Longitudinal deformation of a third generation zotarolimus eluting stent: “The concertina returns!”

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Author contributions: Panoulas VF and Demir OM contributed equally to this work; Panoulas VF and Demir OM wrote the case report; Ruparelia N edited the manuscript and contributed to image formatting; Malik I reviewed and edited the manuscript.

Institutional review board statement: The study was reviewed and approved by the Imperial College Healthcare NHS Trust Institutional Review Board. Both patients provided informed consent.

Informed consent statement: All study participants provided informed written consent prior to study enrollment.

Conflict-of-interest statement: All authors declare no conflict of interest.

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Manuscript source: Invited manuscript

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Received: August 4, 2016
Peer-review started: August 5, 2016
First decision: September 2, 2016
Revised: September 15, 2016
Accepted: November 16, 2016
Article in press: November 17, 2016
Published online: January 26, 2017

Abstract

In the current case series we describe two cases of longitudinal stent deformation in ostial lesions treated with a new generation zotarolimus eluting stent and review current literature on longitudinal stent deformation. Historically not a common occurrence, longitudinal deformation occurred mainly in Promus Element everolimus eluting stents, which had only two rather than the commonly used 3 links between stent rings. Longitudinal deformation commonly occurs secondary to compression of the proximal edge of the stent by either the guide catheters, or intravascular balloons and imaging catheters. The degree of deformation however, depends on the longitudinal strength and design of the stent.

Key words: Coronary angioplasty; Longitudinal stent deformation; Coronary stents; Procedural complications

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Core tip: In the current case series we describe two cases of longitudinal stent deformation in ostial lesions treated with a new generation zotarolimus eluting stent and review current literature on longitudinal stent deformation.
Case 1
A 78-year-old male with diabetes mellitus, hypertension, hypercholesterolemia, and chronic renal failure presented with stable angina. Elective coronary angiography demonstrated significant lesions in the ostial and mid segments of a tortuous right coronary artery (RCA) (Figure 1A). The Judkins Right 4 catheter did not provide sufficient support therefore a 3 dimensional (3D) right coronary (Williams) guide catheter was used to intubate the RCA. This provided improved, yet suboptimal support, therefore a buddy wire was used to facilitate the implantation of a ZES (Resolute Onyx 2.75 mm × 38 mm, Medtronic) in the mid segment. Subsequently, the ostial-proximal lesion was pre-dilated, stented with a Resolute Onyx 3.0 mm × 22 mm ZES and post-dilated with a 3.0 mm non-compliant (NC) balloon (Figure 1B). Following stent deployment and removal of the buddy wire, significant longitudinal deformation was noted (Figure 1C), which was treated with non-compliant balloon dilatation and third ZES insertion (Resolute Onyx 3.0 mm × 18 mm) all the way to the ostium (Figure 1D).

Case 2
A 85-year-old male with end stage renal failure and severely impaired left ventricular systolic function presented with rapidly conducted atrial fibrillation and raised troponin (6000, upper normal limits 30) with inferolateral ST depression on electrocardiogram. On coronary angiography, he had severe three-vessel and left main disease. After being turned down for surgery, his proximal tight RCA lesion was stented with a Resolute Onyx 4.5 mm × 12 mm and post-dilated with a 5 mm non-compliant balloon. In view of the heavily calcified ostial/distal left main stem (LMS) and proximal left anterior descending (LAD) disease (Figure 2A) and bursting non-compliant balloons, decision was made to rotablate the lesions first with 1.5 mm burr. Subsequently a 3.0 mm non-compliant balloon was used to pre-dilate all lesions successfully. The proximal LAD was stented with a Resolute Onyx 3.0 mm × 26 mm stent. The whole length of the LMS into the proximal LAD was stented with a 3.5 mm × 34 mm Resolute Onyx covering the LMS ostium (Figure 2B and C). The LMS segment of the stent was post-dilated with a 4.5 mm NC balloon. Immediately after and despite taking care in removing the trapped LCx wire, there was longitudinal deformation of the stent, which no longer covered the LMS ostium (Figure 2D). After ballooning the deformed stent with a 4.5 mm non-compliant balloon the ostium was covered with another 4.0 mm × 8 mm stent and post-dilated with 4.5 mm NC balloon with an excellent final result (Figure 2E).

DISCUSSION
To our knowledge this case series demonstrates the first reported cases of longitudinal deformation in patients treated with the new Resolute Onyx stent platform.

Traditionally, longitudinal strength was not considered standard characteristics for stent performance. However, recent evidence has highlighted possible complications since longitudinal deformation was first reported by Hanratty et al[10], describing 3 cases where longitudinal compression of a previously deployed stent resulted in

INTRODUCTION
Over the last decade there has been remarkable progress in coronary stent design and materials. Early generation stents were stainless steel followed by the use of cobalt chromium and platinum chromium alloys, enabling stents to be thinner and more flexible, hence improving deliverability and conformability. In addition, thin stent struts have been associated with improved outcomes in drug eluting stents[1-3]. With the reduction in strut thickness, innovative designs have enabled maintenance of radial strength, however, longitudinal stent strength may have been compromised.

A recently observed complication includes longitudinal stent deformation, defined as the distortion or shortening of a stent in the longitudinal axis following successful stent deployment[4]. Longitudinal stent strength is dependent on the architectural composition - number, orientation, shape, thickness and material of the crest (or ring) and links. For example, the reduction in number of links between cells may enhance deliverability (by allowing more lateral bending) but as a consequence may compromise longitudinal strength. Longitudinal deformation can result in protrusion of struts into lumen. Extensive malapposition of struts may result in disruption of blood flow and increased risk of stent thrombosis[5,6]. In addition, longitudinal deformation of drug eluting stents may result in uneven drug delivery that can result in higher rates of instant restenosis[7].

