Emergence of Pathogenic Coronaviruses in Cats by Homologous Recombination between Feline and Canine Coronaviruses

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Abstract

Type II feline coronavirus (FCoV) emerged via double recombination between type I FCoV and type II canine coronavirus (CCoV). In this study, two type I FCoVs, three type II FCoVs and ten type II CCoVs were genetically compared. The results showed that three Japanese type II FCoVs, M91-267, KUK-H/L and Tokyo/cat/130627, also emerged by homologous recombination between type I FCoV and type II CCoV and their parent viruses were genetically different from one another. In addition, the 3′-terminal recombination sites of M91-267, KUK-H/L and Tokyo/cat/130627 were different from one another within the genes encoding membrane and spike proteins, and the 5′-terminal recombination sites were also located at different regions of ORF1. These results indicate that at least three Japanese type II FCoVs emerged independently. Sera from a cat experimentally infected with type I FCoV was unable to neutralize type II CCoV infection, indicating that cats persistently infected with type I FCoV may be superinfected with type II CCoV. Our previous study reported that few Japanese cats have antibody against type II FCoV. All of these observations suggest that type II FCoV emerged inside the cat body and is unable to readily spread among cats, indicating that these recombination events for emergence of pathogenic coronaviruses occur frequently.

Introduction

Coronaviruses (CoVs) (order Nidovirales, family Coronaviridae, subfamily Coronavirinae) are enveloped and have large single-stranded, positive-sense RNA. Most CoVs cause enteric and/or respiratory diseases in mammals and birds. The 5′ two-thirds of the CoV genome consists of two overlapping open reading frames (ORFs 1a and 1b) that encode a non-structural polyprotein, including RNA-dependent RNA polymerase (RdRp). The other third of the genome consists of ORFs encoding structural proteins, spike (S), membrane (M), envelope (E) and the nucleocapsid (N), and some non-structural proteins (nsps) 3a, 3b, 3c, 7a and 7b [1]. Transcription regulatory sequences (TRS) are located at 5′-distal position in each mRNA and play an important role in the RNA replication of CoV [2], [3].

CoVs frequently undergo mutation and recombination, and there are three reasons for this [4]. First, CoV RdRp has low fidelity. Although CoV encodes nsp14, which possesses 3′→5′ exonuclease activity for proofreading, the mutation rate approaches 2.0×10⁻⁶ mutations per site per round of replication [5]. Second, there is a unique RNA replication mechanism using the TRS motif that is known as the “copy choice” mechanism, which induces homologous RNA recombination in CoVs [3], [6]. Third, CoV possesses the largest genome (26–32 kb) among RNA viruses. Furthermore, heterologous recombination that Betacoronavirus subgroup A has the hemagglutinin esterase gene originated from influenza C virus [7], [8]. These mutation and/or recombination events change viral properties, host range and pathogenicity.

Feline CoV (FCoV) is classified into genus Alphacoronavirus, species Alphacoronavirus 1, and includes canine CoV (CCoV), transmissible gastroenteritis virus (TGEV) and porcine respiratory CoV (PRCoV). FCoV is distributed worldwide in cats and mainly induces enteric disease is known as feline enteric coronavirus (FECV). On the other hand, cats infected with FCoV rarely develop the more severe disease feline infectious peritonitis (FIP), which is caused by a mutant virus that is referred to as FIP virus (FIPV). In addition, FCoVs can be divided into two serotypes, types I and II, based on antigenicity [10–12]. These serotypes differ primarily in growth characteristics in cell culture and in receptor usage. Type II FCoV
is able to use feline aminopeptidase N (fAPN) as its receptor, but type I FCoV cannot [9], [13]. Recently, it was revealed that the S protein was solely responsible for the differences in types I and II FCoV with regard to growth characteristics in cell culture and fAPN usage [14].

CCoV was first isolated in 1971 from dogs with moderate to severe enteritis in Germany [15]. CCoV is widespread in the dog population and is one of the most important canine enteropathogens [16–22]. CCoVs were also divided into two genotypes: I and II. Before 2000, it was thought that CCoV had only one genotype, but strain Elmo/02 with a type I FCoV-like S gene was detected in Italy [23]. The Elmo/02 strain possessed a novel ORF3 gene that was absent from other *Alphacoronavirus* I between the S and ORF3a genes [24]. Finally, this type I FCoV-like-CCoV was designated type I CCoV and the reference CCoV was designated type II CCoV. Surprisingly, 36.9%–76.8% of dogs with diarrhea were co-infected with both types I and II CCoV [25–27]. Furthermore, type II CCoV was divided into two subtypes, Ila and IIb [28]. In type II CCoV, the 5’-terminal region of the S gene was similar to that of TGEV and it was thought that type IIb CCoV emerged via recombination between type Ila CCoV and TGEV [28]. Recently, a type Ila CCoV strain CB/05 with high virulence was reported in Europe [29]. CB/05-infected pups showed clinical signs such as lethargy, vomiting, diarrhea and acute lymphopenia, and the viral genome was observed in extraintestinal tissues including brain [29], [30]. Furthermore, immune response induced by enteric CCoV did not protect dogs from infection with CB/05 [31]. However, there is little genetic information on CCoV in Japan.

In this study, to clarify the mechanisms of emergence of type II FCoV, three type II FCoVs isolated in Japan were genetically and antigenetically compared with ten Japanese type II CCoVs and three type II FCoVs isolated in Japan. All efforts were made to minimize pain and suffering.

**Materials and Methods**

All animal procedures were conducted according to the Yamaguchi University Animal Care and Use guidelines and were approved by the Institutional Animal Care and Use Committee of Yamaguchi University. All efforts were made to minimize pain and suffering.

