Coronary Artery Dissection and Perforation Complicating Percutaneous Coronary Intervention – A Review

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Abstract:
After its introduction by Andrew R. Gruentzig in 1977 percutaneous coronary intervention (PCI) is widely utilized in the treatment of symptomatic coronary artery disease. Though it has numerous benefits, serious and potentially life-threatening complications of PCI can occur, including iatrogenic coronary artery dissection and perforation. The incidence of these complications has been augmented by the development of coronary interventional devices intended to remove or ablate tissue. Here we review the classification, incidence, pathogenesis, clinical sequelae and management of coronary artery dissection and perforation in the current era due to PCI. Specifically, the current angiographic classifications of coronary artery dissections and perforations are reviewed. The findings of several recent, registries of PCI-related coronary artery perforations and dissection are summarized. The management of coronary artery dissection and perforation is discussed in details, including the application of newer modalities such as covered stents.

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Keywords: Percutaneous coronary intervention (PCI), Coronary artery dissection.

Introduction:
Percutaneous coronary intervention (PCI) is the most popular and increasingly utilized in the treatment of coronary artery disease. PCI has numerous benefits. Serious and potentially life-threatening complications of PCI can also occur, including iatrogenic coronary artery dissection and perforation. Up to 30% of all conventional balloon angioplasties result in angiographically significant coronary artery dissection.1,2 In recent studies, perforation has been reported to occur in 0.3–0.6% of all patients undergoing PCI.3,4 The incidence of these complications was high in Stenting with predilatation but subsequent studies show no differences between these two procedures. Perforations has been augmented by the development of coronary interventional devices intended to remove or ablate tissue, such as transluminal extraction coronary atherectomy, directional coronary atherectomy, excimer laser coronary angioplasty and high-speed mechanical rotational atherectomy. Here we review the incidence, causes, clinical sequelae and management of coronary artery dissection and perforation in the current era.7

Coronary Artery Dissection:
Percutaneous coronary intervention, which depends upon mechanical dilatation of the artery or ablation of atherosclerotic plaque, is requisitely associated with plaque fracture, intimal splitting and localized medial dissection—these tears may extend into the media for varying distances, and may even extend through the adventitia resulting in frank perforation.

Incidence of coronary artery dissection:
Up to 30% of all conventional balloon angioplasties result in angiographically significant coronary artery dissection.5,6 But all dissections are not clinically significant.

Type of dissection:
The National Heart, Lung and Blood Institute (NHLBI) classification system for intimal tears, developed by the Coronary Angioplasty Registry, was put forth in the pre-stent era for the classification of dissection types after balloon angioplasty. Dissections in this scheme are graded based upon their angiographic appearances as types A through F.12 (Figure 1).

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Type A dissections represent minor radiolucent areas within the coronary lumen during contrast injection with little or no persistence of contrast after the dye has cleared.

Type B dissections are parallel tracts or a double lumen separated by a radiolucent area during contrast injection, with minimal or no persistence after dye clearance.

Type C dissections appear as contrast outside the coronary lumen (“extraluminal cap”) with persistence of contrast after dye has cleared from the lumen.

Type D dissections represent spiral (“barber shop pole”) luminal filling defects, frequently with excessive contrast staining of the dissected false lumen.

Type E dissections appear as new, persistent filling defects within the coronary lumen.

Type F dissections represent those that lead to total occlusion of the coronary lumen without distal antegrade flow. In rare cases, a coronary artery dissection may propagate retrograde and involve the ascending aorta.13

Coronary Artery Perforation:
Coronary perforation occurs when a dissection or intimal tear propagates outward sufficient to completely penetrate the arterial wall. A significant risk factor for perforation during PTCA is the balloon to artery ratio. In the report by Ajluni et al, balloon perforations occurred from a measured balloon to artery ratio of 1.3 ± 0.3, which was significantly larger than the balloon to artery ratio of 1.0 ± 0.3 for other lesions in which perforation did not occur (p < 0.001).3

Similarly, in a registry by Ellis et al, the balloon to artery ratio of those patients undergoing PTCA complicated by perforation was 1.19 ± 0.17 versus 0.92 ± 0.16 for those without perforation (p = 0.03).4 This observation has been confirmed in another large randomized study in which a balloon to artery ratio > 1.1 was shown to result in a 2–3 fold increase in severe dissection leading to abrupt closure compared to a balloon/artery ratio < 1.1.14 In addition, balloon rupture, particularly those associated with pinhole leaks (as opposed to longitudinal tears), may create high pressure jets that increase the risk of dissection or perforation.(Figure 2)

Coronary Artery Dissection and Perforation Complicating Z Rahman et al.

