NOTES

JC Virus Regulatory Region Tandem Repeats in Plasma and Central Nervous System Isolates Correlate with Poor Clinical Outcome in Patients with Progressive Multifocal Leukoencephalopathy

LUZ-ANDREA PFISTER,1 NORMAN L. LETVIN, 1 AND IGOR J. KORALNIK1,2*

Division of Viral Pathogenesis, Department of Medicine,1 and Department of Neurology,2 Beth Israel Deaconess Medical Center, Harvard Medical School, Boston, Massachusetts 02215

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JC virus (JCV), the causative agent of progressive multifocal leukoencephalopathy (PML), has a hypervariable regulatory region (JCV RR). A conserved archetype form is found in the urines of healthy and immunocompromised individuals, whereas forms with tandem repeats and deletions are found in the brains of PML patients. Type I JCV RR, seen in MAD-1, the first sequenced strain of JCV, contains two 98-bp tandem repeats each containing a TATA box. Type II JCV RR has additional 23-bp and 66-bp inserts or fragments thereof and only one TATA box. We cloned and sequenced JCV RR from different anatomic compartments of PML patients and controls and correlated our findings with the patients’ clinical outcome. Twenty-three different sequences were defined in 198 clones obtained from 16 patients. All 104 clones with tandem repeats were type II JCV RR. Patients with poor clinical outcome had high proportions of JCV RR clones with both tandem repeats in plasma (54%) and brain or cerebrospinal fluid (85%). In those who became survivors of PML, archetype sequences predominated in these anatomic compartments (75% and 100%, respectively). In patients with advanced human immunodeficiency virus infection without PML, only 8% of JCV RR clones obtained in the plasma contained tandem repeats. These data suggest that the presence of tandem repeats in plasma and CNS JCV RR clones is associated with poor clinical outcome in patients with PML.

The human polyomavirus JC virus (JCV), the etiologic agent of progressive multifocal leukoencephalopathy (PML), is highly conserved in healthy and immunocompromised individuals and has been called the archetype (1, 20, 32). The archetype has no duplications and contains a 23-bp insert and a 66-bp insert localized at nucleotides (nt) 36 and 92 compared to the first sequenced brain isolate of JCV, MAD-1. A hypervariable form of JCV RR is found in the brain and cerebrospinal fluid (CSF) of PML patients. In the MAD-1 isolate, the RR contains two identical 98-bp tandem repeats.

It has been hypothesized that rearrangements of JCV RR occur in the setting of immunosuppression, leading to JC viremia, hematogenous spread of the virus to the central nervous system (CNS), and the development of PML. Indeed, JC virus DNA is rarely found in the blood of healthy individuals, but it becomes more readily detectable in the blood of human immunodeficiency virus (HIV)-infected people who have <200 CD4+ T cells/µl of blood (16). Analyses of RR from JCV isolates obtained from blood samples have been limited and, for the most part, restricted to direct sequencing of PCR products (2, 5, 26, 27). In addition, the relationship between the types of JCV RR in the blood and disease progression has not been investigated.

The clinical course of PML is variable. Some patients have a fulminant evolution and die within 1 to 6 months of their diagnosis, whereas others have a protracted course and become PML survivors. We sought to determine the variability of JCV RR in the blood and CNS and if patterns of JCV RR found in the CNS arise in the blood. Moreover, we sought to correlate our findings with the patients’ clinical outcome.

Urine, plasma, CSF, or brain samples were obtained from 16 patients. Of these patients, eight who were HIV positive (HIV+) and two who were HIV negative (HIV−) died from PML. One HIV+ patient and one HIV− patient were PML survivors and were still alive more than 2 years after the diagnosis of their neurologic disease. Four patients were HIV+ with other neurological diseases (HIV+/OND) including HIV encephalopathy, cytomegalovirus polyradiculopathy, and other non-PML leukoencephalopathies. These patients were also long-term survivors of their neurologic diseases.

DNA extraction from CSF, plasma, urine, and brain was performed as previously described (16). PCR amplification was performed using the external primers JCRS (5′-ATTAGTGCG...
We first characterized JCV RR in various anatomic compartments in PML progressors, PML survivors, and HIV+ OND patients. Analysis of CSF and/or brain samples from the eight HIV+ patients who died of PML (Fig. 1A) showed that six of them had a JCV RR with two 98-bp tandem repeats (patients no. 1 through 6). All contained 23- and/or 66-bp inserts (at positions nt 36 and 134 and positions nt 92 and 190, respectively) which had variable deletions. In contrast, two patients had only archetype-like JCV RR with one single 98-bp unit (patients no. 7 and 8). Surprisingly, one patient (no. 1) also had an archetype form of JCV RR in 1 of 12 CSF clones sequenced.