**Cells**

*Felis catus* whole fetus-4 cells (fcwf-4 cells; ATCC Number: CRL-2787) [32] were grown in Dulbecco’s modified Eagle’s medium (DMEM; Life Technologies, Carlsbad, CA) containing 10% fetal calf serum (FCS), 100 U/ml penicillin and 100 μg/ml streptomycin (Life Technologies). Cells were maintained in a humidified 5% CO₂ incubator at 37°C.

**Viruses**

Type I FCoV strains C3663 and Yayoi, type II FCoV strains M91-267, KUK-H/L and Tokyo/cat/130627 and type II CCoV strains fc1, fc4, fc7, fc9, fc76, fh100, fh94-039, fh97-022, fh00-089 and fh00-016 were analyzed in this study (Table 1). Type I and II FCoVs, excluding Tokyo/cat/130627, were characterized by indirect fluorescence assay (IFA) using monoclonal antibodies (MAbs) that were kindly provided by Dr. Hohdatsu [11], [33]. Yayoi strain was isolated from a cat with a non-erosive form of FIP in Tokyo by serial passage in suckling mouse brain, and was then adapted to fcwf-4 cells [34]. C3663 strain was isolated from a cat with an erosive form of FIP in Tokyo by serial passage in suckling mouse brain, and was then adapted to fcwf-4 cells [34]. C3663 strain was isolated from a cat with an erosive form of FIP in Kagoshima in 1994 [35]. The pathogenicity of C3663 and Yayoi in cats was characterized [36]. M91-267 strain was isolated from a cat with an erosive form of FIP in Miyazaki in 1991 [33]. Three SPF cats were experimentally infected with M91-267, and all of these died from FIP (unpublished data). KUK-H strain was isolated from a cat with an erosive form of FIP in Kagoshima in 1907, and KUK-H/L that formed large plaques was plaque-purified from the KUK-H strain [35]. KUK-H/L caused lethal FIP in cats [35]. RNA sequences of Tokyo/cat/130627 were obtained from FIP ascites in a cat in Tokyo in 2013. The FIPV spread quickly in a cattery, and more than twenty cats developed FIP. Type II CCoV strains, fc1, fc4, fc7, fc9, fc76, fh100, fh94-039 and fh97-022, were isolated between 1990 and 1997 in Japan [19], and fh00-016 and fh00-087 were isolated in 2000 in Japan [37].

**Reverse transcription (RT)-polymerase chain reaction (PCR)**

Each virus, excluding Tokyo/cat/130627, was inoculated onto an fcwf-4 cell monolayer and was incubated until cytopathic effects (CPEs) were observed. RNA was then extracted from fcwf-4 cells using an RNase Mini kit (Qiagen, Hilden, Germany) and RT reaction was carried out at 30°C for 10 min, 42°C for 30 min, 70°C for 15 min and 5°C for 5 min with random 9-mer oligonucleotide primers or 42°C for 30 min, 70°C for 15 min and 5°C for 5 min with oligo dT-adaptor primer using a TaKaRa RNA LA PCR kit (AMV) Ver.1.1 (TaKaRa, Shiga, Japan).

For amplification of partial S genes of type II CCoVs and type II FCoVs, primers CCVSF (5’-AGCACTTTCCTATT-GATTG-3’) and CCVSR (5’-GCTTATTGTTGCATAATACACCAACACC-3’) were used [38]. For amplification of the N gene, primers NF (5’-CTAAAGCTGGGATCTAGTCAACAG-3’) and NR (5’-TAATAAATAAGGCTTTGAGAAAC-3’) were used [39]. PCR was carried out at 94°C for 2 min, followed by 40 cycles at 94°C for 30 s, 55°C for 30 s, 72°C for 2 min and final extension at 72°C for 10 min using a TaKaRa RNA LA PCR kit (AMV) Ver.1.1 (TaKaRa). PCR products were analyzed electrophoretically and amplified products were purified using a QIAquick PCR Purification kit (Qiagen) for sequence analysis.

In order to amplify the subgenomic mRNA of CCoV Vc1, PCR was performed using 52F (5’-ACTAGGCTTTGCTAGATTATT-3’) as a forward primer and CCVSF (5’-CCAGTTTATTATAAAGCCTG-3’) and NR2 (5’-GGCCAATAACGCTGCCAA-3’) and M13 primer M4 as reverse primers. Primer 52F and NR2 recognized the TRS conserved among *Alphacoronavirus* sequences [36]. The reaction was carried out under the same conditions as mentioned above.

For sequence analysis of ORFs M, N, 7a and 7b of M91-267 and KUK-H/L, we carried out TA cloning. RNA was extracted from fcwf-4 cells infected with M91-267 or KUK-H/L using an RNase Mini kit (Qiagen). Extracted RNA was reverse-transcribed with oligo dT-Adaptor primer using a TaKaRa RNA LA PCR kit (AMV) Ver.1.1 (TaKaRa). PCR products were analyzed electrophoretically and amplified products were purified using a QIAquick PCR Purification kit (Qiagen) for sequence analysis.

