Audiovisual Processing is Abnormal in Parkinson's Disease and Correlates with Freezing of Gait and Disease Duration

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8 Abstract.

- Background: Sensory and perceptual disturbances progress with disease duration in Parkinson's disease (PD) and probably contribute to motor deficits such as bradykinesia and gait disturbances, including freezing of gait (FOG). Simple reaction time tests are ideal to explore sensory processing, as they require little cognitive processing. Multisensory integration is the ability of
- the brain to integrate sensory information from multiple modalities into a single coherent percept, which is crucial for complex motor tasks such as gait.
- **Objectives:** The aims of this study were to: 1. Assess differences in unisensory (auditory and visual) and multisensory processing speed in people with PD and age-matched healthy controls. 2. Compare *relative* differences in unisensory processing in people
- with PD with disease duration and freezing of gait status taking into account the motor delays, which are invariably present in
 PD. 3. Compare relative differences in multisensory (audiovisual) processing between the PD cohort and age-matched controls.
 Methods: 39 people with PD (23 with FOG) and 17 age-matched healthy controls performed a reaction time task in response
- Methods: 39 people with PD (23 with FOG) and 17 age-matched healthy controls performed a reaction time task in resp to unisensory (auditory-alone, visual-alone) and multisensory (audiovisual) stimuli.
- Results: The PD group were significantly slower than controls for all conditions compared with healthy controls but auditory reaction times were significantly faster than visual for the PD group only. These relative unisensory differences are correlated with disease duration and divide the PD group by FOG status, but these factors are co-dependent. Although multisensory facilitation occurs in PD, it is significantly less enhanced than in healthy controls.
- Conclusion: There are significant unisensory and multisensory processing abnormalities in PD. The relative differences in unisensory processing are specific to PD progression, providing a link between these sensory abnormalities and a motor feature
- of PD. Sensory disturbances have previously been postulated to be central to FOG but this is the first study to predict audiovisual
- processing abnormalities using FOG status. The multisensory processing abnormalities are independent of disease duration and
 FOG status and may be a potential biomarker for the disease.
- 29 Keywords: Parkinson's disease, sensory processing, multisensory, auditory, visual

INTRODUCTION

Sensory and perceptual disturbances are common in Parkinson's disease (PD) [1–3]. Subtle deficits of the sensory system, often not detected by routine examination, occur in people with Parkinson's disease (PwP). From simple anosmia and impaired kinesthetic perception, to more complex visual hallucinations and spatiotemporal perceptual abnormalities, altered 37

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sensory processing is found across multiple modalities 38 [4-8]. Of note, integration of multiple environmen-39 tal sensory inputs is crucial for a refined but complex 40 goal-directed motor output (e.g. locomotion through 41 a crowded environment). There is increasing evidence 42 that these sensory deficits contribute to the pathophys-43 iology of some of the abnormal motor features of 44 PD [9-11], including freezing of gait (FOG), where 45 patients feel as though their feet are momentarily 46 glued to the floor [12], and which is closely associated 47 with falls and nursing home placement [13]. Although 48 the underlying pathophysiology FOG is incompletely 49 understood, sensory mechanisms are likely to be core 50 factors underlying this motor symptom [14]. 51

There are many studies quantifying single modal-52 ity (unisensory) deficits in PD. Simple reaction times 53 are helpful when exploring sensory responses, as they 54 require little cognitive processing (interpretation can 55 be difficult in a patient population where cognitive 56 impairment is common). Simple reaction times to 57 auditory and visual stimuli are delayed in PwP as com-58 pared to healthy controls [15-22]. However, motor 59 output in response to sensory stimuli requires both 60 sensory processing and sensorimotor integration. Sim-61 ple unisensory reaction times are, therefore, delayed 62 in PwP because of bradykinesia, and do not solely 63 assess sensory differences in these patients, as the 64 response is a combination of motor and sensory pro-65 cessing pathways. Quantitative assessment of sensory 66 processing speeds therefore requires examination of 67 relative differences in response times to stimuli, sep-68 arate from common motor output time. Nevertheless, 69 premotor delays in processing have been shown in PwP 70 via movement-related potentials [21, 23] and auditory, 71 visual and somatosensory evoked potentials [24-27], 72 implying that unisensory processing is altered in PD, 73 independent of motor integration. 74

75 Multisensory integration is the brain's ability to integrate sensory information from multiple modalities 76 into a single coherent percept, leading to increased 77 speed and accuracy of response [28]. When reaction 78 times to multisensory stimuli are compared to individ-79 ual component unisensory stimuli, the responses are 80 significantly faster than would be predicted based on 81 the unisensory reaction times. By comparing relative 82 response times to unisensory and multisensory stim-83 uli, quantitative assessment of multisensory integration 84 can be performed, while controlling for variable motor 85 86 response times in PD.

