Diagnostic Accuracy of Combined FP-CIT, IBZM, and MIBG Scintigraphy in the Differential Diagnosis of Degenerative Parkinsonism: A Multidimensional Statistical Approach

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In vivo molecular imaging of pre- and postsynaptic nigrostriatal neuronal degeneration and sympathetic cardiac innervation with SPECT is used to distinguish idiopathic Parkinson disease (PD) from atypical parkinsonian disorder (APD). However, the diagnostic accuracy of these imaging approaches as standalone procedures is often unsatisfying. The aim of this study was therefore to evaluate to which extent diagnostic accuracy can be increased by their combined use together with a multidimensional statistical algorithm. Methods: The SPECT radiotracers 123I-(S)-2-hydroxy-3-iodo-6-methoxy-N-[1-ethyl-2-pyrrodinyl]-methyl]benzamide (IBZM), 123I-N-α-fluoropropyl-2β-carbomethoxy-3β-(4-iodophenyl)nortropan (FP-CIT), and meta-123I-iodobenzylguanidine (MIBG) were used to assess striatal postsynaptic D2 receptor binding, striatal presynaptic dopamine transporter binding, and myocardial adrenergic innervation, respectively. Thirty-one PD and 17 APD patients were prospectively investigated. PD and APD diagnoses were established using consensus criteria and reevaluated after 37.4 ± 12.4 and 26 ± 11.6 mo in PD and APD, respectively. Test accuracy (TA) for PD–APD differentiation was computed for all logical (Boolean) combinations of imaging modalities by receiver-operating-characteristic analysis—that is, after multidimensional optimization of cutoff values. Results: Analysis showed moderate TA for PD–APD differentiation using each molecular approach alone (IBZM, 79%; MIBG, 73%; and FP-CIT, 73%). For combined use, the highest TA resulted under the multidimensional optimization of cutoff values: 1.46 or less for IBZM, less than 2.10 for FP-CIT, and greater than 1.43 for MIBG. This algorithm distinguished APD from PD with a sensitivity of 94%, specificity of 94% (TA, 94%), positive predictive value of 89%, and negative predictive value of 97%. Conclusion: Results suggest that the multidimensional combination of FP-CIT, IBZM, and MIBG scintigraphy is likely to significantly increase TA in differentiating PD from APD. The differential diagnosis of degenerative parkinsonism may thus be facilitated.

Key Words: Parkinson disease; atypical parkinsonian disorder; FP-CIT; IBZM; MIBG

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In idiopathic Parkinson disease (PD), progressive nigrostriatal denervation and degeneration in the peripheral autonomic nervous system are typical features (1,2). Clinically, this is reflected by increasing severity of characteristic motor disturbances and occurrence of axial motor symptoms such as postural abnormalities. In addition, different nonmotor symptoms may develop such as urinary or sleep disturbances (1). Although the diagnosis of PD can be unequivocal in patients with typical clinical symptoms, differentiation from atypical parkinsonian disorder (APD), such as parkinsonian type of multiple-system atrophy (MSA-P) and progressive supranuclear palsy (PSP), may be challenging in other cases (3–5). Because prognosis and treatment of patients with APD differ from those of PD patients (5), appropriate nosologic classification is warranted.

Molecular imaging techniques using PET or SPECT offer a variety of tools for diagnosing patients with parkinsonian syndromes (6–8). SPECT of the nigrostriatal dopaminergic system and of the sympathetic innervation of the heart is widely used in this context (4,9). For example, imaging of dopamine transporter (DAT) binding with 123I-N-α-fluoropropyl-2β-carbomethoxy-3β-(4-iodophenyl)nortropan (FP-CIT) successfully visualizes presynaptic dopaminergic degeneration of the nigrostriatal tract. This procedure allows differentiation of degenerative parkinsonism from movement disorders that are not associated with dopaminergic deficit, such as essential tremor (4,5,10). DAT imaging alone, however, does not differentiate the various types of degenerative parkinsonism (PD vs. APD) with sufficient accuracy (5,11).