Viral RNA of Tokyo/cat/130627 was extracted from FIP ascites in a cat using a QIAamp Viral RNA Mini Kit (Qiagen). For sequence analysis, five fragments of the Tokyo/cat/130627 gene between the 3’-terminus of ORF 1b and poly A were amplified using the following primer pairs: 1bF (5’-TTGATTTGAA-GATTGATTTGGG-3’)-CCVSF; CCVSF-S2cenFR3 (5’-
Table 1. Canine and feline coronaviruses analyzed in this study: nucleotide sequence acquisition numbers and serum cross-neutralizing activity.

| Virus     | Strain | Accession No.         | RdRp       | S - poly A         | Partial S   | N            | VN titers | Type I<sup>a</sup> | Type II<sup>b</sup> |
|-----------|--------|-----------------------|------------|--------------------|-------------|--------------|-----------|---------------------|--------------------|
| Type II CCoV | fc1    | AB781791              | AB781790   | AB781790           | AB781790    | <1:10        | 1:400      |
|           | fc4    | AB907625              | AB781807   | AB781797           | AB781798    | <1:10        | 1:4525     |
|           | fc7    | AB907626              | AB781808   | AB781798           | AB781800    | <1:10        | 1:1600     |
|           | fc9    | AB907627              | AB781809   | AB781799           | AB781800    | <1:10        | 1:1600     |
|           | fc76   | AB907628              | AB781810   | AB781800           | AB781801    | <1:10        | 1:1131     |
|           | fc100  | AB907629              | AB781811   | AB781801           | AB781802    | <1:10        | 1:2263     |
|           | fc97-022 | AB907631             | AB781812   | AB781802           | AB781803    | <1:10        | 1:2325     |
|           | fc94-039 | AB907630             | AB781813   | AB781803           | AB781804    | <1:10        | 1:1600     |
|           | fc00-016 | AB907632             | AB781814   | AB781804           | AB781805    | <1:10        | 1:1600     |
|           | fc00-089 | AB907633             | AB781815   | AB781805           | AB781806    | <1:10        | 1:200      |
| Type II FCoV | M91-267  | AB781792              | AB781788   | AB781788           | AB781788    | <1:10        | 1:25000    |
|           | KUK-H/L  | AB781793              | AB781789   | AB781789           | AB781789    | <1:10        | 1:6400     |
|           | Tokyo/cat/130627    | AB907634              | AB907624   | AB907624           | AB907624    | N.D.        | N.D.       |
| Type I FCoV | C3663    | AB781794              | AB535528   | AB535528           | AB535528    | 1:6400       | 1:80       |
|           | Yayoi    | AB781795              | AB695067   | AB781806           | AB781806    | 1:2000       | 1:160      |

N.D.: Not done.
<sup>a</sup>Serum was collected from the cat that was inoculated intraorally with type I FCoV C3663 [36].
<sup>b</sup>Serum was collected from the cat that was inoculated intraperitoneally with type II FCoV M91-267 (unpublished data).
<sup>c</sup>[36].

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GTGTCAATTCCAGTTACAG-3’; S2revFF2 (5’-GAGTGCT-GATGCAACAAGT-3’); N-RR3 (5’-GCCACCAGTCAATGC-3’); N-RF4 (5’-AGTTAGATGGCTGCTGCT-3’); N4 (5’-CATCTCAACCTGTGGTTCAT-3’); and N1 (5’-MMAAY- AAACACACCTGGAAG-3’)-oligo dT-Adaptor primer. RT-PCR was carried out using a QIAGEN OneStep RT-PCR Kit (Qiagen) according to the manufacturer’s instructions. Reactions were carried out at 45°C for 45 min and 95°C for 15 min, followed by 40 cycles at 94°C for 10 s, 55°C for 30 s, 68°C for 3 min, and a final extension at 68°C for 15 min. For amplification of partial RdRp genes, primer IN-6 (5’-GGTTGGGACTATCTAAGTGTGA-3’) and IN-7 (5’-CCATCATCAGATAGAATCATCAT-3’) were used as described previously [40]. PCR products were analyzed electrophoretically and amplified products were purified using a MinElute PCR Purification Kit (Qiagen) for sequence analysis.

Nucleotide sequences
Sequencing was performed using a BigDye Terminator v3.1 Cycle Sequencing kit (Life Technologies) according to the manufacturer’s instructions. Products were purified by ethanol precipitation and analyzed using an ABI PRISM 310 Genetic Analyzer (Life Technologies). For sequence analysis, primers shown in Table S1 were used and nucleotide sequences were deposited to the DNA database of Japan (DDBJ) under the accession numbers listed in Table 1.

Homology search and phylogenetic analysis
Homologies among strains were analyzed using GENETYX Ver.8 (GENETYX Corporation, Tokyo, Japan) and phylogenetic trees were constructed by the neighbor-joining method [41] using MEGA5.0 software [42] based on nucleotide pairwise distance.

For construction of the phylogenetic tree, we referred to the following sequences; type II FCoV 79-1146 (accession no. DQ109021), 79-1683 (JN634064), DF-2 (JQ408981) and NTU156/F/2007 (GQ152141), type I FCoV C3663 (AB535526), Yayoi (AB695067 for S), UCD1 (AB080222 for S, AB080692 for N), Black (EU166072), NTU2/R/2003 (DQ160294), RM (FJ93051), UCD11a (FJ917519), UCD5 (FJ917522), UCD12 (FJ917521), UCD13 (FJ917529), UCD14 (FJ917524), U2 (FJ930606), U16 (FJ930559), U18 (HQ012360), U20 (HQ992471), U 21 (HQ012969), U23 (GU553362), type II CoCV 1-71 (JQ404409), v1 (AY39024 for S, AY390345 for N), K378 (KC175340), NTU396/F/2008 (GQ477367), 3821 (AB017709 for S), TGEV Purdue (DQ11789), and PRCoV ISU-1 (DQ11789). Analysis of the similarity in the 3’-region of the genome, excluding poly A, was carried out using Simplot version 3.5.1 [43].

Sera from cats
Sera collected from two SPF cats experimentally inoculated with FIPVs were used. One cat was inoculated intra-orally with type I FCoV C3663 (3.9×10^6 PFU/cat) and showed an effusion form of FIP [36]. Another cat was inoculated intraperitoneally with type II FCoV M91-267 (1.0×10^6 PFU/cat) and also showed an effusive form of FIP (unpublished data). When clinical symptoms were severe, cats were euthanized under anesthesia. These sera were obtained in our previous experiments carried out under approval by the ethics committee for animal experiments, Faculty of Agriculture, Yamaguchi University.

