“CHOICES”: An acronym to aid in delineating potential causes of non-metabolic, non-infectious acute toxic leukoencephalopathy

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A B S T R A C T

Purpose: To describe non-metabolic, non-infectious etiologies of acute toxic leukoencephalopathy (ATL) on DWI MRI, and provide a useful acronym to remember them.

Material and Methods: Our PACS archive was reviewed, yielding 185 patients with suspected ATL per MRI reports and clinical followup; infectious or metabolic causes were excluded.

Result/Discussion: The 87 included non-infectious, non-metabolic ATL patients’ etiologies are represented by the acronym 'CHOICES': chemotherapy ('C', n = 34); heroin-induced ('H', n = 6); opioid analogues ('O', n = 14); immunosuppressant ('I', n = 11) or imidazole (n = 2); cocaine ('C', n = 1); environmental or ethanol abuse ('E', n = 5); splenial lesions ('S', n = 9); and ‘other’ (n = 5).

Conclusion: The ‘CHOICES’ acronym delineates various toxic etiologies of ATL.

1. Introduction

The term “leukoencephalopathy” (LE) indicates an abnormality of cerebral white matter (WM) that predominantly damages myelin. Toxic LE predominantly affects the cerebral WM and may occur from an exposure to a variety of agents, including environmental toxins, prescription medications, metabolic substances, and illicit drug usage [1]. Such leukotoxic agents can incite varying degrees of deterioration of higher cerebral function, which can clinically present acutely or chronically along a spectrum ranging from mild confusion, altered mental status, dementia, coma, or even death [1]. Acute toxic leukoencephalopathy (ATL) is a potentially reversible condition that may improve after treatment or following withdrawal of the offending toxin early in the course of disease; on MRI, ATL variably has abnormal signal on FLAIR/T2WI, but the abnormalities are typically visible as bright on DWI and dark on accompanying ADC maps relative to normal-appearing white matter (NAWM) [2,3]. ATL typically affects WM tracts extending from the periventricular white matter (PVWM) out to the subcortical WM, more commonly in a symmetric distribution. In a minority of cases, atypical areas of involvement include the basal ganglia, thalami, brainstem, internal capsules, and cerebellum [2,3]. While the exact pathophysiologic mechanism of ATL is unknown, preliminary histologic evidence suggests that the various forms of endothelial insults subsequently and generally result in intramyelinic edema [2,3]. Regarding potential contributing conditions to the onset of ATL, it is of note that uremia has recently been identified as a potential exacerbating factor in ATL [3].

The most common causes of ATL in adults include chemotherapy agents, immunosuppressant therapy, illicit drug use, and medication overuse, particularly from opioid overuse [2,3]. Environmental toxins such as carbon monoxide (CO) may also cause ATL [1–8]. As reversible splenial lesions (RSL’s) may arise from similar potentially toxic substances and the callosal splenium is in fact periventricular, RSL may be considered a subtype of ATL [2,3].

In this study and accompanying review, a differential diagnosis of potential extrinsic causes of ATL (i.e. non-metabolic and non-infectious) in adults is proposed via the acronym “CHOICES”: ‘C’ = chemotherapy; ‘H’ = heroin-induced (illicit usage); ‘O’ = opioid analogues (medication...
Table 1: Potential etiologies of ATL

<table>
<thead>
<tr>
<th>C</th>
<th>Chemotherapy (n = 34)</th>
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<tr>
<td>H</td>
<td>Heroin-induced (n = 6), whether via intravenous (n = 3) or inhaled (n = 3) routes</td>
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<tr>
<td>O</td>
<td>Opioid analogue (n = 14), abuse via various routes of “non-heroine” medications</td>
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<tr>
<td>I</td>
<td>Immunosuppressant (n = 11) or Imidazole medications (n = 2)</td>
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<tr>
<td>C</td>
<td>“Crack” cocaine abuse (n = 1)</td>
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<tr>
<td>E</td>
<td>Environmental (CO, n = 3) and Ethanol-related (n = 2)</td>
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<tr>
<td>S</td>
<td>Splenial lesions (RSL), total n = 9: which include AEDs (n = 7), chemotherapy (n = 1), immunosuppressant medication (n = 1)</td>
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Note: CO = Carbon monoxide, RSL = reversible splenial lesion, AEDs = anti-epileptic drugs.

