Immunoglobulin Responses at the Mucosal Interface

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Keywords
mucosal immunity, B cells, IgA, IgD, class switching

Abstract

Mucosal surfaces are colonized by large communities of commensal bacteria and represent the primary site of entry for pathogenic agents. To prevent microbial intrusion, mucosal B cells release large amounts of immunoglobulin (Ig) molecules through multiple follicular and extrafollicular pathways. IgA is the most abundant antibody isotype in mucosal secretions and owes its success in frontline immunity to its ability to undergo transcytosis across epithelial cells. In addition to translocating IgA onto the mucosal surface, epithelial cells educate the mucosal immune system as to the composition of the local microbiota and instruct B cells to initiate IgA responses that generate immune protection while preserving immune homeostasis. Here we review recent advances in our understanding of the cellular interactions and signaling pathways governing IgA production at mucosal surfaces and discuss new findings on the regulation and function of mucosal IgD, the most enigmatic isotype of our mucosal antibody repertoire.
INTRODUCTION

Mucosal membranes provide a dynamic interface that separates the sterile milieu of our body from the external environment. A key component of this interface is the mucosal epithelium, which blocks invasion by pathogenic and commensal bacteria by forming multiple layers of physical and immune protection (1). Epithelial protection is particularly sophisticated in the intestine, which contains large communities of commensal bacteria that process otherwise indigestible polysaccharides, synthesize essential vitamins and isoprenoids, stimulate the maturation of the immune system, and form an ecological niche that prevents the growth of pathogenic species (2). Conversely, the intestine provides commensals with a stable habitat rich in energy derived from ingested food.

A fine line exists between the homeostatic balance required to maintain commensals and the destructive response required to repel pathogens (3). Such a balance involves an intimate dialogue between prokaryotic and eukaryotic cells of our body that ultimately generates finely tuned signaling programs that ensure a state of hyporesponsiveness against commensals and a state of active readiness against pathogens (4). In this dialogue, our epithelial cells function as interpreters that continuously translate prokaryotic messages to educate the mucosal immune system as to the composition of the local microbiota (5).

Although needed by the host, commensals represent a potential threat of infection and unrestrained inflammation (6). A major defensive mechanism that excludes commensals from the mucosal surface involves immunoglobulin A (IgA) (7). This antibody class works together with nonspecific protective factors such as mucus to block microbial adhesion to epithelial cells without causing tissue-damaging inflammatory reaction (8). By doing so, IgA establishes a state of armed peace in the homeostatic interaction between the host and noninvasive commensal bacteria. When invasive pathogenic bacteria trespass the epithelial border, a state of open war breaks out and IgA receives help from IgG to repel the invaders. In this life-threatening situation, IgG provides a second line of defense that controls microbial dissemination by eliciting a robust inflammatory reaction (9).

Remarkably, different mucosal districts are characterized by distinct antibody signatures. The intestinal tract contains IgA and some IgM but virtually no IgG, whereas the respiratory and urogenital tracts contain equivalent amounts of IgA and IgG in addition to some IgM (10). In humans, the intestinal and urogenital tracts produce large amounts of an IgA subclass known as IgA2, whereas the respiratory tract contains IgD, the most enigmatic class of our mucosal antibody repertoire (10). This review discusses recent advances in our understanding of the regulation and function of mucosal IgA and IgD.

MUCOSA-ASSOCIATED LYMPHOID TISSUES

General Features

Mucosal surfaces comprise various lymphoid structures collectively referred to as mucosa-associated lymphoid tissue (MALT) (8). This secondary lymphoid organ can be further divided in functionally connected subregions, including the gut-associated lymphoid tissue (GALT), nasopharynx-associated lymphoid tissue (NALT), and bronchus-associated lymphoid tissue (BALT) (11–13). In the MALT, functionally distinct inductive and effector sites can be recognized. Intestinal Peyer’s patches (PPs) and mesenteric lymph nodes (MLNs) exemplify mucosal inductive sites, which contain T and B cells undergoing clonal expansion and differentiation upon activation by antigen (14). The intestinal lamina propria (LP) exemplifies mucosal effector sites, whose main function is to recruit effector T and B cells emerging from inductive sites (15).

Antibodies released by effector B cells, including plasma cells, provide the first line of protection at mucosal surfaces. In the intestinal tract and other mucosal districts, the vast
majority of mucosal plasma cells secrete dimeric or oligomeric IgA and to a lesser extent pentameric IgM, both of which interact with the polymeric Ig receptor (pIgR) expressed on the basolateral surface of epithelial cells through a joining (J) chain (16). This interaction is followed by endocytosis of IgA and IgM, proteolytic cleavage of pIgR, and basolateral-to-apical transcytosis of secretory IgA (SIgA) and SIgM, which comprise a pIg-derived secretory component (SC) providing mucophilic properties to the SIg complex (16). As for IgG, this antibody class gains access to respiratory and urogenital secretions through both nonspecific and specific mechanisms, the latter involving interaction with an antibody transporter known as neonatal Fc receptor (nFcR) (10, 17). The mechanism by which IgD enters respiratory secretions remains unknown (10, 18, 19).

