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Review Article

Are Biodegradable Third Generation Drug Eluting Stents the Answer to Instant Restenosis?

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18 **ABSTRACT**
19

The third generation biodegradable Drug Eluting Stent (DES) are being evaluated and being introduced in clinical practice. They have been designed to overcome limitations associated with durable polymer and a persistent metallic stent scaffold which could be related to late target lesion revascularization (TLR) and very late stent thrombosis (VLST). Although a recent pooled data analysis found that biodegradable polymer stents were superior for TLR and VLST compared with first generation Sirolimus Eluting Stent (SES), superiority has not been demonstrated against second generation Everolimus eluting stents (EES) and is yet to be conclusively proven randomized trials. This paper reviews the key features, recent trial data, and future directions of the third generation of DES technology including stents with fully biodegradable scaffolds, stents with biodegradable polymer, and polymer free stents.

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23

24 **INTRODUCTION** Interventional cardiology is currently in the process of refining the third generation of DES
25 technology. It incorporates a broad mix of technologies ranging from incremental improvements in existing stent
26 scaffolds, antiproliferative coats, polymer free, biodegradable polymer coated scaffolds, fully biodegradable scaffolds,
27 newer nano-material coatings and stem cell therapy.

28

29 Compared with first generation DES, the second generation stents have advantages like having thinner struts and
30 increased flexibility, more biocompatible polymers and new generation antiproliferative agents [1,2]. Even the second
31 generation DES are not free from disadvantages as the persistent presence of a stent scaffold or polymer beyond its
32 short-term function is related to late target lesion revascularisation (TLR) and very late stent thrombosis (ST). The two
33 year pooled results from the SPIRIT II, III, IV and COMPARE trials prove that Everolimus eluting stents (EES) have a
34 superior safety and efficacy profile compared with first generation paclitaxel eluting stents (PES) because of lower
35 rates of myocardial infarction (MI) (RR, 0.57; 95% CI, 0.45–0.73), ST (RR, 0.35; 95% CI, 0.21–0.60) and ischemia
36 driven TLR(RR, 0.59;95%CI, 0.47–0.73) [3-5]. Neither EES nor zotarolimus eluting stents (ZES) have demonstrated
37 superior clinical outcomes to first generation sirolimus eluting stents (SES) [6-9].

38

39 Major concern with second generation DES is very late stent thrombosis (VLST) rates beyond one year. The
40 pathogenesis of late restenosis and stent thrombosis in second generation DES include neointimal hyperplasia,
41 persistent inflammation of the vessel wall, in-stent neoatherosclerosis, uncovered struts and/or polymers with
42 secondary stent malapposition and stent fracture [10-13].

43

44 The Bern-Rotterdam cohort followed 4212 patients treated with EES for four years and reported a definite or probable
45 ST rate of 6.3% and a VLST rate of 2.0%. Although the 2% VLST rate is statically significant and lower than the
46 corresponding VLST rate for first generation PES (4.0%, $p < 0.0001$) and SES (2.8%, $p = 0.02$), it represents an
47 ongoing 0.67% annual risk of ST after one year [14]. The HORIZONS-AMI [15] trial at three years, LEADERS [16] and
48 SYNTAX [17, 18] trials at four years and the SIRTAX LATE [81] trial at five years demonstrated similar annual VLST
49 rates of 0.6–0.85% for PES and SES.

50

51 Long term efficacy in terms of repeat revascularization rates, TLR incidence rate and late lumen loss (LLL) are other
52 major limitations of second generation DES. Four year repeat revascularization rates of up to 28.8% have been
53 reported for first generation PES in high risk patients undergoing PCI for left main stem and triple vessel disease [17].
54 Five year SPIRIT III data of 669 low risk patients treated with EES revealed an annual TLR incidence rate of 1.3%
55 beyond one year with TLR increasing from 3.5% at one year to 8.6% at five years [19]. Second generation DES are
56 also associated with a persistent increase in late lumen loss (LLL). In SPIRIT II EES cohort the mean in-stent LLL
57 increased from $0.17\pm 0.32\text{mm}$ to $0.33\pm 0.37\text{mm}$ [20] while in the ISAR-4 EES cohort [21,22] it increased from
58 $0.14\pm 0.41\text{mm}$ to $0.29\pm 0.51\text{mm}$ between six and 24 months interval. Additional limitations with current generation DES
59 include restrictions to non-invasive imaging with CT and MRI, difficulties with future surgical and transcatheter
60 revascularization, long term disruption of native vascular fluid dynamics and vasoreactivity, chronic inflammation,
61 delayed endothelialization and the need for six or more months of dual antiplatelet therapy (DAPT) [23-28].

62
63 The ultimate dream would be to develop a stent system which has best combination of metallic alloys and/or polymers
64 with all desirable properties favourable combination-drug eluting capabilities. This paper reviews the key features,
65 recent trial data, and future directions of the third generation of DES technology including stents with fully
66 biodegradable scaffolds, stents with biodegradable polymer, and polymer free stents.

67 68 **Fully Biodegradable Scaffolds**

69 Fully biodegradable scaffolds aim to combine the advantages of the first and second generation of DES while
70 additionally targeting their disadvantages and limitations. They provide a stable vascular scaffold in the short term,
71 thereby minimizing constrictive remodeling , preventing restenosis due to vascular recoil, and loose intimal dissection
72 flaps [29-31]. The fully biodegradable scaffolds score over the older generation stents by reducing the limitations
73 including but not limited to long-term in-stent restenosis and stent thrombosis associated with a permanent metallic
74 scaffold.

75
76 They have been associated with the development of a homogenously thickened neointima, suggestive of a thicker,
77 more stable fibrous cap [12], potential for expansive arterial remodeling and a return of normal vasomotion [32],
78 theoretical decrease in paradoxical peri-stent vasoconstriction[33], facilitating improved non-invasive CT and MRI
79 imaging, wider future transcatheter and/or surgical revascularization options, freedom from jail branch obstruction, less
80 impediment to vascular growth in the pediatric population and limit the need for prolonged DAPT [32,34,45].

82 Metallic biodegradable scaffolds can be magnesium or iron based. Magnesium has a shorter degradation period of
83 four to 12 months compared with four or more years for iron [37,38]. A polymer coat is used to contain and control the
84 release of an antiproliferative agent. These are designed to biodegrade by Krebs cycle into carbon dioxide and water
85 over six to 24 months, after the antiproliferative agent has been fully released [33,36].

