Mouse Paneth Cell Secretory Responses to Cell Surface Glycolipids of Virulent and Attenuated Pathogenic Bacteria

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Mouse Paneth cells respond to bacteria and bacterial cell surface antigens by discharging secretory granules into the lumen of small intestinal crypts (T. Ayabe et al., Nat. Immunol. 1:113–118, 2000). To investigate mechanisms regulating these responses, purified surface glycolipid molecules with known acyl chain modifications and attenuated properties were tested for the ability to stimulate Paneth cell secretion. The antigens included lipopolysaccharide (LPS) from wild-type and msbB-null Escherichia coli and phoP-null and phoP-constitutive Salmonella enterica serovar Typhimurium strains, as well as LPS, lipid A, and lipoteichoic acid from Pseudomonas aeruginosa and Listeria monocytogenes grown in Mg2+-limited media. Measurements of total secreted protein, secreted lysozyme, and the bactericidal peptide activities of collected secretions showed that the purified antigens elicited similar secretory responses from Paneth cells in mouse crypts ex vivo, regardless of glycolipid acyl chain modification. Despite their impaired Tlr4 pathway, Paneth cells in ex vivo C3H/HeJ mouse crypts released equivalent amounts of bactericidal peptide activity in response to purified bacterial antigens, including lipid A. Thus, mouse Paneth cells respond equivalently to purified bacterial cell envelope glycolipids, regardless of functional Tlr4, the structural properties of glycolipid acyl chains, or their association with virulence in humans.

The α-defensins are 3- to 4-kDa microbicidal peptides expressed by cells of myeloid origin and by epithelial lineages in mammals (30, 58). These cationic, β-sheet-containing peptides contain a defining trisulfidate array and amphipathicity that contributes to peptide bactericidal activity by inducing target cell membrane disruption (23, 47, 53). In cells of myeloid origin, fully processed α-defensins accumulate in azurophilic granules of phagocytic leukocytes, from which they function in microbial cell killing by nonoxidative mechanisms following phagolysosomal fusion (17). Epithelial cells on mucosal surfaces also express α- or β-defensins that may be secreted by apparent constitutive means or as granule constituents of regulated secretory pathways (39, 40, 44, 52). In mouse small intestinal epithelium, exocytotic Paneth cells located at the base of the crypts of Lieberkühn release activated α-defensins, termed cryptdins, in response to cholinergic and bacterial stimuli (2, 3, 48, 51).

Paneth cells secrete granules in a dose-dependent manner following interaction with bacterial antigens and in response to pharmacologic stimulation (2, 49–51). Live gram-negative and gram-positive bacteria and also commercially available preparations of lipopolysaccharide (LPS), lipoteichoic acid (LTA), lipid A, and muramyl dipeptide all induced rapid secretion by mouse Paneth cells. In contrast, mouse Paneth cells are unresponsive to Candida albicans, Cryptococcus neoformans, and trophozoites of Giardia lamblia (2). Both carbamyl choline and bacterial antigens stimulate Paneth cells in isolated mouse small intestinal crypts to secrete by inducing an increase in cytosolic [Ca2+] by the sequential mobilization of intracellular and extracellular Ca2+ stores (3, 49). The inhibition of Paneth cell secretion by highly selective blockers of mIKCa1, a Ca2+-activated K+ channel (3), provided evidence that bacterial antigens also induce secretion by modulating cytosolic Ca2+ dynamics.

