

## Coincidental detection of genomes of porcine parvoviruses and porcine circovirus type 2 infecting pigs in Japan

Prayuth SAEKHOW<sup>1,2</sup>), Shingo KISHIZUKA<sup>1</sup>), Natsuha SANO<sup>1</sup>), Hiroko MITSUI<sup>1</sup>), Hajime AKASAKI<sup>1</sup>), Takahiro MAWATARI<sup>3</sup>) and Hidetoshi IKEDA<sup>1,4</sup>\*)

<sup>1</sup>)Laboratory of Veterinary Hygiene, Graduate School of Veterinary Medicine and Life Science, Nippon Veterinary and Life Science University, Musashino, Tokyo 180–8602, Japan

<sup>2</sup>)Department of Veterinary Biosciences and Public Health, Faculty of Veterinary Medicine, Chiang Mai University, Chiang Mai 50100, Thailand

<sup>3</sup>)Yamagata Prefectural Central Livestock Health and Sanitation Office, Yamagata, Japan

<sup>4</sup>)National Institute of Animal Health, Tsukuba, Ibaraki 305–0856, Japan

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**ABSTRACT.** The infection status of 15 viruses in 120 pigs aged about 6 months was investigated based on tonsil specimens collected from a slaughterhouse. Only 5 species of porcine parvoviruses and porcine circovirus type 2 (PCV2) were detected at high frequencies; 67% for porcine parvovirus (PPV) (PPV-Kr or -NADL2 as the new abbreviation), 58% for PPV2 (CnP-PARV4), 39% for PPV3 (P-PARV4), 33% for PPV4 (PPV4), 55% for PBo-likeV (PBoV7) and 80% for PCV2. A phylogenetic analysis of PPV3 suggested that Japanese PPV3s showed a slight variation, and possibly, there were farms harboring homogeneous or heterogeneous PPV3s. Statistical analyses indicated that the detection of PCV2 was significantly coincidental with each detection of PPV, PPV2 and PPV3, and PPV and PPV4 were also coincidentally detected. The concurrent infection with PCV2 and porcine parvoviruses in the subclinically infected pigs may resemble the infection status of pigs with the clinical manifestations of porcine circovirus associated disease which occurs in 3–5 months old pigs and is thought to be primarily caused by the PCV2 infection.

**KEY WORDS:** coinfection, porcine circovirus associated disease, porcine circovirus type 2, porcine parvoviruses, prevalence

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A number of new parvoviruses have been identified during the past 15 years and given various names, and thereby, the updated taxonomy of the family *Parvoviridae* was proposed in 2014 [6]. The classical porcine parvovirus (PPV), which was first identified in the 1960s [4] and now present worldwide, causes embryonic death, stillbirths and mummification when embryos or fetuses in seronegative dams are infected. The newly identified porcine parvoviruses have been detected in various areas of the world, but its relationship with any diseases remains unclear. PPV is thought to be one of the cofactors for porcine circovirus associated disease (PCVAD) whose main etiologic agent is porcine circovirus type 2 (PCV2) [1, 10, 13]. The PCV2 infection alone does not cause a clinical disease, but concurrent viral or bacterial infections may augment the severity of PCVAD possibly through stimulating the PCV2 replication or suppressing the PCV2 clearance by altered cytokine regulation [8, 9, 18].

During our screening for known viral genomes and newly identified porcine parvovirus genomes in specimens of apparently healthy pigs, we found and now report that

the genomes of PCV2 and the classical and new porcine parvoviruses were coincidentally detected. The 5 porcine parvoviruses we studied include PPV [4], PPV2 [11], PPV3 [14], PPV4 [5] and porcine bocavirus-like virus (PBo-likeV) [2]. According to the proposed taxonomy of the family *Parvoviridae* [6], most of the virus names have been changed as indicated in Table 1. However, we use the previous abbreviations in this paper to avoid confusion.

### MATERIALS AND METHODS

**Sample collection and viral nucleic acid purification:** Tonsil specimens from 120 pigs were collected from a slaughterhouse in 2010 when most of the pigs were probably not injected with the inactivated PCV2 vaccine in Japan. The pigs were about 6 months old and obtained from 22 farms with 1–10 samples per farm.

