Optimizing olfactory testing for the diagnosis of Parkinson’s disease: item analysis of the university of Pennsylvania smell identification test

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The 40-item University of Pennsylvania Smell Identification Test (UPSIT) is an effective instrument to detect olfactory dysfunction in Parkinson’s disease (PD). It is not clear, however, whether tests of this length are necessary to detect such dysfunction. Several studies have suggested that detection of certain odors is selectively compromised in PD, and that a test comprised of these odors could be shorter and more specific for this purpose. Therefore, we attempted to identify a subset of UPSIT odors that distinguish PD from controls with similar or improved test characteristics compared to the full test. The discriminatory power of each odor was examined using UPSIT data from a discovery cohort of 314 PD patients and 314 matched controls and ranked using multiple methods (including odds ratios, regression coefficients and discriminant analysis). To validate optimally discriminant subsets, we calculated test characteristics using data from two independent cohorts (totaling 306 PD and 343 controls). In the discovery cohort, multiple novel 12-item subsets (and the previously described Brief Smell Identification Test-B) performed similarly or improved upon the UPSIT and were better than 12 random items. However, in validation studies from independent cohorts, multiple subsets retained test characteristics similar to the full UPSIT, but did not outperform 12 random items. Differential discriminatory power of individual items is not conserved across independent cohorts arguing against selective hyposmia in PD. However, multiple 12-item subsets performed as well as the full UPSIT. These subsets could form the basis for shorter olfactory tests in the clinical evaluation of Parkinsonism.

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INTRODUCTION

Olfactory impairment is a common finding in Parkinson’s disease (PD), with estimates of prevalence ranging from 50% to more than 90%.1–6 Neurons of the olfactory system are among the first to display PD-related Lewy pathology and clinical anosmia or hyposmia may be detected years before motor symptoms present, suggesting that olfactory impairment may be one of the earliest manifestations of synucleinopathy.7–9 Whether or not such pathology causes olfactory dysfunction is unknown, as other explanations for the early deficits are possible.10 The high prevalence, persistence throughout disease, and ease of olfactory testing has fostered interest in the use of olfaction as a biomarker for early diagnostic strategies, differential diagnosis and prediction of clinical outcomes of PD and related diseases.11

Numerous tests have been used to measure olfactory function in PD with odor identification tests being the most common.12–15 Among the best-characterized and robust of such tests is the University of Pennsylvania Smell Identification Test (UPSIT).16 The UPSIT is comprised of four booklets, each of which contains 10 pages. An odorized “scratch & sniff” label is present on each page of each booklet. The subject scratches the label and then indicates which of four response alternatives best matches the perceived smell. The UPSIT is a robust measure of olfactory dysfunction in PD and has been described in numerous studies.17 However, use of the UPSIT (and other well-characterized methods such as “Sniffin Sticks”18) can be limited by difficulty of incorporating such a test into routine clinical encounter. Shorter tests would seem to be preferable both from the perspective of the patient and the neurologist, particularly within a busy clinical setting.

Shorter tests have indeed been developed, although, as noted in the discussion, there is a trade-off between test length, sensitivity, and reliability. Among such tests is the 12-item Brief Smell Identification Test (B-SIT),19,20 whose test items, derived from the UPSIT, were designed to be cross-cultural in familiarity. This test has been used to assess the prevalence of, or conversion to, such neurodegenerative diseases as Alzheimer’s disease (AD)21,22 and PD23 and has several parallel forms. These forms include odors and response alternatives potentially more sensitive to specific neurodegenerative diseases [e.g., B-SIT Version A for AD based upon24 and B-SIT Version B for PD based upon25]. Numerous other brief screening tests also have been developed, including