Table 1: Potential etiologies of ATL

As above, the RIS search yielded 185 patients suspected of having ATL, based on MRI. However, 98 were ultimately excluded due to: 1) 73/98 as they either were confirmed not to have ATL (n = 53), or had inadequate imaging such as a lack of ADC maps (n = 14), or were children (n = 4), or were found to be a chronic form of toxic LE (n = 2); 2) the other 25/98 exclusions were due to being either metabolic/intrinsic or infectious causes, including acute hepatic encephalopathy (AHE, n = 14), sepsis-related ATL (n = 5), while the other 14/20 were attributed to morphine (oral, n = 3), inhaled (n = 3), while the other 14/20 were attributed to morphine (oral, n = 3), inhaled (n = 3), acetaminophen (n = 2), dopaminergic (n = 1), oxycodone + cocaine (n = 1), and an unspecified opioid (diagnosed via urine test, n = 9). Thus, in 9/20 patients the specific opioid agent was not discerned, demonstrating how causative opioid etiologies may not be determined.

Among the 34 chemotherapeutic patients, being the largest ATL subcategory, the causative agents were: fludarabine (n = 16), methotrexate (n = 9), combined methotrexate + cytarabine (n = 3), 5-fluourouracil (n = 2), doxorubicin (n = 1), cisplatin (n = 1), erlotinib (n = 1), and cytarabine (n = 1).

Regarding the 20 heroin/opioid-related ATL patients, 6/20 were heroin-induced (inhaled n = 3, intravenous n = 3), while the other 14/20 were attributed to morphine (oral, n = 2), methadone (n = 1), transdermal fentanyl (n = 1), combined oxycodone + cocaine (n = 1), and an unspecified opioid (diagnosed via urine test, n = 9). Thus, in 9/20 patients the specific opioid agent was not discerned, demonstrating how causative opioid etiologies may not be determined.

Among the 11 immunosuppressant-related ATL patients, the causes were: cyclosporine (n = 4), tacrolimus (n = 4), and cyclophosphamide (n = 1); the other 2/11 were an “unknown immunosuppressant”, where the records were too old to obtain (i.e. not electronic).

The etiologies of confirmed RSL-type lesions involving solely the callosal splenium were further sub-categorized by their cause of ATL, which included: anti-epileptic drugs (AEDs) (n = 7), chemotherapy (n = 1), and an immunosuppressant medication (n = 1), as in Table 1.
common effect of certain toxins relative to another entity, posterior reversible encephalopathy syndrome (PRES), which may occur from particular toxins that also cause ATL, most notably chemotherapy, immunosuppressants, and illicit drugs [10]. PRES is a more common acute and potentially reversible encephalopathic toxic and non-metabolic insult, also having an overlap in clinical symptom presentation (encephalopathic) as well as etiologies, but these two entities can be distinguished by their imaging appearances. For example, PRES has cortical and subcortical edema on routine MR imaging sequences such as FLAIR; in comparison, the extent of ATL is best visualized with reduced diffusion within the PVWM on DWI [2,3,9]. Also, the anatomic distribution and appearance on DWI help to differentiate these two potentially reversible entities, as ATL “starts” (i.e. in the mildest forms) within the PVWM (corona radiata and centrum semiovale), while the mildest cases of PRES are typically in the cortex/subcortical WM, most commonly being parietal and occipital in distribution [10–12]. Also, based on DWI, ATL has deep PVWM reduced diffusion, typically in a confluent and symmetric fashion, while PRES has DWI abnormalities in only a minority; when present in PRES, these DWI-positive foci are typically focal, asymmetric, cortical-based, and sometimes gyriform [10–13]. Also, postcontrast imaging findings differ between the two entities, as ATL only rarely has contrast enhancement on T1WI MRI (described as punctate within the PVWM in < 10%), while PRES can have varying degrees of avid enhancement in up to 50%, whether cortical, gyriform, or leptomeningeal [3,11]. Finally, on SWI, ATL has microhemorrhages in 15%, but rarely has macrohemorrhages (> 1 cm); meanwhile PRES has microhemorrhages (< 1 cm in size) in nearly 60%, with macrohemorrhages in up to 10–20% [3,14]. Of note, several patients with combined ATL and PRES have recently been reported, corroborating the concept that these entities may have a certain degree of overlap in both etiology and pathophysiology, thought related to toxin-induced endothelial injury with subsequent loss of blood-brain barrier integrity [13,15]. Hence, the following discussion focuses on letters for the acronym “CHOICES” that represents toxic, non-metabolic, non-infectious causes of ATL, keeping in mind that many (but not all) of these agents may also cause PRES, particularly chemotherapeutic medications, immunosuppressant agents, and illicit drug usage.

