Loneliness and isolated living status in middle-aged and older adults in Taiwan: exploration on stress-related biomarkers, depressive symptoms, and disability

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Abstract

Purpose: Loneliness is a subjective feeling by which an individual perceives a lack of closeness in interpersonal relationships. An isolated living status is linked with higher odds of risky health behavior. The conflicting impacts of loneliness and isolated living status on stress-related biomarkers, depressive symptoms, and disability remain unexplained.

Methods: Six hundred twenty-nine participants aged 66.0 (SD=7.3) separated into four groups: “Lonely and Isolated,” “Not Lonely, but Isolated,” “Lonely, but Not Isolated,” and “Neither Lonely, nor Isolated,” were retrieved from the Social Environment and Biomarkers of Aging Study conducted in 2000. Follow-up health indicators in 2006 included three stress-related biomarkers, depressive symptoms, and two physical disability indicators. A hierarchical regression was performed for the analysis.

Results: Firstly, compared to the “Neither Lonely nor Isolated” group, only the “Lonely, but Not Isolated” participants at baseline retained positive associations with the stress-related biomarkers levels 6 years later (urine cortisol level (B=9.25, 95% CI=3.24-15.27), serum Interleukin-6 level (B=2.76, 95% CI=0.72-4.79) and the serum high sensitivity C-reactive protein (hsCRP) level (B=0.40, 95% CI=0.17-0.62)). However, such associations were not observed in the “Lonely and Isolated” participants. Secondly, only “Lonely and Isolated” participants at baseline were positively associated with depressive symptoms 6 years later (B=1.70, 95% CI=0.11-3.30). Finally, the associations between combinations of loneliness and isolated living status and physical disability were eliminated after adjusting the covariables.

Conclusion: Four combinations of loneliness and isolated living status were associated with different impacts on stress-related biomarkers, depressive symptoms, and physical disability. Further dynamic investigations are warranted.

Keywords: Loneliness, Isolated living status, Stress-related biomarkers, Depressive symptoms, Physical disability

Background

Living with active social participation is considered a modifiable determinant to reduce the adverse health effects of loneliness and social isolation. However, in Taiwan, 9-20% of older people live alone, and 21% of older adults reported loneliness in 2017 [1]. In addition, according to reports from the World Health Organization, the world’s population over 60 years will increase...
Loneliness is defined as a distressing feeling that occurs due to the discrepancy between desired and available relationships. It is a subjective measure, where an individual perceives a lack of closeness or depth in interpersonal relationships [4]. The prevalence of loneliness has been recognized to range from 7% to 49% in the aging population [5]. As previously reported, up to a 50% higher prevalence of loneliness has been observed in individuals older than 80 [6]. Loneliness is predictive of increased morbidity and mortality [7]. A significant impact of loneliness on physical and mental health has also been found, where more severe levels of loneliness are associated with higher risks of coronary heart disease, increases in depressive symptoms, suicide, cognitive impairment, and functional disability [8]. Personality characteristics, such as neuroticism, have been shown to increase the risk of loneliness and to moderate the risk of depression [9, 10]. In addition, greater loneliness is associated with higher levels of Interleukin-6 (IL-6), C-reactive protein (CRP) [11], and lower responsivity of cortisol [12]. Higher levels of IL-6, CRP, and dysregulation of the cortisol response are linked with cardiovascular disease and depression [13]. A person who feels lonely will tend to utilize health services excessively, which will result in an additional financial burden on the medical system [5].

Social isolation is an objective measure of limited social contact between an individual and society, and it is often measured based on social network size, diversity, or frequency of social activity [14]. People suffering from social isolation are at an increased risk of overt diabetes, coronary heart disease, dementia, and increases in the rate of all-cause mortality [15]. Those who experience social isolation will experience an increase in hypothalamic-pituitary-adrenocortical axis (HPA axis) activation and higher levels of inflammatory markers, where such effects are more dependent on the disruption of a social bond between a significant pair than objective isolation per se [16]. In previous studies, researchers reported that about one third of the elderly population lived alone in developed countries from 2000 and 2010 [17]. Living alone in later life may be correlated with a variety of factors, including death of the spouse, divorce, poorer physical or mental health, and personal choice. However, those who begin to live alone after a divorce or the death of a spouse have been shown to have a higher risk of mortality compared with those who live alone for other reasons [18]. In addition, social network structure and function are strongly intertwined with anxiety and depressive symptoms in older adults. For example, one longitudinal mediation analysis showed that social isolation is predictive of higher levels of loneliness and depression among older Americans [19]. The relationship between loneliness and depression may be bi-directional, which often worsens lonely individuals’ health and social activity levels even further [3].

