Coronary artery stents: identification and evaluation

J Butany, K Carmichael, S W Leong, M J Collins

First introduced in the 1980s, the coronary stent has been used to reduce the rate of arterial restenosis. Coronary stent implantation is currently a common procedure performed by interventional cardiologists, and the market for development and design is constantly expanding and evolving. This article was designed to assist pathologists in the accurate identification of coronary stents that are currently available, in addition to some that are no longer being implanted. The stents reviewed here were chosen based on frequency of use and/or occurrence in the literature. Some of the newer models have yet to undergo extensive clinical testing. The summaries accompanying each stent include concise physical descriptions and documented complications, intended to serve as a guide for the investigating pathologist.

The introduction of angioplasty for the treatment of coronary artery disease (CAD) led to the development of a completely new approach to CAD. Until 1994, the use of percutaneous transluminal coronary angioplasty (PTCA) alone as a treatment for coronary artery disease had been increasing steadily. However, the incidence of restenosis of coronary arteries was an important problem, necessitating repeated interventional procedures in 30% of patients treated with PTCA alone.

Restenosis is characterised by neointima formation resulting from the migration of proliferating medial vascular smooth muscle cells to the intima, and the synthesis of matrix components, particularly collagen, with resultant stenosis and constrictive remodelling. Coronary artery stents were developed to provide a metal scaffolding for the angioplastied vessel, in an attempt to limit negative remodelling. Sigwart et al first reported the efficacy of stents in reducing restenosis rates in 1987. By 1994, the Food and Drug Administration (FDA) had approved two stents (Gianturco-Roubin stent and the Palmaz-Schatz® stent), and additional studies by Serruys et al and Fischman et al confirmed that stenting led to improved clinical outcome over PTCA alone. Thus, PTCA with stenting has now become the preferred treatment of CAD for interventional cardiologists. The FDA has currently approved at least 16 models of stents from nine different manufacturers.

Worldwide, $2.5 billion was spent on stents in 2003, and this value is expected to increase to $4.0 billion in 2004. Although a wide array of stent designs and technologies has been introduced into the burgeoning stent market, most coronary stents can still be classified on the basis of: (1) nature of expansion or (2) stent design (table 1).

Characterisation of stents

Nature of expansion
A stent can either be self expanding or balloon expandable. Multiple studies have shown that these separate modes of expansion differ with respect to late stent expansion and neointima proliferation in stented lesions. Practising pathologists must consider such differences when deciding on the type of stent being dealt with, before making a diagnosis.

Stent design
On the basis of design, stents can be divided into three groups: coil, tubular mesh, and slotted tube. Coil stents are characterised by metallic wires or strips formed into a circular coil shape. In contrast, tubular mesh stents consist of wires wound together in a meshwork, forming a tube. Finally, slotted tube stents are made from tubes of metal from which a stent design is laser cut. Any number of stent designs can be created within a coil, tubular mesh, or slotted tube framework. Furthermore, such designs may have differences in strut pattern and width or overall stent diameter, to name just a few variations. Stent design affects many properties of the stent, such as elastic recoil and rigidity—two undesirable stent characteristics.

Previous studies have shown that differences in stent design can have a profound influence on late lumen loss and neointima proliferation, thereby affecting restenosis rates and requirements for post-procedural intervention.

Ultimately, stent design should optimise flexibility, trackability, visibility, and biocompatibility (table 2). Although stent flexibility and trackability are highly dependent on mechanical stent design, radiological visibility and host biocompatibility rely on stent material. Radiopaque gold markers are occasionally added to stents for the sole purpose of improving visualisation, helping to localise the stent during deployment at the target lesion. Although most currently approved stents are made of 316 L stainless steel, current medical practice has shown that the use of cobalt–chromium alloy is becoming more widespread.

Abbreviations: CAD, coronary artery disease; FDA, Food and Drug Administration; MSCTA, multislice computed tomography angiography; PTCA, percutaneous transluminal coronary angioplasty

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stainless steel,15 other materials have been used, such as cobalt chromium in the Multi-Link Vision™ stent, which is more radiopaque and elicits a less vigorous host response than stainless steel (Guidant pamphlet. Multi-Link VISION Coronary Stent System, 2002).

