Reconstituted Human Polyclonal Plasma-derived Secretory-like IgM and IgA Maintain the Barrier Function of Epithelial Cells Infected with an Enteropathogen*

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Background: SigM and SigA are emerging as therapeutic agents, but their mode of action requires further definition.

Results: We provide a functional demonstration of SigM and SigA in maintaining the integrity of intestinal epithelial cells exposed to Shigella flexneri.

Conclusion: SigM is superior to SigA in ensuring the homeostasis of intestinal epithelial cells challenged with Shigella flexneri.

Significance: SigM and SigA have potential as mucosal therapeutic agents.

Intravenous administration of polyclonal and monoclonal antibodies has proven to be a clinically valid approach in the treatment, or at least relief, of many acute and chronic pathologies, such as infection, immunodeficiency, and a broad range of autoimmune conditions. Plasma-derived IgG or recombinant IgG are most frequently used for intravenous or subcutaneous administration, whereas a few IgM-based products are available as well. We have established recently that secretory-like IgA and IgM can be produced upon association of plasma-derived polymeric IgA and IgM with a recombinant secretory component. As a next step toward potential future mucosal administration, we sought to unravel the mechanisms by which these secretory Iggs protect epithelial cells located at the interface between the environment and the inside of the body. By using polarized epithelial Caco-2 cell monolayers and Shigella flexneri as a model enteropathogen, we found that polyspecific plasma-derived SigA and SigM fulfill many protective functions, including dose-dependent recognition of the antigen via formation of aggregated immune complexes, reduction of bacterial infectivity, maintenance of epithelial cell integrity, and inhibition of proinflammatory cytokine/chemokine production by epithelial cells. In this in vitro model devoid of other cellular or molecular interfering partners, IgM and secretory IgM showed stronger bacterial neutralization than secretory IgA. Together, these data suggest that mucosally delivered antibody preparations may be most effective when combining both secretory-like IgA and IgM, which, together, play a crucial role in preserving several levels of epithelial cell integrity.

Among these, specific humoral mucosal immunity is dominated by secretory antibodies (Abs): secretory immunoglobulin A (SigA) and secretory immunoglobulin M (SigM). Both secretory Abs result from the transport across the epithelium of J chain-containing polymeric IgA (mostly dimeric) and pentameric IgM by the polymeric immunoglobulin receptor and the polymeric immunoglobulin F.

In vitro and in vivo studies have established the potential of specific, antigen-induced IgM in the systemic neutralization of viruses (8–10), bacteria (11–13), fungi (14), and parasites (15–17). Important advances have come especially from the use of SigM-deficient mice (18), which exhibited a high sensitivity to bacterial and viral infections (19), a condition that could be partly controlled upon administration of normal mouse immune serum (8).

Immunotherapy on the basis of the passive administration of human plasma-derived IgG has been used for three decades in clinical applications, with improvement of a large panel of disease conditions like immunodeficiencies, infections, or autoimmune diseases (20, 21). Preclinical and clinical studies have underscored the efficacy against various infectious agents of polyclonal IgM-enriched preparations administered by the systemic route (22–26). Similar to SigA, SigM can be seen as a valid candidate immunoglobulin for mucosal application, given the

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its ability to bind antigens with strong avidity, its potential to ensure long term protection (27), its capacity to survive low pH conditions (28), as well as its resistance to proteases (29). We have demonstrated recently that human plasma can serve as a source of polyreactive, polymeric IgA (pIgA) and IgM to generate secretory-like IgA and IgM Abs, the natural molecular form found in secretions. We found that plasma-derived purified pIgA and IgM can associate with a recombinant secretory component (SC) with a 1:1 stoichiometry and that this association delayed the degradation of pIgA or IgM by intestinal washes containing proteases. In addition to these essential biochemical features, we showed that pIgA and secretory-like IgA delayed damage to epithelial polarized Caco-2 cell monolayers induced by a virulent strain of enteropathogenic Shigella flexneri (29). However, how the plasma-derived Ab operates to block the bacteria and contributes to epithelial homeostasis was not addressed in this study.

