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Cell adhesion as a novel approach to determining the cellular binding motif on the severe acute respiratory syndrome coronavirus spike protein

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\textbf{ABSTRACT}

Emerging life threatening pathogens such as severe acute respiratory syndrome-coronavirus (SARS-CoV), avian-origin influenzas H7N9, and the Middle East respiratory syndrome coronavirus (MERS-CoV) have caused a high case-fatality rate and psychological effects on society and the economy. Therefore, a simple, rapid, and safe method to investigate a therapeutic approach against these pathogens is required. In this study, a simple, quick, and safe cell adhesion inhibition assay was developed to determine the potential cellular binding site on the SARS-CoV spike protein. Various synthetic peptides covering the potential binding site helped to minimize further the binding motif to 10–25 residues. Following analyses, 2 peptides spanning the 436–445 and 437–461 amino acids of the spike protein were identified as peptide inhibitor or peptide vaccine candidates against SARS-CoV.

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1. Introduction

Severe acute respiratory syndrome (SARS) first appeared in Asia in 2002, caused by the SARS-coronavirus (SARS-CoV) (Drosten \textit{et al.}, 2003; Holmes, 2003; Peiris \textit{et al.}, 2003). The illness quickly spread worldwide and affected more than 8000 people with a high case-fatality rate (Kuehn, 2013). Newly emerging pathogens such as the avian influenza A virus H7N9 (Wu and Gao, 2013) and MERS-CoV (Middle East Respiratory Syndrome coronavirus, also known as HCoV-EMC) (Kuehn, 2013) are severe public health problems that cause a high case-fatality rate and widespread psychological effects on society and the economy. A simple method is required to investigate a therapeutic approach to these diseases without contacting the biohazardous virus.

Previous studies have completed whole genome sequences (29,751 base pairs) (Marra \textit{et al.}, 2003; Ruan \textit{et al.}, 2003). The data reveal SARS-CoV to be a novel coronavirus that does not belong to any of the 3 known classes of coronavirus (Marra \textit{et al.}, 2003). The genomes encode 23 putative proteins, including known coronavirus proteins such as replicase 1A, replicase 1B, spike glycoprotein (S), small envelope protein, membrane glycoprotein, and nucleocapsid protein (Marra \textit{et al.}, 2003; Ruan \textit{et al.}, 2003). The S gene [nucleotide (nt) 21,492–25,259] encodes a transmembrane glycoprotein on the viral surface and is responsible for binding and entry into the host cells (Marra \textit{et al.}, 2003). The S protein can be divided into 2 functional domains: S1 and S2. The S1 domain contains a receptor-binding domain (RBD) responsible for specific binding to the host receptor (Gallagher and Buchmeier, 2001). The angiotensin-converting enzyme 2 (ACE2) is a functional receptor for SARS-CoV (Li \textit{et al.}, 2003). After binding to ACE2, conformational change of the S protein induces exposure of the S2 domain and assists the virus in fusing to the host cells (Matsuyama and Taguchi, 2002). Therefore, the RBD within the S1 domain is the target for development of virus-entry blocking peptides and neutralizing antibodies to prevent SARS-CoV infections. Previous studies have used various methods to investigate the potential RBD from amino acids spanning 303–537 (Xiao \textit{et al.}, 2003), 270–510 (Babcock \textit{et al.},...
2.2. In this study, a cell-adhesion inhibition assay was established to characterize further the potential RBM of the SARS-CoV S protein. Synthetic peptides spanning the potential binding site were used and the RBM was further narrowed to approximately 10–25 residues. This method is useful for characterizing the potential RBM in the RBD of newly emerged coronaviruses in a quick, simple, and safe manner without contacting live threatening pathogens.

2. Materials and methods

2.1. Construction of S1-1/pGEX-6P-3, S1-2/pGEX-6P-1, and S1-3/pGEX-6P-1 expression plasmids

Using 2 plasmids encoding the SARS-CoV spike S protein-TW1-#18 [nucleotide (nt) (nt 20,704–22,087)] and TW1-#19 (nt 21,826–22,087) as templates, an S1-1 encoding SARS-CoV S protein 1–446 amino acid was produced and amplified by polymerase chain reaction (PCR), using one pair of primers: SS-F1-2 containing the EcoRI site (5′-GGAATTCATGTTATTTCTATTTCTACTCTACTGTTGATG-3′) and SS-R4-3 containing the HindIII site (5′-GAAGGGCTACGGTGCATGCTAACATATTTATATTAAT-3′). Using 2 plasmids encoding the SARS-CoV spike S protein-TW1-#19 (nt 21,826–22,087), the S1 protein and the S2 protein-TW1-#20 (nt 23,077–24,454) as templates, the S1-2 encoding SARS-CoV S protein 447–641 amino acid was produced and amplified by PCR, using one pair of primers: SS-F3-2 containing the HindIII site (5′-GACATGGCAAGCTTGGCCCTT-3′) and SS-R3 containing the Sall site (5′-GAATTGTGGCAGCTCAGCTC-3′). The PCR products of S1-1 and S1-2 were subsequently cloned into pGEX-6P-1 and pGEX-6P-3 (Amersham Biosciences, NJ, USA) through EcoRI/HindIII and HindIII/Sall digestion and ligation, respectively. S1-1 fragment was subcloned into S1-2/pGEX-6P-1 through EcoRI/HindIII digestion and ligation to form S1-3/pGEX-6P-1, using standardized molecular cloning technologies (Sambrook et al., 1989).