**GALT and NALT**

Although equipped with a general architecture resembling that of systemic lymphoid organs, the MALT has several unique features, including absence of afferent lymphatics, which requires sampling of antigen directly from the epithelial surface (8). In the GALT, antigen sampling involves specialized microfold (M) cells lodged in the follicle-associated epithelium (20). M cells filter bacteria through a complex glycocalix and sample some of these bacteria via poorly defined receptors (21). IgA receptors allow M cells to sample IgA-coated commensals (22), whereas glycoprotein 2 receptor samples IgA-free commensals as well as pathogens (23). Sampled antigen is eventually transferred to dendritic cells (DCs), which occupy large basolateral invaginations of M cells (21). Alternatively, DCs directly sample antigen from the lumen through transepithelial projections (24). These antigen-loaded DCs migrate to the perifollicular area of PPs to present antigen to CD4+ T cells and initiate antigen-specific T and B cell responses (8).

The NALT has antigen-sampling strategies similar to those present in the GALT, including M cells (11). In humans, the NALT consists of a set of lymphoid aggregates known as Waldeyer’s ring that occupy strategic areas of the oropharynx and nasopharynx and include pharyngeal tonsils (or adenoids) as well as tubal, palatine, and lingual tonsils (10). NALT organogenesis markedly differs from GALT organogenesis in terms of both kinetics and cytokine requirements (11, 25). For example, PPs begin their development during embryonic life and require signals from interleukin-7 (IL-7) and lymphotoxin receptors, whereas tonsils initiate their development shortly after birth and require signals from environmental antigens (11, 25). Another important difference between GALT and NALT is that NALT-targeted immunization preferentially induces antigen-specific immunity in the respiratory and genital tracts, whereas GALT-targeted immunization mainly elicits protective responses in the gastrointestinal tract (11, 26).

**NATURE AND FUNCTION OF MUCOSAL ANTIBODIES**

**Genesis and Function of Antibody Diversity**

Diversification is essential for the mucosal immune system to mount protective responses. Antibodies diversify through three major DNA-modifying processes known as V(D)J recombination, class switch recombination (CSR), and somatic hypermutation (SHM). Bone marrow B cell precursors generate antigen recognition diversity through V(D)J recombination, an antigen-independent process mediated by recombination-activating gene (RAG) endonucleases that assemble antigen-binding Ig variable regions from individual V (variable), D (diversity), and J (joining) gene segments (27). Mature B cells emerging from the bone marrow colonize peripheral lymphoid organs, where they undergo a second wave of Ig gene remodeling through SHM and CSR, two antigen-dependent processes that require the DNA-editing enzyme activation-induced cytidine deaminase (AID) and mediate antibody affinity maturation and antibody
class (or isotype) switching, respectively (28, 29).

SHM introduces point mutations within V(D)J exons, thereby providing the structural correlate for selection of high-affinity Ig mutants by antigen, whereas CSR replaces constant μ (Cμ) and Cδ exons encoding IgM and IgD with Cγ, Cα, or Cε exons encoding IgG, IgA, or IgE, thereby providing antibodies with novel effector functions without changing their antigen-binding specificity (30, 31). In humans, a noncanonical form of CSR from Cα to Cε also exists and generates B cells specialized in IgD production (18, 32). As already discussed, IgA mediates its effector functions by interacting with pIgR on the basolateral surface of mucosal epithelial cells (16). In addition, IgA binds to a high-affinity IgA receptor known as FcαRI (or CD89), which is expressed on granulocytes, monocytes, macrophages, DCs, natural killer (NK) cells, and mast cells (33). These innate immune cells generate activating or regulatory signals upon FcαRI receptor engagement by IgA, depending on the degree of IgA oligomerization (34).

Additional IgA receptors include transferring receptor (CD71), FcαRII (or CD23) (37–39). As for IgG and IgE, these antibody classes initiate multiple innate and adaptive immune responses by activating high- or low-affinity FcγR and FcεR on various innate immune cells (35). In addition, IgG undergoes bidirectional transcytosis across epithelial cells by binding to nFcR (17, 36). In IgE-mediated allergic disorders, IgE undergoes transcytoses across epithelial cells by utilizing a low-affinity IgE receptor known as FcεRII (or CD23) (37–39). The receptors mediating IgD effector functions and IgD transcytosis remain elusive. Although expressing abundant J chain, IgD-secreting plasma cells seem to release monomeric IgD only, which does not bind to pIgR (10). Considering that the hinge region of IgD has a heparin ligand–like site, heparin and other heparan sulfate proteoglycans may play an important role in at least some IgD effector functions (18).

### Antibody Composition of Mucosal Sites

Differences in epithelial cell expression of specific antibody transporters and plasma cell–recruiting chemokines contribute to determining the antibody class composition of a given mucosal site. In general, IgA is the most abundant antibody isotype in mucosal secretions. Yet, IgA is somewhat less abundant than IgG in the urine, bile, and genital and bronchoalveolar secretions. IgD can be detected in nasal, salivary, lacrimal, and bronchoalveolar secretions (10, 18, 19), whereas IgE is measurable in nasal, bronchoalveolar, and intestinal secretions, at least when allergy is present (37).

