

## Intraperitoneal FT536 in Recurrent Ovarian, Fallopian Tube, and Primary Peritoneal Cancer

**Status:** Recruiting

### Eligibility Criteria

**Sex:** Female

**Age Group:** 18 years and over

This study is NOT accepting healthy volunteers

#### Inclusion Criteria:

- epithelial ovarian cancer, fallopian tube, or primary peritoneal cancer that has recurred after treatment (no limit to the maximum number of prior treatments) - must have received prior bevacizumab - if there is a BRCA mutation, must have received a prior PARP inhibitor - agree to have an intraperitoneal catheter placed before the 1st dose of study drug - see link to [clinicaltrials.gov](http://clinicaltrials.gov) for complete inclusion & exclusion criteria

#### Exclusion Criteria:

- women who are pregnant, breastfeeding or planning on becoming pregnant in the next 6 months - active autoimmune disease requiring systemic immunosuppressive therapy - history of severe asthma and currently on chronic medications (more than inhalers) - received enoblituzumab - CNS disease such as stroke, epilepsy, CNS vasculitis, or neurodegenerative disease or needing medications for these conditions in the past 2 years

### Conditions & Interventions

#### Conditions:

Cancer, Women's Health

#### Keywords:

Clinics and Surgery Center (CSC), Fallopian Tube Cancer, Gynecologic Cancer, Ovarian Cancer, Primary Peritoneal Cavity Cancer

### More Information

**Description:** FT536 is a type of cell product made up of "natural killer" or NK cells. NK cells are a type of immune blood cell that are known to attack cancer cells. FT536 is produced by growing cells that come from a healthy human donor. The primary purpose of this study is to identify a safe dose of FT536 cells when given alone (monotherapy).

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#### IRB

**Number:** STUDY00021809

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