Neurocognitive testing commonly uses the MMSE (Mini–Mental State Examination) to evaluate the overall cognitive function of patients at outpatient clinics, but the MMSE has recently been extensively used in the SNSB II (Seoul Neuropsychological Screening Battery II) for making diagnoses. We retrospectively investigated the results of routine neurocognitive tests and the results of the blood tests of 120 elderly patients who had been referred to a South Central Medical Center from 2017 to 2018 and who had been examined at a public health center. These subjects’ space-time capability was high on the sub-region of the global deterioration scale (GDS). GDS showed a significant increase as the Na decreased on the electrolyte analysis. The subjects’ concentration, their language-based orientation for space and time, their memory, and their scores for the frontal lobe function on GDS showed statistically significant reductions ($P < 0.001$) for the normal and abnormal groups according to the ALT and creatinine levels. The frontal/execute function areas showed statistically significant differences ($P < 0.001$) as well as negative correlation between GDS and ALT ($P < 0.01$). In conclusion, this study provides basic information to develop test items that are important for patient screening and diagnosis, and several routine blood chemistry factors provide basic information for diagnosing and assessing the status and progress of cognitively impaired patients.
and instructions, consideration of the physical and emotional state of the patient is essential for performing and interpreting the neuropsychological test. There are so many neuropsychological tests that can not be counted, but Seoul Neuropsychological Screening Battery (SNSB) is the most commonly used test in Korea.

A comprehensive and in-depth assessment of cognitive function provides useful information on early diagnosis of dementia and causative disease. SNSB was developed for the purpose of assessing the efficacy of therapeutic agents. The test items include five cognitive domain tests, as well as Korean version of Mini–Mental State Examination (K-MMSE), geriatric depression scale [1] and barthel index. K-MMSE reported a satisfactory sensitivity (70.3 ∼ 82.7%) and specificity (91.3%) when applied to a 23-point cut-off point in the validity study of hospital dementia patients [2, 3]. K-MMSE has been mainly used in clinical settings, and is sometimes used as a screening test for the prevalence of dementia led by neurologists.

When compared with comprehensive neuropsychological tests such as the SNSB, data from K-MMSE tests that applied the criteria according to education and age derived false negative rate results over 19.7% [4]. Several previous studies including the SNSB, identified that the MMSE did not adequately evaluate executive function [5].

Recently, high-density lipoprotein cholesterol (HDL-C) levels in midlife are inversely associated with both late-life mild cognitive impairment (MCI) and dementia. Midlife HDL-C thus has the potential to be used as a marker of late-life cognitive impairment [6].

Presbycusis is the general term applied to ARHL (age-related hearing loss). The risk factors for presbycusis include noise exposure, smoking, medication, hypertension, family history and other factors [7]. Dementia is a chronic and progressive deterioration disease characterized by cognitive dysfunction and abnormal mental behavior. It has become the greatest global challenge for health and social care in the 21st century [8].

Also, Vitamin B complex acts on the supply and metabolism of energy. Vitamin B12 and B9 (folic acid) are closely related to the onset and prevention of dementia genetic tests show that family dementia tends to be relic and that there are three genes that cause it, but it accounts for 5% to 10% of all dementia. If the family history of dementia appears, an assessment of hereditary dementia is needed. Sporadic Alzheimer’s disease, which accounts for the rest of the disease, causes dementia by compounding multiple risk perceptions. Among them, having a single Apo protein type 4 genes, a risk factor that causes sporadic Alzheimer’s disease, causes 2 ∼ 3 times the risk of dementia than those without genes, having two increases the risk by more than 10 times. It is difficult to diagnose dementia with one Apo protein gene. However, it provides clinical information on the progression of dementia conversion rate or dementia in mild cognitive impairment.

The literature on minerals and cognitive impairment associated with dementia is limited. Previous reported that increased dietary intake of potassium, calcium, and magnesium reduced the risk of all-cause dementia especially vascular dementia. The risk of AD tended to decrease with higher reported dietary mineral intake, but there was no significant linear progression [9]. The significant interaction between fasting glucose and sex suggested fasting glucose levels were deleterious to short-term verbal memory performance, particularly among men [10].

