Is transcranial sonography useful for diagnosing Parkinson’s disease in clinical practice?

O ultrassom transcraniano é útil para o diagnóstico da doença de Parkinson na prática clínica?

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ABSTRACT

Transcranial sonography (TCS) is an emerging ancillary examination for diagnosing Parkinson’s disease (PD). Objective: To evaluate TCS features in patients with PD and its mimics, and establish their accuracy in predicting the final clinical diagnosis after follow-up. Methods: We retrospectively studied 85 patients with an initial clinical suspicion of PD, atypical parkinsonism or essential tremor, all of whom underwent TCS. Two specialists reviewed the follow-up clinical visit records and determined the final clinical diagnosis. The accuracy analysis of the TCS was determined using Bayesian statistical methods. Results: The finding of substantia nigra hyperechogenicity (> 20 mm²) showed high sensitivity (93.4%) and specificity (86.6%). The positive likelihood ratio showed 6.93-fold greater odds for diagnosing PD than an alternative condition when this finding was present. Conclusions: This study revealed the practical usefulness of TCS in differentiating PD from its prevalent mimics when the clinical diagnosis was initially unclear.

Keywords: Parkinson's disease; parkinsonian disorders; essential tremor; ultrasonography; diagnostic techniques and procedures.

The diagnosis of idiopathic Parkinson’s disease (PD) remains a challenge. Although the clinical opinion of experts remains the gold standard, only 53-75% of their diagnoses agree with the definite pathological diagnosis; the major misdiagnoses being essential tremor (ET) and atypical parkinsonism (AP). As suggested by the European Federation of Neurological Societies/Movement Disorder Society – European Section recommendations, ancillary examinations should be effective in establishing the differential diagnosis of PD. Imaging and biomarkers are urgently required to improve the certainty of the current unsatisfactory clinical parameters.

Transcranial sonography (TCS) is used to evaluate the echogenicity of the midbrain and basal ganglia. The sensitivity of TCS is high (90%) in discriminating PD cases from not only a healthy control group but also major PD mimics, such as ET and AP. The TCS parameters have been standardized in diverse...
suggests that TCS might enable early diagnosis16,18. Thus, we aimed to evaluate the diagnostic performance of TCS, specifically in patients with an initially-undefined etiology for parkinsonian motor symptoms under real clinical practice conditions, i.e., in a less-controlled setting than that of a clinical trial protocol.

METHODS

Study population

We retrospectively studied 126 patients who were consecutively evaluated in the outpatient clinic of our Movement Disorders reference center between January 2015 and June 2016. These patients presented with parkinsonian features (resting tremor, bradykinesia, postural instability or rigidity), but did not fulfill all criteria for the diagnosis of PD (the UK Brain Bank) at the initial visit. Due to the uncertainty of the diagnosis, patients were referred for a TCS examination to exclude or confirm the suspicion of idiopathic PD. Among all patients who underwent the examination during the study period, we excluded 21 patients due to suspicion of other clinical diagnoses that we did not want to evaluate in this study, such as cerebellar ataxia, Huntington’s disease, dystonic tremor, and metabolic or toxic tremor12. We also excluded 20 patients who were not able to be evaluated with TCS due to insufficient temporal bone windows, which technically limited the visualization of the midbrain contour. The local ethical standards committee approved the study.

First diagnostic hypothesis

An initial clinical suspicion was defined for each patient in order to compare the first clinical impression of the neurologist with the final diagnosis after follow-up. This initial clinical suspicion was based on the initial clinical data acquired by the physician before the TCS was performed. None of the patients had a defined clinical diagnosis at the time they underwent TCS. However, all patients had an initial clinical suspicion of PD, AP (including multiple system atrophy and progressive supranuclear palsy) or ET. We selected these groups of patients because previous studies have shown that TCS can reliably distinguish these diagnoses from PD19,20. These patients would thus be the best candidates for the TCS examination in a real clinical scenario.

Clinical assessment and diagnosis

Two movement disorder experts reviewed all clinical records from at least six months of follow up after the TCS examination to determine the definitive clinical diagnosis by consensus. In addition to the UK Brain Bank criteria, nonmotor symptoms, levodopa response, persistent asymmetry, unilateral start, hypsomia, and REM sleep behavior disorder, but not the TCS result, were used to define a patient as having PD. Medical records were also examined for general clinical criteria for Lewy body dementia, progressive supranuclear palsy, multiple system atrophy and ET19,20,21,22. After the follow-up period, the patients were labeled as having ET, PD or AP if a clinical diagnosis was achieved. If clinical uncertainty regarding the diagnosis remained, the clinicians labeled the patients as having an “undefined diagnosis”. In addition, if the exclusion criteria for PD (UK Brain Bank) were met and no established clinical criteria for AP or ET were present, the patients were labeled as “excluded PD”. To perform between-group comparisons, the data analysis procedures combined patients with Lewy body dementia and PD into the same category - PD - as Lewy body dementia and PD are believed to be distinct clinical presentations of the same disease with the same pathological substrate but distinct anatomical distribution23. The cases that certainly did not represent PD cases (namely, the AP, ET and “excluded PD” groups) were labeled altogether as the “non-PD” group for some of the analyses. The methodological process is outlined in Figure 1.