Unlike mouse B cells, human B cells produce two IgA subclasses, IgA1 and IgA2, which have a similar receptor-binding profile but different topography from each other (40). IgA1 is present in both systemic and mucosal districts, whereas IgA2 is mostly present in mucosal districts colonized by a large microbiota, including the distal intestinal tract and the urogenital tract (40). This circumstance could reflect the fact that IgA2 is more resistant than IgA1 to degradation by bacterial proteases because IgA2 has a shorter hinge region than IgA1 does (41, 42).

### Function of Mucosal IgA

Mucosal IgA comprises antibodies that recognize antigen with high- and low-affinity binding modes (8). In general, high-affinity IgA neutralizes microbial toxins and invasive pathogens, whereas low-affinity IgA confines commensals in the intestinal lumen. Yet this distinction is not absolute, as growing evidence indicates an important role for high-affinity IgA in the control and regulation of the commensal microbiota (43). Conversely, additional evidence shows that low-affinity IgA protects against some pathogens (44, 45). High-affinity IgA is thought to emerge from follicular B cells stimulated via T cell–dependent (TD) pathways, whereas low-affinity IgA likely emerges from extrafollicular B cells stimulated via T cell–independent (TI) pathways (14). However, this view is rapidly changing. Indeed, recent
IgA also promotes the maintenance of appropriate bacterial communities in specific intestinal segments (47). Indeed, IgA modulates the gene-expression profile of commensal bacteria, selecting species with less inflammatory activity on the host’s tissues (48). By preventing overstimulation of mucosal B cells, this IgA-mediated selection process may impede bacteria-induced expansion of autoreactive, inflammatory, proallergic, and neoplastic B cell clones (7, 14, 40). Furthermore, IgA facilitates the establishment of specific symbiotic relationships not only in the mucosal lumen, but also in PPs (49). As discussed above, IgA also facilitates sampling of luminal antigens by binding to poorly defined receptors on M cells (22, 50). Moreover, IgA neutralizes inflammatory microbial products inside epithelial cells (51). Finally, if bacteria trespass the epithelial barrier, IgA transports these bacteria back into the lumen via plgR or promotes their phagocytosis through FcαRI (34, 52).

The in vivo relevance of IgA can be best seen in patients with SlgAD, common variable immunodeficiency (CVID), and hyper-IgM (HIGM) syndrome. In these primary immunodeficiencies, impaired IgA production is associated with recurrent respiratory and gastrointestinal infections as well as allergy and autoimmunity (53, 54). Some IgA-deficient patients also develop intestinal inflammation and small intestinal nodular lesions, possibly resulting from excessive B cell stimulation by aberrantly expanded commensals (14, 55). Yet SlgAD, CVID, and even some cases of HIGM syndrome can be asymptomatic or mildly symptomatic, possibly because of compensatory increases of unaffected antibodies such as IgM and IgD (10, 18, 53, 54).

Function of Mucosal IgD

The function of IgD has puzzled immunologists over the past several decades. Originally thought to be a recently evolved isotype, IgD is now recognized to be an ancestral molecule that has been conserved throughout evolution to complement the functions of IgM (18, 56). IgD would afford protection to the respiratory mucosa by binding to pathogenic bacteria such as Moraxella catarrhalis and Haemophilus influenzae as well as to their virulence factors (32, 57). In addition to crossing epithelial cells, IgD binds to circulating basophils, monocytes, and neutrophils as well as to mucosal mast cells through unknown receptors (18, 58).

Consistent with recently published data showing the important role of basophils in T helper type 2 (Th2) cell responses and antibody production (59–63), IgD cross-linking induces basophil release of B cell–activating cytokines such as interleukin (IL)-4 and IL-13, which in turn facilitate IgM as well as IgG and IgA production (32). Furthermore, IgD cross-linking triggers basophil release of antimicrobial peptides such as cathelicidin, inflammatory cytokines such as IL-1β and TNF, and various chemokines such as CXCL10 (32, 58). Therefore, IgD may contribute to mucosal immunity not only by neutralizing pathogens and excluding commensals, but also by recruiting basophils as well as other immune cells with antimicrobial and immune-augmenting functions (18).

PATHWAYS INDUCING MUCOSAL IgA RESPONSES

T Cell–Dependent Pathways

Most antigens initiate mucosal IgA responses through a TD reaction that takes place in mucosal lymphoid follicles (8, 15), such as intestinal PPs and MLNs (Figure 1a, b). These organized structures comprise a germinal center that fosters antibody diversification and affinity maturation, including SHM and CSR, through antigen–specific cognate interactions between B cells that express the CD40 receptor and CD4+ Th cells expressing CD40 ligand (CD40L) (13, 40). Together with cytokine receptors and B cell antigen receptor (BCR), CD40 is critical for the induction of AID expression and the initiation of SHM and CSR.
Engagement of CD40 by CD40L leads to (a) the recruitment of TNF receptor-associated factor (TRAF) adapter proteins to the cytoplasmic tail of CD40 (65), followed by (b) activation of an NF-κB inhibitory protein (IkB) kinase (IKK) complex, which triggers (c) phosphorylation and degradation of IkB that retards NF-κB in an inactive state (66, 67). The resulting IkB-free NF-κB proteins translocate from the cytoplasm to the nucleus to initiate transcription of the AICDA gene promoter that encodes AID (40, 68). In contrast, NF-κB is not required for the activation of the intronic α (Iα) promoter upstream of the Cα gene and therefore has little or no role in germ-line Cα gene transcription (40, 69). This circumstance may explain why additional signals from cytokines such as transforming growth factor-β (TGF-β) are needed to elicit IgA CSR, at least in mice (13).

