Edwards
Transcatheter Aortic Valve Replacement (TAVR)

A guide for patients with severe aortic stenosis
This booklet is not intended to explain everything you need to know about your treatment options for aortic stenosis or about the TAVR procedure. Please discuss any questions you have with your doctor. Only a TAVR Heart Team can decide which treatment option is right for you.

This patient booklet is for those who are suffering from severe aortic stenosis and need treatment.

The information in this booklet will help you understand more about your heart, aortic stenosis, and a less invasive procedure called transcatheter aortic valve replacement (TAVR).

Be sure to ask your TAVR Heart Team to explain all of your treatment options and the possible risks and benefits of each.

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There are two problems that can occur in heart valves:

**Stenosis:** when your valve narrows and does not open completely

**Regurgitation:** when your valve does not close completely and blood can leak backwards

It is important that your valves are always working properly. Your valves should:

- Be properly formed and flexible
- Open all the way so that the right amount of blood can pass through
- Close tightly so that no blood leaks back into the chamber

Aortic stenosis is a progressive disease meaning that over time, the leaflets become stiff. This reduces their ability to fully open and close. When the leaflets don't fully open, your heart must work harder to push blood through the aortic valve to your body. As a result, less oxygen-rich blood flows from the lungs to the brain and the rest of the body, which may cause symptoms.

**Causes:**
- Age
- Calcium build-up
- Radiation therapy
- Infection of the heart
- Birth defects
- Rheumatic fever

**What Are the Symptoms of Aortic Stenosis?**
The symptoms of aortic stenosis are commonly misunderstood by patients as “normal” signs of aging.

**Symptoms:**
- Chest pain
- Rapid, fluttering heartbeat
- Trouble breathing or feeling short of breath
- Feeling dizzy or light-headed even fainting
- Difficulty walking short distances
- Swollen ankles or feet
- Not doing activities you used to enjoy
- Difficulty sleeping or the need to sleep sitting up

It’s important to know that heart valve disease may occur with no outward symptoms.
Understanding Your Treatment Options For Severe Aortic Stenosis

If you have been diagnosed with severe symptomatic aortic stenosis, TAVR is an option for you. Only a TAVR Heart Team can tell you if TAVR is right for you.

Medication
Your doctor may prescribe certain medications to help ease some of the symptoms of aortic stenosis; however, it will not cure or fix the valve.

Surgical Aortic Valve Replacement (SAVR)
Open heart surgery for aortic valve replacement is where the doctor will open your chest and will completely remove the damaged valve and replace it with an artificial valve. You will be connected to a heart-lung machine that does the work of your heart and keeps the blood flowing throughout your body. Patients usually need to stay in the hospital for a week or more, before beginning a long period of recovery.

Transcatheter Aortic Valve Replacement (TAVR)
TAVR is a less invasive approach to aortic valve replacement compared to open heart surgery. With the TAVR procedure, the doctor will make a small cut, usually in your groin. A thin, flexible tube is inserted into the artery to guide the heart valve up to your heart, and the valve is expanded into place. It does not remove your old valve, it fits within the diseased valve.

What Is the Best Treatment Option for You?
Seeing a specialized doctor on a TAVR Team will ensure you will be evaluated for all treatment options. They will consider all factors about your health to decide the best treatment option for you.
Your doctor will consider these factors:
• Your medical history
• Your age
• Your current health status
• Your ability to undergo the procedure and recover from it
• The overall condition of your heart

Quality of Life Improvement:
Quality of life studies with the Edwards SAPIEN 3 TAVR® have shown patient health improvements within 30 days, including the ability to take care of themselves and participate in everyday activities.

What Are the Benefits of Transcatheter Aortic Valve Replacement?
If you have severe aortic stenosis and need your valve replaced, transcatheter valve replacement may help your heart to work better. Other benefits may include:
• Better clinical outcomes
• Less invasive, with minimal scarring
• Shorter hospital stay
• Shorter recovery time to getting back to everyday activities
• Less pain and anxiety
• Improved quality of life
• Relief of symptoms

Your doctor will tell you which valve you will receive.
The Edwards SAPIEN 3 and SAPIEN 3 Ultra Valves

The Edwards SAPIEN 3 and SAPIEN 3 Ultra valves are bioprosthetic, balloon-expandable valves. The frame of the valves is made from cobalt chromium to help with strength and durability. The leaflets in the valves are made from the same bovine pericardial tissue (from a cow’s heart) as Edwards surgical valves. An outer sealing skirt surrounds the bottom of the valve, to help stop any possible leakage around the valve.