Acquisition and assessment of TCS images

We performed the TCS examinations according to the guidelines of the European Society of Neurosonology and Cerebral Hemodynamics23. An Esaote MyLab25Gold ultrasound machine (Genoa, Italy) was used along with a 2-3.5MHz phased array transducer. The transtemporal acoustic bone window was used to assess the midbrain bilaterally, and after a 10-degree tilt of the ultrasound beam, the examiner also scanned the thalamic axial plane. The tissue equalization function and gain adjustments were used if necessary23. Characteristic findings are shown in Figure 2.

Based on the published consensus, the SN-TCS was scored as “positive” (SN+) if the SN displayed an increased echo intensity on at least one side relative to the surrounding brainstem tissue with an area value > 20 mm²18; and scored as “markedly positive” with an area value > 25 mm²18. The largest SN planimetric value for each patient was chosen for analysis regardless of the side of the measurement. The examiner qualitatively classified the lentiform nucleus as isoechogenic (LN-) or hyperechogetic (LN+) compared with the surrounding tissues. In addition, the third ventricle (V3) and lateral ventricle widths were measured.

Statistical analysis

Due to the limited number of participants, we used Bayesian methods to analyze our data in order to improve the veracity of the results. We applied t-distribution, the Bayesian one-way ANOVA24, receiver operating characteristic (ROC) curve analysis and the Bayesian test of accuracy as required by each situation and as described in the results section.
Patients with undefined clinical diagnosis referred to TCS between 01/2015 and 06/2016
N = 126

No TCS due to unavailable temporal sonographic window
N = 20 (16%)

Transcranial sonography exams acquired
N = 106 (84%)

Reviewed Initial suspicion of PD, AP or ET
N = 85 (81%)

Reviewed Consensus diagnosis after follow-up
N = 65 (77%)

Excluded due to other movement initial suspicion
N = 21 (20%)

Diagnosis not defined due to not meeting all the criteria
N = 20 (23%)

Clinical Defined non-PD
N = 26 (40%)

Clinical Defined PD
N = 39 (60%)

Clinical AP
N = 9 (35%)

Clinical ET
N = 11 (42%)

Clinical Excluded-PD
N = 6 (23%)

Clinical Idiopathic PD
N = 34 (87%)

Clinical Lewy Body Dementia
N = 5 (13%)

Figure 1. Trial schematic. Detailed schematic showing the patient inclusion process.

Figure 2. TCS images. (A) Delineation of the hyperechogenic SN in the left hemisphere and mesencephalic plane. (B) Delineation of the hyperechogenic SN in the right hemisphere and mesencephalic plane. (C) Delineation of the hyperechogenic LN in the thalamic plane. III Ventr: Third ventricle; and S. Nigra: substantia nigra.
RESULTS

We assessed 85 patients (51 men and 34 women) with a mean age of 67 (17-88) years (Table 1). The average clinical follow-up time after the TCS exam was 17 months. A definitive clinical diagnosis was determined for 65 (76%) of the recruited patients (Figure 1).

The initial clinical suspicion was AP in 18 (21%), ET in 15 (18%) and PD in 52 (61%) considering all the evaluated patients. A definitive diagnosis was achieved in 65 of these patients. Regarding the 65 patients with a final diagnosis, the initial diagnostic hypothesis was confirmed in 55 of them, representing a diagnostic consistency of 85% (Table 2). Twenty-one percent (n = 7) of the patients who were initially suspected to have PD had a different final diagnosis, including ET (n = 2) and "excluded PD" (n = 5). Moreover, two (18%) of the patients initially classified as ET were later diagnosed as PD.

We applied Bayesian statistics using a robust (t-distribution) one-way ANOVA to compare the demographic characteristics among the groups with definitive clinical diagnoses, and the results did not show statistically significant differences in the mean ages.

