

Relationship of Platelet Reactivity With Bleeding Outcomes During Long-Term Treatment With Dual Antiplatelet Therapy for Medically Managed Patients With Non-ST-Segment Elevation Acute Coronary Syndromes

Jan H. Cornel, MD, PhD; E. Magnus Ohman, MD; Benjamin Neely, MS; Joseph A. Jakubowski, PhD; Deepak L. Bhatt, MD, MPH; Harvey D. White, MB, ChB, DSc; Diego Ardissino, MD; Keith A.A. Fox, MB, ChB; Dorairaj Prabhakaran, MD, DM, MSc; Paul W. Armstrong, MD; David Erlinge, MD, PhD; Udaya S. Tantry, PhD; Paul A. Gurbel, MD; Matthew T. Roe, MD, MHS

Background—The relationship between "on-treatment" low platelet reactivity and longitudinal risks of major bleeding dual antiplatelet therapy following acute coronary syndromes remains uncertain, especially for patients who do not undergo percutaneous coronary intervention.

Methods and Results—We analyzed 2428 medically managed acute coronary syndromes patients from the Targeted Platelet Inhibition to Clarify the Optimal Strategy to Medically Manage Acute Coronary Syndromes (TRILOGY ACS) trial who had serial platelet reactivity measurements (P2Y₁₂ reaction units; PRUs) and were randomized to aspirin+prasugrel versus aspirin+clopidogrel for up to 30 months. Contal's method was used to determine whether a cut point for steady-state PRU values could distinguish high versus low bleeding risk using 2-level composites: Global Use of Strategies to Open Occluded Coronary Arteries (GUSTO) severe/life-threatening or moderate bleeding unrelated to coronary artery bypass grafting (CABG) and non-CABG Thrombolysis In Myocardial Infarction (TIMI) major or minor bleeding. Exploratory analyses used 3-level composites that incorporated mild and minimal GUSTO and TIMI events. Continuous measures of PRUs (per 10-unit decrease) were not independently associated with the 2-level GUSTO (adjusted hazard ratio [HR], 1.01; 95% CI, 0.96—1.06) or TIMI composites (1.02; 0.98—1.07). Furthermore, no PRU cut point could significantly distinguish bleeding risk using the 2-level composites. However, the PRU cut point of 75 differentiated bleeding risk with the 3-level composites of GUSTO (26.5% vs 12.6%; adjusted HR, 2.28; 95% CI, 1.77—2.94; P<0.001) and TIMI bleeding events (25.9% vs 12.2%; adjusted HR, 2.30; 95% CI, 1.78—2.97; P<0.001).

Conclusions—Among medically managed non-ST-segment elevation acute coronary syndromes patients receiving prolonged dual antiplatelet therapy, PRU values were not significantly associated with the long-term risk of major bleeding events, suggesting that low on-treatment platelet reactivity does not independently predict serious bleeding risk.

Clinical Trial Registration—URL: http://www.clinicaltrials.gov. Unique identifier: NCT00699998. (*J Am Heart Assoc.* 2016;5: e003977 doi: 10.1161/JAHA.116.003977)

Key Words: DAPT • hemorrhage • platelet

C linical practice guidelines recommend dual antiplatelet therapy (DAPT) with aspirin+a P2Y₁₂ inhibitor for at least 12 months for patients with acute coronary syndromes (ACS),

given the consistent benefits of DAPT demonstrated in large randomized trials.^{1,2} Although P2Y₁₂ inhibitors have been shown to reduce ischemic events, there has been a consistent

From the Medisch Centrum Alkmaar, Alkmaar, The Netherlands (J.H.C.); Duke Clinical Research Institute, Durham, NC (E.M.O., B.N., M.T.R.); Division of Cardiology, Department of Medicine, Duke University Medical Center, Durham, NC (E.M.O., M.T.R.); Eli Lilly and Company, Indianapolis, IN (J.A.J.); Brigham and Women's Hospital Heart & Vascular Center and Harvard Medical School, Boston, MA (D.L.B.); Green Lane Cardiovascular Service, Auckland City Hospital, Auckland, New Zealand (H.D.W.); Division of Cardiology, Azienda Ospedaliero—Universitaria di Parma, Italy (D.A.); Centre for Cardiovascular Science, University of Edinburgh, Scotland, UK (K.A.A.F.); Centre for Chronic Disease Control and Public Health Foundation of India, New Delhi, India (D.P.); Canadian VIGOUR Centre and Division of Cardiology, University of Alberta, Edmonton, Alberta, Canada (P.W.A.); Department of Cardiology, Lund University, Lund, Sweden (D.E.); Sinai Center for Thrombosis Research, Baltimore, MD (U.S.T., P.A.G.).

Accompanying Figure S1 and Tables S1 through S4 are available at http://jaha.ahajournals.org/content/5/11/e003977/DC1/embed/inline-supplementary-material-1.pdf

Correspondence to: Matthew T. Roe, MD, MHS, 2400 Pratt St, Room 7035, Durham, NC 27705. E-mail: matthew.roe@duke.edu Received May 31, 2016; accepted September 30, 2016.

© 2016 The Authors. Published on behalf of the American Heart Association, Inc., by Wiley Blackwell. This is an open access article under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs License, which permits use and distribution in any medium, provided the original work is properly cited, the use is non-commercial and no modifications or adaptations are made.

signal of increased bleeding with DAPT treatment compared with aspirin monotherapy, and with DAPT regimens that include more-potent P2Y12 inhibitors (prasugrel and ticagrelor) compared with clopidogrel.^{3–5} These latter observations may indicate that enhanced platelet inhibition is associated with increased bleeding risk.

Given the consistent association of bleeding events with an increased risk of subsequent mortality and other ischemic outcomes, the focus of DAPT treatment is shifting toward finding the optimal risk/benefit balance for patients with ACS to mitigate the risk of major bleeding while maintaining a significant reduction of ischemic events.⁶ In this regard, past studies have suggested that patients undergoing percutaneous coronary intervention (PCI) who have a robust response to a P2Y₁₂ inhibitor (termed low on-treatment platelet reactivity [LPR] to ADP) have a higher risk of long-term bleeding events following the procedure.^{7,8} Based on the results of these observational studies, a therapeutic window concept for P2Y₁₂ receptor reactivity, in which a cut-off value for high ontreatment platelet reactivity and LPR to ADP associated with post-PCI ischemic and bleeding event risk, has been recently proposed. However, the relationship of platelet reactivity measurements and LPR with long-term bleeding risk in patients with ACS treated with DAPT and managed without revascularization has not been prospectively evaluated.

We analyzed data from the Platelet Function Substudy (PFS) of the Targeted Platelet Inhibition to Clarify the Optimal Strategy to Medically Manage Acute Coronary Syndromes (TRILOGY ACS) trial to evaluate the relationship between measurements of platelet reactivity and the longitudinal risks of predominantly spontaneous bleeding events among medically managed patients with unstable angina/non-ST-segment elevation myocardial infarction (UA/NSTEMI; collectively referred to as non-ST-segment elevation acute coronary syndrome, or NSTE ACS) who were treated with DAPT (aspirin+clopidogrel vs aspirin+prasugrel) for up to 30 months and to determine whether a threshold of LPR could be established that significantly delineated bleeding risk.

Methods

Study Population

The study design and results of the TRILOGY ACS trial have been described. ^{10,11} TRILOGY ACS was a double-blind, active-controlled, randomized trial in high-risk patients with NSTE ACS who were managed medically without planned revascularization. Participants had at least 1 of 4 enrichment criteria (age ≥60 years, diabetes mellitus, past myocardial infarction [MI], or past coronary revascularization at least 30 days before index ACS hospitalization). Patients with a history of transient ischemic attack/stroke, renal failure requiring

dialysis, or concomitant oral anticoagulant treatment were excluded. The TRILOGY ACS study was approved by regulatory authorities in all participating countries and by participating sites' institutional review boards. All participants provided written informed consent.

In the overall trial, 9326 participants at 966 sites in 52 countries were enrolled. Patients were randomly assigned to prasugrel or clopidogrel therapy in a double-blind, double-dummy fashion. The daily prasugrel maintenance dose was 10 mg in participants <75 years of age and 5 mg for study participants ≥75 years of age or who weighed <60 kg. The daily clopidogrel maintenance dose was 75 mg for all patients. Concomitant daily treatment with aspirin was strongly recommended, with low-dose aspirin strongly recommended. Treatment duration was up to 30 months, with a median treatment duration of 15 months and a median follow-up of 17 months. ¹⁰ Patients who required treatment with an oral anticoagulant (OAC) were excluded, and the study drug was stopped if a patient required treatment with an OAC during follow-up.

