Content-Based 3D Shape Retrieval System

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1 INTRODUCTION

1 Introduction

This is a technical report detailing the implementation of a content-based 3D shape retrieval system. The aim of this implementation is the exploration of existing techniques for shape querying using meaningful 3D descriptors, either scalar or histogram-based. To this effect, the Princeton Shape Database is used to evaluate the system in an environment closer to a real life application than if a smaller database would be used.

1.1 Overview

This introductory section concludes with a short overview of existing related work. The technologies used for the application are then listed and briefly described. The Pre-processing section explains the choices of dataset, granularity and vertex count before detailing the database creation and normalization pipeline. The Feature Extraction section lists and explains the features that are implemented in our system, as well as some reasoning behind the design of the feature vector. The Matching section explains our distance function and how shape matching is actually performed using a kd-tree and k-nearest neighbors. Before evaluation, a t-SNE-powered visualization of the feature space of our database is provided. After this, the database is evaluated using MAP and precision at k and some query examples are given. Finally, the conclusions sum up our system's behavior and explore the limitations within our project, as well as the possibilities for improvement and future work.

1.2 System Specifications

All testing has been performed on a Dell Inspiron 7577 with a Windows 10 Pro 64-bit operating system, 16 GB of RAM, an Intel(R) Core(TM) i7-7700HQ CPU @ 2.80 GHz (8CPUs) and NVIDIA Geforce GTX 1060 with Max-Q Design video card.

1.3 Motivation

This project was created as part of the *Multimedia Retrieval* course held by Prof. Dr. Alexandru Telea at the University of Utrecht.

1.4 Literature review

Vranić & Saupe (2003), Körtgen et al. (2003) and Tangelder & Veltkamp (2007) provide a thorough overview of classical 3D shape retrieval methods using shape descriptors. The features used in this project are also

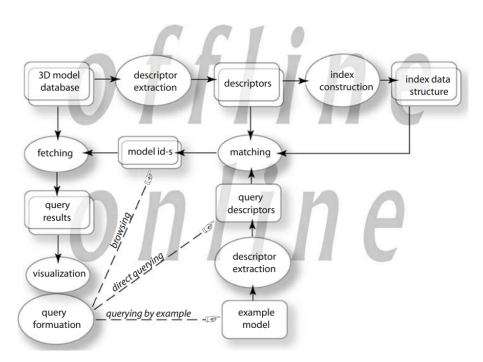


Figure 1: Tangelder & Veltkamp (2007), Conceptual framework for shape retrieval.

explained in their work. Surveying the existing methods, Tangelder & Veltkamp generated a diagram showing the conceptual framework for shape retrieval applications (*Figure 1*). This diagram also describes how our application is structured on a high level, minus any implied online functionality. Additionally, only querying by example is possible.

Feng et al. (2016) use a different approach than the one described in this paper, using voxel shapes and skeleton descriptors for shape matching. Even so, some alternative histogram matching techniques are presented, including Earth Mover's Distance and Multilevel Distance, which could represent viable alternatives to the histogram distance function used in our implementation.

Song (2017) describes some advanced indexing techniques for multimedia data retrieval which we make use of in our project in order to speed up searching. Most notably, he provides some details of kd-tree functionality, as well as some alternative approaches to dimensionality reduction.

2 Technologies Used

This section will list the external libraries used in developing the application. The two programming languages used are C++ for the shape retrieval system and Python for plots and visualization. C++ was selected for its speed and Python for its ease of use.

2.1 C++

ALGLIB v3.15.0 Free Edition ALGLIB is a cross-platform numerical analysis and data processing library. It contains tools for data analysis, optimization and nonlinear solvers, interpolation and linear/non-linear least-squares fitting, linear algebra and many other algorithms. In our implementation, it is used for PCA. Details can be found here: https://www.alglib.net/.

ANN v1.1.2 ANN (A Library For Approximate Nearest Neighbor Searching) is a library written in C++, which supports data structures and algorithms for both exact and approximate nearest neighbor searching in arbitrarily high dimensions. This is used in our implementation to speed up shape matching using the kd-tree structure and k-nearest neighbors. More details can be found here: https://www.cs.umd.edu/~mount/ANN/.

nupengl.core This NuGet package contains various libraries required to build OpenGL applications, including FreeGLUT, GLFW and GLEW. Details can be found here: https://www.nuget.org/packages/nupengl.core/.

PMP Library v1.1.0 The Polygon Mesh Processing (PMP) library contains three main modules used for mesh processing: core (used for data representation), algorithm (used for algorithm implementations) and visualization (used for OpenGL-based visualization). Out of these, the algorithm module is used in our system for mesh decimation. More information, as well as a download link, can be found at: https://www.pmp-library.org/.

3D Unstructured Grid Implementation The application uses a 3D unstructured grid implementation provided by the University of Groningen as part of their Data Visualization Book. Basic visualization using OpenGL as well as a PLY file reader are also included. The implementation can be found here: http://www.cs.rug.nl/svcg/uploads/DataVisualizationBook/sp3_grids.zip

ZPR v0.4 Zoom, Pan and Rotate (ZPR) is a mouse manipulation module for C or C++ GLUT/OpenGL programs. In our project, ZPR enables mouse zooming, panning and rotation when visualizing a 3D shape. The implementation may be found here: http://www.nigels.com/glt/gltzpr/.

2.2 Python

matplotlib v3.1.1 Matplotlib is a Python 2D plotting library. It is a requirement for the Seaborn library which was used for generating the 2D plot of the t-SNE result. Website: https://matplotlib.org/.

numpy Numpy is a fundamental package for scientific computing with Python, as it contains a powerful N-dimensional array object used by other scientific python libraries, such as pandas. Details can be found here: https://numpy.org/.

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pandas v0.25.3 Pandas in an open source library providing high-performance, easy-to-use data structures and data analysis tools for Python. The data frame structure it contains can be used with seaborn to simplify plotting. Website: https://pandas.pydata.org/.

seaborn v0.9.0 Seaborn is a Python data visualization library based on matplotlib. It provides a high-level interface for drawing graphs. Seaborn was used to draw the scatter plot resulting from our t-SNE implementation. Details about the library can be found here: https://seaborn.pydata.org/.

t-SNE Implementation T-Distributed Stochastic Neighbor Embedding (t-SNE) is a technique for dimensionality reduction that is well suited for high-dimensional dataset visualization. Our project uses the implementation by Laurens van der Maaten for the Python language, which can be downloaded here: https://lvdmaaten.github.io/tsne/code/tsne_python.zip. More details about t-SNE can be found on the official web page: https://lvdmaaten.github.io/tsne/.

3 Pre-processing

3.1 Database selection

For the purpose of this project, the Princeton Shape Benchmark (PSB) database of 3D models was selected. Details about this set of 3D models can be found in (Shilane, P. et al, 2004) or on the official website: https://shape.cs.princeton.edu/benchmark/.

