Development of a Novel In Vitro Assay (ALS Assay) for Evaluation of Vaccine-Induced Antibody Secretion from Circulating Mucosal Lymphocytes

H. SUNNY CHANG* and DAVID A. SACK†

Vaccine Testing Unit, Department of International Health, Johns Hopkins University School of Hygiene and Public Health, Baltimore, Maryland 21205

Received 24 April 2000/Returned for modification 4 December 2000/Accepted 17 January 2001

We describe here a novel method for measuring in vitro antibody secretion from the tissue culture of human B lymphocytes in peripheral blood mononuclear cells (PBMC) after oral vaccination with a killed cholera vaccine. Enzyme-linked immunosorbent assay (ELISA) titers of the antibody secreted in the cell supernatant were determined. The validation results demonstrated that human PBMC remained viable and continued to secrete antibodies (total Immunoglobulin A [IgA] and IgG) for up to 4 days of incubation at 37°C with 5% CO2 in cell cultures. The secreted antibody concentration correlated positively with the PBMC concentration and incubation time in the tissue culture and correlated negatively with the storage time of the whole blood at room temperature. In vitro assay of secreting antibody in the lymphocyte supernatant (i.e., the ALS assay) is capable of detecting specific antibody response after oral vaccination with a killed whole-cell-plus-B-subunit cholera vaccine (WC-B) in healthy adults in a phase I clinical trial. Postimmunization PBMC secreted antibodies to cholera toxin in the cell supernatants. Antibody production did not require any in vitro antigen stimulation. In the ALS assay, antigen-specific antibody titers of prevaccination samples were barely detectable, whereas serum antitoxin ELISA titers in background of prevaccine samples were significantly higher than the ALS titers. We conclude that, without any in vitro antigen stimulation after vaccination, PBMC secrete antibodies into the supernatants in the ALS assay. This assay can quantitatively measure the antigen-specific antibody production from the PBMC culture in postvaccination blood samples.

Postvaccination immunity is generally assessed via the use of antibodies in serum, but it is impossible to distinguish between recently produced antibodies and preexisting antibodies. Antibody levels in serum do not represent the latest immune responses accurately, because serum antibodies include the accumulated soluble antibodies that were induced by previous exposure to antigens.

Recent antigen exposure of mucosal T and B cells induces proliferation and differentiation of these cells (14, 25). The activated T and B cells circulate through the thoracic duct into the blood and eventually return to common mucosal sites, such as the lamina propria of the intestine, as matured plasma cells (2, 17, 20, 22, 23, 26).

To develop a sensitive surrogate for assaying local immunity, the lymphocytes traveling from local mucosal areas to the systemic blood circulation are used by methods for in vitro laboratory evaluations such as ELISPOT (6–10, 12, 15, 21; P. W. Lowry, L. M. McFarland, and H. K. Threefoot, Letter, J. Infect. Dis. 154:730, 1986). In its final step, ELISPOT measures the results of specific antibody-secreting cells (ASC) on a spot-forming gel (11–13, 15, 18; Lowry et al., letter). ELISPOT measures the number of antibody producing cells per 10⁶ PBMC following oral vaccination (11, 16). The quantification of antibodies secreted by a fixed concentration of PBMC is as important as the enumeration of ASC.

*Corresponding author. Mailing address: 10326 Champions way, Laurel, MD 20723. Phone: (410) 955-7937. Fax: (301) 604-2076. E-mail: hhz123@yahoo.com
†Present address: ICDDR, Mohakhali, Dhaka 1000, Bangladesh.
ferred to a new tube and washed with 1 X PBS. The cells were centrifuged at 1,200 rpm for 5 min in 40 ml of PBS. The cell pellet was resuspended in 10 ml of PBS. To determine the PBMC concentration, PBMC were stained with trypan blue and counted with a hemocytometer.

The cells were pelleted by centrifugation (1,200 rpm [290 × g], 5 min) and adjusted to a concentration of 10^7 cells per ml with complete RPMI 1640 medium.

Complete RPMI 1640 medium for ALS assay and T-cell proliferation. A total of 50 ml of 10% fetal calf serum (FCS; C-Six Diagnostics, Inc.), 10 ml of 2% t-glutamine (Quality Biologicals, Inc. [catalog no. 118-084-000]), and 5 ml of antibiotics (amphotericin B-penicillin-streptomycin, 1% [Mixed]; Quality Biologicals, Inc. [catalog no. 120-096-050]) was added to every 500 ml of RPMI medium using a sterile technique. The medium was filtered via a 0.22-μm-pore-size filter if a precipitate appeared. The complete medium was stored at 4°C for up to 3 days. To do the T-cell proliferation assay, the same RPMI 1640 complete medium was used except that FCS was replaced with 25 ml of human serum.

