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Investigation into the accuracy and measurement methods of sequential 3D dental scan alignment

Investigation into the accuracy and measurement methods of sequential 3D dental scan alignment

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Abstract

Objectives: Alignment procedures have yet to be standardised and may influence the measurement outcome. This investigation assessed the accuracy of commonly used alignment techniques and their impact on measurement metrics.

Methods: Datasets of 10 natural molar teeth were created with a structured-light model-scanner (Rexcan DS2, Europac 3D, Crewe). A 300µm depth layer was then digitally removed from the occlusal surface creating a defect of known size. The datasets were duplicated, randomly repositioned and re-alignment attempted using a "best-fit" alignment, landmark-based alignment or reference alignment in Geomagic Control (3D Systems, Darmstadt, Germany). The re-alignment accuracy was mathematically assessed using the mean angular and translation differences between the original alignment and the re-aligned datasets. The effect of the re-alignment on conventional measurement metrics was calculated by analysing differences between the known defect size and defect size after re-alignment. Data were analysed in SPSS v24(ANOVA, post hoc Games Howell test, p<0.05).

Results: The mean translation error (SD) was $139\mu m$ (42) using landmark alignment, $130\mu m$ (26) for best-fit and $22\mu m$ (9) for reference alignment (p<0.001). The mean angular error (SD) between the datasets was 2.52 (1.18) degrees for landmark alignment, 0.56 (0.38) degrees for best-fit alignment and 0.26 (0.12) degrees for reference alignment (p<0.001). Using a reference alignment statistically reduced the mean profilometric change, volume change and percentage of surface change errors (p<0.001).

Significance: Reference alignment produced significantly lower alignment errors and truer measurements. Best-fit and landmark-based alignment algorithms significantly underestimated the size of the defect. Challenges remain in identifying reference surfaces in a robust, clinically relevant method.

1. Introduction

Digital 3D scanning, superimposition and comparison, has been used to quantify changes in orthodontics [1,2], periodontics [3,4] and tooth wear [5] measurements, with varying degrees of accuracy. When assessing the accuracy of the 3D comparison process, the majority of literature has focused on the reproducibility error of obtaining the 3D datasets either indirectly via sequential dental models [5–8] or directly using digital scanners. However, the superimposition or alignment of the two datasets is not trivial and is also prone to error [9]. The mathematical complexities of dataset alignment are often hidden from the operator to make software easier to use and may not be immediately obvious to the operator. Errors introduced at this crucial first stage in the digital workflow have rarely been acknowledged in the dental literature [1,10,11] and the complexities in these alignments are under-explored. Comparison is made difficult by the lack of standardisation of measurement metrics. The maximum profilometric change [12], mean profilometric loss [7,13], volume change [6,7]

and percentage of surface change [8] have all been used as outcome measures. No single metric has been universally decided upon and it may be that errors in alignment affect measurement metrics differently.

Traditionally, three different types of scan alignment have been used landmark-based alignment, best-fit alignment and then a reference best-fit. A "landmark based alignment" is performed by the operator manually selecting common landmarks or common points on each dataset which are then aligned by the software. Landmark alignment is relatively straightforward and widely used in medical applications where precision at micron level is not required. However, this method is highly subjective and dependent on the skill and comprehension of the alignment by the operator. In situations when the initial alignment guess is poor or a manual error in a landmark is made, the alignment process will only be partially complete [1].

A standard "best-fit alignment" uses an iterative closest point (ICP) algorithm to align scans, with each software using a slightly different algorithm and do not involve operator-based decisions. The alignment is performed by minimising the mesh distance error between each corresponding data point. By the very nature of the iterative algorithm's termination criteria, alignment will minimise mesh distance error and spread errors evenly over positive and negative deviations. If there is a large defect, the algorithm will attempt to minimise the absolute distance between the two datasets, regardless of the clinical outcome. This may explain erroneous results in tooth wear progression analysis where the tooth appears to have grown over the measurement period [6].

To circumvent this error, researchers have attempted to align on surface areas which have experienced change below a predefined threshold [8] and a recent systematic review on wear measurements has recognised this as a superior approach [14]. A "reference best-fit alignment" aligns datasets by restricting alignment to operator-identified sections of the dataset which are least likely to have undergone change [6,15,16]. This avoids the error of minimising the defect of interest to be measured but introduces an operator error when selecting sections of the dataset.

