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CHI study: protocol for an observational cohort study on ageing and mental health in community-dwelling older adults

Rachael Zhi Yi Lee, Junhong Yu, Iris Rawtaker, Patrick Finbarr Allen, Zhiming Bao, Lei Feng, Qiushi Feng, Jeong Kyu Lee, Chin Tat Lim, Lieng Hsi Ling, Leng Leng Thang, Thet Naing, D Y Wang, Kai Zhen Yap, EH Kua, Rathi Mahendran

ABSTRACT

Introduction Ageing is associated with a multitude of healthcare issues including dementia, depression, frailty, morbidity associated with chronic disease and high healthcare utilisation. With Singapore’s population projected to age significantly over the next two decades, it has become increasingly important to understand the disease burden and etiological process among older adults. The Community Health and Intergenerational study aims to holistically examine ageing in place by investigating the resilience and vulnerability factors of the ageing process in the biological, psychological and social domains within the environment.

Methods and analysis Using a cohort multiple randomised controlled trial design, comprehensive health profiles of community-dwelling older adults will be collected. The objective is to recruit 1000 participants (aged 60–99 years) living in the western region of Singapore within a period of years (2018–2020). Assessments include basic sociodemographic, physical health and function (cardiac, oral and blood profiles and visual function), cognitive functioning, daily functioning, physical fitness, emotional state, free-flowing speech, sleep quality, social connectedness, caregiver burden, intergenerational communication, quality of life, life satisfaction, attitudes to ageing and gratitude and compassion. Results from the cohort will enable future studies to identify at-risk groups and develop interventions to improve the physical and mental health and quality of life of older adults.

Ethics and dissemination Approval of the cohort study by the National University of Singapore Institutional Review Board (NUS-IRB Reference code: H-17-047) was obtained on 12 October 2017. Written consent will be obtained from all participants. Findings from the cohort study will be disseminated by publication of peer-reviewed manuscripts, presentations at scientific meetings and conferences with local stakeholders.

INTRODUCTION

Background

The WHO estimates that the global population of older adults aged 60 years and above will rise from 900 million in 2015 to 2 billion in 2050. In Singapore, the proportion of residents aged above 65 years nearly doubled from 8.8% in 2009 to 14.4% in 2019 and is projected to be 25% by 2050. This poses a challenge as ageing is associated with a plethora of healthcare issues and high healthcare utilisation. Over the years, researchers have conducted nation-wide studies in Singapore to understand age-related diseases and modifiable factors to promote healthy ageing. Previous research has adopted a multidimensional framework (eg, WHO’s definition of health) to better understand the ageing process and healthcare-related needs. Using a similar framework, the Community Health and Intergenerational (CHI) study adopts Engel’s biopsychosocial model of health and disease to holistically examine ageing in place by...
collecting comprehensive health profiles of older adults in Singapore.

To date, cohort studies are shifting towards using holistic frameworks to observe ageing and health in the community. The Healthy Older People Everyday study (n=1051) is one such study that sought to assess physical and mental health among community-dwelling older adults (aged ≥65 years) through basic health screening as well as a health survey.9 Although the study used objective screening tools (eg, mini-mental state exam and physical fitness tests), it comprised mostly of self-reported measurement. The authors also suggested the need for more robust and comprehensive tools to be considered such as the Geriatric Depression Scale. Although other larger age-related cohort studies such as the Australian Imaging Biomarkers and Lifestyle study10 and Alzheimer’s Disease Neuroimaging Initiative11 are notable studies that have collected a wide range of measures (eg, clinical, cognitive, neuroimaging, lifestyle and genetic data), nonetheless the focus was largely on the treatment and progression of Alzheimer’s disease (AD). Further research is needed to assess other health-related determinants of older adults in the healthy ageing spectrum such as oral health assessments, cardiovascular investigations, speech analysis and olfactory measures. The Well-being of the Singapore Elderly study (n=2565) was another comprehensive study that included face-to-face interviews and physical examination; however, it lacked laboratory measurements and did not assess cardiovascular and other physical health risks.12,13 Research using objective measures to determine physical (eg, blood markers or echocardiography) and cognitive (eg, neurocognitive assessments) health status is needed to complement self-reported data. In terms of oral health, there appears to be an association between dental disease, tooth loss and onset of frailty.14 It is plausible that this may be mediated through triggering of inflammatory processes by pathogens from periodontal tissues, but there is a lack of longitudinal data to confirm this hypothesis.