86 87 **ABSORB BVS**

88 **ABSORB A and ABSORB B** : The bioabsorbable everolimus-eluting stent system ABSORB BVS (Abbot Vascular,
89 Santa Clara, CA, USA). The ABSORB BVS stent is based on a poly-L-lactic acid (PLLA) scaffold with a poly-D,L-
90 lactide (PDLLA), everolimus impregnated polymer coat. The device has been assessed in two small single arm
91 industry sponsored non-randomized trials, ABSORB A and ABSORB B. Both studies were restricted to lesions with a
92 RVD of 2.5–3mm and length less than 14mm. Patients received a minimum of six months DAPT post stent insertion.

93
94 Five year data from the ABSORB A trial, a 30 patient study using the first iteration BVS 1.0 [34,39,40], revealed a
95 MACE rate of 3.4%, representing a single non-q wave MI at 46 days, and TLR and ST rates of 0%. LLL increased to
96 $0.48\pm 0.28\text{mm}$ at 24 months. Mean in-stent LLL was $0.43\pm 0.37\text{mm}$ at six months which was largely attributed to
97 scaffold recoil.

98 Optical coherence tomography (OCT) at 24 months showed a smooth endoluminal lining appearance with virtually
99 indiscernible struts suggested almost complete stent biodegradation.

100 Intravascular ultrasound (IVUS) results suggested expansile arterial remodeling with the minimum lumen area (MLA)
101 increasing from $3.92\pm 0.98\text{mm}^2$ to $4.34\pm 1.74\text{mm}^2$ from six to 24 months. There was evidence of a return of normal
102 arterial vasomotion at two years with five of nine patients demonstrating arterial vasodilatation on acetylcholine
103 administration [41].

104
105
106 ABSORB B trial assessed the BVS 1.1 stent, a revision of the BVS 1.0 designed to improve radial support beyond six
107 months and allow stent storage at room temperature in 100 patients [36]. The 24 month MACE rate was 9%,
108 comprising a TLR rate of 6% and non-q-wave MI rate of 3%. There were no ST events [40]. LLL increased from
109 $0.19\pm 0.18\text{mm}$ at six months to $0.27\pm 0.25\text{mm}$ at 12 months and was stable at $0.27\pm 0.20\text{mm}$ out to 24 months [42].
110 Between six and 24 months, mean lumen area by IVUS increased from 6.36mm^2 to 6.85mm^2 with a small increase in
111 MLA from 5.12mm^2 to 5.13mm^2 . Vasoreactivity was demonstrated at 12 months on administration of
112 methylegonovine and acetylcholine [40].

113

114 **ABSORB EXTEND & ABSORB II** : Two larger trials with less restrictive inclusion criteria are currently enrolling
115 patients. ABSORB EXTEND is a 1000 patient multinational single arm trial and ABSORB II is a 500 patient RCT
116 comparing the ABSORB BVS against the second generation DES, Xience PRIME (Abbot Vascular, Santa Clara, CA,
117 USA) [43,44]. Six month data from the first 200 patients enrolled in the ABSORB EXTEND trial revealed a MACE rate
118 of 2.5% comprising an MI rate of 2% and TLR rate of 0.5% [45].

119

120 Despite significant recent interest in biodegradable scaffolds, clinical and trial experience is limited. Only two devices,
121 the bioabsorbable everolimus-eluting stent system ABSORB BVS (Abbot Vascular, Santa Clara, CA, USA) and the
122 Igaki-Tamai stent (Kyoto Medical Planning Co., Kyoto, Japan) have had trial results published in peer reviewed
123 journals. Both of these stents have the European C.E. mark although the Igaki-Tamai is currently only used in
124 peripheral arteries. There is no randomized data and trials have less restrictive inclusion criteria with respect to
125 reference vessel diameter (RVD) and lesion length. Complex lesions including left main coronary artery (LMCA), left
126 main stem, ostial lesions, saphenous vein graft disease and bifurcations have been excluded [32,34,46-49].

127

128 **Igaki-Tamai stent**: The Igaki-Tamai stent was the first ever fully biodegradable stent. The device was also based on a
129 PLLA polymer scaffold but required contrast heated to 80 °C to self expand. It was first implanted in 1999 and 10 year
130 data for 50 patients was reported in 2012 [1, 46]. The study was non-randomised and industry sponsored. At 10 years,
131 rates of TLR, ST and MI were 28%, 4% and 8% respectively. Mean in-stent LLL reduced from 0.91±0.69mm at six
132 months to 0.59±0.50mm at three years while MLA increased from 3.64±1.68mm² to 5.18±2.09mm² over the same
133 period, suggestive of expansile arterial remodeling. At three years, IVUS echogenicity had returned to
134 pre-stent levels, indicating complete stent degradation [46].

135

136 **ReZolve stent** (Reva Medical, San Diego, CA, USA): The ReZolve device is based on a tyrosine polycarbonate rather
137 than PLLA scaffold and has the advantage of being radio-opaque [33]. It elutes sirolimus and is being assessed in the
138 RESTORE single arm clinical trial which is currently enrolling a target cohort of 50 patients [49]. An earlier iteration of
139 the stent was assessed in 27 patients in the 2008 RESORB trial which reported a six month TLR rate of 67% and 30
140 day q-wave-MI rate of 7% [1,35].

141

142 **DESolve stent** (Elixir Medical Corporation, Sunnyvale, CA, USA): DESolve has a PLLA scaffold with a myolimus
143 eluting PLA coat. Six month clinical data of a 16 patient FIM trial revealed a TLR rate of 7%, MI and cardiac death rate

144 of 0% and LLL of 0.19 ± 0.19 mm [47]. A larger trial with the DESolve Nx novolimus eluting stent is underway with a
145 target enrolment of 120 patients [50].

146
147 **ART bioresorbable stent** (Arterial Remodelling Technologies, Paris, France): The ART non-drug eluting
148 bioresorbable stent is based on a PLA scaffold and has recently started enrolling patients in the ARTDIVA FIM trial
149 [51].

150
151 **DREAMS drug eluting absorbable metal stent** (Biotronik, Berlin, Germany) is the only metal biodegradable stent
152 currently undergoing trial assessment. It comprises a magnesium alloy scaffold with a paclitaxel impregnated PLGA
153 coat. It was evaluated in the BIOSOLVE-1 46 patient FIM trial which reported a 12 month TLR rate of 4.7%, MI rate of
154 2.3% and no ST events. Mean LLL was 0.64 ± 0.50 mm at six months and 0.52 ± 0.39 at 12 months [48].

