Central blood pressure lowering effect of telmisartan-rosuvastatin single-pill combination in hypertensive patients combined with dyslipidemia: A pilot study

JungMin Choi MD | Ki-Chul Sung MD, PhD | Sang-Hyun Ihm MD, PhD
Chang-Hwan Yoon MD, PhD | Seung Woo Park MD, PhD | Sung-Ha Park MD, PhD
Jang-Young Kim MD, PhD | Sung-Uk Kwon MD, PhD | Hae-Young Lee MD, PhD

1 Department of Internal Medicine, Seoul National University Hospital, Seoul, Republic of Korea
2 Division of Cardiology, Department of Internal Medicine, School of Medicine, Kangbuk Samsung Hospital, Sungkyunkwan University, Seoul, Republic of Korea
3 Division of Cardiology, Department of Internal Medicine, College of Medicine, The Catholic University of Korea, Seoul, Republic of Korea
4 Cardiovascular Center & Department of Internal Medicine, Seoul National University Bundang Hospital, Seongnam, Republic of Korea
5 Division of Cardiology, Department of Medicine, Samsung Medical Center, Heart Vascular Stroke Institute, Sungkyunkwan University School of Medicine, Seoul, Republic of Korea
6 Division of Cardiology, Department of Internal Medicine, Severance Cardiovascular Hospital, Yonsei University College of Medicine, Seoul, Republic of Korea
7 Department of Cardiology, Wonju College of Medicine, Yonsei University, Wonju, Republic of Korea
8 Institute of Genomic Cohort, Yonsei University, Wonju, Republic of Korea
9 Department of Medicine, Inje University Ilsan Paik Hospital, Goyang, Republic of Korea

Abstract
This multicenter, phase 4, Prospective Randomized Open, Blinded End-point (PROBE) study aimed to evaluate safety and efficacy of telmisartan/rosuvastatin single-pill combination (SPC) therapy on lowering central blood pressure (BP) compared with telmisartan monotherapy in hypertensive patients with dyslipidemia in Korea. Study was terminated earlier than planned due to COVID-19 pandemic, thus should be considered as a pilot study. Among 125 patients who met the inclusion criteria of hypertension and dyslipidemia (defined as 10-year Atherosclerotic Cardiovascular Disease risk score over 5%), 80 patients went through 4-week single-group run-in period with telmisartan 40–80 mg, then randomized to telmisartan 80 mg + rosuvastatin (10 or 20 mg) SPC group or telmisartan 80 mg monotherapy group. The central/brachial BP, brachial-ankle pulse wave velocity (baPWV), and augmentation index (AIx) were assessed at baseline and 16 weeks later. Mean brachial SBP changed from 135.80 ± 14.22 mmHg to 130.69 ± 13.23 mmHg in telmisartan/rosuvastatin group and from 134.37 ± 12.50 mmHg to 133.75 ± 12.30 mmHg in telmisartan monotherapy group.
without significant difference (between-group difference \( p = .149 \)). Mean central SBP were reduced significantly in the telmisartan/rosuvastatin group with change from 126.72 \( \pm \) 14.44 mmHg to 121.56 \( \pm \) 14.56 mmHg while telmisartan monotherapy group showed no significant change (between-group difference \( p = .028 \)). BaPWV changed from 1672.57 \( \pm \) 371.72 m/s to 1591.75 \( \pm \) 272.16 m/s in telmisartan/rosuvastatin group and from 1542.85 \( \pm \) 263.70 m/s to 1586.12 \( \pm \) 297.45 m/s in telmisartan group with no significance (between-group difference \( p = .078 \)). Change of Alx had no significant difference (between-group difference \( p = .314 \)). Both groups showed excellent compliance rate of 96.9 \( \pm \) 4.5% with no significant difference in adverse rate. Telmisartan/rosuvastatin SPC therapy was more effective in lowering central BP compared with the telmisartan monotherapy. The results of this study showed benefit of additive statin therapy in hypertensive patients combined with dyslipidemia.

**KEYWORDS**
angiotensin II receptor blocker, dyslipidemia, hypertension, rosuvastatin, telmisartan

### 1 | INTRODUCTION

Hypertension (HTN) is a known leading cause of death and disability over the world.\(^1\) Although lowering the blood pressure (BP) has been proven to reduce cardiovascular events,\(^2\) recent studies have suggested that the central BP might be more closely related to future cardiovascular events than brachial BP.\(^3\) Studies have shown stronger association of central BP with surrogate markers of cardiovascular diseases (eg, carotid intima media thickness,\(^4\) diastolic function,\(^5\) left ventricular mass\(^6\)) and target organ damage (eg, renal failure,\(^7\) cognitive deficiency\(^8\)) than brachial BP.

The central BP is a sum of incident wave from left ventricle and reflected wave from peripheral vessels at the ascending aorta. The measured contribution of reflected wave on the incident wave at the ascending aorta is defined as augmentation index (Alx).\(^9\) The Alx is affected by pulse wave velocity (PWV), which is defined by distance between two arterial sites divided by pulse transmit time.\(^10\) The Alx and PWV both have a predictive power on cardiovascular events.\(^11,12\) Thus, current studies are also focusing on the predictive role of the Alx and PWV measurement as well as central BP.