Although the use of stents reduces restenosis rates when compared with PTCA alone, restenosis still remains a serious problem in patients with CAD. The stenosis of a stented segment can be attributed primarily to the neointima.14 Stent design, surface material, and vessel damage related to the procedure have been implicated as predictors of neointima proliferation.18 19 Because antithrombotic treatment has been shown to have little influence on restenosis rates, the restenosis problem persists in many stented vessels.11

In response to this dilemma, manufacturers are now producing drug eluting stents. As of April 2004, at least 35 drug eluting stents were either in use or under investigation worldwide.7 Drug eluting stents reduce stent restenosis by delivering drugs directly to the site of vessel injury. The drug, often sirolimus or paclitaxel, acts to inhibit neointimal hyperplasia. Sirolimus (Rapamycin) reduces neointimal hyperplasia by increasing p27 concentrations, inhibiting mitogen stimulated serine/threonine kinase p70S6, and increasing retinoblastoma protein phosphophorylation and cyclin/cyclin dependent kinase complexes.18 These changes lead to the reduction of smooth muscle cell proliferation, and reduced inflammatory cell action.19 In contrast, paclitaxel is an antiproliferative, microtubule stabilising agent that has also been shown to prevent smooth muscle cell proliferation.20 Drugs are released from drug eluting stents by several methods, the most popular involving the use of a polymer matrix.21 To illustrate this, many sirolimus eluting stents are made by first coating the stent metal with a primer, followed by coating with a matrix of synthetic polymers mixed with sirolimus. Finally, a layer containing only the polymer is often sirolimus or paclitaxel, acts to inhibit neointimal hyperplasia. Sirolimus (Rapamycin) reduces neointimal hyperplasia by increasing p27 concentrations, inhibiting mitogen stimulated serine/threonine kinase p70S6, and increasing retinoblastoma protein phosphophorylation and cyclin/cyclin dependent kinase complexes.18 These changes lead to the reduction of smooth muscle cell proliferation, and reduced inflammatory cell action.19 In contrast, paclitaxel is an antiproliferative, microtubule stabilising agent that has also been shown to prevent smooth muscle cell proliferation.20 Drugs are released from drug eluting stents by several methods, the most popular involving the use of a polymer matrix.21 To illustrate this, many sirolimus eluting stents are made by first coating the stent metal with a primer, followed by coating with a matrix of synthetic polymers mixed with sirolimus. Finally, a layer containing only the polymer is sometimes added, which acts as a diffusion barrier.21 22 The CYPHER™ sirolimus eluting stent attempts to achieve 1.40 μg/cm² over the stent surface, and the release of sirolimus should last for 30 days.19 Clinical studies of drug eluting stents have produced very promising results with regard to the reduction of restenosis,20 and their use in clinical practice is becoming common.

"Drug eluting stents reduce stent restenosis by delivering drugs directly to the site of vessel injury."

Because of the large number of stents implanted each year, the pathologist will probably encounter many stents explanted at autopsy. The purpose of this review is to provide the pathologist with the ability to identify many of the commonly used and older stents in an easy to use guide. We also aim to acquaint the pathologist with the different physical characteristics of these stents, in addition to the major known complications associated with them (table 3). Many such complications of first generation stent models are well known and have been documented in previous studies. For instance, the Wallstent, Palmaz-Schatz stent and Gianturco-Roubin stent each have large bodies of literature describing their clinical complications. Longterm follow up results for many of the more contemporary stents are not yet available, so that pathological investigations of new stent models will help assess the clinical efficacy of these devices. A multitude of different stents has been available on the market in the short time since stents have gained widespread acceptance. This paper characterises 21 different stent models that are in current use, in addition to some that are no longer being implanted. We have chosen this list of stents based on frequency of occurrence in the literature and hence popularity in clinical trials, but we also include some new and alternative stent models that have yet to undergo extensive testing. They have been divided into non-drug eluting (18) and drug eluting (3) stents, and are listed alphabetically by model name. We have no bias for or against any of the devices listed or a particular stent manufacturer.

**Table 1** Stent classifications

<table>
<thead>
<tr>
<th>Self expanding</th>
<th>Balloon expandable</th>
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<tbody>
<tr>
<td>Tubular mesh</td>
<td>Tubular mesh</td>
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<tr>
<td>Wallstent</td>
<td>Driver</td>
</tr>
<tr>
<td></td>
<td>Coil</td>
</tr>
<tr>
<td></td>
<td>Gianturco-Roubin II</td>
</tr>
<tr>
<td></td>
<td>Wiktor</td>
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<tr>
<td></td>
<td>Slotted tube</td>
</tr>
<tr>
<td>beStent</td>
<td>BiodivViso®</td>
</tr>
<tr>
<td>beStent2</td>
<td>OY/SV/AS</td>
</tr>
<tr>
<td>Bx VELOCITY®</td>
<td>(CYPHER™)</td>
</tr>
<tr>
<td>Express™</td>
<td>(TAXUS™ Express)</td>
</tr>
<tr>
<td>JOSTENT Flex</td>
<td>Multi-Link PENTA™</td>
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<tr>
<td></td>
<td>/RX™/Vision™</td>
</tr>
<tr>
<td>NIR™ (TAXUS™)</td>
<td>(NIRFLEX™)</td>
</tr>
<tr>
<td>Palmaz-Schatz</td>
<td>153/154/Crown</td>
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</tbody>
</table>

