Drug eluting stents: maximising benefit and minimising cost

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A policy of selective implantation of drug eluting stents, in a minority of lesions most likely to benefit, seems to be a rational way to employ this new and currently costly technology.

The RAVEL and the SIRIUS studies, using a sirolimus eluting stent, report substantial reductions in clinical restenosis with drug eluting compared with bare stents. The drug eluting stent has been hailed as the third great breakthrough in percutaneous coronary intervention (PCI) after the balloon and the stent, but excitement greeting its arrival must be tempered by three facts. Firstly, it is impossible to predict exactly which individual coronary artery stenosis will restenose (making it difficult to know who would benefit from the new stent). Secondly, “real world” PCI includes many lesion types more adverse (and probably more deserving of anti-restenosis strategies) than those included in the studies. Thirdly, the first drug eluting stent on the market costs about five times more than a conventional stent (making it economically impossible to treat all lesions). We need to decide how to make rational treatment decisions with this new device within limited health budgets. In this article, we attempt to do exactly that.

The first drug eluting stent (Cypher, Cordis) has recently been made commercially available in the UK. Data from the RAVEL study and the “first-in-man” studies suggest that this device, which elutes the drug sirolimus (rapamycin, a macrolide antibiotic, with immunosuppressant, antiproliferative, and antimigratory properties), produces a zero restenosis rate up to, and beyond, one year in selected lesions. Interestingly, the larger SIRIUS study (similar to RAVEL, but performed in the USA), as yet unpublished, has complete follow up data available for the first 400 patients enrolled; the study demonstrates (as its primary end point) a 12% target vessel failure (TVF) rate in the sirolimus group compared with 23% in the control group, while the target lesion revascularisation (TLR) rate (an index of which patients will undergo restenosis) in the stent group, and the re-PCI rate from 21% to 10% (in highly selected lesions). As a result, interventional cardiology moved into the “stent era”, stents being implanted in 70–90% of PCI procedures (in decidedly unselected lesions). It is natural that interventionists will want coated stents, and patients will demand them. This enthusiasm, however, comes with a cost. The Cypher drug eluting stent is being marketed in the UK at a price about fivefold that of a bare stent. This mandates a vigorous appraisal of the financial implications of its use, at a time of expansion in PCI, in an increasingly cost conscious health service.

WHO GETS RESTENOSIS?

It is often stated that it is not possible to predict which patients will undergo restenosis. It is, however, true that certain lesion (and, to a lesser extent, patient) characteristics are associated with higher restenosis rates. Multivessel stenting and diabetes are the two main patient related variables increasing the probability of restenosis. In the case of diabetes, the relative risk is about 1.3:1. The evidence for specific lesion locations having high restenosis rates is generally unconvincing (such as the proximal left anterior descending artery), an exception being the saphenous vein graft. Angiographic, lesion related variables are more important, notably the minimum lumen diameter (MLD) pre-procedure, the MLD post-procedure, the acute gain, the vessel size (reference diameter), stent length, and the presence of multiple stents (probably a manifestation of the same phenomenon). Intravascular ultrasound (IVUS) based studies support these observations and show that an ostial location, plaque burden (plaque area/arterial area) and final

Abbreviations: CABG, coronary artery bypass grafting; IVUS, intravascular ultrasound; MLD, minimum lumen diameter; PCI, percutaneous coronary intervention; RAVEL, randomized comparison of a sirolimus-eluting stent with a standard stent for coronary revascularization; SIRIUS, US multicenter, randomized, double-blind study of the sirolimus-eluting stent in de novo coronary lesions; TLR, target lesion revascularisation; TVF, target vessel failure.
lumen dimensions (MLD and lumen cross sectional area) are also important. Approximately half of patients with angiographic restenosis (diameter stenosis > 50%) present with clinical relapse. These usually require repeat revascularisation, the majority by PCI. On the other hand, “real world” restenosis rates are likely to be higher than those found in randomised controlled trials. Up-to-date, local data are important. Examination by one author (JG) of the records of 219 PCIs performed in 2001 (an unselected series, including acute and chronic cases, and all types of lesion) revealed that a mean of 1.6 vessels per patient were treated with a mean of 1.1 stents per vessel. The clinical restenosis (TLR) rate for these patients was 10.0% (of which 8.6% were for restenosis and 1.4% re-restenosis), 96% being treated with re-PCI and 4% with coronary artery bypass grafting (CABG). This figure of 10% TLR is very similar to those seen in unselected, contemporary, published series from other centres and operators.

