Impaired prepulse inhibition of acoustic and tactile startle response in patients with Huntington’s disease

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Abstract
The corpus striatum serves a critical function in inhibiting involuntary, intrusive movements. Striatal degeneration in Huntington’s disease results in a loss of motor inhibition, manifested by abnormal involuntary choreiform movements. Sensorimotor inhibition, or “gating”, can be measured in humans using the startle reflex: the startle reflex is normally inhibited when the startling stimulus is preceded 30–500 ms earlier by a weak prepulse. In the present study, prepulse inhibition (PPI) was measured in patients with Huntington’s disease to quantify and characterise sensorimotor gating. Compared with age matched controls, patients with Huntington’s disease exhibit less PPI. Startle gating deficits are evident in patients with Huntington’s disease when startle is elicited by either acoustic or tactile stimuli. Even with stimuli that elicit maximal PPI in normal subjects, patients with Huntington’s disease exhibit little or no PPI, and their pattern of startle gating does not show the normal modulatory effects usually elicited by changing the prepulse interval or intensity. Startle amplitude and habituation and latency facilitation are largely intact in these patients, although reflex latency is significantly slowed. In patients with Huntington’s disease, startle reflex slowing correlates with cognitive impairment measured by the dementia rating scale, and with the performance disruptive effects of interference measured by the Stroop test. These findings document a profound disruption of sensorimotor gating in patients with Huntington’s disease and are consistent with preclinical findings that identify the striatum and striatopallidal GABAergic efferent circuitry as critical substrates for sensorimotor gating of the startle reflex.

Keywords: Huntington’s disease; startle response; prepulse inhibition

The corpus striatum is thought to facilitate the maintenance of posture and ongoing motor behaviours, and to inhibit or gate inappropriate, involuntary movements.1 These critical functions are accomplished via striatal γ-aminobutyric acid (GABA) containing efferent projections to portions of the globus pallidus and zona reticulata of the substantia nigra, which in turn modulate thalamic motor nuclei directly or indirectly via the subthalamic nucleus.2 Patients with Huntington’s disease experience a progressive degeneration of intrinsic striatal cells, particularly the medium spiny, or spiny I, GABAergic efferent neurons.3,4 As a result, striatal gating mechanisms fail and adventitious choreiform movements develop in Huntington’s disease, progressing to athetosis.

Impaired gating of motor responses can be quantified from the startle reflex. This reflex is normally inhibited when the stimulus—typically a loud noise or air puff—is preceded about 100 ms earlier by a weak prepulse.5 Prepulse inhibition (PPI) is an operational measure of sensorimotor gating: the degree to which the involuntary startle response is inhibited by a weak prepulse reflects the amount of sensorimotor gating. The PPI seems to largely reflect hardwired inhibitory gating processes: it is involuntary and not learned, occurring on the first stimulus presentation and persisting thereafter without habituation.6 Startle reflex and PPI can be quantified in humans from the eyelink component of the reflex with surface EMG recordings from orbicularis oculi. Preclinical measures of startle and PPI in laboratory animals can be accomplished with EMG or accelerometer based measures of individual muscle groups or whole body movement.

As described by Davis and colleagues,4 the acoustic startle reflex is controlled by serial connections between the auditory nerve, cochlea, lateral lemniscus, nucleus reticularis pontis caudalis, and the spinal motoneuron. Whereas the primary startle response is controlled within the pontine reticular formation, startle reflex plasticity—including PPI—is modulated by forebrain circuitry. In rats, PPI seems to be modulated by neural circuitry connecting the hippocampus,7,8 ventral striatum,9,10 subpallidum,11 and pedunculopontine tegmental nucleus,12 which may access the primary startle circuit via pontine tegmental efferents to the nucleus reticularis pontis caudalis.4 Manipulations at several levels of this forebrain gating circuitry in rats can modify the normal inhibitory effects of a prepulse on the startle reflex.11,12

