Fast Nearest Neighbor Search in Medical Image Databases

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Abstract

We examine the problem of finding similar tumor shapes. Starting from a natural similarity function (the so-called ‘max morphological distance’), we showed how to lower-bound it and how to search for nearest neighbors in large collections of tumor-like shapes.

Specifically, we used state-of-the-art concepts from morphology, namely the ‘pattern spectrum’ of a shape, to map each shape to a point in $n$-dimensional space. Following [19, 36], we organized the $n$-d points in an R-tree. We showed that the $L_\infty$ ($= \max$) norm in the $n$-d space lower-bounds the actual distance. This guarantees no false dismissals for range queries. In addition, we developed a nearest-neighbor algorithm that else guarantees no false dismissals.

Finally, we implemented the method, and we tested it against a testbed of realistic tumor shapes, using an established tumor-growth model of Murray Eden[15]. The experiments showed that our method is up to 27 times faster than straightforward sequential scanning.

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1 Introduction

This paper proposes an algorithm which rapidly searches for “similar shapes”. Such an algorithm would have broad applications in electronic commerce (e.g., ‘find shapes similar to a screw-driver’), photo-journalism [20], etc., but would be particularly useful in medical imaging. During the past twenty years, the development of new modalities such as Computed Tomography (CT) and Magnetic Resonance Imaging (MRI) have substantially increased the number and complexity of images presented to radiologists and other physicians. Additionally, the recent introduction of large scale PACS (Picture Archival and Communication Systems) has resulted in the creation of large digital image databases. A typical radiology department currently generates between 100,000 and 10,000,000 such images per year. A filmless imaging department such as the Baltimore VA Medical Center (VAMC) generates approximately 1.5 terabytes of image data annually.

An algorithm that would be able to search for similar shapes rapidly would have a number of useful applications in diagnostic imaging. Both “experts” such as radiologists and non-experts could use such a system for the following tasks:

1. Diagnosis/Classification: distinguish between a primary or metastatic (secondary) tumor based on shape and degree of change in shape over time correlating this with data about diagnoses and symptoms. Computer-aided diagnosis will be especially useful in increasing the reliability of detection of pathology, particularly when overlapping structures create a distraction or in other cases where limitations of the human visual system hamper diagnosis [37].

2. Forecasting/Time Evolution Analysis: predict the degree of aggressiveness of the pathologic process or try to distinguish a particular histology based on patterns of change in shape. In this setting, we would like to find tumors in the database with the similar history as the current tumor.

3. Data Mining: detect correlations among shapes, diagnoses, symptoms and demographic data, and thus form and test hypotheses about the development and treatment of tumors.

In all of the above tasks, the central problem is similarity matching: ‘find tumors that are similar to a given pattern’ (including shape, shape changes, and demographic patient data). We mainly focus on matching similar shapes. In section 6, we discuss how our approach can be naturally extended to handle more complicated queries.

Some terminology is necessary. Following [19], we distinguish between (a) range queries (e.g., find shapes that are within distance $c$ from the desirable query shape) and (b) nearest neighbor queries (e.g., find the first $k$ closest shapes to the query shape) An orthogonal axis of classification distinguishes between whole-matching and sub-pattern matching. In whole-matching queries, the user specifies an
S × S query image and requires images of S × S that are similar; in sub-pattern matching queries, the user specifies only a small portion and requires all the (arbitrary-size) images that contain a similar pattern.

In this work we focus on whole-matching, because this is the stepping stone for the sub-pattern matching, and all the problems listed above, as discussed in section 6.

For the whole matching problem, there are two major challenges:

- How to measure the dis-similarity/distance between two shapes. In the tumor application, as well as in most other shape applications, the distance function should be invariant to rotation and translation. Moreover, we would like a function that pays attention to details at several scales, as we explain later.

- Given such a distance function, how can we do better than sequential scanning of the whole database? This faster method, however, should not compromise the correctness: it should have no false dismissals; that is, it should return exactly the same response set as sequential scanning would do.

Next we provide solutions to the above two challenges. This paper is organized as follows: Section 2 gives the survey. Section 3 gives an introduction to morphology and tumor-shape modeling. Section 4 presents our main result: the lower-bounding of the ‘max-morphological’ distance, as well as a k-nearest neighbor algorithm, without false dismissals. Section 5 gives the experiments. Section 6 discusses how to use the proposed algorithms for more general problems. Section 7 gives the conclusions.

2 Survey

2.1 Multimedia Indexing

The state of the art in multimedia indexing is based on feature extraction [36, 19]. The idea is to extract n numerical features from the objects of interest, mapping them into points in n-dimensional space. Then, any multi-dimensional indexing method can be used to organize, cluster and efficiently search the resulting points. Such methods are traditionally called Spatial Access Methods (SAMs). A query of the form find objects similar to the query object Q becomes the query find points that are close to the query point q, and thus becomes a range query or nearest neighbor query. Thus, we can use the SAM to identify quickly the qualifying points, and, from them, the corresponding objects. Following [1], we call the resulting index as the ‘F-index’ (for ‘feature index’). This general approach has been used in several settings, such as searching for similar time-sequences [1] (e.g., trying to find quickly stock prices that move like MacDonalds), color images [18, 20], etc.

