Complex and Simple Clinical Reaction Times Are Associated with Gait, Balance, and Major Fall Injury in Older Subjects with Diabetic Peripheral Neuropathy

ABSTRACT


Objective: The aim of this work was to identify relationships between complex and simple clinical measures of reaction time (RTclin) and indicators of balance in older subjects with and without diabetic peripheral neuropathy (DPN).

Design: Prospective cohort design. Complex RTclin accuracy, simple RTclin latency, and their ratio were determined using a novel device in 42 subjects (mean ± SD age, 69.1 ± 8.3 yrs), 26 with DPN and 16 without. Dependent variables included unipedal stance time (UST), step width variability and range on an uneven surface, and major fall-related injury over 12 months.

Results: In the DPN subjects, the ratio of complex RTclin accuracy to simple RTclin latency was strongly associated with longer UST (R/P = 0.653/0.004), and decreased step width variability and range (R/P = −0.696/0.001 and −0.782/−0.001, respectively) on an uneven surface. Additionally, the 2 DPN subjects sustaining major injuries had lower complex RTclin accuracy:simple RTclin latency than those without.

Conclusions: The ratio of complex RTclin accuracy:simple RTclin latency is a potent predictor of UST and frontal plane gait variability in response to perturbations and may predict major fall injury in older subjects with DPN. These short latency neurocognitive measures may compensate for lower limb neuromuscular impairments and provide a more comprehensive understanding of balance and fall risk.

Key Words: Reaction, Balance, Gait, Accidental Falls
Accidental falls are a public health priority, with the estimated lifetime costs of more than $34 billion in 2010. The risk of fall-related injury in older people with diabetic peripheral neuropathy (DPN) is at least twice that in older people without DPN, and this increased risk impedes the engagement of this population in meaningful life experiences as well as walking exercise programs, which are the foundation for management of type II diabetes mellitus. Moreover, a distal symmetric polyneuropathy is common in older age groups, with approximately 2/3 related to diabetes mellitus, so that an estimated 20 million Americans aged 55 to 85 are affected.

Falls occur most frequently while older adults are walking on uneven or irregular surfaces, with lateral falls appearing to have greater injury potential than other falls. Therefore, the ability to withstand (or reject) a frontal plane perturbation is necessary for safe ambulation. Prior work confirms that frontal plane hip strength is essential to maintain control of the head/arms/trunk during single limb stance and allows appropriate swing limb foot placement. Accordingly, our prior research in older subjects with varying degrees of DPN found that frontal plane neuromuscular factors (laboratory measured hip strength as normalized rate of torque in the frontal plane and ankle inversion/eversion proprioceptive thresholds as a ratio: $\text{Hip}^{\text{STR}}/\text{Ank}^{\text{PRO}}$) strongly predicted unipedal stance time (UST), sagittal plane responses to perturbation while walking, and prospectively recorded falls and fall-related injury.

An increasing volume of research links cognitive functions and gait/fall risk. Of the cognitive domains, evidence suggests that executive function, specifically inhibitory executive function, is the most important with reference to gait, balance, and fall risk. Intact inhibitory executive function provides the ability to attend selectively to a specific stimulus while diminishing others, and the ability to “withhold pre-potent (automatic) responses.” In contrast, impairments in executive function lead to the inability to rapidly selectively focus on relevant afferent stimuli and/or alter routine motor patterns when they become inappropriate or unnecessary. As such, older subjects with inhibitory executive impairments demonstrate decreased balance in the setting of multiple afferent stimuli and are at increased risk for falls and injury.

We developed and validated a clinical reaction time device and method ($\text{RT}_{\text{clin}}$Dev; Fig. 1) that allows measurement of simple and complex reaction times in a clinical setting. The clinical reaction time device ($\text{RT}_{\text{clin}}$Dev) is used in a ruler-drop testing paradigm, where simple clinical reaction time ($\text{RT}_{\text{clin}}$) latency is determined by the time a vertically oriented rod falls before it is caught by subject hand closure. Complex $\text{RT}_{\text{clin}}$ accuracy requires the subject to catch the same device solely on the randomly determined 50% of trials in which lights affixed to it illuminate. Simple $\text{RT}_{\text{clin}}$ latency is measured in millisecond (ms), and complex $\text{RT}_{\text{clin}}$ accuracy is measured as the percentage of trials in which the subject responds correctly. Simple $\text{RT}_{\text{clin}}$ latency requires sustained attention and rapid reaction. A high level of complex $\text{RT}_{\text{clin}}$ accuracy requires rapid processing and response, as the subject must interpret the signal and make a decision whether to catch the $\text{RT}_{\text{clin}}$Dev within the approximately 400 ms prior to it striking the floor. The challenging component of complex $\text{RT}_{\text{clin}}$ testing is to selectively attend to the lights on the $\text{RT}_{\text{clin}}$Dev and to withhold catching the rapidly descending rod.