Virus-neutralization test
Virus-neutralization (VN) test was performed by 75% plaque-reduction neutralization test (PRNT75) using cat sera inactivated at
56°C for 30 min [12], [36]. Equal volumes of two-fold serially diluted sera and 2.0 × 10^6 PFU/ml virus were mixed and incubated at 37°C for 1 h. Then, 50 μl of this mixture was inoculated onto an fcwf-4 cell monolayer in 24-well plates (Sumitomo Bakelite, Tokyo, Japan). After adsorption at 37°C for 1 h, inoculum was removed and 0.8% agarose (Seaplaque GTG Agarose; Lonza, Switzerland) in DMEM containing 10% FCS was overlaid. Infected cells were incubated at 37°C until CPE was observed, followed by fixing with phosphate-buffered formalin and staining with crystal violet.

Results

Comparison of 3’-region among type II CCoVs, and type I and II FCoVs

Nucleotide sequences of the 3’-region of the genomes, excluding the poly A, of type II CCoV fc1 (8,959b) and type II FCoVs, M91-267 (8,889b), KUK-H/L (8,930b) and Tokyo/cat/130627 (8,831b), were determined (DDBJ) Accession No. AB781790 for fc1, AB781790 for M91-267, AB781790 for KUK-H/L and AB907624 for Tokyo/cat/130627 [Table 1]. Because of a mutation in the start codon (ATG→ACG), Tokyo/cat/130627 lacked ORF3b. In addition, type II FCoVs, M91-267 and Tokyo/cat/130627 possessed a truncated ORF 3c (Fig. 1). When compared with KUK-H/L, M91-267 had a 25-nucleotide deletion in the ORF 3c gene, resulting in a truncated ORF 3c. In comparison with C3663, Tokyo/cat/130627 showed a 25-nucleotide deletion in the ORF 3c gene, resulting in a truncated ORF 3c gene. Deduced amino acid sequences for ORFs S, 3a, 3b, 3c, E, M, N, 7a and 7b in type II FCoVs were compared with those of type I FCoV C3663 and type II CCoV fc1 [Table S2, S3]. Both M91-267 and KUK-H/L showed low identities with type I FCoV C3663 in ORFs S, 3a, 3b, 3c and E and high identities in ORFs N, 7a and 7b (Table S2). In contrast, the two strains showed high identities with type II CCoV fc1 in ORFs S, 3a, 3b, 3c and E and low identities in ORFs N, 7a and 7b (Table S3). In ORF M, the identities among type I FCoV, type II FCoV and CCoV were neither high nor low (Table S2, S3). Interestingly, comparison between Tokyo/cat/130627 and type I FCoV showed low identities in ORF S and high identities in ORFs S, 3a, 3c, E, M, N, 7a and 7b, while comparison with type II CCoV fc1 showed high identity only in ORF S and low identities in ORF 3a, 3c, E, M, N, 7a and 7b (Table S2, S3).

Figure 2. Phylogenetic trees using partial RdRp(A), partial S (B) and N (C) genes. Type I FCoVs, type II FCoVs and type II CCoVs are shown in red, green and blue, respectively. Swine CoV (TGEV and PRCoV), ferret CoV (FRCoV) and human CoV (HCoV) are shown in black. GenBank accession numbers are shown in parentheses.

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Comparison of partial RdRp genes among type II CCoVs and type I and II FCoVs

Nucleotide sequences of partial RdRp gene in ORF1b (394b) of 15 Japanese CoVs were determined and deduced amino acid sequences were compared (Tables S2, S3, S4 and Fig. 2A). In comparison with type I FCoVs, C3663, KUK-H/L and Tokyo/cat/130627 showed higher identity in RdRp than M91-267 (Table S2). On the other hand, the sequence of RdRp of M91-267 was more similar to that of type II CCoV fc1 than type I FCoV C3663 (Table S3). All CCoV strains possessed high homology with fc1 strain and M91-267, but showed low homology with KUK-H/L and Tokyo/cat/130627 (Table S4).

Phylogenetic analysis using partial RdRp genes showed that Japanese type II strains could be divided into two different groups; feline CoV and canine CoV (Fig. 2A). KUK-H/L and Tokyo/cat/130627 belonged to feline CoV group and M91-267 belonged to canine CoV group. The other foreign type II FCoVs belonged to the type II CCoV group.

Comparison of partial S genes among type II CCoVs and type I and II FCoVs

Nucleotide sequences of partial S genes (692b) of 15 Japanese CoVs were determined and deduced amino acid sequences were compared (Tables S5 and Fig. 2B). In comparison with type I FCoV C3663, all type II FCoVs showed low identity. All CCoV strains possessed high homology with fc1 strain, but showed low homology with types I and II FCoV (Table S6).

Phylogenetic analysis using partial S genes showed that all type II FCoVs were more similar to Japanese type II CCoV than type II FCoVs and type II CCoVs from other countries. In addition, Japanese FCoVs belonged to different subgroups; KUK-H/L belongs to a cluster with fc1. M91-267 belongs to the other cluster with fc76 and fc94-039. Tokyo/cat/130627 belongs to the cluster with Taiwanese strain NTU156/P/2007 (Fig. 2B).

Comparison of N genes among type II CCoVs and type I and II FCoVs

Nucleotide sequences of N genes (1149b) of 15 Japanese CoVs were determined and deduced amino acid sequences were compared (Table S2, S3, S6 and Fig. 2C). In comparison with type I FCoV C3663, all type II FCoVs showed higher identity than type II CCoV fc1 (Table S2). All CCoV strains possessed high homology with fc1 strain, but showed low homology with types I and II FCoV (Table S6).

