INTRODUCTION

MELAS (mitochondrial myopathy, encephalopathy, lactic acidosis, and stroke-like episode) is a mitochondrial disorder that is characterized by delayed growth, episodic vomiting, seizures, muscle weakness, and recurrent cerebral injuries that resemble strokes. Although the neurological deficits in MELAS resemble features of ischemic stroke and multifocal cerebral infarct-like lesions are seen on the CT and MR images, they differ with a vascular stroke: MELAS lesions do not follow vascular territories or a border zone and the cerebral angiography fails to demonstrate any steno-occlusive lesion. Therefore, we can assume that the underlying mechanism of MELAS is different from that of an ischemic infarction (1). The main hypothesis explaining the pathophysiology of the infarct-like lesions of MELAS syndrome is angiopathy leading to ischemia (2) and the energy failure from defective oxidative metabolic pathways or the increased energy demand due to neuronal hyperexcitability (3).

Diffusion and Perfusion Characteristics of MELAS (Mitochondrial Myopathy, Encephalopathy, Lactic Acidosis, and Stroke-Like Episode) in Thirteen Patients

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Objective: We analyzed the diffusion and perfusion characteristics of acute MELAS (mitochondrial myopathy, encephalopathy, lactic acidosis, and stroke-like episode) lesions in a large series to investigate the controversial changes of the apparent diffusion coefficient (ADC) that were reported in prior studies.

Materials and Methods: We analyzed 44 newly appearing lesions during 28 stroke-like episodes in 13 patients with MELAS. We performed a visual assessment of the MR images including the ADC and perfusion maps, comparison of the ADC between the normal and abnormal areas, comparison of % ADC between the 44 MELAS lesions and the 30 acute ischemic infarcts. In addition, the patterns of evolution on follow-up MR images were analyzed.

Results: Decreased, increased, and normal ADCs were noted in 16 (36%), 16 (36%), and 12 (27%) lesions, respectively. The mean % ADC was 102 ± 40.9% in the MELAS and 64 ± 17.8% in the acute vascular infarcts (p < 0.001), while perfusion imaging demonstrated hyper-perfusion in six acute MELAS lesions. On follow-up images, resolution, progression, and tissue loss were noted in 10, 4, and 17 lesions, respectively.

Conclusion: The cytotoxic edema gradually evolves following an acute stroke-like episode in patients with MELAS, and this may overlap with hyper-perfusion and vasogenic edema. The edematous swelling may be reversible or it may evolve to encephalomalacia, suggesting irreversible damage.

Index terms: Mitochondrial encephalopathy; Diffusion weighted imaging; Perfusion MR imaging; Stroke
of a stroke-like episode as to whether it is vasogenic or cytotoxic in origin (4). There are conflicting reports regarding the diffusion characteristics of the infarct-like lesions of MELAS. Several reports (5–13) have described a normal or increased apparent diffusion coefficient (ADC), indicating the presence of vasogenic edema. These studies even suggested an increased ADC as a differential point from the cytotoxic edema of an ischemic origin. On the other hand, other studies (14–19) have described that ADC declines during an acute episode, suggesting the presence of cytotoxic edema even though the proposed pathophysiology was different from that of an ischemic stroke.

Because MELAS is a rare metabolic disorder, the majority of the previous studies have described only a small number of cases or the ADC description was not adequate. Therefore, a comprehensive analysis of the diffusion characteristics of acute MELAS lesions linked with a perfusion study in a large number of patients is necessary for a better understanding of the nature of edema in patients with MELAS, which might be different from that of an ischemic stroke.

We analyzed serial MR images, including DWI and perfusion images in 13 MELAS patients, with a particular emphasis on the changes of the ADC as compared with that of ischemic stroke.

MATERIALS AND METHODS

Our Institutional Review Board approved this retrospective study and informed consent was waived.

Subjects

We evaluated 13 patients with MELAS syndrome confirmed by gene analysis, which demonstrated the presence of a point mutation in the mitochondrial DNA (3243A>G). The subjects included five male and eight female patients ranging in age between 10 and 57 years (mean age: 32 years). The presenting symptoms included gaze deviation, mental state changes, tremors, seizures, motor weakness, aphasia, sensory change, and visual or hearing disturbance.