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Table 1. Examples of currently available olfactory tests used in PD and previously proposed discriminant subsets of odors

| Test/Author | # Odors | Comment | Ref |
|-------------|---------|---------|-----|
| Scratch and Sniff-based | | | |
| UPSIT | 40 | Odor identification. Used in >100 PD studies | 50 |
| B-SIT | 12 | Designed to be shorter and cross-culturally valid. Not intended to be PD specific | 19 |
| B-SIT-B | 12 | Based on the BSIT. Modified with the intention to be more specific for PD | 25 |
| Double | 5 | Gasoline, banana, pineapple, smoke, cinnamon identified 82% of PD cases correctly | 33 |
| Pocket Smell Test | 3 | Lemon, lilac, smoke. Not intended to be PD-specific | 26 |
| Bohnen | 3 | Banana, licorice, dill pickle. 75% accurate in identifying PD. Better correlated with dopamine transporter imaging than total UPSIT score | 33 |
| Hawkes | 2 | Pizza, wintergreen. 90% sensitivity 86% specificity for PD | 34 |
| Odor pen-based | | | |
| Sniffin' Sticks | 16 | Odor identification (modules for threshold and discrimination as well). Well-characterized in PD | 18 |
| Mahlknecht | 8 | Licorice, anise, mint, cinnamon, banana, pineapple, rose, coffee. 84% sensitivity, 88% specificity for PD | 29 |
| Casjens | 3 | Coffee, peppermint, anise. Similar classification for rate for PD compared to using 16 odors | 36 |
| Hummel | 3 | Cloves, coffee, rose. 96% sensitivity and 66% specificity for olfactory dysfunction in the general population. Not intended to be PD-specific | 30 |

Table 2. Different sets of odors distinguish between PD and control subjects

| Test | Items | Control mean (SD) | PD mean (SD) | p | AUC (95% CI) | Sen | Spe | Cut |
|------|-------|------------------|--------------|---|--------------|-----|-----|-----|
| UPSIT | 40 | 28 (8.7) | 19 (7.2) | <0.001 | 0.78 (0.74–0.82) | 0.84 | 0.66 | 27 |
| B-SIT | 12 | 8.7 (2.7) | 5.8 (2.6) | <0.001 | 0.78 (0.74–0.82) | 0.85 | 0.62 | 9 |
| B-SIT-B | 12 | 8.3 (3.0) | 5.0 (2.3) | <0.001 | 0.80 (0.76–0.83) | 0.86 | 0.67 | 8 |
| Double | 5 | 3.5 (1.3) | 2.2 (1.4) | <0.001 | 0.75 (0.71–0.79) | 0.79 | 0.58 | 4 |
| Bohnen | 3 | 2.0 (1.0) | 1.0 (0.85) | <0.001 | 0.75 (0.71–0.79) | 0.73 | 0.70 | 2 |
| PST | 3 | 2.2 (0.89) | 1.6 (1.0) | <0.001 | 0.69 (0.65–0.74) | 0.78 | 0.54 | 3 |
| Hawkes | 2 | 1.3 (0.75) | 0.85 (0.75) | <0.001 | 0.67 (0.63–0.71) | 0.79 | 0.50 | 2 |

Data are mean(SD) of the number of correctly identified odors. Area under the receiver-operator characteristic curve (AUC), sensitivity (Sen) and specificity (Spe). Cut = cut-off number of correct answers used for point sensitivity and specificity.

Results

One of the problems with using olfactory tests in PD is that the results can vary widely across studies. This can be due to a variety of factors such as the use of different test materials, the presence of head trauma or aging, or the specific odors used. However, as summarized in Table 1, the odors used in these tests have been found to be effective in distinguishing PD patients from controls.

