



Collateral network concept in 2023

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Extensive thoracoabdominal aortic aneurysm repair can cause spinal cord ischemia which significantly impacts survival and quality of life. Although this complication is uncommon, it is important to recognize the pathophysiology and preventative measures. In the 1990s, Dr. Griep and colleagues proposed the existence of an extensive collateral network that supports spinal cord perfusion, “the collateral network concept”. This includes an interconnecting complex of vessels in the intraspinal, paraspinal, and epidural spaces, and in the paravertebral muscles, involving the intercostal and lumbar segmental arteries as well as the subclavian and hypogastric (iliac) arteries. In this concept, as opposed to the one major segmental input model such as the Adamkiewicz artery, recognition of the importance of multiple inputs to the spinal circulation is paramount to maintaining the spinal blood flow and preventing spinal cord ischemia. In this article, we review the current evidence of the collateral concept and its application in aortic surgery.

Keywords: Collateral network; thoracoabdominal aorta; spinal cord injury



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Introduction

Spinal cord ischemia (SCI) in aortic surgery

SCI is rare (0.26% of all thoracic aneurysm repair) (1), but is the most serious complication after thoracic or thoracoabdominal aortic aneurysm (TAAA) repair, carrying a poor life expectancy regardless of type of procedure (one-year survival 63% in open repair, and 64% in endovascular repair) (2-5). In the original series of TAAA repair, Svensson *et al.* demonstrated the significant relation between the incidence and the extent of the aortic involvement using the Crawford classification of thoracoabdominal aneurysms and associated risk of paraplegia, where the more extensive type I (paraplegia or paraparesis, 15%) and II aneurysms (31%) that require removal of many of the sources of spinal cord blood supply are associated with the highest risk of perioperative neurological morbidity (2). Although the outcomes of TAAA repair have significantly improved over the past decades in experienced centers (paraplegia or paraparesis: Crawford I, 7.8%, and Crawford II, 13.9%) (4), the theory remains the same; the incidence of SCI increases

with the removal of more aortic segments and branches of the descending to abdominal aorta, so-called segmental arteries (SAs). Below the cervical level, the SAs originate from the descending aorta, and consist of the intercostal arteries at the thoracic level and the lumbar arteries at the lumbar level. Understanding the anatomy and physiology of the blood supply to the spinal cord is paramount to performing TAAA repair (6-8). Intraoperative SCI can develop for several reasons; ischemia from intraoperative clamping, incomplete restoration of blood flow to the spinal cord, and embolism (2-4,9,10). On the other hand, there is another mechanism of temporary or permanent paraplegia/paraparesis, known as delayed SCI (4,10). Numerous strategies of spinal cord protection have been discussed which include: operative technique (staged operation, open *vs.* endovascular, extent of repair, revascularization of subclavian artery or internal iliac artery, distal aortic perfusion, hypothermia), management of SAs [coil embolization as a staged procedure, intentional endoleak during endovascular approach, direct SA perfusion including prevention of steal phenomena during open repair,

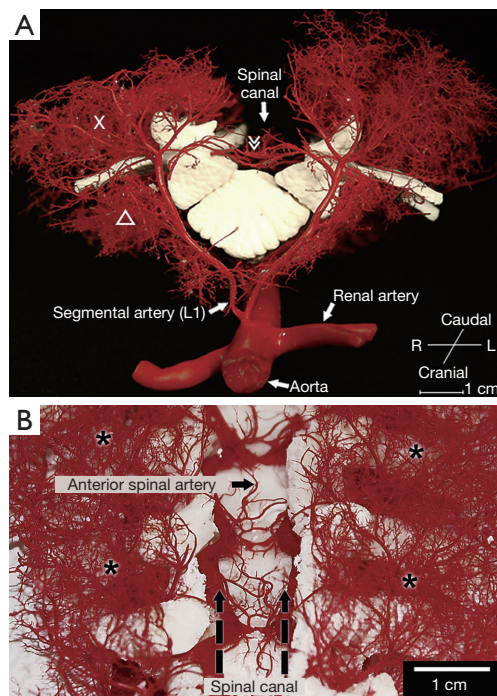


Figure 1 Anatomy of the collateral network, sagittal (A) and dorsal (B) views. Macroscopic appearance of the pair of dorsal segmental vessels at L1. The dorsal process is removed. In (A), X designates the paraspinous muscular vasculature providing extensive longitudinal arterioarteriolar connections in (A) and (B); the triangle indicates iliopsoas muscle; the double arrow indicates anterior spinal artery [reproduced with permission from Etz *et al.* (7); Copyright © 2011, Elsevier]. R, right; L, left.

reimplantation of intercostal arteries (ICAs)], hemodynamic status including collateral network concept [permissive hypertension, correction of anemia, cerebrospinal fluid (CSF) drainage, neuromonitoring], pharmacologic agents and so on (11-33). In this regard, cumulative laboratory studies and clinical observations suggest that a robust collateral network must exist to explain preservation of spinal cord perfusion when segmental vessels are interrupted (6-8,34,35). In this review, we will focus on the collateral network concept and its application.

