ADAPTIVE HYBRID PICTURE CODING

INTERIM SCIENTIFIC REPORT

by

Richard A. Jones
Principal Investigator

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In order to recognize 3-dimensional objects in 2-dimensional scenes a shape description, complete enough to determine the 3-dimensional object, must be recovered from the scene. Although it is possible to recover some 3-dimensional information, for instance shape from shade (21) or shape from texture (13), in general the information recovered will be a 2-dimensional representation of the object. Many methods are available for interpreting 3-dimensional objects in 2-dimensional arrayed range images (3), (20); however this is considered as a somewhat different problem, since depth information can be derived from the scene with little or no ambiguity.
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# TABLE OF CONTENTS

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PART ONE</strong></td>
<td></td>
</tr>
<tr>
<td>Recognition of Partially Occluded 3-Dimensional Objects Using Information From Concavities</td>
<td>1</td>
</tr>
<tr>
<td>I Introduction</td>
<td>1</td>
</tr>
<tr>
<td>II Shape Space And Feature Vectors</td>
<td>2</td>
</tr>
<tr>
<td>The Measurement Vector</td>
<td>4</td>
</tr>
<tr>
<td>Rotation Translation and the Measurement Vector</td>
<td>5</td>
</tr>
<tr>
<td>The Size Variable and The Effects of Scaling</td>
<td>6</td>
</tr>
<tr>
<td>Generalized Description for Irregular 2-Dimensional Shapes</td>
<td>7</td>
</tr>
<tr>
<td>III Critical Points From Real Data</td>
<td>14</td>
</tr>
<tr>
<td>Shape Dictionary for 3-Dimensional Objects</td>
<td>21</td>
</tr>
<tr>
<td>Critical Point Determination for the Problem Text</td>
<td>23</td>
</tr>
<tr>
<td>IV Cognitive Stage</td>
<td>24</td>
</tr>
<tr>
<td>V Results</td>
<td>32</td>
</tr>
<tr>
<td>VI Discussion</td>
<td>33</td>
</tr>
<tr>
<td>VII Conclusions</td>
<td>34</td>
</tr>
<tr>
<td>Appendix</td>
<td>36</td>
</tr>
<tr>
<td>Bibliography</td>
<td>39</td>
</tr>
<tr>
<td><strong>PART TWO</strong></td>
<td></td>
</tr>
<tr>
<td>A Minimax Risk Quantizer for Noisy Sources</td>
<td>62</td>
</tr>
<tr>
<td>I Introduction</td>
<td>62</td>
</tr>
<tr>
<td>II General Quantization</td>
<td>63</td>
</tr>
<tr>
<td>III Risk Quantization</td>
<td>65</td>
</tr>
<tr>
<td>IV Minimax Risk Quantization</td>
<td>67</td>
</tr>
<tr>
<td>Known Source Distribution</td>
<td>70</td>
</tr>
<tr>
<td>Known Noise Distribution</td>
<td>78</td>
</tr>
<tr>
<td>V Results and Conclusions</td>
<td>80</td>
</tr>
<tr>
<td>References</td>
<td>83</td>
</tr>
</tbody>
</table>
PART ONE
RECOGNITION OF PARTIALLY OCCLUDED 3-DIMENSIONAL OBJECTS
USING INFORMATION FROM CONCAVITIES

I INTRODUCTION

In order to recognize 3-dimensional objects in 2-dimensional scenes a shape
description, complete enough to determine the 3-dimensional object, must be
recovered from the scene. Although it is possible to recover some 3-dimensional
information, for instance shape from shade [21] or shape from texture [13], in
general the information recovered will be a 2-dimensional representation of the
object. Many methods are available for interpreting 3-dimensional objects in 2-
dimensionally arrayed range images [3], [20]; however this is considered as a
somewhat different problem, since depth information can be derived from the scene
with little or no ambiguity. The representation recovered from regular 2-dimensional
intensity scenes will consist of a mapping from an object-centered coordinate space
to a viewer-centered coordinate space [2]. This mapping represents any reorientation
of the 3-dimensional object within its coordinate system which results in a change
with regard to the viewer.

Currently there are two approaches to forming an object description under these
conditions. In the first approach [4], a 3-dimensional representation of all the
objects to be recognized is stored. Then when a 2-dimensional scene is to be
analyzed, the 3-dimensional model is transformed to obtain its 2-dimensional
equivalent. The corresponding 2-dimensional versions are then matched against those
extracted from the scene. The main difficulty with this approach is the amount of
upfront processing required to compute an adequate number of transformations to
required to compute an adequate number of transformations to represent the possible
object views.

In the second approach, the transformed object representations are calculated
a-priori and the corresponding 2-dimensional descriptions stored for the scene
analysis algorithm. Obviously the 2-dimensional descriptions must be compact enough so that a sufficient number of views can be stored without exceeding memory restrictions. Fourier descriptors [18], [19], and boundary curvature [26] have been used to describe 3-dimensional objects in this manner however, they are inadequate when conditions of occlusion and boundary noise respectively are allowed. In this paper, a method for 3-dimensional object identification similar to the two above but with much greater immunity to conditions of occlusion and boundary noise is presented. The method is believed to be applicable in many industrial and military systems where efficient and reliable identification is necessary.

In section II, the concepts of shape space and shape vector are introduced. The formalization of these concepts provides a basis upon which shape contours can be analyzed, even in the context of incomplete knowledge (occlusion). The underlying structure for constructing the shape vector is the critical point. In section III a new method for defining critical points is described, which in many cases offers improved performance versus those defined on the basis of curvature. In section IV a procedure for extracting these new critical points from discrete data is presented, and in section V the corresponding shape identification algorithm or cognitive stage is described. Finally, in section VI the procedure's ability to recognize partially obscured, complex 3-dimensional objects in 2-dimensional scenes is demonstrated.

II SHAPE SPACE AND FEATURE VECTORS

In order to distinguish one shape from another, or a partial shape as a part of a whole shape, some mechanism which examines the relationship between shape measurements is needed. One such mechanism which is ideal for this application is a shape space [24]. By introducing a shape space defined on the measurements, relationships between these measurements can be quantified and thereby compared.
The basic element of shape space is the shape vector $\frac{kZ}{j}$, where $j$ indicates the $j$th set of measurements from the $k$th shape. If there are $K$ shapes to be analyzed, then $k$ is integer valued from 1 to $K$. The range of $j$ depends on the total number of measurements possible on a given shape $k$. The shape vector $\frac{kZ}{j}$ consists of a measurement vector, $\frac{kM}{j}$, normalized by a scalar factor $\frac{kS}{j}$ referred to as the size variable. Thus,

$$\frac{kZ}{j} = \frac{kM}{j} / \frac{kS}{j} \quad 1$$

The measurement vector consists of $I$ measurements between predetermined points on the shape $kS$, represented notationally by

$$\frac{kM}{j} = \left( \frac{k_{m1}}{j}, \frac{k_{m2}}{j}, \ldots, \frac{k_{mI}}{j}, \ldots, \frac{k_{m1}}{j} \right) \quad 2$$

Combining (1) and (2) yields the final form of the shape vector

$$\frac{kZ}{j} = \left( \frac{k_{m1}}{j}, \frac{k_{m2}}{j}, \ldots, \frac{k_{mI}}{j}, \ldots, \frac{k_{m1}}{j} \right) \quad 3$$

For notational purposes it is convenient to rewrite (3) as

$$\frac{kZ}{j} = \left( \frac{k_{z1}}{j}, \frac{k_{z2}}{j}, \ldots, \frac{k_{z1}}{j}, \ldots, \frac{k_{z1}}{j} \right) \quad 4$$

where each $\frac{k_{zj}}{j}$ represents the $j$th measurement component normalized by the size variable. When constructing the shape vector two important properties must be preserved: 1) all measurements are made between a set $C$ of predefined points, 2)
the shape vector is independent of the size variable. If these conditions are met, then any two shapes which have the same shape vector are said to be equal with respect to the measurements.

The Measurement Vector

The exact nature of the measurement vector is described next. Let each element of $C$ be represented by the ordered pair $c_n = (x_n, y_n)$, where $x_n$ and $y_n$ are the spatial coordinates in $\mathbb{R}^2$. The displacement vector between any critical point $c_n$ and any reference point $c' = (x', y')$ is given as

$$c_n + c' = (x_n + x', y_n + y')$$

the normal Euclidean definition for displacement in $\mathbb{R}^2$. The resulting vector yields a displacement with respect to the origin. With this definition in mind, consider $I$ arbitrary critical points which constitute the $j$th measurement vector formed from $C$. Now form a reference point, $j\in = (x_j, y_j)$, by calculating the centroid of the $I$ points as

$$\bar{x}_c = \frac{1}{I} \sum_{i=1}^{I} x_i; \quad \bar{y}_c = \frac{1}{I} \sum_{i=1}^{I} y_i.$$ 

Then form the components of the measurement vector by calculating the displacement between each critical point and the feature centroid. Mathematically this is given by

$$j.m_i = ( jx_i - \bar{x}_c, jy_i - \bar{y}_c ); \quad i = 1 \text{ to } I.$$ 

This in effect translates all measurements to the origin of $\mathbb{R}^2$, in other words, the
measurement vector is an l-tuple of vectors originating from the origin of $R^2$. The relationships described by (6) and (7) are demonstrated graphically in figure 1.

**Rotation, Translation and the Measurement Vector**

It is a desirable goal to remove the effects of rotation, translation and scaling from the shape representation or to at least account for them. An examination of these effects for the first two transformations on the measurement vector is now discussed (the effects of scaling are closely related to the definition of the size variable and will be described later).

Since the rotation $R_0$ of the shape by an angle $\theta$ is linear [10], the rotation of the arbitrary set $\{c_1, c_2, c_3, ..., c_l\}$ by an angle $\theta$ about the origin results in a rotation of the $m_i$'s about the origin plus a displacement to the centroid of the rotated feature. This can be shown mathematically as

$$ j_{c_i}^* = R_\theta \{ j_{c_i} \} = R_\theta \{ j_{m_i} + \bar{c} \} = R_\theta \{ j_{m_i} \} + R_\theta \{ j_{\bar{c}} \} \quad 8 $$

where the latter two rotations are about the origin. The latter displacement term in (8) will be factored out by the subtraction in (7). Thus the angle of rotation between the rotated measurement vector and the original measurement vector can be calculated for any component $i$ as

$$ \theta = \text{SIGN}(6) \left[ \cos^{-1} \left( \frac{(j_{m_i}, j_{m_i}^*)}{||j_{m_i}|| \cdot ||j_{m_i}^*||} \right) \right] \quad 9 $$

where $(\cdot, \cdot)$ is the standard inner product and $||\cdot||$ is the standard norm for vectors.
in $\mathbb{R}^2$. From this it can be seen that the only difference between measurement vectors from the rotated set of critical points and the unrotated set is a rotation $\theta$ about the origin.

Similarly, a translation $T$, of the set $\{c_1, c_2, c_3, \ldots, c_I\}$ by a vector $c_t$ can be written as

$$j'_{c_i} = T\left\{jc_i\right\} = T\left\{jm_i + c\right\} = jm_i + jc + c_t$$ \hspace{1cm} 10

However, $j + c_t$ is just a new centroid which will be factored out in (7) leaving only the original $m_i$'s. Therefore, translation has no effect on the measurement vector.

