# Non-steroidal anti-inflammatory drug (NSAID) use in breast surgery: prospective randomised trial

Submission date

15/05/2005

Recruitment status

No longer recruiting

Registration date

12/09/2005

Overall study status

Completed

**Last Edited** 

Condition category

30/07/2008 Cancer Retrospectively registered

Protocol not yet added

SAP not yet added

Results added

Raw data not yet added

Study completed

### **Plain English Summary**

Not provided at time of registration

## Contact information

### Type(s)

Scientific

### Contact name

Dr Samir Hidar

#### Contact details

71, rue Ch Kallala Sousse **Tunisia** 4011

## Additional identifiers

**EudraCT/CTIS** number

IRAS number

ClinicalTrials.gov number

Protocol/serial number

N/A

## Study information

### Scientific Title

### **Study hypothesis**

Modification of drainage volume after NSAID administration

### Ethics approval required

Old ethics approval format

### Ethics approval(s)

Not provided at time of registration

### Study design

Randomised controlled trial

### Primary study design

Interventional

### Secondary study design

Randomised controlled trial

### Study setting(s)

Not specified

### Study type(s)

Treatment

### Participant information sheet

#### Condition

Breast cancer

#### **Interventions**

Before entering operating room for breast cancer patients are randomised into two groups:

- 1. NSAID
- 2. Placebo

### Intervention Type

Drug

#### Phase

**Not Specified** 

### Drug/device/biological/vaccine name(s)

**NSAID** 

### Primary outcome measure

Total volume drained after modified radical mastectomy

### Secondary outcome measures

### Breast surgery complications

## Overall study start date

01/04/2005

## Overall study end date

30/11/2005

## **Eligibility**

### Participant inclusion criteria

Eligible patienst are women with a breast cancer undergoing modified radical mastectomy

### Participant type(s)

**Patient** 

### Age group

Adult

#### Sex

Female

### Target number of participants

100

### Participant exclusion criteria

- 1. Known allergy or contraindication to NSAID
- 2. Planned immediate or late breast reconstruction

### Recruitment start date

01/04/2005

### Recruitment end date

30/11/2005

## Locations

### Countries of recruitment

Tunisia

## Study participating centre

71, rue Ch Kallala

Sousse

Tunisia 4011

## Sponsor information

### Organisation

Farhat Hached University Teaching Hospital (Tunisia)

### Sponsor details

Boulevard M. Karoui Sousse Tunisia 4001

### Sponsor type

Hospital/treatment centre

### **ROR**

https://ror.org/0059hys23

## Funder(s)

### Funder type

Hospital/treatment centre

### **Funder Name**

Farhat Hached University Teaching Hospital (Tunisia)

## **Results and Publications**

## Publication and dissemination plan

Not provided at time of registration

Intention to publish date

Individual participant data (IPD) sharing plan

### IPD sharing plan summary

Not provided at time of registration

### Study outputs

Output type Details Date created Date added Peer reviewed? Patient-facing?

Abstract results Abstract: 17/11/2006 No No