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AN ANALYTICAL APPROACH TO THE DESIGN OF SPECTRAL MEASUREMENTS IN THE DESIGN OF MULTISPECTRAL SENSOR

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ABSTRACT

The purpose of the research which led to this paper is to develop an analytical procedure for the design of the spectral channels for multispectral remote sensor systems. An optimum design based on the criterion of minimum mean-square representation error using the Karhunen-Loeve expansion was developed to represent the spectral response functions from a stratum. From the overall pattern recognition system perspective the effect of the representation accuracy on a typical performance criterion, the probability of correct classification, is investigated. Although the analytical technique was developed primarily for the purpose of sensor design it was found that the procedure has potential for making important contributions to scene understanding. It was concluded that spectral channels which have narrow bandwidths relative to current sensor systems may be necessary to provide adequate spectral representation and improved classification performance. The optimum sensor design provides a standard against which sub-optimum operational sensors can be compared.

I. INTRODUCTION

A pattern recognition system as used in a remote sensing system for earth resources consists of three fundamental components - the scene, the sensor, and the processor (Figure 1). The scene is that portion of the earth's surface observed by the sensor. The desired information is contained in the spectral, spatial, and temporal variations of the electromagnetic energy emanating from the scene. The sensor collects the energy and measures its features. The processor is typically a digitally implemented classification algorithm which makes an appropriate decision based on the feature measurements provided by the output from the sensor. Various types of ancillary data are also now typically used in the decision making process.

At present the design of the processor algorithms is quite advanced and provides variety and flexibility for optimal performance given a feature set (Fukunaga, 1972; Duda and Hart, 1973). However, the design of the best set of features is a complex matter which is not well understood. In the current work we limit considerations to the design of the spectral aspects of features to make the problem more tractable, leaving other aspects to later occasions. We will attempt herein to advance the knowledge of the spectral representation of the scene and to provide an analytical basis for the design of spectral channels for an operational sensor.

II. SPECTRAL REPRESENTATION AND OPTIMUM SENSOR DESIGN

The multispectral scanner is a complex system that is designed infrequently and has very little built-in flexibility. It must serve a wide variety of users with little opportunity for specialization. As a result careful consideration must be given to the specification of the parameters in the sensor design. The parameters may be grouped into five categories - spectral representation, spatial representation, signal-to-noise ratio, ancillary data or information not contained in the spectral response function itself, and the information classes desired (Landgrebe, 1978). As we have indicated, the principal category of parameters dealt with here is that of the spectral representation, since the goal is to select spectral bands; however, all of the parameters are interrelated such that specifying one of them places constraints on the others.

The ultimate goal is to optimize the overall system performance with respect to some criterion. A typical criterion, and the one used here, is the probability of correct classification, an intricate function of the parameters.
Let us begin by considering the information bearing aspects of the spectral response function \( x(\lambda) \) (Holmes and MacDonald, 1969). This response function is proportional to the electromagnetic energy received by the sensor as a function of wavelength \( \lambda \) (Figure 2). Many factors determine the spectral response function for a given observation. The irradiance of the sun, the conditions of the atmosphere, and the reflectance of the surface features are all known to have important effects on the response. Since a deterministic relationship between the response function and the factors affecting it would be very complex, the set of functions which are received will be modeled as a stochastic process.

The ensemble of the stochastic process (Papoulis, 1965) will be defined in terms of the stratification necessary to apply pattern recognition methods to the earth observational problem. A stratum, \( S \), is defined as the largest contiguous area which can be classified to an acceptable level of performance with a single training of the classifier. It is noted that the sensor must be designed to operate satisfactorily over a large number of such strata, which vary greatly with time, location and application. The collection of all possible strata which a sensor may observe is denoted by \( S \). Since the set \( S \) is quite large, it is necessary to select a smaller subset which is representative in a statistical sense in order to perform the analysis.

The random experiment for the stochastic process consists of the observation of a point in a stratum \( S \). Each point in the stratum is mapped into a spectral response function (Figure 3). The collection of all response functions from a stratum defines an ensemble. The ensemble plus the corresponding probability measure defines the stochastic process (Papoulis, 1965). It is appropriate to assume for this process a Gaussian probability measure (Crane et al., 1972).