The Size Variable and the Effects of Scaling

The second condition required for the formulation of the shape vector is that the shape vector must be independent of the size variable. This can be achieved by making the size variable a function of the measurement vector. One such function can be formulated as

$$j^s = \sum_{i=1}^I ||jm_i||$$ \hspace{1cm} 11

This formulation can be interpreted as the spatial energy contained in the measurement vector. With this value as the size variable the sum of all spatial energy in any shape vector always equals 1.

As a result, the effects of scale change can be removed. As an example, consider the previously mentioned set of arbitrary points. If they are scaled by a factor $\alpha$, their measurement vector will become
\[ J^M = \left( a_1 m_1, a_2 m_2, a_3 m_3, a_4 m_4, a_5 m_5, a_6 m_6 \right) \]

which has total energy given by

\[ E_t = ||a|| \sum_{i=1}^{I} ||m_i||. \]

This is equivalent to the size variable for \( J^M \) multiplied by \( a \). When the corresponding shape (feature) vector is calculated the scale factor \( a \) will be canceled by the size variable.

**GENERALIZED DESCRIPTION FOR IRREGULAR 2-DIMENSIONAL SHAPES**

Since initially only 2-dimensional shapes are considered the domain for a shape, called \( S \), is restricted to \( \mathbb{R}^2 \). The shape \( S \) consists of a bounded, simple, closed curve \( B \) [22] and the area enclosed by this curve, called the interior \( I \). In other words, \( S = B \cup I \). The shape description will consist of a finite set of measurements extracted from \( S \). These measurements will be made between a set of points \( C \), which are referred to as critical points. The set of critical points is a subset of \( B \).

In order to define this set \( C \), another structure called a concavity tree must first be described.

According to Sklansky [24], the concavity tree is a structure for describing simply connected silhouettes. If the shape \( S \) is convex, then the concavity tree is trivially the shape \( S \) itself. In this work, it is assumed that the shape is non-convex, this will be the case in most applications where irregular objects are to be processed. The concavity tree is described for the shape \( S \) by the following
procedure.

1) Compute the convex hull of the shape \( S \), called \( S^c \). This is defined operationally by

\[
S^c = C_h(S).
\]

The convex hull is the smallest convex set which contains \( S \).

2) Subtract the original shape \( S \) from its convex hull \( S^c \). This is equivalent to finding the intersection of \( S^c \) with the complement of \( S \) (\( S^c \)). The set formed by \( S^c \cap \overline{S} \) is most likely a disconnected subset of \( S^c \). Consider each connected element of this subset as a branch of the concavity tree (subshape). Denote each branch by \( S_i \), which indicates the \( i \)th component of \( S^c \cap \overline{S} \). Each \( S_i \) will have the same properties as \( S \).

3) To find the next level of the concavity tree, replace \( S \) by \( S_i \) and \( S_i \) by \( S_{ij} \). Repeat the above procedure for each level 1 branch of the concavity tree. The index \( j \) will denote the branches of the concavity tree at level 2.

4) To complete the concavity tree, continue with the above procedure progressing from \( \{S_i\} \) to \( \{S_{ij}\} \) to \( \{S_{ijk}\} \) etc. until all residual subshapes from the second step are convex. The set of shapes \( \{S_i, S_{ij}, S_{ijk}, ...\} \) for all \( i,j,k,... \) forms the concavity tree. Figure 2 shows an arbitrary shape \( S \) and its resulting concavity tree.

Since the concavity tree is based on the relationship between a shape (subshape) and its convex hull there is only one unique tree for each shape (subshape). However, it is possible for this tree to have a countably infinite number of levels and an uncountable infinite number of branches, for example see Koch curves [65]. In this work the class of shapes examined will be restricted to those which are sufficiently smooth as to have only a finite number of levels and branches, and thereby a finite number of critical points. This restriction still includes all shapes found in real applications and all but the most abstract found in any application.

Now that the concavity tree has been described, the definition for the critical
points can be considered. Let the set \( C \) represent the critical points for a shape \( S \) and \( \Psi(\bullet) \) the operation which yields \( C \) from \( S \),

\[
C = \Psi( S ) .
\]

The set \( C \) is equivalent to the union of all critical points extracted from all branches of the concavity tree of \( S \), i.e.,

\[
C = \bigcup_{n \text{ all levels}} C_n
\]

where,

\[
C_1 = \bigcup_{i \text{ all branches}} \Psi( S_i ) ,
\]

\[
C_2 = \bigcup_{j \text{ all branches}} \Psi( S_{ij} ) ,
\]

\[
C_3 = \bigcup_{k \text{ all branches}} \Psi( S_{ijk} ) ,
\]

etc...

The operation \( \Psi(\bullet) \) for any shape (subshape) must now be described. Consider any branch of the concavity tree \( ( S_{ijk...} ) \) and find its intersection with the boundary \( B \), so that

\[
B_{ijk...} = B \cap S_{ijk...}
\]

This intersection defines a line segment for each branch of the concavity tree. The
collection of all such line segments formed at each level of the tree is the basis for the critical point sets $C_n$. The critical points are finally defined as the endpoints of the $B_{ijk...}'s$. Figure 3 illustrates the boundary segments and critical points for the first level concavities of the shape in figure 2. Finally, the operation $\Psi(\cdot)$ can be summed up by the following hierarchy of point set relationships.

$$
C = \Psi(S),
= \bigcup_{n} C_n',
= \bigcup_{n} \left( \bigcup_{m} \bigcup_{i,j,k,...} \psi(S_{i,j,k,...m}) \right),
= \bigcup_{n} \left( \bigcup_{m} \left\{ \text{endpoints } \left[ B_{i,j,k,...m} \right] \right\} \right),
= \bigcup_{n} \left( \bigcup_{m} \left\{ \text{endpoints } \left[ B \cap S_{i,j,k,...m} \right] \right\} \right).
$$

These relationships, between all allowable shapes $S$ with their boundaries and their concavity trees, are unique and well defined. Therefore, the relationships between any shape $S$ and its corresponding critical point set $C$ are also unique and well defined.

Although these relationships are unique and well defined, there exists two special cases which can lead to some ambiguity if not handled properly. The first case occurs when two branches from the same level of the concavity tree, $S_{ij...mn}$ and $S'_{ij...mn}$, have the following property:

$$
S_{ij...mn} \cap S'_{ij...mn} = \text{a single point}
$$
but the intersection of the open sets formed by $S_{ij...m} - S_{ij...m'}$ is empty. When this happens the subsequent boundary segments associated with the two branches of the concavity tree, $S_{ij...mn}$ and $S_{ij...mn'}$, will in fact be a single line segment. To remove any ambiguity, the single point described in (20) will be used to partition this line segment into two separate ones. This single point will be called "critical" and will be considered as two separate points, each with singular associations to $S_{ij...mn}$ and $S_{ij...mn'}$ respectively.

The second case is similar to the first except involves an overlapping of critical points from different levels of the concavity tree. In other words, the sets $C_n$ are not mutually exclusive. This will not be a problem as long as the membership information within the $C_n$'s is retained along with spatial information in $\mathbb{R}^2$. Consequently, the location of critical points within $\mathbb{R}^2$ and the knowledge of the levels and branches of the concavity tree from which they originated are of equal importance.

As a rule of thumb, each branch of the concavity tree will generate two critical points. Each of these critical points can be represented by a triplet $(x,y,l)$, where $x$ and $y$ represent the location of the points in $\mathbb{R}^2$ and $l$ indicates the level of the concavity tree at which the points were generated; $x$ and $y$ are real valued numbers; $l$ is integer valued.

Since initially, only shapes which are invariant in $\mathbb{R}^2$ are allowed three transformations must be considered. The three transformations - scaling, rotation and translation - are all well known mappings from $\mathbb{R}^2$ to $\mathbb{R}^2$, and are represented symbolically by $S_a(S)$, $R_\theta(S)$ and $T(S)$ respectively. They are defined mathematically in the following. The functions $S_a(S)$, $R_\theta(S)$, $T(S)$ map a shape $S$
into a shape $S'$ such that all points $(x,y)$, elements of $S$ are mapped to points in $S'$ by

$$S_{\alpha} \left\{ (x,y) \right\} = (\alpha x, \alpha y) = (x', y'),$$

$$R_{\theta} \left\{ (x,y) \right\} = (x \cos \theta - y \sin \theta, x \sin \theta + y \cos \theta) = (x', y'),$$

$$T \left\{ (x,y) \right\} = (x + x_t, y + y_t) = (x', y'),$$

where $\alpha, \theta, x_t, y_t \in \mathbb{R}$, $(x,y) \in S \subseteq \mathbb{R}^2$, $(x',y') \in S' \subseteq \mathbb{R}^2$.

If $\alpha$ is restricted to being positive and finite, $\theta$ to being greater than zero and less than two pi, and $x_t, y_t$ to being finite, then all three mappings are not only onto but one-to-one [10].

Using the definitions for the transforms of scale, rotation and translation the following theorem can be formulated.

**Theorem:** the set of critical points $C$ obtained for an arbitrary shape $S$ which has undergone one of the transformations $S_{\alpha}$, $R_{\theta}$ or $T$ is identical to the set of critical points for the untransformed shape, subjected to the same transform. This can be stated mathematically as:

$$\Psi \left( S_{\alpha} \left( S \right) \right) = S_{\alpha} \left( \Psi \left( S \right) \right),$$

$$\Psi \left( R_{\theta} \left( S \right) \right) = R_{\theta} \left( \Psi \left( S \right) \right),$$

$$\Psi \left( T \left( S \right) \right) = T \left( \Psi \left( S \right) \right).$$

The proof of this theorem appears in the appendix.
In general compositions of $S_a(\bullet)$, $R_\theta(\bullet)$ and $T(\bullet)$ are not commutative because of the nonlinearity of $T(\bullet)$; therefore their order must be preserved. However, combinations involving only $S_a(\bullet)$ and $R_\theta(\bullet)$ are commutative [10].

The next question to be addressed involves the invariance of critical point determination when only partial shape information is available. This problem can be divided into two simpler and similar type problems, namely, truncated shape representation and composite shape representation. In the first case, it is desirable to determine a source shape $S$ from a given shape $S'$ where $S'CS$ and $B'\nabla B=\emptyset$. In the second case, it is desirable to determine two or more source shapes $S_i$ from $S'$ where $S'=\bigcup_{i=1}^{n}S_i$ and $B'\nabla B=\emptyset$ for all $i$.

In order to accomplish either of the above task it is necessary for a subset of critical points from each source shape to remain unaltered in $S'$. Since the critical points are fundamentally linked to the concavity tree of the source shape, this is equivalent to requiring that a portion of the concavity tree remain unaltered.

In the case of truncation, the portion of the concavity tree which is distorted can be identified by examining the convex hulls of the whole and partial shapes, or $S$ and $S'$ respectively. Those subshapes or branches of the concavity tree which do not intersect a portion of $(S^C-S'^C)$ remain unaltered. An example of this relationship is illustrated in figure 4 for an arbitrary shape $S$ and subshape $S'$. From this it can be seen that each branch of the concavity tree acts as a generating node for all subsequent branches in that portion of the tree. Any node which remains unaltered will generate sub-level branches identical to those of the original tree. A comparison between the branches of the original shape $S$ from figure 4 and its corresponding truncated shape $S'$ is shown in figure 5. The branches inscribed by squares are...
identical in both trees.