Based on these preclinical findings, it is possible that striatal degeneration in Huntington’s disease would be accompanied by dysfunction within startle gating circuitry.
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Briefly, each subject was seated comfortably adjacent to the recording equipment and in view of a trained technician. Two mandibular silver/silver chloride electrodes (Sensor Medics, Anaheim, CA, USA) were positioned below and to the right of the subject’s right eye, over the orbicularis oculi muscle. Placement was selected such that when subjects were asked to move their eye position virtually no activity was recorded with oscilloscope monitoring. Electrode resistances were <5 kohm. Recorded EMG activity was bandpass filtered (1–1000 Hz), and a 60 Hz notch filter was used to eliminate 60 Hz interference. A ground electrode was placed behind the right ear over the mastoid; EMG activity was digitised and 250 readings of 1 ms were recorded starting at startle stimulus onset. Acoustic startle stimuli were delivered by Telephonics (TDH-39-P, Maico, Minneapolis, MN, USA) headphones, and tactile air puff startle stimuli were delivered from a compressed air tank to the subject’s neck via a small rubber tube. Subjects sat upright and were directed to look straight ahead and to stay awake.

Subjects were tested in two separate paradigms. One paradigm (session 1; 11 controls and 11 patients with Huntington’s disease) assessed startle amplitude and PPI with two pulse modalities (acoustic and tactile) and three different intervals between prepulse and pulse (30, 60, and 120 ms). In the initial part of this session, a background 70 dB(A) white noise was followed five minutes later by 73 trials in which a startle response was elicited by a 116 dB(A) 40 ms noise burst. An initial pulse (P-ALONE) trial was followed by 72 trials consisting of six blocks of 12 trials each. Each of these 12-trial blocks included three P-ALONE trials, and nine trials in which pulse was preceded 30, 60, or 120 ms earlier by a prepulse (20 ms burst) that was 15 dB(A) above background (“30 ms”, “60 ms”, or “120 ms” respectively). In the later part of this session, 31 tactile trials were given. An initial PUFF trial consisted of a 30 pounds/square inch air puff delivered through a rubber tube placed about 5 cm from the subject’s suprasternal notch. Tube diameter was 4 mm with air pressure delivery regulated at the air supply. The initial PUFF trial was followed by five blocks of six trials that were either PUFF-only or PUFF preceded 120 ms by an acoustic prepulse as described earlier (prepulse + PUFF). For acoustic and tactile trials, a variable between trial interval averaged 15 seconds. Session 1 used one stationary EMG system located in a research laboratory (DLB) and covered a period of about 1-5 years.

A second paradigm (session 2; 11 controls and 11 patients with Huntington’s disease, none of whom were tested in session 1) assessed startle amplitude and PPI using acoustic stimuli. Slightly, a single prepulse interval (60 ms), and four different prepulse intensities (2, 4, 8, and 16 dB(A) above background). A background 70 dB(A) white
noise was followed five minutes later by 72 startle trials consisting of three conditions: (1) a 118 dB(A) 40 ms noise burst presented alone (P-ALONE); (2) the same 118 dB(A) 40 ms noise burst preceded 60 ms by pre-pulses (20 ms noise bursts) that were 2, 4, 8, or 16 dB(A) above background (PP2, PP4, PP8, or PP16); or (3) no stimulus. These six trial types were repeated six times in pseudorandom order; this block of 36 trials was repeated twice for a total of 72 trials. A variable between trial interval averaged 30 seconds. Session 2 used a portable EMG system in the UCSD Huntington’s disease clinic, over roughly a 1.5 year period. Startle sessions are not painful, and are approved by our Human Subjects Institutional Review Board.

To assess the level of “cognitive” inhibitory deficits in these patients with Huntington’s disease, the Stroop test29 was given to most subjects tested in session 2. Briefly, subjects read for 45 seconds from a standardised list of the words, “blue”, “green”, or “red”, presented in random order, and the number of words correctly read in that 45 second period was recorded (word task score). Next, for 45 seconds subjects named the colour of patterns within a standardised list of blue, green, or red coloured patterns presented in random order, and the number of colours correctly identified in that period was recorded (colour task score). Finally, subjects identified the colour of ink that was mismatched to a word (for example, the word “red” printed in blue ink should be identified as “blue”), and the number of correct responses in a 45 second period was recorded (interference task score). The Stroop test has been used as a measure of cognitive inhibition, because the interference task requires the subject to inhibit or gate the response to the semantic value of the word and to attend to its colour content.29 Our previous studies have found no significant correlation between PPI and Stroop interference scores in a normal population.20