To provide answers to these open questions, we here dissect the mechanisms of protection conferred by plasma-derived pIgA and secretory-like IgA and then extend this study by evaluating the functionality of human plasma IgM and secretory-like IgM in the same experimental setting. We found that IgM or secretory-like IgM demonstrates a superior ability to maintain transepithelial electrical resistance (TER) and to forestall damage of cell monolayers resulting from S. flexneri infection when compared with pIgA or secretory-like IgA. Bacterial aggregates formed with both plasma pIgA and secretory-like IgA. This phenomenon was amplified upon association with IgM and secretory-like IgM, consistent with the capacity of all polyreactive Ab molecules to recognize S. flexneri. The diminished intracellular bacterial load varied as a function of the Ab isotype and resulted in differential production of proinflammatory mediators by the Caco-2 cell monolayers. In addition, incubation of secretory-like IgA and IgM resulted in reduced secretion of the S. flexneri virulence factors invasion plasmid antigen (Ipa) B and IpaC, overall suggesting a dual mode of action of the Abs, combining disabling of the bacteria and shielding of the target epithelium.

**EXPERIMENTAL PROCEDURES**

Preparation of Human Plasma IgA-, IgM- and IgG-enriched Fractions—IgA and IgM were purified from process intermediates of immunoglobulins manufactured from human plasma (30) by affinity chromatography using CaptureSelect human IgA and CaptureSelect human IgM resins (Bioaffinity Co.). The starting material used was a chromatographic side fraction consisting of the strip fraction from an ion exchange chromatography column used in the large scale manufacture of IgG from human plasma. The starting material was diluted in PBS to a target protein (IgA or IgM) concentration of ~1 mg/ml and then loaded onto a CaptureSelect human IgA or IgM column pre-equilibrated with PBS without exceeding the IgA- or IgM-binding capacity of the column. After loading, the column was washed with PBS, and IgA or IgM was eluted with glycine buffer at pH 3.0. The eluate was adjusted with 0.5 M Tris base (pH 8.0) to pH 4.5 and concentrated up to 16 mg/ml protein. Human plasma IgG preparations (IgPro10, Privigen) were prepared as described previously (30). Monoclonal IgAC5 specific for S. flexneri LPS serotype 5a was produced via hybridoma cells as described previously (31).

Separation of Plasma-derived pIgA and Monomeric IgA (mIgA) and Purification of Plasma-derived IgM—IgA-enriched preparations containing a mixture of mIgA and pIgA were diluted in PBS to a final volume of 10 ml, suitable for injection onto the AKTAPrime chromatography system (GE Healthcare). The flow rate was set at 1 ml/min, with PBS as mobile phase for all runs. Separation of the two molecular forms of IgA was performed on two serially coupled 1–m-long columns filled with Sephacryl S–300 high resolution (HR) beads (32). The IgA content of 3.5-ml fractions was verified by immunodetection using polyclonal rabbit anti-human IgA/HRP (1/3000, Dako), and pools of mIgA and pIgA were obtained. IgM-enriched preparations run under identical conditions yielded a single fully excluded peak. The IgM content of 3.5-ml fractions was verified by immunodetection using polyclonal rabbit anti-human IgM/HRP (1/3000, Dako). Concentration was performed using the Labscale system (Millipore) connected to a 100-kDa cutoff cartridge and stored at 4 °C until further use.

In Vitro Association of Polymeric Ig and Free SC—Recombinant human SC was produced from a CHO clone stably transfected with an expression cassette coding for the protein (33). Plasma-derived SlgA molecules were obtained by combining, in vitro, 10 μg of purified pIgA or IgM molecules with 2.5 or 1.5 μg of recombinant human SC, respectively. Mouse monoclonal SlgAC5 was obtained by combining, in vitro, 10 μg of purified pIgAC5 molecules with 2.5 μg of recombinant mouse SC. Association and characterization of SlgA and SlgM Abs were performed in PBS for 30 min at room temperature as described previously (29).

