# International Expert Consensus on Switching Platelet P2Y<sub>12</sub> Receptor– Inhibiting Therapies

**ABSTRACT:** Dual antiplatelet therapy with aspirin and a P2Y<sub>12</sub> inhibitor is the treatment of choice for the prevention of atherothrombotic events in patients with acute coronary syndromes and for those undergoing percutaneous coronary interventions. The availability of different oral P2Y<sub>12</sub> inhibitors (clopidogrel, prasugrel, ticagrelor) has enabled physicians to contemplate switching among therapies because of specific clinical scenarios. The recent introduction of an intravenous P2Y<sub>12</sub> inhibitor (cangrelor) further adds to the multitude of modalities and settings in which switching therapies may occur. In clinical practice, it is not uncommon to switch  $P2Y_{12}$  inhibitor, and switching may be attributed to a variety of factors. However, concerns about the safety of switching between these agents have emerged. Practice guidelines have not fully elaborated on how to switch therapies, leaving clinicians with limited guidance on when and how to switch therapies when needed. This prompted the development of this expert consensus document by key leaders from North America and Europe with expertise in basic, translational, and clinical sciences in the field of antiplatelet therapy. This expert consensus provides an overview of the pharmacology of P2Y<sub>12</sub> inhibitors, different modalities and definitions of switching, and available literature and recommendations for switching between P2Y<sub>12</sub> inhibitors.

ual antiplatelet therapy with aspirin and a platelet P2Y<sub>12</sub> receptor antagonist (P2Y<sub>12</sub> inhibitor) is the treatment of choice for the prevention of atherothrombotic events in patients with acute coronary syndromes (ACS) and for those undergoing percutaneous coronary intervention (PCI).<sup>1,2</sup> Clopidogrel, prasugrel, and ticagrelor are the most commonly used oral platelet P2Y<sub>12</sub> inhibitors; the use of ticlopidine, the first available P2Y<sub>12</sub> inhibitor, has been largely abandoned.<sup>3</sup> Clopidogrel is the only oral P2Y<sub>12</sub> inhibitor indicated for the treatment of patients with stable coronary artery disease.<sup>1,2</sup> Although all 3 agents have an indication for use in ACS, current guidelines support the preferential use of prasugrel and ticagrelor over clopidogrel because of their superior net clinical benefits.<sup>1,2,4–6</sup> Nevertheless, clopidogrel remains widely prescribed.<sup>7,8</sup>

The availability of different oral P2Y<sub>12</sub> inhibitors has enabled physicians to contemplate switching among therapies because of specific clinical scenarios.<sup>9</sup> The recent introduction of an intravenous P2Y<sub>12</sub> inhibitor (ie, cangrelor) further adds to the multitude of modalities and settings in which switching therapies may occur.<sup>6</sup> A variety of factors may contribute to the decision to switch, including the clinical setting, patient characteristics, concomitant therapies, costs, social issues, development of side effects, medication adherence, and patient/physician preference.<sup>9</sup> Dominick J. Angiolillo, MD, PhD et al

The full author list is available on page 1969.

Correspondence to: Dominick J. Angiolillo, MD, PhD, University of Florida College of Medicine– Jacksonville, 655 West 8th Street, Jacksonville, FL 32209. E-mail dominick.angiolillo@jax.ufl.edu

Key Words: drug interactions ■ hemorrhage ■ pharmacodynamics ■ platelet aggregation inhibitors ■ P2Y<sub>12</sub> receptor inhibitors ■ thrombosis

© 2017 American Heart Association, Inc. Therefore, it is not uncommon to change  $P2Y_{12}$  inhibitor. However, concerns about the safety of switching between these agents have emerged.

At present, data from large-scale clinical studies to guide the optimal approach to switching  $P2Y_{12}$  inhibitors are limited, and most data are derived from pharmacodynamic studies. In turn, practice guidelines have not fully elaborated on how to switch therapies, leaving clinicians with limited guidance on when and how to switch therapies when needed, which prompted the development of this expert consensus document. Key leaders from North America and Europe with expertise in basic, translational, and clinical sciences in the field of antiplatelet therapy who have contributed to the scientific literature of switching antiplatelet therapies were identified by the document chairs (D.J.A. and M.J.P.). All invited experts agreed to partake in the development of this document and endorse the recommendations provided. This was an academic collaboration between the identified experts and was free from any type of industry support. This expert consensus provides an overview of the pharmacology of P2Y<sub>12</sub> inhibitors, different modalities and definitions of switching, available literature, and recommendations for switching between P2Y<sub>12</sub> inhibitors.

### PHARMACOLOGICAL PROPERTIES

Concerns surrounding the safety of switching between P2Y<sub>12</sub> inhibitors have emerged because of the potential

for drug-drug interactions (DDIs). A DDI is defined as a modification of the effect of a drug when administered with another drug. In particular, because of a DDI, the effects of a P2Y<sub>12</sub> inhibitor can be decreased, leading to inadequate platelet inhibition and increasing the risk for thrombotic complications; alternatively, there may be a potential for overdosing as a result of an overlap in drug therapy that could lead to excessive platelet inhibition and predispose to bleeding complications. Although to date no studies have shown a clinical impact of DDIs occurring as a result of switching, there is robust evidence associating different levels of platelet reactivity with adverse clinical outcomes.<sup>10,11</sup> The potential for DDIs when switching P2Y<sub>12</sub> inhibitors rests on differences in their pharmacological properties. Key pharmacological properties to consider include drug half-life, the site and mechanism of P2Y<sub>12</sub> receptor binding, and the speeds of onset and offset of pharmacodynamic effects (Table 1).<sup>3,6,9</sup>

Clopidogrel, a second-generation thienopyridine, is a prodrug that is largely (up to 85%) hydrolyzed into an inactive metabolite by human carboxylesterase-1 after intestinal absorption.<sup>12</sup> The remaining prodrug ( $\approx$ 15%) requires a 2-step oxidation process with multiple hepatic cytochrome P-450 (CYP) isoenzymes, mainly CYP2C19, to generate the active thiol metabolite that irreversibly blocks the ADP-binding site on the P2Y<sub>12</sub> receptor (Figure 1). Prasugrel is a third-generation thienopyridine and is also a prodrug. However, the generation of the active metabolite of prasugrel is more efficient compared with clopidogrel because ultrarapid hydrolysis by human

	Clopidogrel	Prasugrel	Ticagrelor	Cangrelor
Receptor blockade	Irreversible	Irreversible	Reversible	Reversible
Prodrug	Yes	Yes	No	No
Half-life of parent drug	≈6 h	<5 min	6–12 h	3–6 min
Half-life of active metabolite	30 mins	Distribution half-life, 30–60 mins	8–12 h	NA
		Elimination half-life, 2–15 h		-
Binding site	ADP-binding site	ADP-binding site	Allosteric binding site	Undetermined*
Administration route	Oral	Oral	Oral	Intravenous
Frequency	Once daily	Once daily	Twice daily	Bolus plus infusion
Onset of action†	2–8 h	30 min–4 h	30 min–4 h	≈2 min
Offset of action	5–10 d	7–10 d	3–5 d	60 min
CYP drug interaction‡	CYP2C19	No	СҮРЗА	No
Approved settings	ACS (invasive and noninvasively managed), stable CAD, PCI, PAD, and ischemic stroke	ACS undergoing PCI	ACS (invasive or noninvasively managed) or history of MI	PCI in patients with or without ACS

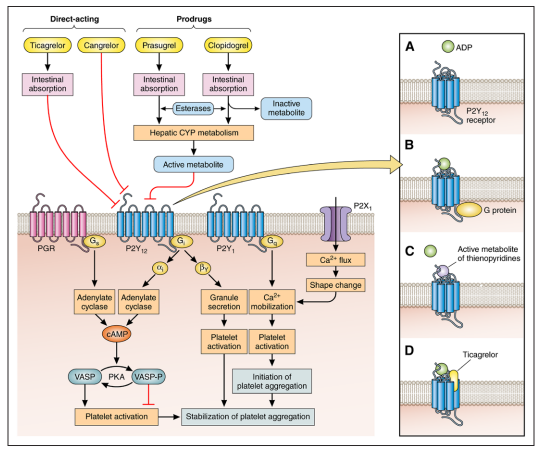
 Table 1. Pharmacological Properties of P2Y<sub>12</sub> Receptor Inhibitors

ACS indicates acute coronary syndrome; CAD, coronary artery disease; CYP, cytochrome P450; MI, myocardial infarction; PAD, peripheral arterial disease; and PCI, percutaneous coronary intervention.

\*The binding site of cangrelor at the P2Y<sub>12</sub> receptor level is not clearly defined; nevertheless, cangrelor is associated with high levels of receptor occupancy, preventing ADP signaling.

t Indicates times after loading dose and bolus administration for oral and intravenous agents, respectively. Times for oral agents refer to clinically stable subjects and may be prolonged in patients with ST-segment–elevation myocardial infarction or treated with opioids.

*‡Indicates clinically significant drug interactions.* 



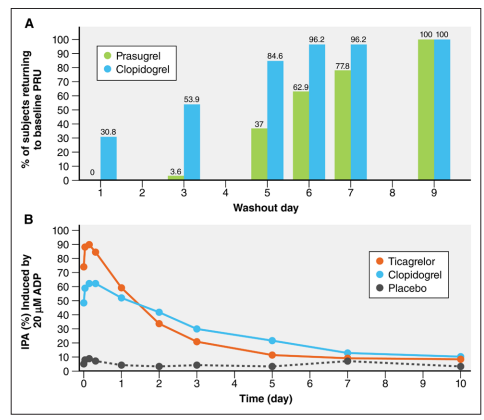
#### Figure 1. Mechanisms of action and binding properties of P2Y<sub>12</sub> inhibitors.

**Left**, Mechanism of action. Activation of the P2Y<sub>12</sub> receptor inhibits adenylyl cyclase, causing a decrease in cAMP and phosphorylated (P) vasodilator-stimulated phosphoprotein (VASP) levels, and activation of P2Y<sub>12</sub> causes an increase in intracellular Ca<sup>2+</sup> levels. These changes promote platelet aggregation by altering the ligand-binding properties of the glycoprotein IIb/IIIa receptor. Inhibition of the P2Y<sub>12</sub> receptor therefore suppresses platelet activation. Clopidogrel and prasugrel are oral prodrugs requiring hepatic metabolism to generate an active metabolite that irreversibly inhibits the P2Y<sub>12</sub> receptor. Ticagrelor is a direct-acting (no metabolism required) oral agent that reversibly inhibits the P2Y<sub>12</sub> receptor. Cangrelor is a direct-acting intravenous agent that reversibly inhibits the P2Y<sub>12</sub> receptor. **Right**, Binding properties. **A**, ADP binds to the P2Y<sub>12</sub> receptor, which (**B**) leads to a conformational change of the receptor and to G-protein activation. **C**, The active metabolite of thienopyridines occupies the ADP-binding site on the P2Y<sub>12</sub> receptor. Binding is irreversible, which renders the receptor nonfunctional for the life of the platelet. **D**, Ticagrelor binds reversibly to the P2Y<sub>12</sub> receptor; and PKA, protein kinase A. Adapted from Rollini et al<sup>9</sup> with permission. Copyright ©2016, Mcmillan Publishers Ltd.

carboxylesterase-2 forms an intermediate metabolite, which subsequently requires only a single-step hepatic CYP oxidation to generate its active metabolite. The active metabolite of prasugrel also irreversibly blocks the ADP-binding site on the P2Y<sub>12</sub> receptor.<sup>12</sup> Although the active metabolite of prasugrel is equipotent to that derived from clopidogrel, its plasma concentration is higher, which translates into more prompt, potent, and predictable platelet inhibitory effects compared with clopidogrel.<sup>12</sup> The active metabolite of clopidogrel is unstable, has a very short half-life ( $\approx$ 30 minutes), and is rapidly eliminated from the circulation if it does not bind to the P2Y<sub>12</sub> receptor. The active metabolite of prasugrel is more stable, but plasma levels fall rapidly as a result of distribution to extravascular compartments (distribution

half-life, 30–60 minutes), after which levels may be insufficient to achieve effective levels of P2Y<sub>12</sub> blockade.<sup>12</sup> These low levels of active metabolite are detectable in the circulation for an extended time compared with clopidogrel as a result of the much longer elimination half-life (2–15 hours).<sup>12</sup> Given the irreversible binding of the active metabolites, recovery time after treatment discontinuation approximates the life span of platelets. Although subject to variability, this is longer after prasugrel (7–10 days) compared with clopidogrel (5–7 days) discontinuation because of the enhanced level of platelet inhibition achieved (Figure 2A).<sup>13,14</sup>

Ticagrelor is an oral cyclopentyl-triazolopyrimidine that reversibly binds the P2Y<sub>12</sub> receptor.<sup>15</sup> It is a direct-acting agent and does not require hepatic metabolism



#### Figure 2. Offset of antiplatelet effects of oral P2Y<sub>12</sub> inhibitors.

**A**, Cumulative proportion of patients returning to baseline reactivity after thienopyridine discontinuation: the RECOVERY trial (Recovery of Platelet Function Following Discontinuation of Prasugrel or Clopidogrel Maintenance Dosing in Aspirin-Treated Subjects With Stable Coronary Disease). Baseline platelet reactivity defined as within 60 P2Y<sub>12</sub> reaction units (PRUs) of the reactivity measured before study drug exposure. Adapted from Price et al<sup>14</sup> with permission. Copyright ©2012, American College of Cardiology. **B**, Offset of inhibition of platelet aggregation (IPA) on ticagrelor, clopidogrel, and placebo: the ONSET/OFFSET study (A Study of the Onset and Offset of Antiplatelet Effects Comparing Ticagrelor, Clopidogrel, and Placebo With Aspirin). IPA after 20 µmol/L ADP (final extent) measured after last ticagrelor, clopidogrel, and placebo maintenance dose (day 0) and followed up for 10 days. Adapted from Gurbel al<sup>16</sup> with permission. Copyright ©2009, American Heart Association, Inc.

to exert its effect. However, ≈30% of the antiplatelet effect of ticagrelor derives from an active metabolite (AR-C124910XX) generated through CYP3A4/5 enzymes. This active metabolite has pharmacological properties similar to those of the parent compound.<sup>15</sup> Ticagrelor requires twice-daily dosing because of its reversible receptor binding and half-life of 6 to 12 hours. Ticagrelor reversibly binds to a distinct site on the P2Y<sub>12</sub> receptor and acts through a noncompetitive, allosteric mechanism to prevent G-protein-mediated signal transduction after ADP binding.<sup>15</sup> The pharmacodynamic effects of ticagrelor are more prompt, potent, and predictable compared with those of clopidogrel. However, because of its reversible binding and relatively short half-life, ticagrelor has a faster offset of antiplatelet effect (3-5 days) compared with thienopyridines<sup>16</sup> (Figure 2B).