T Cell–Independent Pathways

The conventional TD pathway requires 5 to 7 days to initiate protective antibody responses in systemic lymphoid tissues (70, 71). Such kinetics may not be appropriate to afford optimal mucosal protection because mucosal surfaces are constantly exposed to dietary and bacterial antigens. In addition, the TD pathway is often associated with an inflammatory reaction that could disrupt the mucosal epithelial barrier. To compensate for these limitations, the intestinal mucosa has developed a faster TI pathway that generates IgA in response to highly conserved microbial signatures recognized by Toll-like receptors (TLRs) (14, 40), a family of germ-line gene-encoded antigen receptors involved in the activation of both innate and adaptive arms of the immune system (72, 73). In mice, TI IgA production involves B-1 cells from the peritoneal cavity and intestinal LP as well as conventional B-2 cells from isolated lymphoid follicles (ILFs) (13). These B cells release low-affinity IgA (and IgM) in the absence of help from CD4+ T cells via CD40L (74–76). The human counterpart of mouse B-1 cells remains unknown.

TLRs facilitate TI IgA responses either by activating B cells directly or by inducing release of the B cell–activating factor of the TNF family (BAFF, also known as BLyS) and its homolog a proliferation-inducing ligand (APRIL) from innate immune cells (7, 40). Engagement of TLRs by microbial ligands triggers activation of NF-κB (72, 73). This activation requires recruitment of the adaptor protein MyD88 to a cytoplasmic Toll-interleukin-1 receptor (TIR) domain that subsequently elicits formation of an IKK-activating signaling complex composed of IL-1 receptor–associated kinase (IRAK)1, IRAK4, TRAF6, and TGF-β-activated kinase (TAK)-1 (73).

In addition to inducing AID expression in B cells (77, 78), TLR signaling via NF-κB elicits BAFF and APRIL expression in DCs, macrophages, granulocytes, and epithelial cells, including intestinal epithelial cells (IECs) (79–84). In the presence of appropriate cytokines, BAFF and APRIL initiate Cα...
germ-line expression and Cμ-to-Cα CSR by engaging a CD40-related receptor known as transmembrane activator and calcium modulator and cyclophylin ligand interactor (TACI) (84–88). One important property of this receptor is to establish a close cooperation with B cell–intrinsic signals from TLRs (85).

IgA RESPONSES IN MUCOSAL FOLLICLES

Role of CD4+ T Cells

Growing evidence indicates that TD IgA responses involve a heterogeneous population of CD4+ T cells (Figure 1c), including T follicular helper (Tfh) cells, Th2 cells, and T regulatory (Treg) cells (13, 40). These CD4+ T cell subsets express CD40L and release large amounts of IgA-inducing cytokines (89–92). But how do IgA-inducing Tfh, Th2, and Treg cells differentiate from naive CD4+ T cells? A key role is played by DCs strategically positioned in the subepithelial area (21). In addition to receiving antigen from M cells, these DCs directly sample antigen in the intestinal lumen by emanating transepithelial projections through a process controlled by signals from TLRs in IECs (24, 93). Overall, M cells and DCs capture only small amounts of bacteria because intestinal IgA responses have a much higher induction threshold (108–109 bacteria) than do systemic IgG responses (43).

Antigen sampling through transepithelial projections involves nonmigratory gut-resident DCs expressing the fractalkine receptor CX3CR1 (94) developmentally distinct from migratory DCs expressing the αE integrin CD103 (95, 96). As they sample antigen in the lumen, DCs receive powerful conditioning signals from IECs, thereby becoming primed for the induction of noninflammatory CD4+ T cells with an IgA-inducing function (7, 92, 97). Antigen-loaded DCs migrate from subepithelial to perifollicular areas, where they induce Treg and Th2 cell differentiation (95, 96, 98, 99). Although Th2 cells may elicit CD40-dependent IgA CSR and production by releasing IL-4, IL-5, IL-6, and IL-10 (89, 92, 100, 101), Treg cells would do so by releasing TGF-β (90, 91). Some Treg cells would further differentiate into Tfh1 cells, which induce IgA CSR and production via IL-21 and TGF-β1 (91, 102). In general, CD4+ T cells that provide help to B cells in PPs are clearly functionally different from their counterparts that initiate IgG responses in systemic lymphoid follicles, which may explain why intestinal IgA responses have a slower onset than systemic IgG responses (>14 days versus 5–7 days) (43).

IgA-producing B cells generated via TD pathways further differentiate into IgA-secreting plasma blasts in the intestinal LP (7). Here, Th17 cells might facilitate transepithelial release of SlgA through an IL-17-dependent mechanism involving upregulation of pIgR expression (103), although thus far this mechanism has only been described in the respiratory mucosa. Despite requiring CD4+ T cells for germinal center formation, PPs are capable of producing IgA independently of canonical T-B cell interactions (104). Indeed, PPs retain TD IgA responses in the absence of BCR, which is instead required by systemic follicles to initiate TD IgG responses (104). Thus, PPs may generate IgA through an alternative TD pathway that involves activation of B cells by microbial TLR ligands (105). Supporting this model, lack of MyD88 impairs IgA production in intestinal PPs (46, 106).