Dependent measures for the startle reflex included startle amplitude, peak startle latency, reflex habituation, and PPI. All amplitude measurements were based on the first component (R1) of the startle response. For tactile trials, latencies were corrected for the 18 ms delay between computer generated stimulus onset and delivery of the air puff stimulus. The PPI was defined as the % reduction in startle amplitude in the presence of the prepulse compared with the amplitude in the absence of the prepulse (100 – (100 × amplitude on prepulse trial/amplitude on P-ALONE trial)). Thus a large “% score” indicates more PPI, whereas a smaller “% score” indicates less PPI. Statistical analyses consisted of mixed design analyses of variance (ANOVA) with appropriate post hoc tests. Amplitude of P-ALONE and PPI were analysed by two-way ANOVA with repeated measures on trial type (within subject) and diagnosis (between subject). The α value was 0.05.

As in other studies of human startle response,18,21,26 some subjects in each group were characterised by a relative lack of startle stimulus elicited blink-EMG activity and were therefore eliminated from the subsequent analyses (the “non-responders” were then defined as “non-responders” based on knowledge of their P-ALONE values only and cut off values were determined by inspection of the overall sample distribution. Specifically, we examined a histogram of mean startle amplitudes from a large population of control subjects tested in each session. Values tended to form a normal distribution, with the exception of a small cluster of very low values. For each session, criteria for “non-responders” were selected to exclude from analysis subjects with these very low values, leaving for analysis most subjects, whose values were normally distributed. These same criteria were then applied to the patients with Huntington’s disease. Consistent with previous studies completed with this apparatus,19 in session 1, a non-responder was defined by an average acoustic startle P-ALONE amplitude <25 units or an average PUFF amplitude <10 units (1 unit = approximately 7.7 μV). With these criteria, non-responder rates in patients with Huntington’s disease were comparable with those in controls tested with this apparatus for both acoustic startle (control rate = 32%, Huntington’s disease rate = 30-86%) and tactile startle reflexes (control rate = 12%, Huntington’s disease rate = 16-66%). Session 2 was divided into two trial blocks. Data were analysed only for trial blocks where P-ALONE amplitude exceeded 10 units. With these criteria, two trial blocks were excluded from patients with Huntington’s disease (9.1%), comparable with exclusion rates in controls tested with this apparatus (9.0%).

Results
Figures 1–4 present the results. In session 1, startle amplitude on P-ALONE trials did not differ between controls or patients with Huntington’s disease (mean amplitudes (SEM): control = 56.73 (8.35); Huntington’s disease = 49.11 (9.92); F < 1). Similarly, startle amplitude on PUFF trials did not differ between controls or patients with Huntington’s disease (mean amplitudes (SEM): control = 66.33 (17.12); Huntington’s disease = 98.89 (28.85); F < 1). Thus patients with Huntington’s disease exhibited normal startle amplitude in both acoustic and tactile modalities.

Prepulse inhibition was impaired in patients with Huntington’s disease compared with controls in both acoustic and tactile modalities (fig 1A). An ANOVA with repeated measures on trial type for acoustic trials showed an effect of diagnosis (F = 7.76, df 1,16, p < 0.015), no effect of trial type (F < 1), and no diagnosis × trial type interaction (F < 1). The two-way modulatory pattern of PPI, with more inhibition elicited by 120 ms intervals than by 30 ms intervals,21 whereas patients with Huntington’s disease failed to show this modulatory effect of increasing prepulse interval
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Figure 1  (A) Prepulse inhibition of acoustic and tactile startle reflex in patients with Huntington’s disease (HD; n = 11) and normal controls (n = 11) in session 1. *Significant effect of diagnosis, p < 0.05. (B) EMG waveforms from a control subject and a patient with Huntington’s disease, recorded from the orbicularis oculi. Responses in both are from identical trials (“PUFF” or “prepulse + PUFF”, trials No 104 and 103 from session 1 respectively). Near complete startle gating is evident in the waveform recorded from the control subject, whereas an absence of startle gating is evident in the waveform recorded from the patient with Huntington’s disease. Also evident is the increased reflex latency in the patient with Huntington’s disease compared with the control subject, as well as the presence of latency facilitation (reduced latency on “prepulse + PUFF” trial compared to “PUFF” alone trial) in both control subjects and patients with Huntington’s disease. Finally, the waveforms from both controls and patients with Huntington’s disease exhibit several late components that follow the initial reflex peak (R). This R component is the element of the startle waveform that is traditionally used in the calculation of PPI. 

