Thrombocytopenia in a Patient undergoing Primary Percutaneous Coronary Intervention

by Yudi Her Oktaviono
Case report:
THROMBOCYTOPENIA IN A PATIENT UNDERGOING PRIMARY PERCUTANEOUS CORONARY INTERVENTION

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ABSTRACT

Thrombocytopenia is a common abnormality in patients presenting with acute coronary syndrome. Baseline thrombocytopenia in patients with ST-elevation myocardial infarction undergoing primary percutaneous coronary intervention is associated with early adverse events, related to both ischemia and bleeding. Treatment for acute coronary syndrome usually involves antiplatelet, anticoagulant, and antithrombotic therapy, and the performance of percutaneous coronary intervention. The safety of antiplatelet therapy and percutaneous coronary intervention patients who have acute coronary syndrome and thrombocytopenia is unknown, and there are no guidelines or randomized studies that specifically suggest a treatment approach in such patients. One of the institutions in Italy recommends medical and interventioninal strategy with radial as first choice for access site, bare metal stent (BMS) implanation, followed by double antiplatelet therapy (DAPT) for one month. After DAPT discontinuation, at least one antiplatelet drug (aspirin) is recommended for life.

Keywords: Thrombocytopenia; percutaneous coronary intervention; acute coronary syndrome

INTRODUCTION

Platelets have an important role in the pathogenesis of acute coronary syndrome (SKA). Thrombocytopenia, an abnormality that is sometimes found in people with SKA, is one of the risk factors for bleeding and other cardiac events in patients with SKA who are treated with antithrombotic (Hakim et al 2011). Standard therapy for acute coronary syndrome includes antiplatelet, anticoagulant, and thrombolytic agents, as well as percutaneous coronary intervention (PCI). All of these therapies are associated with a risk of bleeding and are usually delayed in patients with thrombocytopenia. The safety level of antiplatelet and PCI therapy in people with SKA with thrombocytopenia is unknown, and there are no randomized guidelines or studies that suggest a therapeutic approach in these patients (Hakim et al 2011). Here we report the case of a patient with acute coronary syndrome with thrombocytopenia performed by a primary PCI.

CASE REPORT

Mr. J, 52 years, went to the IRD Cardiac section with complaints of chest pain that penetrated his back 1 hour
before being hospitalized. Pain was felt when the patient
was washing the car. Complaints were accompanied by
palpations and cold sweat. The patient did not
comeplain of nausea and vomiting. Previous medical
history showed that he did not have hypertension and
stroke, but the patient had suffered from diabetes mellitus
for 10 years, erosive gastritis and liver cirrhosis for the
previous 3 years. From the physical examination, it was
found that the general condition was weak, awareness
was composed of blood pressure 100/60 pulse 95
x/minute breathing 24 x/minute temperature 36.5°C.

Fig. 1. Electrocardiographic examination.

Electrocardiographic examination at IRD showed a 95
x/minute sinus rhythm, normal frontal and horizontal
axis, acute anteroseptal myocardial infarction with
ventricular extracystole episodes, bigemini.

Normal head and neck physical examination, no
increase in JVP, thorax examination obtained a single
S1, S2 heart auscultation, no murmurs, gallops, and
extrastoles were obtained. Pulmonary auscultation
sounded vesicular breath in all lung fields, neither
rhonchi nor wheezing from examination of abdomen and
extremities within normal limits.

From history, physical examination, and electrocardio-
graphic, the diagnosis of acute myocardial ST-ante-
roseptal infarction was established. Initial therapy was
given, namely oxygen with nasal 3 liters per minute,
intravenous infusion and administration of 300 mg aspirin, 600 mg
clopidogrel, and 5 mg sublingual ISDN. Furthermore,
the patient was prepared for primary PCI.