The valves are available in four sizes: 20, 23, 26, and 29 mm in diameter. Your TAVR Heart Team will determine which valve and which size is right for you.

The Edwards SAPIEN 3 Transcatheter Heart Valves

Edwards Lifesciences transcatheter heart valves are designed to work like your native heart valve. The Edwards SAPIEN 3 valves are expanded into place with the help of a balloon, and begin working immediately when they are implanted.

Your doctor may refer to your heart valve by a few different names including SAPIEN 3 or SAPIEN 3 Ultra valve. Your doctor can help you decide which Edwards TAVR heart valve is right for you.

TAVR is a less invasive, technique that uses a catheter to replace your diseased aortic valve. An interventional cardiologist (specializes in catheter procedures), along with a cardiothoracic surgeon (specializes in surgical procedures of the heart), will work together during the procedure. They will guide a new valve into the heart while the heart is still beating, using guidance from X-ray and echocardiography.

TAVR by Edwards Lifesciences

The Edwards SAPIEN transcatheter heart valve was the first of its kind to get FDA approval in the United States for patients who were too sick to undergo open heart surgery. The first transcatheter heart valve from Edwards was approved commercially in Europe in 2007 and in the United States in 2011. To date, Edwards transcatheter heart valves have treated hundreds of thousands of patients in over 70 countries around the world. This era represents a major achievement in the treatment of patients with aortic stenosis.

*The SAPIEN 3 and SAPIEN 3 Ultra valves are both commercially available in the United States. Your doctor will tell you which valve you will receive.
The Edwards SAPIEN 3 TAVR Procedure

What Do You Need to Do Before the Procedure?
Be sure to talk with your TAVR Heart Team about any medication you may be taking. Your doctor may tell you to stop taking certain medication up to one week before the procedure. You should plan on making arrangements for a ride to and from the hospital, and arrange for help at home after the procedure.

Steps of the TAVR Procedure
TAVR allows a new valve to be inserted through a catheter.

1. Before your procedure, you may be placed either under general anesthesia (sleep medicine) or conscious sedation (medicine that helps you relax and block pain but you will remain awake).

2. A small cut will be made where your doctor will insert a short, hollow tube called a sheath.

3. Your new valve will be placed on the delivery system tube and squeezed on the balloon to make it small enough to fit through the sheath.

4. The delivery system carrying the valve will be inflated, expanding the new valve within your diseased valve. The new valve will push the leaflets of your diseased valve aside. The frame of the new valve is strong and it will use the leaflets of your diseased valve to secure itself in place.

5. Your doctor will make sure your new valve is working properly.

On average, the TAVR procedure lasts about 1 hour, compared to 4 hours with open heart surgery.

Watch a video on the Edwards TAVR procedure at NewHeartValve.com/video
NewHeartValve.com
What Happens After the TAVR Procedure?
After your procedure, you may spend a day or two in the hospital. Every patient is different in how they recover. Most patients should begin walking very soon after their Edwards TAVR procedure. Before you leave the hospital, your doctor will discuss your aftercare plan with you. They will give you specific instructions to help you with your recovery. This may include a special diet, when to return to exercise, and any medicine you may need to take.

It is important to carefully follow your doctor’s directions, especially if you need to take any blood thinning medication.

TAVR Follow-Up Visits
Regular check-ups with your doctor are very important. You will probably be asked to return to see your TAVR doctor to have your heart valve checked at 30 days and 1 year after your procedure. However, call or see your doctor whenever you have questions or concerns about your health.

Average length of stay with the TAVR procedure: 3 days¹ compared to 7 days with open heart surgery

Your Edwards TAVR Implant Card
As you leave the hospital, your valve clinic coordinator or nurse should give you a temporary implant card. A permanent card will be sent to you in approximately 6-8 weeks. This card has information about your Edwards TAVR heart valve. Share this card with all members of your healthcare team, including your dentist. It is important to share about your heart valve replacement before any medical, dental, or MRI (magnetic resonance imaging) procedures. If you need an MRI, tell your doctor that you have an Edwards TAVR heart valve.

