First, let's identify the unstructured content:

**Research Article**

**Generation of Monoclonal Antibodies against Immunoglobulin Proteins of the Domestic Ferret (Mustela putorius furo)**

Greg A. Kirchenbaum¹ and Ted M. Ross¹²

¹Center for Vaccines and Immunology, University of Georgia, Athens, GA, USA
²Department of Infectious Diseases, University of Georgia, Athens, GA, USA

Correspondence should be addressed to Ted M. Ross; tedross@uga.edu

Received 7 November 2016; Revised 20 December 2016; Accepted 12 January 2017; Published 14 February 2017

Academic Editor: Kristen M. Kahle

Copyright © 2017 Greg A. Kirchenbaum and Ted M. Ross. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

The domestic ferret (Mustela putorius furo) serves as an animal model for the study of several viruses that cause human disease, most notably influenza. Despite the importance of this animal model, characterization of the immune response by flow cytometry (FCM) is severely hampered due to the limited number of commercially available reagents. To begin to address this unmet need and to facilitate more in-depth study of ferret B cells including the identification of antibody-secreting cells, eight unique murine monoclonal antibodies (mAb) with specificity for ferret immunoglobulin (Ig) were generated using conventional B cell hybridoma technology. These mAb were screened for reactivity against ferret peripheral blood mononuclear cells by FCM and demonstrate specificity for CD79αβ+B cells. Several of these mAb are specific for the light chain of surface B cell receptor (BCR) and enable segregation of kappa and lambda B cells. Additionally, a mAb that yielded surface staining of nearly all surface BCR positive cells (i.e., pan ferret Ig) was generated. Collectively, these MαF-Ig mAb offer advancement compared to the existing portfolio of polyclonal anti-ferret Ig detection reagents and should be applicable to a wide array of immunologic assays including the identification of antibody-secreting cells by FCM.

1. Introduction

The domestic ferret (Mustela putorius furo) serves as an animal model for the study of several viruses that cause human disease [1, 2]. Most notably, ferrets are naturally susceptible to human influenza virus and are capable of viral transmission [3–6]. Their application to influenza research began in 1933, when throat washings from human subjects were administered intranasally into ferrets [6]. These animals went on to exhibit outward symptoms of influenza including fever, sneezing, and lethargy. Transmission of influenza-like disease was also observed following transfer of nasal washings from an infected ferret or cohousing with a naïve contact. Due to the expression of both α2,6 and α2,3 sialic acid moieties along the respiratory tract, ferrets are permissive to infection with human seasonal and pre-pandemic avian influenza isolates [7]. Moreover, they recapitulate the extrapulmonary replication of highly virulent avian influenza subtypes such as H5N1 and H7N7 [8, 9]. Collectively, the ferret model has provided invaluable insights into the antigenicity, virulence and transmissibility of circulating and newly emerging influenza isolates [1, 10–12].

Despite the usefulness of the ferret animal model, the lack of ferret-specific reagents has severely hampered the ability to perform in-depth immunologic profiling. Recent studies have implemented more detailed methods for interrogating the immune response elicited in the ferret, including quantitative RT-PCR to measure cytokine and chemokine transcript levels and flow cytometric analysis of leukocyte populations utilizing cross-reactive monoclonal antibody (mAb) reagents [13–17]. However, mAb with defined specificity for B cell lineage surface markers, such as CD19 [18], that would facilitate identification of ferret B cells at various developmental stages by flow cytometry (FCM) are currently lacking. To circumvent this issue, others have identified ferret B cells on the basis of surface immunoglobulin (Ig) and/or CD79α.
expression [14, 16]. While CD79α is an excellent marker for identification of B lineage cells because it is an obligate chaperone for surface expression of B cell receptor (BCR) [19], the epitope recognized by this mAb (clone HM47) requires intracellular staining and thus does not permit isolation of viable B cells [20]. Alternatively, polyclonal antibodies with reactivity against ferret Ig are conducive for surface staining but do not exclusively define B cells due to binding of ferret Ig by myeloid cells via Fc receptors [14]. Moreover, neither approach enables segregation of ferret B cells on the basis of heavy chain usage. Collectively, the suite of available reagents is still insufficient for an in-depth characterization of the humoral immune response, and specifically the identification of antibody-secreting cells by FCM. To begin to address this unmet need, several novel mAb with specificity for ferret immunoglobulin (Ig) were generated and characterized to define their specificity.

2. Materials and Methods

2.1. Animals. BALB/c mice (female, 8–10 weeks of age) from Jackson Laboratory (Bar Harbor, ME, USA) and Fitch ferrets (Mustela putorius furo, male or female, 6 to 12 months of age) from Triple F Farms (Sayre, PA, USA) were housed in cage units and fed ad libitum. All animals were handled in accordance with protocols approved by Institutional Animal Care and Use Committees and were cared for under USDA guidelines for laboratory animals.