Laboratory tests showed 9.5 g/dL Hb, 8,560
leukocytes/μL, 79,000/μL platelets, 9 mg BUN/dL,
serum creatinine 0.77 mg/dL, glucose 192 mg/dL,
albumin 2.74 g/dL, SGOT 62 U/L, SGPT 27 U/L,
potassium 3.7 mmol/L, sodium 141 mmol/L, chloride
111 mmol/L, APTT 20.7 seconds with control 26.0
seconds, PTT 15.4 seconds with control 12 seconds.
Increased cardiac enzymes, namely CKMB 171 U/L,
and Troponin-I 0.096 ng/mL.

Examination of the chest x-ray shows less inspiring
results, with a large heart impression normal, and
pulmonary congestion.

Fig. 2. Examination of chest x-rays.

Furthermore, primary PCI actions with angiographic
results, namely Left main coronary artery, are
performed. The left anterior descending artery appears
diffuse disease from the proximal-distal LAD with a
maximum stenosis of 95% at the proximal LAD. The
non-dominant Left circumflex artery shows a significant
70% stenosis in the distal part. The right coronary artery
is dominant, there is a significant 80% stenosis in the
ostial, a significant 80% stenosis in the mid section.

After angiography is complete, proceed with thrombus
aspiration, but no thrombus is obtained. Then stenting
DES Firebird II (Rapamycin) 3.0 x 18 mm at the
proximal LAD directly stenting. PCI is done with good
results.

During the PPCI action, it was found that hypotension
improved after dopamine pump was given 5
μg/kg/minute, and the patient complained of nausea
and then was given Ondansetron 1 ampoule iv. After the
action there were no complaints felt by the patient,
physical examination showed the presence of fine wet
rhonchi on the basal of both lung fields which were then
given a 2 ampoule furosemide bolus followed by a
pump of 5 mg/hour. The next therapy is Aspirin 1 x 100
mg, Clopidogrel 1 x 75 mg, Simvastatin 1 x 20 mg,
ISDN 3 x 5 mg, Captopril 3 x 12.5 mg, Spironolactone
1 x 25 mg. Ranitidine 2 x 1 ampoule iv, KSR 3 x 1 tab,
Lactulose 2 x 15 cc syrup, 3 x 15 cc Sucralfate syrup.

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Fig. 3. Coronary angiography.

The patient was discharged without complaint with Aspirin 1 x 100 mg, Clopidogrel 1 x 75 mg, ISDN 3 x 5 mg, Simvastatin 1 x 20 mg, Captopril 3 x 12.5 mg, Bisoprolol 1 x 2.5 mg, Furosemide 1 x 20 mg, Spironolactone 1 x 25 mg, Lansoprazole 1 x 30 mg.

Fig. 4. After installation of stents.

The patient is consulted with the internal medicine department. Patients complain of sometimes nausea. From the examination it was found that patients with anemia (Hb 9.5 g/dL), thrombocytopenia (71,600 g/dL), hypoalbuminemia (albumin 2.74 g/dL). The patient has performed a previous endoscopic examination with grade I-II esophageal varices results, and abdominal ultrasound examination with the results of splenomegaly, parenchymal liver disease, cholelithiasis, and ascites. Conclusion of diagnoses from internal medicine, namely cirrhosis of the liver, esophageal varices of grade I-II, and controlled diabetes mellitus. Therapy given is Diet H1 1900 kcal/day, injection of Omeprozole 2 x 40 mg, Lactulose syrup 2 x 15 cc, Sucralfate syrup 3 x 15 cc, and monitor for signs of bleeding.

During hospital treatment, patients have no complaints of chest pain, tightness, or palpitations. The patient also did not experience bleeding. Physical examination found a blood pressure of 120/70, pulse 84 x/minute, breathing 16 x/minute. Thoracic examination or wheezing were not obtained. Electrocardiography shows a sinus rhythm of 85 x/minute, normal axis, and pathological Q waves are obtained in V1-V3 stumbling block.

Fig. 5. Electrocardiographic examination during treatment.

