Role of the Alpha/Beta Interferon Response in the Acquisition of Susceptibility to Poliovirus by Kidney Cells in Culture

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Replication of poliovirus (PV) is restricted to a few sites, including the brain and spinal cord. However, this neurotropism is not conserved in cultured cells. Monkey kidney cells become susceptible to PV infection after cultivation in vitro, and cell lines of monolayer cultures from almost any tissue of primates are susceptible to PV infection. These observations suggest that cellular changes during cultivation are required for acquisition of susceptibility. The molecular basis for the cellular changes during this process is not known. We investigated the relationship between PV susceptibility and interferon (IFN) response in primary cultured kidney and liver cells derived from transgenic mice expressing human PV receptor and in several primate cell lines. Both kidneys and liver in vivo showed rapid IFN response within 6 h postinfection. However, monkey and mouse kidney cells in culture and primate cell lines, which were susceptible to PV, did not show such rapid response or showed no response at all. On the other hand, primary cultured liver cells, which were partially resistant to infection, showed rapid IFN induction. The loss of IFN inducibility in kidney cells was associated with a decrease in expression of IFN-stimulated genes involved in IFN response. Mouse kidney cells pretreated with a small dose of IFN, in turn, restored IFN inducibility and resistance to PV. These results strongly suggest that the cells in culture acquire PV susceptibility during the process of cultivation by losing rapid IFN response that has been normally maintained in extraneural tissues in vivo.

Poliovirus (PV), belonging to the Picornaviridae, is the causative agent of poliomyelitis (44). The replication of PV is limited to a few sites, including the brain and spinal cord. Severe pathological lesions are not observed in most extraneural tissues, despite the presence of virus in many tissues during the viremic phase of infection (2, 52). PV was initially isolated by Landsteiner and Popper (32) in 1908 as a transmissible pathogen. In those days, PV could be transferred by inoculating a suspension of the spinal cord of a paralyzed monkey into another, because of the neurotropic nature of PV. In 1949, Enders et al. (8) for the first time succeeded in propagating PV in primary cultured cells from various human embryonic tissues. Later, Dulbecco and Vogt (7) found that monkey kidney developed permissivity to infection after cultivation in vitro and that PV titers could be quantified by plaque assay. PV, therefore, is able to replicate in monolayer cells in primary culture and in cell lines derived from almost any tissue of primates, although PV cannot replicate well in the extraneural tissues in vivo. This new technology for propagating PV in cell cultures in vitro had a great impact on virology and allowed revolutionary progress in PV studies. Attenuated Sabin strains were developed by a number of passages of virulent PV strains in cultured cells (53). Large-scale production of PV vaccines has been done using monkey kidney cells. The molecular mechanisms of PV replication have been studied using cultured cells with strict control of the experimental conditions (49). Although we have benefited greatly from this, the molecular basis for this paradoxical change in susceptibility is still unknown.

The factors that control PV susceptibility have been studied. The research focused on identifying the determinants of tissue and cell tropism. Holland and colleagues thought that susceptibility was determined at the level of virus entry into the cell. They proposed that the PV receptor (PVR) is a major determinant of tissue and cell tropism based on observations that a single round of replication occurred in nonpermissive mouse cells after transfection of PV genomic RNA (20) and that PV was adsorbed by a homogenate of neural tissues (19). Molecular cloning of the human PVR gene revealed that PVR is a membrane protein that belongs to the immunoglobulin superfamily (29, 40). Transgenic (tg) mice that carry the human PVR gene were produced. PV selectively replicated in the central nervous system (CNS), and the mice showed paralytic disease that resembled human poliomyelitis upon PV infection (31, 51). If Holland’s hypothesis were correct, one could expect that PV would not be expressed in the extraneural tissues at high levels in vivo but would be expressed in the cells after cultivation in vitro. However, PVR mRNA was detected in various tissues of human and PVR-tg mice (43). Furthermore, when Ren and Racaniello (50) investigated the distribution of PVR...
transcripts in the kidneys of tg mice by in situ hybridization, they showed that epithelial cells in the Bowman’s capsule, podocytes in the glomerulus, and some of the tubular epithelial cells in the medulla expressed human PVR mRNA at high levels, although these cells were not susceptible to PV in vivo. They also observed that the freshly dispersed kidney cells expressed PVR at the cell surface, as judged by a PV binding assay, but susceptibility developed after cultivation in vitro for 24 h. They concluded that expression of PVR is not sufficient for PV susceptibility and predicted that other factors that might change during the cultivation process are also needed.

