The impact of cantilever direction on the clinical outcome of implant-supported fixed dental prostheses

By

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ABSTRACT

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It has been shown that dental implants tend to lose bone over time, which ultimately results in soft tissue loss. Recent studies, however, suggest that the design of implant-supported prostheses may contribute to peri-implant tissue stability.

Aim: To evaluate retrospectively radiographic bone loss around implants supporting cantilevered pontics with either mesial or distal direction, and to identify the technical complications that may occur with mesial/distal cantilever fixed dental prosthesis.

Material and Methods: Records of 14 partially dentate patients, aged between 45-83 years (mean age 69.4), who were treated from March 2003 to March 2015, with mesial/distal cantilever implant-supported fixed dental prostheses were reviewed. For each implant, the radiographs from the time of implant loading were compared to radiographs from the last follow-up visit.
There were evaluated regarding:

1- The distance from widest diameter of the abutment to the crest of the peri-implant bone.
2- The radiographic changes of marginal hard tissue height from the time of implant loading compared to the time of the last follow-up appointment.

Technical complications were noted as (screw-loosening, prosthesis de-cementation and prosthesis loosening).

**Results:** A total of 28 cantilever implant-supported fixed dental prostheses supporting 32 cantilever units were evaluated. Of these 10 (35.7%) had mesial cantilevers, while 18 (64.7%) had a distal cantilever. There was no significant difference in the distribution of the cases between males and females (7 males and 7 females). The non-smokers were 43%. And the non-recorded were 29%. All technical complications associated with implants adjacent to distal cantilever pontics (prosthesis loosening P= 0.114, prosthesis de-cementation P= 0.114 and implant abutment screw-loosening: P= 0.37). Furthermore, all technical complications occurred with cantilever arm length < 10 mm. Mesial and distal bone loss on implants adjacent to the cantilevered units was not state different if cantilever direction was mesial or distal (mesial cantilever P= 0.533, distal cantilever: P= 0.82)
**Conclusion:** Within the limitation of this study, marginal bone loss does not seem to be influenced by the presence of mesial or distal cantilever extensions. Minor technical complications were found with a distal cantilever (prosthesis loosening, prosthesis de-cementation and implant abutment screw-loosening).
DEDICATION

I would like to dedicate this thesis to my mum, my husband and to my brothers and sisters. Their boundless love and guidance have encouraged me to accomplish all my professional and personal goals.
DECLARATION

I declare that all the content of the thesis is my own work. There is no conflict of interest with any other entity or organization.

Name: Hayat Taresh Alaleeli

Signature:
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To my mum, brothers and sisters, thank you for your support and taking care of my baby.
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1. Introduction:

Today, partially edentulous individuals represent the main and steadily increasing group of patients being considered for the rehabilitation with oral implants\(^1\). Most of these patients are middle-aged, that is, between 40 and 50 years when provided with oral implants\(^2\). Thus, with increasing life expectancy, it is assumable that the patients will be in need of their implant-supported reconstructions to function for decades\(^3\). When advising the patient on different treatment options, survival rates and the incidence of biological and technical events should thus base on mean observation periods of at least 5 years to supply the patient with reliable information\(^4\). Furthermore, the treatment decision should ideally be based on systematic reviews of the available evidence\(^5\). It may be stated that implant-supported fixed dental prostheses (FDPs) are a safe and predictable treatment method with high survival rates. However, biological and technical complications are frequent. This, in turn, means that substantial amount of chair time has to be accepted by the patient and the dental service following the incorporation of implant-supported FDPs\(^5\).
To minimize the time needed to address complications, dental professionals should make great efforts in choosing reliable components and materials for implant-supported FDPs and the patients should be part of a well-structured maintenance program after treatment with implant-supported FDPs. Biological complications for implant-supported restorations should be defined by (1) the threshold level of pocket probing depth (PPD), (2) the presence/absence of bleeding on probing (BOP)/suppuration assessed at any examination interval, and (3) marginal bone loss over time in relation to the bone level assessed at the time of prosthesis installation. Technical complications should be divided into (1) major, such as implant fracture, abutment fracture, and loss of supra-structures, (2) medium, such as abutment fracture, veneer or framework fractures, and phonetic complications, and (3) minor, such as abutment and screw loosening, loss of retention, de-bonding, loss of screw hole sealing, veneer chipping (to be polished), and occlusal adjustments. Esthetic impairments should be reported.\textsuperscript{6} \textit{Biologic complications} refer to disturbances in implant function that affect the supporting peri-implant tissues. These include early and late implant failures and adverse reactions in the peri-implant hard and soft tissues. \textit{Technical complications} are collective term for mechanical damages of the implant, implant components, and suprastructures.\textsuperscript{7}

Several reports have been published, showing favorable long-term outcomes with different implant systems.\textsuperscript{8} The term “long term” has been defined as a follow-up of at least 5 years.\textsuperscript{2} Traditionally, life table analyses and Kaplan–Meier statistics have been applied in implant survival studies. Often patient cohorts were analyzed prospectively or retrospectively with periods up to a certain observation time. As an example, a long-term evaluation of non-submerged ITI implants reported an 8-year life table analysis in which
the mean follow-up time was 3.1 years for individual observation periods that ranged from 1 to 8 years\(^2\). The term “Survival” was defined as FDP remaining in-situ with or without modifications. “Success” was defined as the FDPs remaining in-situ free of all complications over the entire observation period. Even with follow-up periods of at least 5 years, some clinicians may argue that this is still too short to obtain reliable information on survival and complication rates\(^9\). Due to the fact that the use of dental implants for rehabilitation of partially edentulous patients is relatively new, a mean follow-up period of at least 5 years was a necessary compromise. The majority of older longitudinal studies on dental implants concentrated on implants survival and did not address the reconstructions at all. Nor did they distinguish between Different types of reconstructions such as implant-supported FDPs, implant-supported single crowns, implant-supported fixed complete dentures, combined tooth-implant-supported FDPs, or implant-supported overdentures (Figure 1)\(^3\).

**Figure 1: Annual failure rates of implant supporting FDPs after 5 years**
2. Literature Review:

2.1. History of Dental Implants
Dental implants are fabricated in a style that mimics the tooth root shape and form as alternate for lost tooth. In the beginning, dental implants were made of commercially pure titanium and made as a screw-in configuration that contained a smooth surface in between the threads. Dr. Per-Ingvar Branemark, discovered that a titanium dental implant could establish a direct host bone-to-implant connection, a phenomenon known as “osseointegration”. Osteointegration describes direct structural and functional connection between living bone and the surface of the dental implant. Normally the implant is connected to the abutment via an external/internal hex (figure 2). Particularly external hex abutment implant has had documentation for more than 30 original implant design.

