The Differential Diagnosis and Treatment of Atypical Parkinsonism

Johannes Levin, Alexander Kurz, Thomas Arzberger, Armin Giese, Günter U. Höglinger

SUMMARY

Background: Aside from idiopathic Parkinson syndrome (Parkinson’s disease), there are a number of other, so-called atypical parkinsonian syndromes: dementia with Lewy bodies (DLB), multiple system atrophy (MSA), progressive supranuclear palsy (PSP), and corticobasal degeneration (CBD). DLB is a common disease, with a prevalence of 0.4% (400 cases per 100 000 persons) in the elderly; MSA and PSP both have a prevalence of 5 to 10 per 100 000 persons, while the prevalence of CBD is about 1 per 100 000.

Methods: This review is based on pertinent publications retrieved by a selective literature search.

Results: The atypical parkinsonian syndromes are synucleinopathies and tauopathies, i.e., disorders characterized by the abnormal deposition of the proteins α-synuclein and tau. The site of deposition is correlated with the clinical features. In DLB, synuclein is mainly deposited in neocortical neurons, with some brain stem involvement as well. The main clinical features are dementia and, later on, Parkinsonism. In MSA, synuclein is deposited in oligodendrocytes, mainly in the cerebellum but also in the brain stem; the main clinical feature is autonomic dysfunction combined with parkinsonism or cerebellar ataxia. Synucleinopathies often impair REM (rapid eye movement) sleep, PSP and CBD, on the other hand, are primary tauopathies. PSP usually causes predominantly supranuclear vertical gaze palsy and early postural instability with falls, less commonly Parkinsonism (PSP-P) or frontotemporal dementia (PSP-FTD) as its most prominent feature. CBD typically manifests itself as markedly asymmetrical parkinsonism with apraxia or cortical sensory disturbance. At present, there is no accepted causal treatment for any of these disorders; the available symptomatic treatments are of limited efficacy and are supported only by low-level evidence.

Conclusion: Causal treatments for neurodegenerative diseases are now being developed and tested, and thus a molecular diagnosis is desirable. This will require the cooperation of primary care physicians with specialized centers.

Cite this as:
CBD. The different disease entities typically affect characteristic regions in the brain. The misfolding and aggregation of these proteins can lead to degeneration of the affected cell populations on the one hand, but on the other hand they can also facilitate their spread into anatomically connected regions of the brain and thereby enable disease progression (4). The increasingly better understanding of the pathophysiology provides new approaches to causally oriented treatment, in order to stop the progression of these currently incurable disorders (4).

Clinical symptoms
Firstly, we will describe the two parkinsonian syndromes characterized by synucleinopathy and then the two tauopathies, in descending order of prevalence.

Dementia with Lewy bodies
Epidemiology: DLB is the second most common form of progressive cognitive decline in old age due to neurodegeneration after dementia in Alzheimer’s disease. Its prevalence in the population older than 65 years is approximately 0.4%. Symptom onset usually occurs between the 50th and 80th year of life. Men are affected slightly more often than women (5, 6).

Clinical symptoms: The initial symptoms consist of a reduction in cognitive performance with fluctuating episodes of poor and better cognitive performance (Table 1). Deficits involve the naming of objects, verbal fluency, visuospatial abilities (as shown by the clock-drawing test), and executive functions. Executive problems become apparent in tasks requiring planning, shifting between strategies, organizing, and self-monitoring. In contrast to Alzheimer’s dementia, memory impairment is not a common or predominant feature in DLB at the early stage of the disease (7).

On the other hand, vivid, recurring or persistent visual hallucinations are frequent, as are delusions, lack of initiative and motivation, depressed moods, and anxiety (8). Akinetic-rigid movement disorder may already be present at the time of diagnosis, but develops over the course of the illness in the majority of cases.