Principles of collateral network concept

In 1881, Dr. Albert Adamkiewicz reported an artery originating from T8-L1, also known as the arteria radicularis magna or Adamkiewicz artery (AKA), as the principal vessel that feeds the lower thoracic, lumbar, and

sacral portions of the spinal cord (36,37). Based on this theory, some groups have been trying to identify AKA by imaging modality and reimplant this to prevent SCI during TAAA repair. In a multi-center analysis from Japan utilizing preoperative computed tomography (CT) angiography, AKA was found in 97.6% of patients who underwent TAAA repair predominantly on the left side of SAs (36). However, reimplantation of these identified SAs did not eliminate paraplegia including delayed SCI. Importantly, the concept of AKA does not explain this pathophysiology (6-8,10,38). Furthermore, reconstruction of SAs does not seem essential to prevent paraplegia given the lower rate of SCI which has been demonstrated in endovascular repair (17-22,39). Other studies showed that SCI is correlated more with the number of intercostal pairs sacrificed rather than the sacrifice of a single, important blood supply such as the AKA (37,38).

In the 1990s, Dr. Griep and colleagues proposed instead the existence of an extensive collateral network that supports spinal cord perfusion (*Figure 1*) (6-8,35,38,39). This collateral network includes an interconnecting complex of vessels in the intraspinal, paraspinous, and epidural spaces, and in the paravertebral muscles, including the intercostal and lumbar SAs as well as the subclavian and hypogastric arteries. In this concept, as opposed to the one major segmental input model (AKA), recognition of the importance of multiple inputs to the spinal circulation is paramount to preventing SCI. In a pig model, Etz *et al.* showed only a 50% paraplegia rate even after sacrificing all arteries which may support the additional blood supply to the spinal cord other than SAs (8). Another important result from Dr. Griep's group was that the incidence of SCI is correlated more with the number of intercostal pairs sacrificed rather than their location; more than 10 pairs or 8-12 pairs sacrificed beginning in the lower thorax (Th6 or below) showed a higher risk of SCI (37,40,41).

The maintenance of adequate spinal cord perfusion is crucial to the success of open and endovascular repair of extensive TAAA to prevent critical SCI when blood flow to the SAs is interrupted. Spinal cord perfusion pressure (SCPP) represents the net pressure gradient that drives oxygen delivery and blood supply to the cord, and is calculated as the difference between the collateral network pressure (CNP) and the pressure within the intrathecal space [CSF pressure $\approx$ central venous pressure (CVP)] (6-8,13,42).

$$\text{SCPP (mmHg)} = \text{CNP (mmHg)} - \text{CSF pressure} (\approx \text{CVP}) \quad [1]$$

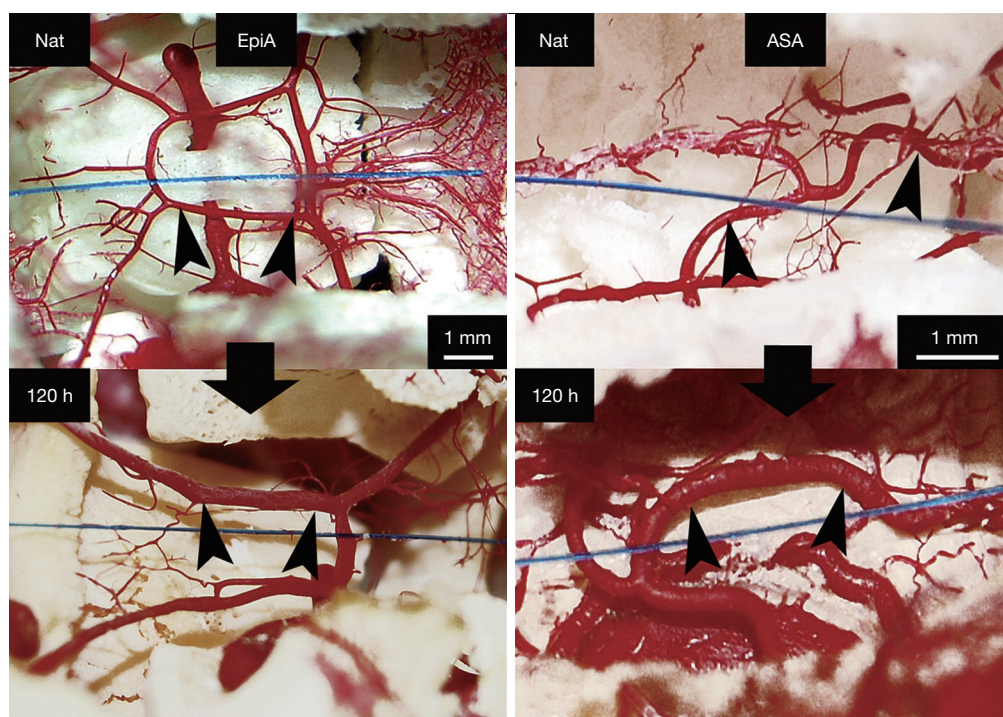


Figure 2 Increase in diameters of epidural arcades (left, black arrowheads) and ASA (right, black arrowheads) in the native pig compared with a pig perfused 5 days after extensive SA occlusion (120 hours). Soft tissue was removed completely from the samples, but the bony vertebral column was preserved. Note the specific enlargement of the longitudinally oriented vessels of the epidural arcade on the left. The blue line is a segment of prolene suture to impart an additional sense of scale [reproduced with permission from Etz *et al.* (35); Copyright © 2011, Elsevier]. Nat, Native; EpiA, epidural arcade; ASA, anterior spinal artery.