Despite the rising importance of central BP, current antihypertensive medications are mainly focused on controlling brachial BP. However, it is well known that antihypertensive classes have different impact on central BP reduction. For example, beta-blocker atenolol, was inferior in reducing central BP to brachial BP to other antihypertensive classes.\(^13\) And angiotensin-converting enzyme inhibitors (ACEIs) and angiotensin receptor blockers (ARBs) have revealed more promising effect of lowering central BP than other classes of antihypertensives.\(^14\)

Along with HTN, dyslipidemia is also known to increase the risk of cardiovascular events.\(^15\) The co-existence of HTN and dyslipidemia is relatively abundant, with its prevalence of at least over 15%.\(^16\) The co-existence leads to a higher risk of cardiovascular event than a simple sum of the two factors.\(^17\) Among the treatments of dyslipidemia, hydroxyl-methyl-glutaryl-coenzyme A reductase inhibitors, statins, are the fundamental drug.\(^18\) Statins were reported to have modest BP lowering effect and is expected to have a beneficial role in arterial stiffening.\(^19\)

Telmisartan is a highly selective blocker to angiotensin type-1 receptor with a long elimination half-life that effectively reduces BP for 24-h dosage interval.\(^20\) Rosuvastatin is highly effective in lowering low-density lipoprotein-cholesterol (LDL-C) and increasing high-density lipoprotein-cholesterol (HDL-C).\(^21\) Previously, the telmisartan/rosuvastatin single-pill combination (SPC) has shown non-inferiority compared to the telmisartan or rosuvastatin monotherapy in lowering brachial BP and LDL-C levels, respectively.\(^22\) However, the role of telmisartan/rosuvastatin combination therapy in central BP has not been evaluated yet.

Thus, this study investigated the efficacy and safety of telmisartan/rosuvastatin SPC therapy in patients with HTN and mild dyslipidemia of 10-year Atherosclerotic Cardiovascular Disease (ASCVD) risk score over 5% compared with telmisartan monotherapy.

### 2 | METHODS

#### 2.1 | Study population

The study population consisted of hypertensive patients aged 50–75 years old, with a calculated ASCVD risk of more than 5%. The participants should also be untreated with dyslipidemia (not under medication or stopped medication for at least 4 weeks prior to randomization). The exclusion criteria were the following: (1) ARB hypersensitivity, (2) secondary HTN, (3) history of myocardial infarction/unstable angina/stroke within 6 months, (4) history of uncompensated congestive heart failure or left ventricular ejection
fraction < 40% within 6 months, (5) hemodynamically significant (more than a moderate degree) valvular heart disease, and (6) current serum creatinine > 3 mg/dl, etc.

2.2 Study design and procedures

This study was a randomized, multicenter, phase 4, Prospective, Randomized, Open-label, Blinded End-point (PROBE) study, which took place from January 11, 2018 to September 2, 2020. All patients were informed thoroughly about the trial and agreed upon written consent. The study protocols were approved by the Institutional Review Board (SNUH 2017-0412) and were done according to the principles of the Declaration of Helsinki. This study was registered in Clinicaltrials.gov: NCT03267329.

The patients were randomized after 4 weeks of run-in period with telmisartan 40 or 80 mg if they met the inclusion criteria. During the run-in period, the patients who were already taking anti-hypertensive therapy had their medicine changed to telmisartan 80 mg monotherapy. Others with no experience of anti-hypertensive therapy were prescribed with telmisartan 40 or 80 mg. After the run-in period, the patients were randomized via 1:1 stratified randomization by the institute. Depending on the ASCVD risk score during the screening, those with a risk score between 5% and 7.4% received rosuvastatin 10 mg, and those with a risk score above 7.5% received rosuvastatin 20 mg. All medications were taken orally once a day, and the patients were evaluated about the efficacy and safety three times during the 16 weeks of the treatment period (Figure 1).

2.3 BP measurement

Electronic BP monitors (Omron HEM-7080IC, Omron Corporation, Kyoto, Japan) were used to measure brachial BP, and all laborato-ries used the same product. The BP of both arms was measured by a trained study coordinator at the screening visit after an initial 5 min of seated rest, and the arm with the higher average SBP obtained by three measurements was selected as the reference arm. The average value of the three measurements at screening visit was used as a baseline.

The central hemodynamic parameters were evaluated in the sitting position after 10 min of resting with overnight fasting using a SphygmoCor software version 7.0 (AtCor Medical, Sydney, Australia). The overnight fasting state was defined as fasting over 8 h. Mean arterial pressure was determined by mathematical integration of the radial pressure waveform and calibrated using the oscillographic value of brachial SBP and DBP. Pulse pressure (PP) amplification was calculated as the ratio of brachial PP: central PP. Brachial-ankle pulse wave velocity (baPWV) was measured according to the manufacturer’s protocol using VP-1000 (Omron Healthcare CO., Ltd., Kyoto, Japan) after participants were supine for 5–10 min.

2.4 Efficacy outcomes

The change in central SBP at 16 weeks from the baseline was evaluated as the primary efficacy outcome (Figure 1). The secondary outcome consisted of the followings: (1) changes of mean brachial SBP (2) changes of AIx (3) changes of baPWV (4) changes of mean brachial PP, and (5) changes of mean brachial DBP at 4 and 16 weeks. The tolerability was evaluated with the occurrence of the adverse reaction, clinical laboratory results, and vital signs.

2.5 Sample size and statistical analysis

To estimate the appropriate sample size, the non-inferiority margin of change of central SBP at 16 weeks was assumed as 4 mmHg with the
TABLE 1  Baseline characteristics

