Research article

A standard procedure for creating a frailty index
Samuel D Searle¹, Arnold Mitnitski¹,²,³, Evelyne A Gahbauer⁴, Thomas M Gill⁴ and Kenneth Rockwood*¹,²,⁵

Address: ¹Geriatric Medicine Research Unit, Dalhousie University & Capital District Health Authority, Halifax, Canada, ²Department of Medicine, Dalhousie University, Halifax, Canada, ³Department of Mathematics & Statistics, Dalhousie University, Halifax, Canada, ⁴Department of Internal Medicine, Yale University School of Medicine, New Haven, CT 06504, USA and ⁵Division of Geriatric Medicine, Dalhousie University, Halifax, Canada

Email: Samuel D Searle - ssearle@dal.ca; Arnold Mitnitski - arnold.mitnitski@dal.ca; Evelyne A Gahbauer - evelyne.gahbauer@yale.edu; Thomas M Gill - thomas.gill@yale.edu; Kenneth Rockwood* - kenneth.rockwood@dal.ca

* Corresponding author

Abstract

Background: Frailty can be measured in relation to the accumulation of deficits using a frailty index. A frailty index can be developed from most ageing databases. Our objective is to systematically describe a standard procedure for constructing a frailty index.

Methods: This is a secondary analysis of the Yale Precipitating Events Project cohort study, based in New Haven CT. Non-disabled people aged 70 years or older (n = 754) were enrolled and re-contacted every 18 months. The database includes variables on function, cognition, co-morbidity, health attitudes and practices and physical performance measures. Data came from the baseline cohort and those available at the first 18-month follow-up assessment.

Results: Procedures for selecting health variables as candidate deficits were applied to yield 40 deficits. Recoding procedures were applied for categorical, ordinal and interval variables such that they could be mapped to the interval 0–1, where 0 = absence of a deficit, and 1 = full expression of the deficit. These individual deficit scores were combined in an index, where 0 = no deficit present, and 1 = all 40 deficits present. The values of the index were well fit by a gamma distribution. Between the baseline and follow-up cohorts, the age-related slope of deficit accumulation increased from 0.020 (95% confidence interval, 0.014–0.026) to 0.026 (0.020–0.032). The 99% limit to deficit accumulation was 0.6 in the baseline cohort and 0.7 in the follow-up cohort. Multivariate Cox analysis showed the frailty index, age and sex to be significant predictors of mortality.

Conclusion: A systematic process for creating a frailty index, which relates deficit accumulation to the individual risk of death, showed reproducible properties in the Yale Precipitating Events Project cohort study. This method of quantifying frailty can aid our understanding of frailty-related health characteristics in older adults.

Background

Frailty is a state of increased vulnerability to adverse outcomes. How best to operationalize frailty is controversial [1-3], but one method uses a frailty index [4]. The principle is to count deficits in health (which can be symptoms, signs, diseases, disabilities or laboratory, radiographic or...
electrocardiographic abnormalities) on the grounds that the more deficits a person has, the more likely that person is to be frail. The index is often expressed as a ratio of deficits present to the total number of deficits considered. For example, if 40 deficits were considered, and 10 were present in a given person, that person's frailty index would be 10/40 = 0.25.

Although the idea and approach are relatively simple, the results yielded by the frailty index have been consistent between surveys evaluated by our group [4-7] and by others [8-11] even though not every frailty index considers the same deficits, or even the same number of deficits. For example, across several frailty index measures, people accumulate deficits, on average, at about 0.03/year [4,5]. In each study, the frailer the person is (the higher the deficit count) the more vulnerable they are to adverse outcomes [5,8,11]. The frailty index is strongly associated with the risk of death, institutionalization and worsening health status, especially when at least 30 variables are included [5]. The frailty index also shows a consistent, sub-maximal limit at about 2/3 of the deficits that are considered. For example, if a frailty index is composed of 60 items, the most that anyone will have wrong with them is not 60, but 40 [5].

