Porcine epidemic diarrhea (PED) is an enteric disease that causes the significant loss for the swine industry. The disease first occurred in Europe in the early 1970s and the virus was first isolated in Belgium in 1978 [1]. Subsequently, the disease was also reported in the United States [2] and several countries in Asia, including China [3], Korea [4], Taiwan [5], Japan [6], and Thailand [7]. PED is characterized by intense diarrhea, vomiting, weight loss, and dehydration. These symptoms result in the death of newborn piglets and weight loss in all pig ages, which adversely affects growth performance of growing pigs [1, 8]. PED diagnosis cannot be made purely on the basis of clinical signs and histopathological lesions. Therefore, the diagnosis to confirm the presence of porcine epidemic diarrhea virus (PEDV) or its antigens must be conducted in the laboratory.

PEDV is an enveloped virus with a single-stranded positive-sense RNA and belongs to the *Alphacoronavirus* genus in the *Coronaviridae* family (ICTV, 2012). Its genome is approximately 28 kb encoding seven viral proteins: ORF1A, ORF1B, spike (S),...
OFR3, small envelope protein (E), membrane (M), and nucleocapsid (N) [9]. Among these viral proteins, S- protein is the key protein responsible for the viral entry into the host cell. It is the envelope type I glycoprotein that can bind to the receptor on the host cell, subsequently fuse the viral membrane to the host membrane, and then release the nucleocapsid protein into the host cell [10]. Vaccination with the recombinant S-protein was shown to protect the piglets against PEDV infection [11]. Thus, S-protein is one of the targets for PEDV vaccine development.

There are several neutralizing epitopes on the S-protein [12–14]. Among these epitopes, 2C10 is one of the peptides that are recognized by neutralizing monoclonal antibody (mAb) against PEDV [15]. The recombinant 2C10 single-chain antibody, consisting of variable regions of heavy (VH) and light chains (VL), was previously expressed in *Escherichia coli* and also showed the PEDV neutralization [16]. This can be developed to use as PEDV diagnostic and prophylaxis treatment.

The goal of this study was to develop 2C10 mAb in plants. Recently, plants have been used as factories to effectively produce several recombinant proteins because of several advantages over other protein expression systems, including low manufacturing cost, high protein expression level, scalability, and lack of human and animal pathogen [17,18]. Moreover, plants contain the post-translational modification, which are often critical to the proper protein functions [19,20]. The two main systems for producing proteins in plants are stable transgenic expression and transient expression. The establishment of transgenic plants is a time-consuming process with limitation of relatively low protein expression level. For transient expression, plant viral vectors were developed to increase the protein expression level [21]. Among different viral vectors, geminiviral vector was developed to produce several proteins in plants such as antigens and antibodies [22–24].

In this study, we used geminiviral vectors to produce 2C10 mAb in *Nicotiana benthamiana* Domin. (Solanaceae) and *Lactuca sativa* L. var. *longifolia* (Asteraceae) by transient expression. The seeds of *N. benthamiana* were provided from Julian Ma’s Lab (London, UK). *N. benthamiana* is the most common host plant for transient expression because of its high biomass yield. Lettuce is another rapid-growing plant that also produces high biomass and contains lower amount of phenolics and alkaloids, compared to tobacco plants. This is the advantage for the protein purification downstream process [24]. Lettuce was obtained commercially from a local grocery store with romaine lettuce labeled on the package. We tested the 2C10 mAb expressed in *N. benthamiana* and lettuce for PEDV binding and neutralizing activity. This plant-produced mAb can be applied to use as diagnostic kit or prophylactic treatment for PEDV infection, which will be very useful for the swine industry.

**Results**

The genes optimized for plant codon usage encoding the VL and VH chains of 2C10 (Fig. 1) were amplified and fused with constant regions of IgG1 mAb [25]. The fused genes were inserted into geminiviral replicon vector (pBY030.2R) (Fig. 2) and introduced to *Agrobacterium tumefaciens* strain GV3101. The 2C10 mAb was expressed in *N. benthamiana* by co-infiltration with the mixture of *A. tumefaciens* containing pBY2C10-kappa, pBY2C10-kamma, and p19. The p19 used in this study was the gene silencing inhibitor p19 from tomato bushy stunt virus using the non-replicating expression vector. At day 5 post-infiltration, the 2C10 mAb in plant extract was detected by western blot analysis using mouse anti-human IgG antiserum, FC specific and mouse anti-kappa light chain antiserum. The result showed that 2C10 IgG was produced in *N. benthamiana*, containing both heavy chain and light chain at approximately 50 and 25 kDa in reducing condition, respectively (Fig. 3A). The assembly of 2C10 IgG was determined after protein A affinity purification by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) analysis. The 2C10 mAb purified from plant proteins were analyzed under reducing and non-reducing conditions. Under reducing condition, two bands of heavy and light chains were detected at 50 kDa and 25 kDa, respectively (Fig. 3B), while the purified protein had the band at 150 kDa (Fig. 3B), which confirms the assembly of the whole IgG molecule.

