Original Article

Immunological Detection of Severe Acute Respiratory Syndrome Coronavirus by Monoclonal Antibodies

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SUMMARY: In order to establish immunological detection methods for severe acute respiratory syndrome coronavirus (SARS-CoV), we established monoclonal antibodies directed against structural components of the virus. B cell hybridomas were generated from mice that were hyper-immunized with inactivated SARS-CoV virion. By screening 2,880 generated hybridomas, we established three hybridoma clones that secreted antibodies specific for nucleocapsid protein (N) and 27 clones that secreted antibodies specific for spike protein (S). Among these, four S-protein specific antibodies had in vitro neutralization activity against SARS-CoV infection. These monoclonal antibodies enabled the immunological detection of SARS-CoV by immunofluorescence staining, Western blot or immunohistology. Furthermore, a combination of monoclonal antibodies with different specificities allowed the establishment of a highly sensitive antigen-capture sandwich ELISA system. These monoclonal antibodies would be a useful tool for rapid and specific diagnosis of SARS and also for possible antibody-based treatment of the disease.

INTRODUCTION

The outbreak of severe acute respiratory syndrome (SARS) in 2003, caused by SARS coronavirus (SARS-CoV)(1,2), ultimately led to 8,000 people becoming infected, 916 of whom died (3; http://who.int/csr/sars/country/en/country2003_08_15.pdf). Even though the WHO announced an end to the epidemic (4; http://www.who.int/entity/csr/sars/resources/en/SARSReferenceLab1.pdf), the threat of re-emergence persists due to the absence of a vaccine, and inability of health services to rapidly detect and specifically diagnose the disease. One of the critical issues in the management of clinical patients and control of the pandemic is a system of early diagnosis that distinguishes SARS from other types of pulmonary infections. As an epidemiological history of contact with SARS patients is not always provable and there are no clinical signs unique to SARS patients (5), confirmatory diagnosis relies primarily on laboratory tests.

To date, viral shedding of SARS-CoV has been extensively studied to improve diagnosis and infectious control (6-8). Maximum virus shedding takes place between day 12 and day 14 of disease onset. For most acute respiratory viral infections, viral shedding occurs within the first few days from the nasopharyngeal tissue and soon after at the upper respiratory tract, but seldom lasts for more than 10 days (6-8). The peak of shedding in stools occurs a few days after respiratory shedding and remains high even after 3 weeks (7, 8). SARS-CoV was detected in patients’ plasma samples within several days of the onset of fever, sometimes at levels equivalent to those recorded for nasopharyngeal aspirates (6, 9).

Previously, during the outbreak in Hong Kong (8), laboratory diagnosis for SARS virus infection was based on a combination of serologic tests, reverse transcription-polymerase chain reaction (RT-PCR), and virus isolation. IgG seroconversion among those infected was 93% by day 28 (5), suggesting that while antibody seroconversion provides reliable proof of infection (5,10); it is, however, not suitable for early diagnosis (11). Among patients in whom the serological evidence could be retrospectively examined, RT-PCR provided about 60% of the diagnostic yield using tracheal aspirates and stools for the first 2 weeks after the onset of illness (8). Although the availability of data that compares the diagnostic yield of various specimen types is still limited, it has been suggested that a combination of stool samples and pooled throat and nasal swab specimens provides reagents for safe and high-yield SARS-CoV detection (8). Furthermore, in addition to RT-PCR on respiratory and fecal samples, serology is needed to confirm the diagnosis of SARS-CoV infection in most cases.

Based on clinical experience, several options have been considered in the quest to develop the capacity to accurately diagnose SARS-CoV infection, including molecular biology techniques and serological tests such as antigen-captured ELISA assay and immunofluorescence assay to detect virus-infected cells in respiratory swabs (5-12). The preparation of monoclonal antibodies (mAbs) is considered to be valuable especially for serological testing.