Another possibility was that susceptibility was determined at the level of translation initiation of the viral protein. Translation initiation of picornaviruses is controlled by an internal ribosome entry site (IRES) located in the 5’ non-coding region of the genome. The PV IRES interacts with canonical translation initiation factors and non-canonical translation initiation factors, IRES trans-activating factors (ITAFs), to achieve efficient translation initiation (17). To date, polyuridine tract binding protein (PTB) (18, 46), neural cell-specific PTB (27, 35, 37, 48), La autoantigen (38, 39), poly(rC) binding protein 2 (PCBP-2) (1), and upstream of N-ras (Unr) (3) have been identified as ITAFs for PV. There is some evidence for tissue-specific or cell-type-specific translation initiation mediated by the IRES in picornaviruses. Chimeric PVs containing replacements of IRES sequences with corresponding sequences from human rhinovirus type 2 or hepatitis C virus did not propagate in the CNS of PVR-tg mice (14, 63). The foot-and-mouth disease virus IRES was not active in neurons because it required ITAFs, which was expressed only in proliferating cells (47). The PV Sabin 3 strain did not grow efficiently in the CNS, in which PTB was not expressed at high levels, and a translation deficit of the Sabin 3 IRES was rescued by increased expression of PTB in the CNS (15). These results may be good examples demonstrating that a lack or shortage of ITAFs resulted in tissue- or cell-specific failure of replication of viruses. If the expression of ITAFs determines susceptibility, we would expect that IRES activity would not be observed in the kidneys in vivo. Kauder and Racaniello (26) reported contradictory results. They constructed a recombinant adenovirus that expressed bicsrticrin reporter luciferase genes under the control of PV IRESs. They infected mice intravenously with the recombinant adenovirus vector and determined reporter gene expression by luciferase assay. They showed that the PV IRES was active in neural, as well as in extraneural, tissues, including the kidneys. This implies that kidney cells in vivo express sufficient levels of all ITAFs for PV. Furthermore, they did not observe the neural-tissue-specific reduction of translational activity of the attenuated Sabin IRES. Sabin 3 virus and hepatitis C virus/PV recombinant virus propagated efficiently in the CNS of neonatal PVR-tg mice and caused paralysis. They then concluded that PV tropism and attenuation are determined after internal ribosomal entry. Since direct measurement of expression profiles of ITAFs in adult and developing tissues has not been reported, the contribution of the ITAFs to tissue- or cell-specific infection of PV is still controversial (56; see reference 55 for a review).

Finally, we have recently demonstrated that PV tissue tropism is strongly influenced by antiviral activity mediated by alpha/beta interferons (IFN-α/β) (21). PVR-tg mice deficient in the alpha/beta IFN receptor 1 (Ifnar-1) gene were produced by crossing PVR-tg mice and Ifnar knockout mice (41). Although PV replication sites were restricted to the CNS in the wild-type PVR-tg mice, extensive PV replication was observed in a wider range of tissues in the PVR-tg/Ifnar knockout mice. This result suggests that extraneuronal tissues, such as the liver, spleen, and pancreas, are potentially susceptible to PV infection and that they are normally protected by the IFN response. In the wild-type PVR-tg mice, neural tissues expressed very low levels of IFN-stimulated genes (ISGs) and did not show rapid IFN response upon PV infection. However, extraneuronal tissues expressed slightly higher levels of ISGs, even in the uninfected mice. They showed sufficient IFN response and were protected from PV infection. From these results, we consider that the difference in IFN responses among the tissues influences differential PV susceptibility.

We hypothesized that the acquisition of PV susceptibility by kidney cells after cultivation may be a consequence of changes in either the IFN response, PVR expression, or ITAF expression. Using primary cultured kidney cells and liver cells from PVR-tg mice, we investigated the relationship between PV susceptibility and expression of the factors mentioned above. Here, we present evidence that the loss of rapid IFN inducibility associated with the decrease in expression of ISGs involved in IFN response during the cultivation process plays an important role in the change in susceptibility of kidney cells.
kidney and liver RNAs were purchased from BD Biosciences. Real-time PCR was performed using an ABI PRISM 7500. The quantification of mouse 18S rRNA, mIFN-α mRNA, and human IFN-α mRNA was performed by the SYBR green method using 18S rRNA-F (5'-GTA ACC CGT TGA ACC CCA TT-3'), 18S rRNA-R (5'-CCA TCC AAT CGG TAG TAG CG-3'), mIFNα-F (5'-CTG TGA ACC TCT TCA CAT CAA A-3'), mIFNα-R (5'-ACA GCC TTG CAG GTC ATT GAG-3'), hIFN-α-F (5'-GTA CTG CAG AAT CTC TCT TCT CTC-3'), and hIFN-α-R (5'-GTG TCT AGA TCT GAC AAC CTC CCA GGG CAC A-3') as primers. The mIFNα-F and mIFNα-R primers amplified all mouse IFN-α mRNA species, and hIFN-α-F and hIFN-α-R primers amplified all human IFN-α mRNA species. The quantification of mouse IFN-β, 2.5'-oligo(deoxy)nucleotyl synthetase (OAS), RIG-I, MDA-5, IRF-3, IRF-7, IRF-9, STAT-1, STAT-2, TBK-1, IKKe, IFNAR-1, La, PTB, PCBP-2, UNR, glyceraldehyde-3-phosphate dehydrogenase (GAPDH), PVR mRNAs, and 18S rRNA was performed using Assay-on-Demand PCR probes (Applied Biosystems). The amounts of mRNAs were determined by comparison with the standard templates of cloned cDNAs of known copy number. The expression levels were then normalized to the levels of 18S rRNA. The data are represented as the copy number of mRNA per 10⁶ copies of rRNA.