| Mean ± SD                      | Telmisartan (N = 33) | Telmisartan/rosuvastatin (N = 36) | Total (N = 69) | p-value |
|--------------------------------|-----------------------|-----------------------------------|----------------|---------|
| Age (years)                    | 62.15(10.35)          | 60.92(8.54)                       | 61.51(9.40)    | .417a   |
| Sex (male %)                   | 22(66.67)             | 23(63.89)                         | 45(65.22)      | .809b   |
| Height (cm)                    | 162.00(8.47)          | 162.53(8.76)                      | 162.28(8.56)   | .800c   |
| Weight (kg)                    | 69.67(12.72)          | 67.97(12.35)                      | 68.78(12.47)   | .575c   |
| Body mass index (BMI, kg/m²)   | 26.47(4.07)           | 25.60(3.30)                       | 26.01(3.69)    | .333c   |
| Brachial SBP (mmHg)            | 134.37(12.50)         | 135.80(14.22)                     | 135.28(13.76)  | .648a   |
| Brachial DBP (mmHg)            | 82.88(8.75)           | 82.37(9.50)                       | 82.68(9.52)    | .862a   |
| Pulse rate (BPM)               | 75.49(10.74)          | 76.39(11.40)                      | 74.47(11.13)   | .736c   |
| Smoking (%)                    | 10(30.30)             | 8(22.22)                          | 18(26.09)      | .445b   |
| Previous antihypertensive medication (%) | 28(84.85)         | 27(75.00)                         | 55(79.71)      | .310b   |
| RAS inhibitors                 | 25(75.76)             | 24(66.67)                         | 49(71.01)      | .406b   |
| Calcium channel blocker        | 4(12.12)              | 4(11.11)                          | 8(11.59)       | .896b   |
| Diuretics                      | 0(0.00)               | 1(2.78)                           | 1(1.45)        | .335b   |
| Other medication (%)           | 18(54.54)             | 17(47.22)                         | 35(50.72)      | .543b   |
| Statins                        | 10(30.30)             | 8(22.22)                          | 18(26.09)      | .445b   |
| Antithrombotic agents          | 11(33.33)             | 8(22.22)                          | 19(27.54)      | .302b   |
| Antidiabetic agents            | 6(18.18)              | 6(16.67)                          | 12(17.39)      | .868b   |

Abbreviations: BPM, beats per minute; DBP, diastolic blood pressure; RAS inhibitors, renin-angiotensin system inhibitors including angiotensin receptor blockers and angiotensin converting enzyme inhibitors; SBP, systolic blood pressure; SD, standard deviation.

aWilcoxon rank sum test.
bChi-square test.
cTwo sample t-test.

standard deviation (SD) as 6.8 mmHg. With a significance level of 5% and a power of 80%, the estimated sample size was 46 participants. Assuming the dropout rate of 10%, minimum of 52 participants for each group (a total of 104) was needed for the enrollment.

The intention to treat (ITT) population was defined as participants who had at least one measurement of primary efficacy endpoint among those enrolled after evaluating inclusion/exclusion criteria. Per protocol (PP) set was defined as participants who completed the study without major protocol deviation among the ITT population. The safety set was defined as any of the participants who were treated at least once after randomization. The analysis of baseline characteristics was done at the ITT population. All statistical analyses were done via SAS version 9.4 (SAS Institute, Inc, NC, USA) with a two-tailed test with a significance level of 5%.

The primary efficacy endpoint of change from baseline to central SBP at 16 weeks was analyzed at the ITT population with paired t-test (or Wilcoxon signed-rank test) in each treatment group. For between-group comparison, analysis of covariance was used with baseline central SBP as a covariate. In addition, the between-group difference of all the adverse events and the adverse drug reaction was analyzed using the Chi-square test (or Fisher’s exact test).

3  | RESULTS

3.1  | Patient recruitment and flow

Total 125 patients were screened throughout eight hospitals, and 80 of the patients remained for randomization. In response to the COVID-19 pandemic from February 2020, there was considerable difficulty in patients’ enrollment. Due to the limitation of study budget, we decided to finish patient enrollment by April 2020. As a result, the number of participants randomized for each group was about 77% of the original plan (A minimum of 52 participants for each group).

Among the screened 125 patients, 45 failed to pass the screening (36 did not qualify inclusion criteria, 8 withdrew consent, 2 other reasons unclarified) and 80 patients went through single-drug run-in. Of total 80 single-drug run-in population, 71 patients (36 from the telmisartan/rosuvastatin SPC group, 35 from the telmisartan monotherapy group) completed the clinical trial, and 9 (five from the telmisartan/rosuvastatin SPC group, four from the telmisartan monotherapy group) were withdrawn for the following reasons: consent withdrawal (six patients), adverse events (two patients), and protocol violation (one patient) (Figure 2).
3.2 | Baseline characteristics

Two treatment groups were comparable for all the baseline demographic and clinical characteristics (Table 1). The mean (SD) age was 61.51(9.40), the male proportion of 65%. Mean baseline office SBP/DBP were 135.28(13.76)/82.68(9.52) mmHg. Among the patients, 79.71% were previously treated with anti-hypertensive medications, most frequently with renin-angiotensin system inhibitors (71.01%), followed by calcium channel blockers (11.59%). None of the patients in this study population were treated with beta-blockers. There was no significant difference in previous anti-hypertensive medication history between two groups. Statins (26.09%), anti-thrombotic agents (27.54%), oral hypoglycemic agents (17.39%) were the most prevalent drug among other non-antihypertensive drugs. The dyslipidemia medication, including statin, was discontinued since visit 1 to avoid concomitant usage with clinical trial medications.

3.3 | Changes in brachial and central BP

The primary efficacy endpoint, change of mean central SBP at 16 weeks from baseline, significantly decreased in the telmisartan/rosuvastatin SPC group by $-5.17 \pm 14.9$ mmHg in ($p = .045$), whereas there was no significant change in the telmisartan monotherapy group ($+1.94 \pm 8.9$ mmHg, $p = .219$) (Figure 3A). The least square (LS) mean (standard error) in telmisartan/rosuvastatin SPC group was $-4.83(1.96)$ mmHg in comparison to telmisartan monotherapy group $1.57(2.05)$ mmHg, resulting in a significant decrease only in the telmisartan/rosuvastatin group. The LS mean difference between the two groups was $-6.41$ mmHg, indicating a significant reduction in the telmisartan/rosuvastatin group ($p = .028$) (Figure 3B).

The brachial BP, one of the secondary efficacy endpoints, was also evaluated at week 4 and 16. The brachial SBP in telmisartan/rosuvastatin SPC group were significantly reduced in within-group analysis at week 16 ($p = .044$) but did not show significant difference in between-group analysis with telmisartan monotherapy group (Table 2). The central DBP and brachial DBP both failed to show significant reduction in either group.