Phylogenetic analysis using N genes showed that FCoV strains and type II CCoV strains were genetically divided into different groups. In the feline CoV group, Japanese type II FCoVs M91-267, KUK-H/L and Tokyo/cat/130627 belonged to feline CoV group and M91-267 belonged to canine CoV group. The other foreign type II FCoVs belonged to the type II CCoV group.

Recombination sites of type II FCoVs

Simplot analysis showed that the similarity of Tokyo/cat/130627 to CCoV fc1 changed at the 3’-terminal region of the S gene (Fig. 2B). Furthermore, Japanese type II FCoVs were more similar to Japanese type II CCoV than type II FCoVs and type II CCoVs from other countries. In addition, Japanese FCoVs belonged to different subgroups; KUK-H/L belongs to a cluster with fc1. M91-267 belongs to the other cluster with fc76 and fc94-039. Tokyo/cat/130627 belongs to the cluster with Taiwanese strain NTU156/P/2007 (Fig. 2B).
gene, and those of M91-267 and KUK-H/L changed within the M gene (Fig. 3).

The M genes were compared among types I and II FCoV and type II CCoV (Fig. 4A). The alignment showed that the 5'-terminal region of the M genes of M91-267 and KUK-H/L was similar to that of CCoV fc1, but the 3'-terminal region was similar to type I FCoV C3663 (Fig. 4A). The M gene of Tokyo/cat/130627 was similar to type I FCoV C3663. Furthermore, the nucleotide sequences indicated that the recombination sites of these two viruses, M91-267 and KUK-H/L, were different. Among these CoVs, two conserved regions were located at 133–177 and 325–366 in the M gene. KUK-H/L was similar to type II CCoV upstream of the first conserved region (region 133–177), but was similar to type I FCoV downstream of the region. On the other hand, M91-267 was similar to type II CCoV upstream of the second conserved region (region 325–366), and was similar to type I FCoV downstream of the region.

The alignment data using type I FCoV C3663, type II FCoV M91-267, KUK-H/L and Tokyo/cat/130627, and type II CCoV fc1 showed that the recombination site of Tokyo/cat/130627 was in the 3'-terminal of the S gene. Among these FCoVs and CCoVs, region 4183–4202 of the S gene was completely conserved (Fig. 4B). Upstream of the conserved region, Tokyo/cat/130627 was more similar to type II CCoV fc1 than type I FCoV C3663, and downstream of the conserved region, Tokyo/cat/130627 was more similar to type I FCoV C3663 (Fig. 4B).

Cross-neutralization activity to CCoV by sera collected from cats infected with FCoV

In order to examine whether cats with VN antibody against type I FCoV can be infected with type II CCoV, cross-neutralizing activity of sera from cats experimentally infected with FCoVs was examined (Table 1). Cat serum against type I FCoV C3663 was able to neutralize infection of type I FCoV strains C3663 and Yayoi (1:6400 and 1:2000, respectively), but not those of type II CCoV and type II FCoV (less than 1:10) (Table 1). On the other hand, cat serum against type II FCoV M91-267 was able to...
It is interesting that all FCoVs, both types I and II, possessed the 5'- and 3'-termini of the FCoV genome, but not CCoV. These regions may be essential for growth of FCoV in cats and double recombination may be required to maintain both the 5'- and 3'-termini of FCoV. Type II FCoV possessed two types of RdRp element derived from type I FCoV or type II CCoV (Fig. 2A), suggesting both types of RdRp were able to function during replication and transcription in cat body. Furthermore, the region upstream of RdRp might be essential for FCoV infection in cats. In addition, it has been reported that N protein is important for viral particle production [52], and the N gene is conserved among FCoVs. Therefore, the N protein of FCoV, but not CCoV, may be essential for replication of FCoV in cats. Interestingly, simplot analysis showed four other candidate recombination sites, one in the 3c gene, two in the N gene and one in the 7a gene, which showed high identity between CCoV and type I FCoV (Fig. 3). If the M or N genes of type I FCoV are not necessary for growth of FCoV in cats, other recombinant type II FCoVs using these possible recombination sites must occur. Further analysis of type II FCoV is necessary to clarify the recombination events of CoV in cats.

Four full genome sequences of type II FCoVs (79–1146, 79–1683, DF-2 and NTU156/P/2007) are deposited in GenBank. We also reported one-third of the full genome of three type II FCoV strains (M91-267, KUK-H/L and Tokyo/cat/130627) and one type II CCoV fc1. Six of seven type II FCoV strains emerged by recombination events at the E or M gene. However, the recombination event of Tokyo/cat/130627 occurred at the 3'-terminal of the S gene. The nucleotide sequences indicated that M91-267, KUK-H/L and Tokyo/cat/130627 originated from type I FCoV strains similar to C3663, Yayoi and NTU2/R/2003, respectively, and that the central region, including the S gene, was acquired from type II CCoV strains similar to fc94-039, fc1 and fc00-089, respectively. In addition, the recombination sites were clearly different (Fig. 3 and 4A, B). These results indicate that the recombination events between type I FCoV and type II CCoV occurred independently. In addition, original viruses of foreign type II FCoVs, 79-1146, 79-1683 and NTU156/P/2007 differed from those of these three Japanese type II FCoVs, indicating that the recombination events occurred among cat populations all over the world.

Sera from cats experimentally inoculated with type I FCoV C3663 could not neutralize type II CCoV infection (Table 1), thus suggesting that the cat infected with type I FCoV could not prevent type II FCoV infection. On the other hand, the cat infected with type II FCoV could neutralize type I FCoV infection (Table 1). In addition, many sera from type II FCoV-infected cats in the outbreak could cross-neutralize type I FCoV infection, and those from type I FCoV-infected cats in the field could not cross-neutralize type II FCoV infection (our unpublished data). These results suggest that the cross-reactivity to type I FCoV in type II FCoV-infected cats might be induced by viral proteins other than S protein. Further analysis will be required to clarify the cross VN activity in type II FCoV-infected cats.