Cangrelor is an intravenous ATP analog that directly and reversibly inhibits ADP binding to the P2Y<sub>12</sub> receptor in a dose-dependent manner, achieving immediate potent platelet inhibition after a bolus dose.<sup>6,17</sup> Although its binding site at the P2Y<sub>12</sub> receptor level is not clearly defined, cangrelor is associated with high levels of receptor occupancy and prevents ADP binding. Cangrelor is promptly inactivated through dephosphorylation by ectonucleotidase and has a very short plasma half-life (3–6 minutes). Therefore, recovery of platelet function is rapid ( $\approx$ 60 minutes) after discontinuation of cangrelor infusion.<sup>6,17</sup>

# SWITCHING MODALITIES AND DEFINITIONS

This expert consensus provides uniform definitions to describe the various modalities of switching of  $P2Y_{12}$  inhibitors. In particular, switching can occur between the oral agents and between the oral agents and an intravenous agent. Moreover, the timing of switching with respect to the index event that led to the initiation of  $P2Y_{12}$  inhibitor therapy may also vary. Ultimately, switching may occur between different classes of  $P2Y_{12}$  inhibitors, which may have potential implications for the occurrence of DDI between the 2 overlapped

Type of Pharmacodynamic Switch	Type of Drug Class Switch	Potential for DDI
Oral*		
Escalation		
Clopidogrel to prasugrel	Intraclass	No
Clopidogrel to ticagrelor	Interclass	No
De-escalation	·	
Prasugrel to clopidogrel	Intraclass	No
Ticagrelor to clopidogrel	Interclass	Yes
Change		
Prasugrel to ticagrelor	Interclass	No
Ticagrelor to prasugrel	Interclass	Yes
Intravenous	• •	
Bridge		
Clopidogrel to cangrelor	Interclass	No
Prasugrel to cangrelor	Interclass	No
Ticagrelor to cangrelor	Interclass	No
Transition		
Cangrelor to clopidogrel	Interclass	Yes
Cangrelor to prasugrel	Interclass	Yes
Cangrelor to ticagrelor	Interclass	No

# Table 2.Modalities of Switching Between P2YReceptor Inhibitors and Potential for DDI

DDI indicates drug-drug interaction.

\*Switching between oral agents may be classified according to relationship from the index event a defined as acute (<24 hours), early (1–≤30 days), late (>30 days–1 year), and very late (>1 year).

agents. Switching modalities between  $P2Y_{12}$  inhibitors and their potential for DDI are summarized in Table 2.

### Switching Between Oral P2Y<sub>12</sub> Inhibitors

Prasugrel and ticagrelor are characterized by enhanced pharmacodynamic effects compared with clopidogrel.<sup>3,6,12–16</sup> Therefore, switching between oral P2Y<sub>12</sub> inhibitors may result in a variation from a less intensive to a more intensive agent (ie, clopidogrel to prasugrel or ticagrelor) or vice versa from a more intensive to a less intensive agent (ie, prasugrel or ticagrelor to clopidogrel). These modalities of switching are defined as escalation and de-escalation, respectively. Although studies comparing the pharmacodynamic effects of prasugrel versus ticagrelor have yielded some inconsistent findings, the overall levels of P2Y12 inhibition are markedly reduced and not that dissimilar between these agents.<sup>18</sup> Switching between prasugrel and ticagrelor is referred to as change. Such terminology (escalation, de-escalation, and change) should be considered only when referring to the pharmacodynamic effects associated with switching and should not imply any clinical correlate (efficacy or safety).

Switching may be also classified according to the  $P2Y_{12}$  inhibitor class. Two different classes of oral  $P2Y_{12}$  inhibitors are available for clinical use: thienopyridine

(ie, clopidogrel or prasugrel) and cyclopentyl-triazolopyrimidine (ie, ticagrelor).<sup>3,6</sup> In some circumstances, an interclass switch (ie, between agents from 2 different classes) may lead to a DDI, which is unlikely to occur from an intraclass switch (ie, between 2 different agents of the same class). Overall, escalation of therapy has not been associated with DDI, regardless of class. However, there is a potential for a DDI with de-escalation therapy, particularly when switching from ticagrelor to clopidogrel.<sup>9,19</sup> A DDI, with an increase in platelet reactivity, has been suggested when switching from ticagrelor to prasugrel but not from prasugrel to ticagrelor.<sup>20,21</sup>

Switching may occur at different times from the index event that led to initiation of oral P2Y<sub>12</sub>-inhibiting treatment. A main concern with switching oral P2Y<sub>12</sub>inhibiting therapy is that if this is associated with inadequate platelet inhibition, it may lead to stent thrombosis.<sup>10,11</sup> Because thrombotic risk is highest in the early weeks after an ACS or PCI, the timing of switching from the index event may have therapeutic implications. Definitions from the Academic Research Consortium have been provided to define stent thrombosis according to timing of occurrence.<sup>22</sup> In line with the therapeutic implications associated with switching according to the time from PCI, this expert consensus believes that incorporating well-known and established definitions would be practical. Accordingly, the timing of switching with respect to the duration since the initiating event may be defined as acute (<24 hours), early (1–30 days), late (>30 days-1 year), or very late (>1 year).

# Switching to and From an Intravenous P2Y<sub>12</sub> Inhibitor

Cangrelor, the only available intravenous P2Y<sub>12</sub> inhibitor, provides more prompt and greater  $P2Y_{12}$  inhibition than any of the oral agents.<sup>17,23–27</sup> Switching may occur from an oral agent to cangrelor or vice versa. Patients are typically switched from an oral P2Y<sub>12</sub> inhibitor to cangrelor while awaiting cardiac or noncardiac surgery. This modality of switching is defined as bridging. Patients are typically switched from cangrelor to an oral P2Y<sub>12</sub> inhibitor in the setting of PCI when cangrelor is used to achieve immediate potent platelet inhibition during the peri-PCI period. Because of the need to continue P2Y<sub>12</sub> inhibition with an oral agent after discontinuation of cangrelor, this type of switching is defined as transition. Because cangrelor is of a different class from all oral P2Y<sub>12</sub> inhibitors, all switches involving cangrelor are by definition interclass. Bridging from oral to intravenous P2Y<sub>12</sub>-inhibiting therapy with cangrelor is associated with sustained P2Y<sub>12</sub> inhibitory effects and does not lead to a DDI.<sup>28</sup> However, transitioning from cangrelor to a thienopyridine (clopidogrel and prasugrel), but not ticagrelor, can be associated with a DDI.<sup>26–31</sup>

### SWITCHING BETWEEN ORAL P2Y<sub>12</sub> INHIBITORS

In this section, a summary of the available data from clinical trials, registries, and pharmacodynamic studies on escalation, de-escalation, and change in oral  $P2Y_{12}$  inhibitors is provided.

### **Escalation (Switching From Clopidogrel** to Prasugrel or Ticagrelor)

Escalating from clopidogrel to prasugrel or ticagrelor therapy commonly occurs in patients presenting with an ACS, above all those undergoing PCI, who may have been pretreated with clopidogrel at the time of clinical presentation. This is particularly frequent among patients who get transferred to a PCI-capable center. Occurrence of an ACS while on clopidogrel is also a reason for escalating therapy. To date, most data on escalation therapy derive from subgroup analyses of large clinical trials, registries. and pharmacodynamic studies.

The TRITON-TIMI 38 trial (Therapeutic Outcomes by Optimizing Platelet Inhibition With Prasugrel-Thrombolysis in Myocardial Infarction 38) demonstrated the superiority of prasugrel over clopidogrel in reducing ischemic events, albeit at the expense of increased bleeding, including fatal bleeding, in patients with ACS undergoing PCI. However, this trial cannot address the impact of switching from clopidogrel to prasugrel because patients with previous exposure to a P2Y<sub>12</sub> inhibitor were excluded.<sup>5</sup> On the contrary, the ACCOAST trial (Comparison of Prasugrel at the Time of Percutaneous Coronary Intervention or as Pretreatment at the Time of Diagnosis in Patients With Non-ST Elevation Myocardial Infarction), which tested the effects of administering prasugrel 30 mg at the time of diagnosis plus 30 mg after coronary angiography versus administering 60 mg after coronary angiography if PCI was indicated in patients with non-ST-segment-elevation myocardial infarction, allowed patients receiving a 75-mg maintenance dose (MD) of clopidogrel at the time of randomization to be enrolled.<sup>32</sup> However, pretreatment with prasugrel increased major bleeding complications without any ischemic benefit, with consistent findings regardless of clopidogrel pretreatment. The TRILOGY-ACS trial (Targeted Platelet Inhibition to Clarify the Optimal Strategy to Medically Manage ACS) assessed the impact of long-term use of prasugrel compared with clopidogrel in patients with non–ST-segment–elevation ACS selected for medical management without revascularization. Prasugrel was initiated with an MD, without a loading dose (LD), in ≈95% of the population; ≈70% of patients randomized to prasugrel had received clopidogrel administered as an LD. Although there were no differences in major bleeding complications between treatment groups, these results need to be interpreted

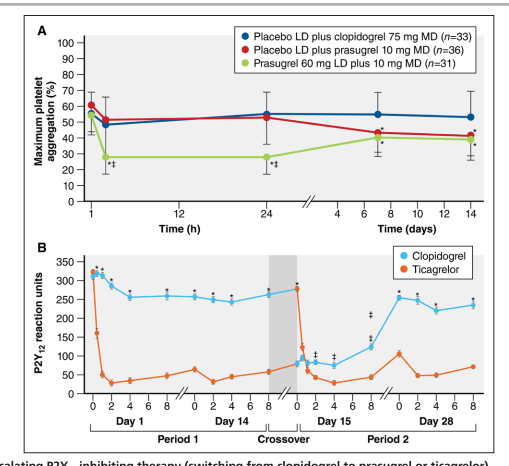
with caution because the trial did not reach its primary efficacy end point.  $^{\mbox{\tiny 33}}$ 

The PLATO trial (Study of Platelet Inhibition and Patient Outcomes) demonstrated the superiority of ticagrelor over clopidogrel in reducing ischemic events without an increase in the rate of overall major bleeding but with an increase in non-coronary artery bypass graft surgery-related bleeding across the spectrum of patients with ACS regardless of the planned management strategy (invasive or noninvasive).<sup>4</sup> Approximately 50% of patients randomized to ticagrelor were previously treated with clopidogrel, and the efficacy and safety of ticagrelor 180-mg LD followed by an MD of 90 mg twice daily were consistent regardless of previous clopidogrel exposure.<sup>4</sup> In the ATLANTIC trial (Administration of Ticagrelor in the Cath Laboratory or in the Ambulance for New ST Elevation Myocardial Infarction to Open the Coronary Artery), which showed that prehospital administration of ticagrelor in patients with acute ST-segment-elevation myocardial infarction appeared to be safe but did not improve pre-PCI coronary reperfusion, patients who were on clopidogrel at the time of presentation were excluded.<sup>34</sup> The PEGASUS trial (Prevention of Cardiovascular Events in Patients With Prior Heart Attack Using Ticagrelor Compared to Placebo on a Background of Aspirin) evaluated the safety and efficacy of a long-term MD of ticagrelor 60 or 90 mg twice daily, initiated without an LD, compared with placebo in patients with a myocardial infarction in the previous 1 to 3 years. Treatment with ticagrelor significantly reduced ischemic events, albeit at the expense of increased major bleeding.<sup>35</sup> Approximately one third of patients were on a P2Y<sub>12</sub> inhibitor (mostly clopidogrel) at the time of randomization.

A number of registries have evaluated escalating from clopidogrel to prasugrel or ticagrelor, showing a prevalence that varied from 5% to 50%, depending on the clinical setting and the period of observation (in-hospital versus after discharge; Table I in the onlineonly Data Supplement).<sup>36–47</sup> The reasons for switching included primarily clinical factors such as ST-segmentelevation myocardial infarction presentation, in-hospital reinfarction, high-risk angiographic characteristics, younger age, higher body weight, sex, and socioeconomic factors. In the majority of cases, the switch occurred in the catheterization laboratory at the time of or immediately after PCI. Although registries did not identify any major safety concerns associated with switching, these findings should be interpreted with caution because the studies were not designed or powered to assess clinical outcomes.