In the latter regard, SPECT of dopamine D2 receptors with radioligands such as 123I-(S)-2-hydroxy-3-iodo-6-methoxy-N-[1-ethyl-2-pyrrodinyl]-methyl]benzamide (IBZM) may be
helpful, because APD patients usually display lower D₂ receptor binding than do PD patients (12–14). Similarly, scintigraphy with meta-¹²³I-iodobenzylguanidine (MIBG) has recently attracted interest in this context, because it also may differentiate APD and PD based on displaying pronounced cardiac sympathetic denervation in PD (15).

However, all these molecular imaging techniques are limited in terms of their test accuracy (TA). The use of FP-CIT SPECT, IBZM SPECT, or MIBG scintigraphy as stand-alone diagnostic means is therefore currently not recommended for differentiating different types of degenerative parkinsonism (11). The aim of the present study was thus to evaluate whether and to what extent the combination of FP-CIT SPECT, IBZM SPECT, and MIBG scintigraphy improved in vivo PD–APD differentiation. We here report on a first series of 48 patients with degenerative parkinsonism for whom a multidimensional statistical algorithm was applied and resulted in a marked improvement in the PD–APD discriminatory power when compared with each of the stand-alone molecular approaches.

MATERIALS AND METHODS

Patients

Between October 2005 and October 2007, a consecutive series of 48 patients with suspected degenerative parkinsonism attending the Movement Disorder Center of the University Hospital Düsseldorf was prospectively recruited. Patients underwent detailed neurologic examination (including testing of the autonomic nervous system and neuropsychologic testing), routine laboratory tests, and diagnostic brain MRI to exclude symptomatic parkinsonism, for example, multiple arteriosclerotic changes or hydrocephalus. Additionally, the Unified Parkinson Disease Rating Scale (UPDRS) (16) was determined 12 h after discontinuation of antiparkinsonian medication (off-state). Levodopa responsiveness was assessed by a short-term levodopa test (oral administration of 200–250 mg levodopa) or by a clear qualitative levodopa response during clinical long-term follow-up examinations. The levodopa test was considered positive if patients revealed improvement of more than 30%, compared with off-baseline.

All patients underwent molecular in vivo brain and cardiac diagnostics at initial presentation comprising presynaptic striatal DAT scintigraphy with FP-CIT, postsynaptic dopamine D₂ receptor scintigraphy with IBZM, and assessment of myocardial sympathetic innervation with MIBG. Exclusion criteria were history of heart disease, including coronary artery disease, infarction, or heart failure; diabetes mellitus; polyneuropathy; dementia; current neuroleptic or tricyclic antidepressive medication; and vascular cerebral lesions.

After molecular in vivo diagnostics, all patients underwent clinical follow-up examinations for 37.4 ± 12.4 (PD) and 26 ± 11.6 (APD) months. Final diagnosis of PD or APD was made by the consensus of 3 movement disorder specialists who were unaware of the results of molecular in vivo diagnostics, taking into consideration data of carefully assessed longitudinal neurologic follow-up examinations, possible response to dopaminergic treatment, and diagnostic MRI. Thirty-one patients were classified as having PD according to the U.K. Brain Bank criteria (17). Further, 11 patients were diagnosed as probable MSA-P according to established criteria (18,19), and 6 patients fulfilled the National Institute of Neurologic Disorders and Stroke criteria for probable PSP (20,21).

Scintigraphic Imaging

After the thyroid was blocked with sodium perchlorate, the tracers FP-CIT (ioflupane [¹²³I]; DaTSCAN, 184 ± 5 MBq), IBZM (iolopride [¹²³I]; IBZM, 184 ± 8 MBq), or MIBG (iobenzguane [¹²³I] injection: AdreView, 134 ± 20 MBq), all from GE Healthcare, were injected intravenously as a slow bolus in a resting state. In all patients, the 3 examinations were performed within 3 wk in randomized order, with a time interval of at least 2 d (2.8 ± 1.1 d) between MIBG scintigraphy and subsequent FP-CIT SPECT or IBZM SPECT, and an interval of, at minimum, 3 d (4.7 ± 2.3 d) after FP-CIT SPECT or IBZM SPECT. IBZM SPECT was performed 12 h after discontinuation of antiparkinsonian medication. Planar and SPECT images were acquired using a dual-head γ-camera (Prism 2000; Philips) equipped with low-energy high-resolution collimators. The imaging data were independently analyzed by 2 raters, who were unaware of the patients’ identity and clinical diagnosis.

