

Non surgical peri-implantitis treatment

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Ethische beoordeling	Positief advies
Status	Werving gestopt
Type aandoening	-
Onderzoekstype	Interventie onderzoek

Samenvatting

ID

NL-OMON23755

Bron

NTR

Verkorte titel

NSPT

Aandoening

Peri-implantitis

Ondersteuning

Primaire sponsor: University Medical Center Groningen

Overige ondersteuning: Graduate School of Medical Sciences, Kolff institute

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

- Mean peri-implant bleeding score (%)
- Mean peri-implant and full-mouth periodontal suppuration on probing score (%);
- Mean peri-implant and full-mouth periodontal probing pocket depth;
- Mean peri-implant and full-mouth periodontal plaque score (%);

Toelichting onderzoek

Achtergrond van het onderzoek

Peri-implantitis is an infectious condition of the tissues around osseointegrated implants resulting in loss of supporting bone and clinical signs of inflammation (bleeding and/or suppuration on probing). Various non-surgical and surgical treatment modalities have been described in the literature including mechanical debridement and/or pharmaceutical therapy (chlorhexidine, local or systemic antibiotics), aimed at removing bacteria and decontamination of the implant surface (Heitz-Mayfield 2014). Despite these various treatment strategies, the most effective treatment option for treating peri-implantitis lesions in a non-surgical way remains unclear. Therefore the search for a potentially beneficial treatment modality is still imperative. One such potentially beneficial treatment might be the use of air polishing. Modern air polishing devices and their specific powders for subgingival application are becoming increasingly significant in the context of maintenance therapy. It has been shown that supportive therapy consisting of debridement and decontamination of implants and suprastructures with air polishing leads to better clinical results than conventional mechanical supportive therapy (Muthukuru et al. 2012). For non-surgical treatment of peri-implantitis, air polishing has only scarcely been investigated (Sahm et al. 2011 (mild to moderate peri-implantitis), Renvert et al. 2010 (severe peri-implantitis)). Sample sizes were small and study designs varied among the studies.

If peri-implantitis is left untreated it may ultimately lead to implant loss. Moreover, it is thought that peri-implantitis, like periodontitis, may exert systemic effects. The inflammatory burden, consisting of bacteria and inflammatory mediators entering the systemic circulation, is thought to be related to the amount of inflamed peri-implant tissue. The greater the amount of inflamed peri-implant tissue, the greater the amount (and the chance) of bacteria and inflammatory mediators entering the systemic circulation may be. On the basis of these considerations the aim of the present study is to investigate the clinical, microbiological and radiographical effectiveness of decontamination of the implant surface during non-surgical treatment of peri-implantitis using air polishing. In addition, immunological samples will be analyzed in order to evaluate the effect of therapy on inflammatory parameters.

Doel van het onderzoek

The primary objective of this randomized controlled trial is to compare the clinical effect of decontamination of the implant surface during the non-surgical treatment of peri-implantitis using air polishing or ultrasonic treatment. Secondary objectives are to assess the microbiological and radiographical effects of these treatment options of peri-implantitis and to evaluate the influence of peri-implantitis and its treatment on inflammatory parameters.

Onderzoeksopzet

Inclusion: 2016-2019
End of followup: end 2020
Statistical analysis: beginning 2021
Writing manuscript: 2021
Submission 2021

Onderzoeksproduct en/of interventie

Airpolishing (Perioflow® EMS), Ultrasonic device (Piezon® EMS)

Contactpersonen

Publiek

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Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

The patient is ≥ 18 years of age;
The patient has at least one endosseous implant in the oral cavity with clinical and radiographical signs of peri-implantitis. Peri-implantitis is defined as progressive loss of marginal bone ≥ 2 mm, as compared to the baseline radiograph (after placing the definitive restoration) in combination with bleeding and/or suppuration on probing;
The implants have been in function for at least two years;
The patient is capable of understanding and giving informed consent.

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

- Medical and general contraindications for the procedures;
- A history of local radiotherapy to the head and neck region;
- Pregnancy and lactation;
- Uncontrolled diabetes mellitus (HbA1c < 7% or < 53 mmol/mol);
- Use of antibiotics during the last 3 months;
- Known allergy to chlorhexidine;
- Long-term use of anti-inflammatory drugs;
- Incapability of performing basal oral hygiene measures as a result of physical or mental disorders;
- Implants with bone loss exceeding 2/3 of the length of the implant or implants with bone loss beyond the transverse openings in hollow implants;
- Implant mobility;
- Implants at which no position can be identified where proper probing measurements can be performed;
- Previous surgical treatment of the peri-implantitis lesions;
- Previous non-surgical treatment of the peri-implantitis lesions during the last 3 months (scaling or curettage)
- Chronic bronchitis and asthma
- Presents of natural dentition (only for patients in heamatological study)

Onderzoeksopzet

Opzet

Type:	Interventie onderzoek
Onderzoeksmodel:	Parallel
Toewijzing:	Gerandomiseerd
Blindering:	Enkelblind
Controle:	Geneesmiddel

Deelname

Nederland	
Status:	Werving gestopt
(Verwachte) startdatum:	27-10-2016
Aantal proefpersonen:	80
Type:	Werkelijke startdatum

Voornemen beschikbaar stellen Individuele Patiënten Data (IPD)

Wordt de data na het onderzoek gedeeld: Nog niet bepaald

Ethische beoordeling

Positief advies

Datum: 29-01-2020

Soort: Eerste indiening

Registraties

Opgevolgd door onderstaande (mogelijk meer actuele) registratie

Geen registraties gevonden.

Andere (mogelijk minder actuele) registraties in dit register

Geen registraties gevonden.

In overige registers

Register	ID
NTR-new	NL8339
Ander register	METC UMCG : METc 2016.356

Resultaten