Clinical and Imaging Features of Multiple System Atrophy: Challenges for an Early and Clinically Definitive Diagnosis

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ABSTRACT

Multiple system atrophy (MSA) is an adult-onset, progressive neurodegenerative disorder. Patients with MSA show various phenotypes during the course of their illness, including parkinsonism, cerebellar ataxia, autonomic failure, and pyramidal signs. Patients with MSA sometimes present with isolated autonomic failure or motor symptoms/signs. The median duration from onset to the concomitant appearance of motor and autonomic symptoms is approximately 2 years but can range up to 14 years. As the presence of both motor and autonomic symptoms is essential for the current diagnostic criteria, early diagnosis is difficult when patients present with isolated autonomic failure or motor symptoms/signs. In contrast, patients with MSA may show severe autonomic failure and die before the presentation of motor symptoms/signs, which are currently required for the diagnosis of MSA. Recent studies have also revealed that patients with MSA may show nonsupporting features of MSA such as dementia, hallucinations, and vertical gaze palsy. To establish early diagnostic criteria and clinically definitive categorization for the successful development of disease-modifying therapy or symptomatic interventions for MSA, research should focus on the isolated phase and atypical symptoms to develop specific clinical, imaging, and fluid biomarkers that satisfy the requirements for objectivity, for semi- or quantitative measurements, and for uncomplicated, worldwide availability. Several novel techniques, such as automated compartmentalization of the brain into multiple parcels for the quantification of gray and white matter volumes on an individual basis and the visualization of α-synuclein and other candidate serum and cerebrospinal fluid biomarkers, may be promising for the early and clinically definitive diagnosis of MSA.

Key Words
Atypical symptom; diagnostic criteria; biomarker; early diagnosis.
INTRODUCTION

Multiple system atrophy (MSA) is defined as an adult-onset, rare, sporadic, and fetal neurodegenerative disease. The term MSA was proposed as encompassing three unrelated diseases: olivopontocerebellar atrophy (OPCA),1 Shy-Drager syndrome (SDS),2 and striatonigral degenerative disease (SND)3 in 1969 by Graham and Oppenheimer.4 In the same year, Takahashi et al.5 independently reported similar concepts: 1) considerable common clinical and pathological features exist between OPCA and SDS, 2) both are nosologically allied conditions based on a detailed review of the literature, and 3) apparent striatonigral changes were similar to those implicated in the pathology of SND.

Although there have been several discussions as to whether OPCA, SDS, and SND comprise a single disease or not, the discovery of the pathological hallmark, argyrophilic aggregates in the cytoplasm of oligodendrocytes, termed glial cytoplasmic inclusions (GCIs), resolved the dispute.6,7 Subsequently, α-synuclein (α-syn) was proven to be the main component of GCIs in 19978 and MSA was classified as an α-synucleinopathy together with Parkinson’s disease (PD) and dementia with Lewy bodies (DLB).

Based on these findings, the first consensus statement on the diagnosis of MSA was proposed in 1998.9 In this statement, three levels of certainty were established: possible, probable, and definite. Definite MSA requires neuropathological confirmation. The combination of severe autonomic failure/urinary dysfunction plus poorly levodopa-responsive parkinsonism or cerebellar ataxia is required for a diagnosis of probable MSA. Patients with predominant parkinsonism are designated MSA-P and those with predominant cerebellar ataxia as MSA-C. It was also recommended to not use the term SDS.

In 2008, the second consensus statement was established.10 These revised criteria also required rigorously defined autonomic failure and poorly levodopa-responsive parkinsonism or cerebellar ataxia for the diagnosis of probable MSA, similar to the first consensus statement (Figure 1). Possible MSA required parkinsonism or cerebellar ataxia and at least one feature suggesting autonomic dysfunction that did not fulfill the level required for probable MSA plus one other feature defined by clinical findings such as a Babinski sign with hyperreflexia, stridor, rapid progression rate, poor response to levodopa, and ataxia or characteristic neuroimaging abnormalities based on magnetic resonance imaging (MRI), 18 fluoro-2-deoxyglucose positron emission tomography (FDG-PET), or presynaptic nigrostriatal dopaminergic PET/single photon emission computed tomography (SPECT) (Figure 2). Features supporting (red flags) and nonsupporting a diagnosis of MSA were also proposed (Table 1). Red flags are other clinical features based on the literature reviews and expert opinion that may raise the clinical suspicion of MSA, particularly for differentiating it from PD.11 Classic pill-rolling rest tremor, clinically significant neuropathy, hallucinations not induced by medications, older age at onset (> 75 years), family history, dementia (as defined by the Diagnostic and Statistical Manual of Mental Disorders fourth edition, DSM-IV), and white matter lesions suggesting multiple sclerosis are nonsupporting features.

