Analysis of a pair of END$^+$ and END$^-$ viruses derived from the same bovine viral diarrhea virus stock reveals the amino acid determinants in N$^{pro}$ responsible for inhibition of type I interferon production

Takashi KOZASA$^{1,2}$, Yuri ABE$^3$, Kazuya MITSUHASHI$^3$, Tomokazu TAMURA$^3$, Hiroshi AOKI$^4$, Masatoshi ISHIMARU$^2$, Shigeyuki NAKAMURA$^3$, Masatoshi OKAMATSU$^3$, Hiroshi KIDA$^{3,5,6}$ and Yoshihiro SAKODA$^{3,6}$*

$^1$Food Safety and Consumer Affairs Bureau, Ministry of Agriculture, Forestry and Fisheries, Chiyoda-ku, Tokyo 100–8950, Japan
$^2$National Veterinary Assay Laboratory, Ministry of Agriculture, Forestry and Fisheries, Kokubunji, Tokyo 185–8511, Japan
$^3$Laboratory of Microbiology, Department of Disease Control, Graduate School of Veterinary Medicine, Hokkaido University, Sapporo 060–0818, Japan
$^4$Department of Basic Science, School of Veterinary Nursing and Technology, Faculty of Veterinary Science, Nippon Veterinary and Life Science University, Musashino, Tokyo 180–8602, Japan
$^5$Research Center for Zoonosis Control, Hokkaido University, Sapporo 001–0020, Japan
$^6$Global Station for Zoonosis Control, Global Institution for Collaborative Research and Education (GI-CoRE), Hokkaido University, Sapporo 001–0020, Japan

(Received 18 August 2014/Accepted 16 December 2014/Published online in J-STAGE 28 December 2014)

ABSTRACT. The Exaltation of Newcastle disease virus (END) phenomenon is induced by the inhibition of type I interferon in pestivirus-infected cells in vitro, via proteasomal degradation of cellular interferon regulatory factor (IRF)-3 with the property of the viral autoprotease protein N$^{pro}$. Reportedly, the amino acid residues in the zinc-binding TRASH motif of N$^{pro}$ determine the difference in characteristics between END-phenomenon-positive (END$^+$) and END-phenomenon-negative (END$^-$) classical swine fever viruses (CSFVs). However, the basic mechanism underlying this function in bovine viral diarrhea virus (BVDV) has not been elucidated from the genomic differences between END$^+$ and END$^-$ viruses using reverse genetics till date. In the present study, comparison of complete genome sequences of a pair of END$^+$ and END$^-$ viruses isolated from the same virus stock revealed that there were only four amino acid substitutions (D136G, D3148G and D3502Y) between two viruses. Based on these differences, viruses with and without mutations at these positions were generated using reverse genetics. The END assay, measurements of induced type I interferon and IRF-3 detection in cells infected with these viruses revealed that the aspartic acid at position 136 in the zinc-binding TRASH motif of N$^{pro}$ was required to inhibit the production of type I interferon via the degradation of cellular IRF-3, consistently with CSFV.

KEY WORDS: bovine viral diarrhea virus, END phenomenon, interferon regulatory factor-3, N$^{pro}$, type I interferon

doi: 10.1292/jvms.14-0420; J. Vet. Med. Sci. 77(5): 511–518, 2015

The genus Pestivirus of the family Flaviviridae comprises 4 recognized species that are economically important pathogens in the veterinary field: bovine viral diarrhea viruses (BVDVs) 1 and 2, classical swine fever virus (CSFV) and border disease virus of sheep [30]. Pestiviruses are quasispecies; a single strain consists of different populations showing various characteristics. Two biotypes of pestiviruses, cytopathogenic (cp) and noncytopathogenic (ncp) viruses, are distinguished by their ability to induce a cytopathic effect (CPE) in tissue cultures [16]. Pestiviruses have been reported to inhibit the production of type I interferon in infected cells in vitro in coordination with the property of viral protein N$^{pro}$ [1, 4, 8, 28]. Subsequently, Newcastle disease virus (NDV) and some orbiviruses replicate efficiently and induce distinguishable CPE in cells infected with pestiviruses [15, 20]. The exaltation of NDV (END) phenomenon is a well-known property of BVDV and CSFV that has been used as a biological marker of vaccine virus and for the titration purpose. Nep pestiviruses are further divided into two types by their ability to induce END phenomenon: END-phenomenon-positive (END$^+$) and END-phenomenon-negative (END$^-$) viruses. A pair of END$^+$ and END$^-$ viruses can be cloned from the same virus stock using reverse plaque formation techniques [5, 19]. END$^-$ viruses do not enhance NDV, but induce intrinsic interference against Western equine encephalitis virus and vesicular stomatitis virus (VSV) [5, 19]. One of the END$^-$ viruses, the GPE strain of CSFV, attenuated and cloned from the virulent ALD strain [29], is a Japanese live vaccine strain against classical swine fever (CSF) that contributed to the eradication of CSF in Japan.