The major challenge is to find feature extraction functions that preserve the dis-similarity/distance between the objects as much as possible. In [1, 19] we showed that the F-index method can guarantee
that there will not be any false dismissals, if the actual distance is lower-bounded by the distance in feature space.

Mathematically, let $O_1$ and $O_2$ be two objects (e.g., time sequences, bitmaps of tumors, etc.) with distance function $D_{\text{object}}(\cdot)$ (e.g., the sum of squared errors) and $F(O_1), F(O_2)$ be their feature vectors (e.g., their first few Fourier coefficients), with distance function $D_{\text{feature}}(\cdot)$ (e.g., the Euclidean distance, again). Then we have:

**Lemma 1 (Lower-Bounding)** To guarantee no false dismissals for range queries, the feature extraction function $F(\cdot)$ should satisfy the following formula:

$$D_{\text{feature}}(F(O_1), F(O_2)) \leq D_{\text{object}}(O_1, O_2) \tag{1}$$

**Proof:** In [19].

Thus, the search for range queries involves two steps. For a query object $Q$ with tolerance $\epsilon$, we

1. Discard quickly those objects whose feature vectors are too far away. That is, we retrieve the objects $X$ such that $D_{\text{feature}}(F(Q), F(E)) < \epsilon$.

2. Apply $D_{\text{object}}(\cdot)$ to discard the false alarms (the clean-up stage).

### 2.2 Spatial access methods

Since we rely on spatial access methods as the eventual indexing mechanism, we give a brief survey of them. These methods fall in the following broad classes: methods that transform rectangles into points in a higher dimensionality space [32]; methods that use linear quadtrees [23] [3] or, equivalently, the $z$-ordering [51] or other space filling curves [17] [35]; and finally, methods based on trees (R-tree [27], k-d-trees [6], k-d-B-trees [56], hB-trees [41], cell-trees [26], etc.)

One of the most promising approaches in the last class is the R-tree [27] and its numerous variants (Greene's variation [25], the $R^+$-tree [58], R-trees using Minimum Bounding Polygons [34], the $R^*$-tree [5], the Hilbert R-tree [38], etc.). We use R-trees, because they have already been used successfully for high-dimensionality spaces (10-20 dimensions [18]); in contrast, grid-files and linear quadtrees may suffer from the 'dimensionality curse'.

### 2.3 Tumor growth models

Our target class is a collection of images of tumor-like shapes. As a preliminary testbed, we use artificial data generated by a certain stochastic model of simulated tumor growth. Our particular model is a discrete-time version of Eden's tumor growth model [15], whose idea is illustrated in Figures 1 and 2. At time $t=0$, only one grid-cell is 'infected'; each infected grid-cell may infect its four non-diagonal neighbors with equal probability $p$ at each time-tick.
On the basic Eden model, we have added the notion of East-West/North-South bias, to capture the effects of anisotropic growth patterns, due to anisotropies in the surrounding tissue (e.g., lesions shaped by their location within the lung, breast, or liver.) Thus, in our model, an infected grid-cell has probability \( p_{NS} \) to infect its North and South neighbors, and probability \( p_{EW} \) to infect its East/West ones, with \( p_{NS} \) not necessarily equal to \( p_{EW} \).

\[
\begin{array}{ccccccc}
6 & 9 & 9 \\
8 & 5 & 4 & 7 & 4 & 6 & 9 \\
6 & 3 & 2 & 1 & 6 & 8 \\
9 & 9 & 9 \\
\end{array}
\]

Figure 1: Lattice at \( t = 9 \). The infection time of each infected cell is marked.

\[
\begin{array}{ccccccc}
\text{t = 1} & \text{t = 5} & \text{t = 10} & \text{t = 25} & \text{t = 50} & \text{t = 100}
\end{array}
\]

Figure 2: Initial seed (left column) and snapshots of tumor at later time steps, with \( p = .7 \)

The Eden model is the simplest of a hierarchy of models: in increasing generality, we have the Williams-Bjerkness model [62], the Contact Process [31], and the Interacting Particle Systems [13, 40]. See [21, 30, 55, 10] for a more comprehensive survey on tumor growth models.

### 2.4 Shape Representation and Matching

Shape representation is an interesting enough problem to have attracted many researchers and generated a rich array of approaches [52]. There are two closely related problems: (a) how to describe a single shape compactly and (b) how to measure the difference of two shapes, so that it corresponds to the humanly perceived difference.

With respect to representations, the most popular methods are:

- representation through ‘landmarks’: for example, in order to match two faces, information about the eyes, nose, etc., are extracted manually [4] or automatically. Thus, a shape is represented by a set of landmarks and their attributes (area, perimeter, relative position etc). Again, the distance between two images is the sum of the penalties for the differences of the landmarks.
• representation through numerical vectors, such as (a) samples of the ‘turning angle’ plot [33] (that is, the slope of the tangent at each point of the periphery, as a function of the distance traveled on the periphery from a designated starting point) (b) some coefficients of the 2-d Discrete Fourier Transform (DFT), or, more recently, the (2-d) Discrete Wavelet Transform [43] or (c) the first few moments of inertia [20, 18]. In these cases, we typically use the (weighted) Euclidean distance of the vectors.