Platelet Function Substudy Protocol

A total of 25 countries participated in the TRILOGY ACS PFS. 12 All patients randomized into the main trial were included in the PFS at participating sites in those countries. The VerifyNow P2Y₁₂ assay (Accriva Diagnostics, San Diego, CA) was used to assess platelet reactivity to ADP measured in P2Y₁₂ reaction units (PRUs) to the randomized therapy, as previously described. 12 Sites were instructed to collect samples only for those patients taking blinded study drug. Platelet reactivity was assessed at baseline; at 2 hours after first dose of study drug; at 30 days; and at 3, 6, 12, 18, 24, and 30 months after randomization, independent of the occurrence of a bleeding event. Patients with at least 1 valid PRU measurement at 30 days or later were included in the analysis. Previous analyses from the TRILOGY ACS PFS demonstrated little inter- and intraindividual changes in serial PRU values over time. 12

Study Endpoints

All bleeding endpoints were prespecified in the trial protocol and were prospectively ascertained. 10,11 An independent cardiovascular adjudication committee adjudicated all suspected bleeding endpoints using the TIMI (Thrombolysis In Myocardial Infarction) bleeding classification scale. Bleeding endpoints were determined algorithmically from case report form data elements using the GUSTO (Global Use Strategies to Open Occluded Coronary Arteries) classification scale. Among participants who received at least 1 dose of study drug during the "at-risk" interval of actual study drug

treatment through 7 days after study drug discontinuation, non-coronary artery bypass graft (CABG)-related bleeding events were classified by the GUSTO bleeding scale as GUSTO severe/life-threatening, moderate, or mild bleeding, and by the TIMI bleeding scale as major, minor, or minimal, as previously defined. 11 The primary analyses used the 2-level composite GUSTO and TIMI bleeding endpoints (GUSTO severe/life-threatening or moderate bleeding; TIMI major or minor bleeding), given that we chose to focus upon consequential and clinically meaningful bleeding events that typically result in hospitalization. Further exploratory analyses extended to the 3-level composite bleeding endpoints for each classification scale (GUSTO severe/life-threatening, moderate, or mild bleeding; TIMI major, minor, or minimal bleeding), given the potential effects of mild/minimal bleeding events on study drug compliance. All bleeding analyses included only the 9240 patients who received at least 1 dose of the study drug.

Statistical Analysis

For this analysis, the "steady-state" PRU values were defined as those occurring at 5 days postrandomization, given that the first 2 PRU measurements obtained (at baseline and 2 hours following first study drug administration) did not reflect steadystate PRU values that would only be expected to occur after at least 5 days of treatment with maintenance doses of prasugrel or clopidogrel (there was no "reloading" of clopidogrel and prasugrel at the time of randomization for the 95% of patients who had been receiving clopidogrel before randomization). 10 To account for events that occurred between 5 and 30 days postrandomization, we assumed that the 30-day PRU value (the next value assessed after the 2-hour value per the study protocol) represented "steady-state treatment" at 5 days (when it was impractical to require patients to have an additional study visit solely for PRU measurement). Missing PRU values with a valid value after day 30 were used as the PRU value at 5 days (backward imputation). Forward imputation was used for patients randomized to clopidogrel who were already taking clopidogrel at home and had missing PRU values at 30 days or later (patient exclusions and imputation details are contained in Figure S1). 12

Baseline characteristics were compared by tertiles of steady-state PRU values to demonstrate how patient clinical characteristics differed by 3 categories of PRU response to the randomized study drug (clopidogrel vs prasugrel). Continuous variables are presented as medians and interquartile ranges. Categorical variables are presented as counts and percentages. Differences in baseline characteristics were tested among tertiles of steady-state PRU values. Continuous variables were compared using ANOVA when the assumption of normality was satisfied; otherwise, the Kruskal–Wallis test

was used. Categorical variables were compared using the chisquare test when cell frequencies were sufficient; otherwise, an exact test was used. Kaplan–Meier plots for the bleeding endpoints by PRU tertiles were analyzed for the 2-level composite bleeding endpoints.

To determine whether a PRU cut point existed that distinguished between high- and low-risk bleeding patients, we used the method of Contal and O'Quigley. 13 This method considers all possible observed values of steady-state PRU values and derives a standardized score statistic that can be used to test the null hypothesis that all observed values have equally likely risks of bleeding using the 2-level composites of GUSTO severe/life-threatening or moderate bleeding and TIMI major or minor bleeding. This test was used to determine whether the cut point that maximizes the score value is statistically different from other cut points with similar score values. However, given results from a past study that only demonstrated associations with clopidogrel metabolizer genomic variants and composite bleeding outcomes that incorporated mild bleeding events, we also separately performed analyses for PRU cut points that incorporated the 3-level composite bleeding endpoints for each classification scale (GUSTO severe/life-threatening, moderate, or mild bleeding; and TIMI major, minor, or minimal bleeding) to comprehensively assess the relationship between PRU values and bleeding risk. 14 As a result, 4 separate PRU cut points were determined.

To explore the unadjusted relationship between PRU values and bleeding outcomes, we grouped individuals according to the PRU value that maximized the score statistic regardless of whether it was a significant cut point. We then used these groups to create Kaplan—Meier plots of the cumulative distribution function and used the log-rank test to determine whether the survival functions (for bleeding endpoints) differed significantly between the groups. This testing procedure was analyzed completely separately for each of the 2- and 3-level composite GUSTO and TIMI bleeding composite outcomes (as previously described) to determine whether each of the 4 derived PRU cut points could reliably distinguish high versus low bleeding risk using the different composite outcomes from both bleeding classification scales.

To account for potential imbalances in baseline characteristics, we derived Cox proportional hazards models to assess the adjusted association between steady-state PRU values and time to first bleed using the GUSTO and TIMI bleeding composite endpoints, as previously described. Based upon previous analyses, we chose to use the following variables for adjustment: weight, age, clopidogrel stratum at time of randomization, aspirin dose category, time from randomization to treatment start, sex, disease classification, Killip class, previous peripheral arterial disease, previous peptic ulcer disease, systolic blood pressure, baseline hemoglobin, baseline

creatinine, baseline (prerandomization) PRU values, and concomitant beta-blocker use. 15-17 Additionally, we included a variable unique to TRILOGY ACS (use of angiography before randomization) given a previous analysis that demonstrated higher rates of bleeding for patients who underwent angiography before randomization. 18 To explore the relationship between steady-state PRU and time to first bleeding event, we constructed a series of models to evaluate the relationship between steady-state high versus low PRU values using the cut points we derived and PRU values (in a continuous fashion) with the 2- and 3-level GUSTO and TIMI composite bleeding endpoints. 13 We also analyzed the adjusted risks of bleeding in a restricted population of patients aged <75 years who were included in the primary efficacy analysis population of the overall TRILOGY ACS trial given that an exploratory treatment regimen (prasugrel 5 mg/day vs clopidogrel 75 mg/day) was studied in the elderly population (age ≥75 years). 10,11 Also, we performed a sensitivity analysis to evaluate the interactions between day 5 PRU values and randomized treatment with respect to bleeding outcomes.

All statistical tests were performed at a significance level of 0.05. All analyses were performed using SAS (version 9.3; SAS Institute Inc., Cary, NC) and R (version 2.14.1; R Foundation for Statistical Computing, Vienna, Austria) software by statisticians at the Duke Clinical Research Institute (Durham, NC), with an independent copy of the database. Dr Roe, the principal investigator for the TRILOGY ACS trial, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analyses.

Results

Platelet Function Substudy Participation

Among 9326 patients enrolled in TRILOGY ACS, 2690 (28%) were initially enrolled in the PFS. After database lock, it was determined that 13 of these patients were inaccurately listed as included in the PFS at randomization and 126 did not have a valid PRU measurement recorded, leaving a total of 2564 patients. Among the patients who received at least 1 dose of study drug, 2428 (26% of the total population) had a valid PRU measurement recorded at 30 days (for imputation of day 5 PRU results), and these patients were included in our analysis (Figure S1).

As previously published, the baseline clinical characteristics and efficacy (ischemic) outcomes were similar for patients who did versus did not participate in the PFS, and bleeding composite outcomes were also similar. ¹² Frequencies of GUSTO severe/life-threatening or moderate bleeding events and TIMI major or minor bleeding events were lower for patients who did versus did not participate in the PFS (Table S1).

Baseline Characteristics

Among the 2428 participants included in this analysis, baseline characteristics stratified by tertiles of baseline PRU values are shown in Table 1. Compared with participants in the middle and highest tertiles, participants in the lowest PRU tertile (PRU <105) were younger; more likely to be male; less likely to have diabetes mellitus; had higher body weight, higher baseline hemoglobin levels, and higher baseline creatinine clearance values; had a lower median Global Registry of Acute Coronary Events (GRACE) risk score; more commonly received the prasugrel 10-mg dose; and had the lowest median baseline PRU values assessed at the time of randomization before the first dose of study drug was administered (when pprox95% of the participants were being treated for the index ACS event with prerandomization clopidogrel). More elderly patients (≥75 years) and those with low body weight (<60 kg) were present in the highest PRU tertile (PRU >211), likely attributed to the use of a lower dose of prasugrel (5 mg) for these key subgroups. Baseline characteristics by the PRU cut point of <75 are detailed in Table S2.

