Clinical update

Stent thrombosis and restenosis: what have we learned and where are we going? The Andreas Grünzig Lecture ESC 2014

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Modern-day stenting procedures leverage advances in pharmacotherapy and device innovation. Patients treated with contemporary antiplatelet agents, peri-procedural antithrombin therapy and new-generation drug-eluting stents (DES) have excellent outcomes over the short to medium term. Indeed, coupled with the reducing costs of these devices in most countries there remain very few indications where patients should be denied treatment with standard-of-care DES therapy. The two major causes of stent failure are stent thrombosis (ST) and in-stent restenosis (ISR). The incidence of both has reduced considerably in recent years. Current clinical registries and randomized trials with broad inclusion criteria show rates of ST at or < 1% after 1 year and ~0.2–0.4% per year thereafter; rates of clinical ISR are 5% respectively. Angiographic surveillance studies in large cohorts show rates of angiographic ISR of ~10% with new-generation DES. The advent of high-resolution intracoronary imaging has shown that in many cases of late stent failure neoatherosclerotic change within the stented segment represents a final common pathway for both thrombotic and restenotic events. In future, a better understanding of the pathogenesis of this process may translate into improved late outcomes. Moreover, the predominance of non-stent-related disease as a cause of subsequent myocardial infarction during follow-up highlights the importance of lifestyle and pharmacological interventions targeted at modification of the underlying disease process. Finally, although recent developments focus on strategies which circumvent the need for chronically indwelling stents—such as drug-coated balloons or fully bioresorbable stents—more data are needed before the wider use of these therapies can be advocated.

Keywords
- Bioresorbable stents
- Coronary artery disease
- Drug-eluting stents
- In-stent restenosis
- Neoatherosclerosis
- Stent thrombosis

Historical background

On 16 September 1977 Andreas Grünzig performed the first percutaneous coronary intervention (PCI) in an awake human at the Universitätsspital in Zürich, Switzerland.¹ Using a rudimentary balloon angioplasty catheter fashioned on his kitchen table, he treated a high-grade stenosis in the proximal left anterior descending artery of a 38-year-old man with very satisfactory acute and late results (figure 1).² Although his work would not have been possible without the pioneering endeavours of earlier physicist investigators, this procedure is rightly remembered as a landmark in the history of cardiovascular medicine.³

The limitations of balloon angioplasty however included an unpredictable acute result—due to early abrupt vessel closure—and a relatively high rate of restenosis at the site of the treated lesion—due mainly to plaque prolapse, vessel recoil, and constrictive remodelling. In this respect, it was the modification of this procedure to include the implantation of a metallic stent in the treated vessel that proved the tipping point to enable widespread uptake of PCI therapy. First human coronary stent implantation was performed in Toulouse and Lausanne within weeks of each other in March and April 1986. (Figure 1).³ By splitting angioplasty-induced arterial dissections and sealing disrupted plaques stent implantation resulted in less acute vessel thrombosis. Moreover, the additional advantage in
terms of mechanical strength with stent implantation resulted in
greater acute gain in luminal calibre and negation of the effects of
vessel recoil and constrictive remodelling. This translated into a sig-
ificantly lower rate of subsequent restenosis.

As the advantages of stent implantation saw its evolution from a
‘bail out’ after complicated balloon angioplasty to a standard treat-
ment strategy, two important limitations were recognized. First, a
not insignificant number of cases continued to result in early acute
vessel closure due to stent thrombosis (ST). An early study re-
ported complete occlusion occurred ≈25% of cases mostly within
the first 14 days after implantation. Moreover, these complications
occurred despite the fact that early stent procedures were often
undertaken with very large doses of heparin (up to 15 000
units)—sometimes with dextran or urokinase infusions—as well
as overlapping oral anti-coagulation. This in turn resulted in

significant morbidity and mortality due to haemorrhagic complica-
tions—with major bleeding occurring in 9% of patients in an early
study at our centre. Indeed, arguably one of the most important de-
velopments in the evolution of stent therapy was the demonstration
that dual antiplatelet therapy (DAPT) with aspirin and an
ADP-receptor inhibitor could reduce both ST and bleeding compli-
cations in comparison with oral antithrombotic therapy. Together
with technical refinements such as the use of routine
high-pressure stent deployment, these developments facilitated
the widespread adoption of coronary stenting for the treatment
of a broad range of obstructive coronary artery disease.

