This Summary of Safety and Clinical Performance (SSCP) is intended to provide public access to an updated summary of the main aspects of the safety and clinical performance of the device.

The SSCP is not intended to replace the Instructions for Use as the main document to ensure the safe use of the device, nor is it intended to provide diagnostic or therapeutic suggestions to intended users or patients.

The following information is intended for users/healthcare professionals.

NuMED

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1. Device identification	and general information
Device trade name(s)	NuMED RVOT/CoA Stent Family Covered CP Stent Covered Mounted CP Stent G-Armor Covered Stent G-Armor Covered Mounted Stent
Model Number	NuMED RVOT/CoA Stent Family – Model 1650         Covered CP Stent – Model 427.1         Covered Mounted CP Stent – Model 428.1         G-Armor Covered Stent – Model 432         G-Armor Covered Mounted Stent – Model 434
Manufacturer's name and address	NuMED, Inc. 2880 Main Street Hopkinton, NY 12965 USA
Manufacturer's single registration number (SRN)	US-MF-000010948
Basic UDI-DI	08877141650TH
Medical device nomenclature description / text	EMDN – P070401020199 - PTFE VASCULAR ENDOPROSTHESES, STRAIGHT - OTHER
Class of device	ш
Year when first certificate (CE) was issued	2004 (Covered CP Stent) 2009 (Covered Mounted CP Stent) G-Armor Devices – Not yet CE Marked
Authorised Representative (AR)	G. van Wageningen B.V. Hallenweg 40, 5683 CT Best, The Netherlands
AR SRN	NL-AR-000010437
Notified Body	SGS Belgium NV
Notified Body ID number	1639

2. Intended use of the device				
Indications for use	<b>INTENDED USE</b> Permanent implants to treat Coarctation of the Aorta, and/or RVOT disruptions.			



	INDICATIONS
	<ul> <li><u>Coarctation of the Aorta (CoA)</u></li> <li>Indicated for implantation in the native and/or recurrent coarctation of the aorta on patients with the following clinical conditions:</li> <li>Stenosis of the aorta resulting in significant anatomic narrowing as determined by angiography or</li> </ul>
	<ul> <li>non-invasive imaging, i.e. echocardiography, magnetic resonance imaging (MRI), CT scan;</li> <li>Stenosis of the aorta resulting in hemodynamic alterations, resulting in systolic pressure gradient, systemic hypertension or altered left ventricular function;</li> <li>Stenosis of the aorta where balloon angioplasty is ineffective or contraindicated;</li> <li>Stenosis diameter &lt;20% of adjacent vessel diameter. Stenosis that would present increased risk of vascular damage or disruption; or aneurysm associated with coarctation of the aorta.</li> </ul>
	<u>Right Ventricular Outflow Tract (RVOT)</u> Indicated for treatment of right ventricle to pulmonary artery (right ventricular outflow tract) conduit disruptions that are identified during conduit pre-dilatation procedures performed in preparation for transcatheter pulmonary valve replacement.
Contraindications and/or limitations	<ul> <li>Contraindications include:</li> <li>Patients too small to allow safe delivery of the stent without compromise to the systemic artery used for delivery;</li> <li>Unfavorable aortic anatomy that does not dilate with high pressure balloon angioplasty (CoA only);</li> <li>Occlusion or obstruction of systemic artery precluding delivery of the stent (CoA only);</li> <li>Clinical or biological signs of infection;</li> <li>Active endocarditis;</li> <li>Known allergy to aspirin, other antiplatelet agents, or heparin (CoA only);</li> <li>Pregnancy.</li> </ul>

3. Device description	
	The Stents are balloon expandable and intended for permanent implant. The Stents are composed of heat treated 90% platinum/10% iridium wire that is arranged in laser welded rows with a "zig" pattern. The number of zigs in a row can be varied and will impact the strength of the stent as well as the eventual expanded diameter and percent stent shortening, while the number of rows will determine the unexpanded length of the stent. The Stents have an ePTFE covering attached to the stent framework. This covering acts as a fluid barrier creating a fluid tight conduit through the stent length.
Description of the device	The BIB Stent Placement Catheter is triaxial in construction with two lumens being used to inflate the balloon while one lumen is being used for tracking over a guidewire. The inner balloon is $\frac{1}{2}$ of the outer balloon diameter and 1 cm shorter. The purpose of the double balloon catheter is to apply an incremental inflation for the purpose of dilating a stent. The inner balloon provides initial expansion of the stent and also acts as a tool to hold the stent on the catheter prior to the outer balloon being inflated. The outer balloon is then inflated, providing the remainder of the expansion. There are radiopaque platinum marker bands under the balloon shoulders, to aid during placement. The balloons are designed to inflate to the diameter and length listed on the label at a specific pressure. Thus, it is recommended that the device be used in conjunction with a mechanism to monitor pressure, an inflation device with pressure gauge.
	The Stents are supplied sterile, by ethylene oxide gas, and are intended for single use only. The stents are invasive and intended for permanent implant by an adequately trained/experienced healthcare professional.
Reference to previous generation(s) or variants	N/A



Accessories which are intended to be used in combination with the device	All Stents are designed to be used with the hemostasis valve tools that are provided with the stents.
Description of any other devices and products which are intended to be used in combination with the device	All Stents are designed to be used with a balloon catheter, introducer, and guidewire.

4. Risks and Warning	
	All significant risks were considered, mitigated as far as possible (AFAP), and are acceptable in regard to the clinical benefit of the device.
	The clinical investigations performed on the CP Stent family of devices reported the following side effects: COAST: aortic aneurysm, stent fracture COAST II: iliac artery dissection PARCS: stent malposition, stent embolization
	The literature reported the following side effects: Acute wall rupture / dissection, aortic aneurysm / pseudoaneurysm, balloon rupture, death, stroke, stent embolization, groin hematoma, late lumen loss, left hemothorax, stent displacement, stent fracture, stent malposition, transitory arrhythmia, and cardiogenic / septic shock.
	Known and foreseeable clinical risks have been considered in accordance with risk management (RM) procedure AP-346 and through the RM files and mitigated as far as possible (AFAP).
	POTENTIAL COMPLICATIONS/ADVERSE EFFECTS
Residual risks and undesirable effects	NOTE: Circumferential tear of the delivery balloon catheter prior to complete expansion of the stent may cause the balloon to become tethered to the stent, requiring surgical removal. In case of rupture of an adequately sized balloon after stent expansion, it can be withdrawn and a new balloon catheter exchanged over a guidewire to complete expansion of the stent.
	Cardiac catheterization carries certain risks. Potential complications & adverse effects associated with device use and indication include:
	<ul> <li>Femoral artery injury</li> <li>Stent Migration</li> <li>Stent Fracture</li> <li>Aortic Rupture/Tear</li> <li>Hematoma</li> <li>Thrombosis</li> <li>Embolization</li> <li>Death</li> <li>Endocarditis</li> <li>Stent Stenosis</li> <li>Aneurysm / Pseudoaneurysm</li> <li>Stent Malposition</li> <li>Sepsis/infection</li> <li>Transitory arrhythmia</li> <li>Bleeding</li> <li>Cerebrovascular Incident</li> </ul>
Warning and Precautions	<ul> <li>The following Warnings and Precautions have been identified and are called out in the Instruction for Use:</li> <li>COVERED STENT WARNINGS</li> <li>Radiofrequency heating during MRI scans on overlapped, 10 zig Stents has not been evaluated.</li> <li>As with any type of implant, infection secondary to contamination of the stent may lead to aortitis, or</li> </ul>



abscess. The platinum/iridium stent may migrate from the site of implant. Over-stretching of the artery may result in rupture or aneurysm formation.
<ul> <li>When the stent is crimped onto a balloon delivery catheter, the maximum balloon inflation pressure</li> </ul>
must not exceed the recommended inflation pressure specified in the manufacturer's instructions.
• The inflated diameter of the stent should at least equal the diameter of the intended implant site.
• Excessive force while crimping may weaken welds of the stent.
• Crimping the 8 zig stent on a balloon catheter smaller than 12mm, and the 10 zig on a balloon catheter
<ul><li>smaller than 26mm, may cause damage to the stent.</li><li>Excessive handling and manipulation of the covering while crimping the stent may cause the covering</li></ul>
to tear off of the stent.
• Crimping the device in the opposite direction of the folds in the covering may cause the covering to
catch while inserting into the hemostasis valve tool and introducer. This could cause the covering to
tear off of the stent.
• Pulling the Covered Stent back through the introducer and/or hemostasis valve may cause the covering
<ul><li>to catch and tear off of the stent.</li><li>This device is intended for single use only. Do not resterilize and/or reuse it, as this can potentially</li></ul>
result in compromised device performance and increased risk of cross contamination.
result in compromised de rice performance and mercased risk of cross containmaton.
COVERED MOUNTED STENT WARNINGS
• Radiofrequency heating during MRI scans on overlapped, 10 zig Stents has not been evaluated.
• As with any type of implant, infection secondary to contamination of the stent may lead to aortitis, or
<ul><li>abscess.</li><li>The platinum/iridium stent may migrate from the site of implant.</li></ul>
<ul> <li>Over-stretching of the artery may result in rupture or aneurysm formation.</li> </ul>
• The inflated diameter of the stent should at least equal the diameter of the intended implant site.
Excessive force while crimping may weaken welds of the stent.
• Crimping the 8 zig stent on a balloon catheter smaller than 12mm, and the 10 zig on a balloon catheter
smaller than 26mm, may cause damage to the stent.
• Excessive handling and manipulation of the covering while crimping the stent may cause the covering to tear off of the stent.
• Crimping the device in the opposite direction of the folds in the covering may cause the covering to
catch while inserting into the hemostasis valve tool and introducer. This could cause the covering to
tear off of the stent.
• Pulling the Covered Stent back through the introducer and/or hemostasis valve may cause the covering
to catch and tear off of the stent.
BIB STENT PLACEMENT WARNINGS
• Do not exceed the RBP. An inflation device with pressure gauge is recommended to monitor pressure.
Pressure in excess of the RBP can cause balloon rupture and potential inability to withdraw the catheter
through the introducer sheath.
<ul> <li>Confirm that the distal end of the introducing sheath is at least 2.5cm back from the most proximal image markers before inflating the outer balloon. Failure to do so may stretch the outer tubing and</li> </ul>
severely hinder balloon deflation.
<ul> <li>Use two appropriate size inflation devices with pressure gauges for inflation.</li> </ul>
• Do not advance the guidewire, balloon dilatation catheter, or any other component if resistance is met,
without first determining the cause and taking remedial action.
• This catheter is not recommended for pressure measurement or fluid injection.
• Do not remove the guidewire from the catheter at any time during the procedure except when the procedure has been completed.
<ul> <li>This device is intended for single use only. Do not resterilize and/or reuse it, as this can potentially</li> </ul>
result in compromised device performance and increased risk of cross contamination.
COVERED STENT PRECAUTIONS
• Use of an inflation device with pressure gauge is highly recommended during this procedure.
• Stents are delicate devices. Care should be exercised while handling to help prevent the possibility of breakage
breakage.



	<ul> <li>The stent is rigid and may make negotiation through vessels difficult.</li> <li>Dilatation procedures should be conducted under fluoroscopic guidance with appropriate x-ray equipment.</li> <li>Guidewires are delicate instruments. Care should be exercised while handling to help prevent the possibility of breakage.</li> <li>Careful attention must be paid to the maintenance of tight catheter connections and by aspiration before proceeding to avoid air introduction into the system.</li> <li>Under no circumstances should any portion of the catheter system be advanced against resistance. The cause of the resistance should be identified with fluoroscopy and action taken to remedy the problem.</li> </ul>
	<ul> <li>COVERED MOUNTED STENT PRECAUTIONS</li> <li>Use of an inflation device with pressure gauge is highly recommended during this procedure.</li> <li>Stents are delicate devices. Care should be exercised while handling to help prevent the possibility of breakage</li> </ul>
	<ul> <li>breakage.</li> <li>The stent is rigid and may make negotiation through vessels difficult.</li> <li>Dilatation procedures should be conducted under fluoroscopic guidance with appropriate x-ray equipment.</li> <li>Guidewires are delicate instruments. Care should be exercised while handling to help prevent the</li> </ul>
	<ul> <li>Outdownes are deleate instruments. Care should be excretised while handling to help provent the possibility of breakage.</li> <li>Careful attention must be paid to the maintenance of tight catheter connections and by aspiration before proceeding to avoid air introduction into the system.</li> <li>The inflation diameter of the balloon used during stent delivery should approximate the diameter of the</li> </ul>
	<ul> <li>obstructive vessel and the intended implant site.</li> <li>Under no circumstances should any portion of the catheter system be advanced against resistance. The cause of the resistance should be identified with fluoroscopy and action taken to remedy the problem.</li> <li>If resistance is felt upon removal, then the balloon, guidewire and the sheath should be removed together as a unit, particularly if balloon rupture or leakage is known or suspected. This may be accomplished by firmly grasping the balloon catheter and sheath as a unit and withdrawing both together, using a gentle twisting motion combined with traction.</li> <li>The balloons must be completely deflated before retracting into the sheath.</li> <li>Proper functioning of the catheter depends on its integrity. Care should be used when handling the catheter.</li> </ul>
Other relevant aspects of safety, including a summary of any field safety corrective actions (FSCA including FSN) if applicable	There have not been any Field Safety Corrective Actions or Field Safety Notices on any versions of the Stents listed in this SSCP.

#### 5. Summary of clinical evaluation and post-market clinical follow-up (PMCF)

#### Summary of clinical data related to equivalent device:

An equivalent device was not used for the clinical evaluation.

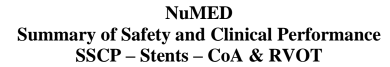
#### Summary of clinical data from conducted investigations of the device:

#### 1. Study name: COAST

Purpose: to provide information that will support labeling of both the CP bare metal and covered stents to treat native and recurrent

CoA in selected children, adolescents and adult.

**Clinical Study Methodology:** Single arm interventional study (open label). The COAST is a prospective, multicenter, single-arm clinical study involving 19 pediatric cardiology centers in the United States. The study includes patients with native or recurrent CoA treated by physicians at the participating institutions. A total of 105 patients underwent attempted implantation, with 104 successes. *Reference to the clinical study plan (and amendment) n*<sup>•</sup>: NCT00552812



