Functional screen for secreted proteins by monoclonal antibody library and identification of Mac-2 binding protein as a potential therapeutic target and biomarker for lung cancer

Lichao Sun, 1 Lizhao Chen, 1 Lixin Sun, 1 Jian Pan3, Long Yu, 1 LuLu Han, 1 Zhihua Yang2, Yuanming Luo4*, and Yuliang Ran1*

1State Key Laboratory of Molecular Oncology, Cancer Institute (Hospital), Chinese Academy of Medical Sciences, Peking Union Medical College, Beijing, China.
2Department of Cell and Molecular Biology, Cancer Institute (Hospital), Chinese Academy of Medical Sciences, Peking Union Medical College, Beijing, China.
3Department of Hematology and Oncology, Children's Hospital of Soochow University, Suzhou, China.
4State Key Laboratory of Microbial Resources, Institute of Microbiology, Chinese Academy of Sciences, Beijing, China.

*Correspondence:

Yuliang Ran, State Key Laboratory of Molecular Oncology, Cancer Institute (Hospital), Chinese Academy of Medical Sciences, Peking Union Medical College, No.17 Panjiayuan Nanli, Chaoyang District, Beijing 100021, People's Republic of China. Phone: 86-10-87788749; Fax: 86-10-67783169; E-mail: ran_yuliang@yahoo.com.cn

Yuanming Luo, State Key Laboratory of Microbial Resources, Institute of Microbiology, Chinese Academy of Sciences, No.1 West Beichen Road, Chaoyang District, Beijing 100101, People's Republic of China. Phone: 0086-10-64807432; Fax:
Abbreviations:

SCLC, small cell lung cancer
NSCLC, non-small-cell lung cancer
SCC, squamous cell carcinoma
AD, adenocarcinoma
LCLC, large cell lung carcinoma
Mac-2BP, Mac-2 binding protein
NSE, neuron-specific enolase
CEA, carcinoembryonic antigen
CYFRA21-1, cytokeratin 19 fragment
TPA, tissue polypeptide antigen
ProGRP, progastrin releasing peptide
ROC, receiver operation characteristics
AUC, area under the curve
Summary: Identification of secreted proteins of lung cancer could provide new candidates of serum biomarkers for cancer diagnosis or targets for therapeutic intervention. In this study, we developed a novel strategy that combined functional monoclonal antibody library screening technique and mass spectrometry to identify functional secreted proteins. BALB/c mice were immunized with cancer cells isolated from fresh human lung cancer tissues. The monoclonal antibody library containing 1160 mAbs was established with the mouse spleen cells, whose serum had most anti-proliferative effect on lung cancer cells. Monoclonal antibodies were subjected to an immunoreactive and functional screen and monoclonal antibodies that reacted strongly with secreted proteins in condition medium and lung cancer tissues with high inhibition of cell proliferation were selected. Antigens that recognized by antibodies were obtained by immunoprecipitation and then identified by mass spectrometry. Mac-2-binding protein (Mac-2BP), the antigen of 13H3 antibody, was identified using this approach. Functional studies demonstrated that the 13H3 antibody suppressed lung cancer cell lines ANIP-973 and A549 proliferation in vitro and inhibit ANIP973 xenograft tumors growth in vivo by inducing cell-cycle arrest at G1 phase, with up-regulation of p27 and downregulation of cyclin D1. Moreover, the serum level of Mac-2BP was significantly higher in lung cancer patients than healthy controls. At a cutoff value of 6ug/ml, Mac-2BP might be a diagnostic biomarker of lung cancer, especially for SCLC. Mac-2BP concentrations of 6ug/ml or higher was associated with poor overall survival in univariate analysis, and was an independent predictor in the multivariate COX analysis. Together, these results firstly demonstrated that
Mac-2BP can be used as a therapeutic target and potential biomarker for lung cancer. Our strategy is feasible, which may facilitate the identification of novel secreted biomarkers of lung cancer.

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**Running title:** Mac-2BP is a potential therapeutic target, biomarker for lung cancer

**Key words:** Lung cancer. Antibody library. Mass spectrometry. Mac-2 binding protein.
INTRODUCTION

Lung cancer is the leading cause of cancer-related death worldwide[1]. Despite diagnostic and therapy improvements over the past decade, the 5-and 10-year patient survival rates remain very low at 14 and 8%, respectively[2]. However, most people diagnosed with cancer confined to the primary site could survive more than 5 years[3]. Current serum protein biomarkers for lung cancer diagnosis are mainly neuron-specific enolase (NSE), carcinoembryonic antigen (CEA), cytokeratin 19 fragment (CYFRA21-1), tissue polypeptide antigen (TPA), progastrin releasing peptide (ProGRP), and tumor M2 pyruvate kinase[4-6]. However, the roles of these tumor markers in the diagnosis of lung cancer are still controversial and remain to be determined due to their relatively low sensitivity. Thus, there is an urgent need to identify lung cancer biomarkers that might be useful for diagnostic purposes.

Many secreted proteins can enter the blood circulation, with potential clinical utility for therapeutic targets and diagnostic biomarkers. From a biomarker discovery perspective, serum is the ideal sample to investigate, but it is difficult to analyze because of large amounts of albumin and other proteins[7]. Recently, analysis of conditioned media has proven to be a very successful strategy for identifying candidate biomarkers. It allows researchers not only to identify candidate biomarkers for the detection of cancer, but also to obtain potential therapeutic targets[8, 9].

In the present study, we developed and used a novel antibody library-based proteomic technology to identify lung cancer-associated secreted functional biomarkers. A monoclonal antibody library was established by immunizing mice with lung cancer
cells isolated from carcinoma tissues. Monoclonal antibodies that reacted with secreted proteins from human lung cancer cells and specifically recognized lung cancer tissues were selected. And the corresponding antigens were identified by immunoprecipitation and mass spectrometry. Using this strategy we successfully identified Mac-2BP as a potential therapeutic target and biomarker for lung cancer.