Unadjusted Bleeding Outcomes

Using the 2-level composite bleeding endpoints for the primary analyses, 28 GUSTO severe/life-threatening or moderate bleeding events and 39 TIMI major or minor bleeding events not related to CABG occurred from randomization through the end of study follow-up. Starting at the landmark of 5 days postrandomization (the starting point for this analysis that corresponds with the steadystate day 5 PRU values), there were 27 GUSTO severe/lifethreatening or moderate bleeding events and 37 TIMI major or minor bleeding events not related to CABG that were included in these analyses. Gastrointestinal bleeding was the most common location for both GUSTO and TIMI bleeding events (Table 2). Bleeding event curves through 30 months by PRU tertiles overlapped during the first 12 months. The highest rates of bleeding through 30 months were observed for the middle PRU tertile (PRU 106-211) for both GUSTO and TIMI 2-level composite bleeding events (Figure 1A and 1B).

Using the 3-level composite bleeding endpoints, there were 297 GUSTO severe/life-threatening, moderate, or mild bleeding events and 290 TIMI major, minor, or minimal bleeding events, with bleeding locations shown in Table S3.

PRU Cut Points to Define Bleeding Risk

Using the method of Contal and O'Quigley, the best PRU cut points identified for GUSTO severe/life-threatening or

Table 1. Baseline Characteristics Stratified by Tertiles of P2Y₁₂ Reaction Unit (PRU) Values

	Day 5 PRU Tertiles			
Variable	PRU ≤105 (n=817)	PRU 106 to 211 (n=803)	PRU >211 (n=808)	P Value
Demographics		,	·	
Age, y*	63 (57, 70)	66 (59, 73)	67 (60, 75)	<0.001
≥75 y (%)	84/817 (10.3)	167/803 (20.8)	217/808 (26.9)	<0.00
Female sex (%)	277/817 (33.9)	293/803 (36.5)	376/808 (46.5)	<0.00
Weight, kg*	76.0 (65.8, 87.5)	75.0 (64.2, 87.0)	74.0 (62.3, 85.0)	0.002
<60 kg (%)	86/817 (10.5)	139/803 (17.3)	149/808 (18.4)	<0.00
Disease classification (%)			'	
NSTEMI	555/817 (67.9)	524/803 (65.3)	545/808 (67.5)	0.476
History (%)			'	
Diabetes mellitus	270/816 (33.1)	291/801 (36.3)	341/808 (42.2)	<0.00
Past MI	375/810 (46.3)	343/802 (42.8)	340/801 (42.4)	0.224
Past PCI	225/815 (27.6)	220/801 (27.5)	199/805 (24.7)	0.335
Past CABG	100/817 (12.2)	111/803 (13.8)	132/806 (16.4)	0.054
Past PAD	42/804 (5.2)	37/790 (4.7)	50/790 (6.3)	0.337
Past atrial fibrillation	51/802 (6.4)	76/791 (9.6)	78/791 (9.9)	0.021
Past heart failure	148/808 (18.3)	168/795 (21.1)	166/801 (20.7)	0.313
Past peptic ulcer disease	50/809 (6.2)	51/800 (6.4)	39/802 (4.9)	0.371
Baseline risk assessment		1	1	
Systolic BP, mm Hg*	127 (115, 138)	127 (116, 139)	130 (120, 140)	0.14
Killip class II to IV (%)	80/817 (9.8)	83/803 (10.3)	120/807 (14.9)	0.002
GRACE risk score*	115 (42, 201)	122 (54, 189)	126 (59, 205)	<0.00
Creatinine, mg/dL*	1.0 (0.8, 1.2)	1.0 (0.8, 1.2)	1.0 (0.8, 1.2)	0.548
CrCl, mL/min*	80.5 (61.3, 104.2)	73.9 (56.2, 97.8)	68.9 (51.1, 91.8)	<0.00
Hemoglobin, g/dL*	14.0 (13.1, 15.1)	13.8 (12.8, 14.9)	13.2 (12.2, 14.1)	<0.00
Prerandomization procedures (%)				
Angiography performed	334/817 (40.9)	313/803 (39.0)	295/808 (36.5)	0.193
Medications at randomization (%)		, ,	, ,	
Aspirin, daily dose, mg				
<100	325/817 (39.8)	343/803 (42.7)	300/808 (37.1)	0.073
100 to 250	361/817 (44.2)	353/803 (44.0)	394/808 (48.8)	0.091
>250	59/817 (7.2)	59/803 (7.3)	56/808 (6.9)	0.946
Beta-blocker	645/817 (78.9)	620/803 (77.2)	606/808 (75.0)	0.166
ACE-I/ARB	571/817 (69.9)	582/803 (72.5)	603/808 (74.6)	0.102
Statin	682/817 (83.5)	657/803 (81.8)	662/808 (81.9)	0.618
Proton pump inhibitor	164/817 (20.1)	210/803 (26.2)	193/808 (23.9)	0.014
Randomization-specific information			, ,	1
Clopidogrel stratum (%)				0.08
No prerandomization clopidogrel	35/817 (4.3)	38/803 (4.7)	40/808 (5.0)	
Clopidogrel started in-hospital; continued to randomization	578/817 (70.7)	516/803 (64.3)	537/808 (66.5)	
, , , , , , , , , , , , , , , , , , , ,	1	()	1 ()	1

Continued

Table 1. Continued

	Day 5 PRU Tertiles			
Variable	PRU ≤105 (n=817)	PRU 106 to 211 (n=803)	PRU >211 (n=808)	P Value
Randomized to prasugrel (%)	643/817 (78.7)	359/803 (44.7)	200/808 (24.8)	<0.001
Prasugrel 5-mg dose [†]	98/643 (15.2)	156/359 (43.5)	102/200 (51.0)	<0.001
Baseline, pre-randomization PRU*	181 (120, 250)	215 (163, 274)	273 (219, 315)	<0.001

ACE-I indicates angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker; BP, blood pressure; CABG, coronary artery bypass grafting; CrCl, creatinine clearance; GRACE, Global Registry of Acute Coronary Events; MI, myocardial infarction; NSTEMI, non-ST-segment elevation myocardial infarction; PAD, peripheral arterial disease; PCl, percutaneous coronary intervention; PRU, P2Y₁₂ reaction unit.

moderate bleeding events (PRU <106) and TIMI major or minor bleeding events (PRU <46) for the primary analyses did not significantly distinguish longitudinal bleeding risks using these 2-level bleeding composite endpoints (Figure 2A and 2B). For the exploratory analyses, the separately determined PRU cut points that maximized the score statistic were <75 both for the 3-level composite of GUSTO severe/lifethreatening, moderate, or mild bleeding events (unadjusted bleeding rates=26.5% for PRU values <75 vs 12.6% for PRU values ≥75) and for the 3-level composite of TIMI major, minor, or minimal bleeding events (unadjusted bleeding rates=25.9% vs 12.2%, respectively). Bleeding event curves distinguished by this cut point of <75 PRU (using the 3-level composite bleeding endpoints) separated early and continued to separate during the trial follow-up period (Figure 2C and 2D).

Table 2. Distribution of Bleeding Locations for the Primary Analyses (2-Level Bleeding)

Location	GUSTO Severe/Life-Threatening or Moderate Bleeding	TIMI Major or Minor Bleeding
Epistaxis	_	1
Gastrointestinal	11	22
Hematuria	_	1
No site identified	4	_
Other	4	4
Subdural hematoma	2	2
Surgical incision site	2	2
Urethral	1	1
Vaginal	1	2
Vascular access site	1	1
Missing	1	1
Total	27	37

GUSTO indicates Global Use of Strategies to Open Occluded Coronary Arteries; TIMI, Thrombolysis In Myocardial Infarction.

Adjusted Bleeding Outcomes

For the primary analyses, no significant association was found between continuous measures of PRU (per 10-unit decrease) with the adjusted risk of the 2-level composites of GUSTO severe/life-threatening or moderate bleeding or with TIMI major or minor bleeding (Table 3). For the exploratory analyses, using the 3-level GUSTO and TIMI composite bleeding endpoints that incorporated GUSTO mild and TIMI minimal bleeds, respectively, there was a significant increase in bleeding risk with continuous measures of PRU (per 10-unit decrease).