Clinically the mean arterial pressure (MAP) is a landmark of SCPP although MAP and SCPP are not equal. In an experimental model using a pig, Etz and colleagues demonstrated that CNP is 60–80% of MAP at baseline (39). When all SAs were taken in 2 stages a week apart, CNP fell only to 50% to 70% of baseline, and SCI was rare. In human beings, baseline CNP was 75% of mean aortic pressure, fell in proportion to the number of SAs ligated, and began recovery within 24 hours (39). Interestingly, they found that CNP was lower with non-pulsatile distal bypass than with pulsatile perfusion. Other experimental models by Geisbüsch *et al.* and Etz *et al.* showed that after experimental SA ligation, an early increase in collateral network flow of existing native vessels was observed for up to 48 hours, likely due to vasodilatation, followed by a definite increase in the number, density, and size of small arteries (Figure 2) (20,35). Furthermore, remodeling of paraspinal microvessels (arterioles up to 40 μ m) was observed, not just in the density or the number of microvessels, but also in the

spatial orientation of these vessels 5 days after completing SA occlusion (T4–L5) (20,35,40). These results suggest that the collateral network concept includes not only robust blood circulation to the spinal cord but also a dynamic remodeling process of the spinal cord perfusion where collateral supply is received from both from the cranial (the subclavian artery) and caudal (hypogastric arteries/internal iliac arteries) pathways to the area of the spinal cord that has been deprived of direct SA perfusion (Figure 3) (40).

Clinical application of the collateral network concept

Staged approach and revascularization of the branches

In line with the dynamic remodeling process of increased spinal cord circulation after sacrifice of SAs, a staged repair strategy for extensive pathologies has emerged. Staged repair can be done by either open or endovascular approaches. Hybrid staged repair (open + endovascular) yields a lower incidence and less severity of SCI compared