**EXPERIMENTAL PROCEDURES**

**Samples-** All tissue and blood specimens were collected from patients in the Department of Pathology in Cancer Hospital, Chinese Academy of Medical Sciences, Beijing, China. Inclusion criteria for the lung cancer patients were: No receiving any treatment prior to surgery; no metastatic disease at diagnosis; no previous diagnosis of carcinoma; no evidence of disease within 1 month after primary surgery. All tissue samples were taken by experienced surgeons and examined independently by two experienced pathologists. Fresh tumor tissues and paired tumor-adjacent normal lung tissues at least 5 cm away from tumor were obtained immediately after surgery, and necrotic tissue was excluded and repeatedly washed with saline in aseptic condition. And normal lung tissues were confirmed to contain no tumor cells by histopathologic evaluation. All the blood samples were collected 24 hours or less before surgery by peripheral venous puncture and were centrifuged at 1500g at 4°C for 10 minutes. The separated plasma was aliquoted and stored at -80°C for future analysis. For immunization, 20 fresh primary lung cancer tissues including 8 SCCs, 9 ADs, 1 LCLC and 2 SCLCs were obtained during 2001-2002(Table1). For immunohistochemical analysis, 105 paraffin-embedded lung tumors and paired
adjacent normal lung tissues were randomly obtained from patients during 1997-2002. Further characteristics of patients and tumors are listed in Table S1. For ELISA study, blood samples were obtained from 320 lung cancer patients (median age at 60 with a range of 50 to 70 years) during 2005-2008 including 115 SCCs, 119 ADs, 10 LC and 76 SCLC. 80 specimens of healthy individuals with similar age (median age at 58 with a range of 48 to 68 years) and sex distribution were analyzed, which were donated on a voluntary basis. All patients were routinely examined every 6 months during 48 months of follow-up. During that period, 480 patients were retrieved. 69 cases were excluded because of lack in clinical and pathological information. 91 cases were excluded because of insufficient follow-up information after treatment. The remaining 320 lung patients were eligible for this study. For all the specimens, clinical data were obtained retrospectively for all lung cancer patients, including age, gender, differentiation, and TNM stage. The study was approved by the medical ethics committee of Cancer Institute and Hospital, CAMS.

**Cell Culture**-Human lung cancer cell lines (A549, ANIP-973) and mouse myeloma cell line (SP2/0) were maintained in the complete growth medium DMEM (Gibco) containing 10% FBS, 2mmol/L glutamine, 100 U/mL penicillin and 100 U/mL streptomycin. All the cell lines were grown at 37 °C in an atmosphere of 5% CO₂. The cells were detached with 0.2% trypsin and 0.1% EDTA and the medium was changed once every other day.

**Conditioned Medium Collection**-For the primary lung cell cultures, we harvested lung cancer tissues from ten lung cancer patients into washing buffer (DMEM
containing 1% antibiotic) immediately after surgery and then transferred them to the laboratory within 30 min. We isolated and cultured the primary cancer cells according to the previous procedures[10, 11]. Briefly we washed and minced the tissues in washing buffer immediately after harvesting and digested them in DMEM medium containing DNase I, collagenase type I and hyaluronidase at 37°C for 15 min and cell suspension was passed through a 40 mM nylon mesh and cancer cells were cultured on each 100-mm dish. After removing the spent medium, we rinsed the cultures three times with Hank’s balanced salt solution and then incubated them in the serum-free DMEM at the 60-70% cell confluence. After 24 h of incubation, we collected and centrifuged the CM for 5 min at 4,000×g at 4 °C to remove cell debris and ultrafiltration was used to concentrate the proteins for 60 fold.

**Monoclonal antibody library construction and screen**- A library of monoclonal antibodies was generated from mice immunized with cancer tissue homogenates in lung cancer using established procedures[12]. The antibody subtypes were identified using the Clonotyping system (SouthernBiotech). Antigen immunoreaction was performed to scan for antibodies capable of reacting with secreted protein. Briefly, Conditioned medium (CM) was collected as mentioned above. The plates were coated with the 200µl CM and were incubated 4°C overnight. Block the remaining protein-binding sites in the coated wells by adding 200 µl blocking buffer, 5% BSA for overnight at 4°C. Wash the plate twice with PBS. Then 100ul of hybridoma supernatant were transferred to an ELISA plate and were incubated for 2 h at room temperature. Then wash the plate four times with PBS, and add the HRP conjugated
goat anti-mouse IgG, IgM(H+L). After sufficient color development add 100 µl of stop solution to the wells. Read the absorbance (optical density) of each well with a plate reader.

**SDS-PAGE Separation and Western blot analysis**—For western blot analysis, proteins from lung cancer xenograft or Anip973 and A549 cells were extracted by RIPA (50 mM Tris pH 7.4, 150 mM NaCl, 1% NP-40, 1% Triton-X100, 0.1% SDS, 0.5% sodium deoxycholate) buffer supplemented with a cocktail of protease inhibitors (Sigma). We dissolved the lyophilized proteins in sampling buffer and separated them with 10% SDS-PAGE. Proteins were transferred to PVDF membranes, and then probed with primary antibodies. The immunoreaction was visualized by super ECL detection reagent (Applygen, Beijing, China) following incubated with horseradish peroxidase-conjugated secondary antibodies. 13H3 antibody was screened from monoclonal antibody library. Antibodies against Galectin-3 (sc-32790, used at 1:800), cyclin D1 (sc-753, used at 1:2500), cyclin E (sc-481, used at 1:2500), CDK2 (sc-163, used at 1:2500), CDK4 (sc-260, used at 1:2500), and p21(sc-817, used at 1:1000) were purchased from Santa Cruz Biotechnology. p27(610241, used at 1:2500) antibody was purchased from BD Biosciences. Mac-2BP (AF2226, used at 1:2000) antibody was purchased from R&D Systems, Inc.