It is necessary to choose a mathematical model for the sensor to represent the spectral response function for each observation. Let the sensor consist of a set of \( N \) filter functions or basis functions \( \{ \phi_1(\lambda) \} \) such that the output of each filter is given by (Figure 4)

\[
x_1 = \int_A x(\lambda) \phi_1(\lambda) d\lambda
\]

(1)

The output of the sensor model is a sequence, \( x_1, x_2, \ldots, x_N \), which represents the spectral response by the approximation

\[
x(\lambda) \approx x_1 \phi_1(\lambda) + x_2 \phi_2(\lambda) + \ldots + x_N \phi_N(\lambda) = \tilde{x}(\lambda)
\]

(2)

For a small number of simple functions, the approximation is very coarse (Figure 5). It is desired to choose a (ordered) set which will be optimum in some sense over a stratum. For the purpose of achieving an optimal set, no restrictions will be placed on the physical realizability of \( \{ \phi_1(\lambda) \} \).

A key consideration is the choice of the criterion for optimality. Because the sensor must function over a varied collection of strata using any of a large collection of classifiers, a criterion was chosen which is a measure of the fidelity with which the output of the sensor represents the input. We will choose the set \( \{ \phi_1(\lambda) \} \) such that for a given \( x(\lambda) \) the approximation \( \tilde{x}(\lambda) \) is as close as possible to the true spectral response function. Since the uses and therefore the specific spectral attributes needed by the various users cannot be predicted, this approach insures that all of the information in \( x(\lambda) \) will still be available in and recoverable from the \( \{ x_1 \} \) produced by the sensor. A common criterion for representation accuracy is the expected mean-square representation error given by

\[
E\{ \varepsilon \} = E \left\{ \int_A [x(\lambda) - \tilde{x}(\lambda)]^2 d\lambda \right\}
\]

(3)

However, it is desirable at this point to generalize this criterion by introducing a weight function \( w(\lambda) \) on the spectral interval. As will be seen presently, the weight associated with each \( \lambda \) can be used to introduce a priori knowledge concerning the spectrum into the analysis. Thus equations (1) and (3) become, (Wiersma, 1979).

\[
x_1 = \int_A x(\lambda) \phi_1(\lambda) w(\lambda) d\lambda
\]

(1a)

\[
E\{ \varepsilon \} = E \left\{ \int_A [x(\lambda) - \tilde{x}(\lambda)]^2 w(\lambda) d\lambda \right\}
\]

(3a)

We want to choose the set of basis functions \( \{ \phi_1(\lambda) \} \) which is optimum with respect to the spectral representation criterion of expected mean-square error \( E \{ \varepsilon \} \). More specifically, it is desired the representation be complete in the sense that the expected mean-square error for any function in the ensemble be made arbitrarily small simply by including enough terms, that convergence of the approximation to the original response be rapid in the first few terms, and that the basis functions be orthogonal to each other.

A technique for determining the set of optimum basis functions for an ensemble which satisfies the desired properties is based on the weighted Karhunen-Loeve expansion. (Davenport and Root, 1958; Van Trees, 1968; Ash, 1967; Wiersma, 1979). The solution to the homogeneous linear integral equation

\[
\gamma_1 \phi_1(\lambda) = \int_A K(\lambda, \xi) \phi_1(\xi) w(\xi) d\xi
\]

(4)

with the covariance function of the stochastic process, \( K(\lambda, \xi) \), as kernel is a set of eigenfunctions \( \{ \phi_1(\lambda) \} \) with corresponding eigenvalues.
If the eigenvalues are arranged in descending order, the corresponding sequence of eigenfunctions can be used to form a sequence of linear combinations of the eigenfunctions which converges to the original spectral response function with arbitrarily small expected mean-square error. Furthermore, because of the ordering of the eigenvalues, convergence in the first few terms is very rapid. This rapid convergence allows one to truncate the series expansion after a finite number of terms N with mean-square error minimized over all possible choices of N basis functions. The mean square error is given by

$$E(e_r) = \sum_{i=N+1}^{\infty} Y_i$$

Since the Karhunen-Loeve expansion is a well-studied technique and satisfies the desired properties for finding the basis functions, it provides a sound analytical method for determining the optimum set of basis functions.