In the current case series and review we describe two cases of longitudinal deformation with the new Resolute Onyx (Medtronic Inc., United States) zotarolimus eluting stents (ZES) and review the literature on longitudinal deformation. This new generation ZES has a novel design, manufactured from a single strand of core wire (platinum iridium) shaped into a continuous sinusoidal waveform. The strut thickness is 81 μm, rendering it extremely trackable and conformable. Every 4th crown is laser fused to provide uniform longitudinal strength across the length of the stent.

CASE REPORT

Case 1
A 78-year-old male with diabetes mellitus, hypertension, hypercholesterolemia, and chronic renal failure presented with stable angina. Elective coronary angiography demonstrated significant lesions in the ostial and mid segments of a tortuous right coronary artery (RCA) (Figure 1A). The Judkins Right 4 catheter...
Panoulas VF et al. Concertina of third generation drug-eluting stents

Figure 1  Longitudinal deformation of a stent implanted at the ostium of the right coronary artery. A: Initial angiogram showing severe calcific lesions in proximal and mid right coronary artery; B: Proximal lesion stented all the way to cover the ostium with a Resolute Onyx 3.0 mm × 22 mm stent; C: Longitudinal deformation of proximal stent treated with 3.0 non-compliant balloon and another 3.0 mm × 18 mm Resolute Onyx stent; D: Final angiographic result.

Figure 2  Longitudinal deformation of an ostial left main stem stent. A: Initial angiogram showing significant calcific ostial and distal left main lesions with further significant proximal LAD calcific disease; B: After 1.5 burr rotablation, LMS into proximal LAD was stented with a 3.5 mm × 34 mm Resolute Onyx (B) to 14 atm covering the LMS ostium (C). The LMS segment of the stent was post-dilated with a 4.5 NC balloon and the jailed LCx wire removed; D: Longitudinal deformation of the ostial LMS stent (yellow arrow pointing at the proximal deformed edge of the stent); E: Final angiographic result after covering the ostial LMS with a 4.0 mm × 8 mm Resolute Onyx stent. LAD: Left anterior descending; LMS: Left main stem.
stent deformation. Two were identified angiographically and one with the aid of intravascular imaging. It was first documented with the Promus Element (Boston Scientific) stent which was related to guide catheter compression of stents deployed in an ostial location[9]. However, Hanratty et al[10] have observed this phenomenon in other drug eluting stents. A retrospective analysis of 4455 interventional cases over a four-year period showed stent deformation occurred in 0.2% of patients affecting 0.097% of stents deployed[10]. In 6 cases, Promus Element was involved and there was 1 case each involving Endeavor (Medtronic), Biomatrix (Biosensors Interventional Technologies), and TAXUS Liberté (Boston Scientific) stents. The rate of stent deformation varied from 0% in several other stent types to 0.86% in the case of the Promus Element stent. In the same series, there was one case of late stent thrombosis attributable to longitudinal stent deformation[10]. In the DUTCH-PEERS study 906 patients were assigned to receive third generation zotarolimus-eluting stents (Resolute Integrity, Medtronic) and 905 to receive everolimus-eluting stents (Promus Element, Boston Scientific)[9]. Longitudinal stent deformation was seen only in the everolimus-eluting stent group [nine (1.0%) of 905 vs 0 of 906, \( P = 0.002 \); nine of 1591 (0.6%) everolimus-eluting stents implanted became deformed], but was not associated with any adverse events.

Despite drug eluting stents having improved remarkably the safety and efficacy of revascularization procedures, stent design is a continuously developing field that aims to balance numerous performance attributes such as stent flexibility, shortening on expansion, trackability, scaffolding, radiopacity, longitudinal strength, radial strength and recoil. An experimental evaluation of longitudinal strength of four commercially available stent design families demonstrated that a 50 g force resulted in longitudinal compression of 1.25-5.30 mm (4.46%-18.93%, compared with the nominal expanded stent length). The Promus Element stent platform had an average longitudinal compression of 13.20 mm (47.07%), demonstrating marked lower resistance to longitudinal compression (Table 1)[11]. Newer stent platforms with ultrathin struts (down to 60 \( \mu \text{m} \)) have shown non-inferiority to established everolimus platforms[11] but their longitudinal strength has yet to be assessed on the bench.

In our series both cases of longitudinal deformation occurred in the hands of very experienced operators implanting the scaffold at an ostial location. Potential reasons behind the deformation could include guide catheter compression of the proximal edge of the ostial stent post removal of jailed wires (case 1 or 2) or aggressive post-dilatation of the stent with a significantly larger NC balloon causing longitudinal shortening (case 2). It is unclear whether laser fusion provides less support compared to traditional links and further evidence is required prior to drawing any conclusions.

In conclusion, longitudinal stent deformation can occur even with new generation ZES and identification is important as, if left untreated, it may associate with a risk of future stent thrombosis, restenosis, and challenges in rewiring and retreatracting these lesions in the future.

### COMMENTS

#### Case characteristics

The authors presented two cases, one with stable angina and one presenting with a non ST elevation myocardial infarction, requiring treatment of ostial lesions with new generation drug eluting stents.

#### Treatment

Both cases were treated with the third generation zotarolimus eluting stent
Resolut Onyx, and in both cases longitudinal deformation of the stents was observed. This was managed with further ballooning and stenting.

**Experiences and lessons**

Longitudinal stent deformation can occur even with new generation ZES and identification is important.

**Peer-review**

This manuscript reports two cases of longitudinal deformation of a 22-mm and 34-mm third generation zotarolimus eluting stent. The issue brought up by the authors is interesting and the cases well documented.

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