Sensor-based Characterization of Daily Walking: A New Paradigm in Frailty Assessment

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Abstract

Background Frailty is an increasingly recognized geriatric syndrome resulting in decline in reserve across multiple physiological systems. Impaired physical function is a prime indicator of frailty.

Objective The goal of this study was to develop an algorithm that discriminates between frailty groups (non-frail, pre-frail, and frail) based on gait performance parameters derived from unsupervised daily physical activity (DPA).

Methods DPA was acquired for 48 hours from older adults (≥65 years) using a tri-axial accelerometer motion-sensor. Purposeful continuous bouts of walking (≥60s) without pauses were identified from acceleration data. These were then used to extract qualitative measures (gait variability, gait asymmetry, and gait irregularity) and quantitative measures (total continuous walking duration and maximum number of continuous steps) to characterize gait performance. Association between frailty and gait performance parameters was assessed using multinomial logistic models with frailty as the dependent variable, and gait performance parameters along with demographic parameters as independent variables.

Results 126 older adults (44 non-frail, 60 pre-frail, and 22 frail, based on the Fried index) were recruited. Step- and stride-times, frequency domain gait variability, and purposeful walking quantitative measures were significantly different between non-frail and pre-frail as well as non-frail and frail groups ( p <0.05). Using the logistic model pre-frail/frail group (vs. non-frail) was identified with 76.8% sensitivity and 80% specificity.

Discussion Everyday walking characteristics were found to be associated with frailty. Along with quantitative measures of physical activity, qualitative measures are critical elements representing the early stages of frailty. In-home gait assessment offers an opportunity to screen for and monitor frailty.
Background

According to World Health Organization (WHO) reports, the global population of older adults aged 60 years or older was 600 million in 2000, and is expected to reach two billion by 2050 (1). Among this population, frailty is an increasingly recognized syndrome that is associated with decline in function and reserve across multiple physiologic systems (2-4). Frailty is characterized by a high vulnerability to adverse health outcomes such as disability, falls, hospitalization, institutionalization, and mortality (2). Reduction or impairment of physical function is a prime indicator of frailty (3), and frailty is one of the major reasons for falls in old age (5-11). However, there is currently no objective method for assessing frailty that incorporates assessment of daily physical activity (DPA). DPA data has been recently used to assess physical function, especially with the help of wearable sensor technology. Using wearable devices, it is possible to continuously measure DPA in the least intrusive manner and for longer durations of time in the participants’ natural environment. Among several DPA, motion analysis of the trunk during walking is known to provide insights regarding neuromuscular deficits caused by frailty and aging (11). In our previous studies, among DPA measures (walking, standing, sitting, and lying), quantitative parameters related to walking, such as total continuous walking duration and maximum number of continuous steps, best discriminated between non-frail and pre-frail groups, with highest effect sizes for the number of steps and the percentage of walking duration within a 24-hour time period (12). While promising, we found that none of these outcomes (or their combination in a multinomial logistic analysis) could significantly discriminate between frailty groups when adjusted for age (12).

In this study, we aimed to improve detection of frailty-related neuromuscular deficits based on gait performance parameters derived from unsupervised DPA. Previous studies have used trunk motion data from supervised in-lab gait tests to characterize
sensorimotor gait performance among frail elders including gait variability, asymmetry, initiation, and irregularity (10,11,13–17). We hypothesized that using more robust measures of unsupervised DPA gait performance (instead of number of steps) it would be possible to distinguish between non-frail and pre-frail older adults.

Methods

This observational cross-sectional study was performed at Arizona Center on Aging, Tucson, AZ. Participants in this study were from primary, secondary, and tertiary healthcare settings within our academic network and also from community providers and aging service organizations. DPA was recorded from eligible volunteers for 48 hours and the walking data from the DPA was processed to study the gait performance parameters and associate these characteristics with frailty.

Participants

Older adults (65 years or older), without severe mobility disorder and the ability to walk at least 10m with or without an assistive device, were considered eligible for the study. Participants with dementia identified by a Mini-Mental State Examination (MMSE) (18) score of <23 or terminal illness were excluded. All the eligible participants signed a written consent form according to the principles expressed in the Declaration of Helsinki (19), approved by the Institutional Review Board of the University of Arizona.