It is well documented that END$^+$ pestiviruses subvert host innate immune defenses by preventing the production of type I interferon. This occurs via the proteasomal degradation of cellular interferon regulatory factor (IRF)-3 in infected cells through its interaction with viral autoprotease.
ase Npro [1, 4, 8, 28]. Npro, the first functional unit of the pestivirus polyprotein, is followed by structural proteins (C, E\textsuperscript{\textit{E1}} \textit{E2}) and nonstructural proteins (p7, NS2, NS3, NS4A, NS4B, NS5A and NS5B) [3, 16, 25, 31, 35]. Npro is a unique protein in pestiviruses that autocatalytically cleaves itself from the nascent polyprotein to generate the N terminus of the viral capsid (C) protein [26]. Npro has two domains: a catalytic N-terminal domain and a C-terminal domain containing a zinc-binding TRASH motif. The zinc-binding TRASH motif, which includes the zinc-binding residues Cys112-Cys134-Asp136-Cys138, is required for binding TRASH motif, which includes the zinc-binding domain containing a zinc-binding TRASH motif. The zinc-domains: a catalytic N-terminal domain and a C-terminal domain containing a zinc-binding TRASH motif.

Recent studies have shown that Npro of CSFV interacts with IRF-7 and dampens the production of type I interferon in plasmacytoid dendritic cells [4]. It has also been reported for CSFV that deletion of 19 amino acids from the N terminus of Npro does not abolish its capacity to inhibit the production of type I interferon [7, 24] and that amino acid substitutions C112A/R, C134A, D136N or C138A in Npro result in the disappearance of this function [24, 32]. In contrast, Ruggli et al. and Tamura et al. succeeded in restoring the Npro functions of wild-type END viruses, the Ames-END and GPE strains of CSFV, by mutating a single amino acid residue (R112C and N136D, respectively) using reverse genetics techniques [24, 33]. Npro of both cp BVDV (e.g., the NADL strain) and ncp BVDV (e.g., the pe515 strain) blocks type I interferon production in vitro via proteasomal degradation of cellular IRF-3 [2, 8]. It has been reported for BVDV that nearly the entire Npro is required for the prevention of type I interferon production because removal of 30 residues from the N terminus or the removal of 24 or 88 residues from the C terminus abolishes this function [2, 8]. Deletion mutants expressing residues 1–69 or 70–168 of Npro also lack the ability to inhibit the production of type I interferon [2]. In addition, the previous reports revealed that substitutions of amino acid residue L8P, E22V or H49L of Npro abolish its capacity to function as an interferon antagonist [2, 6].

The BVDV GBK strain is an adventitious BVDV2 isolated from cells of the bovine kidney cell line GBK [9]. The BVDV GBK strain was grown in BT cells. BVDV GBK E+ and GBK E− strains were grown in MDBK-HS cells. The NDV Miyadera strain was propagated in 10-day-old embryonated hens’ eggs. The New Jersey serotype strain of VSV was grown in SK-L cells.

Cloning of a pair of END and END viruses from viral stocks: A reverse plaque formation technique [5] was used to obtain a pair of END+ and END− viruses from viral stocks as described previously [19]. Viruses were used after three rounds of limited dilution.

Sequencing: BVDV GBK E+ and GBK E− strains, the full-length cDNA clones and the in vitro-rescued viruses were completely sequenced as described previously [34]. In brief, the nucleotide sequences of cDNA clones and PCR fragments from viral RNA were determined using the BigDye Terminator v3.1 Cycle Sequencing Kit (Life Technologies) and a 3130 Genetic Analyzer or a 3500 Genetic Analyzer (Life Technologies) according to the manufacturer’s protocol. Sequencing data were analyzed using GENETYX version 10 software (GENETYX, Tokyo, Japan).