**Figure 2: External and Internal hex abutment**

At the time, within the first year after implant placement the (accepted bone loss) was 1-1.5 mm, followed by 0.2 mm annually using external hex abutment implant system. Despite the long-term documentation of external hex abutment
type of implant, some major problems were recorded, mainly that there was a continuous bone loss over time\textsuperscript{12-13}. From aesthetic point of view, bone loss becomes an important issue when related to aesthetic zones that are mainly visible during smiling. Once bone loss occurs the superimposing soft tissue will follow the bone, which will result in an unsatisfactory aesthetic complication\textsuperscript{14}. Other complications that were detected in this implant system include screw loosening and/or fracture\textsuperscript{15}. An external hex configuration, displayed a hex on top of the implant to which the abutment could be connected. Thus, there have been some design modifications to the root form dental implant in order to overcome the existing limitations. These modified designs of implant aspects included the replacement of an external hex connection with an internal hex connection, as well as the roughening of the implant surface. The roughened surface was thought to increase the contact surface area of implant-to-bone in comparison to the smooth surface implant. Furthermore, this design promoted better mechanical stability following placement, better retention of the blood clot, and finally, greater stimulation of the bone healing\textsuperscript{15,16}. The internal hex connection was created to reduce the number of prosthetic complications that were experienced with the external hex system\textsuperscript{14}. Recent designs of dental implants have overcome some of the inherent problems that were experienced with the smooth surface implant by incorporating a roughened surface, as well as an internal hex abutment connection design. Long-term studies supporting the original smooth surface implants are lacking\textsuperscript{17}.

As stated earlier dental implants are widely used to restore missing teeth as well as to serve as abutments for fixed dental prostheses, removable dental prostheses and complete
dentures. In the U.S the most common causes of the loss of at least one permanent tooth are trauma, periodontitis, failed endodontics, or tooth decay and this was reported in 69% of people between ages 35 to 44 years. As a result, by age 74, 26% of adults have lost all of their permanent teeth. In 2008, the global dental implant market increased to $3.4 billion dollars, while the market for traditional crowns and bridges decreased to $4.4 billion dollars. The market value of dental implants is anticipated to reach $8.1 billion by 2015.

2.2. Major Clinical Challenges for Dental Implants

Dental implants require sufficient alveolar bone, both in width and in height, to acquire adequate primary stability, and to eventually exert its support function. In some cases such as severely atrophic edentulous mandibles and thin maxillary ridges near to the sinus floor, implant treatment is not an option without bone augmentation. In addition, bone loss also results in esthetic problems in the anterior maxilla. On the other hand, patients with implant placement should wait 3 to 6 months for successful osseointegration and final permanent restoration. Therefore, methods to augment alveolar bone and shorten the waiting time are two major clinical challenges for dental implantology.

2.3. Rationale for Dental Implant Application
Dental implants have many advantages over conventional crowns, fixed dental prostheses and removable dental prostheses. Dental implants are able to preserve adjacent tooth structure because abutment tooth preparation is not required as in single crowns and FDPs. The risk of recurrent caries in dental implants is not considered, while caries is considered to be the most frequent reason for failure of existing restorations such as onlays, crowns, and bridges. Implants can provide much more stability and retention of implant-supported prostheses than traditional tooth/tissue-borne partial dentures and tissue borne complete dentures. In addition, the most important aspect of dental implants is to preserve alveolar bone. It has been reported that marginal peri-implant bone loss over a 10-year observation period was less than 1 mm for both mandible and maxillae. Because of alveolar bone preservation, dental implants can be used to restore and maintain the gingival tissue emergence profile in the maxillary esthetic zone after anterior tooth extraction. Furthermore, the preservation of alveolar bone implants may be a key rationale for its 90% long-term survival rate. However, Schopped et al. (2003) reported the loss of alveolar bone 1 year following tooth extraction to be 6 mm in width and 1.2 mm in height.

2.4. Osteointegration

Dental implantology, is currently the most intensively developing field of dentistry. The success, survival and long-term prognosis of implant prosthetic therapy depend primarily on the anchorage of the implant to the jawbone, i.e. on osseointegration.

In the 1969s, Brånemark et al. stumbled upon the phenomenon of osteointegration when using titanium (Ti) in animal models, with little idea of the impact this discovery
would have on the rehabilitation of future medical and dental patients. “Osseointegration”, was characterized by a number of clinical and ultra-structural observations. Osseointegration may broadly be defined as the dynamic interaction and direct contact of living bone with a biocompatible implant in the absence of an interposing soft tissue layer\textsuperscript{24-26}.

Although the clinical term osseointegration describes the anchorage of endosseous implants to withstand functional loading, it provides no insight into the mechanisms of bony healing around such implants. However, in the last decade it has become clear that the long-term success of dental implants also depends on the complex bio-integration of these alloplastic materials, which is determined by the responses of the surrounding host tissues (the alveolar bone, the conjunctival part of the oral soft tissues and the gingival epithelium). Nevertheless, an understanding of the sequence of bone healing events around endosseous implants is believed to be critical in developing biologic design criteria for implant surfaces. Bone growth on the implant surface can be phenomenologically subdivided into three distinct phases that can be addressed experimentally\textsuperscript{27}. The first, \textit{osteoconduction}, relies on the migration of differentiating osteogenic cells to the implant surface, through a temporary connective tissue scaffold. Anchorage of this scaffold to the implant surface is a function of the implant surface design. The second, \textit{de novo bone formation}, results in a mineralized interfacial matrix, equivalent to that seen in cement lines in natural bone tissue, being laid down on the implant surface. The implant surface topography determines whether the interfacial bone formed is bonded to the implant. A third tissue response, \textit{bone remodelling}, creates a bone-implant interface comprising \textit{de novo} bone formation. Treatment outcomes in
dental implantology depend critically on the implant surface designs that optimize the biological response during each of these three distinct integration phases\textsuperscript{28}.

Today, much effort is devoted to the design; synthesis and fabrication of Ti dental implants in order to achieve long term secure anchoring in the bone. Basically, this means the ability of implants to carry and sustain the dynamic and static loads that they are subjected to. The bulk structure of the material governs this ability. Evidently, it is important to obtain adequate function in the shortest possible healing time, with a very small failure rate and with minimal discomfort for the patient. These factors are also important for cost reasons. A wide variety of materials have been used to produce endosseous implants\textsuperscript{28, 29}.

2.5. Implant materials:

Currently, Ti and its alloys are the most commonly utilized dental and orthopedic implant materials that meet the most important requirements\textsuperscript{30, 31}. The properties of Ti and its surface, which is covered by a native oxide layer, are appropriate to allow its use as a biocompatible material\textsuperscript{30}. At a cellular level, the relationship of an implant with the surrounding tissue is highly dependent on the interaction between the passive titanium oxide (TiO\textsubscript{2}), which is formed on the surface of a Ti implant, and biological elements such as collagen, osteoblasts, fibroblasts and blood constituents\textsuperscript{22, 31}. The TiO\textsubscript{2} layer is very stable, corrosion-resistant and may be manipulated to have variable thickness.

2.6. Success interface of osteointegration:

The clinician is often faced with the challenge of identifying successful osseointegration
of a dental implant. Clinical success is determined by a lack of mobility and by the ability of the implant to resist functional loading (chewing force) without mechanical deformation and to transfer the load to the alveolar bone without deterioration of the bony interface\textsuperscript{32}. Radiographically, the bone should appear to be closely apposed to the implant surface. The resolution currently achievable in medical imaging, however, is several orders of magnitude less than what is required to observe a soft tissue cell. Accordingly, radiographic assessment alone is unsuitable to determine with certainty whether soft tissue is present\textsuperscript{33}. A number of studies have analyzed this bone to Ti interface histologically and ultra-structurally, often with inconsistent findings. The difficulty arises primarily with the need to prepare and section the specimens without changing or damaging the interface. Recent studies have utilized CT scanning to obtain a 3-dimensional picture of the implant interface\textsuperscript{34,35}.