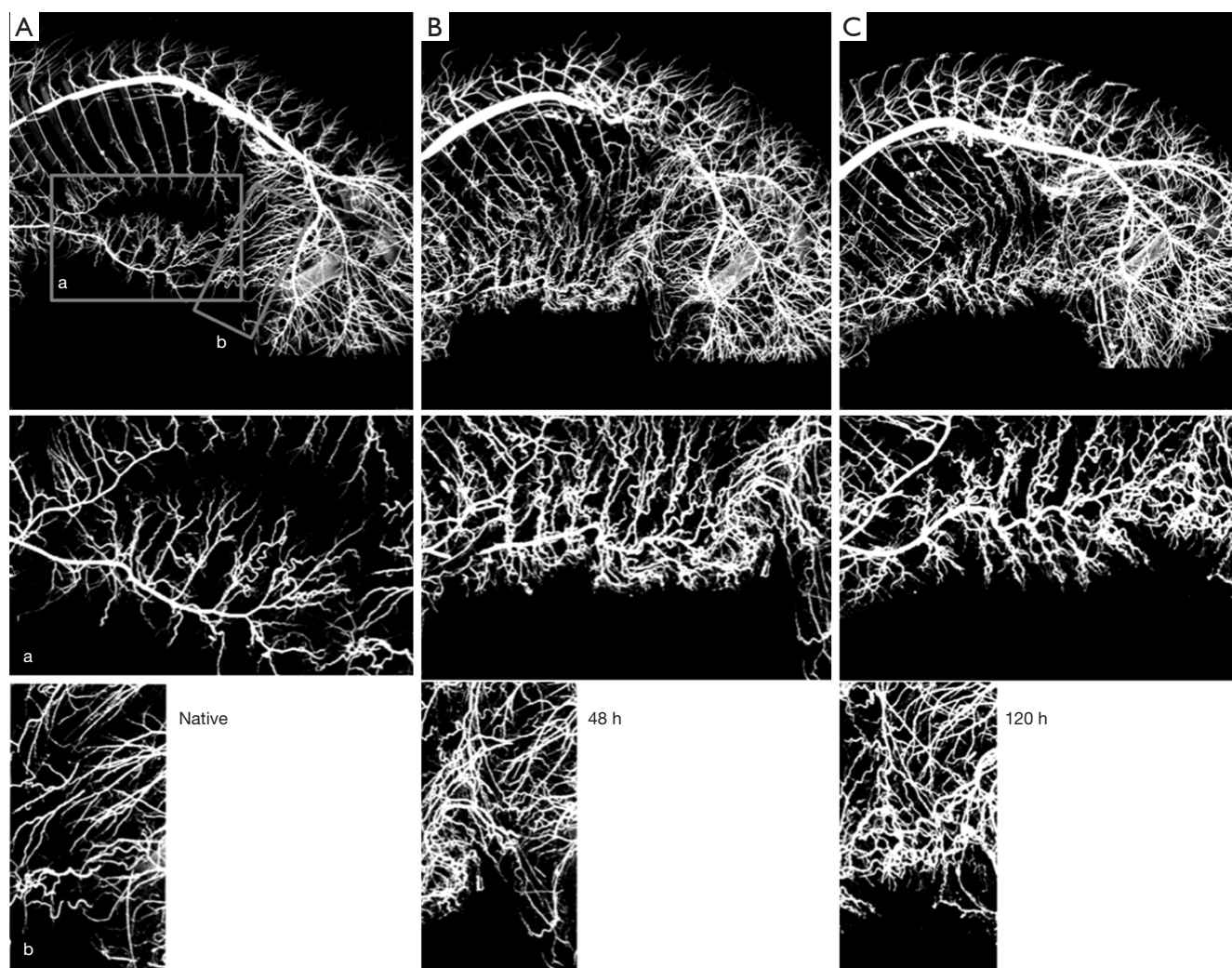


Figure 3 Computed tomographic image of the arterial vessel system of the entire pig. Bones, intestines, and other tissues were extracted from the image. (A-C) Show examples of vessel casting for a native pig at 48 and 120 hours after segmental artery ligation, respectively. A frame around the internal thoracic artery (a) and around the iliolumbar vessel (b) was cropped, and vessel density was compared between the groups. An impressive increase in density of small vessels could be seen, but no statistically significant differences could be demonstrated with the small number of pigs examined at each time point [reproduced with permission from Geisbüsch *et al.* (40); Copyright © 2012, Elsevier].

with single-stage repair (11,13). In patients with extensive TAAA involving proximal descending or arch pathology, open stage I arch repair, classic or frozen elephant trunk, can be used to facilitate stage II procedure by either open or endovascular repair including hybrid approach, to reconstruct visceral branches (14,43-48). Although hybrid or total endovascular arch repair is available, open arch repair remains a gold standard for aortic arch with extensive

distal aortic pathology to provide a reliable proximal repair in this setting.

In Crawford type II TAAA repair, several staged approaches have been reported (12,13,27,28,43-48). Over the past decade, unique applications of the collateral network concept have emerged as staged approaches utilizing endovascular techniques (*Figure 4*). Minimally invasive segmental artery coil embolization (MISACE)

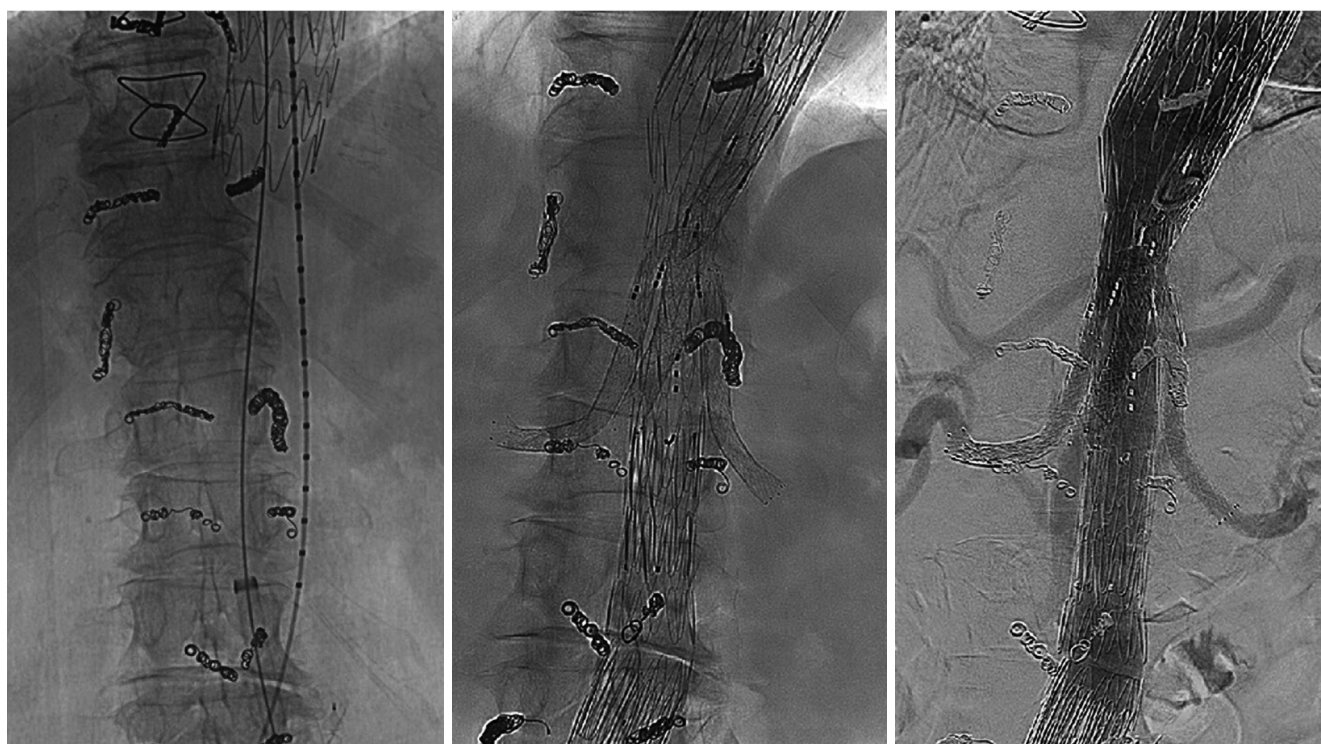


Figure 4 MISACE at the thoracoabdominal level followed by endovascular exclusion of the thoracoabdominal aneurysm and completion angiography. First clinical series of MISACE prior to total endovascular repair of TAAA [reproduced with permission from Branzan *et al.* (19); Copyright © 2018, CCC Marketplace]. MISACE, minimally invasive segmental artery coil embolization; TAAA, thoracoabdominal aortic aneurysm.

prior to endovascular TAAA repair is a novel approach to reduce SCI (19-22); SA occlusion is performed using endovascular coiling to trigger arteriogenic preconditioning of the collateral network, thereby allowing for recruitment of otherwise redundant arterial collaterals to the spinal cord (19-22). A University of Leipzig group reported their initial experience and its efficacy (19). Utilization of MISACE is now under investigation as a randomized study, Paraplegia Prevention in Aortic Aneurysm Repair by Thoracoabdominal Staging (PAPA-ARTiS) trial (ClinicalTrials.gov Identifier: NCT03434314). Patient enrollment was closed in June 2022.