Demographic and Clinical Measures

The recorded clinical measures included self-reported history of falls, use of assistive device and the number of prescriptions. Interviewer-administered questionnaires included the MMSE, Mobility-Tiredness Scale (20), Center for Epidemiologic Studies Depression Scale (CES-D) (21), Falls Efficacy Scale-International (22), and Barthel Activity of Daily Living (ADL) Scale (23).

Gold Standard Frailty Assessment
Frailty was assessed using the five criteria proposed by Fried et al (3), including: self-reported weight loss, weakness measure by the grip strength, self-reported exhaustion, slowness measure by the walking test, and self-reported low energy expenditure. A score of one point was given for each criterion recorded, totaling a score in the range of 0–5. Frailty was categorized as follows: non-frail (score 0), pre-frail (score 1–2), and frail (score 3–5).

Sensor-Based Daily Physical Activity Assessment

DPA was quantified for two consecutive days (48 hours) using a tri-axial accelerometer sensor (PAMSys, BioSensics Cambridge, MA, USA) fixed in a tee-shirt, with a device pocket located at the sternum. PAMSys is a small (5.1 x 3 x 1.6 cm), light (24g) recording system containing inertial sensors. We used a previously validated algorithm to identify walking bouts (PAMWare, BioSensics Cambridge, MA, USA). Briefly, walking bouts were defined by a minimum of three successive steps (24), where steps were estimated by the detection of an acceleration peak beyond a predefined threshold after filtering the signal (25). Using this software several gait parameters were derived including: the duration of walking, walking bout times (duration of each walking episodes), number of steps per walking bout, and walking cadence per bout. PAMWare is 87% sensitive and 87% specific for gait detection (24,26). Within the current study we further refined the PAMSys algorithm to derive assess purposeful walking, as well as gait performance parameters.

Purposeful Walking Bouts

To provide sufficient sample size, purposeful walking bouts, defined as 60 seconds or longer, were used for extraction of gait performance parameters (27,28). This assured that for each analyzed walking event, the participant is commuting with the purpose of getting to a certain destination point, rather than random daily walking. Gait performance parameters including time- and frequency-domain gait variability, gait asymmetry, and
gait irregularity were extracted from purposeful walking bouts, continuous for 60 seconds or longer with no pauses longer than 1.7s between gait cycles (12). Allowable 1.7s pause between gait cycles was conservatively selected based on the average plus standard deviation stride time duration observed in frail participants (12). All the sensor-based gait performance outcome measures are shown in Table 1. For each purposeful walking bout, the raw vertical acceleration signal was filtered using a second order Butterworth filter (cut-off frequency of 2.5Hz (29)), and the peaks of the filtered acceleration signal were detected using a peak-detection algorithm. The time-interval between two consecutive peaks was defined as the step-time, and the time-interval between alternate peaks was defined as the stride-time.

**Gait Variability:** We defined gait variability as the stride-to-stride fluctuation in walking cycles, which has been associated with high risk of fall and cognitive impairments in elders (5,30–32). Gait variability reflects inconsistency in physiological systems that regulate walking, including neuromuscular, reflexive postural control, and cardiovascular systems (33). We used two methods to assess gait variability: 1) step- and stride-time variability using time-domain; and 2) power spectral density (PSD) using frequency-domain analysis (34,35). Step- or stride-time variability was calculated as the coefficient of variation of the series of step- or stride-times for each purposeful walking bout. For PSD analysis, the power spectrum of the acceleration data was calculated using Welch’s averaged modified periodogram method (36), to represent the frequency components of the acceleration signal (37). We used a window size of 512 samples and an FFT length of 2-times the next higher power of the window size (36). An overlap of 50% was considered between the windows. The locomotion band between 0.5–3.0Hz was analyzed (36). PSD components were extracted from the raw acceleration signal, including: maximum PSD peak, PSD width (full width at half maximum height), PSD slope (PSD width to the peak)
and dominant walking frequency. A higher variability in walking was identified by a shorter and wider PSD peak.