Plasmid constructs: The cDNA fragments from the GBK E− strain, obtained by reverse transcription polymerase chain reaction (RT-PCR), were cloned into plasmid pCR®-Blunt II-TOPO® (Life Technologies) using the ZeroBlunt® TOPO® Cloning Kit (Life Technologies). The cDNA sequence was flanked by a modified T7 promoter sequence at the 5′ end and a Pst I restriction site at the 3′ end. Subclones were assembled into a full-length cDNA clone, termed pGBK E−, by replacing the CSFV genome of the full-length cDNA clone of the CSFV Alfıort187-1 strain pA187-1 [23] with the genome of the GBK E− strain using appropriate restriction enzymes and the In-Fusion HD Cloning Kit (Clontech, Mountain View, CA, U.S.A.) according to the manufacturer’s protocol. Details of the constructions may be obtained on request. The full-length cDNA clone pGBK E− was transformed and propagated in competent cell Stbl3 cells (Life Technologies) and purified using the QIAGEN Plasmid Plus Midi Kit (QIAGEN, Hilden, Germany) according to the manufacturer’s protocol.

Three full-length cDNA clones with combinations of the four amino acid substitutions D136G, I263V, D3148G and D3502Y were constructed in the pGBK E+ backbone using...
the QuickChange Lightning Multi Mutagenesis Kit (Agilent Technologies, Santa Clara, CA, U.S.A.) and the In-Fusion HD Cloning Kit. The plasmid pGBK_E+/D136G has a single amino acid substitution at position 136 of GBK_E+, whereas the plasmids pGBK_E+/I2623V; D3148G; D3502Y and pGBK_E+/D136G; I2623V; D3148G; D3502Y have multiple amino acid substitutions at positions 2623, 3148 and 3502 of GBK_E+ and at positions 136, 2623, 3148 and 3502 of GBK_E+, respectively.

**Full-length PCR amplification, in vitro RNA transcription, transfection and viral recovery:** The cDNA-derived viruses were rescued as described previously [34] with some modifications. The full-length genome amplification strategy [22] was employed to obtain a full-length PCR amplicon for in vitro RNA transcription. The cDNA clones were amplified using primers 5GBKE +_T7 (5′-taa tac gac tca cta ta GTA TAC GAG ATT AGC TAA AGT ACT CG −3′, T7 promoter sequence underlined) and 3GBKE + (5′-GGG GCT GTT AGA GGC ATC CTC TAG TC −3′) with AccuPrime Taq DNA polymerase High Fidelity (Life Technologies). Viral RNA was transcribed in vitro from the purified full-length PCR amplicon using the MEGAscript T7 Kit (Life Technologies), the remaining PCR amplicon was digested using DNase (Life Technologies), and the viral RNA was then purified with a MicroSpin S-400 column (GE Healthcare, Buckinghamshire, U.K.). MDBK-HS cells (3 × 10^6 cells) were transfected with 10 µg of viral RNA in a 0.4 cm cuvette using the Gene Pulser Xcell Electroporation system (Bio-Rad, Hercules, CA, U.S.A.) set at 180 V and 950 µF. The cells were incubated at 37°C in 5% CO₂ for 3 days, and the supernatants were then transferred onto naive MDBK-HS cells to obtain infectious viruses.

The viruses were named according to the plasmid from which they were rescued, replacing “p” with “v” in the nomenclature. The cDNA-derived viruses generated in the present study are shown in Fig. 1.

**END assay and measurements of type I interferon production:** The END assay was conducted using BFM cells as described previously [10]. In brief, BFM cells grown in 96-well plates were infected with BVDV and incubated for 5 days at 37°C in 5% CO₂. After removal of the supernatants, the cells were superinfected with 1 HA/ml of the NDV Miyadera strain. The END phenomenon was regarded as positive, if strong CPE was observed in NDV-infected cells.