When the derived LPR cut points of PRU <46 for TIMI bleeding and PRU <106 for GUSTO bleeding were analyzed for the primary analyses, there was no significant association with the adjusted risk of the 2-level composites of GUSTO severe/life-threatening or moderate bleeding for PRU values below versus above the LPR cut point, and there was a marginally significant association with the adjusted risk of TIMI major or minor bleeding. For the exploratory analyses, there was an association with PRU values below versus above the LPR cut point of 75 for both the adjusted risks of the 3-level composites of GUSTO severe/life-threatening, moderate, or mild bleeding and for the TIMI major, minor, or minimal bleeding. Similar adjusted results were observed in the sensitivity analysis of the restricted population of patients aged <75 years (Table S4). Additional modeling showed no significant interactions between day 5 PRU values, randomized treatment, and bleeding outcomes.

Discussion

These hypothesis-generating findings demonstrate no clear relationship between LPR and the longitudinal risks of serious bleeding events (using both the GUSTO and TIMI bleeding classification scales) among patients with NSTE ACS who were managed without revascularization and treated with

^{*}Median (25th, 75th percentiles).

[†]Percentage of the overall patient group from each PRU tertile who received the prasugrel 5 mg/day maintenance dose.

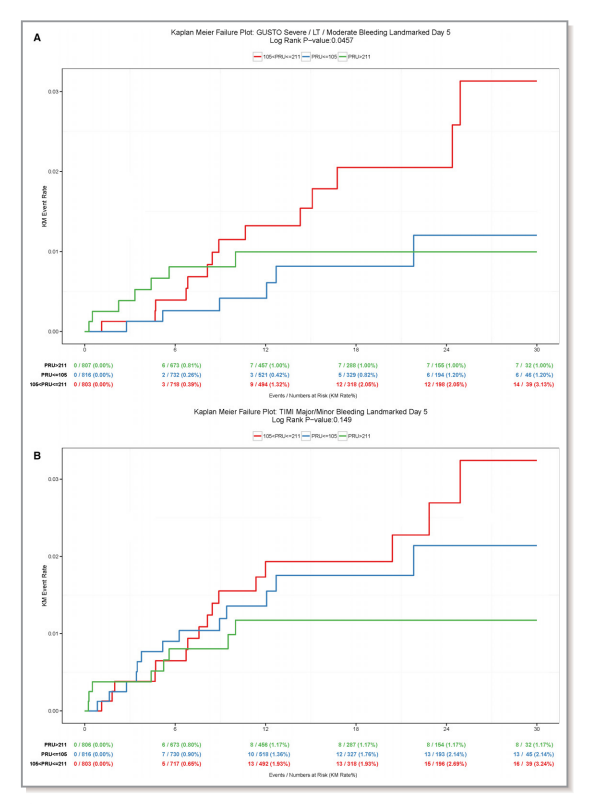


Figure 1. Cumulative Kaplan–Meier (KM) estimates of Global Use of Strategies to Open Occluded Coronary Arteries (GUSTO) severe/life-threatening (LT) or moderate (A) and Thrombolysis In Myocardial Infarction (TIMI) major or minor (B) bleeding events by P2Y₁₂ reaction unit (PRU) tertiles of distribution.

prolonged DAPT for up to 30 months. Only when mild/minimal events were incorporated into composite bleeding endpoints was an association with low PRU values and

bleeding risk demonstrated. Frequency of TIMI major or minor bleeding over 30 months was low (1.5%), however, and bleeding was primarily gastrointestinal in origin.

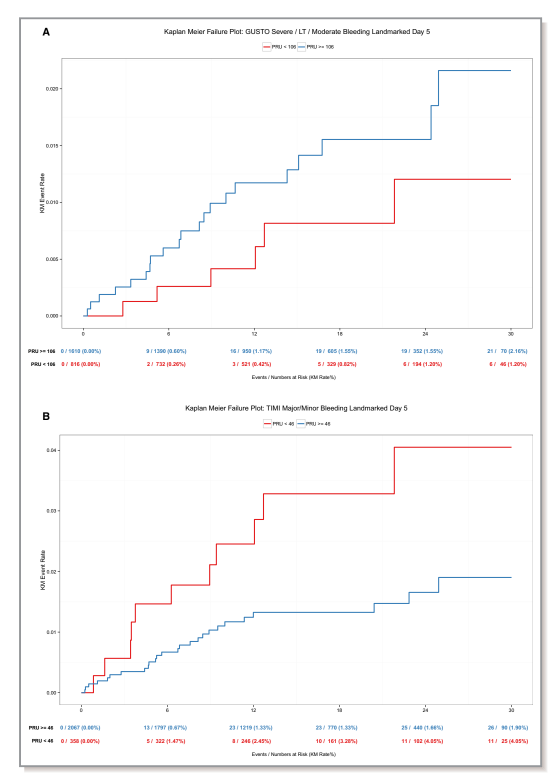


Figure 2. Cumulative Kaplan—Meier (KM) estimates of Global Use of Strategies to Open Occluded Coronary Arteries (GUSTO) severe/life-threatening (LT), or moderate bleeding (A); Thrombolysis In Myocardial Infarction (TIMI) major or minor bleeding (B); GUSTO severe/LT, moderate, or mild bleeding (C); and TIMI major, minor, or minimal bleeding (D) events by the derived low platelet reactivity cut point in P2Y₁₂ reaction units (PRUs).

This is the first large study that evaluated the 5-mg prasugrel dose used to mitigate bleeding risk and the relationship of PRU values with bleeding risk in patient

populations that are vulnerable and eligible for this dose (ie, those with low body weight and the elderly). However, our findings highlight how clinical characteristics associated with

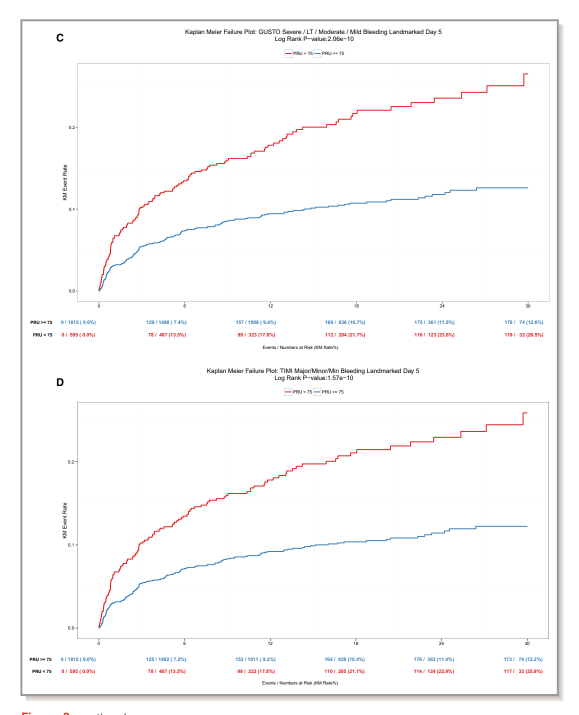


Figure 2. continued.

bleeding risk strongly influence platelet response to P2Y $_{12}$ inhibitors and thus may confound any potential relationship between PRU values and risks of serious bleeding events. In the current study, patients in the lowest PRU tertile (PRU \leq 105) were younger, had higher body weight, and had higher baseline creatinine clearance and hemoglobin values—all factors that are known to be associated with a lower risk of short- and intermediate-term bleeding among patients with ACS. $^{19-23}$ Whereas patients in the lowest PRU tertile were

more likely to be randomized to prasugrel and receive the prasugrel 10-mg maintenance dose (as expected from our previous evaluation of the PFS data according to randomized treatment assignment), the unadjusted risks of GUSTO severe/life-threatening or moderate and TIMI major or minor bleeding were highest among patients in the middle PRU tertile (PRU 106–211). Additionally, we have separately shown that elderly patients (≥75 years) from the TRILOGY ACS study population had a 2- to 3-fold increased risk of both

Table 3. Adjusted Associations of GUSTO and TIMI Composite Bleeding Definitions With Continuous PRU Distributions and the Derived Cut Points for Low Versus High Platelet Reactivity in All Patients

	Adjusted HR (95% CI)	P Value		
GUSTO severe/life-threatening or moderat	e non-CABG bleeding			
Continuous day 5 PRU (per 10-unit decrease)	1.01 (0.96–1.06)	0.82		
Dichotomous (<106) day 5 PRU (LPR vs HPR)*	0.68 (0.25–1.87)	0.46		
GUSTO severe/life-threatening, moderate,	or mild non-CABG blo	eeding		
Continuous day 5 PRU (per 10-unit decrease)	1.04 (1.02–1.05)	<0.001		
Dichotomous (<75) day 5 PRU (LPR vs HPR)*	2.30 (1.72–3.07)	<0.001		
TIMI major or minor non-CABG bleeding				
Continuous day 5 PRU (per 10-unit decrease)	1.02 (0.98–1.07)	0.37		
Dichotomous (<46) day 5 PRU (LPR vs HPR)*	2.35 (1.00–5.52)	0.05		
TIMI major, minor, or minimal non-CABG bleeding				
Continuous day 5 PRU (per 10-unit decrease)	1.04 (1.02–1.06)	<0.001		
Dichotomous (<75) day 5 PRU (LPR vs HPR)*	2.34 (1.74–3.14)	<0.001		

CABG indicates coronary artery bypass graft; GUSTO, Global Use of Strategies to Open Occluded Coronary Arteries; HPR, high platelet reactivity; HR, hazard ratio; LPR, low platelet reactivity; PRU, P2Y₁₂ reaction unit; TIMI, Thrombolysis In Myocardial Infarction. *The 4 derived cut points to determine bleeding risk were separately determined for each of the 2- and 3-level TIMI and GUSTO composite bleeding outcomes.