Although several developments and refinements were incorporated in the second consensus criteria compared to the first, improvements in early diagnosis of MSA were limited (sensitivity of first clinical visit: 41% of possible MSA based on the second consensus criteria, 28% of possible MSA based on the first consensus criteria).12 Only 18% of patients fulfilled the second consensus criteria for probable MSA at the first clinical visit, although the positive predictive value was high and ranged from 86% to 100%. However, a more recent study showed that 62% of patients clinically diagnosed with MSA had
Early Diagnosis of MSA

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Table 1. Supporting and nonsupporting features of multiple system atrophy

| Supporting features (Red flag signs) |
|-------------------------------------|
| Orofacial dystonia                   |
| Disproportionate antecollis          |
| Camptocormia and/or Pisa syndrome    |
| Contractures of hands or feet        |
| Inspiratory sighs                    |
| Severe dysphonia                     |
| Severe dysarthria                    |
| New or increased snoring            |
| Cold hands and feet                  |
| Pathologic laughter or crying        |
| Jerky, myoclonic postural/action tremor |

| Nonsupporting features               |
|-------------------------------------|
| Classic pill-rolling rest tremor     |
| Clinically significant neuropathy    |
| Hallucination not induced by drug    |
| Onset after age 75 years             |
| Family history of ataxia or parkinsonism |
| Dementia (on DSM-IV)                 |
| White matter lesions suggesting multiple sclerosis |

DSM-IV: Diagnostic and Statistical Manual of Mental Disorders fourth edition.

Figure 2. Criteria for the diagnosis of probable MSA. Possible MSA requires parkinsonism or cerebellar ataxia and at least one feature suggesting autonomic dysfunction that does not fulfill the level required for fulfilling probable MSA plus one of other features defined by clinical findings such as Babinski sign with hyperreflexia, stridor, rapid progression rate, poor response to levodopa and ataxia or characteristic neuroimaging abnormalities based on MRI, FDG-PET, and presynaptic nigrostriatal dopaminergic PET/SPECT depending on MSA-P or MSA-C. MSA: multiple system atrophy, MRI: magnetic resonance imaging, FDG-PET: 18 fluoro-2-deoxyglucose positron emission tomography, SPECT: single photon emission computed tomography, MSA-P: MSA predominantly parkinsonism, MSA-C: MSA predominantly cerebellar ataxia, OH: orthostatic hypotension, MCP: middle cerebellar peduncle, DA: dopaminergic.

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received the correct diagnosis as shown at autopsy. The presence of autonomic failure and cerebellar ataxia were the leading causes of misdiagnosis of DLB and progressive supranuclear palsy (PSP), respectively. Thus, a revision of the second consensus criteria for the early and clinically definitive diagnosis of MSA is urgently needed.

In this article, we review the clinical features that can become obstacles for the early diagnosis of MSA and discuss the development of recent diagnostic biomarkers.

**MONOSYSTEM ATROPHY**

**Time from onset to probable MSA**

The median time from initial symptom presentation to combined motor and autonomic dysfunction in MSA (probable MSA) is 2 years but ranges from 1 to 10 years. Initial onset to the presence of concomitant motor and autonomic manifestations at 2, 4, and 6 years occurred in 57.4, 83.5, and 96.5% of patients, respectively. In other words, more than 40% of patients will visit the hospital with isolated motor
or autonomic impairment and not be diagnosed as probable MSA within 2 years of onset. Since the median time from onset to death is 9 years but that from receiving a probable MSA diagnosis to death is 6 years, patients with probable MSA may be at severely advanced stages at diagnosis. Thus, it is important to focus on the clinical characteristics of the isolated autonomic failure or motor impairment (mono system atrophy) stage for early diagnosis.