The procedures for the viral DNA and RNA isolation were previously described [23]. Briefly, the tonsil homogenates were prepared using a Micro Smash machine (Tomy Seiko, Tokyo, Japan), and after centrifugation at 15,000 g for 15 min, aliquots of the supernatant were stored at –80°C. The viral DNA and RNA were isolated by a DNA/RNA purification machine, Magstration System 6GC (Precision System Science, Chiba, Japan) and a solution kit, GC series Magstration-MagaZorb RNA Common Kit (Precision System Science). The isolated nucleic acids were reverse-transcribed by Superscript II reverse transcriptase and primers of random hexamers according to the manufacturer's instructions (In-

\*CORRESPONDENCE TO: IKEDA, H., Laboratory of Veterinary Hygiene, Graduate School of Veterinary Medicine and Life Science, Nippon Veterinary and Life Science University, 1–7–1 Kyonan-cho, Musashino, Tokyo 180–8602, Japan. e-mail: hikeda@nvlu.ac.jp

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Table 1. Prevalence of 15 virus genomes in 120 pigs

Virus	Abbreviation	Prevalence (%; n=120)
porcine parvovirus (porcine parvovirus) <sup>a)</sup>	PPV	67
porcine parvovirus 2 (porcine Cn virus) <sup>a)</sup>	PPV2 (CnP-PARV4) <sup>a)</sup>	58
porcine parvovirus 3 (porcine hokovirus) <sup>a)</sup>	PPV3 (P-PARV4) <sup>a)</sup>	39
porcine parvovirus 4 (porcine parvovirus 4) <sup>a)</sup>	PPV4 (PPV4) <sup>a)</sup>	33
porcine boca-like virus (porcine bocavirus 7) <sup>a)</sup>	PBo-likeV (PBoV7) <sup>a)</sup>	55
porcine circovirus 2	PCV2	80
suid herpesvirus 1	SuHV1	0
hepatitis E virus	HEV	0
swine influenza virus	SIV	0
porcine reproductive and respiratory syndrome virus	PRRSV	0
Japanese encephalitis virus	JEV	0
porcine epidemic diarrhea virus	PEDV	0
porcine rotavirus A	PoRV-A	0
transmissible gastroenteritis virus	TGEV	0
Getah virus	GETV	0

a) New names recently proposed for the family Parvoviridae [6]. The prevalence of PPV2 was previously described [23], but for convenience, the data were included in this table.

vitrogen, Carlsbad, CA, U.S.A.) and used as templates for the various PCRs detecting the genomes of both the DNA and RNA viruses. For the validation of reverse transcription-PCR assay, control RNA and PCR primers of the kit were used in each experiment.

**Detection of viral genome by PCR:** Two multiplex PCRs for 3 DNA viruses (porcine circovirus type 2 (PCV2), suid herpesvirus 1 and porcine parvovirus (PPV)) and 6 RNA viruses (porcine reproductive and respiratory syndrome virus (PRRSV), Japanese encephalitis virus, porcine rotavirus A (PoRV-A), porcine epidemic diarrhea virus (PEDV), transmissible gastroenteritis virus (TGEV) and Getah virus) were performed in separate tubes according to the published method [17]. Other PCR primer pairs included; NP1200 and NP1529 for swine influenza virus [15], HE5-1 and HE5-4m for hepatitis E virus [26], Q1 F and Q2 R for porcine parvovirus 2 [11], PPV3 F and PPV3 R for porcine parvovirus 3 [25], PPV4 F and PPV4 R for porcine parvovirus 4 [25], and SbocaF and SbocaR for PBo-likeV [32].

Viral genomes were amplified by PCR using Quick Taq HS DyeMix (Toyobo, Osaka, Japan) including Taq polymerase. The PCR consisted of an initial enzyme activation step at 94°C for 5 min, followed by 35 cycles of denaturation at 94°C for 30 sec, annealing at 55°C for 30 sec, extension at 72°C for 30 sec and a final extension at 72°C for 7 min.