Role of Conventional Dendritic Cells

Intestinal DCs maintain homeostasis not only by inducing noninflammatory IgA responses, but also by dampening inflammatory Th1 and Th17 cell responses (7, 13, 99, 107–109). Strikingly, both processes implicate DC-induced differentiation of antigen-specific Th2, Treg, and Tfh1 cells (90–92, 98). Intestinal DCs are particularly skilled in eliciting these homeostatic CD4+ T cell responses (Figure 1c) because they receive conditioning signals from IECs (97, 107). One of these signals is provided by thymic stromal lymphopoietin (TSLP), an IL-7-like epithelial cytokine that shifts the
Th1/Th2 balance toward Th2 polarization by attenuating DC production of IL-12 but not of IL-10 (92, 110).

Importantly, IECs release TSLP in response to TLR-mediated signals from bacteria (111). Accordingly, genetic disruption of TLR signaling via NF-κB reduces TSLP expression by IECs and augments IL-12 production by DCs (112). In addition to TSLP, IECs release TGF-β and retinoic acid, which stimulate the development of CD103+ DCs, at least in vitro (113). These DCs promote the formation of Treg cells via TGF-β and retinoic acid and suppress the development of inflammatory Th1 and Th17 cells (99, 114, 115). As discussed earlier, Treg cells emerging from this pathway could induce IgA CSR and production through TGF-β (90, 91).

It must be noted that thus far no DC subset has been formally assigned to the induction of IgA responses in PPs. One notable exception is a TNF-α-inducible nitric oxide synthase–producing DC (tipDC) subset that enhances IgA CSR and production by upregulating the expression of TGF-β receptor on B cells from PPs via nitric oxide (106). The ontogenetic and functional relationships of tipDCs with other intestinal DC subsets remain unclear.

Role of Follicular Dendritic Cells

Germinal centers from PPs and MLNs contain a meshwork of antigen-trapping follicular dendritic cells (FDCs) ontogenetically different from DCs. Indeed, FDCs originate from nonhematopoietic precursors that may include mesenchymal cells (116). One of the main functions of FDCs is to facilitate the positive selection of high-affinity follicular B cells by antigen (117). During this process, antigen arrays on the surface of FDCs activate B cells by extensively cross-linking BCR (118). As shown by recent studies (46), FDCs from PPs and MLNs efficiently induce IgA CSR and production (Figure 1). This response occurs in a TI manner and involves TLR-mediated sensing of bacteria by FDCs, followed by FDC release of TGF-β, BAFF, and APRIL (46).

Role of Lymphoid Tissue Inducer Cells

Together with PPs and MLNs, IFLs represent another important site for IgA induction (14). These lymphoid structures are scattered throughout the intestine and consist of solitary B cell clusters built on a scaffold of stromal cells with interspersed CD4+ T cells and abundant perifollicular DCs (119). Unlike PPs, IFLs appear after birth in response to bacteria colonization (119). Recent findings indicate that TLR signals from commensal bacteria initiate a crosstalk centered on RORγt+ lymphoid tissue inducer (LTi) cells (74). These cells recruit DCs and B cells through various chemokines and stimulate release of active TGF-β, BAFF, and APRIL by activating DCs and stromal cells via a signaling loop involving lymphotixin (74). Together with microbial TLR ligands, TGF-β, BAFF, and APRIL induce IgA CSR in the absence of help from CD4+ T cells (74). Of note, LTi cells also release the IEC-activating cytokine IL-22 and the B cell–attracting chemokine CXCL13, which may enhance IgA production in addition to promoting homeostasis (120).

Homing of IgA-Expressing B Cells

IgA-producing B cells generated in PPs and MLNs upregulate the expression of critical gut-homing receptors such as the α4β7 integrin as well as CCR9 and CCR10 chemokine receptors upon being primed by retinoic acid that is released by local DCs (121). These retinoic acid–primed B cells enter the general circulation via the thoracic duct and thereafter gain access to the LP by binding mucosal addressin-cell adhesion molecule-1 (MAdCAM-1) expressed on LP-based high endothelial venules through α4β7 (122). Retinoic acid–primed B cells further respond to CCL25 and CCL29, two IEC chemokines that bind to CCR9 and CCR10, respectively (122). Once in the LP, IgA-expressing B cells terminally differentiate into IgA-secreting plasma cells, possibly through the help of local signals from IECs, DCs, and macrophages (40). Similar cells would
also provide robust activation and survival signals to B cells emerging from PPs and MLNs, which may explain why intestinal IgA responses do not always correlate with a florid germinal center reaction in PPs and yet show a sustained half-life (>16 weeks) (43).