	e: 19 pediatric		Approval: Institutional	Regulatory Authority Approvals:		
cardiology centers in United States Review Board a		Review Board appr	ovals from all	Investigational Device Exemption from US FDA		
participating instituti		tions	(August 3, 2007)			
Patient Populati	on: Patients with n	ative or recurrent CoA	A. A total of 105 patients u	nderwent attempted implantation, median age 16		
years (range from	n 8 to 52 years) and	with 69.5% male.				
Clinical Study R	esults: Results held	d on file by Sponsor				
Purpose	Criteria		Results			
Performance	Blood pressure gradient and coarctation minimum diameter: cardiac catheterization before and after CP Stent placement		29±14 mmHg at baselin	pressure difference (mmHg) changed from ne to $-3\pm15$ mmHg at 24 months follow-up. um diameter reported at $7.9 \pm 2.7$ mm at fter implantation.		
Safety	Adverse events			nts reported, 7% of the patients experienced		
				ts. ): 5 were successfully treated with covered resolved without intervention.		
			Stent fractures were see	en in 2 patients after one year, 11 patients at onal fractures above 2 years.		
	• •	ort n•: NCT00552812				
Device Used: Ba	re CP Stent and BII	B catheter; covered ste	ents were available in case	of aortic wall injury.		
Conclusion: The	CP stent is safe and	d associated with pers	istent relief of aortic obstr	uction. Stent fracture and progression of fracture		
occur but have no	ot resulted in clinics	lly important carnela	. D.:			
	n resulted in chillea	my important sequera	e. Reintervention is comm	on and related to early and late aortic wall injury		
			e. Reintervention is comm	on and related to early and late aortic wall injury		
and need for re-ea	xpansion of small-d		e. Reintervention is comm	on and related to early and late aortic wall injury		
and need for re-e: 2. Study name: Purpose: To eva	xpansion of small-d : COAST II	liameter stents.		reventing aortic wall injury in patients with aortic		
and need for re-ex 2. Study names Purpose: To eval coarctation Clinical Study M aortic wall injury and angiography reported.	xpansion of small-d : COAST II luate safety and sho fethodology: Singl (Treatment group) were reported. A co	rt-term efficacy of the e arm interventional s or with increased risk	e CP Stent in treating or pr tudy. Patients were enrolle of aortic wall injury (Pre- red standardized review of			
and need for re-e: 2. Study name: Purpose: To eval coarctation Clinical Study M aortic wall injury and angiography reported.	xpansion of small-d : COAST II luate safety and sho fethodology: Singl (Treatment group) were reported. A co clinical study plan	rt-term efficacy of the e arm interventional s or with increased risk ore laboratory perform	e CP Stent in treating or pr tudy. Patients were enrolle of aortic wall injury (Pre- ned standardized review of : NCT01278303	reventing aortic wall injury in patients with aortic ed if they had a history of CoA with pre-existing vention group). Pre/post-implant hemodynamics		
and need for re-ex 2. Study names Purpose: To evaluation Clinical Study M aortic wall injury and angiography reported. Reference to the Investigation site	xpansion of small-d : COAST II luate safety and sho fethodology: Singl (Treatment group) were reported. A co clinical study plan e: 19 pediatric	iameter stents. ort-term efficacy of the e arm interventional s or with increased risk ore laboratory perform (and amendment) n <sup>•</sup> . Ethics Committee	e CP Stent in treating or pr tudy. Patients were enrolle of aortic wall injury (Pre- ned standardized review of : NCT01278303	reventing aortic wall injury in patients with aortic ed if they had a history of CoA with pre-existing vention group). Pre/post-implant hemodynamics all angiograms. One month follow-up was		
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and need for re-ex 2. Study names Purpose: To evaluation Clinical Study M aortic wall injury and angiography reported. Reference to the Investigation situ cardiac centers in	xpansion of small-d : COAST II luate safety and sho fethodology: Singl (Treatment group) were reported. A co clinical study plan e: 19 pediatric United States	iameter stents. rt-term efficacy of the e arm interventional s or with increased risk ore laboratory perform (and amendment) n <sup>•</sup> . Ethics Committee Hopkins Institution Institutional review participating center	e CP Stent in treating or pr tudy. Patients were enrolle of aortic wall injury (Prev ed standardized review of : NCT01278303 Approval: Johns al Review Board and boards of all s.	reventing aortic wall injury in patients with aortic ed if they had a history of CoA with pre-existing vention group). Pre/post-implant hemodynamics all angiograms. One month follow-up was <b>Regulatory Authority Approvals:</b> Investigational Device Exemption from US FD/		
and need for re-ex 2. Study names Purpose: To evaluation Clinical Study M aortic wall injury and angiography reported. Reference to the Investigation site cardiac centers in Patient Populati	xpansion of small-d : COAST II luate safety and sho fethodology: Singl (Treatment group) were reported. A co clinical study plan e: 19 pediatric United States on: Patients with ac	iameter stents. ort-term efficacy of the e arm interventional s or with increased risk ore laboratory perform (and amendment) n <sup>•</sup> . Ethics Committee Hopkins Institution Institutional review participating center ortic coarctation at risk	e CP Stent in treating or product of a ortic wall injury (Prevated standardized review of NCT01278303 Approval: Johns al Review Board and boards of all s.	reventing aortic wall injury in patients with aortic ed if they had a history of CoA with pre-existing vention group). Pre/post-implant hemodynamics all angiograms. One month follow-up was <b>Regulatory Authority Approvals:</b> Investigational Device Exemption from US FD4 with existing aortic wall injury. A total of 158		
and need for re-ex 2. Study names Purpose: To eval coarctation Clinical Study M aortic wall injury and angiography reported. Reference to the Investigation site cardiac centers in Patient Populati patients (83 treatu	xpansion of small-d : COAST II luate safety and sho fethodology: Singl (Treatment group) were reported. A co clinical study plan e: 19 pediatric United States on: Patients with ac	iameter stents. ort-term efficacy of the e arm interventional s or with increased risk ore laboratory perform (and amendment) n <sup>•</sup> . Ethics Committee Hopkins Institution Institutional review participating center ortic coarctation at risk	e CP Stent in treating or product of a ortic wall injury (Prevated standardized review of NCT01278303 Approval: Johns al Review Board and boards of all s.	reventing aortic wall injury in patients with aortic ed if they had a history of CoA with pre-existing vention group). Pre/post-implant hemodynamics all angiograms. One month follow-up was <b>Regulatory Authority Approvals:</b> Investigational Device Exemption from US FD/		
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and need for re-e: 2. Study name: Purpose: To evaluation Clinical Study Marcel aortic wall injury and angiography reported. Reference to the Investigation site cardiac centers in Patient Populati patients (83 treatu females. Clinical Study R Purpose	xpansion of small-d : COAST II luate safety and sho fethodology: Singl (Treatment group) were reported. A co clinical study plan e: 19 pediatric United States on: Patients with ac ment cohort and 75 Results: Results held Cri	iameter stents. ort-term efficacy of the e arm interventional s or with increased risk ore laboratory perform (and amendment) n <sup>•</sup> . Ethics Committee Hopkins Institution Institutional review participating center ortic coarctation at risl prevention cohort, me d on file by Sponsor iteria	e CP Stent in treating or provide cP Stent in treating or provide a standardized review of a critic wall injury (Previded standardized review of a standardized review of a NCT01278303 <b>Approval:</b> Johns al Review Board and a boards of all s. k of a ortic wall injury or we adian age 19 years (range for a standard sta	reventing aortic wall injury in patients with aortic ed if they had a history of CoA with pre-existing vention group). Pre/post-implant hemodynamics all angiograms. One month follow-up was <b>Regulatory Authority Approvals:</b> Investigational Device Exemption from US FDA with existing aortic wall injury. A total of 158 from 5 to 70 years) and with 103 males and 55		
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Device Used: Covered CP Stent by NuMED, pre-mounted on BIB stent delivery catheter.

NuMED

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**Conclusion:** The CP Stent can effectively treat and potentially prevent aortic wall injury associated with aortic coarctation. Access site arterial injury is the most common important complication. Longer-term follow up is necessary to define mid- and late-term outcomes.

#### 3. Study name: PARCS

Purpose: Evaluation of the Covered CP Stent for repair of tears that occur in the pulmonary artery during dilation (enlargement) of a

conduit (passageway) connecting the right ventricle of the heart to the pulmonary arteries.

**Clinical Study Methodology:** Single arm prospective study. The PARCS trial was a prospective, multicenter, single-arm pivotal clinical trial. Forty US centers participated in either the pivotal trial (22 centers) or the continued access protocol, which immediately followed the pivotal trial during Food and Drug Administration submission. If no conduit wall injury occurred during the procedure, the patient was considered a screen failure. If at any point during the procedure, including before intervention, the implanting physician identified an area of wall injury, a CCPS could then be selected and implanted.

Reference to the clinical study plan (and amendment) n<sup>•</sup>: NCT01824160

Patient Population: Participants receiving a Covered CP stent for repair of pulmonary artery injury. In the pivotal trial, fifty patients

met the inclusion criteria, mean age 17 years (range from 6 to 44 years) and 56% of male patients. In the continued access, seventy

patients with mean age of 16 years (range from 7 to 49 years) and 57% of male patients.

Clinical Study Results: Results held on file by Sponsor

Purpose	Criteria	Results
Pivotal (n=50): Severity of illness	Median improvement by at least 1 level	Median improvement of 1 level in Severity
	from baseline to post-procedure	of Illness Score
Pivotal (n=50): Procedure success	$\geq$ 75% patients, based on both device	Procedure success achieved in 68% of
	success and lesion success	patients
Pivotal (n=50): Successful implantation	Coverage of conduit disruption defined	Successful implantation achieved in 83%
of the Melody TPV	as either no residual disruption or	of patients
	contained disruption, followed by	
	successful implantation of Melody valve	
	in $\ge 80\%$ patients	
Pivotal (n=50): Adverse events attributed	$\geq$ 80% patients free of adverse events	At least 80% were free of an adverse event
to covered CP Stent within 30 days	attributed to the covered CP Stent within	attributed to the covered CP Stent. There
	30 days	was 1 report of stent malposition where the
		stent became dislodged and migrated into
		the pulmonary arteries
All patients (n=120): Performance	Covered CP Stent Implant success	CCPS implants successfully treated 95% of
		conduit injuries with either no or minimal
		residual conduit wall injury.
		Melody TPVR was successfully performed
		in 94% of the enrolled cohort, and TPV
		function was not adversely affected by
		placement within the CCPS substrate, with
		6-month follow-up data comparing
		favorably with other previously published
		cohorts.
All patients (n=120): Safety	Stent-related AEs	AEs that specifically related to the CCPS
		and its implantation were uncommon. One
		serious (stent malposition) and one
		somewhat serious (stent embolization) AE
		occurred (both in the same patient who is
		described above). A device usage issue wa
		identified whereby the expanded poly
		tetrafluoroethylene covering separated
		from the stent during attempts to load the
		CCPS device into the delivery sheath. This
		was identified before deployment; the sten
		was removed and replaced with a new
		CCPS without consequence to the patient.



*Reference to the Clinical Study Report n*<sup>•</sup>: NCT01824160

Device Used: Covered CP Stent pre-mounted on BIB

**Conclusion:** The study results demonstrate the safety and efficacy of use of the covered CP Stent when used for pre-stenting in the RVOT prior to Melody TPV implantation.



Summary of clinical data f	rom other sources:							
First Author (Year)	Appraisal/Results							
	Transcatheter Pulmona	its the results from the PAR ary Valve Replacement (NCT ne state of the art informati	Г01824160). Please refer		-			
	Medical condition	Alternatives	Risk/benefit	Side-effects		Equivalence	Surrogate	endpoints
	Yes 1 No 2	Yes 1 No 2	Yes 1 No 2	Yes 1	No 2	Yes 1 No 2		No 2
	Overall SOA Appraisal SOA Grade (Range 6-12)	and Disposition		Disposition	(select)		Accepted, < 1 Excluded, 12	2
1. Delaney et al. (2018) Contribution S&P x SOA x	Relevant SOA Results SOA data	<ul> <li>required t the condu</li> <li>Stenting o</li> <li>Covered s</li> <li>RVOT reconstrution</li> <li>RVOT reconstrution</li> <li>RVOT reconstrution</li> <li>All valved</li> <li>50% and 8</li> <li>RVOT dyst</li> <li>Transcath</li> <li>been utilizion</li> <li>Successfution</li> <li>Successfution</li> <li>Successfution</li> <li>Melody transca</li> </ul>	conduit injury can occur o dilate conduits effectiv it out of concern for exte of the conduit before valv tents have been used in t	ely for TPVR. Con ension of the area e implantation in the vascular space conduit or biopreset of patients with less of type, have nt by 10 years. ed with substant ilitation using hig need for surgical successful conduit raphy. ty often requires with a higher rat vith hemodynam	nduit injury, o a of injury. mproves the o ce to isolate a rosthetic pulm ith congenital e been associa cial patient mo gh-pressure a l pulmonary v it dilation, altl s the use of ul te of recogniza- nic compromis	once identified, cou durability of the im reas of injury. nonary valve placed l heart disease. ated with functiona orbidity and even n ngioplasty with or ralve replacement. hough minor injuri trahigh pressure ne ed conduit injury (se	h pressure angiop Id preclude furth planted valve. ment is necessary al deterioration, v nortality. without stent pla An injury within t es may not be clin oncompliant ballo ≤33%). The vast n	er dilation of during vith between cement has he wall of the nically pons to najority of



	Comments	<ul> <li>with a high rate of progressive valve</li> <li>Conduit wall injury is a known comp Although bare metal stents may pro allow for safe, continued dilation of hemodynamically important residu catastrophic conduit injuries.</li> <li>Covered CP Stent (NuMED) is a ball applications for vascular wall injury Stent outside of the United States is valve implantation. The European e conduit injury.</li> <li>Some US centers did have access to Stent Trial) and could apply for eme could apply for a single-patient com injury.</li> <li>High-pressure balloon and stent angiopli before transcatheter pulmonary valve re patient or prevent successful TPVR. Seve was effective in either treating or mitiga</li> </ul>	t reinforcement of the conduit before valve implant, hav e deformity and stent fracture leading to valvular dysfum olication of isolated or serial balloon angioplasty of the R povide some reinforcement of a damaged conduit wall, th f an injured RVOT conduit that has not been fully prepare al stenosis) for TPVR, and they are not anticipated to be oon-expandable, large-diameter, covered stent whose co , tears, or leak have been reported previously. Experience s extensive and has included its routine use in the pre-ste experience has suggested that this practice may reduce th of the Covered CP Stent as participants in the COAST (Coal ergency use if an unexpected RVOT wall injury occurred. Inpassionate use exemption if they felt a patient was at hi esty are frequently necessary to prepare the dysfunction eplacement (TPVR). Conduit injury can result, which may ere conduit injury was found to be rare but unpredictable ting this problem. The vast majority of patients, even wite eplacement procedure. The covered stent did not interfer up.	ction. VOT con ey are n ed (e.g., effective onstruct e with the enting p ne clinica rctation Non-CO. gh risk f al RVOT be catas e. The co ch identi	iduit. ot likely left with ion and ne Cove rocess fo al impac of the A AST cent or conduit strophic vered st fied con	to ting red CP or t of orta ters uit to the cent duit
	Safety & Performance (for	or safety only)				
	Appraisal					
	Level of Evidence	Study Method/Design	Question Applied	Oxfo	rd LOE 2	011
2. Baykan et al. (2018) Contribution S&P X (S		Control study. Study group was composed of 20 CoA patients who were treated with CP Stent between the dates October 2008 and February 2015, and control group was composed of 20 healthy children with age and sex matched.	To address the presence of hypertension and risk for cardiovascular diseases in patients with CoA who were treated with endovascular stent placement.	1 2	2 3	4 5
only)	Suitability	Relevant Data			Grading	g
SOA -	Device	- CP Stents (Bare and Covered)		D1	D2	, D3
		<ul> <li>Unknown whether pre-mounted on BIB</li> </ul>				
	Application	- CoA		A1	A2	A3
			ment for CoA compared with control group (beatthe			
	Patient	<ul> <li>Patients who had undergone stent place children with age and sex matched).</li> </ul>	ement for CoA compared with control group (healthy	P1	P2	Р3



Report	<ul> <li>Sampling: n=20 CoA and n=20 healthy children</li> <li>Mean age: <ul> <li>CoA group: 14.2 (SD: 3.9) years</li> <li>Control group: 13.7 (SD: 2.7) years</li> </ul> </li> <li>Sex: <ul> <li>CoA group: 16M; 4F</li> <li>Control group: 15M; 5F</li> </ul> </li> <li>High quality <ul> <li>A the late to a to be</li> </ul> </li> </ul>	R1	R2	R3
	Suitability Grade (Range 4-12)		4	
Data Contribution	Relevant Data		Gradin	Ig
Outcomes/Endpoints	Ambulatory blood pressure	Yes 1		No 2
Follow-up	- 6 months and 6 years	Yes 1		No 2
Statistical analysis	<ul> <li>Student t-test was used if the two independent group comparisons were normal and the Mann-Whitney U test was used if the normal distribution was not present. Pearson chi-square analysis was performed to determine whether there was a difference in categorical variables between the case and control groups.</li> </ul>	Yes 1	1	No 2
Clinical significance	<ul> <li>It was shown that hypertension incidence as demonstrated by ambulatory blood pressure monitorization and risk for cardiovascular diseases as indicated by carotid intima media thickness and pulse wave velocity were higher than those in healthy population even after CoA is corrected.</li> <li>CoA should be carefully monitored for hypertension, even if it has been completely corrected by any method. This study suggests that CoA is a part of generalized vasculopathy rather than being a localized narrowing.</li> </ul>	Yes 1		No 2
	Data Contribution Grade (Range 4-8)		4	
Overall S&P Appraisal, Di S&P Grade (Range 9-25)	Image: Seposition and Weighting       LOE (3) + Suitability (4) +       Disposition and Weighting (select)       Accepted and Accepted but Excluded, 22-         Data Contribution (4) = 11       Excluded, 22-       Excluded, 22-	not Pivo		-21
Relevant S&P Results				
Safety data	<ul> <li>Hypertensive:</li> <li>Daytime: 5% were hypertensive and 20% were pre-hypertensive in the study group compare</li> </ul>	ed to 0%	in the	
	<ul> <li>control group.</li> <li>Night: 15% were hypertensive and 15% were pre-hypertensive in the study group compared group.</li> </ul>	to 0% ii	n the co	ontrol
Benefits/claims data	<ul> <li>Night: 15% were hypertensive and 15% were pre-hypertensive in the study group compared group.</li> <li>N/A</li> </ul>	to 0% ii	n the co	ontrol
Benefits/claims data Strengths Weaknesses/	<ul> <li>Night: 15% were hypertensive and 15% were pre-hypertensive in the study group compared group.</li> </ul>			



			nitoring with Holter device) in pre-operat	tive period could	not be	used	
		because at that time they did not have	a blood pressure Holter device.				
	Safety & Performance						
	Appraisal					1.0-	
	Level of Evidence	Study Method/Design	Question Applied			rd LOE	-
		Retrospective data collected of the first	To evaluate the first-in-man use of a n		1 2	3	4 5
		NuDEL delivery systems used in patients	(NuDEL) for implantation of CP Stent (	•			
		from three centers (UK and Ireland).	patients with complex structural and C	.HD.			
	Suitability	Relevant Data				Gradir	ופ
	Device	- NuDEL Delivery System			D1	D2	D3
	Application	- CoA and RVOT			A1	A2	A3
	Patient	- Patients with COA and RVOT			P1	P2	P3
		- Sampling: n=12 (13 CP Stents, Covered	, delivered via 12 NuDELs); with 6 CoA, 5 I	RVOT, and 1			
			emic venous baffle. Note: "P2" due to one				
		stenosis of a Mustard systemic venous					
		- Age: 10-43 years					
		- Sex: Not reported					
<ol> <li>Morgan et al.</li> </ol>	Report	- High quality with minor deficiency as d	evice performance is based on descriptive	e information.	R1	R2	R3
(2017)			Suitability Gra	de (Range 4-12)		6	
					1		
Contribution	Data Contribution	Relevant Data				Gradir	-
S&P x	Outcomes/Endpoints	- Procedure complications.			Yes 1		No 2
SOA x		- Ease of use.					
	Follow-up	- Not reported.			Yes 1		No 2
	Statistical analysis	- Not reported.			Yes 1		No 2
	Clinical significance		eans of covered stent deployment in cha	llenging	Yes 1		No 2
		anatomy.					
			Data Contribution Gra	ade (Range 4-8)		6	
	Overall S&P Appraisal, Dis	position and Weighting					
	S&P Grade	LOE (4) + Suitability (6) +	Disposition and Weighting (select)	Accepted and	Pivotal	9-12	
	(Range 9-25)	Data Contribution (6) = 16	, , ,	Accepted but			3-21
				Excluded, 22-2			
	Relevant S&P Results			•			
	Safety data	- No procedural complications and no re	ports of equipment failure or dysfunction				
	Performance data	- The system required minimal preparati	on – flushing only; therefore, despite a lac	ck of familiarity, i	it was re	eady fo	r
		deployment in each case in under two					
			were that the assembly tracked well thro		ite and	throug	gh
		tortuous and narrowed anatomy in eith	er the outflow tract or the descending ac	orta			



	laims data	-	deploying it. T NuDEL reporte anatomy.	part of the car o this end, the ed to require r	therization p e NuDEL syst ninimal prep	rocedure is g em has been paration and t	etting the ste developed. racked well t	ent into the re	equired ana access site a	tomical positic	d narrowe
Strengths		-	patients with o associated wit Stent was easy	complex outfl h stent moun y to uncover a	ow tract ster ting and mar	nosis and thos nagement of t	se with CoA. the stent–bal	Using this sys Iloon–deliver	stem avoids y system co		
Weakness Potential		-	Conflict of inte Financial supp sectors		received no	specific grant	t from any fu	nding agency	or from cor	mmercial or no	ot-for-profi
<u>State of the</u> Appraisal	<u>e Art</u>										
Medical c	ondition	Alternat	ives	Risk/bene	fit	Side-effec	ts	Equivale	nce	Surrogate	endpoints
Yes 1	No 2	Yes 1	No 2	Yes 1	No 2	Yes 1	No 2	Yes 1	No 2	Yes 1	No 2
SOA Grad	e	and Disposit	ion			Dispositic	on (select)			Accepted, < 1 Excluded, 12	2
SOA Grad (Range 6-:	e 12)		ion			Dispositio	on (select)			• •	2
SOA Grad	e 12) DA Results	10	Conduit ruptu system may pr of large-calibe The range of s make semi-qu Safety and acc balloon and pa catherization p taken in gettin step. Slipping o the balloon or target areas, lo major safety c	rovide an attra r stent proced tents available antitative dec curacy of deple assing it into a procedure is g og these essen of the stents of stent during i ead to vascula onsequences.	active emerge dures and arc e for these the isions about oyment are a nd along the etting the st tial steps rig off balloons l mounting arc r risk of rem	potential con ency backup. e not converse herapies has o stent choice at least partia e delivery she ent into the r ht, and there eading to mig e some of the oving and re-	nplication; th This may be ant with the developed we for each indiv Ily dependen ath to its req equired anat is potential f gration of the problems er inserting a lo	of benefit to techniques ir ell over the la vidual case. at on the prec uired position omical position for safety, eff stent before noountered, v ong, large-cali	operators winvolved, even ast 10–15 yes cise mountin n. The most on before during d or during d which may re- iber sheath,	• •	d stent low volum ve setting. uthors to on its deliv f the ot of time i ems at eve d damage t sing of the an have