• representation through a simpler shape, such as polygonalization [24, 50, 54, 60, 39] and Mathematical Morphology [63, 47, 44, 11, 7], which we shall examine in detail next.

Among them, representations based on morphology are very promising, because they have two very desirable characteristics for our applications:

• they can be easily designed to be essentially invariant to rotation and translation (= rigid motions),

• they are inherently ‘multi-scale’, and thus they can highlight differences at several scales, as we explain next.

The multi-scale characteristic is important, especially for tumors, because the ‘ruggedness’ of the periphery of a tumor contains a lot of information about it [9]. Thus, given two tumor-like shapes, we would like to examine differences at several scales before we pronounce the two shapes as ‘similar’.

Even for general shapes, there exists substantial evidence that scale-space behavior is an important and highly discriminating shape “signature” [61, 42, 8].

3 Introduction to Morphology

Our goal is to choose a distance function between shapes which will be invariant to translation and rotation, and which will ‘give attention’ to all levels of detail. One such function is founded on ideas from the field of Mathematical Morphology. See [12] for a very accessible introduction. Next, we present the definitions and concepts that we need for our application. Table 3 lists the symbols and their definitions.

Some definitions are in order: Consider black-and-white images in 2-d space; the ‘white’ points of an image are a subset of the 2-d address space, while the background is, by convention, black. More formally, let \( \mathcal{X} \) (the “shape space”) be a set of compact subsets of \( \mathbb{R}^2 \), and \( \mathcal{R} \) be the group of rigid motions \( \mathcal{R} : \mathcal{X} \rightarrow \mathcal{X} \).

3.1 Introduction to Morphology

Mathematical Morphology is a rich quantitative theory of shape, which incorporates a multiscale component. It has been developed mainly by Matheron [48, 49], Serra [59, 14], and their collaborators.
Since the 1980’s, morphology and its applications have become extremely popular.

In mathematical morphology, mappings are defined in terms of a *structural element*, a “small” primitive shape (set of points) which interacts with the input image to transform it, and, in the process, extract useful information about its geometrical and topological structure. The basic morphological operators are: \( \oplus, \ominus, \odot, \bullet \) (dilation, erosion, Morphological Opening, and Closing, respectively), and they are defined as follows.

Let \( X_h \) denote the translate of shape \( X \) by the vector \( h \), and let \( H^* \) denote the symmetric of shape \( H \) with respect to the origin:

\[
H^* = \{-h \mid h \in H\}
\]  

**Definition 1** The dilation, \( X \oplus H^* \), of a shape \( X \subset \mathbb{R}^2 \) by a structural element \( H \), is defined as

\[
X \oplus H^* = \bigcup_{h \in H} X_{-h} = \left\{ (x, y) \in \mathbb{R}^2 \mid H_{(x,y)} \cap X \neq \emptyset \right\}
\]
Figure 3 shows a ‘butterfly’ shape $X$, dilated by the unit circle $H$. Intuitively, a dilation ‘blows-up’ the original shape $X$ by tracing its perimeter (and all the internal points) with a ‘brush’ of foot $H$.

We also use $mH$, $m \in \mathbb{Z}^+$ to denote $H \oplus H \oplus \cdots \oplus H$ ($m - 1$ times), i.e., a structural element of size $m$. Intuitively, if the structural element $H$ is the unit ball, then the shape $mH$ is a ball of radius $m$.

**Definition 2** The erosion, $X \ominus H^s$, of a shape $X \subset \mathbb{R}^2$ by a structural element $H$, is defined as

$$X \ominus H^s = \bigcap_{h \in H} X_{-h} = \left\{ (x, y) \in \mathbb{R}^2 \mid H_{(x,y)} \subseteq X \right\}$$  \hspace{1cm} (4)

Figure 3 shows the butterfly shape $X$ eroded by the unit circle $H$, which is a dilation of the complement of $X$. Intuitively, an erosion deletes part of the original shape $X$ by tracing its perimeter with an ‘eraser’ of foot $H$.

Two important composite Morphological operators are opening and closing.

**Definition 3** The opening, $X \circ H$, of a shape $X \subset \mathbb{R}^2$ by a structural element $H$, is defined as an erosion followed by a dilation:

$$X \circ H = (X \ominus H^s) \oplus H$$  \hspace{1cm} (5)

Figure 3 shows the opening of shape $X$. Intuitively, the opening is the set of points that a brush of foot $H$ can reach, when the brush is confined inside the shape, and is barely allowed to touch the periphery of the shape.

**Definition 4** The closing, $X \bullet H$, of a shape $X \subset \mathbb{R}^2$ by a structural element $H$, is defined as a dilation followed by an erosion:

$$X \bullet H = (X \oplus H^s) \ominus H$$  \hspace{1cm} (6)

Figure 3 shows the closing of shape $X$, which is the opening of the complement of $X$. Intuitively, the closing is the set of points that remain after an eraser of foot $H$ sweeps the outside of the dilated $X$.

Thus, the opening by a circle of radius $n$ in effect ‘cuts the corners’; that is, it eliminates the protruding details of the shape $X$, with radius less than $n$. Symmetrically, the closing ‘fills the concavities’ of the appropriate scale.