2.7. Implant success and failure criteria:

Marginal bone loss considered as a key criterion of long-term success following dental implants. Since bone-anchored prostheses are constructed in the oral environment for a lifetime, a pathologic decrease of marginal bone could lead to loss of bone anchorage of the implant, and it is important to know what factors causing the bone resorption. In general, reasons for bone loss can be divided into the following categories: The size of the implants, age and gender of patients, and the presence of cantilevers as influencing factors are subject of debate\textsuperscript{36}. Recent studies claimed that smoking was a factor in identifying a subject with progressive bone loss around an implant with 69% accuracy\textsuperscript{37}. In a long-term follow-up of implant treatment, it was concluded that patients who had a
history of periodontal disease and patients that smoked are more likely to develop bone loss around implants\textsuperscript{38}. Patient’s ability in performing proper oral hygiene procedures has also been shown to influence bone levels around implants\textsuperscript{37}. Furthermore, occlusion is a primary factor for implants success in the long run. Thus, occlusal overload can lead to mechanical stresses on both dental implants and implant prostheses. There are a myriad of variables in a patient population, so no one occlusal scheme can fit all implant patients. Thus, if a clinical condition is likely to increase biomechanical stresses, dentists should implement occlusal mechanisms to decrease the stresses and develop an occlusal scheme that minimizes risk factors and allows the restoration to function in harmony with the rest of the stomatognathic system. This is what we call implant-protected occlusion\textsuperscript{39}.

Occlusal overload is often regarded as one of the main causes for peri-implant bone loss and implant prosthesis failure, because it can cause crestal bone loss, thus increasing the anaerobic sulcus depth and peri-implant disease states if patients cannot clean well. So a harmonious occlusal scheme is a primary objective\textsuperscript{40}.

\textbf{2.8. Differences between natural teeth and implants}

The basic difference between natural teeth and endosseous dental implants is that the natural tooth has a support design that reduces the force to the surrounding crest of bone compared to the same region around an implant. Natural tooth is suspended by the periodontal ligament (PDL), while an endosseous dental implant is in direct contact with the bone through osseointegration. The PDL absorbs shocks and distributes occlusal
stresses away along the axis of natural teeth. However, an endosseous dental implant connected to the bone by osseointegration lacks those advantages of the PDL\textsuperscript{41} (Table 1).

Periodontal tissues are uniquely innervated and structured to retain and support teeth. When natural teeth are lost, both occlusion and attachment with its proprioceptive feedback mechanism are lost. When loaded, the distribution of forces begins with the primary phase of periodontal compliance that is primarily non-linear and complex, followed by the secondary movement phase, which occurs with involvement of the alveolar bone. In contrast, the movement of an implant under loading is dependent on linear and elastic deformation of the bone. The PDL in a natural tooth can produce differences in force adaptation compared with osseointegrated implants due to its shock-absorbing and stress-distributing functions Table 1\textsuperscript{41}. 
Table 1: Natural tooth Vs. Implants

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Tooth</th>
<th>Implant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Composition</td>
<td>Calcium and phosphorus (hydroxyapatite)</td>
<td>Primarily titanium and titanium based alloys</td>
</tr>
<tr>
<td>Nature</td>
<td>Living</td>
<td>Nonliving</td>
</tr>
<tr>
<td>Gingival sulcus depth</td>
<td>Shallow</td>
<td>Depends upon abutment length and restoration margin</td>
</tr>
<tr>
<td>Junctional epithelium</td>
<td>On enamel</td>
<td>On titanium</td>
</tr>
<tr>
<td>Connectivity issue</td>
<td>Perpendicular to tooth surfaces</td>
<td>Parallel and circular fibers; no attachment to implant or bone attachment</td>
</tr>
<tr>
<td>Gingival fibers</td>
<td>Complex array inserted into cementum above crestal bone</td>
<td>No organized collagen fiber attachment</td>
</tr>
<tr>
<td>Crest of bone</td>
<td>1 to 2 mm apical to cementoenamel junction</td>
<td>According to implant design</td>
</tr>
<tr>
<td>Nerve supply</td>
<td>Present</td>
<td>Absent</td>
</tr>
<tr>
<td>Proprioception</td>
<td>Highly sensitive</td>
<td>No ligament receptors</td>
</tr>
<tr>
<td>Physical characteristics</td>
<td>Physiologic mobility caused by viscoelastic properties of the ligament</td>
<td>Rigid connection to bone, as if ankylosed</td>
</tr>
<tr>
<td>Adaptive characteristics</td>
<td>Width of ligament can alter to allow more mobility with increased occlusal forces</td>
<td>No adaptive capacity to allow mobility; orthodontic movement impossible</td>
</tr>
<tr>
<td>Connection</td>
<td>Cementum, bone, periodontium</td>
<td>Osseointegration, bone functional ankylosis ligament</td>
</tr>
<tr>
<td>Junctional epithelium</td>
<td>Lamina lucida and lucida, lamina dense zones</td>
<td>Lamina, lamina densa, and sublamina lucida zones</td>
</tr>
<tr>
<td>Connective tissue</td>
<td>Thirteen groups: perpendicular to tooth surfaces</td>
<td>Two groups: parallel and circular fibers</td>
</tr>
<tr>
<td>Biological width</td>
<td>2.04 to 2.91 mm</td>
<td>Increased collagen, decreased fibroblasts</td>
</tr>
<tr>
<td>Vascularity</td>
<td>Greater, supraperiosteal and periodontal ligament</td>
<td>Less, periosteal</td>
</tr>
<tr>
<td>Probing depth</td>
<td>3 mm in health</td>
<td>2.5 to 5.00 mm</td>
</tr>
<tr>
<td>Bleeding on probing</td>
<td>More reliable</td>
<td>Less reliable</td>
</tr>
</tbody>
</table>

Non-vertical forces are potentially harmful to natural teeth, but during function can cause damage to teeth and the PDL. Non-axial or non-vertical forces with lead to primary trauma from occlusion manifest as a widen PDL and hypermobile tooth. Whereas in implants, the non-axial loads involves the crest of the bone, which is usually traumatic to the supporting structures. According to Parfitt, a lateral force on a healthy natural tooth is rapidly dissipated away from the crest of bone toward the apex of the tooth due to the natural tooth rapidly moving 56~108 μm and rotating around the apical 1/3 of the root. On the other hand, movement of an implant occurs gradually, reaching up to about 10~50 μm under a similar lateral force. So greater forces are concentrated on the crest of the surrounding bone of dental implants in the absence of rotation. Under similar lateral
loads, an implant does not pivot as much as a tooth toward the apex, but instead concentrates greater forces at the crest of the surrounding bone. Therefore, if an initial load of equal magnitude and direction is placed on both an implant and natural tooth, the implant must be protected.

