Establishment of anti-HBcAg Monoclonal Antibodies for Sandwich ELISA Application by Iliac Method Utilizing Incomplete Adjuvant

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Abstract—The prevalence of hepatitis B virus (HBV) infection in Indonesia was moderate in 2013. This makes an appropriate action has to be taken immediately. In an attempt to evaluate our candidate HBV vaccine efficacy, we generated monoclonal antibodies specific to HBcAg for sandwich ELISA application for research purposes. Monoclonal antibodies are generally developed using hybridoma technology by isolating B cells from spleen or lymph nodes followed by fusion with myeloma cells. This study generated hybridoma by modifying the iliac lymph node method and used incomplete Freund’s adjuvant to emulsify the antigen. For this purpose, an eleven-week-old BALB/c mouse was immunized with a single shot recombinant HBcAg water in oil emulsion intramuscularly at the mouse tail base. After one month, besides enlargement of the medial iliac lymph nodes, we also found that the sub-iliac lymph nodes were enlarged.

In two different fusion attempts, the single B cells were then isolated from each lymph node and fused with SP2/0—Ag14 mouse myeloma cells. Hybridoma cells derived from both lymph nodes were screened by ELISA coated with HBcAg peptide. As a result, we obtained two positive wells of hybridoma polyclones from each medial iliac and sub-iliac-derived B lymphocyte. After monoclonalization, we obtained candidate anti-HBcAg monoclonal antibodies as either capture or detection antibodies. Furthermore, we successfully retrieved hybridoma clones 9.3, 9.4, and 9.5, which could produce monoclonal antibodies for sandwich ELISA application.

Keywords—HBcAg; iliac method; incomplete adjuvant; monoclonal antibodies; sandwich ELISA.

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I. INTRODUCTION

Hepatitis B is a disease generated by hepatitis B virus (HBV) infection. The HBVs target hepatocytes to replicate, which may induce liver inflammation or enlargement [1]. In 2007, the prevalence of HBV infection in Indonesia was high in endemicity based on national surveillance of hepatitis B surface antigen (HBsAg), anti-hepatitis B core antigen (anti-HBcAg), and anti-HBsAg [2]. In 2013, the prevalence of HBV decreased to 7.1% of the Indonesian population, changing the prevalence status from high to moderate [3]. HBV is transmitted by percutaneous and mucosal exposure to infectious blood or body fluids. Major modes of HBV transmission include sexual or close household contact with an infected person, the transmission of the perinatal mother to child, injection of drug use, and nosocomial infection [4].

As a disease that causes global health problems, hepatitis B can cause complications that can endanger lives [5]. Long-term hepatitis B infection may promote more serious effects such as cirrhosis, liver failure, or hepatocellular carcinoma [1], [6]–[8]. In an attempt to reduce new cases of HBV, the Indonesian government supports hepatitis B immunization which may reduce the risks of vertical transmission [9]. Evaluation of anti-HBsAg titers in younger subjects (younger than 19-year-old) was increasing among the Indonesian subjects tested, indicating successful vaccination attempts. However, decreasing titers of anti-HBcAg among those subjects may indicate the need for vaccination using HBcAg based vaccine [10].

HBsAg, which is located in the outer part of the virus, both in the mature virus and DNA-negative virus particle, serves as a common target for detecting HBV infection, determining viral load, and vaccine development [11]–[13]. It represents active viral infection [14]. However, HBsAg-negative serological samples determined by immunoassays could generate false-negative results due to possible mutations at the S gene or its promoter region [15]–[17]. In addition, those HBsAg negative samples could be detected as positive by
purposes. We modified the iliac method developed by Sado et al. [12] for patients with spontaneous HBsAg loss and the patients with HBsAg-based immunoassay for HBV detection [24], especially in combination with HBsAg to provoke better immune response by recruiting cytotoxic T lymphocytes. HBcAg peptide expressed in yeast for mouse immunization. Seventy microlitres of 2 µg/µl HBcAg was emulsified in 300 µl incomplete Freund’s adjuvant (ICFA) (Sigma). About 46 µg of peptide in a total of 100 µl emulsion was injected intramuscularly at the base tail of an 11-week-old BALB/c mouse. One month after single-shot immunization, the mouse was sacrificed to isolate both the enlarged medial iliac and sub-iliac lymph nodes. Single B lymphocytes suspension was obtained by passing the lymph nodes through a 70 µm cell strainer (Corning).

