I. POLICY/Criteria

A. Carotid artery stenting

Priority Health will cover carotid artery stenting when all of the following are present:

1. Device is FDA approved for indications of use, AND
2. Patient must have a reference vessel diameter within the range of 4.0 mm and 9.0 mm at the target lesion, AND
3. Patient is high risk* for carotid endarterectomy (CEA), AND
4. Either of the following:
   a. Patient with neurological symptoms and a ≥ 70% stenosis of the common or internal carotid artery by ultrasound or ≥ 50% stenosis of the common or internal carotid artery by angiogram.
   OR
   b. Patient without neurological symptoms and a ≥ 70% stenosis of the common or internal carotid artery by ultrasound or ≥ 60% stenosis of the common or internal carotid artery by angiogram.

*Patients at high risk for CEA are defined as having significant comorbidities and/or anatomic risk factors (i.e., recurrent stenosis and/or previous radical neck dissection), and would be poor candidates for CEA. Significant comorbid conditions include but are not limited to:

- Congestive heart failure (CHF) class III/IV;
- Left ventricular ejection fraction (LVEF) < 30 %;
- Unstable angina;
- Contralateral carotid occlusion;
- Recent myocardial infarction (MI);
- Previous CEA with recurrent stenosis;
- Prior radiation treatment to the neck; and
- Other conditions that were used to determine patients at high risk for CEA in the prior carotid artery stenting trials and studies, such as ARCHER, CABERNET, SAPPHIRE, BEACH, and MAVERIC II.

B. Intracranial angioplasty and stenting, with or without stenting for the prophylaxis and treatment of atherosclerotic lesions, intracranial vasospasm,
or any other indication, is considered investigational and not a covered benefit.

C. Extra-cranial vertebral artery percutaneous transluminal angioplasty, with or without stent implantation and embolic protection, is considered medically necessary for persons with at least 50% stenosis of the vertebral artery who are symptomatic despite optimal medical treatment (e.g., antithrombotic agents, statins, and other risk factor modifications).

II. MEDICAL NECESSITY REVIEW

☐ Required ☑ Not Required ☐ Not Applicable

III. APPLICATION TO PRODUCTS

Coverage is subject to member’s specific benefits. Group specific policy will supersede this policy when applicable.

- HMO/EPO: This policy applies to insured HMO/EPO plans.
- POS: This policy applies to insured POS plans.
- PPO: This policy applies to insured PPO plans.
- ASO: For self-funded plans, consult individual plan documents. If there is a conflict between this policy and a self-funded plan document, the provisions of the plan document will govern.
- INDIVIDUAL: For individual policies, consult the individual insurance policy. If there is a conflict between this medical policy and the individual insurance policy document, the provisions of the individual insurance policy will govern.
- MEDICARE: Coverage is determined by the Centers for Medicare and Medicaid Services (CMS); if a coverage determination has not been adopted by CMS, this policy applies.
- MEDICAID/HEALTHY MICHIGAN PLAN: For Medicaid/Healthy Michigan Plan members, this policy will apply. Coverage is based on medical necessity criteria being met and the appropriate code(s) from the coding section of this policy including the Michigan Medicaid Fee Schedule located at: [http://www.michigan.gov/mdch/0,1607,7-132-2945_42542_42543_42546_42551-159815--00.html](http://www.michigan.gov/mdch/0,1607,7-132-2945_42542_42543_42546_42551-159815--00.html). If there is a discrepancy between this policy and the Michigan Medicaid Fee Schedule, the Michigan Medicaid Fee Schedule will govern. If there is a discrepancy or lack of guidance in the Michigan Medicaid Provider Manual, the Priority Health contract with Michigan Medicaid will govern. For Medical Supplies/DME/Prosthetics and Orthotics, please refer to the Michigan Medicaid Fee Schedule to verify coverage.

IV. DESCRIPTION

Carotid artery angioplasty with stenting (CAS) is an endovascular procedure that has been proposed as an alternative treatment to Carotid Endarterectomy (CEA) in high-risk patients and in patients with contraindications to surgery.
Percutaneous access to the carotid artery is gained through the femoral or brachial arteries. A balloon catheter is passed into the carotid artery under fluoroscopic guidance. Once the catheter has reached the area of blockage, the balloon is inflated to open the blocked artery. To keep the artery open, a semi-rigid tube-like device, or stent, is placed in the carotid artery. The increased dislocation of microemboli during CAS is thought to be the underlying cause for the increased risk of neurologic complications. This risk may be reduced with the use of embolic protection devices, intravessel filtration systems designed to retain the stenotic debris.