Fig.-1: National Heart, Lung and Blood Institute (NHLBI) classification system for coronary artery dissection types. Types A and B are generally clinically benign, whereas types C through F have clinically significant morbidity and mortality if untreated.

Fig.-2: Coronary artery dissection as a result of balloon perforation. (A) Initial cutting balloon inflation for treatment of diffuse in-stent restenosis of the right coronary artery. Examination of the cutting balloon ex vivo revealed a pinhole perforation. This dissection was treated with prolonged balloon inflation with complete sealing of the intimal entry site.
Incidence of coronary perforation:
In agreement with previous reports with the use of non-debulking procedures, one study demonstrated that the incidence of coronary perforation was 0.1%.\textsuperscript{15,16} Since new interventional methods are currently used to overcome the limitations of conventional PCI along with the adjunctive use of GP IIb/IIIa inhibitors, the risk of coronary perforation is reported to be much higher.\textsuperscript{17-20} The wide range of reported incidence of perforations may be attributable to different definitions of vessels perforations, the clinical spectrum of cases, the prevalence of chronic coronary occlusion interventions and the frequency of aggressive debulking use in each institution.

Causes:
The development of debulking devices which cut, vaporize, or drill the vessel wall, such as directional coronary atherectomy (DCA), excimer laser angioplasty and rotational or extraction atherectomy devices may be associated with the increase of perforations (0.5—3%).\textsuperscript{21,22} Again, the use of devices intended to remove or ablate tissue were associated with higher perforation rates than PTCA alone. Two recent registries of cutting balloon angioplasty report a low incidence of perforation with this device.\textsuperscript{23,24} Additionally, the use of the recently developed hydrophilic and heavy-weight guidewires has increased the frequency of this complication.\textsuperscript{25} However, guidewire-related perforations can occur by the distal wire itself or by a fractured guidewire. The use of GP IIb/IIIa inhibitors confers no significantly increased risk of perforation or tamponade, once these complications occur but they are related to a higher risk of fatal bleeding.\textsuperscript{6} Risk increases with complex lesion morphology such as chronic total occlusions (especially long standing with bridging collaterals), angulated calcific lesions, tortuous vessels, bifurcation or ostial lesions and eccentric or long lesions (>10 mm).

The characteristics of specific stents (i.e. stiff stents, requiring high pressure inflation for sufficient expansion) increase the risk of perforation considerably even though these stents is preferable in lesions where the possibility of restenosis is considered to be high.\textsuperscript{26} Specific PCI techniques such as kissing balloons and high pressure inflation increase the risk of perforation.

Type of perforation:
Ellis et al evaluated a novel angiographic classification scheme for coronary artery perforations as a predictor of outcome.\textsuperscript{14} In a multicenter registry of 12,900 PCIs, 62 (0.5%) perforations were reported and categorized as:
Type I, extraluminal crater without extravasation;
Type II, epicardial fat or myocardial blush without contrast jet extravasation;
Type III, extravasation through frank (> 1 mm) perforation; or Type III “cavity spilling” (CS) referring to Type III perforations with contrast spilling directly into either the left ventricle, coronary sinus or other anatomic circulatory chamber. (Figures 3 and 4)

Fig.-3: Ellis classification scheme for coronary artery perforations. Types I and II are “contained” perforations, whereas Type III is a “free” perforation with continuous extravasation of contrast. A more benign Type III “cavity spilling” perforation is also described in which contrast spills directly into an anatomic circulatory chamber.
Interestingly, the Ellis Type I perforation is angiographically identical to the previously described NHBLI Type C dissection, reinforcing the notion that a continuum exists between dissection and perforation.

In another large retrospective analysis, Ajluni et al. reviewed 8,932 PCIs in which coronary artery perforation was reported in 35 (0.4%). Perforations were classified angiographically as a: 1) free perforation, defined as free contrast extravasation into the pericardium (Ellis Type III); or 2) contained perforation, defined as a contained extraluminal blush or localized rounded crater of contrast extending outside the contrast-filled vascular lumen (Ellis Type I or II).

Clinical outcome after perforation:
The frequency of adverse clinical outcomes relates to the severity of the coronary perforation. Perforations with cardiac tamponade are closely associated with mortality. In cases of coronary perforation, myocardial infarction rates occur in 16.7-50%, emergency surgery in 50% and death results in 9-19%, despite more aggressive treatment. Clinical follow up for more than a year after the procedure demonstrated that clinical events were relatively uncommon, including only one patient requiring surgery. Limited studies have reported the occurrence of late pseudoaneurysm formation following coronary perforation most commonly after DCA and cutting balloon use.