The reproducibility of the findings in relation to the frailty index is of some interest because none of the samples in which the frailty index has been operationalized has considered the same deficits. To be clear, it does not matter if study A considered 40 deficits from set X of deficits and study B considered 60 deficits from set Y of deficits; the estimates from each (e.g. the rate of deficit accumulation, the relationship between deficit accumulation and mortality, or the limit to deficit accumulation) appear to be similar. This finding suggests that frailty is a real phenomenon, which is a property of a biologically complex system. It indicates that frailty can be measured in many ways, and therefore can be studied in many existing datasets that might not have set out to measure frailty per se. To encourage more widespread evaluation of frailty – a goal encouraged by many groups [12-14] – we present a detailed, step-by-step procedure to describe which potential variables can be included in a frailty index, and how to establish cut-points for continuous variables. Here, frailty indexes were newly created using baseline and follow-up samples from an existing cohort study, and their properties (e.g. rate of increase, limit, and relationship with mortality) were compared with each other, and with earlier work.

Methods
The Study Sample
The Yale Precipitating Events Project (PEP) is a cohort study based in New Haven CT that enrolled individuals aged 70 years or older. Its methods have been published elsewhere [15,16]. Briefly, 754 community dwelling, English speaking, non-disabled persons with life expectancy and plans to stay in the area for more than 12 months were enrolled in the study. Comprehensive home-based assessments were completed at baseline and every 18 months. The 18-month assessment included 681 participants aged 72 to 98 years. This report uses the baseline and 18 month follow up data to contrast with each other and to compare the properties of the new indexes created in this data set with previously reported frailty indexes. At baseline, most participants (n = 487, 64.6%) were women, and most (n = 682, 90.5%) were white, with a mean Mini-Mental State Examination (MMSE) [17] score of 26.8 (SD = 2.50). Mortality was checked monthly for nine years from the baseline interview and was confirmed by obituaries and death certificates.

Selecting candidate deficits for the Frailty Index
A frailty index counts deficits in health. These deficits were defined as symptoms, signs, disabilities and diseases [5]. All health deficits, including continuous, ordinal and binary variables, were taken from the PEP survey data dictionary. Restricted activity, disability in Activities Daily Living (ADL) and Instrumental ADL, impairments in general cognition and physical performance (e.g. impaired grip strength, impaired walking), co-morbidity, self-rated health, and depression/mood were evaluated.

Variables can be included in a frailty index if they satisfy the following 5 criteria:

1) The variables must be deficits associated with health status. Attributes such as graying hair, while age-related, are attributes and therefore not included. 2) A deficit’s prevalence must generally increase with age, although some clearly age-related adverse conditions can decrease in prevalence at very advanced ages due to survivor effects. 3) Similarly, the chosen deficits must not saturate too early. For instance, age-related lens changes resulting in problems with accommodation (presbyopia) are nearly universal by age 55; in other words, as a variable, presbyopia saturates too early to be considered as a deficit here. 4) When considering the candidate deficits as a group, the deficits that make up a frailty index must cover a range of systems – if all variables were related to cognition, for example, the resulting index might well describe changes in cognition over time, but would be a cognitive impairment index [18] not a frailty index. 5) If a single frailty index is to be used serially on the same people, the items that make up the frailty index need to be the same from one iteration to the next [19]. The requirement to use the same items need not apply to comparisons between samples – i.e. samples that use difference frailty indexes appear to yield similar results [5].
Deficits should be added until there are at least 30–40 total deficits. There needs to be a minimum number of deficits. In general, the more variables that are included in a frailty index, the more precise estimates become. Similarly, estimates are unstable when the number of deficits is small – about 10 or less. Even so, an index with 30–40 variables has been shown to be sufficiently accurate for predicting adverse outcomes [6,14]. Furthermore, a frailty index can be constructed using information that is readily available in most health surveys, and is clinically tractable – i.e. it uses an amount that would be gathered in many routine health assessments of older adults [5].

Coding of individual variables
All binary variables were recoded, using the convention that '0' indicated the absence of the deficit, and '1' the presence of a deficit. For variables that included a single intermediate response (e.g. 'sometimes' or 'maybe'), we used an additional value of '0.5'.

