Pharmacodynamic Effects of Vorapaxar in Patients With and Without Diabetes Mellitus

Results of the OPTIMUS-5 Study

Francesco Franchi, MD, Fabiana Rollini, MD, Victor Kairouz, MD, Jose Rivas Rios, MD, Andrea Rivas, MD, Malhar Agarwal, MD, Maryuri Briceno, MD, Mustafa Wali, MD, Ahmed Nawaz, MD, Gabriel Silva, MD, Zubair Shaikh, MD, Naji Maaliki, MD, Latonya Been, AAS, Jason Piraino, DPM, Andres M. Pineda, MD, Malhar Agarwal, MD, Daniel Soffer, MD, Martin M. Zenni, MD, Lisa K. Jennings, PhD, Theodore A. Bass, MD, Dominick J. Angiolillo, MD, PhD

JACC: BASIC TO TRANSLATIONAL SCIENCE CME/MOC/ECME

This article has been selected as this month’s JACC: Basic to Translational Science CME/MOC/ECME activity, available online at http://www.acc.org/jacc-journals-cme by selecting the JACC Journals CME/MOC/ECME tab.

Accreditation and Designation Statement

The American College of Cardiology Foundation (ACCF) is accredited by the Accreditation Council for Continuing Medical Education (ACCME) and the European Board for Accreditation in Cardiology (EBAC) to provide continuing medical education for physicians. The ACCF designates this Journal-based CME/MOC/ECME activity for a maximum of 1 AMA PRA Category 1 Credit or 1 EBAC Credit. Physicians should only claim credit commensurate with the extent of their participation in the activity.

Successful completion of this CME activity, which includes participation in the evaluation component, enables the participant to earn up to 1 AMA PRA Category 1 Credit or 1 EBAC Credit. Physicians should only claim credit commensurate with the extent of their participation in the activity.

Method of Participation and Receipt of CME/MOC/ECME Certificate

To obtain credit for JACC: Basic to Translational Science CME/MOC/ECME, you must:
1. Be an ACC member or JACBTS subscriber.
2. Carefully read the CME/MOC/ECME-designated article available online and in this issue of the journal.
3. Answer the post-test questions. At least 2 questions provided must be answered correctly to obtain credit.
4. Complete a brief evaluation.
5. Claim your CME/MOC/ECME credit and receive your certificate electronically by following the instructions given at the conclusion of the activity.

CME/MOC/ECME Objective for This Article: Upon completion of this activity, the learner should be able to: 1) compare the mechanism of action of various antiplatelet agents used for the management of acute coronary syndromes; 2) identify the indications and contraindications for antiplatelet agents used in acute coronary syndromes; and 3) discuss the benefits and risks of preventative aspirin in diabetic patients.

CME/MOC/ECME Editor Disclosure: CME/MOC/ECME Editor L. Kristin Newby, MD, is supported by research grants from Amylin, Bristol-Myers Squibb Company, GlaxoSmithKline, Sanofi, Verily Life Sciences (formerly Google Life Sciences), the MURDOCK Study, NIH, and PCORI; receives consultant fees/honoraria from BioKier, DemeRx, MedScope/TheHeart.org, Metanomics, Philips Healthcare, Roche Diagnostics, CMAC Health Education & Research Institute; serves as an Officer, Director, Trustee, or other fiduciary role for the AstraZeneca HealthCare Foundation and the Society of Chest Pain Centers (now part of ACC); and serves in another role for the American Heart Association and is the Deputy Editor of JACC: Basic to Translational Science.

Author Disclosures: The study was originally funded by a grant from Merck until transfer of marketing rights of vorapaxar, including grant responsibilities, to Arazel. At the time of transfer, Merck had funded 60% of overall grant costs. Following such transfer, Arazel filed for bankruptcy and did not cover any of the residual grant costs, which was completed using research funds from the Division of Cardiology, University of Florida College of Medicine–Jacksonville. Upon completion of the study, Deerfield acquired marketing rights of vorapaxar and provided a nominal fee for the final phases of investigator-initiated research studies conducted with vorapaxar at our institution (NCT02548650 and NCT02545933). Dr. Franchi has received payment as an individual for consulting fees or honoraria from AstraZeneca and Sanofi. Dr. Rollini has received payment as an individual for consulting fees or honoraria from Chiesi. Dr. Angiolillo has received payment as an
individual for consulting fees or honoraria from Amgen, Arazel, AstraZeneca, Bayer, Biosensors, Boehringer Ingelheim, Bristol-Myers Squibb, Chiesi, Daiichi-Sankyo, Eli Lilly, Haemonetics, Janssen, Merck, PhaseBio, PLx Pharma, Pfizer, Sanofi, and The Medicines Company; participation in review activities from CeloNova and St. Jude Medical; and institutional payments for grants from Amgen, AstraZeneca, Bayer, Biosensors, CeloNova, CSL Behring, Daiichi-Sankyo, Eisai, Eli Lilly, Gilead, Idorsia, Janssen, Matsutani Chemical Industry Co., Merck, Novartis, Osprey Medical, and Renal Guard Solutions. Dr. Jennings has received payments as an individual for consulting fees or honoraria from Bayer, Janssen, PhaseBio, and Portola; and institutional payments for grants from Janssen and In Motion Musculoskeletal Institute. All other authors have reported that they have no relationships relevant to the contents of this paper to disclose.

Medium of Participation: Online (article and quiz).

