Review article

Heart Transplant and Mechanical Circulatory Support in Patients With Advanced Heart Failure

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ABSTRACT

Patients with advanced heart failure have a poor prognosis and heart transplant is still the best treatment option. However, the scarcity of donors, long waiting times, and an increasing number of unstable patients have favored the development of mechanical circulatory support. This review summarizes the indications for heart transplant, candidate evaluation, current immunosuppression strategies, the evaluation and treatment of rejection, infectious prophylaxis, and short and long-term outcomes. Regarding mechanical circulatory support, we distinguish between short- and long-term support and the distinct strategies that can be used: bridge to decision, recovery, candidacy, transplant, and destination therapy. We then discuss indications, risk assessment, management of complications, especially with long-term support, and outcomes. Finally, we discuss future challenges and how the widespread use of long-term support for patients with advanced heart failure will only be viable if their complications and costs are reduced.

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Trasplante cardiaco y soporte circulatorio mecánico para pacientes con insuficiencia cardiaca avanzada

RESUMEN

Los pacientes con insuficiencia cardiaca avanzada tienen mal pronóstico y el trasplante cardiaco es actualmente la mejor opción de tratamiento disponible. Sin embargo, la escasez de donantes, los largos tiempos de espera y un número creciente de pacientes inestables han favorecido el desarrollo del soporte circulatorio mecánico. Esta revisión resume las indicaciones del trasplante cardiaco, cómo evaluar a los posibles candidatos, las estrategias actuales de inmunosupresión, cómo evaluar y tratar el rechazo, la profilaxis infecciosa y los resultados a corto y largo plazo. Respecto al soporte circulatorio mecánico, se diferencia entre las asistencias ventriculares de corto y largo plazo, así como las diferentes estrategias disponibles: puente hasta la decisión, recuperación, candidatura, trasplante y terapia de destino. Posteriormente se resumen las indicaciones, la valoración del riesgo previo al implante, el manejo de las complicaciones, especialmente de las asistencias de largo plazo y los resultados. Finalmente se plantean los retos futuros y cómo el uso generalizado de las asistencias ventriculares de largo plazo para pacientes con insuficiencia cardiaca avanzada solo será viable si se reducen sus complicaciones y costes.

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Abbreviations

BTT: bridge to transplant
DT: destination therapy
HF: heart failure
HT: heart transplant
LTVAD: long-term ventricular assist device
LVAD: left ventricular assist device
MCS: mechanical circulatory support
STVAD: short-term ventricular assist device

INTRODUCTION: STATUS OF HEART FAILURE

Heart failure (HF) is a clinical syndrome caused by reduced cardiac output and/or elevated intracardiac pressures. Its prevalence is 1% to 2% of adults in developed countries, constituting a major health problem with a high economic burden. Despite the availability of disease-modifying drugs and implantable device therapy, prognosis remains poor.

Approximately 5% of patients are in advanced HF as defined in Table 1. In a very small proportion of these patients, heart transplant (HT) is the only available treatment. Unfortunately, donors are limited, resulting in 250 to 300 HT per year in Spain and 2000 in the United States. Long waiting times and an increasing number of unstable patients have favored the development of mechanical circulatory support (MCS) as bridge to recovery, bridge to transplant (BTT), and bridge to candidacy or decision; initially with short-term ventricular assist devices (STVADs), and in the last decades with long-term ventricular assist devices (LTVADs). The development of LTVADs has created the possibility of destination therapy (DT) in patients who are not candidates for HT (Table 2).

HEART TRANSPLANT

HT is the gold standard for the treatment of end-stage HF because it improves survival, functional status, and quality of life.

Indications

Although HT is the best option, it carries a mortality of approximately 15% in the first year. Therefore, assessing prognosis in patients with advanced HF is mandatory. The most often used scores are the Heart Failure Survival Score and the Seattle Heart Failure Model. A high- to medium-risk range in the first score or an estimated 1-year survival < 80% by the second are cutoff points for listing for HT. The BCNbioHF calculator provides prognostic information derived from clinical parameters but also incorporates biomarkers.

Functional status evaluated with the cardiopulmonary exercise test is frequently used to determine HT eligibility. A peak oxygen consumption of <14 mL/kg/min or <12 mL/kg/min in patients on β-blockers at maximal exertion has been established as the cutoff point for HT. If exercise is submaximal, a ventilation to carbon dioxide slope of >35 also has prognostic value. A 6-minute walk test <300 meters also indicates high risk.

