Abstract. The hepatitis E virus (HEV) capsid protein pORF2 comprises three potential N-linked glycosylation sites. One site, N562, is located at the cell attachment and neutralizing antigenic regions. The present study performed detailed analyses of the effects of specific amino acid substitutions at position 562 in the homodimerization, glycosylation, antigenicity, immunogenicity and neutralization activities of HEV pORF2. Recombinant HEV pORF2 glycoprotein E1 (amino acids 439-617) and three mutant variants (N562L, N562C and N562K) were expressed in Pichia pastoris (P. pastoris) and SDS-PAGE, Western blot analysis, tunicamycin assay, double-antibody sandwich ELISA and in vitro PCR-based neutralization assay were performed to characterize the different constructs. All proteins were indicated to be secreted by P. pastoris and formed homodimers. Tunicamycin assay revealed the glycosylated status of the wild-type protein, but the mutants were indicated to be non-glycosylated. All proteins were immunoreactive with a neutralizing monoclonal antibody but were not recognized by the antibody after denaturation into monomers. An in vitro PCR-based neutralization assay using mouse antibodies indicated efficient neutralization against N562L, whereas antibodies against N562C and N562K were revealed to be non-neutralizing. Collectively, the present study indicated that specific amino acid substitutions at position 562 serve crucial roles in the activity of the HEV neutralizing epitope.

Introduction

Hepatitis E virus (HEV) is an important causative agent of severe hepatitis in humans (1). However, it is a poorly understood pathogen, and has been indicated to cause high mortality during pregnancy (2) and result in chronic hepatitis in immunocompromised individuals, such as transplant recipients and HIV-infected patients with low CD4 cell counts (3,4). HEV belongs to the Orthohepevirus genus of the Hepeviridae family (5). The 7.2-kb HEV genome comprises three open reading frames (ORFs). ORF1 is a polyprotein required for HEV RNA replication, ORF2 is a capsid protein and ORF3 is a small phosphoprotein involved in virion morphogenesis and egress (6). ORF2 is involved in the formation of particle assembly, binding to host cells and elicitation of neutralizing antibodies (7-9). Its capsid protein comprises three domains: Domain S (amino acids 118-313) builds the viral shell, domain M (amino acids 314-453) contributes to a surface plateau at the 3-fold axes of the virus capsid and domain P (amino acids 454-606) forms a protruding spike from the shell (10). It also includes a cell attachment region and neutralizing antigenic region (11).

In cultured cells and samples from infected patients, HEV produces three forms of the ORF2 protein: Intracellular, glycosylated and cleaved ORF2 (12). The ORF2 protein sequence comprises three potential positions for N-linked glycosylation: N137 and N310 in the S domain and N562 in the P domain (13). N562 is highly conserved and located at the apical center of the protruding spike, which comprises a cell attachment region and neutralizing antigenic region (14,15). The location of N562 suggests that it may serve an important role in viral morphogenesis and immune responses; however, several wild strains contain an N562D mutation and it was reported that these strains successfully infected PLC/PRF/5 cells (16). Therefore, the biological functions of the other mutants at N562, and whether other types of amino acid mutations will stimulate the body to produce more efficient neutralizing antibodies, requires further elucidation. The effects of specific amino acid substitutions at position 562 serve crucial roles in the activity of the HEV neutralizing epitope.
acid substitutions at N562 on the dimerization, antigenicity, immunogenicity and production of neutralizing antibodies in the P domain remain poorly understood. The present study performed site-directed mutagenesis on the truncated capsid protein HEV pORF2 E1 (amino acids 439-617) expressed in Pichia pastoris (P. pastoris) to investigate the effects of specific amino acid substitutions at N562 in the dimerization, antigenicity, immunogenicity and production of neutralizing antibodies in the P domain.