2.2. Isolation of Ferret Peripheral Mononuclear Cells. Peripheral blood was collected via cardiac puncture into vacuum collection tubes containing sodium heparin (Becton Dickinson, Cat #367874) and gently inverted to prevent coagulation. Blood was then combined with phosphate buffered saline (PBS) (Corning, Cat #21-040-CV) to a final volume of 35 mL and overlaid on 10 mL Ficoll-Paque PLUS (GE Healthcare, Cat #17-1440) before centrifugation at 500 × g for 25 min with the brake reduced to its lowest setting using a Sorvall Legend XTR (Thermo Scientific, Grand Island, NY, USA). Peripheral blood mononuclear cells (PBMC) at the interface were collected, washed with PBS, and then pelleted (400 × g for 10 min). Following an additional wash with PBS, total cell number and viability was determined by Trypan blue exclusion using the Countess™ (ThermoFisher, Cat #CI0227). Ferret PBMC were used immediately or resuspended in fetal bovine serum (FBS) (HyClone, Cat #SH30396.03) containing 10% DMSO (Thermo Scientific, Cat #20688) for long-term storage. Aliquots of 1-2 × 10⁷ viable ferret PBMC were stored in the vapor phase of liquid nitrogen until use and thawing of cells was according to similarly described methods [22].

2.3. Purification of Ferret Immunoglobulin. Serum from two ferrets (female, 7 months of age) was pooled and immunoglobulin (Ig) precipitated by drop-wise addition of an equal volume of saturated ammonium sulfate solution (4.1 M) (Sigma, Cat #A4418) while maintaining the solution under constant agitation at 4°C. Precipitated protein was pelleted by centrifugation at 11,500 rpm for 20 min at 4°C and then dissolved in PBS. The protein solution was then transferred into a Slide-A-Lyzer dialysis cassette (ThermoFisher, Cat #66030) and dialyzed against PBS at 4°C for three days with daily buffer exchanges. Subsequently, the protein solution was clarified by centrifugation at 6000 rpm for 10 min at 4°C and then passed through a 0.2 μm syringe filter (ThermoScientific, Cat #09-719C). This material is referred to as crude ferret Ig. Ferret Ig was further purified by affinity chromatography using Protein A/G (ThermoFisher, Cat #20423). Briefly, crude ferret Ig protein solution was applied to Protein A/G and the column (Bio-Rad, Cat #7311550) was washed by gravity flow with PBS. Fractions (2 mL) were collected and absorbance (280 nm) was monitored using a PowerWave XS spectrometer (Biotek, Winooski, VT, USA). Once wash fractions returned to baseline, ferret Ig protein was eluted by addition of 0.1 M glycine, pH 2.5 (Amresco, Cat #M103). Eluted protein was immediately neutralized with 200μl 1.5 M Tris, pH 8.8 (Amresco, Cat #M151) and protein containing elution fractions were pooled, buffer exchanged into PBS containing 0.05% sodium azide (Sigma, Cat #S2002), and concentrated using a Spin-X Ultraflow (Biotek, Winooski, VT, USA). Once concentrated, the Ig containing the Imject alum adjuvant (Thermo Scientific, Cat #20688) was included in all gels and used for molecular marker reference.

2.4. Protein Gel Electrophoresis. To assess purity of the respective ferret Ig containing protein solutions, 5 μg of crude or purified ferret Ig was loaded into Bolt™ 10% Bis-Tris Plus precast protein gels (ThermoFisher, Cat #NW00102) and resolved at 150 V for 50 min. Protein samples were diluted in either 2x Laemmli sample buffer (Bio-Rad, Cat #161-0737) with or without β-mercaptoethanol (JT Baker, Cat #P62405) or 6x SDS-sample buffer (reducing) (Boston BioProducts, Cat #BP-111R). Reduced samples were heated at 100°C for 10 min and placed on ice prior to loading. Gels were stained with PageBlue™ protein staining solution (ThermoFisher, Cat #24620) and then destained in water before imaging using the myECL Imager (ThermoFisher, Waltham, MA, USA). Spectra™ Multicolor Broad Range Protein Ladder (ThermoFisher, Cat #26634) was included in all gels and used for molecular marker reference.

2.5. Generation of Murine Monoclonal Antibodies. Female BALB/c mice were immunized with 100 μg of purified ferret Ig containing the Imject adjuvant (Thermo Scientific, Cat #77161) via the intraperitoneal route. Serum was collected via the submandibular vein on day 21 and assayed for antibody reactivity (refer to ELISA subsection). The two mice with the highest antibody titer received a booster immunization, via the intraperitoneal route, containing 100 μg of purified ferret Ig in PBS on Day 28. Three or four days after the booster immunization, splenocytes were harvested and used to perform a fusion with the SP2/O myeloma (kindly provided by Dr. Lawrence Wysocki, University of Colorado Denver) using polyethylene glycol 1450 (ATCC, Cat #50-X). Hybridomas were selected by addition of hypoxanthine (Acros Organics, Cat #1220100) and azaserine (Sigma, Cat #A4142) at a final concentration of 200 μM and 11.5 μM respectively in RPMI 1640 (Sigma, Cat #31980.02).
Cat #R6504) containing 10% FBS, 23.8 mM sodium bicarbonate (Fisher Scientific, Cat #BP328), 7.5 mM HEPES (Amresco, Cat #0485), 170 μM Penicillin G (Tokyo Chemical Industry, Cat #PI770), 137 μM Streptomycin (Sigma, Cat #S9137), 50 μM β-mercaptoethanol, and 1 mM sodium pyruvate (ThermoFisher, Cat #11360070). Eleven days after the respective fusions, culture supernatants were screen for reactivity by ELISA (refer to ELISA subsection). Positive wells were further expanded and maintained under drug selection, and hybridoma cell lines of interest were subcloned by limiting dilution. Hybridomas were subsequently expanded in media containing 5% IgG-stripped PBS (Hyclone, Cat #SH30898.03) and monoclonal antibody (mAb) purified by affinity chromatography (Protein A/G) as described above.

