Specific antibody secreting cells from chickens can be detected by three days and memory B cells by three weeks post-infection with the avian respiratory coronavirus

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Abstract

Infectious bronchitis virus (IBV), the first coronavirus described, has been a continuing problem in poultry for more than 70 years. IBV, causing a highly contagious respiratory disease in chickens, resembles the recently described severe acute respiratory syndrome virus in pathogenesis and genome organization. While previous studies demonstrated that effector and memory CD8⁺ T lymphocytes are critical in controlling acute IBV infection and disease in chickens, here chicken anti-IBV antibody (IgG) secreting cells (ASC) in both peripheral blood mononuclear cells (PBMC) and spleens collected following IBV Gray infection were evaluated using an ELISPOT assay. The ASC in peripheral blood and spleens can be detected from 3 to 7 days post-infection (p.i.), which is 3–7 days earlier than anti-IBV IgG detected in the serum. The ASC frequency reached a maximum at 7–10 days p.i., and decreased more than 90% in the spleen and 70% in PBMC by 14 days p.i. The ASC levels in the PBMC then decreased gradually to 0.5 ASC/10⁶ over the next 8 weeks. The higher concentration of about 20 ASC/10⁶ cells in spleens may, at least partially, account for the presence of antibody in the serum although bone marrow ASC were not determined. In vitro stimulation of PBMC and splenocytes with IBV antigen demonstrated that memory B cells can be activated to secrete antibody by 3 weeks p.i. ELISPOT detection of primary B cells could be useful in the early detection of infection following infection with respiratory coronaviruses.

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1. Introduction

It has been recently shown that the highly contagious severe acute respiratory syndrome (SARS) in humans is caused by a coronavirus (CoV) that resembles infectious bronchitis in transmission, pathogenesis and genome structure [10,19,26]. Infectious bronchitis virus (IBV) infection causes a highly contagious respiratory disease in chickens, especially in young chicks [3,5]. The disease was first described in 1931, and has since remained a major problem in the poultry industry worldwide [3,13].

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Vaccines are available, but they are not effective long-term in controlling IBV infection, especially for variant strains. Genetic variations are common in new strains because of both point mutations and recombinants [14,15,40,41]. The many years of experience dealing with IBV should provide a valuable model for understanding SARS CoV infection in humans.

Recent studies have shown that effector CD8\(^+\) T cells are critical in controlling acute IBV infection [6, 9,30]. Adoptive transfer of T cells collected at 10 days post-infection (p.i.) protected syngenic chicks from clinical illness [30]. IBV specific memory T cells can be generated at 3 weeks p.i., and adoptive transfer of the memory T cells protected to the recipient chicks [22]. Innate immunity may also be instrumental in controlling IBV infection. Chicken interferon type I (ChIFN-I) inhibits IBV replication in vitro and in vivo [23]. Local administration of ChIFN-I inhibited IBV associated respiratory illness [23]. The importance of humoral immunity was indicated by Cook et al. (1991), who demonstrated that after IBV infection, bursectomised chicks suffered more severe and longer illness than intact chicks. The viral titers in tissues were also higher and lasted longer in bursectomised chicks than in normal chicks [7].

Individual antibody secreting cells (ASC) and activated T cells can be detected using ELISPOT assays, powerful tools for quantifying individual cell responses [1,27,28,35,42]. In the current experiments, IBV specific IgG secreting cells were detected in peripheral blood and spleens using an ELISPOT assay, while memory B cells were detected after antigen stimulation.