2.9. Implant-supported cantilever fixed dental prosthesis:

Implant-supported cantilever prosthesis is the pontic attached to the implant crown. In daily practice, treatment-planning decisions are influenced by many technical and biological factors that increase the challenge of making the final treatment plan. A further consideration is the cost-benefit ratio of implants treatment. With regard to prosthesis rehabilitation, implant-supported cantilevers provide simpler rehabilitation procedures compared to surgical reconstructive procedures. The inclusion of a cantilever may be a simpler option in situations where local conditions of the residual edentulous ridge preclude the possibility to place an implant. This is preferable than surgical reconstructive procedures because of long treatment time, high morbidity and increasing cost. Cantilevers were used to extend implant supported full arch FDPs since the early applications of the Branemark method, with promising long-term results. Since then, the use of cantilevers in full arch, multiple unit or even single unit FDPs has been relatively common in implant reconstructions. In a cohort of periodontally treated and maintained patients at Berne University, implant-supported cantilever FDPs (cFDPs) represented 8.6% of all FDPs and 27.78% of implant-supported FDPs. In other groups, cFDPs represented 6.9%, 12.35%, 20.53%, and 67% of implant-supported FDPs. However, it has been claimed that cantilever extensions increase the risk of flexion
overload and that this in turn may compromise the prognosis of the prosthetic rehabilitation and the survival of the implants\textsuperscript{48}. It has been demonstrated longer cantilevers resulted in higher stress at implant sites, thus triggering greater marginal bone loss around implants. Consequently, a higher incidence of biologic and prosthetic complications was expected for this type of rehabilitation, when compared with those without cantilevers. Cantilevers with less-favorable crown/implant ratios can increase the possibility of overloading, possibly resulting in peri-implant bone loss and prosthesis failure\textsuperscript{49}. In terms of cantilever length, a clinical study demonstrated that long cantilevers (≥ 15 mm) induced more implant-prostheses failures compared to cantilevers < 15 mm long\textsuperscript{50}. Duyck et al\textsuperscript{51}; also reported that when a biting force was applied to a distal cantilever, the highest axial forces and bending movements were recorded on the distal implants, which were more pronounced in prostheses supported by only 3 implants, compared to prostheses with 5 or 6 implants. The above study indicated that a shorter cantilever length is more favorable for the success of implant-supported prostheses, particularly for prostheses with fewer implants\textsuperscript{52}. The occlusal contact position can determine the direction of force, which may result in overloading of supporting implants, especially during parafunction\textsuperscript{53}. Cantilevers can cause screw loosening and/or prosthetic screw or abutment screw breakage and should be eliminated. Therefore, periodic evaluation of occlusion is necessary\textsuperscript{52}. Other studies showed equal success rates in cases with or without a cantilever. Thus, the use of cantilevered restorations is controversial and requires clear evidence for their application. Because the use of cantilevered implant restorations could still be a viable treatment option in many instances, their effect on marginal bone loss and the survival rate of the supporting implants and the prosthetic
maintenance should be clarified. Previous studies, reported a high number of technical complications that might be expected in the presence of a cantilever extension. On the contrary, findings from the statistical analysis showed that there were no differences in prosthetic complications between cantilever and non-cantilever prostheses. Previous systematic reviews pointed out the same issue; technical complications are common for either implant or tooth-supported fixed partial prostheses, independently of the presence or absence of cantilevers. Factors such as the type of restoration used or the number of implants supporting the cantilever in the included studies might justify this observation. Additionally, studies have observed the relationship between prosthetic complications, such as implant fracture, and the combination of cantilevers with bruxism, heavy occlusal forces, or this relationship suggests that besides the presence of a cantilever extension, there are other factors that may influence the appearance of prosthetic complications. Further biases that may cause misleading results are that marginal bone loss could be influenced by many factors not limited only to the presence/absence of a cantilever extension. Another confounding factor that might be considered when assessing peri-implant bone loss is the crown-implant ratio. More recently, it have been demonstrated that prosthesis height does not influence the crestal stress concentration, as displayed by no significantly greater marginal bone loss. Thus, assessing contributing factors for marginal bone loss display show limitations because many confounding elements are may be present and thus difficult to assess. In the aesthetic region in particular, the use of single implant cFDP may help resolve compromising situations, when a limited mesio-distal dimension may jeopardize aesthetics. In such cases inter-implant distance of >3 mm is recommended to minimize...
crestal bone loss in between implants\textsuperscript{61}, which achieved papillary height of 3–4 mm at most of the time (Tarnow 2003)\textsuperscript{62}. The use of cantilevers, however, has not been without controversy. Some authors have suggested that occlusal forces on cantilevers are amplified by leverage action, which might result in damaging strain. Finite element analysis (FEA) and in vitro studies have shown cantilever FDPs to develop highest stress/strain levels\textsuperscript{63}, which have been shown to increase with cantilever length above 7 mm\textsuperscript{64}. This increased strain has been hypothesized to have a detrimental effect not only on the longevity of the FDP but also on the peri-implant bone and osseointegration and some authors have discouraged the use of cantilevers. However, no clear evidence of the impact of cantilevers on bone and peri-implant tissues has been documented in animal studies, in the presence of adequate plaque control\textsuperscript{65}. Systematic reviews showed failure rate per 100 cFDP years of 1.18\textsuperscript{66} comparable to non-cantilever FDPs at 1.03\textsuperscript{43} and no evidence for increase marginal bone loss or biological complications\textsuperscript{66}. Furthermore, clinical studies have shown little statistical or clinical difference in the biological outcomes in full arch prosthesis with distal cantilevers\textsuperscript{43} as well as multi-unit FPDs\textsuperscript{68}, in medium term follow-up of 4–10 years. With respect to studies involving technical complications, higher complication rate appears to occur in cFDPs\textsuperscript{45}, although few authors have shown higher success rate for cFDPs\textsuperscript{57}. A study showed that much of this increased technical failure is related to minor screw loosening and veneer fracture\textsuperscript{55}. There is lack of reporting of biological complications involving clinical examinations. Some clinical studies showed a trend but lacked statistical power to detect differences in mean marginal bone loss smaller than 1 mm\textsuperscript{57,69} or lacked cases with >5-year follow-up or appropriate analysis for drop out\textsuperscript{68}. With respect to technical complications, there is
considerable variability in the outcomes reported\textsuperscript{66}, even when published by the same group. Furthermore, the impact of variables such as cantilever dimensions, number of implants, occlusal scheme and dimensions of prosthesis on survival and success rate remain underexplored\textsuperscript{66}. 
3. **Aim:**

   (1) To determine if bone height changes are different on mesial and distal surface of implant adjacent to cantilevered crown units.

   (2) To study the impact of cantilever direction on technical outcomes of implant-supported fixed dental prostheses (FDPs) and bone height.
4. Material and methods:

4.1. Design: Retrospective study:

This study is a retrospective review of the records of 14 partially dentate patients who were restored with dental implants placed by residents, periodontists and prosthodontists working in Dubai Health Authority and Hamdan bin Mohammed Dental College.