The few tests that have merit as broader screening tests in asymptomatic patients include serum glucose, blood urea nitrogen (BUN), creatinine, and urinalysis. Patients on psychotropic medications should be monitored for side effects of that particular therapy. Further prospective data are needed to develop cost-efficient, population-specific diagnostic strategies [11, 12].

Recently reported studies have been conducted to estimate the stage of disease by finding biomarkers through blood tests in cognitive dysfunction groups such as dementia and Alzheimer’s disease [7-10]. However, blood analysis of biomarkers is rarely performed in patients with cognitive dysfunction. Therefore, we conducted the study from the early 90’s [11, 12] to investigate the relationship between routine blood chemistry test and GDS level. Ultimately, routine blood
chemistry factors provide basic information about the diagnosis and progress of cognitive impairment subjects.

**MATERIALS AND METHODS**

1. **Subjects**

The subjects' personal information was not collected and retrospectively investigated through routine blood chemistry results and GDS. In the results (2017~2018 years), Among the 120 patients who first visited the neurologist for cognitive function test and measured the GDS and performed routine blood chemistry tests. This study is a retrospective study. Subjects were the elderly, including the elderly who were referred to the hospital from the public health center to the hospital. This study has been conducted according to the principles expressed in the Declaration of Helsinki (approved and exemption by Institutional Review Board No. 1041479-HR-201805-019 at Namseoul University).

2. **Analysis of routine blood chemistry**

This was a retrospective cohort study conducted on data from the total cholesterol (TC), total bilirubin (TB), uric acid (UA), aspartate aminotransferase (AST), alanine aminotransferase (ALT), blood urea nitrogen (BUN), creatinine (Cr), sodium (Na), potassium (K), chloride (Cl), in Gyeongi province medical center Ansung hospital (2017), which is the most recent data that measured. Laboratory values determined using standard laboratory techniques on a TBA-2000FR (Toshiba LTD., Tokyo, Japan) The data was a randomized survey to conducted by outpatients in Gyeongi province medical center Ansung hospital.

3. **Seoul neuropsychological scanning battery**

The SNSB-II evaluates four cognitive factors such as memory, language skills, space-time functions, and executive function. The evaluation of each cognitive domain conformed to the criteria of SNSB-II. Memory was evaluated by delayed recall using Seoul Verbal Learning Test (SVLT), and the Rey Complex Figure Test (RCFT). Language skills were assessed by naming Korean-Boston Naming Test (K-BNT), and time and space function was evaluated using the RCFT. Executive function was evaluated using a variety of tests, including the motor control and perseveration test (motor impersistance, contrasting program, go-no-go, fist-edge-palm, alternating hand movement, alternating square and triangle, and Luria loop) the controlled word association test (COWAT) and the stroop color reading forward reaction of Korean-Color Word Stroop Test (KCWST).

The SNSB has been modified and complemented to produce the SNSB-II. The SNSB-II expands the age criteria to 45 years old to 90 years old from its previous range of 55 years to 80 years of age.

For SNSB-II assessment, area-specific disability criteria were defined as follows, in accordance with the standards of cognitive dysfunction defined by SNSB-II.

(1) Memory: at least one is abnormal in A and B. A (Verbal memory): SVLT delayed recall scores are abnormal (<15%). B (Visual memory): RCFT delayed recall scores are abnormal (<15%).

(2) Language skills: K-BNT (full) is abnormal (<15%).

(3) Space-time function: the RCFT copy is abnormal (<15%).

(4) Executive dysfunction: two or more are abnormal in A, B, and C.

A (Motor examination): three kinds or more are abnormal in the following tests: motor impersistance, contrasting program, go-no-go, fist-edge-palm, alternating hand movement, alternating square & triangle, and Luria loop.

B (COWAT): at least one or more is abnormal in the following three tests: animal, supermarket, and Hangul consonants.

C: Stroop color reading forward reaction is abnormal (<15%).