This dataset was chosen due to its size, variety, cleanliness and its good options of granularity in terms of classes. A bigger, more diverse database will more accurately reflect the more difficult real-world applications. In addition, the chosen granularity of 53 leaf classes permits for mostly homogeneous shapes within the same class, e.g. the winged vehicle class, while also a flying object, should be very different from the balloon class. Conversely, a finer granularity would mean too many classes that would be too specific, which would interfere with the evaluation step.

3.2 File reading

The files from the PSB were all provided in the OFF file format and an OFF reader was written to accommodate the files in the database. A PLY reader was also utilized in testing and is available within the source code, but the OFF file type is our application's standard.

3.3 Database evaluation

The database contains 1815 models in the OFF file format. All of the models contain only triangles as faces. The distribution of faces is shown in *Figure 2*. The average number of vertices is 4221, the minimum 10, and

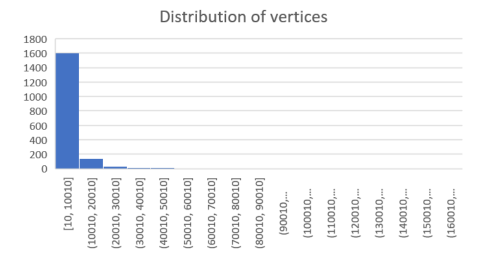


Figure 2: Histogram showing the distribution of vertices in the PSB database

3.4 Supersampling 5

the maximum 160940. The histogram, along with the average, shows that most of the models have a vertex count of smaller than 20,000. In order to successfully perform the upcoming steps of feature extraction the models need to have approximately the same number of vertices. Two actions can be undertaken to achieve this, supersampling and subsampling. The models with a low vertex count are supersampled and the ones with a large vertex count are subsampled. A target number of vertices has to be chosen for this. One solution could be to choose the average. However because several of the meshes have a very high resolution (a hundred thousand vertices), subsampling this to reach the 4000 required vertices would ruin the original shape. Therefore the target vertex count is chosen to be higher than the average, so that the higher resolution models retain their characteristics. The only downside for choosing a higher target number of vertices is an increase in computation time. The target number of vertices is set at 20,000, and the methodology for supersampling and subsampling is explained in the following two sections.

3.4 Supersampling

Supersampling is done in this implementation by splitting a larger triangle into 4 smaller ones via adding the midpoints of the original triangle's edges as vertices and using them to form new triangles with the original vertices. An example is shown below:

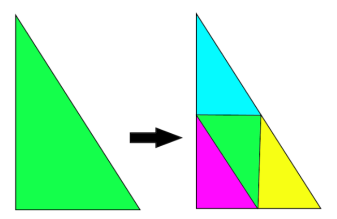


Figure 3: Supersampling Example

To ensure the triangles obtained like this are not outliers in terms of their size, the cells in the model were first sorted by their area inside a vector, from smallest to largest. The area of each cell was calculated using the following formula:

$$S = \frac{|AB||AC|sin(\theta)}{2} \tag{1}$$

Thus, at each step, only the triangle with the largest surface is split up into smaller triangles. The original triangle is removed from the vector, and the smaller triangles are inserted at positions that maintain the ordered property of the data structure.

3.5 Subsampling

For the task of subsampling the Polygon Mesh Processing (PMP) library is used. The SurfaceSimplification class is used to perform mesh decimation, taking as input the mesh and a target number of vertices. To demonstrate that this algorithm works correctly for our models a pre- and post- visualization is shown in figure 4. This is the largest model in the dataset with 316.498 faces and 160.940 vertices. The post decimation model contains 78.473 faces and 40.000 vertices in the middle picture and 20.000 vertices in the right picture. Even though the model only has a fourth of its original vertices in the second case, it clearly retains its characteristics. And even when the vertex count is brought down to 20.000 vertices it is hard to distinguish them. Therefore we can conclude that the algorithm is a suitable tool for this project.

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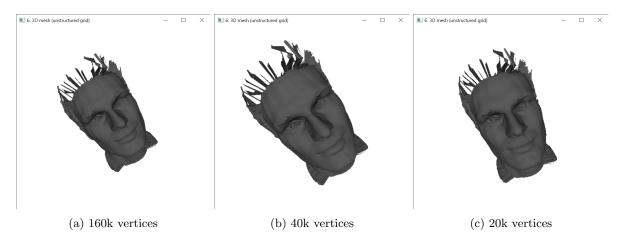


Figure 4: Results of mesh decimation for model m303

3.6 Four step normalization

Next each mesh will go through the four step normalization pipeline so that it is ready to be used in upcoming tasks. This four step normalization is also called pose normalization, which refers to the pose of the models. The goal of this is so that similar models will have the same orientation. Figure 5 shows a visualization of what each step does, the red, green, and blue represent the x, y, and z axes respectively from zero to one. The first step is to center on the barycenter, then, in step 2, PCA is done to orient the object in an intuitive way. Step 3 performs a fliptest which results in the majority of the mass in the object to be located in the negative side of the axis. Finally step 4 normalizes the model, which is excluded in the figure because the model was already normalized and thus would show no difference.

3.6.1 Center on the barycenter

For the first step, the barycenter b is calculated, which is the average x, y, and z coordinates of all vertices in the mesh. Next, each vertex v is translated, by subtracting these averages from each of its points resulting in the new vertex position u.

$$\begin{bmatrix} u_x \\ u_y \\ u_z \end{bmatrix} = \begin{bmatrix} v_x \\ v_y \\ v_z \end{bmatrix} - \begin{bmatrix} b_x \\ b_y \\ b_z \end{bmatrix}$$

The result can be verified by calculating the average coordinates again, which should be 0 after the normalization.

3.6.2 PCA

Principal component analysis (PCA) is done using the ALGLIB library. The three eigenvectors e_1 , e_2 , and e_3 are returned by the algorithm. These three vectors are orthogonal to each other. Therefore these eigenvectors can easily be made to coincide with the x, y, and z axis, because the model only has to be rotated. A rotation matrix M is than created using the vectors x, y, and z, that represent the axes of the normal coordinate system. The eigenvector corresponding to the largest eigenvalue will coincide with the x axis, the next eigenvector with the y axis, and the final one with the z axis.

$$x = \begin{bmatrix} 1 \\ 0 \\ 0 \end{bmatrix} y = \begin{bmatrix} 0 \\ 1 \\ 0 \end{bmatrix} z = \begin{bmatrix} 0 \\ 0 \\ 1 \end{bmatrix} M = \begin{bmatrix} x \cdot e_1 & x \cdot e_2 & x \cdot e_3 \\ y \cdot e_1 & y \cdot e_2 & y \cdot e_3 \\ z \cdot e_1 & z \cdot e_2 & z \cdot e_3 \end{bmatrix}$$

The new vertex u is obtained by multiplying this matrix with the old vertex v.

$$\begin{bmatrix} u_x \\ u_y \\ u_z \end{bmatrix} = M \cdot \begin{bmatrix} v_x \\ v_y \\ v_z \end{bmatrix}$$

The result of the translation can easily be verified. Doing PCA again on the translated mesh returns three eigenvectors that now correspond with the x, y, and z axis.