Frailty index variables can also accommodate ordinal and continuous variables as deficits. To do so requires grading the continuum or rank into a score between 0 (where no deficit is present) and 1 (where the deficit is maximally expressed by the given variable). For some variables, this re-coding is self-evident. Consider the widely used Self-rated Health Question (“How would you rate your health? Excellent, Very Good, Good, Fair, Poor”). To grade this between '0' and '1', each lower self-rating of health was coded to represent a larger deficit (“Excellent = 0”, “Very Good = 0.25”, “Good = 0.5”, “Fair = 0.75” and “Poor = 1”). Similarly, recognized cut-points can be used for ordinal and continuous variables, such as the rapid walk test [15]. For the MMSE, we recoded deficits according to severity of impairment [20]. We assigned a 1 for scores less than 10, denoting severe dementia, 0.75 for scores ≥ 10 and ≤ 17, denoting moderate dementia, 0.5 for scores ≥ 18 and ≤ 20, denoting mild dementia, 0.25 for scores >20 and <24, denoting mild cognitive impairment (MCI), and 0 for scores ≥ 24, denoting no cognitive impairment [20]. Some readers might object that a score of '1' seems something of a discount (not a sufficiently high count) for severe dementia, and that losing only 1 point for it, compared with 0.25 points for MCI is not valid on its face. Consider, however, that a person with severe dementia is likely to have many more deficits than a person with MCI, e.g. more disability, poorer physical performance, higher degrees of behavioural problems and so forth.

Because not all ordinal or continuous variables have published or self-evident cut-points, additional work is required to establish the least arbitrary cut-points for these variables. Methods to address this can be broadly categorized as those based on characteristics of the distribution and those based on judgment (e.g. in relation to some clinically relevant hazard) [21]. Here, we employ both approaches. We used all existing previously coded deficits to establish an interim frailty index, whose purpose was to help provide cut-points for the remaining variables. This interim/nearly completed index was then plotted against the remaining ordinal and continuous variables to understand where their cut-points might be determined. The value of the individual variable that corresponded to 0.2 on the interim frailty index, i.e. the value of the variable at which, on average people had a frailty score of 0.2 or higher, was denoted as that deficit's cut-point. The value 0.2 on the frailty index is recognized by multiple frailty measures as approaching a frail state [7,8,22], so that this method met the convention of defining deficit cut-points. In addition, setting the value at, say, 0.3 seems unreasonably high, as this is consistently well into the range of frailty, however, defined (including by an increased hazard) so would be insensitive. Greater sensitivity is obtained at a cut-point of 0.1, but with less specificity.

Analysis of Baseline and Follow up Cohorts
The rate of accumulation of deficits was calculated by evaluating the slope of a best fit log of the frailty index in relation to age. To evaluate the impact of a given variable on the frailty index, we used an iterative, re-sampling process, similar to “bootstrapping” as detailed elsewhere and elsewhere [19,23]. We performed 1000 iterations where each iteration calculated the baseline and follow up frailty indexes using 80% of their variables, plotted the log of these two frailty indices versus age, and recorded the slope. By analyzing the range of the slopes, we were able to calculate 95% confidence intervals.

To observe the upper limit of the frailty index, the 99th percentiles of each cohort's frailty index was plotted against age. A flattening of this curve (i.e. its approach to a 0 slope) would suggest a common maximum to the frailty index at every age, consistent with earlier observations [24]. Statistical distributions of the frailty index were compared with theoretical models (Goodness of fit by least squares).

Survival analyses were done using bi-variate and multivariate Cox Regression analyses with the frailty index as the independent variable and age and gender as covariates on each of the two survey waves. The survival calculations were based on the available nine year mortality data from the baseline survey. The 18 month follow up survival calculations based from the available seven and a half year (from 18 month interview) mortality data.
Comparison of the Frailty Index calculated for the PEP Study with earlier estimates

The two calculated indexes, one from baseline and one at follow-up, were compared with previously published indexes, to see how well each fit the following characteristics: 1) The Frailty Index should have a skewed density distribution (histogram) that is well approximated by a gamma distribution [4,8,10] 2) The rate of deficit accumulation (prior estimate is 0.03 per year) [4-6]; 3) The presence of a sub-maximal, age-invariant limit to the Frailty Index (prior estimate is ~0.67) [5,6,8]; and 4) Association of the mean value of the Frailty Index with mortality [4-6,8,10,25].

Ethics

The study protocol was approved by the Yale Human Investigation Committee, New Haven; all participants provided informed consent at baseline and at follow-up. Ethical approval for secondary analyses was obtained from the Capital District Health Authority, Halifax, Nova Scotia.

