Dementia features prominently as a common disability among the elderly and is only expected to increase with the aging population of developed countries worldwide. Dementia constitutes diverse combinations of a loss of normal executive function in decision making, a decline in memory encoding and retrieval, and an impairment of the ability to perform cognitive tasks. In the first part of this 2-part review of rapidly progressive dementias, the neurodegenerative conditions that feature prominently in the differential diagnosis of a presenile or rapidly progressive dementia were discussed. These include the common, inexorable dementias typically associated with old age such as Alzheimer dementia and dementia with Lewy bodies. As with any clinical evaluation, imaging cannot replace accurate and thorough physical examination and history, but it contributes greatly to discerning entities within the differential diagnosis of a rapidly progressive dementia.

Rapidly progressive dementias can result from a variety of etiologies, and in this second review article, we discuss the remaining etiologies summarized in the differential diagnosis On-line Table cited in Part 1. One striking characteristic of the differential diagnosis for this clinical presentation of dementia is the tremendous diversity of entities encountered. This review discusses the role of neuroimaging, particularly MR imaging, in the diagnosis of a wide array of dementia-inducing conditions, including the following: prion disease such as Creutzfeldt-Jakob disease; infectious entities such as HIV-associated dementia; autoimmune and inflammatory conditions such as Hashimoto encephalopathy; malignancy-related conditions such as primary CNS lymphoma; and rare manifestations of other causes such as Wilson disease. In many of these clinical scenarios, MR imaging plays a deciding role in clarifying the clinical diagnosis. Other authors have recently reviewed, in greater detail, the clinical work-up of rapidly progressive dementias; this article will focus on the consideration of these diseases when examining neuroimaging studies of rapidly progressive dementias.

**Prion Disease**

There are several rare disorders in which the pathophysiologic entity is the prion, a misfolded form of a normally found protein within the brain, which propagates itself by promoting changes in the conformation of the normal prion protein. The most common set of conditions involving the prion is Creutzfeldt-Jakob disease, which is either inherited in a familial fashion, sporadically occurring, or acquired through exposure. CJD may fall under both the neurodegenerative and infectious classifications; the latter is thought to occur in variant CJD or new-variant CJD, related to ingestion of mad-cow-infected beef. Of these types, the spo-
radic form is the most common, followed by genetic and acquired forms. This condition may differ clinically from many other conditions in the differential on the basis of its cerebellar dysfunction and common presence of myoclonus. Nevertheless, its clinical overlap with corticobasal degeneration and progressive supranuclear palsy necessitates the use of imaging to differentiate it from other rapidly progressive dementias.

The imaging findings typical of CJD, while observed on T2-weighted and FLAIR sequences, are best seen by using high-b-value DWI and are confirmed with reciprocal changes on ADC images. DWI typically demonstrates hyperintensities within the basal ganglia, the thalamus, and cortex (Figs 1 and 2). The “hockey stick” or “pulvinar” sign, which is formed by the confluent hyperintensity within the dorsomedial thalamus and the posterior thalamus (pulvinar), is thought to be indicative of variant (infectious acquired) CJD, though these signs may be seen in other forms of CJD. These imaging changes are much more sensitive and specific than previous noninvasive clinical methods for CJD diagnosis, including the presence of CSF protein 14–3–3 and periodic sharp waves on electroencephalography or neuron-specific enolase. For this reason, imaging findings are essential in making the diagnosis of CJD, despite their not being included in earlier clinical criteria.

Infection-Related Etiologies
In younger patients with dementia symptoms, both infectious and inflammatory-related conditions are much more frequent than the neurodegenerative diseases. Encephalitis and other infections of the CNS are generally hyperacute in onset; when cognitive decline is present, these cognitive symptoms are likewise
hyperacute in onset. In addition, encephalopathic symptoms with marked confusion and neuropsychiatric manifestations also point clinically to this diagnosis, as do non-neurologic manifestations of infection such as cutaneous or respiratory presentations. Readers are referred to the work of Almeida and Lautenschlager for a thorough clinical discussion of dementia associated with infections.

Encephalitis. The diagnosis of encephalitis in the acute setting may prove problematic: Clinical symptoms are generally nonspecific and may involve a variety of neurologic symptoms or none at all. While rarely presenting with dementia acutely, rapidly progressive dementia in younger individuals merits consideration of encephalitis. Even rapid polymerase chain reaction laboratory tests of herpes simplex may have negative findings in the very early stages of encephalitis, which make ancillary work-up imperative.