**Table 2** Desirable stent characteristics

<table>
<thead>
<tr>
<th>Low crossing profile</th>
<th>High flexibility</th>
<th>High stent/host biocompatibility</th>
<th>High radial strength</th>
<th>Low metallic surface area</th>
<th>Favourable radiographic properties</th>
<th>Good trackability (ability to conform to tortuous vessels)</th>
<th>Easy deployment</th>
</tr>
</thead>
</table>

**Table 3** Common stent complications

<table>
<thead>
<tr>
<th>In stent restenosis via neointimal hyperplasia</th>
<th>Side branch occlusion</th>
<th>Poor stent/host biocompatibility</th>
<th>Poor radiographic properties</th>
<th>Post-implantation stent foreshortening</th>
<th>Post-implantation stent expansion</th>
<th>Post-implantation stent migration</th>
<th>Stent related thrombosis</th>
</tr>
</thead>
</table>

www.jclinpath.com
Physical characteristics
Serpentine design composed of sinusoidal ring modules linked via sigmoidal link elements. The laser cut stainless steel has two radiopaque gold markers incorporated at the proximal and distal ends. No weld points.15

Stent material
316 L stainless steel.15

Radiographical characteristics
A single gold marker at each end provides the means for easier radiographical localisation.

Additional comments
First human implant in 1996.15 The radial strength of the slotted tube is optimised by an orthogonal locking mechanism and there is virtually no shortening after expansion.15 This particular stent was found to display a very low elastic recoil percentage (2.35% under 0.2 bar pressure) among 23 tested stents.15 In addition, the late minimal lumen diameter of the stent is similar to the initial dilated diameter.11 The stent was shown experimentally to collapse at a pressure of 0.70 atm.23

beStent2 Model
The successor of the beStent is available mounted on Rapid Exchange with or without Perfusion and Over-the-Wire balloon platforms.22

Type
Balloon expandable; slotted tube.

Technical information
Manufactured and sold by Medtronic.22

Size range
Length, 9–30 mm; diameter, 2.5–4.0 mm; maximum expansion, 4.5 mm.23

Strut dimensions
Radial, 0.095 mm wide × 0.085 mm thick; longitudinal, 0.07 mm wide × 0.085 mm thick.22

Physical characteristics
Flexible radial “S” crowns and longitudinal “V” crowns crossing at a junction that rotates during expansion with almost no foreshortening (fig 1). The laser cut stainless steel has two radiopaque gold markers incorporated at the proximal and distal ends.22

Stent material
316 L stainless steel.22

Radiographical characteristics
A single gold marker at each end provides means for easier radiographical localisation.

FDA approval
The beStent2 stent was approved in 2000.22

BiodivYsio® Model
The models include the Open Cell (BiodivYsio OC), Small Vessel (BiodivYsio SV), and the now unavailable Added Support (BiodivYsio AS) designs.22

Type
Balloon expandable; slotted tube.

Technical information
Manufactured and sold by Biocompatibles Ltd (an Abbott Laboratories company, Abbott Park, Illinois, USA).22

Size range
- OC: length, 8–28 mm; diameter, 2.75–4.0 mm.
- SV: length, 7–18 mm; diameter, 2.0–2.5 mm.
- AS: length, 8–15 mm; diameter, 2.75–4.0 mm.22

Strut dimensions
- OC: 0.05–0.08 mm wide × 0.09 mm thick.
- SV: 0.03–0.05 mm wide × 0.06 mm thick.
- AS: 0.05–0.08 mm wide × 0.09 mm thick.23

Physical characteristics
The BiodivYsio OC and SV stents share an identical strut design of alternating sinusoidal rings consisting of either rectangular or rounded edges. The rings are linked via two turn (S shaped) articulations (fig 2). The BiodivYsio AS stent design includes additional articulations in between those present in the OC and SV models. All BiodivYsio stents are coated with phosphorylcholine (PC) TechnologyTM to increase stent biocompatibility by mimicking the lipid head group of erythrocyte membranes.24

Stent material
316 L stainless steel coated with polymers containing phosphorylcholine (PC TechnologyTM).22

Additional comments
The phosphorylcholine coating of the BiodivYsio is designed to reduce thrombosis by lowering platelet adhesion and at the same time decreasing the adhesion of proteins.25

FDA approval
The BiodivYsio AS stent was approved in September 2000.3

Bx Velocity® Model
The Bx Velocity is also available in a drug eluting form (CYPHER®TM) or a drug coated form with covalently bound heparin (Hepacoat®TM). Available mounted on Rapid Exchange or Over-the-Wire balloon platforms.22

Type
Balloon expandable; slotted tube.