Brain Imaging: FP-CIT SPECT and IBZM SPECT

The brain SPECT scans were acquired according to the European Association of Nuclear Medicine procedure guidelines for brain neurotransmission SPECT using ¹²³I-labeled dopamine transporter ligands (FP-CIT) or ¹²³I-labeled dopamine D₂ receptor ligands (IBZM) (22,23). SPECT image acquisition was started 4 h after injection for FP-CIT and 90 min after injection for IBZM in all patients, with a 128 × 128 image matrix and 120 projections over 360°. Transversal slices were reconstructed by filtered back-projection using a low-pass filter (cutoff, 0.22 Nyquist; order, 7) and attenuation-corrected by Chang’s method (24). Orbital-mesetal parallel slices were created by reangulation of the dataset and 3-dimensional coregistration to a normalized MRI template. Images were evaluated semiquantitatively using template-based regions of interest (ROIs) and calculation of regional binding potentials as striatum–posterior cortex (S/P) ratios (Supplemental Fig. 1A; supplemental materials are available online only at http://jnm.snmjournals.org).

Cardiac Imaging: MIBG Scintigraphy

For the assessment of myocardial MIBG uptake, planar and SPECT scans were acquired. The planar images were acquired at 4 h after injection (5 min, 128 × 128 matrix). Immediately before or after the acquisition of planar images, a SPECT scan of myocardial sympathetic innervation was obtained for each patient to exclude focal deficits of tracer accumulation. The SPECT images were analyzed visually after iterative reconstruction without attenuation correction. Myocardial MIBG uptake was evaluated using standardized ROIs as heart-to-mediastinum (H/M) ratio on planar images 4 h after injection (Supplemental Fig. 1B).

Statistical Evaluation

Group data are presented as mean ± SD. For statistical comparison, the 2-sided t test was used for normally distributed data and the 2-sided Mann–Whitney U test for nonnormally distributed data. Normality of data distribution was assessed using the Kolmogorov–Smirnov test. Differences with a P value of less than 0.05 were considered biologically significant. Descriptive statistics and basic comparisons were analyzed using the Medcalc software package (version 11.1). To evaluate interrater variance, the interrater agreement statistic was computed (weighted k).
RESULTS

At the time of molecular diagnostics, the mean age of PD patients \((n = 31)\) was 64 ± 8.5 y (range, 41–74 y), and the mean age of APD patients \((n = 17)\) was 65.9 ± 8.3 y (range, 49–78 y, \(P = 0.49\), Mann–Whitney \(U\) test, 2-sided). In the PD group, disease duration (6.7 ± 6.7 y) and disease severity (UPDRS-off score, 29.6 ± 19.5) were similar to those of the APD group (disease duration, 5.8 ± 3.3 y, \(P = 0.48\); UPDRS-off score, 32.5 ± 14.2, \(P = 0.42\), Mann–Whitney \(U\) test, 2-sided). PD patients displayed a positive response to levodopa or robust clinical improvements of motor symptoms during continuing dopaminergic therapy in all cases, whereas only 3 of 11 (27%) MSA-P patients showed a levodopa responsiveness. PSP patients revealed no relevant short- or long-time levodopa response (Supplemental Table 1).

During the follow-up period, 14 PD patients (45%) developed motor fluctuations; 8 of these patients (26%) underwent deep brain stimulation. Three patients (10%) died from non-PD–related disorders (gastric carcinoma, heart attack). The APD patients demonstrated a much more rapid worsening of clinical symptoms and lower long-time survival rates. All MSA-P patients \((n = 11)\) developed severe urinary or fecal incontinence and hypotension, and the PSP patients \((n = 6)\) demonstrated a marked postural instability with spontaneous falls and vertical supranuclear palsy. Eleven APD (9 MSA-P, 2 PSP) patients died (65%; median survival rate, 7.81 y) from disease-related complications (pneumonia, suffocation, sepsis).