Caco-2 Cell Culture and Growth as a Polarized Monolayer—The human colonic adenocarcinoma epithelial Caco-2 cells (ATCC) were grown in complete DMEM consisting of DMEM-Glutamax (Invitrogen) supplemented with 10% FCS (Sigma), 10 mM HEPES (Invitrogen), 1% nonessential amino acids (Invitrogen), 1% sodium pyruvate (Invitrogen), 1% l-glutamine (Sigma), 1% penicillin/streptomycin (Sigma), and 0.1% transferrin (Invitrogen) and used between passages 32 and 40. Cells cultivated up to 80% confluency were seeded on polyester Snapwell filters (diameter, 12 mm; pore size, 0.4 μm; Corning Costar) at a density of 0.8 × 10⁵ cells/cm². At week 3, the Caco-2 cell monolayer integrity was checked by measuring the TER using the Millicell-electrical resistance system (ERS) device (Millipore) (34). TER values of well differentiated monolayers were in the range of 400–500 ohm/cm².

Bacterial Strain and Culture Conditions—Bacteria used were the virulent strain of serotype 5a LPS S. flexneri M90T constitutively expressing GFP (35). Bacteria from frozen stock were grown on a Luria-Bertani (LB) agar plate containing 0.1% Congo red (Applichem) and 50 μg/ml ampicillin (Sigma-Aldrich) for 30 h at 37 °C. Three red colonies were amplified in 10 ml of LB liquid broth supplemented with 50 μg/ml ampicillin for 16 h at 37 °C and 200 rpm. The culture was centrifuged at 2000 × g for 5 min, resuspended in PBS, diluted 1:10 in 10 ml of LB liquid broth/ampicillin, and then cultured for 2 h at 37 °C with shaking (200 rpm). Finally, bacteria in the exponential phase were washed twice in PBS by centrifugation at 2000 × g
for 5 min and resuspended in PBS. Assessment of cfu/ml was carried out by measurement of the optical density at 600 nm with the knowledge that 1 optical density unit at 600 nm corresponds to 5 × 10^6 cfu/ml.

**Incubation of Bacteria with Different Ab Preparations**—2 × 10^7 bacteria were mixed with 0.049 μM SlgAC5 specific for *S. flexneri* LPS serotype 5a or with human plasma-derived plgA (0.61 μM), reconstituted SlgA (0.61 μM), mlgA (0.61 μM), IgM (0.61 μM), reconstituted SlgM (0.61 μM), or IgG (0.61 μM). All mixtures were prepared in a final volume of 500 μl of plain DMEM (DMEM complemented with 10 mM HEPES, 1% non-essential amino acids, 1% sodium pyruvate, 1% L-glutamine, and 0.1% transferrin). The mixtures were incubated for 1 h at room temperature under gentle agitation.

**Protection Assay**—One hour before the use of polarized Caco-2 cell monolayers, complete DMEM was replaced by plain DMEM in both the apical and basolateral compartments. Polarized Caco-2 cell monolayers were infected apically with *S. flexneri* serotype 5a alone or in combination with the Ab preparations. Exposure of Caco-2 cells to antigens or the various immune complexes was performed overnight (O/N), and pathogen-induced damage was tracked by measuring TER decrease over time.

**Counting of Associated Bacteria with Cell Monolayers**—To count internalized bacteria, cells on Snapwell filters were washed with PBS and incubated for 30 min with 50 μg/ml gentamicin, followed by incubation in 500 μl of cold lysis buffer (10 mM Tris-HCl (pH 7), 0.2% Nonidet P-40, 50 mM NaCl, and 2 mM EDTA (pH 8)) for 5 min on ice and lysis by up-and-down pipetting. *S. flexneri* present in cell lysates were counted from serial dilutions seeded onto LB agar plates. In control experiments, we checked that Ab-mediated aggregation did not result in an artifactual decrease in bacterial numbers, as determined by plating of *S. flexneri* alone or in complex with SlgA or SlgM (data not shown).

**Laser-scanning Confocal Microscopy (LSCM) Observation of Caco-2 Cell Monolayers**—To examine the integrity of Caco-2 cell monolayers, Snapwells were washed twice with PBS prior to fixation O/N with 5 ml of 4% paraformaldehyde at 4 °C. After washing, filters were permeabilized, and nonspecific binding sites were blocked with PBS containing 5% FCS and 0.2% Triton X-100 for 30 min at room temperature. All samples and Ab dilutions were performed in PBS/0.05% Tween 20 containing 0.1% BSA. Washing reactions were stopped with 1M H2SO4. Absorbance was read at 490 nm with 630 nm as a reference.