3.6.3 Fliptest

The eigenvectors used for the translation in the previous step are unoriented and thus give no information about to which side the model should be directed. Using the fliptest it is ensured that the majority of the mass resides in the negative half-space, the important part of this step is that all models are consistently oriented, i.e. two planes should be facing the same way. Mass in this case is not indicated by the number of the vertices but also takes momentum into consideration, i.e. vertices farther away from the origin have a higher weight. Three variables are introduced: w_x , w_y , and w_z that indicate the total weight of each of the three coordinates.

$$w_i = \sum sign(C_{t,i})(C_{t,i})^2 \tag{2}$$

where $C_{t,i}$ is the i^{th} coordinate of triangle t $(i \in x, y, z)$. The latter part of the summation gives coordinates far away from the origin an higher weight, while the former part gives it either a negative or positive weight. These values are than used for a new scaling matrix M, which flips the coordinates if necessary. In case the mesh is already properly oriented, M will simply be the identity matrix.

$$M = \begin{bmatrix} sign(w_x) & 0 & 0\\ 0 & sign(w_y) & 0\\ 0 & 0 & sign(w_z) \end{bmatrix}$$

3.6.4 Normalization

The last step scales the model in the unit volume, i.e. it can fit in a unit cube. First the min and max of the x,y, and z coordinates of the axis-aligned bounding box are found. Next the largest distance δ between the min and max of these coordinates is used for the scaling factor $s = \frac{1}{\delta}$. Finally each vertex v is multiplied with this factor to obtain the new vertex u.

$$\begin{bmatrix} u_x \\ u_y \\ u_z \end{bmatrix} = \begin{bmatrix} v_x \\ v_y \\ v_z \end{bmatrix} \cdot s$$

A visualization is excluded, however the effectiveness of this step can easily be verified by looking at the resulting bounding box of the model.

3.7 Verifying the normalization

Next, for each model in the database the following steps are performed:

- 1. If the model has less than 20,000 vertices use mesh decimation with target number of vertices set to 20,000.
- 2. If the model has more than 20,000 vertices use supersampling with target number of vertices set to 20.000.
- 3. Center the model on its barycenter.
- 4. Perform PCA and align the eigenvectors with the x, y, and z axis.
- 5.Flip the model if necessary.
- 6. Normalize in a unit cube.

In total 19 files were unable to processed, leaving a total of 1796 models in the new dataset. To verify that all the steps were performed correctly a new overview of the database has been created. The minimum number of vertices is 20,000, the maximum is 22,963, the average is 20,006, and the standard deviation 116.61. These numbers show that the subsampling and supersampling worked correctly. There are a few models with a higher than average vertex count, because they were unable to processed by the PMP mesh decimation algorithm. However they are still included in the final dataset because their vertex count is close enough to 20,000.

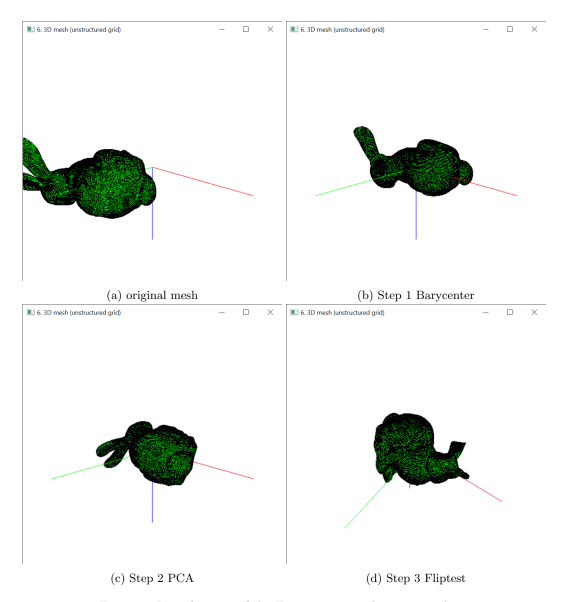


Figure 5: Visualization of the Four step normalization pipeline.

4 Feature Extraction

There are two different types of features used to describe the meshes. The first are features which consist of a single float value, the scalar features. And the second are the histogram features, which for a histogram with n bins contain n float values representing the fraction of values contained in a bin. The definition of these features is discussed in the following subsections.

4.1 Scalar Features

4.1.1 Surface area

The surface area of the mesh is the sum of all the face areas. In the dataset all the faces are triangles, therefore the surface area is the sum of all these triangles. The area of a triangle is calculated using the $triangle_area$ function provided by the pmp library. The area of a triangle t with vertices u, v, and w is defined as follows:

$$area(t) = 0.5 * N((t_v - t_u) \times (t_w - t_u))$$
(3)

Where N(x) is the Euclidean norm of a vector x. The surface area feature F_{SA} for a given mesh M is than

$$F_{SA}(M) = \sum_{i} area(t_i). \tag{4}$$

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where t_i is the i^{th} triangle of mesh M. The minimum value this feature can have is 0, but it does not have an obvious theoretical maximum. The maximum as is displayed in $Table\ 1$ is 32.96, with an average of 1.54 and standard deviation of 1.65, which shows that this feature has large outliers. Therefore the feature cannot be normalized by extent normalization, and instead standardization has to be used.

Feature encoding	Min	Max	Avg	Stddev
SA	0.02194	32.961956	1.53768275	1.648700958
BBV	0.001874	0.986091	0.261523862	0.2203638
ECC	0.000004	0.956016	0.148089259	0.182812364
CIR	0	0.978213	0.075533213	0.137364272
DI	0.74698	1.519733	1.055276205	0.088772773

Table 1: Scalar feature summary pre standardization

4.1.2 Bounding Box Volume

The bounding box of a mesh is a rectangle that can be represented as two points, the opposite corners bot and top. The volume V is easily calculated as the $width \cdot length \cdot height$.

$$bot = \begin{bmatrix} x_{min} \\ y_{min} \\ z_{min} \end{bmatrix} top = \begin{bmatrix} x_{max} \\ y_{max} \\ z_{max} \end{bmatrix}$$

$$V(bot, top) = (x_{max} - x_{min}) \cdot (y_{max} - y_{min}) \cdot (z_{max} - z_{min})$$
(5)

The bounding box volume feature F_{BBV} for a mesh M with its bounding box opposite corners M_{bot} and M_{top}

$$F_{BBV}(M) = V(M_{bot}, M_{ton}). (6)$$

The minimum value a bounding box can have is 0 when the mesh has no faces, and 1 when the bounding box is equal to the unit cube. Which is verified by the numbers in *Table 1* for the BBV feature.