4.1. ‘C’ = chemotherapy (intravenous or intrathecal)

ATL is a clinical consideration in cancer patients exposed to antineoplastic agents who develop acute neurologic symptoms [4]. Chemotherapy is likely the most common cause of ATL (39% ATL patients in this study), based on this and prior studies, with fludarabine potentially being the most common agent in this study (47% of chemotherapy-related ATL), followed by methotrexate (26.5%) [3,4]. Notably, fludarabine-related ATL has relatively poorer outcomes as compared to other chemotherapeutic agents, as it can cause severe, late-onset symptoms even months after the exposure; hence, an exposure to fludarabine is important to recognize as it is likely the most common cause of chemotherapy-related death from ATL [2,16] (Fig. 1).

With methotrexate, the second most common cause of chemotherapy-induced ATL in the current study, the neurotoxicity arises from disruption of CNS folate homeostasis or via direct toxic neuronal damage, typically involving the PVWM with focal or multifocal reduced diffusivity; notably, intrathecal methotrexate neurotoxicity is relatively well-described [2,6,10,17]. The cerebral and cerebellar cortices, subcortical WM, and thalami are other atypical sites of involvement by methotrexate, which, incidentally, can also cause PRES [2,6,10]. Various other agents, such as 5-fluorouracil and anti-metabolite agents such as cytarabine, seem to be more reversible and often have better outcomes if promptly discontinued. Of note, these agents can potentiate each other to cause ATL, as noted in this study’s 3 patients with combined methotrexate + cytarabine toxicity [2]. Also of interest is 5-fluorouracil neurotoxicity, where there is often reduced diffusion in the centrum semiovale, callosal splenium, and PVWM that usually reverses clinically and on imaging after removal of the offending agent; hence,
5-FU-related ATL seems less likely to result in permanent sequelae if recognized promptly (Fig. 2) [6,18].

4.2. 'H' = heroin-induced ATL

Myelin is vulnerable to lipophilic substances and lipid peroxidation, due to its high lipid content [19]. Thus, as opioids are lipophilic, they can traverse the blood-brain barrier and damage myelin. Notably, the word “opioid” is a broad term encompassing both natural (i.e. true opiates such as heroin, morphine and codeine), semi-synthetic (i.e. hydrocodone, oxycodone), and synthetic substances (i.e. methadone, fentanyl) [20]. Hence, for the purposes of the ‘CHOICES’ acronym, we separate heroin-induced LE (usually intravenous or inhaled) from the opioid analogues (typically oral or intravenous), as the original descriptions of toxic LE from illicit opiates are largely based on heroin usage, as opposed to medication abuse [21–23]. A confounding issue in identifying the etiology of ATL can be when heroin or other opiates occur in tandem with other illicit drugs (such as cocaine), making it difficult to discern which is the causative etiology; such instances with a positive urine sample having an ‘unspecified opioid’ occurred as the cause in 9/87 patients (10.3%) in this study [2,3].

Heroin is one of the most commonly abused opiates, seen in about 5% of ATL patients, in accord with the finding of occurring in 6/87 patients (6.9%) in this study [3]. Clinically, whether intravenous or inhaled, heroin-induced ATL may progress through stages, beginning with a cerebellar syndrome, while the other end of the spectrum is a final clinical stage of symptomatology that includes spasms, akinetic mutism, and possibly death [20]. Characteristic areas of inhaled heroin-induced ATL (so-called “chasing the dragon”) include the posterior cerebral WM, the internal capsule’s posterior limbs, and occasionally cerebellar WM, whereas the subcortical U fibers and the internal capsules’ genu are more frequently involved in intravenous heroin-induced neurotoxicity [2,3,5,22]. Depending on the severity, lesions progress symmetrically from the deep PVWM to the subcortical WM, often spreading posterior to anterior; this can be entirely reversible, particularly in intravenous exposures [2,24]. Inhaled exposures (‘chasing the dragon’) tend to have poorer outcomes relative to intravenous exposures, and such severe cases may be result in diffuse cerebral atrophy.
A 19 year old male with oral methadone overdose-related ATL presented with acute left sided weakness and upper motor neuron signs. 4A-C: The reduced diffusion bilaterally involves the PVWM (arrows) on DWI MRI (A) and ADC map (B), with bilateral symmetric periventricular hyperintensity on FLAIR (C). 4D-F: There is also reduced diffusion within the optic radiations on DWI MRI (D) and ADC map (E), also having abnormal signal on FLAIR (F).

