



Randomized Clinical Evaluation of the AccuCinch Ventricular Restoration System in Patients who Present with Symptomatic Heart Failure with Reduced Ejection Fraction (HFrEF)

Status: Recruiting

Eligibility Criteria

Sex: Male or Female

Age Group: 18 years and over This study is NOT accepting healthy

volunteers

Inclusion Criteria:

- at least 18 years old - Ejection Fraction: between 20% and 40% measured by transthoracic echocardiography (TTE) - diagnosis and treatment for heart failure should be established at least 90 days before entering the study & should be on stable, optimal medical therapy for at least 30 days

Exclusion Criteria:

- myocardial infarction or any percutaneous cardiovascular intervention, cardiovascular surgery, or carotid surgery within 90 days prior to consent - any planned cardiac surgery or interventions within the next 180 days - women who are pregnant, planning to become pregnant, or are breast feeding - additional cardiac and medical diagnosis will exclude participation (study staff will review)

Conditions & Interventions

Conditions:

Heart & Vascular

Keywords:

Clinics and Surgery Center (CSC), Dilated Cardiomyopathy, Heart Failure, Heart Failure With Reduced Ejection Fraction (HFrEF)

More Information

Description: The objective of this study is to evaluate the safety and efficacy of the AccuCinch Ventricular Restoration System in patients with symptomatic heart failure with reduced ejection fraction (HFrEF).

Study Contact: Melanie Farinella - crawl027@umn.edu

Principal Investigator: Greg Helmer

Phase: NA

IRB Number: STUDY00013236

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