MR imaging can reveal pathologic changes manifest as FLAIR and T2 hypointensity within the temporal (particularly, medial), insular, and inferior frontal cortices in patients with herpes encephalitis. In the much less common Japanese encephalitis, these abnormalities tend to be seen in subcortical gray matter regions. Patchy enhancement with gadolinium administration indicates nonspecific inflammatory changes generally seen in encephalitis. DWI augments the evaluation of encephalitis with earlier detection of changes than on conventional imaging, though it may not be as sensitive later in the clinical course. Restricted diffusion occurs in herpes encephalitis during the acute phase and appears less often in Japanese encephalitis, though other authors demonstrated diffusion changes in the thalamus in this condition. Moreover, magnetic-susceptibility-sensitive sequences (eg, gradient recalled-echo, susceptibility-weighted, and susceptibility angiography) may detect focal hemorrhage associated with sequelae of CNS infection.

HIV and AIDS-Related Dementias
The occurrence of dementia in HIV-infected individuals ranges from 1 in 10 to 1 in 3. HIV dementia may be considered a subcortical dementia with cognitive slowing, motor retardation, and impaired memory. Fortunately, successful treatment with antiretroviral therapy has the potential to ameliorate cognitive symptoms and sometimes reverse cognitive deficits in HIV dementia. For this reason, identification and establishment of an appropriate treatment regimen is of paramount importance. Imaging is also vital to ensure that no other AIDS-related CNS conditions are present, such as primary CNS lymphoma, progressive multifocal leukoencephalopathy (seen as demyelinating white matter lesions on MR imaging), or toxoplasmosis (with multiple deep gray matter lesions). With cognitive decline in the setting of HIV infection, volumetric studies observed atrophy by using MR imaging. These changes are not generally apparent on conventional MR imaging on an individual basis. DTI may, however, show reduced fractional anisotropy inversely correlated with viral loads. Magnetization transfer ratios also appear different when comparing HIV-infected patients to controls.

MR spectroscopy can also detect changes that reflect dementia symptoms in HIV-infected patients, regardless of neuroanatomic changes; a recent study by Ernst et al demonstrated reductions in parietal gray matter glutamate in subjects with HIV with cognitive deficits. Others have implicated decreased concentrations of NAA and combined glutamate and glutamine in HIV-infected patients with reductions in cortical, hippocampal, and subcortical volume, respectively.

Neuroborreliosis. CNS manifestations related to Lyme disease caused by Borrelia burgdorferi are rare; contentious; and largely related to cranial nerve involvement, polyradiculopathy, or neuropsychiatric manifestations. Acute dementia in this setting has been previously reported, and though infrequent, may place this condition on the differential list of rapid-onset dementias. An extensive review by Hildenbrand et al discussed the neuroimaging findings and the clinical syndrome of neuroborreliosis in greater detail. Rarely, encephalomyelitis may develop from spirochete infiltration into the brain, as indicated by tumefactive white matter lesions. Traditionally, more commonly seen in Lyme disease with CNS involvement are nonspecific FLAIR abnormalities predominantly within the frontal cortex; on the other hand, more recent, systematic investigation suggests that this is an uncommon occurrence with only 10% of patients with neuroborreliosis demonstrating T2 hyperintense foci. These lesions may mimic a demyelinating process; thus, in cases in which multiple sclerosis is considered a reasonable alternate explanation for T2 foci, DTI and magnetization transfer imaging are helpful in demonstrating the absence of occult white matter injury in Lyme disease in juxtaposition to multiple sclerosis.

Other Infectious Dementias. Of importance but rarely rapid in progression is tertiary syphilis, which should be clinically determined by using appropriate laboratory testing with rapid plasma reagin testing performed in all suspected patients with rapid or presenile dementia. Cognitive decline is the most common neurologic symptom in tertiary syphilis. Early MR imaging reports describe focal hyperintensities on T2-weighted images in an arterial distribution, in keeping with meningovascular pathophysiology. Multiple areas of ischemia and infarction are thought to result from endarteritis in some cases of fulminant neurosyphilis. Cerebral gummatous disease may also occur in neurosyphilis, with lesions that are hypointense on T1- and hyperintense on T2-weighted images.