1 Based on PARTNER 3 Low Risk Trial Outcomes
What Are the Risks of Edwards SAPIEN 3 TAVR?

As with any medical procedure, there is a possibility of risks. The Edwards TAVR procedure’s most serious risks are:

- Death
- Stroke
- Serious damage to the arteries
- Serious bleeding

The Edwards SAPIEN 3 TAVR Cannot Be Used for People Who:

- Cannot take blood thinning medications
- Have an active infection in the heart or elsewhere

Additional potential risks associated with the procedure include:

- Heart attack
- Failure of your heart to pump enough blood to the body’s organs
- Irregular heart rate
- Problems with the electrical pathway of your heart that requires a pacemaker
- Collection of fluid or blood around your heart
- Having an abnormal particle (air or blood clots) floating in the bloodstream or attached to an object, including the valve
- Infection in your heart, blood, or other areas
- Injury to your blood vessels or heart that requires treatment
- Blocking, narrowing, or bulging of a blood vessel
- Blood clot, including a blood clot on the valve
- Trouble or inability to breathe
- Fluid buildup in your lungs
- Anemia
- Lab values that are not normal
- Abnormally high or low blood pressure
- Pain, inflammation, or fever
- Pain or changes at the incision site
- Problems with the valve or accessories that do not allow it to work well, including but not limited to, wear, tear, or movement forward (prolapse) or backward (retraction) from the normal position of the valve leaflets, calcium buildup on the leaflets, or a break in the frame
- Incorrect position of valve or valve movement
- Blood leak around the valve
- Additional cardiac surgery, vascular surgery, or intervention, including removal of the transcatheter heart valve
- Fainting or dizziness
- Weakness or trouble exercising
- Allergic reaction
- Inability to move (paralysis)
- Permanent disability
- Kidney failure
- Chest pain
- Damage to blood cells
- Repeat hospitalization
- Sudden or unexpected loss of heart function
- Injury to nerve
- Partial or complete blockage of coronary artery (artery supplying blood to the heart)
- Extra or unusual sound during heartbeat (heart murmur)
Warnings

- X-ray used during the procedure may cause radiation injury to the skin.
- Younger patients, or patients with a disease that results in more calcium in their blood, may have early wear of their valve.
- Talk to your doctor if you are allergic to any of the following materials: anesthesia, contrast media, chromium, nickel, molybdenum, manganese, copper, silicon, and plastics.

Precautions

- TAVR patients should stay on blood-thinning medication and/or aspirin as recommended by their doctor. Patients who do not may be at increased risk of a stroke. Blood-thinning medication may increase the risk of bleeding in the brain (stroke).
- Patients who need a dental procedure should talk to their doctor about risk of infection and needing antibiotics.
- The safety of the transcatheter heart valve is not known for patients who have:
  - A heart that does not pump properly
  - An enlarged heart
- The Edwards TAVR valve has not been studied in patients:
  - Who have an aortic heart valve that has NO build-up of calcium
  - Who only have one leaflet (unicuspid) in their aortic valve
  - Who are low surgical risk and have two leaflets (congenital bicuspid) in their aortic valve
  - Who have a prosthetic ring in any heart valve
  - Who have a low white or red blood cell count, or other irregularities in the blood
  - Who have unusual ultrasound images of the heart that show possible irregularities, such as a blood clot
  - Who have allergies to blood-thinning medications
  - Who are allergic to dye that is injected during the procedure
  - Whose diseased aortic valve is too small or too big to fit the transcatheter heart valve
  - Whose femoral arteries in the legs are too diseased or too small for the delivery device
  - Whose aortic valve leaflets have large pieces of calcium that may block the arteries that supply blood to the heart

How long your tissue valve will last depends on many patient factors and medical conditions. Follow all care instructions to ensure the best possible results. The Edwards transcatheter valve has been tested in a laboratory to mimic 5 years of use without failure. Regular follow-ups will help your doctor know how your valve is working.
The PARTNER 3 Low-Risk Trial

The risks with the procedure may depend on the overall health of the patient. If you are at low-risk for open heart surgery, the clinical data shown in these charts could be what you would expect.

As part of the PARTNER 3 Trial, the SAPIEN 3 valve was studied in patients at low risk for open heart surgery. The trial enrolled about 1,000 patients, mostly in the United States. Patients were randomly chosen for either TAVR with an Edwards valve or open heart surgery (SAVR). Patients were examined at 30 days and 1 year after the procedure and will continue to be followed every year for 10 years.