There have been a few cross-sectional cohort studies carried out on older adults in the local context; these include the Singapore Longitudinal Ageing Study (SLAS) and the Diet and Healthy Ageing (DaHA) cohort. Conceived in 2003, the SLAS (n=6183) aimed to provide a community-based cohort of older adults (aged 55 years and above) for subsequent clinical-based interventions.7 Results from the ongoing study found new prevalence rates15 and associations.6,16 Participants from SLAS were also identified to join subsequent intervention studies such as a computer training randomised controlled trial (RCT) that was found to improve cognitive functioning.17 On the other hand, the DaHA (cohort) study placed emphasis on dietary factors and its association with healthy ageing and reduced risk of age-related medical conditions.5 For instance, the bioactive compounds found in mushroom and long-term tea consumption were associated with delay in cognitive impairment and reduced depression/anxiety symptoms, respectively.5,18 The cohort study later invited suitable subjects to participate in subsequent non-pharmacological RCTs that aimed to improve cognitive and psychological health—art therapy and music reminiscence activity,19 mindfulness awareness programme20 and horticultural therapy.21

Although the aforementioned studies on older adults documented valuable findings, some of them mainly focused on the treatment and progression of AD, while most studies did not incorporate other important measures of health such as detailed oral health examination, cardiovascular assessments and biomarkers, olfactory measures or speech analysis. More observational studies using in-depth and culturally relevant assessments of older adults in the healthy ageing spectrum are needed. This calls for greater integration of health, psychosocial and environmental resources through close collaborations among clinicians, researchers and community partners. Thus, the CHI study aims to holistically investigate factors associated with healthy ageing in a community setting using a broad range of health-related measures.

Research aims
The primary goal is to examine the health profiles of older adults and form meaningful associations based on the following:

a. Biological factors such as the physical health condition (eg, cardiac, oral, blood profiles, vital signs and visual function), physical fitness/function, medical history and medication use, and nutritional status.

b. Psychological factors such as the cognition, emotional state (anxiety and depression symptoms), sleep quality, attitudes, values, satisfaction with life and quality of life.

c. Social factors such as social support, intergenerational relationships and the impact on family members.

Second, this study also acts as a recruitment platform for future interventional studies (eg, feasibility or full-scale trial) to identify at-risk groups or normal ageing participants. The cohort data will enable the development and evaluation of pharmacological and psychosocial interventions targeted at improving health outcomes for older adults. Specifically, data will be used to identify at-risk groups such as (but not limited to) older adults with subsyndromal depression or anxiety, mild cognitive impairment, medical conditions (eg, hyperlipidemia, diabetes, hypertension), at-risk of cardiovascular diseases, oral diseases, speech impairment or sleep apnea. Other future substudies will also explore culturally relevant psychosocial factors related to healthy ageing such as intergenerational communication, attitudes to ageing, social networks, satisfaction with life and many more.

METHODS AND ANALYSIS

Study design
The CHI study adopts a cohort multiple RCT (cmRCT) design, whereby the cohort provides capacity for multiple RCTs over time.22 Using a cmRCT design may increase...
efficiency in trial recruitment and potentially lower attri-
Bution rates.23 Hence, the cmRCT design is adopted to
determine biopsychosocial factors involved in the ageing
process and subsequently introduce interventions that
can mitigate ageing-related issues such as cognitive and
psychological health, diet, medication adherence, speech
impairments, oral hygiene. The ongoing CHI study has
started data collection on 1 February 2018. It will be
conducted in two phases. Phase I comprises of a cross-
sectional cohort study (baseline) detailed in this paper.
Participants enrolling for the CHI cohort will be given
the option of allowing their data to be used for analyses
and in identifying them for future research interventions
(i.e., Phase II) or for comparison purposes for interven-
tion trials. Details of the Phase II interventions will be
published in full manuscripts separately.