### Table 1

<table>
<thead>
<tr>
<th>Areas</th>
<th>Findings/recommendations</th>
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<tbody>
<tr>
<td>Neuropathology</td>
<td>Aggregates of α-synuclein in neuronal somata (Lewy bodies) and processes (Lewy neurites) in typical distribution (neocortex and brain stem)</td>
</tr>
<tr>
<td>Clinical syndrome</td>
<td>Demential cognitive impairments (attention, executive functions, visuospatial skills) with relevance to activities of daily living</td>
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<td>At least one of the following key characteristics:</td>
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<td>– Parkinsonian syndrome, onset after or at most 12 months before onset of dementia</td>
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<td>– Fluctuations in alertness and attention</td>
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<td></td>
<td>– Repeated visual hallucinations</td>
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<tr>
<td>MRI results</td>
<td>Normal medial temporal lobes, in contrast to Alzheimer’s dementia</td>
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<td></td>
<td>Rule out symptomatic causes, for example, vascular encephalopathy, normal pressure hydrocephalus, aqueductal stenosis, encephalitis, tumor</td>
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<tr>
<td>Basic diagnostic evaluation</td>
<td>Neuropsychological tests, early visuospatial impairment (clock-drawing test, figure copying test), impaired speech (naming, verbal fluency), executive dysfunction, recommended Montreal Cognitive Assessment (MoCA), Frontal Assessment Battery (FAB); memory impairments may occur at a later stage</td>
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<tr>
<td>Extended diagnostic evaluation</td>
<td>$^{127}$I-FP-CIT-SPECT/$^{18}$F-Dopa-PET: striatal dopaminergic denervation</td>
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<td></td>
<td>$^{99}$Tc-HMPAO-SPECT/$^{18}$FDG-PET: occipital hypometabolism</td>
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<tr>
<td></td>
<td>$^{123}$I-MIBG-SPECT: cardiac noradrenergic denervation may occur</td>
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<td>Sleep laboratory: where applicable, rapid eye movement (REM) sleep behavior disorder</td>
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<td>Symptomatic treatment</td>
<td>Cholinesterase inhibitors, for example, donepezil (1 × 5–10 mg) or rivastigmine (2–3 × 6–12 mg), for dementia (evidence level 1+, recommendation grade A)</td>
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<td>Memantine (1 × 5–20 mg) for behavioral symptoms (evidence level 1+, recommendation grade A)</td>
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<td></td>
<td>Levodopa (3–4 × 100–200 mg) for akinetic-rigid symptoms (evidence level 2+, recommendation grade B)</td>
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<td>Atypical neuroleptics, e.g., clozapine (6.25–100 mg/day, 2/3 of this at nighttime) or quetiapine (off-label, 25–100 mg at nighttime), for psychosis (evidence level 4, recommendation grade 0)</td>
</tr>
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Evidence level 1 = randomized controlled study (studies); 2 = case-control study (studies); 3 = non-analytic case study (studies); 4 = expert opinion; recommendation grade A = strong recommendation [should]; B = recommendation; 0 = open recommendation [can]

MRI, magnetic resonance imaging; SPECT, single photon emission computed tomography; PET, positron emission tomography
Diagnosis: Typical clinical features include progressive dementia, fluctuating cognition, vivid visual hallucinations, and a parkinsonian syndrome (5).

Magnetic resonance imaging (MRI) may reveal atrophy of the caudate nucleus, putamen, and thalamus. In contrast to Alzheimer’s dementia, little atrophy is seen in the cortex, while the medial temporal lobe in particular is not atrophied (eFigure) (9).

In specialized centers, functional brain imaging techniques can be used to improve diagnostic accuracy. These technologies include $^{123}$I-FP-CIT single-photon emission computed tomography (SPECT) for dopamine transporter visualization (10), positron emission tomography (PET) with $^{18}$F-FDG as a tracer for the assessment of regional cerebral glucose metabolism (11), and PET in

