Detectability of Osseous Lesions with a Pre-Programmed Low-Dose Protocol for Cone-Beam Computed Tomography

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Abstract: The present study aimed to compare the diagnostic reliability of a pre-set, manufacturer-specific, low-dose mode against a standard-dose mode in the detection of four different osseous lesions in the mandible with cone-beam computed tomography (CBCT). Four types of lesions (periapical lesion, extended periodontal gap, recession of the buccal lamella, sequestrum/fracture) were prepared on 40 pig mandibles. CBCT images were obtained from each mandible, with both the low-dose and standard-dose modes using the Orthophos SL CBCT device (Dentsply-Sirona, Bensheim, Germany). Twelve assessors quantitatively (detection of lesions) and qualitatively (assessment of detectability) evaluated the CBCT images in SIDEXIS 4 (Dentsply-Sirona) using a study-specific digital examination tool. A correct diagnosis was achieved in almost 71% (LD: 70.8%; SD: 70.9%) of 1920 lesions, without a statistically significant difference between the low-dose and standard-dose mode. This finding was consistent across all four lesion types. In conclusion, while low-dose mode and standard-dose mode CBCT scans performed similarly in the detection of four prepared lesions of the mandible, the former may be a promising, user-friendly alternative method of obtaining radiation-optimized, three-dimensional images in accordance with the As Low As Diagnostically Acceptable (ALADA) principle.

Keywords: cone-beam computed tomography; radiation; low-dose protocols; dentomaxillofacial radiology; ALADA

1. Introduction

Low-dose protocols for cone-beam computed tomography (CBCT) have become a promising instrument for dose optimization in three-dimensional (3D) radiological diagnostics of the dentomaxillofacial region [1,2]. Since its introduction in 1998 [3], CBCT imaging has evolved into a popular radiographic modality in all fields of modern dental medicine [4]. CBCT has demonstrated advantages in the radiological diagnosis of osseous lesions—especially in the bucco-oral dimension. Dehiscence or fenestration defects and defect morphology can be detected significantly better with 3D imaging than with conventional radiographs [5,6]. CBCT also provides relatively more surgically relevant information than conventional two-dimensional (2D) imaging when applied to oral and maxillofacial traumatology and the assessment of bone pathologies such as osteomyelitis [7]. While superimposition-free imaging of the hard tissue of the facial skull induces less radiation relative to multi-sliced computed tomography [8,9], the applied radiation dose is still higher than that emitted by conventional 2D imaging modalities (panoramic or cephalometric radiography) [10,11].
Hence, whether the higher radiation risk justifies the diagnostic benefits of CBCT must be considered prior to its administration [12]. Further, the diagnostic reliability of radiation-optimized CBCT methods, such as low-dose protocols, warrants analysis [13].

Significant dose optimization can be achieved by limiting the field of view (FOV) to the region of interest (ROI) and by modulating tube voltage (kV) and tube current (mA) [1]. However, the modulation of the scanning parameters (e.g., kV, mA) always depends on the CBCT scanner in use [12]. In 2013, approximately 50 CBCT devices from 20 companies were available [14], confounding their adjustment to a standardized (dose-optimized) scanning protocol for a specific diagnostic task [15]. Furthermore, the modification of tube voltage (kV) and tube current (mA) and the reduction in the exposure time result in altered image contrast and image noise [12]. Nevertheless, low-dose CBCT protocols are reportedly sufficient for performing different diagnostic tasks in 3D dentomaxillofacial radiologic practice, and could constitute an important part of the dose optimization process in terms of the As Low As Diagnostically Acceptable (ALADA) principle [2,15–17].

Predefined device-specific low-dose settings can represent a simple, user-friendly possibility for predictable image quality—especially for radiographic examination, where a greater FOV is needed. The aim of this study was to evaluate the diagnostic accuracy of such a predefined device-specific, low-dose protocol compared to the standard-dose protocol. Due to ethical and safety concerns, however, pursuing these avenues of exploration cannot be performed in vivo [13]. Because the human and pig jaws feature related macroscopic anatomy and tissue composition, the pig model has been proven as a suitable and popular in vitro model in dentistry [18,19]. Hence, the present study induced four osseous lesions in pig mandibles to perform the aforementioned investigation.