Quantification of the error of each alignment method has yet to be performed. We are also unaware of what impact an alignment error will have on the measurement outcome. The aim of this investigation was to assess the accuracy of each alignment technique and the subsequent influence on different measurement metrics. The following null hypotheses were

formulated: There is no difference in the position or angulation of the dataset following alignment between the different techniques. Secondly, there will be no difference in the maximum profilometric gain, maximum profilometric loss, mean profilometric gain, mean profilometric loss, percentage of surface change and volume change measurements of a known defect between the different alignment techniques.

2. Method

Ten randomly chosen lower molar teeth were scanned in a dental model scanner (Rexcan DS2, Europac 3D, Crewe) with a stated accuracy of <10 μ m. Duplicates of the datasets were created and an arbitrary defect was created by digitally removing a 300 μ m layer from the occlusal surface using Meshlab [17], leaving a 1 mm intact perimeter. To repair the gaps created in the dataset during this process, a Meshlab Poisson Reconstruction Filter (subdivision level=11) was used. The "eroded step" can be seen in the grey data set in Figure 1. This produced two scans in perfect alignment, with the latter exhibiting a known defect which was quantified in Geomagic Control (version 2.0) with the following metrics: the maximum profilometric gain (μ m) and maximum profilometric loss (μ m) defined as the maximum difference (positive and negative) in the Z axis within the analysed area; the mean profilometric gain (μ m) and mean profilometric loss (μ m) defined as the average difference (positive and negative) in the Z axis within the analysed area; the percentage of the surface with profilometric loss >50 μ m and the volume change of the defect (mm³).

Datasets with the defect were randomly repositioned from their perfect alignment using custom software written in C++. Each dataset underwent a random rotation about an axis by an amount varying 0 to 360 degrees, and a displacement along X, Y and Z between -10 and +10 mm.

Re-alignment with the original, unaltered dataset was then performed using one of three methods in Geomagic Control. For the landmark based alignment algorithm, ten convenient carefully chosen and corresponding landmarks on each dataset were selected by a single operator. The software then aligned the datasets by superimposing these landmarks. The best-fit algorithm was performed using the entire dataset by aligning 1,000 randomly selected data points, which was then refined with an alignment on 5,000 data points. For the reference alignment, the area of the defect and surrounding occlusal surface we wished to measure were manually selected by the operator and deleted from the dataset (visible in light blue in Figure 1). The best-fit alignment process as described above was performed using this

reduced dataset, which was assessed by the operator as having not experienced change. The transformation matrix was then applied to the complete displaced dataset to realign it with the same orientation.

Transformation matrices for all alignments were used to robustly calculate the deviation from the known perfect alignment. Custom software was written using Singular Value Decomposition in the Point Cloud Library (www.pointclouds.org) to calculate the absolute separation in microns of the geometric centre of the dataset (translation error) and the absolute difference in angulation/rotation in degrees (angular error) between the original dataset and re-aligned dataset. The impact of the alignment errors on measurement outcome was assessed by obtaining the mean profilometric gain, mean profilometric loss and volume change measurements using the realigned dataset and subtracting it from the known true defect size.

Data were analysed in SPSS version 24 and initially tested for normality using histograms, boxplots and Shapiro-Wilk's test. The variance between groups was statistically significant using Levene's test. The data were normally distributed therefore differences were assessed using a one-way ANOVA with post hoc Games Howell test for multiple comparisons.

3. Results

The mean translation error or separation between the original dataset and re-aligned data set was 139 μ m (SD 42) using landmark alignment, 130 μ m (SD 26) for best-fit and 22 μ m (SD 9) for reference best-fit alignment. Reference best-fit had statistically significant reduced translation error compared to both landmark and best-fit alignment (p<0.001).

The landmark alignment resulted in a mean angular error of 2.52 degrees (SD 1.18), 0.56 degrees (SD 0.38), for best-fit alignment and 0.26 degrees (SD 0.12) for reference best-fit alignment. Reference best-fit alignment had reduced angular error compared to landmark alignment (p<0.001) but not best-fit alignment (p=0.094).