In the case of composite shapes, the effects are more complicated. Not only are branches of the original tree truncated, but new branches may be created and others augmented. This occurs because 1) the union of the convex hulls of the combining shape is not necessarily convex, thereby creating new concavities and 2) concavities from one shape may overlay interior areas of another. These possibilities are illustrated in figure 6 for a composition of the shape used previously in figure 4.

In general any branch of the concavity tree which intersects a portion of the overlap region or intersects a concavity which includes the difference between $S^c$ and $\bigcup_i S^c_i$ will be distorted. In certain cases it is possible for certain branches, a subset of the distorted portion of the tree, to remain unaltered. However the conditions under which this occurs are complex and specific to the particular shape.

In the next section, the gap between the theoretical shape description developed in the previous section and a discrete implementation for digitized shapes is bridged. The effects of finite boundary representation, quantization noise and additive observation noise are also considered.

III CRITICAL POINTS FROM REAL DATA

The definition of critical points advanced in the previous section can be restated as follows: critical points are the endpoints of the line segments formed by the intersection of a shape’s boundary $B$ with the branches (subshapes) of its concavity tree. If the shape boundary is continuous and can be observed without error, then these critical points can be exactly determined for all combinations of scaling, rotation and translation. However, in the discrete case the shape is usually represented by a finite, integer-valued sequence. This sequence is almost always in
the form of a chain code or a thinned boundary code [12]. In this work it is assumed that the sampled boundary has been properly preprocessed so that it meets the discrete definitions of being closed and single-connected. These definitions are described in [22] along with various techniques for processing images which guarantee that these conditions are met.

With these requirements in mind, let the processed shape be represented by the sequence \( \{x_i, y_i\}_{i=1}^N \), which corresponds to the quantized estimate of the shape boundary. The coordinates of the sequence, \( x_i \) and \( y_i \), should be eight-connected and integer-valued. Because only boundary information is available the previous definition for critical points will have to be reinterpreted.

The general procedure for extracting critical points is illustrated by the flow chart in figure 7. At the front end of the system is a spatial filter which is chosen to minimize the effects of the quantization and observation noises. At the next stage, the convex hull information about the shape is obtained. A decision process is then required to identify and confirm critical points. Depending on the output of the decision stage, the shape is either segmented into the next level of branches of its concavity tree or the procedure is terminated. This procedure is recursive, with each pass around the loop corresponding to the processing of one level of the concavity tree.

The purpose of the spatial filter is to reduce the effects of the quantization noise and the additive noise in the boundary sequence. The type of filter chosen should be optimized for the statistics of the noise processes and the source shapes [1]. In general these statistics depend on the imaging equipment and the set of source shapes, and cannot be determined globally. In fact, they are most likely nonstationary random processes. On the other hand, if the filter is made robust, good results can still be obtained. Therefore, an averaging filter is chosen.

The output of the filter will be a real-valued sequence \( \{x_i, y_i\}_{i=1}^N \), with the same
number of elements as the input sequence. By increasing the length of the filter window the effects of the noise can be made arbitrarily small. However, the smoothing effects may cause smaller features of the shape to be eliminated. In many cases, this may be considered advantageous because a simultaneous reduction in feature processing and increased noise immunity is obtained.

The next step is to find those points from the sequence \( \{x_i, y_i\}_{i=1}^{N} \) which are elements of the boundary of the convex hull of the digitized version of \( S \). This boundary is equivalent to the Minimum Perimeter Convex Polygon (MPCP) enclosing \( S \). It can be calculated using a modified version of the routine described by Freeman and Shapira [7]. This modified version is adapted to process real data instead of integer data. It requires a longer running time but still iterates to a final solution in all cases. The actual convex hull is not required, only the perpendicular distance from each point of the shape boundary to the MPCP. This distance is referred to as the difference sequence \( \{d_i\}_{i=1}^{N} \).

The procedure for determining the critical points from the difference sequence is now described. As a first step, an interpretation of the definition of critical points must be applied in the context of the difference sequence. Since the difference sequence measures the distance between the boundary of \( S \) and the boundary of \( S^c \), it can be used to find the intersection defined in (18). This intersection for the discrete case corresponds to those points of \( \{d_i\}_{i=1}^{N} \) which equal zero. The endpoints (critical points) of the associated line segments are just those points where the difference sequence goes from a zero value to a nonzero value. If the shape boundary is observable without error, then the critical points can be determined exactly; however, this is not the case when applied to any discrete data. Therefore, the
critical points will be determined by finding certain transition regions in the
difference sequence. This will compensate for most of the errors due to observation
noise as well as errors incurred by the resampling of shapes which have undergone
one or more of the transformations discussed previously. Of these three, only the
transformation of scale seriously affects the difference sequence (the maximum
error resulting from the other two transformations is limited to $2/\sqrt{2}$ times the
quantization grid size). Since the spatial energy contained in the shape is directly
proportional to the area of the quantization grid which it covers, any reduction in
scale by a factor $\alpha$ results in a simultaneous reduction in signal to noise ratio of $\alpha^2$.
Because of this, certain compromises must be made in the performance of the
critical point extraction algorithm, specifically, it will be designed to operate over a
limited range of scale factors. To be more precise, the resulting procedure tries to
predict the behavior of the critical points over the design range of scale factors. This
reduces to a more basic problem of distinguishing concavities formed by the noise
processes from those concavities due to the shape structure. The exact procedure for
this is described next.

The input shape for extracting critical points for the shape dictionary is evaluated
at $\alpha$ equal to one (the maximum scale factor). This corresponds to the highest signal
to noise ratio expected for the shape. The first step is to isolate a set of potential
critical points. This is accomplished by comparing the difference sequence to a
threshold, $r_1$. When the absolute value of the difference sequence is greater than $r_1$
a concavity is formed. The points inside the interval $[0, r_1]$ closest to the position
where the $r_1$ threshold is crossed are considered as potential critical points and
added to the critical point list. These points are in direct correspondence with the
endpoints of the line segments in (18). They determine certain subsequences of
$(d_i^N)_{i=1}^N$ called concavity sequences. The concavity sequences $(d_i^c)$ are the sequential
portions of $(d_i^N)_{i=1}^N$ which are greater than $r_1$. Associated with each concavity
sequence are a pair of critical points, one at each end. The first threshold simply identifies possible critical points. A second threshold is used to evaluate the most probable underlying cause of a concavity.

The second threshold $r_2$, which is greater than $r_1$, is used to eliminate critical points associated with small concavities from the list. It also helps remove most concavities which are formed strictly from observation noise. For each sequence of points within a concavity, $\{d_i^c\}$, the maximum absolute value for the $d_i^c$'s are found. If this value, called $d_{max}^c$, is less than then the critical points associated with the concavity in question are eliminated from the critical points list. Since it is impossible to distinguish small shape concavities from those produced by noise with just one sample, the alternative is to eliminate them all. The reasoning being that the total number of shape concavities eliminated will be small compared to the number of noise produced concavities. This can be justified theoretically by assuming that the average maximum depth of the actual shape concavities is much greater than the maximum depth of the noise concavities. This is illustrated in figure 8 for a generalized representative shape. This type of decision is analogous to the classical maximum-likelihood decision criteria [16]. Note that the depth of the true concavities can be considered as the signal energy, and the depth of the noise related concavities as the noise energy. From figure 8 it can be seen that for appropriate thresholds $r_1$ and $r_2$ most of the effects of the noise can be eliminated, provided the signal to noise ratio is sufficient. The matching algorithm, to be described later in this chapter, is designed to be very tolerant of missed critical points as long as they are listed in the dictionary. However, it is susceptible to error if unexpected critical points are found in the problem text. Therefore, the threshold $r_2$ is biased by an amount $r_b$ when the shape dictionary is being prepared. The direction of the bias is in the direction of higher false alarm rates for concavities due to noise.

Now that the majority of potential critical points have been isolated, it is
necessary to find a method of predicting their behavior in the problem text over the
design range of scale factors. The effects of changes in $\alpha$ on the $|d_i^c|$'s falls into two
categories: 1) concavities found at the maximum value of $\alpha$ may collapse below the
$r_2$ threshold and their associated critical points eliminated from the problem text;
2) segments of the difference sequence, which are outside the interval $[0,r_2]$ for
large $\alpha$, may cross the $r_2$ threshold and create new critical points. Of these two
possibilities, the first case has been observed to happen most frequently. This can
be explained as follows. Because the depth of a concavity is determined from two
extremal points of opposite direction, namely, the outer most points on the shape
boundary contained in the convex hull and the inner most shape points corresponding
to the maximum depth of a concavity, the smoothing effect of the spatial filter
converges these two points approximately twice as fast as distances based on only
one of the extremal points, i.e. case two.

In order to predict the occurrence of these two changes, two additional thresholds
are introduced. Their purpose is to assign a measure to regions of the difference
sequence which may cause one of the above changes. This measure, called the
confidence number, will be assigned individually to each critical point in the shape
dictionary. The confidence number indicates the percentage of the scale factor range
in which a particular critical point is detectable as well as the order in which it
enters (or leaves) the problem text.

The first of these two thresholds, $r_3$, is used to identify candidate critical points
which may be detected in the problem text for smaller values of $\alpha$. If any sequential
portion of $|d_i^c|$ for any concavity has the property that

$$r_2 < |d_i^c| \leq r_3$$

for all $d_i^c$ in that group, then a potential critical point is indicated. The two points
where the sequence \( \{d^c_i\} \) indicated by (25) crosses into the interval \([r_2, r_3]\) are added to the shape dictionary and assigned a confidence number, \( #_c \), given by

\[
#_c = \frac{|r_3 - d^c_{\min}|}{|r_2 - r_3|}
\]

where \( d^c_{\min} \) is the minimum absolute value of the difference sequence in region \([r_2, r_3]\). An example of this situation is illustrated in part a of figure 9.

The second new threshold, \( r_4 \), is used to measure the proportion of scale factors at which a particular concavity, and its associated critical points, can be detected. For each concavity sequence with \( d^c_{\max} \) greater than \( r_2 \) assign a confidence number according to the following:

\[
\text{if } d^c_{\max} > r_4, \quad \text{then } \quad #_c = 1;
\]

\[
\text{else } \quad #_c = \frac{|d^c_{\max} - r_2|}{|r_4 - r_3|}
\]

This situation is illustrated in part b of figure 9. Note that \( r_4 \) is generally twice as large as \( r_3 \) and that \( r_3 > r_2 > r_1 \).

At this point the algorithm for extracting critical points and assignment of confidence numbers for a single level of the concavity tree has been described. To extend the procedure to the rest of the concavity tree, the recursive nature of the algorithm must be defined. Since the concavity sequences correspond to the next level of concavity tree branches, they can be processed individually in the same manner as the original shape. The only modification required is the reversal of the value for \( s \) in (9) at each new level. New critical points are added to the shape dictionary as previously described. Confidence numbers are assigned using the same criteria as before with the following exception: concavities which have confidence numbers less
than 1, have all sublevel critical points assigned a confidence number calculated as a product of the current level confidence numbers with the values determined according to (26) and (27). The process is propagated through the concavity tree until no new points are found. Figure 10 and 11 show the difference sequence and critical points respectively from an arbitrary shape for levels 1, 2 and 3 of the concavity tree respectively. The resulting dictionary page contains the following information extracted sequentially from the shape boundary for each critical point: 1) its x and y coordinate values, 2) its confidence number, 3) the level of the concavity tree at which it was found.