We reviewed a total of 43 sets of MR images and medical records over a 13 year period and we identified 48 acute brain lesions which appeared during 31 stroke-like episodes. Among them, four lesions that developed after three stroke-like events were excluded from the analysis since the corresponding DWI was not obtained. Ultimately, a total of 44 lesions that occurred over 28 stroke-like events were analyzed. The time interval between the acute onset of symptoms and the MRI scan ranged from 0–18 days (mean: 4.2 ± 3.6 days). Details of the individual lesions are summarized in Table 1. Perfusion MR imaging was performed in three patients (patients 5, 10, and 11) during five episodes and in one institution. The interval between symptom onset and perfusion imaging ranged between 0–3 days (mean: 0.8 ± 1.3 days).

Image Acquisition

The MRI examinations, including DWI, were subjected to 1.5- or 3.0-Tesla MRI units (Siemens, Erlangen, Germany/General Electric Medical Systems, Milwaukee, WI/Philips Medical Systems, Best, The Netherlands) in 13 patients from three institutions. A quadrature head coil was used as a receiver and transducer. The spin-echo T1-weighted axial and sagittal images, the fast spin-echo T2-weighted axial images and the fast fluid attenuated inversion recovery (FLAIR) axial images were obtained following the acquisition of diffusion-weighted axial images. A DWI was performed using a multi-slice, single shot, echo planar, spin echo pulse sequence. Two or three sets of images with different b-values (0, 500, and 1000 sec/mm$^2$) were acquired and the ADC maps were created with the obtained signal.

Perfusion imaging was performed using an EPI spin-echo sequence (repetition time 2000 msec, echo time 60 msec), which targeted the central region of infarct-like lesions. Seven axial slices (slice thickness, 5 mm; interslice gap, 2 mm; field of view 260 mm; matrix size 128 x 128) were acquired. The positions of the slices were selected to match the acute lesions on DWI. Ten slices were imaged 50 times per 2 second, with a total imaging time of 1 minute and 40 seconds. Four sets of baseline images were collected before gadopentetate dimeglumine (0.2 mmol/kg) was injected at a speed of 3 mL/s using a MRI-compatible power injector. The signal intensity-time curves were obtained from the area of acute lesion and the contralateral unaffected area. In addition, the maps for regional cerebral blood flow (CBF), cerebral blood volume (CBV), time to peak enhancement, and mean transit time were generated.

Image Analysis

Visual assessment of the cortical and white matter involvement in conventional MR images and the signal
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changes on DWI, ADC, and the perfusion maps, were performed for each new edematous lesion by two radiologists (with 15 years and 16 years of neuroimaging interpretation experience, respectively) in consensus.

To quantify the changes of ADCs independent of the tissue, three regions of interest (ROIs) were positioned in each of the affected areas and in the control regions in the contralateral hemisphere (Fig. 1D). In addition, the mean ADC of the lesions and the control area were compared using paired t-tests. The size of the ROIs positioned at the lesion and control area were the similar, ranging from 5–15 mm² according to the size of the involved area. The ROI measurement was performed by one radiologist who did not participate in the visual assessment. In the cases with bilateral lesions, the corresponding gray or white matter of the normal appearing parenchyma was selected as a control. In addition, the % ADC ([the mean ADC in the infarct-like lesion/the mean ADC in the control area] x 100) was calculated for each lesion. To compare the decline of the ADC between the MELAS lesions and the ischemic infarct, we selected 30 acute vascular infarcts that were detected on the MR images obtained from 0 to 18 days after a stroke (mean: 2.2 ± 4.2 days) in 25 patients, and their mean % ADCs were compared with those of the MELAS lesions by way of independent t-tests. In addition, scatterplots of the % ADC values versus time were drawn up to evaluate all the lesions in the patients with MELAS and ischemic stroke.

The changes of each newly identified acute lesion seen on the follow-up MR images were reviewed and we evaluated whether there was any significant difference in the patterns of evolution between the lesions with a decreased ADC and the lesions with a normal or increased ADC by performing the Fisher’s exact test.

Statistical analyses were performed with the SPSS

Fig. 1. 11-year-old girl with gradually progressive stroke lesions.
A. T2-weighted image obtained three days after first attack shows mild swelling (arrows) of right temporocipital lobe. B. Follow-up T2-weighted image obtained 11 days after first symptom onset and one day after mental change revealed progression of edema in right temporocipital lobe and newly appearing thalamic lesion (arrowhead). C. Affected areas are seen as having high signal intensity (arrowhead) on diffusion-weighted MR imaging. D. Apparent diffusion coefficient decline is prominent at white matter, whereas swollen cortex and right thalamic lesion (arrowhead) show relatively increased or iso-signal intensity on apparent diffusion coefficient map. E. T1-weighted image shows high signal rim (arrows) along cortex of swollen right temporocipital lobe, suggesting cortical laminar necrosis.
software (SPSS, version 17.0; Chicago, IL) and all of the statistical evaluations were considered significant for $p$ values less than 0.05.

