Mild Cognitive Impairment in Republic of Georgia

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Abstract

Objective: The goal of this study was to estimate the prevalence of mild cognitive impairment (MCI) in Georgia. Method: A population-based study was conducted using Georgian version of the Montreal Cognitive Assessment (MoCA) and its cognitive domain index score. Results: Of the initial cohort of 1,000 subjects, 851 met inclusion criteria. The prevalence of MCI was 13.3%, and it was associated with age >65 years (odds ratio [OR] = 4.51, 95% confidence interval [CI] = [3.00, 6.75]), urban residence (OR = 0.53, 95% CI = [0.33, 0.88]), lower education (OR = 3.99, 95% CI = [2.66, 5.93]), and hypertension (OR = 2.51, 95% CI = [1.68, 3.76]), while amnestic MCI was documented in 9.3%, with higher risk in older subjects (OR = 2.69, 95% CI = [1.66, 4.20]), and diabetics (OR = 2.69, 95% CI = [1.25, 5.98]). Conclusion: In this first population-based study of MCI in Georgia, prevalence was comparable with those reported from the United States and Europe. Observed association of MCI with cardiovascular risk factors has important clinical implication for dementia prevention in Georgia.

Keywords

mild cognitive impairment, dementia, epidemiology, Georgia

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Introduction

Mild cognitive impairment (MCI) is a known risk factor for dementia (Petersen at al., 1999). Despite its current pronounced heterogeneity, the concept of MCI permits timely identification of patients at high risk of developing dementia, thus opening a potential therapeutic window and increasing the significance of controlling modifiable risk factors (Winblad et al., 2004).

Published prevalence rates for MCI vary from as low as 2% to 4% to greater than 20%. A number of prospective population-based studies in the United States, France, and Germany estimate the prevalence among older adults to be between 14% and 18% (Busse, Bischkopf, Riedel-Heller, & Angermeyer, 2003; Ganguli, Dodge, Shen, & DeKosky, 2004; Larrieu et al., 2002; Luck et al., 2007). However, epidemiological data on MCI in low and median income countries are sparse. Similar to Western countries, a population-based study in Kolkata, India, showed an overall prevalence rate of MCI at 14.9%, while a study in Brazil was half as common at 7.1% (Das et al., 2007; Herrera, Caramelli, Silveira, & Nitini, 2002).

Results of the Delphi consensus study revealed a huge gap in data on cognitive impairment epidemiology, and therefore there is an urgent need for epidemiological research in Eastern Europe including Georgia (Ferri et al., 2005).

Given the lack of epidemiological data on MCI in Georgia, we sought to estimate the prevalence of MCI and to characterize its demographics and risk factors in a population-based study.

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**Method**

A cross-sectional one-phase study was conducted to identify subjects with MCI among an urban and rural population of Georgia in individuals aged 40 years or older. The study was conducted during a 1 year period from March 1, 2014, to March 1, 2015. Institutional ethical approval was obtained from the Tbilisi State Medical University (Tbilisi, Georgia) before initiation of the study. Individual consent was obtained before enrollment in the study.

Georgia is a country in the South Caucasian region, bordering the Black Sea, with an area of 69,700 km² and a population of 3.8 million according to 2014 census data (Results of General Population Census of Georgia, 2014). Georgian is a Kartvelian language spoken by Georgians and is the most pervasive of the family of Kartvelian languages. Georgian is written in its own Georgian scripts that is unique in their appearance and consists of a 33-letter alphabet. The predominant ethnic group are Georgians whom form about 86.8% of Georgia’s current population (Results of General Population Census of Georgia, 2014). The overall literacy rate in Georgia is 98.8%, while 53% of population resides in an urban area (Results of General Population Census of Georgia, 2014). Tbilisi is the capital of Georgia, is a major urban center, and almost 28.9% of the 1.1 million persons of Georgia reside in Tbilisi (Results of General Population Census of Georgia, 2014).

