Effects of Azelnidipine plus Olmesartan versus amlodipine plus olmesartan on central blood pressure and left ventricular mass index: the AORTA study

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Purpose: The aim of this study was to compare the effects of olmesartan combined with either azelnidipine or amlodipine on central blood pressure (CBP) and left ventricular mass index (LVM) in hypertensive patients.

Patient and methods: Patients with brachial systolic BP ≥ 140 mmHg and/or diastolic BP ≥ 90 mmHg received olmesartan monotherapy (20 mg daily) for 12 weeks. The patients were then randomly assigned to fixed-dose add-on therapy with azelnidipine (16 mg daily) or amlodipine (5 mg daily) (25 patients/group) for a further 24 weeks. CBP and LVM were measured at baseline and at the end of the study.

Results: Baseline characteristics were similar in both groups. The decrease in brachial BP was similar in both groups. CBP and LVM decreased significantly in both groups (both, \( P < 0.001 \)). However, the decreases in CBP and LVM were significantly greater with olmesartan/azelnidipine than with olmesartan/amlodipine (CBP, \( P < 0.001 \); LVM, \( P = 0.002 \)).

Conclusions: These findings indicate that olmesartan/azelnidipine had greater effects on CBP and LVM than did olmesartan/amlodipine, even though the reduction in brachial BP was similar in both groups. These differential effects on CBP and LVM may have important implications for cardiovascular risk reduction.

Keywords: central blood pressure, left ventricular mass index, augmentation index, brachial-ankle pulse wave velocity, olmesartan/azelnidipine

Introduction

When blood pressure (BP) control is inadequate with a single antihypertensive drug, the use of two or three drugs in combination is often necessary to achieve the target blood pressure. Combination therapy with an angiotensin II receptor blocker (ARB) and a diuretic or an ARB plus a calcium channel blocker (CCB) are recommended in the current Japanese Society for Hypertension guidelines.1 Several studies, including the ASCOT-CAFÉ2 and the Strong Heart Study,3 have emphasized the importance of targeting central blood pressure (CBP) rather than brachial BP in terms of cardiovascular disease outcomes. For example, the ASCOT-CAFÉ study compared the efficacy of amlodipine with that of atenolol, a \( \beta \)-blocker, and showed that both regimens had very different effects on central aortic pressures and hemodynamics despite similar effects on brachial BP.4 Meanwhile, in the Strong Heart Study,1 central pulse pressure was more strongly related to vascular hypertrophy, extent of atherosclerosis, and cardiovascular events than was brachial BP.
Several studies have also shown that an ARB in combination with a CCB significantly improves CBP.\(^5\) Notably, however, the influence of CCBs on BP cannot be explained as a class effect because of differences in the effects of azelnidipine and amlodipine.\(^6\) \(^8\)

Unfortunately, unlike amlodipine, few studies have examined the effects of azelnidipine on CBP. Moreover, to our knowledge, no studies of combination therapy have been conducted in Japan. In this context, we conducted the current study (Azelnidipine plus OlmesaRTAn versus amlodipine plus olmesartan; AORTA study) to determine the effects of adding either azelnidipine or amlodipine to ongoing olmesartan in patients with inadequate BP control on a standard dose of olmesartan.

**Methods**

**Patients**

Patients aged 36–75 years were recruited from among outpatients of the Department of Internal Medicine at Clinic Jingumae (Kashihiara, Japan) between March 2007 and October 2008. We initially enrolled consecutive hypertensive patients with or without current treatment who agreed to participate in this study. Hypertension was defined as clinic-measured systolic BP (SBP) ≥ 140 mmHg and/or diastolic BP (DBP) ≥ 90 mmHg on two different occasions or by a previous diagnosis of hypertension with current antihypertensive therapy. Patients were excluded if they had secondary hypertension, arrhythmia, current treatment for congestive heart failure, a history of stroke or coronary artery disease, clinically significant valvular heart disease, renal insufficiency (serum creatinine ≥ 2 mg/dL), mental disorders, severe noncardiovascular disease (eg, cancer or liver cirrhosis), or chronic inflammatory disease. Patients who were already being treated with olmesartan were also excluded. Written informed consent was obtained from all of the patients participating in the study. The study protocol was approved by the Institutional Ethical Committee.