The posterior median of the SN area robust mean was defined as the central measure. With the intent of comparing the PD and non-PD groups, a difference in the distributions of the central tendencies was obtained. Compared to all other clinically diagnosed groups, PD patients had the highest SN hyperechogenicity area values, which were significantly different compared with those of the other groups. The value of the SN hyperechogenic area among all groups that comprised the non-PD group did not differ in the between-group comparisons (Table 3).

The distribution of the frequencies of SN hyperechogenicity (area > 20 mm²) in patients with a final diagnosis of PD and in patients who were not diagnosed with PD (non-PD) is shown in Table 4. The Bayesian proportions test was applied to evaluate the chances of SN hyperechogenicity (> 20 mm²) occurrence in the PD and non-PD groups. The PD group had an estimated relative frequency of SN hyperechogenicity of 0.93 [0.85, 0.99], and the non-PD group of 0.13 [0.03, 0.27]. The estimated group difference (PD minus non-PD) was statistically significant (-0.79 [-0.92, -0.63]), showing that the relative frequency of SN hyperechogenicity was higher for patients diagnosed with PD.

Lentiform nucleus hyperechogenicity (LN+) was observed in 10% (n = 4) of patients diagnosed with PD and 22% (n = 2) of patients diagnosed with AP. Moreover, enlargement of the V3 was observed in 11% (n = 1) of patients in the AP group and 5% (n = 2) of patients in the PD group. These features have been used in other studies to help distinguish between PD and AP. However, when the Bayesian proportions test was used to assess the association of LN+ or V3 enlargement with the diagnoses of AP and PD, a between-group difference was not observed.

The Bayesian accuracy test was applied to evaluate the diagnostic accuracy of TCS based on the following central question: "Is SN hyperechogenicity a finding that could predict a clinical diagnosis of PD in this cohort after follow-up?". Using the cut-off value of 20 mm², the finding of SN hyperechogenicity showed robust positive likelihood ratios (6.93) for a prospective PD clinical diagnosis when the group with a definitive clinical diagnosis of PD was compared with the group diagnosed with alternative clinical disorders (non-PD group). When using the cut-off value of

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**Table 1. Epidemiological data.**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Frequency (n)</th>
<th>Percentage (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Women</td>
<td>34</td>
<td>40.0</td>
</tr>
<tr>
<td>Men</td>
<td>51</td>
<td>60.0</td>
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<tr>
<td>Initial suspicion</td>
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<tr>
<td>Atypical parkinsonism</td>
<td>18</td>
<td>21.2</td>
</tr>
<tr>
<td>Parkinson’s disease</td>
<td>52</td>
<td>61.2</td>
</tr>
<tr>
<td>Essential tremor</td>
<td>15</td>
<td>17.6</td>
</tr>
<tr>
<td>Final diagnosis</td>
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<td></td>
</tr>
<tr>
<td>Atypical parkinsonism</td>
<td>9</td>
<td>10.6</td>
</tr>
<tr>
<td>Undefined</td>
<td>20</td>
<td>23.5</td>
</tr>
<tr>
<td>Excluded Parkinson’s disease</td>
<td>6</td>
<td>7.1</td>
</tr>
<tr>
<td>PD or Lewy body dementia</td>
<td>39</td>
<td>45.9</td>
</tr>
<tr>
<td>Essential tremor</td>
<td>11</td>
<td>12.9</td>
</tr>
<tr>
<td>Symptom side</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right side</td>
<td>45</td>
<td>52.9</td>
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<td>Left side</td>
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<td>Both sides</td>
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<td>Family history</td>
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<td>Positive</td>
<td>14</td>
<td>16.4</td>
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<tr>
<td>Negative</td>
<td>29</td>
<td>34.1</td>
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<tr>
<td>Unknown</td>
<td>42</td>
<td>49.5</td>
</tr>
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</table>

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**Table 2. Diagnosis distribution.**

<table>
<thead>
<tr>
<th>Initial clinical suspicion</th>
<th>Final clinical diagnosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parkinson’s disease (n = 52), 61%</td>
<td>Parkinson’s disease (n = 37), 71%</td>
</tr>
<tr>
<td></td>
<td>Essential tremor (n = 2), 4%</td>
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<tr>
<td></td>
<td>Excluded Parkinson’s (n = 5), 10%</td>
</tr>
<tr>
<td></td>
<td>Undefined (n = 8), 15%</td>
</tr>
<tr>
<td>Atypical parkinsonism (n = 18), 21%</td>
<td>Atypical (n = 9), 50%</td>
</tr>
<tr>
<td></td>
<td>Undefined (n = 9), 50%</td>
</tr>
<tr>
<td>Essential tremor (n = 15), 18%</td>
<td>Essential tremor (n = 9), 60%</td>
</tr>
<tr>
<td></td>
<td>Parkinson’s disease (n = 2), 13%</td>
</tr>
<tr>
<td></td>
<td>Excluded Parkinson’s (n = 1), 7%</td>
</tr>
<tr>
<td></td>
<td>Undefined (n = 3), 20%</td>
</tr>
</tbody>
</table>
DISCUSSION

