Frailty Assessment Using a Novel Approach Based on Combined Motor and Cardiac Functions: A Pilot Study

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Abstract

Background: Previous research showed association between frailty and an impaired autonomic nervous system; however, the direct effect of frailty on heart rate (HR) behavior during physical activity is unclear. The purpose of the current study was to determine the association between HR increase and decrease with frailty during a localized upper-extremity function (UEF) task to establish a multimodal frailty test.

Methods: Older adults aged 65 or older were recruited and performed the UEF task of rapid elbow flexion for 20 seconds with the right arm. Wearable gyroscopes were used to measure forearm and upper-arm motion, and electrocardiography were recorded using leads on the left chest. Using this setup, HR dynamics were measured, including time to peak HR, recovery time, percentage increase in HR during UEF, and percentage decrease in HR during recovery after UEF.

Results: Fifty-six eligible participants were recruited, including 12 non-frail (age=76.92±7.32 years), and 44 pre-frail/frail (age=81.23±8.14 years). Analysis of variance models showed that the percentage increase in HR during UEF and percentage decrease in HR during recovery were both 47% smaller in pre-frail/frail older adults compared to non-frails ($p<0.01$, effect size = 1.42 and 1.17 for increase and decrease percentages). Using logistic models with both UEF kinematics and HR parameters as independent variables, frailty was predicted with a sensitivity of 0.82 and specificity of 0.83.

Conclusion: Current findings showed evidence of strong association between HR dynamics and frailty. It is suggested that combining kinematics and HR data in a multimodal model may provide a promising objective tool for frailty assessment.

Introduction

The concept of frailty is used to identify older adults with low physiological reserves, leading to vulnerability to illness, and increased risk of institutionalization and mortality [1, 2]. Muscle loss and weakness (sarcopenia and dynapenia) are the main symptoms of frailty, caused by inflammatory (elevated interleukin 6 (IL-6), C-reactive protein (CRP), tumor necrosis factor alpha (TNFα)), metabolic (deficiencies of various mitochondrial subunits), and hormonal derangements (cortisol and testosterone) that shift homeostasis from an anabolic to a catabolic state [3–11]. Previous research also showed association between frailty and an impaired autonomic nervous system (ANS) because of alterations in electrical conduction and action potential morphology [12, 13]. The presence of a compromised neurohormonal homeostasis associated with frailty as measured by ANS dysfunction is, in turn, associated with health complications and mortality [14–17].

Heart rate variability (HRV: variability in RR intervals) and HR complexity (entropy analysis) during resting have been used for assessing ANS dysfunction and proposed as a vital sign [18–20]. Although resting HRV provides information about abnormal ANS performance, it may not be directly associated with HR increase or recovery in response to physical activities, which show a natural decline with age [21]. Further, between-subject variability exists in resting HR/HRV data (e.g., due to breathing regulation and...
environmental factors [22–24]). In our previous research we measured HR dynamics defined as HR increase and recovery parameters during and after walking, and investigated the association between these parameters with frailty [25]. We observed that non-frail participants had significantly larger and faster increases in HR during walking, compared to pre-frail/frail older adults [25].

Based on the previous evidence, the aim of the current study, was to establish and validate a platform for simultaneous assessment of motor and cardiac function to assess HR dynamics and predict frailty in community dwelling older adults. For the motor function we have previously validated an upper-extremity function (UEF) test, including rapid elbow flexion, to accurately detect systematic decrements in function, including slowness, weakness, inflexibility, and fatigue.[26, 27]. We have validated the UEF motor test for discriminating between frailty groups, among both community dwelling older adults and bed-bound trauma patients, using the Fried frailty index and the short-version Rockwood questionnaire as a comparators [28–30]. The hypotheses for the current work were: 1) HR dynamics due to UEF would be significantly associated with frailty; and 2) a combined model including both motor and HR parameters would more strongly be associated to frailty compared to models incorporating only one of these individual measures.