To test the effectiveness of geminiviral vector in producing the 2C10 mAb in lettuce, we investigated the transient expression of 2C10 mAb in romaine lettuce from the local store. *Agrobacterium* cultures using for lettuce were similar to the experiment above. At day 4 post-infiltration, western blot analysis of infiltrated lettuce leaves confirmed the expression of both heavy chain and light chain of 2C10 IgG at the expected size (Fig. 4A), similar to the 2C10 expressed in *N. benthamiana* (Fig. 3A). The 2C10 mAb was purified from the lettuce proteins with protein A affinity chromatography (Fig. 4B).

The PEDV binding ability of plant produced 2C10 mAb was tested by indirect ELISA (enzyme-link immunosorbent assay). Plant-produced 2C10 IgG from tobacco and lettuce was diluted 1:1000 before incubated in wells containing immobilized PEDV. Detection with HRP (horseradish peroxidase) labeled anti-human IgG antiserum yielded OD450 measurement. The results showed that 2C10 mAb produced in tobacco and lettuce could bind to PEDV approximately 1.7 and 1.6 times more than the negative control (Table 1). The 2C10 mAb produced in tobacco and lettuce showed no binding to PEDV expressing inhibitor p19 from tomato bushy stunt virus using the non-replicating expression vector. At day 5 post-infiltration, the 2C10 mAb in plant extract was detected by western blot analysis using mouse anti-human IgG antiserum, FC specific and mouse anti-kappa light chain antiserum. The result showed that 2C10 IgG was produced in *N. benthamiana*, containing both heavy chain and light chain at approximately 50 and 25 kDa in reducing condition, respectively (Fig. 3A). The assembly of 2C10 IgG was determined after protein A affinity purification by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) analysis. The 2C10 mAb purified from plant proteins were analyzed under reducing and non-reducing conditions. Under reducing condition, two bands of heavy and light chains were detected at 50 kDa and 25 kDa, respectively (Fig. 3B), while the purified protein had the band at 150 kDa (Fig. 3B), which confirms the assembly of the whole IgG molecule.

The PEDV binding ability of plant produced 2C10 mAb was tested by indirect ELISA (enzyme-link immunosorbent assay). Plant-produced 2C10 IgG from tobacco and lettuce was diluted 1:1000 before incubated in wells containing immobilized PEDV. Detection with HRP (horseradish peroxidase) labeled anti-human IgG antiserum yielded OD450 measurement. The results showed that 2C10 mAb produced in tobacco and lettuce could bind to PEDV approximately 1.7 and 1.6 times more than the negative control (Fig. 5). However, glutathione S-transferase (GST) fused with truncated S-protein of PEDV showed no binding to PEDV; the binding ratio to PEDV is approximately 1 compared to negative control. These results suggested that the plant-produced 2C10 mAb might be developed for use as a diagnostic for PEDV infection.

The 2C10 mAb showed some binding activity to PEDV (Fig. 5). To evaluate the neutralization potential of 2C10 mAb, the PEDV neutralization assay was performed using 2C10 produced from *N. benthamiana*. PEDV virus (100–300 TCID50/50 µL) was incubated with the 2C10 mAb for 1 h at 37°C. The antibody-PEDV mixture was added to Vero cell and incubated for 1 h at 37°C. The result showed that 1:32 dilution ratio of 2C10 mAb was able to neutralize PEDV infection in Vero cell (Table 1) because there is no cytopathic effect. However, after the antibody was diluted to 1:64, the cytopathic effect was shown. Thus, plant-produced 2C10 mAb retained neutralizing activity against infectious PEDV, which suggested the potential to use as PEDV prophylaxis treatment.
Discussion

PED is a major reason for the economic loss in swine industry [26, 27]. PEDV infection induces death in neonatal pigs due to watery diarrhea and dehydration. Previous study reported that there are over one million piglet deaths from PEDV infection during the outbreak in China in 2010, which created large economic losses [28]. For PEDV prevention, there are different types of PEDV vaccine commercially, including live viral vaccine and inactivated virus [29, 30]. However, these vaccines did not significantly reduce the...
morbidly rate due to diarrhea, because several factors including route of administration and strains of vaccine virus affected the poor lactogenic immunogenicity of these commercial vaccines [31]. Another potential approach to protection of neonatal piglets is passive immunity induced by effective quantities of protective antibodies obtained from sow colostrum and milk. Artificial passive immunization by oral administration of specific antibodies represents an attractive approach against gastrointestinal pathogens such as PEDV [26]. Several PEDV neutralizing antibodies were developed previously [15,16,32,33], and they might be good candidates for use as PEDV prophylactic agents. However, an effective antibody production system with low production cost is required for the animal industry.