### 3.2 Granulometries and the Pattern Spectrum

The concept of the *Pattern Spectrum* as a compact shape-size descriptor has been developed by Maragos [46], based on earlier seminal work on openings of sets in Euclidean spaces by Matheron [48, 49, 28, 29], who called them *Granulometries*. Serra [59, 14] and his collaborators have used Lebesgue measures of openings by a size-parameterized family of structural elements to develop shape-size sensitive measurements of shape attributes which they called *Size Distributions*. 

8
Definition 5 The size distribution $y_X^H$ of a shape $X \in X$, with respect to a structural element $H$ is defined as

$$y_X^H = [f_{-M}^H(X), \ldots, f_1^H(X), |f_0^H(X)|, |f_1^H(X)|, \ldots, |f_M^H(X)|]^T$$

with

$$f_m^H(X) \triangleq \begin{cases} X \circ mH & 1 \leq m < M \\ X & m = 0 \\ X \bullet mH & -M \leq m \leq -1 \end{cases}$$

where $H$ is some structural element.

Intuitively, $|f_m^H(X)|$ is the area of a smoothed version of $X$ at scale $m$, i.e., for $|f_0^H(X)|$ is the area of $X$, $|f_1^H(X)|$ is the area of $X \circ H$, etc. In other words, the vector $y_X^H$, contains measurements of the area of $X$ at different scales, or degrees of shape smoothing, thus constituting the size distribution.

The pattern spectrum, as discussed by Maragos [46] contains exactly the same information. Its elements are backward differences of the size distribution. In other words, the size distribution can be thought of as the ‘cumulative pattern spectrum’. The intuitive meaning of the pattern spectrum is the amount of detail (= additional area) that the next closing will add, or that the next (larger-radius) opening will subtract. Figure 4 shows the pattern spectrum of a circular disc of radius 5, as well as of a square of side 10, with respect to a a unit disc structural element $H$. Notice that the disc has only one 'spike', at $m=5$, while the pattern spectrum of the square has details at several scales. (Of course, the situation would be reversed, if the structural element $H$ was the unit square).

The importance of the size distribution and the pattern spectrum is that they summarize important shape characteristics in the sense that they possess high discriminatory power, as reported in [2, 53].

3.3 Distance functions

Given two shapes $X_1$ and $X_2$, a natural distance function involves penalizing the non-common areas. Formally,
Definition 6 Let \( d(\cdot, \cdot) \) denote the area of the symmetric set difference distance measure, i.e., for \( X_1, X_2 \in \mathcal{X} \):

\[
d(X_1, X_2) = |X_1 \setminus X_2| = |X_1 \cup X_2| - |X_1 \cap X_2|
\]  

(9)

We can show that \( d(\cdot, \cdot) \) is a distance metric over \( \mathcal{X} \times \mathcal{X} \). However, we need a distance function that allows rotations and translations. This is achieved by requiring that the two shapes are first optimally aligned by allowable motions. Formally, we have a new distance function:

Definition 7 Define the floating shape distance \( d^*(\cdot) \) of two shapes \( X_1 \) and \( X_2 \) as

\[
d^*(X_1, X_2) = \inf_{R \in \mathcal{R}} d(X_1, R(X_2))
\]  

(10)

where \( \mathcal{R} \) is the set of rigid motions. The process of optimal alignment of two shapes is called registration.

The \( d^*(\cdot) \) distance is very natural and intuitive; it only fails in one account, namely, to consider details at several levels. Figure 5 illustrates the point: \( X_1 \) is a square, \( X_2 \) is an identical square, with a line segment coming out of its left side, and \( X_3 \) is identical to \( X_1 \), with a line segment cutting into it. At the current scale, the distance \( d^*(\cdot) \) among any pair of them is small. For example, if \( X_1 \) and \( X_2 \) are optimally aligned, making the two squares to coincide, then the area of the disjoint part is the area of the protruding line segment, which is negligible. However, the visual difference between the two is non-negligible. The same is true for \( X_1 \) and \( X_3 \). These counter-intuitive results can be remedied by applying the newly introduced tools of morphology: after applying a closing (see third column), we see that the protruding line segment in \( X_2 \) will make its presence more obvious. Similarly, after applying an opening (second column), the ‘cut’ in \( X_3 \) will become more obvious.

Thus, given any two shapes, each opening and closing will emphasize different details of their differences, resulting in a different value of \( d^*(\cdot) \). The question is how to combine all these scale-dependent penalties to arrive at a single number. The solution we propose is to take the maximum difference. More formally,
Definition 8 Define the Max Morphological Distance

\[ d^H_\infty : X \times X \mapsto \mathbb{R}_+ \]  

(11)

as

\[ d^H_\infty (X_1, X_2) \triangleq \max_{-M \leq m \leq M} d^a \left( f^H_m (X_1), f^H_m (X_2) \right) \]  

(12)

with \( f^H_m (X) \) defined in Eq. 8.

For the remainder we assume some fixed structural element \( H \) (e.g., the unit ball), and we drop these indices.

The intuitive meaning of the \( d_\infty () \) distance function is the following:

- compute \( d^a () \), that is take the two shapes \( X_1 \) and \( X_2 \), align them optimally, and compute the area of the disjoint parts

- take their closings using a disk of radius 1, 2, ... \( M \); in each case, compute the \( d^a () \) of the resulting shapes

Figure 5: Three different scales of shapes \( X_1 \), \( X_2 \) and \( X_3 \)
do the same for openings, with a disk of radius 1, 2, ... M

pick the maximum difference, and report it as the distance of the two shapes.