**Methods**

**Participants**

Older adults were recruited from the primary, secondary, and tertiary health care settings, community providers, assisted living facilities, retirement homes, and aging service organizations between October 2016 and March 2018. Inclusion criteria were: 1) being 65 years or older; and 2) the ability to walk a minimum distance of 9.14 m (30 feet) with or without an assistive device (for the frailty assessment). Exclusion criteria were: 1) severe motor disorders (Parkinson's disease, multiple sclerosis, or recent stroke); 2) severe upper-extremity disorders (e.g., elbow bilateral fractures or rheumatoid arthritis); 3) cognitive impairment identified by a Mini-Mental State Examination (MMSE) score $\leq 23$ [31]; 4) terminal illness (i.e., progressive disease where death within six months is expected as a consequence); 5) diseases/disorders that can directly influence HR (including arrhythmia and use of pacemaker) and 6) usage of $\beta$-blockers or similar medications that can influence HR. Written informed consent was obtained according to the principles expressed in the Declaration of Helsinki [32]. The study was approved by the University of Arizona Institutional Review Board.

**Frailty assessment**

Frailty was assessed using the five-component Fried phenotype as an extensively validated and reliable tool [2]. This frailty test included: 1) self-reported unintentional weight loss of 4.54 kg (10 pounds) or more in the previous year; 2) weakness based on grip strength test; 3) slow walking speed; 4) self-reported exhaustion; and 5) self-reported low energy expenditure. Participants were categorized as non-frail if they met none of the criteria, pre-frail if they met one or two criteria, and frail if they met three or more criteria.
UEF test

Details of UEF validation and index development have been explained comprehensively within our previous work [28–30], and only crucial aspects of UEF regarding the measurement procedure and frailty category assessment were presented here. For UEF, while sitting on a chair, participants performed one trial of full elbow flexion and extension as fast as possible for 20 seconds using the right arm. Of note, we have shown UEF results are similar on both sides [28]. Before the test, participants performed a short practice trial with their non-dominant arm to become familiar with the protocol. The protocol was explained to participants, and they were encouraged only once, before elbow flexion, to do the task as fast as possible. To assure consistency, exact same verbal instruction was used, and participants were not further encouraged during the task. Wearable motion sensors (triaxial gyroscope sensors, BioSensics LLC, Cambridge, MA, sampling frequency = 100Hz) were used to measure forearm and upper-arm motion, and ultimately the elbow angular velocity.

Motor function outcomes representing physical frailty features were derived, including slowness (speed of elbow flexion), flexibility (range of motion), weakness (strength of upper-extremity muscles), speed variability (motor accuracy), speed reduction (fatigue), and flexion number. Based on these parameters a motor score (range: resilient=0; extremely frail=1) was calculated for each participant, based on the previously established model [30].

HR assessment

HR was measured using a wearable system with synchronized electrocardiogram (ECG) and accelerometer sensors (360° eMotion Faros, Mega Electronics, Kuopio, Finland; ECG sampling frequency = 1000Hz and accelerometer sampling frequency = 100Hz). One channel ECG was recorded using two electrodes. Electrodes were placed on the left chest, one on the upper mid-thorax, and the other one inferior to the left rib cage. Using the synchronized accelerometer data, the exact starting and endpoints of the UEF task were selected manually. Then a period of 5 seconds before and 10 seconds after the activity were selected, respectively, as baseline and recovery periods. To extract RR intervals, QRS peak detection was performed using the Pan-Tompkins algorithm [33], and detected peaks were manually inspected by two researchers (NT and ME).

Two types of HR measures were extracted, representing: 1) resting-state HR and HRV during baseline; and 2) HR dynamics including HR increase during UEF and HR recovery after UEF. HR baseline parameters included: 1) HR mean; 2) beat-to-beat (RR) interval mean; 3) RR CV: the coefficient of variation (standard deviation divided by mean) of RR intervals; and 4) RMSSD: root mean square of successive heartbeat interval differences. HR dynamics parameters explain the amount and timing of HR changes in response to UEF, which include: 1) time to peak HR: elapsed time to reach maximum HR during the task with reference to minimum baseline HR; 2) HR recovery time: elapsed time to reach minimum HR during the recovery with reference to maximum HR; 3) HR percent increase: percentage increase in HR during the task compared to minimum baseline HR; and 4) HR percent decrease: percent decrease in HR during the recovery compared to maximum HR during the task.
Statistical analysis and power analysis

