The MGuard coronary stent: safety, efficacy, and clinical utility

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Abstract: Atheromatous and thrombotic embolization during percutaneous coronary revascularization is a feared complication that may cause impaired myocardial reperfusion even with a patent epicardial vessel. The MGuard stent is a cobalt chromium bare metal stent with a porous net attached to its outer surface that has been designed to prevent thrombus fragmentation and distal embolization during stent implantation. This review summarizes the available evidence supporting the use of the MGuard stent in different scenarios such as lesions with high thrombus burden, saphenous vein graft interventions, coronary perforations, or carotid lesions.

Keywords: Coronary artery disease, myocardial infarction, coronary stent, complication, thrombus, no-reflow phenomenon

Introduction
Atheromatous and thrombotic embolization during percutaneous coronary intervention (PCI) is a feared complication that may cause impaired myocardial reperfusion even with a patent epicardial vessel. Patients with acute coronary syndromes and those undergoing PCI of saphenous vein graphs (SVGs) are at highest risk of distal embolization, which frequently occurs after stent implantation due to thrombus fragmentation and protrusion by the stent struts. Inadequate myocardial reperfusion may occur in up to 50% of cases despite the use of optimal evidence-based coronary revascularization and is associated with a worse in-hospital and long-term prognosis.

Prevention of embolization rather than treatment should be the way forward as treatment is unlikely to succeed if no reflow is established. Thus, several pharmacological and technical strategies have been tested to minimize the distal embolization risk. Distal protection devices failed to show reductions in infarct size or mortality in patients undergoing primary PCI, although they seem to improve angiographic and clinical outcomes in PCI of degenerated SVG. Similarly, catheter-based thrombus aspiration has been associated with controversial results; although early single-center studies suggested a potential benefit, recent trials have shown no reductions in infarct size or mortality.

The concept of a stent covered with a membrane attached to its outer surface (to create a mechanical barrier to prevent thrombus fragmentation and filtration through the struts) has also been tested previously. However, covered stents, compared with bare metal stents, were associated with an increased risk of myocardial infarction, thus prohibiting their use in this scenario. Importantly, the barrier created by the membrane was related to delayed endothelialization of the stent and also to side branch occlusion, which could explain the increased rate of myocardial infarction.
The MGuard stent is a novel bare metal stent with a porous net attached to its outer surface that has been designed to trap thrombus debris reducing distal embolization, while pores allow normal endothelization of the stent through the membrane. The purpose of this review is to summarize all the relevant literature available about the use of this device.

The MGuard stent

The MGuard™ stent consists of a balloon-expandable close-cell design bare metal stent with a polyethylene terephthalate microfiber sleeve attached to its outer surface (Figure 1). The first-generation stent used a 316L stainless steel frame with strut thickness of 100 μm, while the new MGuard™ Prime platform is manufactured of L605 cobalt chromium alloy (strut thickness 80 μm). The Micronet™ is identical on both stents, with a fiber width of 20 μm and an expanded aperture size of 150x180 μm. Thus, the net creates custom-designed pores that are 5- to 40-fold smaller than the stent cells and appears to effectively act as a mechanical barrier decreasing or even preventing the thrombus protrusion and embolization to the distal vessel. MGuard™ and MGuard™ Prime diameters range from 2.5 mm to 4.0 mm, while lengths range from 11 to 39 and 13 to 38 mm, respectively.

Technical considerations

Guiding catheter, deliverability, and stent dislodgement

The MGuard stent is compatible with 0.014 in guide wires and 6 Fg guiding catheters. Crossing profile is slightly higher than newer-generation bare metal stents (1.1–1.3 mm for the stainless steel platform and 1.0–1.2 mm for the cobalt chromium platform), which together with the close-cell design may impair the deliverability. In a recent trial, the MGuard stent was unable to reach or cross the culprit lesion in nine of 217 cases (4.1%). Of note, all nine cases occurred with the stainless steel platform. Similarly, a recent single-center experience reported 1.8% of cases in which the stent could not reach the lesion.