GUSTO and TIMI bleeding (using 2-level bleeding composite endpoints) when treated with either clopidogrel 75 mg/day or prasugrel 5 mg/day, as compared to younger patients.²⁴ The underlying factors associated with increased bleeding risks for elderly patients are likely multifactorial (lower body weight, lower baseline creatinine clearance, and lower hemoglobin values compared to younger patients) and inter-related, but we observed similar findings in our adjusted analysis of the relationship of PRU values with bleeding risks when elderly patients were excluded. We previously observed that elderly patients had a less-robust PRU response to clopidogrel 75 mg daily compared to younger patients, so older age may be a much stronger contributor to bleeding risk irrespective of on-treatment PRU response to a P2Y₁₂ inhibitor. 12 Finally, our study is the first large study that included both a third-generation P2Y₁₂ inhibitor (prasugrel) and clopidogrel when assessing the association of bleeding risk with PRU values. Further investigation is therefore needed to ascertain how interactions between clinical characteristics, the dose/type of $P2Y_{12}$ inhibitor chosen for an individual patient, and ontreatment PRU values influence serious bleeding rates.

In contrast to the medically managed population studied in TRILOGY ACS, observational studies in patients treated with PCI have suggested that LPR during DAPT treatment may be associated with major bleeding risk. 7-9,25,26 A prospective, randomized trial that leveraged bedside PRU monitoring to inform antiplatelet treatment decisions did not confirm this relationship, but findings from the ADAPT-DES prospective registry demonstrated an inverse relationship between high platelet reactivity (PRU >208) and clinically relevant bleeding in patients undergoing PCI. 23,27,28 After successful PCI, lower platelet reactivity on clopidogrel was an independent predictor of postdischarge bleeding, and these bleedings had a strong relationship with mortality at the 2-year follow-up point.²³ Another recent study in a cohort of patients who underwent elective PCI suggested that LPR provided incremental predictive value for bleeding events through 30 days compared with a bleeding risk score.²⁹ Although the influence of platelet reactivity on bleeding risk may differ for patients who undergo PCI versus ACS patients who are managed without revascularization, the primary 2-level composite bleeding events in TRILOGY ACS occurred infrequently and were primarily spontaneous and unrelated to cardiovascular procedures. The present analysis from TRILOGY ACS thus provides novel evidence for the relationship of platelet reactivity measurements with bleeding risk for ACS patients treated with DAPT who did not undergo PCI.

Limitations

A number of limitations to our analysis should be noted. First, PRU values were missing across all time periods, and multiple imputation techniques were used to account for missing values. The back-imputation technique used to estimate day 5 PRU values requires assumptions about the stability of drug effects and steady state at 5 days postbaseline that may not be accurate. Second, the number of serious bleeding events was small, so this study was underpowered to determine whether there was a significant difference in bleeding risk using the 2-level composite GUSTO and TIMI bleeding outcomes. However, this is the largest platelet function substudy that has been embedded within a randomized clinical trial comparing post-ACS DAPT regimens beyond clopidogrel, so it is unlikely that a larger study will be conducted in the future to capture more-serious bleeding events. Third, the frequencies of GUSTO severe/life-threatening or moderate bleeding events and TIMI major or minor bleeding events were lower for patients who did versus did not participate in the PFS. These findings could be attributed to regional differences in the reporting and/or querying of suspected bleeding events that were further confounded by

the choice of countries that participated in the PFS, but we were not able to investigate these potential assumptions. Finally, we did not analyze how clopidogrel metabolizer genomic variants influenced the relationship of bleeding risk with DAPT treatment in this analysis because we chose to focus solely upon the relationship of platelet reactivity measurements (regardless of genomic status and type/dose of P2Y $_{12}$ inhibitor used). 14

Conclusions

Among NSTE ACS patients managed without revascularization and receiving prolonged DAPT treatment, PRU values were not significantly associated with long-term serious bleeding risk. These hypothesis-generating results suggest that LPR does not independently predict the risk of serious bleeding during the period of DAPT treatment post-ACS.

Acknowledgments

The authors thank the following: Karen Pieper, MS, for expert coordination and management of the statistical analytic team; Jonathan McCall, MS, for expert editorial assistance; and Kerry Stenke for expert graphics assistance. Pieper, McCall, and Stenke are employees of the Duke Clinical Research Institute (Durham, NC); none received any compensation for their work on this manuscript other than their usual salaries.

Sources of Funding

The TRILOGY ACS study was supported by Daiichi Sankyo Incorporated and Eli Lilly and Company. An employee of Eli Lilly (Dr Jakubowski) participated as an author and provided review and comments for drafts of the manuscript. The decisions regarding the design and conduct of the study; the collection, management, analysis, and interpretation of the data; the drafting of the manuscript; the determination of the final content of the manuscript; and the decision to submit the manuscript were made independently by the investigators. All data analyses were performed independently by statisticians from the Duke Clinical Research Institute (Durham, NC), utilizing an independent copy of the database.

Disclosures

Cornel reports receiving consulting payments from Eli Lilly, Merck Sharp and Dohme, AstraZeneca, and Merck. Ohman reports receiving grant support and travel expenses from Daiichi Sankyo and Eli Lilly; consulting fees from AstraZeneca, Boehringer Ingelheim, Bristol-Myers Squibb, Gilead Sciences, Janssen Pharmaceuticals, Liposcience, Merck, Pozen, Hoffmann-La Roche, Sanofi-Aventis, The Medicines Company, and

Web MD; grant support from Gilead Sciences; and lecture fees from Gilead Sciences, Boehringer Ingelheim, and The Medicines Company. Jakubowski is an employee and minor shareholder of Eli Lilly and Company. Bhatt discloses the following relationships: Advisory board: Cardax, Elsevier Practice Update Cardiology, Medscape Cardiology, and Regado Biosciences; board of directors: Boston VA Research Institute, Society of Cardiovascular Patient Care; chair: American Heart Association Get With The Guidelines Steering Committee; data monitoring committees: Duke Clinical Research Institute, Harvard Clinical Research Institute, Mayo Clinic, Population Health Research Institute; honoraria: American College of Cardiology (senior associate editor, Clinical Trials and News, ACC.org), Belvoir Publications (editor in chief, Harvard Heart Letter), Duke Clinical Research Institute (clinical trial steering committees), Harvard Clinical Research Institute (clinical trial steering committee), HMP Communications (editor in chief, Journal of Invasive Cardiology), Journal of the American College of Cardiology (associate editor), Population Health Research Institute (clinical trial steering committee), Slack Publications (chief medical editor, Cardiology Today's Intervention), and WebMD (CME steering committees); other: Clinical Cardiology (deputy editor); research funding: Amarin, AstraZeneca, Biotronik, Bristol-Myers Squibb, Eisai, Ethicon, Forest Laboratories, Ischemix, Medtronic, Pfizer, Roche, Sanofi Aventis, St. Jude Medical, and The Medicines Company; trustee: American College of Cardiology; and unfunded research: FlowCo, PLx Pharma, and Takeda. White reports receiving grant support from Sanofi-Aventis, Eli Lilly, The Medicines Company, NIH, Pfizer, Roche, Johnson & Johnson, Schering-Plough, Merck Sharp & Dohme, AstraZeneca, GlaxoSmithKline, Daiichi Sankyo Pharma Development, and Bristol-Myers Squibb; he also participates in advisory boards for Merck Sharpe & Dohme, Roche, and Regado Biosciences. Ardissino reports receiving consulting payments from Eli Lilly. Fox reports receiving research grants from Lilly, Bayer, Johnson & Johnson, and AstraZeneca; speakers bureau payments from Bayer, Johnson & Johnson, AstraZeneca, and Sanofi-Aventis; and consulting/other payments from Lilly, Bayer, Johnson & Johnson, AstraZeneca, Sanofi-Aventis, Boehringer Ingelheim, and Eli Lilly. Prabhakaran reports receiving research grants from Eli Lilly and the Medtronic Foundation and honoraria from Eli Lilly. Armstrong reports receiving consulting fees from Eli Lilly, Hoffmann-La Roche, Merck, Axio Research, and Orexigen; grant support from Boehringer Ingelheim, Hoffmann-La Roche, Sanofi-Aventis, Scios, Ortho Biotech, Johnson & Johnson, Janssen Pharmaceuticals, GlaxoSmithKline, Amylin Pharmaceuticals, and Merck; and payment for developing educational presentations from AstraZeneca and Eli Lilly and Company. Erlinge reports receiving consulting payments from Eli Lilly. Gurbel reports serving as a consultant for Daiichi Sankyo, Sankyo, Lilly, Pozen, Bayer, AstraZeneca, Accumetrics, Nanosphere, Sanofi-Aventis,