Study sample
This 3-year cohort study targets to recruit 1000 community-
dwelling older adults in Singapore. Participants aged
between 60 and 99 years of any gender and ethnic group
will be eligible for the study. Illiterate participants are
also eligible for the study; however, they will be excused
from the Cambridge Neuropsychological Test Automated
Battery. Assessments will be conducted in languages such
as English, Mandarin, Malay and Chinese dialect (e.g.,
Hokkien), depending on the subject’s preference.

In collaboration with Presbyterian Community Services
(PCS), the main study site will be held at a local seniors’
activity centre, Hannah Seniors Activity Centre (HSAC),
located within the community in the central west district
of Singapore. Confirmation of the sociodemographic
data of over 3000 older adults in the Anak Bukit area (i.e.,
a subzone of the central west district) has been verified
through the Department of Statistics Singapore.24 Data
from PCS also indicated an estimate of 1000 older adults
within a 20-block radius from the activity centre at Toh
Yi. Recruitment is restricted to the central west district
of Singapore.

Patient and public involvement
Participants in this study were not involved in the devel-
lopment of the study design or objectives. The research
design and objectives were developed by the investigators
of this study and underwent review by a board of academic
advisors affiliated to the National University of Singa-
pore Mind-Science Centre. PCS, a community partner,
provided the study site (e.g., quiet rooms in HSAC). In
addition, information about the study procedures and
recruitment process were shared with staff from PCS
prior to data collection.

Procedure
Older adults will be recruited from residences via door-to-
door visits by research nurses and research assistants in
the Toh Yi, Anak Bukit area and other areas within the district
encompassed by a 10 km radius from the HSAC. Eligible
individuals will also be recruited onsite from HSAC,
community centres, resident corners, senior activity
centres and residences within the recruitment area—ad-
vertisement flyers will be made available for visitors to the
respective centres and word of mouth. Interested individu-
als will be invited to HSAC at their convenience. Non-
ambulant individuals who are keen to participate in the
study will have their consent taken in their own homes.
A member of the research team will explain the study in
detail and time will be given for individual to consider
before written consent is given.

Participants will be invited to complete up to six sepa-
rate visits, estimating to a total of 11 hours; they will be
scheduled to complete five visits to HSAC and one to
NUS Cardiovascular Imaging Core Lab at their earliest
convenience. Data will be obtained through semi-strauc-
tured face-to-face interviews (visits 1 and 2), neuropsycho-
logical assessments (visits 3 and 4), biological specimen
collection (blood and dental samples), dental examina-
tion (visit 5) and cardiovascular examination (visit
6), details of which are given in table 1. Non-ambulant
participants will be assessed in their own homes and will
only have to complete visits 1–3. Trained research assis-
tants and certified nurses will conduct visits 1–4 and
blood venepuncture, while certified dentists and medical
sonographers will conduct visits 5 and 6, respectively.
Moreover, experts will provide referral letters to partici-
pants for further follow-up if incidental findings should
arise from assessments such as neuropsychological tests,
depression/anxiety screening, oral health examination,
blood tests and/or cardiovascular examination.

Outcome measures
A wide range of data, spanning across several health
domains, will be collected. Instruments and physical
examinations were introduced by the investigators of this
study that comprises of experts in psychiatry, cardiology,
dentistry, otolaryngology, sociology, pharmacy, family and
population research, linguistics, public health, ortho-
paedics and ophthalmology. These measures have been
validated in the local context.15 25–42 It also comprises
novel scales that will be used to test for validity in this
sample. Table 1 provides an overview of the measures to
be collected.