### Table 2

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<thead>
<tr>
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<tr>
<td><strong>Neuropathology</strong></td>
<td>Aggregates of α-synuclein in cytoplasm of oligodendrocytes (glial cytoplasmic inclusions) and neurons in a typical distribution (cerebellum, pons, and basal ganglia)</td>
</tr>
</tbody>
</table>
| **Clinical syndrome**      | Dysautonomia: urinary incontinence, erectile dysfunction, orthostatic hypotension At least one of the following syndromes:  
– Parkinsonian syndrome (MSA-P, about 60% of cases, PPV about 90%): bradykinesia with rigidity, tremor, or postural instability  
– Cerebellar syndrome (MSA-C, about 40% of cases, PPV about 90%): gait ataxia with cerebellar dysarthria, limb ataxia, or cerebellar oculomotor dysfunction |
| **MRI results**            | Atrophy of putamen, middle cerebellar peduncle, pons, or cerebellum  
Cross shaped hyperintensity in the pons (T2 axial)  
Hypointense putamen with hyperintense rim (T2 axial) |
| **Basic diagnostic evaluation** | Levodopa test: poor response to levodopa  
Schellong test: orthostatic drop in blood pressure after 3 minutes’ standing up from supine position (≥ 20 mm Hg systolic or ≥ 10 mm Hg diastolic)  
Urological examination: exclusion of other causes of erectile dysfunction and urinary incontinence |
| **Extended diagnostic evaluation** | $^{123}$I-FP-CIT-SPECT/$^{18}$F-Dopa-PET: symmetric striatal dopaminergic denervation  
$^{99m}$Tc-HMPAO-SPECT/$^{18}$FDG-PET: hypometabolism in putamen, brainstem, or cerebellum  
$^{123}$I-IBZM-SPECT/$^{18}$FDMPF-PET: symmetric postsynaptic, striatal degeneration  
$^{123}$I-MIBG-SPECT: intact cardiac noradrenergic innervation (DD PD with dysautonomia)  
Oculomotor function: cerebellar pattern (saccadic pursuit movement, gaze-evoked or downbeat nystagmus)  
Sleep laboratory: rapid eye movement (REM) sleep behavior disorder |
| **Symptomatic treatment**  | Levodopa (3–4 × 100–200 mg): mild to moderate improvement of akinetic-rigid symptoms (evidence level 2–, recommendation grade B)  
Aramantidin (3 × 100–200 mg): mild positive effect on akinetic-rigid symptoms (evidence level 1–, recommendation grade 0)  
Botulinum toxin A: positive effect on focal dystonia (evidence level 3, recommendation grade 0)  
Droxidopa (2 × 200–600 mg) (not approved in Germany for this indication): positive effect on orthostatic hypotension (evidence level 1, recommendation grade A)  
Ephedrine (3 × 15–45 mg): positive effect on orthostatic hypotension (evidence level 3, recommendation grade 0)  
Midodrine (3 × 5–10 mg): positive effect on orthostatic hypotension (evidence level 3, recommendation grade 0)  
Fludrocortisone (1–3 × 0,1 mg): positive effect on orthostatic hypotension (evidence level 3, recommendation grade 0)  
Octreotide (25–50 mg s. c. 30 minutes before meals): positive effect on postprandial hypotension (evidence level 2–, recommendation grade 0)  
Desmopressin (10–40 µg spray, 100–400 µg tablets): positive effect on nocturnal polyuria (evidence level 3, recommendation grade 0)  
Oxybutinin (2–3 × 2.5–5 mg): for urge incontinence (evidence level 4, recommendation grade 0) |

Evidence level 1 = randomized controlled study (studies); 2 = case-control study (studies); 3 = non-analytic case study (studies); 4 = expert opinion; recommendation grade A = strong recommendation [should]; B = recommendation; 0 = open recommendation [can]  
MSA, multiple system atrophy; MRI, magnetic resonance imaging; PPV, positive predictive value; DD, differential diagnosis; PD, Parkinson’s disease; s. c., subcutaneous; SPECT, single photon emission computed tomography; PET, positron emission tomography
combination with other appropriate tracers for demonstrating beta-amyloid deposition (12). In addition to a dopaminergic deficit, patients show a characteristic pattern of hypometabolic areas, especially involving the occipital regions of the primary visual cortex and the visual association cortex. In contrast, the medial temporal lobe and the posterior cingulum are not affected. Deposits of beta-amyloid may be present, but are less pronounced than in Alzheimer’s dementia (10–12).
motor disturbance occurs before or concomitantly with cognitive impairment, the condition is usually classified as Parkinson’s disease with dementia (5).