**Immunohistochemistry.** Detection of PV antigens in infected mice was performed as described previously (31).

**Effect of IFN treatment of kidney cells.** Mouse kidney cells were treated with recombinant mouse IFN-β (Tora) at concentrations of 0.1 to 100 IU/ml for 6 h. The cells were washed with MEM three times. The IFN-treated cells were infected with PV (at a multiplicity of infection [MOI] of 0.01) in the presence or absence of anti-mouse IFN-α (17.5 μg/ml) and anti-mouse IFN-β (1.8 μg/ml) monoclonal antibodies (Yamasa Shoyu Co. Ltd, Choshi, Japan). The antibodies at the above concentrations can block 10 and 1.000 IU of IFN-α and IFN-β, respectively. The expression levels of the IFN-α, IFN-β, and Oas1a mRNAs at the indicated times were determined by reverse transcription (RT)-PCR. The PV titer was determined by plaque assay. Mouse IFN activity was measured by the cytopathic effect (CPE) dye uptake method using L929 cells (22, 67). The NIH research reference reagent for mouse IFN-β (National Institute of Allergy and Infectious Diseases, NIH, Bethesda, Md.) was used as the standard for unit definition.

**RESULTS**

**PV susceptibility and IFN response in the kidneys and liver in vivo.** We previously reported that most of the extraneural tissues in the PVR-tg mice are potentially susceptible to PV infection but that they are normally protected by the IFN response. We investigated PV replication and IFN response in the kidneys and liver in vivo in detail. Mice were intravenously inoculated with 10⁷ PFU of PV and were sacrificed daily. Virus titers in the tissues were determined. The PV titers in the kidneys (Fig. 1A) and livers (Fig. 1B) of PVR-tg mice decreased similarly to those of non-tg mice, which suggests that PV cannot replicate well in these tissues of PVR-tg mice. However, the virus titers in the kidneys of the PVR-tg/Ifnar knockout mice increased over time (Fig. 1A). The PV titers in the livers of the PVR-tg/Ifnar knockout mice remained at approximately 10⁷ PFU/g tissue until 3 days postinfection (p.i.), with only a slight decrease observed (Fig. 1B).

We inoculated 10⁸ PFU of PV intravenously into PVR-tg and PVR-tg/Ifnar knockout mice and detected PV antigens by immunohistochemistry. PV antigens were detected in the glomeruli of the kidneys obtained from the PVR-tg/Ifnar knockout mice (Fig. 1D), but not from those of PVR-tg mice (Fig. 1C). This indicates that at least cells in the glomeruli of kidneys in vivo are also potentially susceptible to PV infection. These cells may be exposed most easily to PV circulating in the blood. Consistent with the profile of the PV titer, a few PV antigen-positive cells were detected in the livers of the infected PVR-tg mice (Fig. 1E), but a large number of PV antigen-positive cells were detected in those of the PVR-tg/Ifnar knockout mice (Fig. 1F). These results indicate that PV can replicate in these sites if IFN signaling is disrupted.

In order to determine the IFN response in the kidneys and livers of the PVR-tg mice, we examined the expression of IFN-α, IFN-β, and Oas1a mRNAs after intravenous inoculation of PV (10⁸ PFU) (Fig. 1G to J). We detected very low levels of IFN-α and IFN-β mRNAs in the uninfected kidneys and in the kidneys at 4 h p.i. However, at 6 h p.i., we observed expression of IFN-α and IFN-β mRNAs and induction of Oas1a mRNA by 14.4-fold compared to the value for the uninfected mice, suggesting that an antiviral state had been established by that time (Fig. 1G and H). In the liver, expression of IFN-α and IFN-β mRNAs and induction of Oas1a mRNA were clearly detected at 6 h p.i. (Fig. 1I and J). It should be noted that both kidneys and liver establish the antiviral state within 6 h p.i. Considering the number of cells in the living animal and the amount of PV inoculated, infection in the tissues must occur at a very low MOI. Therefore, only a small number of cells are infected at the beginning. IFN detected before 6 h p.i. might have been produced by these cells. Although the amounts of IFN mRNAs detected by RT-PCR are small, the IFN mRNA per infected cell must be significant. Since a single round of PV replication in the cells takes approximately 6 h, it is likely that the multiple rounds of PV replication are strongly inhibited in the kidneys and liver due to the IFN response.