In addition to the development of briefer tests is the question as to whether a pattern of smell loss can be identified that is more specific to PD relative to aging or other disorders that impact smell function. Double and colleagues identified a set of 5 B-SIT items that correctly differentiated 82% of PD cases (ref. 33, and an early study by Hawkes suggested that 2 UPSIT items alone could effectively distinguish PD patients from controls (ref. 34). Bohnen and colleagues identified three odors that were 75% accurate in differentiating PD from controls and were better correlated with dopamine transporter imaging than total UPSIT scores. Other studies using Sniffin' Sticks have similarly proposed odors that are selectively affected in PD compared to other causes of hyposmia including head trauma or aging (ref. 29,30). However, as summarized in Table 1, the putative “PD-specific” items vary widely across studies, raising questions about their reliability and validity in the wider PD population. Other studies have found no such selectivity (ref. 6,37). Whether there is a selective pattern of hyposmia in PD that can be observed across different cohorts is an unanswered question that has important implications for the development of shorter, more sensitive and specific assays.

The objective of this study was to determine whether a shorter version of the UPSIT could be developed that retained or improved the sensitivity and specificity in detecting hyposmia in PD. Our approach was to comprehensively analyze the discriminatory power of individual UPSIT items using a variety of statistical methods to identify subsets of odors that robustly distinguish PD patients from controls. We first derived candidate subsets in a large matched discovery cohort and then examined their performance in two independent populations of PD patients and controls.

| Table 2. |
|----------|
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| Items | Control mean (SD) | PD mean (SD) | p | AUC (95% CI) | Sen | Spe | Cut |
| UPSIT | 40 | 28 (8.7) | 19 (7.2) | <0.001 | 0.78 (0.74–0.82) | 0.84 | 0.66 | 27 |
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| B-SIT-B | 12 | 8.3 (3.0) | 5.0 (2.3) | <0.001 | 0.80 (0.76–0.83) | 0.86 | 0.67 | 8 |
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| Hawkes | 2 | 1.3 (0.75) | 0.85 (0.75) | <0.001 | 0.67 (0.63–0.71) | 0.79 | 0.50 | 2 |

Data are mean(SD) of the number of correctly identified odors. Area under the receiver-operator characteristic curve (AUC), sensitivity (Sen) and specificity (Spe). Cut = cut-off number of correct answers used for point sensitivity and specificity.

University of Pennsylvania smell identification test, B-SIT brief smell identification test, B-SIT-B brief smell identification test, version B.
items (smoke, soap, licorice, bubblegum) appeared on all lists. Eleven of the items appeared on at least 3 of the 12-item lists. Four top 12 of at least 1 of the 4 initial ranking approaches (Table 3). The worst 12 items using the difference method (Worst) are shown for comparison. To highlight odors that were found as highly discriminant for differentiating PD from controls using multiple different ranking methods, items appearing on the all of the “Difference,” “Odds Ratio,” “Discriminant;” “Regression” and “Combined” lists are shown as bold. 12 item lists were used to facilitate comparison with existing, commercially available smell tests such as the B-SIT.

Development of novel UPSIT subsets for the detection of hyposmia in PD

We attempted to identify novel subsets of UPSIT odors that might outperform the full test using different statistical ranking strategies (see methods for full details). This approach narrowed the 40 UPSIT items to a total of 22 unique items that were in the top 12 of at least 1 of the 4 initial ranking approaches (Table 3). Eleven of the items appeared on at least 3 of the 12-item lists. Four items (smoke, soap, licorice, bubblegum) appeared on all lists. Multiple 12-item subsets had test characteristics similar to the full UPSIT (Table 4). Some, such as the 12-item Combined list, had slightly better test characteristics compared to full UPSIT (Sens/Spec, 0.84/0.77 vs. UPSIT 0.84/0.71, Table 4). Relatively poorer test characteristics were observed for 12-item subsets derived at random (0.78/0.65) or from the worst ranking items (0.72/0.53, Table 4) in the discovery cohort. Further shortening the top-12 items lead to steady declines in AUC and/or the optimal combination of sensitivity and specificity as items beyond the top 11 were removed (Table 4).