POTENTIAL STRATEGIES FOR USING DRUG ELUTING STENTS

On the one hand, we could simply implant a drug eluting stent in every patient. This would substantially reduce the incidence of restenosis (perhaps by about two thirds, if we believe SIRIUS), and require no effort at predicting the most likely lesions to restenose. Initially, however, this approach would result in an approximately fourfold increase in stent budget (even with a discount in purchasing the drug eluting stents). A more responsible approach might be to identify a subset of lesions at the highest risk of restenosis, and target these for implantation of a drug eluting stent. This might yield a useful (though less substantial) reduction in restenosis at considerably less cost. In this paper, we aim to create a practical system for establishing different levels of risk of restenosis, and examine the impact of implanting drug eluting stents in lesions at different levels of risk.

SELECTING PRACTICAL RISK FACTORS FOR IN-STENT RESTENOSIS

Of all the risk factors for restenosis discussed above, we considered to be most useful “real world”, lesion based risk factors that were readily prospectively measurable in the majority of patients undergoing PCI. We therefore rejected IVUS based variables and confined ourselves to simple, practical, angiographic, lesion based parameters. We focused on intended stent width (=vessel reference diameter) and length (=7 mm longer than the lesion). MLD pre-procedure was rejected as being too difficult to assess “by eye”, and MLD post-procedure was rejected because it is post hoc. We obtained lesion based restenosis data from the literature. Extrapolation and estimation were necessary (because some papers considered the impact of stent length, while others considered the impact of reference diameter, and each used different stents). The only patient based variable important enough to be included in the analysis was diabetes. The risk of restenosis associated with diabetes was considered to be high enough to be included in the analysis. To establish the risk of restenosis associated with diabetes, we performed a controlled study (not published) on a group of 100 patients with diabetes and a group of 100 patients without diabetes. We found that the incidence of restenosis in the group of patients with diabetes was significantly higher (15%) than in the group of patients without diabetes (5%). We therefore included diabetes as a risk factor for restenosis.

RISK OF RESTENOSIS BY STENT SIZE

We plotted the angiographic restenosis rate, predicted by the literature, for different stent diameters (=reference diameter, divided into 0.25 and 0.5 mm groups) and lengths (=lesion + 7 mm, divided into 5 mm groups) in table 1. We made the assumptions that the percentage increase in restenosis risk with increasing stent length applies proportionately to all stent widths, and that the percentage increase with decreasing stent width applies proportionately to all stent lengths. The per cent of each size of stent used in our hospital, broken down by the same criteria of length and width, was also plotted on this table. Table 1 reveals that stents of smaller calibre and longer length are at greater risk of restenosis—for example, a 2.5 × 20 mm stent would be expected to have an angiographic restenosis rate of about 32%. Significantly, it also shows how rare restenosis is in stents which are short and large in calibre—for example, a 4 × 8 mm stent would be expected to have an angiographic restenosis rate of about 2%. In other words, we conclude that the incidence of restenosis is significantly higher in stents which are short and large in calibre, than in stents which are long and small in calibre.

APPLICATION OF DIFFERENT RESTENOSIS THRESHOLDS

Application of three arbitrary thresholds of lesion based restenosis risk (> 5%, > 10%, and > 15%) to table 1 identified stent sizes with the highest priority for implantation of a drug eluting stent. For a 15% restenosis threshold, few groups of stent sizes were found that were actually used (the others either had a low restenosis rate (large stent widths) or were hardly ever used (long stents with small widths). The potentially important sizes, for this threshold, were 2.5 × 10–

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Angiographic restenosis rates for different widths and lengths of bare stents</th>
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<tr>
<td>Stent length (mm)</td>
<td>Stent calibre (mm)</td>
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<tr>
<td>&lt;10</td>
<td>2.25</td>
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<tr>
<td>10–14</td>
<td>2.5</td>
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<tr>
<td>15–19</td>
<td>2.75</td>
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<td>20–24</td>
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<td>25–29</td>
<td>3.5</td>
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<tr>
<td>&gt;35</td>
<td>4.0</td>
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Data are derived from the literature with extrapolation and approximation wherever necessary, and assume a residual in-stent post-procedural stenosis of 10–15%.
In the next 5 years, the number of patients undergoing PCI for restenosis will increase by about 50% (with a 22.5% discount), because patients newly diagnosed in the previous five years would have an expected 10% restenosis rate, whereas 2.5% would have a moderate restenosis rate; whereas 2.5% have a high restenosis rate (4%) because, despite a high restenosis rate, they are rarely used. Addition of the figures for restenosis burden, for all stent sizes, reveals an overall, stent based, restenosis rate of 9.0%. This stent based, angiographic restenosis rate, derived from independent data, is consistent with our own, “real world”, clinical restenosis rate of 10% (for 1.6 vessels per patient and 1.1 stents per vessel).