Bacteria regulate and modify lipid A, the glycolipid anchor of their cell surface LPS, in response to the host microenvironment. For example, growth under Mg2+ limitation results in a modification of lipid A acylation in several bacterial species (13, 19–21), and the extent of lipid A acylation modulates LPS-mediated bacterial recognition when human host cells are exposed to viable bacteria. pagP, a gene activated by the phoP-phoQ regulon of salmonellae, increases lipid A acylation (22, 59), and pagP mutants show increased outer membrane permeability to α-helical peptides (22), suggesting that lipid A modification may influence sensitivity to endogenous cationic antimicrobial peptides, including α-defensins. Also, the prevailing lipid A synthesized by wild-type PAK and PAO-1 strains of Pseudomonas aeruginosa grown in high concentrations of Mg2+ is a penta-acylated form, and growth of P. aeruginosa strains in low concentrations of Mg2+ results in lipid A modification that contain aminoarabinose (4-aminoc-4-deoxy-l-arabinose) on the 1′ and/or 4′ phosphates and also palmitate at the 3′-3-oxo-C10:0 position (31, 32). In addition, the bacteria that colonize the human gingival crevice also produce modified cell surface glycolipids with attenuated proinflammatory and chemotactic properties (8, 9).
This study was initiated to test whether bacterial antigens with modified lipid A or lipoateichoic acid moieties would elicit differential secretory responses in Paneth cells. Paneth cells in isolated mouse small intestinal crypts were exposed to a variety of purified gram-positive and gram-negative bacterial cell surface antigens containing well-characterized glycolipid modifications. The data show that Paneth cells respond to effective doses of all structurally distinct antigens to approximately the same extent.

MATERIALS AND METHODS

Purification of bacterial antigens. LPS and LTA samples were isolated by using the Mg2+/H9004 precipitation procedure of Darveau and Hancock or by using the phenol extraction procedure (11). In addition, LPS preparations were subjected to Folch extraction (15) to remove residual lipid contamination and the procedure of Manthey and Vogel to remove LPS contaminating proteins (24, 33). Preparations were tested for activity in 293 cells transfected with human or mouse TLR2 to exclude contaminating bacterial pathogen-associated molecular patterns.

Preparations of LPS. LPS preparations from P. aeruginosa came from the following sources: P. aeruginosa grown in a low concentration (8 μg/mL) of Mg2+/P; P. aeruginosa grown in a high concentration (1 mM) of Mg2+/P. P. aeruginosa wild-type (Con) serovar Typhimurium, which was generously provided by Nancy Freitag (University of Washington). Salmonella enterica serovar Typhimurium was from the following sources: phoP-constitutive [phoP(Con)] serovar Typhimurium, which was isolated from clinical samples, was grown in LB, hemocytometry. The small intestine was removed from adult mice euthanized by CO2 inhalation, the lumen was rinsed with water, and crypts were eluted from segments everted and shaken in Ca2+- and Mg2+-free phosphate-buffered saline (PBS) containing 30 mM EDTA (2). Villi and crypts eluted during 5-min intervals were deposited by centrifugation at 700 × g and resuspended in PBS, and the numbers of crypts in enriched fractions were estimated by hemocytometry. Crypts were resuspended in isotonic piperazone-N,N'-bis(2-ethanesulfonic acid) (PIPES) buffer consisting of 10 mM PIPES (pH 7.4) and 137 mM NaCl (iPIPES) for exposure to microbial antigens. All procedures performed on mice were performed in compliance with the policies of the Institutional Animal Care and Use Committee of the University of California, Irvine.

For preparation and isolation of single Paneth cells, crypt preparations were incubated in 2 ml of Hanks balanced salt solution with 150 U of collagenase (Sigma-Aldrich, St. Louis, Mo./ml) at 37°C for 15 min. Cells were deposited by centrifugation at 1,500 rpm for 5 min in a Beckman GS-6R centrifuge, resuspended, and washed twice with ice-cold PBS, and the preparation was centrifuged through a 30-μm-pore-size filter Cell Strainer cap (Falcon model 352235). Intact crypts, villi, Paneth cells, and agranular epithelial cells were identified under phase microscopy, drawn into capillary tubes, and transferred to PCR tubes for amplification.

