Protected PCI with the Impella® Heart Pump

Effective Hemodynamic Support in Elective and Urgent High-Risk PCI
What is Protected PCI?

A Protected PCI (percutaneous coronary intervention) is a minimally invasive cardiac procedure supported with an Impella heart pump. Impella support helps maintain hemodynamic stability and provides left ventricular unloading during the procedure, which may allow for a more complete revascularization in a single session, reducing long-term incidences of major adverse cardiac and cerebrovascular events (MACCE).

Growing Population Appropriate for PCI

Protected PCI makes complete revascularization a safe and effective option for your patients who are poor candidates for surgery or conventional PCI.

Which of Your Patients May Benefit from Protected PCI?

Impella heart pumps provide safe and effective hemodynamic support in elective and urgent high-risk PCI for a broad range of patients with complex coronary artery disease, hemodynamic compromise and comorbidities.

Patients benefit from this minimally invasive option that may:
- Enable more complete revascularization
- Reduce symptoms and class of heart failure and improve LVEF
- Significantly reduce the overall incidence of post-procedural acute kidney injury (AKI)
- Reduce number of days in the hospital
- Significantly reduce post-discharge MACCE events (composite of death/stroke/MI/repeat revascularization)

- Surgical Ineligibility
- Prior Cardiac Surgery
- Heart Failure
- Diabetes
- Advanced Age
- Unstable Angina/NSTEMI
- Renal Insufficiency
- Mild, Moderate, Severely Depressed Ejection Fraction
- High LVEDP

Growing Population Appropriate for PCI

- Multi-Vessel Disease
- Distal Left Main Disease
- Complex Lesions (Bifurcation, Calcification)
- CTO Retrograde
Surgical Turndown is Common and Associated with Higher Mortality

Protected PCI may be an option when surgery is not. Consider Protected PCI for patients who are considered too high-risk and therefore turned down for surgery, and for patients who decline surgical treatment.

Clinical Guidelines for Protected PCI with Impella Heart Pumps

The following clinical societies reinforce the benefit of using Impella for appropriate patient populations

2020 SCAI Position Statement on Optimal PCI Therapy for Complex Coronary Artery Disease
- MCS devices aim to improve the safety and efficacy of PCI in patients at very high risk for revascularization.
- Observational studies demonstrate improved procedural cardiovascular hemodynamics and more complete revascularization in the presence of MCS (mechanical circulatory support) devices despite higher-risk patient profiles.

2011 ACCF/AHA/SCAI Guideline for Percutaneous Coronary Intervention (J Am Coll Cardiol)
- High-risk patients: Class IIb
- CLASS III: HARM without hemodynamic support; for PCI at hospitals without on-site cardiac surgery

2014 AHA/ACC Guideline for the Management of Patients With Non–ST-Elevation Acute Coronary Syndromes (Circulation)
- Revascularization in Heart Failure: Class I
  - Revascularization strategy based on degree, severity, and extent of CAD; cardiac lesions; extent of LV dysfunction; prior revascularization
  - PVADs: Large amount of ischemic territory/poor LV function
Safety and Efficacy

Impella is the only FDA-approved, non-surgical heart pump proven safe and effective to assist the pumping function of the heart during stent placement and ensure blood flow is maintained to critical organs.

During a Protected PCI or protected stenting procedure, the Impella heart pump is placed temporarily in the heart to assist the pumping function while the physician performs the necessary treatment (angioplasty, stent placement, etc.).

Clinical Evidence

- Extensive data, including randomized controlled trial data and FDA-reviewed studies, support Impella® safety and efficacy.\textsuperscript{10,11,12}
- Impella 2.5\textsuperscript{®} and Impella CP\textsuperscript{®} maintain patient hemodynamics which may allow for more complete revascularization and reduction in major adverse coronary and cerebrovascular events (MACCE).\textsuperscript{8}
- Impella heart pumps are associated with reduction of kidney injury during high-risk PCI.\textsuperscript{4}
- PROTECT III, the most contemporary data, continues to show a reduction in MACCE with Impella\textsuperscript{11}
Benefits of Impella

**Complete Revascularization Associated with Improved Outcomes**

**Benefits in high-risk acute coronary syndrome (ACS) patients at 1 year**

- **71%** risk reduction in all-cause death
- **27%** risk reduction in major adverse cardiovascular events (MACE)
- **41%** risk reduction in myocardial infarction (MI)

**Benefits in high-risk STEMI patients at 3 years**

- **26%** risk reduction in cardiovascular (CV) Death/MI
- **47%** risk reduction in CV Death/MI/Revascularization

**Meta-analysis of benefits in 11 high-risk STEMI studies**

- **29%** risk reduction in CV Death/MI

**Resources**

For a detailed list of resources on the Impella Heart Pumps, please scan the QR code or visit the link below.

[www.heartrecovery.com/resources/downloads](http://www.heartrecovery.com/resources/downloads)

**Fewer Days in the Hospital**

Fewer post-hospitalization adverse events within 90 days after your PCI procedure compared to traditional therapy such as the intra-aortic balloon pump.

**Fewer Post-Hospitalization Adverse Events**

Meta-analysis of benefits in 11 high-risk STEMI studies.

