Effect of Rivaroxaban and Clopidogrel Combination Therapy on In-Stent Responses After Everolimus-Eluting Stent Implantation in a Porcine Coronary Model

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Aim: According to recent clinical trials, a combination of direct oral anticoagulants with antiplatelet drugs is often recommended for atrial fibrillation patients who receive drug-eluting stents (DESs). Although the optimal combination comprises direct factor Xa inhibitors and a P2Y₁₂ receptor antagonist (or aspirin), their influence on vascular responses to DESs remains unclear.

Methods: Pigs were given either aspirin and clopidogrel (dual antiplatelet therapy [DAPT] group), aspirin and rivaroxaban (AR group), or clopidogrel and rivaroxaban (CR group), followed by everolimus-eluting stent (Promus Element) implantation into the coronary artery. Stented coronary arteries were evaluated via intravascular optical coherence tomography (OCT) and histological analysis at 1 and 3 months.

Results: OCT revealed lower neointimal thickness in the DAPT group and comparable thickness among all groups at 1 and 3 months, respectively. Histological analyses revealed comparable neointimal area among all groups and the smallest neointimal area in the CR group at 1 and 3 months, respectively. In the DAPT and AR groups, the neointima continued to grow from 1 to 3 months. A shortened time course for neointima growth was observed in the CR group, with rapid growth within a month (maintained for 3 months). A higher incidence of in-stent thrombi was observed in the AR group at 1 month; no thrombi were found in either group at 3 months. More smooth muscle cells with contractile features were found in the CR group at both 1 and 3 months.

Conclusions: Our results proved the noninferiority of the combination of rivaroxaban with an antiplatelet drug, particularly the dual therapy using rivaroxaban and clopidogrel, compared to DAPT after DES implantation.

Key words: Atrial fibrillation, Rivaroxaban, Clopidogrel, Drug-eluting stent, Porcine model, Optical coherence tomography, Histology

Introduction

The widespread use of drug-eluting stents (DESs) represents a significant advancement in interventional cardiology, offering a solution to the long-standing problem of restenosis after coronary revascularization. Although improved DESs provide a reduction in in-stent restenosis and a reduced risk of stent thrombosis, prolonged antiplatelet therapy is still necessary to prevent these complications¹, ². Dual
Materials and Methods

All animal care and experiments were performed following the Basic Guidelines for Conduct of Animal Experiments published by the Ministry of Health, Labor and Welfare, Japan, and were approved by the Institutional Committee for Use of Laboratory Animals of Nihon University School of Medicine (AP15M011).

Animal Preparation

Three-way cross pigs [Landrace, Large White, and Duroc (male, 3–4 months old, and weighing 35–40 kg)], supplied by the National Federation of Agricultural Cooperative Associations (ZEN-NOH, Tokyo, Japan), were orally administered one of the following antithrombotic therapy regimes (for 3 days before stent implantation and until the end of the study): aspirin (81 mg/day) and clopidogrel (75 mg/day) (DAPT group, \(n = 10\)), aspirin (81 mg/day) and rivaroxaban (2 mg/kg/day) (AR group, \(n = 10\)), or clopidogrel (75 mg/day) and rivaroxaban (2 mg/kg/day) (CR group, \(n = 10\)). The study protocol is shown in Fig. 1A. The dose of rivaroxaban was derived from both our preliminary experiments and previous data published by Becker et al.\(^{10}\); the dose of 2 mg/kg/day in pigs is equivalent to the anticoagulant activity of 15 mg/day in humans (data not shown).

After sedation with midazolam (0.5 mg/kg, im), followed by inhaled sevoflurane (5%) and heparinization (5,000 U, ia) to inhibit blood clotting, we performed coronary angiography (CAG) and implanted an everolimus-eluting stent (Promus Element\(^{TM}\), Boston Scientific, Marlborough, MA, USA) in the left anterior descending coronary artery. Stents of 3.0\(^{\text{w}}\)16 mm were deployed with a 12-atm inflation pressure to reach a 1.3:1 stent-to-artery ratio under the guidance of intravascular ultrasound (Opticross catheter and iLab system, Boston Scientific). Follow-up CAG was performed at 1 (1-month observation subgroup, \(n = 5\) each) and 3 months (3-month observation subgroup, \(n = 5\) each) after implantation. After CAG, Optical coherence tomography (OCT) was performed to evaluate the in-stent neointimal formation; before stent deployment and at 1 and 3 months after stent deployment, blood was drawn for analysis.

Aim

When considering bleeding risk, less antithrombotic therapy, such as a dual therapy comprising antiplatelets and DOACs may be preferred at the time of stenting; the duration of antithrombotic therapy should be shortened if the in-stent environments are stabilized during the early phase after DES implantation. However, in-stent tissue responses to the dual therapy comprising antiplatelets and DOACs early after DES implantation are not yet elucidated; moreover, the impact of dual therapy on the in-stent tissue responses compared to DAPT remains unclear. In this study, we examined whether the combination of rivaroxaban (a direct FXa inhibitor) with aspirin or clopidogrel (a P2Y\(_{12}\) receptor antagonist) was noninferior to DAPT regarding in-stent vascular responses after DES implantation in a porcine model, using an intracoronary imaging modality and histological assessment.
computed. Lumen and stent areas were drawn in each analyzed cross-section; neointimal areas were calculated as [stent area–lumen area]. The neointimal thickness was only measured at each stent strut and was determined based on automated measurements performed from the center of the luminal surface of each strut blooming to the lumen contour\(^{13}\).