2.2. Protein expression and purification

The Escherichia coli culture was performed based on standard methods (Sambrook et al., 1989; Liu et al., 2011; Chang et al., 2012; Tsegv et al., 2013). The glutathione-S-transferase (GST)-containing recombinant protein expression plasmids S1-1/pGEX-6P-3, S1-2/pGEX-6P-1, S1-3/pGEX-6P-1, and pGEX-2KS (Chang et al., 1993, 2002) were transformed into the E. coli strain BL21 (DE3) for expression. The induction and purification of the recombinant proteins were performed using methods modified from previous studies (Chang et al., 1993, 1997b, 2000; Chang and Lo, 2000; Sun et al., 2007). Bacteria containing the expression plasmids were grown in a 2× YT broth (1.6% trypton, 1% yeast extract, and 0.5% NaCl) at 37°C. The recombinant proteins were induced by adding isopropyl β-D-1-thiogalactopyranoside (IPTG) (1 mM) to the medium after the optical density (OD) of the bacterial culture reached 1.0. The optimal induction conditions were incubated at 37°C for 2 h (S1-2/pGEX-6P-1, S1-3/pGEX-6P-1, and pGEX-2KS) and 4 h (S1-1/pGEX-6P-3). For purification, the centrifuged pellets of E. coli cells from a 1-L culture were resuspended in 50 mL 1× phosphate-buffered saline (PBS) containing protease inhibitors-1 mM phenylmethylsulfonyl fluoride (PMSF), 0.7 μg/mL of pepstatin, 1 μg/mL of leupeptin, and 2 mg/mL of lysozyme. The bacterial cells were subsequently disrupted using a French Press. The cell lysates were centrifuged at 10,000 × g for 30 min and passed through 0.45-μm filters to remove insoluble cell components. The filtered-cell lysates were loaded into a glutathione sepharose 4B column and the flow-through was again loaded into the column to ensure saturated binding. The proteins were eluted using an elution buffer (10 mM glutathione; 50 mM Tris–HCl, pH 9.6; 1 mM PMSF; 0.7 μg/mL of pepstatin, and 1 μg/mL of leupeptin) and dialyzed with 1× PBS containing 1 mM PMSF. Consequently, all recombinant proteins used in this study were GST tagged. The protein concentration was determined using a protein assay kit (Bio-Rad, Hercules, CA, USA).

2.3. Antibody generation and purification

New Zealand rabbits were purchased from the National Laboratory Animal Center (Taipei, Taiwan) and maintained in a specific pathogen-free condition in the experimental animal center of Tzu Chi University. The research methods involving the experimental rabbits were performed under strict adherence to the guidelines of the Institutional Animal Care and Use Committee (Tzu Chi University) and the guidance of Animal Protection Law (Taiwan). Using methods modified from previous studies (Harlow and Lane, 1988; Chang et al., 2002; Sun et al., 2007), 250–500 μg of purified recombinant proteins were mixed with complete Freund’s adjuvant and injected into New Zealand rabbits for the first immunization. For boosting, 100–250 μg proteins were mixed with incomplete Freund’s adjuvant and injected in immunization cycles in 3-week intervals. After 5 immunizations, sera from rabbits were isolated and passed through the Protein A column. Immunoglobulin IgG fractions were eluted using an elution buffer (0.1 M citric acid, pH 3.0) and dialyzed with 1× PBS.

2.4. Synthetic peptides

Synthetic peptides GA91 (corresponding to SARS-CoV Spike protein 437–461 amino acids) were produced in linear form. Synthetic peptides GA101 (corresponding to SARS-CoV S protein 436–445 amino acids), GA107 (corresponding to SARS-CoV S protein 466–475 amino acids), and GA142 (corresponding to SARS-CoV S protein 100–109 amino acids) were produced in branched multiple antigenic peptide (M8) form. All synthetic peptides were provided by Genesis Biotech (TaipeiTaiwan).