Results

Construction and characteristics of the Frailty Index at baseline and at follow-up

Of the variables considered, 40 variables that met all frailty index criteria at both baseline and follow-up were chosen (Table 1). Variables were eliminated because they

Table 1: Health Variables and Cut-points for the Frailty Index

| List of 40 Variables included in the frailty index | Cut Point |
|--------------------------------------------------|-----------|
| Help Bathing                                     | Yes = 1, No = 0 |
| Help Dressing                                    | Yes = 1, No = 0 |
| Help getting in/out of Chair                     | Yes = 1, No = 0 |
| Help Walking around house                        | Yes = 1, No = 0 |
| Help Eating                                      | Yes = 1, No = 0 |
| Help Grooming                                    | Yes = 1, No = 0 |
| Help Using Toilet                                | Yes = 1, No = 0 |
| Help up/down Stairs                              | Yes = 1, No = 0 |
| Help lifting 10 lbs                              | Yes = 1, No = 0 |
| Help Shopping                                    | Yes = 1, No = 0 |
| Help with Housework                              | Yes = 1, No = 0 |
| Help with meal Preparations                      | Yes = 1, No = 0 |
| Help taking Medication                           | Yes = 1, No = 0 |
| Help with Finances                               | Yes = 1, No = 0 |
| Lost more than 10 lbs in last year               | Yes = 1, No = 0 |
| Self Rating of Health                            | Poor = 1, Fair = 0.75, Good = 0.5, V. Good = 0.25, Excellent = 0 |
| How Health has changed in last year              | Worse = 1, Better/Same = 0 |
| Stayed in Bed at least half the day due to health (in last month) | Yes = 1, No = 0 |
| Cut down on Usual Activity (in last month)       | Yes = 1, No = 0 |
| Walk outside                                     | <3 days = 1, , ≤ 3 days = 0 |
| Feel Everything is an Effort                     | Most of time = 1, Some time = 0.5, Rarely = 0 |
| Feel Depressed                                   | Most of time = 1, Some time = 0.5, Rarely = 0 |
| Feel Happy                                       | Most of time = 0, Some time = 0.5, Rarely = 1 |
| Feel Lonely                                      | Most of time = 1, Some time = 0.5, Rarely = 0 |
| Have Trouble getting going                       | Most of time = 1, Some time = 0.5, Rarely = 0 |
| High blood pressure                              | Yes = 1, Suspect = 0.5, No = 0 |
| Heart attack                                     | Yes = 1, Suspect = 0.5, No = 0 |
| CHF                                              | Yes = 1, Suspect = 0.5, No = 0 |
| Stroke                                           | Yes = 1, Suspect = 0.5, No = 0 |
| Cancer                                           | Yes = 1, Suspect = 0.5, No = 0 |
| Diabetes                                         | Yes = 1, Suspect = 0.5, No = 0 |
| Arthritis                                        | Yes = 1, Suspect = 0.5, No = 0 |
| Chronic Lung Disease                             | Yes = 1, Suspect = 0.5, No = 0 |
| MMSE                                             | <10 = 1, 11–17 = 0.75, 18–20 = 0.5, 20–24 = 0.25, >24 = 0 |
| Peak Flow                                        | See Table 2 |
| Shoulder Strength                                | See Table 2 |
| BMI                                              | See Table 2 |
| Grip Strength                                    | See Table 2 |
| Usual Pace                                       | See Table 2 |
| Rapid Pace                                       | See Table 2 |

The list of health deficit variables included in the FI and how they were coded as deficits.
Table 2: Continuous Variable Cut-points