2. Materials and Methods

2.1. Preparation of Bone Defects

The present study used 40 mandibles of pig cadavers, bought from a local butcher shop. A declaration of non-responsibility from the cantonal veterinary office was obtained from the Department of Animal Welfare and 3R at the University of Zurich.

After partial soft-tissue removal, four types of osseous bone defects were prepared using drilling instruments (rose head bur H141, diamant bur 859, bur (“Lindemann”) H162, Komet Dental, DENTAL Brasseler GmbH, Lemgo, Germany; handpiece Kavo Expertmatic E10C, KaVo Dental AG, Kloten, Switzerland): periapical lesion (PL), extended periodontal gap (EP), recession of the buccal layer (BL), and sequestrum/fracture (SF). All lesions within each lesion type were prepared with the same type of drilling instrument to ensure lesions were as identical as possible (PL/BL, rose head bur H141; EP, diamant bur 859; SF, “Lindemann” bur H162).

The two-rooted first premolars were separated and extracted from all pig jaws prior to the preparation of the PL. Subsequent to the drilling of the artificial apical bone defect, the roots were repositioned into the alveoles. To simulate a BL, the buccal bone layer around the cervical part of the tooth was removed. EP were dissected by drilling along the upper third of the tooth root. Cortical bone defects were created in the jaw angle to simulate SF.

Each lesion type was prepared 20 times within the 40 mandibles in a randomized order, resulting in up to four different types of lesions per mandible (Figure 1).

2.2. CBCT Examination

The Orthophos SL 3D (Dentsply-Sirona, Bensheim, Germany) was used for the radiographic examination. All mandibles were placed and aligned centrally on a created platform by using the positioning lights of the scanner. Every mandible was scanned with the low-dose protocol (85 kV, 13 mA, exposure time 2.2 s; pixel size 0.160 mm) and the standard-dose protocol (85 kV, 13 mA, exposure time 4.4 s; pixel size 0.160 mm) at a selected FOV of 11 × 10 cm (Table 1).
To mimic in vivo conditions, soft tissue was simulated with a cold-pack (12 × 29 cm, GELLO Geltechnik GmbH, Ahaus, Germany) placed in the middle of the jaw.

The 80 CBCT scans (40, low-dose; 40, standard dose) were uploaded to a modified version of Sidexis 4 - Software (Dentsply Sirona, York, PA, USA) and randomly grouped into 16 datasets, each of which contained five CBCT scans.

![Figure 1. Screenshots of the four lesion types imaged with low-dose (upper row) and standard-dose mode (lower row) of cone-beam computed tomography (CBCT).](image)

| Mode | FOV (cm) | kV/mA | Radiation Time (s) | Voxel Size (µm) | Effective Dose (µSv) |
|------|----------|-------|--------------------|-----------------|---------------------|
| LD   | 11 × 10  | 85/13 | 2.2                | 160             | 20                  |
| SD   | 11 × 10  | 85/13 | 4.4                | 160             | 145                 |

Abbreviations: FOV, field of view.

### 2.3. Image Evaluation

All images were assessed by 12 examiners (eight oral surgeons, four maxillofacial surgeons) with at least two years of training in oral radiology and diagnosing CBCT images. All evaluators were calibrated by receiving instructions (by the primary investigator QD) during the examination of five test mandibles. Assessment was performed in three steps: the marking of a lesion with a cursor, the indication of the lesion type, and the qualitative visibility grading of the lesion: from 1, very low to 10, very high.

All scans were assessed in Sidexis 4 using the same workstation (Supermicro, Windows 10 Professional Edition 64 Bit; Intel®Core™ i7-6700 CPU © 3.40GHz Intel64 Family 6 Model 94 Stepping 3, 3400 MHz/x64) and display (HP Z23n 58, 4 cm, 23 Zoll, IPS LED Backlight).

Each mark was registered in a digital coordinate system within Sidexis 4, and the coordinates were saved in an Excel sheet (Microsoft Excel 2016, Microsoft Corporation, Redmond, WA, USA) along with the lesion type and the qualitative visibility grading information.
2.4. Statistical Analysis

Mixed-effects logistic regression was fitted to the dataset in order to estimate the differences in the detection rate between the low-dose and standard-dose protocols with respect to the four osseous lesions. The target variable (correct detection) was modelled by the fixed variables (radiation dose and lesion type) using random intercepts (pig jaw and examiner). Model diagnosis with residuals showed no severe violations of model assumptions. Thus, marginal means were computed with respect to the fixed effects, and post hoc pairwise comparisons were performed and adjusted for multiple testing according to Tukey’s method. The level of significance was set to $\alpha = 0.05$.