**Study design**

The AORTA study was a 24-week, prospective, randomized, open-label parallel-group study comprising a 12-week run-in period followed by a 24-week randomized treatment period. The design of the study is summarized in Figure 1. Patients who were already being treated with antihypertensive drugs switched their current antihypertensive medications to olmesartan monotherapy. If the clinic SBP exceeded 200 mmHg and/or DBP exceeded 115 mmHg at any time during the run-in period, the patient was withdrawn from the study.

Both regimens were given at fixed doses for 24 weeks; dose titration was not permitted. The patients were instructed to take their medications after breakfast and were not permitted to use any antihypertensive drugs other than the study drugs. Other drugs that had the potential to interfere with the safety or efficacy of the study drugs were also not permitted. Brachial BP, heart rate (HR), CBP, normalized augmentation index (Alx@75), brachial-ankle pulse wave velocity (baPWV), and left ventricular mass index (LVMI) were measured at baseline and at the end of the 24-week treatment phase.

**Measurement of CBP, Alx@75, and baPWV**

The pulse pressure waveform of the radial artery was recorded using an automated tonometry system (HEM-9000AI; Omron Healthcare, Kyoto, Japan) with the patient in the sitting position after resting for ≥5 minutes. The waveform was automatically calibrated using the built-in oscillometric brachial sphygmomanometer, and the peak and trough of the radial pressure wave were adjusted to brachial SBP and DBP, respectively. An algorithm programmed into the HEM-9000AI system performed automatic online detection of the second peak (late systolic inflection) based on the second maxima of the fourth derivative of the radial pressure waveform to determine the radial AI as well as the late or second SBP (SBP2). This algorithm is described in more detail elsewhere.\(^9\) The height of the second peak corresponds to the SBP2 value obtained by the HEM-9000AI. SBP2 is very close to invasively recorded aortic CBP.\(^10\) and was used...
as an estimate of CBP. CBP determined by the HEM9000-AI was reported to be comparable with CBP estimated using a generalized aorta–radial transfer function.\textsuperscript{11,12} The AI was calculated using the formula \((SBP_2 - DBP)/(the \ first \ peak \ SBP - DBP) \times 100.\) Because AI is influenced by the HR, it was normalized for an HR of 75 bpm (Aix@75) as proposed by Wilkinson et al.\textsuperscript{13} baPWV was also used to assess arterial stiffness. baPWV was performed as described previously.\textsuperscript{14}

Briefly, baPWV was determined from the pulse waveforms recorded from both forearms and both ankles using the formula PWV (Omron Healthcare). This parameter was measured in patients who had been lying in the supine position for at least 5 minutes. baPWV measurements were repeated twice to confirm reproducibility, and the deviation between measurements was within 5%. The mean value on the right side was used as the baPWV value in each patient.

### Measurement of LVMi

M-mode echocardiography was performed under 2-dimensional guidance using a Vivid S6 ultrasound system (GE Healthcare, Milwaukee, WI) with a 3-MHz transducer. LVMi was measured by standard M-mode echocardiography and determined using the formula reported by Devereux et al.\textsuperscript{15}

### Statistical analysis

Data are shown as means ± SD. Differences between the two groups at baseline were analyzed using unpaired \(t\) tests for continuous variables and \(\chi^2\) tests for categorical variables. Paired \(t\) tests were used for within-group comparisons while unpaired \(t\) tests were used for between-group comparisons. Values of \(P < 0.05\) (2-sided) were considered statistically significant. SAS software (v 8.2; SAS Institute Inc, Cary, NC) was used for all analyses.