In patients with hemodynamic instability, HT may be performed emergently, preceded or not by MCS. To stratify patients in advanced HF, the Interagency Registry for Mechanical Assisted
Table 3

<table>
<thead>
<tr>
<th>Profile description</th>
<th>Features</th>
<th>Type of MCS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Critical cardiogenic shock (“crash and burn”)</td>
<td>Life-threatening hypotension and rapidly escalating inotropic/pressor support, with critical organ hypoperfusion often confirmed by worsening acidosis and lactate levels</td>
<td>Short-term VAD or VA-ECMO</td>
</tr>
<tr>
<td>Progressive decline (“sliding fast” on inotropes)</td>
<td>Patient with declining function despite intravenous inotropic support, which may be manifest by worsening renal function, nutritional depletion, and inability to restore volume balance. Also describes declining status in patients unable to tolerate inotropic therapy</td>
<td>Short-term VAD or LT VAD</td>
</tr>
<tr>
<td>Stable but inotrope-dependent</td>
<td>Patient with stable blood pressure, organ function, nutrition, and symptoms on continuous intravenous inotropic support (or a temporary circulatory support device or both), but demonstrating repeated failure to wean from support due to recurrent symptomatic hypotension or renal dysfunction. “Dependent stability”</td>
<td>LT VAD</td>
</tr>
<tr>
<td>Resting symptoms on oral therapy at home (“frequent flyer”)</td>
<td>Patient can be stabilized close to normal volume status but experiences daily symptoms of congestion. Diuretic doses generally fluctuate at very high levels. More intensive management and surveillance strategies should be considered, which may in some cases reveal poor adherence that would compromise outcomes with any therapy. Some patients may struggle between 4 and 5</td>
<td>LT VAD</td>
</tr>
<tr>
<td>Exertion intolerant (“housebound”)</td>
<td>Patient who is comfortable at rest but unable to engage in any activity, living predominantly within the house or housebound</td>
<td>LT VAD could be considered</td>
</tr>
<tr>
<td>Exertion limited (“walking wounded”)</td>
<td>Patient who is comfortable at rest without evidence of fluid overload but who is able to undertake some mild activity. Activities of daily living are comfortable and minor activities outside the home such as visiting friends or going to a restaurant can be performed, but fatigue results within a few minutes or with any meaningful physical exertion</td>
<td>LT VAD could be considered</td>
</tr>
<tr>
<td>“Placeholder” Advanced NYHA class III</td>
<td>Patient who is clinically stable with a reasonable level of comfortable activity, despite a history of previous decompensation that is not recent. This patient is usually able to walk more than a block. Any decompensation requiring intravenous diuretics or hospitalization within the previous month should make this person a patient profile 6</td>
<td>LT VAD not considered</td>
</tr>
</tbody>
</table>

LT VAD, long-term ventricular assist device; MCS, mechanical circulatory support; NYHA, New York Heart Association; VAD, ventricular assist device; VA-ECMO, venoarterial extracorporeal membrane oxygenation.

Adapted from Ponikowski et al. and Stevenson et al., with permission.

Circulatory Support (INTERMACS) created a classification that is prognostic and clinically useful regarding the need and type of MCS (Table 3). This classification has also been applied to patients transplanted in an emergency situation and showed worse prognosis for patients transplanted in INTERMACS 1-2 compared with INTERMACS 3-4. Therefore hemodynamic stabilization, either with medication or MCS, is strongly recommended prior to HT.

A decision-making algorithm for HT is provided in Figure 1.

Evaluation for Heart Transplant

A holistic approach is imperative for selecting the best recipients for a scarce resource. The expected mortality after HT can be calculated with the IMPACT model. Comorbidities that increase the morbidity and mortality of HT can amount to absolute or relative contraindications and are described in Table 4.

Pulmonary Hypertension

Pulmonary hypertension develops in response to a passive backward transmission of elevated filling pressures in the left ventricle that with time causes vascular remodelling. Right heart catheterization is recommended before HT, as irreversible pulmonary hypertension is associated with right HF and higher mortality.

Reactive pulmonary hypertension is defined as the presence of a transpulmonary gradient > 12 mmHg and a pulmonary vascular resistance > 3 Wood Units. Pulmonary vasoactivity testing with inotropes, vasodilators, and diuretics is necessary to demonstrate reversibility, defined as a reduction of transpulmonary gradient to < 12 mmHg and pulmonary vascular resistance < 3 Wood Units. Unless these values are achieved, HT is contraindicated. In patients listed for HT, the reversibility of pulmonary hypertension should be reassessed at 3- to 6-month intervals.