“PD-specific” subscales derived in one population do not retain discriminatory power across independent cohorts

We examined the performance of our putative PD-specific subsets with individual item UPSIT data from two independently derived validation cohorts. As in the discovery cohort, test characteristics including sensitivity, specificity and AUC for multiple 12-item subsets were similar to those for the UPSIT indicating that smaller subscales can maintain comparable discriminatory power (Figure). However, when tested in the independent samples, the most highly discriminatory subsets from the discovery cohort did not perform better than a random subset or, in fact, the worst ranking 12 items derived from the discovery cohort. For example, AUCs for the Combined-12 subset, full UPSIT and Worst-12 subset calculated with data from the discovery cohort were 0.83 (95% CI 0.80–0.87), 0.78 (95% CI 0.74–0.82), and 0.66 (95% CI 0.62–0.74) respectively (Tables 2, 3). Using data from the Barts cohort, however, these values were essentially identical to one another (AUCs: Combined-12 = 0.85, UPSIT = 0.87, Worst-12 = 0.86, Figure). Similar results were observed using data from the UCL cohort (AUCs: Combined-12 = 0.82, UPSIT = 0.90, Worst-12 = 0.87, Figure).

Effect of age and gender on olfactory test performance

As age and sex are important determinants of olfactory function, we examined the test characteristics of the UPSIT, B-SIT-B and 12 UPSIT items (“Combined” list) we defined as most highly discriminatory in the discovery cohort (Table 5) as a function of age and sex using data from all three cohorts (N = 1279). We found that although the test AUCs were fairly similar in men and women, higher cut-off values were required for optimal sensitivity/specificity in women. Additionally, the tests effectively distinguished between PD and controls in all age groups but, generally, we observed higher AUCs in subjects less than 74 years old (Table 5). The pattern of age/sex influence was similar across the different tests.

DISCUSSION

Detecting anosmia or hyposmia is of significant interest for early identification and differential diagnosis of PD and related disorders. Although the 40-item UPSIT has been found to be an effective instrument to detect anosmia or hyposmia in PD, it is not clear whether tests employing fewer UPSIT items are equally useful in detecting such olfactory dysfunction. Several studies have suggested that certain odors are selectively compromised in PD and, that a test comprised of these odors could be shorter, easier to administer, and more specific for this purpose. However, little uniformity exists across studies. Some of the candidate subsets identified using “scratch and sniff” tests (UPSIT, B-SIT versions) include gasoline, banana, pineapple, smoke and cinnamon, licorice, banana, dill pickle and wintergreen and pizza (Table 1). Studies using Sniffin Sticks have similarly

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### Table 3. Putative subsets highly discriminant of UPSIT items

| Rank | Difference | Odds ratio | Discriminant | Regression | Combined | Worst |
|------|------------|------------|--------------|------------|----------|-------|
| 1    | Smoke      | Smoke      | Lime         | Grass      | Smoke    | Rootbeer |
| 2    | Motor oil  | Grass      | Turpentine   | Lime       | Grass    | Watermelon |
| 3    | Soap       | Licorice   | Smoke        | Banana     | Turpentine | Leather |
| 4    | Gasoline   | Lemon      | Banana       | Turpentine | Smoke    | Onion |
| 5    | Paint thinner | Motor oil | Bubbles | Smoke | Lime | Gingerbread |
| 6    | Peanut     | Turpentine | Grape        | Bubblegum  | Bubblegum| Peach |
| 7    | Grass      | Dill pickle | Soap | Cherry | Motor oil | Cheddar cheese |
| 8    | Lemon      | Bubbles | Dill pickle | Grape | Banana | Cinnamon |
| 9    | Wintergreen | Soap       | Licorice   | Soap       | Licorice | Chocolate |
| 10   | Grape      | Gasoline   | Pine         | Mint       | Grape    | Mint |
| 11   | Licorice   | Lime       | Cedar        | Cinnamon   | Lemon    | Cherry |
| 12   | Bubbles    | Paint thinner | Gasoline | Licorice | Gasoline | Strawberry |

Summary of subsets of UPSIT items that were identified as highly discriminant for differentiating PD from control subjects in the discovery cohort. Listed are the top 12 most discriminant UPSIT items ranked by five different methods: 1) the absolute difference in percentage of PD and control subjects answering incorrectly (Difference), 2) odds ratio, 3) discriminant function analysis (Discriminant), 4) logistic regression (Regression), 5) a weighted average combining the first four methods (Combined). The worst 12 items using the difference method (Worst) are shown for comparison. To highlight odors that were found as highly discriminant for differentiating PD from controls using multiple different ranking methods, items appearing on all of the “Difference,” “Odds Ratio,” “Discriminant,” “Regression” and “Combined” lists are shown as bold.