Plant-based expression is becoming attractive for recombinant protein production. Several conventional systems, such as E. coli, yeast, mammalian cells, insect cells, etc., are available commercially. Among these production platforms, plants have several advantages over other systems, including low production cost, large scalability, lack of human and animal pathogen, and post-translational modification, which affect the function of many recombinant glycoproteins. In the present study, tobacco (N. benthamiana) and lettuce (L. sativa) were used to produce 2C10 mAb, which previously showed the PEDV neutralization [16].

Among all PEDV neutralizing mAb, 2C10 is one of a good candidate for using to inhibit PEDV infection [15,32,34–38]. The 2C10 mAb recognizes GPRLQPY motif found on carboxy-terminal region of PEDV S-protein [12]. Moreover, this mAb also showed strong binding to the peptides SHRLP(Y/Q)(P/V) or GPVPVTH on the g3p minor coat protein and strong neutralizing activity against KPED-9 strain [15].

The nucleotide sequences of 2C10 VH and VL from previous work [24] were codon usage optimized for N. benthamiana and synthesized. After fusing both VH and VL to the constant region of heavy chain (CH) and light chain (CL), the 2C10 gamma and kappa gene were cloned into geminiviral replicon vector, separately. To test whether 2C10 mAb produced from plants can function, we used the human IgG constant region which is available in the lab. If we get the promising result, the porcine IgG constant region will be used for further study. Geminiviral replicon systems have been used for production of several recombinant proteins in plants, such as vaccine antigens [22–24,39,40] and antibodies [22,24,25]. The transient expression in plants using geminiviral vectors can rapidly produce high level of recombinant protein expression.

In this study, 2C10 mAb was transiently expressed in tobacco and lettuce on day 5 and day 4 after agroinfiltration, respectively (▶Fig. 3A and Fig. 4A). This expression process generated 2C10 mAb in a short period of time. After purification process, the assembly of full-size 2C10 IgG (150 kDa) produced from both plants was detected in SDS-PAGE under non-reducing condition (▶Fig. 3B and Fig. 4B). The quantity of corrected IgG was found in tobacco more than lettuce. However, the antibodies produced from both plants clearly showed binding to PEDV in vitro (▶Fig. 5). Our result showed that the binding of plant-produced 2C10 mAb to PEDV increased approximately 1.7 times. However, the binding of another protein, GST-truncated S, is similar to negative control (the binding ratio is approximately 1). The binding activity of plant-produced 2C10 mAb strongly suggests that it can be developed to use as a reagent in the assays for detection of PEDV.

To test the neutralization of PEDV by 2C10, PEDV strain ST was used in this study. The 2C10 IgG produced from N. benthamiana showed the neutralizing activity at dilution 1:32 (▶Table 1). In this study, the neutralizing activity of plant-produced 2C10 was not strong compared to the neutralizing activity of 2C10 mAb against KPED-9 strain in previous study [15]. The level of neutral-
izing activity might be different depending on the strain of the virus. This data confirmed that the plant-produced 2C10 mAb neutralized PEDV infection in vitro, suggesting the potential to use as PEDV prophylactic agent. Previous data reported that protecting suckling piglets from PEDV infection also depends on the secretory IgA (sIgA) antibodies in milk [41]. The sIgA is more resistant to protease than IgG, which is suitable to use as the oral vaccine [42]. Therefore sIgA is a good approach to use as a prophylactic and therapeutic treatment. Plants were shown to be an effective platform for sIgA production [43–45]. Thus, we can develop this 2C10 mAb to produce sIgA from plants in the future, which will be more effective PEDV prophylactic.

The results of the present study provided a proof of concept in which the PEDV specific antibody can be produced through plant expression system and the produced antibody has an in vitro efficacy against PEDV. Further in vivo investigation involving oral application of plant produce antibody in piglets against oral PEDV challenge is suggested. The in vivo study, although can address the efficacy of the produced antibody, is needed to control several factors including methods to prevent enzymatic activities in gastrointestinal tracts, pH factors, and delivery system in which is beyond the scope of the study. The level of antibody needed to protect young pigs from the PEDV infection is another study that is needed for further investigation. In this study, the neutralization showed at titer 1:32 and at higher concentration of the antibody. However, at which certain level is beyond the scope of the study as the level of antibody induced by either natural infection or vaccination that provide a complete protection is presently yet to be known.