CME/MOC/ECME Term of Approval

Issue Date: November 2019
Expiration Date: October 31, 2020
Pharmacodynamic Effects of Vorapaxar in Patients With and Without Diabetes Mellitus

Results of the OPTIMUS-5 Study

Francesco Franchi, MD, Fabiana Rollini, MD, Victor Kairouz, MD, Jose Rivas Rios, MD, Andrea Rivas, MD, Malhar Agarwal, MD, Maryuri Briceno, MD, Mustafa Walli, MD, Ahmed Nawaz, MD, Gabriel Silva, MD, Zubair Shaikh, MD, Naji Maaliki, MD, Latonya Been, AAS, Jason Piraino, DPM, Andres M. Pineda, MD, Siva Suryadevara, MD, Daniel Soffer, MD, Martin M. Zenni, MD, Lisa K. Jennings, PhD, Theodore A. Bass, MD, Dominick J. Angiolillo, MD, PhD

HIGHLIGHTS

- Vorapaxar reduces thrombotic cardiovascular events in patients with atherosclerotic disease, with enhanced effects in those with DM.
- Adjunctive vorapaxar therapy reduces platelet-mediated thrombogenicity without affecting clot kinetics in both patients with and those without DM having prior MI/PAD on dual antiplatelet therapy with aspirin and clopidogrel.
- The pharmacodynamic effects of vorapaxar occur via selective blockade of the PAR-1 on the platelet membrane without apparent interplay with other platelet signaling pathways.
- Aspirin withdrawal, which leaves patients on a background of clopidogrel and vorapaxar, increases markers specific to COX-1-mediated blockade, leading to an increase in platelet-mediated global thrombogenicity, particularly among patients with DM.
Vorapaxar reduces thrombotic cardiovascular events at the expense of increased bleeding. However, the differential pharmacodynamic (PD) effects of vorapaxar according to diabetes mellitus (DM) status are unknown. Moreover, although withdrawal of aspirin has emerged as a bleeding reduction strategy, the PD effects of stopping aspirin in patients treated with vorapaxar also are unknown. In this prospective PD investigation, vorapaxar was associated with reduced platelet-mediated thrombogenicity without affecting clot kinetics irrespective of DM status. However, platelet-mediated thrombogenicity increased after aspirin withdrawal, particularly among patients with DM. (Optimizing anti-Platelet Therapy in diabetes Mellitus-5 Study [OPTIMUS-5]; NCT02548650) (J Am Coll Cardiol Basic Trans Science 2019;4:763–75) © 2019 The Authors. Published by Elsevier on behalf of the American College of Cardiology Foundation. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

Patients with diabetes mellitus (DM) remain at increased risk for recurrent atherothrombotic events despite standard-of-care oral antiplatelet therapy (1,2). This risk is in part due to their hyperreactive platelet phenotype, which can contribute to inadequate response to oral antiplatelet agents, including dual antiplatelet therapy (DAPT) with aspirin and clopidogrel, which is commonly used for secondary prevention of ischemic recurrences (2–4). Importantly, platelets of patients with DM are characterized by up-regulation of platelet signaling pathways that are not inhibited by DAPT, including thrombin-mediated signaling (2). Of note, thrombin is the most potent inducer of platelet activation and plays a key role in thrombus formation (5,6). Hence, modulating the effects of thrombin represents an attractive option to reduce the risk of thrombotic complications, particularly in high-risk patients such as those with DM (7).

Vorapaxar is a novel, orally active, highly selective, competitive, slowly reversible protease-activated receptor (PAR)-1 inhibitor, which exerts potent inhibition of thrombin-mediated platelet aggregation (7–9). In a large-scale clinical trial, vorapaxar (as adjunct to standard-of-care antiplatelet therapy, mostly aspirin and clopidogrel) significantly reduced recurrent thrombotic events in patients with previous atherothrombosis, particularly those with prior myocardial infarction (MI) or peripheral arterial disease (PAD), albeit at the cost of increased bleeding (10–12). Notably, the absolute risk reduction of thrombotic complications associated with the adjunctive use of vorapaxar was greater among DM compared with patients without DM (13). These clinical observations in conjunction with the distinctive platelet phenotype of patients with DM make these patients particularly attractive for treatment with vorapaxar. However, the differential pharmacodynamic (PD) effects of vorapaxar in DM compared with patients without DM are unknown. Moreover, despite the proven efficacy of vorapaxar in reducing thrombotic complications, the increased risk of bleeding complications remains of concern. Withdrawal of aspirin when potent adjunctive antithrombotic therapies are used has been suggested as a strategy to reduce the risk of bleeding (14). Reduction of bleeding complications with aspirin withdrawal has been consistently shown among patients undergoing coronary stenting treated with a P2Y12 receptor inhibitor (mostly clopidogrel) and requiring oral anticoagulant therapy (i.e., blockade of circulating thrombin) (15–18). However, the PD effects associated with a combination of vorapaxar and clopidogrel, without aspirin, is unknown.

**METHODS**

**STUDY DESIGN AND PARTICIPANTS.** The OPTIMUS (Optimizing anti-Platelet Therapy In diabetes Mellitus)-5 study (NCT02548650) was a prospective, parallel-design, open-label investigation aimed at assessing the PD effects of adjunctive use of vorapaxar in patients with a history of MI or PAD, with and without type 2 DM, on treatment with DAPT (aspirin and clopidogrel) and assessing the PD impact associated with discontinuation of aspirin therapy. The study was conducted in patients with a history of MI or PAD in line with the approved indication for use of vorapaxar (19,20). In particular, vorapaxar is approved by the United States Food and Drug Administration and the European Medicines Agency for the reduction of thrombotic cardiovascular events in patients with a history of MI or PAD and is required to be used in addition to standard-of-care antiplatelet therapy with aspirin and/or clopidogrel (20,21). The study was performed at the University of Florida Health-Jacksonville (Jacksonville, Florida). Patients...
with a history of MI or PAD older than 18 years on DAPT with aspirin and clopidogrel as part of their standard of care for at least 2 weeks were screened for study eligibility at the outpatient cardiology clinics of our institution (see Supplemental Material for details on study inclusion and exclusion criteria). The study complied with the Declaration of Helsinki and was approved by the Western Institutional Review Board. All patients gave written informed consent.