2.5. Cell culture and cell-adhesion/inhibition assay

Vero E6 cells (ATCC No. C1008) and HeLa cells (ATCC No. CCL-22) were maintained in RPMI 1640 (BioWest, Miami, FL, USA), and NIH3T3 cells (ATCC No. CRL-1658) were maintained in Dulbecco’s modified Eagle’s medium (DMEM) (BioWest) containing 10% heat-inactivated fetal bovine serum (Biological Industries, Beit Haemek, Israel), 2 mM l-glutamine (BioWest), 100 U/L of penicillin (BioWest), 100 mg/mL of streptomycin (BioWest), and non-essential amino acid (BioWest). Cell adhesion and inhibition assays were performed according to our previously described methods (Chang et al., 1993, 1997a; Chang and Lo, 1998; Chang et al., 1999, 2001, 2002, 2003; Chang et al., 2005; Sun et al., 2005; Chang and Lo, 2007). Recombinant proteins (1.2 μg/30 μL/cover slip) were coated on cover slips at 37°C for 1 h in 6-well microtiter plates, and then blocked with 5% bovine serum albumin (BSA) at 37°C for a further hour. After being washed twice with PBS, cells (1 × 10⁶) pre-mixed with antibodies (75 μg) or synthetic peptides (10 nmol) at 37°C
for 30 min were seeded onto pre-coated cover-slips and grown in medium at 37 °C for 45 min. After being washed twice with PBS, cells were fixed with 4% paraformaldehyde (in 1× PBS) at room temperature for 30 min. After being washed with PBS again, the cells were examined using an inverted microscope (Axiovert 40 CFL, Carl Zeiss, Göttingen, Germany) and photographed.

3. Results

3.1. Construction of expression plasmids for the SARS-CoV spike protein

The cell adhesion assay required recombinant proteins of the S1 domain and corresponding antibodies. Previous studies have reported that prokaryotic expressed non-glycosylated recombinant S proteins retained their function (Zhou et al., 2004; Chakraborti et al., 2005; Zhao et al., 2005). In this study, different fragments of the S1 domain (S1-1, S1-2, and S1-3) were constructed into E. coli expression vectors and expressed as glutathione S-transferase (GST)-fusion proteins (Fig. 1). S1-1 and S1-2 recombinant proteins, encompassing amino acids 1–446 and 447–641 of the S protein, were used to immunize rabbits for 5 immunization-cycles to obtain corresponding antibodies.

3.2. Determination of potential binding site of SARS-CoV spike proteins by using a cell adhesion assay

Previous studies have shown that SARS-CoV can infect the kidney cells of the African green monkey (Vero E6) (Li et al., 2003; Marra et al., 2003). Purified GST, S1-3, and S1-2 recombinant protein-coated cover slips and Vero E6 cells were used to perform a cell adhesion assay. The cell-adhesion efficiency of Vero-E6 cells to S1-3 coated cover slips were approximately 1.5 to 1.7-fold higher than that on GST-coated controls (Figs. 2A-a-b, B, 4A-a-b, and B). These results indicate that E. coli-derived S1-3 contains functional RBD for Vero E6 cells. In contrast, the efficiency of cell adhesion on S1-2 coated substrates is less than that of substrates coated with the control protein GST (Fig. 2A-c and B). The cell adhesion of Vero E6 on S1-3 coated substrates could be inhibited by rabbit polyclonal antibodies against S1-1 and S1-2, but cannot be inhibited by control antibodies against GST (Fig. 2A-d-f and B). These results indicated that the RBD potentially locates around the S1-1 and S1-2 junction region.

3.3. The permissive cell line Vero E6 but not the non-permissive cell lines HeLa and NIH 3T3 adhere to SARS-S coated surfaces

To verify whether the S protein-mediated cell adhesion is specific to SARS-CoV permissive cells (e.g., Vero E6 cells), 2 SARS-CoV nonpermissive cell lines, HeLa and NIH3T3 [which cannot be infected by SARS-CoV (Wang et al., 2004)], were used as controls to perform the cell adhesion assay. The cell-adhesion efficiency of NIH3T3 or HeLa cells to GST and S1-3 coated substrates did not show significant differences (Fig. 3). This suggests that...
study might not provide high quantitative and sensitive data that matches the flow cytometry (Babcock et al., 2004; He et al., 2004; Sui et al., 2004; Wong et al., 2004; Zhou et al., 2004) and ELISA-based analyses (Xiao et al., 2003; He et al., 2004; Ho et al., 2006), the analysis only requires recombinant proteins, peptides, virus permissive cells, and an ordinary light microscope to accomplish. Consequently, expensive equipment such as the flow cytometer, fluorescence microscope, and ELISA/fluorescence microplate readers are unnecessary, and this assay can be easily conducted in general laboratories, including those in third world countries. Because cell adhesion can be easily measured using MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide, a yellow tetrazole] assays, this method has the potential of being developed as a high throughput screening system for small molecule inhibitors when combined with automatic and robotic systems, which are unsuitable for flow cytometry and the immunofluorescence microscope-based assay. ELISA is also available for high throughput or fast screening systems; however, the recombinant ACE2 protein may not completely present as the native form of ACE2 on the surfaces of SARS-CoV permissive cells. ELISA-based analyses are only suitable to conduct after identifying the cellular receptor, for example after identifying ACE2; however, this study demonstrated that the novel cell-adhesion approach can be conducted even before identifying the cellular receptor.