**Immunoprecipitation and mass spectrometry**—Immunoprecipitation was done essentially as previously described [12], using 13H3 antibody which was chemical coupled to CNBr activated Sepharose4B (Pharmacia). Then the condition medium was applied through the column, the antibody could bind specifically with the
targeting antigen. After washing away the unbound materials, the target protein was collected by elusion and was analyzed by SDS-PAGE and silver stain. Stained bands were excised and subjected to in-gel-digestion as previously described[12]. Dried tryptic peptide mixtures were dissolved in 3μl of saturated solution of α-cyano-4-hydroxycinnamic acid in 50% acetonitrile, and masses of peptides were determined by using MALDI-TOF mass spectrometer (Applied Biosystems Voyager DE-Pro Workstation, Foster City, USA.). The MALDI-TOF MS was operated in the positive ion delayed extraction reflector mode for highest resolution and mass accuracy. Peptides were ionized/desorbed with a 337-nm laser and spectra were acquired at 20 kV accelerating potential with optimized parameters. The MS spectrum was externally calibrated by a peptide standard mixtures (Applied Biosystems) containing des-Arg1-bradykinin (m/z 904.4675), Angiotensin I (m/z 1296.6851), Glu1-fibrinopeptide B (m/z 1570.6774), ACTH (1-17, m/z 2903.0867), ACTH (18-39, m/z 2465.1989), and ACTH (7-38, m/z 3657.9294). The resulting MS data was searched against the NCBIInr protein sequence database (3717264 sequences, 1278821221 residues) downloaded from ftp://ftp.ncbi.nih.gov/blast/db/FASTA/ on June 18, 2006 using the MASCOT 2.0 search engine (Matrix Science, London, UK). Search parameters were as follows: trypsin digestion with one missed cleavage, fixed modification for carbamidomethylation of cysteine, variable modification for methionine oxidation, and mass tolerance of 0.2 Da for singly charged ions. The taxonomy selection was Homo sapiens. The autolytic peaks of trysin 905.50, 2163.05, and 2273.15, and the common contaminant peaks of keratin such as 1307.6441 and
2705.14 (from gi/239938886), 1475.7379 and 1638.8278 (from gi/119395750), and 2501.23 (gi|239938650) were excluded from the mass list when searching the database. For all proteins identified by peptide mass fingerprint, Mascot scores greater than 64 were considered significant (p-value < 0.05).

**Measurement of Human Plasma Levels of Mac-2BP**-A commercially available ELISA kit (Bender MedSystems GmbH, Vienna, Austria) was used to measure Mac-2BP level. Each serum sample was run in duplicate. Briefly, 100 ml of serum (1:100 dilution) were placed into each well of the ELISA plate and incubated for 45 min at 37°C. The plates were washed four times with buffer and incubated with 100 ml of detection antibody at 37°C for 45 min. After four washes, the plates were incubated with substrate solution for 15 min at room temperature, then the reaction was stopped and the plates were read by a spectrophotometer at wavelength 450 nm. A standard curve was generated with the provided standards and used to calculate the quantity of Mac-2BP in each serum sample.

**Immunohistochemistry and tissue microarray assay**-Immunohistochemical analysis was performed with lung cancer tissues. Tissue microarrays were prepared from archival formalin-fixed, paraffin-embedded tissue blocks. For each tumor, a representative tumor area was carefully selected from a H&E-stain section. Sections 5μm in thickness were obtained and mounted on positively charged slides for immunohistochemical analysis. Standard avidin-biotin complex peroxidase immunohistochemical staining was performed. Briefly, after deparaffinization in xylene and graded alcohols, heated antigen retrieval was done in citrate buffer
(10mmol/L pH 6.0) by water-bath kettle heating for 30min. Endogenous peroxidase was blocked in 0.3% hydrogen peroxide for 10 min. Nonspecific binding was blocked by incubation in 10% normal animal serum for 10min. Sections were incubated at 4°C for 24 h with supernatant of each clone of monoclonal antibodies (10µg/ml). Expression levels of proteins were scored by malignant/epithelial cells staining intensity and the percentage of immunoreactive cells. Tissues with no staining were rated as 0, with faint staining or moderate to strong staining in 25% of cells as 1, with moderate staining or strong staining in 25% to 50% of cells as 2, and with strong staining in >50% of cells as 3. Lung cancer tissues that registered levels 0 and 1 were defined as negative for expression, whereas samples at levels 2 or 3 were defined as positive.

**In vitro siRNA transfection**—Two small interfering RNA (siRNA) based on the Mac-2BP sequences (5'-CCATCAGCGTGAATGTGCA-3' and 5'-GGACCTGTATGCCTATGCA-3') were synthesized by Ribobio Inc (Guangzhou China). Mac-2BP expression was determined by Western blot and reverse transcription polymerase chain reaction (RT-PCR).

**Cell proliferation assay**—Cells were trypsinized and resuspended in complete medium and seeded equally into 96-well plates. Cells were counted with CCK-8 (Dojindo,Laboratories, Japan), according to the manufacturer’s instructions and the optical density (OD) was measured at 450 nm. These experiments were performed twice with similar results. Proliferation was also estimated using the EdU incorporation assay. Briefly, cells were cultured in 6-well plates and exposed to 50
mM EdU (Ribobio, Guangzhou, China) for 4 h at 37°C. The cells were then fixed in 4% formaldehyde for 30 min at room temperature and permeabilized in 0.5% Triton X-100 for 10 min. Cells were washed with PBS, and each well was incubated with 400 ml 1XApolloH reaction cocktail for 30 min. DNA was then stained with 5 mg/ml Hoechst 33342 for 30 min and imaged under a fluorescent microscope. The EdU-labeled cells were counted.

