Percutaneous Coronary Interventions over the last three decades: the adjunctive devices (role, development and perspectives)

Leo Finci

Abstract
Percutaneous coronary interventions (PCI) performed on large-scale basis in routine practice with documented long-term clinical benefit for the patient are balloon angioplasty introduced in 1977 and coronary stenting since 1986. Other coronary devices, so-called “adjunctive”, have triggered a great admiration at the time of its initial clinical application but showed to be less successful at the long-term follow-up. However, many adjunctive devices are still necessary in 1-2% of all PCI procedures for complex cases. This article is a short, comprehensive review retracing the conception and destiny of these devices. Coronary balloon catheters, a standard device with advance “therapeutic” features comprise: cutting, drug eluting and high pressure non-compliant balloon catheters. Atherectomy devices can be directional for lesions “debulking”, rotational (Rotablator) for calcified, long lesions and recently Orbital supposed to have less no-reflow phenomenon. Excimer Laser angioplasty for treatment of “uncrossable” lesion was claimed much in the past. Brachytherapy is used almost exclusively for in-stent restenosis. A numerous thrombectomy and embolic protection devices exist but none has shown a proven clinical benefit in randomized studies. With an advancement in medical technology adjunctive devices will play more important role in the future, especially for recanalization of chronic total occlusion and during acute myocardial infarction.

Key words
percutaneous coronary interventions, stents, adjunctive devices.

Introduction

Coronary balloon angioplasty (PTCA) introduced in 1977, followed by coronary stenting in 1986 represent the only two percutaneous interventions (PCI) that have documented and confirmed clinical benefit for the patient and are being performed in current practice at the large-scale basis. Many other coronary devices, still remaining as a part of interventional cardiologist’s arsenal, have triggered a great admiration at the time of initial clinical application but turned out to be less successful at the long-term follow-up. The author of this article has been in the field of interventional cardiology from the begging of its routine application and believes that these so-called “adjunctive” devices need to be mentioned. It is like reading a medical paper that shows negative result - but from it, one can learn a lot, sometimes even more than reading a study with positive, significant result. During last three decades, the level of performed PCI has progressively increased reaching currently a steady usage level while new device technologies are still in progress (Figure 1). For this analysis, Switzerland was chosen as sample country for comparisons due to its stable politi-
Figure 1. Correlation of PCI performed in Switzerland with publications on new coronary devices (number per year) during last three decades.
The plaque is gently displaced into the window housing and “shaved-off”, using the cutter, into the collecting nose cone at the distal tip of the device. In an early multi-center registry, DCA was successful in 85% of cases, and as such was subsequently comparable to success in balloon angioplasty. After having been approved by FDA, it gained confidence worldwide as an alternative to balloon angioplasty, and by 1990’s accounted for approximately 10% of PCI procedures in the United States. In the following years several non-randomized studies showed an improved intraluminal coronary gain after DCA as compared to PTCA that generated an extreme enthusiasm for the device. However, in 1993 after publication of a randomized CAVEAT study showing higher restenosis (57% versus 50%) and complication rates (11% versus 5%) with DCA compared to PTCA, the use of DCA rapidly decreased. The protagonists of atherectomy welcomed the new multicenter USA clinical trial with Orbital atherectomy device on 443 consecutive patients with severely calcified coronary lesions.

Rotablator TM (Boston Scientific USA) is the most known device of RA. It can effectively ablate calcified plaques, facilitating stent delivery and expansion. Practice guidelines recommend its use for preparation of heavily calcified or severely fibrotic lesion that cannot be crossed by a balloon or adequately dilated. However, late restenosis remains high when it is used as a stand-alone therapy or with bare-metal stents. Some observational study suggested a favorable long-term results of RA followed by DES implantation, but a randomized study including 240 patients showed that routine lesion preparation using RA did not reduce late lumen loss even after implantation of DES at 9 months follow-up.

Laser

The potential advantages of intracoronary laser angioplasty are to ablate the plaque material and vaporize all atherosclerotic plaque along the arterial wall. The bulk removal of plaque material could improve acute procedural success rates, decrease complication rates, take care of “untreatable” lesions, and decrease restenosis rates. Several types of lasers have been used in the past (Argon, Holmium), but only the excimer laser is still in practice (ELCA® Laser Ablation Catheter, Spectranetics USA). The coronary laser catheters are offered in sizes ranging from 0.9 to 2.0 mm in diameter and contain up to 250 small, flexible optical fibers mounted within a thin plastic tube.

The great enthusiasm to use the laser angioplasty for the treatment of coronary occlusions was raised in 1986 because laser energy can vaporize atherosclerotic plaque, and there may be no requirement for a pre-existing channel. However, high rate of complications, thermal injury and collateral miss have limited its application. A systemic literature search performed in 2014 by McGill University in Canada, found no benefit even in patients with un-crossable coronary lesion. Occurrence of complications such as coronary dissection (up to 9%), myocardial infarction (0-10%), or major bleeding (0-6%) and increased procedure cost influenced the decision not to recommend its use in Canada.