Additional studies of the more complex later waveform components in patients with Huntington’s disease may provide valuable information.

over the 120 ms range. Thus there is more PPI on 120 ms trials compared with 30 ms trials in controls (paired t test: t = 3.30, df 8, p < 0.015), but not in patients with Huntington’s disease (z < 1). Analysis of tactile PPI also showed less PPI in patients with Huntington’s disease (F = 8.63, df 1,19, p < 0.01). Figure 1B shows examples of EMG waveforms from a control subject and a patient with Huntington’s disease.

Peak startle latency was increased in patients with Huntington’s disease compared with controls for both acoustic and tactile trials, but the modulatory effects of prepulses on reflex latency (latency facilitation) remained intact in patients with Huntington’s disease (figs 1B and 2). For acoustic trials, analysis of peak startle latency showed an effect of group (F = 6.25, df 1,14, p < 0.03), and an effect of trial type (F = 5.65, df 3.42, p < 0.003), but no group × trial type interaction (F = 2.34, df 3.42, NS). Similarly, for tactile trials, ANOVA showed effects of group (F = 20.89, df 1,14, p < 0.0005) and trial type (F = 5.84, df 1,14, p < 0.03), but no group × trial type interaction (F < 1). Thus the startle reflex was slowed in patients with Huntington’s disease in both acoustic and tactile modalities. Most importantly, however, patients with Huntington’s disease retained the modulatory effects of the prepulse on reflex latency, despite the significant loss of the modulatory effects of the prepulse on startle amplitude. Clearly, the prepulse modified startle characteristics (latency) in patients with Huntington’s disease, but did not gate the startle response.

It is possible that the increased startle latency and reduced PPI in patients with Huntington’s disease reflected the non-significantly older age in patients with Huntington’s disease (mean age 52.00 (SEM 4.04) compared with controls (mean age 49.36 (3.12); F < 1). Spearman rank correlations were used to test this hypothesis and showed no significant correlation between age and PPI in control subjects (R = 0.03, 0.08, -0.18 and -0.20 for 30 ms, 60 ms, 120 ms, and tactile trials respectively), consistent with our findings in larger samples of normal controls. More robust negative correlations between age and PPI were noted in patients with Huntington’s disease (R = 0.33, -0.78, -0.55, and -0.64 for 30 ms, 60 ms, 120 ms, and tactile trials respectively; p < 0.05 for 60 ms and tactile comparisons). These negative correlations in patients with Huntington’s disease suggest that increased age is associated with reduced PPI. Positive correlations between age and peak startle latency were noted in patients with Huntington’s disease (R = 0.64, 0.79, 0.64, 0.29, 0.64 and 0.76 for P-ALONE, 30 ms, 60 ms, 120 ms, PUFF and prepulse + PUFF trials respectively; p < 0.05 for 30 ms and prepulse + PUFF trials), whereas no significant positive correlations between age and peak startle latency were noted in controls (R < 0.20, all comparisons). The positive correlations between age and peak startle latency in patients with
Huntington’s disease suggest that increased age is associated with slowing of the startle reflex in these subjects. Age and startle amplitude did not correlate significantly in controls or patients with Huntington’s disease (R = 0.16, p < 0.20, all comparisons). Thus whereas increased age alone cannot account for the significant loss in PPI and increased startle latency in patients with Huntington’s disease some combination of advanced age and a diagnosis of Huntington’s disease may be related to the abnormalities in startle response. Although these findings must be interpreted with caution due to small sample size, one explanation for these correlations is that the PPI loss and reflex slowing varies with the progression of CNS pathology in patients with Huntington’s disease.

Results in session 2 were similar to those in session 1. Startle amplitude on P-ALONE trials did not differ between groups (mean amplitudes (SEM): control = 61.36 (8.15); Huntington’s disease = 57.32 (13.30); F < 1). Startle habituation was evident in controls and patients with Huntington’s disease: there was an effect of trial block (F = 22.08, df 1,20, p < 0.0001), and no block x diagnosis interaction (F < 1). As in session 1, patients with Huntington’s disease exhibited normal acoustic startle amplitude; also, the division of session 2 into two equal blocks showed that patients with Huntington’s disease exhibit normal startle habituation.

