Frailty in Older People

Summary

Frailty is the most problematic expression of population ageing. It is a state of vulnerability to poor resolution of homeostasis following a stress and is a consequence of cumulative decline in multiple physiological systems over a lifespan. This cumulative decline erodes homeostatic reserve until relatively minor stressor events trigger disproportionate changes in health status, typically a fall or delirium. Landmark studies have developed valid models for frailty and these have allowed epidemiological studies that demonstrate the association of frailty with adverse health outcomes. New research is needed to develop more efficient methods to detect and severity grade frailty as part of routine clinical practice, particularly methods with utility for primary care. This would greatly inform the appropriate selection of older people for invasive procedures or medications and would be the basis for a paradigm shift in the care of frail older people towards a more appropriate goal-directed care.

Background

Population ageing worldwide is rapidly accelerating from 461 million people aged over 65 years in 2004 to an estimated 2 billion people by 2050 (1, 2), which has profound implications for the planning and delivery of health and social care. The most problematic expression of population ageing is the clinical condition of frailty. Frailty develops as a consequence of age-related decline in multiple physiological systems, which collectively results in a vulnerability to sudden health status changes triggered by relatively minor stressor events. It is estimated that a quarter to a half of people over 85 years are frail and these people have significantly increased risk of falls, disability, long-term care and death (3, 4). Importantly, up to three quarters of people over 85 years might not be frail, raising the questions of how frailty develops, how it might be prevented and how it can be detected reliably. This review addresses these issues.

Frailty definition and presentations

Frailty is a state of increased vulnerability to poor resolution of homeostasis following a stress, which increases the risk of adverse outcomes including falls, delirium and disability (3, 5, 6). It is a long established clinical expression that implies concern over an older person’s vulnerability and prognosis. This is shown diagrammatically in figure 1, in which

Contributors

AC and JY initiated and coordinated the review. All authors contributed to the writing of the review.

Conflict of interest

KR has applied to various Canadian government schemes to commercialise a new version of the frailty index based on a comprehensive geriatric assessment and a new screening tool, adapted from the Clinical Frailty Scale of the Canadian Study of Health and Aging. To do this, with colleagues he has formed a company called Videx Canada. He has not asserted copyright on any frailty measure discussed here.

AC, JY, SI and MOR declare that they have no conflicts of interest.
an apparently small insult (e.g. a new drug; “minor” infection; or “minor” surgery) results in a dramatic and disproportionate change in health state: from independent to dependent; mobile to immobile; postural stability to falling; lucid to delirious. The dependency oscillations observed in frail older people has been referred to as “unstable disability” to reflect the often marked changes in functional ability that are familiar to practitioners working with older people (7). The common clinical presentations of frailty are given in table 1.

Pathophysiology of frailty

Frailty is a disorder of multiple inter-related physiological systems. There is a gradual decline in physiological reserve with ageing but, in frailty, this decline is accelerated and homeostatic mechanisms start failing (8, 9). An important perspective for frailty, therefore, is to consider how the complex mechanisms of ageing promote cumulative decline in multiple physiological systems, consequent erosion of homeostatic reserve and vulnerability to disproportionate changes in health status following relatively minor stressor events. These complex ageing mechanisms are influenced by underlying genetic and environmental factors (10) in combination with epigenetic mechanisms, which regulate the differential expression of genes in cells and may be especially important in ageing (11, 12).

A schematic representation of frailty is provided in figure 2.

The pathway to frailty

Ageing is considered to result from the lifelong accumulation of molecular and cellular damage caused by multiple mechanisms under the regulation of a complex maintenance and repair network (10). There is uncertainty regarding the precise level of cellular damage required to cause impaired organ physiology but, importantly, many organ systems exhibit considerable redundancy, which provides the physiological reserve required to compensate for age and disease-related changes (13). For example, the brain contains more neurons and skeletal muscle more myocytes than are required for survival (13). Therefore, a key question is whether there is a critical threshold of age-related, cumulative decline in multiple physiological systems beyond which frailty becomes evident. A 2009 cross-sectional study involving 1,002 female participants investigated cumulative physiological dysfunction in six different systems (haematological, inflammatory, hormonal, adiposity, neuromuscular and micronutrients) using 12 measures and reported a non-linear relationship between the number of abnormal systems and frailty, independent of age and comorbidity (14). The presence of abnormal results in three or more systems was a significant predictor of frailty. Importantly, the number of abnormal systems was more predictive than abnormalities in any particular system. This provides evidence to suggest that when physiological decline reaches an aggregate critical mass, frailty becomes evident (14).