Materials and methods

Anti-HEV IgG assay. Serum samples were collected from 18 hospital patients with HEV (10 males and 8 females; age, 36-65 years), admitted to Jinan Central Hospital Affiliated to Shandong First Medical University and Shandong Academy of Medical Sciences (Jinan, China) between January and October 2019. Patients with psychiatric illness, malignancy, pregnancy, HIV infection, herpes zoster, bowel disease and celiac disease were excluded. All patients signed informed consent, and the study was approved by the Bioethics Committee of Jinan Central Hospital Affiliated to Shandong First Medical University and Shandong Academy of Medical Sciences. Serum samples were tested for anti-HEV IgG antibodies according to the manufacturer’s protocol [HEV IgG antibody detection kit (Chemiluminescence microparticle immunoassay); cat. no. 20190904; Xiamen InnoDx. Biotech Co., Ltd.] using an automated chemiluminescent microparticle immunoassay system (Xiamen InnoDx Biotech Co., Ltd.). The result was considered positive when the signal-to-cut-off value of the sample was >1.20 (according to the manufacturer’s recommendations). The anti-HEV IgG-positive human sera were subsequently pooled and stored at -80°C until further use.

Homology modeling and in silico protein simulation. The SWISS-MODEL server accessible via the ExPASy website (http://swissmodel.expasy.org/) (17,18) was used for homology modeling. This prediction server can use the known 3D protein structure with amino acid sequence homologous to the protein of interest and predict its 3D structure based on this homology, and automatically score the 3D protein structures. The best scoring models were used to simulate the pORF2 homodimer using the SymmDock server, which is an online server for the prediction of complexes with Cn symmetry by geometry based docking (http://bioinfo3d.cs.tau.ac.il/SymmDock/), and the symmetry order used for docking was 2 (19).

Site-directed mutagenesis and construction of recombinant plasmids. Recombinant E1 encompassing residues 439-617 of genotype 4 HEV ORF2 was cloned into pET-28a (+) plasmid (Invitrogen; Thermo Fisher Scientific, Inc.), which has been previously constructed in the laboratory (20), and site-directed mutagenesis was performed to generate three mutant variants at N562. Amino acids with molecular weights similar to that of asparagine were selected (leucine, cysteine and lysine), representing aliphatic, sulfur-containing and alkaline amino acids, respectively. Regarding site-directed mutagenesis, the different E1 mutants were generated using the overlap PCR approach. The primers used for mutagenesis are listed in Table I, and PCR was performed using Expand High Fidelity PCR System (Roche) as follows: Denaturation at 94°C for 45 sec, annealing at 54°C for 30 sec and extension at 72°C for 60 sec for a total of 35 cycles. The amplicons were separated by 1% agarose gel electrophoresis and visualized by ethidium bromide.

Expression of recombinant proteins in the P. pastoris strain SMD1168. The E1 gene and three mutants carried EcoRI and XbaI restriction sites at their 5' and 3' ends, respectively. All genes were cloned into EcoRI-XbaI-digested pPICZaA (Invitrogen; Thermo Fisher Scientific, Inc.). P. pastoris SMD1168 cells (Invitrogen; Thermo Fisher Scientific, Inc.) were cultured in 3 ml Yeast Extract-Peptone-Dextrose (YPD) medium (1% yeast extract, 2% peptone, 2% dextrose) at 30°C and when the cells were in the log phase (100 µl) they were mixed with 10 µg plasmid linearized by SacI and were subsequently transferred to an electroporation cuvette and pulsed for 6 msec at 1,000 V using BioRad Gene Pulser (Gene Pulser Xcell; Bio-Rad Laboratories, Inc.). In parallel, a control P. pastoris clone harboring the empty parent vector pPICZaA was generated. The transformation mixture (200 µl) was plated in increasing concentrations of Zeocin™ (Invitrogen; Thermo Fisher Scientific, Inc.) to select putative multicopy recombinants at 30°C for 3 days.

