JC virus (JCV) causes progressive multifocal leukoencephalopathy (PML), a demyelinating disease in humans. The disease, once considered fatal, is now managed with immune reconstitution therapy; however, surviving patients remain severely debilitated. Until now, there has been no animal model to study JCV in the brain, and research into treatment has relied on cell culture systems. In this issue of the JCI, Kondo and colleagues developed a mouse model in which human glial cells are engrafted into neonatal mice that are both immunodeficient and deficient for myelin basic protein. When challenged intracerebrally with JCV, these mice exhibit some of the characteristics of PML. The establishment of this chimeric mouse model is a significant advance toward understanding the mechanism of JCV pathogenesis and the identification of drugs to treat or prevent the disease.

Background
Progressive multifocal leukoencephalopathy (PML) was first described as a rapidly progressive demyelinating disease of unknown origin in the nervous system of patients with underlying immunological abnormalities (1, 2). At the time, the cause of PML was not known, but viral infection was suspected, based on the pathological finding of inclusion bodies in highly condensed oligodendrocyte nuclei. This hypothesis was supported by subsequent work demonstrating the presence of papovavirus-like particles in the nuclei of oligodendrocytes (3). In some cases, astrocytes and cells with the morphological appearance of granule cell neurons were enlarged in patient samples. Perivascular infiltration of lymphocytes and mononuclear cells were found infrequently. Thirteen years after these morphologic observations, the causative agent of PML was identified as a papovavirus and was named for the initials of the patient from whose extracts it was isolated (4). Identification of the virus was made by inoculating cultures of primary human fetal glial cells with extracts made from the brain of a PML patient at necropsy. These cultures contained a mixture of astrocytes and glial precursors, which were referred to at the time as spongioblasts. While both astrocytes and spongioblasts were affected by inoculation of the brain extracts and both populations contained virus, the spongioblasts seemed most susceptible to infection. The virus did not react with antisera against SV40, mouse polyomavirus, or human papillomavirus particles, nor was it capable of propagating in cell types known to support the growth of other papovaviruses. Cultures of primary human fetal glial cells remained the only cell culture system capable of propagating JC virus (JCV; also known as JCPyV), until cell lines derived from these tissues were established by transformation with SV40 or JCV T antigen (5, 6). These established cell lines maintained markers characteristic of astrocytes, although these cell lines have likely undergone numerous changes over time, given the ability of T antigen to induce genetic instability in transformed cells. Results from studies with these cell cultures made it clear that astrocytes and glial precursors could both support viral replication. It should be noted that none of these cultures contained oligodendrocytes, as these postmitotic cells were likely eliminated during cell passaging and the technology to derive them from glial precursors had not yet been developed.

Diagnosis and natural history of the virus
Definitive diagnosis of PML was markedly advanced in 1987 by the development of an in situ hybridization assay that used labeled JCV-specific DNA probes (7, 8). These new assays confirmed that oligodendrocytes were the major target of virus infection in the brain, as replicating JCV DNA was readily detected in these cells. Astrocytes occasionally scored positive in these assays, and in one case, cells from the bone marrow of a PML patient tested positive for the virus (9), leading to the hypothesis that lymphoid cells maytraffic virus from the periphery to the CNS to cause PML. Peripheral sites of viral persistence include the kidney and possibly bone marrow (10, 11). In the kidney, the virus has an archetypal regulatory region that predominates in this tissue, but is rarely seen in blood or in brain. Moreover, in bone marrow and blood, the virus exhibits numerous and diverse rearrangements of the regulatory region, and it is this form of the virus that is associated with CNS disease and has been referred to as PML-type JCV (12, 13). Additional mutations occur in the coding region of the major capsid protein VP1 during high-level replication of the virus in CNS tissue, and these mutations are specific for PML. Thus, persistent virus in kidney is thought to undergo numerous rearrangements that are perhaps driven by recombinase enzymes present in lymphoid cells which then traffic the virus to the CNS, where high-level replication leads to additional mutations that