Temporary aneurysm sac perfusion as intentional non-completion of at least one stentgraft branch is expected to prevent complete aneurysm sac thrombosis and maintain spinal cord perfusion through patent intercostal and/or lumbar arteries (16,18,24,25,43). The delayed bridging stent method (temporary aneurysm sac perfusion) was

reported by Kaszpac *et al.* in 2014 (Figure 5) (24). In this technique, a branched stentgraft is placed in a preliminary intervention, whereby a target vessel, usually the celiac artery, is not connected in order to maintain a temporary endoleak for the purposes of spinal cord perfusion. In the follow-up procedure, simultaneously to the sac perfusion branches, the procedure is completed by connecting the side branch to the target vessel. A similar concept was originally reported by Lioupis and colleagues in 2011 (25), known as the perfusion branch technique, where a branched custom-made endograft is made with a separate perfusion branch in the body of stentgraft to maintain a temporary “intentional” endoleak in order to perfuse the aneurysm and SAs followed by stage II endovascular completion. These novel methods are examples of clinical applications of the collateral network concept, and have demonstrated favorable SCI rates compared with conventional endovascular approaches covering the long aortic segment.

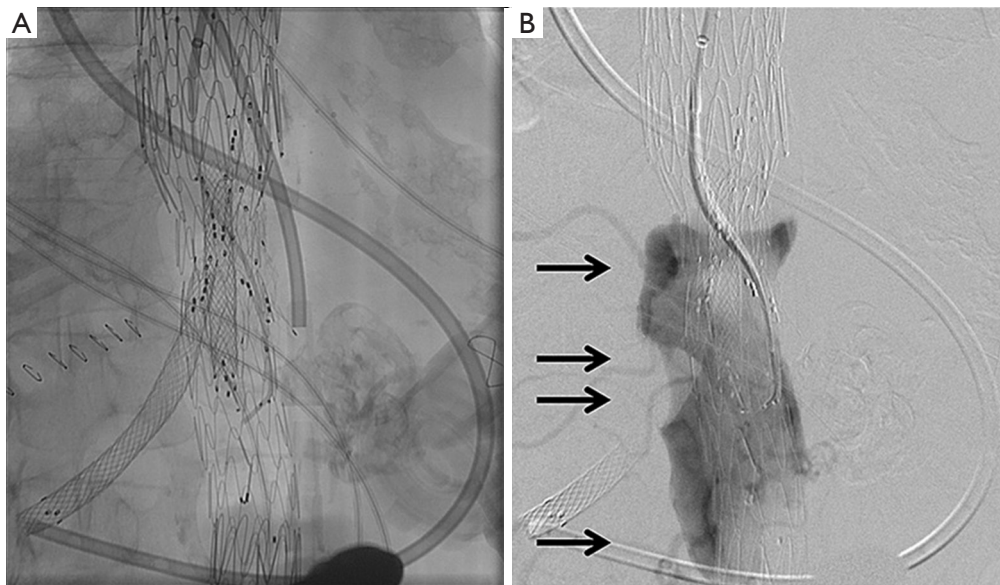


Figure 5 Endovascular thoracoabdominal aneurysm repair using temporary aneurysm sac perfusion technique. (A) Intraoperative angiogram showing implantation of branched endograft with connecting stent grafts to the celiac trunk and the superior mesenteric artery. (B) Temporary aneurysm sac perfusion via the right renal artery with flow to the aneurysm and at least four lumbar arteries (black arrows) [reproduced with permission from Kasprzak *et al.* (24); Copyright © 2014, Elsevier].

According to the collateral network concept, the left subclavian artery (LSCA) is an important collateral source via the internal mammary artery and other branches connected to the SAs (6-8). A meta-analysis by Huang and colleagues reported that revascularization of the LSCA is associated with a lower incidence of SCI after extensive coverage of SAs (44). This favorable outcome in the setting of Zone 1 or 2 thoracic endovascular aortic repair (TEVAR) with LSCA revascularization may suggest that revascularization of *in situ* SAs such as the anterior spinal artery is not necessary for SCI prevention. There are several options for revascularization of LSCA prior to TAAA repair: debranching the LSCA during open arch repair, left carotid artery to LSCA bypass, right axillary artery to left axillary artery bypass, or TEVAR (one-branch, fenestration, chimney, scalloped graft, etc.) (29,43-46). Recent data from the Society for Vascular Surgery Vascular Quality Initiative showed comparable outcomes between open and endovascular LSCA revascularization (29). Currently, LSCA revascularization is recommended for Zone 2 arch repair when patients have the following risk factors for SCI (30): (I) prior infrarenal aortic repair with occlusion of lumbar and middle sacral arteries; (II) planned long-segment (20 cm) coverage of the descending thoracic aorta where

critical intercostal arteries originate; (III) hypogastric artery occlusion, and (IV) presence of early aneurysmal changes that may require subsequent therapy involving the distal thoracic aorta. A history of either repaired or unrepaired abdominal aortic aneurysm (AAA) is known as a risk of SCI during TAAA repair as lumbar or iliac arteries are occluded or narrowed in these patients (the number of patent SAs: 4.5 ± 3.0 pairs in patients with AAA *vs.* 7.6 ± 1.3 pairs in patients without AAA) (31).