2. Materials and methods

2.1. Animals and virus

SPAFAS specific, pathogen-free (SPF) chickens were hatched in our laboratory and housed in an SPF environment at the Laboratory Animal Resources and Research Facility (Texas A&M University, College Station, TX). Immune chickens were generated by inoculating 1-week-old chickens with \(10^7\) EID\(_{50}\) of the IBV Gray strain by the eye–nasal routes. The virus, propagated by inoculating the allantoic sac of 11-day-old chicken embryos with the Gray strain of IBV and harvesting allantoic fluid 36 h p.i., was used for in vivo inoculation [34]. The IBV antigen used in the ELISA and ELISPOT assay was purified by polyethylene glycol (PEG) 8000 precipitation. Briefly, allantoic fluid collected at 36 h p.i. was centrifuged at 10,000 rpm for 25 min to remove any cells and cell debris. Sodium chloride (2.33%) and PEG 8000 (7%) were added to the supernatant and incubated overnight at 4 °C. The virus was collected by centrifuging at 12,000 rpm for 40 min and resuspended in PBS (pH 7.4).

2.2. Cell culture and antigen stimulation

Peripheral blood mononuclear cells (PBMC) and spleen cells were prepared at varying times p.i. [22]. Briefly, 0.5 ml of blood was collected from each of three chicks at each time point through a wing vein with a 1-ml syringe containing anticoagulant EDTA-K\(_3\) and mixed with an equal volume of PBS. PBMC were isolated by centrifuging the diluted blood for 20 min at 2000 rpm through Histopaque-1077 (Sigma). The cells collected from the interface were pooled for each group, washed three times and cultured in complete RPMI-1640 in six-well plates at a density of \(10^7\) cells/ml. Single splenocyte suspensions were prepared as described [22,23]. To detect memory responses, the PBMC and splenocytes were stimulated with UV-inactivated IBV Gray strain (\(10^{7.3}\) EID\(_{50}\)/well). One hundred microliters of cell suspension were collected at 3, 6, 9, and 12 days after stimulation. The cells were washed three times with RPMI-1640 before using in an IgG ELISPOT assay. The supernatants were saved for antibody detection.

2.3. Detection of antibody by ELISA

One-week-old chicks were infected with the IBV Gray strain through the eyes and nose. Sera were collected from three chicks at various times p.i., and antibody responses were determined using ELISA [21]. Briefly, 96-well microtiter ELISA plates (Nunc Maxisorb) were coated with 100 μl/well of purified IBV antigen, diluted in bicarbonate/carbonate buffer (2.93 g NaHCO\(_3\), 1.59 g Na\(_2\)CO\(_3\), 0.203 g MgCl\(_2\) in 11 of distilled water, pH 9.6) at a concentration of 10 μg/ml. The plates were incubated overnight at 4 °C before
blocking with 5% non-fat milk in PBS (pH 7.4) for 1 h at
room temperature or overnight at 4 °C. After washing
the plates three times with PBS (pH 7.4) and 0.1%
Tween 20 (PBS-T), the chicken sera were diluted to
1:200 in PBS-T with 5% FBS and added to the ELISA
plates. The plates were incubated at room tempera-
ture for 1 h and washed again with PBS-T. Secondary
antibody (goat anti-chicken IgG horseradish peroxi-
dase conjugate) diluted to 1:5000 in PBS-T with 5% FBS
was added and the plates were incubated at room tem-
perature for 1 h before thoroughly washing with PBS-T.
The color was developed by adding 100 µl/well of ABTS
one component microwell peroxidase substrate (Kirke-
gaard and Perry Laboratories, Inc.) and optical densi-
ties (OD) were read at 630 nm with an ELISA reader (Emax
Precision Microplate Reader, Molecular Devices
Corporation).

2.4. ELISPOT assay

Nitrocellulose bottomed 96-well multiscreen fil-
tration plates (Millipore Corporation) were coated
with 100 µl/well of purified IBV antigen (5 µg/ml)
and incubated overnight at 4 °C [31]. Plates were then
washed with PBS and blocked with 10% FCS in PBS
for 1 h at room temperature. The cells were diluted in
100 µl complete RPMI-1640 were added to each well
and incubated overnight at 37 °C, 5% CO2. The cells
were removed by rinsing with PBS, and secondary
antibody (goat anti-chicken IgG–horseradish peroxi-
dase conjugate), diluted to 1:2000 in PBS-T with 5%
FBS, was added before incubating at room tempera-
ture for 1 h. TMB membrane peroxidase substrate
(Kirkegaard and Perry Laboratories, Inc.) was used to
develop the spots representing ASC. The spots were
counted under a dissecting microscope [31]. The ASC
frequency (ASC/10^6 cells) is equal to ASC number
from infected chickens minus the ASC number from
uninfected chickens (background).