4.2. Population and location of study:

Fourteen patients (7 males and 7 females), aged 45-83 years (mean age 69.42 years), treated from March 2003 to March 2015 with implant-supported FDPs with a cantilever extension were included in the study. The implant sites included all mandibular and maxillary positions. Data recorded for all patients included the patient’s: age, gender and smoking status. The implant data included: the diameter and length of the implant, age and region (FDI). The prosthesis data included: number, direction and length of cantilever pontics, and age.

Only 4 patients (3 males and 1 female) were treated in Hamdan bin Mohammed Dental College and all the others were treated in Albeda dental center, in Dubai Health Authority. The patients received 28 bridges, 18 with a distal cantilever and 10 with a mesial cantilever supported by at least two implants (some patients received more than one implant).
4.3. Inclusion criteria:

- Partial edentulism
- Implant-supported fixed dental prosthesis with at least one cantilever unit, which has been under functional loading a minimum of one year
- Smokers and non-smokers
- Controlled diabetes and Controlled hypertension

4.4. Exclusion criteria:

- Systemic diseases (such as heart, coagulation, and leukocyte diseases or metabolic disorders)
- A history of radiation therapy in the head and neck region.
- Current treatment with steroids
- Neurological or psychiatric handicap that could interfere with good oral hygiene
- Immuno-compromised status, including infection with HIV
- Severe clenching or bruxism
- Full-arch fixed prostheses
- Drug or alcohol abuse
- Poor attendance record.

4.5. Method of Data collection:

Data collection was carried out from patient records available in the software systems of Dubai Health Authority and Hamdan Bin Mohammed Dental College.
4.6. Sampling measures:

All the patients who had treatment at Dubai Health Authority and Hamdan bin Mohammed Dental College from March 2003 to March 2015 for implant-supported cantilever prostheses and met the inclusion criteria were selected for the study. Patient profiles included: age, gender, medical history, smoking status), implant evaluation (implant length, implant diameter, date of placement), as well as cantilever fixed dental prosthesis supported by implant (cantilever pontics number, cantilever length, date of loading and cantilever direction) were recorded.

Radiographic images were analyzed with a software program (Digora) to measure the following parameters:

- Cantilever length: from the most outer point of adjacent implant abutment crown to the most distal or mesial point of the cantilever pontic (figure 3)

- Peri-implant bone height: from the implant abutment to the crest of the peri implants bone. The implant that is adjacent to the cantilever pontic, on both mesial and distal aspects of the implant was assessed. Intraoral peri-apical radiographs were measured, first at baseline (time of implant loading) and second at the last follow-up appointment evaluation.

Bone loss was measured around the implant adjacent to the cantilever pontic to serve as a standard and compared to the actual implant length to correct for magnification error (figure 3). Bone loss = (Actual implant length x Distance from abutment to bone crest / Radiographical implant length)
Figure 3: Radiograph of cFDP showing measurement landmarks  (BL: bone loss, CL: cantilever length)
4.7. Ethical Approval:

The Scientific Research Ethics Committee of Dubai Health Authority approved the study protocol. (Appendix 1)
5. **Results**

Fourteen patients were included in the study (7 males and 7 females), with a mean age of 69.42(9.03) years. The age range of the patients was between 45-83 years. All patients were rehabilitated with cantilever implant-supported fixed dental prostheses. A total of 44 implants were placed with a mean age of 3.72 years SD=(2.51). The implants supported 28 bridges with a mean age of 2.36 years SD= (2.59). The bridges consisted of 76 units including 32 cantilever units (Table 2).

Therefore, some of the bridges had multiple implants and more than 1 cantilever pontic. Most of the bridges, however, had just a single pontic. In addition, some of the patients had more than one cantilever fixed dental prosthesis.
Table 2: Descriptive variables in the study

<table>
<thead>
<tr>
<th>Items</th>
<th>No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. Of patients (male/female)</td>
<td>14(7/7)</td>
</tr>
<tr>
<td>No. Of implants</td>
<td>44</td>
</tr>
<tr>
<td>No. Of Bridges</td>
<td>28</td>
</tr>
<tr>
<td>No. Of Cantilever units</td>
<td>32</td>
</tr>
<tr>
<td>Mean (SD) age of patient</td>
<td>69.42 (9.03)</td>
</tr>
<tr>
<td>Mean (SD) age of bridges</td>
<td>2.36 (2.59)</td>
</tr>
<tr>
<td>Mean (SD) age of implant</td>
<td>3.72 (2.51)</td>
</tr>
</tbody>
</table>

All implant material was titanium and all cantilever fixed dental prostheses were metal-ceramic. Twenty-eight cantilever implant-supported fixed dental prostheses were evaluated radiographically. The majority of the cantilever pontics were distally directed 23(71.9%), while 9(28.1%) were mesially directed.

Ten (35.7%) cantilever FDPs were located in the upper right quadrant and 10(35.7%) in upper left quadrant. While 5(17.9%) located in lower right quadrant and 3(10.7%) in lower left quadrant. Table 3 shows the distribution of implants according to their position in the jaw (Table 3).
Table 3: Distribution of implant-supported cantilever prosthesis according to their position in the jaw (N=28)

<table>
<thead>
<tr>
<th>Region implant/cantilever</th>
<th>No. (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lower left</td>
<td>3 (10.7)</td>
</tr>
<tr>
<td>Upper left</td>
<td>10 (35.7)</td>
</tr>
<tr>
<td>Lower right</td>
<td>5 (17.9)</td>
</tr>
<tr>
<td>Upper right</td>
<td>10 (35.7)</td>
</tr>
</tbody>
</table>

Smoking habits are shown in (Figure 4). Most of the cases were non-smokers 43%, while 29% were not recorded.

Figure 4: Smoking status

Diagram showing smoking status distribution with non-smokers 43%, smokers 28%, and not-recorded 29%.
5.1. Technical Complications:

Technical complications were recorded at all ages between 45-83 years. At prosthesis level, technical complications were recorded in 4 FDPs: Two with prosthesis loosening (7.1%) and 2 with de-cementation (7.1%). The 2 cases of prosthesis loosening occurred in males while prosthesis de-cementation occurred in 2 females. Two cases had de-cementation of the prosthesis (1 in upper right and 1 in upper left). While loosening occurred twice, both were in the lower jaw, 1 case in lower right and 1 in lower left (Table 4).

One of the 2 cases occurred in male, smoker, 9.5mm implant length and 3.5mm implant diameter and located in lower left quadrant. And the second case occurred in female patient, non-smoker, 11mm length of implant and 3.5mm implant diameter. Both were distal cantilevers, 1-year age of cantilever bridge and 2 years age of implant. In addition, de-cementation occurred in 2 cases, first case in male patient, non-smoker, 11mm length of implant and 3.5mm implant diameter, located in upper right quadrant and age of cantilever prosthesis and implant were 12-years and 13 years respectively. Second case of de-cementation was female patient, not-recorded smoking status; age of implant was 7 years and 6 years of cantilever prosthesis, located in upper left quadrant. Both were distal cantilever.
Table 4: Technical complications of bridges (N=28)

<table>
<thead>
<tr>
<th>Technical complications</th>
<th>Absent (%)</th>
<th>Present (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prosthesis loosening</td>
<td>26 (92.9)</td>
<td>2 (7.1)</td>
</tr>
<tr>
<td>De-cementation</td>
<td>26 (92.9)</td>
<td>2 (7.1)</td>
</tr>
<tr>
<td>Veneer chipping</td>
<td>28 (100)</td>
<td>0</td>
</tr>
<tr>
<td>Framework fracture</td>
<td>28 (100)</td>
<td>0</td>
</tr>
</tbody>
</table>

At the implant level, technical complications were mainly associated with implants of 9.5mm length and 3.5mm width except 1 case, which was 11mm implant length and 3.5mm width.