**Measurement of Virulence Factors IpaB and IpaC Secreted by *S. flexneri***—To examine the impact of Abs on the expression of virulence factors, the immune complexes bacteria-Abs were incubated in wells for 2 h at room temperature, washed with PBS/0.05% Tween 20, and then detection was performed by incubation with isotype-specific Abs (mouse anti-human IgA1/IgA2 biotinylated IgG (BD Biosciences, 1:1000), goat anti-human IgA2 biotinylated IgG (BD Biosciences, 1:1000), goat anti-human IgM to bacteria, 96-well plates (MaxiSorp, Nunc) were coated with 4 × 10^7 cfu/well of *S. flexneri* serotype 5a in PBS O/N at 4 °C. After three washes with PBS/0.05% Tween 20, wells were blocked PBS/0.05% Tween 20 containing 1% BSA (Fluka) for 1 h at room temperature. Serial dilutions of human plasma IgA, SlgA, IgM, SlgM, IgG, or mouse SlgAC5 (from 0.61 μM) were incubated in wells for 2 h at room temperature, washed with PBS/0.05% Tween 20, and then detection was performed by incubation with isotype-specific Abs (mouse anti-human IgA1/IgA2 biotinylated IgG (BD Biosciences, 1:1000), goat anti-human μ chain biotinylated IgG (KPL, 1:1000), goat anti-human γ chain biotinylated IgG (Sigma, 1:1000), or goat anti-mouse α chain biotinylated IgG (KPL, 1:1000 dilution) for 2 h at room temperature followed by Extravidin-HRP (Sigma, 1:5000 dilution) for 1 h at room temperature. All samples and Ab dilutions were performed in PBS/0.05% Tween 20 containing 0.1% BSA. Finally, detection was performed with citrate/phosphate solution (44.4 mM citric acid, 103 mM Na2HPO4 (pH 5.0)) containing 1 mg/ml O-phenylenediamine (Sigma) and 0.01% H2O2. The reactions were stopped with 1 M H2SO4. Absorbance was read at 490 nm with 630 nm as a reference.
**Homeostatic Properties of Secretory-like Plasma IgA and IgM**

**Statistical Analysis**—Results are expressed as mean ± S.E. of n determinations. Differences in TER, cytokine secretion, number of foci, infected areas, and secretion of IpaB and IpaC between the control group (Sf alone) and in the presence of Abs were tested by analysis of variance (parametric, two-tailed). Differences were considered as significant when p < 0.05. Graph generation and statistical analyses were performed using GraphPad Prism software version 6.

**RESULTS**

**Human Plasma plgA and SlgA Abs Agglutinate Bacteria Lead to a Decrease of the Bacterial Load in Caco-2 Cell Monolayers and of Proinflammatory Cytokines/Chemokines**—We demonstrated previously that human plasma-derived plgA and reconstituted SlgA, but not plasma mlgA, significantly maintained Caco-2 cell intestinal epithelial monolayer integrity after O/N infection with *S. flexneri* by preventing the reduction of TER, the disruption of the tight junction network, and the detachment of filter-bound Caco-2 cells (29). However, the underlying mechanisms explaining protection were not addressed. To better define the mode of action of human plasma plgA and SlgA, we first focused on the nature of the interaction between the bacteria and the Abs. The binding capacity of the various molecular forms of plasma IgA to *S. flexneri* was compared by ELISA. All molecular forms of IgA demonstrated a concentration-dependent ability to bind bacteria. At identical concentrations, the signal for plgA or SlgA was 3- to 4-fold stronger than the signal observed with mlgA (Fig. 1A), emphasizing the avidity effect associated with polymeric Abs. To gain further insight into the nature of the interaction, immune complexes between cyanine 5-labeled Abs and GFP-expressing bacteria were formed and visualized directly by LSCM. Specific SlgAC5 and plasma-derived IgG were assessed for comparison. All molecular forms bound to *S. flexneri*, but bacterial aggregates of increasing size formed upon association with human plasma plgA and SlgA, demonstrating that only polymeric IgA was capable of engaging bacteria in complex lattices (Fig. 1B). Interestingly, the large aggregate pattern resembles that formed with *S. flexneri* LPS-specific SlgAC5 (31).