155
156 Biodegradable polymeric scaffolds have a number of limitations including but not limited to thicker struts with an
157 increased crossing profile, limited post-dilatation options which mandates quantitative vessel sizing, radio-lucency with
158 more challenging angiographic visualization.

159 There is also a scarcity of trials testing complex anatomy and challenging lesion subsets including ostial, bifurcation
160 and heavy calcified disease [24]. Potential risk like strut fracture secondary to post-dilatation was observed in one
161 patient at 46 days post stent insertion in the ABSORB A trial. It was hypothesized that fracture resulted from the
162 3.0mm×12mm stent being over expanded post dilation with a 3.5mm×9mm balloon [52].

163
164 Biodegradable polymer DES have demonstrated non-inferiority to both first and second generation DES for safety and
165 efficacy. Although a recent pooled analysis of the ISAR-TEST 3, ISAR-TEST 4 and LEADERS trial data found that
166 biodegradable polymer stents were superior for TLR and VLST compared with first generation SES, superiority has not
167 been demonstrated against second generation EES and is yet to be proven in any single substantial randomized trial
168 [53].

169 **Non-polymeric Drug Eluting Metallic Stents**

170
171 Non-polymeric DES comprises of a metal alloy scaffold directly impregnated with an anti-proliferative agent. The
172 absence of a polymer coat offers a theoretical basis to minimize the duration of DAPT in patients with a high bleeding
173 risk to one month or less based on the BMS guidelines [27] while still providing the established late safety of a BMS

174 and the antiproliferative effects comparable to polymer based DES. Table 1 gives a brief outline of non-polymeric
175 Drug eluting metallic stents.

176
177 **LEADERS-FREE trial** is comparing the BioFreedom with the Gazelle BMS in 2500 randomized patients at high risk of
178 bleeding with the primary endpoints of non-inferiority for MACE and superiority for clinically driven TLR. Importantly,
179 patients will be treated with only one month of DAPT [54].

180 Yukon SES (Translumina, Hechingen, Germany) has been examined in two independently funded, assessor blinded,
181 randomized trials, the ISAR-TEST and ISAR-TEST 3. The ISAR-TEST trial included 450 patients across two centers
182 and reported non-inferiority of the Yukon SES compared with the durable polymer-based TAXUS PES [55] for six
183 month in-stent LLL ($0.48\pm 0.61\text{mm}$ vs $0.48\pm 0.58\text{mm}$, $p = 0.98$) and death & MI (4.4% vs 4.0% , $p = 0.81$). Despite the
184 encouraging early results, it performed poorly in the subsequent three-arm ISAR-TEST 3 study, failing to demonstrate
185 non-inferiority with the first generation Cypher stent in 650 patients for the primary endpoint of in-stent LLL at six to
186 eight months ($0.47\pm 0.56\text{mm}$ vs $0.17\pm 0.45\text{mm}$ vs $0.23\pm 0.46\text{mm}$, $p = 0.94$) [56]. At two years, however, there was no
187 difference for a composite endpoint of death or MI (7.0% vs 6.9% vs 6.4% $p = 0.97$); for TLR (13.9% vs 8.4% vs 10.4 p
188 = 0.19); or for ST (1.0% vs 0.5% vs 1.0%, $p = 0.82$) [57].

189
190 A non-polymeric dual-DES utilises the Yukon stent platform, but incorporates a second antiproliferative agent –
191 probucol, a potent liposoluble antioxidant which reduces neointimal hyperplasia. The stent has been examined in the
192 independently funded, assessor blinded, multicentre randomized ISAR-TEST 2 and ISAR-TEST 5 trials.

193
194 **ISAR-TEST 2 trial** compared this dual-DES (n=333) with the first generation Cypher SES (n=335) and the second
195 generation Endeavour zotarolimus eluting stent (ZES) (n=339)(Medtronic Inc., Santa Rosa, CA, USA) [58] with
196 promising results. The dual-DES was superior to the Endeavour stent at six months for binary angiographic restenosis
197 (dual-DES 11.0% vs ZES 12.0%, $p=0.68$ vs SES 19.3%, $p = 0.002$), in-stent LLL $0.23\pm 0.50\text{mm}$ vs 0.24 ± 0.51 ($p =$
198 0.78) vs $0.58\pm 0.55\text{mm}$, ($p < 0.001$), and TLR (6.8% vs 7.2% ($p = 0.83$) vs 13.6%, $p = 0.001$)); its results were
199 comparable with the Cypher stent. At two years, there was no significant difference in clinical outcomes including
200 cardiovascular death or MI (dual-DES 7.8% vs ZES 9.2% vs SES 10.2%, $p = 0.88$); TLR 7.7% vs 10.7% vs 14.3% ($p =$
201 0.009); BR 13.9% vs 18.6% vs 20.9% ($p = 0.047$) and LLL 0.30 ± 0.54 vs 0.35 ± 0.60 vs 0.57 ± 0.57 ($p < 0.001$) [59].

204 **ISAR-TEST 5 trial** compared the dual-DES (n=2002) with the Resolute ZES (n=1000) (Medtronic Inc., Santa Rosa,
205 CA, USA) and demonstrated the dual-DES to be non-inferior with regards to the BR 13.3% vs 13.4% ($p = 0.95$); LLL
206 0.31 ± 0.58 vs 0.30 ± 0.56 ($p = 0.50$) and primary endpoint of MACE at 12 months (13.1% vs 13.5%, $p = 0.74$) and ST
207 1.1% vs 1.2% ($p = 0.80$) [60].

208
209 **BioFreedom BES** (Biosensors Europe SA, Morges, Switzerland) The Biolimus-A9 eluting BioFreedom stent is
210 currently being assessed in a first in man (FIM) randomized, three arm trial of 182 patients [61]. It was shown to be
211 non-inferior to the TAXUS Liberte for mean in-stent LLL at 12 months (0.17 ± 0.22 mm vs 0.35 ± 0.22 mm, $p = 0.001$) and
212 for MACE at two years (6.8% vs 10.0%, $p =$ not significant).