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Identified by Double et al., odors from the 3-item Pocket Smell Test, 3 items previously identified by Bohnen et al. and two items suggested by Hawkes were significantly lower in PD patients compared to controls (Table 2). Sensitivity and specificity were similar between the UPSIT and each of the 12 item tests (Table 2). The 5 item scale based on odors from Double et al., 3-item subsets and the 2 items proposed by Hawkes had lower sensitivity and/or specificity compared to the full test (Table 2).
Table 4. Test characteristics of putative subsets of highly discriminant UPSIT items from the discovery cohort

| Test     | Items | AUC (95% CI) | Sensitivity | Specificity | Cut-off |
|----------|-------|--------------|-------------|-------------|---------|
| UPSIT    | 40    | 0.78 (0.74–0.82) | 0.84 | 0.66 | 27 |
| Difference | 12  | 0.82 (0.78–0.85) | 0.77 | 0.74 | 8 |
| Odds Ratio | 12  | 0.81 (0.79–0.86) | 0.84 | 0.67 | 8 |
| Regression | 12  | 0.80 (0.76–0.83) | 0.82 | 0.68 | 8 |
| Discriminant | 12  | 0.81 (0.78–0.75) | 0.83 | 0.69 | 8 |
| Combined  | 12  | 0.83 (0.80–0.86) | 0.84 | 0.71 | 8 |
| Random   | 12  | 0.76 (0.72–0.79) | 0.78 | 0.65 | 8 |
| Worst    | 12  | 0.65 (0.61–0.70) | 0.72 | 0.53 | 9 |

Items from the "Combined" List
- 11 item list, AUC=0.80, Sensitivity=0.70, Specificity=0.65
- 10 item list, AUC=0.76, Sensitivity=0.76, Specificity=0.82
- 9 item list, AUC=0.70, Sensitivity=0.77, Specificity=0.70
- 8 item list, AUC=0.75, Sensitivity=0.74, Specificity=0.76
- 7 item list, AUC=0.85, Sensitivity=0.65, Specificity=0.79
- 6 item list, AUC=0.72, Sensitivity=0.67, Specificity=0.82
- 5 item list, AUC=0.74, Sensitivity=0.74, Specificity=0.70
- 4 item list, AUC=0.71, Sensitivity=0.80, Specificity=0.71
- 3 item list, AUC=0.74, Sensitivity=0.74, Specificity=0.70
- 2 item list, AUC=0.70, Sensitivity=0.84, Specificity=0.57

Data are area under the receiver-operator characteristic curve (AUC), sensitivity and specificity for differentiating PD from control subjects in the discovery cohort. Cut-off = number of correct answers used to determine the point sensitivity and specificity. The subsets of highly discriminant items were determined by ranking odors using five different methods: 1) the absolute difference in percentage of PD and control subjects answering incorrectly (difference), 2) odds ratio, 3) discriminant analysis (discriminant), 4) logistic regression (regression), 5) a weighted average combining the first four methods (combined). For comparison, test characteristics for 12 random items and the worst 12 items using the difference method (worst) are shown. 12 item lists were used to facilitate comparison with existing, commercially available smell tests such as the B-SIT. In the second half of the Table, test characteristics for subsets containing decreasing numbers of the 12 most highly discriminatory items from the discovery cohort are shown.

Table 4. Olfactory tests for PD: JF Morley et al.