**Gait Asymmetry:** When gait becomes less automatic due to sarcopenia and cognitive aging, left-right step coordination may require more effort, especially among frail individuals (12,38,39). Further, studies showed that no strong association between gait variability and asymmetry exists, suggesting that asymmetry reflects an independent measure of gait impairments due to distinct pathological causes (12,38). Here, step asymmetry was obtained from the autocorrelation function of the vertical acceleration signal (12,38), represented by a sequence of autocorrelation coefficients over increasing time lags.

\[
\text{Asymmetry}_1 = \frac{A_{d1}}{A_{d2}}, \quad \text{Asymmetry}_2 = \frac{|A_{d1} - A_{d2}|}{\max(A_{d1}, A_{d2})}
\]  

(1)

where Ad1 and Ad2 are the prominence of the first and the second peaks respectively after the central (zero lag) peak (40).

**Gait Irregularity:** Results from supervised gait studies showed that irregularity measures can describe predictability of walking cycles, which can be influenced by both neurological and neuromuscular diseases (41–44). Further, within in-lab settings, it has been demonstrated that gait irregularity can differ between non-frail and pre-frail older adults (16). We used Sample Entropy (SampEn) assessment defined as equation (2), where A was defined as the number of matches in the filtered acceleration signal length m+1 (distance function smaller than tolerance r: d[Xm+1(i),Xm+1(j)]<r) and B as the number of matches of length m: (d[Xm(i),Xm(j)]<r) (45–49).

\[
\text{SampEn} = -\log\left(\frac{\sum_{i=1}^{N-m} A_i}{\sum_{i=1}^{N-m} B_i}\right) = -\log A/B
\]  

(2)
The time-delay of the signal was calculated using mutual information method for all the purposeful walks (50), and the average time-delay of all the purposeful walks was used to calculate the SampEn for each volunteer. We used embedding dimension $m = 3$, and tolerance $r = 0.2$ times the standard deviation of the signal, which are commonly used to compute sample entropy of gait signal (45–49).

Table 1. Sensor based outcome measures
| Parameter                        | Description                                                                 |
|---------------------------------|-----------------------------------------------------------------------------|
| Step/stride time                | Time-interval between two consecutive/alternate acceleration peaks           |
| **Gait Variability**            |                                                                             |
| Step/stride time variability    | Coefficient of variation (\%), standard deviation of step/stride time over   |
|                                 | mean step/stride time                                                       |
| PSD max                         | Maximum height of the PSD distribution curve representing the amount of walking that occurs at the dominant frequency |
| PSD width                       | The width of the PSD curve at half of the maximum height representing the range of walking frequencies |
| PSD slope                       | The slope of the PSD curve from the peak to the width representing the rate of change of walking frequencies |
| Dominant frequency              | The frequency at which the PSD curve attains its peak, representing the frequency at which most of the walking cycles occur |
| **Gait Asymmetry**              |                                                                             |
| Unbiased auto-correlation       | Unbiased auto-correlation coefficients of gait signal, representing left-right step coordination |
| **Gait Irregularity**           |                                                                             |
| Sample entropy                  | Sample entropy, representing the predictability of walking cycles            |
| **Purposeful Walking Quantitative Measures** |                               |
| No. of purposeful walks         | Total number of purposeful walks during 48 hours                             |
| Total continuous walking duration| Total duration of continuous purposeful walks during 48 hours                |
| Max walking bout                | Maximum bout of purposeful continuous walking in 48 hours                    |
| Max no. of continuous steps     | Maximum number of continuous steps in the longest continuous walking bout in 48 hours |
| Walking bout variability        | Coefficient of variation (\%), standard deviation of walking bouts over mean walking bout |

**PSD – Power Spectral Density**

*Purposeful Walking Quantitative Measures*: In addition to the above-mentioned features,
we extracted the following parameters in each purposeful continuous walking event: maximum walking bout, maximum number of continuous steps, walking-bout variability (coefficient of variation in walking bouts duration within 48 hours), and total duration of purposeful walks. Of note, the parameters extracted here were only obtained for continuous purposeful walking events with no pause of 1.7s or longer, as described above.