Shape Dictionary Calculation for 3-Dimensional Objects

Consider an arbitrary, rigid object as shown in figure 12, where $\theta_x$, $\theta_y$ and $\theta_z$ describe angles of rotation in the three axes of the rectangular, object-centered coordinate system. The z-axis of the object-centered coordinate system is restricted to parallel alignments with the z'-axis of the viewer-centered coordinate system. This restriction results in the following interpretations for motion attributed to the object. The imaging plane shown in figure 12 is the space in which the object is projected.

1) Translation of the object (or the object-centered coordinate system) with respect to the z'-axis results in a scale change in the imaging plane.

2) Translation of the object (or object-centered coordinate system) along either or both of the x' and y' axes results in a translation in the imaging plane.

3) Rotation of the object by an angle $\theta_z$ about the z-axis of the object-centered coordinate system results in a rotation by angle $\theta_z$ in the imaging plane.

4) Rotation by the angles $\theta_x$ and $\theta_y$ in the x and y axis of the object-centered coordinate system results in a new projection of the 3-dimensional object in...
The first three transformations are the same transformations which have been shown not to effect the critical point/feature vector formation procedure. Therefore, only the fourth transformation will necessitate a modification in the procedure previously described for 2-dimensional shape recognition.

The necessary modifications will result in an expanded shape dictionary. Now instead of one set of critical points for each shape, there will be several sets; each representing a different rotation about object-centered axes $x$ and $y$. Therefore the following procedure is suggested for assembling the 3-dimensional shape dictionary.

1) With the object at its minimum distance from the $z'$ plane (maximum scale factor) obtain a projection of the object in the viewer imaging plane with $\theta_x$ and $\theta_y$ at zero degrees.

2) Next, obtain the outer contour of the projected shape and extract its critical points using the procedure detailed earlier. Store as the first dictionary page for this shape.

3) Increment $\theta_x$ by $\theta_{inc}$ (a divisor of $180^\circ$) and repeat steps 2 and 3 to obtain the next page in the dictionary for this shape. Continue incrementing $\theta_x$ until a rotation of $180^\circ$ has been reached.

4) Reset $\theta_x$ and increment $\theta_y$ by $\theta_{inc}$. Keeping $\theta_y$ fixed at the new value, repeat steps 1 thru 3 until $\theta_y$ has been incremented to $180^\circ$.

The shape dictionary will now contain $(180^\circ / \theta_{inc})^2$ views of the 3-dimensional object. The other views (for $\theta_x$ and $\theta_y$ between $180^\circ$ and $360^\circ$) are mirror images of those already calculated, and may be extracted from the existing dictionary by processing the critical points in reverse order.
The choice of \( \theta_{\text{inc}} \) directly affects the accuracy and completeness with which the 3-dimensional object is described as well as the amount of memory needed. As \( \theta_{\text{inc}} \) is reduced the accuracy of the description is increased along with the amount of memory required to store the dictionary. With regard to the accuracy of the 3-dimensional shape description, an unintended but beneficial side effect is obtained from the extraction of the critical point's using the concavity tree. As the 3-dimensional object is rotated, various sections of the object become visible. The projections to these sections onto the imaging plane manifest themselves as new concavities in the corresponding 2-dimensional silhouette. Depending on the direction of rotation, these new concavities are either entering or leaving the concavity tree. Since this is the essence of the information measured by the confidence number, additional accuracy is gained. This is especially useful when the actual angle of rotation for the problem text is not a multiple of \( \theta_{\text{inc}} \).

**Critical Point Determination for the Problem Text**

When the boundary of an unknown shape is obtained a critical point list (problem text) must be configured for it. The procedure by which this is done is quite similar to the one described in the previous section; however, no confidence number is calculated. This is due to the fact that an exact value of the scale factor \( \alpha \) is unknown. As a consequence only those portions of the difference sequence partitioned by the thresholds \( r_1 \) and \( r_2 \) are used to determine concavities. These concavities may be formed in three ways which are illustrated in figure 13, parts a, b and c. In the first case the \( r_1 \) and \( r_2 \) thresholds are both crossed in forming a concavity; for this case, the critical points for this concavity are chosen as the points nearest to threshold \( r_1 \) inside the interval \([0,r_1]\). In the second case only the \( r_2 \) threshold is crossed; therefore, the nearest points to \( r_2 \) inside the interval \([0,r_2]\) are chosen as
critical. In the last case, only threshold $r_1$ is crossed; therefore, no critical points are chosen since the concavity is considered too small to register. As a final result the problem text contains the following information extracted sequentially from the unknown shape's boundary for each critical point: 1) its $x$ and $y$ coordinate values, 2) the level of the concavity tree at which it was detected.

IV COGNITIVE STAGE

Once a problem text has been defined it must be matched against all the dictionary pages (shapes). Since the algorithm is required to recognize partial shapes as a part of a larger shape, features must be formed and matched accordingly. The basis for comparison is the shape or feature vector described previously. It consists of measurements made from 1 arbitrary critical points selected from the shape dictionary or problem text. As a first step, the process by which each feature vector is selected must be made non-arbitrary. Four general conditions are suggested for choosing the individual critical points; they are as follows:

1) The set of all feature vectors should give near equal representation to all parts of the shape.
2) The total number of feature vectors is kept as small as possible.
3) Each feature vector represents a feature likely to appear under certain conditions of translation, scale or rotation.
4) Each feature represents a localized characteristic of the shape.

The conditions for the representative feature set can be met by combining the concavity tree information with the confidence number information contained in the shape dictionary. The procedure for this is as follows. Let each pair of critical
points associated with a given concavity act as seed points for a feature vector. Choose the two most reliable neighboring critical points to each side of the seed points to complete the sixtuple. This method guarantees at least one feature vector for each branch of the concavity tree. Furthermore, it restricts each feature to three adjacent concavities for a given scale factor. Finally, a linear growth rate is achieved - if there are N critical points then there will be N/2 feature vectors. The method for selecting the most reliable, neighboring critical points is described next.

Recall that the confidence number not only indicates the percentage of scale factors for which a critical point is detectable but also the order in which they are detected. For example, if \( #a \) and \( #b \) represent the confidence numbers for two critical point pairs - a and b respectively, and \( #a > #b \); then if pair b is detected, pair a should also be detected. Therefore, for a given critical point pair with confidence number \( #c \), choose the two nearest neighboring critical points which have confidence numbers greater than or equal to \( #c \). When considering the problem text, information concerning scaling is unavailable; therefore, the simplest scheme possible is derived, i.e, each consecutive grouping of six critical points forms a feature vector.

Let the feature vectors from the shape dictionary be denoted by \( d_{k}^{m} \) where k indicates the page of the dictionary and m the mth feature vector within page k. Similarly denote the nth feature vector of the problem text by \( d_{n}^{p} \).

Step 1) Associated with each feature vector of the dictionary and the problem text is a sixtuple called the level code vector. Each component of the level code vector is integer valued and indicates the level of the concavity tree at which the corresponding critical point was detected. The symbol for the level code vector associated with feature vector \( d_{k}^{m} \) \( (n, Z) \) is \( d_{k}^{m} \) \( (l, Z) \). For feature vectors to match it is necessary that their level code vectors also match. Therefore, in the first stage of the procedure all of the level code vectors from the shape dictionary page and the problem text are cross matched. The distance defined by
\[ d_1(m,n) = \sum_{i=1}^{I} \left| \frac{d_k}{m^i_1 - n^i_1} \right| \]

is the basis for each level code vector comparison. A value of zero for \( d_1(m,n) \) is a necessary condition for two feature vectors to match. This distance is calculated for all \( m \in M_k \) and all \( n \in N \). This cross referenced set is referred to as \( NXM_k \). Each pair \((m,n)\) such that \( d_1(m,n) = 0 \), is added to the reduced feature set called \( NXM_k' \). Since \( d_1(m,n) \) requires only \((2I-1)\) integer operations, compared to \((2I(I-1))\) real valued operations for a complete feature vector comparison, computational overhead is significantly reduced.

Step 2) For each pair of features associated with the elements of the reduced set \( NXM_k \), calculate a pair of feature vectors using the definitions methods outlined in chapter III. Next evaluate the distance \( d_1(m,n) \) defined by

\[ d_1(m,n) = \left| \left| \frac{d_k}{m^1 z_1} \right| - \left| \left| n^1 z_1 \right| \right| \]  

for each component of the feature vector pair. In the noiseless case the distance \( d_1(m,n) \) is zero for all components of the feature vectors \( m \) and \( n \) if they match. Since this is never the case a finite positive distance \( r_m \) must be allowed when evaluating \((29)\). Therefore, if \( d_1(m,n) < r_m \) for \( i = 1 \) to \( I \), then the feature vectors are said to match. The set of matching feature vector pairs from this stage form the new set \( NXM_k'' \). It should be mentioned at this point that a zero value for all components \( i \) with respect to the distance \( d_1(m,n) \) is not a sufficient condition for a match, since only the magnitude information from each component is considered in \((29)\). On the other hand the angle information is examined in the next phase of the algorithm and errors rarely occur due to this in the final evaluation.

Step 3) Encoded within each group of matching feature vectors is information about
the scale and rotation parameters of the problem text, $\alpha$ and $\theta$ respectively. If a noiseless case is assumed, then the $\alpha$ parameter is equal to the ratio of the two size variables and the $\theta$ parameter is equal to the angle between the feature vectors. Since perfect knowledge of the feature vectors is never obtained the values of $\alpha$ and $\theta$ can only be estimated. The equations for the estimates of $\alpha$ ($\hat{\alpha}$) and $\theta$ ($\hat{\theta}$) are given by (30) and (31) respectively.

$$\hat{\theta}_{m,n} = \sum_{i=1}^{6} \frac{d_k}{||m_i||} \theta_i$$

where

$$\theta_i = \cos^{-1}\left(\frac{d_k \cos\theta_0}{||m_i|| \cdot ||n_i||}\right)$$

and

$$\hat{\alpha}_{m,n} = \frac{\sum_{i=1}^{6} d_k p_i}{\sum_{i=1}^{6} m_i n_i}$$

These estimates are obtained for each pair of matching feature vectors in the set $N_{X \times M}^k$. Note that the estimate in 30 is weighted by the magnitude of the individual components so that large errors in small components do not overly bias the estimate.