Bosentan and sildenafil can sometimes reverse pulmonary hypertension in 3 to 4 months and make the patient suitable for HT. However, guidelines do not support their use in patients with left HF. Left ventricular assist devices (LVADs) can also lower pulmonary hypertension and are used as a bridge to candidacy in patients with irreversible pulmonary hypertension refractory to medical treatment.

Current Picture

In the last few years, a trend has been observed toward older recipients with complex clinical profiles, suboptimal donors (54%) and relatively long ischemia times. There is a worrying tendency toward an increase in emergency transplants (the criteria in Spain can be seen in Table 5) representing up to 40% (20% with MCS) of transplants in Spain; in 2013, 50% of candidates were bridged with MCS according to the international registry. These figures reflect an era of donor shortage and an increased acceptance of more complex candidates for HT. Despite this situation, early mortality after HT remains similar to that of previous periods.

Information regarding surgery, organ preservation, and donor selection can be found elsewhere.

Immunosuppression

Long-term outcome depends on the maintenance of the minimum immunosuppression levels necessary to avoid rejection and minimize adverse effects.

Induction Therapy

Induction therapy guarantees rapid and profound immunosuppression immediately after HT. Thymoglobulin has traditionally been used and is highly effective in depleting lymphocytes. The
lack of randomized trials and increased infection rate has currently limited its use to sensitized patients and those at high risk of acute rejection. Dosing thymoglobulin according to lymphocyte count reduces the cumulative dose without compromising efficacy.16

Currently, in patients with a lower rejection risk, induction therapy is performed with interleukin-2 receptor antagonists, such as basiliximab, which has a better safety profile.17 Differences in the use of induction therapy per protocol exist between centers and, while induction therapy consists of basiliximab in 85% of patients in Spain, it is only used in 30% of patients in the international cohort.4,5

Guidelines recommend induction therapy in patients at high risk for acute rejection or renal dysfunction with the aim of delaying the use of calcineurin inhibitors.17–19 Although widely employed, their impact on survival and long-term adverse effects are unknown.

Immunosuppression Maintenance Strategies

Immunosuppression must be higher in the first 3 to 6 months. Over the years, it can be reduced according to individual risk and the results of endomyocardial biopsy. Initially, standard therapy includes the synergistic combination of 3 groups of drugs.17

- Calcineurin inhibitors: cyclosporine or tacrolimus. Both are metabolized by CYP-450 and consequently have many drug interactions. Adverse effects (arterial hypertension, diabetes, dyslipidemia, neurotoxicity) depend on blood levels, and therefore close monitoring is mandatory. Regarding efficacy, tacrolimus may be associated with lower rejection rates without differences in infection or survival.20 Arterial hypertension and dyslipidemia are less frequent with tacrolimus, whereas diabetes risk is increased. Currently, 85% of patients receive tacrolimus during the first year.4,5
- Antiproliferative drugs inhibit de novo synthesis of purines and block cytotoxic and B lymphocyte proliferation. Added to calcineurin inhibitors, they allow a reduction of their levels. Mycophenolate mofetil is chosen in 96% of patients1 since it exhibits better survival, less rejection, and less cardiac allograft vasculopathy than azathioprine.21 The most frequent adverse effects are leukopenia and gastrointestinal intolerance (the delayed-release formulation is better tolerated in this regard).
- Corticosteroids. A wide range of adverse effects (arterial hypertension, diabetes, hyperlipidemia, osteopenia, myopathy, emotional instability, infection) requires their reduction and, if possible, their discontinuation after 6 months. Withdrawal must be closely monitored by biopsy, as a high rate of acute rejection

Figure 1. Decision-making algorithm for patients with advanced HF as defined in Table 1 after appropriate optimization of medical, device, and surgical treatment. BTC, bridge to candidacy; BTD, bridge to decision; BTR, bridge to recovery; BTT, bridge to transplant; DT, destination therapy; INTERMACS, Interagency Registry for Mechanically Assisted Circulatory Support; HF, heart failure; HT, heart transplant; LTVAD, long-term ventricular assist device; MCS, mechanical circulatory support; VO₂, oxygen consumption. In patients in INTERMACS 1 a short-term ventricular assist device should be placed, preferably ventoarterial extracorporeal membrane oxygenation in conditions such as unclear neurological status, unstable hemodynamics and severe coagulopathy. In less catastrophic situations and in INTERMACS 2, a uni- or biventricular short-term ventricular assist device such as the Centrimag can be implanted, as it can provide up to 1 month of support. After resuscitation of the patient, a weaning trial of the device must be performed and, if not possible, assessment for HT is crucial. The next step should be exchange to a LTVAD as BTT or in some cases as DT. In patients in INTERMACS 3, a LTVAD, preferably only supporting the left ventricle, is recommended. After bridge to candidacy, if the contraindication (pulmonary hypertension, time free of cancer or excess weight) is resolved, the patient should be listed.
Table 4
Contraindications for Heart Transplant