### Histological Analysis

At 1 and 3 months after stent implantation and after CAG and intracoronary imaging analysis, the pigs were sacrificed. The stented coronary arteries were harvested and perfused with saline and perfusion-fixed with 10% buffered formalin to wash away circulating blood components. As previously described\(^{14,15}\), the stented vessels were embedded in methyl methacrylate resin (FUJIFILM Wako Pure Chemical Industries, Osaka, Japan). Sections (4 µm each) were cut using a cemented tungsten carbide knife (RM2255, Leica, Germany) and stained with hematoxylin and eosin (HE) or Masson’s trichrome (MT). Three sections each (proximal, mid, and distal) were acquired\(^{16}\) with an optical microscope (CX41,
Olympus, Tokyo, Japan) equipped with a DP27 camera (Olympus) and analyzed using the ImageJ software (ver. 1.52, National Institutes of Health, Bethesda, MD, USA)\(^{17}\).

The histological evaluation included measurements of the vessel injury, neointimal area, and fibrin deposition area, the evaluation of neointimal characteristics and inflammation, and the assessment of the incidence of in-stent thrombi. Vessel injury caused by stent deployment was quantified using the vessel injury score as previously reported\(^{18,19}\).

In brief, at each stent strut, vessel wall damage was scored as Grade 0, internal elastic lamina was intact; Grade 1, internal elastic lamina lacerated and media compressed; Grade 2, internal elastic lamina and media lacerated and external elastic lamina compressed; and Grade 3, external elastic lamina was disrupted. As with OCT analysis, the neointimal thickness was only measured at the stent struts. Neointimal areas were measured around the stent and above the inner membrane\(^{20}\), and a morphometric analysis of the smooth muscle cells (SMCs) was performed. Neointimal SMCs were divided into two subtypes: SMCs with contractile features and SMCs with synthetic features\(^{21-23}\). Both subtypes were distinguished according to cell morphology, surrounding extracellular matrix (ECM), and their role in the formation of in-stent restenosis. Similar to cultured SMCs, SMCs with synthetic features in the neointima contain a high number of organelles involved in producing more ECM; however, these are largely replaced by contractile filaments in SMCs with contractile features. The rich contractile filaments comprising SMCs with contractile features are elongated and stain these cells more densely by HE or MT than SMCs with synthetic features. The decreased ECM production also promotes a tight alignment of SMCs with contractile features, while the less stretched SMCs with synthetic features are sparsely alienated between sufficient ECM (Fig.1B)\(^{24-26}\). As shown in Fig.1B and 1C, SMCs with contractile features are located close to the lumen and layered on top of the neointima, whereas those with synthetic features are scattered within the ECM below the layer of SMCs with contractile features, close to the media. The area of the layer of SMCs with contractile features was measured; the relative area percentage of SMCs with contractile features was calculated as the area of SMCs with contractile features divided by the total neointimal area.

Defined by MT staining as a reddish area around the stent struts, the area of fibrin deposition was measured as previously described\(^{27}\). Inflammation around the struts was semi-quantitatively graded via five high-power field observations around the struts per section. Inflammation scores were defined as Grade 0, few inflammatory cells except some foreign body giant cell infiltrations; Grade 1, mild, less than approximately 20 inflammatory cells; Grade 2, moderate, less than approximately 50 inflammatory cells; and Grade 3, severe, more than approximately 50 inflammatory cells per high-power field. In-stent thrombi were determined according to the existence of a heterogeneous composition, such as platelets and fibrin attached to the neointimal wall (arrow in Fig.1D). Heparin was intravenously injected before sacrifice to inhibit blood clotting, and the stented coronary arteries were perfused with saline to wash away circulating blood. Images that were confused with an artifact or blood clots were not included in the measurement.

Statistical Analysis

Data were expressed as mean ± standard error of the mean (SEM) and median [interquartile range] for normally and non-normally distributed variables, respectively. One-way analysis of variance or the Kruskal–Wallis test was performed as appropriate to compare differences across the three groups; post-hoc analyzes with the Steel–Dwass post-hoc analysis or Tukey’s honestly significant difference test were performed to compare differences between groups. A chi-squared test was used to analyze the categorical variables between the three groups. All statistical analyzes were performed with RStudio (Version 1.1.463, RStudio, Inc., Boston, MA, USA), an integrated development environment for R (Version 3.5.2, The R Foundation for Statistical Computing, Vienna, Austria). A p-value of less than 0.05 was considered statistically significant.

Results

Biochemical Analysis

Laboratory data of the study animals are summarized in Table 1. There were no significant differences in initial blood samples among the three groups; in addition, parameters at 1 and 3 months after stent deployment did not differ among the three groups.

OCT Findings

As shown in Fig.2, Fig.3A, and Table 2, compared with a heterogeneous or layered pattern, homogeneous patterns accounted for the majority in all groups at 1 month; in addition, compared with AR and CR groups, the DAPT group consisted of the highest proportions of homogeneous and
Table 1. Laboratory data of study animals per treatment group