Many studies have been specifically conducted to provide insights into levels of platelet reactivity associated with switching from clopidogrel to prasugrel or ticagrelor (Figure 3). In the SWAP study (Switching Antiplatelet), conducted in patients receiving mainte-

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# Figure 3. Escalating P2Y<sub>12</sub> inhibiting therapy (switching from clopidogrel to prasugrel or ticagrelor). **A**, Pharmacodynamic profile of switching from clopidogrel to prasugrel therapy: the SWAP study (Switching Antiplatelet). Time course of platelet inhibition as measured with maximum platelet aggregation in response to 20 $\mu$ mol/L ADP in patients with an acute coronary syndrome whose therapy was switched from clopidogrel to prasugrel. Patients were randomly assigned to 1 of the 3 study groups. \**P*<0.0001 vs results with 75-mg maintenance dose (MD) of clopidogrel. ‡*P*<0.0001 vs results with

10-mg MD of prasugel. **B**, Pharmacodynamic profile of switching between clopidogrel and ticagrelor therapy: results from the RESPOND study (Response to Ticagrelor in Clopidogrel Nonresponders and Responders and the Effect of Switching Therapies). P2Y<sub>12</sub> reaction units in clopidogrel-nonresponsive patients before and after crossover. Patients treated with ticagrelor in period 1 received a 600-mg clopidogrel loading dose (LD) followed by 75-mg daily maintenance therapy in period 2; patients treated with clopidogrel in period 1 received a 180-mg ticagrelor LD followed by 90-mg twice-daily maintenance therapy in period 2. \**P*<0.0001. ‡*P*<0.05. Adapted from Rollini et al<sup>9</sup> with permission. Copyright ©2016, Mcmillan Publishers Ltd.

nance clopidogrel therapy after an ACS event, escalation from clopidogrel to prasugrel was associated with further reduction in platelet reactivity within 2 hours with the administration of a 60-mg prasugrel LD and by 1 week with 10-mg prasugrel as an MD (Figure 3A).<sup>48</sup> In the RESPOND study (Response to Ticagrelor in Clopidogrel Nonresponders and Responders and the Effect of Switching Therapies), conducted among patients with stable coronary artery disease, ticagrelor therapy (180-mg LD followed by MD) overcame nonresponsiveness to clopidogrel, and its antiplatelet effect was the same in clopidogrel responders and nonresponders (Figure 3B).<sup>49</sup> Many other studies exploring the pharmacodynamic profiles of switching from clopidogrel to prasugrel or ticagrelor have been conducted (Tables II and III in the online-only Data Supplement).<sup>18,48–66</sup> All studies have consistently shown enhanced platelet inhibition when escalating from clopidogrel to prasugrel or ticagrelor, regardless of clinical setting, as well as a reduction in rates of high on-treatment platelet reactivity (HPR),18,48-70 a well-defined marker of risk of ischemic recurrences, including stent thrombosis.<sup>10,11</sup> These effects are achieved more promptly after administration of an LD compared with an MD regimen. These pharmacodynamic studies did not suggest any type of DDI or concerns of overdosing. This may be attributed to the fact that in patients treated with clopidogrel, even after an LD, a substantial number of P2Y<sub>12</sub> receptors remain uninhibited, allowing additional blockade by the administration of an LD of prasugrel or ticagrelor. The degree of P2Y<sub>12</sub> receptor blockade after prasugrel or ticagrelor administration is similar regardless of previous exposure to clopidogrel.

# De-escalation (Switching From Prasugrel or Ticagrelor to Clopidogrel)

Despite the evidence for the sustained efficacy and safety of prasugrel and ticagrelor with long-term treatment, many physicians limit treatment duration with these agents to the early weeks or months after the index event.<sup>36–41</sup> Reduced costs associated with a generic formulation of clopidogrel and concerns about increased risk of bleeding with prasugrel and ticagrelor remain the most important reasons for de-escalation. Nonbleeding side effects such as dyspnea also represent a potential reason for interrupting ticagrelor therapy.<sup>4,35,71,72</sup>

Overall, registry data indicate that the prevalence of in-hospital de-escalation ranges from 5% to 14% (Table I in the online-only Data Supplement).<sup>36–41</sup> These patients are less likely to be privately insured and have risk factors associated with increased bleeding risk such as older age, lower body weight, previous transient ischemic attack/stroke, in-hospital treatment with coronary artery bypass graft surgery, atrial fibrillation/flutter, and use of oral anticoagulants (OACs).<sup>36–41</sup> Switching between  $P2Y_{12}$  inhibitors after hospital discharge occurs in 5% to 8% of patients, with most cases represented by de-escalation.<sup>45</sup> The SCOPE registry (Switching From Clopidogrel to New Oral Antiplatelet Agents During Percutaneous Coronary Intervention) showed that deescalation of P2Y<sub>12</sub> inhibitors early after the index event in patients with ACS was associated with an increased risk of recurrent ischemic events with no differences in bleeding.<sup>47</sup> These findings are likely attributed to the increase in platelet reactivity and HPR rates, with patients being particularly vulnerable if de-escalation occurs too soon after the acute event.

Recently, randomized trials of de-escalation have been reported (Table I in the online-only Data Supplement).73,74 The randomized TOPIC trial (Timing of Optimal Platelet Inhibition After Acute Coronary Syndrome) showed that in patients who have been event free for the first month after an ACS on a combination of aspirin plus a new-generation P2Y<sub>12</sub> inhibitor, de-escalation to aspirin plus clopidogrel was associated with reduced bleeding complications, mostly minor.<sup>73</sup> Although this study did not show any differences in ischemic events between groups, play of chance cannot be ruled out given the limited sample size of the trial. The TROP-ICAL-ACS trial (Testing Responsiveness to Platelet Inhibition on Chronic Antiplatelet Treatment for ACS) randomized patients with ACS undergoing PCI to either standard treatment with prasugrel for 12 months or a de-escalation regimen (1 week of prasugrel followed by 1 week of clopidogrel and platelet function testing-guided maintenance therapy with clopidogrel or prasugrel from day 14 after hospital discharge).74 The trial showed that a strategy of guided de-escalation of antiplatelet treatment was noninferior to standard treatment with prasugrel at 1 year in terms of net clinical benefit. The strategy did not show any increase in ischemic events, although there was a numeric but not statistically significant reduction in bleeding. The moderate impact on bleeding risk reduction could be explained by the considerably high percentage of patients (40%) who required escalation back to prasugrel therapy because of developing HPR after de-escalation. Thus far, TROPICAL-ACS is the only randomized trial using results of platelet function testing to adjust antiplatelet therapy (escalation or de-escalation) to meet its primary end point.53,75-77 There are limited data assessing the clinical impact of escalation and de-escalation of antiplatelet therapy on the basis of the results of genetic testing, which is currently being evaluated in several randomized trials, including the use of rapid genetic testing.78

Overall, there is a paucity of studies assessing the pharmacodynamic effects associated with de-escalation to clopidogrel therapy that have consistently shown an increase in platelet reactivity and HPR rates, with some reporting lower bleeding events (Table IV in the onlineonly Data Supplement).<sup>19,49,51,52,56,79,80</sup> However, these findings, as well as the absence of increased thrombotic events despite a higher rate of patients developing HPR, should be interpreted with caution because none of these studies were powered for clinical outcomes. It is important to note that although switching from prasugrel or ticagrelor to clopidogrel is intuitively associated with an increase in platelet reactivity and HPR rates, the different speed of offset of the drugs may have important therapeutic implications, particularly with regard to the timing of clopidogrel administration and whether it should be given as an LD.<sup>9,19</sup> The rationale for switching should further influence whether an LD should be given, especially if there are concerns for bleeding.

# Change (Switching Between Prasugrel and Ticagrelor)

To date, there is limited information on switching between the newer-generation P2Y<sub>12</sub> inhibitors prasugrel and ticagrelor. The few available registry data indicate that the rate of switching between these agents ranges from 2% to 4% (Table I in the online-only Data Supplement).<sup>36–39</sup> Although ticagrelor can be administered in patients with ACS upstream before the coronary anatomy is known, physicians might consider switching to prasugrel because of its once-daily administration, which may improve adherence. Another reason to consider switching from ticagrelor to prasugrel is ticagrelor-associated dyspnea. Data from real-world clinical practice show that some patients may be treated with prasugrel despite having a relative or absolute contraindication but are candidates for ticagrelor therapy and may therefore switch treatment. These include patients with ACS who are pretreated with prasugrel before their coronary anatomy is defined but do not undergo PCI and those who have a previous cerebrovascular event.<sup>36,39,41</sup>

There are limited studies on the pharmacodynamic effects associated with change between newer P2Y<sub>12</sub> inhibitors.<sup>20,21</sup> The SWAP-2 study investigated the pharmacodynamic effects of switching from ticagrelor to prasugrel. In this study, patients were switched to prasugrel (with or without a 60-mg LD) 12 hours after the last MD of ticagrelor.<sup>20</sup> Platelet reactivity was higher in patients treated with prasugrel compared with patients treated with ticagrelor at 7 days, not meeting the noninferiority primary end point. Moreover, at 24 hours and even more so at 48 hours, platelet reactivity increased in patients switched to prasugrel compared with preswitch levels, and the use of an LD of prasugrel appeared to be essential to mitigate the increase in platelet reactivity after switching (Figure 4A).<sup>20</sup> The mechanisms for these observations remain unknown but might be the result of prolonged binding of ticagrelor and its major metabolite to the P2Y<sub>12</sub> receptor after plasma levels have fallen, which may potentially impede the active metabolites of thienopyridines to access their binding site. These changes may also explain why, despite being a reversible agent with an 8- to 12 hour half-life, ticagrelor has effects that may persist for several days after drug discontinuation.<sup>16</sup> For these reasons, it has been suggested that switching at a later time after MD (eg, after 24 hours) should limit increases in platelet reactivity by providing more time for P2Y<sub>12</sub> receptor blockade by ticagrelor to decline.

The SWAP-3 study investigated the pharmacodynamic effects of switching from prasugrel to ticagrelor.<sup>21</sup> The study showed that in patients who were on maintenance prasugrel therapy, changing to ticagrelor was associated with a transient reduction in platelet reactivity. These pharmacodynamic findings were observed as early as 2 hours after switching therapy, without any signs of DDI during the entire study time course and with no increase in HPR rates. Of note, these findings were observed when switching to ticagrelor with the 90-mg (not 60-mg) dosing regimen and occurred regardless of the use of an LD (Figure 4B).<sup>21</sup>

### SWITCHING BETWEEN INTRAVENOUS AND ORAL P2Y<sub>12</sub> INHIBITORS Bridging From Oral P2Y<sub>12</sub> Inhibitors to Cangrelor

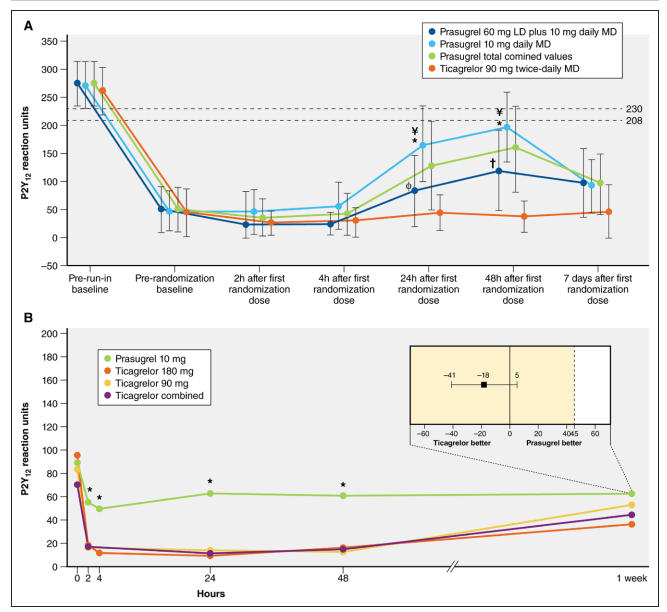
Although cangrelor is being used in real-world clinical practice as a bridging strategy, there are limited data to

support the safety and efficacy of this approach.<sup>81</sup> The BRIDGE trial (Maintenance of Platelet Inhibition With Cangrelor After Discontinuation of Thienopyridines in Patients Undergoing Surgery) showed that among patients who discontinue thienopyridine therapy before cardiac surgery, the use of cangrelor compared with placebo resulted in a higher rate of maintenance of platelet inhibition.<sup>28</sup> The dose of cangrelor used for bridging (0.75– $\mu$ g·kg<sup>-1</sup>·min<sup>-1</sup> infusion without a bolus) derives from a dose-finding study that identified levels of platelet inhibition similar to those achieved in patients with a good response to clopidogrel and is substantially lower than that used in PCI  $(30-\mu g/kg bolus)$ and  $4-\mu g \cdot k g^{-1} \cdot m i n^{-1}$  infusion). The pharmacodynamic results from the BRIDGE study do not suggest any type of DDI, likely because there are still unoccupied receptors in patients treated with oral P2Y<sub>12</sub> inhibitors that can be bound and inhibited by cangrelor. This is in line with in vitro and ex vivo investigations showing no interaction when cangrelor is administered on top of thienopyridines or ticagrelor and is associated with enhanced antiplatelet effects.24-27,30,82

### Transition From Cangrelor to Oral P2Y<sub>12</sub> Inhibitors

Cangrelor was approved for clinical use on the basis of the results of the CHAMPION PHOENIX trial (Cangrelor versus Standard Therapy to Achieve Optimal Management of Platelet Inhibition), which showed that cangrelor significantly reduced the rate of ischemic events, driven by a reduction in stent thrombosis and myocardial infarction, with no significant increase in severe bleeding in patients undergoing PCI.83 In patients treated with cangrelor, a clopidogrel LD was administered immediately after discontinuation of cangrelor infusion. This approach was used because pharmacodynamic studies with cangrelor demonstrated a rapid platelet inhibitory effect during cangrelor infusion and a rapid offset of action after treatment discontinuation.<sup>23</sup> The approach of administering clopidogrel after cangrelor was stopped was used across the cangrelor trial development program to avoid a potential DDI between cangrelor and clopidogrel, as described later. To date, no clinical outcomes study has investigated the safety and efficacy of cangrelor in patients subsequently treated with prasugrel or ticagrelor, although single-center observational data have been published.<sup>84,85</sup>

Given the different pharmacological properties of cangrelor and the oral P2Y<sub>12</sub> inhibitors, several studies have investigated the potential for DDI when these agents are concomitantly administered (Supplemental Table V in the online-only Data Supplement).<sup>23-31</sup> These potential DDIs are concerning because they can result in reduced platelet inhibition and subsequent lack of protection from thrombotic complications in the peri-



**Figure 4. Change between newer-generation oral P2Y**<sub>12</sub> **inhibitors (switching between prasugrel and ticagrelor). A**, Pharmacodynamic profile of switching from ticagrelor to prasugrel: the SWAP-2 study (Switching Antiplatelet 2). Time course of platelet inhibition as measured with P2Y<sub>12</sub> reaction units (PRUs) in patients with stable coronary artery disease. After the run-in phase with ticagrelor was completed, patients were randomly assigned to 1 of 3 regimens. \**P*<0.001 for change from prerandomization baseline for prasugrel maintenance dose (MD) at 24 and 48 hours.  $\phi P$ =0.002 for change from prerandomization baseline for prasugrel LD group at 24 hours. **†***P*<0.001 at 48 hours. **¥***P*<0.001 for the difference between the prasugrel MD group and the prasugrel LD group at 24 and 48 hours. Adapted from Angiolillo et al<sup>20</sup> with permission. Copyright ©2014, American College of Cardiology. **B**, Pharmacodynamic profile of switching from prasugrel to ticagrelor: the SWAP-3 study. Time course of platelet inhibition as measured with PRUs in patients with a recent acute coronary syndrome on maintenance prasugrel 10 mg. The box in the top right corner represents the primary end point: 1-week PRU absolute difference and 2-sided 95% confidence interval between ticagrelor combined and prasugrel 10 mg (tinted area indicates zone of noninferiority. Adapted from Franchi et al<sup>21</sup> with permission. Copyright ©2016, American College of Cardiology.

