Selection and Characterization of a Single-Chain Variable Fragment against Porcine Circovirus Type 2 Capsid and Impedimetric Immunosensor Development

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ABSTRACT: Porcine circovirus type 2 (PCV2) is the primary causative agent of porcine circovirus-associated disease (PCVAD) that causes huge global economic losses for the swine industry. Effective strategies or rapid detection of PCV2 in pig are essential to control PCVAD. Here, single-chain variable fragments (scFvs) were selected and characterized against the PCV2 capsid using phage display technology. Phage scFv clones were selected from the human scFv phagemid library (Tomlinson I + J) for direct panning against the PCV2 capsid. Eighty-four monoclonal phage scFvs were individually tested for binding to the PCV2 capsid by ELISA. Eight scFv clones showed significant binding to the PCV2 capsid and only three clones (clone nos. 13, 37, and 81) contained both VHCDRs and VLCDRs in the sequence. Clone scFv no. 81 had the highest reactivity to the PCV2 capsid and was constructed in the pET22b (+) expression vector. The recombinant was transformed to Escherichia coli BL21(DE3) for expression and purification. The scFv showed appropriate affinity to the PCV2 capsid by western blot analysis. Kinetics of scFv and the PCV2 capsid were determined using surface plasmon resonance and showed binding affinity in the nanomolar range (K_D = 57.2 nM). Our scFv was first applied in the development of an impedimetric immunosensor for PCV2 capsid detection, and results showed that impedance increased with increasing PCV2 capsid expression with limit of detection = 114 nM. Findings demonstrated that our scFv has potential for use as a receptor for biosensor devices.

1. INTRODUCTION

Porcine circovirus type 2 (PCV2) is a small non-enveloped virus, with a single-stranded circular DNA genome approximately 1.7 kb in length as a member of the genus Circovirus, family Circoviridae.1 PCV2 causes circovirus disease and endemic infection of swine. Diseases associated with PCV2 have a significant impact on swine populations worldwide.2–5 Epidemiologic research on endemic infection has shown the rapid global spread of virulent PCV2.6–8 PCV2 is the primary causative agent of porcine circovirus-associated disease (PCVAD) composed of post-weaning multisystemic wasting syndrome (PMWS),9 porcine dermatitis and nephropathy syndrome (PDNS), porcine respiratory disease complex (PRDC), and reproductive failure.10 PMWS-affected piglets between 8 and 16 weeks old exhibit symptoms including jaundice, weight loss, difficulty in breathing, enlarged lymph nodes, and diarrhea.11 Multiple PCV2 vaccines are available for disease prevention that generate an immune response against the viral capsid.12–14 However, antibody levels of piglets at the time of vaccination may be low or non-existent, and a protocol is required to determine the optimal time for early inoculation.
The PCV genome has two major open reading frames (ORFs). ORF1 is essential for viral replication, while ORF2 encodes the major capsid as the protein structural shell of the virus (21). The ORF2 protein contains type-specific epitopes (22). PCV2 has a supramolecular structure formed by twelve pentamers of a single protein called Cap. The primary structure of this protein consists of 233−234 amino acid residues, with a molecular mass of approximately 28 kDa.15−17 The PCV2 capsid is involved in the host protective response system and is a target for vaccine and diagnostic reagent development.

Diagnostic testing of the PCV2 capsid is useful for monitoring infection and helps swine farmers to assess whether there are subpopulations and implement strategies to homogenize the immune status of the breeding herd. Many tools are available for detection of PCV2 including quantitative polymerase chain reaction (PCR) that is widely used to detect PCV2 in serum samples,18 immunoperoxidase monolayer assay (IPMA), western blotting, enzyme-linked immunosorbent assay (ELISA),9,20 and surface plasmon resonance (SPR).21 These methods require antibodies specific to the PCV2 capsid protein. There are polyclonal antibodies which sometimes show nonspecific reactivity, while monoclonal antibodies provide highly specific reactions. Thus, selection of high-quality antibodies is an important issue for increasing the specificity and sensitivity in immunoassay methods.

Phage display is a selection technique whereby a peptide or protein is fused with a bacteriophage coat protein and displayed on the surface of a virion22 as a popular approach for high-throughput screening of protein interactions. This technique is used to produce the specific antibody fragments such as single-domain variable heavy chain nanobodies (VHHs)31 and single-chain variable fragment (scFv). scFv is the most commonly used type of the recombiant antibody. It composes of VH and VL domains that function in antigen−binding activities.23 In a previous report, scFv has been used for detection of various targets such as Bacillus thuringiensis Cry1F toxin,24 Heliobacter pylori VacA toxin,25 tumor necrosis factor−receptor; 4-1BB,26 and Chickpea chlorotic dwarf virus.27 However, to the best of our knowledge, phage display scFv for the PCV2 capsid has never been reported.