All screw-loosening of implant abutments occurred in males and in the upper jaw; 2 in right quadrant and 1 in left quadrant. From a total of 44 implants, only 3(6.8%) of implants had abutment-screw loosening (Table 5). Abutment screw-loosening occurred in 3 implants, all of them are in male patients, 1 was a smoker and the other 2 patients were not recorded smoking status. All of them had equal implant diameter (3.5mm) but different implant lengths. The cantilever length was between 5-9 mm, in all FDPs with implant abutment-screw-loosening.

All cases, which had technical complications, were distal cantilever prostheses. First case was 11mm length, in a non-smoker, placed in the upper right quadrant, cantilever prosthesis was 12 years old and implant was 13 years old. While Second and third cases
have almost the same characteristics, both had 9.5mm implant length and 3.5 mm implant diameter, distal direction of cantilever implant-supported prosthesis, but the smoking status was not-recorded. In both cases implants were placed in 2011 and loaded in the same year. Both cases were placed in the upper jaw, one in the right and one in the left quadrant.

Table 5: Technical complications of implants (N=44)

<table>
<thead>
<tr>
<th>Technical complications</th>
<th>Absent (%)</th>
<th>Present (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abutment screw-loosening</td>
<td>41 (93.2)</td>
<td>3 (6.8)</td>
</tr>
<tr>
<td>Implant screw-fracture</td>
<td>44 (100)</td>
<td>0</td>
</tr>
<tr>
<td>Implant fracture</td>
<td>44 (100)</td>
<td>0</td>
</tr>
</tbody>
</table>
Table 6: Comparison of implant screw-loosening between mesial and distal abutments on bridges according to whether 2 units or > 2 units fixed dental prostheses and cantilever direction

<table>
<thead>
<tr>
<th>Technical complication</th>
<th>Distal Cantilever Direction</th>
<th>Mesial Cantilever Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2 units FDP</td>
<td>&gt;2 units FDP</td>
</tr>
<tr>
<td>Implant Screw-loosening</td>
<td>Present</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Absent</td>
<td>1</td>
</tr>
<tr>
<td>P-value</td>
<td>0.371</td>
<td>X² = 0.8</td>
</tr>
</tbody>
</table>

Table 7: Comparison of prosthesis de-cementation between mesial and distal cantilever prostheses:

<table>
<thead>
<tr>
<th>Technical complication</th>
<th>Distal Cantilever Direction</th>
<th>Mesial Cantilever Direction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2 units FDP</td>
<td>&gt;2units FDP</td>
</tr>
<tr>
<td>Prosthesis Decementation</td>
<td>Present</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Absent</td>
<td>4</td>
</tr>
<tr>
<td>P-value</td>
<td>0.7353</td>
<td>X² = 0.1148</td>
</tr>
</tbody>
</table>
### Table 8: Comparison of prosthesis loosening between mesial and distal cantilever prostheses:

<table>
<thead>
<tr>
<th>Technical complication</th>
<th>Distal</th>
<th>Mesial</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2 units FDP</td>
<td>&gt;2 units FDP</td>
</tr>
<tr>
<td>Prosthesis loosening</td>
<td>Present</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Absent</td>
<td>4</td>
</tr>
</tbody>
</table>

P-value 0.7353

$X^2 = 0.1148$

Most of the technical complications 5(71.4%) were associated with single cantilever pontic bridges supported by double implants, in contrast 1 case of prosthesis loosening occurred in single implant abutment supporting double cantilever pontics and 1 case of (implant abutment screw-loosening) occurred in single implant supporting single cantilever pontic. In addition, no complication occurred with double implants supporting a double cantilever (Figure 5). Nevertheless, all technical complication at prosthesis level occurred in FDPs with < 10 mm cantilever length (prosthesis de-cementation and prosthesis loosening).
Figure 5: Technical complication in relation to implant/cantilever number

- Implant abutment screw-loosening
- Prosthesis de-cementation
- Prosthesis loosening
Figure 6: Radiographs shows single implant supporting double cantilever pontics (left), single implant supporting single cantilever pontic (middle) and image of double implants supporting single cantilever pontic (right)
5.2. Bone height changes:

From the total of 28 bridges, only 22 were available for radiographical analysis of bone height, while the rest of the patients missed their follow-up appointments and some of the follow-up radiographs were not clear. Marginal bone level was measured around implants at the time of prosthesis loading and at the last recorded recall appointment (Table 9).

Bone loss was tested for normality by using Kolmogorov-Sminov test. The data showed normal distribution of the bone loss data around both sides of implants adjacent to mesial and distal cantilevered pontics. Univariate analysis of variance was used to test for significance of bone loss around implants adjacent to mesially or distally directed cantilever and number of implant(s) (single or multiple).
Table 9: Marginal bone height from time of cantilever prostheses loading to the last recall visit
Table 10: Bone Loss Around Implants Supporting Distal Cantilever

<table>
<thead>
<tr>
<th>Direction</th>
<th>Distal side of implant</th>
<th>Mesial side of implant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Implanta</td>
<td>N</td>
<td>Mean</td>
</tr>
<tr>
<td>Single</td>
<td>4</td>
<td>0.35</td>
</tr>
<tr>
<td>Multiple</td>
<td>8</td>
<td>0.93</td>
</tr>
<tr>
<td>Total</td>
<td>12</td>
<td>0.74</td>
</tr>
</tbody>
</table>

The mean of bone loss at distal side is lower than mesial side of implant supporting distal cantilevers (Table 10). However, there was no significant difference on bone loss at mesial and distal sides of implants adjacent to distal cantilivered pontic and (P= 0.82).

