3D superimposition and understanding temporomandibular joint arthritis

Structured Abstract

Objectives – To investigate the 3D morphological variations in 169 temporomandibular joint (TMJ) condyles, using novel imaging statistical modeling approaches.

Setting and sample population – The Department of Orthodontics and Pediatric Dentistry at the University of Michigan. Cone beam CT scans were acquired from 69 subjects with long-term TMJ osteoarthritis (OA, mean age 39.1 ± 15.7 years), 15 subjects at initial consult diagnosis of OA (mean age 44.9 ± 14.8 years), and seven healthy controls (mean age 43 ± 12.4 years).

Materials and methods – 3D surface models of the condyles were constructed, and homologous correspondent points on each model were established. The statistical framework included Direction–Projection–Permutation (DiProPerm) for testing statistical significance of the differences between healthy controls and the OA groups determined by clinical and radiographic diagnoses.

Results – Condylar morphology in OA and healthy subjects varied widely with categorization from mild to severe bone degeneration or overgrowth. DiProPerm statistics supported a significant difference between the healthy control group and the initial diagnosis of OA group ($t=6.6$, empirical $p$-value = 0.006) and between healthy and long-term diagnosis of OA group ($t=7.2$, empirical $p$-value = 0). Compared with healthy controls, the average condyle in OA subjects was significantly smaller in all dimensions, except its anterior surface, even in subjects with initial diagnosis of OA.
Conclusion – This new statistical modeling of condylar morphology allows the development of more targeted classifications of this condition than previously possible.

Key words: bone degeneration; bone overgrowth; temporomandibular joint condyle

Introduction

When the temporomandibular joint (TMJ) presents with initial signs of destruction and inflammation, it is essential to monitor active disease, either localized to the TMJ or systemic, before loading the joints with orthodontic/orthopedic forces or undertaking jaw surgery.

The TMJ differs from other joints because a layer of fibrocartilage, and not hyaline cartilage, covers it (1). The bone of the mandibular condyles is located just beneath the fibrocartilage, making it particularly vulnerable to inflammatory damage and a valuable model for studying arthritic bony changes. The condylar bone is the site of numerous dynamic morphological transformations, which are an integral part of the initiation/progression of arthritis, not merely secondary manifestations to cartilage degradation. Thus, a strong rationale exists for therapeutic approaches that target bone resorption and formation (2–7).

The Research Diagnostic Criteria for temporomandibular disorders validation project (8, 9) concluded that clinical criteria alone, without the use of imaging, are inadequate for valid diagnosis of TMJ arthritis. The application of cone beam CT (CBCT) to craniofacial imaging, with an adequate acquisition protocol, provides a clear visualization of the hard tissues of the TMJ and markedly reduces radiation and cost compared with medical CT (10) (Fig. 1).

Methods for the registration of 3D condylar morphology are essential for the measurement of subtle bony differences in condylar morphology. The regional superimposition techniques, used in this study for across subject comparisons, have been validated by Schilling et al. (11). The objective of this study is to determine the 3D morphological variations in asymptomatic controls, subjects at initial TMJ OA diagnosis and subjects with long-term history of TMJ OA, using 1002 imaging biomarkers. The working hypothesis is that bone morphology is characteristically different in OA compared with controls even at early diagnosis.

Methods

Sixty-nine subjects with long-term TMJ OA (mean age 39.1 ± 15.7 years), 15 subjects at initial consult diagnosis of TMJ OA (mean age 44.9 ± 14.8 years), and seven healthy controls (mean age 43 ± 12.4 years), recruited from the university clinic and through advertisement,
underwent a clinical exam by an orofacial pain specialist. For the initial diagnosis group, only subjects with recent histories of pain, within the last 2 months, were included. For the long-term diagnosis, 13 subjects had only one condyle included as the other side joint had prior history of interventions (joint injections, arthrocentesis, or other pathologies, such as ankylosis) (Fig. 2). Recruitment and clinical exam diagnoses of TMJ osteoarthritis or health were confirmed by CBCT images (8, 9). The imaging protocol consisted of a 20-s scan, using a large field of view to include both TMJs and the same machine (i-Cat® CBCT, 120 kV, 18.66 mA, Imaging Sciences, Hatfield, PA, USA). The university institutional review board approved this study, and all subjects were consented.

**Image analysis methods**

The flow chart in Fig. 3 describes an overview of the image analysis procedures introduced by Schilling et al. (11). All scan volumes were sliced to a voxel size of 0.5 mm³ (Fig. 3B) to standardize voxel size, decrease the computational power and time required to compute the automated registration, using open-source software (3D Slicer v. 4.3.1, [http://www.slicer.org](http://www.slicer.org)) (12).