The brain, endocrine system, immune system and skeletal muscle are intrinsically inter-related and are currently the organ systems best studied in the development of frailty (5). These systems will be considered in greater detail, but it is important to recognise that frailty has also been associated with loss of physiological reserve in the respiratory (15).
cardiovascular (16), renal (17) and haemopoietic and clotting systems (18, 19) and that nutritional status can also be a mediating factor (3, 20-22).

The frail brain

Ageing is associated with characteristic structural and physiological changes in the brain. The loss of individual neurons in the majority of cortical regions is minimal (23), but neurons with high metabolic demands, for example the hippocampal pyramidal neurons, may be disproportionately affected by altered synaptic function, protein transport and mitochondrial function (23). The hippocampus has been identified as an important mediator in the pathophysiology of cognitive decline and Alzheimer’s dementia (24) and is a key component of the stress response, sensing increased glucocorticoid levels and relaying information to the hypothalamus in a negative feedback loop (25).

The ageing brain is also characterised by structural and functional changes to microglial cells, which are the resident immune cell population of the central nervous system (CNS) and are the CNS equivalent of macrophages. They are activated by brain injury and local and systemic inflammation and become primed (hyper-responsive) to small stimuli with ageing, which can potentially cause damage and neuronal death (26-28). Primed microglia are postulated to play an important role in the pathophysiology of delirium (28, 29). A prospective cohort study involving 273 hospitalised older people identified that frailty is associated with both increased risk of developing delirium (odds ratio, OR, 8.5, 95% confidence interval, CI, 4.8-14.8) and subsequent reduced survival (median survival in frail older patients with delirium 88 days (95% CI 5-171 days); median survival in non-frail older patients with delirium 359 days (95% CI 118-600 days)) (6). This indicates that the combination of delirium and frailty identifies older people at particularly high risk of adverse outcomes.

There is accumulating evidence from observational studies to support a temporal association between frailty, cognitive impairment and dementia. A prospective cohort study (n=750) of older people without cognitive impairment at baseline reported that frailty was associated with an increased risk of developing mild cognitive impairment over 12 years of follow-up (hazard ratio, HR, 1.63, CI, 1.27-2.08) (30). Increasing degree of frailty was also associated with a faster rate of cognitive decline. An independent association between frailty and dementia has been reported in two large prospective cohort studies (31, 32).

The frail endocrine system

The brain and endocrine system are intrinsically linked through the hypothalamo-pituitary axis, which controls metabolism and energy use via the signaling action of a series of homeostatic hormones (23). During ageing, there is a decline in production of three major circulating hormones. Firstly, a decline in growth hormone synthesis by the pituitary causes a reduction in insulin-like growth factor-1 (IGF-1) production by the liver and other organs. IGFs are a family of small peptides that increase anabolic activity in many cells. Promotion of neuronal plasticity and increased skeletal muscle strength are considered particularly important effects (33). Secondly, decreased oestradiol and testosterone cause increased release of luteinising hormone (LH) and follicle stimulating hormone (FSH). Thirdly, the
adrenocortical cells that produce the major sex steroid precursor dehydroepiandrosterone (DHEA) and DHEA sulphate (DHEAS) decrease in activity, often mirrored by a gradual increase in cortisol release (34, 35).

Changes to IGF signaling, sex hormone, DHEA/DHEAS production and cortisol secretion are considered important in frailty, although the exact relationships remain uncertain and require further investigation. One cross-sectional study reported significantly lower levels of IGF-1 in people identified as frail when compared to age-matched controls (36). However, if IGF-1 has a key aetiological role in frailty an association between IGF-1 and mortality might be anticipated but a series of observational studies have reported inconsistent associations (37-41). Furthermore, although the muscles of frail older people appear to retain the capability to respond to IGF-1 (42) trials of IGF-1 supplementation in older people have failed to demonstrate benefit (43).