**FIGURE 1** Clinical reaction time device ($\text{RT}_{\text{clin}}$Dev; panel A) used to determine simple $\text{RT}_{\text{clin}}$ latency and complex $\text{RT}_{\text{clin}}$ accuracy (panels B and C) as described in text.
device when the lights do not illuminate, requiring the ability “to inhibit a prepotent (automatic) response”. We, and others have found this action-based technique of determining simple and recognition reaction times to be superior (faster and less variable) than perception/computer-based techniques. Therefore, the RT\textsubscript{clin,Dev} allows the measurement of short latency inhibitory executive function by measuring the frequency with which subjects can withhold a prepotent/automatic response. Complex RT\textsubscript{clin} accuracy and simple RT\textsubscript{clin} latency are of interest separately and also as a ratio (complex RT\textsubscript{clin} accuracy:simple RT\textsubscript{clin} latency), which reflects both accuracy of decision making and speed of response. The ratio allows the greatest (best) scores to occur in subjects who were accurate as well as quick, whereas the lowest (worst) scores would occur in subjects who were neither.

Given the inability of frontal plane lower limb neuromuscular attributes (Hip\textsuperscript{STR}:Ank\textsuperscript{PRO}) to predict frontal plane responses to perturbation and identify the subjects who sustained a major injury during follow-up, we performed a secondary analysis to determine whether the short latency neurocognitive attributes complex RT\textsubscript{clin} accuracy and simple RT\textsubscript{clin} latency as determined by RT\textsubscript{clin,Dev} were related to these important outcomes. More specifically, we hypothesized that decreased/inaccurate baseline complex RT\textsubscript{clin} accuracy and/or increased/slow simple RT\textsubscript{clin} latency would be associated with (1) decreased unipedal stance time (UST); (2) increased frontal plane gait variability in older subjects with and without DPN while walking on smooth and uneven surfaces; and (3) major fall-related injuries prospectively determined over 1-year follow-up.

METHODS

Design

We performed a prospective cohort study, with baseline evaluations of independent variables followed by UST testing, gait evaluation, and then 1-year prospective follow-up for falls and fall-related injuries. The work presented here is a secondary analysis to determine whether complex RT\textsubscript{clin} accuracy and simple RT\textsubscript{clin} latency as determined by RT\textsubscript{clin,Dev} were related to UST, frontal plane gait variability, and major fall-related injuries, outcomes which had not been predicted by lower limb neuromuscular factors. To minimize bias, the study team member evaluating UST and recording falls and fall-related injuries, was blinded with regard to baseline lower limb neuromuscular and RT\textsubscript{clin} variables. The original sample size of 42 subjects, approximately 2/3 with DPN of varying severity, was recruited so as to create a group of older adults with a spectrum of lower limb peripheral neuromuscular function that would provide a high likelihood of significant correlation coefficients between lower limb neuromuscular variables and UST and perturbed gait measures. Therefore, sample size represents a study limitation with regard to the falls and fall-related injury outcomes.

Subjects

The research protocol was approved by the Institutional Review Board, and all participants provided informed written consent. Subjects were recruited between July of 2009 and December of 2011, and prospectively followed until January of 2013. Ninety-nine subjects were screened by phone, with 23 failing to meet inclusion/exclusion criteria and 20 not interested in participating, usually owing to the time commitment or not staying in the area. Of the remaining 56, 6 did not pass the physical examination screen and 4 did not show for scheduled appointments. Of the remaining 46, 4 canceled before the onset of testing owing to medical concerns within their families, to provide the final cohort of 42 subjects. Of these, 26 subjects had varying degrees of DPN and 16 were without (Table 1). All 42 subjects were included in the UST analyses. However, 10 subjects (7 with DPN and 3 without) were preferred not to participate in the gait, falls, and fall-related injury portion of the study, citing concerns with the associated time commitment, and so those

<table>
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<th>TABLE 1 Study subject characteristics</th>
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<td>All Subjects (n = 42)</td>
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<td>Sex, women, n (%)</td>
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<td>Age, years</td>
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<td>MDNS</td>
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The only significant group differences are subjects with DPN having greater MDNS scores (worse neuropathy) than subjects without DPN (P < 0.001) and a trend toward greater BMI (P = 0.052).

MDNS, Michigan Diabetes Neuropathy Score.
analyses include 32 subjects. The 10 subjects who dropped out after baseline and UST testing were not significantly different from those who remained with regard to age, sex, body mass index, or neuropathy severity.

