A Closer Investigation of the Synchronous Bilateral Pattern of MRI Lesions in Acute Necrotizing Encephalopathy Type 1

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ABSTRACT

SUMMARY: We observed a lesion pattern in a series of 4 cases of RANBP2-mutation–linked acute necrotizing encephalopathy, which appears to be specific for this condition. The setting of synchronous bilateral mammillary, amygdaloid, and lateral geniculate lesions, along with claustro-parahippocampal lesions, can serve as a diagnostic tool in this condition. We add several further details to the MR imaging features of the typical brain lesions encountered in this disease.

ABBREVIATIONS: ANE1 = acute necrotizing encephalopathy type 1; RANBP2 = RAN binding protein 2

As a subgroup of the more heterogeneous entity named acute necrotizing encephalopathy of childhood, acute necrotizing encephalopathy type 1 (ANE1) is an underrecognized, infection-induced condition with autosomal dominant inheritance of several pathogenic mutations of the gene RANBP2, showing incomplete penetrance. This condition is also named autosomal dominant acute necrotizing encephalopathy or, less commonly, infection-induced/triggered acute encephalopathy type 3. The gene encodes RAN binding protein 2 (RANBP2), a nuclear pore protein with many intracellular functions. Its dysfunction causes alterations in intracellular metabolism, mitochondrial distribution, nucleocytoplasmatic trafficking, and cytokine overproduction. On the basis of current scientific data, it is not clear how the pathophysiological alterations predispose to attacks or why certain brain areas are specifically targeted.

MR imaging manifestations of encephalopathy and myelopathy are described in the literature.

MATERIALS AND METHODS

We registered 4 pediatric cases of RANBP2-mutation–linked acute necrotizing encephalopathy. Patient 1 is male and was 10 months old at the time of the first of his 2 attacks. Patient 2 is male and was 10 months old at the first of 3 attacks. Patient 4 is female, and was 4 years old at the first attack, and had 2 attacks. Patients 1 and 2 are related (half-siblings). All our patients were treated in the intensive care unit during all their attacks. Clinical follow-up was 6–18 years: Patient 1 has mild cerebral palsy, well-controlled epilepsy, and only minor school problems. Patient 2 has no neurocognitive sequelae and maintains excellent grades at school. Patients 3 and 4 have minor motor coordination issues and moderate mental retardation based on neurocognitive testing.

We used 3T Achieva and 3T Ingenia (Philips Healthcare) scanners for our work. We performed our standard head imaging protocol during the attacks (axial DWI/ADC, T2, FLAIR, SWI, 3D TOF-MRA, non-enhanced, and contrast-enhanced 3D T1 sequences), and an additional 3D FLAIR sequence for assessing small necrotic lesions in the chronic phase.

RESULTS

In the acute phase, restricted diffusion according to DWI and ADC map images, increased signal intensity on T2, FLAIR, SWI, 3D TOF-MRA, non-enhanced, and contrast-enhanced 3D T1 sequences, and an additional 3D FLAIR sequence for assessing small necrotic lesions in the chronic phase.
Many authors refer to bilateral temporomedial involvement, but our current investigation shows that it is more precisely located at the most anterior, subcortical aspect of the parahippocampal gyri (Fig 1). These lesions were confluent with those in the claustrum (hence the above term “claustro-parahippocampal”) during the acute phase but became isolated lacunar necrotic lesions in the chronic stage.

The lesions were exceptional in targeting a specific area of white matter, while being similar in appearance to the other bilateral lesions targeting gray matter.

We found that the thalamic lesions were localized in the ventromedial and/or dorsomedial and/or dorsolateral (also named ventral posterior) nuclei. This was clearly demonstrated in the chronic, postnecrotic stage (Fig 2). In the acute phase, the perifocal edema may be more widespread (involving the whole thalamus or even the surrounding regions like the internal capsule).

We found that the core of the pontine lesions in 3 of our patients arose at the lateral and, in 1 patient (during a different attack), at the central parts of the pontine base (Fig 2). One patient did not present pontine lesions. When the bilateral central parts were involved, they had a heart-shaped appearance on transverse sections (Fig 2).

Finally, we observed that typical bulbar lesions were located symmetrically in the regions of the vestibular nuclei.

**DISCUSSION**

To our best knowledge, we are the first to describe bilateral symmetric lesions of the lateral geniculate bodies in acute necrotizing encephalopathy type 1. Bilateral lesions of these structures are demonstrated by MR imaging exclusively in the setting of another condition in literature. In this case of pancreatitis and microangiopathy, the rest of the associated lesions typically found in ANE1 were absent.

The lateral geniculate body is an anatomic part of the thalamus (located inferolateral to the thalamic pulvinar). Therefore, we emphasize that the typical thalamic lesions considered to be most common in the radiologic setting of this disease are located in other parts of the thalami, as described in the Results section.

We clarified that the symmetric lesions previously described in the temporal medial regions are actually situated at the most anterior subcortical aspect of the parahippocampal gyri and are unique in involving the white matter. Attention must be paid to differentiate them from the amygdaloid lesions because the proximity of the 2 small structures can lead to misinterpretation, appearing confusingly in figure captions of otherwise excellent articles.

The “heart appearance” of the pontine lesions on MR imaging was previously described in a case of bilateral medial pontine...
We present ours as a variation of this radiologic sign in type 1 acute necrotizing encephalopathy.

CONCLUSIONS

On the basis of the analysis of the MR imaging studies of 4 patients, we describe several observations that can improve the diagnostic accuracy in ANE1 and may help the understanding of the pathomechanisms of this disease. The pattern of a synchronous appearance of bilateral amygdaloid, mammillary, lateral geniculate, and claustrum-parahippocampal lesions appears to be constant and is potentially highly specific for this disease. This pattern, in combination with the features of thalamic and brainstem lesions described in detail, might play a role in further development of the neuroradiologic diagnostics or prognostic scoring. To estimate the specificity and sensitivity rates, further investigation on a broader patient group is needed.

Disclosure forms provided by the authors are available with the full text and PDF of this article at www.ajnr.org.

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