The Food and Drug Administration (FDA) approved the first carotid stent/companion carotid embolic protection device system in early September 2004 - The RX ACCULINK™ Carotid Stent System and RX ACCUNET™ Embolic Protection System. This device is available only to participants in a multicenter post-approval study sponsored in the practices of physicians at both academic and private hospitals, who will have a mixture of high, medium and low annual carotid stent implant volumes. The post-approval study will gather data on patient outcomes including death, stroke, myocardial infarction and rare adverse events.

**Background:**

Stroke is the third leading cause of death in the United States and the number one cause of disability in adults. In the United States, about 700,000 people have a new or recurrent stroke each year, and 280,000 will die. Cerebral infarctions account for about 80% to 85% of all strokes. Of all cerebral infarctions, about 20% to 30% are due to atherothrombosis or thromboembolism from the extracranial or intracranial vessels including the carotid arteries. Lifetime cost of stroke exceeds $90,000 per patient in the United States.

Treatment for carotid artery stenosis depends on the degree of blockage and the presence of symptoms. Asymptomatic patients with less than 70% stenosis are treated medically with antiplatelet therapy (e.g., aspirin) to decrease the likelihood of a blood clot and decrease the risk of stroke. Patients with severe symptomatic stenosis (≥ 70%) are referred for surgery. The procedure, carotid endarterectomy (CEA), involves the surgical removal of stenotic plaque from the carotid artery. A multicenter randomized controlled trial (NASCET; the North American Symptomatic Carotid Endarterectomy Trial) demonstrated that CEA significantly improved outcome and reduced the risk of stroke and death compared with medical treatment alone. However, CEA is associated with increased mortality and morbidity in patients with significant comorbidity (e.g., coronary artery disease).
Carotid artery angioplasty with stenting (CAS) is an endovascular procedure that has been proposed as an alternative treatment to CEA in these patients and in patients with contraindications to surgery. Percutaneous access to the carotid artery is gained through the femoral or brachial arteries. A balloon catheter is passed into the carotid artery under fluoroscopic guidance. Once the catheter has reached the area of blockage, the balloon is inflated to open the blocked artery. To keep the artery open, a semi-rigid tube-like device, or stent, is placed in the carotid artery. The proponents of this modality suggest that CAS may reduce the risk and side effects associated with general anesthesia. Furthermore, CAS may reduce the risk of complications for patients with significant comorbidities and may be an alternative for patients with contraindications to surgery or who have anatomically inaccessible lesions. However, early clinical studies reported stroke and death rates ranging from 10% to 12%. This rate did not compare favorably with the CEA stroke and death rate reported in the NASCET trial (5.8%), and raised concerns about the relative safety of the procedure. The increased dislocation of microemboli during CAS is thought to be the underlying cause for the increased risk of neurologic complications. This risk may be reduced with the use of embolic protection devices, intravessel filtration systems designed to retain the stenotic debris.

Guidance from the National Institute for Health and Clinical Excellence NICE (2011) concluded that "current evidence on the safety and efficacy of carotid artery stent placement for symptomatic extracranial carotid stenosis is adequate to support the use of this procedure provided that normal arrangements are in place for clinical governance and audit or research. During the consent process, clinicians should ensure that patients understand the risk of stroke and other complications associated with this procedure. Clinicians should also ensure that patients understand the reasons for advising carotid artery stent placement rather than endarterectomy in their particular case."

In this pivotal randomized trial, carotid endarterectomy (n=151) was no more effective than carotid artery angioplasty with stent placement performed with an embolic protection device (n=159) in reducing the incidence of major cardiovascular events among patients with severe carotid artery stenosis who also had at least one coexisting condition associated with increased risk for complications from endarterectomy. The Stenting and Angioplasty with Protection in Patients at High Risk for Endarterectomy (SAPPHIRE) trial enrolled symptomatic patients with 50% carotid artery stenosis or asymptomatic patients with 80% luminal diameter loss who were poor candidates for endarterectomy. However, only those patients judged by study investigators, both surgeons and interventional physicians, not to be at higher risk for stroke or death if treated with either one of the two procedures were eligible for randomization to undergo endarterectomy or stenting. Therefore, patients determined to be unsuitable for
endarterectomy but for whom stenting was considered feasible were excluded from randomization and enrolled in a stent registry-type study. Baseline characteristics were well balanced between treatment arms with a few exceptions. A significantly greater percentage of patients allocated to receive a stent, than those assigned to undergo endarterectomy, had coronary disease and a history of previous coronary angioplasty. The primary measure of efficacy was a composite endpoint, including 30-day cumulative incidence of death, stroke, or myocardial infarction (MI), or incidence of death or ipsilateral stroke between 31 and 365 days postprocedure. This outcome did not statistically differ between treatment groups in intention-to-treat analysis, and occurrence favored stenting in analysis of those actually treated despite significantly more patients with coronary disease and a history of angioplasty. In addition, target vessel revascularization rates, as well as incidence of major ipsilateral stroke within 1 year of treatment, were significantly lower in the stent versus endarterectomy group. The SAPPHIRE trial was not sufficiently powered to detect significant mortality differences between the two treatment arms. Due to low patient enrollment, the trial was ended early. Authors suggest that the advent of carotid stent registries made the procedure more readily available to patients and hampered recruitment for the randomized comparison with endarterectomy. It seems unlikely that future randomized head-to-head studies would have greater success with enrollment, and the data provided by SAPPHIRE may be among the best published evidence available. Findings support carotid stenting and endarterectomy equivalence in high-risk patients, even hinting that the less invasive treatment may have an edge over the standard surgical approach.

