Switching of Ticagrelor to Clopidogrel at 3 Months in Patients Treated for Acute Coronary Syndrome: Single Centre Experience

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Abstract

Objective: ESC guidelines recommend Dual Anti-Platelet Therapy (DAPT) with aspirin and ticagrelor for 12 months following Non-ST Elevation Myocardial Infarction (NSTEMI). We aimed to assess the safety of switching DAPT with aspirin and ticagrelor to aspirin and clopidogrel after 3 months in patients following NSTEMI, with the maximum duration of DAPT of 12 months.

Material and methods: Patients admitted with NSTEMI between 2011-2012 were identified using the ICD-10 and OPCS-4 coding systems. Retrospective analysis was then performed using electronic records for additional information.

Results: 98 patients were treated with aspirin and ticagrelor following admission with MI. 64% (63/98) were male, 55.1% (54/98) were hypertensive, 66.3% (65/98) with hyperlipidaemia, 20.4% (20/98), had diabetes and 33.7% (33/98) had previous known ischemic heart disease. 30.7% (30/98) had BMI > 30. 74.5% (73/98) underwent percutaneous coronary intervention with stenting of the target lesions, 20.4% (20/98) treated medically while 4.1% (4/98) referred for coronary bypass surgery. 8.2% (8/98) patients were re-admitted within 90 days of NSTEMI before the switchover of DAPT (3 for angina, 2 for non-cardiac chest pains and 3 for non-cardiac conditions), and none after that period. In 51% (50/98) patients DAPT was switched to clopidogrel at 3 months with 49% (48/98) staying on aspirin and ticagrelor. There were three non-cardiac deaths in the follow-up period.

Conclusion: This study shows the potential for the safe switchover of DAPT to clopidogrel following 3 months therapy with ticagrelor for NSTEMI, whilst enhancing cost-savings.

Keywords: Ticagrelor; Clopidogrel; Acute coronary syndrome

Introduction

Treatment with dual antiplatelet therapy including aspirin and a P2Y12 inhibitor is gold standard therapy in the management of patients with Acute Coronary Syndromes (ACS) [1,2]. Ticagrelor, a reversible oral P2Y12 receptor antagonist provides faster, greater and consistent platelet inhibition than Prasugrel and clopidogrel [3-5]. In the PLATO trial (The Platelet Inhibition and Patient Outcomes) [6], ticagrelor demonstrated superiority in preventing death from cardiovascular causes, non-fatal MI and stroke and without an increase in rate of overall major bleeding. Prasugrel is an irreversible P2Y12 inhibitor. Prasugrel has shown reduced ischemic events in patients with ACS planned for Percutaneous Coronary Interventions (PCI) in the TRITON Trial [7], however it showed no benefit over clopidogrel in Non-ST-Elevation Acute Coronary Syndrome (NSTEMI) patients who are for management without revascularization (TRILOGY-ACS trial) [8]. 12 months of Dual-Antiplatelet Therapy (DAPT) with a P2Y12 inhibitor and aspirin is accepted to offer a more clinically-effective option than treatment with clopidogrel and aspirin, however there has been little research into the switching of anti-platelet agents, especially from ticagrelor to clopidogrel, at intervals earlier than 12 months.

Occasionally, in clinical scenarios such as following the onset of 'unpleasant' side effects associated with the newer P2Y12 inhibitors, the recommendation is to switch DAPT. Such 'switching' of DAPT is often to the older clopidogrel-aspirin combination. Admittedly, such 'switchover' of DAPT is based on case-by-case recommendation of the supervising clinician and is rather uncommon. Nevertheless, there are no reported adverse outcomes from such 'downgrading' of DAPT after an initial period of more aggressive DAPT.
time whereby approval for ticagrelor therapy was limited to 3 months only due to high cost. Additional information was collected through follow-up clinic letters, patient electronic records and procedural databases. Recorded data fields included patient demographics, comorbidities, treatment strategy, duration and type of DAPT, hospital re-admissions and mortality.

Patients admitted with ACS were treated with a loading dose of aspirin 300 mg, followed by 75 mg once daily. The second antiplatelet agent was ticagrelor with a loading dose of 180 mg followed by 90 mg twice daily. In addition to optimal medical therapy, patient were treated with Percutaneous Coronary Interventions (PCI) with Bare Metal or Drug-Eluting Stents based on standard practice of the operator, surgical revascularization with Coronary Artery By-Pass Surgery (CABG) or medical therapy only.