3. Results

3.1. Anti-IBV IgG responses detected in serum

It has been shown that cell-mediated immunity can
control acute IBV infection in chickens [6,9,22,29].
However, the role of humoral immune responses in
resolving IBV infection is not clearly defined [7,8,11,
20,24]. To detect B cell antibody responses, 50 one-
week-old chicks were divided into two groups.
Twenty-five chicks in Group 1 were infected with
the IBV Gray strain and 25 uninfected chicks in Group
2 were used as controls. Sera were collected randomly
from three of the 25 chicks from each group at varying
times p.i., and anti-IBV IgG was determined by
ELISA (Fig. 1). Low levels of serum anti-IBV IgG
were detected at 7 days p.i. Relatively low levels of
antibody specific for IBV were detected in all infected
chicks by 10 days p.i. However, the antibody titers
reached maximum levels at 2 weeks p.i. and remained
positive until at least week 10 p.i., the last week
examined (Fig. 1).

3.2. ASC detected in peripheral blood and spleens

To determine the time course of specific ASC in
peripheral blood and spleen, PBMC and splenocytes
were prepared from three infected and uninfected chicks
at various times p.i. Concentrations of 10^7 and
10^6 cells/ml were prepared with complete RPMI-
1640. One hundred microliters of the cell suspensions
were added to IBV antigen coated plates, and the ASC
were detected using an antibody ELISPOT assay
(Fig. 2). Low numbers of specific ASC were identified
in spleens and PBMC by 3 and 7 days p.i., respectively
(Fig. 3), 3–7 days earlier than specific IgG was detected
in sera (Fig. 1). The ASC frequencies were maximum
(592.5 ± 364.2 ASC/10^6 cells) in spleens at 7 days p.i.,
then sharply decreased more than 20-fold to 25 ± 9.9
Fig. 1. Serum anti-IBV IgG. Sera were collected from three infected
chicks at indicated times p.i. and anti-IBV IgG levels were detected
by ELISA. The values represent the means ± SD (n = 3) of OD_{630-}

ASC/10⁶ cells by 14 days p.i. This level of ASC was maintained in spleens until week 10 p.i., the last week examined (Fig. 3). The numbers of ASC in PBMC followed the same kinetics as in the spleens, but at much lower frequencies, reaching a maximum at 10 days p.i. with only 69 ± 4.2 detectable ASC/10⁶ PBMC. The levels of IBV specific IgG producing cells declined to 15 ± 7.1 ASC/10⁶ cells at 14 days p.i. and continued to gradually decrease to 0.5 ASC/10⁶ cells at 10 weeks p.i. (Fig. 3).

3.3. Memory B cells can be detected at 3 weeks p.i.

Memory B cells are identified on the basis of their ability to proliferate and differentiate, and their dependence on stimulation by antigen for recall of antibody production [31]. To detect memory B cells in IBV infected chickens, PBMC were collected from the chickens at 3 and 10 weeks p.i., and stimulated in vitro with the IBV Gray whole virus antigen. The antibody titers in the cell supernatants were detected with ELISA at 3, 6, 9, and 12 days post-stimulation (p.s.) (Fig. 4A). To determine the frequency of the ASC in the cell cultures, 10⁶ cells in 100 μl of media were collected and washed with RPMI-1640 after stimulating in vitro for 3, 6, 9, and 12 days (Fig. 4B).