The Enroute Transcarotid Neuroprotection System (NPS) received 510(k) FDA clearance in February 2015. A compatible stent system, the Enroute Transcarotid Stent System (TSS), was FDA approved in May 2015. However, the Enroute Transcarotid NPS may be used with any FDA-approved carotid stent; a proprietary compatible stent system is not required.

The Enroute NPS is a single-use device intended to provide embolic protection during minimally invasive transcarotid angioplasty and stenting of the carotid artery by reversing blood flow away from the brain.

It protects the brain from plaque and debris dislodged during carotid artery stenting by reversing blood flow and capturing the material in its in-line filter. The Enroute system may provide an option for certain patients whose tortuous or diseased vasculature does not permit access via the groin.

V. CODING INFORMATION

ICD-10 Codes that may apply:
I63.031 - I63.039 Cerebral infarction due to thrombosis of posterior cerebral artery
I63.131 - I63.139  Cerebral infarction due to embolism of carotid artery
I63.231 - I63.239  Cerebral infarction due to unspecified occlusion or stenosis of carotid arteries
I63.59       Cerebral infarction due to unspecified occlusion or stenosis of other cerebral artery
I65.21 – I65.29 Occlusion and stenosis of carotid artery
I65.8         Occlusion and stenosis of other precerebral arteries

CPT/HCPCS Codes:
37215  Transcatheter Placement Of Intravascular Stent(S), Cervical Carotid Artery, Percutaneous; With Distal Embolic Protection
37216  Transcatheter Placement Of Intravascular Stent(S), Cervical Carotid Artery, Percutaneous; Wo Distal Embolic Protection *(Not covered for Priority Health Medicare or Medicaid)*
37217  Transcatheter placement of an intravascular stent(s), intrathoracic common carotid artery or innominate artery by retrograde treatment, via open ipsilateral cervical carotid artery exposure, including angioplasty, when performed, and radiological supervision and interpretation
37218  Transcatheter placement of intravascular stent(s), intrathoracic common carotid artery or innominate artery, open or percutaneous antegrade approach, including angioplasty, when performed, and radiological supervision and interpretation

0075T  Transcatheter placement of extracranial vertebral or intrathoracic carotid artery stent(s), including radiologic supervision and interpretation, percutaneous; initial vessel *(Not covered for Priority Health Medicaid)*
0076T  Transcatheter placement of extracranial vertebral or intrathoracic carotid artery stent(s), including radiologic supervision and interpretation, percutaneous; each additional vessel (List separately in addition to code for primary procedure) *(Not covered for Priority Health Medicaid)*

Not Covered:
61630  Balloon angioplasty, intracranial (e.g., atherosclerotic stenosis), percutaneous
61635  Transcatheter placement of intravascular stent(s), intracranial (e.g., atherosclerotic stenosis), including balloon angioplasty, if performed
61640  Balloon dilatation of intracranial vasospasm, percutaneous; initial vessel
61641  Balloon dilatation of intracranial vasospasm, percutaneous; each additional vessel in same vascular family (List separately in addition to code for primary procedure)
61642  Balloon dilatation of intracranial vasospasm, percutaneous; each additional vessel in different vascular family (List separately in addition to code for primary procedure)

Special Note: Priority Health’s Technology Assessment Committee reviewed Carotid Artery Stenting on December 3, 2004. This medical policy follows the recommendations of the committee.
VI. REFERENCES

“Carotid Artery Stenting for Carotid Artery Disease”, Hayes, Inc, March 2004
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