|                | DAPT      | AR        | CR        |
|----------------|-----------|-----------|-----------|
|                | Baseline  | 1 month   | 3 months  | Baseline  | 1 month   | 3 months  | Baseline  | 1 month   | 3 months  |
| Total protein (g/dl) | 4.7±0.1   | 5.2±0.2   | 5.1±0.1   | 4.7±0.1   | 5.7±0.3   | 5.2±0.2   | 4.6±0.1   | 4.9±0.1   | 4.9±0.2   |
| Total bilirubin (mg/dl) | 0.05±0.02 | 0.03±0.00 | 0.04±0.01 | 0.04±0.00 | 0.02±0.00 | 0.06±0.02 | 0.07±0.01 | 0.06±0.01 | 0.08±0.02 |
| Triglyceride (mg/dl) | 13.1±1.9  | 18.3±4.1  | 19.0±7.5  | 17.3±2.2  | 21.8±4.4  | 10.0±4.0  | 9.3±2.3   | 8.0±4.1   | 9.0±2.9   |
| Free fatty acid (µEQU) | 263.8±60.7 | 223.2±71.2 | 257.3±114.3 | 116.0±23.5 | 128.3±62.5 | 420.5±189.4 | 189.6±51.0 | 245.8±48.8 | 242.2±54.5 |
| Total cholesterol (mg/dl) | 77.0±4.8 | 65.7±4.8 | 80.0±5.8 | 76.0±1.8 | 84.5±6.7 | 73.5±2.0 | 75.6±6.5 | 73.5±4.7 | 61.5±1.4 |
| Free type cholesterol (mg/dl) | 14.3±1.0 | 12.8±1.8 | 15.8±1.1 | 15.5±0.5 | 16.0±1.1 | 13.5±0.9 | 14.1±1.0 | 15.5±1.3 | 12.3±0.2 |
| Blood urea nitrogen (mg/dl) | 10.3±1.0 | 10.7±0.6 | 9.9±1.3 | 9.7±0.6 | 10.4±0.8 | 9.1±0.3 | 8.4±0.7 | 7.7±0.3 | 7.6±0.9 |
| Creatine (mg/dl) | 1.05±0.08 | 1.46±0.13 | 1.92±0.03 | 1.10±0.09 | 1.19±0.11 | 1.78±0.22 | 1.35±0.15 | 1.77±0.18 | 2.05±0.06 |
| Sodium (mEQU/L) | 145.5±0.7 | 143.8±0.7 | 144.3±1.8 | 145.2±1.8 | 143.3±0.9 | 143.0±2.2 | 146.1±0.5 | 145.8±0.6 | 140.8±1.4 |
| Chloride (mEQU/L) | 106.4±2.0 | 104.8±1.6 | 106.8±2.0 | 105.9±1.2 | 106.5±0.9 | 104.3±3.1 | 107.9±2.1 | 112.3±1.1 | 103.3±2.5 |
| Potassium (mEQU/L) | 4.0±0.1 | 3.8±0.2 | 4.1±0.3 | 4.0±0.1 | 4.1±0.2 | 4.1±0.1 | 4.1±0.2 | 3.9±0.1 | 4.1±0.3 |
| AST (IU/L) | 20.0±1.2 | 20.3±1.8 | 22.5±1.7 | 20.8±1.9 | 22.5±1.6 | 26.8±3.8 | 23.6±2.4 | 26.5±4.6 | 26.8±2.6 |
| ALT (IU/L) | 31.0±2.0 | 24.7±1.4 | 33.5±1.6 | 26.7±3.6 | 27.0±1.2 | 28.8±2.4 | 29.8±2.6 | 28.0±1.4 | 26.5±1.9 |
| LDH (IU/L) | 537.0±30.9 | 513.5±44.1 | 567.5±17.5 | 478.2±28.5 | 466.0±30.3 | 554.0±64.2 | 484.0±25.0 | 580.8±78.8 | 481.2±43.0 |
| ALP (IU/L) | 441.1±3.6 | 541.2±65.0 | 364.3±28.1 | 455.2±38.2 | 382.0±52.9 | 416.8±28.1 | 482.9±22.5 | 422.5±51.5 | 351.0±25.9 |

Data are presented as mean±standard error of the mean. AST, aspartate aminotransferase; ALT, alanine aminotrasferase; LDH, lactate dehydrogenase; ALP, alkaline phosphoatase; DAPT, dual antiplatelet therapy (consisting of aspirin and clopidogrel); AR, aspirin and rivaroxaban; CR, clopidogrel and rivaroxaban.

heterogeneous patterns and the lowest proportion of layered patterns. At 3 months, the proportions of neointimal patterns were comparable among the three groups. With regard to neointimal thickness at the stent struts, the DAPT group showed the lowest neointimal thickness at 1 month; however, at 3 months, this had increased to be comparable with the other two groups. In the AR and CR groups, the neointimal thickness was similar between 1 and 3 months (Table 3). Similar to the neointimal thickness, the DAPT group also showed the smallest neointimal area at 1 month (Fig.3E and Table 2). At 3 months, although the neointimal area values were similar in the three groups, the statistical analysis revealed a significantly larger area in the CR group than in the DAPT group (2.48 mm² in CR versus 2.44 mm² in DAPT, p<0.045). In the AR and CR groups, the neointimal area was similar between 1 and 3 months (Fig.3E, 3F, and Table 2).

Histological Findings

Vessel injury scores were comparable among the three groups at both 1 and 3 months (Table 3). The neointimal area was comparable among the three groups at 1 month; however, this value was smallest in the CR group at 3 months (Fig. 5A, 5B, and Table 3). The neointimal area in the DAPT and AR groups increased from 1 to 3 months, whereas in the CR group remained similar.

The proportion of SMCs with contractile features, which indicates stable status, was significantly greater in the CR group at both 1 and 3 months (Fig.4, Fig.5C-D, and Table 3). Fibrin deposition around the stent struts tended to be greater in the AR and CR groups at 1 month and comparable among the three groups at 3 months (Table 3). The semi-quantitative inflammation score tended to be lower in the CR group at 1 month and comparable among the three groups at 3 months (Table 3). In-stent thrombi were more frequent in the AR at 1 month, whereas the incidence between CR and DAPT, as well as CR and AR, was comparable (Fig.5E); however, no in-stent thrombi were seen in any of the three groups at 3 months.