Multisensory integration is enhanced in healthy
 elderly populations [29] but it is unknown if this
 multisensory facilitation is present in PwP. Inefficient

multisensory integration is linked with falls in older adults, highlighting the importance of controlled multisensory processing in balance and locomotor control [30]. Given that locomotion is highly multisensory task and that progressive gait impairment frequently occurs in PD, abnormal multisensory processing may occur in PD. Single cell animal studies have highlighted the basal ganglia as an important multisensory hub [31, 32]. As PD is a basal ganglia disorder and has widespread sensory abnormalities, we hypothesized that multisensory integration is altered in PD.

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Few studies have reported multisensory abnormalities in PD [33]. The multisensory interactions between auditory and visual stimuli have not been studied in PD. We studied PwP and age-matched healthy controls performing a reaction time task in response to unisensory (auditory-alone, visual-alone) and multisensory (audiovisual) stimuli. In this study we have made efforts to limit the effect of attention by comparing relative differences between audio, visual and audiovisual response times. In this way, each participant acts as his or her own control. Thus any differences in performance represent relative differences in either processing of different modalities or shifts in modality-specific attention between groups. Given the widespread sensory abnormalities in PD, we hypothesized that multisensory integration is also altered in PwP. The reaction time task was used in order to:

- 1. Assess differences in unisensory (auditory and visual) processing speed in PwP and age-matched healthy controls.
- 2. Correlate *relative* differences in unisensory (auditory vs visual) processing in PwP with disease duration and FOG status taking into account the known motor delays in PD.
- 3. Compare relative differences in multisensory processing between PwP and age-matched controls.

METHODS

Participants

39 patients with idiopathic PD (as defined by the UK Brain Bank Criteria [34]; Modified Hoehn and Yahr stage II–IV) were recruited from the Movement Disorder Clinic at the Dublin Neurological Institute. Ethical approval was granted from the hospital ethics committee and informed consent was obtained from all participants. All patients underwent clinical and neuropsychological testing including Montreal

Cognitive Assessment (MoCA), Frontal Assessment 138 Battery (FAB) and Unified Parkinson's Disease Rating 139 Scale III (UPDRS III). FOG status was recorded for all 140 patients based on Question 1 of the New Freezing of 141 Gait Questionnaire ("Did you experience a freezing 142 episode over the past month?") [35]. All participants 143 had normal corrected vision and hearing and were 144 tested in the "on"-state. A group of 17 age-matched 145 healthy controls were recruited among hospital staff 146 and relatives of participants for comparison. The con-147 trol group had no neurological comorbidities and 148 normal cognition. 149

150 Stimuli

Participants performed a simple reaction time task 151 consisting of three stimulus conditions: "auditory" (A), 152 "visual" (V) and "audiovisual" (AV). Stimuli were pre-153 sented using Presentation software (Neurobehavioral 154 Systems, Inc., Albany CA). The auditory condition 155 consisted of a 1000-Hz tone (duration 60 msecs; 75 dB; 156 rise/fall time 5 msecs), presented from via inbuilt 157 speakers of a Dell laptop (Latitude E5530). The visual 158 condition consisted of a red disc with a diameter of 159 3.2 cm (subtending 1.5 degrees in diameter at a viewing 160 distance of 122 cm) appearing on a black background, 161 presented on the screen for 60 milliseconds. The audio-162 visual condition consisted of the auditory and visual 163 conditions presented simultaneously. 164

165 Procedure

Participants were seated in front the laptop and 166 instructed to press a button as quickly as possible when 167 they saw the red circle, or heard the tone, or saw the 168 circle and heard the tone together. The stimulus con-169 ditions were presented with equal probability and in 170 171 random order in blocks of 100 trials. Inter-stimulusinterval (ISI) varied randomly between 1000 and 3000 172 milliseconds according to a uniform (square wave) dis-173 tribution. Participants completed 3 blocks, resulting in 174 100 repetitions per stimulus condition. These meth-175 ods are also presented in detail elsewhere [36-41]. The 176 range of reaction times accepted was determined at the 177 individual participant level with the slowest cut off at 178 150 milliseconds and fastest 2.5% of trials excluded. 179

180 Statistical analysis

Data were processed and analyzed using custom MATLAB (Mathworks, Natick, MA) scripts and
SPSS 22.