The assessment methods for global deterioration scale (GDS) cognitive impairment include GDS 1 (no cognitive impairment), GDS 2 (very mild cognitive impairment), GDS 3 (minor cognitive impairment), GDS 4 (middle cognitive impairment), GDS 5 (initial cognitive impairment),
GDS 6 (severe cognitive impairment) and GDS 7 (the last stage severe cognitive impairment)[13]. GDS is accurately reflecting the progression of the disorder of the primary degenerative dementia. Not influenced by educational, cultural, socioeconomic, and other biases.

4. Statistical analysis

The level of statistical significance was defined as having a P-value of less than 0.05 or 0.01. Data were analyzed using PAWS version 22.0 (SPSS Inc., Chicago, IL, USA). The results of score and each blood chemistry analysis value by region of neurocognitive test were calculated by average and standard deviation. The analysis was conducted by t-test and ANOVA (Scheffe Post Hoc test) according to the distribution of neurocognitive test, seasonality, and blood chemistry test factors (including electrolytes). The correlation between blood chemical factors, seasonality, and blood chemistry test factors was analyzed as Pearson Correlation Coefficient. We used the parametric method to identify correlations between the indicators.

RESULTS

1. Distribution of characteristics parameters in clinical patients

The frequency of characteristics parameters of the participants are shown in Table 1. The distribution of total cholesterol was 28 (23.3%) out of reference, and 46 (38.3%) was also distributed. Most of the remaining items were less than 10%. GDS ratings were the highest at 3.0 with 45 (37.5%) and 5.0, 4.0, 2.0 and 6.0.

2. Correlation between blood test factors and GDS grade

The changes in blood test factors according to the GDS grade showed that the GDS showed a significant decline in GDS as the GDS increased (P<0.05), and AST and ALT

| Characteristic | Variables | N  | %  | Reference value | Mean±SD  |
|---------------|-----------|----|----|-----------------|---------|
| TC (mg/dL)    | Abnormal  | 28 | 23.3| <200            | 190.0±38.6|
| Normal        | 92        | 76.7|     |                 |         |
| TB (mg/dL)    | Abnormal  | 2  | 1.7 | <1.3            | 0.6±0.2 |
| Normal        | 118       | 98.3|     |                 |         |
| UA (mg/dL)    | Abnormal  | 14 | 11.7| 2.4~7.0         | 4.8±1.5 |
| Normal        | 106       | 88.3|     |                 |         |
| AST (U/L)     | Abnormal  | 8  | 6.7 | <40             | 24.9±8.8|
| Normal        | 112       | 93.3|     |                 |         |
| ALT (U/L)     | Abnormal  | 6  | 5.0 | <40             | 18.9±10.9|
| Normal        | 114       | 95.0|     |                 |         |
| BUN (mg/dL)   | Abnormal  | 10 | 8.3 | 6.0~20.0        | 15.9±5.6|
| Normal        | 110       | 91.7|     |                 |         |
| Cr (mg/dL)    | Abnormal  | 46 | 38.3| 0.50~1.30       | 0.9±0.4 |
| Normal        | 74        | 61.7|     |                 |         |
| Na (mmol/L)   | Abnormal  | 2  | 1.7 | 136~145         | 138.9±2.7|
| Normal        | 118       | 98.3|     |                 |         |
| K (mmol/L)    | Abnormal  | 3  | 2.5 | 3.5~5.5         | 4.2±0.4 |
| Normal        | 117       | 97.5|     |                 |         |
| Cl (mmol/L)   | Abnormal  | 4  | 3.3 | 98~107          | 103.2±3.1|
| Normal        | 116       | 96.7|     |                 |         |
| GDS (grade)   | 2.0       | 8  | 6.7 |                 | 3.85±1.0|
|               | 3.0       | 45 | 37.5|                 |         |
|               | 4.0       | 27 | 22.5|                 |         |
|               | 5.0       | 37 | 30.8|                 |         |
|               | 6.0       | 3  | 2.5 |                 |         |

Abbreviations: TC, total cholesterol; TB, total bilirubin; UA, uric acid; AST, aspartate aminotransferase; ALT, alanine aminotransferase; BUN, blood urea nitrogen; Cr, creatinine; Na, sodium; K, potassium; Cl, chloride; GDS, global deterioration scale.
showed a significant decrease in GDS 4.0 group compared to 5.0 group in the total Bilirubin group. Creatinine also significantly decreased levels of GDS 3.0, 4.0, and 5.0 to 6.0 (P<0.05). In the analysis of electrolytes, the levels of Na from 2.0 to 5.0 decreased from 6.0 (P<0.05), and the levels of K from 2.0 to 5.0 increased from 6.0 (P<0.05). Overall, the one-way analysis showed statistically significant distribution of the GDS grade (P<0.05) (Table 2).