It became apparent during this research that it may be advantageous to incorporate certain a priori information about the spectral interval into the analysis. The generalization of the Karhunen-Loeve expansion to include a weight function provided a convenient and appropriate means to incorporate such a priori knowledge into the design process.

The optimum sensor design problem may be solved on a digital computer using empirical data taken by field measurements. Some approximations must be made in order to take into consideration some practical constraints. First the response functions are not available as continuous functions but are obtained in the field by sampling the spectrum with an instrument that uses very narrow spectral windows. Secondly, the parameters of the process are not known a priori; hence, it is necessary to estimate the mean and covariance functions using a representative sample from the ensemble. Finally, because the data will be stored and processed digitally it is necessary to quantize the amplitude of the response at each of the spectral sample points. Each of these constraints can potentially contribute to the representation error. It has been shown that with reasonable care in selecting a sufficiently high spectral sampling rate, a large enough sample from the ensemble, and a large number of quantization intervals that the contribution of these factors to the representation error is small (Wiersma, 1979). The integral equation (4) becomes the matrix equation

$$\Phi' = KW\Phi$$

where \(\Phi\) is the matrix of eigenvectors, \(\Gamma\) is the diagonal matrix of eigenvalues, \(K\) is the covariance matrix and \(W\) is the diagonal matrix of weight coefficients.

III. RELATIONSHIP BETWEEN THE SPECTRAL REPRESENTATION AND SYSTEM PERFORMANCE

The performance of the overall system is ultimately what we wish to optimize. For remote sensing problems, an often used criterion for performance is the probability of correct classification \(P_c\). If the vector \(X\) is an observation from one of \(M\) classes \(C_i, i = 1, 2, \ldots, M\) with a priori probabilities \(P_i\), the probability of correct classification, using the maximum likelihood rule is given by

$$P_c = \int\max_i \{P_i p(X|C_i) dX\}$$

where \(p(X|C_i)\) is the conditional joint probability density function for class \(i\). The integral in (7) is over the observation space \(\Omega\).

The analytical procedure based on the weighted Karhunen-Loeve expansion has prescribed a sensor design which minimizes the mean-square error. One would like to know how the ability to represent a process influences the classification performance. To study this relationship the graph of the probability of correct classification vs. expected mean-square error is introduced (Figure 6). We will briefly discuss some of its characteristics.

The addition of terms to the series expansion causes a decrease in the spectral representation error, but the effect of the additional terms on the overall system performance has to be determined. It can be shown that increasing the number of terms in the representation will never decrease the performance provided that the stochastic process is completely known. If after \(N\) terms the improvement in performance is small compared to the reduction in representation error, then the representation is sufficient. This is illustrated by case A of Figure 6 in which the threshold \(T\) indicates the minimum required \(E(e_r)\). However, if the performance is showing significant improvement for a small decrease in the mean-square error, case B of Figure 6, more terms are necessary to complete the representation.

Since the parameters of the stochastic process must be estimated from a sample of the ensemble, the effect of the size of the sample relative to the dimensionality of the system becomes important. Hughes (1968) has shown that if the sample size is too small, the classification performance may actually be degraded by adding terms to the representation. Thus it is necessary to maintain a large set of sample functions from which to estimate the statistics.
The choice of information classes also influences the performance of the pattern recognition system. For purposes of classifying the data into distinct classes it is required that the class list have the following properties simultaneously (Landgrebe, 1978):

- Each class must be of interest to the user, i.e. of informational value.
- The classes must be separable in terms of the features available.
- The list must be exhaustive i.e. there must be a class to which it is logical to assign each pixel in the scene.

The classes may be arranged in a hierarchical tree structure such that classes deeper in the tree require more accurate representation to achieve a given level of classification performance.

The area of the ground resolution element, which is determined by the instantaneous field-of-view (IFOV), the altitude of the sensor, the scan rate, and the velocity of the sensor, are examples of spatial representation parameters. The size of the objects which can be identified and the energy available are influenced by the choice of ground resolution element size. If a typical object which one wishes to identify is smaller than the ground resolution element size, then it is very difficult to classify that object. Mobasseri (1978) has investigated the effect of the area of the resolution element on classification performance. Increasing the area often improves the signal-to-noise ratio which in turn improves the classification performance.

