

Intermacs

Screening Log

Implant Date

MM/DD/YYYY

Inclusion: Patient must meet all inclusion criteria:

If patient meets all inclusion criteria then check **ALL** inclusion reasons below.

- Patient receives a durable mechanical circulatory support device (MCSD) which is FDA approved
- Implanted on or after March 1, 2006 (The device does not need to be the first implant for the patient)
- Patient signed informed consent for the registry

Exclusion: Any exclusion will disqualify the patient for entry into INTERMACS®

If patient meets **ANY** exclusion criteria then check any of the appropriate exclusion reasons below (check all that apply).

- Patient receives a durable mechanical circulatory support device (MCSD) which is not FDA approved
- Patient is incarcerated (prisoner)
- Patient did not sign the informed consent

Device type

- LVAD
- RVAD
- Both (LVAD + RVAD in the same OR visit)
- Total Artificial Heart

Device brand

- Berlin Heart EXCOR (paracorporeal)
- HeartWare HVAD
- HeartMate II LVAS
- HeartMate III
- HeartMate IP
- HeartMate VE
- HeartMate XVE
- Micromed DeBakey VAD - Child
- Novacor PC
- Novacor PCq
- Thoratec IVAD
- Thoratec PVAD
- Other, Specify

Specify brand

Device brand (RVAD)
Specify brand (RVAD)

Age Range

- 19 to 39
- 40 to 59
- 60 to 79
- 80+

Race

- American Indian or Alaska Native

- Asian
- African-American or Black
- Hawaiian or other Pacific Islander
- White
- Unknown / Undisclosed
- Other / none of the above

Ethnicity: Hispanic or Latino

- Yes
- No
- Unknown

Gender

- Male
- Female
- Unspecified

Did death occur within 2 days post implant?

- Yes
- No

Is this VAD an investigational device?

- Yes
- No

Is patient involved in a VAD related study?

- Yes
- No
- Unknown

What is the name of the study?

Is this an industry sponsored post approval study?

- Yes
- No
- Unknown