The second important limitation was late stent failure due to
in-stent restenosis (ISR). Although stenting was important in re-
sisting acute and late constrictive mechanical forces, the stent im-
plantation procedure resulted in increased acute vessel injury at

Figure 1 Historical perspectives on the development of percutaneous coronary intervention and coronary stenting. The first coronary angio-
plasty in an awake human was performed by Andreas Grünzig (A) on 16 September 1977 using a balloon catheter fashioned on his kitchen table
(B). The patient had a high-grade stenosis of the proximal left anterior descending artery and the initial and late follow-up result (D) was very
satisfactory. The first coronary stents implanted in man were performed by Ulrich Sigwart (E) in Lausanne and Jaques Puel (F) in Toulouse in March
and April 1986. An angiogram from an initial patient shows high grade stenosis of the proximal left anterior descending artery (G), which was
treated with a bare metal Wallstent (H) with a good acute result (I).
the time of PCI and an enhanced healing response leading to varying
degrees of neointimal hyperplasia. It was this issue that prompted
research leading to the development of drug-eluting stents (DES).

Drug-eluting stent devices have proved highly effective in redu-
cing the incidence of stent failure and enabled the expansion of
PCI to treat high-risk patient and lesion subsets. Iterative develop-
ment has focused on thinner stent struts and more biocompatible
polymer coatings as well as stents that are fully bioresorbable.

A wide range of DES is currently available. A recent systematic re-
view of DES devices by a European Society of Cardiology-European
Association of Percutaneous Coronary Intervention Task Force
identified at least 68 DES with CE-mark approval as of June
2014. The key characteristics of selected DES devices with pub-
lished large-scale randomized clinical trial data are shown in Figure 2.
Indeed, although it is a subject of considerable debate, the high effi-
cacy and excellent safety of current generation devices may be asso-
ciated with not just improvement in quality of life but also in survival
in treated patients.

Stent thrombosis

An overview of histopathology, risk factors, incidence, and intravas-
cular imaging features of ST is provided in Figure 3A–D, respectively.

Incidence and time course

Stent thrombosis is typically characterized by angiographic or post-
mortem evidence of recently formed thrombus in a previously
stented segment (Figure 3A). Study of thrombus aspirates from pa-
tients presenting with ST have shown a mix of thrombotic and in-
flammatory components including platelet-rich thrombus, fibrin
fragments, and leukocytes of both neutrophil and eosinophil lin-
eage. In order to standardize reporting across clinical trials univer-
sal definitions were agreed upon in 2006 by a group of experts
known as the Academic Research Consortium. This definition
classified evidence of ST as definite, probable, or possible as well
as according to timing after the initial stent implantation (Table 1).
In practice, because of differing pathophysiology and risk factors,
it can be useful to dichotomize events into two major categories:
early ST is defined as thrombosis within the first 30 days and late
ST is thrombosis occurring beyond 30 days. In general, early ST is
more common than late, accounting for ~50–70% of all cases de-
pending on the overall time frame of reference.

In recent years, important progress has been made in reducing the
incidence of ST. Recent large-scale registries show that with contem-
porary antithrombotic therapies and modern generation DES the rate
of early ST is <1% (Figure 3C). Moreover, a recent systematic
review of randomized trials with DES reporting results at 9–12
months showed a median incidence of definite ST of 0.61%. In add-
ition, overall rates of early and late ST out to 3 years have halved in
recent years from ~3.0 to 1.5% (Figure 3C).