Data storage and analysis
Participants will be assigned a subject identification
number (SID). Their corresponding data including ques-
tionnaire responses, audio recording, assessment records
and biological samples will be kept anonymous and coded
with the same assigned SID for consistency. All hardcopy
data responses will be checked by two personnel to mini-
mise missing data. Hardcopy coded data collected will be
entered and stored on a standalone computer, and soft-
copy data will be password protected. Furthermore, all
hardcopy data will be stored in a designated secured and
locked space, accessible only to the selected personnel.
In general, demographic variables will be presented
as descriptive summaries such as mean±SD, median,
### Table 1 Outcome measures

<table>
<thead>
<tr>
<th>Variables</th>
<th>Instrument/scale</th>
<th>Visit</th>
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<tbody>
<tr>
<td><strong>Sociodemographic</strong></td>
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<tr>
<td>Age</td>
<td>Age will be measured based on the date of birth stated on the National Registration Identification Card (NRIC) or long-term visit pass.</td>
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<tr>
<td>Sex</td>
<td>Male or female stated on NRIC or visit pass.</td>
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<tr>
<td>Language use</td>
<td>Measured by language of interview, language participant is able to speak and common language spoken at home.</td>
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<tr>
<td>Marital status</td>
<td>Self-report of marital status; single, married, widowed or divorced/separated.</td>
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<tr>
<td>Ethnicity</td>
<td>Ethnicity as recorded in NRIC or self-report; categorised as Chinese, Malay, Indian or others.</td>
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<tr>
<td>Religion</td>
<td>Religion will be classified as Taoism/Buddhism, Christianity/Catholicism, Hinduism, Islam or others. Participants will also be asked, ‘How important is your religion to you?’ via a 4-point Likert scale response.</td>
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<tr>
<td>Citizenship</td>
<td>Based on citizenship recorded on NRIC or visit pass, which will be categorised into Singapore citizen, Permanent Resident (PR) or others. For PRs, previous citizenship and year of PR status will be collected.</td>
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<tr>
<td>Education</td>
<td>Measured by years of formal schooling and highest education level, which will be categorised as none, primary, secondary/technical education, preuniversity/polytechnic or university.</td>
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<tr>
<td>Employment status</td>
<td>Determined by self-reported employment status; categorised as retired, housewife, full-time, part-time or self-employed. Participants will also be asked to state their previous and current occupation.</td>
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<tr>
<td>Living arrangement and family support</td>
<td>Questions from previous surveys centred on older adults and their children’s living arrangement, and sources of support and care will be used.</td>
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<tr>
<td>Financial status</td>
<td>Financial status is determined by housing type, current gross personal monthly income, current gross household income, insurance coverage and expenditure on medical expenses per month. Participants will also be asked whether their income/allowances are adequate to cover their monthly expenses, reasons if it is not adequate and if their financial resources are adequate to meet their future needs. Various sources of support will be recorded, such as private savings, borrowing money from relatives/friends, etc.</td>
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<tr>
<td>Spouse demographic</td>
<td>Spouse’s age, ethnicity, education, citizenship and employment will also be collected.</td>
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<tr>
<td>Medical history</td>
<td>The medical history of participants and their family will be collected using the self-reported questionnaire from the Diet and Health Aging study. Medical conditions such as hypertension, stroke, diabetes, hyperlipidaemia, cancer, cataracts, mental health illnesses and many more will be recorded.</td>
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<tr>
<td><strong>Biological factors</strong></td>
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<tr>
<td>Body measures</td>
<td>Blood pressure, pulse rate, height, weight, neck circumference and abdominal girth will be measured. In addition, body mass index (BMI) will be calculated.</td>
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<tr>
<td>Visual function</td>
<td>Visual acuity will be measured using a standardised tumbling E distance vision chart of 3 m, while colour blindness is measured using the Ishihara 38-plate colour test. To measure visual functioning, five items were selected from the original 25-item National Eye Institute Visual Functioning Questionnaire. One item was selected from five subscales on general vision, near vision, distance vision, social function and driving and will be scored on a 3-point Likert scale (good, acceptable or poor).</td>
<td>1</td>
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<tr>
<td>Speech</td>
<td>Participants will be asked to speak freely about their life story and experiences for 15–20 min using a language of their choice. Their speech will be recorded using an audio recorder and they will be instructed to remain anonymous in the recording.</td>
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<tr>
<td>Functional status</td>