**Phylogenetic analysis:** For the phylogenetic analysis of PPV3, the 622 bp of the VP gene were examined after amplifying the 713 bp fragment of the VP region (nucleotide positions from 3,359 to 4,072 of the strain AB916464) with the PCR primers PPV3 P7F (3'-GGGGCACTCATTTCTCTGAT-5') and PPV3 P7R (3'-CTGGCCTTTTCCACTTAGGA-5') [25] and sequencing with both primers and the two internal primers PPV3 7F2 (3'-GGAGAATAATGTTCTTCCTC-5') and PPV3 7R2 (3'-TCGTACTCATCAAGCAGCTG-5').

The sequence data and phylogenetic tree were compiled and analyzed using MEGA 5.1 [27] and Genetyx (Genetyx Co., Tokyo, Japan). The phylogenetic trees were generated

by the maximum likelihood method.

The partial sequences of the PCR products of PPV3 have been deposited in DDBJ under accession numbers from LC011459 to LC011478.

**Statistical analysis:** Chi-square tests were used to evaluate the statistical significance of co-isolation of two viral genomes among PPV, PPV2, PPV3, PPV4, PBo-likeV and PCV2 by dividing two categories, PCR-positive and PCR-negative individuals for each virus. *P* values of <0.05 were considered statistically significant.

## RESULTS

**Prevalence of porcine parvoviruses and porcine circovirus 2 in 120 Japanese pigs:** We previously analyzed the prevalence of the PPV2 genomes in the tonsil specimens from 120 pigs [23]. With the same specimens, we extended such a screening for 14 other viral genomes as listed in Table 1. Five of the 14 viral genomes were detected; four were members of the family *Parvoviridae* and another one was PCV2. The prevalences were 67% for PPV, 39% for PPV3, 33% for PPV4, 55% for PBo-likeV and 80% for PCV2 (Table 1), in addition to 58% for PPV2 [23]. Multiple viral genomes were detected from the individual pigs, and thereby, as for the 5 examined porcine parvoviruses, 3%, 23%, 53%, 78% and 93% of the pigs were positive for more than 5, 4, 3, 2 and 1 virus (es), respectively. Only 9 pigs of various farms were negative for the 5 parvovirus DNAs, and 4 of the 9 pigs were negative for PCV2 DNA. Among the 8 farms with larger sample numbers (8 to 10 samples per farm), 7 farms were positive for all 5 parvoviruses, and one farm was negative (0/10) for only one (PPV4) of the 5 parvoviruses. The results suggested that a high proportion of the pigs in most farms were co-infected with the five parvoviruses and PCV2.

We tested the possibility that the detections of these highly prevalent viruses were random or coincidental. The chi square analyses indicated that PCV2 was coincidentally de-

Table 2. Chi square analysis for coincidental detection among genomes of 4 parvoviruses and PCV2

Relationship between two viruses		Number of pigs				$\chi^2$ value	P value	Significance
		+/+	-/+	+/-	-/-			
PPV /	PPV2	47	22	33	18	0.153	0.695	* <sup>a</sup>
	PPV3	33	14	47	26	0.437	0.508	
	PPV4	32	7	48	33	6.154	0.013	
	PBo-likeV	46	20	34	20	0.606	0.436	
	PCV2	69	27	11	13	5.859	0.015	
PPV2 /	PPV3	28	19	41	32	0.136	0.712	*
	PPV4	25	14	44	37	1.031	0.310	
	PBo-likeV	39	27	30	24	0.152	0.697	
	PCV2	60	36	9	15	4.910	0.027	
PPV3 /	PPV4	15	24	32	49	0.012	0.913	*
	PBo-likeV	28	38	19	35	0.653	0.419	
	PCV2	42	54	5	19	4.232	0.040	
PPV4 /	PBo-likeV	26	40	13	41	3.177	0.075	*
	PCV2	32	64	7	17	0.152	0.697	
PBo-likeV /	PCV2	53	43	13	11	0.008	0.927	

a, \*: significant ( $0.01 < P < 0.05$ ). Others without asterisk mean not significant ( $P > 0.05$ ).

tected with PPV ( $\chi^2=5.86$ ,  $P<0.02$ ), PPV2 ( $\chi^2=4.91$ ,  $P<0.03$ ) or PPV3 ( $\chi^2=4.23$ ,  $P<0.04$ ) and that PPV and PPV4 were also coincidentally detected ( $\chi^2=6.15$ ,  $P<0.02$ ) (Table 2).