Analysis of variance (ANOVA) models were used to evaluate the differences in demographic parameters between frailty groups, except sex; chi-square ($\chi^2$) test was used to assess differences in sex categories among frailty groups. HR parameters were compared between groups using ANOVA models; age, sex, and body mass index (BMI) were considered as covariates, and Cohen's effect size ($d$) was estimated. Age, sex, and BMI were selected as adjusting variables, since they have been previously associated with HR measures and frailty [30, 34–36]. In the next step of the analysis, HR and motor parameters, separately and combined, were used in multiple logistic regression models as independent variables to identify frailty status. A stepwise parameter selection based on Akaike information criterion (AIC) values was implemented to identify predictive independent variables. For each predicting model, the area under the curve (AUC) with 95% CI was calculated using receiver operator characteristics (ROC) curves. Power calculation was performed to detect differences in HR dynamic parameters between frailty groups for the sample size obtained for the current study using G*Power, ANOVA, Fixed-effect, one-way analysis. The number of participants in the current study (12 participants in each frailty group) was sufficient to provide 85% power to detect differences in HR dynamics (and UEF score) parameters with effect size of 0.65 and larger.

Results

Participants

Fifty-six eligible participants were recruited, including 12 non-frail (age = 76.92±7.32 years), 40 pre-frail (age = 80.53±8.12 years), and four frail (age = 88.25±4.42 years). For statistical analyses, the pre-frail and frail groups were merged due to the small number of included frail older adults in this study. None of the demographic information was significantly different between the two frailty groups ($p>0.10$, Table 1).

| Demographic information          | Non-frail (n=12) | Pre-frail/Frail (n=44) | $p$-value (effect size) |
|----------------------------------|------------------|------------------------|------------------------|
| Male, n (% of the group)         | 5 (42%)          | 10 (23%)               | 0.20                   |
| Age, year (SD)                   | 76.92 (7.32)     | 81.23 (8.14)           | 0.10 (0.55)            |
| Height, cm (SD)                  | 164.36 (9.13)    | 164.23 (10.27)         | 0.97 (0.01)            |
| Weight, kg (SD)                  | 66.58 (14.69)    | 75.56 (19.60)          | 0.15 (0.52)            |
| Body mass index, kg/m$^2$ (SD)   | 24.67 (5.55)     | 27.74 (5.71)           | 0.10 (0.55)            |

SD: standard deviation

UEF motor and HR Parameters
UEF motor score was significantly different between the two frailty categories; UEF motor score was 0.32±0.18 on average for non-frail and 0.53±0.23 for pre-frail/frail participants ($p<0.01$, Table 2). For HR dynamic parameters, pre-frail/frail older adults showed almost half of the amount of HR increase during UEF, and HR decrease during the recovery compared to non-frail participants ($p<0.01$, Table 2 and Fig. 1). Time to peak and recovery of HR, however, were not significantly different between the frailty groups ($p>0.49$, Table 2). Although trends of higher HR and smaller HRV for pre-frail/frail were observable, none of the baseline HR parameters were significantly different between the frailty groups ($p>0.23$, Table 2).

Table 2

| Parameters                        | Non-frail (n=12) | Pre-frail/Frail (n=44) | p-value (effect size) |
|-----------------------------------|------------------|------------------------|-----------------------|
| UEF motor score, 0-1 (SD)         | 0.32 (0.18)      | 0.53 (0.23)            | <0.01 (1.02) *        |
| HR baseline                       |                  |                        |                       |
| HR mean, BPM (SD)                 | 71.52 (11.38)    | 77.59 (15.97)          | 0.23 (0.44)           |
| RR mean, second (SD)              | 0.86 (0.13)      | 0.80 (0.14)            | 0.23 (0.44)           |
| RR CV, % (SD)                     | 1.70 (1.39)      | 1.58 (1.49)            | 0.77 (0.08)           |
| RMSSD, millisecond                | 16.80 (18.14)    | 15.80 (14.73)          | 0.81 (0.06)           |
| HR dynamics                       |                  |                        |                       |
| Time to peak HR, second (SD)      | 16.84 (6.46)     | 16.11 (5.82)           | 0.49 (0.12)           |
| HR recovery time, second (SD)     | 13.71 (6.22)     | 14.04 (5.76)           | 0.54 (0.06)           |
| HR percent increase, % (SD)       | 19.28 (7.55)     | 10.29 (4.79)           | <0.001 (1.42) *       |
| HR percent decrease, % (SD)       | 15.24 (7.65)     | 8.13 (3.97)            | <0.01 (1.17) *        |