Although loneliness and social isolation have been linked with depression, cardiovascular disease and declined physical function, when researchers simultaneously examine loneliness and social isolation with health, the results are often mixed [20]. People can feel lonely without being isolated and can feel lonely despite living with others. A comparison of subjective individual-level factors and objective environment-related factors showed that loneliness in older adults is higher in the most deprived environments independent of individual-level factors [21]. However, loneliness and social isolation have only been found to be moderately correlated [22]. The potential grouping of people characterized by this viewpoint have not been explored in detail, and measurements of social isolation remain inconsistent [22]. There is a lack of research examining loneliness and an isolated living status (living alone and unmarried), and how these groups might be linked with health.

Assuming that loneliness and an isolated living status may act independently and lead to different health trajectories through their effects on health-risk behaviors, this study is aimed toward exploring the inconsistent findings concerning the role of risk of loneliness and isolated living status on stress-related biomarkers, depressive symptoms, and disability. The participants were separated into four groups [22]: lonely and isolated; lonely but not isolated; not lonely, but isolated; not isolated and not lonely to elaborate on the different impacts of loneliness and isolated living status on health outcomes.

Methods

Participants

Six hundred twenty-nine participants were enrolled from the Social Environment and Biomarkers of Aging Study (SEBAS 2000 and 2006) in this study. The SEBAS is an extension of the Taiwan Longitudinal Study on Aging (TLSA), which began in 1989 and has undergone repeated follow-ups every 3-4 years with a nationally representative sample of adults aged 60 and older. Younger refresher cohorts of the TLSA were added in 1996 and 2003, and as of 1996, participants in the TLSA were representative of older adults aged 50 and older in Taiwan. The first wave of the SEBAS, based on a sub-sample of participants from the 1999 TLSA, was conducted in 2000. A total of 1,713 participants aged 54
and over in 27 townships were selected from the TLSA 1999. There were 1,023 participants who had been interviewed and completed a hospital-based health examination in the first wave of the SEBAS in 2000. The second wave of SEBAS was conducted in 2006 using a protocol similar to that for SEBAS 2000. In both SEBAS waves, health status, health behavior, exposure to stressors, and social relationships were collected. With the exception of participants who passed away or were lost to follow-up, there were 757 participants who had been interviewed for the SEBAS 2006, and 639 of them had completed the health and hospital-based examination assessment [23]. To ensure the reliability of the self-report questionnaire used in the present study, we had to exclude 8 participants who might have had cognitive function impairments based on Short Portable Mental State Questionnaire (SPMSQ) scores $\leq 7$, as well as 2 participants with missing documents. Therefore, there was a total of 629 participants aged 54 and over at baseline and with interview and biomarker data in both 2000 and 2006 analyzed in this study (Fig. 1).

**Measures**

*Explanation of the variables, including loneliness and isolated living status*

Loneliness was assessed in 2000 and 2006 by asking the participants “In the past week, have you experienced the following situations or feelings of loneliness (Felt isolated, with no companions)?” The four answer options were “never,” “rarely (1 day),” “sometimes (2-3 days),” and “often (>4days).” Responses were recorded as a 4-point Likert-type response, ranging from “never” to “often.” Participants who answered “never” were categorized as the non-lonely group. Those who answered “rarely,” “sometimes (2-3 days),” or “often” were grouped together as the “lonely” group.

Multidimensional measures of the Isolated Living Status Index indicate that different aspects of social interaction may have a relationship with depressive symptoms and stress-related biomarkers [24]. Indicators of social isolation in a previous study included living alone, being unmarried, low participation in social activities, and infrequent contact with network members [25]. In the present study, two items (married and living alone) were
used to develop an index of isolated living status in 2000. One point was assigned to participants who were not married/separated/divorced/widowed. One point was assigned to participants who were living alone. These two items were added together to obtain an overall isolated living status index ranging from 0-2. All participants who responded ‘married, and not living alone’ were classified as “non-isolated living status” (isolated living status index=0). All participants who had Isolated Living Status Index scores ≥1 were classified as experiencing isolation.

Combining loneliness with living status, four categories of loneliness and living status were identified: 1. Lonely and Isolated, 2. Not Lonely, but Isolated, 3. Lonely, but Not Isolated, 4. Neither Lonely, nor Isolated. To compare within-group differences, Neither Lonely, nor Isolated was used as a reference category.

The three domains of the outcome variables: stress-related biomarkers (Cortisol, IL-6, and hsCRP), depressive symptoms, and disability (Mobility; IADL)

Firstly, in the SEBAS 2000 and 2006, overnight and 12-hour urine specimens (7pm to 7am) was collected to minimize person-to-person variations and diurnal variations, which provided a more accurate measurement of baseline levels of stress-related biomarkers. The participants provided the urine specimen, and a phlebotomist drew a blood sample. Data from duplicate samples indicated intra-lab correlations of 0.8 or higher and inter-lab correlations of 0.6 or higher.

