

## A Multiple-Map Model for Pattern Classification

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A characteristic feature of vertebrate sensory cortex (and midbrain) is the existence of multiple two-dimensional map representations. Some workers have considered single-map classification (e.g. Kohonen 1984) but little work has focused on the use of multiple maps. We have constructed a multiple-map classifier, which permits abstraction of the computational properties of a multiple-map architecture. We identify three problems which characterize a multiple-map classifier: classification in two dimensions, mapping from high dimensions to two dimensions, and combination of multiple maps. We demonstrate component solutions to each of the problems, using Parzen-window density estimation in two dimensions, a generalized Fisher discriminant function for dimensionality reduction, and split/merge methods to construct a "tree of maps" for the multiple-map representation. The combination of components is modular and each component could be improved or replaced without affecting the other components. The classifier training procedure requires time linear in the number of training examples; classification time is independent of the number of training examples and requires constant space. Performance of this classifier on Fisher's iris data, Gaussian clusters on a five-dimensional simplex, and digitized speech data is comparable to competing algorithms, such as nearest-neighbor, back-propagation and Gaussian classifiers. This work provides an example of the computational utility of multiple-map representations for classification. It is one step towards the goal of understanding why brain areas such as visual cortex utilize multiple map-like representations of the world.

### 1 Introduction

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One of the most prominent features of the vertebrate sensory system is the use of multiple two-dimensional maps to represent the world. The observational data base for cortical maps is excellent, and this area represents one of the better-understood aspects of large-scale brain architecture. Recently, through the use of a system for computer-aided neu-

roanatomy, we have been able to obtain high-precision reconstructions of primary visual cortex map and column architectures, have constructed accurate models of both columnar and topographic architecture of primary visual cortex, and have suggested several computational algorithms which are contingent on the specific forms of column and map architecture which occur in this first visual area of monkey cortex (Schwartz et al. 1988). We expect to be able to extend these methods and ideas to other cortical areas. There is thus good progress in the areas of measuring, modeling, and computing with single-map representations. However, the problem of how to make use of *multiple* maps has been little explored.

Other workers have considered the application of single-map representations to classification. Kohonen (1984) has developed an algorithm for representing a feature space in a map; this algorithm constructs a space-variant representation, in rough analogy to the space-variant nature of primate visual cortex. However, this work does not provide a computational model for computing with multiple maps.

We believe that a classifier utilizing a multiple-map architecture must incorporate the following modules:

An efficient algorithm for classification in two dimensions.

A projection of high dimensional data into a two-dimensional representation.

An algorithm for combining multiple two-dimensional representations.

Our strategy in this work has been to use simple components to construct our multiple-map classifier. In particular, we were seeking algorithms which require one pass through the data and which are not sensitive to convergence issues (e.g. local minima in an energy function). We are interested in the overall properties of the classifier, and we are trying to deemphasize the role of the individual components, which are modular and hence subject to improvement or replacement.

## 2 Classification in Two Dimensions

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We assume that the items, or *instances*, we wish to classify are represented as vectors  $x \in \mathbf{R}^d$ , where each component of  $x$  is a feature measurement. Each instance belongs to a class  $k$ . We also have a *training set*, a set of instances of known class (*training examples*). We refer to the set of training examples in class  $k$  as  $X_k$ . Our problem is to construct a set of discriminant functions  $f_k : \mathbf{R}^d \rightarrow \mathbf{R}$ ,  $k = 1, \dots, c$ . An arbitrary instance  $x$  is assigned to the class  $k$  for which  $f_k(x)$  is maximal.

Because the instances are represented as vectors, we can refer to the distance between an instance and a training example as  $\|x - x'\|$ . We compute discriminant functions

$$f_k(x) = \frac{1}{N} \sum_{x' \in X_k} g(\|x - x'\|), \quad (2.1)$$

where  $g(r)$  is some function which decreases as  $r$  increases. If we let  $g$  be a probability density (i.e. nonnegative and integrating to one over its support) this is the *Parzen-window* estimate (Parzen 1962) for the *a posteriori* density; i.e.  $f_k(x) \approx p(k|x)$ . Since this is the same term which is maximized in the Bayes classifier, our classifier performance approaches the Bayesian limit as the approximation above approaches the actual probability density. This algorithm is related to the nearest-neighbor classifier. Its principal novelty is to use maps to store  $f_k(x)$ . Then, given the training examples, we can compute  $f_k(x)$  by convolution in one pass.