Clinical definitions

All deaths were considered as cardiac unless a specific alternate cause of death was demonstrated.

Post procedural MI was defined as the presence of at least 2 of the following: Cardiac chest pains, rise in cardiac enzymes (creatine kinase or troponin) and new ECG changes.

MACE was defined as: all-cause mortality, cardiac mortality, myocardial infarction (MI), stroke and Target Vessel Revascularization (TVR).

The Target Lesion Revascularization (TLR) was defined as a repeated intervention within the stent or in the immediate distal or proximal segments adjacent to the stent.

TVR was defined as any repeat PCI or CABG of the target vessel.

The successful PCI for the target coronary lesion was defined as restoration of TIMI flow grade 3 without major complications after stent deployment.

Follow-up time was defined as the interval from the index procedure to the last contact.

Follow-up

Patients were followed up routinely at 3-6 months and/or until discharged (discretion of clinician) to the community based on the symptoms. At follow-up patients had a history review and clinical examination. Re-admissions with cardiac events were checked using hospital electronic records, catheter-lab database, and all other clinical letters (admissions & outpatients).

Statistical analyses were performed using SPSS 20 software (SPSS Inc., Chicago, IL, USA). A p-value of <0.05 was considered statistically significant.

Results

98 patients with a mean age of 67.7 years were treated with aspirin and ticagrelor following admission with MI as per guidelines. Patient’s demographics are shown in Table 1.

Treatment strategy varied amongst patients; 73 (74.5%) underwent PCI with stenting of the target lesions. 20 (20.4%) were treated medically while 4 (4.1%) were referred for CABG. In 50 patients (51%) DAPT was switched to aspirin and clopidogrel at 3 months (Group 2), and the remaining 48 patients (49%) stayed on aspirin and ticagrelor (Group 1) for a total of 12 months. The medications were changed by the local family physicians as per local policy at the time.

In total, 9 patients (8.2%) were re-admitted within 90 days of NSTE-ACS, before the switchover of DAPT (3 for angina and 6 with non-coronary related conditions) and none after that period. 4/9 (44%) patients had been treated with PCI, 4 (44%) treated medically and 1 (11%) treated with CABG. At full follow up of the study population, there were 3 deaths (3%) (All of these patients were >80 years old, treated medically and had non-cardiac deaths). There was no other documented MACE (MI, TVR, stroke, major bleed) recorded in either group (ticagrelor vs clopidogrel).

Discussion

Currently both NICE and ESC guidelines recommend aspirin and a P2Y12 inhibitor (clopidogrel/ticagrelor/Prasugrel) as DAPT for 12 months post ACS. The newer antiplatelets have significant cost implications, as they are considerably more expensive. We report absence of any significant adverse outcomes in an observational all-comer study of patients admitted with a clinical presentation of IHD following the ‘switchover’ of DAPT after a 3 month period of aggressive DAPT with the newer P2Y12 inhibitor (ticagrelor). In particular, the downgrading of DAPT to clopidogrel-aspirin did not lead to any significant ischaemic complications.

It is plausible that there are some significant differences between the active/unstable forms of IHD and the stable form of IHD. For example, it is acknowledged that during the period of acute presentation of IHD, there is a high level of residual platelet and thrombin activation, leading to a pro-thrombotic/coagulant coronary state, and that this would be significantly less once the disease has reached a more stable status following achievement of improved coronary flow from the combination of optimal medical and mechanical revascularization therapies [9-12]. The fact that patients in the present study, across all three treatment groups; medical therapy only, PCI with stents and CABG, benefitted from the ‘switchover’ would support achievement of a stable IHD status as being the main influence for this regardless of the supporting revascularization strategy. Admittedly, therefore, whilst the acute and unstable nature of the IHD at clinical presentation would benefit from the more aggressive and greater platelet inhibition, in the form of the newer P2Y12 antagonists, the stabilized IHD disease (after 3 months) would not necessarily require the same ‘more aggressive’ therapy. The clopidogrel-aspirin combination may therefore be sufficient, after 3 months, in maintaining adequate level of platelet inhibition. Our study data supports such a possibility without any concerns regarding patient safety i.e., adverse MACE outcomes from such a ‘switchover’.