**PV susceptibility and IFN response in kidney cells in vitro.** We prepared primary cultured cells by dispersing the mouse kidneys with trypsin. Approximately 6 × 10⁸ viable cells per mouse, as judged by trypsin blue staining, were obtained after trypsinization. The cells were seeded in plates at a density of 3 × 10⁴ cells/cm². Approximately 1% of the cells began to grow and formed a monolayer 1 week later. As shown in Fig. 2B, C, and D, we observed at least three kinds of cells with different morphologies. When PV was inoculated into cultured cells at an MOI of 0.001, the PV titer increased rapidly (Fig. 2A), and all types of cells showed CPE by 24 h p.i. (Fig. 2E and F). When cells were infected with PV at an MOI of 10, rounding and detachment of the cells were observed at 6 h p.i. The results indicate that the kidney cells derived from the PVR-tg mice were susceptible to PV, although the kidneys in vivo are not the sites of PV replication. This result further shows that the kidneys of the PVR-tg mice acquire PV susceptibility after cultivation in vitro, which is the same as monkey kidneys. We then prepared kidney cells in culture from PVR-tg/Ifnar knockout mice. The increase in the PV titer and the appearance of CPE in the kidney cells derived from PVR-tg/Ifnar knockout mice were indistinguishable from those in the kidney cells derived from wild-type PVR-tg mice (Fig. 2A, E, and F). This, in turn, suggests that replication of PV in kidney cells in culture derived from wild-type PVR-tg mice is not affected by the IFN response.

In order to confirm the conclusions mentioned above, we infected kidney cells with PV at an MOI of 10 and measured the IFN response (Fig. 2G and H). We detected very low levels of IFN-α and IFN-β mRNAs in uninfected kidneys and in the kidneys at 4 h p.i. At 6 h p.i., expression of IFN-β mRNA increased slightly. However, we did not detect induction of Oas1a mRNA or IFN activity in the supernatants of the infected cells by a standard IFN assay (data not shown). It is
possible that the amounts of IFN-α and IFN-β mRNAs detected at 6 h p.i. were not sufficient to bring about ISG induction. Alternatively, it is also likely that the IFN mRNA detected at 6 h p.i. and later was not translated or secreted as IFN protein, since the shutoff of host protein synthesis begins to inhibit the translation of IFN mRNAs approximately 2 h after PV infection (9) and the PV 3A protein disrupts normal ER-Golgi trafficking, preventing the secretion of cytokines (4, 6).
FIG. 2. PV replication and IFN response in cultured kidney cells. (A) Time course of PV titers in cultured kidney cells. Primary cultured kidney cells from PVR-tg mice and PVR-tg/Ifnar knockout (KO) mice were infected with PV at an MOI of 0.001. At each time point, the cells were disrupted by three cycles of freezing and thawing, and virus titers were determined by plaque assay. The PV propagation profiles in the two types of cell were indistinguishable. (B to D) Morphology of uninfected primary cultured kidney cells. At least three different kinds of cells were present in the uninfected culture. (E and F) Kidney cells from PVR-tg mice (E) and from PVR-tg/Ifnar knockout mice (F) were infected with PV at an MOI of 1.0. CPE was observed at 24 h p.i. (G and H) IFN response in cultured kidney cells from PVR-tg mice. The cells were infected with PV at an MOI of 10. Total RNA of the cells was isolated at the indicated times p.i., and the IFN-α/β (G) and OAS1a (H) mRNA levels were determined by quantitative real-time PCR analysis. The asterisks indicate a significant difference (*P < 0.05; Student’s t test) in comparison with the uninfected (noninf.) samples. The error bars indicate SEM. Note that the vertical scales in panels G and H are different from those in Fig. 1G to J.

Thus, they did not contribute to establishing the antiviral state. We therefore postulate that the acquisition of PV susceptibility in the kidney cells after cultivation is due to the loss of IFN response during the process of cultivation.

PV susceptibility in liver cells in vitro. We then prepared liver cells in culture by collagenase treatment. We obtained 1 × 10⁷ viable liver cells per mouse and plated them onto plastic plates at a density of 1.25 × 10⁵ cells/cm². The live cells readily attached to the plates and could be maintained for several days, after which the cells were deformed and died. Based on the morphology, most of the cells were hepatocytes, and other cells were not included in the culture (Fig. 3B and D). We then compared the PV susceptibilities of liver cells derived from PVR-tg mice and of those derived from PVR-tg/Ifnar knockout mice. The cells from wild-type PVR-tg mice also acquired susceptibility to PV but, unexpectedly, were not fully susceptible. After PV inoculation at an MOI of 0.001, the titer in the liver cells derived from wild-type PVR-tg mice increased much more slowly than in cultured kidney cells and other cell lines (compare Fig. 2A, 3A, and 4A). Cells with CPE were rarely observed at 24 h p.i. (Fig. 3C). The PV titer continued to increase until 72 h p.i. We stopped observation at that time, since the cells were starting to deteriorate, and therefore, it was difficult to distinguish cellular damage from CPE caused by PV infection. On the other hand, PV replication in the cells from PVR-tg/Ifnar knockout mice was as efficient as in the cultured kidney cells and other cell lines (Fig. 2A, 3A, and 4A). The infected cells showed CPE with shrinkage and loss of the nuclear membrane at 24 h p.i. (Fig. 3E). When the cells of PVR-tg mice were infected with PV at an MOI of 50, the titer reached a plateau at 6 h p.i., but it was approximately 10 times lower than that in liver cells of PVR-tg/Ifnar knockout mice (data not shown). These results indicate that the liver cells derived from wild-type PVR-tg mice are partially resistant to PV.