**Isolated autonomic failure phase in MSA**

PD, DLB, and pure autonomic failure (PAF), classified as α-synucleinopathies, are the most important neurodegenerative diseases to consider in the differential diagnosis of MSA. Autonomic failure is commonly observed in all α-synucleinopathies. Severe autonomic failure that was classically considered an exclusion criterion for the diagnosis of PD is one of the red flags but not an absolute exclusion criterion according to the Movement Disorders Society. In the findings of a meta-analysis, orthostatic hypotension (OH) was observed in more than 30% of patients with PD. Patients with DLB show autonomic failure more frequently than PD. Interestingly, a prospective cohort study on PAF reported that 25 out of 74 subjects with severe OH (34%) developed DLB (n = 13), PD (n = 6), or MSA (n = 6) within 4 years of follow up, supporting the idea that PD, DLB, PAF, and MSA can show isolated autonomic failure prior to reaching the full-blown stage.

We should also pay attention to the nonmotor MSA phenotype. Patients with nonmotor MSA are rare but present with severe autonomic failure leading to sudden death or respiratory failure in the absence of diagnostic motor symptoms/signs of MSA. In our autopsy series from 1976 to 2014, four of 161 patients fulfilled the criteria for nonmotor MSA. The time from onset to death was 1.3 to 2 years. A pathological study showed severe involvement of the medullary serotonergic neurons, particularly in the nucleus raphe obscurus and nucleus of the ventrolateral medulla. A group of serotonergic neurons of the nucleus raphe obscurus projects to phrenic and other respiratory motoneurons and is intrinsically sensitive to levels of CO2 associated with respiratory dysfunction, such as impaired sensitivity to hypercapnia, suggesting an increased risk of sudden death.

Cell loss of serotonin-positive pre-Bötzinger complex cells as an essential part of the medullary respiratory network located in a circumscribed area of the ventrolateral medulla may also contribute to breathing disorders in MSA.

Interestingly, patients with minimal change MSA with sudden death also showed marked involvement of medullary serotonergic neurons in our study. Minimal change MSA presented with definite parkinsonism and dysautonomia but showed selective cell loss in the substantia nigra and locus coeruleus in combination with widespread GCI and astrogliosis. Significant respiratory dysfunction and early OH were observed in minimal change MSA. The clinicopathological findings of minimal change MSA were closely related to those of nonmotor MSA, suggesting that these two phenotypes belong to a continuous pathological spectrum. Nonmotor MSA is a noteworthy clinicopathological variant associated with a risk of unexpected death prior to fulfilling the current diagnostic criteria, which may arise from pathological involvement of the medullary serotonergic system.

Although risk factors for a poor prognosis differ among reports, several natural history studies have found that severe autonomic failure and time from onset to concomitant presence of autonomic failure and motor involvement are commonly associated with rapid progression and survival. It is urgent to establish a strategy for the early diagnosis of MSA during the isolated autonomic failure phase.

**Isolated parkinsonism or cerebellar ataxia phase in MSA**

Patients with MSA often show isolated parkinsonism or cerebellar ataxia. Particularly, the diagnosis of patients with a long duration of the isolated phase without red-flag signs is difficult. Petrovic et al. reported that four of 135 autopsy cases showed MSA-P with slow progression and prolonged survival in the Queen Square Brain Bank. Although the mean survival time of patients with MSA is approximately 9 years, the disease duration of these four patients with MSA-P was 15 years or longer. Interestingly, the latency from the onset of parkinsonism to autonomic failure ranged from 9 to 14 years. The initial levodopa response was good in one, moderate in two, and poor in one patient. All patients developed dyskinesia after a mean duration of levodopa treatment of 4.3 years with generalized chorea. Red-flag signs were observed in all four pa-
tients. The mean latency of red-flag signs from onset was 9 years. Thus, the early diagnosis of these patients would not have been possible using the current consensus statement.