Urine cortisol, measured by using high-performance liquid chromatography (HPLC) in both the 2000 and 2006 SEBAS, were used to assess HPA activity. Depression was linked with higher degrees of dysregulation of HPA activity and higher basal cortisol levels. By contrast, higher levels of cortisol were noted when older adults felt acutely lonely, where their HPA activity would be blunted in the chronic phase [26, 27]. IL-6 was measured using enzyme-linked immunoassays (EIA; Endogen, Pierce Biotechnology) in the SEBAS 2000, as well as enzyme-linked immunosorbent assays (ELISA; R&D Systems). Measurements using EIAs and ELISAs are virtually the same able Mental State Questionnaire (SPMSQ, Cronbach's alpha =0.77) in 2000 [31]. Participants were asked, “Tell me your address or where this is?”, “What is your mother's surname?”, “Who is the President?”, “Who were the previous presidents of Taiwan?” along with the serial 3s subtraction task. Higher scores indicated higher cognitive ability. If the
participants had more than three errors, they were suspected of having cognitive impairment. Participants who had SPMSQ scores ≥7 were excluded in the SEBAS 2000 because of potential cognitive impairment.

Personal stress was assessed by using the 10-item Perceived Stress Scale (PSS, Cronbach’s alpha = 0.81) in 2006 [32]. The PSS is a 5-point scale ranging from 0 to 4 (0 = never, 4 = always). Participants were asked, during the last month, “How often upset by unexpected events?”, “How often felt unable to control important things?”, “How often felt nervous or stress?”, “How often could you not cope with all had to do?”, “How often have been angered by things outside your control?”, “How often felt difficulties so bad they could not be overcome?”, “How often felt confident about handling personal problems?”, “How often felt thing going your way?”, “How often been able to control irritations in your life?”, and “How often felt you were on the top of things?” The last 4 question scores have been reversed as below: 0 = 4, 1 = 3, 2 = 2, 3 = 1, 4 = 0. Higher 10-item PSS scores indicated greater perceived stress.

Socio-demographic variables included age, sex, education (No schooling, 1-6 years, 7-9 years, 9-12 years, ≥12 years).

Statistical analyses
Analyses were carried out with SAS 9.4. Statistical software and included descriptive statistics and a hierarchical linear regression. Descriptive analyses were performed for all variables at baseline and 6 years later. We used Student’s t-tests and Pearson’s correlations to assess the associations between the demographic data and the outcome variables (cortisol, IL-6, and hsCRP, CES-D, mobility, and IADL). Furthermore, paired t-tests and McNemar’s test were used to assess associations between the two waves of the SEBAS. All tests were evaluated at a 0.05 level of statistical significance.

We applied hierarchical regression models to investigate the association between different combinations of baseline loneliness and isolated living status and three outcome variable domains 6 years later: stress-related biomarkers (cortisol, IL-6, and hsCRP), depressive symptoms, and physical disability (mobility/IADL disability). For each outcome variable, three models were estimated. In Model 1, we regressed the combinations of baseline loneliness and isolated living status on the outcome variables 6 years later. In Model 2, we then repeated the analyses, adjusting for the baseline outcome variables. In Model 3, we repeated the analyses adjusting for confounding variables such as age, sex, educational attainment, baseline depressive symptoms, PSS score 6 years later, and comorbidity 6 years later. In terms of statistical power, only the comorbidity item accounting for more than 5% of participants was treated as an independent variable in Model 3.

The results are presented as regression coefficients (B) with a 95% confidence interval.