We then examined the IFN response in primary cultured liver cells of PVR-tg mice infected with PV at an MOI of 10. Significant levels of IFN-α and IFN-β mRNAs were observed at 4 h p.i., and induction of OAS1a mRNA was observed at 6 h p.i. (Fig. 3F and 3G). It is worth noting that IFN response occurred in liver cells as early as 6 h p.i. These results suggest that the primary cultured liver cells derived from PVR-tg mice still retain the capability for a rapid IFN response, as they did in vivo. Although the amounts of IFN mRNAs detected in the liver cells in culture are larger than those detected in the liver in vivo (Fig. 1I), this does not necessarily mean that IFN inducibility in the cultured cells is higher than that in the liver in vivo. Since the cells were infected with PV at an MOI of 10 in the in vitro experiment, the IFN mRNAs produced per infected cell may be lower than those in vivo. It is possible that this early response contributes to the partial resistance of the liver cells to PV infection.

IFN induction in primate cells after PV infection. The above-mentioned results suggested that primary cultured mouse cells were susceptible to PV infection when they could not establish the antiviral state mediated by rapid IFN response. We therefore investigated whether this is true for the cultured AGMK cells and several other cell lines. AGMK cells at the third to fifth passages from the primary culture were used in the experiments. JVK-03 is a cell line spontaneously immortalized from a primary culture of AGMK cells (30).
AGMK cells, JVK-03 cells, COS-7 cells, HEK293 cells, HepG2 cells, and HeLa cells were tested for PV propagation and IFN response. All the cells were highly susceptible, and PV infection spread easily when the cells were infected at an MOI of 0.001 (Fig. 4A). Almost no IFN-α/H9251 or IFN-β/H9252 mRNA expression was detected within 6 h after PV infection at MOIs of 0.001, 0.1, and 1 (data not shown). An increase in OAS1 mRNA levels was not observed upon PV infection at an MOI of 10 (Fig. 4B) or at the different MOIs (data not shown). The results suggest that these cells do not respond to PV rapidly or do not respond at all.

Reduced expression levels of ISGs in kidney cells. We hypothesized that kidney cells have lost the rapid inducibility of IFNs because of the decrease in the expression levels of genes involved in IFN production. We compared the expression levels of various genes in the kidneys in vivo and kidney cells in

**FIG. 3.** PV replication and IFN response in cultured liver cells. (A) Time course of PV titers in cultured liver cells. Primary cultured liver cells from PVR-tg mice and PVR-tg/Ifnar knockout (KO) mice were infected with PV at an MOI of 0.001. At each time point, PV titers were determined by plaque assay. Note that the titer in the cells from PVR-tg mice increased much more slowly than that in cells from PVR-tg/Ifnar knockout mice. (B to E) Morphology of primary cultured liver cells. Liver cells from PVR-tg mice (B and C) and from PVR-tg/Ifnar knockout mice (D and E). Uninfected liver cells are monotonic (B and D). The cells were infected with PV at an MOI of 1.0 and observed at 24 h p.i. (C and E). The CPE was rarely observed in the liver cells from PVR-tg mice (C) but was clearly observed in liver cells from PVR-tg/Ifnar knockout mice (E). (F and G) IFN response in liver cells from PVR-tg mice. The cells were infected with PV at an MOI of 10. Total RNA of the cells was isolated, and the IFN-α/β (F) and OAS1α (G) mRNA levels were determined by quantitative real-time PCR analysis. The asterisks indicate a significant difference (P < 0.05; Student's t test) in comparison with the uninfected (noninf.) samples. The SEM are indicated by vertical bars. Note that the vertical scales in panels F and G are different from those in Fig. 1G to 1J.