Electrochemical biosensors are now an essential component of medical diagnostics.28 Electrochemical impedance spectroscopy (EIS) is a sensitive electrochemical technique used for detecting biomolecular interactions occurring at the electrode surface. Impedance changes occur from incremental deposits on the working electrode surface such as bioreceptor and target−bioreceptor complexes.29 The EIS technique has shown promise in quantifying molecules using antibodies as recognition molecules on electrodes or cell impedimetric immunosensors including bacteria, viruses, parasites, and inflammatory markers.30−34 EIS offers a powerful, informative, and fast electrochemical response, with non-destruction of the target, simple operation, and low cost. Thus, this technique has gained widespread application.

This study selected scFvs against the PCV2 capsid from the human scFv phagemid library (Tomlinson I + J) using phage display technology. The scFv reactivity was characterized by ELISA, western blot, and SPR analysis to develop scFv impedimetric immunosensors based on EIS for PCV2 capsid detection.

2. MATERIALS AND METHODS

2.1. scFv Phage Library Biopanning. One hundred microliters of 5 μg/mL PCV2 capsid protein (Sino Biological Inc.) were coated onto a 96-well plate and incubated at 4 °C overnight. Blocking was performed with 1% BSA in 10 mM PBS and incubated at room temperature for 2 h. Then, 100 μL of phage (1 × 1012) from human scFv phagemid library (Tomlinson I + J) was added into the wells and incubated for 1 h. The plate was washed with 0.1% Tween-20 in PBS to remove unbound phage 10 times. The bound phages were eluted by adding 100 μL of trypsin (1 mg/mL) and incubated for 15 min on a shaker. The bound phage was filtrated using a 0.22 μm filter membrane and infected with Escherichia coli XL1 Blue. The bacteria were cultured in the 2xTY agar plate (1.6% tryptone, 1% g yeast extract, and 0.5% NaCl) with 10 μg/mL tetracycline and 1% glucose, incubated at 37 °C for 30 min, and then plated in 2xTY (containing 100 μg/mL ampicillin) at 37 °C for 16 h. The polyclones from the first round of selection were amplified for further rounds (Figure S1).

Monoclonals found after the first round of selection were amplified for ELISA screening as follows. Individual phage-infected E. coli XL1 Blue colonies were grown in a 96-well plate containing 100 μL of 2xTY (containing 100 μg/mL ampicillin, 0.1% glucose) and shaken at 37 °C until the OD600 reached 0.4. The KM703 helper phage was infected into the cell and incubated at 37 °C for 30 min. The pellet was collected by centrifugation at 3000 rpm for 30 min, plated into 2xTY (containing 100 μg/mL ampicillin, 50 μg kanamycin and 0.1% glucose), and incubated at 30 °C overnight with shaking. The next day, purification was performed by centrifugation of the phage suspension at 4 °C, 3000 rpm for 2 h. The supernatant was collected and precipitated by adding 4% polyethylene glycol in 25 mM NaCl and incubated at 4 °C for 1 h. The precipitated phage was further centrifuged at 3000 rpm for 30 min. The phage pellet was collected, resuspended with PBS, and maintained at 4 °C until required for the next experiment.

2.2. ELISA Screening for Phage scFv Specific to PCV2 Capsid. A 96-well plate was coated with 100 μL PCV2 capsid (5 μg/mL) and incubated overnight at 4 °C. The next day, 200 μL of washing buffer (0.05% Tween-20 in 1x PBS pH 7.4) was added to remove excess protein and washed three times. Then, 200 μL of blocking buffer (1% BSA in 1x PBS pH 7.4) was further added for 2 h. Each phage scFv was added to the blocked well and incubated for 1 h. Then, 100 μL of HRP-conjugated anti-M13 antibody (1:2000 dilution) was added and the sample was incubated at room temperature for 1 h. To indicate the reactivity of individual phages of scFv binding to PCV2, 100 μL TMB with 1% of H2O2 as the substrate for HRP was added in each well and incubated in a dark area to develop the blue color. Then, 100 μL of 1 M of HCL was added to stop the peroxidation reaction. The blue color turned to yellow and absorbance at wavelength 450 nm was measured using an ELISA plate reader. All incubation steps were performed at room temperature, except for protein coating on the first day. Positive monoclonal bacteria were isolated for sequencing. The phage-scFv81, one of the potential phages, was diluted to concentration 2.6 × 10−4 to 5.0 × 108 cfu/mL and subjected to ELISA testing for PCV2 capsid compared with BSA as the negative control.