Type II CCoV was able to use “feline” aminopeptidase N as a receptor to infect cats [53], [54] and type I FCoV-infected cats did not possess VN antibody against type II CCoV infection (Table 1), indicating that cats infected with type I FCoV could be superinfection with type II CCoV from dogs. Our hypothesis on the mechanism of emergence of type II FCoV is shown in Fig. 5. Cats infected with type I FCoV were unable to produce VN antibody against type II CCoV. Hence, cats had the possibility of superinfection with type II CCoV. The recombination event between type I FCoV and type II CCoV occurred inside the cat body, leading to emergence of type II FCoV.

Discussion

Type II FCoV emerged as a result of recombination events between type I FCoV and type II CCoV [44], [45]. Recently, one additional full genome sequence of type II FCoV NTU156/P/2007 was determined, and this facilitated understanding of the mechanisms responsible for emergence of type II FCoV [46]. The prevalence of type II FCoV in the cat population is lower than that of type I, but the reasons for this remain uncertain [12], [47–51]. In this study, numerous FCoV and CCoV isolates from Japan were genetically characterized, and the emergence of type II was discussed.

Our phylogenetic and sequence analysis clearly indicated that type II FCoVs emerged by different recombination events between type I FCoV and type II CCoV. In addition, other type II FCoVs isolated from the USA (79–1683 and 79–1146) and Chinese Taipei (NTU156/P/2007) also showed different origins [45], [46]. These results indicate that type II FCoV independently emerged in different cats and did not spread very easily. Our previous study also showed that many cats possess VN antibody to type I FCoV, but few cats in Japan possess VN antibody against type II FCoV [12], supporting the notion that type II FCoV does not readily spread among the cat population. The reasons why type II FCoV is unable to spread among the cat population are unclear.

Two of three stains of Japanese type II FCoV, M91-267 and Tokyo/cat/130627, possessed the truncated ORF 3c gene (Fig. 1). An intact 3c gene is apparently essential for efficient replication of FCoV in the intestinal tract, resulting in the secretion of FCoV from feces and transmission of FCoV among cat populations [55], [56]. On the other hand, many FIPV possessed truncated 3c gene and cats with FIP did not excrete virus in feces [57–59]. Furthermore, one strain of type II FIPV with intact ORF 3c gene occurred in Taiwan. In early stages of the outbreak, the type II FIPV possessed intact 3c gene, but lost it in later stages [60]. Therefore, these type II FCoVs with truncated 3c gene, M91-267 and Tokyo/cat/130627, might not spread well among cats.
CoVs, such as SARS-CoV, tend to change their host range by mutation and/or recombination [61]. Homologous recombination is a significant factor for change of host range. Therefore, investigations into homologous recombination of CoVs may help to clarify the mechanisms responsible for changes in host range.

Supporting Information

Table S1 Primers used in this study.

Table S2 Comparison of ORF identities between C3663 and other coronaviruses.

Table S3 Comparison of ORF identities between f1 and other coronaviruses.