All statistical analyses and plots were performed using the statistical software R [21], including the packages lmerTest [22], emmeans [23], and DHARMa [24].

3. Results

The detectability rates of the four lesion types differed independently of the radiation dose (Figure 2). Nearly 71% of lesions were correctly detected (LD = 70.8%; SD = 70.9%).

The PL was detected correctly by the examiners in 62% of the cases (LD = 62.7%; SD = 61.3%); EP, 60% (LD = 60.4%; SD = 59.4%); BL, 79% (LD = 79.2%; SD = 78.8%); and SF, 83% (LD = 80.8%; SD = 84.2%). These results constitute descriptive values derived from a total of 480 observations per lesion type.

As calculated with the estimated marginal means method, predictive values for each of the four lesion types did not differ significantly between lesion detectability with the low-dose or standard-dose protocols ($p = 0.9393$). Differences in predictive values between the detection of the BL, SF, PL, and EP differed significantly ($p \leq 0.0001$; Table 2).
The dose-modes also differed little in terms of visibility (Figure 3): PL featured average low-dose and standard-dose visibilities of 6.9 and 7.0, respectively; EP, 7.4 and 7.5; BL, 8.5 and 8.8; and SF, 9.1 for both.

Table 2. Comparison of the detection rates of the different lesions.

| Lesions   | Odds Ratio | p-Values  |
|-----------|------------|-----------|
| PL/BL *   | 0.4026341  | 0.04344587 | <0.0001 |
| PL/EP     | 1.1017147  | 0.10824976 | 0.7575  |
| PL/SF *   | 0.3153958  | 0.03543323 | <0.0001 |
| BL/EP *   | 2.7362678  | 0.29447974 | <0.0001 |
| BL/SF     | 0.7833310  | 0.09380023 | 0.1736  |
| EP/SF *   | 0.2862772  | 0.03208903 | <0.0001 |

PL and EP were detected significantly less than BL and SF (* statistically significant). Abbreviations: PL, periapical lesion; BL, recession of the buccal layer; EP, extended periodontal gap; and SF, sequester/fracture.

Figure 3. Assessment of the visibility of the four lesion types according to dose-mode (1 = very poor, 10 = perfect). Abbreviations: PL, periapical lesion; BL, recession of the buccal layer; EP, extended periodontal gap; SF, sequestrum/fracture; LD, low dose; and SD, standard dose.

4. Discussion

The increasing use of CBCT in dentistry underscores the need to minimize the patients’ exposure to radiation in accordance with the ALADA principle [12]. While low-dose CBCT protocols allow for minimal exposure, their diagnostic reliability relative to that of standard protocols must be determined in order to define the indication areas [13]. To investigate the diagnostic reliability of a pre-programmed, low-dose protocol with respect to typical osseous lesions, as well as fractures and sequestra, we employed an in vitro model widespread in dental medicine; the porcine jaw is well-suited to the investigation of dento-maxillofacial questions due to its resemblance to the macroscopic anatomy and composition of the human jaw [18,19].

A total of 1920 lesions were evaluated in this study. On average, almost 71% of all four lesions were correctly detected with a negligible difference between the low-dose and standard-dose modes (LD = 70.8% (1359); SD = 70.9% (1361)). Great importance was given to a cautious preparation.
technique when generating the lesions—e.g., although only 20 jaws were used to generate the PL simulations, all first premolars were extracted. By doing so, it became possible to exclude extraction signs as a possible confounder influencing the evaluation. Missing buccal lamellae were prepared on multi-rooted teeth, and fractures/sequestra were prepared in the jaw angle in the absence of alveolar foramina, which could influence the evaluation. The aim was to create a discreet finding to detect a possible difference between the applied radiation doses in small lesions. This may be an explanation for the overall detection rate of only 71%. However, these results compare favorably to those of Hedesiu et al., who observed detection rates of 67.5–74.5% when examining prepared osseous lesions in pig mandibles using three different CBCT devices at a standard dose range [25].