**DISCUSSION**

Acute STEMI myocardial infarction is a clinical syndrome defined by the symptoms of myocardial ischemia associated with persistent features of electrocardiography (ECG) namely ST segment elevation and an increase in biomarkers as a result of myocardial necrosis (O’Gara et al 2013). Patients have typical chest pain, with risk factors for diabetes mellitus. An ECG examination shows ST segment elevation in the V1-V3 stain which shows an anterosetal STEMI.
There are two therapeutic concepts for patients with STEMI, namely PCI and thrombolitics. Primary PCI has become the main choice in the act of reperfusion in STEMI. Compared with thrombolytic therapy in randomized clinical trials, PPCI produced higher patency rates in arteries related to infarction, Thrombolysis in Myocardial Infarction (TIMI) flow grade 3, and lower rates of recurrent ischemia, recurrent infarction, recurrent emergency revascularization procedures, intracranial bleeding, and death (Keeley et al 2003). In these patients, primary PCI therapy is used.

Patients with acute myocardial infarction can be accompanied by other abnormalities that can affect the management of acute coronary syndrome. In this case, the patient is also accompanied by hepatic cirrhosis which results in thrombocytopenia. Treatment of acute coronary syndrome includes antiplatelet, anticoagulant, and PCI where these therapies are associated with the risk of bleeding which is generally delayed in patients with thrombocytopenia. The safety of antiplatelet and PCI therapy in patients with SKA and thrombocytopenia is still unknown, there are no guidelines or randomized studies that suggest a therapeutic approach for these patients (Yusuf et al 2010).

**Thrombocytopenia in patients who have primary PCI**

Thrombocytopenia results from abnormal production, platelet distribution, or increased platelet destruction due to hereditary or acquired disease (Overgaard et al 2008). Severe thrombocytopenia (<50 x 10^9/L) is a finding that is rarely found especially in patients with coronary heart disease (Campo et al 2012). Thrombocytopenia is known to be significantly associated with the occurrence of cardiovascular events, major bleeding, and cardiovascular mortality significantly in 30 days in patients with SKA who underwent primary PCI (Judge DA, et al., 2011). In general, the risk of bleeding is still low if the platelet count is still around 50 x 10^9/L, but this risk increases sharply if platelets reach 10-20 x 10^9/L (Towse et al 2009). The risk of bleeding in patients with thrombocytopenia varies, depending on the underlying disease. The most common causes of thrombocytopenia are malignancy, liver disease, and immune-related diseases (Spoon et al 2013). For example, thrombocytopenia caused by chemotherapy, spontaneous bleeding is only found in patients with platelet counts below 10 x 10^9/L (Goldberg et al 1994). In this case, the cause of thrombocytopenia is hepatic cirrhosis. There are several factors in liver cirrhosis which result in thrombocytopenia, including splenic sequestration, decreased thrombopoietic activity, bone marrow suppression due to chronic hepatitis C infection, antiviral therapy with interferon (Hayashi et al 2014).

The cause of coronary thrombosis in patients with thrombocytopenia is still debatable and multifactorial. Even so, from the results of the autopsy the fibrin-platelet thrombus is the same as the thrombus found in the atherosclerosis process. Patients may predispose to the occurrence of coronary thrombosis because of the larger and more adhesive platelet shape to the vascular surface. In patients with acute myocardial infarction or stroke, the average volume of platelets is significantly increased even though the platelet count decreases. Due to natural regulation of platelet production to maintain platelet mass (platelet count and platelet volume on average), it can be concluded that larger platelets not only cause blockage of hemostasis and prevent bleeding due to high activity, but may also be associated with coronary thrombosis in patients with thrombocytopenia (Yusuf et al 2010).

**Therapy management**

The safety and tolerability of anticoagulants and antiplatelets have been well described in the general population of SKA, but few data have shown the safety and tolerability of these therapies in patients with thrombocytopenia. Patients with thrombocytopenia undergoing PCI have been shown to undergo longer hospitalizations, higher mortality rates when hospitalized, higher rates of bleeding (Overgaard et al 2008). Bleeding is the most common side effect of therapy with anticoagulants and antiplatelets. Therefore, the benefits and risks of giving this therapy must be considered well in order to achieve optimal outcomes and the possibility of minimal bleeding in patients with thrombocytopenia (Yusuf et al 2010).