Technical information
Manufactured and sold by Cordis (Miami Lakes, Florida, USA; a Johnson and Johnson Company).22

Figure 1 beStent2 by Medtronic.
The gold radiopaque marker is indicated by a white arrow. Original image taken from website (http://local.teklan.net/medim/eng/products_details.asp?id=64).

Figure 2 BiodivYsio OV/SV by Biocompatibles Ltd. Original image taken from website (http://www.user.cityline.ru/~drgtech/Biocompatibles.htm).
Size range
Length, 8–33 mm; diameter, 2.25–5.0 mm.

Strut dimensions
Radial, 0.13 mm wide × 0.14 mm thick.

Physical characteristics
Sinusoidal ring strut modules linked by flexible “N” shaped flex segments that reduce foreshortening during expansion (< 1.5 mm) (fig 3).

Stent material
316 L stainless steel.

Potential complications
The ISAR-STEREO-2 trial showed that the thick strut (140 μm) Bx Velocity stent has a high rate of restenosis: 31.4% of stented vessels had ≥ 50% diameter stenosis at follow up, compared with 17.9% for the thin strut (50 μm) ACS Multi-Link RX™ stent.

FDA approval
The first human implants were performed in 1999 (France and Italy). In 2000 the drug coated Bx Velocity (Bx Hepacoat) became the first FDA approved drug coated stent.

Driver Model
The Driver model includes the Zipper MX. Available mounted on Discrete Technology™, Rapid Exchange, and Over-the-Wire balloon platforms.

Type
Balloon expandable; tubular mesh.

Technical information
Manufactured and sold by Medtronic AVE. The Driver is a variation of the previous S7 design. They share an identical sinusoidal strut pattern but the Driver has a smaller strut diameter of 0.09 mm (Medtronic Vascular Driver website, 2004. Medtronic website).

Size range
Length, 9–30 mm; diameter, 2.8–4.0 mm; maximum expansion, 4.4 mm.

Strut dimension
Diameter, 0.091 mm.

Physical characteristics
The smooth, edgeless, modular design consists of 10 crown sinusoidal rings linked via 1.0 mm elements (fig 4).

Stent material
Cobalt alloy finished with a chrome rich oxide. The cobalt alloy used in the Driver is more flexible and stronger than 316 L stainless steel, allowing thinner stent struts to be used.

Radiographical characteristics
Cobalt alloy provides increased radiographical visualisation compared with stainless steel stents.

FDA approval
The Driver stent was approved in October 2003 with no reports of implantation outcomes beyond 270 days.

Express™ Model
The Express model includes the Express™ and TAXUS Express™ (paclitaxel drug eluting version). Available mounted on Rapid Exchange, Over-the-Wire, and Monorail™ balloon platforms.

Type
Balloon expandable; slotted tube.

Technical information
Manufactured and sold by Boston Scientific (Natick, Massachusetts, USA). All Express stents share an identical strut pattern regardless of delivery system or drug eluting properties.

Size range
Length, 8–32 mm; diameter, 2.25–5.0 mm.

Strut dimensions
Radial, 0.09 mm wide × 0.13 mm thick; longitudinal, 0.06 mm wide × 0.13 mm thick.

Physical characteristics
The modular ring strut pattern consists of two separate module designs: short, narrow sinusoidal Micro™ elements linked via straight articulations to long, wide sinusoidal Macro™ elements (Boston Scientific Express website, 2004. Boston Scientific website) (fig 5).

Stent material
316 L stainless steel.

FDA approval
Approved in September 2002.

Gianturco-Roubin II
Type
Balloon expandable; coil.

Radiographical characteristics
Cobalt alloy provides increased radiographical visualisation compared with stainless steel stents.
Technical information
Manufactured and sold by Cook Inc (Bloomington, Indiana, USA).\textsuperscript{15}

Size range
Length, 12–60 mm; diameter, 2.5–5.0 mm.\textsuperscript{15}

Strut dimensions
0.172 mm wide × 0.076 mm thick.\textsuperscript{15, 27}

Physical characteristics
Flat wire coil attached to a single longitudinal strut (fig 6).