213
214 **VESTASync SES** (MIV Therapeutics, Atlanta, GA, USA): This SES is currently being assessed in the small, industry
215 funded, double blinded, multicentre VESTASync II study ($n = 75$; NP $n=50$ vs BMS $n = 25$). It has been shown to be
216 non-inferior to the GenX durable polymer stent (MIV Therapeutics, Atlanta, GA, USA) with regards to in-stent late
217 lumen loss at nine months (0.39 ± 0.20 mm vs 0.74 ± 0.52 mm, $p = 0.03$) [62].

218 **Biodegradable Polymer Drug Eluting Stents**

219
220
221 Durable polymers of first and second generation DES remain within the coronary artery environment long after their
222 purpose is fulfilled, and have deleterious effects by causing inflammation, delayed vascular healing, as well as
223 providing a platform for accelerated neoatherosclerosis [1,63]. They are also considered to play a pivotal role in late
224 stent thrombosis (ST) [10-13]

225
226 Biodegradable polymers have been the focus of active research and development. The scientists continue to be
227 challenged by issues like composition, degradation time of the polymer, biocompatibility, interaction and
228 pharmacokinetic profile of the antiproliferative agents. Table 2 gives a brief outline of biodegradable Polymer DES.

229 **BioMatrix** (Biosensors Inc., Newport Beach, CA, USA)

230
231 Biolimus-A9 is a sirolimus analogue with extreme lipophilicity that enables targeted tissue uptake and minimizes
232 systemic exposure. It has been combined with an abluminal polylactic acid (PLA) polymer that biodegrades within six
233 to nine months, eluting 45% of the antiproliferative agent within the first 30 days.

235 **LEADERS study** was an industry funded, multicentre, non-inferiority powered randomized controlled trial (RCT) that
236 examined the use of Biomatrix-Flex BES against the durable polymer first generation Cypher SES (Cordis, Miami
237 Lakes, FL, USA) [16, 64, 65]. 1707 patients (BES n = 857 vs SES n = 850) were enrolled and 96.5% were followed to
238 five years. Patients as well as assessors of angiographic films and staff involved with clinical follow-up were blinded to
239 the assigned stent. Operators involved with stent insertion were not blinded. Non-inferiority was demonstrated for the
240 primary endpoint of major adverse cardiovascular events (MACE) at nine months (9.2% vs 10.5%, $p = 0.39$) and at
241 five years (22.3% vs 26.1%, $p = 0.071$). The definite VLST at five years was also found to be significantly low (0.66%
242 vs 2.5% $p = 0.003$).

243
244 **COMFORTABLE AMI trial** was an industry funded, assessor blinded, multicentre study of 1161 patients randomized
245 to either the BioMatrix- Flex or the Gazelle BMS (Biosensors Europe SA, Morges, Switzerland) (BES n = 575 vs BMS
246 n = 582). It showed that Biomatrix- Flex BES had lower rates of definite VLST from one to five years compared with the
247 Cypher SES (0.66% vs 2.5%, $p = 0.003$) [65]. Its efficacy and safety has also been validated in primary PCI for acute
248 ST elevation myocardial infarction (STEMI) [66]. This showed superiority for MACE at 12 months in favor of the BES
249 (4.3% vs 8.7%, $p = 0.004$). There was no significant difference in the rate of definite or probable late ST (2.5% vs
250 3.7%, $p = 0.25$) at 12 months.

251
252 **Nobori** (Terumo, Somerset, NJ, USA) stents

253 The Nobori BES has also reported encouraging results in both the NOBORI 1 and NOBORI CORE trials [67,68] and
254 more recently in the ongoing, large, industry funded, randomized, all-comers COMPARE II trial (BES $n = 1795$ vs EES
255 $n = 912$) [69]. At 12 months, the stent was non-inferior for MACE compared with a durable polymer EES (5.2% vs
256 4.8%, $p = 0.69$) and had very low but similar rates of definite or probable late ST (0.8% vs 1.0%, $p = 0.58$). BASKET-
257 PROVE II completed recruitment of 2400 all-comer patients randomized to either the Nobori BES, the Xience Prime
258 EES, or the PRO-Kinetic BMS in 2012[70]. They will be followed over five years for MACE and other clinical end
259 points.

260
261 **NOBORI 2 and eNOBORI** are two large, prospective, single-arm, multicenter, registries that enrolled 3067 and 7750
262 patients respectively, out of which 248 and 703 were STEMI patients. All adverse events were adjudicated by an
263 independent clinical event committee in NOBORI 2, while adjudication in eNOBORI (including stent thrombosis) is
264 ongoing. At 1-month, there were no MIs observed. Total of 5 patients died because of cardiac reasons (0.9%) and one
265 TLR (0.17%) and one TVR (0.4%) were found. The TLF rate was 1.0%. In the cohort of patients followed at 3-year, 2

266 patients suffered a cardiac death (0.8%), 10 had an MI (4.0%) and TLF rate was 6.1%. A total of 96% of the patients
267 were angina free. Regarding stent thrombosis (ST), occurring up to 3 years, total of 4 cases have been detected
268 (1.6%), out of which 3 cases were subacute (1.2%) and one case of late ST (0.4%). There was no very late ST
269 detected at 3 years follow up. [71]

270 271 **Supralimus** (Sahajanand Medical)

272 PAINT trial, an industry funded, multicentre, unblinded trial with 274 randomised patients to the Supralimus stent, the
273 Infinnium bioabsorbable polymer PES (Sahajanand Medical Technologies Pt. Ltd., India), or the Millennium Matrix
274 BMS (Sahajanand Medical Technologies Pt. Ltd., India) groups (SES $n = 106$ vs PES $n = 111$ vs BMS $n = 57$)
275 examined the Supralimus stent [72,73]. The polymers included PLLA, PLGA, PLC and PVP. Clinical events were
276 adjudicated by an independent committee. At nine months angiographic follow-up, the Supralimus stent had
277 significantly less in-stent LLL than the BMS (0.32 ± 0.43 mm vs 0.90 ± 0.45 mm, $p < 0.001$) and the Infinnium stent
278 (0.32 ± 0.43 mm vs 0.54 ± 0.44 mm, $p = 0.001$). The Supralimus stent also had superior rates of MACE compared with
279 the BMS at 12 months (8.6% vs 21.1%, $p = 0.01$) and three years (12.5% vs 33.3%, $p < 0.01$).

280 281 282 **Excel** (JW Medical System, Weihai, China)

283 The industry funded CREATE study was a large single-arm, multicentre, prospective registry of 2077 patients
284 implanted with the Excel stent. It reported a MACE rate of 4.5% and definite or probable ST in 1.0% of patients at three
285 year follow-up, half of which occurred beyond one year [74,75].