Results
The characteristics of the sample are summarized in Table 1. The mean age of this study population was 66.0 years (SD = 7.3) at baseline, and 58.8% of the sample was male, 77.9% of the participants were married; 5.7% lived alone, and 14.5% were experiencing loneliness. The outcome variables at baseline and 6 years later were as follows: urine cortisol level (baseline: mean = 19.7 ± 18.3 μg/L, 6 years later: mean = 12.1 ± 17.2, p < 0.001), serum IL-6 level (baseline: mean = 2.9 ± 3.3 pg/mL, 6 years later: mean = 4.0 ± 6.0, p < 0.001), serum hsCRP (baseline: mean = 0.3 ± 0.6 mg/dL), CES-D score (baseline: mean = 4.7 ± 4.8, 6 years later: mean = 4.9 ± 5.2, p = 0.39), mobility disability (baseline: mean = 2.7 ± 4.2, 6 years later: mean = 5.1 ± 6.2, p < 0.001), and IADL disability (baseline: mean = 0.6 ± 1.8, 6 years later: mean = 2.0 ± 3.6, p < 0.001). To examine for potential bias and collinearity, a Pearson’s correlation of all variables, including age and education at baseline, and PSS, CES-D, mobility, IADL index scores, cortisol, IL-6, and hsCRP 6 year later were performed, as summarized in Table 2. The baseline descriptive statistics for the four groups, Lonely and Isolated, Not Lonely, but Isolated, Lonely, but Not Isolated, and Neither Lonely, nor Isolated, are presented in Table 3. Also, Table 3 shows the group differences, adjusted using the Bonferroni correction for pairwise comparisons of the four groups at baseline, on age (F = 12.2, p < 0.001), sex (χ^2 = 30.5, p < 0.001), comorbidity (F = 4.3, p < 0.001), urine cortisol level (F = 4.2, p = 0.001), serum IL-6 level (F = 0.8, p = 0.51), CES-D score (F = 51.4, p < 0.001), mobility disability (F = 20.3, p < 0.001), and IADL disability (F = 10.8, p < 0.001). The group differences, adjusted using the Bonferroni correction for pairwise comparisons of the four groups 6 years later, were as follows: PSS score (F = 9.1, p < 0.001), urine cortisol level (F = 4.5, p = 0.004), serum IL-6 level (F = 3.5, p = 0.02), serum hsCRP level (F = 7.7, p < 0.001), CES-D score (F = 11.0, p < 0.001), mobility disability (F = 9.6, p < 0.001), and IADL disability (F = 7.0, p < 0.001).

Effect of baseline loneliness and isolated living status on stress-related biomarkers 6 years later
First of all, as shown in Table 4, Model 1 on the levels of urine cortisol and serum IL-6 indicated that only the Lonely, but Not Isolated group was positively associated with greater levels of urine cortisol (B = 9.11, 95% CI = 3.47-14.75) and serum IL-6 (B = 2.96, 95% CI =
1.02-4.91) 6 years later. Model 2 further added the baseline levels, and the positive association remained (Cortisol levels: \( B = 9.34, 95\% \text{ CI} = 3.97-14.95 \); IL-6 levels: \( B = 3.20, 95\% \text{ CI} = 1.31-5.09 \)). After adjusting for age, sex, education, depressive symptoms at baseline, and comorbidity (high blood pressure, diabetes, heart disease, lower respiratory disease, and arthritis/rheumatism) 6 years later in Model 3, the Lonely, but Not Isolated group retained a consistently positive association with urine cortisol levels (\( B = 9.25, 95\% \text{ CI} = 3.24-15.27 \)) and serum IL-6 levels 6 years later (\( B = 2.76, 95\% \text{ CI} = 0.72-4.79 \)).

Lastly, because of limited data on serum hsCRP levels in the SEBAS 2000, we performed two steps in the regression model to assess the associations between combinations of loneliness and isolated living status at baseline and the serum hsCRP levels 6 years later. As shown in Table 4, Model 1 on the serum hsCRP levels indicated that only the Lonely, but Not Isolated (\( B = 0.50, 95\% \text{ CI} = 0.30-0.71 \)) group was positively associated with greater serum hsCRP levels 6 years later. After adjusting for other controls, including age, sex, education, depressive symptoms at baseline, PSS 6 years later, and comorbidities 6 years later in the second step, the Lonely, but Not Isolated group at baseline remained consistently positively associated with serum hsCRP levels 6 years later (\( B = 0.40, 95\% \text{ CI} = 0.17-0.62 \)).

Table 1 (continued)

| Variable | SEBAS2000 | SEBAS2006 | Paired t or \( \chi^2 \) | P |
|----------|-----------|-----------|-------------------------|---|
| Yes | 27 (4.3) | 40 (6.4) | | |
| No | 602 (95.7) | 589 (93.6) | | |
| Perceived Stress Scale | | | - | - |
| Stress-related biomarkers | | | | |
| Cortisol levels | | | | |
| IL-6 levels | 2.9±3.3 | 4.0±6.0 | -4.06 | <0.001 |
| hsCRP | - | 0.3±0.6 | | - |
| Depression (CES-D) | | | | |
| Disability | | | | |
| Motility | 2.7±4.2 | 5.1±6.2 | -11.8 | <0.001 |
| IADLs | 0.6±1.8 | 2.0±3.6 | -10.8 | <0.001 |

Note. Numbers are Mean±SD or N (%). CES-D: The Center for Epidemiologic Studies Depression Scale; IADLs: Instrumental Activities of Daily Living; IL-6: Interleukin-6; hsCRP: High sensitivity C-reactive protein. #9-item CES-D score (excluding lonely). *\( p < 0.05 \); **\( p < 0.01 \); ***\( p < 0.001 \).
Model 2 further added the CES-D scale at baseline, and we observed that only when loneliness and isolated living status occurred together (the Lonely and Isolated group) were the CES-D scores higher ($B = 2.34$, $95\%\, CI = 0.72-3.97$). The positive association between loneliness and isolated living status and the CES-D scores remained robust after further adjustment for age, sex, education, PSS 6 years later, and comorbidities 6 years later ($B = 1.70$, $95\%\, CI = 0.11-3.30$).