Other infectious etiologies such as fungal encephalitides may present with dementia, but as with most infections of the central nervous system, they also tend to present with neurologic dysfunction and other clinical features more often than simple dementia. There are reports of Cryptococcus neoformans as a causative agent of rapidly progressive cognitive dysfunction, with 1 case even misdiagnosed as Alzheimer disease. Neurocysticercosis is rare in the developed world, but in places where it is endemic, it may present with dementia in a small fraction of cases.

Another infectious form of sudden dementia, typically seen in a much younger population of children and adolescents (in comparison with the remainder of conditions discussed in this article) is subacute sclerosing panencephalitis, which is a rare condition caused by the measles paramyxovirus, occurring months to years after infection. MR imaging findings are generally normal early in SSPE and may progress to involve high T2 signal within white matter or hyperintense lesions in the parietal and temporal lobes in the acute stage and then progress to involve atrophy and deep
and cortical gray matter changes.\textsuperscript{31-33} MR spectroscopy in later stages of SSPE reveals decreased NAA, consistent with neuronal loss and elevated choline, lipid, and lactate concentrations consistent with inflammation.\textsuperscript{34} More recent work indicates the use of DTI; notably, DTI detects reductions in fractional anisotropy even within normal-appearing white matter on conventional imaging, suggesting subtle disruption of white matter tracts in the early stages of SSPE.\textsuperscript{35}

**Inflammatory and Autoimmune Conditions**

Inflammatory and autoimmune conditions are often associated with diverse clinical symptoms and an amorphous presentation. They constitute the second largest group of rapidly progressive dementias in 2 large cohorts but are a much smaller portion of another group of presenile rapid dementias.\textsuperscript{3,36}

**Multiple Sclerosis.** Nearly 1 in 20 patients with MS manifest dementia, but the prevalence of other cognitive disturbances is much more common and subtle.\textsuperscript{37} Most imaging reports of this phenomenon attribute the cognitive decline to the presence of nonspecific neuroanatomic atrophy, without particular observation of an affected structure.\textsuperscript{38} The reason for this generalized atrophy has been discussed and examined at great length—even in normal-appearing white matter in MS, there are detectable correlates of neuronal loss with decreased NAA on MR spectroscopy within both normal white and gray matter in patients with MS.\textsuperscript{39} Moreover, these measures of atrophy in patients with MS correlate with the progression of disability and were reviewed at great length by Anderson et al.\textsuperscript{39} Nevertheless, dementia in the setting of patients with known MS should be evaluated as with any other rapidly progressive dementia, with MS-related dementia as a diagnosis of exclusion.

**Hashimoto Encephalopathy.** This condition related to Hashimoto thyroiditis is also referred to as steroid-responsive encephalopathy with autoimmune thyroiditis, which explains the nature of this syndrome that is believed to be related to a cross-reaction between autoimmune antithyroid antibodies and neurons, resulting in generalized CNS inflammation and neurologic impairment.\textsuperscript{3,40} Most interesting, patients may not demonstrate overt thyroid dysfunction at the time of neurologic presentation. MR imaging findings, as with many of these autoimmune conditions, are frequently normal (60% of cases) or nonspecific in nature, involving FLAIR and T2 abnormalities within subcortical white matter and contrast enhancement of the meninges.\textsuperscript{40,41} Making the diagnosis of SREAT is important because early initiation of corticosteroid pharmacotherapy yields symptom resolution and reversal of imaging findings.\textsuperscript{41}

**Neurosarcoidosis.** As a systemic granulomatous disease, sarcoidosis may simulate many neurologic disorders, and likewise, imaging of neurosarcoidosis is complex.\textsuperscript{42} Neurologic involvement is also relatively common in neurosarcoidosis, with approximately 1 in 4 patients with sarcoidosis demonstrating CNS involvement at postmortem examination.\textsuperscript{43} When present, the most frequent abnormalities observed on MR imaging are periventricular T2-weighted hyperintense lesions, enhancing granulomas, and leptomeningeal enhancement similar to that seen in meningitis.\textsuperscript{42,44} Dural masses may be seen, which are T2 hypointense; contrast enhancement may be mistaken for meningioma, without a clinical history of sarcoidosis.\textsuperscript{43}