286 287 **SYNERGY** (Boston Scientific)

288 Everolimus Eluting Stents As durable polymer EES have become the most widely used DES worldwide, it is not
289 surprising that advancement continues in this direction through clinical investigation of the Synergy stent (Boston
290 Scientific Corp., Natick, MA, USA). Clinical experience with the stent is limited but the industry funded, assessor
291 blinded EVOLVE randomized trial recently demonstrated non-inferiority for its primary endpoint of in-stent late loss at
292 six months when compared with the PROMUS Element durable polymer EES (Boston Scientific Corp., Natick, MA,
293 USA) (0.10 ± 0.25 mm vs 0.15 ± 0.34 mm, $p = 0.19$) [76]. MACE was also comparable between the stents.

294 295 **PLATINUM Study**

296 In this prospective single blind trial (NCT00823212) 1,530 patients undergoing PCI of 1 or 2 de novo native lesions
297 were randomized at 132 worldwide sites to CoCr-EES (n = 762) or PtCr-EES (n = 768). It was found that novel PtCr-
298 EES was noninferior to the predicate CoCr-EES for TLF, with nonsignificant differences in measures of safety and
299 efficacy through 12-month follow-up after PCI. The 12 month TLF was 2.9% in CoCr-EES and 3.4% in PtCr-EES (p
300 noninferiority =0.001, p superiority = 0.60). By intention-to-treat, there were no significant differences between CoCr-
301 EES and PtCr-EES in the 12-month rates of cardiac death or MI (2.5% vs. 2.0%, p = 0.56), TLR (1.9% vs. 1.9%, p =
302 0.96), TLF (3.2% vs. 3.5%, p = 0.72), or Academic Research Consortium definite or probable stent thrombosis (0.4%
303 vs. 0.4%, p = 1.00). [77]

304
305 **The JACTAX Liberte Paclitaxel Eluting Stents (PES)** (Boston Scientific, Natick, MA, USA) is the effort to advance
306 the initial success of the first generation TAXUS PES into a third generation bioabsorbable polymer DES. The industry
307 funded, single centre OCTDESI pilot study examined 60 patients randomized to either a JACTAX high dose stent
308 (n=20), a JACTAX low dose stent (n=21), or a TAXUS Liberte stent (n=19), with percentage of strut coverage as the
309 primary endpoint. Angiographic endpoints were assessed by an independent core laboratory. At six months, the
310 results were comparable across the three stents for both percentage of uncovered struts (7.0±12.2% vs 4.6±7.3% vs
311 5.3±14.7%, p = 0.81) and for in-stent late loss (0.25±0.32mm vs 0.39±0.43mm vs 0.24±0.44mm, p = 0.39) [78].

312
313 **Combo stent** (OrbusNeich, Fort Lauderdale, FL, USA)

314 The Combo stent is a novel biodegradable polymer SES that utilizes endothelial progenitor cell (EPC) capture
315 technology in addition to low-dose abluminal sirolimus. This EPC capture technology is a luminal coating of immobile
316 CD34 antibodies and aims to capture EPCs that differentiate into endothelial cells to form mature endothelial coverage
317 of stent struts. Early data from the small, industry funded, non-randomized REMEDEE trial showed non-inferiority for
318 its primary angiographic endpoint of in-stent late loss at nine months when compared with the TAXUS Liberte durable
319 polymer PES (0.39±0.45mm vs 0.44±0.56mm, p = 0.55) [79]

320
321 **ISAR-TEST 4** was an independently funded, assessor blinded trial that randomized 2603 patients from two centers to
322 a novel, non-commercially available biodegradable polymer SES or a durable polymer DES, either the first generation
323 Cypher SES or the second generation Xience EES [80]. Non-inferiority of the biodegradable polymer SES was
324 demonstrated for the primary endpoint of MACE at 30 days (4.4% vs 4.5%, p = 0.87) and at one year (13.8% vs
325 14.4%, p = 0.66), as well as for definite or probable late ST at one year (1.0% vs 1.5%, p = 0.29).

326

327 **Fourth Generation Stents**

328 Wayne et al successfully modified a standard bioresorbable terpolymer with the covalent incorporation of lovastatin, as
329 seen on NMR, into a backbone comprised of lactide, glycolide, ε-caprolactone, and lovastatin (60 : 15 : 10 : 15 parts
330 by weight), respectively. Thus a fourth-generation bioresorbable stent was produced that has the potential to deliver
331 two drugs to the site of the procedure-related vessel lumen injury. [82]

333 **Ongoing Clinical trials**

334 The database of the clinicaltrials.gov was searched for biodegradable coronary stents and 14 open trials were
335 identified. Table 3 gives the brief outline of identifier number, design types, primary outcomes and current recruitment
336 status of the “open studies”.

338 **Duration of Dual Antiplatelet Therapy (DAPT)**

339
340 Multiple studies have shown that early discontinuation of clopidogrel after the DES as one of the strong predictors for
341 stent thrombosis (83,84) and hence prolonged dual antiplatelet therapy (DAPT) is historically recommended to
342 prevent stent thrombosis (83,85). However Long term DAPT does not come without complications. There have
343 been reports from several trials of the zotarolimus-eluting stent (Endeavor [E-ZES], Medtronic, Santa Rosa, California)
344 that have shown beneficial efficacy and safety, despite a relatively short duration of DAPT (86-88). Kim et al
345 showed using optical coherence tomography that there is sufficient strut coverage following implantation with the E-
346 ZES as early as 3 months post-procedure (89). A recent registry study with 661 low-risk patients who received DAPT
347 for 3 months following E-ZES implantation has shown favorable long-term clinical outcomes and lower incidence of
348 stent thrombosis after cessation of clopidogrel 3 months post-intervention (90).