Effect of baseline loneliness and isolated living status on physical disability 6 years later
As shown in Table 4, Model 1 on mobility disability indicated that the Lonely and Isolated ($B=4.07$, $95\%\, CI=2.15-5.98$), Not Lonely, But Isolated ($B=2.38$, $95\%\, CI=1.07-3.70$), and Lonely, Not Isolated ($B=2.02$, $95\%\, CI=0.23-3.81$) groups had a positive association with greater levels of mobility disability 6 years later. Model 2 further added mobility disability at the baseline, and the previous associations were eliminated. Furthermore, after adjusting for age, sex, education, depressive symptoms at baseline, PSS 6 years later, and comorbidities 6 years later ($B = 1.70$, $95\%\, CI = 0.11-3.30$).

Discussion
In this study, compared to the Neither Lonely, Nor Isolated group, only people who lived with others and/or got married still experienced loneliness and were prone to having higher levels of cortisol, IL-6, and hsCRP. Secondly, people experiencing both loneliness and an isolated living status were found to have more depressive symptoms, independent of their age, sex, education, number of comorbidities, stress, and baseline depressive symptoms. However, neither loneliness nor isolated living status was found to be related to the levels of mobility disability or IADL disability.

Synergic effects of loneliness and isolated living status on stress-related biomarkers, depressive symptoms, and physical disability
Loneliness and isolated living status have been linked with the Hypothalamic-Pituitary-Adrenal axis and systemic inflammation. In our study, although we found a combination of loneliness and isolated living status to be associated with higher levels of depressive symptoms, this association between loneliness and isolated living status at baseline and the stress-related biomarkers was not observed 6 years later. The theory of allostatics and the allostatic load in social dynamics, stress, and physiological responses may explain this phenomenon [33]. Participants who are socially isolated exhibit dysregulated

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### Table 2 Correlation Matrix for the Continuous Variables in the Study

|          | 1  | 2  | 3  | 4  | 5  | 6  | 7  | 8  | 9  |
|----------|----|----|----|----|----|----|----|----|----|
| 1. Age   | -  | -0.06 | -0.05 | 0.20*** | 0.43*** | 0.37*** | 0.03 | 0.10 | 0.03 |
| 2. Education | -0.12 | -<0.01 | -0.13* | -0.17** | -0.11* | -0.07 | -0.02 | -0.09 |
| 3. PSS   | -0.03 | -0.001 | -<0.01 | 0.30*** | 0.16** | 0.09 | 0.07 | 0.05 | 0.09 |
| 4. CES-D$^a$ | 0.21*** | -0.17** | 0.39*** | -<0.01 | 0.43*** | 0.37*** | <0.01 | 0.10 | 0.02 |
| 5. Mobility | 0.48*** | -0.22*** | 0.15* | 0.45*** | -<0.01 | 0.76*** | <0.01 | 0.16** | 0.08 |
| 6. IADLs | 0.48*** | -0.21*** | 0.18** | 0.40*** | 0.83*** | -<0.01 | 0.06 | 0.18** | 0.05 |
| 7. Cortisol | -0.04 | -0.07 | 0.20** | 0.12 | -0.01 | 0.08 | -0.06 | 0.07 | 0.26*** |
| 8. IL-6  | 0.18** | -0.17* | 0.04 | 0.07 | 0.02 | 0.06 | -0.03 | -0.54*** |
| 9. hsCRP | -0.001 | -0.07 | 0.10 | 0.16* | 0.14* | 0.20** | -0.01 | 0.10 | - |

Note. The upper diagonal was based on men ($N=370$); the lower diagonal was based on women ($N=259$); PSS, The Perceived Stress Scale; CES-D, The Center for Epidemiologic Studies Depression Scale; IADLs, Instrumental Activities of Daily Living; IL-6, Interleukin-6; hsCRP, High sensitivity C-reactive protein. $^a$9-item CES-D score (excluding lonely); $^*p < 0.05$; $^{**}p < 0.01$; $^{***}p < 0.001$
patterns of mood and physiology. The theory of allostatic load means that chronic, sustained stress creates wear and tear on regulatory systems [34]. Our findings suggest that lonely and isolated participants may have failed to regulate their emotions within groups due to an increased allostatic load, which led to depression and attenuated inflammatory responses.