Reaction time analysis

Mean reaction times for each condition were calcu-185 lated for all participants. A mixed one-way analysis of 186 variance (ANOVA), with the factors of stimulus con-187 dition (auditory-alone, visual-alone, audiovisual) and 188 group (PwP and control participants) was performed to 189 compare the reaction times of the three stimulus condi-190 tions between PwP and controls. Post-hoc comparisons 191 between the conditions were performed to test for the 192 presence of relative differences between the unisen-193 sory conditions as a well as faster reaction times in the 194 multisensory condition. In order to examine whether 195 differences in capacity for focused attention differed 196 between groups, reaction times and hit rates were cal-197 culated for the first and last blocks of trials in each 198 group. 199

Relative sensory processing and FOG status

To investigate the relationship between relative sen-201 sory processing (controlling for motor delays) and 202 FOG status, the PwP group was subdivided by Ques-203 tion 1 of the New Freezing of Gait Questionnaire, as 204 described above [35]. A mixed repeated ANOVA was 205 performed with the within-participant factor of relative 206 reaction time (auditory-visual vs audiovisual-visual vs 207 audiovisual-auditory) and between-participant factor 208 of FOG status (freezers vs non-freezers). The reaction 209 times were subtracted to account for variable motor 210 delays in PwP. In this way, the results relate to relative 211 changes in sensory processing rather than reflecting 212 slower motor responses with disease progression. The 213 Greenhouse-Geisser correction was used to adjust F-214 values and probabilities when sphericity was violated. 215 The original degrees of freedom are presented for each 216 analysis. 217

Correlation analysis of disease duration

Correlation analyses were performed on the PwP group to assess the extent to which the relative differences of reaction times for the three conditions, (auditory-visual, audiovisual-visual, audiovisualauditory), are associated with disease duration (years since symptoms onset).

Miller race model

In order to quantitatively assess the degree to which multisensory integration contributes to response times for the audiovisual condition, the Miller race model was employed [42]. Faster reaction times to the multisensory stimuli could be the result of participants responding to whichever stimulus is processed fastest, even in the absence of any interaction between the

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individual sensory stimuli. In this way, sensory pro-233 cessing could be considered a race between two 234 modalities (auditory and visual in this case) on a trial-235 by-trial basis. The race model proposed by Miller is a 236 commonly used behavioral index of multisensory inte-237 gration which takes this effect into account [36-41]. 238 According to Miller's race model, reaction times are 239 still expected to be faster in the multisensory condition 240 compared with the unisensory state. This is because 241 there are now two inputs, which can trigger a response, 242 as opposed to just one. Whichever input is fastest, trig-243 gers a response, making a faster response more likely 244 in the multisensory condition than if only a single stim-245 ulus was present. Miller's race model defines an upper 246 limit for multisensory responses in this simple linear 247 model based on the sum of the cumulative probabilities 248 of each unisensory stimulus triggering a response. If 249 the recorded multisensory reaction time is faster than 250 this upper limit then violation of the race model has 251 occurred and it must be assumed that the unisensory 252 inputs interacted during processing (i.e. multisensory 253 integration occurred). Failure to violate the race model, 254 however, does not prove that the unisensory inputs 255 did not integrate, but implies that the recorded mul-256 tisensory reaction time could be explained by simple 257 summation of unisensory probabilities. To control for 258 false positives resulting from the multiple compar-259 isons, p-values were corrected using the false discovery 260 rate (FDR). The FDR is a sequential Bonferroni-type 261 procedure. 262

263 **RESULTS**

264 Demographics

The demographic and neurocognitive data for the PD cohort (divided by FOG status) is given in Table 1. The 17 healthy control participants (10 Male) had a mean age of 66 +/- 9.7 years (range 52–80).

269 *Hit rate analysis*

Hit rates (proportion of stimuli responded to) were
consistently high across all groups (Table 2). No significant hit rate differences were found between first
and last blocks of trials for any group.

274 Reaction time

PwP were significantly slower than controls for all
conditions. Table 3 and Fig. 1 show the mean reaction times and standard deviations for each condition

Table 1 Patient Demographics by FOG status. Means shown with standard deviation in parentheses (unless median stated)

| | All PD | Freezers | Non-Freezers |
|---------------------------|------------|-------------|--------------|
| N | 39 | 23 | 16 |
| Age | 67.4 (9.8) | 68.7 (9.7) | 66.7 (10.05) |
| Gender (M:F) | 23:16 | 15:8 | 8:8 |
| H&Y stage (median) | 2.5 (0.7) | 3.0 (0.6) | 2.5 (0.3) |
| Disease Duration (years)* | 10.1 (9.4) | 14.0 (10.5) | 5.2 (4.6) |
| UPDRS | 34.1 (14) | 38 (13) | 30 (14) |
| MOCA | 24.7 (4.8) | 24.4 (3.3) | 26.3 (3.6) |
| FAB | 15.7 (3.3) | 15.4 (2.8) | 17.1 (1.5) |
| | | | |

*indicates statistically significant difference between groups. H&Y stage = Modified Hoehn & Yahr stage; UPDRS III = Unified Parkinson's Disease Rating Scale III total; MOCA = Montreal Cognitive Assessment total; FAB = Frontal Assessment Battery total; PD = Parkinson's disease.