PBS overnight. The plates were then washed twice with 1 ml of PBS. The cells were centrifuged at 37°C for up to 25°C. The supernatants were collected, and the cell concentrations were adjusted to 106 cells per ml in RPMI 1640. A total of 100 μl of ALS supernatant sample with the desired dilutions was then inoculated into a 24-well cluster tissue culture and incubated at 37°C with 5% CO2 for 48 h. Cell supernatants were collected for the measurement of total IgA and IgG by ELISA.

Titration of total antibody secretion versus PBMC incubation time in the ALS assay. The effect of different incubation times of PBMC cultures on antibody production in the ALS assay was measured in freshly isolated PBMC at 10^7 cells/ml. Blood samples from two healthy adults were processed for PBMC isolation and adjusted to 10^7, 10^6, and 10^5 cells/ml. Next, 1 ml of each concentration was inoculated into a 24-well cluster tissue culture and incubated at 37°C with 5% CO2. Antibody titers were determined via ELISA for total IgA and IgG.

Measurement of total IgA and IgG secretion abilities for ALS samples processed at days 0, 1, 2, and 3 at room temperature. Titration of the effect of blood storage on the ability of PBMC to secrete the antibodies was done with blood samples from two healthy adults. The blood was stored at day 0, day 1, day 2, and day 3 at room temperature. These same aliquots of blood were processed for PBMC isolation and adjusted to 10^7 cells per ml in complete RPMI 1640 medium. Then, 1 ml of each sample in the 24-well tissue culture plate was inoculated and incubated at 37°C with 5% CO2 for 48 h. Cell supernatants were collected for the measurement of total IgA and IgG by ELISA.

Vaccine AB and consisted of 1.25 × 10^11 V. cholerae organisms. The heat-inactivated bacteria included Inaba-classical (Caio 48; 2.5 × 10^10) and Ogawa-classical (Caio 50; 2.5 × 10^10). The formalin-inactivated bacteria included Inaba ETor (Phil 6973; 5 × 10^10) and Ogawa-classical (Caio 50; 2.5 × 10^10) plus 1.0 mg of the recombiant B subunit of cholera toxin (3).

The dry vaccine was prepared from the same lot of vaccine as the liquid vaccine. To prepare the dry formulation, the same vaccine was mixed with syrup of the CeraVac buffer and spray dried. One dose contained 10 g of dry powder, which was dissolved in 200 ml of water at the time of immunization (3).

To administer the liquid formulation, 3 ml of liquid vaccine (one dose) was mixed with 150 ml of Samarín buffer in a cup and was ingested orally, according to the instructions on the packet. To administer the dry vaccine, one dose (10 g of dry powder) was mixed with 200 ml of water and ingested orally. Eating and drinking was not allowed for 1 h before and after vaccination (3).

The volunteers were randomly assigned to receive two doses of either the dry vaccine or the liquid vaccine. Six volunteers in each group. Peripheral blood was collected by using a Vacutainer on days 0, 14, 21, and 24. Samples (20 ml) of blood were obtained by using a Vacutainer into sodium citrate tubes (Blue Top; Becton Dickinson), and 10 ml without coagulant was placed in a Red Top tube and stored at 25°C.

ALS assay for V. cholerae antitoxin IgA and IgG during an oral cholera vaccine clinical trial. To apply the ALS assay for measurement of the antigen-induced specific T-cell response, PBMC samples were collected during a safety and immunogenicity trial of an oral killed cholera vaccine in healthy adult volunteers. A standard liquid formulation of the vaccine and a spray dry formulation of the vaccine were compared. The liquid formulation was stored continuously at 4°C, but the dry vaccine was placed at room temperature for 30 days. Volunteers were randomized to receive two doses of either vaccine in a double-blind manner. Healthy volunteers between 18 and 50 years of age were recruited from the Baltimore area. Each volunteer received two doses of vaccine. Serum and PBMC were collected at days 0, 14, 21, and 24 after administration of dose one. The ALS assay was used to measure antitoxin IgA and IgG. (The complete evaluation of the vaccine trial will be reported elsewhere.)
Blood samples (data not shown). Days for the total viable cell counts. Wells. The cells were harvested and counted for H3 incorporation. The results are increased linearly with time of incubation (2 to 4 days) when 10^7 PBMC inoculation concentration and incubation time. A log of the total IgA was used as a dependent variable. The period of regression relationship between antibody secretion with the variables of incubation time and PBMC concentration in the ALS assay was accessed quantitatively. The geometric mean and standard deviation were calculated using graphs. A t-test was performed (P) between day 0 and any of the other days for the total viable cell counts. Three blood samples from healthy volunteers were collected (no vaccination) and stored at 25°C. These samples were processed at day 0, day 1, day 2, and day 3, and the cell concentrations were adjusted to 10^6 cells/ml. On a 96-well tissue culture plate, 100 μl of each PBMC sample was added to each well. To stimulate PBMC, 100 μl of a 2-ng/ml concentration of ConA per ml was added to each well. The cells were harvested and counted for H3 incorporation. The results are expressed as the ratio of sample counts with ConA to corresponding control counts without ConA.