**Nucleotide sequence diversity of PPV3:** In order to know the genetic diversity of the Japanese PPV3s, a phylogenetic analysis based on the 622 bases of the VP gene was performed using 20 Japanese samples, 5 samples each from 4 farms and 87 reference sequences from around the world. The Japanese PPV3s were slightly diverged in the phylogenetic tree with 1.6% (10/622 bases) of the maximum nucleotide difference and closely related to the other PPV3s detected in Europe, North America, South America and Hong Kong (Fig. 1).

To characterize the variation in the nucleotide sequence among the farms or within a farm, the 20 Japanese PPV3 sequences were tentatively separated into 6 sequence groups based on their phylogenetic branch and % nucleotide difference (Figs. 1, 2 and Table 3). The sequence group 1 was a major one to which 10 of the 20 sequences belonged. Farms C and D appeared homogeneous, having 4 and 5 sequences of the sequence group 1, respectively, while farms A and B had relatively heterogeneous PPV3 sequences. Farm A had the sequence groups 2 and 6 with a 1.3% nucleotide difference (8/622 bases), and farm B had the sequence groups 1, 3 and 4 with 0.6–1.4% nucleotide differences (4–9/622 bases) (Table 3).

## DISCUSSION

The present study, together with our previous study [23], examined the tonsil specimens of 120 apparently healthy pigs for the screening of 15 viruses which can infect pigs. Only the five porcine parvoviruses, i.e., PPV, PPV2, PPV3, PPV4 and PBo-likeV, and PCV2 were detected, and their prevalences were quite high, ranging from 33% to 80% (Table 1) [23]. The high prevalences of the classical PPV and PCV2 at the age of about 6 months are common in most

pig-producing countries, whereas the prevalences of PPV3, PPV4 and PBo-likeV are the first observations in Japanese pigs.

The PPV3 DNA was detected in 39% of the 120 Japanese pigs (Table 1). Since the first identification of PPV3 [14], the prevalence has been reported in several countries and appears to widely vary from lower frequencies (6–20%) in Hungary [7], China [21], the U.S.A. [29] and Germany [25] to higher frequencies (44–73%) in Hong Kong [14], China [16] and Thailand [22].

The prevalence of the PPV4 genome was 33% in this study which is comparable to the prevalence (44%) in Thailand [22], but higher than those of several other countries, that is, 1% in China [12], 6% in Hungary [7], 7% in Germany [25] and 4% in the U.S.A. [30].

PBo-likeV [2], which was also called PBoV (PBoV-SX) [31] or PBoV1 [24, 33], is one of several porcine bocaviruses which have recently been discovered [28]. The PBo-likeV infection was initially supposed to be associated with respiratory tract diseases in pigs due to the remarkable difference in the prevalences between sick (39% (74/191)) and healthy (7% (3/41)) pigs [32]. The prevalence of PBo-likeV was 55% in our study (Table 1), in contrast to 18% in Thailand [22], 7% in China [32], 63% in different areas of China [24], 2% in Romania [7] and 13% in the wild boars of Romania [3].

Although the prevalences of PPV3, PPV4 and PBo-likeV show some variation among countries, the available data suggest that these newly identified parvoviruses have already spread worldwide.

The phylogenetic analysis of PPV3 suggested that, compared to the variation of 87 sequences deposited from around the world, the 20 Japanese sequences were less variable and belonged to limited branches (Fig. 1). In the 4 farms we analyzed, 2 farms appeared to have heterogeneous PPV3s (Table 3 and Fig. 2). Although the variations within the farms were

Fig. 1.