For each series of antigens, preparations of crypts, numbering 2,000 to 4,000 crypts, were divided equally, resuspended in 500 μl of nominal Ca2+/Mg2+-free isotonic PIPES buffer, and exposed to antigens (100 ng of LPS/ml, 10 μg of LTA/ml, and 1 μg of lipid A/ml) for 30 min at 37°C. After incubation, secretions were obtained by the collection of supernatants from crypts deposited by low-speed centrifugation (2). Protein concentrations were determined by using the Bradford assay, and laosymy activity was measured with the Micrococcos luteus turbidometric assay (18, 42). For each set of determinations, a standard curve was determined by using hen egg white lysozyme (Sigma) in sample tubes containing 175 μl of Micrococcos lysodeikticus (Sigma) suspended in PBS at 200 μg/ml, and activity was measured as the absorbance at 530 nm at 25°C. Dilutions of collected crypt secretions were assayed similarly, and the laosymy concentrations in samples of secretions were extrapolated from the linear region of the standard curves.

RT-PCR detection of Toll-like receptor mRNA's in mouse Paneth cells. For reverse transcription-PCR (RT-PCR) experiments, isolated villi, intact crypts, individual Paneth cells, or individual agranular crypt epithelial cells were transferred to individual microfuge tubes, sonicated in 175 μl of 1:100 dilutions of the primary RT-PCR products were used as templates for nested amplification using primer set 1 (Table 2). PCR mixes lacking AmpliTaq DNA polymerase were preheated at 94°C for 2 min, and then complete reaction mixes were run for 40 cycles as described before, except for

### TABLE 1. Primers for primary TLR amplification

<table>
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<tr>
<th>Receptor</th>
<th>Primer (sequence)</th>
<th>Accession no.</th>
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<td>mMD2</td>
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<td>mCD14</td>
<td>mCD14-S50 (5'-ATGGGTTGTTGCTTCTCGG)</td>
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<td>mTLR1</td>
<td>mTLR1-S50 (5'-AGTTGCTTCTCTCGGAGT)</td>
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<td>mTLR2</td>
<td>mTLR2-S330 (5'-ATGGGTTGTTGCTTCTCGG)</td>
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<td>mTLR3</td>
<td>mTLR3-S880 (5'-ATCTCTCGTTCACACACCG)</td>
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<td>mTLR4</td>
<td>mTLR4-S82 (5'-ACACCGAAGCCTTGAATCC)</td>
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<td>mTLR5</td>
<td>mTLR5-S225 (5'-ACACCGAAGCCTTGAATCC)</td>
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<td>mTLR6</td>
<td>mTLR6-S223 (5'-ATGTCTCAGTGGATGCTGAG)</td>
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<td>mTLR7</td>
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<td>mTLR8</td>
<td>mTLR8-S862 (5'-CTCTACATCCATCACATAC)</td>
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<tr>
<td>mTLR9</td>
<td>mTLR9-S999 (5'-AAGTGGTCTGTGCTCACTCC)</td>
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a The primer pairs shown were used to amplify TLR, MD-2, and CD14 mRNAs by RT-PCR from samples (500 ng) of total RNA from mouse bone marrow, stomach, small intestine, and colon and from isolated villi, intact crypts, individual Paneth cells, or individual agranular crypt epithelial cells (see Materials and Methods).

Lm10403S-L. LTA from L. monocytogenes 10403S cells grown in 8 μM Mg2+.

Lm10403S-H. LTA from L. monocytogenes 10403S cells grown in a high concentration (1 mM) of Mg2+.

Preparation and analysis of isolated mouse mumps and Paneth cells. Small intestine was removed from adult mice euthanized by CO2 inhalation, the lumen was rinsed with water, and crypts were eluted from segments everted and shaken in Ca2+- and Mg2+-free phosphate-buffered saline (PBS) containing 30 mM EDTA (2). Villi and crypts eluted during 5-min intervals were deposited by centrifugation at 700 × g and resuspended in PBS, and the numbers of crypts in enriched fractions were estimated by hemocytometry. Crypts were resuspended in isotonic piperazone-N,N'-bis(2-ethanesulfonic acid) (PIPES) buffer consisting of 10 mM PIPES (pH 7.4) and 137 mM NaCl (iPIPES) for exposure to microbial antigens. All procedures performed on mice were performed in compliance with the policies of the Institutional Animal Care and Use Committee of the University of California, Irvine.