3.4 | Changes in baPWV and aortic functional parameters

AIx did not show significant change at week 16 compared with the baseline in both in-group ($p = .201$ in telmisartan monotherapy group, $p = .854$ in telmisartan/rosuvastatin SPC group) and
between-group analysis (p = .314). The baPWV showed decrease in telmisartan/rosuvastatin SPC group although insignificant (from mean 1672.57 m/s to 1591.75 m/s p = .134). There was no statistical significance between the two groups in baPWV (between-group p = .078) (Table 3).

3.5 | Changes in biochemical parameters

Serum creatinine, serum potassium, hemoglobin A1C, liver enzyme (aspartate aminotransferase, alanine aminotransferase), high-density lipoprotein did not show significant difference after 16 weeks compared to the baseline between the two groups. One patient in telmisartan/rosuvastatin SPC group showed abnormal increase in liver enzyme and was reported as ‘hepatic enzyme increase’ adverse effect. The total cholesterol, triglyceride, low-density lipoprotein of both groups showed significant difference between the two groups at week 16 (p < .001,.047, <.001) (Table 4). However, none of the patients showed significant change at the shift table, indicating insignificance of the reduction in the above three values.

3.6 | Compliance and tolerability

The overall compliance rate of two groups were 96.14 (SD 4.86)% for the telmisartan group and 97.53 (SD 4.03)% for the telmisartan/rosuvastatin SPC group with no significant difference (p = .274). Of the 77 patients treated with the medication at least once after enrollment, 10 patients experienced 16 adverse events in total. The telmisartan group had five participants with nine events, and the telmisartan/rosuvastatin group had four participants with seven events. The incidence rate between the two groups did not differ significantly (p = 1.000). A major adverse event was found once at one participant in the telmisartan group: a renal cyst (not an adverse drug reaction). The adverse event that resulted in discontinuation of the treatment was found once each for both telmisartan and telmisartan/rosuvastatin group (p = 1.000). The most common adverse event was asthenia, which was found in a total of two participants, one from each group. The adverse event that resulted in the treatment’s discontinuation was a headache in the telmisartan group, and hepatic enzyme increase in the telmisartan/rosuvastatin group. The headache was considered as an adverse drug reaction related to the clinical trial medication.

4 | DISCUSSION

In this study, we found that the telmisartan/rosuvastatin SPC group had more significantly reduced mean central SBP than the telmisartan monotherapy group in patients with HTN and mild dyslipidemia. The brachial SBP were also reduced in telmisartan/rosuvastatin SPC group but without significance compared to telmisartan monotherapy group. The between-group difference of Alx and baPWV were statistically insignificant despite the baPWV being reduced only in telmisartan/rosuvastatin SPC group.

Although statin’s effect on lowering BP has been identified previously, its effect on lowering central BP is still controversial. In a small-scale randomized controlled trial, the statin usage reduced central BP and aortic stiffness. However, in the CAFE-LLA study, statin did not lower central BP. Most recently, in the study from the CARTaGENE cohort, the statin use was significantly associated only in patients who were targets of primary prevention. This result is in line with our study as all the participants in this study were at least over 5% of the ASVCD risk factor, which makes them the target of primary prevention.
TABLE 2  Adjusted changes in central BPs and brachial BPs in the telmisartan/rosuvastatin SPC and telmisartan monotherapy groups

| Variables            | Visit  | Telmisartan (N = 33) | Telmisartan/rosuvastatin (N = 36) | LS mean difference [95% CI] | p-value |
|----------------------|--------|----------------------|-----------------------------------|-----------------------------|---------|
| Mean central SBP     |        |                      |                                   |                             |         |
| (mmHg)               |        | Mean (SD)            | Mean (SD)                         |                             |         |
| Baseline             |        | 124.33(14.20)        | 126.72(14.44)                     |                             | .491b   |
| Week 16              |        | 126.27(16.36)        | 121.56(15.64)                     |                             |         |
| p-value              |        | .219d                | .045d                             |                             |         |
| Changea              |        | 1.57(2.05)           | -4.83(1.96)                       | -6.41[−12.08, −0.73]        | .028    |
| Mean central DBP     |        |                      |                                   |                             |         |
| (mmHg)               |        | Mean (SD)            | Mean (SD)                         |                             |         |
| Baseline             |        | 81.36(10.66)         | 81.11(12.70)                      |                             | .929b   |
| Week 16              |        | 82.48(13.67)         | 79.92(9.92)                       |                             |         |
| p-value              |        | .455e                | .245e                             |                             |         |
| Changea              |        | 1.73(1.89)           | -1.26(1.78)                       | -2.99[−8.25, 2.26]          | .260    |
| Mean brachial SBP    |        |                      |                                   |                             |         |
| (mmHg)               |        | Mean (SD)            | Mean (SD)                         |                             |         |
| Baseline             |        | 134.37(12.50)        | 135.80(14.22)                     |                             | .648c   |
| Week 16              |        | 133.75(12.30)        | 130.69(13.23)                     |                             |         |
| p-value              |        | .711d                | .044d                             |                             |         |
| Changea              |        | -1.99(1.13)          | -2.52(1.27)                       | -2.73[−6.39, 0.94]          | .149    |
| Mean brachial DBP    |        |                      |                                   |                             |         |
| (mmHg)               |        | Mean (SD)            | Mean (SD)                         |                             |         |
| Baseline             |        | 82.85(8.75)          | 82.37(9.50)                       |                             | .862c   |
| Week 16              |        | 83.04(11.82)         | 79.87(10.76)                      |                             |         |
| p-value              |        | .886d                | .064d                             |                             |         |
| Changea              |        | 0.21(1.33)           | -2.52(1.27)                       | -2.73[−6.39, 0.94]          | .143    |

Abbreviations: DBP, diastolic blood pressure; LS, least square; PP, pulse pressure; SBP, systolic blood pressure; SD, standard deviation; SE, standard error.