References

1. Woo PC, Huang Y, Lau SK, Yuen KY (2010) Coronavirus genomics and bioinformatics analysis. Viruses 2: 1804–1820.
2. Makino S, Joo M, Makino JK (1991) A system for study of coronavirus mRNA synthesis: a regulated, expressed subgenomic defective interfering RNA results from intergenic site insertion. J Virol 65: 6031–6041.
3. Pasternak AO, Spaan WJ, Suijder EJ (2006) Nidovirus transcription: how to make sense... J Gen Virol 87: 1403–1421.
4. Böllis M, Donaldson E, Baric R (2011) SARS-CoV and Emergent Coronavirus: Viral Determinants of Interspecies Transmission.Curr Opin Virol 1: 624–634.
5. Eckerle LD, Becker MM, Halpin RA, Li K, Venter E, et al. (2010) Infidelity of SARS-CoV Nsp14-exonuclease mutant virus replication is revealed by complete genome sequencing. PLoS Pathog 6: e1000966.
6. Lai MM, Baric RS, Makino S, Keck JG, Egbert J, et al. (1985) Recombination between nonsegmented RNA genomes of murine coronaviruses. J Virol 56: 449–456.
7. Luytjes W, Bredenkamp PJ, Noten AF, Horzinek MC, Spaan WJ (1988) Sequence of mouse hepatitis virus A59 mRNA 2: indications for RNA recombination between coronaviruses and influenza virus. Virology 160: 415–422.
8. Zeng Q, Langereis MA, van Vliet AL, Huizinga EG, et al. (2008) Structure of coronavirus hemagglutinin-esterase offers insight into corona and influenza virus evolution. Proc Natl Acad Sci U S A 105: 9065–9069.
9. Pedersen NC, Evermann JF, McKeirnan AJ, Ott RL (1984) Pathogenicity of canine coronavirus: antigen detection in parvovirus, coronavirus and rotavirus infections in dogs in Minnesota. J Am Vet Med Ass 184: 573–575.
10. Fiscus SA, Teramoto YA (1987) Antigenic comparison of feline coronavirus and canine coronavirus. J Virol 61: 2139–2151.
11. Hohdatsu T, Sasamoto T, Okada S, Koyama H (1991) Antigenic analysis of feline infectious peritonitis virus with a large deletion in the 5' terminal region of genome. Vet Microbiol 28: 13–24.
12. Stiba N, Maeda K, Kato H, Mochizuki M, Iwata H (2007) Differentiation of feline coronavirus type I and II infections by virus neutralization test. Vet Microbiol 124: 348–352.
13. Hohdatsu T, luniyama Y, Yokoyama Y, Kida K, Koyama H (1998) Differences in virus receptor for type I and type II feline infectious peritonitis virus. Arch Virol 143: 839–850.
14. Tekey G, Hofmann-Lehmann R, Bank-Wolf B, Maier R, Thiel HJ, et al. (2010) Chimeric feline coronaviruses that encode type II spike protein on type I genetic background display accelerated viral growth and altered receptor usage. J Virol 84: 3326–3333.
15. Binn LN, Lazar EC, Keenan KP, Huxsoll DL, Marchwicki RH, et al. (1974) Sequence of mouse hepatitis virus A59 mRNA 2: indications for RNA recombination between coronaviruses and influenza virus. Virology 160: 415–422.
16. Carmichael LE (1978) Infectious canine enteritis caused by corona-like virus: virology and pathology. J Virol 23: 449–456.
17. Pasternak AO, Spaan WJ, Suijder EJ (2006) Nidovirus transcription: how to make sense... J Gen Virol 87: 1403–1421.
18. Bolles M, Donaldson E, Baric R (2011) SARS-CoV and Emergent Coronavirus: Viral Determinants of Interspecies Transmission. Curr Opin Virol 1: 624–634.
19. Bandia C, Ishiguro S, Massuy N, Hohdatsu T, Mochizuki M (1999) Canine coronavirus infections in Japan: virological and epidemiological aspects. J Vet Med Sci 61: 731–736.
20. Naylor MJ, Monckton RP, Lehrbach PR, Deane EM (2001) Canine coronavirus in Australian dogs. Aust Vet J 79: 116–119.
21. Yeşil automátik Yılmaz Z, Torun S, Pratefi A (2004) Canine coronavirus infection in turkey dog population. J Vet Med B Infect Dis Vet Public Health 51: 353–355.
22. Schulz BS, Strauch G, Mueller RS, Eichhorn W, Hartmann K (2008) Comparison of the prevalence of enteric viruses in healthy dogs and those with acute haemorrhagic diarrhoea by electron microscopy. J Small Anim Pract 49: 84–90.
23. Pratefi A, Martella V, Decaro N, Tinelli A, Camero M, et al. (2003) Genetic diversity of a canine coronavirus detected in pups with diarrhoea in Italy. J Virol Methods 110: 9–17.
24. Lorusso A, Decaro N, Schellen P, Rottier PJ, Buonavoglia C, et al. (2000) Gain, preservation, and loss of a group I coronavirus accessory glycoprotein. J Virol 82: 10312–10317.
25. Pratefi A, Decaro N, Tinelli A, Martella V, Elia G, et al. (2004) Two genotypes of canine coronavirus simultaneously detected in the fecal samples of dogs with diarrhoea. J Clin Microbiol. 42: 1797–1799.
26. Decaro N, Mari V, Elia G, Addie DD, Camero M, et al. (2010) Recombinant canine coronaviruses in dogs, Europe. Emerg Infect Dis 16: 41–47.
27. Soma T, Ohbata T, Ichii H, Takahashi T, Talaraquah S, et al. (2011) Detection and genotyping of canine coronavirus RNA in diarrheic dogs in Japan. Res Vet Sci 90: 203–207.
28. Decaro N, Mari V, Campolo M, Lorusso A, Camero M, et al. (2009) Recombinant canine coronaviruses related to transmissible gastroenteritis virus of Swine are circulating in dogs. J Gen Virol 83: 1527–1538.
29. Buonavoglia C, Decaro N, Martella V, Elia G, Campolo M, et al. (2006) Canine coronavirus highly pathogenic for dogs. Emerg Infect Dis 12: 492–494.
30. Decaro N, Buonavoglia C (2008) An update on canine coronaviruses: viral evolution and pathobiology. Vet Microbiol 132: 221–234.
31. Decaro N, Elia G, Martella V, Campolo M, Mari V, et al. (2010) Immunity after natural exposure to enteric canine coronavirus does not provide complete protection against infection with the new pantropic CB/05 strain. Vaccine 28: 724–729.
32. Jacobs-Gee FS, Horzinek MC (1983) Expression of feline infectious peritonitis coronavirus antigens on the surface of feline macrophage-like cells. J Gen Virol 64: 1859–1866.
33. Hohdatsu T, Okada S, Koyama H (1991) Characterization of monoclonal antibodies against feline infectious peritonitis virus type II and antigenic relationship between feline, porcine, and canine coronaviruses. Arch Virol 117: 85–95.
34. Hayashi T, Yanei T, Touroumd M, Nakayama H, Watake Y, et al. (1981) Serodiagnosis for feline infectious peritonitis by immunofluorescence using infected suckling mouse brain sections. J Vet Med Sci 43: 669–676.
35. Mochizuki M, Mitsutake Y, Miyahora Y, Higashihara T, Shimizu T, et al. (1997) Antigenic and plaque variations of serotype II feline infectious peritonitis coronavirus. J Vet Med Sci 59: 253–258.
36. Terada Y, Shiozaki Y, Shiozawa H, Mahmoud HY, Nomura K, et al. (2012) Feline infectious peritonitis virus with a large deletion in the 5’ terminal region of spike gene retains its virulence for cats. J Gen Virol 93: 1930–1934.
37. Mochizuki M, Hashimoto M, Ishida T (2001) Recent epidemiological status of canine viral enteric infections and Giardia infection in Japan. J Vet Med Sci 63: 573–575.
38. Naylor MJ, Galia CS, McOrist S, Lehrbach PR, Deane EM, et al. (2002) Molecular characterization confirms the presence of a divergent strain of canine coronavirus (UWSMN-1) in Australia. J Clin Microbiol 40: 3518–3522.
39. Wang YY, Lu CP (2009) Analysis of putative recombination hot sites in the S gene of canine coronaviruses. Acta Virol 53: 111–112.