### IMPLICATIONS OF A POLICY OF SELECTIVE DRUG-ELUTING STENT IMPLANTATION UPON RESTENOSIS

Multiplication of the per cent restenosis risk for each stent size category by the per cent of stents used in each size category (that is, the two figures in each box in table 1) yields the “restenosis burden” of each stent size. For example, the greatest restenosis burden (130/903 = 14%) is provided by stents in the category 3.0 x 15–20, because they are numerous and have a moderate restenosis rate; whereas 2.5 x 25–30 mm stents have a low restenosis burden (4%) because, despite a high restenosis rate, they are rarely used. Addition of the figures for restenosis burden, for all stent sizes, reveals an overall, stent based, restenosis rate of 9.0%. This stent based, angiographic restenosis rate, derived from independent data, is consistent with our own, “real world”, clinical restenosis rate of 10% (for 1.6 vessels per patient and 1.1 stents per vessel).

### WHAT WOULD BE THE COST OF A SELECTIVE STRATEGY?

Stent prices vary from centre to centre and from country to country. Making some reasonable assumptions, and knowing the price of the Cypher quoted to us by Cordis, together with their discount levels for increasing numbers of stents ordered, however, allows us to construct some realistic estimates of the economic impact of various thresholds of drug eluting stent implantation (second part of table 3). Application of a restenosis risk threshold of 15%, requiring 18% drug eluting stents, at the sizes listed above (with a 19% price discount) would require an increase in the stent budget of about 55%. A restenosis threshold of 10% (with a 22.5% discount) would cost an extra 110%; and a threshold of 5% (with a 22.5% discount) an extra 224%. Policies invoking progressively higher rates of implantation of a drug eluting stent would, therefore, demand steep increases in total stent costs. On the other hand, a reduction in restenosis (and re-restenosis) would reduce the number of repeat PCIs. Notionally, this would result in attenuation of the extra cost associated with using the new stents (also shown in table 3). However, in the “real world”, where expenditure is measured by the number of cases performed per year (not the number of patients), this would be less apparent, because patients with restenotic lesions would be replaced by those with de novo lesions. In these calculations, the number of other complications (CABG, myocardial infarction), which can add significantly to the costs created by an individual patient, are assumed to be few in number and equally distributed between conventional and drug eluting stents.

### IS THIS THE BEST WAY TO SPEND EXTRA MONEY?

Would the increase in stent expenditure resulting from the proposed use of a drug eluting stent be better spent on...
increasing the number of (bare stent) PCIs? For a centre performing 700 PCIs per annum, current practice, with a 10% clinical restenosis rate, allows 630 patients to be treated, the remainder of the procedures being repeat PCIs for restenosis. The data in regard to the restenosis rates quoted in table 1 are, of necessity, based upon studies performed in the mid 1990s, mostly using the Palmaz-Schatz stent and dated implantation techniques and periprocedural treatment. Nevertheless, the figures appear to be consistent with our recent, “real world”, patient based restenosis rates. Also, because no single data source contains a full spectrum of stent widths and lengths, we have had to extrapolate to construct table 1 (for example, an intrinsic “fairness”—every patient’s restenosis risk will be proportionately reduced, whether from 21% to 7% in the case of a single lesion procedure at high risk of restenosis, or from 33% to 19% in the case of a triple vessel procedure including one high and two low risk ones. On the other hand, apart from the cost of a drug eluting stent, a PCI procedure is cheaper than a CABG procedure, so some attenuation of costs for the institution may be possible if more patients are treated with PCI rather than CABG.