Patients on DAPT with aspirin and clopidogrel who met study entry criteria were divided into 2 cohorts according to the presence or absence of type 2 DM. Type 2 DM status was defined according to the World Heart Organization criteria, and patients needed to be on treatment with oral hypoglycemic agents and/or insulin for at least 2 months, without any changes in regimen (22). Vorapaxar (2.5 mg once daily [o.d.]) was added to the standard DAPT regimen of aspirin (81 mg o.d.) plus clopidogrel (75 mg o.d.), also known as triple therapy. Triple therapy was maintained for 30 ± 5 days. Patients then stopped taking aspirin and maintained dual therapy with vorapaxar (2.5 mg o.d.) plus clopidogrel (75 mg o.d.) for 30 ± 5 days. Blood sampling for PD testing was conducted at 3 time points: baseline (while patients were on standard DAPT); after 30 ± 5 days of triple therapy; and 30 ± 5 days after dual therapy. At each time point, blood was collected before the morning dose of clopidogrel and vorapaxar, in order to measure trough levels of platelet inhibition. Laboratory personnel were blinded to treatment assignments. Compliance to randomized treatment was assessed by pill count and patient interview. After completing the study, patients resumed their standard DAPT regimen. A flow diagram of the study design is illustrated in Figure 1.

**BLOOD SAMPLING AND LABORATORY ASSESSMENTS.** Peripheral venous blood samples were drawn through a short venous catheter inserted into a forearm vein and collected in citrate, EDTA, and serum tubes as appropriate for assessments. The first 2 to 4 ml of blood was discarded to avoid spontaneous platelet activation. Blood sampling for PD assessments was performed at 3 time points as indicated in the study design section. Multiple assays were used, including light transmittance aggregometry (LTA); whole blood vasodilator-stimulated phosphoprotein (VASP); TEG 6s (Haemonetics Corp., Braintree, Massachusetts) thrombelastograph coagulation analyzer, which also included the platelet mapping assay using adenosine diphosphate (ADP); and enzyme-linked immunosorbent assay–based assessment of serum thromboxane.
### Baseline Characteristics of the PD Population

|                        | DM (n = 30) | Non-DM (n = 34) | p Value |
|------------------------|-------------|-----------------|---------|
| **Age, yrs**           | 61 ± 8      | 56 ± 9          | 0.015   |
| **Male**               | 23 (77)     | 23 (67)         | 0.579   |
| **BMI, kg/m²**         | 31 ± 5      | 32 ± 8          | 0.757   |
| **Race**               |             |                 |         |
| White                  | 15 (50)     | 22 (65)         |         |
| Black                  | 14 (47)     | 11 (32)         |         |
| Other                  | 1 (3)       | 1 (3)           |         |

Enrollment criteria:
- Prior MI: 13 (43) vs 28 (82), p = 0.002
- PAD: 17 (57) vs 6 (18), p = 0.002
- CKD: 2 (7) vs 2 (6), p = 1
- Hypertension: 29 (97) vs 25 (74), p = 0.015
- Dyslipidemia: 27 (90) vs 22 (65), p = 0.020
- Active smoking: 10 (33) vs 14 (41), p = 0.408
- CAD: 22 (73) vs 20 (67), p = 0.593
- Prior PCI: 20 (67) vs 25 (73), p = 0.534
- Prior CABG: 7 (23) vs 4 (12), p = 0.322
- Creatinine, mg/dl: 1.1 ± 0.3 vs 0.9 ± 0.2, p = 0.073
- CrCl, ml/min: 102 ± 40 vs 117 ± 58, p = 0.232
- Platelet count, 10^9/μl: 249 ± 62 vs 227 ± 59, p = 0.160
- Hematocrit, %: 39 ± 4 vs 41 ± 4, p = 0.182
- Hemoglobin, g/dl: 12.9 ± 1.3 vs 13.5 ± 1.6, p = 0.145

Medications:
- Insulin therapy: 16 (53) vs 0 (0), p < 0.001
- OAD: 23 (77) vs 0 (0), p < 0.001
- Beta-blockers: 27 (90) vs 30 (88), p = 1
- ACE inhibitor/ARB: 25 (83) vs 21 (62), p = 0.093
- Statins: 30 (100) vs 32 (94), p = 0.494

Values are mean ± SD or n (%). *2 patients categorized as MI also had PAD (see Supplemental Material for definition of study entry criteria).

ACE = angiotensin-converting enzyme; ARB = angiotensin receptor blocker; BMI = body mass index; CABG = coronary artery bypass graft; CAD = coronary artery disease; CKD = chronic kidney disease; CrCl = creatinine clearance; DM = diabetes mellitus; MI = myocardial infarction; OAD = oral antidiabetic drug; PAD = peripheral arterial disease; PCI = percutaneous coronary intervention; PD = pharmacodynamic.

B₂ (TXB₂) (23–26). A detailed description of the assays is provided in the Supplemental Material. Assessments were performed and described with the following objectives: 1) to define the PD effect of vorapaxar on thrombin-mediated effects on platelets and systemically; to this extent, LTA following thrombin receptor activating peptide (TRAP) (15 μM) stimuli and markers of clot kinetics using the TEG 6s system were utilized, respectively; 2) to define the PD effect of vorapaxar on platelet-mediated thrombogenicity; to this extent, LTA following stimuli with combination of 2 μg/ml collagen-related peptide + 5 μM ADP + 15 μM TRAP (CAT) was used; and 3) to define the PD effect of vorapaxar on P2Y₁₂ inhibition induced by clopidogrel and the impact of aspirin withdrawal; to this extent LTA following stimuli with ADP (20 μM) and VASP as well as markers sensitive to cyclooxygenase (COX)-1 blockade, including LTA following arachidonic acid (1 mM) and collagen (3 μg/ml) stimuli, and measurement of serum TXB₂ levels, respectively, were assessed. LTA results are reported as maximum platelet aggregation (MPA%), VASP results as platelet reactivity index, and serum TXB₂ levels in ng/ml.