Table 5. Effect of age and sex on olfactory test characteristics

| Test        | Age | AUC (95% CI) | Cut-off | Sensitivity | Specificity |
|-------------|-----|--------------|---------|-------------|-------------|
| UPSIT       | <63 | 0.86 (0.82–0.90) | 0.79 | 0.74 | 0.83 |
|             | 63–73 | 0.88 (0.84–0.92) | 0.88 | 0.83 | 0.79 |
|             | >73  | 0.76 (0.70–0.82) | 0.82 | 0.71 | 0.83 |
| BSIT-B      | <63 | 0.85 (0.81–0.90) | 0.80 | 0.74 | 0.77 |
|             | 63–73 | 0.87 (0.83–0.91) | 0.88 | 0.82 | 0.85 |
|             | >73  | 0.77 (0.71–0.83) | 0.82 | 0.71 | 0.85 |
| Combined    | <63 | 0.85 (0.81–0.89) | 0.76 | 0.70 | 0.74 |
|             | 63–73 | 0.86 (0.82–0.91) | 0.86 | 0.80 | 0.77 |
|             | >73  | 0.78 (0.72–0.84) | 0.82 | 0.72 | 0.72 |

Data are area under the receiver-operator characteristic curve (AUC), sensitivity and specificity for differentiating PD from control subjects in the discovery cohort. Cut-off = number of correct answers used to determine the point sensitivity and specificity. Subjects from all three cohorts (N = 1279) were divided by sex and age tertile (<63 years old, 63–73 years old and >73 years old). UPSIT University of Pennsylvania smell identification test, BSIT-B brief smell identification test, version B. Combined: Top 12 items found most highly discriminatory in the discovery cohort.
patients involved in the study were not autopsy verified. The study has several limitations that are important to consider. Most of these are related to the use of multiple independent validation cohorts. We feel that this is a strength, this study is not conclusive. There are likely other odor sets that might be specific to PD and that could be discovered using the full 40-item test in a discovery cohort but not in independent replication cohorts. Data are area under the receiver-operator characteristic curve (error bars represent the 95% confidence interval) for the 40 UPSIT items (red), or items from the BSIT-B (green), “combined” subset (blue), random 12 items (white) or worst 12 items (from the training cohort) when tested using data from the training cohort (Penn) or two independent cohorts (Barts, UCL).

Fig. 1   Novel UPSIT subsets do no retain discriminatory power across independent cohorts. Data are area under the receiver-operator characteristic curve (error bars represent the 95% confidence interval) for the 40 UPSIT items (red), or items from the BSIT-B (green), “combined” subset (blue), random 12 items (white) or worst 12 items (from the training cohort) when tested using data from the training cohort (Penn) or two independent cohorts (Barts, UCL).

sets of receptor cells. Even if some subset of receptors were damaged specifically by PD, the gestalt of a given smell, like the perception of visual objects, can likely resist the loss of some segments of the olfactory "object" and still retain identification ability via feature-detection processes. Third, the search for odorants specific to PD is further complicated by the fact that most if not all of the odorants employed in the extant olfactory tests are comprised of multiple chemicals. Until there is a better understanding of the relative distribution numbers of the ~400 classes of receptor types within the epithelium and the nature and range of ligands that activate each receptor type, finding sets of odorants that might be specifically damaged by PD or any other disease is unlikely. Finally, the quest is further confounded by attempting to compare results across studies using different tests with seemingly the same “odors”. Even if the qualitative “odor” from one test appears to be the same qualitative “odor” as that from another, different chemicals and combinations of chemicals can make up the same “odor”. In other words, different odorants or combinations of odorants often are being compared.