Results from logistic models showed that percent increase and decrease in HR, as well as UEF motor score were all significantly associated with frailty ($p<0.01$). Using previously developed UEF motor score in the logistic model, an area under curve (AUC) of receiver operating characteristic (ROC) of 0.78 was achieved. Combining both UEF HR dynamic (i.e., HR percent increase) and motor score, the AUC was improved to 0.87 (Table 3). Using this model, pre-frailty/frailty was predicted with a sensitivity and specificity of 0.82 and 0.83 (Table 3).
Table 3
Results for logistic models using HR dynamics and UEF motor score. A significant association is represented by the asterisk.

| Independent variable | Parameter estimate | Standard error | Chi-square ($\chi^2$) | p-value (95% CI) |
|-----------------------|--------------------|----------------|-----------------------|------------------|
| Model 1 - UEF motor score (AUC=0.78; Sensitivity=0.75; Specificity=0.75) | | | | |
| Intercept             | 0.61               | 0.73           | 0.70                  | 0.4 (-0.81:2.12) |
| UEF motor score       | -0.05              | 0.02           | 6.85                  | <0.01 (-0.08:-0.01) * |
| Model 2 - HR dynamics (AUC=0.84; Sensitivity=0.80; Specificity=0.75) | | | | |
| Intercept             | -4.91              | 1.27           | 14.97                 | <0.001 (-7.92:-2.81) * |
| HR percent increase   | 0.25               | 0.08           | 10.38                 | <0.001 (0.12:0.44) * |
| Model 3 - Combined UEF (AUC=0.87; Sensitivity=0.82; Specificity=0.83) | | | | |
| Intercept             | -3.21              | 1.55           | 4.28                  | 0.04 (-6.68:-0.45) * |
| HR percent increase   | 0.23               | 0.08           | 7.73                  | <0.01 (0.09:0.42) * |
| UEF motor score       | -0.03              | 0.02           | 2.67                  | 0.1 (-0.07:0.01) |

HR: heart rate; UEF: upper-extremity function; AUC: area under curve; CI: confidence interval

Discussion

HR dynamics and frailty

As hypothesized, significant associations were observed between frailty status and HR changes during the activity and afterwards during the recovery period. During physical activity, an increase in sympathetic outflow increases HR and stroke volume to match demand [37, 38]. During recovery from the physical activity, parasympathetic activity increases to reduce HR to baseline [39–41]. Lack of resilience in changing HR in pre-frail/frail older adults can be explained by both a compromised ANS performance or lack of cardiac reserve. Previous research provided evidence of ANS dysfunction with frailty. Focusing on resting state differences in HRV as an indicator of ANS performance, a smaller HRV has been observed among pre-frail and frail older adults compared to non-frails [42]. On the other hand, lack of cardiac reserve during resting, can move pre-frail/frail individuals to a more imbalance (less homeostatic) and already stressed state, causing an inability to respond to additional stress such as a simple task of arm movement. In confirmation of this theory, although not significant, we observed trends of higher mean HR during resting among pre-frail/frail participants compared to non-frails (Table 3).

Only a few studies exist to assess HR dynamics during activity across frailty groups. Smaller changes in HR has been reported previously for lying-to-standing and seated step test [43, 44]. Also, in our previous
research we observed that pre-frail/frail older adults had 46% smaller and 49% slower increase in HR during walking compared to pre-frail/frail older adults [45]. One noticeable difference between our previous and current findings is that time to peak HR during activity was significantly different in 15 feet walk test, while this parameter was not different in the current study. One possible explanation is that for pre-frail and frail older adults performing a walking test with a set distance takes longer than non-frails, which consequently may lead to a slower HR increase. This explanation needs to be further assessed by executing walking test with a set duration rather than distance. Nevertheless, based on current findings, assessing changes in HR magnitude, rather than timing of HR changes (both increase and recovery) may provide a more robust way of measuring HR dynamics.