PBMC collected from chickens at 3 weeks p.i. secreted IBV specific antibody even without in vitro antigen stimulation, suggesting that plasma cells were present in the peripheral blood at this time.
(Fig. 4A). However, higher levels of antibody were detected after antigen stimulation (Fig. 4A), indicating that memory B cells were generated at 3 weeks p.i. The antibody levels in the stimulated cultures reached a plateau at 9 days p.s. that lasted for at least 3 days. The ELISPOT assay indicated that the plasma cells replicated after in vitro incubation (without antigen stimulation), but died within 3 days of incubation (Fig. 4B). After in vitro antigen stimulation, ASC were detected at 3–9 days, but were not detected by 12 days p.s. The frequency of the ASC in IBV antigen stimulated cultures was similar to that in unstimulated cultures; however, the ASC survived longer in the presence of in vitro antigen stimulation (Fig. 4B). Although the numbers of ASC were detectable prior to day 9 of stimulation, the anti-IBV antibody could be detected in low levels from cell cultures at 3 days p.s. and reached a maximum after 9 days of antigen stimulation. Therefore, most of the detected antibody was likely produced and accumulated before 9 days p.s. (Fig. 4).

In contrast, antibody was not detected in the supernatants of PBMC collected at 10 weeks p.i. in the absence of IBV antigen stimulation (Fig. 4A). As expected, antibody was detected in the supernatants of cells collected from chicks at 10 weeks p.i., following in vitro stimulation with IBV antigen (Fig. 4A). The ASC frequency detected by ELISPOT assay correlated with antibody production in the supernatants with in vitro stimulation (Fig. 4A, B).

3.4. Memory B cells detected in spleens

Previous studies showed that memory T cells can be detected in peripheral blood and spleens [22]. In the present study, low frequencies of IBV specific plasma cells were detected in the spleens after the infection was resolved (Fig. 3), and the memory B cells were detected even at 10 weeks p.i. (Fig. 4). To detect memory B cells in spleens, splenocytes were collected at 10 weeks p.i. and stimulated in vitro with the IBV Gray strain antigen. The results indicated that the memory spleen cells secreted antibody only after antigen stimulation (Fig. 5A). Antibody could neither be detected from splenocytes without antigen stimulation nor from splenocytes of uninfected chickens (Fig. 5A). Furthermore, ELISPOT assays demonstrated that ASC could be detected only in antigen stimulated cultures (Fig. 5B), not in unstimulated cultures (data not shown).

4. Discussion

Effector and memory CD8$^+$ T cells play critical roles in controlling IBV acute infection [22]. However, the precise role of antibody in the control of IBV infection remains controversial. Some reports have shown that circulating antibody titer did not correlate with protection from IBV infection [11,12,24]. However, other studies demonstrated that humoral immunity plays an important role in disease recovery and virus clearance [7,37,38]. In our own studies, although the transfer of MHC compatible immune T cells eliminated initial illness and viral replication, mild illness was observed following the acute stage of infection [22,30]. Therefore, total recovery involves other mechanisms in addition to T cells. The current study revealed that the ASC frequency in spleens and peripheral blood reached maximum levels between 7 and 10 days p.i., correlating with the maximum virus load in the lungs [29], suggesting that ASC are available early in chicks after IBV infection, although their role in antibody production at this time is not know.