The term idiopathic late onset cerebellar ataxia (ILOCA) was proposed in 1981. ILOCA is a slowly progressive, adult-onset ataxia and includes both ataxia and urinary dysfunction, abnormal reflexes, and dementia. Currently, ILOCA is considered to include MSA-C, spinocerebellar ataxias (SCAs), fragile X-associated tremor ataxia syndrome, Friedreich’s ataxia, autosomal recessive cerebellar ataxia type 1, and other novel genetic disorders. Thus, it is very important to distinguish early in the disease course between patients with MSA-C who show isolated cerebellar ataxia and those with other causes of ILOCA. Gilman et al. reported that the median time from isolated cerebellar ataxia (ILOCA phenotype) to the combination of cerebellar ataxia and autonomic failure (MSA) was 4.5 years.

Figure 3 summarize the many faces of MSA. To diagnose patients with MSA who show isolated parkinsonism or cerebellar ataxia during the early course of the illness, other specific imaging modalities or biomarkers will be necessary.

ATYPICAL SYMPTOMS IN MSA

Dementia and hallucination in MSA

Dementia based on the DSM-IV is a nonsupporting feature of MSA. However, patients with MSA frequently have fronto-executive dysfunction and may show impairment in memory and visuospatial function. Imaging and neuropathological findings support the concept that striatofrontal deafferentation, cortical degeneration, and cerebellar pathology could be associated with cognitive impairments in MSA.

A recent multicenter study using gray- and white-matter voxel-based morphometry and fully automated subcortical segmentation showed that only focal volume reduction in the left dorsolateral prefrontal cortex was observed in patients with MSA with cognitive decline compared to those with normal cognition, suggesting only a marginal contribution of cortical pathology to cognitive deficits. Previously, we reported that brain SPECT showed a significant correlation between neuropsychological impairment and decreased perfusion in the prefrontal cortex in patients with MSA-P. 18F-FDG brain PET showed that glucose metabolism in the striatum was the most powerful determinant of glucose metabolism in the frontal cortex in MSA-P. Deafferentation due to focal degeneration of striatofrontal circuits may play an important role in frontal-executive dysfunction in some patients with MSA-P. However, it remains undetermined whether putamen- or white matter-tract involvement correlates with deafferentation and cognitive decline in MSA-P.

In contrast, longitudinal studies have shown that progressive cerebral atrophy could occur in patients with MSA. Kitayama et al. reported that 10 of 58 patients with MSA (17%) had dementia fulfilling the DSM-IV criteria. More recently, autopsy-confirmed MSA cases that presented a clinical disease type of frontotemporal dementia (one with behavioral variant frontotemporal dementia, one with progressive nonfluent aphasia, and two with corticobasal syndrome) were reported. These results also support the view that cortical changes may influence cognitive decline in MSA.
Patients with MSA-C could also show similar cognitive decline compared to patients with MSA-P, but the pathogenic mechanism of neural correlates causing the cognitive impairments in MSA-C remains undetermined. It is well known that the greater part of the human cerebellum is associated with cerebral networks involved in cognition. Since the frontal cortex and the cerebellar networks are composed of polysynaptic circuits, analysis of functional imaging such as of resting-state networks could provide insight into subcortical cognitive decline in MSA-C.

Hallucinations not induced by drugs are also nonsupporting features of MSA. According to the results provided by the European Multiple System Atrophy registry, the frequency of patients experiencing hallucinations was only 5.5%. A retrospective autopsy study reported that 9% of patients with MSA had hallucinations. More recently, Koga et al. demonstrated that visual hallucinations were observed in 10 of 79 (13%) patients with autopsy-confirmed MSA. The Neuropsychiatric Inventory revealed that 15% of patients with MSA had hallucinations. Since these reports did not include detailed information about the use of dopaminergic medication, it is difficult to speculate on the pathophysiology of the hallucinations. Particularly, the influence of concomitant Lewy body pathology should be considered since Lewy body-spectrum pathology was observed in 22.7% of patients with MSA.