2.6. ELISA. The enzyme-linked immunosorbent assay (ELISA) was used to assay antibody reactivity against purified ferret Ig and to determine the IgG subclass and concentration of the respective mAb. To measure antibody binding against the purified ferret Ig antigen, CoStar high binding ELISA plates (Corning, Cat #3590) were coated overnight at 4°C with 2 μg/mL purified ferret Ig in carbonate buffer pH 9.4 containing 5 μg/mL fraction V bovine serum albumin (BSA) (Equitech-Bio, Cat#BAC69). Alternatively, plates were coated with 5 μg BSA in carbonate buffer alone. Plates were blocked with ELISA block buffer, PBS containing 0.2% BSA, 0.1% bovine gelatin (Sigma, Cat #G9391), and 0.05% Tween 20 (Sigma, Cat #P7949), for 90 min at 37°C prior to addition of culture supernatant or antibody solutions. Culture supernatants were diluted 1:2 for hybridoma screening, and purified mAb were diluted to 3 μg/mL in ELISA blocking buffer prior to 3-fold serial dilution. Plates were incubated for 90 min at 37°C and washed with PBS to remove unbound antibody. For hybridoma screening, horseradish peroxidase conjugated goat anti-mouse IgG (γ-specific) (Southern Biotech, Cat #1030-05) secondary antibody diluted in blocking buffer was added to ELISA plates. Alternatively, binding of mAb to ferret Ig was revealed by addition of horseradish peroxidase conjugated goat anti-mouse IgGl (γ1-specific) (Southern Biotech, Cat #1070-05) secondary antibody. After addition of secondary antibody, plates were incubated for 60 min at 37°C and then washed extensively with PBS prior to addition of 2.2-azino-bis (3-ethylbenzothiazoline-6-sulfonic acid) (ABTS) (Amresco, Cat #0400) substrate. Plates were incubated at 37°C for development and colorimetric conversion was terminated by addition of 5% sodium dodecyl sulfate (SDS) (Teknova, Cat #S0294). Optical density was measured at 414 nm (OD414) using a PowerWave XS spectrophotometer.

To determine the IgG subclass or determine the concentration of the respective mAb, CoStar high binding ELISA plates were coated overnight at 4°C with 1 μg/mL goat anti-mouse IgG (γ-specific) capture antibody (Sigma, Cat #MI397) in carbonate buffer pH 9.4 containing 5 μg/mL BSA. Plates were then blocked with ELISA blocking buffer for 90 min at 37°C. Culture supernatants from clonal hybridoma lines of interest, diluted mouse anti-ferret Ig (Mab-F-Ig) mAb or mouse IgGlκ (Biologen, Cat #401402), were serially diluted in ELISA blocking buffer and plates incubated for 90 min at 37°C. Plates were washed five times with PBS, horseradish peroxidase conjugated goat anti-mouse IgGl (γ1-specific) secondary antibody diluted in ELISA blocking buffer added and the plates incubated for 60 min at 37°C. Following extensive washing with PBS, ABTS substrate was added and plates incubated at 37°C for development. Colorimetric conversion was terminated by addition of 5% SDS solution, and OD414 was measured using a PowerWave XS spectrophotometer. The concentration of individual mAb was then interpolated based on a nonlinear regression of the IgGlκ standard using PRISM 6.0 (GraphPad Software, La Jolla CA, USA).

2.7. Competitive ELISA. A competitive ELISA was performed using unlabeled and biotinylated MaF-Ig mAb (refer to Protein Conjugation) to identify overlapping epitope recognition. CoStar high binding ELISA plates were coated overnight at 4°C with 2 μg/mL purified ferret Ig in carbonate buffer pH 9.4 containing 5 μg/mL BSA. Plates were then blocked with ELISA blocking buffer for 90 min at 37°C. Unlabeled MaF-Ig mAb (5–50 μg/mL starting concentration, determined using BCA assay kit) were serially diluted in ELISA blocking buffer, followed by addition of biotinylated MaF-Ig mAb, and plates incubated overnight at 4°C. Plates were washed five times with PBS, horseradish peroxidase conjugated streptavidin diluted in blocking buffer added, and the plates incubated for 60 min at 37°C. Following extensive washing with PBS, ABTS substrate was added and plates incubated at 37°C for development. Colorimetric conversion was terminated by addition of 5% SDS solution, and OD414 was measured using a PowerWave XS spectrophotometer. The percent of maximal signal was determined using the formula 100 × [OD414 experimental sample – OD414 blank/OD414 maximal signal – OD414 blank].

2.8. Western Blot. To characterize the specificity of individual MaF-Ig mAb, 1 μg of purified ferret Ig or 0.5 μl of ferret serum was reduced and resolved by protein gel electrophoresis as described previously. Protein transfer to polyvinylidene difluoride (PVDF) membranes was performed using the Trans-Blot Turbo RTA Mini PVDF transfer kit (Bio-Rad, Cat #1704272) and a Trans-Blot Turbo Blotting system (Bio-Rad, Hercules, CA, USA) according to the manufacturer's instructions. The membrane was blocked with PBS + Tween 20 (0.1% v/v) (PBST) containing 5% BSA (VWR, Cat #0332) at room temperature (RT) with constant agitation. The PVDF membranes were then cut into strips and probed with 15 mL of PBST containing 0.1 μg/mL of individual MaF-Ig mAb overnight at RT. The following day, PVDF membranes were washed three times with PBST before incubation for 60 min at RT with 10 mL PBST containing horseradish peroxidase conjugated goat anti-mouse IgGl (γ1-specific). Following extensive washing with PBS, membranes were treated with 4 mL Clarity Western ECL Substrate (Bio-Rad, Cat #1705060) and imaged using myECL Imager. Postacquisition analysis was performed using myImageAnalysis™ Software (ThermoFisher).