2.3. Construction of scFv Expression Vectors. The scFv/pIT2 phagemids were extracted, and scFv was amplified using a Kapa HiFi Kit with LMB3 as forward primer.
(S’CAGGAA ACA GCT ATG AC 3’) and pHEN as the reverse primer (S’CTA TGC GGC CCC ATT CA 3’). The amplified scFv products were confirmed for size by 2% agarose gel. To prepare the inserted gene and vector, the scFv gene and pET22b (+) vector were cut using restriction enzyme Ncol and NotI and then confirmed for size by 2% agarose gel. The scFv and pET22b (+) vector contained in the gel were further purified by a purification kit, and concentration was measured by a nanodrop spectrophotometer. The ligation reaction of recombinant scFv/pET22b+ was performed with ratio (µL) of insert gene/pET22b (+)/T4 ligase/buffer A/buffer B (0.3:6.7:1:1:1), incubated at 20 °C for 30 min. The recombinant gene was further transformed into the DH5α competent cell. Individuals of inserted scFv in pET22b (+) clone were confirmed by colony PCR with T7 as forward and reverse primer and confirmed for product size by 2% agarose gel. The clones with scFv genes were further grown in the 2xTY agar plate, and the scFv/pET22b (+) plasmid was extracted by sequencing confirmation.

2.4. scFv Expression and Purification. The recombinant scFv with 6xHis tag was expressed in E. coli BL21(DE3). The strains containing the expression plasmid were propagated overnight at 37 °C in 3 mL Luria—Bertani (LB) (seed) medium containing 100 µg/mL of ampicillin. Then, 1% of seed was added to 500 mL LB broth containing 100 µg/mL of ampicillin and incubated at 37 °C for around 4 h to reach OD600 = 1.0. Then, isopropyl β-d-thiogalactopyranoside (IPTG) was added to a final concentration of 0.4 mM and incubated overnight for 20 h at 16 °C. Next day, the cells were collected by centrifugation at 4 °C, 3000 rpm for 15 min. Cell pellets were then resuspended in 20 mL of 1X PBS buffer pH 7.4 and further sonicated for 30 min on ice to disrupt the cells. The cell homogenate was centrifuged at 4 °C, 10,000 g for 30 min. Then, the supernatant was collected and purified using cation exchange, followed by HisPurTM Ni-NTA Resin. scFv fragments were analyzed for size by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) under reduced condition, with bands observed after staining with Coomassie Blue.

2.5. Western Blot Analysis. To confirm the size, scFv in SDS-PAGE gel was transferred into the polyvinylidene fluoride (PVDF) membrane for 30 min and blocked with TSW buffer pH 7.4 (10 mM Tris, 0.9% NaCl, 0.25% gelatin, 0.1% Triton x–100, 0.02% SDS) for 1 h. The scFv fragments were detected with 3000X diluted anti His antibody for 1 h, then washed and further incubated in 3000X diluted anti mouse IgG HRP for 1 h, and washed to remove the unreacted antibody. HRP was visualized with enhanced chemiluminescence (ECL), a luminol-based substrate for HRP.

To prove scFv binding to the PCV2 capsid, equal volumes of the PCV2 capsid (150 ng/well) were electrophoresed onto SDS-PAGE, with 4 wells of PCV2 capsid loaded for different antibody detection and transferred into the PVDF membrane. The PCV2 capsid antigen in the membrane was then detected by antibody mentioned as follows: (1) used 1° antibody: purified scFv (0.05 mg/mL), 2° antibody: anti His antibody (mouse monoclonal), 3° antibody: anti mouse IgG-HRP, (2) used 1° antibody: phage scFv (2.6 × 10⁸ cfu/mL), 2° antibody: anti M13 HRP, and (3) used 1° antibody: commercial anti PCV2 (rabbit polyclonal), 2° antibody anti rabbit IgG–HRP.

2.6. Interaction Analysis. The SWISS-MODEL (https://swissmodel.expasy.org/) was used to obtain the 3D structure of scFv, with the 3D structure of the PCV2 capsid obtained from the protein databank (PDB ID: 3JCI). The quality of the 3D scFv model was checked by PROCHECK. Identification of the potential binding interface of scFv to the PCV2 capsid and protein–protein docking was performed by the program Frodock version 2.0. Molecular interaction between scFv and the PCV2 capsid was visualized by PyMOL software version 1.3.