Second, it is not uncommon to rationalize the need for more aggressive DAPT regime in patients following PCI, especially in view of an enhanced risk of ‘stenst thrombosis’. It is now widely acknowledged that in cases that re-present with ‘stenst thrombosis’ following PCI whilst on DAPT, it is often not just the underlying disease and the pro-coagulant state of blood flow, but also PCI-related factors that influence such re-admissions. Therefore, whilst the choice of more aggressive DAPT with newer P2Y12 antagonists would be an important adjuvant for such scenarios, the underlying issues of untreated residual disease, small edge dissections, possible inadvertent inadequate stent expansion and perhaps the use of extensive stent metal (long stents, stent overlap, bifurcation stenting etc.) also need addressing (more than the choice of switching to aggressive DAPT regime). Nevertheless, once the early period has lapsed, it might still be possible to switchover from a ‘more aggressive’ to the ‘less aggressive’ DAPT regime without jeopardizing
clinical outcome. Once again, our data implies that such a ‘switchover’ of DAPT therapy to maintain less intense platelet inhibition might not necessarily be unsafe after 3 months when the localized coronary injury site is well on its way to healing.

Third, in some cases, there is a greater need for tailoring the anti-platelet regime to the individual patient. For example, the very elderly (>80 years) and frail patients are often acknowledged to be at higher risk of bleeding. For this reason alone, DAPT with Prasugrel in the elderly (>75 years) and the frail (<60 kg weight) is not recommended. DAPT with ticagrelor, whilst not contraindicated in such high-risk patients, is often still used with caution. Likewise, the use of the newer P2Y12 inhibitors in patients on formal anti-coagulant regimes (Warfarin/New Oral Anticoagulants (NOACs) used in conditions such as systemic thrombosis, chronic atrial fibrillation, prosthetic valve disease etc.) remains unclear. Longer periods of more aggressive DAPT (ticagrelor-aspirin or Prasugral-aspirin) are often not considered in such cases. In addition, even after newer DAPT regimes have been initiated in some such cases, there is lack of consensus on the duration of such DAPT as well as whether it should be with or without formal anticoagulant use (i.e., triple therapy). Finally, the question of when to initiate the ‘switchover’ to less intense platelet inhibition in such cases on formal anti-coagulant regimes is yet to be answered. Often in cases where there is a need to continue or even resume formal anticoagulation, the choice is in favor of antiplatelet monotherapy, and rarely DAPT with newer P2Y12 inhibitors. More often, clopidogrel-aspirin is used, but again for a short duration of a few months only. The data from the present study, whilst not specifically looking at such high-risk patients, does however suggest the safety of such ‘switchover’ after a period of 3 months.

The optimal dosing and timing of switching after the last dose of ticagrelor to clopidogrel is unclear. It has been found that a loading dose of clopidogrel reduces the risk of High On-Treatment Platelet Reactivity (HTPR) with no increase in bleeding risk [13]. A waiting period of 24 h after the last dose of ticagrelor before loading with 600 mg of clopidogrel has been suggested in literature to allow sufficient time for ticagrelor and its metabolites to be eliminated as well as for new platelets to be exposed to the active metabolite of clopidogrel. Data in this area is very limited [14].

One year treatment with ticagrelor has been associated with an approximate 0.18 and 0.16 gain in Life-Years (LY) and Quality-Adjusted Life Years (QALY) respectively [15,16]. However, ticagrelor comes at a cost; with a 28-pack of ticagrelor costing £34.60 compared to a 30-pack of clopidogrel at £3.40. This additive cost to cash-strapped public services such as the National Health Service (NHS) in the UK is of particular importance. By switching over from ticagrelor to clopidogrel at 3 months, this results in a £1042.00 saving per patient per annum.

This study highlights the potential for safely switching from ticagrelor to clopidogrel at an interval of 3 months post ACS. This would be a more economically viable option in the current financially stretched institutes, which may in turn lead to the delivery of better healthcare as it would reduce the funding issues associated with ticagrelor, and enable more patients to receive it as treatment post ACS.

Limitations of Study

This is a non-randomized retrospective observational registry of a single center experience. The statistical data derived from such a small study sample population would not be without its limitations. However, it would be sufficient to rationalize a recommendation for a larger trial on the issue of ‘switchover’ of DAPT from the newer more expensive and aggressive therapies to the cheaper but older DAPT regime. A longer-term follow up (MACE outcomes) beyond 1 year would also be necessary to ensure long-term safety of a ‘switchover’ regime.

Conclusion

This study demonstrates the potential for safe switchover of DAPT to aspirin and clopidogrel following initial 3-month therapy with aspirin and ticagrelor independent of the revascularization treatment strategy. This would enhance cost-savings with no compromise in clinical care.

References