4.1.3 Eccentricity

The eccentricity of a mesh the ratio of the biggest eigenvalue of the covariance matrix to the smallest. The alglib library is once again used to do PCA and obtain the eigenvalues of the shape covariance matrix. For the eigenvalues e_1 , e_2 , and e_3 where e_1 is the largest value and e_3 the smallest, the eccentricity is

$$F_{EC}(M) = M_{e_3}/M_{e_1} \tag{7}$$

Where M_{e_3} and M_{e_1} are the smallest and largest eigenvectors of mesh M respectively. This describes the relation between the biggest and smallest eigenvectors, or also between the number of points spread in the x direction and the z direction. An object like a sphere or a head will have a high eccentricity, because their vertices are not spread out in a single direction. The picture of the head in figure 6 has an eccentricity of 0.72, while the leopard on the right has an eccentricity of 0.03.

The eccentricity can go close to 0 for long objects or reach 1 for a perfect spherical form. Furthermore, this feature could also be extended to also describe the relation ratio between the other eigenvalues, M_{e_2} / M_{e_1} or M_{e_3} / M_{e_2} .

4.1.4 Circularity and Compactness

Circularity is the inverse of compactness, a measure which represents how compact a shape is. From this perspective, circularity shows how "circular" a shape is. In 2D, compactness is minimized by a perfect circle, where its value is 1. The formula for 2D compactness is constructed as the dimensionless ratio between a shape's perimeter (P) and area (A), with constants added to the denominator such that it produces the value 1 for a circle. The formula is the following:

$$C_{2D} = \frac{P^2}{4\pi A} \tag{8}$$

Because the system requires such a metric to be used for 3D shapes, a similar formula has been derived. We begin by replacing perimeter and area with surface area (S) and volume (V), their equivalent 3D metrics:

$$\frac{S}{V}$$
 (9)

As the result must be dimensionless, these measures must be raised to adequate powers so that their units of measurement cancel out:

$$\frac{S^3}{V^2} \tag{10}$$

Lastly, the constant (x) for the denominator must be determined by ensuring the derived formula will give 1 as the result if the shape in question is a perfect sphere:

$$\frac{(4\pi R^2)^3}{x(\frac{4}{3}\pi R^3)^2} = 1\tag{11}$$

Solving for x, we obtain the result 36π . Thus, the final compactness formula is:

$$C_{3D} = \frac{S^3}{36\pi V^2} \tag{12}$$

As mentioned at the start of this question, we use the inverse metric of compactness, circularity, due to the fact that it produces values only between 0 and 1; 1 if the shape is a sphere, and very close to 0 if the shape has a very uneven contour.

$$Circularity = \frac{36\pi V^2}{S^3} \tag{13}$$

In the program, the two models in *Figure 7* below return expected results for circularity. For the human head (a), the computation returns a value of 0.311646, much larger than the one for the leopard (b), 0.059514.

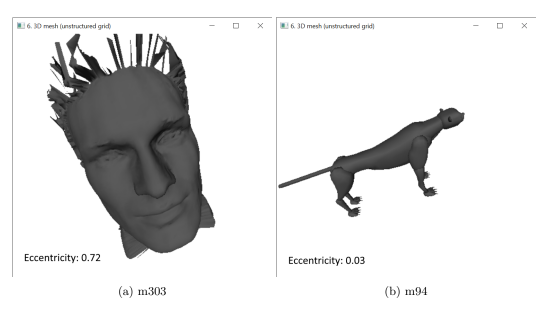


Figure 6: Two models with different eccentricity values

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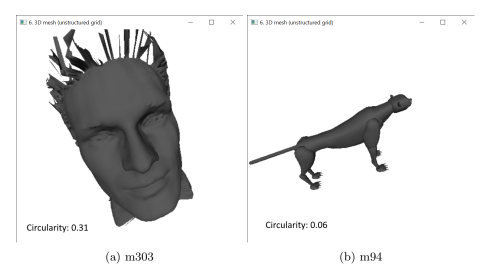


Figure 7: Two models with different circularity values

4.1.5 Diameter

The diameter feature represents the maximum distance between two points within the shape. As our shapes have been normalized to fit within a unit cube, the maximum value for the diameter is $\sqrt{2}$, its diagonal.

Computing the diameter can be done through multiple approaches. The naive solution is to try every possible pair of points, resulting in a complexity of $O(n^2)$. As our shapes have around 20,000 vertices, this is not a scalable approach (a single computation can take up to 1 minute).

An approximation for the diameter value is obtained instead by using random samples. By sampling 2000 points from the mesh and running the brute force method, we obtain an approximation of the diameter much faster, reducing the number of steps to just 2000^2 .

While this approximation is fast, it's accuracy is not at all guaranteed. To increase the chances of reaching the ground truth diameter value, we can repeat this technique several times and only keep the maximum value of each approximation. Running the sampling 10 times has proven to almost always reach the ground truth within 7 seconds, down from 58 (*Table 2*).

Shape	Method	Result	Time
	Brute Force	1.01355	58s
m303	2000 samples	1.01022	<1s
111303	5 * (2000 samples)	1.01022	4s
	10 * (2000 samples)	1.01355	7s
	Brute Force	1.01043	58s
m94	2000 samples	1.00442	<1s
11194	5 * (2000 samples)	1.01043	4s
	10 * (2000 samples)	1.01043	7s

Table 2: Diameter result and timing on two models.

4.1.6 Normalizing features

The three features bounding box volume, eccentricity, and circularity all fall in the range [0,1]. If the surface area feature is also to be normalized in this range, extent normalization has to be used through its minimum and maximum value. However this feature has large outliers. For example, if the minimum of 1 and maximum of 36 is used to normalize, the distance between a surface area of 3 and 1 would become 0.0571. Even though these 2 meshes would be very different in terms of surface area, their distance does not reflect this. Therefore standardization is used for this feature, and to be consistent, it will also be used for all the other features. Each

new feature value will be calculated with the following formula

$$f'_{i,j} = \frac{f_{i,j} - \mu_i}{\delta_i}, \quad i \in \{SA, BBV, ECC, DI, C\}$$

$$\tag{14}$$

Where μ_i is the average for feature i, δ_i the standard deviation, and $f_{i,j}$ the feature value for the *jth* mesh. This standardization changes the average of a feature to 0 and the standard deviation to 1. Meaning that if two different feature values differ one standard deviation from each other, there distance will be one.

4.2 Histogram features

For extracting the histogram features random vertices pairs are chosen to calculate the different features. For each histogram feature a histogram is created using a minimum and maximum value based on their theoretical maximum. These results will than be used to adjust the minimum and maximum so that there wont be any bins that are universally unused. The number of bins used bins that will finally be used is set at 12.

For the histogram features several graphs are displayed showing the histograms of two different classes. This is only for an indication of how the feature might work, it certainly does not mean that the differences in values is the same for all classes, or that all classes share a similar distribution within their own class. It can however be used to show whether the feature itself works correctly when it gives similar results for similar models, as is the case in several of the histograms. A prediction could be made that these features will help differentiate between different classes.

