cerebral hemispheres, including the olfactory bulbs, septal nuclei, cingulate gyrus, hippocampus and hypothalamus. Infection of 4-week-old rats resulted in a persistent infection in the CNS. Animals developed a transient immunologically specific encephalitic response which caused bizarre, hyperactive and aggressive behavior. Rats recovered from the encephalitis with hydrocephalic pathological deficits resulting from lysis of some infected neurons. Behavior hence was characterized by inactivity and passiveness. Infection in the brain persisted. Immunosuppression of the mice resulted in very similar virus-cell interaction but no encephalitis occurred and animals did not become ill. The behavioral disease was therefore due to responses of infected neurons triggered by specifically sensitized inflammatory cells.


The behavior of 6 marmosets (Callithrix jacchus) inoculated intracerebrally with CSF from schizophrenic patients was compared over a 2½ year period with the behavior of marmosets inoculated with CSF from patients with neurodegenerative disease or from patients undergoing spinal anaesthesia who suffered from no neurological or psychiatric disorders. Animals inoculated with CSF from psychiatric or neurological patients became progressively and significantly less active than animals inoculated with CSF from other patients although, post-mortem, no consistent differences between groups of animals were found in light- or electron-microscopic brain examination, brain biochemical analysis or viral isolation.

In another experiment all 4 marmosets inoculated intracerebrally with brain material from a patient with the Gerstmann-Sträussler syndrome (G.S.S.) developed a spongiform encephalopathy which was indistinguishable from that shown by another 4 marmosets inoculated with brain material from a typical case of Creutzfeldt-Jakob disease in a comparable time course of 20-32 months. The familial occurrence of a viral disease in G.S.S. will be discussed.


To test the possibility that schizophrenia might result from a transmissible agent analogous to the CJD agent, homogenized brain from various cortical and subcortical regions of 10 patients with chronic schizophrenia was inoculated intracerebrally into 37 primates and 22 rodents. No gross behavioral peculiarities were noted in routine observations by veterinarians and animal technicians over a five year follow-up period. One squirrel monkey and one guinea pig developed ataxia and tremor at 44 and 20 months respectively. Reinoculation with original inoculum has been negative at 40 months. Nineteen rodents and 11 primates died during the follow-up period. Histopathological examination of rodent brains showed cerebellar and hippocampal gliotic changes in 5/9 experimental cases and 3/9 controls. Similar findings were not observed in primate brains which could not be differentiated from controls. While these results do not support a transmissible agent model for schizophrenia analogous to CJD, it is not conclusive evidence against a virus as certain known viral illnesses also cannot be transmitted (e.g. SSPE).

Abstract 35. Coronavirus infection in rats: induction of an autoimmune response against brain antigen H. Wege, R. Watanabe, R. Dörries, P. Massa and V. ter Meulen Institute for Virology and Immunobiology, Würzburg, West Germany
Persistent virus infections of the CNS tissue may lead to functional and structural alterations of the host cell accompanied by psychological and neurological changes. In coronavirus infection of rats a persistent infection is the basis for the development of a subacute demyelinating encephalomyelitis (SDE). Animals surviving this infection may develop a relapsing course accompanied by old and fresh demyelinating lesions. The age of the animal, the time of infection, the response of the immune system, the genetic background of the host as well as the properties of the virus mutant used for infection determine the outcome and development of the CNS disease process. Of particular interest is the finding that spleen cells from diseased rats are sensitized against myelin basic protein (MBP) and adoptive transfer of such cells causes CNS changes similar to experimental allergic encephalomyelitis (EAE). In addition, rats which develop a remission of SDE are resistant to subsequent induction of EAE when challenged with MBP. Analysis of cerebrospinal fluid from rats at different stages of the disease reveal intrathecal IgG synthesis which are in part specific for corona virus antigen. These data indicate that in the course of coronavirus infection in rats an autoimmune reaction against brain antigen develops which may be of pathogenetic importance. This phenomenon may also play a role in the development of other CNS disorders.

Abstract 36. In vivo and in vitro models of demyelinating disease: factors influencing the disease process caused by coronavirus infection of the rat
O. Sorensen, S. Beushausen, M. Coulter-Mackie, S. Dales
University of Western Ontario, London, Canada.

The coronaviruses, ubiquitous in mammals, including man, manifest serotype-related predilection for different tissues. The murine viserotropic strain is MHV3 and neurotropic in JHMV. JHMV when inoculated into neonatal rats, can cause either a rapidly fatal acute encephalomyelitis or, after longer incubation periods, a paralytic disease. High anti-JHMV IgG ratios of CSF/serum, indicative of local antibody production in the CNS, were noted only where disease was demonstrable, suggesting that local antibody production accompanied the infection but did not prevent the neurological disease. Among animals in which neurologic symptoms had not become manifest, only those with elevated CSF/serum ratios were found to have histological CNS lesions. Immunofluorescent microscopy indicated that viral antigens were present in both glia and neurons. Antigen-positive cells were frequently present in histologically normal CNS tissue, while regions of necrosis were antigen-negative. Testing for the presence of viral RNA with JHMV cDNA probes revealed that the virus was rapidly disseminated throughout the CNS, presumably establishing centers of infection prior to the development of recognizable tissue damage. Viral RNA was also detected in the CNS following recovery from paralysis and as late as 5 months post-infection, where no disease occurred. These findings indicate that, although infection by JHM virus can spread rapidly throughout the CNS, formation of lesions during chronic disease is a slower process. The current data and previous observations suggest that in rats JHMV can remain in a latent state for periods of at least several months, without apparent neurologic disease, despite the absence of any known provirus phase in the replicative strategy of coronaviruses.

Concerning the serotype specificity for explanted cells from the CNS of newborn, inbred, Wistar-Furth rats, an unambiguous tropism of MHV3 for astrocytes and JHMV for oligodendrocytes could be demonstrated. With the latter cell-virus interaction, relatively small differences in spatial density of oligodendrocytes influence profoundly the duration of persistence and virus yield.

The in vitro temporal programme of oligodendrocyte differentiation, monitored by induction of a myelin related enzyme, 2'3'-cyclic nucleotide-3' phosphohydrolase, corresponds to that occurring in vivo and is coincident with the onset of insusceptibility to disease caused by JHMV. On the basis of these data it is concluded that in-vitro interaction of JHMV with oligodendrocytes reflects accurately the in-vivo host control over the tropism and expression of this virus, thereby effecting the progressive, demyelinating disease process.