<td>Barthel's Index of Activities of Daily Living and Lawton's Instrumental Activities of Daily Living scale (iADL) are common scales used in Singapore to assess older adults' ability to perform basic and complex self-care tasks independently. This study adopts five items (bowels, bladder, grooming, toilet use and feeding) from the 10-item Barthel Index and the original 8-item iADL scale. Both scales are scored on a 3-point ordinal scale (independent, some help required or dependent).</td>
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<tbody>
<tr>
<td>Medication use/adherence</td>
<td>Participants will be asked to bring along any prescribed medication, supplements and/or over-the-counter medication. Name of medication, dosage form, dosing instructions, frequency of use, duration used, purpose and source of medication will be recorded.</td>
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<tr>
<td>Physical fitness</td>
<td>Physical fitness is determined by results from five tasks; handgrip test using a calibrated Jamar dynamometer, modified functional reach test, 30 s chair stand, timed up and go, 6 m fast gait speed test. These tests will assess grip strength, balance, lower extremity strength, functional mobility and gait speed, respectively. In addition, levels of physical activity will be measured by the 4-item international physical activity (short form) questionnaire.</td>
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<tr>
<td>Blood profile*</td>
<td>23.5 mL of blood will be obtained through venepuncture performed by certified nurses. 13.5 mL of blood will be tested for general health markers; alkaline phosphatase, alanine aminotransferase, phosphate, calcium, uric acid, full blood count panel without erythrocyte sedimentation rate, total cholesterol, high-density lipoprotein cholesterol, low-density lipoprotein cholesterol, triglycerides, fasting blood glucose, glycated haemoglobin A1c, free thyroxine (T4), thyroid-stimulating hormone, thyroid peroxidase antibody, intact parathyroid hormone, 25-hydroxyvitamin D, sodium, potassium, chloride, urea, creatinine, estimated glomerular filtration rate. The remaining 10 mL of blood will be used for near-term assays of candidate cardiovascular biomarkers that include (but not limited to) N-terminal-proB-type natriuretic peptide, high-sensitivity cardiac troponin, growth differentiation factor-15 and ST2 protein.</td>
<td>2/3/4/5</td>
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<tr>
<td>Olfactory status</td>
<td>Using a recently developed olfactory test kit, participants will be tasked to smell nine locally developed scents (almond, lemon, orange, pineapple, banana, coconut, rose, cinnamon and mushroom). They will then be asked to identify the scent and rate the intensity and pleasantness of the scent scored on a 5-point Likert scale.</td>
<td>2/3/4/5</td>
</tr>
<tr>
<td>Oral health status</td>
<td>Participants will receive intraoral and extraoral clinical examinations by three calibrated dentists. Similar to previous studies, the oral health examination includes examining and recording of oral mucosa status, periodontal status, tooth ( coronal and root) status and treatment needs, tooth wear, occlusal contacts, and prosthodontic status. In addition, oral samples (dental plaque and saliva) will be collected for DNA extraction and microbiome analyses. Specifically, supragingival plaque will be removed with sterile cotton pellets and dried prior to sampling of subgingival plaque. Full mouth subgingival plaque will be removed using a sterile curette and resuspended in a microcentrifuge tube containing 500 µl of sterile saline. Participants will then be asked to chew a paraffin gum to transfer bacteria from teeth to saliva and subsequently drool into a receptacle.</td>
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</tr>
<tr>
<td>Nutritional status</td>
<td>The widely used Mini Nutritional Assessment—Short Form will be used to assess nutritional status. It consists of six items; appetite, weight loss, mobility, psychological stress or current illness, BMI and neuropsychological problems. Total weighted screening scores range from 0 to 14 points.</td>
<td>5</td>
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<tr>
<td>Cardiovascular status</td>
<td>Six non-invasive cardiovascular procedures will be performed: (1) echocardiography (ultrasound scanning) will be conducted to assess the morphology and function of the heart using a scanning transducer lightly applied to the chest, (2) echocardiography of the carotid and femoral arteries, and modified applanation tonometry at the radial artery of the wrist will also be used to determine carotid intima-media thickness and arterial stiffness properties, (3) skin autofluorescence scanning to detect dermal deposition of advanced glycation end products (AGEs) will be measured by an AGE Reader SU device, which requires participants to place their forearm on the reader, (4) echocardiography of flow-mediated dilation at the brachial artery will be conducted using a 10 MHz linear array probe, steadied by a stereotactic clamping image the brachial artery and position electronic tracking gates at the media-adventitia interface of opposing arterial walls as well as the use of the E20 rapid cuff inflator, to induce reactive hyperaemia by inflation of a pneumatic cuff placed around the participant’s proximal forearm to a pressure of 50mm Hg above the systolic blood pressure for 5 min, (5) ECG will also be performed, so as to record the electrical activity of the participant’s heart at resting state using electrodes with adhesive pads attached to the chest, arms and legs and last (6) ambulatory ECG (Holter) monitoring will be conducted to detect arrhythmias, including atrial fibrillation, assess heart rate variability and heart rate complexity using a portable monitor attached by wires to electrode patches placed on the chest for 24 hours. During the 24 hours monitoring, participants will also be tasked to fill in a diary sheet (ie, type of activities and heart-related symptoms experienced) as accurately as possible. These cardiovascular procedures will adhere to strict local standards, and reports will be reviewed by cardiologists.</td>