Statistical Analysis

Separate analysis of variance (ANOVA) models were performed to compare sociodemographic parameters between the three Fried frailty groups. To explore differences in gait performance parameters among frailty categories, univariate ANOVA models were used with each gait performance parameter as the dependent variable, and the Fried frailty categories (non-frail, pre-frail, and frail) as the independent variable. Subsequently, gait performance parameters were used in a single multivariable nominal logistic model to assess the association between frailty categories and DPA gait performance parameters. In this model we combined pre-frail and frail groups, due to the limited number of frail participants. The model was developed following these steps: 1) univariate nominal logistic model analysis of the gait performance parameters as independent variables was performed. Gait performance parameters with significant association with frailty were considered for subsequent steps; 2) collinearity between the various gait performance parameters was tested using the variance inflation factor (VIF) index. VIF value greater than 10 represented the presence of collinearity (55); and 3) gait performance and demographic parameters were selected based on Akaike information criterion (AIC) values. Participants who exhibited no 60-second purposeful walk were automatically categorized as frail. All analyses were done using JMP (Version 11; SAS Institute Inc., Cary, NC, USA), and statistical significance was concluded when \( p < 0.05 \).

Results
Demographic and Clinical Measures

The study involved a total of 126 participants, among whom 44 were non-frail, 60 were pre-frail and 22 were frail according to the Fried frailty criteria (3). Table 2 shows demographic and clinical characteristics. Pre-frail and frail participants were significantly older than non-frail participants, used assistive devices more frequently, and had higher BMI, history of falls, number of prescriptions, perceived tiredness, and fear of falling score (p< 0.05). When compared to non-frail and pre-frail participants, frail individuals had significantly higher levels of depressive symptoms, fear of falling, perceived tiredness, and lower performance scores in ADL (p< 0.05).

Table 2. Demographic and clinical characteristics

| Characteristics                  | Non-frail (N) | Pre-frail (P) | Frail (F) | p N vs P |
|----------------------------------|---------------|---------------|-----------|----------|
| Age (years)                      | 74.6 ± 6.5    | 79.72 ± 8.6   | 82.81 ± 9.7 | 0.001    |
| Height (cm)                      | 161.42 ± 6.30 | 161.62 ± 9.32 | 161.16 ± 11.28 | 0.901    |
| Weight (kg)                      | 66.67 ± 12.82 | 74.53 ± 18.27 | 78.83 ± 20.99 | 0.032    |
| Body mass index                  | 25.53 ± 4.25  | 28.49 ± 6.41  | 30.17 ± 6.30  | 0.019    |
| Gender                           |               |               |           | 0.262    |
| Male                             | 7(15.9)       | 15(25.0)      | 3(14.3)    |          |
| Female                           | 37(84.1)      | 45(75.0)      | 18(85.7)   |          |
| History of falls                 | 13(33.3)      | 26(47.3)      | 11(57.9)   | 0.177    |
| Falls Efficacy Scale - International | 20.8 ± 4.2 | 28.0 ± 9.5   | 39.5 ± 12.3 | <0.001   |
| Use of assistive devices         | 4(10.0)       | 26(47.3)      | 14(73.7)   | <0.001   |
| Mobility-tiredness scale         | 5.6 ± 0.8     | 4.7 ± 1.4     | 2.7 ± 1.8  | <0.001   |
| Number of prescriptions          | 2.5 ± 1.8     | 4.1 ± 3.8     | 6.0 ± 3.4  | 0.021    |
| MMSE                             | 29.2 ± 1.1    | 28.6 ± 1.6    | 28.7 ± 1.7 | 0.121    |
| CES-D                            | 6.6 ± 5.7     | 6.9 ± 6.8     | 14.0 ± 7.0 | 0.958    |
| Barthel ADL Scale                | 95.5(14.0)    | 96.2(5.5)     | 88.1(9.0)  | 0.955    |

Results presented as mean ± SD or number (%). Bold-faced values show statistical significance (p<0.05).

Sensor-Based Daily Physical Activity Assessment

Table 3 shows the sensor-based DPA parameters for purposeful walking with resulting p-values of one-way ANOVA test (Figure 1, Table 3). Of note, among participants, four non-frail (9.09%), 17 pre-frail (28.33%), and 11 frail (50%) had no walking bout equal or longer
than 60 seconds.