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In this paper we report the successful establishment and
the characterization of mAb against SARS-CoV structural
components. These mAbs enabled the general immunological
detection of SARS-CoV, by the methods such as immuno-
fluorescent staining, Western blotting, and immunohistology,
in addition to the construction of highly sensitive antigen-
capture sandwich ELISA.

MATERIALS AND METHODS

Virus and cell culture: SARS-CoV (HKU-39849) was
kindly supplied by Dr. J. S. M. Peiris, Department of Micro-
biology, the University of Hong Kong. The live virus was
manipulated under the physical containment level P3. For
the purification of the virion, the day-2 culture supernatant
was centrifuged at 8,000 g for 30 min to remove cell
debris. The virion in the supernatant was precipitated with
8% polyethylene glycol/0.5 M NaCl, and further purified by
20%/60%-discontinuous sucrose density centrifuga-
tion. This fraction was inactivated by UV-irradiation (260 nm,
4.75 J/cm²), and used as UV-inactivated SARS-CoV fraction.
We and others confirmed that this condition completely
inactivates SARS-CoV(13,14).

Production of mAbs: BALB/c mice (9-week old females,
Japan SLC) were immunized subcutaneously with 20 µg
of UV-inactivated SARS-CoV using Freund’s Complete Ad-
juvant (FCA, Sigma, St. Louis, Mo., USA). After 2 weeks,
the mice were boosted with a subcutaneous injection of 5 µg
of UV-inactivated SARS-CoV using Freund’s Incomplete
Adjuvant (FIA, Sigma). On day-3 after the boost, sera
were tested by ELISA for the antibody titer against
SARS-CoV. The two mice showing highest antibody titer were
further boosted intravenously with 5 µg of the inactivated
virus 14 days after the previous boost. This immunization
schedule was called protocol-1. In protocol-2 the booster
injection was repeated two more times before the final boost.
Three days after the final boost, spleens from two mice were
excised and the splenocytes were fused with Sp2/O-Ag14
myeloma by the polyethylene glycol method of Kozbor
and Roder (15). The fused cells from the two spleens were
cultured and HAT-selected on twenty 96-well plates. The
first screening was conducted by ELISA using SARS-CoV
infected Vero E6 cell lysate as the antigen. In this first screen-
ing, the ELISA with uninfected Vero E6 cell lysate was used
as the negative control. After the virus was inactivated by
UV-irradiation, cell lysates were prepared by NP-40 lysis
buffer (1% NP-40/150 mM NaCl/ 50 mM Tris, pH 7.5)
followed by centrifugation at 15,000 rpm for 20 min to
remove the cell debris. The supernatant was diluted 100-fold
using ELISA-coating buffer (50 mM sodium bicarbonate,
pH 9.6) and the ELISA plates (Dynatech, Chantilly, Va., USA)
were coated at 4°C overnight. After blocking with 1% ovalbumin in PBS-Tween (10 mM phosphate buffer, 140
mM NaCl, 0.05% Tween 20, pH 7.5) for 1 h, the culture
supernatants from HAT-selected hybridomas were added and
incubated for 1 h. After washing with PBS-Tween, the
bound antibodies were detected with alkaline phosphatase-
conjugated anti-mouse IgG (1:2000, Zymed, South San
Francisco, Calif., USA) using p-nitrophenyl phosphate
(PNPP) as a substrate. The second screening was conducted by
ELISA using the cell lysates of chick embryonic fibroblast
(CEF) cells that were transfected by vaccinia virus vector
containing the gene either of SARS-CoV spike (S) or
nucleocapsid (N) proteins.

Recombinant virus proteins: Genomic RNA was extracted
from SARS-CoV strain HKU39849 and reverse transcribed
to cDNA. The corresponding open reading frames (ORF)
were amplified by PCR and cloned into the transfer vector, pDIsgptmH5, which also harbored Escherichia coli xantine-guanine phosphoribosyltransferase
under the control of vaccinia virus p7.5 promoter in the
cloning site of pUC/DIs (16). The recombinant clones of
attenuated vaccinia virus, DIs, which harbored each ORF were
obtained by homologous recombination induced in DIs-
infected-, pDIsgptmH5-transfected CEF cells. The detailed
protocol will be published elsewhere.