**STUDY ENDPOINTS AND SAMPLE SIZE CALCULATION.**

The primary endpoint of our study was the comparison of CAT-induced MPA measured by LTA between triple (vorapaxar plus DAPT) and dual (vorapaxar plus clopidogrel) therapy. The rationale for choosing CAT-induced MPA for the primary endpoint was that this combination of agonist is more reflective of global thrombogenicity as it stimulates multiple platelet signaling pathways. We hypothesized that dual therapy would be noninferior to triple therapy after 30 ± 5 days of treatment in both patients with and without DM. Under the null hypothesis that the mean CAT-induced MPA between dual and triple therapy is not equal to 0 and a common standard deviation of 13%, a sample size of 28 patients per group with a valid primary endpoint time point allowed for the 95% confidence interval (CI) to stay within ±10% with 80% power and 2-sided \( z = 0.05 \). Considering the 2 groups (patients with and without DM), a total of 56 patients with valid primary endpoint data needed to be included. Assuming up to 40% rate of invalid results due to hemolysis or dropout, we estimated that up to 79 patients would need to be enrolled. Noninferiority was assessed using a 95% CI of the difference in mean MPA between the 2 arms. As there were no preliminary data in this setting, the 10% noninferiority margin was arbitrarily defined. Mean values of platelet aggregation and variability were estimated based on previous data of vorapaxar (26). Our approach for the statistical assumption is in agreement with recommendations for pilot investigations (27).

Other exploratory objectives included comparisons between patients with and without DM of all PD parameters measured by multiple assays, and comparisons between levels of platelet inhibition achieved by DAPT (baseline therapy) versus levels achieved by adding vorapaxar. The effects of additive inhibition of the thrombin-mediated platelet activation pathway, with or without aspirin therapy, on serum thromboxane levels also were evaluated.

**STATISTICAL ANALYSIS.**

Categorical variables are expressed as frequencies and proportions. Continuous variables were analyzed for normal distribution with the Kolmogorov-Smirnov test and are expressed as mean ± SD. Comparisons between categorical variables were performed using the 2-tailed Fisher exact test or the Pearson chi-square test. The
Student’s t-test was used to compare continuous variables. A full-factorial repeated measure analysis of variance method for dependent variables with a general linear model was used to evaluate intragroup comparisons between time points (3). Least square mean differences in MPA between groups and the corresponding 2-sided 95% CI for the difference were obtained to assess noninferiority based on the linear model. The Student’s t-test was used to compare continuous variables between groups. Platelet reactivity results are reported as mean ± SD for the detailed analyses. p Values are used to report superiority testing, and CIs are used to determine noninferiority. A 2-tailed p < 0.05 is considered to indicate a statistically significant difference for all the analyses performed. Given the exploratory nature of superiority intragroup comparisons, adjustment for multiple comparisons was not performed. Statistical analysis was performed using SPSS version 25.0 software (SPSS Inc., Chicago, Illinois).

The safety population was composed of all patients exposed to at least 1 dose of study medication (any time from enrollment until completion of the study). Any adverse event during the study period was recorded. The PD population included all patients with PD data and without a major protocol deviation thought to affect the PD effects of vorapaxar, aspirin, and clopidogrel. The PD population was used for analysis of all primary and secondary PD variables.

RESULTS

PATIENT POPULATION. Between March 25, 2016 and October 22, 2018, 132 patients were screened. A total of 71 patients on maintenance therapy agreed to participate in the study; 5 patients were not eligible for randomization due to the presence of an exclusion criteria. A total of 66 patients (30 with DM, 36 without DM) were exposed to at least 1 dose of study medication, representing the safety population. Two patients were not compliant with medications and therefore were excluded from the PD analysis. Thus, a total of 64 patients (30 with DM, 34 without DM) represented the PD population of the study. Of these patients, 56 (28 with DM, 28 without DM) had valid primary endpoint data. Prior MI and PAD were the enrollment criteria for 41 (64%) and 23 (36%) patients, respectively. Baseline characteristics of the study population are summarized in Table 1. Patients with DM were older and had higher rates of PAD, hypertension, and hyperlipidemia. No ischemic or Bleeding Academic Research Consortium type 2 to 5 bleeding events were observed. One DM patient had a Bleeding Academic Research Consortium type 1 bleeding
(hematuria) while on triple therapy that led to study drug discontinuation, and 11 patients (7 with DM, 4 without DM) had nonbleeding adverse events (see Supplemental Material for details).

**PD FINDINGS. Effects of vorapaxar on thrombin-mediated effects.** Adjunctive treatment with vorapaxar was associated with complete blockade of TRAP-induced platelet aggregation ($p < 0.001$). Complete suppression of TRAP-induced platelet aggregation persisted after discontinuation of aspirin therapy ($p < 0.001$). Overall, such effects were consistent irrespective of DM status (Figure 2A). On the contrary, vorapaxar did not affect markers of clot kinetics, including speed of thrombin generation, which remained unvaried even after discontinuation of aspirin therapy (Figure 2B). Findings were consistent irrespective of DM status (Supplemental Figures 1 to 4). These observations are supportive of the platelet-specific, and not systemic, effects of vorapaxar on modulating thrombin-mediated effects.