Lemma 2 The function $d_{\infty}$ is indeed a distance metric between elements of $X$.

Proof: See Appendix A.

4 Problem Definition - Proposed solution

The problem we focus on is the design of fast searching methods that will operate on the tumor database to locate the most similar object to the query object. The (dis-)similarity is measured by the max-morphological distance (Eq. 12). We focus on both range queries as well as on nearest-neighbor queries. We have three obstacles to overcome:

- what features to use (i.e., how to map tumor-shapes into n-d points)
- how to prove that the above mapping is contractive, that is, it obeys the Lower-Bounding Lemma (Lemma 1).
- how to use the resulting F-index on the feature space, so that we can answer nearest-neighbor queries with respect to the actual distance (as opposed to the distance in feature space)

Next we present the proposed solutions to these three problems.

4.1 Features

Our goal is to derive features that will capture a lot of the shape information, that will be rotation and translation invariant, and that will lead to a feature-distance function that fulfills the Lower-Bounding Lemma. Given the success of the pattern spectrum as a means to capture shape information [2, 46, 45, 63], we started with its coefficients as good features. More specifically, we use the coefficients $y_x$ of the size distribution (Eq. 7), which contains exactly the same information as the pattern spectrum.

Thus, we ‘penalize’ two shapes for differences at several scales. The question is, what is the best way to combine the penalties of each scale? A natural choice is to pick the maximum among the penalties. This is identical to the $L_{\infty}$ norm of the two feature vectors, and it leads to the following distance function:

Definition 9 Define the Max Granulometric Distance $\delta_{\infty}(\cdot)$ of two shapes $X_1$, $X_2$ as

$$\delta_{\infty}^H(X_1, X_2) = \max_{-M \leq m \leq M} |y_{X_1}(m) - y_{X_2}(m)|$$ (13)
4.2 Lower-Bounding

Our next challenge is to show that the distance in feature space (i.e., the max-granulometric distance $\delta_\infty()$) lower-bounds the actual distance $d_\infty()$. This is necessary to guarantee no false dismissals.

**Lemma 3 (Max-Morphological Distance Bounding)** The max-granulometric distance $\delta_\infty()$ lower-bounds the max-morphological distance $d_\infty()$:

$$\delta_\infty(X_1, X_2) \leq d_\infty(X_1, X_2), \quad \forall X_1, X_2 \in \mathcal{X}$$ (14)

**Proof:** Observe that

$$d^*(X_1, X_2) \geq ||X_1| - |X_2||$$ (15)

with equality achieved if and only if there exists some rigid motion $R \in \mathcal{R}$ which brings all points in $X_2$ (or $X_1$) in registration with points in $X_1$ ($X_2$, respectively). Then

$$d^*(f_m(X_1), f_m(X_2)) \geq ||f_m(X_1)| - |f_m(X_2)||$$ (16)

and

$$\max_{-M \leq m \leq M} d^*(f_m(X_1), f_m(X_2)) \geq \max_{-M \leq m \leq M} ||f_m(X_1)| - |f_m(X_2)||$$ (17)

Recall that the left-hand side is the definition of $d_\infty$ and the right-hand side is the definition of $\delta_\infty$. Thus, the proof is complete. QED

The latter result guarantees completeness of range queries.

By keeping the dimensionality of the spectra space small (say $M = 5 \rightarrow 2M+1 = 11$ features) we can use an F-index which, as we show later, results in considerably faster access of large image databases.

4.3 Nearest-neighbor algorithm

We have just described a good set of features, namely, the $2M+1$ entries of the cumulative pattern spectrum (≡ size distribution) of an image, as well as proved that the resulting $\delta_\infty()$ distance lower-bounds the actual distance. Thus, the resulting ‘F-index’ will guarantee no false dismissals upon range queries.

The next problem is to find the $k$-nearest neighbors of a query image, given that the images of the collection have already been mapped into $n$-d points and organized in a SAM. Algorithms to find the $k$-nearest neighbors of a given point already exist, using a branch-and-bound algorithm [22], and have been applied to R-trees recently [57].

However, the SAM search will return the $k$-nearest neighbors with respect to the max-granulometric distance $\delta_\infty()$, as opposed to the max-morphological distance $d_\infty()$ that we really want.

Here we describe the algorithm that finds the actual $k$-nearest neighbors in any F-index where the Lower-Bounding Lemma (Lemma 1) holds:
Algorithm 1 (k-nn)
/*uses an F-index to return the k-nearest neighbors according to the actual distance*/
/*input: query object Q; # of nn k*/
/*output: k-nearest objects X_1, X_2, ..., X_k*/
1. Search the SAM to find the k-nn wrt the feature distance $D_{\text{feature}} (\delta_\infty$ in our case).
2. Compute the actual distance $D_{\text{object}}(Q, X)$ ($d_\infty(Q, X)$ in our case) for all the k candidates $X$, and return the maximum $\epsilon_{\text{max}}$.
3. Issue a range query with the feature vector $F(Q)$ of the query object $Q$ and $\epsilon_{\text{max}}$ on the SAM, retrieve all the actual objects, compute their actual distances $D_{\text{object}}()$ from $Q$, and pick the nearest $k$.

