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Microsimulation model for the health economic evaluation of osteoporosis interventions: study protocol

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ABSTRACT

Introduction Osteoporosis is a systemic skeletal disease that is characterised by reduced bone strength and increased fracture risk. Osteoporosis-related fractures impose enormous disease and economic burden to the society. Although many treatments and health interventions are proven effective to prevent fractures, health economic evaluation adds evidence to their economic merits. Computer simulation modelling is a useful approach to extrapolate clinical and economic outcomes from clinical trials and it is increasingly used in health economic evaluation. Many osteoporosis health economic models have been developed in the past decades; however, they are limited to academic use and there are no publicly accessible health economic models of osteoporosis.

Methods and analysis We will develop the Australian osteoporosis health economic model based on our previously published microsimulation model of osteoporosis in the Chinese population. The development of the model will follow the recommendations for the conduct of economic evaluations in osteoporosis by the European Society for Clinical and Economic Aspects of Osteoporosis, Osteoarthritis and Musculoskeletal Diseases and the US branch of the International Osteoporosis Foundation. The model will be a state-transition semi-Markov model with memory. Clinical parameters in the model will be mainly obtained from the Dubbo Osteoporosis Epidemiology Study and the health economic parameters will be collected from the Australian arm of the International Costs and Utilities Related to Osteoporotic Fractures Study. Model transparency and validates will be tested using the recommendations from Good Research Practices in Modelling Task Forces. The model will be used in economic evaluations of osteoporosis interventions including pharmaceutical treatments and primary care interventions. A user-friendly graphical user interface will be developed, which will connect the user to the calculation engine and the results will be generated. The user interface will facilitate the use of our model by people in different sectors.

Ethics and dissemination No ethical approval is needed for this study. Results of the model validation and future economic evaluation studies will be submitted to journals. The user interface of the health economic model will be publicly available online accompanied with a user manual.

Strengths and limitations of this study

We will develop a state-transition microsimulation for health economic evaluation of osteoporosis interventions.

The model will be based on the Australian setting and will be adapted and adaptable to other settings.

The model will be publicly available with a web interface to allow access for future users to evaluate new osteoporosis interventions of interest in their own settings.

The model will be developed following the consensus statement endorsed by the Scientific Advisory Board on Economic Aspects of Osteoporosis, Osteoarthritis and Musculoskeletal Diseases, the Committee of Scientific Advisors and the Committee of National Societies of International Osteoporosis Foundation, the US National Osteoporosis Foundation and Osteoporosis Canada.

INTRODUCTION

Osteoporosis is a common disease that is usually silent until a fracture occurs. Fracture is a global health concern particularly in women. Australian women aged 60 years were estimated to have a residual lifetime fracture risk of 44% and the risk for Chinese women was approximately 33%.1,2 Despite a lower residual lifetime risk of having a fracture in men, the overall mortality HR in men after a fracture was higher than women.3 For those who survived fractures, quality of life (QoL) deteriorated dramatically up to 6 months after fracture and the loss of QoL persisted in the remaining lifetime.4 With an accelerated ageing global population, the number of fractures will increase in the coming decades.5,6

Bone strength is mainly reflected by bone mineral density (BMD). Individuals with reduced BMD are at the greatest risk of osteoporotic fracture. The assessment of fracture risk is largely based on clinical risk factors with and without BMD.7 With the...
predicted fracture risk, treatment decisions can be made following primary or secondary fracture prevention.8,9

Numerous therapeutic regimes, including pharmaceutical interventions, have been developed to treat osteoporotic patients by improving their BMD. While treatment of osteoporosis has proven effective in preventing future fractures, the associated cost must be considered against the costs of not preventing fractures. In 2011/2012, the Australian government spent over 179 million Australian dollars in pharmaceuticals for osteoporosis management.10 Similar costs of osteoporosis medications have been noted elsewhere.6,11–12 With a range of osteoporosis treatments available in the community, some governments employ health economic evaluation in treatment reimbursement policy making, such as the Pharmaceutical Benefits Advisory Committee (PBAC) in Australia and the National Institute for Health and Care Excellence (NICE) in the UK. These government agencies conduct health economic evaluations of osteoporosis treatments to select those that represent good value for money.