<td>6</td>
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<tr>
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<tbody>
<tr>
<td><strong>Psychological factors</strong></td>
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<tr>
<td>Psychiatric symptoms</td>
<td>Depressive and anxiety symptoms are assessed using the 15-item Geriatric Depression Scale, and the 20-item Geriatric Anxiety Inventory, respectively. Both scales have been validated in the local context and shown good psychometric properties in older adult populations. Participants with scores above the local cut-off point that signifies risk of depression and anxiety will undergo an assessment by a psychiatrist and referred for follow-up.</td>
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<tr>
<td>Lifestyle factors</td>
<td>Lifestyle factors are assessed by a previously developed lifestyle questionnaire. In addition, participants will be asked about the type of leisure activity who have participated before and number of hours spent/week participating in; mindfulness, artwork, exercise, social activities, musical activities or others (same as previous work).</td>
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<tr>
<td>Perceived oral health and QoL</td>
<td>The 15-item Oral Health Attitudes Questionnaire will assess attitudes to oral health, while oral health-related quality of life (QoL) will be assessed using the short form Oral Health Impact Profile—14 items. PDQ has also been used to measure SCD in a Singapore cohort.</td>
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<tr>
<td>Subjective cognitive decline (SCD)</td>
<td>The 20-item Perceived Deficits Questionnaire (PDQ) is part of the Multiple Sclerosis Quality of Life Inventory that assesses self-perceived cognitive decline. PDQ has also been used to measure SCD in a Singapore cohort.</td>
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<tr>
<td>Cognitive functioning</td>
<td>Measures of cognitive functioning include: (1) A locally modified and validated 30-point Mini-Mental State Examination (MMSE) with stratified education and ethnic cut-offs to assess global cognitive function; (2) the 5-point Clinical Dementia Rating (CDR); (3) assessing verbal and learning memory (Rey Auditory Verbal Learning Test) attention and working memory (Digit Span Forward and Backward Task), divided attention and sequencing (Colour Trails Test), visual-spatial abilities (Wechsler's Block Design) and verbal fluency (Semantic Verbal Fluency—Animals); (4) eight computerised language-independent cognitive tests of the Cambridge Neuropsychological Test Automated Battery (Cambridge, UK) will be administered: Motor Screening Task (sensorimotor), Paired Associates Learning—12 patterns (visual memory and learning), Verbal Recognition Memory—immediate and delayed recall (verbal memory), Stockings of Cambridge (spatial planning), Emotion Recognition Test—short (social cognition), Rapid Information Visual Processing—three targets (sustained attention), Spatial Span—forward and reverse (visuospatial working memory) and Multitasking Test (executive function). Participants will be diagnosed as either cognitively normal, having mild cognitive impairment or dementia by a three-member expert panel of psychiatrists who will review scores from the MMSE, CDR and neurocognitive battery based on Petersen's mild cognitive impairment criteria and the Diagnostic and Statistical Manual of Mental Disorder, DSM-V for dementia criteria.</td>
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<tr>
<td>Caregiver burden</td>
<td>The widely used 22-item Zarit Burden Interview will measure caregiver burden and will be conducted through a phone interview with participants’ caregiver/family member.</td>
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<tr>
<td>QoL</td>
<td>QoL will be determined by the 13-item WHO QoL assessment for older adults (WHOQoL-AGE) scored on a 5-point Likert scale, with higher scores indicating higher QoL.</td>
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<tr>
<td>Sleep quality</td>
<td>Sleep quality will be assessed by locally validated 19-item Pittsburgh Sleep Quality Index. Used in both clinical and general population, the STOP-Bang questionnaire will also be used to screen for obstructive sleep apnea.</td>
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<tr>
<td>Life satisfaction</td>
<td>Life satisfaction will be measured by a 5-item Satisfaction With Life Scale scored on a 7-point Likert response scale as well as extended questions on nine different areas of life such as health, financial situation, community elderly service, government elderly policy, friendships, spouse, children, leisure activities and current work status.</td>
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<tr>
<td>Perceived health state</td>
<td>Perceived health state will be assessed by EuroQol-5D-3L (EQ-5D-3L), that was developed by EuroQol Group: (1) the EuroQol-5D (EQ-5D) descriptive system that measures five dimensions of health (mobility, self-care, usual activities, pain/discomfort and anxiety/depression) and (2) the EuroQol (EQ) visual analogue scale where participants indicate their health state ranging from 0 (worst imaginable health) to 100 (best imaginable health).</td>
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Variables | Instrument/scale |
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Attitudes to ageing | The 24-item Attitudes to Ageing Questionnaire\(^7\) will be used to measure older adults’ attitudes to the ageing process using a 5-point Likert scale. |
Gratitude | Dispositional gratitude will be determined by the 6-item Gratitude Questionnaire\(^7\)\(^8\) scored on a 7-point Likert scale (strongly disagree to strongly agree). |
Compassion | The 10-item Compassion Scale\(^7\)\(^9\) is used to measure five dimensions of compassion: generosity, hospitality, objectivity, sensitivity and tolerance across social networks and relationships using a 1 (none) to 7 (all) response scale. |