**Vasculitides.** Primary CNS vasculitis is a rare entity, but systemic vasculitides may also present with CNS manifestations with cognitive deficits in half of patients.\textsuperscript{45,46} Vasculitides can be differentiated from many other dementias in that there are frequently concomitant cerebrovascular events seen on MR imaging, with hemorrhage and infarct more likely present.\textsuperscript{3} FLAIR and T2-weighted hyperintensities are also prevalent in vasculitis, thickening of the vessels and extramural enhancement are more definitive findings, and findings of gadolinium enhancement within the meninges are not as commonplace as generally thought on the basis of the results of a large observational study.\textsuperscript{45}

**Other Inflammatory and Autoimmune Conditions.** Celiac disease, an intolerance of food products containing gluten, often with the presence of autoantibodies to the enzyme tissue transglutaminase, may present with neurologic symptoms, including dementia, though those reported are quite varied, ranging from chronic headaches to ataxia.\textsuperscript{47,48} Oddly, these symptoms can occur without gastrointestinal involvement and may improve with dietary changes.\textsuperscript{49} MR imaging of the brain may reveal hyperintense T2-weighted posterior white matter foci with a reported prevalence of 1 in 5 in patients with celiac disease in 1 study, independent of the presence of neurologic symptoms.\textsuperscript{3,50} Most interesting, measured cerebral perfusion on SPECT imaging in 1 small study was significantly lower within several brain regions in patients with untreated celiac disease, while there was no significant difference between patients adhering to dietary restriction and controls.\textsuperscript{51} These findings suggest a process by which patients may sustain hypoperfusion with resultant brain injury, generating lesions visible on conventional MR imaging. Nevertheless, the evidence for and mechanism of neurologic involvement and the importance of MR imaging findings in celiac disease are controversial topics without conclusions.

**Malignancy-Related Conditions**

**Paraneoplastic CNS Involvement.** There are a number of paraneoplastic neurologic disorders with a diverse set of clinical presentations that may present with cognitive impairment or dementia. The most common symptoms that may suggest this condition in the differential diagnosis of rapidly progressive dementia involve retrograde and anterograde amnesia. The diagnosis of this condition is relatively straightforward in patients with established cancer (particularly, small cell lung cancer, germ cell tumors, thymoma) but must also be considered in the setting of occult malignancy as well. Paraneoplastic limbic encephalitis (Fig 3)\textsuperscript{52} is the most classic paraneoplastic disorder with anterograde memory impairment being most prominent; imaging reveals increased signal within the mesial temporal lobes.\textsuperscript{41} FLAIR hyperintensities within the caudate, anterior putamen, and sometimes the medial temporal lobe have been reported in the setting of paraneoplastic syndrome in small cell lung cancer with anti-CV2 antibodies.\textsuperscript{53} Half of those patients with autoimmune dementia diagnosed with CSF autoantibodies will have a paraneoplastic syndrome. Other authors recommend the use of FDG-PET for any patient sus-
FIG 3. Acute limbic encephalitis. A 26-year-old man with underlying precursor T-cell acute lymphoblastic leukemia developed generalized seizures, had short-term memory loss, and was disoriented in time and place 1 month after undergoing unsuccessful allogeneic bone marrow transplantation. At autopsy, multifocal subacute polioencephalomyelitis in the brain regions that were shown as affected on MR images confirmed the diagnosis of paraneoplastic limbic encephalitis with neuronal loss. Initial MR images were obtained 1 day after the first generalized seizure occurred. A, Axial FLAIR image (TR/TE, 9000/110 ms; TI, 2261 ms) shows a slightly elevated signal intensity of both hippocampal formations (black arrows) and amygdala (white arrows). B, Coronal conventional T2-weighted turbo spin-echo image (TR/TE/NEX, 4462 ms/120 ms/3) shows no signal-intensity abnormality.52

