyzoites (23). Enterocytes lack the salvage enzyme, uracil phosphoribosyltransferase, so that $[^3H]$uracil incorporation is confined to the parasite. Although possibly introducing a degree of artificiality to the assay, we selected a nonmalignant rat small intestinal cell line. At the time of these studies, the only human intestinal cell line available was a malignant colonic line. A nonmalignant, well-differentiated human small intestinal cell line has now been propagated. With the rat enterocyte assay system, we found that incubation of tachyzoites with either human whey or purified secretory IgA 1 subtype from this whey (of the acutely, some of the chronically, but not of the uninfected mother[s]) reduced subsequent intracellular replication of *T. gondii*. It will be of interest to determine whether human secretory IgA also blocks uptake into human cells.

Secretory IgA may protect mucosal surfaces by prevention of attachment to or colonization of mucosal surfaces and entry into susceptible cells. In other systems, (e.g., as demonstrated for reo and influenza viruses) (8–10), but not studied in our experiments, this is by sterically blocking the association of an organism with cellular receptors. Our experiments indicate that *T. gondii*-specific IgA in whey does not inhibit invasion by *T. gondii* through complement-dependent lysis. This is consistent with the known low efficiency of IgA for complement-dependent lysis. Two of three inhibitory whey samples agglutinated tachyzoites. Thus, agglutination could be one mechanism of inhibition, but not the only one. In separate studies (26a, Mineo, J. R., D. Mack, I. Kahn, K. Ely, J. Smith, R. McLeod, and L. Kasper, manuscript submitted for publication), we and others have demonstrated that polyclonal, monospecific, and monoclonal antibodies to SAG1 (p30) inhibit infection within enterocytes, fibroblasts, and MDBK cells (Mineo, J. R., D. Mack, I. Kahn, K. Ely, J. Smith, R. McLeod, and L. Kasper, manuscript submitted for publication). Also, Fab's of the polyclonal, monospecific antibody to SAG1 block infection of fibroblasts by wild-type *T. gondii* tachyzoites that have surface SAG1 but not by mutant tachyzoites that do not have SAG1 (Mineo, J. R., D. Mack, I. Kahn, K. Ely, J. Smith, R. McLeod, and L. Kasper, manuscript submitted for publication). These studies also demonstrate that SAG1 participates in *T. gondii*'s invasiveness through interaction with a host cell glycosylated molecules (Mineo, J. R., D. Mack, I. Kahn, K. Ely, J. Smith, R. McLeod, and L. Kasper, manuscript submitted for publication). Antibody to the major 30-kD epitope (SAG1) is present in three of three inhibitory whey samples from acutely infected mothers (demonstrated in Figs. 2 and 3, summarized in Table II). Dot blot with affinity-purified SAG1 confirmed that this antibody in whey was to SAG1 (our unpublished data). Thus, agglutination is one mechanism whereby subsequent replication of *T. gondii* could be inhibited. Other mechanisms also are operative because the Fab portion of antibody to p30 blocks infection of host cells, a host cell receptor for SAG1 has been partially characterized (Mineo, J. R., D. Mack, I. Kahn, K. Ely, J. Smith, R. McLeod, and L. Kasper, manuscript submitted for publication), and we have demonstrated that antibody in human whey recognizes this epitope.

For studies of the initial samples (Fig. 2), antiserum to secretory IgA, IgG, and secretory piece were used to demonstrate that intact secretory IgA was reactive rather than just IgA monomers. Antibodies to IgA1 and IgA2 were not further differentiated by Western blotting. As jacaoin isolates only subtype IgA1, there may be other substances in whey (e.g., IgA2 or IgG) from infected women that contribute to differences in uracil incorporation.

As we have found that human whey contains a *T. gondii*-specific secretory IgA antibody that can reduce infection in enterocytes in vitro, it may be possible to stimulate such an antibody in vivo. This could then contribute to protection against peroral acquisition of *T. gondii*. There are a number of other examples of pathogen-specific IgA in milk (17, 18) and protection provided by slgA or milk that presumably contains such IgA (3–7, 16, 18), as well as protection against certain pathogens (e.g., polio virus) conferred by oral immunization. The studies described herein, as well as work with other microorganisms (3–7, 16–18), provide a rationale for future studies to determine whether such *T. gondii*-specific IgA antibody also protects in vivo. They provide a basis to further characterize the *T. gondii* epitope(s) that stimulate(s) such protective antibody and to determine whether intestinal immunization (27) can protect against acquisition of *T. gondii*.

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