4.2.1 Sampling vertices

Ideally, for the histogram features, every possible combination of vertices should be considered. This means that for the histogram features that need two points, the number of combinations is approximately n^2 , where n is the total number of vertices. And for the triangle area, or angle, approximately n^3 combinations are possible. Because calculating the features for all these combinations is not doable, especially not with 20,000 vertices, only a sample of these combinations will be chosen. Obviously, the higher the sample size, the better, as the only constraint is obtaining a reasonable computation time.

It is important that each combination of vertices has the same probability to be selected as a sample. This is done by randomly sampling points from the uniform distribution using the c + + standard library function rand(). If a given pair of vertices is identical, the pair will be discarded. Note that for each model and each feature the same seed is used, so that the results are reproducible.

If for the distance between two pairs of vertices, n samples are generated, the probability that one of these combinations is included is $1 - (\frac{1}{n^2})^n$. However, if this same probability has to be achieved for the features that require 3 or 4 random samples, the sample size has to be increased by an order of magnitude, i.e. n^2 vertex combination samples have to be generated. This is obviously not achievable in a reasonable time, therefore the same sample size is used for all the histogram features.

The actual sample size that is used in the system is determined by maximizing the sample size while keeping a manageable computation time. The number of samples generated are 100,000 resulting in a computation time of 400ms. This time is considered reasonable because it allows the features of new models to be calculated in real time which is important for when a query is submitted, because a new input mesh has to be normalized and its features need to be calculated.

4.2.2 D1 - distance to barycenter

This feature calculates the distance between a randomly generated point and the barycenter. The distance D between two vectors u and v is calculated as follows:

$$D(u,v) = \sqrt{\sum_{i=1}^{3} (u_i - v_i)^2}.$$
 (15)

The minimum distance is obviously 0, and the maximum distance is equal to the diameter of a unit cube, $\sqrt{3}$, which is an unrealistic scenario. In order to evaluate the quality of the features, the histogram distributions of different classes will be plotted on a graph. The blue lines represent the histogram distributions of models m93, m94, m95, m96, and m97, which are all leopard-like animals, and the yellow lines are models m1120, m1121, m1122, m1123, and m1124, which are air planes. These graphs will be made for each of the histogram features. Figure 8 indicates that the feature will not be useful for differentiating between classes. There is both large variance between the two classes, as well as within the classes.

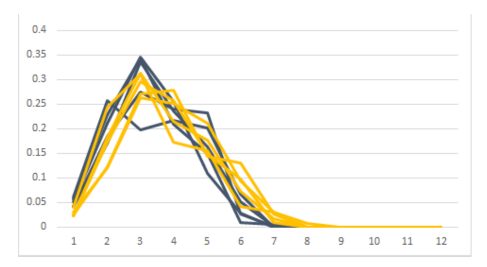


Figure 8: Histogram distributions for feature D1 for different classes. Yellow = air planes, Blue = leopards

4.2.3 D2 - distance between two points

This time two random points are generated and the same distance function as for the D1 feature is used. The minimum and maximum distances are the same as for the previous feature, although the distances will generally be higher.

Figure 9 shows that the two classes have different types of distributions, however there is a lot of variation within the classes themselves.

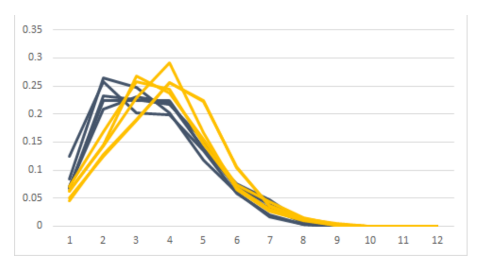


Figure 9: Histogram distributions for feature D2 for different classes. Yellow = air planes, Blue = leopards

4.2.4 D3 - area of a random triangle

To get a random triangle, three random points are generated. The area of this triangle is calculated the same way as when calculating the total surface area. The square root of this value is then added to the histogram, since this feature is no longer the length of a line segment, but a 2D area. Figure 10 shows a difference between the two classes, however there is also some within-class variation.

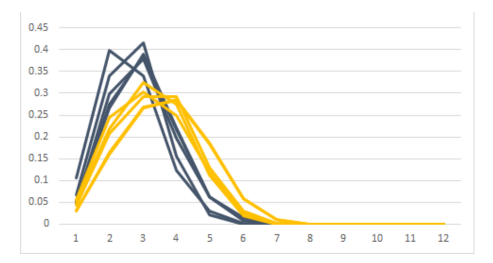


Figure 10: Histogram distributions for feature D3 for different classes. Yellow = air planes, Blue = leopards

4.2.5 D4 - volume of a random tetrahedron

Four random points a, b, c, and d are generated so that together they will be the vertices of the tetrahedron. The formula used for calculating the volume is:

$$V(a, b, c, d) = \frac{|(a - d) \cdot ((b - d) \times (c - d))|}{6}$$
(16)

Then, the cube root of this value is taken, as it is a 3D volume. Figure 11 shows a significant difference between distributions of air planes and leopards. Furthermore, the within-class differences are minimal. Therefore, this might indicate that D4 is a good feature, at least for distinguishing planes from leopards.

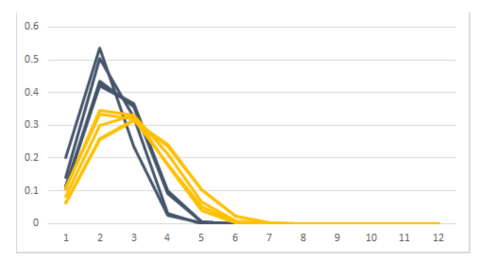


Figure 11: Histogram distributions for feature D4 for different classes. Yellow = air planes, Blue = leopards

4.2.6 A3 - angle between 3 random vertices

The last histogram feature uses three random points u, v, and w to calculate the angle between them. These three points create a triangle that has three angles, however the order in which they are generated determines which angle is chosen. Therefore, all three possible angles are calculated, and the maximum of those is chosen. For each angle the distance is represented by using two vectors that share a similar point. For example, the vectors a = v - u and b = w - u give one possible angle of the triangle. The formula used for calculating an

4.3 Feature vector 15

angle given two vectors is as follows:

$$Angle(a,b) = atan2(N(a \times b), (a \cdot b)), \tag{17}$$

where N gives us the euclidean norm of the vector, and atan2 the angle in the euclidean plane. The minimum value that the angle could take is 0, and the maximum is π .

In Figure 12 it can be seen that the two classes are quite dissimilar, however one of the air planes seems to be an outlier compared to the rest of its class.