Table 11: Bone Loss Around implants Supporting Mesial Cantilever

<table>
<thead>
<tr>
<th>Direction</th>
<th>Distal side of implant</th>
<th>Mesial side of implant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Implanta</td>
<td>N</td>
<td>Mean</td>
</tr>
<tr>
<td>Single</td>
<td>7</td>
<td>0.73</td>
</tr>
<tr>
<td>Multiple</td>
<td>3</td>
<td>1.01</td>
</tr>
<tr>
<td>Total</td>
<td>10</td>
<td>0.92</td>
</tr>
</tbody>
</table>

No significant difference was detected on bone loss with both sides mesial and distal around implants adjacent to mesial cantilevered pontic (P = 0.533). No statistical significant difference in marginal bone loss between mesial cantilever (P- value 0.53), and distal cantilever (P-value 0.82) on both sides of implant ( Table 11).
Table 12: Comparison of marginal bone loss between mesial and distal cantilevers on mesial and distal sides of implant supporting cantilever unit(s)

<table>
<thead>
<tr>
<th>Cantilever direction</th>
<th>Mesial cantilever</th>
<th>Distal cantilever</th>
</tr>
</thead>
<tbody>
<tr>
<td>Implant-supporting cantilever</td>
<td>Distal side of implant</td>
<td>Mesial side of implant</td>
</tr>
<tr>
<td>Mean</td>
<td>0.91</td>
<td>0.76</td>
</tr>
<tr>
<td>SD</td>
<td>0.84</td>
<td>0.76</td>
</tr>
<tr>
<td>P-value</td>
<td>0.67</td>
<td></td>
</tr>
</tbody>
</table>

There was 1 distal cantilever, which showed marginal bone loss >1 mm (1.52) and mesial and distal sides of implant adjacent to cantilever and technical complication (prosthesis loosening) in smoker male 73 years old. While 1 female 57 years old, non-smoker with no technical complications and 2 FDPs showed marginal bone loss > 1 mm, the distal cantilever (mesial: 2.17mm, distal: 3.08mm) marginal bone loss, and the mesial cantilever showed marginal bone loss (mesial: 2.23mm, distal: 0.58mm). Third case was female 62 years old, non-smoker with mesial cantilever, marginal bone loss was 1.19 mm in mesial and distal sides of implant supporting the cantilever unit.

All cases which recorded marginal bone loss > 1 mm occurred in prosthesis supported by implant with 9.5 mm length and 3.5 mm width. Two out of three cases showed bone loss
occurred in non-smoker patient. All marginal bone loss occurred in different age group, which showed no relation between patient age and marginal bone loss rate.

The number of implants placed to support the prosthesis and number of cantilevered pontics number (at least 1 implant) appeared to have significant impact on bone height. More than 1.2 mm bone loss was recorded around 1 implant supported 2-cantilever pontics. As well as, 2 technical complication of prosthesis (de-cementation, prosthesis loosening) recorded in (1 implant supported 2-cantilevers).
5.3. Gender:

Both technical complications of prosthesis loosing; 2 cases and De-cementation; 2 cases exist equally among gender. For abutment screw loosening the 3 cases were a male.

5.4. Age:

De-cementation complication occur on the youngest age 45 and 59 years

Prosthesis loosing occur on the above elderly group age 62 and 73 years old

For the screw loosening the 3 cases had lower elderly age 54, 59 and 67

5.5. Region:

For the abutment screw loosening all of them in maxilla 2 on right and 1 on left side

All of them were distal cantilevers

For the Decementation cases two were on maxilla and one on the right and one on the left

5.6. Implant Length and Diameter:

For the abutment screw loosening the average length is 10 (0.87) and the average of the diameter was 3.67 (0.289).

For the Decementation the average of the length of implant was 11 mm and the average of diameter was 4 mm.
For the prosthesis loosening the average length of implant was 10 mm and the average of diameter was 4 mm.

5.7. Age Of Cantilever prosthesis:

For the abutment screw loosening the average age of cantilever was 7 (0.33) and the average age of implant was 7.67 (4.62).

For the Decementation the average age of cantilever was 9 (4.25) and the average age of implant was 10 (4.24).

For the prosthesis loosening the average age of cantilever was 1 year and the average age of implant was 2 years.
6. Discussion:

Albrektsson et al\textsuperscript{12} reported 1mm peri-implant bone loss during the first year of function, followed by an annual loss <0.2mm after the first year in service as a criteria for implant success. Albrektsson & Isidor\textsuperscript{13} also proposed the criterion for implant success and they suggested an average peri-implant marginal bone loss of less than 1.5 mm the first year after insertion of the prosthesis and less than 0.2 mm annual bone loss after that as a standard for successful therapy. In the present study implants had been in function for a mean of 48 months, minimal marginal peri-implant bone loss was reported with average less than 2 mm, which is considered as insignificant. Aligetta et al.\textsuperscript{70} reported that the majority of studies showed that presence of cantilever extensions does not affect negatively on marginal bone height\textsuperscript{70}. Although the data was limited, drawing a definitive conclusion from such study in problematic, as well as, other factors should be considered important such as: periodontal status, smoking status and the presence of adequate plaque control. In general, causes of marginal bone loss depend on patient factors and implant design factors. Nevertheless, smoking had a determined effect on peri-implant bone loss\textsuperscript{39}. In contrast, in the present study, none of the smokers showed more that 1 mm bone loss or any technical complication, although 29% of the cases were not reported and most probably they are smokers (figure 4).

In our study, the radiographical examination was done on the existing radiographs at the time of implant loading and at last follow-up visit. Most of the patients who completed their treatment for more than 5 years are not attending regularly for follow-up visits. There are many reasons for missing follow-up visits at Dubai Health Authority. First,
most of the patients are elderly, poorly educated, and not aware of the importance of maintenance and follow-up visits. Second, since they do not have pain or symptoms of discomfort, they refused to attend for maintenance.

Smoking tobacco has been shown to have a deleterious effect on both osseointegration and crestal bone loss, Bain 1996\textsuperscript{71}. Our results indicate that smokers do not show more crestal bone loss than non-smokers, however, these finding should be interpreted with caution, as only 22.2\% of the implants included in our study were placed in non-smokers.

It has been demonstrated that a higher incidence of biologic and prosthetic complications was expected for cantilever FDPs, when compared with those without cantilevers\textsuperscript{72}, as claimed that cantilever extension increases the risk of flexion overload, and consequently compromise the prognosis of prosthetic rehabilitation. Further, in term of cantilever length. A clinical study demonstrated that long cantilevers (≥ 15 mm) induced more implant-prostheses failures compared to cantilevers < 15 mm long\textsuperscript{50}. The above study indicated that a shorter cantilever length is more favorable for the success of implant-supported prostheses, particularly for prostheses with fewer implants. In current study, all technical complication recorded with cantilever length less than 10 mm and (2-cantilever supported by 1 implant).

The present data indicated that technical complications are uncommon with cantilever FDPs. The present study demonstrated that cantilever implant-supported fixed dental prosthesis directed mesial/distal to implant restoration causes minor technical complication such as abutment screw-loosening, prosthesis de-cementation and prosthesis loosening. Further, no implant failure has been recorded. In present data the
number of patients was limited, which has equally distributed, 7 males and 7 females. Screw-loosening of abutments N= 3 (0.7%) and prostheses de-cementation N= 2 (7.4%), occurred in implants placed at least 5 years beforehand. On the other hand, 2 cases that showed prosthesis loosening, the implants were placed in 2014 and loaded in 2015. These results corroborate the most frequently cited technical complications for cantilever implant-supported fixed dental prostheses, which were veneer fracture, screw loosening and loss of retention55. It has been shown that the more presence of a cantilever extension does not increase the mechanical/technical risks for implants supporting short span cantilever FDPs3. According to our observations, distal or mesial bone loss on implant adjacent to cantilevered units was not statistically different if the cantilever pontic mesial or distal directed.