For illustration, see figure 1. We depict a two-class, one-dimensional classifier. The “map” is simply a segment of the real line. The training examples are shown on the  $x$ -axis as boxes. The weighting function  $g(x)$  is a Gaussian function. The individual convolutions  $g(x) * \delta(x - x')$  are shown as dotted lines. The class-specific density estimates, which are also the discriminant functions, are shown as a solid and broken line, respectively.

We consider a two-dimensional, three-class problem in figure 2 and figure 3. The weighting function is a circular two-dimensional Gaussian function. The instances have been drawn from prespecified two-dimensional multivariate normal distributions; this permits construction of a Bayes classifier to determine minimal error rate. In figure 2, we show a comparison between the Parzen-window density estimates  $f_k(x)$  and the actual probability density functions for each class. In figure 3, the classifier is compared to a Bayesian classifier. The visual comparison indicates that the classifier is capturing much of the character of the Bayesian classifier. When the classifier was trained on 400 samples from each class, and tested on 300 (different) instances, its error rate was 16.0%, which may be compared to 14.4% for the Bayesian classifier.

One important issue in the application of this method is the choice of the weighting function (or kernel). We have typically used Gaussian kernels, in which case we need to choose the kernel variance  $\sigma^2$  (or covariance matrix  $\Sigma_m$  in higher dimensions). This is a difficult problem in general; we have used heuristic algorithms. For example, if we desire an isotropic kernel, we might use  $\sigma = N^{-1/m} \sqrt{\lambda_1}$ , where  $\lambda_1$  is the largest eigenvalue of the covariance matrix resulting from the projection of the data into an  $m$ -dimensional map. The factor  $N^{-1/m}$  arises from the heuristic decision to give each training instance an equal amount of map volume; since the kernel is  $m$ -dimensional, the volume scales as  $\sigma^m$ . More generally, we could use  $\Sigma_m = \frac{1}{N} P \Sigma$ , where  $P$  is the projection into the map (see below). In experimental studies, we have found that the performance of the classifier is insensitive to small changes in the kernel size or shape.

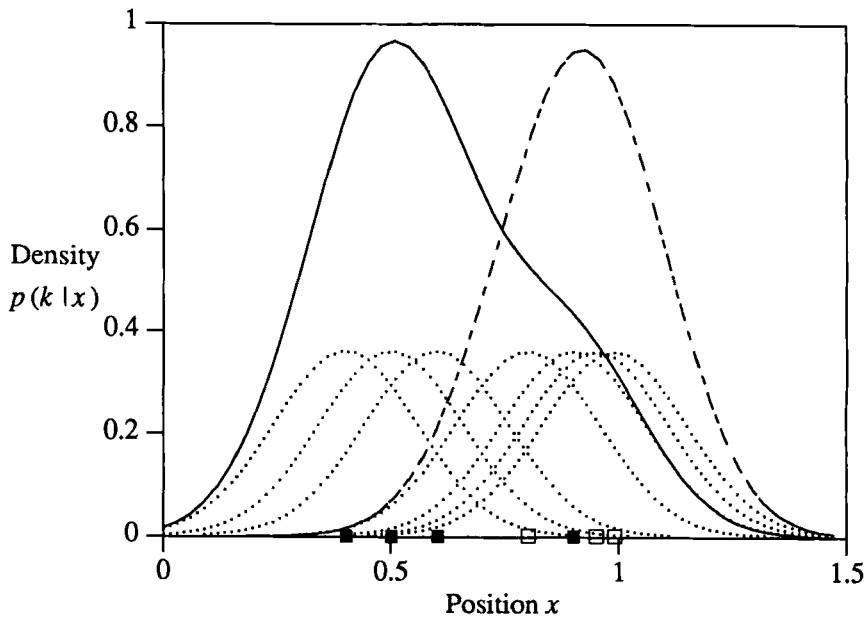


Figure 1: A one-dimensional two-class classifier. Class 0 instances are shown as solid boxes; class 1 instances are shown as open boxes. The estimated *a posteriori* density  $p(k|x)$  (here, for one-dimensional  $x$ ) is shown as a solid line for class 0, and a broken line for class 1. The Parzen-window function  $g(\|x - x'\|)$  for each paradigm  $x'$  is shown as a dotted line. The classifier operates by choosing the class for which the estimated *a posteriori* density is maximized. Thus, samples drawn with feature measurements below 0.7 would be assigned to class 0. Samples drawn with  $x > 0.7$  would be assigned to class 1.