### Results

#### Baseline characteristics

During the run-in period, 95 patients received 20 mg olmesartan monotherapy once daily. At the end of the run-in period, 43 patients discontinued because they did not meet the BP criteria. Thus, 52 patients with a clinic SBP ≥ 140 mmHg and/or DBP ≥ 90 mmHg were eligible and were randomized to receive add-on azelnidipine (16 mg daily) or amlodipine (5 mg daily) to ongoing olmesartan. Patients were randomized using the permuted block method. Of the 26 patients randomized to the olmesartan/azelnidipine regimen, one was excluded after missing the final assessment visit. Of the 26 patients randomized to the olmesartan/amlodipine regimen, one was excluded after missing the final assessment visit. Thus, the population included 25 patients assigned to each regimen. The characteristics of the patients at the start of the randomized phase of the study (ie, baseline) are shown in Table 1. The baseline BP and other hemodynamic parameters were similar in both groups.

#### Changes in brachial BP, CBP, and HR

Brachial BP and CBP decreased significantly in both treatment groups (all, \(P < 0.001\); Table 2). However, the decrease in CBP was significantly greater in the olmesartan/azelnidipine group than in the olmesartan/amlodipine group (−14.0 ± 4.3 vs −8.3 ± 3.7 mmHg, respectively, \(P < 0.001\); Figure 2A), whereas the decrease in brachial BP was similar in both groups. The magnitude of the decrease in HR was significantly greater in the olmesartan/azelnidipine group than in the olmesartan/amlodipine group (−4.4 ± 4.3 vs 1.1 ± 3.3 bpm, respectively, \(P < 0.001\); Figure 2B).

#### Changes in Aix@75

The Aix@75 decreased significantly in both treatment groups between baseline and endpoint (both \(P < 0.001\); Table 2). The magnitude of the decrease in Aix@75 was significantly greater in the olmesartan/azelnidipine group than in the olmesartan/amlodipine group (−8.4 ± 5.4 vs −4.3 ± 3.7%, respectively, \(P < 0.001\); Figure 2C).

### Table 1 Baseline characteristics

<table>
<thead>
<tr>
<th></th>
<th>Olmesartan/ Azelnidipine (n = 25)</th>
<th>Olmesartan/ Amlodipine (n = 25)</th>
<th>(P)-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>65.8 ± 4.1</td>
<td>67.5 ± 4.6</td>
<td>0.17</td>
</tr>
<tr>
<td>Sex (male, %)</td>
<td>17 (68%)</td>
<td>18 (72%)</td>
<td>0.75</td>
</tr>
<tr>
<td>BMI (kg/m(^2))</td>
<td>25.8 ± 1.0</td>
<td>25.8 ± 1.3</td>
<td>0.97</td>
</tr>
<tr>
<td>SBP (mmHg)</td>
<td>151.8 ± 5.6</td>
<td>151.6 ± 4.4</td>
<td>0.89</td>
</tr>
<tr>
<td>DBP (mmHg)</td>
<td>85.8 ± 5.2</td>
<td>86.2 ± 4.7</td>
<td>0.75</td>
</tr>
<tr>
<td>CBP (mmHg)</td>
<td>147.3 ± 5.4</td>
<td>146.3 ± 3.5</td>
<td>0.46</td>
</tr>
<tr>
<td>HR (bpm)</td>
<td>73.5 ± 5.8</td>
<td>73.7 ± 7.0</td>
<td>0.93</td>
</tr>
<tr>
<td>Aix@75 (%)</td>
<td>91.9 ± 4.5</td>
<td>92.7 ± 3.3</td>
<td>0.48</td>
</tr>
<tr>
<td>baPWV(cm/sec)</td>
<td>1895.9 ± 201.0</td>
<td>1848.4 ± 174.0</td>
<td>0.38</td>
</tr>
<tr>
<td>LVMi (g/m(^2))</td>
<td>122.8 ± 3.0</td>
<td>122.8 ± 2.8</td>
<td>0.96</td>
</tr>
<tr>
<td>eGFR (ml/min/1.73 m(^2))</td>
<td>66.4 ± 4.1</td>
<td>66.4 ± 4.5</td>
<td>0.97</td>
</tr>
<tr>
<td>HbA1c (%)</td>
<td>5.8 ± 1.2</td>
<td>5.7 ± 1.1</td>
<td>0.92</td>
</tr>
<tr>
<td>LDL-C (mg/dL)</td>
<td>110.3 ± 16.5</td>
<td>112.0 ± 12.3</td>
<td>0.67</td>
</tr>
<tr>
<td>HDL-C (mg/dL)</td>
<td>55.9 ± 9.0</td>
<td>55.4 ± 8.3</td>
<td>0.83</td>
</tr>
<tr>
<td>TG (mg/dL)</td>
<td>152.7 ± 75.4</td>
<td>170.4 ± 61.7</td>
<td>0.37</td>
</tr>
<tr>
<td>UA (mg/dL)</td>
<td>5.4 ± 1.0</td>
<td>5.6 ± 0.8</td>
<td>0.53</td>
</tr>
</tbody>
</table>

Note: Values are means ± standard deviation or n (%).