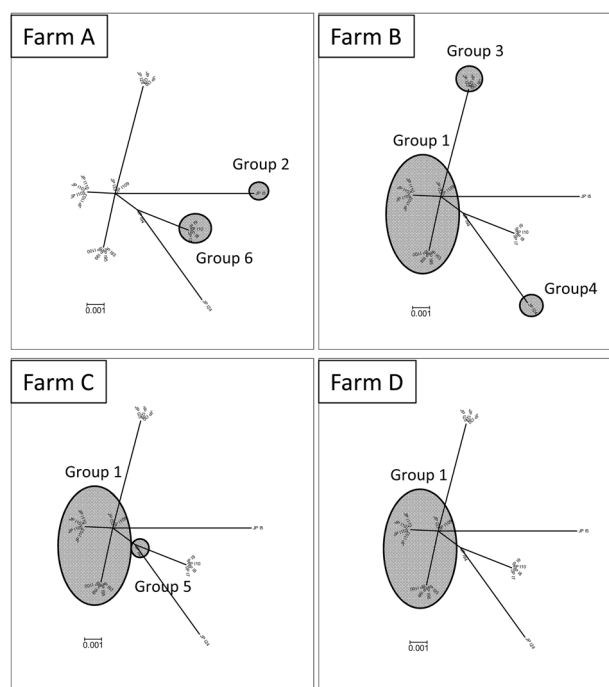
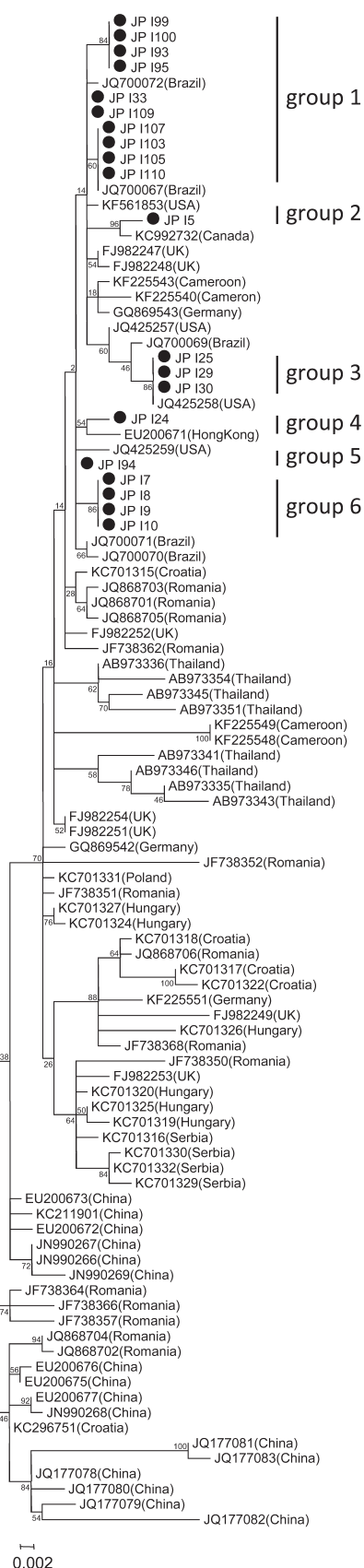


Fig. 2. The phylogenetic tree was constructed with the 20 Japanese PPV3s detected from the 4 farms based on the 622 bases of the PPV3 VP gene. The 6 sequence groups were tentatively defined by the phylogenetic branch and % nucleotide difference, i.e., <0.5% (3/622) within each sequence group. The 6 sequence groups detected from the 4 pig farms are indicated. The relationship among the sequence data, the sequence group and the farm is indicated in Table 3.

not high, this raises the possibility that the observed variation within a farm resulted from multiple invasions of different strains rather than natural mutations within a farm after the invasion of one strain. The coexistence of different strains in a farm and coinfection of a pig with different strains must be risk factors for vaccine strategies and generation of a new recombinant virus strain.

The PCV2 genome was detected at a high frequency (80%) (Table 1) which is common worldwide. Interestingly, PCV2 was coincidentally detected along with PPV, PPV2 or PPV3, and PPV and PPV4 were also coincidentally detected (Table 2). These associations were weak, but statistically significant ( $0.01 < P < 0.05$ ). Recently, similar associations were observed in pigs with PCVAD; the prevalences of the PPV and PPV2 DNAs were significantly higher in the

Fig. 1. The phylogenetic tree was constructed, based on the 622 bases of the PPV3 VP gene, with the 20 Japanese PPV3s and 87 PPV3s currently deposited in the data bank. For the Japanese sequences, the 6 tentative sequence groups (Sequence groups 1–6) were defined by phylogenetic branch and % nucleotide difference, i.e., <0.3% (2/622) within each sequence group. The relationship between the farm and the sequence group of the detected PPV3 sequences is indicated in Table 3.