**Effects of vorapaxar on global thrombogenicity.** Adding vorapaxar to DAPT significantly reduced CAT-induced aggregation in both patients with DM (mean difference $\approx 15$; 95% CI: 7 to 23; $p < 0.001$) and patients without DM (mean difference $\approx 20$; 95% CI: 15 to 25; $p < 0.001$). However, stopping aspirin was associated with an increase in CAT-induced aggregation in both patients with DM (mean difference $\approx 12$; 95% CI: 3 to 21; $p = 0.010$) and patients without DM (mean difference $\approx 10$; 95% CI: 3 to 16; $p = 0.003$), thus not meeting the primary endpoint of non-inferiority (Figure 3). After aspirin withdrawal, CAT-induced aggregation was significantly lower compared with baseline in patients without DM (mean difference $\approx 15$; 95% CI: 7 to 23; $p < 0.001$) but not in DM (mean difference $\approx 3$; 95% CI: -6 to 11; $p = 0.542$) patients. Overall, the magnitude of increase in CAT-induced aggregation after aspirin withdrawal in the presence of vorapaxar and clopidogrel therapy was higher in DM patients compared with patients without DM ($p = 0.036$) (Figure 3).

**Effects of vorapaxar on modulating P2Y$_{12}$ inhibition induced by clopidogrel in the presence and absence of aspirin.** Adding vorapaxar to DAPT did not affect markers assessing P2Y$_{12}$ inhibition using all assays, including LTA and VASP. After aspirin withdrawal, there were no significant differences in levels of these
markers among patients without DM, but these levels modestly increased in patients with DM (Figure 4, Supplemental Figure 5). Aspirin withdrawal was associated with a marked increase in makers sensitive to COX-1 blockade, including arachidonic acid– and collagen-induced aggregation as well as serum TXB₂ levels, in both patients with and without DM (Figure 5, Supplemental Figure 6).

**DISCUSSION**

Vorapaxar is a PAR-1 inhibitor clinically approved for reduction of thrombotic cardiovascular events in patients with a history of MI or PAD treated with standard-of-care antiplatelet therapy with aspirin and/or clopidogrel (19–21). The OPTIMUS-5 study was conducted to provide insights on the PD effects of vorapaxar in this setting, particularly exploring profiles among patients with and without DM treated with DAPT (aspirin and clopidogrel), which to date have not been explored. Moreover, in light of the emerging interest in an aspirin-free approach as a strategy to reduce the risk of bleeding complications in patients treated with more potent antithrombotic therapies, this investigation also explored the PD effects associated with aspirin withdrawal in the presence of vorapaxar and clopidogrel therapy. The results of OPTIMUS-5 can be summarized as follows: 1) adjunctive treatment with vorapaxar reduces platelet-mediated thrombogenicity without affecting clot kinetics in both patients with and without DM treated with DAPT (aspirin and clopidogrel); 2) reduction of platelet-mediated thrombogenicity induced by vorapaxar occurs selectively via PAR-1 blockade without apparent interplay with other platelet signaling pathways (e.g., P2Y₁₂ or thromboxane); and 3) platelet-mediated thrombogenicity is increased after aspirin withdrawal, particularly among patients with DM.

The TRA 2P-TIMI 50 (Thrombin Receptor Antagonist in Secondary Prevention of Atherothrombotic Ischemic Events-Thrombolysis In Myocardial Infarction 50) trial, conducted in patients with previous atherothrombosis, demonstrated that, when added to standard-of-care treatment, including antiplatelet therapy with aspirin and clopidogrel, vorapaxar significantly reduces recurrent thrombotic events (10). This benefit was limited to patients with a history of prior MI or PAD but not to patients with a prior cerebrovascular event, in whom vorapaxar was associated with increased harm (i.e., increased rates of intracranial hemorrhage) (10–12). Notably, in the cohort of patients with DM and prior MI included in the TRA 2 P trial, vorapaxar reduced the primary composite endpoint at 3 years by 27% (p = 0.002) and led to a greater absolute risk reduction (absolute risk difference: −3.50%) compared with those without DM (absolute risk difference: −1.36%), with a number needed to treat of 29 (13). The benefit of vorapaxar...
also was consistent in patients with PAD, although no specific subgroup data according to DM status were available (12,13). These observations make patients with DM an attractive population for treatment with vorapaxar. The reason for the enhanced benefit of vorapaxar among patients with DM has been hypothesized to be attributed to an up-regulated status of thrombin-mediated platelet activation, which makes these patients more susceptible to the antithrombotic effects of the drug (13). However, in our investigation we found that vorapaxar completely abolished TRAP-induced aggregation irrespective of DM status. In addition, the effects on global thrombogenicity were similar in patients with and without DM. Thus, it is more likely that the clinical observations from TRA 2P-TIMI 50 can be attributed to the greater baseline risk of patients with DM, which allows for a greater magnitude of treatment effects with vorapaxar. This observation is consistent with other secondary prevention studies in patients with DM using potent antiplatelet therapies (28).

The increased risk of bleeding complications with vorapaxar remains a clinical concern. PD investigations have suggested that in the presence of potent P2Y$_{12}$ blockade, aspirin provides limited adjunctive antithrombotic effects (14,29,30). Although the GLOBAL LEADERS trial, which tested a strategy of ticagrelor monotherapy after only 1 month of DAPT versus standard DAPT in a large all-comers population undergoing percutaneous coronary intervention, failed to meet its primary endpoint for superior efficacy of aspirin withdrawal, such an experimental strategy was not associated with any safety concerns (31). Several ongoing investigations are evaluating the safety and efficacy of dropping aspirin in the presence of potent P2Y$_{12}$ receptor blockade in patients undergoing percutaneous coronary intervention (14,32). However, a number of other studies, mostly conducted in patients with atrial fibrillation undergoing coronary stenting and requiring treatment with both oral anticoagulant (OAC) and antiplatelet therapy, have shown that stopping aspirin and maintaining treatment with OAC and a P2Y$_{12}$ inhibitor (mostly clopidogrel) significantly reduced bleeding complications without any apparent trade-off in ischemic events (15-18,33). Accordingly, current recommendations are to minimize the duration of DAPT in patients also taking OAC (34,35). OPTIMUS-5 also explored the effects of dropping aspirin in the setting of a platelet-specific, rather than systemic, modulation of thrombin-mediated effects. We showed that stopping aspirin was associated with an increase in markers specific to COX-1-mediated blockade leading to an increase in platelet-mediated global thrombogenicity. Importantly, the magnitude of increase was