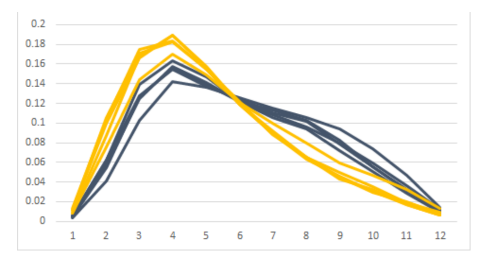


Figure 12: Histogram distributions for feature A3 for different classes. Yellow = air planes, Blue = leopards

4.3 Feature vector

A model i can be represented in the system using its feature vector F_i . This is created by combining all of the previously discussed features $f_{i,j}$.

$$F_i = \{f_{1,j}, f_{2,j}, ..., f_{10,j}\}$$
(18)

Each $f_{i,j}$ is a list of k values, where k is 1 for the scalar values and 12 for the histogram values. The feature vector is thus a list of 65 values (Table 3).

i	feature name	k	\min	max
1	surface area	1	-0.919356	19.06002
2	bounding box volume	1	-1.178278	3.28805
3	eccentricity	1	-0.810042	4.419431
4	circularity	1	-0.549875	6.57143
5	diameter	1	-3.472866	5.231975
6	distance to barycenter	12	0	1.060660
7	distance between two points	12	0	$\sqrt{2}$
8	area of a random triangle	12	0	0.848528
9	volume of a random tetrahedron	12	0	0.565685
10	angle between 3 random vertices	12	0.942478	π

Table 3: Feature vector format and standardized extreme values.

5 Matching

All the features discussed in the previous section are calculated for each mesh and put in a single feature vector $x = (x_1, ..., x_n)$ where each x_i is a single feature. To compare different meshes the distance is to be calculated between their two feature vectors. If feature vector x is compared with a different feature vector $y = (y_1, ..., y_n)$, each of their features will be compared individually and the sum of these distances will be regarded as the distance between the two feature vectors.

5.1 Distance calculation

Each feature has k values; for scalar features k = 1 because it only has one value. For histograms k is equal to the number of bins of that histogram. The distance function L(x, y) gives the total euclidean distance between two feature vectors x and y:

$$L(x,y) = \left(\sum_{i=1}^{n} d(x_i, y_i)^2\right)^{1/2},\tag{19}$$

where d is the distance function between two features:

$$d(x_i, y_i) = \frac{\left(\sum_{j=1}^k |x_{i,j} - y_{i,j}|^2\right)^{1/2}}{k}.$$
 (20)

 $x_{i,j}$ is the j^{th} value of the i^{th} feature of feature vector x, and the same goes for y. The sum of all the distances between feature values is divided by k so that each feature has the same weight, i.e. a histogram with 8 values should not be weighted 8 times as much as a scalar feature which only has 1 value.

5.2 Query result

These distances are calculated for each mesh in the database, including the query item, if it exists. The result is thus a list of distances for a query item i $L_i = d_1, ...d_k$, where k is the number of items in the database, and is ordered in ascending order. With an input query size s, the first s elements of L are then returned as the query result. An alternative approach that uses a kd-tree as a data structure to facilitate faster searches.

5.3 kd-tree

The system, should return a number of relevant models k as specified by the end user. In the previously described method, the distances are being sorted first resulting in a time complexity of $\mathcal{O}(n\log n)$. Alternatively, for a small size of k, instead of sorting the result, the unsorted list of distances could simply be traversed k times and finding a new nearest neighbour each time. This has a time complexity of $\mathcal{O}(kn)$. In this system, the size of the dataset is not so high that this becomes unreasonable, however, to be able to scale up with an even larger dataset, a search structure can be created. The data structure used is called a kd-tree and is a way to partition the data to allow for fast searches. The cost of creating such a tree is $\mathcal{O}(n\log n)$, however this can be precomputed and stored server side so that this overhead cost is irrelevant for the user. Searching for the k nearest neighbours in this tree has an average time complexity of $\mathcal{O}(k\log n)$, which thus greatly accelerates the search process.

The library used in this system for creating and searching kd-trees is ANN. In addition to the previously mentioned search, ANN has the functionality of an approximate search, which can further accelerate the search in exchange for a user specified loss in accuracy. In this system however, because the number of models and dimensions is manageable, the exact k-nearest neighbours are returned instead.

6 Dimensionality reduction

In order to assess the quality of our feature vectors, a means to visualize the entire dataset and how distinguishable each class is from the others is desirable. As it is impossible to directly plot our 65-dimensional feature vectors into a scatter plot, the dimensionality reduction technique known as t-Distributed Stochastic Neighbor Embedding (t-SNE) has been used to transcribe our feature space into two dimensions. (van der Maaten & Hinton, 2008) describe the technique in detail. This section discusses how this was achieved, then the resulting plot is shown.

6.1 Perplexity Parameter

T-SNE takes as input, aside from the data itself, three parameters: the number of desired dimensions (2 in our case), the dimensionality of the data (65 dimensions) and perplexity. The latter is defined in (van der Maaten & Hinton, 2008) as 2 to the power of the Shannon entropy of the probability distribution represented by our multidimensional space. In practice, however, perplexity is a parameter that must be experimented with to determine the best results.

Several values between 0 and 100 have been tested in an attempt to select the one that minimizes the "error" reported by the algorithm, which is actually the Kullback-Leibler divergence between the original multidimensional data and the 2D result. The table below displays the results:

Perplexity	K-L Divergence
25.0	0.927
50.0	0.915
75.0	0.905
100.0	0.909

Table 4: t-SNE perplexity and K-L divergence results.

6.2 Resulting Plot

However, t-SNE is not deterministic, as it initializes its gradient function randomly. Instead of finding a perfect perplexity, it could be more appropriate to run the algorithm multiple times with the same perplexity and choosing the result with minimal error. After several attempts, the lowest error obtained was 0.660 on 100.0 perplexity. This can be seen in Figure 13.

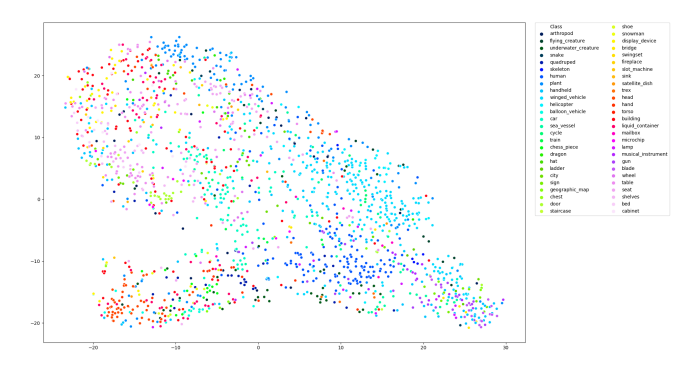


Figure 13: t-SNE plot on feature vector space: 100.0 perplexity and 0.660 error

While the figure is hard to read due to the large amount of points and classes, some information that confirms the feature values are not random can be derived from it. For example, the clusters of humans and winged vehicles are fairly well delimited, and so are the clusters for heads and doors.