Health economic evaluation of osteoporosis interventions is commonly conducted using disease modelling approaches,13 such as Markov cohort models and individual-level state-transition models.14–16 Health economic modelling using different sources of information is able to capture long-term cost and health benefits of osteoporosis interventions. Health economic analyses of osteoporosis interventions may be limited due to the validity, transparency and accessibility of health economic models that are used in the analyses. Making health economic models transparent via thorough documentation and making them available for public use are potential solutions to minimise these limitations; moreover, these solutions provide opportunities to update existing models with newly available epidemiological and health economic data as they become available.17 To date, there is no publicly accessible health economic model of osteoporosis.

OBJECTIVE

We aim to develop a publicly available health economic model to evaluate the cost-effectiveness of osteoporosis interventions. The model will be based on the Australian setting and will be adapted and adaptable to other settings. Moreover, we will develop a web interface to allow access for future users to evaluate new osteoporosis interventions of interest in their own settings.

METHODS AND DESIGN

Model structure and parameters

We have published a microsimulation model of osteoporosis in the Chinese population and will develop the Australian model using the existing model structure.18 The model will incorporate Australian fracture risk equations for estimating absolute risk of osteoporosis fractures based on patients’ characteristics (eg, age, sex, family history, BMD, history of fractures and falls as well as comorbidities), and time-varying factors such as exposure to primary and/or secondary fracture prevention therapy and patients’ accumulated history of fractures. These equations will form the basis of the simulation model for estimating outcomes for Australian patients with osteoporosis.

The development of our osteoporosis health economic model will follow the recently published modelling guideline in osteoporosis.19 Simulations will be based on a state-transition individual patient simulation model (figure 1). Simulated patients in the model are either alive or dead based on annual transitions. Outcomes will be analysed over patients’ lifetimes but time horizons may be varied according to the study of interest. Simulated patients will transit in the model until they are absorbed in the ‘death’ state.

Fragility fracture is a key outcome of osteoporosis and can occur in hip (proximal femur), vertebrae (spine), wrist (distal radius) and other non-hip non-vertebral sites such as humerus, pelvis, ribs and shoulder. Simulated patients can have multiple fractures at different fracture sites. Fracture probabilities will be simulated from one of three risk engines. First, for countries that report fracture incidence rates, fracture probabilities will be calculated from incidence rates using the following equation:

\[ p = 1 - \exp(-r) \]

where \( p \) is the probability and \( r \) is the annual fracture incidence rate by sex, fracture site and BMD level. In the model for the Australian population, annual fracture incidence rates by sex, fracture site and BMD level will be obtained from the Dubbo Osteoporosis Epidemiology Study.20 For countries without reported fracture incidence rates, users will be able to choose the formula for calculation of annual fracture probabilities. Ten-year risk for hip and all fragility fractures are predicted by the FRAX or Garvan Fracture Risk Calculator (GARVAN-FRC).21–24 The annual probability of having a fracture will be calculated as:

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**Figure 1** Structure of the osteoporosis state-transition model. Simulated patients transit in the model following the arrow direction. Simulation is concluded when all simulated patients transit to the ‘death’ state. Reprinted by permission from Springer Nature Customer Service Centre GmbH: Springer Nature.18
\[ p = 1 - \exp \left( \frac{\ln(1-P)}{t} \right) \]

where \( P \) is the cumulative risk of having a fracture over a period of \( t \) years. For example, for a given set of clinical risk factors, a group of patients is estimated to have a 5\% 10-year probability of having a hip fracture. In this scenario, the annual probability of having a hip fracture is calculated as 0.51\%.