Imaging Results

The individual binding data are plotted in Figure 1 for all 3 molecular imaging modalities. The interrater reliability was high (for FP-CIT SPECT: weighted \(\kappa = 0.972\), and sensitivity, 0.158; for IBZM SPECT: weighted \(\kappa = 0.943\), and sensitivity, 0.155; and for MIBG scintigraphy: weighted \(\kappa = 0.955\), and sensitivity, 0.143).

Mean binding of FP-CIT was significantly higher in PD patients (contralateral, 1.96 ± 0.39; ipsilateral, 2.09 ± 0.50) than in APD patients (contralateral, 1.59 ± 0.22, Mann–Whitney \(U\) test, \(P = 0.001\); ipsilateral, 1.68 ± 0.32, \(t\) test, \(P = 0.004\); Fig. 1A).

Mean \(D_2\) receptor binding ratios were significantly lower in APD patients (contralateral, 1.44 ± 0.16, \(t\) test; ipsilateral, 1.46 ± 0.13, \(t\) test; Fig. 1B) than in PD patients (contralateral, 1.60 ± 0.11, \(P = 0.0007\); ipsilateral, 1.59 ± 0.10, \(P = 0.001\)).

Neither FP-CIT SPECT nor IBZM SPECT revealed significant side-to-side differences in PD or APD \((t\) test).

Mean myocardial MIBG accumulation was significantly lower in the PD group (ratio, 1.34 ± 0.27) than in the APD group (1.60 ± 0.29, Mann–Whitney \(U\) test, \(P = 0.0039\); Fig. 1C).

Although statistical analysis of FP-CIT, IBZM, and MIBG values revealed significant group differences between PD and APD for each molecular imaging modality, the overlap in modalities was considerable. Thus, in individual cases,
differentiation between PD and APD may be impossible, as evidenced in Figure 2, which depicts FP-CIT, IBZM, and MIBG accumulation in PD (A and C) and APD (B and D) patients. A typical combination of results is demonstrated in Figures 2A and 2B: decreased FP-CIT binding in PD and APD but maintained or even increased D₂ receptor density and severely decreased cardiac MIBG accumulation in PD is shown in Figure 2A (patient 26: FP-CIT, 1.60; IBZM, 1.68; MIBG, 1.13), and maintained cardiac innervation but reduced D₂ receptor binding in APD is shown in Figure 2B (patient 45: FP-CIT, 1.40; IBZM, 1.38; MIBG, 2.01).

In Figures 2C (patient 3, PD: FP-CIT, 2.33; IBZM, 1.72; MIBG, 1.75) and 2D (patient 38, APD: FP-CIT, 1.66; IBZM, 1.54; MIBG, 1.76), the results of MIBG scintigraphy and IBZM SPECT are inconclusive. In both cases, diagnoses were correctly made when considering FP-CIT SPECT results.

ROC Analyses

To identify the optimally discriminating cutoff values in the differentiation of APD and PD, we calculated ROC curves for the 3 individual imaging modalities (Fig. 3) and for all logical combinations of these molecular imaging modalities.

**Individual Imaging Modalities.** Using ROC analysis, we determined the following cutoff values for differentiating APD from PD: for FP-CIT SPECT, 1.73 or less (AUC, 0.775; 95% CI, 0.632–0.883; Fig. 3A); for IBZM SPECT, 1.47 or less (AUC, 0.772; 95% CI, 0.628–0.881; Fig. 3B); and for MIBG scintigraphy, more than 1.34 (AUC, 0.755; 95% CI, 0.610–0.868; Fig. 3C). The highest TA (79%) was evident for IBZM SPECT, with a sensitivity of 53%, specificity of 94%, PPV of 82%, and NPV of 78%. For MIBG scintigraphy, TA was 73% (sensitivity, 88%; specificity, 65%; PPV, 58%; NPV, 91%), and for FP-CIT SPECT, TA was 73% (sensitivity, 76%; specificity, 71%; PPV, 59%; NPV, 85%).

**Combined Imaging Modalities.** We further optimized cutoff values for the possible logical combinations of the different imaging techniques. The highest TA (94%) was reached by combining IBZM SPECT, FP-CIT SPECT, and MIBG scintigraphy under the assumption that a diagnosis of APD has to be made whenever this is suggested by the results of at least 2 of the 3 methods. Calculated cutoff values for this combination were as follows: IBZM SPECT, 1.46 or less; MIBG scintigraphy, more than 1.43; and FP-CIT SPECT, less than 2.10 (Table 1, middle column). These cutoff values distinguished APD from PD with a sensitivity of 94%, a specificity of 94%, a PPV of 89%, and an NPV of 97%.