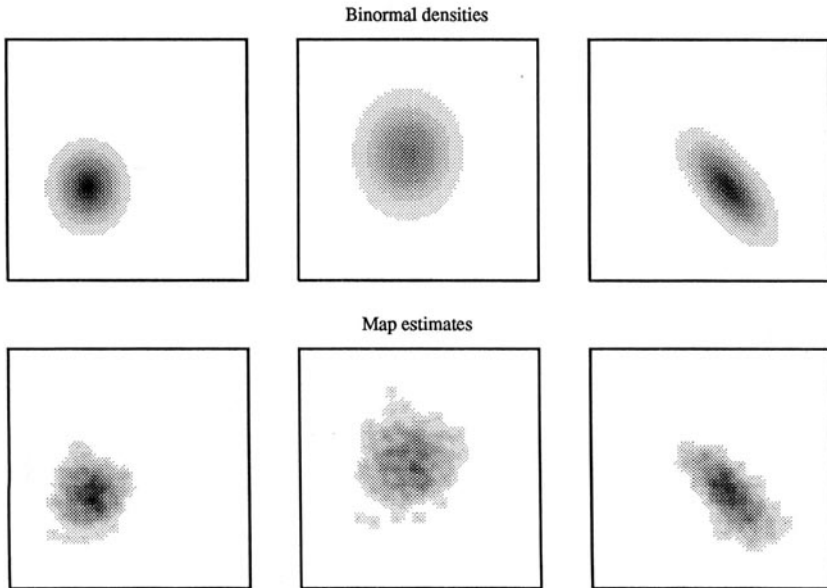


Figure 2: Comparison of actual binormal density against estimated density  $f_k(x)$  computed by our classifier. Here a higher density corresponds to a darker region of the plot. The top row shows density plots for three binormal distributions in  $R^2$ . The bottom row shows the estimates  $f_k(x)$  computed by the classifier for 400 samples drawn from each of these three distributions. The weighting function used is a circular Gaussian with a variance approximately  $1/60$  the width of the figure.

### 3 From $d$ Dimensions to Two Dimensions

In the previous section, we showed that the performance (as measured by error rate) approaches that of the Bayesian classifier for two-dimensional

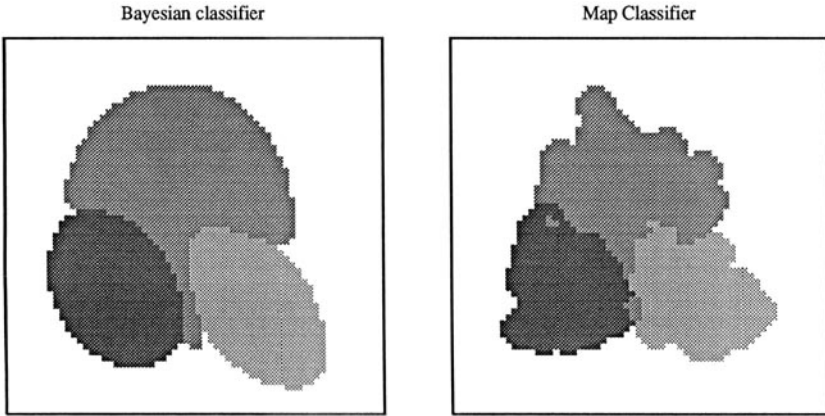


Figure 3: Comparison of decision regions computed by our classifier against decision regions of a hypothetical Bayes classifier which had complete knowledge of the underlying class distributions. Regions in the Bayes classifier are clipped due to round-off.