A 37 year old male with myelodysplastic syndrome underwent a normal-appearing baseline brain MRI prior to bone marrow transplantation (BMT), which was nearly normal on FLAIR (A), DWI (B) and ADC map (C). Nineteen days post-BMT, while the patient was on cyclosporine therapy (an immunosuppressant medication), regions of hypointensity (arrows) from ATL developed within the bilateral posterior PVWM on FLAIR (D) and DWI (E); note mild, asymmetric reduced diffusion in the left posterior PVWM on the ADC map (F, arrow). The patient’s symptoms promptly resolved over several days after the cessation of cyclosporine.
and neurologic sequelae [2,3,5] (Fig. 3). Recent research has suggested that, of the major subtypes of etiologies of ATL involving the PVWM, opiates typically have the worst outcomes, followed by chemotherapy [2,3].

4.3. ‘O’ = opioid analogues (i.e. oxycodone, hydrocodone, methadone, fentanyl)

There are a number of opioid analogues which have been more recently described to cause ATL in contrast to that traditionally described of heroin-induced ATL [2,3]. For example, methadone hydrochloride is a synthetic opioid receptor agonist, where its primary symptoms of neurotoxicity are sedation, dizziness, abnormal sensation, respiratory depression, and coma [25]. On neuroimaging, a number of cases of methadone-induced LE have been described, having bilaterally high signal within the cerebral WM on FLAIR or T2WI, with sparing of the cortical U-fibers; in addition, symmetrical involvement of the cerebellar gray and WM, basal ganglia, hippocampi, central segmental tracts, have also been reported [25–27].

Oxycodone- and fentanyl-induced ATL are less well described. As mentioned previously, in opioid-related ATL it can be difficult to discern the exact causative drug or medication since illicit usage may involve several drugs ingested together. In the current study, there were 9 patients categorized as an ‘unspecified opioid’, for which the exact causative opioid was unclear. Oxycodone is an orally ingested semisynthetic opioid which has been reported to involve the PVWM, but also may involve the cerebellum and globi pallidi [3,28]. Fentanyl is a synthetic analogue, typically administered intravenously, but also can be transdermal, which can present on DWI and FLAIR MRI as PVWM and deep WM abnormalities in cerebral or cerebellar distributions [2,3]. It is important to note that in the setting of an opioid overdose such as fentanyl, it may be difficult to initially differentiate the MRI findings of ATL involving the PVWM from that of fentanyl-related (or other opioid-induced) delayed or subacute post-anoxic/hypoxic-ischemic encephalopathy (HIE), which can occur from respiratory depression or from the opioid’s toxic effects, or even via a combination of both ATL and HIE occurring simultaneously [2,29]. In pure ATL, the cerebral cortex overlying the PVWM injury remains normal signal intensity on FLAIR, while in patients with purely HIE the overlying cortex is mildly hyperintense [2] (Fig. 4).

4.4. ‘I’ = immunosuppressant medications (e.g. cyclosporine, tacrolimus, etc.) and imidazoles

Immunosuppressant-related ATL is a less common subtype, typically due to medications such as cyclosporine, tacrolimus, and less commonly mycophenolate. In the current study, this occurred in 12.6% of patients, where cyclosporine and tacrolimus were the most common
causes. As with chemotherapeutic agents, immunosuppressant-related neurotoxicity more commonly results in PRES, but a minority (about 20–28%) of patients with MRI findings of neurotoxicity may suffer purely ATL [2–4,30]. The neurotoxicity is thought to result from endothelial injury that leads to blood–brain barrier dysfunction and subsequent PVWM injury, most often in the parietooccipital regions, usually with symmetric involvement of the deep and subcortical WM (Fig. 5) [2,3,31]. On DWI, there is often a milder degree of reduced diffusion relative to other causes of ATL, and the clinical severity and outcomes are also typically milder and more reversible in immunosuppressant-related ATL, as compared to chemotherapy and opiate-related ATL [3,6].