Social factors

Parenting style | A self-developed 13-item Personal and Parents’ Parenting Style Scale\(^42\) is used to examine the relationship between participants’ parenting strategies and how they were parented as children, scored on a 1 (never) to 5 (always) scale. |
Social connectedness | Social connectedness/isolation will be measured by the 6-item Friendship Scale\(^21\)\(^80\) scored on a 5-point Likert scale, whereby higher scores indicate higher level of connectedness (\(\alpha = 0.83\)). |
Perceived social support | Perceived social support is determined by a self-developed scale that consists of an open-ended question (‘How many close friends/relatives do you have?’) and seven items on perceived social support scored on a 5-point Likert scale. |
Intergenerational communication | The Perceptions of Intergenerational Communication Scale\(^81\) will be used to measure perceived communication between generations scored on a 7-point Likert-type scale, ranging from 1 (strongly disagree) to 7 (strongly agree). |

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*Venipuncture procedure will be scheduled in conjunction with another visit.
†Available to the first 300 literate participants only as the test was added in at a later stage of the study.
percentages for continuous variables and proportions for categorical variables. Univariate and multivariate linear regression analyses will be conducted to determine associations between continuous outcomes; logistic regressions will be used for dichotomous outcomes. Specifically, group differences (between cognitive diagnoses or self-reported medical conditions) will be analysed using independent-sample t-tests and analysis of variance. Relationships between physical and mental health, psychosocial and demographic variables will be analysed using multiple regressions and structural equation models. In addition, mixture models will be used to identify subgroups of participants based on their psychosocial, physical and mental health characteristics. Prediction modelling will also be attempted using the new covariates under study (eg, attitudes to ageing and intergenerational influence). P values of <0.05 will be considered statistically significant.

ETHICS AND DISSEMINATION
Ethics approval from the National University of Singapore Institutional Review Board (NUS-IRB Reference code: H-17-047) was obtained on 12 October 2017. The CHI study will be conducted in accordance with the principles of Good Clinical Practice and adhere to the Human Biomedical Research Act which provides a legal framework for researchers in Singapore to conduct research and the use of human tissue.