| Variable               | Deficit for Men                  | Deficit for Women               | Source of cut point               |
|------------------------|----------------------------------|----------------------------------|-----------------------------------|
| Peak Flow (liters/min) | ≤340                             | ≤310                             | Plotted versus frailty index      |
| Body Mass Index (BMI)  | <18.5, ≥ 30 as a deficit.        | <18.5, ≥ 30 as a deficit.        | Published [34]                    |
|                        | 25–<30 as a ’half deficit’       | 25–<30 as a ’half deficit’       |                                   |
| Shoulder Strength (kg) | ≤12                              | ≤9                               | Plotted versus frailty index      |
| Grip Strength (GS in kg) | For BMI ≤ 24, GS ≤ 29          | For BMI ≤ 23, GS ≤ 17            | Published [15.22]                 |
|                        | For BMI 24.1–28, GS ≤ 30         | For BMI 23.1–26, GS ≤ 17.3       |                                   |
|                        | For BMI >28, GS ≤ 32             | For BMI 26.1–29, GS ≤ 18         |                                   |
|                        |                                  | For BMI>29, GS ≤ 21              |                                   |
| Rapid pace Walk (sec)  | >10                              | >10                              | Published [15]                    |
| Usual pace Walk (sec)  | >16                              | >16                              | Plotted versus frailty index      |

Deficit cut off values for continuous variables by sex and source of cut off.
frailty index (nature of the distribution; slope in relation to age; presence of a limit) seem to be broadly replicable. Several reproducible characteristics (e.g. the distribution, the slope and limit of deficit accumulation) of each Frailty Index were provided so that they maybe used, as in previous papers [4-11], to describe the overall frailty state of the group. The baseline Frailty Index showed a rate of accumulation to be 0.020 per year (per 1 year increase in age) with an upper limit to the frailty index of about 0.60 while the follow up showed a rate of 0.026 deficits accumulated per year with a limit around 0.70 (Figure 2).

We used a re-sampling by variable procedure to construct confidence intervals for the slopes of the lines (Figure 3). This procedure gives us information about the frailty construct, showing that a range of deficits can in fact be combined to give a result that is informative in the aggregate. In other words, the slope depends on the overall behaviour of the deficit accumulation, and is not driven by a small number of variables. In this regard, earlier work has shown reasonable consistency of the rate of deficit accumulation across community-dwelling random samples [6]. Here, we noted that the follow up cohort had frailty index characteristics – frailty index values, rate and limit similar to those of previously studied community dwelling random samples. Most notable is the 0.03 accumulation of deficits and the age independent limit to frailty of 0.67. The baseline sample had lower estimates – a lower average Frailty Index and a lower maximal limit. This suggests that the baseline cohort was not as frail as the follow up cohort.

The relationship between the Frailty Index and mortality is of interest on several grounds, but here is presented chiefly because it represents a relevant and non-arbitrary test of predictive validity. This is important because predictive validity is one of two types of so-called criterion validation, the other being validation against a so-called “gold standard” [21]. As there is no gold standard for frailty assessment, predictive validation is an important method of validating any approach to frailty operationalization. Note that our intent in checking the ability of the frailty index to predict mortality is validation of the index, rather than developing a mortality prediction index that included frailty. If the frailty index were meant to be a mortality prediction instrument, there might be a rationale for weighting several items, particularly age [26]. One notable result from the Cox analyses is that including the Frailty Index increased the impact of being male on mortality. This likely reflects the observation from earlier stud-
ies that while men accumulate fewer deficits than do women, any given level of deficit accumulation is more lethal for them and at any given age, females seem to be more frail than males [6,11].

The relationship with mortality is also important in understanding how deficit accumulation might operate. Classically, Gompertz described the rate of mortality being exponentially related to age [27]. Equally unsurprisingly, mortality exponentially increases with the accumulation of deficits [5,8,19]. In addition, acceleration of deficit accumulation is characteristic of older people prior to death [8].

Our data must be interpreted with caution. Not all items had established cut-points. In addition, cut-points can be difficult to apply across a sample that covers many ages, as the effects of continuous traits can be age-specific.[28] Our approach derived cut-points based on the “interim frailty index” procedure described above. In addition, the sample is small, so that any individual estimates can be unstable; this is where aggregation of items in a frailty index can be helpful, and where the re-sampling strategy is useful.