Other atypical symptoms in MSA

Patients with MSA may show vertical gaze palsy, which is a characteristic finding of PSP. Koga et al. reported that 17.6% of patients with autopsy-confirmed MSA presented vertical gaze palsy. However, the detailed clinical characteristics and pathological background of vertical gaze palsy in MSA have not been elucidated, since PSP, particularly PSP with a predominant cerebellar ataxia type, shows a combination of cerebellar ataxia and vertical gaze palsy. Age at onset, early falls, and no dysautonomia can increase the accuracy of the diagnosis of PSP-C.

WHY DO WE NEED NEW DIAGNOSTIC CRITERIA FOR MSA?

Toward the development of disease-modifying therapy or symptomatic interventions for MSA

The etiology and pathogenesis of MSA remains unresolved but understanding of the intrinsic mechanism has steadily improved. Relocation of phosphoprotein-25α (p25α), which is a myelin stabilizing protein, from the myelin sheath to the oligodendroglial cell soma followed by formation of cytoplasmic p25α inclusions has been found to be an early event in patients with MSA. Although the presence of α-syn in oligodendrocytes remains under debate, aberrant α-syn mRNA expression and α-syn prion-like propagation are considered to be closely associated with disease onset and progression. Decreased glial cell line-derived neurotrophic factor expression, autophagy disruption, mitochondrial failure, and neuroinflammation can also play an important role in the pathogenesis of MSA. Although MSA is reportedly a sporadic disease, rarely, patterns of familial aggregation following autosomal dominant and autosomal recessive inheritance have been reported, which supports the hypothesis of a genetic contribution. Tsuji et al. demonstrated that functionally impaired variants of COQ2 were associated with an increased risk of MSA in multiplex families and patients with sporadic disease. A recent meta-analysis reported an association of the COQ2 V393A variant with an increased risk of MSA in patients of East Asian ancestry.

Clinical trials on disease-modifying therapy (DMT) have been conducted. A randomized, double-blind, placebo-controlled trial of autologous mesenchymal stem cells (MSCs) administered through the carotid artery revealed a significantly slower progression of the Unified Multiple System Atrophy Rating Scale (UMSARS) scores compared to placebo. Although there were several limitations including a small number of patients, a single-center design, and increased signal abnormalities on brain MRI indicating small strokes, this study provided encouragement for researchers to conduct additional clinical trials. Consequently, a phase 1 study is underway at the Mayo Clinic in Rochester to evaluate the safety and tolerability of intrathecal injection of autolo-
gous MSCs in a dose-escalation study with patients with MSA (NCT02315027). The results of studies on myeloperoxidase inhibitors for ameliorating microglial activation (NCT02388295) and on two vaccines (PD01A and PD03A) against α-syn (NCT02270489) are also expected.

For the successful development of DMT or symptomatic interventions for MSA, we need early diagnostic criteria and clinically definitive categorization. However, as we discussed in the previous section, early diagnosis of MSA can still be challenging based exclusively on clinical findings because patients with MSA can show isolated parkinsonism, cerebellar ataxia, and autonomic failure, particularly during the early course of the illness, as well as other atypical findings including cognitive impairment and vertical gaze palsy (Figure 4). Thus, it is crucial to establish well-validated biomarkers for a very early and clinically definitive diagnosis.

Diagnostic challenges for isolated autonomic failure phase

According to a United States prospective cohort study on PAF,19 the presence of a probable rapid-eye movement sleep behavior disorder is strongly correlated with the development of DLB, PD, or MSA. Patients whose diagnosis converted from PAF to MSA had a younger age at onset, severe bladder/bowel dysfunction, preserved olfactory function, and a cardiac chronotropic response to tilt more than 10 beats per minute. Patients with PAF that converted to DLB or PD showed impaired olfactory function, a lesser chronotropic response to tilt, and a longer duration of illness. The patients with retained PAF had very low plasma noradrenaline (NA) levels, no rapid eye movement (REM) sleep behavior disorder (RBD), preserved olfactory function, and a slow resting heart rate. The combination of olfactory function and plasma NA levels successfully classified PD/DLB (impaired olfaction, tended to have lower NA levels), PAF (normal olfaction and lower NA levels), and MSA (normal olfaction and tended to have plasma NA levels that were not low). Thus, the combination of age at onset of autonomic failure, bladder/bowel function, olfactory function, RBD, plasma NA levels, and a cardiac chronotropic response to tilt may serve as a good indicator for predicting conversion from PAF to MSA.