We next examined whether sequestering bacteria in immune complexes would influence the number of infecting bacteria found in Caco-2 cells. Although specific SlgAC5 monoclonal Abs were most effective at blocking bacterial internalization, we found that the presence of either of the plasma-derived Abs reduced the number of internalized bacteria 2-fold on average (Fig. 2A). The effect has to be considered as a trend only because statistical significance was not reached. In the same experimental setting, we then studied the effect of the molecular forms of plasma IgA on the inflammatory response of infected cell monolayers. In the presence of plgA or SlgA, basolateral secretion of TNF-α, CCL3, and CXCL8 was less than half the level of production measured upon infection with *S. flexneri* alone (Fig. 2B). This inhibition of cytokine/chemokine production was in the same range as that obtained with specific SlgAC5, whereas mlgA and (monomeric) IgG had no effect on the secretion of the three proinflammatory mediators (Fig. 2B). We conclude that the various molecular forms of plasma IgA and plasma IgG directly bind to bacteria but with distinct consequences on

![FIGURE 1. Association of human plasma-derived IgA/SlgA with *S. flexneri*.](image)

**A**—binding of equimolar concentrations of plgA, reconstituted SlgA, or mlgA to immobilized *S. flexneri* as determined by ELISA. Successive dilutions of the various molecular forms of IgA were assessed, with the 1:1 ratio corresponding to 0.61 μM of each respective Ab. Data are representative of two independent experiments performed in duplicates. **B**, LSCM images of immune complexes of bacteria associated with human plasma-derived plgA, SlgA, mlgA, IgG, or anti-*S. flexneri* LPS-specific SlgAC5 monoclonal Abs. Bacteria constitutively expressing GFP appear in green. Bound Abs were detected by antisera directed against the α or γ chain, followed by Abs conjugated to fluorophores, yielding red signals after image processing. Images are representative of one representative field obtained from 15 observations from three independent slides per experiment. Scale bars = 10 μm.
infection or responsiveness of target epithelial cells, depending on the size of formed immune complexes.

Human plasma IgM and SlgM Abs Induce Bacterial Agglutination, Reducing Interactions between S. flexneri and Caco-2 Cell Monolayers—Because the avidity of polyspecific pIgA/SIgA seems to be crucial to limit Caco-2 cell monolayer infection by S. flexneri, we sought to determine whether plasma-derived IgM would achieve similar functions. As shown in Fig. 3A, IgM and SIgM demonstrated concentration-dependent binding to S. flexneri. LSCM imaging revealed large aggregates comprising bacteria and either IgM or SIgM Abs (Fig. 3B), whose sizes were well above that detected previously after incubation with SIgA. This prompted us to speculate that such a strong agglutination capacity may result in blocking the internalization of S. flexneri by Caco-2 cell monolayers. After O/N infection with the bacteria alone, or in complex with IgM, SlgM...
or SIgA, cells were treated with gentamicin for 30 min, lysed, and then the lysate was plated on a selective medium for numeration. In comparison with bacteria alone, an at least 15-fold reduction in the number of cell-internalized bacteria was measured when associated with either IgM or SIgM (Fig. 4A), whereas a 2- to 3-fold decrease was observed in the presence of SIgA (Figs. 2A and Fig. 4A). These results illustrate a strong ability of IgM and SIgM to neutralize the bacteria via aggregation.