Written informed consent will be obtained from all the participants after objectives and procedures of the research are fully explained to them by a member of the research team. Participants will also be informed that they can withdraw from the research at any time without giving any reasons. In addition, participants will be given the option of (1) providing their coded human biological materials and data for use in future research, (2) being recorded for the free flow speech segment and (3) being contacted for future intervention studies, incidental findings, changes to the research and follow-up appointment for memory concerns. Participants with dementia will also be asked to invite their legally acceptable representatives to the consent-taking process and data collection. The research team will ascertain that any persons making a decision on behalf of the participant with dementia acts in the best interest of the participant and takes into account of the participant’s wishes and feelings.

Results from the cohort study will be disseminated by publication of peer-reviewed manuscripts, presentations at scientific meetings and/or conferences with local stakeholders. The researchers may also communicate aggregated results to members of the public and clinical professionals through ad hoc meetings/events or mass media releases.

DISCUSSION
Using a cmRCT design, the CHI study seeks to explore vulnerability and resiliency factors associated with ageing with subsequent clinical trials of interventions and community programmes that could potentially hold translational significance. The study intends to recruit 1000 older adults and collect a comprehensive set of biological, psychological and social data. Meaningful associations between outcome measures found will provide significant information on the physical and mental health of older adults in Singapore. Results will also help identify at-risk groups of older adults and test out subsequent interventions targeted at improving health outcomes. In addition, having an interdisciplinary team of investigators enables the introduction of in-depth and novel health assessments such as oral examination, cardiovascular investigations, olfactory test and speech analysis. Given the limited sample size and cost considerations, this study excluded genetic and other in-depth measures (eg, neuroimaging and Structured Clinical Interview for Diagnostic and Statistical Manual of Mental Disorders Fifth Edition, SCID-5) which could have added value to findings. Moreover, several ageing cohort studies in Singapore have previously collected the above-mentioned data; hence, due to the limited resources, these measures were excluded in favour of other novel measures. Recruitment of participants in a confined area may also affect generalisability of the results. Nevertheless, the CHI cohort is culturally relevant and will provide clinicians, researchers and policymakers with information on improving the physical and mental health of older adults in Singapore.

Author affiliations
1Department of Psychological Medicine, Yong Loo Lin School of Medicine, National University of Singapore, Singapore
2Department of Psychiatry, Sengkang General Hospital, Singapore
3Dean, Faculty of Dentistry, National University of Singapore, Singapore
4Centre for Oral Health, National University Health System, Singapore
5Department of English Language and Literature, Faculty of Arts and Social Science, National University of Singapore, Singapore
6Department of Sociology, National University of Singapore, Singapore
7Centre for Family and Population Research, National University of Singapore, Singapore
8Saw Swee Hock School of Public Health, National University of Singapore, Singapore
9Department of Orthopaedic Surgery, Yong Loo Lin School of Medicine, National University of Singapore, Singapore
10Department of Medicine, Yong Loo Lin School of Medicine, National University of Singapore, Singapore
11Department of Cardiology, National University Heart Centre, Singapore
12Department of Japanese Studies, National University of Singapore, Singapore
13Department of Ophthalmology, National University Health System, Singapore
14Department of Otolaryngology, Yong Loo Lin School of Medicine, National University of Singapore, Singapore
15Department of Pharmacy, Faculty of Science, National University of Singapore, Singapore
16Department of Psychological Medicine, National University Hospital, Singapore
17Academic Development Department, Duke-NUS Medical School, Singapore

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**Contributors**
RM, EHK and IR made a significant contribution to the conception of the study and implementation of the protocol. LF, PFA, LHL, ZB, DW, DYW, LTT, KZY, QF, JKL, CTL and TN participated in the design of the study. RM initiated and conducted the study with a team of researchers (including PFA, LHL and RZYL) and is the primary author of the manuscript. RM, RZYL and JY prepared the first draft of the manuscript. RM, JY, RZYL, ZB and PFA made revisions to the manuscript. All authors read and approved the final version of the manuscript.

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**ORCID iD**
Rachael Zhi Yi Lee http://orcid.org/0000-0001-9844-941X

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