Kikuchi et al.26 reported that an olfactory function test can be valuable as a clinical instrument for differentiating PD from MSA in the early stages with high specificity. Low plasma NA levels correspond to postganglionic involvement in PD, DLB, and PAF.57 If available, 123I-metiodobenzylguanidine (MIBG) cardiac scintigraphy may provide additional information for differentiating MSA from PD, DLB, and PAF.56,59 In general, PD, DLB, and PAF show predominantly postganglionic involvement, but MSA shows preganglionic involvement.19 However, to varying degrees, both preganglionic and postganglionic involvements can be observed in MSA.60 Nagayama et al.61 found that there was significant longitudinal decreased cardiac MIBG accumulation in patients with MSA. Patients with MSA may have Lewy body pathology including cardiac sympathetic nervous system involvement.62,63 Thus, we should consider the possibility of a complex pathology and differences in the spread of the autonomic involvement in MSA.

Although PAF is a rare condition, further studies are expected to prove the usefulness of a composite score for these indices, to compare them with the
autopsy results and to develop additional semi- or quantitative biomarkers that will improve the objective efficacy of DMT.

**Diagnostic challenge for an isolated parkinsonism phase**

Conventional MRI (cMRI) is a reliable and widely available test for differentiating MSA from PD, DLB, and PSP. Based on a detailed review of brain MRI as a diagnostic tool for MSA, several characteristic MRI findings have been incorporated in the current diagnostic criteria as supportive of ‘possible MSA.’ However, the utility of these findings has not been investigated for patients with MSA during the early stages, when the clinical diagnosis remains uncertain.

Recently, Mestre et al. reported that 30% of patients with MSA-P showed characteristic MRI signs preceding the clinical diagnosis. T2 posterior putaminal atrophy affected 91.7% of the patients (n = 11/12), T2 slit-like putaminal hyperintensity 50% (n = 6/12), and hypointensity and/or definite atrophy of the posterior putamen on T2 imaging 66.7% (6/9), which were suggested as characteristic abnormal MRI findings for MSA. These results support the view that the diagnostic criteria should be reconsidered based on cMRI findings.

However, these cMRI findings may also be observed not only in patients with PD, PSP, SCA 17, and adult GM1 gangliosidosis but also in control subjects. In addition, putaminal MRI findings may change among different magnetic-field strengths (Figure 5A). A T2 hyperintense putaminal rim without atrophic changes at 3 T may be non-specifically observed in patients with PD or healthy controls but the configuration of PD or controls and MSA can be different. Finally, we should also consider that approximately 60% of patients with MSA-P had normal cMRI findings within 2 years of disease onset.

**123I-MIBG cardiac scintigraphy can also be useful for differentiating PD from other neurodegenerative parkinsonism conditions such as MSA and PSP. However, it is difficult to differentiate MSA from PSP and corticobasal degeneration (CBD) since patients with MSA, PSP, and CBD will show preserved MIBG accumulation.** In addition, there are no data regarding the results of MIBG cardiac scintigraphy findings in patients with early MSA who did not fulfill the diagnostic criteria.

**Diagnostic challenges for isolated cerebellar ataxia phase**

Pontine signal alterations called a hot cross bun (HCB) sign and T2 hyperintensity in the bilateral middle cerebellar peduncle (MCP) are characteristic MRI findings in patients with MSA (Figure 5C). Interestingly, SCA 2, SCA 3, SCA 7, and SCA 8 may also show an HCB sign. T2 hyperintensity in the MCP is also observed in patients with fragile X associated tremor/ataxia syndrome, Wilson’s disease, liver cirrhosis, multiple sclerosis, stroke and other...
Kim et al. demonstrated that SCA 1, 2, 3, 6, 17, or dentatorubral-pallidoluysian atrophy was confirmed in 22 of 302 (7.3%) of clinically diagnosed MSA cases. Thus, we should consider both MSA and SCAs when patients show a combination of an HCB sign and T2 hyperintensity in the MCP.