Our paper also has some strengths. In replicating many of the characteristics of a frailty index in a new sample, we can give additional assurance of the robustness of the approach. By spelling out in detail how each step in constructing a frailty index can be undertaken, and by submitting to an open access journal, we are aiming to make the method widely available. We have also made more precise a method for establishing cut-points for variables that were not constructed for inclusion in a frailty index, thereby further allowing the method to be used. In this regard, the relationship of any given variable to a mean frailty index score of 0.2 might seem arbitrary. In a study that related the frailty index approach to the phenotypic definition of frailty popularized from the Cardiovascular Health Study [22], 0.2 corresponded to the mean frailty index value for persons defined as "pre-frail" [7,22]. A more recent paper from another group used the 0.2 cut-point on a so-called "deficit index" to distinguish people who were "robust" from those who were "pre-frail"[29].
Finally, like many health surveys, the PEP study has many more variables than are needed to construct a 40-item frailty index. Several eligible variables were not included only because we had reached our target of a 40-item Frailty Index. There is no scientific reason not to include more – we have constructed an frailty index of 70 items. On the other hand, a recurring concern about the frailty index has been the feasibility of calculating it if a lot of variables are used [30]. Here, as in some earlier studies, [7,19,31] we have selected variables at random (bootstrapping) from a list of eligible variables to make up the Frailty Index and have again shown that the results are insensitive to the precise composition of the index.

Change in the health status of elderly people is an obvious concern to clinicians and to population planners. In the next round of analyses, we will be interested to know whether the changes in the frailty states (baseline frailty state versus the follow up state) can be described using a so called "stochastic" transition model [32] which we have evaluated in other community-dwelling elderly samples, although not with ones that include as many performance measures as the PEP study [33]. This intriguing possibility is motivating further inquiries by our group.

Conclusion
A systematic process for creating a Frailty Index was presented for the Yale Precipitating Events Project, a well studied cohort in which deficit accumulation previously had not been evaluated. The process allows operationalization of the frailty index to be carried out in other data-

![Figure 3: Variance in the Slope of the Frailty Index](image)

**Figure 3**
Variance in the Slope of the Frailty Index. The Bootstrapping of the frailty index. The frailty index was created and plotted 1000 times, each time randomly picking 80% of the variables of the index. Twenty iterations are shown here. The experimental and best fit regression lines of the average index values are shown in the baseline (red) and follow up (blue).

Table 3: Cox Analyses

| Analysis   | Variable  | Baseline   | Follow up  |
|------------|-----------|------------|------------|
|            |           | HR 95% CI  | HR 95% CI  |
| Bi-variate | Age       | 1.09 1.07 – 1.11 | 1.09 1.06 – 1.11 |
|            | Frailty Index | 1.03 1.02 – 1.04 | 1.05 1.04 – 1.05 |
|            | Male Sex   | 1.46 1.16 – 1.82 | 1.37 1.07 – 1.74 |
| Multi-variatea | Age       | 1.08 1.06 – 1.10 | 1.06 1.04 – 1.09 |
|            | Frailty Index | 1.03 1.02 – 1.04 | 1.04 1.04 – 1.05 |
|            | Male Sex   | 1.80 1.42 – 2.27 | 1.71 1.33 – 2.20 |

Cox Regression analyses of the baseline and follow up. Calculations were based on monthly follow ups for nine years from the baseline interview (or seven and a half years from the 18 month follow up). The frailty index hazard ratios (HR) are calculated with % levels of the index (i.e. the HR measures a change of 0.01 on the index).

* Analysis done with Age, Sex and Frailty Index as covariates.
sets. The frailty index reveals how frailty, understood as a vulnerability state with an increased risk of adverse outcomes, can be quantified. This method of quantifying frailty can aid our understanding of health and frailty-related health characteristics and outcomes in older adults.

Competing interests
No sponsor had a role in the decision to undertake these analyses or to submit the study for publication. Each author asserts no proprietary interest in the result and no financial conflict of interest.

Authors’ contributions
Sam Seale carried out the analyses and wrote the first draft as part of his PhD program. Arnold Mitnitski supervised these analyses. Evelyne Gahbauer provided the datasets. Thomas Gill is the PI of the PEP study and critiqued each draft. Kenneth Rockwood conceived of the idea with Arnold Mitnitski, with whom he arranged funding and co-wrote the first and subsequent drafts. All authors reviewed and approved the final draft of the paper.