---

**FIGURE 5** Markers Sensitive to Cyclooxygenase-1 Blockade

(A) Arachidonic acid (AA)-induced maximal platelet aggregation (MPA%) measured by light transmittance aggregometry (LTA). The p values represent comparisons between the 2 groups at each time point. Error bars indicate SD. Intratreatment comparisons for non-DM: Baseline versus DAPT + vorapaxar: mean difference $\pm$ 2; 95% CI: $\pm$ 10 to 15; $p = 0.734$. Baseline versus clopidogrel + vorapaxar: mean difference $\pm$ 42; 95% CI: 26 to 58; $p < 0.001$. DAPT + vorapaxar versus clopidogrel + vorapaxar: mean difference $\pm$ 44; 95% CI: 28 to 60; $p < 0.001$. Intratreatment comparisons for DM: Baseline versus DAPT + vorapaxar: mean difference $\pm$ 2; 95% CI: $\pm$ 2 to 21; $p = 0.108$. Baseline versus clopidogrel + vorapaxar: mean difference $\pm$ 40; 95% CI: 25 to 54; $p < 0.001$. DAPT + vorapaxar versus clopidogrel + vorapaxar: mean difference $\pm$ 49; 95% CI: 36 to 63; $p < 0.001$. (B) Serum thromboxane B$_2$ (TXB$_2$). The p values represent comparisons between the 2 groups at each time point. Error bars indicate SD. Intratreatment comparisons for non-DM: Baseline versus DAPT + vorapaxar: $p = 0.545$. Baseline versus clopidogrel + vorapaxar: $p = 0.001$. DAPT + vorapaxar versus clopidogrel + vorapaxar: $p < 0.001$. DAPT + vorapaxar versus clopidogrel + vorapaxar: $p < 0.001$. Abbreviations as in Figure 1.
higher in patients with DM compared with patients without DM, and patients with DM achieved levels of global thrombogenicity while on vorapaxar and clopidogrel that were similar to those while on aspirin and clopidogrel.

Our finding that markers specifically assessing COX-1 blockade increase with aspirin withdrawal provides support that alternative antithrombotic treatment regimens cannot replace the selective effects of aspirin on platelet COX-1 blockade. The increase in platelet-mediated global thrombogenicity could also be attributed to the loss of synergism that is known to occur when aspirin and clopidogrel are concomitantly used (36). Such synergism may be less relevant in the presence of more potent P2Y12 blockade (29,30). It may be hypothesized that the enhanced increase in platelet-mediated thrombogenicity among patients with DM may be attributed to the fact that these patients are more susceptible to such loss of synergism as reflected by the increase in markers of P2Y12 signaling. Overall, these findings should caution against strategies of aspirin withdrawal in the absence of effective alternative antithrombotic treatment (14). Indeed, our PD observations question any potential clinical advantage of dual therapy with vorapaxar and clopidogrel compared with standard-of-care DAPT with aspirin and clopidogrel, especially in patients with DM. The use of OAC, which provides a block of systemic levels of thrombin, may be more effective in this regard, as modulation of circulating thrombin can also indirectly affect platelet reactivity (37). Thus, this approach would result in more wide-ranging antithrombotic effects compared with vorapaxar, which only selectively inhibits the effects of thrombin on platelets via selective PAR-1 blockade (7). The benefits of the antithrombotic efficacy of an OAC in addition to a single antiplatelet agent (e.g., aspirin) was recently supported by a large-scale secondary prevention study conducted in patients with CAD and PAD, which showed a significant reduction in ischemic recurrences and reduced cardiovascular mortality associated with a very low dosing regimen of rivaroxaban in adjunct to aspirin (38).

Despite the known interplay between thrombin and P2Y12-mediated signaling, adjunctive treatment with vorapaxar does not interfere with markers of P2Y12 receptor blockade (39). This is consistent with other studies modulating systemic levels of thrombin with oral anticoagulant therapies (40-42). These findings may be attributed to the fact that these patients already are on P2Y12-inhibiting therapy, which may not allow unraveling such interplay as previously observed in in vitro investigations and in platelets from nonmedicated patients (43-45).

**STUDY LIMITATIONS.** Our study was not designed to assess clinical outcomes. Moreover, our study was conducted in patients treated with aspirin and clopidogrel, and it may be argued that many patients with a history of MI are currently treated with more potent P2Y12 inhibitors (prasugrel or ticagrelor). However, in the TRA 2P study, clopidogrel was the P2Y12 inhibitor utilized in 99.3% of patients, and the approved indication for vorapaxar was based on this background antiplatelet therapy. Accordingly, our PD study was conducted to mimic how vorapaxar was approved for clinical use based on the TRA 2P trial (10,20,21). Moreover, clopidogrel is still the most frequently used P2Y12 inhibitor and is the only agent of this class approved for treatment of PAD (46). If similar findings would have been observed in patients on DAPT with aspirin and prasugrel or ticagrelor is unknown. The PD effects associated with the addition of vorapaxar to patients with a history of MI, treated with potent P2Y12 inhibitors (prasugrel or ticagrelor) and the effects associated with aspirin withdrawal in these patients are currently under investigation (NCT02545933).

**CONCLUSIONS**

Adjunctive treatment with vorapaxar reduces platelet-mediated thrombogenicity without affecting clot kinetics in both patients with DM and patients without DM, while on DAPT. However, platelet-mediated thrombogenicity is increased after aspirin withdrawal, particularly among patients with DM. These PD observations do not support any potential clinical advantage of dual therapy with vorapaxar and clopidogrel compared with standard-of-care DAPT with aspirin and clopidogrel, especially in patients with DM. The PD findings of this study suggest that the enhanced clinical benefit among patients with DM associated with adjunctive treatment with vorapaxar in addition to standard-of-care oral antiplatelet therapy, including aspirin and clopidogrel, is more likely due to the greater baseline risk profile of these patients, which allows for a greater magnitude of treatment effects, rather than a differential effect on platelets from patients with DM compared to patients without DM.