Subjects were recruited from the University of Michigan Orthotics and Prosthetics Clinic, Endocrinology Clinic, and the Older Americans Independence Center Human Subjects Core between November 2010 and January 2012. Written informed consent was obtained from all subjects after review from the institutional review board. Eligible subjects were between the ages of 50 and 85 years, weighed less than 136 kg (as required for harness support system during gait evaluation), were free of central neurologic disease, including Parkinson disease or Parkinsonism, vestibular disorders, symptomatic coronary artery disease, plantar skin sores or joint replacement within the previous year, symptomatic postural hypotension, severe musculoskeletal deformity (eg, amputation or Charcot changes), lower extremity or back pain that limited standing to less than 10 minutes, were able to walk 1 block or more, had greater than antigravity ankle strength (>grade 3/5 by manual muscle testing), and corrected vision not worse than 20/50. Subjects with DPN had a history of type 2 diabetes mellitus confirmed by review of records and the ongoing use of oral hypoglycemic agents or insulin. The presence of DPN was confirmed by the following: (a) symptoms (subject reported altered sensation in the distal lower limbs); (b) signs, Michigan Diabetes Neuropathy Score (MDNS; a 46 point scale with increasing score relating to more severe DPN) >1031; and (c) bilaterally abnormal fibular motor nerve conduction studies (recording over the extensor digitorum brevis, defined as amplitude <2.0 mV, latency >6.2 ms, and/or conduction velocity <41.0 m/s, using Nicolet Viking 4). Subjects without DPN had no history of diabetes mellitus, no symptoms or signs of DPN (MDNS <10), and normal fibular nerve conduction studies. Subjects were excluded if they reported a fall within 1 month of testing.

**Independent Variables**

**Simple RT<sub>clin</sub> Latency**

Simple RT<sub>clin</sub> latency was determined using the RT<sub>clin.Dev</sub>. This is a custom-built device that consists of a 107-cm collapsible, rigid, lightweight dumbbell-shaped shaft affixed to an 11 × 6 × 2.5-cm spacer box housing a linear accelerometer, timing circuit, microprocessor, battery, liquid crystal display, and 2 light-emitting diodes. Using a “ruler drop” test paradigm, participants stood with their dominant forearm resting on an adjustable table surface so that the hand was positioned at the edge. The examiner suspended the device vertically so that the spacer box rested between the thumb and other digits (Fig. 1). After predetermined random delay times ranging from 2 to 5 seconds, the examiner released the device. The participants were instructed to catch the falling device as quickly as possible every trial. Participants were given 2 simple RT<sub>clin</sub> practice trials before 10 data collection trials. Simple RT<sub>clin</sub> latency was defined as the mean of the 10 trials and reported in milliseconds.

**Complex RT<sub>clin</sub> Accuracy**

Subjects were positioned in the same manner as for determining simple RT<sub>clin</sub> latency. To determine complex RT<sub>clin</sub> accuracy, the light-emitting diodes on the device illuminated randomly during 50% of the trials at the instant the device accelerated upon release. Participants were instructed to catch the falling RT<sub>clin.Dev only</sub> during those trials when the light-emitting diodes illuminated, and to resist catching it when the light-emitting diodes did not turn on, consistent with a standard go/no-go testing paradigm. Verbal instructions emphasized response accuracy, not speed. Participants completed 6 practice trials before 20 data collection trials. Complex RT<sub>clin</sub> accuracy was recorded as the percentage of correctly performed trials/total number of trials.

**Complex RT<sub>clin</sub> Accuracy:Simple RT<sub>clin</sub> Latency**

The ratio of complex RT<sub>clin</sub> accuracy to simple RT<sub>clin</sub> latency was also of interest, given that it reflects both accuracy of decision making and speed of response, so that the greatest (best) scores occur in subjects who were both accurate and quick, whereas the lowest (worst) scores occur in subjects who were neither. The resulting value was multiplied by 1000 given that simple RT<sub>clin</sub> latency was reported in millisecond.

**Dependent Variables**

**Unipedal Stance Time (UST)**

As per prior protocol, 3 trials were performed on subject’s foot of choice, and then 3 trials on the opposite foot. One practice trial was allowed for each foot before data collection. The mean UST in seconds of all 6 trials was the outcome of interest.

**Gait Analysis of Frontal Plane Gait Variability**

As illustrated in prior reports subjects were fitted in a safety harness that housed a cable fastened to an overhead track. The cable was secured high enough to catch subjects should they...
experience an accidental fall. Subjects wore flat-soled, standard athletic shoes. Kinematic data were collected through 2 optoelectronic markers (infrared-emitted diodes) positioned 5 cm apart on an aluminum strip (10 cm × 1.5 cm) that was bent at a 90-degree angle and inserted under the laces of each shoe at the midline. The top marker was located anterior to the center of the malleoli. The subjects also wore a waist marker positioned on a belt at the level of the umbilicus.