B. Cell Fusion

SP 2/0-Ag14 mouse myeloma cells were used for cell fusion. The cells were obtained from the European Collection of Authenticated Cell Culture (ECACC 85072401). The cells were maintained in RPMI medium (Sigma) supplemented with 10% FBS (Sigma) and antibiotics 100 IU/ml penicillin-100 µg/ml streptomycin in humidified 5% CO2 incubator at 37°C. Confluent cells were harvested using 0.25% trypsin/EDTA (Sigma). The B cells were fused with SP 2/0-Ag14 cells for hybridoma generation by using polyethylene glycol (Roche). Next, the fused cells were resuspended in a selection medium containing 10% FBS (Sigma), 10% BM-condimmed H1 (Roche), as well as hypoxanthine aminopterin and thymidine (HAT) (Sigma) and seeded onto 96-well plate. Non-fusion cells will not survive under an HAT-containing medium, leaving the successfully generated hybridoma to grow.

C. Screening of Hybridoma and Monoclonalization

Two weeks after hybridoma selection in HAT medium, the hybridoma conditioned medium was used for ELISA to screen the clones which produce antibodies against HBcAg. For this purpose, 50 ng/well of recombinant HBcAg peptide in sodium carbonate buffer pH 9.6 was used to coat ELISA plate and incubated overnight at 4°C. The next day, the plates were washed with PBS-T and incubated with 5% skim milk/PBS-T for one hour. Following the next washing, 20 µl of each conditioned medium was added to each well containing 80 µl PBS-T and incubated at 4°C. After 24 hours, the plates were washed and incubated with HRP-conjugated anti-mouse IgG γ-chain secondary antibody (1:5,000) for one hour. Then, the antibody solution was removed and washed with PBS-T, followed by the addition of 80 µl ABTS chromogenic substrate (Invitrogen). The absorbance of the developed color reaction was measured at OD 405 nm. The positive clones were then harvested and gradually transferred onto 12 and 6 well-plates. HAT containing medium was replaced with HT containing medium for cell recovery. To obtain monoclones, the candidate hybridoma cells were seeded back onto 96-well plate after a limiting dilution of 1 cell/well. Two weeks later, ELISA again examined the clones to retrieve monoclones that secrete anti-HBcAg antibodies. The successful clones were then propagated to collect more conditioned medium and make the frozen stocks.

II. MATERIAL AND METHOD

A. Mouse Immunization

Healthy male BALB/c mice aged 8-week with an average of 20 g body weight were obtained from PT. BLST, Bogor, Indonesia. Food and water were given to the mice ad libitum. Fitzgerald obtained a 183 aa of 19 kDa recombinant HBcAg peptide expressed in yeast for mouse immunization. Seventy microlitres of 2 µg/µl HBcAg was emulsified in 300 µl incomplete Freund’s adjuvant (ICFA) (Sigma). About 46 µg of peptide in a total of 100 µl emulsion was injected intramuscularly at the base tail of an 11-week-old BALB/c mouse. One month after single-shot immunization, the mouse was sacrificed to isolate both the enlarged medial iliac and sub-iliac lymph nodes. Single B lymphocytes suspension was obtained by passing the lymph nodes through a 70 µm cell strainer (Corning).
10% BM-condimned H1. Three days later, the conditioned medium was collected, and the cells were replenished with a new medium and continue incubated for three days to collect another conditioned medium. The collected serum-free conditioned medium was used for sandwich ELISA to retrieve candidate monoclonal antibodies which are applicable for either capture or detection antibodies. Conditioned medium from 5#21 derived monoclones were used to coat ELISA plate and represent capture antibody. After blocking with skim milk in PBS-T, HBcAg peptide was added as antigen. Then, a conditioned medium from 9#4 derived monoclones was added to represent detection antibodies. As for the control wells, the second conditioned medium was not added to the wells. To visualize the immuno-reaction, HRP conjugated anti-mouse secondary antibody was added, followed by the addition of ABTS substrate to generate a turquoise color. The signal observed by the microplate reader was analyzed by subtracting the OD405 of wells containing both conditioned medium of 5#21 and 9#4 derived monoclones with OD405 of each control well without a conditioned medium of 9#4 derived monoclones.