**ADDRESS FOR CORRESPONDENCE:** Dr. Dominick J. Angiolillo, University of Florida College of Medicine-Jacksonville, 655 West 8th Street, Jacksonville, Florida 32209. E-mail: dominick.angiolillo@jax.ufl.edu.
Vorapaxar and Diabetes

COMPETENCY IN MEDICAL KNOWLEDGE:
Vorapaxar is a PAR-1 inhibitor clinically approved for the reduction of thrombotic cardiovascular events in patients with a history of MI or PAD treated with standard-of-care antiplatelet therapy with aspirin and/or clopidogrel. This study provides evidence that adjunctive treatment with vorapaxar in addition to DAPT reduces platelet-mediated thrombogenicity without affecting clot kinetics in both patients and patients without DM. Notably, this reduction occurs selectively via PAR-1 blockade without apparent interplay with other platelet signaling pathways, and platelet-mediated thrombogenicity is increased after aspirin withdrawal, particularly among patients with DM. This has clinical implications as it demonstrates that the enhanced benefit of vorapaxar shown in clinical trials among patients with DM is likely just attributed to the greater baseline risk of patients with DM, thus under-scoring the need for more aggressive treatment of these patients. In addition, the study shows that alternative antithrombotic treatment regimens cannot replace the selective effects of aspirin on platelet COX-1 blockade and cautions against strategies of aspirin withdrawal in the absence of effective alternative antithrombotic treatment.

TRANSLATIONAL OUTLOOK: Our study was conducted in patients treated with aspirin and clopidogrel, and it may be argued that many patients with a history of MI are currently treated with the more potent P2Y12 inhibitors prasugrel and ticagrelor. The PD effects associated with the addition of vorapaxar to treatment with prasugrel or ticagrelor and the effects associated with aspirin withdrawal in these patients need to be investigated in specifically designed studies. Use of direct oral anticoagulants, which inhibit systemic thrombin, may be more effective in reducing platelet-mediated global thrombogenicity, as modulation of circulating thrombin (rather than just its selective effect on the platelet PAR-1 receptor) can also indirectly affect platelet reactivity and result in more wide-ranging antithrombotic effects. The association between PD findings and clinical outcomes in patients with vascular disease manifestations treated with strategies that modulate the effects of thrombin warrants further investigation.