The clinical diagnosis of CNS lymphoma is frequently problematic, with nondiagnostic studies leading ultimately to the necessity of brain biopsy for definitive diagnosis; imaging may provide ancillary support to clinical evidence.5 MR spectroscopy may be particularly helpful in identifying a neoplastic pattern of metabolite concentrations evidenced by large choline concentrations with lipid and lactate present.14 MR perfusion may clarify diagnosis in contrast to other tumors in that primary lymphoma in the brain has lower relative cerebral blood volume.56 Likewise, DWI with ADC maps demonstrate low ADC values compared with glioblastoma and metastatic lesions.58 Brain thallium-201 SPECT may be combined with toxoplasma serology to improve diagnostic accuracy in excluding this potential infectious etiology with focal CNS mass lesions.18

Intravascular lymphoma, a rare non-Hodgkin lymphoma, may likewise generate neurologic symptoms, including subacute dementia, concomitantly with cerebrovascular events.2 As a result of its intravascular source, CNS involvement is characterized by diffuse T2-weighted hyperintensity with patchy enhancement and, on occasion, edema.2,59 There are some diffusion changes consistent with ischemia in the setting of intravascular lymphoma.60 Angiography in these patients does not aid in the diagnosis because intravascular lymphoma appears consistent with vasculitis as a result of the intravascular malignancy.2 An antemortem diagnosis of intravascular lymphoma is particularly difficult.

Lymphoma. Primary CNS lymphoma is a malignancy of increased recognition and prevalence related to its presence in AIDS. Most of its neurologic manifestations are more related to mass-effect symptoms, but dementia has also been observed, thereby adding this neoplasm to the differential diagnosis of rapid-onset dementias.55 Early diagnosis is important for averting the need for extensive resection with poorer outcomes.56 MR imaging categorically reveals solitary or multiple supratentorial mass lesions with minimal edema, generally isointense-to-mildly hyperintense on T1-weighted imaging and hypointense on T2-weighted images, and almost always with enhancement.14,57 Solitary lesions are more common in immunocompetent individuals.14 Of importance, the lesion is capable of crossing the corpus callosum, forming a “butterfly” lesion.57 There may also be diffuse extension of the white matter signal change classified as “lymphomatosis cerebri.” This form of diffusely infiltrating CNS lymphoma would be more likely to present as a progressive dementia.3

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Metastatic CNS involvement. As with any intracranial process and as the most common cause of brain tumors, metastatic disease may present with rapidly progressive dementia, particularly in the setting of multifocal cortical involvement as has been reported in a few cases of rare miliary cerebral metastases presenting with subacute dementia.61-63 Some patients have been reported to develop dementia during treatment as a result of metastatic disease, which could be mistaken for a side effect of therapy (particularly radiation therapy).64 In each of these cases, diagnosis of the underlying cause of the dementia is often elusive until postmortem neuropathology. The imaging of most brain lesions consistent with metastatic cancer, aside from miliary spread, is more straightforward, and recognition is important so that patients may benefit from radiation therapy.65 Imaging of brain metastases requires a dedicated review of its own.65 Briefly, MR imaging generally reveals multiple lesions with a propensity for the gray-white matter junctions with T2 hyperintensity consistent with extensive edema.67 Some of these lesions may hemorrhage, as may be evidenced by T1 hyperintensity. DWI and DTI aid in distinguishing metastases from other lesions such as abscesses in the setting of cystic metastasis with a relative lack of restricted diffusion (bright on ADC, dark on DWI); DTI demonstrates displacement of white matter fibers in metastasis.68

Lymphomatoid Granulomatosis. Lymphomatoid granulomatosis, a B-cell lymphoma caused by the Epstein-Barr virus with prominent CNS involvement as a result of angiodestructive pathophysiology, is one condition that has been reported to pres-
ent with dementia in a presenile population. Generally, pulmonary manifestations exist in almost all patients with CNS manifestations of LG; this unusual constellation may provide the clinical clue necessary to make this diagnosis, whereas those cases with exclusively neurologic involvement are particularly difficult to diagnose. Multifocal involvement on brain MR imaging is commonplace, with punctate or linear T2 hyperintensities within perivascular spaces with corresponding enhancement on T1 post-contrast images, particularly within frontal and parietal regions—a finding not exclusive to LG and seen in angiitis and neurosarcoidosis but distinguishable from CNS lymphoma, which most often presents as a solitary lesion. These regions of enhancement correspond on histopathology to granulomatous infiltrates of lymphoid cells and precede hemorrhage. Ring-like enhancement may also appear in the lesions of LG. Imaging may demonstrate temporal lobe atrophy, perhaps related to lymphocytic infiltration into cerebral blood vessels causing microvascular occlusion and resultant ischemia. Most interesting, researc