Even in specialized movement disorder clinics, approximately 10–25% of PD patients may be misdiagnosed using standard clinical diagnostic criteria when compared with autopsy findings. This rate of misdiagnosis emphasizes the need for complementary methods and biomarkers for diagnosing PD. A similar demand exists for reliable statistical approaches, such as Bayesian methods, to enable more accurate measurements of the power of these markers for diagnosing PD.

Currently, TCS is consolidated as a practical diagnostic tool that can discriminate between PD and PD mimics in late clinical stages. However, in clinical practice, one of the difficulties in a movement disorder clinic is distinguishing between PD, AP and ET in early clinical stages. Likewise, TCS also seems to be valid in the differential diagnosis of PD even in the early stages. If the patient presents with an incomplete clinical syndrome, the finding of a hyperechogenic SN suggests a diagnosis of PD, whereas the absence of this result indicates an alternative condition.

The Bayesian estimate for a ROC curve revealed an area under the curve of 0.945, displaying high sensitivity for identifying AP (0.884) and ET (0.934), respectively, with low specificity (0.163). Moreover, the positive predictive value, sensitivity, and specificity of this measure for PD diagnosis were also high (greater than 90%), as shown in Table 5.

In addition, when applying the same accuracy test, hyperechogenic SN or V3 enlargement showed high sensitivity for identifying AP (0.884) and ET (0.934), respectively, with low specificity (0.163). Moreover, the positive predictive value, sensitivity, and specificity for identifying AP were 0.93 [0.83, 0.98] and 0.83 [0.70, 0.93], respectively. The Bayesian estimate for a ROC curve revealed an area under the curve of 0.945, displaying high sensitivity for identifying AP (0.884) and ET (0.934), respectively, with low specificity (0.163).

Although the clinical follow-up period in our study was a minimum of six months, most of the patients had complained of symptoms for more than three years. However, not all patients in the definitively-diagnosed group had a five-year follow-up period, and some patients had not received a final diagnosis by the end of this study (n = 20) because they still did not fulfill all the diagnostic criteria. The rate of disagreement between the first clinical suspicion and the definitive diagnosis was 14% (n = 9), which is consistent with the rates previously described for this scenario.

The TCS results considering the SN > 20 mm² agreed with the final clinical diagnosis in all of the cases in which the clinical diagnosis had to be changed from PD to non-PD (n = 7) or from ET to PD (n = 2). Although the TCS results were not considered when these final clinical diagnoses were considered to be statistically significant if the highest density intervals (HDI) did not cross the value zero.
determined, our study results highlight the suitability of TCS in enabling clinicians to reach the proper diagnosis earlier, as SN hyperechogenicity was highly predictive of the subsequent fulfillment of the UK Brain Bank criteria. Likewise, the absence of SN hyperechogenicity predicted the subsequent clinical characterization of a non-PD clinical diagnosis.

The high positive predictive value reported in the present study (91.2%) is consistent with the results from a study using a prospective methodology and preselected patients (92.9%). It is also a plausible argument for the early use of this test in clinical practice, namely, when most of the motor and nonmotor symptoms have not yet manifested. Besides being a study carried out in a clinical practice scenario, our results also reproduced a high negative predictive value (85.7%), revealing the efficacy of TCS in excluding a PD diagnosis based on negative results.

The LR is a measure of how the test would influence the diagnostic decision, and considers sensitivity and specificity, but is not affected by the disease prevalence. Although underused, the LR can refine the clinical diagnosis. The LR considers the probability that the patient will present the characteristic of interest before the test (in our case, a PD diagnosis) and the extent to which the test can increase the probability of the characteristic occurrence. A positive likelihood ratio (LR+) value of 6, for example, represents an increase of 35% over the pretest probability that the characteristic will occur based on the test results. A positive LR ranging from 2-5 represents a small increase in post-test probability that the characteristic will occur, whereas values from 5-10 indicate a moderate increase in the post-test probability, and values greater than 10 indicate a large increase in the post-test probability. In this study, a significant increase (6.93 times) was observed in the odds that an undiagnosed case of parkinsonism would receive a subsequent PD diagnosis (based on UK Brain Bank criteria) if a hyperechogenic SN (> 20 mm²) was observed. This finding reflects an increase of approximately 35% from the previous post-test probability of diagnosis. With findings of SN > 25 mm², the LR+ is even higher (12.87) and represents an increase of more than 45% for the probability of receiving a PD diagnosis. Similar positive LR values have also been described when evaluating the odds of an asymptomatic subject prospectively developing specific PD symptoms when SN hyperechogenicity was observed. These data reveal how a clinician can apply the results of SN hyperechogenicity evaluation in a simple and accurate manner when assessing a possible case of PD.