**FIG. 4.** PV replication and IFN response in cultured cells. (A) Time course of PV titers in AGMK cells and primate cell lines. AGMK cells, JVK-03 cells, COS-7 cells, HEK293 cells, HepG2 cells, and HeLa cells were tested for PV propagation and IFN response. All the cells were highly susceptible, and PV infection spread easily when the cells were infected at an MOI of 0.001. The cells were disrupted at the indicated time points, and PV titers were determined. (B) Expression of OAS1 mRNA in infected cells. Cells were infected with PV at an MOI of 10. At 4 and 6 h p.i., total RNA was isolated from the cells, and mRNA levels were determined. Note that OAS1 mRNA levels did not change during the course of infection in all cells. The SEM are indicated by vertical bars. noninf., uninfected.
vitro by RT-PCR. RIG-I, MDA-5, STAT-1, STAT-2, IRF-7, and IRF-9 are involved in the IFN response and are induced by IFNs (25, 33, 36, 54, 65). As shown in Fig. 5A, the levels of MDA-5, STAT-1, STAT-2, IRF-7, and IRF-9 mRNAs were reduced to less than 50% of their original levels after cultivation. TBK-1, IKKε, IRF-3, and IFNAR-1 are also involved in the IFN response (41, 57, 66) but are not IFN inducible. Their expression levels, and that of a housekeeping gene, the
FIG. 6. Effect of priming on the susceptibility of cultured mouse kidney cells. (A) Experimental procedure. Kidney cells were pretreated with IFN-β at the indicated doses for 6 h. The cells were washed with MEM three times and challenged with PV with or without anti-IFN-α and -β antibodies (Abs) to neutralize the effects of newly synthesized IFNs. The cells were tested for survival, PV titers, and IFN response at the indicated time points. (B) Resistance of mouse kidney cells to PV infection after pretreatment with IFN-β. The cells in duplicate were treated with IFN-β at the indicated doses (left column). The cells were then infected with PV at an MOI of 0.1 for 3 days, and surviving cells were visualized by staining them with Amido black 10B. The IFN activities in the culture medium at 72 h after challenge were measured by the CPE dye uptake method (right). Mean values (± SEM) of two wells are indicated. Note that IFNs were induced only in the pretreated cells. noninf., uninfected. (C) Time course of PV replication in pretreated kidney cells. The cells pretreated with 1 IU/ml of IFN-β were infected with PV at an MOI of 0.001 in the presence (+) or absence (−) of anti-IFN antibodies, and the PV titer was determined at each time point. Open circles indicate the PV titer of untreated kidney cells infected at the same MOI. Note that the PV titer did not increase after 48 h p.i. if the anti-IFN antibodies...
GAPDH gene, did not show considerable change. Notably, a decrease in the level of IRF-7 during cultivation was most evident in the kidneys. Similarly, in the liver, the expression levels of IFN-inducible mRNAs also decreased (Fig. 5B). RIG-I, MDA-5, STAT-1, STAT-2, and IRF-7 mRNA levels decreased to less than 50% after cultivation. The decrease in the expression levels of these genes may be responsible for the acquisition of susceptibility.

With regard to the ITAFs and PVR, we directly measured the mRNA levels of PTB, La, PCBP-2, UNR, and PVR and compared these expression levels in the kidneys and livers of PVR-tg mice in vivo and in vitro (Fig. 5A and B). The expression levels of these mRNAs in vivo and in vitro did not differ significantly. In situ hybridization and immuno-fluorescence studies revealed that PVR mRNA is not expressed in all cells in the kidneys of PVR-tg mice in vivo (23, 50). However, all the primary cultured cells are susceptible to PV infection. We therefore could not exclude the possibility that additional expression of PVR in the kidney cells that had no PVR expression in the kidneys in vivo also occurred during the cultivation process.

In addition, we determined the levels of ISGs involved in the IFN response in primate cells. The human TaqMan probes worked sufficiently in measuring mRNA for these genes of the African green monkey. Since it was not possible to compare the expression levels of ISGs before and after cultivation directly, we used human kidney and liver RNAs that were commercially available as controls. The mRNA levels for each ISG in AGMK cells, JVK-03 cells, COS-7 cells, and HepG2 cells were different from cell to cell. However, in general, they were lower than those of the human kidney and liver in vivo (Fig. 5C). In the HEK293 cells and HeLa cells, IRF-7 mRNA levels were much higher than or similar to those in human kidneys. In these cells, however, the levels of other ISG mRNAs tested were lower than those in the human kidneys. The fact that HEK293 cells and HeLa cells failed to induce rapid IFN response upon PV infection suggested that high-level expression of IRF-7 alone was insufficient for rapid IFN response. Rapid and robust IFN response was not achieved unless a positive feedback loop of IFN response was formed (60). This idea is supported by the experimental results showing that depletion of the gene products involved in the IFN response using RNA interference or dominant-negative techniques resulted in a decrease in IFN response (5, 64). Our results also suggest that the shortage of one of the ISGs may be rate limiting for the whole IFN response.

Expression levels of ITAF mRNAs in cultured human and monkey cells were also determined. The expression levels of ITAF mRNAs were not greatly different from cell to cell (Fig. 5C). The expression levels of La autoantigen, PCBP-2, and UNR mRNAs were almost the same as those in human kidneys in vivo. The levels of PTB mRNA were approximately twofold higher than those in human kidneys in vivo. However, the absolute values for PTB mRNA levels were the same as those for PTB mRNA levels in mouse kidneys and livers in vivo (4,000 to 10,000 copies/10^7 copies of 18S rRNA). Thus, the PTB levels observed in the cultured cells were not exceptionally high. We therefore consider that the changes in mRNA levels for La, PCBP-2, PTB, and UNR do not contribute to the acquisition of susceptibility to PV infection in these cells in culture. We could not measure the PVR mRNA levels for monkey cells by the TaqMan probe designed for the human PVR gene. In the HEK293, HepG2, and HeLa cells, the PVR mRNA levels were not significantly higher than those in the human kidney or liver. Thus, the acquisition of PV susceptibility is associated with a decrease in ISG expression levels. These results strongly suggest that the kidney cells in culture, as well as many cell lines of primate origin, have lost rapid IFN response due to the loss of factors that regulate this response.