The PPI in patients with Huntington’s disease was also impaired in session 2 (fig 3). An ANOVA showed effects of diagnosis (F = 17.84, df 1,20, p < 0.0005) and trial type (F = 5.80, df 3,60, p < 0.002), and a diagnosis x trial type interaction (F = 4.37, df 3,60, p < 0.008). Patients with Huntington’s disease showed significantly less PPI than controls for 4, 8, and 16 dB trials (F = 6.92, 13.83, and 17.78 respectively). Control subjects showed the normal modulatory pattern of PPI, in this case with more inhibition elicited by more intense prepulses, whereas patients with Huntington’s disease failed to show this modulatory effect of increasing prepulse intensity. The PPI in patients with Huntington’s disease in session 2 averaged negative values: startle amplitude on prepulse trials was actually increased compared with amplitude on P-ALONE trials in this Huntington’s disease group. Prepulse potentiation has been reported in humans using 2000 ms prepulse intervals, and in rats treated with the D2 dopamine agonist quinpirole. Whereas several individual patients with Huntington’s disease in session 1 exhibited prepulse potentiation, the average positive PPI values in the session 1 Huntington’s disease group may reflect differences from session 2 either in the trial parameters, session design, patient characteristics, or EMG recording systems.

Figure 4 shows the startle latency in session 2. As in session 1, peak latency was significantly increased in patients with Huntington’s disease compared with controls, and again, the modulatory effects of prepulses on reflex
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An ANOVA showed an effect of group (F = 11.85, df 1, 20, p < 0.003), an effect of trial type (F = 14.63, df 4, 80, p = 0.0001), but no group × trial type interaction (F < 1). In session 2, unlike session 1, age differed significantly between patients with Huntington’s disease (mean age 48.09 (SEM 4.39)) and controls (mean age 30.73 (SEM 2.51)); F = 11.79, df 1, 20, p < 0.003). For this reason, a separate analysis compared PPI in the six oldest controls (M:F = 3:3; mean age 35.67 (SEM 3.19)) and the six youngest patients with Huntington’s disease (M:F = 5:1; mean age 38.67 (1.63)) tested in session 2 (age comparison control v Huntington’s disease, F < 1). The PPI in these patients with Huntington’s disease was also significantly less than PPI in the oldest controls. An ANOVA showed an effect of group (F = 16.277, df 1, 10, p < 0.003), an effect of trial type (F = 5.02, df 3, 30, p < 0.007), and no group × trial type interaction (F < 1). As PPI is greater in male than female controls,23 the over-representation of women among the age matched controls should bias against the PPI deficit found in patients with Huntington’s disease. With the small sample size and relatively uniform absence of PPI, no significant correlations were noted between PPI and age (p > 0.05, all comparisons), age at diagnosis (p > 0.05 all comparisons), or interval between first diagnosis and testing (p > 0.05, all comparisons) (for example, R, for PPI on 16 dB trials—most comparable with 60 ms trials in session 1—and age, at diagnosis, and interval between diagnosis and testing were 0.46, −0.47, and −0.18 respectively).

We assessed the relation between cognitive impairment—as reflected by dementia rating scale scores—and startle abnormalities in patients with Huntington’s disease by simple regression analyses. There was no significant correlation between dementia rating scale scores and PPI for any prepulse condition in session 1 or session 2 (0.38 > r > −0.58; p > 0.05 all comparisons). As nearly identical P-ALONE stimuli were used in session 1 and session 2, P-ALONE latencies were pooled from these two sessions. A simple regression analysis showed a significant correlation between startle latency and dementia rating scale scores, with longer reflex latencies associated with lower dementia rating scale scores (r = −0.77, p < 0.0005). Similar correlations were noted when latencies from session 1 and session 2 were analysed separately (session 1: r = −0.71; session 2: r = −0.71). As age was significantly correlated with reflex latency in session 1, it is important that there was no significant relation between age and dementia rating scale score (r = −0.24, NS).