Discussion

In this study, we demonstrated the in-stent responses to a dual therapy of antiplatelet drugs and the FXa inhibitor rivaroxaban after DES implantation using intracoronary imaging analysis and histological examination. Our study revealed evidence supporting the noninferiority of combination therapy using rivaroxaban with an antiplatelet drug, particularly the dual therapy of clopidogrel and rivaroxaban, compared with traditional aspirin/clopidogrel therapy after DES implantation in a porcine coronary artery. Therefore, our results provide pathological evidence verifying the efficacy of the combination of a P2Y12 receptor antagonist with the direct FXa inhibitor rivaroxaban.
made this therapy more effective than dual therapy using aspirin and rivaroxaban (AR group). Dual therapy using aspirin and rivaroxaban did not show sufficient antithrombotic effects when compared with DAPT; this indicates that a combination of aspirin and OAC is not sufficient, as previously reported\(^3\). On the other hand, CR treatment was similar to DAPT and could suppress in-stent thrombi. This may be due to the strong antiplatelet effect of clopidogrel as an adenosine diphosphate-P2Y\(_{12}\) receptor antagonist\(^3\). Furthermore, the anti-inflammatory effect of clopidogrel might contribute to the tendency toward a lower inflammation score at 1 month in the CR group\(^3\). Especially, these results indicate that the dual therapy of clopidogrel and rivaroxaban might be an effective antithrombotic therapy after DES implantation.

A previous porcine study revealed that neointimal growth reached a peak at 3 months after coronary artery DES implantation; this was maintained until 6 months, after which, neointimal growth began to decline\(^3\). In our DES implantation model, the neointima had fully covered the surface of the DES at both 1 and 3 months in all three groups.
However, the patterns of vascular response after DES implantation were somewhat different among the three groups. Despite inconsistencies between OCT and pathological results regarding the measurement of the intimal thickness and area, both OCT and pathological results indicated a different time course for neointimal growth between DAPT and AR or CR treatments. In the DAPT group, as reported in a previous study, the neointima continued to grow from 1 to 3 months. However, in the CR group, the neointima grew rapidly and reached its peak within 1 month; subsequently, the neointima almost stopped growing and was maintained until 3 months (Fig. 3, Fig. 5, Table 2, and Table 3). The shortened time course of neointimal growth may be favorable for rapid neointimal stability and quick vascular healing. The inconsistencies between the OCT and pathological results regarding the measurements of the intimal thickness and area might be due to the small sample size, resolution of OCT, and distortion of the tissue–stent interface during pathological processing.

As the neointimal thickness was measured only at the stent struts, a lack of information on neointimal hyperplasia between struts makes its evaluation function different from that of the neointimal area. The DAPT group showed a different distribution of the three neointimal OCT pattern types between 1 and 3 months, while there was a similar distribution between 1 and 3 months in the AR and CR groups. This was consistent with the abovementioned data.
After DES implantation: SMCs with contractile features and SMCs with synthetic features. With less proliferation, lower ECM production, and rich, tightly aligned contractile filaments, SMCs with contractile features are considered mature and stable\(^2\) 4, playing a beneficial role after DES implantation. The proportionally greater number of SMCs with contractile features in the CR group at both 1 and 3 months indicated a rapid and more stabilized in-stent neointimal environment using a dual therapy of clopidogrel and rivaroxaban.

Recent clinical trials have demonstrated that rivaroxaban reduces mortality and cardiovascular events, including stent thrombosis, in patients with neointimal growth and indicated different time courses for vascular response between DAPT and CR or AR treatments. Since the heterogeneous pattern is thought to be linked to poor long-term prognoses\(^4\)-\(^6\)-\(^2\), a lower proportion of heterogeneous patterns in AR and CR might indicate a beneficial effect. The high proportion of layered patterns in the AR and CR groups at 1 month might reflect a clearer boundary between SMCs with contractile and synthetic features.

SMC heterogeneity has been described in atherosclerotic lesions and restenosis after angioplasty or stenting\(^2\)-\(^2\). Consistent with previous studies, two types of SMCs were distinguished in the neointima after DES implantation: SMCs with contractile features and SMCs with synthetic features. With less proliferation, lower ECM production, and rich, tightly aligned contractile filaments, SMCs with contractile features are considered mature and stable\(^2\) 4, playing a beneficial role after DES implantation. The proportionally greater number of SMCs with contractile features in the CR group at both 1 and 3 months indicated a rapid and more stabilized in-stent neointimal environment using a dual therapy of clopidogrel and rivaroxaban.

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### Table 2. Comparison of OCT findings between three groups

|                     | DAPT          | AR            | CR            | Overall   | p-value |
|---------------------|---------------|---------------|---------------|-----------|---------|
| At 1 month          |               |               |               |           |         |
| Mean neointimal     | 0.23 [0.20, 0.28] | 0.30 [0.24, 0.47] | 0.34 [0.26, 0.46] | 0.000     |         |
| area (mm²)          | 1.92 [1.65, 2.17] | 2.40 [2.04, 3.40] | 2.48 [1.93, 3.16] | 0.000     |         |
| Homogeneous (%)     | 81.1          | 58.6          | 51.1          | <0.001    |         |
| Heterogeneous (%)   | 11.3          | 0.9           | 4.3           | 0.000     | NS      |
| Layered (%)         | 7.5           | 40.5          | 44.7          | 0.000     | NS      |
| At 3 months         |               |               |               |           |         |
| Mean neointimal     | 0.33 [0.28, 0.44] | 0.32 [0.25, 0.41] | 0.35 [0.29, 0.42] | 0.301     | NA      |
| area (mm²)          | 2.44 [2.13, 3.19] | 2.52 [2.07, 3.13] | 2.48 [1.79, 2.92] | 0.021     | 0.459   |
| Homogeneous (%)     | 66.3          | 60.2          | 50.8          | 0.113     | NA      |
| Heterogeneous (%)   | 4.0           | 2.2           | 2.3           | 0.004     | NA      |
| Layered (%)         | 29.7          | 37.6          | 46.9          | 0.462     | NA      |

Data are presented as median [interquartile range] or number. OCT, optical coherence tomography; DAPT, dual-antiplatelet therapy (consisting of aspirin and clopidogrel); AR, aspirin and rivaroxaban; CR, clopidogrel and rivaroxaban.