Effect of priming of mouse kidney cells with IFN. We tested if pretreatment of mouse kidney cells with a small amount of IFN reconverted the cells to PV resistance by acquisition of rapid IFN inducibility. The experimental procedures are shown in Fig. 6A. Kidney cells were treated with various concentrations of recombinant mouse IFN-β for 6 h. They were then infected with PV at an MOI of 0.1. The culture medium was removed for IFN assay, and the cells were fixed with 4% paraformaldehyde at 3 days p.i. and stained with Amido black 10B. The cells were infected with more than 0.3 IU of IFN-β protected from PV infection (Fig. 6B). Consistent with the protection phenomenon, IFN activity was observed in the culture medium of the protected cells (Fig. 6B). Similarly, cells were protected when they were infected with PV at an MOI of 1 (data not shown). The infected cells had been alive for more than a week on the day uninfected control cells died because of overgrowth. However, the cells were killed by PV infection at an MOI of 10 (data not shown). The cells pretreated with IFNs became resistant to PV infection, except when they were infected at a very high MOI.

We compared the virus growth in the pretreated (1 IU/ml for 6 h) and untreated kidney cells after PV infection at an MOI of 0.001 (Fig. 6C), and the viabilities of the cells were monitored by Amido black 10B staining (Fig. 6C, bottom). PV replication was greatly inhibited in the pretreated cells compared to the untreated cells. Notably, the increase in PV titer was not observed after 48 to 72 h p.i. (Fig. 6C, middle), despite the presence of many uninfected cells (Fig. 6C, bottom). This indicated that the infection did not spread after 48 h p.i.,
possibly because newly synthesized IFNs accumulated in the culture medium. Pretreated cells were also infected in the presence of anti-IFN antibodies. In the presence of anti-IFN antibodies, the PV titer increased over time, indicating that the PV infection could spread. In parallel with the increase in PV titer, protection was not observed after 48 h p.i. The results suggest that the pretreatment of IFN-β confers rapid IFN inducibility on the cells and that the antiviral state is maintained by the newly produced IFNs. We then determined the mRNA levels of the IFNs and ISGs after pretreatment with 1 IU of IFN-β and subsequent PV infection. As expected, all the ISG mRNA levels increased due to the pretreatment and became higher than the mRNA levels of ISGs in the kidney in vivo (Fig. 6D). Furthermore, induction of IFN-α and IFN-β mRNAs at high levels was detected as early as 4 h p.i. (Fig. 6E). Consistent with this result, IFN activity was detected in the supernatant of PV-infected cells at 6 h p.i. (data not shown). OAS1a mRNA was also induced at high levels by the pretreatment, and the levels were maintained during the infection period (Fig. 6F). OAS1a mRNA levels in the pretreated cells were not maintained unless IFN was continuously supplied. In fact, the levels decreased over time after removal of IFN-β without PV infection or after PV infection in the presence of anti-IFN antibodies (Fig. 6F). This suggests that the expression levels are maintained by newly synthesized IFNs during this period. All these results indicate that loss of rapid IFN response after cultivation in kidney cells is reversible and that primed kidney cells restore rapid IFN inducibility and resistance to PV infection.

**DISCUSSION**

In this study, we demonstrated that the change in susceptibility of cells during the process of cultivation in vitro is controlled most strongly by the level of IFN response. Kaufer and Racaniello and Ren and Racaniello predicted that a factor(s) other than PVR is needed for the acquisition of PV susceptibility after cultivation. They also showed that the PV IRES is functional in the mouse kidney in vivo, suggesting that a factor(s) other than ITAFs is important for the change in susceptibility (26, 50). In the cases of kidney cells and primate cell lines, rapid IFN response is lost in association with a decrease in at least some ISG expression. Although it is obvious that the expression of PVR and ITAFs influences the pathogenicity of PV in vivo in some specific situations, it did not change significantly during the cultivation process, and thus, it may not have a strong influence on this phenomenon.