349
350 **RESET Trial** (NCT01145079) randomly assigned 2,117 patients with coronary artery stenosis into 2 groups according
351 to DAPT duration and stent type: 3-month DAPT following Endeavor zotarolimus-eluting stent (E-ZES) implantation (E-
352 ZES+ 3-month DAPT, n=1,059) versus 12-month DAPT following the other DES implantation (standard therapy,
353 n=1,058). E-ZES+3-month DAPT was noninferior to the standard therapy with respect to the occurrence of the primary
354 endpoint (difference: 0.0%; 95% confidence interval [CI]: -2.5 to 2.5; p 0.84; p < 0.001 for noninferiority). The
355 composite rates of any death, myocardial infarction, or stent thrombosis were 0.8% and 1.3%, respectively (difference:
356 -0.5%; 95% CI: -1.5 to 0.5; p 0.48). The rates of stent thrombosis were 0.2% and 0.3%, respectively (difference: -0.1%;
357 95% CI: -0.5 to 0.3; p 0.65) without its further occurrence after cessation of clopidogrel in the E-ZES+3-month DAPT

358 group. The rates of target vessel revascularization were 3.9% and 3.7%, respectively (difference: 0.2%; 95% CI: -2.3
359 to 2.6; p = 0.70). (REal Safety and Efficacy of a 3-month dual antiplatelet Therapy following E-ZES implantation
360 [RESET]. [91])

362 DISCUSSION

363 The field of interventional cardiology is experiencing a great deal of cutting edge research especially in order to reduce
364 the disadvantages of second generation stents. Although the second generation stents have come a long way and
365 offer significant benefits including a large evidence base, good deliverability and operator familiarity, long term definite
366 or probable ST rates of up to 0.67% per annum and TLR rates of 1.3% per annum suggest a scope for improvement.

367
368 Although a recent pooled analysis of the ISAR-TEST 3, ISAR-TEST 4 and LEADERS
369 trial data found that biodegradable polymer stents were superior for TLR and VLST compared with first generation
370 SES, superiority has not been demonstrated against second generation EES and is yet to be proven in any single
371 substantial randomized trial [53]. Trials to date have been small, non- randomized and exclusively industry funded.
372 Early trial data has shown the promise of longer term expansile remodeling and restoration of vasoreactivity but the
373 clinical implication of this is uncertain and there is no large study to backup this hypothesis. Moreover, deliverability,
374 expansion constraints together with an absence of data in complex lesions suggests the need for further research.

375
376 Two larger trials with broader inclusion criteria are currently underway and should provide a greater indication of
377 performance of third generation stents. There is a need for developing a technology which can provide excellent
378 efficacy and safety, deliverability in broad range of clinical settings, minimal limitations on non-invasive imaging and
379 future revascularization procedures, and limit the need for prolonged DAPT.

381 **Table 1: Non-polymeric drug eluting stents.**

Study (n) Current status	Stent (Manufacturer)	Drug	Results/endpoints

<p>ISAR-TEST [52] (NP $n = 225$ vs PES $n = 225$)</p> <p>Completed</p>	<p>Yukon (Translumina)</p>	<p>Sirolimus</p>	<p>9 months</p> <p>LLL 0.48 ± 0.61 vs 0.48 ± 0.58 ($p = 0.98$)</p> <p>Death and MI 4.4% vs 4.0% ($p = 0.81$)</p>
<p>ISAR-TEST 2 [55] (DD $n = 333$ vs SES $n = 335$ vs ZES $n = 339$)</p> <p>Completed</p>	<p>Dual DES</p>	<p>Sirolimus and probucol</p>	<p>6–8 months</p> <p>BR 11.0% vs 12.0% ($p = 0.68$) vs 19.3% ($p = 0.002$)</p> <p>LLL 0.23 ± 0.50 vs 0.24 ± 0.51 ($p = 0.78$) vs 0.58 ± 0.55 ($p < 0.001$)</p> <p>TLR 6.8% vs 7.2% ($p = 0.83$) vs 13.6% ($p = 0.001$)</p> <p>2 years</p> <p>Death and MI 7.8% vs 10.2% vs 9.2% ($p = 0.61$)</p>

			<p>TLR 7.7% vs 10.7% vs 14.3% ($p = 0.009$)</p> <p>BR 13.9% vs 18.6% vs 20.9% ($p = 0.047$)</p> <p>LLL 0.30±0.54 vs 0.35±0.60 vs 0.57±0.57 ($p < 0.001$)</p>
<p>ISAR-TEST 3 [53] (NP $n = 201$ vs BP $n = 202$ vs PP $n = 202$)</p> <p>Completed</p>	<p>Yukon (Translumina)</p>	<p>Sirolimus</p>	<p>6–8 months</p> <p>LLL 0.47±0.56 vs 0.17±0.45 vs 0.23±0.46 ($p = 0.94$)</p> <p>2 years</p> <p>TLR 13.9% vs 8.4% vs 10.4% ($p = 0.19$)</p> <p>Death and MI 7.0% vs 6.9% vs 6.4% ($p =$ 0.97)</p> <p>ST 1.0% vs 0.5% vs 1.0% ($p = 0.82$)</p>
<p>ISAR-TEST 5 [57] (DD $n = 2002$ vs ZES</p>	<p>Dual DES</p>	<p>Sirolimus and</p>	<p>6–8 months</p>

<p><i>n</i> = 1000)</p> <p>Completed</p>		<p>probucol</p>	<p>BR 13.3% vs 13.4% (<i>p</i> = 0.95)</p> <p>LLL 0.31±0.58 vs 0.30±0.56 (<i>p</i> = 0.50)</p> <p>1 year</p> <p>MACE 13.1% vs 13.5% (<i>p</i> = 0.74)</p> <p>ST 1.1% vs 1.2% (<i>p</i> = 0.80)</p>
<p>VESTASync II [58] (NP <i>n</i>=50 vs BMS <i>n</i> = 25)</p> <p>Ongoing</p>	<p>VESTASync (MIV Therapeutics)</p>	<p>Sirolimus</p>	<p>9 months</p> <p>LLL 0.39±0.20 vs 0.74±0.52 (<i>p</i> = 0.03)</p>
<p>FIM [59] (NP SD <i>n</i>=60 vs PES <i>n</i> = 60)</p> <p>Ongoing</p>	<p>BioFreedom (Biosensors)</p>	<p>Biolimus A9</p>	<p>1 year</p> <p>LLL 0.17±0.22 vs 0.35±0.22 (<i>p</i> = 0.001)</p> <p>2 years</p> <p>MACE 6.8% vs 10.0% (<i>p</i> = NS)</p>

383 BR, binary restenosis; DD, non-polymeric dual DES; FIM, first-in-man; LLL, in-stent late lumen loss (mm); MACE, major adverse
 384 cardiovascular events; NP, non-polymeric DES; ST, definite/probable stent thrombosis; TLF, target lesion failure; PP, permanent
 385 polymer; BP, biodegradable polymer; SD, standard dose; NS, not significant.