Results of General Population Census of Georgia a stratified survey method was used for the study. Georgia is divided into 12 territories (nine regions, one city, and two autonomous republics), and comprised the sampling frame for the study. Primary survey regions were stratified according to population and region type (urban vs. rural). Based on this strategy, one urban area, Tbilisi (central location, six administrative districts, population 1.1 million), and two rural areas—Kakheti (eastern location, eight administrative districts, population 318,000) and Imereti (western location, 12 administrative districts, population 534,000)—were selected. Survey subregions included all administrative districts of Tbilisi (six administrative districts, population 1.1 million) and one administrative district in both the Kakheti region (Sagarejo municipality rural settlement with population 681), and the Imereti region (Sachkhere municipality rural settlement with population 815; Results of General Population Census of Georgia, 2014). Both these settlements in Kakheti and Imereti regions are listed as rural settlements in 2014 Results of General Population Census of Georgia. We randomly selected these two villages: one in eastern and one in western Georgia because of cultural differences.

Study investigators contacted random households within each survey region. Where there was no response, the household was replaced by the next in order.

Sample size calculation was made based on previously reported MCI prevalence that was in a range of 2% to 20% (Busse et al., 2003; Das et al., 2007; Ganguli et al., 2004; Herrera et al., 2002; Larrieu et al., 2002; Luck et al., 2007). Averaging these data, 10% was used as an expected prevalence, allowable margin of error 2%, and 95% confidence interval (CI). Using formula \( n = \left( \frac{z^2 \cdot P (1 - P)}{d^2} \right) \) where \( n \) = sample size, \( z = z \) statistic for the level of confidence, \( P = \) expected prevalence, and \( d = \) allowable margin of error (Arya, Antonisamy, & Kumar, 2012), calculation yielded sample size of \( n = 864 \) subjects. Given about 15% possibility of incomplete data, we targeted total of \( N = 1,000 \) subjects. To have rural population representation in the study, we decided to recruit 20% (\( n = 200 \)) of aforementioned sample of \( N = 1,000 \) from two rural regions of Georgia described above. No information was collected on households that either refused to be enrolled or were unavailable to participate.

**Cognitive Testing**

All individuals in these households were evaluated using the Montreal Cognitive Assessment (MoCA; Nasreddine et al., 2005). The MoCA was previously translated into Georgian language and was validated showing its reliability and accuracy for evaluation of MCI (Janelidze et al., 2017).

MoCA cognitive domain index score (CDIS) was used to evaluate memory, executive function, visuospatial, language, attention, and orientation abnormalities (Julayanont, Brousseau, Chertkow, Phillips, & Nasreddine, 2014). CDIS was calculated as follows:

\[
\text{CDIS} = \left( \frac{\text{MIS} + \text{VIS} + \text{EIS} + \text{LIS}}{4} \right)
\]

The memory index score (MIS) was calculated by adding the number of words remembered in free delayed recall with a score ranging from 0 to 5.

The executive index score (EIS) was calculated by adding raw scores for the modified Trail-Making Test Part B, clock drawing, digit span forward and backward, letter A tapping, serial-7 subtraction, letter fluency, and abstraction, with a score ranging from 0 to 13.

The visuospatial index score (VIS) was determined by adding the raw scores of the cube copy, clock drawing, and naming, with a score ranging from 0 to 7.

The language index score (LIS) was obtained by adding the raw scores for naming, sentence repetition, and letter fluency, with a score ranging from 0 to 6.

The attention index score (AIS) was obtained by adding the raw scores for digit span forward and backward, letter A tapping, serial-7 subtraction, sentence repetition, and the words recalled in both immediate recall trials, with a score ranging from 0 to 18.

The orientation index score (OIS) was calculated as a sum of points for the orientation section of the MoCA, with a score ranging from 0 to 6 (Julayanont et al., 2014).

MIS, VIS, EIS, and LIS were used to categorize the MCI subtypes into single domain amnestic and nonamnestic, as well as multidomain amnestic (memory impairment plus one other impaired domain) and nonamnestic (Julayanont et al., 2014). Participants who scored less than 1.5 SD below the age- and education-adjusted mean.
value in MIS, VIS, EIS, and LIS were considered as being impaired in that cognitive domain.

**Definition of MCI.** We used International Working Group on MCI consensus criteria which defines MCI as follows: (a) the individual is neither normal nor demented (see the *Diagnostic and Statistical Manual of Mental Disorders* [4th ed.; *DSM-IV*; American Psychiatric Association, 1994] criteria); (b) there is evidence of cognitive deterioration, shown by objectively measured (>1.5 \( \text{SD} \) below mean value) decline; and (c) activities of daily life are preserved and complex instrumental functions are either intact or minimally impaired (Gauthier et al., 2006; Winblad et al., 2004).