The smooth surface was constructed of flat linoleum tile. The uneven surface, which has been described in prior reports, was created by placing a 1.5 × 10-m piece of dark industrial carpet over randomly distributed prism-shaped blocks of wood (height, 1.5 cm; width, 3.5 cm; length, 6–16 cm). The blocks of wood were located within the mid 6.5-m section of carpet and were not changed between trials. For trials on both the smooth and uneven surfaces, subjects were instructed to walk down the runway at their own pace, as if they were “walking to mail a letter.” Subjects completed 10 trials on each walkway, with the first 2 used for accommodation and the last 8 for data collection. The subjects ambulated down the walkway toward an optoelectronic camera system (Optotak 3020, Northern Digital Corp, Waterloo, Ontario), which recorded marker positions at 100 Hz. To detect heel strike and toe off, each subject wore rearfoot and forefoot foot switches in each shoe (force-sensing resistors made by FlexiForce, Tekscan Inc, South Boston, MA). These sensors were connected to the data acquisition hardware. A custom C++ program operating in conjunction with the Optotak Application Programming Interface was then used to track the timing of heel strike and toe off for each step. Once the heel strike and toe off information were known, then the timing of double support was known and step width and step length was taken from the kinematic marker data as previously described. Kinematic data were quantified by using a custom algorithm written in MATLAB. Frontal plane gait variability was measured using the standard deviation (SD) and range of step width and referred to as step width variability and step width range, respectively. The latter is of interest given that falls are unusual events as compared to the number of steps most people take each day, and therefore “outlier” steps in the frontal plane likely have clinical relevance.

Recording Falls and Fall-related Injuries

Falls and fall-related injuries were recorded through 1 year of follow-up using methods described by Tinetti et al. Twenty-six calendars (each spanning a 2-week period) were provided to each of the 32 subjects so that data could be collected prospectively for each subject for 1 year. Subjects assessed themselves daily, and if a fall or fall-related injury occurred, they checked a box on the calendar and recorded a description of the circumstances. Subjects returned the calendars every 2 weeks, and in the few cases where a subject did not return a calendar, the study coordinator contacted the subject to determine the occurrence of a fall or fall-related injury during the missed time period. Falls were defined as unintentional changes in body posture that resulted in the subject coming to rest on the ground or other lower level that was not a consequence of a physical blow or loss of consciousness.

Fall-related injuries were separated into 2 groups: major and minor. Major injuries were defined as an Abbreviated Injury Score greater than 2, and minor injuries were defined as abrasions, bruises, and lacerations that did not require sutures but interfered with the subject’s activities of daily living for at least 24 hours.

Statistical Analyses

Descriptive statistics were generated for the independent variables, Complex RT, accuracy and simple RT, latency, and also for the laboratory-based dependent variables, UST, step width variability and range on the smooth and uneven surfaces, and inspected for normality. Bivariate relationships between these were determined using Pearson correlation coefficients. These relationships were evaluated, separately, for subjects with and without DPN. Significance was set at \( P = 0.05/3 \) or \( P = 0.016 \), and a trend set at \( P = 0.032 \), to adjust for 3 correlations being performed for each group.

To determine the independence of the relationships identified between complex RT, accuracy, Simple RT latency and their ratio (referred to collectively as “short latency neurocognitive attributes”) and the outcomes UST, step-width variability, and range, multiple regression analyses were performed by singly entering other variables with known relationships to the outcome variables. More specifically, age, Michigan Diabetes Neuropathy score, and the composite measure of lower limb neuromuscular function previously demonstrated to predict UST and falls (Hip\(^{STR}\), Ank\(^{PRO}\)) were evaluated.

Major fall-related injury group differences in simple RT, latency, complex RT, accuracy, and their ratio among the subjects with DPN were not
subjected to statistical analyses given the small number of subjects sustaining such an injury.

RESULTS

Subjects

Study subject characteristics are provided in Table 1. Forty-two subjects were available for the UST testing. Ten subjects had barriers to continuing participation and so 32 subjects were included in the gait testing and prospective evaluation of fall-related injuries.