 Table 2

 Mean hit rate and standard deviation for control group and people with Parkinson's disease (PwP) group

| Group | А | V | AV |
|--------------------------|-------------|-------------|-------------|
| PwP(N=39) | 0.94 (0.08) | 0.92 (0.09) | 0.97 (0.03) |
| Controls $(N=17)$ | 0.98 (0.05) | 0.94 (0.06) | 0.98 (0.02) |
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A = auditory, V = visual, AV = audiovisual.

| Table 3 |
|---|
| Mean and standard deviation of reaction times for control group and |
| people with Parkinson's disease (PwP) group |

| Group | А | V | AV |
|-------------------|--------------|--------------|--------------|
| PwP(N=39) | 374.1 (74.0) | 403.8 (67.6) | 325.2 (68.0) |
| Controls $(N=17)$ | 295.2 (47.9) | 315.1 (36.9) | 245.1 (29.7) |

A = auditory-alone, V = visual-alone, AV = audiovisual.

(auditory-alone, visual-alone, audiovisual) and group (PwP and control participants). The mixed repeated ANOVA revealed a significant difference between the conditions' reaction times ($F_{2,108} = 84.32$, P < 0.001) with the fastest reaction times for the audiovisual condition. The analysis revealed significant difference between groups ($F_{1,53} = 24.1$, P < 0.001) with faster reaction times for all stimulus conditions in the control participants than in the participants with PD.

To investigate the significant effect of condition (auditory, visual, audiovisual), the data were submitted to a follow-up within-group between-stimulus conditions analysis. The paired *t*-tests revealed that the reaction times in the audiovisual condition (AV) were significantly faster than the reaction times for the auditory-alone (A) and visual-alone (V) conditions in the control group (auditory-alone vs audiovisual p < 0.001; visual-alone vs audiovisual p < 0.001; visual-alo

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Fig. 1. Reaction times for the audio (blue), visual (green) and Audiovisual (red) conditions for both the people with Parkinson's disease (PWP, circles) and control participants (squares). The horizontal line and errorbars depict the mean and standard error of the mean.

the patients with PD revealed significant differences between the unisensory conditions; auditory-alone vs visual-alone (p < 0.001), while in the control participants there was no significant difference between the unisensory auditory-alone and visual-alone conditions (p = 0.26).

304 FOG status and disease duration analysis

To investigate the relationship between *relative* sensory processing (controlling for motor delays) and FOG status, the PD group was subdivided by

Question One of the New Freezing of Gait Question-308 naire [35], as described above (Table 1). A mixed 309 repeated ANOVA was performed with the within-310 participant factor of relative reaction time (A-V, 311 A-AV vs A-AV) and between-participant factor of 312 FOG status (freezers vs non-freezers). The reaction 313 times were subtracted to account for variable motor 314 delays in PwP, which allows for the analysis of 315 relative sensory reaction times, taking into account 316 variable motor delays seen in PwP. In this way, 317 the results reflect true changes in sensory process-318 ing rather than slower motor responses in freezers. 319 Of note, no significant reaction time differences were 320 found between first and last trial blocks for either 321 group. The analysis revealed a significant difference 322 between the relative reaction times ($F_{2.74} = 67.663$, 323 P < 0.001). There was a significant interaction of 324 FOG status and relative reaction time ($F_{2,74} = 3.37$, 325 P < 0.05). The analysis revealed no significant differ-326 ence between groups across relative reaction times 327 $(F_{1,37} = 2.39, P = 0.131)$. The interaction effect was 328 driven by a statistical difference ($t_{37} = 2.037, p < 0.05$) 329 of the relative difference between the auditory and 330 visual unisensory reaction times (i.e. A-V) in the 331 freezers (M = -43.3, SD = 55.13 ms) compared with 332 non-freezers (M = -10.32, SD = 40.23 ms). As FOG 333 tends to occur late in the course of the idiopathic PD, 334 efforts were made to address this strong relationship 335 inherent in FOG studies. A follow-up Kruskal-Wallis 336 test of disease duration (years since symptom onset) 337 between the freezers and non-freezers was performed 338 which revealed a statistical difference between the 339 groups (H(1) = 11.84, p < 0.001). 340 341

