Monoclonal antibody preparation of Golgi phosphoprotein 2 and preliminary application in the early diagnosis of hepatocellular carcinoma

QIANG JU*, YANJIE ZHAO*, YANHONG LIU, GUOHUA ZHOU, FENG LI, PINGLI XIE, YUEHUI LI and GUAN-CHENG LI

Cancer Research Institute, Key Laboratory of Carcinogenesis and Cancer Invasion Ministry of Education, Key Laboratory of Carcinogenesis Ministry of Health, Central South University, Changsha, Hunan 410078, P.R. China

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Abstract. Golgi phosphoprotein 2 (Golph2) is a type II Golgi-specific membrane protein, which has been found to be overexpressed in hepatocellular carcinoma (HCC) patients. The sensitivity of diagnosis of HCC using Golph2 (76%) was markedly elevated compared with alpha-fetoprotein (AFP) (70%), and Golph2 is expected to be a novel and effective serum biomarker for the diagnosis of HCC. The aim of this study was to prepare monoclonal antibodies against Golph2 and to establish double-antibody sandwich enzyme-linked immuno-sorbent assay (s-ELISA), which will be used in diagnostics, therapeutics and as a tool in understanding the role of Golph2 in the pathogenesis of liver diseases and cancer. In this study, fusion protein TRX-Golph2 was expressed and purified using an Escherichia coli system. BALB/c mice were immunized with TRX-Golph2 recombinant protein. The hybridoma technique was used for the production of anti-Golph2 monoclonal antibody. Hybridoma clones were screened using indirect ELISA and anti-Golph2 monoclonal antibody was produced in the ascites of BALB/c mice. The specificity of anti-Golph2 monoclonal antibody was detected by western blot analysis and immunocytochemistry. s-ELISA was established using horseradish peroxidase (HRP)-labeled anti-Golph2 monoclonal antibody and used to detect the antigen in the serum of HCC patients. As a result, five stable hybridoma cell clones (5C6D5, 5B7F5, 7F5F3, 8A7B4, 8C9E8) producing anti-Golph2 monoclonal antibody were established. The highest titer of anti-Golph2 monoclonal antibody (5C6D5) was 1:51,200. Western blot analysis revealed that anti-Golph2 monoclonal antibody had a high specificity for Golph2 protein. Anti-Golph2 monoclonal antibody was HRP-labeled and the optimal working concentration was found to be 1:500. The levels of antigen in a proportion of HCC patients were shown to be significantly higher compared to those found in healthy controls.

Introduction

Hepatocellular carcinoma (HCC) is one of the most common malignant tumors with high malignancy and mortality due to lack of early diagnosis and its resistance to conventional chemotherapy (1). HCC affects approximately one million people every year worldwide, with the incidence equal to the mortality. In 2008, HCC was listed as the third most common cause of cancer-related mortality (2). Thus, early diagnosis is crucial in order to increase the survival rate for patients (3). To date, alpha-fetoprotein (AFP) together with imaging and pathology detection are commonly used in early clinical diagnosis of liver cancer. However, the specificity and sensitivity of AFP for liver cancer screening are not satisfactory. With the development of molecular biology, a number of new types of tumor markers have been discovered.

Golgi phosphoprotein 2 (Golph2) is a type II Golgi-specific membrane protein that is predominantly expressed in the epithelial cells of a number of human tissues (4). In normal human liver, Golph2 is only expressed in biliary epithelial cells and is almost undetected in liver cells. However, increased expression of Golph2 has been reported to be correlated with numerous viral or non-viral infectious liver diseases (5). It was first identified in a search for upregulated hepatic genes in acute giant-cell hepatitis (4q) and then in patients with acute and chronic hepatitis (6). Studies revealed that Golph2 was overexpressed in the serum of HCC patients (7,8). In China, Mao et al first observed that the level of Golph2 in the serum of patients with HCC infected by hepatitis B virus (HBV) was significantly higher than HBV carriers, patients without hepatic diseases and healthy adults (9). In addition, other studies reported that the sensitivity of diagnosis of HCC by Golph2 (76%) was higher than AFP (70%), indicating that
Expression, purification and identification of recombinant protein. The recombinant plasmid pET21a(+)–TRX–Golph2, which was constructed in the laboratory, was transformed into Escherichia coli Rosetta (ATCC, Manassas, VA, USA) and optimal expression of recombinant proteins was achieved through controlling the concentration of isopropyl-β-D-thiogalactopyranoside (IPTG) and growth conditions. After induction, bacteria were harvested and centrifuged at 4°C, 4,449 x g for 10 min, and the pellet was resuspended in 50 mmol/l sodium phosphate with 0.5 mol/l NaCl (pH 8.0). The resuspended cells were then lysed by sonication and centrifuged at 4°C, 10,012 x g for 10 min. The expression form of fusion protein TRX-Golph2 was analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE, 12% gel). The recombinant protein TRX-Golph2 was further purified by immobilized metal affinity chromatography (MagneHis™ Protein Purification System; Promega, Madison, WI, USA) under native conditions. The purified protein was dialyzed in 1X phosphate buffered saline (PBS) at 4°C overnight and condensed to a high concentration.