While the large number of subjects in the discovery cohort and use of multiple independent validation cohorts are strengths, this study has several limitations that are important to consider. Most patients involved in the study were not autopsy verified so that some of the PD subjects likely had non-Lewy body Parkinsonism and some of the controls may have had pre-motor PD other than PD. Our analysis of individual items and novel combinations was retrospective using existing UPSIT data and, therefore, cannot account for item ordering or the effect of distractor choices that would be present if the proposed UPSIT subsets had been presented together as independent tests. Smoking history was not available for all subjects, but smoking has a relatively small impact on olfactory function, compared to factors such as age, sex or the presence underlying neurologic disease, such as PD (ref.44). Indeed, age and sex are significant determinants of olfactory function such that optimal UPSIT cut-off scores can differ between men and women or among different age groups.45 Similarly, we found that higher cut-off values were required for optimal sensitivity/specificity in women, reflecting generally better olfactory performance compared to men. The tests effectively distinguished between PD and controls in all age groups but performed best in subjects less than 74 years old.

However, the influence of age and sex were similar using the full-length UPSIT or subsets of UPSIT items (Table 5).

Finally, this study examined multiple international cohorts for discovery and validation but only included subjects from the US and UK. Cultural factors influencing recognition of certain odors are known to affect performance on olfactory identification tests in other populations, possibly limiting generalizability of these results to other cultures.40 Similarly, cultural heterogeneity between the discovery and validation cohorts could explain some of the variable performance of different subsets of UPSIT items between the cohorts (Fig. 1).

While our results, along with those of earlier studies, argue against selective anosmia or hyposmia in PD, they do suggest that shorter versions of the UPSIT or Sniffin’ Sticks retain much of the discriminatory power of the full tests for detecting olfactory dysfunction in PD. The decision to employ a short or long test for a given clinical or research purpose depends on a number of factors, including the setting of the administration, proposed indication, and pre-test probability of PD in the population studied. As discussed in detail, shorter tests may maintain suitable test characteristics for a binary outcome (diagnosis). However, longer tests are more sensitive to subtle alterations in function and allow for distinctions between degrees of dysfunction, which can be critical for counseling patients regarding prognosis, including patients with non-neurodegenerative disorders such as head trauma.46 Longer tests also allow for the detection of malingering on the basis of improbable forced-choice responding,47 which cannot be discerned from shorter tests, and are clearly more reliable than shorter tests.48 We found that decreasing even the most discriminatory set of items to fewer than 11 odds resulted in steadily decreasing test performance. This can have an impact when small samples are being tested or when individual patients or subjects are being assessed. It must be kept in mind, of course, that while olfactory testing can be a very sensitive aid in diagnosing PD, e.g., in differentiating between PD from progressive supranuclear palsy and essential tremor, it is not specific to PD.49

Our findings that 12-item UPSIT subsets performed better that the full 40-item test in a discovery cohort but not in independent replication cohorts has several practical implications for the use of olfactory tests for PD. First, 12-item tests are sufficient and may save time and cost compared to the full UPSIT. Second, attempts to discover new “PD-specific” odor sets may be ill-advised as they can be defined by chance in any cohort but are unlikely to generalize to the broader PD population. Further, we found that AUC, sensitivity and specificity declined as items were removed from the 12-item subsets suggesting that significantly shorter tests would lack sufficient diagnostic utility. Additionally, any such shorter or “PD-specific” test would lack normative data for categorizing individual patients and would need prospective validation in new cohorts. Overall, the balance of evidence suggests that shorter versions of the UPSIT—particularly the currently available B-SIT-B—should be employed with confidence to allow decreased time of administration and cost of olfactory assessment in a variety of clinical and research applications for the evaluation of Parkinsonism.