Therapy: The treatment of DLB usually needs to address all three domains of the syndrome: cognitive deficits, neuropsychiatric symptoms, and movement disorder (13). A particular problem is posed by the fact that medication with levodopa can improve the parkinsonian symptoms but may worsen the neuropsychiatric symptoms. Dopamine agonists should not be used for treating movement disorder in DLB. Moreover, patients with DLB are particularly sensitive to the adverse effects of antipsychotic drugs.

Table 1 shows the main options for symptomatic treatment. The evidence levels reported for the individual therapies refer to the 2008 guidelines of the Scottish Intercollegiate Guidelines Network (SIGN). The recommendation grades are

Differential diagnosis: The major differential diagnoses include Alzheimer’s dementia and Parkinson’s disease with dementia. DLB differs from Alzheimer’s disease in respect to the following features: in DLB, memory impairments are relatively mild and occur at a later stage (Mini Mental State Examination); patients exhibit prominent visuo-spatial deficits (Clock Drawing Test), fluctuating cognitive performance; early parkinsonism, and mild atrophy of the medial temporal lobe including the hippocampus. They may also show rapid eye movement (REM) sleep behavior disorder (eFigure). DLB and PD with dementia form a spectrum lacking clear clinical or neuropathological boundaries. To distinguish between the two entities, the temporal sequence of symptom onset is usually employed. If motor disturbance occurs before or concomitantly with cognitive impairment, the condition is usually classified as Parkinson’s disease with dementia (5).

Therapy: The treatment of DLB usually needs to address all three domains of the syndrome: cognitive deficits, neuropsychiatric symptoms, and movement disorder (13). A particular problem is posed by the fact that medication with levodopa can improve the parkinsonian symptoms but may worsen the neuropsychiatric symptoms. Dopamine agonists should not be used for treating movement disorder in DLB. Moreover, patients with DLB are particularly sensitive to the adverse effects of antipsychotic drugs. Table 1 shows the main options for symptomatic treatment. The evidence levels reported for the individual therapies refer to the 2008 guidelines of the Scottish Intercollegiate Guidelines Network (SIGN). The recommendation grades are
reproduced in accordance with the Association of the Scientific Medical Societies in Germany (AWMF) 2012. These definitions are also used for the other atypical parkinsonian syndromes described in the following chapters.

Multiple system atrophy

**Epidemiology:** Multiple system atrophy (MSA) has a prevalence of about 5/100 000. The mean age at disease onset is in the sixth decade of life, and the sex distribution is even. After an average of 6 to 10 years, patients die mostly from aspiration or nocturnal cardiorespiratory arrest (14).

**Clinical symptoms:** MSA has two different subtypes, either with predominant parkinsonism (MSA-P) (in Europe, this applies to 60% of cases) or with predominant cerebellar symptoms (MSA-C, 40%). In addition to motor impairment, vegetative symptoms are an obligatory component of MSA—especially urinary incontinence, erectile dysfunction, or orthostatic hypotension (Table 2). Isolated pure autonomic failure (PAF) may precede the motor impairments in MSA.

A hypokinetic-rigid parkinsonian syndrome is dominant in MSA-P, but as a rule its presentation tends to be more symmetrical and less responsive to levodopa than in PD. The classical, slow, pill-rolling resting tremor of PD is rare in MSA. What is mostly found is an irregular, higher-frequency postural tremor.

In MSA-C, the most common symptom is gait ataxia with wide-based movements. Furthermore, ataxia of the limbs is common, as are cerebellar oculomotor impairments, scanning dysarthria, and intention tremor.

Some further symptoms are typical of MSA but do not affect every patient and can occur in both predomi- nance types (15). Patients in advanced stages of the disease often have dysarthria, inspiritional stridor, and dysphagia, for example. Extrapyramidal signs are seen in 30–50% of patients with MSA, but not in PD patients. Focal dystonias and posture abnormalities (antecollis, Pisa syndrome, contractures of the hands/feet) are also common (14). Many patients with MSA have a comorbid diagnosis of REM sleep behavior disorder (14).