<table>
<thead>
<tr>
<th>Absolute</th>
<th>Relative</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systemic disease with life expectancy &lt; 2 years:</td>
<td>Age &gt; 70 years (carefully selected patients may be considered)</td>
</tr>
<tr>
<td>Active neoplasm (if preexisting, evaluation with an oncology specialist is necessary to stratify the risk of recurrence and establish a time to wait after remission)</td>
<td>Diabetes with end-organ damage (except nonproliferative retinopathy) or persistent poor glycemic control (HbA1c &gt; 7.5%) despite treatment</td>
</tr>
<tr>
<td>Systemic disease with multiorgan involvement (systemic lupus erythematosus, amyloidosis, sarcoidosis)</td>
<td>Active infection, except VAD infection. Patients with HIV, hepatitis, Chagas disease and tuberculosis can be considered with strict management</td>
</tr>
<tr>
<td>Severe chronic obstructive pulmonary disease (FEV1 &lt; 1 L)</td>
<td>Severe peripheral arterial or cerebrovascular disease not suitable for treatment</td>
</tr>
<tr>
<td>Renal or hepatic severe dysfunction, if associated renal or liver transplant is not performed</td>
<td>Other serious comorbidity with poor prognosis, such as neuromuscular diseases</td>
</tr>
<tr>
<td>Irreversible pulmonary hypertension</td>
<td>Obesity: BMI &gt; 35 kg/m²</td>
</tr>
<tr>
<td>Pulmonary artery systolic pressure &gt; 50 mmHg</td>
<td>Cachexia: BMI &lt; 18 kg/m²</td>
</tr>
<tr>
<td>Transpulmonary gradient &gt; 12 mmHg</td>
<td>Frailty: when 3 of 5 possible symptoms (including unintentional weight loss of &gt; 5 kg within the past year, muscle loss, fatigue, slow walking speed, and low levels of physical activity) are present</td>
</tr>
<tr>
<td>Pulmonary vascular resistance &gt; 3 Wood units despite treatment</td>
<td>Current alcohol or drug abuse</td>
</tr>
<tr>
<td></td>
<td>Insufficient social support</td>
</tr>
<tr>
<td></td>
<td>Elevated panel-reactive antibody test defined as &gt; 10%</td>
</tr>
</tbody>
</table>

BMI, body mass index; FEV1, forced expiratory volume in 1 second; HbA1c, glycated hemoglobin; HIV, human immunodeficiency virus; VAD, ventricular assist device.

Table 5
Emergent Transplant Criteria in Spain

<table>
<thead>
<tr>
<th>Grade 0 emergency (national priority)</th>
<th>Advanced HF with short-term MCS (including ECMO)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Dysfunctional long-term VAD</td>
</tr>
<tr>
<td></td>
<td>- Mechanical dysfunction</td>
</tr>
<tr>
<td></td>
<td>- Infection</td>
</tr>
<tr>
<td></td>
<td>- Tromboembolism</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Grade 1 emergency (regional priority)</th>
<th>Advanced HF with 1 of the following:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>- Vasoactive drugs and invasive mechanical ventilation</td>
</tr>
<tr>
<td></td>
<td>- Intra-aortic balloon pump</td>
</tr>
<tr>
<td></td>
<td>- Long-term VAD</td>
</tr>
<tr>
<td></td>
<td>Arrhythmogenic storm defined as 3 or more sustained ventricular tachycardia or ventricular fibrillations in 24 h that require intervention to finish them despite maximal antiarrhythmic treatment</td>
</tr>
</tbody>
</table>

ECMO, extracorporeal membrane oxygenation; HF, heart failure; MCS, mechanically circulatory support; VAD, ventricular assist device.

has been observed. Currently, 80% of patients remain on corticosteroids after 1 year and 62% at 7 years.