**RESULTS**

Human blood sample storage and viability counts as determined with a hemocytometer under a ×36 lens at room temperature. A 10-ml portion of blood yielded about 150×10^6 PBMC after storage at room temperature up to 48 h. By day 3, the PBMC yield dropped by about 30% on average (Table 1 and Fig. 1).

**Negative control: antibody level in PBMC.** There was no detectable antigen-specific or nonspecific IgA or IgG in 10^7 sonicated PBMC from the _V. cholerae_-vaccinated-volunteer blood samples (data not shown).

**Effect of blood storage on total IgA secretion at room temperature.** The results indicated that storage was a negative factor for IgA production in the ALS assay. After 24 h of storage, the same sample's total IgA yield dropped from 450 to 50 μg/ml in the in vitro cultures. However, samples stored for 1, 2, and 3 days produced similar and significant amounts of total IgA (50 μg/ml) in the supernatants.

The average total IgA level of two normal human blood samples, which were stored at 25°C before PBMC processing, was determined. The supernatants were taken after 96 h of incubation, and the initial concentrations were 7 logs of the cells. The total IgA levels on storage days 0, 1, 2, and 3 were 459 (range, 417 to 500), 62 (range, 62 to 62), 49 (range, 38 to 60), and 51 (range, 44 to 59), μg/ml, respectively.

**Effect of blood storage on T-cell proliferation at room temperature.** PBMC from blood samples stored up to 24 h were highly sensitive to ConA stimulation. These PBMC produced a high and sustained level of T-cell proliferation (a 33-fold increase in H3 incorporation compared to non-ConA-stimulated controls). However, after blood storage for 24 h, the T-cell proliferation ability of the PBMC dropped significantly from a titer of 33-fold to 7-fold in H3 (1/20; stock, 1 μCi) incorporation compared to non-ConA-stimulated controls (Table 1 and Fig. 2).

**Total IgA secretion increases with the increase of PMBC concentrations in the ALS assay.** When cells were incubated for 96 h in tissue culture, the total IgA production increased exponentially with the increase in the concentration of PBMC. At from 10^5 to 10^7 cells, the log of the IgA concentration was linearly related to the log of the initial cell concentration. The slope appeared to increase with higher cell concentrations (Table 2).

**Total IgA secretion increases with the increase of PBMC incubation time in the ALS assay.** Total IgA secretion increased linearly with time of incubation (2 to 4 days) when 10^7 PBMC of day 0 blood were used in the ALS assay. The slope appeared to increase with longer incubation days. The total IgA from the samples of two normal human volunteers was measured in the supernatants of an ALS assay following incubation for 2, 3, and 4 days, and the results, obtained with a 1-ml

**Table 1. Stability of human PBMC at 25°C for up to 72 h**

<table>
<thead>
<tr>
<th>Time (h)</th>
<th>Avg PBMC yielda ± SD (10^6/10 ml of blood)</th>
<th>P</th>
<th>Avg T-cell proliferationa ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>151 ± 26</td>
<td></td>
<td>33.9 ± 12.7</td>
</tr>
<tr>
<td>24</td>
<td>132 ± 46</td>
<td>0.29</td>
<td>37.6 ± 19.2</td>
</tr>
<tr>
<td>48</td>
<td>142 ± 27</td>
<td>0.35</td>
<td>6.9 ± 10.2</td>
</tr>
<tr>
<td>72</td>
<td>103 ± 57</td>
<td>0.13</td>
<td>9.8 ± 12.8</td>
</tr>
</tbody>
</table>

a Three human normal blood samples were obtained with citrate anticoagulant and stored at 25°C. Each day, 10 ml of each sample was processed for PBMC isolation up to 72 h via hemocytometer under a ×36 microscope lens. Average PBMC yields and standard deviations of the three volunteers are reported in the table. A one-tailed t test was performed (P) between day 0 and any of the other days for the total viable cell counts.

b Three blood samples from healthy volunteers were collected (no vaccination) and stored at 25°C. Each day, 10 ml of each sample was processed for PBMC isolation up to 72 h via hemocytometer under a 36 microscope lens. Average PBMC yield dropped by about 30% on average (Table 1 and Fig. 1).
portions of PBMC (10^7 cells per ml), were averaged together. The IgA levels were determined to be 116 (range, 74 to 157), 197 (range, 120 to 274), and 459 (417 to 500) mg/ml for incubation days 2, 3, and 4, respectively.