This significant difference in disease duration with respect to FOG status prompted the exploration of the



Fig. 2. Correlation of disease duration and relative sensory processing. Scatterplots displaying on the x-axis years since symptom onset and on the y-axis of the left panel, the subtraction of visual from auditory reaction times (RTs); middle panel, the subtraction of visual from auditory from audiovisual reaction times; and right panel, the subtraction of auditory from audiovisual reaction times. Each circle represents a person with Parkinson's disease (with freezers indicated in blue and non-freezers indicated in black), r-values and *p*-values are shown for significant (solid lines) and non-significant (dashed lines) regression analyses. A = auditory-alone, V = visual-alone, AV = audiovisual.

relationship between relative sensory processing (con-343 trolling for motor output delays) and disease duration, 344 three *post-hoc* correlation analyses were performed 345 on the PD group (Fig. 2). Correlation analyses were 346 performed between years since symptom onset (x-347 axis) versus 1) auditory-alone reaction times minus visual-alone reaction times (A-V); 2) audiovisual reac-349 tion times minus visual-alone reaction times (AV-V); 350 and 3) audiovisual reaction times minus auditory-alone 351 reaction times (AV-A). Again, the reaction times were 352 subtracted to account for variable motor speed in PwP. 353 Thus any differences are due to true sensory process-354 ing differences rather than slower motor responses with 355 disease progression. 356

The correlation between the subtraction of mean 357 reaction time of auditory from visual (A-V) conditions 358 and years since symptom onset revealed a signifi-359 cant relationship ($r_{37} = -0.351$, P < 0.05). A similar 360 significant relationship was found between the sub-361 traction of mean reaction time of audiovisual from 362 visual (AV-V) conditions and years since symptom 363 onset ($r_{37} = -0.415$, P < 0.0125). In contrast, there was no significant correlation between the subtraction of 365 mean reaction time of auditory and visual (A-V) con-366 ditions and years since symptom onset $(r_{37} = 0.0952)$, 367 P = 0.56). The analysis suggests that relative delays 368

in visual processing correlate with disease duration. A follow-up ANOVA with the within-participant factor of *relative* reaction time (A-V, A-AV vs A-AV) and between-participant factor of FOG status (freezers vs non-freezers) resulted no significant interaction of FOG status and relative reaction times ($F_{2,74} = 0.931$, P = 0.195). This further highlights the intricate link between FOG status and disease duration and further work is required to separate these effects.

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Miller Inequality

To test the Miller race model, reaction time range was calculated across the three stimulus types for each participant. Reaction times were sorted from fastest to slowest and the reaction time distribution was then divided into quantiles from the 5th to the 100th percentile in increments of 5% (e.g. as shown in Fig. 3A and Fig. 3B). At the individual level, a participant was said to have shown race model violation if the cumulative probability of their reaction times to the audiovisual stimulus was larger than that predicted by the race model at any quantile. We expect violations to occur in the quantiles which contain the fastest reaction times since, the faster the multisensory response, the more likely it is that multisensory



Fig. 3. A) & B) Cumulative Probability distributions for the auditory-alone (blue), visual-alone (green), audio-visual (red) and the cumulative probability predicted by the race model (black dotted) as a function of reaction time for people with Parkinson's diseases (PwP) and aged matched controls, respectively. C) & D) illustrate the subtraction of the multisensory cumulative probability and the cumulative probability predicted by the race model, known as the Miller inequality, as a function of reaction times for PwP (left) and aged matched controls (right), the errorbars depict standard error of the mean. The shaded areas indicate miller inequality values statistically greater than zero (dashed horizontal line) and signify race-model violation. E) The Miller inequality as a function of percentiles for PwP (dark grey) and aged matched controls (light grey). The shaded area indicates percentiles where the miller inequality is greater than zero (dashed horizontal line) for the control group and that are also significantly greater than PwP.

facilitation has occurred. Conversely, the quantiles 393 relating to slower multisensory reaction times are less 394 likely to violate the race model. Testing of the Miller 395 race model outlined above is also independent of vari-396 able motor responses as the multisensory response 397 times are compared directly to the individual unisen-398 sory response times. 399