Stent material
316 L stainless steel.\textsuperscript{15}

Potential complications
Studies report a restenosis rate of up to 57%,\textsuperscript{27} and have also shown that the restenosis rate is greater than that of the Palmaz-Schatz stent.\textsuperscript{28} The increased restenosis rate is attributed to greater stent recoil and increased levels of tissue prolapse seen with the Gianturco-Roubin II stent compared with the Palmaz-Schatz stent. Among 22 stents tested in vitro for elastic recoil (under 0.2 bar pressure), the Gianturco-Roubin II had the highest elastic recoil (mean, 18.34%; SD, 2.41%), and may need to be overdilated by the interventional cardiologist to compensate for this.\textsuperscript{11}

Additional comments
The Gianturco-Roubin II stent showed a higher procedural success rate than conventional angioplasty with a significantly higher event free survival (83% vs 65%) at one year, and better clinical outcomes in a randomised multicentre study.\textsuperscript{19}

FDA approval
The premier Gianturco-Roubin model was the first coronary stent approved by the FDA in June 1993.\textsuperscript{30} The Gianturco-Roubin II stent was approved in June 1996.\textsuperscript{51, 5}

JOSTENT\textsuperscript{TM} Flex
Model
The JOSTENT models include the older JOSTENT Plus. In addition, the JOSTENT Coronary Stent Graft model is composed of a layer of PTFE sandwiched between two JOSTENT Flex stents.\textsuperscript{22}

Type
Balloon expandable; slotted tube.

Technical information
Originally manufactured and sold by JOMED (an Abbott Laboratories company) (Abbott Vascular Devices website, 2004).

Size range
Length, 9–32 mm; diameter, 2.0–5.0 mm.\textsuperscript{22}

Strut dimension
0.09 mm wide × 0.09 mm thick.\textsuperscript{22}

Physical characteristics
The JOSTENT Flex sinusoidal ring modules are linked via alternating flexible spiral and slightly curved link elements (fig 7). The struts are also smooth with rounded edges.\textsuperscript{22}

Stent material
316 L stainless steel.\textsuperscript{22}

Potential complications
The JOSTENT Coronary Stent Graft has been reported to cause a high side branch occlusion rate of up to 18.6%, in some instances resulting in major complications, such as myocardial infarction.\textsuperscript{31} Restenosis at the proximal and distal edges of the JOSTENT Coronary Stent Graft remains a serious limitation of this stent, despite low observed neointimal proliferation at the stent centre.\textsuperscript{29} Studies have shown that the lumen can be completely obscured by artefacts when being visualised by multislice computed tomography angiography (MSCTA).\textsuperscript{14}

Additional comments
Studies have shown that the stent is not superior to conventional angioplasty for small coronary arteries.\textsuperscript{33} Stent graft systems such as the JOSTENT Coronary Stent Grafts may prove to be the preferred treatment for perforations in the coronary arteries, although more studies are needed.\textsuperscript{31}

FDA approval
The JOSTENT Coronary Stent Graft was approved in January 2001.\textsuperscript{1, 22}

Multi-Link PENTA\textsuperscript{TM} Type
Balloon expandable; slotted tube.

Technical information
Manufactured and sold by Guidant (Indianapolis, Indiana, USA).\textsuperscript{22}

Size range
Length, 8–38 mm; diameter, 2.75–4.0 mm; maximum expansion, 4.5 mm.\textsuperscript{22}

Strut dimensions
Radial, 0.097 mm wide × 0.124 mm thick; longitudinal, 0.076 mm wide × 0.091 mm thick.\textsuperscript{22}

Figure 6 Gianturco-Roubin II by Cook Inc. The longitudinal strut is indicated by a black arrow. Original image taken from website (http://www.cis.tugraz.at/biomech/Proj1.htm).

Figure 7 JOSTENT Flex by JOMED. Original image taken from website (http://www.ami.co.il/html/cardio/in-cardio/JostentFlexMaster.htm).

Figure 8 Multi-Link PENTA by Guidant. The five turn ACCESS-LINK element is shown. Original image taken from website (http://www.guidant.jp/products/penta/reach.shtml).
Physical characteristics
Corrugated Multi-Link five crown rounded corner zigzag ring module design linked via five turn ACCESS-LINK™ elements (fig 8).

Stent material
316 L stainless steel.22

Potential complications
Side branch occlusion immediately after stenting is not uncommon (up to 25%) because of the specific slotted tube design.34 The Multi-Link stent has a large mounted profile (1.53 mm) compared with many other stents, such as JOSTENT (1.15 mm), and ranked highest in a study of profile size, among eight other stents.12

Additional comments
In clinical use it was shown to have a higher implantation success rate compared with the Palmaz-Schatz stent,35 and also produced lower in stent tissue proliferation as a result of its corrugated ring design.17 In a comparison with seven other slotted tube stents, the Multi-Link stent ranked second in a longitudinal flexibility test.12 In addition, reopening of stenosed side branches by balloon dilatation is feasible using the Multi-Link, because a 3.5 mm balloon inflated to 12 atm through the struts of the Multi-Link can achieve a 94% balloon expansion, compared with 80% for the Palmaz-Schatz and 60% for the NIR stent.36

FDA approval
The Multi-Link PENTA model was approved in 2001.52

Multi-Link RX

Model
The models include the Multi-Link RX PIXEL™ and ULTRA™ used in small and large coronary vessels, respectively.22

Type
Balloon expandable; slotted tube.