PV is a highly cytopathogenic virus, and the infected cells are destroyed by lytic replication of the virus. PV has several mechanisms to inhibit IFN action. The 2A protease cleaves eIF-4G, leading to the suppression of cap-dependent protein synthesis (9). Host proteins, including IFNs, cannot be translated after shutoff. This shutoff phenomenon occurs as early as approximately 2 h p.i. in HeLa cells. The 3C protease cleaves the p65-RelA subunit of the NF-κB complex (42). Since the IFN-β gene is also positively regulated by NF-κB through positive regulatory domain II on the IFN-β gene promoter (10, 34, 62), cleavage of p65-RelA suppresses the NF-κB-mediated expression of IFN-β. Cleavage of p65-RelA is observed 3 h p.i. in HeLa cells. The 3A gene product inhibits the trafficking of secreted proteins, including IFNs (4, 6). The PV-infected cells in monolayer cultures either do not produce a large amount of IFNs or do not produce them at all (59) (Fig. 2G), probably due to inhibition by these viral proteins. Because of these anti-IFN effects of PV, the effects of IFNs on susceptibility to PV have not been taken into consideration.

As the inhibition by PV proteins operates within a few hours p.i., host cell defense must operate rapidly. It is known that treatment with a small dose of IFNs can enhance the amount of IFN produced and bring about IFN production at an earlier time in many cell lines. This phenomenon is known as priming (59). Using embryonic fibroblasts, Hata et al., Sato et al., and Taniguchi and Takaoka have shown that cells expressing very low levels of IRF-7 cannot induce a rapid and strong IFN response upon virus infection, while the cells expressing certain levels of IRF-7, possibly together with other ISGs, by a spontaneous weak IFN stimulation are able to form a positive-feedback loop of IFN action and induce a rapid and strong IFN response (Fig. 7A and B) (16, 54, 60). The former and the latter states may correspond to the “unprimed state” and “primed state,” respectively. If PV infection occurs in the unprimed cells, the viral proteins can accumulate to the threshold levels quickly, IFN action is strongly inhibited by the viral proteins, and the cells will not be able to establish the antiviral state (Fig. 7A). Conversely, if PV infection occurs in the primed cells, the ISGs have already accumulated above the threshold level before infection, and a rapid and strong IFN response will occur, strengthening the cellular antiviral state (Fig. 7B). Since both expression of viral proteins after PV infection and expression of ISG induction by IFNs occur rapidly, which of them gains the initiative first is very important in determining susceptibility. The cells must be primed to produce IFNs in response to PV infection (Fig. 6) (24). From this point of view, “primed or not primed” is a very important factor in determining the fate of the cells and tissues.

We previously showed that the cells in some extraneural tissues are normally protected by the IFN response (21). The cells in the extraneural tissues in vivo expressed ISGs to a certain degree before PV infection. Expression of OASs and PKR may inhibit viral replication. In addition, ISGs that are involved in IFN production, like RIG-I and IRF-7, support a rapid and strong IFN response. Therefore, it is possible to consider that cells in the extraneural tissues in vivo are normally in the “primed state.” Indeed, constitutive expression of IFNs under physiological conditions is known in humans (13, 61). It is likely that the cells in the extraneural tissues may be exposed to the constitutive IFNs. This weak IFN signal may be important for maintaining the basal ISG levels (60). On the other hand, down-regulation of ISG expression in cultured cells occurred during the process of cultivation (Fig. 5). The acquisition of susceptibility was associated with this decrease in expression of ISGs. Priming of the kidney cells with a small amount of IFN increased the basal ISG levels (Fig. 6C). This, in turn, restored rapid IFN response and resistance to PV (Fig. 6). The results suggest that cultured cells in vitro do not receive constitutive IFNs or do not produce constitutive IFNs by themselves. Therefore, we consider that the acquisition of PV susceptibility by cells in vitro is due to the transition of the cells from the “primed state” to the “unprimed state.”

PV exhibits a paradoxical tropism. The replication sites of
PV in vivo are restricted to a few tissues, including the CNS. However, PV is able to propagate in monolayer cultures derived from almost any tissue of primates. By taking the IFN response of the cells and tissues into consideration, it is now possible to explain this situation. The behavior of the cells and tissues after PV infection is largely dependent on the expression levels of ISGs, namely, the degree of priming. Infection in the tissues occurs at a very low MOI in natural infection in humans and experimental infection in tg mice in vivo. Therefore, initial infection in the tissues occurs in only a small number of cells (Fig. 1B). In the tissues where the cells are primed, these small numbers of infected cells become victims to produce IFNs rapidly (Fig. 1G and H). PV cannot spread easily, and severe pathological lesions are not produced, since the surrounding cells receive IFNs and strengthen the antiviral state before they are exposed to PV (Fig. 7D). Thus, the IFN response prevents the chain of events leading to massive infection. This was also true for the primed kidney cells in vitro (Fig. 6). On the other hand, in susceptible tissues, PV will spread from the initial infectious center to surrounding cells.
because the IFN response is not sufficient (Fig. 7C). Similarly, PV can spread very easily in cultured cells because they are not in the primed state (Fig. 2). Therefore, the IFN response has a strong influence on the susceptibility of the cells both in vivo and in vitro. The paradox is now elucidated, since both tissue and cell specificities of PV infection are regulated, at least in part, by the same mechanism mediated by the IFN response.

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