Brachytherapy

The radiation is believed to inhibit the cellular proliferation. Radiation for treatment of in-stent restenosis has been using two sources: gamma and beta radiation. Currently approved are the Checkmate System (Cordis Corporation) that uses gamma radiation and the Beta-Cath System (Novoste Corporation) that uses beta radiation. Approval by the FDA in USA for both of these devices is limited their use in stents that had been implanted in the past, and then re-stenosed. In the years following 2000, the treatment of restenosis after bare metal stent implantation using brachytherapy raised a great hope, but rapid developments in drug eluting stents had progressively pushed the brachytherapy aside. A meta-analysis in 2011 of 12 studies and comparing the outcomes of drug-eluting stents versus intracoronary brachytherapy for in-stent restenosis suggests that the use of drug-eluting stents for in-stent restenosis is associated with reduced occurrences of target-vessel revascularization and binary restenosis. The American College of Cardiology Guidelines (2011) do not recommend brachytherapy for the prevention of restenosis as the lower rates of restenosis occur with the use of drug-eluting stents in comparison to bare metal stents or vascular brachytherapy. Many technical limitations are present within brachytherapy, such as the operator’s protection against gamma radiation, coronary lesion geographical miss (unwilling exposure of healthy tissue to the radiation) and the absence of distinguished long-term benefit.

Thrombectomy

A numerous devices with different mechanisms of action exist all having the same objective to reduce distal thrombus embolization and improve myocardial perfusion. In a meta-analysis there was a significant improvement in ST-segment resolution, myocardial blush and TIMI grade 3flow as parameters of myocardial perfusion, as well as clinical parameters such as reduction in mortality. Non-manual, mechanical thrombectomy may have a role in selected patients with large caliber vessels and heavy thrombus burden. Catheter aspiration thrombectomy uses a catheter that is advanced over a guidewire to the thrombus whereby syringe suction is used to aspirate the debris. Devices used for this procedure include the DiverTM, Export®, ProntoTM, RescueTM, Thrombuster®, and TransVascular Aspiration Catheter®. Mechanical thrombectomy devices apply energy directed through
saline jets or a rotating catheter head to facilitate breakup of the thrombus prior to its active aspiration. These devices include the AngioJet® and X-Sizer®.

**Embolic protection**

Embolic protection devices can be proximal and distal, using either balloon or filters. Distal device employs an occlusion balloon advanced over a guide-wire distal to the thrombus in order to trap and aspirate thrombotic debris released during angioplasty and stenting procedures such as the PercuSurge GuardWire®, FilterWire EXT™, SpideRX™, AngioGuard™ and Filtrap. A study concerning these devices conducted in 2011 by the USA Agency for Healthcare Research and Quality examined 53 randomized trials (n = 8,185) and 9 observational studies (n = 1,479) and found no significant impact on mortality, myocardial infarction, stroke, or MACE versus a control when using the longest duration of follow-up.

**Stents**

There are several so called “dedicated” stents, designed for specific lesion categories such as bifurcations or vein grafts, which are on the market. None of them has showed clinical benefit as compared to the standard technique. Covered stent are of real utility in case of coronary artery perforation such as the Grafmaster RX, Abott, and mPK Papyrus, Biotronik, Switzerland - a more flexible stent. Specially designed drug eluting stents capable to accommodate multiple drugs in a special reservoirs incited a great hope at the beginning. The example is the NEVO® Cordis, USA, coronary stent but it was retrieved from the market after Johnson & Johnson Company announced halt of their research activities in interventional cardiology. The MGuard (Inspire MD, Boston USA) stent utilizes MicroNet™ technology, which is a circular knitted mesh that wraps around the stent to protect patients from plaque debris flowing downstream upon deployment (during acute myocardial infarction). Biodegradable vascular scaffolds (BVS) are very promising devices and will certainly expand in the future.

**CTO**

Recanalization of Chronic total coronary occlusions (CTO) has been for years and is still now a real battlefield for use of adjunctive devices (Figure 2). A recent meta-analysis study including 18,061 patients showed an angiographic success rate of 77% with following complications: contrast nephropathy 3.8%, coronary perforation 2.9%, myocardial infarction 2.5%, death 0.2%, emergent coronary surgery 0.1%, tamponade 0.3% and stroke 0.01%. The concept of mechanical recanalization with consequent angioplasty is still applied with different possibilities such as sub intimal tracking, re-entry technique or retrograde approach. Micro catheters represent a real advancement, the most known are Transit® (Cordis, Miami, United States), Finecross®, Pro Great® (Terumo, Japan). They have the advantage of large lumen and trackability for selective injection and tortuosity, whereas Corsair® (Asahi Intecc, Aichi, Japan) has more support, even beyond a calcified or tortuous segment. The Tornus device (Asahi) is a catheter made of 8 stainless steel strands woven together to enhance flexibility and strength in exchanging wires, delivering balloons and providing support for CTO procedures. The Frontrunner (Lumend, Cordis, USA) device is designed to create intraluminal blunt micro dissection to facilitate penetration of the fibrous cap. Two other recent systems for lesion crossing and lumen re-entry technologies include CrossBoss catheter (BridgePoint Medical, Plymouth, Minnesota), a metal micro catheter with a rounded tip that can advance through a CTO eventually into the sub intimal space. Then Stingray balloon and Stingray guide wire systems (BridgePoint Medical) which are employed to penetrate the distal true lumen for re-entry. In a catheterization laboratory that performs these interventions, other adjunctive safety devices are necessary like: covered stents, thrombectomy devices, snares, embolization coils and delivery systems to manage possible complications. With the advancement of medical technology, adjunctive devices will play more important role in the future, especially for recanalization of chronic total occlusion and for patients with acute coronary syndrome undergoing percutaneous coronary intervention.
References