Another important observation was that HR increase can characterize cardiac imbalance behavior, similar or even better than HR recovery. Most previous research has focused on HR recovery for disease diagnosis; studies showed prognostic value in measuring HR recovery one minute after cardiopulmonary exercise testing for heart failure prediction [17, 46]. Nevertheless, all these HR assessments were performed after the physical activity, since performing whole body exercise makes accurate HR assessment complex due to motion artifacts. In the implemented UEF approach, participants performed elbow flexion with the right arm while HR data measurement electrodes were placed on the left side. This ECG placement provided minimal motion artifact from the right-side arm movement to permit accurate dynamic HR assessment.

**Combined HR and motor model**

In confirmation of our hypothesis, current results suggest that combining HR and motor function in a single model can enhance frailty prediction in comparison to models involving each of these physiological systems individually. It is believed now that frailty is caused by loss of homeostasis not necessary in one domain, but multiple physiological systems; in other words, frailty is the result of a compromised dynamic interaction between several physiological systems, rather than one specific pathway [47]. Accordingly, the concept of frailty assessment across multiple physiological systems and their interactions has recently drawn more attention. Ghachem et al., for instance, assessed dysregulation of six physiological systems including oxygen transport, kidney/liver function, leukopoiesis, micronutrients, lipids, and electrolytes in association with frailty [48]. They have provided evidence that frailty is more strongly associated with the number of dysregulated systems, rather than the type of dysregulation [48]. In confirmation to previous research, current findings support the hypothesis that assessing multiple physiological systems would improve frailty assessment. Unlike previous work, our approach involved one testing, within which, both cardiac and motor performance were evaluated, to efficiently balance the accuracy and the burden of testing.

**Limitation and future direction**

Although current findings were promising, there are limitations that warrant future research. First, the sample of community dwelling older adults chosen for the current study was small and may not reflect condition of hospitalized older adults. Due to the small sample of participants additional analyses were
not performed, including assessment of interaction effect of frailty and HR on motor function performance. Further, in the current study the association between baseline HR and HR dynamic parameters were not reported. Since these results were similar to our previous work, we encourage readers to read previously reported findings regarding HR measure analysis during gait tests [45]. One other limitation of the current study is the lack of long-term resting HR measurement. Although five seconds of rest before UEF would be enough for short-term HRV assessment, several other analyses related to regularity (complexity analysis) of HR data could not be accomplished here due to limited number of samples for nonlinear dynamic analysis. Previous studies demonstrated significant association between HR complexity and frailty [49–51], and therefore, it would be interesting to explore how the HR complexity during basal condition is related to HR dynamics in response to physical activity, especially across frailty groups. Lastly, older adult participants with arrhythmia and those who used β-blockers and pacemakers were excluded from the study. Therefore, effects of these disorders and medications on HR measures need to be studies in future.

**Conclusion And Clinical Implications**

Current findings showed that HR changes due to physical activity was smaller among pre-frail/frail individuals during the activity and afterwards during the recovery period. We also showed that by combining HR and motor function we may improve frailty prediction compared to models incorporating each of these measures individually. The proposed multimodal HR and motor frailty assessment approach is objective and easy to perform. Due to its simplicity, compared to gait test, this test can be performed on hospitalized patients to predict therapy complications and identify patients with treatment-responsive frailty for directing appropriate care, with potential implications for older adults with heart diseases.

**Declarations**

**Ethics approval and consent to participate:** Written informed consent was obtained according to the principles expressed in the Declaration of Helsinki. The study was approved by the University of Arizona Institutional Review Board (Protocol ID: 1412591150R002).

**Consent for publication:** Not applicable

**Availability of data and materials:** The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request. The implemented code for HR data analysis will be available for research purposes. Please contact the corresponding author for request access to UEF codes.