A simple fed-batch protocol developed for high-level production of recombinant proteins by P. pastoris was used, as described previously (21). Briefly, recombinant clones were grown in 4 ml YPD medium at 30°C to an optical density of 1 at 600 nm (OD600). Subsequently, 0.5 ml of the cells were inoculated into 50 ml buffered complex glycerol medium [1% yeast extract, 2% peptone, 100 mM potassium phosphate pH 6.0, 1.34% yeast nitrogen base (YNB), 4x10⁻⁶ M biotin, 1% glycerol] grown at 30°C until culture reached OD600=20. The cells were harvested by centrifugation at 1,500 g for 5 min at room temperature, washed with sterile double-distilled water, and resuspended in 10 ml buffered methanol-complex medium [1% yeast extract, 2% peptone, 100 mM potassium phosphate pH 6.0, 1.34% YNB, 4x10⁻⁶ M biotin, 0.5% methanol]. The expression of recombinant proteins was induced for 72 h at 30°C, with 100% methanol being added to a final concentration of 0.5% every 12 h to maintain the promoter activity. After induction for 72 h, the cells were harvested by centrifugation at 13,800 x g for 20 min at room temperature. Both the cells and supernatant were stored at -80°C.

SDS-PAGE and western blotting. A total of 20 µl from 10 ml culture supernatant was mixed with 4 µl 6x SDS-PAGE loading buffer, boiled (100°C) for 5 min, analyzed on 15% SDS-PAGE and stained with Coomassie blue. For non-reducing SDS-PAGE, the buffer did not contain β-mercaptoethanol and the samples were analyzed without boiling. For western blot analysis, the separated proteins were electro transferred onto a nitrocellulose membrane. The membrane was incubated for 1 h at room temperature with blocking buffer (5% nonfat milk powder in 0.05% Tween-20-TBS). The monoclonal antibody (mAb) 5G5, which was produced in our laboratory, and anti-HEV IgG-positive human sera were used as primary antibodies (1:200) incubated at 4°C overnight. HRP-conjugated goat anti-mouse (cat. no. 5220-0287) and anti-human IgG (cat. no. 5220-0277; both from KPL) were used as secondary antibodies (1:2,000) for 2 h at room temperature. Color
SMD1168 cells were divided into two cycle and had free access to autoclaved food pellets and autoclaved water. The culture supernatants were harvested after 3 days of induction and subjected to SDS-PAGE analysis, which was the same protocol reported in the previous paragraph.

Double-antibody sandwich ELISA. Double-antibody sandwich ELISA was performed to compare the antigenicity of the four recombinant HEV capsid proteins at the same concentration. Microtiter plates were coated with mAb 4E9 [100 µl/well, 1:200 dilution in CBS buffer (pH 9.6; 0.05 M; 1.59 g NaHCO3 and 2.93 g NaHCO3 up to 1,000 ml double-distilled water) at 37˚C for 2 h. The proteins were serially diluted from 1:100 to 1:51,200 and incubated at 37˚C for 45 min. After incubation with HRP-5G5 [1:5,000; KPL USA] and mixed in equal concentrations] in 1M urea (200 µl) were collected from the ocular venous plexus of each mouse at 0, 3, 5, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36 and 38 weeks. The ocular venous plexus was selected to collect blood from the mice, as this method was indicated to cause only minimal injury to the animals and is easy to perform. Prior to blood collection, the mice were anesthetized by an intraperitoneal injection of 400 mg/kg chloral hydrate. The immune serum samples obtained from each group of mice were pooled. At the end of the animal experiment, the mice were euthanized by CO2 inhalation (60%; 3 l/min) in a 10-liter volume euthanasia chamber (23). After the animals gradually lost consciousness, CO2 concentration was increased to 100% and maintained for >10 min to ensure the animals' death. At that time, the animals did not breathe spontaneously for 2-3 min without blinking reflex. The animals died entirely unconscious. All procedures involving animals were approved and performed according to the guidelines for animal experimentation of the Jinan Central Hospital Affiliated to Shandong First Medical University and Shandong Academy of Medical Sciences Ethics Service Committee (Jinan, China). Anti-HEV IgG antibodies in mouse sera were detected by indirect ELISA, as described previously (24). Briefly, microtiter plates were coated with 100 µl/well of p166 mix [previously produced in our laboratory; four p166 recombinant proteins (amino acids 452-617 of HEV) were generated from W1 strain]. The culture supernatants of immune sera in 10% ethylene glycol were added to each well, and subsequently incubated for 45 min at 37°C, followed by incubation with an HRP-conjugated goat anti-mouse IgG (cat. no. 5220-0287, 1:5,000; KPL USA). The reaction was developed using 100 µl 3,3',5,5'-tetramethylbenzidine at 37°C for 2 h. The optical density was measured at 450 nm with a reference wavelength of 630 nm.