However, previous studies have not given due consideration to disease duration from onset to MRI. In general, the progression rate is significantly rapid in MSA compared to SCAs. Importantly, more than 60% of patients with MSA-C had an HCB sign within 2 years from the onset of ataxia. In contrast, patients with SCAs with an HCB sign showed significantly longer disease duration compared to MSA-C, although available data are limited. Careful neurological evaluation to exclude nonneurodegenerative diseases will be necessary, but an early HCB sign may strongly support the diagnosis of MSA.

**DEVELOPMENT OF NOVEL DIAGNOSTIC BIOMARKERS**

Currently, conventional MRI, olfactory test, 123I-MIBG cardiac scintigraphy, plasma NA level, cardiac chronotropic response upon tilt, REM sleep behavior disorder, and genetic tests and so on are available for the differential diagnosis of MSA from other diseases in clinical practice. However, there is no established test for the early and clinically definitive diagnosis that can satisfy the criteria of objectivity, semi- or quantitative measurement, and uncomplicated and worldwide availability.

**MRI**

Diffusion-weighted MRI (DWI) measures the random Brownian motion of water molecules within a voxel. Patients with MSA-P show increased putaminal diffusivity compared to those with PD. A meta-analysis on putaminal diffusivity showed a sensitivity of 90% [95% confidence interval (CI): 76.7–95.8%] and specificity of 93% (95% CI: 80.0–97.7%) for discriminating MSA-P from PD. More recently, Péran et al. demonstrated that multimodal MRI, including DWI, is able to discriminate patients with PD from those with MSA with high accuracy. Further prospective multicenter studies with standardized regions of interest and imaging protocols across institutions or different MRI equipment vendors of DWI will be needed, particularly of cases in the early disease stages prior to fulfilling the diagnostic criteria.

Voxel-based morphometry analysis is a whole-brain, unbiased, objective technique. Recent advances in imaging algorithms have enabled automatic compartmentalization of the brain into multiple parcellations and provide for quantification of gray and white matter volumes in these regions on an individual basis. Scherfler et al. demonstrated that the diagnostic accuracy for PD vs. MSA-P or PSP was 97.4% using the midbrain and putaminal volumes as well as cerebellar gray matter volume calculated by automated and observer-independent volumetric MRI analysis, although the diagnostic accuracy based on validated clinical consensus criteria was 62.9%. Compared to DWI, multicenter studies have widely used volumetry. Individual volumetric T1-weighted MRI can be a good candidate for a novel diagnostic marker for novel diagnostic criteria for MSA. We should confirm the usefulness of this procedure for the diagnosis of patients with isolated autonomic failure and in the cerebellar ataxia phase.

**Visualization of α-syn**

Biopsy and PET are two modalities used to detect α-syn accumulation in MSA. Doppler et al. demonstrated that phospho-α-syn (p-α-syn) in dermal nerve fibers was found in 67% of patients with MSA and PD, but not in those with tauopathy or controls when analyzing 15 consecutive sections. Analyzing serial sections increased the sensitivity to 75% and 73%, respectively. Interestingly, p-α-syn clustered in autonomic fibers in PD but mainly in unmyelinated somatosensory fibers in MSA.

BF-227 can bind aggregated amyloid beta protein and p-α-syn. The PET data of BF 227 demonstrated high distribution volumes in the subcortical white matter, putamen and posterior cingulate cortex, globus pallidus, primary motor cortex and anterior cingulate cortex, and substantia nigra in MSA in accordance with GCI-rich brain areas compared to the normal controls. A pathological study showed that BF-227 histofluorescence was observed in most of the GCIs. However, Verdurand et al. reported that 18F-BF-227 did not bind to GCIs of MSA brain tissue using state-of-the-art autoradiography.

More recently, Koga et al. demonstrated that the tau tracer 11C-PBB3 showed significant autoradiographic binding to the striatopallidal fibers in two...
MSA cases with high densities of GCIs. One MSA case showed increased binding of PBB3 in the frontal lobe, globus pallidus, midbrain, parietal lobe, putamen, temporal lobe, substantia nigra, thalamus, and ventral striatum.99

Although there are several candidate PET radiotracers for imaging aggregated α-syn, they are not easy to develop. The amount of insoluble α-syn protein in the MSA brain is 10-fold or lower than the amount of Aβ in the Alzheimer’s disease brain. The tracer must readily pass through the glial cell membrane to access the binding site.90 There is still no known radioligand that selectively binds to α-syn with high affinity.