data drawn from Gaussian distributions. Actually there is nothing in our derivation which restricts us to two dimensions; a  $d$ -dimensional classifier is defined as above, except that the density estimate  $p_k(x)$  will require a  $d$ -dimensional map. In practice we limit ourselves to two dimensions for three reasons. First, our original motivation is to understand the functional utility of laminar structures such as neocortex for pattern classification in the brain. Second, two-dimensional maps can be processed

by conventional image processing software and hardware, and the user of the classifier gets the benefit of visual displays of the intermediate structures in the classifier (e.g.  $p_k(x)$  and the computed partition). Finally, the number of bins required to store the maps  $p_k(x)$  grows exponentially with the dimensionality  $d$ , and thus favors small  $d$ .

Restriction to two dimensions introduces an interesting aspect to the classification problem. Although our original data is in  $d$  dimensions, i.e. the data is composed of  $d$  measurements, we must somehow extract only *two* measurements or combinations of measurements with which to construct our classifier. The classification problem then spawns a problem of feature derivation. We can formulate the dimensionality reduction problem as construction of a function  $P : \mathbf{R}^d \rightarrow \mathbf{R}^2$  which maps  $d$ -dimensional instances to two-dimensional map positions. The two dimensions of the map constitute the two derived features. We need to specify what kind of function  $P$  we will allow. To date, we have only considered linear projections, but nonlinear functions could also be used (e.g. Kohonen's self-organizing feature map; Kohonen 1984).

We apply the generalized Fisher discriminant which was first introduced for a projection to  $\mathbf{R}$  (Fisher 1936) and later generalized to a domain of arbitrary dimensionality (Bryan 1951). A discussion of the technique may be found in (Duda and Hart 1973). The two vectors which comprise  $P$  turn out to be the eigenvectors associated with the two largest eigenvalues in the generalized eigenvalue system

$$S_b u = \lambda S_w u, \quad (3.1)$$

where  $S_w$  is the "within-class" scatter matrix, given by

$$S_w = \sum_{k \in \Omega} \frac{1}{N_k} \sum_{x' \in X_k} (x' - \bar{x}_k)(x' - \bar{x}_k)^T \quad (3.2)$$

with  $\bar{x}_k$  the class mean and  $N_k$  the number of training examples representing class  $k$ , and  $S_b$  is the "between-class" scatter matrix, given by

$$S_b = \frac{1}{N} \sum_{k \in \Omega} N_k (\bar{x}_k - \bar{x})(\bar{x}_k - \bar{x})^T \quad (3.3)$$

Here,  $\bar{x}$  is the mean over all the training examples. In practice, we have found that  $S_w$  is nonsingular, so the system can be reduced to a standard eigenvalue problem

$$S_w^{-1} S_b u = \lambda u. \quad (3.4)$$

The extraction of the principal eigenvalue and its eigenvector are realizable using a typical Hebb synapse model with a fixed-length weight vector (Oja 1982).

The performance of the discriminant can be observed with Fisher's classical iris data. This data describes a four-dimensional, three-class

problem. Figure 4 depicts the classifier constructed from the projection of the iris data into the two-dimensional subspace which maximizes the ratio described above. The classes can be seen to be fairly well separated. The classifier was tested by splitting the 150 instances into a 100-instance training set and a 50-instance test set. After training on the 100 instances, the classifier achieved 98% correct classification on the 50 instances in the test set. By comparison, the nearest-neighbor classifier operating on the same training and test sets achieved 98% correct classification, the Gaussian classifier achieved 94% correct classification, and a multilayer perceptron trained using back-propagation<sup>1</sup> achieved 96% correct classification.

#### 4 Using Multiple Maps: A Tree of Maps

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The previous example showed that for a relatively easy four-dimensional three-class problem, the generalized Fisher discriminant analysis was adequate to obtain a map which permitted good classifier performance. But in general, the discriminant analysis does not yield enough separation. For example, consider a regular five-dimensional simplex; this is a set of six equidistant points on the unit sphere in  $\mathbf{R}^5$ . Locate a spherical multivariate normal distribution at each vertex of the simplex. This is a point swarm whose density declines as  $\exp(-r^2)$ , where  $r$  is the distance from the vertex. We construct the classifier by utilizing discriminant analysis to find a projection  $P : \mathbf{R}^5 \rightarrow \mathbf{R}^2$ . With 600 training points and 300 test points, the error rate is 36%.