Regarding imidazole medications, metronidazole (an antimicrobial) is a nitroimidazole derivative that can easily cross the blood-brain barrier. The characteristic pattern of involvement on MRI is the cerebellar dentate nuclei, but other affected areas may include the vestibular nuclei, cerebral PVWM, or callosal splenium, with or without dentate involvement (Fig. 6) [32]. In the current study, there were two such patients with ATL from metronidazole, one with characteristic dentate involvement, and the other having PVWM, callosal splenium, and bilateral corticospinal tract involvement. Hence, ATL should be considered in patients with PVWM DWI abnormalities receiving high doses of metronidazole, even if the dentate appears normal. This phenomenon can be entirely reversible both on imaging and clinically upon removal of the medication [32].

4.5. ‘C’ = “crack” or cocaine

The neuroimaging findings of cocaine toxicity can have either of two appearances related to differing underlying mechanisms: 1) ischemic events due to vasospasm, or 2) direct neurotoxicity causing diffuse PVWM injury [33]. Regarding direct WM toxicity, the abnormalities are best visualized on DWI or FLAIR MRI, with or without reduced diffusion [33]. While heroin-induced LE usually involves the posterior cerebral and cerebellar WM, cocaine-related LE usually involves the frontal lobes, often sparing the brainstem and cerebellum [34,35]. In the current study, only one such patient was diagnosed with cocaine-related ATL (Fig. 7).

4.6. ‘E’ = environmental (carbon monoxide) or ethanol abuse

Carbon monoxide (CO) is a colorless and odorless gas, the toxic levels of which may be present in several scenarios, such as smoke inhalation, suicide attempts, or abandoned buildings. The symptoms of acute CO intoxication vary, and may include chest or abdominal pain, dizziness, or headaches, or even coma and death [6]. CO toxicity can result from induced hypoxia, respiratory chain blockage, or direct cellular toxicity; such direct toxic effects on WM can result when CO activates polymorphonuclear leukocytes, causing lipid peroxidation and resultant acute demyelination [7,36]. In CO-related encephalopathy, there are three general patterns of involvement: PVWM (most common), basal ganglial, and hippocampal (least common), usually with ADC reduction in the acute phase (Fig. 8) [6,7,37,38]. The WM abnormalities are usually reversible, however basal ganglial insults are more prone to permanent damage due to the higher metabolic activity of those nuclei [6,38]. Accordingly, in this study, three such patients with CO-related ATL were noted, each having PVWM involvement. Interestingly, both CO and inhaled opiate neurotoxicity may appear similar, due to either toxic demyelination or spongiform degeneration, as both syndromes can involve the PVWM, basal ganglia, or rarely the hippocampi [2]. Thus, in patients presenting as acutely obtunded with PVWM involvement on MRI, toxicity screening for both CO and opiate toxicity may be warranted if the clinical history is limited. Ethanol (EtOH) neurotoxicity uncommonly causes ATL, albeit rare, as hippocampal edema (from withdrawal seizures) is the most common MRI finding of EtOH neurotoxicity; less common findings are Wernicke encephalopathy from thiamine deficiency, and rarely ATL [39]. The rare form of ATL that is EtOH-related is Marchiafava-Bignami Disease (MBD), which is an uncommon acute presentation of chronic EtOH abuse, with a rapid onset of altered mental status, gait abnormalities,
loss of consciousness, and speech impairment [40]. In the acute form on MRI, the genu and splenium of the corpus callosum are more commonly involved on FLAIR and DWI with reduced diffusion, and lesions may extend from the corpus callosum into adjacent PVWM, occasionally with involvement of other regions such as the cerebral cortex, middle cerebellar peduncles, and internal capsules [41]. In the late phase, there is hyperintensity on ADC map and FLAIR, with loss of the DWI-bright signal. In this regard, in the current study there were two EtOH-related ATL patients, one of which had the typical ATL-type of PVWM affection; the other appeared in the distribution of MBD with callosal and PVWM involvement (Fig. 9). Hence, MBD should be kept in mind in any acutely encephalopathic patients with a history of alcohol abuse and ATL [40,41].