E. Purification of Monoclonal Antibodies by Protein A/G Agarose

Hybridoma conditioned medium was incubated with protein A/G agarose (Abcam) at 4°C at the rotating platform. One hour later, the suspension was transferred into an empty polypropylene column (Biorad), and the beads were settled down by gravity. After collecting the flow-through, the beads were washed with PBS. Next, the bound antibodies were eluted with 100 mM glycine pH 2.7 and collected in microtubes containing one-tenth of the expected final volume of 1 M Tris pH 9 to neutralize the acidic pH. To observe the purified antibodies, flow-through, washed and elution fractions were subjected to SDS-PAGE and Coomassie brilliant blue staining. Elution fractions that contained the most concentrated antibodies were then dialyzed with PBS. Finally, the antibody concentration was determined by a nano-spectrophotometer.

F. Preparation of HRP-conjugated Monoclonal Antibody

Purified monoclonal antibodies in PBS were conjugated to HRP by using an HRP-lightning link kit (Abcam). Briefly, one kit of 10 µg HRP-lightning link was mixed with 9.3, 9.4, and 9.5 monoclonal antibodies at a final volume of 20 µl. Modifier was added two microliters per 20 µl volume before the addition of HRP. After overnight incubation at room temperature in the dark, a quencher was added two microliters per 20 µl reaction. The HRP-conjugated mAbs were used for sandwich ELISA 30 minutes later.

G. Sandwich ELISA application using HRP-conjugated monoclonal antibodies

Each well of the ELISA plate was coated with 50 ng of each purified 9.3, 9.4, or 9.5 monoclonal antibodies, conditioned medium of hybridoma clone 5.2, 25 ng mouse IgG isotype control, and commercial HBcAg (1:2,000). After coating and blocking with 3% skim milk, the HBcAg peptide was added 36 ng or four ng per well. Then, HRP-linked 9.3, 9.4, or 9.5 monoclonal antibodies (1:300) were added, followed by the addition of ABTS substrate for color development.

III. RESULTS AND DISCUSSION

HBV infection can be detected either by DNA isolation and amplification by PCR or immunoassays [28]. PCR is more sensitive, but it is more difficult and expensive [28]–[30]. On the other hand, immunoassay such as ELISA is easier and faster to detect a higher number of samples [31], [32]. Most ELISA developed for HBV infection utilized HBsAg-based detection. However, mutation of HBsAg may lead to a false negative analysis [15], [33]. Thus, anti-HBcAg immunoassay is also combined with HBsAg detection for more comprehensive analyses [34]. More recently, a sensitive immunoassay based on HBcrAgs was developed utilizing monoclonal antibodies against HBcAg and HBeAg [35], [36]. The assay is more sensitive than HBsAg-based immunoassay, while the results are associated with HBV DNA levels in serological samples [35]. Moreover, HBcAg is one of HBcrAgs, which also developed as candidate HBV vaccines, especially in combination with HBsAg [22]. Therefore, we generated our monoclonal antibodies against HBcAg to develop homemade sandwich ELISA for research purposes.

Fig. 1 Mouse immunization and isolation of lymph nodes. A. Schematic representation of hybridoma generation. B. Enlargement of medial iliac (left panel) and sub-iliac (right panel) lymph nodes of immunized mouse after single shot of HBcAg peptide emulsified in incomplete Freund’s adjuvant (ICFA). Open circle represents the location of lymph node.
Fig. 2 Screening of candidate hybridomas by using ELISA plate coated with HBcAg retrieved two candidate polyclonal hybridomas 5#21 and 9#4.