We prove that the above algorithm will give no false dismissals:

Lemma 4 Algorithm 1 guarantees no false dismissals for k-nn queries.

Proof: Let $X_k$ be the shape returned as the k-th nearest neighbor by the algorithm (step 3); let $Y$ be an object that is the j-th nearest neighbor (with $j < k$). Then

$$D_{\text{object}}(Q, Y) \leq D_{\text{object}}(Q, X_k)$$ (18)

Suppose for a moment that the algorithm fails to return $Y$. We will show that this leads to a contradiction: After the range query is issued in step 3, all eliminated shapes (including $Y$) must have feature distance $D_{\text{feature}}$ greater than $\epsilon_{\text{max}}$, that is

$$D_{\text{feature}}(Q, Y) > \epsilon_{\text{max}}$$ (19)

and, from the Lower-Bounding Lemma (Lemma 1),

$$D_{\text{object}}(Q, Y) \geq D_{\text{feature}}(Q, Y) > \epsilon_{\text{max}}$$ (20)

However,

$$\epsilon_{\text{max}} \geq D_{\text{object}}(Q, X_k)$$ (21)

(since $X_k$ was obviously retrieved by step 3). Combining the last two inequalities, we obtain

$$D_{\text{feature}}(Q, Y) > D_{\text{object}}(Q, X_k)$$ (22)

which contradicts Eq. 18. QED
5 Experiments

To test the speed of our approach, we implemented our method and ran experiments. Next we describe the set up, as well as our results and observations, for range queries and for nearest neighbor queries.

Testbed: We generated 20,000 black-and-white 128 × 128 pixel images of tumor shapes based on Eden’s model of tumor growth. Each image contains a tumor that either (a) grows uniformly in all eight directions, (b) is biased vertically and horizontally with slower growth along the diagonals, (c) is restricted along one direction (blocked by a barrier such as a bone), or (d) is restricted along two directions (cone-shaped). Within each of these four classes of growth, we vary

- the number of iterations, which affects the size of the tumor;
- the directional bias \( p_{NS}/p_{EW} \), which affects the ratio of height to width.

We performed experiments for varying database sizes \( N \), by choosing \( N \) images among the 20,000 ones.

Competing methods:

- straightforward sequential scan: This is the simple brute force algorithm. Given a query image, the algorithm goes through all images in the database and computes its max-morphological distance from the query image, keeping track of the images with the minimum distance. Because the algorithm is comparing images on a pixel-by-pixel basis, it is extremely inefficient.

- F-index with an \( n \)-d R-tree: On insertion, the size distribution, or cumulative pattern spectrum \( y_X \), of each image of the database has been computed and the \( n \)-dimensional vector has been inserted into an R-tree. Given a query image, its cumulative pattern spectrum is computed, and is then submitted for a range or \( k \)-nearest neighbor search in the R-tree, as we discussed before.

Measurements: We are interested in the response time, that is, the time until the last actual hit is returned to the user (after the system has discarded possible false alarms). For some small settings we report actual (wall-clock) time, from the \texttt{time} utility of \textit{UNIX} \( ^{TM} \). However, the time \( t_{mm} \) to compute the max-morphological distance between two images is high \( (t_{mm} = 12.69 \text{ sec on average}) \) and shows small variance (standard deviation of 0.036sec). Thus, to accelerate the execution of experiments on large databases, we \texttt{time} all the other steps of the algorithms involved, and simply ‘charge’ a delay of \( t_{mm} \) seconds for each max-morphological distance computation that we omit.

Hardware and software The methods were implemented in 'C' and \textit{KornShell} under \textit{UNIX} \( ^{TM} \). The experiments ran on a dedicated Sun SPARCstation 5 with 32Mb of main memory, running SunOS
4.1.3. The disk drive was a FUJITSU M2266S-512 model ‘CRANEL-M2266SA’ with minimum positioning time of 8.3 ms and maximum positioning time of 30 ms.

We present experiments on range queries as well as nearest neighbor queries. We also give some pictures of the images that have been returned.

5.1 Range queries

Figure 6: Response time vs. response-set size $a$ for range queries (a) with seq. scanning (b) without seq. scanning

We asked 20 queries on a database of $N = 1,000$ images for both methods. Figure 6(a) plots the response time for the F-index as a function of the response-set size $a$ (i.e., number of actual hits, after the false-hits have been eliminated), for several values of the tolerance. It also shows the response time for sequential scanning for comparison, which is estimated to take 12697.6 seconds. Figure 6(b) shows only the proposed F-index method, in more detail. Figure 7 displays this data in a table. The performance gap between the two methods is very large: our method achieves 15-fold to 27-fold savings.

5.2 Nearest Neighbor Queries

We ran queries with $k=2,3,4$, and 10 for several $N$. Figure 8 shows (a) the results of $k$-nearest neighbor queries with $k = 10$, for varying $N$, for the F-index method compared to the sequential scan algorithm, and (b) the results of $k=2,3,4$, and 10 for the F-index method only. Each data point represents the average response time (in seconds) for 100 random query images taken from the database.