3. Correlation of GDS subcategory factors according to GDS grade

The changes in sub-domains according to the GDS grade showed statistical significance in all sub-domains (P<0.001), significantly lower in concentration at GDS 5.0 and 6.0, and lower in language than GDS 2.0 at 3.0, 4.0 and 5.0. Visuospatial function was significantly lower than GDS 2.0 and 3.0 at 4.0, 5.0, and 6.0, and memory and frontal/executive function decreased as GDS levels increased. They were all statistically significant (P<0.05) (Table 3).

4. Difference of blood factors according to GDS subcategory

The association of blood factors with GDS subcategory showed statistical differences in the levels of visual function in the normal and abnormal groups of total cholesterol (P<0.05). In addition, statistical significance was shown in the frontal/execute function areas in the summit and abnormal groups of ALT and Creatinine (P<0.001, P=0.019). The electrolyte factor has a significantly smaller population of abnormal groups, but it has statistically significance in Na, Cl, language, visuospatial function, and frontal/executive function (P<0.05) (Table 4).

5. Correlation of blood factors and GDS

Table 5 showed the significant correlation between

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### Table 2. Difference of parameters in according to the level of GDS

| Variables | GDS grade (N) | Schéffe | F   | P   |
|-----------|--------------|---------|-----|-----|
|           | 2.0* (8) | 3.0* (45) | 4.0* (27) | 5.0* (37) | 6.0* (3) |
| TC        | 199.1±40.0 | 192.9±40.9 | 187.9±40.2 | 187.7±32.3 | 169.0±69.3 | 0.440 | 0.779 |
| TB        | 0.5±0.1   | 0.6±0.2   | 0.6±0.2   | 0.5±0.2   | 0.6±0.1   | c>d  | 1.345 | 0.258 |
| UA        | 4.3±0.5   | 4.9±1.5   | 4.9±1.7   | 4.6±1.5   | 3.7±1.3   | a>d  | 0.875 | 0.481 |
| ALT       | 27.0±14.8 | 20.6±11.8 | 18.2±8.5  | 16.1±9.8  | 13.6±5.7  | a>c>d| 2.277 | 0.065 |
| BUN       | 15.2±3.6  | 15.2±4.9  | 16.5±5.9  | 16.5±6.7  | 12.6±1.5  | a>b  | 0.617 | 0.651 |
| Cr        | 0.8±0.1   | 0.9±0.2   | 1.0±0.4   | 0.7±0.0   | b>e, c>d, e>d | 0.979 | 0.422 |
| Na        | 139.1±1.9 | 139.4±2.0 | 139.4±2.8 | 138.4±2.4 | 133.3±7.3 | a>e, b>e, c>d, e>d | 4.665 | 0.002* |
| Cl        | 103.5±1.4 | 103.4±2.3 | 103.1±3.8 | 103.4±3.2 | 98.3±7.0  | a>e, b>e, c>d, e>d | 0.319 | 0.865 |

Abbreviations: See Table 1.
*P<0.05.