Abbreviations: BMI, body mass index; SBP, systolic blood pressure; DBP, diastolic blood pressure; CBP, central blood pressure; HR, heart rate; Aix@75, normalized augmentation index; baPWV, brachial-ankle pulse wave velocity; LVMi, left ventricular mass index; eGFR, estimated glomerular filtration rate; LDL-C, low-density lipoprotein cholesterol; HDL, high-density lipoprotein cholesterol; TG, triglyceride; UA, uric acid.
### Table 2 Changes in hemodynamic parameters

<table>
<thead>
<tr>
<th></th>
<th>Olmesartan/Azelnidipine</th>
<th>Olmesartan/Amlodipine</th>
<th>P-value*</th>
<th>P-value (between-group difference)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline</td>
<td>Week 24</td>
<td>Change</td>
<td>Baseline</td>
</tr>
<tr>
<td>SBP (mmHg)</td>
<td>151.8 ± 5.6</td>
<td>136.7 ± 3.1</td>
<td>−15.2 ± 5.8</td>
<td>&lt;0.001</td>
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<tr>
<td>DBP (mmHg)</td>
<td>85.8 ± 5.2</td>
<td>82.6 ± 3.1</td>
<td>−3.2 ± 3.1</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>CBP (mmHg)</td>
<td>147.3 ± 5.4</td>
<td>133.2 ± 4.3</td>
<td>−14.0 ± 4.3</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>HR (bpm)</td>
<td>73.5 ± 5.8</td>
<td>69.1 ± 4.8</td>
<td>−4.4 ± 4.3</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Aix@75 (%)</td>
<td>91.9 ± 4.5</td>
<td>83.5 ± 2.8</td>
<td>−8.4 ± 5.4</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>baPWV (cm/sec)</td>
<td>1895.9 ± 201.0</td>
<td>1620 ± 159.8</td>
<td>−275.9 ± 161.2</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>LVMI (g/m²)</td>
<td>122.8 ± 3.0</td>
<td>116.2 ± 3.3</td>
<td>−6.6 ± 3.4</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

**Notes:** Values are means ± SD. P: baseline vs week 24. **Abbreviations:** SBP, systolic blood pressure; DBP, diastolic blood pressure; CBP, central blood pressure; HR, heart rate; Aix@75, normalized augmentation index; baPWV, brachial-ankle pulse wave velocity; LVMI, left ventricular mass index.

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Changes in baPWV but not LVMI decreased significantly in both treatment groups (P < 0.001; olmesartan/azelnidipine, P = 0.003; Table 2). The magnitude of the decrease in baPWV was significantly greater in Olmesartan/Azelnidipine than Olmesartan/Amlodipine (P = 0.003).
In this study, we found that, during 24 weeks of combination therapy, olmesartan/azelnidipine elicited significantly greater decreases in CBP than olmesartan/amlodipine, even though the decrease in brachial BP was similar with both treatments. In addition, AIx@75, baPWV, and HR all showed significantly greater decreases with olmesartan/azelnidipine than with olmesartan/amlodipine. Different hemodynamic effects of the two regimens may explain why the decrease in CBP was greater with olmesartan/azelnidipine than with olmesartan/amlodipine, a concept supported by the greater decrease in AIx@75 with olmesartan/azelnidipine than with olmesartan/amlodipine. AI is a measure of the contribution of wave reflection to the aortic waveform and depends on the pulse wave velocity, and magnitude and site of the reflected pressure wave, and thus provides a composite measure of large artery (systemic) stiffness.16 Increases in CBP and the AIx@75 are influenced by pressure waves that are reflected back toward the heart from branch points throughout the arterial tree. PWV is also a measure of arterial stiffness. When arteries become sclerosed, PWV is elevated, promoting an acceleration of the waves reflected back. Renin–angiotensin system (RAS) inhibitors and CCBs act directly on peripheral sites, alter the arterial smooth muscle, and thus reduce pressure wave reflection.2,17–21 In fact, RAS inhibitors and CCBs have been shown to decrease CBP by reducing pressure wave reflection.22 Compared with amlodipine, azelnidipine may elicit greater reductions in CBP by inducing greater decreases in pressure wave reflection. Thus, the main mechanism responsible for CBP lowering associated with olmesartan/azelnidipine may involve reductions in the magnitude of and potential delays in pressure wave reflection.