METHODS

Subjects and olfactory assessment

For the initial (discovery cohort) phase of the UPSIT item analysis, we examined individual UPSIT item results from a convenience sample of PD patients (N = 314) and age-matched controls (N = 314) that had been administered in several protocols at the Michael J. Crescenz VA Medical Center in Philadelphia and the University of Pennsylvania. The mean (SD) age in each group was 67.4 (10.0) years and each was comprised of 83% males and were 94% Caucasian. Among PD patients, the median (interquartile range) Hoehn and Yahr stage and mean (SD) UPDRS motor score was 2.2 (1.7), and mean (SD) disease duration was 11.8 (6.8) years. Subjects were recruited from the outpatient clinic and comprised 70% of newly diagnosed patients (N = 222) and 30% of patients previously treated for PD (N = 92). Patients were enrolled sequentially until 314 PD and 314 age-matched control subjects had been recruited. Controls were recruited from the community, as well as family members of PD patients, and were matched by age, gender, and education. All participants were interviewed in English and written informed consent was obtained from all before any testing. Among the 314 PD and age-matched controls included in the analysis, 9% were Hispanic/Latino and 7% had a body mass index (BMI) > 30. No significant differences were found in the distribution of other characteristics among control subjects as a result of matching, and control subjects scored within the normal range on all olfactory tests.

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scores were 2(2–3) and 22 (10.1), respectively. In an attempt to validate the performance of putative PD-specific UPSIT subsets, we used individual item data from two independent validation cohorts of PD patients and control subjects derived at University College, London (UCL Cohort) and Barts & The London School of Medicine and Dentistry (Barts Cohort). The Barts cohort was comprised of 176 PD patients with a mean age 60 (9.8) years and 177 control subjects with a mean age of 62 (10.7) years (p = 0.15). Subjects in the Penn cohort were only 6% non-white. Race data were not collected for all of the Barts/UCL subjects but they were largely drawn from the Oldchurch /Queens and UCL hospital patients. The vast majority were middle class Caucasian British. There were 167 PD subjects (mean age = 63 (9.9)) and 130 controls (mean age = 65(9.5)) in the UCL cohort. Most subjects were screened extensively for nasal disease. However, some subjects were particularly controls that were tested in community settings such as malls or state fairs, did not undergo rigorous screening, though subjects with clear active rhinitis of any etiology were not included. All studies from which UPSIT results were analyzed were approved by Institutional Review Boards (IRB) at Crenszcz VA Medical Center, University of Pennsylvania, Barts and The London School of Medicine and Dentistry and University College London. Methods were performed in accordance with relevant regulations and guidelines. Informed consent was obtained from patients before participation in protocols.

Statistical analysis
Individual responses to each of the 40 items were recorded as correct or incorrect. Discriminatory power of individual odors to differentiate between PD patients and control subjects was tested using several statistical approaches. First, individual odors were ranked by the difference between the percentages of PD patients versus controls answering incorrectly (Difference). A complimentary approach ordered odors by odds ratio of PD versus controls grouping for each item (Odds Ratio). The third method used discriminant function analysis, a method based on ANOVA that generates models incorporating all items into one or more weighted functions to come up with two sets, one that best discriminated PD versus controls (Discriminant) and one that least discriminated PD versus controls (Worst). We also used logistic regression to identify items that best explained variation in outcome using diagnosis of PD versus controls as the dependent variable and ranking individual odors by the calculated for candidate subsets. Cut-offs for point sensitivities and specificities in Parkinson’s disease.

ADDITIONAL INFORMATION
Competing interests: R.L.D. is President and major shareholder in Sensonics International, the manufacturer of smell and taste tests, some of which were assessed in this study. The remaining authors declare no competing financial interests.

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AUTHOR CONTRIBUTIONS
J.F.M.: conception and design, statistical analysis, drafting of the manuscript and revision. Guarantor. A.C.: design and execution of statistical analysis, revision of the manuscript. L.S.M.: data acquisition, statistical analysis, revision of the manuscript. A.J.L.: data acquisition, revision of the manuscript. D.R.W.: data acquisition, revision of the manuscript. R.K.: data acquisition, revision of the manuscript. J.A.: data acquisition, revision of the manuscript. D.W.: data acquisition, revision of the manuscript. C.H.: data acquisition, revision of the manuscript. R.L.D.: data acquisition, revision of the manuscript. J.E.D.: conception and design, revision of the manuscript.

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