The second best results were found for the assumption that all 3 methods must indicate APD, using the following cutoff values: IBZM SPECT, 1.73 or less; MIBG scintig-
raphy, 1.31 or more; and FP-CIT SPECT, less than 2.10 (Table 1, first column). Under these circumstances, TA amounts to 92%, sensitivity to 88%, specificity to 94%, PPV to 88%, and NPV to 94%. The 1-of-3 combination (Table 1, last column) reached a TA of only 83% for the APD diagnosis (sensitivity, 94%; specificity, 77%; PPV, 70%; NPV 96%) using the following cutoff values: IBZM SPECT, 1.46 or less; MIBG scintigraphy, 1.41 or more; and FP-CIT SPECT, 1.41 or less.

**Specified Results of 2-of-3 Combination**

On closer examination of the 2-of-3 combination yielding the highest TA, the results for IBZM SPECT and MIBG scintigraphy were in agreement for 29 patients (4 correctly positive, APD; 24 correctly negative, PD; and 1 falsely negative). In 19 patients, the results of IBZM SPECT and MIBG scintigraphy were discordant. Considering the results of FP-CIT SPECT in these cases led to a correctly positive result (APD) in 12 patients, a correctly negative result (PD) in 5 patients, and an incorrectly positive result in 2 patients.

**DISCUSSION**

This study explores to which extent the combined use of IBZM SPECT, MIBG scintigraphy, and FP-CIT SPECT and their multidimensional statistical evaluation may improve diagnostic accuracy in the differential diagnosis of degenerative parkinsonism. The analysis shows moderate TA for PD versus APD differentiation using each molecular approach alone (IBZM SPECT, 79%; MIBG scintigraphy, 73%; and FP-CIT SPECT, 73%). The highest TA resulted if at least 2 of the 3 biologic markers were positive for APD using the following cutoff values: IBZM SPECT, 1.46 or less; FP-CIT SPECT, less than 2.10; and MIBG scintigraphy, more than 1.43. This algorithm distinguished APD from PD with a sensitivity of 94%, a specificity of 94% (TA also 94%), a PPV of 89%, and a NPV of 97%.

**TABLE 1**

Diagnosis of APD: Evaluative Statistical Parameters for Combined Use of FP-CIT SPECT, IBZM SPECT, or MIBG Scintigraphy for Differentiation of APD from PD

<table>
<thead>
<tr>
<th>Parameter</th>
<th>3 of 3</th>
<th>At least 2 of 3</th>
<th>Any 1 of 3</th>
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<tbody>
<tr>
<td>Cutoff values</td>
<td>FP-CIT &lt; 2.10</td>
<td>FP-CIT &lt; 2.10</td>
<td>FP-CIT ≤ 1.41</td>
</tr>
<tr>
<td></td>
<td>IBZM ≤ 1.73</td>
<td>IBZM ≤ 1.46</td>
<td>IBZM ≤ 1.46</td>
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<tr>
<td></td>
<td>MIBG ≥ 1.31</td>
<td>MIBG &gt; 1.43</td>
<td>MIBG ≥ 1.41</td>
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<tr>
<td>Sensitivity (%)</td>
<td>88</td>
<td>94</td>
<td>94</td>
</tr>
<tr>
<td>Specificity (%)</td>
<td>94</td>
<td>94</td>
<td>77</td>
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<tr>
<td>NPV (%)</td>
<td>94</td>
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<td>PPV (%)</td>
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<td>89</td>
<td>70</td>
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<tr>
<td>TA (%)</td>
<td>92</td>
<td>94</td>
<td>83</td>
</tr>
</tbody>
</table>

Cutoff values were calculated for all possible logical combinations of 3 imaging techniques and optimized using maximum TA as optimization criterion. Accordance of at least 2 of 3 biologic markers using indicated cutoff values yielded highest TA of 94% for diagnosing APD (middle column).
In clinical terms, the ability of widely available molecular imaging techniques combined with this algorithm to distinguish APD from PD with such high accuracy is remarkable, possibly allowing the correct discrimination between APD and PD to occur earlier in the disease course.