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389 **Table 2. Biodegradable polymer drug eluting stents.**

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Study (n) Current status	Stent (Manufacturer)	Drug	Polymer type	Results/endpoints
LEADERS [37] (BES n = 857 vs SES n = 850) Completed	BioMatrix (Biosensors)	Biolimus A9	Abluminal PLA	5 years MACE 22% vs 26% (p = 0.07) Definite VLST 0.66% vs 2.5% (p = 0.003)
COMFORTABLE AMI [38] (BES n = 575 vs BMS n = 582)	BioMatrix (Biosensors)	Biolimus A9	Abluminal PLA	1 year MACE 4.3% vs 8.7% (p = 0.004) ST 2.5%

Completed				vs 3.7% ($p = 0.25$)
COMPARE II [41] (BES $n = 1795$ vs EES $n = 912$) Ongoing	Nobori (Terumo)	Biolimus A9 Abluminal	PLA	1 year MACE 5.2% vs 4.8% ($p = 0.69$) ST 0.8% vs 1.0% ($p = 0.58$)
BASKETPROV E- II [42] (target $n = 2400$, BES vs EES vs BMS) Recruiting	Nobori (Terumo)	Biolimus A9 Abluminal	PLA	Primary endpoint of MACE at 2 years
PAINT [46] (SES $n = 106$ vs PES $n = 111$ vs BMS $n = 57$)	Supralimus (Sahajanand Medical)	Sirolimus	PLLA, PLGA, PLC, PVP	9 months LLL 0.32 ± 0.43 vs 0.54 ± 0.44 vs 0.90 ± 0.45 ($p < 0.001$)

Completed				3 years MACE 12.5% vs 16.6% vs 33.3% ($p < 0.01$)
CREATE registry [45] ($n = 2077$) Completed	Excel (JW Medical System)	Sirolimus	PLA	3 years MACE 4.5% ST 1.0%
REMEDEE [49] (SES $n = 124$ vs PES $n = 59$) Ongoing	Combo (OrbusNeich)	Sirolimus + EPC	Abluminal	9 months LLL 0.39 ± 0.45 vs 0.44 ± 0.56 ($p =$ 0.55)
EVOLVE [43] (SYNERGY	SYNERGY (Boston Scientific)	Everolimus	PLGA Rollcoat Abluminal	6 months LLL 0.10 ± 0.25

<p><i>n</i>=94 vs SYNERGY half-dose <i>n</i>=99 vs PROMUS Element <i>n</i> = 98) Completed</p>				<p>vs 0.13±0.26 vs 0.15±0.34 (paired <i>p</i> = ns) TLF 2.2% vs 4.1% vs 3.1% (<i>p</i> = NS) ST 0.0% vs 0.0% vs 0.0%</p>
<p>OCTDESI [50] (JACTAX <i>n</i>=20 vs JACTAX low-dose <i>n</i>=21 vs TAXUS <i>n</i> = 19) Ongoing</p>	<p>JACTAX Liberte (Boston Scientific)</p>	<p>Paclitaxel</p>	<p>Juxtaposed Abluminal Coating technology</p>	<p>6 months USS 7.0±12.2% vs 4.6±7.3% vs 5.3±14.7% (<i>p</i> = 0.81) LLL 0.25±0.32 vs 0.39±0.43 vs 0.24±0.44 (<i>p</i> = 0.39)</p>

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LLL, in-stent late lumen loss (mm); MACE, major adverse cardiovascular events; PLA, poly-L-lactide; PLC, 75:25 poly-L-lactide-co-caprolactone; PLGA, 50:50 poly-D,L-lactide-co-glycolide; PLLA, poly-L-lactic acid; PVP, polyvinyl pyrrolidone; ST, definite/probable stent thrombosis; TLF, target lesion failure; USS, uncovered stent struts; NS, not significant.

Clinical Trial (NCT identifier)	Official Title	Study Type	Primary Outcome Measures	De sig ne d as Sa fet y Iss ue	Esti mat ed Stu dy Co mpl etio n Dat e	Curr ent Stat us
DESTINY TRIAL (Inspiron x Biomatrix) (NCT0185608 8)	Stents Coated With the Biodegr adable Polymer on Their Faces and Elution of Sirolimu s Ablumin ais Versus	Interventional Allocation: Rand omized	Lumen Loss [Time Fra me: 9 months after the procedure]	Ye s	Feb ruar y 201 8	Recr uitin g Last verifi ed: May 201 3

	Elution Biolimus for the Treatme nt of Coronar y Lesions Again - Random ized Destiny					
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<p>PONTINA (NCT01060306)</p>	<p>Prospective Optical coherence Tomography Evaluation of neointimal Coverage of a biodegradable Polymer-based Drug-eluting Stent</p>	<p>Observational: Case Control Time Perspective: Prospective</p>	<p>Assessment of neointimal coverage of the biodegradable polymer-based Biolimus A9-eluting stent (Biomatrix stent) after full drug elution and polymer biodegradation [Time Frame: 6 months]</p>	<p>No</p>	<p>January 2011</p>	<p>Unknown Last verified: January 2010</p>
<p>BESS (NCT01268371)</p>	<p>Comparison of Biolimus-eluting Biodegradable Polymer</p>	<p>Interventional. Allocation: Randomized</p>	<p>MACE</p>	<p>Yes</p>	<p>July 2015</p>	<p>Recruiting Last verified:</p>

	Everoli mus- eluting and Sirolimu s-eluting Coronar y Stents					April 201 3
BIO-RESORT (NCT0167480 3)	Compari son of BIOdegr adable Polymer and DuRabl E Polymer Drug- eluting Stents in an All COmeR s PopulaT ion: Random ized Multicen	Interventional. Allocation: Rand omized	Target vessel failure (TVF) [Time Fra me: 1 year]	Ye s	Nov emb er 201 6	not yet ope n for parti cipa nt recr uitm ent Last verifi ed: Aug ust 201 2

	ter Trial in an All Comers Populati on Treated Within the NeThErl ands 3 (TWEN TE 3)					
EVOLUTION (NCT0082577 3)	A Random ized Study to Evaluat e Safety and Efficacy of the ExcelT M Sirolimu s Eluting Stent With a Biodegr adable	Interventional. Allocation: Rand omized	Ischemia- driven Target Vessel Failure which is a composite of cardiac death, myocardial infarction (Q and non-Q wave) and target vessel revasculari	Ye s	April 201 4	Recr uitin g Last verifi ed: Jan uary 200 9