**Competing interests:** The authors declare that they have no competing interests

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References

1. Clegg A, Young J, Iliffe S, Rikkert MO, Rockwood K: Frailty in elderly people. The lancet 2013, 381(9868):752-762.
2. Fried LP, Tangen CM, Walston J, Newman AB, Hirsch C, Gottdiener J, Seeman T, Tracy R, Kop WJ, Burke G: Frailty in older adults: evidence for a phenotype. The Journals of Gerontology Series A: Biological Sciences and Medical Sciences 2001, 56(3):M146-M157.
3. Yao X, Li H, Leng SX: Inflammation and immune system alterations in frailty. Clinics in geriatric medicine 2011, 27(1):79-87.
4. Hubbard RE, O’Mahony MS, Savva GM, Calver BL, Woodhouse KW: Inflammation and frailty measures in older people. Journal of cellular and molecular medicine 2009, 13(9b):3103-3109.
5. Serviddio G, Romano AD, Greco A, Rollo T, Bellanti F, Altomare E, Vendemiale G: Frailty syndrome is associated with altered circulating redox balance and increased markers of oxidative stress. International Journal of Immunopathology and Pharmacology 2009, 22(3):819-827.
6. Schaap LA, Pluijm SM, Deeg DJ, Visser M: Inflammatory markers and loss of muscle mass (sarcopenia) and strength. The American journal of medicine 2006, 119(6):526. e529-526. e517.
7. Visser M, Pahor M, Taaffe DR, Goodpaster BH, Simonsick EM, Newman AB, Nevitt M, Harris TB: Relationship of interleukin-6 and tumor necrosis factor-α with muscle mass and muscle strength in elderly men and women: the Health ABC Study. The Journals of Gerontology Series A: Biological Sciences and Medical Sciences 2002, 57(5):M326-M332.
8. Cesari M, Penninx BW, Pahor M, Lauretani F, Corsi AM, Williams GR, Guralnik JM, Ferrucci L: Inflammatory markers and physical performance in older persons: the InCHIANTI study. The Journals of Gerontology Series A: Biological Sciences and Medical Sciences 2004, 59(3):M242-M248.
9. Johar H, Emeny RT, Bidlingmaier M, Reincke M, Thorand B, Peters A, Heier M, Ladwig K-H: Blunted diurnal cortisol pattern is associated with frailty: a cross-sectional study of 745 participants aged 65 to 90 years. The Journal of Clinical Endocrinology & Metabolism 2014, 99(3):E464-E468.

10. O'Connell M, Tajar A, Roberts SA, Wu F: Do androgens play any role in the physical frailty of ageing men? International journal of andrology 2011, 34(3):195-211.

11. López-Armada MJ, Riveiro-Naveira RR, Vaamonde-García C, Valcárcel-Ares MN: Mitochondrial dysfunction and the inflammatory response. Mitochondrion 2013, 13(2):106-118.

12. Varadhan R, Chaves PH, Lipsitz LA, Stein PK, Tian J, Windham BG, Berger RD, Fried LP: Frailty and impaired cardiac autonomic control: new insights from principal components aggregation of traditional heart rate variability indices. Journals of Gerontology Series A: Biomedical Sciences and Medical Sciences 2009, 64(6):682-687.

13. Moghtadaei M, Jansen HJ, Mackasey M, Rafferty SA, Bogachev O, Sapp JL, Howlett SE, Rose RA: The impacts of age and frailty on heart rate and sinoatrial node function. The Journal of physiology 2016, 594(23):7105-7126.

14. Adamson PB, Smith AL, Abraham WT, Kleckner KJ, Stadler RW, Shih A, Rhodes MM: Continuous autonomic assessment in patients with symptomatic heart failure: prognostic value of heart rate variability measured by an implanted cardiac resynchronization device. Circulation 2004, 110(16):2389-2394.

15. Scalvini S, Volterrani M, Zanelli E, Pagani M, Mazzuero G, Coats AJ, Giordano A: Is heart rate variability a reliable method to assess autonomic modulation in left ventricular dysfunction and heart failure?: Assessment of autonomic modulation with heart rate variability. International journal of cardiology 1998, 67(1):9-17.

16. De Jong MMJ, Randall DC: Heart rate variability analysis in the assessment of autonomic function in heart failure. Journal of Cardiovascular Nursing 2005, 20(3):186-195.

17. Arena R, Guazzi M, Myers J, Peberdy MA: Prognostic value of heart rate recovery in patients with heart failure. American heart journal 2006, 151(4):851. e857-851. e813.

18. Morris JA, Norris PR: Role of reduced heart rate volatility in predicting death in trauma patients. Advances in surgery 2005, 39:77-96.