2.7. Kinetics and Affinity Determination by SPR. Binding kinetics of scFv to the PCV2 capsid was measured by SPR (OpenSPR, Nicolea Lifesciences) at 25 °C. The ligand was activated for COOH groups by mixing 100 µL of 0.030 mg/mL scFv in 10 mM PBS pH 7.4 (running buffer), 50 µL of 0.2 M EDC, and 50 µL of 0.05 M NHS, incubated at 4 °C for 1 h. All measurements were performed in running buffer. The NH2 sensor chip surface (Nicoya Lifesciences) was loaded into an OpenSPR instrument and performed for baseline. To immobilize scFv on NH2, 200 µL of coupled scFv/EDC/NHS was injected at a flow rate of 10 µL/min and dissociated for 5 min. Then, 200 µL of blocking was injected to inactive non-immobilized scFv. Various concentrations of the PCV2 capsid were performed (5.52–176 nM) and injected at 200 µL with a flow rate of 10 µL/min and dissociated for 2 min. Kinetic data were fitted and analyzed by Trace Drawer version 1.6.1 using the 1:1 binding model, and respective rate constant values were calculated. All measurements were performed in triplicate.

2.8. Impedimetric Immunosensor. EIS was detected using a PalmSens4 Potentiostat (Zahner, Germany). The modified process was adapted from Cordeiro and Chinna-dayala. bare gold electrodes (Metrohm) were initially cleaned with DI water, pre-treated by immersion in 0.1 M NaCl for 5 min, and measured for impedance as a reference electrode. Then, 5 µL of 20 mM disodium phosphate (DSP) in 99% DMSO was dropped onto the working electrode for 2 h and rinsed in DI water to remove unbound DSP. The modified electrode was further immobilized with 5 µL of 0.121 mg/mL scFv for 1 h at 4 °C and washed to remove unreacted scFv. A schematic diagram of immunosensor fabrication is shown in Figure 7A. Then, different concentrations of the PCV2 capsid 35–704 nM in 10 mM PBS (pH 7.4) were dropped onto the modified electrode for 30 min at room temperature and washed to remove the unbound PCV2 capsid. All measurements were performed by dropping 100 µL of the 5 mM [Fe(CN)₆]₃⁻/₄⁻ redox probe covering three electrodes (working, reference and counter) as the background electrolyte. The impedance was scanned by inserting the modified electrode into a potentiostat, and the result was displayed by program PSTrace version 5.7 with parameters set as T equilibration:3 s, Edc:0.08961 V, Eac:0.005 V, N frequency:73:12/dec, Max impedance was scanned by inserting the modified electrode into a potentiostat, and the result was displayed by program PSTrace version 5.7 with parameters set as T equilibration:3 s, Edc:0.08961 V, Eac:0.005 V, N frequency:73:12/dec, Max

3. RESULTS

3.1. Screening of scFvs by Biopanning. Monoclones of the scFv phage-bounded PCV2 capsid were found on the 2xTY agar plate after the first round of biopanning that was performed in triplicate, with approximately 20 clones on each plate. Eighty-four individual colonies from the three plates were collected for propagation by helper phage KM703.
The ELISA was used for screening positive clones using an HRP-conjugated anti-M13 antibody since the scFv proteins which were displayed on the phages also have M13-tag on the C-terminal of the sequences. Binding of selected scFv phages with the PCV2 capsid was quantitatively confirmed using an HRP-conjugated anti-M13 antibody by measuring the absorbance at 450 nm. Results showed significant binding of the scFv phage to PCV2 capsid, especially phage nos. 13, 22, 24, 32, 37, 81, 83, and 84 (Figure 1A). The eight positive clones were then collected for plasmid extraction sequencing using agarose gel, as shown in Figure 1B. Clone no. 81 showed the highest binding to the PCV2 capsid (OD450 = 1.115) and less interaction with BSA. Various concentrations of phage scFv81 from $2.6 \times 10^{-4}$ to $5.0 \times 10^6$ cfu/mL against the fixed concentration of the PCV2 capsid (100 ng/well) were measured by ELISA. Results showed that binding activity of phage scFv to PCV2 capsid was significantly higher than phage scFv bound to BSA control (Figure 1C). Furthermore, increasing phage scFv concentration increased the binding complex of the scFv/PCV2 capsid, as shown by the increase in OD450. Thus, concentration of $2.6 \times 10^5$ cfu/mL phage scFv was selected for further investigation.