Measurements of type I interferon production in BVDV-infected cells were performed using a previously described plaque reduction method with VSV as a challenge virus [20], with some modifications. In brief, the supernatants from BVDV-infected cells were inactivated by exposure to UV light (254 nm) in the UV Crosslinker (ATTO Corporation, Tokyo, Japan) under the condition of 500 mJ/cm². Viral inactivation of samples was confirmed using indirect FA techniques with anti-BVDV NS3 monoclonal antibody #46/1 [13] after inoculation of samples onto MDBK-HS cells and incubation for 2 days at 37°C in 5% CO₂. Then, monolayers of SK-L cells in 12-well plates were inoculated with 1 ml of a four-fold dilution of UV-inactivated supernatant and incubated for 24 hr at 37°C in 5% CO₂. The supernatants were removed, and the cells were inoculated with VSV. Interferon titers were expressed as reciprocals of dilutions that reduced the number of challenge viral plaques by 50%.

**Detection of IRF-3 in BVDV-infected cells by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and western blotting:** SDS-PAGE and western blotting were performed as described previously [12]. BFM cells were infected with BVDV (m.o.i. of 1) in six-well plates and incubated for 2 days at 37°C in 5% CO₂. Then, monolayers of SK-L cells in 12-well plates were inoculated with 1 ml of a four-fold dilution of UV-inactivated supernatant and incubated for 24 hr at 37°C in 5% CO₂. The supernatants were removed, and the cells were inoculated with VSV. Interferon titers were expressed as reciprocals of dilutions that reduced the number of challenge viral plaques by 50%.

![Fig. 1. Schematic representation of the genomic differences between GBK_E+ virus and GBK_E− virus, and mutant viruses derived from cDNA clones generated in the present study. (A) Four amino acid substitutions, D136G, I2623V, D3148G and D3502Y, were found in the GBK_E+ virus in comparison with the GBK_E− virus. (B) Three recombinant viruses with combinations of amino acid substitutions were generated in the GBK_E+ backbone by site-directed mutagenesis and In-Fusion techniques. The white and gray boxes indicate the nonstructural and structural proteins, respectively.](image-url)
treated with anti-human IRF-3 rabbit polyclonal antibodies (GeneTex, Hsinchu City, Taiwan), goat anti-rabbit IgG-HRP conjugate (Bio-Rad) and Immunoblotting Western Detection Reagents (Millipore), in that order. The membranes were read using Lumi Vision PRO (Aishin Seiki, Kariya, Japan), and specific bands for IRF-3 were detected.

RESULTS

Cloning of a pair of END+/END− viruses from the BVDV GBK strain and determining their whole genome sequences: A pair of END+ and END− viruses (GBK_E+ and GBK_E−, respectively) was cloned from the BVDV GBK strain by means of reverse plaque formation techniques and limited dilution. The genomes of GBK_E+ and GBK_E− viruses were both 12,284 nucleotides in length and encoded 3,897 deduced amino acids. Comparison of the complete genome sequences of both viruses revealed only six nucleotide and four amino acid differences (D136G, I2623V, D3148G and D3502Y, the numbers refer to the amino acid position in GBK_E+) between the two viruses (Table 1). The complete genome sequences of the GBK_E+ and GBK_E− strains were deposited in the DDBJ/EMBL/GenBank databases under accession numbers AB894423 and AB894424, respectively.

Generation and characterization of in vitro-rescued viruses: The infectious BVDV vGBK_E+ was successfully rescued by electroporation of MDBK-HS cells with viral RNA transcribed in vitro from a full-length PCR amplicon obtained from a cDNA clone pGBK_E+. The complete sequence of vGBK_E+ was entirely identical to that of GBK_E+. The mutant viruses vGBK_E+/D136G, vGBK_E+/I2623V, D3148G; and vGBK_E+/D136G; I2623V; D3148G; and D3502Y were also recovered from full-length cDNA clones. Sequencing of the complete genomes of these three viruses confirmed the mutations at the desired amino acid positions and demonstrated the lack of any other mutations in comparison with the parental GBK_E+ virus.

To investigate the biological properties of the mutant viruses, BFM cells were inoculated with both the parental GBK_E+ and the in vitro-rescued vGBK_E+ viruses. The results revealed that both GBK_E+ and vGBK_E+ were ncp (data not shown) and exhibited the END phenomenon (END+). In addition, titration of both viruses in MDBK-HS cells revealed that vGBK_E+ exhibited the same growth characteristics as wild-type GBK_E+ (data not shown). Moreover, compared with the wild-type GBK_E+ strain, other in vitro-rescued viruses grew equally in MDBK-HS cells (data not shown).