Step 4) By this point there are only a few possible matches remaining, the problem now is to pick the best match of the remaining few, or to reject all of them if they are false matches. Since feature vectors from the same shape should all have the same scale factor and angle of rotation, clusters in the estimates of $\alpha$ and $\theta$ should be formed. The following procedure is used to define such a cluster. Let $p$ be the index on set $N_{X \times M}^k$ with values from 1 to $nm$ and form the vectors $[\hat{\alpha}_1, ..., \hat{\alpha}_p, ..., \hat{\alpha}_{nm}]^T$ and $[\hat{\theta}_1, ..., \hat{\theta}_p, ..., \hat{\theta}_{nm}]^T$ from the estimates in step 3. Now
calculate the two circulant difference matrices $T_\alpha$ and $T_\theta$ and normalize them by $r_\alpha$ and $r_\theta$ so that

$$T_\alpha = \begin{bmatrix} a_1 - a_1 & \cdots & a_1 - a_{nm} \\ \vdots & \ddots & \vdots \\ a_{nm} - a_1 & \cdots & a_{nm} - a_{nm} \end{bmatrix}$$

and

$$T_\theta = \begin{bmatrix} \hat{\theta}_1 - \hat{\theta}_1 & \cdots & \hat{\theta}_1 - \hat{\theta}_{nm} \\ \vdots & \ddots & \vdots \\ \hat{\theta}_{nm} - \hat{\theta}_1 & \cdots & \hat{\theta}_{nm} - \hat{\theta}_{nm} \end{bmatrix}$$

The values $r_\alpha$ and $r_\theta$ represent the maximum allowable intra-cluster distances for the estimates of parameters $\alpha$ and $\theta$ respectively. Next calculate $T_D$ the total difference matrix between all estimates as

$$T_D = \sqrt{\left(T_\alpha^2 + T_\theta^2\right)}$$

where each operation is performed element wise. $T_D$ will be an $nm \times nm$ difference matrix normalized by $\sqrt{\frac{r_\alpha^2 + r_\theta^2}{r_\alpha + r_\theta}}$. To find the clusters find the matrix formed by setting all elements of $T_D > 1$ to zero and all others to 1. Now each row (or column) contains a 1 for each estimate that is clustered with the indexing estimate. In general there should be only one cluster if there is a match; however, the above procedure may also produce several subclusters of the larger cluster. In order to pick the best cluster, the number of terms and the strength of the
individual matches between each feature vector should be considered. The following
decision scheme is chosen to select the final cluster.

1) Total the number of members in each cluster, call this number GCNT
2) Multiply each group member by the weighting factor

\[ 1 - \sum_{i=1}^{1} d_i(m,n) \]

this indicates the degree of match between each feature vector in the
cluster. Sum this number and call it GSUM

3) Add the value \( w \cdot (GCNT - 1) \) to GSUM and find the average value of this
total for each group; \( GAVG = \frac{(w \cdot (GCNT - 1) + GSUM)/GCNT} \)

4) Pick the cluster for which \( GAVG \) is maximum.

To get the final estimates of \( \alpha \) and \( \theta \), find the average of the individual \( \hat{\alpha}_i \) and \( \hat{\theta}_i \) for
the chosen cluster. If there is no cluster the above procedure reduces to picking the
individual feature vector with the highest degree of match.

Unlike \( \alpha \) and \( \theta \) the translation of a whole shape is not equivalent to the same
translation in each of its feature vectors (recall that translation is not a linear
operation). Therefore, scale and rotation information is required to estimate \( z_t \) (the
translation vector between the centroid of the dictionary shape and the centroid of
the problem shape which have been matched). To estimate \( z_t \) the matched shape
dictionary is scaled and rotated with respect to its centroid using the final estimates
obtained for \( \alpha \) and \( \theta \). The rotated and scaled versions of the dictionary coordinates
are calculated as

\[
\hat{d}_{k,m}^{x'} = \hat{x}_k + \alpha \left[ \cos(\hat{\theta}) \left( \frac{d_k}{m} \cdot \hat{x}_k \right) + \sin(\hat{\theta}) \left( \frac{d_k}{m} \cdot \hat{y}_k \right) \right]
\]
\[
\begin{align*}
\hat{d}_{k,x} &= \hat{y}_k + \alpha \frac{d_{k,x}}{m} \cos(\theta) \left( \frac{d_{k,y}}{m} - \hat{y}_k \right) + \alpha \frac{d_{k,y}}{m} \sin(\theta) \left( \frac{d_{k,x}}{m} - \hat{x}_k \right) \\
\end{align*}
\] 35

where \( m = 1 \) to \( M_k \) and \( (x_k, y_k) \) is the centroid of the dictionary. Note if the dictionary is prepared so that its centroid is \((0,0)\) then (35) can be reduced to

\[
\begin{align*}
\hat{d}_{k,x} &= \alpha \cos(\theta) \left( \frac{d_{k,x}}{m} \right) + \alpha \sin(\theta) \left( \frac{d_{k,y}}{m} \right) \\
\hat{d}_{k,y} &= \alpha \cos(\theta) \left( \frac{d_{k,y}}{m} \right) - \alpha \sin(\theta) \left( \frac{d_{k,x}}{m} \right) \\
\end{align*}
\] 36

Since translations from the same shape should be equal (except for certain cases of symmetry) a clustering of translation parameters, similar to that just described for \( \alpha \) and \( \theta \), is expected for the estimates of \( x_t \) and \( y_t \). Therefore, an additional step can be implemented to further reduce the possibility of false alarms and large errors in the estimate of \( z_t \).

Consider the final cluster from which \( \alpha \) and \( \theta \) were obtained. Calculate the values \( c_{x_t} \) and \( c_{y_t} \), the translation coordinates between each matched feature vector from the cluster, where

\[
\begin{align*}
-\frac{1}{6} \sum_{i=1}^{6} c_{x_i} &= \frac{1}{6} \sum_{i=1}^{6} c_{y_i} \\
\end{align*}
\] 37
where

\[ x'_1 = c x'_1 - \hat{x}'_1 \]

\[ y'_1 = c y'_1 - \hat{y}'_1 \]

with

\( c \) = the index over the cluster obtained from the estimates of \( \alpha \) and \( \theta \)  
\((c=1 \text{ to } C)\),

\( i \) = the index on the individual components of each feature vector from the cluster,

\( x' \) = the \( x \) coordinate from the transformed dictionary page,

\( y' \) = the \( y \) coordinate from the transformed dictionary page.

Next form the two vectors \([x'_1, \ldots, x'_t, \ldots, x'_C]^T\) and \([y'_1, \ldots, y'_t, \ldots, y'_C]^T\) from the values obtained from (37). Then calculate the two circulant difference matrices \( T_x \) and \( T_y \) normalized by \( \tau_x \) and \( \tau_y \) (the maximum allowable intra-cluster distance for \( x_t \) and \( y_t \)) as

\[
T_x = \begin{bmatrix}
1 x'_t - x'_1 & \ldots & 1 x'_t - C x'_t \\
\vdots & \ddots & \vdots \\
1 x'_t - x'_1 & \ldots & 1 x'_t - C x'_t \\
\end{bmatrix}
\]

\[
T_y = \begin{bmatrix}
1 y'_t - y'_1 & \ldots & 1 y'_t - C y'_t \\
\vdots & \ddots & \vdots \\
1 y'_t - y'_1 & \ldots & 1 y'_t - C y'_t \\
\end{bmatrix}
\]

Next calculate \( T_z \) the total normalized translation difference, as
where each operation is performed element wise. $T_z$ can be checked for clustering in the same manner used previously with $\hat{\alpha}$ and $\hat{\theta}$. In general all elements of $T_z$ are expected to cluster; however, there is a small probability (dependent on the values of $r_x$ and $r_y$) that they will not. In this case, the non-clustering terms are discarded and the final estimates of $x_t$ and $y_t$ are obtained by averaging over the remaining cluster elements.

V RESULTS

A 3-dimensional shape dictionary was constructed. It consisted of two airplanes, an F-15 and an F-4. A value of 15 degrees was chosen for $\theta_{\text{inc}}$ and a total of 20 views generated for each plane. The range of views covered $\pm 30$ degrees pitch and 0 to -45 degrees roll. The boundary curves were obtained by thresholding a 512X512 pixel image of each view to obtain a binary representation. A contour was then obtained from this image and processed for critical points. The contours obtained for each plane are shown in figure 14.

A value of 13 was chosen for the window length of the averaging filter used in the first stage of the critical point extraction routine. The values used for the segmenting thresholds were $r_1=-.3$ and $r_2=-1.2$. The values of the thresholds used to determine the confidence numbers were $r_3=-2.4$ and $r_4=-4.8$ respectively. This corresponds to a design range of .5 to 1 for the scale factor $\alpha$. A bias value of $\tau_b$ equal to .48 was calculated based on a maximum deviation due to grid rotation of $(\sqrt{2}-1)r_2$. A maximum normalized distance of $r_m=.04$ is allowed between feature vector components. Maximum cluster distances of $r_a=\pm .075$ and $r_\theta=\pm 5$ degrees were
selected for the scale, rotation cluster portion of the algorithm. Translation parameter clustering was implemented with a maximum scattering of ±10 pixels for both $r_x$ and $r_y$. A cluster weight of $r_w = .1$ was chosen for the cluster decision procedure.

The problem texts were then extracted from the scene shown in figure 15. This scene contains three planes, an F15, an F-4 and a HARRIER, placed at arbitrary orientations within the range covered by the shape dictionary. The HARRIER was used as a control to check the algorithm's ability to discriminate non-dictionary shapes from dictionary shapes. The video camera has been moved away from the planes to provide a scale change and adverse lighting has been employed to add uncertainty in the determination of the planes' boundaries. The three problem texts shown in figure 16 were extracted from the scene using binary thresholding. They were then matched with the shape dictionary. From figure 16 it can be seen that only partial shape information is recovered for each plane (the shadows are interpreted as part of the object information).

Each problem text was cross-matched with the two objects in the shape dictionary. The results of the match are demonstrated in figure 17. The white contours superimposed over the original scene indicate the view from the dictionary matched to each problem text with its estimated scale, rotation and translation parameters. The first problem text is identified as an F-15 at -15 degrees pitch and -15 degrees roll. The second problem text was identified as an F-4 at 0 degrees pitch and -30 degrees roll. The third problem text was not recognized as a member of the shape dictionary. The procedure generated no false alarms.

VI DISCUSSION

If additional identification power is required a second stage can be implemented to
eliminate portions of any composite shape problem text as they are identified, thereby reducing the composite problem text to a truncated problem text which has a higher recognition rate. Also false alarm rates can be significantly reduced by implementing a similar procedure to compare boundary points once a possible match has been indicated. It is also possible to extend the feature set chosen for each dictionary shape thus reducing the probability of a miss. Unfortunately this results in an increase in match times required for each dictionary shape. This increase in time will be linearly related to the increase in the feature vector set. Miss rates may also be reduced by relaxing the maximum distance allowed between feature vector components. This is recommended only if the previously discussed verification procedure is implemented because relaxation also increases the rate of false alarms.

If a sequence of dynamic images (scenes with objects in motion) is being analyzed, it is believed that the overall recognition rates will improve over the entire course of the sequence. This is because the amount of occlusion will not remain constant during the entire sequence, thus revealing more object information in the totality of the sequence than in any one frame.