Unipedal Stance Time

When all subjects were considered, UST demonstrated significant correlations with simple RT\textsubscript{clin} latency ($R/P = -0.421/0.005$) and complex RT\textsubscript{clin} accuracy:simple RT\textsubscript{clin} latency (0.386/0.011), but not complex RT\textsubscript{clin} accuracy (0.215/0.172). However, when considered separately, the subjects without DPN demonstrated no significant relationships between complex RT\textsubscript{clin} accuracy, simple RT\textsubscript{clin} latency, and their ratio, and UST (Table 2B). The limitation of UST to a maximum of 30 seconds resulted in a ceiling effect for this group, and the resulting skewing of the data may have obscured an association between the 2 variables (Fig. 2B). In contrast, the subjects with DPN demonstrated robust relationships with UST, with decreased (quicker) simple RT\textsubscript{clin} latency and increased (more accurate) complex RT\textsubscript{clin} accuracy, and greater complex RT\textsubscript{clin} accuracy:simple RT\textsubscript{clin} latency ratio significantly related to longer UST (Table 2A and Fig. 2A).

Given prior work strongly correlating the ratio of Hip\textsuperscript{STR}:Ank\textsuperscript{PRO} and UST in the same cohort, this variable was included with complex RT\textsubscript{clin} accuracy: simple RT\textsubscript{clin} latency in a regression model. When subjects without DPN were considered, the ratio of Hip Strength to Ankle Proprioceptive precision was the only significant predictor. However when subjects with DPN were analyzed, complex RT\textsubscript{clin} accuracy: simple RT\textsubscript{clin} latency and Hip\textsuperscript{STR}:Ank\textsuperscript{PRO} were both significant predictors, with a resultant $R^2$ of 0.680 (Table 2B). No other variables contributed to the model.

Frontal Plane Gait Variability

When all subjects were considered, step width range demonstrated significant correlations with simple RT\textsubscript{clin} latency, complex RT\textsubscript{clin} accuracy and complex RT\textsubscript{clin} accuracy:simple RT\textsubscript{clin} latency on the uneven surface ($R/P = 0.541/0.001$; $R/P = -0.523/0.002$; and $R/P = -0.709/0.001$, respectively) but not the even surface ($R/P = 0.137/0.441$; $R/P = -0.183/0.300$; and $R/P = -0.201/0.255$, respectively). However, when subjects without DPN were considered separately, there were no significant relationships between the short latency neurocognitive attributes and frontal plane gait variability or range on the even or uneven surfaces (Table 3). The subjects with DPN demonstrated a trend between complex RT\textsubscript{clin} accuracy:simple RT\textsubscript{clin} latency and step width variability on the even surface. However, the subjects with DPN demonstrated strong relationships

| TABLE 2A | Relationships (Pearson correlation coefficients, $R$, and $p$ values) between simple RT\textsubscript{clin} latency (SRT), complex RT\textsubscript{clin} accuracy (RRTAcc), and their ratio (RRTAcc:SRT), with UST in subjects with and without DPN |
|---|---|---|---|
| **Subjects Without DPN ($R/P$ Values)** | **Subjects with DPN ($R/P$ Values)** |
| SRT | $-0.473/0.064$ | $-0.520/0.008$ |
| RRTAcc | $0.026/0.924$ | $0.472/0.017$ |
| RRTAcc:SRT | $0.319/0.229$ | $0.653/0.004$ |

| TABLE 2B | Results of multivariate analyses with UST as the dependent variable |
|---|---|---|---|---|---|
| **Subjects Without DPN** | **Subjects with DPN** |
| RRTAcc:SRT | Hip\textsuperscript{STR}:Ank\textsuperscript{PRO} | RRTAcc:SRT | Hip\textsuperscript{STR}:Ank\textsuperscript{PRO} |
| Beta | NS | 3.82 | 0.505 | 0.525 |
| $p$ value | NS | 0.002 | 0.001 | 0.001 |
| $R^2$ | 0.580 | 0.680 | | |
FIGURE 2  Relationships between complex RT<sub>clin</sub> accuracy/Simple RT<sub>clin</sub> latency and UST in subjects with, and without, DPN. A, Subjects with DPN. B, Subjects without DPN.

TABLE 3  Relationships (Pearson correlation coefficients, R<sub>i</sub>, and P values) between SRT, RRTAcc, and their ratio (RRTAcc:SRT), and step width variability and range on even and uneven surfaces