PCI period. In a study conducted in healthy volunteers, clopidogrel administration during cangrelor infusion was associated with an impaired antiplatelet effect of clopidogrel after cangrelor discontinuation.<sup>30</sup> This reflects the fact that the clopidogrel active metabolite,

like the prasugrel active metabolite, cannot bind to the P2Y<sub>12</sub> receptors if already largely occupied by cangrelor.<sup>86</sup> In turn, the plasma concentrations of the unbound thienopyridine active metabolites fall rapidly to subtherapeutic levels as a result of distribution to other

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compartments and systemic clearance. Therefore, after cangrelor infusion is stopped, when receptors become available for binding, most of the active metabolite of thienopyridines has already been eliminated from the circulation. In contradistinction, the antiplatelet effects of clopidogrel are not diminished when it is administered after cangrelor infusion because of the very fast offset of action of cangrelor and subsequent availability of P2Y<sub>12</sub> receptors for binding by clopidogrel active metabolite.<sup>23,30,31</sup> The transition from cangrelor to prasugrel is associated with transient recovery of platelet reactivity, in particular within 1 hour after cangrelor discontinuation.<sup>27</sup> However, it was observed that recovery of platelet function was attenuated when prasugrel was administered 30 minutes before the cangrelor infusion was stopped.<sup>27</sup> Conversely, administration of clopidogrel 30 minutes or 1 hour before cangrelor infusion discontinuation did not prevent recovery of platelet reactivity more effectively than administration at the end of the infusion.<sup>31</sup> Similar findings were observed when platelets were incubated with cangrelor before the addition of the active metabolites of either prasugrel or clopidogrel, when the ability of thienopyridines to inhibit platelet aggregation was strongly reduced.<sup>82</sup> However, the ExcelsiorLOAD2 study (Impact of Extent of Clopidogrel-Induced Platelet Inhibition During Elective Stent Implantation on Clinical Event Rate-Advanced Loading Strategies) showed that a 60-mg LD of prasugrel given at the start of a 2-hour infusion of cangrelor was associated with sufficient platelet inhibition after cangrelor, with only rare cases of HPR.87 These observations may be attributed to the relatively higher concentration and longer half-life of the active metabolite of prasugrel compared with that of clopidogrel.<sup>12</sup> However, whether similar findings would be observed with longer infusions of cangrelor (eg, up to 4 hours) is unknown.

Unlike that observed with thienopyridines, no interaction was shown for the transition from cangrelor to ticagrelor, allowing more versatile use of ticagrelor with respect to timing of administration in relation to the start of cangrelor therapy.<sup>26</sup> The presence of an interaction between thienopyridines, in particular clopidogrel, and cangrelor, but not between ticagrelor and cangrelor, is probably the result of the different half-lives of these drugs, as well as the different sites and types of binding to the P2Y<sub>12</sub> receptor.<sup>6,12,15,17</sup> Ticagrelor reversibly binds the P2Y<sub>12</sub> receptor at a site distinct from the ADP-binding site and has a half-life of 6 to 12 hours. Although it is unknown whether ticagrelor can bind with the P2Y<sub>12</sub> receptor during cangrelor infusion, its half-life (which exceeds that of the duration of cangrelor infusion) is such that drug is still systemically available to bind with the P2Y<sub>12</sub> receptor after discontinuation of cangrelor infusion. On the basis of these observations, ticagrelor can be administered before, during, or after cangrelor infusion.<sup>26</sup>

## P2Y<sub>12</sub> INHIBITORS: EXPERT CONSENSUS RECOMMENDATIONS ON SWITCHING

This expert consensus group developed recommendations on when and how to switch between P2Y<sub>12</sub> inhibitors, taking into consideration the pharmacological profiles of oral and intravenous P2Y<sub>12</sub> inhibitors; data from clinical trials, registries, and pharmacodynamic studies; and the potential for thrombotic complications based on the time elapsed from the index event leading to initiation of P2Y<sub>12</sub>-inhibiting therapy. In line with the limited safety and efficacy data in this field, these recommendations are to be considered mostly consensus based rather than evidence based. In general, switching approaches that have shown to be associated with DDI should be avoided or minimized unless clinically necessary. The provided recommendations are to be considered as guidance for the practicing clinician, who may consider alternative approaches based on the clinical context of the patient. The considerations made here are proposed under the assumption that these patients are also, for the most part, treated with concomitant low-dose aspirin in line with guideline recommendations. The expert consensus recommendations on switching P2Y<sub>12</sub> inhibitors are described in detail in the following sections and summarized in Figures 5 and 6.

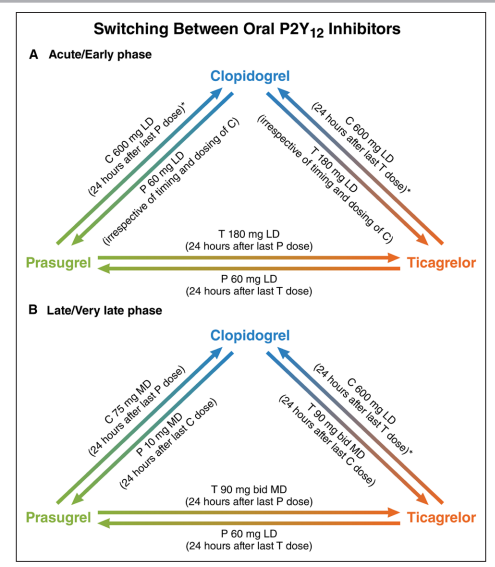
### Switching Between Oral P2Y<sub>12</sub> Inhibitors

# Escalation (Switching From Clopidogrel to Prasugrel or Ticagrelor)

Escalation from clopidogrel to prasugrel or ticagrelor in the early, particularly acute, phase of treatment should occur with the use of a 60- or 180-mg LD, respectively. Administration of an LD regimen may occur regardless of the timing of the last dose of clopidogrel. This should be followed by standard MD regimens (prasugrel 10 mg daily or ticagrelor 90 mg twice-daily). Beyond the early phase, it is reasonable to escalate with a 10-mg daily or 90-mg twice-daily MD regimen of prasugrel or ticagrelor, respectively, without an LD. It is also reasonable and practical for the patient to start the new MD regimen at the time of the next scheduled dose of P2Y<sub>12</sub>-inhibiting therapy (eg,  $\approx 24$  hours from last dose of clopidogrel). Similar considerations on timing of switching should apply for elderly or low-body-weight patients in whom a 5-mg MD of prasugrel is being used.

# De-Escalation (Switching From Prasugrel or Ticagrelor to Clopidogrel)

There was a lack of group consensus on the appropriate approach to de-escalate from prasugrel to clopidogrel in the acute/early phase (ie, with an MD or an LD) given the limited data on therapy de-escalation. The pro-

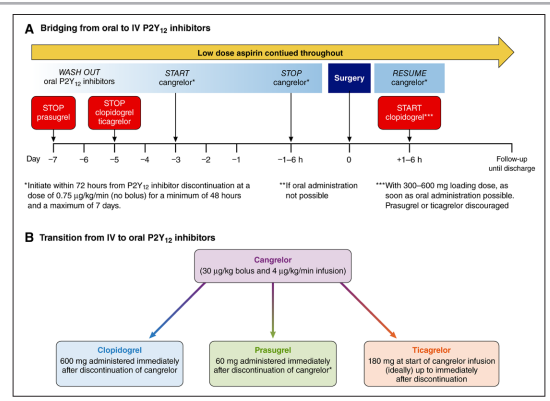


#### Figure 5. Consensus recommendations on switching between oral P2Y<sub>12</sub> inhibitors.

**A**, Switching between oral agents in the acute/early phase. In the acute/early phase ( $\leq$ 30 days from the index event), switching should occur with the administration of a loading dose (LD) in most cases, with the exception of patients who are deescalating therapy because of bleeding or bleeding concerns, in whom a maintenance dose (MD) of clopidogrel (C) should be considered. Timing of switching should be 24 hours after the last dose of a given drug, with the exception of when escalating to prasugrel (P) or ticagrelor (T), when the LD can be given regardless of the timing and dosing of the previous clopidogrel regimen. \*Consider de-escalation with clopidogrel 75-mg MD (24 hours after last prasugrel or ticagrelor dose) in patients with bleeding or bleeding concerns. **B**, Switching between oral agents in the late/very late phase. In the late/very late phase (>30 days from the index event), switching should occur with the administration of an MD 24 hours after the last dose of a given drug, with the exception of patients changing from ticagrelor to prasugrel therapy, for whom an LD should be considered. De-escalation from ticagrelor to clopidogrel should occur with administration of an LD 24 hours after the last dose of ticagrelor (but in patients in whom de-escalation occurs because of bleeding or bleeding concerns, an MD of clopidogrel should be considered). \*Consider de-escalation with clopidogrel 75-mg MD (24 hours after last prasugrel or ticagrelor dose) in patients with bleeding or bleeding or bleeding concerns, an MD of clopidogrel should be considered.

longed offset of prasugrel (7–10 days) has the advantage of allowing clopidogrel to reach its full antiplatelet effects during this time even if initiated with a 75-mg MD regimen. Moreover, because of the high receptor occupancy rates induced by prasugrel, it may be argued that administration of an LD of clopidogrel would not provide further pharmacodynamic effects. These pharmacological considerations suggest that de-escalation with an MD might be appropriate. Switching from prasugrel to clopidogrel with a 75-mg MD is also a reasonable option in patients in whom switching occurs as a result of a bleeding event or concerns about bleeding. Therefore, defining the reason for de-escalation may have an impact on the strategy (LD versus MD) used.

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#### Figure 6. Consensus recommendations on switching between oral and intravenous P2Y<sub>12</sub> inhibitors.

**A**, Bridging from oral to intravenous agents. For both cardiac and noncardiac surgery, if withdrawal of P2Y<sub>12</sub>-inhibiting therapy is needed, clopidogrel and ticagrelor should be discontinued for 5 days and prasugrel for 7 days. It is reasonable to start cangrelor bridging up to 3 to 4 days after prasugrel discontinuation and 2 to 3 days of clopidogrel and ticagrelor discontinuation. Platelet function testing may be considered to help guide timing of starting cangrelor infusion. After surgery, regardless of bridging strategy, clopidogrel should be resumed with a loading dose (LD) as soon as oral administration is possible and the risk of severe bleeding is acceptable (prasugrel and ticagrelor administration should be discouraged). If the use of oral P2Y<sub>12</sub>-inhibiting therapy is not possible, postsurgery bridging with an intravenous agent should be considered. **B**. Transition from intravenous to oral agents. An LD should always be used when transitioning from cangrelor to an oral agent. In the case of thienopyridines (clopidogrel or prasugrel), this should be administered immediately after discontinuation of cangrelor infusion. Ticagrelor can be administered before, during, or immediately after cangrelor infusion, although earlier administration (eg, at the time of percutaneous coronary intervention) should be considered. \*According to the package insert of the European Medical Agency, but not that of the US Food and Drug Administration, prasugrel may also be administered 30 minutes before infusion is stopped. Preliminary studies have shown that prasugrel given at the start of a 2-hour infusion of cangrelor results in sufficient platelet inhibition, but this strategy cannot be routinely recommended until more data are available.

However, it may also be argued that in the acute phase of treatment of patients with ACS, recovery of platelet function after discontinuation of prasugrel therapy may be shortened given their high platelet turnover rates, which may potentially not allow clopidogrel to reach its full platelet inhibitory effects before washout of prasugrel-mediated inhibition has been completed. Recovery of 37% and 63% of platelet function has been shown after 5 and 6 days, respectively, in patients with stable coronary artery disease.<sup>14</sup> Moreover, the onset of clopidogrel effect is variable, unpredictable, and often delayed. Therefore, in the early and, in particular, the acute phases of de-escalation, it may be also reasonable to administer a 600-mg LD of clopidogrel. This clopidogrel LD should be given at the time of the next scheduled dose of  $\text{P2Y}_{12}\text{-inhibiting}$  therapy (eg,  $\approx\!\!24$ hours from last dose of prasugrel) for practical reasons and because this would allow some offset of the effects of prasugrel and allow new uninhibited platelets to be released into circulation. Beyond the early phase or in more stabilized patients, the use of a 75-mg MD of clopidogrel (without an LD) at the time of the next scheduled dose (eg,  $\approx$ 24 hours from last dose of prasugrel) should be considered.