Shoulon and Fahn ratings of total functional capacity (13 = normal, 0 = require total care)24 were available on 10 subjects at the time of testing for session 2. Correlations between total functional capacity and PPI were positive, but not statistically significant, for all prepulse conditions (0.50 > r > 0.19; p > 0.05 all comparisons). It is interesting that the only two patients with Huntington’s disease who exhibited consistent PPI (mean PPI across all conditions = 32.7%) had the highest total functional capacity scores (10 and 12); the remaining eight exhibited no overall prepulse inhibition (mean PPI across all conditions = −25.7) and had low total functional capacity scores (mean = 5.38). Chorea ratings,25 available for 10 patients, did not correlate significantly with mean PPI (r = −0.12). Results from the Stroop test (table) showed expected deficits in all three Stroop conditions in patients with Huntington’s disease.26 Compared with controls, patients with Huntington’s disease recorded lower scores in the word task (F = 43.54, df 1, 16, p < 0.0001), the colour task (F = 180.42, df 1, 16, p < 0.0001), and the interference task (F = 86.55, df 1, 16, p < 0.0001). The relative interference score, calculated as the word score divided by the interference score, was significantly greater in patients with Huntington’s disease than in controls (F = 4.52, df 1, 16, p < 0.05). This suggests that when compared with controls, patients with Huntington’s disease had more difficulty in the interference condition relative to their own “baseline” word scores.

Correlations of Stroop test scores, dementia rating scale scores, and startle latencies showed that slowing of the startle reflex in patients with Huntington’s disease was significantly correlated with performance in the word task (r = −0.76, p < 0.05), and with the relative interference score (r = −0.80, p < 0.02), but not with performance in the colour or interference tasks (r = 0.26 and −0.05, respectively; p > 0.05, both comparisons). By contrast, Stroop test scores in control subjects did not correlate significantly with any startle measure, as we have reported previously (0.16 > r > −0.55; p > 0.05 all comparisons).20 Other studies have shown strong correlations between Stroop deficits, overall cognitive impairment, and caudate atrophy in patients with Huntington’s disease.24 In our small sample, dementia rating scale scores in patients with Huntington’s disease were significantly correlated with performance in the word, colour, and interference tasks (r = 0.95, 0.72, and 0.71; p < 0.0005, 0.05, and 0.05 respectively) but were not significantly correlated with the relative interference score (r = 0.16, NS).

### Mean Stroop test score (SEM)

<table>
<thead>
<tr>
<th>Condition</th>
<th>Control (n = 10)</th>
<th>Huntington’s disease (n = 8)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Word</td>
<td>108.40 (7.21)</td>
<td>46.00 (5.46)‡</td>
</tr>
<tr>
<td>Colour</td>
<td>78.40 (2.46)‡</td>
<td>26.25 (3.08)‡</td>
</tr>
<tr>
<td>Interference</td>
<td>49.00 (2.74)</td>
<td>15.13 (2.18)*</td>
</tr>
<tr>
<td>Relative interference</td>
<td>2.28 (0.19)</td>
<td>3.28 (0.47)*</td>
</tr>
</tbody>
</table>

* p < 0.05 v control group.
†Word score/interference score.
Discussion

The degree to which a weak sensory “pre-pulse” inhibits or “gates” a potent motor reflex is an operational measure of sensorimotor gating. We reported previously that patients with Huntington’s disease are impaired in their ability to inhibit normally a reflex response to an intense auditory or tactile stimulus. There are several possible explanations that might account for these abnormalities in sensorimotor gating.

Firstly, it could be argued that the loss of PPI in patients with Huntington’s disease reflects factors not directly related to the pathophysiology of Huntington’s disease, such as generalised weakness or a reduced ability to detect an auditory prepulse. The loss of PPI in these patients cannot be easily attributed to generalised debilitation or weakness, as startle amplitude—a measure of motor reactivity—is not reduced. This suggests that the “primary” pontine reticular startle circuit is intact in patients with Huntington’s disease, and is consistent with our previous findings from preclinical and clinical studies that that startle amplitude and PPI are controlled by separate neural mechanisms. A failure to “detect” the prepulse cannot account for reduced PPI in patients with Huntington’s disease, as the patients were screened for hearing deficits, and because prepulse facilitation was intact in these same patients. Thus there is quantitative evidence of the modulatory influence of pre-pulses on the startle reflex in these patients, despite the loss or absence of PPI.