2.9. Flow Cytometry. To evaluate the specificity of commercially available anti-ferret Ig reagents by flow cytometry
then dG-tailed with TdT (NEB, Cat #M0315S) and dGTP using QIAquick PCR spin columns (Qiagen, Cat #28104) and using oligo-dT primer. First-strand cDNA was isolated using SuperScript III RT (ThermoFisher, Cat #18080051) and first-strand cDNA synthesis performed using oligo-dT primer. First-strand cDNA was isolated using QIAquick PCR spin columns (Qiagen, Cat #28104) and then dG-tailed with TdT (NEB, Cat #M0315S) and dGTP (ThermoFisher, Cat #10218014). Variable IgH or Igk genes were then amplified from dG-tailed cDNA templates using Phusion (NEB, Cat# M0530S). A poly-A tail was added to products following completion of the second round of PCR by addition of 5 units recombinant Taq polymerase (ThermoFisher, Cat #EP0402) directly into the reaction and incubation at 72°C for 10 min. Products from Igk PCR were further purified with QIAquick PCR spin columns before digestion with restriction enzymes PflmI (NEB, Cat #R0595S) or PfmI (NEB, Cat #R0590S) to disrupt the rearranged Vκ21-I2 gene from the SP2/0 fusion partner. After 2% agarose electrophoresis, the uncut Vκ products were isolated using the QIAquick gel extraction kit (Qiagen, Cat #27804) and eluted with autoclaved water. Variable region genes were cloned into pCR-TOPO (ThermoFisher, Cat #K4575J10) or pSC-A (Agilent, Cat #240205) plasmids according to the manufacturer's instructions. Plasmid DNA was purified using QIAprep spin columns (Qiagen, Cat #27104) and submitted to Macrogen (Rockville, MD, USA) for sequencing. Heavy and kappa variable region genes were identified using IMGT V-Quest [21].

2.12. Statistics. Statistical analyses were performed using PRISM 6.0.

3. Results

3.1. Commercial Reagents against Ferret Immunoglobulin. Lack Heavy Chain Specificity. Expression of a class-switched B cell receptor (BCR), such as IgG or IgA, can be used as a marker of memory B cells, while naïve B cells express an IgM BCR [25]. As a first attempt to segregate ferret B cells into naïve and memory compartments on the basis of surface BCR expression, ferret PBMC were stained with polyclonal goat anti-ferret IgM (GaF-IgM) or goat anti-ferret IgG (GaF-IgG) antisera. Additionally, the mouse anti-human CD79β mAb (clone CB3-1), which cross-reacts with ferret leukocytes (Supplementary Materials available online at https://doi.org/10.1155/2017/5874572), was included in the staining solution to identify surface BCRα⁺ cells [16]. The GaF-IgM antisera labeled ~99% of the CD79β⁺ population, while the GaF-IgG antisera contained ~66% of the CD79β⁺ population (Supplementary Materials). To extend these observations, ferret PBMC were next stained with both the GaF-IgM and GaF-IgG simultaneously. The majority of CD79β⁺ ferret PBMC exhibited staining with both the GaF-IgM and GaF-IgG reagents. Collectively, these findings indicate that surface staining with anti-CD79β enables identification of ferret B cells and currently available anti-ferret Ig reagents are insufficient to discriminate B cells on the basis of heavy chain expression.

3.2. Purification of Ferret Immunoglobulin. Ferret Ig was first crudely enriched from serum through ammonium sulfate precipitation and the resulting protein solution was predominantly IgG (Figure 1, lanes 2 and 3). Next, ferret Ig was further purified by affinity chromatography using Protein A/G. This second purification step removed the majority of contaminating proteins and produced a highly pure ferret Ig preparation (Figure 1, lanes 4 and 5). Reduction of the purified ferret...
reactivity with the light chain (Igλ) and heavy (Igμ or Igγ) chain components of ferret Ig.

3.3. Immunization with Purified Ferret Immunoglobulin. Mouse IgG2a mAb have increased nonspecific binding to ferret leukocytes relative to IgG1 (data not shown). In order to elicit an antibody response utilizing the IgG1 subclass, BALB/c mice were immunized with purified ferret Ig prepared with the Imject alum adjuvant [27]. Following a single immunization with ferret Ig and adjuvant, antigen-specific reactivity was detected by ELISA and western blot (Supplementary Materials). Moreover, reduction of ferret Ig in the SDS-PAGE enabled discrimination between antibody reactivity with the light (Igκ/Igλ) and heavy (Igμ or Igγ) chain components of ferret Ig.

3.4. Characterization of Monoclonal Antibodies by ELISA. Based on their reactivity with ferret Ig, two mice (№ 3 and № 5) were chosen for mAb generation. Eight IgG1+ mAb, collected from two independent fusions, reacted with purified ferret Ig. Each of these mouse anti-ferret Ig (MaF-Ig) mAb reacted with ferret Ig by ELISA (Figure 2). Additionally, normalization of the input IgG1 concentration highlighted the distinct binding curves of several MaF-Ig mAb.