Neutralization assay: The known tissue culture infectious
dose (TCID) of SARS-CoV was incubated for 1 h in the
presence or absence of the purified mAb serially diluted 10-
fold, and then added to Vero E6 cell culture grown to confluence
in a 96-well microtiter plate. As a control, mAb against N
protein was added to the culture. After 48 hr, cells were fixed
with 10% formaldehyde and stained with crystal violet to
visualize the cytopathic effect induced by the virus (17). Neutralization antibody titers were expressed as the mini-
mum concentration of purified immunoglobulin that inhibits
cytopathic effect.

Western blot: UV-inactivated purified SARS-CoV virion
(0.5 µg/lane) (13) was loaded on SDS-PAGE under reduced
conditions. Proteins were transferred to the PVDF membrane
(Genetics, Tokyo, Japan). After blocking with BlockAce
(Snow Brand Milk Products Co., Ltd., Tokyo, Japan) reagent,
the membranes were reacted with the mAb or the diluted sera (1:1000) that had been obtained from mice inoculated
with UV-irradiated SARS-CoV. After washing, the membrane
was reacted with peroxidase-conjugated Fab’ (γ-galwashed
mouse IgG (H+L) (1:20,000 Jackson Immuno Research, West
Grove, Pa., USA), and the bands were visualized using chemi-
luminescent reagents (Amasham Biosciences, Piscataway,
N.J., USA) on the X-ray film (Kodak, Rochester, N.Y., USA).

Purification and biotinylation of mAbs: Hybridomas
were grown in Hybirdoma-SFM medium (Invitorgen, Carlsbad,
Calif., USA) supplemented with recombinant IL-6 (18) and
cenulin (100 U/mL)/streptomycin (100 µg/mL). The culture
supernatants were harvested, added with 1/100 volume of 1 M Tris-HCl (pH 7.4) and 1/500 volume of 10% NaN³,
and directly loaded on the Protein G-Sepharose 6B column
(Amersham Biosciences). The column was washed with PBS
and eluted with Glycine/HCl (pH 2.8). After measuring the
OD280 of the fractions, protein containing fractions were
poled and added with an equal volume of saturated
(NH₄)₂SO₄. Precipitated proteins were dissolved in PBS,
dialysed against PBS and stored at −20°C. The purified
antibodies were biotinylated using sulfo-NHS-LC-biotin
(Pierce, Rockford, Ill., USA) according to the manufacturer’s
protocol.

Antigen-capture ELISA: The purified mAb for the antigen-
capture was immobilized on the microplate (Immulon 2,
Dynatech) by incubating 4 µg/mL antibody in 50 mM sodium
bicarbonate buffer (pH 8.6) at 4°C overnight. The microplate
was blocked with 1% BSA, washed with PBS-Tween, and
reacted with serial dilution of UV-inactivated purified SARS-
CoV for 1 h at room temperature. After washing with PBS-
Tween, wells were reacted with biotinylated probing mAb
(0.1 µg/mL) for 1 h at room temperature. After washing, wells
were reacted with β-D-galactosidase-labeled streptavidine
(Zymed) for 1 h at room temperature. After washing,
fluorescent substrate 4-methylumberyferyl-β-D-galactoside (Sigma-Aldrich, St. Louis, Mo., USA) was added and the substrate was incubated for 2 h at 37°C. The reaction was stopped by adding 0.1M Glycine-NaOH (pH 10.2) and the fluorescence (FU) of the reaction product, 4-methylumberyferyl, was measured using FluoroScan (Flow Laboratories Inc., Inglewood, Calif., USA).