The marginal bone loss could be influenced by many factors and that it is not limited only to the presence/absence of a cantilever extension45,54. Another confounding factor that might be considered when assessing peri-implant bone loss is the crown-implant ratio56, which leads to misleading results. Recent studies claimed that prosthesis height does not influence the crestal stress concentration, as displayed by no significantly greater marginal bone loss58. Thus, assessing contributing factors for marginal bone loss display show limitations because many confounding elements are may be present and thus difficult to assess60. Our results also suggest that the use of single-unit cantilevered extensions in the distal aspect of the implant restorations may influence technical complications at implant and prostheses level. Only one implant placed in 2011 and loaded in 2012 showed a mesial bone loss in 2015 around 1 implant and the grade III mobility. The reason as recorded was poor oral hygiene. With maintenance, the mobility
reduced to grade I at last follow up visit. Also from the ethical point of view, it is not allowed to take further radiograph for research analysis purpose so that not all cases were measured accurately. Moreover, regarding the data that collected from Dubai Health Care Authority most of the cases are done in periodontics department so that most of the prosthetical details are missing or not recorded. It is difficult to collect all required information in the retrospective study because of missing data in the clinical notes. These records should be viewed with caution since our study did not report on large sample size and our data depended mainly on the clinical records, which were not well recorded.

In the present study, no significant bone loss was formed for implants with a mean age of 48 months. In cases that showed around 2 mm bone loss; this 2 mm bone loss was a cumulative amount after 2 years of loading cantilever prostheses. While the majority showed no excessive marginal bone loss, which was in agreement with some studies, particularly Aglietta et al. 2012\textsuperscript{70}, bone level changes in the posterior maxilla and mandible and also concluded no difference between cantilever and non-cantilever implants, even when small FDPs were compared with implants supporting single crowns with cantilever.

Stress distribution in the surrounding bone has been reported to depend on the dimensions of the implant\textsuperscript{73}, which directly affects the area of possible bone retention. It has been recommended that implants to be as long and as wide as possible within the anatomic limitations of the patient\textsuperscript{74}. Although a number of investigators reported higher rates of bone loss in shorter implant\textsuperscript{75} length and the importance of implant length in sustaining the load of fixed restorations has been pointed out\textsuperscript{76}. There are also studies
where implant length was reported to have little influence on the amount of stress in vertical loading and to have a smaller effect on stress distribution in the bone than the implant diameter\textsuperscript{77}, finite element method (FEM) analysis of implants has shown stress reduction with increasing diameter and radiographically measured bone loss for 4.0-mm diameter implants has been found to be less than 3.5-mm\textsuperscript{14}. However, in the present study, 9.5 mm implant length and 3.5 mm width of the implant caused a significant impact on MBL as well as the technical complications.

No major detrimental effects with respect to peri-implant tissues were observed at implants in the proximity of cantilever extensions. However, in another recent systematic review it was pointed out that the incorporation of cantilevers into implant-borne prostheses might be associated with a higher incidence of minor technical complications\textsuperscript{78}. 

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7. Conclusion:

(1) Within the limitation of this study, marginal bone loss does not seem to be influenced by the presence of cantilever extensions mesial or distal.

(2) Moreover, minor implant technical complications were found when a distal cantilever was present (screw-loosening, prosthesis de-cementation and prosthesis loosening).

(3) Most of technical complications were associated with 9.5mm implant length and 3.5mm width, which represent the majority of included implants.
8. Data collection: Case Report Form

8.1. Personal profile

- Age:
- Gender:
- Medical record:
- Smoker/ non-smoker:
- Para functional habits:
- Periodontal status:

8.2. Implant evaluation

- Implant position & region: FDI#
- Number of implant(s):
- Implant material: Titanium/ ceramic / others
- Implant restoration fixation: screw-retained/ cemented
- Implant design: parallel/ tapered / one piece / two piece
- Implant length:
- Implant diameter:
- Date of delivery of implant (year):
- Implant – Abutment connection type:
- Implant placed by: GP / Specialist
8.3. Prosthesis:

- Prosthesis material: MC / all ceramic
- Prosthesis extension type: mesial / distal
- Crown units/implant ratio:
- Date of delivery:
- Prosthesis delivered by: GP / specialist
- Length of cantilever (mm):
- FDP Supported by:
- No of units supporting FDP:
9. Bibliography:


3. Pjetursson BE, Thoma D, Jung R., Zwahlen M, Zembic A. A systematic review of the survival and complication rates of implant-supported fixed dental prostheses (FDPs) after a mean observation period of at least 5 years, Clin. Oral Implants Res 2012; 23(Suppl. 6), 22–38.


43. Romeo E., Storelli S. Systematic review of the survival rate and the biological, technical and esthetic complications of fixed dental prostheses with cantilevers on implants report edin longitudinal studies with a mean of 5 years follow-up. *Clin. Oral Implants Res* 2012; 23(Suppl. 6) 39–49


52. Yi-Bing Wang. Implant occlusion: biomechanical considerations for implant-supported prostheses, Department of Dentistry, Tri-Service General Hospital 2008; No. 325, J Dent Sci · Vol 3 · No 2


10. APPENDIX

10.1 Ethics approval letter from DHA

<table>
<thead>
<tr>
<th>From:</th>
<th>Dubai Scientific Research Ethics Committee (DSREC) Dubai Health Authority</th>
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<tr>
<td>To:</td>
<td>Ms. Hayat TareeshAlaleed, Hamdan Bin Mohammed College of Dental Medicine</td>
</tr>
<tr>
<td>Study Site</td>
<td>Dental Department in Dubai Health Authority</td>
</tr>
<tr>
<td>Date:</td>
<td>01 May 2016</td>
</tr>
<tr>
<td>Ref:</td>
<td>DSREC-SR-05/2016_02</td>
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</table>

Subject: Approval for the research proposal, "The impact of cantilevers position on clinical outcome of implant-supported fixed dental prosthesis."

Dear Ms. Hayat TareeshAlaleed,

Thank you for submitting the above mentioned research proposal to Dubai Scientific Research Ethics Committee, DHA. Dubai Scientific Research Ethics Committee has been organized and operates in accordance with the ICH/GCP guidelines.

Your request was discussed with Dubai Scientific Research Ethics Committee. We are pleased to advice you that the committee has granted ethical approval for the above mentioned study to be conducted in Dubai Health Authority. However, you will have to approach the Medical Director of the Hospitals to secure permission to review any hospital records and to carry out your study in the hospital.

Please note that it is DSREC’s policy that the principal investigator should report to the committee of the following:

1. Anything which might warrant review of ethical approval of the project in the specified format, including:
   - any serious or unexpected adverse events and
   - unforeseen events that might affect continued ethical acceptability of the project
2. Any proposed changes to the research protocol or to the conduct of research
3. Any new information that may affect adversely the safety of the subjects
4. If the project is discontinued before the expected date of completion (reason to be specified)
5. Annual report to DSREC about the progress of the study
6. A final report of the finding on completion of the study

Please note that this approval is valid for one year from the date of this letter. It is your responsibility to ensure that an application for continuing review approval has been submitted at the required time.