	Appraisal					
	Level of Evidence	Study Method/Design	Question Applied	Oxfo	rd LOE	2011
		Retrospective review of incidence and potential predictors of conduit disruption.	To assesses the frequency of RVOT conduit disruption during transcatheter pulmonary valve replacement (TPVR) and the effectiveness and safety of NuMED Covered Mounted CP Stentsfor its prevention or treatment.	1	2 3	4 5
	Suitability	Relevant Data			Gradi	ng
	Device	<ul> <li>Covered Mounted CP Stents (12 to 22m)</li> </ul>	im)	D1	D2	D3
	Application		tion or treatment of RVOT conduit disruption during	A1	A2	A3
Bishnoi et al. (2015) Contribution S&P x SOA x	Patient	<ul> <li>Population: Patients undergoing TPVR r (patients with pre-existing tears, patien and patients developed tears after tran prophylactically placed in patients of pe and/or severity of homograft stenosis).</li> <li>Sampling: 50 patients receiving 69 Cove (comparative cohort: 251 implants US N for bare metal stenting of supported co</li> </ul>	ered CP Stents during TPVR/PPVI procedures Melody transcatheter pulmonary valve IDE Trial, planned nduit) ption requiring intervention was 6% in the study.	P1 R1	P2 R2 <b>R</b> 2 <b>4</b>	P3 R3
		1	, , ,			
	Data Contribution	Relevant Data			Gradi	0
	Outcomes/Endpoints	<ul> <li>Peak-to-peak RVOT gradient</li> <li>Mean Doppler RVOT Gradient at 6 mor</li> <li>Valve competence with no or trivial pu</li> <li>Safety</li> </ul>		Yes 1		No 2
	Follow-up	- 6 months		Yes 1		No 2
	Statistical analysis	- Not provided		Yes 1		No 2
	Clinical significance	- Covered Mounted CP Stent implantation without negative impact on the transca	on can successfully treat RVOT conduit disruption attraction attra	Yes 1		No 2
			Data Contribution Grade (Range 4-8)		6	



(Range 9-2	25)	Data Contribution (	6) = 13						ed but not Piv d, 22-25	otal, 13-2
Relevant S&	&P Results									
Safety dat	a	<ul> <li>Stent fracture:</li> <li>implantation. I</li> <li>portions of the</li> <li>No Covered CF</li> </ul>	Patient was CP Stent (C	successfully t overed).	reated with in	mplantation c				
Performar	nce data	<ul> <li>Peak-to-peak F</li> <li>Mean Doppler</li> <li>IDE trial.</li> <li>Valve compete</li> <li>comparator gr</li> <li>Conduit tears:</li> </ul>	RVOT gradie RVOT Gradi ence with no oup.	nt: Decreased ent at 6 mon or trivial pul	l from 45.5 ± ths: 12.86 ± 5 monary regur	17.5 mm Hg t .0 mmHg con gitation: At fo	npared to 20.	0 ± 8.6 mm	-	
Benefits/c	laims data	- N/A		•						
Strengths		CCPS implanta     CCPS implanta     This retrospect     during PPVI.     Retrospective	ive analysis	suggests hig	n RVOT condu	uit systolic pre	essure gradier	nt is a risk fa	actor for cond	uit tears
Potential I State of the Appraisal		- Kreutzer	small and in rest reporte N: none esearch grar : research g	d: It and consul rant support	tant for Medt from Medtro	ronic; resear	ch grant supp de Medical; c	ort from Nu		Inc.
Medical co	andition	Alternatives	Risk/ben	ofit	Side-effe	rte	Equivale	nce	Surrogate	e endpoint
Yes 1	No 2	Yes 1 No 2	Yes 1	No 2	Yes 1	No 2	Yes 1	No 2	Yes 1	No 2
SOA Grade	2	nd Disposition		-		on (select)		-	Accepted, < 2	12
SOA Grade (Range 6-2 Relevant SC SOA data	12)	9 - Surgical mana arteries, trunci implantation o	us arteriosus	s, and those ι	CHD such as Indergoing Ro	tetralogy of oss procedure			Excluded, 12	on of th



	Comments	<ul> <li>compression and somatic outgrowth on</li> <li>Endovascular treatment using balloon</li> <li>lifespan and reduce a patient's need for</li> </ul>	dilatation and bare stent implantation has been show	vn to e	xtend	conduit
	Safety & Performance					
	Appraisal					
	Level of Evidence	Study Method/Design	Question Applied	Oxfor	d LOE	2011
		Prospective randomized controlled trial.	To evaluate outcomes of treatment with Covered versus Bare NuMED CP Stents.	1 2	3	4 5
	Suitability	Relevant Data			Gradir	Ig
	Device	<ul> <li>NuMED CP Stent (Bare and Covered)</li> <li>Stent was hand-crimped down onto BIE</li> </ul>	3	D1	D2	D3
	Application	- Severe native CoA		A1	A2	A3
5. Sohrabi et al.	Patient	<ul> <li>Patients with severe native CoA</li> <li>Sampling: n=120 (60 CP Stents versus 6</li> <li>Mean age: 23.6±10.99 (range 12 to 58)</li> <li>Sex: 79 M; 41 F</li> </ul>		P1	P2	Р3
(2014)	Report	- High quality.		R1	R2	R3
Contribution			Suitability Grade (Range 4-12)		4	
S&P x	Data Cantaliantian	Delevent Dete			Cure all'a	-
SOA -	Data Contribution	Relevant Data			Gradir	-
	Outcomes/Endpoints	<ul> <li>Procedural success</li> <li>Reduction in systolic blood pressure graves</li> <li>Reduction in mean diameter of coarcta</li> <li>Adverse effects</li> </ul>		Yes 1		No 2
	Follow-up	- 31.1 ± 19.2 months		Yes 1		No 2
	Statistical analysis	- A p-value <0.05 was considered signific	cant.	Yes 1		No 2
	Clinical significance	<ul> <li>remarkable hemodynamic effects in se</li> <li>complication during the procedure and</li> <li>Patients undergoing CP Stent (Covered</li> </ul>	) implantation experienced a non-significantly lower re- ce of pseudoaneurysm formation with respect to CP	Yes 1		No 2



		- These findings indicate that CoA stentir	g is a safe procedure.				
			Data Contribution Gra	ade (Range 4-8)		4	
	Overall S&P Appraisal, Di	sposition and Weighting					
	S&P Grade (Range 9-25)	LOE (2) + Suitability (4) + Data Contribution (4) = 10	Disposition and Weighting (select)	Accepted and Accepted but r Excluded, 22-2	not Pivo		-21
	Relevant S&P Results				5		
	Safety data	<ul> <li>Pseudoaneurysms: 0 (CP Stent, Bare) ve</li> <li>Mortality: 1 (CP Stent, Bare) versus 0 (CF</li> </ul>					
	Performance data	<ul> <li>and 3.36 mmHg respectively; no signification</li> <li>Mean diameter of coarctation segment and 15.82 mm respectively; no signification</li> </ul>	atients eduction: from 54.61 (CP Stent, Bare) and ant difference between the two types of reduction: From 3.34 (CP Stent, Bare) and at difference between the two types of st versus 0 (CP Stent, Covered), non-signifi	stent, P<0.001 d 3.30 (CP Stent, 9 ent, P<0.001			
	Benefits/claims data	<ul> <li>Reduction in mean systolic blood pressu</li> <li>Reduction in diameter of coarctation seg</li> </ul>	re gradient				
	Strengths	stent delivery	nto a balloon-in-balloon catheter (NuME				
	Weaknesses/ Potential bias	patients did not undergo 24-hour ambul	al in this respect, study was limited in so atory blood pressure monitoring, which o cond, evaluation of the blood pressure re procedure outcome.	could have diagno	osed th	е	
	Safety & Performance Appraisal						
	Level of Evidence	Study Method/Design	Question Applied			d LOE 2	-
Vanagt et al. (2014)		Single-center retrospective study (CHD database of all CP Stent, Covered, during 2003-2012)	To evaluate possibilities and safety of ( (Covered) in CHD.	CP Stent	1 2	3	4
a sa tu ila sa ti a sa							
Contribution &P x OA -	Suitability Device	Relevant Data         -       Covered CP Stent         -       The stent was hand-crimped on BIB			D1	Gradin D2	g D3
	Application	<ul> <li>CoA and RVOT pre-stenting for percutar</li> </ul>	neous revalvulation		A1	A2	A
	Patient	- Patients with CoA and RVOT pre-stentin CP Stent (Covered) was chosen for deliv	ng for percutaneous revalvulation. For th /ery balloon protection after rupture of t (81%) because tear, rupture, or fracture	he pre-dilation	P1	P2	P3



	<ul> <li>Sampling: n= 51 (CoA group), n=37 (</li> <li>Mean age: <ul> <li>CoA group: 19 (range from 8 to</li> <li>RVOT group: 16 (range from 6 for 5 ex:</li> <li>CoA group: 38M; 13F</li> </ul> </li> </ul>	69) years	ed.			
Poport	- RVOT group: 26M; 11F - High quality.			R1	R2	R
Report		Suitability Grac	$A = (Range A_12)$	N1	4	К
<u> </u>			ie (nalige 4-12)		4	
Data Contribution	Relevant Data				Gradir	ופ
Outcomes/Endpoints	Increase in diameter at coarctation	(CoA group)		Yes 1	1	<u>ю</u> No 2
	- Decrease in peak to peak gradient (	CoA group) ting and pulmonary valve delivery (RVOT Gro	up)			
Follow-up	- Not specified.			Yes 1		No 2
Statistical analysis	- Two-sided p<0.05 was considered s	ignificant.		Yes 1		No 2
Clinical significance		e applied in CHD patients. The covering all s, thereby increasing the safety margin with r	more complete	Yes 1		No 2
		Data Contribution Gra	de (Range 4-8)		5	
S&P Grade	LOE (4) + Suitability (4) +	Disposition and Weighting (select)	Accepted and			
			Accepted and Accepted but Excluded, 22-2	not Piv		3-21
(Range 9-25) Relevant S&P Results	LOE (4) + Suitability (4) +		Accepted but	not Piv		3-21
S&P Grade (Range 9-25)	LOE (4) + Suitability (4) + Data Contribution (5) = 13 - CoA Group: - No acute bleeding, aneurysm fo - Mild procedure related-complic present in left ventricle), and tr present in left ventricle). - During follow-up: no stent fract ischemia or signs of vessel occlu - RVOT group: - No procedure-related complica	Disposition and Weighting (select) prmation or life-threatening complications. cations included groin hematoma (n = 3), tran ansient atrioventricular block with nodal esca cures, nor stent recompression occurred, and usion at the puncture site.	Accepted but Excluded, 22-2	not Pive 25 hm (n = 1, while	1, no vire v	wire was



	Benefits/claims data Strengths Weaknesses/ Potential bias Safety & Performance	<ul> <li>22/37 single procedure and 15/37</li> <li>Graft diameter (RVOT Group)         <ul> <li>Increased from graft stenosis diar</li> <li>Increase in luminal diameter in CoA pa</li> <li>CP Stent (Covered) frame is made from additional gold soldering. The strut thi relatively atraumatic.</li> <li>CP Stent (Covered) was hand-crimped performed with a 10 ml syringe on the inflation pressures to 4–6 atmosphere</li> <li>In this retrospective study, there are n</li> </ul> </li> </ul>	i) mm Hg, P<0.001. g and pulmonary valve delivery (RVOT Group): ' in a second procedure. neter of 13 (5-22) mm to 22 (16-26) mm at pre-revalvulation tients. n 90% platinum and 10% iridium 0.013" wire, welded in a zi ckness is slightly larger than most other stents, but makes to on a balloon-in-balloon (BIB, Numed). Hand-inflation of the inner balloon and 20 ml syringe on the outer balloon, auto	ig patte the ster e balloo omatica rently re	ern with nt edges on was Ily limit elated t	ing to the
	Appraisal					
	Level of Evidence	Study Method/Design	Our attack Accelted	0.4-	rd LOE	2011
		, , , ,	Question Applied			-
		Retrospective and observational study.	Question Applied           To investigate reduction in aortic wall rupture and dissection, as well as aneurysms by implanting covered stents.		2 3	<b>4</b> 5
		Retrospective and observational study.	To investigate reduction in aortic wall rupture and dissection, as well as aneurysms by implanting		2 3	<b>4</b> 5
	Suitability	Retrospective and observational study.	To investigate reduction in aortic wall rupture and dissection, as well as aneurysms by implanting	1	2 3 Gradin	4 5
7. Alcibar et al. (2013)		Retrospective and observational study.	To investigate reduction in aortic wall rupture and dissection, as well as aneurysms by implanting covered stents.		2 3	<b>4</b> 5
	Suitability	Retrospective and observational study.         Relevant Data         -       Covered CP Stent         -       BIB or Z-Med balloons (NuMED) – 9 of	To investigate reduction in aortic wall rupture and dissection, as well as aneurysms by implanting covered stents.	1	2 3 Gradin	4 5
(2013)	Suitability Device	Retrospective and observational study.         Relevant Data         -       Covered CP Stent         -       BIB or Z-Med balloons (NuMED) – 9 of         -       Hand crimped         -       CoA and re-coarctation	To investigate reduction in aortic wall rupture and dissection, as well as aneurysms by implanting covered stents. the 17 patients had BIB ation (2 adolescents and 15 adults treated between	1 2 D1	2 3 Gradin D2	4 5
(2013) Contribution S&P x	Suitability Device Application	Retrospective and observational study.         Retrospective and observational study.         Relevant Data         - Covered CP Stent         - BIB or Z-Med balloons (NuMED) – 9 of         - Hand crimped         - CoA and re-coarctation         - Patients treated for CoA and re-coarct         November 2005 and January 2012).         - Sampling: n=17 (11 native CoA and 6 r         - Mean age: 35 (range 14-65) years	To investigate reduction in aortic wall rupture and dissection, as well as aneurysms by implanting covered stents. the 17 patients had BIB ation (2 adolescents and 15 adults treated between	1 2 D1 A1	2 3 Gradin D2 A2	4 5 g D3 A3
(2013) Contribution S&P x	Suitability Device Application Patient	Retrospective and observational study.         Retrospective and observational study.         Relevant Data         - Covered CP Stent         - BIB or Z-Med balloons (NuMED) – 9 of         - Hand crimped         - CoA and re-coarctation         - Patients treated for CoA and re-coarct         November 2005 and January 2012).         - Sampling: n=17 (11 native CoA and 6 r         - Mean age: 35 (range 14-65) years         - Sex: 4 M; 13 F	To investigate reduction in aortic wall rupture and dissection, as well as aneurysms by implanting covered stents. the 17 patients had BIB ation (2 adolescents and 15 adults treated between	1 7 D1 A1 P1	Gradin D2 A2 P2	4 5 D3 A3 P3
(2013) Contribution S&P x	Suitability Device Application Patient	Retrospective and observational study.         Retrospective and observational study.         Relevant Data         - Covered CP Stent         - BIB or Z-Med balloons (NuMED) – 9 of         - Hand crimped         - CoA and re-coarctation         - Patients treated for CoA and re-coarct         November 2005 and January 2012).         - Sampling: n=17 (11 native CoA and 6 r         - Mean age: 35 (range 14-65) years         - Sex: 4 M; 13 F	To investigate reduction in aortic wall rupture and dissection, as well as aneurysms by implanting covered stents. the 17 patients had BIB ation (2 adolescents and 15 adults treated between e-coarctation)	1 7 D1 A1 P1	Gradin D2 A2 P2 R2	4 5 g D3 A3 P3 R3