3.5. Assessment of Monoclonal Antibody Reactivity by Western Blot. As an additional technique to further characterize the reactivity of the respective mAb with ferret Ig, individual MaF-Ig mAb were used to probe reduced ferret Ig antigen and serum from three naive ferrets in a western blot screen.

Three of the eight MaF-Ig mAb demonstrated specific reactivity with ferret Ig antigen using this assay (Figure 3). Both MaF-Ig mAb 4-B10 and 8-H9 reacted with ferret Ig light chain protein (Figures 3(b) and 3(c)). Additionally, mAb II-E3 reacted with a protein species corresponding to the Igγ chain (Figure 3(d)). Of note, these MaF-Ig mAb reacted with both purified ferret Ig antigen and serum samples, suggesting the epitopes recognized by the respective mAb are not polymorphic.

3.6. Assessment of Monoclonal Antibody Reactivity by Flow Cytometry. In spite of earlier observations suggesting that the GaF-Ig antisera was unlikely to define IgG(heavy) ferret B cells exclusively (Supplementary Materials), this reagent was incorporated into a flow cytometric screening assay to evaluate the ability of each MaF-Ig mAb to bind ferret PBMC because it enabled resolution of these cells into distinct populations on the basis of staining intensity (Figure 4(b)). The two populations of ferret PBMC that demonstrated staining with the GaF-Ig antisera are referred to as IgG(int) and IgG(heavy), respectively, for the sole purpose of detailing the staining patterns observed for the respective MaF-Ig mAb (Figure 4 and data not shown). Using this flow cytometric screening approach, both 4-B10 and 8-H9 had similar staining patterns of ferret PBMC and labeled ~50% of the IgG(heavy) population (Figures 4(c) and 4(d)). Moreover, both of these MaF-Ig mAb also reacted with a small population of IgG(heavy) cells. Three additional MaF-Ig mAb (4-E3, 4-DII, and 6-H5) had similar reactivity patterns (data not shown). A distinct staining pattern was observed using two MaF-Ig mAb (6-B5 and 6-B7) (Figure 4(e) and data not shown). Specifically, pretreatment of ferret PBMC with these mAb resulted
in the disappearance of the IgG$_{\text{hi}}$ population, likely due to BCR internalization or blocking of epitopes targeted by the GaF-Ig antibodies. Additionally, a subset of the IgG$_{\text{int}}$ population was labeled by 6-B7 on the basis of GaF-Ig staining (Figure 4(e)). Finally, the II-E3 mAb had a third staining pattern with low-level binding to IgG$_{\text{int}}$ cells but did not bind to IgG$_{\text{hi}}$ cell population.

As a next step to further characterize the reagents, directly labeled MaF-Ig mAb were used to stain ferret PBMC in combination with anti-CD79β. Based on previously observed Western blot and flow cytometric staining patterns (Figures 3 and 4), the MaF-Ig mAbs were segregated into two classifications. The first group of MaF-Ig mAb comprised 6-B5, 6-B7, and II-E3 and these mAb had a reactivity profile consistent with Ig heavy chain specificity. Despite evidence of reactivity with IgG$_{\text{hi}}$ cells in the indirect screen (data not shown), biotinylated 6-B5 did not stain CD79β$^+$ ferret B cells (Figure 5(a)). By contrast, biotinylated 6-B7 stained almost all of the CD79β$^+$ cell population (Figure 5(b)). In addition, II-E3 stained the entire CD79β$^+$ cell population with a low-level of fluorescence, as well as a small population of CD79β$^{\text{neg}}$ cells (Figure 5(c)). Similar results were also observed using these same mAb following Dylight conjugation (data not shown).

The second group of MaF-Ig mAb (4-B10 and 8-H9) exhibited a reactivity profile consistent with light chain specificity. However, double-labeling CD79β$^+$ ferret PBMC with these mAb revealed staining of distinct populations of cells (Figure 6(a)). Moreover, nearly the entire CD79β$^+$ population stained positive with either 4-B10 or 8-H9, with few cells reacting with both mAb. Collectively, these findings suggested that 4-B10 and 8-H9 define distinct populations of ferret B cells on the basis of light chain expression.

The remaining MaF-Ig mAb (4-E3, 4-DII, and 6-H5) also had light chain reactivity. These mAb were categorized on the basis of co-staining with 4-B10 or 8-H9 single-positive ferret B cells (Figures 6(b)–6(d)). Using a multilabeling approach, 4-E3 was found to co-stain with nearly the entire 4-B10$^{\text{pos}}$ population, while simultaneously not labeling 8-H9$^{\text{pos}}$ cells (Figure 6(b) and data not shown). Similarly, 8-H9$^{\text{pos}}$ cells were labeled by 4-DII or 6-H5, and these mAb exhibited minimal reactivity with 4-B10$^{\text{pos}}$ cells (Figures 6(c) and 6(d)). Of note, we routinely observed a small population of 8-H9$^{\text{pos}}$ cells that were not stained by 4-DII (Figure 6(c)). Collectively, these data indicate that 4-B10 and 4-E3 recognize a common light chain protein that is distinct from the light chain recognized by the other mouse anti-ferret light chain (MaF-IgL) mAb (8-H9, 4-DII, and 6-H5).