The concentrated recombinant protein was separated by SDS-PAGE on a 12% polyacrylamide gel and then transferred onto a nitrocellulose membrane (Millipore, Billerica, MA, USA) by electroblotting. Fat-free milk (5%) was used to block the membrane at 37°C for 2 h. The membrane was then incubated with anti-His tag antibody (1:2,000; Santa Cruz Biotechnology, Inc., Santa Cruz, CA, USA) overnight in a 4°C refrigerator. The membrane was then washed with PBS-Tween-20 (PBST) three times and incubated with the horseradish peroxidase (HRP)-conjugated goat anti-mouse secondary antibody (1:1,000, Sigma, St. Louis, MO, USA) in a 37°C shaker for 1 h. After washing three times with PBST, the membrane was visualized with an enhanced chemiluminescence (ECL) system (Pierce Biotechnology, Inc., Rockford, IL, USA).

Immunization of mice. Eight-week-old Balb/c mice were immunized three times with purified TRX-Golph2 fusion protein at days 0, 21 and 42 with 100, 50 and 50 µg, respectively, by peritoneal injection. Complete and incomplete Freund's adjuvant (1:1, Sigma) were used to emulsify the protein in the first and other immunizations, respectively. Ten days after the third immunization, mice were bled from the caudal vein and the serum titer was detected by indirect ELISA. Three days after the last booster immunization (100 µg), the spleen of the mouse with the highest titer was removed under sterile conditions to prepare a splenic lymphocyte suspension for cell fusion.

Cell culture and fusion. Mouse myeloma cell line SP2/0 was cultured in 10% fetal bovine serum (Hyclone, Tauranga, New Zealand) and RPMI-1640 medium (Gibco-BRL, Grand Island, NY, USA) in a 37°C, 5% CO₂ incubator. The feeder cells, obtained from the peritoneal cavity of a normal BALB/c mouse by peritoneal injection of RPMI-1640 medium, were plated in 96-well plates and cultured in a 37°C, 5% CO₂ incubator for 1 day prior to fusion. Splenic lymphocytes (1x10⁶) were fused with SP2/0 cells (1x10⁵) in 50% polyethylene glycol 1500 (PEG 1500, Sigma-Aldrich, St. Louis, MO, USA). The hybridoma cells were cultured in 96-well plates containing feeder cells and screened by hypoxanthine-aminopterin-thymidine (HAT; Sigma-Aldrich) and cultured in a 37°C, 5% CO₂ incubator. Two weeks later, indirect ELISA using purified TRX-Golph2 protein and HepG2 cells as coating proteins, respectively, was performed to detect the positive clones and then screened clones were recloned by limiting dilution in hypoxanthine-thymidine (HT; Sigma-Aldrich) medium for a further 2 weeks.

Screening of positive clones and cloning. By observing the 96-well plates, we selected a single clone to perform the indirect ELISA using purified TRX-Golph2 protein and HepG2 cells as coating proteins, respectively. The optical density (OD) was read at a wavelength of 405 nm (microplate reader, Tianshi, Beijing, China) and considered positive when the ratio of OD test to OD control was greater than 3 times. The wells that were positive for both purified TRX-Golph2 protein and HepG2 cells were considered as positive wells and then recloned 3-5 times by limiting dilution until their antibody secretion was 100%. Finally, the positive clones in the 96-well plates were transferred to 24-well plates in 10% fetal bovine serum RPMI-1640 medium. The strain of hybridoma cells that secreted specific mAb was established.