IgA RESPONSES IN MUCOSAL EXTRAFOXLICULAR AREAS

IgA CSR in the Intestinal Lamina Propria

The diffuse tissue of the LP is an additional site for IgA production and diversification (Figure 1c), although it is less important than PPs and MLNs (123). Consistent with this possibility, genetically engineered mice lacking PPs, MLNs, and even ILFs retain some antigen-specific IgA plasma cells, which are mostly located in the LP (124, 125). In both mice and humans, a fraction of B cells from the intestinal LP contain molecular hallmarks of ongoing IgA CSR, including AID, H2AX protein (a nuclear protein associated with double-strand DNA breaks generated by AID within S regions), excised S\(\alpha\)-S\(\mu\) switch circles, and switch circle I\(\alpha\)-C\(\mu\) transcripts (111, 126–128). Of note, AID and IgA remain detectable in LP B cells from mice and humans lacking CD4\(^+\)T cells or CD40L (74, 111, 129). In general, LP B cells are less activated and more scattered than PP B cells, and therefore LP IgA CSR can pass unrecognized unless highly sensitive and accurate methodologies are used (126, 130, 131). In this regard, a green fluorescent protein–AID reporter mouse model has proven very helpful (129). The mechanisms underlying IgA CSR in the LP remain poorly understood but likely involve DCs and IECs (111, 132).

Role of DCs

Abundant evidence demonstrates that DCs can initiate TI CSR and antibody production by activating B cells through antigen and cytokines, including BAFF and APRIL (84, 133–138). In the intestine, antigen-sampling CX3CR1\(^+\) DCs or CD103\(^+\) DCs may activate LP B cells through similar mechanisms (Figure 1c). A LP DC subset with clearer B cell–stimulating function is that of tipDCs (106). These DCs initiate TI IgA production by releasing BAFF and APRIL through a mechanism involving TLR-induced iNOS-mediated nitric oxide production (Figure 1c). In the LP, another DC subset with B cell–licensing functions is that of CD11c\(^+\)CD11b\(^+\) DCs (139). These DCs induce TI IgA production upon sensing bacteria through TLR5, a process that elicits release of retinoic acid and IL-6 (Figure 1c).

Role of Intestinal Epithelial Cells

IECs and respiratory epithelial cells can deliver IgA-inducing signals to LP B cells by releasing BAFF, APRIL, and IL-10 in response to TLR signals (79, 111, 127, 140). Similar epithelial cells can also amplify DC production of BAFF, APRIL, and IL-10 by stimulating DCs through TSLP (79, 111). In humans, APRIL is particularly effective at inducing IgA2, an IgA subclass particularly abundant in the distal intestine (111). In addition to triggering direct IgM-to-IgA1 CSR (111), APRIL elicits sequential IgA1-to-IgA2 CSR in the LP of the distal intestine (Figure 1c). This process would allow B cells arriving from PPs to acquire an IgA2 subclass more resistant than IgA1 to degradation by bacterial proteases (42).

IgD RESPONSES IN THE RESPIRATORY MUCOSA

Geography of IgD Production

IgD constitutes a significant fraction of the antibodies produced in the upper segments of the human respiratory and digestive tracts (Figure 2a). The mucosal IgD class originates predominantly from IgD\(^+\) IgM\(^-\) B cells bearing morphologic and immunophenotypic features of plasmablasts (32). Indeed, IgD\(^+\) IgM\(^-\) B cells display canonical plasma cell traits such as an eccentric nucleus, a large basophilic cytoplasm filled with IgD (Figure 2b), and J chain as well as IgD-secreting activity, in addition to features typical of mature B cells such as expression of surface IgD and CD19 (32, 141–143). Of note, IgD\(^+\) IgM\(^-\) plasmablasts originate from a
process of C_μ_−to-C_δ__ CSR that leads to the loss of IgM expression (142). This process takes place in the aerodigestive mucosa because this site contains various molecular hallmarks of ongoing C_μ_−to-C_δ__ CSR (32). In general, the respiratory mucosa expresses chemokines and vascular adhesion molecules capable of promoting the recruitment of IgD^+IgM^− plasmablasts from the periphery (144). In this regard, the peripheral blood contains some IgD^+IgM^− plasmablasts, which may be in transit to reach distant mucosal effector sites (32, 145). IgD^+IgM^− plasmablasts are rarely found in the GALT, probably because these B cells express little or no gut homing receptors such as α4β7 and CCR9 (144).

**Regulation of IgD Production**

In humans and other higher mammals such as cows, a rudimentary S-like intronic DNA region known as σ_5 is present upstream of the C_δ_ exon (18). Like canonical S regions, σ_5 contains guanosine-cytosine repeats and serves as an acceptor DNA region for donor S_μ_ to mediate nonhomologous C_μ_−to-C_δ__ CSR (18). Alternatively, virtually identical I_μ_ and Σ_μ_ intronic DNA regions located 5′ of C_μ_ and C_δ_ exons, respectively, could mediate homologous C_μ_−to-C_δ__ CSR (18). S_μ_−to-σ_5 CSR requires AID because HIGM2 patients with AID deficiency are completely devoid of IgD^+IgM^− plasmablasts (32). In addition, naive B cells from HIGM2 patients are unable to undergo S_μ_−to-σ_5 CSR when stimulated with appropriate stimuli in vitro (32). HIGM1 and HIGM3 patients with deleterious substitutions of CD40L and CD40, respectively, and CVID patients with deleterious substitutions of TACI have a reduced and yet detectable fraction of IgD^+IgM^− plasmablasts in mucosal sites (32), suggesting that IgD CSR and production proceed through both TD and TI pathways (**Figure 2e**). Consistent with this possibility, CD40L, BAFF, or APRIL can induce IgD CSR and production when combined with IL-15 plus IL-21 or IL-2 plus IL-21 (32).