From January 2011 to April 2014, the manufacturer received 54 complaints of stent dislodgement from the delivery system, most occurring with the MGuard Prime stent. By contrast, no cases of net detachment from the stent were seen. Stent dislodgement was reported in two of 217 cases (0.9%) of the prospective, randomized, multicenter evaluation of a polyethylene terephthalate micronet mesh–covered stent (MGuard) in ST-segment elevation myocardial infarction (MASTER) trial. Similarly, stent dislodgement occurred in six of 155 cases with the MGuard Prime stent (3.87%) at the IDE-MGuard for acute ST-elevation reperfusion II (MASTER II) trial, which led the manufacturer in 2014 to a voluntary field safety corrective action. This included the temporary suspension of the MASTER II trial and the addition of a new step during the manufacturing process so-called thermoforming in which the balloon-shaping process forms “shoulders,” which are intended to provide additional securement of the stent to the delivery system.

Side branch occlusion

Due to its double-layer design (metallic platform + mesh sleeve fibers), the MGuard stent may increase the risk of side branch (SB) occlusion. SB compromise may be related to two mechanisms: first, the summation of both layers (the metallic struts and the sleeve fibers) may act as a mechanical barrier blocking the coronary flow to the SB; second, the thrombus nearly located to the bifurcation may shift from the main vessel to the SB. It is possible that SBs compromised by the second mechanism could slightly improve with pharmacological treatment. Nevertheless, if the occluded SB is clinically relevant, a bifurcation technique may be considered (Figure 2). In our experience, wiring the SB should not be extremely complicated. By contrast, to advance a balloon across the net may require additional support measures such as the use of extension catheters.

MGuard stent in patients with ST-segment elevation myocardial infarction

After promising preliminary results in two small studies, efficacy and safety of the MGuard stent in patients undergoing primary PCI were first assessed by Piscione et al11 (Table 1). This was a multicenter Italian prospective registry that included 100 patients with ST-segment elevation myocardial infarction (STEMI). Thrombectomy was performed in 10% of the patients and predilation in 42% of cases. Complete myocardial reperfusion including myocardial blush grade =3 and ST-segment resolution at 60 minutes was achieved in 90%
The MGuard coronary stent

of patients. In hospital mortality occurred in seven patients (five presenting with cardiogenic shock preprocedure). Two subacute stent thromoses were also reported. Major adverse cardiovascular events (MACEs) up to 2 years were observed in 7.9% of the patients. Similarly, the MAGICAL study was a prospective multicenter international trial that enrolled 60 consecutive patients with STEMI. In this study, a high rate of complete epicardial and myocardial perfusion was also achieved (thrombolysis in myocardial infarction [TIMI] 3 flow in 93% of patients, myocardial blush grade 3 in 73%, ST-segment resolution in 68%). Cardiac mortality up to 3 years was very low (7%).

As opposed to previous studies in whom manual aspiration was performed in <20% of cases, our group first reported results of the MGuard stent in selected cases of failed manual aspiration. The study included 56 consecutive patients in whom a high thrombus burden persisted after aggressive manual aspiration. After MGuard stent deployment, >89% of cases had thrombus score =0, and >70% ST resolution was observed in 58.7% of patients. Only one stent thrombosis (acute) was reported, and 9-month MACE rate was very low (3.6%).

Importantly, a recent retrospective study compared results of consecutive procedures performed with either bare metal stents or MGuard stents. Due to differences in baseline characteristics between groups, a propensity matching was performed. After adjustment, patients treated with the MGuard stent presented with restenosis more frequently than those treated with bare metal stents. These results, however, have not been confirmed in other recent studies.

Two large-scale randomized trials have compared the MGuard stent to conventional PCI. The MASTER I trial was a prospective, multicentre, international, randomized study in which 433 subjects presenting with STEMI were randomized in a 1:1 fashion to the MGuard stent versus bare metal or drug-eluting stents. Device success was greater in the control arm (95.9% for MGuard vs 99.1% for control; P=0.003), but better reperfusion indices were obtained in the study group, as shown by the primary endpoint of the study, ST segment resolution (57.8% vs 44.7%; P=0.008), and incidence of TIMI 3 flow (91.7% vs 82.9%, P=0.006). Consequently, mortality at 30 days (0% vs 1.9%, P=0.06) and 1 year (1% vs 3.3%, P=0.09) tended to be lower in the MGuard group. Subgroup analyses showed the highest differences favoring the MGuard stent in cases with large thrombus burden or long delay time. However, incidence of repeat revascularization was higher than the control group (8.6% vs 0.9%, P=0.0003). The MASTER II trial was an international, multicenter, randomized trial designed to enrol 1,114 patients to show superiority of the MGuard stent vs

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**Figure 2** Coronary angiogram (30° cranial oblique projection) showing a totally occluded right coronary artery before intervention (A). Despite exhaustive thrombus aspiration. Red arrow points to the thrombotic total occlusion. (B), a high thrombus burden (TIMI thrombus score =4) persisted at the bifurcation (red arrows). After MGuard deployment (green line), TIMI 3 flow was restored to the posterolateral branch, although the posterior descending artery persisted totally occluded (C). Following "reverse T stenting," final TIMI flow was 3 in both the posterior descending artery and the posterolateral branch branches (D, final result).