**Linear regression model.** There was a significant linear relationship between the log IgA and log cell concentrations and the incubation time. The log IgA in (micrograms per milliliter) is the dependent variable. The independent variables are the incubation time and the log cell concentration (based on day 0 storage data). The regression equation is as follows: log IgA (mg/ml) = 2.4632 + 0.424 incubation day + 0.794 log con.

According to this model, if the incubation time increases by 1 day, 0.424 log of total IgA will be secreted. If the cell concentration is increased by 1 log, 0.794 log of total IgA would be secreted. The linear regression model was highly significant, with an F score of 61.9, a P value of 0.000, and a regression coefficient of 0.89 (Table 3). The model was validated by its randomness residual distribution (Fig. 3).

**ALS IgA anti-CTB for cholera vaccine volunteers.** Immunization with either formulation of the oral killed cholera vaccine in humans induced specific IgA anti-CTB 14 days after the first dose and 7 days after the second dose in the ALS assay. The peak of the IgA ALS titer was at day 21 and started to decrease by day 24 (Table 4). The liquid formulation of the oral vaccine induced significantly higher IgA to CTB than the dry formulation in the ALS assay (Table 4).

However, the titers in serum showed a very different result from those of the ALS assay. The titers in serum showed higher titers with the dry formulation. An antibody titer continuously increased until day 24 (Table 4).

**ALS IgG anti-CTB.** In comparison to the IgA response, the ALS IgG anti-CTB response of the liquid formulation was significantly higher than that of the dry formulation. Both formulations induced significant ALS antibody response 14 days after the first dose and 7 days after the second dose. The peak of ALS IgG anti-CTB response was at day 21 and dropped at day 24 (Table 4). Similar to the IgA response, the dry vaccine induced IgG titers in serum that were higher than those with the liquid vaccine. The titers in serum continuously increased until day 24 (Table 4).

**DISCUSSION**

ALS, a specific, reliable, and accurate immunoassay, was developed for the evaluation of fresh antibody production from circulating mucosal secreting B lymphocytes. In the human trial described here, the ALS assay detected the significant antitoxin increases induced by either formulation of the oral vaccine. The ALS results indicated a peak booster antitoxin response at day 21, which is 7 days after the second dose, that started to decrease at day 24. (Complete results for this clinical trial will be reported separately.)

By assaying only antibodies secreted by circulating cells, the ALS method controlled the confounding effect of accumulative...
antibody in the serum samples, which contain both recent and preexistent soluble antibodies. Since the serum portion of the blood sample has been removed in the ALS assay, this assay measures only the secreting antibodies. When the ALS assay was performed, antibody titers from prevaccination samples were barely detectable, but background titers in serum were found in prevaccination samples. In the ALS assay, vaccine-activated mucosal lymphocytes

TABLE 4. Antitoxin response in serum and ALS assays after oral cholera vaccination

<table>
<thead>
<tr>
<th>Antitoxin assay and day</th>
<th>Dry formulation</th>
<th>Liquid formulation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>GMT (range)</td>
<td>Fold increase to day 0</td>
</tr>
<tr>
<td>Serum IgA antitoxin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>35 (6–219)</td>
<td>1</td>
</tr>
<tr>
<td>14</td>
<td>89 (23–347)</td>
<td>2.5</td>
</tr>
<tr>
<td>21</td>
<td>269 (148–490)</td>
<td>7.7</td>
</tr>
<tr>
<td>24</td>
<td>251 (132–479)</td>
<td>7.2</td>
</tr>
<tr>
<td>Serum IgG antitoxin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>120 (58–251)</td>
<td>1</td>
</tr>
<tr>
<td>14</td>
<td>191 (78–468)</td>
<td>1.6</td>
</tr>
<tr>
<td>21</td>
<td>389 (123–1230)</td>
<td>3.2</td>
</tr>
<tr>
<td>24</td>
<td>575 (263–1230)</td>
<td>4.8</td>
</tr>
<tr>
<td>ALS IgA antitoxin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>1 (0.2–1.12)</td>
<td>1</td>
</tr>
<tr>
<td>14</td>
<td>0.91 (0.43–2.34)</td>
<td>0.9</td>
</tr>
<tr>
<td>21</td>
<td>1.92 (0.66–5.75)</td>
<td>1.9</td>
</tr>
<tr>
<td>24</td>
<td>0.66 (0.55–1.02)</td>
<td>0.7</td>
</tr>
<tr>
<td>ALS IgG antitoxin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>0.28 (0.07–1.05)</td>
<td>1</td>
</tr>
<tr>
<td>14</td>
<td>0.47 (0.09–2.4)</td>
<td>1.7</td>
</tr>
<tr>
<td>21</td>
<td>1.30 (0.63–2.69)</td>
<td>4.6</td>
</tr>
<tr>
<td>24</td>
<td>0.68 (0.32–1.48)</td>
<td>2.4</td>
</tr>
</tbody>
</table>