REFERENCES
1. Donahoe SM, Stewart GC, McCabe CH, et al. Diabetes and mortality following acute coronary syndromes. JAMA 2007;298:765-75.
2. Ferreiro JL, Angiolillo DJ. Diabetes and antiplatelet therapy in acute coronary syndrome. Circulation 2011;123:798-813.
3. Angiolillo DJ, Jakubowski JA, Ferreiro JL, et al. Impaired responsiveness to the platelet P2Y12 receptor antagonist clopidogrel in patients with type 2 diabetes and coronary artery disease. J Am Coll Cardiol 2014;64:1005-14.
4. Angiolillo DJ, Fernandez-Ortiz A, Bernardo E, et al. Platelet function profiles in patients with type 2 diabetes and coronary artery disease on combined aspirin and clopidogrel treatment. Diabetes 2005;54:2430-5.
5. Davi G, Patrono C. Platelet activation and atherothrombosis. N Engl J Med 2007;357:2482-94.
6. Angiolillo DJ, Ueno M, Goto S. Basic principles of platelet biology and clinical implications. Circ J 2010;74:597-607.
7. Angiolillo DJ, Capodanno D, Goto S. Platelet thrombin receptor antagonism and atherothrombosis. Eur Heart J 2010;31:17-28.
8. Franchi F, Angiolillo DJ. Novel antiplatelet agents in acute coronary syndrome. Nat Rev Cardiol 2015;12:30-47.
9. Franchi F, Rollini F, Park Y, Angiolillo DJ. Platelet thrombin receptor antagonism with vorapaxar: pharmacology and clinical trial development. Future Cardiol 2015;11:547-64.
10. Morrow DA, Braunwald E, Bonaca MP, et al. Vorapaxar in the secondary prevention of atherothrombotic events. N Engl J Med 2012;366:1404-13.
11. Scirica BM, Bonaca MP, Braunwald E, et al. Vorapaxar for secondary prevention of thrombotic events for patients with previous myocardial infarction: a prespecified subgroup analysis of the TRA 2P-TIMI 50 trial. Lancet 2012;380:1317-24.
12. Bonaca MP, Scirica BM, Creager MA, et al. Vorapaxor in patients with peripheral artery disease: results from TRA2P-TIMI 50. Circulation 2013;127:1522-9.
13. Cavender MA, Scirica BM, Bonaca MP, et al. Vorapaxor in patients with diabetes mellitus and previous myocardial infarction: findings from the thrombin receptor antagonist in secondary prevention of atherothrombotic ischemic events-TIMI 50 trial. Circulation 2015;131:1047-53.
14. Capodanno D, Mehran R, Valgimigli M, et al. Aspirin-free strategies in cardiovascular disease and cardioembolic stroke prevention. Nat Rev Cardiol 2018;15:480-96.
15. Dewilde WJ, O’Riabns T, Verheugt FW, et al. 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;381:1107-15.
16. Gibson CM, Mehran R, Bode C, et al. Prevention of bleeding in patients with atrial fibrillation undergoing PCI. N Engl J Med 2016;375:2423-34.
17. Cannon CP, Bhatt DL, Oldgren J, et al. Dual antithrombotic therapy with dabigatran after PCI in atrial fibrillation. N Engl J Med 2017;377:1513-24.
18. Lopes RD, Heizger G, Aronson R, et al. Antithrombotic therapy after acute coronary syndrome or PCI in atrial fibrillation. N Engl J Med 2019;380:1509-24.
19. Magnani G, Bonaca MP, Braunwald E, et al. Efficacy and safety of vorapaxor as approved for clinical use in the United States. J Am Heart Assoc 2015;4:e001505.
20. U.S. Food and Drug Administration (FDA). Vorapaxor prescribing information. Available at: http://www.fda.gov/drugs/informationondrugs/ucm423935.htm. Accessed June 1, 2019.
21. European Medicines Agency (EMA). Vorapaxor prescribing information. Available at: http://www.ema.europa.eu/ema/index.jsp?curl=pages/medicines/human/medicines/002814/
2016;27:642
Gurbel PA, Bliden KP, Tantry US, et al. First Study. Circulation 2016;134:780
and coronary artery disease: the OPTIMUS (Opti-
macodynamic comparison of prasugrel versus
Franchi F, Rollini F, Aggarwal N, et al. Phar-
expression in non-ST-elevation acute coronary
26. Storey RF, Kotha J, Smyth SS, et al. Effects of
dosing regimens in type 2 diabetes mellitus pa-
capodanno D, Patel A, Dharmashankar K, et al. Pharm-
and analysis of pilot studies: recommendations for
28. Armstrong PC, Leadbeater PD, Chan MV, et al. In the presence of strong P2Y12 receptor
blockade, aspirin provides little additional inhibition
of platelet aggregation. J Thromb Haemost 2011;9:562–61.
29. Traby L, Kollars M, Kaidar A, Eichinger S, Woltz M, Kyrle PA. Effects of P2Y12 receptor in-
hibition with or without aspirin on hemostatic system activation: a randomized trial in healthy
subjects. J Thromb Haemost 2016;14:273–81.
30. Vranckx P, Valgimigli M, Jüni P, et al. Ticag-
relor plus aspirin for 1 month, followed by ticag-
relor monotherapy for 23 months vs aspirin plus
clopидогрел or ticagrelor for 12 months, followed by aspirin monotherapy for 12 months after im-
plantation of a drug-eluting stent: a multicentre, open-label, randomised superiority trial. Lancet 2018;392:940–9.
31. Baber U, Dangs G, Cohen DJ, et al. Ticagrelor
with aspirin or alone in high-risk patients after coronary intervention: rationale and design of the
TWILIGHT study. Am Heart J 2016;182:125–34.
32. Lopes RD, Hong H, Harshkamp RE, et al. Safety and efficacy of antithrombotic strategies in
patients with atrial fibrillation undergoing percu-
taneous coronary intervention: a network meta-
analysis of randomized controlled trials. JAMA Cardiol 2019 June 19 [E-pub ahead of print].
33. Angiolillo DJ, Goodman SG, Bhatt DL, et al. Antithrombotic therapy in patients with atrial
fibrillation treated with oral anticoagulation un-
dering percutaneous coronary intervention. Circulation 2018;138:527–36.
34. Capodanno D, Huber K, Mehran R, et al. Management of antithrombotic therapy in atrial
fibrillation patients undergoing PCI: current status and future directions. J Am Coll Cardiol 2019;74:
83–99.
35. Cadroy Y, Bossavy JP, Thalames C, Sagnard L,
Sakariassen K, Boneu B. Early potent antith-
rombotic effect with combined aspirin and a
loading dose of clopidogrel on experimental
arterial thrombogenesis in humans. Circulation 2000;101:2823–8.
36. Berissoff JI, Sporkin HM, ten Cate H. The he-
mostatic system as a modulator of atherosclerosis.
N Engl J Med 2011;364:1746–60.
37. Eikelboom JW, Connolly SJ, Bosch J, et al. Rivaroxaban with or without aspirin in stable car-
diovascular disease. N Engl J Med 2017;377:
1319–30.
38. Nylander S, Mattsson C, Ramström S, Lindahl TL. Synergistic action between inhibition of
P2Y12/P2Y1 and P2Y12/thrombin in ADP- and
thrombin-induced human platelet activation. Br J Pharmaco 2004;142:1325–31.
39. Franchi F, Rollini F, Cho JR, et al. Effects of
dabiaglutan on the cellular and protein phase of
coaulation in patients with coronary artery
disease on dual antiplatelet therapy with aspirin and clo-
pidogrel: results of the EDOX-APT study. Thromb Haemost 2019. Aug 30 [Epub ahead of print].
40. Furugohi T, Isobe K, Honda Y, et al. DU-176b,
a potent and orally active factor Xa inhibitor: in
vitro and in vivo pharmacological profiles. J Thromb Haemost 2006;4:1542–9.
41. Honda Y, Kamisato C, Morishima Y. Edox-
aban, a direct factor Xa inhibitor, suppresses
tissue-factor induced human platelet aggregation
and clot-bound factor Xa in vitro: comparison
with an antithrombin-dependent factor Xa in-
hibitor, fondaparinux. Thromb Res 2016;141:
17–21.
42. Wong PC, Jiing X, Apachea, a direct factor Xa
inhibitor, inhibits tissue-factor induced human
platelet aggregation in vitro: comparison with
direct inhibitors of factor Vila, Xia and thrombin.
Thromb Haemost 2010;104:302–10.
43. Jones WS, Patel MR. Antithrombotic therapy
in peripheral artery disease: generating and
translating evidence into practice. J Am Coll Car-
diol 2018;71:352–62.

**KEY WORDS** dual antiplatelet therapy,
pharmacodynamics, platelets, thrombin, vorapaxar

**APPENDIX** For an expanded Methods section
as well as supplemental figures and references,
please see the online version of this paper.