VII CONCLUSIONS

The definition of critical points using the concavity tree structure and the corresponding identification algorithm described in this paper have been shown to offer the following features and advantages when the identification of 3-dimensional shapes in 2-dimensional scenes is considered. The theoretical definition of critical points using the concavity tree structure of a shape is invariant to transforms of scale, rotation and translation within the 2-dimensional scene. In the actual implementation, invariance to the latter two transforms is maintained and
invariance over a specific range of scale factors is obtained. The critical point
definition is reasonably stable under conditions of shape occlusion and partial
framing, such that partial shape information is usually sufficient to identify a shape.
The algorithm requires only one pass per concavity tree branch to extract the
critical points. These critical points are the same regardless of starting point. The
critical points are concentrated in areas of high shape dynamics, i.e. the highest
areas of shape quantification occur in those regions of the shape of highest
qualitative interest. The algorithm is considered robust in the sense that, the more
complex a shape becomes the higher the probability of identifying it in a scene. This
is due to the fact that more critical points are allocated to the more complex
shapes. The algorithm is flexible in the sense that it can be modified to accommodate
restrictions on memory or computation. Also as a byproduct of the matching
procedure, estimates of the scale, rotation and translation parameters of an
identified shape are automatically generated.
The proof of the theorem in section III follows. It has already been established that the operation \( \Psi(\cdot) \) is a point set relationship between a shape (subshape) and its concavity tree, or to be more exact a shape (subshape) and its convex hull. Therefore it is sufficient to show that this relationship is maintained under the three transformations. The convex hull \( S_c \) of a shape \( S \) is defined mathematically as the set of all points \( r \) such that

\[
r = \lambda p + (1 - \lambda) q ; \quad 0 < \lambda < 1 \quad \text{a1}
\]

where \( p, q \) are points in \( S \). This includes all points on all line segments joining every pair of points in \( S \). For points \( r, p, q \) in \( \mathbb{R}^2 \) (a1) may be rewritten as

\[
(r_x, r_y) = \lambda (p_x, p_y) + (1-\lambda) (q_x, q_y) , \quad \text{a2}
\]

where \( r_x = \lambda p_x + (1-\lambda) q_x ; \ r_y = \lambda p_y + (1-\lambda) q_y ; \ 0 < \lambda < 1 \). The effect of each transformation is to map individual points of \( S \) into \( S' \), therefore the definition of \( S'_c \) is simply the above definition with \( p, q \) replaced by \( p', q' \). Consider first the convex hull of a shape under the transformation \( S_d(\cdot) \). It becomes the set of all points \( r' \) such that

\[
r' = \lambda p' + (1 - \lambda) q' ; \quad 0 < \lambda < 1
\]

or

\[
(r'_x, r'_y) = \lambda (\alpha p_x, \alpha p_y) + (1-\lambda) (\alpha q_x, \alpha q_y) , \quad \text{a3}
\]

where

\[
\begin{align*}
r'_x &= \lambda \alpha p_x + (1-\lambda) \alpha q_x ; \\
r'_y &= \lambda \alpha p_y + (1-\lambda) \alpha q_y . \quad \text{a4}
\end{align*}
\]

the terms in (a4) can be rearranged so that
\[ r'_x = \alpha \left[ \lambda p_x + (1-\lambda) q_x \right]; \]
\[ r'_y = \alpha \left[ \lambda p_y + (1-\lambda) q_y \right] \]

which is equivalent to the transformation \( S_{\alpha}(\bullet) \) on the whole line segment, or

\[ r' = S_{\alpha} \left\{ \lambda p + (1-\lambda) q \right\}. \]

Taken over the entire set \( S \), this can be rewritten as

\[ C_h(S_{\alpha}(S)) = S_{\alpha}(C_h(S)). \]

which can be interpreted as "the convex hull of the scaled shape \( S \) is equal to the scaled convex hull of \( S \). A similar argument can be used for the transformations \( R_\vartheta(\bullet) \) and \( T(\bullet) \) with (a4), (a5) and (a6) replaced by (a8), (a9), (a10) and (a11), (a12), (a13) for \( R_\vartheta(\bullet) \) and \( T(\bullet) \) respectively. Specifically,

\[ (r'_x, r'_y) = \lambda \left( p_x \cos \theta - p_y \sin \theta, p_x \sin \theta + p_y \cos \theta \right) + (1-\lambda) \left( q_x \cos \theta - q_y \sin \theta, q_x \sin \theta + q_y \cos \theta \right) \]

where,

\[ r'_x = \lambda \left( p_x \cos \theta - p_y \sin \theta \right) + (1-\lambda) \left( q_x \cos \theta - q_y \sin \theta \right); \]
\[ r'_y = \lambda \left(-p_x \sin \theta + p_y \cos \theta \right) + (1-\lambda) \left(-q_x \sin \theta + q_y \cos \theta \right) \]

which can be rewritten as,

\[ r'_x = [\lambda p_x + (1-\lambda) q_x] \cos \theta - [\lambda p_y + (1-\lambda) q_y] \sin \theta; \]
\[ r'_y = [\lambda p_x + (1-\lambda) q_x] \sin \theta + [\lambda p_y + (1-\lambda) q_y] \cos \theta \]

and finally,

\[ r' = R_\vartheta \left\{ \lambda p + (1-\lambda) q \right\}. \]
Similarly,

$$(r_x', r_y') = \lambda (p_x + x_t, p_y + y_t) + (1-\lambda) (q_x + x_t, q_y + y_t)$$

where,

$$r_x' = \lambda (p_x + x_t) + (1-\lambda) (q_x + x_t);$$
$$r_y' = \lambda (p_y + y_t) + (1-\lambda) (q_y + y_t);$$

which can be rewritten as,

$$r_x' = \lambda p_x + \lambda x_t + q_x - \lambda q_x + x_t - \lambda x_t = [\lambda p_x + (1-\lambda) q_x] + x_t$$
$$r_y' = \lambda p_y + \lambda y_t + q_y - \lambda q_y + y_t + \lambda y_t = [\lambda q_y + (1-\lambda) q_y] + y_t$$

and finally,

$$r' = T \left\{ \lambda p + (1-\lambda) q \right\}.$$  

These arguments can be summed up, as in (a7) for the scaling transform, by (a14) and (a15) for the rotation and translation transforms respectively.

$$C_h(R_\theta(S)) = R_\theta(C_h(S)).$$
$$C_h(T(S)) = T(C_h(S)).$$

It has been shown that the relationship between the convex hull of a shape with the shape itself is preserved under the defined transformations. The only other operations involved with the determination of C from S are intersections. However, since $S_{\sigma}(\bullet), R_\theta(\bullet)$ and $T(\bullet)$ are restricted to be one-to-one, all intersections are preserved by the nature of the transforms, i.e., any two points which coincide in the image of the transforms will also coincide in the preimage of transforms. Therefore the relationships in (24) are true.
BIBLIOGRAPHY


Figure 1 - Feature Vector Representation $i = 6$
Original Shape Level 1 Concavities

Level 2 Concavities Level 3 Concavities

Concavity Tree

Figure 2 - Concavity Tree for Arbitrary Shape S
Figure 3 - Boundary Segmentation at Level 1 for Arbitrary Shape S
Whole Shape S

Partial Shape S'

Convex Hull of Whole Shape S (\(S^c\))

Convex Hull of Partial Shape S' (\(S'^c\))

- indicates \(S^c - S'^c\)
- indicates unaltered subshapes
- indicates altered subshapes

Figure 4 - Concavity Tree for Partial Shape S'
Figure 5 - Concavity Tree Comparison for Whole and Partial Shapes $S$ and $S'$
Figure 6 - Regions of Distortion in Convex Hull Due to Shape Composition
Input Boundary Sequence → Boundary Filter

Calculate Convex Hull for Shape (Subshape) → Segment Shape or (Subshape) at Critical Points and Update Critical Point List

Calculate Difference Sequence for Shape (Subshape)

Check Thresholds

New Points

No New Points

Store in Shape Dictionary END

Figure 7 - Flow Diagram for Critical Point Extraction Procedure
Generalized Distribution of Concavity Depths Due to Noise

Generalized Distribution of Concavity Depths Due to Shape

Maximum Concavity Depth

Distribution

Shape Concavities missed due to thresholding

Concavities Detected Due to Noise

Maximum Concavity Depth

Thresholding

Figure 8 - Concavity Thresholding
(a) Case 1

(b) Case 2

Figure 9 - Potential Concavities
Figure 10 - Levels 2 and 3 Difference Sequences
Figure 11 - Critical Points for Levels 1, 2 and 3
Figure 12 - 3-Dimensional Object Orientation
Figure 13 - Types of Concavities
Figure 14-6 - 3-Dimensional Object Model: "F-16"
Figure 16 - Problem Texts Extracted from the Scene
Figure 17 - Scene with Identified Objects
I. Introduction

Classical quantizer design theory has focused on determining the optimal quantization mapping which minimizes the expected value of a distortion measure between the quantizer input and output. These techniques usually require a priori information concerning the statistics (probability distribution, 1st and 2nd moments, etc.) of the signal to be quantized. The need for quantizing sources with incomplete knowledge of the source statistics is well recognized, and a number of efforts [1][6][12] have been made to address the problem. Bath and VandeLinde [1] have presented the fundamental development for minimax quantization of signals with distributions from a unimodal generalized moment constrained class. The minimax quantizer is said to be
that quantizer with a maximum distortion between the quantizer input and output less than that of any other quantizer for the given distribution class.

Minimum risk quantization theory [4] has been applied to the problem of quantizing a source signal corrupted by independent additive noise (figure 1) and has shown the promise of using risk theory to determine an optimal quantization scheme. This approach also uses considerable a priori information concerning the source and noise. The general risk theory analysis however, allows the use of a number of estimation schemes to solve the corruptive noise problem. Minimax estimation can be applied to the quantization problem to produce a quantizer which guarantees a maximum risk less than that of any other quantizer. In section II the general quantization problem is stated. Section III follows with a description of the risk theory approach to quantization. Section IV develops the minimax risk quantization theory for an additive noise corrupted source with parallels drawn to the foundation provided by Bath and VandeLinde. Section V describes some simulation results followed by conclusions.

II. General Quantization

Most of the prior work in quantizer design
characterize the quantizer mapping as an estimator of the quantizer input. A decision is made regarding within which region of the domain an input sample falls, and a representative value is determined which best estimates elements of that region. Denote the quantizer domain space $\mathbf{X}$, and the finite range space $\mathbf{Y}$. Then the quantizer mapping, $y = q(x)$, produces $y \in \mathbf{Y}$, $\forall x \in \mathbf{X}$. The design of a quantizer consists of completely specifying $q(x)$ and $\mathbf{Y}$, when $\mathbf{X}$ is known. The mapping is surjective since every element in $\mathbf{Y}$ is a possible outcome of $q(x)$, $\forall x \in \mathbf{X}$. Since $\mathbf{Y}$ is finite, call $N$ the number of elements in $\mathbf{Y}$. Define an index set on the elements of $\mathbf{Y}$ so that $\mathbf{Y} = \{y_1, \cdots, y_N\}$.

The domain can be partitioned into $N$ disjoint sets, $X_i = \{x \mid q(x) = y_i, x \in \mathbf{X} \}$, $i=1, \cdots, N$, with $\bigcup_i X_i = \mathbf{X}$.