Human Plasma IgM and SIgM Abs Diminish the Secretion of Proinflammatory Chemokines and Cytokines by Caco-2 Cell Monolayers—The finding that plasma IgM and SIgM Abs appear superior to SIgA for all parameters tested so far led us to hypothesize that this should hold true when examining the proinflammatory responsiveness of Caco-2 cell monolayers exposed to the bacterium alone or in complex with Abs. Measurement of the basolateral secretion of TNF-α, CCL3, and CXCL8 by ELISA after O/N incubation showed that cell monolayers infected by SIgA-, IgM- or SIgM- S. flexneri complexes released at least 3-fold less TNF-α, CXCL8, and CCL3 than monolayers infected by the bacteria alone (Fig. 4B). Hence, neutralization of S. flexneri by either IgM or SIgM had the most marked effect on the proinflammatory response of the polarized Caco-2 cell monolayer of all Abs tested, and, importantly, this occurs in the absence of any other cell partner that could have biased the analysis.

Human Plasma IgM and SIgM Abs Efficiently Prevent Damage to Epithelial Caco-2 Cell Monolayers Infected by S. flexneri—Human plasma-derived IgM and reconstituted SIgM at the same molar concentration as SigA, serving as a reference control, were combined with S. flexneri and incubated O/N with Caco-2 cell monolayers. The integrity of the cell monolayers was assessed by TER measurement, cell morphology, and the number of, and total surface of, infected foci on the whole filters. In contrast to S. flexneri alone, TER was maintained when the bacteria were mixed with IgM or SIgM in a range similar to that observed with SIgA (Fig. 5A). Representative snapshots of transversal sections obtained along the x axis of the monolayers showed that the infected areas were systematically smaller after incubation with IgM and SIgM compared with incubation with SIgA or with S. flexneri alone (Fig. 5B). Detection of the preserved, well organized actin network in IgM-treated samples confirmed the improved monolayer integrity (Fig. 5B), as exemplified by limited binding and spreading of S. flexneri. When compared with bacteria alone, complexes with plasma SIgA diminished the total surface of infection foci 2-fold while slightly reducing their number (Fig. 5, C and D). Plasma IgM and SIgM showed a 15-fold and 10-fold reduction for these two parameters, respectively, reflecting lower damage of the monolayer (Fig. 5, C and D). Therefore, both IgM and SIgM prevent the destruction of Caco-2 cell monolayers exposed to infectious S. flexneri to a degree superior to SIgA.

**FIGURE 4. Modulatory effect of various human plasma-derived IgM/SIgM preparations on Caco-2 cells infected by S. flexneri.** A, bacteria internalized within Caco-2 cell monolayers determined after O/N infection by S. flexneri alone or in complex with human plasma IgM, reconstituted SigM, or SIgA. Bacterial counts were carried out after addition of gentamicin for the last 30 min of incubation. Data are expressed on a per filter basis and correspond to a pool of two independent experiments \((n = 4 \text{ or } 5)\) for each tested condition. B, basolateral secretion of TNF-α, CXCL8, and CCL3 by polarized Caco-2 cell monolayers after O/N incubation with S. flexneri alone or associated with plasma-derived IgM, SigM, or SIgA. The concentration of proinflammatory mediators was determined by ELISA. Non-infected Caco-2 cell monolayers \((\text{No bacteria})\) served as a control. Data are the pool of two experiments performed in triplicates \((n = 6)\). When obtained, significant statistical differences were calculated by comparison with the condition Sf. *, \(p < 0.01\); **, \(p < 0.01\); ***, \(p < 0.001\); nd, non-detectable.
Human Plasma Secretory-like Abs Impact the Secretion of the Virulence Factors IpaB and IpaC Released by S. flexneri—In addition to its protective properties, the specific anti-S. flexneri IgAC5 monoclonal Ab is known to mediate a transient suppression of the type 3 secretion system of the bacterium, leading to a decrease in secretion of the virulence factor known as IpaB (37). To examine whether polyreactive human plasma-derived IgA or IgM could act via a similar mechanism, we evaluated the secretion of IpaB and IpaC under identical conditions as those used to form immune complexes in protection experiments. S. flexneri was associated with human plasma pIgA, SIgA, mIgA, IgM, SIgM, IgG, or specific IgAC5 monoclonal Ab for 1 h or left as such. Although bacteria not exposed to Congo red secreted a low basal level of IpaB and no IpaC, exposure to Congo red led to secretion of various levels of IpaB and IpaC in supernatants as a function of the complexing Ab. The level of secretion of both IpaB and IpaC by bacteria in complex with human plasma SIgA or SIgM was decreased (Fig. 6, A and B). In comparison with the bacterium alone, the other molecular forms of IgA, IgM, as well as IgG and SIgAC5, did not lead to measurable changes (Fig. 6C). This result suggests that the secretory form of plasma-derived IgA or IgM may contribute indirectly to protection of target epithelial surfaces through its impact on bacterial virulence.