19. Muller JE, Tofler G, Stone P: Circadian variation and triggers of onset of acute cardiovascular disease. Circulation 1989, 79(4):733-743.

20. Kleiger RE, Miller JP, Bigger Jr JT, Moss AJ: Decreased heart rate variability and its association with increased mortality after acute myocardial infarction. The American journal of cardiology 1987, 59(4):256-262.

21. Javorka M, Zila I, Balharek T, Javorka K: Heart rate recovery after exercise: relations to heart rate variability and complexity. Brazilian Journal of Medical and Biological Research 2002, 35(8):991-1000.

22. Cipryan L, Litschmannova M: Intra-day and inter-day reliability of heart rate variability measurement. Journal of Sports Sciences 2013, 31(2):150-158.
23. Pinna GD, Maestri R, Torunski A, Danilowicz-Szymanowicz L, Swoch M, La Rovere MT, Raczak G: Heart rate variability measures: a fresh look at reliability. *Clinical Science* 2007, **113**(3):131-140.

24. Taelman J, Vandeput S, Spaepen A, Van Huffel S: Influence of mental stress on heart rate and heart rate variability. In: *4th European conference of the international federation for medical and biological engineering: 2009*. Springer: 1366-1369.

25. Toosizadeh N, Ehsani H, Parthasarathy S, Carpenter B, Ruberto K, Mohler J, Parvaneh S: Frailty and Heart Response to Physical Activity. *Archives of Gerontology and Geriatrics* 2020:104323.

26. Ehsani H, Mohler MJ, Golden T, Toosizadeh N: Upper-extremity function prospectively predicts adverse discharge and all-cause COPD readmissions: a pilot study. *International Journal of Chronic Obstructive Pulmonary Disease* 2019, **14**:39.

27. Toosizadeh N, Berry C, Bime C, Najafi B, Kraft M, Mohler J: Assessing upper-extremity motion: An innovative method to quantify functional capacity in patients with chronic obstructive pulmonary disease. *PloS one* 2017, **12**(2):e0172766.

28. Toosizadeh N, Mohler J, Najafi B: Assessing upper extremity motion: an innovative method to identify frailty. *Journal of the American Geriatrics Society* 2015, **63**(6):1181-1186.

29. Toosizadeh N, Joseph B, Heusser MR, Jokar TO, Mohler J, Phelan HA, Najafi B: Assessing upper-extremity motion: an innovative, objective method to identify frailty in older bed-bound trauma patients. *Journal of the American College of Surgeons* 2016, **223**(2):240-248.

30. Toosizadeh N, Wendel C, Hsu C-H, Zamrini E, Mohler J: Frailty assessment in older adults using upper-extremity function: index development. *BMC geriatrics* 2017, **17**(1):1-7.

31. Folstein MF, Folstein SE, McHugh PR: "Mini-mental state": a practical method for grading the cognitive state of patients for the clinician. *Journal of psychiatric research* 1975, **12**(3):189-198.

32. Association GAotWM: World Medical Association Declaration of Helsinki: ethical principles for medical research involving human subjects. *The Journal of the American College of Dentists* 2014, **81**(3):14-18.

33. Pan J, Tompkins WJ: A real-time QRS detection algorithm. *IEEE transactions on biomedical engineering* 1985(3):230-236.

34. Anderson R, Jönsson P, Sandsten M: Effects of Age, BMI, Anxiety and Stress on the Parameters of a Stochastic Model for Heart Rate Variability Including Respiratory Information. *BIOSIGNALS* 2018, **4**:17-25.

35. Gordon E, Peel N, Samanta M, Theou O, Howlett S, Hubbard R: Sex differences in frailty: a systematic review and meta-analysis. *Experimental gerontology* 2017, **89**:30-40.

36. Hubbard RE, Lang IA, Llewellyn DJ, Rockwood K: Frailty, body mass index, and abdominal obesity in older people. *Journals of Gerontology Series A: Biomedical Sciences and Medical Sciences* 2010, **65**(4):377-381.

37. Evrengul H, Tanriverdi H, Kose S, Amasyali B, Kilic A, Celik T, Turhan H: The relationship between heart rate recovery and heart rate variability in coronary artery disease. *Annals of Noninvasive Electrocardiology* 2006, **11**(2):154-162.
38. Shephard RJ: Exercise physiology. Toronto; Philadelphia: BC Decker; Saint Louis, Mo.: Sales and distribution ...; 1987.