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small intestinal and bone marrow amplifications, which were run for 30 cycles. Samples of the PCRs were analyzed by separation in 2% agarose gels to visualize and to size amplification products. Amplification products purified with the QIAEX II gel extraction kit (QIAGEN, Valencia, Calif.) were cloned into the pCR2.1 TOPO vector (Invitrogen, Burlington, Calif.), and their identities were confirmed by DNA sequencing (data not shown).

**Bactericidal peptide assays.** Two types of bactericidal assays were performed on secretions collected from crypts exposed to bacterial antigens, both using the defense-sensitive ΔphoP strain of serovar Typhimurium as the test species (14, 36). In the first assay, 500 to 1,000 CFU of exponentially growing cells were deposited by centrifugation in 45 μl of PIPES buffer, and combined with 5 μl of secretion collected from crypts or villi. After 60 min at 37°C, the surviving bacteria were quantitated by plating incubation mixtures on nutrient plates, and their numbers in individual experiments varied slightly, groups were normalized by expressing bacterial cell killing as a percentage (± standard deviation) relative to bacteria incubated for 1 h at 37°C in PIPES alone. In the second assay, secretions were collected from 10^6 crys, and the bactericidal activities of samples (0.1 to 10 μl) of secretions were assayed against 5 × 10^8 ΔphoP serovar Typhimurium. Exponential-phase bacteria grown in trypticase soy broth at 37°C were deposited by centrifugation at 1,700 × g for 10 min, washed in 10 mM PIPES (pH 7.4), and resuspended in 10 mM PIPES (pH 7.4) supplemented with 0.01 volume of TSA. Test ΔphoP serovar Typhimurium cells were incubated with samples of secretions in a total volume of 50 μl for 1 h in a shaking incubator at 37°C. 20-μl samples of incubation mixtures were diluted 1:200 with 10 mM PIPES (pH 7.4), and 50 μl of the diluted samples were plated on trypticase soy agar plates by using an Autoplate 4000 plater (Spiral Biotech Inc., Bethesda, Md.). Surviving microorganisms were quantitated as CFU per milliliter after incubation at 37°C for 12 to 18 h.

**RESULTS**

**Paneth cell secretion in response to purified antigens.** When isolated mouse small intestinal crypts are exposed to live bacteria ex vivo, Paneth cells secrete bactericidal peptides that kill bacteria in the medium (2). This fact is illustrated in Fig. 1: isolated crypts or villi were exposed to ~1,000 CFU of serovar Typhimurium (ΔphoP) for 30 min and then tested for survival of the exposed bacteria. As reported previously (2, 3), bacterial cell survival was unaffected by incubation with villi (Fig. 1), but bacteria exposed to intact crypts were killed in a dose-dependent fashion (Fig. 1). A previous study has shown that the majority of the bactericidal peptide activity released by Paneth cells in response to bacteria is attributable to cryptdins (2). In these earlier studies (2, 3), LPS, lipid A, LTA, and muramyl dipeptide had been obtained from commercial sources, and the possibility that potential contaminants may have been influencing secretion was a concern.