Figure 3A and B shows the cumulative probabil-400 ity for the auditory-alone (blue), visual-alone (green), 401 audiovisual (red) and the cumulative probability pre-402 dicted by Miller's race-model (black-dotted) for PwP 403 and aged matched controls, respectively. The PD group 404 had a broader cumulative probability distribution for 405 all three conditions with onsets later than their aged 406 matched controls. Figure 3C and D shows the subtrac-407 tion of the value predicted by the race model from the 408 audiovisual cumulative probability curve, known as the 409 Miller inequality, as a function of reaction time divided 410 into percentiles. Miller inequality values statistically 411 greater than zero (dashed horizontal line) signify race-412 model violation. To test for within-group violation of 413 the race model, the Miller inequality values at each of 414 the reaction times were submitted to one-tailed *t*-tests 415 (greater than 0, dashed line). The analysis revealed sig-416 nificant violation of the race model (shaded areas) for 417 PwP (Fig. 3C) and aged-matched controls (Fig. 3D), 418 thus both groups showed multisensory reaction time 419 benefits. Interestingly, there was no significant dif-420 ference in race model violation between freezers and 421 non-freezers. 422

Figure 3E illustrates the Miller inequality as a func-423 tion of percentile for the PD group (dark grey) and 424 control group (light grey). To investigate differences 425 in multisensory processing between PwP and con-426 trols, taking into account reaction time differences, the 427 Miller inequalities at each percentile were submitted 428 to unpaired *t*-tests. The analysis revealed significantly 429 430 larger Miller inequality and a larger number of percentiles violating the race model (dashed line) in the 431 control group (shaded area) than the PD group. Thus, 432 the PD group has less enhanced multisensory process-433 ing compared with aged matched controls, as measured 434 by violation of the race model. 435

DISCUSSION 436

Sensory and perceptual disturbances are promi-437 nent in PD and probably contribute to bradykinesia 438 and gait disturbances [9–11]. Our results show delays 439 in response times to visual, auditory and audiovi-440 sual stimuli in PwP compared with age-matched 441

healthy controls. This is not surprising, given the 442 prominence of bradykinesia in PD. However, by com-443 paring auditory-alone, visual-alone and audio-visual 444 responses, differences in relative sensory processing 445 between PwP and controls suggest that sensory pro-446 cessing is inherently altered in PD. These changes 447 correlate with both FOG status and disease duration, 448 suggesting an effect that is specific to PD progression 449 and providing a link between these sensory abnormal-450 ities and a motor feature of PD. Specifically, there is 451 a significant difference between auditory and visual 452 reaction times in PwP which is not present in age-453 matched healthy controls. This relative difference is 454 significantly greater in those with FOG and correlates 455 with disease duration. Although multisensory facili-456 tation occurs in PD, it is significantly less enhanced 457 than in healthy controls. Reaction time tests represent a 458 simplistic model for assessing sensorimotor and cross-459 sensory function but it allows quantitative assessment 460 of deficits which underpin more complex abnormal-461 ities of sensorimotor function in PD using a simple 462 portable paradigm. 463

There is an extensive literature describing sensory 464 deficits in PD, predominantly in response to a sin-465 gle sensory modality. Few studies have quantitatively reported on multisensory integration in PD and no 467 study to date has investigated the interaction of audi-468 tory and visual modalities and their effect on reaction time. Our study has shown that both unisensory and multisensory processing abnormalities are present in 471 patients with PD. We will discuss the unisensory and 472 multisensory findings of the current study separately. 473

Unisensory processing

Our study showed that unisensory responses to both auditory and visual stimuli are slower than healthy controls. In the PD group (but not in controls) the responses to visual stimuli were significantly slower than in the auditory modality.

There is extensive clinical, behavioral, electrophys-480 iological and imaging evidence, showing abnormal 481 visual processing with PD progression at multiple 482 levels from retina to visual cortex [43, 44]. Gait param-483 eters of PwP deteriorate significantly in the absence 484 of visual feedback [1] and FOG occurs most often 485 when visual feedback is lacking (e.g. in dark envi-486 ronments) [14]. Retinal nerve fibre layer thickness 487 [45], functional neuroimaging [44, 46] and visual 488 evoked potential studies [25, 47] all provide evidence 489 that visual processing deficits correlate with both dis-490 ease duration and specific motor symptoms in PD, 491

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consistent with the findings of our study. Auditory
processing deficits are less extensive in PD but auditory evoked potentials are abnormal in PD, suggesting
both early and late information processing deficits
[27, 48–52].