In four of these HIV+ PML progressors, the JCV RR could be amplified and cloned from plasma samples. Clones were obtained from one of these individuals (no. 1) that had tandem repeats identical to his most prominent CSF JCV RR, as well as a distinct archetype-like JCV RR. Patient no. 4 had clones with only a tandem repeat JCV RR, identical to the one found in his CSF. Patients no. 7 and 8 had archetype-like JCV RR. Patient no. 7 had two distinct plasma JCV RR that were different from the one found in his brain, whereas the single pattern found in the plasma of patient no. 8 was identical to the CSF sequence. Urine specimens were analyzed for three of these patients. Two had archetype (no. 1 and 2) and one had tandem repeat (no. 6) JCV RR.

For the two HIV− PML progressor patients (Fig. 1B), plasma could be analyzed from one and brain from the other. We obtained two distinct sequences from the plasma of patient no. 9 and one type of sequence from the brain of patient no. 10. In all of them, tandem repeats were present in the JCV RR.

CSF and plasma were analyzed from one HIV+ PML survivor (no. 11, Fig. 1C). In both samples, we found an archetype JCV RR with a deletion of 40 bp (nt 208 to 247) downstream of the 98-bp unit. This deleted region encodes a nuclear factor 1 (NF-1) binding site. The analysis of plasma from one HIV+ PML survivor (no. 12) showed JCV RR with tandem repeats (Fig. 1C).

Plasma samples were analyzed from four HIV− individuals with neurologic disorders other than PML (Fig. 1D). One of them, no. 13, had an archetype JCV RR. Two others, no. 14 and no. 16, had an archetype-like JCV RR, and no. 15 had two different sequences, an archetype-like JCV RR and a JCV RR with tandem repeats. These patients had undetectable JCV DNA in their CSF. Interestingly, the sequence found in the plasma of patient no. 14, which has a 10-bp deletion at the 3′ end of the 23-bp insert, is identical to that of a strain (PNG-1A) recently identified in the urine sample from a healthy individual from Papua New Guinea. Similar 10-bp deletions were also found in the same region in a few Central and East African strains (24). Patient no. 14 is originally from Haiti.

We then correlated the patterns of JCV RR in different anatomic compartments with the patients’ clinical outcomes. Patients with poor clinical outcomes had a high proportion of tandem repeats in the JCV RR in the plasma (54%) and CNS (85%) (Table 1). In contrast, PML survivors had a high proportion of archetype JCV RR in the plasma (75%) and CNS (100%). Finally, HIV+ individuals with low CD4+ T-cell counts without PML had a low proportion of tandem repeats in JCV RR in the plasma (8%).

These sequence data are in agreement with previous reports (2, 23, 31) demonstrating that JCV RR is hypervariable in the
FIG. 1. Sequencing results of the JCV RR. On the top is a representation of the MAD-1 and archetype JCV RR. The nucleotide numbers are based on the prototype MAD-1 sequence. The known transcription regulation factor binding sites are indicated and include the lytic control element (Lytic E), nuclear factor 1 (NF-1), and c-Jun. Each 98-bp unit is represented by an open box. The TATA box is represented by TATA. The archetype contains only one 98-bp unit with two inserts. Black box, 23-bp insert; checkered box, 66-bp insert; dotted lines, deletions in the 98-bp units or in the 23- or 66-bp inserts; grey box, region downstream of the 98-bp units. Some clones contain fragments of this region inserted.
body. We isolated two to four distinct JCV RR in 6 of 12 PML patients and in 1 of 4 patients without PML. These six PML patients were progressors. The presence of dual infection with different JCV strains was also detected in 6 of 21 (28.6%) PML patients in a recent study of JCV genotype, and this was found to be an additional risk factor for the development of PML (8).

The precise anatomic compartment in which tandem repeats first appear has not been unequivocally determined, since they can be found both in blood and in CNS samples. It is also unclear if these tandem repeats are the cause or the consequence of JCV neurotropism. Patient no. 1 was the only one for whom it was possible to amplify, clone, and sequence JCV RR from four different anatomic compartments, (brain, CSF, plasma, and urine). We obtained four different JCV RR for this particular patient and saw an evolutionary gradient from archetype to tandem repeat going from the urine to the blood and then to the CNS. For the four HIV+ PML progressors whose plasma and CSF or brain samples could be analyzed (patients 1, 4, 7, and 8), JCV RR in the CNS samples had deletions of the 23- and 66-bp inserts in numbers either similar to or greater than those of the plasma (Fig. 1A). These data suggest that tandem repeats first originate in the plasma and that further rearrangements may occur once the virus enters the CNS.