Because ticagrelor has a relatively fast offset of action, the use of a clopidogrel 600-mg LD should be considered when de-escalating from ticagrelor to avoid any significant gap in platelet inhibition, regardless of the timing of switching (ie, acute, early, or late), However, de-escalation to clopidogrel with an MD is a reasonable option, particularly in patients in whom switching occurs as a result of bleeding. Although the optimal timing of switching after the last dose of ticagrelor is unknown, waiting 24 hours after the last dose of ticagrelor should be considered because this not only exceeds the half-life of ticagrelor but also allows new platelets to be released into circulation and exposed to the active metabolite of clopidogrel, thus preventing a potential DDI. Furthermore, the level of platelet inhibition 24 hours after discontinuation of ticagrelor therapy is similar to the average level of inhibition provided by MD clopidogrel,<sup>14</sup> so a significant window of undertreatment is unlikely with this approach.

#### Change (Switching Between Prasugrel and Ticagrelor)

On the basis of pharmacodynamic data suggesting a potential DDI, a 60-mg LD of prasugrel should always be used when changing from ticagrelor to prasugrel, regardless of timing (early or late), and switching with a 10-mg MD should be avoided.<sup>20</sup> Waiting 24 hours after the last MD of ticagrelor to administer the 60-mg LD of prasugrel should be considered because this allows more time for ticagrelor and its metabolite to be eliminated and new platelets to enter into systemic circulation. Pharmacodynamic studies do not suggest DDI when changing from prasugrel to ticagrelor therapy.<sup>19</sup> Therefore, this change can be performed with a standard 90-mg twice-daily MD dose regimen, without the need for an LD, which should be started at the time of the next scheduled dose (eg, ≈24 hours from last dose of prasugrel), particularly in stabilized patients. However, the use of an LD administered 24 hours after the last dose of prasugrel can be considered when the change occurs in the acute phase of patients with ACS.

# Switching Between Intravenous and Oral P2Y<sub>12</sub> Inhibitors

#### Bridging From Oral P2Y<sub>12</sub> Inhibitors to Cangrelor

Because the effects of the oral agents persist with meaningful levels of P2Y<sub>12</sub> inhibition after drug discontinuation, it is reasonable to wait to start cangrelor bridging ( $0.75-\mu g\cdot kg^{-1}\cdot min^{-1}$  infusion without a bolus) for up to 3 to 4 days after prasugrel discontinuation and 2 to 3 days of clopidogrel and ticagrelor discontinuation to minimize the duration of infusion. Platelet function testing might also help time the initiation of cangrelor bridging in an efficient fashion. For example, cangrelor infusion can be started once the pharmacodynamic effect is close to the threshold of HPR. This may also have cost implications linked to hospitalization and the drug and potentially may minimize the risk of bleeding complications associated with prolonged treatment with parenteral therapies.

#### Transition From Cangrelor to Oral P2Y<sub>12</sub> Inhibitors

In patients undergoing PCI, cangrelor (30– $\mu$ g/kg bolus and 4– $\mu$ g·kg<sup>-1</sup>·min<sup>-1</sup> infusion) should be initiated before PCI and continued for ≥2 hours or for the duration of the procedure, whichever is longer; the infusion can be continued for up to 4 hours at the discretion of the physician. Infusions up to 4 hours might be considered particularly in patients treated with opiates such as morphine (terminal half-life varies from 1.5–4.5 hours) and possibly in patients undergoing primary PCI, which are settings known to reduce the pharmacodynamic onset of oral antiplatelet agents.<sup>88–91</sup> These observations are likely attributed to impaired gastrointestinal motility and drug absorption, which can be accentuated in patients undergoing primary PCI.<sup>92</sup>

In the transition from cangrelor to a thienopyridine, the thienopyridine should be administered immediately after discontinuation of cangrelor with an LD (clopidogrel 600 mg or prasugrel 60 mg) to avoid a potential DDI.93,94 According to the package insert of the European Medical Agency, but not that of the US Food and Drug Administration, prasugrel may also be administered 30 minutes before the infusion is stopped.93,94 Although preliminary studies have shown that prasugrel given at the time a 2-hour infusion of cangrelor is started results in sufficient platelet inhibition,<sup>87</sup> this strategy cannot be routinely recommended until more data are available. Although cangrelor is approved for use in patients who have not received an oral P2Y<sub>12</sub> inhibitor before the PCI procedure, for those patients who have been pretreated with a thienopyridine, if the pretreatment was shortly before the initiation of cangrelor or unknown, an LD at the end of the infusion should be considered.

The US Food and Drug Administration indicates that ticagrelor can be administered before, during, or immediately after cangrelor infusion,<sup>93,94</sup> whereas the European Medical Agency indicates that ticagrelor should be administered immediately after discontinuation of cangrelor infusion or up to 30 minutes before the end of the infusion, Ticagrelor should be administered as a 180-mg LD. This expert consensus recommends that earlier administration of ticagrelor (eg, at the time of PCI) should be considered over administration at the end of cangrelor infusion because it would minimize the potential gap in platelet inhibition during the transition phase.

#### SPECIAL CONSIDERATIONS

A number of settings represent clinical conundrums with regard to the management of antithrombotic therapy. Accordingly, there are a number of scenarios in which there may be a need to switch antiplatelet therapy but the modality to do this has not been studied. This expert consensus recognizes that there are some settings that may be unique and require specific recommendations.

 Patients undergoing cardiac and noncardiac surgery. Preoperative and postoperative management of antiplatelet therapy is described in detail

elsewhere.95 The decision to withdraw P2Y<sub>12</sub>inhibiting therapy should take into account the thrombotic and bleeding risks of the individual patient according to the specific surgery being performed and timing from PCI.<sup>96</sup> Similarly, the need for bridging should be individualized as described previously.95,96 For patients with ACS requiring coronary artery bypass surgery, unless recent PCI was conducted, P2Y<sub>12</sub>-inhibiting therapy should be withdrawn before surgery but restarted postoperatively if the bleeding risk is low. For both cardiac and noncardiac surgery, if withdrawal of P2Y12-inhibiting therapy is warranted, clopidogrel and ticagrelor should be discontinued for 5 days and prasugrel for 7 days. If bridging with cangrelor, it is reasonable to wait up to 3 to 4 days after prasugrel discontinuation and 2 to 3 days after clopidogrel and ticagrelor discontinuation to minimize duration of cangrelor infusion. After noncardiac surgery, regardless of bridging strategy, clopidogrel should be resumed with an LD as soon as oral administration is possible and the risk of severe bleeding is acceptable. Prasugrel and ticagrelor administration should be discouraged in the early period after major noncardiac surgery when there is an ongoing risk of serious bleeding. If oral administration of clopidogrel is not possible, postsurgery bridging with an intravenous agent should be considered.

- Patients with bleeding or at high risk for bleeding complications. Management of bleeding complications in patients on dual antiplatelet therapy goes beyond the scope of this document and is described elsewhere.<sup>97</sup> In dual antiplatelet therapy-treated patients who develop a bleeding complication, there is commonly a desire for de-escalation therapy. This should start with an MD regimen (ie, clopidogrel 75 mg), unless there has been a gap of therapy for  $\geq 5$  days, in which case a 300-mg LD might be used. Similar approaches should be considered for patients at high risk for bleeding complications such as those who have or develop thrombocytopenia, patients who develop a cerebrovascular event, and elderly patients, among others.
- Switching after thrombolysis. Clopidogrel therapy is the standard of care in patients treated with thrombolytics who require P2Y<sub>12</sub> inhibitor therapy. Escalation of P2Y<sub>12</sub> inhibitors is discouraged within 24 hours of thrombolysis because the combination of lytics with potent platelet inhibitors (ie, glycoprotein Ilb/IIIa inhibitors) increases bleeding<sup>98</sup>; after this duration, any escalation to a more potent regimen should occur with an LD regimen (prasugrel 60 mg or ticagrelor 180 mg).

- Patients requiring OAC. In patients requiring OAC who also undergo PCI requiring dual antiplatelet therapy, clopidogrel should be the P2Y<sub>12</sub> inhibitor of choice.<sup>99,100</sup> If patients are already on a newer-generation P2Y<sub>12</sub> inhibitor (eg, patients who already had PCI and develop atrial fibrillation requiring OAC), de-escalation therapy is recommended, and clopidogrel should be started with a 75-mg MD regimen. If patients are P2Y<sub>12</sub> inhibitor naïve, clopidogrel should be initiated with a 600-mg LD regimen (eg, patients with atrial fibrillation already on OAC who undergo PCI). Details of the management of PCI patients requiring OAC are given elsewhere.<sup>99,100</sup>
- Patients undergoing very late (>1 year) switch. De-escalation should occur with an MD regimen (no LD). Recently, a ticagrelor 60-mg twice-daily dosing regimen has been approved for post– myocardial infarction patients >1 year from their index event. When ticagrelor therapy is initiated for post–myocardial infarction patients >1 year from their index event, a switch should be made directly to 60-mg twice-daily MD (no LD) regardless of the prior P2Y<sub>12</sub> inhibitor used.<sup>35</sup>
- Patients on unknown therapy. It is not uncommon that patients are referred with unknown medication status. These patients should be treated as naïve, and an LD should be used.

### **CONCLUSIONS**

The current availability of a variety of P2Y<sub>12</sub> inhibitors provides clinicians with flexibility to optimize antiplatelet therapy for the individual patient. Although clinical data support the initiation and treatment of antiplatelet therapy with specific P2Y<sub>12</sub> inhibitors, clinical circumstances often arise that require the clinician to switch among the available therapies. Robust clinical outcomes data for specific switching strategies are lacking, but strategies can be guided by the different pharmacological profiles of these inhibitors, which may lead to DDIs that have potential implications for safety and efficacy. Therefore, this expert consensus document provides recommendations derived largely from pharmacodynamic and registry data, integrated with an understanding of the pharmacological principles of the agents involved. Ongoing dedicated studies will provide important insights into this topic.

### **AUTHORS**

Dominick J. Angiolillo, MD, PhD; Fabiana Rollini, MD; Robert F. Storey, MD; Deepak L. Bhatt, MD, MPH; Stefan James, MD, PhD; David J. Schneider, MD; Dirk Sibbing, MD; Derek Y.F. So, MD; Dietmar Trenk, PhD; Dimitrios Alexopoulos, MD; Paul A. Gurbel, MD; Willibald Hochholzer, MD; Leonardo De Luca, MD; Laurent Bonello, MD; Daniel Aradi, MD, PhD; Thomas Cuisset, MD, PhD; Udaya S. Tantry, PhD; Tracy Y. Wang, MD, MHS, MSc; Marco Valgimigli, MD, PhD; Ron Waksman, MD; Roxana Mehran, MD; Gilles Montalescot, MD; Francesco Franchi, MD; Matthew J. Price, MD

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# AFFILIATIONS

Division of Cardiology, University of Florida College of Medicine, Jacksonville (D.J.A., F.R., F.F.). Department of Infection, Immunity and Cardiovascular Disease, University of Sheffield, United Kingdom (R.F.S.). Brigham and Women's Hospital Heart & Vascular Center, Harvard Medical School, Boston, MA (D.L.B.). Department of Medical Sciences, Cardiology and Uppsala Clinical Research Center, Uppsala University, Sweden (S.J.). Department of Medicine, Cardiology Unit, Cardiovascular Research Institute, University of Vermont, Burlington (D.J.S.). Department of Cardiology, Ludwig-Maximilians-Universität München, Germany (D.S.). DZHK (German Center for Cardiovascular Research), partner site Munich Heart Alliance, Germany (D.S.). Division of Cardiology, University of Ottawa Heart Institute, Ontario, Canada (D.Y.S.F.). Department of Cardiology & Angiology II, University Heart Center Freiburg-Bad Krozingen, Germany (D.T., W.H.). Second Department of Cardiology, National and Capodistrian University of Athens, Attikon University Hospital, Greece (D. Alexopoulos). Inova Center for Thrombosis Research and Drug Development, Inova Heart and Vascular Institute, Falls Church, VA (P.A.G., U.S.T.). Division of Cardiology, Laboratory of Interventional Cardiology, San Giovanni Evangelista Hospital, Tivoli-Rome, Italy (L.D.L.). Assistance Publique-Hôpitaux de Marseille, Department of Cardiology, Hôpital Nord, Marseille, France (L.B.). Mediterranean Academic Association for Research and Studies in Cardiology, Marseille, France (L.D.L.). Aix-Marseille University, INSERM UMRS 1076, Marseille, France (L.D.L.). Heart Center Balatonfüred and Semmelweis University Budapest, Hungary (D. Aradi). Department of Cardiology, CHU Timone, and Aix-Marseille Université, Faculté de Médecine, Marseille, France (T.C.). Duke Clinical Research Institute, Duke University Medical Center, Durham, NC (T.Y.W.). Swiss Cardiovascular Center Bern, Bern University Hospital, Switzerland (M.V.). Section of Interventional Cardiology, MedStar Washington Hospital Center, DC (R.W.). Icahn School of Medicine at Mount Sinai, New York City, NY (R.M.) Sorbonne Université Paris 6, ACTION Study Group, Hôpital Pitié-Salpêtrière, France (G.M.). Division of Cardiovascular Diseases, Scripps Clinic, La Jolla, CA (M.J.P.).