Alternatives to the first 2 drugs are mammalian target of rapamycin (m-TOR) inhibitors (everolimus, sirolimus), which inhibit the proliferation of lymphocytes and smooth muscle cells. Everolimus combined with calcineurin inhibitors has been shown to reduce rejection and cardiac vasculopathy compared with mycophenolate mofetil, without differences in survival. The SCHEDULE trial demonstrated a reduction in cardiac vasculopathy progression in the group with early everolimus introduction and calcineurin inhibitor withdrawal compared with standard calcineurin inhibitor therapy, but a higher rejection rate was also seen. While a combination of an m-TOR inhibitor and calcineurin inhibitor is recommended in cardiac vasculopathy, a strategy of calcineurin inhibitor withdrawal or minimization is recommended in renal dysfunction and cancer. Even though m-TOR inhibitors have been shown to be useful, a significant decrease in their use has been noticed in the last few years mainly driven by their numerous adverse effects: infections, pneumonitis, proteinuria, effusions, hyperlipidemia, diarrhea, and myelotoxicity.

Single use of tacrolimus in the long-term has also been proposed, without differences in rejection in the first year. However, this is not a common approach and further studies are necessary.

Rejection

Graft rejection is one of the main causes of death in the first few years after HT. Hyperacute rejection due to preformed antibodies against AB0 or human leucocyte antigens is rare at present.

Cellular Acute Rejection

Cellular acute rejection is mediated by T-lymphocytes and is characterized by inflammatory infiltration with myocyte damage. Its severity is classified according to the pathology findings. Only symptomatic patients or asymptomatic patients with severity ≥ 2R (multifocal myocyte damage) are treated with corticosteroids +/- thymoglobulin if there is hemodynamic instability. The incidence of treated rejection in the first year is around 15%.

Surveillance biopsies for detecting rejection are routinely performed in most centers, more frequently in the first 3 months; they are then tapered until 1 year post-transplant and afterwards only if rejection is clinically suspected.

Allomap is a noninvasive gene-expression profiling test for rejection surveillance in HT recipients. This test has a high negative predictive value, identifies patients at low risk for cellular rejection, and can avoid routine biopsies a few months after HT.
Antibody-mediated Rejection

This is a recently described entity, present in 10% to 20% of HT. B-lymphocytes produce antibodies against human leucocyte antigens activating an inflammatory response that causes endothelial dysfunction. Prevention by minimizing exposure to alloantigens (avoiding nonessential blood transfusions) and maintaining appropriate immunosuppression is paramount. Diagnosis is challenging and suspicion is key. The usual clinical picture is a patient with HF, left ventricular dysfunction without cellular infiltration in the biopsy, female sex, allo sensitized, previous transfusion, retransplanted, and prior LVAD or history of Cytomegalovirus infection. Confirmation is made by pathology findings (vasculitis, edema), C4d or C3d deposition, and determination of antihuman leucocyte antibodies in serum. Therapy and its duration are not well established but there is a consensus in treating patients with biopsy findings and dysfunction or antibodies. A more detailed review of diagnosis and treatment can be found elsewhere.29

Cardiac Allograft Vasculopathy

This entity is characterized by diffuse and concentric thickening of the intima of the epicardial and intramural coronary arteries. Its etiology is not clear but it is considered to be the manifestation of chronic rejection influenced by nonimmunological factors, such as diabetes mellitus, hypertension, smoking, and Cytomegalovirus infection. Its clinical expression includes angina, myocardial infarction, or sudden death but it usually manifests as HF with or without ventricular dysfunction. Statin therapy has been shown to reduce graft vasculopathy and mortality and therefore it is indicated in all HT patients irrespective of lipid levels.17 A baseline coronary angiography 1 month after HT and another at 1 year are recommended. After that, the need for repeat coronary angiographies or noninvasive tests is variable.17,19

Infectious Complications: Prophylaxis

Diagnosis of infection can be challenging in HT recipients and treatment must be aggressive. Prevalence is higher in the initial 6 months. In the very early period, previous recipient infections can be exacerbated and donor-transmitted or surgery-related infections may appear. Between 0.5 to 6 months, opportunistic infections emerge: viral (Cytomegalovirus), fungal (Aspergillus and P. jiroveci) and bacterial (Nocardia and Lysteria). After 6 months, the risk diminishes and community-acquired infections are the most common.