The different result between the two groups might partly be explained by the withdrawal of statin in 30% of participants who were previously prescribed with statin in the telmisartan monotherapy group. Previous studies have shown the withdrawal of statin to be associated with increased risk of stroke or acute coronary syndrome. However, those studies were focused on the acute phase of vascular stress with inclusion criteria of patients who had stroke or coronary artery disease in the previous 24 h, respectively. On the other hand, study by McGowan showed no clinically important increase of acute coronary syndrome in the stable cardiac patients who went through short-term statin withdrawal. As our study population excluded any participants who had acute coronary syndrome or stroke within 6 months, the withdrawal of statin would not have affected as much as the previous studies by Blanco and colleagues or Heeschen and colleagues but rather be similar to study by McGowan. Also, approximately 75% of study population were never treated with statin before, thus the difference cannot solely be explained by the effect of withdrawal of statin.

TABLE 3  Adjusted changes in arterial stiffness assays in the telmisartan/rosuvastatin SPC and telmisartan monotherapy groups

| Variables            | Visit  | Telmisartan (N = 33) | Telmisartan/rosuvastatin (N = 36) | LS mean difference [95% CI] | p-value |
|----------------------|--------|----------------------|-----------------------------------|-----------------------------|---------|
| Brachial-ankle pulse | Baseline| Mean (SD)            | Mean (SD)                         |                             | .130d   |
| wave velocity (cm/s) | Week 16| 1542.85(263.70)      | 1672.57(371.72)                   |                             |         |
| Changea              |        | 19.70(34.15)         | -66.40(33.14)                     | -86.10[−182.09, 9.90]       | .078    |
| Augmentation index   | Baseline| Mean (SD)            | Mean (SD)                         |                             | .943h   |
| (%)                  | Week 16 | 35.91(26.15)         | 33.67(18.32)                      |                             |         |
| Changea              |        | -1.99(1.57)          | 0.22(1.50)                        | 2.21[−2.14, 6.56]           | .314    |

Abbreviations: DBP, diastolic blood pressure; LS, least square; PP, pulse pressure; SBP, systolic blood pressure; SE, standard error.

The different result between the two groups might partly be explained by the withdrawal of statin in 30% of participants who were previously prescribed with statin in the telmisartan monotherapy group. Previous studies have shown the withdrawal of statin to be associated with increased risk of stroke or acute coronary syndrome. However, those studies were focused on the acute phase of vascular stress with inclusion criteria of patients who had stroke or coronary artery disease in the previous 24 h, respectively. On the other hand, study by McGowan showed no clinically important increase of acute coronary syndrome in the stable cardiac patients who went through short-term statin withdrawal. As our study population excluded any participants who had acute coronary syndrome or stroke within 6 months, the withdrawal of statin would not have affected as much as the previous studies by Blanco and colleagues or Heeschen and colleagues but rather be similar to study by McGowan. Also, approximately 75% of study population were never treated with statin before, thus the difference cannot solely be explained by the effect of withdrawal of statin.
TABLE 4 Changes of lipid profile at week 16 compared with baseline in the telmisartan/rosvastatin SPC and telmisartan monotherapy groups

| Variables                 | Visit       | Telmisartan (N = 38) | Telmisartan/rosvastatin (N = 39) | Total (N = 77) |
|---------------------------|-------------|----------------------|----------------------------------|---------------|
| Total cholesterol (mg/dl) | Baseline    | Mean (SD) 203.44(35.71) | 205.81(40.40) | 204.64(37.93) |
|                           | Week 16     | Mean (SD) 210.68(40.96) | 147.23(31.99) | 178.52(48.45) |
|                           | Changea     | Mean (SD) 7.09(32.30)  | -59.96(36.34) | -26.90(48.03) |
|                           | p-value     | .196b                | <.001b              | <.001c         |
| Triglyceride (mg/dl)      | Baseline    | Mean (SD) 219.87(158.51) | 179.18(77.77) | 199.26(125.20) |
|                           | Week 16     | Mean (SD) 193.60(102.82) | 155.28(104.41) | 174.17(105.68) |
|                           | Changea     | Mean (SD) 6.57(96.78)  | -27.33(73.05) | -10.62(86.66) |
|                           | p-value     | .949c                | <.001c              | .047e           |
| HDL-cholesterol (mg/dl)   | Baseline    | Mean (SD) 44.29(9.78)  | 49.38(11.94) | 46.87(11.15) |
|                           | Week 16     | Mean (SD) 46.22(11.23) | 53.83(12.81) | 50.03(12.56) |
|                           | Changea     | Mean (SD) 1.36(6.13)  | 4.25(6.55) | 2.81(6.46) |
|                           | p-value     | .192b                | <.001b              | .057d           |
| LDL-cholesterol (mg/dl)   | Baseline    | Mean (SD) 117.09(44.74) | 123.93(35.57) | 120.56(40.24) |
|                           | Week 16     | Mean (SD) 129.10(32.74) | 65.32(28.05) | 96.30(44.07) |
|                           | Changea     | Mean (SD) 3.82(33.13) | -59.84(34.75) | -28.92(46.53) |
|                           | p-value     | .506b                | <.001b              | <.001d          |

Abbreviations: HDL, high density lipoprotein; LDL, low density lipoprotein; LS, least square; SD, standard deviation.

aChange = (Value at week 16) – (Value at baseline).

Although insignificant in between-group analysis, the baPWV of the telmisartan/rosvastatin SPC group showed greater reduction at week 16 from the baseline compared to the telmisartan monotherapy group. Such reducing tendency of baPWV might have contributed to the decrease in central BP. There are not many studies that reveal the association between reduced PWV and reduced central BP. Study by Hirata et al. (2005) and Raff et al. (2015) are one of the few studies showing decrease in PWV and central BP after use of ramipril and olmesartan, respectively.31,32

PWV is the most known measure of arterial stiffness, along with Alx.33 Arterial stiffness is caused by hemodynamic forces, extrinsic factors including hormones, and ARBs can significantly reduce Alx.35 Furthermore, meta-analysis by Yen and colleagues suggested that ARBs can also reduce PWV.26 Not only ACEIs and ARBs but also meta-analyses on statin can significantly reduce the aortic stiffness, defined by PWV and Alx.37-39 Such effect of the drugs is explained by renin-angiotensin-axis system mechanism of ARBs and antioxidant and anti-inflammatory activity of statin.36,40