Risk factors for stent thrombosis

In general, it can be useful to classify risk factors as patient-, proce-
procedure-, or device specific (Figure 3B).
Early stent thrombosis

In terms of early ST procedural risk factors are the most important. Stent undersizing, presence of residual dissection, impaired TIMI flow and residual disease proximal or distal to the stent lesion are important predictors of ST. A French study showed that lesion complexity and index PCI in the setting of acute myocardial infarction were strong predictors of subsequent ST. In addition, patient-specific risk factors such as reduced left ventricular function and impaired response to ADP-antagonist therapy confer important increased risk. Indeed, premature discontinuation of antiplatelet therapy in the initial 30 days after stenting is arguably the most important predictor of ST. Moreover, considerable interest has focused on predicting risk based on response to ADP-antagonist therapy. First, pharmacogenetic testing seems to be able to identify patients-at-risk based on genetic polymorphism related to enzymes required for clopidogrel metabolism. Second, many studies have shown an association between high on-treatment platelet reactivity in platelet function testing and subsequent ST. For example, a registry study from our centre showed a 9-fold risk of early ST in patients with high on-treatment platelet reactivity. Importantly, however, while both pharmacogenetic and platelet function testing are attractive for identifying patients at risk no trial has yet been able to show significant improvement in outcomes if treatment is modified (i.e. intensity increased) on the basis of this data.

Finally, although device-specific factors were thought to be of lesser importance in determining the risk of early ST, recent studies suggest that there may be important differences. Analysis of large datasets suggests that rates of early ST seem to be slightly higher

Figure 3 Stent thrombosis: central illustration of histopathology, risk factors, incidence, and intravascular imaging features. (A) Representative case showing late stent thrombosis with uncovered struts following drug-eluting stent implantation. Histologic section from a 47-year-old male who had overlapped drug-eluting stents (paclitaxel-eluting stent in the proximal segment and sirolimus-eluting stent in the distal segment) implanted 10 months prior to death. A low-power image (i) shows a platelet-rich occlusive thrombus in the lumen in paclitaxel-eluting stent. A high-power image (ii) of boxed area in (i) shows uncovered struts with peri-strut fibrin. Image (iii) also shows a platelet-rich occlusive thrombus. A high-power image (iv) of boxed area in (iii) shows partially covered struts with neointima (Asterisk indicates stent strut). (B) Principal risk factors for stent thrombosis classified according to patient-related, stent type-related, and procedure-related risk factors. (C) Incidence of stent thrombosis after bare metal stents, early-generation drug-eluting stents (G1 DES), and new-generation drug-eluting stents (G2 DES); adapted from Tada et al. (D) Representative optical coherence tomography findings from patients presenting with stent thrombosis: (i) persistent uncovered stent struts late after implantation; (ii) marked stent malapposition in the target vessel, this may have been present at the time of implantation or acquired due to late positive remodelling; (iii) neatherosclerotic plaque formation: diffuse low-signal intensity with higher backscatter in deeper neointimal layers may indicate underlying lipid-rich atherosclerotic tissue; (iv) severe stent underexpansion at site of overlap of multiple stent layers.
Late stent thrombosis

Although significant technical shortcomings in the index procedure will more likely manifest as early stent failure, such factors can also play an important role in late ST where significant mechanical issues—e.g. stent undersizing or underexpansion—remain after the time point of DAPT discontinuation. Malapposition (or incomplete stent apposition) is often observed on intravascular imaging in patients with ST. However, definitive evidence of its clinical importance based on appropriately designed case–control studies is lacking at present. In particular, the threshold at which malapposition distance and extent become clinically relevant is not well defined. Patient-specific risk factors also remain important for late ST. In particular, reduced left ventricular function and diabetes mellitus are associated with increased risk. In addition, impaired response to ADP-antagonist therapy also confer important increased risk for late ST.

An important role in late ST is played by stent type-related factors. Controversy generated by presentations at the European Society of Cardiology annual meeting in 2006 focused attention on a possible increased risk of cardiac death with early-generation DES devices, mediated through higher rates of ST. Indeed, a number of meta-analyses that appeared shortly afterwards showed evidence of a small but significant increase in the risk of ST with both sirolimus- and paclitaxel-eluting stents. Moreover, registry reports showed evidence of an ongoing risk of ST out to 4–5 years with no clear evidence of an attenuation of this effect with time. The underlying substrate for this excess risk was identified in autopsy studies to be delayed arterial healing, a pathophysiological process characterized by impaired endothelial coverage, persistent fibrin deposition, and ongoing vessel wall inflammation. Although clearly multifactorial in origin, it seemed to be that inflammatory reaction to the durable polymer coatings used in early-generation devices played an important role. Indeed, the passage of time has shown us that delayed healing likely underlies a spectrum of adverse clinicopathological entities including not just late ST but also delayed late luminal loss (which may contribute to late re-stenosis), persistent vasomotor dysfunction proximal and distal to the stented segment, and de novo in-stent atherosclerosis. Newer generation DES seem to have addressed this healing problem in a meaningful way by incorporating thinner stent struts (which reduce acute vessel injury), more biocompatible polymer coatings (both nonerodable and biodegradable), and lower dosages of sirolimus-analogue drugs (Figure 2). Although each of these iterative developments may be clinically relevant in isolation, it should not be forgotten that overall clinical performance of DES is due to aggregate effects of both the backbone and the drug-matrix coating.