IgD^+IgM^− plasmablasts are biased toward the use of IgA light chain, harbor hypermuted V(D)J genes, and release polyreactive as well as monoreactive IgD (32, 142, 143). Secreted IgD would exert its protective function not only by binding to antigen, but also by interacting with innate immune cells, including basophils (18, 32). By arming basophils with IgD highly reactive against respiratory bacteria, mucosal IgD^+IgM^− plasmablasts may educate our immune system as to the antigenic composition of the upper respiratory tract (18). Upon sensing respiratory antigen, IgD-activated basophils would initiate or enhance innate and adaptive immune responses both systemically and at mucosal sites of entry (18). This possibility is consistent with recent evidence showing that activated basophils can migrate to secondary lymphoid organs to initiate Th2 and B cell responses (59, 61, 63, 146).
IgD

Antimicrobial factors

Migration to upper aerodigestive mucosa and blood

Circulating basophil

Salivary glands

Nasal mucosa

Lachrymal gland

Salivary glands

Tonsil (normal)

IgD DAPI

Tonsil (HIGM1)

IgD DAPI

Tonsil (PFAPA)

IgD DAPI

Nasal mucosa (normal)

IgD DAPI

Tonsil (normal)

IgD AID BAFF

Lumina

Mucosal infiltration

CD40

CD40L

MHC

TCR

Bacteria

IgM+IgD+

B cell

IgM+IgD+

B cell

CD40

CSR, SHM

FOLLCLE

LUMEN

FOLLCLE

Migration to upper aerodigestive mucosa and blood

Circulating basophil

Antimicrobial factors

Mucosal infiltration

IgD

IgM– IgD+

B cell

IgM– IgD+

B cell

TH

TH

Migration to upper aerodigestive mucosa and blood

Circulating basophil

Antimicrobial factors

Mucosal infiltration

IgD

IgM– IgD+

B cell

IgM– IgD+

B cell

TH

TH

Migration to upper aerodigestive mucosa and blood

Circulating basophil

Antimicrobial factors

Mucosal infiltration

IgM

IgM

IgD

IgD

IgM

IgM

IgD

IgD
COMPLEXITY OF MUCOSAL HUMORAL IMMUNITY

Despite a wealth of sensing and effector mechanisms capable of triggering inflammation in response to microbial sensing and intrusion, our mucosal immune system establishes homeostatic conditions based on a fine discrimination between commensals and pathogens (3). Epithelial cells play a key role in this process by sensing bacteria through a complex arsenal of pattern-recognition receptors, including TLRs (1). These innate antigen receptors educate the immune system as to the composition of the local microbiota and thereafter instruct the generation of effector and regulatory lymphocytes whose main function is to dampen inflammation and elicit massive IgA production (5).

A key aspect of this response relates to the intertwined nature of the signaling networks involved in the induction of mucosal IgA. In these networks, the adaptor protein MyD88 seems to form a critical hub because its deletion leads to a profound impairment of intestinal IgA responses, at least in mice (46, 106). This finding likely relates to the fact that TLRs are critical for the activation and differentiation of multiple immune and nonimmune cells involved in the production and release of IgA, including B cells, T cells, DCs, FDCs, and IECs. Yet some studies suggest that mucosal signaling pathways may be even more interconnected and integrated than currently thought.

As discussed earlier, TLR signaling via MyD88 induces IEC, DC, and FDC production of BAFF and APRIL, two innate factors that deliver IgG and IgA CSR signals by engaging TACI on B cells (40). In addition to generating TACI ligands, TLRs cooperate with TACI to optimize IgA and IgG CSR and production (79, 111, 147). Such cooperation involves upregulation of TACI expression on B cells (148). However, TLRs and TACI further cooperate at the signaling level (Figure 3). Indeed, TACI engagement triggers recruitment of MyD88 to a highly conserved cytoplasmic domain of TACI distinct from the cytoplasmic TIR domain of TLRs (85). Interaction of TACI with MyD88 is followed by activation of a TLR-like pathway that elicits AID expression and CSR via NF-κB (85). These findings could provide an alternative explanation to previously published in vivo data demonstrating an essential role of MyD88 in systemic TI IgG responses induced by BAFF (46, 106, 149, 150), suggest that TACI and TLRs may converge on MyD88 to generate mucosal IgA.