Fig. 3 ELISA screening of 5#21 and 9#4-derived monoclonal hybridomas. The black bars indicated candidate monoclonal hybridomas, which were retrieved for further experiment.

A. Hybridoma Generation and Screening

Hybridoma techniques are commonly used to generate monoclonal antibodies (mAbs), the essential tools for research and diagnostic. There are a variety of procedures that can be applied to immunize the animals and isolate spleen or lymph nodes to prepare hybridoma. In this study, we modified the procedure established by Sado and colleagues [27], [37]. We utilized a single injection of HBcAg peptide emulsified in ICFA applied intramuscularly at the base tail of 11-week-old BALB/c mouse to enlarge iliac lymph nodes (Fig. 1.A.). Unlike complete Freund’s adjuvant (CFA), which is generally used for first immunization, ICFA, which is usually used as booster adjuvant, does not contain M. tuberculosis cell extract to induce Toll-Like Receptor (TRL) response. However, CFA induces additional severe lesions at the injection sites. On the other hand, ICFA still maintains its function as a water-in-oil emulsion adjuvant to release the antigen slowly from the injection spot and prolong antigen exposure to the host immune system [38] while mediating the activation of dendritic cells, which sequentially generate follicular T helper cells and promote antibody production by follicular B cells [39]. Our protocol found enlargement of medial iliac lymph nodes and sub-iliac lymph nodes (Fig. 1.B.). One month after a single immunization, we isolated both enlarged medial iliac and sub-iliac lymph nodes located close to the injection sites to retrieve antibody-secreting B cells. The intramuscular injection has been widely applied for vaccination to generate
antibody responses of local lymph nodes [40]. After about two weeks, we selected 96 growing clones under a HAT-containing medium to screen hybridoma conditioned medium by ELISA using an HBCAg coated plate. As a result, we retrieved two candidates polyclonal hybridoma clones, clone 5#21, which derived from medial iliac B cell, and clone 9#4, which derived from sub-iliac B cell. Furthermore, conditioned medium from clone 5#21 demonstrated a higher signal for HBCAg detection than the conditioned medium from clone 9#4 (Fig. 2). Because these two hybridomas were observed as polyclones, we performed monoclonalization of hybridomas by limiting dilution to seed one cell per well in a 96-well cell culture plate, 48 wells for each clone. After the new clones reached confluence, next round ELISA was carried out to screen the positive monoclonal clones. As the results, we obtained several candidate clones 5B5, 5C3, 5D3, 5F4 and 5H5 derived from clone 5#21 and named as 5.1, 5.2, 5.3, 5.4 and also candidate clones 9A7, 9A8, 9D8, 9D11, 9G7, 9G10 and 9H8 derived from clone 9#4 and renamed as 9.1, 9.2, 9.3, 9.4, 9.5, 9.6, 9.7 (Fig. 3).

Fig. 4 Screening of candidate anti-HBcAg monoclonal antibodies in conditioned medium for sandwich ELISA.

To establish homemade sandwich ELISA, we selected the candidate monoclonal antibodies by applying them as both capture and detection antibodies. The sandwich consists of candidate capturing antibodies, recombinant HBCAg peptides, candidate detection antibodies, and HRP-conjugated anti-mouse IgG γ-chain secondary antibody. The antibodies in wells that produce higher signals than control well without detection antibodies were selected as candidate antibodies. As the results, the wells coated with clone 5.2 conditioned medium representing capture antibody and used clone 9.1, 9.3, 9.4, 9.5 or 9.6 conditioned media representing detection antibodies showed higher signals of HBCAg detection than the control without detection antibodies (Fig. 4). Next, we retrieved clones 9.3, 9.4, and 9.5 conditioned media for antibody purification and further sandwich ELISA after conjugation with HRP.

