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An Infected Drug-Eluting Stented Coronary Aneurysm Forming Intracardiac Fistula

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1. Introduction

Many cardiologists have usually operated percutaneous coronary intervention (PCI) for coronary artery disease practically. Sometimes, complications following stent implantation rarely include thrombosis, rupture, sepsis and infected coronary aneurysm [Baddour, 2004; Berkalp, 1999]. Particularly, infected coronary aneurysm causes fatal outcome. In addition to contamination at the time of catheterization, there are other mechanisms how stents can become infected, including transient bacteremia from skin flora via access-site hematomas, pseudoaneurysms, delayed bleeding, prolonged arterial sheath insertion, and several procedures performed from the same access site over a short time period [Baddour, 2004]. Previous report demonstrated that 18% of patients who underwent complex PCIs had detectable bacteremia [Ramsdale, 2004]. However, prophylactic antibiotics administrations are not conducted routinely prior to coronary stenting in the current PCI procedures, because the incident rate of stent infections was reported to be less than one in 10,000 cases [Myles, 2000].

The drug-eluting stent (DES) era was ushered in with the first published human study by Sousa et al. in 2001, showing a nearly complete abolition of neointimal hyperplasia by use of sirolimus-eluting stent (SES) [Sousa, 2001]. The role of DES is very questionable, because of long term anticoagulant therapy, but in the patients with comorbidity (diabetes mellitus) DES are recommend. Despite the dramatic capacity of SES to reduce the restenosis rate after PCI, several SES-related problems have been raised [Ong & Serruys, 2005]; 1) the requirement of a prolonged dual anti-platelet regimen to avoid the risk of DES thrombosis [McFadden, 2004], 2) the occurrence of acquired late malapposition 3) the late formation of coronary artery aneurysm (SES having an antiproliferative action may be responsible for the delayed and inappropriate healing, which should lead to weakening of the arterial wall and delayed aneurysm formation) [Abreu, 2005; Degertekin, 2003], 4) a severe localized hypersensitivity consisting predominantly of T lymphocytes and eosinophils, which was caused by the metallic stent, polymer, or sirolimus [Virmani, 2004; Nebeker, 2005; Stabile, 2004].

2. Case report

A 70-year-old man with hypertension, type 2 diabetes mellitus, mild renal dysfunction and old cerebral infarction, was admitted to our hospital with sustained angina. The patient
suffered an inferior wall ST-depression unstable angina that required immediate PCI with two SESs (Cypher®, Cordis Johnson & Johnson, Japan) in the proximal right coronary artery (RCA) (Figure 1). Satisfactory angiographic results were obtained with smooth luminal outline, and then the arterial sheath was removed (Figure 1). The patient had other significant stenosis lesions in the proximal left anterior descending artery (LAD) and the middle circumflex artery (LCX).

Next day after the procedure, this patient had fever without suppuration on the access site, and grew *Staphylococcus aureus* on repeat blood cultures. Despite exhaustive screenings for a potential source of the infection, no infectious focus was detected. The patient was treated with intravenous adapted antibiotics, however, remained poor.

Three weeks after PCI, the patient had fever, convulsion and syncope attack with the torsades de pointes in ECG. The causes were considered due to trouble of the residual left coronary arteries (LCAs) lesions, and the patient was successfully implanted two SESs in the target LAD stenosis and a SES in the target LCX stenosis. The postprocedural course was uneventful, and the inflammatory signs were improved by the treatment with adapted antibiotics.

Two months later, the patient has recurrence fever with eosinophilia and an inferolateral wall ST-depression angina. Coronary angiography revealed no obvious changes in the LCAs, but an occluded proximal RCA stent, a large aneurysm off the stent, and a fistula into the right ventricular chamber with rapid clearance (Figure 2). A 16-slice multidetector computed tomographic angiogram scan confirmed both an occluded proximal RCA, and the aneurysm (50x30 mm in size) forming fistula (10 mm in size) into right ventricle (Figure 3). The patient underwent a resection of the RCA stents and aneurysm, and a reconstruction of the right ventricular wall without coronary bypass grafting. Microscopic specimen from the resected aneurysmal wall revealed an extensive inflammatory reaction with a predominance of neutrophils consisten with the micro-abscessn (Figure 4). The patient is doing well 5 years after operation [Kishida, 2007].
Angiography of the RCA demonstrated occlusion of proximal stent site, an entry point to aneurysm, and the saccular aneurysm in the area of proximal stents and its coronary artery fistula originating from the RCA stents, with contrast spillage into the right ventricular chamber through the fistula. Rapid clearance of contrast was observed, with none in the pericardial space [Reproduced Kishida, 2007].

Fig. 3. The 16-slice multidetector computed tomographic angiogram disclosed huge RCA aneurysm (10x10 mm in diameter) and draining image into right ventricular chamber via fistula [Reproduced Kishida, 2007].

Fig. 4. Histopathology from pseudoaneurysm after surgical excision. Its histopathologic examination revealed marked inflammatory reaction with a predominance of neutrophils consistent with an abscess. Hematoxylin and eosin staining. (original magnification, x200) [Reproduced Kishida, 2007].
2.1 Comments for case report

To date, there have been only 7 reported cases of infected drug-eluting stented coronary aneurysm, including our case [Kishida, 2007; Marcu, 2005; Alfonso, 2006; Singh, 2005; Jang, 2007; Le, 2007; Furutado, 2011] (Table 1). In six cases Staphylococcus aureus and one Pseudomonas aeruginosa bacteremia was responsible for causing mycotic stent complications. Of these cases, there were two cases of an infected drug-eluting stented coronary aneurysm forming fistula to cardiac chamber. The mechanism of infection at the site of drug-eluting stenting is not well understood. Potential causes for drug-eluting coronary stent infections include local suppression of immune response and endothelialization which should lead to weakening of the arterial wall and delayed aneurysm formation [Degertekin, 2003]. The potential role of SES in locally blunting the innate response to bacterial agents may be considered. Formation of mycotic aneurysmal fistula in our case may be partly accounted for by this mechanism.

<table>
<thead>
<tr>
<th>Case</th>
<th>Vessel</th>
<th>Stent</th>
<th>Fistula</th>
<th>Organism</th>
<th>Days</th>
<th>Max size</th>
<th>Ope</th>
<th>Outcome</th>
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<tr>
<td>55yo</td>
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<tr>
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<td>+</td>
<td>Alive</td>
<td>2011 Furutado</td>
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Table 1. Infected drug-eluting stented coronary aneurysm

3. Conclusion

Infections specificity, related to the use of intracoronary DESs, is exceedingly rare. However, stent infection should be considered in the differential diagnosis of patients presenting unexplained fever and relapsing bacteremia at any time following drug-eluting stenting.

4. References


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In this book we examined a periprocedural complication of coronary angiography, and coronary intervention. That includes related to cardiac catheterization and diagnostic coronary angiography, and those that occur as a consequence of the specific equipment. However, improvements in devices, the use of stents, and aggressive antiplatelet therapy have significantly reduced the incidence of major periprocedural complications. This book giving knowledge and experiences many of interventional cardiologists from all over the world, and provide possibility to recognize new approach in this domain. Book gives lecture on how we image and how we decide on what to treat, how to treat it, and then results of that treatment. They offer many answers to what we have today and what we will have tomorrow.

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