Motor responses to sensory stimuli test sensory 497 processing, sensorimotor integration and motor perfor-498 mance. Existing reaction time studies which examine 499 each modality in isolation, therefore, reflect senso-500 rimotor effects rather than pure sensory ones. By 501 comparing relative differences between reaction times 502 to auditory and visual stimuli over a large number of 503 trials, the current study examines sensory responses 504 independent of a common motor output. Our study 505 shows that visual reaction times were significantly 506 slower compared with auditory reaction times in PD, 507 although both were slower compared with controls. 508 Moreover, the difference between auditory and visual 509 response times was correlated with FOG and disease 510 duration. The relative differences between freezers and 511 non-freezers appears to be due to a greater reduction in 512 auditory reaction time (i.e. faster response) in the freez-513 ers compared with controls, rather than being driven 514 by differences in visual reaction times. This suggests 515 a possible adaptive response in PwP where auditory 516 processing becomes faster relative to visual process-517 ing. This difference increases with disease duration 518 and the development of FOG. Such an adaptive pro-519 cess is consistent with a recent neuroimaging study which found functional reorganization of locomotor 521 networks in PD patients with FOG which is postu-522 lated to be a maladaptive compensatory mechanism in 523 freezers [53]. 524

Since FOG occurs more commonly in late stage PD,
 it is important to be cautious when interpreting associations involving disease duration and FOG as they are
 closely correlated. This confounder is present to some
 degree in all studies of FOG. Nevertheless, our results
 support a disease-specific effect, independent of motor
 performance, rather than a corollary of multiple other
 neurological deficits seen in this group.

533 Multisensory processing

A number of studies have implicitly examined mul-534 tisensory integration in PD. Studies on interactions 535 between proprioceptive and visual information and 536 537 their effect on spatial estimation have focused on spatial orientation and inherently invoked the investigation 538 of spatial working memory, which complicates the 539 effect of multisensory integration in PD [1, 10, 11, 540 54–57]. This is the first study to explicitly examine 541

audiovisual multisensory integration in PD and we have shown that, although multisensory facilitation occurs in PwP, it is significantly less enhanced compared with age-matched healthy controls.

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Animal studies have shown that kinesthetic sensory processing deficits correlate with degree of basal ganglia dopamine loss. With minor dopamine loss (e.g. in caudate nucleus only), this deficit can be overcome by integrating with visual information [58]. This effect has similarly been seen in clinical studies in PwP [11]. It is proposed that, as striatal dopamine loss worsens, the ability to compensate using sensory information is also lost. Single-cell recordings in mouse and cat have isolated large populations of multisensory neurons in the caudate and substantia nigra (cat) and dorsomedial striatum (mouse) [31, 32]. These suggest that the basal ganglia is a multisensory hub, crucial for integration of complex sensory stimuli from multiple modalities during execution of motor output. The striatal multisensory responses can be facilitatory or inhibitory. It is probable that a similarly large proportion of human striatal neurons have the capacity for multisensory integration, refining the response to multisensory stimuli and allowing fine motor control with complex sensory inputs. The progressive loss of striatal dopaminergic innervation affects these neurons explaining the reduced multisensory facilitation in PD. Furthermore, as progressive loss of these neurons occurs over time, the sensorimotor responses become less and less refined, eventually approaching an all-or-nothing response. In this case, certain complex sensory environments could lead to dramatic augmentation of motor output by leading to a net crude facilitatory response whereas others (e.g. doorways, noise, crowds) could cause dramatic inhibition of motor output by leading to a net crude inhibitory response, causing akinesia or freezing of gait. This is consistent with existing models of FOG, which suggest that intense sensory stimulation overloads integrated parallel processing network within the basal ganglia leading to overactivity of the output nuclei of the basal ganglia causing FOG [59-61]. Cowie et al. compared the gait of PwP and healthy controls walking through doorways and showed progressive scaling of gait parameters as PwP walked through increasingly narrow doorways [62]. As FOG frequently occurs at doorways [63], it is possible that a perceptual deficit underpins the pathophysiology of FOG [14, 64]. We posit that these sensorimotor effects occur due to multisensory interactions between visual and nonvisual sensory inputs, rather than simple unisensory deficits.

The most dramatic multisensory effect seen in PD 594 is that of sensory cueing on gait [65] and, in partic-595 ular, on FOG [66]. Sensory cueing (i.e. the use of a 596 temporal or spatial stimulus to facilitate motor output) 597 is used widely in PD as a strategy to improve gait. 598 The fact that FOG can be strikingly relieved by the 599 addition of rhythmical sensory stimuli provides fur-600 ther evidence that there are significant sensory effects 601 in PD. Given that locomotion is a highly complex mul-602 tisensory task, the improvements in gait using specific 603 sensory stimuli are probably mediated via alterations 604 in sensory integration with motor output [67]. It should 605 be noted that attention is a powerful modulator of these 606 sensory effects, in particular, sensory cueing. Indeed, 607 attentional cues alone can reduce freezing and improve 608 gait. Our findings that multisensory integration is less 609 enhanced in PD patients than in healthy controls could 610 be considered to be at odds with the observation that 611 patients with PD get significant benefit from additional 612 sensory information such as in rhythmical cueing. It is 613 important to highlight that the results of the current 614 study show that multisensory integration is reduced 615 but present in PD. We must consider the possibility 616 that intact but diminished multisensory integration may 617 be beneficial, as the over-integration of multisensory 618 information seen in older adults has been linked with 619 falls [30]. Finally, the multisensory changes seen here 620 do not correlate with either disease duration or FOG 621 status. This suggests that altered multisensory process-622 ing may occur even in early PD and may be a potential 623 biomarker for the disease. Multisensory deficits have 624 similarly been suggested as a potential biomarker in 625 other neurodegenerative disorders, such as Niemann 626 Pick Type C, using a similar paradigm [36]. 627