**Unimodal Investigations**

A series of studies has been conducted using singular molecular components of synapses as potentially facilitating the differential diagnosis of degenerative parkinsonism.

**Presynaptic Nigrostriatal System**

Imaging of presynaptic DAT can be assessed by a variety of SPECT and PET tracers (25). FP-CIT SPECT is thus of widespread clinical use and allows highly accurate differentiation between degenerative parkinsonism and movement disorders not associated with a dopamine deficit such as essential tremor. Plotkin et al., for example, report a sensitivity, specificity, and TA of 93%, 100%, and 94%, respectively, in this context (12). In patients with degenerative parkinsonism, however, striatal DAT accumulation shows moderate TA in differentiating PD from APD, because loss of dopaminergic neurons in the substantia nigra and reduction of striatal dopamine projections are the histopathologic hallmark in all of these disorders. A sensitivity and a specificity of 81% were reported for distinguishing PD from MSA (26); these were close to our results of 76% sensitivity and 71% specificity.

Several different analyses to distinguish PD from APD based on presynaptic properties of the dopaminergic system reflect the current clinical need for differentiation. First, neuropathologic studies suggest that, compared with PD, the substantia nigra in APD is more uniformly affected in MSA and PSP, with disturbed projections to both putamen and caudate nucleus. Because in PD more ventrolateral areas of the substantia nigra and its putaminal projections are targeted, several SPECT studies have tried to objectify a specific pattern or degree of striatal DAT uptake in this area, allowing improvement of the TA of the differential diagnosis in degenerative parkinsonism (5,9,27–31). On the basis of these results, some groups have reported that in early untreated PD patients the relative sparing of the caudate nucleus in FP-CIT SPECT might be helpful in distinguishing patients with PD from patients with APD (31,32), whereas others have found these results of limited value (9,12,33).

Second, a few studies suggest that a left-to-right asymmetry in DAT binding may be helpful for the diagnostic characterization of PD (28,34). In this context, Knudsen et al. observed that an asymmetry index of striatal DAT binding greater than 15% allowed discrimination between early-to-moderate PD and MSA, with a sensitivity of 86% (34). However, these results were obtained in a small sample and are contradictory to other findings (9,12,31), including our own results, which do not reveal significant asymmetry indices in drug-treated PD patients.

Third, quantification of striatal subregions and calculation of specific DAT binding indices have not consistently been reported to distinguish different degenerative parkinsonian syndromes with high TA and therefore should be used with caution in standard clinical practice (14).

Fourth, in a study using $^{125}$I-2 β-carbomethoxy-3β-(4-iodophenyl)tropane (β-CIT) SPECT, a highly accurate classification of patients with APD and PD has been reported for the use of an operator-independent statistical parametric mapping, which has succeeded in localizing a specific reduction of specific binding in mid brain and pons of APD patients (30,35). However this voxel-based analysis technique, unlike ROI-based methods, has not yet been validated in routine clinical use and is therefore not generally recommended (4).

Moreover, the results probably cannot be transferred to FP-CIT SPECT, because, compared with β-CIT, FP-CIT has a much higher affinity to the DAT and a much lower affinity to serotonin transporters, the latter being mainly responsible for accumulation of β-CIT in pons and mid brain.

Finally, the progression of DAT decline is 2 times faster in APD than in PD; therefore, longitudinal DAT imaging might become a helpful diagnostic biomarker (29).

**Dopamine D$_2$ Receptors**

Upregulation of posterior putamen D$_2$ receptor binding in early and drug-naive degenerative parkinsonism was reported to be indicative of PD, whereas reduction could potentially be specific for APD (9,12). The use of specific striatal D$_2$ receptor binding as a stand-alone procedure has thus been shown as a reasonable discriminating parameter in differential diagnosis of degenerative parkinsonism. Findings of a symmetrically decreased mean D$_2$ receptor binding in APD (9,34) correspond to the SPECT results of our study, but as with FP-CIT SPECT, there is a varying degree of overlap between APD and PD. For detection of APD, a TA of 70%–90%, a specificity of 70%–100%, and a sensitivity of 60%–87% are reported (10,12,14,36). Our ROC analysis results for the isolated use of IBZM SPECT agree with these data, yielding a TA of 79%, a high specificity of 94%, but only a moderate sensitivity of 53%.