For the multiple domain MCI type, we defined as deficits evidenced by scores >1.5 \( \text{SD} \) below mean values in more than one areas of cognitive functioning with or without memory impairment. Based on these criteria, MCI was divided in five categories: any MCI (MoCA >1.5 \( \text{SD} \) below mean value), amnestic, nonamnestic, multidomain amnestic, and multidomain nonamnestic.

We excluded individuals with MoCA < 16 from analysis classifying them moderately to severely impaired similar to Ganguli, Chang, Snitz, Saxton, Vanderbilt, & Lee (2010) study that excluded subjects with Mini Mental Status Examination (MMSE) < 21. This MoCA score threshold was chosen based on the evidence from Alzheimer’s Disease (AD) Imaging Initiative study, where MoCA cut-off score of 16 corresponded to MMSE score of 21 (Trzepacz, Hochstetler, Wang, Walker, & Saykin, 2015).

**Data Collection**

Demographic information was collected on each participant including: age, gender, level of education, and region (urban vs. rural). We also obtained information on the following medical conditions: smoking, hypertension (HTN), diabetes mellitus (DM), and hyperlipidemia (HLD). These medical conditions were reported by subjects and no medical record review or testing was performed. Only current smoking was documented.

Because of the fact that until recently, full school education in Georgia lasted 11 years, subjects were classified into two groups: (a) General (school graduation) with <12 years (11 years) of education (11 years); (b) Higher-with ≥12 years of education. Based on these, study subjects were asked simple question if they have general education (11 years) or higher education which means >12 years.

**Statistical Analysis**

Values were reported as mean (±SD), percentage, and the two-tailed \( p \) or Fisher’s exact two-sided tests were performed to compare the means or the distributions of variables as appropriate. One-way ANOVA was used for comparison of multiple groups. Spearman \( r \) correlation was used for correlations; \( p < .05 \) was considered statistically significant. SPSS version 22 and GraphPad Prism7 were used for statistical analysis.

**Results**

In all, 1,000 subjects were evaluated for the study, 149 were excluded for following reasons: \( n = 46 \) with MoCA < 16, \( n = 31 \) with incomplete data, \( n = 25 \) could not complete the test due to unknown reasons and \( n = 47 \) due to neurologic or psychiatric problems. In all, 851 subjects were enrolled in the study. Participants had a mean age of 56.5 ± 11.8 years, 63% were women, and 71% had ≥12 years of education. The rates were similar to the Georgian population >40 years of age, except for a higher rate of women (63.3% vs. 56%; Table 1). The prevalence of medical disorders among subjects included the following: HTN 26%, diabetes 4.6%, HLD 8.4%, and current smoking 23.6%.

The prevalence of any MCI was 13.3%. There was significantly higher prevalence of any MCI in males (odds ratio \( \text{OR} = 1.54, 95\% \text{CI} = [1.03, 2.27], p < .05 \)), older subjects with age >65 years (\( \text{OR} = 4.51, 95\% \text{CI} = [3.00, 6.75], p < .0001 \)) and general versus higher education group (\( \text{OR} = 3.99, 95\% \text{CI} = [2.66, 5.93], p < .0001 \); Table 2), as well as among those with HTN (\( \text{OR} = 2.51, 95\% \text{CI} = [1.45, 4.40], p < .0001 \).
Table 2. Prevalence of Different Types of MCI According to Demographics.