Archetype JCV RR is usually not found in the CSF of PML patients. However, 1 of the 12 clones obtained from the CSF of HIV+ PML progressor patient no. 1 showed an archetype JCV RR while 10 of 10 clones from HIV+ PML survivor patient no. 11 had this sequence. In addition, HIV+ PML progressors no. 7 and no. 8 had archetype-like JCV RR in the brain or CSF. To our knowledge, this is the first report of fatal PML cases in which no tandem repeats were found in the CNS. Finally, one HIV+ PML progressor had a tandem repeat JCV RR isolated from the urine. Therefore, the dogma suggesting that archetype JCV RR is present only in the urine and that tandem repeats are restricted to the CNS is not absolute.

We also aimed to determine if additions or deletions of known protein binding sites in JCV RR correlated with clinical outcomes. Complex rearrangements of the JCV RR with additional inserts containing new predicted NF-1 or c-Jun binding sites were found only in patients with fatal outcomes (no. 1, 3, and 6). In contrast, deletions of predicted NF-1 or c-Jun binding sites were found only in patients with good outcomes (no. 11 and 14). NF-1 is a family of proteins, and the one that specifically binds to the JCV promoter is the NF-1 class D. This protein is predominantly expressed in glial cells (25).

Most of the studies of JCV promoter activity have used the prototype MAD-1 regulatory region, which was the first JCV that was sequenced and has no 23-bp or 66-bp inserts. This JCV RR pattern was called type I. Subsequent studies have shown that the vast majority of JCV strains isolated in vivo contain these inserts or fragments thereof, as well as a deletion of the second TATA box. Such JCV RR were called type II (12, 22). The 23-bp insert contains a binding site for the transcription factor SP-1 (13). The promoters of the oligodendrocyte-specific cellular genes, myelin basic protein and proteolipid protein, contain similar binding sites. Together with the paired NF-1 and c-Jun binding sites, the SP-1 binding site in the 23-bp insert is part of a motif that is conserved between several glial-specific promoters (13). These findings are in agreement with in vitro studies demonstrating that the number of NF-1 binding sites is directly proportional to the level of viral transcription in glial cell lines (18, 19).

The function of the 66-bp insert has not yet been elucidated. No predicted binding sites of transcription regulators have been found in this region. Such sites have also not been shown in the region where it is inserted. Interestingly, deletions of \(\geq 85\%\) of the 66-bp insert correlated in our study with a fatal outcome. In contrast, the complete deletion of the 66-bp insert did not correlate with the clinical outcome.

The present study shows that PML progressors have a higher frequency of archetype-like and tandem repeat JCV RR in the

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<th>Clinical outcome and/or patient characteristic</th>
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<td>9</td>
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<td>1</td>
<td>10</td>
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<td>24</td>
<td>8 (33)</td>
<td>14 (58)</td>
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a A, archetype; AL, archetype like; TR, tandem repeats.
plasma and CNS than do PML survivors. In these latter patients, archetype JCV RR predominates. In the present series of patients, the number of PML progressors studied was larger than the number of PML survivors. This is due to the fact that PML survivors are quite rare and account only for approximately 10% of all patients with this disease. They also usually have a lower JC virus load in the CSF and undetectable JCV DNA in the blood. CSF JCV DNA often becomes undetectable following initiation of antiretroviral treatment. These changes parallel their higher CD4+ T-lymphocyte counts. Therefore, analysis of JCV RR from these patients’ samples is often impossible. Attempts to amplify JCV RR from the plasma of two additional PML survivors were unsuccessful. However, analyses of plasma samples from four HIV+ patients without PML also showed only 8% tandem repeats.

Since MAD-1 was the first JCV strain to be isolated, the type I MAD-1 JCV RR has been employed in the study of JCV type I JCV RR was found only in three possible PML cases (28). Therefore, more emphasis should be directed to the study type I JCV RR in order of appearance in Fig. 1: patient no. 1, AF354666, AF354667, and AF354671; patient no. 2, AF354569, AF354570, and AF354571; patient no. 3, AF354572; patient no. 4, AF354573, and AF354574; patient no. 5, AF354575; patient no. 6, AF354576 and AF354577; patient no. 7, AF354578, AF354579, and AF354580; patient no. 8, AF354581; patient no. 9, AF354582 and AF354583; patient no. 10, AF354584; patient no. 11, AF354585; patient no. 12, AF354587; patient no. 13, AF354588; patient no. 14, AF354589; patient no. 15, AF354590 and AF354591; patient no. 16, AF354592.

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REFERENCES