Concurrent with prior reports, Ajluni et al. found that overall clinical outcomes were worse for patients with free perforations (tamponade 20%, CABG 60%, death 20%) than with contained perforations (tamponade 6%, CABG 24%, death 6%). The use of devices intended to remove or ablate tissue were associated with higher perforation rates than PTCA alone (1.3% vs. 0.1%). Women and the elderly were also at increased relative risk. The investigators reported that Type I perforations were associated with no deaths or myocardial infarction, and tamponade in 8%. Type II perforations, when treated with a prolonged balloon inflation, resulted in no deaths and a low incidence of adverse sequelae (myocardial infarction in 14%, tamponade in 13%). Type III perforations were associated with the rapid development of cardiac tamponade (63%), the need for urgent bypass surgery (63%) and a high mortality (19%). Type III “cavity spilling” perforations, however, were associated with less catastrophic consequences (no deaths, myocardial infarction or tamponade resulted).

Of 2991 conventional PCI procedures performed in our department, there were only 5 cases of coronary perforation (0.1%). Guidewire perforations constituted 20% of all perforations and 80% were caused during or after stent implantation. It is important to note that most cases (80%) were managed with prolonged balloon occlusion. Haemopericardium occurred in 60% of patients with coronary perforation and pericardial drainage was performed in 2 cases because of haemodynamic instability. There was one procedural-related death in these series.

Spontaneous coronary artery dissection:
Spontaneous coronary artery dissection (SCAD) is a rare condition that usually occurs in relatively young patients who are predominantly female. Seldom could it be a cause of acute myocardial ischemia leading to a sudden cardiac death. SCAD...
consists of intramural hematoma formation or, rarely, intimal tears that initiate and propagate the dissection in the vessel wall. In rare cases, the SCAD occurs in male patients. Many strategies could be considered in patients with SCAD, such as PCI, bypass surgery, or conservative medical management. In general, the long-term prognosis of patients with SCAD is considered favorable if they survive the acute phase.

**Risk factors for spontaneous coronary artery dissection:**
The most frequent were pregnancy or postpartum state, coronary artery disease, and exercise. Less common risk factors included Marfan’s syndrome, hypersensitivity vasculitis, Kawasaki’s disease, and alpha-1-antitrypsin deficiency oral contraceptives, and hypertension.

Coronary artery perforation has been reported to be more frequent in elderly female patients, probably due to their small vasculature.

The incidence of spontaneous coronary dissection occurs at rates of 0.1–0.28% of all angiographic studies. Blunt trauma, cocaine use, and extension of aortic dissection have also been reported to result in coronary dissection.

In general, spontaneous coronary dissection is a life-threatening condition; however, in those surviving the initial event, the survival rate is reported to be 78%–82%.

**Diagnosis:**
Coronary perforation can be easily diagnosed by coronary angiography and echocardiography and it is usually accompanied by new episodes of chest pain, haemodynamic deterioration and electrocardiographic changes.

Angiographic evidences of perforation are the presence of blush, jet, coronary sinus compression and contrast in pericardium. The most difficult type of diagnosis is when there is extramural blood collection on the vessel wall, which can only be detected indirectly from coronary angiography. Of interest, there is no angiographic evidence of perforation in 10 to 20%. Delayed tamponade can occur several hours to several days after angioplasty and is common in perforations induced by guidewire or GP IIb/IIIa. Thus this diagnosis should remain high on the list of the differential diagnosis of post-PCI hypotension.

MRI can demonstrate acute myocardial infarction due to posttraumatic coronary artery dissection.

**Prevention and management of coronary perforation:**
Since the occurrence of perforation is highly associated with the selection of an appropriate device for the lesion, the morphologic assessment of the lesions by intravascular ultrasound (IVUS) would be very helpful. IVUS can provide not only the location of calcification but also the severity, the depth and the eccentricity of calcification, which are thought to influence the risk of perforation.