|                | aMCI   | naMCI  | mdaMCI | namdMCI | Any MCI |
|----------------|--------|--------|--------|---------|---------|
|                | n (%)  | n (%)  | n (%)  | n (%)   | n (%)   |
| All            |        |        |        |         |         |
| n = 851        | 79 (9.3) | 90 (10.5) | 23 (2.7) | 31 (3.6) | 114 (13.3) |
| Tbilisi (n = 719) | 64 (8.9) | 74 (10.2) | 18 (2.5) | 22 (3.0) | 87 (12.1) |
| Rural (n = 132)  | 15 (11.3) | 16 (12.1) | 5 (3.8)  | 9 (6.8)  | 27 (20.4) |
| OR (95% CI)     | 0.78 [0.41, 1.32] | 0.83 [0.47, 1.49] | 0.83 [0.25, 1.63] | 0.43 [0.19, 0.98] | 0.53 [0.33, 0.88] |
| Age ≥65 (n = 238) | 38 (16.0) | 29 (9.9)  | 8 (3.3)  | 12 (5.1) | 66 (27.7) |
| Age <65 (n = 613) | 41 (6.7)  | 61 (12.1) | 15 (2.4) | 19 (3.1) | 48 (7.8)  |
| OR (95% CI)     | 2.65 [1.66, 4.20] | 1.25 [0.79, 1.99] | 1.39 [0.61, 3.18] | 1.67 [0.81, 3.56] | 4.51 [3.00, 6.75] |
| Male (n = 312)  | 34 (10.9) | 29 (9.3)  | 11 (3.5) | 10 (3.2) | 52 (16.6) |
| Female (n = 539) | 45 (8.3)  | 61 (11.3) | 12 (2.2) | 21 (3.8) | 62 (11.5) |
| OR (95% CI)     | 1.34 [0.83, 2.13] | 0.80 [0.51, 1.26] | 1.60 [0.73, 3.74] | 0.82 [0.40, 1.76] | 1.54 [1.03, 2.27] |
| General education (n = 243) | 22 (9.0) | 40 (16.4) | 11 (4.5) | 17 (6.7) | 64 (26.3) |
| Higher education (n = 608) | 57 (9.3) | 50 (8.2)  | 12 (2.0) | 14 (2.3) | 50 (8.2)  |
| OR (95% CI)     | 0.96 [0.58, 1.60] | 2.20 [1.40, 3.40] | 2.35 [1.07, 5.51] | 3.19 [1.60, 6.68] | 3.99 [2.66, 5.93] |

Note. Any MCI = MoCA < 22. MCI = mild cognitive impairment; aMCI = amnestic MCI; naMCI = nonamnestic MCI; mdaMCI = multidomain MCI; namdMCI = nonamnestic multidomain MCI; OR = odds ratio; CI = confidence interval; MoCA = Montreal Cognitive Assessment.

*significant p < .05. †p < .01. ‡p < .001. §p < .0001. ‡‡p < .0001.
95% CI = [1.68, 3.76], p < .0001; Table 3), while residents of Tbilisi compared with rural population had lower (OR = 0.53, 95% CI = [0.33, 0.88], p < .05) prevalence of any MCI (Table 2).

The mean MoCA score of the study population was 25.2 ± 3.1. There was a statistically significant negative correlation between age and MoCA score (r = −.252, p < .0001), as well as age and CDISs including MIS (r = −.225, p < .0001), EIS (r = −.091, p = .008), LIS (r = −.215, p < .0001), VIS (r = −.089, p = .009), AIS (r = −.268, p < .0001), and a negative trend with OIS, although not significant (r = −.065, p = .058).

The prevalence of amnestic MCI (aMCI) was 9.3% (Table 2), and it was significantly higher with subjects older than 65 years, as well as those with diabetes (Table 2). On the contrary, smokers had significantly lower prevalence of aMCI (Table 3). Smokers were significantly younger (54.6 ± 11.1 vs. 57.1 ± 11.9, p = .008), but there was no difference in proportion of subjects with degree between smokers and nonsmokers (66.3% vs. 72.0%, p = .77).

Nonamnestic MCI (naMCI) was documented in 10.5% of subjects with significantly higher prevalence in high school nondegree graduates (Table 2). Multidomain amnestic MCI (mdaMCI) was observed in 2.7% of subjects with significantly higher prevalence among high school nondegree graduates (Table 2). Finally, nonamnestic multidomain MCI (namdMCI) prevalence was 3.4% with significant increase in high school nondegree graduates and significant decrease among Tbilisi residents (Table 2).

Figure 1 shows prevalence of MCI types according different age groups. There was significant increase in prevalence of all types of MCIs according age groups except naMCIs including both single and multidomain (Figure 1).