B. Antibody Purification

We purified the antibodies by using protein A/G agarose to purify general IgG based antibodies. All candidate mAb from clones 9.3, 9.4, and 9.5 conditioned media were purified using this method. As we can see in figure 5, the mAbs were successfully purified from serum-free conditioned medium and mostly concentrated in eluate fraction 2. Determination of mAbs concentration by nanodrop results in mAbs concentration 214, 100, and 164 ng/µl for 9.3, 9.4, and 9.5 mAbs, respectively, which represent in the SDS-PAGE gel after coomassie blue staining. The purified antibodies were concentrated in elution fraction 2, which appeared as denatured heavy and light chains (Fig. 5). Purified antibodies were then conjugated with HRP for the development of homemade sandwich ELISA. Unfortunately, the conditioned medium of clone 5.2 was not purified by protein A/G agarose, which indicates that mAb secreted by this clone may not be IgG. We retrieved IgG-based antibodies derived from clone 9#4 for sandwich ELISA for a further experiment.

![Fig. 4 Screening of candidate anti-HBcAg monoclonal antibodies in conditioned medium for sandwich ELISA.](image1)

![Fig. 5 Profile of anti-HBcAg monoclonal antibodies 9.3, 9.4 and 9.5 in reducing sample buffer after purification by protein A/G sepharose. The antibodies appeared as denatured heavy chains and light chains. E: eluate, W: wash, FT: flow through.](image2)
C. Establishment of Home-Made Sandwich ELISA

All antibodies 9.3, 9.4 and 9.5 can be used in ELISA during the screening of mAbs for recombinant HBcAg detection. Furthermore, for sandwich ELISA application, we conjugated those antibodies with HRP to establish easy and fast ELISA consistent with several steps: coating of ELISA plate with purified mAbs, blocking with skim milk/PBS-T, the addition of antigen, the addition of HRP-linked mAbs, the addition of ABTS substrate with additional PBS washing before each step. As a result, HRP-linked monoclonal antibody 9.3 showed the highest signal as detection antibody with 9.4 and 9.5 mAbs as capture antibodies (Fig. 6). In addition, the commercial HBcAg antibody showed a positive signal as our candidates mAbs, while on the contrary, control IgG as capture antibody did not generate a significant signal.

Sandwich ELISA is an easy immunoassay-based method for high-throughput applications. By obtaining monoclonal antibodies, which function as either capture or detection antibodies, the antigen-containing samples can be sandwiched between those antibodies. Detection antibodies can be conjugated to HRP enzyme for further direct color development after chromogenic substrate addition. Here, we successfully retrieved hybridoma clones that produced anti-HBcAg mAbs 9.3, 9.4, and 9.5 for sandwich ELISA application. However, further studies are needed to examine the serological samples in comparison to available commercial kits.

Immunostaining of the liver tissue section is another general method to measure chronic hepatitis B patients [41]. The presence of HBcAg in nuclear hepatocytes indicates active viral replication. Moreover, HBcAg also is detected in the cytoplasm representing the current regenerative activity of HBcAg protein expression and the viral assembly [42]. To expand the research applications of our anti-HBcAg monoclonal antibodies, we will further investigate the capability of those antibodies for immunostaining of tissue section or in vitro cell culture expressing ectopic HBcAg.

IV. CONCLUSION

We successfully isolated monoclonal hybridomas capable of producing anti-HBcAg monoclonal antibodies for sandwich ELISA application. Those monoclonal antibodies produced by clones 9.3, 9.4, and 9.5 are IgG-based antibodies that can be purified using Protein A/G agarose and conjugated with HRP for easy detection with a chromogenic substrate. HRP-linked monoclonal antibody 9.3 had the highest signal of HBcAg detection for detection antibody with 9.4 and 9.5 monoclonal antibodies as capture antibodies.

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CONFLICT OF INTEREST

The authors declare that there was no conflict of interest related to this research.

AUTHOR CONTRIBUTIONS

Conceptualization and methodologies by EPS. Research investigation by EPS, NH, AK, PWP, AP. Funding acquisition by AZM, EPS, and AS. Project administration and ethical clearance document by NH. Writing original draft by
read and approved the final manuscript.

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