Acknowledgements
Funds for secondary analysis came from the Canadian Institutes of Health Research, through an operating grant to Dr. Mitnitski (MOP62823) and by a grant from the Fountain Innovation Fund of the Queen Elizabeth II Health Sciences Foundation. Dr. Gill is the recipient of a Midcareer Investigator Award in Patient-Oriented Research (K24AG021507) from the National Institute on Aging. Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation, and the National Institute on Aging Research, the Patrick and Catherine Weldon Donaghue Medical Institute on Aging. Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation. Dr. Gill is the recipient of a Midcareer Investigator Award in Patient-Oriented Research (K24AG021507) from the National Institute on Aging. Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation, and the National Institute on Aging. Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation. Dr. Gill is the recipient of a Midcareer Investigator Award in Patient-Oriented Research (K24AG021507) from the National Institute on Aging. Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation, and the National Institute on Aging. Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation. Dr. Gill is the recipient of a Midcareer Investigator Award in Patient-Oriented Research (K24AG021507) from the National Institute on Aging. Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation, and the National Institute on Aging. Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation. Dr. Gill is the recipient of a Midcareer Investigator Award in Patient-Oriented Research (K24AG021507) from the National Institute on Aging. Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation, and the National Institute on Aging. Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation. Dr. Gill is the recipient of a Midcareer Investigator Award in Patient-Oriented Research (K24AG021507) from the National Institute on Aging. Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation, and the National Institute on Aging.