	- Reduction in lumen diameter				
	- Reduction of hypertensive medications at follow-up				
E a Ula su sua	- Adverse effects				N
Follow-up	- 2.5 years			Yes 1	No 2
Statistical analysis	- Significance was considered as P<0.05.			Yes 1	No 2
Clinical significance	<ul> <li>CP Stents (Covered) are effective in treating CoA and are the treatment of choice in patients with complex operating room as a rescue device when implanting a</li> </ul>	anatomy, and must be ava		Yes 1	No
	· · · · · · · · · · · · · · · · · · ·	Data Contribution Gra	ade (Range 4-8)		4
Overall S&P Appraisal, Di S&P Grade		on and Weighting (select)	Accepted and	Pivotal 9-	12
(Range 9-25)	Data Contribution (4) = 12		Accepted but		
(Range 5 25)			Excluded, 22-2		, 13 21
Relevant S&P Results					
Safety data Performance data	<ul> <li>One death: patient died two days post-op due to mas increase in blood pressure and an existing aneurysm.</li> <li>No local complications occurred, except one hemator</li> <li>No patient had any complication at the iliac-femoral I</li> <li>Blood pressure gradient: Reduced from 40 to 2 mmH</li> </ul>	ma that resolved spontaneo level that required stenting.	ously.		
	<ul> <li>Lumen diameter: Increased from 4 to 15 mm (P&lt;0.00</li> <li>At follow-up (2.5 years):         <ul> <li>All good initial outcome persisted without any sig</li> <li>13/17 patients underwent imaging study; no and observed.</li> <li>Medication for hypertension was reduced in 5 page</li> </ul> </li> </ul>	1) gns of re-obstruction. eurysms, dissections, and/o			e
Benefits/claims data	<ul> <li>Increased in luminal diameter</li> <li>Decreased in antihypertensive medication use</li> </ul>				
Strengths	<ul> <li>Having observed the case of aortic rupture, and with had CoA and re-coarctation since their youth, the aut York, United States) ePTFE CP Stent (Covered). This st wall when expanded.</li> </ul>	hors decided to electively in	mplant a NuMED	(Hopkinto	n, New
Weaknesses/ Potential bias	<ul> <li>Retrospective and observational study with no contropatients underwent clinical follow-up, this did not inc determine with certainty the incidence of potential and</li> </ul>	lude an imaging study in all			



	Safety & Performance Appraisal						
	Level of Evidence	Study Method/Design	Question Applied		Oxfo	rd LOE	2011
		Single arm interventional study.	To evaluate the use of CP Stent (Cover	ed) as the	1 2	2 3	4
		,	primary modality in the treatment for				
	Suitability	Relevant Data				Gradi	ng
	Device	- CP Stent (Covered and Bare) 25 cov	vered stents and 2 bare stents in 25 patients (	one patient	D1	D2	D
		had 3 stents (2 bare, one covered)	for native CoA with aortic arch hypoplasia (co	ombination			
		covered + stents = a new approach	)				
		- The covered CP stent was hand-cri	mped down onto BIB				
	Application	- Native CoA			A1	A2	A
	Patient	- Patients with native CoA without p	revious treatment		P1	P2	PB
		- Sampling: n=25					
		- Mean age: 22.5 (range 14-46) year	5				
		- Sex: 16 M; 9 F					
	Report	- High quality.			R1	R2	R
Chang et al.			Suitability Gra	de (Range 4-12)		4	
(2012)							
	Data Contribution	Relevant Data				Gradi	ng
Contribution	Outcomes/Endpoints	- Decrease in systolic gradient			Yes 1		No 2
S&P x		- Increase in stenotic segment diam	eter				
SOA -	Follow-up	- 32 (7-72) months			Yes 1		No 2
	Statistical analysis	<ul> <li>P&lt;0.05 was set as statistically signi</li> </ul>			Yes 1		No 2
	Clinical significance		as the primary modality is safe and effective	in the	Yes 1		No 2
		treatment for native CoA in adoles					
			in adolescents and adults acquired excellent r				
			ic gradient across CoA, successful relief of ana	atomic			
		stenosis, and reduction of systemi		the fellow			
			encountered during the procedure or during	the follow-up			
		period of up to 72 months.	Data Contribution Gra	ada (Banga 1 9)		4	
			Data contribution dia	aue (Ralige 4-6)		4	
	Overall S&P Appraisal, Dis	position and Weighting					
	S&P Grade	LOE (4) + Suitability (4) +	Disposition and Weighting (select)	Accepted and	Divotal	0 1 2	
	(Range 9-25)	Data Contribution $(4) = 12$	Disposition and weighting (select)	Accepted and			
		Data Contribution (4) – $12$				Jiai, 1	5-21
	(Ralige 9-25)						
				Excluded, 22-2	.5		
	Relevant S&P Results Safety data	- No acute complications were obser	ved	Excluded, 22-2	.5		



	Performance data Benefits/claims data	<ul> <li>In the patient with the implantation encountered, the left subclavian ar left arm ischemia was not detected</li> <li>Peak systolic gradient across the left</li> <li>Decreased from median 67.5 m</li> <li>Stenotic segment diameter</li> <li>Increased from median 5.0mm</li> <li>At follow-up (up to 72 months):</li> </ul>		hrombo	sis, and t	:he
	Strengths	- BIB offered precise and safe contro	l over the stent implantation without any stent migration nary treatment modality may reduce the risk of significant co	omplicat	ions rela	ted
	Weaknesses/	- Conflict of interest: not reported.				
	Potential bias					
	Cofoty & Dorformanco (fo	r cafatu anlu)				
9. Erdem et al.	Safety & Performance (fo Appraisal Level of Evidence	r safety only) Study Method/Design Single arm interventional study.	Question AppliedTo present author's institutional experience of endovascular CP Stent implantation in children and adults with native and recurrent CoA.		rd LOE 2 2 3	<b>4</b> 5
9. Erdem et al. (2011)	Appraisal Level of Evidence	Study Method/Design Single arm interventional study.	To present author's institutional experience of endovascular CP Stent implantation in children and adults with native and		2 3	<b>4</b> 5
(2011) Contribution S&P X (S	Appraisal	Study Method/Design         Single arm interventional study.         Relevant Data         -       CP Stent (16 Covered or 31 Bare) –	To present author's institutional experience of endovascular CP Stent implantation in children and adults with native and recurrent CoA. n=47 ingle balloon catheter (n=18) (not subject device), Z-med			<b>4</b> 5
(2011) Contribution S&P X (S only)	Appraisal Level of Evidence Suitability	Study Method/Design         Single arm interventional study.         Relevant Data         -       CP Stent (16 Covered or 31 Bare) –         -       BIB (n=29) (not subject device) or state	To present author's institutional experience of endovascular CP Stent implantation in children and adults with native and recurrent CoA.	1	2 3 Gradinį	<b>4</b> 5
(2011) Contribution S&P X (S	Appraisal Level of Evidence Suitability Device	Study Method/Design         Single arm interventional study.         Relevant Data         -       CP Stent (16 Covered or 31 Bare) –         -       BIB (n=29) (not subject device) or s (not subject device); manually crim         -       Patients with native or recurrent C	To present author's institutional experience of endovascular CP Stent implantation in children and adults with native and recurrent CoA. n=47 ingle balloon catheter (n=18) (not subject device), Z-med ped oA grecurrent CoA and/or aneurysm developed after either sup 2) ered or Bare)	1 : D1	2 3 Gradina D2	4 5
(2011) Contribution S&P X (S only)	Appraisal         Level of Evidence         Suitability         Device         Application	Study Method/Design         Single arm interventional study.         Relevant Data         -       CP Stent (16 Covered or 31 Bare) –         -       BIB (n=29) (not subject device) or ss (not subject device); manually crim         -       Patients with native or recurrent C         -       Patients with native CoA (Group 1) surgery or balloon angioplasty (Group 1)         -       Sampling: n=45 (47 CP Stents, Cover -         -       Median age: 11 (range: 5-33) years	To present author's institutional experience of endovascular CP Stent implantation in children and adults with native and recurrent CoA. n=47 ingle balloon catheter (n=18) (not subject device), Z-med ped oA grecurrent CoA and/or aneurysm developed after either sup 2) ered or Bare)	1 : D1 A1	Grading Grading D2 A2	4 5 3 D3 A3



Data Contribution	Relevant Data			Gr	ading	
Outcomes/Endpoints	<ul> <li>Decrease in invasive and echocardi</li> <li>Increase in lesion diameter</li> <li>Adverse effects</li> </ul>	ographic gradients		Yes 1	No	
Follow-up	- 12.1±7.1 months; median 11 month	h (range 2-29)		Yes 1	No	
Statistical analysis	- A p value <0.05 was considered sta			Yes 1	No	
Clinical significance	<ul> <li>Early and short- term follow-up res effective in reducing coarctation gr recurrent CoA.</li> <li>Some serious complications do occ</li> </ul>	Some serious complications do occur and hypertension remains in some patients. Aortic disruption and stent displacement are potentially catastrophic complications of stenting			No	
		<ul> <li>Aortic disruption and stent displacement are potentially catastrophic complications of stenting but implanting a second covered stent can seal the ruptured wall and parking in a safe area or</li> </ul>				
		ent can seal the ruptured wall and parking in ried by half-inflated balloon could solve the p				
		Data Contribution Gra			4	
			. 0 -/			
S&P Grade	LOE (4) + Suitability (5) +	Disposition and Weighting (select)	Accepted and			
(Range 9-25)	Data Contribution (4) = 13		Accepted but Excluded, 22-2	<b>but not Pivotal, 13</b> - 22-25		
				-		
Relevant S&P Results				-		
Safety data	<ul> <li>covered stent</li> <li>One stent was displaced before</li> <li>balloon and long sheath, and response to the sterial complications</li> <li>No difficulty in catheter manipulation</li> <li>None of the patients required intered ay except the patient with aortic response to the steries of the steries</li></ul>	cessfully managed immediately in the same some it was completely opened. It was carried epositioned into the correct place. On. Insive care following the procedure, and all was upture and after stenting with covered stent	session with impl ed with support were discharged	antation o of partial home the	lly inflat	
Safety data Benefits/claims data	<ul> <li>Two immediate complications relations of the second stent</li> <li>One an acute wall rupture, successed stent</li> <li>One stent was displaced before balloon and long sheath, and response of the second stent relations</li> <li>No femoral arterial complications</li> <li>No difficulty in catheter manipulation</li> <li>None of the patients required interday except the patient with aortic relations</li> <li>Increase in luminal/lesion diameter</li> </ul>	cessfully managed immediately in the same some it was completely opened. It was carried epositioned into the correct place. On. Insive care following the procedure, and all supplure and after stenting with covered stent	session with impl ed with support were discharged	antation o of partial home the	lly infla	
Safety data	<ul> <li>Two immediate complications relations of the stent</li> <li>One an acute wall rupture, successed stent</li> <li>One stent was displaced before balloon and long sheath, and response of the sterial complications</li> <li>No difficulty in catheter manipulation</li> <li>None of the patients required intered day except the patient with aortic response of the steries of</li></ul>	cessfully managed immediately in the same some it was completely opened. It was carried epositioned into the correct place. On. Insive care following the procedure, and all was upture and after stenting with covered stent	session with impl ed with support were discharged this patient was	antation o of partial home the followed t	ly infla e follow wo day:	
Safety data Benefits/claims data	<ul> <li>Two immediate complications relations of the stent with a covered stent</li> <li>One stent was displaced before balloon and long sheath, and response of the patients required interest day except the patient with a ortic response of the most complication diameter.</li> <li>Increase in luminal/lesion diameter.</li> <li>CP stent is the one of the most complications that sexcellent radial strengs.</li> <li>Some limitations have to be noted a streng streng streng strengs.</li> </ul>	cessfully managed immediately in the same some it was completely opened. It was carried epositioned into the correct place.	session with impl ed with support were discharged this patient was liant visibility on	antation o of partial home the followed t fluoroscop	ly infla e follow wo day by.	



			ood pressure monitoring before stenting wa eurysm was done in limited number of pati	•		v patier	nts.
	Safety & Performance						
	Appraisal	Cturdue Mathead /Design	Question Applied		Oxford LOE 20		
	Level of Evidence	Study Method/Design Prospective single arm interventional study.	Question Applied           To evaluate the management of aneur associated with CoA by covered stent			2 <b>3</b>	4 5
	Suitability	Relevant Data					ng
	Device				D1	D2	D3
	Application	- Patients with native CoA associated w			A1	A2	A3
10. Butera et al.	Patient	<ul> <li>Patients with CoA associated with aou</li> <li>Sampling: n=11 (3 native CoA, 3 with angioplasty, and 2 with previous bare</li> <li>Median age: 13 (range: 6-66) years</li> <li>Sex: Not reported</li> </ul>	previous surgical repair, 3 with previous ba	illoon	P1	P2	Р3
(2011)	Report	- High quality.			R1	R2	R3
Contribution			Suitability Grad	de (Range 4-12)		6	
S&P x	Data Contribution	Relevant Data			Grading		
SOA -	Outcomes/Endpoints	<ul> <li>Systolic pressure gradient reduction</li> <li>Increase in aortic diameter</li> <li>Adverse effects</li> </ul>			Yes 1		No 2
	Follow-up	- Median follow-up 50 (16-61) months			Yes 1	L	No 2
	Statistical analysis	- P-value less than 0.05 was considered			Yes 1		No 2
	Clinical significance	treatment of CoA associated with ao	onsidered the treatment of choice for nati		Yes 1	L	No 2
			Data Contribution Gra	ade (Range 4-8)		4	
	Overall S&P Appraisal, Dis	sposition and Weighting					
	S&P Grade (Range 9-25)	LOE (3) + Suitability (6) + Data Contribution (4) = 13	Disposition and Weighting (select)	Accepted and Accepted but Excluded, 22-		.3-21	

	Relevant S&P Results							
	Safety data	- No early complications observed.						
	Performance data	- Successful device deployment: Ach						
		<ul> <li>Successful relief of stenoses and co</li> </ul>						
			median 6 (0.5 – 11) to 12 (10-22) mm, P<0.001					
			: four patients developed systemic hypertension (one intraste					
			al growth, three showed restenosis secondary to somatic grow	rth). Re-	dilatatio	on		
	Deve fits / states a data		ed without complication in all cases.					
	Benefits/claims data	- Increase in luminal diameter						
		- Reduce systolic pressure gradient	(notionts associated with partic wall anounism)					
	Strengths		patients associated with aortic wall aneurysm) ed with an alloy of 90% platinum and 10% iridium. Theoretical	h, this a	ombing	tion		
	Strengtris		adial strength, which is enhanced by being designed in a "zig"					
			ng the risk of balloon rupture or injury to the vessel wall and, i					
			e radio-opaque. Furthermore, the e-PTFE protects the stenotic					
		segment.		e and an				
	Weakposses/							
		Weaknesses/ - No conflict of interest reported.						
	Potential bias Safety & Performance	- No connict of interest reported.						
	Potential bias Safety & Performance Appraisal		Outstien Apriliad			2011		
	Potential bias Safety & Performance	Study Method/Design	Question Applied		rd LOE			
	Potential bias Safety & Performance Appraisal		To determine the safety and efficacy of the CP Stent		rd LOE 2 3			
	Potential bias Safety & Performance Appraisal	Study Method/Design						
	Potential bias Safety & Performance Appraisal	Study Method/Design	To determine the safety and efficacy of the CP Stent			4		
	Potential bias         Safety & Performance         Appraisal         Level of Evidence	Study Method/Design Single arm interventional study. Relevant Data	To determine the safety and efficacy of the CP Stent		2 3	<b>4</b>		
. Tanous et al. (2010)	Potential bias         Safety & Performance         Appraisal         Level of Evidence         Suitability	Study Method/Design Single arm interventional study. Relevant Data	To determine the safety and efficacy of the CP Stent (Covered) in adults with CoA.		2 3 Gradin	4 1		
(2010)	Potential bias         Safety & Performance         Appraisal         Level of Evidence         Suitability         Device	Study Method/Design Single arm interventional study. Relevant Data - Covered CP Stent hand-crimped o	To determine the safety and efficacy of the CP Stent (Covered) in adults with CoA.	1 2 D1	2 3 Gradin D2	4 ng D3 A3		
(2010)	Potential bias         Safety & Performance         Appraisal         Level of Evidence         Suitability         Device         Application	Study Method/Design         Single arm interventional study.         Relevant Data         -       Covered CP Stent hand-crimped o         -       Native CoA (n=14) and previous tree	To determine the safety and efficacy of the CP Stent (Covered) in adults with CoA.	1 2 D1 A1	2 3 Gradin D2 A2	4 D3 A3		
(2010) Contribution	Potential bias         Safety & Performance         Appraisal         Level of Evidence         Suitability         Device         Application	Study Method/Design         Single arm interventional study.         Relevant Data         -       Covered CP Stent hand-crimped o         -       Native CoA (n=14) and previous tradements with native CoA and CoA with native CoA	To determine the safety and efficacy of the CP Stent (Covered) in adults with CoA.	1 2 D1 A1	2 3 Gradin D2 A2	4 D3 A3		
(2010) Contribution &P x	Potential bias         Safety & Performance         Appraisal         Level of Evidence         Suitability         Device         Application	Study Method/Design         Single arm interventional study.         Relevant Data         -       Covered CP Stent hand-crimped o         -       Native CoA (n=14) and previous troped on the study of the state of th	To determine the safety and efficacy of the CP Stent (Covered) in adults with CoA.	1 2 D1 A1	Gradin D2 A2 P2	4 D3 A3 P3		
(2010) Contribution &P x	Potential bias         Safety & Performance         Appraisal         Level of Evidence         Suitability         Device         Application	Study Method/Design         Single arm interventional study.         Relevant Data         -       Covered CP Stent hand-crimped o         -       Native CoA (n=14) and previous trop         -       Patients with native CoA and CoA m         -       Sampling: 14 native CoA; 8 CoA (w         -       Mean age: 39±14 (range 19 to 67)	To determine the safety and efficacy of the CP Stent (Covered) in adults with CoA.	1 2 D1 A1	2 3 Gradin D2 A2	4 D3 A3 P3		
(2010) Contribution	Potential bias         Safety & Performance         Appraisal         Level of Evidence         Suitability         Device         Application         Patient	Study Method/Design         Single arm interventional study.         Relevant Data         -       Covered CP Stent hand-crimped o         -       Native CoA (n=14) and previous tradements         -       Patients with native CoA and CoA mean age: 39±14 (range 19 to 67)         -       Sex: 11 M; 11 F	To determine the safety and efficacy of the CP Stent (Covered) in adults with CoA.	1 2 D1 A1 P1	Gradin D2 A2 P2	4 2 09 03 A3 P3		
(2010) Contribution	Potential bias         Safety & Performance         Appraisal         Level of Evidence         Suitability         Device         Application         Patient         Report	Study Method/Design         Single arm interventional study.         Relevant Data         -       Covered CP Stent hand-crimped o         -       Native CoA (n=14) and previous trophone         -       Patients with native CoA and CoA or CoA; 8 CoA (w)         -       Sampling: 14 native CoA; 8 CoA (w)         -       Sex: 11 M; 11 F         -       High quality.	To determine the safety and efficacy of the CP Stent (Covered) in adults with CoA.	1 2 D1 A1 P1	Gradin Gradin D2 A2 P2 R2 5	4 2 08 03 03 A3 P3 P3 R3		
Contribution S&P x	Potential bias         Safety & Performance         Appraisal         Level of Evidence         Suitability         Device         Application         Patient	Study Method/Design         Single arm interventional study.         Relevant Data         -       Covered CP Stent hand-crimped o         -       Native CoA (n=14) and previous tradements         -       Patients with native CoA and CoA mean age: 39±14 (range 19 to 67)         -       Sex: 11 M; 11 F	To determine the safety and efficacy of the CP Stent (Covered) in adults with CoA. an Z-Med II (NuMED) or Covered Mounted CP Stent eatment (n=8) with previous treatment ith previous treatment) years Suitability Grade (Range 4-12)	1 2 D1 A1 P1	Gradin Gradin D2 A2 P2 R2 S Gradin	4 9 08 03 03 03 03 03 03 03 03 03 03 03 03 03		