Table 3. Nucleotide sequence diversity of PPV3 within 4 farms

Farm	"sequence group"						Nucleotide difference
	1 <sup>a)</sup>	2	3	4	5	6	
A		JP 15				JP 17 JP 18 JP 19 JP 110	Sequence groups 2 vs 6: 1.3% (8/622)
B	JP 133		JP 125 JP 129 JP 130	JP 124			Sequence groups 1 vs 3: 0.8% (5/622) Sequence groups 1 vs 4: 0.6% (4/622) Sequence groups 3 vs 4: 1.4% (9/622)
C	JP 193 JP 195 JP 199 JP 1100				JP 194		Sequence groups 1 vs 5: 0.5% (3/622)
D	JP 1103 JP 1105 JP 1107 JP 1110 (JP 1109) <sup>b)</sup>						

PCR products were directly sequenced, and the sequence data of the 622 bases of the PPV3 VP gene were subjected to a phylogenetic analysis. The "sequence group" in this analysis was defined by the phylogenetic branch (Fig. 1) and % nucleotide difference. In this table, nucleotide sequences in a box were identical, except for the JP 1109 sequence of the sequence group 1 in farm D. a), In the "sequence group 1", the JP 133 sequence of farm B was identical to the JP 1109 sequence of farm D, which are located at the center of the phylogenetic tree of Fig. 2. The 4 sequences of farm C (JP 193, JP 195, JP 199 and JP 1100) were identical, and the 4 sequences of farm D (JP 1103, JP 1105, JP 1107 and JP 1110) were also identical. The sequence of JP 133 and JP 1109 differed by 2 bases from the 4 identical sequences of farm C and by 1 base from the 4 identical sequences of farm D. The 4 sequences of farm C and the 4 sequences of farm D differed by 3 bases. b), the JP 1109 sequence was slightly different (0.2% (1/622)) from the other 4 sequences at the same farm.

PCVAD cases containing high amounts of PCV2 DNA than in the non-PCVAD cases, while, in contrast to our data, PPV3, PPV4 and PPV5 were not correlated with the amount of PCV2 [20]. The major difference between the two studies is that they analyzed the lungs of pigs with PCVAD probably aged 3–5 months while we used the tonsils of subclinical pigs aged about 6 months.

PCV2 is recognized as a causative agent of PCVAD. The clinical features of PCVAD or formerly called postweaning multisystemic wasting syndrome (PMWS) caused by PCV2 are systemic including enlargement of the lymph nodes, progressive loss of body weight or wasting combined with difficulty in breathing, diarrhea, pale skin and jaundice [9, 19]. The histopathologic changes in the affected lymphoid tissues are a severe lymphoid depletion, a diffuse infiltration of histiocytic cells and various inflammatory lesions. The pathogenesis of PCVAD or PCV2-induced diseases is complex, probably involving PCV2 infection and cofactors, such as other infections and altered cytokine or immune responses [8]. Particularly, the concurrent infection of PCV2-infected pigs by viruses (PPV, PRRSV, etc.), bacteria (*Mycoplasma hyopneumoniae*) or parasites may not be only a secondary infection after PCV2-induced depletion of lymphocytes, but could be important for the disease manifestation [18]. The experimental inoculation with PCV2 and PPV, but not PCV2 alone, could reproduce lesions similar to those of the field cases of PMWS [1, 10, 13]. The mechanism for the synergistic effect of coinfection was proposed that coinfection may promote the PCV2 infection by stimulating immune

cells and providing target cells for the PCV2 replication or suppressing the PCV2 clearance by alteration of the cytokine production and profiles [1, 18].

The coincidental detections of PCV2 and PPVs in various combinations have been observed in both pigs with PCVAD at 3–5 months old and healthy pigs at about 6 months old. Therefore, the two stages may share a common mechanism for the proliferation of these viruses regardless of the presence or absence of PCVAD. Since circovirus and parvovirus are both DNA viruses, which require actively proliferating cells for efficient viral replication, lymphoproliferation or immunosuppression induced by infection with a virus could support the growth of other viruses.

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