### Edwards TAVR Clinical Data for Low-Risk Patients

#### Low-Risk Clinical Data – TAVR

<table>
<thead>
<tr>
<th>Event</th>
<th>Risk Within 30 Days</th>
<th>Risk Within 1 Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>Death From Any Cause</td>
<td>1 out of 100</td>
<td>1 out of 100</td>
</tr>
<tr>
<td>Death From Heart Related Cause</td>
<td>1 out of 100</td>
<td>1 out of 100</td>
</tr>
<tr>
<td>Disabling Stroke</td>
<td>0 out of 100</td>
<td>1 out of 100</td>
</tr>
<tr>
<td>New Permanent Pacemaker</td>
<td>7 out of 100</td>
<td>8 out of 100</td>
</tr>
<tr>
<td>Life-Threatening or Disabling Bleeding</td>
<td>2 out of 100</td>
<td>3 out of 100</td>
</tr>
<tr>
<td>Major Vascular Complications</td>
<td>3 out of 100</td>
<td>3 out of 100</td>
</tr>
<tr>
<td>Heart Attack (Myocardial Infarction)</td>
<td>1 out of 100</td>
<td>2 out of 100</td>
</tr>
</tbody>
</table>

The frequency is shown as the number of patients out of every 100.

### Low-Risk Clinical Data – Open Heart Surgery

#### SAVR Patients

<table>
<thead>
<tr>
<th>Event</th>
<th>Risk Within 30 Days</th>
<th>Risk Within 1 Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>Death From Any Cause</td>
<td>2 out of 100</td>
<td>3 out of 100</td>
</tr>
<tr>
<td>Death From Heart Related Cause</td>
<td>1 out of 100</td>
<td>2 out of 100</td>
</tr>
<tr>
<td>Disabling Stroke</td>
<td>1 out of 100</td>
<td>1 out of 100</td>
</tr>
<tr>
<td>New Permanent Pacemaker</td>
<td>4 out of 100</td>
<td>6 out of 100</td>
</tr>
<tr>
<td>Life-Threatening or Disabling Bleeding</td>
<td>12 out of 100</td>
<td>13 out of 100</td>
</tr>
<tr>
<td>Major Vascular Complications</td>
<td>2 out of 100</td>
<td>2 out of 100</td>
</tr>
<tr>
<td>Heart Attack (Myocardial Infarction)</td>
<td>2 out of 100</td>
<td>3 out of 100</td>
</tr>
</tbody>
</table>

The frequency is shown as the number of patients out of every 100.

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18
The risks with the TAVR procedure may depend on the overall health of the patient. If you are at intermediate risk for open heart surgery, the clinical data shown in these charts could be what you would expect.

As a single-arm study, the SAPIEN 3 Ultra Valve (the latest TAVR valve available from Edwards) was studied in patients at intermediate risk for open heart surgery. The study enrolled about 40 patients in Canada and the United Kingdom, who had severe aortic stenosis. This chart summarizes 30-day data of a study with 40 patients to confirm the procedural safety and performance of the SAPIEN 3 Ultra valve in treating patients with severe aortic stenosis who are at intermediate operative risk for open heart surgery. The results from this trial confirmed the safety and effectiveness of the design updates of the SAPIEN 3 Ultra system compared to its previous generation of the SAPIEN 3 valve (see clinical data on page 20).

Intermediate-Risk Clinical Data

<table>
<thead>
<tr>
<th>SAPIEN 3 Ultra Valve</th>
<th>Risk at Discharge From Hospital</th>
<th>Risk Within 30 Days</th>
</tr>
</thead>
<tbody>
<tr>
<td>Death From Any Cause</td>
<td>0 out of 100</td>
<td>0 out of 100</td>
</tr>
<tr>
<td>Death From Heart Related Cause</td>
<td>0 out of 100</td>
<td>0 out of 100</td>
</tr>
<tr>
<td>Disabling Stroke</td>
<td>0 out of 100</td>
<td>0 out of 100</td>
</tr>
<tr>
<td>New Permanent Pacemaker</td>
<td>18 out of 100</td>
<td>18 out of 100</td>
</tr>
<tr>
<td>Major Vascular Complications</td>
<td>0 out of 100</td>
<td>0 out of 100</td>
</tr>
<tr>
<td>Heart Attack (Myocardial Infarction)</td>
<td>0 out of 100</td>
<td>0 out of 100</td>
</tr>
</tbody>
</table>

The frequency is shown as the number of patients out of every 100.