3.2. Sequencing and Complementarity-Determining Region Analysis. Sequencing the eight positive clones with specific primers found only three genes: scFv13, scFv37 and scFv81 that contained both VHCDR and VLCDR regions (Figure 2A). However, only scFv81 with the highest binding affinity was chosen for further study. The full-length sequence of scFv81 consisted of 248 amino acid residues with a predicted molecular weight of 27.2 kDa, including a flexible amino acid linker of Gly-Ser between the VH and VL regions. The homology identities and coverages for VH and VL were analyzed through the international ImMunoGeneTics information system (IMGT) (www.imgt.org), as shown in Table 1. VH was the closest to human V region frameworks by accession no. Z12367 Homsap IGHV4-38-2*01F with amino acid homology covering 100%, calculated as a percentage of the identical residues for their FR1-3 sequences (a total of 120 residues). VL is closest to the human V region by accession no. X59315 Homsap IGKV1-39*01F with amino acid homology covering 92.03% for FR1 and 100% for FR2-3 (Table 1). The predicted CDR1, CDR2, and CDR3 of the VH domain were located at positions 27–35, 53–59, and 98–111 in the primary sequence, while predictions of CDR1, CDR2, and CDR3 of the VL domain were specified at positions 164–169, 187–189, and 226–233, respectively, as depicted in Figure 2B. The 3D structure of scFv was modeled by SWISS-MODEL (Figure 2C). The Ramachandran plot of the 3D model that indicated a
good quality of structure showed 59.5% of residues in most favored regions, 34.5% of residues in additional allowed regions, 5.0% of residues in generously allowed regions, and only 1.0% of residues in disallowed regions (Figure S2).

The interaction model was predicted by Frodock version 2.0. Results showed a possible binding interface between CDR3 of VH and VL of scFv with the surface of the PCV2 capsid protein (Figure 2D). This binding used the hydrophobic interaction between residues VAL103, TYR104 and VAL39, TYR169, 229, and TRY119 (Figure S3A). Hydrogen bonds were found between residues TYR54 and ARG173 (OH-NH2), SER230 and SER117 (OG-OG), and TYR169 and TYR119 (OH-OH) (Figure S3B) with ionic interaction between residues LYS66 and ASP169, ASP108, and ARG7 (Figure S4A), aromatic–aromatic interactions between residues TYR104 and PHE4, TYR169 and TYR119 (Figure S4B), and cation–Pi interaction between residues TYR169 and ARG7, TYR229 and LYS61, and TYR54 and ARG173 (Figure S5). The scFv was found to use multiple residues to form interface contact with the PCV2 capsid, especially residue TYR54 from VHCDR1, residues VAL103, TYR104, and ASP108 from VHCDR3, residue TYR169 from VLCDR1, and residues TYR229, SER230, from VLCDR3 (Figure 2B).

Table 1. Percent Amino Acid Homology of the ScFv Sequence with the Closest Human V Region Frameworks

| closest human V region | amino acid homology with human FRs |
|------------------------|-----------------------------------|
|                       | FR1 (%) | FR2 (%) | FR3 (%) |
| VH | Z12367 Homsap IGHV4-38*2*01 F | 100.0    | 100.0   | 100.0   |
| VL | X59315 Homsap IGKV1-39*01 F   | 92.03    | 100.0   | 100.0   |

Asterisk followed by two number indicates the allele polymorphism.

3.3. Construction of scFv 81 in pET22b (+) Expression Vector. The scFv81 was constructed according to Figure 3A–C. scFv genes from pIT2 phagemid were amplified by PCR using LMB3 as the forward primer and pHEN as the reverse primer (Figure 3A). The scFv genes performed double digestion by NcoI and NotI, with the size of scFv products before and after digestion confirmed by 2% agarose gel as 926 and 731 bp, respectively (Figure 3D). The pET22b (+) expression vector was performed by restriction enzyme, similar to the scFv gene. Sizes of pieces of pET22b (+) before and after digestion were 5947 and 5439 bp, respectively (Figure 3E). The purified scFv fragments were then inserted into the open reading frame of the pET22b (+) vector to generate the recombinant plasmid scFv/pET22b (+), with an expected size of 6177 bp (Figure 3C) under regulation of the T7 promoter using NcoI (bp 904–907) and NotI (bp 166–169) restriction sites. At the C-terminal, the constructs were fused to a hexahistidine-tag. The recombinant scFv/pET22b (+) was then transformed into E. coli DH5α competent cells, and the insertion was confirmed by colony PCR with the T7 primer, designed for up-stream amplification of the VH and down-stream amplification of the VL genes. The corrected size of the scFv gene was found to be 973 bp, while no scFv band was found in pET22b (+) without scFv insertion (Figure 3F). This result indicated that our positive transformants had an scFv antibody gene inserted into the pET22b (+) expression. Positive colonies showing the presence of scFv gave further plasmid extraction and transformed into E. coli BL21 competent for further expression.