As the telmisartan monotherapy group showed no reduction in PWV at all, the reducing tendency shown at telmisartan/rosvastatin SPC group might have been due to the addition of statin. The effect of statin on PWV is expected to be dose dependent.41 Thus, the statistical significance of statin on PWV in this trial might have been attenuated due to the mixture of rosvastatin dosage 10 and 20 mg. In this study, the reduction of PWV showed marginal significance with Alx having no statistically significant difference between the two groups. Such result could be due to different effect of treatment duration on PWV and Alx. The meta-analysis by Sahbekar and colleagues found positive association between statin treatment duration and Alx.39 The other hand, the meta-analysis by D’elia and colleagues and Upala and colleagues found reduction of PWV despite the difference of statin treatment duration.37,38 Thus, the discordance on the change of PWV and Alx may be due to baPWV being independent from treatment duration but Alx being more dependent on the duration. Also, although study by Van Doornum and colleagues showed reduction of Alx in rheumatoid arthritis patients after 12 weeks of atorvastatin, the potency of statin is thought to differ between the type of statin.39,42 Thus, appropriate duration needed for the reduction of Alx might differ in rosuvastatin. Furthermore, Alx is known to be affected by age, showing different response according to age, reaching plateau at age of 60 years.43 This difference makes Alx more sensitive marker in younger age group and less feasible to elders. Study by Hayashi and colleagues even showed that Alx might not be an appropriate marker to identify coronary artery disease in elderly.44 On the other hand, PWV increases along with age even after 50 years.45 And the aging influences on PWV more prominently in female.46 As the mean age of our study population was 61.5 years, the Alx might not have been a suitable marker leaving baPWV as a more feasible one. This is one of the limitations of our study, and further study comprising younger age group would be necessary to clear this point.

The cause why BP lowering at week 16 compared to baseline in telmisartan monotherapy group was insignificant, might come from the
4-week single group run-in period with telmisartan 40–80 mg. As our study protocol comprised of 4-week run-in period with telmisartan 40 or 80 mg, most participants in telmisartan monotherapy group continued their run-in period treatment without change. Therefore, it is not surprising that there was no significant change in BP between the baselines to week 16. Actually, among the 33 participants who were in telmisartan monotherapy group, only five participants were prescribed with telmisartan 40 mg in the run-in period and up-titrated to telmisartan 80 mg during the 16 weeks of treatment. Other 28 participants were on telmisartan 80 mg since run-in period. Among the five participants who had telmisartan dosage changed, three participants showed further BP reduction whereas two did not show any reduction of BP (data not shown). This proportion of patients who had telmisartan dosage up-titrated in the treatment period might be too small to affect significance of within-group reduction.

Pulse pressure amplification (PPA), defined by ratio of brachial PP to central PP, is another factor known to be inversely associated with cardiovascular risk and mortality. Decrease in central SBP caused to central PP, is another factor known to be inversely associated with cardiovascular risk and mortality.7,47 Decrease in central SBP caused by statin as shown in this study would eventually give rise to PPA ultimately resulting in reduced cardiovascular risk and mortality.

The main limitation of this study was a premature closure of enrolment, restricting the study as a pilot study. Although the investigators tried every effort to complete enrollment, the COVID-19 crisis, which has continued in the current situation, significantly disabled patient’s visit in the tertiary hospitals, which were recruited to COVID-19 patient’s care. Therefore, after deep consideration, the investigators and the sponsor decided to finish the patient enrollment in April 2020. This premature termination of patient enrolment unfulfilled the statistically calculated population, which is the major limitation of this study.

Because of the small study population, the result was interpreted with careful consideration of the possible weak points derived by the small study design. As a small study, the probability of false positive (type 1 error) is increased and over-estimation of the magnitude of association can be found more easily.48 Not only type 1 error, but also previous evaluation on randomized controlled trials also showed a need for caution to the occurrence of type 2 error in underpowered randomized controlled trials.49 Consequently, the result of our study cannot be confirmative and necessitates the need for larger confirmatory study.

The marginal significance of PWV mostly came from the smaller sample size than originally planned. Besides underpowered study, the marginal significance of PWV may be due to the other factors that influence PWV. The PWV is known to be influenced by PP, age, sex, BP, body mass index, triglyceride, blood glucose, salt intake, electrocardiogram voltage, urine albumin, and genetic factors.50–52 In this study, the two groups did not differ between the PP, age, and sex. However, other factors including salt intake, urine albumin, genetic factors were not considered. Thus, such factors might have attenuated the PWV difference in this study. This is one of the limitations of our study, and further study comprising younger age group would be necessary to clear this point.

On the contrary, there is also a chance that the effect has been exaggerated due to higher baseline baPWV in telmisartan/rosuvastatin SPC group. Although the participants were randomly assigned to each group, there was numerical difference in baseline PWV yet without statistical significance (p = .130). We did not exclude the possibility that the difference might be significant if the original enrolment was completed (type 2 error). However, telmisartan the monotherapy group showed numerically increased PWV by 2.9%, whereas the telmisartan/rosuvastatin SPC group showed decreased PWV by 4.8%. Therefore, even though the extent of change might be influenced by the baseline values, we think that the difference between two groups remains significant.

5 | CONCLUSIONS

In this study, telmisartan/rosuvastatin combination therapy showed a reduction in both central and brachial SBP when the telmisartan monotherapy group failed to show significant reduction in patients with HTN and mild dyslipidemia. The results of this study showed another benefit of statin therapy in hypertensive patients combined with dyslipidemia.
3. McEniery CM, Cockcroft JR, Roman MJ, Franklin SS, Wilkinson IB. Central blood pressure: current evidence and clinical importance. Eur Heart J. 2014;35(26):1719-1725.