**Cardiac MIBG Uptake**

For MIBG scintigraphy, the overall H/M ratio (1.34) of the PD patients in the present study confirms previous results reporting H/M ratios between 1.19 and 1.57 depending on disease duration, PD subtype, and (probably) also on the exact method of quantification such as size, shape, and positioning of ROIs and the use of maximum compared with mean count values (15,28,37–39). Nevertheless, in any case considerable overlap exists for the PD–APD differentiation, suggesting also that MIBG scintigraphy alone is not sufficient to distinguish these entities and that low MIBG uptake alone does not necessarily indicate PD (40).

**Multimodal investigations**

In summary, the present results on the stand-alone analyses of FP-CIT SPECT, IBZM SPECT, and MIBG scintigraphy concur with previous studies reporting presynaptic nerve terminal affection, striatal D$_2$ receptor involvement, and myo-
cardiac sympathetic dysfunction in both PD and APD patients (9,10,12,14,31,34,41). Specifically, the present data reproduce the lower cardiac MIBG uptake in PD (vs. APD). Similarly, DAT and D2 receptor binding were lower in APD than in PD (5,9,12,14,15,39). However, as also previously shown (5,12,40), there is a substantial overlap between PD and APD for each of these 3 biologic markers that markedly limits their respective diagnostic accuracy. We therefore examined to what extent the combination of these 3 tracers would lead to an improvement of TA. Our data show that by combining FP-CIT SPECT, IBZM SPECT, and MIBG scintigraphy and using a multidimensional statistical approach, TA can be raised to 94%, assuming that APD diagnosis is indicated by at least 2 of the 3 molecular approaches.

Only a few studies have evaluated the potential diagnostic benefit of combining pre- and postsynaptic dopaminergic imaging in patients with PD versus APD (9,12,34). These studies reported specific binding ratios and regional differences in striatal uptake patterns for each imaging technique alone but did not develop specific cutoff values for combined pre- and postsynaptic quantitative measures. Only Koch et al. calculated the diagnostic accuracy for the combined use of IBZM SPECT and FP-CIT SPECT using an automated and observer-independent quantification method (14). They reported a higher diagnostic power for the combined use that could be further increased if a combination of pre- and postsynaptic radiotracer binding asymmetries was considered in conjunction with striatal D2 receptor binding (14). However, this approach requires a complex calculation algorithm, and sensitivity (90.3%), specificity (73.9%), and TA (79%) were inferior to that of our study.

**Practical Considerations**

To minimize costs and radiation exposure, we suggest first combining IBZM SPECT and MIBG scintigraphy. A combination of well-preserved striatal D2 receptor binding (S/P ratio $\geq$1.46) and a marked reduction of myocardial MIBG accumulation (H/M ratio $\leq$1.43) are typical for PD and almost certainly exclude APD. On the other hand, if a combination of reduced D2 receptor density (ratio <1.46) and normal or only mildly reduced cardiac sympathetic innervation, as evidenced by an H/M ratio greater than 1.43, is indicative of APD. In patients with borderline or contradictory results based on IBZM SPECT and MIBG scintigraphy, an additional FP-CIT SPECT scan (S/P ratio <2.10) will be helpful to confirm or exclude APD diagnosis.

Factors such as instrumentation, reconstruction, filtration, and image processing technique may influence the quantitative results of these scintigraphic measures (22,23). Therefore, the ratios reported here may need adjustments according to the acquisition and processing specifics of the individual imaging facility.

**CONCLUSION**

Results suggest that the combination of FP-CIT SPECT, IBZM SPECT, and MIBG scintigraphy imaging may lead to a marked improvement in distinguishing PD from APD. This approach is completely based on commercially available tracers and can therefore be used in a clinical setting in the absence of short-living PET tracers. Because the present data are based on a small sample size from 1 center, confirmation of this approach in a large population is warranted using a multicenter design.

**DISCLOSURE STATEMENT**

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