To investigate mechanisms regulating Paneth cell secretory responses, we tested whether pure bacterial surface glycolipids with known acyl chain modifications and attenuated properties would differ in the extents to which they stimulate Paneth cell secretion. Subsequent to the exposure of the crypts to purified antigens, secretions were collected from the crypts and assayed for protein, lysozyme, and bactericidal peptide activity. In the first series of experiments, the antigens consisted of LPS molecules purified (i) from WT and ΔmsbB mutants of E. coli strains (55); (ii) from WT, ΔphoP, phoP(Con), and ΔwaaP mutants of serovar Typhimurium (57); and (iii) from clinical isolates of P. aeruginosa, as well as WT and ΔphoP strains of P. aeruginosa (13). Relative to crypts incubated under identical conditions but not exposed to antigens, Paneth cells in all crypt preparations released protein and lysozyme to similar extents in response to LPS (Fig. 2A through C). Unexposed crypt preparations secreted 50 to 100 μg of total protein/ml and 0.4 to 2 μg of lysozyme/ml, approximately 2 ng of total protein and 30 pg of lysozyme on a per crypt basis. For example, with a

**TABLE 2. Primers for nested Tlr amplification**

<table>
<thead>
<tr>
<th>Receptor</th>
<th>Sense (sequence)</th>
<th>Antisense (sequence)</th>
<th>Product size (bp)</th>
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<tbody>
<tr>
<td>mMD2</td>
<td>mMD2-S117 (5’-AGTGGTCTCTGCAACTCCTCC)</td>
<td>mMD2-AS338 (5’-GGCACAGAATCTCCTTACGC)</td>
<td>222</td>
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<tr>
<td>mCD14</td>
<td>mCD14-S317 (5’-GCGATTCTTTAGCACTGGT)</td>
<td>mCD14-AS951 (5’-GGTTCCCTACCCGCTGTG)</td>
<td>635</td>
</tr>
<tr>
<td>mTLR-1</td>
<td>mTLR1-S722 (5’-AATCCTCTGCAGCAGTATG)</td>
<td>mTLR1-AS1049 (5’-ATCTCTCTAGAAGCTTGACC)</td>
<td>328</td>
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<tr>
<td>mTLR-2</td>
<td>mTLR2-S658 (5’-TTATCTCCCCCTGGTCCGC)</td>
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<td>mTLR-3</td>
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<td>mTLR-4</td>
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<td>mTLR4-AS968 (5’-ACACCTGCGAGGACACATGG)</td>
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<td>mTLR5-AS807 (5’-GGCGACAGGCTCTGATGGTCC)</td>
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<td>mTLR-6</td>
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<td>mTLR9-AS1548 (5’-ATGATCAAGGTGTGGTACAC)</td>
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* The primer pairs shown were used in nested amplification reactions for Tlr, MD-2, and CD14 mRNAs. Samples consisted of 5 μl of 1:100 dilutions of the primary RT-PCR products (Table 1; see Materials and Methods). PCR mixes lacking AmpliTaq DNA polymerase were preheated at 94°C for 2 min, and then complete reaction mixes were run as described in Materials and Methods.
given set of antigens, e.g., LPS from *P. aeruginosa*, the preparation of crypts was divided equally among the five experimental conditions so that the quantities of peptides secreted in response to antigens are comparable within a given experiment. On the other hand, because the actual numbers of crypts differed between experiments, secretions released by crypts exposed to different antigens cannot be compared directly. Whether measured by protein secretion or lysozyme activity, WT and variant LPS molecules elicited equivalent Paneth cell secretion at levels similar to the release induced by commercial LPS. Although all bacterial glycolipids tested induced similar levels of secretion, previous studies had shown that Paneth cells in intact mouse crypts did not respond to live fungi *C. neoformans* and *C. albicans* or to live trophozoites of *G. lamblia*. Under the conditions of these assays, secretion was independent of the species from which the LPS was isolated and the extent or quality of acyl chain modification (Fig. 2A through C), but we cannot exclude the possibility that differential secretory responses may occur at submaximal antigen concentrations.