**Diagnosis:** The diagnostic criteria of MSA were revised in 2008 (16). The diagnosis requires at least one symptom of vegetative dysfunction (orthostatic hypotension, urinary incontinence, erectile dysfunction) after the exclusion of symptomatic causes. Furthermore,
sporadic progressive parkinsonian symptoms (MSA-P) or ataxia (MSA-C) and at least one further typical symptom or a characteristic imaging finding are needed. Atrophy in the putamen, the middle cerebellar peduncle, the pons, and the cerebellum are visible on MRI (Figure 2), and hypometabolism in the putamen, brain stem, or cerebellum on FDG-PET scans. Not included in the diagnostic criteria but still indicative of MSA are signaling anomalies in T2-weighted MRI scans with 1.5 Tesla (eFigure): a cross-shaped hypointensity in the pons (“hot cross bun” sign) and a hypointense putamen with a hyperintense rim (putaminal rim sign) (17).

**Therapy:** In about one-third of patients, treatment with levodopa will improve the hypokinetic-rigid symptoms in MSA-P (18). To date, no effective treatment exists for ataxia. Symptomatic treatment of the vegetative symptoms is recommended (13) (Table 2), as these notably impair patients’ quality of life (19). Furthermore, speech therapeutic, occupational therapeutic, and physiotherapeutic measures are indicated in order to maintain patients’ independence.

**Progressive supranuclear palsy**

**Epidemiology:** Progressive supranuclear palsy has a prevalence of 5–10/100 000 persons. The mean age at onset is 65 years. Patients die after an average of 8 years, often as a result of aspiration. The sex ratio is for the most part evenly balanced (20).

**Clinical symptoms:** The clinical manifestations of PSP are diverse (Table 3). The most common of these (ca 40%) is Richardson’s syndrome, with levodopa-resistant akinetic-rigid symptoms affecting the axial muscles, a tendency to fall backwards that occurs early in the disease course, and vertical gaze paresis (21). Patients are unable to move their eyes upwards or downwards at will, but vertical eye movements can still be triggered by the vestibulo-ocular reflex (passive head movement applied by the investigator). A frontal lobe syndrome is also typical, with apathy, impaired executive functions, and a positive “applause sign” (the inability to stop clapping after being asked to imitate the examiner’s three handclaps). Additionally, the Luria test yields a pathological result: patients are unable to correctly perform the sequence of hand–fist–palm at least six times in succession. Verbal fluency is reduced to <9 words beginning with S in 60 seconds. Spastic (pseudobulbar) speech and swallowing impairment is also characteristic. Often PSP is initially characterized by asymmetric levodopa-responsive parkinsonism (PSP-P, about 20%), which is clinically indistinguishable from Parkinson’s disease, until the later onset of vertical gaze paresis. A rare syndrome with sudden and transient gait freezing without accompanying rigidity or tremor is known as pure akinesia with gait freezing (PAGF, <5%) and is considered typical of an underlying PSP neuropathology. Sometimes the onset of the disease may be dominated by clinical symptoms in accordance with the behavioral variant of frontotemporal dementia (bvFTD, 15%) or progressive non-fluent aphasia (PNFA, 5%), which can be distinguished from other forms of frontotemporal dementia only once oculomotor problems appear. Similarly, PSP pathology can clinically trigger a corticobasal syndrome (CBS, 10%), which is described in the chapter on CBD (21).