# FOOTNOTES

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#### International Expert Consensus on Switching Platelet P2Y<sub>12</sub> Receptor–Inhibiting Therapies

Dominick J. Angiolillo, Fabiana Rollini, Robert F. Storey, Deepak L. Bhatt, Stefan James, David J. Schneider, Dirk Sibbing, Derek Y.F. So, Dietmar Trenk, Dimitrios Alexopoulos, Paul A. Gurbel, Willibald Hochholzer, Leonardo De Luca, Laurent Bonello, Daniel Aradi, Thomas Cuisset, Udaya S. Tantry, Tracy Y. Wang, Marco Valgimigli, Ron Waksman, Roxana Mehran, Gilles Montalescot, Francesco Franchi and Matthew J. Price

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## SUPPLEMENTAL MATERIAL

Name (Acronym)	Study Population (N)	Switch from Clopidogrel to new P2Y <sub>12</sub> receptor inhibitors (Prevalence)	Switch from new P2Y <sub>12</sub> receptor inhibitors to clopidogrel	Switch between new P2Y <sub>12</sub> receptor inhibitors	Clinical outcomes (exploratory)
Alexopoulos et al <sup>1</sup>	ACS	C → P: 40.1%	P → C: 1%	Between P and T:	Outcomes at 1 month:
(GRAPE)	undergoing PCI (N=1794)	C → T: 50.3%	T → C: 4.3%	4.3%*	higher risk of bleeding and less MACE in patients switched from C to P or T compared with those treated with C only. No differences in MACE and bleeding when compared with those initially treated with P or T.
Clemmensen et al <sup>2</sup>	STEMI	C → P: 48.7%	P → C: 8.3%	$P \rightarrow T: 2.8\%$	In-hospital outcomes: no
(MULTIPRAC)	(N=2053)	C → T: 11.6%	T → C: n/a	$T \rightarrow P: n/a$	differences in MACE and non-CABG related bleeding in patients switched from C to P vs patients on P only.
Bagai et al <sup>3</sup>	NSTEMI,	C → P: 10.4%	$P \rightarrow C: 11.5\%$	$P \rightarrow T: 0.3\%$	Outcomes at 6 months:
(TRANSLATE-ACS)	STEMI (N=11999)	C → T: 1%	T → C: 2.1%	T → P: 3.4%	no significant differences in MACE and bleeding in switched (any switching) vs non switched patients.

Supplemental Table 1. Switching Oral P2Y12 Receptor Inhibitors: Findings from Registries and Randomized Clinical Trials

Schiele et al <sup>4</sup>	NSTEMI,	C → P: 16.5%	$P \rightarrow C: 4.6\%$	n/a	n/a
(FAST-MI)	STEMI	$C \rightarrow T: n/a$	$T \rightarrow C: n/a$		
	(N=4101)				
De Luca et al <sup>5</sup>	NSTE-ACS	In cath-lab switch:	In cath-lab switch:	In cath-lab switch:	n/a
(EYESHOT)	STEMI,	$C \rightarrow P/T: 3\%$	P/T → C: 0.3%	Between P and T:	
	(N=2585)			0.3%	
		At discharge switch:	At discharge switch:		
		$C \rightarrow P/T: 9.6\%$	P/T → C: 3.2%	At discharge switch:	
				Between P and T:	
		Switch in medically	Switch in medically	1.3%	
		managed patients:	managed patients:		
		$C \rightarrow P/T: 2\%$	$P/T \rightarrow C: 3.4\%$	Switch in medically	
				managed patients:	
				Between P and T:	
				0.3%	
Bagai et al <sup>6</sup>	NSTEMI and	C → P: 5.2%	P → C: 11.5%	n/a	n/a
(ACTION Registry-	STEMI	$C \rightarrow T: n/a$	$T \rightarrow C: n/a$		
GWTG and CathPCI)	undergoing PCI				
	(N=47040)				
De Luca et al <sup>7</sup>	NSTEMI,	150 patients switched	n/a	n/a	Outcomes at 30-days: no
	STEMI, UA	from C to P matched			differences in MACE,
	underwent PCI	with 300 C only			NACE and bleeding
	(N=450)	treated patients			between groups.
Loh et al <sup>8</sup>	ACS	C → P: 14.8%	n/a	n/a	In-hospital outcomes: no
	undergoing PCI	$C \rightarrow T: n/a$			differences in bleeding
	(N=606)				between patients who
					switched to P compared
					those on P only. MACE
					were greater in patients

Almendro-Delia et al <sup>9</sup>	ACS (N=468)	$C \rightarrow P: 25\%$ $C \rightarrow T: n/a$	n/a	$P \rightarrow T: 0.3\%$ $T \rightarrow P: n/a$	<ul> <li>who switched to P</li> <li>compared to those on P</li> <li>only.</li> <li>In-hospital outcomes: no</li> <li>difference in bleeding</li> <li>and MACCE between</li> <li>patients who switched to</li> <li>P compared with those</li> </ul>
					on C only.
Zettler et al <sup>10</sup> (TRANSLATE-ACS)	NSTEMI, STEMI undergoing PCI (N= 8672)	C → P: 91.7% C → T: 8.3%	P → C: 97.4% T → C: 87.5%	P → T: 2.6% T → P: 12.5%	<ul> <li>Before switching: MACE: 18.5% in the C group, 2.1% in the P group, 1.5% in the T group.</li> <li>GUSTO moderate/severe Bleeding: 0.9% in the C group, 0.3% in the P group, 0% in the T group.</li> <li>After switching: MACE: 1.9% in the group who switched from C, 0.8% in the group who switched from P, 0 in the group who switched from T. No bleeding events were observed.</li> </ul>

Bagai et al <sup>11</sup>	NSTEMI,	C → P: 30.7% (HPR)	n/a	n/a	Outcomes at 1 year:
(TRANSLATE-ACS)	STEMI	C → P: 4.4% (w/o			MACE:
	undergoing PCI	HPR)			- HPR who switched:
	with HPR on	,			10%,
	clopidogrel	$C \rightarrow T: n/a$			- HPR who did not
	(N=261)				switch: 22.7%.
					- no HPR who switched:
					22.2%,
					- no HPR who did not
					switch: 12.8%.
					BARC type 2 or higher
					bleeding:
					- HPR who switched:
					23.8%,
					- HPR who did not
					switch: 22.1%.
					- no HPR who switched
					22.2%,
					- no HPR who did not
					switch: 19.4%.
De Luca et al <sup>12</sup>	ACS	In cath-lab switch:	In cath-lab switch:	In cath-lab switch:	• In-hospital outcomes:
(SCOPE)	undergoing PCI	$C \rightarrow P: 2.1\%$	n/a	n/a	NACE: 22.7% in patients
	(N=1363)	$C \rightarrow T: 0.2\%$			who switched from P/T
					to C, 0% in patients who
		At discharge switch:	At discharge switch:	At discharge switch:	switched from C to P/T
		$C \rightarrow P: 0.4\%$	$P \rightarrow C: 0.8\%$	P → T: 0.3%	and between P and T.
		C → T: 0.6%	T → C: 1%	$T \rightarrow P: 0.2\%$	MACE: 20.4% in
					patients who switched
					from P/T to C, 0% in

After discharge	After discharge	After discharge	patients who switched
switch:	switch:	switch:	from C to P/T and
$C \rightarrow P: 0.3\%$	$P \rightarrow C: 0.7\%$	P → T: 1.6%	between P and T.
$C \rightarrow T: 0.3\%$	T → C: 0.8%	T → P: 1.4%	Bleeding: 3.8% in
			patients who switched
From admission to	From admission to	From admission to	from P/T to C, 0% in
follow-up switch:	follow-up switch:	follow-up switch:	patients who switched
C → P: 1.2%	P → C: 1.3%	P → T: 1.9%	from C to P/T and
C → T: 1.4%	T → C: 1.8%	T → P: 1.6%	between P and T.
			• Follow up outcomes
			(~45days):
			NACE: 22.7% in patients
			who switched from P/T
			to C, 0% in patients who
			switched from C to P/T,
			4.3% in patients who
			switched between P and
			Т.
			MACE: 20.4% in
			patients who switched
			from P/T to C, 0% in
			patients who switched
			from C to P/T, 2.2% in
			patients who switched
			between P and T.
			Bleeding: 3.8% in
			patients who switched
			from P/T to C, 0% in
			patients who switched

					from C to P/T, 2.2% in
					patients who switched
					between P and T.
Cuisset et al <sup>13</sup>	ACS	n/a	P/T → C: 50% <sup>†</sup>	n/a	Outcomes at 1 year:
(TOPIC)	undergoing PCI				Net clinical benefit:
	(N=646)				13.4% in patients who
					switched, 26.3% in
					patients who did not
					switch.
Sibbing et al <sup>14</sup>	Biomarker-	$C \rightarrow P: 39\%^{\#\$}$	$P \rightarrow C: 50\%^{\dagger}$	n/a	Outcomes at 1 year:
(TROPICAL-ACS)	positive ACS				Net clinical benefit: 7%
	with a	$C \rightarrow T: n/a$	$T \rightarrow C: n/a$		in patients who switched
	successful PCI				(guided de-escalation
	(N=2610)				group), 9% in patients
					who did not switch
					(control group, 12
					months on P).

\*: cumulative rate only available.

<sup>†</sup>: mandated by the trial per randomization.

<sup>#</sup>: 19% of total trial population, 39% of patients de-escalated to clopidogrel.

<sup>\$</sup>: mandated by the trial in patients with high platelet reactivity on clopidogrel.

ACS: acute coronary syndrome; C: clopidogrel; CABG: coronary artery bypass graft; MACE: major adverse cardiac events; MACCE: Major Adverse Cardiac and Cerebrovascular event; NACE: net adverse clinical events; NSTEMI: non-ST-elevation myocardial infarction; STEMI: ST-elevation myocardial infarction; P: prasugrel; PCI: percutaneous coronary intervention; T: ticagrelor; UA: unstable angina. ACTION Registry-GWTG and CathPCI: National Cardiovascular Data Registry Acute Coronary Treatment and Intervention outcomes Network Registry-Get With the Guidelines and the National Cardiovascular Data Registry CathPCI Registry; EYESHOT: EmploYEd antithrombotic therapies in patients with acute coronary Syndromes HOspitalized in iTalian cardiac care units; GRAPE: GReek AntiPlatElet Registry; HPR: high on treatment platelet reactivity; MULTIPRAC: MULTInational non-interventional study of patients with ST-segment elevation myocardial infarction treated with PRimary Angioplasty and Concomitant use of upstream antiplatelet therapy with prasugrel or clopidogrel; SCOPE: Incidence and Outcome of Switching of Oral Platelet P2Y12 Receptor Inhibitors in Patients with Acute Coronary Syndromes Undergoing Percutaneous Coronary Intervention: The SCOPE Registry; TOPIC: Benefit of switching dual antiplatelet therapy after acute coronary syndrome: the TOPIC (timing of platelet inhibition after acute coronary syndrome) randomized study; TRANSLATE-ACS: TReatment with ADP receptor iNhibitorS: Longitudinal Assessment of Treatment Patterns and Events after Acute Coronary Syndrome.

Modified with permission from Rollini F et al. Nat Rev Cardiol 2016;13:11-27 [ref 15].

Study	Study	Study	PD test	Key PD switching findings	Clinical outcomes
(Acronym)	Design	Population			(exploratory)
		(N)			
Payne et al <sup>16</sup>	Open-label,	Healthy	LTA	MPA 37% on C $\rightarrow$ 5% 1 hr	Outcomes at 22 days: no
	randomized,	subjects		after P 60 mg	differences in bleeding
	fixed sequence	(N=35)			episodes or other adverse
				MPA 37% on C $\rightarrow$ 28% 1 hr	events.
				after P 10 mg	
Wiviott et al <sup>17</sup>	Multicenter,	Planned PCI	LTA	IPA 45.4% on C 150 mg $\rightarrow$	Outcomes at 29 days:
(PRINCIPLE-TIMI44)	randomized,	(N=201)	VASP	60.8% after 15 days on P 10	bleeding occurred in 4
	double-blind,		VN P2Y12	mg.	subjects switching from C
	double-				to P.
	dummy, active			PRI 39.7% on C 150 mg $\rightarrow$	
	comparator-			25.1% after 15 days on P 10	
	controlled,			mg.	
	crossover				
				VN: consistent finding (data	
				not reported)	
Montalescot et al <sup>18</sup>	Double-blind,	UA/NSTEMI	LTA	MPA 38.6% after C 900 mg→	Outcomes at 60 days:
(ACAPULCO)	randomized,	(N=56)	VN P2Y12	28.9% after 15 days on P 10	there were no differences
	crossover		VASP	mg	in any non-CABG-related
				MPA 38.6% after C 900 mg $\rightarrow$	TIMI major or
				38.2% after 15 days on C 150	GUSTO severe/life-
				mg $\rightarrow$ 25% after 15 days on P	threatening bleeding
				10 mg.	events. Five subjects (3 P
					and 2 C) experienced

# Supplemental Table 2. Pharmacodynamic studies of switching from clopidogrel to prasugrel

				PRU 96.3 ± 67.6 on C 150	bleeding during MD
				$mg \rightarrow 47.1 \pm 32.4$ after 15 days	treatment.
				on P 10 mg	
				PRI $40.6 \pm 22.5\%$ on C 150	
				$mg \rightarrow 22.8 \pm 15.7\%$ after 15	
				days on P 10 mg	
Angiolillo et al <sup>19</sup>	Multicenter,	Prior ACS	LTA	MPA 60.2% on C 75 mg→	Outcomes at 15 days:
(SWAP)	randomized,	(30-330 days)	VN P2Y12	41.1% after 7 days on P 10 mg.	bleeding by
	double-blind,	(N=139)	VASP		TIMI criteria was
	double-			MPA 55.5% on C 75 mg→	reported in 12.5% of the
	dummy, active-			41% after a dose of P 60 mg +	С
	control			7 days of P 10 mg.	75 mg MD group, 8.5%
					of the P 10 mg MD group,
				MPA 53.8% on C 75 mg $\rightarrow$	and 13.6% of the P
				55% after 7 days on C 75 mg.	LD+MD group.
				VN and VASP: consistent	
				finding (data not reported)	
Trenk et al <sup>20</sup>	Randomized,	Stable CAD	VN P2Y12	Observed a substantial	Outcomes at 6 months:
(TRIGGER-PCI)	parallel	with HPR		decrease in PRU in the P arm	TIMI major non-coronary
	assignment,	undergoing		compared to C.	artery bypass graft
	double blind	PCI		176 (94.1%) patients of the P	bleeding occurred in 3
		(N=212)		arm reached a PRU $\leq$ 208.	patients on P and 1 patient
					on C.
Diodati et al <sup>21</sup>	Randomized,	ACS	VN P2Y12	PRU 57.9, 6 hours after	Outcomes at 72 hours:
(TRIPLET)	double-blind,	undergoing		placebo/P 60 mg.	treatment-emergent
	double-	planned PCI			adverse events were 3 in
	dummy,	(N=282)			the placebo LD/P 60 mg