**Flow cytometric assay for cell cycle** - Cancer cells were transfected with Mac-2BP siRNAs as described above. 48 h later, the cells were digested with trypsin, washed twice in PBS, and centrifuged at 1200 rpm for 5 min. The cells were resuspended in cold ethanol and maintained at 4°C for 24 h. After being washed twice in PBS, the cells were stained with propidium iodide (10μg/ml) for 30 min and subjected to flow cytometric analysis.

**Antibody treatment of tumor bearing mice** - Tumor-bearing mice were size-matched and divided into groups. The weights of mice were similar within each treatment cohort. Treatment began 3 days after injection of the tumor cells. Four of these groups received i.p. injections of (i) at 50 mg/kg of mAb 13H3 for high dose treatment, (ii) 25 mg/kg of mAb 13H3 for medium dose treatment, (iii) 12.5 mg/kg of mAb 13H3 for low dose treatment or (iv) PBS, respectively. The respective dosages were administered once daily for two weeks followed by four times weekly for four weeks. The remaining group of 8 BALB/c nude mice did not receive any treatment and used as a negative control of the tumor model. Tumor volumes were measured 3-4 times weekly with a caliper and calculated as \( \pi/6 \times \text{length} \times \text{width} \). Tumors were weighed
35 days after inoculation.

**Statistical Analysis**-The SPSS 15 software package (SPSS, Inc., Chicago, IL) was used for statistical analysis. Mean between-group values were compared using the $\chi^2$-test. The independent sample t test was used for ELISA group analysis. The association between the markers and clinicopathologic features was analyzed using $\chi^2$-test or two-sided t-test as appropriate. Receiver operating characteristics (ROC) curves were generated to compare the predictive sensitivity and specificity, and the area under the curve (AUC). The survival rates were assessed by the Kaplan–Meier method and compared by the log-rank test. Statistical significance was set at $P < 0.05$ (two-tailed). All comparisons were two-tailed, and p values of $< 0.05$ were considered significant.

**RESULTS**

The functional antibody library was established by immunizing mouse with lung cancer cells from lung cancer patients-We used lung cancer cells, which were isolated from 20 fresh primary lung cancer tissues, for the immunization of five mice. At 12 months after immunization, the serum from immunized mice was collected for cell immunofluorescence and functional identification. The serum of #1 mouse showed the highest immunofluorescent intensity with lung cancer cell line A549 in vitro, which was selected for proliferation assay(Figure.1A). The serum of mouse #1 of 1:200 dilution can significantly suppressed ANIP-973 and A549 cell growth
To identify whether antibodies in the serum could recognize the secreted proteins, we collected the culture medium of lung cancer cells for Western Blot analysis and probed with immunized serum. The results revealed that proteins in the condition medium were identified by the SDS-PAGE, which were also recognized by the immunized serum of #1 mouse (Figure.1C). Then, the splenic cells of #1 mouse were used to fuse with SP2/0 cells, which were maintained in HAT medium supplemented with 2.5% methylcellulose in an atmosphere of 5% CO₂ at 37°C. After cultivating for 8-10 days, the monoclonal library containing 1160 clones were established for screening secreted proteins.

**Functional secreted Protein Screen by Antibody Library against lung cancer cells**

We screened the library for monoclonal antibodies that bound specifically to secreted protein by ELISA, which were pre-coated with the proteins in the condition medium. From the 1160 mAbs, a total of 47 mAbs showed reactivity with the secreted protein, which were subjected to immunohistochemical analysis with 40 carcinoma tissues or normal human lung tissues. From them, 20 of these mAbs can specifically react with lung cancer tissues not the normal lung tissues. Next, the remaining 20 mAbs were selected for further analysis with all cases the remaining cases. Six clones of these antibodies including 5B8, 5D4, 4D10, 2B4, 2F9 and 13H3 that strongly reacted with cancerous tissues and seldom exhibited normal lung tissue staining(<10%) (Table2). Figure.1D shows the typical view of immunohistochemical staining of 13H3 with four types of lung cancer tissues. To get the functional monoclonal antibodies, 20 mAbs were also chosen for proliferation analysis. Among
the 6 clones, 13H3 antibody demonstrated the highest inhibitory rate of proliferation, which was selected for identifying the target antigen.

**The antigen of mAb 13H3 is identified as Mac-2BP.**

We collected the CM samples harvested from two lung cancer cell lines (A549 and ANIP-973) and lung cancer cells from cancer specimens for western blot detected by 13H3 antibody. The result revealed that 13H3 could recognize the corresponding protein in three kinds of CM. Thus we collected the CM form lung cancer cell line A549 for identifying target antigen (Figure.2A). The molecular weight of antigen recognized by 13H3 is approximately 85 kDa as determined by Western blotting. After immunoprecipitation, the protein was separated by SDS-PAGE (Figure.2B). Then, the aimed protein bands were individually excised, in-gel digested with trypsin, and analyzed by MALDI-TOF peptide mass fingerprinting (PMF). Three bands bound by 13H3 antibody were identified as Mac-2BP (gi|5031863) (Figure.2C), Vimentin (gi|340219) (Figure S1) and Integrin-linked Kinase-2 (gi|8648885) (Figure S2), respectively. To verify the result of mass spectrometry, commercial anti-human Mac-2BP antibody and 13H3 were subjected to immunoprecipitation and Western blot analysis. The result revealed that the immunoprecipitate by 13H3 mAb could be recognized by commercial anti-human Mac-2BP, and the immunoprecipitate by anti-Mac-2BP could be recognized by 13H3 (Figure.2D). The above results verified that 13H3 can specifically recognize the Mac-2BP in the CM and cancer specimens.