Simulated patients could stay without any fracture for the entire life (represented as ‘no history of fracture’), sustain a fracture (represented as ‘fractured’), stay in postfracture state (represented as ‘postfracture’) or sustain another fracture (represented as ‘fractured’). In the case of death, Australian life tables combined with fracture site-specific relative risks adjusted by time since fracture will be used to calculate the probability of death during the course of the simulation.3

The model will primarily take the societal perspective, which is a broader perspective accounting for all stakeholders. In the Australian model, we will use data from the Australian arm of the International Costs and Utilities Related to Osteoporotic Fractures Study (AusICUROS) to calculate the costs of direct health and non-healthcare utilisation.10 Specifically, direct healthcare utilisation included hospitalisation, ambulance, imaging, medical services, pharmaceuticals and supplements, non-admitted, subacute/rehabilitation and community-based services including GP and physiotherapy services. Direct non-healthcare utilisation included residential care, meals on wheels and other community services.10 Relevant unit costs of the above healthcare utilisation will be taken from Pharmaceutical Benefit Scheme (PBS), Medical Benefit Scheme (MBS) and other government reporting system such as the Australian-refined diagnosis-related groups (AR-DRGs). In addition, we will include indirect costs of productivity loss. While societal perspective will be used primarily, the cost data will be categorised to be adapted for a different perspective for future use. For example, when evaluating the cost-effectiveness of an osteoporosis medication for listing on the PBS, only costs borne by the Australian government will be included.

While the model can estimate the life expectancy, it will primarily use quality-adjusted life years (QALYs) to quantify health outcomes, since this is the most commonly used metric in health economic evaluations. The QALY, which adjusts life expectancy by the degree of morbidity, is usually measured on a health state utility (HSU) scale where zero represents death and one represents full health in each year of life. HSUs for each level of disease severity will be incorporated from the AusICUROS and an international meta-analysis of HSU changes before and after osteoporosis-related fractures.10 25

The overview of the osteoporosis model algorithm is illustrated in figure 2.
with a prior fracture was higher and varied by sex, fracture site and BMD level.\textsuperscript{27} Likewise, mortality was also higher after fracture.\textsuperscript{28} Therefore, we will use an individual-level state-transition model to incorporate the memory of events occurring for simulated patients in the model. Tracker variables will be used to record events of interest during simulation. In our model, we will define the following tracker variables to represent the heterogeneity of simulated patients: ‘site of each fracture’, ‘number of fracture by fracture site’, ‘time of the first fracture’, ‘time since the last fracture’ and ‘time after treatment onset’.

**Management with uncertainties and reporting uncertain analyses**

Four types of uncertainties are commonly considered in health economic modelling studies, namely stochastic uncertainty, parameter uncertainty, heterogeneity and structural uncertainty.\textsuperscript{28} These uncertainties will be dealt with during the development of the model and analyses. For stochastic uncertainty, sufficient number of first-order Monte Carlo simulations will be conducted. For parameter uncertainties, CI of parameters along with point estimates will be included in the model. For instance, gamma or log normal distribution for cost data, log normal distribution for relative risks or hazard ratios will be used.\textsuperscript{29} Both deterministic and probabilistic sensitivity analyses will be conducted to report uncertainties. One-way sensitivity analysis will be performed to evaluate how results change by changing the value of individual parameters and the results will be presented in tornado diagrams. Probabilistic sensitivity analysis (PSA) will be conducted by simultaneous sampling of all parameters that are defined by distributions. Cost-effectiveness acceptability curves (CEAC) and distribution of net monetary or net health benefits will be presented to illustrate the cost-effectiveness decisions given a range of willingness-to-pay (WTP) thresholds. Handling uncertainties and reporting uncertainty analyses will be conducted in line with the Good Research Practices in Modelling Task Force-6.\textsuperscript{28}

**Model transparency and validation**

Cost-effectiveness of health interventions from modelling studies informs policy decision makers when they prioritise funding for scarce healthcare resources. Therefore, the validity and transparency of a health economic model are keys in assisting policy decision making. Transparency of health economic models is subject to how well the model is documented including its structure, parameters, equations and key assumptions used in the model. Validity refers to how well the model represents the real world.\textsuperscript{30} To ensure our model will be developed following recommendations from the Good Research Practices in Modelling Task Force-7,\textsuperscript{30} we will make the non-technical documentation available following the example of the previous work in diabetes from our team.\textsuperscript{31} There are five aspects of validity, namely face validity, internal and external validities, cross validity and predictive validity.\textsuperscript{30} In our study, we will primarily deal with face, internal and external validities of our osteoporosis model.