Aspirin remains an important antiplatelet therapy for patients with SKA. Aspirin therapy in thrombocytopenia is still being debated, there is no widely accepted protocol. One study of people with SKA with cancer-induced thrombocytopenia showed that aspirin was associated with a significant increase in seven-day survival of SKA in patients with SKA with cancer, with or without thrombocytopenia, and not associated with more severe bleeding (Sarkis et al 2007). Aspirin has a short plasma half-life, but has a long biological effect due to irreversible platelet inhibition, as well as P2Y12 antagonists. Because there is no specific reversal agent, the treatment or prevention of bleeding is by stopping antiplatelet therapy. When antiplatelet drugs are stopped, as soon as the function of hemostasis is stable, antiplatelet must be given back immediately. If life-threatening bleeding is found, platelet transfusion can be considered for the reversal of the antiplatelet effect, but
the risk of arterial thrombosis is necessary (Makris et al 2012).

The American Heart Association recommends that patients with DES stents be implanted and have a high risk of bleeding so P2Y12 antagonist therapy can be stopped after 6 months of administration (recommendation IIb) (Levine et al 2016). The use of bare metal stents (BMS) is recommended for patients with a high risk of bleeding, who cannot take dual antiplatelet therapy for 12 months. Because the risk of stent thrombosis with BMS is greatest in the first 14 to 30 days, dual antiplatelet therapy is at least 30 days after implantation with BMS (Levine et al 2011).

The study conducted by G. Campo and colleagues showed that there were 35 patients with severe thrombocytopenia who underwent PCI. The use of drug-eluting stent (DES) is sometimes still used for certain cases such as stenosis on the left playing artery, overlap on the left anterior descending artery, or small blood vessels. Antiplatelets commonly used are unfractionated heparin (UFH), only 3 people use bivalirudin. There were 20 patients who were recommended for treatment with dual antiplatelet for 6 months, but only 5 patients could meet the target. Nearly half of the study population experienced bleeding complications, the majority of which were superficial bleeding, namely petechiae and ecchymosis. When bleeding occurs, antiplatelet drugs are temporarily delayed. Based on this study, recommendations for medical strategies and interventions, namely first choice arterial access, namely the radial artery, BMS implantation, followed by dual antiplatelet use for 1 month. After that, a minimum of 1 antiplatelet (aspirin 75 mg) is recommended for life. Periprocedural antiplatelets are operator dependent, and the use of Glycoprotein (GP) IIb-IIIa inhibitors is not recommended. Research conducted by Darcy and colleagues showed that patients with thrombocyopenia (<100x10^9/L) who were subsequently performed for coronary angiography had a moderate or large hematoma nine times more often than patients with platelet counts above that value. To minimize this complication, radial arterial access is recommended. In addition, they also stated that being careful in providing therapy, monitoring and follow-up properly can reduce PCI-related bleeding events and can improve compliance with antiplatelet (Darcy et al 1996, Campo et al 2012).

SUMMARY

Patients with acute myocardial infarction can be accompanied by other abnormalities that can affect the management of acute coronary syndrome. Thrombocytopenia is known to be significantly associated with significant cardiovascular events, major bleeding, and cardiovascular mortality in 30 days in patients with SCA who underwent primary PCI. There is still little data that shows the safety and tolerability of antiplatelet and antiplatelet therapy in patients with thrombocyopenia. One recommendation from one institution is for medical strategies and interventions, namely first choice arterial access, namely the radial artery, BMS implantation, followed by dual antiplatelet use for 1 month. After that, a minimum of 1 antiplatelet (aspirin 75 mg) is recommended for life. Periprocedural antiplatelets are operator dependent, and the use of Glycoprotein (GP) IIb-IIIa inhibitors is not recommended.

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