	<p>Polymer Versus SirOlimus ELUting Stent With Non-Biodegradable Polymer in the Treatment of Patlents With de nOvo Coronar y Artery LesioNs</p>		<p>zation (TVR) at 12 months. [Time Fra me: 12 months]</p>			
<p>OCTOBER(N CT01012583)</p>	<p>Optical Coheren ce TomOgr aphy Assess ment of the Excel</p>	<p>Observational: Case Control Time Perspectiv e: Prospective</p>	<p>To quantitate the presence of neointimal stent strut coverage at 6 month</p>	<p>Ye s</p>	<p>Oct ober 201 0</p>	<p>Unk now n Last verifi ed: Nov emb er</p>

	Drug-Eluting Stent With Biodegradable Polymer vs. the Cypher Drug-Eluting Stent With Permanent Polymer		via Optical Coherence Tomography follow-up. [Time Frame: 6 month]			2009
Pro-HOPE (NCT01880879)	A Prospective Multicenter Trial Evaluating Helios Biodegradable Polymer Sirolimu	Interventional Single Group Assignment	1 year incidence of target lesion [Time Frame: 1year]	No	January 2015	Recruiting Last verified: January 2013

	s-eluting Stent Safety and Effective ness in Treatme nt of Coronar y Artery Disease					
Evaluate Safety And Effectiveness Of The Tivoli® DES and The Firebird2® DES For Treatment Coronary Revasculariza tion (NCT0168138 1)	A Prospec tive, Open Label, Random ized Study to Evaluat e Safety And Effective ness Of The Tivoli® Biodegr adable Polymer	Interventional. Allocation: Rand omized	Ischemia- driven Target Lesion Failure (TLF) which is a composite of cardiac death, myocardial infarction (Q and non-Q wave) and target lesion revasculari	Ye s	Sep tem ber 201 8	Recr uitin g Last verifi ed: Nov emb er 201 2

	<p>Rapamycin-Eluting Stent and The FIREBIRD2® Rapamycin-Eluting Coronary CoCr Stent For Treatment of Coronary Revascularization</p>		<p>zation (TLR) at 12 months post-procedure. [Time Frame: 12 months]</p>			
<p>CREDIT-I (NCT01909869)</p>	<p>A PILOT First-In-Man Study to Evaluate Safety and Efficacy</p>	<p>Interventional Single Group Assignment</p>	<p>MACE</p>	<p>Yes</p>	<p>March 2018</p>	<p>Recruiting Last verified: July 2018</p>

	of the EXCEL- II With Cobalt Chromium Alloys Sirolimus Eluting Biodegradable Polymer Stent in the Treatment of Patients With de Novo Coronary Artery Lesions(CREDIT -I)					3
DISCOVERY1 23 (NCT0184484 3)	Evaluation With OFDI of Strut Coverage of	Interventional Single Group Assignment	OFDI assessed percent stent strut coverage [Time Fra	No	December 2014	Recruiting Last verified:

	Terumo New Drug Eluting Stent (Development Code TCD- 10023) With Biodegradable Polymer at 1, 2 and 3 Months		me: 3 months post procedure.]			April 201 3
OPTIMA (NCT0113701 9)	Optical Coherence Tomography Assessment of Intimal Tissue and Malapposition: A	Interventional. Allocation: Randomized	Rate of stent strut malapposition [Time Frame: 0 Days]	No	October 201 2	Unknown Last verified: May 201 0

	Random ized Compari son of the Biolimus A9- eluting and Everoli mus- eluting Coronar y Stents					
ORIENT (NCT0182655 2)	Compari son of the Angiogr aphic Result of the Orsiro Hybrid Stent With Resolut e Integrity Stent	Interventional. Allocation: Rand omized	Late Lumen Loss [Time Fra me: 9 months]	Ye s	Dec emb er 201 5	Not yet recr uitin g Last verifi ed: May 201 3

	(ORIENT)					
FIREHAWK (NCT01412164)	A Prospective Multicenter Single-Arm Observational Registry Study Assessing the Safety and Efficacy of FIREHAWK Biodegradable Polymer Target-release Rapamycin-	Observational Non-Randomized	Device related cardiovascular composite endpoint [Time Frame: 12 months]	Yes	February 2013	Not recruiting Last verified: April 2012

	<p>eluting Stent for the Treatme nt of Coronar y Artery Disease : TARGE T II</p>					
CREDIT -III	<p>A Prospec tive Multicen ter Single- Arm Observa tional Registry Study to Assess the Safety and Efficacy of EXCEL-</p>	<p>Observational [Patient Registry]</p>	<p>The Target Lesion Failure(TL F) as the primary endpoint at 12-month [Time Fra me: 12mon ths]</p>	<p>Ye s</p>	<p>Jun e 201 5</p>	<p>Recr uitin g Last verifi ed: Jan uary 201 4</p>

	<p>II With Sirolimu s Eluting Stent for the Treatme nt of Patients With de Novo Coronar y Artery ! (CREDI T-III Trial)</p>					
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401 www.clinicaltrials.gov; As accessed on 2/4/2014)

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408
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410

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412

413 All authors were equally involved in preparation of the manuscript. All authors read and approved the final manuscript.

414

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667

668 **ABBREVIATIONS**

669

670 PLLA, poly-L-lactic acid;

671 PDLLA, poly-D,L-lactide;

672 PLGA, polylactic-co-glycolic acid;

673 PLA, polylactide derivative.

674 BR, binary restenosis;

675 DD, non-polymeric dual DES;

676 FIM, first-in-man;

677 LLL, in-stent late lumen loss (mm);

678 MACE, major adverse cardiovascular events;

679 NP, non-polymeric DES;

680 ST, definite/probable stent thrombosis;

681 TLF, target lesion failure;

682 PP, permanent polymer;

683 BP, biodegradable polymer;

684 SD, standard dose;

685 NS, not significant.

686 PLA, poly-L-lactide;

687 PLC, 75:25 poly-L-lactide-co-caprolactone;

688 PLGA, 50:50 poly-D,L-lactide-co-glycolide;

689 PLLA, poly-L-lactic acid;

690 PVP, polyvinyl pyrrolidone;

691 USS, uncovered stent struts;

692 NS, not significant.

693 RCT, Randomized control trial.