Table 3. Purposeful gait performance parameters and one-way ANOVA results between non-frail(N), pre-frail(P), and frail(F) groups

| Parameter                                      | Non-frail (N) | Pre-frail (P) | Frail (F) |       |
|------------------------------------------------|---------------|---------------|-----------|-------|
| **Temporal Gait Parameters**                   |               |               |           |       |
| Step-time (s)                                  | 0.56± 0.05    | 0.61± 0.06    | 0.62± 0.036 | <     |
| Stride-time (s)                                | 1.13± 0.09    | 1.22± 0.12    | 1.24± 0.13 | <     |
| **Time Domain Gait Variability**               |               |               |           |       |
| Step variability (%)                          | 10.79± 2.80   | 10.83± 3.48   | 11.43± 2.96 | (     |
| Stride variability (%)                        | 9.16± 2.94    | 8.83± 3.19    | 8.40± 2.43 | (     |
| **Frequency-domain Gait Variability**          |               |               |           |       |
| PSD max (W/Hz)                                 | 0.17± 0.16    | 0.07± 0.06    | 0.07± 0.10 | <     |
| PSD width (Hz)                                 | 0.22± 0.10    | 0.21± 0.03    | 0.21± 0.02 | (     |
| PSD slope (W)                                  | 1.24± 1.21    | 0.49± 0.46    | 0.44± 0.62 | <     |
| Dominant frequency (Hz)                        | 1.90± 0.16    | 1.74± 0.18    | 1.70± 0.18 | <     |
| **Gait Asymmetry**                             |               |               |           |       |
| Asymmetry 1                                    | 1.10± 0.14    | 0.24± 0.07    | 0.07± 0.03 | (     |
| Asymmetry 2                                    | 0.09± 0.07    | 0.08± 0.07    | 0.07± 0.03 | (     |
| **Gait Irregularity**                          |               |               |           |       |
| Time delay (ms)                                | 145.39± 14.90 | 156.93± 25.85 | 153.59± 14.21 | (     |
| Sample entropy (bits)                          | 0.93± 0.28    | 0.99± 0.29    | 1.07± 0.27 | (     |
| **Purposeful Walking Quantitative Measures**    |               |               |           |       |
| Number of purposeful walking                   | 13.25± 11.22  | 10.63± 10.43  | 6.36± 5.87 | (     |
| Total continuous walking duration (s)           | 4042.3± 3012.8 | 2436.7± 1988.4 | 1795.9± 1659.4 | (     |
| Max walking bout (s)                           | 3 ± 6         | 9 ± 12        | 2 ± 2      | (     |
| Max number of continuous steps                 | 475.62± 512.27 | 216.98± 228.95 | 115.85± 47.61 | (     |
| Walking bout variability (%)                   | 1867.5± 1735.9 | 896.63± 1055.5 | 461.45± 420.15 | (     |
|                                                   | 8 ± 8         | 3 ± 3         |           | (     |
|                                                   | 252.74± 110.51 | 195.88± 73.98 | 178.52± 31.43 | (     |

PSD – Power Spectral Density

Among gait parameters, frequency domain gait variability parameters and quantitative measures including maximum number of continuous steps and total continuous walking duration best discriminated non-frail and pre-frail, as well as non-frail and frail individuals. Step- and stride-time were significantly different between non-frail vs pre-frail and frail groups ($p<0.001$). Further, among the frequency domain gait variability parameters, PSD amplitude, PSD slope, and the dominant frequency were significantly different between non-frail and pre-frail, and non-frail and frail groups ($p<0.05$). However, step- and stride-time variabilities did not show statistical significance between the three groups ($p>0.1$).
Gait asymmetry, representing left and right step coordination, was not statistically significant between frailty groups ($p>0.1$). Similarly, gait irregularity, measuring predictability of walking cycles, displayed an increasing trend towards pre-frailty and frailty; but the observed between-group differences were not significant ($p>0.1$). All the purposeful walking quantitative measures showed differences between the three groups, and most of these differences were statistically significant between non-frail and pre-frail, as well as non-frail and frail groups ($p<0.05$).

**Frailty Prediction using Gait Performance Parameters**

A multinomial logistic regression model was developed using gait performance parameters extracted from DPA along with age and BMI to predict frailty. A step-wise logistic model was developed with frailty groups as the dependent variable (non-frail vs. pre-frail/frail), for which, age (years), BMI (kg/m$^2$), stride-time variability (%), dominant frequency (Hz), and maximum number of continuous steps were selected as independent variables (Table 4). The logistic regression model developed with these features was able to predict pre-frail/frail category with an improved receiver operating characteristics (ROC) area under curve (AUC) of 0.84 compared to age (ROC AUC: 0.71) and total number of steps for 48 hours (ROC AUC: 0.77, Table 5). The ROC curves are shown in Figure 2.