**Diagnosis:** The current criteria of the National Institute of Neurological Disorders and Stroke (NINDS) and the Society for Progressive Supranuclear Palsy (SPSP) for the diagnosis of PSP (22) are based on the clinical confirmation of the tendency to falls and oculomotor impairments. They are thus very sensitive for Richardson’s syndrome, but less suitable for detecting the other clinical courses that PSP takes (21). MRI will reveal midbrain and frontal lobe atrophy (Figure 2) (23). The parkinsonian symptoms generally improve only slightly in the levodopa test. In case of doubt, nuclear medicine methods can be used to confirm symmetric presynaptic nigrostriatal dopaminergic denervation (for example, FP-CIT-SPECT), postsynaptic striatal degeneration (for example, IBZM-SPECT), or hypometabolism in the frontal and midbrain regions (FDG-PET) to ascertain the diagnosis.
Therapy: The treatment of PSP is currently exclusively symptomatic (Table 3) (13, 20). Therapeutic target symptoms include akinetic-rigid symptoms, oculomotor impairments, neuropsychological deficits, and the occasionally occurring dystonia and sleep impairment. The fundamentals of pharmacological treatment are summarized in Table 3. Furthermore, speech therapy and physiotherapy are indicated to prevent aspiration and falls.

Corticobasal degeneration
The term corticobasal degeneration (CBD) is reserved for the neuropathological diagnosis. Clinically, CBD mostly manifests with a corticobasal syndrome (CBS) (24). Patients with CBD pathology may, however, also have other clinical syndromes than CBS (24) (Table 4). Similarly, patients with clinical CBS may in terms of neuropathology also have underlying PSP, Alzheimer’s disease, or other pathologies (25). The clinicopathological assignment is therefore possible to a very limited degree only, which underlines the necessity of a strict distinction of the neuropathological diagnosis of CBD from the clinical diagnosis of CBS.

Epidemiology: The prevalence of CBD is about 1/100 000. It usually affects persons in their 6th and 7th decades of life and affects both sexes in equal measure. Patients die after an average of eight years with the disease, mostly from aspiration due to dysphagia (26).

Clinical symptoms: CBS is a progressive constellation of symptoms that comprise functional damage to the basal ganglia and the cerebral cortex and mostly appear asymmetrically (27).

The basal ganglia symptoms include akinesia, rigidity, dystonia, and myoclonus. The hypokinetic-rigid parkinsonian syndrome is typically resistant to levodopa. Tremor may be present, which is superimposed on myoclonus and therefore appear irregular and jerky. As the disease progresses, many patients develop a characteristic, fixed dystonia-related posture of the affected arm, which usually consists of flexion of the hand and forearm and adduction of the upper arm at the shoulder joint (28). During the disease course, most patients develop rigid immobility in just a few years and require care.

The following cortical symptoms can develop:

- Apraxia (difficulty in executing intentional movements despite intact upper and lower motor neuron function)
- Cortical sensory loss (although primary sensation is intact, topographic assignment is impaired in two-point discrimination, graphesthesia, stereognosis)
- Alien limb phenomenon (experiencing one’s limb as not belonging to one’s body)
- Pyramidal signs
- Behavioral disorders
- Aphasia (24).

Diagnosis: The current consensus criteria (24) describe different clinical phenotypes that may be associated with CBD pathology (Table 4). The corticobasal syndrome is defined as described above. Imaging techniques can be used to demonstrate asymmetric parietal atrophy (eFigure) and hypometabolism, as well as presynaptic and postsynaptic nigrostriatal degeneration.

Therapy: Symptomatic treatment (13) (Table 4) of the akinetic-rigid syndrome with levodopa yields a slight transient improvement in about 30% of patients with CBD. Myoclonus responds to clonazepam, among others. In focal dystonia, the local injection of botulinum toxin may be useful (28). Anticholinergic drugs are useful for tremors. Physiotherapy, occupational therapy, and speech therapy should also be prescribed.