As mentioned above, it is common practice to require that a representative value $y$ estimate the sample $x$. To this end, $\mathbf{Y}$ and $q(x)$ are defined so that $y$ closely approximates $x$. This will require that each subset $X_i$ be a connected subset with $y_i \in X_i$. The use of risk theory for quantization analysis will describe the quantization process in such a fashion that the membership requirement, $y_i \in X_i$, may no longer be applicable and will not be required.
III. Risk Quantization

The problem of quantizing noisy sources, as illustrated in figure 1, requires new techniques to determine optimal quantization schemes. Here an independent identically distributed (i.i.d.) information source has been corrupted by an i.i.d. additive noise source, and it is required to quantize the resulting noisy signal. The classical quantizer design procedure yields the quantizer mapping which produces range elements that best estimate the noisy input samples -- which may produce unsatisfactory results. The problem is caused by the perception of the quantizer as an input estimator. By changing this perception, it will be shown that the problem may be solved using risk and estimation theory.

Risk quantization is based upon the state space representation of figure 2. Here the source, $\Theta$, is mapped through some probabilistic transition mechanism, $P_{X|\Theta}(x|\Theta)$, from the parameter space $\Theta$ to an observation space $\mathcal{X}$. The quantizer mapping $q(x)$ is the decision rule which determines a representative element $y$ from the decision space $\mathcal{Y}$ which estimates the source, $\Theta$, based upon the observation $x \in \mathcal{X}$. Since the quantizer maps onto the decision space $\mathcal{Y}$, and the quantizer range must consist of a finite number of elements, the elements of $\mathcal{Y}$ may be
indexed so that \( y_i, i=1, \ldots, N \), and \( \bigcup_i y_i = \mathcal{Y} \) as before.

Although the observation \( x \) is an element of the set \( X_i \), \( y_i \) representing the observation set will not be required to be an element of that set. This is because the probabilistic transition (due to the noise) might corrupt the original signal source such that the best estimate does not lie within the observation set \( X_i \). The concept of quantization as estimation is the key to risk quantization, with the specific requirement of source estimation rather than input estimation. This will provide the necessary quantization schemes for noise corrupted systems, while incorporating the classical quantizer definition of input estimation for the noiseless case.

To determine an estimation rule, some form of quality measure is required. This may be given by a cost function \( C(\theta, y) \) over \( \Theta \times \mathcal{Y} \) which represents the cost associated with estimating the parameter \( \theta \) by decision \( y \), based on observation \( x \), \( \forall \theta \in \Theta, x \in \mathcal{X}, y \in \mathcal{Y} \). The cost is generally expressed as a function of the difference between the parameter and its estimate,

\[
C(\theta, y) = C(e = \theta - y)
\]

With the specified cost function, the risk is defined as the expected value of the cost:
Estimation theory techniques may be employed to determine the quantization mapping \( q(x) \) which minimizes the risk function.

IV. Minimax Risk Quantization

The risk theory approach has been applied to the additive noise problem depicted in Figure 1 for well defined \textit{a priori} source and noise distributions to obtain a minimum risk quantization solution [4]. It is shown in [4] that the risk approach is a viable one to quantizer design for such systems, and is equivalent to the classical result for noiseless systems.

It is often likely, however, that the quantizer design must be based on incomplete knowledge of the signal statistics. For example, a problem may arise when a quantizer is needed for a source where the statistics are fairly well known, but various unknown noise distributions can corrupt the source signal. Likewise, the noisy characteristics of a sensor may be well defined, but the exact probability distribution of the source signal to be quantized with the sensor unknown. These types of problems suggest the use of minimax estimation to
determine the quantization mapping. Minimax estimation seeks to minimize the maximum possible distortion by the estimator, and is directly applicable to the risk approach to quantizer design.

This is demonstrated by examining the problem of an additive noise source using risk theory and minimax estimation. Specifically, the source statistics and probability distribution will be assumed known a priori. The exact distribution of the noise, however, will be assumed to be unknown, but belonging to a generalized moment constrained class of distributions, C. A risk analysis of the system will be applied using minimax estimation to produce a minimax risk quantization scheme for the system. Further, it will be shown that the analysis is reversible and may be applied to the additive noise problem when the noise probability distribution is well defined, but the knowledge of the source distribution is incomplete.

A minimax estimator \( t^* \) is one which guarantees a maximum risk no greater than that for any other estimator [11]. That is,

\[
\sup_{\theta} \mathcal{R}(\theta) \leq \sup_{\theta} \mathcal{R}(\theta) \tag{3}
\]

Likewise, the minimax quantizer will be defined as one which guarantees the minimum maximal risk for all quantizers. Minimax quantization theory has been
developed for the classical quantizer problem of noiseless sources by Bath and VandeLinde [1]. The approach developed will be applied in the risk theory development of an appropriate quantizer for the additive noise problem.

**Known Source Distribution**

Consider the sequence of independent, identically distributed random variables \( \{ \theta_k \} \), from the known cumulative probability distribution function (c.p.d.f) \( F_\theta \). The source sequence has been corrupted by a noise sequence \( \{ N_k \} \), with unknown c.p.d.f.s \( F_{N_k} \), to produce the observation sequence \( \{ X_k \} \) to the quantizer.

The sequence \( \{ F_{N_k} \} \), while unspecified, are constrained to belong to the set \( C \) of all possible c.p.d.f.s with whatever *a priori* information of the noise characteristics is available. The set to be considered here is the set of c.p.d.f.s belonging to the generalized moment constrained class. These distributions are required to have a generalized moment less than or equal to some finite constant. This may be viewed as a restriction of the noise power to be finite.
The minimax risk is defined as:

$$R^* = \min_{q \in Q_N} \max_{F_N \in \mathcal{C}} R(q, F_\theta, F_N)$$

(4)

where $R(q, F_\theta, F_N)$ is the risk for an $N$-level quantizer $q$ from the set of all possible $N$-level quantizers $Q_N$ and the c.p.d.f.s $F_\theta$, $F_N$, described above. The minimax risk quantizer, $q^*$, is that quantizer which provides the minimax risk $R^*$. The properties of the minimax risk quantizer, analogous to the minimax quantizer, are:

1) $\forall F_N \in \mathcal{C}, \exists R^* > R_q^*$

   where $R_q^*$ is the absolute maximal risk for quantizer $q^*$.

2) $\exists q \in Q_N, q \neq q^* \exists \max_{F_N \in \mathcal{C}} R(q, F_\theta, F_N) < R^*$

   where $q^*$ guarantees a maximum risk no greater than the maximum risk of any other quantizer $q \in Q_N$.

The quantizer mapping, $q(x) \in Q_N$, maps an $x \in X_j$ to a quantizer level $y_j \in Y$, $j = 1, \cdots, N$. For the scalar quantizer, $X_j = [x_i, x_{i+1})$, where $x_1 = -\infty$, $x_{N+1} = \infty$, and the quantizer thresholds comprise the set $\{x_i\}$, $i = 1, \cdots, N+1$. Note that the set of quantizers $Q_N$ does not
require the quantizer level $y_i$ be an element of $X_i$.

The risk for any particular quantizer mapping may be defined as:

$$ R(q, F_\theta, F_N) = \int \int C(\theta, q(x)) \ dF_N \ p_\theta(\theta) \ d\theta $$  \hspace{1cm} (5)

where the inner integral is a Lebesgue-Stieltjes integral over the sequence of noise c.p.d.f.s $\{F_{N_k}\}$. The order of integration is changed to yield:

$$ R(q, F_\theta, F_N) = \int \int C(\theta, q(x)) \ p_\theta(\theta) \ d\theta \ dF_N $$ \hspace{1cm} (6)

The purpose of the interchange is to facilitate the minimax operation procedure that occurs later.

It is apparent that if the system is noiseless, the risk function is precisely that of the classical quantizer distortion function. The minimax quantizer then is the same as the results of Max [9] since the source distribution is assumed known.

$$ R^* = \inf_q \int C(\theta, q(x)) \ p_\theta(\theta) \ d\theta, \ \theta = x $$

$$ \Rightarrow \ R^* = \inf_q \int C(\theta, q(x)) \ p_\theta(\theta) \ d\theta $$ \hspace{1cm} (7)

The cost function used is restricted to some bounded distortion measure, $d(\theta, q(x))$, between the source and quantizer output.
The distortion measure \( d(\theta, q(x)) \) is required to be even, continuous, monotonic strictly increasing in \( |\theta - q(x)| \), and zero for perfect estimation. The bound on the cost implies no additional penalty for a distortion larger than the limit, \( L \).

The generalized moment constraint is given by some function \( \rho(n) \) satisfying the same conditions as those for the distortion function such that:

\[
\int_{N} \rho(n) \, dF(n) \leq c. \tag{9}
\]

The constraint function \( \rho(n) \) simply implies that the noise signal power is bounded. The distortion and constraint functions considered here are:

\[
d(\theta, q(x)) = |\theta - q(x)|^{n} \tag{10}
\]

\[
\rho(n) = |n|^{m} \quad m \geq n \geq 1.
\]

The conditions of bounded cost and those imposed on \( d(\theta, q(x)) \) and \( \rho(n) \) are required for the Lagrange minimization of the next section.

The problem then, is to determine the minimax risk quantizer \( q^{*} \) for the worst case c.p.d.f., \( F_{N}^{*} \), which achieves:
The set $Q_N$ is the set of all symmetric quantizers where the levels are symmetric about the source mean, and the thresholds are symmetric about the observation mean. The set $C$ is the set of c.p.d.f.s belonging to the generalized moment class, and $R$ is the risk function using the cost function $C(\Theta, q(x))$. To determine the minimax risk quantizer, it first necessary to find the maximum risk due to a particular quantizer mapping $q \in Q_N$, then determine which quantizer provides the minimum of all of the maximum risks.

It has been shown [1] that the set $C$ is a weak* compact subset of a normed Banach vector space (N.B.V.) of normalized functions of bounded variation. Also with any linear functional on N.B.V. $[0, \infty]$ (such as $R(q, F_\Theta, F_N)$), which is weak* continuous in $F_N$, then:

$$\forall q \in Q_N, \exists F^*_N \in C, R^*_N(q) \exists:\$$

$$R^*_N(q) = R(q, F^*_\Theta F_N) = \max_{F_N \in C} R(q, F_\Theta, F_N)$$

(12)

Furthermore, a method for determining a minimax quantizer via a constrained minimization in a Lagrange multiplier space ($\mathbb{R}^2$) through the use of the Lagrange duality theorem has been developed [1]. The technique will be
paraphrased here to determine a minimax risk quantizer. Let $C = \{ F_N \in N.B.V.[0, \infty) \}$ where $F_N$ exhibits the following properties:

1) $F_N$ is nonnegative, monotonically nondecreasing.
2) $\int \limits_{N} dF_N \leq 1,$
3) $\int \limits_{N} \rho(n) F_N(n) \leq c.$