**Neutralization of Mac-2 BP with mAb 13H3 suppressed lung cancer cells proliferation in vitro and tumor growth in vivo.**
We purified 13H3 antibody to evaluate the effect of 13H3 on human lung cancer cells A549 and ANIP-973 growth in vitro. As shown in Figure 3A, 13H3 led to strong inhibition of cell proliferation in 2 lung cancer cell lines in a dose-dependent manner. Then, nude mice bearing Anip973 xenografts were treated by 13H3 mAb. As shown in Figure 3B, 13H3 mAb effectively reduced the tumor weights in a dose-dependent manner with the inhibition rates of 60.14%, 54.86% and 43.84% for the high(50mg/ml), moderate(25mg/ml) and low (12.5mg/ml) dose of 13H3, respectively. These results indicated that 13H3 was a functional mAb, which might be used as potential target for lung cancer treatment.

**Knockdown of Mac-2BP Expression in lung Cancer Cells Inhibits Proliferation.**

We examined the effects of Mac-2 BP knock-down on cell proliferation. Firstly, we detected the level of Mac-2BP in all six lung cancer cell lines including Glc-P, Glc-82, ANIP-973, A549, H1299 and H157(Figure.3C). Then, we employed the siRNAs technology to specifically decrease the level of Mac-2BP in ANIP-973 and A549 cells(Figure.3D). Knockdown of Mac-2BP in cell clones induced drastic growth retardation as compared with the control cells (Figure 3E). Furthermore, we used the 5-ethynyl-2-deoxyuridine (EdU) DNA Cell Proliferation Kit to provide an indication of cell proliferation rate. The result showed that the number of EdU+ cells was approximately 34% or 38% lower in A549 SiRNA-Mac2-BP cells and ANIP-973 Si-RNA-Mac2-BP cells than in control cells (Figure 3F).

**Downregulation of Mac-2BP lead to G1 arrest and overexpression of p27.**

To unravel the mechanism of cell growth inhibition induced by Mac-2BP depletion,
we examined the cell cycle by FACS analysis. A significant increase of G1 peak (72.3%, 66.2%) was observed in ANIP-973 or A549 SiRNA-Mac-2BP cells, compared with the control cells (49.1%, 47.6%), suggesting G1 cell cycle arrest (Figure 4A). Therefore, we assessed the effect of knockdown of Mac-2BP on cell cycle regulatory molecules involving in G1 phase. For one thing, Cyclin D1 is over-expressed in many tumors, and Cyclin D1 with the Cdk4 and Cdk6 are important for positive regulators of G1 to S-phase transition[13]. Cyclin E/CDK2 is also active in mid-G1 close to the restriction point. For another, the CDKi’s including p21 and p27 can cause a blockade of the G1 to S transition[14]. Hence, we detected the level of cyclin D1, cyclin E, CDK2, CDK4, CDK6 and p27, p21 in parent cells and Mac-2bp SiRNA cells. Our results indicated downregulation of Mac-2BP was coupled with high expression of p27 and low expression of cyclin D1. There was no significant alteration in the expression levels of CDK2, CDK4, CDK6 in Mac-2BP SiRNA cells as compared with the control cells.

Mac-2BP has been identified independently as a ligand of galactin-3, which may play important roles in adhesion, invasion and metastasis through interaction with galectin-3. Then, we determined the expression of galactin-3 in ANIP-973 and A549 Mac-2BP SiRNA cells. The result showed that down-regulation of Mac-2BP was coupled with low expression of galactin-3 and that the expression of p21 could not be detected. And, there were no changes in cyclin E expression compared with in control cells (Fig. 4B, 4C). In addition, we collected cancer tissues from antibody treatment experiments and assessed the expression Mac-2BP, galectin-3, cyclinD1 and p27. As
expected, we found the expression of Mac-2BP, cyclinD1 and galectin-3 was decreased and the expression of p27 was highly expressed (Figure 4D).

**Elevated Serum Levels of Mac-2 BP in lung cancer Patients.**

As mentioned, Mac-2 BP is a secreted protein, which may be detectable in serum. To investigate the potential of Mac-2BP as a serological marker for lung cancer, we analyzed the level of Mac-2 BP in 320 lung cancer patients and 80 healthy donors by ELISA. Serum concentrations of Mac-2BP in lung cancer patients (6.72, 0.14-19.89ug/mL) were significantly higher than those in healthy controls(1.28, 0.16-28.64ug/mL) (P<0.05). When classified according to histologic type of lung cancer, the serum levels of Mac-2BP were 10.31ug/mL (0.14-26.9ug/mL) in small cell lung cancer, and 5.61ug/mL (0.91-19.89ug/mL) in non-small cell lung cancer and the differences were significant (P<0.05)(Figure.5A).

**ROC analysis of Serum Mac-2BP.**

ROC curves for serum Mac-2BP concentrations were constructed to determine the cutoff values. The approximate area under the ROC curve assessing serum Mac-2BP as a diagnostic tool for the detection of lung cancer against normal controls was 0.874, at a cutoff value of 6ug/ml. (Figure 5B). Then, we analyzed the values of Mac-2BP serum level in discriminating NSCLC from lung cancer patients. As the Figure 5C shows the ROC curves for SCLC versus NSCLC and the area under the ROC curve was 0.724. These observations indicated that Mac-2BP might be a diagnostic biomarker of lung cancer, especially for SCLC.