39. Carter III R, Watenpaugh DE, Wasmund WL, Wasmund SL, Smith ML: Muscle pump and central command during recovery from exercise in humans. Journal of Applied Physiology 1999, 87(4):1463-1469.

40. Nishime EO, Cole CR, Blackstone EH, Pashkow FJ, Lauer MS: Heart rate recovery and treadmill exercise score as predictors of mortality in patients referred for exercise ECG. Jama 2000, 284(11):1392-1398.

41. Savin WM, Davidson DM, Haskell WL: Autonomic contribution to heart rate recovery from exercise in humans. Journal of Applied Physiology 1982, 53(6):1572-1575.

42. Parvaneh S, Howe CL, Toosizadeh N, Honarvar B, Slepian MJ, Fain M, Mohler J, Najafi B: Regulation of cardiac autonomic nervous system control across frailty statuses: a systematic review. Gerontology 2016, 62(1):3-15.

43. Romero-Ortuno R, Cogan L, O’Shea D, Lawlor BA, Kenny RA: Orthostatic haemodynamics may be impaired in frailty. Age and ageing 2011, 40(5):576-583.

44. Weiss CO, Hoenig HH, Varadhan R, Simonsick EM, Fried LP: Relationships of cardiac, pulmonary, and muscle reserves and frailty to exercise capacity in older women. Journals of Gerontology Series A: Biomedical Sciences and Medical Sciences 2010, 65(3):287-294.

45. Toosizadeh N, Ehsani H, Parthasarathy S, Carpenter B, Ruberto K, Mohler J, Parvaneh S: Frailty and heart response to physical activity. Archives of gerontology and geriatrics 2021, 93:104323.

46. Maeder MT, Ammann P, Rickli H, Brunner-La Rocca HP: Impact of the exercise mode on heart rate recovery after maximal exercise. European journal of applied physiology 2009, 105(2):247-255.

47. Fried LP, Xue Q-L, Cappola AR, Ferrucci L, Chaves P, Varadhan R, Guralnik JM, Leng SX, Semba RD, Walston JD: Nonlinear multisystem physiological dysregulation associated with frailty in older women: implications for etiology and treatment. Journals of Gerontology Series A: Biomedical Sciences and Medical Sciences 2009, 64(10):1049-1057.

48. Ghachem A, Fried LP, Legault V, Bandeen-Roche K, Presse N, Gaudreau P, Cohen AA: Evidence from two cohorts for the frailty syndrome as an emergent state of parallel dysregulation in multiple physiological systems. Biogerontology 2021, 22(1):63-79.

49. Chaves PH, Varadhan R, Lipsitz LA, Stein PK, Windham BG, Tian J, Fleisher LA, Guralnik JM, Fried LP: Physiological complexity underlying heart rate dynamics and frailty status in community-dwelling older women. Journal of the American Geriatrics Society 2008, 56(9):1698-1703.

50. Katayama PL, Dias DPM, Silva LEV, Virtuoso-Junior JS, Marocolo M: Cardiac autonomic modulation in non-frail, pre-frail and frail elderly women: a pilot study. Aging clinical and experimental research 2015, 27(5):621-629.

51. Takahashi AC, Bonjorni LA, Buto MS, Vassimon-Barroso V, Minatel V, Rocha SM, Ribeiro FH, Montano N, Porta A, Catai AM: Short-term complexity of cardiovascular oscillations in frailty syndrome. In:
**Figures**

![Bar charts showing HR dynamic parameters between non-frail (NF) and pre-frail/frail (PF/F) participants](chart.png)

**Figure 1**

Differences in HR dynamic parameters between non-frail (NF) and pre-frail/frail (PF/F) participants. *p*-values for ANOVA model, adjusted with age, sex, and body mass index are presented.

- **HR Increase (%)**
  - NF: 20 ± 2
  - PF/F: 10 ± 2
  - *p* < 0.001
  - Effect size = 1.42

- **HR Decrease (%)**
  - NF: 16 ± 2
  - PF/F: 8 ± 2
  - *p* < 0.01
  - Effect size = 1.17