Prophylaxis should be started 10 to 15 days post-HT: ganciclovir or valganciclovir for Cytomegalovirus for 3 months or preemptive therapy guided by polymerase chain reaction determinations. In the case of Cytomegalovirus donor positive/recipient negative recipients, prophylaxis for 6–12 months is recommended. Trimethoprim/sulfamethoxazole is recommended for P. jiroveci and Toxoplasma gondii for 6 months in all patients, or for up to 1 year if treated with everolimus; antifungal prophylaxis with nystatin to prevent candidiasis while the patient is on high dose corticosteroids and inhaled amphotericin B for Aspergillus during the initial hospitalization. Further information about dosage and regimens can be found in various guidelines.17–19

Causes of Death After Heart Transplant

Overall, survival at 1, 5, and 10 years is 81%, 68%, and 51%, respectively. In the Spanish cohort, median survival is 10.9 years. Survival is worse in older recipients, with older donors, emergency transplant, and recipients supported by venoarterial extracorporeal membrane oxygenation (VA-ECMO).4,5

Primary graft failure is the leading cause of death in the first month, while infection is the most common cause during the first year. Primary graft failure is defined as ventricular dysfunction without a clear etiology. The RADIAL score can help stratify risk.30 After 1 year, the main causes of death are cardiac vasculopathy and malignancy.4 Globally, the main cause of death in Spain is cardiac vasculopathy (20%), followed by infection (16%), primary graft failure (14%) and tumors (13%).4

MECHANICAL CIRCULATORY SUPPORT

The use of MCS has grown exponentially over the past 15 years, mostly as BTT. However, other strategies after implantation of MCS exist, as noted previously.

Type of Mechanical Circulatory Support

The type of MCS will depend on the clinical situation, defined by the INTERMACS classification. Two groups are distinguished:

- Short-term ventricular assist devices aim to support hemodynamically unstable HF patients for days or weeks. Initially used for postcardiotomy shock, their use was expanded to stabilize a shocked patient and gain time either for recovery, decision, or BTT.
- Long-term ventricular assist devices are designed to assist patients with advanced HF during a period of months to years. While awaiting candidacy, transplant and, in very few cases, recovery. However, with the appearance of continuous-flow LVADs and their increased durability, DT has become an option. Currently, many patients who receive a LVAD are DT equivalent, because only 30% of patients with MCS implanted as BTT will receive an organ within the first year of listing.31

Additionally, VADs can be classified according to various criteria: a) ventricle supported: left, right or both (biventricular VAD or total artificial heart); b) location: extracorporeal or intracorporeal; c) flow provided: pulsatile-flow or continuous-flow, and d) pump: pneumatic, axial, or centrifugal.

Short-term Ventricular Assist Devices

Percutaneous STVAD will not be discussed, as these devices are reviewed in another article published in Revista Española de Cardiología.

In our environment, VA-ECMO and Centrimag (St. Jude Medical, Pleasanton, California, USA) are the preferred options in INTERMACS 1–2. Venoarterial ECMO is a modified cardiopulmonary bypass that supports both ventricles (3.5–4.5 L/min), with the possibility of peripheral cannulation, even outside of the operating room. The Centrimag is a central continuous-flow pump (4–7 L/min) that can be used as left, right, or biventricular support via stenotomy. When to use one or the other is discussed in Figure 1.

Despite the use of MCS in cardiogenic shock, mortality is still around 50%, mainly due to shock prior to implant and less frequently secondary to complications of the STVAD.32 What is important is to design a strategy after implanting a STVAD, aiming for recovery when possible, but it is also important to have the possibility of bridging from VA-ECMO to Centrimag, from STVAD to HT if waiting times are acceptable, or from STVAD to LT VAD.
Implantation of a STVAD in a community hospital and then bridging to a LTVAD in a tertiary hospital can be performed with similar results to those achieved when the whole process is performed in a tertiary hospital, emphasizing the importance of early stabilization with STVADs in cardiogenic shock.

In Spain, 47% of patients undergo HT in an emergency situation: > 50% of them with an intra-aortic balloon pump, 23% with ECMO, 16% with continuous-flow VADs, and 6.5% with pulsatile-flow VADs. Venoarterial ECMO is clearly associated with worse survival after HT compared with the remaining options. However, the registry does not distinguish between STVADs and LTVADs and therefore outcomes after HT in patients with a STVAD, such as the Centrimag, are unknown. This issue will be clarified by an analysis of STVAD as BTT that is currently underway (ASIST-TC study). The most common STVADs are shown in Figure 2.

**Figure 2.** Short-term ventricular assist devices. A: CARDIOHELP (Maquet, Bridgewater, New Jersey, United States) extracorporeal membrane oxygenation system. B: Centrimag (St. Jude Medical, Pleasanton, California, United States) is a continuous-flow centrifugal pump that can support one or both ventricles. VAD, ventricular assist device. Reproduced with the permission of Maquet and St. Jude Medical.