Adverse events
All of the patients who entered the randomized phase of the study completed the study without experiencing any serious adverse events or drug-related adverse events.

Discussion
In this study, we found that, during 24 weeks of combination therapy, olmesartan/azelnidipine elicited significantly greater decreases in CBP than olmesartan/amlodipine, even though the decrease in brachial BP was similar with both treatments. In addition, AIx@75, baPWV, and HR all showed significantly greater decreases with olmesartan/azelnidipine than with olmesartan/amlodipine. Different hemodynamic effects of the two regimens may explain why the decrease in CBP was greater with olmesartan/azelnidipine than with olmesartan/amlodipine, a concept supported by the greater decrease in AIx@75 with olmesartan/azelnidipine than with olmesartan/amlodipine. AI is a measure of the contribution of wave reflection to the aortic waveform and depends on the pulse wave velocity, and magnitude and site of the reflected pressure wave, and thus provides a composite measure of large artery (systemic) stiffness.16 Increases in CBP and the AIx@75 are influenced by pressure waves that are reflected back toward the heart from branch points throughout the arterial tree. PWV is also a measure of arterial stiffness. When arteries become sclerosed, PWV is elevated, promoting an acceleration of the waves reflected back. Renin–angiotensin system (RAS) inhibitors and CCBs act directly on peripheral sites, alter the arterial smooth muscle, and thus reduce pressure wave reflection.2,17–21 In fact, RAS inhibitors and CCBs have been shown to decrease CBP by reducing pressure wave reflection.22 Compared with amlodipine, azelnidipine may elicit greater reductions in CBP by inducing greater decreases in pressure wave reflection. Thus, the main mechanism responsible for CBP lowering associated with olmesartan/azelnidipine may involve reductions in the magnitude of and potential delays in pressure wave reflection.
Another mechanism that may explain the decrease in CBP in this study is that olmesartan/azelnidipine decreased the HR whereas olmesartan/amldipine did not. These results are consistent with previous reports in which azelnidipine decreased clinic-measured HR and 24-hour HR, whereas amldipine increased both of these parameters, despite similar BP reductions.9,25 It was also reported that, after switching from amldipine, azelnidipine exhibited hypotensive effects compared with amldipine, and significantly decreased HR and the total number of extrasystole phases.24 Noradrenaline levels and the LF/HF ratio were significantly decreased, and the washout rate was significantly reduced on 123I-MIBG myocardial scintigraphy. These findings suggest that azelnidipine inhibits the enhancement of sympathetic nervous activity.24 These differences in the effects of azelnidipine and amldipine can also be explained by the finding that azelnidipine exerts stronger inhibition of sympathetic activity in the vasodilation-induced baroreceptor reflex than does amldipine.7

Experimental studies have also shown that azelnidipine dose-dependently reduces HR.25 In another study, azelnidipine suppressed cardiac hypertrophy, fibrosis, NADPH oxidase, and superoxide levels in stroke-prone spontaneously hypertensive rats more potently than amldipine, and was associated with lower HR than amldipine.26 Azelnidipine also caused a greater reduction than amldipine in the beat rate of the sinus node/atrial preparation of these rats.26 Based on these findings, olmesartan/azelnidipine may reduce CBP via the HR-lowering effect of azelnidipine.