Figure 2
IgD responses in the aerodigestive mucosa. (a) Scheme of human NALT, including tonsillar mucosa. (b) Immunofluorescence analysis of nasal and tonsillar mucosal surfaces from healthy, HIGM1, and PFAPA (periodic fever-aphthous stomatitis-pharyngitis-cervical adenitis) donors stained for IgD (green), AID (red), and BAFF or nuclei (DAPI staining, blue). Dashed lines demarcate follicles. Original magnification, ×10. (c) Scheme of mucosal IgD responses. Antigen-sampling DCs initiate IgD CSR by activating follicular or extrafollicular B cells through T cell–dependent (CD40L, IL-2, IL-15, IL-21) or T cell–independent (BAFF, APRIL, IL-15, IL-21) pathways, respectively. The resulting plasmablasts secrete IgD reactive against respiratory bacteria that exert protective functions either locally or systemically by interacting with an elusive IgD receptor (IgDR) on circulating basophils. In the presence of IgD-binding antigens, basophils migrate to systemic or mucosal lymphoid tissues, where they enhance immunity by releasing antimicrobial factors as well as B cell–stimulating and proinflammatory mediators such as BAFF, IL-4, IL-1β, and TNF. As compared to tonsil tissues of healthy subjects, there are decreased (and yet detectable) numbers of IgD class switched (IgD⁺ IgM⁻) plasmablasts in follicular and extrafollicular areas in tonsils of patients with Hyper-IgM syndrome type 1 (HIGM1) caused by loss-of-function mutations in the CD40L gene. Increased numbers of IgD class switched (IgD⁺ IgM⁻) plasmablasts are found in tonsils of a patient with PFAPA syndrome, with increased levels of IgD in tonsillar epithelium. (Additional abbreviations used in figure: APRIL, a proliferation-inducing ligand; BAFF, B cell–activating factor; CSR, class switch recombination; NALT, nasopharynx-associated lymphoid tissue; SHM, somatic hypermutation; TNF, tumor necrosis factor.)
Interconnectivity of signaling pathways emanating from TLRs and TACI. DCs activate B cells by releasing BAFF, APRIL, and cytokines upon sensing microbial TLR ligands. Engagement of TACI by BAFF and/or APRIL triggers association of the adaptor MyD88 to a TACI highly conserved (THC) domain that activates NF-κB via IRAK-1, IRAK-4, TAK-1, and IKK-mediated degradation of IκB. Additional NF-κB activation involves binding of TRAF2, TRAF5, and TRAF6 to a TRAF-binding site (TBS) in the cytoplasmic domain of TACI and calcium modulator and cyclophilin ligand (CAML), a transmembrane TACI-interacting protein. NF-κB initiates class switch recombination (CSR) by binding to κB motifs on AICDA and C4 gene promoters. Engagement of TLRs by microbial ligands enhances CSR through a TIR-dependent pathway that shares MyD88 with the TIR-independent pathway emanating from TACI. Further CSR-inducing signals are provided by cytokines via signal transducer and activator of transcription (STAT) proteins that bind to γ-interferon-activated site (GAS) motifs on AICDA and C4 gene promoters. (Additional abbreviations used in figure: APRIL, a proliferation-inducing ligand; BAFF, B cell–activating factor; DC, dendritic cell; IKK, IκB kinase; IRAK, IL-1 receptor–associated kinase; TACI, transmembrane activator and calcium modulator and cyclophilin ligand interactor; TAK, TGF-β-activated kinase; TIR, Toll-interleukin-1 receptor; TLR, Toll-like receptor.)
CONCLUSIONS

The past decade has seen copious new discoveries regarding the regulation and function of mucosal antibodies and on the lineage, functional heterogeneity, and plasticity of mucosal cell types with B cell–modulating and antibody-inducing function. In addition, it is becoming increasingly clear that mucosal antibody responses follow dynamics quite different from those characterizing systemic IgG responses. For instance, intestinal IgA responses show additive increases after each antigenic challenge instead of prime-boost synergistic increases typical of systemic IgG responses (43). In addition, intestinal IgA responses to a given bacterial species are rapidly attenuated by colonization with a different species (43). Such IgA attrition could reflect the need of the gut immune system to rapidly adapt itself to the dominant microbial species present in the lumen at any given time (43). The lack of cardinal IgG memory characteristics in intestinal IgA responses has important implications with respect to the development of effective mucosal vaccines. One prediction is that induction of long-lasting IgA-mediated protection will require the development of creative vaccine delivery strategies capable of ensuring sustained stimulation of mucosal B cells, including the embedding of appropriate immunogens in stable components of our microbiota, edible probiotic bacteria, or genetically modified foods such as transgenic plants. Another critical step toward the generation of an effective mucosal vaccine is to acquire more detailed knowledge of the multiple pathways involved in mucosal antibody responses. A better understanding of these pathways will be critical in devising mucosal vaccines that can provide rapid, robust, and sustained protection without causing excessive inflammation or inappropriate tolerance.

DISCLOSURE STATEMENT

The authors are not aware of any affiliations, memberships, funding, or financial holdings that might be perceived as affecting the objectivity of this review.

ACKNOWLEDGMENTS

The authors are supported by U.S. National Institutes of Health grants R01 AI-05753 and R01 AI-074378 (to A. Cerutti), Ministerio de Ciencia e Innovación grant SAF 2008-02725 (to A. Cerutti), funds from Catalan Institute for Research and Advanced Studies (to A. Cerutti), funds from the Municipal Institute of Medical Research Foundation (to A. Cerutti), and a Sara Borrell fellowship (to A. Chorny).

LITERATURE CITED


90–91. Demonstrate that intestinal Treg cells induce IgA production.

106. Shows that a subset of intestinal DCs promotes IgA production in Peyer’s patches and lamina propria through nitric oxide.


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