Preparation and titer determination of anti-Golph2 mAb. Balb/c mice were injected with incomplete Freund’s adjuvant (0.5 ml) by peritoneal injection. Three days later, the mice were peritoneally injected with 5x10⁶ hybridoma cells diluted by D-Hank’s. Ten days later, ascitic fluid was collected and centrifuged at 278 x g for 5 min to remove the cellular deposition. Anti-Golph2 mAb in supernatant was diluted...
into different gradients (1:100; 1:200; 1:400; 1:800; 1:1,600; 1:3,200; 1:6,400; 1:12,800; 1:25,600; 1:51,200) and added into 96-well plates coated with TRX-Golph2 antigen, respectively. Indirect ELISA was performed to detect the titer of anti-Golph2 mAb. The OD was read at 405 nm.

Subclass determination and purification of anti-Golph2 mAb. The subclass of anti-Golph2 mAb was determined using the SBA Clonotyping System (SouthernBiotech, Birmingham, AL, USA) according to the manufacturer’s instructions. The anti-Golph2 mAb (3 ml) was added with equivalent precooled ammonium sulfate overnight in a 4°C refrigerator. They were then centrifuged at 4°C, 1,738 x g for 15 min. The supernatant was decanted, solubilized by adding PBS (0.02 M, 3 ml), and then filtered by Sephacryl S-300 HR gel filtration (Pharmacia, USA). PBS containing 0.02% NaN₃ was used to wash the column and collected components. The positive components confirmed by indirect ELISA were dialyzed in PBS at 4°C overnight and condensed to a high concentration. The purity was identified by SDS-PAGE (12%).

Determination of the relative mAb affinity. A 96-well immunoplate was coated with the TRX-Golph2 fusion protein (4 μg/ml) at 4°C overnight, and then was blocked with 2% fetal bovine serum at 37°C for 2 h. Serial dilutions of the purified mAb were incubated at 37°C for 2 h. The plate was rinsed and incubated with the HRP-conjugated goat anti-mouse antibody (1:4,000 dilution) at 37°C for 1 h. After washing, TMB (3,3',5,5'-tetramethylbenzidine) substrate was used for color development. The OD value was measured at 405 nm in order to determine the relative affinity.

Western blot analysis of anti-Golph2 mAb specificity. Total proteins extracted from HepG2 cells were separated by SDS-PAGE on a 12% polyacrylamide gel and then transferred onto a nitrocellulose membrane (Millipore) by electroblotting. Fat-free milk (5%) was used to block the membrane at 37°C for 2 h. The membrane was then incubated with purified anti-Golph2 mAb (1:2,000) overnight in a 4°C refrigerator. The membrane was then washed with PBS-Tween-20 (PBST) three times and incubated with the HRP-conjugated goat anti-mouse secondary antibody (1:1,000) in a 37°C shaker for 1 h. After washing three times with PBST, the membrane was visualized with an ECL system (Pierce Biotechnology, Inc.).

Immunocytochemistry. Appropriate amounts of cells were cultured on microslides overnight. When the cells were adherent, slides were fixed with 4% paraformaldehyde for 15 min. PBST (0.5%) was used to wash the slides three times. Endogenous peroxidase activity was blocked by 3% H₂O₂/methanol and non-specific binding was blocked with 2% BSA-PBS. Diluted anti-Golph2 mAb (50 μl; 1:50) was added and incubated overnight in a 4°C refrigerator. After three 0.5% PBST washes, the appropriate (1:200) HRP-conjugated goat anti-mouse secondary antibody was added and incubated at room temperature for 1 h. Slides were washed with 0.5% PBST three times. Diaminobenzidine (DAB) was added and allowed to react for 1 min. The reaction was stopped by adding water and hematoxylin was added to counterstain for 2 min. Slides were dehydrated with alcohol and mounted with neutral gummi.

Preparation of enzyme-labeled anti-Golph2 mAb. Modified sodium periodate oxidation was used for labeling anti-Golph2 mAb (8A7B4) with high purity. The result and the optimal working concentration of labeled anti-Golph2 mAb with HRP were analyzed by direct ELISA.