**RESULTS**

**Visual Assessment**

The newly affected lesions appeared as infarct-like gyral swelling (Figs. 1–3) in all the hemispheric lesions ($n = 40$) except in the case of a small separated white matter lesion and adjacent white matter involvement in 24 of the 40 (60%) cortical lesions. The three remaining lesions were noted in the thalamus ($n = 2$) and putamen ($n = 1$).

Diffusion weighted MR imaging revealed the presence of hyperintense edema in 37 lesions (84%, Figs. 1C, 2D) and iso-signal intensity in seven lesions (16%). The ADC maps revealed a dark area within the infarct-like lesion in 16 (36%, Figs. 1D, 3C) lesions and a bright area in 16 (36%) lesions (Fig. 2E). The remaining 12 (27%) lesions exhibited iso-signal intensity compared to the control area. The MRI findings, including the diffusion characteristics of each lesion, are summarized in Table 1.

A total of 11 lesions were identified on perfusion images obtained during five episodes in three patients. Perfusion imaging demonstrated increased CBF (Figs. 2B, 3D, 3E) and CBV in six newly appearing lesions, and this suggested a hyper-perfused state regardless of the ADC findings (the ADC was increased in four, decreased in one, and not changed in one), while the five chronic lesions exhibited decreased perfusion (Fig. 3E).

**Quantitative Analysis**

The mean ADC in the acutely affected areas and the normal control regions were $0.733 \pm 0.32$ and $0.758 \pm 0.16 \times 10^{-3} \text{mm}^2/\text{s}$, respectively ($p = 0.596$). Each mean % ADC at the lesions exhibiting different signal intensities on DWI and ADC map are summarized in Table 2. The mean % ADC was $103 \pm 41.2\%$ in the acute lesions of MELAS, compared to $65 \pm 33.3\%$ in the acute vascular infarct ($p < 0.001$). The distribution of the % ADC according to the elapsed time since the onset of acute symptoms in the MELAS lesions (Fig. 4A) showed a random distribution, below or above 100%, with the majority being more than 50% until the 18th day after the onset of acute symptoms. However, many of the % ADCs in the vascular infarcts were less than 50% at the stroke onset and then gradually increased to remain less than 100% until the 5th day (Fig. 4B).

**Evolution of the Lesions**

The follow-up MR images were available for 31 of the 44 lesions (70%) and the total follow-up period ranged between eight days and 67 months (mean: 11 ± 16 months). On the follow-up MR images, complete resolution or regression with a small area of altered signal intensity

### Table 1. Summary of the MRI Findings in 44 Acute Edematous Lesions

| Case | Number of Episodes | Timing of MRI* | Number of Lesions | DWI High/Low/Iso | ADC High/Low/Iso |
|------|--------------------|----------------|------------------|-----------------|-----------------|
| 1    | 3                  | 3, 0, 5D       | 3                | 3/0/0           | 1/2/0           |
| 2    | 1                  | 3D             | 4                | 4/0/0           | 0/1/3           |
| 3    | 3                  | 2, 6, 3D       | 5                | 5/0/0           | 0/2/3           |
| 4    | 1                  | 14D            | 2                | 2/0/0           | 0/2/0           |
| 5    | 4                  | 0, 5, 1, 0D    | 6                | 6/0/0           | 5/0/1           |
| 6    | 2                  | 3, 1D          | 3                | 3/0/0           | 2/1/0           |
| 7    | 3                  | 2, 7, 2D       | 6                | 1/0/5           | 2/1/3           |
| 8    | 2                  | 7, 3D          | 3                | 3/0/0           | 1/2/0           |
| 9    | 3                  | 3, 11, 1D      | 4                | 4/0/0           | 2/2/0           |
| 10   | 2                  | 18, 0D         | 2                | 2/0/0           | 1/1/0           |
| 11   | 2                  | 1, 3D          | 2                | 2/0/0           | 1/1/0           |
| 12   | 1                  | 5D             | 1                | 1/0/0           | 0/1/0           |
| 13   | 1                  | 0D             | 3                | 1/0/2           | 1/0/2           |