Boehringer Ingelheim, Merck, Medtronic, CSL, and Haemonetics; receiving grants from NIH, Daiichi Sankyo, Lilly, Pozen CSL, AstraZeneca, Sanofi-Aventis, Haemoscope, Harvard Clinical Research Institute, and Duke Clinical Research Institute; receiving payment for lectures, including service on speakers' bureaus, from Lilly, Daiichi Sankyo, Nanosphere, and Merck; receiving payment for development of educational presentations from Merck, the Discovery Channel, and Pri-Med; holding stock or stock options in Merck and Pfizer; and holding patents in the area of personalized antiplatelet therapy and interventional cardiology. Roe reports research grants from Eli Lilly and Company, Janseen Pharmaceuticals, Sanofi-Aventis, Daiichi-Sankyo, Familial Hypercholesterolemia Foundation, and Ferring Pharmaceuticals; educational activities or lectures for Amgen and Bristol-Myers Squibb; and consulting or other services for AstraZeneca, Eli Lilly and Company, Merck & Co, Elsevier Publishers, Amgen, Boehringer-Ingelheim, and PriMed. All conflicts of interest are listed at https://www.dcri.org/ about-us/conflict-of-interest. The remaining authors have no conflicts to disclose.

References

- 1. Roffi M, Patrono C, Collet JP, Mueller C, Valgimigli M, Andreotti F, Bax JJ, Borger MA, Brotons C, Chew DP, Gencer B, Hasenfuss G, Kjeldsen K, Lancellotti P, Landmesser U, Mehilli J, Mukherjee D, Storey RF, Windecker S, Baumgartner H, Gaemperli O, Achenbach S, Agewall S, Badimon L, Baigent C, Bueno H, Bugiardini R, Carerj S, Casselman F, Cuisset T, Erol Ç, Fitzsimons D, Halle M, Hamm C, Hildick-Smith D, Huber K, Iliodromitis E, James S, Lewis BS, Lip GY, Piepoli MF, Richter D, Rosemann T, Sechtem U, Steg PG, Vrints C, Luis Zamorano J. 2015 ESC guidelines for the management of acute coronary syndromes in patients presenting without persistent ST-segment elevation: Task Force for the Management of Acute Coronary Syndromes in Patients Presenting without Persistent ST-Segment Elevation of the European Society of Cardiology (ESC). Eur Heart J. 2016;37:267–315.
- 2. Amsterdam EA, Wenger NK, Brindis RG, Casey DE Jr, Ganiats TG, Holmes DR Jr, Jaffe AS, Jneid H, Kelly RF, Kontos MC, Levine GN, Liebson PR, Mukherjee D, Peterson ED, Sabatine MS, Smalling RW, Zieman SJ; American College of Cardiology; American Heart Association Task Force on Practice Guidelines; Society for Cardiovascular Angiography and Interventions; Society of Thoracic Surgeons; American Association for Clinical Chemistry. 2014 AHA/ACC guideline for the management of patients with non-ST-elevation acute coronary syndromes: executive summary: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines. J Am Coll Cardiol. 2014;64:e139—e228.
- Yusuf S, Zhao F, Mehta SR, Chrolavicius S, Tognoni G, Fox KK; Clopidogrel in Unstable Angina to Prevent Recurrent Events Trial Investigators. Effects of clopidogrel in addition to aspirin in patients with acute coronary syndromes without ST-segment elevation. N Engl J Med. 2001;345:494–502.
- Wiviott SD, Braunwald E, McCabe CH, Montalescot G, Ruzyllo W, Gottlieb S, Neumann FJ, Ardissino D, De Servi S, Murphy SA, Riesmeyer J, Weerakkody G, Gibson CM, Antman EM; TRITON-TIMI 38 Investigators. Prasugrel versus clopidogrel in patients with acute coronary syndromes. N Engl J Med. 2007;357:2001–2015.
- Wallentin L, Becker RC, Budaj A, Cannon CP, Emanuelsson H, Held C, Horrow J, Husted S, James S, Katus H, Mahaffey KW, Scirica BM, Skene A, Steg PG, Storey RF, Harrington RA; PLATO Investigators, Freij A, Thorsén M. Ticagrelor versus clopidogrel in patients with acute coronary syndromes. N Engl J Med. 2009;361:1045–1057.
- Kearney JF Jr. Balancing the risks and benefits of dual platelet inhibition. N Engl J Med. 2015;372:1854–1856.
- Sibbing D, Schulz S, Braun S, Morath T, Stegherr J, Mehilli J, Schömig A, von Beckerath N, Kastrati A. Antiplatelet effects of clopidogrel and bleeding in patients undergoing coronary stent placement. J Thromb Haemost. 2010;8:250–256.