Intermediate-Risk Clinical Outcomes

<table>
<thead>
<tr>
<th>SAPIEN 3 Valve</th>
<th>Risk Within 30 Days</th>
<th>Risk Within 1 Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>Death From Any Cause</td>
<td>2 out of 100</td>
<td>8 out of 100</td>
</tr>
<tr>
<td>Death From Heart Related Cause</td>
<td>1 out of 100</td>
<td>5 out of 100</td>
</tr>
<tr>
<td>Disabling Stroke</td>
<td>1 out of 100</td>
<td>3 out of 100</td>
</tr>
<tr>
<td>New Permanent Pacemaker</td>
<td>11 out of 100</td>
<td>N/A</td>
</tr>
<tr>
<td>Major Vascular Complications</td>
<td>7 out of 100</td>
<td>N/A</td>
</tr>
<tr>
<td>Heart Attack (Myocardial Infarction)</td>
<td>1 out of 100</td>
<td>N/A</td>
</tr>
</tbody>
</table>

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The PARTNER II Intermediate-Risk Trial

The risks with the TAVR procedure may depend on the overall health of the patient. If you are at intermediate risk for open heart surgery, the clinical data shown in this chart could be what you would expect.

As part of the PARTNER II Trial, the SAPIEN 3 valve was studied in patients at intermediate risk for open heart surgery. The trial enrolled about 1,000 patients, mostly in the United States. The outcomes in this trial were compared to those patients who participated in another trial and were treated with surgery. Patients were examined at 30 days and 1 year after the procedure and will continue to be followed every year for 10 years.
The PARTNER II High-Risk/Inoperable Study

The risks with the TAVR procedure may depend on the overall health of the patient.

If you are at high risk or cannot have open heart surgery, the clinical data shown in this chart could be what you would expect.

The SAPIEN 3 valve was studied in approximately 600 US patients that were either high risk or too sick for open heart surgery.

Patients were examined at 30 days and 1 year after the procedure.

All data presented in charts are the transfemoral approach population only.

<table>
<thead>
<tr>
<th>High-Risk and Inoperable Clinical Data</th>
<th></th>
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</thead>
<tbody>
<tr>
<td><strong>TAVR</strong></td>
<td><strong>Risk Within 30 Days From TAVR</strong></td>
</tr>
<tr>
<td>Death From Any Cause</td>
<td>2 out of 100</td>
</tr>
<tr>
<td>Death From Heart Related Cause</td>
<td>1 out of 100</td>
</tr>
<tr>
<td>All Stroke</td>
<td>2 out of 100</td>
</tr>
<tr>
<td>New Permanent Pacemaker</td>
<td>14 out of 100</td>
</tr>
<tr>
<td>Life-Threatening or Disabling Bleeding</td>
<td>6 out of 100</td>
</tr>
<tr>
<td>Major Vascular Complications</td>
<td>6 out of 100</td>
</tr>
<tr>
<td>Heart Attack (Myocardial Infarction)</td>
<td>1 out of 100</td>
</tr>
</tbody>
</table>

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TVT Registry* Data With TAVR for Bicuspid Patients

If you have been diagnosed with having a bicuspid aortic valve, your aortic valve has two valve leaflets as opposed to three. This is a rare hereditary abnormality that occurs in 0.5% to 2% of the general population. Aortic stenosis is a frequent complication in this population and may occur at a younger age in patients with a bicuspid valve, when compared to patients with a tricuspid aortic valve.

The TVT Registry collects safety and efficacy data from hospitals in the US treating patients with TAVR. The registry showed results of 545 patients with bicuspid aortic stenosis who received a SAPIEN 3 valve.

Over 90% of these patients were either at high risk or ineligible for open heart surgery.

<table>
<thead>
<tr>
<th>Bicuspid Aortic Valve Clinical Data</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Risk Within 30 Days From TAVR</strong></td>
<td><strong>Risk Within 1 Year From TAVR</strong></td>
</tr>
<tr>
<td>Death From Any Cause</td>
<td>3 out of 100</td>
</tr>
<tr>
<td>Death From Heart Related Cause</td>
<td>2 out of 100</td>
</tr>
<tr>
<td>All Stroke</td>
<td>2 out of 100</td>
</tr>
<tr>
<td>Aortic Valve Reintervention or Reoperation</td>
<td>1 out of 100</td>
</tr>
</tbody>
</table>

The frequency is shown as the number of patients out of every 100.

*The STS/ACC TVT RegistryTM is the main repository for clinical data related to transcatheter aortic valve replacement (TAVR). The Registry, created by a collaboration between STS and the American College of Cardiology, monitors patient safety and real-world outcomes related to TAVR.
The SAPIEN 3 and SAPIEN 3 Ultra valve are both commercially available in the United States. Your doctor will tell you which valve you will receive.

CAUTION: Federal (United States) law restricts these devices to sale by or on the order of a physician. See Instructions for Use for full prescribing information, including indications, contraindications, warnings, precautions, and adverse events.

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