Clinically, the memory loss and cortical symptoms seen in Wernicke encephalopathy are accompanied by ophthalmoplegias and ataxia; however, as already mentioned, several conditions within the spectrum of rapidly progressive dementia share these symptoms. Moreover, acute Wernicke encephalopathy is underappreciated, particularly in nonalcoholic patients. Imaging clues may improve recognition of this underdiagnosed clinical entity, with excellent specificity of findings on MR imaging. There may be T2-weighted and FLAIR hyperintensity within the area around the third ventricle, periaqueductal regions, and bilateral thalami (particularly dorsomedial) in Wernicke encephalopathy—a finding that has been reported even in nonalcoholic cases. Moreover, serial imaging illustrates remission of these changes with successful treatment. DWI improves the neuroimaging evaluation and appears more sensitive to the presumed ischemic and cytotoxic pathophysiology of Wernicke encephalopathy in these regions, evidenced by restricted diffusion. Therefore, clinical imaging demonstrating gadolinium enhancement or DWI abnormalities within the mammillary bodies and periaqueductal regions and thalami can be of clinical utility in clarifying the diagnosis. These imaging findings may conveniently correlate with the presence of hemorrhagic necrosis within these structures. Ancillary imaging with MR spectroscopy may show decreased or normal NAA with the notable presence of lactate.

**Wilson Disease.** Impaired copper transportation in this autosomal recessive disorder may precipitate either liver disease or neurologic dysfunction, which in the absence of liver disease, presents later in early adulthood. For this reason, progressive dementia with or without other neurologic findings may occur in an atypical age group with Wilson disease. Classically seen on MR imaging is the accumulation of copper within the putamen and thalami reflected by T2-weighted hyperintensity without evidence of restricted diffusion. A few reports have described the presence of high signal intensity in T2-weighted images of the tegmentum and red nuclei to form the “face of the giant panda” and, inferiorly within the pons, to form the “face of the panda cub,” with relative hypointensity of the central portion with adjacent hyperintensity within the aqueduct. While these imaging findings may suggest impaired copper transportation as a cause, ultimately laboratory studies of copper and ceruloplasmin facilitate a definitive diagnosis.

**Other Conditions.** Metabolic diseases such as mitochondrial disease may also present with a characteristically subcortical dementia; there is impairment in attention and rousability early in the disease process that differentiates them from dementias. Prominent potential causes of subacute dementia are mitochondrial myopathy, encephalopathy, lactic acidosis, and stroke-like episodes (MELAS). Imaging findings early in this disease may reveal cortical hyperintensities on FLAIR and T2-weighted images in a gyriform pattern. DWI also depicts these lesions characteristic of MELAS as hyperintense regions of the cortex.

Osmotic myelinolysis is another potential cause of rapidly progressive dementia but will generally present more acutely and involve decreased levels of consciousness that exclude it from the rapidly progressive dementias. Central pontine myelinolysis is visible as a trident-shaped increased T2 signal intensity with sparing of the corticospinal tracts. Extra-pontine myelinolysis, while rarer, may have T2 prolongation and DWI signal increases in the basal ganglia, thalamus, or cerebellum. The involvement of subcortical structures is an uncommon occurrence, but nevertheless 1 has been reported.

**CONCLUSIONS**

Presenile and rapidly progressive dementias are a diverse collection of conditions ranging from reversible to intransigent and
rapidly progressive. Distinctions within this heterogeneous group are best accomplished with a careful and thorough clinical and laboratory examination and may be greatly aided by the use of neuroimaging, particularly MR imaging. We discussed, in the first part of this review, the more frequently encountered neurodegenerative causes of dementia, and in this second installment, we conclude our discussion of rapidly progressive dementias with a review of prion, infectious, inflammatory, autoimmune, neoplastic, and other conditions. The need for further investigation in several areas of clinical investigation of all these conditions is particularly important, especially because a large number of these conditions have overlapping imaging and clinical findings that obfuscate accurate diagnosis. Both the clinician and radiologist alike should be open to the possibility of alternate diagnoses in the setting of rapidly progressive presenile dementias, particularly because several diseases within this group benefit from urgent and specific treatment.

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