Discussion

In this population-based, cross-sectional study of MCI in Republic of Georgia the prevalence of MCI was 13.3%. This is similar to 16.0% reported by Petersen et al. (2010), 17.7% observed by Ganguli et al. (2010), 18.8% by Lopez et al. (2003; all in U.S. cohorts), and 19.4% by Busse, Hensel, Gühne, Angermeyer, & Riedel-Heller (2006) in Germany. However, 13.3% reported here is much lower compared with 42.0% documented in France (Artero et al., 2008), 28.3% in Northern Manhattan (Manly et al., 2005), and 39.1% in Australia (Sachdev et al., 2012). On the contrary, our finding is almost twice higher compared with 6.3% found in some U.S. studies, 6.5% in Finland, and 5.1% in Germany (Busse et al., 2003; Ganguli et al., 2004; Hänninen, Hallikainen, Tuomainen, Vanhanen, & Soininen, 2002). There are several explanations for varying estimates of MCI prevalence reported in these studies including age, MCI criteria used, and cut-points for abnormality for neuropsychological test scores (Rosebud & Knopman, 2013). In the current study, among older subjects >65 years the overall prevalence of MCI was 27.7%, which is similar to studies from U.S. reporting 22.2% and 23.4% (Fisk, Merry, & Rockwood, 2003; Plassman et al., 2008; Unverzagt et al., 2001).

We documented aMCI in 9.9% of studied subjects, which is comparable with the prevalence of 14.9% reported in Kolkata, India (Das et al., 2007). Higher prevalence of aMCI in India most likely reflects older age, as mean age of their cohort was 10 years older compared with the current study. Lower prevalence of aMCI ranging from 2.4% to 5.3% was reported in population-based cohorts from Canada and Finland (Fisk et al., 2003; Hänninen et al., 2002). In Finnish study, it is possible that exclusion of subjects older than 76 years resulted an underestimation of the prevalence, while Canadian study used more strict aMCI criteria that might yield relatively lower population prevalence estimate (Fisk et al., 2003; Hänninen et al., 2002).

In this study, multiple domain MCIs including amnestic and nonamnestic ranged from 3.3% to 5.3%. Similar to aMCI, these rates were comparable with 8.8% reported from Kolkata, India (Das et al., 2007). As in aMCI, slightly lower prevalence of multiple domain MCIs in the current study most likely is a reflection of younger age.

Age is the strongest associated factor with MCI (Busse et al., 2006; Das et al., 2007; Ganguli et al., 2010, 2011; Hänninen et al., 2002; Trzepacz et al., 2015; Unverzagt et al., 2001). In the current study, age >65 years was associated with 4.5-fold increase in any MCI prevalence (Table 2), and each 10-year increase in age was associated with statistically significant increase in prevalence of any and aMCIs including single and multidomain (Figure 1). In addition, age negatively correlated with CDIS scores including memory, language, executive, visual, and attention scores, while orientation was nearly significant.

This study found statistically significant 1.5-fold increase in prevalence of any MCI among males compared with female subjects. This is similar to studies from India, Finland, and the United States (Das et al., 2007; Hänninen et al., 2002; Petersen et al., 2010; Plassman et al., 2008) but opposite to report of Luck et al. (2007) showing 1.4-fold increase of MCI among male residents (Table 2). Most likely explanation of this finding is the fact that proportion of subjects with higher education in Tbilisi was more than three-fold higher compared with rural regions (Table 1). However, those with general education compared with
### Table 3. Prevalence of Different Types of MCI According Risk Factors.