### Table 3. Comparison of histological findings between three groups

|                     | DAPT          | AR            | CR            | Overall   | p-value |
|---------------------|---------------|---------------|---------------|-----------|---------|
| At 1 month          |               |               |               |           |         |
| Neointimal area     | 1.75 [1.52, 2.16] | 1.81 [1.59, 2.21] | 2.13 [1.77, 2.96] | 0.398     | NA      |
| (mm²)               | 33.9 ± 3.1    | 12.3 ± 2.4    | 43.2 ± 4.2    | 0.000     | 0.000   |
| Fibrin deposition   | 1971.4 [1767.3, 2922.8] | 3771.0 [2047.3, 6102.2] | 3895.2 [2957.6, 5834.4] | 0.059     | NA      |
| area (µm²)          | 1.07 ± 0.08   | 1.33 ± 0.14   | 0.91 ± 0.13   | 0.054     | NA      |
| Inflammation score  | 0.95 ± 0.07   | 0.87 ± 0.08   | 1.02 ± 0.09   | 0.462     | NA      |
| Vessel injury score |               |               |               |           |         |
| At 3 months         |               |               |               |           |         |
| Neointimal area     | 2.47 [2.42, 2.89] | 2.32 [1.92, 2.83] | 2.05 [1.80, 2.29] | 0.016     | 0.401   |
| (mm²)               | 47.5 ± 3.1    | 33.2 ± 1.9    | 60.9 ± 5.3    | 0.000     | 0.002   |
| Fibrin deposition   | 273.3 [56.9, 520.1] | 15.7 [0.0, 106.3] | 93.0 [0.0, 260.1] | 0.137     | NA      |
| area (µm²)          | 1.04 ± 0.13   | 1.08 ± 0.12   | 1.32 ± 0.11   | 0.209     | NA      |
| Inflammation score  | 1.17 ± 0.07   | 1.08 ± 0.07   | 1.25 ± 0.08   | 0.314     | NA      |
| Vessel injury score |               |               |               |           |         |

Data are presented as mean ± standard error of the mean or median [interquartile range]. DAPT, dual antiplatelet therapy (consisting of aspirin and clopidogrel); AR, aspirin and rivaroxaban; CR, clopidogrel and rivaroxaban; cSMC, contractile smooth muscle cell.
atherosclerosis; therefore, the reaction to antiplatelet drugs and anticoagulants may be different in actual atherosclerotic lesions. However, as our results reflected a fundamental vessel response after mechanical injury with DES implantation, they are considered a meaningful examination of the effect of antiplatelet drugs and DOACs on neointimal progression and thrombus formation. In this study, given that only a single dose of rivaroxaban could be tested, the dose–response relationship for rivaroxaban in a swine cardiovascular system remains unclear. Since the biological responses between pigs and humans are different (more specifically, the reaction in pigs is more rapid than that in humans\(^48\)), our results cannot precisely predict the optimal duration of antithrombotic therapy in clinical practice. It would, therefore, be meaningful to perform long-term observations that exceed 6 months, to verify the peak of neointimal hyperplasia and prove the noninferiority or benefit of CR or AR treatment regarding neointimal growth and vascular healing. However, the acute coronary syndrome\(^4, 43, 44\). A previous study showed that aspirin and rivaroxaban therapy could lead to better cardiovascular outcomes than aspirin alone\(^49\). Moreover, combination therapy using P2Y\(_{12}\) receptor antagonists and FXa inhibitors may have more benefits related to clinical efficacy\(^46, 47\). Our study is consistent with previous clinical results and provides supporting evidence of a pathological vascular response for the results of these clinical trials. The shortened time course of neointimal growth and the increased number of mature/stable SMC with CR dual therapy indicated quick vascular healing and might contribute to the shortening of the administration period of antiplatelet drugs and DOACs.

**Study Limitations**

Our study has some limitations; this study was performed using a healthy coronary animal model. In brief, we implanted a stent in a healthy coronary artery and not in one that had developed atherosclerosis; therefore, the reaction to antiplatelet drugs and anticoagulants may be different in actual atherosclerotic lesions. However, as our results reflected a fundamental vessel response after mechanical injury with DES implantation, they are considered a meaningful examination of the effect of antiplatelet drugs and DOACs on neointimal progression and thrombus formation. In this study, given that only a single dose of rivaroxaban could be tested, the dose–response relationship for rivaroxaban in a swine cardiovascular system remains unclear. Since the biological responses between pigs and humans are different (more specifically, the reaction in pigs is more rapid than that in humans\(^48\)), our results cannot precisely predict the optimal duration of antithrombotic therapy in clinical practice. It would, therefore, be meaningful to perform long-term observations that exceed 6 months, to verify the peak of neointimal hyperplasia and prove the noninferiority or benefit of CR or AR treatment regarding neointimal growth and vascular healing. However, the
DES implantation. The patterns of vascular response were distinct, depending on the different drug combinations. The shortened time course of neointimal growth and the predominantly stable type of SMCs with CR dual therapy indicated quick vascular healing and may contribute to the shortening of the administration period for antiplatelet drugs and DOACs.