For face validity, the clinicians in our team (JAE, TW, JRC and AP) will evaluate whether the model presents the osteoporosis clinical pathway and the current osteoporosis management algorithms in Australia. For internal and external validity, we will conduct goodness-of-fit analyses on the model results against values used in developing the model (ie, internal validity) and that from literature (external validity). In particular, we will compare the following values: annual fracture rates by age and sex; life expectancies by age and sex; residual lifetime fracture risks by age, sex and BMD level. Goodness-of-fit analysis will be conducted using linear regression. The slope of the regression line along with the R-squared of the regression line and the root mean square error (RMSE) will be used to assess the goodness-of-fit of the model predictions and that from the literature.\textsuperscript{32} In addition, the Bland-Altman statistics will be reported to evaluate the agreement between model results and real-world observed results.\textsuperscript{33}

**Patient and public involvement**

Patients and public will not be involved in this study.

**APPLICATION OF THE MODEL**

The first focus of the model application will be to quantify the cost-effectiveness of all first-line osteoporosis secondary prevention fracture medications currently reimbursed by the PBS in order to identify the most cost-effective medications and to further identify currently reimbursed medications that are not cost-effective indicating the need for disinvestment in these medications.\textsuperscript{34} The second focus will be to identify the most cost-effective osteoporosis screening and treatment strategy in Australia, as has been performed in China.\textsuperscript{35}

The model will be used to evaluate the costs and incremental cost-effectiveness of alternative new interventions for primary and/or secondary fracture prevention. Modelling is usually required in economic evaluations of therapies for osteoporosis fracture prevention, as most trials end before long-term outcomes are known. Evaluating alternative management strategies will comprise the following steps:

i. Development of the scenario to be simulated. This involves choosing an intervention or management guideline for evaluation. Evidence must be available on its effect on fracture risks and synthesis undertaken to provide inputs for the simulation model. In the case of this analysis, the intervention to be evaluated may include denosumab, bisphosphonates and other commonly used medical treatments presently available to confirm cost-effectiveness, or to identify for disinvestment if no longer found to be cost-effective; and any new medications not yet reimbursed under the MBS.

ii. Simulating long-term outcomes and costs. This will involve carrying out two simulations: in the first, the
target population is assumed to receive standard of care and in the second, the target population is assumed to receive standard of care plus the alternative treatment;

iii. Estimation of the incremental cost-effectiveness ratio. Results from the simulation model will be combined with other information (eg, the cost of treatment) in order to undertake an incremental cost-effectiveness analysis in which the net cost (ie, treatment costs minus savings due to fractures avoided) is divided by the net effectiveness (QALYs gained).

To ensure that the work undertaken has relevance to policy, we intend to hold a workshop on osteoporosis modelling in Australia during the course of this project. The aim of the workshop will be to obtain the opinions of key stakeholders (eg, officers from the Department of Health & Aging, representatives from Osteoporosis Australia (patients’ perspectives), clinicians from the Royal Australian College of General Practitioners and representatives from pharmaceutical companies) to assist in prioritising the work programme.

Model interface and availability

We will capitalise on our team member AJP’s extensive experience in his previous CORE Diabetes Model to ensure similar successful translation of the model into general use. A user-friendly graphical user interface will be developed, which will connect the user to the calculation engine and the results are generated. Input data, such as cohort characteristics, key cost and utility parameters, and key treatment characteristics (costs and effects) will be accessible and editable by the user. They will be stored in a structured query language (SQL) database. Results generated by the model will also be written to and stored in a database for easy future access by the user. A user instruction manual and help files will be developed.

Like the CORE Diabetes Model, the osteoporosis model will be made available to university or other public sector research groups at no cost, while those seeking to use it for commercial use (eg, such as pharmaceutical companies preparing a regulatory submission) will pay a license fee that will support maintenance and further development of the web-based model.

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Contributors

LS and AJP contributed to the initial idea of the study. LS, JE, TW, KMS, JRC, TN and AJP contributed to the design of the work. LS drafted the manuscript. LS, JAE, TW, KMS, JRC, TVN and AJP contributed critical review for the intellectual content and approved the final version of the manuscript submitted for publication.

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Competing interests

None declared.

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Not required.

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