In general, AI and CBP increase with HR lowering. In this study, however, both parameters decreased despite HR lowering. This result is consistent with a previous report showing that olmesartan/azelnidipine significantly decreased HR, AI, CBP, and Aortic PWV (AoPWV) by more than olmesartan/hydrochlorothiazide.5 The HR-lowering effect of azelnidipine through its sympathetic inhibition may be at least partly responsible for the greater effect of the olmesartan/azelnidipine combination on AoPWV.3 We measured baPWV rather than AoPWV, but it has been reported that baPWV correlates well with AoPWV.27

In this study, we also found that the decrease in LVMl was significantly greater with olmesartan/azelnidipine than with olmesartan/amldipine. The strong correlation between change in LVMl and change in CBP (Figure 3), suggests that the greater reduction in LVMl in the olmesartan/azelnidipine group may be related to the greater reduction of CBP in that group. In hypertension the pressure reflection wave during systole augments central ascending aortic BP.17,28,29 As a result, the LV must provide greater force during ejection to overcome the augmented pressure.30,31 AI, CBP and wasted LV effort were positively associated with LV hypertrophy in untreated hypertensive patients.32 It was also reported that an increase in the ascending aortic AI caused by a reflected wave may be involved in the formation of LV hypertrophy.33 Moreover, in the REASON study, the greater change in LV mass was linked to CBP but not brachial BP.34 Thus, LV lowering associated with olmesartan/azelnidipine may have been due to reductions in CBP and AI.

It has been reported that lowering BP with the additional use of azelnidipine is associated with improvements in LV diastolic performance (ie, an increase in the e’ velocity), a reduction in LV filling pressure (a decrease in the E/e’ ratio) and a decrease in the brain natriuretic peptide level in patients with hypertension and a preserved systolic function.35 Among patients in whom amldipine was switched to azelnidipine, BP and HR decreased significantly, and these reductions were associated with an increase in the e’ velocity.35 Therefore, regression of LVMI may be related to the improvements in LV diastolic function elicited by azelnidipine.

CBP, AI, and LV hypertrophy are independent predictors of cardiovascular morbidity in patients with hypertension.3,16,37 Epidemiological studies have shown that an increased HR is associated with a higher incidence of cardiovascular diseases and a worse prognosis.38,39 Tachycardia caused by reflex activation of the sympathetic nervous system is a major adverse effect of CCB therapy. However, unlike amldipine, azelnidipine has been shown to inhibit sympathetic activity,49 suggesting that tachycardia is unlikely to occur with azelnidipine. Indeed, we found that the HR decreased with azelnidipine, but not with amldipine.

Some limitations of this study should be discussed. First, this was not a double-blind randomized trial, meaning the possibility of a significant bias cannot be excluded. However, measurements were performed by a single clinical investigator who was blinded to the treatment allocation; therefore, there was no bias in the measurement and evaluation of laboratory data. Second, the study period was relatively short and the sample size was small; longer and larger studies are necessary to evaluate the long-term effects of these combination therapies on arterial structure. Third, the dose of azelnidipine used might also have influenced the results. However, we used the standard clinical dose of azelnidipine (ie, 16 mg/day), a dose that achieved similar reductions in BP to that achieved by a standard dose of amldipine (ie, 5 mg/day). Finally, the CBP results are based solely on radial tonometry data. Radial tonometry is comfortable for patients...
and easy to use in a clinical setting, but it is subject to inter-observer variability and the results may not be an accurate reflection of actual pressures. Therefore, future studies should validate these results using direct pressure measurements.

**Conclusion**

In conclusion, the AORTA study revealed that olmesartan in combination with azelnidipine elicited greater reductions in CBP and LVMI compared with olmesartan in combination with amlodipine, even though the reduction in brachial BP was similar with both regimens. CBP is an independent predictor of cardiovascular morbidity in hypertensive patients. Therefore, the superior effects of olmesartan/azelnidipine therapy on central hemodynamics may be associated with more favorable cardiovascular outcomes than with olmesartan/amlodipine. Such differential effects may be important for cardiovascular risk reduction and warrant large-scale, long-term clinical trials to confirm this hypothesis.

**Acknowledgment**

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**Disclosure**

The authors declare no conflicts of interest in this work.

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