Identification of the amino acid determinants responsible for inhibition of type I interferon production: As expected, the mutant viruses vGBK_E+/D136G, vGBK_E+/I2623V; D3148G; D3502Y and vGBK_E+/D136G; I2623V; D3148G; D3502Y remained ncp (data not shown). An END assay revealed that the vGBK_E+/D136G and vGBK_E+/D136G; I2623V; D3148G; D3502Y viruses were changed to END−; BFM cells infected with these viruses did not exhibit CPE after NDV superinfection. In contrast, the vGBK_E+/I2623V; D3148G; D3502Y virus remained END+; BFM cells infected with this virus exhibited clear CPE after super-infection with NDV, as observed in BFM cells infected with GBK_E+ (Fig. 2).

The amount of type I interferon in supernatants from BFM cells infected with the parent or one of the mutant viruses GBK_E+, vGBK_E+, vGBK_E+/D136G, vGBK_E+/I2623V; D3148G; D3502Y or vGBK_E+/D136G; I2623V; D3148G; D3502Y was measured in SK-L cells. Secretion of type I interferon was inhibited in BFM cells infected with vGBK_E+, vGBK_E+ or the virus carrying the mutations without Npro (vGBK_E+/I2623V; D3148G; D3502Y), which is in consistent with the results from the END assay. Cells infected with GBK_E+ or viruses carrying mutations in Npro (vGBK_E+/D136G and vGBK_E+/D136G; I2623V; D3148G; D3502Y) did induce type I interferon (Table 2).

These results clearly indicate that the aspartic acid of Npro (at position 136 of the GBK_E+ strain) is the key amino acid residue that determines the capacity to inhibit the production of type I interferon and induce the END phenomenon.

Detection of IRF-3 in cells infected with the parental or mutant viruses: To investigate the expression levels of IRF-3 in BVDV-infected cells, IRF-3 in the lysates of BFM cells infected with GBK_E+, vGBK_E+ or one of the three mutant viruses was detected by western blotting. IRF-3 was not detected in the lysates of cells infected with GBK_E+ or vGBK_E+ or vGBK_E+/I2623V; D3148G; D3502Y, whereas clear bands of IRF-3 appeared in the lysates of cells infected with the END− GBK_E+ virus or one of the Npro mutant viruses (vGBK_E+/D136G and vGBK_E+/D136G; I2623V; D3148G; D3502Y) (Fig. 3).

Comparison of amino acid sequences of zinc-binding TRASH motif in Npro: The amino acid sequences of the zinc-binding TRASH motifs in the Npro proteins from various BVDV strains deposited in the DDBJ/EMBL/GenBank databases were compared with those of the GBK_E+ and GBK_E− strains. At least one strain per subgenotype [1a−1b (except for 1f) and 2a−2c] [11, 18] was chosen in the present study. As a result, BVDV strains, except for the G strain, contain amino acid residues Cys112-Cys134-Asp136-Cys138 in the zinc-binding TRASH motif of Npro, as observed in GBK_E+ (Fig. 4).

DISCUSSION

Both BVDV and CSFV inhibit the production of type I interferon in vitro through proteasomal degradation of cellular IRF-3 [1, 4, 8, 28]. Cells infected with these (END+) viruses exhibit clear CPE after super-infection with NDV.
viruses exhibit strong CPE after superinfection with NDV or some orbiviruses [15, 20]. One of the two domains of pestivirus Npro, the C-terminal domain containing a zinc-binding TRASH motif consisting of Cys112-Cys134-Asp136-Cys138, is required for IRF-3 binding and prevention of type I interferon production [7, 32]. For CSFV, it has been reported that mutations at positions 112, 134, 136 or 138 in Npro of END+ viruses abolish the inhibition of type I interferon production, while mutations at positions 112 or 136 in Npro of END− viruses restore this function [24, 32, 33]. For

Fig. 2. Results of END assays. BFM cells infected with GBK_E+, GBK_E− or the in vitro-rescued viruses were superinfected with Newcastle disease virus (NDV). Cells infected with GBK_E+, vGBK_E+ or vGBK_E+/I2623V; D3148G; D3502Y exhibited distinguishable CPE after superinfection with NDV (END+), whereas those infected with GBK_E−, vGBK_E+/D136G or vGBK_E+/D136G; I2623V; D3148G; D3502Y did not exhibit CPE (END−). Scale bar, 5 µm.
T. KOZASA ET AL.