**Long-term Ventricular Assist Devices**

In the seventh annual report of the INTERMACS database, more than 15,000 LTVADs have been reported with a mean rate of 2500 patients per year in the last 2 years. Although initially conceived for as BTT, their usefulness as DT was first evaluated in the REMATCH trial. This trial used pulsatile-flow LTVADs and, although they provided adequate support, their use was limited by their short durability, cost, and large size. The introduction of continuous-flow LTVADs led to better survival free from stroke and device failure. Currently > 90% of LTVADs are continuous-flow. This improvement in technology has also led to an increase in DT to 46% of all implants in the United States.

In Spain, the most commonly employed LTVAD until recent years was the Excor (Berlin Heart, Berlin, Germany). However, because of the need for high levels of anticoagulation and antiplatelet therapy to avoid thromboembolic complications, most centers currently use it only for mid-term support or as biventricular support. Currently, continuous-flow LTVAD are the preferred option for long-term support.

The VADs currently being used are shown in Figure 3. The INCOR was the first LTVAD used in Spain. The HeartMate II is the most frequently used continuous-flow LTVAD, and the only one approved for DT. The others are smaller centrifugal pumps, avoiding the need for an abdominal pocket. In the ADVANCE and ENDURANCE trials, the HVAD was shown to be noninferior to the HeartMate II. Initial experience with HeartMate III has shown better outcomes at 6 months compared with HeartMate II, primarily because of a lower rate of pump thrombosis.

**Indications, Risk and Outcomes**

Heart failure patients eligible for LVAD implantation must fulfill the criteria described in Table 6. A thorough evaluation of risk is mandatory and overall, outcomes of patients with continuous-flow LVAD are satisfactory. The current survival rates are approximately 80% and 70% at 1 and 2 years as BTT and 75% and 65% as DT, but are worse with biventricular support (50% at 1 year). Transplant outcomes in patients bridged with continuous-flow LTVAD are similar to those without bridging. For patients in INTERMACS 1, LTVAD implantation should be avoided since their survival is lower. Short-term ventricular assist device should be considered in these patients instead.

Regarding ambulatory HF, the ROADMAP trial studied patients in INTERMACS > 4. The 1-year freedom from death, urgent HT, or delayed LVAD was better for the LTVAD group compared with medical treatment (80% vs 63%, P = .024). However, adverse events were twice as common with LTVADs, indicating the need for caution regarding too early LTVAD placement.
Patients Potentially Eligible for Implantation of a Left Ventricular Assist Device

<table>
<thead>
<tr>
<th>Patients with &gt; 2 mo of severe symptoms despite optimal medical and device therapy and more than 1 of the following:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left ventricular ejection fraction &lt; 25%, and if measured, peak VO₂ &lt; 12 mL/kg/min</td>
</tr>
<tr>
<td>≥ 3 hospitalizations for HF in the previous 12 mo without an obvious precipitating cause</td>
</tr>
<tr>
<td>Dependence on intravenous inotropic therapy</td>
</tr>
<tr>
<td>Progressive end-organ dysfunction (worsening renal and/or liver function) due to reduced perfusion and not to inadequate ventricular filling pressure (PCWP &gt; 20 mmHg and SBP &lt; 80-90 mmHg or CI ≤ 2 L/min/m²)</td>
</tr>
<tr>
<td>Absence of severe right ventricular dysfunction together with severe tricuspid regurgitation</td>
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CI, cardiac index; HF, heart failure; PCWP, pulmonary capillary wedge pressure; SBP, systolic blood pressure; VO₂, oxygen consumption.

Adapted from Ponikowski et al., with permission.
Complications and Management of Left Ventricular Assist Device Patients

Initially, the standard treatment to prevent thromboembolic complications is aspirin 81 to 325 mg once daily to achieve an arachidonic acid inhibition > 70%, and vitamin K antagonists to achieve an international normalized ratio of 2 to 3. Clinical practice has established a maximum goal of an international normalized ratio of 2.5 in the absence of other thromboembolic risk factors, as bleeding occurs more frequently than thrombosis, although antithrombotic therapy must be tailored to the specific device and patient. Bridging with heparin in the initial postoperative period is generally recommended.