Note.— *Time interval between acute symptom onset and MRI scan.
High, low, and iso indicate increased, decreased, and iso-signal intensity as compared to normal control region.
ADC = apparent diffusion coefficient, D = day(s), DWI = diffusion-weighted MR imaging.
was seen in 10 lesions (32%), whereas 17 lesions (55%) evolved to encephalomalacia with or without residual abnormal signal intensity (Fig. 2F). For three lesions, the progressive spread of the edema occurred over 5–8 days, which eventually evolved to atrophy on the later follow-up images (Figs. 1, 2). An apparently progressive extent of the edematous lesion was seen in the remaining patient (patient 11) who underwent a second MR imaging 17 months after the initial scan. However, the different diffusion and perfusion characteristics within the lesion (ADC decline and hyper-perfusion at the posterior acute lesion, increased ADC

Table 2. ADC Findings in 44 Acute Lesions on DWI

| SI* on ADC map | Number of Lesions (%) | Mean % ADC ± SD |
|----------------|-----------------------|-----------------|
| Low            | 16 (36)               | 61 ± 16.7       |
| Iso            | 12 (27)               | 102 ± 5.3       |
| High           | 16 (36)               | 145 ± 28        |
| Total          | 44 (100)              | 103 ± 41.2      |

Note.— *SI means signal intensity. Low, iso, and high indicate increased, decreased, and iso-signal intensity as compared to normal control region. ADC = apparent diffusion coefficient, SD = standard deviation.
and hypo-perfusion at the anterior old lesion) suggested that
the newly appearing acute lesion overlapped with a chronic
lesion (Fig. 3). Cortical laminar necrosis was seen in four
lesions (Fig. 1E). On the follow-up MR images, the majority
of the lesions became faint or disappeared on DWI, and
exhibited increased ADC except for three progressive lesions
that exhibited a further decrease in ADC over the short-term
follow-up period. There was no significant difference in the
patterns of evolution (resolved, tissue loss, or progressed)
of the edematous lesions according to the ADC profile (high,
low, or iso-signal intensity compared to the control area)
during the acute stage ($p = 0.44$) (Table 3).

**DISCUSSION**

Diffusion-weighted MR imaging reflects the random
translational motion of water molecules and it is sensitive
to changes in the biological environment associated with
various pathophysiological processes. Increased ADC values

| Table 3. Evolution of Edematous Lesions according to Initial Apparent Diffusion Coefficient Findings |
|---------------------------------------------------------------|
| Low | Iso | High | Total |
| Resolved | 4  | 3  | 3  | 10 |
| Progressed edema | 1  | 1  | 24 |
| Cortical laminar necrosis | 4  | 0  | 0  | 4 |
| Residual lesion and tissue loss | 10 | 3  | 4  | 17 |

Note.—Some of lesions exhibited more than one evolution
pattern on follow-up images.
Low, iso, and high indicate increased, decreased, and iso-signal
intensity as compared to normal control region respectively.

![Fig. 3. 57-year-old man with two stroke-like episodes.](image)

A. Fluid attenuated inversion recovery image obtained 18 days after first symptom onset shows infract-like lesion at left temporal lobe (arrow).
B. T2-weighted image obtained on day of second attack and 17 months after prior episode shows high signal intensity in left temporoparietal lobe, including prior lesion.
C. Apparent diffusion coefficient map shows diffusion restriction in posterior region of lesion (arrow), while anterior old lesion shows increased apparent diffusion coefficient (arrowhead).
D. Perfusion curves (relaxivity on vertical axis and time on horizontal axis) show increased blood volume (which is proportional to signal loss) in left occipital lesion (green curve) compared to that of contralateral normal tissue (purple curve).
E. Cerebral blood flow map demonstrates hyper-perfusion at newly appearing left occipital lesion (arrow), while anterior chronic lesion shows decreased perfusion (arrowheads).
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correspond to increased extracellular space in vasogenic edema while cytotoxic edema in acute ischemic stroke causes reduced ADC values (4). The decline in ADC was also described to occur in association with severe hypoglycemia (20), status epilepticus (21), and cortical spreading depression (22).