References
1. Bergman H, Ferrucci L, Guralnik J, Hogan DB, Hummel S, Karunan- than S, Wolfson C. Frailty: an emerging research and clinical paradigm–issues and controversies. J Gerontol A Biol Sci Med Sci 2007, 62:731-737.
2. Fisher AL: Just what defines frailty? J Am Geriatr Soc 2005, 53:2229-2230.
3. Whitson HE, Purser JL, Cohen HJ: Frailty thy name is ... Frailty? J Gerontol A Biol Sci Med Sci 2007, 62(7):728-730.
4. Mitnitski AB, Mogilner AJ, Rockwood K: Accumulation of deficits as a proxy measure of aging. The Scientific World 2001, 1:323-336.
5. Rockwood K, Mitnitski A: Frailty in relation to the accumulation of deficits. J Gerontol A Biol Sci Med Sci 2007, 62A:722-727.
6. Mitnitski A, Xiaowei S, Skoog I, Broe GA, Cox JL, Grunfeld E, Rock- wood K: Relative fitness and frailty of elderly men and women in developed countries and their relationship with mortality. J Am Geriatr Soc 2005, 53(12):2184-2189.
7. Rockwood K, Andrew M, Mitnitski A: A Comparison of two approaches to measuring frailty in elderly people. J Gerontol A Biol Sci Med Sci 2007, 62A(No 7):738-743.
8. Kulminski A, Yashin A, Arbeek K, Akushevich I, Ukraintseva S, Land K: Cumulative index of health disorders as an indicator of aging-associated processes in the elderly: Results from analyses of the National Long Term Care Survey. Mech Ageing Dev 2007, 128:250-258.
9. Kulminski A, Ukraintseva SV, Akushevich IV, Arbeek KG, Yashin A: Cumulative index of health deficiencies as a characteristic of long life. J Am Geriatr Soc 2007, 55(4):925-46.
10. Goggins WB, Woo J, Sham A, Ho SC. Frailty index as a measure of personal biological age in a Chinese population. J Gerontol A Biol Sci Med Sci 2005, 60A:1046-1051.
11. Woo J, Goggins W, Sham A, Ho SC. Social determinants of frailty. J Am Geriatr Soc 2005, 51:402-408.
12. Bergman H, Ferrucci L, Guralnik J, Hogan DB, Hummel S, Karunan- than S, Wolfson C. Frailty: an emerging research and clinical paradigm–issues and controversies. J Gerontol A Biol Sci Med Sci 2007, 62(7):731-737.
13. Walston J, Hadley EC, Ferrucci L, Guralnik JM, Newman AB, Stude- nski SA, Ershler WB, Harris T, Fried L: Research agenda for frailty in older adults: toward a better understanding of physiology and etiology: summary from the American Geriatrics Societies/National Institute on Aging Research Conference on Frailty in Older Adults. J Am Geriatr Soc 2006, 54(4):991-1001.
14. Ferrucci L, Guralnik JM, Studenski S, Fried LP, Cutler GB Jr, Walston JD: Interventions on frailty working group. Designing rand- omized, controlled trials aimed at preventing or delaying functional decline and disability in frail, older persons: a consensus report. J Am Geriatr Soc 2004, 52(4):625-634.
15. Gill TM, Gahbauer EA, Allore HG, Ham L: Transitions between frailty states among community-living older persons. Arch Intern Med 2006, 166:418-423.
16. Gill TM, Desai MM, Gahbauer EA, Holford TR, Williams CS: Restricted activity among community-living older persons: incidence, precipitants, and health care utilization. Ann Intern Med 2001, 135:331-332.
17. Folstein M, Folstein S, McHugh P: “Mini-mental state”. A practical method for grading the cognitive state of patients for the clinician. J Psychiatr Res 1975, 12:189-198.
18. Mitnitski A, Rockwood K: Transitions in cognitive test scores over 5 and 10 years in elderly people: evidence for a model of age-related deficit accumulation. BMC Geriatr 2008, 8(1):3.
19. Rockwood K, Mitnitski A, Song X, Steen B, Skoog I: Long-term risks of death and institutionalization of elderly people in relation to deficit accumulation at age 70. J Am Geriatr Soc 2006, 54(6):975-979.
20. Reisberg B: Functional Assessment Staging (FAST). Psychopa- thological Bull 1988, 24:65-69.
21. Streiner D, Norman G: Health Measurement Scales: A guide to their development and use 3rd edition. Oxford, Oxford University Press; 2003:172-193.
22. Fried LP, Tangen CM, Walston J, Newman AB, Hirsch C, Gottdiener J, Seeman T, Tracy R, Kop WJ, Burke G, McBurnie MA: Cardiovascular Health Study Collaborative Research Group: Frailty in older adults: evidence for a phenotype. J Gerontol A Biol Sci Med Sci 2001, 56A:M146-M156.
23. Efron B, Tibshirani R: An Introduction to the bootstrap New York, Chapman & Hall; 1993.
24. Rockwood K, Mitnitski A: Limits to deficit accumulation in elderly people. Mech Ageing Dev 2006, 127(5):494-6.
25. Rockwood K, Mitnitski A: frailty, fitness and the mathematics of deficit accumulation. Rev Clin Gerontol 2007, 17:1-12.
26. Lee SJ, Lindquist K, Segal MR, Covinsky KE: Development and vali- dation of a prognostic index for 4-year mortality in older adults. JAMA 2006, 295(7):801-8. Erratum in: JAMA 2006, 295(16):1900.
27. Gomperz B: On the nature of the function expressive of the law of human mortality and on a new mode of determining life contingencies. In Philosophical Transaction of the Royal Society of London 1825 San Diego: Academic Press.
28. Yashin AI, Arbeek KG, Akushevich I, Kulminski A, Ukraintseva S, Ukraintseva SV: Stochastic model for analysis of longitudinal data on aging and mortality. Math Biosci 2007, 208:538-551.
29. Kulminski AM, Ukraintseva SV, Kulminskaya IV, Arbeek KG, Land K, Yashin AI: Cumulative deficits better characterize susceptibil- ity to death in elderly people than phenotypic frailty: lessons from the Cardiovascular Health Study. J Am Geriatr Soc 2008, 56:898-903.
30. Martin FC, Brighton P: Frailty: different tools for different pur- poses? Age Ageing 2008, 37(2):129-31.
31. Rockwood K, Abeysundera MJ, Mitnitski A: How should we grade frailty in nursing home patients? J Am Med Dir Assoc 2007, 8(9):595-603.
32. Mitnitski A, Bao L, Rockwood K: Going from bad to worse: a stochastic model of transitions in deficit accumulation, in relation to mortality. Mech Ageing Dev 2006, 127:490-493.
33. Mitnitski A, Song X, Rockwood K: Improvement and decline in health status from late middle age: modeling age-related changes in deficit accumulation. Exp Gerontol 2007, 42(11):1109-15.
34. Flegal KM, Graubard BI, Williamson DF, Gail MH: Cause-specific excess deaths associated with underweight, overweight and obesity. JAMA 2007, 298(17):2028-2037.

Pre-publication history
The pre-publication history for this paper can be accessed here:

http://www.biomedcentral.com/1471-2318/8/24/prepub