Although an association between testosterone levels and frailty has been identified (44) this may be a sensitive marker rather than a pathological mechanism (45). A cross-sectional study reported an association between DHEAS and frailty but the influence of comorbid conditions could not be confidently excluded (46). A U-shaped association between DHEAS and mortality has been reported in disabled older women (47).

One cross sectional study (n=214) reported that frailty was independently associated with chronically elevated diurnal cortisol levels (48). A link between chronically elevated cortisol and frailty is plausible, as persistently high levels of cortisol are associated with increased catabolism, leading to loss of muscle mass, anorexia, weight loss and reduced energy expenditure - cardinal clinical features of frailty (49).

The frail immune system

The ageing immune system is characterised by a decline in stem cells, alterations in T-lymphocyte production, blunting of the B-cell led antibody response and reduced phagocytic activity of neutrophils, macrophages and natural killer cells (50, 51). This senescent immune system may function adequately in the quiescent state but fail to respond appropriately to the stress of acute inflammation (50). There is evidence that inflammation has a key role in the pathophysiology of frailty through an abnormal, low-grade inflammatory response that is hyper-responsive to stimuli and that persists for a prolonged period following removal of the initial inflammatory stimulus (19, 52-57). A range of inflammatory cytokines have been independently associated with frailty including interleukin-6 (IL-6), C-reactive protein (CRP), tumour necrosis factor-α (TNFα) and CXC chemokine ligand-10 (CXCL-10), a potent pro-inflammatory mediator (52, 54-58). However, higher CRP levels in very old people have also been associated with better memory function (59). Advanced glycation end products (AGEs) are a group of molecules produced by the glycation of proteins, lipids and nucleic acid that have potential to cause widespread cellular damage by upregulation of inflammation (60). They have been associated with ageing, chronic disease and mortality and may have an important role in frailty (60).

Inflammation is associated with anorexia and catabolism of skeletal muscle and adipose tissue, which may contribute to the nutritional compromise, muscle weakness and weight
loss that characterise frailty (41, 61, 62). Furthermore, frailty is associated with an impaired antibody response to influenza (63) and pneumococcal vaccine (64), which helps explain the observation that vaccination in older people is associated with only relatively modest clinical effectiveness (65).

Frail skeletal muscle - sarcopenia

Sarcopenia has been defined as progressive loss of skeletal muscle mass, strength and power and is considered a key component of frailty (66, 67). Loss of muscle strength and power may be more important than changes to muscle mass (68). Under normal circumstances, muscle homeostasis is maintained in a delicate balance between new muscle cell formation, hypertrophy and protein loss. This delicate balance is coordinated by the brain, endocrine system and immune system and is influenced by nutritional factors and level of physical activity. The adverse neurological, endocrine and immune components of frailty have the potential to upset this delicate homeostatic balance and accelerate the development of sarcopenia. Inflammatory cytokines including IL-6 and TNFα activate muscle breakdown to generate amino acids for energy and cleave antigenic peptides (69). This fundamentally protective response may become pathological in the presence of an overactive, insufficiently regulated inflammatory response that characterises frailty, leading to loss of muscle mass and strength, with attendant decline in functional ability.

Frailty models

Reliable frailty models should be assessed against their success in predicting both natural history and response to therapeutic interventions and be underpinned by biological principles of causality (70). The two principal emerging models of frailty are the phenotype model (3) and the cumulative deficit model underpinning the Canadian Study of Health and Aging (CSHA) Frailty Index (71).

The phenotype model

In a landmark study, Fried and colleagues (3) undertook a secondary analysis of data obtained from a prospective cohort study (the Cardiovascular Health Study (CHS)) involving 5210 men and women aged 65 years and older. A frailty phenotype was operationalised using a cluster of variables: unintentional weight loss; self-reported exhaustion; low energy expenditure; slow gait speed; weak grip strength (table 2). The lowest quintile values were used to define absence/presence of these variables. People with Parkinson’s disease, previous stroke, cognitive impairment or depression were excluded. People with three of the five factors were considered frail, one or two factors as pre-frail, and no factors as robust older-people. The population so defined was categorised as 7% frail, 47% pre-frail and 46% not frail. Follow-up assessments were undertaken at 3 and 5 years with the outcomes of falls; mobility and function; hospitalisation; and death.