All mathematical and measurement errors are shown in Figure 2. Negative results indicate an underestimation of the defect size while positive results indicate an overestimate of the defect size. The mean profilometric gain error was -60.4 μ m (SD 23.2) for landmark alignment, -47.3 μ m (SD 7.0) for standard best fit alignment and 1.2 (SD 5.3) μ m for reference best fit alignment. These differences were statistically significant (p<0.001). The mean profilometric

loss error was 63.4 μ m (SD 33.3) for landmark alignment, 81.2 μ m (SD 14.8) for standard best-fit alignment and 26.4 μ m (SD 14.7) using reference best fit alignment. Statistical differences were observed between reference best fit and standard best fit alignment (p<0.001) and between landmark alignment and standard best fit alignment (p=0.006). The volume change error was -6.2 mm³ (SD 5.3) for landmark alignment, -9.0 mm³ (SD 2.9) for best fit alignment and -1.1 mm³ (SD 0.5) for reference best fit alignment. Reference best fit alignment had statistically significant reduced measurement error compared to both landmark alignments (p=0.028) and reference best fit alignment (p<0.001). There were also statistical differences between landmark alignment and best fit alignment (p=0.009). Maximum profilometric errors were statistically similar for all methods of alignment with large standard deviations within groups.

The best-fit alignment resulted in the greatest underestimation of volume change and mean profilometric loss compared to reference best-fit alignment (p<0.001) and landmark based alignment (p<0.05). Best-fit alignment was the only alignment which resulted in overestimation of mean profilometric gain (p<0.001).

Representative colour maps from two examples of the alignments are shown in Figure 3. The scale ranges from +0.4mm to -0.4mm. The true defect colour maps (first column) show a clear negative deviation occlusally as expected (blue), with unchanged buccal and lingual tooth tissue (green). A slight peripheral swelling (yellow/red) can be seen on the circumference of the occlusal table due to the Poisson surfacing algorithm closing the mesh defect caused during the creation process. The second column shows the colour maps after landmark alignment displaying tilts in the data set. The third column displays colour maps after a best-fit alignment. The defect appears to have been pulled occlusally decreasing the size of the defect. A profilometric gain (gain in tooth structure) in yellow on the buccal and lingual surfaces is also visible. The fourth column shows the colour maps after reference best-fit alignment. The appearance is very similar to the true defect, with the full degree of loss recorded.

4. Discussion

Perfect re-alignment will be difficult to obtain with digital comparison software. However, the technique is developing, and many researchers are quoting outcomes for measuring change. The technique using reference best-fit alignment significantly improved the alignment accuracy and decreased the measurement error. In contrast, landmark-based

alignment and standard best-fit alignment resulted in statistically significant increased alignment errors. This resulted in significant underestimation of the defect size and errors consistent with profilometric gain over the data set.

Landmark alignment had the largest angular error which also resulted in the largest mean profilometric gain error. The large standard deviations reflect the difficulty of manually selecting convenient landmarks accurately at a micron level. This method creates greater inconsistencies in the data and poor inter-examiner reliability. In contrast, the best-fit alignment resulted in better angulation of the datasets but the greatest underestimation of the defect as the software minimised the difference between the two datasets. There was also a substantial amount of surface "gain" as the errors were spread evenly across the data set. This may explain why clinical investigations using this method of alignment have failed to show significant differences in the tooth wear progression over time between groups of different risk levels [8,13] compared to those who have used reference best-fit alignment [6].

Reference best-fit alignment significantly reduced the error for each of the measurement metrics. The absolute mathematical translation errors were 6 times smaller in the reference alignment group, while the angular (rotational) errors were half those of the full alignment group. The profilometric gain, profilometric loss and volume change error was also significantly reduced and not statistically different from the true defect. The reference surfaces used for this study represented a small section of the dataset (blue area in Figure 2). Intuitively, one might think that a reduced dataset would adversely influence the alignment algorithm. However, as only one true fit exists for the minimal dataset, the alignment process was unaffected. Challenges remain in identifying these surfaces in an objective, robust method while minimising operator error. It may be that a best fit alignment could be used to identify areas of change before a more accurate selective surface alignment could be performed. A suggested approach may be to initially manually select reference areas which are unlikely to have experienced change based upon clinical knowledge. Alignment could then be restricted to sections on the dataset which have not experienced beyond a predefined threshold. A measurement process error of 15 microns have been observed by our group [5] and others [15,16] and thus a threshold of 20-25 microns as identified by a recent systematic review would seem reasonable [14]. Markers such as gingival margins, adjacent teeth movement and soft tissues are susceptible to change outside of this threshold and thus cannot be assumed to be reliable references.

Maximum positive and negative change measurements led to large standard deviations with no statistically significant differences between alignment methods and a lack of clinically relevant information. As the metric is reliant upon a single data point, any outliers present will corrupt the data. For this reason, maximum profilometric changes should be used with caution unless complete trueness of the dataset can be assumed, which is rare in clinically collected data. The mean positive and negative profilometric changes are well-accepted measures and yet resulted in large underestimations or overestimations of change depending the type of alignment used. When an average value for the entire dataset is taken, it may not always reflect the change being clearly visible in the colour maps. In contrast, volume change measurements are not reduced or averaged when substantial sections of the dataset have not experienced change. This may explain why authors using volumetric analysis to investigate wear progression observed differences in groups [6,7], while those relying on mean profile differences did not [13,18]. If the size of the surface area to be analysed is standardised (for example a 4x4mm section of the dataset) then this may facilitate more accurate comparison of profilometric change measurements.