**Abbreviation:** TIMI, thrombolysis in myocardial infarction.
The performance of the MGuard stent has been tested in two small multicenter prospective registries. The first-in-man study included 29 patients (17 with SVG intervention, mean SVG age =12.6 years). Device and procedural success was 100%. Final TIMI 3 flow was observed in all cases, and periprocedural myocardial infarction was observed in a single case. However, an extension of the study with 23 SVG interventions showed a high rate of MACE (23%), although no stent thrombosis or cardiac deaths were observed up to 20 months of follow-up. Similarly, the INSPIRE trial reported a high rate of procedural success of the MGuard stent in 16 SVGs, achieving final TIMI 3 flow in all cases. Nevertheless, in agreement with the previous study, a high conventional PCI in clinical events such as death or myocardial infarction. However, due to a higher than expected frequency of stent dislodgement, the study was voluntarily suspended in April 2014 after enrolment of 310 patients (155 patients per group). Analysis of the enrolled cohort showed no benefit of the MGuard stent in any of the analysed endpoints (ST-segment resolution was 56.9% vs 59.3% in the control group, P=0.68; mortality was 0.6% vs 1.9% in the control group, P=0.62). Similarly, a pooled analysis of the MASTER I and II trials (n=743) failed to show better myocardial reperfusion (ST-segment resolution 57.5% for the MGuard vs 50.7% for the control group, P=0.07), although mortality at 30 days was lower (0.3% vs 1.9%, P=0.03).

Thus, data from randomized trials suggest that the use of MGuard stent may reduce distal embolization and may improve survival at 30 days compared with other commercially available coronary stents, although further randomized studies are warranted to confirm this hypothesis. Also, further research is needed to understand if the restenosis rate is only related to the lack of drug elution or, by contrast, if there is also an antigenic stimulus caused by the polyethylene terephthalate mesh.

### MGuard stent in saphenous vein grafts

Compared with atherosclerotic plaques in native coronary arteries, SVG plaques are softer, longer, more friable, and contain necrotic debris and cholesterol crystals. Thus, percutaneous revascularization of SVG is associated with high rates of distal embolization, no-reflow phenomena, and periprocedural myocardial infarction. To avoid this complication, several drugs and devices have been tested. Graft stents have shown discouraging results and distal protection devices, although effective, are associated with an increase of procedural length and costs.

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### Table 1 Summary of studies with the MGuard stent in patients with STEMI

| Study | n | Device | Follow-up | TIMI flow achieved | Stent thrombosis | Mortality 30 days | TLR 30 days | Stent dislodgement | Deaths | Notes |
|-------|---|--------|-----------|-------------------|-----------------|------------------|-------------|-------------------|--------|-------|
| MASTER I | 122 | BMS/DES | 6 months | 57.9 ± 44.7 | 0.4% | 0.6% | 2% | 0.6% | 2% | Values expressed as %.
| MGuard | 60 | BMS | 9 months | 57.5 ± 50.7 | 0.9% | 0.9% | 1% | 0.9% | 1% | Includes 16 SVG patients, whom five events of in-hospital death occurred.
| MGuard II | 18 | BMS | 6 months | 56.9 ± 59.3 | 0.6% | 0.6% | 0% | 0.6% | 0% | Deaths reported in the control group.
| INSPIRE | 10 | BMS/DES | 12 months | 57.5 ± 68 ± 11.4 vs | 0 vs 0 | 1 vs 1 | 1 vs 1 | 1 vs 1 | 1 vs 1 | Includes 16 SVG patients, whom five events of in-hospital death occurred.

### Abbreviations

- BMS: bare metal stent
- DES: drug-eluting stent
- NA: not available
- NS: not significant
- STEMI: ST-segment elevation myocardial infarction
- TLR: target lesion revascularization
- TIMI: thrombolysis in myocardial infarction
- TV: target vessel
- SVG: saphenous vein graft
frequency of repeated revascularization of the target vessel was observed at 12 months of follow-up (19%).