Establishment and optimization of double-antibody s-ELISA. A 96-well immunoplate was coated with purified anti-Golph2 mAb (5C6D5, 20 μg/ml) at 4°C overnight and then blocked with 2% BSA-PBS at 4°C overnight. After washing plates with PBST three times, serum dilutions (1:2, 1:4, 1:8), strong positive (50 ng/ml antigen), weak positive (3 ng/ml antigen) and negative, were incubated at 37°C for 2 h, respectively. The plates were washed three times and enzyme-labeled anti-Golph2 mAb (1:300, 1:400, 1:500, 1:600) was added and the plates were incubated at 37°C for 1 h. The plates were washed again and colored with 100 μl 2,2-Azinoibis-(3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt (ABTS; Boehringer Mannheim GmbH, Germany) for 25 min at 37°C. The OD was read at a wavelength of 405 nm. A chessboard titration method was designed to examine the optimal working concentration of HRP-labeled mAb and the dilution ratio of serum.

Analysis of serum samples by s-ELISA. The expression level of Golph2 in serum dilutions was analyzed by s-ELISA.

Statistical analysis. All of the values which presented as the mean of three paralleled individual experiments were entered into SPSS 13.0 software. The mean and standard deviation (SD) of the OD of individual groups were calculated. The expression of Golph2 in serum was described using box-and-whisker plots. The difference between groups was analyzed using Student t-tests. P<0.05 was considered to indicate a statistically significant result.

Results

Expression, purification and identification of recombinant protein. The recombinant fusion protein was expressed under the optimizing prokaryotic expression conditions of 1 mM IPTG, 30°C, 2 x g, for 10 h and purified by immobilized metal affinity chromatography under native conditions. The expressed and purified protein were identified by 12% SDS-PAGE and subsequently stained with Coomassie Brilliant Blue R250 (Fig. 1). The molecular weight of TRX-Golph2 protein was ~73 kDa. The protein was then further identified by western blot analysis using anti-His tag antibody (Fig. 2).

Establishment of hybridoma cell lines and preparation of anti-Golph2 mAb. Through the procedures of immunization, fusion and clone selection, five hybridoma cell lines (5C6D5, 5B7F5, 7F5F3, 8A7B4, 8C9E8) that stably secreted anti-Golph2 mAb were obtained. The identity, subclass and titer are shown in Table I.

Investigation of mAb relative affinity. The relative affinities of mAbs were defined by the antibody concentration at which the OD value reached half the maximal signal at the plateau stage of antigen-antibody binding. The results revealed that
the order of relative affinity of the five selected mAbs was 8A7B4>5C6D5>5B7F5>8C9E8>7F5F3 (Fig. 3).

**Purification of anti-Golph2 mAb.** The anti-Golph2 mAb was purified by Sephacryl S-300 HR gel filtration. After dialyzing and condensing, the purity was identified by SDS-PAGE (12%) (Fig. 4).

**The specificity of anti-Golph2 mAb.** Immunocytochemistry (HepG2) revealed that the membrane of the Golgi was positively stained (Fig. 5A and B). By western blot analysis, anti-Golph2 mAb (8A7B4) was shown to react specifically with Golph2 protein (45kDa) in accordance with the theoretical value (Fig. 5C).

**Preparation of enzyme-labeled anti-Golph2 mAb.** The anti-Golph2 mAb (8A7B4) was labeled with HRP by modified sodium periodate oxidation. By direct ELISA, the working concentration of labeled mAb was shown to be 1:500.

**Establishment and optimization of double-antibody s-ELISA.** One serum sample from a known patient with overexpression of Golph2 in serum was used for the determination of the dilution ratio of serum. The optimal working concentration of HRP-labeled mAb and the dilution ratio of serum were fixed on 1:500 and 1:2, respectively, by the chessboard titration method (Table II).