In conclusion, we developed a novel plant-produced PEDV 2C10 mAb, which showed binding and neutralizing activity against PEDV. This technology provides an effective and low-cost platform to produce PEDV mAb, which could be developed for PEDV diagnostic and prophylactic agents. The platform developed here will be beneficial, especially to the swine industry.

Materials and Methods

Design of the constructs for producing 2C10 mAb in plants

We designed plant-optimized DNA sequences encoding 2C10 VH and VL based upon previous study [16], using codons that are preferred in tobacco. The VH was fused to the N-terminus of a gene encoding constant region of the humanized 62-71-3 mAb [46]. The synthetic VH gene was amplified using primers Ncol-VH-F and PmlI-VH-R whereas the 62-71-3 CH was amplified using PmlI-CH-F and Sacz-CH-R. All primer sequences are in Table 2. The overlapped extension PCR between VH and CH was performed using primers Ncol-VH-F and Sacz-CH-R. For the cloning of 2C10 light chain, the synthetic VL gene was amplified using primers Ncol-VL-F and XhoI-VL-R whereas the 62-71-3 CL was amplified using primers XhoI-CL-F and Sacz-CL-R. The overlapped PCR between VL and CL was performed using primers Ncol-VL-F and Sacz-CL-R. The genes of 2C10 heavy and light chain were inserted into the geminiviral vector via Ncol and Sacz sites. Two geminiviral vectors, 2C10 heavy chain and light chain genes, was co-infiltrating with p19 vector.

Plant inoculation and protein expression

The seeds of N. benthamiana were provided from Julian Ma’s Lab (London, UK). N. benthamiana plants were co-infiltrated with Agrobacterium GV3101 strains containing pBY2C10-gamma, pBY2C10-kappa, and p19 that protects gene silencing of transient protein production in plant (Fig. 2). The final OD600 of mixed Agrobacterium in equal amount of the three strains for vacuum infiltration is 0.25. Plants were maintained in the growth chamber for 5 d before protein extraction. Romaine lettuce (L. sativa) was obtained commercially from a local grocery store. The agroinfiltration of lettuce was performed as previously described [24]. After vacuum infiltration with Agrobacterium, lettuce heads were covered with Saran wrap and kept in the growth chamber for 4 d before protein extraction.

Protein purification

Protein purification Infiltrated tobacco and lettuce leaves were homogenized by using a blender with 1× phosphate-buffered saline (1× PBS: 137 mM NaCl; 2.7 mM KCl; 4.3 mM Na2 HPO4; 1.47 mM KH2PO4 at pH 7.4), pH 7.5. Crude extract was filtered through Miracloth and centrifuged at 26,000 g for 20 min. The solution was filtered with 0.2 μm filter before loading into the protein A bead column. After washing the column with PBS, pH 7.5, for 10 bed volumes, the protein was eluted with 100 mM glycine, pH 2.5. After the protein was eluted from the column, 1 M Tris base was added to neutralize to a final pH of 7.5.
**Table 1** Result of viral neutralization assay. Different dilutions of plant-produced 2C10 IgG were incubated with PEDV 1 h before transfer to Vero cells. After 1 h, the mixed of viral and 2C10 IgG were removed. The Vero cells plate was continuous incubated at 37℃ for 5–7 days for CPE determination. Positive control: colostrum sample with PEDV neutralizing antibody; negative control: colostrum sample without PEDV neutralizing activity; +: cytopathic effect; --: no cytopathic effect.

<table>
<thead>
<tr>
<th>mAb/ dilution</th>
<th>1:4</th>
<th>1:8</th>
<th>1:16</th>
<th>1:32</th>
<th>1:64</th>
<th>1:128</th>
<th>1:256</th>
</tr>
</thead>
<tbody>
<tr>
<td>N. benthamiana 2C10</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>1× PBS</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Negative control</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Positive control</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