**MGuard stent in other scenarios**

**Coronary perforations**

Graft stents seem to be an effective tool to seal severe perforations that have persisted despite prolonged balloon inflation. However, the high profile and low flexibility of previous covered stents compromise the deliverability of these devices, particularly in emergency situations. Moreover, the mechanical barrier created by the membrane may preclude the endothelization of the stent, which eventually may be associated with stent thrombosis or with restenosis at the stent edges. Our experience, although limited, suggests that the MGuard stents can also be successfully used to seal coronary perforations. The net, although pored, appears to effectively act as a mechanical barrier, decreasing or even preventing the leakage to the pericardial cavity. Nevertheless, additional mechanism might be involved, such as the stretching of the vessel or the compression of the disrupted tissue. Moreover, the cobalt platform would be more deliverable than the graft stent allowing also successful deployment in tortuous vessels, whereas the porous net would allow endothelization of the stent. Importantly, when MGuard stents are used in this bailout situation, high rates of restenosis may be expected. The mechanism, although unknown, might also be related to the inflammatory process involving the perforation.

**Coronary aneurysms**

Coronary artery aneurysms are a rare condition with a reported incidence of 0.14%–4.9% of patients undergoing coronary angiography and 0.3%–5.3% in patients after PCI. The majority of coronary aneurysms are atherosclerotic in origin and, in most cases, they are asymptomatic. However, potential complications associated with these abnormalities include thrombus formation, distal embolization, and, rarely, rupture.

Current available evidence for MGuard stent is limited to two case reports, in which the MGuard stent was successfully implanted, achieving complete exclusion of the aneurysm. However, the percutaneous exclusion of coronary aneurysm is not generally recommended, and therefore the use of the MGuard stent in this scenario should not be advised. Moreover, in our experience, although limited to two cases, MGuard stent implantation did not achieve total exclusion of the aneurysm in the early phase and was related to severe restenosis at late follow-up.

**Carotid arteries**

To avoid microembolization after stent implantation in carotid artery stenosis, a mesh-covered stent may be implanted. Notably, for carotid arteries, the manufacturer has mounted the same porous net over a nitinol self-expandable platform (so-called CGuard stent). Although the clinical experience is very limited, recent results of the CARENET trial have shown a very low rate of new ischemic ipsilateral lesions (as assessed by diffusion-weighted magnetic resonance imaging) as well as no major adverse cardiac and cerebrovascular events at 30 days. However, although these results are promising, more data are needed before making recommendations about the safety/efficacy of this device.

**Perspectives**

Considering the clinical benefits associated with drug-eluting stents in patients with STEMI and also due to the relatively high rate of restenosis of the MGuard stent, the manufacturer decided in 2014 to definitely cancel the MASTER II trial and to start development of the drug-eluting version of the MGuard stent. This new device will elute sirolimus from the stent platform (rather than the net) and will be available in late 2015.

**Conclusion**

The MGuard stent is a useful tool to avoid distal embolization during PCI of thrombotic lesions. The micronet attached to its outer surface prevents thrombus fragmentation and filtration during stent deployment. The enhanced myocardial reperfusion achieved with the use of the MGuard stent may be associated with a reduced mortality compared with other commercially available bare metal or drug-eluting stents. However, its use is associated with some limitations that must be highlighted: 1) the stent deliverability (even with the cobalt chromium version) may be slightly worse than other coronary stents that may be related to a higher crossing profile, and it must be taken into account when treating severely calcified or tortuous vessels; 2) due to its double-layer design (metallic platform + mesh sleeve fibers), the MGuard stent may compromise side branches, although bifurcation techniques can be performed to recover the compromised side branch; and 3) restenosis rate may be higher than with other commercially available coronary stents.

The MGuard stent can also be used in SVGs when conventional PCI with distal protection devices cannot be performed, as well as to seal coronary perforations that persist despite prolonged balloon inflation. In these cases, however, a high rate of repeated revascularization may be expected.
The new sirolimus-eluting version of the MGuard stent, which is currently under development, is invented to minimize the restenosis rate of the MGuard Prime stent and will probably be available in late 2015 in Europe.

Disclosure
The authors report no conflicts of interest in this work.

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