People categorised as frail were observed to have more adverse outcomes compared to people categorised as not frail, with the pre-frail group having outcomes intermediate between the two. Observed mortality at 7 years was 12%, 23% and 43% for not frail, pre-
frail and frail groups respectively. The 7-year adjusted hazard rate for mortality was 1.63 (95% CI 1.27-2.08) for the frail group (3).

This work is important as it suggests a frailty phenotype can be defined and might be a basis to detect frailty in routine care. However, it is not presently clear how the variables might reliably be translated into clinical practice. Additionally, the five factors were fortuitously available and selected from a prospective cohort study that was not designed to investigate frailty. Other potentially important factors such as cognitive impairment, a highly prevalent condition associated with functional decline and disability, were not included as a component of the phenotype (72). Nonetheless, despite these criticisms, the general approach of clusters of variables to define a frailty phenotype has been independently validated (73, 74).

The cumulative deficit model

The Frailty Index (FI) was developed as part of the CSHA (71); a five year prospective cohort study (n=10,263) designed to investigate the epidemiology and burdens of dementia in older people in Canada (mean age: 82 years). Ninety-two baseline parameters of symptoms (e.g. low mood), signs (e.g. tremor) and abnormal laboratory values, disease states and disabilities, collectively referred to as deficits, were used to define frailty (75). The FI was a simple calculation of the presence or absence of each variable as a proportion of the total (e.g. 20 deficits present out of a possible 92 gives a FI of 20/92 = 0.22). Thus frailty is defined as the cumulative effect of individual deficits - ‘the more individuals have wrong with them, the more likely they are to be frail’ (76).

The statistical distribution of the FI (a gamma distribution) was consistent with a probability model that typically describes systems with inbuilt redundancy. This is an attractive mathematical model for frailty as it implies that the FI has properties that fully support the concept of reduced reserve. Thus, although each individual deficit carries no obvious or imminent threat for mortality (e.g. hearing impairment), the deficits cumulatively contribute to an increased risk of death. This is consistent with the increased vulnerability and threat of impending homeostatic failure that is essential to the frailty concept. Importantly, the cumulative deficit model expresses the notion of a gradation of frailty with progressive accumulation of deficits each of which has an equal weight in mathematical modeling of the FI. This is attractive clinically because it allows frailty to be considered as gradable rather than present/absent. Moreover, it is the number of equally weighted deficits, as a measure of accumulated vulnerability, that is related to adverse outcomes, rather than particular clusters of deficits (77). Importantly, a value of 0.67 appears to identify a level of frailty beyond which further deficit accumulation is not sustainable and death is likely to supervene (78). This value may represent the warning sign of being close to the ‘tipping point’ that characterises a complex system on the brink of collapse (79).

Subsequent work has demonstrated that the rather daunting initial list of 92 variables can be reduced to a more manageable 30 or so without loss of predictive validity (4). The criteria for variable inclusion into the FI are: biologically sensible; accumulates with age; does not saturate too early (80). This makes the FI very adaptable as a conceptual approach.
Several studies using data from the CSHA demonstrated that the FI was strongly related to the risk of death and institutionalisation (4, 77). The 10-year adjusted hazard rate for mortality was 1.57 (95% CI 1.41-1.74) for the frail group (4), a statistically similar estimate to the phenotype model (3). Mortality has previously been demonstrated to be exponentially related to the value of the FI (81).

**Comparing the phenotype model with the cumulative deficit model**

The phenotype and cumulative deficit models demonstrate overlap in identification of frailty (82) and considerable statistical convergence (83). This is especially important because the demonstration of convergent predictive validity for adverse health outcomes between two conceptually different models of frailty may help advance the debate regarding whether frailty is best considered as a syndrome or a state by lending support for recognition of the condition as a unified construct.

The continuous FI demonstrated greater discriminatory ability for people with moderate and severe frailty when compared to the categorical phenotype model, a finding that has been independently validated (84). Use of continuous models may facilitate more precise identification of frail older people for interventions to improve outcomes and a 2011 study re-scaled the phenotype model to make it more continuous, providing better discriminatory capacity (85).

**Epidemiology**

The importance of frailty as a leading cause of death in older people comes from a ten-year prospective cohort study involving community dwelling older people (n=754) (86). Cause of death was based on clinical home-based assessments conducted at 18-month intervals and death certificates. The most common condition leading to death was frailty (27.9%) compared to organ failure (21.4%), cancer (19.3%), dementia (13.8%) and other causes (14.9%).