3.4. Expression and Purification of Soluble scFv. scFv/pET22b (+) was expressed in E. coli BL21, and 0.4 mM IPTG for 20 h at 16 °C induced soluble scFv. After pelleting the cells and sonication, soluble scFvs containing His-tags were purified from the cell lysate by HisPurTM Ni-NTA resin. After washing with PBS, the His-tagged scFvs were eluted with increasing
concentration gradient of imidazole (50–400 mM) and the size of scFv fragments was evaluated by SDS-PAGE with Coomassie Blue staining (Figure 4). Results showed high insoluble scFv in cell lysate compared to soluble scFv in the supernatant (with a band at 27.2 kDa as expected) (Figure 4A). The scFv was confirmed by western blot analysis with anti-His antibody and presented the band as expected (Figure 4A). However, the scFv was contaminated by other proteins when using only the Ni\(^{2+}\) column. Thus, to remove the remaining contaminants, two columns of cation exchange followed by Ni\(^{2+}\)NTA were performed. Single bands of scFv proteins with high purity appeared in imidazole concentration at 100 mM (Figure 4B). Fractions of scFv were further dialyzed into PBS, while scFv concentration was determined at 0.121 mg/mL by a nanodrop spectrophotometer at 280 nm.

### 3.5. Western Blot Analysis

To confirm the binding of scFv, the PCV2 capsid was loaded in equal protein (150 ng/well) in 4 wells of SDS-PAGE; one for Coomassie Blue staining as positive control protein and three for western blotting and probed with different antibodies of purified scFv, phage scFv, and commercial anti PCV2 antibody. Results showed that SDS-PAGE contained the PCV2 capsid with a molecular weight of 23.8 kDa (Figure 5A). The purified scFv and phage scFv had specific binding to the PCV2 capsid,
presenting in specific bands on western blot membranes (Figure 5C,D). This result of the probed PCV2 capsid with the anti PCV2 antibody (rabbit polyclonal) showed the specific band and one tiny non-specific band on the membrane (Figure 5B).

3.6. Kinetics and Affinity Determination by SPR. To evaluate the apparent affinities of purified scFv antibodies to the PCV2 capsid, binding kinetics using the immobilized scFv antibody were determined by SPR. The scFv was immobilized on an NH\textsubscript{2} sensor chip through covalent interaction with the activated carboxyl functional group of scFv, allowed by injected different concentration PCV2 capsids (5.52–176 nM). SPR binding profiles are presented in Figure 6. The scFv antibody displayed high binding affinity to the PCV2 capsid, with K\textsubscript{D} values in the nanomolar range (57.2 nM) and exhibited parameters with association rate (k\textsubscript{a}) = 8.29 × 10\textsuperscript{4} M\textsuperscript{-1} S\textsuperscript{-1} and dissociation rate (k\textsubscript{d}) = 0.0474 S\textsuperscript{-1}.

3.7. Impedimetric Immunosensor. The purified scFv was applied for the immunosensor based on EIS for detection of the PCV2 capsid. Quantitative behavior of the immunosensor was assessed by measuring the electron transfer resistance (R\textsubscript{et}) of different concentrations of the PCV2 capsid. The immunosensor was performed according to Figure 7A, which shows the modification steps of gold electrodes with DSP, scFv antibody, and PCV2 capsid, respectively. The impedance spectra were characterized by modified processes on the electrode, with results presented as Nyquist plots in Figure 7B using bare gold electrodes (curve a, R\textsubscript{et} = 42 Ω). The gold electrodes were further attached by DSP (curve b, R\textsubscript{et} = 138 Ω), scFv immobilized (curve c, R\textsubscript{et} = 167 Ω), and the R\textsubscript{et} value increased. The scFv/DSP/gold electrodes were further incubated with PCV2 capsid, and the R\textsubscript{et} value increased significantly (curve c, R\textsubscript{et} = 234 Ω). Figure 7C shows that the EIS signal increased with increasing PCV2 capsid concentration from 35, 176, 352, and 704 nM, with R\textsubscript{et} values increasing as 323, 359, 419, and 546 Ω, respectively. A plot of the ΔR\textsubscript{et} value against the concentration of the PCV2 capsid in the range 35–704 nM exhibited a linear relationship with a correlation coefficient of 0.9963 and LOD = 114 nM (Figure 7D). These experimental results confirmed the feasibility of the
proposed immunosensor using scFv as a probe for the diagnosis of the PCV2 capsid.