Paneth cell secretion also was measured in response to purified cell surface glycolipids from gram-negative and gram-positive bacteria grown under conditions of *phoP* activation or repression. Because activity of the *phoP*/*phoQ* regulon is associated with resistance to defensins and polymyxin (12, 22), lipid A molecules isolated from WT, *phoP*(Con), and *ΔphoP* mutants of serovar Typhimurium grown under low-Mg\(^2+\) (*phoP*-activated) or high-Mg\(^2+\) (*phoP*-repressed) conditions were tested for the ability to elicit Paneth cell secretion. At the concentrations tested, all lipid A molecules stimulated equivalent secretion regardless of acyl chain modifications associated with *phoP* regulation as judged by the quantity of protein and lysozyme released by Paneth cells (Fig. 2D). Similarly, analyses of LTA-containing glycolipid fractions from *L. monocytogenes* grown in low and high concentrations of Mg\(^2+\) also showed that the extent of Paneth cell secretion was unaffected by modifications associated with *phoP* activity (Fig. 2E).

**Bactericidal activities of induced Paneth cell secretions.** Although measurements of secreted protein and lysozyme supported the view that glycolipid modification did not affect Paneth cell secretory responses, we tested whether secretions induced by different antigens differed in their bactericidal properties. The bactericidal activities of all secretions were measured against defensin-sensitive *ΔphoP* serovar Typhimurium, and they were not distinguishable (Fig. 3). Samples (10 μl) of Paneth cell secretions induced by LPS from *ΔmsbB* strains of *E. coli* had the same bactericidal activities as those elicited by corresponding WT strains (Fig. 3A, upper panel). In addition, secretions elicited by LPS from *P. gingivalis*, a species of Mg\(^2+\) (bars 1), PAK cells grown in a high concentration of Mg\(^2+\) (bars 2), *phoP*(Con) PAK cells (bars 3), *ΔphoP* PAK cells (bars 4), and Sigma (commercial lipid A; bars 5). Also shown are results for control crypts incubated in iPIPES without antigen (bars 6). (E) Results for control crypts incubated in iPIPES without antigen (bars 6).

![FIG. 2. Protein and lysozyme secretion by Paneth cells in crypts stimulated with varied bacterial cell surface antigens.](http://iai.asm.org/)
recognized for its attenuated proinflammatory activity (4, 7, 10), were indistinguishable from secretions stimulated by LPS from B. forsythus and E. coli (Fig. 3A, lower panel). Similarly, secretions collected from crypts stimulated by serovar Typhimurium LPS had equivalent bactericidal activities, regardless of whether phoP was active or repressed (Fig. 3B).

To test whether purified glycolipids with known acyl chain modifications elicit differential Paneth cell secretion, crypts were exposed to lipid A from P. aeruginosa or LTA-containing fractions from L. monocytogenes to induce the release of Paneth cell bactericidal peptide activity (Fig. 3C). Regardless of the Mg$^{2+}$-mediated status of phoP in the cells from which the glycolipids were isolated, the overall bactericidal activities of the collected secretions were similar to each other and to secretions induced by LPS (Fig. 3B and C). Thus, by these independent criteria, purified LPS and lipid A molecules and
LTA-containing fractions induce equivalent levels of secretion from mouse Paneth cells, regardless of the covalent modifications that are associated with virulence in humans or diminished sensitivity to defensins.