	Follow-up	- 12 (9-15) months			Yes 1		No 2
	Statistical analysis	<ul> <li>A P-value &lt;0.05 was considered sig</li> </ul>	nificant.		Yes 1		No 2
	Clinical significance	- Covered stents are safe, durable, a	nd efficacious in the management of CoA.		Yes 1		No 2
			Data Contribution Gra	ade (Range 4-8)		4	
	Overall S&P Appraisal, Di	isposition and Weighting					
	S&P Grade	LOE (4) + Suitability (5) +	Disposition and Weighting (select)	Accepted and	Pivotal	9-12	
	(Range 9-25)	Data Contribution (4) = 13		Accepted but Excluded, 22-2	not Pivo		8-21
	Relevant S&P Results						
	Safety data	<ul> <li>One pseudoaneurysm. Patient was Note: this problem may have been used the problem did not reoccur.</li> </ul>	treated successfully. caused because the stent was hand crimped. \	When pre-moun	ted ster	nts wer	e
	Performance data	<ul> <li>Reduction in peak systolic gradient intervention and 6 ± 9 mmHg at fol</li> </ul>	across coarctation site: From average 29 ± 17 low up, P<0.001	to 3 ± 5 mmHg i	mmedia	itely po	ost
	Benefits/claims data	- Reduction in peak systolic gradient					
	Strengths	- N/A					
	Weaknesses/	- This review is limited by the small s	ample size and lack of a randomized comparis	on group			
		<ul> <li>This study was not intended to demonstrate the efficacy of percutaneous therapy, or the superiority of stents, but rather to document a single-center experience as an alternative and safe treatment option i</li> </ul>					
	Potential bias	- This study was not intended to dem	nonstrate the efficacy of percutaneous therapy ngle-center experience as an alternative and s	y, or the superior			
		- This study was not intended to dem stents, but rather to document a sin spectrum of patients with aortic co	nonstrate the efficacy of percutaneous therapy ngle-center experience as an alternative and s	y, or the superior			
	Potential bias Safety & Performance (for	- This study was not intended to dem stents, but rather to document a sin spectrum of patients with aortic co	nonstrate the efficacy of percutaneous therapy ngle-center experience as an alternative and s	y, or the superior	otion in		d
	Potential bias           Safety & Performance (for Appraisal	<ul> <li>This study was not intended to dem stents, but rather to document a sin spectrum of patients with aortic co</li> </ul>	nonstrate the efficacy of percutaneous therapy ngle-center experience as an alternative and s arctation.	y, or the superior afe treatment op	otion in	a broa	d
. Moltzer et al. (2010)	Potential bias Safety & Performance (for Appraisal Level of Evidence	This study was not intended to dem stents, but rather to document a sin spectrum of patients with aortic co or safety only)     Study Method/Design     Prospective observational study.	A provide the efficacy of percutaneous therapy ngle-center experience as an alternative and starctation.	y, or the superior afe treatment op	Oxfor 1 2	a broa	d 2011 4
(2010)	Potential bias           Safety & Performance (for Appraisal	<ul> <li>This study was not intended to dem stents, but rather to document a sin spectrum of patients with aortic co</li> <li>br safety only)</li> <li>Study Method/Design         Prospective observational study.     </li> <li>Relevant Data         CP Stent (Bare and Covered) – 6 of     </li> </ul>	An output to the efficacy of percutaneous therapy of the percutaneous	y, or the superior afe treatment op	Oxfor 1 2	a broa	2011 4
(2010) Contribution	Potential bias           Safety & Performance (for           Appraisal           Level of Evidence           Suitability           Device	<ul> <li>This study was not intended to dem stents, but rather to document a sin spectrum of patients with aortic co</li> <li>or safety only)</li> <li>Study Method/Design         Prospective observational study.     </li> <li>Relevant Data         <ul> <li>CP Stent (Bare and Covered) – 6 of - BIB (manually crimped)</li> </ul> </li> </ul>	An output to the efficacy of percutaneous therapy of the percutaneous	y, or the superior afe treatment op	Oxfor 1 2 D1	d LOE Gradin D2	d 2011 4 D: D:
	Potential bias           Safety & Performance (for           Appraisal           Level of Evidence           Suitability	<ul> <li>This study was not intended to dem stents, but rather to document a sin spectrum of patients with aortic co</li> <li>br safety only)</li> <li>Study Method/Design         <ul> <li>Prospective observational study.</li> </ul> </li> <li>Relevant Data         <ul> <li>CP Stent (Bare and Covered) – 6 of - BIB (manually crimped)</li> <li>Native CoA and re-coarctation</li> <li>Patients with native CoA and re-coarctation</li> <li>Sampling: n=24</li> <li>Mean age: 36 (18-60) years</li> </ul> </li> </ul>	Ourstrate the efficacy of percutaneous therapyingle-center experience as an alternative and starctation.         Question Applied         To evaluate the intermediate-term out implantation for CoA in adults.         the 24 patients had covered stents	y, or the superior afe treatment op	Oxfor 1 2	a broa	d 2011 4 Dg A3
(2010) Contribution 5&P X (S only)	Potential bias           Safety & Performance (for           Appraisal           Level of Evidence           Suitability           Device           Application	<ul> <li>This study was not intended to dem stents, but rather to document a sin spectrum of patients with aortic co</li> <li>Dr safety only)</li> <li>Study Method/Design         <ul> <li>Prospective observational study.</li> </ul> </li> <li>Relevant Data         <ul> <li>CP Stent (Bare and Covered) – 6 of BIB (manually crimped)</li> <li>Native CoA and re-coarctation</li> <li>Patients with native CoA and re-coarctation</li> <li>Sampling: n=24</li> <li>Mean age: 36 (18-60) years</li> <li>Sex: 12 M; 12 F</li> </ul> </li> </ul>	Ourstrate the efficacy of percutaneous therapyingle-center experience as an alternative and starctation.         Question Applied         To evaluate the intermediate-term out implantation for CoA in adults.         the 24 patients had covered stents	y, or the superior afe treatment op	Oxfor 1 2 D1 A1 P1	d LOE 3 Gradin D2 A2 P2	d 20111 4 D3 P3
(2010) Contribution 5&P X (S only)	Potential bias           Safety & Performance (for           Appraisal           Level of Evidence           Suitability           Device           Application	<ul> <li>This study was not intended to dem stents, but rather to document a sin spectrum of patients with aortic co</li> <li>br safety only)</li> <li>Study Method/Design         <ul> <li>Prospective observational study.</li> </ul> </li> <li>Relevant Data         <ul> <li>CP Stent (Bare and Covered) – 6 of - BIB (manually crimped)</li> <li>Native CoA and re-coarctation</li> <li>Patients with native CoA and re-coarctation</li> <li>Sampling: n=24</li> <li>Mean age: 36 (18-60) years</li> </ul> </li> </ul>	Ourstrate the efficacy of percutaneous therapyingle-center experience as an alternative and starctation.         Question Applied         To evaluate the intermediate-term out implantation for CoA in adults.         the 24 patients had covered stents	y, or the superior afe treatment op come of stent	Oxfor 1 2 D1 A1	a broa	d 2011 4



	Data Contribution	Relevant Data				Grading
	Outcomes/Endpoints	- Decrease in systolic gradient			Yes 1	No 2
		- Increase in minimum aortic diameter				
		- Adverse effects				
	Follow-up	- 24 hours post intervention and 33 (8-			Yes 1	No 2
	Statistical analysis		d a p-value <0.05 was considered statistica		Yes 1	No 2
	Clinical significance		blood pressure gradient decrease and inc		Yes 1	No 2
			ions do occur and hypertension remains ir	n the majority		
		of patients.				
			Data Contribution Gra	ade (Range 4-8)		4
	Overall S&P Appraisal, Dis	sposition and Weighting				
	S&P Grade	LOE (3) + Suitability (4) +	Disposition and Weighting (select)	Accepted an	<b>Pivotal</b>	9-12
	(Range 9-25)	Data Contribution (4) = 11		Accepted but Excluded, 22-2		tal, 13-21
	Relevant S&P Results	•				
	Safety data	<ul> <li>One death due to aorta ruptured.</li> <li>Two groin hematoma post-op.</li> </ul>				
	Benefits/claims data	Reduced in systolic gradient     Increased in minimum aortic diameter				
	Strengths	- N/A				
	Weaknesses/ Potential bias	This was a single-center report and part 24-hour blood pressure monitoring be	undergone stent implantation since the au tients were not compared with surgery or fore stenting was not performed in the ma toring is therefore difficult to translate in t	balloon angiopla ijority of the pati	sty alone ients. Po:	e. Finally, st-stent 24
	Safety & Performance (fo	r safety only)				
	Appraisal	Study Method/Design	Question Applied		Ovfor	d LOE 2011
L3. Agnoletti et al. (2009)		Two arms comparative interventional study.	To compare the CP Stent and the Palm treatment of native and postoperative		1 2	
Contribution			patients.			
Contribution S&P X (S	Suitability	Relevant Data				Grading
SOA -	Device	- CP Stent (Bare & Covered), crimped or     - 96 CP Stents (34 covered), 77 Palma St			D1	D2 D3
		- Palmaz stent, crimped on BIB and simp				
	Application	<ul> <li>Patients with CHD (including CoA/re-content)</li> </ul>			A1	<b>A2</b> A3
	Patient	- Patients with CHD (including CoA/re-c	oarctation, RVOT and other CHD condition	s such as	P1	P2 P3



Report	<ul> <li>transposition of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, septal defect, single version of the great arteries, ventricular septal defect, single version of the great arteries, ventricular septal defect, septal defect</li></ul>	ons)	R1	R2	R3
	Su	iitability Grade (Range 4-12)		6	
Data Contribution	Relevant Data			Grad	ing
Outcomes/Endpoints	<ul> <li>Blood pressure gradient reduction</li> <li>Vessel diameter reduction</li> <li>Adverse effects</li> </ul>				No 2
Follow-up	- Not reported.		Yes 1		No 2
Statistical analysis	- A P-value less than 0.05 was considered statistically significant for st	tent group comparison.	Yes 1		No 2
Clinical significance	<ul> <li>The use of the CP Stents to treat stenotic lesions of CHD is effective and relatively safe. The overall efficacy of CP Stents for the treatment of stenotic lesions is superior to that of the Palmaz stent.</li> <li>CP Stents' overall safety is higher than that of the Palmaz stent; but Palmaz stents have a lower profile when inserted.</li> </ul>				No 2
	Data Con	tribution Grade (Range 4-8)		5	
Overall S&P Appraisal, Dis					
S&P Grade (Range 9-25)	LOE (3) + Suitability (6) + Disposition and Weightin Data Contribution (5) = 14	ng (select) Accepted and Accepted but Excluded, 22-	t not Piv		13-21
		·			
elevant S&P Results	<ul> <li>Stent-related complications:</li> <li>CP Stents (n=89 patients): 4 mild, 1 moderate, 1 severe.</li> </ul>				



	Benefits/claims data Strengths Weaknesses/ Potential bias	<ul> <li>Decreased in blood pressure gradient.</li> <li>Increased in vessel diameter.</li> <li>Efficacy of CP Stents was similar to that of that of Palmaz for the stenting of aorta,</li> <li>Study presented retrospective results ob</li> <li>CP stents were used for patients weighing</li> </ul>	alloons, and 1 on pre-mounted stent). of Palmaz stent for stenting of the right ventricular outflo but the difference was not statistically.	rent cor	ncerni	ng age	,
	Safety & Performance	<ul> <li>Subgroup analyses were not performed.</li> </ul>		compi		13.	
	Appraisal			· · ·			
14. Bruckheimer et al. (2009) Contribution S&P x SOA -	Level of Evidence Suitability Device Application Patient	Study Method/Design         Prospective interventional single arm study to evaluate placement of CP Stent (Covered) with smallest possible balloon sizes, then dilation with larger diameter balloons until pressure gradient was <20 mmHg and the stent was opposed to the aortic wall.         Relevant Data         -       Covered CP Stents         -       Manually crimped on a balloon         -       Native CoA         -       Patients with native CoA         -       Sampling: n=22	Question Applied To report on the early results of treatment of native CoA by implantation and serial dilations of covered stents.		rd LOI           2         3           Grad         D2           A2         P2		3
	Report	<ul> <li>Mean age: 15.5 (7.8 – 38.6) years</li> <li>Sex: 14 M; 8 F</li> <li>High quality.</li> </ul>		R1	R2	R3	5
			Suitability Grade (Range 4-12)		4		
				1			
	Data Contribution	Relevant Data			Grad	ing	
	Outcomes/Endpoints	<ul> <li>Increase of coarctation diameter</li> <li>Reduction of peak pressure gradient</li> <li>Adverse effects</li> </ul>		Yes 1	1	No 2	
	Follow-up	<ul> <li>Median 18.5 (1.6-31.4) months</li> </ul>		Yes 1		No 2	



	Statistical analysis	<ul> <li>P-values reported.</li> </ul>			Yes 1	I	No 2
	Clinical significance	- Serial dilation of CP Stents (Covere	ed) is feasible, safe and an effective percutane	ous method	Yes 1		No 2
		for the treatment of native CoA.					
			Data Contribution Gra	ade (Range 4-8)		4	
	Overall S&P Appraisal, Di	sposition and Weighting					
	S&P Grade	LOE (3) + Suitability (4) +	Disposition and Weighting (select)	Accepted and	Pivotal	9-12	
	(Range 9-25)	Data Contribution (4) = 11		Accepted but Excluded, 22-2	not Pivo		-21
	Relevant S&P Results						
	Safety data	<ul> <li>One small tear at the distal stent ed</li> <li>One femoral pseudoaneurysm white</li> </ul>					
	Performance data	- Increase of coarctation diameter:	ention to 12.6 ±1.9 mm post-intervention, P=0	).001			
		<ul> <li>Reduction of peak pressure gradier</li> <li>From 29.4 ± 8.5 to 6.7 ± 5.7 m</li> </ul>	nt:				
	Benefits/claims data	<ul> <li>Increase of coarctation diameter</li> <li>Reduction of peak pressure gradier</li> </ul>					
		neddetion of peak pressure gradier					
	Strengths	- N/A					
	Strengths Weaknesses/ Potential bias	<ul> <li>N/A</li> <li>No conflict of interest reported.</li> </ul>					
	Weaknesses/ Potential bias Safety & Performance						
	Weaknesses/ Potential bias	- No conflict of interest reported.	Question Applied		Oxfor	d LOE	2011
	Weaknesses/ Potential bias Safety & Performance Appraisal		Question Applied To evaluate the use of Covered CP Ster of CoA.	nts in treatment	Oxfor 1 2	d LOE	2011 4
	Weaknesses/ Potential bias Safety & Performance Appraisal Level of Evidence	No conflict of interest reported.     Study Method/Design     Single arm interventional study.	To evaluate the use of Covered CP Ster	nts in treatment	1 2	3	4
5. Tzifa et al. (2006)	Weaknesses/ Potential bias Safety & Performance Appraisal	- No conflict of interest reported.      Study Method/Design     Single arm interventional study.      Relevant Data     - Covered CP Stent	To evaluate the use of Covered CP Ster	nts in treatment	1 2		<b>4</b>
(2006) Contribution	Weaknesses/ Potential bias Safety & Performance Appraisal Level of Evidence Suitability Device	No conflict of interest reported.     Study Method/Design     Single arm interventional study.      Relevant Data     Covered CP Stent     BIB (hand-crimped)	To evaluate the use of Covered CP Ster	nts in treatment	1 2 D1	3 Gradin D2	<b>4</b>
(2006) Contribution	Weaknesses/ Potential bias Safety & Performance Appraisal Level of Evidence Suitability	No conflict of interest reported.     Study Method/Design     Single arm interventional study.      Relevant Data     Covered CP Stent     BIB (hand-crimped)     CoA     Patients with CoA (fully grown pati	To evaluate the use of Covered CP Ster of CoA.	nts in treatment	1 2	3 Gradin	4 ng Di Ai
(2006) Contribution S&P x	Weaknesses/         Potential bias         Safety & Performance         Appraisal         Level of Evidence         Suitability         Device         Application	<ul> <li>No conflict of interest reported.</li> <li>Study Method/Design         <ul> <li>Single arm interventional study.</li> </ul> </li> <li>Relevant Data         <ul> <li>Covered CP Stent</li> <li>BIB (hand-crimped)</li> <li>CoA</li> <li>Patients with CoA (fully grown pati</li> <li>Sampling: n=30</li> <li>Mean age: 28±17.5 (range 8 to 65)</li> </ul> </li> </ul>	To evaluate the use of Covered CP Ster of CoA.	nts in treatment	1 2 D1 A1	3 Gradin D2 A2	4 ng Di Ai
Contribution S&P x	Weaknesses/         Potential bias         Safety & Performance         Appraisal         Level of Evidence         Suitability         Device         Application	<ul> <li>No conflict of interest reported.</li> <li>Study Method/Design         <ul> <li>Single arm interventional study.</li> </ul> </li> <li>Relevant Data         <ul> <li>Covered CP Stent</li> <li>BIB (hand-crimped)</li> <li>CoA</li> <li>Patients with CoA (fully grown patients are started or start</li></ul></li></ul>	To evaluate the use of Covered CP Ster of CoA.	nts in treatment	1 2 D1 A1	3 Gradin D2 A2	4