**Clinical significance and prognostic value of Mac-2BP as a serologic biomarker**
for lung cancer.

The clinical profiles of patients with a serum Mac-2BP level above the cut-off level (6ug/ml) are shown in Table 3. At the cutoff of 6ug/mL, the positive rates in Small-cell lung cancer, Adenocarcinoma, Squamous cell carcinoma and Large cell carcinoma were 82.9% (63 of 76), 45.4% (54 of 119), 62.6% (72 of 115) and 70% (7 of 10), respectively. The serum concentrations of Mac-2BP did not differ significantly with age (P=0.482), gender (P=0.77). The concentrations of Mac-2BP were significantly correlated with tumor histology (P=0.000), lymph node metastases (P = 0.022), and distant metastases (P= 0.001)(Table3). Thus, the elevation of serum Mac-2BP levels appears to be closely associated with lung cancer progression. Multivariate survival analysis performed using a Cox proportional hazard model showed that Mac-2BP was an independent prognostic factor for lung cancer overall survival (Table4). Kaplan–Meier survival analysis showed that increased Mac-2BP concentrations of 6ug/ml or higher in lung cancer serum were correlated with poor overall survival times (P = 0.000) (Figure.5D).

DISCUSSION

Lung cancer is a leading cause of cancer death worldwide. Early detection of lung cancer greatly improves patient survival. Serum biomarker tests have great potential to facilitate the early detection. Over the past several decades, some biomarkers including Cancinoembryonic antigen (CEA), cytokeratin 19 fragment antigen 21-1 (CYFRA21-1), squamous cell carcinoma antigen (SCC), neuron specific enolase (NSE) are commonly used in diagnosing lung cancer [15-18]. Unfortunately, most
biomarkers are limited by their low specificity and/or sensitivity. Therefore, there is an urgent need to find out other potential biomarkers in clinical practice.

Proteomic technologies have been introduced to identify markers associated with cancer. The current proteomic strategy is just comparing serum from cancer patients with those from normal controls. However, the prospects of blood proteomics are challenged by the fact that blood is a very complex body fluid containing large amounts of proteins. Some abundant blood proteins including albumin immunoglobulin may mask the low abundance proteins, which are usually potential markers [19]. To avoid the major limitations in blood proteomics, recent studies also focused on analyzing the conditioned media from cancer cells with different phenotype to identify secreted proteins associated with drug resistance or metastasis. For example, Chen et al. analyzed the secretomes of a primary NSCLC cell line and its brain metastatic subline and found L-lactate dehydrogenase B chain (LDHB) was associated with the metastasis[20]. Unfortunately present methods easily got the differential expression protein and rarely obtained the functional protein, which might also be the targets in cancer treatment.

In the present study, we isolated primary lung cancer cells from four histological types of lung cancer tissues for a short time (24 hour) culture, which were used for immunization. Then, we constructed a large capacity of hybridoma monoclonal antibody library containing 1260 monoclones. To specifically obtain the monoclonal antibodies recognizing secreted protein, we condensed the conditioned medium (CM) of lung cancer primary cells from cancer tissues, which was coated with the plate for
ELISA screening. From it, we identified 47 mAbs, which could react with the secreted protein. This result might ascribe to the low concentrations of secreted proteins in the culture medium, which could not be detected by ELISA method. To acquire lung cancer specific antibody, the candidate monoclonal antibodies were firstly subjected to immunohistochemical assay in 40 lung cancer tissues. Then 20 of these antibodies predominantly reacted with lung cancer tissues were selected for immunohistochemistry in the remaining 65 specimens. The result showed that the positive rate of antigen recognized by 13H3 was 91.4% (96 of 105) and seldom reacting with lung tissues. Further studies also showed 13H3 antibody could inhibit lung cancer cells proliferation in vitro and effectively reduce the tumor weights in vivo in a dose-dependent manner.

Using immunoprecipitation and mass spectrometry, the antigen of 13H3 mAb was identified as Mac-2BP, a ligand of galectin-3[21]. Mac-2BP plays key roles in proliferation in regulating growth and motility of OSCC cells and mediating homotypic adhesion of melanoma cells and the formation of multicellular aggregates[22, 23]. Galectin-3 is involved in cell cycle regulation through induction of cyclin D1 and c-Myc when translocated to the nucleus[24, 25]. Recent studies also showed that Mac-2BP and galectin-3 were found to be deposited in extracellular matrix and the interaction between them is associated with cancer progression [26, 27]. In this study, we firstly elucidated the mechanism of Mac-2BP in lung cancer proliferation. We evaluated the expression of Mac-2BP in 6 lung cancer cells and selected ANIP-973 and A549 for the following study. As mentioned in Results, we
knocked down its expression using SiRNA targeting Mac-2BP and found that down-regulation of Mac-2BP significantly decreased the proliferation of lung cancer cells. Flow cytometric analysis proved that the lower proliferation rate of knockdown cells seems to be associated with G1 phase arrest. Previous studies have reported that Cyclin D1 is over-expressed in many tumors, and Cyclin D1 with the Cdk4 and Cdk6 are important for positive regulators of G1 to S-phase transition. Cyclin E/CDK2 is active in mid-G1 close to the restriction point. On the contrary the CDKi’s including p21 and p27 can cause a blockade of the G1 to S transition [28, 29]. Therefore, we detected the level of cyclin D1, cyclin E, CDK2, CDK4, CDK6 and p27, p21 in parent cells and Mac-2bp SiRNA cells. Our results indicated G1 cell cycle arrest by knockdown of Mac-2BP was mediated through the increased expression of p27 and a simultaneous decrease in cyclin D1. There was no significant alteration in the expression levels of CDK2, CDK4, CDK6 in Mac-2BP SiRNA cells as compared with the control cells. The results suggested that Mac-2BP might take an active part in cell cycle regulation.