Table I. Primers used for mutagenesis.

| Construct | Sequence (5'-3') |
|-----------|----------------|
| E1N562    | F: CCC GAA TTC GTT ATC CAG GAC TAT GAT AAT C |
| E1N562L   | R: TCT TCT AGA TCA AGG GTA AAC AGT GTC C |
| E1N562C   | F: ATC ACT AGC AGT AGT TTG ATA ATT GTA TGG G |
| E1N562K   | R: CCA TAC AAT TAT tgt ACT ACT GCT AGT G |

The restriction sites introduced in the primer sequences are underlined. Lowercase letters indicate modified codons in the ORF2 protein.

Tunicamycin assays. For the detection of protein glycosylation, the P. pastoris SMD1168 cells were divided into two equal aliquots during induction periods: One aliquot was supplemented with tunicamycin, which was dissolved in 1% (v/v) DMSO at a final concentration of 50 µg/ml according to Ito et al (22), whereas the other aliquot was incubated without tunicamycin. The culture supernatants were harvested after 3 days of induction and subjected to SDS-PAGE analysis, which was the same protocol reported in the previous paragraph.

Immunization schedule and anti-HEV antibody detection. Each protein was emulsified with Freund's complete adjuvant (MilliporeSigma; Merck KGaA) and was used to immunize a group of six specific pathogen-free female BALB/c mice at 6 weeks of age with 20-25 g weight (obtained from the Animal Resource Centre of Yangzhou University, Yangzhou, China). During the animal experiment, the mice were maintained at 20-26°C, 40-70% relative humidity with a 12/12-h light/dark cycle and had free access to autoclaved food pellets and autoclaved water. The mice were subcutaneously inoculated at two locations on the back with 20 µg of the target protein, quantified using the Bradford method. After 3 and 5 weeks, the mice were administered an intraperitoneal booster injection with 20 µg of the target protein without the adjuvant. Blood samples (200 µl) were collected from the ocular venous plexus of each mouse at 0, 3, 5, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36 and 38 weeks. The ocular venous plexus was selected to collect blood from the mice, as this method was indicated to cause only minimal injury to the animals and is easy to perform. Prior to blood collection, the mice were anesthetized by an intraperitoneal injection of 400 mg/kg chloral hydrate. The immune serum samples obtained from each group of mice were pooled. At the end of the animal experiment, the mice were euthanized by CO2 inhalation (60%; 3 l/min) in a 10-liter volume euthanasia chamber (23). After the animals gradually lost consciousness, CO2 concentration was increased to 100% and maintained for >10 min to ensure the animals' death. At that time, the animals did not breathe spontaneously for 2-3 min without blinking reflex. The animals died entirely unconscious. All procedures involving animals were approved and performed according to the guidelines for animal experimentation of the Jinan Central Hospital Affiliated to Shandong First Medical University and Shandong Academy of Medical Sciences Ethics Service Committee (Jinan, China). Anti-HEV IgG antibodies in mouse sera were detected by indirect ELISA, as described previously (24). Briefly, microtiter plates were coated with 100 µl/well of p166 mix [previously produced in our laboratory; four p166 recombinant proteins (amino acids 452-617 of HEV) were generated from W1 strain]. The culture supernatants of immune sera in 10% ethylene glycol were added to each well, and subsequently incubated for 45 min at 37°C, followed by incubation with an HRP-conjugated goat anti-mouse IgG (cat. no. 5220-0287, 1:5,000; KPL USA). The reaction was developed using 100 µl 3,3',5,5'-tetramethylbenzidine at 37°C for 2 h. The optical density was measured at 450 nm with a reference wavelength of 630 nm.