One of the most feared complications is stroke, which can occur in 7% to 15% of patients with an LVAD. Predictors of ischemic stroke with the HVAD are aspirin ≤ 81 mg and atrial fibrillation, whereas predictors of hemorrhagic stroke are a mean arterial pressure > 90 mmHg, aspirin ≤ 81 mg, and an international normalized ratio > 3. To decrease the incidence of hemorrhagic strokes, it is crucial to strictly control blood pressure so that the mean arterial pressure is < 90 mmHg.50

Bleeding events are favored by shear stress on blood components and reduced pulse pressure in continuous-flow technology. The current incidence (7.79 events/100 patient-months) is lower than in previous periods.51 The most prevalent is gastrointestinal bleeding, which is a leading cause of readmission but does not affect survival. Described predisposing factors are age, lower albumin levels, and lower body mass index.53 Pathophysiology is explained by acquired von Willebrand factor deficiency, impaired platelet aggregation, and gastrointestinal angiodysplasias due to reduced pulse pressure akin to Heyde syndrome.52,53

The TRACE US study54 analyzed 100 patients with reduced antithrombotic therapy after a bleeding episode. Despite this, subsequent bleeding occurred in 52%, although rates of ischemic stroke were similar.

The incidence of pump thrombosis rose from 2.2% at 3 months after implantation in 2011 to 8.4% in 2013,55 probably due to changes in clinical management with lower goals of anticoagulation and antiaggregation and lower LVAD flows in order to achieve opening of the aortic valve. Pump thrombosis is a daunting complication that depends on device characteristics, operative technique, antithrombotic management, and patient factors. This complication must be suspected in the presence of hemolysis (lactate dehydrogenase elevation and/or high plasma free hemoglobin), a transient pump power increase, or left HF. The best diagnostic tool for Heartmate II is the Columbia ramp study,56 in which blunted reductions in left ventricular end diastolic diameter in response to increases in pump speed indicate an obstruction to flow through the device. These slope parameters cannot be directly applied to HVAD patients, as the left ventricle end diastolic diameter is drastically smaller.57 Treatment is based on increasing antithrombotic medication and if there is hemodynamic instability, fibrinolysis or pump exchange may be performed.

Another frequent complication is aortic insufficiency, which is noted in 25% to 52% of patients after 1 year of continuous-flow LVAD support and is cumulative over time. The reasons for the development of aortic insufficiency are thought to be: a) lack of opening of the aortic valve, which may lead to leaflet fusion, and b) altered flow dynamics in the ascending aorta, which may contribute to aortic sinus dilatation. Therefore, when more than mild aortic insufficiency is detected prior to LVAD implantation, it is recommended to repair or replace the aortic valve. To prevent aortic insufficiency after LVAD implantation, it is recommended to optimize speed to eliminate more than mild MR and position the septum at the midline. If both are achieved, speed may be reduced to allow intermittent aortic valve opening. If aortic insufficiency secondary to LVAD is asymptomatic, speed reduction to maximize aortic valve opening is recommended. If the patient is symptomatic, an increase in speed is recommended. If symptoms persist after hemodynamic assessment, repair, replacement or closure of the aortic valve with a patch or by sewing the leaflets may be considered.58

Driveline infection is a feared complication present in up to 40% of patients over time. The incidence of infection can be decreased by patient education and the use of a standardized kit with a silver dressing and an anchoring device.59

Finally, the development of human- leukocyte antigen antibodies during MCS has been described. Younger age, pre-VAD panel-reactive antibodies, and female sex were independent predictors of elevated antibodies post-VAD. Although the development of these antibodies is associated with a longer waiting time for HT due to the need for virtual cross-match, no association with increased rejection, graft failure or death after HT was found.60

FUTURE PROSPECTS

The incipient use of donor hearts after circulatory death may increase the donor pool.61 However, this increase might not be sufficient to meet the needs of all the patients with advanced HF. To increase the use of LTVADs in the future, we would need to reduce their cost, diminish thrombotic and bleeding complications and avoid the driveline to minimize infections.

In Spain, the use of LTVADs as BTT or bridge to candidacy is slowly increasing, but their use as DT is still anecdotal and is restricted to young patients with an absolute contraindication for HT and those aged 65 to 75 years with comorbidities that may limit graft survival. The main reason for the low rates of LVAD use is the high cost of the device, which is approximately 94 600 €. With a current threshold of 30 000 €/quality-adjusted life years 62 for end-of-life care interventions, by reducing the cost by 15%, the incremental cost-effectiveness ratio may be acceptable.

CONCLUSIONS

HT remains the best available therapy for patients with advanced HF but, given the shortage of donors and long waiting lists, LVADs are increasingly being used to save lives and enhance quality of life. In Spain, a shift from STVADs to LTVADs as BTT or bridge to candidacy is occurring but the pace is slow. Implantation of LTVADs as DT can be an alternative for highly selected patients, but access to these devices is limited by their cost. Widespread use of LVADs will only be viable if their complications are reduced and become cost-effective.

CONFLICTS OF INTEREST

U.P. Jorde is a consultant to St. Jude Medical (no honoraria).

REFERENCES