Figure 9(a) shows response time vs. $k$ (= 2,3,4,10) for $N = 10,000$ and $N = 20,000$ for both methods. Figure 9(b) shows response time vs. $k$ for $N = 10,000$ and $N = 20,000$ for the proposed method only.
<table>
<thead>
<tr>
<th>response-set size</th>
<th>seq scan (1)</th>
<th>F-index (2)</th>
<th>ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>$a$</td>
<td>time (sec)</td>
<td>time (sec)</td>
<td>(1) : (2)</td>
</tr>
<tr>
<td>2</td>
<td>1269.76</td>
<td>720.25</td>
<td>17.63</td>
</tr>
<tr>
<td>4</td>
<td>1269.76</td>
<td>660.83</td>
<td>19.21</td>
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<td>7</td>
<td>1269.76</td>
<td>499.15</td>
<td>25.44</td>
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<td>9</td>
<td>1269.76</td>
<td>613.03</td>
<td>20.71</td>
</tr>
<tr>
<td>12</td>
<td>1269.76</td>
<td>564.96</td>
<td>22.48</td>
</tr>
<tr>
<td>14</td>
<td>1269.76</td>
<td>705.85</td>
<td>17.99</td>
</tr>
<tr>
<td>15</td>
<td>1269.76</td>
<td>520.10</td>
<td>24.41</td>
</tr>
<tr>
<td>17</td>
<td>1269.76</td>
<td>528.76</td>
<td>24.01</td>
</tr>
<tr>
<td>17</td>
<td>1269.76</td>
<td>621.38</td>
<td>20.43</td>
</tr>
<tr>
<td>17</td>
<td>1269.76</td>
<td>688.00</td>
<td>18.46</td>
</tr>
</tbody>
</table>

Figure 7: Table of (a) response-set size $a$ (b) response time for seq. scan (c) response time for F-index (d) ratio of seq. scan time to F-index algorithm time

Again, each data point represents the average response time over 100 queries.

The observations are the following:

- Our proposed algorithm is 3-7 times faster than sequential scanning, even for a large value of $k$ (e.g., 10) for the nearest neighbors;
- The savings of the proposed method compared to sequential scan seems to increase with the database size $N$;
- Response time grows slowly with $k$.

### 5.3 Sample output

Here we illustrate that the max-morphological distance function $d_{\infty}(\cdot)$ seems to capture the perceptual distance between two shapes. Figure 10 shows a few query images (left column) and their corresponding 3-nearest neighbors according to the max-morphological distance. Since the query images were drawn from the database, the first nearest neighbor is identical to the query shape (which is a sanity check for our algorithms and implementations). Notice how similar the other 2 nearest neighbors are, for both query shapes.

Finally, Figure 11 illustrates the realism of Eden’s model. Figure 11(a) shows the whole mamogram, highlighting the tumor shape; (b) shows the tumor magnified; (c) shows the tumor shape after it has been thresholded (and thus becomes a black-and-white image); and (d) shows the nearest neighbor that was retrieved from our testbed of 20,000 synthetic tumor shapes. The similarity of the real tumor with its synthetic nearest neighbor is striking.
Figure 8: (a) Response time vs. db size(N), for \( k = 10 \) nm queries for both seq. scan and F-index. (b) Response time vs. \( N \) for \( k=2,3,4,\)and 10.

6 Discussion

As mentioned in the introduction, we focus on whole-matching queries for 2-d shapes. In this section we discuss how to use the proposed algorithms as a stepping stone to more complicated problems, specifically (a) sub-pattern matching (b) time-evolution pattern matching and (c) matching on additional data (demographic, etc.)

1. Sub-pattern matching: The problem is to detect a similar \( s \times s \) shape within a collection of \( S \times S \) bitmaps \( (s \leq S) \). The idea is to use sliding, overlapping windows, as we suggested in [19] for time sequences. That is, we assume that the smallest query will be at least of size \( s_{\text{min}} \times s_{\text{min}} \) and we pre-process each \( S \times S \) data image by considering all the possible \( s_{\text{min}} \times s_{\text{min}} \) windows in it; then we extract \( n \) features from each window, mapping each window into a point in \( n \)-d space. Thus, the problem is reduced to the whole-matching problem, which we have just solved. Algorithms to minimize the space requirements of the feature vectors, as well as to handle queries of arbitrary sizes, have been suggested in [19] for 1-d time sequences; all these techniques can be easily extended to 2-d images (and, in fact, arrays of arbitrary dimensionality).