**Prevalence of frailty**

A recent systematic review investigated the prevalence of frailty (87). Twenty-one community based cohort studies involving 61,500 older people were identified. The operational definitions for frailty and the inclusion/exclusion criteria varied between the studies, which largely explained the considerable variation in reported frailty prevalence rates of 4.0 to 59.1%. When the reported rates were restricted to the studies that used the phenotype model, the weighted average frailty prevalence rate was 9.9% (95% CI 9.6-10.2), and 44.2% for pre-frailty prevalence (95% CI 44.2-44.7). Frailty was statistically more prevalent in women (11 studies, 9.6%; 95% CI 9.2-10) than men (5.2%; 95% CI 4.9-5.5). Frailty increased steadily with age: 65-69 years: 4%; 70-74 years: 7%; 75-79 years: 9%; 80-84 years: 16%; ≥85 years: 26%. Rates appear to be higher in studies that employed the graded frailty index, which would count as frail some people whose increased risk is captured in the pre-frail category of the phenotype model (4).
Most frailty models were developed in Caucasian populations and the prevalence of frailty may be higher in inhabitants of Southern Europe (88) and older Hispanic and African Americans (3, 89) so different cut-offs for frailty may be required in different populations.

**Adverse outcomes of frailty**

Associations between frailty and adverse outcomes reported in four large prospective cohort studies (3, 90-92) are presented in table 3.

**Frailty transitions**

Frailty is a dynamic process (93) but a transition to a level of greater frailty is more common than improvement and the development of frailty frequently results in a spiral of decline that leads to increasing frailty and risk of worsening disability, falls, admission to hospital and death (94). Risk of admission to long-term care is also increased in those with mild frailty (adjusted risk ratio (RR) 2.54, 95% CI 1.67-3.86) and moderate/severe frailty (RR 2.60, 95% CI 1.36-4.96) (95).

**Frailty, comorbidity and disability**

The CHS population was used to investigate the overlap between frailty, comorbidity and disability (3, 96). Frailty and comorbidity (defined as two or more of the following nine diseases: myocardial infarction; angina; congestive heart failure; claudication; arthritis; cancer; diabetes; hypertension; chronic obstructive pulmonary disease) was present in 46.2% of the population, frailty and disability (defined as the presence of restriction in at least one activity of daily living) was present in 5.7%, and the combination of frailty, disability and comorbidity was present in 21.5% of the study group. Importantly, frailty was present without comorbidity or disability in 26.6% of the study group. This finding provides support for frailty as an independent concept, distinct from comorbidity and disability. However, more recent work suggests that the overlap is more frequent and increases with greater frailty (97). The contribution of subclinical disease may be particularly important and physiological measurements to identify older people at risk of frailty may help guide the development of earlier preventative interventions (85).

**The instrumentation of frailty**

The demonstration of large between-group differences for people who are frail compared to not frail (4) is important because it entreats clinicians away from judgments based on chronological age towards the notion of frailty. Researchers and clinicians, therefore, require simple, valid, accurate and reliable tools to detect frailty. Monitoring outcomes of interventions in frail people additionally requires tools that are sensitive to change (98).

**Standardised questionnaires to identify frailty**

A systematic review with a broadly based study selection criteria identified 20 candidate tools for frailty (99). However, most of the included studies described either primary research to investigate models of frailty or the focus was on functional restriction, which, although an important manifestation of frailty, is insufficient for reliable identification of the condition. The Frail Elderly Functional Questionnaire (19 items) was identified as a
potential outcome measure for frailty intervention studies as it is suitable for use by telephone or proxy, valid and reliable (100), and is sensitive to change (101).

The Groningen Frailty Indicator (102) and the Tilburg Frailty Indicator (103) are simple and similar (104) questionnaire based approaches to detecting people with frailty. Aspects of validity have been investigated but, importantly, studies of diagnostic accuracy against well-defined community populations of older people are not yet available. Moreover, both these questionnaires require new information to be gathered. The option of using existing patient data held in primary care records to construct a frailty index consistent with the cumulative deficit model requires investigation.

