

An open, parallel-group, randomized, early feasibility trial to evaluate the initial safety and performance of the INGA catheter in labor induction

Status: Recruiting

Eligibility Criteria

Sex: Female

Age Group: 18 years and over

This study is NOT accepting healthy volunteers

Inclusion Criteria:

- pregnant adults ages 18–56 - at least 37 weeks pregnant - pregnant with one baby - planning to have labor induced using a balloon catheter

Exclusion Criteria:

- pregnant with more than one baby (twins, triplets, etc.) - baby is not in a head-down position - previous cesarean section or other surgery that left a scar on the uterus - you or your baby need immediate delivery for medical reasons

Conditions & Interventions

Conditions:

Women's Health

Keywords:

catheter, Labor, pregnant

More Information

Description: The purpose of this study is to evaluate the safety and effectiveness of a new device used during labor induction. We will enroll pregnant adults who are at least 37 weeks pregnant and are scheduled to have labor induced using a balloon catheter. The study will evaluate whether the device can improve monitoring of the pregnant person and baby during labor.

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