	3-arm, parallel, active- comparator controlled			PRU 35.6, 6 hours after C 600 mg/P 60 mg. PRU 53.9, 6 hours after C 600 mg/P 30 mg.	group; 4 in the C 600 mg LD/P 60 mg LD group; and 7 in the C 600 mg LD/P 30 mg. There were 2 deaths in the placebo/P 60 mg LD treatment group.
Rollini et al <sup>22</sup>	Prospective, randomized, parallel design, open-label	CAD (N=110)	LTA VN P2Y12 VASP	MPA 50.3±% on C $\rightarrow$ 32±6% 30 min after P 60 mg, $\rightarrow$ 18.8±4% 2 hours after P 60 mg, $\rightarrow$ 16.6±2% 24 hours after P 60 mg, $\rightarrow$ 28.6±4% 1 week after P 60 mg. PRU 182±19 on C $\rightarrow$ 105±23 30 min after P 60 mg, $\rightarrow$ 33±13 2 hours after P 60 mg, $\rightarrow$ 15±11 24 hours after P 60	Outcomes at 1 week: 2 minor bleeding.

				mg, $\rightarrow$ 73±14 1 week after P	
				60 mg.	
				oo mg.	
				PRI 60.8 $\pm$ 7% on C $\rightarrow$	
				33.6±7% 30 min after P 60	
				mg, $\rightarrow$ 13.4±6% 2 hours after	
				P 60 mg, $\rightarrow$ 14.2±4% 24 hours	
				after P 60 mg, → 32.1±5% 1	
				week after P 60 mg.	
Sardella et al <sup>23</sup>	Open-label,	Stable CAD	MEA	AUC 576 on C→180.5 after 15	Outcomes at 3 months: 3
(RESET GENE)	crossover	with HPR		days on P 10 mg	minor bleedings in
	randomized	undergoing			patients initially treated
		PCI		AUC 380.5 on C 150 mg $\rightarrow$	with P and 1 minor
		(N=32)		256 after 15 days on P 10 mg	bleeding while on C.
Lhermusier et al <sup>24</sup>	Prospective,	ACS	VN P2Y12	PRU 143 (53-199) after C 600	n/a
	open-label,	(N=48)	VASP	$mg \rightarrow 111 (7-127) 4$ hours after	
	randomized			P 10 mg→ 97 (41-145) 24	
				hours after initial P dose.	
				PRU 122 (82-149) after C 600	
				mg $\rightarrow$ 7 (6-31) 4 hours after P	
				30 mg→ 27 (7-20) 24 hours	
				after initial P dose.	

				PRI 44 (29-58)%, after C 600 mg $\rightarrow$ 35 (16-41)%, 4 hours after P 10 mg $\rightarrow$ 21 (19-58)%, 24 hours after initial P dose. PRI 59 (16-68)%, after C 600 mg $\rightarrow$ 15 (2-30)%, 4 hours after P 30 mg $\rightarrow$ 10 (7-20)%, 24 hours after initial P dose.	
Alexopoulos et al <sup>25</sup>	Randomized, single-center, single-blind, crossover	ACS with HPR (N=44)	VN P2Y12	PRU 280.3 on C → 90.8 after 15 days on P 10 mg → 32.1 after 15 days on T 90 mg/b.i.d.	Outcomes at 30 days: no patient exhibited a major adverse cardiovascular event or a major bleeding event; 4 patients (2 P and 2 T) reported minimal bleeding events. Allergic reactions (n=2), dyspepsia, (n=2), dyspnea (n=4) occurred with T.
Koul et al <sup>26</sup>	Prospective, observational registry	STEMI undergoing PCI (N=223)	VASP	PRI 79% after C 600 (pre-PCI) mg→74% after P 60 mg (after PCI)→ 17% 1 day after PCI	Outcomes in-hospital: 1.1% major bleeding.
Cuisset et al <sup>27</sup>	Prospective, observational registry	NSTE ACS with DM undergoing PCI (N=107)	VASP	PRI 47±21% after C 600 mg→ 31±13% after 1 month on P 10 mg. On C: HPR 50%	Outcomes at 30 days: one stent thrombosis; 10 BARC bleeding complications.

Nührenberg et al <sup>28</sup>	Non- randomized, observational	STEMI undergoing PCI	VN P2Y12 LTA MEA	LPR 13% On P: HPR 8% LPR 22% PRU 10 (8-31) after C 600mg + P 60 mg MPA 1 after C 600mg + P 60	n/a
		(N=47)		mg AU*min 214 (184-250) after C 600mg + P 60 mg	
Parodi et al <sup>29</sup>	Non- randomized, observational	CAD undergoing PCI (N=454)	LTA	HPR group: MPA 72±11% on C→ 43±16% on P	Outcomes in hospital and at 6 months: there were no differences in major or minor TIMI bleeding rates or in BARC bleeding rates or ischemic events between the patients switching and naïve.
Aradi et al <sup>30</sup>	Prospective, observational registry	ACS undergoing PCI (N=741)	MEA	P 60 mg/10 mg provided significantly more potent platelet inhibition than the repeated LD of C 600 mg. 86% of the P MD treated patients remained below the cut point for HPR	Outcomes at 1 year: rates of thrombotic complications in HPR group were similar to those in the no HPR group without any difference in all-cause death, myocardial

					infarction, stent
					thrombosis, or stroke. No
					excess of major bleeding
					after switching to P in
					HPR group compared
					to those without HPR.
Mayer et al <sup>31</sup>	Prospective	ACS with	MEA	AU*min 651 (543-780) after C	Outcomes at 30 days: 2
(ISAR-HPR)	observational	HPR		600 mg→ 156 (88-261) after P	(1.7%) combined
	registry	undergoing		60 mg	death/stent thrombosis
		PCI			and 10 (8.7%) TIMI
		(N=428)			major bleeding.
Lhermusier et al <sup>32</sup>	Open-label,	ACS with	VN P2Y12	PRU 234 (164-267) after C	Outcomes at discharge:
	multicenter,	planned	VASP	600 mg→ 23 (5-71), 4 hours	no differences in bleeding
	nonrandomized	invasive		after P 60 mg→ 9 (5-47) at	between groups (3 in the
	observational	strategy		discharge on P 10 mg.	C to P group and 3 in the
		(N=75)			P only group).
				PRI 68.4 (31.53-79.61)% after	
				C 600 mg→ 8.67 (4.51-	
				16.85)%, 4 hours after P 60	
				mg→ 8.05 (5.12-13.38)% at	
				discharge on P 10 mg.	

ACS: acute coronary syndrome; AU\*min: aggregation unit \* minutes; BARC: Bleeding Academic Research Consortium; bid: twice a day; C: clopidogrel; GP IIb/IIIa inhibitor: glycoprotein IIb/IIIa inhibitors; GUSTO: Global Use of Strategies to Open Occluded Arteries; HPR: high on-treatment platelet reactivity; LD: loading dose; LPR: low on-treatment platelet reactivity; LTA: light transmission aggregometry; MD: maintenance dose; MEA: multiple electrode platelet aggregometry; MPA: maximal platelet aggregation; NSTEMI: non-ST elevation myocardial infarction; P: prasugrel; PCI: percutaneous coronary intervention; PRU: P2Y12 reaction units; PRI: platelet reactivity index; T: ticagrelor; STEMI: ST-elevation myocardial infarction; TIMI: Thrombolysis in Myocardial Infarction; UA: unstable angina; VNP2Y12: VerifyNow P2Y12; VASP: Vasodilator-stimulated phosphoprotein.

ACAPULCO: Prasugrel compared with high-dose clopidogrel in acute coronary syndrome; ISAR-HPR: A comparative cohort study on personalised antiplatelet therapy in PCI-treated patients with high on-clopidogrel platelet reactivity; PRINCIPLE-TIMI44: prasugrel in Comparison to Clopidogrel for Inhibition of Platelet Activation and Aggregation-Thrombolysis in Myocardial Infarction 44; RESET GENE: The pharmacodynamic effects of switching theRapy in patiEntS with high on-trEatment plateleT reactivity and GENotypE variation with high clopidogrel dose versus prasugrel; SWAP: SWitching Anti Platelet; TRIGGER-PCI: Testing platelet Reactivity In patients underGoing elective stent placement on clopidogrel to Guide alternative thErapy with pRasugrel; TRIPLET: Transferring From Clopidogrel Loading Dose to Prasugrel Loading Dose in Acute Coronary Syndrome Patients. Modified from Rollini F et al. Nat Rev Cardiol 2016;13:11-27 [ref 15].

Study	Study	Study	PD test	Key PD switching findings	Clinical outcomes
(Acronym)	Design	Population			(exploratory)
		(N)			
Gurbel et al <sup>33</sup>	Randomized,	Stable CAD	LTA	Nonresponder cohort:	Outcomes at 30
(RESPOND)	double-blind,	(N=98)	VN P2Y12	MPA 59 $\pm$ 9% on C $\rightarrow$ 35 $\pm$ 11%, 4 hours	days: 4 patients (2
	double-dummy		VASP	after T 180 mg.	nonresponders and
	crossover				2 responders)
				Responder cohort:	experienced the 5
				MPA 47 $\pm$ 15% on C $\rightarrow$ 32 $\pm$ 8%, 4 hours	serious adverse
				after T 180 mg.	events, and all
					events occurred
				VN and VASP: consistent finding (data	during or after T
				not reported)	therapy. One major
					and 3 minor
					bleeding events
					occurred during T
					treatment, and no
					bleeding events
					occurred during C
					treatment. Dyspnea
					was reported in 13
					and 4 patients
					receiving T and C,
					respectively. Two
					nonresponder
					patients had

## Supplemental Table 3. Pharmacodynamic studies of switching from clopidogrel to ticagrelor

					dyspnea during
24					switching.
Caiazzo et al <sup>34</sup>	Randomized,	ACS	MEA	AU 34.4 $\pm$ 1.3 on C $\rightarrow$ 17.6 $\pm$ 7.2, 2 hours	Outcomes at 30
(SHIFT-OVER)	single-blind	(N=50)	LTA	after T 90 mg.	days: no deaths nor
				AU 41.7 $\pm$ 2 on C $\rightarrow$ 18.1 $\pm$ 6, 2 hours	strokes were
				after T 180 mg.	reported after
					switch. A total of 2
				MPA 24 $\pm$ 17% on C $\rightarrow$ 9 $\pm$ 4%, 2 hours	patients underwent
				after T 90 mg.	a new
				MPA 25 $\pm$ 14% on C $\rightarrow$ 9 $\pm$ 3%, 2 hours	hospitalization
				after T 180 mg.	within the shift to
					Т.
Lhermusier et al <sup>24</sup>	Prospective,	ACS	VN P2Y12	PRU 146 (97-236) after C 600 mg→ 12	Outcomes at 24
	open-label,	(N=48)	VASP	(4-46), 4 hours after T 90 mg→ 9 (5-	hours: 1 bleeding
	randomized			10), 24 hours after initial T dose.	event following
				PRU 92 (33-143) after C 600 mg→ 4	switch to T 90mg.
				(2-6) 4 hours after T 180 mg $\rightarrow$ 4 (3-5),	
				24 hours after initial T dose.	
				PRI 57 (18-80)%, after C 600 mg→ 3	
				$(2-6)\%$ , 4 hours after T 90 mg $\rightarrow$ 4 (2-	
				11)%, 24 hours after initial T dose.	
				PRI 47 (10-65)%, after C 600 mg→ 2	
				$(0-5)\%$ , 4 hours after T 180 mg $\rightarrow$ 3 (0-	
				4)%, 24 hours after initial T dose.	
Rollini et al <sup>22</sup>	Prospective,	CAD	LTA	MPA 50.2 $\pm$ 6% on C $\rightarrow$ 42 $\pm$ 6% 30 min	Outcomes at 1
	randomized,	(N=110)	VN P2Y12	after T 180 mg, $\rightarrow$ 21.9 $\pm$ 4% 2 hours	week: 1 minor

pa	arallel design,	VASP	after T 180 mg, $\rightarrow$ 18.9±1.1% 24 hours	bleeding, 26
ор	pen-label		after T 180 mg (12 hours after T 90	dyspnea
			mg), $\rightarrow$ 34±4% 1 week after T 180 mg	
			(12 hours after T 90 mg).	
			PRU 170±24 on C $\rightarrow$ 125±25 30 min	
			after T 180 mg, $\rightarrow$ 28±15 2 hours after	
			T 180 mg, $\rightarrow$ 20±10 24 hours after T	
			180 mg (12 hours after T 90 mg), $\rightarrow$	
			$58\pm15$ 1 week after T 180 mg (12 hours	
			after T 90 mg).	
			PRI 57.6 $\pm$ 9% on C $\rightarrow$ 41.1 $\pm$ 8% 30 min	
			after T 180 mg, $\rightarrow$ 15.6±6% 2 hours	
			after T 180 mg, → 19.7±5% 24 hours	
			after T 180 mg (12 hours after T 90	

				mg), → 32.6±5% 1 week after T 180	
				mg (12 hours after T 90 mg).	
Alexopoulos et al <sup>25</sup>	Prospective,	ACS with	VN P2Y12	PRU 277.4 on C $\rightarrow$ 34.1 after 15 days	Outcomes at 30
	randomized,	HPR		on T 90mg/b.i.d $\rightarrow$ 111.4 after 15 days	days: no patient
	single-blind,	(N=44)		on P 10 mg.	exhibited a major
	crossover				adverse
					cardiovascular
					event or a major
					bleeding event; 4
					patients (2 P and 2
					T) reported
					minimal bleeding
					events.
					Allergic reactions
					(n=2), dyspepsia,
					(n=2), dyspnea
					(n=4) occurred with
					Т.
Hibbert et al <sup>35</sup>	Prospective,	STEMI	VN P2Y12	PRU 252 (233-280) naïve	n/a
(CAPITAL RELOAD)	observational	(N=52)		patients $\rightarrow$ 220(82-269) 2 hours after T	
				180 mg.	
				PRU 255 (233-304) after C 600 mg→	
				90 (5-205) 2 hours after T 180 mg.	
Koul et al <sup>26</sup>	Prospective	STEMI	VASP	PRI 64% after C 600 (pre-PCI)	In-hospital
	registry	undergoing		mg→53% after P 60 mg (after PCI)→	outcomes: the rate
		PCI		29% 1 day after PCI	of major in-hospital
		(N=223)			

		bleeding was 3.3%
		in this cohort.