628 Future directions

Rehabilitation strategies which incorporate sensory 629 feedback have been shown to be of benefit in PD 630 [68-74]. Specific strategies targeting multisensory 631 integration result in behavioral and imaging changes 632 in healthy cohorts [75-78] providing evidence that 633 multisensory deficits can be improved with training. 634 Such multisensory strategies have led to improvements 635 in balance and posture in older adults [79-82] and 636 improvements in rehabilitation following spinal cord 637 injury and stroke [83, 84]. Further exploration of the 638 role of multisensory training in PD may lead to promis-639 ing therapeutic strategies for mobility, safety and FOG. 640

The main limitation of this study is the inability to
separate the effects of disease duration and FOG status.
Freezing and disease duration are intricately linked. By

controlling for one, the effect of the other is lost. This could be overcome by specifically recruiting patients with early FOG or those late in their disease course without FOG. This would, however, select out biologically different subtypes of PD. This may allow a greater understanding of the sensory processes underlying FOG but this subgroup analysis is beyond the scope of the current work.

As mentioned above, multisensory integration is intricately linked with attention and it is likely that attentional effects may contribute to the results seen above. Performance on attentional tasks are correlated with FOG, in particular when performed under temporal pressure [85, 86]. Tard et al. recently examined attention in FOG using unisensory reaction times and showed no difference between freezers and nonfreezers in simple reaction times when corrected for disease duration [87]. However, when a divided attention task was performed freezers were slower. This suggests that divided attention is impaired in FOG. Future work should focus on combining these two paradigms in order to explore the parallel effects of multisensory integration and attention.

Our multisensory findings could be explained by 667 inequality of unisensory response times. It has been 668 shown that equivalence of unisensory responses of 669 individual modalities leads to optimal multisensory 670 facilitation when those modalities are combined [88, 671 89]. If one modality dominates (as auditory does in the 672 PD cohort), then there is less opportunity for multisen-673 sory facilitation. The auditory response times in this 674 study are closely correlated with multisensory facili-675 tation. In contrast, the healthy control group displays 676 approximately equal responses to auditory and visual 677 stimuli, perhaps explaining the greater multisensory 678 integration in controls compared with the PD group. 679 Alterations in unisensory processing in PD described 680 above may, therefore, be contributing directly to the 681 diminished multisensory enhancement seen here. To 682 account for this difference, the visual and auditory 683 stimuli could be titrated for each participant to allow 684 equivalent unisensory response times, thus eliminating 685 this dominance effect. 686

Future work should include examining the effect 687 of dopaminergic therapy on the above findings. All 688 patients were tested in the "on"-medication state. It 689 would be necessary, however, to confirm that our mul-690 tisensory findings are similar off medication. Future 691 studies should also include variation of detectability 692 of unisensory stimuli to allow for optimum multisen-693 sory gain, inclusion of other sensory modalities and 694 more complex stimuli as well as variation of timing 695

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between stimuli to examine the effect of temporal window of integration. Although the discussion here is in
terms of specific modalities (visual and auditory), we
posit that there may be a more global effect of relative
sensory differences also affecting other modalities.

701 CONCLUSION

PD is associated with widespread sensory deficits: 702 peripheral and central; simple and complex; unisen-703 sory and multisensory. The precise interaction that 704 these impairments have with gait and motor con-705 trol is incompletely understood. It is, however, likely 706 that a greater understanding of these processes will 707 have positive implications for therapeutic targets and 708 rehabilitation. 709

- The current study has shown that:
- Both unisensory and multisensory delayed reac tion times exist in patients with PD, in line with
 previous findings.
- Relative differences in auditory and visual pro cessing occur in PwP and correlate with FOG
 and longer disease duration.
- 3. Multisensory integration of auditory and visual stimuli is significantly less enhanced compared with age-matched healthy controls, adding to the literature supporting both simple and higher-order sensory processing abnormalities in PD.

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