Define the convex set $\tilde{C} = \{ F_N \in N.B.V.[0, \infty) \}$, where $F_N$ satisfies 1). Now define a convex functional

$$G: N.B.V.[0, \infty] \to \mathbb{R}^2,$$

$$G = \left[ \int \limits_{N} dF_N - 1 \right], \left[ \int \limits_{N} \rho \, dF_N - c \right]$$

which represents the necessary constraints for $F_N$ to be a c.p.d.f. and have a generalized moment constraint. This implies:

$$C' = \{ F_N \in \tilde{C}: G(F_N) \leq 0 \}$$

By the Lagrange duality theorem [8], an expression for the maximum risk due to the quantization mapping $q(x)$ may be found:
\[ R^*(q) = \sup_{F_N \in C} R(q, F_\theta, F_N) \]

\[
\sup_{F_N \in C} R(q, F_\theta, F_N) = \min_{\lambda_1, \lambda_2 \geq 0} \sup_{F_N \in C} \left\{ R(q, F_\theta, F_N) \right\}_1 \]

\[
- \lambda_1 \left( \int dF_N - 1 \right) - \lambda_2 \left( \int \rho dF_N - c \right)
\]

where the inner maximization is achieved for some worst-case distribution, \( F^*_N \in \overline{C} \), and the outer minimization by \( \lambda_1^* \) and \( \lambda_2^* \). If the cost weighting function is bounded, \( R^*(q) \) is finite and equation (6) can be used to replace \( R(q, F_\theta, F_N) \) of equation (13) to yield:

\[ R^*(q) = \min_{\lambda_1, \lambda_2 \geq 0} \left[ (\lambda_1 + \lambda_2 c) + B(\lambda_1, \lambda_2) \right] \]  

where

\[ B(\lambda_1, \lambda_2) = \max_{F_N \in C} \int I(n) dF_N(n) \]

\[ I(n) = \int_{\theta} C(\theta - q(x)) p_\theta(\theta) d\theta - \lambda_1 - \lambda_2 \rho(n) \]

The minimization of equation (14) need only be considered when \( B(\lambda_1, \lambda_2) \) is finite. This will only occur when \( I(n) \) is nonpositive. If \( I(n) \) is positive at any point, a sequence of noise random variables \( \{N_k\} \) with c.p.d.f.s \( \{F_{N_k}\} \) could occur such that as \( k \) gets large, the
integration tends to infinity.

If \( I(n) \) is required to be nonpositive, then the maximization of equation (15) over all possible \( F_N \in \bar{C} \) implies:

\[
B(\lambda_1, \lambda_2) = 0. \quad (17)
\]

Since \( F_{N_k} \) a constant is an element of \( \bar{C} \) for \( \{N_k\} \), it follows that if \( I(n) \) is made nonpositive, the maximum possible integration of equation (15) is zero.

The minimax risk for any quantizer \( q \in Q_N \) and the specified conditions then reduces to:

\[
R^*(q) = \min_{0 < I(n) \leq 0} \left( \lambda_1, \lambda_2 \geq 0 \right) (18)
\]

Figures 3 through 6 are a series of curves which depict the function \( I(n) \) for various quantizers with

\[
p_\theta(\theta) \sim N(0, 1), \quad \text{and distortion bounds of } L = 0.25, 0.75.
\]

These curves show the effects of the distortion bound on the function, and that it is quantizer dependent. Note the discontinuity near the quantizer threshold.

The minimax risk quantizer \( q^* \) then, is that quantizer which produces:

\[
R^* = \min_{q \in Q_N} \min_{\lambda_1, \lambda_2 \geq 0} (\lambda_1 + \lambda_2 c) \quad (19)
\]

This result is similar to that obtained for minimax
quantization by Bath and Vandelinde for the generalized moment constrained class of quantizer input distributions. It differs primarily in the function $I(n)$ which incorporates the cost of estimating the source by the quantization mapping and the a priori knowledge of the clean source probability distribution function, $F_0(\theta)$.

The minimization of equation (14) over all nonnegative $\lambda_1$, $\lambda_2$, and nonpositive $I(n)$ implies the maximum of $I(n)$ is zero. As can be seen in equation (16), the role played by $\lambda_1$ in $I(n)$ is merely that of a bias term. This implies that the minimization of equation (14) can be determined by solving:

$$\lambda_1^*(\lambda_2) = \max_{\theta} \left\{ \int_\Theta G(\theta, q(x)) \ p_\theta(\theta) \ d\theta - \lambda_2 \rho(n) \right\}$$

(20)

and minimizing over all possible $\lambda_2 \geq 0$. Maximization of $I(n)$ with respect to $\lambda_2$ and $\lambda_1^*(\lambda_2)$ for a specific quantizer $q \in Q_N$ will then yield the minimum $\lambda_2$ and the requisite $\lambda_1$ for $\max_{\lambda_1, \lambda_2 \geq 0} I(n) = 0$.

**Known Noise Distribution**

The minimax risk theory is easily modified for the converse problem. That is, the problem considered thus far
is that of quantizing an i.i.d. source, with known probability distribution, corrupted by a sequence of independent additive noise random variables each of uncertain distribution. If the noise sequence is i.i.d. with known c.p.d.f. $F_N$, but the source sequence $\{\theta_k\}$ is only known to be distributed with the c.p.d.f. sequence $(F_{\theta_k})$ from the set $C$, then the risk may be written:

$$ R(q,F_{\theta},F_N) = \int \int C(\theta,q(x)) p_N(n) \, dn \, dF_{\theta} \quad (21) $$

The minimax risk quantizer development for this problem parallels the development for the previous case. A similar solution is described as the minimax risk quantizer $q^*$ which yields the minimum maximal risk,

$$ R^*(q) $$

where:

$$ R^*(q) = \min_{\lambda_1, \lambda_2 \geq 0, \ I(\theta) \leq 0} (\lambda_1 + \lambda_2 c) \quad (22) $$

and

$$ I(\theta) = \int \int C(\theta - q(x)) p_N(n) \, dn - \lambda_1 - \lambda_2 \rho(\theta) \quad (23) $$

The same restrictions apply to the distortion and constraint functions, $d(\theta,q(x))$ and $\rho(\theta)$ as before. Likewise,
\[
\lambda^*_1(\lambda_2) = \max_{\theta} \left\{ \int_{\mathcal{N}} C(\theta, q(x)) \, p_N(n) \, dn - \lambda_2 \rho(\theta) \right\}
\]  
(24)

and the minimax risk quantizer \( q^* \) is specified by a procedure analogous to that of the previous section.

Figures 7 through 10 show the effects due to the constraint on the noise power and the distortion bound \( L \) on the function \( I(\theta) \). These appear roughly similar to \( I(s) \) in [1], differing through the knowledge of the noise for \( I(\theta) \).

Note that in the noiseless case, this risk definition coincides precisely with the distortion function of an unknown input sequence used by Bath and VandeLinde. The minimax risk quantizer then is identical to the minimax quantizer developed there for the classical quantizer distortion measure.

V. Results and Conclusions

The minimax risk quantizer determination procedure, is described by the following algorithm. No attempt has been made to optimize the computation procedure.

1) Select \( \lambda_2 \geq 0 \).

2) Determine the optimal \( \lambda_1 \) for the specified
\[ \lambda_2: \lambda_1^*(\lambda_2). \]

3) Repeat step 2) over all \( \lambda_2 \geq 0 \) to determine the minimum of \((\lambda_1^*(\lambda_2) + \lambda_2 c)\). This is the maximum risk for the quantizer \( q: R^*(q) \).

4) Repeat steps 1)-3) over all quantizers \( q \in Q_N \) to determine the minimum of the maximum risks: \[ R^* = \min_{q \in Q_N} R^*(q). \]

5) The minimax risk quantizer is that quantizer \( q^* \) which yields the minimax risk:
\[ R^*(q^*) = R^*. \]

It appears from the plots of \( I(n) \) that the areas of maxima occur at or within a \( \delta \)-neighborhood of the quantizer thresholds. A simulation has been performed based on this observation with the results appearing in tables 1 and 2. These results were determined by sampling \( I(n) \) in the region surrounding each threshold. The quantizer step size was fixed at 0.01 and the accuracy of the Lagrange minimization is such that the error in the estimate is at most \( 1 \times 10^{-7} \). The minimax risk quantizer with specific quantizer thresholds were found, and the minimum of these selected as the minimax risk quantizer.

Comparisons of the minimax risk quantizers to the
Max-Lloyd and minimum risk quantizers (G.S.+G.N.) are shown at the bottom of tables 1 and 2. The mean square power constraint, $c$, is fixed such that for the Gaussian noise case, the Signal-to-Noise Ratio (SNR) is 8 dB. This value is shown in table 1. At 8 dB, the thresholds of all three are nearly identical, with the levels for the minimum risk quantizer inside (closer to the mean) those of the Max-Lloyd quantizer, and the minimax risk quantizer levels just slightly inside those of the minimum risk quantizer. The constraint $c$ is fixed for table 2 so that for the Gaussian noise case, the SNR is 4 dB. At 4 dB, the outer threshold of the minimax risk quantizer has migrated well outside of the other two quantizers, which may be from the Max-Lloyd and minimum risk quantizers assumption of Gaussian densities, while the minimax risk quantizer only considers constrained noise power. The levels of the minimax risk quantizer are now between those of the Max-Lloyd and minimum risk quantizers. This may be interpreted from the standpoint of the assumptions also. The minimum risk quantizer assumes both the source and noise are Gaussian, so it arrives at closely compacted estimates for the levels. The minimax risk quantizer, on the other hand, only assumes the source is Gaussian for this example, and the noise power constrained -- therefore the estimates are outside those of the minimum risk
quantizer (for this example). The minimax risk quantizer compensates for the corrupting influence of the noise, whereas the Max-Lloyd quantizer does not. For this reason, the estimates for the quantizer levels of the minimax risk quantizer are then placed within those of the Max-Lloyd quantizer.

References


TABLE I

Minimax Quantizer Determination
\((N = 4, \text{Squared-Error Cost, } p_\theta \sim N(0,1))\)

\(\mu_N = 0, \quad L = 0.25, \quad \text{SNR} = 8\text{dB} \Rightarrow c = 0.1585\)

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<th>(Y_1)</th>
<th>Minimax-Risk</th>
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Max-Lloyd Minimum Risk* Minimax Risk

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<th>(X_i)</th>
<th>(Y_i)</th>
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<td>1.298</td>
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*Minimum Risk for Gaussian Sources.
TABLE 2
Minimax Quantizer Determination
\((N = 4, \text{ Squared-Error Cost, } p_0 \sim N(0,1))\)
\(\mu_N = 0, \quad L = 0.25, \quad \text{SNR} = 4\text{dB} \quad c = 0.398\)

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Max-Lloyd Minimum Risk* Minimax Risk
\[
\begin{array}{ccc}
 x_i & y_j & x_i & y_j & x_i & y_j \\
 0 & 0.5354 & 0 & 0.3830 & 0 & 0.41 \\
 1.160 & 1.785 & 1.160 & 1.277 & 1.35 & 1.57 \\
\end{array}
\]

*Minimum Risk for Gaussian Sources.
Additive Noise Problem

Figure 1
Estimation Problem

Figure 2
Figure 3

PLOT OF I(N)

Quantization:
0.0 0.25
0.5 0.75

$\lambda_2 = 0.1$

$\sigma_\theta = 1.$

$L = 0.25$
Figure 7

PLOT OF I(THETA)
Figure 8

PLOT OF I(\Theta)

Quantiles:
0. 0.5
1. 1.5

\lambda_2 = 0.05

\alpha = 1.

1. 0.25
Figure 9

PLOT OF I(THETA)

Quantizer:
0. 0.8
1. 1.5

$\lambda_2 = 0.05$

$\sigma_N = 0.25$

$\lambda = 0.75$
PLOT OF I(THETA)

Figure 10
END

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