4.7. ‘S’ = splenial lesions

Focal splenial lesions that spare, or only minimally involve, the PVWM are relatively uncommon abnormalities on DWI MRI in acutely encephalopathic patients, usually being related to anti-epileptic drugs (AEDs) [3,42,43]. While AEDs are likely the most commonly reported etiology of a reversible splenial lesion (RSL), RSL’s may also arise from chemotherapy, immunosuppressive medications, sepsis, metabolic disorders, or even antimicrobials such as metronidazole [3,42–44]. In this regard, the term RSL has also been alternated with the term “mild encephalitis with reversible splenial lesions” (MERS), a reversible phenomenon thought related solely to infectious etiologies (i.e. sepsis); however, the term RSL is used here to exclude the “infectious” subtypes, as AEDs have now been found to be the most common likely cause of RSLs, although infectious causes lacking exposure to an AED also occur [43–45]. In RSL, the abnormal signal on DWI MRI (often invisible on FLAIR) is either present solely within the splenium of the corpus callosum near the midline, or occasionally has additional, symmetric involvement of the posterior WM as well; the abnormalities typically do not enhance, and resolve within 2–3 weeks following the insult if the offending medication is withdrawn [2,3,43–45]. In the current study, an AED was the causative etiology in 7/9 patients, while the other two patients having RSL were from chemotherapy and immunosuppressant therapy (Fig. 10). Interestingly, as RSL variably involves the PVWM, it may be considered a focal subtype of ATL; another reason to consider it as a subtype (or relative) of ATL is that there is overlap with the various etiologies that may cause ATL [3,42,44,45]. Again, as sepsis may also cause RSL-type lesions, septic RSL (termed MERS) is discussed under...
Fig. 9. A 37 year old male with alcohol abuse presented with visual changes, who was ultimately diagnosed with ATL from Marchiafava-Bignami disease. 9A-C: On the initial MRI, reduced diffusion is noted in the body (dotted arrows) and the splenium (arrows) of the corpus callosum on DWI MRI (A) and ADC map (B), while these same areas are hyperintense on FLAIR (C). On a follow up MRI 2 months later, subsequent focal atrophy was present within the callosal splenium (arrows), having elevated diffusion that appears dark on DWI (D) that is less visible on SWI (E); focal atrophy is also noted within the callosal body (dotted arrow) on sagittal FLAIR (F).

Fig. 10. A 20 year old male with ganglioglioma on anti-epileptic therapy for 5 years presented with confusion, eventually being diagnosed with RSL-ATL. 10A-C: There is reduced diffusion within the callosal splenium (arrows) on DWI (A) and ADC map (B), with hyperintensity on FLAIR (C). 10C-D: The lesions resolved after 3 months, as on DWI (D), ADC (E) and FLAIR (F) sequences.
4.8. Mimics of ATL and ATL-like appearances

The terms ‘mimics’ or ‘ATL-like’ implies that the abnormalities are confined to PVWM with reduced diffusion in a symmetric and confluent fashion, usually lacking contrast enhancement. However, their characteristic imaging findings usually differ from that on sequences other than DWI, and may involve other anatomic locations. In addition, their clinical presentation is often discordant with ATL. These entities may appear similar to ATL, with corresponding PVWM involvement on DWI that causes their appearance to simulate ATL; such entities with PVWM DWI findings include metabolic entities such as osmotic demyelination syndrome (ODS, or EPM “extrapontine myelinolysis”), acute hepatic (or “hyperammonemic”) encephalopathy (AHE), as well as infectious processes (such as sepsis). However, additional areas of involvement within these disorders will usually hint that these are not true ATL.

As above, the ATL-like entities on DWI of PVWM include ODS/EPM, AHE, and MERS. Regarding ODS, while in “central” pontine myelinolysis, the pons is the predominant site of abnormality on MRI (especially DWI and FLAIR acutely), in “extrapontine myelinolysis”, the basal ganglia and PVWM may be involved, even without the pons (Fig. 11) [46]. Regarding AHE, it is a clinicoradiologic syndrome (usually with elevated plasma ammonia levels) which characteristically involves the insular cortex symmetrically on DWI or FLAIR MRI (both being > 80%), while the thalami (70%, 85%), posterior limb of internal capsule (80%, 75%), and PVWM (85%, 80%) can be involved on DWI and FLAIR, respectively; less commonly, more severe insults of the cortex diffusely or basal ganglia may occur, and the clinical and imaging findings can be reversible with therapy (Fig. 12) [47–49]. Finally, as mentioned above, sepsis-related MERS can mimic AED-related RSL/ATL; as the callosal splenium is part of the PVWM, we include ‘splenial lesions’ (most commonly due to AEDs) in the ‘CHOICES’ acronym, recognizing that MERS-type lesions related to sepsis can resemble RSL [2,3].