Figure 6. Current and future diagnostic biomarkers of MSA. Hitherto, conventional MRI, olfactory test, 123I-metaiodobenzylguanidine (MIBG) cardiac scintigraphy, plasma NA level, cardiac chronotropic response upon tilt, REM sleep behavior disorder, and genetic tests are available for differential diagnosis of MSA from other diseases based on their clinical condition. Conventional MRI may be useful for differentiating MSA from other mimic diseases. Olfactory function tests and autonomic function tests can provide important information for differentiating Lewy body disease from MSA, PSP, and CBD. REM sleep disorders are strongly linked to synucleinopathies that involve the CNS. Genetic tests are useful for differentiating MSA from other hereditary disorders. However, there are no known biomarkers that fulfill the criteria of objectivity, semi- or quantitative, and uncomplicated and worldwide availability. Currently, several biomarkers, such as DWI, automated and observer-independent volumetric MRI, skin biopsy, α-syn PET imaging, and blood and CSF biomarkers that have the potential for overcoming such limitations, are under development. MSA: multiple system atrophy, MRI: magnetic resonance imaging, REM, rapid eye movement, CNS: central nervous system, NA: noradrenaline, PSP: progressive supranuclear palsy, CBD: corticobasal degeneration, PD: Parkinson’s disease, DLB: dementia with Lewy bodies, PAF: pure autonomic failure, DWI: diffusion-weighted imaging, PET: positron emission tomography, CSF: cerebrospinal fluid, α-syn: α-synuclein, SCD: spinocerebellar degeneration, FXTAS: Fragile X-associated tremor/ataxia syndrome.

Other candidate biomarkers

MSA, particularly MSA-P, showed significantly decreased peripapillary retinal nerve fiber layer thickness in the inferior and inferotemporal sectors and significant perifoveal thinning in the superior outer sector compared to healthy controls. Retinal thinning correlated with the UMSARS score and the Geriatric Depression Scale.91

A number of studies have searched for candidate biomarkers using blood and cerebrospinal fluid, including neurofilament light chain, catecholamine metabolite, coenzyme Q10 micro RNA, and disease-related proteins such as total α-syn, DJ-1, amyloid beta and total tau, but there are still no reliable biomarkers for the diagnosis of MSA.92–99

Figure 6 summarizes the current and future diagnostic biomarkers of MSA.

CONCLUSIONS

The several trials assessing potential DMT failed to reach their primary efficacy endpoint, although preclinical disease-model studies have shown positive findings.100–103 As we discussed in this review, probable MSA can only be diagnosed at a substantially advanced stage. Early diagnostic criteria and clinically definitive categorization are needed for the success of DMT or symptomatic interventions for MSA. To develop early diagnostic criteria, prospective image-omics cohort studies focusing on patients with MSA who show isolated autonomic failure or motor involvement will be necessary. Since the MSA phenotype could differ between western and eastern countries (MSA-C vs. MSA-P, proportion of the phenotype could differ between western and eastern countries),104,105 international collaborative studies will be helpful. To establish clinically definitive criteria, the development of a disease-specific marker is urgently needed because patients with MSA may show atypical symptoms during the course of the illness. PD, PSP, DLB, PAF, and other genetic, and immunologic conditions can mimic MSA.106 Molecular PET imaging and biomarkers for detecting α-syn are promising, but we need more specific radioligands and multicenter results. On the other hand, individually automated and observer-independent volumetric MRI analysis seem to be attractive because it will satisfy the requirements for objectivity, semi- or quantitative measurements, and uncomplicated and worldwide availability for
universal use. Further prospective multicenter studies will be necessary to incorporate this technique into future diagnostic criteria. Finally, the combination of multimodal markers, including functional and molecular imaging, as well as cerebrospinal fluid and plasma biomarkers, will be helpful for both establishing early diagnostic criteria and for clinically definitive categorization.

Conflicts of Interest
The authors have no financial conflicts of interest.

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