ACS: acute coronary syndrome; AU: aggregation unit; C: clopidogrel; CAD: coronary artery disease; HPR: high on-treatment platelet reactivity; LTA: light transmission aggregometry; MEA: multiple electrode platelet aggregometry; MPA: maximal platelet aggregation; P: prasugrel; PCI: percutaneous coronary intervention; PRU: P2Y12 reaction units; PRI: platelet reactivity index; T: ticagrelor; STEMI: ST-elevation myocardial infarction; VNP2Y12: VerifyNow P2Y12; VASP: Vasodilator-stimulated phosphoprotein.

CAPITAL RELOAD: A Comparative Pharmacodynamic Study of Ticagrelor versus Clopidogrel and Ticagrelor in Patients Undergoing Primary Percutaneous Coronary Intervention; RESPOND: Response to Ticagrelor in Clopidogrel Nonresponders and Responders and Effect of Switching Therapies; SHIFT-OVER: The Administration of a Loading Dose Has No Additive Effect on Platelet Aggregation During the Switch From Ongoing Clopidogrel Treatment to Ticagrelor in Patients With Acute Coronary Syndrome.

Modified from Rollini F et al. Nat Rev Cardiol 2016;13:11-27 [ref 15].

Study	Study	Study	PD test	Key PD switching findings	Clinical outcomes
(Acronym)	Design	Population			(exploratory)
		(N)			
Gurbel et al <sup>33</sup>	Randomized,	Stable CAD	LTA	Nonresponder cohort:	Outcomes at 30 days:
(RESPOND)	double-blind,	(N=98)	VN P2Y12	MPA 36 $\pm$ 14% on T $\rightarrow$ 56 $\pm$ 9% 4 hours	4 patients (2
	double-dummy		VASP	after C 600 mg.	nonresponders and 2
	crossover				responders)
				Responder cohort:	experienced the 5
				MPA 25 $\pm$ 11% on T $\rightarrow$ 45 $\pm$ 8% 4 hours	serious adverse
				after C 600 mg.	events, and all events
					occurred during or
				VN and VASP: consistent finding	after T therapy. One
				(data not reported)	major and 3 minor
					bleeding events
					occurred during T
					treatment, and no
					bleeding events
					occurred during C
					treatment. Dyspnea
					was reported in 13
					and 4 patients
					receiving T and C,
					respectively. Two
					nonresponder patients
					had dyspnea during
					switching.

## Supplemental Table 4. Pharmacodynamic studies of switching from newer generation P2Y<sub>12</sub> inhibitors to clopidogrel

Wiviott et al <sup>17</sup>	Multicenter,	Planned PCI	LTA	IPA 61.9% on P 10 mg→ 46.8% after	Outcomes at 29 days:
(PRINCIPLE-TIMI44)	randomized,	(N=201)	VASP	15 days on C 150 mg.	no bleeding events in
	double-blind,		VN P2Y12		subject switching
	double-			PRI 21.7% on P 10 mg→ 48% after	from P to C.
	dummy, active			15 days on C 150 mg.	
	comparator-				
	controlled,			VN: consistent finding (data not	
	crossover			reported)	
Montalescot et al <sup>18</sup>	Double-blind,	UA/NSTE	LTA	MPA 28.9% on P 10 mg→ 42.5%	Outcomes at 60 days:
(ACAPULCO)	randomized,	MI	VN P2Y12	after 15 days on C 150 mg.	there were no
	crossover	(N=56)	VASP		differences in any
				PRU $41.3 \pm 43.8$ on P 10 mg $\rightarrow$ 101.3	non-CABG-related
				$\pm$ 60.8 after 15 days on C 150 mg	TIMI major or
					GUSTO severe/life-
				PRI 21.5 $\pm$ 13.4% on P 10 mg $\rightarrow$ 38.1	threatening bleeding
				$\pm$ 19.8% after 15 days on C 150 mg	events. Five subjects
					(3 P and 2 C)
					experienced bleeding
					during MD treatment.
Sardella et al <sup>23</sup>	Open-label,	Stable CAD	MEA	AUC 180.5 after 15 on P 10 mg→330	Outcomes at 3
(RESET GENE)	crossover	with HPR		after 15 days on C 150 mg	months: 3 minor
	randomized	undergoing			bleedings in patients
		PCI			initially treated with
		(N=32)			P and 1 minor
					bleeding while on C.
Pourdjabbar et al <sup>36</sup>	Prospective,	ACS	VN P2Y12	PRU~40 on T*→114±73.1, 48 hrs	Outcomes at 30 days:
(CAPITAL OPTI-	randomized,	(N=60)		after C 600 mg	No differences in
CROSS)	open-label			PRU~40 on T* $\rightarrow$ 165.1±70.5, 48 hrs	MACE, TIMI major
				after C 75 mg	bleeding or stent

				_	thrombosis between
				PRU~40 on T* $\rightarrow$ 165.8±71, 72 hrs	groups.
				after C 600 mg	
				PRU~40 on T* $\rightarrow$ 184.1±68, 72 hrs	
				after C 75 mg	
				HPR rate after C 600 mg: 27%	
				HPR rate after C 75 mg: 57%	
Kerneis et al <sup>37</sup>	Prospective,	ACS	LTA	MPA 21.01±10.47% on P 10 mg→	Outcomes at 30 days:
	observational	(N=31)	<b>VN P2Y12</b>	43.84±15.19% after 15 on C 75 mg.	no major bleedings.
	registry		VASP		
				PRU 14.23±27.98 on P 10 mg→	
				155±87.24 after 15 days on C 75 mg.	
				PRI 12.55±11.9% on P 10 mg→	
				43.63±21.82% after 15 days on C 75	
				mg.	
Deharo et al <sup>38</sup>	Prospective,	ACS with	VASP	PRI 7±2% on P→ 37.8±15.6% on C	Outcomes at 30 days:
(POBA)	observational	LPR			no bleeding events
		(N=20)			after switching to C.

\*: value estimated from the figure.

ACS: acute coronary syndrome; BARC: Bleeding Academic Research Consortium; C: clopidogrel; CAD: coronary artery disease; GP IIb/IIIa inhibitor: glycoprotein IIb/IIIa inhibitors; GUSTO: Global Use of Strategies to Open Occluded Arteries; HPR: high on treatment platelet reactivity; IPA: inhibition of platelet aggregation; LPR: low platelet reactivity; LTA: light transmission aggregometry; MD: maintenance dose; MPA: maximal platelet aggregation; NSTEMI: non-ST elevation myocardial infarction; P: prasugrel; PCI: percutaneous coronary intervention; PRU: P2Y12 reaction units; PRI: platelet reactivity index; T: ticagrelor; TIMI: Thrombolysis in Myocardial Infarction; UA: unstable angina; VNP2Y12: VerifyNow P2Y12; VASP: Vasodilator-stimulated phosphoprotein.

ACAPULCO: Prasugrel compared with high-dose clopidogrel in acute coronary syndrome; CAPITAL OPTI-CROSS: Optimizing crossover from ticagrelor to clopidogrel in patients with acute coronary syndrome: the Capital opti-cross randomized trial; POBA: Predictor of Bleeding with Antiplatelet drugs; PRINCIPLE-TIMI44: prasugrel in Comparison to Clopidogrel for Inhibition of Platelet Activation and Aggregation-Thrombolysis in Myocardial Infarction 44; RESET GENE: The pharmacodynamic effects of switching theRapy in patiEntS with high on-trEatment plateleT reactivity and GENotypE variation with high clopidogrel dose versus prasugrel; RESPOND: Response to Ticagrelor in Clopidogrel Nonresponders and Responders and Effect of Switching Therapies. Modified from Rollini F et al. Nat Rev Cardiol 2016;13:11-27 [ref 15].

Study	Study Design	Study Population (N)	PD test	Key PD switching findings	Clinical outcomes (exploratory)
Schneider et al <sup>39</sup>	Prospective not randomized. Transition between Cang and T	Stable CAD (N=12)	LTA VN P2Y12 VASP	Switching from Cang to T (T administered during Cang infusion): MPA 20µmol/1: 2.3 $\pm$ 2.2% on C $\rightarrow$ 19 $\pm$ 16%, 30 min after stopping infusion. MPA 5µmol/1: 4.1 $\pm$ 2% on C $\rightarrow$ 10 $\pm$ 7%, 30 min after stopping infusion. VNP2Y12: 6.4 $\pm$ 7.5% on C $\rightarrow$ 76 $\pm$ 79%, 30 min after stopping infusion. VASP: 15 $\pm$ 9% on C $\rightarrow$ 48 $\pm$ 49%, 30 min after stopping infusion. No significangt increase in platelet reactivity compared with the pre- defined reference time point (5.25 h after stopping infusion) with each assay <i>Switching from T to Cang</i> (Cang started 12 h after last T dose): MPA 20µmol/1: 12 $\pm$ 9% on T $\rightarrow$ 1.5 $\pm$ 1.9 during Cang infusion. VN and VASP: consistent findings Previous treatment with ticagrelor did not alter the inhibitory effect of canggrelor.	Throughout study participation (telephone interview on study days 10-12): no adverse events

Supplemental Table 5. Pharmacodynamic studies of switching between cangrelor and oral P2Y<sub>12</sub> receptor inhibitors

Schneider et al <sup>40</sup>	Prospective not randomized Transition between Cang and P	Stable CAD (N=15)	LTA VN P2Y12 VASP	Switching from Cang to P (P administered during Cang infusion): MPA 20µmol/l: <4% on C→ 41%, 30 min after stopping infusion when P administered 30 before infusion stop; ~60%, 30 min after stopping infusion when P administered 1 before infusion stop or at the end of infusion. VN and VASP: consistent findings. Switching from P to Cang (Cang started 24 h after last P dose): MPA 20µmol/l: <5% on P→ <5%, during Cang infusion. VN and VASP: consistent findings	Throughout study participation (telephone interview on study days 13-15): no adverse events
Angiolillo et al <sup>41</sup>	Prospective, randomized, double-blind, placebo controlled	ACS (N=210)	VN P2Y12	PRU 210.9±94.0 prior Cang infusion ( thienopyridine administered median 29 h prior), 68.9±67.8 during infusion, 279.7±106.5 pre-CABG (median 3.2 h post infusion)	Study-defined excessive CABG- surgery-related bleeding not significantly different between Cang (11.8%) or placebo (10.4%). Ischemic end points prior to surgery were low: 2.8% with Cang, 4.0% with placebo
Angiolillo et al <sup>42</sup>	Predefined	CAD	VN P2Y12	PRU 93.5 on Cang → ~240 6-12 h	Not reported
	substudy of CHAMPION-	undergoing PCI	LTA VASP	after stopping infusions and administering C.	

	PCI and CHAMPION- PLATFORM randomized trials	(N=167)		Post infusion PRU was not significantly different in patients treated initially with Cang+C or placebo+C LTA and VASP: consistent findings.	
Steinhubl et al <sup>43</sup>	Randomized, open-label	Healthy volunteers (N=20)	Whole- blood impedance aggregomet ry LTA Flow cytometry	<ul> <li>Whole blood aggregometry:</li> <li>ADP-induced aggregation completely inhibited during Cang infusion.</li> <li>C at the end of cang infusion: max inhibition 3 h post C administration.</li> <li>C alone: max inhibition 2 h post C administration.</li> <li>C and Cang simultaneously: platelet reactivity similar to baseline (no antiplatelet therapy) at all time points after C administration.</li> <li>LTA and Flow cytometry: consistent findings.</li> </ul>	n/a
Schneider et al <sup>44</sup>	Prospective not randomized. Transition between Cang and C	Stable CAD (N=12)	LTA VN P2Y12 Flow cytometry	LTA ADP 20µmol/1: MPA ≤2 during Cang inf. MPA 1 h after stop Cang infusion: 56±9% with C at the end of infusion 56±17 with C 30 min prior stop infusion 65±10 with C 1 h prior stop infusion. VN P2Y12 and Flow cytometry: consistent findings	n/a
Hochholzer et al <sup>45</sup>	Prospective, randomized, open-label.	CAD without MI	Whole- blood impedance	HPR 1 h after stopping Cang: C: 35%; P: 6.7%; T: 4.4%	The incidence of 30-day ischemic and bleeding events

Transition	undergoing	aggregomet	Platelet reactivity (median):	was similar among
from Cang to	PCI (n=110)	ry	94 AU x min on Cang $\rightarrow$	groups.
C, P, T*			212 with C, 154 with P, 81 with T 1 h	MI: 0% with C,
			after stopping infusion	11.1% with P and
				4.4% with T.
				TIMI major
				bleeding:
				0% with C, 0%
				with P and 2.2%
				with T.

\* C 600 mg was administered immediately after discontinuation of Cang infusion. P 60 mg and T 180 mg were administered at the start of Cang infusion.

ACS: acute coronary syndrome; ADP: adenosine diphosphate; AU x min: aggregation units per min; C: clopidogrel; CAD: coronary artery disease; Cang: cangrelor; LTA: light transmission aggregometry; MI: myocardial infarction; MPA: maximal platelet aggregation; P: prasugrel; PCI: percutaneous coronary intervention; PRU: P2Y12 reaction units; T: ticagrelor; VNP2Y12: VerifyNow P2Y12; VASP: Vasodilator-stimulated phosphoprotein.

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