The main objective of the present study was to ascertain possible differences in detection rates between low-dose and standard-dose protocols. This investigation failed to observe any statistically significant differences between low-dose and standard-dose CBCT protocols in the detectability of four different osseous lesions. We further found PL and EP to be correctly detected significantly less often than SF and BL; the former pair may have been rendered less visible also on account of the cautious preparation method and macroscopic anatomical differences between the pig and human jaws. Consistent with this finding, the evaluation for the visibility of the lesion types demonstrated that PL and EP were less visible than the BL and SF in both the low-dose and standard-dose modes, with little difference between the two.

The low-dose method employed in our investigation achieved an effective radiation dose reduction of 85% (LD = 20 mSv; SD = 145 mSv [20,26]) by halving the radiation time (LD = 2.2 s; SD = 4.4 s) and using a larger copper diaphragm (LD = 1 mm copper diaphragm; SD = 0.3 mm copper diaphragm). Three-dimensional information can thus be obtained by means of a conventional imaging modality like the panoramic radiograph [27].

Despite the benefit of these changes, limiting the FOV to the ROI remains a significant factor in reducing the effective radiation dose [1,9]. Pauwels et al. showed that a dose reduction of approximately 82% is possible with the selection of the smallest FOV (4 × 4 cm) rather than the largest (17 × 12 cm) [28]. The vertical limitation of the FOV in particular helps to protect radiation-sensitive organs such as the thyroid gland or the eye lenses and thus contributes to a reduction in the effective dose [29].

As a secondary approach for dose optimization, the modulation of tube voltage (kV) and tube current (mA) can reduce the dose used by standard protocols by up to 40% [1].

PL could be diagnosed much more frequently with CBCT than with conventional imaging methods such as panoramic and periapical radiography [30,31]. 3D imaging could be particularly relevant in cases where there is a discrepancy between the diagnostic findings of the intraoral X-ray and the clinical situation. In such cases, possible clinical causes like, e.g., untreated root canals (MB2) or root resorptions, are visualized [32]. As the FOV does not seem to have a significant impact on detectability with CBCT, it should be chosen to be as small as possible [25]. Indeed, our results indicate that a sufficient dose reduction can be achieved with the application of the low-dose protocol without any significant detriment to the detectability of periodontal lesions.

A recent review has shown that CBCT can visualize periodontal lesions with furcation involvement and allows for the adequate examination of periodontal defects in the bucco-oral dimension [6,33]. However, the application of CBCT in the diagnosis of marginal bone resorption is still a second-level imaging method, following primary 2D radiological evaluation [33]; with the application of the low-dose protocol, the clinical examination can benefit from 3D information without risking significant radiation exposure. In support of this argument, Al-Okshi et al. recently showed that dose-optimized CBCT is sufficient for the assessment of periodontal structures [15].

For the assessment of anatomical landmarks such as the buccal lamella and the maxillary sinus or the diagnosis of fractures, 3D information can be crucial for clinical decision-making. Low-dose CBCT has been shown to be sufficient for adequately diagnosing midfacial fractures as well as for pre-surgical implant planning [34–36]. Low-dose protocols also have potential application in the
radiological follow-up of bone pathologies (e.g., fibro-osseous lesions), as 3D data may reveal changes earlier than 2D information collected with conventional radiography.

In addition to the aforementioned limitation of having not employed a clinical study design, the present investigation may have also been limited by our means of evaluating the CBCT datasets; as the evaluators received the same training in recognizing the four lesion types, a learning effect may have influenced our findings, leading to better diagnostics in later performed CBCT analyses. However, as the evaluators in the present study had at least two years of experience with CBCT images at the time of evaluation, this learning effect during the study should have been negligible. Concerning the expected increased use of low-dose CBCT in the future, it should be noted that the reporting of 3D datasets is more complex compared to 2D datasets, and thus requires specific training [37].

5. Conclusions

The present study demonstrated the similarity between the diagnostic reliabilities of a specific pre-programmed low-dose protocol and a standard-protocol for CBCT. However, our results cannot be applied to all low-dose protocols because of device-specific differences. As suggested in a recent narrative review, more devices need to be tested for the reliability of their low-dose protocols [13].

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