Outlook
For a long time, attempts have been made in atypical parkinsonism to diagnose molecular-neuropathologically defined disease entities on the basis of neurological-psychiatric symptom constellations and MRI findings. The clinical syndromes and patterns of regional cerebral atrophy (Figure 2) and functional impairments confirmed by nuclear medical techniques correlate well with each other (eFigure). As it turns out, however, different molecular pathologies can trigger overlapping syndromes and imaging results (Figure 1). CBS, for example, may be due to underlying CBD, PSP, or other pathologies (24). Conversely, PSP pathology may manifest with a multitude of syndromes (21). For this reason, symptoms often do not allow satisfactory attribution to one particular disease entity. Since the established therapies aim for symptom alleviation by means of neurotransmitter modulation, a syndromal classification has been sufficient for routine clinical practice. However, treatment approaches to date have remained unsatisfactory in terms of symptom control and do not delay the progress of the disease. Luckily, spectacular insights into the disease mechanisms have been achieved in recent years, so that causal therapeutic approaches can be studied clinically. These include modulators of pathological protein misfolding, such as the substance Anle138b and antibodies to alpha-synuclein or tau (29, 30). Interventions against defined molecular target structures do, however, require a diagnosis based on molecular pathology, to confirm the presence of the molecular target structures for causal therapy in the individual patient. For this reason, we are currently faced with the challenge of defining and diagnosing neurodegenerative diseases on the basis of their molecular pathologies, not on the basis of clinical syndromes as entities. On this background it is to be welcomed that new diagnostic tools have been developed—for example, tau-PET (eFigure). In this way, molecular processes that could be detected only by neuropathological examination so far, can be put to diagnostic use—for example, deposits of the tau protein. We can therefore be optimistic that relevant innovations will be achieved in the coming years in dealing with these rapidly progressive, fatal diseases.
KEY MESSAGES

- In distinction from Parkinson’s disease, atypical parkinsonism occurs as four diseases in particular: dementia with Lewy bodies (DBL), multiple system atrophy (MSA), progressive supranuclear palsy (PSP), and corticobasal degeneration (CBD).

- The current treatment of the atypical parkinsonian syndromes is symptomatic.

- A detailed and regular clinical examination is important in order to identify problems requiring treatment in good time.

- The syndrome- and imaging-based diagnostic evaluation of the atypical parkinsonian syndromes in specialized centers allows good, but not perfect, classification of the neuropathologically defined disease entities.

- Promising molecular-based diagnostic methods, such as tau position emission tomography (PET), and therapeutic options, such as tau/synuclein antibodies are currently under clinical development.

Conflict of interest statement

Dr Levin has received study support (third party funds) from Parkinson Fonds Deutschland gGmbH (a private organization that provides grants for research into Parkinson’s disease) and lecture fees from Bayer Healthcare and MSD. Prof Giese holds stocks and shares in and has personal links with MODAG. He also holds shares in the patent “EP2367381” and diverse national secondary patents.

Prof Högl is a board member for Asceneuron, Bristol-Myers Squibb, Roche, and UCB Pharma. He has received lecture fees from Abbvie, Bristol-Myers Squibb, Elan, Novocea, Roche, Sellas, Teva, and UCB Pharma.

The remaining authors declare that no conflict of interests exists.

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**eFigure:** Magnetic resonance imaging shows in the midsagittal plane no specific atrophy pattern for Parkinson’s disease (PD) (A), whereas in progressive supranuclear palsy (PSP) (B), midbrain atrophy (*) is visible, and for multiple system atrophy (MSA) (C) the image shows pontine atrophy (*) and atrophy in the upper portion of the cerebellar vermis (#). Furthermore, signal anomalies in T2 weighted MRI with 1.5 Tesla are suggestive of, but not highly sensitive for, MSA—especially a cross-shaped hypointensity in the pons (hot cross bun sign, *, D) and a hypointense putamen (*) with hyperintense putaminal rim (putaminal rim sign, arrow, E). In corticobasal syndrome (CBS) (F), what is typically seen is asymmetric parietal atrophy with atrophy of the gyri and accentuation of the sulci (arrows). In the coronal plane, atrophy of the hippocampus (*) and the temporal lobe (#) with dilatation of the temporal horns of the lateral ventricles (+) are typically seen in Alzheimer’s dementia (AD). However, findings are normal in dementia with Lewy bodies (DLB) (H).

Molecular diagnosis of these disorders is the objective of current research, such as has been shown in the example of a patient with PSP by confirming tau deposition in the basal ganglia and the brainstem on positron emission tomography (PET) using the radiopharmaceutical 18-F-T807, which binds to tau (I, Prof Drzeńga, Clinic and Outpatient Clinic of Nuclear Medicine, University Hospital Cologne)