3.7 Identification of Overlapping Epitope Recognition through Competitive Binding ELISA. To further characterize whether the MaF-Ig mAb were recognizing overlapping or distinct epitopes on ferret Ig, competitive binding assays were performed. Specifically, unlabeled MaF-Ig mAb were used as competitors and evaluated for their ability to inhibit binding of biotinylated mAb (4-B10, 4-E3, 6-B7, 8-H9, and II-E3) to the purified ferret Ig antigen. Consistent with the double-labeling FCM studies in which 4-B10 and 8-H9 recognized distinct populations of ferret PBMC (Figure 6(a)), these MaF-Ig mAb did not exhibit inhibition of each other in the competitive binding ELISA (Figures 7(a) and 7(c)). In addition, despite recognition of the same population as assessed by double-labeling FCM (Figure 6(b)), 4-E3 exhibited only subtle inhibition of 4-B10 at the highest concentration tested (5 μg/mL) (Figure 7(a)). In stark contrast, 4-B10 competed for binding with biotinylated 4-E3 (Figure 7(b)). Moreover,
Figure 4: Flow cytometric assessment of monoclonal antibody reactivity. Reactivity of mouse anti-ferret Ig mAb with ferret PBMC was evaluated by flow cytometry. Binding of mouse Ig to ferret PBMC was revealed by secondary staining with Alexa Fluor 647 conjugated goat anti-mouse IgG (GaM-IgG). Additionally, ferret B cells were identified by costaining with biotinylated goat anti-ferret IgG, which was revealed by secondary staining with phycoerythrin conjugated streptavidin (SA-PE). (a) Reactivity of the GaM-IgG secondary antibody with ferret PBMC. (b–f) 1 μg of an IgG1 control (b) or mouse anti-ferret Ig mAb (c–f) was used for indirect surface staining of ferret PBMC. The presented data were generated using PBMC from a single ferret and are representative of two or more independent experiments that yielded similar results.

Figure 5: Surface staining with heavy chain reactive monoclonal antibodies. Biotinylated mouse anti-ferret Ig mAb were used in combination with anti-CD79β to evaluate surface staining of ferret B cells. Binding of biotinylated (a) 6-B5, (b) 6-B7, or (c) II-E3 was revealed by secondary staining with phycoerythrin conjugated streptavidin (SA-PE). Frequency of CD79β<sup>−/pos</sup> ferret PBMC that costained with the mouse anti-ferret Ig mAb are indicated. Data shown were generated using a single ferret and are representative of two or more independent experiments that yielded similar results.
4-B10 exhibited superior competition with biotinylated 4-E3 relative to the homologous unlabeled competitor. No competition of biotinylated 8-H9 was observed by any MαF-Ig mAb tested. Strikingly, even unlabeled 4-DII at 20 μg/mL failed to compete with 8-H9 despite these mAb labeling a common population of ferret PBMC by FCM (Figures 6(c) and 7(c)). Similarly, none of the MαF-Ig mAb exhibited competition with biotinylated H-E3 (Figure 7(d)). However, a number of MαF-IgL mAb were found to compete with biotinylated 6-B7, despite assignment of this mAb as heavy chain specific (Figure 7(e)). Specifically, both unlabeled 4-B10 and 8-H9 demonstrate definitive competition with biotinylated 6-B7 (Figure 7(e)). Moreover, 4-E3 and 4-DII (only at high concentrations) were also capable of inhibiting binding of biotinylated 6-B7 to the ferret Ig antigen, albeit to a less extent relative to 4-B10 and 8-H9.

3.8. Determination of Light Chain Specificity. Although the individual MαF-IgL mAb were already segregated into two distinct groups on the basis of light chain reactivity, their precise specificity remained undefined. To determine which group of MαF-IgL mAb was specific for the Igβ chain, a fluorescently labeled bacterial protein with specificity for Igγ chains from a variety of species [28, 29], Protein L, was used to label ferret B cells. Approximately 25% of CD79β+ ferret B cells stained brightly with the fluorescently conjugated Protein L reagent (Figure 7(a)) and these cells were putatively assigned as Igκ-expressing B cells. Next, these CD79β+
In this study, eight individual mAb with specificity for ferret Ig were characterized (Table 1). Five of these mAb had specificity for ferret Ig light chain, while the remaining three mAb did not recognize a distinct surface expressed heavy chain. Additionally, we generated a mAb (6-B7) that yielded surface staining of nearly all surface BCR positive cells (i.e., pan ferret Ig). Collectively, these MaF-Ig mAb offer advancement in ferret Ig detection compared to the existing portfolio of polyclonal antisera reagents. Moreover, several of these mAb (4-B10, 8-H9, and 11-E3) are likely to be applicable to a wide array of immunologic assays, including the identification of antibody-secreting cells by FCM.

Pooled serum was chosen as the source of material for subsequent purification of ferret Ig because both IgM and IgG antibody classes were abundant and this biological fluid was readily available [26]. Moreover, the purified ferret Ig preparation had the potential to elicit mAb with specificity for the heavy chain (Ig\(\mu\) or Ig\(\gamma\)) and the light chain (Ig\(\lambda\) or Ig\(\kappa\)) using a single antigen. However, due to the complexity of the ferret Ig antigen, antigen-specific binding in the ELISA format was insufficient to determine the precise specificity of the respective MaF-Ig mAb (Figure 2). To this end, additional screening approaches, such as FCM, western blot and competitive binding assays, were necessary to further define the specificity and categorized the individual MaF-Ig mAb.