**4. DISCUSSION**

PCV2 infection is associated with many diseases. Globally, PCVADs have a serious economic impact on the swine industry.3,5 Thus, effective strategies for rapid detection of PCV2 will help veterinarians and farmers to generate information that can be used to assess clinical disease and make epidemiological decisions to evaluate farm stability and establish vaccines and biosecurity programs.

Phage display screening has previously been used for selecting the specific binder to the PCV2 capsid, but different libraries exist such as peptide libraries38−40 and Bactrian camel sdAb library.41−43 This study used human scFv library. The single-domain antibody (sdAb) has smaller fragments with size 15 kDa44 consisting of either VH or VL domains, while scFv consists of both VH and VL with a size of 28 kDa. In terms of phage display, the small antibody format (sdAb, scFv) is preferred because of the ease for packaging the new phage particles and expression compared to full immunoglobulin, which is large sized and not suitable for *E. coli.*45 Tomlinson I + J libraries are widely used, based on scFv format with diversity of 18 amino acids incorporated in CDRs 2 and 3 in positions involved in antigen binding, and are highly diverse in the natural repertoire. Many studies used this library, and their antibody showed high specificity and binding affinity to the targets.46−48

Selection of scFv against the PCV2 capsid by phage display screening was successful in the first round of panning. The bounded phage infected *E. coli* XL1-Blue, and subsequently put on 2XY agar plate allowed to the monoclonal scFv. Screening of positive clones from libraries generally require at least three to seven rounds of biopanning to select specifically binding clones.49,50 However, we performed panning in triplicate and found approximately 20 individual clones on each plate.

In sequencing and complementary-determining region (CDR) analysis, CDRs form part of the variable chains in immunoglobulins (antibodies) that bind to their specific antigens. Variable regions of a polypeptide chain are composed of CDR1, CDR2, and CDR3 as the most variable.51 Our results showed different amino acids in CDRs of the three clones, and this may contribute to the clones binding to different epitopes on the PCV2 capsid, as shown in the ELISA result (Figure 1A). Clone scFv 81 gave the highest reactivity and was constructed into the pET22b (+) expression vector.

Our recombinant scFv/pET22b (+) was first transformed into the *E. coli* DH5α competent cell and subsequently into *E. coli* BL21 because *E. coli* DH5α strain growth is faster than *E. coli* BL21. *E. coli* DH5α has recA mutation; it does not show heterologous recombination, and this ensures higher insert stability.52

The abilities of the scFv binding PCV2 capsid were evaluated by Western blotting (Figure 5). Purified scFv and phage scFv specific capturing to PCV2 capsid by size 23.8 kDa were displayed on the PVDF membrane. These results were similar to using the anti PCV2 capsid antibody as the control. This suggested that the purified scFv bound specifically to the PCV2 capsid and was effective for diagnostics development. SPR analysis has been widely used to study interactions between immobilized biomolecules on sensor chip and solution-phase analytes. Our scFv showed good binding...
affinity to PCV2 in the nanomolar range (57.2 nM), probably because of the long VHCDR3 loop (Figure 2) as the main factor that determined the character of high binding affinity. Our binding affinities of scFv to PCV2 capsid were similar in the nanomolar range with Hu ($K_D = 40$ ng/mL, 952 nM) with used full length of the anti-PCV2 antibody. Yang ($K_D = 49.4$ nM) observed kinetics binding by using sdAb. Hao ($K_D = 103$ nM) observed kinetics binding using peptides against PCV2. However, no report has evaluated the binding affinity of the scFv antibody to PCV2 using SPR analysis. Our protein–protein docking analysis confirmed the strong binding of scFv with the PCV2 capsid using several interaction bonds including hydrophobic, hydrogen bond, and aromatic and cation–π interaction.