Fortunately, we are not confined to one map. One method of using multiple maps utilizes a split/merge technique to reduce one many-class problem to several problems, each with fewer classes. We merge the original *base* classes into *superclasses*, each of which is represented by the union of training examples from its underlying base classes. We then apply discriminant analysis to the newly formed superclasses. If we can achieve an adequate separation, we proceed as above with a separate map for each superclass. If necessary, we can again perform merges among the elements of a superclass, until we have divided each superclass into component base classes.

The consequence of this approach is to create a tree of maps. From the root of the tree, we project instances into a superclass. If the superclass is a base class, we assign that class to the instance. Otherwise, the instance is assigned to a superclass, which has its own map. We project the instance into that map, and continue as above, until the instance lands in a region assigned to a base class. In the training phase, we construct

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<sup>1</sup>A multilayer perceptron with 4 hidden units was trained using back-propagation (Rumelhart et al. 1986) with 5000 iterations through the training set, with  $\epsilon = 0.02$  and  $\alpha = 0$ .



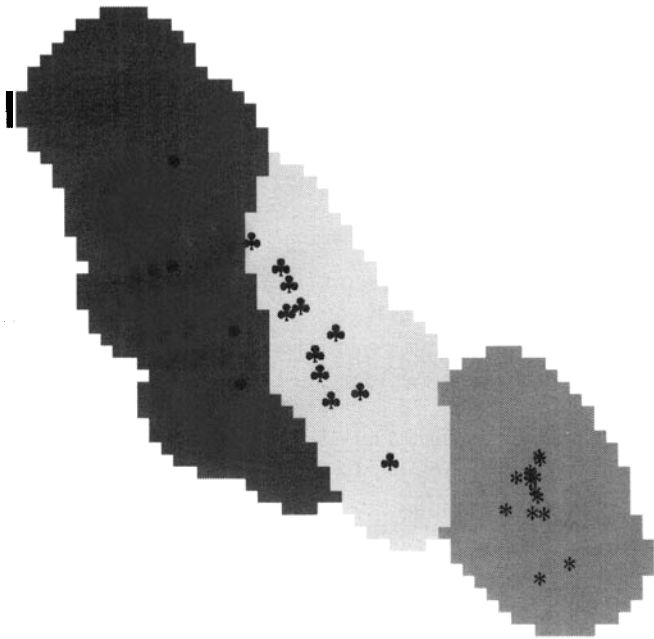


Figure 4: Iris classifier constructed from 100 training points drawn from the iris data (shaded background). Projection of four-dimensional iris data points into the two-dimensional subspace which maximizes the ratio of between-class variance to within-class variance (foreground data points).

a classifier in  $\mathbf{R}^2$  for each internal node in the tree to classify instances into one of the superclasses for that node.

We can illustrate this algorithm with the simplex data. We partition the classes so that three maps are used to classify. In the first map, we merge classes 2–5 into a superclass, letting classes 0 and 1 remain as base classes. In the second map, we will resolve the superclass composed of classes 2–5 into classes 2 and 3 and a superclass formed of 4 and 5. Finally, in the third map, we will resolve the superclass formed from 4 and 5 above into component base classes. The error rate of the three-map classifier is found to be 4.7%, a dramatic improvement over the single-map classifier (36% error rate). This may be compared to error rates of 2%, 6% and 2.7% respectively for the Gaussian, nearest-neighbor and

multilayer perceptron classifiers.<sup>2</sup> We have also applied our classifier to real-world data which consisted of 22 cepstral parameters from digitized speech.<sup>3</sup> Each of 16 data sets represented one speaker; seven classes (monosyllabic words) were present in each data set. Each set consisted of 70 training instances and 112 test instances. The results are summarized:

Classifier:	Multiple-map	Gaussian <sup>4</sup>	Nearest-neighbor	Multilayer perceptron <sup>5</sup>
Average error rate(%):	6.5	6.0	5.9	6.3
Range (%):	1.8–12.5	3.6–10.7	1.8–15.2	1.8–11.6

from which it may be seen that all four classifiers under consideration had closely comparable performance.