3D surface mesh models of the right and left mandibular condyles were constructed by semi-automatic discrimination procedures that outlined the cortical boundaries of the condylar region and allowed manual editing, checking slice by slice in all three planes of space (Fig. 3C) using open-source software (ITK-SNAP software v.2.4, [www.itksnap.org](http://www.itksnap.org)) (13). After generating all 3D surface models, left condyles were mirrored in the sagittal plane to form right condyles to facilitate comparisons (Fig. 3D).

Owing to individual morphological variability across subjects, voxel-based approaches fail, and a landmark-based approach was used to approximate consistently all condyles in the same coordinate system (11). The surface mesh files for each condyle were opened in surface analysis software (VAM v. 3.7.6, Canfield 113 Scientific Inc., Fairfield, NJ, USA), and 25 landmarks were placed on each condylar surface model: four points evenly spaced along the superior surface of the sigmoid notch, four on the medial and four on the lateral portions of the ramus adjacent to the sigmoid notch, three along the posterior surface of the neck, three on medial and three on lateral portions of the neck, and one on the medial, one lateral, one anterior, and one posterior extremes of the condylar head (Figs 3E and 4A). The 25 landmarks were used only for spatial approximation of all condyles in a common arbitrary xyz coordinate system (Fig. 4B). That is, the 25 landmarks were used only for registration of all surface models and not for analysis of morphological variability. After registration, all 3D surface models were simulta-

*Fig. 2. CBCT cross sections and 3D models for a patient with ankylosis on the right side and OA on the left side; these are examples of exclusion and inclusion criteria, respectively. Note that red arrows indicate bony proliferations on the condyle and subchondral lesions in the articular fossa on the ankylosed right side. These findings differ from the left side that presents with osteoarthritic changes exemplified by flattening of the lateral pole (yellow arrows) and a bony projection on the anterior condylar surface (orange arrows), without dysmorphology of the articular fossa (green arrows).*
neously clipped/cropped to define the condylar region of interest and open-source software (SPHARM-PDM software, http://www.nitrc.org/projects/spharm-pdm) (14–17) was used to generate a mesh approximation from the volumes, whose points were mapped to a ‘spherical map’ (Fig. 3F). The homology/correspondence of the mapping of the 1002 points across all subjects (Fig. 4C) was verified with color-coded maps of the surface parameterization (Fig. 3G).

Creation of the average mesh

An average 3D condylar shape was generated for the two TMJ OA groups and control group (Figs 3H, 4A and 5A). The core of the ability to compute the group average and group variability is the establishment of correspondence (homology) between each of the 1002 points in the condylar surface models across all subjects. This allows the association of any of the 1002 points’ locations on the condyle of subject A with the

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Fig. 3. Flowchart of the image analysis procedures. (A). Radiographic diagnoses of TMJ health or disease in multiplanar cross sections; (B). standardization of voxel size; (C). construction of surface models; (D). mirroring of left condyles to form right condyles; (E). landmark registration; (F). SPHARM-PDM spherical parameterization of 1002 correspondent mesh point-based models; (G). correspondence verification; (H). construction of average healthy, initial diagnosis, and long-term history of OA surface meshes; (Ia). computation of surface distances between groups; (Ib). computation of vector differences; (Ic). p-value color-coded maps showing the surfaced with statistically significant condylar morphological differences; (J). diagnostic classification of the severity of condylar morphological changes.
corresponding locations on the condyle of subject B, C... n number of subjects. Considerable intersubject variability is also accounted for in the model that captures the average condylar morphology and variability around that average morphology. Open-source software (Linux MeshMath script, http://www.nitrc.org/projects/spharm-pdm) (17) was used to create average meshes for the control, initial diagnosis, and long-term history of OA groups. The 3D morphological variability of the 1002 surface point correspondences was used for surface mesh averaging. The affine transformations of the 3D morphological variability were then applied to the 1002 points on the condylar surface individually. In geometry, an affine transformation is a transformation which preserves ratios of distances between points lying on a surface model, where parallel lines will remain parallel to each other after an affine transformation. Grouping all the mean points provided the linear and non-linear deformation fields that resulted in the average condyle shape for each group.

Calculation of absolute and signed distances and vector differences

The MeshMath script was then used to calculate 3D point-wise subtractions between each group’s average morphology (Fig. 3Ia, Ib). The computed 1002 vector differences were displayed on the condyle surface and scaled according to the magnitude of difference and pointing in the direction of change. The patterns of variation across TMJ OA and control samples were determined through calculation of signed distances, where the areas of bone resorption were displayed as negative values (blue), no differences (0 mm surface distances, white), or bone proliferation as positive (red).