| Risk Factor | n (%) | aMCI | naMCI | mdsMCI | namdMCI | Any MCI |
|-------------|-------|------|-------|--------|---------|---------|
|             |       | n (%) | n (%) | n (%)  | n (%)   | n (%)   |
| HTN+ (n = 225) | 28 (12.4) | 29 (12.8) | 9 (4.0) | 10 (4.4) | 50 (22.2) |
| HTN– (n = 626) | 51 (8.1) | 61 (9.7) | 14 (2.2) | 21 (3.3) | 64 (10.2) |
| OR (95% CI) | 1.60 [0.97, 2.58] | 1.37 [0.86, 2.18] | 1.84 [0.75, 4.19] | 1.34 [0.65, 2.91] | 2.51 [1.68, 3.76] |
| DM+ (n = 39) | 8 (20.5) | 2 (5.1) | 2 (5.1) | 3 (7.6) | 9 (23.1) |
| DM– (n = 812) | 71 (8.7) | 88 (10.8) | 21 (2.6) | 28 (3.4) | 105 (12.9) |
| OR (95% CI) | 2.69 [1.25, 5.98] | 0.44 [0.10, 1.71] | 2.04 [0.46, 8.31] | 2.33 [0.72, 7.11] | 2.02 [0.93, 4.22] |
| HLD+ (n = 65) | 10 (15.4) | 3 (4.6) | 2 (3.1) | 0 (0.0) | 10 (15.3) |
| HLD– (n = 786) | 69 (8.8) | 87 (11.1) | 21 (2.7) | 31 (4.0) | 104 (13.2) |
| OR (95% CI) | 1.89 [0.93, 3.72] | 0.39 [0.13, 1.19] | 1.16 [0.26, 4.52] | 0.00 [0.00, 1.28] | 1.35 [0.66, 2.70] |
| Smoking+ (n = 201) | 8 (4.0) | 18 (8.9) | 4 (2.0) | 4 (2.0) | 22 (10.9) |
| Smoking– (n = 650) | 71 (10.9) | 72 (11.1) | 19 (2.9) | 27 (4.1) | 92 (14.1) |
| OR (95% CI) | 0.34 [0.16, 0.70] | 0.79 [0.46, 1.35] | 0.67 [0.24, 1.92] | 0.47 [0.17, 1.32] | 0.74 [0.45, 1.21] |

Note. Any MCI = MoCA < 22; MCI = mild cognitive impairment; aMCI = amnestic MCI; naMCI = nonamnestic MCI; mdsMCI = multidomain MCI; namdMCI = nonamnestic multidomain MCI; HTN = hypertension; OR = odds ratio; CI = confidence interval; DM = diabetes mellitus; HLD = hyperlipidemia; MoCA = Montreal Cognitive Assessment.‡ nonsignificant p < .1. *p < .001. †p < .001. ‡p < .01. †p < .05.
higher education cohort had significantly higher prevalence of all types of MCI except aMCI (Table 2). A protective effect of education on cognitive function has been noted in other studies as well (Fisk et al., 2003; Hänninen et al., 2002).

In this study, smoking was associated with 66% lower prevalence of aMCI. Earlier case-control studies also reported reduced risk of AD among smokers (Lee, 1994). The most likely explanation of smoking paradox reported in current study is the fact that smokers were significantly younger. Subsequent cohort study examining smoking in midlife, have found that smoking is a risk factor for AD (Ott et al., 1998).

We found statistically significant more than two-fold higher prevalence of any MCI in subjects with HTN and diabetes (Table 3). In addition, aMCI was strongly associated with diabetes, but not with HTN (Table 3). Several studies have identified association of vascular risk factors with MCI and their role in progression of MCI to dementia (DeCarli et al., 2001; Di Carlo et al., 2000; Kivipelto et al., 2001; Solfrizzi et al., 2004). In a recent positron emission tomography study of individuals without dementia from three U.S. communities, a cumulative number of midlife vascular risk factors was associated with elevated brain amyloid deposition suggesting a role of vascular disease in the development of AD (Gottesman et al., 2017).

In the current study, mean age, proportion of subjects in age subgroups (10-year increments), as well as proportion of individuals with general and/or higher education was comparable with the Georgian population older than 40 years (Table 1). The female prevalence of 63% reported in this study was higher than 56% documented among Georgian population of corresponding age older than 40 years in Georgian Census (2014). However, female prevalence of 50.0% in rural community of studied participants was compatible to 47.7% in rural regions of Georgian population reported in Results of General Population Census of Georgia (2014).

Although 23.6% prevalence of smoking in this study was compatible to previously reported 27.7% (Grim et al., 1999), prevalences of other vascular risk factors were much lower in this study including HTN (26.0% vs. 56%), DM (4.6% vs. 10.0%), and HLD (31.0% vs. 7.6%). However, Grim et al. (1999) study was conducted almost 20 years ago, and recent meta-analysis of HTN in low- and middle-income countries shows 31.5% prevalence in Europe and Central Asia, which is comparable with 26.0% reported here (Sarki, Nduka, Stranges, Kandala, & Uthman, 2015). In addition, prevalence of DM in Georgia recently was reported to be 2.2%, which is close to observed 4.6% in this study, while HLD prevalence of 8.7% is exactly the same as documented here (Wilkins et al., 2017).