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**References**

1) Singh I, Shafiq N, Pandhi P, Reddy S, Pattnaik S, Sharma Y, Malhotra S: Triple antiplatelet therapy vs. dual antiplatelet therapy in patients undergoing percutaneous coronary intervention: an evidence-based approach to answering a clinical query. Br J Clin Pharmacol, 2009; 68: 4-13

2) Harper RW: Drug-eluting coronary stents--a note of caution. Med J Aust, 2007; 186: 253-255

3) Dewilde WJM, Oirbans T, Verheugt FWA, Kelder JC, De Smet BJGL, Herrman J-P, Adriaenssens T, Vrolix M, Heestermans AACM, Vis MM, Tijsen JGP, van't Hof AW, ten Berg JM, WOEST study investigators: Use of clopidogrel with or without aspirin in patients taking oral anticoagulant therapy and undergoing percutaneous coronary intervention: an open-label, randomised, controlled trial. Lancet, 2013; 381: 1107-1115

4) Mega JL, Braunwald E, Wiviott SD, Bassand J-P, Bhatt DL, Bode C, Burton P, Cohen M, Cook-Brunn N, Fox KAA, Goto S, Murphy SA, Plotnikov AN, Schneider D, Sun X, Verheugt FWA, Gibson CM, ATLAS ACS 2–TIMI 51 Investigators: Rivaroxaban in patients with a recent acute coronary syndrome. N Engl J Med, 2012; 366: 9-19

5) Gibson CM, Pinto DS, Chi G, Arbetter D, Yee M, Mehran R, Bode C, Halperin J, Verheugt FWA, Wildgoose P, Burton P, van Eickels M, Korjiaen S, Daaboul Y, Jain P, Lip GYH, Cohen M, Peterson ED, Fox KAA: Recurrent hospitalization among patients with atrial fibrillation undergoing intracorony stenting treated with 2 treatment strategies of rivaroxaban or a dose-adjusted oral vitamin K antagonist treatment strategy. Circulation, 2017; 135: 323-333

6) Cannon CP, Bhatt DL, Oldgren J, Lip GYH, Ellis SG, Kimura T, Maeng M, Merkely B, Zeymer U, Gropper S, Nordaby M, Kleine E, Harper R, Manassie J, Januzzi JL, ten Berg JM, Steg PG, Hohnloser SH, RE-DUAL PCI Steering Committee and Investigators: Dual antithrombotic therapy with dabigatran after PCI in atrial fibrillation. N Engl J Med, 2017; 377: 1513-1524

7) Gibson CM, Mehran R, Bode C, Halperin J, Verheugt FW, Wildgoose P, Birmingham M, Janus J, Burton P, van Eickels M, Korjiaen S, Daaboul Y, Lip GYH, Cohen M, Husted S, Peterson ED, Fox KA: Prevention of bleeding in patients with atrial fibrillation undergoing PCI. N Engl J Med, 2016; 375: 2423-2434

8) Lopes RD, Heizger G, Aronson R, Vora AN, Massaro T, Mehran R, Goodman SG, Windicke S, Darius H, Li J, Averkov O, Bahit MC, Berwanger O, Budaj A, Hijazi Z, Parkhomenko A, Sinnaepe A, Storey RF, Thiele H, Vinereanu D, Granger CB, Alexander JH, AUGUSTUS Investigators: Antithrombotic therapy after acute coronary syndrome or PCI in atrial fibrillation. N Engl J Med, 2019; 380: 1509-1524

9) Lopes RD, Hong H, Harshkan RE, Bhatt DL, Mehran R, Cannon CP, Granger CB, Verheugt FWA, Li J, ten Berg JM, Sarofoff N, Gibson CM, Alexander JH: Safety and efficacy of antithrombotic strategies in patients with atrial fibrillation undergoing percutaneous coronary intervention: A network meta-analysis of randomized controlled trials. JAMA Cardiol, 2019; 4: 747-755

10) Becker EM, Perzborn E, Klipp A, Lücker U, Bütehorn U, Kast R, Badimon JJ, Lax V: Effects of rivaroxaban, acetylsalicylic acid and clopidogrel as monotherapy and in combination in a porcine model of stent thrombosis. J Thromb Haemost, 2012; 10: 2470-2480

11) Kubo T, Akasaka T, Kozuma K, Kimura K, Kawamura M, Sumiyoshi T, Ino Y, Morino Y, Tanabe K, Kadota K, Kimura T, RESET Investigators: Comparison of neointimal coverage between everolimus-eluting stents and sirolimus-eluting stents: an optical coherence tomography substudy of the RESET (Randomized Evaluation of Sirolimus-eluting versus Everolimus-eluting stent Trial). EuroIntervention, 2015; 11: 564-571

12) Kim J-S, Afari ME, Ha J, Tellez A, Milewski K, Conditt G, Cheng Y, Hua Yi G, Kaluza GL, Granada JF, Thiele H, Parkhomenko A, Sinnaepe P, Storey RF, Thiele H, Vinereanu D, Granger CB, Alexander JH, AUGUSTUS Investigators: Antithrombotic therapy after acute coronary syndrome or PCI in atrial fibrillation. N Engl J Med, 2017; 377: 1513-1524

13) Tanigawa J, Barlis P, Di Mario C: Intravascular optical coherence tomography: optimisation of image acquisition and quantitative assessment of stent strut apposition. EuroIntervention, 2007; 3: 128-136