Implementation of an indirect FCM screening method during the initial characterization enabled identification of
MaF-Ig mAb that recognized epitopes present on ferret leukocytes. Moreover, incorporation of the GaF-IgG antisera in this screen enabled resolution of ferret PBMC into distinct populations on the basis of staining intensity, and provided insight into the specificity of the respective MaF-Ig mAb (Figure 4 and data not shown). Specifically, each of MaF-IgG mAb exhibited a similar staining pattern in which approximately half of the slgG
\(^{hi}\) population was intensely labeled by the respective mAb. Moreover, these MaF-IgG mAb also labeled a small subset of the slgG
\(^{int}\) population at a similarly intense level, while the remainder of the slgG
\(^{int}\) population exhibited a low-level of staining (Figures 4(c) and 4(d) and data not shown). This staining pattern indicated that surface Ig
\(^{G}\) B cells constituted a fraction of the slgG
\(^{int}\) population and was consistent with the observation that the GaF-IgG antisera as well as other commercially available polyclonal anti-ferret IgG antisera fail to label all B cells (data not shown and [14]). Additionally, this staining pattern implied that myeloid cells constituted a portion of the slgG
\(^{int}\) population and that IgG acquired via FcR was bound in such manner that the light chain was accessible to the respective MaF-IgG mAb. Furthermore, segregation of the IgG
\(^{int}\) and slgG
\(^{hi}\) populations in the FCM screen also facilitated perception of the distinct binding specificities exhibited by the heavy chain reactive mAb 6-B7 and II-E3 on the basis of their differential reactivity profile with the slgG
\(^{hi}\) population (Figures 4(e) and 4(f)). Lastly, in the absence of the apparent disappearance of the slgG
\(^{hi}\) population that resulted following pretreatment of ferret PBMC with 6-B5 (data not shown), this MaF-IgG mAb would not have been characterized further.

Despite the perceived reactivity of 6-B5 in the antigen-specific capture ELISA and indirect FCM screening during the initial characterization, definitive staining of ferret B cells using this mAb, when directly coupled, was not observed (Figure 5(a) and data not shown). It is likely that biotinylation of 6-B5 was inefficient, rather than that conjugation destroyed the binding specificity of this mAb. Specifically, both unlabeled and biotinylated 6-B5 exhibited comparable binding to ferret IgG in an ELISA when using a goat anti-mouse IgG1 secondary reagent (data not shown). By contrast, the
same biotinylated 6-B5 exhibited a near absence of reactivity against ferret Ig when streptavidin was used to reveal binding. Consequently, it will be necessary to explore alternative chemistries for conjugation of 6-B5 to fully realize the utility of this mAb for flow cytometric applications.

The competitive binding ELISA contributed to the characterization of the respective MaF-Ig mAb. First, the data generated using this approach served to reaffirm prior observations, such as the distinction between 4-B10 and 8-H9 determined by double-labeling FCM. Second, the competitive binding and antigen-specific capture ELISA data were in agreement and reflected differences in functional affinity by several of the MaF-Ig mAb. Most notably, 4-B10 exhibited stronger avidity for the ferret Ig antigen relative to 4-E3 (Figure 2) and also demonstrated superior competition of biotinylated 4-E3 compared to the homologous mAb (Figure 7(b)). Thirdly, the competitive binding ELISA also provided additional insight into the epitopes recognized by the respective MaF-Ig mAb. While it was already presumed that the MaF-Ig mAb were targeting the constant region of kappa or lambda light chain based on their FCM staining patterns (Figure 6), it remained unclear if these anti-kappa or anti-lambda mAb were targeting distinct or overlapping epitopes. On the basis of competitive inhibition, it is likely that 4-B10 and 4-E3 are recognizing similar, but not completely overlapping epitopes on the kappa constant region since the inhibition was unidirectional (Figures 7(a) and 7(b)). By contrast, 4-D11 failed to inhibit binding of biotinylated 8-H9 despite the use of an elevated concentration (20 μg/mL), suggesting that these mAb recognize distinct epitopes on the lambda constant region. However, it remains plausible that the large difference in functional affinity between these mAb contributed to the lack of competition (Figures 2 and 7(c)). Additionally, biotinylated 4-D11 failed to produce a sufficiently high binding signal against the ferret Ig antigen to enable evaluation of reciprocal inhibition by 8-H9 (data not shown). Similar to 8-H9, no inhibition of biotinylated 11-E3 was observed using any of the MaF-Ig mAb tested, further supporting the notion that this mAb recognizes a distinct epitope (Figure 7(d)). Surprisingly, several of the MaF-Ig mAb were capable of inhibiting binding of biotinylated 6-B7 to the ferret Ig antigen. This was unexpected since 6-B7 was categorized as heavy chain reactive based on the observed FCM staining pattern (Figure 5(c)). However, the ability of 4-B10 and 8-H9, as well as 4-E3 and 4-D11 to a less extent, to compete with 6-B7 supports the conclusions that 6-B7 is recognizing an epitope that is restricted to the heavy chain. Moreover, these data suggest that 6-B7 recognizes an epitope on the heavy chain of ferret Ig that is proximal to either the kappa or lambda constant region. To this end, we hypothesize that 6-B7 would maintain reactivity with a Fab fragment of ferret Ig.