The collateral network concept can also be applied to the extent of repair as surgical resection of life-threatening segments only may be an important strategy given the impact of paraplegia. According to the largest experience of open TAAA repairs by Dr. Coselli's group, the incidence of SCI was lower in extent I or IV repair compared to extent II or III repair (4). For chronic aortic dissection with aneurysm cases, we previously reported that the growth rate of distal aorta is favorable after resection of most problematic segments (47). This concept is crucial especially in elective cases because most aneurysm surgeries are prophylactic (48).

Intraoperative blood flow to the SAs during TAAA repair
Steal phenomenon of the aorta or SAs has been recognized

as a potential risk factor for SCI during TAAA repair (9,13). Shiiya and colleagues reported based on clinical experiences that aortic steal phenomenon was a major cause of intraoperative SCI, and some cases were reversible by clamping the SAs after recognizing neuromonitoring changes (9). Their findings support the importance of both the steal phenomenon and collateral network in demonstrating reversibility of intraoperative SCI by improving SCPP through clamping the SAs. Similarly, distal aortic perfusion while cross-clamping the aorta is another method to maintain SCPP (perfusion to the SAs) assisting spinal cord protection. A University of Texas group reported their experience of TAAA repair using the adjunct of distal aortic perfusion (increasing CNP) and CSF drainage (lowering CSF pressure), providing a low incidence of SCI (15). Recently, Haunschild and colleagues showed the significance of steal phenomenon even during distal aortic perfusion (49). In this report, distal aortic perfusion with blood flow steal from the SAs resulted in decrease in perfusion in the spinal cord regional flow, especially in the lower thoracic and upper lumbar levels (Th8-L2). Interestingly, this decrease of perfusion in the spinal cord increased when clamping the SAs. These results suggest that Th8-L2 are most sensitive to “segmental steal”, and segmental steal during TAAA repair using distal aortic perfusion also plays an important role in causing SCI.

Re-implantation of SAs during open TAAA repair has been discussed as a method of prevention of SCI. Dr. Coselli's group reported their approach using left heart bypass with mild hypothermia (32–34 °C, nasopharyngeal) where one or more pairs of intercostal and lumbar arteries were reattached, especially between T8 and L1, when possible (4). Some centers look for AKA as a preoperative evaluation of TAAA repair with a high rate of reimplantation rate (36). Dr. Estrera and Safi's group reported the efficacy of neuromonitoring-guided reimplantation of SAs where patent important SAs (i.e., T8-T12) or adjacent patent SAs (T4-T7, L1, L2) were reimplanted if changes on somatosensory evoked potentials (SSEPs) and motor evoked potentials (MEPs) were encountered (43). A Japanese multi-center study looking at identification of AKA demonstrated that a risk of not reconstructing AKA segments was not significant in open repair, and whether AKA covered or not was not associated with SCI in endovascular approach (36). Furthermore, Dr. Griep's group reported a favorable SCI rate without any reimplantation of SAs based on the

collateral network concept. Of note, they demonstrated that the number of SAs cut and location of SAs (number of sacrifice >8 with highest T6 or below) were associated with SCI (13,37,38,41).

Regardless of re-implanting SAs or not, the majority of aortic programs use some sort of approach based on the collateral network concept to prevent SCI such as increasing the MAP including distal aortic pressure, decreasing cerebrospinal fluid pressure, increasing hemoglobin levels by transfusion, or completion of repair to establish pulsatile flow to entire “collateral network” as soon as possible (4,15,36,37,42,43).

Importance of SCPP

Maintaining adequate SCPP is the key component of the collateral network concept to secure blood flow to the spinal cord. It is paramount to measure both upper body and lower body arterial pressure during TAAA repair. The MAP needs to be maintained on the high side (>90 mmHg) especially during sacrifice of the SAs and after completion of repair (2,6–8,39,47,49). This can be also achieved by increasing cardiopulmonary bypass flow through the femoral artery to increase a distal aortic perfusion pressure to the lower SAs (49). Care should be taken to maintain pulsatile flow of the upper body during partial cardiopulmonary bypass of the lower body as pulsatile flow provides more blood flow to the spinal cord, as demonstrated by Etz *et al.* (39).

Neuromonitoring such as MEP and/or SSEP is frequently used to detect early SCI during the procedure (4,23,37,42). Once SCI is suspected on MEPs or SSEPs after clamping SAs, the target MAP can be increased and measurement of MEP is repeated in 1 to 5 minutes to observe recovery of evoked potentials (13,14,43,47,49). Dr. Etz's group from Leipzig introduced the efficacy of collateral network near-infrared spectroscopy (NIRS) (17,27) as non-invasive, real-time monitoring of spinal cord oxygenation which can be used during surgical repair and postoperatively. In a clinical series, lumbar collateral network NIRS oxygenation levels dropped significantly after proximal aortic cross-clamping but fully recovered after restoration of pulsatile flow to the baseline (17). They found that lumbar collateral network NIRS reacts to occlusion of SAs in real time and correlates with permanent neurologic deficit in animal models (27). Although further evidence is warranted, this approach has a potential as an adjunct to adjust intraoperative management such as

increasing MAP or distal aortic perfusion pressure during anastomosis to increase SCPP.