Technical information
Manufactured and sold by Guidant.22

Size range
- PIXEL: length, 8–28 mm; diameter, 2.0–2.5 mm; maximum expansion, 3.0 mm.22
- ULTRA: length, 13–38 mm; diameter, 3.5–5.0 mm; maximum expansion, 5.5 mm.22

Strut dimensions
- PIXEL: 0.096 mm wide × 0.099 mm thick.
- ULTRA: 0.1 mm wide × 0.13 mm thick.22


Physical characteristics
Corrugated Multi-Link five crown rounded corner zigzag ring module design linked via straight link elements (fig 9A,B). The PIXEL model contains an additional straight link element between modules22 (fig 9A).

Stent material
316 L stainless steel.22

Additional comments
Scaffolding struts are thinner compared with the Bx Velocity (50 μm v 140 μm), which is said to account for the lower rate of restenosis (18% v 31%).26

FDA approval
The Multi-Link RX ULTRA model was approved in 2000.3 22

Multi-Link Vision

Type
Balloon expandable; slotted tube.

Technical information
Manufactured and sold by Guidant (Guidant pamphlet. Multi-Link VISION Coronary Stent System, 2002).

Size range
Length, 8–38 mm; diameter, 2.75–4.0 mm.

Strut dimensions
0.081 mm thick.

Physical characteristics
Corrugated Multi-Link five crown rounded corner zigzag ring module design linked via single turn link elements (fig 10).

Stent material
Cobalt chromium alloy.

Radiographical characteristics
The low nickel super alloy is more radiopaque than stainless steel.

Additional comments
The cobalt chromium alloy is stronger than stainless steel; therefore, the stent is thinner and made with less material, and occupies less vessel space than the average stainless steel stent. The enhanced biocompatibility of cobalt chromium compared with 316 L stainless steel was proposed in a study in which vessels stented with the Multi-Link VISION had a low lumen loss, comparable to that of the thinner strut Multi-Link device.37

FDA approval
The Multi-Link VISION model was approved in July 2003.3

NIR™

Model
Additional models include the NIROYAL™ stent, a NIR stent that is plated with gold.

Type
Balloon expandable; slotted tube.

Figure 10 Multi-Link Vision by Guidant. Original image taken from website (http://www.guidant.com/products/producttemplates/vi/US_VISION_Intro.shtml).
**Technical information**
Manufactured and sold by Medinol® (Jerusalem, Israel).22

**Size range**
Length, 9–32 mm; diameter, 2.0–5.0 mm.32

**Strut dimension**
0.1 mm thick.23

**Physical characteristics**
Sinusoidal ring modules linked via curved link elements (fig 11).

**Stent material**
316 L stainless steel. The NIROYAL model has an outer layer of gold.22

**Potential complications**
The NIR stent has been described as being very stiff compared with other stents.12 As with other gold plated stents, the NIROYAL is prone to a higher rate of restenosis by neointimal hyperplasia compared with bare metal stents.18

Using MSCTA, the lumen of a vessel treated with the NIROYAL stent was completely obstructed radiographically,32 making clinical angiography follow up difficult using MSCTA. Balloon expansion of side branches through the struts of the NIR stent may be difficult, because a 3.5 mm balloon could only be inflated to 60% by 12 atm pressure through NIR struts.36

**Additional comments**
Believed to perform equivalently to the Palmaz-Schatz stent, but also designed to provide increased longitudinal flexibility and trackability during deployment.23 The stent was shown experimentally to collapse at a pressure of 1.05 atm.23

**Figure 11**
![NIR (TAXUS) by Medinol. Original image taken from website](http://www.hgcardio.com/ptca.htm)

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**NIRFLEX™ Model**
Model premounted on a Rapid Exchange delivery system

**Type**
Balloon expandable; slotted tube.

**Technical information**
Manufactured and sold by Medinol.23

**Size range**
Length, 9–32 mm; diameter, 2.5–4.0 mm; maximum expansion, 5.75 mm.23

**Strut dimension**
0.1 mm thick.23

**Physical characteristics**

**Stent material**
316 L stainless steel.22

**FDA approval**
The NIRFLEX model was approved in October 2003.5

**Palmaz-Schatz 153 Model**
Available mounted on the PAS delivery system.15

**Figure 12**

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**Type**
Balloon expandable; slotted tube.