14) Malik N, Gunn J, Holt C, Shepherd L, Francis S, Newman C, Crossman D, Cumberland D: Intravascular stents: a new technique for tissue processing for histology,
immunohistochemistry, and transmission electron microscopy. Heart, 1998; 80: 509-516
15) Rippstein P, Black MK, Boivin M, Veinot JP, Ma X, Chen Y-X, Human P, Zilla P, O’Brien ER: Comparison of processing and sectioning methodologies for arteries containing metallic stents. J Histochem Cytochem, 2006; 54: 673-681
16) Schwartz RS, Edelman ER, Carter A, Chronos N, Rogers C, Robinson KA, Waksman R, Weinberger J, Wilensky RL, Jensen DN, Zuckerman BD, Virmani R, Consensus Committee: Drug-eluting stents in preclinical studies: recommended evaluation from a consensus group. Circulation, 2002; 106: 1867-1873
17) Schneider CA, Rashband WS, Eliceiri KW: NIH Image to Image: 25 years of image analysis. Nat Methods, 2012; 9: 671-675
18) Schwartz RS, Huber KC, Murphy JG, Edwards WD, Camrud AR, Vlietstra RE, Holmes DR: Restenosis and the proportional neointimal response to coronary artery injury: Results in a porcine model. J Am Coll Cardiol, 1992; 19: 267-274
19) Hofma SH, Whelan DMC, van Beusekom HMM, Verdouw PD, van der Giessen WJ: Increasing arterial wall injury after long-term implantation of two types of stent in a porcine coronary model. Eur Heart J, 1998; 19: 601-609
20) Mori M, Sakata K, Nakanishi C, Nakahashi T, Kawashiri M, Yoshioka K, Takuwa Y, Okada H, Yokawa J, Shimojima M, Yoshimuta T, Yoshida S, Yamagishi M, Hayashi K: Early endothelialization associated with a biolimus A9 bioreabsorbable polymer stent in a porcine coronary model. Heart Vessels, 2017; 32: 1244-1252
21) Orlandi A, Ehrlich HP, Ropraz P, Spagnoli L G, Gabbiani G: Rat aortic smooth muscle cells isolated from different layers and at different times after endothelial denudation show distinct biological features in vitro. Arterioscler Thromb, 1994; 14: 982-989
22) Hao H, Gabbiani G, Bochaton-Piallat ML: Arterial smooth muscle cell heterogeneity: implications for atherosclerosis and restenosis development. Arterioscler Thromb Vasc Biol, 2003; 23: 1510-1520
23) Escuer J, Martinez MA, McGinty S, Peña E: Mathematical modelling of the restenosis process after stent implantation. J R Soc Interface, 2019; 16: 20190313
24) Rensen SSM, Doevendans PAFM, van Eys GJJM: Regulation and characteristics of vascular smooth muscle cell phenotypic diversity. Neth Heart J, 2007; 15: 100-108
25) Turley EA: Extracellular matrix remodeling: Multiple paradigms in vascular disease. Circ Res, 2001; 88: 2-4
26) Sugita S, Mizutani E, Hozaki M, Nakamura M, Matsumoto T: Photoelasticity-based evaluation of cellular contractile force for phenotypic discrimination of vascular smooth muscle cells. Sci Rep, 2019; 9: 3960
27) Wakamati R, Hao H, Imanaka T, Shibuya M, Ueda Y, Tsujimoto M, Ishibashi-Ueda H, Hirota S: Initial pathological responses of second-generation everolimus-eluting stents implantation in Japanese coronary arteries: Comparison with first-generation sirolimus-eluting stents. J Cardiol, 2018; 71: 452-457
28) Costa MA, Simon DI: Molecular basis of restenosis and drug-eluting stents. Circulation, 2005; 111: 2257-2273
29) Pearson JD: Endothelial cell function and thrombosis. Baillieres Clin Haematol, 1994; 7: 441-452
30) Toutouzas K, Colombo A, Stefanadis C: Inflammation and restenosis after percutaneous coronary interventions. Eur Heart J, 2004; 25: 1679-1687
31) Chandrasekar B, Tanguay JF: Platelets and restenosis. J Am Coll Cardiol, 2000; 35: 555-562
32) Ragosta M, Gimple LW, Gertz SD, Dunwiddie CT, Vlaskus GP, Haber HL, Powers ER, Roberts WC, Sarembock IJ: Specific factor Xa inhibition reduces restenosis after balloon angioplasty of atherosclerotic femoral arteries in rabbits. Circulation, 1994; 89: 1262-1271
33) Kaiser B, Painz M, Scholz O, Kunitada S, Fareed J: A synthetic inhibitor of factor Xa, DX-9065a, reduces proliferation of vascular smooth muscle cells in vivo in rats. Thromb Res, 2000; 98: 175-185
34) Cheung WM, D’Andrea MR, Andrade-Gordon P, Damiano BP: Altered vascular injury responses in mice deficient in protease-activated receptor-1. Arterioscler Thromb Vasc Biol, 1999; 19: 3014-3024
35) Damiano BP, D’Andrea MR, de Garavilla L, Cheung WM, Andrade-Gordon P: Increased expression of protease activated receptor-2 (PAR-2) in balloon-injured rat carotid artery. Thromb Haemost, 1999; 81: 808-814
36) Leon MB, Baim DS, Popma JJ, Gordon PC, Cutlip DE, Ho KK, Giambartolomei A, Diver DJ, Lasorda DM, Williams DO, Pocock SJ, Kuntz RE: A clinical trial comparing three antithrombotic–drug regimens after coronary-artery stenting. N Engl J Med, 1998; 339: 1665-1671
37) Dorsam RT, Murugappan S, Ding Z, Kunapuli SP: Clopidogrel: interactions with the P2Y12 receptor and clinical relevance. Hematology, 2003; 8: 359-365
38) Li M, Zhang Y, Ren H, Zhang Y, Zhu X: Effect of clopidogrel on the inflammatory progression of early atherosclerosis in rabbits model. Atherosclerosis, 2007; 194: 348-356
39) Virmani R, Kolodgie FD, Farb A, Lafont A: Drug eluting stents: are human and animal studies comparable? Heart, 2003; 89: 133-138
40) Kilickesmez K, Dall’Ara G, Rama-Merchan JC, Ghione E, Notarniccolo S, Zuckerman BD, Virmani R, Consensus Committee: Drug-eluting stents in preclinical studies: are human and animal studies comparable? Heart, 2003; 194: 348-356
41) Shi S-Y, Chen K-L, Gu J, Xu C, Chen Q-R, Chen Y-Q, Li M, Zhang Y, Ren H, Zhang Y, Zhu X: Effect of clopidogrel on the inflammatory progression of early atherosclerosis in rabbits model. Atherosclerosis, 2007; 194: 348-356
42) Kilickesmez K, Dall’Ara G, Rama-Merchan JC, Ghione E, Notarniccolo S, Zuckerman BD, Virmani R, Consensus Committee: Drug-eluting stents in preclinical studies: are human and animal studies comparable? Heart, 2003; 89: 133-138
43) Shi S-Y, Chen K-L, Gu J, Xu C, Chen Q-R, Chen Y-Q, Li M, Zhang Y, Ren H, Zhang Y, Zhu X: Effect of clopidogrel on the inflammatory progression of early atherosclerosis in rabbits model. Atherosclerosis, 2007; 194: 348-356
44) Streeter J, Narula J, Narula S, Abu-Yousef M, Chen Q-R, Li M, Zhang Y, Ren H, Zhang Y, Zhu X: Effect of clopidogrel on the inflammatory progression of early atherosclerosis in rabbits model. Atherosclerosis, 2007; 194: 348-356
45) Virmani R, Kolodgie FD, Farb A, Lafont A: Drug eluting stents: are human and animal studies comparable? Heart, 2003; 89: 133-138
46) Kilickesmez K, Dall’Ara G, Rama-Merchan JC, Ghione E, Notarniccolo S, Zuckerman BD, Virmani R, Consensus Committee: Drug-eluting stents in preclinical studies: are human and animal studies comparable? Heart, 2003; 89: 133-138
47) Shi S-Y, Chen K-L, Gu J, Xu C, Chen Q-R, Chen Y-Q, Li M, Zhang Y, Ren H, Zhang Y, Zhu X: Effect of clopidogrel on the inflammatory progression of early atherosclerosis in rabbits model. Atherosclerosis, 2007; 194: 348-356
48) Kilickesmez K, Dall’Ara G, Rama-Merchan JC, Ghione E, Notarniccolo S, Zuckerman BD, Virmani R, Consensus Committee: Drug-eluting stents in preclinical studies: are human and animal studies comparable? Heart, 2003; 89: 133-138
49) Shi S-Y, Chen K-L, Gu J, Xu C, Chen Q-R, Chen Y-Q, Li M, Zhang Y, Ren H, Zhang Y, Zhu X: Effect of clopidogrel on the inflammatory progression of early atherosclerosis in rabbits model. Atherosclerosis, 2007; 194: 348-356
43) Gibson CM, Chakrabarti AK, Mega J, Bode C, Bassand J-P, Verheugt FWA, Bhatt DL, Goto S, Cohen M, Mohanavelu S, Burton P, Stone G, Braunwald E, ATLAS-ACS 2 TIMI 51 Investigators: Reduction of stent thrombosis in patients with acute coronary syndromes treated with rivaroxaban in ATLAS-ACS 2 TIMI 51. J Am Coll Cardiol, 2013; 62: 286-290