Kawanishi *et al.* from Kobe University demonstrated that the duration of intraoperative hypotension after separation from cardiopulmonary bypass is an independent risk factor for SCI in patients with TAAA which supports the importance of maintaining SCPP (50). Etz *et al.* nicely demonstrated that SCPP is lower at completion of cardiopulmonary bypass or repair compared to baseline SCPP level (approximately one-third of the baseline) which returns to normal level by 120 hours postoperatively (39).

CSF drainage is another important adjunct to increase SCPP by reducing the CSF pressure/CVP. In a single-center randomized clinical study, Coselli and colleagues showed the efficacy of CSF drainage during open TAAA repair to reduce SCI (33). CSF drainage can also be used during endovascular repair for patients with risk factors for SCI such as long segmental coverage, occlusion of SAs or aortic branches, etc. (23,43–47). Dr. Estrera's group reported the importance of protocol-based management of CSF drain and hemodynamics, known as CSF drain status/oxygen delivery/patient status (COPS) protocol (51). In this protocol, the goal parameters are: CSF drain pressure <5–10 mmHg, correcting oxygen saturation, goal hemoglobin >12 g/dL, cardiac index >2.5 L/min/m², and MAP >90 mmHg with SCPP >80 mmHg.

Hypothermia

Mild hypothermia is frequently used as an adjunct of spinal cord protection during open repair, regardless of femoro-femoral bypass or left heart bypass (4,9,13,15). In an experimental model, Strauch and colleagues demonstrated the protective effect of mild hypothermia (32 °C) over normothermia (36.5 °C) which allows for 50 minutes of safe ischemic time of the spinal cord when cross-clamping the aorta (34). Deep hypothermia can be utilized in extensive TAAA repair expecting prolonged SCI during the procedure. Although there may be concern of complications related to deep hypothermia such as longer cardiopulmonary bypass time, bleeding, respiratory failure, renal failure, or infection, there are numerous benefits such as reliable organ protection including the spinal cord, no steal phenomenon, bloodless surgical field, and less complicated anastomosis without any clamping (4,36,43,52,53). Previous studies comparing open TAAA repair using deep hypothermic circulatory arrest versus beating hearts with warmer

temperature (mild to moderate hypothermia) reported lower incidence of SCI utilizing deep hypothermic circulatory over mild to moderate hypothermia (52,53). In terms of ischemic tolerance of the spinal cord during deep hypothermia, Etz and colleagues studied the safe limit of lower body ischemic time using selective antegrade cerebral perfusion model during aortic arch repair, showing that 120 minutes of deep to low-moderate hypothermia (20–23 °C) falls into the safe lower body ischemic time (54). Although their model was designed to assess the relationship between temperature and spinal cord protection during aortic arch surgery using antegrade cerebral perfusion, this result may provide significant insights into TAAA repair in terms of temperature management.

Delayed SCI

SCI can present in a delayed fashion with progressive lower extremity weakness (2,10,38). In the largest experience of 3,309 TAAA repairs from Dr. Coselli's group, the proportion of delayed SCI (permanent paraplegia + paraparesis) was greater than immediate SCIs (delayed SCI, 55.6%, N=99/178 *vs.* immediate SCI, 44.4%, N=79/178) (4). They also reported a wide range of time frames regarding delayed SCI where paraplegia and paraparesis occurred 13 hours to 91 days postoperatively (10). In Dr. Griep's group's experience, delayed SCI accounted for 85% of SCI after TAAA repairs where the breakdown of delayed SCI was further divided to intermediate-delayed (up to 48 hours postoperatively) and late-delayed SCI (>48 hours postoperatively) with almost equal distribution (38). A similar trend was observed in studies by Conrad *et al.* and Maeda *et al.* comparing open versus endovascular TAAA repair, where a greater percentage of patients showed delayed SCI in patients who received endovascular repair (55,56).

There are several potential mechanisms associated with delayed SCI, including unstable hemodynamics resulting in low SCPP, spinal cord edema, inflammatory response including ischemic-reperfusion injury, or occlusion of re-implanted SAs (2,9,10,38,57). Occurrence of delayed SCI itself strongly suggests the presence of the collateral network concept because acute occlusion or severe stenosis of one of the SAs or embolism is less likely as a cause of SCI in the postoperative period as a delayed onset. This late presentation of SCI also shows the importance of maintaining SCPP not only in the operative room but also