BVDV, previous studies have revealed that amino acid substitutions L8P, E22V or H49L in Npro abolish its capacity to interfere with type I interferon production [2, 6]. However, to the best of our knowledge, there were no studies with BVDV that had approached the basic mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quasispecies; single strains consist of populations with various characteristics, such as END + and END − [17]. We cloned a pair of END + and END − viruses (GBK_E+ and GBK_E−) from the BVDV GBK strain using reverse plaque formation techniques [5, 19]. Determination of complete genome sequences of these viruses revealed that there were only four amino acid differences between them (Table 1). To clarify the molecular mechanism of the inhibition of type I interferon production using the amino acid differences between a pair of END + and END − viruses as well as viruses genetically engineered on the basis of these differences. Pestiviruses are quas
in cellular IRF-3 (Fig. 3). Therefore, the results indicate that the inhibition of type I interferon production and the END phenomenon displayed by GBK_E- occurs by the degradation of cellular IRF-3 caused by combining with the zinc-binding TRASH motif. Furthermore, this function was abolished by mutation of Npro at position 136. Comparison of vGBK_E+/D136G; I2623V; D3148G; D3502Y (the same amino acid sequence as GBK_E-) with vGBK_E-/I2623V; D3148G; D3502Y revealed that the function of Npro was restored by a single G136D mutation in Npro of the GBK_E- virus.

In the present study, a comparison of the amino acid sequences of GBK_E+, GBK_E- and viruses from DDBJ/EMBL/GenBank databases revealed that amino acid residues Cys112-Cys134-Asp136-Cys138 in the zinc-binding TRASH motif of Npro were well conserved among field BVDV isolates (Fig. 4). It is reported that these four residues are required for the inhibition of type I interferon production [7, 32]. Taken together, the above-mentioned results suggest that field isolates inhibit the type I interferon production in infected cells, although there remains a possibility that amino acid residues other than those of TRASH motif contribute to this phenomenon. The G strain contains the amino acid residue Glu136 in the TRASH motif of Npro whereas two isolates (GBK_E- and END- virus) contain amino-terminal domain of bovine viral diarrhea virus inhibits DNA binding by interferon regulatory factor 3 and targets it for proteasomal degradation. J. Virol. 80: 11723–11732. [Medline] [CrossRef]

4. Fiebach, A. R., Guzyłack-Piriou, L., Python, S., Summerfield, A. and Ruggli, N. 2011. Classical swine fever virus Npro limits type I interferon induction in plasmacytoid dendritic cells by interacting with interferon regulatory factor 7. J. Virol. 85: 8002–8011. [Medline] [CrossRef]

5. Fukusho, A., Ogawa, N., Yamamoto, H., Sawada, M. and Sazawa, H. 1976. Reverse plaque formation by hog cholera virus of the GPE strain inducing heterologous interference. Infect. Immun. 14: 332–336. [Medline]

6. Gil, L., Hansri, I. H., Vassilev, L., Liang, D., Lai, V. C., Zhong, W., Hong, Z., Dubovi, E. J. and Donis, R. O. 2006. The amino-terminal domain of bovine viral diarrhea virus Npro protein is necessary for alpha/beta interferon antagonism. J. Virol. 80: 900–911. [Medline] [CrossRef]

7. Gottipati, K., Ruggli, N., Gerber, M., Tratschin, J. D., Benning, M., Bellamy, H. and Choi, K. H. 2013. The structure of classical swine fever virus Npro: a novel cysteine autoprotease and zinc-binding protein involved in subversion of type I interferon function. J. Virol. 87: 3828–3838. [Medline] [CrossRef]

8. Inaba, Y., Tanaka, Y., Kumagai, T., Omori, T. and Ito, H. 1968. Bovine diarrhea virus. II. END phenomenon: exaltation of Newcastle disease virus in bovine cells infected with bovine diarrhea virus. Jpn. J. Microbiol. 12: 35–49. [Medline] [CrossRef]

9. Jenckel, M., Hoper, D., Schirrmeier, H., Reimann, I., Goller, K. and Hoffmann, B. and Beer, M. 2014. Mixed triple: allied viruses in unique recent isolates of highly virulent type 2 bovine viral diarrhea virus detected by deep sequencing. J. Virol. 88: 6983–6992. [Medline] [CrossRef]