Build Construct

E1N562

F: CCC GAA TTC GTT ATC CAG GAC TAT GAT AAT C
R: TCT TCT AGA TCA AGG GTA AAC AGT GTC C

E1N562L

F: ATC ACT AGC AGT AGT TTG ATA ATT GTA TGG G
R: CCA TAC AAT TAT tgt ACT ACT GCT AGT G

E1N562C

F: ATC ACT AGC AGT AGT ATC ATA ATT GTA TGG G
R: TAC CCA TAC AAT TAT tgt ACT ACT GCT AGT G

E1N562K

F: ATC ACT AGC AGT AGT TTG ATA ATT GTA TGG G
R: TAC CCA TAC AAT TAT aag ACT ACT GCT AGT G

The restriction sites introduced in the primer sequences are underlined. Lowercase letters indicate modified codons in the ORF2 protein.

F, forward; R, reverse.
In vitro PCR-based neutralization assay. The titers of the neutralizing antibodies were tested by an in vitro PCR-based neutralization assay (25,26). The inactivated (60°C for 30 min) serum samples were serially diluted from 1:10 to 1:80 by two-fold steps using Hanks' Balanced Salt solution (HBBS). The NJ703 HEV genotype 4 strain (GenBank accession no. AY789228; isolated from the stool of an acute HEV-infected patient in our laboratory) was diluted at 1:500 using HBBS and was mixed with equal volumes (100 µl) of the diluted serum. The mixture was incubated at 37°C for 1 h and inoculated onto an A549 cell monolayer (cat. no. CCL-185; American Type Culture Collection) in DMEM medium supplemented with 10% FBS (Invitrogen; Thermo Fisher Scientific, Inc.). After incubation for 5 days at 37°C in 5% CO₂, the cells were washed three times with HBBS, followed by immediate RNA extraction using TRIzol® reagent (Invitrogen; Thermo Fisher Scientific, Inc.). One-step reverse transcription (RT) PCR was conducted, as described previously (21). OneStep RT-PCR kit (Qiagen GmbH) and universal HEV PCR primers, forward, 5'-CGGACAGAATTGATTCGCGGC-3', and reverse, 5'-TCGCGGGCGGTGAGAGAGGCCA-3', were used as follows; RT, 50°C for 45 min and 95°C for 15 min; PCR, denaturation at 94°C for 30 sec, annealing at 62°C for 30 sec and extension at 72°C for 45 sec for 55 cycles. The amplicons were separated by 1% agarose gel electrophoresis and visualized using ethidium bromide. Neutralization was defined by the absence of detectable HEV RNA in the inoculated cell culture.

Statistical analysis. The double-antibody sandwich ELISA was repeated twice and the results are presented as the mean ± SD. Differences in the immunoreactivity with mAb 4E9 between wild-type (WT) E1 and its three mutant groups were examined by one-way ANOVA followed by Tukey's post hoc test. SPSS v20.0 (IBM Corp.) was used for statistical analysis and the results were graphically illustrated using GraphPad Prism v5.0 (GraphPad Software, Inc.). P<0.05 was considered to indicate a statistically significant difference.

Results

Predicted secondary structure models and in silico simulations of homodimerization. The WT protein and its mutants exhibited similar secondary structures, with the WT and the three mutated residues all located on the coils (Fig. 1A). The best ranked 3D models were selected among the available E1 protein structures using the SWISS-MODEL server. Subsequently, dimerization was simulated using the SymmDock server. The best simulations (the molecular simulations closest to the spatial structure of the target protein) indicated that N562 was engaged in the formation of homodimers, which is the case in the proposed quaternary structure of the HEV-like particle (Protein Data Bank ID: 2ZTN). To evaluate the engagement of N562 in the four predicted E1 homodimers, (E1)2, (E1 N562L)2, (E1 N562C)2 and (E1 N562K)2, the distance between two monomers at N562 was measured and compared with those in the 2ZTN model (Fig. 1B). The distances between N562 and N562 (the same position in the second monomer) was 20.9 Å in the 2ZTN model. This distance allows N562 to engage in hydrogen bonding with the neighboring amino acids in the opposite monomer and participate in the assembly of the two monomers. Accordingly, concerning (E1 N562L)2, the distance between L562 and L562' was 43.1 Å, which was closer to the distance measured in the 2ZTN model compared with that in the WT, E1 N562C and E1 N562K, which were predicted to exhibit distances of 53.3, 55.8 and 73.4 Å, respectively. This indicated that the amino acid at position 562 was better positioned to engage in homodimerization in the N562L model compared with the WT, N562C and N562K models.