Secondly, it is possible that cognitive impairment in patients with Huntington’s disease might be accompanied by attentional deficits that weaken the inhibitory effects of pre-pulses on the startle reflex. Such an explanation is incompatible with the interpretation that PPI reflects largely “preattentive” processes, and that a pre-pulse interval of 30 ms—which produces robust PPI in controls but not patients with Huntington’s disease—is too short to evoke attentional mechanisms. In fact, PPI in patients with Huntington’s disease was not correlated significantly with cognitive impairment as measured by dementia rating scale scores. It is possible that a loss of PPI, or perhaps the increase in reflex latency, reflects drug induced changes in startle gating caused by the use of dopamine antagonists in patients with Huntington’s disease. Such an interpretation is not compatible with findings that neuroleptic drugs increase PPI in rats and humans with schizophrenia, and that startle latency is not increased in schizophrenic patients treated with much higher doses of antipsychotic drugs than were the patients in the present study.

Thirdly, it could be argued that the loss of PPI in patients with Huntington’s disease reflects the slightly increased age in these patients, compared with controls. This interpretation is not supported by the finding of lower PPI in patients with Huntington’s disease than in similar age controls in session 2. The possibility that age alone accounts for the reflex abnormalities in patients with Huntington’s disease is refuted by the finding that PPI and startle latency do not significantly correlate with age in normal controls in the present study and in previous studies with larger samples. The stronger correlation of age with gating failure and increased reflex latency in patients with Huntington’s disease suggests that these abnormalities are linked to the disease progression in Huntington’s disease. This interpretation must be made with caution given the small samples in the present study.

Previous studies have reported abnormalities in the blink reflex in patients with Huntington’s disease. Increased latency and reduced habituation of electrically evoked blinks were noted in patients with Huntington’s disease compared with normal controls and unaffected family members. This is consistent with our present findings of increased latency in both acoustic and tactile startle responses in patients with Huntington’s disease. Reflex slowing in such patients was correlated significantly with the degree of cognitive impairment measured by the dementia rating scale, and was also correlated significantly with the performance disruptive effects of interference on the Stroop test. The failure to detect habituation deficits in our study might reflect the fact that the paradigm used is not designed to be optimally sensitive to changes in habituation, which are best detected by multiple presentations of a single stimulus type. Others, however, have reported normal or even enhanced habituation of electrically stimulated blinks in patients with Huntington’s disease.

We have no direct evidence that the loss of PPI in our patients reflects degenerative changes within the striatum. Such evidence would include significant correlations between PPI and neuroimaging detected volume changes or metabolic changes in the striatum in these patients. This experimental approach has been initiated in studies of PPI in patients with schizophrenia, in whom there is a significant negative correlation (R = −0.77) between PPI and lenticular nucleus volume assessed by MRI. As patients with Huntington’s disease also exhibit atrophy and hypometabolism in cortical areas, it is possible that reduced PPI might reflect cortical dysfunction rather than striatal degeneration. Such a mechanism is not supported by animal studies, in which PPI in adult rats is not significantly reduced by large lesions of the medial prefrontal cortex, the frontal poles, the entire frontal cortex, the dorsal hippocampus and overlying parietal cortex, the parietal cortex alone, or the ventral hippocampus.

Although the cause of striatal degeneration in Huntington’s disease is not well understood, several investigators have hypothesised that loss of striatal medium spiny cells results from an endogenous excitotoxin, similar to kainic or quinolinic acid. Quinolinic acid is a product of the kynurenine pathway. Increased activity of a quinolinic acid synthetic enzyme
Impaired prepulse inhibition of acoustic and tactile startle response in patients with Huntington’s disease has been reported in postmortem brain tissue from patients with Huntington’s disease. In rats, infusion of quinolinic acid results in a selective destruction of neurons, leaving fibres of passage intact. Recently, we found that PPI is impaired in rats after quinolinic acid lesions of the anteroventral or dorsal/posterior striatum (Kodsi and Swerdlow, unpublished data) and after microinjection into corresponding regions of the anteroventral or dorsal/posterior globus pallidus (Kodsi and Swerdlow, unpublished data). Impaired startle gating in patients with Huntington’s disease can thus be mimicked in rats by striatal damage caused by a neurotoxin that has been implicated in the pathophysiology of Huntington’s disease, or by pharmacological interruption of striatopallidal GABAergic neurotransmission.