Fig. 4. (A and B) landmark-based registration used to approximate condyles from all subjects in the group comparisons. (A). 25 points in the ramus and condyle surfaces used for the landmark-based registration; (B). reference condylar model (red) with the overlays of multiple condyles approximated in the same coordinate system. Note that the xyz coordinate system is common to all models after registration and uses as reference an arbitrary condyle in the sample to standardize the coordinate system across all condyles; (C). parameterization of 1002 homologous or correspondent surface mesh points for statistical comparisons and detailed phenotypic characterization of the surface models.
Control of the quality of the signed distances

Semitransparent overlays between the average models in 3D Slicer software were used to compare visually the signed distance patterns (14).

Statistical analysis

The statistical framework for testing condylar morphological variations included a Hotelling T-squared test in a multivariate analysis of covariance open-source software (Shape Analysis MANCOVA, Fig. 3Ic) (18,19) corrected for false discovery rate at 0.05. For testing a high-dimensional hypothesis, Direction–Projection–Permutation (DiProPerm) (20), rigorously tested lower dimensional graphical visual differences: direction (projected samples onto an appropriate direction), projection (calculated univariate two sample statistics), and permutation (assessed significance using 1000 permutations of group membership). The approach of distance weighted discrimination (DWD) calculated a direction vector to classify high-dimensional data sets, and their principal components (PC) were graphically plotted. The vector space is a mathematical structure formed by the collection of surface vectors that is called the feature space. Because the control and OA samples had different sample sizes, an appropriately weighted version of DWD, wDWD, was used to find a direction vector in the feature space separating the diagnostic groups.

Results

The semitransparent overlays revealed that compared with healthy controls, the initial TMJ OA diagnosis group showed marked condylar bone changes and that these bone changes were even more marked in the long-term TMJ OA group (Fig. 5A,B). The mean OA models were of smaller size in all dimensions, and areas of statistically significant differences were in the superior articular surface of the condyles, particularly in the anterior and superior portion of the lateral pole. That is, average bone resorption in this area was $-2.7$ mm in the initial TMJ OA diagnosis group and $-4.2$ mm in the long-term TMJ OA group as compared to the healthy control group (Fig. 5C). In the anterior surface of the condyle, on average, a small area of $1.2$ mm of bone apposition was noted in the initial TMJ OA...
diagnosis group and of 3.5 mm in the long-term TMJ OA group compared with the healthy control group (Fig. 5C). When the initial diagnosis and long-term OA average condylar models were compared, statistically significant differences indicative of more bone resorption in the long-term OA group were noted along the whole condylar surface except at the superior surface of the lateral pole. Subgroups were established by comparison of each individual condyle and the healthy controls' average condylar morphology (Fig. 6).

The DiProPerm test found statistically significant morphological differences between the healthy control and the OA groups (Fig. 7A; \( p \)-value = 0.001), between the healthy controls and the initial OA group (Fig. 7B; \( p \)-value = 0.006), between the initial diagnosis and long-term diagnosis of OA groups (Fig. 7C; \( p \)-value = 0.0009), and between the healthy controls and the long-term diagnosis of OA groups (Fig. 7D; \( p \)-value = 0). The projection plots of healthy control condyles (red circles in Fig. 7A) tend to cluster and are clearly separated from the OA groups. Most projected plots of initial OA diagnosis condyles are located within the bounds of the plots of the long-term OA diagnosis condyles (Fig. 7A–C). The maximal partition of condylar morphology, as established by 1002 points in each individual condyle, can be observed in the graphic plots of the principal component refined in the wDWD direction. The wDWD direction onto the PC was shown by the angle in the PC analysis score plots shown in Fig. 7D. Figure 8, in the manuscript online version, shows that while individual morphology variability occurs in the OA patients, the principal component of deformation markedly reveals the flattening of the lateral pole (sequence of images as the .gif file animation plays).

Discussion

These study findings revealed imaging biomarkers of the bone morphology differences at the articular surfaces of the condyle that can now be quantitatively tested by statistical imaging methods. The novel aspect of this study is that, even though the ability to predict progression was not addressable in the cross-sectional study design, these biomarkers can be reasonable surrogate biomarkers of tissue destruction and/or repair overtime.
The methods used in this study included open-source image analysis software. Commercial software packages produce adequate surface reconstructions and/or offer landmark, surface, and/or voxel-based registration methods, but they are not open source, cannot be modified, do not interact well with each other, do not provide flexibility for customization, and, moreover, do not address the 3D correspondence problem across subjects with different facial morphology or from pre- and post-surgery scans of the same patient. Due to its open licensing, 3D Slicer has allowed specific implementation of modules for assessment of condylar changes and their detailed tutorials are now available (e.g., at www.youtube.com/channel/UCQUtGe5KrpBt2k4mrNeHeUQ). 3D Slicer version 4 provides powerful tools for multimodal imaging, volume rendering, registration, and visualization.