Thus, we collected cancer tissues from antibody treatment experiments and assessed the expression galectin-3. As expected, we found the expression of Mac-2 BP was decreased and the expression of galectin-3 was highly downregulated. We speculated that 13H3 mAb could inhibit the interaction between Mac-2 BP and galectin-3 and both of them were degrade quickly, which might suppress the proliferation of lung cancer cells by inducing G1 phase arrest. But the precise mechanisms should be further explored in the future.
Mac-2BP is a secreted glycoprotein and elevated levels of Mac-2BP have been observed in patients with different types of cancer including breast cancer[30], biliary tract carcinoma[31], colon cancer[32] and non-Hodgkin's lymphoma[33]. Although the association between Mac-2BP in the tumor tissue and the presence of metastasis has been confirmed in a series of 72 NSCLC patients by immunohistochemistry, there is no information about serum Mac-2 BP status in lung cancer and its potential clinical application. Therefore, we analyzed sera obtained from 320 lung cancer patients and 80 healthy donors using ELISA to investigate the potential value of Mac-2 BP as a serological marker for lung cancer. Serum concentration of Mac-2BP was found to be elevated in four lung cancer types, especially in NSCLC. Mac-2 BP might be a diagnostic biomarker of lung cancer and area under the ROC curve (AUC) was 0.874. Importantly, Mac-2 BP also showed the diagnostic ability for detecting SCLC (AUC=0.724). According to our study, the diagnostic power of Mac-2 BP is super to the present four tumor markers including SCC, CEA, Cyfra21-1and NSE for lung cancer reported by others[16, 34, 35]. In addition, the concentrations of Mac-2 BP were significantly correlated with tumor histology (P=0.000), lymph node metastases (P=0.022), distant metastases (P=0.001) and was an independent prognostic factor. Our results indicate that Mac-2 BP is a powerful diagnostic biomarker of lung cancer.

In conclusion, functional monoclonal antibody library screening technique is an effective antibody library-based proteomics approach. Using this approach, we successfully identified a functional lung cancer gene Mac-2 BP and a functional
monoclonal antibody 13H3. Both of Mac-2 BP and 13H3 are of great significance in
diagnosis and treatment of lung cancer.
FIGURE LEGENDS:

Figure 1. The establishment and screen of antibody library. A. The immunized serum of #1 mouse showed the highest immunofluorescent intensity with lung cancer cells. B. The proliferation assay was performed with serum of #1 mice at 1:200 dilutions. C. Left panel, the conditioned media of lung cancer cells (25 μg protein) were resolved on 9-15% gradient SDS gels and coomassie stained. Right panel, the proteins in conditioned media were recognized by immunized serum using western blot. D. Immunocytochemistry of mAb 13H3 with four types of human lung cancer tissues and normal lung tissues (n=40).

Figure 2. Mac-2BP is identified as the antigen of mAb 13H3. A. Western blot analysis (20 μg protein) of the conditioned medium from A549, ANIP-973 and lung cancer tissues using 13H3. B. Left, specific tumor antigen that was visualized on silver stained method; Right, validated by Western blotting using 13H3 as the primary antibody. C. Mass spectrometric analysis of the tumor antigen. Upper panel showed the peptide mass fingerprint of Mac-2BP; lower panel showed the result for database search showing the detected peptide fragments and the peptide coverage. D. The immunoprecipitate by anti-Mac-2BP could be recognized by 13H3.

Figure 3. Mac-2BP involved in proliferation in vitro and tumor growth in vivo.

A. Proliferation assay of ANIP-973 and A549 cells. The 13H3 antibody significantly suppresses ANIP-973 and A549 cells proliferation in a dose-dependent manner.

B. The expression of MAC2BP expression on six lung cancer cell lines, including Glc-P, Glc-82, ANIP-973, A549, H1299 and H157. E. Semiquantitative reverse
transcription-PCR was performed for amplification of Mac-2BP and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) using specific primers after siRNA transfection. Amplified bands were analyzed by agarose gel electrophoresis. Expression level of Mac-2BP was determined by densitometric analysis after normalizing RNA load. C. EdU labeling showing proliferation of Mac-2BP silenced and control cells. The percentage of positive cells was derived from triplicate samples. D. The effect of 13H3 treatment on the xenograft tumor growth. E. ANIP-973 and A549 cells transfected with Mac-2BP Si-RNA or NC control were seeded in 96-well plates for Cell Counting Kit-8 assay. The results are expressed as the mean optical density of absorbance of two independent experiments.

**Figure 4. Induction of G1 arrest by downregulation of Mac-2BP**

A. The cell clones were cultured in complete medium for 48 h. Cells were then harvested, fixed with ethanol, stained with propidium iodide and analyzed by flow cytometry. The percentages of different cell cycle stages are shown in each panel. B. The levels of Mac-2BP, galectin-3, CDK2, CDK4, CDK6, CyclinD1, CyclinE, p21, p27 and were analyzed by Western blotting in A549 cells transfected with si-RNA-Mac-2 BP or NC control. C. The levels of Mac-2BP, galectin-3, CDK2, CDK4, CDK6, CyclinD1, CyclinE, p21, p27 and were analyzed by Western blotting in ANIP-973 cells transfected with si-RNA-Mac-2 BP or NC control. D. The levels of CyclinD1, p27 and galectin-3 were analyzed by Western blotting in xenograft tumor treated with different dose of 13H3 mAb.
Figure 5. Elevated Mac-2 BP levels in lung cancer serum samples.