Furthermore, we reaffirmed the hypothesis that people who are both lonely and isolated rather than lonely or isolated individually are at higher risk for the development of depression. Some investigators have suggested that gender differences in individuals living alone may be a predictor of the gender differences found in depression. Older women are more likely than older men to be unmarried (widowed/divorced/separated) and to live alone than older men. However, older men who live alone have more depressive symptoms than older women who live alone [35]. Men also reported more loneliness than women [36]. In our study, older men who were lonely and socially isolated tended to have higher levels of depressive symptoms ($B = 3.15, p = 0.01$), but this association was not

Table 3  Sociodemographic and Health-related Characteristics of Participants in the Four Loneliness and Isolated Living Status Groups

|                                | aNeither Lonely, nor Isolated (N=438) | bNot Lonely, but Isolated (N=100) | cLonely, but Not Isolated (N=49) | dLonely and Isolated (N=42) | F or $\chi^2$ Pairwise comparisons among means |
|--------------------------------|---------------------------------------|----------------------------------|---------------------------------|-----------------------------|------------------------------------------|
| Age                            | 65.0±7.2                              | 69.2±7.0                         | 66.2±7.1                        | 69.1±6.8                    | 12.2*** a=b, a=c                       |
| Sex                            |                                       |                                  |                                 |                             | 30.5***                                  |
| Male                           | 287(65.5)                             | 39(39)                           | 27(55.1)                        | 17(40.5)                    |                                         |
| Female                         | 151(34.5)                             | 61(61)                           | 22(44.9)                        | 25(59.5)                    |                                         |
| Education                      |                                       |                                  |                                 |                             | 20.8                                    |
| No schooling                   | 97(22.2)                              | 29(29)                           | 20(40.8)                        | 16(38.1)                    |                                         |
| Elementary (<6 years)          | 195(44.5)                             | 46(46)                           | 17(34.7)                        | 20(47.6)                    |                                         |
| Junior High (7-9 years)        | 53(12.1)                              | 10(10)                           | 2(4.1)                          | 3(7.1)                      |                                         |
| Senior High (9-12 years)       | 50(11.4)                              | 11(11)                           | 6(12.2)                         | 2(4.8)                      |                                         |
| College (>12 years)            | 43(9.8)                               | 4(4)                             | 4(8.2)                          | 1(2.4)                      |                                         |
| Comorbidity†                   | 0.8±0.9                               | 0.9±1.0                          | 1.1±1.1                         | 1.2±1.2                     | 4.3*** c=d                            |
| Loneliness2006                 |                                       |                                  |                                 |                             |                                         |
| Yes                            | 43(9.8)                               | 17(17.0)                         | 10(20.4)                        | 17(40.5)                    |                                         |
| No                             | 395(90.2)                             | 83(83.0)                         | 39(79.6)                        | 25(59.5)                    |                                         |
| Isolated living status2006     |                                       |                                  |                                 |                             |                                         |
| Yes                            | 39(9.8)                               | 99(99.0)                         | 10(20.4)                        | 41(97.6)                    |                                         |
| No                             | 399(91.1)                             | 1(1.0)                           | 39(79.6)                        | 1(2.4)                      |                                         |
| Perceived Stress Scale         | 8.6±5.9                               | 8.6±6.9                          | 13.4±7.1                        | 11.5±7.7                    | 9.1*** c=d                            |
| Stress-related Biomarkers      |                                       |                                  |                                 |                             |                                         |
| Cortisol levels2000            | 21.2±20.4                             | 16.6±13.3                        | 18.3±10.6                       | 12.7±7.9                    | 4.2*** c=d                            |
| Cortisol levels2006            | 11.9±15.7                             | 9.2±13.2                         | 21.1±26.7                       | 10.1±23.6                    | 4.2*** a=c, b=c                       |
| IL-6 levels2000                | 2.9±3.3                               | 2.9±2.4                          | 2.7±2.7                         | 3.7±4.7                      | 0.8                                    |
| IL-6 levels2006                | 3.7±5.3                               | 3.9±4.3                          | 6.6±11.2                        | 5.1±7.4                      | 3.5** a=c                             |
| hsCRP2000                      | -                                     | -                                | -                               | -                           |                                         |
| hsCRP2006                      | 0.3±0.5                               | 0.3±0.5                          | 0.8±1.6                         | 0.3±0.3                      | 7.67*** a=c, b=c, d=c                 |
| Depression†                    |                                       |                                  |                                 |                             |                                         |
| CES-D2000                      | 3.8±4.0                               | 4.3±3.8                          | 10.5±6.1                        | 9.2±6.0                      | 51.4*** a=c, a=d, b=c, b=d            |
| CES-D2006                      | 4.3±4.7                               | 5.4±5.5                          | 6.6±5.4                         | 8.5±7.1                      | 11.0*** a=c,d, b=d                   |
| Disability                     |                                       |                                  |                                 |                             |                                         |
| Mobility2006                   | 2.0±3.4                               | 3.7±4.3                          | 5.3±6.0                         | 5.3±5.5                      | 20.3*** a=b, a=c,d                   |
| Mobility2006                   | 4.3±5.6                               | 6.7±6.7                          | 6.3±7.0                         | 8.3±7.3                      | 9.6*** a=b, a<d                      |
| IADLs2000                      | 0.4±1.5                               | 1.0±2.1                          | 1.5±2.6                         | 1.4±2.1                      | 10.8*** a,b, a=c, a=d                |
| IADLs2006                      | 1.6±3.2                               | 2.9±4.4                          | 2.5±4.4                         | 3.5±4.3                      | 7.0*** a,b, a<d                      |