| Parameter | Parameter Estimate | Std. Error | $\chi^2$ | $p$-value |
|-----------|--------------------|------------|---------|-----------|
| Age (years) | -0.1191           | 0.04       | 8.27    | 0.004     |
| BMI (kg/m$^2$) | -0.1772           | 0.06       | 8.49    | 0.004     |
| Stride variability (%) | -0.2507           | 0.11       | 5.39    | 0.020     |
| Dominant frequency (Hz) | 6.6265            | 2.14       | 9.62    | 0.002     |
| Max no. of continuous steps | 0.0001           | 0.00       | 0.33    | 0.565     |

**Table 5. Logistic model performance comparison of different parameters for 80% specificity**

| Model Features | Accuracy | Sensitivity | Specificity | AUC  |
|----------------|----------|-------------|-------------|------|
| Age            | 65.1%    | 58.1%       | 80%         | 0.71 |
| Total number of steps | 74.6%    | 46.5%       | 80%         | 0.77 |
| Gait performance parameters | 77.7%    | 76.8%       | 80%         | 0.84 |

**Discussion**

As hypothesized, several sensor-based gait performance parameters significantly
discriminated between non-frail and pre-frail groups, even when adjusted with age. In our previous studies the total number of steps and walking duration could not significantly discriminate pre-frail participants from non-frail when adjusted with age (12). This improvement in frailty assessment occurred due to utilization of the novel concept of purposeful walking for obtaining both qualitative and quantitative gait performance parameters.

**Purposeful Continuous Walks**

The ability to walk longer distances is instrumental for humans to perform various activities of daily living and lead an independent life. Walking requires complex mechanisms within the human sensory motor system to provide the necessary timing, coordination, and balance such that the interplay between the center of mass and the base of support are regulated in a repetitive manner (46,56). Previous studies that explored the gait characteristics of non-disabled adults for two weeks to define walking duration, found that 81% of all walking bouts lasted about 60 seconds (27,28). Accordingly, in our study, we used purposeful continuous walking bouts of 60 seconds or longer to provide sufficient sample for extracting gait performance parameters. Previous studies also showed that accelerometer-derived gait performance measures based on three-day daily activities could improve fall risk evaluation in older adults, when 60-second continuous walking periods were implemented compared to overall number of steps (28). Similarly, in the current study, purposeful walking with no pauses resulted in better discrimination between the frailty groups ($p<0.05$), when compared to walks that included pauses.

**Advantages of Qualitative Gait Parameters**

In addition to previously reported quantitative parameters of gait (number of steps, mean walking bout duration, and longest walking bout duration) (12), here we extracted
qualitative gait performance parameters (gait variability, asymmetry, and irregularity). Specifically, gait variability represented a promising measure for differentiating gait deficits among the three frailty categories. Gait variability, defined as the stride-to-stride fluctuation in walking cycles, has been previously associated with high risk of fall and cognitive impairments in elders (5,30–32,57). Gait variability reflects inconsistency in physiological systems that regulate walking, including neuromuscular, reflexive postural control, and cardiovascular systems. We used two methods to assess gait variability: step/stride time variability using time-domain and power spectral density (PSD) using frequency-domain analyses. We observed that the frequency-domain parameters were significantly different between frailty groups, while time-domain parameters were not. Owing to the low sampling frequency in this study, some of the information content may be lost due to filtering for peak detection (58). Hence, the PSD analysis performed on the entire raw acceleration signal may provide a more efficient tool for assessing gait variability for low sampling frequency motion sensor data.

Additionally, gait asymmetry and irregularity were also investigated here, as parameters that representing gait deficits independent of gait variability. Gait asymmetry, representing left-right step coordination, has been used as a metric to observe walking patterns in older individuals. Cognitive aging and sarcopenia render gait to be less automatic and left-right symmetry co-ordination is expected to require additional effort, especially in frail individuals (39). Although gait asymmetry showed a decreasing trend from non-frail to pre-frail and frail samples, differences were not significant (Table 3). Since loss in step-coordination may also happen due to hip, knee, or ankle impairments (59), these confounding variables can mask the effect of frailty. Further, gait irregularity, representing the predictability of walking cycles, and can be influenced by both neurological and neuromuscular diseases (13,42–44). Previous studies have used sample
entropy to obtain gait predictability or repeatability to investigate differences in the relationship between executive function, and gait variability and stability during single and dual-task walking in persons with and without dementia (43). We computed gait irregularity using the sample entropy method and observed an increase in irregularity in pre-frail/frail population but there was no statistical significance seen between the groups (Table 3).