**Technical information**
Manufactured and sold by Cordis (a Johnson and Johnson Company).15

**Size range**
Length, 15 mm; diameter, 3.0–5.0 mm.15

**Strut dimensions**
0.07 mm thick.22

**Physical characteristics**
The Palmaz-Schatz 153 model design consists of expandable diamond mesh slotted tubes (fig 13) articulated at specific points by straight link elements.

**Stent material**
316 L stainless steel.15

**Potential complications**
Only one length (15 mm) was manufactured. More than one stent was needed for the treatment of long lesions.25 Delivery of the long stent was hindered (or made impossible) by its rigidity.40 The treatment of lesions involving major side branches is not recommended using this particular stent because of its unique tight metal strut design.38 The high radiolucency of the Palmaz-Schatz stent makes the accurate assessment of stent diameter by angiography very difficult.41 Increased neointimal proliferation and dissection at the proximal and distal ends of the stented segment result from the high pressure required for stent deployment (20 atm).11

**Additional comments**
Implanted in over 600 000 patients, the Palmaz-Schatz stent was found to perform as well as its early successors,27 exhibiting a low percentage of subacute thrombosis.42 It was first used to introduce the concept of the balloon expandable stent in 1985.43 The stent was shown experimentally to collapse at a pressure of 0.65 atm.23

**FDA approval**
The first approval of the Palmaz-Schatz stent was in August 1994, further approval was granted to the medium and long models in June 1997.5

**Palmaz-Schatz 154 Model**
The models include the Spiral Coronary Palmaz-Schatz stent and its successor the Improved Spiral Coronary Palmaz-Schatz stent. Available mounted on the PowerGrip™ Rapid Exchange delivery system.15

**Type**
Balloon expandable; slotted tube.

**Figure 13**
Technical information
Manufactured and sold by Cordis (a Johnson and Johnson Company).\textsuperscript{15}

Size range
- Spiral: length, 8–18 mm; diameter, 3.0–5.0 mm.\textsuperscript{15}
- Improved Spiral: length, 15 mm; diameter, 3.0–5.0 mm.\textsuperscript{15}

Strut dimensions
0.09 mm thick.\textsuperscript{22}

Physical characteristics
Expandable diamond mesh identical to the 153 model (fig 13), articulated at specific points via spiral link elements. The initial Spiral model was modified to create the Improved Spiral by cutting the stent with 10 rows and five articulations rather than 12 rows and six articulations.\textsuperscript{15}

Stent material
316 L stainless steel.\textsuperscript{15}

Additional comments
Found to possess twice the mechanical strength of the PS-153 precursor and was shown experimentally to collapse at a pressure of 1.26 atm.\textsuperscript{23}

Palmaz-Schatz Crown
Model
Available mounted on the PowerGrip Rapid Exchange delivery system.\textsuperscript{15}

Type
Balloon expandable; slotted tube.

Technical information
Manufactured and sold by Cordis (a Johnson and Johnson Company).\textsuperscript{15}

Size range
Length, 15–30 mm; diameter, 3.0–4.0 mm.\textsuperscript{15}

Strut dimensions
0.07 mm thick.\textsuperscript{22}

Physical characteristics
The slotted tube is designed with a continuous sinusoidal pattern showing no articulation sites.

Stent material
316 L stainless steel.\textsuperscript{15}

Additional comments
The Palmaz-Schatz Crown ranked highest in a stiffness test among 13 other stents tested.\textsuperscript{44}

Potential complications
Difficulties in precise positioning as a result of foreshortening of up to 20% on expansion.\textsuperscript{27} In addition, the mesh design is known to cause side branch entrapment.\textsuperscript{15} Because of the self-expanding design, there is a chronic radial pressure exerted on the vessel walls that can exacerbate neointima proliferation. It has been stated that this proliferation is balanced by post-procedural stent expansion.\textsuperscript{16}

Additional comments
First patented in 1982,\textsuperscript{43} the Wallstent was the first clinically tested stent (1986).\textsuperscript{15} It was found to maintain the same radiographical characteristics over a 10 year period of implantation, with no calcification or lesion progression in those who were restenosis free after the first six months.\textsuperscript{45} In addition, a study showed an 18% loss in diameter of the stented segment over a five year period followed by virtually no loss (0.3%) in diameter over the subsequent five years.\textsuperscript{45}

FDA approval
The first human implant was performed in 1986\textsuperscript{15}; FDA approval was given for the Boston Scientific Magic Wallstent in September 1993.\textsuperscript{8}

Wiktor Model
The Wiktor stent is also available with Hepamed\textsuperscript{TM} coating.\textsuperscript{15}

Type
Balloon expandable; coil.

Technical information
Manufactured and sold by Medtronic Interventional Vascular.\textsuperscript{15}

Size range
Length, 16 mm; diameter, 3.0–4.5 mm.\textsuperscript{15}

Strut dimensions
Diameter, 0.127 mm.\textsuperscript{15}

Physical characteristics
Single wire helical sinusoidal coil.