Note. Numbers are Mean±SD or N (%). CES-D The Center for Epidemiologic Studies Depression Scale; IADLs Instrumental Activities of Daily Living; IL-6 Interleukin-6; hsCRP High sensitivity C-reactive protein. †total 9-item comorbidity score. ‡9-item CES-D score (excluding lonely). *p < 0.05; **p < 0.01; ***p < 0.001
significant in women ($B = -0.15, p = 0.90$) after adjusting for the confounding variables.

The relationships between loneliness, isolated living status, and physical disability in older individuals have remained unclear and inconsistent. Some studies have reported that having a large number of social relationships is associated with fewer physical disabilities [37], but some studies have reported limited or no significant associations [38]. In one cross-sectional study examining the combined effect of marital status and living arrangement, married older adults living with children had better IADL scores than those who were unmarried and living with children [39]. In addition, feelings of loneliness may exacerbate existing vulnerabilities in health that lead to disabilities, either through poor health behavior or through an inflammatory or cardiovascular pathway. A later prospective study showed both social isolation and loneliness to be associated with a decrease in gait speed [40]. In our study, we found that loneliness and isolated living status at baseline were positively correlated with mobility disabilities and IADL disabilities 6 years later. However, these associations were not significant after adjusting for the baseline conditions. One consideration is that socio-economic status may act as a buffer against the effects of social relationships on functional disabilities. Greater social resources were associated with better self-rated health as well as a composite measure of physical function. Even with these findings, the mechanisms remain unclear, and further investigation may be needed.

**Effects of loneliness without an isolated living status on stress-related biomarkers, depressive symptoms, and physical disability**

Loneliness affects people at any stage of life. Some participants tend to widen their social network to achieve...
a desirable level of social interaction, but some of them do not [41]. In our study, we found that the Lonely, but Not Isolated participants at baseline were positively associated with higher levels of depressive symptoms and mobility disabilities 6 years later. However, this association was not observed after adjusting for the baseline conditions. Interestingly, we found a consistent positive association between loneliness and the stress-related biomarker levels (Table 4). A lonely participant with unsatisfactory levels of social support had a high likelihood of psychological distress and an inflammatory response [42]. After a sex stratification analysis, older women who felt lonely but were not socially isolated had higher levels of Cortisol, IL-6, and hsCRP, but such associations were not observed in older men. Loneliness was associated positively with demographic and environmental factors such as physical illness, a small social network, and a lack of a spousal confidant. On the contrary, the same objective social relationships (e.g., spouse) and higher levels of education were found to be protective factors for health. Loneliness and isolated living status were independently associated with lower levels of self-rated physical health. The association between loneliness and isolated living status was mediated by the perceived quality of social relationships. For example, even when these individuals had a spouse, the marital relationship could be tense, which can have negative consequences for individuals [43]. In contrast, active social participation can lead to an increase in physical exercise, alleviated loneliness, and lower levels of physical disability. Active participation in social activities plays an important role in maintaining mental and physical well-being [41].

Effects of isolated living status without loneliness on stress-related biomarkers, depressive symptoms, and physical disability

Some gaps between loneliness and isolated living status were found for the Only Isolated group just as was the case with the Only Lonely group. A previous longitudinal mediation analysis suggested that social disconnectedness (e.g., unmarried with infrequent social interaction) predicts higher levels of subsequent loneliness, which in turn predict higher levels of depressive symptoms in a general older adult population [19]. A comparison of loneliness and social isolation indicated that loneliness has a stronger association with depressive symptoms than social isolation [44, 45]. In this study, we found that people experiencing isolated living status without loneliness at baseline were not associated with depressive symptoms 6 years later after adjusting for baseline depression. Older adults who are living alone and unmarried may be able to optimize social relationships or perceived deficits in support [46]. Also, if participants tended to prefer being alone to being with others, this desire for solitude may actually reduce stress levels and enhance mental balance [33]. This might imply why we did not observe significant associations between isolated living status and stress-related biomarkers.