Protein dimerization. Non-reducing SDS-PAGE was performed to verify the dimerization of E1 N562L, E1 N562C and E1 N562K mutants compared with that of the WT protein. E1 N562L, E1 N562C and E1 N562K were assembled into 39-kDa homodimers (Fig. 2; lanes 3, 5 and 7), whereas the WT protein was assembled into a 44-kDa homodimer (Fig. 2; lane 1). Analysis of the denatured proteins indicated that the E1 N562L, E1 N562C and E1 N562K mutant proteins were 19-kDa in size (Fig. 2; lanes 4, 6 and 8), whereas WT E1 migrated as a 21-kDa protein on the gel (Fig. 2; lane 2). Proteins from pPICZaA/SMD1168 were used as negative control.

WT protein is a glycoprotein, but its mutants are not glycosylated. The observation of a higher molecular weight of the WT protein suggested that it may be glycosylated, whereas the mutant proteins were either non-glycosylated or a relatively smaller fraction was glycosylated. Tunicamycin, which is an inhibitor of bacterial and eukaryote N-acetylglucosamine transferases, is known to prevent the glycosylation of newly synthesized glycoproteins (27). Therefore, tunicamycin assays were performed to evaluate the glycosylation status of the WT protein and its mutants. In the presence of tunicamycin, the glycosylation of the WT protein was inhibited, leading to a shift from the higher molecular weight to the lower molecular eight form (Fig. 3; lanes 1 and 2). The WT protein was validated to be a glycoprotein, as it was susceptible to tunicamycin treatment. However, the electrophoretic mobility of the three mutants was unaffected by tunicamycin treatment, suggesting that the mutants were not glycosylated (Fig. 3; lanes 3 to 8).

Immunoreactivity of the WT protein and its three mutants. Western blot analysis demonstrated that the WT protein and N562L, N562C and N562K immunoreacted with the HEV-neutralizing mAb 5G5 as homodimers but were not reactive as monomers (Fig. 4A). Western blot analysis was also performed using anti-HEV IgG-positive human sera, and the results were indicated to be similar to those obtained with mAb 5G5 (Fig. 4B). Proteins from pPICZaA/SMD1168 were used as negative control.

Antigenicity. The antigenicity of the expressed proteins was assessed using double-antibody sandwich ELISAs. The immunoreactivity levels were indicated to be decreased as the dilution increased. The immunoreactivity of WT E1 was significantly lower compared with that of the three mutant variants (Fig. 5A; P<0.05). Moreover, pPICZaA/SMD1168 exhibited no reactivity to mAb 4E9. These results indicated that a sugar chain at N562 may mask the partially neutralizing
epitopes, thereby reducing the reactivity with the neutralizing mAb.

**Immunoreactivity.** All mice vaccinated with the recombinant proteins presented HEV-specific antibodies in sera at 3 weeks post inoculation (wpi). The E1-, E1 N562L-, E1 N562C- and E1 N562K-vaccinated mice exhibited peak levels of antibody titers at 10 wpi. Overall, the kinetics of antibody responses were highly similar among the four groups, indicating that the amino acid substitutions and glycosylation status resulted in similar immunogenic properties.

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**Figure 1.** Mutational studies on the dimer interface. (A) Schematic representation of wild-type E1 and three-point mutations targeting the dimer interface region. Secondary structural elements are presented for the E1 region. The mutated residues were located on the coils, and β-strands and coils are indicated in red and black, respectively. (B) Distances between the two aa 562 in each monomer measured on the (2ZTN)2 model and the predicted homodimers (E1N562)2, (E1N562L)2, (E1N562C)2 and (E1N562K)2. aa, amino acid; HEV, hepatitis E virus.

**Figure 2.** Western blot analysis of the indicated expressed proteins. U, unheated; H, heated; M, molecular weight marker.