### Clinical consequences and treatment of stent thrombosis

Stent thrombosis is a very serious clinical event typically resulting in ST-elevation myocardial infarction in the majority of cases and mortality rates that may be as high as 20–40%. Although detailed consideration of management of ST is beyond the scope of this review, most registries of ST report that thrombus aspiration and balloon angioplasty are frequently used with repeat stenting in ~30–50% of cases.

### Dual antiplatelet therapy duration and prevention of stent thrombosis

Key to the prevention of ST is the prescription of an appropriate duration of DAPT after PCI. Randomized clinical trials in the 1990s demonstrated conclusively that DAPT was superior to anti-coagulation for the prevention of complications after bare metal stenting. Early randomized clinical trials with DES implantation included a recommendation for 3–6 months of DAPT after PCI, though concerns soon emerged about a possible increase in late ST after DES implantation. Two important consequences of these were that (i) guideline authorities recommended—on the basis of expert opinion—a more prolonged duration of DAPT of typically at least 12 months and (ii) a number of large-scale clinical trials were initiated to define more precisely the optimal duration of DAPT after DES implantation (see Table 2). Data from all of these studies have now been reported permitting reappraisal of recommendations relating to DAPT duration.

Results from the initial studies to report data showed that prolongation of DAPT did not reduce ischaemic adverse events but did lead to an excess of major bleeding events. However, none
of the individual studies were powered to show a reduction in ST, all were open-label, and by virtue of randomizing patients at the time of stent implantation rather than of treatment divergence, many of these trials may have been biased towards a null effect. In this respect, the recently published results of the DAPT trial are of high clinical relevance. Overall, 9921 patients treated with DES were randomized to prolonged duration (30 months) or standard duration (12 months) DAPT. The main findings were that prolonged DAPT duration significantly reduced ST [0.4 vs. 1.4%; hazard ratio, 0.29 (95% CI, 0.17–0.48); P < 0.001]. Results were consistent according to clinical presentation at the time of index stenting. Moreover, a number of aspects should be considered when interpreting the data. First, although no interaction with stent type was observed, a clustering of ST events occurred in patients treated with early-generation DES, stents which have now fallen out of use—e.g. 27% of enrolled patients were treated with paclitaxel-eluting stents, while 57% of all ST occurred in patients treated with this stent. Second, there is some cause for concern due to a higher rate of death in patients treated with prolonged duration DAPT. However, an association with bleeding events is not clear and it remains possible that this is a chance observation. Third, despite the best efforts of investigators only selected patients were included: the majority of screened patients were not represented in the randomized controlled trial. Fourth, it must of course be recognized that prolongation of DAPT not only offers the possibility to mitigate the risk of ST but also to prevent ischaemic complications not related to the stented lesion; in this respect, the demonstrated reduction in the overall incidence of myocardial infarction [2.1 vs. 4.1%; hazard ratio, 0.47 (95% CI, 0.37–0.61); P < 0.001] is an important finding. Finally, an update meta-analysis of published trials on DAPT duration supports the individual findings of the DAPT trial (Figure 4). The clear reduction in ST at the expense of increased major bleeding highlights the importance of clinical judgement in individualizing treatment duration for our patients in clinical practice. Accordingly, while a general time window for optimal duration of DAPT—between 6 months and 30 months duration—might be recommended, a one size fits all approach for DAPT duration is likely not optimal. Moreover, DAPT duration in clinical practice is a dynamic decision, which might be re-assessed at regular intervals during follow-up.