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increased during the AIDS pandemic, with the disease occurring in 2%–5% of HIV-infected individuals (17). In 2005, the first cases of PML appeared in patients that were being treated with humanized monoclonal antibodies that block leukocyte trafficking into inflamed tissues for the treatment of inflammatory diseases, such as multiple sclerosis, rheumatoid arthritis, and Crohn’s disease (18–20). The rapid rise in PML incidence led many investigators to develop and test a variety drugs for efficacy against this devastating disease (21–23). As no animal models existed, these investigations were restricted to cultures of either primary or established cell lines (24–29). The lack of an animal model for PML has been a major limitation for the development of effective therapies for people who have or are at risk for developing PML.

The model

In this issue, Kondo and colleagues have succeeded in developing a mouse with humanized glia (14). This remarkable achievement was brought about by implantation of primary human glial progenitor cells into neonatal immunodeficient, myelin-deficient (Rag2−/− Mbp−/−) mice. These mice developed white matter derived from the human glial cell precursors. Intracerebral inoculation of these chimeric mice with JCV led to virus replication and spread within the human white matter tracts. In this model, JCV infects human-derived oligodendrocytes but does not readily replicate in them, instead inducing apoptosis that results in demyelination.

likely favor viral spread within the brain. The current model developed by Kondo and colleagues involves direct inoculation of the rearranged or PML-type JCV into brain parenchyma, thus bypassing the early stages of persistence and viral evolution that lead to pathogenesis (14). The fact that the authors observed and documented mutations in VP1 from the injected virus make it clear that these changes can come about after viral penetration of CNS tissue and are likely due to a high level of replication. It would be interesting to examine whether archetype virus injected into the CNS of these mice could undergo the rearrangements necessary to evolve into the pathogenic form, or whether these rearrangement events are restricted to extraneural sites of replication.

Rising incidence of PML

In 1979, the incidence of PML was estimated to be 1 in 10 million people (15, 16). The occurrence of PML dramatically
ably, the authors were able to demonstrate an evolution of viral sequences during disease progression in these mice that reflected the biology of disease in PML patients. Moreover, infection of the chimeric mice with JCV mutants derived from human patients also led to viral spread and disease (30, 31). This latter point is interesting, as mutant virus arising in human patients seems unable to recognize sialic acid–containing receptors for the virus (32, 33). These viruses must evolve to use alternative means of infection and spread, the mechanisms of which are under investigation in several labs.

The power of the model developed by Kondo et al. (14) is the ability to finally test specific drugs and other treatments for efficacy against PML. In order for drugs to be efficacious, it will be imperative for them to cross the blood–brain barrier of the mouse to inhibit viral spread and, ultimately, disease progression. The authors make a very strong argument that JCV does not replicate in oligodendrocytes, but rather kills them by inducing apoptosis. This observation is intriguing; however, it may be unique to the authors’ specific mouse model, as years of work with human tissues clearly indicates that the primary target for lytic replication of JCV in the human brain is the mature oligodendrocyte. Kondo and colleagues also demonstrated that the virus does not need oligodendrocytes to spread (14). This point is not surprising, as JCV has been propagated for years in primary or established cell cultures that do not contain oligodendrocytes. It is important to note that while the engrafted cells in this mouse model are human, all of the other cells, such as kidney and lymphoid cells, remain of mouse origin and are therefore not susceptible to infection with the virus. This limitation will preclude studies to define routes of primary infection, factors involved in the establishment of latency or persistence, and contributions of different virus strains to disease manifestations. Ultimately, this model will not be able to elucidate how the virus is spread from the periphery to the CNS. In future studies, one could imagine humanizing the immune system of these mice and, for example, investigating the hypothesis that lymphoid cells are responsible for trafficking the virus to the CNS. Although an animal system to study JCV pathogenesis from the initial site of infection through to PML remains elusive, the model generated by Kondo and colleagues is a major breakthrough in the field.

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