In the present study, most likely due to our small sample size, the isolated finding of LN hyperechogenicity in the TCS examination was not a unique feature that could differentiate PD from AP as already described in other studies. However, in LN hyperechogenicity, this feature, when combined with the

**Table 6. Bayesian ROC curve.**

<table>
<thead>
<tr>
<th>Group</th>
<th>N</th>
<th>ROC area</th>
<th>Cut (mm²)</th>
<th>Sensitivity</th>
<th>Specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td>ET x PD</td>
<td>50</td>
<td>0.963</td>
<td>20.0</td>
<td>0.939</td>
<td>0.952</td>
</tr>
<tr>
<td>(n = 11 x n = 39)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AP x PD</td>
<td>48</td>
<td>0.858</td>
<td>21.9</td>
<td>0.910</td>
<td>0.691</td>
</tr>
<tr>
<td>(n = 9 x n = 39)</td>
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<tr>
<td>Non-PD x PD</td>
<td>65</td>
<td>0.928</td>
<td>21.4</td>
<td>0.896</td>
<td>0.890</td>
</tr>
<tr>
<td>(n = 26 x n = 39)</td>
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</table>

absence of SN hyperechogenicity, which occurred in two cases, was identified only in patients with a final clinical diagnosis of AP. This finding supports previous reports claiming that the combination of LN without SN hyperechogenicity is a good predictor of an AP diagnosis\(^1\). In addition, also probably due to the sample size, the combination of V3 enlargement and LN hyperechogenicity (applied here with the intent to predict a diagnosis of PD or AP) showed low sensitivity and high specificity, which is inconsistent with published data, showing that the association of LN+ and V3 enlargement has a sensitivity and specificity of 77% in distinguishing between AP and PD\(^2,3\).

Regarding the weaknesses of the method, patients with corticobasal syndrome and Lewy body dementia, two other relevant PD mimics, also typically show SN hyperechogenicity; thus, TCS is not a useful tool for differentiating among diagnoses of PD, Lewy body dementia, and corticobasal syndrome\(^4\). Nevertheless, some authors consider PD and Lewy body dementia as two entities on the same spectrum of Lewy body disorders\(^5\). Furthermore, corticobasal syndrome has clinical features (e.g., myoclonus, asymmetric dystonia, apraxia, and alien hand phenomenon) that are distinguishable from those of PD solely on a clinical basis, which minimizes the issue from this limitation of TCS. The clinical use of TCS might also be limited by the need for operator experience and training in addition to the technical difficulties in imaging through the temporal window. When trained operators use the correct technique, the proportion of inaccessible bone windows does not exceed 20%\(^6\). In our sample, this proportion was 16% (n = 20).

Other neuroimaging methods, such as MRI (nigro-striatal and neuromelanin imaging, DTI, and SWI), dopamine transporter brain SPECT, MiBG cardiac SPECT or\(^7\) F-DOPA PET, are expensive and not readily available and may involve exposure to radiation\(^8\). The absence of an accurate, economical and practical method (or biomarker) for early confirmation of a PD diagnosis is a current medical issue; meaning that a significant proportion of patients with early-stage PD are not diagnosed until specific milestones of the natural history of the disease are reached\(^9\).

We recognize that the present study is a retrospective cohort study with a small sample size and relatively short follow-up time. Despite these limitations, the strength of the association between SN hyperechogenicity and the prediction of the clinical diagnosis of PD suggests that TCS is an ancillary technique that can be used, particularly in countries with limited resources. This was the first Brazilian study to use TCS to distinguish PD from AP and ET in a clinical practice scenario. In addition to the other case series of PD patients evaluated with TCS in Brazil\(^10,11\), our study highlights TCS as a nonradiation-dependent, accessible and safe diagnostic tool that can be utilized in Brazilian reality.

In conclusion, our results reinforce the previously-reported power of TCS in hastening the diagnosis of PD. This retrospective cohort study suggests that TCS is suitable for use in a real-world clinical scenario and provides supporting evidence for the use of TCS as a complementary diagnostic tool in the initial neurological evaluation of patients who present with bradykinesia, rigidity, and tremor\(^12\).

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### References