**Histology:** Formaldehyde-fixed human lung tissue that was RT-PCR positive for SARS-CoV (19) and lung from a SARS-CoV infected macaque were embedded in paraffin and sectioned using the standard method. After de-paraffinization by standard method, the sections were soaked with 0.1 M citrate-buffer (pH 6.0) and autoclaved for 10 min at 121°C to inactivate viruses. Endogenous peroxidase was inactivated by 0.3% hydrogen peroxide for 30 min at room temperature. After blocking with 5% normal goat serum for 10 min, sections were incubated with the mAb at 4°C overnight. The bound antibody was detected by biotinylated anti-mouse IgG followed by peroxidase-labeled streptavidin (LSAB2 kit, DakoCytomation, Kyoto, Japan) and visualized with 0.2 mg/mL 3,3’-diaminobenzidine in 0.015% hydrogen peroxide/0.05M Tris-HCl (pH 7.6). The sections were counterstained with hematoxylin.

**RESULTS**

In order to establish the hybridomas that secrete specific mAbs to SARS-CoV, we immunized BALB/c mice with purified SARS-CoV whole virion fraction. The virus was inactivated by UV-irradiation to avoid a change in antigenicity presumably caused by aldehyde-fixation or detergent-solubilization. The immunization protocols used were those of the standard method in which the boost administrations were repeated twice (protocol-1) or four times (protocol-2) with 2-week intervals using FCA/FIA as an adjuvant (see Materials and Methods). Three days after the final boost, a single cell suspension was prepared from two spleens of immunized mice and fused with SP-2/O myeloma by a polyethylene-glycol method, the fused cells were then HAT-selected (15).

In the experiment with immunization protocol-1, we found that the culture supernatants from 28 of the 1,920 wells were strong-positive in ELISA testing in which the cell-lysate of SARS-CoV infected Vero E6 cells was used as a coated antigen (Table 1). As a negative control, we used uninfected Vero E6 cell-lysatse as the antigen. Wells that showed a positive reaction were omitted from the count. Among the 28 wells, 19 reacted to vaccinia vector-based recombinant-S-protein and three reacted to recombinant-N-protein. These hybridomas were successfully cloned by a repeated limiting dilution method. The remaining six wells did not give rise to a significant positive signal to recombinant-S, -N or -M proteins. One anti-S mAb cross-reacted to porcine transmissible gastroenteritis virus (TGEV) and this clone was also characterized further.

In a previous report that studied human IgG avidity maturation after rubella vaccination, high-avidity antibodies were shown to be very low avidity in this assay system. The avidities of these cloned mAbs were tested by avidity-ELISA in the presence of urea. Although in the presence of 6M urea some anti-S mAbs retained 18–35% of the original reactivity, less than 10% of the original reactivity remained in the presence of 8M urea (Table 2). Three anti-N mAbs showed a very low avidity index in this assay system.

In order to confirm the negative results for normal lungs and several specimens from pneumonia patients including cases complicated by measles, influenza type A, herpes-simplex and herpes zoster.

The mAbs that worked for immunohistochemistry, i.e.,

<table>
<thead>
<tr>
<th>Immobilized antigen</th>
<th>Experiment-1</th>
<th>Experiment-2</th>
<th>Total</th>
</tr>
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<tbody>
<tr>
<td>(Total wells assayed)</td>
<td>1,920</td>
<td>960</td>
<td>2,880</td>
</tr>
<tr>
<td>SARS-CoV infected Vero cell-lysate</td>
<td>28</td>
<td>14</td>
<td>42</td>
</tr>
<tr>
<td>Recombinant - S</td>
<td>19</td>
<td>7</td>
<td>26</td>
</tr>
<tr>
<td>- N</td>
<td>3</td>
<td>0</td>
<td>3</td>
</tr>
</tbody>
</table>