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<tr>
<th>Step Width Variability Even Surface</th>
<th>Step Width Variability Uneven Surface</th>
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<tr>
<td>Subjects Without DPN R/P Values</td>
<td>Subjects with DPN R/P Values</td>
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<tr>
<td>SRT  -0.025/.936</td>
<td>0.331/.143</td>
</tr>
<tr>
<td>RRTAcc -0.197/.519</td>
<td>-0.397/.075</td>
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<tr>
<td>RRTAcc:SRT -0.119/.698</td>
<td>-0.481/.027</td>
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<tr>
<th>Step Width Range Even Surface</th>
<th>Step Width Range Uneven Surface</th>
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<tr>
<td>Subjects Without DPN R/P Values</td>
<td>Subjects with DPN R/P Values</td>
</tr>
<tr>
<td>SRT  -0.134/.662</td>
<td>0.232/.311</td>
</tr>
<tr>
<td>RRTAcc -0.037/.905</td>
<td>-0.245/.285</td>
</tr>
<tr>
<td>RRTAcc:SRT -0.040/.896</td>
<td>-0.308/.174</td>
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between complex RT\textsubscript{clin} accuracy and complex RT\textsubscript{clin} latency for both step width variability and range on the uneven surface (Table 3). The relationship between the ratio of neurocognitive attributes and step width range on the uneven surface was particularly robust, with an $R^2$ of 0.611 (Fig. 3). Notably, none of the demographic variables or hip strength:ankle proprioceptive precision demonstrated significant or near significant relationships with step width variability or range on the uneven surface. DPN sustained a major injury in the 1-year follow-up. These 2 subjects demonstrated increased (worse) simple RT\textsubscript{clin} latency and decreased complex RT\textsubscript{clin} accuracy, as well as decreased complex RT\textsubscript{clin} accuracy:simple RT\textsubscript{clin} latency compared to subjects with DPN not sustaining major injury. The mean complex RT\textsubscript{clin} latency:simple RT\textsubscript{clin} latency of the 2 DPN subjects with major injuries were approximately 2 SDs less than those subjects without major injury (Table 4).

**DISCUSSION**

The results of this research demonstrate that in older adults with DPN there appear to be relationships between the short latency neurocognitive attributes, complex RT\textsubscript{clin} accuracy, simple RT\textsubscript{clin} latency, and their ratio, and 3 important mobility-related outcomes: UST, frontal plane gait variability on an uneven surface, and major prospective fall-related injuries. These findings support at least 3 novel, clinically relevant concepts: (1) when the data reported here are considered with prior work,\textsuperscript{27} RT\textsubscript{clin} seems to be an innovative, clinically accessible method for

**Major Injury**

There were no significant differences in complex RT\textsubscript{clin} accuracy or simple RT\textsubscript{clin} latency, or their ratio, between subjects sustaining minor injury compared to those who did not, with all $p$ values $> 0.40$ (Table 4). One subject without DPN sustained a major injury, an 80-year-old woman with complex RT\textsubscript{clin} accuracy and simple RT\textsubscript{clin} of 0.76 and 192 ms, respectively, with a ratio of 3.95, all of which were within 1 SD of the mean values for all subjects without DPN (0.72 $\pm$ 0.09, 173 $\pm$ 22, and 4.2 $\pm$ 0.8, respectively). Two of the subjects with DPN sustained a major injury in the 1-year follow-up. These 2 subjects demonstrated increased (worse) simple RT\textsubscript{clin} latency and decreased complex RT\textsubscript{clin} accuracy, as well as decreased complex RT\textsubscript{clin} accuracy:simple RT\textsubscript{clin} latency compared to subjects with DPN not sustaining major injury. The mean complex RT\textsubscript{clin} accuracy:simple RT\textsubscript{clin} latency of the 2 DPN subjects with major injuries were approximately 2 SDs less than those subjects without major injury (Table 4).

**FIGURE 3**  Relationship between complex RT\textsubscript{clin} accuracy:simple RT\textsubscript{clin} latency and step width range on the uneven surface in subjects with DPN ($R^2 = 0.611$).

**TABLE 4**  Simple RT\textsubscript{clin} Latency (SRT), Complex RT\textsubscript{clin} Accuracy (RRTAcc), and their ratio RRTAcc:SRT) in subjects with and without major and minor injuries

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<thead>
<tr>
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<th>Subjects with DPN</th>
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<tr>
<td></td>
<td>+Major Injury (n = 2)</td>
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<tr>
<td>SRT</td>
<td>229 $\pm$ 68</td>
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<tr>
<td>RRTAcc</td>
<td>0.55 $\pm$ 0.14</td>
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<tr>
<td>RRTAcc:SRT*</td>
<td>2.5 $\pm$ 0.8</td>
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*Multiplied by 1000.
measuring inhibitory executive function over a brief time interval; (2) the data link RT_{clin}-based short latency neurocognitive attributes to UST and frontal plane responses to sequential perturbations during walking and, in so doing, provide a mechanism by which poor inhibitory executive function increases fall risk; and (3) the relationships between complex RT_{clin} accuracy: simple RT_{clin} latency and measures of balance, frontal plane gait control, and major injuries were identified predominantly in the subjects with DPN, suggesting that short latency neurocognitive capability is of greater importance in the setting of lower limb neuromuscular impairment. These points are addressed in sequence below.