Previous studies have evaluated antibody and not the B cells producing specific antibody. This is the first report enumerating the effector and memory ASC specific for a respiratory coronavirus. In this study, the frequency of anti-IBV ASC in PBMC and spleens, detected by an ELISPOT assay, implicated anti-IBV IgG secreting cells and subsequent accumulation of antibody in recovery. Memory B cells in peripheral blood and spleens could be detected by 3 weeks, and as late as 10 weeks, p.i. This is also the first report of direct detection of avian memory B cells, and the evaluation of the frequencies of avian antibody secreting B cells in peripheral blood. Wu et al. detected ASC in chicken spleens after infectious bursal disease virus (IBDV) infection using ELISPOT assay [42]. Although they did not determine the dynamics of the generation of ASC after virus infection, the IgG ASC frequency at 14 days p.i. (22.67 ± 6.11 ASC/10$^6$ splenocytes) was similar to our results (25 ± 9.9 ASC/10$^6$ splenocytes).
Because by definition, memory B cells do not secret antibody without recall antigen stimulation [31], the ASC we detected in the PBMC and spleen cells without IBV antigen stimulation would be plasma cells (Fig. 2). Similar to the T cell responses described in chickens [22,29], three distinct stages can be observed in the chicken B cell response: expansion, decreases in numbers, and memory (Fig. 3). At the first stage, the IBV specific ASC expanded dramatically and reached a maximum between 7 and 10 days p.i. During this time, IBV in vivo infection increases to its maximum titer [6]. After 10 days infection, the virus was cleared by the host immune system. As in T cell responses, following antigen depletion, the IBV specific ASC response entered the second stage when the numbers decreased precipitously, perhaps as a result of apoptosis. In the third stage, a low level of residual memory B cells were generated and could be detected at 3 weeks p.i., similar to the maintenance of memory T cells [22].

It has been reported in mammals that long-lived plasma cells can be detected in bone marrow and spleen [18,33,36]. The antibody level in serum was maintained by these long-lived plasma cells [33] that survive independent of antigen stimulation [16,17]. In this study, we also detected a low level (~20 ASC/10^6) of plasma cells in chicken spleens up to at least 18 weeks p.i. (data not shown), and their survival was likely independent of antigen stimulation because IBV was cleared after 2 weeks p.i. [6]. Therefore, we presume that these cells were long-lived plasma cells as in mammals. These plasma cells could maintain the anti-IBV IgG in the serum. However, future studies should quantify the ASC, especially long-lived plasma cells, in chicken bone marrow [32].

IBV infected chicks showed the greatest clinical illness between 5 and 10 days p.i., and the viral load in the lungs reached maximum levels by 7–10 days p.i. [29]. Interestingly, the ASC frequency reached a maximum at 7–10 days p.i., while the antibody in the sera was barely detectable at 7 days p.i. and became unquestionably positive only by 10 days p.i. or beyond. When the ASC frequencies were decreasing between 10 and 14 days, the serum antibody was increasing. Therefore, serum antibody did not correlate with clinical illness or viral titers. However, the ASC frequency in this study and effector T cells shown in previous studies did correlate [6]. Although the lower levels of antibody, detected in the sera at 7–10 days p.i., could be saturated by the presence of maximum levels of virus in vivo, plasma B cells may not have matured until later. The decrease in primary ASC may be a consequence of the decreasing levels of antigen stimulation as the virus is eliminated from the host by cellular mechanisms. It is possible that continued presence of initial B, as well as T, cell numbers depends on the presence of antigen or that activated B cells have migrated elsewhere. While B cell function certainly depends on the levels of secreted specific antibody. Local specific IgA and IgM in the early infection might also be involved in controlling IBV infection, and their role should be
addressed in the future [25,29,37,39]. Furthermore, the levels of neutralizing antibody were not examined.

The recent pandemic of SARS has demonstrated that human coronaviruses are potentially a major public health problem [10,19,26]. The SARS associated clinical illness, epidemiology, and viral genome structure resemble those that have been described for IBV in chicken for more than 70 years [2,4,19,26]. IBV infection should be the most practical animal model for SARS. RT-PCR is a sensitive method for diagnosis of SARS CoV. Serological testing for specific antibody, including indirect immunofluorescence assays and ELISA, has been developed [4]. However, it typically takes 2–3 weeks for patients to produce detectable antibody. In the current study, we demonstrated that the ELISPOT assay detected ASC 3–7 days earlier than antibody. Therefore, this technique could be used in earlier diagnosis of specific infections including SARS CoV.

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