Thirdly, optimal management of patients receiving OAC who are treated with stent implantation is unclear. Industry-support for...
randomized trials has been lacking and only two modest-sized investigator-initiated RCTs have been completed. The WOEST investigators showed overall comparable outcomes between OAC plus clopidogrel vs. OAC plus dual antiplatelet therapy for 12 months. In the ISAR-TRIPLE trial, we also showed broadly similar outcomes between a strategy of 6 weeks DAPT vs. 6 months DAPT. However, both of these trials were significantly underpowered to detect differences in safety endpoints. Presently, a number of large-scale industry-supported RCTs are examining the best treatment for this important patient subgroup. Finally, more potent oral ADP-receptor antagonists significantly reduced the incidence of ST in comparison with clopidogrel; however, these agents are currently recommended only in patients undergoing stenting following presentation with an acute coronary syndrome. Moreover, application of novel intravenous ADP-receptor antagonists may further reduce rates of ST in the acute phase after stenting.

In-stent restenosis

An overview of histopathology, risk factors, incidence, and intravascular imaging features of ISR is provided in Figure 5A–D, respectively.

The higher degree of vessel injury with stent implantation in comparison with balloon angioplasty alone increased the extent of neointimal hyperplasia in the intervened segment and this is the dominant cause of restenosis after bare metal stent implantation. Restenosis after PCI has been characterized as a distinct pathophysiological process rather than merely an accelerated form of post-intervention atherosclerosis. In general, terms inflammatory response to vessel wall injury during PCI plays a central role in restenosis after stenting with vessel wall inflammation driving fibroblast growth and smooth muscle cell hyperplasia. Mechanistically contributing factors to restenosis after vascular intervention may be divided into five categories: (i) acute or subacute prolapse of the disrupted plaque, (ii) elastic recoil of the vessel wall, (iii) constrictive remodelling, (iv) neointimal hyperplasia (due to extracellular matrix deposition and smooth muscle cell hyperplasia), and (v) de novo in-stent atherosclerosis (neoatherosclerosis).

Angiographic restenosis is commonly adjudicated as a binary event defined as a re-narrowing of >50% of the vessel diameter as determined by coronary angiography. Intravascular imaging modalities acquire data in three dimensions, using these modalities restenosis is defined as a re-narrowing of >75% of the reference vessel area in cross-section. The term clinical restenosis is sometimes used to refer to restenosis of the treated lesion accompanied by requirement for re-treatment, for example, due to symptoms or signs of ischaemia. Rates of clinical restenosis are usually considerably lower than rates of restenosis detected by imaging as not all restenotic lesions cause ischaemia or elicit symptoms.

From a pathological standpoint, it appears that there are considerable differences between restenosis that occurs after bare metal stenting vs. after DES (Table 3; Figure 5). The main difference is that restenosis after bare metal stenting is typically characterized by neointimal hyperplasia consisting of a proteoglycan matrix and high

![Figure 4](http://eurheartj.oxfordjournals.org/)

**Figure 4** Summary results of meta-analysis of trials investigating prolonged duration vs. standard duration dual antiplatelet therapy after drug-eluting stent implantation. Odds ratio with (95% confidence interval) associated with prolonged vs. control dual antiplatelet therapy accounting for events occurred at the longest follow-up available in each included studies. The diamonds and the horizontal lines indicate the odds ratio and the (95% confidence interval) derived from meta-analysis. DAPT, dual antiplatelet therapy; figure based on analysis of data from Cassese et al.
proportion of vascular smooth muscle cells. In contrast, restenosis after DES is typically characterized by a proteoglycan-rich neointimal hyperplasia with relatively few smooth muscle cells. Moreover, neoatherosclerotic change within the restenotic tissue is seen earlier and more frequently in DES restenosis. Intravascular imaging with OCT also reveal distinct differences between the two processes: imaging of bare metal stents, early-generation drug-eluting stents, and new-generation drug-eluting stents over time and rates of binary angiographic restenosis (red line) in a large registry of patients with angiographic surveillance after stent implantation. Adapted from Cassese et al.\(^ {85} \) (D) Optical coherence tomography imaging of patients with in-stent restenotic tissue during surveillance after stenting; tissue with homogeneous-signal intensity (i) is typical after bare metal stenting; heterogeneous, attenuated, or layered signal intensity tissue (ii–iv) is typical after drug-eluting stents.