**Detection of serum.** The levels of antigen in the samples were detected by s-ELISA using HRP-labeled anti-Golph2 mAb (Fig. 6). The median serum level of Golph2 in the HCC group

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**Table I. Characterization of anti-Golph2 protein mAbs.**

<table>
<thead>
<tr>
<th>Clone name</th>
<th>Titer</th>
<th>Subclass</th>
</tr>
</thead>
<tbody>
<tr>
<td>5B7F5</td>
<td>1:25600</td>
<td>IgM (λ)</td>
</tr>
<tr>
<td>5C6D5</td>
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</tr>
<tr>
<td>7F5F3</td>
<td>1:25600</td>
<td>IgM (κ)</td>
</tr>
<tr>
<td>8A7B4</td>
<td>1:12800</td>
<td>IgG1 (κ)</td>
</tr>
<tr>
<td>8C9E8</td>
<td>1:25600</td>
<td>IgM (λ)</td>
</tr>
</tbody>
</table>

Golph2, Golgi phosphoprotein 2; mAbs, monoclonal antibodies.
As shown in Fig. 6, the average serum content of Golph2 in HCC patients is >100 µg/l, consistent with other studies using s‑ELISA (8).

Discussion

Since it was first identified in acute giant-cell hepatitis, tissue-based Golph2 overexpression has been confirmed in HCC (8,14), lung adenocarcinoma (15), seminomas (16), renal cell cancer (17) and other malignant diseases. Notably, a large number of studies have indicated that the serum level of Golph2 may be a more reliable biomarker for the early diagnosis of HCC than AFP. Previous studies have showed that Golph2 is an adenovirus‑induced cellular protein whose expression is regulated through E1A protein (4,18). Golph2 overexpression, intracellular trafficking between the Golgi and plasma membrane through an endosomal pathway, and the cleavage of Golph2 may explain the secretion of Golph2 (19). However, the exact mechanism involved remains unclear. mAbs against Golph2 prepared in this study lay a foundation for further investigation of the interaction between antigen epitopes and antibodies, and provide a tool for determining the Golph2 secretion mechanism.

In our study, the E. coli system Rosetta was selected to express Golph2 protein. When using a prokaryotic system, it is difficult to express the full length of Golph2. The Rosetta system contains rare codon tRNA that improves protein expression, allowing Golph2 to be expressed. We selected pET21a(+)-TRX as the prokaryotic expression vector. TRX is a thioredoxin and is widely used in molecular cloning expression. TRX protein prevents the target protein from being degraded by endoproteinase and provides the ligand binding sites for affinity purification. Furthermore, TRX contributes to folding of the target protein and increases its soluble expression. The fusion protein TRX-Golph2 can be expressed under native conditions.
conditions. SDS-PAGE showed that the fusion protein had a 78 kDa band. This is not in accordance with the theoretical value, as Golph2 has abundant acidic amino acids (PI=4.72), which causes the migration rate of the recombinant protein to decline in the denatured SDS-PAGE (20). After induction and purification, we successfully obtained TRX-Golph2 in the soluble form.

Using mAb technology with high purity and good reproducibility (21,22), we selected five hybridoma cell lines that stably secreted anti-Golph2 mAb. Two strains of relatively high affinity mAb were selected and analyzed by western blot analysis and immunocytochemistry. Western blotting showed that the mAb recognized Golph2 protein specifically. Immunocytochemistry revealed that Golph2 is located in the membrane of the Golgi. The mAb with higher purity was labeled with HRP. The concentration of the coated antibody was fixed at 20 μg/ml, and the optimal working concentration of HRP-labeled mAb and the dilution ratio of serum was confirmed to be 1:500 and 1:2, respectively, using the chessboard titration method in s-ELISA.

Double-antibody s-ELISA is commonly used for testing antigens. In the present study, HRP-labeled mAb with different epitopes from coated mAb was produced for the first time. Previous studies on Golph2 in serum used enzyme-labeled anti-Golph2 polyclonal antibody or a combination of anti-mAb with enzyme-labeled anti-IgG to perform the double-antibody s-ELISA (8). Polyclonal antibody has a high level of cross reaction and can generate false positive results, while mAb has high specificity and accuracy. Therefore, our s-ELISA had significantly higher sensitivity and specificity than the previous studies. Results showed that the expression level of serum Golph2 in patients with HCC was markedly higher than that in healthy individuals.

In conclusion, mAbs and s-ELISA against Golph2 may be useful tools for the sensitive and specific diagnosis of HCC. The preparation of humanized antibody has been published previously (23), whereas the optimization of humanized antibody requires further research. This study provides the basis for optimization of humanized antibody.

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References