Stent material
Tantalum.\textsuperscript{15}

Potential complications
Using MSCTA, the Wiktor stent showed strong artefacts and the lumen of the stent was totally obstructed—this was attributed to the tantalum material from which the stent is made.\textsuperscript{32} Only negligible artefacts are expected when using magnetic resonance imaging.\textsuperscript{12} Protrusion of the intima into the stented lumen may be more common in the Wiktor stent
than in slotted tube stents because of its open wire design. Insertion of an intravascular ultrasound catheter into the Wiktor stent must be performed with extreme care, because the increased radiopacity of the tantalum material can lead to stent damage during intravascular ultrasound. Among 22 stents tested in vitro for elastic recoil (under 0.2 bar pressure), the Wiktor had the third highest elastic recoil percentage.

Additional comments
The open design of the Wiktor may allow more access to side branches and provide the stent with additional flexibility and conformity. In a three point bend test, the unexpanded Wiktor had a mean (SD) measured stiffness of 0.5 (0.2) g force/mm, the lowest among 13 stents tested, conferring high flexibility to this stent.

FDA approval
The Wiktor stent was approved in June 1997.

DRUG ELUTING STENTS
CYPHER
Model
The CYPHER drug eluting stent is available mounted on Over-the-Wire or Rapid Exchange balloon platforms (CYPHER website, 2004. Cordis (J&J) website).

Type
Balloon expandable; slotted tube.

Technical information
Manufactured and sold by Cordis (a Johnson and Johnson Company).

Size range
Length, 18–33 mm; diameter, 2.50–3.50 mm.

Strut dimensions
Radial, 0.13 mm wide × 0.14 mm thick.

Physical characteristics
Maintains all the characteristics of the Bx Velocity, sinusoidal ring strut modules linked by flexible N shaped Flex Segments (fig 3).

Eluted drug
Sirolimus. Initial drug concentration of 1.4 μg/mm² of stent, eluted over 30 days.

Stent material
316 L stainless steel.

Potential complications
Reports of post-implantation thrombosis (including up to 20% resulting in death) to the FDA within 10 months after its approval led to the release of an official FDA statement (February 2004). Professionals are asked to adhere to the manufacturer’s recommendations when intending to use this stent, including matching stent and vessel diameters, implanting only within previously untreated vessels, and maintaining an adequate antiplatelet regimen.

FDA approval
The CYPHER stent became the first drug eluting stent approved by the FDA in April 2003.

TAXUS™
Type
Balloon expandable; slotted tube.

Technical information
Manufactured and sold by Medinol.

Size range
Length, 15 mm; diameter, 3.0 and 3.5 mm.

Strut dimension
0.1 mm thick.

Eluted drug
Paclitaxel. The initial paclitaxel concentrations are 1.02 μg/mm² (85 μg total) and 2.04 μg/mm² (171 μg total) for the slow and fast release formulations, respectively.

Physical characteristics
Maintains all the characteristics of the NIR stent, sinusoidal ring modules linked via curved link elements (fig 11).

Stent material
316 L stainless steel.

Additional comments
Studies showed a reduced occurrence of myocardial infarction nine to 12 months after implantation when using the TAXUS model compared with the bare metal stent. In addition, 70% lower angiographic restenosis was seen, together with decreased target lesion revascularisation, nine and 12 months after implantation, respectively.

TAXUS Express²
Type
Balloon expandable; slotted tube.

Technical information

Size range
Length, 8–32 mm; diameter, 2.25–5.00 mm.

Strut dimensions
Radial, 0.091 mm wide × 0.13 mm thick; longitudinal, 0.06 mm wide × 0.13 mm thick.

Eluted drug
Paclitaxel. The slow release formulation has an initial drug concentration of 1.0 μg/mm² of stent.

Physical characteristics
Maintains the characteristics of the Express models. The modular ring strut pattern consists of two separate module designs: short, narrow sinusoidal Micro elements linked via straight articulations to long, wide sinusoidal Macro elements (Boston Scientific Express website, 2004. Boston Scientific website) (fig 5).

Stent material
316 L stainless steel.

FDA approval
The TAXUS Express² drug eluting stent was approved in March 2004.
REFERENCES

15. Serruys PW, Rensing B. Handbook of coronary stents. London: Dunitz; Distributed in the USA by Fulfilment Center Taylor and Francis (Kentucky, USA), 1997.
22. Serruys PW, Rensing B. Handbook of coronary stents. London: Dunitz; Distributed in the USA by Fulfillment Center Taylor and Francis (Kentucky, USA), 2000, viii, 364.