## 5 Automatic Generation of the Map Tree

In the preceding examples of multiple map usage, we interactively chose a map tree. In this section we explore a simple approach to automatic generation of the map tree. This is a *clustering* problem; we want to group classes into superclasses which in some way reflect the natural similarity between classes.

We introduce the distance matrix  $A$  for the classes. For any interclass distance measure  $\text{dist}(i, j)$ ,  $A_{ij} = A_{ji} = \text{dist}(i, j)$ . We use a very simple tree generation algorithm. We treat  $A$  as a graph with each class represented by a node, and each edge weighted according to interclass distance. We then compute the minimal spanning tree. We form superclasses by recursively removing the largest edge in the tree, yielding two subtrees, each of which forms a superclass. We can use a variety of interclass distance measures; we have experimented with distances between class means, overlap of the one-dimensional Fisher discriminant projections, and overlap of the two-dimensional Fisher discriminant projections.

<sup>2</sup>A multilayer perceptron with 7 hidden units was trained using back-propagation (Rumelhart et al. 1986) with 5000 iterations through the training set, with  $\epsilon = 0.02$  and  $\alpha = 0$ .

<sup>3</sup>We are grateful to R. Lippmann of MIT Lincoln Laboratory for providing this data.

<sup>4</sup>Covariance matrix estimates were obtained by pooling data from all 16 speakers for each class.

<sup>5</sup>Multilayer perceptrons for each speaker used 15 hidden units.

## 6 Discussion

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Our classifier is inspired by the prevalence of maps in the vertebrate brain. Its components are two-dimensional map units which implement Parzen-window density estimation, a dimensionality reduction methodology, and a scheme for decomposing a problem so that it can be solved by a system of maps. The training and running costs are favorably low. It admits an easy formal description. The intermediate results of classifier construction, e.g. the density estimates  $p_k(x)$  and the partition computed by the classifier, are easily observed by a human user. This allows insight into the structure of the data that is hard to gain from other algorithms. Many of the operations can be implemented with conventional image processing operations (and thus can take advantage of special-purpose image processing hardware). The error rate is comparable to popular parametric, nonparametric, and neural network classifiers.

Only a few other workers have considered the role of maps in pattern classification. In particular, Kohonen (1984) has considered iterative algorithms for "self-organizing feature maps." We wish to distinguish his work from ours. In our classifier, the map function comprises a linear projection of a data instance to determine a position in the map followed by a reference to that position. Kohonen maps an instance via a distance computation at each node of his map, followed by a winner-take-all cycle to obtain the nearest-neighbor to the instance among all the map nodes. The projection we use is computed with one pass through the training set to compute second-order statistics, which are diagonalized in a step which has a cost related only to the dimensionality of the data, but not the number of samples. Kohonen uses a very large number of iterations through the training set. Most importantly, we emphasize the use of multiple maps, which is not considered in (Kohonen 1984). We could use a Kohonen-type feature map as a module in our classifier (replacing the Fisher discriminant analysis) although we would then sacrifice these advantages.

Tree classifiers have been considered at length in Breiman et al. (1984). There are similarities between their classifiers and ours at classification time, although the training algorithms are quite distinct. The principal difference in classifier operation is that we use two-dimensional density estimation at each node, while they use one-dimensional linear discriminants. Their discriminant is typically a threshold comparison of one feature value, although they also describe an iterative technique for obtaining a discriminant from a linear combination of a subset of the feature variables. There are much larger differences in the classifier training algorithms; we present an example of a simple heuristic for generating map trees (based on minimal spanning trees) whereas they examine a large set of possible splits in the data to generate trees. We wish to emphasize that our tree classifier is one possible technique for utilizing multiple maps; examination of alternative approaches is an important research problem.

Perceptual (and probably cognitive) functions of the brain are mediated by laminar cortical systems. Three carefully investigated systems (monkey vision, bat echolocation, and auditory localization in the owl) are committed to multiple two-dimensional spatial maps. The present paper describes the first attempt to construct a pattern classification system which has high performance and which is based on a multiple parallel map-like representation of feature vectors. The algorithms described in this paper allow us to begin to investigate the pattern classification and perceptual performance of such map-based architectures.

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