### Table 3. Difference of GDS subcategory in according to the level of GDS

| Variables | GDS grade (N) | Schéffe | F   | P   |
|-----------|--------------|---------|-----|-----|
|           | 2.0* (8) | 3.0* (45) | 4.0* (27) | 5.0* (37) | 6.0* (3) |
| Attention | 27.9±28.6 | 29.4±22.7 | 14.9±18.2 | 9.8±11.3 | 6.5±11.0 | a>d, b>c>d, c>e | 6.530 | 0.000* |
| Language  | 52.2±27.6 | 48.2±23.7 | 32.2±28.2 | 25.2±30.3 | 33.1±46.8 | a>d, b>c>d | 4.342 | 0.003* |
| VF        | 46.7±34.3 | 34.1±32.7 | 9.0±16.1  | 6.1±15.4  | 0.7±1.1   | a>c,d b,d,e | 10.648 | 0.000* |
| Memory    | 45.9±17.2 | 14.2±19.8 | 9.3±9.9   | 5.4±7.4   | 0.3±0.4   | a>b,c,d,e b,d> | 13.993 | 0.000* |
| FE        | 33.0±27.4 | 17.3±21.5 | 7.3±19.8  | 3.6±9.5   | 0.03±0.05 | a>b,c,d,e | 6.151 | 0.000* |

Abbreviations: VF, Visuospatial function; FE, Frontal/Executive function.
*P<0.05.
Table 4. Difference of GDS subcategory in accordance to the level of parameters (abnormal/normal)

| Variables | Attention | Language | VF | Memory | FE |
|-----------|-----------|----------|----|--------|----|
| TC Abnormal | 25.2±23.5 | 41.4±25.6 | 29.8±29.7 | 13.6±18.6 | 11.5±19.0 |
| Normal | 17.7±19.6 | 36.2±30.2 | 16.8±27.6 | 11.7±16.9 | 11.4±20.3 |
| TB Abnormal | 2.126 (0.097) | 2.011 (0.407) | 2.328 (0.035*) | 0.602 (0.616) | 0.048 (0.975) |
| Normal | 19.7±20.9 | 37.5±29.4 | 20.0±28.7 | 12.2±17.3 | 11.4±20.1 |
| UA Abnormal | 1.767 (0.341) | 3.950 (0.508) | 0.853 (0.749) | 0.696 (0.660) | 0.071 (0.973) |
| Normal | 24.6±26.9 | 27.2±23.8 | 15.7±18.5 | 7.2±8.7 | 10.1±17.4 |
| AST Abnormal | 3.104 (0.330) | 3.147 (0.167) | 5.516 (0.065) | 1.9±3.0 | 0.136 (0.795) |
| Normal | 19.9±20.9 | 37.1±29.3 | 19.8±28.6 | 12.0±17.2 | 10.9±19.1 |
| ALT Abnormal | 17.7±16.7 | 32.7±22.9 | 21.9±29.0 | 9.1±11.3 | 1.9±3.0 |
| Normal | 19.5±21.0 | 37.6±29.5 | 19.8±28.6 | 12.3±17.5 | 11.9±20.3 |
| BUN Abnormal | 0.026 (0.309) | 0.534 (0.722) | 0.494 (0.962) | 0.05±0.07 | 2.247 (0.298) |
| Normal | 19.4±20.7 | 36.6±29.9 | 14.0±22.3 | 8.5±11.4 | 7.3±17.6 |
| Cr Abnormal | 0.705 (0.019) | 3.147 (0.167) | 1.749 (0.999) | 1.553 (0.492) | 0.791 (0.500) |
| Normal | 19.6±22.6 | 35.6±29.9 | 14.0±22.3 | 8.5±11.4 | 7.3±17.6 |
| Na Abnormal | 2.094 (0.199) | 2.666 (0.880) | 9.420 (0.202) | 6.146 (0.107) | 12.134 (0.019)* |
| Normal | 11.8±13.5 | 0.3±0.3 | 0.1±0.0 | 3.1±4.2 | 0.05±0.07 |
| K Abnormal | 19.6±20.9 | 38.0±29.0 | 20.2±28.6 | 12.3±17.3 | 11.6±20.1 |
| Normal | 2.384 (0.604) | 5.905 (0.000*) | 4.974 (0.000*) | 1.563 (0.456) | 2.260 (0.417) |
| CI Abnormal | 0.901e-04 (0.199) | 1.702 (0.787) | 2.820 (0.477) | 2.774 (0.368) | 2.807 (0.363) |
| Normal | 9.7±10.7 | 51.1±41.5 | 0.6±0.9 | 6.9±12.7 | 0.3±0.4 |

Abbreviations: See Table 1, VF, Visuospatial function; FE, Frontal/Executive function.
*P<0.05.

blood factors and GDS. In the correlation between blood factors and GDS, BUN and Creatinine showed negative correlation to TB (P<0.05), and uric acid showed positive correlation (P<0.01). The Uric acid also showed positive correlations with potassium (P<0.05), and GDS and ALT showed negative correlation (P<0.01).