As for the large-scale epidemiological study of the MCI complete neuropsychological testing in the field is not feasible, we used MoCA with its CDIS (Julayanont et al., 2014; Nasreddine et al., 2005). The Georgian version of MoCA was previously validated showing reliability and accuracy of this test for evaluation of MCI (Janelidze et al., 2017). In that study, MoCA < 22 was optimal to detect MCI with 100% sensitivity and 69% specificity (Janelidze et al., 2017). In the current study, based on 1.5 SD below normative values, the same cutoff MoCA < 22 was found to be an optimal threshold for MCI. Lam et al. (2013) have shown that MoCA is a valid tool for assessment of cognition that shows good agreement with existing neuropsychological screening tests and global measures. In addition, MoCA subscores for different cognitive domains also demonstrated validity when compared with neuropsychological testing–derived measures (Lam et al., 2013). In that study, in the case of memory, executive, and visuospatial dysfunction, the MoCA’s subscores were reasonable screens for domain-specific impairment (Lam et al., 2013). In another study, individuals with MCI with a low MoCA MIS score were at high risk of conversion to AD (Julayanont et al., 2014).

The study has some limitations. One possible limitation is inclusion of subjects with relatively younger age >40 years. We considered this age cut-off because of previously reported high prevalence of vascular risk factors in a similar age group (mean age of 55 years) of Georgian population with a prevalence of HTN 56%, DM 10%, and HLD 31% (Grim et al., 1999). Another potential shortcoming of this study which is a common problem with cross-sectional studies is nonresponse and lack of the data on nonresponse rate. We cannot completely rule out possibility that those who refuse to participate are more impaired than those who agree to participate. The lack of information on nonresponders and those that refused to participate might create possible bias. However, given the fact that demographic profile of the studied population is comparable with Georgian population of the same age, sampling biases are less likely and the results are generalizable to whole Georgian population. Representability of...
rural community in the study can be potential methodologi-
cal shortcoming. As 20% rate of rural residence is lower
than 42% reported in recent 2014 census, this might be
potential limitation of the study. However, this sample size
allowed appropriate subgroups analysis. Another potential
limitation is the selection of rural sites for the study. The
rate of MCI was lower in Tbilisi as compared with rural
regions. The two rural regions were selected for the study
were based on study investigators, geographic region (east
and west), and population. Each region was defined as a
rural settlement based on a 2014 census (Results of
General Population Census of Georgia, 2014). We regis-
tered cardiovascular comorbidities based on survey with-
out confirmation in medical or pharmacy records.
However, in a recent study, self-reported diagnosis sensi-
tivity for HTN was 83%, for diabetes 73%, and for hyper-
cholesterolemia 59% while specificity was >80% for all
three conditions indicating that self-reports are reasonably
accurate for certain chronic conditions and can provide a
useful estimate for broad measures of population preva-
ience (Martin, Leff, Calonge, Garrett, & Nelson, 2000).
In conclusion, in this population-based cross-sectional
study, prevalence of MCI in Georgia was 13.3%, and it was associated with advanced age, male gender,
rural residence, lower education, HTN, and diabetes.
These findings have significant implications not only for
better understanding of MCI profile in Georgia, but also
for public health planning in this country as aggressive
vascular risk factor control intervention potentially may
prevent progression of MCI to dementia.

Author Contributions
All authors made substantial contributions to the conception
and design of the study and analysis and interpretation of data.
Marina Janelidze: study concept and design, contributed to
drafting the paper and revising it critically for intellectual
content; Nino Mikeladze: data acquisition, analysis and interpreta-
tion of data, contributed to drafting the paper and revising it
critically for intellectual content; Nazibrola Bochorishvili:
data acquisition, contributed to drafting the paper and revising it
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tion, contributed to drafting the paper and revising it critically for
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contributed to drafting the paper and revising it critically for
intellectual content; Ekaterina Mirvelashvili: study design,
statistical analysis of the data; Nino Shikashvili: data collec-
tion, statistical analysis of the data; John K. Lynch: design,
analysis and interpretation of data and revising the manuscript
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concept and design, analysis and interpretation of data, draft-
ing the paper and revising it critically for intellectual content,
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