In this study, the acute infarct-like lesions frequently exhibited a decrease in ADC, and although there were many reversible lesions, a large proportion of the edematous lesions eventually evolved to localized tissue loss, suggesting irreversible damage. Indeed, we increasingly observed areas of restricted diffusion during the early stage of MELAS (Table 4), and cytotoxic edema is the most likely cause of the apparent diffusion restriction. Prolonged mitochondrial energy failure may lead to ion pump failure and then cytotoxic edema will eventually develop.

However, an increase in ADC during the early phase also occurred in our study as well as in many prior reports (Table 4). In these settings, the occurrence of vasogenic edema due to a breakdown of the blood-brain barrier has been implicated as a cause of stroke-like lesions. The presence of vasogenic edema is supported by hyper-perfusion foci in the acute lesions of MELAS on the perfusion MR imaging in our study, which was also demonstrated in a recent study (23), which contrasts with the typical hypo-perfusion in acute vascular stroke. We can speculate that the increased regional lactate level might cause a change of the local vascular permeability or auto-regulation at the beginning of a stroke-like episode, and eventually leads to vasogenic edema and increased perfusion. This theory is also supported by the hyper-perfusion of the affected area on SPECT (single photon emission computed tomography) (3), abnormal mitochondria within the vascular smooth muscle and endothelial cells (suggesting impaired auto-regulation) (24), and vasodilation seen on angiography or dynamic CT (25).

In addition, we can assume that those patients exhibiting a normal or increased ADC during the early period of a stroke-like event were possibly examined after the time point of pseudonormalization of the cytotoxic edema because the symptoms in MELAS are often nonspecific or they are not sufficiently severe or acute enough to receive medical attention. The described conflicting findings may also reflect the observation that some of the lesions disappear after a couple of weeks, whereas others are newly developed or persist.

Although the lack of diffusion restriction is not the criteria to differentiate MELAS from the classic ischemic stroke anymore, there are several different patterns of ADC decline in MELAS compared to that of ischemic stroke. Our data indicates that the % ADC was significantly higher in MELAS compared to the observed % ADC reduction in acute ischemic stroke. Iizuka et al. (15) also reported only a slight difference in ADC values between the stroke-like lesions and the non-affected area, which suggests concomitant vasogenic edema. The initial neuronal injury may cause cytotoxic edema and a low ADC, while the subsequent hyperperfusion causes increased blood in the voxels, and the extracellular edema may gradually increase the ADC. As the lesions evolve and new areas get involved, both processes overlap temporally, which leads to a mixture of cytotoxic and vasogenic edema (18).

Another different pattern of ADC reduction in MELAS is manifested as the time point of ADC decline being much

![Graph A](image1.png)  
**Fig. 4.** Scatter plots of the % apparent diffusion coefficient (vertical axis) according to elapsed time (days) after symptom onset (horizontal axis) in mitochondrial myopathy, encephalopathy, lactic acidosis, and stroke-like episode lesions (A) and lesions of ischemic infarcts (B).
| Author                  | Year | Case | Episode | Timing of MRI | DWI      | ADC       |
|------------------------|------|------|---------|---------------|----------|-----------|
| Yoneda et al. (5)      | 1999 | 1    | 1       | 3D            | NR       | High      |
|                        |      |      |         | 10D           | NR       | High      |
|                        |      |      |         | 17D           | NR       | High      |
|                        |      |      |         | 25D           | NR       | Iso       |
| Oppenheim et al. (6)   | 2000 | 2    | 1       | 2D            | High     | High/Iso  |
| Ohshita et al. (7)     | 2000 | 3    | 1       | 34D           | High     | NR        |
|                        |      |      |         | 2             | 15D      | High      |
|                        |      |      |         | 30D           | High     | High      |
|                        |      |      |         | 150D          | High     | Iso       |
|                        |      |      | 4       | 1             | 240D     | High      |
|                        |      |      |         | 2             | 5D       | High      |
|                        |      |      |         | 30D           | High     | High      |
|                        |      |      |         | 150D          | High     | Iso/Iso   |
| Yonemura et al. (8)    | 2001 | 5    | 1       | 2D            | High     | High      |
|                        |      |      |         | 2             | 2D       | High      |
| Kolb et al. (9)        | 2003 | 6    | 1       | NR            | High     | High      |
|                        |      |      |         | 7             | 1        | 2D        | High      |
| Abe et al. (10)        | 2004 | 8–19 | NR      | within 14D    | High     | High*     |
| Jian-Ren (11)          | 2005 | 20   | 1       | 10D           | High     | High      |
| Alemdar et al. (12)    | 2007 | 21   | 1       | NR            | High     | High      |
| Ito et al. (13)        | 2008 | 22   | 1       | 0             | High     | High      |
| Wang et al. (14)       | 2003 | 23   | 1       | 5D            | High     | Low       |
|                        |      |      |         | 11D           | High     | Low       |
|                        |      |      |         | 194D          | High     | High      |
| Iizuka et al. (15)     | 2003 | 24   | 1       | 4/9D          | High     | Low       |
|                        |      |      |         | 25             | 12/17D   | High      |
|                        |      |      |         | 26             | 1        | 1D        | High      |
|                        |      |      |         | 2             | 6D       | High      |
| Mizrachi et al. (16)   | 2006 | 27   | 1       | Days to weeks | High     | Iso       |
|                        |      |      |         | Days          | High     | Low       |
|                        |      |      |         | 30D           | High     | Iso       |
| Bi et al. (17)         | 2006 | 28   | 1       | 1D            | High     | Iso       |
|                        |      |      |         | 3D             | High     | Low       |
|                        |      |      |         | 21D           | High     | High/Low  |
|                        |      |      |         | 150D          | High     | High      |
|                        |      |      |         | 2             | 2D       | High      |
| Tzoulis et al. (18)    | 2008 | 29   | 1       | 8D            | High     | High/Low  |
|                        |      |      |         | 11D           | High     | Low       |
|                        |      |      |         | 58D           | High     | High      |
|                        |      |      | 2       | 13D           | High     | High/Low  |
| Sheerin et al. (19)    | 2008 | 30   | 1       | NR            | High     | Low       |