Table S4 Amino acid sequence identities of partial RdRp among type II CCoV and types I and II FCoV.

Table S5 Amino acid sequence identities of partial S protein among type II CCoV and types I and II FCoV.

Table S6 Amino acid sequence identities of N protein among type II CC0V and types I and II FC0V.

Author Contributions

Conceived and designed the experiments: KM. Performed the experiments: YT NM KN RK HS TM MM KM. Analyzed the data: YT KM. Contributed reagents/materials/analysis tools: MM TS KM. Contributed to the writing of the manuscript: YT NM KM.
40. Terada Y, Minami S, Noguchi K, Mahmoud HY, Shimoda H, et al. (2014) Genetic characterization of coronaviruses from domestic ferrets, Japan. Emerg Infect Dis 20: 284–287.
41. Saitou N, Nei M (1987) The neighbor-joining method: a new method for reconstructing phylogenetic trees. Mol Biol Evol 4: 406–425.
42. Tamura K, Peterson D, Peterson N, Stecher G, Nei M, et al. (2011) MEGA5: Molecular Evolutionary Genetics Analysis using Maximum Likelihood, Evolutionary Distance, and Maximum Parsimony Methods. Mol Biol Evol 28: 2731–2739.
43. Lole KS, Bollinger RC, Paranjape RS, Gadkari D, Kulkarni SS, et al. (1999) Full-length human immunodeficiency virus type 1 genomes from subtype C-infected seroconverters in India, with evidence of intersubtype recombination. J Virol 73: 152–160.
44. Motokawa K, Hohdatsu T, Hashimoto H, Koyama H, et al. (1996) Comparison of the amino acid sequence and phylogenetic analysis of the peplomer, integral membrane and nucleocapsid proteins of feline, canine and porcine coronaviruses. Microbiol Immunol 40: 425–433.
45. Herrewegh AA, Smeenk I, Horznicek MC, Rottier PJ, de Groot RJ (1998) Feline coronavirus type II strains 79–1683 and 79–1146 originate from a double recombination between feline coronavirus type I and canine coronavirus. J Virol 72: 4508–514.
46. Lin CN, Chang KY, Su BL, Chueh LL (2013) Full genome analysis of a novel type II feline coronavirus NTU156. Virus Genes 46: 316–322.
47. Vennema H (1999) Genetic drift and genetic shift during feline coronavirus evolution. Vet Microbiol 69: 139–141.
48. Addie DD, Schaap IA, Nicolson L, Jarrett O (2003) Persistence and transmission of natural type I feline coronavirus infection in cats; a possible role in feline infectious peritonitis. J Gen Virol 84: 2735–2744.
49. Benetka V, Kühler-Heus A, Kolodziejek J, Nowotny N, Hofmann-Parisot M, et al. (2004) Prevalence of feline coronavirus types I and II in cats with histopathologically verified feline infectious peritonitis. Vet Microbiol 99: 31–42.
50. Kumiron M, Meli ML, Harsig M, Gröenzi E, Polan A, et al. (2005) Feline coronavirus serotypes 1 and 2: seroprevalence and association with disease in Switzerland. Clin Diag Lab Immunol 12: 1209–1215.
51. Hohdatsu T, Okada S, Ishizuka Y, Yamada H, Koyama H (1992) The prevalence of types I and II feline coronavirus infections in cats. J Vet Med Sci 54: 557–562.
52. Masters PS (2006) The molecular biology of coronaviruses. Adv Virus Res 66: 193–292.
53. Tresnan DB, Levis R, Holmes KV (1996) Feline aminopeptidase N serves as a receptor for feline, canine, porcine, and human coronaviruses in serogroup I. J Virol 70: 8669–8674.
54. McArdle F, Bennett M, Gaskell RM, Tennant B, Kelly DF, et al. (1990) Canine coronavirus infection in cats; a possible role in feline infectious peritonitis. Adv Exp Med Biol 276: 475–479.
55. Pedersen NC (2009) A review of feline infectious peritonitis virus infection: 1963–2008. J Feline Med Surg 11: 225–250.
56. Pedersen NC, Liu H, Scarlett J, Leutenegger GM, Golevko L, et al. (2012) Feline infectious peritonitis: role of the feline coronavirus 3c gene in intestinal tropism and pathogenesis based upon isolates from resident and adopted shelter cats. Virus Res 165: 17–20.
57. Chang HW, de Groot RJ, Egberink HF, Rottier PJ (2010) Feline infectious peritonitis: insights into feline coronavirus pathobiogenesis and epidemiology based on genetic analysis of the viral 3c gene. J Gen Virol 91: 415–420.
58. Vennema H, Poland A, Foley J, Pedersen NC (1998) Feline infectious peritonitis viruses arise by mutation from endemic feline enteric coronaviruses. Virology 243: 150–157.
59. Pedersen NC, Liu H, Dodd KA, Pesavento PA (2009) Significance of coronavirus mutants in feces and diseased tissues of cats suffering from feline infectious peritonitis. Viruses 1: 166–184.
60. Wang YT, Su BL, Hsieh LE, Chueh LL (2013) An outbreak of feline infectious peritonitis in a Taiwanese shelter: epidemiologic and molecular evidence for horizontal transmission of a novel type II feline coronavirus. Vet Res. 44: 57.
61. Graham RL, Baric RS (2010) Recombination, reservoirs, and the modular spike: mechanisms of coronavirus cross-species transmission. J Virol 84: 3144–3146.