Deficits in PPI have been reported in patients with schizophrenia and in patients with obsessive compulsive disorder; similar findings have been reported in patients with Tourette’s syndrome. Importantly, the pattern of PPI deficits varies between different gating disorders. For example, whereas the total amount of PPI is reduced in these patients with Huntington’s disease compared with matched controls, PPI in patients with obsessive compulsive disorder and schizophrenia exhibits “normal” patterns of stimulus modulation: PPI increases with increasing prepulse intensity, with maximal inhibition seen at short (30–60 ms) prepulse intervals. By contrast, the modulatory influences of prepulse parameters on PPI are totally abolished in Huntington’s disease. This suggests that the loss of PPI in patients with Huntington’s disease reflects a complete “disconnection” of the primary startle circuit from forebrain elements that modulate the inhibitory effects of the prepulse. By contrast, PPI deficits in patients with schizophrenia or obsessive compulsive disorder seem to reflect a reduced sensitivity to the inhibitory effects of the prepulse, or a diminished influence of inhibitory forebrain circuitry on the primary startle circuit. As our preclinical studies suggest that the modulatory effects of prepulse intensity and interval on PPI are controlled at discrete sites within the startle gating circuitry, it might ultimately be possible to link different patterns of PPI deficits to separate substrates within limbic motor gating circuitry.

Genetic studies have associated a gene on the G8 fragment of chromosome 4 with the autosomal dominant transmission of Huntington’s disease. More specific characterization of the Huntington’s disease gene and its relation to the pathophysiology of the disease will benefit from the identification of objective phenotypic markers of impaired striatal function and gating. The PPI may serve as such a marker, because it is an objective and highly quantitative measure of sensorimotor gating with a discrete underlying neural basis that is impaired in Huntington’s disease. Startle reflex latency is also an objective, quantifiable variable that seems to correlate significantly with cognitive impairment and the performance disruptive effects of interference in Huntington’s disease. As startle measures are non-invasive, brief (a typical session lasting 25 minutes), and can be obtained with portable testing units, it would be realistic to pursue larger studies of sensorimotor gating and startle reflex latency in families of Huntington’s disease probands. A similar strategy is being used with the event related potential P50 gating paradigm in a large proband of schizophrenic patients.

In summary, sensorimotor gating is impaired in patients with Huntington’s disease compared with controls. The loss of sensorimotor gating is evident in both acoustic and tactile modalities, but is accompanied by normal startle amplitude, normal habituation, and normal prepulse latency facilitation. The pattern of impaired gating in Huntington’s disease is distinct from the pattern of deficits previously reported in patients with schizophrenia or obsessive compulsive disorder, and suggests a complete or near complete disconnection of forebrain modulatory influences on “primary” startle circuitry in the reticular formation of the brain.

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14 Madan S. Mental status examination for organic mental syndrome in the elderly patient. In: Bellack LS, Karasu...
Some speech disorders

Continued from page 191.

Marcel Proust, 1913, Remembrance of things past. Vol 1 Swann's way.
Moreover, the name Swann, with which I had for so long been familiar, had now become for me (as happens with certain aphasics in the case of the most ordinary words) a new name.

James Joyce, 1922, Ulysses
Your attention! We're née that fou. The Leith Police dismiss us.

Sinclair Lewis, 1923, Babbitt
"And there's no if, and or but about it!"

Marcel Proust, 1927, Time regained, Vol 3
Of the two, one, the intellectual one, passed over in the time in complaining that he suffered from progressive aphasia, that he constantly pronounced one word, one letter by mistake for another.

PG Wodehouse, undated, The story of Webster, from Mr Mulliner's relations
Webster, like the Stag at Eve, had now drunk his fill. He had left the pool of alcohol and was walking round in slow, meditative circles. From time to time he mewed tentatively, as if he were trying to say "British Constitution." His failure to articulate the syllables appeared to tickle him...