**Note.**— ADC = apparent diffusion coefficient, D = days, DWI = diffusion-weighted MR imaging, MELAS = mitochondrial myopathy, encephalopathy, lactic acidosis, and stroke-like episode, NR = not reported.
delayed and prolonged compared to that of ischemic stroke. The DWI changes in the MELAS syndrome did not appear until two days after the acute attack in a large series study (10), while restricted diffusion was seen after several days or even weeks after the event in previously published studies (Table 4) as well as in our study. Conversely, the DWI shows diffusion changes within hours following the vascular stroke, which is followed by a gradual increase in ADC up to reach a normal value (pseudonormalization) at about a week after the ischemic event. Therefore, it is postulated that unlike the acute ischemia of a vascular infarct, the cytotoxic edema of MELAS perhaps comes from gradually evolving metabolic cell death associated with mitochondrial dysfunction (15, 17).

Another important feature of the ADC change of MELAS is the gradual spread of the core of the edematous lesion, which is another reason for the delayed manifestation of a declined ADC. Several reports have demonstrated a slowly progressive and spreading pattern of this infarct-like lesion (14, 15, 17, 18), which was also observed in our series (Figs. 1, 2). Such a slowly progressive and spreading pattern has never been described for ischemic stroke.

Slow progression of the edema can be also seen from the cortical gray matter to the adjacent white matter. The gray matter of the cortex is particularly sensitive to changes in energy balance with its large energy demand (19) and the majority of the infarct-like lesions in MELAS preferentially involve the gray matter, with or without white matter extension as a function of the severity and acuity of the injury. Such a temporal progression can be seen both on the serial conventional MR images and DWI, and is evidenced by a shift in the declining ADC to the white matter, while the fully swollen cortical grey matter shows normal or increased ADC on short-term follow-up images (Fig. 1).

The limitations of this retrospective study included the variable timing of the MR imaging after symptom onset and the irregular follow-up time. Moreover, the technique of DWI was not uniform because the data was collected from three institutions for a long time period. To overcome this limitation, we used the % ADC rather than the absolute ADC value for comparison. Another limitation is that the perfusion imaging was performed in only five of the 28 episodes. Further study on both diffusion and perfusion analysis during acute MELAS lesions in a large series may provide more generalized information.

In conclusion, diverse ADC changes suggesting the gradual evolution of the cytotoxic edema that overlapped with hyper-perfusion and vasogenic edema following a stroke-like episode are the distinguishable features of MELAS, as compared with the features of an ischemic stroke. Although there are some reversible lesions, the edematous swelling may evolve to the progressive expansion of extensive cortical laminar necrosis and eventually to irreversible encephalomalacia with chronic abnormal signal intensity. This ultimately suggests the presence of irreversible damage, regardless of the initial ADC changes.

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