**Fracture risk assessment in nursing homes**

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The enormous health burden consequent to an increasingly ageing demographic is a clear health priority worldwide, recognised by the World Health Organization in its recent “WHO Report on Ageing and Health”[1]. As populations age, the number of individuals who suffer from noncommunicable chronic diseases such as dementia, cardiovascular conditions, osteoporosis and sarcopenia will increase. Using osteoporotic fracture as an example, these demographic changes are projected to result in a doubling in the number of individuals at high fracture risk from around 150 million in 2010 to nearly 300 million in 2040[2] and the numbers of hip fractures worldwide similarly are expected to rise from 1.66 million in 1990 to 6.26 million annually by 2050.[3] Osteoporotic fractures cost the EU €39 billion annually[4] and are associated with substantial morbidity and excess mortality.[5] Frail older persons represent a particularly high risk group, who may require institutional care in residential or nursing homes. This population are known to be at greater risk of fracture than comparable free-living individuals for a variety of reasons, including generally impaired musculoskeletal health resulting in decreased mobility, increased falls risk and lower BMD, with the adjunctive effects of comorbidities such as dementia and cardiovascular disease.[5, 6]

Whilst this institutionalized population is known to be at high fracture risk, there is little consensus about how best to risk stratify individuals within such an environment. Ihama et al., presented in this issue of European Geriatric Medicine, undertook a study of available fracture risk calculators in a nursing home setting in order to ascertain their predictive value for the outcomes of falls and fractures.[7] As is not unusual, recruitment presented some difficulties in this old frail, often demented population, and it is notable that only 217 (35%) of the 618 residents in 18 homes were enrolled and that of these 217 enrolled, 70% did not have mental capacity and so consent was obtained from a relative. This of course has implications for the ascertainment of self-reported clinical risk factors and indeed for incident events which were reported by the care home at 12 months. Importantly there were 325 incident falls amongst the enrolled participants but only 10 fractures. Thus, the statistical power available to investigate the predictive value of the three risk calculators, together with that for the individual clinical risk factors, is modest. Unsurprisingly, the authors did not find that either FRAX, QFracture or the GARVAN calculator were associated with incident falls or fracture events; indeed only BMI and timed up and go test (TUGT) appeared to have predictive value, although these findings must be appreciated in the context of the large number of statistical tests undertaken. An intriguing further question is whether inclusion of age and sex in the models with BMI or TUGT would alter the associations, an important point in the comparison with the three risk calculators since they all include these two attributes as input variables.

The paper from Ihama et al.[7] thus raises important questions about the general approach to risk stratification in frail older people. The issue ultimately is whether we simply take the view that this is a group at very high risk of fracture (the subsequent question then being whether other factors influence the decision to treat with antiosteoporosis medications) or whether we attempt to stratify within this high-risk population. Whilst traditional risk factors for fracture are still highly relevant here, there are other considerations more specific to the nursing home setting, for example greater mobility may be a risk factor for fracture, since there may be a greater risk of falling in this state compared with being bedbound.[8] There may be clear difficulties with the use of oral medications in the setting of dementia, with parenteral therapies such as subcutaneous denosumab providing a potential solution. An important consideration here is the competing hazard of mortality. Frail older nursing home populations typically have a median survival around 2 years.[9] Indeed the mortality rate amongst those enrolled to the present study was 25% over 12 months, and these were likely to be the fitter individuals amongst the overall population. There is evidence for a benefit of antiresorptive therapies on vertebral fracture risk at 12 months,[10] and over recent years increasing evidence of greater magnitude, and more rapid, effects of anabolic agents, demonstrated head-to-head for risedronate versus teriparatide[11] and alendronate versus romosozumab.[12] Indeed the International Osteoporosis Foundation and the European Society for the Clinical and Economic Aspects of Osteoporosis, Osteoarthritis and Musculoskeletal Diseases have recently released guidance on the targeting of specific therapies by risk level, recommending that those at highest fracture risk should receive anabolic therapies, which will most rapidly reduce their risk.[13] Clearly in the setting of care towards the end-of-life, the likelihood of an event happening relative to the chance of dying from another cause must be considered, together with the likelihood of mortality resulting from a fracture.[5] These are difficult considerations in general, and particularly in the specific, and are driven as much by factors such as philosophy and religion as by empirical science.

The issue of fracture versus mortality risk is relevant to the methodology employed in the three risk calculators tested by Ihama et al. QFracture and Garvan calculators are country specific (UK and Australia respectively), generated in single cohorts, and yield an estimate of cumulative fracture risk over a particular time horizon.[14-16] In contrast, FRAX® was derived through meta-analyses of prospective cohort studies from Europe, North America, Asia and Australia including nearly 45,000 individuals, has subsequently been validated in a similar number of patients in other cohorts, and integrates the risk of fracture with risk of death to generate a probability of fracture over a 10-year period.[17, 16] It is important to note therefore that the metric of risk generated by FRAX differs from that from QFracture and Garvan calculators and that they cannot be used interchangeably. The metric used is particularly relevant when considering frail older persons, who will have a very high fracture risk (in terms of predicted cumulative incidence) but also a high risk of dying over the time period considered. The QFracture solution to this point is to allow calculation of fracture risk over any time period from 1 to 10 years.[14] The Garvan calculator allows calculation over a 5- or 10-year time horizon.[15] In using this approach, the assessing physician must make an assumption regarding likely survival time. The problem then is that there is no consensus on what level of cumulative fracture risk constitutes the threshold for antiosteoporosis medication use by different time horizons. One answer might be to multiply up the risk assessed over a shorter time horizon to the equivalent over 10 years. Unfortunately in the case of QFracture, whilst this would allow assessment over a globally established time horizon, there are still no nationally agreed thresholds for intervention on the basis of this particular risk calculator.[18] The question then is whether FRAX probability could or should be subdivided. The critical difference here is that the metric from FRAX, as described above, integrates risk of fracture and risk of death. Thus, for the same cumulative fracture risk, the probability of a fracture over 10 years will be lower if survival is expected to be 5 years compared with 10 years. This approach means that the same metric can be used at all ages as it accommodates the increased risk of death during the time horizon at older ages. Indeed in the “younger old” population, subdivision of the 10-year probability into smaller time periods, for example deriving a 2-year time horizon by dividing the 10-year period by five, gives broadly proportionate answers.[19] However in the oldest old, there can be substantial deviation from 2 years being 1/5 of the 10-year probability because a fracture cannot occur after the time of predicted death. This problem is even more apparent when considering the effect of a very recent prior fracture on FRAX probability.[20]

Taken as a whole, we are still left with the question of whether to consider the institutionalised oldest old as a group generally at high risk of fracture (and then aim to consider the majority for treatment) or whether to attempt to individually risk assess within this population. Given the current global standards for fracture risk assessment, and the linkage of FRAX, but not other risk calculators, to agreed risk thresholds in many set settings internationally,[21-24, 13] together with the accommodation for competing mortality in the risk model, use of FRAX would seem a reasonable way forward if this were desired. However, its performance characteristics specifically within this population remain to be assessed. For the moment then, particularly given the woefully low proportion of frail older persons, not to mention of the high risk population as a whole, who are appropriately assessed and treated for osteoporosis,[25, 26] a pragmatic approach of aiming to treat, unless there are reasons not to, may well be the order of the day.

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