Assessments to identify frailty

The timed-up-and-go test (TUGT), a simple standardised measure of gait speed that requires a stop watch (105), and hand grip strength that requires a hand-held dynamometer (106), have been investigated as potential single assessments to detect frailty. Pulmonary function is associated with frailty (15) and may have utility as a straightforward detection test. However, diagnostic accuracy of these assessments has not been confirmed. A systematic review identified nine prospective studies (n=34,485) investigating slow gait speed (107). Slow gait speed successfully characterised the sub-group of older people who had adverse outcomes and had similar accuracy to complex multivariate models that included itemising chronic conditions.

The Edmonton Frail Scale is a multi-dimensional assessment instrument that includes the TUGT, and a test for cognitive impairment (108). It is quick to administer (less than 5 minutes) and is valid, reliable and feasible for routine use by non-geriatricians but the diagnostic accuracy has not been investigated.

The various interRAI instruments are widely used internationally to standardise the assessment of older people. Nine items that are embedded in many of the instruments can be extracted and comprise the Changes in Health, End-Stage Disease and Signs and Symptoms (CHESS) scale. Although not explicitly a frailty measure (109), CHESS has been demonstrated to be a strong predictor of mortality and further validation studies are ongoing.

Comprehensive geriatric assessment (CGA) has become the internationally established method to assess older people in clinical practice. It is a multidisciplinary diagnostic process to determine an older person’s medical, psychological and functional capability to develop a plan for treatment and follow up (110). The process, provided it is closely linked to interventions, is associated with superior outcomes (111) and has been applied successfully beyond elderly care medicine (112, 113). Two studies embedded in the CSHA programme were used to investigate the predictive validity of CGA as conducted by over 70 clinicians (71, 114). In both studies, the clinically obtained CGA results were highly correlated with the research standard CSHA FI and were predictive of death and need for institutional care. These studies are the first objective confirmation that CGA is sensitive to the reliable detection of degrees of frailty. CGA is currently the gold standard to detect frailty and it should be more widely deployed. The practical limitation of CGA is the time and expertise required for the process.
Interventions for frailty

Reducing the prevalence or severity of frailty is likely to have large benefits for the individual, their families and for society. Several approaches have been investigated in clinical trials.

Interventions based on comprehensive geriatric assessment

Frail older people receiving inpatient CGA are more likely to return home, less likely to experience cognitive or functional decline and have lower in-hospital mortality (111). Complex interventions based on CGA delivered to older people in the community can increase the likelihood of continuing to live at home, principally through a reduced need for care home admission and reduced falls (115, 116), but those who are most frail appear to receive least benefit (115).

Exercise interventions

Exercise has physiological effects on the brain, endocrine system, immune system and skeletal muscle (42, 117-120). Three systematic reviews of home-based and group-based exercise interventions for frail older people concluded that exercise can improve outcomes of mobility and functional ability (121-123). Meta-analysis identified that the effect sizes are likely to be small to moderate (pooled standardised mean difference (SMD) for mobility 0.18, 95% CI 0.05-0.30; pooled SMD for functional ability 0.27, 95% CI 0.08-0.46) (121). The most effective intensity (duration and frequency) of exercise intervention remains uncertain, but adherence was characteristically high across a range of interventions.

Most trials did not use a validated measure or operationalised model to record frailty at baseline or follow-up but, where results were stratified by frailty, those who were most frail appeared to gain least benefit (124). However, this is contradictory to the results from a Cochrane review that incorporated 49 RCTs of exercise interventions for long-term care (LTC) residents (a group of older people who are likely to be very frail) and concluded that interventions, particularly those involving strength and balance training, can successfully increase muscle strength and functional abilities (125). It is therefore possible that even small gains in strength of LTC residents translate into important functional gains.