- Patti G, Pasceri V, Vizzi V, Ricottini E, Di Sciascio G. Usefulness of platelet response to clopidogrel by point-of-care testing to predict bleeding outcomes in patients undergoing percutaneous coronary intervention (from the Antiplatelet Therapy for Reduction of Myocardial Damage During Angioplasty-Bleeding Study). Am J Cardiol. 2011;107:995–1000.
- Tantry US, Bonello L, Aradi D, Price MJ, Jeong YH, Angiolillo DJ, Stone GW, Curzen N, Geisler T, Ten Berg J, Kirtane A, Siller-Matula J, Mahla E, Becker RC, Bhatt DL, Waksman R, Rao SV, Alexopoulos D, Marcucci R, Reny JL, Trenk D, Sibbing D, Gurbel PA; Working Group on On-Treatment Platelet Reactivity. Consensus and update on the definition of on-treatment platelet reactivity to ADP associated with ischemia and bleeding. J Am Coll Cardiol. 2013;62:2261– 2273.
- 10. Roe MT, Armstrong PW, Fox KA, White HD, Prabhakaran D, Goodman SG, Cornel JH, Bhatt DL, Clemmensen P, Martinez F, Ardissino D, Nicolau JC, Boden WE, Gurbel PA, Ruzyllo W, Dalby AJ, McGuire DK, Leiva-Pons JL, Parkhomenko A, Gottlieb S, Topacio GO, Hamm C, Pavlides G, Goudev AR, Oto A, Tseng CD, Merkely B, Gasparovic V, Corbalan R, Cinteză M, McLendon RC, Winters KJ, Brown EB, Lokhnygina Y, Aylward PE, Huber K, Hochman JS, Ohman EM; TRILOGY ACS Investigators. Prasugrel versus clopidogrel for acute coronary syndromes without revascularization. N Engl J Med. 2012;367:1297–1309.
- 11. Chin CT, Roe MT, Fox KA, Prabhakaran D, Marshall DA, Petitjean H, Lokhnygina Y, Brown E, Armstrong PW, White HD, Ohman EM; TRILOGY ACS Steering Committee. Study design and rationale of a comparison of prasugrel and clopidogrel in medically managed patients with unstable angina/non-ST-segment elevation myocardial infarction: the TaRgeted platelet Inhibition to cLarify the Optimal strateGy to medicallY manage Acute Coronary Syndromes (TRILOGY ACS) trial. Am Heart J. 2010;160:16–22.e1.
- 12. Gurbel PA, Erlinge D, Ohman EM, Neely B, Neely M, Goodman SG, Huber K, Chan MY, Cornel JH, Brown E, Zhou C, Jakubowski JA, White HD, Fox KA, Prabhakaran D, Armstrong PW, Tantry US, Roe MT; TRILOGY ACS Platelet Function Substudy Investigators. Platelet function during extended prasugrel and clopidogrel therapy for patients with acs treated without revascularization: the TRILOGY ACS platelet function substudy. JAMA. 2012;308:1785–1794.
- Contal C, O'Quigley J. An application of changepoint methods in studying the effect of age on survival in breast cancer. Comput Stat Data Anal. 1999;30:253–270.
- 14. Bhatt DL, Paré G, Eikelboom JW, Simonsen KL, Emison ES, Fox KA, Steg PG, Montalescot G, Bhakta N, Hacke W, Flather MD, Mak KH, Cacoub P, Creager MA, Berger PB, Steinhubl SR, Murugesan G, Mehta SR, Kottke-Marchant K, Lincoff AM, Topol EJ; CHARISMA Investigators. The relationship between CYP2C19 polymorphisms and ischaemic and bleeding outcomes in stable outpatients: the CHARISMA genetics study. Eur Heart J. 2012;33:2143–2150.
- 15. Ferguson JJ, Califf RM, Antman EM, Cohen M, Grines CL, Goodman S, Kereiakes DJ, Langer A, Mahaffey KW, Nessel CC, Armstrong PW, Avezum A, Aylward P, Becker RC, Biasucci L, Borzak S, Col J, Frey MJ, Fry E, Gulba DC, Guneri S, Gurfinkel E, Harrington R, Hochman JS, Kleiman NS, Leon MB, Lopez-Sendon JL, Pepine CJ, Ruzyllo W, Steinhubl SR, Teirstein PS, Toro-Figueroa L, White H; SYNERGY Trial Investigators. Enoxaparin vs unfractionated heparin in high-risk patients with non-ST-segment elevation acute coronary syndromes managed with an intended early invasive strategy: primary results of the SYNERGY randomized trial. JAMA. 2004;292:45–54.
- 16. Becker RC, Bassand JP, Budaj A, Wojdyla DM, James SK, Cornel JH, French J, Held C, Horrow J, Husted S, Lopez-Sendon J, Lassila R, Mahaffey KW, Storey RF, Harrington RA, Wallentin L. Bleeding complications with the P2Y12 receptor antagonists clopidogrel and ticagrelor in the PLATelet inhibition and patient Outcomes (PLATO) trial. Eur Heart J. 2011;32:2933–2944.
- 17. Giugliano RP, White JA, Bode C, Armstrong PW, Montalescot G, Lewis BS, van't Hof A, Berdan LG, Lee KL, Strony JT, Hildemann S, Veltri E, Van de Werf F, Braunwald E, Harrington RA, Califf RM, Newby LK; EARLY ACS Investigators. Early versus delayed, provisional eptifibatide in acute coronary syndromes. N Engl J Med. 2009;360:2176–2190.
- 18. Wiviott SD, White HD, Ohman EM, Fox KA, Armstrong PW, Prabhakaran D, Hafley G, Lokhnygina Y, Boden WE, Hamm C, Clemmensen P, Nicolau JC, Menozzi A, Ruzyllo W, Widimsky P, Oto A, Leiva-Pons J, Pavlides G, Winters KJ, Roe MT, Bhatt DL. Prasugrel versus clopidogrel for patients with unstable angina or non-ST-segment elevation myocardial infarction with or without angiography: a secondary, prespecified analysis of the TRILOGY ACS trial. Lancet. 2013;382:605–613.
- Moscucci M, Fox KA, Cannon CP, Klein W, López-Sendón J, Montalescot G, White K, Goldberg RJ. Predictors of major bleeding in acute coronary syndromes: the Global Registry of Acute Coronary Events (GRACE). Eur Heart J. 2003;24:1815–1823.
- 20. Subherwal S, Bach RG, Chen AY, Gage BF, Rao SV, Newby LK, Wang TY, Gibler WB, Ohman EM, Roe MT, Pollack CV Jr, Peterson ED, Alexander KP. Baseline risk of major bleeding in non-ST-segment-elevation myocardial infarction: the CRUSADE (Can Rapid risk stratification of Unstable angina patients Suppress

- ADverse outcomes with Early implementation of the ACC/AHA Guidelines) Bleeding Score. *Circulation*. 2009;119:1873–1882.
- 21. Mehran R, Pocock SJ, Nikolsky E, Clayton T, Dangas GD, Kirtane AJ, Parise H, Fahy M, Manoukian SV, Feit F, Ohman ME, Witzenbichler B, Guagliumi G, Lansky AJ, Stone GW. A risk score to predict bleeding in patients with acute coronary syndromes. *J Am Coll Cardiol*. 2010;55:2556–2566.
- 22. Mathews R, Peterson ED, Chen AY, Wang TY, Chin CT, Fonarow GC, Cannon CP, Rumsfeld JS, Roe MT, Alexander KP. In-hospital major bleeding during ST-elevation and non-ST-elevation myocardial infarction care: derivation and validation of a model from the ACTION Registry[®]-GWTG[™]. Am J Cardiol. 2011;107:1136-1143.
- Genereux P, Giustino G, Witzenbichler B, Weisz G, Stuckey TD, Rinaldi MJ, Neumann FJ, Metzger DC, Henry TD, Cox DA, Duffy PL, Mazzaferri E, Yadav M, Francese DP, Palmerini T, Kirtane AJ, Litherland C, Mehran R, Stone GW. Incidence, predictors, and impact of post-discharge bleeding after percutaneous coronary intervention. J Am Coll Cardiol. 2015;66: 1036–1145.
- 24. Roe MT, Goodman SG, Ohman EM, Stevens SR, Hochman JS, Gottlieb S, Martinez F, Dalby AJ, Boden WE, White HD, Prabhakaran D, Winters KJ, Aylward PE, Bassand JP, McGuire DK, Ardissino D, Fox KA, Armstrong PW. Elderly patients with acute coronary syndromes managed without revascularization: Insights into the safety of long-term dual antiplatelet therapy with reduced-dose prasugrel versus standard-dose clopidogrel. Circulation. 2013;128:823–833.
- Cuisset T, Cayla G, Frere C, Quilici J, Poyet R, Gaborit B, Bali L, Morange PE, Alessi MC, Bonnet JL. Predictive value of post-treatment platelet reactivity for occurrence of post-discharge bleeding after non-ST elevation acute coronary

- syndrome. Shifting from antiplatelet resistance to bleeding risk assessment? *EuroIntervention*. 2009;5:325–329.
- 26. Aradi D, Kirtane A, Bonello L, Gurbel PA, Tantry US, Huber K, Freynhofer MK, ten Berg J, Janssen P, Angioililo DJ, Siller-Matula JM, Marcucci R, Patti G, Mangiacapra F, Valgimigli M, Morel O, Palmerini T, Price MJ, Cuisset T, Kastrati A, Stone GW, Sibbing D. Bleeding and stent thrombosis on P2Y12-inhibitors: collaborative analysis on the role of platelet reactivity for risk stratification after percutaneous coronary intervention. Eur Heart J. 2015;36:1762–1771.
- 27. Collet JP, Cuisset T, Rangé G, Cayla G, Elhadad S, Pouillot C, Henry P, Motreff P, Carrié D, Boueri Z, Belle L, Van Belle E, Rousseau H, Aubry P, Monségu J, Sabouret P, O'Connor SA, Abtan J, Kerneis M, Saint-Etienne C, Barthélémy O, Beygui F, Silvain J, Vicaut E, Montalescot G; ARCTIC Investigators. Bedside monitoring to adjust antiplatelet therapy for coronary stenting. N Engl J Med. 2012;367:2100–2109.
- 28. Stone GW, Witzenbichler B, Weisz G, Rinaldi MJ, Neumann FJ, Metzger DC, Henry TD, Cox DA, Duffy PL, Mazzaferri E, Gurbel PA, Xu K, Parise H, Kirtane AJ, Brodie BR, Mehran R, Stuckey TD; ADAPT-DES Investigators. Platelet reactivity and clinical outcomes after coronary artery implantation of drugeluting stents (ADAPT-DES): a prospective multicentre registry study. *Lancet*. 2013;382:614–623.
- 29. Mangiacapra F, Ricottini E, Barbato E, Demartini C, Peace A, Patti G, Vizzi V, De Bruyne B, Wijns W, Di Sciascio G. Incremental value of platelet reactivity over a risk score of clinical and procedural variables in predicting bleeding after percutaneous coronary intervention via the femoral approach: development and validation of a new bleeding risk score. Circ Cardiovasc Interv. 2015;8: e002106.