**Incidence and time course**

Systematic angiographic surveillance in \(>10,000\) patients undergoing coronary stenting at our centres showed rates of angiographic restenosis of \(\approx 30\%\) after bare metal stenting.\(^ {81} \) By inhibiting vascular smooth muscle migration the introduction of DES led to a significant reduction in rates of angiographic restenosis to \(\approx 15\%\) with early-generation devices and 12% with newer generation devices (Figure 5). On the other hand, contemporary randomized trials without angiographic surveillance typically demonstrate rates of clinically relevant restenosis of \(< 5\%\) at 12 months.\(^ {82} \)

angiographic surveillance studies have shown that neointimal formation after bare metal stenting tends to peak at 6 months after stenting and thereafter remain stable or regress somewhat over the medium term.\(^ {12,83} \) This is in keeping with completion of vessel healing, contraction of neointima, and positive remodelling of the vessel wall. Interestingly, more prolonged follow-up of series out to 7–11 and 15–20 years indicated some further luminal narrowing beyond 4 years.\(^ {84,85} \) After DES implantation however, the time course of restenosis seems to be rather different. In a large serial angiographic follow-up registry, we found that ongoing erosion of luminal calibre between 6 and 8 months and 2 years post-stenting is a feature of DES therapy.\(^ {86} \) Raber et al. showed incremental late loss in patients treated with first-generation DES who had surveillance angiography at 6–8 months and 5 years.\(^ {86} \)
observation of ongoing delayed late loss with DES beyond the 6- to 8-month time window supports the hypothesis of DES-associated delayed arterial healing seen in autopsy and preclinical studies and suggest that the temporal course of restenosis with DES may be significantly right-shifted compared with bare metal stents. Moreover, some of this late erosion of luminal calibre may be attributable to the higher incidence of in-stent neoatherosclerosis formation in DES.\(^{48}\)

### Risk factors for in-stent restenosis

The occurrence of ISR may have an important impact on long-term prognosis after PCI\(^{87}\) and identification of patients at risk is an important undertaking. Risk factors for ISR can also be classified as patient-, procedure-, or device-specific (see Figure 5).\(^{88,89}\)

Early studies in the DES era showed that the major risk factors for restenosis after DES were vessel size, final diameter stenosis, and type of DES (i.e. SES were more effective in preventing ISR than paclitaxel-eluting stent).\(^{90}\) By reducing the extent of injury at the time of implantation thinner stent struts are also associated with a reduced restenotic risk in comparison with thicker struts.\(^{91}\) In the largest analysis to date, we investigated the risk factors for restenosis in a series of 10,004 patients with angiographic follow-up after coronary stenting.\(^{81}\) Binary restenosis was detected in 26% of patients overall. At multivariate analysis, smaller vessel size (odds ratio 1.59 [95% confidence interval, 1.52–1.68] for each 0.5-mm decrease), total stented length (1.27 [1.21–1.33]), complex lesion morphology (1.35 [1.20–1.51]), diabetes mellitus (1.32 [1.19–1.46]), and history of bypass surgery (1.38 [1.20–1.58]) were independently associated with restenosis. Moreover, use of first-generation DES vs. bare metal stents (0.35 [0.31–0.39]) and second-generation DES vs. first-generation DES (0.67 [0.58–0.77]) were independent predictors of lower rates of restenosis. Overall in terms of therapeutic measures to reduce restenosis, the most important issues are likely to be meticulous attention to procedural detail and use of high-performance DES. Other approaches including systemic pharmacotherapy have been associated with mixed results and are discussed in detail elsewhere.\(^{92}\)

In terms of risk prediction using biomarkers, a number of studies have investigated an association between inflammatory biomarkers at the time of stenting and subsequent restenosis though the clinical utility of such an approach is unclear. At our centre, we showed that although baseline CRP levels did not seem to correlate with restenosis, the change between baseline and peak post-intervention CRP values strongly correlated with angiographic restenosis.\(^{93}\) A subsequent study from Park et al. also failed to show an association between baseline CRP and restenosis\(^{94}\) as did a larger analysis of four ISAR randomized trials.\(^{95}\)