For a given remote sensing problem the signal is the part of the received response which is information bearing, and the noise is that part which is non-information bearing. The influence of the signal-to-noise ratio where the noise is white, Gaussian and additive was demonstrated which is non-information bearing. The influence by (Ready et al., 1971). Results show that overall classification performance decreased with an increase in the noise level. A class which was difficult to identify under low noise level conditions suffered the most degradation when noise was added.

IV. EXPERIMENTAL SYSTEM

An experimental software system has been set up to implement the analytical procedure that has been developed. The software system has been implemented on an IBM 370 computer at the Laboratory for Applications of Remote Sensing (LARS) at Purdue University.

A collection of field data consisting of spectral response functions on three dates from Williams County, North Dakota and three dates from Finney County, Kansas was available from the field measurements library at LARS. More than one thousand spectra were available from each location and collection date. The response functions were sampled in wavelength using narrow windows of .02μm.

The optimum set of basis functions is found numerically by estimating the covariance matrix from the sample response function. Maximum likelihood estimates of the mean and covariance matrices are given by

\[ \hat{\mu} = \frac{1}{N_s} \sum_{i=1}^{N_s} x_i \]  

and

\[ \hat{K} = \frac{1}{N_s} \sum_{i=1}^{N_s} (x_i - \hat{\mu}) (x_i - \hat{\mu})^T \]  

where \( N_s \) is the number of sample functions available and \( x_i \) is the \( i \)th sample vector. The covariance is the kernel in the linear integral equation whose solutions are the optimum basis functions or eigenvectors. A remarkably stable and accurate method of numerically computing the eigenvalues and eigenvectors was published by Grad and Brebner (1968).

The eigenvectors are used to perform the linear transformation

\[ y_i = \phi_i^T (X - \bar{X}) \]  

on the data set. The class conditional statistics are computed using the transformed data.

In order to compare the performances of two systems an algorithm which estimates the probability of correct classification for an \( M \) class problem given the class conditional multivariate Gaussian statistics was used (Wiersma, 1979). This algorithm, which is based on the stratified posterior estimator, (Whitsitt and Landgrebe, 1977) was found to be accurate within one-half of one percent.

The experimental system also included the ability to simulate (suboptimal) practical sensors. Although nearly any sensor characteristic could be simulated, most of the sensors which were simulated consisted of a small set of rectangular basis functions, i.e.

\[ \psi_k(\lambda) = \begin{cases} 1.0 & \lambda_k \leq \lambda \leq \lambda_{k+1} \\ 0.0 & \text{otherwise} \end{cases} \]  

where the \( \lambda_k \) are endpoints of the spectral channels. The endpoints of two suboptimum sensors which were implemented are listed in Table 1.

V. RESULTS

One of the first tasks in using the spectral parameter design system was to select a weight function. The uniform weight function of Figure 7 was tried first implying that no knowledge about signal and noise regions is available a priori. Plots of the first four eigenvectors for this weight function are shown in Figure 8. It was observed that the eigenvectors were dominated in several cases by components in the bands near
The dimensionality can be defined as the number of terms necessary to represent the original waveform to the desired accuracy. For the information classes used on the six data sets the dimensionality was about six to eight. Some strata required fewer terms to obtain a good representation and good performance, while others required more terms.

The graph of classification performance as a function of expected mean-square error proved valuable for studying the relationship between representation accuracy and classification performance for the family of functions \( \bar{\xi}_n(\lambda) \). It is possible to see which terms contributed to both the representation and the classification performance. By examining the asymptotic performance as \( E(c_n) \) approached zero, estimates of the maximum classification performance for the June 29 data set the maximum value of probability of correct classification was about 0.96, which is the value for 10 optimal features.

Each of the optimum basis functions was ranked according to its ability to classify points in the stratum. The ranking was based on single feature classification performance and multiple feature performance using divergence calculations and the performance estimator. The rankings for the June 29 data set are in parenthesis in Figure 11. It was found that the ranking on performance is not drastically different from the ranking based on representation accuracy. In general the first five eigenvectors tend to have a higher ranking than eigenvectors six through ten.