18 7 EVALUATION

7 Evaluation

The final step is to evaluate the performance of the retrieval system. This is done by analysing the query result Q which is a list of items $x_1, ..., x_s$, where s is the size of the query result and each x is an item in the database. As it was explained in the matching section, this list is ordered by ascending distance to the query item l. First off, there are several values that describe a query result, namely the true positives (TP), false positives (FP), true negatives (TN), and false negatives (FN). TP is the number of items with the correct class label in a query, FP the number of wrong items in a query, TN the number of wrong items not in the query result, and FN the number of correct items not in query.

$$TP = |\{y \in Q \mid c_y = c_l\}|, \\ FP = s - TP, \\ TN = k - TP - FP - FN, \\ FN = N_{c_l} - TP,$$

where c_i is the class label of item i, k the number of items in the dataset, and N_{c_l} the number of items in the class of query item l.

7.1 Evaluation metric

There are many evaluation metrics that can be used. Furthermore, there are different aspects of the system for which the performance can be evaluated. For example, the performance for a single item in the dataset, the average performance of a class, and the overall performance of the program. In the evaluation of this system, the performance for each item will be calculated, and from those measurements the class average, and overall performance will be derived. The overall performance is calculated as the average of all items, not an average of all classes.

The performance metric chosen are mean average precision (MAP) and precision at k. For evaluation purposes, the model for which the querying is performed is not included in its own query result list. The distance of a model to itself will always be very close to 0 and would thus give a misleading precision if it were to be included. The definition of these performance metrics and why they are chosen is discussed in the following two sections. Furthermore the results of the evaluation are shown in table 5.

7.1.1 MAP

Two interesting evaluation metrics are precision and recall. Precision shows how many of the returned items in the query are of the correct class, while recall shows how many of the correct class are represented in the query. The problem with recall however is that it is highly dependent on the size of the query, and therefore not useful in evaluating the performance of a system that produces a ranked query. A query of size 1 can never have a recall of 1, assuming that the class size is higher than 1. Therefore the recall is not a useful performance metric for this system, because it gives little insight. A large query size will always result in a high recall and a low query size in low recall. The more interesting metric is precision whose definition is given below.

$$P = \frac{TP}{TP + FP} \tag{21}$$

Another problem that arrives is what query size the evaluation metric should be calculated for. A universal query size is undesirable, because there are large discrepancies between the sizes of different classes. A class with only five items will have a terrible performance when the query size is set to 100, because the maximum precision is $\frac{5}{100}$. Furthermore, even for a specific class, choosing the query size is an arbitrary decision. Therefore the Mean Average Precision (MAP) is used.

The query result that is returned by the matching algorithm is an ordered list of ascending distance values of size d-1, where d is the size of the database. The MAP for a class is the mean of all the average precisions of its models' query results. The average precision for a query result of a given model is defined as:

$$AvgP = \frac{\sum_{k=1}^{n} (P(k) \cdot rel(k))}{m}$$
 (22)

Where rel(k) is the indicator function and equals to 1 if the k^{th} item in the query result has the correct label and 0 otherwise, and m is the number of models in the relevant class excluding the model itself. The MAP for a class is then the mean of all of these average precision values.

7.1 Evaluation metric 19

MAP might punish some classes too much when they contain an outlier. If it does, then, for each model in the class, the outlier will yield a precision close to 0. In this case, it might be possible that even a perfect system cannot obtain a precision of 1, because of an outlier or even a wrongly sampled mesh. Furthermore, calculating precision for items that are very far in the query result can be useless when the user only wants a small result. For example, perhaps a user would not want to look at more than 20 models, therefore only the first 20 results would be relevant for them. If a system would return 20 items that have the correct class label, but the other 20 items in the class appear in the back of the query result, the average precision would be close to 0.5, while the returned query is a perfect result for the user. Despite these downsides of MAP, it is still chosen as a strict metric to judge the quality of the entire query results. One of the upsides of MAP is that it considers the position of the correct classes, which is not the case for the regular precision.

7.1.2 Precision at k

In addition to MAP, another performance metric is used, the precision at k. For this a value for k has to be decided, which will be the length of the query result. Then, for this smaller sized query, its precision will be calculated. As it was previous discussed, a user might only want to look at a certain number of items. Precision at k then measures how many correct items the user will see. In the evaluations in this report, the value for k will be set to 10. However because some classes contain less than 10 models, k will be $min(k, n_c - 1)$ where $n_c - 1$ is the number of models in class c, excluding the model that is queried for.

A downside of precision at k is that it does not account for the position of models. For example a result query with only the first five items being the correct class and a second result query with only the last five items being the correct class, will have the same precision at k. Intuitively, however, the first query result is better.

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class name	MAP	Precision at k	Observed model count
winged vehicle	0.416308	0.666527	239
balloon vehicle	0.116555	0.14375	16
helicopter	0.123383	0.22	35
arthropod	0.0545013	0.103704	27
human	0.394029	0.648993	149
trex	0.0641384	0.0666667	6
flying creature	0.0340165	0.05	26
quadruped	0.120444	0.232258	31
snake	0.0491144	0.0833333	4
underwater creature	0.131797	0.231429	35
blade	0.20551	0.257895	19
head	0.292354	0.496104	77
hand	0.232334	0.176471	17
skeleton	0.409468	0.170471	5
	0.409408	0.4	4
torso			_
bridge	0.103303	0.1	10
building	0.08743	0.161225	98
chess piece	0.203151	0.276471	17
chest	0.0761211	0.047619	7
city	0.0492937	0.0894737	19
display device	0.111166	0.18	40
door	0.301497	0.439286	28
dragon	0.0524088	0.0666667	6
fireplace	0.0110214	0	6
bed	0.0463207	0.0535714	8
cabinet	0.0725405	0.0833333	9
seat	0.159862	0.327273	77
shelves	0.172277	0.276923	26
table	0.167187	0.342308	78
geographic map	0.0562935	0.1	11
gun	0.112295	0.178571	28
hat	0.152967	0.2125	16
ladder	0.0446827	0	4
lamp	0.0475397	0.0863636	22
liquid container	0.0473331	0.19661	59
mailbox	0.0373931	0.19001	7
	0.022003	0.047619	7
microchip musical instrument			22
	0.0932159	0.15	
plant	0.194478	0.391111	135
satellite dish	0.0023526	0	4
sea vessel	0.098209	0.229545	44
shoe	0.0831799	0.107143	8
sign	0.0635423	0.1125	16
sink	0.00389432	0	4
slot machine	0.0768283	0	4
snowman	0.15229	0.166667	6
swingset	0.0991632	0	4
staircase	0.00529275	0	7
handheld	0.0996813	0.202459	122
car	0.24536	0.483636	110
cycle	0.125922	0.138462	13
train	0.0980175	0.0952381	7
wheel	0.0992868	0.158824	17
total	0.200032	0.343097	1796
	3.20002	1.02000.	1 3 0

Table 5. Performance evaluation using MAP and precision at $\mathbf k.$

7.2 Dissimilar classes 21

7.2 Dissimilar classes

The performance results show that there is a large discrepancy between the performances of different classes. The worst classes even have a precision at k of 0, meaning that for none of the models in its class the result query gave even a single relevant model in the first 10 results. This leads to the questions of what causes this bad result, and what can be done to improve performance for this class. For the first question, four of the models of the staircase class are shown in figure 14. It is clear that the overall shapes of these models are very different, which leads to their feature values being very different as well.