2. Time-evolution: Currently, we considered static, 2-d images. By introducing time, we have sequences of images, which can be viewed as 3-d arrays, if we envision time as one more dimension. We could extend our 2-d algorithms to handle such 3-d shapes: The morphological concepts that we mentioned before (opening, closing, pattern spectrum) can be trivially extended to 3-d shapes. Therefore, our 2-d algorithms can be trivially extended to accommodate temporal patterns.
<table>
<thead>
<tr>
<th>db size</th>
<th>seq scan time (sec)</th>
<th>F-index time (sec)</th>
<th>ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>200</td>
<td>2539.52</td>
<td>675.74</td>
<td>3.76</td>
</tr>
<tr>
<td>400</td>
<td>5079.04</td>
<td>1084.19</td>
<td>4.68</td>
</tr>
<tr>
<td>800</td>
<td>10158.68</td>
<td>2422.04</td>
<td>4.19</td>
</tr>
<tr>
<td>1000</td>
<td>12697.60</td>
<td>2375.44</td>
<td>5.35</td>
</tr>
<tr>
<td>2000</td>
<td>25395.20</td>
<td>4814.56</td>
<td>5.27</td>
</tr>
<tr>
<td>4000</td>
<td>50790.40</td>
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<td>14743.66</td>
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</tr>
<tr>
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<td>126976.00</td>
<td>20002.95</td>
<td>6.35</td>
</tr>
<tr>
<td>12000</td>
<td>152371.20</td>
<td>33973.27</td>
<td>4.49</td>
</tr>
<tr>
<td>20000</td>
<td>253952.00</td>
<td>39340.32</td>
<td>6.46</td>
</tr>
</tbody>
</table>

Table 2: Values for 10-nn queries: (a) db size N (b) response time for seq. scan (c) response time for the proposed algorithm (d) ratio of seq. scan time to proposed algorithm time

3. Inclusion of non-spatial data: The diagnosis for a tumor-like shape has to take into account not only the shape, but also additional information: e.g., whether/how close is it to an organ (lung, liver, etc.), demographic characteristics of the patient (age, gender, etc.) Our algorithm can easily incorporate such data by augmenting the n-dimensional feature vectors with a few more numerical features (age of patient, distance of tumor from liver, etc). Thus, our feature space will have n spatial features (the size-distribution coefficients of the tumor shape) plus n' non-spatial features; the rest of our algorithms will remain intact.

7 Conclusions

We have focused on fast searching for similar shapes with emphasis on tumor-like shapes. To solve the problem, we used a multi-scale distance function, the so-called ‘max-morphological’ distance. This distance function is based on modern signal processing methods, and specifically mathematical morphology. The distance is invariant to rotations and translations, and gives similar attention to all levels of detail (‘scales’). From the database end, we used the ‘Feature index’ (F-index) approach [1, 19], which is the latest in multimedia indexing.

The main contribution of this work is that it manages to couple the max-morphological distance with the F-index. This is done by using the coefficients of the size distribution as features, and by showing that the $L_\infty (= \text{max})$ distance in the resulting feature space lower-bounds the max-morphological distance. Given the Lower-Bounding Lemma (Lemma 1), this guarantees no false dismissals for range queries.

Additional contributions are the following:
Figure 9: Response time vs. $k$ for $N = 10,000$ and $N = 20,000$ (a) with seq. scanning (b) without seq. scanning

- The design and implementation of a nearest neighbor algorithm on an F-index, which provably guarantees no false dismissals
- The implementation of the proposed method and the experimentation on a synthetic, but realistic, database of tumor-like shapes. There, the proposed method achieved dramatic speed-ups (up to 27-fold) over the straightforward sequential scanning.
- A survey of the literature on tumor-growth modeling. Given the administrative and patient-privacy problems in obtaining large collections of real X-rays, we believe that the above citations and algorithms will provide an excellent starting point for database researchers who want to work in this area and need a realistic testbed for their algorithms.
- The introduction of the basic morphological concepts (dilation, erosion, pattern spectrum, etc.) in an intuitive way, so that these powerful tools will become more accessible to database researchers.

Future research should focus on applications and extensions of the proposed method for several modalities including Computed Radiography, CT, MRI, Ultrasound, and Nuclear Medicine, as well as non-radiologic images in areas such as dermatology and pathology. The algorithm could be incorporated for general use in a large-scale PACS and serve as a powerful tool for both diagnostic and research purposes.

References

[1] Rakesh Agrawal, Christos Faloutsos, and Arun Swami. Efficient similarity search in se-
Figure 10: Query images (left column) and their nearest neighbors, according to the max-morphological distance.


Figure 11: (a) a real tumor within a mamogram (b) magnification of the tumor (c) its black-and-white (thresholded) version (d) the most similar synthetic tumor


A Proof that $d_\infty$ is a distance metric

Lemma: The function $d_\infty$ is indeed a distance metric between elements of $X$

Proof: We start with $d_\infty$. We have to verify the following properties:

- $d_\infty(X_1, X_2) \geq 0$
- $d_\infty(X_1, X_2) = 0$ if and only if $X_1 = X_2$: trivial because $m$ can equal 0.
- $d_\infty(X_1, X_2) = d_\infty(X_2, X_1)$: follows from symmetry of $R$, and $d(\cdot, \cdot)$.
- Triangle Inequality: $d_\infty(X_1, X_2) \leq d_\infty(X_1, X) + d_\infty(X, X_2)$.

\[
d_\infty(X_1, X_2) \leq \max_{-M \leq m \leq M} \left\{ d^*(f_m^H(X_1), f_m^H(X)) + d^*(f_m^H(X), f_m^H(X_2)) \right\} \tag{23}
\]

\[
\leq \max_{-M \leq m \leq M} d^*(f_m^H(X_1), f_m^H(X)) + \max_{-M \leq m \leq M} d^*(f_m^H(X), f_m^H(X_2)) \tag{24}
\]

\[
= d_\infty(X_1, X) + d_\infty(X, X_2) \tag{25}
\]

Thus, the proof is complete. QED
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