1) Immunization protocol-1
2) Immunization protocol-2

Table 1. Summary of the first hybridoma screening by ELISA

<table>
<thead>
<tr>
<th>Clone</th>
<th>Epitope</th>
<th>Avidity Index (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>4M urea</td>
<td>6M urea</td>
</tr>
<tr>
<td>Experiment-1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SKOT-7</td>
<td>N</td>
<td>1.6</td>
</tr>
<tr>
<td>SKOT-8</td>
<td>N</td>
<td>2.3</td>
</tr>
<tr>
<td>SKOT-9</td>
<td>N</td>
<td></td>
</tr>
<tr>
<td>SKOT-10</td>
<td>S</td>
<td>45.5</td>
</tr>
<tr>
<td>SOAT-5</td>
<td>S</td>
<td>73.4</td>
</tr>
<tr>
<td>SOAT-13</td>
<td>S</td>
<td>63.8</td>
</tr>
<tr>
<td>Experiment-2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SOAT-5</td>
<td>S</td>
<td>51.3</td>
</tr>
<tr>
<td>SOAT-13</td>
<td>S</td>
<td>77.0</td>
</tr>
</tbody>
</table>

Table 2. Avidity ELISA
SKOT-7, -8 and -9 also worked for Western-blot detection of the viral proteins (Fig. 3). Anti-N mAbs detected a band of 50 kDa that corresponds to the calculated molecular weight of SARS-CoV N-protein. In some experiments with longer exposure, a band with an apparent molecular weight of 120 kDa was also detected. None of the anti-S mAbs worked in the Western blot, suggesting that the major antigenic determinants of the S-protein are ‘conformational’ epitopes.

We tested the in vitro neutralizing activities of anti-S mAbs. As shown in Fig. 4, SKOT-20 neutralized in vitro SARS-CoV infection to Vero E6 cells at an antibody concentration of 1 μg/mL. Another anti-S mAb, SKOT-19, which had a low avidity value, also showed similar neutralizing activity. SKOT-10 and -3 also had neutralization activity but required higher antibody concentrations.

Lastly, we tried to construct an antigen-capture detection system for SARS-CoV by sandwich ELISA. In preliminary experiments, we tested all the combinations of two mAbs from the selected eight mAbs to obtain the highest detection sensitivity for purified SARS-CoV virion, and found that the immobilization of SKOT-8 on the ELISA plate followed by the detection with biotinylated SKOT-9 gave the best result (data not shown; see Materials and Methods). In this sandwich ELISA, SARS-CoV protein was successfully detected in a concentration as low as 40 pg/mL (Fig. 5). Since the mAbs were originally raised against SARS-CoV strain HKU39849, we tested the validity of this system for other strains of SARS-CoV. As shown in Fig. 6, it was confirmed that the strains HK14T1WL, CDC00592 and Frankfurt1 were as detectable as HKU39849 using this system.

**DISCUSSION**

We established mAbs against SARS-CoV, which enable...
Among the originally selected 42 mAbs that were positive in ELISA for SARS-CoV infected Vero E6 cell lysate, 26 reacted to recombinant-S-protein and only three reacted to N-protein. We could not find hybridoma secreting mAb to M-protein or other protein components of SARS-CoV. These results suggest that S protein is the dominant target in the antibody response. We observed that none of the anti-S mAbs established worked in Western blot, suggesting that these mAbs may recognize ‘conformational’ epitopes. In contrast, all three anti-N mAbs worked in Western blot and immunohistochemistry, suggesting that these mAbs recognize ‘linear’ epitopes.

We examined whether our mAbs were applicable for immunofluorescence detection of virus-infected cells. In immunofluorescent staining of Vero E6 cells infected with SARS-CoV, anti-N mAbs stained the Golgi body and anti-S mAbs stained the Golgi body and surface membrane. This difference in localization of N- and S-proteins may reflect the common assembly process of coronaviruses (21). Further analysis is needed to clarify sensitivity and specificity in infected cells for clinical use.