**Table 2** Primer list.

<table>
<thead>
<tr>
<th>Primer Name</th>
<th>Sequencing (5′-3′)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NcoI-VL-F</td>
<td>CCAATGGACATGAGAGTTCCAGCGATATTG</td>
</tr>
<tr>
<td>XhoI-VL-R</td>
<td>CTCCGAGACCCCTGTACCTCC</td>
</tr>
<tr>
<td>XhoI-CL-F</td>
<td>CTCGCCGAGTGTGCTGCTCC</td>
</tr>
<tr>
<td>SacI-CL-R</td>
<td>GACGCTTACCTCCGCCCCTATTTG</td>
</tr>
<tr>
<td>NcoI-VH-F</td>
<td>CCACTGGAACCTGGACTTTGG</td>
</tr>
<tr>
<td>PmlI-VH-R</td>
<td>CACGTGTGAGTCTTATCGCAG</td>
</tr>
<tr>
<td>PmlI-CH-F</td>
<td>CACGTGTCCACCATGGTCAG</td>
</tr>
<tr>
<td>SacI-CH-R</td>
<td>GAGCCTTCATTGCAGGGGACAAAG</td>
</tr>
</tbody>
</table>

**SDS-PAGE and western blotting**

The purified plant 2C10 mAb protein was mixed with SDS loading sample buffer (250 mM Tris–HCl pH 6.8, 10% [w/v] SDS, 30% [v/v] glycerol, 5% [w/v] β-mercaptoethanol, and 0.02% [w/v] bromophenol blue) for reducing SDS-PAGE or SDS loading sample buffer without β-mercaptoethanol for non-reducing SDS-PAGE and boiled for 5 min. Proteins were separated using 10% SDS-PAGE. Proteins were stained with Coomassie Brilliant Blue G-250. For western blot, proteins were transferred to a nitrocellulose membrane (Bio-Rad) and probed with HRP-conjugated mouse anti-human IgG1 antiserum or Fc specific or HRP-conjugated mouse anti-kappa light chain antiserum, diluted at 1:1000 in 1% non-fat milk (Bio-Rad) and probed with HRP-conjugated mouse anti-human IgG1 antiserum or Fc specific, (diluted 1:1000 in 1% skim milk) at 37 ℃ for 2 h. The plate was washed three times with PBST and incubated with HRP-conjugated anti-human IgG antiserum, Fc specific, (diluted 1:1000 in 1% skim milk) at 37 ℃ for 2 h. The plates were washed again and developed using 3, 3′, 5′, 5′-tetramethy benzidine liquid substrate for 15 min at 23 ℃ in dark condition. The reactions were stopped with 1 N H2SO4 and determined with micro-plate reader at OD450.

**Binding affinity analyzed by ELISA**

After titration and optimal dilution of PEDV, polystyrene 96 well microtiter plate were coated with solution from the media from Vero cell as negative control and viral solution and incubated at 4 ℃ overnight. The plate was blocked with 5% skim milk in 1× PBS at 37 ℃ for 1 h and then subsequently incubated with 1:1000 plant 2C10 mAb diluted in 1% skim milk at 37 ℃ for 2 h. The plate was washed three times with PBST and incubated with HRP-conjugated anti-human IgG antiserum, Fc specific, (diluted 1:1000 in 1% skim milk) at 37 ℃ for 2 h. The plates were washed again and developed using 3, 3′, 5′, 5′-tetramethy benzidine liquid substrate for 15 min at 23 ℃ in dark condition. The reactions were stopped with 1 N H2SO4 and determined with micro-plate reader at OD450.

**Neutralization assay**

Vero cells were seeded into the 96-well plates and grown until 80% coverage of the plate. PEDV virus was diluted to TCID50 (50% tissue culture infectious dose) in MM medium and incubated with 100 µl of different dilutions of plant-made 2C10 mAb (1:4, 1:8, 1:16, 1:32, 1:64, 1:128, and 1:256) at 37 ℃ for 1 h. The Vero cells were washed with PBS one time and MM containing 10 µg/mL trypsin one time. The viral mixed solutions were transferred to the Vero cells plate and incubated at 37 ℃ for 1 h. Following the incubation, the solution was removed and MM plus 2% FBS was added. The Vero cells plate was incubated at 37 ℃ in a humidified 5% CO2 for 5–7 d. Colostrum samples with and without neutralizing activity were used as positive and negative control, respectively. The assays were performed in duplicate. The cytopathic effect (CPE) were determined and compared with positive and negative control.

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thank Professor Julian Ma (St. George’s Medical School, University of London, UK) for providing the facility to produce 2C10 mAb.

**Conflict of Interest**

On behalf of my co-authors, I declare that there are no financial or other conflicts of interest in the present manuscript. This work was original research, and the data presented in this article have not been published in elsewhere.

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[18] Van Nieuwstadt AP, Zetstra T. Use of two enzyme-linked immunosorbent assays to monitor antibody responses in swine with experimentally