Some studies have suggested that isolated living status is positively associated with physical disabilities. For instance, older people living in substandard neighborhoods have significantly higher incident mobility difficulties than those in less-deprived neighborhoods [47]. In our study, we did not find significant associations between different combinations of loneliness and isolated living status at baseline and mobility/IADL disability 6 years later after adjusting for baseline disabilities and covariates. The associations between isolated living status and physical disability in an aging population may be partly but not fully explained by correlated social and economic circumstances and social relationships [48].

Strengths and limitations

The strengths of this analysis include the longitudinal design with a nationally representative sample cohort in Taiwan. The dataset provided multiple measures of health, demographic factors, and biological indicators for controlling for potential confounding variables. However, there are some limitations that should be noted. First, loneliness was assessed by using one question regarding the perception of loneliness in the past week. This measurement may be less reliable than a composite measurement of loneliness from multiple perspectives [49]. Secondly, compared to the complexity and inconsistency of social isolation measurements, the measurement of isolated living status in the present study may be a simpler indicator by which to explore the effects of objective isolation status on health [50]. However, the different effects of loneliness, isolated living status, and social isolation on health need more examination in future studies. Thirdly, some variables such as depressive symptoms and physical disability were addressed using self-reported rating scales, which may have led to response bias due to personality traits and anxiety [51]. Also, because the way people think about loneliness can be affected by age, sex, cross-cultural differences, and the cognitive/affective process of each individual, more research is needed to investigate whether our findings can be generalized to other populations [52]. Finally, this study only employed two waves of self-reported measurements and biomarkers, which may have fluctuated, so causality cannot be confirmed. Enrolling more waves for the purpose of measurement and checking diurnal changes in stress-related biomarkers may be a more convincing method by which to measure the effects of loneliness and isolated living status on health.
Conclusion
This study simultaneously examines the effects of four combinations of loneliness and isolated living status on physical and mental health in longitudinal data, where each has been shown to have unique associations with levels of stress-related biomarkers (Cortisol, IL-6, and hsCRP), depressive symptoms, and physical disabilities (mobility and IADL). The findings suggested that those who felt lonely without being socially isolated had higher levels of cortisol and inflammatory markers than those who felt lonely and objectively isolated. However, only in the presence of both loneliness and isolated living status did depressive symptoms become more severe. In terms of physical disabilities, a positive association between loneliness, isolated living status, and physical disabilities did not exist after controlling for baseline and confounding variables. Based on our findings, we suggest that both loneliness and isolated living status be included in future studies to explore broader pathophysiological indicators for both physical and mental health.

Abbreviations
hsCRP: High-sensitivity C‑reactive protein; IL‑6: Interleukin‑6; CRP: C‑reactive protein; SEBAS: Social Environment and Biomarkers of Aging Study; TLSA: Taiwan Longitudinal Study on Aging; SPMSQ: Short Portable Mental State Questionnaire; PSS: Perceived Stress Scale; HPLC: High‑performance liquid chromatography; EIA: Enzyme‑linked immunosassays; ELISA: Enzyme‑linked immunosorbent assays; CES‑D: Center for Epidemiologic Studies Depression Scale; IADL: Instrumental Activities of Daily Living; SD: Standard deviation; CI: Confidence interval.

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Authors’ contributions
Study concept and design: CJC, TYT. Acquisition of data: CJC. Analysis and interpretation of data: CJC, TYT, TYW. Drafting of the manuscript: TYT, KKC, CJC. Critical revision of the manuscript for important intellectual content: TYT, CJC, PSC, YKJ, HHT. All authors contributed to and reviewed the final version of the manuscript. The author(s) read and approved the final manuscript.

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Availability of data and materials
Data was publicly accessible with a request to the Ministry of Health and Welfare, Taiwan.

Declarations
Ethics approval and consent to participate
The SEBAS study protocols and procedure were authorized by the Institutional Review Board at Antai Medical Care Cooperation Antai Tian-Sheng Memorial Hospital in Taiwan (Official Approval Code: NIFP-IRB-2000-01), Georgetown University (Washington, D.C., USA. Official Approval Code: 1999–195), and Princeton University (Princeton, New Jersey, USA. Official Approval Codes: #1848, #2193, #2791, #5391)). The Health Data Science Center at National Kung University Hospital granted permission to use the raw data. The Ethical Committee for Human Research at National Cheng Kung University Hospital approved this study (A-ER-106-499) and waived informed consent: Not applicable.

Consent for publication
Not applicable.

Competing interests
All authors declare that they have no conflicts of interest.

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