10. Kameyama, K., Sakoda, Y., Tamai, K., Igarashi, H., Tajima, M., Mochizuki, T., Namba, Y. and Kida, H. 2006. Genetic recombination at different points in the Npro-coding region of bovine viral diarrhea viruses and the potentials to change their antigenicities and pathogenicities. Virus Res. 116: 78–84. [Medline] [CrossRef]

11. Kameyama, K., Sakoda, Y., Tamai, K., Igarashi, H., Tajima, M., Mochizuki, T., Namba, Y. and Kida, H. 2006. Development of an immunochromatographic test kit for rapid detection of bovine viral diarrhea virus antigen. J. Virol. Methods 138: 140–146. [Medline] [CrossRef]

12. Kozasa, T., Aoki, H., Nakajima, N., Fukusho, A., Ishimaru, M. and Nakamura, S. 2011. Methods to select suitable fetal bovine serum for use in quality control assays for the detection of adventitious viruses from biological products. Biologicals 39: 242–248. [Medline] [CrossRef]

13. Kumagai, T., Shimizu, T. and Matumoto, M. 1958. Detection ACKNOWLEDGMENT. We are grateful to Dr. Nicolas Ruggli (Institute of Virology and Immunology, Switzerland) for providing the full-length CSFV cDNA clone pA187-1.

REFERENCES

1. Bauhofer, O., Summerfield, A., Sakoda, Y., Tratschin, J. D., Hofmann, M. A. and Ruggli, N. 2007. Classical swine fever virus Npro interacts with interferon regulatory factor 3 and induces its proteasomal degradation. J. Virol. 81: 3087–3096. [Medline] [CrossRef]

2. Chen, Z., Rijnbrand, R., Jangra, R. K., Devaraj, S. G., Qu, L., Ma, Y., Lemon, S. M. and Li, K. 2007. Ubiquitination and proteasomal degradation of interferon regulatory factor-3 induced by Npro from a cytopathic bovine viral diarrhea virus. Virology 366: 277–292. [Medline] [CrossRef]

3. Collett, M. S., Larson, R., Belzer, S. K. and Retzel, E. 1988. Proteins encoded by bovine viral diarrhea virus: the genomic organization of a pestivirus. Virology 165: 200–208. [Medline] [CrossRef]
of hog cholera virus by its effect on Newcastle disease virus in swine tissue culture. Science 128: 366. [Medline] [CrossRef]

16. Meyers, G. and Thiel, H. 1996. Molecular characterization of pestiviruses. Adv Virus Res. 47: 53–118. [Medline] [CrossRef]

17. Muhsen, M., Aoki, H., Ikeda, H. and Fukusho, A. 2013. Biological properties of bovine viral diarrhea virus quasispecies detected in the RK13 cell line. Arch Virol. 158: 753–763. [Medline] [CrossRef]

18. Nakamura, S., Fukusho, A., Inoue, Y., Sasaki, H. and Ogawa, N. 1993. Isolation of different non-cytopathogenic bovine viral diarrhoea (BVD) viruses from cytopathogenic BVD virus stocks using reverse plaque formation method. Vet Microbiol. 38: 173–179. [Medline] [CrossRef]

19. Nakamura, S., Shimazaki, T., Sakamoto, K., Fukusho, A., Inoue, Y. and Ogawa, N. 1995. Enhanced replication of orbiviruses in bovine testis cells infected with bovine viral diarrhoea virus. J Vet Med Sci. 57: 677–681. [Medline] [CrossRef]

20. Tamura, T., Nagashima, N., Ruggli, N., Summerfield, A., Kida, H. and Choi, K. H. 2009. Zinc binding in pestivirus Npro is required for interferon regulatory factor 3 interaction and degradation. J Virol. 83: 817–829. [Medline] [CrossRef]

21. Nishine, K., Aoki, H., Sakoda, Y. and Fukusho, A. 2014. Field distribution of END phenomenon-negative bovine viral diarrhea virus. J Vet Med Sci. 76: 1635–1639. [Medline] [CrossRef]

22. Rasmussen, T. B., Reimann, I., Uttenenthal, A., Leifer, I., Depner, K., Schirrmeyer, H. and Beer, M. 2010. Generation of recombinant pestiviruses using a full-genome amplification strategy. Vet Microbiol. 142: 13–17. [Medline] [CrossRef]

23. Ruggli, N., Tratschin, J. D., Mittelholzer, C. and Hofmann, M. A. 1996. Nucleotide sequence of classical swine fever virus strain Alfort/187 and transcription of infectious RNA from stably cloned full-length cDNA. J Virol. 70: 3478–3487. [Medline] [CrossRef]