**Complex RT_{clin} Accuracy as a Measure of Short Latency Inhibitory Executive Function**

There seem to be 2 inhibitory challenges to the patient or subject attempting to achieve high complex RT accuracy, one afferent in nature and the other efferent. The afferent task is to selectively attend to the lights on RT_{clin}Dev while ignoring the dominant visual stimulus, the rapidly descending device. This requires the ability to quickly prioritize and/or suppress incoming stimuli. The efferent task is to withhold the urge to catch the falling device. Accordingly, Giordani and Persad\(^2\) describe inhibitory executive function as the ability to “prevent distracting information from...causing interference,” and “prevent pre-potent (automatic) responses that may not be appropriate...” and both seem present when complex RT is tested. There are a variety of other methods for evaluating executive function, most commonly through Stroop or Trails B testing, but neither requires the subject to make a decision within approximately 420 ms or evaluates visuomotor pathways. This may represent an advantage for RT_{clin}Dev when evaluating fall risk, as falls are events that occur, or are prevented, over a brief time interval and often include visuomotor responses. Furthermore, the time available for decision making is similar to the time interval available for altering swing limb trajectory (435 and 480 ms for our subjects on smooth and uneven surfaces, respectively), possibly making RT_{clin}Dev particularly suitable for gait evaluation. Another potential advantage over standard methods of measuring executive function is that RT_{clin}Dev is a 3-dimensional object rather than a screen image. We, along with Montare,\(^{29,30}\) have found that simple and complex reaction time tests using falling objects yield quicker responses and decreased variability compared to screen-based measurements, possibly owing to the fact that moving objects activate the visuomotor pathways, whereas the latter works through visuo perceptual pathways.\(^{29}\) In one of the only other studies to evaluate short latency inhibitory executive function and prospective falls, Schoene et al.\(^{36}\) evaluated a large group of older subjects with respect to their ability to perform a Stroop-like test on a computer-controlled mat, which provided cues. Consistent with our findings, subjects with prolonged and/or inaccurate stepping responses were more likely to report a history of accidental falls.

**The Relationships Between Complex RT_{clin} Accuracy: Simple RT_{clin} Latency and Mobility Outcomes Provide a Mechanism By Which Poor Executive Function Increases Fall Risk**

The second clinically relevant aspect of this research is that the data link complex RT_{clin} accuracy, simple RT_{clin} latency, and their ratio to UST and frontal plane responses to sequential walking surface perturbations in older subjects with DPN. Although prior work has linked executive function with gait speed and gait variability,\(^37\) a recent review\(^32\) found minimal research investigating the influence of executive function on balance and response to perturbation as reported here. One potential explanation for this relationship is that the subjects with poor complex RT_{clin} accuracy: simple RT_{clin} latency are unable to rapidly inhibit attention to irrelevant external stimuli and distracting internal cognitive processes within the 420 ms available, and so are similarly unable to inhibit preplanned lower limb responses while walking and cannot quickly adjust swing limb trajectories within the time available. This uneven surface would then lead to intermittent missteps in relation to the center of mass with the subsequent development of lateral momentum and the need for a lateral recovery step. In support of this reasoning, Asai et al.\(^38\) found that lateral, but not sagittal, momentum was less effectively attenuated during a distracting task while walking, suggesting that lateral control is most heavily reliant on attention. Perhaps most compelling, Sturnieks et al.\(^39\) found that executive functioning was an independent predictor of the need to take a step after lateral waist pull perturbations, whereas standing sway and lower limb strength predicted anterior perturbations. This mirrors our findings that HipST\(^{56}\);Ank\_PRO predict sagittal plane characteristics of step length and speed on the uneven surface,\(^19\) whereas complex RT_{clin} accuracy: simple RT_{clin} latency predict frontal plane gait variability. Finally, the increase in
extreme lateral step placement on the uneven surface associated with poor complex RT<sub>clin</sub> accuracy:simple RT<sub>clin</sub> latency provide a mechanism by which poor or slow inhibitory executive function can lead to frontal plane instability and predispose to more severely injurious falls.  