**DISCUSSION**

The appropriate cut-off score and validity for the screening of dementia were presented based on the results of clinical studies. However, it is not calculated by surveying the residents of the community. Therefore, these tools should be used appropriately for the purpose of screening dementia. It is necessary to verify the validity of the questionnaire to the residents of the community.

The pathology of dementia, especially AD, which is focused on in the following section, includes: (1) loss of neurons in the temporal lobes and hippocampus; (2) NFTs; (3) SPs that consist of amyloid-β (Aβ); and (4) amyloidopathy of the cerebrovasculature [14].

Increased β-amyloid load mediated by the APOE ε4 allele may lead to an increase in a-synuclein aggregation and toxicity via stabilization in the formation of hybrid nanopores [15] or through a direct connection between intracellular a-synuclein and extracellular APOE and β-amyloid [16].

A typical rating scale for patients with dementia is

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[14] [Link to the reference]
[15] [Link to the reference]
[16] [Link to the reference]
Clinical Dementia Rating (CDR) scale [17] and Global Deterioration Scale (GDS) [18]. GDS and CDR are the criteria for the severity of dementia in clinical studies. Clinical trials are the most widely used criteria for evaluating the efficacy of dementia drugs. GDS is a detailed classification of early cognitive impairments, so GDS can be used as a test tool for clinical trials of dementia treatments or early diagnosis of dementia.

Brain-derived biomarkers are typically present at relatively low concentrations in the blood because of the blood–brain barrier preventing free passage of molecules, targeted blood-based biomarkers (plasma Aβ, described in the AlzBiomarker database (https://www.alzforum.org/alzbiomarker), plasma tau, plasma neurofilament light, hsa-miR-125b, plasma Aβ42/Aβ40 ratio [19]). Both low and high serum magnesium levels are associated with an increased risk of all-cause dementia [20].

Recently, experiments have also shown that their expression levels are altered in both presbycusis and AD mouse models. Therefore, we propose that exploring the specific molecular link between presbycusis and AD may provide new ideas for their prevention and treatment [21].

A strength of this study is that the measures used with both groups were very similar, facilitating comparisons. Limitations of this study include a small sample size and data that are cross-sectional in nature. Additionally, although many investigators stress the role of diet in regulating potassium levels, information was not available for these participants.

Association between sUA (serum uric acid) and dementia/cognitive impairment was weak across undifferentiated dementia groups [22]. Higher homocystein levels, lower educational attainment, and decreased physical activity were particularly strong predictors of incident AD [23]. Higher levels of the minerals iron and copper were identified as being related to increased risk for AD but the role of other minerals (such as potassium) in cognitive decline was not addressed. Different lab abnormalities at the time of peak metabolic derangement accounted for unique patterns of neuropsychological impairment.

Recently, they found that a higher potassium level in MCI participants was predictive of MCI status. These findings contradict our premise of lower potassium levels...
that are related to MCI status. A plausible explanation may be the different socioeconomic status and biological components in our population [24].

Various risk or protective factors for dementia syndrome or cognitive impairment have been suggested by numerous epidemiological studies [25]. However, we still do not know much about the mechanisms through which various lifetime experiences such as physical and mental activity, social relationships, stress experiences, personality, diet and nutrition, and sleep, and many aging-related physiological or pathological changes in the human body, such as vascular status, hormonal changes, glucose or lipid metabolism, and body mass index (BMI).

In future, proteomical factors, biochemical and genetic analyses using blood and DNA specimen are also ongoing. Additionally, external validity of candidate biomarkers will be tested by using the data and samples collected from an independent cohort that is established for validation of candidate biomarkers discovered in cognitive impairments.

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