in the intensive care unit to enhance collateral network circulation and its remodeling. Etz and colleagues analyzed delayed SCI patients and found that CVP was significantly higher in the paraplegic patients from 1 to 5 hours postoperatively compared to patients whose motor function recovered (38). Their result supports the importance of lower CSF pressure or CVP in the early postoperative period by utilizing CSF drainage to maintain sufficient SCPP. Although the duration of spinal drainage after TAAA repair is usually 24–48 hours after surgery (57), it needs to be determined based on individual SCI risk including the length of aorta replaced, the number of sacrificed SAs, patency of SAs, history of aortic surgery, or other factors as discussed above (4,28–30,37,38,57). When delayed SCI happens, several treatments are considered: vasopressors, systemic corticosteroids, naloxone, and osmotic agents (e.g., mannitol, glycerol), blood transfusion, and CSF drainage (10,38,57). In the largest experiences from the Texas Heart Institute, outcomes of delayed SCI were not significantly different compared to immediate SCI where both groups showed poor survival in the long-term (five-year survival was 31% in both groups) (10). According to the COPS protocol proposed by the University of Texas group, a CSF drain is maintained for 3 days given the onset of delayed SCI based on their experience (51). Cheung *et al.* reported increased chances of recovery of delayed SCI (5 out of 8 full recovery, and 3 had partial recovery) using the COPS protocol (57). As some delayed SCIs are reversible by appropriate treatment, frequent neurological evaluation and maintenance of SCPP is crucial after TAAA repair.

On the other hand, the role of routine insertion of CSF drain during TEVAR is unclear in terms of benefits and risks related to CSF drain insertion (intra cranial hypotension, intracranial hemorrhage, headache, local bleeding, and SCI related to spinal hematoma). A Duke University's group reported the efficacy of a restrictive lumbar CSF drain algorithm in the setting of TEVAR for TAAA repair, where permissive hypertension and collateral revascularization showed 0% incidence of SCI (58). Of note, their algorithm is based on the collateral network concept.

Pharmacological effect on collateral network

Many pharmacologic options have been reported which include anti-inflammatories, antioxidants, antiapoptotic agents, drugs that reduce neuronal excitotoxicity, drugs that induce or mimic ischemic preconditioning, medications

that reduce metabolic demand, and osmotic agents to reduce tissue swelling (59), but discussing all of these mechanisms would be beyond the purpose of our review. In terms of collateral circulation, systemic vasopressors (e.g., norepinephrine, epinephrine, vasopressin, or phenylephrine) are a common approach to increase the MAP although this causes vasoconstriction of the arteries, so it is important to maintain the balance between suitable MAP and appropriate use of vasopressors in the perioperative period.

Although clinical application of these drugs is not yet common, Svensson *et al.* reported the efficacy of topical approaches using intrathecal papaverine as a local vasodilator which results in lower SCI rate in patients with TAAA repair (SCI rate of intrathecal papaverine group, 3.6% *vs.* control, 7.5%) (60). The idea of this technique is to increase blood flow to the spinal cord according to the collateral network concept.

Conclusions

The collateral network concept includes not only the robust, anatomical blood supply to the spinal cord but also a dynamic, temporal and spatial remodeling process of the spinal circulation, and was suggested by Dr. Griep and colleagues in 1990s based on both basic experiments and clinical experiences. This concept is now widely accepted when treating TAAA regardless of open endovascular or hybrid repair to prevent perioperative SCI.

Based on the current evidence available for the collateral network concept, the following recommendations for the management of spinal cord protection are proposed (*Table 1*): staged TAAA repair regardless of open or endovascular repair, if possible, resection of life-threatening segments only to minimize the length of repair, revascularization of the LSCA if it needs to be covered by stent graft during proximal repair, neuromonitoring during TAAA repair, CSF drainage if open repair or endovascular repair for long segment coverage, CSF pressure <12 mmHg for 48 hours after operation, maintaining high MAP (>90 mmHg) perioperatively, selective reconstruction of SAs with extensive aortic resection, avoidance of steal phenomenon of SAs, distal aortic perfusion during cross-clamping if indicated, mild or deep hypothermia depending on the extent of open repair, securing hemostasis to avoid unstable hemodynamics, and maintaining hemoglobin level >10–12 g/dL to maintain stable hemodynamics and oxygen delivery.

Table 1 Consideration of spinal cord protection in thoracoabdominal aneurysm repair based on the collateral network concept

Management	Considerations
Preoperative management	
Surgical strategy	Staged procedure if possible for both open and endovascular repair
Length of repair	Resection of life-threatening segments only
Revascularization of the left subclavian artery	Yes
Identification of the artery of Adamkiewicz by imaging study	Not necessary
Intraoperative management	
Neuro monitoring	MEP or SSEP
Spinal drainage	Yes if patients have risks of SCI, open repair, or covering long segment TEVAR (>15–20 cm)
MAP	>90 mmHg
Cerebrospinal fluid pressure	<12 mmHg
Segmental arteries	Ligation before opening the aneurysm with neuromonitoring Any change during clamping large SAs needs repeat MEP with increased MAP Re-implantation of SAs is controversial but can be done with neuro monitoring guidance or intraoperative findings
Perfusion strategy (open repair)	Distal perfusion after distal anastomosis Selective perfusion of SAs is not necessary Mild hypothermia (32 °C) using partial bypass or deep hypothermic circulatory arrest
Postoperative management	
Spinal drainage	Keep for 24–48 hours Drainage does not exceed 15–20 mL/hour to maintain CSF pressure or CVP <12 mmHg
Mean arterial pressure	>90–100 mmHg for first 7 days
Neurological assessment	–
Avoidance of anemia	Hemoglobin >10–12 g/dL

MEP, motor evoked potential; SSEP, somatosensory evoked potential; SCI, spinal cord ischemia; TEVAR, thoracic endovascular aortic repair; SA, segmental artery; MAP, mean arterial pressure; CSF, cerebrospinal fluid; CVP, central venous pressure.

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Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

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