The Relationships Between Complex RT<sub>clin</sub> Accuracy:Simple RT<sub>clin</sub> Latency and Mobility Were Prominent Only in Subjects With DPN, Suggesting Greater Importance in the Setting of Neuromuscular Impairments

The third novel feature of this research is that the relationships between the RT<sub>clin</sub>Dev-derived short latency neurocognitive attributes and UST, response to gait perturbations on the uneven surface, and major injuries were identified solely in the subjects with DPN. These relationships did not seem to be present in the older subjects with normal lower limb neuromuscular function and minimally in subjects with DPN on a smooth surface. One possible explanation for this may be that subjects with DPN and its associated lower limb neuromuscular limitations are more reliant on cortical control of balance and gait and posture in challenging situations than subjects without DPN. If so, the ability to quickly inhibit attention to less relevant stimuli and internal cognitive processes as measured by complex RT<sub>clin</sub> accuracy:simple RT<sub>clin</sub> latency would offer an advantage with regard to immediacy of response to perturbation. In support of this, the prefrontal cortex, the region associated with executive cognitive functions, shows increased activity during challenging activities such as adapting to different walking speeds on a treadmill, maintaining balance while standing on a suddenly translating surface, and in patients with cerebellar/brainstem strokes while walking at a uniform speed compared to controls. Other supportive work finds that the effect of executive function on mobility is greatest in older adults with lower limb sensorimotor changes and that distraction, a sign of executive impairment, interferes with the calibration of ankle muscle response to perturbation in older subjects approximately 400 ms after perturbation and so increases the likelihood of taking multiple steps. Together, these studies suggest that the prefrontal cortex and its associated executive functions are most essential during challenges to balance, particularly when subcortical systems are disrupted or desynchronized by altered peripheral sensory function or weakness, as is the case for the DPN subjects reported here.

The results may have clinical use. The ability to respond quickly and the capacity to rapidly suppress irrelevant stimuli are difficult for the clinician to quantify. However, RT<sub>clin</sub>Dev may offer a clinically accessible method for doing so. If so, measurement of simple RT<sub>clin</sub> latency and complex RT<sub>clin</sub> accuracy may be useful in the evaluation of patients with lower limb neuromuscular impairments, which increase their fall risk. The presence of neuromuscular as well as neurocognitive impairments may then mark the patient as high risk for falls and injury. Potential interventions would target proximal strengthening as well as cognitive procedures that enhance attention and mental focus. There is evidence that interventions such as mindfulness training, discontinuing of sedating medications, and/or the addition of activating medications, may lead to laudable changes in gait that suggest reduced fall risk. If neither is possible, then environmental modification with reduction in surface irregularities and distracters while walking is indicated.

Although the study’s strengths and innovative features have been described, enthusiasm must be reserved given the study’s limitations. The greatest weakness is the limited subject numbers, asymmetry of subjects with and without DPN, and the absence of power calculations, consistent with this being a secondary analysis of data obtained previously for other purposes. Furthermore, complex RT<sub>clin</sub> accuracy and simple RT<sub>clin</sub> latency have not been validated specifically within older people with DPN. However, it seems unlikely that the presence of DPN had influence, as post hoc testing revealed no significant or near-significant DPN/non-DPN group differences for any of the 3 measures (p values of 0.992, 0.129, and 0.464 for, respectively, complex RT<sub>clin</sub> accuracy, simple RT<sub>clin</sub> latency and their ratio). Formal neuropsychologic testing was not performed and so the inclusion of subjects with mild cognitive impairment is possible. Additionally, subject moods and personality traits were not evaluated, and so these may have influenced the results. Despite these, the strength of the associations between UST and step width variability, and complex RT<sub>clin</sub> accuracy and simple RT<sub>clin</sub> latency and their ratio are so strong in the subjects with DPN that the likelihood of a spurious relationship seems small. In contrast, with only 2 DPN group major injuries during the 1 year of prospective follow-up, the possibility of a chance association with respect to that outcome is clearly present, and the results need to be replicated before acceptance is considered. Weak associations between short latency neurocognitive attributes and the three mobility outcomes may have
been missed in the non-DPN group, a problem likely accentuated by the limiting of UST to a maximum of 30 seconds, which could have obscured correlational analyses for these subjects. However, it should be noted that Hip^{STR,Ank}_{PRE} demonstrated a significant relationship with UST in this group, despite the same limitations of this outcome (Table 2B).

In summary, this preliminary study suggests that complex RT_{clin} accuracy, simple RT_{clin} latency, and their ratio were potent predictors of UST, frontal plane gait variability in response to perturbations while walking, and associated with major fall injury in older subjects with DPN. The findings suggest a plausible mechanism by which impairments in inhibitory executive function can increase fall and injury risk. These neurocognitive variables, which can be obtained at the bedside or in the clinic using RT_{clin,Dev}, may be combined with evaluation of critical lower limb neuromuscular attributes^45 so as to allow the clinician a more comprehensive understanding of fall risk.

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Supplementary Checklist

STROBE Checklist: http://links.lww.com/PHM/A315

REFERENCES