### Clinical consequences and treatment of in-stent restenosis

Although ISR may be associated with a recurrence of stable angina symptoms, it is well recognized that up to a third of patients present with myocardial infarction or unstable angina\(^{96}\) and in contemporary clinical trials of patients with ISR \(\sim 20\%\) of patients have biomarker positive acute coronary syndrome. Most patients requiring treatment are amenable to repeat catheter intervention and the most effective strategies seem to be repeat stenting with new-generation DES or angioplasty with drug-coated balloons.\(^{97}\)

### Neoatherosclerosis as a common pathway in late stent failure

Neoatherosclerosis is a term coined to describe the development of atherosclerotic plaque inside an implanted coronary stent. Histopathologically, the process is characterized by three main stages: (i) early foamy macrophage infiltration, (ii) manifest atherosclerotic plaque development, and (iii) necrotic core plaque formation with or without thin fibrous caps (Figure 6). Although neoatherosclerosis is also observed after bare metal stenting, it occurs earlier and more frequently after stenting with DES. The first systematic report of this process described findings in a series of autopsy segments after drug-eluting and bare metal stenting.\(^{98}\) Neoatherosclerotic change inside the stent was seen in a higher proportion of cases after DES when compared with after bare metal stent implantation (35 vs.
Moreover, early neoatherosclerotic changes—such as foamy macrophage infiltration—were as early as 4 months after DES when compared with only beyond 2 years after bare metal stent implantation. These findings were confirmed in a subsequent large series of 384 autopsy specimens. Moreover, in spite of iterative development in device technology, the incidence of neoatherosclerosis seems comparable between early- and new-generation DES. The observations may be explained by the fact that under
normal circumstances, arterial walls are protected from infiltration by circulating lipid particles by a healthy endothelial cell barrier. However, DES are well known to cause anatomical and functional endothelial impairment—features characteristic of delayed arterial healing. Accordingly, it can be hypothesized that the presence of incompetent endothelium after DES is more likely to lead to accelerated and perhaps more frequent neatherosclerosis.

In parallel with these reported autopsy series, the increasing availability of high-resolution intravascular imaging with optical coherence tomography (OCT) permitted the improved characterization of in-stent tissue in patients presenting with late stent failure. Evaluation of tissue extent and homogeneity, signal attenuation, and neointimal tissue contours permits identification of possible areas of neatherosclerosis with or without necrotic core formation and plaque rupture or erosion in clinical practice.\(^{78}\) Moreover, development of techniques for quantitative analysis of OCT signal intensity offers potential for more accurate identification of these tissue types in future studies.\(^{99}\) A series of 33 patients with OCT imaging during intervention for late ST showed neatherosclerotic plaque in \(~70\%\) of cases.\(^{100}\) Initial reports in patients with ISR suggested that ca. 50\% of cases had tissue type consistent with neatherosclerosis.\(^{101}\) In another study, predictors of neatherosclerosis in patients with in-stent intimal hyperplasia included stent age (\(>48\) months), DES as stent type, current smoking, and chronic renal insufficiency.\(^{102}\) Taken together, both human autopsy and clinical imaging studies suggest that in many cases neatherosclerosis is the final common pathway for late stent failure.\(^{48}\) Important limitations of existing datasets are a paucity of histopathological correlation studies and the absence of an established preclinical model of neatherosclerosis.\(^{38}\) Although OCT imaging is intuitively attractive, the routine use of this imaging modality for guiding treatment of stent failure is presently not supported by clinical trial data.