Previous work by our group [13] and others [12,19] have used best-fit alignments to measure changes in tooth tissue over time or assess the reproducibility of dental materials and tools. This data would suggest that we may be underestimating changes which have occurred. This study is limited in that we have only assessed the performance of Geomagic. Other software packages may result in a more accurate alignment and this will be a focus of future work. The findings from this paper highlight the importance of understanding the datasets which are being compared and choosing the most suitable measurement metric to ensure accurate, clinically relevant conclusions are reached.

5. Conclusions

Reference best-fit alignment resulted in significantly lower alignment errors and truer measurements. Both standard best-fit and landmark-based alignment algorithms significantly underestimated the size of the defect. Aligning data using a best-fit algorithm on selected surfaces which are unaffected by change can significantly improve the measurement accuracy. However challenges remain in identifying these surfaces with a clinically relevant, robust method and require validation with clinical, longitudinal data.

6. Funding

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Figures

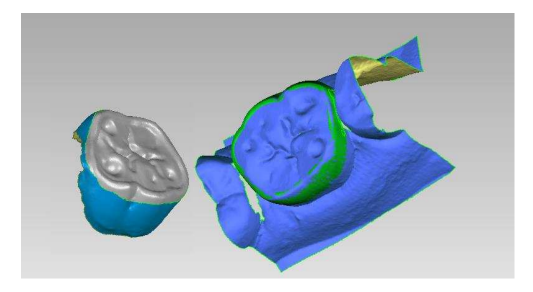


Figure 1: Image of the four surfaces used for analysis. The original surface (purple) is overlaid by the digitally eroded surface in green in perfect alignment allowing true quantification of the defect. The same surface was then duplicated and displaced to test re-alignment. For reference best-fit alignment, the area used for alignment is shown in blue. The area excluded from alignment in grey.

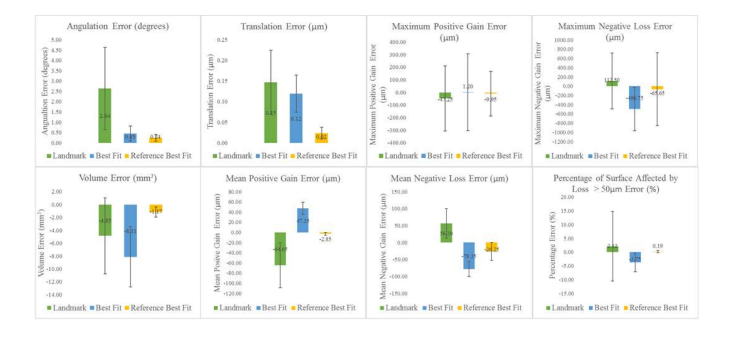


Figure 2: Figure showing errors introduced after the alignment. The angular error and translation errors (mathematical differences between the true alignment and re-alignment) are shown in the top right corner. The remaining graphs represent the resultant measurement errors (differences from known true value of the defect).

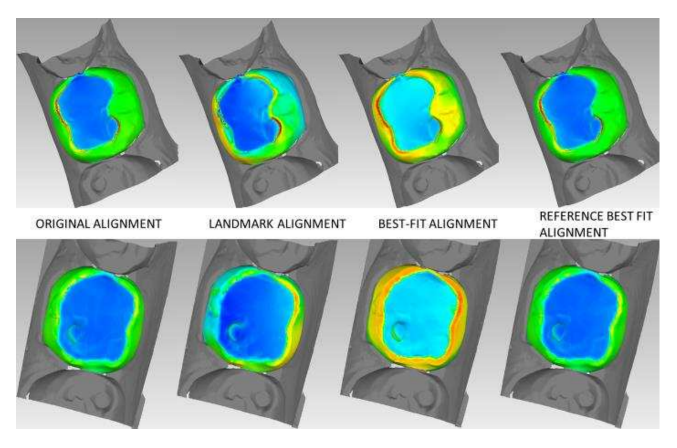


Figure 3: Representative images showing defects after alignment. Areas in green depict no change, deepening areas of blue indicate deepening areas of loss and deepening areas of red indicate areas of gain.

Figure Captions

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