In spite of strong reactivity with purified ferret Ig by ELISA (Figure 2) and the Igγ chain by western blot (Figure 3(d)), intense staining of ferret B cells using directly labeled 11-E3 was not observed (Figure 5(c)). However, low-level staining of the entire CD79β+ B cell population, as well as a small population of non-B cells in bulk ferret PBMC, was observed using 11-E3. Collectively, these observations indicate that the epitope recognized by 11-E3 may be restricted to secreted ferret IgG. Further experimentation will be necessary to resolve the utility of 11-E3 for studying ferret B cells and derived Ig.

The assignment of lambda light chain specificity to 8-H9 was based upon several independent observations. First, the near absolute segregation between populations of 4-B10+ and 8-H9+ ferret B cells indicated that these mAb recognized fundamentally distinct light chains (Figure 6(a)). Of note, this observation also indicates that allelic exclusion of dual light chain expression is largely intact in the ferret model [30]. Secondly, 8-H9+ ferret B cells were severely underrepresented in the population of B cells that bound the kappa-specific Protein L reagent at high levels (Figures 8(a) and 8(b)). While there was a population of 8-H9+ B cells in this gate (~20%), it is plausible that these Igλ+ B cells acquired labeling with Protein L either due to inherent BCR specificity or through the association of secreted Igκ with surface expressed FcR [31, 32]. In addition, within the total B cell population, the frequency of 8-H9+ B cells detected from multiple independent ferrets closely resembled the distribution of Igλ+ B cells observed in humans [33].

While a draft sequence of the ferret genome is currently available [34], the Ig loci have not been annotated. Based on the findings presented in this report, we hypothesize that the ferret Igλ locus more closely resembles the human Igλ locus than the mouse Igλ locus due to the increase proportion of B cells that express a lambda light chain (Figure 8(c)). In laboratory mice, the Igλ locus is comprised of 3 functional Vλ gene elements that rearrange with associated Jλ-CA gene elements [35]. Additionally, in mice the prevalence of serum immunoglobulin utilizing a lambda light chain is severely reduced relative to immunoglobulin utilizing a kappa light chain [36]. By contrast, the human Igλ locus is more complex and encodes greater than 35 Vλ genes belonging to 10 subgroups [37]. Moreover, the distribution of kappa and lambda immunoglobulin is more balanced in human serum [38]. Collectively, these findings provide the first evidence that lambda light chain usage in ferrets closely resembles that seen in humans and further supports the use of ferrets for studying the antibody response elicited by influenza virus infection or influenza vaccination.

To more accurately mimic influenza infection and vaccination in humans, ferrets can be infected with various influenza viruses from different subtypes to establish a preimmune state in the animal. This model is useful to study both inactivated and live attenuated influenza vaccine (LAIV) candidates. In young children, an IgA recall response occurred following a second administration of LAIV [39]. However, ferrets previously infected with LAIV had a robust IgG antibody-secreting cell (ASC) response in the absence of an accompanying IgA ASC population following experimental challenge with seasonal influenza [40]. While it remains plausible that the immune response elicited by influenza infection of preimmune ferrets was distinct from that observed in young children following LAIV inoculation, it is more likely that the discrepancy between these two models originates from the use of polyclonal anti-ferret Ig detection reagents that possess an inherent lack of heavy chain specificity. This apparent lack of heavy chain specificity was observed...
using a variety of polyclonal anti-ferret Ig reagents (Supplementary Materials and data not shown). While mAb with specificity for discrete ferret Ig heavy chain determinants were unfortunately not generated in this study, this example serves as motivation for the continued development and characterization of ferret Ig-specific reagents.

In addition, a polyclonal goat anti-ferret Ig reagent with specificity for IgA, IgG, and IgM was used to identify ferret B cells in the context of influenza infection using FCM [14]. In this study, the polyclonal goat anti-ferret Ig (A, G, and M) failed to label a substantial population of ferret B cells that exhibited intracellular staining with the anti-CD79α mAb (clone HM47). The MxF-Ig mAb reported here are an improvement over existing reagents and enable the identification of nearly all surface Ig F-ferret B cells (Figure 5(b)) and discrimination between IgG and IgA-expressing cells (Figure 6(a)).

We envision that several of the MxF-Ig mAb generated in this study will have applications in basic science and veterinary medicine. Due to their strong affinity for purified ferret Ig and reactivity with denatured heavy (IgG or light chain proteins, αβ10, αβ9-H9, and IgM may prove to be most useful. Specifically, these MxF-Ig mAb should enable the identification of antibody-secreting cells by FCM. Additionally, these mAb are likely to have utility as secondary detection reagents in an array of assays such as ELISA, ELISPOT, and western blot. Implementation of these mAb in a variety of immunologic assays will also contribute towards the assessment of the next-generation of broadly-reactive influenza vaccines in this highly relevant model. In summary, the generation of these MxF-Ig mAb is an improvement over existing reagents available to immunologists and opens the door to more sophisticated study of ferret B cells.

Competing Interests
The authors report no conflict of interests with the results reported in this study.

Acknowledgments
This work was supported by the Vaccine and Gene Therapy Institute of Florida and the University of Georgia. Additionally, the authors thank Lawrence Wysocki for providing protocols and the SP2/0 myeloma and Rayleigh Chan and Thomas Penrose for their assistance in the production and purification of reagents used in this work.

References


Submit your manuscripts at www.hindawi.com