DISCUSSION

We established previously that human plasma-derived plgA and IgM can be assembled into secretory-like Abs (29). However, how the various molecular forms of the Ab displayed differential protection was not evaluated at the cellular and molecular levels. This study addresses these issues and further extends the analysis to plasma-derived IgM and SIgM. We found that plgA and IgM or the secretory form of the Ab recognized S. flexneri to the same extent and that the interaction led to the formation of aggregates with a size dependent on the valence of the Ab. Although reduced bacteria internalization into the cell monolayer occurred with all Ab isotypes and molecular forms tested, bacteria-induced damage to the monolayer organization was diminished significantly with polymeric and secretory-like Abs only, as identified by TER measurement, cell imaging, quantification of infected foci, and assessment of areas exhibiting monolayer destruction as well as secretion of proinflammatory mediators. Moreover, plasma-derived Abs were found to disable S. flexneri in its capacity to produce IpaB
Homeostatic Properties of Secretory-like Plasma IgA and IgM

A

| S. flexneri | + | + | + | + | + | + | + |
|-------------|---|---|---|---|---|---|---|
| Antibody    | - | SlgAC5 | plgA | SlgA | mlgA | IgG | IgM |
| IpaB        |   |       |       |       |       |     |     |
| OspF        |   |       |       |       |       |     |     |

B

| S. flexneri | + | + | + | + | + | + | + |
|-------------|---|---|---|---|---|---|---|
| Antibody    | - | SlgAC5 | plgA | SlgA | mlgA | IgG | IgM |
| IpaC        |   |       |       |       |       |     |     |
| OspF        |   |       |       |       |       |     |     |

C

FIGURE 6. Impact of human plasma-derived IgA and IgM preparations on the secretion of virulence factors IpaB and IpaC by *S. flexneri*. Immune complexes between bacteria and human plasma-derived plgA, SlgA, mlgA, IgG, IgM, SigM, or anti-Shigella LPS-specific SlgAC5 monoclonal Ab were formed for 1 h as described under “Experimental procedures,” and the expression of virulence factors was induced by Congo red. A and B, the secretion of IpaB (A) and IpaC (B) was examined by immunoblot analysis using monoclonal Abs directed against IpaB and IpaC, with immunodetection of OspF serving as a loading control. The images are representative of one individual experiment performed three times in duplicates. C, densitometric analysis of replicated immunoblots (n = 6) exposed for optimal times to avoid saturation of the photographic film. The intensity of the signals reached with *S. flexneri* alone was fixed arbitrarily at 100%. When obtained, significant statistical differences were calculated by comparison with the condition Sf. *, p < 0.02; **, p < 0.005.

and IpaC, two proteins involved in the invasion of epithelial cells. A direct comparison with SlgA led to the conclusion that IgM and SlgMAb molecules demonstrate a superior activity in preserving the integrity and responsiveness to infection of polarized Caco-2 cell monolayers used as a mimic of the gut mucosal epithelium.

Similar in vitro models have been used previously to compare the neutralizing function of antigen-specific monoclonal IgG and IgA/SlgA (31, 39, 40). Although one has to acknowledge that the Caco-2 cell line may not recapitulate all functions of normal intestinal epithelial cells, their pattern of cytokines following exposure to *S. flexneri* was consistent with that of other cell lines and that found in vivo (41, 42). The availability of polyreactive IgA and IgM Abs with well characterized biochemical properties (29) allowed us to directly address their modes of action in the absence of any other cellular and molecular partners involved in clearance of *S. flexneri*, therefore allowing a comparison of their respective functional characteristics.