**Figure 3.** Effect of tunicamycin on the expression of the indicated glycoproteins at non-denaturing conditions. (−), untreated protein; (+), treated protein; M, molecular weight marker.
The HEV capsid protein encoded by ORF2 is responsible for the formation of particle assembly, binding to host cells and elicitation of neutralizing antibodies (28,29). Considerable effort has been made for the expression of recombinant HEV capsid proteins elicited similar anti-body responses and morphogenesis (38). In the present study, a set of mutant proteins containing L, C and K in place of N at position 562 was constructed and tested for homodimerization, glycosylation, antigenicity, immunogenicity and neutralization activity.

In the current study, E1 was indicated to form a 44-kDa homodimer. This molecular weight is slightly higher compared with the predicted molecular weight of 40 kDa based on the 179-amino acid sequence. The slightly higher molecular weight resulted from a facultative N-linked glycosylation site (N-X-S/T) at N562 of the P domain. Tunicamycin is an inhibitor of bacterial and eukaryotic N-acetylglucosamine transferases, preventing the formation of N-acetylglucosamine lipid intermediates and the glycosylation of newly synthesized glycoproteins (27). Tunicamycin has been indicated to block the formation of N-glycosidic linkages by inhibiting the transfer of N-acetylglucosamine-1-phosphate to dolichol monophosphate (27). Tunicamycin assay revealed that the 44-kDa protein homodimer was glycosylated. The N562L, N562C and N562K mutations completely abrogated the glycosylation, thereby supporting the importance of the asparagine residue for glycosylation at this position, as reported previously (39).

Li et al. (38) reported that the dimerization of the capsid E2s domain (ORF2; amino acids 455-602) was a prerequisite for the virus-host interaction and for the binding of certain neutralizing antibodies to HEV. The findings of the present study indicated that E1 WT, E1 N562L, E1 N562C and E1 N562K formed a homodimer, suggesting that replacing N562 with leucine, cysteine or lysine exhibited no effect on dimerization. It may be possible that the P. pastoris expression-secretion system used in the current study was conducive to the formation of homodimers.

E1, E1 N562L, E1 N562C and E1 N562K were revealed to interact with mAb 5G5 and anti-HEV IgG-positive human sera as homodimers, whereas their denaturation abrogated this immunoreactivity.

Furthermore, determination of immune response rates and antibody neutralization capacities demonstrated that all four recombinant HEV capsid proteins elicited similar antibody responses in immunized mice, but not all antibodies were neutralized in the in vitro PCR-based neutralization assay. The observation that E1 N562L elicited the strongest neutralizing antibody response among the tested proteins suggested that specific amino acid substitutions at position 562 resulted in differential effects on the immunological properties and neutralization of these proteins. However, additional experiments are required to support these results. A comparison experiment between E1 N562L and E1 N562 should be performed by measuring neutralizing antibody...
levels at different doses of E1 N562L and E1 N562, such as 5, 10 and 20 µg, to further support these conclusions.

In conclusion, analyses of the four recombinant HEV capsid proteins comprising different mutations at position 562 revealed differential effects of mutations at this position on the neutralizing capacity of the examined antibody. This finding contributed to a greater understanding of the antigenic characteristics of the glycosylated and non-glycosylated form of capsid protein ORF2 and may be helpful in the development of novel vaccines and diagnostic methods.
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Availability of data and materials

All data generated or analyzed during the present study are included in this published article.

Authors' contributions

MX performed site-directed mutagenesis and was a major contributor in writing the manuscript. LS detected and analyzed the patient with HEV for anti-HEV IgG antibody in the patient's serum, performed expression of recombinant proteins in the P. pastoris strain SMD1168. YW performed SDS-PAGE and western blotting. SG performed the immunization schedule and anti-HEV antibody detection. WY performed for the purification of proteins. ML designed the research, performed in vitro PCR-based neutralization assay and statistical analysis. All authors read and approved the final manuscript.

Ethics approval and consent to participate

All patients signed an informed consent, and the study was approved by the Bioethics Committee of Jinan Central Hospital Affiliated to Shandong First Medical University and Shandong Academy of Medical Sciences (Jinan, China). All procedures involving animals were approved and performed according to the guidelines for animal experimentation of the Jinan Central Hospital Affiliated to Shandong First Medical University and Shandong Academy of Medical Sciences Ethics Service Committee (Jinan, China).

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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