Fig. 7. DiProPerm graphic results. The left panels show the distribution of the data projected onto the DWD direction, illustrating how well the groups listed in the right panels can be separated. The curves in the left panel are smooth histograms, with each color showing the subhistograms for that group. The center panel shows principal component graphics, where each condyle is plotted in the first principal direction. The horizontal x-axis is the projected value, and the vertical y-axis reflects order in the data set, to avoid overplotting.

Due to great individual morphological variability across subjects, rigid voxel-based approaches fail to register anatomic structures from different subjects in populational or group average studies. Other voxel-based registration approaches, which are fluid or semifluid (21), lead to changes and deformation of the morphology of the surface models, known as morphing, and the morphing hampers adequate evaluation of individual morphology. For these reasons, the registration across subjects in this study used an initial landmark-based approach to approximate all condyles in a common arbitrary xyz coordinate system (as shown in Fig. 4). This initial procedure only aimed to approximate all condyles and allow the automatic computation of 1002 correspondent surface points for all condyles in a consistent way.
Our findings shown in Fig. 7A reveal that most plot projections of initial OA diagnosis condyles were located within the bounds of the long-term diagnosis condyles. This finding may be explained by the fact that both groups present characteristic osteoarthritic changes. The characteristic osteoarthritic changes observed in this study consisted of flattening of the lateral pole and bony projections in the anterior condylar surface, at initial diagnosis and significantly more marked at long-term diagnosis. It is possible that, even though the initial OA diagnosis group included only subjects with histories of pain within the last 2 months, the condition could have been of long-standing but asymptomatic for the individual. Interestingly, some of the plots of the condyles at initial diagnosis were projected toward the healthy control group. This may be explained by the fact that in patients with diagnosis of OA, the disease progression usually affects one joint first, while morphological changes may be less evident in the contralateral TMJ, particularly at initial diagnosis.

The present study findings represent a preliminary step toward an index of osteoarthritic changes. Our research group’s long-term goals are to implement statistical measurements that allow early detection of the degree of bone destruction and/or bone proliferation in any individual joint (22,23). Such an early diagnosis index is beyond the scope of the present work and will require larger control and OA samples, longitudinal studies and further statistical description of combinatorial biomarker assessments, such as receiver operating characteristic (ROC) curves on disease vs. health, as well as classification-based schemes for computer-aided diagnosis of TMJ OA.

Importantly, an emerging theme in OA is a broadening of focus from a disease of cartilage to a disease of the entire joint and the multiple biological systems that interact with one another in this disease (4). The cross-talk that occurs between the components of the joint, which takes place over years, results in degradation of the articular cartilage and disk, bony changes, synovial proliferation, muscle and tendon weakness, and fatigue. The TMJ condyle is the site of numerous dynamic morphologic transformations in the initiation/progression of OA (24–27), which are not merely manifestations secondary to cartilage degradation. Thus, a strong rationale exists for therapeutic approaches that target bone resorption and formation and take into account the complex cross-talk between all of the joint tissues. While this study focused on bone morphological variability, greater understanding of mechanisms in this multitissue disease requires a combination of imaging technologies, using methods that facilitate superimposition of soft and hard tissue data sets.

Conclusions

This new statistical modeling of condylar morphology allows the development of more targeted statistical classifications of TMJ OA than previously possible. Compared with healthy controls, the average condyle in OA subjects was significantly smaller in all dimensions, except its anterior surface, even in subjects with initial diagnosis of OA.

Clinical relevance

This study investigated morphological variations in 169 temporomandibular joint (TMJ) condyles,
using novel imaging modeling approaches. Direction–Projection–Permutation statistics supported significant differences between the healthy control and TMJ osteoarthritis (OA) groups, even at initial diagnosis, with targeted classification of the severity and location of morphological differences. The TMJ condylar bone is the site of numerous dynamic morphological transformations, which are an integral part of the initiation/progression of OA, not merely secondary manifestations to cartilage degradation.

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Conflict of interests

This is an original work that has not been submitted for publication or presentation elsewhere. All authors had no financial and personal relationships with other people or organizations that could inappropriately influence (bias) their work.

References