Nutritional interventions

Nutritional interventions may have potential to address the impaired nutrition and weight loss of frailty. However, there is a paucity of evidence. One RCT that investigated the effects of exercise and nutritional supplementation in 100 frail older people living in long-term care reported that nutritional supplementation had no effect on muscle strength, gait speed, stair climbing or physical activity (126). A Cochrane review of nutritional interventions for preventing and treating pressure ulcers in older hospital patients, a group who are likely to be frail, reported that it was not possible to draw any firm conclusions due to the absence of trials of high methodological quality (127).
Pharmacological agents

Few pharmacological agents have been investigated in frailty. Angiotensin converting enzyme (ACE) inhibitors have been demonstrated to improve the structure and biochemical function of skeletal muscle (128) and there is evidence that ACE inhibitors may halt or slow the decline in muscle strength in older age (129) and improve exercise capacity and quality of life (130). Testosterone improves muscle strength but also increases adverse cardiovascular and respiratory outcomes (131). IGFs have direct effects on skeletal muscle (5) but IGF-1 does not appear to improve muscle strength or bone density in healthy older women (132). Low vitamin D levels have been associated with frailty (133) and vitamin D has been demonstrated to improve neuromuscular function (134). Although vitamin D prescription for older people who are deficient may reduce falls (135) and use of calcium/vitamin D supplements for older people in long-term care can reduce fractures (136), the general use of vitamin D as treatment for frailty remains controversial (43). The use of pharmacological agents for the prevention and treatment of frailty is an important area for future research.

Conclusions

Modern healthcare systems are largely organised around a single system illness (137). Many older people, however, have multi-organ problems. Frailty is a practical, unifying concept in the care of these older people that directs attention away from organ specific diagnoses towards a more holistic viewpoint of the patient and their predicament. It is a state of vulnerability to poor resolution of homeostasis following a stress and is strongly associated with adverse outcomes. Distinguishing older people who are frail from people who are not frail should therefore form an essential aspect of assessment in any health care encounter that might result in an invasive procedure or potentially harmful medication. It allows practitioners to weigh up benefits and risks, and for patients to make properly informed choices. Failure to detect frailty potentially exposes patients to interventions from which they may not benefit and indeed may be harmed. Conversely, to exclude physiologically well (non-frail) older people simply on the basis of age is unacceptable.

The most evidence-based process to detect and severity grade frailty is the process of CGA. This is a resource intensive process and new research is urgently required to find equally reliable but more efficient and responsive methods for routine care. The necessary requirements of future frailty instruments, including the important issue of clinical sensibility, have been defined (138). This might be achieved by development and further validation of the currently available frailty-specific multi-dimensional questionnaires, but the utility of existing clinical data sets, particularly in primary care, is attractive. This approach would be underpinned by the cumulative deficits frailty model and implies that frailty could be both positively identified and severity graded. Such a simple tool would facilitate essential research to gain deeper insight into the complex mechanisms of frailty and aid the development and evaluation of interventions to improve outcomes. It would also have considerable clinical merit as it would be the basis for a paradigm shift in the care of frail older people towards a more appropriate goal-directed care in which individually framed clinical outcomes that span organ systems are negotiated with patients (139).
Search strategy and selection criteria

We developed a structured search strategy with the assistance of an expert librarian at the University of Leeds. We searched the Cochrane Library (2000-2012), CINAHL (2000-2012), MEDLINE (2000-2012), EMBASE (2000-2012), PSYCINFO (2000-2012) and PEDRO (2000-2012). We used the search terms “frailty”, “frail elderly” or “sarcopenia” with terms “aged” or “aged, 80 and over” or “aging/genetics” or “longevity” or “centenarian” or “oldest old” or “very old” or “very elderly”. We performed a further search limiting the results to systematic reviews of interventions for the prevention and treatment of frailty. Additional papers were identified from personal libraries and reference lists of retrieved articles.

“It is more important to know what sort of person has a disease than to know what sort of disease a person has” Hippocrates c400BC

Acknowledgments

We would like to express our gratitude to Deirdre Andre, research support librarian at the University of Leeds, for her assistance in devising and running the search strategy for this review.

References


et al.


Lancet. Author manuscript; available in PMC 2014 July 15.


Lancet. Author manuscript; available in PMC 2014 July 15.