A. Serum levels of Mac-2 BP in healthy controls and lung cancer patients. The serum levels of Mac-2 BP in healthy controls (n=80) and Lung cancer patients (n=320) were measured by ELISA. B. ROC curve analysis of the diagnostic efficacy of Mac-2 BP for lung cancer. C. ROC curve analysis of the diagnostic efficacy of Mac-2 BP for SCLC. D. Kaplan Meier survival curve showed correlation between Mac-2 BP serum level and overall survival.
## Table 1: Clinical and pathologic information of 20 lung cancer patients

<table>
<thead>
<tr>
<th>Patients</th>
<th>Gender</th>
<th>Age</th>
<th>TNM stage</th>
<th>Differentiation</th>
<th>Type</th>
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<tr>
<td>1</td>
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<td>Middle</td>
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<tr>
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### Table 2: The immunohistochemical screen of monoclonal antibodies

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<th>Clonal number</th>
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<th>CCK-8 assay</th>
<th>Subtype</th>
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<tr>
<td></td>
<td>lung cancer lung inhibitory rate</td>
<td>heavy chain</td>
<td>light chain</td>
</tr>
<tr>
<td>5B8</td>
<td>69.5% (73/105) 4.7% (5/105)</td>
<td>15%</td>
<td>IgM k</td>
</tr>
<tr>
<td>5D4</td>
<td>78% (82/105) 2.9% (3/105)</td>
<td>19%</td>
<td>IgG1 k</td>
</tr>
<tr>
<td>4D10</td>
<td>80% (84/105) 2.9% (3/105)</td>
<td>20%</td>
<td>IgG1 k</td>
</tr>
<tr>
<td>2B4</td>
<td>85.7% (90/105) 8.6% (9/105)</td>
<td>21%</td>
<td>IgM k</td>
</tr>
<tr>
<td>2F9</td>
<td>92.4% (97/105) 7.6% (8/105)</td>
<td>25%</td>
<td>IgM k</td>
</tr>
<tr>
<td>13H3</td>
<td>98.1 (103/105) 1.9% (2/105)</td>
<td>29%</td>
<td>IgM k</td>
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### Table 3: Clinicopathological characteristics of the 320 lung cancer patients

<table>
<thead>
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<th>Serum Mac-2BP</th>
<th>negative</th>
<th>positive</th>
<th>p value</th>
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<tbody>
<tr>
<td>Sex (Male: Female)</td>
<td>97:27</td>
<td>156:40</td>
<td>0.77</td>
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<tr>
<td>Age</td>
<td>59.58±10.29</td>
<td>60.39±9.81</td>
<td>0.482</td>
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<tr>
<td>Type</td>
<td></td>
<td></td>
<td>0.000*</td>
</tr>
<tr>
<td>SCC</td>
<td>43</td>
<td>72</td>
<td></td>
</tr>
<tr>
<td>AD</td>
<td>65</td>
<td>54</td>
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</tr>
<tr>
<td>LC</td>
<td>3</td>
<td>7</td>
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</tr>
<tr>
<td>SCLC</td>
<td>13</td>
<td>63</td>
<td></td>
</tr>
<tr>
<td>Differentiation</td>
<td></td>
<td></td>
<td>0.008*</td>
</tr>
<tr>
<td>Well</td>
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<td></td>
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<tr>
<td>Moderate</td>
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<td>82</td>
<td></td>
</tr>
<tr>
<td>poor</td>
<td>31</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td>Depth of invasion</td>
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<td></td>
<td>0.006*</td>
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<tr>
<td>T1+T2+T3</td>
<td>96</td>
<td>144</td>
<td></td>
</tr>
<tr>
<td>T4</td>
<td>23</td>
<td>59</td>
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<td>Lymph node involvement</td>
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<td>Metastasis</td>
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<td>M0</td>
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### Table 4: Cox multivariate analysis

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<th>Variables</th>
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<td>Gender</td>
<td>0.901(0.642-1.263)</td>
<td>0.544</td>
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<tr>
<td>Age</td>
<td>1(0.986-1.013)</td>
<td>0.971</td>
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<tr>
<td>Mac2-BP</td>
<td>1.868(1.382-2.524)</td>
<td>0.000*</td>
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<tr>
<td>Type</td>
<td>0.729(0.496-1.071)</td>
<td>0.107</td>
</tr>
<tr>
<td>T</td>
<td>0.958(0.686-1.338)</td>
<td>0.803</td>
</tr>
<tr>
<td>N</td>
<td>1.24(0.899-1.712)</td>
<td>0.19</td>
</tr>
<tr>
<td>M</td>
<td>1.051(0.722-1.529)</td>
<td>0.796</td>
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</table>
REFERENCE


**Figure 2**

**A**

<table>
<thead>
<tr>
<th>Protein</th>
<th>Mass (kDa)</th>
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<tr>
<td>A549</td>
<td>170</td>
</tr>
<tr>
<td>ANIP-973</td>
<td>130</td>
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<tr>
<td>Cancer cells isolated from specimens</td>
<td>95</td>
</tr>
<tr>
<td>13H3</td>
<td>59</td>
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</tbody>
</table>

**B**

Western blot

**C**

[Graph showing protein mass and peptide distribution]

**D**

170KD, 130KD, 95KD, 59KD, 55KD, 43KD, 34KD

**13H3 mouse IgG**

**Mac-2BP goat IgG**
Figure 5

A

Serum Mac2-bp (ug)

P < 0.05

Normal  AD  LC  SCC  SCLC

B

Sensitivity

AUC = 0.874

1-Specificity

0.0  0.2  0.4  0.6  0.8  1.0

C

Sensitivity

AUC = 0.742

1-Specificity

0.0  0.2  0.4  0.6  0.8  1.0

D

Cum Survival

Overall survival

Mac2-BP > 6 ug/ml

Mac2-BP ≤ 6 ug/ml

p = 0.000