**Toll-like receptor mRNAs in mouse Paneth cells.** To test for potential mediators of antigen recognition by Paneth cells, the distribution of Toll-like receptor (Tlr) mRNAs was investigated in mouse intestinal epithelial cells. By a nested RT-PCR approach, mRNAs for MD-2, CD14, and Tlr1 to Tlr9 were detected in whole-organ mRNAs from mouse bone marrow, stomach, small bowel, and colon RNAs (Fig. 4), demonstrating that all primer sets supported amplification of the target sequences. Similar nested RT-PCR amplifications of isolated intact crypts and villi showed that MD-2, CD14, and all Tlr mRNAs except Tlr4 were present in isolated crypts. Individual intact villi lacked MD-2 but appeared to express Tlr4, although the cell lineages expressing the sequences remained unknown (Fig. 5A and B). Accordingly, Tlr mRNA was assayed in triplicate pools, each consisting of five individual Paneth cells or villus enterocytes (3). Tlr1 was not found in villus enterocytes. Tlr4 was not detected in either epithelial cell population, and MD-2 was absent from two of the three replicates in each case (Fig. 5C and D). The fact that Tlr mRNAs could be detected only by a nested PCR approach is an indication of their low levels of gene expression in these epithelial cell lineages. Widespread Tlr expression by small intestinal epithelia differs from the expression of the Ca$$^{2+}$$-activated K$$^+$$ channel mIKCa1, which occurs only in Paneth cells and not in villus enterocytes (3). These data support the view that mouse Paneth cells respond to bacteria by Tlr4-independent mechanisms.

**Tlr4 is not required for Paneth cell secretion in response to bacteria.** To test whether the absence of Tlr4 gene expression in Paneth cells (Fig. 5D) meant that Tlr4 was not involved in the induction of Paneth cell secretion by cell surface antigens, secretion was tested in the Tlr4-deficient C3H/HeJ mouse strain (5, 43). As Western blot analyses of small intestinal protein extracts from C3H/HeJ and outbred Swiss mice showed, the Tlr4 mutation had no effect on overall cryptdin peptide levels (Fig. 6A). As expected, if secretions were medi-
cells in C3H/HeJ crypts should not be responsive to those antigens. However, C3H/HeJ mouse crypts released bactericidal peptide activity when exposed to lipid A, and the responses were similar to secretions elicited by varied LPS molecules and LTA (Fig. 6C). Although the mechanisms by which lipid A could induce secretion from Tlr4-null cells remain to be defined, Paneth cell secretion in response to bacteria is independent of Tlr4, consistent with the absence of Tlr4 mRNA in Paneth cells.

**DISCUSSION**

In these studies, we tested the hypothesis that mouse Paneth cells would exhibit differential sensitivity to bacterial glycolipids associated with reduced human virulence and greater sensitivity to defensins. The findings show that Paneth cells in intact crypts respond equivalently to all forms of lipid A and LTA tested, suggesting that they recognize common components of these glycolipids rather than secrete differentially in response to particular acyl chain modifications. Because intact crypts were exposed to antigens, we cannot exclude the possibility that the antigens act on crypt epithelial cells, which release mediators that stimulate Paneth cells to release granules. Because secretion is rapid (having been detected within 5 to 15 min following exposure to antigen [2]), we speculate that the mechanism would involve components associated with vesicular fusion mechanisms at the apical cell surface. However, whether antigens interact with Paneth cells directly or at the apical or basolateral surface remains to be established.

Paneth cells release α-defensins and additional secretory granule constituents in response to pharmacologic stimulation and exposure to bacteria and bacterial antigens (2). In mouse small intestinal crypts stimulated with carbamyl choline, the cytosolic calcium dynamics change only in Paneth cells and in a biphasic pattern consistent with the mobilization of intracellular Ca$^{2+}$ stores, followed by the influx of extracellular Ca$^{2+}$ (49). Also, the mouse Ca$^{2+}$-activated K$^+$ channel mIKCa1 modulates mouse Paneth cell secretion, because channel blockade with highly selective mIKCa1 inhibitors diminished Paneth cell secretion in intact crypts by ~50%. In mouse small intestines, the expression of mIKCa1 is restricted to Paneth cells, identifying the channel as a functional Paneth cell marker that appears to regulate the influx of extracellular Ca$^{2+}$ essential for sustaining the secretory response. In preliminary studies of LPS-induced secretion from crypts of mice null for the phosphoinositide 3-kinase p85-α subunit (16, 56), defective crypts released bactericidal peptide activity at levels similar to those of isogenic control crypts, suggesting that secretion in response to LPS does not require phosphoinositide 3-kinase (T. Ayabe, D. Fruman, and A. J. Ouellette, unpublished data). Thus, the responses that we have characterized to pure cell surface glycolipids from several bacterial species must initiate increased cytosolic Ca$^{2+}$, but, despite the exclusion of Tlr4-mediated signaling, those mechanisms remain obscure. Recently, mouse Paneth cells were shown to be immunopositive for Tlr9 (45), immunolocalizing Tlr9 to apical secretory granules. The injection of mice with CpG-containing oligonucleotides downregulated Tlr9 and appeared to stimulate Paneth cell degranulation, although the effects of CpG injection may have been an indirect action rather than a direct action on