Data Contribution	Relevant Data			Grading		
Outcomes/Endpoints	- Reduction in blood pressure gradient			Yes 1	No 2	
	- Reduction in coarctation diameter					
Follow-up	- 11 months			Yes 1	No 2	
Statistical analysis	<ul> <li>Statistical significance was defined as I</li> </ul>			Yes 1	No 2	
Clinical significance		e therapy of choice in patients with comp		Yes 1	No 2	
		afe alternative to conventional stenting in	patients with			
	severe and complex CoA lesions or adv					
		Data Contribution Gra	ide (Range 4-8)		4	
	entities and Minishains					
Overall S&P Appraisal, Dis S&P Grade	LOE (4) + Suitability (5) +	Disposition and Weighting (select)	Accepted and P	ivotal 0 1	<b>ว</b>	
	Data Contribution (4) = 13	Disposition and weighting (select)	Accepted and P			
(Range 9-25)			Excluded, 22-25		, 13-21	
Relevant S&P Results				,		
Safety data	- Two stent fractures in the "old" design	of the stent, no fractures in the "new" ste	nt design			
.,	-	overed) have been produced with reinford	-	ing joints a	as the	
	"new" stent design		5	5,		
Performance data	- Blood pressure gradient: From 36 + 20	mmHg to 4 + 4 mmHg, P<0.0001				
	- Diameter at coarctation: From 6.4 +3.8	mm to 17.1 + 3.1 mm, P<0.0001				
Benefits/claims data	- Reduction in blood pressure gradient					
	- Reduction in coarctation diameter					
	- BIB allows readjustment of position aft	er inflation of the inner balloon.				
Strengths	- Covered stents were chosen:					
		ts with CoA aneurysms or previous stent-r				
		ons because of complex CoA anatomy or a				
		work of platinum iridium wire welded in a ids caused by the welding and transfers th				
		ate the welded area, once again adding to		-		
		PTFE to achieve a solid tubular structure the				
	-	eter and will stretch over the range of diar	neters of expansion			
	is initially approximately 7 mm in diame	eter and will stretch over the range of diar taut over the stent when expanded. Whe	•			
	is initially approximately 7 mm in diame to 24 mm diameter), and will always be	e taut over the stent when expanded. Whe	en the covering is			
	is initially approximately 7 mm in diame to 24 mm diameter), and will always be folded over the crimped stent and expa	e taut over the stent when expanded. Whe ands uniformly when the balloon is inflated	en the covering is			
Weaknesses/	is initially approximately 7 mm in diame to 24 mm diameter), and will always be folded over the crimped stent and expa	e taut over the stent when expanded. Whe	en the covering is			



16. Meadows et al. (2015) Contribution S&P X (S only) SOA -	adolescents and adult (NCTC	e results from the COAST trial for CP Stent (Bar	e and Covered) to treat native and recurrent CoA in select in <b>Table G-1</b> for safety and performance of the subject de			). 1.
17. Taggart et al. (2016) Contribution S&P x SOA -			e safety and short-term efficacy of the CP Stent in treating presented in <b>Table G-1</b> for safety and performance of the s			
	Safety & Performance (for s	afety only)				
	Appraisal           Level of Evidence         Study Method/Design         Question Applied					2011
		Retrospective study.	To study the early and late outcomes after stenting of native and recurrent CoA with uncovered and covered stents.	1	2 <b>3</b>	4 5
	Suitability Relevant Data					Ig
18. Sasikumar et al. (2020)	Device	<ul> <li>CP Stent (Bare and Covered) – "D1" for</li> <li>Unknown if pre-mounted on BIB</li> </ul>	subject devices tent (covered), Andra XL and XXL stents, Palmaz XL	D1	D2	D3
Contribution	Application	- CoA (native and recurrent)		A1	A2	A3
S&P X (S only) SOA x	Patient	<ul> <li>Patients with CoA (native and recurrent)</li> <li>Sampling: n=45 (20 covered stents, 25 non-covered stents)</li> <li>Covered stents used were covered 7 CP Stent; 13 Advanta V12 Stent</li> <li>Non-covered stents used were 17 CP Stent, 6 Andra XL and XXL stents, and 2 Palmaz XL.</li> <li>Mean age: 28±17.5 (range 8 to 65) years. Age per device group was not reported.</li> <li>Sex: 32 M, 13 F. Sex per device group was not reported.</li> </ul>			P2	Ρ3
	Report	<ul> <li>High quality with deficiencies</li> </ul>		R1	R2	R3
		• • • •	Suitability Grade (Range 4-12)		6	
	Data Contribution	Relevant Data			Gradin	a
	Outcomes/Endpoints	- Safety		Yes 1	Gradin	ng No 2
	Follow-up	- Covered stent group: 57 months		Yes 1		No 2



		ent group: 35 months					
Statistical analysis				ge for Social Sciences (ver		Yes 1	No 2
				s median and range and o			
				egorical parameters were			
				ompared by Student t tes			
				test for nonparametric of	data.		
Clinical significance	<ul> <li>Not reported sp</li> </ul>	ecifically for subject	devices.	Data Cantaila tian	Cuarla (Davaga 4.0)	Yes 1	No 2
				Data Contribution	Grade (Range 4-8)		5
Overall S&P Appraisal, Disp	sposition and Weighting						
S&P Grade	LOE (3) + Suitability (6	5) +	Dispositio	on and Weighting (select)	) Accepted and	d Pivotal 9-1	2
(Range 9-25)	Data Contribution (5)	= 14			Accepted bu	t not Pivota	l, 13-21
	Excluded, 22-2		-25				
Relevant S&P Results							
Safety data	Outcomes		Covered (n=1	8)	Uncovered	(bare meta	l) (n=8)
	Late lumen loss (no or mild)		2 (Advanta 1,	2 (Advanta 1, CP 1)		4 (CP 3, Palmaz 1)	
	Late lumen loss (moderate)		12 (Advanta 7, CP 4, Andra 1)		4 (CP3, Palı	4 (CP3, Palmaz 1)	
	Late lumen loss (se	evere)	4 (Advanta 3,	CP 1)	0		
	Fracture		1 Advanta		0		
Performance data	- Not reported spe	ecially for subject dev	rices.		l		
Benefits/claims data	- Not reported						
Strengths	- Not reported.						
Weaknesses/	- Not reported.						
Potential bias							
o							
<u>State of the Art</u> Appraisal							
	Alternatives	Risk/benefit	Side-effec	ts Equivale	0000	urrogate en	dnointe
	Yes 1 No 2	Yes 1 No 2	Yes 1	No 2 Yes 1		es 1	No 2
ICS I NU Z	ICJI INU Z	NO Z	163 1	inoz iest		COT	
Overall SOA Appraisal and	Disposition						
SOA Grade	8		Dispositio	on (select)	Acce	pted, < 12	
SUA GIAUE						• •	



	SOA data SOA ments Safety & Performance	<ul> <li>stent group had residual gradient &gt;10</li> <li>No mortality or aortic wall injury in eit</li> <li>Mean number of anti-hypertensive was</li> <li>Greater incidence of severe late lume the authors, this phenomenon was brain lumen obstruction was also noted in consequent less radial strength.</li> <li>A previous study on Advanta stent in However, the median period of follow re-coarctation or aneurysm formation</li> <li>Another study described 2 patients was the stent on follow-up and both the collapse in a patient who had Advan residual gradient was 5 mm Hg imm follow-up and he underwent a repeat</li> <li>Uncovered stents can be safely implication</li> </ul>	ther group. as 1.38 ± 0.74 in the covered goup and 1+0.7 in the uncover en loss (>30% lumen loss) in the covered stent group on for and specific (Advanta V12 stent). Single strut fracture which n one Advanta V12 stent. The stents have an open cell uplantation in 25 patients did not show any complications youp in that study was only 4.9 months and longer follow-up to ith Advanta stent implantation who developed in-folding of cases were managed by re-stenting. The authors had a sta stent implantation, which was managed by balloon an nediately after the balloon angioplasty, the gradient incre-	red grou illow-up h was r stent s relate b is nee f the pr similar gioplas eased t	up b. Accor not caus geomet d to the ded to l oximal proxima ty. Thou o 25mr	rding to sing any ry with e stent. look for edge of al stent ugh the mHg on
	Appraisal					
	Level of Evidence	Study Method/Design	Question Applied		rd LOE	-
		Single center retrospective study.	The aim of this study was to investigate the impact and safety of covered stent placement for treatment of (re)CoA during a longer follow-up period.	1	2 3	4 5
19. Stassen et al.	Suitability	Relevant Data			Gradin	p
(2021)	Device		n (CCP) stents (NuMED Inc. Hopkinton, NY, USA) were	D1	D2	D3
Contribution S&P x		, .	28 (18.0%), 8z34 (25.8%), 8z39 (25.8%), 8z45 (28.1%) and		52	23
SOA x	Application	- CoA (recurrent)		A1	A2	A3
		<ul> <li>Patients with CoA who were treated with 102 covered stents from 2003 to 2017</li> <li>All patients with a covered stent implantation for a native CoA or reCoA after surgical or transcatheter repair were included.</li> <li>89 patients with 102 covered stents in 93 procedures</li> </ul>			-	P3
	Patient	- All patients with a covered stent impl transcatheter repair were included.	antation for a native CoA or reCoA after surgical or	P1	P2	F3



				Gi						
Data Contribution	Relevant Data									
Outcomes/Endpoints	<ul> <li>Short-term pre/post-implant hemodynamics and angiographic data were reported. Changes in blood pressure, the use of antihypertensive drugs and complications were recorded during follow-up.</li> </ul>									
Follow-up	- Mean follow-up time was 6.6±3.7 years (min max range 0.2-15.7 years).									
Statistical analysis	<ul> <li>Continuous variables are presented as mean plus minus standard deviation (range minimum- maximum). In case of an asymmetric distribution of data, results are reported as median (interquartile range (IQR)). Proportions are noted as number and percentage. Comparison of individual parameters before and after stenting was performed using the two-tailed paired t test. Categorical data were compared with a McNemar. A p value of less than 0.05 was considered statistically significant. Statistical analysis was done using the SPSS software version</li> </ul>									
	26 package (SPSS Inc., Chicago, IL USA).									
Clinical significance	- The magnitude of the treatment effect ob	· •		Yes 1						
		Data Contribution Gra	ade (Range 4-8)							
Relevant S&P Results			( 2)							
Safety data	<ul> <li>Long-term adverse events were found in 4.</li> </ul>	.5% of patients (covered stent fracture	e (n=3), aneurysn	n formatio						
Performance data	<ul> <li>The procedural success rate was 100%.</li> <li>The mean invasive ascending-to-descending aorta systolic gradient under general anaesthesia decreased from 25 16mmHg to 4 ± 7mmHg (p&lt;0.001). After a mean follow-up time of 6.6 ± 3.7 years, there was a persistent improvement of the mean systolic blood pressure gradient between right arm and leg (~7 ± 18 vs 38 ± 24mmHg; p&lt;0.001). A larger proportion of patients required antihypertensive medication (33.7% vs 50.0%, p=0.017) and needed ≥ two drugs (20.2% vs 27.4%, p=0.066) to control blood pressure.</li> <li>Covered stent implantation for CoA is highly successful, safe and results in a persistent hemodynamic improvement is mandatory to maintain favourable hemodynamic results after stenting.</li> </ul>									
Benefits/claims data										
			- Patients were followed for a mean period of 6.6±3.7 years (maximum follow-up time 15.7 years). To knowledge, this is the largest study with the longest follow-up of the use of covered stents in (re)CoA.							
Strengths	- Patients were followed for a mean peri									



	State of the	e Art										
	Appraisal	••••										
		Medical condition		Alternatives		efit	Side-effects		Equivalence		Surrogate endpoints	
	Yes 1	No 2	Yes 1	No 2	Yes 1	No 2	Yes 1	No 2	Yes 1	No 2	Yes 1	No 2
	Overall SO	A Appraisal	and Disposit	tion								
	SOA Grad		7	7			Disposition (select)			Accepted, < 12 Excluded, 12		
	(Range 6-	12)										
	Relevant S											
	SOA data	OA Results	CoA:									
Holzer et al.	Comment Safety & Pe		- 1 - 1 - 5 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1	<ul> <li>CoA is a congenital cardio-vascular malformation, characterised by a restriction of the lumen of the thoracic aorta. occurs in approximately 4 of 10,000 live births and comprises 5% to 8% of CHD.</li> <li>Mostly, CoA is detected in childhood and repaired surgically or by endovascular therapy. Occasionally it is diagnosed i adolescence or adulthood by investigations done for systemic hypertension.</li> <li>The natural history of CoA carries a poor prognosis due to complications such as left ventricular failure, intracrania haemorrhage, aortic rupture or dissection, premature coronary artery disease and sudden death.</li> <li>Smaller and younger infants are typically treated surgically but remain at risk for recurrent obstruction with up to 100 requiring further intervention during adulthood.</li> <li>In older children and adults, the preferred treatment method depends on the individual anatomy and nature of the lesion, but endovascular therapy with either balloon angioplasty or stent implantation is commonly preferred over surgery. Although balloon angioplasty results in excellent acute hemodynamics, it is associated with a high rate or aortic wall injury and recurrent obstruction. Because of these concerns, stent implantation is usually favoured to avoid overdilation or the elastic recoil of the aorta. Bare stent implantation has become a worthy alternative to surgery an balloon angioplasty and seems to lead to better results and fewer complications. However, although interventions wit bare stent implantation seem efficient and generally safe, major complications such as local aneurysm formation aortic rupture, dissection and even death may occur. To prevent these aortic wall injuries (AWI) during the ster procedure, covered stents are increasingly used and their safety and efficacy for immediate and intermediate follow up have been demonstrated. However, long-term results remain limited.</li> <li>Covered stents are increasingly used in severe and complex coarctations of the aor</li></ul>								
(2021)	Appraisal											
ontribution	Appraisal	vidence	Study	y Method/De	esign		Question	Applied			Oxfo	rd LOE 2011
		TUCINC	Juu	, meenou/De								



SOA x		follow-up data from patients in these 2 trials and those treated through their Continued Access protocols. Data was prospectively collected during the 2 multi-compare this to earlier follow-up data. For the purpose of this study, follow-up was defined as: Immediate (1 month), Early (12 months),			
		center studies Late (48 or 60 months). To identify possible predictors of late-term outcome post-stent implantation.			
	Suitability	Relevant Data		Gradin	σ
	Device	- CP Stent (Bare and Covered)	D1	D2	D3
		<ul> <li>52% received covered stents and 48% received bare stents.</li> <li>No data if pre-mounted or not with BIB</li> <li>The minimum stent diameter was 14.4mm (interquartile range (IQR), 12.6-16.0mm) with a minimum stent diameter to the aorta at diaphram ratio of 0.87 (IQR, 0.77-1.0).</li> </ul>			
	Application	<ul> <li>CoA (native or recurrent)</li> <li>Native coarctation was present in 49%, postsurgical in 24% and postcatheterization in 27%.</li> <li>The minimum coarctation diameter was 8.0mm (IQR, 5.4-10.5mm), and median aortic diameter at the diaphragm was 16.0mm (IQR, 14.0-19.0mm).</li> </ul>	A1	A2	A3
	Patient	<ul> <li>All patients enrolled in the COAST or COAST II trials and their Continued Access extensions were included. Patients without late follow-up data were excluded from analysis, except for analyzing the estimated cumulative incidence of stent fractures, aortic wall injury, and reinterventions.</li> <li>Cohort of 248 patients</li> <li>COAST: 105 patients enrolled in COAST with 16 Continued Access patients (n=121)</li> <li>COAST II: 82 participants from COAST II with an additional 45 Continued access patients (n=127).</li> <li>From the 180 patient cohort, the median age at implant was 17 years (IQR, 13-28 years), the median weight (66.3kg, IQR, 53.8-78.1kg).</li> </ul>	P1	P2	P3
	Report	- High quality report	R1	R2	R3
		Suitability Grade (Range 4-12)		4	
	Data Contribution	Relevant Data		Gradin	g
	Outcomes/Endpoints	<ul> <li>Parameters used to assess aortic stent outcomes:         <ul> <li>Hemodynamic</li> <li>Systemic systolic hypertension</li> <li>Use of antihypertensive medication</li> <li>Upper limb to lower limb blood pressure difference of ≥20mm Hg</li> <li>Reinterventions</li> <li>Stent fractures</li> <li>Aortic wall injury</li> </ul> </li> </ul>	Yes 1		<u>в</u> No 2