The eigenvectors were used to evaluate the spectrum to determine effective ways of sampling. It was observed that the first few eigenvectors had relatively wide subintervals. Eigenvectors later in the sequence exhibited high frequency (i.e. high spectral resolution) variations in magnitude. Examining several of the performance vs. representation curves it was found that these later terms were often significant thus indicating a need for high resolution sampling bands in certain portions of the spectrum. In particular it was found that narrow bandwidths on the order of .02 to .05 micrometers were needed in the visible region between 0.60 and 0.70 \( \mu \mbox{m} \).

The primary purpose of designing an optimal sensor was to use it as a standard to compare and evaluate practical sensor systems. With the present state-of-the-art in sensor system design functions such as those in Figure 11 are not practical. Practical designs such as the ones described by equation 11 and Table 1 can be evaluated and their performances compared to the optimal system. Based on the comparisons, one can make a decision as to whether the candidate design is satisfactory or some modifications need to be made.

The comparisons between the two simulated sensors of Table 1 and the optimum set were made on the basis of representation accuracy (Table 2) and classification performance (Figure 13).
Sensors 1 and 2 provide much poorer representation accuracy chiefly because they do not represent the whole spectral interval. For the information

Table 2. Comparison of Expected Mean-Square Error (in relative units) for Data Taken Over Williams County, North Dakota on June 27, 1977

<table>
<thead>
<tr>
<th>Sensor</th>
<th>Error (in relative units)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensor 1</td>
<td>17320</td>
</tr>
<tr>
<td>Sensor 2</td>
<td>16380</td>
</tr>
<tr>
<td>First 4 optimal basis functions</td>
<td>26.31</td>
</tr>
<tr>
<td>First 6 optimal basis functions</td>
<td>11.37</td>
</tr>
<tr>
<td>First 10 optimal basis functions</td>
<td>5.253</td>
</tr>
</tbody>
</table>

classes used for this data set much of the information in the response is not required for discrimination between the classes. The improvement in representation by sensor 1 over sensor 2 is very significant. The optimal sensor consists of the first 10 eigenvectors, where 10 provides a small mean-square error yet keeps the computational problems involved with high dimensional systems to a reasonable level. As shown in Figure 13 the four band sensor 1 compares very poorly with the best four optimal $\psi_i$, showing sensor 1 to be far from optimal for these classes. On the other hand the six band sensor 2 compares very favorably with the best six optimal $\psi_i$, the difference in estimated $\hat{P}$ being certainly less than the uncertainty of $P$ estimation. The slight further improvement of the first ten optimal $\psi_i$ is shown for comparison. However, since there is a large difference in representation accuracy, this favorable comparison may not be true for a different set of information classes.

VI. CONCLUSIONS

An analytical procedure has been developed for the design of the spectral characteristics for a multispectral remote sensor system. The procedure provides a standard for comparison of suboptimum sensors and a basis for the selection of spectral channels for operational sensor systems.

Significant contributions to the understanding of the scene were gained from this investigation. In particular knowledge about important spectral bands was extracted and efficient spectral sampling techniques were developed. Determination of scene properties such as signal dimensionality and maximum probability of correct classification was performed.

VII. REFERENCES


Figure 1. Pattern Recognition System

Figure 2. Spectral Response Function for Mature Wheat Collected on August 4, 1977 Over Williams County, North Dakota
Figure 3. Realization of a Stratum as the Ensemble of Spectral Sample Functions

\[ x[\lambda] = x_1 \Phi_1[\lambda] + x_2 \Phi_2[\lambda] + ... + x_n \Phi_n[\lambda] \]

Figure 4. Sensor System Model

Figure 5. Approximation of the Spectral Response Function by a Set of Four Basis Functions
Figure 6. Probability of Correct Classification as a Function of Expected Mean-Square Representation Error

Figure 7. Weight Function Number 1

Figure 8. First Four Optimum Basis Functions Using Weight Function Number 1 Over Williams County, May 8, 1977 Data

Figure 9. Weight Function Number 2

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Figure 10. Expected Mean-Square Error as a Function of the Number of Terms in the Karhunen-Loeve Expansion for Williams County, June 29, 1977

Figure 11. First Eight Eigenvectors for Williams County, June 29, 1977
Figure 12. Estimate of Probability of Correct Classification vs Expected Mean-Square Error for Williams County, June 29, 1977

Figure 13. Comparison of Probability of Correct Classification for Several Sensors for Williams County, North Dakota, June 29, 1977