For unknown reasons, the cell adhesion of Vero E6 cells to SARS-S (S1-2) substrates, which do not contain the RBD, with a lower efficiency than that of control protein GST (Fig. 2). The functional comparisons between different but related cell-adhesion proteins are difficult. For example, previously we found that even a single alanine insertion in various places around the cell-binding motif RGD (R: arginine; G: glycine; D: aspartic acid) of the snake-venom-protein rhodostomin could result a dramatically difference of cell-adhesion property. An alanine insertion at position 56 (between 56th and 57th amino acid of rhodostomin) could disrupt the cell-adhesion function of rhodostomin, while single insertions at positions 45–55 were not (Chang et al., 2001). Because the remaining protein backbone is unchanged in this case, these results suggest that the protein structure of rhodostomin is the key determinant of cell-adhesion function. Even though, the protein backbone may play an important role in other conditions. For example, evidences indicate that positive charge poly-amino acids such as poly-lysine can serve as a cell adhesion component (Mazia et al., 1975; Rajnicek et al., 1998; Brynda et al., 2005). In addition, net positive charge of cell-attached substratum seems to play a positive role on lamellipodia, filopodia formation, cell-adhesion and cell-growth (Rajnicek et al., 1998; Brynda et al., 2005). As a result, after analyzing the amino acid sequences of recombinant GST-S1-2 and GST proteins, which encoded by S1-2/pGEX-6P-1 and pGEX-2KS, respectively, we found that pGEX-2KS-encoded GST contains more positive-charge amino acids than S1-2/pGEX-6P-1-encoded GST-S1-2 (GST: 13%; GST-S1-2: 10%). These phenomena, which include differences on the protein structures and positive changes, may explain the reason why cell adhesion to GST-substrates was more efficient than that of GST-S1-2-substrates. Nonetheless, further functional and structural investigations are needed to address the possibilities. Despite of this, the assay in our condition has a great resolution power (with high reproducibility and significantly) to distinguish the levels of cell adhesion when RBD-containing proteins were used. This suggests that the analysis system has great potentials on the analysis of RBD of viral proteins.

Using this method, our data suggest that peptides spanning residues 436–445 and 437–461 of the S protein are the minimal size of peptides to block the binding of the SARS-CoV S protein to Vero-E6 cells. These peptides are located within the hundred-amino-acids RBDs previously identified using other methods (Xiao et al., 2003; Babcock et al., 2004; He et al., 2004; Sui et al., 2004; Wong et al., 2004; Zhou et al., 2004).
Wong et al., 2004) and are within the RBM (residues 424–494) characterized by the crystal structure of the SARS-CoV S protein (residues 306–527) and the ACE2 complex (Li et al., 2005). These peptides also overlap with a hexapeptide (residues 438–443), which blocks viral entry into host cells through ACE2 (Struck et al., 2012). These peptides cover most parts of the putative high antigenicity site (residues 426–456) reported by the Hsiang CY group (Ho et al., 2004). These peptides are useful as peptide inhibitors and are optimal candidates for developing the SARS peptide vaccine.

Acknowledgments

This work was supported by grants from Tzu-Chi University (TCMRC9220). The authors are grateful to Professor Chen PJ (National Taiwan University, Hepatitis Research Center) for generously providing SARS-CoV spike-gene containing plasmids TW1-#18, TW1-#19, and TW1-#20. We also greatly appreciate Genesis Biotech (Taipei, Taiwan) for kindly providing all synthetic peptides. The authors also want to thank Professor Wang MH and his team (Experimental Animal Center, Tzu-Chi University) for their help in maintaining the experimental animals and the pathogen-free environments. We deeply appreciate Professor Chen JH (Department of Molecular Biology and Human Genetics, Tzu-Chi University), Professor Pang CY (Institute of Medical Sciences, Tzu-Chi University), and Professor Lo SY (Department of Laboratory Medicine and Technology, Tzu-Chi University) for kindly providing Hela, NIH3T3, and the Vero E6 cell lines, respectively.

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