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	<ul> <li>Predictor variables used to assess late-term results:</li> </ul>		
	-Demographics		
	-Type of coarctation		
	-Preimplantation clinical data		
	-Baseline characterization data		
	-Type of stent		
	-Poststent catherization data		
	-Postcatheterization data		
Follow-up	- Follow-up data was collected at 1, 6, 12, 24, 36, 48 and 60 months and included MRI at 12 and	Yes 1	No 2
	24 months, and fluoroscopy at 12, 24, 48 and 60 months.		
	- 96% of patients returned for 1-month follow-up, 86% for 12-month follow-up, and 63% for 60-		
	month.		
	- A total of 180 patients (73%) had either 48- or 60-month follow-up data.		
	- Out of the 180 patients with late follow-up, 177 (98%) had also immediate and 180 (100%)		
	early follow-up data available for analysis.		
	- Aortic imaging (either MRI, computed tomography, or angiography) was available for 180/180		
	(100%) at immediate follow-up, 177/180 (98%) at intermediate follow-up, and 41:180 (23%) at		
	late follow-up. Fluoroscopy was available for 180/180 (100%) at immediate follow-up, 178/180		
	(99%) at intermediate follow-up, and 136/180 (76%) at late follow-up.		
Statistical analysis	<ul> <li>Categorical variables are summarized as frequencies and percentages, and continuous</li> </ul>	Yes 1	No 2
,	variables as either means and SDs or medians with interquartile range (IQR) as noted. For the		
	entire cohort, the cumulative incidence of stents fractures, aortic wall injury, and		
	reinterventions at immediate, early and late follow-up was estimated using the Kaplan-Meier		
	method. Patients who did not have an outcome event were censored at time. Changes in		
	hemodynamic measures over time were evaluated using tests of trend. For patients with late		
	follow-up, associations between patient and procedure characteristics and 4 binary outcome		
	variables – suboptimal hemodynamic outcome, stent fractur, catheter reintervention, and		
	aortic wall injury – were assessed using Fisher exact test. Characteristics significant at the 0.20		
	level were considered for inclusion in multivariable logistic regression models. Forward		
	selection was used, and P <0.05 was required for retention in the final model. To assess		
	generalizability, characteristics of patients with and without late follow-up were compared		
	using Fisher exact and Wilcoxon rank sum tests; no significant differences were found. All		
	analytics were performed using SAS software version 9.4.		
Clinical significance	<ul> <li>Coarctation stenting is effective at maintaining obstruction relief up to 60 months postimplant</li> </ul>	Yes 1	No 2
	with reduction in the number of patients requiring antihypertensive medication. However, an		
	increase in-stent fractures and reinterventions were observed between medium and long-term		
	follow-up. Covered stents appear to confer some protection from the development of stent		
	fractures but do not provide complete protection from late aneurysm formation.		
	Data Contribution Grade (Range 4-8)		4
1	Data Contribution Grade (Kange 4-6)		4

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Overall S&P Appraisal	, Disposition and Weighting		
S&P Grade	LOE (3) + Suitability (4) +	Disposition and Weighting (select)	Accepted and Pivotal 9-12
(Range 9-25)	Data Contribution (4) = 11		Accepted but not Pivotal, 13-21 Excluded, 22-25
Relevant S&P Results			
Safety data	<ul> <li>No dissections were found.</li> <li>The cumulative incidence was 1.2%</li> <li>In 3 patients, the aneurysm was protter reminder, the aneurysm was witter in 4 of 13 patients, aneurysms were patients the aneurysms were diagnonelective stent re-expansion.</li> <li>17 patients had covered stents implated by univariate analysis, coarctation mwall injury (12% versus 2%, P=0.007)</li> <li>There was a borderline relationship</li> </ul>	ximal to the implanted stent, in one patient t hin the borders of the implanted stent. identified on MRI or computed tomography used by angiography during catheterization po anted to treat the aneurysm; 2 did not. inimum diameter <6mmm was the only factor	he location was not specified, and in before reintervention, while in 9 erformed for other reasons such as or significantly associated with aortic
	<ul> <li>implanted. As such, the notion that of aneurysm, may not be the case. Dincidence of aneurysm formation whe cohort. Also, the median follow-up in covered stents (85 versus 35 months) aneurysms were not identified until</li> <li>Most aneurysms developed within the pressure within the aorta distributes formation. Another possibility is that implantation.</li> <li>Current study did not investigate the stent implantation because cases had stent implantation and received covered the Adverse Events:</li> <li>Over the follow-up period, 2 patient</li> </ul>	he borders of the stent, including covered ste s flow between the stent and the aortic wall, t the expanded polytetrafluoroethylene beca e benefit of a covered stent to reduce the risk ve not been randomly assigned and high-risk ered stents. s had additional adverse events that were ca	a protection from the development d show a significant difference in the stents, albeit in a much smaller e with bare stents compared with onstrates that the majority of ents. One possible explanation is that eventually leading to aneurysm me damaged during initial c of acute aortic wall injury during patients were excluded for bare ptured in the data set. One patient
	without any clear relationship to the	erse event (possible transient ischemic attac procedure itself. Another patient developed is adverse events were documented in any pa	cardiogenic/sceptic chock 7 months

<sup>&</sup>lt;sup>1</sup> Butera G, Manica JL, Marini D, Piazza L, Chessa M, Filho RI, Sarmento Leite RE, Carminati M. From bare to covered: 15-year single center experience and follow-up in transcatheter stent implantation for aortic coarctation. Catheter Cardiovasc Interv. 2014 May 1;83(6):953-63. doi: 10.1002/ccd.25404. Epub 2014 Feb 4. PMID: 24459104.



Renefits											
Denents	/claims data		<ul> <li>At late follow-up, freedom from surgical intervention was 100%, catheter reintervention 78.7%, stent fracture 75.6%, and freedom from aortic wall injury 93.7%.</li> </ul>								
					, ,						
			•	ts had subopti	-	•			C - 11	and Chart	
										period. Stent f medium and lo	
						•				t fractures, but	-
			•	lete protection	•	•			it of steri		they do in
Strength	ıs							o 60 months po	ost-proced	dure.	
Weakne	sses/	- S	mall sample :	size							
Potentia	ıl bias				power to eva	iluate all para	meters conti	ributing to long	-term mo	orbidity in these	e patients,
			uch as aortic								
						•			or additior	nal data regard	ing stent
			,	cations for rei					tions and	the way some	of the data
			vas collected.		IICES DELWER			onnent muica		the way some	
					to 60 montl	is follow-up a	s long-term,	this is still a rel	atively sh	ort time period	Ι.
										s. Ot did not co	
							odalities, as v	vas done in the	Congenit	tal Cardiovascu	lar
			nterventional	Study Consor	tium Report	.2					
State of t	<u>he Art</u>										
Appraisal						-					
Appraisal Medical	condition	Alternati		Risk/bene	1	Side-effec		Equivalenc	1		endpoints
Appraisal		Alternati Yes 1	ves No 2	Risk/bene Yes 1	fit No 2	Side-effec Yes 1	ts No 2	Equivalence Yes 1	ce No 2	Surrogate	endpoint: No 2
Appraisal Medical Yes 1	condition No 2	Yes 1	No 2	-	1				1		1
Appraisal Medical Yes 1 Overall So	condition No 2 OA Appraisal ar	Yes 1	No 2	-	1	Yes 1	No 2		1	Yes 1	No 2
Appraisal Medical Yes 1 Overall SC SOA Gra	condition No 2 OA Appraisal ar	Yes 1	No 2	-	1	Yes 1			1		No 2
Appraisal Medical Yes 1 Overall So	condition No 2 OA Appraisal ar	Yes 1	No 2	-	1	Yes 1	No 2		1	Yes 1 Accepted, < 1	No 2
Appraisal Medical Yes 1 Overall SO SOA Gra (Range 6	condition No 2 OA Appraisal ar	Yes 1	No 2	-	1	Yes 1	No 2		1	Yes 1 Accepted, < 1	No 2
Appraisal Medical Yes 1 Overall SO SOA Gra (Range 6	condition No 2 OA Appraisal an ade 5-12) SOA Results	Yes 1 nd Dispositi 7 CoA:	No 2	Yes 1	No 2	Yes 1       Disposition	No 2	Yes 1	No 2	Yes 1 Accepted, < 1 Excluded, 12	No 2
Appraisal Medical Yes 1 Overall SC SOA Gra (Range 6 Relevant	condition No 2 OA Appraisal an ade 5-12) SOA Results	Yes 1 nd Dispositi 7 CoA: - C	No 2	Yes 1	No 2	Yes 1 Dispositio	No 2 on (select) ancy by surg	Yes 1 ery. Beyond in	No 2	Yes 1 Accepted, < 1 Excluded, 12 ercutaneous tre	No 2
Appraisal Medical Yes 1 Overall SO SOA Gra (Range 6 Relevant	condition No 2 OA Appraisal an ade 5-12) SOA Results	Yes 1 nd Dispositi 7 CoA: - C e	No 2 on oA is repaire ither balloon	d during the angioplasty or	No 2 neonatal pe	Yes 1 Dispositio	No 2 on (select) ancy by surg	Yes 1 ery. Beyond in tly employed to	fancy, pe	Yes 1 Accepted, < 1 Excluded, 12	No 2 2 eatment unt coarctat

<sup>&</sup>lt;sup>2</sup> Forbes TJ, Kim DW, Du W, Turner DR, Holzer R, Amin Z, Hijazi Z, Ghasemi A, Rome JJ, Nykanen D, Zahn E, Cowley C, Hoyer M, Waight D, Gruenstein D, Javois A, Foerster S, Kreutzer J, Sullivan N, Khan A, Owada C, Hagler D, Lim S, Canter J, Zellers T; CCISC Investigators. Comparison of surgical, stent, and balloon angioplasty treatment of native coarctation of the aorta: an observational study by the CCISC (Congenital Cardiovascular Interventional Study Consortium). J Am Coll Cardiol. 2011 Dec 13;58(25):2664-74. doi: 10.1016/j.jacc.2011.08.053. PMID: 22152954.



	coarctation. It has rounded edges to reduce the incidence of aortic wall injury and can be expanded to 24mm diameter.
	- Stent implantation, balloon angioplasty, and surgery are all treatment options for coarctation in patients beyond
	<ul> <li>infancy.</li> <li>Treated coarctation is associated with long-term morbidity irrespective of treatment strategy.</li> </ul>
	COAST Trials:
	<ul> <li>The FDA pivotal trials COAST (Coarctation of the Aorta Stent Trial; 2007-2016) and COAST II (Covered Cheatham- Platinum Stents for Prevention or Treatment of the Aorta; 2010-2016) demonstrated safety and efficacy of the bare and Covered CP Stents when used to treat aortic coarctation. Short- and medium-term results have been previously</li> </ul>
	reported (Meadows et al. (48) and Taggart et al. (49)).
	<ul> <li>The Covered CP Stent is a CP stent covered by a 0.28" sleeve of 0.005" thick expanded polytetrafluoroethylene tubing and was available to centers participating in the COAST trial for compassionate and emergency use for aortic wall injury occurring during aortic interventions.</li> </ul>
	- COAST II included patients who received a Covered CP stent as an emergency or compassionate use during the initial COAST trial (legacy arm) and prospectively enrolled patients between 2010 and 2011.
	<ul> <li>COAST II included higher-risk groups, such as patients with aortic wall injuries and those with nearly atretic descending aorta of 3mm or less diameter.</li> </ul>
Comments	Hemodynamic Outcome:
	<ul> <li>Study corroborates the results from the largest multi-center study of stenting for coarctation from the Congenital Cardiovascular Interventional Study Consortium, which reported 23% systolic hypertension at 12 to 60 months of follow-up, 9% arm-leg blood pressure gradient ≥20 mm Hg, 23% need for antihypertensive medication and the presence of any of these 3 in 37%.<sup>3</sup></li> </ul>
	Stent Fractures: - Previous studies of the bare metal CP stent documented stent fractures of 2% at 12 months, and 12% at 24 months
	<ul> <li>Previous studies of the bare metal CP stent documented stent fractures of 2% at 12 months, and 12% at 24 months (Meadows et al. (48)). While the design and metallic composition of the CP stent may contribute, stents fractures are not limited to CP stents.<sup>4</sup> Boe et al.<sup>5</sup> reported a 21% fracture rate for Palmaz Genesis XD stents when used for coarctation therapy in children &lt; 20Kg at a mean follow-up of 75 months.</li> </ul>
	- It is unclear whether somatic growth can add additional force and loading conditions to the implanted stent, or
	<ul> <li>whether participation in contact sports might impact the incidence of stent fractures.</li> <li>Bare metal stents have a s significantly higher fracture rate than covered CP stent. Possible explanations could be that</li> </ul>
	the struts of a bare stent become more solidly embedded into the aortic wall, and that the expanded
	polytetrafluoroethylene covering more equally distributes the radial force to multiple struts or that it reduces the

<sup>&</sup>lt;sup>3</sup> Holzer R, Qureshi S, Ghasemi A, Vincent J, Sievert H, Gruenstein D, Weber H, Alday L, Peirone A, Zellers T, Cheatham J, Slack M, Rome J. Stenting of aortic coarctation: acute, intermediate, and long-term results of a prospective multi-institutional registry--Congenital Cardiovascular Interventional Study Consortium (CCISC). Catheter Cardiovasc Interv. 2010 Oct 1;76(4):553-63. doi: 10.1002/ccd.22587. PMID: 20882661.

<sup>&</sup>lt;sup>4</sup> McElhinney DB, Marshall AC, Schievano S. Fracture of cardiovascular stents in patients with congenital heart disease: theoretical and empirical considerations. Circ Cardiovasc Interv. 2013 Oct 1;6(5):575-85. doi: 10.1161/CIRCINTERVENTIONS.113.000148. PMID: 24129934.

<sup>&</sup>lt;sup>5</sup> Boe BB, Loccoh E, Stockmaster K, Holzer RJ, Cheatham SL, Cheatham JP, Armstrong A, Berman DP. Median and long-term outcomes of stent implantation for coarctation of the aorta in small patients (<20 kg). [Abstract presented at PICS 2019]. *J Struct Heart Dis.* 2018;4:140.



<ul> <li>Reinterventions:</li> <li>Previously reported data documented transcatheter reinterventions of about 5% by 24 months follow-up (Meadows et al. (48)).</li> <li>There is no expert consensus defining when a reintervention should be performed.</li> <li>Reinterventions in this patient population are not unexpected and do not represent a poor outcome.</li> <li>Aortic Wall Injury:</li> <li>Aneurysms did not just occur in patients with bare metal stents, but equally in patients who had covered stents implanted. As such, the notion that covered stent implantation confers long-term protection from the development of aneurysm, may not be the case. Data are in contrast with Butera et al.<sup>6</sup> who did show a significant difference in the incidence of aneurysm formation when comparing patients bare versus covered stents, albeit in a much smaller cohort. Also, the median follow-up in that study was significantly longer for those with bare stents compared with covered stents (85 versus 35 months). This is important as the current study demonstrates that the majority of aneurysms were not identified until late follow-up.</li> <li>Most aneurysms developed within the borders of the stent, including covered stents. One possible explanation is that pressure within the aorta distributes flow between the stent and the aortic wall, eventually leading to aneurysm formation. Another possibility is that the expanded polytetrafluoroethylene became damaged during initial implantation.</li> <li>Current study did not investigate the benefit of a covered stent to reduce the risk of acute aortic wall injury during stent implantation and received covered stents.</li> </ul>

<sup>&</sup>lt;sup>6</sup> Butera G, Manica JL, Marini D, Piazza L, Chessa M, Filho RI, Sarmento Leite RE, Carminati M. From bare to covered: 15-year single center experience and follow-up in transcatheter stent implantation for aortic coarctation. Catheter Cardiovasc Interv. 2014 May 1;83(6):953-63. doi: 10.1002/ccd.25404. Epub 2014 Feb 4. PMID: 24459104.

### An overall summary of the clinical performance and safety:

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A comprehensive, systematic, and critical evaluation of the pertinent clinical data and pre-clinical study data in relation to the Covered Stents has been carried out and documented in this report. Based on the results of this evaluation, it is considered that:

- a) Conformity with relevant general safety and performance requirements set out in MDR Annex I under the normal conditions of the intended use of the device has been confirmed.
- b) Undesirable side-effects and acceptability of the benefit-risk ratio have been evaluated and are acceptable according to the current knowledge/the state of the art in the medical fields concerned and according to available medical alternatives.
- c) The information materials supplied by NuMED, and the risk reduction measures are adequate taking into account the intended purpose of the device.
- d) Usability aspects have been adequately considered and the Covered Stents and NuDEL including the IFUs, is suitable for the intended users.
- e) The claims foreseen in the information materials provided with the CER are adequate taking into account the intended purpose of the device.
- f) The information materials supplied and the RM documentation for the device under evaluation are consistent with the clinical data and pre-clinical study data presented in this CER and with the current knowledge/state of the art.