Meticulous attention to guidewire position, to device sizing and to the ‘blind high pressure technique’ is required since they are all highly related to the risk of coronary perforation. Another cause of cardiac perforation occurs as a consequence of the use of a temporary pacing wire, of which interventional cardiologists should always be aware. So careful manipulation of pacing wire can prevent these devastating complications. Early management is crucial for plugging the perforation without resorting to surgery. If the perforation site is visible, the blood flow of the perforated branch should be blocked immediately by the plain balloon, which was used for the procedure. Balloon inflation should be performed at the lowest possible pressure to promote haemostasis, generally less than 2 atm and should last for 5 to 15 min. If myocardial ischaemia develops, this should be exchanged by a perfusion balloon, which enables the blood to perfuse to the distal portion of the vessel. Prolong balloon inflation may avoid surgery in 60-70% of the cases. In the present series, prolonged balloon inflations were used in 80% of patients and they were successful in sealing the coronary perforations in all but 1 patient. (Figure 5 and 6)

In the meantime, if cardiac tamponade and low blood pressure ensue, pericardiotentesis should be performed immediately. In a study, a case of delayed pericardial haemorrhage occurred in a patient of perforation type I, which did not require pericardiotentesis. Two out of three cases of
patients with perforation type III developed cardiac tamponade, of which one required pericardiocentesis. Administration of fluids could prevent cardiac tamponade and hypovolemic shock.

For haemostasis, protamine should be administered to negate the effects of heparin and in a dosage to achieve a partial thromboplastin time of less than 60 seconds or an ACT of less than 150 second.48,49 The administration of protamine seems to be safe with no vessel thrombosis complications. Platelet transfusion is useful in patients treated with abciximab, but not with tirofiban or eptifibatide 50

Delivery of covered stents (autologous vein cover or polytetrafluoroethylene (PTFE) covered stents) may be attempted as an optional treatment before resorting to emergency surgery, especially for perforation sites located in the proximal or mid-portion of the index vessel.51-53 However, preparing these stents is technically difficult, time-consuming and the restenosis rates of these devices are known to be high.

If a large perforation is causing serious ischaemia, the guidewire cannot be re-crossed or the bleeding continuous, emergency surgery is the only option. Coronary perforation accounts for 20% of referrals for urgent surgery.54 If the situation allows, this should be done with a perfusion balloon in place and dilated at low pressure.

Embolization of gel foam or thrombogenic metallic (stainless steel/platinum) coils into the leaking vessels is another treatment option when perforation persists.55 If the perforation is too distal, or in a small vessel or it is near to a chronic total occlusion or surgery is not possible, there is always the possibility of coil embolization onto the site through an infusion catheter. However, this technique is complicated, requiring a special delivery system and adds to the cost of the procedure.

Treatment options for SCAD have increased significantly, roughly paralleling the development of coronary angiography. These include medical therapy, percutaneous intervention, and coronary bypass. It is not possible to define the optimal treatment from in our review. Of all the treatment options, no one option stands out as superior to the others. Our observations suggest that no matter what treatment is chosen, more than 95% of patients are likely to survive if the patient survives to the time of diagnosis.

**Fig.-5:** Type II perforation. (A) A calcified lesion of the mid-LAD (B) A conventional balloon angioplasty was performed. (C) After several balloon inflations, epicardial staining by contrast extravasation occurred. (D) Extravasation was terminated after prolonged balloon inflation.

**Fig.-6:** Type III perforation. (A) A calcified lesion of the mid-LCX. (B) A conventional balloon angioplasty was performed. (C) After several balloon inflations, epicardial staining by contrast extravasation occurred. (D) A long inflation of the balloon at a very low pressure was performed. (E) Extravasation was terminated after prolonged balloon inflation.

**Clinical outcome after perforation:**
The frequency of adverse clinical outcomes relates to the severity of the coronary perforation. Perforations with cardiac tamponade are closely associated with mortality.56,57 In cases of coronary perforation, despite more aggressive treatment,
myocardial infarction rates occur in 16.7-50%, emergency surgery in 50% and death results in 9-19%. Clinical follow up for more than a year after the procedure demonstrated that clinical events were relatively uncommon, including only one patient requiring surgery. Limited studies have reported the occurrence of late pseudoaneurysm formation following coronary perforation most commonly after DCA and cutting balloon use.57

Conclusion:
Coronary artery dissections and perforations remain common complications during PCI but clinical sequelae have been minimized by early diagnosis and routine use of coronary stents. Perforation, although rare, can often be a life-threatening complication. Coronary perforation remains a dreadful complication but with low incidence in the current interventional era. Prevention is the best treatment of this complication.58

Early and spot recognition and attention to the angiographic appearance of the perforation is essential for the successful management of this complication. Those perforations which are “contained” angiographically carry a more benign prognosis than those which are freely extravagating contrast into the pericardium. Treatment should be aimed at sealing the perforation with low pressure prolonged conventional or perfusion balloon inflation, prudent reversal of anticoagulation and use of covered stents. Echocardiography should be performed in all cases of coronary perforation and urgent pericardiocentesis if tamponade develops. In cases where sealing of the perforation by conservative measures cannot be achieved, emergency bypass surgery must be performed.59

References:


