

# De verbreding van de bovenkaak met schedelbot of bekkenkambot: Een vergelijking.

No registrations found.

|                              |                            |
|------------------------------|----------------------------|
| <b>Ethical review</b>        | Not applicable             |
| <b>Status</b>                | Suspended                  |
| <b>Health condition type</b> | -                          |
| <b>Study type</b>            | Observational non invasive |

## Summary

### ID

NL-OMON26526

### Source

NTR

### Health condition

retentieverlies bovenprothese  
atrofie van de bovenkaak

problems with retention of denture  
atrophic maxilla

## Sponsors and support

**Primary sponsor:** UMCG

**Source(s) of monetary or material Support:** UMCG

## Intervention

## Outcome measures

### Primary outcome

Pain levels.

## Secondary outcome

1. Donor site morbidity;
2. Intra oral morbidity;
3. Bone growth (histology).

## Study description

### Background summary

N/A

### Study objective

Augmentation of the maxilla with calvarial bone has a lower complication rate than iliac crest bone.

### Study design

First postoperative days, 2 weeks, 6 weeks, 12 weeks, 16 weeks, half year, one year, two years.

### Intervention

Two standard treatments to augment the maxilla are compared.

## Contacts

### Public

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## Eligibility criteria

### Inclusion criteria

1. Edentulous patients;
2. Cannot wear an upper denture satisfactorily;
3. Have an atrophied maxilla with insufficient bone volume to support dental implants;
4. Written informed consent.

### Exclusion criteria

1. Unable to be operated on due to severe health problems;
2. The use of intravenous bisphosphonates.

## Study design

### Design

|                     |                            |
|---------------------|----------------------------|
| Study type:         | Observational non invasive |
| Intervention model: | Parallel                   |
| Allocation:         | Non controlled trial       |
| Masking:            | Open (masking not used)    |
| Control:            | N/A , unknown              |

### Recruitment

NL

|                           |             |
|---------------------------|-------------|
| Recruitment status:       | Suspended   |
| Start date (anticipated): | 01-07-2013  |
| Enrollment:               | 40          |
| Type:                     | Anticipated |

## Ethics review

|                   |                |
|-------------------|----------------|
| Not applicable    |                |
| Application type: | Not applicable |

## Study registrations

### Followed up by the following (possibly more current) registration

No registrations found.

### Other (possibly less up-to-date) registrations in this register

No registrations found.

### In other registers

| Register | ID                                  |
|----------|-------------------------------------|
| NTR-new  | NL3795                              |
| NTR-old  | NTR3968                             |
| Other    | :                                   |
| ISRCTN   | ISRCTN wordt niet meer aangevraagd. |

## Study results

### Summary results

N/A