Overall, it is concluded that the risks associated with the use of the Covered Stents and NuDEL are acceptable when weighed against the benefits to the patient and are compatible with a high level of protection of health and safety, taking into account the generally acknowledged state of the art; that the intended clinical performances are achieved by the device; and that known and foreseeable risks and undesirable side-effects are considered acceptable when weighed against the benefits from performance achieved by the device.

#### Ongoing planned post-market clinical follow-up:

The Stent Device Family has been on the market since 2004 in the EU and 1999 in other markets. Over time variants of the Stent Device Family have been introduced to these markets. Since then, the devices are likely to have been used in a variety of patients and populations. The Stents have been subjected to several clinical investigations where efficacy and safety has been demonstrated.

For the original Stent Device Family, a PMCF study is not warranted at this time due to the fact that the long-term safety and clinical performance has been established via device use and ample clinical experience. This experience would likely have identified any rare complications or problems that would become apparent only after widespread device use. Continued PMS activities will provide sufficient data to adequately address clinical risks, and detect emerging risks on the basis of evidence.

A PMCF study was initiated in 2018 for the additional sizes that were added to the product line, and another one in 2021 for the new G-Armor devices, to determine if there were any new complications which were previously not addressed through actual clinical use, or if any new risks are introduced. Each study had a target size of 59 patients, based on a confidence level of 95%. The studies were conducted by issuing a form to the treating physician and collecting data. The results of the 2018 study are included in the clinical data that is used for the clinical evaluation. The 2021 study for the G-Armor stent line is still going.

### 6. Possible diagnostic or therapeutic alternatives

Alternative treatments for CoA include surgery or balloon angioplasty.

Alternative treatments for RVOT include surgery, transcatheter pulmonary valve replacement, or balloon valvuloplasty / angioplasty (to delay the need for replacement only).

## 7. Suggested profile and training for users

The RVOT/COA Stent Device Family is intended for use by a Cardiac surgeon and/or interventionalist.

#### 8. Reference to any harmonised standards and CS applied

There are no Common Specifications for this type of device.

The following harmonised standards are followed for this device:

- EN ISO 10993-10: 2023 Biological Evaluation of Medical Devices Part 10: Tests for Skin Sensitization
- EN ISO 10993-23: 2021 Biological Evaluation of Medical Devices Part 23: Tests for Irritation



- EN ISO 11135: 2014 / A1:2019 Sterilization of health-care products Ethylene oxide Requirements for the development, validation and routine control of a sterilization process for medical devices.
- EN ISO 11737-1: 2018 / A1:2021 Sterilization of medical devices Microbiological methods Part 1: Determination of a population of microorganisms on products
- EN ISO 13485: 2016 / A11:2021 Medical devices Quality management systems Requirements for regulatory purposes
- EN ISO 14971: 2019 / A11:2021 Medical Devices Application of Risk Management to Medical Devices
- EN ISO 15223-1: 2021 Medical devices Symbols to be used with medical device labels, labelling and information to be supplied Part 1: General requirements

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SSCP revision number	Date Issued	Change Description	Revision validated by Notified Body
00	21 June 2022	Initial implementation	☐ Yes Validation Language: English ⊠ No
01	06 July 2023	Updated sections 4, 5, 7, 8, and 9 for CER Update.	☐ Yes Validation Language: English ☑ No
02	28 July 2023	Updated Section 2.	Ves Validation Language: English



Document Revision: 02 Date issued: 28 July 2023

This Summary of Safety and Clinical Performance (SSCP) is intended to provide public access to an updated summary of the main aspects of the safety and clinical performance of the device. The information presented below is intended for patients or lay person. A more extensive summary of its safety and clinical performance prepared for healthcare professionals is found in the first part of this document.

The SSCP is not intended to give general advice on the treatment of a medical condition. Please contact your healthcare professional in case you have questions about your medical condition or about the use of the device in your situation. This SSCP is not intended to replace an Implant card or the Instructions for Use to provide information on the safe use of the device.

1. Device identification	n and general information
Device trade name(s)	Covered CP Stent Covered Mounted CP Stent G-Armor Covered Stent G-Armor Covered Mounted Stent
Manufacturer's name and address	NuMED, Inc. 2880 Main Street Hopkinton, NY 12965 USA
Year when first certificate (CE) was issued	2004 (Covered CP Stent) 2009 (Covered Mounted CP Stent) G-Armor Devices – Not yet CE Marked
Basic UDI-DI	08877141650TH

2. Intended use of the	device
	The Stents are intended for implantation in the native and/or recurrent coarctation of the aorta.
	An aortic coarctation is a partial blockage or narrowing in the aorta, the body's main blood vessel distributing blood to all parts of the body. This blockage of the aorta makes the heart work harder to pump blood to your body and can weaken the heart muscle. Furthermore, this blockage can cause severe upper body hypertension (high blood pressure), increasing the risk of stroke. This blockage is present from birth.
Intended purpose	The Stents are also intended for treatment of right ventricle to pulmonary artery (right ventricular outflow tract) conduit disruptions that are identified during conduit pre-dilatation procedures performed in preparation for transcatheter pulmonary valve replacement.
	A Right Ventricular Outflow Tract (RVOT) is also known as a pulmonary conduit, and it is a tube that connects the heart to the lungs. Placement of an RVOT is typically associated in patients that have one of the following conditions: Pulmonary Atresia, Tetralogy of Fallot, or Double Outlet Right Ventricle. These three conditions can lead to pulmonary conduit failure.
Indications and intended patient groups	The device is used to treat any patients that have an aortic coarctation or RVOT conduit disruptions as long as none of the below listed contraindications and/or limitations are applicable.
Contraindications and/or limitations	<ul> <li>The following patients should NOT receive the Stent:</li> <li>Patients who are too small to allow the stent to pass through their arteries without damaging the artery;</li> <li>Patients with a stiff aorta that does not get larger with balloon dilation. (CoA only)</li> </ul>



<ul> <li>Patients with blocked leg arteries making it difficult or unsafe to move the catheter and stent to the narrowed aorta; (CoA only)</li> <li>Patients with any signs of infection;</li> <li>Patients with active infection in the heart or blood vessels (endocarditis);</li> <li>Patients with a known allergy to aspirin, other antiplatelet agents, or heparin; (CoA only)</li> </ul>
Pregnancy.

3. Device description	
	The Stents are balloon expandable and intended to permanently stay in your body. The Stents are used for coarctation of the aorta or treatment of right ventricle to pulmonary artery (right ventricular outflow tract) conduit disruptions that are identified during conduit pre-dilatation procedures performed in preparation for transcatheter pulmonary valve replacement (TPVR). The covering acts as a fluid barrier creating a fluid tight conduit through the stent length. Blood cannot flow across the covering.
Description of the device	The BIB Stent Placement Catheter is triaxial in construction with two lumens being used to inflate the balloon while one lumen is being used for tracking over a guidewire. The inner balloon is $\frac{1}{2}$ of the outer balloon diameter and 1 cm shorter. The purpose of the double balloon catheter is to apply an incremental inflation for the purpose of dilating a stent. The inner balloon provides initial expansion of the stent and also acts as a tool to hold the stent on the catheter prior to the outer balloon being inflated. The outer balloon is then inflated, providing the remainder of the expansion. There are radiopaque platinum marker bands under the balloon shoulders, to aid during placement. The balloons are designed to inflate to the diameter and length listed on the label at a specific pressure. Thus, it is recommended that the device be used in conjunction with a mechanism to monitor pressure, an inflation device with pressure gauge.
	The Stents are composed of heat-treated metal (90% platinum and 10% iridium) wire that is arranged in laser welded rows with a "zig" pattern. The number of rows determines the unexpanded length of the stent. The Covered versions have an ePTFE covering that is attached to the metal wire frame
Medicinal Substances	The Stents do not contain any medicinal substances.
Mode of Action	The Stents are implanted using a thin hollow tube (catheter) with a balloon on the end. Your physician will place the stent on the balloon at the start of your procedure. The catheter with the stent is then placed through the skin, typically into the artery in your upper leg. The balloon and stent are moved to the appropriate position at the narrowed part of your aorta or in the RVOT. Once in place, the balloons are inflated to expand the stent. The catheter is then removed from the body and the stent stays in place.
Description of Accessories	All Stents are packaged and shipped to the physician with hemostasis valve tools. These tools are hollow tubes that are placed in the valve of the introducer to help the Stent move through that valve without any issues. The valve of the introducer is very tight to prevent blood loss during the procedure, so the tools help the Stent move through the valve without causing damage to the stent or moving the stent on the catheter.

## 4. Risks and Warning

Contact your healthcare professional if you believe that you are experiencing side effects related to the device or its use or if you are concerned about risks. This document is not intended to replace a consultation with your healthcare professional if needed.

How potential risks have been controlled or managed	The Stent Device Family has been developed in accordance with documented processes to ensure that it is designed, manufactured, packaged, and labelled in accordance with the current state of the art and meets all requirements of the appropriate regulations. Design verification activities were performed and include pre-clinical testing and clinical investigations. A clinical literature review has
	also been performed on the Stent Device Family. All risks identified during these activities were



	mitigated as far as possible and are considered acceptable in regards to the clinical benefit of the device. Continued review of all Post Market Surveillance and Post Market Clinical Follow-up Data is performed to identify any additional risks that may be identified after the device was placed on the market.
Remaining risks and undesirable effects	<ul> <li>Cardiac catheterization and stent insertion carry certain risks. Potential complications &amp; adverse effects associated with device use and indication include:</li> <li>Femoral Artery Injury</li> <li>Stent Migration – movement of the stent away from original implant site</li> <li>Stent Stenosis – growth of tissue within the stent, leading to return of the blockage</li> <li>Stent Fracture – break in the frame of the stent</li> <li>Aneurysm/Pseudoaneurysm – weakening or injury of the aorta wall</li> <li>Aortic Rupture/Tear – perforation or tearing of the aorta, causing internal bleeding</li> <li>Stent Malposition – poor position of stent, requiring a 2nd stent</li> <li>Hematoma – bruising at the site where the device is introduced into the body</li> <li>Sepsis/infection – Infection</li> <li>Thrombosis – formation or presence of a blood clot</li> <li>Embolization – passage and lodging of an embolus within the bloodstream</li> <li>Transitory arrhythmia – Irregular heartbeat</li> <li>Endocarditis - infection within the stent</li> <li>Bleeding - at the site of where the device is introduced into the body</li> <li>Cerebrovascular Incident - stroke</li> <li>Death</li> </ul>
Warning and Precautions	The majority of warnings and precautions listed for the Stents pertain to the placement and use of the device in the cath. lab by the physician. MRI Conditional information is applicable to the Stents after they are implanted. This information should be used by any MRI technician that is performing an MRI procedure on any patient with a NuMED Stent implanted. All patients will be provided with an Implant Card after their procedure. This Implant Card will give the location of where to find the most up to date MRI parameters to be used for patients that have a NuMED Stent implanted.
Summary of any field safety corrective actions (FSCA including FSN) if applicable	There have not been any Field Safety Corrective Actions or Field Safety Notices on any versions of the Stents listed in this SSCP.

5. Summary of clinical evaluation and post-market clinical follow-up		
Clinical background of the device	The NuMED Stent Device Family has been sold globally since 1999.	
	The NuMED Stent was tested and found to be safe and effective to repair aortic wall injuries and to widen the narrow part of the aorta related to coarctation of the aorta. A study was conducted with 82 patients weighing more than 31 lbs at the time of implant. Most of the patients (89%) were treated with one Covered CP stent, 11% needed more than one to complete the repair.	
	On average arm systolic blood pressure was 25 mmHg higher than the leg pressure before the procedure. A reduction of a gradient to 15mmHg or less following the procedure suggests that the blockage is reduced effectively. By one month after covered stent placement the average arm pressure was only 1 mmHg higher than the leg pressure. Two years after implant, 85% of patients had arm blood pressures less than 15 mmHg above their leg pressure, which suggests that most of the treated aortas did not re-narrow. Repair of aortic wall injury was successful in all of the 49 patients	



	who received their Covered CP Stent to repair their weakened aortic wall. An overview of complications and additional treatments provided after the stenting
	<ul> <li>procedure is shown below:</li> <li>Serious complications related to the Covered CP Stent or implant procedure, such as: causing injury to the aortic wall or damage to the leg artery used for Stent insertion, were identified in 6 out of 100 (6%) of patients within the first month of implant.</li> </ul>
	• No patients needed surgery to repair the aorta or to remove the stent.
	• One patient required stent repair of the leg artery damaged during insertion of the implant catheter.
	• One patient required surgical repair of the leg artery damaged during insertion of the implant catheter.
	• 1 out of 20 (5%) patients developed small aneurysms (weakened areas of the aorta) in the area of stent placement in the years following stent therapy, making CT or MRI imaging an important part of follow up care. However, none of the patients who developed aneurysms demonstrated symptoms or required surgery. All were successfully treated with additional covered stent placement.
	<ul> <li>Overall, 16% of patients required repeat cardiac catheterization for a second dilation of the stent, mostly to keep up with the size of the patient as he/she grew and for some to repair aortic wall injuries as noted above.</li> </ul>
	The NuMED Covered Stent was tested and found to be safe and effective to use as a Treatment of right ventricle to pulmonary artery (right ventricular outflow tract) conduit disruptions that are identified during conduit pre-dilatation procedures performed in preparation for transcatheter pulmonary valve replacement (TPVR). A study was conducted with 50 patients weighing an average of 58 kg. at the time of implant. Most patients (80%) were treated with one Covered CP stent.
	Out of 49 patients treated with the Covered CP Stent (CCPS), 81.6% of them had device and lesion success with no adverse events attributed to the CCPS. Out of 49 patients treated with the CCPS, 93.9% of the patients had successful coverage of conduit disruption followed by successful implantation of an artificial valve. An overview of complications and additional treatments provided after the stenting procedure is shown below:
	<ul> <li>Serious complications related to the CCPS or stent implant procedure, such as: stent embolization was identified in 1 out of 50 (2%) patients.</li> <li>7 (14%) of the patients required a second CCPS, and (3) 6% of the patients required a third CCPS during the procedure. Of these 10 patients, 4 (40%) of them planned on having the second CCPS implanted before the procedure.</li> </ul>
The clinical evidence for the CE marking	The CE marking was based on data from three clinical studies, a review of published literature, and a review of post market surveillance data provided by NuMED. Additional pre-clinical testing was performed as part of the development and design of the device. In vitro (on the bench) testing was performed on the devices as part of the Design History File. Biocompatibility testing was also performed on the materials used to manufacture this device to determine if it met the requirements for an implant in the human body. The device passed all tests.
Safety	The clinical data and pre-clinical study data demonstrated that the device performed as intended by NuMED in the clinical setting; the device does not pose unacceptable safety concerns in the clinical setting; and any risks associated with clinical use of the device are acceptable when weighed against the benefits to the patient.



## 6. Possible diagnostic or therapeutic alternatives

When considering alternative treatments, it is recommended to contact your healthcare professional who can take into account your individual situation.

### **Coarctation of the Aorta**

Your cardiologist believes that relief of the blockage is important for your health and safety. There are three ways to relieve the blockage: by surgery, by stent implantation without surgery, or by balloon angioplasty.

### **Surgical Therapy**

Surgical treatment of the blockage is usually performed through an incision on the side of the chest, approaching the aorta by spreading the ribs. The narrowed portion of the aorta is removed and then the aorta is sewn back together. For more complicated coarctation, surgery might be performed from the front of the chest, opening the breast bone and using heart lung bypass. For some patients a benefit of a surgical approach is that the repair can be performed without the use of man-made materials. However, for other (especially adult) patients a man-made tube graft or patch may be needed. Please consult with your surgeon regarding his or her approach. For younger patients, surgery results in a lower need for a second procedure to keep up with growth when compared to balloon or stent therapy.

Risks of surgery include: pain from the surgical incision, prolonged fluid drainage from the chest after surgery, chest or wound infection, longer recovery time compared to stent therapy, prolonged postoperative rib discomfort and increased risk of very high blood pressure occurring after immediately after surgery, requiring intravenous therapy in an ICU, compared to stent repair. There is a low risk, probably less than 5%, of developing an aneurysm (weakened areas of the aorta) in the area of surgery in the years following stent therapy, making CT or MRI imaging an important part of follow up care.

### **Stent Therapy (without surgery)**

A stent is an expandable metal tube that is implanted into your aorta to keep it open. Surgery is not required for this procedure. The stent is implanted using a thin hollow tube (catheter) with a balloon on the end. The catheter with stent is inserted through the artery in the upper leg. The balloon and stent are then moved to the appropriate position to the narrowed part of your aorta. Once in place, the balloons are inflated to expand the stent against the aortic wall. The catheter is then removed from the body and the stent remains in place.

### **Balloon Angioplasty**

A specially designed catheter with a tiny balloon is carefully guided through the artery to the blockage, then inflated to widen the opening and increase blood flow to the heart.

## <u>RVOT</u>

There are three ways to treat pulmonary conduit failure. One is a surgical conduit replacement, one is Transcatheter Pulmonary Valve Replacement, and the last is Balloon Valvuloplasty / Angioplasty.

### Surgical Replacement:

Surgical replacement of a pulmonary valve conduit involves a physician removing the narrow or leaking conduit and replacing it with an artificial valve.

### **Transcatheter Pulmonary Valve Replacement:**

An artificial valve is mounted on a thin hollow tube (catheter) with a balloon on the end, and is inserted into the artery in your upper leg. It is then advanced to the pulmonary conduit and the balloon inflated to place the new artificial pulmonary valve. The catheter is then removed from the body.

### **Balloon Valvuloplasty / Angioplasty:**

A thin hollow tube (catheter) with a balloon on the end is inserted into the artery in your upper leg and advanced to the pulmonary conduit. The balloon is then inflated to a specified pressure to open your conduit so that the blood will flow better. The catheter is then removed from the body.

### 7. Suggested profile and training for users

The Stent Device Family is intended for use by cardiology and surgical professionals undertaking stent implantation.