4. Stamatakopoulos KS, Manios E, Barlas G, et al. Time rate of blood pressure variation is superior to central hemodynamics as an associate of carotid intima-media thickness. J Hypertens. 2010;28(1):51-58.

5. Kim HL, Lim WH, Seo JB, Kim SH, Zo ZH, Kim MA. The comparison of the impact of arterial stiffness and central pressure on left ventricular geometry and diastolic function. Clin Hypertens. 2019;25:18.

6. Weber T, Wasserteurer S, Schmidt-Trucksäss A, et al. Relationship between 24-hour ambulatory central systolic blood pressure and left ventricular mass: a prospective multicenter study. Hypertension. 2017;70(6):1157-1164.

7. Safar ME, Blacher J, Pannier B, et al. Central pulse pressure and mortality in end-stage renal disease. Hypertension. 2002;39(3):735-738.

8. Dias Eda M, Giollo LT Jr, Martinelli DD, et al. Carotid intima-media thickness is associated with cognitive deficiency in hypertensive patients with elevated central systolic blood pressure. Cardiovasc Ultrasound. 2012;10:41.

9. Trudeau L. Central blood pressure as an index of antihypertensive control: determinants and potential value. Can J Cardiol. 2014;30(5 Suppl):S23-S28.

10. O'Rourke MF, Staessen JA, Vlachopoulos C, Drezep D, Plante GE. Clinical applications of arterial stiffness: definitions and reference values. Am J Hypertens. 2002;15(5):442-444.

11. Willum-Hansen T, Staessen JA, Torp-Pedersen C, et al. Prognostic value of aortic pulse wave velocity as index of arterial stiffness in the general population. Circulation. 2006;113(5):664-670.

12. Vlachopoulos C, Aznaouridis K, O'Rourke MF, Safar ME, Boua K, Stefanadis C. Prediction of cardiovascular events and all-cause mortality with central haemodynamics: a systematic review and meta-analysis. Eur Heart J. 2010;31(15):1865-1871.

13. Dhakam Z, McEniery CM, Yasmin, Cockcroft JR, Brown MJ, Wilkinson IB. Atenolol and eprosartan: differential effects on central blood pressure and aortic pulse wave velocity. Am J Hypertens. 2006;19(2):214-219.

14. Boutouyrie P, Aoucha E, Trunet P, Laurent S. Amlodipine-valsartan combination decreases central systolic blood pressure more effectively than the amlodipine-atenolol combination: the EXPLOR study. Hypertension. 2015;65(6):1314-1322.

15. Yusuf S, Hawken S, Ounpuu S, et al. Effect of potentially modifiable risk factors associated with myocardial infarction in 52 countries (the INTERHEART study): case-control study. Lancet. 2004;364(9438):937-952.

16. Onat E, Hergenc G, Sanli I, Turkmens E, Can G, Sango V, V. Dyslipidemic hypertension: distinctive features and cardiovascular risk in a prospective population-based study. Am J Hypertens. 2005;18(3):409-416.

17. Stamler J, Wentworth D, Neaton JD. Prevalence and prognostic significance of hypercholesterolemia in men with hypertension. Prospective data on the primary screeners of the Multiple Risk Factor Intervention Trial. Am J Med. 1986;80(2a):33-39.

18. Reiter-Brennan C, Osei AD, Iftekhar Uddin SM, et al. ACC/AHA lipid guidelines: personalized care to prevent cardiovascular disease. Cleve Clin J Med. 2020;87(4):231-239.

19. Golomb BA, Dimsdale JE, White HL, Ritchie JB, Criqui MH. Reduction in blood pressure with statins: results from the UCSDD Statin Study, a randomized trial. Arch Intern Med. 2008;168(7):721-727.

20. Smith DH, Cramer MJ, Neutel JM, Hettiarachchi R, Koval S. Comparison of telmisartan versus losartan: meta-analysis of titration-to-response studies. Blood Press Monit. 2003;8(3):111-117.

21. Nissen SE, Nicholls SJ, Sipahi I, et al. Effect of very high-intensity statin therapy on regression of coronary atherosclerosis: the ASTEROID trial. JAMA. 2006;295(13):1556-1565.

22. Oh GC, Han J-K, Han KH, et al. Efficacy and safety of fixed-dose combination therapy with telmisartan and rosuvastatin in korean patients with hypertension and dyslipidemia: TELSTA-YU (TELmisartan-rosuvaSTAtin from YUhan), a multicenter, randomized, 4-arm, double-blind, placebo-controlled, phase III study. Clin Ther. 2018;40(5):675-691.e1.

23. Matsu Y, Eguchi K, O'Rourke MF, et al. Differential effects between a calcium channel blocker and a diuretic when used in combination with angiotensin II receptor blocker on central arterial pressure in hypertensive patients. Hypertension. 2009;54(4):716-723.

24. Munakata M, Nunokawa T, Taya J, Yoshinaga K, Toyota T. Brachial-ankle pulse wave velocity as a novel measure of arterial stiffness: present evidences and perspectives. Curr Hypertens Rev. 2005;1(3):223-234.

25. Ferrier KE, Muhlmann MH, Baguet JP, et al. Intensive cholesterol reduction lowers blood pressure and large artery stiffness in isolated systolic hypertension. J Am Coll Cardiol. 2002;39(6):1020-1025.

26. Williams B, Lacy PS, Cruickshank JK, et al. Impact of statin therapy on central aortic pressures and hemodynamics: principal results of the Conduit Artery Function Evaluation-Lipid-Lowering Arm (CAFE-LLA) Study. Circulation. 2009;119(1):53-61.

27. Lamarche F, Agharazi M, Nadeau-Fredette AC, Madore F, Goupil R. Central and brachial blood pressures, statins, and low-density lipoprotein cholesterol: a mediation analysis. Hypertension. 2018;71(3):415-421.

28. Blasco M, Nombela F, Castellanos M, et al. Statin treatment withdrawal in ischemic stroke: a controlled randomized study. Neurology. 2007;69(9):904-910.