Mimics of ATL include the subacute phase (7–21 days) of hypoxic-ischemic encephalopathy (HIE, typically with cortical reduced diffusion and mild FLAIR hyperintensity in the acute phase, Fig. 13), acute disseminated encephalomyelitis (ADEM, typically asymmetric, multifocal, contrast-enhancing and possibly involving the basal ganglia, with only mild DWI abnormalities, Fig. 14), progressive multifocal leukoencephalopathy (PML, typically asymmetric, with elevated diffusivity on ADC within the PVWM, while having contrast enhancement in > 90%). Also, less common ATL mimics are PML-IRIS (an inflammatory response occurring after treatment of PML due to recovery of the immune system in the immunocompromised) (Fig. 15), acute necrotizing encephalopathy (usually T1-bright foci with hemorrhage involving the basal ganglia and cerebellum), and acute hemorrhagic leukoencephalitis (AHL, typically having cystic necrosis with only peripheral reduced diffusion).
diffusion and enhancement extending beyond the PVWM (Fig. 16) [2,3,50–55]. The latter three entities are rather rare, even relative to ATL, and while involving the PVWM, these syndromes have imaging appearances quite different from ATL on other MR sequences, and usually enhance avidly after the use of intravenous contrast. In practice, of these mimics, ADEM may be the one entity that might truly mimic ATL on neuroimaging, as a small percentage of ADEM patients resemble ATL by having confluent and symmetric abnormalities on DWI, and being limited to the PVWM, without contrast enhancement [51]. Another discriminating point regarding the subacute phase of HIE is that it

Fig. 12. A 53 year old male with chronic hepatic failure presented with acute confusion from acute hepatic/hyperammonemic encephalopathy (AHE). The patient had an elevated serum level of NH₄⁺ 101 µg/dl (normal: < 20-40 µg/dl). **12A-D**: bilateral symmetric PVWM involvement of the centrum semiovale is present on DWI (A) and FLAIR (B), also with involvement of the thalami (D arrows); involvement of the internal capsules’ posterior limbs (D dotted arrows) is noted on the ADC map (C) and FLAIR (D), being characteristic of AHE.
could mimic ATL on DWI and ADC maps, but the clinical history and imaging in the acute phase (1–6 days’ post-insult) usually clearly discern this entity from ATL [2,3,50]. Hence, these six entities were not included in the CHOICES acronym, as their typical MRI appearances are usually readily identifiable on imaging sequences other than DWI, they can occur in locations other than the PVWM, and the clinical history is usually discordant with a toxic insult.

This study is limited by its retrospective nature, descriptive/observational design, and the small number of patients; for example, only one patient had cocaine-related ATL. However, as this observational study is for the purposes of developing an acronym and a review about PVWM involvement of ATL, it should be sufficient to have a small number of patients for each letter in the acronym to aid in memorization of this potentially difficult differential diagnosis. Additionally,
another potential limitation is that in many patients, there were overlapping causes that gave rise to uncertainty of an exact etiology, such as with the ‘unspecified’ category for opioids.

5. Conclusion

In summary, we describe an acronym (“CHOICES”) for non-metabolic and non-infectious etiologies of ATL-related PVWM injury on DWI, and provide relevant examples. The intent is to increase awareness of possible non-metabolic and non-infectious etiologies of ATL due to its potentially reversible nature after treatment or removal of the toxin. Mimics of ATL and ATL-like disorders are described. Hopefully this improves recall of associated etiologies, which may result in prompt diagnosis and therapy.

Fig. 15. A 62 year old immunocompromised male, with PML, has “T2-shine through” (i.e. highly elevated diffusion) on DWI (A) and elevated diffusivity on ADC map (B); typically asymmetric lesions of PML are also seen (arrows) on FLAIR (C). Such multifocal lesions usually have an incomplete rim of enhancement (arrows) on postcontrast T1WI, suggestive for PML-IRIS (D).
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