Limitations
There are limitations to consider in the interpretation of these findings. The first limitation is adherence to the wearable sensor equipment. Though the participants did not express any obvious discomfort while wearing the sensors, it is possible that a few of the participants forgot to wear them immediately after a shower. Second, our sample was predominantly women. Although we did not observe a gender specific difference in gait performance, the model developed here may have limited generalizability to a population with a more balanced gender composition. Finally, not all participants, especially pre-frail and frail, had 60 seconds or longer purposeful continuous walking during the 48 hours of data collection. This in turn, reduced our frail group sample size, which more likely resulted in non-significant gait performance parameter differences between pre-frail and frail groups. To overcome this limitation, we combined pre-frail and frail participants and all the participants who did not exhibit purposeful continuous walking were automatically categorized as pre-frail/frail while developing the logistic model. Since detecting the onset of frailty at the pre-frail stage is most crucial in recovering health status (60), our findings suggested a promising method for pre-frailty identification using DPA data.

Summary And Conclusion
Using gait performance parameters within 48 hours of daily monitoring, pre-frailty/frailty was identified with a sensitivity of 76.8% and specificity of 80% among elders. Findings
suggest that purposeful walking parameters, including gait variability and the amount of continuous walking, may noticeably improve gait deficit assessment compared to previous methods of step counting. The proposed gait performance characterization based on sensor-based daily physical activity provides potential for being integrated into clinical care for in-home screening of frailty (much as a holter monitor is used) to provide information pertaining to an individual’s condition before hospital admission, or when frailty is suspected. This method is advantageous over its in-clinic counterpart as it is objective rather than subjective self-reported measures of physical function, and is measured in real-world walking activities rather than an artificial clinical setting. Healthcare research in wearable devices has been constantly growing in various areas like remote patient monitoring and healthcare (61–65), wearable sensor-based systems for health monitoring (66–69), and ambient-assisted living tools for older adults (70). For elderly populations requiring continuous health monitoring, sensor-based wearables and remote monitoring will help eliminate the hassle of periodic commute to diagnostic centers, reduce the amount of recurring admissions to the hospital, and facilitate more efficient clinical visits with objective results (71).

**Abbreviations**

ADL: Barthel activity of daily living; AIC: Akaike information criteria; ANOVA: Analysis of variance; AUC: Area under the curve; BMI: Body mass index; CES-D: Center for Epidemiologic Studies Depression Scale; DPA: Daily physical activity; MMSE: Mini-mental state examination; PSD: Power spectral density; ROC: Receiver operating characteristic; SampEn: Sample entropy; SD: Standard deviation; VIF: Variance inflation factor; WHO: World health organization.

**Declarations**
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Availability of data and materials
The datasets used and/or analysed during the current study available from the corresponding author on reasonable request.

Authors’ contributions
DPK: data analysis, interpretation of findings, statistical analysis, manuscript writing and critical review of manuscript. NT: concept, study design, subject recruitment, study management, data analysis and interpretation, statistical analysis, manuscript writing and editing, critical review of manuscript and approval of final manuscript. JM: subject recruitment, study management, study design, interpretation of findings, critical review of manuscript and approval of final manuscript. HE: interpretation of findings, critical review of manuscript, comments and approval of final manuscript. KL: concept, study design, data analysis and interpretation, statistical analysis, manuscript writing and editing, critical review of manuscript and approval of final manuscript.

Competing interests
The authors declare that they have no competing interests.

Consent for publication
Not applicable.

Ethics approval and consent to participate
The study was approved by the University of Arizona and Banner Sun Health Research Institute’s Institutional Review Boards. Before participation, written informed consent according to the principles expressed in the Declaration of Helsinki was obtained from all subjects.

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Figures
Figure 1

Comparison of purposeful gait performance parameters between non-frail (N), pre-frail (P), and frail (F) groups (*p<0.05)
Figure 2

Logistic regression model ROC curves for age, total number of steps, and gait performance parameters.