* Anti-CTB IgG and IgA levels from volunteer sera were measured via ELISA of GM1-CTB-coated plates. Titers were determined by use of a hyperbolic curve. Then the geometric mean titer (GMT) of same-day determinations for each group of volunteers were calculated by using Excel software. The fold increase values are also reported. Anti-CTB ALS IgG and IgA levels from volunteer PBMC samples were measured via ELISA of GM1-CTB-coated plates. Titers were determined by use of a hyperbolic curve. Then, the GMT values of same-day determinations for each group of volunteers were calculated by use of Excel software. The fold increase values are also reported.
were cultured in vitro for 2 to 4 days. The secreted antigen-specific immunoglobulins in the supernatants of tissue culture were qualitatively and quantitatively measured. These PBMC were believed to be the circulating mucosal lymphocytes. ALS measures the change in host antibody response with the amount of nonstimulated, in vitro antibody produced at different postvaccination time points. This test allowed us to monitor the magnitude of the mucosal B-cell’s antibody production strength during the course of immunization.

Antibody production of lymphocytes requires multiple signals and optimal cognitive interactions, such as receptor engagement between antigen-presenting cells (APC), T cells, and B cells. The isolated PBMC layer from blood samples contains a mixture of these components. In the ALS system, antibody production is enhanced by cell concentration and incubation time synergistically. High concentrations of cells in a contained space enhances the cognitive distance of cell-to-cell interaction. As the efficiency of cell interaction increased, antibody secretion increased exponentially. When the blood samples were subjected to a long storage condition such as at room temperature for 2 days, it is possible that some key components, such as the cytokines necessary for antibody secretion, start to deteriorate. Although T-cell proliferation had been effective after 24 h of storage, the total antibody production ability dropped significantly after 24 h. Therefore, the storage of blood samples was certainly a sensitive factor for the ALS assay. If one could accurately determine which components were defective, preservation measures could be taken and/or supplements could be added to further extend the in vitro antibody secretion for a longer time period, which would have tremendous practical value for the processing of large numbers of blood samples.

Since 1963, immunoglobulin secretion at the cellular level has been assayed by hemolytic plaque assay (1). Hemolytic assay can detect cells secreting complement-binding antibodies against erythrocytes (4, 5, 24). This assay has limitations when it is applied to soluble antigens passively adsorbed to red blood cells (24). Inconsistent results had generally been associated with difficulty in coupling antigen efficiently to red cells (4, 5, 24). Additionally, the hemolytic assay did not permit quantitation of secreted molecules. ELISPOT is a qualitative assay for ASC. It requires a subjective reading of the formed spots.

The ALS assay quantifies the amount of antibody secreted and the strength of the antibody production for a fixed number of PBMC. Logistically, ALS assay does not require live bacteria during testing as the vibriocidal tests. Compared to the ASC assay, the ALS assay uses antibody supernatants of the PBMC as its final specimen rather than the PBMC. In terms of the storage of samples, cells may be stored at −70°C for up to 6 months for the ASC assay, whereas ALS supernatants can be stored at 4°C or −20°C for a much longer time. The ALS assay final result is based on readings from the ELISA reader rather than the subjective determinations of spot formation on gels in the ASC assay and turbidity in the vibriocidal assay. The major limitation of the ALS assay is the requirement of the use of fresh blood to yield a high quantity and quality of PBMC.

This assay is specifically useful for the determination of a recent immune response during vaccine trials in areas where the disease is endemic and where the population already has preexisting serum titers. It could also be used as a diagnostic method for identifying recent infections.

ACKNOWLEDGMENTS

We thank Lou Bourgeois and Janet Shimko for their excellent technical review of the manuscript and Keerti Shah, Mark Steinhoff, and W. B. Greenough III for their technical advice.

REFERENCES