Previously, scFvs have been used in electrochemical immunosensors based on EIS for detection of many targets such as prostate specific antigen and toxic microalga *Alexandrium minutum* accepted PCV2 capsid. Here, scFv was applied on gold electrodes as an immunosensor to detect the PCV2 capsid by measuring electrode transfer resistance changes. EIS fabrication is simple with little time required to prepare scFv-attached electrodes. DSP, a protein crosslinker, was first attached on the gold electrode via a disulfide bond and then reacted with α-amine at the N-terminal of the scFv antibody. The EIS spectra, presented as Nyquist plots, showed increasing impedance when components changed in each step of modification (Figure 7B) because transfer resistance at the electrode surface gradually increased. Thus, in the measuring step, different concentrations of the PCV2 capsid were incubated with the scFv-modified electrode, corresponding to increase in electron transfer resistance (Figure 7C). This result suggested scFv binding to the target with LOD = 114 nM (2.7 μg/mL). There is no report that evaluated the sensitivity of the scFv antibody binding to PCV2 by the impedimetric immunosensor. Our study was compared to Luo’s study. They developed a label-free, antibody-based PCV2 biosensor; Staphylococcus protein A (SPA) was employed to modify a silanized fiber surface to enhance the bioactivity of a specific anti-PCV2 antibody, resulting in LOD = 4.06 ng/mL (96 nM); “our LOD value is higher than this immunoassay.

Our study demonstrated that scFv constructs provided improved binding affinity but also functioned as tracers of target antigens and signal transduction molecules in EIS immunosensor systems for PCV2 detection, with potential for diagnostic application of biosensors in the swine industry. However, the further study about the cross-reactive test, for example, test scFv activity with porcine reproductive and respiratory syndrome virus, pseudorabies virus, and also PRDC, porcine respiratory disease complex; PDNS, porcine dermatitis and nephropathy syndrome; PMWS, post-weaning multisystemic wasting syndrome; PI3, porcine immunodeficiency syndrome; PRRS, porcine respiratory and reproductive syndrome; ECL, enhanced chemiluminescence; CDR, complementarity-determining region; CFU, colony forming unit; EIS, electrochemical impedance spectroscopy; ELISA, enzyme-linked immunosorbent assay; IPMA, immunoperoxidase monolayer assay; IPTG, isopropyl β-D-thiogalactopyranoside; $K_D$, equilibrium dissociation constant; ORFs, open reading frames; PCV2, porcine circovirus type 2; PCVAD, porcine circovirus-associated disease; PDNS, porcine dermatitis and nephropathy syndrome; PMWS, post-weaning multisystemic wasting syndrome; PRDC, porcine respiratory disease complex; $R_{et}$, electron transfer resistance; scFv, single-chain variable frag-

### 5. CONCLUSIONS

We successfully performed screening and isolation of scFv clones from Human Single Fold scFv libraries I + J against the PCV2 capsid. The recombinant scFv gene/pcET22b (+) showed potential for use in antibody production of E. coli BL21. Binding behavior of scFv to the PCV2 capsid was evaluated by ELISA, western blot, and SPR. Results of SPR assays showed affinity binding of the scFv antibody to PCV2 capsid in the nanomolar range ($K_D = 57.2$ nM). Our preliminary investigation of scFv as an immunosensor based on EIS for PCV2 capsid detection with a modified gold electrode showed LOD in nanomolar range (LOD = 114 nM). This suggested that scFvs has the potential to perform as a PCV2 real virus. However, further research is required to confirm these findings.

### ASSOCIATED CONTENT

- **Supporting Information**
  The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.1c03894.

Planning phage display cycle to screen the scFv-specific PCV2 capsid, Ramachandran plot of the scFv-specific PCV2 capsid, interaction between scFv and the PCV2 capsid via hydrophobic interaction, main chain-side chain hydrogen, ionic interaction, aromatic–aromatic interaction, and cation–π interaction (PDF)

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### Notes

The authors declare no competing financial interest.

### ACKNOWLEDGMENTS

This project was supported by Research and Researchers for Industries (RRI), the Thailand Research Fund (PHD6010080). This research was also supported by the Postdoctoral Fellowship from Kasetsart University and Kasetsart University Research and Development Institutes (KURDI (F(KU) 6.64)).

### ABBREVIATIONS

ECL, enhanced chemiluminescence; CDR, complementarity-determining region; CFU, colony forming unit; EIS, electrochemical impedance spectroscopy; ELISA, enzyme-linked immunosorbent assay; IPMA, immunoperoxidase monolayer assay; IPTG, isopropyl β-D-thiogalactopyranoside; $K_D$, equilibrium dissociation constant; ORFs, open reading frames; PCV2, porcine circovirus type 2; PCVAD, porcine circovirus-associated disease; PDNS, porcine dermatitis and nephropathy syndrome; PMWS, post-weaning multisystemic wasting syndrome; PRDC, porcine respiratory disease complex; $R_{et}$, electron transfer resistance; scFv, single-chain variable frag-
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