**New developments: bioresorbable stents**

Bioresorbable stents (BRS) are an important technological development with potential to enhance the outcomes of patients treated by PCI and radically change future stenting practices. The basic concept is based upon the degradation of the stent backbone to inert particles after its useful function is served; once the stent has been fully degraded, this theoretically removes the risks associated with both ST and restenosis. Moreover, these devices offer additional potential benefits, including restoration of normal vasomotor tone of the stented segment and increase in lumen calibre due to positive vessel remodelling associated with stent degradation. This might in turn translate into improvements in coronary physiology and reduction in angina symptom burden.\(^{14}\)

Clinical trial reports in selected patients with comparatively straightforward lesion morphology have shown some encouraging results with BRS technology.\(^{103}\) However, the generalizability of these observations is unclear. Recently, a number of registry studies from real-world practice have been published and these data are of interest for a number of reasons.\(^{104}\) First, the overall clinical performance of these stents seems satisfactory—at least over short-term follow-up—though more data are needed in larger patient numbers. Secondly, restenosis rates seen with these stents in clinical practice seems low and in line with observations seen in early clinical trials. Thirdly, the rates of ST seen with BRS in the first 6–12 months seems higher than that observed with current generation metallic DES. Moreover, the majority of these events occur within the first 30 days. This means that their occurrence is likely related to the implantation procedure and could be influenced by the expertise of the operator. Specifically, careful selection of patients and lesions is critical and meticulous attention to implantation detail is essential, including a low threshold to use intravascular imaging for optimization of stent deployment. Overall, although BRS is undoubtedly a potential breakthrough technology, concern exists regarding greater complexity in implantation technique and a possible excess of early ST.\(^{107}\) Current devices likely represent relatively immature iterations of the technology requiring careful patient selection and meticulous attention to implantation technique. Further device iteration—with improved backbones and optimized radial strength—will likely be required before widespread adoption can be recommended.

**Perspective**

The great progress made with coronary stenting and antithrombotic therapies over the course of the last 25–30 years means that the vast majority of our patients who require PCI—including those with unstable presentations, complex disease patterns, and multiple co-morbidities—can be successfully treated in a safe and effective manner. Although nowadays PCI almost always involves implantation of a permanent metallic prosthesis, the excellent clinical outcomes over the short- to medium-term support the effectiveness of this approach. However, a number of areas of unmet need still exist. First, we need to remain focused on the fact that while stent implantation relieves symptoms and may improve prognosis, it remains a downstream therapy which targets the final common pathway of cardiovascular disease rather than the underlying disease process. Indeed, even at a follow-up interval as short as 3-years after stenting, natural history studies show that progression in non-target lesions starts to predominate as a cause of subsequent myocardial infarction.\(^{103}\) In this respect, the cornerstone of therapy remains risk modification, though lifestyle intervention and treatment with disease modifying agents; the impact of application of PCSK9 inhibitors—potent systemic therapies for lipid lowering—on both overall disease progression as well as late stent failure will be potentially considerable.\(^{109}\) Secondly, the availability of high-resolution intravascular imaging has increased our awareness of neatherosclerosis—a final common pathway in many cases of thrombotic or restenotic late stent failure.\(^{48}\) A better understanding of this disease process as well as the development of targeted therapies to prevent its occurrence will likely have a significant impact on the late outcomes of our patients. Thirdly, there remain lesion and patient subtypes in need of improved interventional device options: these include stent implantation in the setting of acute myocardial infarction, bifurcation lesions, chronically occluded vessels, lesions with severe calcification, ISR, and lesions in patients with diabetes. For this reason, we need to ensure that safe and effective methods of evaluation of innovative coronary stent devices remain available.
to our patients.\footnote{16} Finally, although recent developments focusing on strategies which circumvent the need for chronically indwelling stents—a so-called leave nothing behind approach—are promising, more data are needed before the wider use of these therapies can be advocated. In particular, though drug-coated balloon therapy seems effective for ISR,\footnote{10} it remains unestablished for the treatment of de novo coronary disease. In addition, though treatment with fully bioresorbable stents is feasible and intuitively attractive, concerns related to unacceptable rates of early stent failure need to be addressed most likely through further iterative development. These issues and concerns notwithstanding, as we look back at the progress of the last decades, we must acknowledge that it has been a remarkable journey from the kitchen table of Andreas Grünzting to the contemporary procedures and excellent outcomes available to our patients today. It remains a great pity that his premature death meant that he did not live to bear witness to these important developments.

**Authors’ contributions**

R.A.B., M.J., A.K.: handled funding and supervision, acquired the data, conceived and designed the research, drafted the manuscript, and made critical revision of the manuscript for key intellectual content.

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**Conflict of interest**:


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Stent thrombosis and restenosis


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