Remarkably, for most of the parameters examined, in comparison with the specific protective SlgAC5 monoclonal Ab, polyreactive plgA and SlgA displayed similar degrees of protection at a 10-fold higher concentration only. Both IgM and SlgM proved to be even more potent. This is particularly true for blocking of internalization, for maintenance of the polarized Caco-2 cell monolayer integrity, and, to a lesser extent, for reduction in the production of proinflammatory TNF-α. This may be explained by the observation that apical immune exclusion appears to be the most potent function of multivalent IgM Abs. Strikingly, agglutination mediated by plgA and SlgA did not reduce the internalization of bacteria with a better efficiency than mlgA and IgG. This contrasts with the observation that LPS-specific monomeric IgAC5 and IgGC20 monoclonal Abs perform more poorly than their SlgA counterpart recognizing the same epitope (31). This suggests that the polyclonal nature of plasma-derived mlgA and IgG masking both LPS and bacterial adhesins impairs bacterial entry more efficiently. Alternatively, it may block intracellular bacterial proliferation without impacting epithelial cell responsiveness, such as morphological changes and secretion of proinflammatory mediators. In addition, this reveals that the sensing of antigens by epithelial cells might differ depending on the molecular form of the Abs participating in the immune complexes as, for instance, intracellular processing pathways (43).

The presence of SC in reconstituted SlgA and SlgM did not modify in any way the function of the Ab molecule in our model, with the notable exception of the effect on IpaB and IpaC secretion by *S. flexneri*. However, in vivo, the presence of bound SC is essential for the stability and anchoring of the molecule at mucosal surfaces (3). Furthermore, in addition to interfering with bacterial targeting of epithelial cells via carbohydrates abundantly found on its surface (44), SC associated with plgA and IgM appears to intervene negatively in the secretion of *S. flexneri* virulence factors crucial for the infection of epithelial cells (45, 46). These results suggest the unexpected role of polyclonal SlgA and SlgM mostly in altering bacterial metabolism, which adds to recognized mechanisms of protection effective at the level of the mucosal epithelium.

Previous animal studies have demonstrated that administration of polyclonal IgM molecules, especially derived from human plasma, could be beneficial in case of sepsis. Lachmann et al. (25) showed that IgM-enriched preparations reduced *Klebsiella pneumoniae* infection using a distress syndrome rat model. Stehr et al. (26) emphasized the benefits of polyclonal IgM-enriched solution by using a rabbit model of bacteremia. Clinical trials showed that IgM-enriched preparations were able to significantly decrease endotoxin levels in plasma (23) and even reduce mortality (22) during the early phase of septic shock. Norrby-Teglund et al. (47) showed that IgM-enriched...
preparations were able to inhibit specific streptococcal antigens. However, as a limitation to the interpretation of these results, IgM-enriched preparations contained significant amounts of IgG and/or IgA Ab molecules, making it difficult to strictly assign the protective effect to the IgM moiety of the product. These in vivo studies, together with our current demonstration of epithelial cell protection with IgM/SigM, may suggest that mucosal passive delivery of IgM via the gastrointestinal and nasal routes are worth considering.

Although systemic IgM-based immune complexes may lead to the activation of complement, mucosal delivery along the gut is not expected to trigger such a process because the antibody distribution will be restricted to the lumen. In addition, previous research has shown that epithelial cells and follicular DCs in germinal centers producing cell membrane complement regulatory proteins (protectin CD59), membrane cofactor protein, and decay-accelerating factor (CD55) are protected from the action of complement (48, 49). Together with SlgA exhibiting an important reduction in cytokine release as well, this would combine protection and low proinflammatory responses by the epithelial cells located at the interface between the environment and the inside of the body. In addition to potential clinical applications against infectious agents, it may be appealing to apply IgA and IgM preparations topically to patients with primary immune deficiency, which is associated with chronic infection and autoimmunity (50).

In the in vivo context, one can argue that commensal bacteria may act as a “trap” for exogenously delivered SlgA/SlgM. However, the fact that a large percentage of bacteria are coated with endogenous SlgA (51) at a steady state suggests that administration of milligram amounts of either SlgA or SlgM should leave enough operative Ab molecules available. A plausible explanation would be that both non-antigen-specific (possibly mediated by carbohydrate moieties (52) and specific interactions contribute to the recognition of bacterial antigens by the polyreactive Abs.

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