Supplemental Material

Figure S1. Consort diagram demonstrating patient flow and PRU imputation approaches

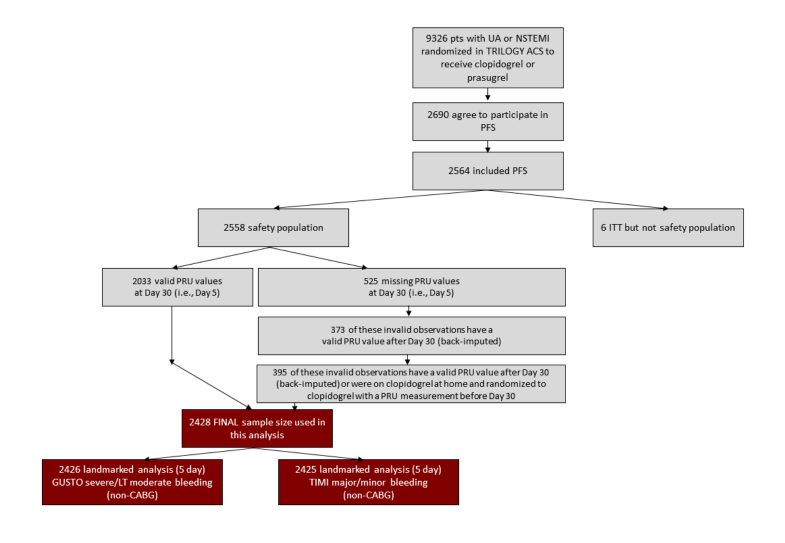


Table S1. Bleeding event rates by participation in the PFS*

	Included in PFS (N=2428)	Not Included in PFS (N=6812)
GUSTO severe/life-threatening or moderate bleeding (%)	1.83%	3.63%
TIMI major or minor bleeding (%)	2.24%	3.81%

^{*}Kaplan-Meier estimates of bleeding rates through 30 months PFS, Platelet Function Sub-Study

Table S2. Baseline characteristics stratified by PRU values

	PRU <75	PRU ≥75	P-
	(N=601)	(N=1827)	value
Demographics			
Age, yrs	62.0 (56.0,	66.0 (60.0,	<0.00
	69.0)	74.0)	1
Female sex	212/601	734/1827	0.033
	(35.3)	(40.2)	
White race	388/601	1120/1827	0.153
	(64.6)	(61.3)	
Weight, kg	76.0 (66.7,	75.0 (63.0,	0.004
	87.5)	86.1)	
NSTEMI	403/601	1221/1827	0.919
	(67.1)	(66.8)	
Killip class II-IV	52/601 (8.7)	231/1826	0.008
		(12.7)	
Time from FMC to treatment start, hrs	99.8 (54.9,	108.9 (63.0,	0.211
	157.8)	160.9)	
CV risk factors			
Family history of CAD	179/536	503/1640	0.238
	(33.4)	(30.7)	
Hypertension	480/598	1508/1823	0.174
	(80.3)	(82.7)	
Hyperlipidemia	318/541	1011/1699	0.765
	(58.8)	(59.5)	
Diabetes mellitus	190/600	712/1825	0.001
	(31.7)	(39.0)	
Current/recent smoking	118/594	322/1808	0.261
	(19.9)	(17.8)	
Prior peptic ulcer disease	38/596 (6.4)	102/1815	0.494

	PRU <75	PRU ≥75	P-
	(N=601)	(N=1827)	value
		(5.6)	
Angiography performed	252/601	690/1827	0.069
	(41.9)	(37.8)	
CV disease history			
Prior myocardial infarction	282/595	776/1818	0.044
	(47.4)	(42.7)	
Prior PCI	170/599	474/1822	0.256
	(28.4)	(26.0)	
Prior CABG	72/601 (12.0)	271/1825	0.080
		(14.8)	
Prior PAD	28/590 (4.7)	101/1794	0.410
		(5.6)	
Prior atrial fibrillation	33/589 (5.6)	172/1795	0.003
		(9.6)	
Prior chronic heart failure	110/593	372/1811	0.293
	(18.5)	(20.5)	
Baseline labs and measurments			
GRACE risk score	114.0 (42.0,	123.0 (54.0,	<0.00
	201.0)	205.0)	1
Creatinine	1.0 (0.8, 1.1)	1.0 (0.8, 1.2)	0.094
CrCL, mL/min	82.3 (62.5,	71.8 (54.0,	<0.00
	105.6)	94.7)	1
Systolic BP, mmHg	127.0 (115.0,	129.0 (118.0,	0.334
	138.0)	140.0)	
Heart rate, bpm	68.0 (61.0,	70.0 (62.0,	0.068
	75.0)	76.0)	
Hemoglobin	14.0 (13.0,	13.5 (12.5,	<0.00
	15.1)	14.6)	1

Concomitant medications at

	PRU <75	PRU ≥75	P-
	(N=601)	(N=1827)	value
randomization			
Daily dose <100 mg	233/601	735/1827	0.526
	(38.8)	(40.2)	
Daily dose 100-250 mg	266/601	842/1827	0.435
	(44.3)	(46.1)	
Daily dose >250 mg	43/601 (7.2)	131/1827	0.990
		(7.2)	
Beta-blocker	476/601	1395/1827	0.150
	(79.2)	(76.4)	
ACE-I/ARB	419/601	1337/1827	0.100
	(69.7)	(73.2)	
Statin	502/601	1499/1827	0.408
	(83.5)	(82.0)	
Proton pump inhibitor	121/601	446/1827	0.032
	(20.1)	(24.4)	
Randomization specific information			
Clopidogrel strata			0.014
1	24/601 (4.0)	89/1827 (4.9)	
2	433/601	1198/1827	
	(72.0)	(65.6)	
3	144/601	540/1827	
	(24.0)	(29.6)	
Randomized treatment	503/601	699/1827	<0.00
	(83.7)	(38.3)	1
Duration of clopidogrel use before	108.3 (62.8,	107.9 (65.0,	0.794
treatment start, hrs	149.3)	156.6)	
Age ≥75 yrs	41/601 (6.8)	427/1827	<0.00
		(23.4)	1
Weight <60 kg	50/601 (8.3)	324/1827	<0.00

	PRU <75	PRU ≥75	P-
	(N=601)	(N=1827)	value
		(17.7)	1
Prasugrel maintenance 5 mg	54/503 (10.7)	302/699	<0.00
		(43.2)	1
Baseline PRU values	179.0 (115.0,	238.0 (179.0,	<0.00
	249.0)	295.0)	1

Data are presented as medians (25th, 75th percentiles) or n/N (%).

ACE-I/ARB, angiotensin-converting enzyme inhibitor/angiotensin receptor blocker; BP, blood pressure; CAD, coronary artery disease; CABG, coronary artery bypass graft; CrCl, creatinine clearance; CV, cardiovascular; FMC, first medical contact; GRACE, Global Registry of Acute Coronary Events; PAD, peripheral artery disease; PCI, percutaneous coronary intervention; PFS, Platelet Function Substudy; PRU, P2Y12 reaction unit; NSTEMI, non-ST-segment elevation myocardial infarction.

Table S3. Distribution of bleeding locations for the exploratory analyses (3-level bleeding)

Location	GUSTO severe/life- threatening or moderate or mild bleeding	TIMI major or minor or minimal bleeding
Breast	1	1
Epistaxis	47	50
Gastrointestinal	63	59
Hematuria	10	10
Hemoptysis	6	6
Intraocular	3	3
No site identified	9	
Other	132	136
Subdural hematoma	1	1
Surgical incision site	8	7
Urethral	2	2
Vaginal	5	5
Vascular access site	9	9
Missing	1	1
Total	297	290

GUSTO indicates Global Use of Strategies to Open Occluded Coronary Arteries; TIMI, Thrombolysis In Myocardial Infarction.

Table S4. Adjusted associations of GUSTO and TIMI composite bleeding definitions with continuous PRU Distributions and the derived cut-points for low vs. high platelet reactivity restricted to patients aged <75 years

	Adjusted HR (95% CI)	P
GUSTO severe/life-threatening or moderate non-CABG bleeding		
Continuous day 5 PRU (per 10-unit decrease)	1.01 (0.95–1.06)	0.85
Dichotomous (<106) day 5 PRU (LPR vs. HPR)	0.61 (0.20–1.84)	0.38
GUSTO severe/life-threatening, moderate, or mild non-CABG bleeding		
Continuous day 5 PRU (per 10-unit decrease)	1.04 (1.03–1.06)	<0.001
Dichotomous (<75) day 5 PRU (LPR vs. HPR)	2.19 (1.61–2.98)	<0.001
TIMI major or minor non-CABG bleeding		
Continuous day 5 PRU (per 10-unit decrease)	1.02 (0.98–1.07)	0.35
Dichotomous (<46) day 5 PRU (LPR vs. HPR)	1.99 (0.81–4.90)	0.13
TIMI major, minor, or minimal non-CABG bleeding		
Continuous day 5 PRU (per 10-unit decrease)	1.04 (1.03–1.06)	<0.001
Dichotomous (<75) day 5 PRU (LPR vs. HPR)	2.21 (1.62–3.02)	<0.001

CABG, coronary artery bypass graft; CI, confidence interval; GUSTO, Global Use of Strategies to Open Occluded Coronary Arteries; HPR, high platelet reactivity; HR, hazard ratio; LPR, low platelet reactivity; PRU, P2Y₁₂ reaction unit; TIMI, Thrombolysis In Myocardial Infarction.