Some characteristics of staircases are present in these pictures, namely the large number of 90 degree edges leading to the top. A feature could be added that would encapsulate this characteristic so that all these models will have a distance of 0 for this feature. However, even in this case, all the other feature values will be very different, making it very unlikely that their overall distances will be small enough to yield good results.



Figure 14: Models m1733, m1734, m1735, and m1736 of the staircase class.

Looking at another bad performing class, the mailboxes, a less obvious difference in models is visible. They look a lot more alike, however their distances are still too large. The seemingly small differences in shape have a large impact on their feature values. For example the first and fourth mailboxes are shorter than the second and fourth. Furthermore the volume of the actual box is different in each model. Finally there are some more small differences, such as the rectangular shape below the box in the fourth model, and the extended rectangles in the third shape.

It could thus be said that even though these models look alike, their actual shapes are still too different for the system. The small differences cause the feature values differences to be too large. Both the mailbox class and the staircase class have a precision at k of 0, however, the mailbox class has an MAP value four times that of the staircase class, meaning that the relative distances of mailboxes are smaller than that of the staircases. This makes sense because, intuitively, one would say that the mailboxes are more similar to each other than the staircases.

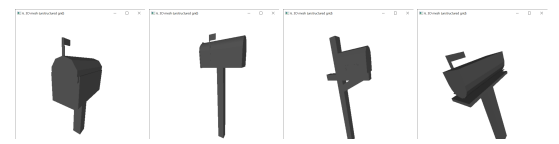


Figure 15: Models m515, m516, m517, and m517 of the mailbox class.

7.3 Query examples

In this section several query results are shown for several different classes. Firstly, we will query for a model from the quadruped class which has an MAP of 0.12, and precision at k of 0.23. For this specific model, five out of the nine other models are of the same class. Three of these wrong models, the hands, can be explained by them having a similar elongated shape and their fingers resembling legs. The turtle does not have a similar shape however it does also have four legs.

Secondly a query result for a model of the flying creature class is shown, which has a MAP of 0.034, and precision at k of 0.05. In this case, only two out of the nine models of the query result are of the same class.

22 8 CONCLUSION

Six of these, however, are airplanes and have a very similar shape to the bird. The ship is the only of these models that is an unreasonable result. The system thus returns models that are arguably very similar to the input shape, although not from the same class.

Finally the query result for one of the best performing classes, humans, shows a perfect query result. The good results for humans show that they are somewhat clustered in the dataset. This means that for a certain amount of models they exist in the middle of the clusters and thus all of their neighbours are other humans. This cluster can be visualized in the above t-SNE plot, *Figure 13*.

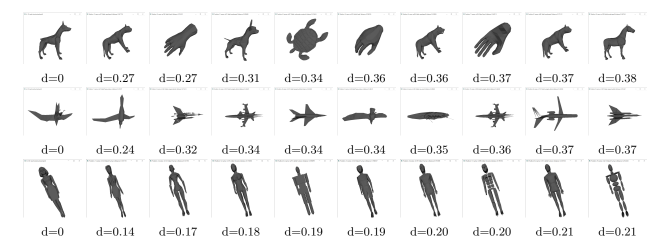


Figure 16: Query results and their squared distances for models m92, m42, and m134 respectively.

8 Conclusion

This section begins by presenting a summary of our work. Some limitations of our project are then discussed, followed by possible improvements to our implementation and future work.

8.1 Summary

This report presented our attempt at a content-based 3D shape retrieval system using scalar features that, given a 3D shape, returns the 10 most similar shapes (excluding itself) from a preprocessed Princeton Shape Benchmark database of 1796 models that are grouped into 53 leaf classes. This search is sped up by the use of a kd-tree and k-nearest neighbours. A visualization of the 65-dimensional feature space is also provided as a 2D plot created using t-SNE and seaborn.

Our system performs much better on some classes, such as humans and winged vehicles, than on some other classes, like staircases and mailboxes, with an average precision at k=10 of 0.34 and an average MAP of 0.20. We discussed these results in section 7, and provided some query examples.

8.2 Limitations

Due to the choice at the start of the project to use 20,000 vertices and the 1800+ shape database, database-wide evaluation, recomputing features and reprocessing the database were very slow tasks. In addition, because of the database size, bugs with individual shapes were much harder to detect. For example, some of our shapes started looking like in *Figure 17*.

This turned out to be a problem with the PMP library. Some of the shapes were not correctly read by the library, resulting in some of their feature values being affected (i.e. smaller area due to less triangles). Due to the nature of the problem and the time required to solve it, we elected to repair as much of the database as possible and accept that meshes with too many samples would risk to lose some of their faces. This is clearly a priority for any future version of this system.

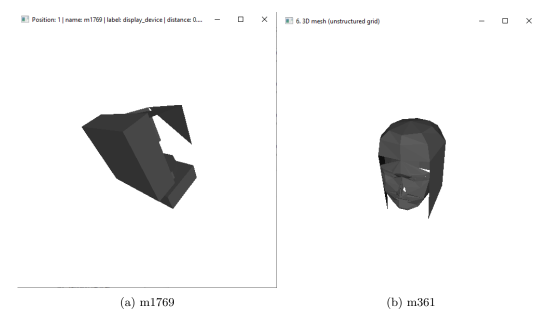


Figure 17: Models with faces missing.

8.3 Improvements & future work

There are several parts of our system that could be improved. First and foremost, a different mesh decimation library should be used if we cannot identify what causes the problem with the PMP mesh decimation.

Secondly, we could improve the speed of our application by changing the way we compute the diameter. While the method we use right now is much faster than the brute force approach, there are at least two better options to consider. First would be the convex hull approximation which is then used to calculate the diameter much faster, as mentioned in (Xue, J. et al., 2019). Another approach would be to use the technique described by (Har-Peled, S., 2001).

Thirdly, we could improve the quality of our matching function by experimenting with different distance functions for our feature vectors. For example, the effectiveness of our histogram features can be increased by computing their distance a different way, for instance by using the Earth Mover's Distance suggested by (Rubner, Y. et. al., 2000).

Lastly, the weights for each feature are set to 1 in the current system. In future work these weight parameters could be optimized to yield the best results, i.e. bad features would be weighted less and good features would weight more.

On the visualization side, it would be possible to make a plot that better resembles our system's performance by using a distance matrix as input for the t-SNE algorithm. This was omitted due to time constraints and not being possible in the Python implementation we decided on for ease, but could be much more insightful on how our system actually behaves. In addition to this, it could be possible to perform the entire matching on a simple 2D feature space if our dimensionality reduction resulting plot was reliable enough (with well defined clusters).

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