44) Ohman EM, Roe MT, Steg PG, James SK, Povsic TJ, White J, Rockhold F, Plotnikov A, Mundl H, Strony J, Sun X, Husted S, Tendera M, Montalescot G, Bahit MC, Ardissino D, Bueno H, Claesys MJ, Nicolau JC, Cornel JH, Goto S, Kiss RG, Güray U, Park D-W, Bode C, Welsh RC, Gibson CM: Clinically significant bleeding with low-dose rivaroxaban versus aspirin, in addition to P2Y12 inhibition, in acute coronary syndromes (GEMINI-ACS-1): a double-blind, multicentre, randomised trial. Lancet, 2017; 389: 1799-1808

45) Eikelboom JW, Connolly SJ, Bosch J, Dagenais GR, Hart RG, Shestakovska O, Diaz R, Alings M, Lonn EM, Anand SS, Widimsky P, Hori M, Avezum A, Piegaas LS, Branch KRH, Probstfield J, Bhatt DL, Zhu J, Liang Y, Maggioni AP, Lopez-Jaramillo P, O'Donnell M, Kakkar AK, Fox KAA, Parkhomenko AN, Ersl G, Störk S, Keltai M, Ryden L, Pogosova N, Dans AL, Lanas F, Commerford PJ, Torp-Pedersen C, Guzik TJ, Verhamme PB, Vinereanu D, Kim J-H, Tonkin AM, Lewis BS, Felix C, Yusoff K, Steg PG, Metsarinne KP, Cook Bruns N, Misselwitz F, Chen E, Leong D, Yusuf S, COMPASS Investigators: Rivaroxaban with or without aspirin in stable cardiovascular disease. N Engl J Med, 2017; 377: 1319-1330

46) Gurbel PA, Tantry US: GEMINI-ACS-1: toward unearthing the antithrombotic therapy cornerstone for acute coronary syndromes. Lancet, 2017; 389: 1773-1775

47) Yasuda S, Kaikita K, Akao M, Ako J, Matoba T, Nakamura M, Miyauchi K, Hagiwara N, Kimura K, Hirayama A, Matsui K, Ogawa H, AFIRE Investigators: Antithrombotic therapy for atrial fibrillation with stable coronary disease. N Engl J Med, 2019; 381: 1103-1113

48) Chaabane C, Otsuka F, Virmani R, Bochaton-Piallat ML: Biological responses in stented arteries. Cardiovasc Res, 2013; 99: 353-363