29. Heeschen C, Hamm CW, Laufs U, Snapinn S, Böhm M, White HD. Withdrawal of statins increases event rates in patients with acute coronary syndromes. Circulation. 2002;105(12):1446-1452.

30. McGowan MP. There is no evidence for an increase in acute coronary syndromes after short-term abrupt discontinuation of statins in stable cardiac patients. Circulation. 2004;110(16):2333-2335.

31. Hirata K, Vlachopoulos C, Adjii A, O'Rourke MF. Benefits from angiotensin-converting enzyme inhibitor ‘beyond blood pressure lowering’: beyond blood pressure or beyond the brachial artery? J Hypertens. 2005;23(3):551-556.

32. Raff U, Walker S, Ott C, Schneider MP, Schmieder RE. Olmesartan improves pulse wave velocity and lowers central systolic blood pressure and ambulatory blood pressure in patients with metabolic syndrome. J Clin Hypertens (Greenwich). 2015;17(2):98-104.

33. Jannas I, Sondej T, Targowski T, Dobrowolski A, Olzewska R. [Pulse wave velocity - a useful tool in assessing the stiffness of the arteries]. Pol Merkur Lekarski. 2014;47(276):257-262. Pomiar prędkości fal tętna – użyteczne narzędzie w ocenie sztywności tętnic.

34. Ziemia SJ, Melenovsky V, Kass DA. Mechanisms, pathophysiology, and therapy of arterial stiffness. Arterioscler Thromb Vasc Biol. 2005;25(5):932-943.

35. McGaughy TJ, Fletcher EA, Shah SA. Impact of antihypertensive agents on central systolic blood pressure and augmentation index: a meta-analysis. Am J Hypertens. 2016;29(4):448-457.

36. Yen CH, Lai YH, Hung CL, et al. Angiotensin receptor blockers effect on peripheral muscular and central arterial stiffness: a meta-analysis of randomized controlled trials and systematic review. Acta Cardiol Sin. 2014;30(2):98-107.

37. Upala S, Wirunswanyawka K, Jaruvongvanich V, Sanguankeo A. Effects of statin therapy on arterial stiffness: a systematic review and meta-analysis of randomized controlled trial. Int J Cardiol. 2017;227:338-341.

38. D’elia L, La Fata E, Iannuzzo A, Rubba PO. Effect of statin therapy on pulse wave velocity: a meta-analysis of randomized controlled trials. Clin Exp Hypertens. 2018;40(7):601-608.

39. Sahebkar A, Pećin I, Tedeschi-Reiner E, Derosa G, Maffioli P, Reiner Ž. Effects of statin therapy on augmentation index as a measure of arterial stiffness: a systematic review and meta-analysis. Int J Cardiol. 2016;212:160-168.
40. Shishehbor MH, Brennan ML, Aviles RJ, et al. Statins promote potent systemic antioxidant effects through specific inflammatory pathways. Circulation. 2003;108(4):426-431.

41. Mitsiou E, Boutari C, Kotsis V, et al. Effect of low (5 mg) vs. high (20-40 mg) rosvastatin dose on 24h arterial stiffness, central haemodynamics, and non-alcoholic fatty liver disease in patients with optimally controlled arterial hypertension. Curr Vasc Pharmacol. 2018;16(4):393-400.

42. Van Doornum S, McColl G, Wicks IP. Atorvastatin reduces arterial stiffness in patients with rheumatoid arthritis. Ann Rheum Dis. 2004;63(12):1571-1575.

43. McEniery CM, Yasmin, Hall IR, Qasem A, Wilkinson IB, Cockcroft JR. Normal vascular aging: differential effects on wave reflection and aortic pulse wave velocity: the Anglo-Cardiff Collaborative Trial (ACCT). J Am Coll Cardiol. 2005;46(9):1753-1760.

44. Hayashi S, Yamada H, Bando M, et al. Augmentation index does not reflect risk of coronary artery disease in elderly patients. Circ J. 2014;78(5):1176-1182.

45. Diaz A, Tringler M, Wray S, Ramirez AJ, Cabrera Fischer EI. The effects of age on pulse wave velocity in untreated hypertension. J Clin Hypertens (Greenwich). 2018;20(2):258-265.

46. Tomiyama H, Yamashina A, Arai T, et al. Influences of age and gender on results of noninvasive brachial–ankle pulse wave velocity measurement—a survey of 12517 subjects. Atherosclerosis. 2003;166(2):303-309.

47. Benetos A, Thomas F, Joly L, et al. Pulse pressure amplification a mechanical biomarker of cardiovascular risk. J Am Coll Cardiol. 2010;55(10):1032-1037.

48. Hackshaw A. Small studies: strengths and limitations. Eur Respir J. 2008;32(5):1141-1143.

49. Abdullah L, Davis DE, Fabricant PD, Baldwin K, Namdari S. Is there truly “no significant difference”? Underpowered randomized controlled trials in the orthopaedic literature. J Bone Joint Surg Am. 2015;97(24):2068-2073.

50. Aatola H, Hutri-Kähönen N, Juonala M, et al. Lifetime risk factors and arterial pulse wave velocity in adulthood: the cardiovascular risk in young Finns study. Hypertension. 2010;55(3):806-811.

51. Sougawa Y, Miyai N, Utsumi M, Miyashita K, Takeda S, Arita M. Brachial-ankle pulse wave velocity in healthy Japanese adolescents: reference values for the assessment of arterial stiffness and cardiovascular risk profiles. Hypertens Res. 2020;43(4):331-341.

52. Sonoda H, Takase H, Dohi Y, Kimura G. Factors associated with brachial-ankle pulse wave velocity in the general population. J Hum Hypertens. 2012;26(12):701-775.

How to cite this article: Choi JM, Sung K-C, Ihm S-H, et al. Central blood pressure lowering effect of telmisartan-rosuvastatin single-pill combination in hypertensive patients combined with dyslipidemia: A pilot study. J Clin Hypertens. 2021;23:1664-1674. https://doi.org/10.1111/jch.14345