**FIG. 6.** Tlr4-null Paneth cells from C3H/HeJ mice release bactericidal peptide activity in response to bacterial antigens. (A) Intestinal proteins (≥10 kDa) prepared from outbred Swiss (lane 1) and C3H/HeJ (lane 2) mice by Centricon 30 centrifugal filtration were separated by acid-urea polyacrylamide gel electrophoresis, transferred to a nitrocellulose membrane, and probed with anti-cryptdin-1 rabbit serum (2, 3). Arrows at the left denote the presence of immunoreactive procryptdin (upper arrow) and mature cryptdins 1 through 3 and 6 (lower arrow), respectively. (B) Approximately 5,000 crypts were incubated with 10$^3$ CFU of live P. aeruginosa PAK cells grown in 8 μM MgCl₂ (bar 1), LPS (100 ng/ml) from P. aeruginosa Δphop PAK cells grown in 8 μM MgCl₂ (bar 2), lipid A (1 μg/ml) from Δphop serovar Typhimurium (bar 3), lipid A (1 μg/ml) from Δphop serovar Typhimurium (bar 4), LTA (10 μg/ml) from WT L. monocytogenes Scott A-M (see Materials and Methods) (bar 5), commercially available LPS (Sigma, 100 ng/ml) from serovar Typhimurium (bar 6), and control crypts incubated in iPIPES without antigen (bar 7).
Paneth cell Tlr9. Also, the finding that C3H/HeJ mouse crypts are responsive to pure LPS and lipid A molecules was unexpected but consistent with the lack of Tlr4 mRNA in Paneth cells and also excludes Tlr4-mediated signaling in Paneth cell responses to lipid A. In mice and humans, the NOD2 cytoplasmic muramyl dipeptide receptor has been associated with susceptibility to Crohn’s disease (6, 28, 35, 37), and that pattern recognition receptor is localized specifically in Paneth cells (29, 38). Thus, it is possible that Paneth cells may harbor additional, as yet uncharacterized receptors that may mediate secretory responses to glycolipids. Also, the possibility that glycolipid acyl chains may disrupt Paneth cell membranes or otherwise physically induce vesicular fusion at the apical membrane cannot be excluded.

Paneth cell α-defensin levels have been found to be generally invariant and unresponsive to the presence or absence of the resident microflora (1, 25–27, 41). Oral infection of mice with WT strains of serovar Typhimurium reduces the Paneth cell α-defensin mRNA concentration to approximately 35% of the normal level, and peptide levels also appear to decline. On the other hand, attenuated ΔphoP strains of serovar Typhimurium do not diminish cryptdin mRNA levels (46). The observed effects may be mediated by direct interactions between virulent Salmonella and Paneth cells or indirectly by inducing villus enterocytes or mesenchymal cells to release mediators via a p38-signaling pathway (46). In either case, there appears to be an association between virulence and the effects on α-defensin mRNA levels. That finding is in contrast to the absence of differential effects of glycolipids associated with attenuation or virulence in humans and Paneth cell responses. Thus, our findings appear to exclude the possibility that the selective inhibition of Paneth cell secretion by WT serovar Typhimurium cell surface antigens or phoP-regulated cell envelope modifications contributes to virulence.

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REFERENCES