24. Ruggli, N., Summerfield, A., Fiebach, A. R., Guizzly-Piriou, L., Bauhofer, O., Lamm, C. G., Waltersperger, S., Matsuno, K., Liu, L., Gerber, M., Choi, K. H., Hofmann, M. A., Sakoda, Y. and Tratschin, J. D. 2009. Classical swine fever virus can remain virulent after specific elimination of the interferon regulatory factor 3-degrading function of N\(^{\text{pro}}\). J Virol. 83: 817–829. [Medline] [CrossRef]

25. Rümenapf, T., Unger, G., Strauss, J. H. and Thiel, H. J. 1993. Processing of the envelope glycoproteins of pestiviruses. J Virol. 67: 3288–3294. [Medline] [CrossRef]

26. Rümenapf, T., Stark, R., Heimann, M. and Thiel, H. J. 1998. N-terminal protease of pestiviruses: identification of putative catalytic residues by site-directed mutagenesis. J Virol. 72: 2544–2547. [Medline] [CrossRef]

27. Sakoda, Y. and Fukusho, A. 1998. Establishment and characterization of a porcine kidney cell line, FS-L3, which forms unique multicellular domes in serum-free culture. In Vitro Cell. Dev. Biol. Anim. 34: 53–57. [Medline] [CrossRef]

28. Seago, J., Hilton, L., Reid, E., Doceul, V., Jeyatheesan, J., Moosneradj, K., McCauley, J., Charleston, B. and Goodbourn, S. 2007. The N\(^{\text{pro}}\) product of classical swine fever virus and bovine viral diarrhea virus uses a conserved mechanism to target interferon regulatory factor-3. J. Gen. Virol. 88: 3002–3006. [Medline] [CrossRef]

29. Shimizu, Y., Furuuchi, S., Kumagai, T. and Sasahara, J. 1970. A mutant of hog cholera virus inducing interference in swine testicle cell cultures. Am J Vet Res. 31: 1787–1794. [Medline] [CrossRef]

30. Simmonds, P., Becher, P., Collett, M. S., Gould, E. A., Heinz, F. X., Meyers, G., Monath, T., Pletnev, A., Rice, C. M., Stiasny, K., Thiel, H.-J., Weiner, A. and Bukh, J. 2012. Family Flaviviridae. pp. 1003–1020. In: Virus Taxonomy. Ninth Report of the International Committee on Taxonomy of Viruses (King, A. M. Q., Adams, M. J., Carstens, E. B. and Lefkowitz, E. J. eds.). Elsevier Academic Press, San Diego.

31. Stark, R., Meyers, G., Rümenapf, T. and Thiel, H. J. 1993. Processing of pestivirus polyprotein: cleavage site between autoprotease and nucleocapsid protein of classical swine fever virus. J Virol. 67: 7088–7095. [Medline] [CrossRef]

32. Szymanski, M. R., Fiebach, A. R., Tratschin, J. D., Gut, M., Ramanujam, V. M., Gottipati, K., Patel, P., Ye, M., Ruggli, N. and Choi, K. H. 2009. Zinc binding in pestivirus N\(^{\text{pro}}\) is required for interferon regulatory factor 3 interaction and degradation. J Mol Biol. 391: 438–449. [Medline] [CrossRef]

33. Tamura, T., Nagashima, N., Ruggli, N., Summerfield, A., Kida, H. and Nakamura, S. 2014. N\(^{\text{pro}}\) of classical swine fever virus contributes to pathogenicity in pigs by preventing type I interferon induction at local replication sites. Vet Res. 45: 47. [Medline] [CrossRef]

34. Tamura, T., Sakoda, Y., Yoshino, F., Nomura, T., Yamamoto, N., Sato, Y., Okamatsu, M., Ruggli, N. and Kida, H. 2012. Selection of classical swine fever virus with enhanced pathogenicity reveals synergistic virulence determinants in E2 and NS4B. J Virol. 86: 8602–8613. [Medline] [CrossRef]

35. Tautz, N., Elbers, K., Stoll, D., Meyers, G. and Thiel, H. J. 1997. Serine protease of pestiviruses: determination of cleavage sites. J Virol. 71: 5415–5422. [Medline] [CrossRef]