Lancet. Author manuscript; available in PMC 2014 July 15.
Figure 1.
Vulnerability of frail older people to a sudden change in health status following a minor illness. The green line represents a fit older person who, following a minor stress such as an infection, experiences a relatively small deterioration in function and then returns to homeostasis. The red line represents a frail older person who, following a similar stress, experiences a larger deterioration which may manifest as functional dependency and who does not return to baseline homeostasis.
Key: UTI: Urinary tract infection
Figure 2.
A schematic representation of the pathophysiology of frailty.
Ageing is considered to result from the lifelong accumulation of molecular and cellular damage caused by multiple mechanisms regulated by a complex maintenance and repair network under the influence of genetic, environmental and epigenetic mechanisms. There is uncertainty regarding the precise level of cellular damage required to cause impaired organ physiology but, importantly, many organ systems exhibit considerable redundancy, which provides the physiological reserve required to compensate for age and disease-related
changes. The brain, endocrine system, immune system and skeletal muscle are intrinsically inter-related and are currently the organ systems best studied in the development of frailty. Loss of physiological reserve in other systems including the cardiovascular, respiratory and renal systems also contributes. Influenced by level of physical activity and nutritional factors, cumulative loss of physiological reserve in these organ systems can lead to frailty, which is characterised by increased vulnerability to poor resolution of homeostasis following a stress, increasing the risk of adverse outcomes including falls, delirium and disability. These are common clinical presentations of frailty, are common reasons for admission to hospital and can accelerate further decline. Both frailty itself and these common clinical presentations identify those at increased risk of requiring care at home and long-term care admission.
<table>
<thead>
<tr>
<th>Common clinical presentations of frailty.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Non-specific:</strong></td>
</tr>
<tr>
<td><strong>Falls:</strong></td>
</tr>
<tr>
<td><strong>Delirium:</strong></td>
</tr>
<tr>
<td><strong>Fluctuating disability:</strong></td>
</tr>
</tbody>
</table>

*Lancet*. Author manuscript; available in PMC 2014 July 15.
Table 2
The five phenotype model indicators of frailty and their associated measures.

<table>
<thead>
<tr>
<th>Frailty indicator</th>
<th>Measure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight loss</td>
<td>Self-reported weight loss of more than 10 pounds or recorded weight loss of ≥5% per annum</td>
</tr>
<tr>
<td>Self-reported exhaustion</td>
<td>Self-reported exhaustion on CES-D depression score (3-4 days per week or most of the time)</td>
</tr>
<tr>
<td>Low energy expenditure</td>
<td>Energy expenditure &lt;383 KCal/week (males) or &lt;270 KCal/week (females)</td>
</tr>
<tr>
<td>Slow gait speed</td>
<td>Standardised cut-off times to walk 15 feet, stratified for sex and height</td>
</tr>
<tr>
<td>Weak grip strength</td>
<td>Grip strength, stratified by sex and BMI</td>
</tr>
</tbody>
</table>

Key. CES-D, Center for Epidemiological Studies Depression; BMI, body mass index.
Table 3: Covariate adjusted associations between frailty and adverse outcomes (falls, disability, hospitalisation, care home admission and mortality) from four large prospective cohort studies.

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Country</th>
<th>Number of participants</th>
<th>Length of follow-up</th>
<th>Falls HR/OR 95% CI</th>
<th>Worsening disability HR/OR 95% CI</th>
<th>Hospitalisation HR/OR 95% CI</th>
<th>Care home admission HR/OR 95% CI</th>
<th>Mortality HR/OR 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardiovascular Health Study (CHS) (3)</td>
<td>2001</td>
<td>US</td>
<td>5317</td>
<td>7 years</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Canadian Study of Health &amp; Aging (CSHA) (90)</td>
<td>2004</td>
<td>Canada</td>
<td>9008</td>
<td>5 years</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>OR 2.54 1.67-3.86</td>
<td>OR 2.60 1.36-4.96</td>
</tr>
<tr>
<td>Women's Health and Aging Study (WHAS) (91)</td>
<td>2006</td>
<td>US</td>
<td>1438</td>
<td>3 years</td>
<td>HR 0.92 0.63-1.64</td>
<td>NA</td>
<td>NA</td>
<td>HR 5.16 0.81-32.8</td>
<td>HR 2.75 1.46-4.99</td>
</tr>
<tr>
<td>Study of Osteoporotic Fractures (SOF) (92)</td>
<td>2008</td>
<td>US</td>
<td>6701</td>
<td>4.5 years</td>
<td>OR 1.23 1.02-1.48</td>
<td>OR 2.44 1.95-3.04</td>
<td>OR 1.89 1.66-2.14</td>
<td>OR 2.75 1.46-4.99</td>
<td></td>
</tr>
</tbody>
</table>

Key: HR, hazard ratio; OR, odds ratio; CI, confidence interval; US, United States; NA, not available.