During the course of outbreak of SARS-CoV in Hong Kong, it was reported that more than half the patients were not positively diagnosed by RT-PCR (8). Therefore, the diagnosis was finally confirmed by serum specimens in a convalescent-phase at a late stage of illness (8). To overcome this problem, virus shedding patterns have been extensively analyzed, with results showing that respiratory shedding of the virus increases over the first week and viral shedding in stools begins a few days after respiratory shedding (7,8). From this analysis, it is considered that a combination of stool sampling and pooled throat and nasal swab specimens could be good specimens for safe and highly sensitive SARS-CoV detection.

In general, a single diagnostic test is not conclusively reliable, because of the serious potential for false positives and negatives. Considering the limited sensitivity of RT-PCR, serological screening systems other than antibody detection are currently being examined (22,23). ELISA-based antigen captured assays are known to offer high specificity and reproducibility. Antigen-captured assays have been used in the diagnosis and monitoring of disease in cases of infection with dengue virus (24), human immunodeficient virus p24 (25) and Ebola hemorrhagic fever (26) and examined in hepatitis B virus and hepatitis C virus (22,23). In this context, extensive analysis in Ebola hemorrhagic fever suggests that the RT-PCR assay is extremely useful, but should always be utilized in combination with antigen-captured ELISA, which makes the diagnosis more reliable (26).

<table>
<thead>
<tr>
<th>Table 3. Summary of selected hybridoma clones</th>
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<tbody>
<tr>
<td><strong>Clone</strong></td>
</tr>
<tr>
<td>SKOT-7</td>
</tr>
<tr>
<td>SKOT-8</td>
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<tr>
<td>SKOT-9</td>
</tr>
<tr>
<td>SKOT-3</td>
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<tr>
<td>SKOT-10</td>
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<tr>
<td>SKOT-19</td>
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<tr>
<td>SKOT-20</td>
</tr>
<tr>
<td>SOAT-5</td>
</tr>
<tr>
<td>SOAT-13</td>
</tr>
</tbody>
</table>

[^1]: Numbers represent minimum concentration (μg/mL) that exerts neutralization. -, no neutralization activity; ND, not determined.
[^2]: Avidity index at 6M urea (see Materials and Methods).
In the case of SARS-CoV, the assay has been recently evaluated by using mAbs and polyclonal antibodies directed against recombinant SARS-CoV nucleocapsid protein (22,23). A soluble N-protein was observed to be released from infected cells in culture, which led to the opportunity to evaluate the level in serum specimens from infected patients. N-antigen ELISA employing mAbs reproducibly detected 50% of patients on days 3 and 5 after the onset of illness, with a limitation of the detection of the recombinant protein at 50 pg/ml (22). N-antigen ELISA with use of polyclonal antibodies detected 60-50% of nasopharyngeal aspirate and fecal specimens from patients at day 3 to day 24 after the onset of illness, although the signal was relatively weak in fecal samples (22). These results suggest that antigen-captured assay could be useful for the early diagnosis of SARS-CoV infection.

We developed an antigen-capture ELISA system that detects purified SARS-CoV virion at levels as low as 40 pg/mL. The sensitivity of the system, which comprised two anti-N mAbs, seems high enough to detect virus protein in patient sera when compared to a recently reported antigen-capture ELISA system, which detects 100 pg/mL of purified recombinant N protein, successfully determined the virus protein in patient sera (22). We are now improving the sensitivity of the system and checking its applicability in the diagnosis and monitoring of SARS-CoV infection. Although none of our mAbs cross-reacted to human or other animal coronaviruses (229E, TGEV and MHV) by ELISA, it is also important to define the specificity of these mAbs by other techniques such as Western blot and immunofluorescent staining. This issue is currently under investigation.

Two anti-S mAbs, SKOT-19 and -20 demonstrated significant virus neutralizing activity. It would be interesting to address whether these mAbs interfere with the binding of the virion to its recently reported receptor, ACE2 (27). If this were the case, the humanization of these mAbs by means of either CDR-grafting or mouse-human chimeric antibody would be of interest as a possible application for the therapeutic use of these mAbs.

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induces severe acute respiratory syndrome, SARS-CoV.