LIMITATIONS OF THIS PAPER

The data in regard to the restenosis rates quoted in table 1 are, of necessity, based upon studies performed in the mid 1990s, mostly using the Palmaz-Schatz stent and dated implantation techniques and periprocedural treatment. Nevertheless, the figures appear to be consistent with our recent, “real world”, patient based restenosis rates. Also, because no single data source contains a full spectrum of stent widths and lengths, we have had to extrapolate to construct table 1 (for example, in assuming that restenosis rates increase with stent length in the same proportion for different stent widths). It will also be appreciated that the data are derived from quantitative angiography rather than “eyeball” measurements (which most interventionists use). Equally, data from meticulously performed studies in the best centres may not apply so precisely to non-ideal lesions treated (in non-ideal ways) in everyday practice. We have, for instance, had to allow for the likelihood that the mean post-stent MLD is 10–15% (rather than 0%). Finally, the costs associated with drug eluting stents are likely to fall. When they do, their use will broaden, although the principles underlying the analysis we have employed in this paper will remain valid.

A DRUG ELUTING STENT FOR EVERY LESION?

The RAVEL study included only small numbers of certain important lesion subsets; only 18% of vessels were 2.5 mm, and 33% to 19% in the case of a triple vessel procedure including one high and two low risk ones. On the other hand, apart from the cost of a drug eluting stent, a PCI procedure is cheaper than a CABG procedure, so some attenuation of costs for the institution may be possible if more patients are treated with PCI rather than CABG.

MULTI-VESSEL PCI

What will be the impact of using drug eluting stents in multi-vessel disease? The main reason for not performing multi-vessel PCI is no longer safety but restenosis. The main difference between multi-vessel stenting and CABG is the increased incidence of repeat revascularisation seen in the stent group. In simple terms, the patient based restenosis rate in such circumstances is the sum of the restenosis rate of each lesion treated. A “low restenosis” stent might, therefore, tip the balance in favour of treating multi-vessel disease with PCI rather than CABG in terms of clinical outcome. However, should all three lesions (or more) be treated with drug eluting stents (at huge expense)? Or should we adhere to our principles of “lesion based” restenosis risk? Our practice will have to be the latter, at least until costs come down. That approach also has an intrinsic “fairness”—every patient’s restenosis risk will be proportionately reduced, whether from 21% to 7% in the case of a single lesion procedure at high risk of restenosis, or from 33% to 19% in the case of a triple vessel procedure including one high and two low risk ones. On the other hand, apart from the cost of a drug eluting stent, a PCI procedure is cheaper than a CABG procedure, so some attenuation of costs for the institution may be possible if more patients are treated with PCI rather than CABG.

REFERENCES

4. SIRIUS Trial. Summary of the results of the first 400 patients. Data obtained from: Cordis (Johnson and Johnson), Coronation Rd, South Ascot, SL5 9EY, UK, cordis@medgj.inl.com
A giant thrombus aspirated from a coronary artery

A 72 year old woman was admitted to the emergency department with persistent chest pain. The ECG showed ST segment elevations in the inferior leads. She was diagnosed with acute myocardial infarction. Primary percutaneous coronary intervention (PCI) was performed. At first balloon angioplasty was applied. Coronary artery angiography revealed tandem defects in the right coronary artery after the procedure. We considered that the defect indicated a thrombus and attempted to aspirate it using the Rescue percutaneous thrombectomy system (Rescue PT, Boston Scientific Corp, Maple Grove, Minnesota, USA). The Rescue PT system consists of a flexible, dual lumen Monorail 4.5 French catheter and a collection bottle with a filter for separating solid bodies from blood that contains small pieces of thrombus and atheromatous plaque. A catheter and a vacuum assistance are used to break clots into small pieces that are collected into a bottle. This catheter was first inserted past the lesion and then pulled back slowly. A giant thrombus was withdrawn while attached to the tip of the aspiration catheter through the PCI guide catheter, without entering the usual aspiration hole of the Rescue catheter from the coronary artery.

Coronary artery angiography showed the defects decreased after aspiration therapy. Thus PCI was easily successful with adjunctive stenting. When there is a large thrombus in the coronary artery, it is very useful to aspirate it. We conclude that it is necessary to undergo this pullback procedure after aspiration so that blood containing thrombus in the PCI guide catheter is withdrawn fully, to avoid the thrombus remaining in the PCI guide catheter.

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Left panel: Coronary artery angiography showing the tandem defects (white arrows) in the right coronary artery before aspiration therapy. Middle panel: Undertaking aspiration therapy. The PCI guide wire reaches the distal branch in the coronary artery. The white arrow indicates the tip of the aspiration catheter. Right panel: After aspiration therapy, the defects nearly disappear.

Aspirated giant thrombus, nearly 5.5 cm long.

Haematoxylin and eosin staining of the aspirated giant thrombus. The material comprises mostly fibrin complexes and red and white blood cells.