**Fewer Repeat Visits**

Fewer repeat visits to the hospital for heart-related issues than traditional therapy such as the intra-aortic balloon pump.
**Improve Your Patient Outcomes**

Physicians who implement best practices and protocols find significantly improved in-hospital safety in high-risk PCI patients with fewer complications when using the Impella heart pump.

Data from PROTECT III Study demonstrates that evolution and adoption of best practices leads to reduced rates of death, stroke, myocardial infarction and repeat procedures (MACCE) when Impella is used to support high-risk PCI patients.

**Enhance Quality of Life**

**Statistics Show That:**

- **45%** of PCI patients have incomplete revascularization\(^\text{13-16}\)
- **14%** of PCI patients are “staged” and not all staged patients return for a second procedure, leading to an incomplete revascularization\(^\text{17}\)

**8 out of 10 Patients** Experienced Improved Ejection Fraction\(^\text{12}\) or NYHA Heart Failure Class\(^\text{12}\) After a Protected PCI procedure with the Impella Heart Pump.
How Protected PCI with Impella Works

The Impella device is inserted through a small incision and advanced through the arteries and into the heart. Once in place, the Impella heart pump is turned on, pulling blood from the left ventricle and releasing it into the aorta. This active ‘unloading’ of the left ventricle increases blood flow to the brain and other vital organs and may protect the kidneys from acute injury.

When the heart is strong enough to pump on its own and the Impella heart pump is no longer needed, the device will be removed (before the patient goes home from the hospital).

The Insertion Procedure

STEP 1
Access common femoral artery

STEP 2
Pre-dilate, insert a 14 Fr sheath into the common femoral artery

STEP 3
Advance a guidewire with a diagnostic catheter across the aortic valve

STEP 4
Prepare the Impella pump to guide it along the wire

STEP 5
Insert and place across the aortic valve and confirm position placement

STEP 6
Removal of the guidewire and Impella device is started by pulling blood from the left ventricle into the aorta
Protected PCI with Impella

Advancing the way heart failure is treated

To learn more about Protected PCI and how Impella may be an option for your patients visit:

www.HeartRecovery.com

INDICATIONS FOR USE

High-Risk PCI

The Impella 2.5®, Impella CP® and Impella CP® with SmartAssist® Systems are temporary (≤ 6 hours) ventricular support devices indicated for use during high-risk percutaneous coronary interventions (PCI) performed in elective or urgent, hemodynamically stable patients with severe coronary artery disease, when a heart team, including a cardiac surgeon, has determined high-risk PCI is the appropriate therapeutic option. Use of the Impella 2.5, Impella CP, and Impella CP with SmartAssist Systems in these patients may prevent hemodynamic instability, which can result from repeat episodes of reversible myocardial ischemia that occur during planned temporary coronary occlusions and may reduce peri- and post-procedural adverse events.

Cardiogenic Shock

The Impella 2.5®, Impella CP®, Impella CP® with SmartAssist®, Impella 5.0®, Impella 5.5® with SmartAssist® and Impella LD® Catheters, in conjunction with the Automated Impella Controller™ (collectively, “Impella® System Therapy”), are temporary ventricular support devices intended for short term use (≤ 4 days for the Impella 2.5, Impella CP, and Impella CP with SmartAssist, and ≤ 14 days for the Impella 5.0, Impella 5.5 with SmartAssist and Impella LD) and indicated for the treatment of ongoing cardiogenic shock that occurs immediately (< 48 hours) following acute myocardial infarction or open heart surgery or in the setting of cardiomyopathy, including peripartum cardiomyopathy, or myocarditis as a result of isolated left ventricular failure that is not responsive to optimal medical management and conventional treatment measures (including volume loading and use of pressors and inotropes, with or without IABP). The intent of Impella System Therapy is to reduce ventricular work and to provide the circulatory support necessary to allow heart recovery and early assessment of myocardial function.

CONTRAINdications:
The Impella 2.5, Impella CP, Impella CP with SmartAssist, Impella 5.0, Impella 5.5 with SmartAssist and Impella LD are contraindicated for use with patients experiencing any of the following conditions: Mural thrombus in the left ventricle; Presence of a mechanical aortic valve or heart constrictive device; Aortic valve stenosis/calcification (equivalent to an orifice area of 0.6 cm² or less); Moderate to severe aortic insufficiency (echocardiographic assessment graded as ≥ +2); Severe peripheral arterial disease precluding placement of the Impella System; Significant right heart failure*; Combined cardiopulmonary failure*; Presence of an Atrial or Ventricular Septal Defect (including post-infarct VSD)*; Left ventricular rupture*; Cardiac tamponade* *This condition is a contraindication for the cardiogenic shock indication only.

POtential Adverse Events:

Acute renal dysfunction, Aortic valve injury, Bleeding, Cardiogenic shock, Cerebral vascular accident/Stroke, Death, Hemolysis, Limb ischemia, Myocardial infarction, Renal failure, Thrombocytopenia and Vascular injury

In addition to the risks above, there are other WARNINGS and PRECAUTIONS associated with Impella devices.

Visit http://www.abiomed.com/important-safety-information to learn more.