with longer follow-up and are likely to be largely due to changes in behaviour caused by preclinical dementia.

**OP51**

**PROJECTING THE INCIDENCE AND PREVALENCE OF POST-STROKE COGNITIVE IMPAIRMENT AND DEMENTIA IN THE IRISH POPULATION AGED 40+ YEARS FROM 2015–2025**

1E Sexton*, 2NA Donnelly, 3N Merriman, 4M Guzman-Castillo, 5P Bandonz, 2MA Wren, 1A Hickey, 5M O'Flaherty, 1K Bennett. 1Division of Population Health, RCSI, Dublin 2, Ireland; 2Social Research, ESRI, Dublin 2, Ireland; 3Dept of Public Health and Policy, University of Liverpool, Liverpool, UK

**Background** Post-stroke cognitive impairment (PSCI) is a common consequence of stroke, leading to reduced quality of life and increased care needs. However, rehabilitation services for this condition in Ireland are very limited. The aim was to apply estimates of PSCI incidence to the Irish population and project the number with PSCI in the population in 2025.

**Methods** We developed a deterministic Markov model to estimate future incidence of PSCI in the population aged 40–89 years living in Ireland up to 2025. Population data, estimates and projections to 2025 were obtained from the Irish Central Statistics Office. Data from the Irish Longitudinal Study on Ageing were used to estimate age and sex specific stroke prevalence in 2014. Age and sex specific stroke incidence was estimated using 2015 public hospital discharge data (n=6,155). Transition probabilities across six health states defined by cognitive impairment, physical disability, dementia and death were estimated using data from stroke survivors in the English Longitudinal Study on Ageing (n=490) (2002–2011). Published data from the South London Stroke Register were used to estimate annual stroke recurrence.

**Results** The Irish population aged 40–89 years in 2015–2025 (n=2.7m) is projected to have a cumulative incidence of stroke of approximately 2.3% by 2025 (n=63,100). Of these incident strokes, approximately 22.5% are estimated to have died due to stroke (n=14,200), and 23.8% to have died of another cause (n=15,000) by 2025. Of the survivors in 2025 (n=30,600), approximately 50.9% are predicted to have cognitive impairment without dementia (n=15,500), and 19.4% to have dementia (n=5,900). The total number of stroke survivors is projected to increase from 26700 in 2015 to 41400 in 2025, equivalent to a 55% increase in numbers, and the number with post-stroke dementia is projected to more than double from 3900 in 2015 to 8700 in 2025.

**Discussion** In 2025, over two thirds of Irish people who have survived a stroke in the preceding 10 years will have cognitive impairment. The number of people with post-stroke dementia is set to double between 2015 and 2025. The model is limited by its deterministic nature, and the assumption that age-specific disease incidence will remain stable. The model will be further developed to include a probabilistic sensitivity analysis, to model alternative scenarios for trends in disease incidence, and to extend the projections to 2035. The model will also be used in an economic evaluation of alternative strategies for stroke management, including cognitive rehabilitation.
simulation scenarios to explore the relative impact of different intervention approaches across the dementia population.

Background Interest in loneliness and social isolation as risk factors for premature mortality has recently gained increased attention in both the research literature and public discourses. This has resulted in the established of loneliness taskforces in both Ireland and the United Kingdom. Both loneliness and social isolation have previously been linked to a host of adverse health outcomes, including cardiovascular disease, depression, reduced immune and cognitive functioning, and mortality. Loneliness is most often conceptualised as the subjective assessment of an individual’s social relationships while social isolation is the objective quantitative measure of social contacts. Loneliness and social isolation that have been shown to be distinct yet synergistic constructs.

This study examined the association between loneliness, social isolation, and all-cause and cardiovascular mortality in Ireland and also tested the hypothesis that loneliness provides a mechanistic pathway that explains the association between social network size and mortality. This is the first study in Ireland to use linked survey-health assessment-mortality data, combining rich individual-level data from the Irish Longitudinal Study on Ageing (TILDA) with official death certificate data provided by the official mortality register.

Methods The sample included 6,800 participants, including 654 decedents (with 199 deaths due to cardiovascular disease) who had participated in the first round of TILDA. Underlying and contributory causes of death were coded using the International Classification of Diseases, Injuries and Causes of Death (ICD-10). Loneliness was measured using the five-item Social Network Index. Cox proportional hazards ratios were computed for other factors, the most lonely (HR 1.39 95% CI: 1.14–1.70, p<0.01) was more than twice as likely to die as the least lonely. All results were adjusted for age, sex, total income, clustering of individual-level data from the Irish Longitudinal Study on Ageing (TILDA) with official death certificate data provided by the official mortality register.

Results Mean (SD) age at baseline was 74.1 (2.9) years. Mean annual percentage declines for walking speed and grip strength were 2.1% and 1.5% respectively; declines were smaller for hip BMD (0.6%) and lean mass (0.5%). Trajectories from LMEM (applied to raw data) for grip strength, walking speed, lean mass, and hip bone mineral density (BMD) were explored among 3075 men and women from the Health, Aging and Body Composition Study; each measure was assessed at least 5 times during a median 9 year follow-up period. The following techniques were implemented: linear mixed effects models (LMMEM) (applied to raw data and age-specific z-scores from generalised additive models for location, scale and shape); growth mixture models (GMM); and latent class trajectory models (LCTM). LMEM use random effects to capture inter-individual variation in level and change around a population-average trajectory; GMM extend LMEM by identifying clusters of individuals with similar trajectories and deriving cluster-specific average trajectories; and LCTM are simplified GMM with no random effects, assuming all individuals in a cluster have the same trajectory.

Conclusion LMEM enable a more comprehensive analysis of change compared to methods using data from only two time-points. However, inter-individual differences in rates of change regarding musculoskeletal parameters in this age group and duration of follow-up may be too small to be identified using more complex techniques such as GMM or LCTM.